3D Data Processing in Structural Biology Assignment 2

(due May 10th)

Write a program that detects an (almost) largest structural alignment between two proteins or two RNA molecules. Input to the program is two structures in the PDB format.

- Consider only Cα atoms for proteins and only phosphate atoms for RNA molecules
- Define transformations using triangles formed by 3 consecutive $C\alpha$ atoms for proteins or phosphate atoms for nucleic acids i.e. try only |pdb1|x|pdb2| transformations.

Name your program structAlign.cc and submit this file in ex2.tar

Input to the program:

structalign epsilon pdb1 pdb2

where epsilon is the largest allowed distance between two corresponding atoms in the alignment

Output:

{Size of the largest alignment} {RMSD between the aligned atoms} {3D transformation (3 rotational and 3 translational parameters)}
For example:

130 1.32 0.5 0.11 0.7 10.1 21.30 -10.1

The transformation transforms pdb2 on pdb1. In addition, your program should create a transformed PDB file 'transformed.pdb' that contains the transformed pdb2. This file should contain **all** atoms (not only $C\alpha$ or P atoms).

To help you with this task we provide you a program alignRand.cc that performs the alignment task be generating random transformations. It superimposes the centers of mass of the two input PDB files and generates random rotations. You can use this program as a basis for your alignment program.

In addition, we included a number of basic classes, that alignRand.cc uses for alignment and of course you can use them too. These classes are a subset of GAMB library: http://bioinfo3d.cs.tau.ac.il/group/GAMB/GAMB.html

Briefly, this is the relevant functionality:

Vector3 – basic 3D vector class for storing and manipulating xyz

coordinates

- RigidTrans3 basic class for 3D rigid transformations that stores 3D rotation (Matrix3 class) and translation (Vector3 class).
- Triangle basic class for defining triangles and transformations.
- Atom stores the information from the PDB ATOM line that includes index, residue type, coordinates, etc.
- Molecule template container for molecules
- PDB enables efficient PDB parsing. Use PDB::Selector to read specific parts of the PDB.
- GeomHash enables efficient hashing. Useful in building correspondence list, when you look for points that are within short distance from a given point
- Match this class implements a correspondence list. The function calculateBestFit computes a transformation that minimizes RMSD for the correspondence list that is stored in the class object.

Sample input files are in the input folder.

Note: you don't have to implement bipartite matching algorithm to find the maximal correspondence in this assignment. Instead, you can rely on the shortest distance heuristics that is implemented by the Match class.

Please submit your code in ex2.tar: structAlign.cc and a file with a table summarizing alignment size and RMSD for each input pair for epsilon = 1.0 and epsilon = 3.0

Good Luck!