# X-ray sociology

- more exact than NMR
- lots of Nobel prizes
- tells you where atoms are to a fraction (  $\frac{1}{5}$   $\frac{1}{4}$  ) of an Å
- can work on large structures
- Hamburg is full of crystallographers
- not all proteins / nucleotides are happy to crystallise



# **Summary of story**



# Topics

- tiny bit about chemistry / crystals
- calculating electron density
- placing coordinates in density

# **Forming crystals**

Familiar crystals – Kochsalz, sugar...

- crystal formation small molecules
  - rigid, regular
  - soup of unordered molecules  $\rightleftharpoons$  ordered crystal,  $\Delta G$  favourable

Proteins

- lots of internal mobility not rigid
- not nice regular shapes
- soup molecules  $\rightleftharpoons$  ordered crystal,  $\Delta G$  borderline Often can not be crystallised

## **Protein crystals**

- small, 1 to 100  $\mu m$
- not as well ordered as small molecules

# 

We measure an average over all molecules – consequence ..

• right hand side – the average is smeared

# Diffraction

#### Start with 1-dimension





Spacing of peaks on the detector ? Will depend on

- distance  $r_{AB}$
- wavelength  $\lambda$

What we measure (intensity *I*) will be

- periodic (cosinus) in *x*
- change slowly if  $\lambda$  is big

$$I = \cos\left(\frac{2\pi}{\lambda}x + \alpha\right)$$

- $\alpha$  phase important later
- relate this to proteins



# protein crystal and grids

- wavelengths are we talking about ? X-ray  $\approx 1 \text{ Å}$
- do we have grids ?

Work in 3D

- makes maths more difficult
- have to rotate and add up pictures
- we have lots of pairs of atoms (AB)
- X-rays interact with electrons
- hydrogen is almost invisible

What one measures

• sum over many pairs of electron clouds





# **Summing and Fourier transforms**

Simple formula in 1D and one pair

$$I = \cos\left(\frac{2\pi}{\lambda}x + \alpha\right)$$

- what we measure is a sum  $\sum_{reflections} \cos(...)$
- better nomenclature  $\Sigma_{hkl}$  *h*, *k*, *l* are indices of detected spots

#### How do I break a signal into frequencies?

- fourier transform think of pictures of media player
  - from sound in real time to picture of frequencies  $cos(\omega)$
- Can you just take a signal and fourier transform ?
- only if you know the  $\alpha$  's
- There is phase for every reflection, but you cannot measure it

# phase problem

- lots of "reflections" (10<sup>5</sup>)
- you measure their size / intensity
- you cannot measure their phase  $\alpha$
- if you know the  $\alpha$  for each reflection
  - just calculate density

Where do the phases ( $\alpha$ ) come from ?



# Ways to find phases

Can you do it directly?

- for very small molecules yes
- protein ?
  - imagine you have 10<sup>5</sup> reflections
  - try just 4 possibilities in each case
  - 4<sup>10000</sup> possibilities
- Strategies two most important
- 1. if you know some coordinates, substitute them into the equation gets a good starting point (molecular replacement)
- 2. if you know some phases, easy to get the next ones (multiple isomorphous replacement)

+ if you have a reasonable initial guess, it can be optimised

# molecular replacement

- most common <sup>3</sup>/<sub>4</sub> or more of structures in protein data bank
- you need to know some coordinates
- you do not know coordinates
  - you have the coordinates of something close (homologue)
- pretend these coordinates are responsible for the measurements
- substitute into formula and get an initial set of  $\alpha$  's
- refine

# multiple isomorphous replacement (MIR)

Remember

- for small molecules you can find phases directly (few reflections)
- if you know some phases, the next are easier

How to treat protein as if is a small molecule ?

- Heavy atoms have so many electrons they dominate the observations
- give a starting point for phases



- what are heavy atoms ? Au, Pt, Hg, Br, Se, Xe
- should bind at the same position in every protein molecule

# more phasing

How difficult?

• for many proteins – plenty of data but phases take years

Other methods ?

• yes

# resolution - meaning

- most common statistic for X-ray structures resolution
- meaning when do two points look like one?
- Resolution in X-ray.. depends on how scattered the waves are





# waves and resolution

- bottom path = lower path + wavelength  $\lambda$  then they reinforce each other
- difference in lengths is  $n \lambda$

 $n\lambda = \overline{\text{ABC}} = 2(d\sin\theta)$ 

• then  $d = \frac{n\lambda}{2\sin\theta}$ 

#### **Resolution consequences**

- smaller wavelength  $\lambda$ , the better
- angle  $\theta$  you cannot do much but..







In the centre .. reflections

- did not diffract much
- due to low resolution information
- strong

Further out

- high resolution information
- signal weak

# resolution practical

Mostly a function of crystal

- crystal not so regular and / or
- atoms mobile
  - you are seeing an average
  - there is no high resolution information



# 

### disorder - static and dynamic

static

# 

dynamic



model for this - soon

# model fitting - refinement

- you have initial electron density and initial phases
- have to fit atoms (build a model)
- what are the variables ?
  - *x*, *y*, *z* for atoms
  - *B*-factors (mobility) ... next slide
- Given atoms, you can calculate density
- given density, you can calculate reflections (structure factors *F*)

Refinement

 how well do the structure factors from your model agree with measurements ?



# **B**-factors

A model for atom location

• Gaussian (normal) probability



Why Gaussian ?

- pretend a particle moves in a harmonic well  $E(x) = x^2$
- from Boltzmann relation,  $p(x) \propto \exp\left(\frac{-x^2}{kT}\right)$

(formal in Sommersemester)

• *k* Boltzmann constant, *T* temperature



## **B**-factors interpretation

Probability distribution

- how likely are you see an atom at a position or
- what is the typical movement at room temperature ?

 $B = 8 \pi^2 u^2$  where *u* are fluctuations in Å  $u = \left(\frac{B}{8\pi^2}\right)^{\frac{1}{2}}$ 

if  $B = 50 \text{ Å}^2$ , typical displacement  $\approx 0.8 \text{ Å}$ if  $B = 20 \text{ Å}^2$ , typical displacement  $\approx 0.5 \text{ Å}$ 

The connection to fitting...

# fitting to density

The path to this picture...





- water molecules
- atoms have different
  - sizes
    - electron clouds
  - mobilities



## **Refinement – cost function –** *R*

The cost function

- atoms  $\rightarrow$  density  $\rightarrow$  structure factors (*F*)
  - $F_{hkl}^{calc}$  structure factors calculated
  - $F_{hkl}^{obs}$  structure factors observed



very important... *R* factor

$$R = \frac{\sum_{hkl} \left| F_{hkl}^{obs} - F_{hkl}^{calc} \right|}{\sum_{hkl} \left| F_{hkl}^{obs} \right|}$$

# Variables in refinement

What are the variables ?

- x, y, z for every atom +
- *B*-factors
- adding ions and water to fill density



refinement progress						
	data used	# par	# obs	Rw	Rf	
molecular replacement	15-4.0	3	775	44.5		
rigid body	10-2.5	9	2997	46.4	47.4	
first round	10-1.5	3887	13818	30.2	34.3	
after first		3735		26.2	31.4	Jata and J
SHELXWAT		4091		20.2	24.6	data used #nar number of
bld + SHELXWAT		4231		18.7	22.8	#par inumber of parameters in model
include all data	10-1.1	4203	33993	19.1	21.7	#obs number of
ANIS 20		9453		16.0	19.2	reflections
Rebuild SHELXWAT		9557		13.4	15.8	Rw <i>R</i> factor
rebuild		10481		12.3	15.3	Rf R <sub>free</sub> (soon)
rebuild		10819		12.1	15.1	
rebuild		10838		11.6	14.6	
rebuild		11494		11.3	14.4	
rebuild		11576		11.0	14.0	
rebuild		11774		10.7	13.8	

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ANIS 20		9453		16.0	19.2	
Rebuild SHELXWAT		9557		13.4	(15.8 <sup>6 A</sup> )	
rebuild		10481		12.3	15.3	
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- Start: Low resolution data is enough just a few parameters
- #par and #obs
  - where is the molecule ?
  - where is molecule + phases ?
  - add in first atoms
    - number of parameters grows at each step
    - add in more data (#obs)



refinement progress							
	data used	# par	# obs 0	Rw	Rf		
molecular replacement	15-4.0	3	775	44.5			
rigid body	10-2.5	9	2997	46.4	47.4		
first round	10-1.5	3887	13818	30.2	34.3		
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ANIS 20		9453		16.0	19.2		
Rebuild SHELXWAT		9557		13.4	15.8		
rebuild		10481		12.3 <i> </i>	R of bit more than		
rebuild		10819		12.1 2	20 % is typical		
rebuild		10838		11.6	14.0		
rebuild		11494		11.3	14.4		
rebuild		11576		11.0	14.0		
rebuild		11774		10.7	13.8		

11/27/2019 [32]

# What is happening in refinement

Similar problem to NMR refinement

- use a minimizer
  - move *x*, *y*, *z* and *B*-factors
    - until agree with experimental data  $(F_{hkl}^{obs})$
    - maintain known chemistry bond lengths angles
- Different to NMR
- more variables
  - *B*-factors
  - you can add water and ions to fill density
- more data
- search for a single solution (not many possibilities)



# **R** free - overfitting

Overfitting?

- you give me data
- true model is a line (y = ax + m)
  - two parameters
- I fit to a polynomial  $y = ax^4 + bx^3 + cx^2 + dx + m$
- apparently better fit
  - will not predict correctly

How would you detect this ?

- I have half a dozen points
- do fitting on five see how good the fit on the sixth point is In general fit on 90 % the data and test on remaining 10 %



X

	refinement progress					
	data used	# par		Rw	Rf	
molecular replacement	15-4.0	3	775	44.5		
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rebuild		11774		10.7	13.8	

Crystal data – also has noise

- you can always fit to noise
  - add water, ions, move atoms

How to detect?

- fit on 90 % of data
- calculate *R* on remaining data

$$R_{free} = \frac{\sum_{hkl} \left| F_{hkl}^{obs} - F_{hkl}^{calc} \right|}{\sum_{hkl} \left| F_{hkl}^{obs} \right|}$$



X

- but  $F_{hkl}^{obs}$  is from 10% of data not used in fitting
- typical values in protein data bank 20 to 30 %

# **Practical meaning of resolution**

- formally limit when two points become one point
- practical



best	< 1 Å	see H electrons
	1.2 Å	separate atoms
	2 Å	sidechains
worst	>4 Å	overall shape

bl831.als.lbl.gov/~jamesh/movies/

## resolution in PDB



11/27/2019 [38]

# **Errors, uncertainty**

#### Errors ? – Many

- good data you cannot tell O from N
- bad data you may slip by one amino acid
- spectacular errors
  - trace chain backwards, join wrong secondary structure units

#### Uncertainty

- if I have 2 Å data, what is my uncertainty ? Much smaller
  - 1. averaging over observations
  - 2. known bonds





#### coordinate error - synthetic data

Resolution near 1.5 to 2.0 Å

• errors 0.2 to 0.3 Å



estimates of coordinate error – not important for us



# missing atoms

Clear pieces of structure missing

Look at *B*-factors



# NMR vs X-ray

	NMR	X-ray
certainty	spread amongst 50 models	B-factors / Gaussian model
resolution	no meaning	well defined
size	rarely > 200 residues	big

## not for discussion - make sure the ideas are clear

- \* Forming crystals is a question of energy differences
- \* Simple refraction, one dimension
- \* generalising to 3D not discussed
- \* from reflections to density via Fourier Transform
- \* phasing methods
- \* high and low resolution reflections
- \* static vs dynamic disorder
- \* fitting, overfitting
- \* R and R free
- \* B-factors and missing atoms