

Eidgenössische Technische Hochschule Zürich Swiss Federal Institute of Technology Zurich



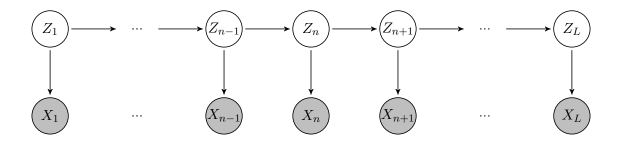
Statistical Models in Computational Biology

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Problem 6: Hidden Markov Models

(2 points)



Consider the HMM represented by the above graph, where X_n are observed variables and Z_n are hidden variables. Each hidden variable can take on K different values and the observed variables can take on M different realisations. The HMM is now parameterized as in the lecture on slide 18.

- (a) What is the maximum number of parameters to define the HMM? (1 point)
- (b) Assume K=2 and the transition matrix is

$$T = \begin{pmatrix} 0.1 & 0.9 \\ 0.4 & 0.6 \end{pmatrix}.$$

What is the stationary distribution π ?

(1 point)

Problem 7: Predicting protein secondary structure using HMMs

(8 points)

Proteins are molecules consisting of chains of amino acids. The different types of amino acids are labeled by different letters of the alphabet. A linear sequence of amino acids is called the primary structure of a protein. The pattern of hydrogen bonds in a biopolymer of a protein is the secondary structure. The Dictionary of Protein Secondary Structure (DPSS) uses seven letters to describe the pattern of hydrogen bonds in a biopolymer. One goal is to predict the secondary structure from the amino acid sequence. In a supervised learning setting, we can use amino acid sequences and matching secondary structures to predict the secondary structure.

Consider the HMM from the previous problem. The observed state space \mathcal{X} for the amino acid sequence consists of the letters A, C, D, E, F, G, H, I, K, L, M, N, P, Q, R, S, T, U, V, W, X, Y. The hidden state space \mathcal{Z} for the secondary structure consists of B, C, E, G, H, I, S, T. (https://en.wikipedia.org/wiki/Protein_secondary_structure).

You will use the Viterbi algorithm in order to predict the secondary structure of a protein. We provided you with files consisting of amino acid and secondary structure sequences, and an R implementation of the Viterbi algorithm. Every row in the tab-separated file represents one protein, where the first column is a protein identifier, the second column is the amino acid sequence X and the third column is the matching secondary structure of the protein Z. We also provide a file with amino acid sequences only for which we want you to predict the secondary structure.

The project consists of the following steps:

- (a) Read proteins_train.tsv, proteins_test.tsv and proteins_new.tsv into the memory and store each in a data.frame. (0 points)
- (b) Estimate the vector of initial state probabilities I, the matrix of transition probabilities T and the matrix for emission probabilities E by maximum likelihood. (1 point)
- (c) Estimate the stationary distribution π of the Markov chain by solving the eigenvalue problem (0.5 points) and by using a brute-force approach (0.5 points). (1 points)
- (d) Having estimated the parameters, i.e., the emission and transition matrices E,T and the vector of initial state probabilities I, you can predict the latent state sequence Z of a protein's amino acid sequence X using the Viterbi algorithm. Use the Viterbi algorithm provided in viterbi.r (carefully read the parameter description!) and iterate over each data.frame of proteins_test.tsv and proteins_new.tsv row by row and use the amino acid sequence to predict its secondary structure, which you add to the data.frame as a new column. Save the extended data.frame of proteins_new.tsv including the predicted secondary structure as a tsv file and hand it in together with your pdf. (1 points)
- (e) Estimate confidence intervals for each parameter in I,E and T with bootstrapping. In a single bootstrap run i estimate the probabilities for I_i,E_i and T_i the same as before, but not on the original data set proteins_train.tsv, but on the resampled data set. I.e., sample with replacement as many rows from proteins_train.tsv as the original data set has. Run a thousand bootstraps and compute the empirical 95% confidence intervals for each single parameter in $\{I_i\}_i, \{E_i\}_i$ and $\{T_i\}_i$. (2 points)
- (f) Use the following measure to compute the accuracy of the predicted secondary structure $P=(p_i)$ for the data.frame of proteins_test.tsv given the real secondary structure $S=(s_i)$:

$$a\left(P,S\right) = \frac{1}{L} \sum_{i} \begin{cases} 1 & \text{if } p_{i} = s_{i} \\ 0 & \text{if } p_{i} \neq s_{i} \end{cases}$$

with sequence length L. Compute the accuracy for every protein in your data.frame and store the accuracies in a vector. What is the accuracy of the Viterbi algorithm over all sequences (i.e. call summary on the vector of accuracies). (2 points)

(g) Instead of using the Viterbi algorithm, now randomly guess secondary structures for all sequences. Compare the global accuracies of the Viterbi and the random approach and plot all accuracy distributions using boxplots. (1 point)

References

[1] Durbin, Richard and Eddy, Sean R and Krogh, Anders and Mitchison, Graeme *Biological sequence analysis: probabilistic models of proteins and nucleic acids*. Cambridge university press, 1998

[2] Protein Structure. https://en.wikipedia.org/wiki/Protein_structure