Comparative / Homology Modelling

Topics
• rotamer optimisation
• loop prediction
• reliability of sequence similarity

Summary
• one protein sequence (protein 1)
• some related protein with structure (protein 2)
• put sequence 1 onto structure 2
The mission

..AADEFGHIKHFEDA.. your sequence

No structure
- will not crystallise, too big for NMR, in a hurry, no money

You want to
- replace a residue for binding to a surface
- guess which residues in your sequence are involved in chemistry
- ...

Andrew Torda
16.12.2019
...AADEFGHIKH-GED...
your sequence
do a blast search ... find
...AQDEF-HIKKGFED
structure 4b49 in PDB

replace original
..AQDEF-H..

with your sequence
..AADEFGH..

replace sidechains
Using model

with substrate

...AADEFGH\textcolor{red}{I}KH-GED...

who is near substrate?
predictions as to active site
You now have coordinates for your sequence
• how accurate?
• does it matter?

May not need to be accurate
• phasing (X-ray crystallography)
• guiding mutagenesis

May or may not be good enough
• docking
Most basic rule

Guiding belief
• similar sequence gives similar structure
  • evolution
  • chemistry

Most important
• closer the sequence is to template (sequence terms) – better the model
Reasonable expectations

- two enzymes (G6Pdh) easy to find homology
- could one have been modelled, knowing the other?
- knowing the structures below, this might be the limit of what could be done
Sequence and structure similarity

Two proteins with similar sequence
- how likely is similar structure? very
  - question of degree (how similar?)

Reasons?
- Intuitive – chemically obvious
- evolution

More on this next semester
Overall modelling protocol

1. decide on template
2. align sequence (unknown structure) to known structure / template / parent
3. replace sidechains of parent with new ones
4. fix
   • gaps
   • insertions
   • loops
5. overall structure
Finding a template / parent

How unique is my sequence?

- given human haemoglobin, you would find horse, pig, and $10^3$ globin structures
- given a strange enzyme from an exotic virus, it may have no obvious homologues – it has evolved too much
- blast / psi-blast / fasta / HMMs

<table>
<thead>
<tr>
<th>high sequence identity</th>
<th>low sequence identity</th>
<th>very low</th>
</tr>
</thead>
<tbody>
<tr>
<td>(&gt; ~20-25 %)</td>
<td>(&lt; ~20-25 %)</td>
<td></td>
</tr>
<tr>
<td>blast, fasta, anything</td>
<td>psi-blast, HMMs</td>
<td>psi-blast, optimism</td>
</tr>
</tbody>
</table>

Why so vague?
Template reliability

Length and degree of similarity

Old rule
  < 20 %, not similar
  > 25 % similar
  otherwise (twilight zone)

..not a good rule
Template reliability

Why is this not enough?

• consider random mixture of amino acids
• add bias of composition (some amino acids are rare)
• compare a lot of proteins and say
  • pairs have 15 % similarity (average)
• we see a pair of 20 % similarity for 50 residues
  • is it significant?
• we see a pair of 20 % similarity for 600 residues
  • more convincing
Quantifying importance of similarity length

We know the size of an alignment how often are the two proteins not structurally related?

but there is more to deciding whether or not similarity is significant
More to reliability

15 % similarity

your sequence  PDB sequence  not so reliable

40 %

your sequence  PDB sequence  much more reliable

3rd sequence

40 %

How significant is the similarity between two proteins?
• does not only depend on the two proteins
• psi-blast in sequence lectures
Summarise

Sequence identity is most important

It is not enough to say 20 – 25 % similarity
Sequence alignment

We have picked a template for our sequence now...

1. decide on template
2. align sequence (unknown structure) to known structure / template / parent
3. replace sidechains of parent with new ones
4. fix
   - gaps
   - insertions
   - loops
5. overall structure

Need an alignment

- difference compared to database searches?
  - not scanning a database (10^7 sequences)
    - we can do best possible alignment
Careful alignments

Computer time not a problem - use
• most expensive alignment algorithm, could be one of
  • Needleman and Wunsch
  • Gotoh
  • Smith and Waterman

Maybe
• fancy substitution matrix
• play with gap penalties

How important?
Alignment errors

two reasonable alignments

- ANDREW or ANDREW
- ANQEWE or AN-QEWE

difference?
- from $C_i^\alpha$ to $C_{i+1}^\alpha$ almost 4 Å
Difficult alignment example

- sequence with unknown structure **ANDREW**
- sequence of structure **ANDRWQANDRKWSANDRWWC**
- reasonable alignments
  
  **ANDREW**------------------  guess 1  [ includes gap
  -------**ANDREW**--------  guess 2
  ------------------------**ANDREW**  guess 3

- Is one correct? More likely to be correct?
- guess 1 - a residue has disappeared (difficult to model)
- guess 2 - K -> E;  guess 3  R -> E
- very dependent on
  alignment quality / scoring / substitution matrix
Sidechains – should we worry?

When do we not care?
- for some residues, not meaningful (ala/gly)
- some residues entirely on surface of protein
  - interact with solvent
  - barriers to rotation? smaller than $kT$
  - all conformations accessible

When is it sensible to worry?
- sidechain is big and buried
- sidechain is charged and buried (salt bridge?)
- example – trp usually
  - big
  - buried
  - hydrophobic
  - not very mobile
Sidechain placement

How to place sidechains

• if identical to parent
  • re-use parent coordinates

Question

• what angle should I have at each rotatable bond?

Reasonable strategies

• initial placement
  • random
  • probabilities from protein data bank?
• fix !..
Fixing sidechains

Considerations
- atoms do not lie on top of each other
- residues like to pack (few holes in proteins – energy arguments)
- hydrophobic residues like each other
- charged and polar residues usually talk to solvent
- buried charges in salt bridges / no free charges in protein core

Can we write this down as a formula?
- almost
  - an energy function should contain this (next Semester)
Optimising sidechains

Basic philosophy
- write down some function for energy +
  - energy minimisation
  - molecular dynamics
  - Monte Carlo / simulated annealing
  - self-consistent mean field methods
  - clique method – our example
- so as to rotate side-chains / make conformations more likely
Rotamers and cliques

Many ways to optimise side chains
• annealing, simulations, self-consistent mean field optimization

Clique detection
• just one example (not best, fastest, ...)

Ingredients
• side-chain rotamers (discretisation)
• score for energies / clashes

Definition
• clique – subgraph where each point is connected to all others
Most sidechains have rotatable angles (more than 1)
- for each angle – usually 2 or 3 angles are more likely
- approximate:
  - pretend each side chain may only exist in one of the preferred positions "rotamers"
  - per sidechain
    - maybe 3, 9, .. rotamers
- crude ? yes
- useful ?
  - transform problem into a smaller search

Rotamers

Fitting rotamers in a protein

Simple quasi-energy function

- atoms may not clash
- imagine 0 is fixed
- 0 does not fit with 1
  - OK with 2 or 3
- 1 is not OK with 0, 2, 3
  - OK with 4, 5, ... 9

What we want – lists of who is compatible with who
Rotamers

Draw as a graph
• lines connect who is compatible with who
• connections for 0 and 1 drawn
• do for all other nodes (rotamers)
• no edges between nodes for 1 residue
Rotamers

Imagine there is only one possible set of rotamers
- every node (rotamer) will be connected to every other
  - is a clique
Imagine there are two solutions
- there will be two cliques

Application
- take protein
- build graph
- find all cliques
- write out lists of sidechain conformations

What was a very difficult problem seems to be tractable but...
Rotamers – problems with cliques

Killer problem
• finding maximal cliques is very very difficult

Rotamer concept
• side chains do not exist at only 0, 120, 240°

Better energy functions are more complicated
• here: only said - not compatible/incompatible
Broken main chain

Typical situation
ANDR–WQANDRKWSANDRWWC parent
ANDREW—DRKWS—DRWWC model
our model...

Basic problem...
• pieces of unknown structure
• endpoints relatively fixed
• should be joined
Loop modelling

Loop problem

• do not want to disturb regular secondary structure
  • more likely to be correct
• ends of loop relatively well known
• composition (sequence) of loop

The problem specifically:

• find an arrangement of backbone and sidechains which
  • is geometrically possible
  • low energy

Possibilities

1. distance geometry
2. database search
3. brute force

Andrew Torda
16.12.2019
Distance geometry
- we know
  - end points and distances
  - sequence of loop
    - all bond lengths and angles
- use distance geometry to generate plausible arrangements (Modeller Übung)

Results?
- arrangement of atoms with
  - correct covalent geometry
  - no atoms on top of each other (set by minimum distances)
- little consideration of torsion angles
2. Loops Database searching

Database searching
- imagine we have a 9 residue loop
- take protein data bank
- collect coordinates of all 9-residue loops
- insert those with correct end to end distance
- refinement...
  - insert those with almost correct distance &
  - similar sequence to loop residues
3. Loops – brute force

Desperation / brute force for small number of residues

- divide angles into pieces (maybe 30°), $360/30 = 12$
- test every combination (joining ends, energy)
- called "grid search"

- How many angles?
- per residue
  - fix $\omega$
  - phi $\phi$, psi $\psi$ $12 \times 12 = 144$
- possibilities = $144^{N_{res}}$
General repairs

What do we have now?
• sidechains placed and maybe optimised
• rough guess coordinates for all residues (including loops)

Broken?
• sidechains and loops often wrong
• small changes in other parts of structure
• time for last refinement .. again
  • energy minimisation / molecular dynamics / ...
Quality

General vs specific
- general
  - energies / geometries (almost the same)
- specific properties of this protein (vague and not for exams)
  - expected residues in active site
  - known reactive residues on surface
  - ... any experimental data
Checking by energy

Use a classical energy function (details next semester)
• if physics were perfect, would include all ideas mentioned
• details good (atom overlap, angles, ..)

Statistical approach
• take features you believe in
  • hydrophobic residue on surface, buried residue in middle..
  • phi / psi distributions
  • count occurrence in databank
• count occurrence in your model
• see if model is statistically plausible
Real world

Recipe on these slides?

- too simple
  - steps combined / repeated
  - usually many models generated and checked multiple templates
  - multiple templates simultaneously?

- automatic methods are very good
What does one achieve?

Very easy cases?
- not much change from parent

Very difficult?
- lots of errors

Why bother?
- good modellers are experts on their systems
- some proteins are so important (money) – no waiting on
  - experiment
  - competitors
- simple predictions
  - which residues may I modify (binding to sensor…)
- consider absolute limits
An Example

2mnr and 4enl

• would be a typical modelling target
• in real world
  • alignment would not be perfect
  • loops may be quite wrong
The sequence alignment

2mnr and 4enl example

• sequence alignment not the same as alignment from structures
Why I do not like modelling

You build a model – decide that the grey residues are important

- Has the model helped you?
known structure

your sequence

but we could look in original structure and mark all the residues near active site

and look for corresponding residues in your sequence
Summarise für Klausur

Ideen der Sequenzähnlichkeit

Technische Probleme
- loops
- Seitenkettenplatzierung

Keine der unscharfen Aussagen
- Qualität