tinselR – An RShiny Application for Annotating Outbreak Trees

Running Title: tinselR for annotating phylogenetic trees Jennafer A. P. Hamlincurrent,1,2,# Teofil Nakov3, and Amanda Williams Newkirk2

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### ABSTRACT

Across the United States, public health laboratories perform whole-genome sequencing for many pathogens. This high-resolution data determines the relationships between isolates via phylogenetics. In combination with other epidemiological data, epidemiologists use this data to inform investigations to confirm an outbreak and identify potential transmission routes. Our goal was to develop an open-source user-friendly graphical user interface (GUI) for phylogenetic tree visualization and annotation. Here, we present tinselR for that purpose and which is availble at [https://github.com/jennahamlin/tinselR.](https://github.com/jennahamlin/tinselR)

### ANNOUNCEMENT

Given that the R programming language contains some of the gold standard packages for phylogenetic analyses and visualization (e.g., ape (Paradis, Claude, and Strimmer 2004), and ggtree (Yu et al. 2017)), we used the Rshiny framework (Chang et al. 2017) to develop **tinselR** (pronounced tinsel-er) to provide GUI access to the tools in ape, ggtree, and other vital packages. tinselR’s minimum input requirement is a Newick formatted phylogenetic tree. Once loaded, user-selected inputs change the appearance of the displayed tree. For example, a user can quickly transform tip label formatting. By adding a genetic distance matrix or metadata file or both, the user can include annotations on the image, relabel tips, or add a heatmap to the phylogenetic tree. These modified tree images are downloadable in various formats (pdf, png, or tiff) for presentations, publications, or other communications with collaborators. Below we detail how to install the application and describe the example data pre-loaded so that new users can familiarize themselves with the application.

The genetic distance matrix file must contain a square matrix of single nucleotide polymorphism (SNP) differences between the tree tips. The metadata file is a table of additional information to be changed or displayed on the tree. The tip labels in the Newick tree, distance matrix, and metadata files must match before upload, or tinselR will report an error. The primary function of the metadata file is to relabel the tips on the tree image. The header of the first column must be Tip.labels, and it must contain the labels for all tree tips in the uploaded Newick file. The alternative identification labels can be provided in the metadata file using the column header Display.labels in column two. If desired, users may include additional columns in the metadata file, such as the collection site, and display the information in a heatmap next to the tree. Headers for these other columns in the metadata file are flexible because they are not automatically recognized and used by tinselR. Acceptable formats include CSV, TSV, and TXT for the genetic distance and metadata files. Users can set file types independently for each input.

### INSTALLATION AND EXAMPLE DATA

To install tinselR from GitHub, users will need to install the R package devtools (Wickham and Chang 2016). The R packages ggtree (Yu et al. 2017) and treeio (Wang et al. 2020) is also required and can be installed from Bioconductor using BiocManager (Morgan 2019). With the installation of these dependencies, tinselR is installable via the install\_github command from devtools. Explicit installation commands are below (Figure 1), and the final command (run\_app()) will launch the application. Note that install\_github will also install other missing R dependencies. tinselR will accept Newick tree files from any program, e.g., RAxML (Stamatakis 2014), as input. Although it is possible to host ShinyR applications on a server, to date tinselR has only been tested by single users running the application locally. We recommend testing to ensure tinselR performs as expected under multi-user conditions before providing access from a server for production purposes.

After launching tinselR, new users can explore the application using one of the pre-loaded datasets located in the ‘Example Data’ tab. We provide three datasets (i.e., Newick formatted tree, genetic distance matrix, and metadata file). These data are either *Escherichia coli* (from NCBI Bioproject: PRJNA218110) or *Salmonella enterica* (from NCBI Bioproject: PRJNA230403) with the number of isolates ranging from 14 - 19. The genomic data used in the example data sets were generated and used under the CDC IRB protocol 7172. After clicking on the ‘Example Data’ tab, users can select one of the datasets (e.g., example data 1, 2, and 3) from the drop-down menu. We highlight the capabilities of tinselR (Figure 1) using example data 1 below. Run the below code in your R console -

**1). Install devtools package**

install.packages("devtools", dep=T)

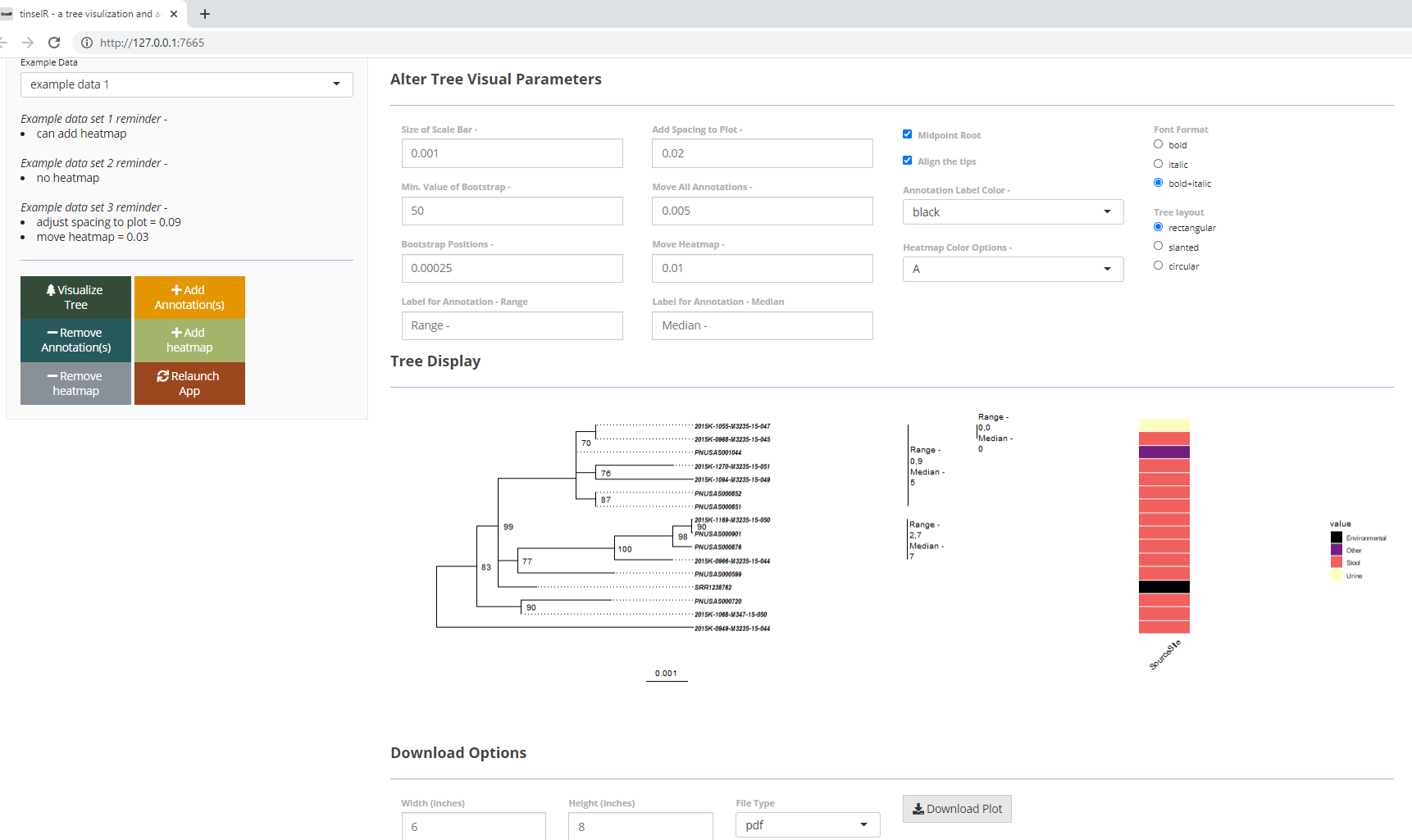
**2). Install ggtree and treeio**

if (!requireNamespace("BiocManager", quietly = TRUE))  
install.packages("BiocManager")  
BiocManager::install("ggtree")

Note that installing ggtree will also install treeio

**3). Install and launch the tinselR shiny application**

devtools::install\_github("jennahamlin/tinselR")  
library(tinselR)  
run\_app()



Example dataset 1 displayed with annotations and a heatmap indicating collection source.

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