Laboratory of Bioinformatics 1

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Course organization

- Lab1 in the first (Lab1a) and second semester are actually separate courses
- In february there is a 1 week intensive course run by professor Allegravia, which is part of Lab1a
 - It is about protein-protein interaction
 - We will have an exam also for this part
- There will be a written test on 09/01/20
- We have to write a report for 10/01/20 or 22/01/20
 - If we submit for the first deadline we will have feedback
- The oral defence will be in the last week of January
 - Probably it will be 29-31/01/20

Introduction

- The reference source in the filed is the Journal of Bioinformatics
- Functional annotation is the core of this course
- Functional annotation studies the relationship between structure and function
- Functional annotation requires data collection, storage and analysis
- Functional annotation is the activity of attributing structural and functional features to translated protein sequences
- Before starting data analysis, be sure of the quality of your data (!)
- A database must be implemented, curated and mined
- Database curation refers to updating the data and to keeping them compliant with the database standard
- A database release is its content at a given date
- Data mining is the retrieval of information from a database
 - It is done with a browser
- In Hamburg there is an EMBL faicility that uses X-rays in a flow citometer

X-ray cristallography

- I can understand that I have a protein crystal by shining light on it and seeing how it diffracts
- Routinely X-ray diffraction is not able to locate hydrogens
- Each crystal has a unit cell
- Electron density is the result of data analysis on diffraction maps
- The main cristallographic techniques are NMR and X-rays
- X-ray sources used for studying protein structure are synchrotrons or more traditional lab sources
- Synchrotrons are located in the US, Europe or Japan
- Sychrothron source give an anisotrope beam, which gives semicircles in the diffraction map instead of focused spot
- In order to have a diffraction pattern you need to have interference between the diffracted beams

- Bragg diffraction law: $2d * \sin \theta = n\lambda$
 - When the same beam is reflected by 2 planes, the part which is reflected by the lower plane travels for a longer distance
 - The distance is exactly $2d * sin\theta$, where θ is the angle of incidence and d is the distance between the planes
 - If this distance is equal to n wavelenghts, we observe reflection because of interference
 - By knowing all the terms except d, I can derive the minimal distance between the diffraction planes
- A typical diffraction maps is organized with 3 coordinates (H, K, L) and an intensity dimension (I)
 - The intesity is proportional to the amplitude, therefore to the amount of costructive interference
 - The 3 coordinates reflect the facts that we are operating in 3 dimensions
 - I can recove the electron densities from the diffraction pattern with the Fourier transform
 - * It is a computation-heavy task
- I cannot recover the phase from the diffraction map
 - In a synchrotron, I can recover the phase of the wave with the anisotropy approach
- Once I have the electron density, I need to fit my molecule in it to determin the conformation
 - This is easier if my electron density has an high resolution
 - I can take advantage of similar proteins with a similar structure to do the fitting
 - This fitting procedure is called refinement, because it reduces noise in the model
- To validate my model, I compute the diffraction pattern of the theoretical protein structure to check if it matches the experimental pattern within a reasonable tollerance

Crio-electron microscopy

- Crio-electron microscopy gives us a diffraction pattern using an electron beam
 - It is really useful for really big complexes that cannot be cristallized
 - Resolution is lower than X-ray diffraction

PDB and other DBs and tools

- The PDB file does not contain the electron density, it is an approximation of the structure
- The resolution of an X-ray diffraction is important
 - 5.0Å resolution is reasonably accurate only for the position of the backbone
 - 1.5Å can be generally trusted, also for drug design
- A ligand in PDB is any molecule co-cristallized with the macromolecule considered
- Signal peptides are 10 to 30 residues in length
 - They are usually cleaved and therefore they do not appear in the protein 3d strucutre
- A PDB file has an unique identifier of 4 letters and numbers
- On PDB I can also find the FASTA file for the protein
 - FASTA contains the covalent structure (i.e the sequence) of the protein
 - FASTA has 60 residues per line
 - It is the sequence derived from the structure, it can be different than the one in uniprot (!)
- Coverage refers to the percentage of protein sequences covered in the protein structure
- PDB files produced using a synchrotron source have 2 spots associated with every atom
- For each ATOM we have the xyz coordinates, the occupancy, and temperature factor
 - Occupancy is how well the atom fits the electron density
 - The temperature factor refers to the mobility of the position
- $\bullet\,$ The PDB file contains the atomic model of a macromolecule
- The CPK colorscheme is a popular set of colors used for the different atoms
- The structure validation window reports the percentile rank of different validation methods
 - Blue is good, red is bad
- DSSP is a program that reads a PDB file and assigns a secondary structure to each PDB coordinate
 - It was made by Sanders, one of the founders of bioinformatics, and Kabsch
- A database can be defined by its statistics

- Data in a DB can be distributed in categories that are relevant for the interpretation of data
- The space group refers to the simmetries of the unit cell
- The Ramachadran plot of a structure can be generated with Procheck (EMBL)
 - It is much more informative than the 3d view generated with Rasmol
- The PDBsum is a pictorial database that provides an at-a-glance overview of the contents of each 3D structure deposited in the Protein Data Bank
 - It is offered by EMBL
 - It shows also the Procheck output for the structure
 - It shows the biological unit instead of the unit cell

PDB101

- A crystal is composed of the unit cell, that is translationally repeated in the crystal
- The unit cell is composed of asymmetric units, that rotated and translated form the unit cell
- The asymmetric unit is the unique part of the crystal structure
- The biological assembly is the biologically relevant form
- Occupancy of an atom is the fraction of times that atom is in the specified position in the crystal
 - The occupancies for an atom always sum to 1, giving the possible alternate conformations
- The R-value is the fit between the theoretical diffraction pattern of the model and the experimental one
 - 0 is a perfect fit, 0.63 is a the fit of a random diffraction pattern
- R-free is another statistic that avoids the bias introduced in the refinement step

Nuclear magnetic resonance

- It does not require the crystal, it is performed in an homogeneous really concentrated protein solution
- The frequencies used are on the radiowave length, so really low energies
- NMR measures contacts among atoms in solution, it measure frequence changes when nuclei come close to each other
- The data produced is a contact map
- The only nuclei considered are mainly O, H, C
- From NMR I get many conformation for each molecule, and if I visualize it in a molecular visualization software I see many superimposed structures
 - The core is usually stable and in agreement with X-ray data, while regions with high temperature factor have many configurations
 - From the many structures I can recover a consensus structure

Time resolved X-ray crystallography

- I use pulsed X-ray instead of a continuous beam
- I can observe the conformational changes of the protein between different pulses

Protein structure allignment

- We wan to compare the structure of 2 proteins
- Hydrogens bond are mainly located at the level of the backbone
- I keep 1 protein fix as a template, and I try to superimpose the other backbone
 - There are many different algorithms to do this
- The first reasonable step would be to compare the position of the backbone of the 2 proteins
- Sequence similarity is meaningful only with an underlying structural similarity
- The reduce representation of a protein contains only the $C-C\alpha-N$ elements of the backbone
- The simplest measure of structural similarity is the root mean squared deviation (RMSD) among atomic coordinates

- One of the algorithm for structural alligment used in the PDB is called java combinatorial extension (JCE)
- JCE was written by Philippe Bourne, the director of the PDB
 - It is one of the best-performing algorithms
- The length of an allignment is the length of the protein sequence, plus the gap introduced
- Sequence identity is a score between 0 and 1 that gives the number of corresponding residues after the allignment
- Sequence similarity is a score between 0 and 1 that gives the number of similar residues after the allignment
- Structure is conserved more than sequence (!)

Protein families

- Multiple structural allignment allow to define protein families
 - We know around 14000 protein families
- They were discovered by M. Dayhoff
- A protein family is the set of proteins that perform the same function in different organism
- Proteins in the same family can also be really different in sequence, but their structure is really similar
- We cannot detect sequence similarity when under 30%, but we can detect structure similarity in those
 cases
 - Under 30% the result is not statistically significant
- A superfamily is a set of protein families with different foldings that can perform the same function

Domains

- A protein domain coincides with the folded protein for small globular proteins (150 aa)
- When the PDB grew, we realised that multi-domain proteins share domains with small globular proteins

Sequence allignment

- It is our only way to compare proteins for which I do not have structures
- Sequence comparison can be pairwise or database search
- Database search is an extension of pairwise sequence alignment
- Sequence allignment can be local or global
- A global allignment optimizes pairing over the whole sequences by introducing gaps
 - A global allignment has a length that is at least as long as the longest sequence
- A local alignment stops the allignment if continuing it makes it worse
- A metric is a set of rules that allow us to score a result
- The Hamming distance metric is, for a pair of sequences equal in length, the number of mismatching positions
 - It is used for ungapped allignments
- The Levensthein or edit distance