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**Protein Folding**

1. *Designed Random Energy Model (REM)*: Consider a protein model in which for a given sequence and structure, the energy is randomly taken from the Gaussian probability density

$$p(E) = \frac{1}{\sqrt{2\pi\Sigma^2}} \exp\left(-\frac{E^2}{2\Sigma^2}\right).$$

The total number of structures is  $\Omega_{str}$ , while the number of sequences is  $\Omega_{seq} \gg \Omega_{str}$ .

(a) A particular *sequence* has a (unique) native structure of energy  $E_N$ . Calculate and plot the energy  $E(T)$  of this sequence as a function of temperature  $T$ .

(b) For a particular *structure*, we attempt to design a good sequence by Monte Carlo sampling of representative sequences at a ‘temperature’  $\tau$ . Calculate and plot the designed native energies  $E_N(\tau)$  as a function of the design temperature  $\tau$ .

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2. *Folding time*: [Adapted from Gutin *et al.*, J. Chem. Phys. **108**, 6466 (1998).] Assume that to change from one compact structure to any other, the protein has to unfold to an intermediate flexible state of (higher) energy  $E^*$ . If the starting configuration is at an energy  $E$ , the typical (activation) time to overcome this energy barrier behaves as

$$t_0(E) = \tau \exp\left(\frac{E^* - E}{k_B T}\right),$$

where  $T$  is the temperature, and  $\tau$  is an elementary time step. The folding time is then related to the number of accessible states (hence entropy) to be explored, by

$$t_F(E) = t_0(E)n(E) = t_0(E) \exp\left(\frac{S(E)}{k_B}\right).$$

(a) Use a random energy model to calculate  $E$  and  $S$  as a function of temperature  $T$ .

(b) Calculate the folding time  $t_F(T)$ , and plot  $\ln t_F(T)$  as a function of  $1/T$ .

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3. *Amino-acid interactions*: What can we learn by combining the Random Energy Model with commonly used interaction potentials between amino acids?

(a) Find a  $20 \times 20$  matrix of interactions  $U(a, a')$  amongst amino acids, and calculate the mean  $\langle U \rangle$  and variance  $\langle U^2 \rangle_c$  of its elements. The commonly used Miyazawa–Jernigan

(MJ) interaction matrix can be found in S. Miyazawa and R.L. Jernigen, *J. Mol. Biol.* **256**, 623 (1996). (Table 3 of this publication is available in the assignments section.)

(b) Model the possible configurations of a protein by the ensemble of compact self-avoiding walks on a cubic lattice. (All lattice sites are visited by compact walks.) Calculate the number  $n$  of non-polymeric nearest neighbor interactions for such configurations on an  $N = L \times L \times L$  lattice, and deduce the ratio  $n/N$  for large  $N$ .

(c) The number of compact walks on a cubic lattice asymptotically grows as  $g^N$ , with  $g \approx 1.85$ . Use this in conjunction with the results from parts (a) and (b) to estimate the folding temperature  $T_c$  of a random sequence of amino-acids, and the corresponding energy  $E_c$ .

**(Optional)** (d) Select a protein, find its amino-acid sequence and construct a contact matrix corresponding to its structure. Use the interaction matrix from part (a) to estimate the energy of the native structure, and calculate the ratio  $E_N/E_c$ .

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**4. Analysis of protein structures:** Calculate  $\phi$  and  $\psi$  torsion angles in `Rasmol` for a given protein (see the commands below). Make  $(\phi, \psi)$  “Ramachandran” diagrams by plotting  $\phi$  along the  $x$  and  $\psi$  along the  $y$  axis; one  $(\phi, \psi)$  point for each amino acid.

(a) Do amino acids that are part of different secondary structure elements (helices, sheets) land in the same or different islands on the  $(\phi, \psi)$  diagram? You can find secondary structure elements in fields `HELIX` and `SHEET` of the protein structure file (aka PDB file). Explain your observations.

(b) Find amino acids that have unusual  $(\phi, \psi)$  angles (i.e. deviate from the many clouds of points). What types of amino acids tend to have “unusual”  $(\phi, \psi)$  conformation? Discuss.

(c) Visualize protein structure in `Rasmol`, following the sequence of commands below, and select those with “unusual”  $(\phi, \psi)$  conformation. Do they tend to be close to the ligand?

Some sample proteins to explore (PDB files provided on the Assignment page):

Hemoglobin (alpha chain) 4HHB\_A.PDB

Immunoglobulin domain 1TEN.PDB

You can use the following sequence of `Rasmol` commands to generate a good view of a protein, and the `fipsi.dat` file of  $(\phi, \psi)$  angles

```
set background white
wireframe off
ribbons
```

```
color structure
```

```
select ligand
```

```
cpk
```

```
color green
```

```
select protein
```

```
write RDF fipsi.dat
```

To select a particular set of amino acids, (e.g. 128 and 156) you can do the following

```
select 128,156
```

```
cpk
```

```
color red
```

```
*****
```