# **Intrinsically Disordered Proteins**

Are proteins always structured ?

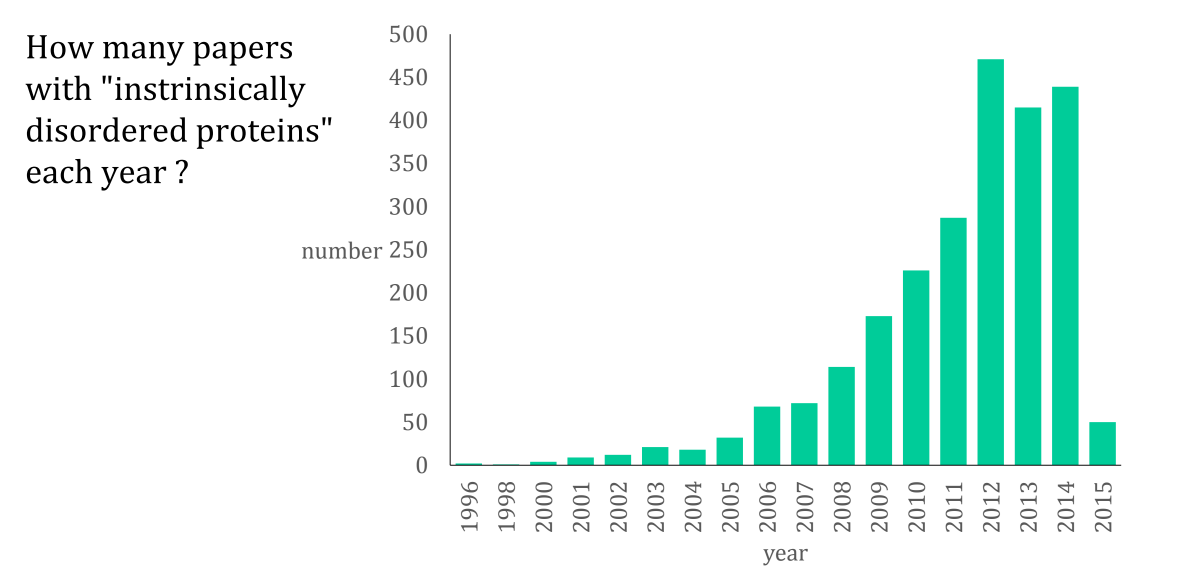
instrinsically disordered, natively disordered, natively unstructured. Can a protein have no structure and be useful ?

The real question

- are there proteins which do something (message, catalysis)
- have almost no fixed structure in solution
- maybe take on a structure when they meet a ligand or other protein ?

Fashion

#### A trendy discovery



28/01/2015 [2]

# Why discuss them?

Claim

- there are proteins whose native structures we will never see
- might adapt a conformation when they bind something
  - DNA, other protein, other ligand
- contradict classic belief function requires well-defined structure

Maybe they are important - claim

44 % of human protein-coding genes have a disordered segment more than 30 residues long

What do they do ?

- molecular recognition, molecular assembly, protein modification, and entropic chains
- sites for modifications (phosphorylations, cleavage, ..)

## What properties would you expect?

The features that you see from non-native, unhappy proteins

- gel filtration not sharp peaks
- proteases very susceptible
- folding / unfolding curves no specific unfolding temperature as with most proteins

The difference ?

• despite appearing unhappy, they still have some function

### **Example functions - enzymes**

Could you have an enzyme with little structure?

- normal picture
  - well-defined structure positions groups in space so they catalyse some reaction
- bad examples
  - RNase E has structured domain + large unstructured region
  - hepatitis C virus NS3 protease Zn<sup>2+</sup> necessary to fold and make active
  - is there evidence that the unstructured protein does anything interesting?

# Example functions - binding

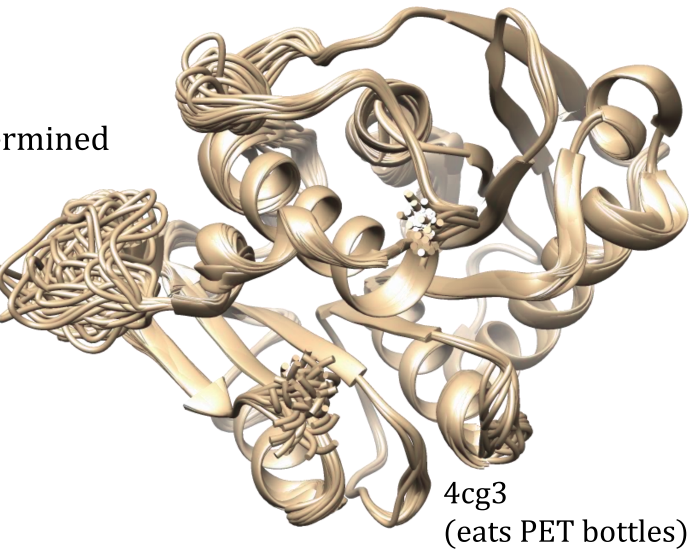
- DNA binding proteins take on structure only after binding
- targets for modification maybe being exposed makes them easier to phosphorylate, acetylate, ...
  - these changes lead to activation / inactivation
- small ligand storage.. Zn<sup>2+</sup> example coming

What is disorder ? When is it significant ?

## uncertainty

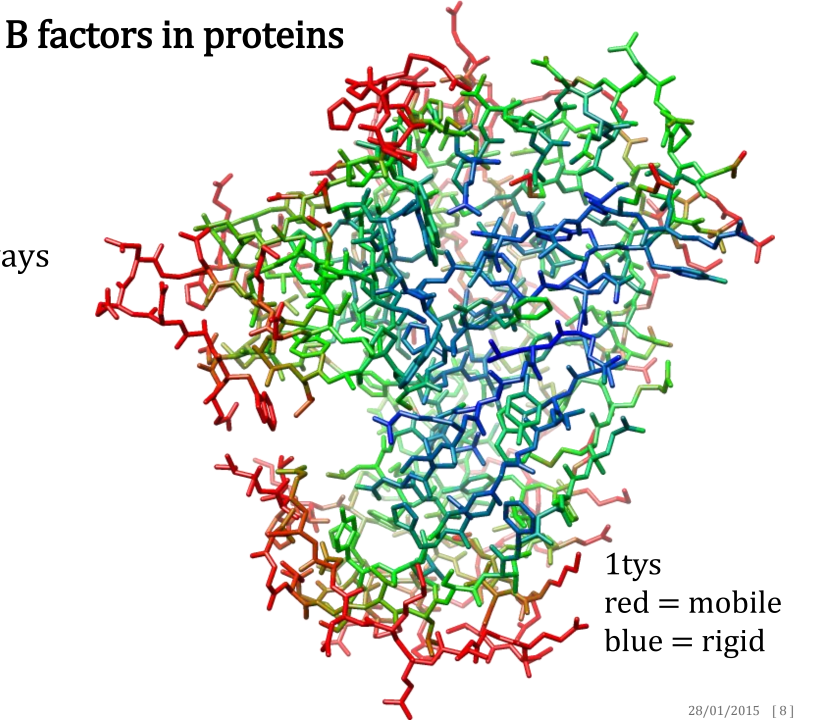
#### typical NMR structure

- 50 models
- some parts of structure well-determined
- some very mobile
- just a feature of NMR ?



#### **B**-factors

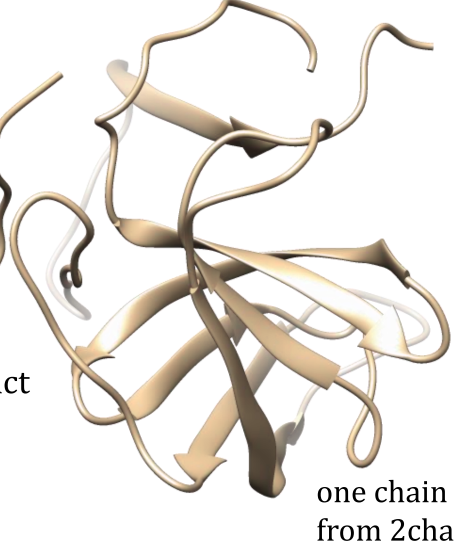
- better measure of mobility
- crystallographers have always known that some parts of structure are mobile
- what about really mobile ?



# Mobile pieces

History of structures like chymotrypsin

- easy to obtain, crystallise
- lots of structures
- always missing regions
- almost all chymotrypsin structures have this region
  - does not seem to be an experimental artefact



#### do intrinsically disordered proteins exist?

In perspective

- all atoms move at room temperature
- some parts of a structure are relatively unrestrained
- are there large parts which are unrestrained under normal conditions?

# Why be suspicious

Why do you not see structure ?

- It is easy to denature some proteins
- most proteins are marginally stable
- speak to a molecular biologist my protein does not fold
  - because the protein is unfolded when native (intrinsically unfolded) or
  - you have wrong conditions ?

What are conditions ?

- cells are very crowded
- sometimes exotic
  - organelles (mitochondria low pH)
  - electric fields near membranes (think of nerve cells)

#### What if you do not see structure

NMR

- NOE's distances between residues
- not easy to measure
- signal to noise
- technical reasons for not seeing NOE ( $\tau_c$  correlation time bad luck)
- your sample is bad

Crystallography

• not easy – most proteins need slightly unusual conditions

What if I do not see structure ?

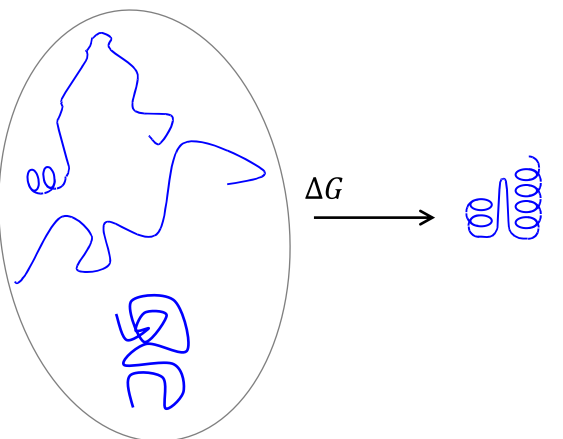
• maybe measurements are under wrong conditions

# What does unstructured mean?

Most chemistry is an equilibrium

- folding  $\Delta G = RT \ln \frac{\text{[folded]}}{\text{[unfolded]}}$
- $\Delta G$  is usually small

• if 
$$K_{fold} = \frac{[folded]}{[unfolded]} = 10^3$$
  
the unfolded is like a 1 in 1000 inpurity



- if  $K_{fold} = 1$ 
  - then your protein is 50% present, might be enough to work
  - may not look structured in NMR, may not crystallise
- does this explain "transient structure" ?

## What would you see experimentally

- crystallography nothing boring
- NMR? What you see from denatured proteins
  - broad lines
  - no NOEs (distances)
  - evidence of mobility ?

#### Better

• measurements that look at overall shape

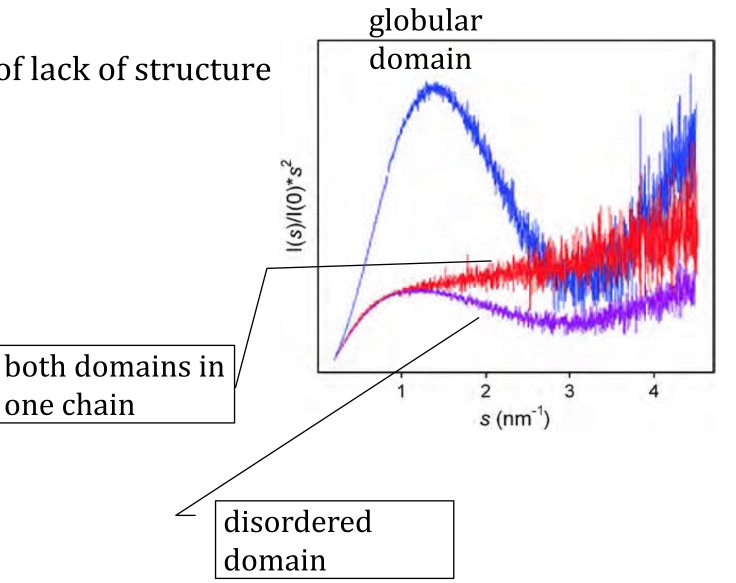
• radius of gyration 
$$r_g = \left(\frac{1}{N}\left(\sum_{i=1}^N |\vec{r}_i - \vec{r}_c|^2\right)\right)^{1/2}$$
 where  $\vec{r}_c$  is centre

• light scattering, small angle x-ray scattering (SAXS)

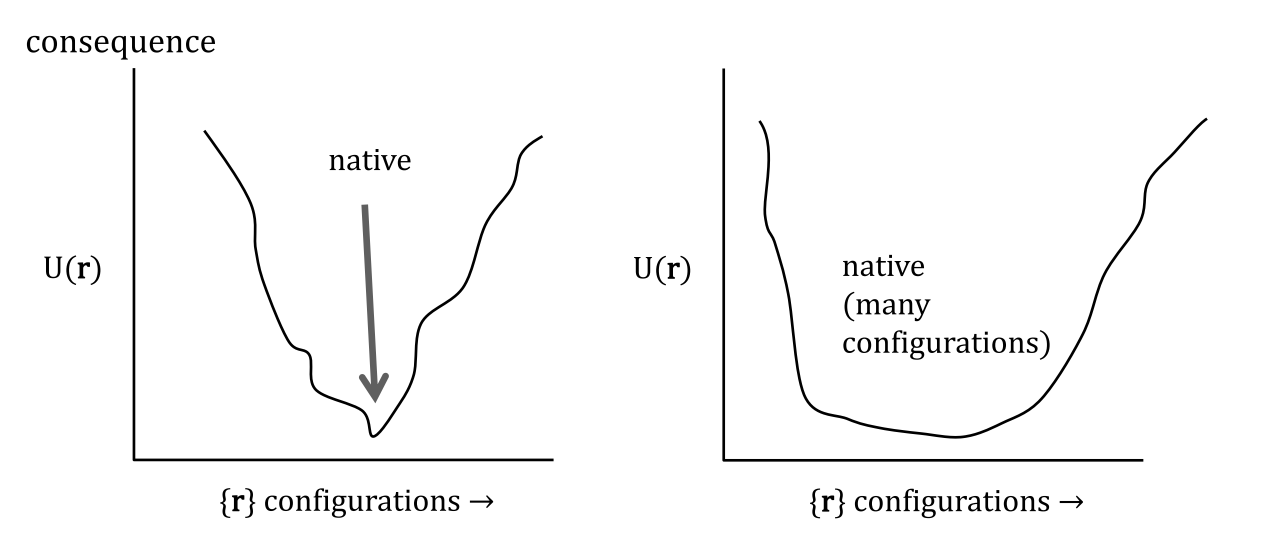
# Example...

src-kinase

- certainly can find evidence of lack of structure
- Krattky plots for
  - globular domain
  - unstructured domain
  - the two in one chain



### Energy landscape



what would one need for this kind of landscape?

# Flat energy landscape

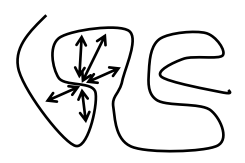
- Two possibilities
- 1. particles like to interact with each other, but they do not care with whom

2. particles interact with solvent .... some detail

#### Particles interact with each other

Non-specific

- a particle interacts equally with many other sites
- simplest extreme example
  - (leu)<sub>n</sub> or (trp)<sub>n</sub> each hydrophobic residue likes to pack with other hydrophobic residues
  - would not be soluble
- maybe realistic
  - "low complexity"
  - similar sets of residues repeat
  - DSKDSKDSK...DSKDSK...
  - lots of interactions of similar energy
  - statistically many proteins have regions of low complexity



#### particles interact with solvent

if side chains are very soluble, there is no hydrophobic core

- does it happen ?
- prothymosin  $\alpha$  (nuclear protein, interactions with chromatin, histones)
  - 109 residues, half are glu or asp, charge -54 at neutral pH
  - normal proteins have a small charge
  - binds lots of metal<sup>+</sup> ions in a cell
  - other histone binding proteins have acidic domains
  - what will happen at low pH?

#### What else do we know?

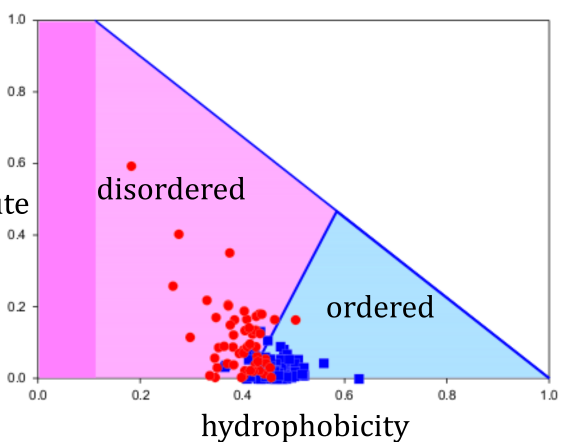
People collect lists – there are databases of disordered proteins

Sequence properties of intrinsically disordered proteins

- rich in polar and charged amino acids
- less than usual bulky hydrophobic

Best property...

o.6 average absolute charge <sup>0.4</sup>



Uversky, Biochim Biophys Acta, 2013, 1834, 932-951

# Can one predict disorder

For building and testing methods, one needs test set

• lists of largely disordered proteins *vs* normal proteins with known structures people collect these

#### OR

- take set of PDB crystal structures with B-factors
  - high B-factors = less rigid
  - try to predict B-factors

What descriptors would you use ?

# Typical prediction method

Features

- 1. sequence complexity
- 2. hydrophobicity
- 3. charge
- 4. single residue propensity

a few words on each ...

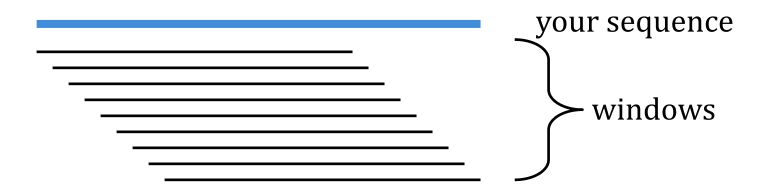
# low complexity

example

something like

 $c = -\sum_i p_i \ln p_i$  visiting each site over window of  $\approx 40$  residues  $p_i$  is the frequency of each amino acid type *i* in the window

• summing over all overlapping windows gives per residue score



# hydrophobicity and charge

- literature values for hydrophobicity (easy)
- charge very easy

## single residue propensity

- forget physics and chemistry just look at statistics
- visit all sites in protein data bank
- for each type of residue
  - collect average B factors
  - call this disorder propensity
  - residues missing in crystal structures are called "very disordered"
- end result ? table with big variation / error bars
- likelihood of an amino acid type to be disordered

# predictions

Methods

- neural networks, support vector machines (SVMs)
- classifiers (trees, bagging + boosting)
- elegant statistics ?

What comes out?

• first, how do you measure it

### **Predictors – ROC curves**

How good is your predictor?

• that a medication works, that a signal is real, that amino acid is disordered?

A good predictor

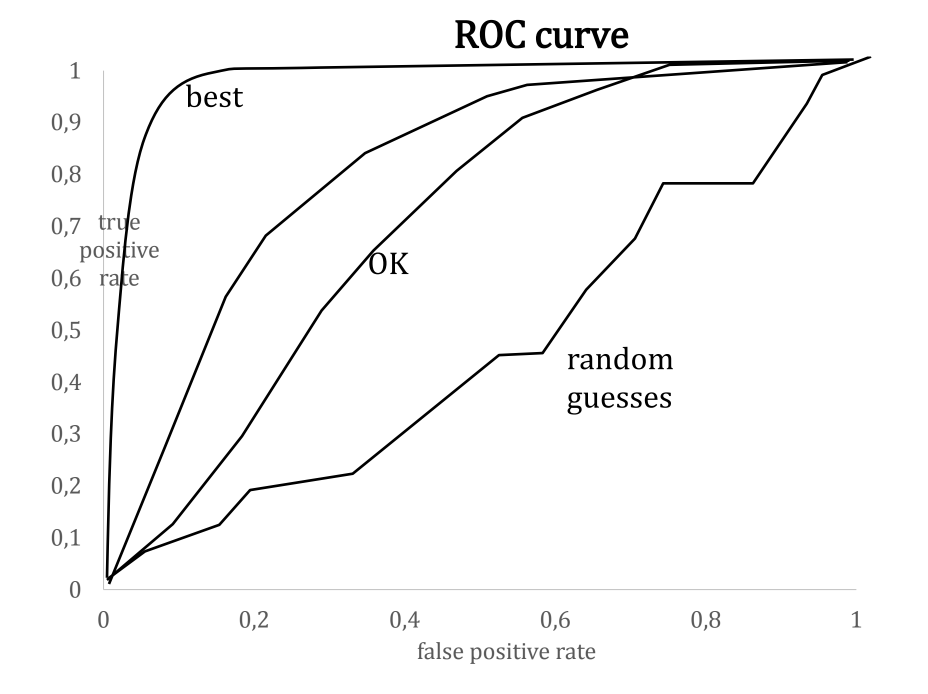
• the first 8 predictions are correct, then one wrong, then...

A bad predictor

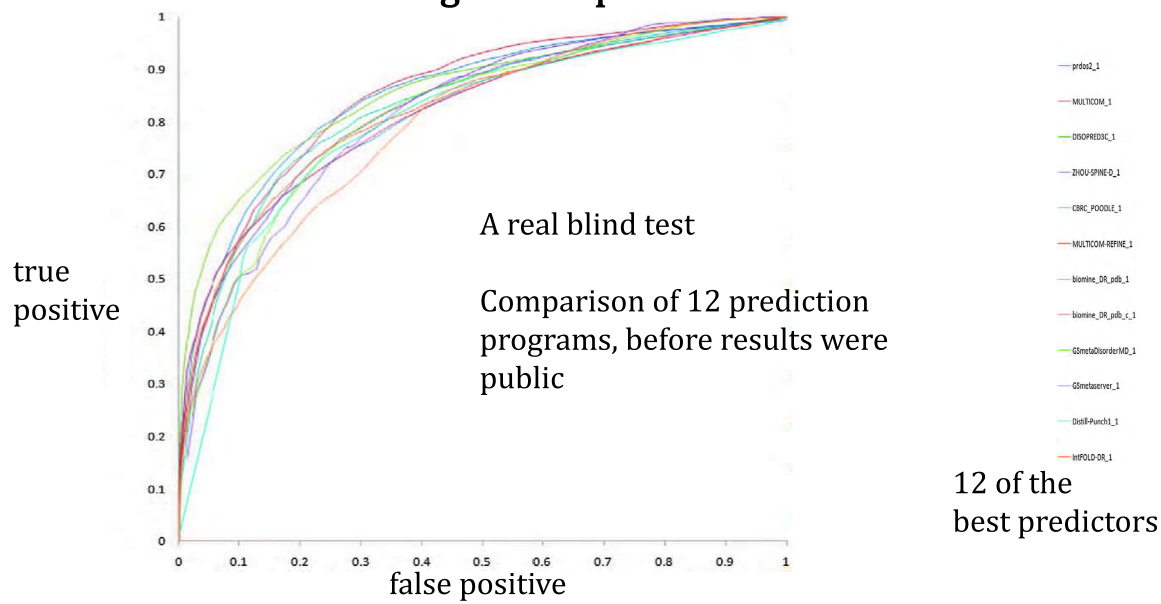
• the first 1 is correct, next is wrong, then 1 correct, then one wrong..

Turn this into a curve

- When 1 % of my predictions are wrong, how many are correct?
  - when 2 % of my predictions ...



#### How good are predictions?



#### interpretation

No surprises

• we already knew that charge and hydrophobicity were good predictors

Is any of this real?

• It is very fashionable (100s of literature articles)

Imagine a protein has

- little structure in test tube, finds structure when binding DNA, other protein
  - is this really different to a protein which is more structured in test tube + changes structure a bit upon binding ?
- What if a protein folds at low pH?

# For Klausur

- what is an intrinsically unstructured protein ?
- why is there no clear definition ?
- what experimental methods would recognise the property ?
- what are implications for the energy landscape ?
- how does a ROC curve work?
- how would one estimate sequence complexity?