BT610 Frontier in Biomolecular Simulation 3-0-0-6

Introduction to computational structural biology; review of protein structure; basic thermodynamics; quantum mechanics; molecular mechanics; computer simulation; free energy calculations; protein structure prediction; protein-ligand interaction; enzyme catalysis; molecular docking: large scale screening of ligands against a target; advanced sampling techniques; molecular modeling softwares.

Texts:

- 1. Andrew Leach, *Molecular Modeling; Principles and Applications*, 2nd Edition Addison-Wesley Publications. 2001
- 2. D. Frenkel and B. Smit, Understanding Molecular Simulation, 2nd Edition, Academic Press, 2001

References:

1. K. A. Dill and S. Bromberg, *Molecular Driving Forces: Statistical Thermodynamics in Chemistry and Biology, Garland Science*, 2003