



INDIAN INSTITUTE OF TECHNOLOGY KHARAGPUR

End-Autumn Semester Examination 2022-23

Date of Examination: 16/11/2022

Session (FN/AN): AN

Duration: 3 hrs.

Full Marks: 80

Subject No.: ES60011

Subject: Application of Machine Learning in Biological Systems

Department/Center/School: School of Energy Science and Engineering

Specific charts, graph paper, log book etc., required

Special Instructions (if any): (1) Answer all questions, (2) In case of reasonable doubt make practical assumptions and write that on your answer script, (3) The parts of each question must be answered together, (4) Calculator is allowed

1. Mention whether the following statements are true or false and justify your answer with a proper explanation.

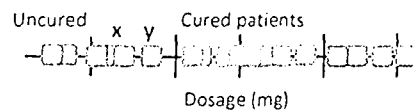
- (a) The threshold value does not allow misclassification for soft margin classifiers in the Support vector machine
- (b) Backpropagation learning is based on gradient descent along the error surface
- (c) Random forest algorithm uses learning Rate as one of its hyper-parameter.
- (d) RPKM normalization method is better than TPM normalization to analyze the gene expression data.
- (e) RNA-seq technique examines the expression of all the genes in different tissues..
- (f) The False Discovery Rate (FDR) can control the number of false positives. [2+2+2+2+2+2= 12]

2. Calculate Quality Score and probability of Error for following bases during sequencing.

- (a) If the probabilities of base call error are 0.2, 0.015, 0.23 and 0.045, then calculate the respective quality score.
- (b) If the quality score is 15, 20, 26 and 34, calculate the probability of base call error. [3+3=6]

3. (a) What is a polynomial kernel? Let's consider a and b represent the observed number of cured and uncured patients with respect to the drug dosage in milligrams. If $a = 3$, $b = 11$, and the polynomial coefficient is $\frac{1}{2}$, calculate the 2-dimensional relationship between the observation 'a' and 'b' using a polynomial kernel.

(b) Suppose we have two dosage measurements of a drug for heart disease i.e., x and y. If $x = 5$ and $y = 6$, what will be the influence value of x and y on each other? Consider the value $\gamma = 2$ for scaling the influence with radial kernel. [3+3=6]



4. (a) What is Convolutional Neural Network (CNN)? Mention the usage of CNN

(b) Explain different layers in Convolutional Neural Network (CNN)

(c) Consider an input image file that has been converted into a matrix of size 12×12 along with a filter of size 3×3 with a Stride of 1. Determine the size of the convoluted matrix.

(d) When we use "valid padding" CNN? Considering an input matrix with 10×10 dimensions and a (2×2) filter, what will be the dimensions of the output matrix after convolution using valid padding? [2+2+2+2=8]

5. (a) Write the BTW transformation for the sequence: ATCCGATC

(b) Calculate the False Discovery Rate (FDR) for the following data set:

False positive = 9 and True positive = 65

(c) Calculate Familywise Error Rate (FWER) for the following data:

alpha level for an individual test = 14%; Number of comparisons = 19

[2+2+2=6]

6. Below table represents the drug effectiveness corresponding to the provided dosage

(a) Considering the threshold of the dosage < 3, calculate the squared residuals (SSR) of the dosage

(b) Calculate the optimal threshold value of the dosage for considering it as the root node in the regression tree.

[2+6=8]

Dosage	Drug effectiveness
2	0
8	12
10	25
13	35
16	98
17	100

7. Consider a data set with 9 mice where 5 are obese and 4 are not obese. We have calculated the log(odds) of obesity for each candidate data by fitting line X.

The log (odds) of each candidate data point for line X is as follows:

log(odds) of a = +.3, log(odds) of b = +1.2, log(odds) of c = +2, log(odds) of d = +2.9

log(odds) of e = +3.8, log(odds) of f = -1.8, log(odds) of g = -1.2, log(odds) of h = -0.7, log(odds) of i = -0.1

Next, we rotate the line (Y) and calculate the log(odds of obesity) for all the candidate data.

The log (odds) of each candidate data point for line Y is as follows:

log(odds) of a = +0.2, log(odds) of b = +0.5, log(odds) of c = +0.8, log(odds) of d = +1.1

log(odds) of e = +1.6, log(odds) of f = -1.1, log(odds) of g = -0.9, log(odds) of h = -0.5, log(odds) of i = -0.2

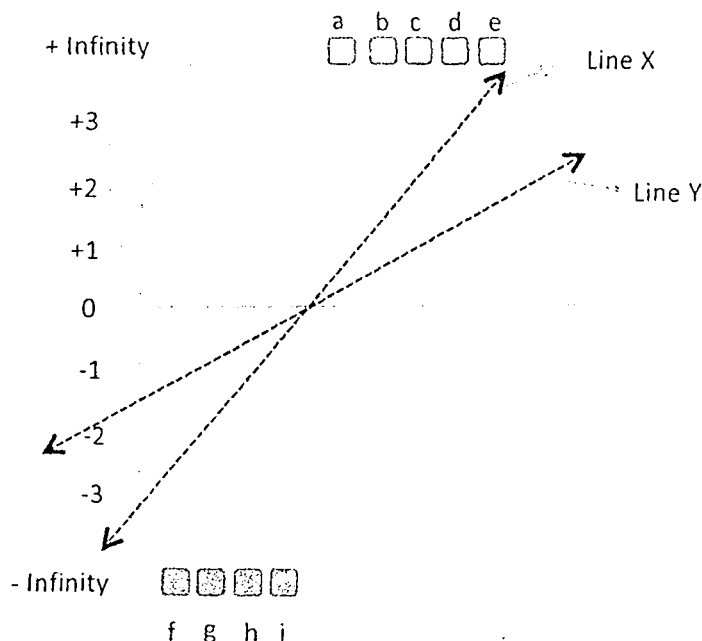
a) What will be the average probability of the mice being obese

b) Calculate the likelihood of the obese mice when fitting line X

c) Calculate the log(likelihood) of all the given data points when fitting line Y

d) Which line can be considered the best fitting line for the above scenario and why?

[2+2+2+4=10]



8. Here we have provided a dataset of 6 patients with colon cancer and its associated symptoms as observed in these patients.

(a) Calculate the total Gini impurity value of **Fatigue** symptoms for separating the patients with or without colon cancer.

(b) Which symptom can be considered the root node for constructing the classification tree and why?

Symptoms			Disease
Diarrhea	Loss of Appetite	Fatigue	Colon Cancer
Yes	No	No	No
No	Yes	Yes	Yes
Yes	Yes	Yes	Yes
No	No	Yes	No
Yes	No	Yes	Yes
No	No	No	Yes

[3+5=8]

9. Suppose we have an artificial neural network having three inputs with corresponding weights, i.e., $w_1 = 2$; $w_2 = -4$; and $w_3 = 1$.

The step function gives the activation of the unit: $\phi(v) = 1$ if $v \geq 0$ otherwise, 0

Consider each of the following input patterns and answer the following questions:

Pattern	P ₁	P ₂	P ₃	P ₄
X ₁	1	0	1	1
X ₂	0	1	0	1
X ₃	0	1	1	1

(a) Calculate the weighted sum for each pattern.

(b) Calculate the output value y of the given neural network

(c) What is the role of the activation functions in Neural Networks?

[4+2+2=8]

10. (a) Normalize the sequence read count genomic data using RPKM normalization.

Gene Name	Gene Lengths	Sample_1	Sample_2	Sample_3
Gene-1	30 kb	3216	1560	1250
Gene-2	14 kb	1452	960	852
Gene-3	18 kb	745	486	856
Gene-4	24 kb	1258	1469	751
Gene-5	28 kb	2548	2984	1589
Gene-6	19 kb	879	1546	1852
Gene-7	6 kb	450	145	94
Gene-8	25 kb	2546	2879	1548
Gene-9	9 kb	198	247	254
Gene-10	4 kb	56	98	105

(b) Normalize the sequence read count genomic data for four samples using 1 PM normalization.

Gene Name	Gene Lengths	Sample_1	Sample_2	Sample_3
Gene-1	40 kb	2100	1240	952
Gene-2	25 kb	1254	540	643
Gene-3	16 kb	950	964	785
Gene-4	24 kb	1120	1840	647
Gene-5	9 kb	582	498	399
Gene-6	7 kb	325	256	478
Gene-7	5 kb	100	350	296
Gene-8	10 kb	507	800	777
Gene-9	4 kb	150	156	198
Gene-10	9 kb	165	289	335

[4+4=8]