Lesson activities

▼ (1) Explore Databases

Databases

- 1. We will be using the National Center for Biotechnology Information (NCBI) Genbank Database in our next activity:
 - NCBI: https://www.ncbi.nlm.nih.gov/
- 2. Interactive Question: What are other public databases that contain genetic information or metadata?
 - Answer after question prompt #1 on the shared google doc.
- 3. How to use NCBI's Genbank database:
 - Introduction to NCBI Basic Local Alignment Search Tool (BLAST) BLAST is a program that can
 detect sequence similarity between a Query sequence and sequences within a database.
 - There are great existing tutorials and slides that have been shared to better understand the NCBI's
 BLAST alignment algorithm.
 - http://www.shodor.org/media/content//petascale/materials/dataIntensive/BLAST/BLAST_Intro_pdf.pdf
 - https://community.gep.wustl.edu/wiki/images/2/28/2011_8b_BLASTrv7_rev.pdf

▼ (2) Mystery Species Activity

Let's get started:

- Open the mystery species data file: mystery_species.fasta.txt
- 2. Navigate to NCBI Blast
- 3. Choose any DNA sequence from the file and copy the sequence.
- 4. Paste the sequence in the "Enter Query Sequence" window.
- 5. Search all "Standard databases (nr etc.)" for "Highly similar sequences (megablast)"
- 6. Investigate the top match. Use a search engine to retrieve a photo of your mystery species!
- ▼ (3) Comand Line Basics

Follow UNIX Setup instructions used in The Carpentries https://carpentries.org/:

- Setup: https://swcarpentry.github.io/shell-novice/setup.html
- Window's users install Git for Windows:

- https://gitforwindows.org/
- or Putty:
 - http://faculty.smu.edu/reynolds/unixtut/windows.html

Useful Links

- · Command Line for Mac and Windows
 - Lesson 1: https://swcarpentry.github.io/shell-novice/01-intro/index.html
 - Lesson 2: https://swcarpentry.github.io/shell-novice/02-filedir/index.html
 - Lesson 3: https://swcarpentry.github.io/shell-novice/03-create/index.html
- ▼ (4) Data Analysis & Data Visualization (usegalaxy.org)

Sign up and Tutorial Steps

- 1. Sign-up for a Galaxy account at usegalaxy.org
- Upload sequences file to UseGalaxy.org instance. (Zika data from Next Strain.org; https://nextstrain.org/zika)
- 3. Run Multiple Sequence Alignment using MAFFT.
- 4. Visualize the alignment.
 - Integrated MSA Viewer vs IGV.
 - Interpret the results.
- 5. Tree building (zika virus; https://nextstrain.org/zika)
 - Generate a Phylogeny (study evolution) using IQtree.
 - Visualize the phylogeny using the integrated "Phylogenetic Tree Visualization" tool.

Useful Links

- Introduction to Galaxy
 - https://galaxyproject.org/learn/
 - https://training.galaxyproject.org/training-material/topics/introduction/
- ▼ (ADVANCED) Nextstrain.org Zika Tutorial
 - 1. Follow the Next Strain tutorial on how to set up a visualization of Zika virus data: https://nextstrain.org/docs/tutorials/zika.
- 2. Follow Installation/Setup Instructions.
- 3. Helpful Tips:
 - Install latest version of Node.js : https://nodejs.org/en/
 - Best to Install Augur & Auspice with Conda.

4. Follow the commands to analyze the data provided.

Note: Update to last command in Zika Tutorial

• "Explore the Results" to work with the latest version of augur.