Nextstrain Tutorial

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In this tutorial we are going to generate a nextstrain.org community build. To do so, we will run a bioinformatics pipeline (augur), which will perform sequence alignment, phylogenetic reconstruction, infer ancestral states (for location and mutations), and finally combine such results to be visualized using a tool named auspice.

- (1) **Create** a folder in your directory of preference: it will be your working directory (workdir).
- (2) **Access** the repository below and **download** its content as a .zip file:

github.com/grubaughlab/nextstrain course

- (3) **Decompress** the downloaded .zip file and **copy** its content into your workdir.
- (4) Now, your workdir should look as follows:

workdir/
config/
auspice_config.json
colors.tsv
dropped_strains.txt
lat_longs.tsv
reference.gb
data/
metadata.tsv
sequences.fasta
LICENSE
README.md
Snakefile
metadata.xlsx
nextstrain_installation.pdf
nextstrain tutorial.pdf

- (5) Open the file metadata.xlsx. This spreadsheet contains three tabs, named after some of the files in workdir, they are: metadata, lat_longs, and colors.
- (6) **Copy** and **paste** the data from each tab in the corresponding files, replacing their existing data.
- (7) The current reference.gb file is included in config/only as an example. **Inspect** the format of this file using a text editor, and **check** the fields 'CDS' and 'gene' in this annotation file. **Access** the link below, and **download** this new reference genome in .gb format (**click** on Send to > File > Format:Genbank > Create File).

ncbi.nlm.nih.gov/nuccore/JX669468

- (8) **Open** this .gb file on a text editor, and **edit** it as follows, keeping all the edited elements aligned, so that it looks like the file inspected at step (7):
 - a. Replace 'mat_peptide' by 'CDS'
 - b. Replace 'product' by 'gene'
 - c. **Rename** long gene names at 'gene' fields by standard gene acronyms (check the literature)
- (9) **Delete** the existing reference.gb; **move** the file created in the previous step to workdir/config, and; **rename** it as your new reference.gb file.
- (10) **Open** the file auspice_config.json using a text editor and **edit** the lines showing the following content:
 - a. 'Add your project title here'
 - b. 'Add your name here'
 - c. 'Add your github link repository here'
- (11) **Open** the file Snakefile using a text editor, and **edit** the following lines as appropriate, keeping their suffixes ('tree' and 'meta'):
 - a. 'addHereYourProjectName tree.json'
 - b. 'addHereYourProjectName_meta.json'
- (12) Activate nextstrain: conda activate nextstrain
- (13) Having done all the edits above, **access** your workdir in the Terminal, and **run** augur using the command below:

snakemake export

- (14) **Watch** for potential error messages: they may indicate issues with file formatting in previous steps.
- (15) If no issues were flagged, now **run** auspice to visualize the results using the command below:

auspice view

(16) **Open** a web browser and **access** the results using the link generated by the command above, which may look like the url below:

http://localhost:4000