Lab sheet#6

ExPASy Tools

Objective:

- Translation of a nucleotide sequence to a protein sequence using ExPASy.
- Analysis of protein primary structure and computing various physical and chemical parameters

A. Translation of a nucleotide sequence to a protein sequence using ExPASy web page:

- 1. Retrieve the FASTA format of homo sapiens X-ray repair cross complementing 1 (XRCC1) mRNA sequence (NM 006297.3) from the NCBI GenBank database.
- **2.** Use the translate tool on ExPASy website:
 - ➤ Paste the retrieved XRCC1 mRNA sequence into the box.
 - ➤ Click the "Translate sequence" button.
 - ➤ Choose the most reliable predicted protein sequence.

B. Primary Structure Analysis of a Protein Using ProtParam on the ExPasy server:

- 1. Search for ProtParm, click the first link.
- **2.** Copy the protein sequence of XRCC1 protein (CAG33009.1), then paste in the ProtParm webpage.
- **3.** Click compute parameters.
- **4.** Find out the Molecular weight of the protein.
- **5.** How many Cysteine amino acids located in the protein sequence.
- **6.** What is the total number of negatively charged residues.
- 7. Find out the Estimated half-life of the protein in mammals and yeast.

✓	Exercise:	
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- ✓ Find the most reliable predicted protein sequence for human TRF2 mRNA (NM 005652.4).
- ✓ What is the molecular weight, Ip and number of positively charged residues in TRF2 (NP 005643.2).

Lab sheet#7

Structure visualization using jmol

Objectives:

- Download protein sequence as PDB format using PBD ID.
- To be familiar with structure visualization program (Jmol).
- Display 3D structure of a protein, change its view, color and select motifs by writing commands.

Use Jmol program to perform the following:

Jmol is a free molecular viewer, used to create and view three dimensional structures of proteins. This viewer is linked to the Protein Data Bank (**PDB**) website, which provide biological macromolecular structures.

Note: in case of any difficulty downloading Jmol as an independent program on a desktop, you can use the web-based version (oftern referred to as "JSmol") that runs on a JavaScript platform (https://chemapps.stolaf.edu/jmol/jsmol/jsmol.htm). Make sure that Java is installed.

Visualization of ATP-binding subunit of the histidine permease from salmonella typhimurium.

Each structure hosted on the Protein Databank has a unique four character long alpha-numeric identifier, referred to as the structure's **PDB ID**.

- 1. Open Protein Data Bank (PDB) website, Search for PDB ID of the protein above, download the structure as PDB format, then answer the following questions:
 - Who are the authors of the PDB file?
 - From what source was this molecule isolated?
 - In which journal was the primary citation published?
 - On what date was the file deposited into the PDB?
 - How many chains are in this file?
 - Are there any heterologous groups within this PDB file? If so, which ones?

- 2. Open Jmol program and get the protein structure by entering PDB ID. (file→ load→open from PDB).
- 3. Change Style display from Atom style to <u>Cartoon</u> scheme style. (Click right > style > scheme > cartoon).
- 4. Open the console window, change the color of the whole structure to grey.

 (Click right → console→type: select all→ press enter→ type: color grey→ press enter).
- **5.** Open the protein page in <u>protein database (NCBI)</u> to know the different motifs of the protein and their location.
- 6. Select the Walker A/P-loop motif (39-46), and color it by blue. (type: select 39-46 → press enter → type: color blue → press enter).
- 7. Select the ABC transporter signature motif 154-163, and color it to green.

 (type: select 154-163 → press enter → type: color green → press enter).
- 8. Select Walker B motif 174-179, and color red.

 (type: select 174-179 → press enter → type: color red → press enter).
- 9. Show the ATP ligand as <u>Ball and stick</u> scheme style.

 (type: select ligand → click right → style→ scheme → ball and stick).
- 10. Change the measurement unit from nm to Angstroms.

 (click right → measurements → distance unites angstroms)
- 11. Show which one is closest to Walker A/P-loop motif and measure the distance between them. (double click on the ligand then drag the mouse towards the motif)
- 12. Save the protein structure as a picture. (right click \rightarrow file \rightarrow save \rightarrow save as PNG)

Here is \rightarrow Video displaying the steps to visualize the protein above:

https://drive.google.com/file/d/1cD4R0XxGxRbi40LI4FJlrbIoz79_3GrW/view?usp=sharing

Exercise

- 1. Open Protein Data Bank (PDB) website, Search for PDB ID of T3R3 Human Insulin Hexamer.
 - Go down to the molecule description to see how many polymers and chains does insulin have.
- 2. Open Jmol program and get the protein structure by entering PDB ID. (file > get PDB).
- **3.** Change Style display from Atom style to <u>Cartoon</u> scheme style.
- **4.** Change Style display to **<u>Backbone 1.5</u>** scheme style.

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(Click right →console→type: Backbone 1.5→press enter).
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- 5. Turn cartoon style off. (type: cartoon off→press enter)
- **6.** Select sheets and color red.

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(type: select sheets → press enter → type: color red → press enter)
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7. Select helix and color blue.

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(type: select helix → press enter → type: color blue → press enter)
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8. Show cysteins (Sulfur) that forms disulphide bridges "showing how the polypeptides hold together through S-S bonds". Change to wireframe 1.25 and color them yellow.

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(type: select sulfur → press enter→ type: wireframe 1.25→ press enter→ type: color yellow).
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- How many disulphide bonds are found in insulin protein?
- Show for each disulphide bridge the position of each Cys and the chain involved.
- 9. Move the structure 0 360 0 0 0 0 0 10. (type: move 0 360 0 0 0 0 0 10 → press enter)
- 10. Save the protein structure.

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