

## Daffodil International University

Faculty of Science & Information Technology Department of Computer Science & Engineering Final Semester Examination, Spring 2025

Course Code: CSE115, Course Title: Introduction to Biology and Chemistry for Computation

Level: 01 Term: 01 Batch: 68

Time: 2:00 Hrs Marks: 40

## **Answer ALL Questions**

[The figures in the right margin indicate the full marks and corresponding course outcomes. All portions of each question must be answered sequentially.]

1.	pre Mo tear	research team is developing a new anti-cancer drug that targets DNA to block the replicant neer cells. The drug binds directly to DNA's nucleotide bases, interfering with its structure eventing the cancer cells from multiplying. The team is considering two computational met blecular Mechanics (MM) and Quantum Mechanics (QM) for the drug-DNA interaction. Since maims to understand electronic interactions at the binding site, they must choose the propriate approach.	s nucleotide bases, interfering with its structure and The team is considering two computational methods: chanics (QM) for the drug-DNA interaction. Since the ms at the binding site, they must choose the most thod for analyzing the binding mechanism four answer by comparing both methods, all energy in a molecular system. Describe attion with a relevant example. In the Global Antibiotic Resistance Project, is gorithm for detecting mutations linked to antibiotic surrows-Wheeler Transform (BWT) and suffix arrayint of sequencing data for efficient DNA sequence attion, they are given the Burrows-Wheeler Transform the geding with read mapping, they must reconstruct the res computational efficiency. It the original genome sequence from the p explanation of the process.  [BWM) from the original sequence and	CO1
,	aj		5+5]	
	b)	the key components involved in the calculation with a relevant example.		-
2.	dev resi bas alig (BV	Emily Roberts, a leading genomics scientist at the Global Antibiotic Resistance Projectiveloping an efficient genome alignment algorithm for detecting mutations linked to antibistance in bacteria. Her team employs the Burrows-Wheeler Transform (BWT) and suffix a sed read mapping to handle the vast amount of sequencing data for efficient DNA sequencement. As part of their computational validation, they are given the Burrows-Wheeler Transform (BWT) sequence: "ATTCG\$GG". Before proceeding with read mapping, they must reconstructional genome and analyze how BWT improves computational efficiency.	escribe  iistance Project, is inked to antibiotic i) and suffix array- nt DNA sequence Wheeler Transform nust reconstruct the  com the [5+5] ce and BWM  to understand how quence of the virus  CO3	
	a) b)	given BWT. Provide a detailed step-by-step explanation of the process.  Generate the Burrows-Wheeler Matrix (BWM) from the original sequence and compute the corresponding Suffix Array (SA(T)). Illustrate the efficiency of BWM		CO2
3.	mu stra from corn Sec	Rachel is studying the genetic variation in several strains of the Influenza virus to understand trations affect its virulence and transmission. He needs to align the entire RNA sequence of the ains to understand the differences and similarities between them. He will compare two sequence and different strains and use an alignment strategy that ensures all parts of both sequence ansidered. Given Sequences:  quence 1 (Virus Strain A): AGTACGGA quence 2 (Virus Strain B): AAGTAGGA	virulence and transmission. He needs to align the entire RNA sequence of the virus d the differences and similarities between them. He will compare two sequences and use an alignment strategy that ensures all parts of both sequences are sequences:  Strain A): AGTACGGA  Strain B): AAGTAGGA	
	gs bs	constructing the alignment score matrix.  Compute the optimal alignment and alignment score to find the best alignment between the two sequences over their entire length.	+21	
4.	Ms	st. Zannatun Nesa, a bioinformatics student, is studying the genetic variations between two sp I wants to identify the similarities and differences between their DNA sequences. She is usin	pecies ng the	

FASTA algorithm to compare the sequences by applying hash tables to efficiently match subsequences. She also needs to evaluate the performance of the FASTA algorithm in terms of its sensitivity and selectivity to ensure that the algorithm is accurately detecting significant sequence matches.			
	ery Sequence: ACGGTAGCTA		
1 ar	get Sequence: TCACGGTCT		
(a)	<b>Apply</b> the FASTA algorithm with K=1 for both the query sequence and the target sequence to find the best matching subsequences.	[6+2+2]	CO3
by	Calculate sensitivity and selectivity specifically for the base G (Guanine) considering the Query sequence as the dataset and the Target sequence as the outcome.		
1	<b>Interpret</b> the results based on the sensitivity and selectivity value obtained and assess how well the FASTA algorithm performs in finding the relevant sequence matches between the two sequences.		