In Silico protein structure and function prediction

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Slides: http://www.trhvidsten.com/Teaching.html

Proteins play key roles in a living system

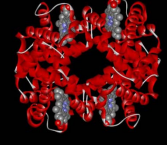
Three examples of protein functions

- Catalysis:
 Almost all chemical reactions in a living cell are catalyzed by protein enzymes
- Transport:
 Some proteins transports various
 substances, such as oxygen, ions, and so on
- Information transfer:For example, hormones



Alcohol dehydrogenase oxidizes alcohols to aldehydes or ketones

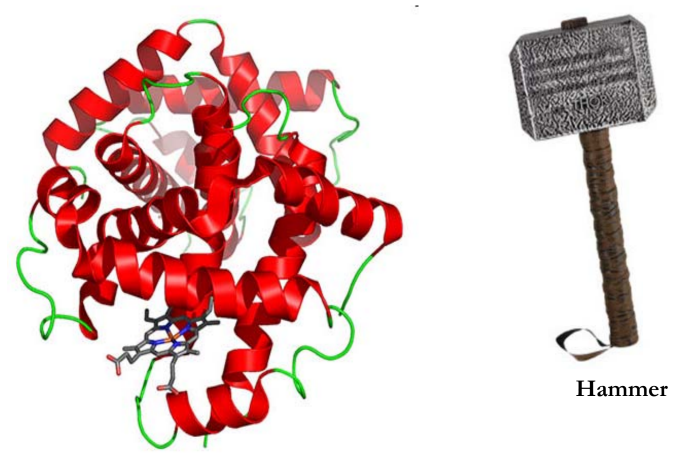
Haemoglobin carries oxygen



Insulin controls the amount of sugar in the blood

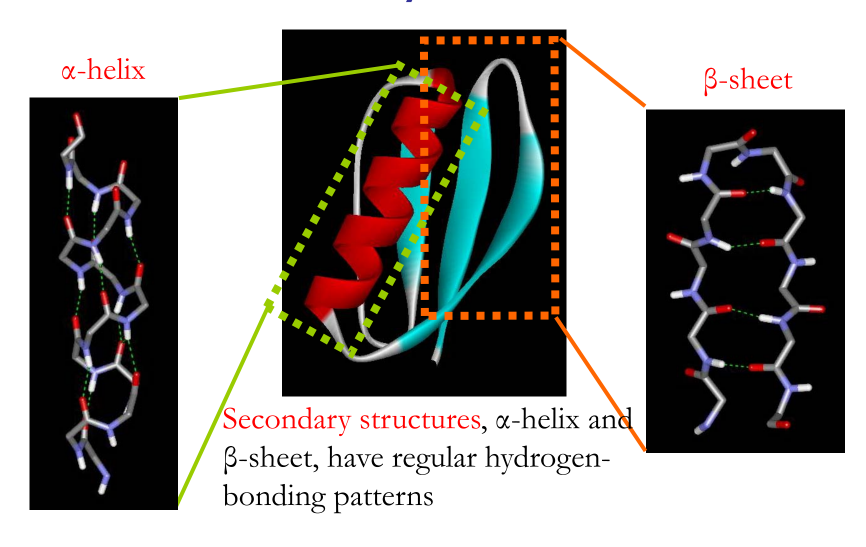
Structure - function

The 3D shape (and chemical properties) of proteins determine their function

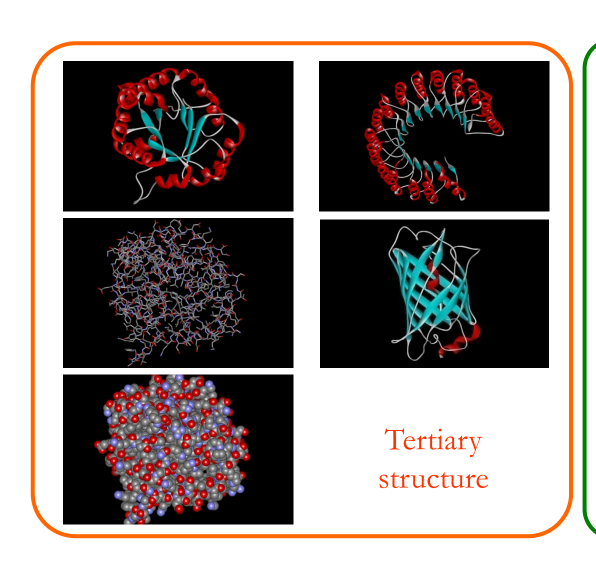


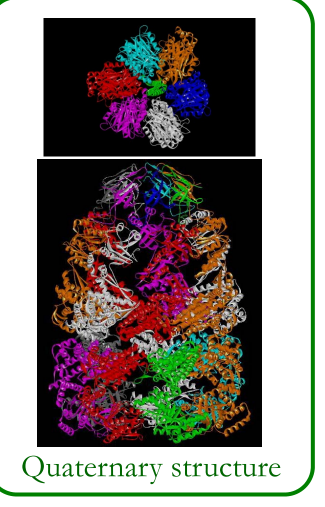
Hemoglobin

Basic structural units of proteins: Secondary structure



Three-dimensional structure of proteins





Hierarchical nature of protein structure

Primary structure (Amino acid sequence) Secondary structure (α -helix, β -sheet) Tertiary structure (Three-dimensional structure formed by assembly of secondary structures) Quaternary structure (Structure formed by more than one polypeptide chains)

Domains: recurrent units of proteins

- The same or similar domains are found in different proteins
- Each domain has a well determined compact structure and performs a specific function
- ➤ Proteins evolve through the duplication and domain shuffling

Protein domains can be defined based on:

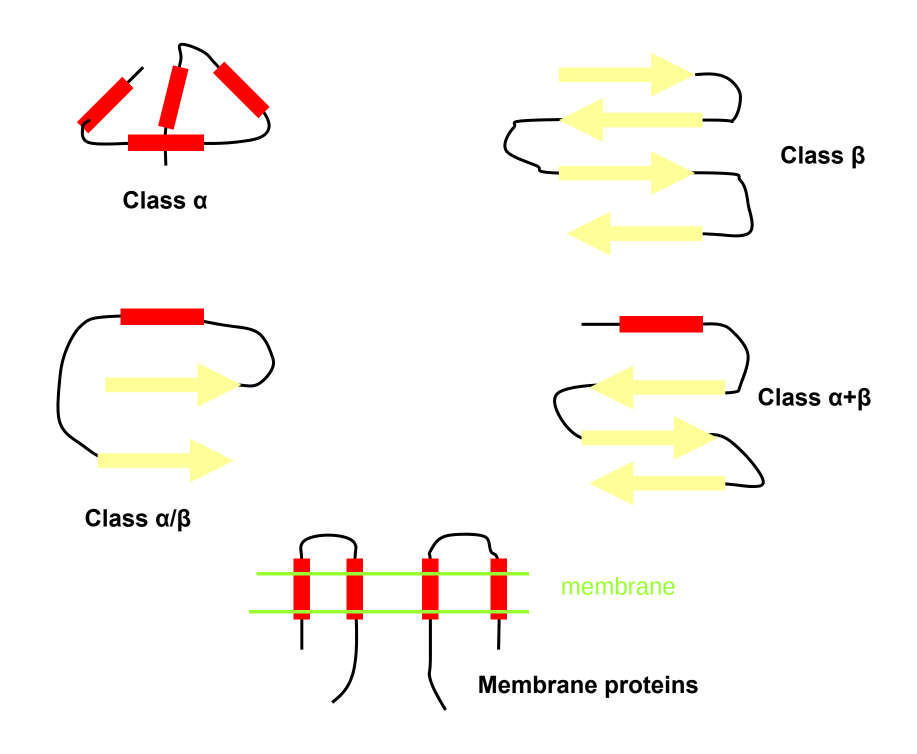
- Geometry: group of residues with a high contact density, number of contacts within domains is higher than the number of contacts between domains
- > Kinetics: domain as an independently folding unit
- Physics: domain as a rigid body linked to other domains by flexible linkers
- ➤ Genetics: minimal fragment of gene that is capable of performing a specific function

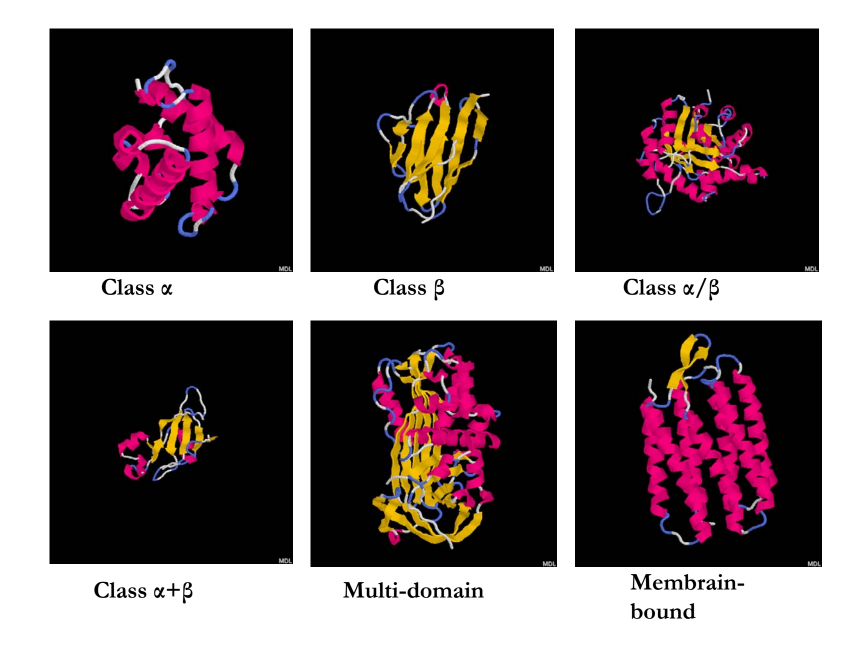
Protein folds

- \triangleright One domain \rightarrow one fold
- Fold definition: two folds are similar if they have a similar topology: arrangement/orientation of secondary structure elements (architecture) and connectivity
 - topology = architecture + connectivity
- Fold classification: structural similarity between folds is found using structure-structure comparison algorithms

Domain/fold classification

- Class α: a bundle of α helices connected by loops on the surface of protein
- \triangleright Class β : antiparallel β sheets
- \triangleright Class α/β: mainly parallel β sheets with intervening α helices
- \triangleright Class α+β: mainly segregated α helices and antiparallel β sheets
- Multidomain proteins: comprise domains representing more than one of the above four classes
- Membrane and cell-surface proteins: α helices (hydrophobic) with a particular length range, traversing a membrane

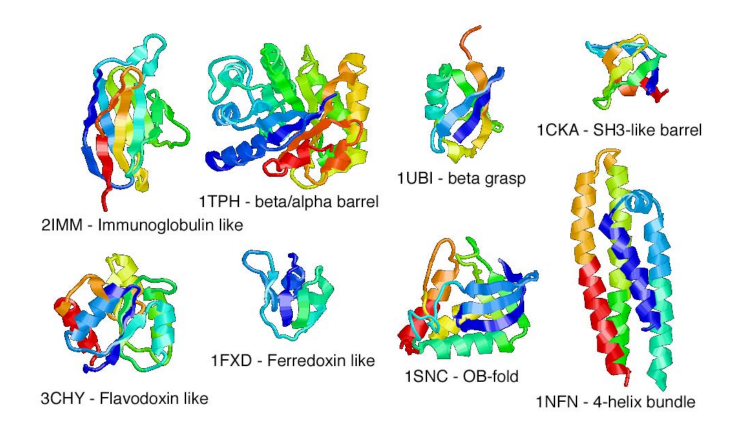




Structural classification of proteins (SCOP)

- The SCOP database aims to provide a detailed and comprehensive description of the structural and evolutionary relationships between all proteins whose structure is known.
- Created by manual inspection and aided by automated methods
- Consists of four hierarchical categories:
 - Class, Fold, Superfamily and Family.

SCOP

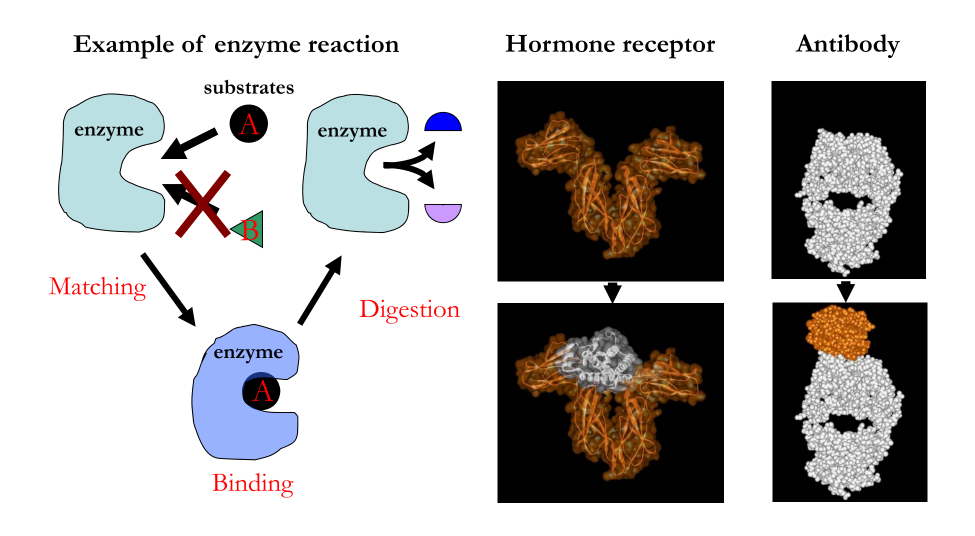


The eight most frequent SCOP folds

Why study structure?

- A full understanding of a molecular system comes from careful examination of the sequence-structure-function triad
- Below 30% protein sequence identity detection of a homologous relationship is not guaranteed by sequence alone
- Structure is much more conserved than sequence
- > However:
- A non-redundant set of sequences is different than a non-redundant set of structures is different than a non-redundant set of functions

The structure-function relationship

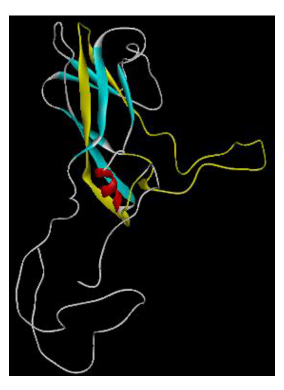


Structure-function relationships

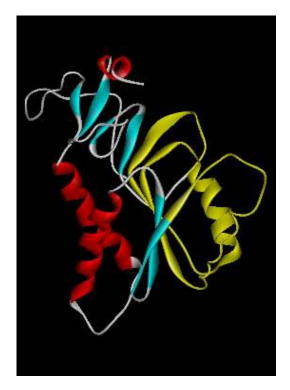
- The golden rule is there are no golden rules George Bernard Shaw
 - Complication comes from one structure multiple functions
 - Some folds are promiscuous and adopt many different functions superfolds
- Above 40% sequence identity, sequences tend to have the same structure and function but there are exceptions
- Structure and function tend to diverge at ~ 25% sequence identity
- The structure-function relationship is even more complex than the relationship between sequence and structure (and not as well understood)

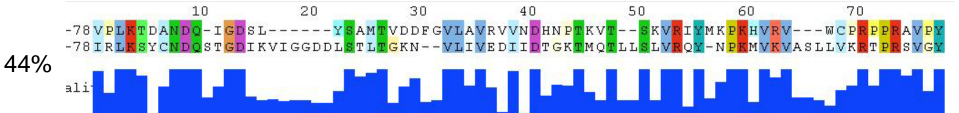
Similar sequences – different structures

1PIV:1 Viral Capsid Protein

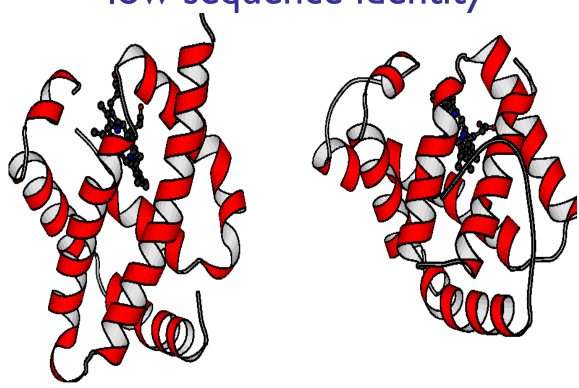


1HMP:A Glycosyltransferase



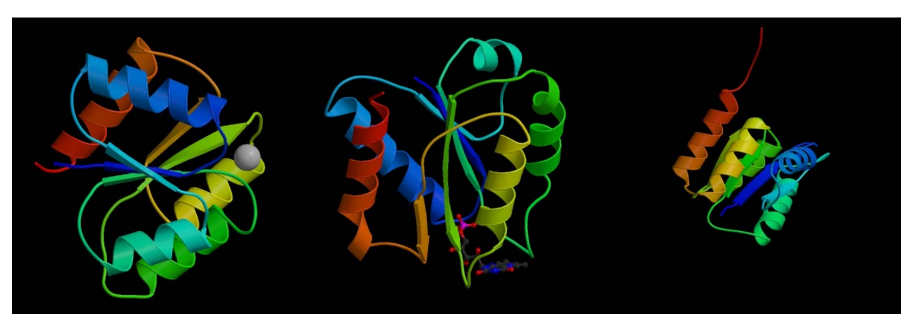


Same structure and function – low sequence identity



The globin fold is resilient to amino acid changes. *V. stercoraria* (bacterial) hemoglobin (left) and *P. marinus* (eukaryotic) hemoglobin (right) share just 8% sequence identity, but their overall fold and function is identical.

Similar structure - different function

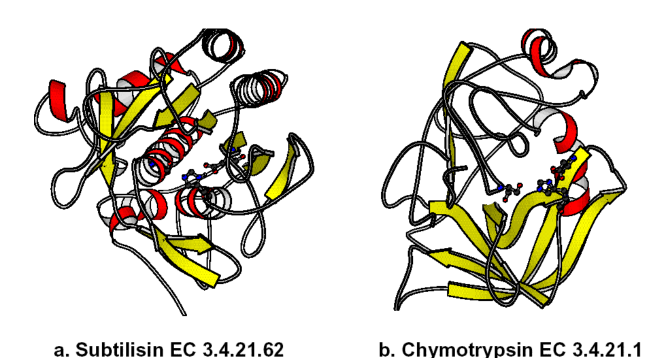


1ymv CheY Signal Transduction 1fla Flavodoxin Electron Transport

1pdo Mannose Transporter

Less than 15% sequence identity

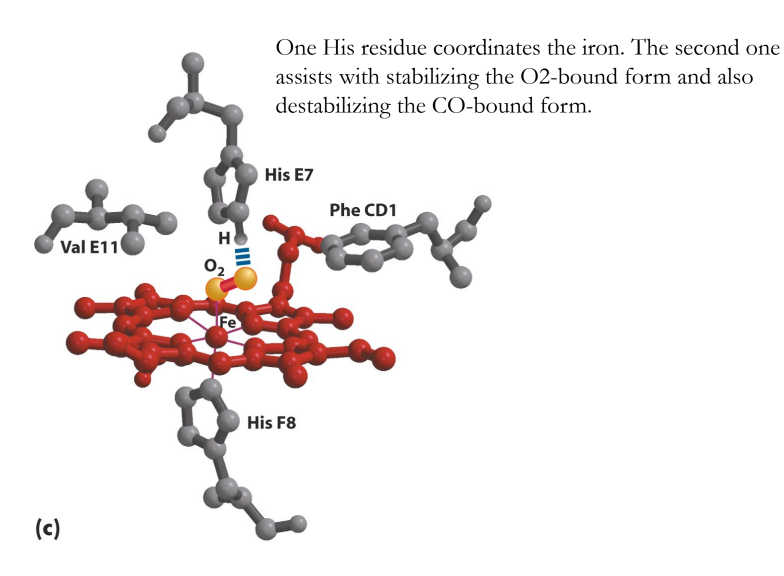
Convergent evolution



Subtilisin and chymotrypsin are both serine endopeptidases. They share no sequence identity, and their folds are unrelated. However, they have an identical, three-dimensionally conserved Ser-His-Asp catalytic triad, which catalyses peptide bond hydrolysis. These two

enzymes are a classic example of convergent evolution.

Functional sites: Oxygen-binding site



Computational function prediction methods

Major challenges

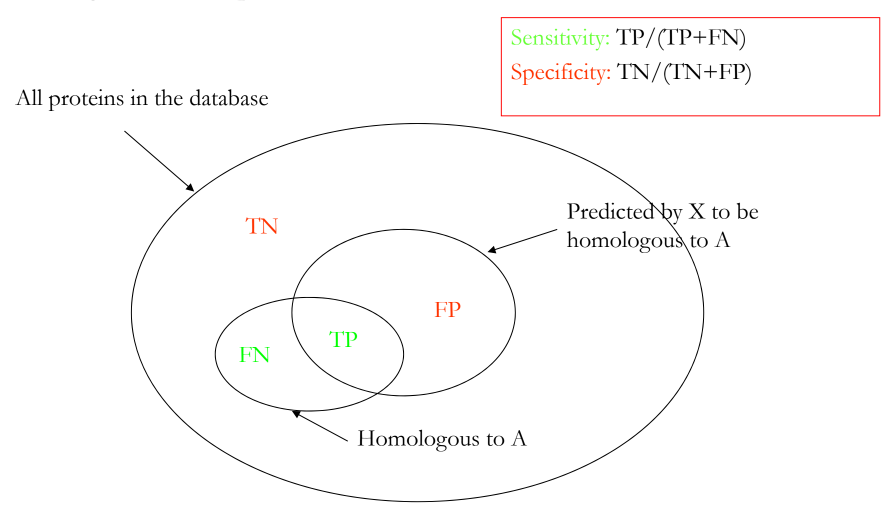
- The multifunctional nature of proteins
 - → proteins have multiple domains hosting different function
 - → some domain host several functions
- The functional sites in proteins may be
 - better conserved than global sequence
 - → low sequence similarity between functionally similar proteins
 - better conserved than global fold
 - → the same function may be hosted by different folds
- ... but in some cases functional sites may be
 - less conserved than global sequence
 - → highly similar sequences do not have the same function
 - less conserved than global fold
 - → the same fold may host different functions

Computational function prediction methods

- > Sequence-based
- Sequence alignment: Transfer function information from a known protein with high sequence similarity to the target
- Sequence-motifs: Extract function-specific sequence profiles from conserved sites and use these to assign functional classes to targets
- > Structure-based
- Structure alignment: Transfer function information from a known protein with high structure similarity to the target
- Structure-motif: Use 3D templates of functional sites, scan the target structure and assign function

Power of computational methods

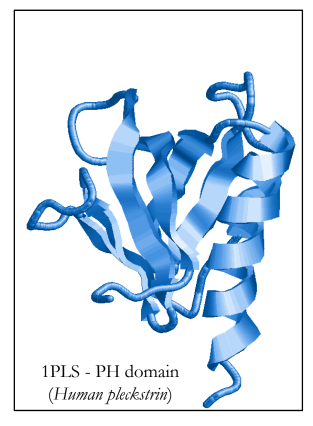
You want to find homologous proteins to a specific protein A using some computational method X:

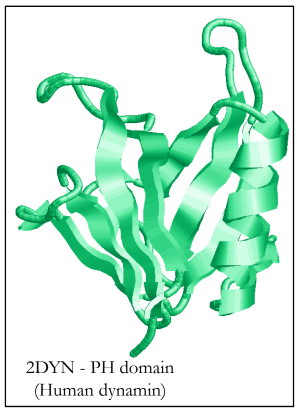


Example method: Global structure similarity

1PLS/2DYN:

23% sequence identity

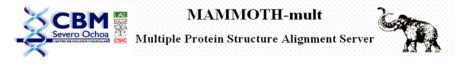




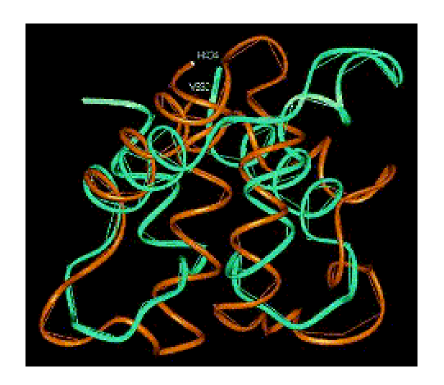
Example method: Global structure similarity

Dali

http://ekhidna.biocenter.helsinki.fi/dali_server/



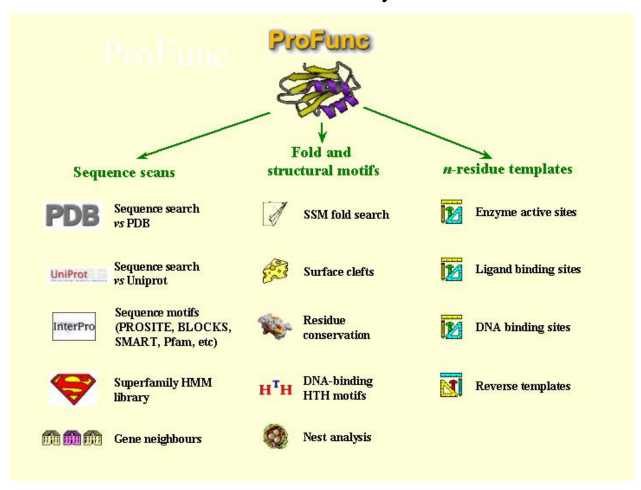
http://ub.cbm.uam.es/mammoth/pair/index3.php



Structural similarity between Calmodulin and Acetylcholinesterase

Example method: ProFunc

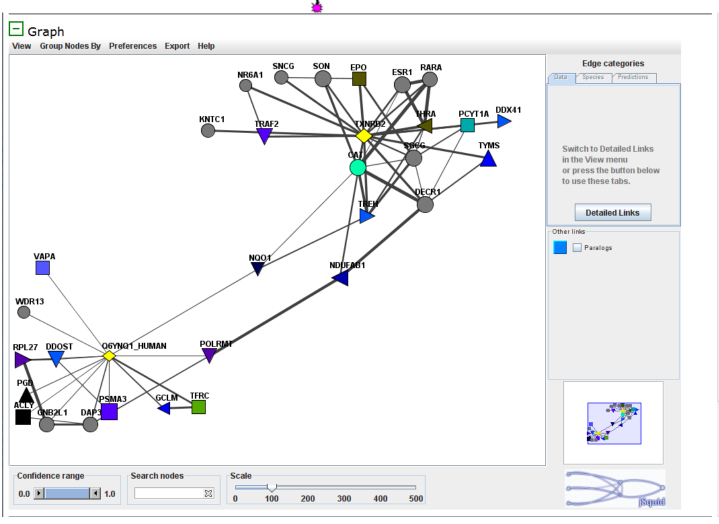
Successful function prediction methods are typically metaservers that combine many methods



http://www.ebi.ac.uk/thornton-srv/databases/ProFunc/

Example method:

FUNL&Up networks of functional coupling

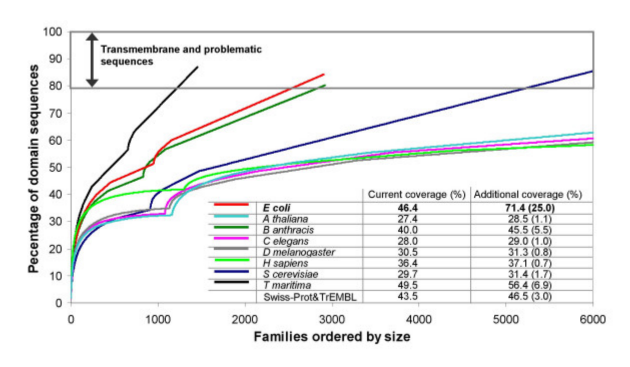


http://funcoup.sbc.su.se

Structural Genomics

- The biggest limitation for predicting function from structure is the low availability of structure information
- ➤ Solution: Structural genomics
 - Solve experimentally the structure for a representative set of all protein sequences, e.g., one or a few proteins from each fold
 - Predict the structure for the remaining sequences using homology modeling, i.e., transfer structure from a structurally solved homology
 - Predict function from structure
- > Structure prediction methods are better at predicting the core of proteins than the loops

Structural Genomics



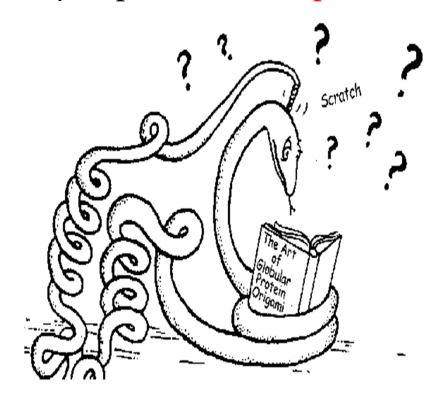
Marsden, Lewis and Orengo. Towards a comprehensive structural coverage of completed genomes: a structural genomics viewpoint. BMC Bioinformatics8: 86, 2007.

A domain sequence is structurally annotated if it can be assigned to a CATH or Pfam-A_struc family through the use of hidden Markov model searches

The protein folding problem

Anfinsen's thermodynamic hypothesis (1973): Protein folding is a strictly physical process that solely depends on the protein sequence





The folding problem:

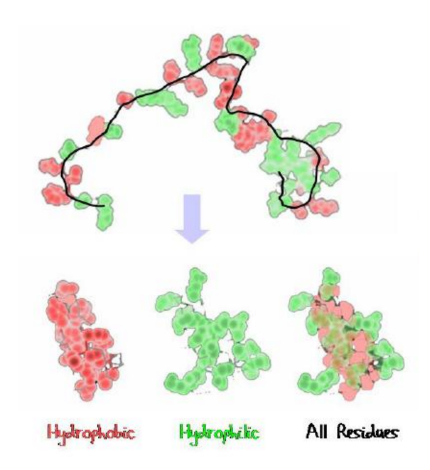
discover nature's algorithm for specifying 3D structure of proteins from their amino acid sequences

Hydrophobic interactions (I)

- Atomic charges dictate how folds occur
- Groups of C-H atoms have little charge
 - Called hydrophobic or non-polar
- > Hydrophobic groups pack together
 - To avoid contact with solvent (aqueous solution)
 - To minimise energy
- Hydrophobic and hydrophilic regions are the main driving force behind the folding process

Hydrophobic interactions (II)

- Hydrophobicity vs. hydrophilicity
- ➤ Van der Waals interaction
- > Electrostatic interaction
- > Hydrogen bonds
- ➤ Disulfide bonds



Folding is directed mainly by internal residues

- Mutations that change surface residues are accepted more frequently and are less likely to affect protein conformations than are changes of internal residues
- This is consistent with the idea of hydrophobic forcedriven folding

Molten globule

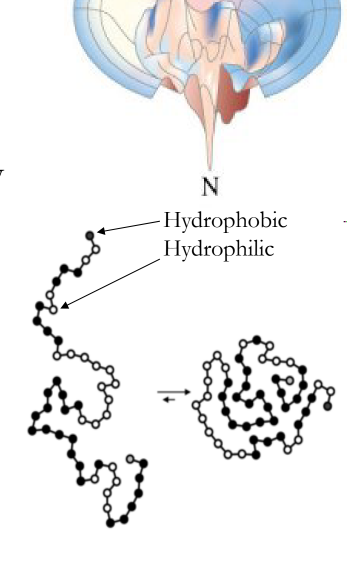
- Phase 1: Much of the secondary structure that is present in a native proteins forms within a few milliseconds
- ➤ Phase 2: Hydrophobic collapse into the Molten globule
 - Slightly larger (5-15% in radius) than the native conformation
 - Significant amount of secondary structure formed
 - Side chains are still not ordered/packed
 - Structure fluctuation is much larger not very thermodynamically stable

Computational folding methods

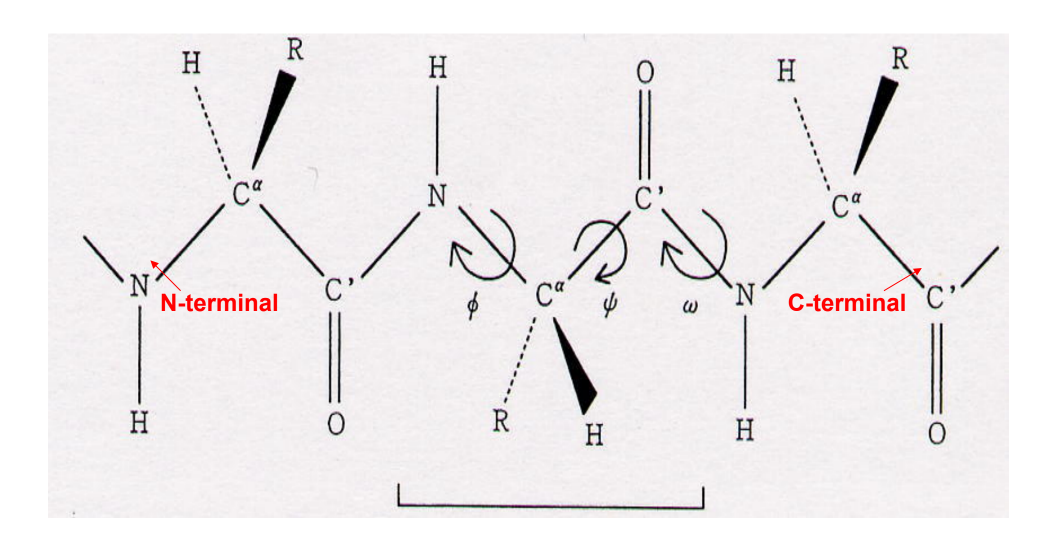
• No effective folding machine exists that is based on physical principles and energy minimization alone

 Current computational methods rely on known protein structures – machine learning approach:

- Template-based modeling
- Template-free modeling



Structure represented by angels



Protein folding

- > Levinthal's paradox
 - If for each residue there are only two degrees of freedom (ψ,ϕ)
 - Assume each can have only 3 stable values
 - This leads to 3^{2n} possible conformations
 - If a protein can explore 10¹³ conformation per second (10 per picosecond)
 - Still requires an astronomical amount of time to fold a protein
- Conclusion: proteins must fold in a way that does not randomly explore each possible conformations!

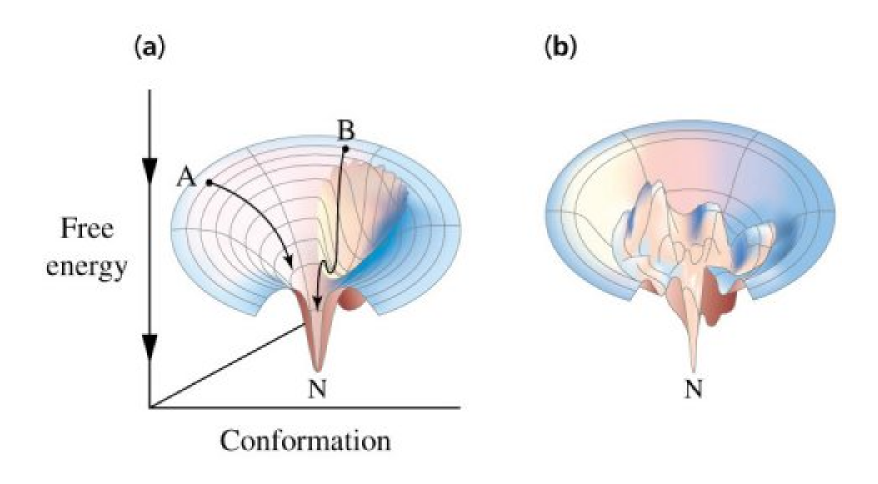
Structure prediction

- Protein structure prediction is the "holy grail" of bioinformatics
- Since structure is so important for function, solving the structure prediction problem should allow protein design, design of inhibitors, etc
- ➤ Huge amounts of genome data what are the functions of all of these proteins?

Assumptions

- Assumption 1: All the information about the structure of a protein is contained in its sequence of amino acids
- Assumption 2: The structure that a (globular) protein folds into is the structure with the lowest free energy
- Finding native-like conformations require:
 - A scoring function (potential)
 - A search strategy.

The free energy surface of a protein



Physics-based protein simulation

- All atom quantum mechanics (QM) calculation is not feasible
- > QM can be applied to a small set of atoms
 - Modeling of an active site
 - Can get total energies (binding vs. non-binding, pK_a etc.), wave function (charge distribution)
 - QM/MM simulations (i.e. remaining atoms are treated with <u>M</u>olecular <u>M</u>echanics)

Problems

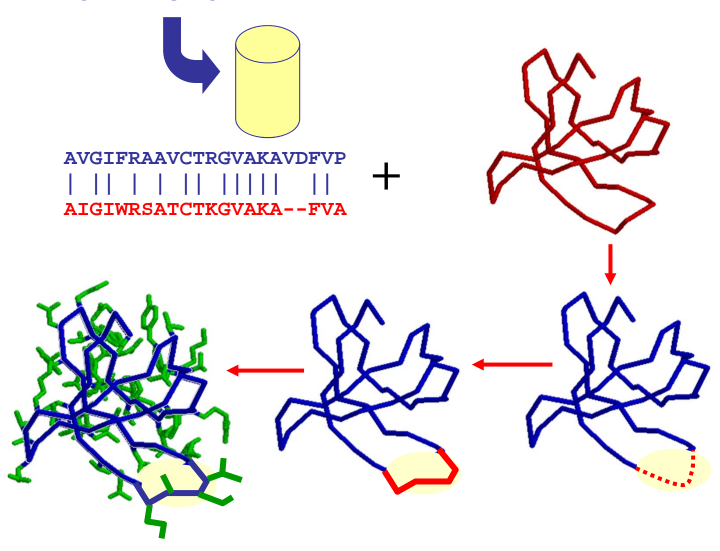
- ➤ Is the energy function correct?
 - Precise enough to discriminate non-native structure.
 - Yet simple enough for computers to carry out efficiently.
- ➤ Is the conformational search good enough to cover the global minimum?
- Protein folding without any prior knowledge about protein structure is a difficult task.
- ➤ Protein structure prediction is often quoted as an "NP complete problem", i.e. the complexity of the problem grows exponentially as the number of residues increases

Flavors of "knowledge-based" structure prediction

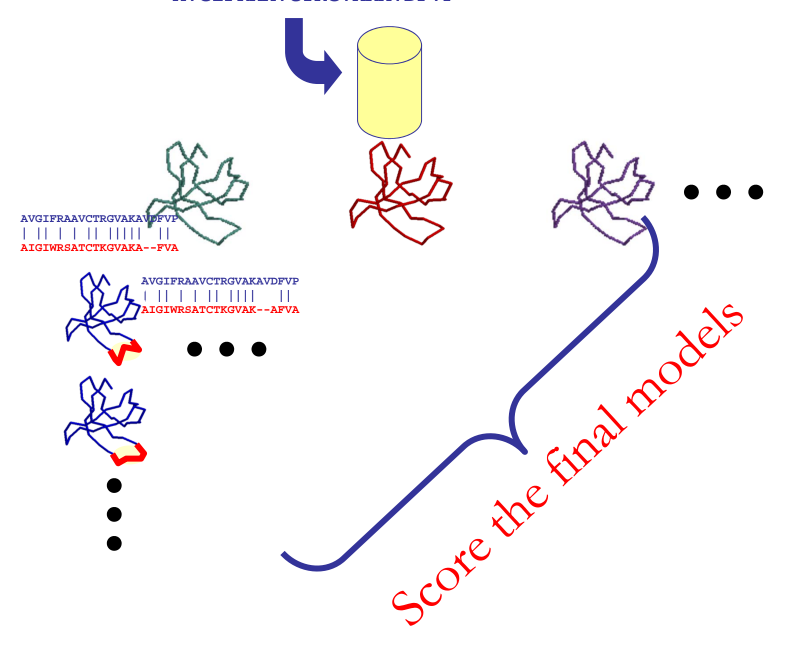
- > Experimental data
 - X-ray crystallography
 - NMR spectroscopy
- > Computational methods
 - Homology/comparative modeling
 - Fold recognition (threading)
 - Ab initio (de novo, new folds) methods (Ab initio: "from the beginning".

Comparative modeling

AVGIFRAAVCTRGVAKAVDFVP

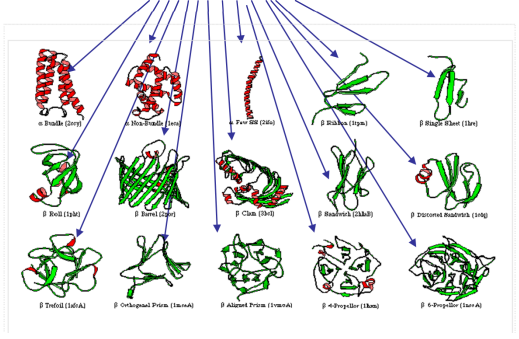


AVGIFRAAVCTRGVAKAVDFVP



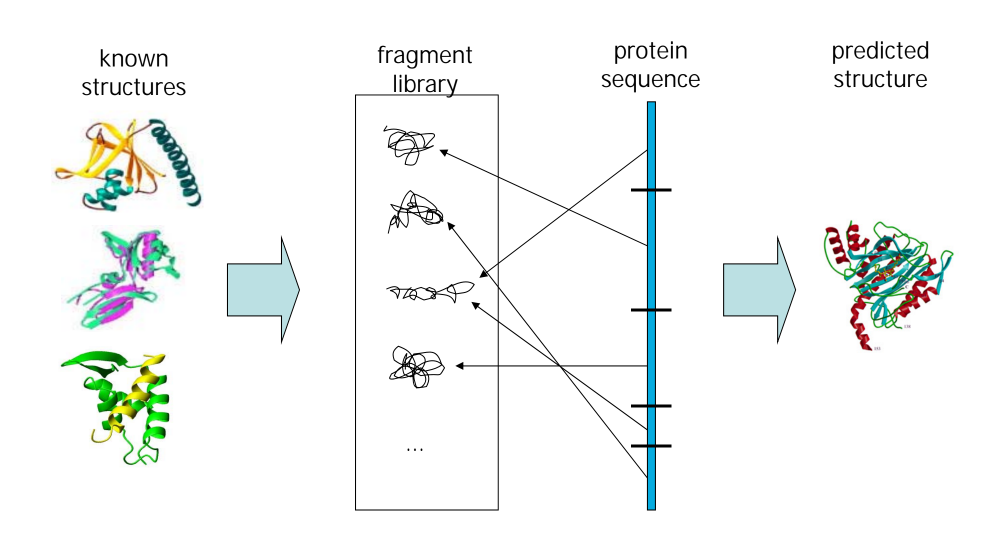
Fold recognition

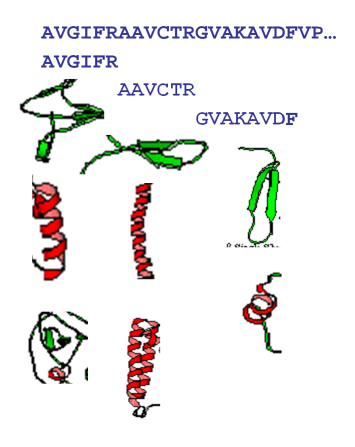
AVGIFRAAVCTRGVAKAVDFVPVESMETTMRSPV FTDNSSPPAVPQSFQVAHLHAPTGSGKSTKVPAA YAAQGYKVLVLNPSVAATLGFGAYMSKAHGIDPN IRTGVRTITTGAPVTYSTYGKFLADGGCSGGAYD IIICDECHSTDSTTILGIGTVLDQAETAGARLVV LATATPPGSVTVPHPNIEEVALSNTGEIP



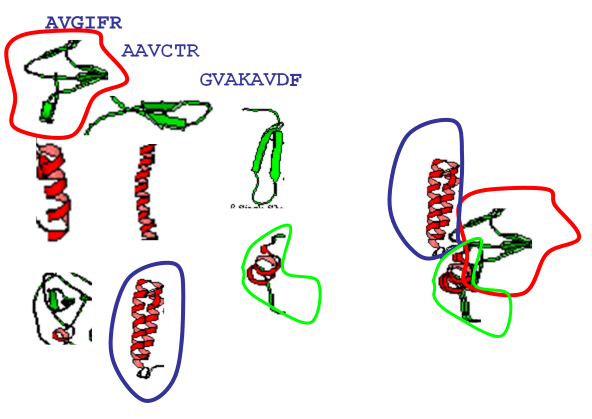
Score and select model

Fragment assembly

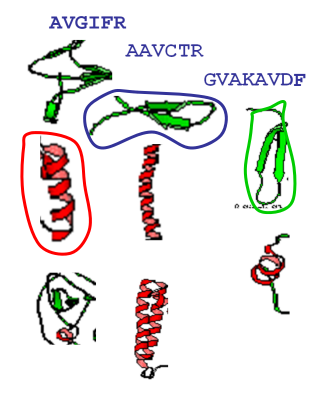


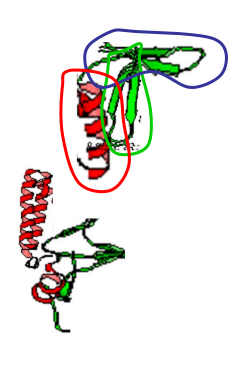


AVGIFRAAVCTRGVAKAVDFVP...



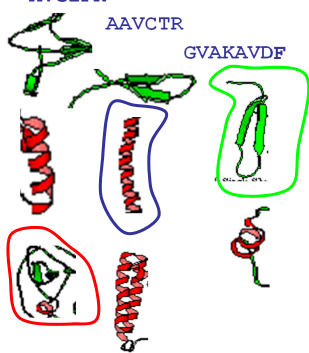
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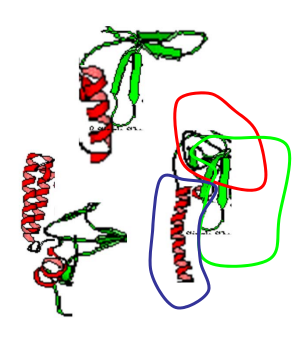


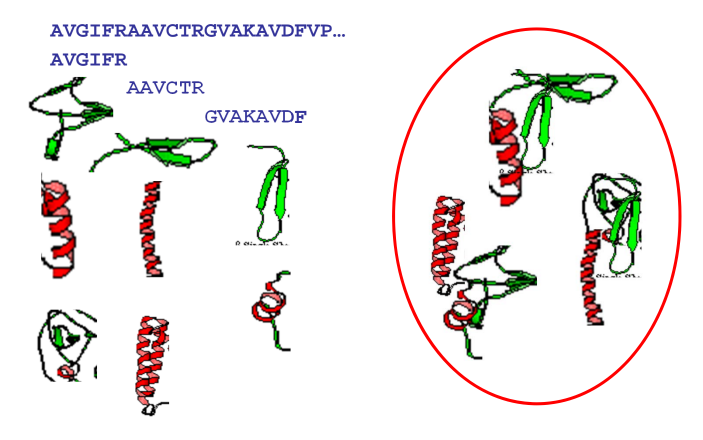


AVGIFRAAVCTRGVAKAVDFVP...

AVGIFR







Score and select model

CASP: Community Wide Experiment on the Critical Assessment of Techniques for Protein Structure Prediction

http://www.predictioncenter.org/

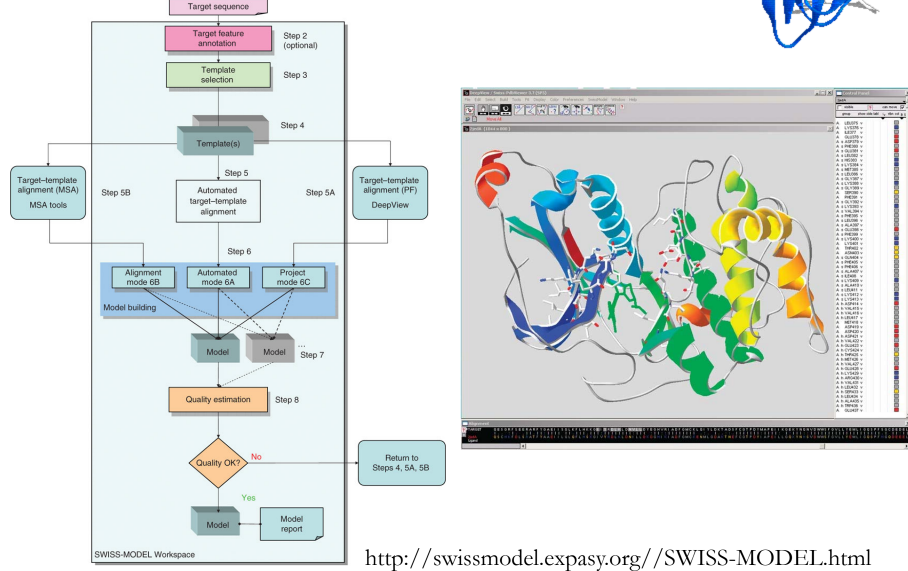
- Aim: obtain an in-depth and objective assessment of our current abilities and inabilities in the area of protein structure prediction
- Participants will predict the structure of a set of sequences soon to be known structures
- These will be true predictions, not 'post-dictions' made on already known structures.

Meta-methods

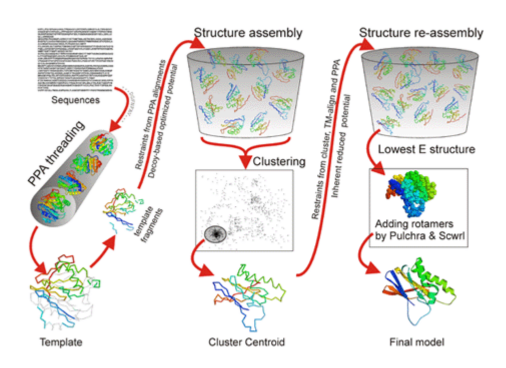
- Meta-methods combine predictions from individual methods
 - E.g. 3D-Jury: http://bioinfo.pl/Meta/
- Range from methods that select the best prediction to methods that improve and combine other predictions
- ➤ Often include methods for all flavors of protein structure prediction

SWISS-MODEL





I-TASSER

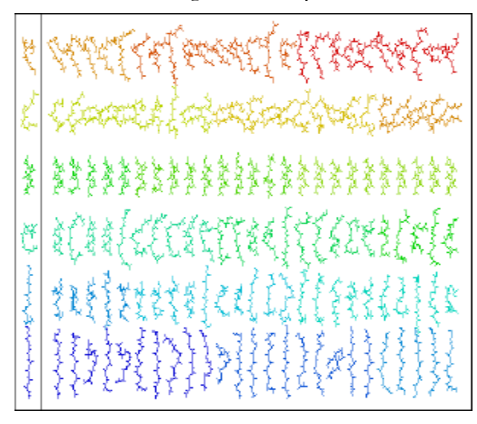


http://zhang.bioinformatics.ku.edu/I-TASSER/

Rosetta/Robetta

- Decoys are assembled from fragments
- Lowest energy model from a set of generated decoys is selected as the prediction
- ➤ Monte Carlo simulated annealing
- Physical energy function with elements of a statistical potential

Fragment library



CASP: progress

- Most progress in the fold prediction category and for servers over humans
- ➤ GDT_TS = (GDT_P1 + GDT_P2 + GDT_P4 + GDT_P8)/4, where GDT_Pn denotes percent of residues under distance cutoff <= nÅ

Kryshtafovych, Venclovas, Fidelis and Moult. Progress Over the First Decade of CASP Experiments. PROTEINS: Structure, Function, and Bioinformatics Suppl 7:225–236, 2005.

