11: Catchup II Machine Learning and Real-world Data (MLRD)

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Lent 2018

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Last session: HMM in a biological application

- In the last session, we used an HMM as a way of approximating some aspects of protein structure.
- Today: catchup session 2.
- Very brief sketch of protein structure determination: including gamification and Monte Carlo methods.
- Related ideas are used in many very different machine learning applications ...

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What happens in catchup sessions?

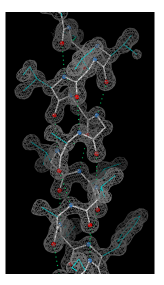
Lecture and demonstrated session scheduled as in normal session.

- Lecture material is non-examinable.
- Time for you to catch-up in demonstrated sessions or attempt some starred ticks.
- Demonstrators help as usual.

Protein structure

- Levels of structure:
 - Primary structure: sequence of amino acid residues.
 - Secondary structure: highly regular substructures, especially α-helix, β-sheet.
 - Tertiary structure: full 3-D structure.
- In the cell: an amino acid sequence (as encoded by DNA) is produced and folds itself into a protein.
- Secondary and tertiary structure crucial for protein to operate correctly.
- Some diseases thought to be caused by problems in protein folding.

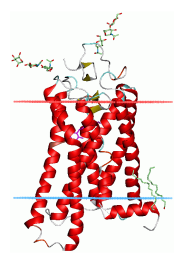
Alpha helix



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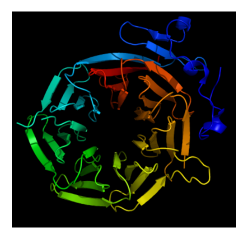
Bovine rhodopsin



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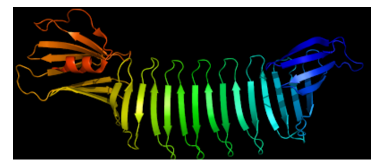
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7-bladed propeller fold



http://beautifulproteins.blogspot.co.uk/

Peptide self-assembly mimic scaffold: an engineered protein



http://beautifulproteins.blogspot.co.uk/

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Protein folding

- Anfinsen's hypothesis: the structure a protein forms in nature is the global minimum of the free energy and is determined by the animo acid sequence.
- Levinthal's paradox: protein folding takes milliseconds not enough time to explore the space and find the global minimum. Therefore kinetic function must be important.

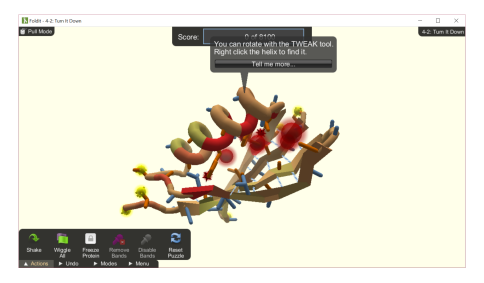
Protein structure determination and prediction

- Primary structure may be determined directly or from DNA sequencing: relatively easy.
- Secondary and tertiary structure can be determined by x-ray crystallography and other direct methods, but difficult, expensive, time-consuming.
- Given amino acid sequence, can we predict the structure? i.e., determine how the protein will fold.
- Secondary structure prediction is relatively tractable: various prediction methods, including HMMs (cf last session).
- Tertiary structure prediction is very difficult.

Protein tertiary structure prediction

- Modelling protein structure fully is hugely computationally expensive.
- Ideally, should model all the water molecules too
- Several approaches, including:
 - 1 Molecular Dynamics (MD): modelling chemistry. folding@home: use home computers to run simulations.
 - 2 Foldit: get lots of humans to work on the problem (an example of **gamification**).
 - 3 Use **Monte Carlo methods** (repeated random sampling) to explore possibilities.

Foldit: combined human-computer intelligence



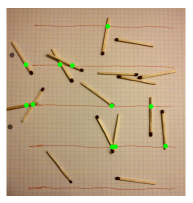
Monte Carlo methods in protein structure prediction

- Objective: find lowest energy state of protein.
- Idea: start with secondary structure, try (pseudo)random move, see if result is lower energy and repeat.
- Problem: local minima locally good move may not be part of best solution.
- So: also sometimes accept a move that increases energy.
- Specific approach Metropolis-Hastings: a type of Markov Chain Monte Carlo method.

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Monte Carlo methods in general

- Using random sampling to solve intractable numerical problems.
- Earliest example: Buffon's needle for estimating π



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Monte Carlo methods

- Physicists developed modern Monte Carlo methods at Los Alamos: programmed into ENIAC by von Neumann.
- Bayesian statistical inference not until 1993 (Gordon et al): essential for many modern machine learning approaches.
- Gibbs sampling is a special case of Metropolis-Hastings.
- More about this in later courses: Mathematical Methods, Machine Learning and Bayesian Inference, Bioinformatics.

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Practical introduction by Geyer in

http://www.mcmchandbook.net/HandbookTableofContents.html