Protein Design

Andrew Torda, wintersemester 2007 / 2008, AST ... 00.904

- What is it?
- Why?
- Experimental methods
- What we need
- Computational Methods
- Extensions

fragt nicht:

Fahrradunfall

What is protein design?

- Assumption
 - you can write a protein sequence on a piece of paper
 - a molecular biologist can produce it
 - clone, express, fold, purify, ...
- Most general
 - you have a protein which is useful (enzyme, binding, ...)
 - you want to make it more stable
 - temperature
 - solvents (tolerate organic solvents)
 - pH

Experimental approach

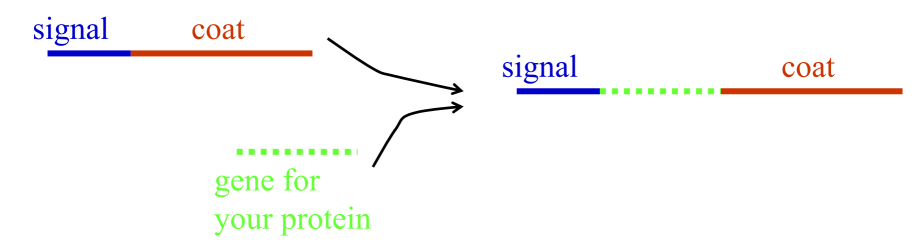
- 1. simple selection
- 2. phage display
- 3. *in vitro* evolution
- 4. manual

Selection

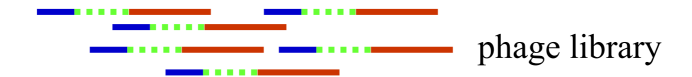
- Want protein that is active and more stable
- need assay for activity
- clone gene into bacteria, (semi-)randomly mutate
- select for bacteria (need assay)

phage display

- aim evolve / select for proteins with better binding
- put gene into phage

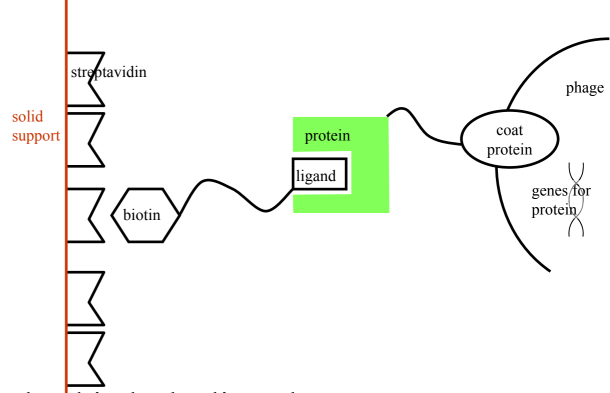


copy many times and mutate gene for your protein



phage display

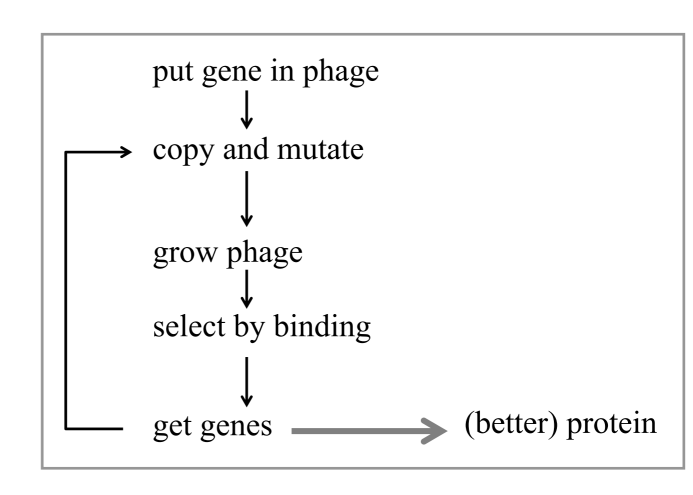
- grow up phage with the library
- selection
 - needs some strong binding like streptavidin+biotin



- if we have a protein that binds the ligand
 - can be selected + its corresponding genes

phage display

• improve binding with each cycle

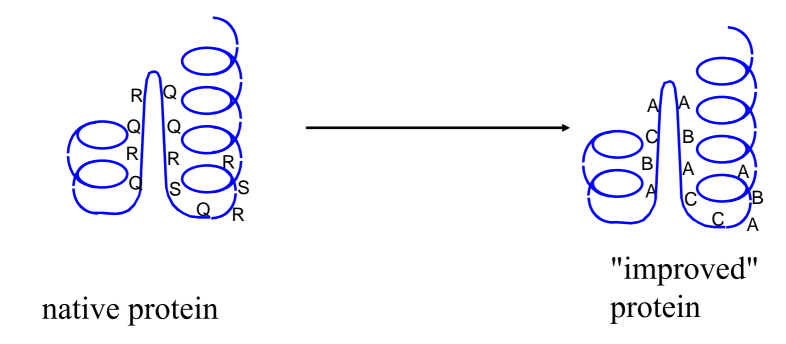


Other experimental methods

- in vitro evolution / ribosome display
 - similar philosophy to phage display
- manual
 - guess and use site directed mutagenesis
 - compare with phage display
 - few mutants instead of 10⁴
- computational methods ...
 - first specify the problem

Formalising the problem

- We have a working structure
 - want to make it more stable (limit to this)

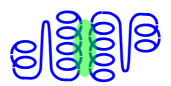


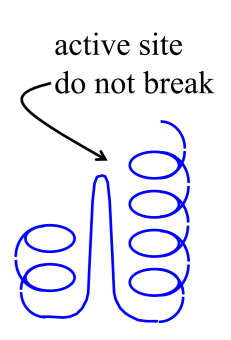
- Rules
 - structure should not change
 - should be able to fix some residues (active site, important)...

Fixing / specifying residues

Examples

- lysine (K) often used for binding
 - change a residue to K and protein does not fold
 - mission:
 - adapt the rest of the residues to be stable
- change all residues, but not those in active site
- change some residues at surface to be soluble
- change some residues at surface to stop dimers





Ingredients

- Score function (like energy)
- Search method

Score function

- how does sequence fit to structure?
- sequence $S=\{s_1, s_2, ...s_N\}$
- coordinates $R = \{ \mathbf{r}_1, \mathbf{r}_2, \dots \mathbf{r}_N \}$
- score = f(S, R) (diffferent nomenclature soon)
- mission
 - adjust S to as to maximise score (minimise quasi-energy)

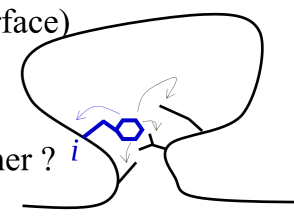
Score function

- how do amino acids
 - suit structure?
 - suit each other?

$$score = \sum_{i=1}^{N_{res}} score_{struct}(s_i, R)$$

$$+ \sum_{i=1}^{N_{res}} \sum_{i>i}^{N_{res}} score_{pair}(s_i, s_j, R)$$

- *score*_{struct} might have
 - backbone preferences (no proline in helices, ..)
 - solvation (penalise hydrophobic at surface)
- score_{pair}
 - are residues too big (clashing)
 - are there holes? charges near each other? i
- messy functions
 - lots of parameters
- discussed more later



Searching

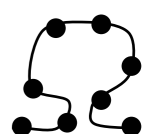
- long topic
- systematic search how long?
- search space for $N_{res} = 20 \times 20 \times ... = 20^{Nres}$
- must it be so bad?

What if there are no correlations?

for (i = 0; i
$$< N_{res}$$
; i++)
find best residue at position i

- search space would be $20 N_{res}$
- is this realistic?
 - not very every time I change a residue, it affects all neighbours
 - changing the neighbours affects their neighbours ...





Searching

- in a dream world could grow linearly with sequence
- in the real world = 20^{Nres}
 - brute force / systematic search not possible
- two methods here
 - 1. Monte Carlo / simulated annealing
 - 2. Pruning / dead end elimination

Monte Carlo

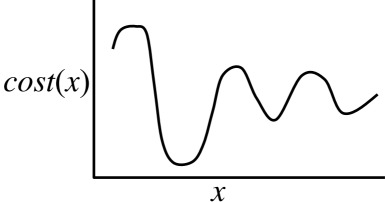
- more formally next semester
- first the problem

The sequence optimisation problem

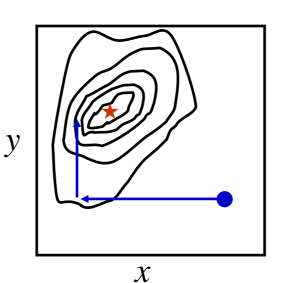
- discrete
- local minima / correlations in surface
- high dimensional

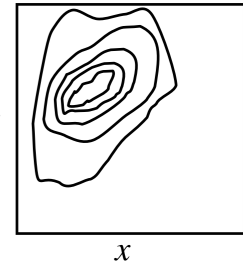
dimensions and correlations

• a 1D problem



- a 2D problem, but easy
 - only one minimum
- difficult correlations
 - the best value for x depends on y



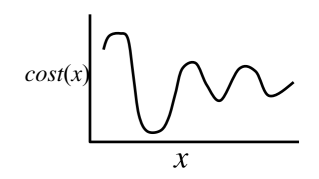


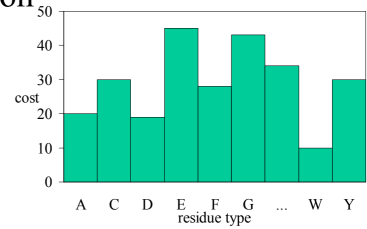
discrete

- for a continuous function use gradients
 - to optimise
 - to recognise minima / maxima
 - continuous functions
 - step in one direction is good
 - try another in same direction₅₀



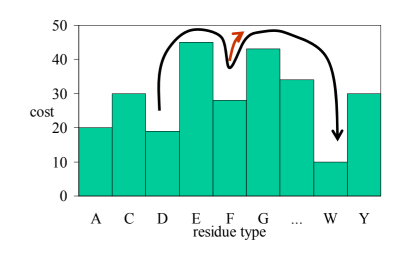
- no gradients
- order of labels arbitrary
 - ACDE or ECAD
- discrete
 - step in one direction may be no predictor of best direction





what do we want?

- from step to step (sequence to sequence)
 - be prepared to move in any direction
 - if the system improves, try not to throw away good properties
 - must be willing to go uphill sometimes
- philosophy
 - take a random move
 - if it improves system
 - keep it
 - if cost becomes worse
 - sometimes keep it
 - sometime reject



Acceptance /rejection

- for convenience, write $cost(S_n)$ neglect the coordinates R Sign convention
- system (sequence) at step n is S_n
- after a random step, cost changes from $cost(S_n)$ to $cost(S_{n+1})$
- $\Delta c = cost(S_{n+1}) cost(S_n)$
- our sign convention: if $\Delta c < 0$, system is better

When to accept?

- if Δc is a bit < 0, maybe OK
- if $\Delta c \ll 0$, do not accept

Formal acceptance rule

- $\Delta c < 0$, $e^{\Delta c}$ is between 0...1
- $\Delta c \approx 0$ then $e^{\Delta c} \approx 1$ as $\Delta c \rightarrow -\infty$ then $e^{\Delta c} \rightarrow 0$
- formalise this rule

```
set up S=S_0 and cost(S_0) while (not finished) S_{trial} = random \ step \ from \ S \Delta c = cost(S) - cost(S_{trial}) if (\Delta c < 0) /* accept */ S=S_{trial} else r = rand \ (0..1) if (e^{\Delta c} \ge r) S=S_{trial}
```

vorsicht! not the final method

why we need temperature

- As described
 - system will run around
 - try lots of new configurations
 - sometimes accept bad moves
 - always take good moves
 - may never find best solution
 - imagine you are at a favourable state
 - most changes are uphill (unfavourable)
 - many of the smaller ones will be accepted
 - if we were to find the best sequence, the system would move away from it
- how to fix?

why we need temperature

- Initial sequence is not so good
 - let the system change a lot and explore new possibilities
- after some searching, make the system less likely to go uphill
- introduce the concept of temperature T
- initially high T means you can go uphill (like a high energy state)
- as you cool the system down, it tends to find lowest energy state
- change acceptance criterion to $e^{\frac{\Delta c}{T}}$

• as
$$T \to \infty$$
, $e^{\frac{\Delta c}{T}} \to 1$ $T \to 0$, $e^{\frac{\Delta c}{T}} \to 0$

• put this into previous description

why we need temperature

```
set up S=S_0 and cost(S_0) set T=T_0
while (not finished)
S_{trial} = random step from S
                                            /* ε bit smaller than 1 */
     T = \epsilon T
     \Delta c = cost(S) - cost(S_{trial})
     if (\Delta c < 0)
             S= S<sub>trial</sub>
     else
             r = rand (0..1)
             if (\exp(\Delta c/T) \ge r)
                      S = S_{trial}
```

- name of this procedure
 - "simulated annealing"

Final Monte Carlo / annealing

- History applications
 - discrete problems travelling salesman, circuit layout
- deterministic? No
- convergence? Unknown
- practical issues
 - what is a random step?
 - change one amino acid? change interacting pairs?
- easy to program
- lots of trial and error
- statistical properties next semester
- can we reduce the search space?

Pruning

- Are there elements of sequence which are impossible?
 - at position 35, no chance of Y, W, I, L, ...
- can one find impossible combinations
 - reduce the search space so it can be searched systematically (brute force)
- ... dead end elimination method
- use an energy-like nomenclature

Nomenclature

- we are not dealing with
 - free energy G or F or potential energy U or E
- but let us pretend
 - score is E
- rule: more negative E, better the system
- structure is fixed so neglect R / r terms
- define a function $s_i(a)$ as the residue type at site i
 - can take on 20 values of "a" why?

 foreach (a in A, C, D, E..., W, Y)

 evaluate energy corresponding to a
- our energies?
 - two parts pairwise and residue with backbone

Nomenclature

- E is (quasi-energy) of whole system
 - label E_1 as the terms that depend on residue + fixed environment
 - E_2 as the energy terms that depend on pairs

$$E = \sum_{i=1}^{N_{res}} E_1(s_i) + \sum_{i=1}^{N_{res}} \sum_{j>i}^{N_{res}} E_2(s_i, s_j)$$

• if we are interested in site *i* and being in state *a* what do we have to look at ?

$$\sum_{i=1}^{N_{res}} E_1(s_i(a)) + \sum_{i=1}^{N_{res}} \sum_{j>i}^{N_{res}} E_2(s_i(a), s_j(b))$$

Nomenclature and rules

- there are $20 (N_{type})$ residues
- which fits best to the fixed environment? $\min_{a} E_1(s_i(a))$
- implies testing each of the N_{type} for a
- what is the best energy type a at site i could have, interacting with one site j? $E_1(s_i(a)) + \min_b E_2(s_i(a), s_j(b))$
- what is the best energy that type *a* at *i* could have considering all neighbours?

$$E_1(s_i(a)) + \sum_{j \neq i} \min_b E_2(s_i(a), s_j(b))$$

- for each a can work out what is the best score it could yield
 - loop over b
 - within loop over *j*

Dead-end elimination method

• worst energy that type c at i could have considering all neighbours? $E_1(s_i(c)) + \sum_{i \neq i} \max_d E_2(s_i(c), s_j(d))$

- when can one eliminate (rule out) residue type a at site i?
- for any residues a, c
- if the best energy for a is worse than the worst for c
 - a cannot be part of the optimal solution ... if

$$E_1(s_i(a)) + \sum_{i \neq i} \min_b E_2(s_i(a), s_j(b)) > E_1(s_i(c)) + \sum_{i \neq i} \max_d E_2(s_i(c), s_j(d))$$

Dead-end elimination method

$$E_1(s_i(a)) + \sum_{j \neq i} \min_b E_2(s_i(a), s_j(b)) > E_1(s_i(c)) + \sum_{j \neq i} \max_d E_2(s_i(c), s_j(d))$$

using this approach

how strong is this condition ?

DEE condition

- much of the time
 - cannot really rule out type a
- example?
 - initial 2×10^{27} final
 - searchable in 90 cpu hr
- deterministic

Dahiyat, B.I, Mayo, S.L. (1997), Science 278, 82-87

Combining ideas

- use DEE to get a list of candidate residues at each position
- search remaining space with Monte Carlo / simulated annealing
- not deterministic

Success

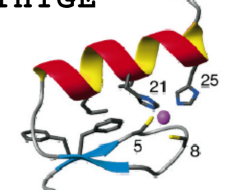
- Method
 - Dead end elimination + systematic search

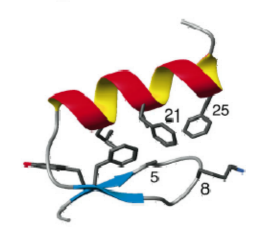
designed QQYTAKIKGRTFRNEKELRDFIEKFKGR

native KPFQCRICMRNFSRSDHLTTHIRTHTGE

New sequence

- about 20 % similar to start
- not related to any known protein (still)
- Structure solved by NMR
- Problem solved?
 - maybe not

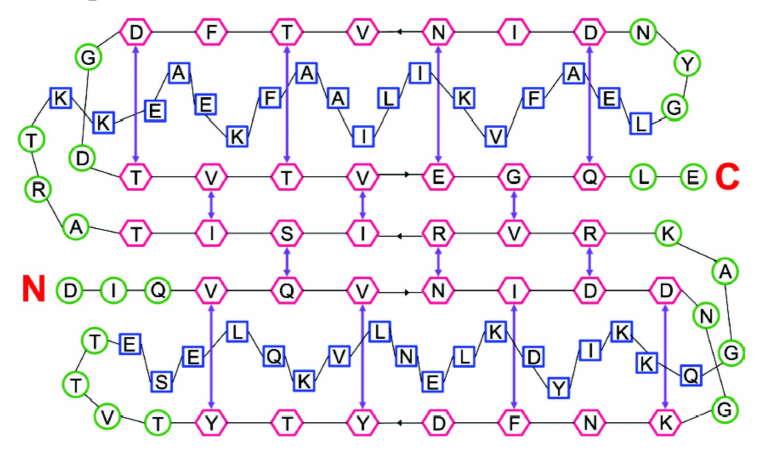




Success

Mission

- sketch a new protein topology
- build a sequence to fit it



Success

Methods

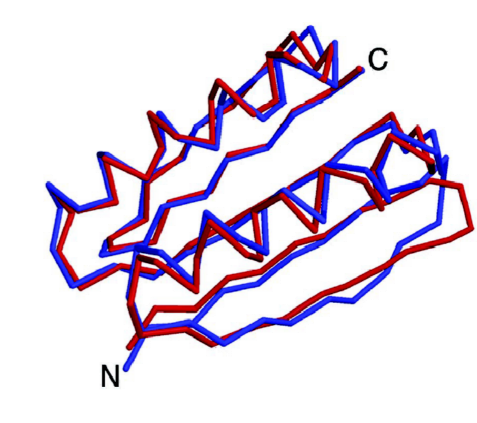
• pure Monte Carlo

Result

apparently new sequence

Structure

- as predicted
- solved by X-ray
 - neat phasing trick!
- Problem solved
 - unclear (how many failures ?)



Methods so far

Methods

	Monte Carlo	Dead-end elimination
guaranteed global optimum	no	does not try
deterministic	no	yes

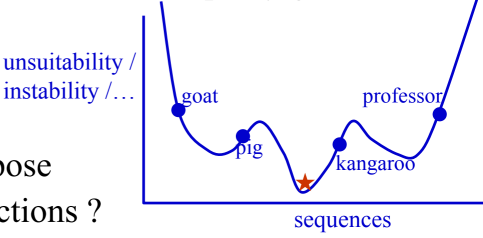
Determinism

May not matter

- consider real proteins compare human, goat, ...
 - all stable all slightly different
- implication
 - there may be many solutions which are equally good

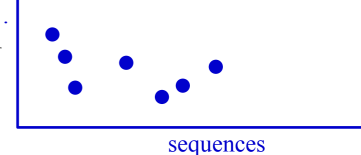
Counter argument

- sequences in nature are
 - not optimal
 - not optimal for our purpose
- How good are our energy functions?



Determinism and energy

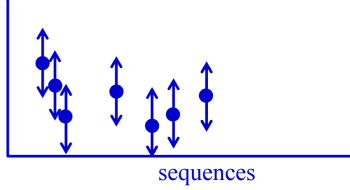
• I have a perfect score / energy function



unsuitability / instability / . . .

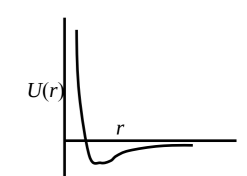
unsuitability /

- I have errors / approximations
 - best answer could be any one



Problems – stability / energy

- energy functions
- what do we mean by energy?
- example two charges $U(r) = \frac{q_1 q_2}{Dr}$
- example two argon atoms $U(r) = 4\varepsilon \left(\sigma^{12}r^{-12} \sigma^6r^{-6}\right)$
- make energy better?
 - replace every amino acid by a larger one (more contacts – more negative energy)
 - silly proteins are not full of large amino acids
- what determines stability?



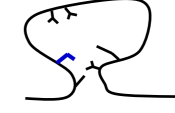
Problems – stability / energy

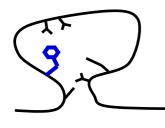
- stability does a molecule prefer to be folded or unfolded?
- what is unfolded?
 or
 ?

- my energy function tells me to change "X" to "Y"
 - it affects both the good and bad
 - has it affected the energy difference?
 - no guarantee
- my score function is like energy (potential or free)
 - certainly not identical

Problems - sidechains

- long topic next semester gross problems here
- side chain positions
 - can I ever calculate the energy if I change X to Y?
 - insert a phe into this structure
 - what interactions does it have?



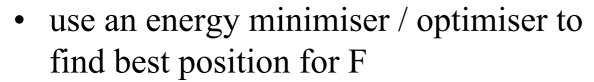




- how to cope with side chain positions in a practical way
 - optimise location of sidechains
 - use average
 - explicit rotamers

Sidechains – optimise at each step

- I start with known protein
 - change $A \rightarrow F$





- we have a gigantic search space
- explicit optimisation of one side chain would be expensive
- silly?
- I change $A \rightarrow F$, but the rest of the side chains may move
- bad idea



Sidechains – use averaging

- ignore the problem of sidechain geometry
- silly?
 - at room temperature, side chains move
 - small (middle of protein) to big (surface)
 - we cannot expect Å accuracy anyway
- implementation
 - functions which care about X interacting with Y
 - no attention to location of each atom
- rather fast searching
- what if we want to worry about atoms?

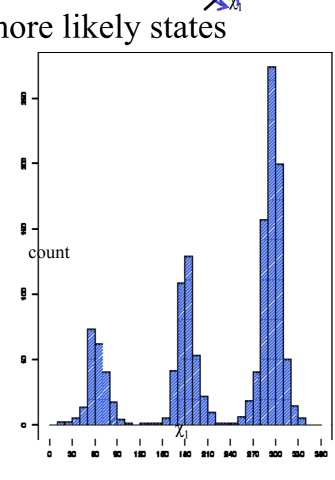
Sidechains – use rotamers

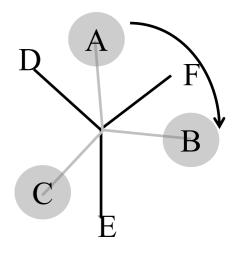
• sidechains can move anywhere but

• there are preferences in diagram – three more likely states

• how many times is the first angle (χ_1) seen at each angle ?

- how to use this?
 - look for most popular angles (60, 180, 300)



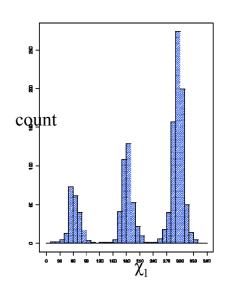


Sidechains – use rotamers

- For this example
 - do not have 1 cys residue
 - replace with cys1, cys2, cys3
 - treat all amino acids similarly
- more complicated because of more angles
- consequence
 - N_{type} of amino acids >> 20
- requires that you have a pre-built rotamer library



- Monte Carlo (random moves between residues or rotamers)
- dead end elimination (will remove impossible rotamers)



Problems – viability

- Designed sequences must
 - fold
 - be expressed + produced

Summary so far

- Experimental approaches
- Nature of the problem discrete (not continuous)
- Optimisation methods (MC, DEE)
 - more genetic algorithms
- Score functions
 - not energy, not free energy, not potential energy
- Success / state of the art
 - not many examples from literature
 - failure rate?
 - cost

More aims

- Useful and possible ?
 - changing solvents?
 - reactions in CH₃0H, ethanol, ...
 - may be possible experimentally
 - pH tolerant
 - washing detergent is basic
- Useful, but difficult
 - change activity / specificity
 - ribonuclease should cut after a different nucleotide