

## ScripTree: scripting phylogenetic graphics

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### ABSTRACT

**Summary:** There is a large amount of tools for interactive display of phylogenetic trees. However, there is a shortage of tools for the automation of tree rendering. Scripting phylogenetic graphics would enable the saving of graphical analyses involving numerous and complex tree handling operations and would allow the automation of repetitive tasks. ScripTree is a tool intended to fill this gap. It is an interpreter to be used in batch mode. Phylogenetic graphics instructions, related to tree rendering as well as tree annotation, are stored in a text file and processed in a sequential way.

**Availability:** ScripTree can be used online or downloaded at [www.scriptree.org](http://www.scriptree.org), under the GPL license.

**Implementation:** ScripTree, written in Tcl/Tk, is a cross-platform application available for Windows and Unix-like systems including OS X. It can be used either as a stand-alone package or included in a bioinformatic pipeline and linked to a HTTP server.

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### 1 PHYLOGENETIC GRAPHICS

Phylogenetic graphics deals with basic operations on trees (e.g. rooting) and tree rendering processes (e.g. annotation) in the context of large trees and/or collection of trees. Phylogenetic graphics resorts to dynamic information visualization techniques such as ‘focus+context’ magnifying features. Tree annotation consists in the highlighting (coloring, posting of text or symbols) of subtrees or leaf labels according to additional information (e.g. taxonomy, geography, gene function, etc.) related to the entities under study (molecular sequences, species, etc.). Tools like TreeJuxtaposer (Munzner *et al.*, 2003), TreeDyn (Chevenet *et al.*, 2006), Dendroscope (Huson *et al.*, 2007) are examples of tree editors with phylogenetic graphic capabilities. A new challenge in the field is the automation of a graphical analysis encoded as a sequence of operations that precisely describes the way to display and tag trees with additional information. These operations are stored in a script, usable on the same or different datasets. Moreover, scripting is a flexible approach allowing computations to be run either as a local stand-alone process or incorporated within a pipeline and potentially accessible through a web interface. Currently, there is a need for such automation of phylogenetic graphics as we see more and more

web sites that provide access to bioinformatic analyses displaying trees, e.g. PhylomeDB (Huerta-Cepas *et al.*, 2007), phylogeny.fr (Dereeper *et al.*, 2008), PhyloExplorer (Ranwez *et al.*, 2009). Yet, few existing tools have scripting capabilities—ATV/Archeopteryx (Zmasek *et al.*, 2001), TreeGraph (Muller *et al.*, 2004), Ape (Paradis *et al.*, 2004), TreeDyn (Chevenet *et al.*, 2006), Dendroscope (Huson *et al.*, 2007) or ETE (Huerta-Cepas *et al.*, 2010). Archeopteryx displays single trees in interactive manipulations. Hence, this is not the tool of choice for automatically rendering tree collections with complex annotations. ETE offers elaborated features for analysis of trees as well as automation and visualization of trees. It is a powerful programmable toolkit, but requires object oriented Python programming skills to annotate trees in an automated way. ScripTree has tree *annotation* features more elaborated than the TreeDyn ones. ScripTree is a higher level interpreter, including numerous and specific annotation commands. It is dedicated to automation and, hence, does not contain a graphical user interface.

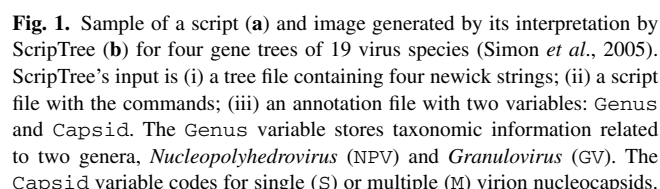
### 2 SCRIPTREE INPUT/OUTPUT

The basic ScripTree command line is `scriptree -tree file.nwk [options]`. The `-tree` argument refers to a file containing one or more newick strings encoding trees. Without any other specification, ScripTree uses default settings for rendering these trees. Different output file formats are available: PostScript, SVG, PNG and TGF. The SVG format can be displayed by web browsers and edited with drawing programs such as InkScape. The TGF format is useful for an interactive post-processing using the TreeDyn editor. A first optional flag is `-script file.txt` indicating the file containing rendering and annotation commands. The specificity of ScripTree is to take into account additional information into the tree rendering process, indicated by a second optional flag, `-annotation file.txt`. This information is given in a tabular CSV format: annotation variables as columns, and rows related to leaves or internal nodes of the trees. Separating annotations from the tree encoding allows the latter to be compatible with the newick format outputted by common phylogenetic inference programs and the former to be reused on other tree collections.

### 3 SCRIPTREE COMMANDS

ScripTree commands are divided into three families: edition, projection and identification. In the following, we present examples of commands belonging to these families (Fig. 1a) and apply them

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The *edition* family acts on trees as a whole, specifying: (i) global tree rendering such tree size, leaf label font, organization of a tree collection into rows and columns; (ii) tree manipulations such as branch swapping and rerooting. For instance, command (1) in Figure 1a sets the size of the trees to  $80 \times 150$  pixels, organizes them as a two-by-two matrix, displays them with a rectangular shape accounting for branch lengths (`-conformation 1`) and finally roots the trees at the smallest subtree containing all leaf labels that begin with Cn (CnA and/or CnB depending on the trees).

The *identification* commands which enable to highlight only parts of the trees, identified by a query, operate either on the

## 4 CONCLUSION

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