# Programming Assignment 3: Sequence Tagging

CSE 156: Statistical NLP: Spring 2019 University of California, San Diego

Due: May 13, 2019

Given a natural language sentence  $\mathbf{x} = \langle x_1, x_2, \dots, x_n \rangle$ , the task of sequence tagging is to find the most likely sequence of tags  $\hat{\mathbf{y}} = \langle y_1, y_2, \dots, y_n \rangle$ .

For this homework you will build a Hidden Markov Model (HMM) to tag gene names in biological text — an example problem of named entity recognition. Under this model the joint probability of a sentence  $x_1, x_2, \ldots, x_n$  and a tag sequence  $y_1, y_2, \ldots, y_n$  is defined as:

$$P(x_1 \dots x_n, y_1 \dots y_n) = q(y_1|*,*) \times q(y_2|*, y_1) \times q(\text{STOP}|y_{n-1}, y_n) \times \prod_{i=3}^n q(y_i|y_{i-2}, y_{i-1}) \times \prod_{i=1}^n q(x_i|y_i)$$

where \* is a padding symbol that indicates the beginning of a sentence and STOP is a special HMM state indicating the end of a sentence. Your task will be to implement this probabilistic model and a decoder for finding the most likely tag sequence for new sentences.

### 1 Preliminaries

#### 1.1 Data

We provide a labeled training data set gene.train, a labeled and unlabeled version of the development set, gene.key and gene.dev, and an unlabeled test set gene.test. Notice that there is no labeled test set for this assignment, reporting performance on the dev set will suffice. The labeled files take the format of one word per line with the word and tag separated by space and a single blank line separates sentences, e.g.

```
Comparison O
with O
alkaline I-GENE
phosphatases I-GENE
and O
5 I-GENE
- I-GENE
nucleotidase I-GENE
Pharmacologic O
aspects O
of O
neonatal O
hyperbilirubinemia O
.O
...
```

The unlabeled files contain only the words of each sentence and will be used to evaluate the performance of your model. The task is to identify gene names within biological text. In this dataset there is one type of entity: gene (GENE). The dataset is adapted from the BioCreAtIvE II shared task.

Several utility scripts are provided to help with the assignment. The scripts can be called at the command-line to pre-process the data and to check results.

### 1.2 Collecting Counts

The script <code>count\_freqs.py</code> handles aggregating counts over the data. It takes a training file as input and produces trigram, bigram and emission <code>counts</code>. To see its behavior, run the script on the training data and pipe the output into a file.

python count\_freqs.py gene.train > gene.counts

Each line in the output contains the count for one event. There are two types of counts:

• Lines where the second token is WORDTAG contain emission counts  $Count(y \leadsto s)$ , for example

13 WORDTAG I-GENE consensus

indicates that consensus was tagged 13 times as I-GENE in the training data.

• Lines where the second token is n-GRAM (where n is 1, 2 or 3) contain unigram counts Count(y), bigram counts  $Count(y_{n-1}, y_n)$  or trigram counts  $Count(y_{n-2}, y_{n-1}, y_n)$ , for example

16624 2-GRAM I-GENE 0

indicates that there were 16624 instances of an O tag following an I-GENE tag and

9622 3-GRAM I-GENE I-GENE O

indicates that in 9622 cases the bigram I-GENE I-GENE was followed by an O tag.

#### 1.3 Evaluation

The script eval\_gene\_tagger.py provides a way to check the output of a tagger. It takes the correct result and a user result as input and gives a detailed description of accuracy.

python eval\_gene\_tagger.py gene.key gene\_dev.p1.out

Found 2669 GENEs. Expected 642 GENEs; Correct: 424.

	precision	recall	F1-Score
GENE:	0.158861	0.660436	0.256116

Results for gene identification are given in terms of precision, recall, and F1-Score. Let  $\mathcal{A}$  be the set of instances that our tagger marked as GENE and  $\mathcal{B}$  be the set of instances that are correctly GENE entities. Precision is defined as  $|\mathcal{A} \cap \mathcal{B}|/|\mathcal{A}|$  whereas recall is defined as  $|\mathcal{A} \cap \mathcal{B}|/|\mathcal{B}|$  F1-score represents the geometric mean of these two values.

# 2 Baseline (35%)

 Using the counts produced by count\_freqs.py, write a function that computes emission parameters

$$e(x|y) = \frac{Count(y \leadsto x)}{Count(y)}$$

• We need to predict emission probabilities for words in the test data that do not occur in the training data. One simple approach is to map infrequent words in the training data to a common class and to treat unseen words as members of this class. Replace infrequent words (Count(x) < 5) in the original training data file with a common symbol \_RARE\_. Then re-run count\_freqs.py to produce new counts.

• As a baseline, implement a simple gene tagger that always produces the tag  $y^* = \arg\max_y e(x|y)$  for each word x. Make sure your tagger uses the <code>\_RARE\_</code> word probabilities for rare and unseen words. Your tagger should read in the counts file and the file <code>gene.dev</code> (which is <code>gene.key</code> without the tags) and produce output in the same format as the training file. For instance

Write your output to a file called gene\_dev.p1.out and evaluate by running

The expected result should match the result above.

• Your tagger can be improved by grouping words into informative word classes rather than just into a single class of rare and unseen words. Propose more informative word classes for dealing with such words.

In your report, describe your design decisions for rare and unseen words. Compare the model performance of a few different design choices on the train and development sets. Pick your best model and report its performance on the dev set.

## 3 Trigram HMM (65%)

• Using the counts produced by count\_freqs.py, write a function that computes parameters

$$q(y_i|y_{i-2}, y_{i-1}) = \frac{Count(y_{i-2}, y_{i-1}, y_i)}{Count(y_{i-2}, y_{i-1})}$$

for a given trigram  $y_{i-2}$   $y_{i-1}$   $y_i$ . Make sure your function works for the boundary cases:  $q(y_1|*,*)$ ,  $q(y_2|*,y_1)$ ,  $q(STOP|y_{n-1},y_n)$ .

• Using the maximum likelihood estimates for transitions and emissions, implement the Viterbi algorithm to compute

$$\underset{y_1...y_n}{\operatorname{arg\,max}} p(x_1...x_n, y_1...y_n)$$

Be sure to replace infrequent words (Count(x) < 5) in the original training data file and in the decoding algorithm with a common symbol \_RARE\_ . Your tagger should have the same basic functionality as the baseline tagger.

Run the Viterbi tagger on the development set. The model should have a total F1-Score of 40.

Your report should include a brief summary (e.g., a short paragraph) of your Viterbi algorithm implementation and any specific challenges or issues that came up. If you could not get your implementation working, describe where you got stuck.

- Here again, your tagger can be improved by grouping words into informative word classes rather than just into a single class of rare and unseen words. Propose more informative word classes for dealing with such words. Using the same strategy in Section 2 is fine.
  - In your report, describe your design decisions for rare and unseen words. Compare the model performance of a few different design choices on the train and development sets. Pick your best model and report its performance on the dev set.
- The Collins notes on HMMs should be very helpful for this part of the assignment.

## 4 Bonus: Non-trivial extensions (+5%)

There are several extensions you can consider making: smoothing the parameters, moving to 4-grams or higher, and so on. Bonus points will be earned for making non-trivial extensions to your tagger.

### 5 Submission Instructions

Submit your work on Gradescope.

- Code: You will submit your code together with a neatly written README file to instruct how to run your code with different settings. We assume that you always follow good practice of coding (commenting, structuring), and these factors are not central to your grade.
- Report: Submit your report, it should be four pages long, or less, in pdf (reasonable font sizes).

## 6 Acknowledgments

Adapted with some changes from a similar assignment by Michael Collins.