

# **Tutorial: Environment for Tree Exploration**

Release 2.3.6

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# Contents

1	Char	ngelog history	3
	1.1	What's new in ETE 2.3	3
	1.2	What's new in ETE 2.2	5
	1.3	What's new in ETE 2.1	9
2	The 1	ETE tutorial	13
	2.1	Working With Tree Data Structures	13
	2.2	The Programmable Tree Drawing Engine	
	2.3	Phylogenetic Trees	
	2.4	Clustering Trees	
	2.5	Phylogenetic XML standards	
	2.6	Interactive web tree visualization	93
	2.7	Testing Evolutionary Hypothesis	95
	2.8	Dealing with the NCBI Taxonomy database	
	2.9	SCRIPTS: orthoXML	
3	ЕТЕ	's Reference Guide	115
	3.1	Master Tree class	115
	3.2	Treeview module	
	3.3	PhyloTree class	
	3.4	Clustering module	
	3.5	Nexml module	
	3.6	Phyloxml Module	
	3.7	Seggroup class	
	3.8	WebTreeApplication object	
	3.9	EvolTree class	
	3.10	NCBITaxa class	
Bi	bliogr	aphy	169
Рy	thon I	Module Index	171
In	dex		173

[Download PDF documentation]  $\parallel$ 

Contents 1

2 Contents

# **Changelog history**

# 1.1 What's new in ETE 2.3

# 1.1.1 Update 2.3.2

- added NCBITaxa.get\_descendant\_taxa()
- added NCBITaxa.get common names()
- ete ncbiquery: dump descendant taxa given a taxid or taxa name. new option '-descendants'\_; renamed '-taxonomy'\_ by '-tree'\_
- fixes <misaligned branches <a href="https://github.com/jhcepas/ete/issues/113">https://github.com/jhcepas/ete/issues/113</a> '\_ in ultrametric tree images using vt\_line\_width > 0
- fixes <windows installation problem <a href="https://github.com/jhcepas/ete/issues/114">https://github.com/jhcepas/ete/issues/114</a>

#### 1.1.2 New Modules

# tools

A collection of command line tools, implementing common tree operations has been added to the ETE core package. All tools are wrapped by the **ete** command, which should become available in your path after installation.

- ete build: Build phylogenetic trees using a using a number of predefined built-in gene-tree and species-tree workflows. Watch example
- ete view: visualize and generate tree images directly form the command line.
- ete compare: compare tree topologies based on any node feature (i.e. name, species name, etc) using the Robinson-Foulds distance and edge compatibility scores.
- ete ncbiquery: query the ncbi taxonomy tree directly from the database.
- **ete mod**: modify tree topologies directly from the command line. Allows rooting, sorting leaves, pruning and more
- ete annotate: add features to the tree nodes by combining newick and text files.
- ete generate: generate random trees, mostly for teaching and testing

# ncbi taxonomy

The new **ncbi\_taxonomy** module provides the class NCBITaxa, which allows to query a locally parsed NCBI taxonomy database. It provides taxid-name translations, tree annotation tools and other handy functions. A brief tutorial and examples on how to use it is available here

#### 1.1.3 New features

#### **News in Tree instances**

- added TreeNode.iter\_edges() and TreeNode.get\_edges()
- added TreeNode.compare() function
- added TreeNode.standardize() utility function to quickly get rid of multifurcations, single-child nodes in a tree.
- added TreeNode.get\_topology\_id() utility function to get an unique identifier of a tree based on their content and topology.
- added TreeNode.expand\_polytomies()
- improved TreeNode.robinson\_foulds() function to auto expand polytomies, filter by branch support, and auto prune.
- improved TreeNode.check\_monophyly() function now accepts unrooted trees as input
- Default node is set to blank instead of the "NoName" string, which saves memory in very large trees.
- The branch length distance of root nodes is set to 0.0 by default.
- newick export allows to control the format of branch distance and support values.
- Tree and SeqGroup instances allow now to open gzipped files transparently.

#### News in the treeview module

- improved SVG tree rendering
- improved random\_color() function (a list of colors can be fetch with a single call)
- improved SeqMotifFace
- Added RectFace
- Added StackedBarFace

# 1.1.4 Highlighted Bug Fixes

- Newick parser is now more strict when reading node names and branch distances, avoiding silent errors when parsing node names containing illegal symbols (i.e. ][)(,: )
- fixes several minor bugs when retrieving extra attributes in PhyloNode.get\_speciation\_trees().
- Tree viewer crashes when redrawing after changing node properties.
- fixed installation problem using pip.

- visualizing internal tree nodes as a circular tree produce crashes
- math domain error in SequencePlotFace.
- Fix likelihood calculation bug in EvolTree
- Fix BarChartFace problem with negative numbers
- Fix problem that produced TreeStyle attributes to be ignored in PhyloTree instances.

# 1.2 What's new in ETE 2.2

#### 1.2.1 BUGFIXES

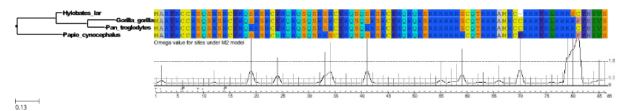
- Fixes in NeXML parser and exporting functions
- Fixed 'paste newick' functionality on the GUI
- Fixed PhyloNode.is\_monophyletic() and moved to TreeNode.check\_monophyly().
- Fixed consistency issued in TreeNode.sort\_descendants() function.

#### 1.2.2 SCRIPTS

- Improvements in the standalone visualization script (a.k.a. ete2)
- Added the *etree2orthoxml script*, which provides conversion between phylogenetic tree and the orthoXML format

#### 1.2.3 NEW MODULES

• New EvolNode tree object type is available as a part of adaptation-test extension recently developed by François Serra (see *Testing Evolutionary Hypothesis* in the tutorial).



# 1.2.4 NEW FEATURES

- News in core Tree instances:
  - Added TreeNode.robinson\_foulds() distance to compare the topology of two trees (i.e. tree.robinson\_foulds(tree2)). It includes automatic pruning to compare trees of different sizes. See tutorial and examples
  - Added new options to TreeNode.copy() function, allowing faster methods to duplicate tree node instances. See tutorial and examples

- Added preserve\_branch\_length argument to TreeNode.prune() and TreeNode.delete(), which allows to remove nodes from a tree while keeping original branch length distances among the remaining nodes.
- Added TreeNode.resolve\_polytomy() function to convert multifurcated nodes into an arbitrary structure of binary split nodes with distance. *See tutorial and examples*
- Added TreeNode.get\_cached\_content() function, which returns a dictionary linking each node instance with its leaf content. Such a dictionary might be used as a cache to speed up functions that require intensive use of node traversing. See tutorial and examples
- Improved TreeNode.get\_ascii() function for text-based visualization of trees. A
  new attributes argument can be passed to display node attributes within the ASCII tree representation.

- Random branch length and support values generation is now available for the TreeNode.populate() function.
- a new argument is\_leaf\_fn is available for a number of traversing functions, thus allowing to provide custom stopping criteria when browsing a tree. This is, any node matching the function provided through the is\_leaf\_fn argument will be temporarily considered as a terminal/leaf node by the traversing function (tree will look as a pruned version of itself). See tutorial and examples
- Added TreeNode.iter\_ancestors() and TreeNode.get\_ancestors() functions.
- Added TreeNode.iter\_prepostorder() tree node iterator.
- Newick parser accepts now the creation of single node trees. For example, a text string such
  as "node1;" will be parsed as a single tree node whose name is node1. By contrast, the
  newick string (node1); will be interpreted as an unnamed root node plus a single child
  named name1.
- TreeNode.write() accepts now a format\_root\_node argument to export root node features as a part of the newick string.
- The new TreeNode.check\_monophyly() method allows to check if a node is mono, poly or paraphyletic for a given attribute and values (i.e. grouped species). Although monophyly is actually a phylogenetic concept, the idea can be applied to any tree, so any topology could be queried for the monophyly of certain attribute values. If not monophyletic, the method will return also the type of relationship connecting the provided values (para- or poly-phyletic). See tutorial and examples

 New TreeNode.get\_monophyletic() method that returns a list of nodes in a tree matching a custom monophyly criteria.

# • News PhyloTree instances:

 Added PhyloNode.get\_speciation\_trees() method, which returns all possible species topologies present in a gene family tree as described in Treeko. See tutorial and examples

See also:



TreeKO: a duplication-aware algorithm for the comparison of phylogenetic trees.

Marcet-Houben M, Gabaldón T.

Nucleic Acids Res. 2011 May;39(10):e66. doi: 10.1093/nar/gkr087.

- Added PhyloNode.split\_by\_dups() method, which returns a list of partial subtrees resulting from splitting a tree at duplication nodes. *See tutorial and examples*
- Added PhyloNode.collapse\_lineage\_specific\_expansions() method, which returns a pruned version of a tree, where nodes representing lineage specific expansions are converted into a single leaf node. See tutorial and examples

# • News on sequence and multiple sequence alignment parsing:

 added the option to disable the automatic correction of duplicated names when loading SeqGroup data from phylip and fasta files.

# • News on tree visualization and image rendering:

- node style attributes can now be modified without the need of initialization by directly accessing the TreeNode.img\_style attribute.
- Multiple layout functions can now be provided to combine their functionality. This way, you
  can keep separate styling templates and combine them as necessary.

```
from ete2 import TreeStyle

def color_leaves(node):
    if node.is_leaf():
        node.img_style["fgcolor"] = "red"

def size_internal(node):
    if not node.is_leaf():
        node.img_style["size"] = 15

ts = TreeStyle()
# provide a list of layout functions, instead of a single one
ts.layout_fn = [color_leaves, size_internal]
```

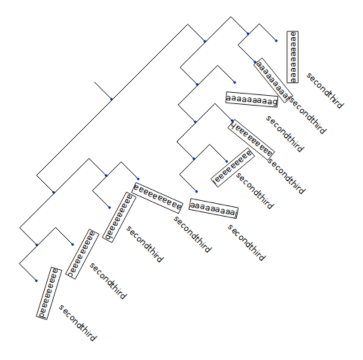
COLOR\_SCHEMES and SVG\_COLORS dictionaries are provided for easy access to color codes and several predefined color schemes. In addition, a random\_color() function is also available as a generator of RGB colors (where saturation and lightness can be fixed).

```
from ete2 import random_color, COLOR_SCHEMES, SVG_COLORS

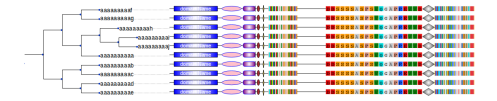
# generate 20 random colors
node_colors = [random_color(s=0.4, l=4) for i in xrange(20)]
```

#### • News on node faces:

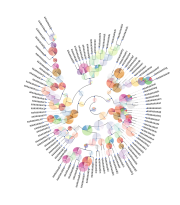
- New face.rotation attribute, that allows to rotate individual faces even when a global treestyle.rotation is used.

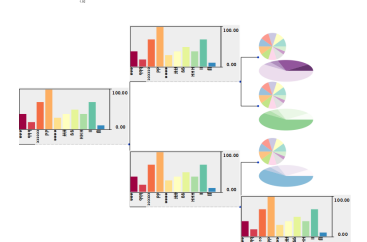


- Improved SequenceFace: Sequence sites are now rendered one by one, allowing interaction with each of them and getting rid of the previous pixmap size limitation. Site image dimensions and colours are now configurable.
- Added new SeqMotifFace class, which represent an enriched version of the former SequenceFace instance. This new face type allows to represent sequences as a succession of domain/motif elements or to represent sequence positions as color points. Gaps can also be taken into account and therefore shown as as a black space or a flat line.



- Added PieChartFace and BarChartFace face types for built-in representation of statistics attached to nodes.
- Improved ImgFace class, now accepting on the fly image scaling.





#### · News on the GUI

- Allows image region selection.
- Allows zooming on selected regions or specific nodes (Z zoomIn, X zoomOut, R focus region).
- C-c will now interrupt the GUI application when started from a terminal.
- Added keyboard-based node navigation (click on a node and play the arrow keys).

# 1.3 What's new in ETE 2.1

- A basic standalone tree visualization program called "ete2" is now installed along with the package.
- The drawing engine has been completely rewritten to provide the following new features:
  - Added TreeStyle class allowing to set the following
    - \* Added circular tree drawing mode
    - \* Added tree *title face block* (Text or images that rendered on top of the tree)
    - \* Added tree legend face block (Text or images that rendered as image legend)
    - \* Added support for tree rotation and orientation

- \* Possibility of drawing aligned faces as a table
- \* Added header and footer regions for aligned faces.
- \* And more! Check TreeStyle documentation
- Added new face positions float, branch-top and branch-bottom. See tutorial (*Node faces*) for more details.
- Added several Face attributes:
  - \* face border
  - \* face background color
  - \* left, right, top and bottom margins
  - \* face opacity
  - \* horizontal and vertical alignment (useful when faces are rendered as table)
- Added support for predefined NodeStyle, which can be set outside the layout function (allows to save and export image rendering info)

# - Added new face types:

- \* CircleFace (basic circle/sphere forms)
- \* TreeFace (trees within trees)
- \* StaticItemFace and DynamicItemFace (create custom and interactive Qt-GraphicsItems)

# - Improved faces:

- \* AttrFace accepts prefix and suffix text, as well as a text formatter function. fstyle argument can be set to italic
- \* TextFace: fstyle argument can be set to italic

# - Save and export images

- \* Added full support for SVG image rendering
- \* Added more options to the TreeNode.render() function to control image size and resolution
- Added support for SVG\_COLORS names in faces and node styles

#### • Core methods:

- Added TreeNode.copy(): returns an exact and independent copy of node and all
  its attributes
- Added TreeNode.convert\_to\_ultrametric(): converts all branch lengths to allow leaves to be equidistant to root
- Added TreeNode.sort\_descendants(): sort tree branches according to node names.
- Added TreeNode.ladderize(): sort tree branches according to partition size
- Added TreeNode.get\_partitions(): return the set of all possible partitions grouping leaf nodes

- Tree nodes can now be fully exported using cPickle
- Newick parser can read and export branch distances and support values using scientific notation
- TreeNode.swap\_childs() method has changed to TreeNode.swap\_children()
- Added ete2.nexml module (read and write nexml format)
- Added ete2.phyloxml module (read and write phyloxml format)
- Added ete2.webplugin module: Allows to create interactive web tree applications
- Tree visualization GUI checks now for newer version of the ETE package.
- Added PhylomeDB3Connector
- Added PhyloNode.get\_farthest\_oldest\_node() function, which allows to find the best outgroup node in a tree, even if it is an internal node.

# • Bug Fixes and improvements:

- Fix: TreeNode.get\_common\_ancestor() accepts a single argument (node or list of nodes) instead of a succession or nodes. It can also return the path of each node to the parent.
- Fix: Fast scroll based zoom-in was producing tree image inversions
- Fix: Phylip parser does not truncate long names by default
- Fix: "if not node" syntax was using a len(node) test, which made it totally inefficient. Now, the same expression returns always *True*
- Improvement: Traversing methods are now much faster (specially preorder and levelorder)
- Improvement: Faster populate function (added possibility of random and non-random branch lengths)
- Improvement: Faster prune function
- Improvement: unicode support for tree files
- Improvement: Added newick support for scientific notation in branch lengths

# • Improved documentation and examples:

- Online and PDF tutorial
- Better library reference
- A set of examples is now provided with the installation package and here

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The ETE tutorial

Contents:

# 2.1 Working With Tree Data Structures

# **Contents**

- Working With Tree Data Structures
  - Trees
  - Reading and Writing Newick Trees
    - \* Reading newick trees
    - \* Writing newick trees
  - Understanding ETE Trees
  - Basic tree attributes
    - \* Root node on unrooted trees?
  - Browsing trees (traversing)
    - \* Getting Leaves, Descendants and Node's Relatives
    - \* Traversing (browsing) trees
    - \* Advanced traversing (stopping criteria)
      - · Collapsing nodes while traversing (custom is\_leaf definition)
    - \* Iterating instead of Getting
    - \* Finding nodes by their attributes
      - · Search\_all nodes matching a given criteria
      - · Search nodes matching a given criteria (iteration)
      - · Find the first common ancestor
      - · Custom searching functions
      - · Shortcuts
  - Checking the monophyly of attributes within a tree
  - Caching tree content for faster lookup operations
  - Node annotation
  - Comparing Trees
    - \* Calculate distances between trees
    - \* Robinson-foulds distance
  - Modifying Tree Topology
    - \* Creating Trees from Scratch
    - \* Deleting (eliminating) and Removing (detaching) nodes
  - Pruning trees
  - Concatenating trees
  - Copying (duplicating) trees
  - Solving multifurcations
  - Tree Rooting
  - Working with branch distances
    - \* Getting distances between nodes
    - \* getting midpoint outgroup

# 2.1.1 Trees

Trees are a widely-used type of data structure that emulates a tree design with a set of linked nodes. Formally, a tree is considered an acyclic and connected graph. Each node in a tree has zero or more child nodes, which are below it in the tree (by convention, trees grow down, not up as they do in nature). A node that has a child is called the child's parent node (or ancestor node, or superior). A node has at most one parent.

The height of a node is the length of the longest downward path to a leaf from that node. The height of the root is the height of the tree. The depth of a node is the length of the path to its root (i.e., its root path).

- The topmost node in a tree is called the root node. Being the topmost node, the root node will not have parents. It is the node at which operations on the tree commonly begin (although some algorithms begin with the leaf nodes and work up ending at the root). All other nodes can be reached from it by following edges or links. Every node in a tree can be seen as the root node of the subtree rooted at that node.
- Nodes at the bottommost level of the tree are called leaf nodes. Since they are at the bottommost level, they do not have any children.
- An internal node or inner node is any node of a tree that has child nodes and is thus not a leaf node.
- A subtree is a portion of a tree data structure that can be viewed as a complete tree in itself. Any node in a tree T, together with all the nodes below it, comprise a subtree of T. The subtree corresponding to the root node is the entire tree; the subtree corresponding to any other node is called a proper subtree (in analogy to the term proper subset).

In bioinformatics, trees are the result of many analyses, such as phylogenetics or clustering. Although each case entails specific considerations, many properties remains constant among them. In this respect, ETE is a python toolkit that assists in the automated manipulation, analysis and visualization of any type of hierarchical trees. It provides general methods to handle and visualize tree topologies, as well as specific modules to deal with phylogenetic and clustering trees.

# 2.1.2 Reading and Writing Newick Trees

The Newick format is one of the most widely used standard representation of trees in bioinformatics. It uses nested parentheses to represent hierarchical data structures as text strings. The original newick standard is able to encode information about the tree topology, branch distances and node names. Nevertheless, it is not uncommon to find slightly different formats using the newick standard.

ETE can read and write many of them:

FOR-	DESCRIPTION	SAMPLE
MAT		
0	flexible with support values	((D:0.723274,F:0.567784)1.000000:0.067192,(B:0.279326,H:0.756049)1.00
1	flexible with internal node names	((D:0.723274,F:0.567784)E:0.067192,(B:0.279326,H:0.756049)B:0.807788)
2	all branches + leaf names + internal supports	((D:0.723274,F:0.567784)1.000000:0.067192,(B:0.279326,H:0.756049)1.00
3	all branches + all names	((D:0.723274,F:0.567784)E:0.067192,(B:0.279326,H:0.756049)B:0.807788)
4	leaf branches + leaf names	((D:0.723274,F:0.567784),(B:0.279326,H:0.756049));
5	internal and leaf branches + leaf names	((D:0.723274,F:0.567784):0.067192,(B:0.279326,H:0.756049):0.807788);
6	internal branches + leaf names	((D,F):0.067192,(B,H):0.807788);
7	leaf branches + all names	((D:0.723274,F:0.567784)E,(B:0.279326,H:0.756049)B);
8	all names	((D,F)E,(B,H)B);
9	leaf names	((D,F),(B,H));
100	topology only	((,),(,));

Formats labeled as *flexible* allow for missing information. For instance, format 0 will be able to load a newick tree even if it does not contain branch support information (it will be initialized with the default value). However, format 2 would raise an exception. In other words, if you want to control that your newick files strictly follow a given pattern you should use **strict** format definitions.

# Reading newick trees

In order to load a tree from a newick text string you can use the constructor TreeNode or its Tree alias, provided by the main module ete2. You will only need to pass a text string containing the newick structure and the format that should be used to parse it (0 by default). Alternatively, you can pass the path to a text file containing the newick string.

```
from ete2 import Tree

# Loads a tree structure from a newick string. The returned variable 't' is the root not
t = Tree("(A:1, (B:1, (E:1,D:1):0.5):0.5);")

# Load a tree structure from a newick file.
t = Tree("genes_tree.nh")

# You can also specify the newick format. For instance, for named internal nodes we will
t = Tree("(A:1, (B:1, (E:1,D:1) Internal_1:0.5) Internal_2:0.5) Root;", format=1)
```

# Writing newick trees

Any ETE tree instance can be exported using newick notation using the Tree.write() method, which is available in any tree node instance. It also allows for format selection (*Reading and Writing Newick Trees*), so you can use the same function to convert between newick formats.

```
# Loads a tree with internal node names
t = Tree("(A:1, (B:1, (E:1,D:1) Internal_1:0.5) Internal_2:0.5) Root;", format=1)
# And prints its newick using the default format
print t.write() # (A:1.000000, (B:1.000000, (E:1.000000, D:1.000000) 1.000000:0.500000) 1.000
# To print the internal node names you need to change the format:
print t.write(format=1) # (A:1.000000, (B:1.000000, (E:1.000000, D:1.000000) Internal_1:0.500
# We can also write into a file
t.write(format=1, outfile="new_tree.nw")
```

# 2.1.3 Understanding ETE Trees

Any tree topology can be represented as a succession of **nodes** connected in a hierarchical way. Thus, for practical reasons, ETE makes no distinction between tree and node concepts, as any tree can be represented by its root node. This allows to use any internal node within a tree as another sub-tree instance.

Once trees are loaded, they can be manipulated as normal python objects. Given that a tree is actually a collection of nodes connected in a hierarchical way, what you usually see as a tree will be the root node instance from which the tree structure is hanging. However, every node within a ETE's tree structure can be also considered a subtree. This means, for example, that all the operational methods that we will review in the following sections are available at any possible level within a tree. Moreover, this feature will allow you to separate large trees into smaller partitions, or concatenate several trees into a single structure. For this reason, you will find that the TreeNode and Tree classes are synonymous.

# 2.1.4 Basic tree attributes

Each tree node has two basic attributes used to establish its position in the tree: TreeNode.up and TreeNode.children. The first is a pointer to parent's node, while the later is a list of children nodes. Although it is possible to modify the structure of a tree by changing these attributes, it is strongly recommend not to do it. Several methods are provided to manipulate each node's connections in a safe way (see *Comparing Trees*).

In addition, three other basic attributes are always present in any tree node instance:

Method	Description	Default
		value
TreeNode.dist	stores the distance from the node to its parent (branch length).	1.0
	Default value = 1.0	
TreeNode.suppoinforms about the reliability of the partition defined by the node 1.0		1.0
	(i.e. bootstrap support)	
TreeNode.name	Custom node's name.	NoName

In addition, several methods are provided to perform basic operations on tree node instances:

Method	Description
TreeNode.is_leaf()	returns True if <i>node</i> has no children
TreeNode.is_root()	returns True if <i>node</i> has no parent
TreeNode.get_tree_root()	returns the top-most node within the same tree structure as <i>node</i>
len(TreeNode)	returns the number of leaves under <i>node</i>
print node	prints a text-based representation of the tree topology under <i>node</i>
if node in tree	returns true if <i>node</i> is a leaf under <i>tree</i>
for leaf in node	iterates over all leaves under <i>node</i>
TreeNode.show()	Explore node graphically using a GUI.

This is an example on how to access such attributes:

```
from ete2 import Tree
t = Tree()
# We create a random tree topology
t.populate(15)
print t
print t.children
print t.get_children()
print t.up
print t.name
print t.dist
print t.is_leaf()
print t.get tree root()
print t.children[0].get tree root()
print t.children[0].children[0].get_tree_root()
# You can also iterate over tree leaves using a simple syntax
for leaf in t:
  print leaf.name
```

#### Root node on unrooted trees?

When a tree is loaded from external sources, a pointer to the top-most node is returned. This is called the tree root, and it will exist even if the tree is conceptually considered as unrooted. This is, the root

node can be considered as the master node, since it represents the whole tree structure. Unrooted trees can be identified as trees in which master root node has more than two children.

# 2.1.5 Browsing trees (traversing)

One of the most basic operations for tree analysis is *tree browsing*. This is, essentially, visiting nodes within a tree. ETE provides a number of methods to search for specific nodes or to navigate over the hierarchical structure of a tree.

# **Getting Leaves, Descendants and Node's Relatives**

TreeNode instances contain several functions to access their descendants. Available methods are self explanatory:

<pre>TreeNode.get_descendants([strategy, is_leaf_fn])</pre>	Returns a list of all (leaves and internal) descendant nodes
TreeNode.get_leaves([is_leaf_fn])	Returns the list of terminal nodes (leaves) under this node
TreeNode.get_leaf_names([is_leaf_fn])	Returns the list of terminal node names under the current
TreeNode.get_children()	Returns an independent list of node's children.
TreeNode.get_sisters()	Returns an indepent list of sister nodes.

# Traversing (browsing) trees

Often, when processing trees, all nodes need to be visited. This is called tree traversing. There are different ways to traverse a tree structure depending on the order in which children nodes are visited. ETE implements the three most common strategies: **preorder**, **levelorder** and **postorder**. The following scheme shows the differences in the strategy for visiting nodes (note that in both cases the whole tree is browsed):

• preorder: 1) Visit the root, 2) Traverse the left subtree, 3) Traverse the right subtree.

- postorder: 1) Traverse the left subtree, 2) Traverse the right subtree, 3) Visit the root
- levelorder (default): every node on a level before is visited going to a lower level

#### Note:

- Preorder traversal sequence: F, B, A, D, C, E, G, I, H (root, left, right)
- Inorder traversal sequence: A, B, C, D, E, F, G, H, I (left, root, right); note how this produces a sorted sequence
- Postorder traversal sequence: A, C, E, D, B, H, I, G, F (left, right, root)
- Level-order traversal sequence: F, B, G, A, D, I, C, E, H

Every node in a tree includes a TreeNode.traverse() method, which can be used to visit, one by one, every node node under the current partition. In addition, the TreeNode.iter\_descendants() method can be set to use either a post- or a preorder strategy. The only different between TreeNode.traverse() and TreeNode.iter\_descendants() is that the first will include the root node in the iteration.

TreeNode.traverse([strategy, is_leaf_fn])	Returns an iterator to traverse the tree structure under the
<pre>TreeNode.iter_descendants([strategy, is_leaf_fn])</pre>	Returns an iterator over all descendant nodes.
<pre>TreeNode.iter_leaves([is_leaf_fn])</pre>	Returns an iterator over the leaves under this node.

strategy can take one of the following values: "postorder", "preorder" or "levelorder"

```
# we load a tree
t = Tree('((((H,K)D,(F,I)G)B,E)A,((L,(N,Q)O)J,(P,S)M)C);', format=1)
for node in t.traverse("postorder"):
    # Do some analysis on node
    print node.name

# If we want to iterate over a tree excluding the root node, we can
# use the iter_descendant method
for node in t.iter_descendants("postorder"):
    # Do some analysis on node
    print node.name
```

Additionally, you can implement your own traversing function using the structural attributes of nodes. In the following example, only nodes between a given leaf and the tree root are visited.

```
from ete2 import Tree
tree = Tree( "(A:1, (B:1, (C:1,D:1):0.5):0.5);" )

# Browse the tree from a specific leaf to the root
node = t.search_nodes(name="C")[0]
while node:
    print node
    node = node.up
```

# Advanced traversing (stopping criteria)

# Collapsing nodes while traversing (custom is\_leaf definition)

From version 2.2, ETE supports the use of the is\_leaf\_fn argument in most of its traversing functions. The value of is\_leaf\_fn is expected to be a pointer to any python function that accepts a node instance as its first argument and returns a boolean value (True if node should be considered a leaf node).

By doing so, all traversing methods will use such a custom function to decide if a node is a leaf. This becomes specially useful when dynamic collapsing of nodes is needed, thus avoiding to prune the same tree in many different ways.

For instance, given a large tree structure, the following code will export the newick of the pruned version of the topology, where nodes grouping the same tip labels are collapsed.

```
from ete2 import Tree
def collapsed_leaf(node):
    if len(node2labels[node]) == 1:
       return True
    else:
       return False
t = Tree("((((a,a,a)a,a)aa, (b,b)b)ab, (c, (d,d)d)cd);", format=1)
print t
# We create a cache with every node content
node2labels = t.get_cached_content(store_attr="name")
print t.write(is leaf fn=collapsed leaf)
             /
#
           /-I--a
        /-/ \-a
#
#
     /-/ \-a
          \-b
#
        /-c
#
#
    \backslash - /
     /
           /-d
        \backslash - /
           \backslash -d
# We can even load the collapsed version as a new tree
t2 = Tree( t.write(is_leaf_fn=collapsed_leaf) )
print t2
        /-aa
    /-1
  /
       \ -b
```

Another interesting use of this approach is to find the first matching nodes in a given tree that match a

custom set of criteria, without browsing the whole tree structure.

Let's say we want get all deepest nodes in a tree whose branch length is larger than one:

# **Iterating instead of Getting**

As commented previously, methods starting with **get**\_ are all prepared to return results as a closed list of items. This means, for instance, that if you want to process all tree leaves and you ask for them using the <code>TreeNode.get\_leaves()</code> method, the whole tree structure will be browsed before returning the final list of terminal nodes. This is not a problem in most of the cases, but in large trees, you can speed up the browsing process by using iterators.

```
Most get_ methods have their homologous iterator functions. Thus, TreeNode.get_leaves() could be substituted by TreeNode.iter_leaves(). The same occurs with TreeNode.iter_descendants() and TreeNode.iter_search_nodes().
```

When iterators are used (note that is only applicable for looping), only one step is processed at a time. For instance, TreeNode.iter\_search\_nodes() will return one match in each iteration. In practice, this makes no differences in the final result, but it may increase the performance of loop functions (i.e. in case of finding a match which interrupts the loop).

# Finding nodes by their attributes

Both terminal and internal nodes can be located by searching along the tree structure. Several methods are available:

method	Description
t.search_nodes(attr=value)	Returns a list of nodes in which attr is equal to value, i.e. name=A
t.iter_search_nodes(attr=value)	Iterates over all matching nodes matching attr=value. Faster when
	you only need to get the first occurrence
t.get_leaves_by_name(name)	Returns a list of leaf nodes matching a given name. Only leaves are
	browsed.
t.get_common_ancestor([node1	, Return the first internal node grouping node1, node2 and node3
node2, node3])	
t&"A"	Shortcut for t.search_nodes(name="A")[0]

# Search\_all nodes matching a given criteria

A custom list of nodes matching a given name can be easily obtain through the TreeNode.search\_node() function.

# Search nodes matching a given criteria (iteration)

A limitation of the TreeNode.search\_nodes() method is that you cannot use complex conditional statements to find specific nodes. When search criteria is too complex, you may need to create your own search function.

```
from ete2 import Tree

def search_by_size(node, size):
    "Finds nodes with a given number of leaves"
    matches = []
    for n in node.traverse():
```

```
if len(n) == size:
    matches.append(n)
  return matches

t = Tree()
t.populate(40)
# returns nodes containing 6 leaves
search_by_size(t, size=6)
```

#### Find the first common ancestor

Searching for the first common ancestor of a given set of nodes it is a handy way of finding internal nodes.

```
from ete2 import Tree
t = Tree( "((H:0.3,I:0.1):0.5, A:1, (B:0.4,(C:0.5,(J:1.3, (F:1.2, D:0.1):0.5):0.5):0.5):
print t
ancestor = t.get_common_ancestor("C", "J", "B")
```

#### **Custom searching functions**

A limitation of the previous methods is that you cannot use complex conditional statements to find specific nodes. However you can user traversing methods to meet your custom filters. A possible general strategy would look like this:

```
from ete2 import Tree
t = Tree("((H:0.3,I:0.1):0.5, A:1, (B:0.4,(C:1,D:1):0.5):0.5);")
# Create a small function to filter your nodes
def conditional_function(node):
    if node.dist > 0.3:
        return True
    else:
        return False
# Use previous function to find matches. Note that we use the traverse
# method in the filter function. This will iterate over all nodes to
# assess if they meet our custom conditions and will return a list of
# matches.
matches = filter(conditional_function, t.traverse())
print len(matches), "nodes have ditance >0.3"
# depending on the complexity of your conditions you can do the same
# in just one line with the help of lambda functions:
matches = filter(lambda n: n.dist>0.3 and n.is_leaf(), t.traverse() )
print len(matches), "nodes have ditance >0.3 and are leaves"
```

#### **Shortcuts**

Finally, ETE implements a built-in method to find the first node matching a given name, which is one of the most common tasks needed for tree analysis. This can be done through the operator & (AND). Thus, TreeNode&"A" will always return the first node whose name is "A" and that is under the tree "MyTree". The syntaxis may seem confusing, but it can be very useful in some situations.

```
from ete2 import Tree
t = Tree("((H:0.3,I:0.1):0.5, A:1, (B:0.4,(C:1,(J:1, D:1):0.5):0.5):0.5):0.5):0.5);")
# Get the node D in a very simple way
D = t\&"D"
# Get the path from B to the root
node = D
path = []
while node.up:
 path.append(node)
 node = node.up
print t
# I substract D node from the total number of visited nodes
print "There are", len(path)-1, "nodes between D and the root"
# Using parentheses you can use by-operand search syntax as a node
# instance itself
Dsparent= (t&"C").up
Bsparent= (t&"B").up
Jsparent= (t&"J").up
# I check if nodes belong to certain partitions
print "It is", Dsparent in Bsparent, "that C's parent is under B's ancestor"
print "It is", Dsparent in Jsparent, "that C's parent is under J's ancestor"
```

# 2.1.6 Checking the monophyly of attributes within a tree

Although monophyly is actually a phylogenetic concept used to refer to a set of species that group exclusively together within a tree partition, the idea can be easily exported to any type of trees.

Therefore, we could consider that a set of values for a given node attribute present in our tree is monophyletic, if such values group exclusively together as a single tree partition. If not, the corresponding relationship connecting such values (para or poly-phyletic) could be also be inferred.

The TreeNode.check\_monophyly() method will do so when a given tree is queried for any custom attribute.

```
# We can check how, indeed, all vowels are not monophyletic in the
# previous tree, but polyphyletic (a foreign label breaks its monophyly)
print t.check_monophyly(values=["a", "e", "i", "o", "u"], target_attr="name")
# however, the following set of vowels are monophyletic
print t.check_monophyly(values=["a", "e", "i", "o"], target_attr="name")
# A special case of polyphyly, called paraphyly, is also used to
# define certain type of grouping. See this wikipedia article for
# disambiguation: http://en.wikipedia.org/wiki/Paraphyly
print t.check_monophyly(values=["i", "o"], target_attr="name")
```

Finally, the TreeNode.get\_monophyletic() method is also provided, which allows to return a list of nodes within a tree where a given set of attribute values are monophyletic. Note that, although a set of values are not monophyletic regarding the whole tree, several independent monophyletic partitions could be found within the same topology.

For instance, in the following example, all clusters within the same tree exclusively grouping a custom set of annotations are obtained.

```
from ete2 import Tree
t = Tree("(((((((4, e), i), o), h), u), ((3, 4), (i, june)));")
# we annotate the tree using external data
colors = {"a":"red", "e":"green", "i":"yellow",
          "o": "black", "u": "purple", "4": "green",
          "3":"yellow", "1":"white", "5":"red",
          "june": "yellow"}
for leaf in t:
    leaf.add features(color=colors.get(leaf.name, "none"))
print t.get_ascii(attributes=["name", "color"], show_internal=False)
                     /-4, green
                     \−e, green
                 \-i, yellow
              \-o, black
           \-h, none
        \-u, purple
           /-3, yellow
#
           \-4, green
#
           /-i, yellow
           \-june, yellow
print "Green-yellow clusters:"
# And obtain clusters exclusively green and yellow
for node in t.get_monophyletic(values=["green", "yellow"], target_attr="color"):
   print node.get_ascii(attributes=["color", "name"], show_internal=False)
# Green-vellow clusters:
```

**Note:** When the target attribute is set to the "species" feature name, associated to any PhyloTree node, this method will accomplish with the standard phylogenetic definition of monophyly, polyphyly and paraphyly.

# 2.1.7 Caching tree content for faster lookup operations

If your program needs to access to the content of different nodes very frequently, traversing the tree to get the leaves of each node over and over will produce significant slowdowns in your algorithm. From version 2.2 ETE provides a convenient methods to cache frequent data.

The method TreeNode.get\_cached\_content() returns a dictionary in which keys are node instances and values represent the content of such nodes. By default, content is understood as a list of leave nodes, so looking up size or tip names under a given node will be instant. However, specific attributes can be cached by setting a custom store\_attr value.

```
from ete2 import Tree
t = Tree()
t.populate(50)

node2leaves = t.get_cached_content()

# lets now print the size of each node without the need of
# recursively traverse
for n in t.traverse():
    print "node %s contains %s tips" %(n.name, len(node2leaves[n]))
```

# 2.1.8 Node annotation

Every node contains three basic attributes: name (TreeNode.name), branch length (TreeNode.dist) and branch support (TreeNode.support). These three values are encoded in the newick format. However, any extra data could be linked to trees. This is called tree annotation.

The TreeNode.add\_feature() and TreeNode.add\_features() methods allow to add extra attributes (features) to any node. The first allows to add one one feature at a time, while the second can be used to add many features with the same call.

Once extra features are added, you can access their values at any time during the analysis of a tree. To do so, you only need to access to the TreeNode.feature\_name attributes.

Similarly, TreeNode.del\_feature() can be used to delete an attribute.

```
import random
from ete2 import Tree
# Creates a tree
t = Tree('(H:0.3,I:0.1):0.5, A:1, (B:0.4,(C:0.5,(J:1.3,(F:1.2,D:0.1):0.5):0.5):0.5):
# Let's locate some nodes using the get common ancestor method
ancestor=t.get_common_ancestor("J", "F", "C")
# the search_nodes method (I take only the first match )
A = t.search_nodes(name="A")[0]
# and using the shorcut to finding nodes by name
C= t&"C"
H= t&"H"
I= t&"I"
# Let's now add some custom features to our nodes. add_features can be
# used to add many features at the same time.
C.add_features(vowel=False, confidence=1.0)
A.add_features(vowel=True, confidence=0.5)
ancestor.add_features (nodetype="internal")
# Or, using the oneliner notation
(t&"H").add_features(vowel=False, confidence=0.2)
# But we can automatize this. (note that i will overwrite the previous
# values)
for leaf in t.traverse():
   if leaf.name in "AEIOU":
      leaf.add_features(vowel=True, confidence=random.random())
   else:
      leaf.add_features(vowel=False, confidence=random.random())
# Now we use these information to analyze the tree.
print "This tree has", len(t.search_nodes(vowel=True)), "vowel nodes"
print "Which are", [leaf.name for leaf in t.iter_leaves() if leaf.vowel==True]
# But features may refer to any kind of data, not only simple
# values. For example, we can calculate some values and store them
# within nodes.
# Let's detect leaf nodes under "ancestor" with distance higher thatn
# 1. Note that I'm traversing a subtree which starts from "ancestor"
matches = [leaf for leaf in ancestor.traverse() if leaf.dist>1.0]
# And save this pre-computed information into the ancestor node
ancestor.add_feature("long_branch_nodes", matches)
# Prints the precomputed nodes
print "These are nodes under ancestor with long branches", \
   [n.name for n in ancestor.long_branch_nodes]
# We can also use the add_feature() method to dynamically add new features.
label = raw_input("custom label:")
value = raw_input("custom label value:")
```

```
ancestor.add_feature(label, value)
print "Ancestor has now the [", label, "] attribute with value [", value, "]"
```

Unfortunately, newick format does not support adding extra features to a tree. Because of this drawback, several improved formats haven been (or are being) developed to read and write tree based information. Some of these new formats are based in a completely new standard (*Phylogenetic XML standards*), while others are extensions of the original newick formar (NHX http://phylosoft.org/NHX/http://phylosoft.org/NHX/).

Currently, ETE includes support for the New Hampshire eXtended format (NHX), which uses the original newick standard and adds the possibility of saving additional date related to each tree node. Here is an example of a extended newick representation in which extra information is added to an internal node:

```
(A:0.35, (B:0.72, (D:0.60, G:0.12):0.64[&&NHX:conf=0.01:name=INTERNAL]):0.56);
```

As you can notice, extra node features in the NHX format are enclosed between brackets. ETE is able to read and write features using such format, however, the encoded information is expected to be exportable as plain text.

The NHX format is automatically detected when reading a newick file, and the detected node features are added using the <code>TreeNode.add\_feature()</code> method. Consequently, you can access the information by using the normal ETE's feature notation: <code>node.feature\_name</code>. Similarly, features added to a tree can be included within the normal newick representation using the NHX notation. For this, you can call the <code>TreeNode.write()</code> method using the <code>features</code> argument, which is expected to be a list with the features names that you want to include in the newick string. Note that all nodes containing the suplied features will be exposed into the newick string. Use an empty features list (<code>features=[]</code>) to include all node's data into the newick string.

```
import random
from ete2 import Tree
# Creates a normal tree
t = Tree('(H:0.3,I:0.1):0.5, A:1,(B:0.4,(C:0.5,(J:1.3,(F:1.2,D:0.1):0.5):0.5):0.5):0.5)
print t
# Let's locate some nodes using the get common ancestor method
ancestor=t.get_common_ancestor("J", "F", "C")
# Let's label leaf nodes
for leaf in t.traverse():
    if leaf.name in "AEIOU":
      leaf.add_features(vowel=True, confidence=random.random())
    else:
      leaf.add features(vowel=False, confidence=random.random())
# Let's detect leaf nodes under "ancestor" with distance higher thatn
# 1. Note that I'm traversing a subtree which starts from "ancestor"
matches = [leaf for leaf in ancestor.traverse() if leaf.dist>1.0]
# And save this pre-computed information into the ancestor node
ancestor.add_feature("long_branch_nodes", matches)
print
print "NHX notation including vowel and confidence attributes"
print
print t.write(features=["vowel", "confidence"])
print "NHX notation including all node's data"
```

print

```
# Note that when all features are requested, only those with values
# equal to text-strings or numbers are considered. "long branch nodes"
# is not included into the newick string.
print t.write(features=[])
print
print "basic newick formats are still available"
print
print t.write(format=9, features=["vowel"])
# You don't need to do anything speciall to read NHX notation. Just
# specify the newick format and the NHX tags will be automatically
# detected.
nw = """
(((ADH2:0.1[&&NHX:S=human:E=1.1.1.1], ADH1:0.11[&&NHX:S=human:E=1.1.1.1])
:0.05[&&NHX:S=Primates:E=1.1.1.1:D=Y:B=100], ADHY:0.1[&&NHX:S=nematode:
E=1.1.1.1], ADHX:0.12[&&NHX:S=insect:E=1.1.1.1]):0.1[&&NHX:S=Metazoa:
E=1.1.1.1:D=N], (ADH4:0.09[&&NHX:S=yeast:E=1.1.1.1], ADH3:0.13[&&NHX:S=yeast:
E=1.1.1.1], ADH2:0.12[&&NHX:S=yeast:E=1.1.1.1], ADH1:0.11[&&NHX:S=yeast:E=1.1.1.1]):0.1
[&&NHX:S=Fungi]) [&&NHX:E=1.1.1.1:D=N];
# Loads the NHX example found at http://www.phylosoft.org/NHX/
t = Tree(nw)
# And access node's attributes.
for n in t.traverse():
    if hasattr(n, "S"):
       print n.name, n.S
```

# 2.1.9 Comparing Trees

#### Calculate distances between trees

The :Tree:'compare' function allows to calculate distances between two trees based on any node feature (i.e. name, species, other tags) using robinson-foulds and edge compatibility distances. It automatically handles differences in tree sizes, shared nodes and duplicated feature names.

- result["rf"] = robinson-foulds distance between the two trees. (average of robinson-foulds distances if target tree contained duplication and was split in several subtrees)
- result["max\_rf"] = Maximum robinson-foulds distance expected for this comparison
- result["norm rf"] = normalized robinson-foulds distance (from 0 to 1)
- result["effective\_tree\_size"] = the size of the compared trees, which are pruned to the common shared nodes.
- result["ref\_edges\_in\_source"] = compatibility score of the target tree with respect to the source tree (how many edges in reference are found in the source)
- result["source\_edges\_in\_ref"] = compatibility score of the source tree with respect to the reference tree (how many edges in source are found in the reference)
- result["source\_subtrees"] = number of subtrees in the source tree (1 if do not contain duplications)
- result["common\_edges"] = a set of common edges between source tree and reference
- result["source\_edges"] = the set of edges found in the source tree
- result["ref\_edges"] = the set of edges found in the reference tree

• result["treeko\_dist"] = TreeKO speciation distance for comparisons including duplication nodes.

#### **Robinson-foulds distance**

Two tree topologies can be compared using ETE and the Robinson-Foulds (RF) metric. The method TreeNode.robinson\_foulds() available for any ETE tree node allows to:

- compare two tree topologies by their name labels (default) or any other annotated feature in the tree.
- compare topologies of different size and content. When two trees contain a different set of labels, only shared leaves will be used.
- examine size and content of matching and missing partitions. Since the method return the list of partitions found in both trees, details about matching partitions can be obtained easily.

In the following example, several of above mentioned features are shown:

```
from ete2 import Tree
t1 = Tree('(((a,b),c), ((e, f), g));')
t2 = Tree('((a,c),b), ((e, f), g));')
rf, max_rf, common_leaves, parts_t1, parts_t2 = t1.robinson_foulds(t2)
print t1, t2
print "RF distance is %s over a total of %s" %(rf, max_rf)
print "Partitions in tree2 that were not found in tree1:", parts_t1 - parts_t2
print "Partitions in tree1 that were not found in tree2:", parts_t2 - parts_t1
# We can also compare trees sharing only part of their labels
t1 = Tree('((a,b),c), ((e, f), g));')
t2 = Tree('(((a,c),b), (g, H));')
rf, max_rf, common_leaves, parts_t1, parts_t2 = t1.robinson_foulds(t2)
print t1, t2
print "Same distance holds even for partially overlapping trees"
print "RF distance is %s over a total of %s" %(rf, max_rf)
print "Partitions in tree2 that were not found in tree1:", parts_t1 - parts_t2
print "Partitions in tree1 that were not found in tree2:", parts_t2 - parts_t1
```

# 2.1.10 Modifying Tree Topology

# **Creating Trees from Scratch**

If no arguments are passed to the TreeNode class constructor, an empty tree node will be returned. Such an orphan node can be used to populate a tree from scratch. For this, the TreeNode.up, and TreeNode.children attributes should never be used (unless it is strictly necessary). Instead, several methods exist to manipulate the topology of a tree:

TreeNode.populate(size[, names_library,])	Generates a random topology by populating current node.
<pre>TreeNode.add_child([child, name, dist, support])</pre>	Adds a new child to this node.
TreeNode.add_child([child, name, dist, support])	Adds a new child to this node.
TreeNode.delete([prevent_nondicotomic,])	Deletes node from the tree structure.
TreeNode.detach()	Detachs this node (and all its descendants) from its parent ar

```
from ete2 import Tree
t = Tree() # Creates an empty tree
A = t.add child(name="A") # Adds a new child to the current tree root
                           # and returns it
B = t.add_child(name="B") # Adds a second child to the current tree
                           # root and returns it
C = A.add_child(name="C") # Adds a new child to one of the branches
D = C.add_sister(name="D") # Adds a second child to same branch as
                             # before, but using a sister as the starting
                              # point
R = A.add_child(name="R") # Adds a third child to the
                           # branch. Multifurcations are supported
# Next, I add 6 random leaves to the R branch names_library is an
# optional argument. If no names are provided, they will be generated
# randomly.
R.populate(6, names_library=["r1","r2","r3","r4","r5","r6"])
# Prints the tree topology
print t
                                                     /-r4
                                                     -r2
                                           \backslash -r1
 a common use of the populate method is to quickly create example
 trees from scratch. Here we create a random tree with 100 leaves.
t = Tree()
t.populate(100)
```

# Deleting (eliminating) and Removing (detaching) nodes

As currently implemented, there is a difference between detaching and deleting a node. The former disconnects a complete partition from the tree structure, so all its descendants are also disconnected from the tree. There are two methods to perform this action: TreeNode.remove\_child() and TreeNode.detach(). In contrast, deleting a node means eliminating such node without affecting its descendants. Children from the deleted node are automatically connected to the next possible parent. This is better understood with the following example:

```
from ete2 import Tree
# Loads a tree. Note that we use format 1 to read internal node names
t = Tree('((((H,K)D,(F,I)G)B,E)A,((L,(N,Q)O)J,(P,S)M)C);', format=1)
print "original tree looks like this:"
# This is an alternative way of using "print t". Thus we have a bit
# more of control on how tree is printed. Here i print the tree
```

```
# showing internal node names
print t.get_ascii(show_internal=True)
#-NoName--|
# Get pointers to specific nodes
G = t.search_nodes(name="G")[0]
J = t.search_nodes(name="J")[0]
C = t.search_nodes(name="C")[0]
# If we remove J from the tree, the whole partition under J node will
# be detached from the tree and it will be considered an independent
# tree. We can do the same thing using two approaches: J.detach() or
# C.remove_child(J)
removed_node = J.detach() # = C.remove_child(J)
# if we know print the original tree, we will see how J partition is
# no longer there.
print "Tree after REMOVING the node J"
print t.get_ascii(show_internal=True)
# however, if we DELETE the node G, only G will be eliminated from the
# tree, and all its descendants will then hang from the next upper
# node.
G.delete()
print "Tree after DELETING the node G"
print t.get_ascii(show_internal=True)
#
```

# 2.1.11 Pruning trees

Pruning a tree means to obtain the topology that connects a certain group of items by removing the unnecessary edges. To facilitate this task, ETE implements the TreeNode.prune() method, which can be used by providing the list of terminal and/or internal nodes that must be kept in the tree.

From version 2.2, this function includes also the *preserve\_branch\_length* flag, which allows to remove nodes from a tree while keeping original distances among remaining nodes.

```
from ete2 import Tree
# Let's create simple tree
t = Tree('((((H,K),(F,I)G),E),((L,(N,Q)O),(P,S)));')
print "Original tree looks like this:"
print t
# Prune the tree in order to keep only some leaf nodes.
t.prune(["H","F","E","Q", "P"])
print "Pruned tree"
print t
```

# 2.1.12 Concatenating trees

Given that all tree nodes share the same basic properties, they can be connected freely. In fact, any node can add a whole subtree as a child, so we can actually *cut & paste* partitions. To do so, you only need to call the TreeNode.add\_child() method using another tree node as a first argument. If such a node is the root node of a different tree, you will concatenate two structures. But caution!!, this kind of operations may result into circular tree structures if add an node's ancestor as a new node's child. Some basic checks are internally performed by the ETE topology related methods, however, a fully qualified check of this issue would affect seriously the performance of the method. For this reason, users themselves should take care about not creating circular structures by mistake.

```
from ete2 import Tree
# Loads 3 independent trees
t1 = Tree('(A, (B, C));')
t2 = Tree('(D,E), (F,G));')
t3 = Tree('(H, ((I,J), (K,L)));')
print "Tree1:", t1
print "Tree2:", t2
print "Tree3:", t3
#
# Locates a terminal node in the first tree
A = t1.search_nodes(name='A')[0]
# and adds the two other trees as children.
A.add_child(t2)
A.add_child(t3)
print "Resulting concatenated tree:", t1
#
```

# 2.1.13 Copying (duplicating) trees

ETE provides several strategies to clone tree structures. The method <code>TreeNode.copy()</code> can be used to produce a new independent tree object with the exact topology and features as the original. However, as trees may involve many intricate levels of branches and nested features, 4 different methods are available to create a tree copy:

- "newick": Tree topology, node names, branch lengths and branch support values will be copied as represented in the newick string This method is based on newick format serialization works very fast even for large trees.
- "newick-extended": Tree topology and all node features will be copied based on the extended newick format representation. Only node features will be copied, thus excluding other node attributes. As this method is also based on newick serialisation, features will be converted into text strings when making the copy. Performance will depend on the tree size and the number and type of features being copied.
- "cpickle": This is the default method. The whole node structure and its content will be cloned based on the cPickle object serialization python approach. This method is slower, but recommended for full tree copying.
- "deepcopy": The whole node structure and its content is copied based on the standard "copy" Python functionality. This is the slowest method, but it allows to copy very complex objects even when attributes point to lambda functions.

```
\Internal_2, 0.5
                         \backslash -D, 0.0
# Newick copy will loose custom node annotations, complex features,
# but not names and branch values
print t.copy("newick").get_ascii(attributes=["name", "dist", "label", "complex"])
                           /-A, 0.0
#
             /Internal_1, 0.7
                          \backslash -B, 0.0
 -NoName, 0.0
                          /-C, 0.0
#
             \Internal_2, 0.5
                           # Extended newick copy will transfer custom annotations as text
# strings, so complex features are lost.
print t.copy("newick-extended").get ascii(attributes=["name", "dist", "label", "complex")
                             /-A, 0.0, custom Value, __0_ 1__ _2_ 3__ _1_ 11__ _1_ 0__
            /Internal_1, 0.7
#
                           -NoName, 0.0
                          /-C, 0.0
             \Internal_2, 0.5
                          # The default pickle method will produce a exact clone of the
# original tree, where features are duplicated keeping their
# python data type.
print t.copy().get_ascii(attributes=["name", "dist", "label", "complex"])
print "first element in complex feature:", (t & "A").complex[0]
#
                         /-A, 0.0, custom Value, [[0, 1], [2, 3], [1, 11], [1, 0]]
#
          /Internal_1, 0.7
                         -root, 1.3
                         /-C, 0.0
           \Internal_2, 0.5
                         \-D, 0.0
# first element in complex feature: [0, 1]
```

# 2.1.14 Solving multifurcations

When a tree contains a polytomy (a node with more than 2 children), the method resolve\_polytomy() can be used to convert the node into a randomly bifurcated structure in which branch lengths are set to 0. This is really not a solution for the polytomy but it allows to export the tree as a strictly bifurcated newick structure, which is a requirement for some external software.

The method can be used on a very specific node while keeping the rest of the tree intact by disabling the recursive flag.

```
from ete2 import Tree
t = Tree("(( (a, b, c), (d, e, f, g)), (f, i, h));")
print t
#
polynode = t.get_common_ancestor("a", "b")
polynode.resolve_polytomy(recursive=False)
print t
# /
# ---/
#
  | /-f
| |
#
          \backslash -h
t.resolve_polytomy(recursive=True)
print t
#
```

### 2.1.15 Tree Rooting

Tree rooting is understood as the technique by with a given tree is conceptually polarized from more basal to more terminal nodes. In phylogenetics, for instance, this a crucial step prior to the interpretation of trees, since it will determine the evolutionary relationships among the species involved. The concept of rooted trees is different than just having a root node, which is always necessary to handle a tree data structure. Usually, the way in which a tree is differentiated between rooted and unrooted, is by counting the number of branches of the current root node. Thus, if the root node has more than two child branches, the tree is considered unrooted. By contrast, when only two main branches exist under the root node, the tree is considered rooted.

Having an unrooted tree means that any internal branch within the tree could be regarded as the root node, and there is no conceptual reason to place the root node where it is placed at the moment. Therefore, in an unrooted tree, there is no information about which internal nodes are more basal than others. By setting the root node between a given edge/branch of the tree structure the tree is polarized, meaning that the two branches under the root node are the most basal nodes. In practice, this is usually done by setting an **outgroup node**, which would represent one of these main root branches. The second one will be, obviously, the brother node. When you set an outgroup on unrooted trees, the multifurcations at the current root node are solved.

In order to root an unrooted tree or re-root a tree structure, ETE implements the TreeNode.set\_outgroup() method, which is present in any tree node instance. Similarly, the TreeNode.unroot() method can be used to perform the opposite action.

```
# Let's define that the ancestor of E and D as the tree outgroup. Of
# course, the definition of an outgroup will depend on user criteria.
ancestor = t.get_common_ancestor("E","D")
t.set_outgroup(ancestor)
print "Tree rooteda at E and D's ancestor is more basal that the others."
print t
#
#
# Note that setting a different outgroup, a different interpretation
# of the tree is possible
t.set_outgroup( t&"A" )
print "Tree rooted at a terminal node"
print t
#
           \backslash -A
```

Note that although **rooting** is usually regarded as a whole-tree operation, ETE allows to root subparts of the tree without affecting to its parent tree structure.

```
# Each main branch of the tree is independently rooted.
node1 = t.get_common_ancestor("A","H")
node2 = t.get_common_ancestor("B","D")
node1.set_outgroup("H")
node2.set_outgroup("E")
print "Tree after rooting each node independently:"
                      \backslash -E
```

# 2.1.16 Working with branch distances

The branch length between one node an its parent is encoded as the TreeNode.dist attribute. Together with tree topology, branch lengths define the relationships among nodes.

### Getting distances between nodes

The TreeNode.get\_distance() method can be used to calculate the distance between two connected nodes. There are two ways of using this method: a) by querying the distance between two

descendant nodes (two nodes are passed as arguments) b) by querying the distance between the current node and any other relative node (parental or descendant).

```
from ete2 import Tree
```

```
# Loads a tree with branch lenght information. Note that if no
# distance info is provided in the newick, it will be initialized with
# the default dist value = 1.0
nw = """(((A:0.1, B:0.01):0.001, C:0.0001):1.0,
(((((D:0.00001:0,I:0):0,F:0):0,G:0):0,H:0):0,
E:0.000001):0.0000001):2.0;"""
t = Tree(nw)
print t
#
#
# Locate some nodes
A = t&"A"
C = t\&"C"
# Calculate distance from current node
print "The distance between A and C is", A.get_distance("C")
# Calculate distance between two descendants of current node
print "The distance between A and C is", t.get_distance("A","C")
# Calculate the toplogical distance (number of nodes in between)
print "The number of nodes between A and D is ", \
    t.get_distance("A","D", topology_only=True)
```

Additionally to this, ETE incorporates two more methods to calculate the most distant node from a given point in a tree. You can use the <code>TreeNode.get\_farthest\_node()</code> method to retrieve the most distant point from a node within the whole tree structure. Alternatively, <code>TreeNode.get\_farthest\_leaf()</code> will return the most distant descendant (always a leaf). If more than one node matches the farthest distance, the first occurrence is returned.

Distance between nodes can also be computed as the number of nodes between them (considering all branch lengths equal to 1.0). To do so, the **topology\_only** argument must be set to **True** for all the above mentioned methods.

```
# Calculate the farthest node from E within the whole structure
farthest, dist = (t&"E").get_farthest_node()
print "The farthest node from E is", farthest.name, "with dist=", dist
# Calculate the farthest node from E within the whole structure,
# regarding the number of nodes in between as distance value
# Note that the result is differnt.
```

```
farthest, dist = (t&"E").get_farthest_node(topology_only=True)
print "The farthest (topologically) node from E is", \
         farthest.name, "with", dist, "nodes in between"
# Calculate farthest node from an internal node
farthest, dist = t.get_farthest_node()
print "The farthest node from root is is", farthest.name, "with dist=", dist
#
# The program results in the following information:
#
# The distance between A and C is 0.1011
# The distance between A and C is 0.1011
# The number of nodes between A and D is 8.0
# The farthest node from E is A with dist= 1.1010011
# The farthest (topologically) node from E is I with 5.0 nodes in between
# The farthest node from root is is A with dist= 1.101
```

#### getting midpoint outgroup

In order to obtain a balanced rooting of the tree, you can set as the tree outgroup that partition which splits the tree in two equally distant clusters (using branch lengths). This is called the midpoint outgroup.

The TreeNode.get\_midpoint\_outgroup () method will return the outgroup partition that splits current node into two balanced branches in terms of node distances.

```
from ete2 import Tree
# generates a random tree
t = Tree();
t.populate(15);
print t
                         -gogjl
                        \-vxbgp
                         -xyewk
                                              /-opben
                                                                     /-xorvn
                                                                                /-wdima
                                                                                \neg qxovz
                                                           -isngq
                                                           -neasc
                                                                                \−exmsn
                                                                                /-udspq
```

```
\-buxpw
                                            \-rkzwd
# Calculate the midpoint node
R = t.get_midpoint_outgroup()
# and set it as tree outgroup
t.set_outgroup(R)
print t
                                  -opben
                                                       /-xoryn
                                                                  /-wdima
                                                                  \neg qxovz
                                             \-isngq
                                             \-vxbgp
                                   -neqsc
                                                                  \-djeoh
                                                       \-exmsn
                                                       /-udspq
                                                       \-buxpw
                       \-rkzwd
```

# 2.2 The Programmable Tree Drawing Engine

#### **Contents**

- The Programmable Tree Drawing Engine
  - Overview
  - Interactive visualization of trees
  - Rendering trees as images
  - Customizing the aspect of trees
    - \* Tree style
      - · Show leaf node names, branch length and branch support
      - · Change branch length scale (zoom in X)
      - · Change branch separation between nodes (zoom in Y)
      - · Rotate a tree
      - · circular tree in 180 degrees
      - · Add legend and title
    - \* Node style
    - \* Node faces
      - · Faces position
      - · Face properties
    - \* layout functions
  - Combining styles, faces and layouts
    - \* Fixed node styles
    - \* Node backgrounds
    - \* Img Faces
    - \* Bubble tree maps
    - \* Trees within trees
    - \* Phylogenetic trees and sequence domains
    - \* Creating your custom interactive Item faces

#### 2.2.1 Overview

ETE's treeview extension provides a highly programmable drawing system to render any hierarchical tree structure as PDF, SVG or PNG images. Although several predefined visualization layouts are included with the default installation, custom styles can be easily created from scratch.

Image customization is performed through four elements: a) TreeStyle, setting general options about the image (shape, rotation, etc.), b) NodeStyle, which defines the specific aspect of each node (size, color, background, line type, etc.), c) node faces.Face which are small pieces of extra graphical information that can be added to nodes (text labels, images, graphs, etc.) d) a layout function, a normal python function that controls how node styles and faces are dynamically added to nodes.

Images can be rendered as **PNG**, **PDF** or **SVG** files using the TreeNode.render() method or interactively visualized using a built-in Graphical User Interface (GUI) invoked by the TreeNode.show() method.

### 2.2.2 Interactive visualization of trees

ETE's tree drawing engine is fully integrated with a built-in graphical user interface (GUI). Thus, ETE allows to visualize trees using an interactive interface that allows to explore and manipulate node's properties and tree topology. To start the visualization of a node (tree or subtree), you can simply call the TreeNode.show() method.

One of the advantages of this on-line GUI visualization is that you can use it to interrupt a given program/analysis, explore the tree, manipulate them, and continuing with the execution thread. Note that **changes made using the GUI will be kept after quiting the GUI**. This feature is specially useful for using during python sessions, in which analyses are performed interactively.

The GUI allows many operations to be performed graphically, however it does not implement all the possibilities of the programming toolkit.

```
from ete2 import Tree
t = Tree( "((a,b),c);" )
t.show()
```

# 2.2.3 Rendering trees as images

Tree images can be directly written as image files. SVG, PDF and PNG formats are supported. Note that, while PNG images are raster images, PDF and SVG pictures are rendered as vector graphics, thus allowing its later modification and scaling.

To generate an image, the <code>TreeNode.render()</code> method should be used instead of <code>TreeNode.show()</code>. The only required argument is the file name, whose extension will determine the image format (.PDF, .SVG or .PNG). Several parameters regarding the image size and resolution can be adjusted:

Argument	Description
units	"px": pixels, "mm": millimeters, "in": inches
h	height of the image in units.
W	weight of the image in units.
dpi	dots per inches.

**Note:** If h and w values are both provided, image size will be adjusted even if it requires to break the original aspect ratio of the image. If only one value (h or w) is provided, the other will be estimated to maintain aspect ratio. If no sizing values are provided, image will be adjusted to A4 dimensions.

```
from ete2 import Tree
t = Tree( "((a,b),c);" )
t.render("mytree.png", w=183, units="mm")
```

### 2.2.4 Customizing the aspect of trees

Image customization is performed through four main elements:

#### Tree style

The TreeStyle class can be used to create a custom set of options that control the general aspect of the tree image. Tree styles can be passed to the TreeNode.show() and TreeNode.render() methods. For instance, TreeStyle allows to modify the scale used to render tree branches or choose between circular or rectangular tree drawing modes.

```
from ete2 import Tree, TreeStyle
t = Tree( "((a,b),c);" )
circular_style = TreeStyle()
```

```
circular_style.mode = "c" # draw tree in circular mode
circular_style.scale = 20
t.render("mytree.png", w=183, units="mm", tree_style=circular_style)
```

**Warning:** A number of parameters can be controlled through custom tree style objetcs, check TreeStyle documentation for a complete list of accepted values.

Some common uses include:

# Show leaf node names, branch length and branch support

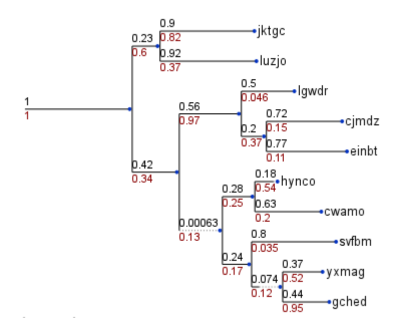


Figure 2.1: Automatically adds node names and branch information to the tree image:

```
from ete2 import Tree, TreeStyle
t = Tree()
t.populate(10, random_dist=True)
ts = TreeStyle()
ts.show_leaf_name = True
ts.show_branch_length = True
ts.show_branch_support = True
t.show(tree_style=ts)
```

Change branch length scale (zoom in X)

Change branch separation between nodes (zoom in Y)

Rotate a tree

46

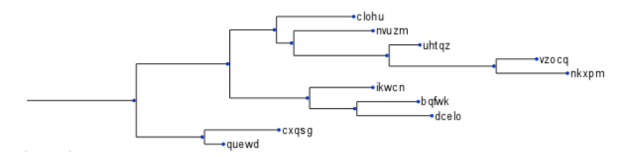


Figure 2.2: Increases the length of the tree by changing the scale:

```
from ete2 import Tree, TreeStyle
t = Tree()
t.populate(10, random_dist=True)
ts = TreeStyle()
ts.show_leaf_name = True
ts.scale = 120 # 120 pixels per branch length unit
t.show(tree_style=ts)
```

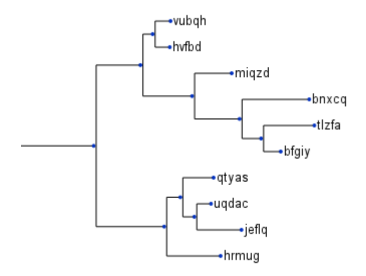


Figure 2.3: Increases the separation between leaf branches:

```
from ete2 import Tree, TreeStyle
t = Tree()
t.populate(10, random_dist=True)
ts = TreeStyle()
ts.show_leaf_name = True
ts.branch_vertical_margin = 10 # 10 pixels between adjacent branches
t.show(tree_style=ts)
```

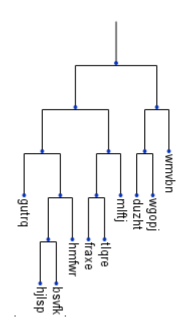


Figure 2.4: Draws a rectangular tree from top to bottom:

```
from ete2 import Tree, TreeStyle
t = Tree()
t.populate(10)
ts = TreeStyle()
ts.show_leaf_name = True
ts.rotation = 90
t.show(tree_style=ts)
```

#### circular tree in 180 degrees

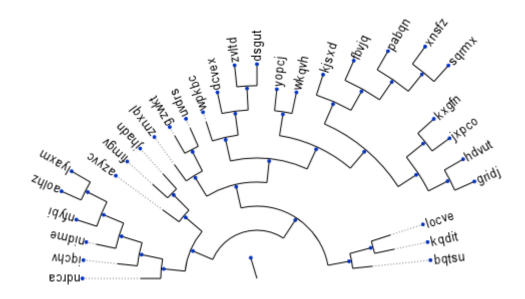


Figure 2.5: Draws a circular tree using a semi-circumference:

```
from ete2 import Tree, TreeStyle
t = Tree()
t.populate(30)
ts = TreeStyle()
ts.show_leaf_name = True
ts.mode = "c"
ts.arc_start = -180 # 0 degrees = 3 o'clock
ts.arc_span = 180
t.show(tree_style=ts)
```

#### Add legend and title

```
from ete2 import Tree, TreeStyle, TextFace
t = Tree( "((a,b),c);" )
ts = TreeStyle()
ts.show_leaf_name = True
ts.title.add_face(TextFace("Hello ETE", fsize=20), column=0)
t.show(tree_style=ts)
```

# Node style

Through the NodeStyle class the aspect of each single node can be controlled, including its size, color, background and branch type.

A node style can be defined statically and attached to several nodes:

If you want to draw nodes with different styles, an independent NodeStyle instance must be created for each node. Note that node styles can be modified at any moment by accessing the TreeNode.img\_style attribute.



Figure 2.6: Simple tree in which the same style is applied to all nodes:

```
from ete2 import Tree, NodeStyle, TreeStyle
t = Tree("((a,b),c);")
# Basic tree style
ts = TreeStyle()
ts.show_leaf_name = True
# Draws nodes as small red spheres of diameter equal to 10 pixels
nstyle = NodeStyle()
nstyle["shape"] = "sphere"
nstyle["size"] = 10
nstyle["fgcolor"] = "darkred"
# Gray dashed branch lines
nstyle["hz_line_type"] = 1
nstyle["hz_line_color"] = "#cccccc"
# Applies the same static style to all nodes in the tree. Note that,
# if "nstyle" is modified, changes will affect to all nodes
for n in t.traverse():
  n.set_style(nstyle)
t.show(tree_style=ts)
```

Figure 2.7: Simple tree in which the different styles are applied to each node:

```
from ete2 import Tree, NodeStyle, TreeStyle
t = Tree("((a,b),c);")
# Basic tree style
ts = TreeStyle()
ts.show_leaf_name = True
# Creates an independent node style for each node, which is
# initialized with a red foreground color.
for n in t.traverse():
  nstyle = NodeStyle()
   nstyle["fgcolor"] = "red"
   nstyle["size"] = 15
   n.set_style(nstyle)
# Let's now modify the aspect of the root node
t.img_style["size"] = 30
t.img_style["fgcolor"] = "blue"
t.show(tree_style=ts)
```

Static node styles, set through the set\_style() method, will be attached to the nodes and exported as part of their information. For instance, TreeNode.copy() will replicate all node styles in the replicate tree. Note that node styles can be also modified on the fly through a layout function (see layout functions)

#### **Node faces**

Node faces are small pieces of graphical information that can be linked to nodes. For instance, text labels or external images could be linked to nodes and they will be plotted within the tree image.

Several types of node faces are provided by the main ete2 module, ranging from simple text (TextFace) and geometric shapes (CircleFace), to molecular sequence representations (SequenceFace), heatmaps and profile plots (ProfileFace). A complete list of available faces can be found at the ete2.treeview reference page..

#### **Faces position**

Faces can be added to different areas around the node, namely **branch-right**, **branch-top**, **branch-bottom** or **aligned**. Each area represents a table in which faces can be added through the TreeNode.add\_face() method. For instance, if two text labels want to be drawn bellow the branch line of a given node, a pair of TextFace faces can be created and added to the columns 0 and 1 of the **branch-bottom** area:

```
from ete2 import Tree, TreeStyle, TextFace
t = Tree( "((a,b),c);")

# Basic tree style
ts = TreeStyle()
ts.show_leaf_name = True

# Add two text faces to different columns
t.add_face(TextFace("hola "), column=0, position = "branch-right")
t.add_face(TextFace("mundo!"), column=1, position = "branch-right")
t.show(tree_style=ts)
```

If you add more than one face to the same area and column, they will be piled up. See the following image as an example of face positions:

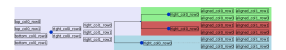


Figure 2.8: Source code used to generate the above image.

**Note:** Once a face object is created, it can be linked to one or more nodes. For instance, the same text label can be recycled and added to several nodes.

#### **Face properties**

Apart from the specific config values of each face type, all face instances contain same basic attributes that permit to modify general aspects such as margins, background colors, border, etc. A complete list of face attributes can be found in the general Face class documentation. Here is a very simple example:

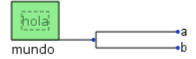


Figure 2.9: Basic use of face general attributes

```
from ete2 import Tree, TreeStyle, TextFace
t = Tree("(a,b);")
# Basic tree style
ts = TreeStyle()
ts.show_leaf_name = True
# Creates two faces
hola = TextFace("hola")
mundo = TextFace("mundo")
# Set some attributes
hola.margin\_top = 10
hola.margin_right = 10
hola.margin_left = 10
hola.margin_bottom = 10
hola.opacity = 0.5 # from 0 to 1
hola.inner_border.width = 1 # 1 pixel border
hola.inner_border.type = 1 # dashed line
hola.border.width = 1
hola.background.color = "LightGreen"
t.add_face(hola, column=0, position = "branch-top")
t.add_face(mundo, column=1, position = "branch-bottom")
t.show(tree_style=ts)
```

# layout functions

Layout functions act as pre-drawing hooking functions. This means, when a node is about to be drawn, it is first sent to a layout function. Node properties, style and faces can be then modified on the fly and return it to the drawer engine. Thus, layout functions can be understood as a collection of rules controlling how different nodes should be drawn.

```
from ete2 import Tree
t = Tree( "((((a,b),c), d), e);")
def abc_layout(node):
```

```
vowels = set(["a", "e", "i", "o", "u"])
if node.name in vowels:

# Note that node style are already initialized with the
# default values

node.img_style["size"] = 15
node.img_style["color"] = "red"

# Basic tree style
ts = TreeStyle()
ts.show_leaf_name = True

# Add two text faces to different columns
t.add_face(TextFace("hola "), column=0, position = "branch-right")
t.add_face(TextFace("mundo!"), column=1, position = "branch-right")
t.show(tree_style=ts)
```

### 2.2.5 Combining styles, faces and layouts

Examples are probably the best way to show how ETE works:

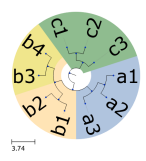
### Fixed node styles



```
from ete2 import Tree, faces, AttrFace, TreeStyle, NodeStyle
def layout(node):
    # If node is a leaf, add the nodes name and a its scientific name
    if node.is_leaf():
        faces.add_face_to_node(AttrFace("name"), node, column=0)
def get_example_tree():
    t = Tree()
    t.populate(8)
    # Node style handling is no longer limited to layout functions. You
    # can now create fixed node styles and use them many times, save them
    # or even add them to nodes before drawing (this allows to save and
    # reproduce an tree image design)
    # Set bold red branch to the root node
    style = NodeStyle()
    style["fgcolor"] = "#0f0f0f"
    style["size"] = 0
    style["vt_line_color"] = "#ff0000"
    style["hz_line_color"] = "#ff0000"
    style["vt_line_width"] = 8
    style["hz_line_width"] = 8
    style["vt_line_type"] = 0 # 0 solid, 1 dashed, 2 dotted
```

```
style["hz_line_type"] = 0
    t.set_style(style)
    #Set dotted red lines to the first two branches
   style1 = NodeStyle()
    style1["fgcolor"] = "#0f0f0f"
    style1["size"] = 0
   style1["vt_line_color"] = "#ff0000"
   style1["hz_line_color"] = "#ff0000"
   style1["vt_line_width"] = 2
   style1["hz_line_width"] = 2
   style1["vt_line_type"] = 2 # 0 solid, 1 dashed, 2 dotted
   style1["hz_line_type"] = 2
    t.children[0].img_style = style1
    t.children[1].img_style = style1
    # Set dashed blue lines in all leaves
   style2 = NodeStyle()
   style2["fgcolor"] = "#000000"
   style2["shape"] = "circle"
    style2["vt_line_color"] = "#0000aa"
    style2["hz_line_color"] = "#0000aa"
   style2["vt_line_width"] = 2
   style2["hz_line_width"] = 2
   style2["vt_line_type"] = 1 # 0 solid, 1 dashed, 2 dotted
    style2["hz_line_type"] = 1
    for l in t.iter_leaves():
        l.img_style = style2
   ts = TreeStyle()
   ts.layout_fn = layout
   ts.show_leaf_name = False
   return t, ts
if __name__ == "__main__":
   t, ts = get_example_tree()
   t.show(tree_style=ts)
    #t.render("node_style.png", w=400, tree_style=ts)
```

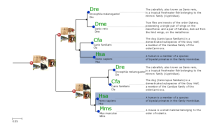
### Node backgrounds



```
from ete2 import Tree, faces, AttrFace, TreeStyle, NodeStyle
def layout(node):
```

```
if node.is_leaf():
       N = AttrFace("name", fsize=30)
        faces.add_face_to_node(N, node, 0, position="aligned")
def get_example_tree():
    # Set dashed blue lines in all leaves
   nst1 = NodeStyle()
   nst1["bgcolor"] = "LightSteelBlue"
   nst2 = NodeStyle()
   nst2["bgcolor"] = "Moccasin"
   nst3 = NodeStyle()
   nst3["bgcolor"] = "DarkSeaGreen"
   nst4 = NodeStyle()
   nst4["bgcolor"] = "Khaki"
   t = Tree("((((a1,a2),a3), ((b1,b2), (b3,b4))), ((c1,c2),c3));")
   for n in t.traverse():
       n.dist = 0
   n1 = t.get common ancestor("a1", "a2", "a3")
   n1.set_style(nst1)
   n2 = t.get_common_ancestor("b1", "b2", "b3", "b4")
   n2.set_style(nst2)
   n3 = t.get_common_ancestor("c1", "c2", "c3")
   n3.set_style(nst3)
   n4 = t.get_common_ancestor("b3", "b4")
   n4.set_style(nst4)
   ts = TreeStyle()
   ts.layout_fn = layout
   ts.show_leaf_name = False
   ts.mode = "c"
   ts.root_opening_factor = 1
   return t, ts
if __name__ == "__main__":
   t, ts = get_example_tree()
    #t.render("node_background.png", w=400, tree_style=ts)
   t.show(tree_style=ts)
```

### **Img Faces**



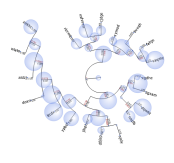
Note that images are attached to terminal and internal nodes.

```
# Import Tree instance and faces module
from ete2 import Tree, faces, TreeStyle
# Loads an example tree
nw = """
(((Dre:0.008339,Dme:0.300613)1.000000:0.596401,
(Cfa:0.640858, Hsa:0.753230)1.000000:0.182035)1.000000:0.106234,
((Dre:0.271621,Cfa:0.046042)1.000000:0.953250,
(Hsa:0.061813, Mms:0.110769)1.000000:0.204419)1.000000:0.973467);
t = Tree(nw)
# You can create any random tree containing the same leaf names, and
# layout will work equally
# t = Tree()
# Creates a random tree with 8 leaves using a given set of names
# t.populate(8, ["Dme", "Dre", "Hsa", "Ptr", "Cfa", "Mms"])
# Set the path in which images are located
img path = "./"
# Create faces based on external images
humanFace = faces.ImgFace(img_path+"human.png")
mouseFace = faces.ImgFace(img_path+"mouse.png")
dogFace = faces.ImgFace(img_path+"dog.png")
chimpFace = faces.ImgFace(img_path+"chimp.png")
fishFace = faces.ImgFace(img_path+"fish.png")
flyFace = faces.ImgFace(img_path+"fly.png")
# Create a faces ready to read the name attribute of nodes
#nameFace = faces.TextFace(open("text").readline().strip(), fsize=20, fgcolor="#009
nameFace = faces.AttrFace("name", fsize=20, fgcolor="#009000")
# Create a conversion between leaf names and real names
code2name = {
        "Dre": "Drosophila melanogaster",
        "Dme": "Danio rerio",
        "Hsa": "Homo sapiens",
        "Ptr": "Pan troglodytes",
        "Mms": "Mus musculus",
        "Cfa": "Canis familiaris"
# Creates a dictionary with the descriptions of each leaf name
code2desc = {
        "Dre":""The zebrafish, also known as Danio rerio,
is a tropical freshwater fish belonging to the
minnow family (Cyprinidae).""",
        "Dme":""True flies are insects of the order Diptera,
possessing a single pair of wings on the
mesothorax and a pair of halteres, derived from
the hind wings, on the metathorax""",
        "Hsa":"""A human is a member of a species
of bipedal primates in the family Hominidae.""",
        "Ptr":"""Chimpanzee, sometimes colloquially
chimp, is the common name for the
two extant species of ape in the genus Pan.""",
```

```
"Mms":"""A mouse is a small mammal belonging to the
order of rodents.""",
        "Cfa": """The dog (Canis lupus familiaris) is a
domesticated subspecies of the Gray Wolf,
a member of the Canidae family of the
orderCarnivora."""
        }
# Creates my own layout function. I will use all previously created
# faces and will set different node styles depending on the type of
# node.
def mylayout(node):
    # If node is a leaf, add the nodes name and a its scientific
    if node.is_leaf():
        # Add an static face that handles the node name
        faces.add_face_to_node(nameFace, node, column=0)
        # We can also create faces on the fly
        longNameFace = faces.TextFace(code2name[node.name])
        faces.add_face_to_node(longNameFace, node, column=0)
        # text faces support multiline. We add a text face
        # with the whole description of each leaf.
        descFace = faces.TextFace(code2desc[node.name], fsize=10)
        descFace.margin_top = 10
        descFace.margin_bottom = 10
        descFace.border.margin = 1
        # Note that this faces is added in "aligned" mode
        faces.add_face_to_node(descFace, node, column=0, aligned=True)
        # Sets the style of leaf nodes
        node.img_style["size"] = 12
        node.img style["shape"] = "circle"
    #If node is an internal node
    else:
        # Sets the style of internal nodes
        node.img_style["size"] = 6
        node.img_style["shape"] = "circle"
        node.img_style["fgcolor"] = "#000000"
    # If an internal node contains more than 4 leaves, add the
    # images of the represented species sorted in columns of 2
    # images max.
    if len(node) >= 4:
        col = 0
        for i, name in enumerate(set(node.get_leaf_names())):
            if i > 0 and i % 2 == 0:
                col += 1
            # Add the corresponding face to the node
            if name.startswith("Dme"):
                faces.add_face_to_node(flyFace, node, column=col)
            elif name.startswith("Dre"):
                faces.add face to node(fishFace, node, column=col)
            elif name.startswith("Mms"):
                faces.add_face_to_node(mouseFace, node, column=col)
            elif name.startswith("Ptr"):
```

```
faces.add_face_to_node(chimpFace, node, column=col)
            elif name.startswith("Hsa"):
                faces.add_face_to_node(humanFace, node, column=col)
            elif name.startswith("Cfa"):
                faces.add_face_to_node(dogFace, node, column=col)
            # Modifies this node's style
            node.img_style["size"] = 16
            node.img_style["shape"] = "sphere"
            node.imq_style["fgcolor"] = "#AA0000"
    # If leaf is "Hsa" (homo sapiens), highlight it using a
    # different background.
    if node.is_leaf() and node.name.startswith("Hsa"):
        node.img_style["bgcolor"] = "#9db0cf"
# And, finally, Visualize the tree using my own layout function
ts = TreeStyle()
ts.layout_fn = mylayout
t.render("img_faces.png", w=600, tree_style = ts)
```

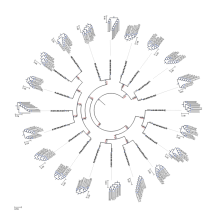
#### **Bubble tree maps**



import random from ete2 import Tree, TreeStyle, NodeStyle, faces, AttrFace, CircleFace def layout(node): if node.is\_leaf(): # Add node name to laef nodes N = AttrFace("name", fsize=14, fgcolor="black") faces.add\_face\_to\_node(N, node, 0) if "weight" in node.features: # Creates a sphere face whose size is proportional to node's # feature "weight" C = CircleFace(radius=node.weight, color="RoyalBlue", style="sphere") # Let's make the sphere transparent C.opacity = 0.3# And place as a float face over the tree faces.add\_face\_to\_node(C, node, 0, position="float") def get example tree(): # Random tree

```
t = Tree()
   t.populate(20, random_branches=True)
    # Some random features in all nodes
   for n in t.traverse():
       n.add_features(weight=random.randint(0, 50))
    # Create an empty TreeStyle
   ts = TreeStyle()
    # Set our custom layout function
   ts.layout_fn = layout
    # Draw a tree
   ts.mode = "c"
   # We will add node names manually
   ts.show_leaf_name = False
    # Show branch data
   ts.show_branch_length = True
   ts.show_branch_support = True
   return t, ts
if __name__ == "__main__":
   t, ts = get_example_tree()
    #t.render("bubble_map.png", w=600, dpi=300, tree_style=ts)
   t.show(tree_style=ts)
```

### **Trees within trees**

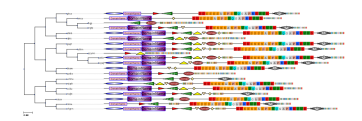


```
import random
from ete2 import Tree, TreeStyle, NodeStyle, faces, AttrFace, TreeFace
# Tree Style used to render small trees used as leaf faces
small_ts = TreeStyle()
small_ts.show_leaf_name = True
small_ts.scale = 10

def layout(node):
    if node.is_leaf():
```

```
# Add node name to laef nodes
        N = AttrFace("name", fsize=14, fgcolor="black")
        faces.add_face_to_node(N, node, 0)
        t = Tree()
        t.populate(10)
       T = TreeFace(t, small_ts)
        # Let's make the sphere transparent
       T.opacity = 0.8
        # And place as a float face over the tree
        faces.add_face_to_node(T, node, 1, position="aligned")
def get_example_tree():
    # Random tree
   t = Tree()
   t.populate(20, random_branches=True)
    # Some random features in all nodes
   for n in t.traverse():
       n.add_features(weight=random.randint(0, 50))
    # Create an empty TreeStyle
   ts = TreeStyle()
    # Set our custom layout function
   ts.layout_fn = layout
    # Draw a tree
   ts.mode = "c"
    # We will add node names manually
   ts.show_leaf_name = False
    # Show branch data
   ts.show_branch_length = True
   ts.show_branch_support = True
    return t, ts
if __name__ == "__main__":
   t, ts = get_example_tree()
    #t.render("tree_faces.png", w=600, dpi=300, tree_style=ts)
   t.show(tree_style=ts)
```

#### Phylogenetic trees and sequence domains



```
import sys
```

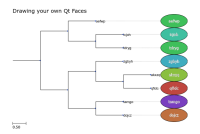
from ete2 import Tree, SeqMotifFace, TreeStyle, add\_face\_to\_node

```
seq = "LHGRISQQVEQSRSQVQAIGEKVSLAQAKIEKIKGSKKAIKVFSSAKYPAPERLQEYGSIFTDAQDPGLQRRPRHR
motifs = [
    # seq.start, seq.end, shape, width, height, fgcolor, bgcolor
    [10, 100, "[]", None, 10, "black", "rgradient:blue", "arial | 8 | white | domain Name
    [101, 150, "o", None, 10, "blue", "pink", None],
    [155, 180, "()", None, 10, "blue", "rgradient:purple", None],
    [160, 190, "^", None, 14, "black", "yellow", None],
    [172, 180, "v", None, 12, "black", "rgradient:orange", None],
    [185, 190, "o", None, 12, "black", "brown", None],
    [198, 200, "<>", None, 15, "black", "rgradient:gold", None],
    [210, 240, "compactseq", 2, 10, None, None, None],
    [300, 320, "seq", 10, 10, None, None, None],
    [310, 420, "<>", None, 30, "black", "rgradient:black", None],
    [1, 420, "()", None, 10, "red", "red", None],
[11, 30, "()", None, 20, "blue", "blue", None],
    [300, 310, "()", None, 40, "green", "green", None],
def layout (node):
    if node.is_leaf():
        seqFace = SeqMotifFace(seq, motifs, scale_factor=1)
        add_face_to_node(seqFace, node, 0, position="aligned")
def get_example_tree():
    # Create a random tree and add to each leaf a random set of motifs
    # from the original set
    t = Tree()
    t.populate(10)
    # for l in t.iter_leaves():
          seq_motifs = [list(m) for m in motifs] #sample(motifs, randint(2, len(mot
          seqFace = SeqMotifFace(seq, seq_motifs, intermotif_format="line",
                                  segtail_format="compactseg", scale_factor=1)
          seqFace.margin_bottom = 4
          f = l.add_face(seqFace, 0, "aligned")
    ts = TreeStyle()
    ts.layout_fn = layout
    return t, ts
if __name__ == '__main__':
    t, ts = get_example_tree()
    t.render("motifs.png", w=1200, dpi=300, tree_style=ts)
    #t.show(tree_style=ts)
```

#### Creating your custom interactive Item faces

Note that item faces shown in this image are not static. When the tree is view using the tree.show() method, you can interact with items.

```
# We will need to create Qt4 items
from PyQt4 import QtCore
from PyQt4.QtGui import QGraphicsRectItem, QGraphicsSimpleTextItem, \
```



QGraphicsEllipseItem, QColor, QPen, QBrush from ete2 import Tree, faces, TreeStyle, NodeStyle # To play with random colors import colorsys import random class InteractiveItem (QGraphicsRectItem): def \_\_init\_\_(self, \*arg, \*\*karg): QGraphicsRectItem.\_\_init\_\_(self, \*arg, \*\*karg) self.node = None self.label = None self.setCursor(QtCore.Qt.PointingHandCursor) self.setAcceptsHoverEvents(True) def hoverEnterEvent (self, e): # There are many ways of adding interactive elements. With the # following code, I show/hide a text item over my custom # DynamicItemFace if not self.label: self.label = QGraphicsRectItem() self.label.setParentItem(self) # This is to ensure that the label is rendered over the # rest of item children (default ZValue for items is 0) self.label.setZValue(1) self.label.setBrush(QBrush(QColor("white"))) self.label.text = QGraphicsSimpleTextItem() self.label.text.setParentItem(self.label) self.label.text.setText(self.node.name) self.label.setRect(self.label.text.boundingRect()) self.label.setVisible(True) def hoverLeaveEvent(self, e): if self.label: self.label.setVisible(False) def random\_color(h=None): """Generates a random color in RGB format.""" if not h: h = random.random() s = 0.51 = 0.5return \_hls2hex(h, l, s)

return '#%02x%02x%02x' %tuple (map (lambda x: int (x\*255),

**def** \_hls2hex(h, l, s):

```
colorsys.hls_to_rgb(h, l, s)))
def ugly_name_face(node, *args, **kargs):
    """ This is my item generator. It must receive a node object, and
    returns a Qt4 graphics item that can be used as a node face.
    # receive an arbitrary number of arguments, in this case width and
    # height of the faces
   width = args[0][0]
   height = args[0][1]
    ## Creates a main master Item that will contain all other elements
    ## Items can be standard QGraphicsItem
    # masterItem = QGraphicsRectItem(0, 0, width, height)
    # Or your custom Items, in which you can re-implement interactive
    # functions, etc. Check QGraphicsItem doc for details.
   masterItem = InteractiveItem(0, 0, width, height)
    # Keep a link within the item to access node info
   masterItem.node = node
    # I dont want a border around the masterItem
   masterItem.setPen(QPen(QtCore.Qt.NoPen))
    # Add ellipse around text
    ellipse = QGraphicsEllipseItem(masterItem.rect())
    ellipse.setParentItem(masterItem)
    # Change ellipse color
    ellipse.setBrush(QBrush(QColor( random color())))
    # Add node name within the ellipse
   text = QGraphicsSimpleTextItem(node.name)
   text.setParentItem(ellipse)
    text.setPen(QPen(QColor("white"))))
    # Center text according to masterItem size
   tw = text.boundingRect().width()
   th = text.boundingRect().height()
    center = masterItem.boundingRect().center()
   text.setPos(center.x()-tw/2, center.y()-th/2)
   return masterItem
def master_ly(node):
    if node.is_leaf():
        # Create an ItemFAce. First argument must be the pointer to
        # the constructor function that returns a QGraphicsItem. It
        # will be used to draw the Face. Next arguments are arbitrary,
        # and they will be forwarded to the constructor Face function.
        F = faces.DynamicItemFace(ugly_name_face, 100, 50)
        faces.add face to node (F, node, 0, position="aligned")
def get_example_tree():
    t = Tree()
```

```
t.populate(8, reuse_names=False)

ts = TreeStyle()
  ts.layout_fn = master_ly
  ts.title.add_face(faces.TextFace("Drawing your own Qt Faces", fsize=15), 0)
  return t, ts

if __name__ == "__main__":
  t, ts = get_example_tree()

#t.render("item_faces.png", h=400, tree_style=ts)
  # The interactive features are only available using the GUI
  t.show(tree_style=ts)
```

# 2.3 Phylogenetic Trees

### **Contents**

- Phylogenetic Trees
  - Overview
  - Linking Phylogenetic Trees with Multiple Sequence Alignments
  - Visualization of phylogenetic trees
  - Adding taxonomic information
    - \* Automatic control of species info
    - \* Automatic (and custom) control of the species info
    - \* Manual control of the species info
  - Detecting evolutionary events
    - \* Species Overlap (SO) algorithm
    - \* Tree reconciliation algorithm
    - \* A closer look to the evolutionary event object
  - Relative dating phylogenetic nodes
    - \* Implementation
  - Automatic rooting (outgroup detection)
  - Working with duplicated gene families
    - \* Treeko (splitting gene trees into species trees)
    - \* Splitting gene trees by duplication events
    - \* Collapse species specific duplications

#### 2.3.1 Overview

Phylogenetic trees are the result of most evolutionary analyses. They represent the evolutionary relationships among a set of species or, in molecular biology, a set of homologous sequences.

The PhyloTree class is an extension of the base Tree object, providing a appropriate way to deal with phylogenetic trees. Thus, while leaves are considered to represent species (or sequences from a given species genome), internal nodes are considered ancestral nodes. A direct consequence of this is, for instance, that every split in the tree will represent a speciation or duplication event.

# 2.3.2 Linking Phylogenetic Trees with Multiple Sequence Alignments

PhyloTree instances allow molecular phylogenies to be linked to the Multiple Sequence Alignments (MSA). To associate a MSA with a phylogenetic tree you can use the PhyloNode.link\_to\_alignment() method. You can use the alg\_format argument to specify its format (See SeqGroup documentation for available formats)

Given that Fasta format are not only applicable for MSA but also for **Unaligned Sequences**, you may also associate sequences of different lengths with tree nodes.

```
from ete2 import PhyloTree
fasta_txt = """
>seqA
MAEIPDETIQQFMALT---HNIAVQYLSEFGDLNEALNSYYASQTDDIKDRREEAH
>seqB
MAEIPDATIQQFMALTNVSHNIAVQY--EFGDLNEALNSYYAYQTDDQKDRREEAH
>seqC
MAEIPDATIQ---ALTNVSHNIAVQYLSEFGDLNEALNSYYASQTDDQPDRREEAH
>seqD
MAEAPDETIQQFMALTNVSHNIAVQYLSEFGDLNEAL------REEAH
"""
# Load a tree and link it to an alignment.
t = PhyloTree("(((seqA, seqB), seqC), seqD);")
t.link_to_alignment(alignment=fasta_txt, alg_format="fasta")
```

The same could be done at the same time the tree is being loaded, by using the alignment and alg\_format arguments of PhyloTree.

```
# Load a tree and link it to an alignment.
t = PhyloTree("(((seqA,seqB),seqC),seqD);", alignment=fasta_txt, alg_format="fasta")
```

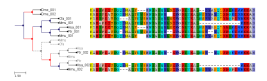
As currently implemented, sequence linking process is not strict, which means that a perfect match between all node names and sequences names is not required. Thus, if only one match is found between sequences names within the MSA file and tree node names, only one tree node will contain an associated sequence. Also, it is important to note that sequence linking is not limited to terminal nodes. If internal nodes are named, and such names find a match within the provided MSA file, their corresponding sequences will be also loaded into the tree structure. Once a MSA is linked, sequences will be available for every tree node through its node.sequence attribute.

```
LTNVSHOFMA LTNVSH
            LTNVSH----
            LTNVSH----
             ----FMA LTNVSH
# Load a tree and link it to an alignment. As usual, 'alignment' can
# be the path to a file or data in text format.
t = PhyloTree("(((seqA,seqB),seqC),seqD);", alignment=fasta_txt, alg_format="fasta")
#We can now access the sequence of every leaf node
print "These are the nodes and its sequences:"
for leaf in t.iter_leaves():
   print leaf.name, leaf.sequence
#seqD MAEAPDETIQQFMALTNVSHNIAVQYLSEFGDLNEAL-----REEAH
#seqC MAEIPDATIQ---ALTNVSHNIAVQYLSEFGDLNEALNSYYASQTDDQPDRREEAH
#seqA MAEIPDETIQQFMALT---HNIAVQYLSEFGDLNEALNSYYASQTDDIKDRREEAH
#seqB MAEIPDATIQQFMALTNVSHNIAVQY--EFGDLNEALNSYYAYQTDDQKDRREEAH
# The associated alignment can be changed at any time
t.link_to_alignment(alignment=iphylip_txt, alg_format="iphylip")
# Let's check that sequences have changed
print "These are the nodes and its re-linked sequences:"
for leaf in t.iter leaves():
   print leaf.name, leaf.sequence
#seqD MAEAPDETIQQFMALTNVSHNIAVQYLSEFGDLNEAL------REEAHQ-----FMALTNVSH
#seqC MAEIPDATIQ---ALTNVSHNIAVQYLSEFGDLNEALNSYYASQTDDQPDRREEAHQFMALTNVSH-------
#seqA MAEIPDETIQQFMALT---HNIAVQYLSEFGDLNEALNSYYASQTDDIKDRREEAHQFMALTNVSHQFMALTNVSH
#seqB MAEIPDATIQQFMALTNVSHNIAVQY--EFGDLNEALNSYYAYQTDDQKDRREEAHQFMALTNVSH-------
# The sequence attribute is considered as node feature, so you can
# even include sequences in your extended newick format!
print t.write(features=["sequence"], format=9)
# (((seqA[&&NHX:sequence=MAEIPDETIQQFMALT---HNIAVQYLSEFGDLNEALNSYYASQTDDIKDRREEAHQF
# MALTNVSHQFMALTNVSH], seqB[&&NHX:sequence=MAEIPDATIQQFMALTNVSHNIAVQY--EFGDLNEALNSY
# YAYQTDDQKDRREEAHQFMALTNVSH-----]), seqC[&&NHX:sequence=MAEIPDATIQ---ALTNVSHNIA
# VQYLSEFGDLNEALNSYYASQTDDQPDRREEAHQFMALTNVSH-----]),seqD[&&NHX:sequence=MAEAPD
# ETIQOFMALTNVSHNIAVOYLSEFGDLNEAL------REEAHQ-----FMALTNVSH]);
# And yes, you can save this newick text and reload it into a PhyloTree instance.
sametree = PhyloTree(t.write(features=["sequence"]))
print "Recovered tree with sequence features:"
print sametree
#
#
           \-seqD
print "seqA sequence:", (t&"seqA").sequence
\# \ \ \mathsf{MAEIPDETIQQFMALT} -- + \mathsf{HNIAVQYLSEFGDLNEALNSYYASQTDDIKDRREEAHQFMALTNVSHQFMALTNVSH}
```

# 2.3.3 Visualization of phylogenetic trees

PhyloTree instances can benefit from all the features of the programmable drawing engine. However, a built-in phylogenetic layout is provided for convenience.

All PhyloTree instances are, by default, attached to such layout for tree visualization, thus allowing for in-place alignment visualization and evolutionary events labeling.



```
from ete2 import PhyloTree, TreeStyle
alg = """
 >Dme_001
MAEIPDETIQQFMALT---HNIAVQYLSEFGDLNEAL--YYASQTDDIKDRREEAH
MAEIPDATIQQFMALTNVSHNIAVQY--EFGDLNEALNSYYAYQTDDQKDRREEAH
 MAEIPDATIQ---ALTNVSHNIAVQYLSEFGDLNEALNSYYASQTDDQPDRREEAH
 >Mms_001
 MAEAPDETIQQFMALTNVSHNIAVQYLSEFGDLNEAL-----REEAH
 >Hsa_001
MAEIPDETIQQFMALT---HNIAVQYLSEFGDLNEALNSYYASQTDDIKDRREEAH
 MAEIPDATIQ-FMALTNVSHNIAVQY--EFGDLNEALNSY--YQTDDQKDRREEAH
 >Mmu 002
 MAEIPDATIQ---ALTNVSHNIAVQYLSEFGDLNEALNSYYASQTDDQPDRREEAH
 >Hsa 002
 MAEAPDETIQQFM-LTNVSHNIAVQYLSEFGDLNEAL-----REEAH
 >Mmu_001
MAEIPDETIQQFMALT---HNIAVQYLSEFGDLNEALNSYYASQTDDIKDRREEAH
MAEIPDATIQ-FMALTNVSHNIAVQY--EFGDLNEALNSY--YQTDDQKDRREEAH
>Mmu 001
MAEIPDATIQ---ALTNVSHNIAVQYLSEFGDLNEALNSYYASQTDDQPDRREEAH
def get_example_tree():
    # Performs a tree reconciliation analysis
    gene_tree_nw = '((Dme_001,Dme_002),(((Cfa_001,Mms_001),((Hsa_001,Ptr_001),Mmu_001)),
    species_tree_nw = "((((Hsa, Ptr), Mmu), (Mms, Cfa)), Dme);"
    genetree = PhyloTree(gene_tree_nw)
    sptree = PhyloTree(species_tree_nw)
    recon_tree, events = genetree.reconcile(sptree)
    recon_tree.link_to_alignment(alg)
    return recon_tree, TreeStyle()
if __name__ == "__main__":
    # Visualize the reconciled tree
    t, ts = get_example_tree()
```

t.show(tree\_style=ts)

#recon\_tree.render("phylotree.png", w=750)

# 2.3.4 Adding taxonomic information

PhyloTree instances allow to deal with leaf names and species names separately. This is useful when working with molecular phylogenies, in which node names usually represent sequence identifiers. Species names will be stored in the PhyloNode.species attribute of each leaf node. The method PhyloNode.get\_species() can be used obtain the set of species names found under a given internal node (speciation or duplication event). Often, sequence names do contain species information as a part of the name, and ETE can parse this information automatically.

There are three ways to establish the species of the different tree nodes:

- Default: The three first letters of node's name represent the species
- The species code of each node is dynamically created based on node's name
- The species code of each node is manually set.

#### **Automatic control of species info**

```
from ete2 import PhyloTree
# Reads a phylogenetic tree (using default species name encoding)
t = PhyloTree("(((Hsa_001,Ptr_001),(Cfa_001,Mms_001)),(Dme_001,Dme_002));")
                                /-Hsa 001
                                \-Ptr 001
                                \-Mms 001
                     /-Dme_001
                      \-Dme_002
# Prints current leaf names and species codes
print "Deafult mode:"
for n in t.get_leaves():
   print "node:", n.name, "Species name:", n.species
# node: Dme_001 Species name: Dme
# node: Dme_002 Species name: Dme
# node: Hsa_001 Species name: Hsa
# node: Ptr_001 Species name: Ptr
# node: Cfa_001 Species name: Cfa
# node: Mms 001 Species name: Mms
```

#### Automatic (and custom) control of the species info

The default behavior can be changed by using the PhyloNode.set\_species\_naming\_function() method or by using the sp\_naming\_function argument of the PhyloTree class. Note that, using the sp\_naming\_function argument, the whole tree structure will be initialized to use the provided parsing function to obtain species name information. PhyloNode.set\_species\_naming\_function() (present in all tree nodes) can be used to change the behavior in a previously loaded tree, or to set different parsing function to different parts of the tree.

```
from ete2 import PhyloTree
# Reads a phylogenetic tree
t = PhyloTree("(((Hsa 001,Ptr 001),(Cfa 001,Mms 001)),(Dme 001,Dme 002));")
# Let's use our own leaf name parsing function to obtain species
# names. All we need to do is create a python function that takes
# node's name as argument and return its corresponding species name.
def get_species_name(node_name_string):
    # Species code is the first part of leaf name (separated by an
    # underscore character)
    spcode = node name string.split(" ")[0]
    # We could even translate the code to complete names
    code2name = {
      "Dme": "Drosophila melanogaster",
      "Hsa": "Homo sapiens",
      "Ptr": "Pan troglodytes",
      "Mms":"Mus musculus",
      "Cfa": "Canis familiaris"
    return code2name[spcode]
# Now, let's ask the tree to use our custom species naming function
t.set_species_naming_function(get_species_name)
print "Custom mode:"
for n in t.get_leaves():
    print "node:", n.name, "Species name:", n.species
# node: Dme_001 Species name: Drosophila melanogaster
# node: Dme_002 Species name: Drosophila melanogaster
# node: Hsa_001 Species name: Homo sapiens
# node: Ptr_001 Species name: Pan troglodytes
# node: Cfa_001 Species name: Canis familiaris
# node: Mms_001 Species name: Mus musculus
```

### Manual control of the species info

To disable the automatic generation of species names based on node names, a None value can be passed to the PhyloNode.set\_species\_naming\_function() function. From then on, species attribute will not be automatically updated based on the name of nodes and it could be controlled manually.

```
from ete2 import PhyloTree
# Reads a phylogenetic tree
t = PhyloTree("(((Hsa_001,Ptr_001),(Cfa_001,Mms_001)),(Dme_001,Dme_002));")
# Of course, you can disable the automatic generation of species
# names. To do so, you can set the species naming function to
# None. This is useful to set the species names manually or for
# reading them from a newick file. Other wise, species attribute would
# be overwriten
mynewick = """
(((Hsa_001[&&NHX:species=Human],Ptr_001[&&NHX:species=Chimp]),
(Cfa_001[&&NHX:species=Dog],Mms_001[&&NHX:species=Mouse])),
(Dme_001[&&NHX:species=Fly],Dme_002[&&NHX:species=Fly]));
"""
t = PhyloTree(mynewick, sp_naming_function=None)
```

```
print "Disabled mode (manual set)"
for n in t.get_leaves():
    print "node:", n.name, "Species name:", n.species

# node: Dme_001 Species name: Fly
# node: Dme_002 Species name: Fly
# node: Hsa_001 Species name: Human
# node: Ptr_001 Species name: Chimp
# node: Cfa_001 Species name: Dog
# node: Mms_001 Species name: Mouse
```

Full Example: Species aware trees.

### 2.3.5 Detecting evolutionary events

There are several ways to automatically detect duplication and speciation nodes. ETE provides two methodologies: One implements the algorithm described in Huerta-Cepas (2007) and is based on the species overlap (SO) between partitions and thus does not depend on the availability of a species tree. The second, which requires the comparison between the gene tree and a previously defined species tree, implements a strict tree reconciliation algorithm (Page and Charleston, 1997). By detecting evolutionary events, orthology and paralogy relationships among sequences can also be inferred. Find a comparison of both methods in Marcet-Houben and Gabaldon (2009).

### Species Overlap (SO) algorithm

In order to apply the SO algorithm, you can use the PhyloNode.get\_descendant\_evol\_events() method (it will detect all evolutionary events under the current node) or the PhyloNode.get\_my\_evol\_events() method (it will detect only the evolutionary events in which current node, a leaf, is involved).

By default the **species overlap score** (**SOS**) **threshold** is set to 0.0, which means that a single species in common between two node branches will rise a duplication event. This has been shown to perform the best with real data, however you can adjust the threshold using the <code>sos\_thr</code> argument present in both methods.

```
\-Mmu 001
                                          \-Mmu_002
# To obtain all the evolutionary events involving a given leaf node we
# use get_my_evol_events method
matches = t.search_nodes(name="Hsa_001")
human_seq = matches[0]
# Obtains its evolutionary events
events = human_seq.get_my_evol_events()
# Print its orthology and paralogy relationships
print "Events detected that involve Hsa_001:"
for ev in events:
    if ev.etype == "S":
       print ' ORTHOLOGY RELATIONSHIP:', ','.join(ev.in_seqs), "<====>", ','.join(ev.
    elif ev.etype == "D":
                PARALOGY RELATIONSHIP:', ','.join(ev.in_seqs), "<====>", ','.join(ev.o
# Alternatively, you can scan the whole tree topology
events = t.get_descendant_evol_events()
# Print its orthology and paralogy relationships
print "Events detected from the root of the tree"
for ev in events:
    if ev.etype == "S":
       print ' ORTHOLOGY RELATIONSHIP:', ','.join(ev.in_seqs), "<====>", ','.join(ev.
    elif ev.etype == "D":
                PARALOGY RELATIONSHIP:', ','.join(ev.in_seqs), "<====>", ','.join(ev.o
# If we are only interested in the orthology and paralogy relationship
# among a given set of species, we can filter the list of sequences
# fseqs is a function that, given a list of sequences, returns only
# those from human and mouse
fseqs = lambda slist: [s for s in slist if s.startswith("Hsa") or s.startswith("Mms")]
print "Paralogy relationships among human and mouse"
for ev in events:
    if ev.etype == "D":
        # Prints paralogy relationships considering only human and
        # mouse. Some duplication event may not involve such species,
        # so they will be empty
        print ' PARALOGY RELATIONSHIP:', \
            ','.join(fseqs(ev.in_seqs)), \
            "<===>",\
            ','.join(fseqs(ev.out_seqs))
# Note that besides the list of events returned, the detection
# algorithm has labeled the tree nodes according with the
# predictions. We can use such lables as normal node features.
dups = t.search_nodes(evoltype="D") # Return all duplication nodes
```

#### Tree reconciliation algorithm

Tree reconciliation algorithm uses a predefined species tree to infer all the necessary genes losses that explain a given gene tree topology. Consequently, duplication and separation nodes will strictly follow the species tree topology.

To perform a tree reconciliation analysis over a given node in a molecular phylogeny you can use the PhyloNode.reconcile() method, which requires a species PhyloTree as its first argument. Leaf node names in the species are expected to be the same species codes in the gene tree (see taxonomic\_info). All species codes present in the gene tree should appear in the species tree.

As a result, the PhyloNode.reconcile() method will label the original gene tree nodes as duplication or speciation, will return the list of inferred events, and will return a new **reconciled tree** (PhyloTree instance), in which inferred gene losses are present and labeled.

```
from ete2 import PhyloTree
```

```
# Loads a gene tree and its corresponding species tree. Note that
# species names in sptree are the 3 firs letters of leaf nodes in
# genetree.
gene_tree_nw = '((Dme_001,Dme_002),(((Cfa_001,Mms_001),((Hsa_001,Ptr_001),Mmu_001))),(Ptr_001)
species_tree_nw = "((((Hsa, Ptr), Mmu), (Mms, Cfa)), Dme);"
genetree = PhyloTree(gene_tree_nw)
sptree = PhyloTree(species_tree_nw)
print genetree
                      /-Dme 001
                      \-Dme_002
                                            -Cfa_001
                                           \-Mms_001
                                                      -Hsa_001
                                                     \-Ptr_001
                                           \-Mmu_001
                                 /-Ptr<u>002</u>
                                           \-Mmu 002
# Let's reconcile our genetree with the species tree
recon_tree, events = genetree.reconcile(sptree)
# a new "reconcilied tree" is returned. As well as the list of
# inferred events.
print "Orthology and Paralogy relationships:"
for ev in events:
    if ev.etype == "S":
        print 'ORTHOLOGY RELATIONSHIP:', ','.join(ev.inparalogs), "<====>", ','.join(ev.
    elif ev.etype == "D":
        print 'PARALOGY RELATIONSHIP:', ','.join(ev.inparalogs), "<====>", ','.join(ev.org)
# And we can explore the resulting reconciled tree
print recon_tree
```

```
# You will notice how the reconcilied tree is the same as the gene
  tree with some added branches. They are inferred gene losses.
                       '-Dme_001
                       \-Dme_002
                                              -Cfa_001
                                             \-Mms 001
                                                        /-Hsa<u> </u>001
                                                       \-Ptr_001
                                            \-Mmu_001
                                            /-Mms
                                             \-Cfa
                                                                  /-Hsa
                                                                  \-Ptr_002
                                                       \-Mmu
                                                                  /-Ptr
                                                                  \-Hsa 002
                                                       \-Mmu 002
# And we can visualize the trees using the default phylogeny
# visualization layout
genetree.show()
recon_tree.show()
```

#### A closer look to the evolutionary event object

Both methods, species overlap and tree reconciliation, can be used to label each tree node as a duplication or speciation event. Thus, the PhyloNode.evoltype attribute of every node will be set to one of the following states: D (Duplication), S (Speciation) or L gene loss.

Additionally, a list of all the detected events is returned. Each event is a python object of type phylo.EvolEvent, containing some basic information about each event (etype, in\_seqs, out\_seqs, node):

If an event represents a duplication, in\_seqs are all paralogous to out\_seqs. Similarly, if an event represents a speciation, in\_seqs are all orthologous to out\_seqs.

# 2.3.6 Relative dating phylogenetic nodes

In molecular phylogeny, nodes can be interpreted as evolutionary events. Therefore, they represent duplication or speciation events. In the case of gene duplication events, nodes can also be assigned to a certain point in a relative temporal scale. In other words, you can obtain a relative dating of all the duplication events detected.

Although **absolute dating is always preferred and more precise**, topological dating provides a faster approach to compare the relative age of paralogous sequences (read this for a comparison with other methods, such as the use of synonymous substitution rates as a proxy to the divergence time).

Some applications of topological dating can be found in Huerta-Cepas et al, 2007 or, more recently, in Huerta-Cepas et al, 2011 or Kalinka et al, 2001.

#### **Implementation**

The aim of relative dating is to establish a gradient of ages among sequences. For this, a reference species needs to be fixed, so the gradient of ages will be referred to that referent point.

Thus, if our reference species is *Human*, we could establish the following gradient of species:

• (1) Human -> (2) Other Primates -> (3) Mammals -> (4) Vertebrates

So, nodes in a tree can be assigned to one of the above categories depending on the sequences grouped. For instance:

- A node with only human sequences will be mapped to (1).
- A node with human and orangutan sequences will be mapped to (2)
- A node with human a fish sequences will be mapped to (4)

This simple calculation can be done automatically by encoding the gradient of species ages as Python dictionary.

```
relative_dist = {
    "human": 0, # human
    "chimp": 1, # Primates non human
    "rat": 2, # Mammals non primates
    "mouse": 2, # Mammals non primates
    "fish": 3 # Vertebrates non mammals
}
```

Once done, ETE can check the relative age of any tree node. The  $PhyloNode.get_age()$  method can be used to that purpose.

For example, let's consider the following gene tree:

```
# | \-fish
#-Dup3
# | /-humanC
# | /---| \-chimpC
# | | |
# \Dup2 \-humanD
# | |
# | /-ratC
# \---|
# \-mouseC
```

the expected node dating would be:

- Dup1 will be assigned to primates (most distant species is chimp). Dup1.get\_age(relative\_distances) will return 1
- Dup2 will be assigned to mammals [2] (most distant species are rat and mouse). Dup2.get\_age(relative\_distances) will return 2
- Dup3 will be assigned to mammals [3] (most distant species is fish). Dup3.get\_age(relative\_distances) will return 3

```
from ete2 import PhyloTree
# Creates a gene phylogeny with several duplication events at
# different levels. Note that we are using the default method for
# detecting the species code of leaves (three first lettes in the node
# name are considered the species code).
nw = """
((Dme_001,Dme_002),(((Cfa_001,Mms_001),((((Hsa_001,Hsa_003),Ptr_001)
,Mmu_001),((Hsa_004,Ptr_004),Mmu_004))),(Ptr_002,(Hsa_002,Mmu_002))));
t = PhyloTree(nw)
print "Original tree:",
print t
#
#
              /-Dme_001
               \-Dme_002
                                   /-Cfa_001
                                   \-Mms_001
                                                                   ′-Hsa 001
                                                                   \-Hsa_003
                                                        \-Ptr_001
                                              \-Mmu 001
                                                         -Hsa_004
                                                        \-Ptr_004
                                              \-Mmu_004
```

```
#
# Create a dictionary with relative ages for the species present in
# the phylogenetic tree. Note that ages are only relative numbers to
# define which species are older, and that different species can
# belong to the same age.
species2age = {
  'Hsa': 1, # Homo sapiens (Hominids)
  'Ptr': 2, # P. troglodytes (primates)
  'Mmu': 2, # Macaca mulata (primates)
  'Mms': 3, # Mus musculus (mammals)
  'Cfa': 3, # Canis familiaris (mammals)
  'Dme': 4  # Drosophila melanogaster (metazoa)
}
# We can translate each number to its correspondig taxonomic number
age2name = {
 1: "hominids",
 2:"primates"
  3:"mammals".
  4:"metazoa"
}
event1= t.get_common_ancestor("Hsa_001", "Hsa_004")
event2=t.get_common_ancestor("Hsa_001", "Hsa_002")
print
print "The duplication event leading to the human sequences Hsa_001 and "+\
    "Hsa_004 is dated at: ", age2name[event1.get_age(species2age)]
print "The duplication event leading to the human sequences Hsa_001 and "+\
    "Hsa_002 is dated at: ", age2name[event2.get_age(species2age)]
# The duplication event leading to the human sequences Hsa_001 and Hsa_004
# is dated at: primates
# The duplication event leading to the human sequences Hsa_001 and Hsa_002
# is dated at: mammals
```

**Warning:** Note that relative distances will vary depending on your reference species.

## 2.3.7 Automatic rooting (outgroup detection)

Two methods are provided to assist in the automatic rooting of phylogenetic trees. Since tree nodes contain relative age information (based on the species code autodetection), the same relative age dictionaries can be used to detect the farthest and oldest node in a tree to given sequences.

PhyloNode.get\_farthest\_oldest\_node() and PhyloNode.get\_farthest\_oldest\_leaf() can be used for that purpose.

# 2.3.8 Working with duplicated gene families

#### Treeko (splitting gene trees into species trees)

Comparisons between tree topologies provide important information for many evolutionary studies. Treeko (Marcet and Gabaldon, 2011) is a novel method that allows the comparison of any two tree topologies, even those with missing leaves and duplications. This is important in genome-wide analysis since many trees do not have exact leaf pairings and therefore most tree comparison methods are rendered useless.

Although Treeko is available as a standalone package, it uses ETE to generate all possible species tree topologies within a duplicated gene family tree.

Thus, the ETE method PhyloNode.get\_speciation\_trees() is expected to provide the core functionality required to perform a Treeko analysis. When used, the method will return a list of all possible species trees observed after combining the different non-duplicated subparts under a gene family tree node.

Duplication events will be automatically identified using the species overlap algorithm described within this manual. However, duplication nodes can be manually labeled and used by disabling the autodetect\_duplication flag.

Because of the combinatorial background of the Treeko method, the number of speciation trees generated by this function may vary enormously (ranging from few hundreds to tens of thousands topologies).

Here is a basic example on how to use it:

```
from ete2 import PhyloTree
t = PhyloTree("((((Human_1, Chimp_1), (Human_2, (Chimp_2, Chimp_3))), ((Fish_1, (Human_3)))
t.set_species_naming_function(lambda node: node.name.split("_")[0] )
print t.get_ascii(attributes=["name", "species"], show_internal=False )
             /-Human_1, Human
#
              \-Chimp_1, Chimp
              /-Human_2, Human
                 /-Chimp_2, Chimp
                 \-Chimp_3, Chimp
              /-Fish_1, Fish
                 /-Human_3, Human
                 \-Fish\_3, Fish
           \-Yeast_2, Yeast
     \-Yeast 1, Yeast
# We obtain a list of species trees inferred from the duplication
# events. Note that species specific duplications are ignored.
ntrees, ndups, sptrees = t.get_speciation_trees()
```

```
print "Found %d species trees and %d duplication nodes" %(ntrees, ndups)
for spt in sptrees:
   print spt
# Found 5 species trees and 4 duplication nodes
#
     /-Human 1
#
#
     \-Chimp 1
     /-Human 2
#
       /-Chimp_2
        \-Chimp_3
#
     /-Fish_1
     \-Yeast 2
        /-Human 3
        \-Fish 3
     \-Yeast_2
# --Yeast_1
```

**Note:** For performance reasons, species trees are created without any link to the original gene family tree, rather than the species name of each node. However, the map\_features attribute can be used to keep certain attributes of the original tree into the generated species trees.

**Note:** Although the efficiency of the method to generate all possible trees has been significantly improved from ETE version 2.2, creating thousands of new PhyloTree objects could affect performance. The flag newick\_only is now available to limit the output to a newick string per generated tree, thus improving the speed they can be processed or dumped into a file.

#### Splitting gene trees by duplication events

A much simpler approach to separate duplicates within the same gene family tree is to split the topology by their duplication nodes. For this, the method PhyloNode.split\_by\_dups() is provided.

```
/-Chimp_2, Chimp
                  \-Chimp_3, Chimp
              /-Fish_1, Fish
                 /-Human_3, Human
                 \-Fish_3, Fish
           \-Yeast_2, Yeast
     \-Yeast_1, Yeast
# Again, species specific duplications are ignored
for node in t.split_by_dups():
    print node
#
     /-Human 1
     \-Chimp 1
#
     /-Human_2
        /-Chimp_2
        \-Chimp_3
# --Yeast_2
# --Fish_1
     /-Human_3
     \-Fish\_3
# --Yeast_1
```

### Collapse species specific duplications

The method PhyloNode.collapse\_lineage\_specific\_expansions() method, which returns a pruned version of a tree, where nodes representing lineage specific expansions are converted into a single leaf node is also available.

From the previous examples, the lineage specific duplication of Chimp\_1 and Chimp\_2 could be easily collapsed into a single node.

```
/ \-Chimp_1, Chimp
             /-Human_2, Human
               /-Chimp_2, Chimp
               \-Chimp_3, Chimp
             /-Fish_1, Fish
            / /-Human_3, Human
             \-Fish\_3, Fish
          \-Yeast_2, Yeast
    \-Yeast_1, Yeast
t2 = t.collapse_lineage_specific_expansions()
print t2.get_ascii(attributes=["name", "species"], show_internal=False )
             /-Human_1, Human
             \-Chimp_1, Chimp
         / /-Human_2, Human
             \-Chimp_2, Chimp
            /-Fish_1, Fish
        \-Fish_3, Fish
          \-Yeast_2, Yeast
    \-Yeast_1, Yeast
```

# 2.4 Clustering Trees

### **Contents**

- Clustering Trees
  - Overview
  - Loading ClusterTrees
  - Visualization of matrix associated Trees
  - Cluster Validation Example

#### 2.4.1 Overview

Cluster analysis is the assignment of a set of observations into subsets (called clusters) so that observations in the same cluster are similar in some sense. Clustering is a method of unsupervised learning, and a common technique for statistical data analysis used in many fields, including machine learning, data mining, pattern recognition, image analysis and bioinformatics. Hierarchical clustering creates a hierarchy of clusters which may be represented in a tree structure called a dendrogram. The root of the tree consists of a single cluster containing all observations, and the leaves correspond to individual observations. [The Wikipedia project Jun-2009].

ETE provides special ClusterNode (alias ClusterTree) instances to deal with trees associated to a clustering analysis. The basic difference between Tree and ClusterTree is that leaf nodes in a cluster-tree are linked to numerical profiles. Such profiles are expected to represent the data used to generate the clustering tree. In other words, trees are bound to numerical arrays.

Based on this, ClusterTree instances provide several several clustering validation techniques that help in the analysis of cluster quality. Currently, inter and intra-cluster distances, cluster std.deviation, Silhouette analysis and Dunn indexes are supported. In addition, ClusterTree nodes can be visualized using the ProfileFace face type, which can represent cluster profiles in different ways, such as line plots, heatmaps or bar plots.

Although this type of trees are intended to be used for clustering results, any tree that can be linked to a table (i.e. phylogenetic profiles) could be loaded using this data type, thus taking advantage of the profile visualization modes, etc.

#### 2.4.2 Loading ClusterTrees

A ClusterTree can be linked to a numerical matrix by using the text\_array argument.

```
from ete2 import ClusterTree
```

```
# Example of a minimalistic numerical matrix. It is encoded as a text
# string for convenience, but it usally be loaded from a text file.
matrix = """
#Names\tcol1\tcol2\tcol3\tcol4\tcol5\tcol6\tcol7
A\t-1.23\t-0.81\t1.79\t0.78\t-0.42\t-0.69\t0.58
B\t-1.76\t-0.94\t1.16\t0.36\t0.41\t-0.35\t1.12
C\t-2.19\t0.13\t0.65\t-0.51\t0.52\t1.04\t0.36
D\t-1.22\t-0.98\t0.79\t-0.76\t-0.29\t1.54\t0.93
E\t-1.47\t-0.83\t0.85\t0.07\t-0.81\t1.53\t0.65
F\t-1.04\t-1.11\t0.87\t-0.14\t-0.80\t1.74\t0.48
G\t-1.57\t-1.17\t1.29\t0.23\t-0.20\t1.17\t0.26
H\t-1.53\t-1.25\t0.59\t-0.30\t0.32\t1.41\t0.77
print "Example numerical matrix"
print matrix
# #Names coll
                  col2
                         co13
                                  col4
                                          co15
                                                  col6
                                                          col7
```

```
1.79 0.78
# A
         -1.23
               -0.81
                                     -0.42
                                            -0.69
                                                    0.58
# B
        -1.76 -0.94 1.16
                             0.36
                                    0.41
                                            -0.35
                                                    1.12
# C
        -2.19 0.13 0.65
                             -0.51 0.52
                                            1.04
                                                    0.36
               -0.98 0.79
# D
        -1.22
                              -0.76 -0.29 1.54
                                                    0.93
# E
               -0.83 0.85 0.07
        -1.47
                                     -0.81
                                            1.53
                                                    0.65
# F
                              -0.14 \quad -0.80
        -1.04
               -1.11 0.87
                                            1.74
                                                    0.48
# G
         -1.57
               -1.17
                      1.29
                            0.23
                                     -0.20
                                            1.17
                                                    0.26
# H
        -1.53
               -1.25 0.59
                              -0.30 0.32
                                            1.41
                                                    0.77
# We load a tree structure whose leaf nodes correspond to rows in the
# numerical matrix. We use the text_array argument to link the tree
# with numerical matrix.
t = ClusterTree("(((A,B),(C,(D,E))),(F,(G,H)));", text_array=matrix)
```

Alternatively, you can re-link the tree (or a sub-part of it) to a new matrix using the ClusterNode.link\_to\_arraytable() method.

```
t = ClusterTree("(((A,B),(C,(D,E))),(F,(G,H)));")
t.children[0].link_to_arraytable(matrix1)
t.children[1].link_to_arraytable(matrix2)
```

Once the tree is linked to a table of profiles, the following node properties will be available: PhyloNode.profile, PhyloNode.deviation, PhyloNode.silhoutte, PhyloNode.intercluster\_dist, PhyloNode.intracluster\_dist, PhyloNode.dunn.

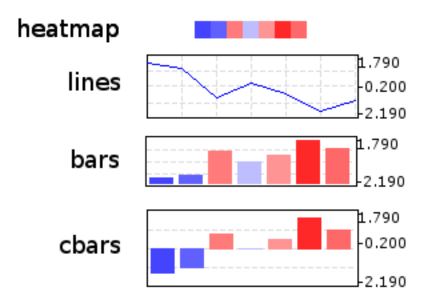
```
Similarly, the following methods are provide for convenience PhyloNode.iter_leaf_profiles(), PhyloNode.get_leaf_profiles(), PhyloNode.get_silhouette() and PhyloNode.get_dunn() methods.
```

#### 2.4.3 Visualization of matrix associated Trees

Clustering or not, any ClusterTree instance, associated to a numerical matrix, can be visualized together with the graphical representation of its node's numeric profiles. To this end, the ProfileFace class is provided by the treeview module. This face type can represent a node's numeric profile in four different ways:

Additionally, three basic layouts are provided that use different styles of ProfileFace instances: **heatmap**, **line\_profiles**, **bar\_profiles**, **cbar\_profiles**.

```
# Import Tree instance and faces module
from ete2 import ClusterTree
# Example of a minimalistic numerical matrix. It is encoded as a text
# string for convenience, but it usally be loaded from a text file.
matrix = """
#Names\tcol1\tcol2\tcol3\tcol4\tcol5\tcol6\tcol7
A\t-1.23\t-0.81\t1.79\t0.78\t-0.42\t-0.69\t0.58
B\t-1.76\t-0.94\t1.16\t0.36\t0.41\t-0.35\t1.12
C\t-2.19\t0.13\t0.65\t-0.51\t0.52\t1.04\t0.36
D\t-1.22\t-0.98\t0.79\t-0.76\t-0.29\t1.54\t0.93
E\t-1.47\t-0.83\t0.85\t0.07\t-0.81\t1.53\t0.65
F\t-1.04\t-1.11\t0.87\t-0.14\t-0.80\t1.74\t0.48
G\t-1.57\t-1.17\t1.29\t0.23\t-0.20\t1.17\t0.26
```



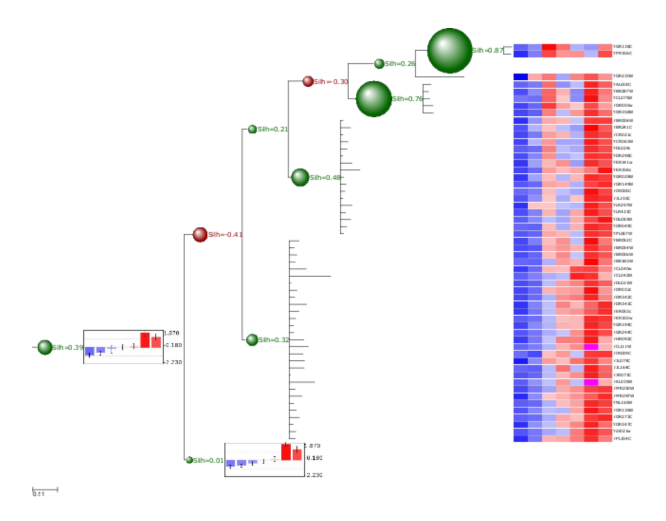
```
H\t-1.53\t-1.25\t0.59\t-0.30\t0.32\t1.41\t0.77
print "Example numerical matrix"
print matrix
# #Names col1
                  col2
                          co13
                                  col4
                                          co15
                                                  col6
                                                           co17
          -1.23
                  -0.81
                          1.79
                                  0.78
                                          -0.42
                                                   -0.69
                                                           0.58
# A
# B
          -1.76
                  -0.94
                          1.16
                                  0.36
                                          0.41
                                                   -0.35
                                                           1.12
# C
                                          0.52
          -2.19
                 0.13
                          0.65
                                  -0.51
                                                   1.04
                                                           0.36
#
 D
          -1.22
                  -0.98
                          0.79
                                  -0.76
                                          -0.29
                                                   1.54
                                                           0.93
#
 E
          -1.47
                  -0.83
                          0.85
                                  0.07
                                          -0.81
                                                   1.53
                                                           0.65
# F
          -1.04
                          0.87
                                          -0.80
                                                           0.48
                 -1.11
                                  -0.14
                                                  1.74
# G
          -1.57
                          1.29
                                  0.23
                                          -0.20
                                                  1.17
                                                           0.26
                  -1.17
# H
          -1.53
                  -1.25
                          0.59
                                  -0.30
                                          0.32
                                                   1.41
                                                           0.77
# We load a tree structure whose leaf nodes correspond to rows in the
# numerical matrix. We use the text array argument to link the tree
# with numerical matrix.
t = ClusterTree("(((A,B),(C,(D,E))),(F,(G,H)));", text_array=matrix)
t.show("heatmap")
t.show("cluster cbars")
t.show("cluster_bars")
t.show("cluster_lines")
```

## 2.4.4 Cluster Validation Example

If associated matrix represents the dataset used to produce a given tree, clustering validation values can be used to assess the quality of partitions. To do so, you will need to set the distance function that was used to calculate distances among items (leaf nodes). ETE implements three common distance methods in bioinformatics: **euclidean**, **pearson** correlation and **spearman** correlation distances.

In the following example, a microarray clustering result is visualized and validated using ETE.

Image resulting from a microarray clustering validation analysis. Red bubbles represent a bad silhouette index (S<0), while green represents good silhouette index (S>0). Size of bubbles is proportional to the Silhouette index. Internal nodes are drawn with the average expression profile grouped by their



partitions. Leaf node profiles are shown as a heatmap.

```
from ete2 import ClusterTree, TreeStyle, AttrFace, ProfileFace, TextFace
from ete2.treeview.faces import add_face_to_node
# To operate with numbers efficiently
import numpy
PATH = "./"
# Loads tree and array
t = ClusterTree(PATH+"diauxic.nw", PATH+"diauxic.array")
# nodes are linked to the array table
array = t.arraytable
# Calculates some stats on the matrix. Needed to establish the color
# gradients.
matrix_dist = [i for r in xrange(len(array.matrix))\
               for i in array.matrix[r] if numpy.isfinite(i)]
matrix_max = numpy.max(matrix_dist)
matrix_min = numpy.min(matrix_dist)
matrix_avg = matrix_min+((matrix_max-matrix_min)/2)
# Creates a profile face that will represent node's profile as a
# heatmap
profileFace = ProfileFace(matrix_max, matrix_min, matrix_avg, \
                                         200, 14, "heatmap")
cbarsFace = ProfileFace(matrix max,matrix min,matrix avg,200,70,"cbars")
nameFace = AttrFace("name", fsize=8)
# Creates my own layout function that uses previous faces
def mylayout (node):
    # If node is a leaf
    if node.is_leaf():
        # And a line profile
        add_face_to_node(profileFace, node, 0, aligned=True)
        node.img_style["size"]=0
        add_face_to_node(nameFace, node, 1, aligned=True)
    # If node is internal
    else:
        # If silhouette is good, creates a green bubble
        if node.silhouette>0:
            validationFace = TextFace("Silh=%0.2f" %node.silhouette,
                                      "Verdana", 10, "#056600")
            node.img_style["fgcolor"]="#056600"
        # Otherwise, use red bubbles
        else:
            validationFace = TextFace("Silh=%0.2f" %node.silhouette,
                                      "Verdana", 10, "#940000")
            node.img style["fgcolor"]="#940000"
        # Sets node size proportional to the silhouette value.
        node.img_style["shape"]="sphere"
        if node.silhouette<=1 and node.silhouette>=-1:
            node.img_style["size"] = 15+int((abs(node.silhouette)*10)**2)
        # If node is very internal, draw also a bar diagram
        # with the average expression of the partition
```

# 2.5 Phylogenetic XML standards

New in version 2.1.

From version 2.1, ETE provides support for NeXML and PhyloXML phylogenetic XML standards, both reading and writing. These standards allow to encode complex phylogenetic data, and therefore they are not limited to trees. Although ETE is mainly focused on allowing transparent interaction with trees, it also provides basic I/O methods to data of different type.

Essentially, NexML and PhyloXML files are intended to encode collections of phylogenetic data. Such information can be converted to a collection Python objects sorted in a hierarchical way. A specific Python class exists for every element encoded documented by the NeXML and PhyloXML formats. This is possible thanks to the the general purpose Python drivers available for both formats (http://etetoolkit.org/phyloxml-and-nexml-python-parsers). ETE will use such drivers to access XML data, and it will also convert tree data into PhyloTree objects. In practice, conversions will occur transparently. NeXML and PhyloXML files are loaded using their specific root classes, provided by the main ETE module, and all the information will become available as a collection of Python objects internally sorted according to the original XML hierarchy.

New in version 2.1.

#### 2.5.1 **NeXML**

NeXML(http://nexml.org) is an exchange standard for representing phyloinformatic data inspired by the commonly used NEXUS format, but more robust and easier to process.

### **Reading NeXML projects**

Nexml projects are handled through the Nexml base class. To load a NexML file, the Nexml.build\_from\_file() method can be used.

```
from ete2 import Nexml

nexml_prj = Nexml()
nexml_prj.build_from_file("/path/to/nexml_example.xml")
```

Note that the ETE parser will read the provided XML file and convert all elements into python instances, which will be hierarchically connected to the Nexml root instance.

Every NeXML XML element has its own python class. Content and attributes can be handled through the "set\_" and "get\_" methods existing in all objects. Nexml classes can be imported from the ete2.nexml module.

```
from ete2 import Nexml, nexml
nexml_prj = Nexml()
nexml_meta = nexml.LiteralMeta(datatype="double", property="branch_support", content=1.0
nexml_prj.add_meta(nexml_meta)
nexml_prj.export()

# Will produce:
#
# <Nexml xmlns:xsi="http://www.w3.org/2001/XMLSchema-instance" xsi:type="Nexml">
# <meta datatype="double" content="1.0" property="branch_support" xmlns:xsi="http://www.w3.org/2001/xmlschema-instance" xmlns:xsi="h
```

#### **NeXML** trees

NeXML tree elements are automatically converted into PhyloTree instances, containing all ETE functionality (traversing, drawing, etc) plus normal NeXML attributes.

In the Nexml standard, trees are represented as plain lists of nodes and edges. ETE will convert such lists into tree topologies, in which every node will contain a <code>nexml\_node</code> and <code>nexml\_edge</code> attribute. In addition, each tree node will have a <code>nexml\_tree</code> attribute (i.e. <code>NEXML->FloatTree</code>), which can be used to set the nexml properties of the subtree represented by each node. Note also that <code>node.dist</code> and <code>node.name</code> features will be linked to <code>node.nexml\_edge.length</code> and <code>node.nexml\_node.label</code>, respectively.

```
from ete2 import Nexml
# Create an empty Nexml project
nexml_project = Nexml()
# Load content from NeXML file
nexml_project.build_from_file("trees.xml")
# All XML elements are within the project instance.
# exist in each element to access their attributes.
print "Loaded Taxa:"
for taxa in nexml_project.get_otus():
             for otu in taxa.get_otu():
                         print "OTU:", otu.id
# Extracts all the collection of trees in the project
tree_collections = nexml_project.get_trees()
# Select the first collection
collection_1 = tree_collections[0]
# print the topology of every tree
for tree in collection_1.get_tree():
             # trees contain all the nexml information in their "nexml_node",
             # "nexml_tree", and "nexml_edge" attributes.
            print "Tree id", tree.nexml_tree.id
            print tree
             for node in tree.traverse():
                         print "node", node.nexml_node.id, "is associated with", node.nexml_node.otu, "One of the other other of the other othe
# Output:
```

# ========

```
# Loaded Taxa:
# OTU: t1
# OTU: t2
# OTU: t3
# OTU: t4
# OTU: t5
# Tree id tree1
                 \ -n6(n6)
                 -n9(n9)
      -n2(n2)
# node n1 is associated with None OTU
# node n3 is associated with None OTU
# node n2 is associated with t1 OTU
# node n4 is associated with None OTU
# node n7 is associated with None OTU
# node n5 is associated with t3 OTU
# node n6 is associated with t2 OTU
# node n8 is associated with t5 OTU
# node n9 is associated with t4 OTU
# Tree id tree2
                 /-tree2n5(n5)
                 \-tree2n6(n6)
                 /-tree2n8(n8)
                 \-tree2n9(n9)
      \t-tree2n2(n2)
# node tree2n1 is associated with None OTU
# node tree2n3 is associated with None OTU
# node tree2n2 is associated with t1 OTU
# node tree2n4 is associated with None OTU
# node tree2n7 is associated with None OTU
# node tree2n5 is associated with t3 OTU
# node tree2n6 is associated with t2 OTU
# node tree2n8 is associated with t5 OTU
# node tree2n9 is associated with t4 OTU
[Download tolweb.xml example] | [Download script]
Node meta information is also available:
from ete2 import Nexml
# Creates and empty NeXML project
p = Nexml()
# Fill it with the tolweb example
p.build_from_file("tolweb.xml")
```

```
# extract the first collection of trees
tree_collection = p.trees[0]
# and all the tree instances in it
trees = tree collection.tree
# For each loaded tree, prints its structure and some of its
# meta-properties
for t in trees:
   print t
   print
   print "Leaf node meta information:\n"
   print
    for meta in t.children[0].nexml_node.meta:
        print meta.property, ":", (meta.content)
# Output
# -----
# ---- /-node3(Eurysphindus)
# Leaf node meta information:
# dc:description :
# tbe:AUTHORITY : Leconte
# tbe:AUTHDATE : 1878
# tba:ANCESTORWITHPAGE : 117851
# tba:CHILDCOUNT : 0
# tba:COMBINATION_DATE : null
# tba:CONFIDENCE : 0
# tba:EXTINCT : 0
# tba:HASPAGE : 1
# tba:ID : 117855
# tba:INCOMPLETESUBGROUPS : 0
# tba:IS_NEW_COMBINATION : 0
# tba:ITALICIZENAME : 1
# tba:LEAF : 0
# tba:PHYLESIS : 0
# tba:SHOWAUTHORITY : 0
# tba:SHOWAUTHORITYCONTAINING : 1
[Download tolweb.xml example] | [Download script]
```

**Creating Nexml project from scratch** Nexml base class can also be used to create projects from scratch in a programmatic way. Using the collection of NeXML classes provided by the:mod:*ete2.nexml* module, you can populate an empty project and export it as XML.

```
import sys
# Note that we import the nexml module rather than the root Nexml
# class. This module contains a python object for each of the
# nexml elements declared in its XML schema.
from ete2 import nexml

# Create an empty Nexml project
nexml_project = nexml.Nexml()
tree_collection = nexml.Trees()
```

```
# NexmlTree is a special PhyloTree instance that is prepared to be
# added to NeXML projects. So lets populate a random tree
nexml_tree = nexml.NexmlTree()
# Random tree with 10 leaves
nexml_tree.populate(10, random_branches=True)
# We add the tree to the collection
tree collection.add tree (nexml tree)
# Create another tree from a newick string
nexml_tree2 = nexml.NexmlTree("((hello, nexml):1.51, project):0.6;")
tree_collection.add_tree(nexml_tree2)
# Tree can be handled as normal ETE objects
nexml_tree2.show()
# Add the collection of trees to the NexML project object
nexml_project.add_trees(tree_collection)
# Now we can export the project containing our two trees
nexml_project.export()
[Download script]
```

**Writing NeXML objects** Every NexML object has its own export () method. By calling it, you can obtain the XML representation of any instance contained in the Nexml project structure. Usually, all you will need is to export the whole project, but individual elements can be exported.

```
import sys
from ete2 import Nexml
# Create an empty Nexml project
nexml_project = Nexml()

# Upload content from file
nexml_project.build_from_file("nexml_example.xml")

# Extract first collection of trees
tree_collection = nexml.get_trees()[0]

# And export it
tree_collection.export(output=sys.stdout, level=0)
```

#### **NeXML** tree manipulation and visualization

NeXML trees contain all ETE PhyloTree functionality: orthology prediction, topology manipulation and traversing methods, visualization, etc.

For instance, tree changes performed through the visualization GUI are kept in the NeXML format.

```
from ete2 import nexml
nexml_tree = nexml.NexMLTree("((hello, nexml):1.51, project):0.6;")
tree_collection.add_tree(nexml_tree)
nexml_tree.show()
```

New in version 2.1.

# 2.5.2 PhyloXML

PhyloXML (http://www.phyloxml.org/) is a novel standard used to encode phylogenetic information. In particular, phyloXML is designed to describe phylogenetic trees (or networks) and associated data, such as taxonomic information, gene names and identifiers, branch lengths, support values, and gene duplication and speciation events.

### Loading PhyloXML projects from files

ETE provides full support for phyloXML projects through the Phyloxml object. Phylogenies are integrated as ETE's tree data structures as PhyloxmlTree instances, while the rest of features are represented as simple classes (ete2.phyloxml) providing basic reading and writing operations.

[Download script] [Download example]

Each tree node contains two phyloxml elements, phyloxml\_clade and phyloxml\_phylogeny. The first attribute contains clade information referred to the node, while phyloxml\_phylogeny contains general data about the subtree defined by each node. This way, you can split, or copy any part of a tree and it will be exported as a separate phyloxml phylogeny instance.

Note that node.dist, node.support and node.name features are linked to node.phyloxml\_clade.branch\_length, node.phyloxml\_clade.confidence and node.phyloxml\_clade.name, respectively.

### Creating PhyloXML projects from scratch

In order to create new PhyloXML projects, a set of classes is available in the ete2.phyloxml module.

```
from ete2 import Phyloxml, phyloxml
import random
project = Phyloxml()

# Creates a random tree
phylo = phyloxml.PhyloxmlTree()
phylo.populate(5, random_branches=True)
phylo.phyloxml_phylogeny.set_name("test_tree")
# Add the tree to the phyloxml project
project.add_phylogeny(phylo)
```

```
print project.get_phylogeny()[0]
           /-iajom
#
#
            \-wiszh
           /-xryqw
# Trees can be operated as normal ETE trees
phylo.show()
# Export the project as phyloXML format
project.export()
# <phy:Phyloxml xmlns:phy="http://www.phyloxml.org/1.10/phyloxml.xsd">
#
      <phy:phylogeny>
          <phy:name>test_tree</phy:name>
          <phy:clade>
              <phy:name>NoName</phy:name>
              <phy:branch_length>0.000000e+00</phy:branch_length>
              <phy:confidence type="branch_support">1.0</phy:confidence>
              <phy:clade>
                   <phy:name>NoName</phy:name>
                   <phy:branch_length>1.665083e-01</phy:branch_length>
                   <phy:confidence type="branch_support">0.938507980435</phy:confidence>
                   <phy:clade>
                       <phy:name>NoName</phy:name>
                       <phy:branch_length>1.366655e-01</phy:branch_length>
                       <phy:confidence type="branch_support">0.791888248212</phy:confidence</pre>
                       <phy:clade>
                           <phy:name>ojnfg</phy:name>
                           <phy:branch_length>2.194209e-01</phy:branch_length>
                           <phy:confidence type="branch_support">0.304705977822</phy:confidence</pre>
                       </phy:clade>
                       <phy:clade>
                           <phy:name>qrfnz</phy:name>
                           <phy:branch_length>5.235437e-02</phy:branch_length>
                           <phy:confidence type="branch_support">0.508533765418</phy:confidence</pre>
                       </phy:clade>
                   </phy:clade>
                   <phy:clade>
                       <phy:name>shngq</phy:name>
                       <phy:branch_length>9.740958e-01</phy:branch_length>
                       <phy:confidence type="branch_support">0.642187390965</phy:confidence</pre>
                   </phy:clade>
              </phy:clade>
               <phy:clade>
                   <phy:name>NoName</phy:name>
                   <phy:branch_length>3.806412e-01</phy:branch_length>
#
                   <phy:confidence type="branch_support">0.383619811911</phy:confidence>
                   <phy:clade>
                       <phy:name>vfmnk</phy:name>
```

**Note:** NeXML and PhyloXML python parsers are possible thanks to Dave Kulhman and his work on the generateDS.py application. Thanks Dave! ;-)

New in version 2.1.

### 2.6 Interactive web tree visualization

Starting at version 2.1, ETE provides a module to interactively display trees within web pages. This task is not straightforward, but ETE tries to simplify it by providing a basic <code>WebTreeApplication</code> class that can be imported in your python web applications.

WebTreeApplication implements a transparent connector between ETE's functionality and web application. For this, a pre-built WSGI application is provided.

Through this application, you will be able to create custom web implementations to visualize and manipulate trees interactively. Some examples can be found at the PhylomeDB tree browser or in the ETE's online treeviewer.

### 2.6.1 NO X system available?

Alternatively, a virtual X system such as XVFB has been reported to work in servers without a proper X backend. Just install XVFB and preface your ETE commands with xvfb-run.

```
xvfb-run python MyETEscript.py
```

### 2.6.2 Installing a X server

All modern linux desktop installations include a graphical interface (called X server). However web servers (in which the ETE plugin is expected to run) may not count with a X server.

#### 2.6.3 Servers

In order to render tree images with ETE, you will need to install, at least, a basic X server. Note that the X server does not require a desktop interface, such as Gnome or KDE.

In Ubuntu, for instance, a basic X server called xdm can be installed as follows:

```
apt-get install xserver-xorg xdm xfonts-base xfonts-100dpi
xfonts-75dpi
```

Once the X server is installed, you will need to configure it to accept connections from the web-server.

In our example, edit the /etc/X11/xdm/xdm-config file and set following values:

```
DisplayManager*authorize: false!
DisplayManager*authComplain: false
```

Do not forget to restart your xdm server.

```
/etc/init.d/xdm restart
```

### 2.6.4 Desktops

If you plan to use web tree application in a linux desktop computer, then the X server is already installed. You will only need to give permissions to the web-server (i.e. apache) to connect your display. Usually, as simple as running the following command in a terminal:

```
xhost +
```

# 2.6.5 Configuring the web sever

You will need to add support for WSGI application to your web server. In the following steps, an Apache2 web server will be assumed.

- Install and activate the modwsgi module in Apache.
- Configure your site to support WSGI.

Configuration will depend a lot on your specific system, but this is an example configuration file for the default site of your Apache server (usually at /ete/apache2/sites-available/default):

```
<VirtualHost *:80>
ServerAdmin webmaster@localhost
DocumentRoot /var/www
<Directory />
       Options +FollowSymLinks
       AllowOverride None
</Directory>
ErrorLog /var/log/apache2/error.log
# Possible values include: debug, info, notice, warn, error, crit,
# alert, emerg.
LogLevel warn
CustomLog /var/log/apache2/access.log combined
# WSGI SPECIFIC CONFIG
WSGIDaemonProcess eteApp user=www-data group=www-data processes=1 threads=1
WSGIProcessGroup eteApp
```

**Note:** /var/www/webplugin/wsgi/ is the folder in which python web application will be located. Make sure that the apache WSGI config enables this folder to run wsgi-scripts.

**Warning:** Important notes:

/var/www/webplugin/ is assumed to be the directory in which your application will run. /var/www/webplugin/tmp/ should be writable by the web-server (i.e. chmod 777)

### 2.6.6 Implementation of WebTreeApplications

ETE's WebTreeApplication uses WSGI in the backend, and a several javascript files in the frontend. Basic files are included as an example in the ETE installation package examples/webplugin. The whole example folder is necessary, and it contains a commented copy of a web-tree implementation examples/webplugin/wsgi/webplugin\_example.py.

# 2.7 Testing Evolutionary Hypothesis

#### **Contents**

- Testing Evolutionary Hypothesis
  - Extension Requirements:
  - Overview
    - \* The working directory
  - Descriptive analysis
    - \* Branch model
    - \* Site model
  - Hypothesis Testing
    - \* Test on sites
    - \* Test on branches
    - \* Test on branch-sites
  - Utilities
    - \* Load precomputed evolutionary model
  - References

### 2.7.1 Extension Requirements:

You should have codeml and slr in your path:

- CodeML, you can download it from http://abacus.gene.ucl.ac.uk/software/paml.html
- SLR, from here: http://www.ebi.ac.uk/goldman-srv/SLR/

Download, compile and install both of the programs, in order to be able to run the examples.

This ETE extension is written on top of these two external programs, so it is strongly recommended to read their corresponding documentation.

#### 2.7.2 Overview

An other aspect in the study of evolutionary history, is the analysis of selective pressures accounting for the conservation or degeneration of **protein coding genes**.

The EvolTree class is an extension of the class PhyloTree that implements mainly bindings to the PAML package [yang2007] but also to the SLR program [massingham2005].

Evolutionary variables that are used to summary selective pressures are, of course the branch-length (bL) already available in PhyloTree, but also the rate of non-synonymous mutations (dN), the rate of synonymous mutations (dS) and finally the  $\omega$  ratio:

$$\omega = \frac{dN}{dS} \tag{2.1}$$

#### The working directory

EvolTree works mainly as PhyloTree, thus it needs a tree and an alignment. However as you are going to run external programs over it, **a working directory needs to be defined**. By default tree.workdiris "/tmp/ete2-codeml/", but it is recommended to change it to a more useful path.

Jobs will be stored in the workdir, and you will be able to load precomputed evolutionary models from there.

## 2.7.3 Descriptive analysis

In order to identify the evolutionary trends in a phylogenetic tree, one can either:

- conduct an analysis over branches and compute the value of  $\omega$  in each of them.
- look at the selective pressures along the alignment.

#### **Branch model**

As for PhyloTree, we first load the tree and alignment (and you working directory, if you want to save a copy of your jobs):

```
from ete2 import EvolTree

tree = EvolTree("((Hylobates_lar, (Gorilla_gorilla, Pan_troglodytes)), Papio_cynocephalus);
```

```
tree.link_to_alignment ('''>Hylobates_lar
ATGGCCAGGTACAGATGCTGCCGCAGCCAGAGCCGGAGCAGATGTTACCGCCAGAGCCGGAGCAGATGTTACCGCCAGAGCCAGAGCCGAGCCAGAGCCAGAGCCGAGCCAGAGCCGAGCCAGAGCCGAGCCAGAGCCGAGCCAGAGCCGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCCAGAGCAGAGCCAGAGCAGAGCCAGAGCCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAG
```

Once loaded we are able to compute selective pressure among the tree according to an evolutionary model. In this case, we will use free-ratio model:

```
tree.run_model ('fb.example')
```

EvolNode.run\_model() allows to run different evolutionary models (follow this link EvolNode.run\_model() to see the full list of models available). By convention, the name of the model called is the first word, the rest of the string, after the dot, corresponds to its identifier in order to differentiate different runs of one model. We can thus either run "fb" model and "fb.something" with the same result, but they will be stored under different names instead of being overwritten.

Once computation is done, we can retrieve the evol.Model object from the tree, with the  $EvolNode.get_evol_model()$  function.

```
my_first_fb_model = tree.get_evol_model('fb.example')
print my_first_fb_model
```

#### This last print statement would ouptut:

```
Evolutionary Model fb.example:
     log likelihood
                   : -521.421323
     number of parameters : 13
     sites inference : None
     sites classes
                      : None
     branches
                            , node_ids: 5  , name: NoName
     mark: #0 , omega: None
     mark: #1 , omega: 0.0001
                            , node_ids: 6 , name: NoName
     , node_ids: 7 , name: NoName
                            , node_ids: 1
                                         , name: Gorilla_gorilla
     mark: #5 , omega: 0.1049
     mark: #6 , omega: 0.5334
                            , node_ids: 3
                                         , name: Pan_troglodytes
```

By default, the free-branch model is run with this parameters:

```
clock = 0
    CodonFreq = 2
        getSE = 0
         icode = 0
    fix_kappa = 0
        kappa = 2
#
       Malpha = 0
#
       method = 0
#
        Mgene = 0
#
        model = 1
        ncatG = 8
        noisy = 0
#
      NSsites = 0
#
    fix\_omega = 0
         omega = 0.7
#
# RateAncestor = 0
     runmode = 0
#
      seqtype = 1
  Small Diff = 1e-6
      verbose = 2
```

However, it is usually recommended to run one model several times with different starting values, and keep only the model with higher likelihood. Here an example, on how to do this.

```
best_model = None
best_lnl = float('-inf')
for starting_omega in [0.2, 0.7, 1.2]:
    tree.run_model('fb.'+str(starting_omega))
    current_model = tree.get_evol_model('fb.'+str(starting_omega))
    print 'current model log likelihood:', current_model.lnL
    if current_model.lnL > best_lnl:
        best_lnl = current_model.lnL
        best_model = current_model
```

Finally in order to have a quick look of the selctive pressures over our phylogeny:

```
tree.show()
```

by default this will the picture obtained:



Node size, and color dependant of the  $\omega$  value. But other displays are available:

```
from ete2.treeview.layouts import evol_clean_layout

tree.show(layout=evol_clean_layout)
```

With here  $\omega$  ratios in red and also in gray the dN and dS values.

#### Site model

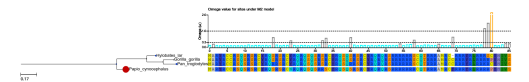
Another way to look at selective pressures, is to compute directly along the alignment, the value of  $\omega$  for a whole column (putting all leaves together). For doing this, we can use for example the model M2 of CodeML or directly use SLR. As before we just have to:

```
tree.run_model('M2')
tree.run_model('SLR.lele')
```

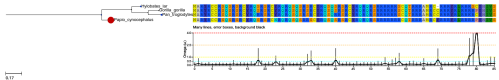
and to display the results:

```
tree.show (histfaces=['M2'])
```

when a site model is computed, an histface is automatically generated. Thus with this call, what we are doing is to draw the default histface corresponding to the model named M2.lala. This is the result:

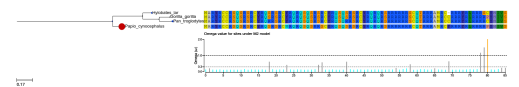


However customizing this face is feasible:



or:

tree.show(histfaces=['M2'])



The col dictionary contains the colors for sites detected to be under positive selection (PS), relaxation (RX), or conserved (CN). However, it is not a good idea to use them now as we do not know if there is indeed positive selection.

To be able to accept M2 results we will have to test this model against a null model.

### 2.7.4 Hypothesis Testing

In order to know if the parameters estimated under a given model a reliable, we have to compare its likelihood to a null model.

Usually, the alternative model is a model that estimates the proportion of sites with  $\omega>1$  and we compare its likelihood with a null model, usually a model that do not (letting  $\omega<=1$ ). This comparison is done through a likelihood ratio test. If the alternative model has the best fit than we are able to accept the possibility of  $\omega>1$ .

To see a non-exhaustive list of famous comparison see the documentation of the function: EvolNode.get\_most\_likely()

#### **Test on sites**

In order to know if some sites are significantly under positive selection, relaxed or conserved we have usually to compare 2 models. However using the model "SLR" we can directly infer positive selection or relaxation through the SLR program [massingham2005].

The most usual comparison, and perhaps the most robust, is the comparison of models M2 and M1.

```
tree.run_model ('M1')
tree.run_model ('M2')
pval = tree.get_most_likely ('M2','M1')
model2 = tree.get evol model('M2')
print model2
# Evolutionary Model M2:
                            : -517.824749
        log likelihood
        number of parameters : 11
        sites inference : BEB, NEB
        sites classes
          proportions: p0=0.98794 p1=0.00000 p2=0.01206
          w: w0=0.09887 w1=1.00000 w2=178.86153
           mark: #0, omega: 2.25526440485 , nodes paml_ids: 6 4 2 7 1 3
if pval < 0.05:
 print 'M2 model wins.'
  for s in range(len(model2.sites['BEB']['aa'])):
    if model2.sites['BEB']['p2'][s] > 0.95:
     print 'positively selected site %s at position: %s, with probability: %s' % (model
else:
 print 'M1 model is not rejected'
# M2 model wins.
# positively selected site P at position: 81, with probability: 0.96293
```

Each sites model, contains a dictionary 'model.sites' in which are stored the results of CodeML. These are displayed through the histface, but they are still accessible by digging a bit. 'site' dictionary usually contains the result of the NEB and BEB analysis (prefer the BEB when available). For each of BEB and NEB, the probability of belonging from one category of site is summarized by 'p0', 'p1' and 'p2' in the case of M2 model that have only 3 class of sites (p0, the probability of belonging to the first class of sites with  $\omega < 1$ ; p1, the probability of belonging to the second class of sites with  $\omega = 1$ ; p2, the probability of belonging to the third class of sites with  $\omega > 1$ ). For details on the models and outputs, it is recommended to read PAML documentation.

#### **Test on branches**

CodeML allows to test models that infer different selective pressures on a set of given branches. To do it, some branches of the tree are marked (usually with strings like this: '#1'). This is how to do it with ETE:

```
marks = ['2', '3', '4']
# mark a group of branches
tree.mark_tree (marks, ['#1', '#1', '#1'])
print tree.write ()
# ((Hylobates_lar, (Gorilla_gorilla #1, Pan_troglodytes #1) #1), Papio_cynocephalus);
```

By doing this a branch model will compute different  $\omega$  values in mark branches (usually called foreground  $\omega_{frq}$ ) and in the rest of the tree (usually called background  $\omega_{bkq}$ ).

#### Two kinds of branch models are usually used:

- the free-branch model: were  $\omega_{frq}$  and  $\omega_{bkq}$  are free
- the neutral-branch model: were  $\omega_{frg}$  is fixed to one.
- the M0 model: were all branches evolve at the same rate.

The comparison between free-branch and M0, will tell us if foreground branches have an  $\omega$  significantly different from the rest of the tree.

And the comparison between free-branch and neutral-branch models will tell us if  $\omega_{frg}$  is significantly higher than 1.

```
tree.run_model ('b_free')
tree.run_model ('b_neut')
tree.run_model ('M0')

b_free = tree.get_evol_model('b_free')

print b_free

# Evolutionary Model b_free.234:
# log likelihood : -525.690213
# number of parameters : 9
# sites inference : None
# sites classes : None
# branches
# mark: #0, omega: 0.157280451975 , nodes paml_ids: 6 4 2
```

```
# mark: #1, omega: 0.118462858241 , nodes paml_ids: 7 1 3

if tree.get_most_likely ('b_free', 'M0') < 0.05:
    # branch models have a branches dictionary were keys corresponds to paml_id of nodes
    # select one of the marked branches
    frg_node = tree.search_nodes(_nid=2)[0]
    frg_pamlid = frg_node.paml_id
    w_frg = bfree.branches[frg_pamlid]['w']
    # select one of the unmarked branches
    bkg_node = tree.search_nodes(_nid=1)[0]
    bkg_pamlid = bkg_node.paml_id
    w_bkg = bfree.branches[bkg_pamlid]['w']
    print 'foreground branches evolving at omega value of %s significantly different from
if tree.get_most_likely ('b_free', 'b_neut') < 0.05:
    print 'foreground branches are significantly different from 1.'</pre>
```

for the given example, background and foreground are not different...

#### Test on branch-sites

An other popular test is the branch-site test contrasting model A and A1. This test allows to detect genes with some sites under positive selection.

this example shows how to run it over all branches in the tree:

```
# also needs the MO model as null model
tree.run_model ('M0')
for leaf in tree:
    leaf.node id
   print '\n----\nNow working with leaf ' + leaf.name
    tree.mark_tree ([leaf.node_id], marks=['#1'])
   print tree.write()
    # to organize a bit, we name model with the name of the marked node
    # any character after the dot, in model name, is not taken into account
    # for computation. (have a look in /tmp/ete2.../bsA.. directory)
    print 'running model bsA and bsA1'
    tree.run_model ('bsA.'+ leaf.name)
    tree.run_model ('bsA1.' + leaf.name)
   print 'p-value of positive selection for sites on this branch is: '
   ps = tree.get_most_likely ('bsA.' + leaf.name, 'bsA1.'+ leaf.name)
    rx = tree.get_most_likely ('bsA1.'+ leaf.name, 'M0')
    print str (ps)
   print 'p-value of relaxation for sites on this branch is: '
   print str (rx)
    if ps<0.05 and float (bsA.wfrg2a)>1:
       print 'we have positive selection on sites on this branch'
    elif rx<0.05 and ps>=0.05:
       print 'we have relaxation on sites on this branch'
    else:
        print 'no signal detected on this branch, best fit for MO'
    print '\nclean tree, remove marks'
    tree.mark_tree (map (lambda x: x.node_id, tree.get_descendants()),
                    marks=[''] * len (tree.get_descendants()), verbose=True)
```

```
print tree.get_evol_model('bsA.Hylobates_lar')
  Evolutionary Model bsA.Hylobates_lar:
        log likelihood : -521.203318
#
        number of parameters : 11
#
         sites inference : BEB, NEB
         sites classes
         foreground w: f0=0.06452 f1=1.00000 f2=3.15727 f3=3.15727
        proportions: p0=0.00415 p1=0.00023 p2=0.94413 p3=0.05150
        background w: b0=0.06452 b1=1.00000 b2=0.06452 b3=1.00000
        branches
                                 , node_ids: 5 , name: NoName
        mark: #0 , omega: None
                                  , node_ids: 6 , name: NoName
         mark: #0 , omega: None
                                  , node_ids: 4 , name: Papio_cynocephalus
         mark: #0 , omega: None
        mark: #1 , omega: None
                                   , node_ids: 2 , name: Hylobates_lar
        mark: #0 , omega: None
                                   , node_ids: 7 , name: NoName
        mark: #0 , omega: None
                                   , node_ids: 1 , name: Gorilla_gorilla
        mark: #0 , omega: None
                                   , node_ids: 3 , name: Pan_troglodytes
```

#### 2.7.5 Utilities

# Load precomputed evolutionary model

When an evolutionary model is computed, the output is stored in *tree.workdir* and can be load afterwards. Inside *tree.workdir*, a new directory is created for each model you compute (if each model has a different name), thus to load one model:

```
from ete2 import EvolTree

tree = EvolTree("((Hylobates_lar, (Gorilla_gorilla, Pan_troglodytes)), Papio_cynocephalus);

tree.link_to_alignment ('''>Hylobates_lar

ATGGCCAGGTACAGATGCTGCCGCAGCCAGAGCCGGAGCAGATGTTACCGCCAGAGCCGGAGCAGATGTTACCGCCAGAGGCAAAGCC
>Papio_cynocephalus

ATGGCCAGGTACAGATGCTGCCGCAGCCAGAGCCGAAGCAGATGCTATCGCCAGAGCCGGAGCAGATGTAACCGCCAGAGACAGAGCC
>Gorilla_gorilla

ATGGCCAGGTACAGATGCTGTCGCAGCCAGAGCCGCAGCAGATGTTACCGGCAGAGCCGGAGCAGGTGTTACCGGCAGAGACAAAGCC
>Pan_troglodytes

ATGGCCAGGTACAGATGCTGTCGCAGCCAGAGCCGGAGCAGATGTTACCGGCAGAGACGAGGTGTTACCGGCAAAGGCAAAGCC

**TGGCCAGGTACAGATGCTGTCGCAGCCAGAGCCGGAGCAGATGTTACCGGCAGAGACGAGGTGTTACCGGCAAAGGCAAAGCCC

**TGGCCAGGTACAGATGCTGTCGCAGCCAGAGCCGGAGCAGATGTTACCGGCAGAGACGAGGTGTTACCGGCAAAGGCAAAGCCC

**TGGCCAGGTACAGATGCTGTCGCAGCCAGAGCCGGAGCAGATGTTACCGGCAGAGACGAGGTGTTACCGGCAAAGGCAAAGCCC

**TGGCCAGGTACAGATGCTGTCGCAGCCAGAGCCGGAGCAGATGTTACCGGCAGAGACGAGGTGTTACCGGCAAAGGCAAAGCCC

**TGGCCAGGTACAGATGCTGTCGCAGCCAGAGCCGGAGCAGATGTTACCGGCAGAGACGAGGTGTTACCGGCAAAGGCAAAGCCC

**TGCCAGGTACAGATGCTGTCGCAGCCAGAGCCGGAGCAGATGTTACCGGCAGAGACGAGGCGAGAGGTGTTACCGGCAAAGGCAAAGCCC

**TGCCAGGTACAGATGCTGTCGCAGCCAGAGCCGGAGCAGATGTTACCGGCAGAGACGAGGTGTTACCGGCAAAGGCAAAGCCC

**TGCCAGGTACAGATGCTGTCGCAGCCAGAGCCGGAGCCGGAGCAGATGTTACCGGCAGAGACGGAGCAGGTGTTACCGGCAAAGGCAAAGCCC

**TGCCAGGTACAGATGCTGTCGCAGCCAGAGCCGGAGCCGGAGCAGATGTTACCGGCAGAGACGAGGCGAGAGCGAAAGGCAAAGCCC

**TGCCAGGTACAGATGCTGCCGCAGAGCCGAGAGCCGGAGCAGATGTTACCGGCAGAGACGAGGCGAAAGGCAAAGCCC

**TGCCAGGTACAGATGCTGCCGCAGCCAGAGCCGGAGCCGAGAGGCAGATGTTACCGGCAGAGCCGAAAGCCAAAGCCC

**TGCCAGGTACAGATGCTGCCGCAGAGCCGAGAGCCGAGAGCAGATGTTACCGGCAGAGAGCAGAGGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGAGCAGAGAGCAGAGCAGAGCAGAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGCAGAGAGAGCAGAGCAGAG
```

Note: EvolNode.link\_to\_evol\_model() is also able to load directly evol.Model objects.

Indeed all output files generated do not need to be kept, only a pickle or cPickle of the model for example.

Thus we can save a tree with all Models computed, after having run some models:

```
# assume we already have run some models:
for model in tree._models:
    print tree.get_evol_model(model)

# Evolutionary Model M2.example:
    log likelihood : -517.824749
    number of parameters : 11
# sites inference : BEB, NEB
# sites classes :
```

```
#
           proportions: p0=0.98794 p1=0.00000 p2=0.01206
           w : w0=0.09887 w1=1.00000 w2=178.86192
         branches
           mark: #0, omega: 2.25526864047 , nodes paml_ids: 6 4 2 7 1 3
#
 Evolutionary Model M1.example:
#
         log likelihood : -521.541845
#
        number of parameters : 9
#
        sites inference : NEB
#
         sites classes
           proportions : p0=0.94075 p1=0.05925
         w : w0=0.07025 w1=1.00000 branches :
           mark: #0, omega: 0.125334579074 , nodes paml_ids: 6 4 2 7 1 3
# we save the whole tree into my_tree.pik
from cPickle import dump
out = open('my_tree.pik', 'w')
dump (tree, out)
out.close()
and load them:
from cPickle import load
out = open('my_tree.pik')
tree = load(out)
out.close()
# and here the same result
for model in tree._models:
   print tree.get_evol_model(model)
# Evolutionary Model M2.example:
        log likelihood : -517.824749
#
        number of parameters : 11
         sites inference : BEB, NEB
#
         sites classes
          proportions: p0=0.98794 p1=0.00000 p2=0.01206
         w : w0=0.09887 w1=1.00000 w2=178.86192 branches :
#
#
           mark: #0, omega: 2.25526864047 , nodes paml_ids: 6 4 2 7 1 3
#
 Evolutionary Model M1.example:
        log likelihood : -521.541845
#
         number of parameters : 9
#
        sites inference : NEB
         sites classes
          proportions : p0=0.94075 p1=0.05925
           w : w0=0.07025 w1=1.00000
         branches
            mark: #0, omega: 0.125334579074 , nodes paml_ids: 6 4 2 7 1 3
```

this also can be done for one model alone:

```
from cPickle import dump, load

m2_to_save = tree.get_evol_model('M2.example')

out = open ('m2.pik', 'w')
dump(m2_to_save, out)
out.close()

# and then load it
out = open ('m2.pik')
m2_to_save = dump(out)
out.close()

tree.link_to_evol_model(m2_to_save)
```

### 2.7.6 References

New in version 2.3.

# 2.8 Dealing with the NCBI Taxonomy database

ETE's *ncbi\_taxonomy* module provides utilities to efficiently query a local copy of the NCBI Taxonomy database. The class NCBITaxa offers methods to convert from taxid to names (and vice versa), to fetch pruned topologies connecting a given set of species, or to download rank, names and lineage track information.

It is also fully integrated with PhyloTree instances through the PhyloNode.annotate ncbi taxa() method.

# 2.8.1 Setting up a local copy of the NCBI taxonomy database

The first time you attempt to use NCBITaxa, ETE will detect that your local database is empty and it will attempt to download the latest NCBI taxonomy database (~300MB) and will store a parsed version of it in your home directory: ~/.etetoolkit/taxa.sqlite. All future imports of NCBITaxa will detect the local database and will skip this step.

```
from ete2 import NCBITaxa
ncbi = NCBITaxa()
```

# 2.8.2 Upgrading the local database

Use the method :NCBITaxa: 'update\_taxonomy\_database' to download and parse the latest database from the NCBI ftp site. Your current local database will be overwritten.

```
from ete2 import NCBITaxa
ncbi = NCBITaxa()
ncbi.update_taxonomy_database()
```

# 2.8.3 Getting taxid information

you can fetch species names, ranks and linage track information for your taxids using the following methods:

```
NCBITaxa.get_rank()
NCBITaxa.get_lineage()
NCBITaxa.get_taxid_translator()
NCBITaxa.get_name_translator()
NCBITaxa.translate_to_names()
```

The so called get-translator-functions will return a dictionary converting between taxids and species names. Either species or linage names/taxids are accepted as input.

```
from ete2 import NCBITaxa
ncbi = NCBITaxa()
taxid2name = ncbi.get_taxid_translator([9606, 9443])
print taxid2name
# {9443: u'Primates', 9606: u'Homo sapiens'}

name2taxid = ncbi.get_name_translator(["Homo sapiens", "primates"])
print name2taxid
# {'Homo sapiens': 9606, 'primates': 9443}
```

Other functions allow to extract further information using taxid numbers as a query.

```
from ete2 import NCBITaxa
ncbi = NCBITaxa()

print ncbi.get_rank([9606, 9443])
# {9443: u'order', 9606: u'species'}

print ncbi.get_lineage(9606)

# [1, 131567, 2759, 33154, 33208, 6072, 33213, 33511, 7711, 89593, 7742,
# 7776, 117570, 117571, 8287, 1338369, 32523, 32524, 40674, 32525, 9347,
# 1437010, 314146, 9443, 376913, 314293, 9526, 314295, 9604, 207598, 9605,
# 9606]
```

And you can combine combine all at once:

```
from ete2 import NCBITaxa
ncbi = NCBITaxa()

lineage = ncbi.get_lineage(9606)
print lineage

# [1, 131567, 2759, 33154, 33208, 6072, 33213, 33511, 7711, 89593, 7742,
# 7776, 117570, 117571, 8287, 1338369, 32523, 32524, 40674, 32525, 9347,
# 1437010, 314146, 9443, 376913, 314293, 9526, 314295, 9604, 207598, 9605,
# 9606]

names = ncbi.get_taxid_translator(lineage)
print [names[taxid] for taxid in lineage]

# [u'root', u'cellular organisms', u'Eukaryota', u'Opisthokonta', u'Metazoa',
```

```
# u'Eumetazoa', u'Bilateria', u'Deuterostomia', u'Chordata', u'Craniata',
# u'Vertebrata', u'Gnathostomata', u'Teleostomi', u'Euteleostomi',
# u'Sarcopterygii', u'Dipnotetrapodomorpha', u'Tetrapoda', u'Amniota',
# u'Mammalia', u'Theria', u'Eutheria', u'Boreoeutheria', u'Euarchontoglires',
# u'Primates', u'Haplorrhini', u'Simiiformes', u'Catarrhini', u'Hominoidea',
# u'Hominidae', u'Homininae', u'Homo', u'Homo sapiens']
```

# 2.8.4 Getting descendant taxa

Given a taxid or a taxa name from an internal node in the NCBI taxonomy tree, their descendants can be retrieved as follows:

### 2.8.5 Getting NCBI species tree topology

Getting the NCBI taxonomy tree for a given set of species is one of the most useful ways to get all information at once. The method NCBITaxa.get\_topology() allows to query your local NCBI database and extract the smallest tree that connects all your query taxids. It returns a normal ETE tree in which all nodes, internal or leaves, are annotated for lineage, scientific names, ranks, and so on.

If needed, all intermediate nodes connecting the species can also be kept in the tree:

# 2.8.6 Automatic tree annotation using NCBI taxonomy

NCBI taxonomy annotation consists of adding additional information to any internal a leaf node in a give user tree. Only an attribute containing the taxid associated to each node is required for the nodes in the query tree. The annotation process will add the following features to the nodes:

- sci\_name
- taxid
- named\_lineage
- lineage
- rank

Note that, for internal nodes, taxid can be automatically inferred based on their sibling nodes. The easiest way to annotate a tree is to use a PhyloTree instance where the species name attribute is transparently used as the taxid attribute. Note that the :PhyloNode:'annotate\_ncbi\_taxa': function will also return the used name, lineage and rank translators.

Remember that species names in *PhyloTree* instances are automatically extracted from leaf names. The parsing method can be easily adapted to any formatting:

```
from ete2 import PhyloTree

# load the whole leaf name as species taxid
tree = PhyloTree('((9606, 9598), 10090);', sp_naming_function=lambda name: name)
tax2names, tax2lineages, tax2rank = tree.annotate_ncbi_taxa()

# split names by '|' and return the first part as the species taxid
tree = PhyloTree('((9606|protA, 9598|protA), 10090|protB);', sp_naming_function=lambda name;
tax2names, tax2lineages, tax2rank = tree.annotate_ncbi_taxa()

print tree.get_ascii(attributes=["name", "sci_name", "taxid"])

# /-9606|protA, Homo sapiens, 9606
# /, Homininae, 207598
#-, Euarchontoglires, 314146
# /-9598|protA, Pan troglodytes, 9598
#-
```

\-10090|protB, Mus musculus, 10090

Alternatively, you can also use the NCBITaxa.annotate\_tree() function to annotate a custom tree instance.

```
from ete2 import Tree, NCBITaxa
ncbi = NCBITaxa()
tree = Tree("")
ncbi.annotate tree(tree, taxid attr="name")
```

# 2.9 SCRIPTS: orthoXML

```
Contents
SCRIPTS: orthoXML
OrthoXML parser
The etree2orthoxml script
* Usage
* Example: Using custom evolutionary annotation
* Example: Automatic detection of species names
* Example: Tree rooting
```

# 2.9.1 OrthoXML parser

etree2orthoxml is a python script distributed as a part of the ETE toolkit package. It uses an automatic python parser generated on the basis of the OrthoXML schema to convert the evolutionary events in phylogenetic tree topologies into the orthoXML format.

ETE OrthoXML parser is a low level python module that allows to operate with the orthoXML structure using python objects. Every element defined in the orthoXML schema has its akin in the parser module, so a complete orthoXML structure can be generated from scratch within a python script. In other words, low level access to the orthoXML parser allows to create orthoxml documents in a programmatic way.

The following example will create a basic orthoXML document

```
from ete2 import orthoxml
# Creates an empty orthoXML object

oxml = orthoxml.orthoXML()

# Add an ortho group container to the orthoXML document
ortho_groups = orthoxml.groups()
oxml.set_groups(ortho_groups)

# Add an orthology group including two sequences
orthologs = orthoxml.group()
orthologs.add_geneRef(orthoxml.geneRef(1))
orthologs.add_geneRef(orthoxml.geneRef(2))
ortho_groups.add_orthologGroup(orthologs)

oxml_file = open("test_orthoxml.xml", "w")
oxml.export(oxml_file, level=0)
oxml_file.close()

# producing the following output
```

```
#<ortho:orthoXML>
# <ortho:groups>
# <ortho:orthologGroup>
# <ortho:geneRef id="1"/>
# <ortho:geneRef id="2"/>
# </ortho:orthologGroup>
# </ortho:groups>
#</ortho:orthoXML>
```

# 2.9.2 The etree2orthoxml script

etree2orthoxml is a standalone python script that allows to read a phylogenetic tree in newick format and export their evolutionary events (duplication and speciation events) as an orthoXML document. The program is installed along with ETE, so it should be found in your path. Alternatively you can found it in the script folder of the latest ETE package release (http://etetoolkit.org/releases/ete2/).

To work, etree2orthoxml requires only one argument containing the newick representation of a tree or the name of the file that contains it. By default, automatic detection of speciation and duplication events will be carried out using the built-in *species overlap algorithm*, although this behavior can be easily disabled when event information is provided along with the newick tree. In the following sections you will find some use case examples.

Also, consider reading the source code of the script. It is documented and it can be used as a template for more specific applications. Note that etree2orthoxml is a work in progress, so feel free to use the etetoolkit mailing list to report any feedback or improvement to the code.

### **Usage**

```
usage: etree2orthoxml [-h] [--sp_delimiter SPECIES_DELIMITER]
                      [--sp_field SPECIES_FIELD] [--root [ROOT [ROOT ...]]]
                      [--skip_ortholog_detection]
                      [--evoltype_attr EVOLTYPE_ATTR] [--database DATABASE]
                      [--show] [--ascii] [--newick]
                      tree file
etree2orthoxml is a python script that extracts evolutionary events
(speciation and duplication) from a newick tree and exports them as a
OrthoXML file.
positional arguments:
 tree_file
                        A tree file (or text string) in newick format.
optional arguments:
 -h, --help
                        show this help message and exit
  --sp_delimiter SPECIES_DELIMITER
                        When species names are guessed from node names, this
                        argument specifies how to split node name to guess the
                        species code
  --sp_field SPECIES_FIELD
                        When species names are guessed from node names, this
                        \hbox{argument specifies the position of the species name} \\
                        code relative to the name splitting delimiter
  --root [ROOT [ROOT ...]]
                        Roots the tree to the node grouping the list of node
```

```
names provided (space separated). In example: '--root
                      human rat mouse'
--skip_ortholog_detection
                      Skip automatic detection of speciation and duplication
                      events, thus relying in the correct annotation of the
                      provided tree using the extended newick format (i.e.
                      '((A, A)[&&NHX:evoltype=D], B)[&&NHX:evoltype=S];')
--evoltype_attr EVOLTYPE_ATTR
                      When orthology detection is disabled, the attribute
                      name provided here will be expected to exist in all
                      internal nodes and read from the extended newick
                      format
--database DATABASE
                      Database name
--show
                      Show the tree and its evolutionary events before
                      orthoXML export
--ascii
                      Show the tree using ASCII representation and all its
                      evolutionary events before orthoXML export
--newick
                      print the extended newick format for provided tree
                      using ASCII representation and all its evolutionary
                      events before orthoXML export
```

# **Example: Using custom evolutionary annotation**

If all internal nodes in the provided tree are correctly label as duplication or speciation nodes, automatic detection of events can be disabled using the <code>--skip\_ortholog\_detection</code> flag.

Node labeling should be provided using the extended newick format. Duplication nodes should contain the label evoltype set to D, while speciation nodes should be set to evoltype=S. If tag names is different, the option evoltype\_attr can be used as convenient.

In the following example, we force the HUMAN clade to be considered a speciation node.

```
# etree2orthoxml --skip_ortholog_detection '((HUMAN_A, HUMAN_B)[&&NHX:evoltype=S], MOUSI
 <orthoXML>
     <species name="A">
         <database name="">
                 <gene protId="HUMAN_A" id="0"/>
             </genes>
         </database>
     </species>
     <species name="B">
         <database name="">
                 <gene protId="HUMAN_B" id="1"/>
                 <gene protId="MOUSE_B" id="2"/>
             </genes>
         </database>
     </species>
     <groups>
         <orthologGroup>
             <geneRef id="2"/>
             <orthologGroup>
                 <geneRef id="0"/>
                 <geneRef id="1"/>
```

You can avoid tree reformatting when node labels are slightly different by using the evoltype\_attr:

```
# etree2orthoxml --evoltype_attr E --skip_ortholog_detection '((HUMAN_A, HUMAN_B)[&&NHX.
```

However, more complex modifications on raw trees can be easily performed using the core methods of the ETE library, so they match the requirements of the etree2orthoxml script.

```
from ete2 import Tree
# Having the followin tree
t = Tree('((HUMAN_A, HUMAN_B)[&&NHX:speciation=N], MOUSE_B)[&&NHX:speciation=Y];')
# We read the speciation tag from nodes and convert it into a vaild evoltree label
for node in t.traverse():
    if not node.is_leaf():
        etype = "D" if node.speciation == "N" else "S"
        node.add_features(evoltype=etype)

# We the export a newick string that is compatible with etree2orthoxml script
t.write(features=["evoltype"], format_root_node=True)

# converted newick:
# '((HUMAN_A:1, HUMAN_B:1)1:1[&&NHX:evoltype=D], MOUSE_B:1)1:1[&&NHX:evoltype=S];'
```

# **Example: Automatic detection of species names**

As different databases and software may produce slightly different newick tree formats, the script provides several customization options.

In gene family trees, species names are usually encoded as a part of leaf names (i.e. P53\_HUMAN). If such codification follows a simple rule, etree2orthoxml can automatically detect species name and used to populate the relevant sections within the orthoXML document.

For this, the sp\_delimiter and sp\_field arguments can be used. Note how species are correctly detected in the following example:

# **Example: Tree rooting**

When evolutionary events are expected to be automatically inferred from tree topology, outgroup information can be passed to the program to root the tree before performing the detection.

```
# etree2orthoxml --ascii --root FLY_1 FLY_2 --sp_delimiter '_' --sp_field 0 '((HUMAN_A,
```

```
/-FLY 1
         /D, NoName
        \-FLY_2
-S, NoName
                           /-HUMAN A
        /D, NoName
         \S, NoName
                      \-HUMAN_B
                  \-MOUSE B
<orthoXML>
    <species name="FLY">
        <database name="">
            <genes>
                <gene protId="FLY 1" id="0"/>
                <gene protId="FLY_2" id="1"/>
            </genes>
        </database>
    </species>
    <species name="HUMAN">
        <database name="">
            <genes>
                <gene protId="HUMAN_A" id="2"/>
                <qene protId="HUMAN_B" id="3"/>
            </genes>
        </database>
    </species>
    <species name="MOUSE">
        <database name="">
                <gene protId="MOUSE_B" id="4"/>
            </genes>
        </database>
    </species>
```

```
<groups>
        <orthologGroup>
            <paralogGroup>
                <geneRef id="0"/>
                <geneRef id="1"/>
            </paralogGroup>
            <orthologGroup>
                <geneRef id="4"/>
                <paralogGroup>
                    <geneRef id="2"/>
                    <geneRef id="3"/>
                </paralogGroup>
            </orthologGroup>
        </orthologGroup>
    </groups>
</orthoXML>
```

# **ETE's Reference Guide**

Current modules:

# 3.1 Master Tree class

class TreeNode (newick=None, format=0, dist=None, support=None, name=None)

TreeNode (Tree) class is used to store a tree structure. A tree consists of a collection of TreeNode instances connected in a hierarchical way. Trees can be loaded from the New Hampshire Newick format (newick).

#### **Parameters**

- **newick** Path to the file containing the tree or, alternatively, the text string containing the same information.
- **format** (0) subnewick format

FORMAT	DESCRIPTION
0	flexible with support values
1	flexible with internal node names
2	all branches + leaf names + internal supports
3	all branches + all names
4	leaf branches + leaf names
5	internal and leaf branches + leaf names
6	internal branches + leaf names
7	leaf branches + all names
8	all names
9	leaf names
100	topology only

**Returns** a tree node object which represents the base of the tree.

# \*\* Examples: \*\*

```
t1 = Tree() # creates an empty tree
t2 = Tree('(A:1, (B:1, (C:1,D:1):0.5):0.5);')
t3 = Tree('/home/user/myNewickFile.txt')
```

add\_child (child=None, name=None, dist=None, support=None)

Adds a new child to this node. If child node is not suplied as an argument, a new node instance will be created.

#### **Parameters**

- **child** (*None*) the node instance to be added as a child.
- **name** (*None*) the name that will be given to the child.
- **dist** (*None*) the distance from the node to the child.
- **support**' (*None*) the support value of child partition.

#### **Returns** The child node instance

# add\_face (face, column, position='branch-right')

Add a fixed face to the node. This type of faces will be always attached to nodes, independently of the layout function.

#### **Parameters**

- face a Face or inherited instance
- column An integer number starting from 0
- **position** ("branch-right") Posible values are: "branch-right", "branch-top", "branch-bottom", "float", "aligned"

# add\_feature (pr\_name, pr\_value)

Add or update a node's feature.

### add\_features (\*\*features)

Add or update several features.

### add\_sister(sister=None, name=None, dist=None)

Adds a sister to this node. If sister node is not supplied as an argument, a new TreeNode instance will be created and returned.

### check\_monophyly (values, target\_attr, ignore\_missing=False, unrooted=False)

Returns True if a given target attribute is monophyletic under this node for the provided set of values.

If not all values are represented in the current tree structure, a ValueError exception will be raised to warn that strict monophyly could never be reached (this behaviour can be avoided by enabling the *ignore\_missing* flag.

### **Parameters**

- values a set of values for which monophyly is expected.
- **target\_attr** node attribute being used to check monophyly (i.e. species for species trees, names for gene family trees, or any custom feature present in the tree).
- **ignore\_missing** (*False*) Avoid raising an Exception when missing attributes are found.

**Parameters unrooted** (*False*) – If True, tree will be treated as unrooted, thus allowing to find monophyly even when current outgroup is spliting a monophyletic group.

**Returns** the following tuple IsMonophyletic (boolean), clade type ('monophyletic', 'paraphyletic' or 'polyphyletic'), leaves breaking the monophyly (set)

#### children

A list of children nodes

```
compare (ref_tree, use_collateral=False, min_support_source=0.0,
    min_support_ref=0.0, has_duplications=False, expand_polytomies=False, un-
    rooted=False, max_treeko_splits_to_be_artifact=1000, ref_tree_attr='name',
    source_tree_attr='name')
```

compare this tree with another using robinson foulds symmetric difference and number of shared edges. Trees of different sizes and with duplicated items allowed.

returns: a Python dictionary with results

```
convert_to_ultrametric (tree_length=None, strategy='balanced')
```

Converts a tree into ultrametric topology (all leaves must have the same distance to root). Note that, for visual inspection of ultrametric trees, node.img\_style["size"] should be set to 0.

```
copy (method='cpickle')
```

Returns a copy of the current node.

Variables method (cpickle) – Protocol used to copy the node

structure. The following values are accepted:

- •"newick": Tree topology, node names, branch lengths and branch support values will be copied by as represented in the newick string (copy by newick string serialisation).
- •"newick-extended": Tree topology and all node features will be copied based on the extended newick format representation. Only node features will be copied, thus excluding other node attributes. As this method is also based on newick serialisation, features will be converted into text strings when making the copy.
- •"cpickle": The whole node structure and its content is cloned based on cPickle object serialisation (slower, but recommended for full tree copying)
- •"deepcopy": The whole node structure and its content is copied based on the standard "copy" Python functionality (this is the slowest method but it allows to copy complex objects even if attributes point to lambda functions, etc.)

# del\_feature(pr\_name)

Permanently deletes a node's feature.

```
delete (prevent_nondicotomic=True, preserve_branch_length=False)
```

Deletes node from the tree structure. Notice that this method makes 'disappear' the node from the tree structure. This means that children from the deleted node are transferred to the next available parent.

Parameters prevent\_nondicotomic (True) – When True (default), delete

function will be execute recursively to prevent single-child nodes.

```
Parameters preserve_branch_length (False) – If True, branch lengths
```

of the deleted nodes are transferred (summed up) to its parent's branch, thus keeping original distances among nodes.

#### **Example:**

```
/ Croot-|
```

> H.delete() will produce this structure:

### describe()

Prints general information about this node and its connections.

#### detach()

Detachs this node (and all its descendants) from its parent and returns the referent to itself.

Detached node conserves all its structure of descendants, and can be attached to another node through the 'add\_child' function. This mechanism can be seen as a cut and paste.

#### dist

Branch length distance to parent node. Default = 0.0

New in version 2.3.

Given a tree with one or more polytomies, this functions returns the list of all trees (in newick format) resulting from the combination of all possible solutions of the multifurcated nodes.

http://ajmonline.org/2010/darwin.php

# get\_ancestors()

versionadded: 2.2

Returns the list of all ancestor nodes from current node to the current tree root.

 $\verb"get_ascii" (show_internal=True, compact=False, attributes=None)$ 

Returns a string containing an ascii drawing of the tree.

### **Parameters**

- **show\_internal** includes internal edge names.
- compact use exactly one line per tip.
- attributes A list of node attributes to shown in the ASCII representation.

get\_cached\_content (store\_attr=None, container\_type=<type 'set'>, \_store=None)

Returns a dictionary pointing to the preloaded content of each internal node under this tree.

Such a dictionary is intended to work as a cache for operations that require many traversal

Such a dictionary is intended to work as a cache for operations that require many traversal operations.

**Parameters store\_attr** (*None*) – Specifies the node attribute that

should be cached (i.e. name, distance, etc.). When none, the whole node instance is cached.

**Parameters** store – (internal use)

```
get_children()
```

Returns an independent list of node's children.

```
get_closest_leaf(topology_only=False, is_leaf_fn=None)
```

Returns node's closest descendant leaf and the distance to it.

**Parameters topology\_only** (*False*) – If set to True, distance between nodes will be referred to the number of nodes between them. In other words, topological distance will be used instead of branch length distances.

**Returns** A tuple containing the closest leaf referred to the current node and the distance to it.

```
get_common_ancestor(*target_nodes, **kargs)
```

Returns the first common ancestor between this node and a given list of 'target\_nodes'.

### **Examples:**

```
t = tree.Tree("(((A:0.1, B:0.01):0.001, C:0.0001):1.0[&&NHX:name=common], (D:0.
A = t.get_descendants_by_name("A")[0]
C = t.get_descendants_by_name("C")[0]
common = A.get_common_ancestor(C)
print common.name
```

```
get_descendants (strategy='levelorder', is_leaf_fn=None)
```

Returns a list of all (leaves and internal) descendant nodes.

**Parameters is\_leaf\_fn** (None) – See TreeNode.traverse() for documentation.

```
get_distance (target, target2=None, topology_only=False)
```

Returns the distance between two nodes. If only one target is specified, it returns the distance bewtween the target and the current node.

#### **Parameters**

- target a node within the same tree structure.
- **target2** a node within the same tree structure. If not specified, current node is used as target2.
- **topology\_only** (*False*) If set to True, distance will refer to the number of nodes between target and target2.

**Returns** branch length distance between target and target2. If topology\_only flag is True, returns the number of nodes between target and target2.

```
get_edges (cached_content=None)
```

New in version 2.3.

Returns the list of edges of a tree. Each egde is represented as a tuple of two elements, each containing the list of nodes separated by the edge.

```
get_farthest_leaf (topology_only=False, is_leaf_fn=None)
```

Returns node's farthest descendant node (which is always a leaf), and the distance to it.

**Parameters topology\_only** (*False*) – If set to True, distance between nodes will be referred to the number of nodes between them. In other words, topological distance will be used instead of branch length distances.

**Returns** A tuple containing the farthest leaf referred to the current node and the distance to it.

```
get_farthest_node (topology_only=False)
```

Returns the node's farthest descendant or ancestor node, and the distance to it.

**Parameters topology\_only** (*False*) – If set to True, distance between nodes will be referred to the number of nodes between them. In other words, topological distance will be used instead of branch length distances.

**Returns** A tuple containing the farthest node referred to the current node and the distance to it.

```
get_leaf_names (is_leaf_fn=None)
```

Returns the list of terminal node names under the current node.

Parameters is  $leaf_fn(None)$  - See TreeNode.traverse() for documentation

```
get_leaves (is_leaf_fn=None)
```

Returns the list of terminal nodes (leaves) under this node.

Parameters is  $leaf_fn(None)$  - See TreeNode.traverse() for documentation.

```
get_leaves_by_name (name)
```

Returns a list of leaf nodes matching a given name.

```
get_midpoint_outgroup()
```

Returns the node that divides the current tree into two distance-balanced partitions.

```
get_monophyletic(values, target_attr)
```

New in version 2.2.

Returns a list of nodes matching the provided monophyly criteria. For a node to be considered a match, all *target\_attr* values within and node, and exclusively them, should be grouped.

#### **Parameters**

- values a set of values for which monophyly is expected.
- target\_attr node attribute being used to check monophyly (i.e. species for species trees, names for gene family trees).

#### get\_sisters()

Returns an indepent list of sister nodes.

```
get_topology_id (attr='name')
```

New in version 2.3.

Returns the unique ID representing the topology of the current tree. Two trees with the same topology will produce the same id. If trees are unrooted, make sure that the root node is not binary or use the tree.unroot() function before generating the topology id.

This is useful to detect the number of unique topologies over a bunch of trees, without requiring full distance methods.

The id is, by default, calculated based on the terminal node's names. Any other node attribute could be used instead.

```
get_tree_root()
```

Returns the absolute root node of current tree structure.

### img\_style

Branch length distance to parent node. Default = 0.0

#### is\_leaf()

Return True if current node is a leaf.

#### is\_root()

Returns True if current node has no parent

#### iter ancestors()

versionadded: 2.2

Iterates over the list of all ancestor nodes from current node to the current tree root.

```
iter_descendants (strategy='levelorder', is_leaf_fn=None)
```

Returns an iterator over all descendant nodes.

```
Parameters is leaf_fn(None) - See TreeNode.traverse() for documentation.
```

```
iter_edges (cached_content=None)
```

New in version 2.3.

Iterate over the list of edges of a tree. Each egde is represented as a tuple of two elements, each containing the list of nodes separated by the edge.

```
iter_leaf_names (is_leaf_fn=None)
```

Returns an iterator over the leaf names under this node.

```
Parameters is_leaf_fn (None) – See TreeNode.traverse() for documentation.
```

```
iter_leaves (is_leaf_fn=None)
```

Returns an iterator over the leaves under this node.

```
Parameters is leaf_fn(None) - See TreeNode.traverse() for documentation.
```

```
iter_prepostorder(is_leaf_fn=None)
```

Iterate over all nodes in a tree yielding every node in both pre and post order. Each iteration returns a postorder flag (True if node is being visited in postorder) and a node instance.

```
iter_search_nodes(**conditions)
```

Search nodes in an interative way. Matches are being yield as they are being found. This avoids to scan the full tree topology before returning the first matches. Useful when dealing with huge trees.

```
ladderize(direction=0)
```

Sort the branches of a given tree (swapping children nodes) according to the size of each partition.

```
t = Tree("(f,((d, ((a,b),c)),e));")
print t
#
# /-f
```

**populate** (size, names\_library=None, reuse\_names=False, random\_branches=False, branch\_range=(0, 1), support\_range=(0, 1))
Generates a random topology by populating current node.

### **Parameters**

- names\_library (*None*) If provided, names library (list, set, dict, etc.) will be used to name nodes.
- **reuse\_names** (*False*) If True, node names will not be necessarily unique, which makes the process a bit more efficient.
- random\_branches (*False*) If True, branch distances and support values will be randomized.
- **branch\_range** ((0,1)) If random\_branches is True, this

range of values will be used to generate random distances.

**Parameters support\_range** ((0,1)) – If random\_branches is True,

this range of values will be used to generate random branch support values.

```
prune (nodes, preserve_branch_length=False)
```

Prunes the topology of a node to conserve only the selected list of leaf internal nodes. The minimum number of nodes that conserve the topological relationships among the requested nodes will be retained. Root node is always conserved.

Variables nodes – a list of node names or node objects that should be retained

**Parameters preserve\_branch\_length** (*False*) – If True, branch lengths

of the deleted nodes are transferred (summed up) to its parent's branch, thus keeping original distances among nodes.

### **Examples:**

```
t1 = Tree('((((((A,B)C)D,E)F,G)H,(I,J)K)root;', format=1)
t1.prune(['A', 'B'])
   /-A
/D /C|
/F| \-B
#
# /H| \-E
# / /
                           /-A
\#-root \-G
                         -root
\ −B
t1 = Tree('(((((A,B)C)D,E)F,G)H,(I,J)K)root;', format=1)
t1.prune(['A', 'B', 'C'])
# /-A
# /D /C|
# /F| \-B
# /H / \−E
# / /
                         -root- C|
#-root \-G
\ -B
t1 = Tree('(((((A,B)C)D,E)F,G)H,(I,J)K)root;', format=1)
t1.prune(['A', 'B', 'I'])
 /D /C|

/F| \-B
# /H| \-E
                            /-I
                          -root
                           / /-A
#-root \-G
# /
# / /-I
                             \C/
                              \ -B
 \backslash K /
t1 = Tree('(((((A,B)C)D,E)F,G)H,(I,J)K)root;', format=1)
t1.prune(['A', 'B', 'F', 'H'])
           /-A
#
         /D /C|
```

### remove\_child(child)

Removes a child from this node (parent and child nodes still exit but are no longer connected).

#### remove\_sister(sister=None)

Removes a sister node. It has the same effect as 'TreeNode.up.remove\_child(sister)'

If a sister node is not supplied, the first sister will be deleted and returned.

Parameters sister – A node instance

**Returns** The node removed

render (file\_name, layout=None, w=None, h=None, tree\_style=None, units='px', dpi=90)

Renders the node structure as an image.

#### Variables

- **file\_name** path to the output image file. valid extensions are .SVG, .PDF, .PNG
- layout a layout function or a valid layout function name
- tree style a *TreeStyle* instance containing the image properties
- units (px) "px": pixels, "mm": millimeters, "in": inches
- h (None) height of the image in units
- w (None) weight of the image in units
- **dpi** (300) dots per inches.

#### resolve\_polytomy (default\_dist=0.0, default\_support=0.0, recursive=True)

Resolve all polytomies under current node by creating an arbitrary dicotomic structure among the affected nodes. This function randomly modifies current tree topology and should only be used for compatibility reasons (i.e. programs rejecting multifurcated node in the newick representation).

- **default\_dist** (0.0) artificial branch distance of new nodes.
- **default support** (0.0) artificial branch support of new nodes.
- **recursive** (*True*) Resolve any polytomy under this node. When False, only current node will be checked and fixed.

```
robinson_foulds (t2, attr_t1='name', attr_t2='name', unrooted_trees=False, expand_polytomies=False, polytomy_size_limit=5, skip_large_polytomies=False, correct_by_polytomy_size=False, min support t1=0.0, min support t2=0.0)
```

Returns the Robinson-Foulds symmetric distance between current tree and a different tree instance.

#### **Parameters**

- **t2** reference tree
- attr\_t1 (name) Compare trees using a custom node attribute as a node name.
- attr\_t2 (False) Compare trees using a custom node attribute as a node name in target tree.
- attr\_t2 If True, consider trees as unrooted.
- **expand\_polytomies** (*False*) If True, all polytomies in the reference and target tree will be expanded into all possible binary trees. Robinson-foulds distance will be calculated between all tree combinations and the minimum value will be returned. See also, NodeTree.expand\_polytomy().

**Returns** (rf, rf\_max, common\_attrs, names, edges\_t1, edges\_t2, discarded\_edges\_t1, discarded\_edges\_t2)

```
search nodes(**conditions)
```

Returns the list of nodes matching a given set of conditions.

#### **Example:**

```
tree.search_nodes(dist=0.0, name="human")
```

## set\_outgroup (outgroup)

Sets a descendant node as the outgroup of a tree. This function can be used to root a tree or even an internal node.

**Parameters outgroup** – a node instance within the same tree structure that will be used as a basal node.

```
set_style (node_style)
```

Set 'node\_style' as the fixed style for the current node.

```
show (layout=None, tree_style=None, name='ETE')
```

Starts an interative session to visualize current node structure using provided layout and TreeStyle.

```
sort descendants(attr='name')
```

This function sort the branches of a given tree by considerening node names. After the tree is sorted, nodes are labeled using ascendent numbers. This can be used to ensure that nodes in a tree with the same node names are always labeled in the same way. Note that if duplicated names are present, extra criteria should be added to sort nodes.

Unique id is stored as a node.\_nid attribute

```
standardize(delete_orphan=True, preserve_branch_length=True)
```

New in version 2.3.

process current tree structure to produce a standardized topology: nodes with only one child are removed and multifurcations are automatically resolved.

# support

Branch support for current node

### swap\_children()

Swaps current children order.

```
traverse (strategy='levelorder', is_leaf_fn=None)
```

Returns an iterator to traverse the tree structure under this node.

#### **Parameters**

- **strategy** ("*levelorder*") set the way in which tree will be traversed. Possible values are: "preorder" (first parent and then children) 'postorder' (first children and the parent) and "levelorder" (nodes are visited in order from root to leaves)
- **is\_leaf\_fn** (*None*) If supplied, is\_leaf\_fn function will be used to interrogate nodes about if they are terminal or internal. is\_leaf\_fn function should receive a node instance as first argument and return True or False. Use this argument to traverse a tree by dynamically collapsing internal nodes matching is\_leaf\_fn.

#### unroot()

Unroots current node. This function is expected to be used on the absolute tree root node, but it can be also be applied to any other internal node. It will convert a split into a multifurcation.

up

Pointer to parent node

```
write(features=None, outfile=None, format=0, is_leaf_fn=None, format_root_node=False, dist_formatter=None, support_formatter=None, name formatter=None)
```

Returns the newick representation of current node. Several arguments control the way in which extra data is shown for every node:

### **Parameters**

- **features** a list of feature names to be exported using the Extended Newick Format (i.e. features=["name", "dist"]). Use an empty list to export all available features in each node (features=[])
- outfile writes the output to a given file
- **format** defines the newick standard used to encode the tree. See tutorial for details.
- **format\_root\_node** (*False*) If True, it allows features and branch information from root node to be exported as a part of the newick text string. For newick compatibility reasons, this is False by default.
- $is_{end} See TreeNode.traverse()$  for documentation.

# **Example:**

```
t.get_newick(features=["species", "name"], format=1)
```

#### Tree

alias of TreeNode

# 3.2 Treeview module

#### **Contents**

- · Treeview module
  - TreeStyle
  - NodeStyle
  - Faces
  - Color names

# 3.2.1 TreeStyle

### class TreeStyle

New in version 2.1.

Contains all the general image properties used to render a tree

- About tree design -

**Parameters layout\_fn** (*None*) – Layout function used to dynamically control the aspect of nodes. Valid values are: None or a pointer to a method, function, etc.

# - About tree shape -

- mode ("r") Valid modes are 'c'(ircular) or 'r'(ectangular).
- **orientation** (0) If 0, tree is drawn from left-to-right. If 1, tree is drawn from right-to-left. This property only makes sense when "r" mode is used.
- rotation (0) Tree figure will be rotate X degrees (clock-wise rotation).
- min\_leaf\_separation (1) Min separation, in pixels, between two adjacent branches
- branch\_vertical\_margin (0) Leaf branch separation margin, in pixels. This will add a separation of X pixels between adjacent leaf branches. In practice, increasing this value work as increasing Y axis scale.
- arc\_start (0) When circular trees are drawn, this defines the starting angle (in degrees) from which leaves are distributed (clock-wise) around the total arc span (0 = 3 o'clock).
- arc\_span (359) Total arc used to draw circular trees (in degrees).
- margin\_left (0) Left tree image margin, in pixels.
- margin\_right (0) Right tree image margin, in pixels.
- margin\_top (0) Top tree image margin, in pixels.
- margin\_bottom (0) Bottom tree image margin, in pixels.

### - About Tree branches -

#### **Parameters**

- scale (*None*) Scale used to draw branch lengths. If None, it will be automatically calculated.
- **optimal\_scale\_level** ("*mid*") Two levels of automatic branch scale detection are available: "mid" and "full". In full mode, branch scale will me adjusted to fully avoid dotted lines in the tree image. In other words, scale will be increased until the extra space necessary to allocated all branch-top/bottom faces and branch-right faces (in circular mode) is covered by real branches. Note, however, that the optimal scale in trees with very unbalanced branch lengths might be huge. If "mid" mode is selected (as it is by default), optimal scale will only satisfy the space necessary to allocate branch-right faces in circular trees. Some dotted lines (artificial branch offsets) will still appear when branch-top/bottom faces are larger than branch length. Note that both options apply only when scale is set to None (automatic).
- root\_opening\_factor (0.25) (from 0 to 1). It defines how much the center of a circular tree could be opened when adjusting optimal scale, referred to the total tree length. By default (0.25), a blank space up to 4 times smaller than the tree width could be used to calculate the optimal tree scale. A 0 value would mean that root node should always be tightly adjusted to the center of the tree.
- **complete\_branch\_lines\_when\_necessary** (*True*) True or False. Draws an extra line (dotted by default) to complete branch lengths when the space to cover is larger than the branch itself.
- extra\_branch\_line\_type (2) 0=solid, 1=dashed, 2=dotted
- extra\_branch\_line\_color" ("gray") RGB code or name in SVG\_COLORS
- **force\_topology** (*False*) Convert tree branches to a fixed length, thus allowing to observe the topology of tight nodes
- **draw\_guiding\_lines** (*False*) Draw guidelines from leaf nodes to aligned faces
- **guiding\_lines\_type** (2) 0=solid, 1=dashed, 2=dotted.
- guiding\_lines\_color ("gray") RGB code or name in SVG\_COLORS

#### - About node faces -

- **allow\_face\_overlap** (*False*) If True, node faces are not taken into account to scale circular tree images, just like many other visualization programs. Overlapping among branch elements (such as node labels) will be therefore ignored, and tree size will be a lot smaller. Note that in most cases, manual setting of tree scale will be also necessary.
- **draw\_aligned\_faces\_as\_table** (*True*) Aligned faces will be drawn as a table, considering all columns in all node faces.
- **children\_faces\_on\_top** (*True*) When floating faces from different nodes overlap, children faces are drawn on top of parent faces. This can be reversed

by setting this attribute to false.

#### - Addons -

#### **Parameters**

- **show\_border** (*False*) Draw a border around the whole tree
- show\_scale (*True*) Include the scale legend in the tree image
- **show\_leaf\_name** (*False*) Automatically adds a text Face to leaf nodes showing their names
- **show\_branch\_length** (*False*) Automatically adds branch length information on top of branches
- **show\_branch\_support** (*False*) Automatically adds branch support text in the bottom of tree branches

# - Tree surroundings -

The following options are actually Face containers, so graphical elements can be added just as it is done with nodes. In example, to add tree legend:

```
TreeStyle.legend.add_face(CircleFace(10, "red"), column=0)
TreeStyle.legend.add_face(TextFace("0.5 support"), column=1)
```

#### **Parameters**

- aligned\_header a FaceContainer aligned to the end of the tree and placed at the top part.
- aligned\_foot a FaceContainer aligned to the end of the tree and placed at the bottom part.
- **legend** a FaceContainer with an arbitrary number of faces representing the legend of the figure.
- **legend\_position=4** (*4*) TopLeft corner if 1, TopRight if 2, BottomLeft if 3, BottomRight if 4
- title A Face container that can be used as tree title

# class FaceContainer

New in version 2.1.

Use this object to create a grid of faces. You can add faces to different columns.

```
add_face (face, column)
add the face face to the specified column
```

# 3.2.2 NodeStyle

```
class NodeStyle (*args, **kargs)
```

New in version 2.1.

A dictionary with all valid node graphical attributes.

- **fgcolor** (#0030c1) RGB code or name in SVG\_COLORS
- **bgcolor** (#FFFFFF) RGB code or name in SVG\_COLORS
- node\_bgcolor (#FFFFFF) RGB code or name in SVG\_COLORS
- partition\_bgcolor (#FFFFFF) RGB code or name in SVG\_COLORS
- faces\_bgcolor (#FFFFFF) RGB code or name in SVG\_COLORS
- vt\_line\_color (#000000) RGB code or name in SVG\_COLORS
- hz\_line\_color (#000000) RGB code or name in SVG\_COLORS
- hz\_line\_type (0) integer number
- vt\_line\_type (0) integer number
- size (3) integer number
- shape ("circle") "circle", "square" or "sphere"
- draw\_descendants (*True*) Mark an internal node as a leaf.
- hz\_line\_width (0) integer number representing the width of the line in pixels. A line width of zero indicates a cosmetic pen. This means that the pen width is always drawn one pixel wide, independent of the transformation set on the painter.
- vt\_line\_width (0) integer number representing the width of the line in pixels. A line width of zero indicates a cosmetic pen. This means that the pen width is always drawn one pixel wide, independent of the transformation set on the painter.

### **3.2.3 Faces**

add\_face\_to\_node (face, node, column, aligned=False, position='branch-right')
Adds a Face to a given node.

Parameters face - A Face instance

#### **Parameters**

- **node** a tree node instance (Tree, PhyloTree, etc.)
- **column** An integer number starting from 0
- **position** ("branch-right") Possible values are "branch-right", "branch-top", "branch-bottom", "float", "float-behind" and "aligned".

#### class Face

Base Face object. All Face types (i.e. TextFace, SeqMotifFace, etc.) inherit the following options:

- margin\_left (0) in pixels
- margin\_right (0) in pixels
- margin\_top (0) in pixels
- margin\_bottom (0) in pixels

- opacity (1.0) a float number in the (0,1) range
- **rotable** (*True*) If True, face will be rotated when necessary (i.e. when circular mode is enabled and face occupies an inverted position.)
- $hz_align(0) 0 left$ , 1 center, 2 right
- **vt\_align** (1) 0 top, 1 center, 2 bottom
- background.color background color of face plus all its margins.
- inner\_background.color background color of the face excluding margins
- border Border around face margins.
- inner\_border Border around face excluding margins.

# border and inner\_border sub-parameters:

#### **Parameters**

- (inner\_)border.type (0) 0=solid, 1=dashed, 2=dotted
- (inner\_)border.width (*None*) a positive integer number. Zero indicates a cosmetic pen. This means that the pen width is always drawn one pixel wide, independent of the transformation set on the painter. A "None" value means invisible border.
- (inner\_)border.color (black) RGB or color name in SVG\_COLORS

See also specific options for each face type.

### **Parameters**

- **text** Text to be drawn
- ftype Font type, e.g. Arial, Verdana, Courier
- **fsize** Font size, e.g. 10,12,6, (default=10)
- fgcolor Foreground font color. RGB code or color name in SVG\_COLORS
- **penwidth** Penwdith used to draw the text.
- fstyle "normal" or "italic"
- tight\_text (False) When False, boundaries of the text are

approximated according to general font metrics, producing slightly worse aligned text faces but improving the performance of tree visualization in scenes with a lot of text faces.

```
class AttrFace (attr, ftype='Verdana', fsize=10, fgcolor='black', penwidth=0, text_prefix='', text_suffix='', formatter=None, fstyle='normal', tight_text=False)

Dynamic text Face. Text rendered is taken from the value of a given node attribute.
```

#### **Parameters**

- attr Node's attribute that will be drawn as text
- ftype Font type, e.g. Arial, Verdana, Courier, (default="Verdana")

131

• **fsize** – Font size, e.g. 10,12,6, (default=10)

3.2. Treeview module

- fgcolor Foreground font color. RGB code or name in SVG\_COLORS
- **penwidth** Penwdith used to draw the text. (default is 0)
- **text\_prefix** text\_rendered before attribute value
- text suffix text rendered after attribute value
- **formatter** a text string defining a python formater to process the attribute value before renderer. e.g. "%0.2f"
- fstyle "normal" or "italic"

class ImgFace (img\_file, width=None, height=None)

Creates a node Face using an external image file.

#### **Parameters**

- img file path to the image file.
- width (*None*) if provided, image will be scaled to this width (in pixels)
- **height** (*None*) if provided, image will be scaled to this height (in pixels)

If only one dimension value (width or height) is provided, the other will be calculated to keep aspect ratio.

class CircleFace (radius, color, style='circle')

New in version 2.1.

Creates a Circle or Sphere Face.

### **Parameters**

- radius integer number defining the radius of the face
- color Color used to fill the circle. RGB code or name in SVG COLORS
- **style** ("circle") Valid values are "circle" or "sphere"

class RectFace (width, height, fgcolor, bgcolor)

New in version 2.3.

Creates a Rectangular solid face.

class StackedBarFace (percents, width, height, colors=None, line\_color=None)

Creates a new molecular sequence face object. :param seq: Sequence string to be drawn :param seqtype: Type of sequence: "nt" or "aa" :param fsize: Font size, (default=10)

You can set custom colors for amino-acids or nucleotides:

- **codon** (*None*) a string that corresponds to the reverse translation of the amino-acid sequence
- **col\_w** (*None*) width of the column (if col\_w is lower than font size, letter wont be displayed)

- **fg\_colors** (*None*) dictionary of colors for foreground, with as keys each possible character in sequences, and as value the colors
- **bg\_colors** (*None*) dictionary of colors for background, with as keys each possible character in sequences, and as value the colors
- alt\_col\_w (3) works together with special\_col option, defines the width of given columns
- **special\_col** (*None*) list of lists containing the bounds of columns to be displayed with alt\_col\_w as width
- interactive (False) more info can be displayed when mouse over sequence

```
class InteractiveLetterItem(*arg, **karg)
```

This is a class

#### hoverEnterEvent(e)

when mouse is over

#### hoverLeaveEvent (e)

when mouse leaves area

New in version 2.2.

Creates a face based on an amino acid or nucleotide sequence and a list of motif regions.

### **Parameters**

- **seq** (*None*) a text string containing an aa or nt sequence. If not provided, seq and compact seq motif modes will not be available.
- **motifs** (*None*) a list of motif regions referred to original sequence. Each motif is defined as a list containing the following information:

### Where:

- seq.start: Motif start position referred to the full sequence
- seq.end: Motif end position referred to the full sequence
- **shape:** Shape used to draw the motif. Available values are:

```
* o = circle or ellipse
* > = triangle (base to the left)
* < = triangle (base to the left)</li>
* ^ = triangle (base at bottom)
* v = triangle (base on top )
```

- \* <> = diamond
- \* [] = rectangle
- \* () = round corner rectangle
- \* seq = Show a color and the corresponding letter of each sequence position
- \* compact seq = Show a color for each sequence position
- width: total width of the motif (or sequence position width if seq motif type)
- height: total height of the motif (or sequence position height if seq motif type)
- fgcolor: color for the motif shape border
- bgcolor: motif background color. Color code or name can be preceded with the "rgradient:" tag to create a radial gradient effect.
- **intermotif\_format** (*line*) How should spaces among motifs be filled. Available values are: "line", "blank", "none" and "seq", "compactseq".
- **seqtail\_format** (*none*) How should remaining tail sequence be drawn. Available values are: "line", "seq", "compactseq" or "none"
- **seq\_format** (*compactseq*) How should sequence be rendered in case no motif regions are provided. Available values are: "seq" and "compactseq"

class BarChartFace (values, deviations=None, width=200, height=100, colors=None, labels=None, min\_value=0, max\_value=None)
New in version 2.2.

### **Parameters**

- values a list of values each representing a vertical bar.
- width (200) width of the bar chart.
- **height** (100) height of the bar chart
- colors (None) a list of colors, one per bar value
- label (*None*) a list of labels, one per bar
- min value (0) min value to set the scale of the chart.
- max\_value (*None*) max value to set the scale of the chart.

**class PieChartFace** (percents, width, height, colors=None, line\_color=None) New in version 2.2.

- percents a list of values summing up 100.
- width width of the piechart
- height height of the piechart
- colors a list of colors (same length as percents)

• line\_color – color used to render the border of the piechart (None=transparent)

### **Parameters**

- max\_v maximum value used to build the build the plot scale.
- max\_v minimum value used to build the build the plot scale.
- **center\_v** Center value used to scale plot and heatmap.
- width (200) Plot width in pixels.
- **height** (40) Plot width in pixels.
- style (lines) Plot style: "lines", "bars", "cbars" or "heatmap".
- **colorscheme** (2) colors used to create the gradient from min values to max values. 0=green & blue; 1=green & red; 2=red & blue. In all three cases, missing values are rendered in black and transition color (values=center) is white.

### class TreeFace (tree, tree\_style)

New in version 2.1.

Creates a Face containing a Tree object. Yes, a tree within a tree :)

#### **Parameters**

- tree An ETE Tree instance (Tree, PhyloTree, etc...)
- tree\_style A TreeStyle instance defining how tree show be drawn

### class StaticItemFace (item)

New in version 2.1.

Creates a face based on an external QtGraphicsItem object. QGraphicsItem object is expected to be independent from tree node properties, so its content is assumed to be static (drawn only once, no updates when tree changes).

Parameters item – an object based on QGraphicsItem

```
class DynamicItemFace (constructor, *args, **kargs)
```

New in version 2.1.

Creates a face based on an external QGraphicsItem object whose content depends on the node that is linked to.

**Parameters constructor** – A pointer to a method (function or class constructor) returning a QGraphicsItem based object. "constructor" method is expected to receive a node instance as the first argument. The rest of arguments passed to Item-Face are optional and will passed also to the constructor function.

### 3.2.4 Color names

SVG\_COLORS

Ap Red colors	art fron	n RGB colo	or codes, the fol	llowing	SVGc	olor	names are s Brown color		ed:	
IndianRed	CD 5C 5C	205 92 92	GreenYellow	AD FF 2F	173 255	47	Cornsilk	FF F8 DC	255 248	220
LightCoral	F0 80 80	240 128 128	Chartreuse	7F FF 00		0	BlanchedAlmond	FF EB CD	255 235	205
Salmon	FA 80 72	250 128 114	LawnGreen		124 252	0	Bisque	FF E4 C4		
DarkSalmon	E9 96 7A		Lime	00 FF 00		0	NavajoWhite	FF DE AD		
LightSalmon	FF AO 7A		LimeGreen	32 CD 32	50 205	50	Wheat	F5 DE B3		
Crimson	DC 14 3C	220 20 60	PaleGreen		152 251 3		BurlyWood	DE B8 87		
Red	FF 00 00		LightGreen	90 EE 90			Tan	D2 B4 8C		
FireBrick	B2 22 22	178 34 34	MediumSpringGreen		0 250		RosyBrown	BC 8F 8F	188 143	
DarkRed	8B 00 00	139 0 0	SpringGreen	00 FF 7F	0 255		SandyBrown	F4 A4 60	244 164	96
Pink colors	05 00 00	133 0 0	MediumSeaGreen	3C B3 71	60 179 3		Goldenrod	DA A5 20	218 165	32
Pink	FF CO CB	255 192 203	SeaGreen	2E 8B 57	46 139	87	DarkGoldenrod	B8 86 0B	184 134	
LightPink		255 182 193	ForestGreen	22 8B 22	34 139	34	Peru	CD 85 3F	205 133	
HotPink	FF 69 B4	255 105 180	Green	00 80 00	0 128	0	Chocolate	D2 69 1E	210 105	30
DeepPink	FF 14 93	255 20 147	DarkGreen	00 64 00		0	SaddleBrown	8B 45 13	139 69	
MediumVioletRed	C7 15 85	199 21 133	YellowGreen	9A CD 32		50	Sienna	A0 52 2D	160 82	
	DB 70 93	219 112 147		6B 8E 23						42
PaleVioletRed	DB 70 93	215 112 147	Olive	80 80 00		35	Brown	A5 2A 2A 80 00 00	165 42 128 0	
Orange colors	DD 30 77	OFF 160 100	Olive			0	Maroon		128 0	0
LightSalmon	FF A0 7A		DarkOliveGreen	55 6B 2F		47	White color	_	255 255	055
Coral	FF 7F 50	255 127 80	MediumAquamarine				White	FF FF FF		
Tomato	FF 63 47	255 99 71	DarkSeaGreen	8F BC 8F			Snow	FF FA FA		
OrangeRed 	FF 45 00	255 69 0	LightSeaGreen	20 B2 AA	32 178 3		Honeydew	FO FF FO		
DarkOrange	FF 8C 00	255 140 0	DarkCyan	00 8B 8B	0 139 3		MintCream	F5 FF FA		
Orange	FF A5 00	255 165 0	Teal	00 80 80	0 128	128	Azure	FO FF FF		
Yellow colors			Blue/Cyan colo				AliceBlue	FO F8 FF	240 248	255
Gold	FF D7 00	255 215 0	Aqua	00 FF FF	0 255 2	255	GhostWhite	F8 F8 FF	248 248	255
Yellow	FF FF 00	255 255 0	Cyan	00 FF FF	0 255 2	255	WhiteSmoke	F5 F5 F5	245 245	245
LightYellow	FF FF E0	255 255 224	LightCyan	EO FF FF	224 255 2	255	Seashell	FF F5 EE	255 245	238
LemonChiffon	FF FA CD	255 250 205	PaleTurquoise	AF EE EE	175 238 2	238	Beige	F5 F5 DC	245 245	220
LightGoldenrodYellow	FA FA D2	250 250 210	Aquamarine	7F FF D4	127 255 2	212	OldLace	FD F5 E6	253 245	230
PapayaWhip	FF EF D5	255 239 213	Turquoise	40 E0 D0	64 224 2	208	FloralWhite	FF FA FO	255 250	240
Moccasin	FF E4 B5	255 228 181	MediumTurquoise	48 D1 CC	72 209 2	204	Ivory	FF FF F0	255 255	240
PeachPuff	FF DA B9	255 218 185	DarkTurquoise	00 CE D1	0 206 2	209	AntiqueWhite	FA EB D7	250 235	215
PaleGoldenrod	EE E8 AA	238 232 170	CadetBlue	5F 9E A0	95 158 3	160	Linen	FA FO E6	250 240	230
Khaki	F0 E6 8C	240 230 140	SteelBlue	46 82 B4	70 130 3	180	LavenderBlush	FF FO F5	255 240	245
DarkKhaki	BD B7 6B	189 183 107	LightSteelBlue	BO C4 DE	176 196 2	222	MistyRose	FF E4 E1	255 228	225
Purple colors			PowderBlue	B0 E0 E6	176 224 2	230	Gray colors			
Lavender	E6 E6 FA	230 230 250	LightBlue	AD D8 E6	173 216 2	230	Gainsboro	DC DC DC	220 220	220
Thistle	D8 BF D8	216 191 216	SkyBlue	87 CE EB	135 206 2	235	LightGrey	D3 D3 D3	211 211	211
Plum		221 160 221	LightSkyBlue	87 CE FA	135 206 2	250	Silver	C0 C0 C0	192 192	192
Violet		238 130 238	DeepSkyBlue	00 BF FF	0 191 2		DarkGray	A9 A9 A9	169 169	169
Orchid		218 112 214	DodgerBlue	1E 90 FF			Gray	80 80 80		
Fuchsia	FF 00 FF	255 0 255	CornflowerBlue	64 95 ED	100 149 2		DimGray	69 69 69	105 105	
Magenta	FF 00 FF		MediumSlateBlue	7B 68 EE			LightSlateGray	77 88 99	119 136	
MediumOrchid	BA 55 D3	186 85 211	RoyalBlue	41 69 E1	65 105 2		SlateGray	70 80 90	112 128	
BlueViolet	8A 2B E2		MediumBlue	00 00 CD		205	Black	00 00 00	0 0	
DarkViolet	94 00 D3		DarkBlue	00 00 8B		139				
DarkOrchid	99 32 CC	153 50 204	Navy	00 00 80		128				
	8B 00 8B		MidnightBlue	19 19 70						
DarkMagenta	80 00 88		MunightBlue	15 15 70	23 23 .	112				
Purple										
Indigo	4B 00 82									
SlateBlue	6A 5A CD									
DarkSlateBlue	48 3D 8B	72 61 139								
MediumSlateBlue	7B 68 EE	123 104 238								

# COLOR\_SCHEMES

 $\verb"random_color" (h=None, l=None, s=None, num=None, sep=None, seed=None)"$ 

returns the RGB code of a random color. Hue (h), Lightness (l) and Saturation (s) of the generated color could be fixed using the pertinent function argument.

# 3.3 PhyloTree class

Bases: ete2.coretype.tree.TreeNode

Extends the standard TreeNode instance. It adds specific attributes and methods to work with phylogentic trees.

### **Parameters**

- **newick** Path to the file containing the tree or, alternatively, the text string containing the same information.
- alignment file containing a multiple sequence alignment.
- alg\_format "fasta", "phylip" or "iphylip" (interleaved)
- **format** sub-newick format

FORMAT	DESCRIPTION
0	flexible with support values
1	flexible with internal node names
2	all branches + leaf names + internal supports
3	all branches + all names
4	leaf branches + leaf names
5	internal and leaf branches + leaf names
6	internal branches + leaf names
7	leaf branches + all names
8	all names
9	leaf names
100	topology only

• **sp\_naming\_function** — Pointer to a parsing python function that receives nodename as first argument and returns the species name (see PhyloNode.set\_species\_naming\_function(). By default, the 3 first letter of nodes will be used as species identifiers.

**Returns** a tree node object which represents the base of the tree.

Add NCBI taxonomy annotation to all descendant nodes. Leaf nodes are expected to contain a feature (name, by default) encoding a valid taxid number.

All descendant nodes (including internal nodes) are annotated with the following new features:

Node.spname: scientific spcies name as encoded in the NCBI taxonomy database

*Node.named\_lineage*: the NCBI lineage track using scientific names

Node.taxid: NCBI taxid number

*Node.lineage*: same as named\_lineage but using taxid codes.

Note that for internal nodes, NCBI information will refer to the first common lineage of the grouped species.

#### **Parameters**

- **taxid\_attr** (*name*) the name of the feature that should be used to access the taxid number associated to each node.
- tax2name (*None*) A dictionary where keys are taxid numbers and

values are their translation into NCBI scientific name. Its use is optional and allows to avoid database queries when annotating many trees containing the same set of taxids.

**Parameters tax2track** (*None*) – A dictionary where keys are taxid numbers and

values are their translation into NCBI lineage tracks (taxids). Its use is optional and allows to avoid database queries when annotating many trees containing the same set of taxids.

Parameters tax2rank (None) – A dictionary where keys are taxid numbers and

values are their translation into NCBI rank name. Its use is optional and allows to avoid database queries when annotating many trees containing the same set of taxids.

:param None dbfile: If provided, the provided file will be used as a local copy of the NCBI taxonomy database.

**Returns** tax2name (a dictionary translating taxid numbers into

scientific name), tax2lineage (a dictionary translating taxid numbers into their corresponding NCBI lineage track) and tax2rank (a dictionary translating taxid numbers into rank names).

## collapse\_lineage\_specific\_expansions (species=None, return\_copy=True)

Converts lineage specific expansion nodes into a single tip node (randomly chosen from tips within the expansion).

**Parameters species** (*None*) – If supplied, only expansions matching the species criteria will be pruned. When None, all expansions within the tree will be processed.

### get\_age (species2age)

Implements the phylostratigrafic method described in:

Huerta-Cepas, J., & Gabaldon, T. (2011). Assigning duplication events to relative temporal scales in genome-wide studies. Bioinformatics, 27(1), 38-45.

# get\_age\_balanced\_outgroup (species2age)

New in version 2.2.

Returns the node better balance current tree structure according to the topological age of the different leaves and internal node sizes.

**Parameters species2age** – A dictionary translating from leaf names into a topological age.

#### get descendant evol events (sos thr=0.0)

Returns a list of **all** duplication and speciation events detected after this node. Nodes are assumed to be duplications when a species overlap is found between its child linages. Method is described more detail in:

"The Human Phylome." Huerta-Cepas J, Dopazo H, Dopazo J, Gabaldon T. Genome Biol. 2007;8(6):R109.

# get\_farthest\_oldest\_leaf(species2age, is\_leaf\_fn=None)

Returns the farthest oldest leaf to the current one. It requires an species2age dictionary with the age estimation for all species.

**Parameters is\_leaf\_fn** (*None*) – A pointer to a function that receives a node instance as unique argument and returns True or False. It can be used to dynamically collapse nodes, so they are seen as leaves.

## get\_farthest\_oldest\_node(species2age)

New in version 2.1.

Returns the farthest oldest node (leaf or internal). The difference with get\_farthest\_oldest\_leaf() is that in this function internal nodes grouping seqs from the same species are collapsed.

## get\_my\_evol\_events (sos\_thr=0.0)

Returns a list of duplication and speciation events in which the current node has been involved. Scanned nodes are also labeled internally as dup=TruelFalse. You can access this labels using the 'node.dup' sintaxis.

Method: the algorithm scans all nodes from the given leafName to the root. Nodes are assumed to be duplications when a species overlap is found between its child linages. Method is described more detail in:

"The Human Phylome." Huerta-Cepas J, Dopazo H, Dopazo J, Gabaldon T. Genome Biol. 2007;8(6):R109.

#### 

Calculates all possible species trees contained within a duplicated gene family tree as described in Treeko (see Marcet and Gabaldon, 2011).

### Parameters autodetect duplications (True) – If True, duplication

nodes will be automatically detected using the Species Overlap algorithm (PhyloNode.get\_descendants\_evol\_events(). If False, duplication nodes within the original tree are expected to contain the feature "evoltype=D".

**Parameters features** (*None*) – A list of features that should be

mapped from the original gene family tree to each species tree subtree.

**Returns** (number\_of\_sptrees, number\_of\_dups, species\_tree\_iterator)

### get\_species()

Returns the set of species covered by its partition.

#### iter\_species()

Returns an iterator over the species grouped by this node.

```
link to alignment (alignment, alg format='fasta', **kwargs)
```

 $\verb"ncbi_compare" (autodetect_duplications=True, cached_content=None)$ 

# reconcile (species\_tree)

Returns the reconcilied topology with the provided species tree, and a list of evolutionary events inferred from such reconciliation.

```
set_species_naming_function(fn)
```

Sets the parsing function used to extract species name from a node's name.

**Parameters fn** – Pointer to a parsing python function that receives nodename as first argument and returns the species name.

```
# Example of a parsing function to extract species names for
# all nodes in a given tree.
def parse_sp_name(node_name):
    return node_name.split("_")[1]
tree.set_species_naming_function(parse_sp_name)
```

### species

split\_by\_dups (autodetect\_duplications=True)

Returns the list of all subtrees resulting from splitting current tree by its duplication nodes.

**Parameters autodetect\_duplications** (*True*) – If True, duplication

nodes will be automatically detected using the Species Overlap algorithm (PhyloNode.get\_descendants\_evol\_events(). If False, duplication nodes within the original tree are expected to contain the feature "evoltype=D".

Returns species\_trees

### PhyloTree

alias of PhyloNode

#### class EvolEvent

Basic evolutionary event. It stores all the information about an event(node) ocurred in a phylogenetic tree.

```
etype: D (Duplication), S (Speciation), L (gene loss), in_seqs: the list of sequences in one side of the event. out_seqs: the list of sequences in the other side of the event node: link to the event node in the tree
```

#### **Contents**

• Clustering module

# 3.4 Clustering module

Creates a new Cluster Tree object, which is a collection of ClusterNode instances connected in a hierarchical way, and representing a clustering result.

a newick file or string can be passed as the first argument. An ArrayTable file or instance can be passed as a second argument.

**Examples:** t1 = Tree() # creates an empty tree t2 = Tree( '(A:1,(B:1,(C:1,D:1):0.5):0.5);' ) t3 = Tree( '/home/user/myNewickFile.txt' )

## get\_dunn (clusters, fdist=None)

Calculates the Dunn index for the given set of descendant nodes.

# get\_leaf\_profiles()

Returns the list of all the profiles associated to the leaves under this node.

# get\_silhouette(fdist=None)

Calculates the node's silhouette value by using a given distance function. By default, euclidean distance is used. It also calculates the deviation profile, mean profile, and inter/intracluster distances.

## It sets the following features into the analyzed node:

- node.intracluster
- node.intercluster
- · node.silhouete

intracluster distances a(i) are calculated as the Centroid Diameter

intercluster distances b(i) are calculated as the Centroid linkage distance

\*\* Rousseeuw, P.J. (1987) Silhouettes: A graphical aid to the interpretation and validation of cluster analysis. J. Comput. Appl. Math., 20, 53-65.

## iter\_leaf\_profiles()

Returns an iterator over all the profiles associated to the leaves under this node.

# link\_to\_arraytable (arraytbl)

Allows to link a given arraytable object to the tree structure under this node. Row names in the arraytable object are expected to match leaf names.

Returns a list of nodes for with profiles could not been found in arraytable.

## set distance function (fn)

Sets the distance function used to calculate cluster distances and silouette index.

## ARGUMENTS:

fn: a pointer to python function acepting two arrays (numpy) as arguments.

## **EXAMPLE:**

# A simple euclidean distance my\_dist\_fn = lambda x,y: abs(x-y) tree.set\_distance\_function(my\_dist\_fn)

### ClusterTree

alias of ClusterNode

New in version 2.1.

# 3.5 Nexml module

## 3.5.1 Nexml classes linked to ETE

class Nexml (\*args, \*\*kargs)

Creates a new nexml project.

build\_from\_file (fname, index\_otus=True)

Populate Nexml project with data in a nexml file.

#### 3.5.2 Generic Nexml classes

A concrete implementation of the AbstractChar element.

AAChar.superclass

alias of AbstractChar

class AAFormat (about=None, meta=None, states=None, char=None, set=None, valueOf =None)

The AAFormat class is the container of amino acid column definitions.

AAFormat.superclass

alias of AbstractFormat

class AAMapping (state=None, valueOf\_=None)

An IUPAC ambiguity mapping.

AAMapping.superclass

alias of AbstractMapping

 $\begin{array}{lll} \textbf{class AAMatrixObsRow} \ (about=None, & meta=None, & label=None, & id=None, & otu=None, \\ & cell=None, \ set=None, \ valueOf\_=None) \end{array}$ 

This is a row in a matrix of amino acid data containing granular observations.

AAMatrixObsRow.superclass

alias of AbstractObsRow

class AAMatrixSeqRow (about=None, meta=None, label=None, id=None, otu=None,  $seq=None, valueOf\_=None$ )

This is a row in a matrix of amino acid data containing raw sequence data.

AAMatrixSeqRow.superclass

alias of AbstractSeqRow

**class AAObs** (about=None, meta=None, label=None, char=None, state=None, valueOf\_=None)

This is a single cell in a matrix containing an amino acid observation.

AAObs.superclass

alias of AbstractObs

**class AAObsMatrix** (about=None, meta=None, row=None, set=None, valueOf\_=None)

A matrix of rows with single character observations.

AAObsMatrix.superclass

alias of AbstractObsMatrix

 $\begin{array}{c} \textbf{class AAPolymorphicStateSet} \ (about=None, \ meta=None, \ label=None, \ id=None, \ symbol=None, \ member=None, \ uncertain\_state\_set=None, \\ \end{array}$ 

*valueOf\_=None*)

The AAPolymorphicStateSet defines a polymorphic ambiguity mapping.

AAPolymorphicStateSet.superclass alias of AbstractPolymorphicStateSet

 $\textbf{class AASeqMatrix} (about=None, meta=None, row=None, set=None, valueOf\_=None)$ 

A matrix of rows with amino acid data as sequence strings.

AASeqMatrix.superclass

alias of AbstractSeqMatrix

**class AAState** (about=None, meta=None, label=None, id=None, symbol=None, valueOf = None)

This is a concrete implementation of the state element, which requires a symbol element, in this case restricted to AAToken, i.e. a single IUPAC amino acid symbol, and optional mapping elements to refer to other states.

AAState.superclass

alias of AbstractState

A container for a set of states.

AAStates.superclass

alias of AbstractStates

class AAUncertainStateSet (about=None, meta=None, label=None, id=None, symbol=None, member=None, valueOf =None)

The AAUncertainStateSet defines an uncertain ambiguity mapping.

AAUncertainStateSet.superclass

alias of AbstractUncertainStateSet

class AbstractBlock (about=None, meta=None, label=None, id=None, otus=None, format=None, valueOf =None)

The AbstractBlock is the superclass for blocks that contain an element structure of type Abstract-Format.

AbstractBlock.superclass

alias of TaxaLinked

class AbstractCells (about=None, meta=None, label=None, id=None, otus=None, format=None, matrix=None, valueOf\_=None)

The AbstractSeqBlock type is the superclass for character blocks that consist of granular character state observations.

AbstractCells.superclass alias of AbstractBlock

The AbstractChar type is the superclass for a column definition, which may have a "states" attribute that refers to an AbstractStates element, a codon attribute of type CodonPosition and an id attribute that may be an actual id (e.g. for categorical matrices where observations explicitly refer to a column definition) or an integer for sequence matrices.

AbstractChar.superclass alias of IDTagged

The AbstractEdge superclass is what concrete edges inherit from by restriction. It represents an edge element much like that of GraphML, i.e. an element that connects node elements.

AbstractEdge.superclass alias of IDTagged

class AbstractFormat (about=None, meta=None, states=None, char=None, set=None, valueOf =None)

The AbstractFormat type is the superclass for the element that defines the allowed characters and states in a matrix, and their ambiguity mapping. It may enclose AbstractStates elements that define states and their mappings, and AbstractChar elements that specify which AbstractStates apply to which matrix columns.

AbstractFormat.superclass alias of Annotated

# class AbstractMapping (state=None, valueOf\_=None)

The AbstractMapping type is the superclass for an ambiguity mapping. In an instance document, a subclass of this type will look like <member state="st1"/>, i.e. an element called "member" with an attribute called "state" whose value is an id reference that refers to an element that subclasses AbstractState. The purpose of AbstractMapping is to specify which other states may be implied, e.g. a nucleotide symbol "N" would have mappings to "A", "C", "G" and "T".

AbstractMapping.superclass alias of Base

The AbstractNetwork superclass is what a concrete network inherits from.

 $\label{local_abstractNetwork.superclass} \textbf{alias of} \; \texttt{IDTagged}$ 

 $\begin{array}{ll} \textbf{class AbstractNode} \ (about=None, & meta=None, & label=None, & id=None, & otu=None, \\ root=False, valueOf\_=None) \end{array}$ 

The AbstractNode superclass is what concrete nodes inherit from by restriction. It represents a node element much like that of GraphML, i.e. an element that is connected into a tree by edge elements.

AbstractNode.superclass alias of OptionalTaxonLinked

class AbstractObs (about=None, meta=None, label=None, char=None, state=None, valueOf =None)

The AbstractObs type is the superclass for single observations, i.e. cells in a matrix. A concrete instance of AbstractObs has a "char" attribute that refers to an explicitly defined character (e.g. in

categorical matrices), and a "state" attribute that either holds a reference to an explicitly defined state, or a raw state value (a continuous value).

AbstractObs.superclass alias of Labelled

class AbstractObsMatrix (about=None, meta=None, row=None, set=None, valueOf=None)

The AbstractObsMatrix super class is the abstract type for a <matrix> element that contains rows which hold granular state observations.

AbstractObsMatrix.superclass alias of Annotated

 $\begin{array}{ll} \textbf{class AbstractObsRow} \ (about=None, & meta=None, & label=None, & id=None, & otu=None, \\ & cell=None, \ set=None, \ valueOf\_=None) \end{array}$ 

The AbstractObsRow represents a single row in a matrix. The row must refer to a previously declared otu element by its id attribute (and must have an id itself, may have a label, and may have meta attachments). The row contains multiple cell elements.

AbstractObsRow.superclass alias of TaxonLinked

 $\begin{array}{c} \textbf{class AbstractPolymorphicStateSet} \ (about=None, \quad meta=None, \quad label=None, \\ id=None, \quad symbol=None, \quad member=None, \quad un-to-symbol=None, \quad member=None, \quad un-to-symbol=None, \quad member=None, \quad un-to-symbol=None, \quad un-to-symbol=None$ 

certain\_state\_set=None, valueOf\_=None)

The AbstractPolymorphicStateSet type is the super-class for a polymorphic state set definition. The element has a required AbstractSymbol attribute that in restricted concrete subclasses must be of a sensible type such as a single IUPAC character. It may enclose zero or more AbstractMapping elements to resolve ambiguities.

AbstractPolymorphicStateSet.superclass alias of AbstractUncertainStateSet

 $\textbf{class AbstractRootEdge} \ (about=None, \ meta=None, \ label=None, \ id=None, \ length=None, \ length=None,$ 

*target=None*, *valueOf\_=None*)

The AbstractRootEdge complex type is a superclass for the edge that leads into a root, i.e. an edge with only a target attribute, but no source attribute. This type of edge is used for coalescent trees, where the initial lineage has a certain length before things start splitting up.

AbstractRootEdge.superclass alias of IDTagged

The AbstractSeqMatrix super class is the abstract type for a <matrix> element that contains rows which hold raw character sequences.

AbstractSeqMatrix.superclass alias of Annotated

 $\begin{array}{ll} \textbf{class AbstractSeqRow} \ (about=None, & meta=None, & label=None, & id=None, & otu=None, \\ & seq=None, \ valueOf\_=None) \end{array}$ 

The AbstractSeqRow represents a single row in a matrix. The row must refer to a previously declared otu element by its id attribute (and must have an id itself, may have a label, and may have meta attachments). The row contains a single seq element with raw character data.

AbstractSeqRow.superclass alias of TaxonLinked

**class AbstractSeqs** (about=None, meta=None, label=None, id=None, otus=None, format=None, matrix=None, valueOf =None)

The AbstractSeqBlock type is the superclass for character blocks that consist of raw character sequences.

AbstractSeqs.superclass

alias of AbstractBlock

class AbstractState (about=None, meta=None, label=None, id=None, symbol=None,  $valueOf\_=None$ )

The AbstractState type is the super-class for a state definition. The element has a required symbol attribute that in restricted concrete subclasses must be of a sensible type such as a single IUPAC character. It may enclose zero or more AbstractMapping elements to resolve ambiguities.

AbstractState.superclass alias of IDTagged

class AbstractStates (about=None, meta=None, label=None, id=None, state=None, polymorphic\_state\_set=None, uncertain\_state\_set=None, set=None, valueOf = None)

A container for a set of AbstractState elements.

AbstractStates.superclass alias of IDTagged

class AbstractTree (about=None, meta=None, label=None, id=None, node=None, rootedge=None, edge=None, set=None, valueOf\_=None)

The AbstractTree superclass is what a concrete tree inherits from.

AbstractTree.superclass alias of IDTagged

class AbstractTrees (about=None, meta=None, label=None, id=None, network=None,  $tree=None, set=None, valueOf\_=None$ )

The AbstractTrees superclass is what concrete trees inherit from.

AbstractTrees.superclass alias of IDTagged

 $\textbf{class AbstractUncertainStateSet} \ (about=None, \ meta=None, \ label=None, \ id=None, \\ symbol=None, member=None, valueOf = None)$ 

The AbstractUncertainStateSet type is the super-class for an uncertain state set definition. The element has a required AbstractSymbol attribute that in restricted concrete subclasses must be of a sensible type such as a single IUPAC character. It may enclose zero or more AbstractMapping

elements to resolve ambiguities.

AbstractUncertainStateSet.superclass alias of AbstractState

class Annotated (about=None, meta=None, valueOf\_=None)

The Annotated complexType is a super class for objects that optionally have metadata annotations of type Meta.

Annotated.superclass alias of Base

class Base (valueOf\_=None)

The base type for all complexType definitions in the nexml schema. This type allows a number of special attributes: xml:lang - for languages codes xml:base - see http://www.w3.org/TR/xmlbase/

xml:id - see http://www.w3.org/TR/xml-id/ xml:space - for whitespace handling xlink:href - for links Also see http://www.w3.org/2001/xml.xsd for more information on the xml and xlink attributes.

class ContinuousCells (about=None, meta=None, label=None, id=None, otus=None, format=None, matrix=None, valueOf=None)

A continuous characters block consisting of granular cells preceded by metadata.

ContinuousCells.superclass alias of AbstractCells

 ${\it class} \ {\it ContinuousChar} \ (about=None, \ meta=None, \ label=None, \ id=None, \ tokens=None, \\ states=None, \ codon=None, \ valueOf\_=None)$ 

A concrete implementation of the char element, which requires only an id attribute.

ContinuousChar.superclass alias of AbstractChar

The Continuous Format class is the container of continuous column definitions.

ContinuousFormat.superclass alias of AbstractFormat

class ContinuousMatrixObsRow (about=None, meta=None, label=None, id=None, otu=None, cell=None, set=None, valueOf\_=None)

This is a row in a matrix of continuous data as granular obervations.

ContinuousMatrixObsRow.superclass alias of AbstractObsRow

class ContinuousMatrixSeqRow (about=None, meta=None, label=None, id=None, otu=None, seq=None, valueOf\_=None)

This is a row in a matrix of continuous data as character sequences.

ContinuousMatrixSeqRow.superclass alias of AbstractSeqRow

class ContinuousObs (about=None, meta=None, label=None, char=None, state=None, valueOf=None)

This is a single cell in a matrix containing a continuous observation.

ContinuousObs.superclass alias of AbstractObs

 $\begin{array}{c} \textbf{class ContinuousObsMatrix} \ (about=None, \quad meta=None, \quad row=None, \quad set=None, \quad val-ueOf\_=None) \end{array}$ 

A matrix of rows with single character observations.

ContinuousObsMatrix.superclass alias of AbstractObsMatrix

A matrix of rows with seq strings of type continuous.

ContinuousSeqMatrix.superclass alias of AbstractSeqMatrix

class ContinuousSeqs (about=None, meta=None, label=None, id=None, otus=None, format=None, matrix=None, valueOf =None)

A continuous characters block consisting of float sequences preceded by metadata.

ContinuousSeqs.superclass alias of AbstractSeqs

**class DNAChar** (about=None, meta=None, label=None, id=None, tokens=None, states=None, codon=None, valueOf\_=None)

A concrete implementation of the AbstractChar element.

DNAChar.superclass

alias of AbstractChar

**class DNAFormat** (about=None, meta=None, states=None, char=None, set=None,  $valueOf\_=None$ )

The DNAFormat class is the container of DNA column definitions.

DNAFormat.superclass

alias of AbstractFormat

class DNAMapping (state=None, valueOf\_=None)

An IUPAC ambiguity mapping.

DNAMapping.superclass

alias of AbstractMapping

 $\begin{array}{lll} \textbf{class DNAMatrixObsRow} \ (about=None, & meta=None, & label=None, & id=None, & otu=None, \\ & cell=None, \ set=None, \ valueOf\_=None) \end{array}$ 

This is a row in a matrix of DNA data containing granular observations.

DNAMatrixObsRow.superclass

alias of AbstractObsRow

This is a row in a matrix of DNA data containing raw sequence data.

DNAMatrixSeqRow.superclass

alias of AbstractSeqRow

**class DNAObs** (about=None, meta=None, label=None, char=None, state=None, valueOf =None)

This is a single cell in a matrix containing a nucleotide observation.

DNAObs.superclass

alias of AbstractObs

class DNAObsMatrix (about=None, meta=None, row=None, set=None, valueOf\_=None)

A matrix of rows with single character observations.

DNAObsMatrix.superclass

alias of AbstractObsMatrix

 $\begin{tabular}{ll} {\bf class\ DNAPolymorphicStateSet}\ (about=None,\ meta=None,\ label=None,\ id=None,\ symbol=None,\ member=None,\ uncertain\_state\_set=None,\ none,\ no$ 

valueOf = None

The DNAPolymorphicStateSet type defines an IUPAC ambiguity mapping. It may enclose zero or more AbstractMapping elements to resolve ambiguities.

```
DNAPolymorphicStateSet.superclass alias of AbstractPolymorphicStateSet
```

 $\textbf{class DNASeqMatrix} \ (about=None, \ meta=None, \ row=None, \ set=None, \ valueOf\_=None)$ 

A matrix of rows with seq strings of type DNA.

DNASeqMatrix.superclass alias of AbstractSeqMatrix

 $\begin{array}{lll} \textbf{class DNAState} \ (about=None, & meta=None, & label=None, & id=None, & symbol=None, & valueOf\_=None) \end{array}$ 

This is a concrete implementation of the state element, which requires a symbol element, in this case restricted to DNAToken, i.e. a single IUPAC nucleotide symbol, and optional mapping elements to refer to other states.

DNAState.superclass

alias of AbstractState

 $\begin{array}{c} \textbf{class DNAStates} \ (about=None, \quad meta=None, \quad label=None, \quad id=None, \quad state=None, \quad poly-\\ morphic\_state\_set=None, \quad uncertain\_state\_set=None, \quad set=None, \quad val-\\ ueOf\_=None) \end{array}$ 

A container for a set of states.

 ${\tt DNAStates.superclass}$ 

alias of AbstractStates

 $\textbf{class DNAUncertainStateSet} \ (about=None, \ meta=None, \ label=None, \ id=None, \ symbol=None, \ member=None, \ valueOf\_=None)$ 

The DNAUncertainStateSet type defines an IUPAC ambiguity mapping. It may enclose zero or more AbstractMapping elements to resolve ambiguities.

DNAUncertainStateSet.superclass alias of AbstractUncertainStateSet

A DNA characters block consisting of granular cells preceded by metadata.

DnaCells.superclass
 alias of AbstractCells

A DNA characters block consisting of sequences preceded by metadata.

DnaSeqs.superclass alias of AbstractSeqs

A concrete network implementation, with floating point edge lengths.

FloatNetwork.superclass alias of AbstractNetwork

FloatTree.subclass alias of NexmlTree

```
FloatTree.superclass
```

alias of AbstractTree

**class IDTagged** (about=None, meta=None, label=None, id=None, valueOf\_=None)

The IDTagged complexType is a super class for objects that require unique id attributes of type xs:ID. The id must be unique within the XML document.

IDTagged.superclass

alias of Labelled

class IntNetwork (about=None, meta=None, label=None, id=None, node=None,

edge=None, set=None, valueOf\_=None)

A concrete network implementation, with integer edge lengths.

IntNetwork.superclass

alias of AbstractNetwork

class IntTree (about=None, meta=None, label=None, id=None, node=None, rootedge=None,

edge=None, set=None, valueOf\_=None)

A concrete tree implementation, with integer edge lengths.

IntTree.subclass

alias of NexmlTree

IntTree.superclass

alias of AbstractTree

**class** Labelled (about=None, meta=None, label=None, valueOf\_=None)

The Labelled complexType is a super class for objects that optionally have label attributes to use as a (non-unique) name of type xs:string.

Labelled.superclass

alias of Annotated

**class** LiteralMeta (datatype=None, content=None, property=None, valueOf =None)

Metadata annotations in which the object is a literal value. If the @content attribute is used, then the element should contain no children.

class NetworkFloatEdge (about=None, meta=None, label=None, id=None, source=None,

*length=None*, *target=None*, *valueOf\_=None*)

A concrete network edge implementation, with float edge.

 ${\tt NetworkFloatEdge.} \textbf{superclass}$ 

alias of AbstractEdge

class NetworkIntEdge (about=None, meta=None, label=None, id=None, source=None,

length=None, target=None, valueOf\_=None)

A concrete network edge implementation, with int edge.

NetworkIntEdge.superclass

alias of AbstractEdge

 ${\bf class\ NetworkNode}\ (about=None,\ meta=None,\ label=None,\ id=None,\ otu=None,\ root=False,$ 

*valueOf* =*None*)

A concrete network node implementation.

NetworkNode.superclass

alias of AbstractNode

class Nexm1 (\*args, \*\*kargs)

Creates a new nexml project.

```
Nexml.build_from_file (fname, index_otus=True)
```

Populate Nexml project with data in a nexml file.

The OptionalOTULinked complexType is a super class for objects that that optionally have an otu id reference.

OptionalTaxonLinked.superclass alias of IDTagged

class ProteinCells (about=None, meta=None, label=None, id=None, otus=None, format=None, matrix=None, valueOf\_=None)

An amino acid characters block consisting of granular cells preceded by metadata.

ProteinCells.superclass alias of AbstractCells

class ProteinSeqs (about=None, meta=None, label=None, id=None, otus=None, format=None, matrix=None, valueOf =None)

An amino acid characters block consisting of sequences preceded by metadata.

ProteinSeqs.superclass alias of AbstractSeqs

A concrete implementation of the AbstractChar element, i.e. a single column in an alignment.

RNAChar.superclass
alias of AbstractChar

class RNAFormat (about=None, meta=None, states=None, char=None, set=None, valueOf =None)

The RNAFormat class is the container of RNA column definitions.

RNAFormat.superclass alias of AbstractFormat

 ${\bf class} \ {\bf RNAMapping} \ ({\it state=None, valueOf}\_{=}None)$ 

An IUPAC RNA ambiguity mapping.

 $\begin{array}{c} {\tt RNAMapping.superclass}\\ {\tt alias\ of\ AbstractMapping} \end{array}$ 

 $\begin{array}{ll} \textbf{class RNAMatrixObsRow} \ (about=None, & meta=None, & label=None, & id=None, & otu=None, \\ & cell=None, \ set=None, \ valueOf\_=None) \end{array}$ 

This is a row in a matrix of RNA data containing granular observations.

RNAMatrixObsRow.superclass alias of AbstractObsRow

 $\begin{array}{ll} \textbf{class RNAMatrixSeqRow} \ (about=None, & meta=None, & label=None, & id=None, & otu=None, \\ seq=None, valueOf\_=None) \end{array}$ 

This is a row in a matrix of RNA data containing raw sequence data.

RNAMatrixSeqRow.superclass alias of AbstractSeqRow

**class RNAObs** (about=None, meta=None, label=None, char=None, state=None, valueOf =None)

This is a single cell in a matrix containing an RNA nucleotide observation.

RNAObs.superclass

alias of AbstractObs

**class RNAObsMatrix** (about=None, meta=None, row=None, set=None, valueOf\_=None) A matrix of rows with single character observations.

RNAObsMatrix.superclass

alias of AbstractObsMatrix

The RNAPolymorphicStateSet describes a single polymorphic IUPAC ambiguity mapping.

RNAPolymorphicStateSet.superclass alias of AbstractPolymorphicStateSet

**class RNASeqMatrix** (about=None, meta=None, row=None, set=None, valueOf\_=None) A matrix of rows with seq strings of type RNA.

RNASeqMatrix.superclass alias of AbstractSeqMatrix

 $\begin{array}{lll} \textbf{class RNAState} \ (about=None, & meta=None, & label=None, & id=None, & symbol=None, & valueOf\_=None) \end{array}$ 

This is a concrete implementation of the state element, which requires a symbol attribute, in this case restricted to RNAToken, i.e. a single IUPAC nucleotide symbol, and optional mapping elements to refer to other states.

RNAState.superclass

alias of AbstractState

 $\begin{array}{c} \textbf{class RNAStates} \ (about=None, \quad meta=None, \quad label=None, \quad id=None, \quad state=None, \quad poly-\\ morphic\_state\_set=None, \quad uncertain\_state\_set=None, \quad set=None, \quad val-\\ ueOf\_=None) \end{array}$ 

A container for a set of states.

 ${\tt RNAStates.superclass}$ 

alias of AbstractStates

class RNAUncertainStateSet (about=None, meta=None, label=None, id=None, symbol=None, member=None, valueOf =None)

The RNAUncertainStateSet describes a single uncertain IUPAC ambiguity mapping.

RNAUncertainStateSet.superclass alias of AbstractUncertainStateSet

 $\textbf{class} \ \texttt{ResourceMeta} \ (\textit{href=None}, \textit{rel=None}, \textit{meta=None}, \textit{valueOf}\_=\textit{None})$ 

Metadata annotations in which the object is a resource. If this element contains meta elements as children, then the object of this annotation is a "blank node".

class RestrictionCells (about=None, meta=None, label=None, id=None, otus=None, format=None, matrix=None, valueOf\_=None)

A standard characters block consisting of granular cells preceded by metadata.

```
RestrictionCells.superclass
```

alias of AbstractCells

A concrete implementation of the char element, which requires a unique identifier and a state set reference.

RestrictionChar.superclass

alias of AbstractChar

class RestrictionFormat (about=None, meta=None, states=None, char=None, set=None, valueOf $_=$ None)

The RestrictionFormat class is the container of restriction column definitions.

RestrictionFormat.superclass

alias of AbstractFormat

class RestrictionMatrixObsRow (about=None, meta=None, label=None, id=None, otu=None, cell=None, set=None, valueOf =None)

This is a row in a matrix of restriction site data as granular obervations.

RestrictionMatrixObsRow.superclass alias of AbstractObsRow

 $\begin{array}{ll} \textbf{class RestrictionMatrixSeqRow} \ (about=None, & meta=None, & label=None, & id=None, \\ otu=None, \ seq=None, \ valueOf\_=None) \end{array}$ 

This is a row in a matrix of restriction site data as character sequences.

RestrictionMatrixSeqRow.superclass alias of AbstractSeqRow

class RestrictionObs (about=None, meta=None, label=None, char=None, state=None, valueOf =None)

This is a single cell in a matrix containing a restriction site observation.

RestrictionObs.superclass alias of AbstractObs

class RestrictionObsMatrix (about=None, meta=None, row=None, set=None, valueOf=None)

A matrix of rows with single character observations.

RestrictionObsMatrix.superclass alias of AbstractObsMatrix

class RestrictionSeqMatrix (about=None, meta=None, row=None, set=None, valueOf\_=None)

A matrix of rows with seq strings of type restriction.

 $\label{eq:matrix.superclass} \textbf{alias of } \textbf{AbstractSeqMatrix}$ 

class RestrictionSeqs (about=None, meta=None, label=None, id=None, otus=None, format=None, matrix=None, valueOf =None)

A restriction site characters block consisting of sequences preceded by metadata.

RestrictionSeqs.superclass alias of AbstractSeqs

This is a concrete implementation of the state element, which requires a symbol element, in this case restricted to 0/1.

RestrictionState.superclass

alias of AbstractState

class RestrictionStates (about=None, meta=None, label=None, id=None, state=None, polymorphic\_state\_set=None, uncertain\_state\_set=None, set=None, valueOf\_=None)

A container for a set of states.

RestrictionStates.superclass

alias of AbstractStates

A RNA characters block consisting of granular cells preceded by metadata.

RnaCells.superclass

alias of AbstractCells

A RNA characters block consisting of sequences preceded by metadata.

RnaSeqs.superclass

alias of AbstractSegs

**class StandardCells** (about=None, meta=None, label=None, id=None, otus=None, format=None, matrix=None, valueOf =None)

A standard characters block consisting of granular cells preceded by metadata.

StandardCells.superclass

alias of AbstractCells

A concrete implementation of the char element, which requires a states attribute to refer to a set of defined states

StandardChar.superclass

alias of AbstractChar

**class StandardFormat** (about=None, meta=None, states=None, char=None, set=None, valueOf =None)

The StandardFormat class is the container of standard column definitions.

StandardFormat.superclass

alias of AbstractFormat

**class** StandardMapping (state=None, valueOf\_=None)

A standard character ambiguity mapping.

StandardMapping.superclass

alias of AbstractMapping

 ${\bf class} \ {\bf Standard Matrix Obs Row} \ (about=None, \quad meta=None, \quad label=None, \quad id=None, \quad id$ 

otu=None, cell=None, set=None, valueOf =None)

This is a row in a matrix of standard data as granular obervations.

label=None,

id=None,

meta=None,

```
StandardMatrixObsRow.superclass
          alias of AbstractObsRow
class StandardMatrixSeqRow (about=None,
                               otu=None, seq=None, valueOf_=None)
     This is a row in a matrix of standard data as character sequences.
```

StandardMatrixSeqRow.superclass alias of AbstractSeqRow

class StandardObs (about=None, meta=None, label=None, char=None, state=None, valueOf = None

This is a single cell in a matrix containing a standard observation.

StandardObs.superclass alias of AbstractObs

class StandardObsMatrix (about=None, meta=None. row=None. set=None. valueOf = None

A matrix of rows with single character observations.

StandardObsMatrix.superclass alias of AbstractObsMatrix

class StandardPolymorphicStateSet (about=None, label=None, meta=None, id=None, symbol=None, member=None, uncertain\_state\_set=None, valueOf\_=None)

The StandardPolymorphicStateSet type is a single polymorphic ambiguity mapping.

StandardPolymorphicStateSet.superclass alias of AbstractPolymorphicStateSet

class StandardSeqMatrix (about=None, meta=None,row=None, set=None, val $ueOf_=None$ 

A matrix of rows with seq strings of type standard.

StandardSeqMatrix.superclass alias of AbstractSeqMatrix

class StandardSeqs (about=None, meta=None, label=None, id=None, otus=None, format=None, matrix=None, valueOf =None)

A standard characters block consisting of sequences preceded by metadata.

StandardSeqs.superclass alias of AbstractSegs

class StandardState (about=None, meta=None, label=None, id=None, symbol=None, val $ueOf_=None$ 

This is a concrete implementation of the state element, which requires a symbol element, in this case restricted to integers, and optional mapping elements to refer to other states.

StandardState.superclass alias of AbstractState

class StandardStates (about=None, meta=None, label=None, id=None, state=None, polymorphic\_state\_set=None, uncertain\_state\_set=None, set=None, valueOf = None

A container for a set of states.

StandardStates.superclass alias of AbstractStates

 $\textbf{class StandardUncertainStateSet} \ (about=None, \ meta=None, \ label=None, \ id=None, \\ symbol=None, \ member=None, \ valueOf\_=None)$ 

The StandardUncertainStateSet type is a single uncertain ambiguity mapping.

StandardUncertainStateSet.superclass alias of AbstractUncertainStateSet

class TaxaLinked (about=None, meta=None, label=None, id=None, otus=None, valueOf\_=None)

The TaxaLinked complexType is a super class for objects that that require an otus id reference.

TaxaLinked.superclass

alias of IDTagged

The TaxonLinked complexType is a super class for objects that require a taxon id reference.

TaxonLinked.superclass

alias of IDTagged

A concrete edge implementation, with float length.

TreeFloatEdge.superclass alias of AbstractEdge

class TreeFloatRootEdge (about=None, meta=None, label=None, id=None, length=None, target=None, valueOf = None)

A concrete root edge implementation, with float length.

TreeFloatRootEdge.superclass
 alias of AbstractRootEdge

A concrete edge implementation, with int length.

TreeIntEdge.superclass alias of AbstractEdge

A concrete root edge implementation, with int length.

TreeIntRootEdge.superclass alias of AbstractRootEdge

**class TreeNode** (about=None, meta=None, label=None, id=None, otu=None, root=False, valueOf=None)

A concrete node implementation.

TreeNode.superclass

alias of AbstractNode

 $\begin{tabular}{ll} {\bf class\ Trees}\ (about=None,\ meta=None,\ label=None,\ id=None,\ otus=None,\ network=None,\ tree=None,\ set=None,\ valueOf\_=None) \end{tabular}$ 

A concrete container for tree objects.

Trees.superclass

alias of TaxaLinked

## class attrExtensions (valueOf\_=None)

This element is for use in WSDL 1.1 only. It does not apply to WSDL 2.0 documents. Use in WSDL 2.0 documents is invalid.

```
class Nexm1 (*args, **kargs)
```

Creates a new nexml project.

Nexml.build\_from\_file (fname, index\_otus=True)

Populate Nexml project with data in a nexml file.

class NexmlTree (newick=None, alignment=None, alg\_format='fasta', sp\_naming\_function=<function\_parse\_species at 0x4062050>, format=0)
Special PhyloTree object with nexml support

New in version 2.1.

# 3.6 Phyloxml Module

# 3.6.1 Phyloxml classes linked to ETE

```
class Phyloxml (*args, **kargs)
```

**class PhyloxmlTree** (phyloxml\_clade=None, phyloxml\_phylogeny=None, \*\*kargs) PhyloTree object supporting phyloXML format.

# 3.6.2 Generic Phyloxml classes

#### class Accession (source=None, valueOf\_=None)

Element Accession is used to capture the local part in a sequence identifier (e.g. 'P17304' in 'UniProtKB:P17304', in which case the 'source' attribute would be 'UniProtKB').

class Annotation (source=None, type\_=None, ref=None, evidence=None, desc=None, confidence=None, property=None, uri=None, valueOf\_=None)

The annotation of a molecular sequence. It is recommended to annotate by using the optional 'ref' attribute (some examples of acceptable values for the ref attribute: 'GO:0008270', 'KEGG:Tetrachloroethene degradation', 'EC:1.1.1.1'). Optional element 'desc' allows for a free text description. Optional element 'confidence' is used to state the type and value of support for a annotation. Similarly, optional attribute 'evidence' is used to describe the evidence for a annotation as free text (e.g. 'experimental'). Optional element 'property' allows for further, typed and referenced annotations from external resources.

class BinaryCharacters ( $lost\_count=None$ ,  $absent\_count=None$ ,  $present\_count=None$ ,  $type\_=None$ ,  $gained\_count=None$ , gained=None, lost=None, present=None, absent=None,  $valueOf\_=None$ )

The names and/or counts of binary characters present, gained, and lost at the root of a clade.

class BranchColor (red=None, green=None, blue=None, valueOf\_=None)

This indicates the color of a clade when rendered (the color applies to the whole clade unless overwritten by the color(s) of sub clades).

Element Clade is used in a recursive manner to describe the topology of a phylogenetic tree. The parent branch length of a clade can be described either with the 'branch\_length' element or the 'branch\_length' attribute (it is not recommended to use both at the same time, though). Usage of the 'branch\_length' attribute allows for a less verbose description. Element 'confidence' is used to indicate the support for a clade/parent branch. Element 'events' is used to describe such events as gene-duplications at the root node/parent branch of a clade. Element 'width' is the branch width for this clade (including parent branch). Both 'color' and 'width' elements apply for the whole clade unless overwritten in-sub clades. Attribute 'id\_source' is used to link other elements to a clade (on the xml-level).

This is used to express a typed relationship between two clades. For example it could be used to describe multiple parents of a clade.

```
class Confidence (type_=None, valueOf_=None)
```

A general purpose confidence element. For example this can be used to express the bootstrap support value of a clade (in which case the 'type' attribute is 'bootstrap').

class Date (unit=None, desc=None, value=None, minimum=None, maximum=None, valueOf =None)

A date associated with a clade/node. Its value can be numerical by using the 'value' element and/or free text with the 'desc' element' (e.g. 'Silurian'). If a numerical value is used, it is recommended to employ the 'unit' attribute to indicate the type of the numerical value (e.g. 'mya' for 'million years ago'). The elements 'minimum' and 'maximum' are used the indicate a range/confidence interval

## **class Distribution** (*desc=None*, *point=None*, *polygon=None*, *valueOf\_=None*)

The geographic distribution of the items of a clade (species, sequences), intended for phylogeographic applications. The location can be described either by free text in the 'desc' element and/or by the coordinates of one or more 'Points' (similar to the 'Point' element in Google's KML format) or by 'Polygons'.

# class DomainArchitecture (length=None, domain=None, valueOf\_=None)

This is used describe the domain architecture of a protein. Attribute 'length' is the total length of the protein

## **class Id** (*provider=None*, *valueOf\_=None*)

A general purpose identifier element. Allows to indicate the provider (or authority) of an identifier.

## class MolSeq (is\_aligned=None, valueOf\_=None)

Element 'mol\_seq' is used to store molecular sequences. The 'is\_aligned' attribute is used to indicated that this molecular sequence is aligned with all other sequences in the same phylogeny for which 'is aligned' is true as well (which, in most cases, means that gaps were introduced, and that all sequences for which 'is aligned' is true must have the same length).

Element Phylogeny is used to represent a phylogeny. The required attribute 'rooted' is used to indicate whether the phylogeny is rooted or not. The attribute 'rerootable' can be used to indicate that the phylogeny is not allowed to be rooted differently (i.e. because it is associated with root

dependent data, such as gene duplications). The attribute 'type' can be used to indicate the type of phylogeny (i.e. 'gene tree'). It is recommended to use the attribute 'branch\_length\_unit' if the phylogeny has branch lengths. Element clade is used in a recursive manner to describe the topology of a phylogenetic tree.

Phylogeny.subclass alias of PhyloxmlTree

class Point (geodetic\_datum=None, alt\_unit=None, lat=None, long=None, alt=None, valueOf\_=None)

The coordinates of a point with an optional altitude (used by element 'Distribution'). Required attributes are the 'geodetic\_datum' used to indicate the geodetic datum (also called 'map datum', for example Google's KML uses 'WGS84'). Attribute 'alt\_unit' is the unit for the altitude (e.g. 'meter').

class Polygon (point=None, valueOf\_=None)

A polygon defined by a list of 'Points' (used by element 'Distribution').

**class Property** (datatype=None, id\_ref=None, ref=None, applies\_to=None, unit=None, valueOf =None, mixedclass =None, content =None)

Property allows for typed and referenced properties from external resources to be attached to 'Phylogeny', 'Clade', and 'Annotation'. The value of a property is its mixed (free text) content. Attribute 'datatype' indicates the type of a property and is limited to xsd-datatypes (e.g. 'xsd:string', 'xsd:boolean', 'xsd:integer', 'xsd:decimal', 'xsd:float', 'xsd:double', 'xsd:date', 'xsd:anyURI'). Attribute 'applies\_to' indicates the item to which a property applies to (e.g. 'node' for the parent node of a clade, 'parent\_branch' for the parent branch of a clade). Attribute 'id\_ref' allows to attached a property specifically to one element (on the xml-level). Optional attribute 'unit' is used to indicate the unit of the property. An example: cproperty datatype="xsd:integer" ref="NOAA:depth" applies\_to="clade" unit="METRIC:m"> 200 /property>

class ProteinDomain (to=None, confidence=None, fromxx=None, id=None, valueOf =None)

To represent an individual domain in a domain architecture. The name/unique identifier is described via the 'id' attribute. 'confidence' can be used to store (i.e.) E-values.

class Reference (doi=None, desc=None, valueOf\_=None)

A literature reference for a clade. It is recommended to use the 'doi' attribute instead of the free text 'desc' element whenever possible.

Element Sequence is used to represent a molecular sequence (Protein, DNA, RNA) associated with a node. 'symbol' is a short (maximal ten characters) symbol of the sequence (e.g. 'ACTM') whereas 'name' is used for the full name (e.g. 'muscle Actin'). 'location' is used for the location of a sequence on a genome/chromosome. The actual sequence can be stored with the 'mol\_seq' element. Attribute 'type' is used to indicate the type of sequence ('dna', 'rna', or 'protein'). One intended use for 'id\_ref' is to link a sequence to a taxonomy (via the taxonomy's 'id\_source') in case of multiple sequences and taxonomies per node.

This is used to express a typed relationship between two sequences. For example it could be used to describe an orthology (in which case attribute 'type' is 'orthology').

Element Taxonomy is used to describe taxonomic information for a clade. Element 'code' is intended to store UniProt/Swiss-Prot style organism codes (e.g. 'APLCA' for the California sea hare 'Aplysia californica') or other styles of mnemonics (e.g. 'Aca'). Element 'authority' is used to keep the authority, such as 'J. G. Cooper, 1863', associated with the 'scientific\_name'. Element 'id' is used for a unique identifier of a taxon (for example '6500' with 'ncbi\_taxonomy' as 'provider' for the California sea hare). Attribute 'id\_source' is used to link other elements to a taxonomy (on the xml-level).

```
class Uri (type_=None, desc=None, valueOf_=None)
```

A uniform resource identifier. In general, this is expected to be an URL (for example, to link to an image on a website, in which case the 'type' attribute might be 'image' and 'desc' might be 'image of a California sea hare').

```
class PhyloxmlTree (phyloxml_clade=None, phyloxml_phylogeny=None, **kargs) PhyloTree object supporting phyloXML format.
```

# 3.7 Seggroup class

SeqGroup class can be used to store a set of sequences (aligned or not).

## **Parameters**

- **sequences** Path to the file containing the sequences or, alternatively, the text string containing the same information.
- **format** (*fasta*) the format in which sequences are encoded. Current supported formats are: fasta, phylip (phylip sequencial) and iphylip (phylip interleaved). Phylip format forces sequence names to a maximum of 10 chars. To avoid this effect, you can use the relaxed phylip format: phylip\_relaxed and iphylip\_relaxed.

```
msf = ">seq1\nAAAAAAAAA\n>seq2\nTTTTTTTT\n"
seqs = SeqGroup(msf, format="fasta")
print seqs.get_seq("seq1")

get_entries()
    Returns the list of entries currently stored.

get_seq(name)
    Returns the sequence associated to a given entry name.
```

```
iter_entries()
```

Returns an iterator over all sequences in the collection. Each item is a tuple with the sequence name, sequence, and sequence comments

```
set_seq (name, seq, comments=None)
Updates or adds a sequence
```

```
write (format='fasta', outfile=None)
```

Returns the text representation of the sequences in the supplied given format (default=FASTA). If "oufile" argument is used, the result is written into the given path.

New in version 2.1.

# 3.8 WebTreeApplication object

## class WebTreeApplication

Provides a basic WSGI application object which can handle ETE tree visualization and interactions. Please, see the webplugin example provided with the ETE installation package (http://pypi.python.org/pypi/ete2).

```
register_action (name, target, handler, checker, html_generator)
```

Adds a new web interactive function associated to tree nodes.

```
set_default_layout_fn (layout_fn)
```

Fix the layout function used to render the tree.

```
set_external_app_handler(handler)
```

Sets a custom function that will extend current WSGI application.

```
set_external_tree_renderer(handler)
```

If the tree needs to be processed every time is going to be drawn, the task can be delegated.

```
set_tree_loader (TreeConstructor)
```

Delegate tree constructor. It allows to customize the Tree class used to create new tree instances.

```
set tree size (w, h, units = px')
```

Fix the size of tree image

```
set_tree_style (handler)
```

Fix a TreeStyle instance to render tree images.

# 3.9 EvolTree class

Re-implementation of the standart TreeNode instance. It adds attributes and methods to work with phylogentic trees.

#### **Parameters**

- **newick** path to tree in newick format, can also be a string
- alignment path to alignment, can also be a string.
- **alg format** (*fasta*) alignment format.
- **sp\_naming\_function** function to infer species name.
- **format** type of newick format

• binpath – path to binaries, in case codeml or SLR are not in global path.

# change\_dist\_to\_evol (evol, model, fill=False)

change dist/branch length of the tree to a given evolutionary variable (dN, dS, w or bL), default is bL.

#### **Parameters**

- evol evolutionary variable
- model Model object from which to retrieve evolutionary variables
- **fill** (*False*) do not affects only dist parameter, each node will be annotated with all evolutionary variables (nodel.dN, node.w...).

# get\_descendant\_by\_node\_id(idname)

returns node list corresponding to a given idname

## get\_evol\_model(modelname)

returns one precomputed model

Parameters modelname – string of the name of a model object stored

Returns Model object

## get\_most\_likely(altn, null)

Returns pvalue of LRT between alternative model and null model.

usual comparison are:

Alternative	Null	Test	
M2	M1	PS on sites (M2 prone to	
		miss some sites) (Yang 2000)	
M3	M0	test of variability among sites	
M8	M7	PS on sites (Yang 2000)	
M8	M8a	RX on sites?? think so	
bsA	bsA1	PS on sites on specific branch	
		(Zhang 2005)	
bsA	M1	RX on sites on specific	
		branch (Zhang 2005)	
bsC	M1	different omegas on clades	
		branches sites ref: Yang	
		Nielsen 2002	
bsD	M3	different omegas on clades	
		branches sites (Yang Nielsen	
		2002, Bielawski 2004)	
b_free	b_neut		
		foreground branch not neutral (w != 1)	
		• RX if P<0.05	
		(means that w on	
		frg=1)	
		• PS if P>0.05 and	
		wfrg>1	
		• CN if P>0.05 and	
		wfrg>1	
		(Yang Nielsen 2002)	
b free			
b_free	M0	different ratio on branches	

## **Parameters**

- altn model with higher number of parameters (np)
- **null** model with lower number of parameters (np)

link\_to\_alignment (alignment, alg\_format='paml', nucleotides=True, \*\*kwargs)
same function as for phyloTree, but translate sequences if nucleotides nucleotidic sequence
is kept under node.nt\_sequence

## **Parameters**

- alignment (*True*) path to alignment or string
- alg\_format one of fasta phylip or paml
- alignment set to False in case we want to keep it untranslated

link\_to\_evol\_model (path, model)

# link EvolTree to evolutionary model

- free-branch model ('fb') will append evol values to tree
- Site models (M0, M1, M2, M7, M8) will give evol values by site and likelihood

3.9. EvolTree class

#### **Parameters**

- path path to outfile containing model computation result
- model either the name of a model, or a Model object (usually empty)

```
mark tree (node ids, verbose=False, **kargs)
```

function to mark branches on tree in order that paml could interpret it. takes a "marks" argument that should be a list of #1,#1,#2 e.g.:

```
t=Tree.mark_tree([2,3], marks=["#1","#2"])
```

#### **Parameters**

- **node\_ids** list of node ids (have a look to node.node\_id)
- verbose (False) warn if marks do not correspond to codeml standard
- **kargs** mainly for the marks key-word which needs a list of marks (marks=['#1', '#2'])

#### **Parameters**

- layout a layout function
- tree style (*None*) tree style object
- **Nonehistface** an histogram face function. This is only to plot selective pressure among sites

```
run_model (model_name, ctrl_string='', keep=True, **kwargs)
```

To compute evolutionnary models. e.g.: b\_free\_lala.vs.lele, will launch one free branch model, and store it in "WORK\_DIR/b\_free\_lala.vs.lele" directory

WARNING: this functionality needs to create a working directory in "rep"

WARNING: you need to have codeml and/or SLR in your path

The models available are:

Model name	Description	Model kind
M1	relaxation	site
M10	beta and gamma + 1	site
M11	beta and normal > 1	site
M12	0 and 2 normal > 2	site
M13	3 normal > 0	site
M2	positive-selection	site
M3	discrete	site
M4	frequencies	site
M5	gamma	site
M6	2 gamma	site
M7	relaxation	site
M8	positive-selection	site
M8a	relaxation	site
M9	beta and gamma	site
SLR	positive/negative selection	site
M0	negative-selection	null
fb_anc	free-ratios	branch_ancestor
bsA	positive-selection	branch-site
bsA1	relaxation	branch-site
bsB	positive-selection	branch-site
bsC	different-ratios	branch-site
bsD	different-ratios	branch-site
b_free	positive-selection	branch
b_neut	relaxation	branch
fb	free-ratios	branch

#### **Parameters**

- model\_name a string like "model-name[.some-secondary-name]" (e.g.: "fb.my\_first\_try", or just "fb") \* model-name is compulsory, is the name of the model (see table above for the full list) \* the second part is accessory, it is to avoid over-writing models with the same name.
- ctrl\_string list of parameters that can be used as control file.
- **kwargs** extra parameters should be one of: verbose, CodonFreq, ncatG, cleandata, fix\_blength, NSsites, fix\_omega, clock, seqfile, runmode, fix\_kappa, fix\_alpha, Small\_Diff, method, Malpha, aaDist, RateAncestor, outfile, icode, alpha, seqtype, omega, getSE, noisy, Mgene, kappa, treefile, model, ndata.

sep = 'n'

**show** (*layout=None*, *tree\_style=None*, *histfaces=None*)

call super show of PhyloTree histface should be a list of models to be displayes as histfaces

#### **Parameters**

- layout a layout function
- **tree\_style** (*None*) tree\_style object
- **Nonehistface** an histogram face function. This is only to plot selective pressure among sites

3.9. EvolTree class

```
write (features=None, outfile=None, format=10)
```

Inherits from Tree but adds the tenth format, that allows to display marks for CodeML. TODO: internal writting format need to be something like 0

Returns the newick representation of current node. Several arguments control the way in which extra data is shown for every node:

#### **Parameters**

- **features** a list of feature names to be exported using the Extended Newick Format (i.e. features=["name", "dist"]). Use an empty list to export all available features in each node (features=[])
- outfile writes the output to a given file
- **format** (10) defines the newick standard used to encode the tree. See tutorial for details.
- **format\_root\_node** (*False*) If True, it allows features and branch information from root node to be exported as a part of the newick text string. For newick compatibility reasons, this is False by default.
- is\_leaf\_fn See TreeNode.traverse() for documentation.

## **Example:**

```
t.get_newick(features=["species", "name"], format=1)
```

#### EvolTree

alias of EvolNode

# 3.10 NCBITaxa class

#### class NCBITaxa (dbfile=None)

Bases: object

versionadded: 2.3

Provides a local transparent connector to the NCBI taxonomy database.

Annotate a tree containing taxids as leaf names by adding the 'taxid', 'sci\_name', 'lineage', 'named\_lineage' and 'rank' additional attributes.

## **Parameters**

- $\mathbf{t}$  a Tree (or Tree derived) instance.
- taxid\_attr (name) Allows to set a custom node attribute containing

the taxid number associated to each node (i.e. species in PhyloTree instances).

Parameters tax2name,tax2track,tax2rank – Use these arguments to provide

pre-calculated dictionaries providing translation from taxid number and names,track lineages and ranks.

## get\_broken\_branches (t, taxa\_lineages, n2content=None)

Returns a list of NCBI lineage names that are not monophyletic in the provided tree, as well as the list of affected branches and their size.

## **CURRENTLY EXPERIMENTAL**

#### get common names (taxids)

 $\begin{tabular}{l} {\tt get\_descendant\_taxa} (parent, intermediate\_nodes=False, rank\_limit=None, collapse\_subspecies=False, return\_tree=False) \\ \end{tabular}$ 

given a parent taxid or scientific species name, returns a list of all its descendants taxids. If intermediate\_nodes is set to True, internal nodes will also be dumped.

## get\_fuzzy\_name\_translation(name, sim=0.9)

Given an inexact species name, returns the best match in the NCBI database of taxa names.

**Parameters sim** (0.9) – Min word similarity to report a match (from 0 to 1).

**Returns** taxid, species-name-match, match-score

### get lineage(taxid)

Given a valid taxid number, return its corresponding lineage track as a hierarchically sorted list of parent taxids.

## get\_name\_translator(names)

Given a list of taxid scientific names, returns a dictionary translating them into their corresponding taxids.

Exact name match is required for translation.

### get\_rank (taxids)

return a dictionary converting a list of taxids into their corresponding NCBI taxonomy rank

#### get\_taxid\_translator(taxids)

Given a list of taxids, returns a dictionary with their corresponding scientific names.

# get\_topology (taxids, intermediate\_nodes=False, rank\_limit=None, collapse subspecies=False, annotate=True)

Given a list of taxid numbers, return the minimal pruned NCBI taxonomy tree containing all of them.

## **Parameters intermediate\_nodes** (*False*) – If True, single child nodes

representing the complete lineage of leaf nodes are kept. Otherwise, the tree is pruned to contain the first common ancestor of each group.

**Parameters rank\_limit** (*None*) – If valid NCBI rank name is provided, the tree is pruned at that given level. For instance, use rank="species" to get rid of sub-species or strain leaf nodes.

**Parameters collapse\_subspecies** (*False*) – If True, any item under the species rank will be collapsed into the species upper node.

## translate\_to\_names (taxids)

Given a list of taxid numbers, returns another list with their corresponding scientific names.

## update\_taxonomy\_database(taxdump\_file=None)

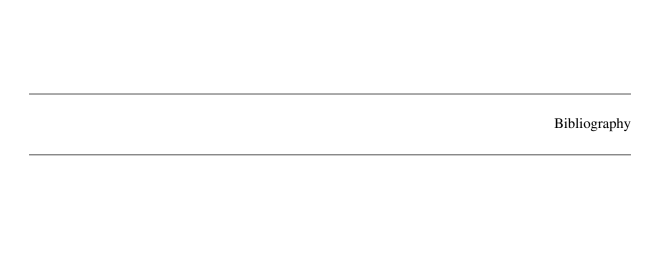
Updates the ncbi taxonomy database by downloading and parsing the latest taxdump.tar.gz file from the NCBI FTP site.

3.10. NCBITaxa class 167

**Parameters taxdump\_file** (*None*) – an alternative location of the taxdump.tax.gz file.

genindex

modindex



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[massingham2005] Massingham T. and Goldman N. Detecting amino acid sites under positive selection and purifying selection. Genetics 169: 1853-1762. (2005)

170 Bibliography

# е

```
ete2.13
ete2.clustering, 80
ete2.evol, 161
ete2.ncbi_taxonomy, 166
ete2.nexml, 141
ete2.parser.seqgroup, 160
ete2.phylo, 137
ete2.phyloxml, 157
ete2.treeview, 127
ete2.webplugin, 93
```

4	ete2.clustering (module), 80, 140
dd_child() (TreeNode method), 115	ete2.evol (module), 161
dd_face() (FaceContainer method), 129	ete2.ncbi_taxonomy (module), 166
dd_face() (TreeNode method), 116	ete2.nexml (module), 141
add_face_to_node() (in module ete2), 130	ete2.parser.seqgroup (module), 160
dd_feature() (TreeNode method), 116	ete2.phylo (module), 137
dd_features() (TreeNode method), 116	ete2.phyloxml (module), 157
dd_sister() (TreeNode method), 116	ete2.treeview (module), 127
nnotate_ncbi_taxa() (PhyloNode method), 137	ete2.webplugin (module), 93, 161
nnotate_tree() (NCBITaxa method), 166	EvolEvent (class in ete2.phylo), 140
AttrFace (class in ete2), 131	EvolNode (class in ete2), 161
	EvolTree (in module ete2), 166
3	expand_polytomies() (TreeNode method), 118
BarChartFace (class in ete2), 134 puild_from_file() (Nexml method), 142	F
ound_from_frie() (Nexim method), 142	Face (class in ete2), 130
	FaceContainer (class in ete2), 129
change_dist_to_evol() (EvolNode method), 162	
heck_monophyly() (TreeNode method), 116	G
hildren (TreeNode attribute), 116	get_age() (PhyloNode method), 138
CircleFace (class in ete2), 132	get_age_balanced_outgroup() (PhyloNode
ClusterNode (class in ete2), 140	method), 138
ClusterTree (in module ete2), 141	get_ancestors() (TreeNode method), 118
collapse_lineage_specific_expansions() (PhyloN-	get_ascii() (TreeNode method), 118
ode method), 138	get_broken_branches() (NCBITaxa method), 166
COLOR_SCHEMES (in module ete2), 136	get_cached_content() (TreeNode method), 118
compare() (TreeNode method), 117	get_children() (TreeNode method), 118
convert_to_ultrametric() (TreeNode method), 117	get_closest_leaf() (TreeNode method), 119
copy() (TreeNode method), 117	get_common_ancestor() (TreeNode method), 119
<b>1</b>	get_common_names() (NCBITaxa method), 167
<b>.</b>	get_descendant_by_node_id() (EvolNode
lel_feature() (TreeNode method), 117	method), 162
lelete() (TreeNode method), 117	get_descendant_evol_events() (PhyloNode
lescribe() (TreeNode method), 118	method), 138
letach() (TreeNode method), 118	get_descendant_taxa() (NCBITaxa method), 167
list (TreeNode attribute), 118	get_descendants() (TreeNode method), 119
DynamicItemFace (class in ete2), 135	get_distance() (TreeNode method), 119
≣	get_dunn() (ClusterNode method), 141
	get_edges() (TreeNode method), 119
ete2 (module), 13, 109	get_entries() (SeqGroup method), 160

get_evol_model() (EvolNode method), 1 get_farthest_leaf() (TreeNode method),		iter_species() (PhyloNode method), 139
get_farthest_node() (TreeNode method)		L
get_farthest_oldest_leaf() (PhyloNode		ladderize() (TreeNode method), 121
139		link_to_alignment() (EvolNode method), 163
get_farthest_oldest_node() (PhyloNode 139	method),	link_to_alignment() (PhyloNode method), 139 link_to_arraytable() (ClusterNode method), 141
get_fuzzy_name_translation() (2 method), 167	NCBITaxa	link_to_evol_model() (EvolNode method), 163
get_leaf_names() (TreeNode method), 1	20	M
get_leaf_profiles() (ClusterNode method), 1		mark_tree() (EvolNode method), 164
get_leaves() (TreeNode method), 120		N
get_leaves_by_name() (TreeNode method	od), 120	IN
get_lineage() (NCBITaxa method), 167		ncbi_compare() (PhyloNode method), 139
get_midpoint_outgroup() (TreeNode me	ethod), 120	NCBITaxa (class in ete2), 166
<pre>get_monophyletic() (TreeNode method)</pre>	, 120	Nexml (class in ete2), 142
get_most_likely() (EvolNode method), 1	162	NexmlTree (class in ete2), 142
get_my_evol_events() (PhyloNode meth		NodeStyle (class in ete2), 129
get_name_translator() (NCBITaxa meth-	-	Р
get_rank() (NCBITaxa method), 167		PhyloNode (class in ete2), 137
get_seq() (SeqGroup method), 160		PhyloTree (in module ete2), 137
get_silhouette() (ClusterNode method),	141	•
get_sisters() (TreeNode method), 120		Phyloxed (class in etc2), 157
get_speciation_trees() (PhyloNode meth	nod), 139	PhyloxmlTree (class in ete2), 157
get_species() (PhyloNode method), 139		PieChartFace (class in ete2), 134
get_taxid_translator() (NCBITaxa method		populate() (TreeNode method), 122
get_topology() (NCBITaxa method), 16		ProfileFace (class in ete2), 135
<pre>get_topology_id() (TreeNode method),</pre>	120	prune() (TreeNode method), 122
get_tree_root() (TreeNode method), 120	)	R
H		random_color() (in module ete2), 136
hoverEnterEvent() (	Sequence-	reconcile() (PhyloNode method), 139
Face.InteractiveLetterItem	method),	RectFace (class in ete2), 132
133	memoa),	register_action() (WebTreeApplication method),
	Sequence-	161
Face.InteractiveLetterItem	method),	remove_child() (TreeNode method), 124
133	memou),	remove_sister() (TreeNode method), 124
133		render() (EvolNode method), 164
		render() (TreeNode method), 124
img_style (TreeNode attribute), 121		resolve_polytomy() (TreeNode method), 124
ImgFace (class in ete2), 132		robinson_foulds() (TreeNode method), 124
is_leaf() (TreeNode method), 121		run_model() (EvolNode method), 164
is_root() (TreeNode method), 121		run_moder() (Evon tode method), 104
iter_ancestors() (TreeNode method), 12	1	S
iter_descendants() (TreeNode method),		search_nodes() (TreeNode method), 125
iter_edges() (TreeNode method), 121	121	sep (EvolNode attribute), 165
iter_entries() (SeqGroup method), 160		SeqGroup (class in ete2), 160
iter_leaf_names() (TreeNode method), 1	21	SeqMotifFace (class in ete2), 133
iter_leaf_profiles() (ClusterNode method), iter_leaf_profiles() (ClusterNode method)		SequenceFace (class in ete2), 132
iter_leaves() (TreeNode method), 121	u), 141	
iter_prepostorder() (TreeNode method),	121	SequenceFace.InteractiveLetterItem (class in ete2), 133
iter_search_nodes() (TreeNode method)		0.02), 155
noi_scarch_nodes() (Treefrode inclied)	, 141	

174 Index

```
set_default_layout_fn()
                           (WebTreeApplication
        method), 161
set distance function() (ClusterNode method),
set_external_app_handler() (WebTreeApplication
        method), 161
set external tree renderer() (WebTreeApplication
        method), 161
set_outgroup() (TreeNode method), 125
set_seq() (SeqGroup method), 160
set_species_naming_function()
                                    (PhyloNode
        method), 139
set_style() (TreeNode method), 125
set_tree_loader() (WebTreeApplication method),
        161
set_tree_size() (WebTreeApplication method), 161
set_tree_style() (WebTreeApplication method),
        161
show() (EvolNode method), 165
show() (TreeNode method), 125
sort_descendants() (TreeNode method), 125
species (PhyloNode attribute), 140
split_by_dups() (PhyloNode method), 140
StackedBarFace (class in ete2), 132
standardize() (TreeNode method), 125
StaticItemFace (class in ete2), 135
support (TreeNode attribute), 126
SVG_COLORS (in module ete2), 135
swap_children() (TreeNode method), 126
Т
TextFace (class in ete2), 131
translate_to_names() (NCBITaxa method), 167
traverse() (TreeNode method), 126
Tree (in module ete2), 126
TreeFace (class in ete2), 135
TreeNode (class in ete2), 115
TreeStyle (class in ete2), 127
U
unroot() (TreeNode method), 126
up (TreeNode attribute), 126
update_taxonomy_database()
                                    (NCBITaxa
        method), 167
W
WebTreeApplication (class in ete2), 161
write() (EvolNode method), 166
write() (SegGroup method), 160
write() (TreeNode method), 126
```

Index 175