# **User manual for Phylobook**

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#### Overview:

As the volume of sequence data from variable pathogen genomes increases, means of analyzing, annotating and extracting specific taxa and taxon groups for study becomes more difficult. To meet these challenges for moderately large datasets (e.g., hundreds to thousands of taxa) the "Phylobook" tool was developed. Phylobook displays phylogenetic tree data adjacent to highlighter plots showing the position of mutations in the alignment. A key feature of Phylobook is that it allows the user to identify lineages and recombinants within a given dataset, annotate them and then export selected subsets of sequences for downstream analysis. Lineage identification can be aided using annotations created by one or more clustering methods.

## **Installation and Configuration:**

Phylobook is distributed as a Docker container. A container is a standard unit of software that packages code and all its dependencies so that the application runs quickly and reliably from multiple computing environments, including Mac, Windows and Linux systems. Docker containers run under a "Docker engine", a virtual machine that is available on most common operating systems.

Install Docker on your host server: <a href="https://docs.docker.com/get-docker/">https://docs.docker.com/get-docker/</a>. Note that Docker Compose is an additional tool that is needed and it is automatically included with Mac and Windows downloads of Docker. However, <a href="Linux">Linux</a> users will need to add it manually. You can do this by running the following command after your Docker installation is complete.

sudo pip install docker-compose

(pip = python installer)

- 2) Install Git on your computer.
  - a. On a Mac, if you have previously installed the developer tools, Git is already installed. If you have not previously installed the developer tools you can obtain Xcode from the app store and run the installer.
  - b. On Windows, git can be obtained at https://gitforwindows.org/

- c. On Linux, Git is specific for the specific version of Linux but most common options can be found at https://git-scm.com/download/linux.
- 3) Create 2 new folders. One is the folder into which Phylobook will be installed, the other is the folder that will contain the data that Phylobook uses. These folders can be anywhere on the host computer.
- 4) Install Phylobook. In a terminal window, navigate to the install folder (/Applications/Phylobook in this example) and type the command:

git clone https://github.com/MullinsLab/phylobook.git

This will install phylobook in the install directory. If you are on a mac and used the example folder name of /Applications/Phylobook, this step will create a folder named /Applications/Phylobook/phylobook and it will populate it with the phylobook software.

5) Create an environment file. Change directory to the newly created phylobook folder by typing

cd phylobook

Create the .env file by copying the .env.Template file to .env using the command:

cp .env.TEMPLATE .env

Note: On most computers, files that start with a "." are hidden from display by default so if you wish to see or edit the file with a GUI based editor (like textedit), you will need to tell your computer to show hidden files (the Mac OS command in the folder you wish to display hidden files is: cmd + shift + "."). If you are comfortable with a command line editor, you can edit .env in the terminal window.

6) Configure Phylobook by editing the .env file for your own environment. Phylobook can be set up to run locally (e.g., just on your laptop or desktop) or it can be run as an internet connected server. As an internet connected server, the login can be configured to be either local, Single Sign-On (aka, "SSO"), or dual, i.e., local and SSO.

Note: SSO authenticates users using an institutional or corporate user authentication service and is most useful if every intended user belongs to the same organization. SSO must be configured to connect to your organization's Open Authorization server and configuration will likely require assistance from a local IT professional. Local authorization will require a username and password for each user and those usernames/passwords will be local to the Phylobook server. Dual login allows for a mix of local usernames/passwords and SSO. This is most convenient when some of the users are from the same organization and other users are collaborators from external organizations. Since SSO configuration is institution specific, the following describes configuration as a local server.

If you are comfortable with a command line editor, you can edit .env in the terminal window.

Some lines (e.g., DEBUG, SERVER NAME, and EMAIL\_\*, should only be edited by developers or to change passwords). The lines that need to be edited in the .env file are:

#### SECRET KEY=PUT A UNIQUE PASSWORD HERE

The password can be anything the user desires. If you need help generating a strong password, a number of websites can be used for that purpose. For example, either <a href="https://djecrety.ir/">https://randomkeygen.com/</a> can be used to generate a strong password. The secret key is used by the Django web framework.

• DJANGO ALLOWED HOSTS=localhost 127.0.0.1

If phylobook will be run local to your computer or from behind a reverse proxy such as NGNIX or Apache (not covered here), you can leave this line unchanged. Otherwise, it should be completed with the server and domain name.

PROJECT PATH=

The PROJECT\_PATH directory is the primary storage location for project data within Phylobook. PROJECT\_PATH should be set to the desired location of the data directory. In our mac example, this might be /Applications/Phylobook/Data

- DB USER=postgres
- DB NAME=postgres
- DB PASS=

In most cases, the DB\_USER and DB\_NAME can be left as PostgreSQL. However, if you have another server that is your research group's primary database server AND you wish to store the database outside of the Docker container, you may wish to configure Phylobook to use that alternate database. We do not recommend this. DB\_PASS should be set to a unique password. Other configuration changes may be needed for your specific environment but just those mentioned above are sufficient to get Phylobook running on your local computer.

- EMAIL HOST=outlook.office365.com
- EMAIL\_USE\_TLS=1
- EMAIL\_PORT=587
- EMAIL HOST USER=
- EMAIL HOST\_PASSWORD=

Passwords can be changed on the Admin panel (the "(Admin)" link at the top of each Phylobook page) by the administrator. In addition, Phylobook has the ability to send password reset emails if a user has forgotten their password. In order for this to work, the .env file needs to contain valid email settings, including the login credentials called for above. Collecting these credentials is outside the scope of this manual.

Note: Although not recommended, but if you chose LOGIN\_TYPE=SSO or LOGIN\_TYPE=dual in config.env, then you must create settings/saml.py by copying settings/saml.py.TEMPLATE:

cp settings/saml.py.TEMPLATE settings/saml.py

Edit settings/saml.py and add your institution's SAML configuration and certificates:

nano settings/saml.py

7) Build and deploy the docker containers by issuing the following command:

docker compose up -d --build

After you have issued that command, you should see phylobook running under docker (viewable in the Docker Desktop), e.g., see **Figure 1**. If this does not appear automatically, simply launch the Docker app.

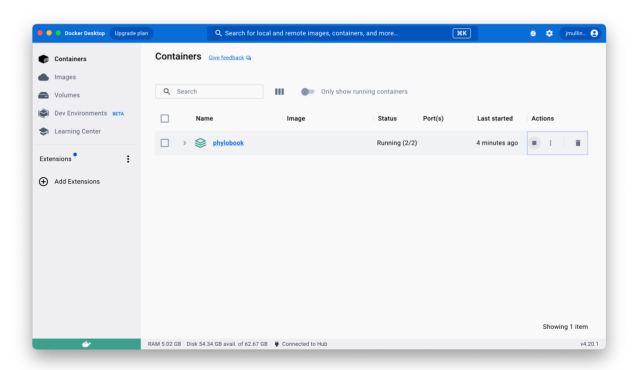


Figure 1. Docker desktop.

Note: Depending on the location of the PROJECT\_PATH directory, you may need to make the directory available to Docker as a shared folder. Do this by going to Settings > Resources > File Sharing in the Docker desktop and adding the directory as shown in **Figure 2**.

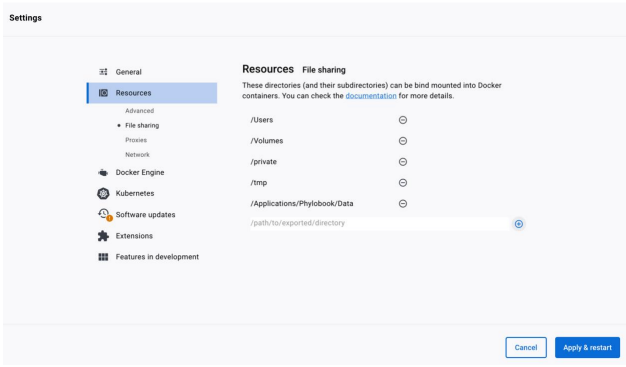


Figure 2. A view of Docker Desktop showing how to share a local directory with a Docker container.

8) The following command will perform initial database migrations:

docker exec -it phylobook python manage.py migrate --settings=phylobook.settings.prod

9) The following command will create the initial superuser/admin for the phylobook system:

docker exec -it phylobook python manage.py createsuperuser --settings=phylobook.settings.prod

After executing this command you will be asked for a username and a password. These credentials will be used for the initial login to Phylobook.

- 10) In a web browser, login to the server http://localhost:8000/ using your super user account credentials. In the upper right corner of the browser window, there is an "Admin" link. Click the link and you will enter the Administrator module.
- 11) Prior to using Phylobook for analysis, data files must be placed in the PROJECT\_PATH directory and Phylobook. Later in this manual we will discuss how to generate data files for use in Phylobook. For now, we will show how to setup Phylobook using sample data provided as part of the Phylobook installation. Sample data files are provided as .tar.gz files open these files to expand into folders.

Note: Multiple datasets may exist in each project folder. Each dataset will be represented by multiple files that start with the dataset name and that end with names associated with the

type of file. While the Phylobook pipeline produces a large number of files (**Figure 3**), only three files are required for each project, with additional files being optional (**Table 1**).

	IV_DEMO	2	Today at 12:11 PM	- <del></del>	Folder
~	run_phyml_logs	9	Today at 12:11 PM		Folder
	run_phyml.log	0	Mar 2, 2023 at 1:45 PM	892 bytes	Log File
	DEMO.log	9	Mar 2, 2023 at 1:45 PM	891 bytes	Log File
- ~	DEMO.phy_phyml_tree.txt.svg	0	Mar 2, 2023 at 2:43 PM	38 KB	SVG image
1	DEMO.json	9	Mar 2, 2023 at 2:43 PM	322 bytes	JSON Documen
	DEMO_highlighter.png	0	Mar 2, 2023 at 1:45 PM	545 KB	PNG image
	DEMO_highlighter.fasta	9	Mar 2, 2023 at 1:45 PM	176 KB	BBEdit Docume
I C	DEMO_highlighter.txt	9	Mar 2, 2023 at 1:45 PM	21 KB	text
	DEMO_highlighter_untrimmed.png	9	Mar 2, 2023 at 1:45 PM	556 KB	PNG image
M	DEMO.phy_phyml_tree.txt_nexus.tre	0	Mar 2, 2023 at 1:45 PM	5 KB	FigTree tree file
a		0	Mar 2, 2023 at 1:45 PM	2 KB	FigTree tree file
10		0	Mar 2, 2023 at 1:45 PM	43 KB	text
100	DEMO.phy_log.txt	9	Mar 2, 2023 at 1:45 PM	231 bytes	text
10	DEMO.phy_phyml.txt	9	Mar 2, 2023 at 1:45 PM	81 KB	text
HC   112	DEMO.phy_phyml_tree.txt	9	Mar 2, 2023 at 1:45 PM	2 KB	text
(1) (1)		9	Mar 2, 2023 at 1:45 PM	3 KB	text
3	DEMO.phy	9	Mar 2, 2023 at 1:45 PM	172 KB	Alignment file
-	DEMO.fasta	9	Mar 2, 2023 at 1:43 PM	174 KB	BBEdit Docume
	DEMO.cluster.km9clusters	9	Dec 7, 2022 at 4:06 PM	656 bytes	Document
173.	DEMO.cluster.km8clusters	9	Dec 7, 2022 at 4:06 PM	656 bytes	Document
	DEMO.cluster.km7clusters	9	Dec 7, 2022 at 4:06 PM	656 bytes	Document
	DEMO.cluster.km/clusters	9	Dec 7, 2022 at 4:06 PM	656 bytes	Document
	DEMO.cluster.km5clusters	9	Dec 7, 2022 at 4:06 PM	656 bytes	Document
	DEMO.cluster.km/sclusters	9	Dec 7, 2022 at 4:06 PM	656 bytes	Document
	DEMO.cluster.km3clusters	9	Dec 7, 2022 at 4:06 PM	656 bytes	Document
	DEMO.cluster.km3clusters	9	Dec 7, 2022 at 4:06 PM		Document
	DEMO.cluster.km10clusters	9	Dec 7, 2022 at 4:06 PM	656 bytes 657 bytes	Document
	DEMO.cluster.Kimoclusters  DEMO.cluster.GAPclusters				Document
- VAVA	_SARS_CoVSpike	9	Dec 7, 2022 at 4:06 PM  Today at 11:13 AM	656 bytes	Folder
	_SARS_COVSpike un_phyml_logs	0	Today at 11:13 AM		Folder
2000					
	run_phyml.log	0	Mar 1, 2023 at 12:35 PM	1 KB	Log File
-	WA_SARS-CoV2_SpikeAA.log	0	Mar 1, 2023 at 12:35 PM	1 KB	Log File
	VA_SARS-CoV2_SpikeAA.phy_phyml_tree.txt.svg	0	Mar 1, 2023 at 2:52 PM	105 KB	SVG image
	VA_SARS-CoV2_SpikeAA.json	0	Mar 1, 2023 at 2:52 PM	489 bytes	JSON Documer
□ W	VA_SARS-CoV2_SpikeAA_highlighter.png	0	Mar 1, 2023 at 12:36 PM	766 KB	PNG image
N III	VA_SARS-CoV2_SpikeAA_highlighter.fasta	0	Mar 1, 2023 at 12:36 PM	187 KB	BBEdit Docume
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Figure 3. Pipeline output files present in the HIV\_DEMO and WA\_SARS\_CoVSpike folders.

Table 1. User files used by Phylobook

File type	Requirement	File naming convention	Source
Image of	Required	DatasetName.phy_phyml_tree.txt.svg	Pipeline
phylogenetic			
tree			
Image of	Required	DatasetName_highlighter.png	Pipeline
highlighter			
plot			
Alignment	Required	DatasetName_highlighter.fasta	Pipeline
FASTA			
Cluster	Optional	DatasetName.cluster.clusters	Clustering algorithm
Assignments			https://github.com/MullinsLab/ClusteringForPhylobook

12) <u>Making a project visible within the phylobook server:</u> Once data is available in the PROJECT\_PATH directory, the Admin tool in Phylobook is used to make that project available within the system. After logging in (step 11 above), you will see an "Admin" link in the upper right corner of the browser window. Click the link and you will enter the Administrator module (Figure 4). The admin module allows an administrator to add projects, project categories, users, and groups to the system.

# Phylobook Admin

# Phylobook Admin

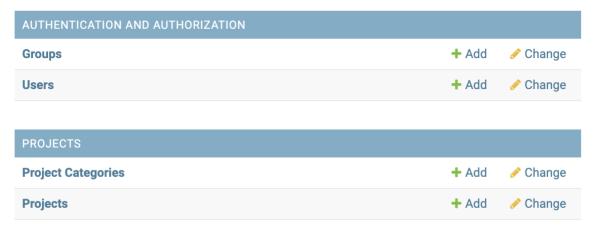


Figure 4. The Phylobook Administration module.

To add a project to Phylobook, click on the Add link next to "Projects" and type the EXACT name of the folder in the Project path directory. For example, to make the sample folder HIV\_DEMO available you would enter it in the dialog below and SAVE (**Figure 5**).



Figure 5. Adding a project to the Phylobook server

Follow the same process to add the WA\_SARS\_CoVSpike dataset. Once added a project will appear in the Projects list in the admin module and on the landing page within Phylobook.

For users who have a large number of projects stored within Phylobook, the landing page can become a bit cluttered with a very long list of projects. The concept of "Project categories" allows projects to be grouped on the landing page by user defined categories, which then behave as subfolders on the landing page. Note that this only affects the display on the landing page for Phylobook. All projects remain in the PROJECT\_PATH directory.

After a project has been created, permissions for that project can be assigned for users within the admin module. Only administrators will see all projects.

13) <u>Setting Access Permissions</u>: Permissions are managed at the level of a project, with all project contents having the same permissions. Once created, users and groups can be assigned permissions within the "Object Permissions" for any given Project object. This is performed by clicking a Project in the list of Projects and then clicking "Object Permissions".

Select the user or group to which you want to assign permissions and click the "Manage user" or "Manage group" button.

Set the permissions for a user or group. Phylobook uses the default permissions provided by a custom permission object in Django. Of these permissions, only two matter to Phylobook: "Can change project" and "Can view project". If a user or group has "Can change project" permission, then they are able to have access to the full set of annotation tools and can edit and save changes. Users or groups with "Can view project" permission are only able to view the projects contents and have no ability to save any new information. Save any permission changes and then click "VIEW SITE" in the upper right corner to return to the main Phylobook site project list.

14) <u>Other configuration considerations:</u> Phylobook can have fairly large POST requests when dealing with large samples. If you are using a reverse proxy server it should be configured to accept POST requests of at least 2M.

## Using Phylobook - viewing and annotating trees:

#### *Initial viewing and navigation of the data:*

Assuming you have followed the installation and configuration steps above, you should have 2 projects within Phylobook – HIV\_DEMO and WA\_SARS\_CoVSpike. Log in to Phylobook and select the project HIV\_DEMO. You should see a display that looks like the image in **Figure 6**.

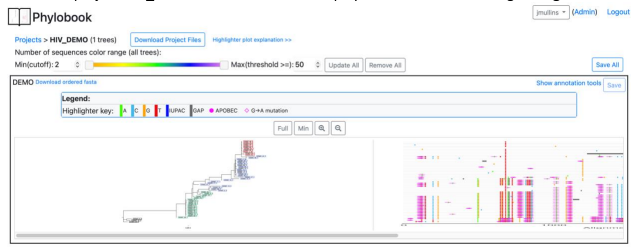


Figure 6. Initial view of the HIV\_DEMO project.

Clicking on the "Full" button will expand the view. Clicking on the magnifying glass icons will further increase or decrease the size of the view window. The upper left corner shows the current project being displayed and the number of trees within that project. If multiple trees are present in the project, multiple panels similar to that in **Figure 6** will be displayed. Each panel corresponds to a dataset within a project consisting of a set of sequences represented by a phylogenetic tree and a highlighter plot. The highlighter plot shows sequence variation relative to a master sequence. The master sequence is marked with a (m) just to the right of the sequence names in the highlighter plot. Highlighter plots are created using the Los Alamos National Laboratories (LANL) Highlighter tool found at

https://www.hiv.lanl.gov/content/sequence/HIGHLIGHT/highlighter\_top.html/.

Just to the right of the project name is the "Download Project Files" button. This will download all files associated with this project. This can be used to archive data at a given state of analysis or to share data with other Phylobook installations. The demo data provided through our GitHub site was downloaded from our own Phylobook server. Just to the right of the Download Project Files button is a link to LANL's page that explains the highlighter application.

Within the phylogenetic tree, sequence names may be surrounded by colored boxes that indicate sequence groupings, e.g., lineage designations. As will be discussed later, groupings can be edited within Phylobook and sequences within a group can be exported for further analysis.

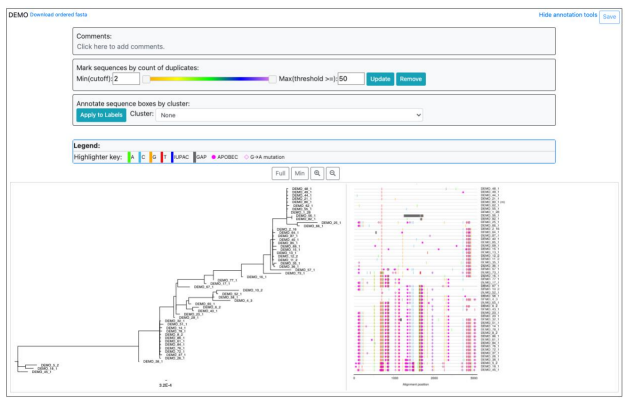


Figure 7. View of the HIV\_DEMO project after pressing the "Full" button and selecting "Show annotation tools".

#### **Annotations**

Annotation tools can be shown or hidden by clicking on the Show/Hide Annotation tools text in the upper right of the window (Figure 7). Trees can be labeled by abundance of sequences within the population, proposed lineage groups, information encoded within the sequence name (such as date or tissue type) and individual markers can be place on the tree and associated with a notes field. Annotation tool availability is context specific. For example, the "Annotate sequence boxes by cluster" tool will only appear when clustering data is available and the "Color sequence names by field" tool will only appear when the sequence names have the required fields available in the sequence names.

#### Labeling sequences by abundance within a population:

When sequences are produced from a variable population (HIV in this example), it is nonetheless common that many sequences are identical to each other. Our standard practice is to collapse identical sequences into one sequence prior to alignment and creation of the phylogenetic tree (). When this is done, the sequence names are modified to end with an underscore followed by the number of identical sequences that were collapsed into this particular sequence ID. In the DEMO\_HIV data, the 8<sup>th</sup> sequence down on the tree (DEMO\_1\_28) is the result of collapsing 28 identical sequences. The default in Phylobook is that all sequence names end with \_n where n is the number of sequences that were collapsed into each sequence ID. Near the top of the project page, below the Comments field (see **Figure 9** for tools available when the Comments field is selected), is a tool labeled "Mark sequences by count of duplicates:" with a slider that permits assignment of the color and number range to be

used. This tool is used to apply a colored square just to the left of each sequence name. The color of the square indicates the number of sequences that were collapsed. Using this tool, individual or all of the trees within a given project can be annotated with the same color range. **Figure 8** shows the DEMO tree after the sequences have been marked by abundance within the sample. Note that after this annotation was added, the sample is marked by a red border. This indicates that the newly created annotation has not yet been saved. Clicking on the save button will store the annotation and change the color of the border back to black. Annotations are stored within the SVG file that represents the tree.



**Figure 8. Sequences marked by abundance.** For each sequence in the tree that represents more than one sequence in the dataset, a small colored square has been placed to the left of the sequence name in the tree. The color of the square indicates the abundance of the sequence in the original dataset. Sequences with no square represent a single sequence in the dataset. A red border indicates that the changes have not yet been saved.

#### Lineage assignment using clustering data:

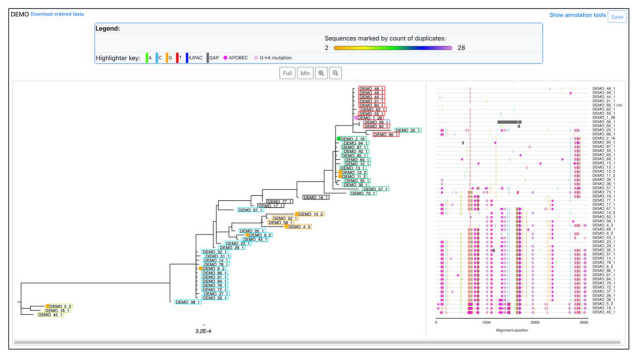
For the DEMO dataset, the "Annotate sequence boxes by clusters" is available since clustering data is present within the project directory. One of the primary applications of Phylobook is to create and store individual groups of sequences from within a dataset. These groupings (or "lineage" assignments) can be created in a semi-automated fashion with an external algorithm that assigns sequences to groupings or "clusters". When data is loaded into Phylobook, one or more initial lineage designations can be provided for each dataset. These consist of files in the dataset with names in the format of DataSetName.cluster.AlgorithmName. For example, in the DEMO dataset in the HIV\_DEMO project, there are files named DEMO.cluster.km2clusters,

DEMO.cluster.km3clusters, etc. Each of these is a text file that contains sequenceID, LineageID for the all of the sequences in the dataset (where lineage ID is an integer). In the DEMO example, the lineage designations of type "km#clusters" were created by running a kmedoid algorithm on the nucleotide distance matrix for all samples in the dataset. Kmedoid () was run with "K" = 2-10 and a lineage designation file was saved for each value of K. Clustering will be discussed in more detail later in the manual.

When lineage designation files are available, the "Annotate sequence boxes by clusters" tool will be available and the pulldown menu "Cluster" will contain the algorithm name (which is parsed from the file name). For example, **Figure 8** shows the pulldown menu with "km7clusters" selected. Selection of this algorithm annotates the tree with colored triangles to the right of each sequence name. The color of the triangle corresponds to an initial lineage designation. Clicking the "Apply to Labels" button will transfer these tentative lineage designations to boxes that correspond to saved lineage designations in Phylobook. **Figure 9** shows the result of this process.



**Figure 8. Using the "Annotating sequence boxes by cluster" tool.** In this example, the km7clusters algorithm has been selected and each sequence in the tree is labeled by a colored triangle at the end of the sequence name. The triangle color corresponds to the initial lineage designation.

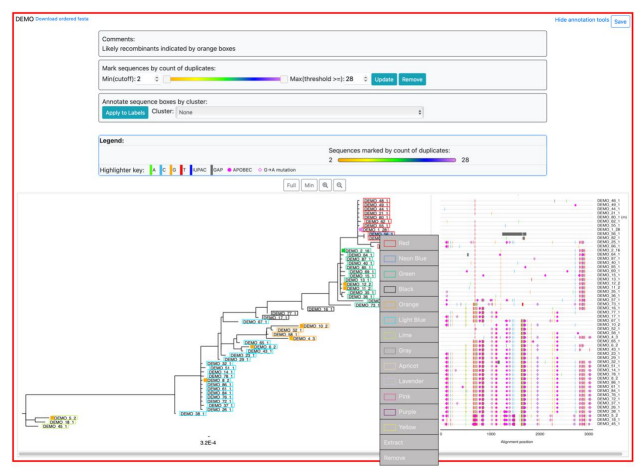


**Figure 9. DEMO data set after km7cluster designations have been Applied to Labels.** After clicking "Apply to Labels" clusters are indicated by the colored boxes around the sequence names in the tree. To simplify the image, the cluster designations (triangles in **Figure 8**) were then turned off by selecting "None" in the Cluster pulldown menu and the annotation tools were hidden.

After applying the K-medoid designations to the data, lineages are indicated by the colored boxes around the sequence names in the trees. For the most part, selecting km7 makes biological sense as the sequence groupings correlate mostly with what appears by eye to be separate lineages. For example, 9 of the first 10 sequences have been grouped into one lineage as indicated by the red boxes around the sequence names. One sequence (DEMO\_56\_1) has been left out of this grouping and inspection of the highlighter plot shows a large gap in this sequence relative to others in the grouping. A second grouping of sequences indicated by the green boxes appears to be a separate lineage but there are two sequences (DEMO\_25\_1 and DEMO\_66\_1) that appear to be recombinants. At the bottom of the tree, there are two unique lineages (lime green and light blue boxes). In between, there are a number of sequences that appear to be recombinants of the other lineages. In such complicated trees, automating lineage assignments via clustering (or other algorithms) is likely to result in some errors so Phylobook provides a mechanism for manual editing of lineages.

#### Manual editing of lineages:

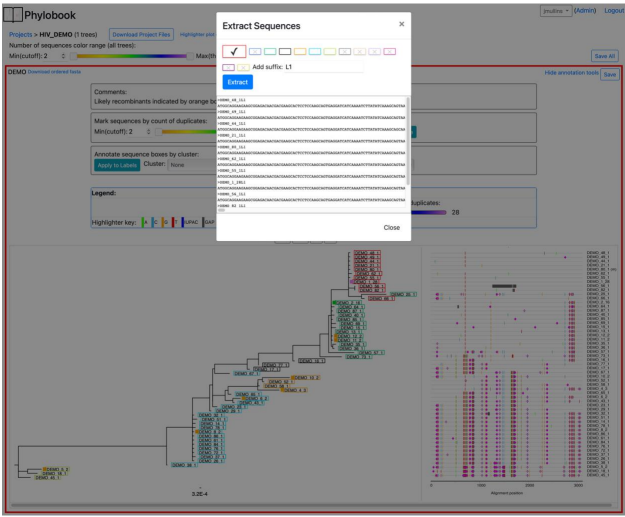
Lineages can be manually edited by hovering over a single sequence name or dragging the mouse to select multiple sequence names in the tree that one wishes to assign to a different lineage. **Figure 10** shows the menu that pops up when this is done. In this example DEMO\_56\_1 has been selected. By clicking on the red box, it will be assigned to the same lineage as the other sequences labeled by red boxes. **Figures 10** also shows the result of additional manual editing to place all likely recombinant sequences in a separate "lineage" designated by orange boxes, as noted in the Comments field.



**Figure 10. Manual editing of lineage designations**. Dragging the mouse around (a) sequence name(s) in the tree brings up a box that allows one to assign the selected names to a new or different lineage.

#### Exporting sequences in lineage groups:

After sequences have been assigned to lineage groups, they may be exported for downstream analysis. **Figure 10** shows the menu used to manually assign sequence groups to lineages. At the bottom of that same menu is a button labeled "Extract". Clicking on Extract pulls up the menu shown in **Figure 11**. Extracted sequences can be copied and pasted into another application for downstream analysis. This process will be automated in future updates to Phylobook.



**Figure 11. Menu for extracting sequences associated with selected lineages.** In this case, the "red" lineage was selected and the system was told to add "L1" (for lineage 1) to the sequence names. The extract button was clicked and the window was populated with the selected sequences that have been renamed as requested. These sequences can then be copied and pasted into another application for downstream analysis.

#### Annotation by fields encoded in the sequence names:

In some cases it is useful to annotate the tree to indicate associations with other parameters such as time, tissue source, partner relationships, etc. Phylobook provides for annotation using additional data encoded within the sequence name. In this annotation method, the font color of the sequence names in the tree is colored based on the values in an annotation field. As discussed above, by default we encode the number of identical sequences in a sample in the sequence name with \_n at the end of the sequence name where n is the number of sequences that were collapsed into each sequence in a dataset. Additional information can be encoded within the sequence names delimited by additional underscores.

Phylobook parses the sequence names in each dataset to identify fields that are delimited by underscores and counts the number of variants for each field. Phylobook allows the user to designate a font color for each variant. Since it is difficult to visually distinguish more than about

10 font colors, the feature is limited to annotations with 10 or fewer variants. For example, in the WA\_SARS-CoV2\_SpikeAA sample data set, the sequence names contain information about the date of sampling. In this data set, the sequence names have the form SequenceID\_x\_Year\_Month\_n. The Year field contains the values 2020, 2021, 2022 and 1 (1 is the Wuhan reference sequence which was collected in 2020) and the month field contains values that run from 1-12. In such a dataset, an additional annotation tool becomes active (**Figure 12**). Fields that can be used to color sequence filenames are highlighted in teal boxes.



Figure 12. Annotation by values encoded in the filename. When appropriate fields in the filename exist, Phylobook will identify fields that are delimited by underscores and identify the number of variants within each such field. In cases where the number of variants is ≤10, the annotation tool "Color sequence names by field:" will appear followed by a representation of a sequence name - MW813275\_FieldA\_FieldB\_3\_1 in this example. In this case FieldA and FieldB are highlighted with teal boxes indicating that either field can used to color the sequence filenames.

Clicking on an eligible field in the sequence name representation brings up a dialogue box similar to that shown in **Figure 13**. In this case, we selected the field year values (except for the reference sequence which is from 2020). We then used the pulldown menus in the dialog box to assign the black font to all sequences from 2020, the magenta font to all sequences from 2021 and the light blue font to all sequences from 2022. The net result of this color assignment is show in **figure 14**. As expected, sequences from 2022 (mostly Omicron variants) cluster together at the bottom of the tree while sequences from 2020 (mostly similar to the early Wuhan sequences) cluster near the top of the tree and 2021 sequences are mostly in the middle of the tree.

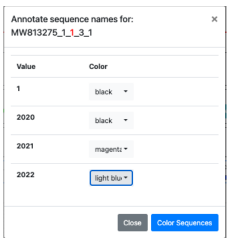


Figure 13. Dialogue box used to assign colors to sequence names based on a field in the sequence name.

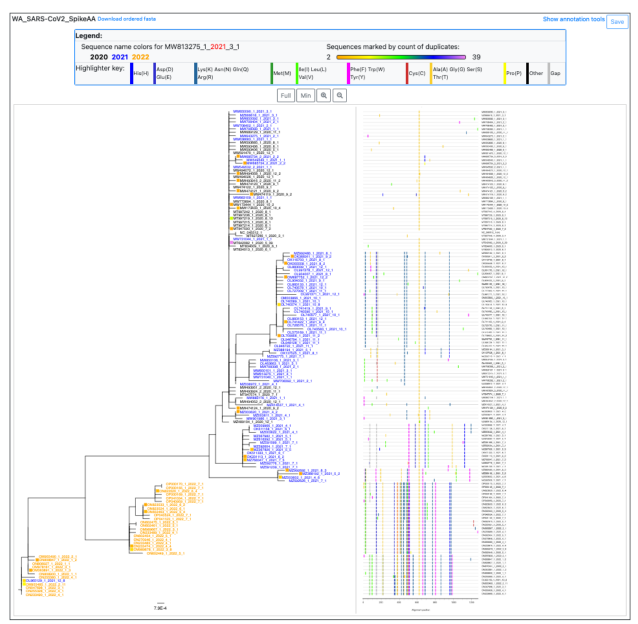
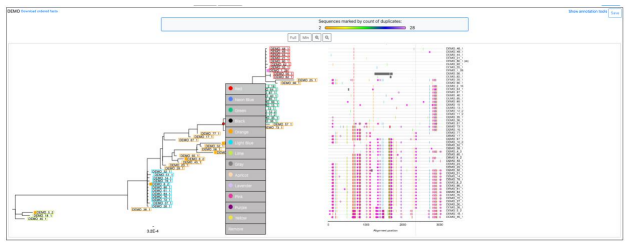


Figure 14. WA\_SARS\_CoV2\_SpikeAA sequences color coded by year. Black corresponds with sequences obtained in 2020, blue  $\rightarrow$  2021 and orange  $\rightarrow$  2022. Note: other annotations (lineage designations) present in the sample dataset have been removed in this diagram for clarity.

#### Annotations with colored dots:

Phylobook also allows the user to place small, colored dots at any location within the tree image. Control clicking within a blank region of the image brings up the annotation tool shown in **Figure 15**. The user can select a colored dot and drag to place it in any desired location on the tree. When combined with a comment, this can be useful to label specific branches within the tree or other features of interest.



**Figure 16**. **Annotation of the tree with colored dots.** Control clicking a blank region of the tree brings up the box shown above. The user can select a colored dot from within this box and place it anywhere on the tree by dragging it to a desired location.

### Preprocessing data and making it available within Phylobook:

Now that you have Phylobook installed and have familiarized yourself with its use, the next step is to populate it with your own data.

The Phylobook pipeline can be installed either directly on your computer or it can be installed within a Docker container. We strongly recommend you install it via a Docker container and will provide detailed instructions for that install below.

- 1. These instructions assume that you have already installed Docker and Git as described above.
- 2. Create a directory to house the pipeline.
- 3. In a terminal window, change to the desired directory and execute the command:

git clone <a href="https://github.com/MullinsLab/phylobook\_pipeline.git">https://github.com/MullinsLab/phylobook\_pipeline.git</a>

This will create a directory called "phylobook\_pipeline" in your designated directory.

Note: If you are installing on a Mac or a Unix operating system, you will then need to modify the docker-compose.yml file to have your userid and groupid. This will assure that files created by the pipeline will be associated with your userid and groupid. To make this modification first discover your userid and groupid by issuing the following terminal command

In the example output from this command below, the userid is 501 and the groupid is 20.

jamesmullins@iMac phylobook\_pipeline % id uid=501(jamesmullins) gid=20(staff) groups=20(staff),12(everyone),61(localaccounts),79(\_appserverusr),80(admin),81(\_appserveradm),98(\_lpadmin),33(\_appstore),100(\_lpoperator),204(\_developer),250(\_analyticsusers),395(com.apple.access\_ftp),398(com.apple.access\_screensharing),399(com.apple.access\_ssh),400(com.apple.access\_remote\_ae)

- 4. Using a text editor open the docker-compose.yml file and edit line 12 to delete the # sign (making it no longer a comment) and change it to the correct user and group ID.
- 5. In the terminal, switch to the phylobook\_pipeline directory and build the new container using the command:

docker compose up -d --build

This should create the new docker container for the pipeline. In docker desktop, you should see a display similar to the image in **Figure 17**.

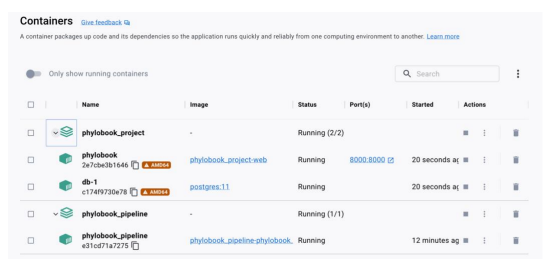


Figure 17. Docker container display after installation of the Phylobook pipeline.

6. With the pipeline running inside the docker container, issue the command:

docker exec -it phylobook\_pipeline bash

This will put you into a terminal inside of the docker and you can operate the pipleline from there (you may get a message similar to: "I have no

- name!@ce1806727fef:/phylobook\_pipeline\$)", which you can ignore). Control-d will return you to the command line on your local computer's OS.
- 7. The phylobook pipeline operates on a working directory that contains one or more sequence alignment fasta files. Those fasta files must be stored in a directory to which the pipeline has access. Under the default setup, the only portion of the host computer's file system that is visible within the Docker container is the phylobook\_pipeline directory (and subfolders therein). Anything that gets created inside the docker's phylobook\_pipeline directory will show up in the host machine. To facilitate this, the docker container includes a directory under the phylobook pipeline directory, pipeline data (Figure 18).

phylobook_pipeline	Today at 4:43 PM		Folder
> <u></u>	Today at 3:56 PM	**	Folder
.gitignore	Today at 3:56 PM	48 bytes	Document
docker-compose.yml	Today at 3:59 PM	352 bytes	YAML Document
□ Dockerfile	Today at 3:56 PM	1 KB	Document
> iii figtree	Today at 3:56 PM		Folder
license	Today at 3:56 PM	2 KB	Document
> phyml	Today at 4:13 PM		Folder
✓ i pipeline_data	Today at 4:43 PM		Folder
WA_SARS-CoV2_SpikeAA.fasta	Mar 1, 2023 at 12:33 PM	185 KB	TextEdit
policy.xml	Today at 3:56 PM	5 KB	XML Document
README.md	Today at 3:56 PM	5 KB	Markdocument
requirements.txt	Today at 3:56 PM	22 bytes	Plain Text
> script	Today at 3:56 PM		Folder

Figure 18. Contents of a Phylobook pipeline directory.

- 8. Next, you must populate the data folder with one or more fasta files that contain sequence alignments of your own data. A sample data file named WA\_SARS-CoV2\_SpikeAA.fasta is provided in the data directory, and can be found at https://github.com/MullinsLab/phylobook\_pipeline if it needs to be redownloaded later. Leave this file in the working directory, or replace it with one of your own.
- 9. Then issue the command:

python3 /phylobook pipeline/script/phylobook.py -d /phylobook pipeline/pipeline data -t aa

This command will run the script phylobook.py on the fasta files in /phylobook\_pipeline/pipeline\_data. Note that in this example, there was only a single fasta file on which to operate and it contained an amino acid sequence alignment. If the directory contains (a) nucleic acid sequence alignment(s), then the command should end with nt instead of aa.

10. After the pipeline has run, the folder will be populated with all of the required files for phylobook. In addition, a number of intermediate results files and a log will be created (see **Figure 3**). These files can now be moved to a project directory folder in phylobook.

run_phyml_logs	Today at 5:20 PM		Folder
WA_SARS-CoV2_SpikeAA_highlighter_untrimmed.png	Today at 5:20 PM	787 KB	PNG image
WA_SARS-CoV2_SpikeAA_highlighter.fasta	Today at 5:20 PM	187 KB	TextEdit
WA_SARS-CoV2_SpikeAA_highlighter.png	Today at 5:20 PM	768 KB	PNG image
WA_SARS-CoV2_SpikeAA_highlighter.txt	Today at 5:20 PM	14 KB	Plain Text
WA_SARS-CoV2_SpikeAA.fasta	Today at 5:18 PM	183 KB	TextEdit
WA_SARS-CoV2_SpikeAA.phy	Today at 5:18 PM	183 KB	TextEdit
WA_SARS-CoV2_SpikeAA.phy_log.txt	Today at 5:19 PM	292 bytes	Plain Text
WA_SARS-CoV2_SpikeAA.phy_phyml_stats.txt	Today at 5:19 PM	2 KB	Plain Text
WA_SARS-CoV2_SpikeAA.phy_phyml_tree.txt	Today at 5:19 PM	6 KB	Plain Text
WA_SARS-CoV2_SpikeAA.phy_phyml_tree.txt_newick.tre	Today at 5:19 PM	5 KB	Dendroocume
WA_SARS-CoV2_SpikeAA.phy_phyml_tree.txt_nexus.tre	Today at 5:19 PM	11 KB	Dendroocume
WA_SARS-CoV2_SpikeAA.phy_phyml_tree.txt.svg	Today at 5:19 PM	67 KB	Scalabls Image
WA_SARS-CoV2_SpikeAA.phy_phyml.txt	Today at 5:19 PM	494 KB	Plain Text
WA_SARS-CoV2_SpikeAA.phy_pwcoldist.txt	Today at 5:19 PM	488 KB	Plain Text

#### 11. Notes:

- a. The pipeline will operate on all the files in the pipeline data folder sequentially. It assumes that all files are either nucleotide alignments OR amino acid alignments. I cannot handle a mixture of both. It also assumes that ALL files are alignment files. So, after running the pipeline, you must move the files out of the pipeline data directory prior to doing another run.
- b. At present, the pipeline <u>does not</u> handle fasta files that have been wrapped at 80 characters. Please assure your files are appropriately formatted.
- c. The highlighter plot uses the first sequence in the fasta alignment file as the reference sequence. E.g. sequence variation relative to the FIRST sequence in the file is shown in the highlighter plot. If the sequences in the fasta file have been created by collapsing identical sequences into representative sequences, ourstandard practice has been to put the most abundant sequence first in the fasta file.
- d. Several files that are not needed by Phylobook are also produced by the pipeline (for example the distance matrix, the newick tree file etc). We typically store these files in the Phylobook project directory as we occasionally find them useful for other analyses.