

**Quiz2 2025**  
**Biophysics(BIOP)**  
**(6. Nov.25)**

**Total Marks:20**

**Time: 45 min**

**Section-A (MCQ 1\*10 Marks)**

**1. During NVT equilibration, which quantity is kept constant?**

- A. Number of particles, volume, and temperature
- B. Number of particles, pressure, and temperature
- C. Volume, pressure, and energy
- D. Temperature, pressure, and density

**2. Which of the following thermostats is *commonly used* in GROMACS?**

- A. Parrinello–Rahman
- B. Berendsen
- C. Nose–Hoover
- D. Both (b) and (c)

**3. Which command is used to compute RMSD in GROMACS?**

- A. gmx rms
- B. gmx gyrate
- C. gmx rmsd
- D. gmx energy

**4. The success of homology modelling relies primarily on the principle that:**

- A. Protein structure is more conserved than protein sequence during evolution
- B. Protein sequence is more conserved than structure during evolution
- C. Sequence and structure evolve independently
- D. Sequence similarity implies identical function

**5. Protein folding leads to a decrease in conformational entropy because:**

- A. The protein adopts many flexible conformations
- B. The number of accessible microstates decreases
- C. The solvent entropy decreases
- D. The energy of the system increases

**6. During protein folding, which thermodynamic quantity primarily drives the process despite an apparent decrease in conformational entropy of the protein chain?**

- A. Increase in enthalpy due to hydrogen bonds
- B. Decrease in Gibbs free energy due to hydrophobic effect
- C. Increase in conformational entropy of backbone
- D. Increase in heat capacity of the solvent

**7. Increasing temperature affects the Boltzmann distribution for protein conformations by:**

- A. Sharpening the distribution
- B. Flattening the distribution
- C. Increasing energy differences
- D. Making all conformations equally probable

**8. In molecular dynamics simulations, the Helmholtz free energy is more relevant than Gibbs free energy because:**

- A. Simulations are typically carried out at constant pressure

- B. Volume fluctuations are negligible  
 C. Solvent compressibility is infinite  
 D. The system is adiabatic

9. Mixing two different amino acids in water increases entropy because:

- A. The amino acids form more covalent bonds  
 B. The number of accessible microstates increases  
 C. Their mass changes  
 D. Hydrogen bonding is eliminated

10. When a charged side chain moves from solvent to protein core, the free energy change is usually:

- A. Favorable ( $\Delta G < 0$ )  
 B. Unfavorable ( $\Delta G > 0$ )  
 C. Zero  
 D. Undefined

**Short Answer Type (2 marks each)**

1. A protein in solution can adopt many unfolded conformations but only a few folded states. Explain how entropy contributes to protein folding being thermodynamically unfavorable at high temperatures.
2. What is the difference between the SPC and TIP3P water models used in GROMACS?
3. How does the Verlet cutoff scheme improve performance in GROMACS?
4. Two protein conformations differ in energy by 5 kJ mol<sup>-1</sup>. Which conformation is more populated at 300 K? Explain without calculation.
5. You are building a homology model for a newly sequenced protein (~310 amino acids). You find the following possible template structures:

Template	Sequence Identity	Coverage (Aligned Length)	Resolution (Å)	Organism Similarity	Known Function Match
A	28%	98%	1.8 Å	Same organism	Function unknown
B	42%	85%	2.5 Å	Different organism	Known same enzymatic function
C	55%	60%	3.2 Å	Same organism	Known same biological pathway
D	33%	100%	1.2 Å	Distant organism	Active site mutated (not conserved)

Which template would you select for homology modeling and why?