PARUL UNIVERSITY

PARUL INSTITUTE OF APPLIED SCIENCES

MID SEMESTER INTERNAL EXAMINATION, SUMMER 2019-20

B.Sc. Microbiology Semester IV

Date: 04/03/2020

Time: 1hr 30min

Paper Name: Bioinformatics

Paper Code: 11102252

Max. Ma Instructi			
	uestions are compulsory and options are	given in first and second question	only
-	bers to the right of question indicate the		. 0111)
Q. 1	Attempt any one question of the follow (i) Explain the classification of biolog Sequence databases, Structure datab with examples. Explain about Primar examples. (ii) Explain in detail Dot Matrix algorithm.	gical databases in detail. Explain ases, and Functional databases y and Secondary databases with	(08)
Q. 2	Attempt any three questions of the following (i) What is SwissProt database? (ii) Short note on PlasmoDB. (iii) Short note on Smith Waterman Algorithm (iv) Difference between global and log (v) What is pairwise alignment and mutations.	gorithm al alignment?	(12)
Q. 3	Do as directed. Attempt all five questions. (i) Give four examples of tools of Multiple Sequence Alignment. (ii) What is sequence alignment? (iii) What are the three main steps in dynamic programming? (iv) The Protein Information Resource (PIR) is established in 1984 by which organization? (v) HOVERGEN is a specialized database for which types of genes?		
Q. 4	Write correct option in your answer sheet for following 15 multiple choice questions. (15)		(15)
MCQ 1	GenBank, the nucleic acid sequence d (A) Brookhaven Laboratory (C) European Molecular Biology Laboratory (EMBL)	atabase is maintained by (B) DNA Database of Japan (D) National Centre for Bio Information (NCBI)	,
MCQ 2	The initiation of FASTA format has sy (A) > (C) /	ymbol? (B) < (D) *	

MCQ 3	Which of the following is not a variant of BLAST?			
	(A) BLASTN	(B) BLASTP		
	(C) BLASTX	(D) TBLASTNX		
MCQ 4	Which of the following is the sequence is a sequence alignment tool provided by NCBI?			
	(A) Chime	(B) BLAST		
	(C) FASTA	(D) Clustal W		
MCQ 5	The procedure of aligning two sequences by searching for patterns that is in the same order in the sequences			
	(A) sequence alignment	(B) pair-wise alignment		
	(C) multiple sequence alignment	(D) all of these		
MCQ 6	The alignment procedure that tries to align the entire sequence is			
	(A) multiple sequence alignment	(B) pair-wise alignment		
	(C) global alignment	(D) local alignment		
MCQ 7	Which is the default scoring matrix used in BLAST?			
	(A) PAM 62	(B) BLOSUM 62		
	(C) BLOSUM 45	(D) BLOSUM 80		
MCQ 8	Literature databases include			
	(A) MEDLINE and PubMED	(B) MEDLINE and PDB		
	(C) PubMED and PDB	(D) MEDLINE and PDS		
MCQ 9	If you want to align two sequences which are more identical in sequence, which of the			
	following scoring matrices would be most appropriate			
	(A) BLOSSUM 35	(B) BLOSSUM 40		
	(C) BLOSSUM 62	(D) BLOSSUM 80		
MCQ 10	Among the following which one is not the approach to the local alignment?			
	(A) Smith-Waterman algorithm	(B) K-tuple method		
	(C) Words method	(D) Needleman-Wunsch algorithm		
MCQ 11	Which of the following statements is false when describing SWISS-PROT?			
	(A) It is a curated protein sequence database	(B) Data is redundant		
	(C) Provides a high level of	(D) It is maintained by Swiss Institute		
	annotations	of Bioinformatics and EBI.		
MCQ 12	Which alignment is used to predict whether two sequences are homologous or not?			
	(A) Local	(B) Global		
	(C) Pair-wise	(D) Multiple		
MCQ 13	The scoring matrix for protein sequences is			
	(A) PAM matrix	(B) BLOSSUM matrix		
	(C) RNA sequences	(D) Thure Etzold		

MCQ 14	Which of the following is not a software for dot plot analysis?		
	(A) SIMMI	(B) DOTLET	
	(C) DOTMATCHER	(D) LALIGN	
MCQ 15	For palindromic sequences, what is the structure of the dot plot?		
	(A) 2 intersecting diagonal lines at the midpoint	(B) One diagonal	
	(C) Two parallel diagonals	(D) No diagonal	

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