HMM instructions

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Background

In this exercise, you will train a hidden Markov Model (HMM) to recognise **domain (D)** and **linker (L)** regions in protein sequences. To simplify the exercise, you will have a three letter amino acid alphabet: **H (hydrophobic)**, **P (polar)** and **C (charged)** amino acids.

Let a HMM be defined by:

- States, Q = {B, D, L, E}
- Alphabet, $\Sigma = \{H, P, C\}$
- Transition probabilities between the states, A1 =

	В	L	D	E
В	0	0.5	0.5	0
L	0	0.7	0.2	0.1
D	0	0.2	0.7	0.1
E	0	0	0	0

• Emission probabilities, *E*1 =

	Н	Р	С
L	0.5	0	0.5
D	0	0.5	0.5

Note: These matrices are also provided as .tsv files in the input directory.

And let an observed sequence be X1 = CCHHPCCPHHCH. This sequence is provided as *.fasta* file in the input directory.

Skeleton code

We provide you with a skeleton script called *hmm.py* and some helper functions in the module *hmm_utility.py*. **Check the directory of the input files.** Invoking the help option for *hmm.py* will produce the following output:

```
usage: python3 hmm.py [-h] [-v] [-o OUT_DIR] [-i MAX_ITER] [-c CONV_THRESH]
                      {viterbi, forward, backward, baumwelch} fasta transition
                      emission
 Perform the specified algorithm, with given sequences and parameters.
 Example syntax:
    python3 hmm.py -vv viterbi seq.fasta A.tsv E.tsv
    python3 hmm.py baumwelch in.fa priorA priorE -o ./outputs -i 1
positional arguments:
  {viterbi, forward, backward, baumwelch}
                        which algorithm to run
                        path to a FASTA formatted input file
  fasta
                        path to a TSV formatted transition matrix
 transition
  emission
                        path to a TSV formatted emission matrix
optional arguments:
                        show this help message and exit
  -h, --help
  -v, --verbose
                        print verbose output specific to the algorithm
                          (print even more output if flag is given twice)
  -o OUT_DIR
                        path to a directory where output files are saved
                          (directory will be made if it does not exist)
                          (file names and contents depend on algorithm)
                        maximum number of iterations (Baum-Welch only, default: 100 )
  -i MAX_ITER
  -c CONV_THRESH
                        convergence threshold
                                                     (Baum-Welch only, default: 0.01)
```

The skeleton script provides all the Input/Output functions so you can focus on the actual algorithms. Where possible most symbols/variables adhere to the formalism that is used in *Durbin et. al.* For any kind of matrix we use two-dimensional dictionaries.

Symbol in Durbin et al.	Usage in the skeleton code
a_{kl}	A[k][1]
$e_l(x_i)$	E[l][X[i]]
$v_l(i), f_l(i), b_k(i)$	V[l][i], F[l][i], B[k][i]

Viterbi algorithm

The Viterbi algorithm has been implemented for you. Inspect the code closely and see if it represents the formulas below. Run it on the sequence X1 = CCHHPCCPHHCH with the HMM defined by A1 and E1. You can find files for the sequence and matrices in the input directory. Make a new output directory and provide it to the -o argument. Run the script with different verbosity options (-v, -vv) and inspect the output. Have a look at the files the are created in the output directory.

Algorithm: Viterbi

Initialisation
$$(i=0)$$
: $v_k\left(0
ight)=1,\ v_k\left(0
ight)=0\ ext{for }k>0$
Iteration ($i=1...L$): $v_l(i)=e_l(x_i) ext{max}_k(v_k(i-1)a_{kl})$
Termination: $P(x,\pi^*)=v_E(L+1)= ext{max}_k(v_k(L)a_{kE})$

Note: This implementation varies slightly from the implementation in Durbin et al.. We introduce an additional end state E because we think it is clearer and more consistent.

Forward and backward algorithms

Implement the **forward** and **backward algorithms** (see <u>Biological sequence analysis</u> (https://canvas.vu.nl/courses/77637/files/8372916?wrap=1

(https://canvas.vu.nl/courses/77637/files/8372916/download?download_frd=1) , p.59-60). What is the probability for P(X1|HMM) in each case? Make different output directories for the different algorithms.

[Modify *hmm.py*; look for the ### START CODING HERE ### blocks in the provided template to see where code is missing.]

Algorithm: Forward

Initialisation (
$$i=0$$
): $f_B(0)=1,\ f_k\left(0
ight)=0\ ext{for }k>0$ Iteration ($i=1...L$): $f_l(i)=e_l(x_i)\sum_k f_k(i-1)a_{kl}$ Termination: $P(x)=f_E(L+1)=\sum_k f_k(L)a_{kE}$

Algorithm: Backward

Initialisation
$$(i = L)$$
: $b_k(L) = a_{kE}$, for all k

Iteration ($i=L-1,\ldots,0$): $b_k(i)=\sum_l a_{kl}e_l(x_{i+1})b_l(i+1)$ Termination: $P(x)=b_0(0)$

Note: we can continue the iteration until i=0. Pay attention to the index of the sequence and the index of table/dict b in Backward Algorithm.

Expected output for the forward algorithm with X1, A1 and E1:

```
>seq1
P = 4.39e-08

- C C H H P C C P H
B 1.00e+00 0.00e+00 0.00e+
```

Expected output for the backward algorithm with X1, A1 and E1:

```
>seq1
P = 4.39e-08

- C C H H P C C P H

B 4.39e-08 9.75e-08 2.17e-07 6.19e-07 6.19e-06 2.35e-05 5.22e-05 1.16e-04 1.16e-03 3.31e-03 1.12e-02 2.50e-02 0.00e+00 0.00e+00
D 3.27e-08 6.06e-08 8.66e-08 2.48e-07 8.66e-06 2.48e-05 6.14e-05 1.62e-04 4.64e-04 1.33e-03 7.00e-03 1.00e-02 1.00e-01 0.00e+00
L 4.62e-08 1.15e-07 3.03e-07 8.66e-07 2.48e-06 1.75e-05 3.25e-05 4.64e-05 1.62e-03 4.64e-03 1.32e-02 3.50e-02 1.00e-01 0.00e+00
E 0.00e+00 0.00e+00
```

Baum-Welch algorithm

Implement the **Baum-Welch algorithm** (see *Biological sequence analysis*, p. 65). Two optional arguments can be provided to the script for the maximum iterations (-i) and convergence threshold (-c). Run the Baum-Welch algorithm on sequence *X1*, and the matrices *A1* and *E1* given above.

[Modify *hmm.py*; look for the ### START CODING HERE ### blocks in the provided template to see where code is missing.]

Algorithm: Baum-Welch

Prepare new matrices in which you keep track of the observed Transition and Emission counts.

Iteration:

- 1. For each sequence j = 1...n:
 - Calculate $f_k(i)$ and P(x) for sequence j using the forward algorithm.
 - Calculate $b_k(i)$ for sequence j using the backward algorithm.
- 2. Add the contribution of sequence *j* to *A* and *E*

$$A^j_{kl}=\sum_i f^j_k(i)a_{kl}e_l(x^j_{i+1})b^j_l(i+1) \ / \ P(x^j)$$
and $E^j_k(s)=\sum_{(i|x^j_i=s)} f^j_k(i)b^j_k(i) \ / \ P(x^j)$

Pay attention to the index of the sequence X and the index of the table/dict b from the Backward Algorithm.

3. Calculate the new model parameters.

$$A_{kl} = \sum_j A^j_{kl}$$
 and $E_k(s) = \sum_j E^j_k(s)$

4. Normalise, so rows add up to 1:

$$a_{kl} = rac{A_{kl}}{\sum_{l'} A_{kl'}}$$
 and $e_k(s) = rac{E_k(s)}{\sum_{s'} E_k(s')}$

Termination: Stop iterating when you've reached either convergence or the maximum number of iterations.

Calculate the Sum Log-Likelihood (SLL) with your posterior parameters $m{ heta}$ and save your results.

SLL =
$$l(x^1,\ldots,x^n| heta)=\log_{10}P(x^1,\ldots,x^n| heta)=\sum_{j=1}^n\log_{10}P(x^j| heta)$$

Note: the termination step is implemented in the main function.

Example output for the Baum-Welch algorithm with one iteration (with sequence_X1.fasta as input):