

# **SplitsTree App User Manual**

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SplitsTree App (version 6.6.5, built 25 Nov 2025)



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# Introduction

The SplitsTree App is new software for exploring and analyzing phylogenetic data, with an emphasis on phylogenetic networks. Offering a comprehensive set of features, the software provides over 100 algorithms for computing distances, phylogenetic trees, split networks, haplotype networks, rooted phylogenetic networks, tanglegrams, consensus trees and consensus networks.

This new software [Huson and Bryant, 2024] is designed to accommodate the increasing scale and intricacy of modern data sets. It extends, integrates and supersedes our earlier applications SplitsTree4 [Huson and Bryant, 2006] for unrooted phylogenetic trees and networks, Dendroscope3 [D. H. Huson, 2012] for rooted trees and networks, and PopArt [Leigh and Bryant, 2015] for haplotype analysis.

If you use this program, the please cite:

Daniel H. Huson and David Bryant. The SplitsTree App: interactive analysis and visualization using phylogenetic trees and networks. Nature Methods (2024) <https://doi.org/10.1038/s41592-024-02406-3>.

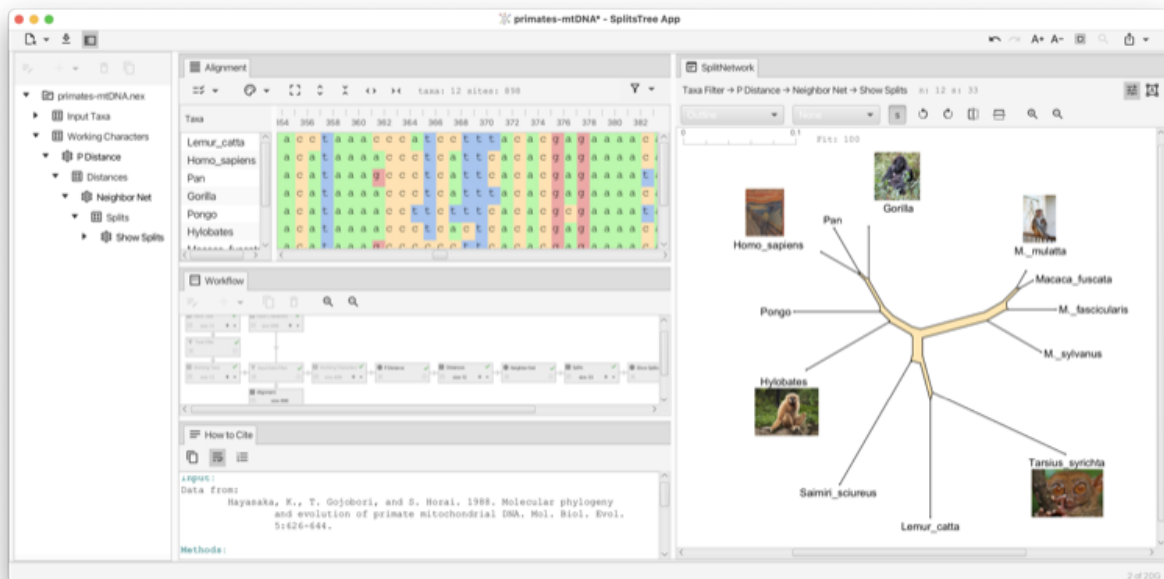


Figure 1: Example of SplitsTree analysis of primate mtDNA.

# Chapter 1

## Using SplitsTree

In this chapter we give an overview of how the interface of the SplitsTree app is organised. Later we provide more details on the actual methods and procedures.

### 1.1 Getting started

To get started using this program, download the latest installer from <https://software-ab.cs.uni-tuebingen.de/download/splitstree6> for Linux, Mac OS X or Windows, and install the program on your computer. Versions for iOS and Android are being tested.

Launch the program by double-clicking the program icon or launch it from the command line (Linux).

Use the File->Open... menu item to open a file containing data in one of the supported formats (see Chapter D).

If the data you provide is set of characters (or multiple sequence alignment), then by default, the SplitsTree app will compute P-distances (see Section C.1) and then will run Neighbor Net (see Section C.2) to obtain a split network (see Section 1.10). If you provide a distance matrix, then this will also result in a split network being displayed. If you supply trees, then the first tree will be displayed (see Section 1.6).

Here is a toy example of characters data. You can copy this text from the manual and then paste it onto the import data button (see Section 1.3) to obtain the network show in the figure (see Fig. 1.1).

6 64

```
Taxon1 TAAGTAGATCGGAGTTTTACTCGTGTGATTTTGGGTATTTTTATTTAGATTATGAAATTATA
Taxon2 CTTAATATATAATGATATTACTTAAACATTATTAATGATACACTAACTATAATTATTGAACAT
Taxon3 AAAATTATATAATATAACATTATATTCATTACCACAAGATTATATTATAAAATATATTGTACAC
Taxon4 TTCAATATATAATGAAACTTTATAAATAACTTTAGAAATCTTATAAAAAAATATCGACGAACAA
Taxon5 TAAGTAGATCTGAGTTTTTACTCGTGTGATTTTGGGTATTTTTATTTAGATTATGAAATTATA
Taxon6 TAAATTGGATAATTTTTATTATGTGTTACTACAGAAATCATTAATTATATAAACGATATACAA
```

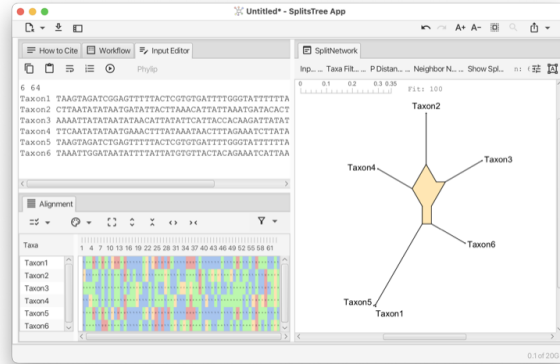


Figure 1.1: Neighbor net computed for toy characters data.

## 1.2 Layout of the main window

In the SplitsTree app, you can open one or more documents and each one has its own main window. Different analyses of the same data in the same document are shown in different tabs in that window (see Fig. 1.2). Tabs can be laid out side-by-side.

Each document opened in SplitsTree has its own main window. The main window has the following parts:

- A menu bar providing menu items to access features of the program. Note that all features of the program can also be accessed from within the main window as well (see Fig. 1.2) (figure part g).
- A toolbar containing both document- and tab-specific items (figure parts e,f).
- A main tab pane that contains all the text and data tabs (figure part a).
- A sidebar providing access to the workflow (figure parts b,c).
- An algorithms tab pane (inside the sidebar) for parameterizing and running algorithms (figure parts b,d).

## 1.3 Main toolbar items

There are three *document-specific toolbar items* on the left side of the main toolbar and several *tab-specific toolbar items* ones on the right (see Fig. 1.2) (see Fig. 1.3).

Document specific toolbar items:

- The *files menu button* - provides a menu that contains file-related items from the main files menu and a list of recently opened documents.
- The *import data button* - use this to open any string or file from the system clipboard, or via drag-and-drop, into a new SplitsTree document.
- The *sidebar toggle button* - use this to show or hide the sidebar.

Tab specific toolbar items (that apply only to the currently selected main tab):

- An *undo button* and *redo button*.



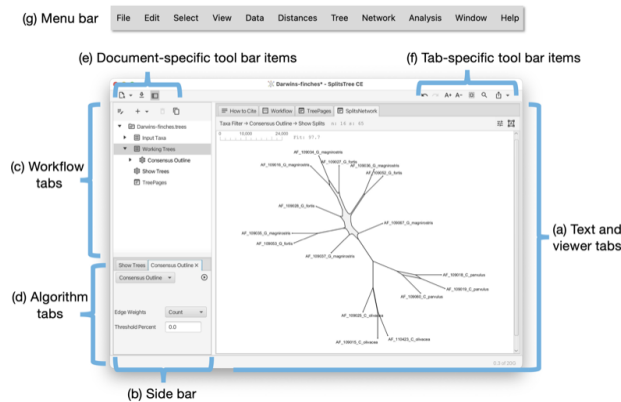


Figure 1.2: The main window. (a) Text and graphic output are presented in the main tabs on the right-hand side of the window. (b) There is a side-bar on the left hand side that shows (c) the workflow at the top and (d) tabs for setting algorithm parameters at the bottom. At the top left (e) there are some document-specific tool-bar items whereas the items at the top right (f) apply to the current main tab. While all program features can be directly accessed from the main window, the menu bar (g) provides alternative access to many of the features.

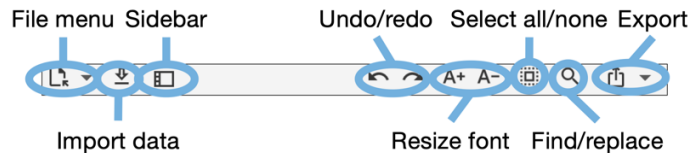


Figure 1.3: The main toolbar has three document-specific items on the left: Files menu button, import data button and sidebar toggle button. It has several tab-specific ones on the right: Undo and redo, font resizing, select all/none, find/replace and export.

- An *increase font size button* and *increase font size button*.
- A *selection button* to select all or none.
- A *find button* that you press to open the find dialog, press again to open the replace dialog (if available) and once again to close the dialog.
- An *export button* that gives access to items for copying, exporting or printing the data or image associated with the current tab. An additional item is provided for showing or hiding the *QR code* associated with a given tree or network.

## 1.4 The main tabs

The main window uses a tabbed pane to present all text and visualizations (see Fig. 1.2). Here is an overview of the supported tabs:

- **Alignment tab** - provides a visualization of the input multiple sequence alignment.
- **Tree-View tab** - shows a phylogenetic tree or rooted network.
- **Tree-Pages tab** - shows pages of phylogenetic trees or rooted networks.
- **Tanglegram tab** - shows a tanglegram of trees or rooted networks [Scornavacca et al., 2011].

- Densi-Tree tab - shows a densi-tree visualization of a profile of trees [Bouckaert, 2010].
- Split-Network tab - show a split network [Dress and Huson, 2004] or phylogenetic outline [Bagci et al., 2021].
- Network tab - shows a network such as a haplotype network [Bandelt et al., 1999].
- World map tab - shows a map of the world and displays traits data that have associated and latitude and longitude coordinates.
- Workflow tab - provides access to the workflow graph.
- How to cite tab - provides a description of the data and algorithms used, and provides the necessary citations.
- Input editor tab - provides an interactive editor for entering and parsing input data.
- Report tabs - these are used to present the results of analyses such as phylogenetic diversity.
- Text tabs - any of the data blocks can be displayed in such a text tab, they provide several different formats.

## 1.5 Alignment tab

The *Alignment tab* provides a visualization of the input characters or multiple sequence alignment (see Fig. 1.4).

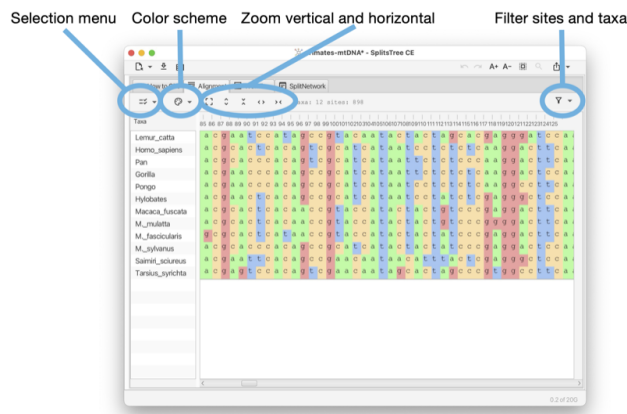


Figure 1.4: The alignment tab for displaying and working with a multiple sequence alignment.

The alignment tab has a drop-down menu button at the left that contains items for selecting sites. There is a button for selecting a color scheme. There is a button for toggling between a close-up view and a total view of the alignment, and buttons for zooming in and out both vertically and horizontally.

## 1.6 Tree-View tab

The *Tree-View tab* shows a phylogenetic tree or a rooted network (see Fig. 1.5).

The tree view tab has a toolbar and side panel that are hidden by default, but can be opened using two toggle buttons at the top right of the tab.

The toolbar provides items for selecting how to draw the tree (or rooted network), the choices are between rectangular, circular and radial cladogram or phylogram. There is a button that toggles the scale bar and addition

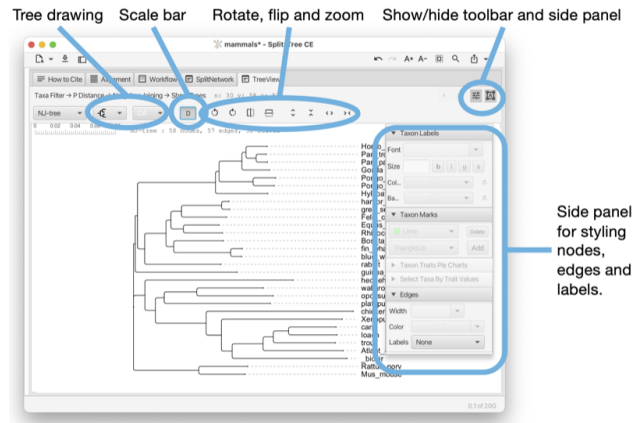


Figure 1.5: The tree-view tab for drawing a phylogenetic tree or rooted network.

information (such as name of the tree, if any, and number of nodes, edges and leaves). In addition, there are buttons for rotating, flipping and zooming.

The side panel (on the right) contains items for styling the taxon labels and for adding marks to the taxa. These are colored shapes that appear next to the taxon label. There are items for displaying taxon traits, if present. Moreover, there are items for setting the line width of edges and to choose whether to label edges by associated weights or confidence values.

## 1.7 Tree-Pages tab

The *Tree-Pages* tab shows pages of phylogenetic trees (see Fig. 1.6).

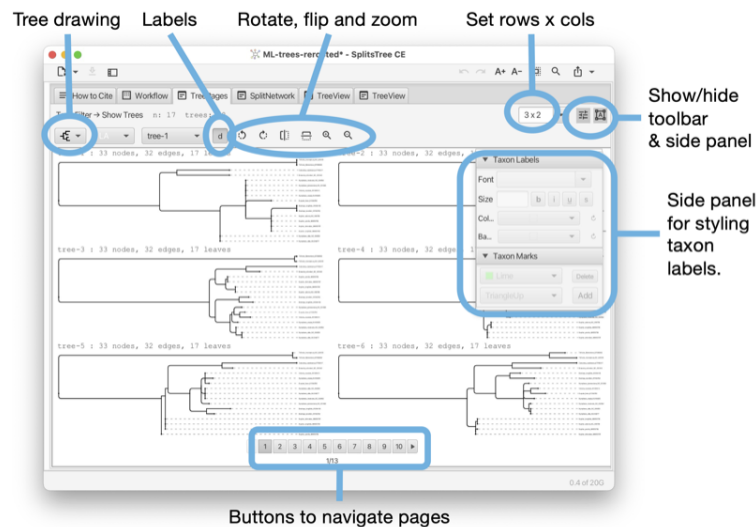


Figure 1.6: The tree-pages tab for displaying a collection of phylogenetic trees or rooted networks.

The tree-pages tab has a toolbar and side panel that are hidden by default, but can be opened using two toggle buttons at the top right of the tab.

The toolbar provides items for selecting how to draw the tree (or rooted network), the choices are between rectangular, circular and radial cladogram or phylogram. There is a button that toggles addition information (such as name of the tree, if any, and number of nodes, edges and leaves). In addition, there are buttons for rotating, flipping and zooming. At the right side of the toolbar, there is a text input field for setting the dimensions of a page in the format *rows x cols*.

The side panel contains items for styling the taxon labels and for adding marks to the taxa.

At the bottom of the tree-pages tab there is a row of buttons that can be used to navigate through the pages.

## 1.8 Tanglegram tab

The *Tanglegram tab* shows a tanglegram of trees or rooted networks, which is calculated based on the objective of displacement optimization [Huson, 2025] (see Fig. 1.7).

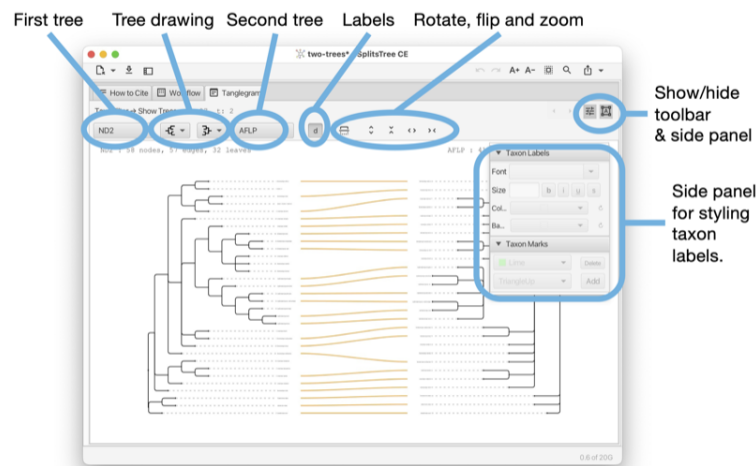


Figure 1.7: The tanglegram tab for comparing two phylogenetic trees or rooted networks.

The tanglegram tab has a toolbar and side panel that are hidden by default, but can be opened using two toggle buttons at the top right of the tab.

The toolbar provides items to determine the first and second trees (from the same file) and for selecting how to draw either tree (or rooted network), the choices are between rectangular phylogram, rectangular cladogram and triangular cladogram (trees only). There is a button that toggles addition information (such as name of the tree, if any, and number of nodes, edges and leaves). There are buttons for rotating, flipping and zooming.

There is a menu that allows one to select precisely what the target of displacement optimization should be, namely taxa or reticulate edges in which of the two phylogenies. In addition, if one of the two input phylogenies has a very unresolved node, that using the NeighborNet-presorting heuristic may help to untangle that situation (but only when both sides are being untangled.)

The side panel contains items for styling the taxon labels and for adding marks to the taxa.

## 1.9 DensiTree tab

The *Densi-Tree* tab shows a densi-tree visualization of a Bayesian profile of trees [Bouckaert, 2010] (see Fig. 1.8).

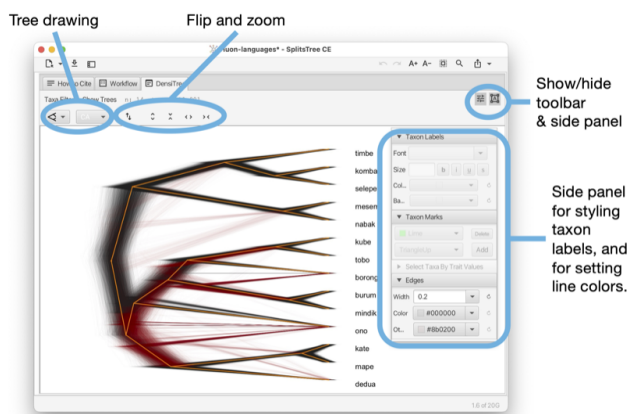


Figure 1.8: The densi-tree tab for displaying a Bayesian profile of phylogenetic trees.

The densi-tree tab has a toolbar and side panel that are hidden by default, but can be opened using two toggle buttons at the top right of the tab.

The toolbar provides items for determining how to draw the trees, choices are between rectangular, triangular, rounded and radial phylogram. (The rounded phylograms are time-consuming to draw and can cause problems.) In addition, there are buttons for rotating, flipping and zooming.

The side panel contains items for styling the taxon labels and for adding marks to the taxa. The line width can be set here. In addition, two colors can be set. The first is used for edges that in the tree profile that are compatible with the displayed greedy consensus tree (default color is black), and the second is used for incompatible edges (default color is red).

## 1.10 Split-Network tab

The *Split-Network* tab shows a split network [Dress and Huson, 2004] or phylogenetic outline [Bagci et al., 2021] (see Fig. 1.9).

To reshape the layout of the network by rotating the edges associated with one or more selected splits, press-and-drag on the network.

The split-network tab has a toolbar and side panel that are hidden by default, but can be opened using two toggle buttons at the top right of the tab.

The toolbar provides a choice box to determine how to draw the splits. The choices are as a split network, to-scale or as a topology (with all edges of uniform length), or as a phylogenetic outline, again, either to-scale or as a topology (with all edges of uniform length). There is a second choice box to determine whether the network is to be drawn unrooted or rooted, using either *mid-point rooting* or *outgroup rooting*. The latter requires that some taxa have been selected; these are treated as the outgroup.

In addition, there are buttons for rotating, flipping, zooming and for setting the *scale ratio* to a specific value to ensure that different networks are drawn to the same scale. Use the rotate buttons to rotate the entire network.

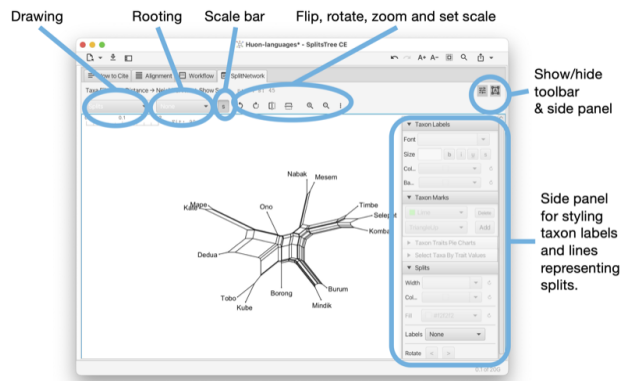


Figure 1.9: The split-network tab for displaying a collection of splits as a split network or phylogenetic outline.

The right side panel contains items for styling the taxon labels and for adding marks to the taxa. The line width can be set here. In addition, the line width and color can be set. The color of the inner area of an outline can be set. You can request to have the splits labeled by their weight, their confidence values (if available) or their internal split ids.

## 1.11 Network tab

The *Network tab* shows a network (see Fig. 1.10). The network tab has a toolbar and side panel that are hidden by default, but can be opened using two toggle buttons at the top right of the tab.

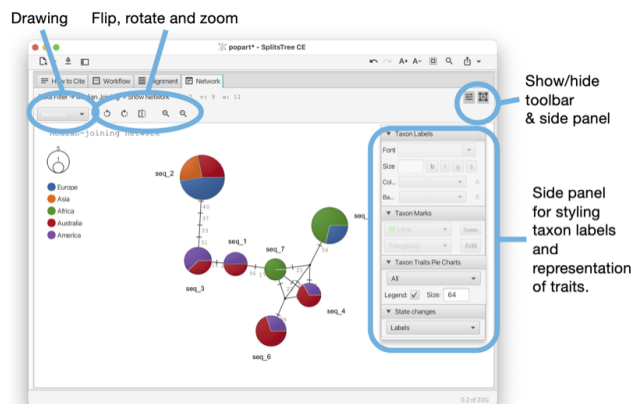


Figure 1.10: The network tab for displaying a haplotype networks and related on constructs.

The toolbar provides a choice box to determine how to draw the network. In addition, there are buttons for rotating, flipping and zooming.

The side panel contains items for styling the taxon labels and for adding marks to the taxa. The line width can be set here. In addition, there are items to determine which traits are to be shown in pie charts and whether to a legend. To change the colors used in pie charts, press on the items in the legend. These are then stored in the traits block in the TRAITCOLOR entry. Also, there is a menu button for determining how to represent character-state changes along an edge. The choices are has hatches (short marks), labels, compact labels and counts.

## 1.12 World map tab

The *World Map tab* shows a map of the world and places any traits data that comes with latitude and longitude assignments on the map (see Fig. 1.11). To change the colors used in pie charts, press on the items in the legend. They are stored in the traits block in the TAXONCOLOR entry. The world tab has a toolbar and side panel, the latter is hidden by default, but can be opened using the toggle button at the top right of the tab.

This tab appears when the input data contains a traits block that has latitude and longitude specifications (see Section B.2).

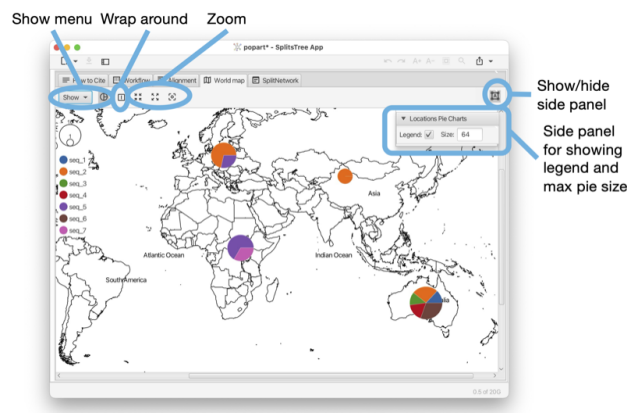


Figure 1.11: The world map tab for displaying haplotype locations of origin.

There is a *Show* menu button to determine whether country names, continent names and/or oceans should appear as labels. There is a button to determine whether to show two copies of the map side-by-side for Pacific-centric data. There is a button to zoom to the shown haplotype data.

## 1.13 Workflow graph tab

The *workflow tab* provides access to the *workflow graph* (see Fig. 1.12).

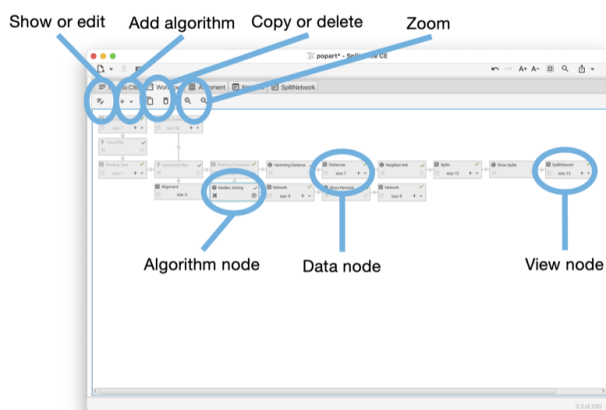
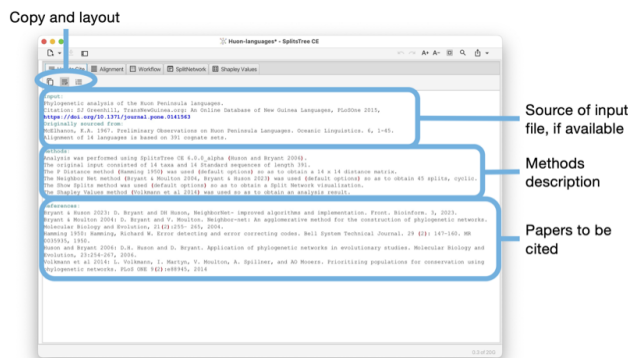


Figure 1.12: The workflow tab provides access to the workflow tab, for advanced users.

There are two items for zooming in and out.

The *How to cite* tab provides a description of the data and algorithms used, and provides the necessary citations (see Fig. 1.13).



If the input is a Nexus or SplitsTree file that contains a comment at the beginning of the file describing the source of the data, then this will be reported at the top of the text area. This is followed by a description of the methods used. Finally, all suggested references are listed.

The program will try to guess to which input format the entered text adheres to and will indicate the name of the format in the toolbar. When a valid format has been detected, then the run button will be enabled. Pressing the run button will parse the data and launch an analysis of the entered data.



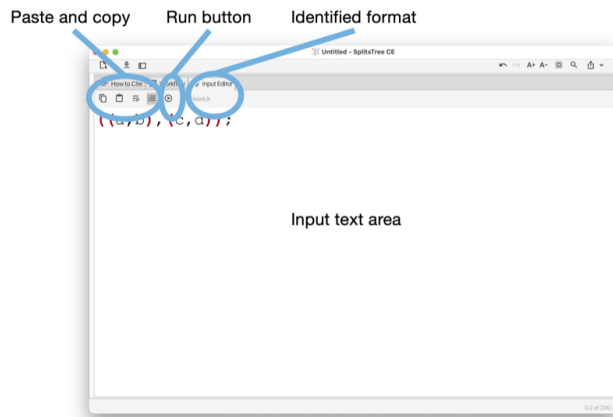


Figure 1.14: The editor tab is used to enter data into a new document. Pressing the run button will parse the data and launch an analysis of the entered data.

The input editor can be opened from the File menu and is automatically open when the user imports a text or file into the program that is not in one of the recognized input formats.

## 1.16 Report tabs

*Report tabs* are used to present the results of analyses such as Tajima' D, phylogenetic diversity or Shapely values as a text (see Fig. 1.15).

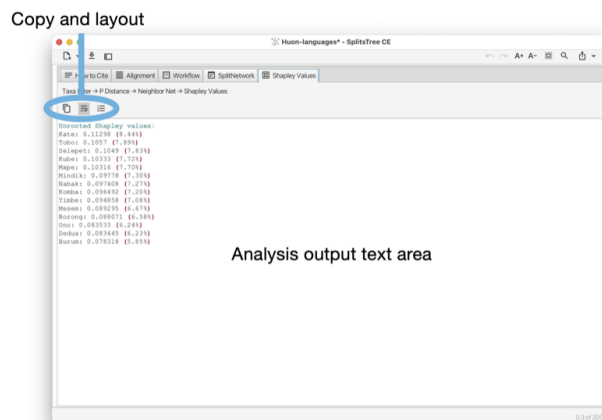


Figure 1.15: Report tabs are text tabs that are used to provide the result of an analysis, here the Shapely values for a set of taxa based on splits.

The toolbar of the report tab contains a button to copy the complete or selected content of the tab. There are buttons to turn line-wrapping and lines numbers on and off.

## 1.17 Text tabs

*Text tabs* are used to show the content of data blocks, in a choice of several different formats (see Fig. 1.16).

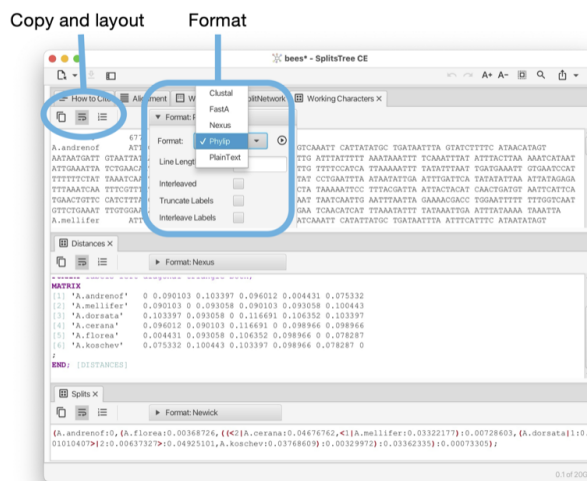


Figure 1.16: Text tabs are used to display the content of any of the data nodes in the workflow. Here we show three such tabs, one for characters data, one for distance data and one for splits data.

The toolbar of any text tab contains a button to copy the complete or selected content of the tab. There are buttons to turn line-wrapping and lines numbers on and off. There is a format pane that can be used to select the desired display format and to specify any options associated with the format.

Such a text tab can be opened by selecting a data node item in the sidebar and then pressing the show/edit button at the top of the sidebar, or by double-clicking on the item.

## 1.18 The sidebar

The sidebar (see Fig. 1.2) (figure part b) contains a representation of the workflow as a tree at the top, and the algorithms tab pane at the bottom (figure parts c,d).

The *workflow tree view* contains a representation of all input data, computed data and algorithms used in the computation. There are three types of nodes:

- *data nodes* that represent data blocks and result blocks,
- *algorithm nodes* that represent algorithms, and
- *view nodes* that represent visualizations.

Double-clicking on a data node will open a text tab displaying the corresponding data, or analysis result, if the data block is a report. Double-clicking on an algorithm node will open the corresponding algorithm tab.

An algorithm tab has a run button (at the right) to execute the algorithm and may contain some optional input items (below) to set parameters of the algorithm.

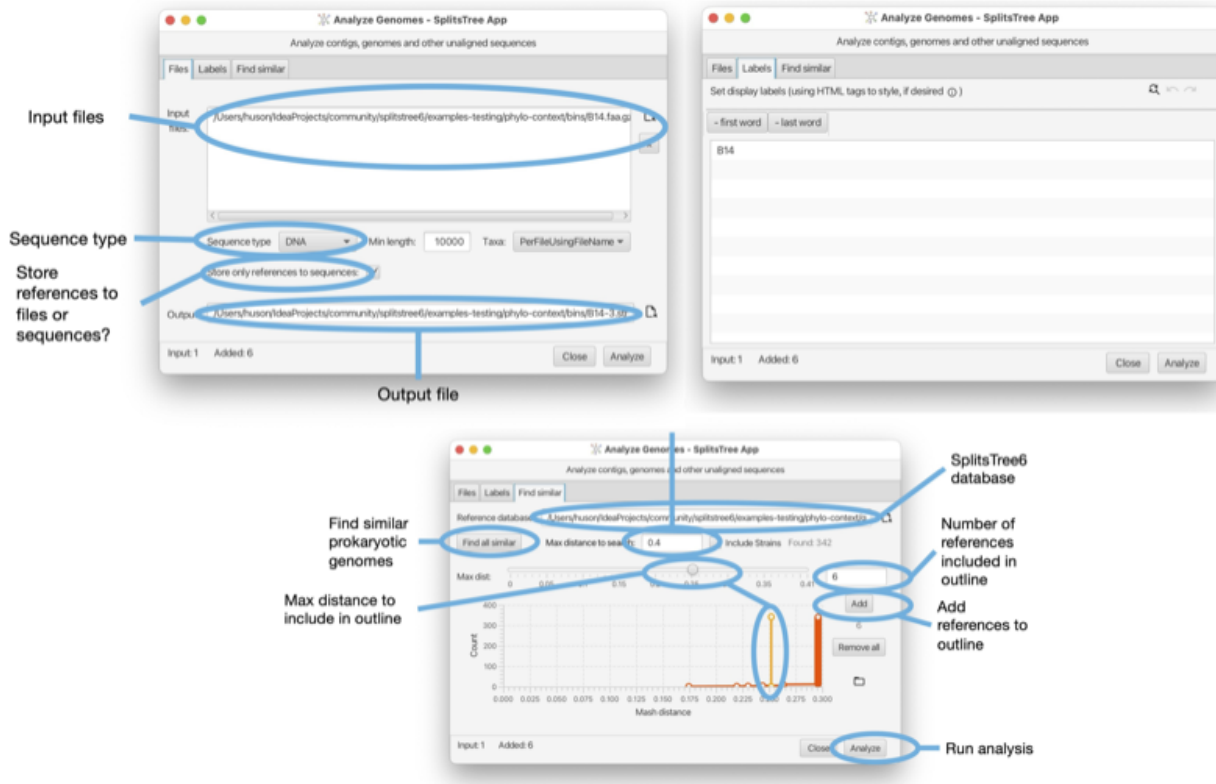


Figure 1.17: The first tab is used to specify the input files, the type of input (DNA or protein sequences) and whether to use files or FastA records as input genomes. Also, specify the output file (and whether to store input sequences as sequences or as references to files). The second tab is used to edit the labels of genomes. The third tab is used to specify the database to compare against (downloaded from the SplitsTree page), the distance to search in, and distance within which to include references.

## 1.19 The draft genome dialog

The SplitsTree App supports the calculation of the *phylogenetic context* of a draft prokaryotic genome [Bagci et al., 2021]. One or more files (FastA format) each containing one or more sequences representing draft genomes (or metagenomic assembly bins) can be imported into the program and then compared against a set of GTDB reference genomes [Parks et al., 2018] using mash distances [Ondov et al., 2016] and then represented as a phylogenetic outline.

The dialog is opened using the File->Analyze Draft Genomes... menu item and is set up using three tabs, as shown in the Figure (see Fig. 1.17).



## Chapter 2

# Building trees and networks

### 2.1 Using the workflow

In SplitsTree4, data analysis was based on a simple linear sequence. To construct a Neighbor Net, for example, one might input character data, apply a transform to infer a distance matrix, apply another transform to produce the set of splits in the Neighbor Net and another transform for convert those splits into a network on the screen.

That simplicity came with limitations. For example, to compare the result of analyses using different parameters or distance methods it was necessary to duplicate the whole file and start again.

The SplitsTree App implements a far more sophisticated system for workflows. It is still straightforward to run a simple linear workflow as in SplitsTree4, but it is now possible to branch that workflow at any point, exploring alternative parameters or methods. The use of frames make it easy to view the results of different analyses side-by-side.

The branching structure of a document's workflow can be viewed in the side panel (as a hierarchy) or in the workflow panel (as a graph). To illustrate, open the example file `ungulates.nex` which can be found in the directory `publications/WelkerEtal2015` in the Examples directory. By default, the SplitsTree App creates a network by running Neighbor Net and using the  $p$ -distance. Switching to the workflow panel displays the (linear) workflow for this initial analysis (see Fig. 2.1).

In this graph, nodes correspond to algorithms (indicated by a ⚙ icon) or data (indicated by an 📄 icon). From the algorithm nodes you can edit the parameters of the method. Selecting an algorithm node and clicking the delete button (top of pane) removes that node and any descendants of that node.

Suppose we want to compare a network computed by the Neighbor Net algorithm with a tree obtained using BioNJ. Assuming both are to be computed from the same distance matrix, we can select the corresponding node and choose BioNJ from the popup menu marked with a plus (either on the node or in the toolbar) (see Fig. 2.2). SplitsTree then constructs and displays the BioNJ tree. Switching back to the workflow panel we see that a new sequence of nodes has branched off the distances node, indicating the revised analysis.

This analysis also creates a new window tab. Right-click on a tab to get a context menu that allows you to split the main tab pane into two parts, then drag the tabs to the left or right panes to view both the Neighbor Net



Figure 2.1: Right hand side of the workflow created when `ungulates.nex` is opened.

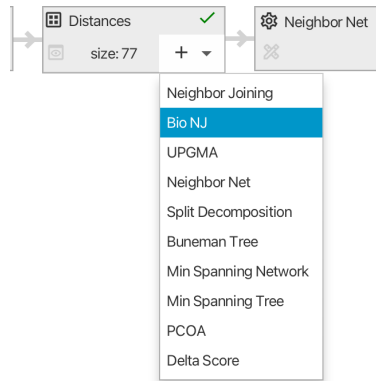


Figure 2.2: Attaching a BioNJ algorithm to an existing Distances block.

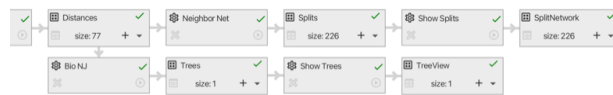


Figure 2.3: Right hand side of the workflow after adding a BioNJ analysis.

network and the BioNJ tree side-by-side.

## 2.2 Building trees

SplitsTree implements four standard tree construction methods:

- NJ (Neighbor-Joining), the original method of Saitou and Nei [Saitou and Nei, 1987].
- BioNJ, the modification of NJ introduced by Gascuel to reduce variance of the node-to-node estimates [Gascuel, 1997].
- UPGMA, the agglomerative method for constructing ultrametric (molecular clock) trees, introduced by Sokal and Sneath [Sokal and Michener, 1958].
- Buneman, a method for inferring compatible splits (and therefore trees) from distances which tends to produce trees with large multifurcations [Bandelt and Dress, 1992].

Each of these can be called from the Trees menu, or added as an algorithm in the workflow. There are several options for displaying trees, available by clicking on one of the two buttons on the right-hand-side of the tree window:



(see Section 1.6).

## 2.3 Neighbor Net and other split network methods

### 2.3.1 Neighbor Net

Given a distance matrix as input, the *Neighbor Net* algorithm operates in three stages. First, an agglomerative method is used to identify a circular ordering of the taxa. The splits computed by the algorithm are a subset of the set of all splits that can be formed from consecutive sets of taxa in that ordering. Second, a heavily customized algorithm is used to efficiently compute split weights. Those with zero weight are removed (use a split filter to remove splits with larger weight). Finally, a planar split network algorithm takes the weighted splits and produces the split network representation. A complete description of the entire process is available in Bryant and Huson [2023].

There is a single option available in Neighbor Net, the method used to infer split weights. We found that the Active Set method performed better than the other methods, and this is the default and recommended option. We have left the other algorithms as options in order to enable a repeat of the analysis in Bryant and Huson [2023].

When Neighbor Net is called, SplitsTree produces a split block and a split network block in the workflow. As we stress in Huson and Bryant [2006], the main information in the network is the set of weighted splits. Think of the network as a means of visualising the splits, in the sense that the same set of splits can be represented in several different ways.

### 2.3.2 Manipulating split networks

To rotate or flip the entire network, use the toolbar revealed by pressing the preferences button at the top right of the split network panel, making sure that none of the nodes or edges in the network are selected (see Section 1.10).

Click on an edge in the split network to select that split. The edges associated to that split can be rotated using the rotate buttons in the toolbar or the arrows in the side panel which appears when you click the button on the right (see Fig. 2.4).

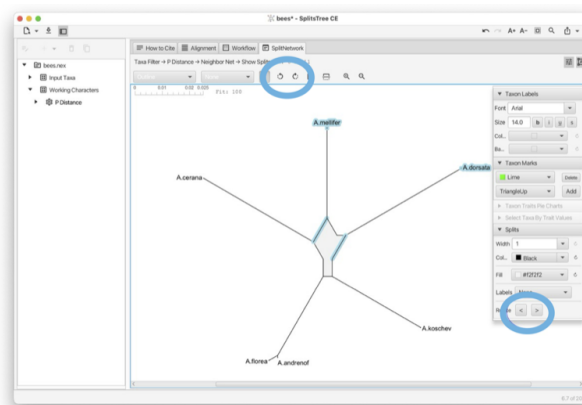


Figure 2.4: If one or more splits are selected, then highlighted buttons can be used to change the angles of the selected splits.

The traditional approach to displaying split networks marks out the splits with a mesh of quadrilaterals and polygons. The outline representation Bagci et al. [2021] constructs just the outer perimeter of the network. This is sufficient to represent all the split weights, and is generally much faster to compute and draw. To switch back

and forward between the graph mode and the outline mode use the pop-up menu at the left of the toolbar (see Section 1.10).

### 2.3.3 Split Decomposition

Given a distance matrix, the *Split Decomposition* method [Bandelt and Dress, 1992] can be selected in the Network menu, or on a distances node in the workflow. *Split Decomposition* is a predecessor of Neighbor Net, though the structures of the two methods are quite different. Split Decomposition works by inferring a set of splits satisfying a quartet condition in the distance matrix. Split Decomposition produces a set of weakly compatible splits and, as such, can produce more complex split networks than those produced by Neighbor Net. The resulting split network will not necessarily be planar. In practice, the conservative nature of the selection criteria means that Split Decomposition produces far fewer splits than Neighbor Net.

### 2.3.4 Splits in characters

SplitsTree includes several methods for extracting splits directly from character data. These methods do not assume any explicit model for sequence evolution. As such they do not correct for hidden mutations. However, they can reveal important structure within sequences from closely related organisms, as well as artefacts resulting from data handling problems.

The simplest is *BinaryToSplits* (see Section C.1), which applies to binary data only. Each binary character determines a split separating those with allele/state 0 and those with allele/state 1. The weight assigned to a split equals the summed weight for all characters inducing that split, defaulting to a count of those characters if weights are not specified. The *BinaryToSplits* algorithm is available via the workflow graph or workflow hierarchy. The user can specify a weight/count threshold on the splits, a cap on the maximum dimension of the split network (see Section C.3) and the option to include all 'trivial' splits separating one taxon from the remainder automatically.

The *DNAToSplits* method (see Section C.1) carries out a similar analysis but on nucleotide data. Splits are either determined via an RY coding (AG vs CT) or by splitting the most frequent state (assumed ancestral) from the other states (assumed derived) at each site.

The *Parsimony Splits* method (see Section C.1), introduced by Bandelt and Dress [1992], produces a set of weakly compatible splits directly from character taxa. The method is quartet based, like Split Decomposition, but for each four taxa, it determines the two most frequent pairings of two taxa versus the other two taxa.

## 2.4 Haplotype networks

A haplotype network is an elegant and efficient way to represent character or sequence data. Each node corresponds to a particular sequence with the size of the node proportional to the number of copies of that sequence in the data. Sequences which differ in one position are connected by an edge which is (optionally) labelled by the exact difference. Different methods for constructing haplotype networks generate different graphs for connecting sequences at larger distances. For them all, a key property is that given one sequence, the network, and the mutations along each edge, the entire alignment can be reconstructed.

SplitsTree provides implementations of two widely-used haplotype network methods, *MinSpanningNetwork* [Excoffier and Smouse, 1994] and *MedianJoining* [Bandelt et al., 1999]. Haplotype networks are drawn as graphs with each edge labelled by marks indicating the number of mutations/differences along that edge. This can be modified using the side panel which appears when clicking the preferences button at the top right of the network panel.



### 2.4.1 Minimum spanning network

A minimum spanning tree for a graph is a connected subgraph of minimum weight. Sometimes there is a unique minimum spanning tree; other times there are multiple.

In this context, the graph contains a node for each input sequence and edge between every pair of nodes. The length of each edge is the Hamming distance between the corresponding sequences. Other distance measures can be used, but the Hamming distance is appropriate for Haplotype Network construction.

The minimum spanning network is formed from all those edges in the graph which appear in every minimum spanning tree (see Section C.2).

A minimum spanning network is constructed from a characters block by first determining Hamming distances (right-click on the characters block and select Add Algorithm -> Hamming distance). Then right-click on the distance block produced and add the Min Spanning Network algorithm.

### 2.4.2 Median Joining network

Median Joining (see Section C.1) is probably the most highly-cited method for constructing phylogenetic networks. The implementation in SplitsTree is based on the method described in Bandelt et al. [1999]. The Median-Joining network method makes repeated use of minimum spanning networks, each time augmenting the set of observed sequences with putative ancestral sequences.

A Median-Joining network is constructed from a characters block via the Network menu, or by adding an algorithm to the workflow. The method comes with a single option  $\epsilon$  that is an integer controlling a threshold determining when two sequences are considered adjacent. In Bandelt et al. [1999],  $\epsilon$  varies between 0, 1 and 2.

## 2.5 Rooted phylogenetic networks

### 2.5.1 Implicit vs explicit trees and networks

A haplotype network is a direct representation of the input data and a split network represents groupings or splits between taxa. Both are examples of so-called *implicit* or *data-display* networks that aim at visualizing evolutionary data. In contrast, an *explicit network* is a representation of the putative evolutionary history, including reticulate events such as speciation-by-hybridization or horizontal gene transfer.

Strictly speaking, unrooted phylogenetic trees, too, are implicit representations of evolutionary data, whereas rooted phylogenetic trees have a direction (away from the root) and this allows branching nodes to be explicitly interpreted as representing speciation events.

Explicit phylogenetic networks are necessarily rooted. The Autumn algorithm [Huson and Linz, 2018] (see Section C.4) produces an explicit rooted phylogenetic network in which reticulations may be interpreted as putative hybridization or HGT events. However, just because a phylogenetic tree has a root does not mean that it is explicit. For example, the Cluster Network algorithm (see Section C.4) takes as input a set of rooted trees and aims at displaying all their clusters as a rooted network (in the hardwired sense [Huson et al., 2012]). Here, the reticulate nodes do not have a direct biological interpretation.

### 2.5.2 Hybridization networks

In mathematical phylogenetics, a *hybridization network* is a rooted phylogenetic network that contains or displays an input set of rooted phylogenetic trees. Usually, the requirement is that such a network minimizes the “hybridization

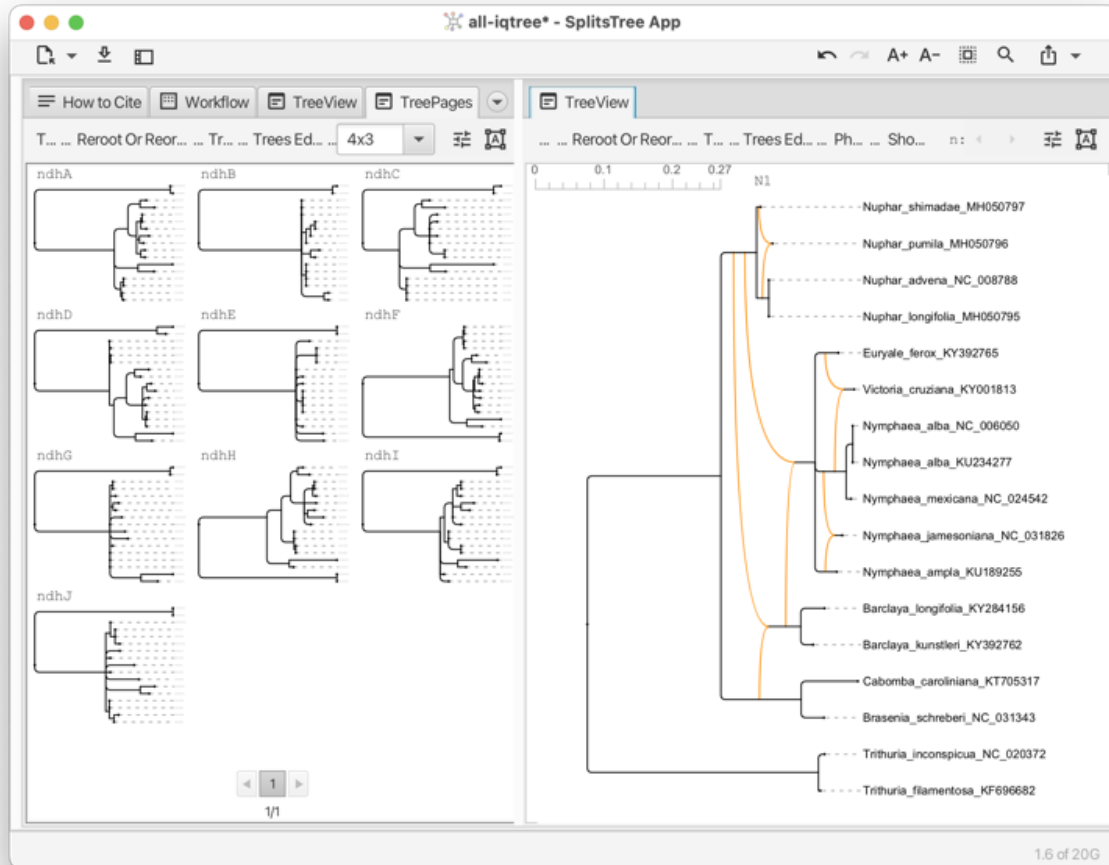


Figure 2.5: On the left we show 10 different gene trees for the NADH dehydrogenase-like complex in waterlilies [Gruenstaedl, 2019] and on the right we show a network that contains all 10 trees, with hybridization number  $h = 5$ , computed using the PhyloFusion algorithm.

number”, that is, the number of reticulations. (To be precise, a reticulation node of indegree  $k$  contributes  $k - 1$  toward the hybridization number.)

SplitsTree currently offers two algorithms for computing such networks for real world data. The Autumn algorithm [Huson and Linz, 2018] (see Section C.4) takes as input two rooted phylogenetic trees and computes, as output the list of all different hybridization networks that contain the two trees. The input trees may have multifurcations and unequal taxon sets. This algorithm aims at providing an exact solution (networks that minimize the hybridization) of a computational hard problem, so it might not terminate if the input trees have too many conflicts.

The PhyloFusion algorithm [Zhang et al., 2023, 2024] takes as input multiple rooted trees and computes one or more rooted phylogenetic networks that display all the input trees. Again, we allow multifurcations and missing taxa. This very fast heuristic aims at minimizing the hybridization number. With this, we provide a versatile method for exploring the practical use of rooted networks in phylogenetics (see Section C.4) (see Fig. 2.5).

### 2.5.3 How to compute a rooted network from rooted trees

Assume that you have a collection of phylogenetic trees for which you would like to explore the use of rooted phylogenetic networks to represent them. To obtain a useful network, you must setup a pipeline consisting of several steps (see Fig. 2.6). In this analysis, incorrect edges are particularly harmful because they generate unnecessary reticulations and so it is important that the input trees have confidence values (such as bootstrap support values, say) associated with the edges so that low-confidence can be ignored.

- First, use the *Reroot or Reorder* algorithm (see Section C.4) to ensure that all trees are correctly rooted (using either midpoint- or outgroup rooting).
- Second, use the *Trees Filter* (see Section C.4) to select the subset of trees that you would like to place into the network.
- Third, use the *Trees Edges Filter* (see Section C.4) to contract any low-confidence edges. By default, the confidence threshold is set to 70.
- Finally, use the *Phylo Fusion* algorithm (see Section C.4) to compute a rooted network that contains all input trees.

### 2.5.4 Cluster networks

The Cluster Network algorithm extracts all clusters from an input set of rooted phylogenetic trees and computes a network using the cluster-popping algorithm [Huson et al., 2012]. This is a fast algorithm that provides a network that contains all input trees. However, it does not aim at minimizing the hybridization number.

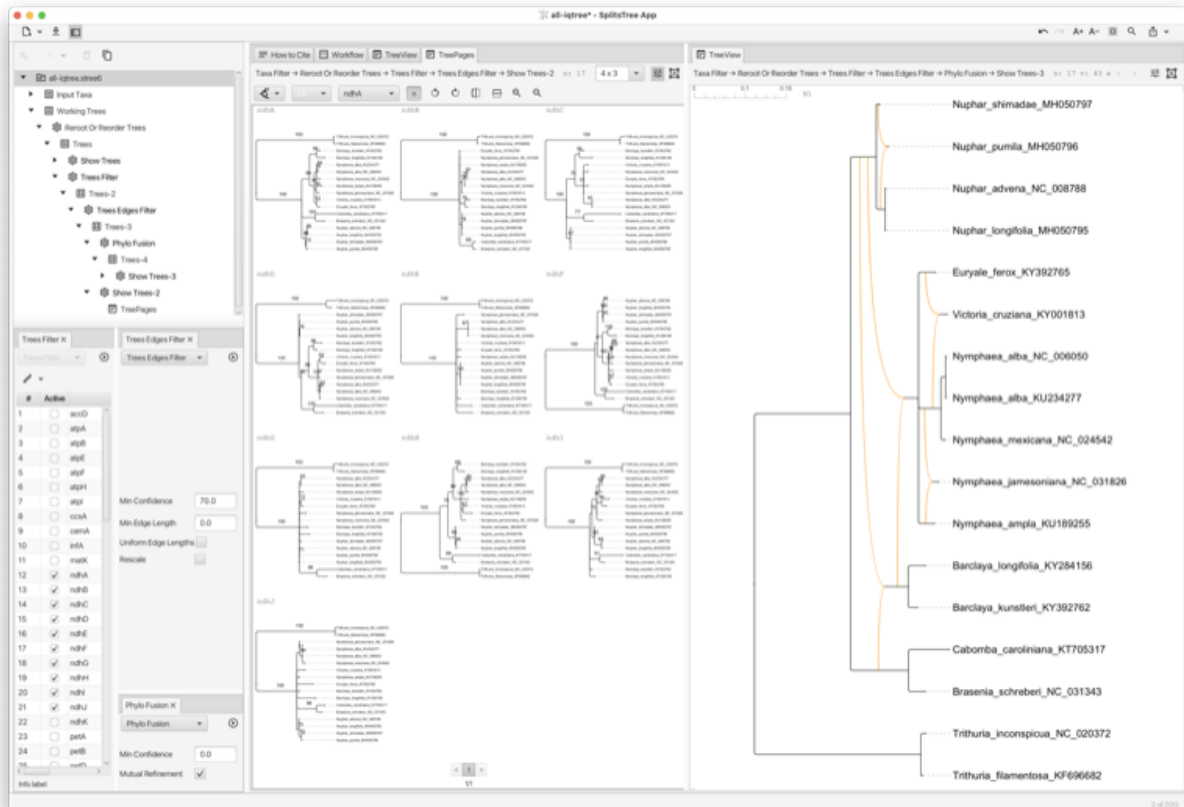


Figure 2.6: For a set of 48 genes in waterlilies [Gruenstaedtl, 2019], on the left we see that 10 different gene trees for the NADH dehydrogenase-like complex have been selected. On the right we see a hybridization network computed using the PhyloFusion algorithm.

## Chapter 3

# Consensus trees and networks

The methods in this chapter all attempt to summarise information contained in a set of trees. (Most also work if the input contains rooted phylogenetic networks, in which case the calculations are based on “hardwired clusters” contained in the networks.) There are several possible sources:

1. Trees returned from different genes or loci.
2. Trees produced from different methods.
3. Trees produced from different bootstrap replicates.
4. Trees sampled from the posterior distribution in a Bayesian analysis.

One of the big improvements with the most recent version of SplitsTree is that the routines for reading in files of trees can now cope with large tree files or large trees.

### 3.1 Consensus trees

A consensus method summarises a set of trees (on the same set of taxa) with a single tree. It can be thought of as analogous to an average tree or median tree.

#### 3.1.1 Average consensus method

The average consensus method implements an idea of Lapointe and Cucumel [1997]. Additive (leaf to leaf) distance matrices are constructed for each tree. This can take some time on larger files. The average of these matrices are then used to construct either a Neighbor-Joining tree or a NeighborNet.

The method can be called from the workflow by selecting a trees block and adding the algorithm ‘Average Consensus’ (see Section C.4). Alternatively, add an ‘Average Distances’ algorithm to the tree block. This creates a new distance block which can be output or analysed using a method of choice.

#### 3.1.2 Strict-, majority- and greedy-consensus methods

The strict consensus, majority rule consensus and greedy consensus are three of the oldest and most widely used consensus methods in phylogenetics.

- The *strict consensus tree* is formed from all splits appearing in all trees;

- The *majority rule tree* is formed from all splits appearing in over half the trees;
- The *greedy consensus tree* is constructed using a greedy algorithm aimed at producing a collection of splits with maximal weight, the weight of each split given by the number of trees containing it.

These methods are available from the Trees menu in the menu bar or by adding an algorithm to the trees block in the workflow.

Note that there is a slight difference in the consensus tree depending on whether the input trees are to be considered rooted or unrooted. For example the two trees

$$((a,b),c,d) \quad \text{and} \quad (a,b,(c,d));$$

share a split  $ab|cd$  which would appear in their unrooted consensus tree, but they share no clusters, so their rooted consensus tree would be completely unresolved.

### 3.1.3 Densi-tree consensus

The densi-tree consensus [Bouckaert, 2010] shows the greedy consensus tree together with a rendering of all input trees (see Section 1.9).

## 3.2 Networks representing trees

### 3.2.1 Consensus networks

Consensus networks are based on the idea of using split networks to represent more splits than can appear in a single tree Bandelt [1995], Holland et al. [2004]. They can be constructed using the menu command `Network>Consensus network`, or by adding an algorithm to a trees block in the workflow. Note that, with the menu command, if there is more than one trees block then SplitsTree will ask the user to select one.

SplitsTree implements several weighting methods for the splits. These are used to determine the split weights used in the output tree or network. A standard analysis consensus network analysis will use the frequency (or count) of a split as the weight used for selecting and displaying splits.

- **Mean** - use the mean of the weights in the input trees. This treats different trees as estimations of the distances between taxa.
- **TreeSizeWeightedMean** - use the mean of the weights in the input trees after normalizing each of the input trees to total length 1. This should be used if the different trees are on different scales, e.g. because they were computed using different methods.
- **Median** - use the median weight. Use as an alternative to mean weights.
- **Count** - use the number of trees that contain a split as its weight. This is useful to emphasize the conflicts in different trees when using a network for consensus.
- **Sum** - use the sum over all weights in the input trees. Similar use-case to counts.
- **Uniform** - give all splits weight 1. This emphasizes the topology of the consensus tree or network.
- **TreeNormalizedSum** - use the sum over all weights in the normalized input trees. Not sure when you would want to use this.

The threshold percent controls how many splits are included in the network. When the weight is computed from split counts it specifies the percentage of trees which a split needs to be contained in for the split to be included in the network. Reducing this threshold will increase the number of splits, giving a more complex network. The

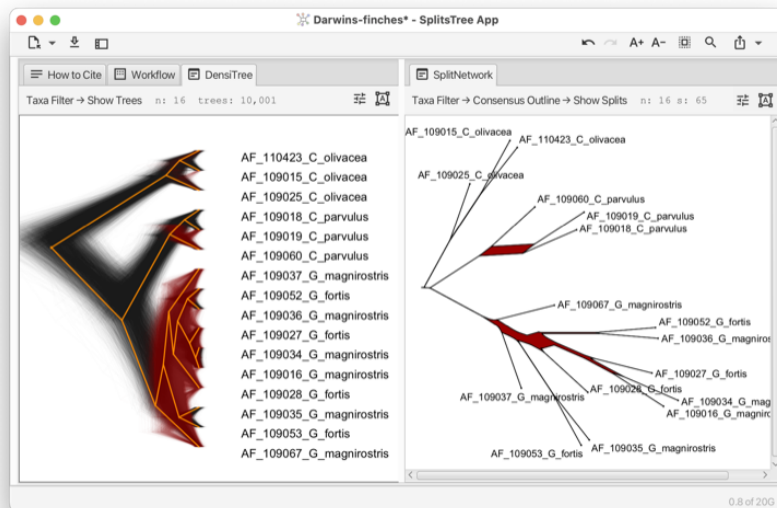


Figure 3.1: For a Bayesian profile of trees (from the Beast examples directory), we show the densi-tree consensus on the left and the outline consensus on the right (rooted by the three taxa at the top of both diagrams).

High Dimension Filter is the same as that used in the split weight filter (see Section C.3), greedily removing splits which generate high dimensional boxes in the diagram.

### 3.2.2 Consensus outline

The consensus outline method (see Section C.4) takes as input a set of trees and produces as output a set of circular splits that are displayed either as a planar split network or as a phylogenetic outline. It operates by greedily selecting a subset of input splits that are compatible with some circular ordering of the input tree, computed using the PQ-tree algorithm [Booth and Lueker, 1976]. One possible application is as an alternative to the densi-tree visualization (see Fig. 3.1).

### 3.2.3 Confidence networks

The idea behind a *confidence network* (see Section C.4) is to choose the threshold in a consensus network so that at least 95% of the trees have *all* their splits contained in that network. The method was originally designed as a way to create confidence intervals from bootstrap distributions Huson and Bryant [2006], however the dimensionality of the problem, and shortcomings of empirical bootstrap distributions, meant that the confidence sets produced were massive. The same machinery can be readily applied to samples from the posterior distribution of trees in a Bayesian analysis, in which case the network represents a confidence set.

The main option in a confidence network is the level, which is 0.95 by default. This is the proportion of input trees which will have their splits contained in the network. Decreasing this number produces smaller networks.





# Appendix A

## The main menu bar

All functionality of the program can be used directly from the main window. In addition, the program provides menus to access the most often used features.

### A.1 The File menu

This menu has the following items:

- New... - Create a new document and open it
- Open... - Open an existing file and process it
- Replace Data... - Replace the current input data
- Edit Input... - Open the input editor tab
- Analyze Draft Genomes... - Open the microbial draft genome analyzer
- Save - Save the current document
- Save As... - Save the current document to a chosen file
- Page Setup... - Page setup for printing
- Print... - Print the current tab
- Close - Close the current document

### A.2 The Edit menu

This menu has the following items:

- Undo - Undo
- Redo - Redo
- Cut - Copy selected text in a text tab or nodes in a workflow tab
- Copy - Copy selected text in a text tab or nodes in a workflow tab
- Copy Newick - Copy tree(s) and networks(s) from the current tab to the system clipboard

- Copy Image - Copy an image of the current tab to the system clipboard
- Paste - Paste text in a text tab or nodes in a workflow tab
- Duplicate - Duplicate selected nodes in a workflow tab
- Delete - Delete selected text in a text tab or nodes in a workflow tab
- Find... - Open the find dialog for the current tab
- Find Again - Find next match in the current tab
- Replace... - Open the replace dialog for the current tab
- Goto Line... - Go to a specific line in current text tab

## A.3 The Select menu

This menu has the following items:

- Select All - Select all taxa or text in the current tab
- Select None - De-select all taxa or text in the current tab
- Select Inverse - Invert the selection in the current tab
- From Previous Window - Select taxa in current tab that were selected in the previously focused window
- Brackets - Select text between matching brackets
- Compatible Sites - Select all compatible sites in the alignment tab

## A.4 The View menu

This menu has the following items:

- Use Dark Theme - Toggle dark mode
- Enter Full Screen - Toggle full-screen mode
- Increase Font Size - Increase the font size in the current tab
- Decrease Font Size - Decrease the font size in the current tab
- Zoom In - Increase the scale factor in the current tab
- Zoom Out - Decrease the scale factor in the current tab
- Zoom In Horizontal - Increase the horizontal scale factor in the current tab
- Zoom Out Horizontal - Decrease the horizontal scale factor in the current tab
- Reset - Reset the scale factor in the current tab
- Rotate Left - Left rotate the tree or network, or selected splits, in the current tab
- Rotate Right - Right rotate the tree or network, or selected splits, in the current tab
- Flip - Flip the tree or network in the current tab
- Show Scale Bar - Show a scale bar in the current tab

- Show QR Code - Show or hide a QR code for the tree or network in the current tab
- Layout Labels - Rerun label layout

## A.5 The Data menu

This menu has the following items:

- Filter Taxa - Show the filter taxa tab
- Filter Characters... - Shows the alignment tab
- Group Identical Haplotypes... - Group identical haplotypes and open as new document
- Filter Trees - Show the filter trees tab
- Filter Splits - Show the filter splits tab
- Splits Slider - Show the splits slider (filter by weight) tab
- Edit Traits - Show the edit traits tab

## A.6 The Distances menu

This menu has the following items:

- P Distances - Compute distances based on the normalized number of character-state differences
- Log Det - Compute distances using the Log-Det method
- Jukes Cantor - Compute distances using the Jukes-Cantor model
- K2P - Compute distances using the Kimura-2P model
- F81 - Compute distances using the Felsenstein-1981 model
- HKY 85 - Compute distances using the Hasegawa-Kishino-Yano model
- F84 - Compute distances using the Felsenstein-1984 model
- Protein ML Dist - Compute distances for proteins using maximum-likelihood estimation
- Gene Content Distance - Compute distances based on the presence/absence of genes

## A.7 The Tree menu

This menu has the following items:

- NJ - Compute a tree from distances using the Neighbor-Joining method
- BioNJ - Compute a tree from distances using the Bio-NJ method
- UPGMA - Compute a rooted tree from distances using the UPGMA method
- Buneman Tree - Compute a tree from distances using the Buneman tree method
- Consensus Tree - Compute a consensus tree

- Minimum Spanning Tree - Compute a minimum spanning tree from distances
- Reroot Or Reorder Trees - Open the reroot or reorder tab
- Show Single Tree - Show a single tree in a tab
- Show Tree Pages - Show multiple trees in a tab
- Show Tanglegram - Show two trees as a tanglegram
- Show DensiTree - Show a Bayesian chain of trees as a densi-tree

## A.8 The Network menu

This menu has the following items:

- Neighbor Net - Compute a planar split network from distances using the Neighbor-Net method
- Split Decomposition - Compute a split network from distances using the split-decomposition method
- Parsimony Splits - Compute a split network from DNA sequences using the parsimony-splits method
- Consensus Splits - Open the consensus splits tab
- Consensus Network - Compute a consensus network from trees
- Consensus Outline - Compute a consensus outline from trees
- Super Network - Compute a super network from trees on unequal taxon sets
- Median Joining Network - Compute a haplotype network from DNA sequences using the median-joining algorithm
- Min Spanning Network - Compute a minimum spanning network from distances
- World Map - Show the world map tab
- Hybridization Network - Compute a minimum hybridization network for two rooted trees using the autumn algorithm
- Cluster Network - Compute a cluster network from trees

## A.9 The Analysis menu

This menu has the following items:

- Bootstrap Tree - Perform bootstrapping on a tree calculation from sequences
- Bootstrap Tree as Network - Perform bootstrapping on a tree calculation from sequences, producing a network
- Bootstrap Splits Network - Perform bootstrapping on a split network calculation from sequences
- Estimate Invariable Sites - Estimate the proportion of invariable sites
- Compute Delta Score - Compute the delta score from distances
- Run Phi Test for Recombination - Run the Phi Test to detect recombination in sequences
- Compute Tajima's D - Run the Phi Test to detect recombination in sequences

- Splits Phylogenetic Diversity - Compute the phylogenetic diversity for selected taxa on splits
- Splits Shapley Values - Compute Shapely values on splits
- Tree Phylogenetic Diversity - Compute the phylogenetic diversity for selected taxa on trees
- Rooted Tree Fair Proportion Diversity Index - Compute rooted-tree fair-proportion diversity index
- Rooted Tree Equal Splits Diversity Index - Compute rooted-tree equal-splits diversity index
- Unrooted Tree Shapley Values - Compute Shapely values on unrooted trees
- PCoA - Perform principal coordinate analysis on distances
- Show Workflow - Show the workflow tab

## **A.10 The Window menu**

This menu has the following items:

- Show Message Window... - Show or hide the message window
- Set Window Size... - Set the window size
- Untitled

## **A.11 The Help menu**

This menu has the following items:

- Check For Updates... - Check whether an update is available for download
- About... - Show splash screen (authors, version number and license).



# Appendix B

## Main data blocks

SplitsTree is organized around data blocks that correspond to “Nexus” blocks [Maddison et al., 1997].

### B.1 Taxa block

This block maintains the list of all taxa in the analysis. There is a fixed number (nTax) of taxa and each has an id 1..nTax and an unique label. Optionally, an info string can be provided for each taxon. Also, a display label may be provided for each taxon. This can include certain HTML tags that are used to render the label.

```
BEGIN TAXA;
  [TITLE title;]
  DIMENSIONS NTAX=number-of-taxa;
  [TAXLABELS
    list-of-labels
  ;]
  [TAXINFO
    list-of-info-items (use 'null' for missing item)
  ;]
  [DISPLAYLABELS
    list-of-html-strings (use 'null' for missing item)
  ;]
END;
```

### B.2 Traits block

This block contains traits associated with the input taxa. Each trait has a label, optional latitude and longitude, and a value, which can either be a number or a string.

```
BEGIN TRAITS;
  [TITLE {title};]
  DIMENSIONS [NTAX=number-of-taxa] NTRAITS=number-of-traits;
  [FORMAT
    [LABELS={YES|NO}]
    [MISSING=symbol]
    [SEPARATOR={COMMA|SEMICOLON|WHITESPACE}]
  ;]
  [TRAITLATITUDE latitude-trait-1 latitude-trait-2 ... latitude-trait-n;
  TRAITLONGITUDE longitude-trait-1 longitude-trait-2 ... longitude-trait-n;]
  [TRAITCOLOR color-trait-1 color-trait-2 ... color-trait-n;]
```

```

[TAXONCOLOR color-taxon-1 color-taxon-2 ... color-taxon-ntax;]

    TRAITLABELS label-trait-1 label-trait-2 ... label-trait-n;
MATRIX
    trait data in specified format
;
END;

```

## B.3 Characters block

This block maintains a set of characters or a multiple sequence alignment. There is a fixed number of taxa and characters or positions. Several different formats are supported. Characters can have weights and both characters and their states can have labels.

```

BEGIN CHARACTERS;
    [TITLE {title};]
    [LINK {type} = {title};]
    DIMENSIONS [NTAX=number-of-taxa] NCHAR=number-of-characters;
    [FORMAT
        [DATATYPE={STANDARD|DNA|RNA|PROTEIN|MICROSAT}]
        [RESPECTCASE]
        [MISSING=symbol]
        [GAP=symbol]
        [MatchChar=symbol]
        [SYMBOLS="symbol symbol ..."]
        [LABELS={NO|LEFT}]
        [TRANSPPOSE={NO|YES}]
        [INTERLEAVE={NO|YES}]
        [TOKENS=NO]
    ;]
    [CHARWEIGHTS wgt_1 wgt_2 ... wgt_nchar;]
    [CHARSTATELABELS character-number [ character-name ] [ /state-name [ state-name... ] ], ...;]
    [CHARLABELS character-name [character-name...];]
MATRIX
    sequence data in specified format
;
END;

```

## B.4 Distances block

This block maintains a distance matrix of size nTax times nTax.

```

BEGIN DISTANCES;
    [TITLE {title};]
    [LINK {type} = {title};]
    [DIMENSIONS NTAX=number-of-taxa;]
    [FORMAT
        [TRIANGLE={LOWER|UPPER|BOTH}]
        [[NO] DIAGONAL]
        [LABELS={LEFT|NO}]
    ;]
MATRIX
    distance data in specified format
;
END;

```



## B.5 Trees block

This block maintains a list of trees. These can be rooted or unrooted phylogenetic trees, or rooted phylogenetic networks. Trees can partial in the sense that they need to contain all taxa.

```
BEGIN TREES;
  [TITLE {title};]
  [LINK {type} = {title};]
[PROPERTIES [PARTIALTREES={YES|NO}] [ROOTED={YES|NO}] [RETICULATED={YES|NO}];]
[TRANSLATE
  nodeLabel1 taxon1,
  nodeLabel2 taxon2,
  ...
  nodeLabelN taxonN
;]
[TREE name1 = tree1-in-Newick-format;]
[TREE name2 = tree2-in-Newick-format;]
...
[TREE nameM = treeM-in-Newick-format;]
END;
```

## B.6 Splits block

This block maintains a set of splits, usually with weights, sometimes with confidence values, and usually with a suitable cycle for layout purposes.

```
BEGIN SPLITS;
  [TITLE {title};]
  [LINK {type} = {title};]
[DIMENSIONS [NTAX=number-of-taxa] [NSPLITS=number-of-splits];]
[FORMAT
  [Labels={LEFT|NO}]
  [Weights={YES|NO}]
  [Confidences={YES|NO}]
  [Intervals={YES|NO}]
  [ShowBothSides={NO|YES}]
;]
[Threshold=non-negative-number;]
[PROPERTIES
  [Fit=non-negative-number]
  [{Compatible|Cyclic|Weakly Compatible|Incompatible}]
;]
[CYCLE [taxon_i_1 taxon_i_2 ... taxon_i_ntax];]
[SPLITSLABELS label_1 label_2 ... label_nsplits;]
MATRIX
[label_1] [weight_1] [confidence_1] split_1,
[label_2] [weight_2] [confidence_2] split_2,
...
[label_nsplits] [weight_nsplits] [confidence_nsplits] split_nsplits[,]
;
END;
```

## B.7 Network block

Maintain a network, such as a haplotype network or just a set of points (for PCoA).

```
BEGIN NETWORK;
  [TITLE {title};]
  [LINK {type} = {title};]
```

```

[DIMENSIONS [NVertices=number-of-nodes] [NEdges=number-of-edges];]
[TYPE {HaplotypeNetwork|Points|Other};]
[FORMAT
;]
[PROPERTIES
[info =' information string to be shown with plot']
;]
VERTICES
ID=number [LABEL=label] [x=number] [y=number] [key=value ...],
...
ID=number [LABEL=label] [x=number] [y=number] [key=value ...]
;
EDGES
ID=number SID=number TID=number [LABEL=label] [key=value ...],
...
ID=number SID=number TID=number [LABEL=label] [key=value ...]
;
END;

```

## B.8 View block

This block represents a viewer.

```

BEGIN VIEW;
[TITLE title;]
[LINK {type} = {title};]
NAME <name>;
INPUT <input-block-name>;
[OPTIONS
<name>=<value>,
...
<name>=<value>
;]
END;

```

## B.9 Algorithms block

This block represents an algorithm.

```

BEGIN ALGORITHM;
[TITLE <title>;]
[LINK <parent-block-type> = <parent-title>;]
NAME <name>;
[OPTIONS
<name>=<value>,
...
<name>=<value>
;]
END;

```

## B.10 Report block

This block represents a textual report.

```

BEGIN REPORT;
[TITLE title;]
[LINK {type} = {title};]

```

```

TEXT
    text...
;
END;

```

## B.11 Sets block

This block represents a collection of taxon sets and/or character sets.

```

BEGIN SETS;
    [TITLE {title};]
    [TAXSET {name}={list of names and/or IDS};]
    ...
    [CHARSET {name}={list of positions};]
    ...
END;

```

## B.12 SplitsTree6 block

This block holds program-specific data and its presence in a file indicates that the file was generated by SplitsTree6 and represents a complete analysis.

```

BEGIN SPLITSTREE6;
    DIMENSIONS nDataNodes=number nAlgorithms=number;
    PROGRAM version=version-string;
    WORKFLOW creationDate=long;
END;

```

## B.13 Genomes block

This block represents a collection of genomes.

```

BEGIN GENOMES;
    [TITLE {title};]
    [LINK {type} = {title};]
    [DIMENSIONS NTAX=number-of-taxa;]
    [FORMAT
        [LABELS={YES|NO}]
        [ACCESSIONS={YES|NO}]
        [MULTIPART={YES|NO}]
        [FILES={YES|NO}]
    ;]
    MATRIX
        [label] [accession] length {sequence | [#parts] length {sequence|{file:... offset}}} .. length {sequence|{file offset}}},
        ...
        [label] [accession] length {sequence | [#parts] length {sequence|{file:... offset}}} .. length {sequence|{file offset}}}
    ;]
END;

```



# Appendix C

## Algorithms

Here we list of all provided algorithms, organized by input data.

### C.1 Algorithms on a Characters Block

#### P Distance

The *P Distance* algorithm takes a Characters block as input and produces a Distances block as output. It computes the normalized Hamming distance. The algorithm has the following options:

`HandleAmbiguousStates = {Ignore | AverageStates | MatchStates}` - choose how to handle ambiguous states (nucleotide data only)

Reference: [Hamming, 1950]

#### Hamming Distance

The *Hamming Distance* algorithm takes a Characters block as input and produces a Distances block as output. It computes the Hamming distance, that is the number of differences between sequences. The algorithm has the following options:

`HandleAmbiguousStates = {Ignore | AverageStates | MatchStates}` - choose how to handle ambiguous states (nucleotide data only)

Reference: [Hamming, 1950]

#### Log Det

The *Log Det* algorithm takes a Characters block as input and produces a Distances block as output. It computes distances using the Log-Det method. The algorithm has the following options:

`PropInvariableSites = <Double>` - proportion of invariable sites

`FudgeFactor = <Boolean>` - input missing matrix entries using LDDist method

`FillZeros = <Boolean>` - replace zeros with small numbers in rows/columns with values

Reference: [Steel, 1994]

## Jukes Cantor Distance

The *Jukes Cantor Distance* algorithm takes a Characters block as input and produces a Distances block as output. It calculates distances under the Jukes-Cantor model. The algorithm has the following options:

PropInvariableSites = <Double> - proportion of invariable sites

SetSiteVarParams = {fromChars | defaultValues} - set site variation parameters to default values, or to estimations from characters

Reference: [Jukes and Cantor, 1969]

## K2P Distance

The *K2P Distance* algorithm takes a Characters block as input and produces a Distances block as output. It calculates distances under the Kimura-2P model. The algorithm has the following options:

TsTvRatio = <Double> - ratio of transitions vs transversions

Gamma = <Double> - alpha value for the Gamma distribution

PropInvariableSites = <Double> - proportion of invariable sites

SetSiteVarParams = {fromChars | defaultValues} - set site variation parameters to default values, or to estimations from characters

UseML\_Distances = <Boolean> - use maximum likelihood estimation of distances (rather than exact distances)

Reference: [Kimura, 1980]

## F81 Distance

The *F81 Distance* algorithm takes a Characters block as input and produces a Distances block as output. It calculates distances under the Felsenstein-81 model. The algorithm has the following options:

BaseFrequencies = <doubleArray> - base frequencies (in order ACGT/U)

SetBaseFrequencies = {fromChars | defaultValues} - set base frequencies to default values, or to estimations from characters (using Capture-recapture for invariable sites)

PropInvariableSites = <Double> - proportion of invariable sites

SetSiteVarParams = {fromChars | defaultValues} - set site variation parameters to default values, or to estimations from characters

UseML\_Distances = <Boolean> - use maximum likelihood estimation of distances (rather than exact distances)

Reference: [Felsenstein, 1981]

## HKY85 Distance

The *HKY85 Distance* algorithm takes a Characters block as input and produces a Distances block as output. It calculates distances under the Hasegawa-Kishino-Yano model. The algorithm has the following options:

TsTvRatio = <Double> - ratio of transitions vs transversions

BaseFrequencies = <doubleArray> - base frequencies (in order ACGT/U)

SetBaseFrequencies = {fromChars | defaultValues} - set base frequencies to default values, or to estimations from characters (using Capture-recapture for invariable sites)

PropInvariableSites = <Double> - proportion of invariable sites

SetSiteVarParams = {fromChars | defaultValues} - set site variation parameters to default values, or to estimations from characters

Reference: [Hasegawa et al., 1985]

## F84 Distance

The *F84 Distance* algorithm takes a Characters block as input and produces a Distances block as output. It calculates distances under the Felsenstein-84 model. The algorithm has the following options:

BaseFrequencies = <doubleArray> - base frequencies (in order ACGT/U)

SetBaseFrequencies = {fromChars | defaultValues} - set base frequencies to default values, or to estimations from characters (using Capture-recapture for invariable sites)

PropInvariableSites = <Double> - proportion of invariable sites

SetSiteVarParams = {fromChars | defaultValues} - set site variation parameters to default values, or to estimations from characters

UseML\_Distances = <Boolean> - use maximum likelihood estimation of distances (rather than exact distances)

Reference: [Felsenstein and Churchill, 1996]

## GTR Distance

The *GTR Distance* algorithm takes a Characters block as input and produces a Distances block as output. It calculates distances under the general time-reversible model. The algorithm has the following options:

PropInvariableSites = <Double> - proportion of invariable sites

SetSiteVarParams = {fromChars | defaultValues} - set site variation parameters to default values, or to estimations from characters

RateMatrix = <doubleSquareMatrix> - rate matrix for GTRDistance (in order ACGT/U)

UseML\_Distances = <Boolean> - use maximum likelihood estimation of distances (rather than exact distances)

Reference: [Tavar, 1986]

## Protein ML Distance

The *Protein ML Distance* algorithm takes a Characters block as input and produces a Distances block as output. It computes distances for proteins using maximum-likelihood estimation. The algorithm has the following options:

Model = {cpREV45 | Dayhoff | JTT | mtMAM | mtREV24 | pmb | Rhodopsin | WAG} - choose an amino acid substitution model

PropInvariableSites = <Double> - proportion of invariable sites

Gamma = <Double> - alpha parameter for gamma distribution. Negative gamma = Equal rates

Reference: [Swofford et al., 1996]

## Dice Distance

The *Dice Distance* algorithm takes a Characters block as input and produces a Distances block as output. It computes distances using the DiceDistance coefficient distance.

Reference: [Dice, 1945]

### **Jaccard Distance**

The *Jaccard Distance* algorithm takes a Characters block as input and produces a Distances block as output. It computes distances based on the JaccardDistance index.

Reference: [Jaccard, 1901]

### **Gene Content Distance**

The *Gene Content Distance* algorithm takes a Characters block as input and produces a Distances block as output. It computes distances based on the presence/absence of genes.

Reference: [Huson and Steel, 2004]

### **Gene Sharing Distance**

The *Gene Sharing Distance* algorithm takes a Characters block as input and produces a Distances block as output. It computes distances using the gene-sharing distance.

Reference: [Snel et al., 1997]

### **Upholt Restriction Distance**

The *Upholt Restriction Distance* algorithm takes a Characters block as input and produces a Distances block as output. It calculates distances for restriction data.

Reference: [Upholt, 1977]

### **Nei Li Restriction Distance**

The *Nei Li Restriction Distance* algorithm takes a Characters block as input and produces a Distances block as output. It calculates distances for restriction data.

Reference: [Nei and Li, 1979]

### **Base Freq Distance**

The *Base Freq Distance* algorithm takes a Characters block as input and produces a Distances block as output. It calculates distances from differences in the base composition.

### **Binary To Splits**

The *Binary To Splits* algorithm takes a Characters block as input and produces a Splits block as output. It converts binary characters directly into splits. The algorithm has the following options:

MinSplitWeight = <Double> - minimum split weight threshold

HighDimensionFilter = <Boolean> - activate high-dimensional filter to avoid exponential graph size

AddAllTrivial = <Boolean> - ensure all trivial splits are present

Reference: [Huson et al., 2012]



## DNA To Splits

The *DNA To Splits* algorithm takes a Characters block as input and produces a Splits block as output. It converts DNA characters directly into splits. The algorithm has the following options:

Method = {MajorityState | RYAlphabet} - use either majority-state-vs-others or RY alphabet

MinSplitWeight = <Double> - minimum split weight threshold

HighDimensionFilter = <Boolean> - activate high-dimensional filter to avoid exponential graph size

Reference: [Huson et al., 2012]

## Median Joining

The *Median Joining* algorithm takes a Characters block as input and produces a Network block as output. It computes a haplotype network using the median-joining method. The algorithm has the following options:

Epsilon = <Integer> - balances accuracy (smaller value) and efficiency (larger value)

Reference: [Bandelt et al., 1999]

## Parsimony Splits

The *Parsimony Splits* algorithm takes a Characters block as input and produces a Splits block as output. It computes weakly-compatible splits directly from DNA characters.

Reference: [Bandelt and Dress, 1992]

## Characters Filter

The *Characters Filter* algorithm takes a Characters block as input and produces a Characters block as output. It filter characters. The algorithm has the following options:

ExcludeGapSites = <Boolean> - exclude all sites that contain a gap

ExcludeParsimonyUninformativeSites = <Boolean> - exclude all sites that are parsimony uninformative

ExcludeConstantSites = <Boolean> - exclude all sites that are constant

ExcludeFirstCodonPosition = <Boolean> - exclude first and then every third site

ExcludeSecondCodonPosition = <Boolean> - exclude second and then every third site

ExcludeThirdCodonPosition = <Boolean> - exclude third and then every third site

## External Program

The *External Program* algorithm takes a Characters block as input and produces a Trees block as output. It runs an external program. The algorithm has the following options:

Name = <String> - specify a name for this calculation

ProgramCall = <String> - specification of external program: replace 'path-to-program' by path to program and

use '%i' and '%o' as place - holders for the program's input and output files

CharactersFormat = {Phylip | Nexus | FastA} - specify the format to write out the current data in

### **Estimate Invariable Sites**

The *Estimate Invariable Sites* algorithm takes a Characters block as input and produces a Report block as output. It estimates the proportion of invariant sites using capture-recapture.

Reference: [Steel et al., 2000]

### **Phi Test**

The *Phi Test* algorithm takes a Characters block as input and produces a Report block as output. It performs a statistical test for detecting the presence of recombination.

Reference: [Bruen et al., 2006]

### **Tajima D**

The *Tajima D* algorithm takes a Characters block as input and produces a Report block as output. It performs Tajima's D test to determine whether a DNA sequence is evolving neutrally. The algorithm has the following options:

ExcludeGapSites = <Boolean> - exclude gapped sites from calculation.

Reference: [Tajima, 1989]

## **C.2 Algorithms on a Distances Block**

### **Neighbor Joining**

The *Neighbor Joining* algorithm takes a Distances block as input and produces a Trees block as output. It computes an unrooted phylogenetic tree using the neighbor-joining method.

Reference: [Saitou and Nei, 1987]

### **Bio NJ**

The *Bio NJ* algorithm takes a Distances block as input and produces a Trees block as output. It computes an unrooted phylogenetic tree using the Bio-NJ method.

Reference: [Gascuel, 1997]

### **UPGMA**

The *UPGMA* algorithm takes a Distances block as input and produces a Trees block as output. It computes a rooted phylogenetic tree using the UPGMA method.

Reference: [Sokal and Michener, 1958]

## Neighbor Net

The *Neighbor Net* algorithm takes a Distances block as input and produces a Splits block as output. It computes a set of cyclic splits using the neighbor-net method. The algorithm has the following options:

InferenceAlgorithm = {GradientProjection | ActiveSet | APGD | SplitsTree4} - the inference algorithm to be used

References: [Bryant and Moulton, 2004, Bryant and Huson, 2023]

## Split Decomposition

The *Split Decomposition* algorithm takes a Distances block as input and produces a Splits block as output. It computes a set of weakly-compatible splits using the split-decomposition method.

Reference: [Bandelt and Dress, 1992]

## Buneman Tree

The *Buneman Tree* algorithm takes a Distances block as input and produces a Splits block as output. It computes a set of compatible splits using the Buneman tree method.

Reference: [Bandelt and Dress, 1992]

## Min Spanning Network

The *Min Spanning Network* algorithm takes a Distances block as input and produces a Network block as output. It computes a minimum spanning network. The algorithm has the following options:

Epsilon = <Double> - weighted genetic distance measure. Low: MedianJoining, High: full median network

MinSpanningTree = <Boolean> - calculate minimum spanning tree

Reference: [Excoffier and Smouse, 1994]

## Min Spanning Tree

The *Min Spanning Tree* algorithm takes a Distances block as input and produces a Trees block as output. It computes a minimum spanning tree.

Reference: [Excoffier and Smouse, 1994]

## PCOA

The *PCOA* algorithm takes a Distances block as input and produces a Network block as output. It performs principal coordinates analysis. The algorithm has the following options:

FirstCoordinate = <Integer> - choose principal component for the x Axis

SecondCoordinate = <Integer> - choose principal component for the y Axis

Reference: [Gower, 1966]

## Delta Score

The *Delta Score* algorithm takes a Distances block as input and produces a Report block as output. It calculates the delta score.

Reference: [Holland et al., 2002]

## C.3 Algorithms on a Splits Block

### Bootstrap Splits

The *Bootstrap Splits* algorithm takes a Splits block as input and produces a Splits block as output. It performs bootstrapping on splits. The algorithm has the following options:

Replicates = <Integer> - number of bootstrap replicates

MinPercent = <Double> - minimum percentage support for a split to be included

ShowAllSplits = <Boolean> - show all bootstrap splits, not just the original splits

RandomSeed = <Integer> - if non-zero, is used as seed for random number generator

HighDimensionFilter = <Boolean> - heuristically remove splits causing high-dimensional network

Reference: [Felsenstein, 1985]

### Greedy Tree

The *Greedy Tree* algorithm takes a Splits block as input and produces a Trees block as output. It produces a phylogenetic tree based on greedily selecting a compatible set of splits.

Reference: [Huson et al., 2012]

### Dimension Filter

The *Dimension Filter* algorithm takes a Splits block as input and produces a Splits block as output. It heuristically remove splits that lead to high-dimensional boxes in a split network. The algorithm has the following options:

MaxDimension = <Integer> - heuristically remove splits that create configurations of a higher dimension than this threshold

### Show Splits

The *Show Splits* algorithm takes a Splits block as input and produces a View block as output. It provides interactive visualizations of split networks. The algorithm has the following options:

View = {SplitNetwork} - the type of splits viewer to use

### Splits Filter

The *Splits Filter* algorithm takes a Splits block as input and produces a Splits block as output. It filter splits. The algorithm has the following options:

WeightThreshold = <Float> - set minimum split weight threshold

`ConfidenceThreshold` = `<Float>` - set the minimum split confidence threshold

`MaximumDimension` = `<Integer>` - set maximum dimension threshold (necessary to avoid computational overload)

`FilterAlgorithm` = {`None` | `GreedyCompatible` | `GreedyCircular` | `GreedyWeaklyCompatible` | `BlobTree`} - set the filter algorithm

`RecomputeCycle` = `<Boolean>` - recompute circular ordering

### Weights Slider

The *Weights Slider* algorithm takes a Splits block as input and produces a Splits block as output. It allows one to interactively filter splits by their weight. The algorithm has the following options:

`WeightThreshold` = `<Double>` - set minimum split weight threshold

### Incompatibility Score

The *Incompatibility Score* algorithm takes a Splits block as input and produces a Report block as output. It calculates an incompatibility score on splits.

### Phylogenetic Diversity

The *Phylogenetic Diversity* algorithm takes a Splits block as input and produces a Report block as output. It calculates the phylogenetic diversity for selected taxa.

Reference: [Volkman et al., 2014]

### Shapley Values

The *Shapley Values* algorithm takes a Splits block as input and produces a Report block as output. It calculates Shapley values on splits.

Reference: [Volkman et al., 2014]

## C.4 Algorithms on a Trees Block

### Autumn Algorithm

The *Autumn Algorithm* algorithm takes a Trees block as input and produces a Trees block as output. It computes all minimum hybridization networks using the Autumn algorithm. The algorithm has the following options:

`FirstTree` = `<Integer>` - index of the first tree

`SecondTree` = `<Integer>` - index of the second tree

`OnlyOneNetwork` = `<Boolean>` - report only one network

`RerootToMinimize` = `<Boolean>` - reroot input trees to minimize hybridization number

Reference: [Huson and Linz, 2018]

## Phylo Fusion

The *Phylo Fusion* algorithm takes a Trees block as input and produces a Trees block as output. It combines multiple rooted phylogenetic trees into a rooted network. The algorithm has the following options:

OnlyOneNetwork = <Boolean> - report only one network

MutualRefinement = <Boolean> - mutually refine input trees

NormalizeEdgeWeights = <Boolean> - normalize input edge weights

SearchHeuristic = {Thorough | Medium | Fast} - fast, Medium, or Thorough search

GroupNonSeparated = <Boolean> - improve performance by grouping taxa that are not separated by a non-trivial edge

CladeReduction = <Boolean> - improve performance using clade reduction

References: [Zhang et al., 2023, 2024]

## Average Consensus

The *Average Consensus* algorithm takes a Trees block as input and produces a Splits block as output. It calculates average consensus tree.

Reference: [Lapointe and Cucumel, 1997]

## Blob Tree

The *Blob Tree* algorithm takes a Trees block as input and produces a Trees block as output. It extract the blob tree from a rooted network The algorithm has the following options:

SeparateBlobs = <Boolean> - for any blob that shares its top node with some other blob, insert an edge above it to keep blobs separate

Reference: [Huson et al., 2012]

## Bootstrap Tree Splits

The *Bootstrap Tree Splits* algorithm takes a Trees block as input and produces a Splits block as output. It performs bootstrapping on trees. The algorithm has the following options:

Replicates = <Integer> - number of bootstrap replicates

MinPercent = <Double> - minimum percentage support for a split to be included

ShowAllSplits = <Boolean> - show all bootstrap splits, not just the original splits

RandomSeed = <Integer> - if non-zero, is used as seed for random number generator

HighDimensionFilter = <Boolean> - heuristically remove splits causing high-dimensional network

Reference: [Felsenstein, 1985]

## Bootstrap Tree

The *Bootstrap Tree* algorithm takes a Trees block as input and produces a Trees block as output. It performs bootstrapping on trees. The algorithm has the following options:

Replicates = <Integer> - number of bootstrap replicates

TransferBootstrap = <Boolean> - use transform bootstrapping (TBE), less susceptible to rogue taxa

MinPercent = <Double> - minimum percentage support for a branch to be included

RandomSeed = <Integer> - if non-zero, is used as seed for random number generator

Reference: [Felsenstein, 1985]

## Cluster Network

The *Cluster Network* algorithm takes a Trees block as input and produces a Trees block as output. It computes the cluster network that contains all input trees (in the hardwired sense). The algorithm has the following options:

EdgeWeights = {Mean | Count | Sum | Uniform} - compute edge weights

ThresholdPercent = <Double> - minimum percentage of trees that a cluster must appear in

Reference: [Huson and Rupp, 2008]

## Consensus Tree

The *Consensus Tree* algorithm takes a Trees block as input and produces a Trees block as output. It provides several methods for computing a consensus tree. The algorithm has the following options:

Consensus = {Majority | Greedy | Strict} - consensus method to use

Reference: [Bryant, 2001]

## Consensus Network

The *Consensus Network* algorithm takes a Trees block as input and produces a Splits block as output. It computes the consensus network. The algorithm has the following options:

EdgeWeights = {Mean | TreeSizeWeightedMean | Median | Count | Sum | Uniform | TreeNormalizedSum} - how to calculate edge weights in resulting network

ThresholdPercent = <Double> - threshold for percent of input trees that split has to occur in for it to appear in the output

HighDimensionFilter = <Boolean> - heuristically remove splits causing high-dimensional consensus network

Reference: [Holland et al., 2004]

## Consensus Outline

The *Consensus Outline* algorithm takes a Trees block as input and produces a Splits block as output. It computes the consensus outline. The algorithm has the following options:

EdgeWeights = {Mean | TreeSizeWeightedMean | Median | Count | Sum | Uniform | TreeNormalizedSum} - how to calculate edge weights in resulting network

ThresholdPercent = <Double> - threshold for percent of input trees that split has to occur in for it to appear in the output

Reference: [Huson and Cetinkaya, 2023]

## Consensus Splits

The *Consensus Splits* algorithm takes a *Trees* block as input and produces a *Splits* block as output. It provides several consensus methods. The algorithm has the following options:

`Consensus = {Strict | Majority | GreedyCompatible | ConsensusOutline | GreedyWeaklyCompatible | ConsensusNetwork}`  
- consensus method

`EdgeWeights = {Mean | TreeSizeWeightedMean | Median | Count | Sum | Uniform | TreeNormalizedSum}` - how to calculate edge weights in resulting network

`ThresholdPercent = <Double>` - threshold for percent of input trees that split has to occur in for it to appear in the output

`HighDimensionFilter = <Boolean>` - heuristically remove splits causing high-dimensional consensus network

Reference: [Huson et al., 2012]

## Filtered Super Network

The *Filtered Super Network* algorithm takes a *Trees* block as input and produces a *Splits* block as output. It computes a super network using the Z-closure method. The algorithm has the following options:

`MinNumberTrees = <Integer>` - set the min number trees

`MaxDistortionScore = <Integer>` - set the max distortion score

`UseTotalScore = <Boolean>` - set the use total score

Reference: [Whitfield et al., 2008]

## LSA Tree

The *LSA Tree* algorithm takes a *Trees* block as input and produces a *Trees* block as output. It extract the LSA tree from a rooted network.

Reference: [Huson et al., 2012]

## Normalize Rooted Networks

The *Normalize Rooted Networks* algorithm takes a *Trees* block as input and produces a *Trees* block as output. It the Normalize Rooted Networks algorithm

Reference: [Francis et al., 2021]

## Rooted Consensus Tree

The *Rooted Consensus Tree* algorithm takes a *Trees* block as input and produces a *Trees* block as output. It provides several methods for computing a rooted consensus tree. The algorithm has the following options:

`Consensus = {Majority | Strict | Greedy}` - consensus method to use



## Reroot Or Reorder Trees

The *Reroot Or Reorder Trees* algorithm takes a *Trees* block as input and produces a *Trees* block as output. It reroot or reorder all trees. The algorithm has the following options:

`RootBy = {Off | MidPoint | OutGroup}` - determine how to reroot

`RearrangeBy = {Off | RotateChildren | RotateSubTrees | ReverseChildren | ReverseSubTrees}` - determine how to rearrange

`Reorder = {Off | ByTaxa | Lexicographically | ReverseOrder | LadderizedUp | LadderizedDown | LadderizedRandom | Stabilize}` - determine how to reorder

`Rescale = <Boolean>` - rescale each tree to total length of 100

## Show Trees

The *Show Trees* algorithm takes a *Trees* block as input and produces a *View* block as output. It provides several types of interactive visualizations of trees. The algorithm has the following options:

`View = {TreeView | TreePages | Tanglegram | DensiTree}` - the type of viewer to use

## Super Network

The *Super Network* algorithm takes a *Trees* block as input and produces a *Splits* block as output. It computes a super network using the Z-closure method. The algorithm has the following options:

`EdgeWeights = {AverageRelative | Mean | TreeSizeWeightedMean | Sum | Min | None}` - determine how to calculate edge weights in resulting network

`SuperTree = <Boolean>` - enforce the strong induction property, which results in a super tree

`NumberOfRuns = <Integer>` - number of runs using random permutations of the input splits

`ApplyRefineHeuristic = <Boolean>` - apply a simple refinement heuristic

`Seed = <Integer>` - set seed used for random permutations

`HighDimensionFilter = <Boolean>` - heuristically remove splits causing high-dimensional network

Reference: [Huson et al., 2004]

## Trees Filter

The *Trees Filter* algorithm takes a *Trees* block as input and produces a *Trees* block as output. It filter trees.

## Trees Filter More

The *Trees Filter More* algorithm takes a *Trees* block as input and produces a *Trees* block as output. It filter trees. The algorithm has the following options:

`RequireAllTaxa = <Boolean>` - keep only trees that have the full set of taxa

`MinNumberOfTaxa = <Integer>` - keep only trees that have at least this number of taxa

`MinTotalTreeLength = <Double>` - keep only trees that have at least this total length

## Trees Edges Filter

The *Trees Edges Filter* algorithm takes a Trees block as input and produces a Trees block as output. It filter edges in trees. The algorithm has the following options:

MinConfidence = <Double> - keep only edges that have this minimum confidence value

MinEdgeLength = <Double> - keep only edges that have this minimum length

UniformEdgeLengths = <Boolean> - change all edge weights to 1

Rescale = <Boolean> - rescale each tree to total length of 100

## Tree Selector

The *Tree Selector* algorithm takes a Trees block as input and produces a Trees block as output. It select a tree to use. The algorithm has the following options:

Which = <Integer> - which tree to use

## Tree Selector Splits

The *Tree Selector Splits* algorithm takes a Trees block as input and produces a Splits block as output. It selects a single tree and extracts its splits. The algorithm has the following options:

Which = <Integer>

## Unique Topologies

The *Unique Topologies* algorithm takes a Trees block as input and produces a Trees block as output. It filters trees or rooted networks returning all unique topologies (using hardwired clusters). The algorithm has the following options:

Unrooted = <Boolean> - ignore location of root

## Confidence Network

The *Confidence Network* algorithm takes a Trees block as input and produces a Splits block as output. It computes a credibility network using Beran's algorithm. The algorithm has the following options:

Level = <Double> - set the level (between 0 and 1)

HighDimensionFilter = <Boolean> - heuristically remove splits causing high-dimensional consensus network

Reference: [Huson and Bryant, 2006]

## Phylogenetic Diversity

The *Phylogenetic Diversity* algorithm takes a Trees block as input and produces a Report block as output. It calculates the phylogenetic diversity for selected taxa. The algorithm has the following options:

Rooted = <Boolean> - interpret trees as rooted?

ApplyTo = {OneTree | AllTrees} - determine whether to apply to one or all trees

WhichTree = <Integer> - the index of the tree that the method will be applied to

Reference: [Faith, 1992]

### Robinson Foulds Distances

The *Robinson Foulds Distances* algorithm takes a Trees block as input and produces a Report block as output. It calculates the Robinson-Foulds distance between each pair of trees. The algorithm has the following options:

Normalize = <Boolean> - normalized distances

Reference: [?]

### Tree Diversity Index

The *Tree Diversity Index* algorithm takes a Trees block as input and produces a Report block as output. It calculates the fair-proportion and equal-splits values on trees. The algorithm has the following options:

Method = {FairProportions | EqualSplits} - choose the type of index calculation

ApplyTo = {OneTree | AllTrees} - determine whether to apply to one or all trees

WhichTree = <Integer> - the index of the tree that the method will be applied to

References: [Redding, 2003, Redding and Mooers, 2006]

### Unrooted Shapley Values

The *Unrooted Shapley Values* algorithm takes a Trees block as input and produces a Report block as output. It calculates unrooted Shapley values on trees. The algorithm has the following options:

ApplyTo = {OneTree | AllTrees} - determine whether to apply to one or all trees

WhichTree = <Integer> - the index of the tree that the method will be applied to

Reference: [Haake et al., 2008]

### Average Distances

The *Average Distances* algorithm takes a Trees block as input and produces a Distances block as output. It calculates the average distances between taxa over a set of trees.

Reference: [Lapointe and Cucumel, 1997]

### Enumerate Contained Trees

The *Enumerate Contained Trees* algorithm takes a Trees block as input and produces a Trees block as output. It enumerates all contained trees. The algorithm has the following options:

RemoveDuplicates = <Boolean> - suppress duplicate trees in output

## Loose And Lacy

The *Loose And Lacy* algorithm takes a Trees block as input and produces a Trees block as output. It computes the 'loose' and 'lacy' species for a given tree and taxon trait. The algorithm has the following options:

SpeciesDefinition = {Loose | Lacy | Both} - species definition to use

TraitNumber = <Integer> - number of specific trait to use

UseAllTraits = <Boolean> - use all traits

Reference: [Hoppe et al., 2019]

## List One RSPR Trees

The *List One RSPR Trees* algorithm takes a Trees block as input and produces a Report block as output. It determines which trees are exactly on rSPR away from each other. The algorithm has the following options:

ApplyTo = {OneTree | AllTrees} - determine whether to apply to one or all trees

WhichTree = <Integer> - the index of the tree that the method will be applied to

# C.5 Algorithms on a Network Block

## Show Network

The *Show Network* algorithm takes a Network block as input and produces a View block as output. It provides interactive visualizations of networks. The algorithm has the following options:

View = {Network | Text} - the type of network viewer to use

## Appendix D

# Supported import and export formats

The program support several widely-used import and export formats.

### D.1 Supported import formats

#### D.1.1 Importers for a characters block

Can import characters data in the following formats: FastA, MSF, Stockholm, Nexus, Phylip.

#### D.1.2 Importers for a distances block

Can import distances data in the following formats: Nexus, Phylip, CSV.

#### D.1.3 Importers for a trees block

Can import trees data in the following formats: Newick, NEXML, Nexus.

#### D.1.4 Importers for a splits block

Can import splits data in the following formats: Newick, Nexus.

#### D.1.5 Importers for a network block

Can import network data in the following formats: Nexus.

The program also import a network given in GML format, if the network was previously exported in the format from SplitsTree. To import GML from other sources, please note that some of the nodes must have taxon names given using the key "label" and edges can be given lengths using the key "weight". The graph must be connected otherwise it cause an exception.

### **D.1.6 Importers for a genomes block**

Can import genomes data in the following formats: Nexus.

## **D.2 Supported output formats**

### **D.2.1 Exporters for a taxa block**

Can export taxa data in the following formats: Nexus.

### **D.2.2 Exporters for a characters block**

Can export characters data in the following formats: Clustal, FastA, Nexus, Phylip.

### **D.2.3 Exporters for a distances block**

Can export distances data in the following formats: Nexus, Phylip.

### **D.2.4 Exporters for a trees block**

Can export trees data in the following formats: NeXML, Newick, Nexus.

### **D.2.5 Exporters for a splits block**

Can export splits data in the following formats: FastA, Newick, Nexus.

### **D.2.6 Exporters for a network block**

Can export network data in the following formats: Nexus.

### **D.2.7 Exporters for a genomes block**

Can export genomes data in the following formats: Nexus.

### **D.2.8 Exporters for a view block**

Can export view data in the following formats: GML, Nexus.

## D.3 Taxon display labels import

Taxon display labels can be imported from a text file. Each line of the file must contain two tab-separated entries. The first entry is the taxon name, as used in the input data, and the second entry is the corresponding display label, which may contain HTML formatting.

Here is an example. The first label has font size 24 and text color blue. The second label is shown in bold. The remaining four labels have a yellow background.

```
Co90-125      <size "24"><c blue>Co90-125
s428          <b>s428
s421          <bg yellow>s421
s433          <bg yellow>s433
s434          <bg yellow>s434
s498          <bg yellow>s498
```

## D.4 Traits import

Traits can be imported from a text file. The first line of the file must define the names of the traits. The line must start with the keyword **Traits** and then must contain a list of the names of the different traits, separated by tabs

Then there must be one line for each taxon. The first entry must be the taxon name and this must be followed by one value for each of the listed traits, separated by tabs.

Here is an example defining five traits, Europe to America, for seven taxa seq\_1 to seq\_7. In this case, the trait values are counts.

The second line starting with the key word **Coordinates** is optional. When present, it provides the latitude and longitude associated with each trait.

| Traits      | Europe   | Asia | Africa      | Australia | America             |
|-------------|----------|------|-------------|-----------|---------------------|
| Coordinates | 53,16.75 |      | 43.68,87.33 | 5.4,26.5  | -25.61,134.35 0,-76 |
| seq_1       | 0        | 0    | 0           | 3         | 3                   |
| seq_2       | 10       | 5    | 0           | 6         | 0                   |
| seq_3       | 0        | 0    | 0           | 3         | 5                   |
| seq_4       | 0        | 0    | 0           | 4         | 2                   |
| seq_5       | 4        | 0    | 10          | 0         | 0                   |
| seq_6       | 0        | 0    | 0           | 7         | 3                   |
| seq_7       | 0        | 0    | 5           | 0         | 0                   |

The program also supports a second way of specifying taxon-trait associations. After specifying the first (one or two) lines, the taxon-trait counts can also be specified by listing a taxon, a trait and then the desired count, like this:

| Traits      | Europe    | Asia | Africa      | Australia | America             |
|-------------|-----------|------|-------------|-----------|---------------------|
| Coordinates | 53,16.75  |      | 43.68,87.33 | 5.4,26.5  | -25.61,134.35 0,-76 |
| seq_1       | Australia | 3    |             |           |                     |
| seq_1       | America   | 3    |             |           |                     |
| seq_2       | Europe    | 10   |             |           |                     |
| seq_2       | Asia      | 5    |             |           |                     |
| seq_2       | Australia | 6    |             |           |                     |
| ...         |           |      |             |           |                     |





# Appendix E

## Workflow

SplitsTree is designed around the concept of a workflow. This is a provenance graph in which nodes explicitly represent data blocks and algorithms.

### E.1 Input and working nodes

The graph is displayed the workflow tab (see Section 1.13) and is also presented in the sidebar (see Section 1.18). While the casual user will use menu items to set up and change the graph (without being aware of the graph), a user more familiar with the program will use controls in the workflow tab and sidebar to explicitly add, delete, duplicate or modify nodes and edges in the workflow graph.

Each main window contains one workflow and the workflow represents one input dataset, all applied algorithms and derived data.

In more detail, the graph is a tree with two root nodes (see Fig. E.1).

The first root node represents the set of input taxa. The number of input taxa is fixed and each taxon has a unique name (a label that does not contain special characters such as a single or double quote and must not be a number.) In addition, each taxon can have a display label, which may contain certain HTML commands, that is used to draw the label associated with the taxon.

The second root node represents the input data. This may be a set of characters (or sequence alignment), a distance matrix, a collection of trees (or rooted networks), or a set of splits.

The input-taxa node has one child, the taxa filter node. This can be used to remove some of the input taxa. This node has one child that contains the set of working taxa. This node represents the set of taxa that are actually used in all computations.

The input data node has one child that contains the working data. The data associated with this node is copied from the input data node, removing any taxa that have been deactivated using the taxa filter node.

All calculations undertaken in the program are based on the set of working taxa and the working data.

If the input data is character data (or a multiple sequence alignment), then the input data is displayed in the alignment tab (see Section 1.5) and controls associated with the tab allow the user to add or remove taxa in the same manner as when using the taxa filter.

The taxa filter also allows the user to edit the display labels associated with taxa.

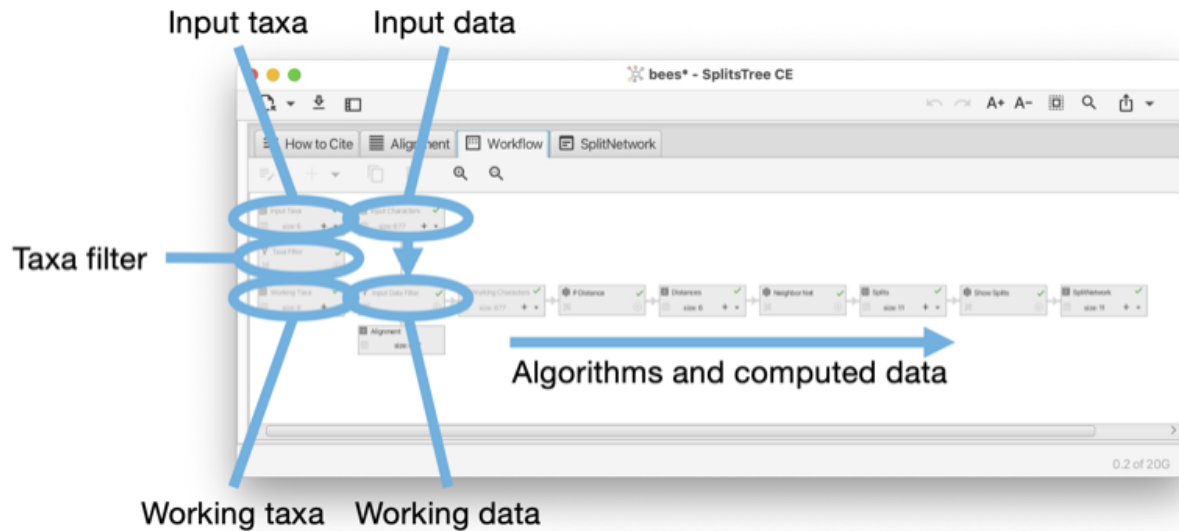


Figure E.1: The workflow is a directed tree with two root nodes, the first representing the set of input taxa and the second representing the input data. The taxa filter node can be used to remove input taxa and thus produces the working taxa node. Filtering taxa also has the effect of removing the corresponding data from the input data node and gives rise to the working data node. Subsequent algorithms are run on the data represented by the working taxa and working data nodes.

## E.2 Data and algorithm nodes

The workflow contains two main types of nodes, algorithm nodes and data nodes. The workflow is a bipartite graph: data nodes only have algorithm nodes as children and vice versa (see Fig. E.2).

In the Figure (see Fig. E.1) it looks like the input data node and working data node are directly connected to each other. However, actually there is a special algorithm node between the two that facilitates the filtering of data when the taxa filter is used. As this node is only used internally, it is not displayed in the workflow tab or sidebar.

Each algorithm node computes data that is based on to a data node, while each data node is provided with the set of working taxa and one parental data node to work with. (Some algorithms, such as those that perform bootstrapping, additionally access other ancestral nodes to perform their calculations.)

The final nodes of the workflow (leaves) are always data nodes and each is of one of two special kinds. First, there are view nodes that represent the graphical visualization of trees or networks. Second, there are reporter nodes that are used to present the textual output of a calculation, such as the computation of Tajima's D.

An algorithm node can represent one of several algorithms, depending on the type of input data and output data. For example, there are three algorithms that take as input a distance matrix and produce, as output, a set of splits, namely neighbor-net, split decomposition and the Buneman tree algorithm. In the Figure (see Fig. E.3) we show such an algorithm node and the controls associated with it.

A data node usually represents a block of data, which can be either characters (aligned sequences), a distance matrix, a set of splits, a collection of trees (and/or rooted networks) or a haplotype network (see Fig. E.4). A view node is a data node that corresponds to a viewer for trees or networks.

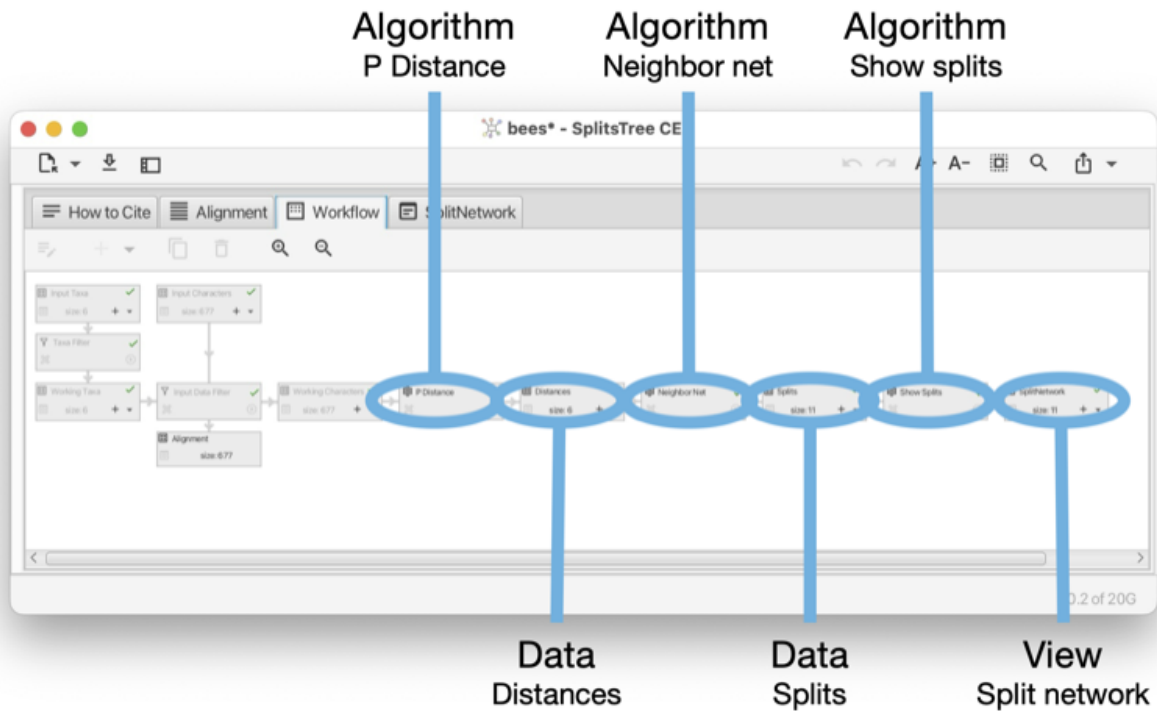


Figure E.2: Here we show the workflow that shows a split network computed from characters data. The workflow contains chains of alternating algorithm and data nodes. The characters data (working data node) is followed by the  $p$ -distance calculation, giving rise to a data node containing distances. The distances are provided to the neighbor-net algorithm, giving rise to a set of splits. The splits are passed to the show splits node, which computes the visualization, which is represented by the splits view node.



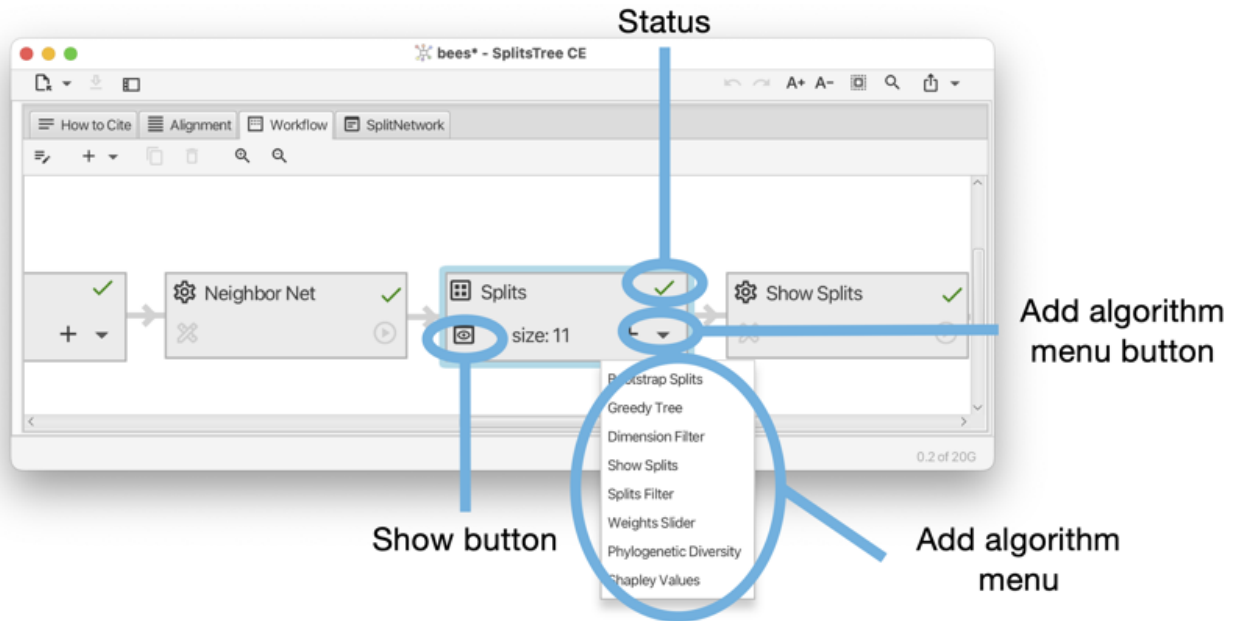


Figure E.4: A data node has a drop-down menu that allows one to attach a new algorithm to the node, thus starting a new chain of nodes. The show button opens a tab that provides a textual representation of the data. The status icon whether the data is up-to-date (green tick), is waiting to be updated (yellow clock) or is missing due to a failed or canceled algorithm (red cross).

### E.3 Exporting the workflow

When saving a document, the workflow, including all input data and computed data, are saved to a file to be opened later. Such a full file has file suffix `.stree6`.

In addition, using the `File->Export->Workflow...` menu item, the user can save just the workflow graph, without the current data, to a file. Such a workflow file has file suffix `.wflow6`.

A saved workflow can be reopened in SplitsTree and data can then be loaded using the `File->Replace Data...` menu item.

### E.4 Running a workflow on multiple datasets

To run an analysis on multiple datasets, the SplitsTree packages comes with a commandline tool called `run-workflow`, located in the `tools` directory. The basic idea is as follows. First, use SplitsTree to interactively set up the desired analysis. Then save export the workflow to a file(see Section E.3). Then use the `run-workflow` program to apply the workflow to multiple datasets.

The program is configured using a number of commandline options, to set the workflow file, to set the input data files, to set the output file or files, and to determine what should be written to the output.

Here is a synopsis of the program:

```
SYNOPSIS
  RunWorkflow [options]
DESCRIPTION
```

Runs a SplitsTree6 workflow on input data  
OPTIONS

Input Output:

|                          |  |
|--------------------------|--|
| -w, --workflow [string]  | File containing SplitsTree6 workflow. Mandatory option.  |
| -i, --input [string(s)]  | File(s) containing input data (or directory). Mandatory option.  |
| -f, --format [string]    | Input format. Default: Unknown. Legal values: 'Unknown' 'FastA' 'MSF' 'Newick' 'Nexml' 'Nexus' 'Phylip' 'Stockholm'          |
| -o, --output [string(s)] | Output file(s) (or directory or stdout). Default value(s): 'stdout'  |
| -n, --node [string]      | Title of node to be exported (if none given, will save whole file).  |
| -e, --exporter [string]  | Name of exporter to use. Legal values: 'Clustal' 'FastA' 'GML' 'NeXML' 'Newick' 'Nexus' 'Phylip' 'PlainText' 'NexusWithTaxa' |

Other:

|                               |   |
|-------------------------------|---|
| -x, --inputExt [string]       | File extension for input files (when providing directory for input).          |
| -r, --recursive               | Recursively visit all sub-directories. Default: false.                        |
| -t, --time [string]           | Maximum wall-clock time to run (e.g. 100s, 2m, 3h or 4d). Default: unlimited. |
| -p, --propertiesFile [string] | Properties file.  |
| -s, --silent                  | Silent mode (hide all stderr output). Default: false.                         |
| -v, --verbose                 | Echo commandline options and be verbose. Default: false.                      |
| -h, --help                    | Show program usage and quit.  |

Here is a summary of the most important options:

- Use the -w option to specify the workflow file.
- Use the -i option to list the input files or a directory that contains the input files.
- Use the -f option to specify the file format of the input files.
- Optionally, use the -x option to specify the file suffixes of the input files, if a specified directories contains some files that should not be used.
- Optionally, use the -r option to recursively visit all directories contained in the specified input directory.
- Use the -o option to specify the output file or files. If you specify one directory, then one output file per input file will be created and written to that directory. If you specify one file, then all output is written to that file. You can also specify one output file per input file. Use the keyword stdout to have all output written to the console. If output files end on .gz then they will be written in gzip format.
- Use the -n option to specify a specific data node to be saved to output. For example, if your analysis generates a trees block called Trees, then you can specify -n Trees to output the trees. If this option is not specified, then the whole workflow is output containing all input and computed data.
- Use the -e option to specify which exporter (and thus format) to use when generating output. This option is ignored if the -n is not specified.

# Appendix F

## Styling labels

In SplitsTree, each taxon is represented by a unique label. These labels are specified in the input taxa block. In addition to these labels, the taxon block may also contain a set of “display labels”, one for each taxon. If provided, the display labels are used in drawings of trees and networks. Display labels can be styled using a set of HTML tags. There are several ways to do this:

- Use the side panel (on the right) to choose a font, font size, text color, background color or mark. Each of these interactive choices produce corresponding HTML tags in the display labels.
- Right-clicking on a taxon label displayed in a tree or network will open an input dialog for the display label. The dialog allows the user to edit the display label any changes to the label are immediately shown in a preview pane.
- In the TaxaFilter tab, there is an entry for each taxon, which can be edited.
- Display labels can be imported from a file (and exported to a file for reuse). Each line of the file must contain a key and a value separated by a tab. The key is the unique taxon name and the value is the desired display label.

Here is a list of all supported HTML tags, most are standard HTML, a few are specific to SplitsTree:

- `<i>text</i>` - display enclosed text in italics,
- `<b>text</b>` - display enclosed text in bold,
- `<sup>text</sup>` - display enclosed text as super-script,
- `<sub>text</sub>` - display enclosed text as super-script,
- `<u>text</u>` - display enclosed text as underlined,
- `<a>text</a>` - display enclosed text using strike-through,
- `<br>` - add a new-line,
- `<font "name">text</font>` - display enclosed text using the named font,
- `<size "value">text</size>` - display enclosed text using the given font size value,
- `<c "value">text</c>` - display enclosed text using the given color value,
- `<bg "value">background-color</bg>` - display enclosed text using the given background color value,
- `<mark shape="value" width="value" height="value" fill="color" stroke="color">` - adds a mark to text, using the provided shape, width, height, fill and stroke colors, and

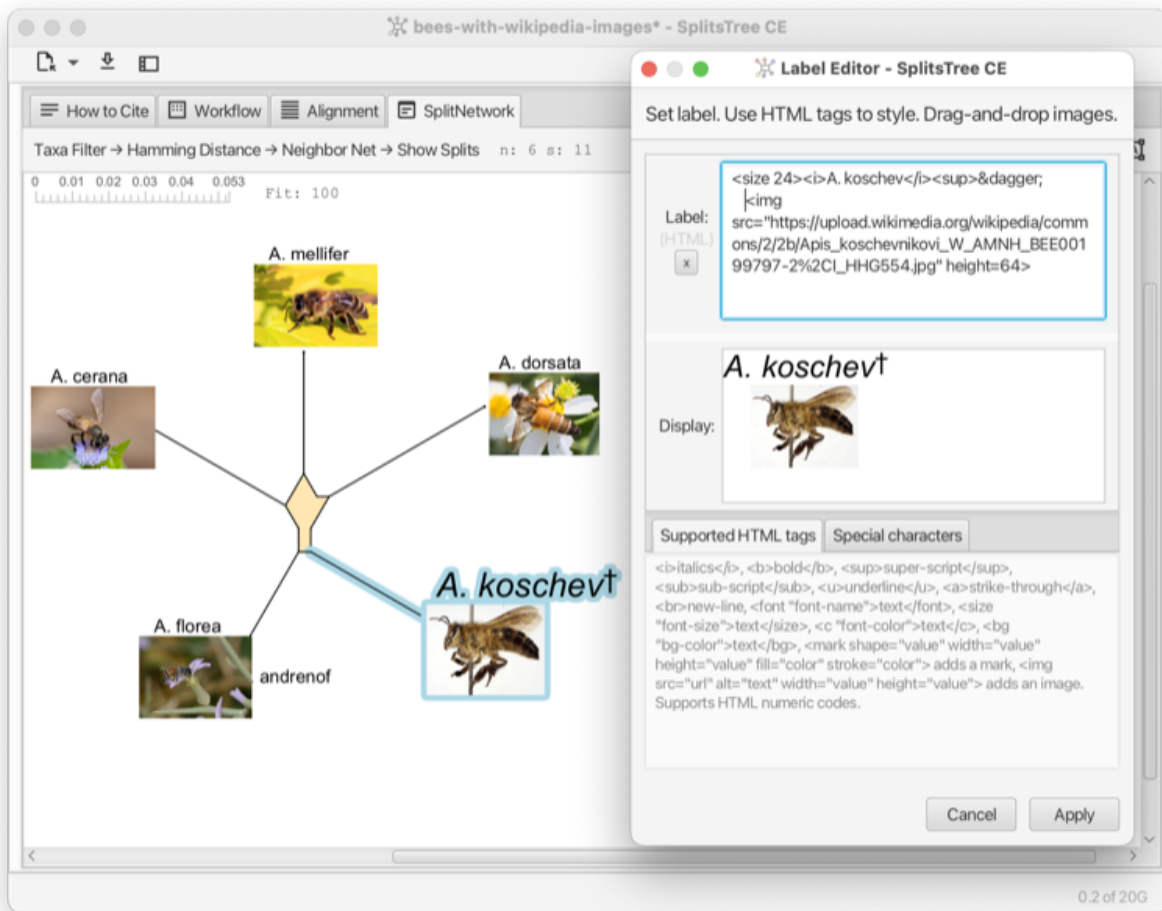


Figure F.1: Here, for all taxa (except for *A. Andrenof*), the display label includes a Wikipedia image of the represented species of bees. By right-clicking on the *A. Koschevt* label, the display-label editor was opened for this label.

- `` - adds an image using the provided URL and width and/or height.

HTML numerical codes can be used to specify characters.

In the Figure (see Fig. F.1) a display label has been set for each taxon (except for *A. Andrenof*). Using HTML, the display labels include a Wikipedia image of the represented species of bees. In addition, the display label editor is shown open for *A. Koschevt*. The display label contains HTML tags for italics and a tag to start a superscript. The HTML entity `&dagger;` is used to display a dagger.



## Appendix G

# Additional parameters

The layout algorithm for rooted phylogenetic networks and for tanglegrams uses simulated annealing for nodes of large outdegree. The default parameters are: start temperature = 1000, end temperature = 0.01, 1000 iterations per temperature step, cooling rate = 0.95.

We do not expose these parameters in the UI, however, if you really want to change these, then edit the properties file `SplitsTree6.def` (its location is system specific, either `~/Library/Preferences/SplitsTree6.def` or `~/SplitsTree6.def`) and write statements like this:

- `SA_DEFAULT_START_TEMPERATURE=2000,`
- `SA_DEFAULT_END_TEMPERATURE=1,`
- `SA_DEFAULT_ITERATIONS_PER_TEMPERATURE=100` and
- `SA_DEFAULT_COOLING_RATE=0.80,`

to change the values to 2000, 1, 100 and 0.8, say, respectively.

Additionally, the computation of tanglegrams for rooted networks use 32 different random starting configurations, by default. This number can be changed by adding this:

- `TANGLEGRAM_PARALLEL_JOBS=1024,`

to use 1024 random starting configurations, say.

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