**Introduction to Bioinformatics**

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**Protein Modeling Methods**

- **Ab initio methods:**
  - solution of a protein folding problem
  - search in conformational space

- **Energy-based methods:**
  - energy minimization
  - molecular simulation

- **Knowledge-based methods:**
  - homology modeling
  - fold recognition

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**Knowledge**

Knowledge is a pattern that exceeds certain threshold of interestingness.

Factors that contribute to interestingness:
- coverage
- confidence
- statistical significance
- simplicity
- unexpectedness
- actionability

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**Knowledge-based methods**

- Finding patterns in known structures
- Deriving rules (usually in the form of PMF)
- Applying the rules

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**Protein representation (Crambin)**

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Neighbor identification in proteins: Voronoi/Delaunay Tessellation in 2D

Delaunay simplex is defined by points, whose Voronoi polyhedra have common vertex.

Delaunay simplex is always a triangle in a 2D space and a tetrahedron in a 3D space.

Delaunay tessellation of Crambin
Dealunay simplices classification

Fold Recognition

Pattern searching
  sequence patterns
  structure patterns
  residue composition patterns

Threading
  sequence-structure compatibility
  structure-sequence compatibility

Threading
  • Only the local environment is taken into account
  • Non-local contacts are assumed with generic peptide
  • No gaps are allowed in the alignment

Homology Modeling

  • Identification of structurally conserved regions (using multiple alignment)
  • Backbone construction (based on SCR)
  • Loop construction (KB or conformational search)
  • Side-chain restoration (KB, rotamer, or MM)
  • Structure verification and evaluation
  • Structure refinement (energy minimization)

Homology Modeling Programs

  Modeller
  (http://guitar.rockefeller.edu/modeller)
  Swiss-Model
  (http://www.expasy.ch/swissmod)
  Whatif
  (http://www.cmbi.kun.nl/whatif)
Swiss-Model

- **Method:**
  Knowledge-based approach.

- **Requirements:**
  At least one known 3D-structure of a related protein.
  Good quality sequence alignments.

- **Procedures:**
  Superposition of related 3D-structures.
  Generation of a multiple alignment.
  Generation of a framework for the new sequence.
  Rebuild lacking loops.
  Complete and correct backbone.
  Correct and rebuild side chains.
  Verify model structure quality and check packing.
  Refine structure by energy minimisation and molecular dynamics.

Methods and Programs used by Swiss-Model

- **Sequence Alignment**

- **Knowledge Based Protein Modelling**

- **Energy Minimisation**
  Gromos96 (van Gunsteren W.F. http://igc.ethz.ch/gromos/)

- **Model evaluation**
  Swiss-PdbViewer
  (http://www.expasy.ch/spdbv/mainpage.html)

Swiss-Model Request Types

- First Approach mode.
- Optimise mode.
- Combine mode.
- GPCR mode.

Model Confidence Factors

The Model B-factors are determined as follows:

- The number of template structures used for model building.
- The deviation of the model from the template structures.
- The Distance trap value used for framework building.

The Model B-factor is computed as:

\[ 85.0 \times \left( \frac{1}{\text{# selected template str.}} \right) \times \left( \frac{\text{Distance trap}}{2.5} \right) \]

and

99.9 for all atoms added during loop and side-chain building