## **Computer Simulations of Chemical Reactions in Solution and Biomolecules**

MSc. and Ph.D. Course

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## TABLE OF CONTENTS (SYLLABUS)

- 1. **Introduction to Molecular Modeling:** *Historical Developments, Literature*
- 2. **Quantum Mechanics:** *Key Concepts, Methods, and Machinery*
- 3. **Molecular Mechanics:** *Key Concepts, Methods, and Machinery*
- 4. **Statistical Thermodynamics:** Essential Concepts (Partition Functions, Boltzmann Population, Entropy, Enthalpy, Free Energy)
- 5. **Solvation Methods:** Polarized Continuum Methods (PCM), Conductor-Like Screening Model (COSMO, COSMO-RS), Explicit Solvation, PBSA/GBSA Methods
- 6. **QM/MM Methods:** Background, Theory, Applications, and Case Studies (Reaction Mechanisms of Enzymes)
- 7. Thermodynamic Cycles for Computation of a Free-Energy Change in Condensed Phase: Concepts, Applications (Reduction Potentials and Acidity Constants)
- 8. Free Energy Perturbation (Thermodynamic Integration) and PMF Methods:

  \*Concept, Theory, Applications\*
- 9. **Transition State Theory:** Eyring Equation (Theory, Applicability and Limitations, Kinetic Isotope Effects, Tunneling Correction), More Advanced Theories (Variational Transition State Theory)
- 10. **Modelling Chemical Reactions in Solution:** *Theory and Applications*
- 11. **Physical Chemistry of Enzyme Catalysis I:** Reaction Rate and Order, Michaelis-Menten Equation, Metals in Enzymology (Stability Constants, Selectivity, Magnetism, Spectroscopy, Redox and Spin-State Activity)
- 12. **Physical Chemistry of Enzyme Catalysis II:** Theoretical Calculations of Reaction Profiles, Electron Transfer Kinetics and Redox Reactions, Marcus Theory, Adiabatic versus Non-adiabatic reaction dynamics (Landau-Zener model and beyond)
- 13. **Physical Chemistry of Enzyme Catalysis III Metaloenzymes:** Crystal/Ligand Field Theory, DFT vs. WFT theories (Advantages and Pitfalls), Relativistic Effects (on Reactivity).
- 14. **Recent Advances in Molecular Modeling.** *Outlook and Summary.*

Talks by LR

Talks by MS