

# BIOSC 1540 - Computational Biology

## Quiz 04

Apr 8, 2025

20 points

Please read the following instructions carefully before beginning your assessment.

- **Time limit:** You have 15 minutes to complete and turn in this assessment.
- **Closed note:** You may not use any notes or additional resources during this assessment.
- **No digital devices:** The use of digital devices, including calculators, is not allowed.

I agree to follow the above instructions. I affirm that all work on this assessment will be my own and that I will not give or receive any unauthorized assistance. To have your assessment graded, you must write your name, sign, and provide your student ID below.

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Name

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Signature

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Student ID

**Bored area:** Draw a scene from your favorite movie, design/name a new animal, or write a micro story from the point of view of your pencil/pen during this quiz.

### Problem 1

In a Ramachandran plot, which of the following best explains why certain regions are “forbidden”?  
(2 points)

- ☐ (A) They correspond to bond lengths that are too short.
- ☐ (B) They would result in steric clashes between atoms.
- ☐ (C) They require impossible electron configurations.
- ☐ (D) They only occur in denatured proteins.

### Problem 2

What interaction is generally considered the primary driving force for the folding of globular proteins?  
(3 points)

- ☐ (A) Disulfide bonds
- ☐ (B) Hydrogen bonds
- ☐ (C) Hydrophobic interactions
- ☐ (D) Ionic (salt bridge) interactions

### Problem 3

In X-ray crystallography, the smallest distance between two points that can be resolved in a protein structure is referred to as the \_\_\_\_\_, which is often measured in units of \_\_\_\_\_.  
(1 point)

### Problem 4

The main source of 3D structures of biomolecules is called the \_\_\_\_\_.  
Hint: It does not start with a “Q”.  
(3 points)

### Problem 5

In the context of binding thermodynamics, which aspect is most likely to be underestimated when relying solely on a single, static structure?  
(2 points)

- ☐ (A) The entropy of the system.
- ☐ (B) The precise orientation of hydrogen bonds.
- ☐ (C) The strength of electrostatic interactions.
- ☐ (D) The direct measurement of binding enthalpy.

### Problem 6

In grid-based protein-ligand binding models, if the number of available binding sites ( $N$ ) increases while the number of ligands ( $L$ ) remains constant, how does this affect the system's entropy?

(1 point)

- Ⓐ The entropy decreases.
- Ⓑ The entropy increases.
- Ⓒ The entropy remains unchanged.
- Ⓓ The entropy first decreases then increases.

### Problem 7

Why is it important for the target and template proteins to share a high level of sequence identity?

(2 points)

- Ⓐ It leads to consistent secondary structure elements.
- Ⓑ It speeds up the energy optimization steps.
- Ⓒ It increases the chance that both proteins undergo the same post-translational processing.
- Ⓓ It helps generate a more accurate alignment.

### Problem 8

Profile-based alignment methods improve the sensitivity of homology detection by:

(1 point)

- Ⓐ Use insertions and deletions to emphasize conserved regions.
- Ⓑ Speeding up the alignment process at the cost of precision in selecting homologs.
- Ⓒ Bypassing the requirement for generating multiple sequence alignments.
- Ⓓ Modeling each position in a sequence as a distribution of potential amino acids.

### Problem 9

Deep learning approaches, such as AlphaFold, derive critical information on residue-residue contacts from \_\_\_\_\_.

(2 points)

### Problem 10

Describe two potential reasons why a ligand might have a good docking score (e.g.,  $\Delta G$  of -30 kcal/mol) but not be a successful inhibitor in real-world applications.

(1 point)

### Problem 11

Which of the following best describes the role of stochastic algorithms in pose optimization during molecular docking?

(1 point)

- ☐ (A) They explore the energy landscape by making random modifications to molecular poses and accepting changes based on probability criteria.
- ☐ (B) They use adaptive energy barriers to guide the sampling of molecular shapes, focusing on maintaining proper transition probabilities.
- ☐ (C) They group similar molecular shapes by analyzing structural differences to identify key representative conformations.
- ☐ (D) They apply evolutionary techniques that iteratively select and improve binding poses using scoring functions.

### Problem 12

Which statement best describes the role of traditional scoring functions in molecular docking?

(1 point)

- ☐ (A) They break down binding interactions into local atomic potentials calibrated with experimental and quantum data.
- ☐ (B) They use machine learning models trained on structural data to predict binding strengths based on geometric and chemical features.
- ☐ (C) They estimate binding energies using physical forces, statistical data, and solvation effects to rank docking poses.
- ☐ (D) They assess how well proteins and ligands fit together by combining multiple scoring methods.