Roll No. Total No. of Pages :02

Total No. of Questions: 09

B.Tech.(BT) (2012 to 2017) (Sem.-5)
BIOINFORMATICS

Subject Code: BTBT-505 M.Code: 70506

Time: 3 Hrs. Max. Marks: 60

INSTRUCTIONS TO CANDIDATES:

- SECTION-A is COMPULSORY consisting of TEN questions carrying TWO marks each.
- 2. SECTION-B contains FIVE questions carrying FIVE marks each and students have to attempt any FOUR questions.
- 3. SECTION-C contains THREE questions carrying TEN marks each and students have to attempt any TWO questions.

SECTION-A

1. Answer briefly:

- a. Sequence similarity vs. sequence identity
- b. EMBL
- c. Global alignment
- d. Affine gap penalties
- e. BIT score
- f. Rasmol
- g. COGS
- h. Bootstrap
- i. Entrez
- j. Phylogram

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SECTION-B

- 2. Write about the CATH database.
- 3. Describe the GOR method of protein secondary structure prediction.
- 4. Explain Smith-Waterman algorithm.
- 5. Describe any one multiple sequence alignment tool.
- 6. Differentiate between PAM and BLOSUM scoring matrices.

SECTION-C

- 7. Blast is used for database searching. Describe the BLAST algorithm and BLAST output.
- 8. Describe different bioinformatics methods of gene prediction. How prokaryotic gene prediction tool is different from eukaryotic gene prediction tool?
- 9. Write the application of phylogenetic tree constructions. Construct the tree based on given matrix using UPGMA method.

	A	В	С	D
A	0			
В	7	0		
C	6	8	0	
D	11	13	10	0

NOTE: Disclosure of Identity by writing Mobile No. or Making of passing request on any page of Answer Sheet will lead to UMC against the Student.

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