EECS545 - Homework 6

March 27th, 2018

Instructions

- This homework is Due Wednesday, April the 11th at 5pm. No late submissions will be accepted.
- As always, submit a write-up and your code for this homework.
- You are expected to use Python for the programming questions in this homework. Use of Python 3 is recommended.
- Submit your write-up to Gradescope under the assignment titled Homework 5. Make sure to tag the pages with the corresponding problems numbers.
- Submit all your python code to Canvas in a compressed zip file named uniqname_hw5_code.zip. Your zip file should contain prob4.py.
- 1. Suppose we have binary states (labeled A and B) and binary observations (labeled 0 and 1) and the initial, transition, and emission probabilities as in the given table. Please answer

State	$P(S_1)$
A	0.80
В	0.20

(a)	Initial	Pro	bs.
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S_1	S_2	$P(S_2 S_1)$
A	A	0.80
A	В	0.20
В	A	0.30
В	В	0.70

(b) Transition Probs.

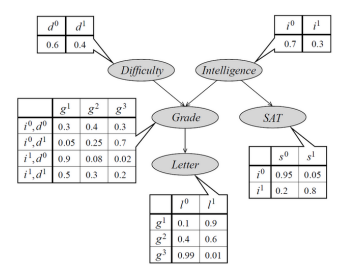
S	O	P(O S)
A	0	0.80
A	1	0.20
В	0	0.10
В	1	0.90

(c) Emission Probs.

the questions below. You need do the calculation by hand instead of using any programs or packages.

- (a) Using the forward algorithm, compute and report the probability that we observe the sequence $O_1O_2O_3 = 010$ and $O_1O_2O_3 = 101$. Briefly justify/explain your answer.
- (b) Using the backward algorithm, compute and report the probability that we observe the sequence $O_1O_2O_3 = 010$ and $O_1O_2O_3 = 101$. Briefly justify/explain your answer.
- (c) Use the Viterbi algorithm to compute and report the most likely sequence of states for $O_1O_2O_3=010$. Show your derivation.
- 2. In this problem, you need to implement in code the forward-backward and Viterbi algorithms, and learn a HMM model with K hidden states from several simple DNA sequences. A DNA sequence is regarded as a series of components from A, C, G, T. Assume the initial state is uniformly-randomly picked from all hidden states. We got two DNA sequences $X_1 = CCTACACGCA$ and $X_2 = CTACGCAAT$, calculate and report:

- the transition probability and emission probability using the forward-backward algorithm for K=2 and K=4.
- the most likely sequence of states of length 4 when the first observation is A, for the two models you have learned.
- 3. Consider the following Bayesian student network. The network models the relationship between the following random variables. The grade received by a student (G) is influenced by the difficulty of the course (D) and the intelligence of the student (I). Our student asks the professor for a recommendation letter. Due to the large class size the professor does not remember the names of students, and hence writes the letter based on the student's course grade. L represents the quality of the recommendation letter. The network also shows how a student's SAT score is influenced by his/her intelligence.



- (a) Write an expression for the joint distribution P(D, I, G, L, S) exploiting the conditional independence relationships in the network.
- (b) What is the probability that a student gets a strong letter $(L = l^1)$?
- (c) Given that the student is intelligent $(I = i^1)$, how likely is it that he/she will get a strong letter?
- (d) If the student did well on his/her SAT $(S=s^1)$, what is the probability that they will do well in the course $(G=g^1)$?
- (e) If the student was intelligent, did well on his/her SAT and got a great letter, how likely is it that the course was difficult $(D = d^1)$?

4. In this problem we will perform **principal component analysis (PCA)** to reduce the dimensionality of data in the Boston Housing dataset from 13 to 2, and visualize the dataset in this lower-dimensional space.

Normalize the features (such that each feature has mean 0 and standard deviation 1 over the data), perform PCA, extract the first two principal components for each datapoint, and include the following in your report:

- Report the values of the first two principal directions of the data. This should be two vectors of dimension 10.
- Create a scatter plot of the first two principal components of each datapoint in the dataset. Color each point based on its target value (see hint #2 for how to do this).

Please submit your code as **prob4.py**.

Hint #1: You can include the boston housing dataset in your code with

```
from sklearn.datasets import load_boston
data, target = load_boston().data, load_boston().target
```

Hint #2: You can create a scatterplot that assigns each point to a color based on it's target value with