

The Hebrew University of Jerusalem

Syllabus

3D DATA PROCESSING IN STRUCTURAL BIOLOGY - 76562

Last update 03-03-2019

HU Credits: 3

<u>Degree/Cycle:</u> 1st degree (Bachelor)

Responsible Department: Computer Science & Computational Biology

Academic year: 0

Semester: 2nd Semester

<u>Teaching Languages:</u> Hebrew

Campus: E. Safra

Course/Module Coordinator: Dina Schneidman

Coordinator Email: dina.schneidman@mail.huji.ac.il

Coordinator Office Hours:

Teaching Staff:

Dr. Dina Schneidman

Course/Module description:

Introductory course to computational structural biology. The course will focus on general methods for processing three-dimensional datasets, such as pattern recognition, surface and volume matching.

We will learn how these algorithms are used in Computational Structural Biology for structural modeling of macromolecules.

Course/Module aims:

Review of methods for processing 3D data with applications in structural modeling of macromolecules.

<u>Learning outcomes - On successful completion of this module, students should be</u> able to:

The course will give the students the tools to read and understand state-of-the-art papers in computational structural biology. In addition the goal is to learn to develop new algorithms for structural modeling of macromolecules, as well as 3D object recognition.

Attendance requirements(%):

100%

Teaching arrangement and method of instruction: Frontal teaching Small groups guidance for final projects

Course/Module Content:

- 1.Introduction to macromolecular structure
- 2. Pattern recognition by point matching, structural alignment of proteins (pairwise and multiple)
- 3. Pattern recognition by surface matching, algorithms for assembly of macromolecular complexes (pairwise and multiple)
- 4. Volumetric matching algorithms, structural modeling with Electron Microscopy (EM) data
- 5.Structural modeling with distance distribution from Small Angle X-ray Scattering (SAXS)experiments
- 6.Structural modeling with distance restraints based on cross-linking mass

spectrometry datasets 7.Integration of multiple datasets 8.Handling uncertainty in the data

Required Reading:

N/A

Additional Reading Material:

Course/Module evaluation:
End of year written/oral examination 0 %
Presentation 0 %
Participation in Tutorials 10 %
Project work 0 %
Assignments 20 %
Reports 0 %
Research project 70 %
Quizzes 0 %
Other 0 %

Additional information:

No biological background is needed. The course is focused on general methods for 3D datasets with applications in Structural Biology.

The grade consists mainly of the research project (70%), 3-4 exercises (20%) and class participation (10%).