Total number of printed pages – 4 CPBT 8309 / PEBT 8301

Sixth Semester Examination - 2008

## BIOINFORMATICS

Full Marks – 70

## Time: 3 Hours

Answer Question No. 1 which is compulsory and any five from the rest. The figures in the right-hand margin indicate marks.

- Answer the following questions : 1.
  - What is PSI-BLAST ? Why it is advan-(a) tageous over BLAST ?
  - (b) Expand EMBL. Which are the nucleotide databank of EMBL ?
  - (c) Define gap penalty. What is its significance in the alignment of sequences ?

P.T.O.

 $2 \times 10$ 

B. Tech

- (d) What is threading ? How threading play significant role in protein structure prediction?
- Differentiate between BLOCKS and PRO-(e) FILES.
- (f) What is PAM matrix ? Show an out put for PAM Matrix.
- Differentiate between Lattice model and (g) Continuous model of the polymer simulation.
- Which one is the potential database for (h) metabolic and regulatory pathways?
- What basic tools of CAMD (Computer (i) aided molecular design) you can adopt for ligand designing of protein ?
- Name three primary nucleotide sequence (i) repositories along with their locations and functions.
- What is an 'E-value'? You do a databank 2. (a) search using FASTA with an aminoacid
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sequence as query. The only reported match has an E-value of 10. What does this mean for the similarity and homology of the sequences ? 5

- (b) What are the major extensions of BLAST?Discuss the algorithms used and applications of these programmes.
- What are the sequence and structure databank of protein ? Briefly explain the PIR-PSD and SWIS-PROT database and their application in proteomics. 4+3+3
- 4. (a) What is sequence alignment ? Differentiate between the algorithms used for local and global alignment study.
  - (b) Calculate the dynamic programming matrix and optical alignment for two DNA sequence 'GAATTC' and 'GATTA', scoring '+2' for matching, '-1' for mismatch and '2' for linear gap penalty.
- 5. What is *In Sillico* secondary structure prediction of protein ? Discuss methods including

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logic neural networking and Chao-Fasman algorithms for protein secondary structure prediction. 2+4+4

- 6. Differentiate between the following : 5×2
  - (a) EST and SNP
  - (b) SCOP and CATH.
- 7. Write short notes on any *two* of the following : 5×2
  - (a) Hidden Markov Model
  - (b) Dynamic Programming using distance matrix
  - (c) Molecular dynamics in drug designing
- Define database and data type in bioinformatics. What are the different data types and databases are used for gene structure and gene product function prediction ? Show the out put format of nucleotide database in NCBI using ENTREZ search. 2+5+3

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