COMP 5970/6970 Project 3: 100 points 20% Credit Final Submission due before 11:59 PM Monday March 16

Instructions:

- 1. This is an individual project. You should do your own work. Any evidence of copying either from a public source or from the works of other without due credits will result in a zero grade and additional penalties/actions.
- Submissions by email or late submissions (even by minutes) will receive a zero grade. No makeup will be
 offered unless prior permission has been granted, or there is a valid and verifiable excuse.
- 3. No show for your project presentation will receive a zero grade. There is also a penalty for missing a presentation day in which you are not presenting.

Submission:

For 5970, you are required to upload the following to canvas before 11:59 PM Monday March 16:

- Source Code: Python source files (upload .zip file in case of multiple files) containing your code only (no test data needed) and ReadMe.txt file (template provided) describing how to run your code. Note that we will NOT debug your code. If your code does not execute as described in ReadMe.txt, you will receive a zero grade.
- 2. **Presentation Slide**: One slide only in PPT/PPTX/PDF format to be used during the oral presentations (see below). If you submitted file spans more than a page, we will extract the first page for the oral presentation.

For 6970, you are required to upload the following to canvas before 11:59 PM Monday March 16:

- 1. **Source Code**: Python source files (upload .zip file in case of multiple files) containing your code only (no test data needed) and ReadMe.txt file (template provided) describing how to run your code. Note that we will NOT debug your code. If your code does not execute as described in ReadMe.txt, you will receive a zero grade.
- 2. **Presentation Slide**: One slide only in PPT/PPTX/PDF format to be used during the oral presentations (see below). If you submitted file spans more than a page, we will extract the first page for the oral presentation.
- 3. **Project Report**: Completed report document in PDF format using template provided. Make sure to have all necessary sections of scientific writing: abstract, introduction, methods, results, discussion, references.

Presentations:

Presentations will be during the class on Wednesday March 18 and Friday March 20.

Attendance is mandatory during all the presentation days. Missed presentation days without university-approved excuse will result in a <u>penalty of 25 points for each missed class</u>. Note that this penalty will be applied when you miss a presentation day in which you are not presenting. **No show for your project presentation will receive a zero grade.**

Everyone is required to deliver 3 minutes flash presentation accompanied by the submitted slide following the Three Minute Thesis (3MT) format, with additional 2 minutes for Q&A:

- 1. Your presentation should at least contain methods (i.e., implementation), results (e.g., output), and conclusion.
- 2. Having appropriate graphics and visuals (e.g., figures, plots) in the presentation slides to help illustrate key concepts or results will be positively graded.
- 3. Any additional scientific insights and/or challenges faced and/or limitations of your implementation and/or efficiency analyses and/or comparisons with alternative approaches will be positively graded.
- 4. Practice your talk not to exceed the time limit or finish too early.
- 5. No need to bring your slides. We will set things up and decide the presentation sequence.

Implementing Gaussian Naïve Bayes for protein Secondary Structure prediction

Objective: Implement Gaussian Naïve Bayes for protein secondary structure prediction.

Note: You must use standard Python programming language. You are NOT allowed to use non-standard packages or libraries (e.g. Biopython, scikit-learn, SciPy, NumPy, etc.).

A: Raw Data:

Two directories (*fasta* and *sa*) are supplied. The *fasta* directory contains 150 protein sequences in FASTA format. A FASTA file is as follows:

>sequenceID

 ${\tt AAGTAGGAATAATATCTTATCATTATAGATAAAAAACCTTCTGAATTTGCTTAGTGTGTATACGACTAGACATATATCAGCTCGCCGATTATTTGGATTATTCCCTG}$

The true 3-class secondary structure (SS) labels of these proteins can be found in the *ss* directory. Files in this directory are also in FASTA format. SS labels have three possible values:

'H': helix
'E': strand
'C': coil

N.B. The true SS labels are calculated using the DSSP (Dictionary of Protein Secondary Structure: Pattern Recognition of Hydrogen-Bonded and Geometrical Features. Kabsch and Sander, 1983) software and the resulting 8-class assignment has been transformed into 3-class.

B: Curating Training and Test Datasets:

Divide the raw data into non-overlapping sets of training (\sim 75%) and test (\sim 25%) datasets using simple random sampling without replacement.

C. Position Specific Scoring Matrix (PSSM) Files:

In addition to the *fasta* and *sa* directories, a *pssm* directory is also provided containing the evolutionary profile of the 150 proteins (one *pssm* file for each *fasta* sequence). Every row in a *pssm* file corresponds to the evolutionary profile of one amino acid. *pssm* files are generated using NCBI PSIBLAST software.

D. Feature Generation:

Use the values in the first 20 PSSM columns for each amino acid in a protein from the *pssm* file generated by PSIBLAST. Fist few lines for a sample *pssm* file are given below. For a protein sequence of N amino acids, there will be N x 20 PSSM values.

```
5 -4 -3 -6 -5
                                     3 -4 -6
      1 -2 -2 2 -4 1 1 0 -4 -3 -1 -2 -2 -5
         4 -2 1 -5 1 0 -3 0 -2 -3
        0 -2 0 -5 2 1 -1 -2 -4 -5
                                    0 -3 -6
                                             6 -1 -1 -6
5 S
      1 -2 1 5 -5 -1 3 -1 -1 -5 -5 0 -3 -6 0 1 -1 -6
      -1 -2 -5 -5 -4 -4 -3 -5 0 3 0 -3 -1 5 -2 -4 -1 -2
7 C
      -2 -6 -5 -6 11 -6 -6 -5 -6 -4 -4 -6 -4 -5 -6 -3 -3 -5
           1 -1 -3 1 0 -3 3 -2 2 0
                                       0 0 -5
      -1 -3 -5 -3 -4 3 2 -5 -3 -2 5 -3 2 -1 -3 -3 -3 -2
10 P
      -2 -3 -4 -3 -5 -1 0 -5 -3 -5 -4 -2 -4 -6
                                            8 -1 -3 -6
      2 0 -3 -3 -5 0 -1 -4 -1 -2 -2 4 1 -5 4
      -1 -2 -1 5 -5 -2 3 -4 -2 -2 -2
      -1 0 -3 -3 -3 0 0 -3 -4 -1 -3
                                    0 -2 -4 3 1 3 -5
      -2 -4 -3 -4 -5 -4 -4 7 -5 -7 -6 -4 -5 -6 -4 -3 -4 -5 -6
      -3 2 -2 -3 -5 -3 -2 -4 1 -1 -2 -1 -2 -3
```

Additionally, use a sliding window of 5 around the central residue (i.e. 2 residues on both sides) for feature generation and use the central residue's true SS type (from the corresponding ss file) for class label assignment (H/E/C). Therefore, there will be $20 \times 5 = 100$ PSSM values for each non-terminal residue. For terminal residues that do not have one or more neighbors on either side, use -1 as dummy PSSM values. E.g., for the first residue at the N-terminal, the feature vector will start with forty (20×2) dummy PSSM values of -1. Similarly, there will be twenty dummy PSSM values of -1 at the beginning of the feature vector for the second residue. There will be symmetrically opposite effect at the C-terminal with feature vectors ending with forty dummy PSSM values of -1 for the last residue and twenty dummy PSSM values of -1 for the second last residue.

E. Gaussian Naïve Bayes Learning on Training Set:

Implement the Gaussian Naïve Bayes learning algorithm that learns class priors and class conditional means and variances, assuming each $P(X_i | Y = y_k)$ to follow Gaussian distribution.

F. Gaussian Naïve Bayes Classification on Test Set:

Implement Gaussian Naïve Bayes classifier that takes the parameters learned during training to calculate the probability of each class and predict the class labels on any given test dataset by selecting the most probable class assignment (or a single sequence).

N.B. Gaussian Naïve Bayes is an offline-learning algorithm. Therefore, training and classification should be implemented separately. The classification algorithm should take a test file in FASTA format as an input (passed as a command line argument to your program) and predict the class labels (i.e. H, E, C) in FASTA format (just like ss file) in a standalone mode. You may save the parameters learned during training in a file that can be fed into the classifier, in an offline mode.

G. Evaluate Accuracy:

Use Q3 accuracy to evaluate the classification performance on the test dataset.