# Practical 6 11 March 2024

## **Questions**

- 1. Using AL2CO server (<a href="http://prodata.swmed.edu/al2co/al2co.php">http://prodata.swmed.edu/al2co/al2co.php</a>), obtain the positional conservation scores from multiple sequence alignment (MSA) of given set of protein sequences (set1 and set2) using the methods given below:
  - (i) Unweighted frequency and entropy-based measure
  - (ii) Unweighted frequency and variance-based measure
  - (iii) Unweighted frequency and sum of pairs measure
  - (iv) Weighted frequency and variance-based measure
  - (v) Normalize the scores obtained with (i)

## **Sequences:**

### Set 1

P69905, P01946, P01942, P01966, P01958, P01959, P01965, P06635, P60529, P80043 and P01980

#### Set 2

TPIS\_HUMAN, TPIS\_YEAST, TPIS\_GRAGA, TPIS\_TRYCR, TPIS\_MAIZE, TPIS\_MOUSE, TPIS\_DROME, TPIS\_RABIT, TPIS\_CAEEL

- 2. Tabulate the topmost 10 residues with highest and lowest conservation scores (in both Set1 and Set 2) obtained with method (i).
- 3. Write a program to compute the conservation score from MSA using unweighted frequency, and entropy, variance and sum of pairs-based measures.
- 4. Using the program written in Q3 (unweighted frequency and entropy-based measure), compare the MSA from Clustal Omega, MAFFT, and MUSCLE. Identify the residues with (i) similar and (ii) different conservation scores among the three alignment methods.
- 5. Check the scores manually at positions 9, 11, 20, 22 and 30 (use MSA from Clustal Omega)
- 6. Obtain the conservation score of 1BTM, A-chain using Consurf server (<a href="https://consurf.tau.ac.il/">https://consurf.tau.ac.il/</a>)

Deadline: 17th March 2024