

(University of Choice)

## MASINDE MULIRO UNIVERSITY OF SCIENCE AND TECHNOLOGY (MMUST)

#### MAIN CAMPUS

### UNIVERSITY EXAMINATIONS 2018/2019 ACADEMIC YEAR

#### FOURTH YEAR SECOND SEMESTER EXAMINATIONS

# FOR THE DEGREE OF BACHELOR OF MEDICAL BIOTECHNOLOGY /LABORATORY SCIENCES (DIRECT ENTRY AND UPGRADING)

#### MAIN EXAMINATION

COURSE CODE: BML 421

COURSE TITLE: BIO-INFORMATICS AND BIO-COMPUTING

**DATE: 27<sup>TH</sup> MAY 2019 TIME: 8.00** -10.00 AM

#### **INSTRUCTIONS TO CANDIDATES**

This paper is divided into three sections, **A B** and **C**, carrying respectively: Multiple Choice questions (**MCQs**), short answer questions (**SAQs**) and long answer questions (**LAQs**).

TIME: 2 Hours

MMUST observes ZERO tolerance to examination cheating

BML 421: Bioinformatics and Biocomputing

#### SECTION A: MULTIPLE CHOICE QUESTIONS (MCQs)

- 1. Bioinformatics and bio-computation field involve the following EXCEPT?
  - a) The discovery of therapeutic products
  - b) Biotechnology and recombinant technology
  - c) The study of molecular pathways and drug resistance
  - d) Improving the genetic code
- 2. Core genome comprises which of the following?
  - a) Set of unique genes that allow speciation and are not conserved across same genus.
  - b) Set of unique genes shared by all genomes that are conserved across all species in a genus.
  - c) A set of non-essential genes in an organism
  - d) The genome that is found in all organisms
- 3. Which of the following best describes the pan-genome of an organism?
  - a) Set of genes forming up a single strain
  - b) Essential genes in a specie
  - c) Complete set of all complement genes present in all strains of a specie
  - d) All sets of genes that are comprised in a genus.
- 4. Which of the following is NOT a functional macromolecule in cellular processes?
  - a) mRNA
  - b) SiRNA
  - c) tRNA
  - d) Ribosomal RNA
- 5. Which of the following is a BLAST program for a protein sequence in translated nucleotide database?
  - a) tBLASTx
  - b) BLASTn
  - c) BLASTx
  - d) tBLASTn
- 6. What does OMIM represent in bioinformatics?
  - a) Multiple sequence alignment tool
  - b) Organization of mammalian genetic trees
  - c) Database for Mendelian inheritance in man
  - d) Objects for microorganism inheritance
- 7. PDB as applied in bioinformatics is a
  - a) Primer design program
  - b) Data bank for proteins and their structures
  - c) Phylogenetic database
  - d) Phosphorylating dimer of biomolecules

- 8. Orthologous genes
  - a) Similar genes in the same genus that result from gene duplication.
  - b) Genes that have lost function over time in ancestry.
  - c) Similar genes in different genus that results from gene ancestral inheritance.
  - d) Acquired genes from other organisms that were not part of the genus ancestry.
- 9. Which of the following best describes a core genome?
  - a) Set of unique genes that allow speciation and are not conserved across same genus.
  - b) Set of unique genes that are conserved across the genus.
  - c) A set of non-essential genes in an organism
  - d) The genome that is found in all organisms
- 10. Which of the following is a BLAST program for a protein sequence in a protein database?
  - a) BLASTp
  - b) BLASTn
  - c) BLASTx
  - d) tBLASTn
- 11. Which of the following is a BLAST program for a nucleotide sequence in a nucleotide database?
  - a) BLASTp
  - b) BLASTn
  - c) BLASTx
  - d) TBLASTn
- 12. Which of the following describes paralogous genes?
  - a) Similar genes in the same genus that result from gene duplication.
  - b) Similar genes in different genus that results from gene ancestral inheritance.
  - c) Genes that have lost function over time in ancestry.
  - d) Acquired genes from other organisms that were not part of the genus ancestry.
- 13. Clustal W is a suitable software for computational analysis of
  - a) Pairwise sequence alignment
  - b) Multiple sequence alignment
  - c) Gene expression profiles
  - d) Protein folding into tertiary structures

#### 14. EXPASY is

- a) Database for expression profiles of genes
- b) Database for gene transcripts
- c) Computational Software for protein analysis

- d) Bank for expression regulatory molecules
- 15. Which of the following is a BLAST program for translated nucleotide sequence in a translated nucleotide database?
  - a) tBLASTx
  - b) BLASTn
  - c) BLASTx
  - d) tBLASTn
- 16. Dispensable genome
  - a) Set of unique genes that are conserved across the genus.
  - b) A set of non-essential genes in an organism
  - c) The genome that is found in all organisms
  - d) Set of unique genes that allow speciation and are not conserved across same genus.
- 17.A bioinformatician would like to study expressed genes and their abundance. Where can he find this information
  - a) ENTREZ
  - b) UNI GENE
  - c) BLAST
  - d) Protein data bank
- 18. During computational translation the software will stop the peptide sequence translation when it encounters the following EXCEPT?
  - a) AUG
  - b) UGA
  - c) UAG
  - d) UAA
- 19. Which of the following is FALSE about RNA splicing?
  - a) It is a process through which bacteria produce mature mRNA
  - b) It involves removal of non-coding regions of RNA
  - c) It is a post-transcription process
  - d) RNA editing process for eukaryotes
- 20. Which of the following is a BLAST program for a translated nucleotide sequence in protein database?
  - a) BLASTp
  - b) BLASTn
  - c) BLASTx
  - d) TBLASTn

#### SECTION B: SHORT ANSWER QUESTIONS (40 MARKSO

- 1. Giving relevant examples, describe the use of databases in bioinformatics. (8 Marks).
- 2. List types of bioinformatics databases giving examples of the databases in each case. (8 Marks).
- 3. In the EMBL database format, what do the following lines represent; ID, AC, NI, DE, KW, DT, OS, OC, RN, FT, SQ, CC, RP, RX, RA, RL. (8 Marks).
- 4. Describe the tasks that the ExPASy database can enable you perform. (8 Marks).
- 5. Describe the stepwise process of performing a blast of a DNA sequence. (8 Marks).

#### SECTION C: LONG ANSWER QUESTIONS (40 Marks)

- 1. Discuss the some of the common algorithms used in sequence analysis. (20 Marks).
- 2. Compare and contrast BLAST and FastA platforms during sequence homology searches. (20 Marks).