# Package 'glottoTrees'

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abridge\_labels

Shorten labels to a glottocode substring

# Description

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Shortens tip and node labels to a glottocode substring within them. If any labels lack a glottocode substring, a warning is given.

# Usage

abridge\_labels(phy)

# **Arguments**

phy

A phylo or multiPhylo object, containing one or more trees to manipulate.

# **Details**

Glottocodes comprise four lowercase letters (or b10b or 3adt) followed by four numbers, and are only identified if they are initial in the string or are preceded by <.

Also recognizes and retains duplicate suffixes, i.e., a hyphen followed by one or more numerals at the end of the string (see apply\_duplicate\_suffixes).

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#### Value

A phylo or multiPhylo object, the manipulated tree(s).

### **Examples**

```
library(ape)
tree <- get_glottolog_trees("Koreanic")</pre>
plot(tree)
nodelabels(tree$node.label)
tree2 <- abridge_labels(tree)</pre>
plot(tree2)
nodelabels(tree2$node.label)
# Retain duplicate suffixes:
tree3 <- clone_tip(tree, "Jollado<chol1278>", n = 2, subgroup = TRUE)
tree3a <- apply_duplicate_suffixes(tree3)</pre>
plot(tree3a)
nodelabels(tree3a$node.label)
tree4 <- abridge_labels(tree3a)</pre>
plot(tree4)
nodelabels(tree4$node.label)
\ensuremath{\text{\#}}\xspace A warning is issued if any label does not contain a glottocode
supertree <- assemble_supertree() # contains nodes without glottocodes</pre>
supertree2 <- abridge_labels(assemble_supertree())</pre>
# Applied to a `multiPhylo` object:
trees <- get_glottolog_trees(c("Kartvelian", "Basque"))</pre>
trees2 <- abridge_labels(trees)</pre>
plot(trees[[1]])
nodelabels(trees[[1]]$node.label)
plot(trees2[[1]])
nodelabels(trees2[[1]]$node.label)
plot(trees[[2]])
nodelabels(trees[[2]]$node.label)
plot(trees2[[2]])
nodelabels(trees2[[2]]$node.label)
```

add\_tip

Add tips to a tree

# Description

Add one or more tips below a parent node specified by its label.

# Usage

```
add_tip(phy, label, parent_label)
```

# Arguments

phy A phylo object. The tree to manipulate. label A character vector containing tip labels.

parent\_label A character string containing the label of the parent node.

#### **Details**

The length of the branches, between the added tips and their parent node is set equal to the longest of the original branches directly below node n.

#### Value

A phylo object containing the modified tree.

# **Examples**

```
library(ape)

tree <- abridge_labels(get_glottolog_trees("LeftMay"))
tree <- ultrametricize(rescale_branches_exp(tree))
plot_glotto(tree)

# Attach one or more new tips to a tree:
tree2 <- add_tip(tree, label = "rockypeak", parent_label = "iter1240")
plot_glotto(tree2)
tree3 <- add_tip(tree, label = c("bo", "kaumifi"), parent_label = "bopa1235")
plot_glotto(tree3)

# Move tips by using remove_tip() and add_tip():
tree4 <- remove_tip(tree, "amap1240")
tree4a <- add_tip(tree4, "amap1240", parent_label = "left1242")
plot_glotto(tree4a)</pre>
```

```
apply_duplicate_suffixes
```

Apply duplicate suffixes to tips and nodes

# **Description**

Suffixes are applied to ensure tip labels and node labels are not duplicates. Suffixes have the form -1, -2, -3, ...

#### **Usage**

```
apply_duplicate_suffixes(phy)
```

# **Arguments**

phy

A phylo object, the tree whose labels are to have copy suffixes applied.

#### Details

The function recognizes existing duplicate suffixes and deals with them in one of two ways. If a label has n duplicates that are already suffixed -1, -2, ... -n, then the suffixes are not changed. Under any other conditions, old suffixes are removed and new ones applied.

Suffixation of tips and of nodes are handled independently of one another.

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#### Value

A phylo object, the same tree but with suffixes applied to the labels.

### **Examples**

```
library(ape)
tree <- abridge_labels(get_glottolog_trees("Koreanic"))</pre>
plot_glotto(tree)
tree2 <- clone_tip(tree, "chol1278", n = 2, subgroup = TRUE)</pre>
plot_glotto(tree2)
# Technically, tree2 is ill-formed because it has duplicate tip labels.
# Note how this causes problems if we try to clone one of them, since it
# is unclear which should be cloned:
## Not run:
tree2a <- clone_tip(tree2, "chol1278")</pre>
## End(Not run)
# Suffixation of duplicate tips
tree3 <- apply_duplicate_suffixes(tree2)</pre>
plot_glotto(tree3)
# Once they are suffixed, these tips can be cloned successfully:
tree4 <- clone_tip(tree3, c("chol1278-2", "chol1278-3"), subgroup = TRUE)</pre>
plot_glotto(tree4)
# Suffixing is applied across all tips that share a glottocode, and
# separately, across all nodes that share a glottocode:
tree5 <- apply_duplicate_suffixes(tree4)</pre>
plot_glotto(tree5)
```

assemble\_rake

Bind trees as a high-level rake

# **Description**

Takes a multiPhylo object containing multiple trees and combines them into a single tree with a rake structure at its root, below which each tree appears on its own branch.

### Usage

```
assemble_rake(phy)
```

# **Arguments**

phy

A multiphylo object containing the trees to be combined.

# Value

A phylo object, a single tree.

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#### **Examples**

```
library(ape)
arnhem_hypothesis <-
    c("Gunwinyguan", "Mangarrayi-Maran", "Maningrida",
        "Kungarakany", "Gaagudju")
trees <- get_glottolog_trees(arnhem_hypothesis)
simple_rake <- assemble_rake(trees)
rake_in_rake <- assemble_rake(c(assemble_rake(trees[1:3]), trees[4:5]))
plot(simple_rake)
plot(rake_in_rake)

# If `phy` contains only one tree, a warning is issued.
mono_rake <- assemble_rake(trees[3])
plot(mono_rake)</pre>
```

assemble\_supertree

Create a glottolog super-tree

# Description

Combining glottolog family trees into one large tree. Families can be assembled directly below a rake structure at the root, or can be grouped, so that the root first branches into groups, and the families then branch out below the group nodes.

## Usage

```
assemble_supertree(macro_groups, glottolog_version)
```

### Arguments

 ${\tt macro\_groups}$ 

A list of character vectors, in which each vector contains the names of one or more macroareas which define a group. Alternatively, setting macro\_groups to NULL causes the tree to be assembled without groups.

glottolog\_version

A character string, specifying which glottolog version to use. Currently available options are '4.0' through to '4.8' and '5.0'. If no value is specified then the newest available version is used.

#### **Details**

Grouping is controlled by the macro\_groups parameter. Groups can comprise a single glottolog macroarea, or multiple macroareas. Current macroareas are Africa, Australia, Eurasia, North America, Papunesia and South America. Setting macro\_groups to NULL causes the tree to be assembled without groups.

```
# Supertree whose first order branches are the glottolog macroareas
supertree <- assemble_supertree()
supertree_v.4.3 <- assemble_supertree(glottolog_version = "4.3")</pre>
```

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clone\_tip

Clone tips

## **Description**

Clones tips as sisters of the original. Optionally, places the new clones and the original in their own subgroup, in which case the node for the new subgroup is assigned the same label as the original tip.

# Usage

```
clone_tip(phy, label, n = 1, subgroup = FALSE)
```

# **Arguments**

phy A phylo object. The tree to manipulate.

label A character vector containing tip labels.

n A numeric vector. The number of clones to make.

subgroup A logical. Whether to create a subgroup containing the new clones and their

original.

## Value

A phylo object containing the modified tree.

```
library(ape)

tree <-
    rescale_branches_exp(abridge_labels(get_glottolog_trees("Tangkic")))
plot_glotto(tree)
tree2 <- clone_tip(tree, "nyan1300")
plot_glotto(tree2)

tree3 <- clone_tip(tree, "nyan1300", subgroup = TRUE)
plot_glotto(tree3)

# Add suffixes to labels, to keep all labels distinct
tree3a <- apply_duplicate_suffixes(tree3)</pre>
```

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```
plot_glotto(tree3a)

tree4 <- clone_tip(tree, "lard1243", n = 3)
plot_glotto(tree4)

tree5 <- clone_tip(tree, "lard1243", n = 3, subgroup = TRUE)
plot_glotto(tree5)

tree6 <- clone_tip(tree, c("lard1243", "nyan1300"), n = 2, subgroup = TRUE)
plot_glotto(tree6)
tree6a <- apply_duplicate_suffixes(tree6)
plot_glotto(tree6a)

## Not run:
# Returns error if any element of `label` is not in `phy`
tree7 <- clone_tip(tree, c("lard1243", "xxxxx1234"))

## End(Not run)</pre>
```

collapse\_node

Collapse one or more nodes rootwards

# **Description**

Collapse one or more nodes rootwards

# Usage

```
collapse_node(phy, label)
```

# Arguments

phy A phylo object. The tree to manipulate.

label A character vector containing node labels.

# Value

A phylo object containing the modified tree.

```
library(ape)
tree <-
    rescale_branches_exp(abridge_labels(get_glottolog_trees("Tangkic")))
plot_glotto(tree)
tree2 <- collapse_node(tree, "gang1267")
plot_glotto(tree2)</pre>
```

convert\_to\_tip 9

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Convert nodes to tips

# **Description**

For nodes identified by their label, remove the clade that they dominate and replace it with a tip of the same name.

# Usage

```
convert_to_tip(phy, label, warn = TRUE)
```

# **Arguments**

phy A phylo object. The tree to manipulate.

label A character vector containing node or tip labels.

warn A logical, whether to issue warning when label cantains tip labels.

# **Details**

Any labels included in label which are both node labels and tip labels are regarded as node label, and that node will be removed and replaced by a tip.

Any labels included in label which are tip labels only are ignored.

#### Value

A phylo object containing the modified tree.

## **Examples**

```
tree1 <- abridge_labels(get_glottolog_trees("GreatAndamanese"))
plot_glotto(tree1)
tree2 <- convert_to_tip(tree1, label = c("okol1242", "sout2683"))
plot_glotto(tree2)</pre>
```

 ${\tt extract\_glottocode}$ 

Extract glottocode substrings

# **Description**

From a character vector, extracts the first glottocode from each element.

# Usage

```
extract_glottocode(label)
```

## **Arguments**

label

A string

#### **Details**

Glottocodes comprise four lowercase letters (or b10b or 3adt) followed by four numbers, and are only identified if they are initial in the string or are preceded by <.

Also recognizes and retains duplicate suffixes, i.e., a hyphen followed by one or more numerals at the end of the string (see apply\_duplicate\_suffixes).

# Value

A string

### **Examples**

```
extract_glottocode("DongoKresh<dong1296>-l-")
extract_glottocode(c("DongoKresh<dong1296>-l-", "Goro-Golo<orlo1238>"))
# Duplicate suffixes are recognised and retained
extract_glottocode(c("Goro-Golo<orlo1238>-l", "Goro-Golo<orlo1238>-2"))
```

```
get_glottolog_families
```

Simple family metadata

# Description

Returns a dataframe of metadata on glottolog's language families.

# Usage

```
get_glottolog_families(glottolog_version)
```

# **Arguments**

```
glottolog_version
```

A character string, specifying which glottolog version to use. Currently available options are '4.0' through to '4.8' and '5.0'. If no value is specified then the newest available version is used.

### **Details**

Returned columns are: tree, tree\_name, n\_tips, n\_nodes and main\_macroarea.

```
head(get_glottolog_families())
head(get_glottolog_families(glottolog_version = "4.3"))
```

```
get_glottolog_languages
```

Simple language metadata

### **Description**

Returns a dataframe of metadata on glottolog's languages.

# Usage

```
get_glottolog_languages(glottolog_version)
```

# **Arguments**

glottolog\_version

A character string, specifying which glottolog version to use. Currently available options are '4.0' through to '4.8' and '5.0'. If no value is specified then the newest available version is used.

#### **Details**

Returned columns are: glottocode, isocodes, name, name\_in\_tree, position, tree and tree\_name.

# **Examples**

```
head(get_glottolog_languages())
head(get_glottolog_languages(glottolog_version = "4.3"))
```

```
get_glottolog_phylo_geo
```

Extended glottolog metadata

# **Description**

Returns a dataframe of glottolog geographical and phylogenetic metadata.

# Usage

```
{\tt get\_glottolog\_phylo\_geo(glottolog\_version)}
```

# Arguments

```
{\tt glottolog\_version}
```

A character string, specifying which glottolog version to use. Currently available options are '4.0' through to '4.8' and '5.0'. If no value is specified then the newest available version is used.

# **Details**

Returned columns are: glottocode, isocodes, name, level, vertex\_type, vertex\_label, vertex\_name, macroarea, latitude, longitude, family\_glottocode, family\_name and tree.

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#### **Examples**

```
head(get_glottolog_phylo_geo())
head(get_glottolog_phylo_geo(glottolog_version = "4.3"))
```

```
get_glottolog_trees Glottolog trees by version
```

# **Description**

Returns a multiPhylo object containing all, or a requested subset, of the glottolog trees.

## Usage

```
get_glottolog_trees(family, glottolog_version)
```

# **Arguments**

family

A character vector. Elements are names of glottolog families whose trees are to be returned. If family is left unspecified, all trees are returned.

glottolog\_version

A character string, specifying which glottolog version to use. Currently available options are '4.0' through to '4.8' and '5.0'. If no value is specified then the newest available version is used.

#### **Details**

By default, trees are returned from the most recent version of glottolog. Alternatively, an older version of glottolog can be specified.

## Value

A phylo object containing one glottolog tree, or a multiPhylo object containing multiple glottolog trees.

```
library(ape)
tree_totonacan <- get_glottolog_trees(family = "Totonacan")
tree_totonacan_v4.3 <- get_glottolog_trees("Totonacan", "4.3")
plot(tree_totonacan)
plot(tree_totonacan_v4.3)
trees <- get_glottolog_trees(family = c("Caddoan", "Tangkic"))
plot(trees[[1]])
plot(trees[[2]])</pre>
```

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glottolog\_geography Geographical data from glottolog, versions 4.0 - 5.0

# **Description**

Datasets of geographical information about the languages in glottolog.

# Usage

```
glottolog_geography_v4.0 glottolog_geography_v4.1 glottolog_geography_v4.2 glottolog_geography_v4.3 glottolog_geography_v4.4 glottolog_geography_v4.5 glottolog_geography_v4.6 glottolog_geography_v4.7 glottolog_geography_v4.8 glottolog_geography_v4.8 glottolog_geography_v5.0
```

## **Format**

A dataframe:

glottocode glottocode of the lect
name name of the lect
isocodes the ISO-639-3 code of the lect
level "language" or "dialect"
macroarea glottolog's geographical macroarea
latitude
longitude
v4.0: A dataframe of 20,049 rows

v4.0: A dataframe of 20,049 rows v4.1: A dataframe of 20,290 rows v4.2: A dataframe of 20,752 rows v4.3: A dataframe of 20,930 rows v4.4: A dataframe of 21,329 rows v4.5: A dataframe of 21,508 rows v4.6: A dataframe of 21,652 rows 14 glottolog\_trees

```
v4.7: A dataframe of 21,746 rows
v4.8: A dataframe of 21,957 rows
v5.0: A dataframe of 22,111 rows
```

#### Source

https://glottolog.org/meta/downloads

glottolog\_trees

Trees from glottolog, versions 4.0 - 5.0

### **Description**

multiPhylo objects, which provide a representation of the phylogenetic relationships of the languages in glottolog.

# Usage

```
glottolog_trees_v4.0
glottolog_trees_v4.1
glottolog_trees_v4.2
glottolog_trees_v4.3
glottolog_trees_v4.4
glottolog_trees_v4.5
glottolog_trees_v4.6
glottolog_trees_v4.7
glottolog_trees_v4.8
glottolog_trees_v4.8
glottolog_trees_v4.8
```

# **Format**

```
v4.0: A multiPhylo object of 421 trees v4.1: A multiPhylo object of 421 trees v4.2: A multiPhylo object of 422 trees v4.3: A multiPhylo object of 418 trees v4.4: A multiPhylo object of 420 trees v4.5: A multiPhylo object of 420 trees v4.6: A multiPhylo object of 420 trees v4.7: A multiPhylo object of 422 trees v4.8: A multiPhylo object of 422 trees v4.8: A multiPhylo object of 422 trees v5.0: A multiPhylo object of 423 trees
```

keep\_as\_tip

#### **Source**

https://glottolog.org/meta/downloads

keep\_as\_tip

Keep tips and convert nodes to tips

# **Description**

Nominate which tips and nodes are to be kept as tips. Others tips are removed. If any element of label is both a node label and tip label, it will be treated as referring to the tip, not the node.

# Usage

```
keep_as_tip(phy, label)
```

# Arguments

phy A phylo object. The tree to manipulate.

label A character vector containing tip and node labels.

### Value

A phylo object containing the modified tree.

# Examples

```
library(ape)

tree <- abridge_labels(get_glottolog_trees("Tangkic"))
plot_glotto(tree)

tree2 <- keep_as_tip(tree, c("lard1243", "kaya1319", "nyan1300", "gang1267"))
plot_glotto(tree2)</pre>
```

keep\_tip

Keep tips

## **Description**

Nominate which tips are to be kept. Others are removed.

# Usage

```
keep_tip(phy, label)
```

# Arguments

phy A phylo object. The tree to manipulate.

label A character vector containing tip labels.

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#### Value

A phylo object containing the modified tree.

# **Examples**

```
library(ape)

tree <- abridge_labels(get_glottolog_trees("Tangkic"))
plot_glotto(tree)

tree2 <- keep_tip(tree, c("lard1243", "kang1283", "kaya1319"))
plot_glotto(tree2)</pre>
```

move\_node

Move a node

# **Description**

Move one node to a position dominated by a new parent node.

# Usage

```
move_node(phy, label, parent_label)
```

# **Arguments**

phy A phylo object. The tree to manipulate.

label A character string containing the label of the node to move.

parent\_label A character string containing the label of the new parent node.

#### **Details**

The branch length above the moved node remains unchanged.

If moving the node would result in any other node(s) having no descendant tips, then those other nodes are removed.

# Value

A phylo object containing the modified tree.

```
library(ape)

tree <- abridge_labels(get_glottolog_trees("LeftMay"))
tree <- ultrametricize(rescale_branches_exp(tree))
plot_glotto(tree)

tree2 <- move_node(tree, "iter1240", parent_label = "left1242")
plot_glotto(tree2)</pre>
```

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move_tip	Move a tip
----------	------------

# **Description**

Move one tip to a new parent node.

# Usage

```
move_tip(phy, label, parent_label)
```

# Arguments

phy A phylo object. The tree to manipulate.

label A character string containing the tip label.

parent\_label A character string containing the label of the parent node.

# **Details**

In the tip's new position, the length of the branch above it is the same as the longest branch above any new sister of the tip.

If moving the tip would result in any node(s) having no descendant tips, then those nodes are removed.

## Value

A phylo object containing the modified tree.

# **Examples**

```
library(ape)

tree <- abridge_labels(get_glottolog_trees("LeftMay"))
tree <- ultrametricize(rescale_branches_exp(tree))
plot_glotto(tree)

tree2 <- move_tip(tree, "amap1240", parent_label = "left1242")
plot_glotto(tree2)</pre>
```

nonbranching\_nodes

Find nodes with a single child

# Description

Find nodes with a single child

## Usage

```
nonbranching_nodes(phy)
```

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## **Arguments**

phy A phylo object

### Value

A vector of node labels

plot\_glotto

Plot a tree in downward, linguistic style

# Description

An attempt is made to choose a reasonable plot width, height and offset of the labels from the tree's tips. If the choices are not satisfactory, manipulate them using positive or negative values of extra\_width, extra\_height and extra\_offset.

# Usage

```
plot_glotto(
  phy,
  nodelabels = TRUE,
  extra_width = 0,
  extra_height = 0,
  extra_offset = 0,
  srt = 0,
  adj = NULL,
  ...
)
```

# Arguments

phy A phylo object. nodelabels A logical, whether to plot node labels. A numeric, extra width to add beyond the default. extra\_width extra\_height A numeric, extra height to add beyond the default. extra\_offset A numeric, extra offset to add beyond the default. A numeric giving how much the labels are rotated in degrees. srt A numeric specifying the justification of the text strings of the labels: 0 (leftadj justification), 0.5 (centering), or 1 (right-justification). Additional arguments passed to plot(). . . .

# **Details**

By default, tip labels will run vertically, except in the case that all tip labels are a maximum of one character long, in which case they are oriented horizontally.

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# **Examples**

```
library(ape)

tree <- abridge_labels(get_glottolog_trees("Tangkic"))
plot_glotto(tree)
tree2 <- rescale_deepest_branches(tree, 7)
plot_glotto(tree2)
plot_glotto(tree2, srt = 90, adj = 0.5)</pre>
```

relabel\_node

Change node labels

# **Description**

Replaces existing tip labels with new labels.

# Usage

```
relabel_node(phy, label, new_label)
```

# Arguments

phy A phylo object.

label A vector of strings, the node labels to be replaced.

new\_label A vector of strings, the same length as label. The corresponding replacement

labels.

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relabel_tip
-------------

Change tip labels

# **Description**

Replaces existing tip labels with new labels.

# Usage

```
relabel_tip(phy, label, new_label)
```

## **Arguments**

phy A phylo object.

label A vector of strings, the tip labels to be replaced.

new\_label A vector of strings, the same length as label. The corresponding replacement

labels.

# **Examples**

 $relabel\_with\_names$ 

Change labels from glottocodes to names

# Description

Looks up glottolog language names corresponding to glottocodes and replaces tip and node labels, which contain glottocodes, with the appropriate names.

# Usage

```
relabel_with_names(phy, glottolog_version)
```

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#### **Arguments**

```
phy A phylo object, the tree to manipulate. glottolog_version
```

A character string, specifying which glottolog version to use. Currently available options are '4.0' through to '4.8' and '5.0'. If no value is specified then the newest available version is used.

## **Details**

Also recognizes and retains duplicate suffixes, i.e., a hyphen followed by one or more numerals at the end of the string (see apply\_duplicate\_suffixes).

Labels without glottocodes are left unchanged and a warning is given. The version of glottolog to use for look-up can be controlled with glottolog\_version.

### Value

A phylo object, the manipulated tree.

#### **Examples**

```
library(ape)
# Replace full glottolog labels with names
tree <- get_glottolog_trees("Kresh-Aja")</pre>
plot(tree)
nodelabels(tree$node.label)
tree2 <- relabel_with_names(tree)</pre>
plot(tree2)
nodelabels(tree2$node.label)
# Replace abridged labels with names
tree3 <- abridge_labels(tree)</pre>
plot(tree3)
nodelabels(tree3$node.label)
tree4 <- relabel_with_names(tree3)</pre>
plot(tree4)
nodelabels(tree4$node.label)
# Use names from earlier glottolog version:
tree5 <- relabel_with_names(tree, glottolog_version = "4.3")</pre>
plot(tree5)
nodelabels(tree5$node.label)
```

remove\_clade

Remove clades

## **Description**

From a tree, remove clacdes, identified by the labels of their deepest node.

## Usage

```
remove_clade(phy, label)
```

remove\_tip

# **Arguments**

phy A phylo object. The tree to manipulate.

label A character vector containing node labels.

# Value

A phylo object containing the modified tree.

 $remove\_tip$ 

Remove tips

# Description

From a tree, remove tips identified by their node labels.

# Usage

```
remove_tip(phy, label)
```

# Arguments

phy A phylo object. The tree to manipulate.

label A character vector containing tip labels.

## Value

A phylo object containing the modified tree.

```
library(ape)

tree <- abridge_labels(get_glottolog_trees("Tangkic"))
plot_glotto(tree)

tree2 <- remove_tip(tree, c("kang1283", "kaya1319"))
plot_glotto(tree2)</pre>
```

rescale\_branches 23

rescale\_branches

Set all branch length to 1

# **Description**

Sets all branch lengths in a tree to the same length.

# Usage

```
rescale_branches(phy, length = 1)
```

# **Arguments**

phy A phylo object, the tree to manipulate.

length A numeric stating the branch length.

# **Details**

By default, sets all branch lengths to 1.

# Value

A phylo object, the manipulated tree.

# **Examples**

```
library(ape)
tree <- abridge_labels(get_glottolog_trees("Tangkic"))
tree2 <- clone_tip(tree, "nyan1300", n = 2, subgroup = TRUE)
plot_glotto(tree2)
tree3 <- rescale_branches(tree2)
plot_glotto(tree3)</pre>
```

rescale\_branches\_constrained

Exponentialize branch lengths using constraints

# Description

Sets the branches furthest from the root to shortest\_length and then scales other branches exponentially in such a way that the total tree has a height of tree\_height.

# Usage

```
rescale_branches_constrained(phy, tree_height, shortest_length)
```

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## **Arguments**

phy A phylo object, the tree to manipulate.

 ${\tt tree\_height} \qquad \text{A positive numeric, the desired height of the whole tree.}$ 

shortest\_length

A positive numeric, the desired length of the shortest branches, i.e., the branches farthest from the root.

#### Value

A phylo object, the manipulated tree.

# **Examples**

```
library(ape)
tree <- abridge_labels(get_glottolog_trees("Siouan"))
plot_glotto(tree)
tree2 <- rescale_branches_constrained(tree, tree_height = 2000, shortest_length = 200)
plot_glotto(tree2)
tree3 <- rescale_branches_constrained(tree, tree_height = 2000, shortest_length = 100)
plot_glotto(tree3)</pre>
```

# **Description**

Sets the deepest branches to length 1/2, the next deepest to 1/4, the next to 1/8, etc., all multiplied by the parameter length.

# Usage

```
rescale_branches_exp(phy, length = 1)
```

# **Arguments**

phy A phylo object, the tree to manipulate.

length A positive numeric, a multiplier for the exponential branch lengths 1/2, 1/4,

1/8...

## Value

A phylo object, the manipulated tree.

```
library(ape)
tree <- abridge_labels(get_glottolog_trees("Siouan"))
plot_glotto(tree)
tree2 <- rescale_branches_exp(tree)
plot_glotto(tree2)</pre>
```

```
rescale_deepest_branches
```

Set length of deepest branches

# **Description**

Sets lengths of branches immediately below the root to the same, user-specified length.

### Usage

```
rescale_deepest_branches(phy, length = 1)
```

# **Arguments**

phy A phylo object, the tree to manipulate. length A numeric stating the branch length.

#### Value

A phylo object, the manipulated tree.

# **Examples**

```
library(ape)
arnhem_hypothesis <-
    c("Gunwinyguan", "Mangarrayi-Maran", "Maningrida",
        "Kungarakany", "Gaagudju")
tree <- assemble_rake(abridge_labels(get_glottolog_trees(arnhem_hypothesis)))
plot_glotto(tree)
# tree now contains five language families. All branch lengths are 1.
# Set the deepest branch lengths to 5, implying a great genealogical
# distance between the families within the tree.
tree2 <- rescale_deepest_branches(tree, length = 5)
plot_glotto(tree2)</pre>
```

ultrametricize

Ultrametricize tree by stretching final edges

# Description

Alters branches ending in a tip in such a way that all tips are equidistant from the root. Does this by lengthening branches above all but the existing, most-distance tip(s).

# Usage

```
ultrametricize(phy)
```

## **Arguments**

phy

A phylo object, the tree to manipulate.

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#### Value

A phylo object, the manipulated tree.

#### **Examples**

```
library(ape)
tree <- rescale_branches_exp(abridge_labels(get_glottolog_trees("Siouan")))
plot_glotto(tree)
tree2 <- ultrametricize(tree)
plot_glotto(tree2)</pre>
```

which\_tree

Tree numbers of glottolog families

# Description

Returns the tree number of one or more glottolog families.

## Usage

```
which_tree(family, glottolog_version)
```

### **Arguments**

family

A character vector. Elements are names of glottolog families whose trees are to be returned. If family is left unspecified, all trees are returned.

glottolog\_version

A character string, specifying which glottolog version to use. Currently available options are '4.0' through to '4.8' and '5.0'. If no value is specified then the newest available version is used.

## Value

A named vector of integers, giving the tree numbers and the family names as the vector names.

```
which_tree("Caddoan")
which_tree(c("Caddoan", "Tangkic"), glottolog_version = "4.3")
# If some family names are unrecognized, a warning is issued
which_tree(c("Caddoan", "Zzz"), glottolog_version = "4.4")
## Not run:
# If no family names are recognized, an error results
which_tree()
which_tree("Zzz")
## End(Not run)
```

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