Molecular Evolution

Andrew Torda summer semester 2015, Struktur & Simulation

Why?
• applications not possible with detailed models

Ingredients in this set of lectures
• models for proteins
  • simple representation - lattices, simple energy functions
• Boltzmann relation and partition function
  • ability to calculate probability of conformations

Aim
• from very few assumptions
• simulation which reproduces physical properties
Why lattice models?

Earlier – building models
• how much detail - rather arbitrary

Here – minimal models
• one does not need serious chemistry to reproduce protein properties
• evolutionary pressure may not be real
Plan

Generalities
- sources of evolutionary pressure
- example of unexpected evolutionary pressures (Darwinian)
- neutral networks
  - alternative explanation
Evolution observables

Phenotypes / population properties

- blue eyes, brown eyes (macroscopic)
- protein /nucleotide functions (molecular)

Consequence?
- mostly look at evolution in terms of pressure on phenotypes
- classic adaptive Darwinism

First - a property to be explained later
Haemoglobin conservation

Look at residues 37, 43, 83 and 87

4 residues stand out as conserved
Sequence variability

Take family of related sequences
• see how conserved / variable they are

Variable sites
• are they unimportant?

• remember this picture!

• return to Darwinism
Adaptive Darwinism

- I see a fish which lives behind a rock and eats seaweed
- A mouse is just the right size to squeeze through the hole in my wall
- Voltaire (1694-1778)

Master Pangloss taught...

dass in dieser besten aller möglichen Welten, ...

„Es ist erwiesen“ sagte er, „dass die Dinge nicht anders sein können: denn da Alles zu einem Zweck geschaffen worden, ist Alles notwendigerweise zum denkbar besten Zweck in der Welt. Bemerken Sie wohl, dass die Nasen geschaffen wurden, um den Brillen als Unterlage zu dienen, und so tragen wir denn auch Brillen“

Two aspects
- adaptation to glasses (evolution is directed)
- best of all possible worlds (we / the world are optimised)
Classic Darwinism – molecular level

Obvious pressures
• function – protein must work
• stability – must be stable under your conditions

Less obvious, but simple
• folding – must fold in reasonable time

Less obvious, but reasonable
• mutation resistance
Other evolutionary pressures

Is it good to be resistant to mutation?
• what if a gamma ray hits me and my children die?
• more formally
  • a sequence (protein) is more likely to propagate if
    • it can be changed
    • it keeps functioning
• can this be modelled?

Plan:
• be Darwinian
• (later) show why it is probabilistic (not Darwinian)
Simulating mutation resistance

Lattice simulations
• 25 residues, 2 dimensional, compact, 5×5 lattice
• 20 residue types (not two or 5 or 6)
• 1081 conformations
• remember we can calculate $Z$ and stability
• for any sequence can say
  • will this sequence fold or not? $\Delta G_{fold}$
    • how different is lowest energy to other energies
• too big to check all sequences

Example calculation
• look at differences with and without evolution

Example evolution calculation

Evolution simulation
• apply mutations infrequently / randomly
• sequence must maintain
  • same structure
  • foldability
• for each member of population
  • check lowest energy configuration
    • if it has changed – sequence dies
  • check $\Delta G_{fold}$ based on Boltzmann probability of lowest energy structure
    • if sequence is not foldable – dies
• of remaining sequences, randomly pick for reproduction
Simulation reminder

Simulations this semester

• system is not at equilibrium at start

Normal procedure

• simulation for \( n \times 1000 \)'s steps ... throw away
• simulation more ... keep for averaging and analysis
Comparing populations

Take a sequence which folds
• copy 3 000 times – initial population

<table>
<thead>
<tr>
<th>initial population</th>
<th>forget (equilibration)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>30 000 generations</td>
</tr>
<tr>
<td>diverse population</td>
<td>keep and sample</td>
</tr>
<tr>
<td></td>
<td>30 000 generations</td>
</tr>
</tbody>
</table>

generated sequences randomly

evolved sequences

random sequences
Properties to look at

• How often does a mutation make a protein more stable?
• How often does
  • a stable protein become more stable? (not often)
  • an unstable protein become more stable? (must be higher)
• Do the fractions differ between
  • random sequences (right hand side previous Folien)
  • evolved sequences (left hand side)

From simulation look at proteins with some ΔG (stability)
• after mutation get new ΔG
• look at large number of mutations, get probability
  \( P(\Delta \Delta G > 0) \) of becoming even less stable
What do you expect?

Evolved sequences must be more stable than random ones (obvious)

Will they also be more resistant to mutations?
Simulation results

Take a sequence and have a look

- when it mutated and survived
  - how often did it become less stable $P(\Delta \Delta G > 0)$?

\[ \Delta G_{wt} = \text{stability before mutation} \]

\[ P(\Delta \Delta G > 0) \]

very stable proteins ↔ less stable

Simulation results

Becomes more stable

probability of becoming less stable

\[ P(\Delta\Delta G > 0) \]

very stable proteins \( \leftrightarrow \) less stable

Interpreting results

random sequence
• unstable ($\Delta G > 0$)
  • not easy to make more stable
• stable ? ($\Delta G < 0$)
  • all mutations make it worse

evolved sequence
• very stable ?
  • cannot make better
• marginally stable ?
  • mutations often OK
Results explanation

Without explicitly adding idea
• evolution makes
  • more stable proteins (obvious)
  • proteins which survive mutations (why ?)

Agree with experiment ?
• small amount of the time
  • mutations have no effect
  • make protein more stable than natural version
Results explanation

Stability was selected
- moves population to left
- it should not change the fraction of stabilising mutations

Simulation selected for stable sequences
- of those stable sequences, did not select for mutation resistance
- \( P(\Delta \Delta G > 0) \) is a probability

- effect must come from somewhere else
Sequence variability interpretation

- Typical part of sequence analysis
  - look at collection of related sequences and see how conserved they are (conservation, profiles, sequence entropy, ..)

- Why are some sites so well conserved?
  - function?
- Why do some sites vary?
  - old view: they do not matter
  - this paper
    - this is a consequence of evolution
  - if they are important and fragile, you die
Subtle evolutionary pressure?

Is this an evolutionary pressure?
- seems like a good idea to not die when mutated
- authors argue that the reason is different
- neutral evolution ...

so far
- very simple lattice model reproduces
- stability, evolutionary pressures

- not Darwin, but what is it?
Simulating at the molecular level

Basic idea
• take a population (maybe $10^3$ or as big as possible)
  • make random changes
  • look at consequences
  • kill or reproduce molecules

Most popular
• RNA
  • for a given mutation, can guess at secondary structure
• Proteins
  • lots of lattice calculations
Simulation machinery

HP model in two dimensions
• length 18
  • one can look at all sequences
  • all conformations
  • ... for any sequence
    • can find minimum energy structure
  • for any structure
    • we can find all sequences which have this as minimum energy

Bornberg-Bauer, E (1997) Biophys J. 73, 2393-2403 How are model protein structures distributed in sequence space?
Calculations

Find popular structures
• which is best for many sequences
• collect these sequences
  • neutral set
Neutral mutations
• which of these sequences are connected by a point mutation?
• example
  • HPHPHHH . . and HPHPPHH . . have same ground state
  • they are connected by one change
  • this change does not cost anything in evolution
    • it is "neutral"
  • in pictures...
Neutral mutations

Look at sites which can be changed
• many possible sequences
Can one mutate each to every other?
• $\text{HPHPHHH}$ . . . and $\text{HPHPPPP}$ are not connected
What can we say about the connected sequences?
• form connected sets

sites where neutral mutations were found

$\text{HPHPHHH}$ and $\text{HPHPPPP}$ may be a set, but not connected

Bornberg-Bauer, E (1997) Biophys J. 73, 2393-2403 How are model protein structures distributed in sequence space?
Connected and non-connected sets

Each dot is one protein sequence/structure

neutral set with two connected sets

neutral set and connected set
Neutral networks

Sequences which can turn into each other are "neutral network"

How big are the neutral sets ?
• about $\frac{1}{4}$ have more than 5 sequences
• most popular has 48 sequences
• lots of very rare structures

Are these sets fully connected ?
(can anyone eventually mutate into anyone else) ?
• about 80 % of time
**Evolutionary consequences**

- a population can quickly spread over a huge number of accessible sequences
- immense variation at molecular level is possible

- Can one hop between different connected networks?
  - in this model – not so easily (≥ 2 mutations)

More interesting consequences
- some structures are hard to find by random moves
- some are very popular
- what does this say about mutation study?
Mutation resistance revisited

Earlier slides
• it seems as if proteins evolve in order to be resistant to mutations (sounds Darwinian)

Alternative
• think of sequence space
• a group of related sequences are a cluster in this space
Networks, probabilities, mutation resistance

huge network
1000's sequences

mutate to here
• seems mutation resistant
  • lots of possibilities to mutate and maintain structure
  • more likely to be found (more sequences)
• mutate here? likely to die

This is the alternative explanation of mutation resistance
• nothing to do with evolutionary pressure
Darwinian versus neutral evolution

Crux of these lectures

• Darwinian evolution – what you see is
  • most fit (selection pressure)
• Neutral evolution – what you see is
  • whatever is most likely to occur

Relevance to mutation resistance

• Darwinian
  • useful trait that will be selected for
• Neutral
  • larger neutral networks
    • by definition – mutation tolerant
    • because they are larger, more likely to be found
Summarise

• simple system lets you simulate long-term behaviour
• simulation selected for folding - found mutation resistance
• explanation comes from neutral networks

• not really an evolutionary trait