

# BINF200 Assignment 2

Multiple sequence analysis, phylogenetics, motif analysis

2023-10-04

## 1 Deadline, grading and report

The assignment is due **13 October 2023**.

The assignment is scored on **20 points** and counts towards **10% of the final grade**.

Your report should be a **single PDF file** that contains your report text, code, and figures in a single document. The easiest workflow is probably to run your analyses in a Jupyter or similar notebook, and save the final notebook as a PDF file.

You *may* work together, but you *must* declare it in your report.

Any use of ChatGPT or other generative AI tools *must* be declared in your report.

## 2 Background

In this compulsory assignment, you will perform bioinformatics analyses covering multiple sequence alignment, phylogenetic tree construction and motif finding. The assignment will cover practical use of state-of-the-art tools, and also questions requiring programming. You may use any programming language: Python, Julia, R, ...

## 3 Data

Download all files in the following OneDrive folder:

[https://universityofbergen-my.sharepoint.com/:f:/r/personal/tom\\_michael\\_uib\\_no/Documents/public/BINF200/Coronavirus?csf=1&web=1&e=D5umem](https://universityofbergen-my.sharepoint.com/:f:/r/personal/tom_michael_uib_no/Documents/public/BINF200/Coronavirus?csf=1&web=1&e=D5umem)

You should find the following files:

- **protein\_N\_data.fasta** - Sequences of the gene coding for coronavirus nucleocapsid (N) protein in a number of coronaviruses
- **GCA\_011537005.1\_partial\_genomic.fasta** - Part of the BetaCoV/Wuhan/IPBCAMS-WH-02/2019 genome

- **motifCountMatrix.csv** - Count matrix of a sequence motif

## 4 Tasks

### 4.1 Multiple sequence alignment and phylogenetic tree construction for the coronavirus nucleocapsid protein (Total points: 5)

We will focus on different genera of corona viruses namely: alpha, beta, gamma and delta. Their genomes, gene and protein sequences, together with annotations and data reports are available from NCBI:

- Alpha coronavirus: <https://www.ncbi.nlm.nih.gov/datasets/taxonomy/693996/>
- Beta coronavirus: <https://www.ncbi.nlm.nih.gov/datasets/taxonomy/694002/>
- Gamma coronavirus: <https://www.ncbi.nlm.nih.gov/datasets/taxonomy/694013/>
- Delta coronavirus: <https://www.ncbi.nlm.nih.gov/datasets/taxonomy/1159901/>

For simplicity, we will use data from only one important gene, that encodes the coronavirus nucleocapsid (N) protein. This is a structural protein that forms complexes with genomic RNA, interacts with the viral membrane protein during virion assembly and plays a critical role in enhancing the efficiency of virus transcription and assembly. You can read more about it in the paper *The SARS-CoV-2 Nucleocapsid Protein and Its Role in Viral Structure, Biological Functions, and a Potential Target for Drug or Vaccine Mitigation*.

The datasets listed in Figure 1 were used to create **protein\_N\_data.fasta**.

#### 4.1.1 Parse the fasta file (1 point)

How many sequences are contained in the file **protein\_N\_data.fasta**?

List the names of the sequences.

#### 4.1.2 Find protein N in a specific coronavirus genome (1 point)

From the sequences in **protein\_N\_data.fasta**, find the sequence for which the first letter in its name is closest in the alphabet to the **first letter in your first name**. If there are multiple sequences starting with the same letter, pick one arbitrarily. For the selected sequence:

- Find the assembly accession ID in the table above.
- Go to the NCBI website (cf. links above) and find the corresponding genome assembly.
- What are the genomic coordinates (start and end position) of gene N in this genome? (Hint: follow the RefSeq link)

Genus	Organism Scientific Name	Organism Qualifier	Taxonomy id	Assembly Accession
Alpha	Human coronavirus NL63	strain: Amsterdam I	277944	GCF_000853865.1
Alpha	Bat coronavirus CDPHE15/USA/2006	strain: bat/USA/CDPHE15/2006	1384461	GCF_000913415.1
Alpha	Mink coronavirus strain WD1127	strain: WD1127	766791	GCF_000919475.1
Alpha	Camel alphacoronavirus	isolate: camel/Riyadh/Ry141/2015	1699095	GCF_001500975.1
Alpha	Ferret coronavirus	isolate: FRCoV-NL-2010	1264898	GCF_001661775.1
Alpha	Lucheng Rn rat coronavirus	isolate: Lucheng-19	1508224	GCF_001962315.1
Beta	Rabbit coronavirus HKU14	strain: HKU14-1	1160968	GCF_000896935.1
Beta	Middle East respiratory syndrome-related coronavirus	strain: HCoV-EMC	1335626	GCF_000901155.1
Beta	Betacoronavirus HKU24	strain: HKU24-R05005I	1590370	GCF_000930095.1
Beta	Betacoronavirus England 1	isolate: H123990006, strain: England 1	1263720	GCF_002816195.1
Beta	Severe acute respiratory syndrome coronavirus 2		2697049	GCF_009858895.2
Gamma	Beluga whale coronavirus SW1	isolate: SW1	694015	GCF_000872845.1
Gamma	Turkey coronavirus	isolate: MG10	11152	GCF_000880055.1
Gamma	Duck coronavirus		300188	GCF_012271565.1
Gamma	Canada goose coronavirus		2569586	GCF_012271745.1
Delta	Sparrow coronavirus HKU17	strain: HKU17-6124	1159906	GCF_000868165.1
Delta	Wigeon coronavirus HKU20	strain: HKU20-9243	1159908	GCF_000895415.1
Delta	Night heron coronavirus HKU19	strain: HKU19-6918	1159904	GCF_000896035.1
Delta	Common moorhen coronavirus HKU21	strain: HKU21-8295	1159902	GCF_000896895.1
Delta	Porcine coronavirus HKU15	strain: HKU15-155	1159905	GCF_002816235.1

Figure 1: Table of coronavirus source datasets

#### 4.1.3 Multiple sequence alignment (1 point)

Build a multiple sequence alignment for the **protein\_N\_data** using a multiple sequence alignment tool of your choice. (Hint: check out the services provided by EMBL's European Bioinformatics Institute (EMBL-EBI).)

#### 4.1.4 Phylogenetic tree reconstruction (1 point)

Based on the results from the previous step, build a phylogenetic tree. (Hint: at this stage it is not required to make an “advanced tree”, providing a simple tree is enough). Save the image of the phylogenetic/guide tree.

#### 4.1.5 Interpretation (1 point)

Based on the results from the previous two steps, what do you see? Elaborate with a small text (3-4 lines): Explain what you observe from the multiple sequence alignment itself (hint: check the number of conserved sites), and give a short interpretation of the phylogenetic tree you have constructed.

### 4.2 Step-by-step multiple sequence alignment and phylogenetic tree construction using UPGMA (Total points: 10)

#### 4.2.1 Compute pairwise similarities (2 points)

Use the Needleman-Wunsch (dynamic programming) pairwise alignment algorithm to build a matrix of global alignment scores for each pair of sequences in **protein\_N\_data.fasta**. You can choose between multiple options:

- Implement the Needleman-Wunsch algorithm yourself. (Hint: You have probably done this already in BINF100)
- Use an existing implementation of the algorithm. (Hint: Check biopython, biojulia)
- Use the *needleall* command line program from the EMBOSS suite. (Hint: You installed the whole EMBOSS suite for Assignment 1.)
- Use a webserver such as EMBL-EBI's EMBOSS Needle service. (Hint: Manually inputting every pair of sequences will be extremely tedious, though they do provide APIs.)

#### 4.2.2 Generate a pairwise distance matrix (4 points)

Generate a distance matrix from the score matrix you have created in the previous step. For this task we will use Feng & Doolittle's formulation, and we will compute the distance  $D$  using formula:

$$D = -\log S_{eff} = -\log \frac{S_{obs} - S_{rand}}{S_{max} - S_{rand}}$$

where

- $S_{obs}$  is the observed pairwise alignment score
- $S_{max}$  is the best alignment score for both sequences, obtained by taking the average of the score of aligning either sequence to itself
- $S_{rand}$  is the expected (average) score for aligning two random sequences of the same length and residue composition, obtained by random shuffling the nucleotide composition of the two sequences. (Hint: more info about the Feng & Doolittle can be found at this URL: <https://rna.informatik.uni-freiburg.de/Teaching/index.jsp?toolName=Feng-Doolittle>)

Compute  $S_{rand}$  by taking the average score of **10** pairwise alignments between random sequences with the same sequence compositions as the original sequences.

#### 4.2.3 Generate a “guide tree” of phylogenetic relationships (2 points)

Generate a “guide tree” of phylogenetic relationships from the pairwise distance matrix you have created in the previous step using the UPGMA method. You can choose between multiple options:

- Implement the UPGMA hierarchical clustering algorithm yourself. (Hint: You can represent the tree as a binary tree, either implementing a tree class yourself, or using an existing data structure.)
- Use an existing implementation of the algorithm. (Hint: UPGMA is more commonly known as hierarchical clustering with average linkage. Check SciPy or similar packages for other languages.)

#### 4.2.4 Interpret your results (2 points)

Visualize your guide tree and compare it to the phylogenetic tree constructed in Section 4.1.4. Elaborate with a small text (3-4 lines) to explain what you observe.

### 4.3 Sequence motifs (Total points: 5)

Do simple motif searching on corona virus sequences using the input dataset (**protein\_N\_data.fasta**) we have already analysed.

### 4.3.1 MEME analysis (1 point)

Connect to the MEME platform at <https://meme-suite.org/>.

- Find the MEME motif discovery tool.
- Input **protein\_N\_data.fasta** to discover enriched motifs in this set of sequences, allowing for zero or one motif occurrence per sequence and finding upto 5 motifs. Which discovery mode, sequence alphabet, and site distribution options do you select?

Open and download the **MEME HTML output file** and include the sequence logos of the motifs found in your report.

### 4.3.2 Convert count matrix to PWM (2 points)

We will work with a 20-nucleotide subset of the first motif found by the MEME software, given by the count matrix:

base\position	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18	19	20
A	0	0	0	0	20	0	0	19	0	11	0	6	0	0	8	20	0	9	0	0
C	0	0	12	0	0	8	0	0	8	8	11	1	0	0	7	0	20	0	0	0
G	0	0	0	0	0	0	0	0	0	0	0	3	20	20	0	0	0	0	20	20
T	20	20	8	20	0	12	20	1	12	1	9	10	0	0	5	0	0	11	0	0

Figure 2: Motif count matrix

The count matrix is also available as a file **motifCountMatrix.csv**.

1. Compare the count matrix against your sequence logos and mark the 20-nucleotide window corresponding to this count matrix in the right logo.
2. Convert the count matrix to a position-specific probability matrix (PPM)  $P$ . To avoid zeros in the PPM, we add *pseudo-counts* and define

$$P_{k,i} = \frac{\text{Count}_{k,i} + 0.25 * \sqrt{N}}{N + \sqrt{N}},$$

where  $\text{Count}_{k,i}$  is the value of the count matrix for nucleotide  $k$  in motif position  $i$ , and  $N$  is the number of sequences in **protein\_N\_data.fasta** (Hint: Count the totals in each column of the count matrix).

3. Convert the PPM matrix to a position-specific weight matrix (PWM)  $W$  using the formula

$$W_{k,i} = \log_2 \frac{P_{k,i}}{0.25}$$

What would be the value of  $W$  for a random background site with equal counts for all nucleotides and using the pseudo-count formula above to compute the random probabilities?

#### 4.3.3 Scan a coronavirus genome for motif occurrences (2 points)

Scan part of the BetaCoV/Wuhan/IPBCAMS-WH-02/2019 genome (the sequence in the file **GCA\_011537005.1\_partial\_genomic.fasta**) and score all possible motif occurrences. Use the sliding window approach presented in the lecture and report (figure) both the log-odds score and the odds of each possible motif starting position in the genome sequence.

Elaborate with a small text (3-4 lines) to explain what you observe.