How does this sequence fold?

VQAVAVLKGDAGVSGVVKFEQASESEPTTVSYEIAGNSPNAERGFHIHEFGDATNGCVSA GPHFNPFKKTHGAPTDEVRHVGDMGNVKTDENGVAKGSFKDSLIKLIGPTSVVGRSVVIH AGQDDLGKGDTEESLKTGNAGPRPACGVIGLTN


## Lattice model

- model a protein as a chain of hydrophobic (H) and polar (P) residues

- a conformation is a self-avoiding walk on a 2D square lattice


Protein folding: schematic


## Conformations

The HP model: H• PO


## Ab initio structure prediction

Kim T. Simons, Charles Kooperberg, Enoch Huang and David Baker
"Assembly of Protein Tertiary Structures from Fragments with Similar Local Sequences using Simulated Annealing and Bayesian Scoring

## Functions

J. Mol. Biol., vol. 268, 209-225 (1997).

A simulated annealing procedure needs:
— method for generating structures

- scoring function

Graham Kemp, Chalmers University of Technolog

## Estimating P(sequence|structure)

Similar to scoring a sequence-fold match when threading
Profiles:

$$
\prod_{i} P\left(a a_{i} \mid E_{i}\right)
$$

Pairwise potentials:

$$
\prod_{i<j} P\left(a a_{i,}, a a_{j} \mid r_{i j}\right)
$$

Simons et al. (1997):

$$
\prod_{i} P\left(a a_{i} \mid E_{i}\right) \times \prod_{i<j} \frac{P\left(a a_{i,} a a_{j} \mid r_{i j}, E_{i,} E_{j}\right)}{P\left(a a_{i} \mid r_{i j}, E_{i,} E_{j}\right) P\left(a a_{j} \mid r_{i j} E_{i,} E_{j}\right)}
$$

The CKY algorithm — natural language

## Parsing natural language vs. folding a protein

Parsing natural language:
a) start with one-dimensional string of words;
b) consider all possible topologies representing possible relationships among words and phrases;
c) chooses the one that conveys the correct single meaning of the sentence.

## Folding a protein:

a) start with one-dimensional string of amino acid residues;
b) consider all possible topologies representing possible native substructures of a protein;
c) chooses the one that has the global minimum free energy.

```
Zipping and assembly
```

3. Extract the trees


Zipping and assembly with constraints: information used
Constraints used in modelling human p8MTCP

## Protein amino acid residue sequence

## Constraints

- Angle constraints:
- torsion angle ranges predicted from chemical shifts
- Distance constraints:
- main chain N and O involved in hydrogen bonds in secondary structures
- HN-HN NOEs from 4D NMR experiments
- from predicted secondary structure
- disulphide bridges
- no steric overlaps
- 


## Actual cells used in constructing one model


residue(1,'PHE'). esidue(2,'PHE') residue(3,'ASP') residue( $($, 'ASP') residue(4,'GLU').
residue(5,'LYS'). \% etc
disulphide_bond $(6,33)$ disulphide_bond $(13,27)$ disulphide_bond $(17,34)$.
alpha_helix(4,8).
antiparallel_bridge $(12,34)$ antiparallel_bridge $(14,32)$ antiparallel_bridge $(22,35)$ antiparallel_bridge $(25,33)$


## Additional rule:

disulphide ( $\mathrm{A}, \mathrm{B}$ ) :- disulphide_bond ( $\mathrm{A}, \mathrm{B}$ ). disulphide $(A, B)$ :- disulphide_bond $(B, A)$.
disulphide_distance_constraints :-
disulphide(A,B),
disulphide (C,D),
1 is C-B,
strand (StrandStart, StrandEnd),
B >= StrandStart,
C $=<$ StrandEnd,
assert (lower_distance_bound (
( $\left.\left.A,{ }^{\prime} \mathrm{CA} \mathrm{A}^{\prime}\right),\left(\mathrm{D}, \mathrm{C}^{\prime} \mathrm{CA}{ }^{\prime}\right), 13.0\right)$ ),
assert (upper_distance_bound (
( $A,{ }^{\prime}$ 'CA'), (D, 'CA'), 15.0)),
fail.


Human $\beta$-defensin 6: 50 best models
Claims made for ZAMDP method


All residues


Core residues: 4-35

- local-first-global-later explains quick folding, and avoidance of vast stretches of conformational space
- reflects parallel nature of physical kinetics
- captures relationship between contact order (whether contacts are mainly local or mainly non-local) and folding rate
- identifies slow- and fast-folding proteins, and slow- and fast-folding routes

