Package 'phytools'

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Title Phylogenetic Tools for Comparative Biology (and Other Things)

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Depends R (>= 2.10), ape (>= 3.0-10), maps

Imports animation, clusterGeneration, graphics, grDevices, methods, mnormt, msm, numDeriv, phangorn (>= 2.0.4), plotrix, scatterplot3d, stats, utils

Suggests geiger, rgl

ZipData no

Description Package contains various functions for phylogenetic analysis. This functionality is concentrated in the phylogenetic analysis of comparative data from species. For example, the package includes functions for Bayesian and ML ancestral state estimation; visual simulation of trait evolution; fitting models of trait evolution with multiple Brownian rates and correlations; visualizing discrete and continuous character evolution using colors or projections into trait space; identifying the location of a change in the rate of character evolution on the tree; fast Brownian motion simulation and simulation under several other models of continuous trait evolution; fitting a model of correlated binary trait evolution; locating the position of a fossil or an recently extinct lineage on a tree using continuous character data with ML; plotting lineage accumulation through time, including across multiple trees (such as a Bayesian posterior sample); conducting an analysis called stochastic character mapping, in which character histories for a discrete trait are sampled from their posterior probability distribution under a model; conducting a multiple (i.e., partial) Mantel test; fitting a phylogenetic regression model with error in predictor and response variables; conducting a phylogenetic principal components analysis, a phylogenetic regression, a reduced major axis regression, a phylogenetic

canonical correlation analysis, and a phylogenetic ANOVA; projecting a tree onto a geographic map; simulating discrete character histories on the tree; and fitting a model in which a discrete character evolves under the threshold model. In addition to this phylogenetic comparative method functionality, the package also contains functions for a wide range of other purposes in phylogenetic biology. For instance, functionality in this package includes (but is not restricted to): adding taxa to a tree (including randomly, everywhere, or automatically to genera); generating all bi- and multi-furcating trees for a set of taxa; reducing a phylogeny to its backbone tree; dropping tips or adding tips to special types of phylogenetic trees; exporting a tree as an XML file; converting a tree with a mapped character to a tree with singleton nodes and one character state per edge; estimating a phylogeny using the least squares method; simulating birth-death trees under a range of conditions; rerooting trees; a wide range of visualizations of trees; and a variety of other manipulations and analyses that phylogenetic biologists may find useful for their research.

License GPL (>= 2)

URL http://github.com/liamrevell/phytools

Repository CRAN

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NeedsCompilation no

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phytools-package	phytools: things)	Phylogenetic	Tools for	comparative	biology	(and	other
	inings)						

Description

phytools provides functions for phylogenetic comparative biology; as well as several other functions for tree inference, manipulation, and analysis that are not implemented in other R packages.

The complete list of functions can be displayed with library(help = phytools).

More information on **phytools** can be found at http://www.phytools.org or http://blog.phytools.org.

Author(s)

Liam J. Revell

Maintainer: Liam J. Revell < liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

add.arrow

Add an arrow pointing to a tip or node on the tree

Description

This function adds an arrow to a plotted tree.

Usage

```
add.arrow(tree=NULL, tip, ...)
```

Arguments

tree	an object of class "phylo", "contMap", or "densityMap". If not supplied, the function will obtain the last plotted phylogeny from the environmental variable last_plot.phylo.
tip	label of tip or tip or node number. If tree=NULL then the tip or node number must be supplied.
	optional arguments to control the shape and size of the arrow including: its length (arrl) in the units of the plot; the length of the arrowhead (hedl); the total angle between the wings in the arrowhead (angle); the line width for the plotted lines (lwd); the offset from the tip or end of tip label, in character widths (offset); and the color (col).

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Details

This function presently works for radial (type="fan") and right facing square phylograms (type="phylogram"). Trees can be plotted using phytools function plotTree, plotSimmap, contMap, densityMap, and ape method plot.phylo.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

nodelabels

add.color.bar

Add color bar to a plot

Description

This function adds a color bar to a plot created by plotBranchbyTrait. A color bar can be added by clicking on a location within the plot (when prompt=TRUE) or by setting prompt=FALSE and supplying x & y coordinates for the object. This function is also used internally by plot.contMap and plot.densityMap.

Usage

```
add.color.bar(leg, cols, title=NULL, lims=c(0,1), digits=1, prompt=TRUE, lwd=4, outline=TRUE, ...)
```

Arguments

leg	numerical value for the length of the legend.
cols	colors for the legend.
title	text to plot above the bar.
lims	range for the bar.
digits	digits for plotted numbers.
prompt	logical value indicating whether the location of the legend should be obtained interactively.
lwd	width of the plotted bar.
outline	logical value indicated whether or not to outline the plotted color bar with a 1 pt line.

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... optional arguments including: x x-coordinate of the legend (if prompt=FALSE); y y-coordinate of the legend; subtitle optional legend subtitle; direction direction of the color bar (i.e., increase from left to right or from right to left).

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2013) Two new graphical methods for mapping trait evolution on phylogenies. *Methods in Ecology and Evolution*, **4**, 754-759.

See Also

contMap, densityMap, plotBranchbyTrait

add.everywhere

Add tip to all edges in a tree

Description

This function adds a tip to all branches to the tree and returns a a list of trees as an object of class "multiPhylo".

Usage

```
add.everywhere(tree, tip.name)
```

Arguments

tree an object of class "phylo".

tip.name a string containing the name of the tip to add.

Value

A list of trees as an object of class "multiPhylo". Since the tip can be added to any branch, the length of the list is equal to the number of edges in tree.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

add.random 9

See Also

allFurcTrees, exhaustiveMP

Description

This function adds new tips at random to a tree with branch lengths. If no edge lengths are provided, and the tree is ultrametric, then edge lengths are assigned to keep the tree ultrametric. The probability that at new tip is added along any branch is directly proportional to the length of the branch.

Usage

```
add.random(tree, n=NULL, tips=NULL, edge.length=NULL, order=c("random","input"))
```

Arguments

tree an object of class "phylo".

n a number of tips to add to the tree. If NULL, will use length(tips).
tips a set of tip names for the added tips. If NULL, names will be supplied

edge.length terminal edge length for the added tips. If NULL, and is.ultrametric(tree) == TRUE,

then edge lengths will be assigned to keep the tree ultrametric. Note that if edge lengths are assigned and n>1, then the assigned terminal edge lengths are not guaranteed as subsequent random tip addition could occur along the new termi-

nal edge.

order addition order for the new tips.

Details

Note that sometimes the resultant tree plotted with plot.phylo or plotSimmap may display with branches crossing. If so, the tree can be 'untangled' using untangle.

Value

An object of class "phylo".

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

10 add.simmap.legend

See Also

allFurcTrees, add.everywhere

add.simmap.legend

Add legend to stochastically mapped tree

Description

This function adds a legend (by default, interactively) to a plotted stochastic character mapped tree.

Usage

```
add.simmap.legend(leg=NULL, colors, prompt=TRUE, vertical=TRUE, ...)
```

Arguments

leg	states for the discrete character in the order of colors.
colors	colors for the legend in the order of leg, or, if leg=NULL, named vector of colors in which names(colors are the states of the mapped discrete character.
prompt	logical value indicating whether the location of the legend should be obtained interactively (i.e., by clicking in the plotting area).
vertical	logical value indiciating whether to plot the legend vertically (if TRUE) or horizontally.
• • •	optional arguments including: x x-coordinate of the legend (if prompt=FALSE); y y-coordinate of the legend; and shape which can be shape="square", the default, or shape="circle".

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

plotSimmap

add.species.to.genus 11

 ${\it add.} {\it species.to.} {\it genus} \ \ {\it a dd species to genus on a phylogeny or bind simulated species subtrees to a backbone genus tree}$

Description

add. species.to.genus adds an additional species to a genus on a phylogeny. genus.to.species.tree simulates pure-birth subtrees and then binds them at a random height along the terminal edge leading to each corresponding genus on a genus-level backbone tree.

Usage

```
add.species.to.genus(tree, species, genus=NULL, where=c("root","random"))
genus.to.species.tree(tree, species)
```

Arguments

tree	an object of class "phylo". In the case of genus.to.species.tree this should be a genus-level backbone tree.
species	string contain species name in the format "Genus_species" or "Genus species".
genus	for add.species.to.genus, optional argument containing the genus to which species is to be attached. If NULL then genus will be extracted from species.
where	for add. species.to.genus, the location to attach species to the tree. where="root" will cause the species to be attached to the MRCA of all members of genus. where="random" will cause species to be attached at random to the subtree descended from the MRCA of all members of genus.

Details

For add. species.to.genus, if genus contains only one species and where="root", then species will be attached midway along the branch leading to the one species. If where="random" then species will be added at a random position along the edge. If genus cannot be found in the tree, then the original tree is returned and a warning printed. If the tree is not ultrametric, then the resultant tree may not contain branch lengths and a warning will be printed. If genus is non-monophyletic then species will be attached to the most inclusive group containing members of genus and a warning will be printed.

Value

An object of class "phylo".

Author(s)

Liam Revell liam.revell@umb.edu>

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References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

```
add.random, bind.tip
```

allFurcTrees

Generate all bi- and multifurcating unrooted trees

Description

This function creates all possible unrooted bi- and multifurcating trees and returns a list of trees as an object of class "multiPhylo".

Usage

```
allFurcTrees(n, tip.label=NULL, to.plot=TRUE)
```

Arguments

n an integer giving the desired number of species.

tip.label an optional vector of length n containing the tip names.

to.plot an optional logical value indicating whether or not to plot the trees.

Details

This function should be used with caution for n greater than about 8, as in this case the number of possible trees is extremely large.

Value

A list of trees as an object of class "multiPhylo".

Author(s)

Liam Revell liam.revell@umb.edu>

References

Felsenstein, J. 2004. Inferring Phylogenies. Sinauer.

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

```
add.everywhere, exhaustiveMP
```

anc.Bayes 13

anc.Bayes	Bayesian ancestral character estimation	

Description

This function uses Bayesian MCMC to sample from the posterior distribution for the states at internal nodes in the tree.

Usage

```
anc.Bayes(tree, x, ngen=10000, control=list())
```

Arguments

tree an object of class "phylo".

x a vector of tip values for species; names(x) should be the species names.

ngen a integer indicating the number of generations for the MCMC.

control a list of control parameters containing the following elements: sig2: starting

value for σ^2 (BM rate); a: starting for the state at the root node; y: starting values for the states at all internal nodes excluding the root (should be labeled with node numbers); pr.mean: means for the prior distributions in the following order - sig2, a, y, note that the prior probability distribution is exponential for sig2 and normal for a and y; pr.var: variances on the prior distributions, same order as pr.mean (but the variance is not used for sig2); prop: variances on the normal proposal distributions in the same order as pr.mean; sample: sample

frequency from the MCMC.

Value

A matrix with number of rows ngen/sample+1 containing the posterior sample and likelihoods. Matrix columns are labeled either sig2 or by the node number of the internal node.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

```
ace, anc.ML, anc.trend, evol.rate.mcmc, fastAnc
```

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Examples

```
tree<-pbtree(n=50)
x<-fastBM(tree,sig2=2) # simulate using fastBM
X<-anc.Bayes(tree,x,ngen=10000) # sample ancestral states
estimates<-colMeans(X[21:nrow(X),]) # get estimates, excluding burnin</pre>
```

anc.ML

Ancestral character estimation using likelihood

Description

This function estimates the evolutionary parameters and ancestral states for Brownian evolution using likelihood. It is also possible (for model="BM") to allow for missing data for some tip taxa.

Usage

```
anc.ML(tree, x, maxit=2000, model=c("BM","OU"), ...)
```

Arguments

tree an object of class "phylo".

x a vector of tip values for species; names(x) should be the species names.

maxit an optional integer value indicating the maximum number of iterations for optimization.

model model of continuous character evolution ont he tree. It's possible that only model="BM" works in the present version as model="OU" has not be thoroughly tested & some bugs were reported for an earlier version.

... other arguments.

Details

Because this function relies on a high dimensional numerical optimization of the likelihood function, fastAnc should probably be preferred for most purposes. If using anc.ML, users should be cautious to ensure convergence. This has been ameliorated in phytools>=0.2-48 by seeding the ML optimization with the result from fastAnc.

Value

A list with the following components:

sig2 the variance of the BM process.
ace a vector with the ancestral states.

logLik the log-likelihood.

convergence the value of \$convergence returned by optim() (0 is good).

anc.trend 15

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

Schluter, D., Price, T., Mooers, A. O., and Ludwig, D. (1997) Likelihood of ancestor states in adaptive radiation. *Evolution* **51**, 1699-1711.

See Also

```
ace, anc. Bayes, fastAnc, optim
```

Examples

```
tree<-pbtree(n=50)
x<-fastBM(tree) # simulate using fastBM
anc.ML(tree,x) # fit model & estimate ancestral states</pre>
```

anc.trend

Ancestral character estimation with a trend

Description

This function estimates the evolutionary parameters and ancestral states for Brownian evolution with directional trend.

Usage

```
anc.trend(tree, x, maxit=2000)
```

Arguments

tree an object of class "phylo".

x a vector of tip values for species; names(x) should be the species names.

maxit an optional integer value indicating the maximum number of iterations for opti-

mization.

Details

Note that this will generally only work and produce sensible results for a phylogeny with some non-contemporary tips (i.e., a tree with some fossil species). The function uses optim with method="L-BFGS-B"; however optimization is only constrained for the sig2 which must be >0.

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Value

A list with the following components:

ace a vector with the ancestral states.

mu a trend parameter per unit time.

sig2 the variance of the BM process.

logL the log-likelihood.

convergence the value of \$convergence returned by optim() (0 is good).

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

```
ace, anc.Bayes, anc.ML, optim
```

Examples

```
\label{tree-rel} $$ tree < -rtree (20) $$ x < -fastBM(tree,mu=2) $$ \# simulate using fastBM with a trend (m!=0) $$ anc.trend(tree,x) $$ # fit model $$ estimate ancestral states $$
```

ancThresh Ancestral character estimation under the threshold model using Bayesian MCMC

Description

This function uses Bayesian MCMC to estimate ancestral states and thresholds for a discrete character under the threshold model from quantitative genetics (Felsenstein 2012).

Usage

```
ancThresh(tree, x, ngen=1000, sequence=NULL, method="mcmc", model=c("BM","OU","lambda"), control=list(), \dots)
```

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Arguments

tree phylogenetic tree.

x a named vector containing discrete character states; or a matrix containing the

tip species, in rows, and probabilities of being in each state, in columns.

ngen number of generations to run the MCMC.

sequence assumed ordering of the discrete character state. If not supplied and x is a vector

then numerical/alphabetical order is assumed; if not supplied and x is a matrix,

then the column order of x is used.

method only method currently available is "mcmc".

model model for the evolution of the liability. Options are "BM" (Brownian motion, the

default), "OU" (Ornstein-Uhlenbeck), or "lambda" (the lambda model).

control list containing the following elements: sample, the sampling interval; propliab

variance of the proposal distribution for liabilities; propthresh variance on the proposal distribution for the thresholds; propalpha variance on the proposal distribution for alpha (for model="OU"); pr.anc prior probability distribution on the ancestral states for each node, in a matrix - not all nodes need to be supplied; pr.th prior density on the thresholds; burnin number of generations to exclude for burn-in when plotting posterior probabilities on the tree; plot logical value indicating whether or not to plot the posterior probabilities; print logical value indicating whether or not to print the state of the MCMC; piecol colors for the posterior probabilities plotted as pie charts at internal nodes; and tipcol which indicates whether the tip colors should be based on the input data ("input") or sampled tip liabilities ("estimated"). These will only differ if

there is uncertainty in the tip states.

... additional arguments to be passed to plotThresh (called internally).

Value

This function returns as list with four elements:

ace posterior probabilities for each character state at each internal node, with the

burn-in excluded.

mcmc full posterior sample for the states.

par full posterior sample for the thresholds, the alpha parameter of the "OU" model

(if applicable), and the likelihood.

liab full posterior sample of the liabilities at internal and tip nodes.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Felsenstein, J. (2012) A comparative method for both discrete and continuous characters using the threshold model. *American Naturalist*, **179**, 145-156.

Revell, L. J. (2014) Ancestral character estimation under the threshold model from quantitative genetics. *Evolution*, bold68, 743-759.

applyBranchLengths

See Also

anc.Bayes, threshBayes

anoletree

Phylogeny of Greater Antillean anole ecomorph species with mapped discrete character

Description

A phylogeny of Greater Antillean anole species with a mapped discrete character - 'ecomorph class.' Data and tree are from Mahler et al. (2010).

Usage

data(anoletree)

Format

The data are stored as a modified object of class "simmap" with a mapped discrete character. (E.g., see read.simmap.)

Source

Mahler, D. L, L. J. Revell, R. E. Glor, and J. B. Losos. (2010) Ecological opportunity and the rate of morphological evolution in the diversification of Greater Antillean anoles. *Evolution*, **64**, 2731-2745.

applyBranchLengths

Applies the branch lengths of a reference tree to a target

Description

This function applies the set of branch lengths from a reference tree to a target tree while reconciling any mappings (as in read.simmap) with the new branch lengths.

Usage

```
applyBranchLengths(tree, edge.length)
```

Arguments

tree target tree.

edge.length number of digits for rounding. Passed to round.

as.multiPhylo

Value

A tree with branch lengths, or modified "phylo" object with a mapped discrete character.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

as.multiPhylo

Conversion to object of class "multiPhylo"

Description

This function converts between object classes.

Usage

```
as.multiPhylo(x, ...)
## S3 method for class 'multiSimmap'
as.multiPhylo(x, ...)
```

Arguments

```
x object to be converted to "multiPhylo". Presently an object of class "multiSimmap".... optional arguments.
```

Value

An object of class "multiPhylo".

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

20 ave.rates

tes Average the posterior rates
·

Description

Primarily used internally by posterior.evolrate.

Usage

```
ave.rates(tree, shift, tips, sig1, sig2, ave.shift, showTree=TRUE)
```

Arguments

tree	a tree.
shift	the shift point for this sample.
tips	tip names tipward of shift.
sig1	rate 1.
sig2	rate 2.
ave.shift	average shift from all samples.
showTree	logical value indicating whether to plot the rate-stretched tree.

Value

A list of the rates.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

```
\verb|evol.rate.mcmc|, \verb|minSplit|, \verb|posterior.evolrate||\\
```

averageTree 21

|--|

Description

These functions compute average trees or consensus trees by various criteria.

Usage

```
averageTree(trees, start=NULL, method="quadratic.path.difference",
tol=1e-12, quiet=FALSE, ...)
ls.consensus(trees, start=NULL, tol=1e-12, quiet=FALSE, ...)
minTreeDist(tree, trees, method="quadratic.path.difference", ...)
```

Arguments

trees	object of class "multiPhylo".
tree	object of class "phylo". For minTreeDist the tree on which to find the edge lengths that minimize the distance to the phylogenies in trees.
start	starting tree for optimization.
method	distance criterion for minimization. Options are "symmetric.difference", "branch.score.difference", "path.difference", and "quadratic.path.difference".
tol	tolerance value for optimization.
quiet	logical value indicating whether to run "quietly" or not.
	other arguments to be passed internally.

Value

An object of class "phylo" with edge lengths.

Author(s)

Liam Revell liam.revell@umb.edu>

bind.tip	Attaches a new tip to a tree	

Description

Functions adds a new tip to the tree. If the tree is ultrametric and no branch length is specified, then edge.length is scaled so that the tree remains ultrametric after the new tip is added.

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Usage

```
bind.tip(tree, tip.label, edge.length=NULL, where=NULL, position=0, interactive=FALSE, \dots)
```

Arguments

tree receptor tree.

tip.label a string containing the species name for the new tip.

edge.length edge length for the new tip (a scalar).

where node number to attach new tip. If position>0 then then tip will be attached

below the specified node. Node numbers can also be tips, in which case the new tip will be added along the terminal edge. To find out the tip number for given

species with name "species" type: which(tree\$tip.label=="species".

position distance *below* node to add tip.

interactive logical value indicating whether or not the species should be added interactively.

(Defaults to FALSE.)

... arguments to be passed to plotTree (for interactive=TRUE.)

Details

Wrapper function for 'ape' bind.tree. Note that interactive=TRUE works only for right-facing phylograms.

Value

An object of class "phylo".

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

bind.tree.simmap

Attaches a new tip to a tree

Description

This function grafts tree y onto tree x at node where.

Usage

```
bind.tree.simmap(x, y, where="root")
```

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Arguments

```
x an object of class "simmap". (The receptor tree.)
y an object of class "simmap". (The tree being grafted.)
where node number to attach new tip, or the root node if where="root".
```

Details

This function wraps around bind.tree for objects of class "simmap"; however it presently only allows y to be grafted at a node of x and it does not allow y to possess a root edge.

Value

```
An object of class "simmap".
```

Author(s)

Liam Revell < liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

bmPlot	Simulates and visualizes discrete-time Brownian evolution on a phylogeny

Description

This function conducts discrete-time Brownian motion simulation on an input tree, plots the outcome, and returns the tip and internal node states to the user as a named vector. The function will first rescale and round the branch lengths to integer length, if they are not already in integer values. If integer branch lengths are provided, the user should also set ngen=max(nodeHeights(tree)). For type="threshold" the visualization is of the threshold model (Felsenstein 2012), in which the evolving character is liability and the segments of evolution are colored by their value for the threshold trait. If type="threshold" is used, the function requires at least one addition input: thresholds, a vector containing the ordered thresholds between states. The user can also provide the colors for plotting in colors. Note that one more color than threshold should be provided as one threshold implies two states; two thresholds, three states; etc. If no value for colors is provided, the function will recycle a set of four colors up to the number of times required by thresholds. Finally, the optional argument return.tree=TRUE will tell the function to return a list with the tip and note states and an object of class "phylo" with (for type="threshold"), the state for the threshold model through time mapped on the branches of the tree in discrete time.

Usage

```
bmPlot(tree, type="BM", anc=0, sig2=1/1000, ngen=1000, ...)
```

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Arguments

tree	a phylogenetic tree in "phylo" format.
type	the type of plot to create. See Description.
anc	the ancestral value for the root node.
sig2	the BM rate (variance of the Brownian evolution process).
ngen	number of generations for the simulation: will rescale the tree to this total length.
	arguments to be passed to different methods.

Value

This function conducts and plots discrete time Brownian simulation and returns a vector containing the simulated states at internal nodes and tips of the tree. It also returns, by default (although this can be turned off) a tree with the branch lengths in discrete time and with a mapped discrete character (for type="threshold").

Author(s)

Liam Revell liam.revell@umb.edu>

References

Felsenstein, J. 2012. A comparative method for both discrete and continuous characters using the threshold model. *American Naturalist*, **179**, 145-156.

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

Revell, L. J. (2014) Ancestral character estimation under the threshold model from quantitative genetics. *Evolution*, bold68, 743-759.

See Also

fastBM, pbtree, phenogram, threshBayes

Examples

```
# plot BM simulation on 20 taxon tree
tree<-pbtree(n=20)
x<-bmPlot(tree)

# plot simulation of a threshold character
tree<-pbtree(n=20)
x<-bmPlot(tree,type="threshold",thresholds=c(0,1,2))</pre>
```

branching.diffusion 25

branching.diffusion Animation of branching random diffusion

Description

This function creates an animation of branching random diffusion (i.e., BM with speciation).

Usage

```
branching.diffusion(sig2=1, b=0.0023, time.stop=1000, ylim=NULL,
smooth=TRUE, pause=0.02, record=NULL, path=NULL)
```

Arguments

sig2 variance of BM process.

b birthrate for branching process.

time.stop number of generations to run.

ylim y limits (for plotting).

smooth no longer used.

pause (in s) between generations.

record filename for video file output (no video if NULL).

path full path to file for video rendering (by default is C:/Program Files/ffmpeg/bin/ffmpeg.exe).

Value

An animated plot and (optionally) a recorded video file. For animation to be recorded to file, the function requires the package "animation" as well as a video renderer.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

```
bmPlot, fastBM
```

26 brownie.lite

	brownie.lite	Likelihood test for rate variation in a continuous trait
--	--------------	--

Description

This function takes a modified "phylo" object with a mapped binary or multistate trait (see read.simmap) and data for a single continuously valued character. It then fits the Brownian rate variation ("noncensored") model of O'Meara et al. (2006; *Evolution*). This is also the basic model implemented in Brian O'Meara's "Brownie" program.

Usage

```
brownie.lite(tree, x, maxit=2000, test="chisq", nsim=100, se=NULL, ...)
```

Arguments

tree	a phylogenetic tree in modified "phylo" format (see read.simmap, make.simmap, or paintSubTree).
X	a vector of tip values for species; names(x) should be the species names.
maxit	an optional integer value indicating the maximum number of iterations for optimization - may need to be increased for large trees.
test	an optional string indicating the method for hypothesis testing - options are "chisq" or "simulation".
nsim	number of simulations (only used if test="simulation").
se	a vector containing the standard errors for each estimated mean in x.
	optional arguments.

Details

Sampling error in the estimation of species means can also be accounted for by assigning the vector se with the species specific sampling errors for x.

Value

A list with the following components:

sig2.single	is the rate for a single rate model - this is usually the "null" model.
a.single	is the estimated state at the root node for the single rate model.
var.single	variance on the single rate estimator - obtained from the Hessian.
logL1	log-likelihood of the single-rate model.
k1	number of parameters in the single rate model (always 2).
sig2.multiple	is a length p (for p rates) vector of BM rates from the multi-rate model.
a.multiple	is the estimated state at the root node for the multi-rate model.

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p x p variance-covariance matrix for the p rates - the square-roots of the diagovar.multiple

nals should give the standard error for each rate.

logL.multiple log-likelihood of the multi-rate model.

number of parameters in the multi-rate model (p+1). k2

P-value for a likelihood ratio test against the χ^2 distribution; or P.chisq P.sim P-value for a likelihood ratio test agains a simulated null distribution. logical value indicating if the likelihood optimization converged. convergence

Author(s)

Liam Revell liam.revell@umb.edu>

References

O'Meara, B. C., C. Ane, M. J. Sanderson, and P. C. Wainwright. (2006) Testing for different rates of continuous trait evolution using likelihood. Evolution, 60, 922–933.

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). Methods Ecol. Evol., 3, 217-223.

See Also

brownieREML, evol.vcv

brownieREML REML version of brownie.lite

Description

This function takes a modified "phylo" object with a mapped binary or multistate trait (see read.simmap) and data for a single continuously valued character. It then uses restricted maximum likelihood (REML) to fit the Brownian rate variation ("noncensored") model of O'Meara et al. (2006; Evolution). This function is similar to brownie.lite but uses REML (which is faster and unbiased) instead of ML. REML optimization takes advantage of Felsenstein's (1985) contrasts algorithm.

Usage

```
brownieREML(tree, x, maxit=2000, ...)
```

Arguments

tree	a phylogenetic tree in modified "phylo" format (see read.simmap and make.simmap).
Χ	a vector of tip values for species; names(x) should be the species names.
maxit	an optional integer value indicating the maximum number of iterations for optimization - may need to be increased for large trees.
	optional arguments.

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Value

A list with the following components:

sig2.single is the rate for a single rate model - this is usually the "null" model.

logL1 log-likelihood of the single-rate model.

sig2.multiple is a length p (for p rates) vector of BM rates from the multi-rate model.

logL2 log-likelihood of the multi-rate model.

convergence numerical value from optim.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Felsenstein, J. 1985. Phylogenies and the comparative method. American Naturalist, 125, 1-15.

O'Meara, B. C., C. Ane, M. J. Sanderson, and P. C. Wainwright. 2006. Testing for different rates of continuous trait evolution using likelihood. *Evolution*, **60**, 922-933.

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

brownie.lite, evol.vcv, evol.rate.mcmc

cladelabels Add labels to subtrees of a plotted phylogeny

Description

This function adds clade labels to a plotted tree.

Usage

```
cladelabels(tree=NULL, text, node, offset=NULL, wing.length=NULL, cex=1,
orientation="vertical")
```

Arguments

tree an object of class "phylo". If not supplied, the function will obtain the	tree	an object of class	"phylo".	If not supplied, the	he function wi	ll obtain the last
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plotted phylogeny from the environmental variable last_plot.phylo.

text desired clade label text.

node number for the most recent common ancestor of members of the clade.

offset offset (as a multiplier of character width) for the label. Defaults to offset=1 if

tree is supplied or offset=8 otherwise.

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wing.length length of the wings to add to the top & bottom of the label bar (in character

widths).

cex character expansion factor.

orientation orientation of the text. Can be orientation = "vertical" (the default) or

"horizontal".

Details

This function presently works only for rightward facing plotted phylogenies - but no warning will be returned if your tree does not conform to this requirement!

Author(s)

Liam Revell liam.revell@umb.edu>

References

Paradis, E., J. Claude, and K. Strimmer (2004) APE: Analyses of phylogenetics and evolution in R language. *Bioinformatics*, **20**, 289-290.

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

nodelabels

collapse.to.star

Collapse a subtree to a star phylogeny

Description

This function collapses a subtree to a star. If the tree has edge lengths, the function will keep the tips at the same height above the root as in the original tree.

Usage

```
collapse.to.star(tree, node)
```

Arguments

tree an object of class "phylo".

node node for the clade to be collapsed.

Value

An object of class "phylo".

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Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

```
splitTree, starTree
```

collapseTree

Interactive tree visualizer

Description

Function creates an interactive visualization of collapsing & expanding clades on the tree.

Usage

```
collapseTree(tree, ...)
```

Arguments

tree an object of class "phylo".

... optional arguments. These *mostly* match the arguments of plotSimmap, but also include the argument drop.extinct which will (if the input tree is ultrametric)

drop any 'extinct' lineages from the tree that is returned by the function.

Details

Function first plots a fan style tree, and then the user collapses node on the tree by clicking on them. Collapsed nodes are collapsed to the common ancestor of the clade. Nodes that have been collapsed can also be expanded by clicking.

Value

Returns the final plotted tree.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

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See Also

```
plotTree, plotSimmap
```

consensus.edges

Compute consensus edges for a tree under some criterion

Description

This function computes consensus edge lengths by different methods.

Usage

```
consensus.edges(trees, method=c("mean.edge", "least.squares"), ...)
```

Arguments

trees object of class "multiPhylo" in which the trees must have edge lengths. This

could be, for instance, a sample from the posterior distribution of trees in a

Bayesian analysis.

method method for computing the edge lengths. Could be the mean of all trees in which

the edge is present, or it could be the least-squares edge lengths computed on the mean patristic distance matrices from the input phylogenies in trees. Note that in the latter case the phangorn function nnls.tree is used and the option

rooted will be set to is.rooted(tree) for the consensus tree.

... optional arguments, the most popular of which is consensus.tree - a user sup-

plied consensus tree. Another optional argument for method="mean.edge" is if.absent which tells the function how to include absent edges in the computation of average edge lengths. Possible values are "zero" (the default) or

"ignore".

Value

An object of class "phylo" with edge lengths.

Author(s)

Liam Revell liam.revell@umb.edu>

32 contMap

contMap	Map continuous trait evolution on the tree	

Description

Function plots a tree with a mapped continuous character. The mapping is accomplished by estimating states at internal nodes using ML with fastAnc, and the interpolating the states along each edge using equation [2] of Felsenstein (1985).

Usage

```
contMap(tree, x, res=100, fsize=NULL, ftype=NULL, lwd=4, legend=NULL, lims=NULL, outline=TRUE, sig=3, type="phylogram", direction="rightwards", plot=TRUE, ...) ## S3 method for class 'contMap' plot(x, ...)
```

Arguments

tree	object of class "phylo".
х	vector of phenotypic trait values for species. names(x) should contain the species names and match tree\$tip.label. Or, for plot.contMap, an object of class "contMap".
res	resolution for gradient plotting. Larger numbers indicate a finer (smoother) gradient.
fsize	relative font size - can be a vector with the second element giving the font size for the legend.
ftype	font type - see options in plotSimmap. As with fsize, this can be a vector with the second element giving font type for the legend.
lwd	line width for branches. Can be a single integer number or a vector. In the latter case, the second number will be taken to be the desired legend width.
legend	if FALSE no legend is plotted; if a numeric value, it gives the length of the legend in units of branch length. Default is 0.5 times the total tree length.
lims	range for the color map. By default, this will be $c(min(x), max(x))$, and should always include this range.
outline	logical value indicating whether or not to outline the branches of the tree in black.
sig	the number of decimal places to show on the legend limits.
type	type of plot desired. Options are "phylogram" for a rightward square phylogram; and "fan" for a circular phylogram.
direction	plotting direction for type="phylogram".
plot	logical value indicating whether or not to plot the tree. If plot=FALSE then an object of class "contMap" will be returned without plotting.

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. . .

optional arguments for plot.contMap which include all the arguments of contMap except for tree, x, res, and lims. Also method, "fastAnc", "anc.ML", or "user" (for user-supplied states) specifying which function to use for ancestral state estimation; hold specifies whether or not to hold output to graphical device before plotting (defaults to hold=TRUE); and anc.states a vector containing some or multiple ancestral user-supplied ancestral states at nodes. Some other plotting arguments, such as xlim and ylim, may also work.

Value

Plots a tree. An object of class "contMap" is returned invisibly.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Felsenstein, J. 1985. Phylogenies and the comparative method. American Naturalist, 125, 1-15.

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

Revell, L. J. 2013. Two new graphical methods for mapping trait evolution on phylogenies. *Methods in Ecology and Evolution*, **4**, 754-759.

See Also

```
anc.ML, densityMap, fastAnc, plotSimmap
```

cophylo

Creates a co-phylogenetic plot

Description

This function creates an object of class "cophylo" or, in the case of plot.cophylo, plots that object. The function can (optionally) first attempt to rotate the nodes of both trees to optimize vertical matching of tips.

Usage

```
cophylo(tr1, tr2, assoc=NULL, rotate=TRUE, ...) ## S3 method for class 'cophylo' plot(x, ...)
```

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Arguments

tr1	object of class "phylo".
tr2	object of class "phylo".
assoc	matrix containing the tip labels in tr1 to match to the tip labels in tr2. Note that not all labels in either tree need to be included; and, furthermore, one label in tr1 can be matched with more than one label in tr2, or vice versa.
rotate	logical argument indicating whether nodes on both trees should be rotated to attempt to match in vertical position.
Х	in the case of plot.cophylo, an object of class "cophylo" to be plotted.
	optional arguments to be passed to tipRotate, or, in the case of plot.cophylo, to the internally used tree plotting function, phylogram. phylogram takes similar arguments to plotSimmap, such as fsize, ftype, lwd, and pts, though not all options from plotSimmap and plotTree are swilched. If foi as is symplical

all options from plotSimmap and plotTree are available. If fsize is supplied as a vector, different size fonts for the left & right facing trees may be used. In addition, the optional argument scale.bar, which should be a vector containing the lengths of the scale bars desired for the right & left trees, will add scale

bars to the plot when supplied to plot.cophylo.

Details

If no matrix of associations, assoc, is provided, then cophylo will look for exact matches of tip labels between trees.

Value

An object of class "cophylo" which includes the following components or a pair of plotted facing phylogenies with links between tips as specified in assoc.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). Methods Ecol. Evol., 3, 217-223.

See Also

```
cophyloplot, plotSimmap
```

Examples

```
tr1<-pbtree(n=26,tip.label=LETTERS)</pre>
tr2<-pbtree(n=26,tip.label=sample(LETTERS))</pre>
obj<-cophylo(tr1,tr2)</pre>
plot(obj)
```

countSimmap 35

countSimmap	Counts the number of character changes on a SIMMAP style tree or set of trees

Description

This function takes a tree or a set of trees with a mapped discrete character (SIMMAP style, e.g., see make.simmap or read.simmap), and computes the total number of character changes as well as the number of character changes between all states.

Usage

```
countSimmap(tree, states=NULL, message=TRUE)
```

Arguments

tree	a single tree or a set of trees with a mapped discrete character (e.g, see make.simmap or read.simmap.
states	optional argument with the states for the mapped character. If not provided, these will be computed from the tree. This is useful if averaging across many trees, some of which may lack certain states.
message	optional logical argument indicating whether or not to return an informative message about the function output.

Value

A list with up to three elements: N is an integer value giving the total number of character changes on the tree; Tr gives the number of of transitions between row and column states (or a matrix containing both N and the transitions between states, in rows, for an object of class "multiPhylo"); and (optionally) message contains an explanatory message about the function output.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

Examples

```
tree<-pbtree(n=100, scale=1)
Q<-matrix(c(-2,1,1,1,-2,1,1,1,-2),3,3)
colnames(Q)<-rownames(Q)<-c("A","B","C")
mtree<-sim.history(tree,Q)
countSimmap(mtree,states=rownames(Q))</pre>
```

36 densityMap

densityMap	Plot posterior density of stochastic mapping on a tree	
densityMap	Plot posterior density of stochastic mapping on a tree	

Description

Function plots a tree with the posterior density for a mapped character from stochastic character mapping on the tree. Since the mapped value is the probability of being in state "1", only binary [0,1] characters are allowed.

Usage

```
densityMap(trees, res=100, fsize=NULL, ftype=NULL, lwd=3, check=FALSE, legend=NULL, outline=FALSE, type="phylogram", direction="rightwards", plot=TRUE, ...)  
## S3 method for class 'densityMap'  
plot(x, ...)
```

Arguments

trees	set of phylogenetic trees in a modified "multiPhylo" object. Values for a two-state discrete character are mapped on the tree. See make.simmap and read.simmap for details.
res	resolution for gradient plotting. Larger numbers indicate a finer (smoother) gradient.
fsize	relative font size - can be a vector with the second element giving the font size for the legend.
ftype	font type - see options in plotSimmap. As with fsize, can be a vector with the second element giving font type for the legend.
lwd	line width for branches. If a vector of two elements is supplied, the second element will be taken to be the desired width of the legend bar.
check	check to make sure that the topology and branch lengths of all phylogenies in trees are equal.
legend	if FALSE no legend is plotted; if a numeric value, it gives the length of the legend in units of branch length. Default is 0.5 times the total tree length.
outline	logical value indicating whether or not to outline the branches of the tree in black.
type	type of plot desired. Options are "phylogram" for a rightward square phylogram; and "fan" for a circular phylogram.
plot	logical value indicating whether or not to plot the tree. If plot=FALSE then an object of class "densityMap" will be returned without plotting.
direction	plotting direction for type="phylogram".
Х	for plot.densityMap, an object of class "densityMap".

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. . .

optional arguments for plot.densityMap. These include all the arguments of densityMap except trees and res. Additional optional arguments include mar (margins), offset (tip label offset), and hold (whether or not to use dev.hold to hold output to graphical device before plotting; defaults to hold=TRUE). Also, the argument states can be used to 'order' the states on the probability axis (that is, which state should correspond to a posterior probability of 0 or 1). Some other plotting arguments, such as xlim and ylim, may also work.

Value

Plots a tree and returns an object of class "densityMap" invisibly.

Author(s)

Liam Revell < liam.revell@umb.edu>

References

Bollback, J. P. 2006. Stochastic character mapping of discrete traits on phylogenies. *BMC Bioinformatics*, **7**, 88.

Huelsenbeck, J. P., R. Neilsen, and J. P. Bollback. 2003. Stochastic mapping of morphological characters. *Systematic Biology*, **52**, 131-138.

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

Revell, L. J. 2013. Two new graphical methods for mapping trait evolution on phylogenies. *Methods in Ecology and Evolution*, **4**, 754-759.

See Also

make.simmap, plotSimmap, read.simmap

densityTree

Plots a posterior sample of trees

Description

Functions plots a posterior sample of trees, including with mapped discrete characters.

make. transparent is used internally and converts a color to transparent with a certain user-specified alpha level.

```
densityTree(trees, colors="blue", alpha=NULL, method="plotTree", fix.depth=FALSE,
use.edge.length=TRUE, compute.consensus=FALSE, ...)
make.transparent(color, alpha)
```

38 describe.simmap

Arguments

trees an object of class "multiPhylo" or "multiSimmap".

colors a color or a named vector of colors in which names correspond to mapped states

in an object of class "multiSimmap".

alpha transparency level for plotted trees which is passed to internally used function,

make.transparent. (0 is fully transparent, which 1 is fully opaque.) By default

will be one divided by the number of trees.

method plotting method to be used internally. Can be "plotTree" or "plotSimmap".

fix.depth logical value indicating whether or not to plot trees with a fixed depth or to

permit plotted trees to have different depths.

use.edge.length

logical value indicating whether to use the edge lengths of the input tree. Defaults to use.edge.length=TRUE unless any input tree edge lengths are NULL.

compute.consensus

logical value indicating whether or not to use the tip order from a consensus tree.

(Defaults to compute.consensus=FALSE.

... arguments to be passed to plotTree or plotSimmap.

color in make. transparent, the color to render transparent.

Value

Function creates a plot.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

describe.simmap Summ

Summarizes a stochastic mapped tree or set of trees

Description

This function summarizes the result of one or more stochastic maps.

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Usage

```
describe.simmap(tree, ...)
## S3 method for class 'simmap'
summary(object, ...)
## S3 method for class 'multiSimmap'
summary(object, ...)
## S3 method for class 'describe.simmap'
plot(x, ...)
```

Arguments

tree a single tree or a set of trees as an object of class "simmap" or "multiSimmap",

respectively.

object of class "simmap" or "multiSimmap".

x for S3 plot method, an object of class "describe.simmap".

.. optional arguments which include: plot, a logical value indicating whether or not to plot the posterior probabilities at nodes (default is plot=FALSE); check equal, a logical value indicating whether or not to check if all trees are equal using

a logical value indicating whether or not to check if all trees are equal using all.equal.phylo (default is check.equal=FALSE); and message, a logical indicating whether or not to print an informative message to the screen (default is

message=TRUE).

Value

An object of class "describe.simmap" with the following elements:

count a matrix containing the number and types of transitions for each tree, if tree is

an object of class "multiSimmap".

times a matrix containg the times spend in each state on each tree.

ace the posterior probabilities of each node being in each state, if tree is an object

of class "multiSimmap".

legend a vector containing the plot legend, if plot=TRUE.

if class(tree)="simmap" then the function simply returns the results of countSimmap combined with the states at each node of the tree and a matrix containing the total and relative times spent in each state on the tree.

Author(s)

Liam Revell < liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

40 di2multi.simmap

di2multi.simmap	Collapse branches of zero length to polytomy in stochastic map style tree

Description

This function collapses branches of zero length (or, more specifically, branches with length shorter than tol) to create a polytomy in a stochastic-map style tree.

Usage

```
di2multi.simmap(tree, tol=1e-08)
```

Arguments

tree	modified object of class	"phylo" containing a	a stochastically mapped discrete

character.

tol length below which edges should be treated as having zero length.

Details

This function should theoretically perform similarly to di2multi in ape.

Value

A tree with a stochastically mapped discrete character

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

```
di2multi, make.simmap, read.simmap
```

dotTree 41

dotTree	Creates a phylogenetic dot plot	

Description

Creates a plot in which different sized dots/circles represent different tip values for a quantitative trait

Usage

```
dotTree(tree, x, legend=TRUE, method="plotTree", standardize=FALSE, ...)
dot.legend(x, y, min, max, Ntip, length=5, prompt=FALSE, method="plotTree", ...)
```

Arguments

tree	an object of class "phylo".
LICE	an object of class physic.

x vector of trait values; or a matrix. In the case of dot.legend, the x coordinate

of the legend.

legend logical value indicating whether or not a legend should be plotted.

method tree plotting method to be used internally. Will switch to method="phylogram"

if the number of traits is greater than one. For dot.legend, it should be the

method that was used for the plot.

standardize a logical value indicating whether or not to standardize x, or each column of x,

to have a mean of zero & variance of one prior to analysis.

y y coordinate of the legend.

min minimum value for dot.legend.
max maximum value for dot.legend.

Ntip number of tips in the plotted tree for dot.legend.

length length of legend.

prompt logical value indicating whether or not to prompt for legend position.

optional arguments. In the case of dotTree, these will be passed to plotTree or a different internally used plotting function for method="phylogram". See phylo.heatmap for more detail on these arguments. Other option for dotTree also include data.type ("continuous" or "discrete"), colors, length, for data type "continuous" the length of the legend in terms of plotted circles, x.space, the spacing of the columns in the plotted data matrix, and leg.space,

the spacing of the legend dots (again, for data.type="continuous" only).

Value

Function creates a plot.

Author(s)

Liam Revell liam.revell@umb.edu>

drop.clade

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

drop.clade

Drop a clade from a tree

Description

Mostly internal function for posterior.evolrate; function drops the clade containing the species in tip.

Usage

```
drop.clade(tree, tip)
```

Arguments

tree object of class "phylo". tip set of tips in a clade.

Details

Probably should not use unless you know what you're doing.

Value

An object of class "phylo".

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

drop.leaves 43

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Drop all the leaves (tips) from a tree

Description

Drops all the leaves from a tree, leaving behind only the structure leading to internal nodes.

Usage

```
drop.leaves(tree, ...)
```

Arguments

tree object of class "phylo".

optional arguments. Presently includes only the logical value keep.tip.labels

which tells the function how to labels the tips on the reduced tree.

Value

An object of class "phylo".

Author(s)

Liam Revell < liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

```
drop.tip.contMap
```

Drop tip or tips from an object of class "contMap" or "densityMap"

Description

This function drops one or multiple tips from an object of class "contMap" or "densityMap". This function is equivalent to drop. tip but for an object of this class.

```
drop.tip.contMap(x, tip)
drop.tip.densityMap(x, tip)
```

44 drop.tip.simmap

Arguments

X	an object of class "contMap" or "densityMap".
tip	name or names of species to be dropped.

Details

For more information about objects of class "contMap" or "densityMap", please refer to the documentation pages for contMap or densityMap, respectively.

Value

```
An object of class "contMap" or "densityMap".
```

Author(s)

```
Liam Revell < liam.revell@umb.edu>
```

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

```
contMap, densityMap, drop.tip, drop.tip.simmap
```

drop.tip.simmap

Drop tips or extract clade from tree with mapped discrete character

Description

This function drops one or multiple tips from the modified "phylo" object with a mapped binary or multistate trait (see read.simmap) while maintaining the matrix \$mapped.edge and list of mappings by branch maps. This function is equivalent to drop.tip but for a tree with a mapped discrete character.

extract.clade.simmap is functionally equivalent to extract.clade but preserves discrete character mappings on the tree.

Usage

```
drop.tip.simmap(tree, tip)
extract.clade.simmap(tree, node)
```

Arguments

tree a modified object of class "phylo" (see read.simmap).

tip name or names of species to be dropped.

node number for the root node of the clade to be extracted.

edgeProbs 45

Value

A modified object of class "phylo" containing the elements maps and \$mapped.edge with the time spent in each state along each edge of the tree.

Author(s)

Liam Revell < liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

brownie.lite, drop.tip, extract.clade, make.simmap, read.simmap, sim.history

edgeProbs

Compute the relative frequencies of state changes along edges

Description

This function computes the relative frequencies of character state changes along edges from a sample of stochastically mapped character histories. This function assumes that all trees in the sample differ only in their mapped histories & not at all in topology or branch lengths. Note that it only asks whether the starting and ending states of the edge differ in a particular way, and thus ignores multiple-hits along a single edge.

Usage

edgeProbs(trees)

Arguments

trees

an object of class "multiSimmap" containing a sample of trees that are identical in topology & branch lengths with different stochastically mapped character histories.

Value

The object that is returned is a matrix with the state changes & the relative frequency of each state change. Rows are in the order of the matrix edge for any of the mapped trees.

Author(s)

Liam Revell liam.revell@umb.edu>

46 estDiversity

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

plotSimmap

estDiversity

Estimate diversity at each node of the tree

Description

This function estimates the lineage density at each node in the tree based on a biogeographic model (based on Mahler et al. 2010).

Usage

```
estDiversity(tree, x, method=c("asr","simulation"), model="ER", ...)
```

Arguments

tree is a phylogenetic tree in "phylo" format.

x a vector containing the biogeographic area for each of the tip taxa.

method method for reconstructing ancestral biogeography.

model model for ancestral character estimation. In theory, any model from ace; however only symmetric models permitted for method="asr".

... optional arguments. So far, this includes only nsim, the number of stochastic mappings to conduct using make.simmap for method="simulation".

Details

Two different methods are implemented in the current version. For method="asr" the state at the current node, and at each position along each co-extant internal edge, is computed as the marginal (empirical Bayesian) ancestral state reconstruction using the re-rooting method of Yang (2006). The lineage density is then computed as the sum of the marginal reconstructions (posterior probabilities) times the summed marginal ancestral reconstructions across co-extant edges. In method="simulation", stochastic character mapping is used to generate optional argument nsim stochastic maps of ancestral biogeography. Then the lineage density at each node is computed as the number of co-existing lineages with the same biogeography as the focal node, averaged acrossed stochastic maps. The importance of this distinction may depend on the degree to which reconstructions at internal nodes are independent, which relates to the distinction between marginal and joint reconstruction (e.g., see Yang 2006).

Value

A vector containing the estimated lineage density at each node

evol.rate.mcmc 47

Author(s)

Liam Revell < liam.revell@umb.edu>

References

Mahler, D. L, L. J. Revell, R. E. Glor, and J. B. Losos. (2010) Ecological opportunity and the rate of morphological evolution in the diversification of Greater Antillean anoles. *Evolution*, **64**, 2731-2745.

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

Yang, Z. (2006) Computational Molecular Evolution. Oxford University Press.

See Also

fitDiversityModel

evol.rate.mcmc

Bayesian MCMC method for identifying exceptional phenotypic diversification in a phylogeny

Description

This function takes a phylogenetic tree and data for a single continuously valued character and uses a Bayesian MCMC approach to identify the phylogenetic location of a shift in the evolutionary rate through time.

Usage

```
evol.rate.mcmc(tree, x, ngen=10000, control=list())
```

Arguments

tree a phylogenetic tree in "phylo" format.

a vector of tip values for species; names(x) should be the species names.

ngen an optional integer value indicating the number of generations for the MCMC.

control a list of control parameters containing the following elements: sig1: starting

value for $\sigma(1)^2$; sig2: starting value for $\sigma(2)^2$; a: starting value for a; sd1: standard deviation for the normal proposal distribution for $\sigma(1)^2$; sd2: standard deviation for the normal proposal distribution for $\sigma(2)^2$; kloc: scaling parameter for tree move proposals - $1/\lambda$ for the reflected exponential distribution; sdlnr: standard deviation on the log-normal prior on $\sigma(1)^2/\sigma(2)^2$; rand.shift: probability of proposing a random shift in the tree (improves mixing); print: print frequency for the MCMC; sample: sample frequency.

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Details

Default values of control are given in Revell et al. (2012).

Value

A list with the following components:

mcmc results from the MCMC run.

tips list of stips in rate $\sigma(1)^2$ for each sampled generation of MCMC (to polarize the

rate shift).

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

Revell, L. J., D. L. Mahler, P. Peres-Neto, and B. D. Redelings. (2012) A new method for identifying exceptional phenotypic diversification. *Evolution*, **66**, 135-146.

See Also

anc.Bayes, brownie.lite, evol.vcv, minSplit, posterior.evolrate

evol.vcv

Likelihood test for variation in the evolutionary VCV matrix

Description

This function takes a modified "phylo" object with a mapped binary or multistate trait and data for an arbitrary number of continuously valued character. It then fits the multiple evolutionary variance-covariance matrix (rate matrix) model of Revell & Collar (2009; *Evolution*).

Usage

```
evol.vcv(tree, X, maxit=2000, vars=FALSE, ...)
```

Arguments

tree a phylogenetic tree in modified "phylo" format (see read. simmap).

X an n x m matrix of tip values for m continuously valued traits in n species - row

names should be species names.

maxit an optional integer value indicating the maximum number of iterations for opti-

mization - may need to be increased for large trees.

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vars an optional logical value indicating whether or not to estimate the variances of

the parameter estimates from the Hessian matrix.

... optional arguments.

Details

This function performs optimization by first optimizing the likelihood with respect to the Cholesky matrices using optim. Optimization is by method="Nelder-Mead". Using box constraints does not make sense here as they would be applied to the Cholesky matrix rather than the target parameters. May have to increase maxit for large trees and more than 2 traits.

Value

A list with the following components:

R. single vcv matrix for the single rate matrix model.

vars.single optionally, a matrix containing the variances of the elements of R. single.

logL1 log-likelihood for single matrix model.

k1 number of parameters in the single marix model.

R.multiple mxmxp array containing the p estimated vcv matrices for the p regimes painted

on the tree.

vars.multiple optionally, an array containing the variances of the parameter estimates in R. multiple.

logL.multiple log-likelihood of the multi-matrix model.

k2 number of parameters estimated in this model. P.chisq P-value of the χ^2 test on the likelihood ratio.

convergence logical value indicating whether or not the optimization has converged.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J., and D. C. Collar (2009) Phylogenetic analysis of the evolutionary correlation using likelihood. *Evolution*, **63**, 1090-1100.

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

```
evol.rate.mcmc, brownie.lite
```

50 evolvcv.lite

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Likelihood test for a shift in the evolutionary correlation between traits

Description

This function takes a modified "phylo" object with a mapped binary or multistate trait and data for two and only two continuously valued character. It then fits four different evolutionary models: common rates and correlation; different rates, common correlation; different correlations, common rates; no common structure.

Usage

```
evolvcv.lite(tree, X, maxit=2000, tol=1e-10)
```

Arguments

tree	a phylogenetic tree in modified "phylo" format (see read.simmap).
X	an n x m matrix of tip values for m continuously valued traits in n species - row names should be species names.
maxit	an optional integer value indicating the maximum number of iterations for optimization - may need to be increased for large trees.
tol	tolerance value for "L-BFGS-B" optimization.

Value

A list with the results summarized for each model.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J., and D. C. Collar (2009) Phylogenetic analysis of the evolutionary correlation using likelihood. *Evolution*, **63**, 1090-1100.

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

```
brownie.lite, evol.vcv
```

exhaustiveMP 51

exhaustiveMP	Exhaustive and branch & bound MP optimization	

Description

This function does exhaustive and branch & bound MP searches.

Usage

```
exhaustiveMP(data, tree=NULL, method="branch.and.bound")
```

Arguments

data is a phyDat (Schliep 2011) object containing DNA or other data.

tree an optional input tree (used only with method="branch.and.bound").

method an optional string indicating method to use: "branch.and.bound" or "exhaustive".

Details

Should probably not be used for more than about 8 species (and definitely not more than 10 species). Performs parsimony calculations using parsimony in the "phangorn" package (Schliep 2011).

Value

A "phylo" or "multiPhylo" object that is the MP tree or set of MP trees. It also returns the parsimony scores in attr(trees, "pscore") or attr(trees[[i]], "pscore") for the ith tree.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Felsenstein, J. (2004) Inferring Phylogenies. Sinauer.

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

Schliep, K. P. (2011) phangorn: phylogenetic analysis in R. *Bioinformatics*, 27, 592-593.

```
mrp.supertree, optim.parsimony, pratchet
```

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expm

Matrix exponential

Description

Wrapper for MatrixExp that retains row/column names.

Usage

expm(Y)

Arguments

Υ

a matrix.

Value

A matrix.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

export.as.xml

Export trees & data in XML format

Description

This function exports trees & character data in XML format.

Usage

```
export.as.xml(file, trees, X)
```

Arguments

file filename for export.

trees a phylogenetic tree or trees in "phylo" or "multiPhylo" format.

X a matrix of class "DNAbin" or a matrix with discretely valued non-DNA charac-

ter data.

fancyTree 53

Details

Can be used to create input file for the program SIMMAP v1.5 (Bollback 2006), also see: http://www.simmap.com.

Value

A file.

Author(s)

Liam Revell < liam.revell@umb.edu>

References

Bollback, J. P. (2006) Stochastic character mapping of discrete traits on phylogenies. *BMC Bioinformatics*, 7, 88.

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

make.simmap, read.nexus, read.simmap, write.simmap

fancyTree

Plots special types of phylogenetic trees

Description

This function plots different types of phylogenetic trees. If type="extinction" (or any unambiguous abbreviation) the function will plot a tree in which branches preceding the MRCA of all extant taxa and branches leading only to extnct lineages are plotted with dashed red lines. If type="traitgram3d" the function will plot a three dimensional traitgram (that is, a projection of the tree into three dimensional morphospace where two dimensions are the phenotypic trait and the third axis is time since the root). In this case, the additional argument X, a matrix containing the tip values of all species (with species IDs as row names) should be supplied. Optionally, the user can also supply the matrix A, which contains the ancestral states in the tree with rows labeled by node number. If type="droptip" the function will create a two panel figure in which the first panel is the tree with lineages to be pruned highlighted; and the second panel is the pruned tree. In this case, the additional argument tip, the tip name or vector of tip names to be dropped, must be supplied. If type="densitymap", a posterior probability density "heat-map" is created based on a set of trees in a "multiPhylo" object containing a binary [0,1] mapped character. (See densityMap for additional optional arguments if type="densitymap".) If type="contmap", reconstructed continuous trait evolution is mapped on the tree. Again, see contMap for additional arguments if type="contmap". If type="phenogram95" a 95-percent phenogram is plotted using transparency to visualize uncertainty at ancestral nodes and along branches. Most of the options of phenogram are available. Finally, if type="scattergram" a phylogenetic scatter plot matrix 54 fancyTree

containing contMap style trees on the diagonal and phylomorphospace plots in non-diagonal panels is produced. For type="scattergram" a trait matrix X must be supplied. The only additional arguments available for this type are fsize, colors, and label. (See phylomorphospace for details.) Presently only type="traitgram3d" uses the list control which can be supplied the same set of control parameters as phylomorphospace3d, as well as the control parameter maxit which will be passed to anc.ML. Finally, the optional argument hold will be passed to multiple methods if supplied. It is a logical value that indicates whether or not the output to the graphical device should be held using dev.hold before plotting (defaults to hold=TRUE).

Usage

```
fancyTree(tree, type=c("extinction","traitgram3d","droptip","densitymap",
"contmap","phenogram95","scattergram"), ..., control=list())
```

Arguments

```
tree a phylogenetic tree in "phylo" format.

type the type of special plot to create. See Description.

arguments to be passed to different methods.
```

control a list of control parameters, depending on type.

Value

This function plots different types of phylogenetic trees. For type="droptip" the function also returns the pruned tree.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

contMap, densityMap, drop.tip, phenogram, phylomorphospace3d, plot.phylo, plotSimmap

Examples

```
# plot tree with extinction
set.seed(10)
tree<-pbtree(b=1,d=0.4,t=4)
fancyTree(tree,type="extinction")

## Not run:
# plot 3D traitgram
tree<-pbtree(n=50,scale=10)
Y<-sim.corrs(tree,vcv=matrix(c(1,0.75,0.75,1),2,2))</pre>
```

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```
fancyTree(tree,type="traitgram3d",X=Y,control=list(spin=FALSE))
# plot with internal nodes from simulation
Y<-sim.corrs(tree, vcv=matrix(c(1, 0.75, 0.75, 1), 2, 2), internal=TRUE)
B<-Y[length(tree$tip)+1:tree$Nnode,]; Y<-Y[1:length(tree$tip),]</pre>
fancyTree(tree,type="traitgram3d",X=Y,A=B,control=list(simple.axes=TRUE,spin=FALSE))
## End(Not run)
# plot with dropped tips
tree<-pbtree(n=30)
tips<-sample(tree$tip.label)[1:10]</pre>
pruned<-fancyTree(tree,type="droptip",tip=tips)</pre>
## Not run:
# plot 95-percent CI phenogram
tree<-pbtree(n=30)</pre>
x<-fastBM(tree)
fancyTree(tree,type="phenogram95",x=x)
## End(Not run)
```

fastAnc

Fast estimation of ML ancestral states

Description

This function performs fast estimation of the ML ancestral states for a continuous trait by taking advantage of the fact that the state computed for the root node of the tree during Felsenstein's (1985) contrasts algorithm is also the MLE of the root node. Thus, the function reroots the tree at all internal nodes and computes the contrasts state at the root each time. The function can also (optionally) compute variances or 95-percent confidence intervals on the estimates.

Usage

```
fastAnc(tree, x, vars=FALSE, CI=FALSE, ...)
```

Arguments

tree	an object of class "phylo".
X	a vector of tip values for species; names(x) should be the species names.
vars	a logical value indicating whether or not to compute variances on the ancestral state estimates. Variances are based on Equation (6) of Rohlf (2001) .
CI	a logical value indicating whether or not to compute 95-percent confidence intervals on state estimates.
	optional arguments. Presently this consists of anc.states, a named vector containing ancestral states to fix. Names should correspond to node numbers in the input tree.

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Value

A named vector containing the states at internal nodes - names are node numbers; or a list containing ancestral state estimates (ace), variances on the estimates (var), and/or 95-percent confidence intervals (CI95).

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

```
ace, anc.Bayes, anc.ML, pic
```

Examples

```
tree<-pbtree(n=50)
x<-fastBM(tree) # simulate using fastBM
fastAnc(tree,x) # estimate states</pre>
```

fastBM

Fast Brownian simulation

Description

This function conducts fast quantitative trait simulation on a phylogeny under several different models: Brownian motion (default), BM with a trend (for mu!=0), bounds (for bounds!=c(-Inf,Inf)), and OU.

Usage

```
fastBM(tree, a=0, mu=0, sig2=1, bounds=c(-Inf,Inf), internal=FALSE, nsim=1, ...)
```

Arguments

tree	is a phylogenetic tree in "phylo" format.
a	a value for ancestral state at the root node.
mu	an optional value for the mean of random normal changes along branches of the tree - can be used to simulate a trend if mu!=0.
sig2	instantaneous variance of the BM process.
bounds	a vector with the lower and upper bounds (respectively) for bounded Brownian simulation - by default simulation is unbounded.

fastMRCA 57

internal	logical value indicating whether or not to return states for internal nodes.
nsim	number of simulations.
• • •	optional arguments alpha and theta used for OU simulation. If alpha is set then mu and bounds are ignored with a warning.

Value

A vector (for nsim=1) or matrix containing the tip states for the n species in the tree, and (optionally) the ancestral states for internal nodes.

Author(s)

```
Liam Revell < liam.revell@umb.edu>
```

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

```
sim.corrs
```

Examples

```
tree<-pbtree(n=1000)
x<-fastBM(tree,sig2=0.1) # Brownian motion
y<-fastBM(tree,mu=1) # with a trend</pre>
```

```
fastMRCA Get the MRCA (or height above the root of the MRCA) of a pair of tip taxa
```

Description

This function returns the most recent common ancestor (node number) for a pair of taxa; or, in the case of fastHeight, the height above the root of the MRCA of a pair of taxa; or, in the case of fastDist, the patristic distance between a pair of taxa.

```
fastMRCA(tree, sp1, sp2)
fastHeight(tree, sp1, sp2)
fastDist(tree, sp1, sp2)
```

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Arguments

tree	a phylogenetic tree as an object of class "phylo".
sp1	species name.
sp2	species name.

Details

This function is mostly redundant with findMRCA (or findMRCA(..., type="height") in the case of fastHeight) but for very large trees will be considerably faster.

Value

The node number of the MRCA or the height above the root (for fastHeight).

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

```
findMRCA, mrca
```

Examples

```
tree<-pbtree(n=2000)
anc<-fastMRCA(tree,"t1","t15")</pre>
```

findMRCA

Get the MRCA of a set of taxa

Description

This function returns the most recent common ancestor (node number) for a set of taxa. If tips=NULL will be redundant with mrca (for type="node") or vcv.phylo, but much slower (for type="height").

```
findMRCA(tree, tips=NULL, type=c("node","height"))
```

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Arguments

tree a phylogenetic tree as an object of class "phylo".

tips a vector containing a set of tip labels.

type either "node" to return the node of the MRCA; or "height" to return the height

above the root of the MRCA of tips.

Details

If tips==NULL will return the result of a normal function call to mrca. If tips=NULL will return a matrix equal to vcv.phylo.

Value

The node number of the MRCA, or a matrix of node numbers (if tips==NULL) - for type="node"; or the height of the MRCA, or a matrix of heights (if tips==NULL) - for type="height".

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

findMRCA, mrca

Examples

```
tree<-pbtree(n=20)
anc<-findMRCA(tree,c("t1","t10","t15"))</pre>
```

fitBayes

Evolutionary model fitting with intraspecific variability using Bayesian MCMC

Description

This function uses Bayesian MCMC to sample terminal states (species means) as well as evolutionary parameters.

```
fitBayes(tree, x, ngen=10000, model="BM", method="reduced", control=list())
```

60 fitBayes

Arguments

tree an object of class "phylo".

x a vector of phenotypic values for individuals; names(x) should contain the

species names (not individual IDs).

ngen a integer indicating the number of generations for the MCMC.

model an evolutionary model: either "BM" or "lambda".

method a method: either "reduced" or "full".

control a list of control parameters containing the following elements: sig2: starting

value for σ^2 (BM rate); lambda: starting value for the λ parameter; a: starting for the state at the root node; xbar: starting values for the states at the tips; intV: starting value for the intraspecific variance (reduced method); or v: starting value for the vector of intraspecific variances for all species (full method); pr.mean: means for the prior distributions in the following order - sig2, lambda (if applicable), a, xbar, intV or v (if applicable), note that the prior probability distribution is exponential for sig2 and normal for a and y; pr.var: variances

on the prior distributions, same order as pr.mean.

Value

A matrix with number of rows ngen/control\$sample+1 containing the posterior sample and likelihoods. Matrix columns are labeled by species (for species means and variances), or by the corresponding evolutionary parameter.

Author(s)

Liam Revell < liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

Revell, L. J. and R. G. Reynolds. (2012) A new Bayesian method for fitting evolutionary models to comparative data with intraspecific variation. *Evolution*, 66, 2697-2707.

See Also

anc.Bayes, brownie.lite, evol.rate.mcmc

fitDiversityModel 61

fitDiversityModel Fit diversity-dependent phenotypic evolution model
--

Description

This function fits a diversity-dependent phenotypic evolution model (based on Mahler et al. 2010).

Usage

```
fitDiversityModel(tree, x, d=NULL, showTree=TRUE, tol=1e-6)
```

Arguments

50	
tree	an object of class "phylo".
x	a vector with tip values for a continuously distributed trait.
d	a vector containing the inferred historical diversity at each node in the tree - if d=NULL (the default) function will treat the diversification as if it occurred in a single geographic area.
showTree	optional logical value indicating whether to plot the tree transformation implied by the model.
tol	some small value by which d is incremented during rescaling of psi for optimization. If R thinks your matrices are singular during optimization, try increasing tol slightly.

Value

A list with the following components:

logL	log-likelihood of the fitted model.
sig0	estimated starting value for the rate at the root of the tree.
psi	the estimated rate of change in the rate associated with the addition of a lineage.
VCV	a matrix with the variances and covariance of the estimated parameters (from the Hessian).

Author(s)

Liam Revell < liam.revell@umb.edu>

References

Mahler, D. L, L. J. Revell, R. E. Glor, and J. B. Losos. 2010. Ecological opportunity and the rate of morphological evolution in the diversification of Greater Antillean anoles. *Evolution*, **64**, 2731-2745.

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

62 fitMk

See Also

brownie.lite, estDiversity, evol.rate.mcmc

fitMk

Fits Mk model

Description

This function fits a so-called Mk model for discrete character evolution. It is primarily designed to be used inside of make.simmap.

Two plot methods are available. plot.fitMk plots an object of class "fitMk" returned by fitMk. plot.gfit plots an object of class "gfit" from geiger's fitDiscrete function. Both plots portray the fitted model using a graph of arrows connecting states.

Usage

```
fitMk(tree, x, model="SYM", fixedQ=NULL, ...)
## S3 method for class 'fitMk'
plot(x, ...)
## S3 method for class 'gfit'
plot(x, ...)
```

Arguments

tree an object of class "phylo".

x a vector of tip values for species; names(x) should be the species names. In the

case of plot.fitMk, an object of class "fitMk".

model. See make.simmap or ace for details.

fixedQ fixed value of transition matrix Q, if one is desired.

... optional arguments, including pi, the prior distribution at the root node (defaults

to pi="equal"). For plot method optional arguments include (but may not be limited to): signif, the number of digits for the rates to be plotted; main, a character vector of length two with the headings for each subplot; cex.main, cex.traits, and cex.rates, font sizes for the various text elements of the plot; and show.zeros, a logical argument specifying whether or not to plot arrows with the ML estimated transition rate is not different from zero (with tolerance

specified by the optional argument tol).

Value

An object of class "fitMk". In the case of plot. fitMk, a plotted Mk model.

Author(s)

Liam Revell liam.revell@umb.edu>

fitPagel 63

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

```
ace, make.simmap
```

fitPagel

Function to test for correlated evolution of binary traits

Description

This function fit's Pagel's (1994) model for the correlated evolution of two binary characters. plot.fitPagel plots the fitted models using arrows.

Usage

```
fitPagel(tree, x, y, method="fitMk", model="ARD", dep.var="xy", ...)
## S3 method for class 'fitPagel'
plot(x, ...)
```

Arguments

tree	an object of class "phylo".
X	a vector of phenotypic values for a binary trait for the species in tree. For S3 plot method, an object of class "fitPagel".
у	a second binary character for the species in tree.
method	function to use for optimization (defaults to method="fitMk"). Other options are "ace" to use the ace function in ape for optimization, or to "fitDiscrete" (if the geiger package is installed) to use geiger's fitDiscrete for optimization.
mode1	model of evolution for the individual characters. Can be model="ER", "SYM" (equivalent to "ER" in this case), and "ARD".
dep.var	dependent variable. If dep.var="xy" than the rate of substitution in x depends on y & vice versa. If dep.var="x" than the substitution rate in x depends on y, but not the converse. Finally, if dep.var="y" than the rate of substitution in y depends on x, but not the converse.
	optional arguments to be passed to fitMk, ace, or fitDiscrete. For plot method optional arguments include (but may not be limited to): signif, the number of digits for the rates to be plotted; main, a character vector of length two with the headings for each subplot; cex.main, cex.sub, cex.traits, and cex.rates, font sizes for the various text elements of the plot; and lwd.by.rate, a logical argument specifying whether or not to scale arrow line widths in proportion to the estimated rates.

64 gammatest

Value

An object of class "fitPagel" which contains the optimized matrices under an independence & a dependence model, log-likelihoods, a likelihood ratio, and a P-value for the independence model based on a chi-squared test.

plot.fitPagel creates a plot showing the different fitted models with arrows.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Pagel, M. (1994) Detecting correlated evolution on phylogenies: A general method for the comparative analysis fo discrete characters. *Proceedings of the Royal Society B*, **255**, 37-45.

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

```
ace, fitMk, make.simmap
```

gammatest

Gamma test of Pybus & Harvey (2000)

Description

Conducts γ -test of Pybus & Harvey (2000).

Usage

```
gammatest(x)
```

Arguments

Х

list from function call of ltt in which ltt(...,gamma=F).

Details

Do not use for object returned by ltt(...,gamma=T).

Value

A list containing:

gamma optionally, a value for the γ -statistic. p two-tailed P-value for the γ -test.

genSeq 65

Author(s)

Liam Revell < liam.revell@umb.edu>

References

Pybus, O. G., and P. H. Harvey (2000) Testing macro-evolutionary models using incomplete molecular phylogenies. *Proc. R. Soc. Lond. B*, **267**, 2267-2272.

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

1tt

Examples

```
tree<-pbtree(n=200)
z<-ltt(tree,gamma=FALSE)
g<-gammatest(z)</pre>
```

genSeq

Simulate a DNA alignment on the tree under a model

Description

Simulates DNA sequence on tree under the specified model. Uses sim.history internally.

Usage

```
genSeq(tree, l=1000, Q=NULL, rate=1, format="DNAbin", ...)
```

Arguments

tree	object of class "phylo".
1	length of desired sequences.
Q	transition matrix for the simulation. Row and column names $c("a","c","g","t")$ (although not necessarily in that order, should be provided. If NULL, a single rate is assumed.
rate	multiplier for Q, or a vector for Gamma rate heterogeneity.
format	format of the output object. Can be "DNAbin", "phyDat", or "matrix".
	optional arguments.

Value

An object of class "DNAbin" or "phyDat", or a matrix of nucleotides.

66 getCladesofSize

Author(s)

Liam Revell < liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

Examples

```
## simulate gamma rate heterogeneity
tree<-pbtree(n=26,tip.label=LETTERS)
gg<-rgamma(n=100,shape=0.25,rate=0.25)
X<-genSeq(tree,l=100,rate=gg)</pre>
```

getCladesofSize

Get all subtrees larger than or equal to a specified size

Description

This function gets all subtrees that cannot be further subdivided into two reciprocally monophyletic subtrees of size >= clade.size.

Usage

```
getCladesofSize(tree, clade.size=2)
```

Arguments

tree is an object of class "phylo".

clade.size subtree size.

Value

An object of class "multiPhylo".

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

```
extract.clade, getDescendants
```

getDescendants 67

mbers

Description

This function returns the set of node & tip numbers descended from node.

Usage

```
getDescendants(tree, node, curr=NULL)
```

Arguments

tree a phylogenetic tree as an object of class "phylo".

node an integer specifying a node number in the tree.

curr the set of previously stored node numbers - used in recursive function calls.

Value

The set of node and tip numbers for the nodes and tips descended from node in a vector.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

paintSubTree

getExtant Returns a list of the extant or extinct lineages in a tree containing non- contemporaneous tips
--

Description

The function getExtant takes a tree as input and returns a vector containing the names of all the tips that have a height above the root that is equal (to a degree of numerical precision determined by tol) to the height of the highest tip. getExtinct returns the complement.

68 getSisters

Usage

```
getExtant(tree, tol=1e-8)
getExtinct(tree, tol=1e-8)
```

Arguments

tree a phylogeny stored as an object of class "phylo" with some tips that are non-

contemporaneous (i.e., end before the present).

tol a tolerance value to account for numerical imprecision.

Value

A vector with the tip names of extant or extinct species in the tree.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

nodeHeights

getSisters

Get the sister node number, label, or set of nodes for a node or tip

Description

This function takes a tree and node or tip number of label and returns the number or label of the sister or sisters to that node or tip.

Usage

```
getSisters(tree, node, mode=c("number","label"))
```

Arguments

tree object of class "phylo".

node a node number, tip number, node label, or tip label.

mode an optional string indicating whether to return the node or tip number(s) or the

node or tip label(s), if available.

getStates 69

Value

If mode="number" this function returns an integer or vector containing the node number of numbers of the sister node or tip. If mode="label" that this function returns a list containing up to two vectors: one for the node numbers of labels of sister nodes (if available); and the other containing the tip labels of the sister tips.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

getStates

Get the states at nodes or tips from a mapped tree

Description

This function gets the states from the nodes or tips of a mapped tree (e.g., make.simmap).

Usage

```
getStates(tree, type=c("nodes","tips"))
```

Arguments

tree is a modified object of class "phylo" or "multiPhylo".

type mode indicating whether to get states at the nodes (type="nodes") or the tips

(type="tips") of the tree.

Value

A named vector (for "phylo") or matrix (for "multiPhylo").

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

```
describe.simmap, make.simmap, read.simmap, sim.history
```

70 lambda.transform

ladderize.simmap

Ladderize a tree with a mapped discrete character

Description

This function "ladderizes" an object of class "phylo" with a mapped discrete character. For more information, please see ladderize.

Usage

```
ladderize.simmap(tree, right=TRUE)
```

Arguments

tree an object of class "phylo" with a mapped discrete character.

right a logical specifying how the tree should be ladderized.

Value

An object of class "phylo" with a mapped discrete character.

Author(s)

Liam Revell < liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

```
make.simmap, ladderize
```

lambda.transform

Lambda transformation of matrix

Description

Function multiplies the off-diagonals of a square matrix by lambda. Used internally in phyl.pca and other functions.

```
lambda.transform(lambda, C)
```

likMlambda 71

Arguments

lambda scalar, usually (but not necessarily) on the interval 0,1.

C matrix probably returned by vcv.phylo.

Details

Do not use unless you know what you're doing.

Value

A matrix.

Author(s)

Liam Revell < liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

likMlambda

Likelihood for joint lambda

Description

Computes the likelihood.

Usage

```
likMlambda(lambda, X, C)
```

Arguments

lambda scalar, usually on the interval 0,1.

X data, in a matrix.

C matrix probably returned by vcv.phylo.

Details

Do not use unless you know what you're doing.

Value

A scalar.

72 locate.fossil

Author(s)

Liam Revell < liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

locate.fossil

Locate a fossil lineage in a tree using continuous characters

Description

This function uses ML to place a fossil lineage into a tree using continuous traits.

Usage

```
locate.fossil(tree, X, ...)
```

Arguments

tree an object of class "phylo".

X a matrix with continuous character data.

.. optional arguments including time.constraint which can be a scalar (positive height above the root of the fossil or negative time before present) or a vector (age range of fossil, either positive or negative); edge.constraint, which is equivalent to constraint in locate.yeti; plot, rotate, and quiet, which have the same interpretation (and defaults) as the equivalent arguments in

locate.yeti.

Value

Optimized tree as an object of class "phylo".

Author(s)

Liam Revell < liam.revell@umb.edu>

References

Felsenstein, J. (1981) Maximum likelihood estimation of evolutionary trees from continuous characters. *American Journal of Human Genetics*, 25, 471-492.

Felsenstein, J. (2002) Quantitative characters, phylogenies, and morphometrics. In: MacLeod, N. and P. Forey (Eds.) *Morphology, Shape and Phylogeny* (pp. 27-44). Taylor and Francis, London.

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

locate.yeti 73

Revell, L. J., D. L. Mahler, R. G. Reynolds, and G. J. Slater. (In press) Placing cryptic, recently extinct, or hypothesized taxa into an ultrametric phylogeny using continuous character data: A case study with the lizard *Anolis roosevelti*. *Evolution*.

locate.yeti

Locate a cryptic, recently extinct, or missing taxon on a tree

Description

This function uses ML (or REML) to place a recently extinct, cryptic, or missing taxon on an ultrametric (i.e., time-calibrated) phylogeny.

Usage

```
locate.yeti(tree, X, ...)
```

Arguments

tree an object of class "phylo".

X a matrix with continuous character data.

optional arguments including: method ("ML" or "REML", defaults to "ML"); search

("heuristic" or "exhaustive", defaults to "heuristic"); constraint, a vector containing the daughter node numbers from tree\$edge for each edge to try; plot a logical argument specifying whether or not to plot the likelihood profile on edges (defaults to FALSE); rotate a logical indicating whether or not to rotate the data based on the input tree; and quiet, which is logical and has an

obvious interpretation.

Value

Optimized tree as an object of class "phylo".

Author(s)

Liam Revell liam.revell@umb.edu>

References

Felsenstein, J. (1981) Maximum likelihood estimation of evolutionary trees from continuous characters. *American Journal of Human Genetics*, 25, 471-492.

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

Revell, L. J., D. L. Mahler, R. G. Reynolds, and G. J. Slater. (In press) Placing cryptic, recently extinct, or hypothesized taxa into an ultrametric phylogeny using continuous character data: A case study with the lizard *Anolis roosevelti*. *Evolution*.

74 ltt

ls.tree

Least squares branch lengths for a given tree

Description

Computes the least squares branch lengths conditioned on a topology and distance matrix. Internal function for optim.phylo.ls.

Usage

```
ls.tree(tree, D)
```

Arguments

tree phylogeny.

D distance matrix.

Details

Do not use unless you know what you're doing.

Value

A tree with branch lengths.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

ltt

Creates lineage-through-time plot (including extinct lineages)

Description

This function computes LTT plot with extant and extinct lineages, and optionally conducts γ -test of Pybus & Harvey (2000). The object returned by 1tt can be plotted or re-plotted using plot.

Usage

```
ltt(tree, plot=TRUE, drop.extinct=FALSE, log.lineages=TRUE, gamma=TRUE, ...)
```

Itt 75

Arguments

tree is a phylogenetic tree in "phylo" format, or an object of class "multiPhylo"

containing a list of phylogenetic trees.

plot a logical value indicating whether or not to create LTT plot.

drop.extinct logical value indicating whether or not to drop extinct tips from the tree.

log.lineages logical value indicating whether LTT plot should be on log-linear (default) or

linear-linear scale.

gamma logical value indicating whether or not to compute equgamma from Pybus &

Harvey (2000; Proc. Roy. Soc. B).

... other arguments to be passed to plotting methods. See plot.default.

Details

Although it is calculated here, it's unclear how to interpret the γ -statistic if not all the tips in the tree are contemporaneous.

Value

An object of class "ltt" which includes the following components:

times a vector of branching times.

1tt a vector of linages.

gamma optionally, a value for the gamma-statistic.

p two-tailed P-value for the gamma-test.

If tree is an object of class "multiPhylo", then an object of class "multiLtt" is returned consisting of a list of object of class "ltt".

Author(s)

Liam Revell liam.revell@umb.edu>

References

Pybus, O. G., and P. H. Harvey (2000) Testing macro-evolutionary models using incomplete molecular phylogenies. *Proc. R. Soc. Lond. B*, **267**, 2267-2272.

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

gammatest, 1tt95

76 ltt95

Examples

```
trees<-pbtree(n=100,scale=100,nsim=10)
obj<-ltt(trees,plot=FALSE)
plot(obj,log="y",log.lineages=FALSE,main="lineage through time plots")
tree<-pbtree(b=1,d=0.25,t=4)
obj<-ltt(tree,gamma=FALSE)
obj</pre>
```

1tt95

Creates a $(1-\alpha)$ -percent CI for a set of LTTs

Description

This function computes LTT plots for a set of trees & plots a $(1-\alpha)$ -percent CI by various methods.

Usage

```
ltt95(trees, alpha=0.05, log=FALSE, method=c("lineages","times"),
mode=c("median","mean"), ...)
## S3 method for class 'ltt95'
plot(x, ...)
```

Arguments

trees	is an object of class "multiPhylo" containing a list of phylogenetic trees.
alpha	confidence level.
log	logical value indicating whether or not to plot on the log-scale.
method	plot the CI on the number of lineages given time ("lineages"); or on times given a number of lineages ("times").
mode	plot the median or mean LTT.
X	object of class "1tt95" for plotting method.
	optional arguments to be used by 1tt95 or the plotting method. So far, res gives the number of time-steps (defaults to res=100. xaxis ("standard", "negative", or "flipped") determines the scale (time from the root, time back from the present, or time from the present) of the x-axis of the plot.

Details

This function creates a plot and invisibly returns an object of class "1tt95".

Author(s)

Liam Revell liam.revell@umb.edu>

make.era.map 77

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

1tt

make.era.map

Create "era" map on a phylogenetic tree

Description

This function creates a temporal map on the tree based on limits provided by the user.

Usage

```
make.era.map(tree, limits, ...)
```

Arguments

tree a phylogenetic tree as an object of class "phylo".

limits a vector containing the temporal limits, in time since the root node of the tree,

for the mappings.

... optional arguments.

Value

A modified phylogenetic tree of class "phylo" with the following additional elements:

maps a list of named vectors containing the times spent in each state on each branch,

in the order in which they occur.

mapped.edge a matrix containing the total time spent in each state along each edge of the tree.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

```
make.simmap, read.simmap, plotSimmap
```

78 make.simmap

Examples

```
tree<-pbtree(n=1000,scale=100)
tree<-make.era.map(tree,c(0,25,50,75))
plotSimmap(tree,pts=FALSE,ftype="off")</pre>
```

make.simmap

Simulate stochastic character maps on a phylogenetic tree or trees

Description

This function performs stochastic mapping using several methods.

For Q="empirical", it first fits a continuous-time reversible Markov model for the evolution of x and then simulates stochastic character histories using that model and the tip states on the tree. This is the same procedure that is described in Bollback (2006), except that simulation is performed using a fixed value of the transition matrix, Q, instead of by sampling Q from its posterior distribution.

For Q="mcmc", it first samples Q nsim times from the posterior probability distribution of Q using MCMC, then it simulates nsim stochastic maps conditioned on each sampled value of Q.

For Q set to a matrix, it samples stochastic mappings conditioned on the fixed input matrix.

Usage

```
make.simmap(tree, x, model="SYM", nsim=1, ...)
```

Arguments

tree a phylogenetic tree as an object of class "phylo", or a list of trees as an object

of class "multiPhylo".

x a vector containing the tip states for a discretely valued character, or a matrix

containing the prior probabilities of tip states in rows.

model a character string containing the model - options as in ace.

nsim number of simulations. If tree is an object of class "multiPhylo", then nsim

simulations will be conducted per tree.

optional arguments. So far, pi gives the prior distribution on the root node of the tree - options are "equal", "estimated", or a vector with the frequencies. If pi="estimated" then the stationary distribution is estimated by numerically solving pi*Q=0 for pi, and this is used as a prior on the root. Defaults to pi="equal" which results in the root node being sampled from the conditional scaled likelihood distribution at the root. message tells whether or not to print a message containing the rate matrix, Q and state frequencies. Defaults to message=TRUE. For optional argument Q="mcmc" the mean value of Q from the posterior sample is printed. tol gives the tolerance for zero elements in Q. (Elements less then tol will be reset to tol). Q can be a string ("empirical" or "mcmc"), or a fixed value of the transition matrix, Q. If "empirical" than a single value of Q, the most likely value, is used for all simulations. If "mcmc",

make.simmap 79

then nsim values of Q are first obtained from the posterior distribution for Q using Bayesian MCMC, then a simulated stochastic character map is generated for each value of Q. vQ a single numeric value or a vector containing the (normal) sampling variances for the MCMC. The order of vQ is assumed to be in the order of the index.matrix in ace for the chosen model. prior a list containing alpha and beta parameters for the gamma prior distribution on the transition rates in Q. Note that alpha and beta can be single values or vectors, if different priors are desired for each value in Q. As for vQ, the order of prior is assumed to be the order of index.matrix in ace. prior can also be given the optional logical value use.empirical which tells the function whether or not to give the prior distribution the empirical mean for Q. If TRUE then only prior\$beta is used and prior\$alpha is set equal to prior\$beta times the empirical mean of Q. burnin and samplefreq are burn-in and sample frequency for the MCMC, respectively.

Details

Uses code modified from ace (by Paradis et al.) to perform Felsenstein's pruning algorithm & compute the likelihood.

As of phytools>=0.2-33 x can be a vector of states or a matrix containing the prior probabilities of tip states in rows. In this case the column names of x should contain the states, and the row names should contain the tip names.

Note that there was a small (but potentially significant) bug in how node states were simulated by make.simmap in versions of phytools<=0.2-26. Between phytools 0.2-26 and 0.2-36 there was also a bug for asymmetric models of character change (e.g., model="ARD"). Finally, between phytools 0.2-33 and phytools 0.2-47 there was an error in use of the conditional likelihoods for the root node, which caused the root node of the tree to be sampled incorrectly. All of these issues should be fixed in the present version.

Q="mcmc" and Q set to a fixed value were introduced to phytools >= 0.2-53. As of the present version of phytools, this method is still somewhat experimental & should be used with caution.

If tree is an object of class "multiPhylo" then nsim stochastic maps are generated for each input tree.

Value

A modified phylogenetic tree of class "phylo" (or a modified "multiPhylo" object, for nsim > 1) with the following additional elements:

maps a list of named vectors containing the times spent in each state on each branch,

in the order in which they occur.

mapped.edge a matrix containing the total time spent in each state along each edge of the tree.

Q the assumed or sampled value of Q.

logL the log-likelihood of the assumed or sampled Q.

Author(s)

Liam Revell liam.revell@umb.edu>

80 map.overlap

References

Bollback, J. P. (2006) Stochastic character mapping of discrete traits on phylogenies. *BMC Bioinformatics*, **7**, 88.

Huelsenbeck, J. P., R. Neilsen, and J. P. Bollback (2003) Stochastic mapping of morphological characters. *Systematic Biology*, **52**, 131-138.

Paradis, E., J. Claude, and K. Strimmer (2004) APE: Analyses of phylogenetics and evolution in R language. *Bioinformatics*, **20**, 289-290.

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

brownie.lite, brownieREML, countSimmap, describe.simmap, evol.vcv, plotSimmap, read.simmap,
write.simmap

map.overlap

Proportional overlap between two mapped character histories on a tree

Description

This function computes the fraction of a stochastic character mapping that is shared between two differently mapped trees. In map.overlap it will compute a single quantitying giving the overall similarity of the maps, consequently this measure only makes sense of some or all of the states are shared between the two mapped tress. In Map.Overlap is computed in which the rows are states observed in tree1 and columns give the states for tree2.

Usage

```
map.overlap(tree1, tree2, tol=1e-6, ...)
Map.Overlap(tree1, tree2, tol=1e-06, standardize=TRUE, ...)
```

Arguments

tree1 an object of class "simmap".
tree2 an object of class "simmap".
tol an optional tolerance value.

standardize for Map.Overlap, a logical value indicating whether or not to standardize over-

lap by dividing by the summed branch length of the tree.

... optional arguments, such as check.equal, a logical value indicating whether

or not to check if tree1 and tree2 match in underlying topology and branch lengths (they should). This value is TRUE by default, but can be set to FALSE if

tree1 and tree2 are known to be equal to speed up calculation.

map.to.singleton 81

Value

A numerical value on the interval 0-1, for map. overlap; or a matrix (Map. Overlap).

Author(s)

Liam Revell < liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

```
make.simmap, read.simmap
```

map.to.singleton

Converts a mapped tree to a tree with singleton nodes

Description

The function map.to.singleton takes an object of class "phylo" with a mapped discrete character and converts it to a tree with singleton nodes, in which edge has only one state. The states for each edge are stored in names(tree\$edge.length). plotTree.singletons plots a tree with singleton nodes. Finally, drop.tip.singleton drops tips from the tree leaving ancestral nodes for all remaining tips as singletons. rootedge.to.singleton converts a tree with a root edge to a tree with a singleton node instead.

Usage

```
map.to.singleton(tree)
plotTree.singletons(tree)
drop.tip.singleton(tree, tip)
rootedge.to.singleton(tree)
```

Arguments

tree a modified object of class "phylo" with a mapped discrete character or (for

plotTree.singletons a tree with singleton nodes.

tip a tip label or vector of tip labels.

Value

An object of class "phylo" with singleton nodes. plotTree.singletons plots a tree. If names (tree\$edge.length)!=NULL it will use a different color from palette for each mapped state.

82 mapped.states

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

```
collapse.singles, drop.tip, make.simmap
```

mapped.states	Returns a vector, matrix, or list of the mapped states on a tree or set of
	trees

Description

Computes and orders a vector, matrix, or list of the unique mapped states on a tree or state of trees of class "simmap" or "multiSimmap".

Usage

```
mapped.states(tree, ...)
```

Arguments

tree a single tree or a set of trees as an object of class "simmap" or "multiSimmap", respectively.
... optional arguments.

Value

A vector, matrix, or list.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

markChanges 83

markChanges	Add marked changes to a plotted tree with mapped discrete character

Description

This function adds the reconstructed changes to a plotted tree with a stochastically mapped discrete character.

Usage

```
markChanges(tree, colors=NULL, cex=1, lwd=2, plot=TRUE)
```

Arguments

tree	an object of class "simmap".	
CICC	an object of class similar.	

colors a named vector of colors used to plot the stochastically mapped character on the

tree.

cex expansion factor for line height.

lwd line width.

plot logical value indicating whether the changes should be plotted or not.

Value

This function returns (invisibly) a matrix containing the x & y coordinates of the marked changes.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

plotSimmap

84 matchNodes

matchNodes

Matches nodes between two trees

Description

This function returns a matrix in which the first column contains all the internal nodes for tr1 and the second column contains the matching nodes from tr2, inasmuch as they can be identified. For method="descendants", pairs of matching nodes are defined by sharing all descendant leaves in common. for method="distances", nodes are considered to matched if the share the same set of distances (or proportional distances, for optional argument corr=TRUE) to all tips. matchLabels is functionally equivalent but matches node (tip) numbers based on the labels only.

Usage

```
matchNodes(tr1, tr2, method=c("descendants", "distances"), ...)
matchLabels(tr1, tr2)
```

Arguments

tr1 first tree. tr2 second tree.

method method to use to match nodes between trees. "descendants" uses the tip

species descended from each node; "distances" uses the distances from the nodes to the tips. Any umambiguous shortening of "descendants" or "distances"

is allowed.

... optional arguments which may or may not be used depending on method. tol

is a tolerance value for the difference from exact matching that is allowed for method="distances". corr, which is FALSE by default, indicates whether to match nodes under method="distances" using the correlation (corr=TRUE) or

the absolute similarity of distances.

Details

Primarily designed to be used internally by fastAnc.

Value

A matrix in which the first column contains the nodes of tr1 with the second column containing matching nodes in tr2, with the criterion for matching defined by method.

Author(s)

Liam Revell < liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

mergeMappedStates 85

mergeMappedStates

Merge two or more mapped states into one state

Description

This function merges two or mapped states on the tree to get one new state. For instance, one could merge the states "C", "G", and "T" to get the state "not-A".

Usage

```
mergeMappedStates(tree, old.states, new.state)
```

Arguments

tree a modified object of class "phylo" or "multiPhylo" with a mapped discrete

character.

old.states states to merge.

new.state name for new state.

Value

A modified object of class "phylo" or "multiPhylo".

Author(s)

Liam Revell < liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

```
make.simmap, read.simmap
```

86 midpoint.root

midpoint.root

Midpoint root a phylogeny

Description

This function midpoint roots a rooted or unrooted tree (Farris 1972).

Usage

```
midpoint.root(tree)
```

Arguments

tree

an object of class "phylo".

Details

Midpoint rooting involves locating the midpoint of the longest path between any two tips and putting the root in that location. This function performs the same operation as midpoint in the phangorn package, but uses no phangorn code internally.

Value

A phylogenetic tree in "phylo" format.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Farris, J. (1972) Estimating phylogenetic trees from distance matrices. *American Naturalist*, **106**, 645-667.

Paradis, E., J. Claude, and K. Strimmer (2004) APE: Analyses of phylogenetics and evolution in R language. *Bioinformatics*, **20**, 289-290.

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

reroot, root

minRotate 87

vector	minRotate	Rotates all nodes of the tree to minimize the difference in order with a vector
--------	-----------	---

Description

This function rotates all the nodes of the tree to try and minimize the different between the order of the tips and the rank-order of a numeric vector x or (in the case of tipRotate the actual integer vector, x.

Usage

```
minRotate(tree, x, ...)
tipRotate(tree, x, ...)
```

Arguments

. . .

tree	tree.
x	numeric vector.

optional arguments to be used by tipRotate. Presently optional arguments can be fn, function to be used to compute the distance between the order of the tip labels in tree and the numeric vector x (presently fn=function(x) x^2 by default); methods, the methods of tree traversal (can be "pre", "post", or c("pre", "post"), for pre-, post-, or both pre- and post-order tree traversal); and print, a logical argument specifying whether to print the search progress or to behave quietly. Only the option print is available for minRotate.

Details

Primarily designed to be used internally by phylo.to.map, in the case of minRotate, or cophylo, in the case of tipRotate.

Value

```
A object of class "phylo".
```

Author(s)

```
Liam Revell liam.revell@umb.edu>
```

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

88 minSplit

minSplit	Finding the minimum (median) split in the posterior sample

Description

This function takes a phylogenetic tree and a list of splits and identifies the split with the smallest summed or summed squared distances to all the other splits. Used to be called min.split() but was changed to avoid conflict with the generic min.

Usage

```
minSplit(tree, split.list, method="sum", printD=FALSE)
```

Arguments

tree a phylogeny stored as an object of class "phylo".

split.list either a matrix with two named columns, "node" and "bp"; a \$mcmc matrix from

evol.rate.mcmc(); or the entire raw output from evol.rate.mcmc().

method an optional string indicating the criterion to minimize: options are "sum" and

"sumsq".

printD logical specifying whether to print distances to screen (FALSE by default).

Value

A list with the following components:

node node for the minimum split.

bp location on the branch leading to node of the minimum split.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

Revell, L. J., D. L. Mahler, P. Peres-Neto, and B. D. Redelings (2012) A new method for identifying exceptional phenotypic diversification. *Evolution*, **66**, 135-146.

See Also

```
evol.rate.mcmc, posterior.evolrate
```

mrp.supertree 89

mrp.supertree	Matrix representation parsimony supertree estimation

Description

This function estimates the MRP (matrix representation parsimony) supertree from a set of trees (Baum 1992; Ragan 1992).

Usage

```
mrp.supertree(trees, method=c("pratchet","optim.parsimony"), ...)
compute.mr(trees, type=c("phyDat","matrix"))
```

Arguments

```
an object of class "multiPhylo" (i.e., a list of trees).

method an argument specifying whether to optimize the tree using pratchet or optim. parsimony.

type for compute.mr, the type of object to return.

optional arguments - mostly to be passed to pratchet or optim. parsimony.
```

Details

Function uses pratchet or optim.parsimony from the "phangorn" package (Schliep 2011) and prop.part from the "ape" package (Paradis et al. 2004). See pratchet or optim.parsimony for optional arguments, which vary slightly depending on the method. All optional arguments of these methods are available to the user with one exception. The argument tree in optim.parsimony is supplied instead as start. In addition to being an object of class "phylo", start can also be assigned the string values of "NJ" or "random", in which case either a neighbor-joining or random tree will be used as the starting tree for optimization.

The function compute.mr computes the matrix-representation matrix of the input trees. It is used internally by mrp.supertree, but can also be used to export an object that can be written to file if desired.

Value

```
A "phylo" or "multiPhylo" object that is the MP or set of MP MRP trees.
In the case of compute.mr, an object of class "phyDat" or a matrix.
```

Author(s)

```
Liam Revell liam.revell@umb.edu>
```

90 multi.mantel

References

Baum, B. R., (1992) Combining trees as a way of combining data sets for phylogenetic inference, and the desirability of combining gene trees. *Taxon*, **41**, 3-10.

Felsenstein, J. (2004) Inferring Phylogenies. Sinauer.

Paradis, E., J. Claude, and K. Strimmer (2004) APE: Analyses of phylogenetics and evolution in R language. *Bioinformatics*, **20**, 289-290.

Ragan, M. A. (1992) Phylogenetic inference based on matrix representation of trees. *Molecular Phylogenetics and Evolution*, *1*, 53-58.

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

Schliep, K. P. (2011) phangorn: phylogenetic analysis in R. Bioinformatics, 27, 592-593.

See Also

exhaustiveMP, optim.parsimony, pratchet

multi.mantel

Multiple matrix regression (partial Mantel test)

Description

This function conducting a multiple matrix regression (partial Mantel test) and uses Mantel (1967) permutations to test the significance of the model and individual coefficients. It also returns the residual and predicted matrices.

Usage

```
multi.mantel(Y, X, nperm=1000)
```

Arguments

Y single "dependent" square matrix. Can be either a symmetric matrix of class

"matrix" or a distance matrix of class "dist".

X a single independent matrix or multiple independent matrices in a list. As with

Y can be a object of class "matrix" or class "dist".

nperm number of Mantel permutations.

Value

A list with the following components:

r. squared multiple R-squared.

coefficients model coefficients, including intercept.

tstatistic t-statistics for model coefficients.

multiC 91

fstatistic F-statistic for the overall model.

probt vector of probabilities, based on permutations, for tstatistic.

probF probability of F, based on Mantel permutations.

residuals matrix of residuals.

predicted matrix of predicted values.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Mantel, N. (1967) The detection of disease clustering and a generalized regression approach. *Cancer Research*, **27**, 209–220.

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

multiC

Returns a list with phylogenetic VCV matrix for each mapped state

Description

This function takes a modified "phylo" object as input and returns a set of so-called phylogenetic covariance matrices as a list: one for each mapped state.

Usage

```
multiC(tree,internal=FALSE)
```

Arguments

tree a phylogeny with mapped discrete state in a modified object of class "phylo"

(e.g., see read.simmap.

internal logical value indicating whether or not internal nodes should be returned.

Value

A list of matrices.

Author(s)

Liam Revell < liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

92 multiRF

See Also

evolvcv.lite, read.simmap, vcvPhylo, vcv.phylo

multiRF

Computes Robinson-Foulds distance between a set of trees

Description

Computes the Robinson-Foulds (Robinson & Foulds 1981) distance between a set of trees in an object of class "multiPhylo". If trees contains a large number of phylogenies (say 100 or 1,000) this will be much faster than calling RF.dist in the phangorn package for all pairwise comparisons because it avoids repeating some internal calculations. Nonetheless for large numbers multiRF is slow, and will use lots of memory.

Usage

multiRF(trees)

Arguments

trees

object of class "multiPhylo" consisting of two or more fully bifurcating, unrooted trees. If trees are rooted, they will be unrooted.

Details

Computes the Robinson-Foulds distance between all phylogenies in an object of class "multiPhylo". Uses prop.part internally for most of the heavy lifting.

Value

A distance matrix.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Robinson, D. R., Foulds, L. R. (1981) Comparison of phylogenetic trees. *Mathematical Biosciences*, **53**, 131-147.

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

nodeHeights 93

nodeHeights	Compute the heights above the root of each node	
-------------	---	--

Description

nodeHeights computes the height above the root for all nodes in the tree. nodeheight computes the height above the root for a single node.

Usage

```
nodeHeights(tree, ...)
nodeheight(tree, node, ...)
```

Arguments

a phylogeny as an object of class "phylo".

for nodeheight, the node for which we want to compute a height about

for nodeheight, the node for which we want to compute a height above the root

(or including the root edge, for root.edge=TRUE).

... optional arguments - presently only root.edge, a logical value indicating whether

or not to include the root edge length in the calculation of node heights.

Details

The function nodeHeights also gives a handy way to get the total length of the tree from the root to the heighest tip which will be given by max(nodeHeights(tree)). Generally speaking, nodeHeights will be faster if the heights of all or a large proportion of nodes is needed, wherease nodeheight will be faster if the height of one or a small number of nodes are needed.

Value

Either a matrix of the same dimensions as tree\$edge containing the height above the root of each node in edge (for nodeHeights); or a single positive number (for nodeheight).

Author(s)

```
Liam Revell liam.revell@umb.edu>
```

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

```
vcvPhylo
```

94 nodelabels.cophylo

Examples

```
tree<-rtree(10)
X<-nodeHeights(tree)</pre>
```

nodelabels.cophylo

Add labels to a plotted "cophylo" object

Description

This function adds node, edge, or tip labels to the plotted trees of a "cophylo" object.

Usage

```
nodelabels.cophylo(..., which=c("left","right"))
edgelabels.cophylo(..., which=c("left","right"))
tiplabels.cophylo(..., which=c("left","right"))
```

Arguments

... arguments to be passed to nodelabels, edgelabels, or tiplabels.

which argument indicated which of the two plotted trees (the "left" or "right" tree)

to be used.

Details

Note that the order of tips, edges, and nodes may be different in the object of class "cophylo" than they are in the original input trees, particularly if cophylo(...,rotate=TRUE) was used.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Paradis, E., J. Claude, and K. Strimmer (2004) APE: Analyses of phylogenetics and evolution in R language. *Bioinformatics*, **20**, 289-290.

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

```
cophylo, edgelabels, nodelabels, tiplabels
```

optim.phylo.ls 95

ontim	phylo.ls
OD CIIII.	DIIATO.T2

Phylogeny inference using the least squares method

Description

This function performs phylogeny inference using least-squares.

Usage

```
optim.phylo.ls(D, stree=NULL, set.neg.to.zero=TRUE, fixed=FALSE,
tol=1e-10, collapse=TRUE)
```

Arguments

D a distance matrix.

stree an optional starting tree for the optimization.

set.neg.to.zero

a logical value indicating whether to set negative branch lengths to zero (default

TRUE).

fixed a logical value indicating whether to estimate the topology - if TRUE only the

branch lengths will be computed.

tol a tolerance value used to assess whether the optimization has converged.

collapse a logical indicating whether to collapse branches with zero length.

Details

Function uses nni from the "phangorn" package (Schliep 2011) to conduct NNIs for topology estimation. Since topology optimization is performed using NNIs, converge to the true least-squares topology is not guaranteed. It is consequently probably wise to start with a very good tree - such as a NJ tree.

Value

An object of class "phylo" that (may be) the least-squares tree with branch lengths; also returns the sum of squares in attr(tree, "Q-score").

Author(s)

Liam Revell liam.revell@umb.edu>

96 orderMappedEdge

References

Cavalli-Sforza, L. L., and A. W. F. Edwards. (1967) Phylogenetic analysis: Modesl and estimation procedures. *American Journal of Human Genetics*, **19**, 233-257.

Felsenstein, J. (2004) Inferring Phylogenies. Sinauer.

Paradis, E., J. Claude, and K. Strimmer. (2004) APE: Analyses of phylogenetics and evolution in R language. *Bioinformatics*, **20**, 289-290.

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

Schliep, K. P. (2011) phangorn: phylogenetic analysis in R. Bioinformatics, 27, 592-593.

See Also

exhaustiveMP, nni

orderMappedEdge

Order the columns of mapped.edge to match across trees

Description

This function takes a modified object of class "multiPhylo" with a mapped discrete character (e.g., see read.simmap and sorts the columns of each tree\$mapped.edge to have the same state ordering. This is handy if we want to, for instance, run brownie.lite on a set of mapped trees, and then average the fitted parameter values across trees. The function also works for a single tree.

Usage

orderMappedEdge(trees, ordering=NULL)

Arguments

trees

object of class "phylo" or "multiPhylo".

ordering

ordering for the columns of \$mapped.edge. If NULL, then an alphabetical order is assumed. Options are "alphabetical", "numerical", or any specific

ordering of the mapped traits (e.g., c("A", "B", "C").

Value

A modified object of class "phylo" or "multiPhylo".

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

paintSubTree 97

paintSubTree	Paint sub-trees with a discrete character	
--------------	---	--

Description

This function maps or "paints" an arbitrary, i.e., user-specified, discrete character history on the tree. paintSubTree paints the clade downstream of node with a particular state; whereas paintBranches paints only a specified branch.

Usage

```
paintSubTree(tree, node, state, anc.state="1", stem=FALSE)
paintBranches(tree, edge, state, anc.state="1")
```

Arguments

tree	a phylogenetic tree as an object of class "phylo" or a modified object with mapped character traits.
node	an integer specifying the node number tipward of which the function should paint the derived state.
edge	an integer or vector of integers specifying the node or tip numbers of the edges that should be painted in paintBranches.
state	a string (or numeric value) specifying the state to paint on the tree tipward of node.
anc.state	the ancestral state to use; will only be applied if there are presently no character values mapped on the tree.
stem	logical or numeric value indicating whether to use the derived state on the stem leading to node (or not, if stem=FALSE), or, alternatively, what fraction of the stem should be assigned to the derived clade. Note that for tip clades stem=FALSE is not allowed.

Value

A modified phylogenetic tree of class "phylo" with the following additional elements:

maps a list of named vectors containing the times spent in each state on each branch,

in the order in which they occur.

mapped.edge a matrix containing the total time spent in each state along each edge of the tree.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

98 paste.tree

See Also

```
make.simmap, read.simmap, plotSimmap, sim.history
```

paste.tree

Paste two trees together

Description

Primarily internal function for posterior.evolrate; can be used to graft a clade into a receptor tree, at the "sticky tip" labeled with "NA".

Usage

```
paste.tree(tr1, tr2)
```

Arguments

tr1 receptor tree. tr2 donor clade.

Details

The donor clade needs to have a root edge, even if it is zero length.

Value

A tree.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

Examples

```
tr1<-rtree(10)
tr2<-rtree(10)
tr1$tip.label[1]<-"NA"
tr2$root.edge<-0
tr3<-paste.tree(tr1,tr2)</pre>
```

pbtree 99

pbtree

Simulate pure-birth or birth-death stochastic tree or trees

Description

This function simulates stochastic birth-death trees. Simulation can be performed conditioning on n, on t, or on both simultaneously. If the both, then (for optional argument method="rejection") rejection sampling is performed whereby trees are simulated given b and t until a tree containing n taxa is found. The giving-up point can be set using the optional argument max.count. Simulations can also be performed in continuous time (the default) or discrete time; the difference being that wait times in the continuous-time simulation come from the exponential distribution; whereas waiting times in discrete-time simulations come from the geometric distribution. In addition, discrete-time simulations allow for the possibility that multiple speciation events can occur at (exactly) the same time, so long as they are on separate branches. Finally, sometimes for stopping criterion n in discrete-time there will be a number of tips different from n. This indicates that the last event contained more than one speciation event, and a warning is printed.

method="direct" is presently experimental. It does not really perform direct sampling; however waiting times & birth or death events are sampled first - with only wait-times consistent with n and t being retained. This rejection sampling occurs one layer earlier than for method="rejection". This results in a significant (several-fold) speed-up of the code and enables sampling conditioned on n and t simultaneously for much higher b and d. At the present time, extant.only=TRUE does not work for this mode, nor does type="discrete".

Note that if ape=FALSE, then the function will run faster, and the tree is theoretically compatible with the ape "phylo" standard; however some downstream errors with functions such as bind. tree have been observed.

Usage

```
pbtree(b=1, d=0, n=NULL, t=NULL, scale=NULL, nsim=1,
type=c("continuous","discrete"), ...)
```

Arguments

b	birth rate or speciation rate for type="continuous"; the probability of speciating per time-step for type="discrete".
d	death rate or extinction rate for type="continuous"; the probability of going extinct per time-step for type="discrete".
n	desired number of species (i.e., taxa-stop criterion).
t	total time for simulation (i.e., time-stop criterion).
scale	if set, rescales tree to have total length scale.
nsim	number of simulated trees to return.
type	string to indicate whether to simulate trees in continuous or discrete time. If the former, then wait times between speciation events are drawn from an exponential distribution; whereas if the latter then wait times comes from a geometric distribution.

100 pgls.Ives

. . .

optional arguments including ape, a logical value indicating whether to return nodes in a 'ape' compatible ordering (default is TRUE); extant.only a logical value indicating whether or not to return only extant species (defaults to FALSE); max.count a numeric value indicating the maximum number of iterations to run is sampling conditioned on both n and t (defaults to 1e5); method gives the method used for simultaneously conditioning on n and t - options are "rejection" and "direct"; tip.label, a vector of tip labels (only works for n!=NULL); and, finally, quiet, a logical value indicating whether or not to suppress certain message (defaults to FALSE).

Value

A tree or list of trees as an object of class "phylo" or "multiPhylo", respectively.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

Examples

```
# simulate a pure-birth tree with 400 tips, scaled to a length of 1.0
tree<-pbtree(n=400,scale=1)
# simulate a pure-birth tree conditioning on n & t
tt<-log(50)-log(2)
tree<-pbtree(n=50,t=tt)</pre>
```

pgls.Ives

Phylogenetic regression with intraspecific sampling error

Description

This function fits the phylogenetic regression model with within-species sampling error following Ives et al. (2007).

Usage

```
pgls.Ives(tree, X, y, Vx=NULL, Vy=NULL, Cxy=NULL, lower=c(1e-8,1e-8))
```

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Arguments

tree a phylogeny as an object of class "phylo".

X a named vector containing a *single* independent variable (multiple independent

variables to be added in future). X can contain the species means, or a single long vector containing the sample of values for each species. In the latter case the names(X) will be repeating - all samples from the same species should have

the same name.

y vector the dependent variable. Can be species means or individual values, as for

Χ.

Vx sampling variances for X. If NULL, then the within-species variance is computed

from the data assuming that individual samples, not species means, have been

provided in X.

Vy sampling variances for y. If NULL, then the within-species variance is computed

from the data assuming that individual samples, not species means, have been

provided in y.

Cxy sampling covariances between X and y. This will also be computed from the

data if Cxy==NULL. Note than in this case - but not for the calculation of Vx and Vy, the same number of observations and the same ordering must be provided for X and y. If this is not the case, then it is assumed that different individuals have been sampled for X and y and thus Cxy is assumed to be zero for all species.

lower vector specifying the lower bounds for estimation for sig2x and sig2y, respec-

tively. (Must be >0.)

Details

Presently only the bivariate regression model is implemented. Uses optim(...,method="L-BFGS-B") for optimization. Note that some problems have been reported with the optimization algorithm for this model, which is simple and thus may fail to find the ML solution.

Value

A list with the following components:

beta a vector or matrix of regression coefficients.

sig2x fitted BM rate for X. sig2y fitted BM rate for y.

a fitted ancestral states for X and y.

logL log-likelihood.

convergence a value for convergence. convergence=0 is good; see optim for more details.

message a message for convergence.

Author(s)

Liam Revell liam.revell@umb.edu>

phenogram phenogram

References

Ives, A. R., P. E. Midford, and T. Garland Jr. (2007) Within-species measurement error in phylogenetic comparative methods. *Systematic Biology*, **56**, 252-270.

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

```
brownie.lite, phylosig, phyl.resid
```

phenogram

Plot phenogram (traitgram)

Description

Function plots a traitgram (Evans et al. 2009), that is, a projection of the phylogenetic tree in a space defined by phenotype (on the y axis) and time (on the x). If a discrete character is mapped on the tree this will also be plotted.

Usage

```
phenogram(tree, x, fsize=1.0, ftype="reg", colors=NULL, axes=list(), add=FALSE, \dots)
```

Arguments

tree	an object of class "phylo", with or without a mapped discrete character.
x	a vector containing the states at the tips or the states at all the tips and the internal nodes of the tree.
fsize	relative font size for tip labels.
ftype	font type - options are "reg", "i" (italics), "b" (bold), or "bi" (bold-italics).
colors	colors for plotting the mapped character (if available) in tree. If no character is mapped on the tree, then a single color for all the branches of the tree can be provided.
axes	list of axis dimensions. Items are time and trait.
add	optional logical value indicating whether to add to an open plot. If TRUE, then new axes will not be plotted.
	optional arguments including xlim, ylim, log, main, sub, xlab, ylab, asp, type, lty, lwd, offset, and digits are as in plot.default or par. Note that axes overrides xlim and ylim. spread.labels is a logical value indicating whether or not to minimize tip label overlap (default is TRUE); spread.cost is a numeric vector indicating the relative penalty to be used for label overlap and deviance, respectively (if spread.labels=TRUE); spread.range is the range over which to (potentially) spread the labels - note that if labels do not overlap,

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not all of that range will be used; finally, link is a numeric value by which to offset the tip labels, linking them to the tips with a dashed line (default is 0, if spread.labels=FALSE, or 10-percent of the total tree length otherwise). The optional argument offsetFudge "fudges" the computation of label offset in scaling xlim. It is 1.37, which is the correct fudge in the Windows R GUI, but this may need to be changed in other systems. hold indicates whether (or not) the output to the graphical device should be held using dev.hold before plotting (defaults to hold=TRUE). quiet suppresses some system messages if set to quiet=TRUE.

Details

For spread.labels=TRUE numerical optimization is performed to optimize the distribution of the labels vertically, where the solution depends on the vector spread.cost containing the cost of overlap (first) and the cost of deviation from the vertical position of the tip. Note that because this is done via numerical optimization, plotting may hang briefly while the best solution is found (especially for large trees).

Value

Plots a traitgram, optionally with a mapped discrete character, and (invisibly) returns a matrix containing the coordinates of the plotted tip labels.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Evans, M. E. K., Smith, S. A., Flynn, R. S., Donoghue, M. J. (2009) Climate, niche evolution, and diversification of the "bird-cage" evening primroses (Oenothera, sections Anogra and Kleinia). *American Naturalist*, **173**, 225-240.

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

Examples

```
## Not run:
tree<-pbtree(n=20,scale=2)
x<-fastBM(tree)
phenogram(tree,x)
# or, simulate a discrete character history
tree<-sim.history(tree,Q=matrix(c(-1,1,1,-1),2,2),anc="1")
# simulate in which the rate depends on the state
x<-sim.rates(tree,c(1,10))
phenogram(tree,x)
# now use spread.labels
tree<-pbtree(n=40)
x<-fastBM(tree)
phenogram(tree,x,spread.labels=TRUE,spread.cost=c(1,0))</pre>
```

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```
## End(Not run)
```

phyl.cca

Phylogenetic canonical correlation analysis

Description

This function performs phylogenetic canonical correlation analysis (e.g., Revell & Harrison 2008; *Bioinformatics*).

Usage

```
phyl.cca(tree, X, Y, lambda=1.0, fixed=TRUE)
```

Arguments

tree a phylogenetic tree in "phylo" format.

X a data matrix with traits in columns.

Y data matrix with traits in columns, to be correlated with X.

lambda optionally, a (fixed) value for lambda.

fixed optionally, a logical value indicating whether or not to estimate lambda using

likelihood.

Details

(Optional) joint optimization of λ is performed using optimize on the interval (0,1).

Value

A list with the following components:

cor canonical correlations.

xcoef coefficients for the canonical variables for X.
ycoef coefficients for the canonical variables for Y.
xscores matrix with the canonical scores for X.

yscores matrix with the canonical scores for Y.

chisq vector of χ^2 values.

p P-values for the hypothesis test that the *i*th and all subsequent correlations are

zero.

Author(s)

Liam Revell liam.revell@umb.edu>

phyl.pairedttest 105

References

Revell, L. J., Harrison, A. S. (2008) PCCA: A program for phylogenetic canonical correlation analysis. *Bioinformatics*, **24**, 1018-1020.

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

```
phyl.pca
```

nhv1	pairedttest
риут.	parreuttest

Phylogenetic paired t-test

Description

This function conducts a phylogenetic paired t-test, roughly following Lindenfors et al. (2010; *J. Evol. Biol.*). This is not a phylogenetic ANOVA, in which we want to compare the means of different sets of species on the tree. Instead, we are interested in the difference between two characters, or two measures of a character within a species, and we want to know if this difference is significantly different from zero controlling for the phylogenetic non-independence of species.

Usage

```
phyl.pairedttest(tree, x1, x2=NULL, se1=NULL, se2=NULL, lambda=1.0, h0=0.0,
fixed=FALSE)
```

Arguments

tree	a phylogeny as an object of class "phylo".
x1	data vector for first trait, or matrix with two traits in columns.
x2	data vector for second trait (or null if x1 is a matrix).
se1	standard errors for x1.
se2	standard errors for x2.
lambda	starting value for Pagel's lambda (or fixed value, if fixed=TRUE).
h0	null hypothesis (to be tested) for the mean difference between $x1$ and $x2$.
fixed	logical value specifying whether or not to optimize lambda.

Details

Likelihood optimization is performed using optim with method="L-BFGS-B" with box constraints on lambda (0,1).

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Value

A list with the following components:

dbar phylogenetic mean difference.

se standard error of dbar.

sig2 estimated evolutionary variance (of the difference).

lambda fitted (or fixed) value of lambda.
logL log-likelihood of the fitted model.

t.dbar t-value ((dbar-h0)/se where se is computed from the Hessian).

P. dbar P-value.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Lindenfors, P., L. J. Revell, and C. L. Nunn (2010) Sexual dimorphism in primate aerobic capacity: A phylogenetic test. *J. Evol. Biol.*, **23**, 1183-1194.

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

phyl.pca

Phylogenetic principal components analysis

Description

This function performs phylogenetic PCA (e.g., Revell 2009; Evolution).

Usage

```
phyl.pca(tree, Y, method="BM", mode="cov", ...)
## S3 method for class 'phyl.pca'
biplot(x, ...)
```

Arguments

tree phylogeny as an object of class "phylo".

Y data matrix with traits in columns.

method method to obtain the correlation structure: can be "BM" or "lambda".

mode is the mode for the PCA: can be "cov" or "corr".

x object of class "phyl.pca" for biplot.phyl.pca.

for S3 plotting method biplot.phyl.pca, other arguments to be passed to

biplot.

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Details

If method="lambda" λ is optimized on the interval (0,1) using optimize. S3 methods (print, summary, and biplot) are modified from code provided by Joan Maspons and are based on the same methods for objects of class "prcomp". Function biplot now permits the argument choices to be supplied, which should be a vector of length two indicated the two PC axes to be plotted.

Value

An object of class phyl.pca which is a list with some or all of the following components:

Eval diagonal matrix of eigenvalues.

Evec matrix with eigenvectors in columns.

S matrix with scores.
L matrix with loadings.

lambda fitted value of lambda (method="lambda" only).

logL log-likelihood for lambda model (method="logL" only).

Author(s)

Liam Revell liam.revell@umb.edu>, Joan Maspons

References

Revell, L. J. (2009) Size-correction and principal components for interspecific comparative studies. *Evolution*, **63**, 3258-3268.

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

```
phyl.cca, phyl.resid
```

phyl.resid

Phylogenetic size-correction via GLS regression

Description

This function fits one or multiple phylogenetic regressions (depending on the number of columns in Y) and computes the residuals. Designed for phylogenetic size correction using GLS regression (e.g., Revell 2009; *Evolution*).

Usage

```
phyl.resid(tree, x, Y, method="BM")
```

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Arguments

tree	a phylogenetic tree in	"phylo" format.
------	------------------------	-----------------

x vector containing the single independent variable (e.g., size), or matrix with

multiple independent variables in columns.

Y vector or matrix with one or multiple dependent variables in columns.

method method to obtain the correlation structure: can be "BM" or "lambda".

Details

Optionally fits λ for each regression model. Likelihood optimization of λ is performed for method="lambda" using optimize on the interval (0,1). This function is redundant with residuals(gls(...,correlation=corPagel(...))] but some users may find this method simpler, and it provides a good way to cross-check your results & make sure that you are using gls correctly.

Value

A list with the following components:

beta a vector or matrix of regression coefficients.

resid a vector or matrix of residuals for species.

lambda a vector of lambda values (method="lambda" only).

logL a vector of log-likelihoods (method="lambda" only).

Author(s)

Liam Revell < liam.revell@umb.edu>

References

Revell, L. J. (2009) Size-correction and principal components for interspecific comparative studies. *Evolution*, **63**, 3258-3268.

Revell, L. J. (2010) Phylogenetic signal and linear regression on species data. *Methods in Ecology and Evolution*, **1**, 319-329.

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

```
phyl.pca, gls
```

phyl.RMA

phyl.RMA	Phylogenetic reduced major axis (RMA) regression

Description

This function performs phylogenetic RMA regression.

Usage

```
phyl.RMA(x, y, tree, method="BM", lambda=NULL, fixed=FALSE, h0=1.0)
```

Arguments

x vector with names.
y vector with names.

tree a phylogenetic tree in "phylo" format.

method method to obtain the correlation structure: can be "BM" or "lambda".

lambda value of lambda for fixed λ .

fixed logical value indicating whether or not λ should be optimized using likelihood. h0 null hypothesis for beta. Defaults to 1.0. Note that a null hypothesis of 0.0 is

not allowed.

Details

Optionally jointly estimates lambda if method="lambda". Likelihood optimization of lambda is performed using optimize on the interval (0,1). Note that some statistician think there is *never* a condition in which a reduced-major-axis regression should be used.

Value

A list with the following components:

RMA. beta a vector of RMA regression coefficients.

V a VCV matrix for the traits.

lambda fitted value of lambda (method="lambda" only).

logL log-likelihood (method="lambda" only).

test a vector containing results for hypothesis tests on beta.

resid a vector of residuals for y given x.

Author(s)

phyl.vcv

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

```
phyl.cca, phyl.pca, phyl.resid
```

phyl.vcv

Compute evolutionary VCV matrix for a tree & dataset

Description

Primarily an internal function for phyl.pca; this can be used to compute the phylogenetic trait variance-covariance matrix given a phylogenetic VCV, lambda, and a data matrix. Should not be confused with vcv.phylo in the "ape" package (although one of the objects returned is the output of vcv.phylo).

Usage

```
phyl.vcv(X, C, lambda)
```

Arguments

lambda value for λ transformation.

X data matrix.

C matrix containing the height above the root of each pair of species in the tree.

Typically this will have been produced by calling vcv.phylo.

Details

Do not use unless you know what you're doing.

Value

A list containing three elements, as follows: C, the matrix vcv.phylo transformed by lambda; R, the among trait variance-covariance matrix for the data in X; and alpha, a vector of ancestral states at the root node of the tree.

Author(s)

Liam Revell < liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

phylANOVA 111

phylANOVA	Phylogenetic ANOVA and post-hoc tests

Description

This function performs the simulation-based phylogenetic ANOVA of Garland et al. (1993) and (optionally) conducts all posthoc comparisons of means among groups (also obtaining the P-values by phylogenetic simulation).

Usage

```
phylANOVA(tree, x, y, nsim=1000, posthoc=TRUE, p.adj="holm")
```

Arguments

tree	a phylogenetic tree in "phylo" format.
x	a vector containing the groups.
у	a vector containing the response variable (continuously valued).
nsim	an integer specifying the number of simulations (including the observed data).
posthoc	a logical value indicating whether or not to conduct posthoc tests to compare the mean among groups.
p.adj	method to adjust P-values for the posthoc tests to account for multiple testing. Options same as p.adjust.

Details

Uses a little bit of code from phy. anova in the "geiger" package as well as pairwise.t.test.

Value

A list containing the following elements:

F	F from observed data.
Pf	P-value for F from simulation.
Т	matrix of t-values.
Pt	matrix of multiple test corrected P-values from posthoc t-tests.

Author(s)

phylo.heatmap

References

Garland, T., Jr., A. W. Dickerman, C. M. Janis, & J. A. Jones. (1993) Phylogenetic analysis of covariance by computer simulation. *Systematic Biology*, **42**, 265-292.

Harmon, L. J., J. T. Weir, C. D. Brock, R. E. Glor, W. Challenger. (2008) GEIGER: investigating evolutionary radiations. *Bioinformatics*, 24, 129-131.

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

```
anova, pairwise.t.test
```

phvlo.	heatmap

Creates a phylogenetic heat map

Description

Functions creates a multivariate phylogenetic heatmap.

Usage

```
phylo.heatmap(tree, X, fsize=1, colors=NULL, standardize=FALSE, ...)
```

Arguments

tree	an object of class "phylo".
X	a matrix containing data for multiple continuous characters in which rownames correspond to the tip labels of the tree.
fsize	an integer or vector of length 3 containing the font size for the tip labels, the trait labels, and the legend text. (If a single integer is supplied, then the value will be recycled.)
colors	a vector of colors to be passed to image. Can be a function call (e.g., heat.colors(n=200)[200:1]).
standardize	a logical value indicating whether or not to standardize each column of X to have the same variance & mean prior to analysis.
	optional arguments. So far these include: legend, a logical value indicating whether or not to plot a figure legend (defaults to legend=TRUE); labels, a logical value indicating whether or not to plot trait labels (defaults to labels=TRUE); split, a numeric vector indicating the fraction of the horizontal dimension to use for the tree & heatmap, respectively (defaults to split=c(0.5,0.5)); xlim, ylim, & mar, defined as in par; and ftype, lwd, and pts as defined in plotSimmap.

Value

Function creates a plot.

113 phylo.to.map

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). Methods Ecol. Evol., 3, 217-223.

phylo.to.map

Plot tree with tips linked to geographic coordinates

Description

Function plots a tree and tips pointing to coordinates on a global map.

Usage

```
phylo.to.map(tree, coords, rotate=TRUE, ...)
## S3 method for class 'phylo.to.map'
plot(x, type=c("phylogram","direct"), ...)
```

Arguments

tree an object of class "phylo".

a matrix containing the latitude (in column 1) and the longitude of all tip species coords

in the tree. The row names should be the same as tree\$tip.label.

rotate a logical value indicating whether or not to rotate nodes of the tree to better

match longitudinal positions.

for plot.phylo.to.map, an object of class "phylo.to.map". Χ

a string indicating whether to map the tips of the tree onto a geographic map type

from a square phylogram (type="phylogram") or to project the tree directly

onto the map (type="direct").

optional arguments. For phylo. to. map, which creates an object of class "phylo. to. map"

and (optionally) plots that object, arguments include: database and regions (see map), as well as any arguments that should be passed to plot.phylo.to.map internally. For phylo.to.map, optional arguments xlim and ylim, which control the plot area for the map; fsize for the font size of plot labels and ftype for the font type (following plotSimmap; split which controls the proportion of vertical (or horizontal) space for the tree (first) and map, in a vector; psize the size of the plotted points on the map - or cex. points, a vector contain the size of the tip points and geographic coordinate points, respectively; from tip a logical value indicating whether to plot the linking lines from the tips (if TRUE) or from the end of the tip label, the default; colors, a single value or a vector of colors for the points and the linking lines; pch a single value or a vector of point types; 1wd and 1ty for the linking lines; and pts a logical value indicating whether or not to plot points at the tips of the tree. mar and asp are as in par.

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Value

phylo.to.map creates an object of class "phylo.to.map" and (if plot=TRUE) plots a phylogeny projected onto a geographic map.

plot.phylo.to.map plots on object of class "phylo.to.map".

Author(s)

Liam Revell < liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). Methods Ecol. Evol., 3, 217-223.

phylo.toBackbone

Converts tree to backbone or vice versa

Description

Converts between "phylo" and "backbonePhylo".

Usage

```
phylo.toBackbone(x, trans, ...)
backbone.toPhylo(x)
```

Arguments

an object of class "phylo" (for phylo. toBackbone); or an object of class backbone. toPhylo Χ

(for backbone.toPhylo).

trans data frame containing the attributes necessary to translate a backbone tree to an

> object of class "backbonePhylo". The data frame should contain the following variables: tip.label: the tip labels in the input tree (not all need be included); clade.label: labels for the unobserved subtrees; N: number of species in each subtree; and depth: desired depth of each subtree. depth for each terminal

taxon in x cannot be greater than the terminal edge length for that taxon.

optional arguments. . . .

Value

Either an object of class "phylo" or an object of class "backbonePhylo", depending on the method.

Author(s)

phyloDesign 115

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

plot.backbonePhylo

phyloDesign

Compute design matrix for least squares analyses

Description

Primarily an internal function for optim.phylo.ls, this function creates a design matrix for least squares phylogenetic analysis.

Usage

phyloDesign(tree)

Arguments

tree

phylogenetic tree.

Details

This function returns a matrix containing the edges in the tree (in columns) and pairs of tip node numbers (in rows). Values in the matrix are either 1 if the edge is on the shortest path between the two tips; and 0 otherwise. Probably do not use unless you know what you're doing.

Value

A matrix.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

116 phylomorphospace

phylomorphospace

Creates phylomorphospace plot

Description

This function creates a phylomorphospace plot (a projection of the tree into morphospace) for two characters following Sidlauskas (2008; Evolution). It will also plot a discrete character mapped on tree.

Usage

```
phylomorphospace(tree, X, A=NULL, label=c("radial","horizontal","off"),
control=list(), ...)
```

Arguments

tree	a phylogenetic tree in "phylo" format, or a modified "phylo" object with a
	mapped discrete character.

Χ an n x 2 matrix of tip values for two characters in n species.

Α an optional m x 2 matrix (for m nodes) of values for two taits at internal nodes in

the tree - if not supplied, these values will be estimated using fastAnc.

string indicating whether to plot the tip labels in the same direction as the terlabel

> minal edge (label="radial"), horizontally label="horizontal", or not at all "off". label=TRUE and label=FALSE are also acceptable, for compatibility

with phytools $\leq 0.3-03$.

a list containing the following optional control parameters: col.edge: a vector control

of edge colors; and col. node: a vector of node colors.

optional arguments for plotting, including xlim, ylim, xlab, ylab, lwd, colors, fsize, and node.by.map. colors is only used when there is a mapped discrete character on the tree, in which case control\$col.edge is ignored. fsize is relative to the default, which is textxy(...,cx=0.75). node.by.map is a logical value (defaults to FALSE which tells the function whether or not to plot the node colors using the colors of the mapped discrete character. Setting this option to TRUE will cause control\$col.node to be ignored. node.size is a vector containing the point size relative to the default (see par for plotted internal nodes and tips, respectively. Defaults to node.size=c(1,1.3). If only one number is provided it will be recycled. axes is a logical value indicating whether or not axes should be plotted (see plot.default. Finally, add indicates whether to

add the phylmorphospace to the current plot.

Value

This function creates a phylomorphospace plot

Author(s)

phylomorphospace3d 117

References

Paradis, E., J. Claude, and K. Strimmer (2004) APE: Analyses of phylogenetics and evolution in R language. *Bioinformatics*, **20**, 289-90.

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

Sidlauskas, B. (2008) Continuous and arrested morphological diversification in sister clades of characiform fishes: A phylomorphospace approach. *Evolution*, **62**, 3135-3156.

Examples

```
tree<-pbtree(n=25)
X<-fastBM(tree,nsim=2)
phylomorphospace(tree,X,xlab="trait 1",ylab="trait 2")</pre>
```

phylomorphospace3d

Creates tree-dimensional phylomorphospace plot

Description

This function creates a phylomorphospace plot for three characters using the 3D visualization package, 'rgl'.

Usage

```
phylomorphospace3d(tree, X, A=NULL, label=TRUE, control=list(),
method=c("dynamic","static"), ...)
```

Arguments

8	
tree	a phylogenetic tree in "phylo" format.
Χ	an n x 3 matrix of tip values for two characters in n species.
Α	an optional m \times 3 matrix (for m nodes) of values for two taits at internal nodes in the tree - if not supplied, these values will be estimated using anc.ML.
label	logical value indicating whether to print tip labels next to terminal nodes in the plot (presently doesn't do anything, but labels can be dropped using control). spin=TRUE,axes=TRUE,box=TRUE,simple.axes=FALSE,lwd=1,ftype="reg"
control	a list containing the following optional control parameters: spin: a logical value indicating whether to animate the plot when created; axes: a logical indicating whether to plot the axes; box: a logical value indicating whether to plot in box; simple.axes: logical value indicating whether to replace box and axes with simpler axes; lwd: line widths; ftype: font type ("off" turns off labels altogether); col.edge a vector of colors of length nrow(tree\$edge).
method	a string either "dynamic" for a dynamic (animated) plot created using rgl; or "static" for a flat 3D plot created using scatterplot3d and base graphics. The latter has the advantage of being very easy to export in standard format.
•••	optional arguments to be passed to scatterplot3d. Most options not available. angle is an important option that does work here.

phylosig phylosig

Value

This function creates a three dimensional phylomorphospace plot. The function returns a function from spin3d (for method="dynamic"); or a series of functions from scatterplot3d (for method="static").

Author(s)

Liam Revell < liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

Sidlauskas, B. (2008) Continuous and arrested morphological diversification in sister clades of characiform fishes: A phylomorphospace approach. *Evolution*, **62**, 3135-3156.

See Also

fancyTree, phenogram, phylomorphospace

Examples

```
tree<-pbtree(n=26,tip.label=LETTERS)
X<-fastBM(tree,nsim=3)
## Not run:
phylomorphospace3d(tree,X,control=list(spin=FALSE))
## End(Not run)
phylomorphospace3d(tree,X,method="static")</pre>
```

phylosig

Compute phylogenetic signal with two methods

Description

This function computes phylogenetic signal using two different methods. It can also conduct the hypothesis tests for significant phylogenetic signal, and estimate phylogenetic signal incorporating sampling error following Ives et al. (2007).

Usage

```
phylosig(tree, x, method="K", test=FALSE, nsim=1000, se=NULL, start=NULL,
control=list())
```

phylosig 119

Arguments

tree a phylogenetic tree in "phylo" format.

x vector containing values for a single continuously distributed trait.

method method to compute signal: can be "K" or "lambda".

test logical indicating whether or not to conduct a hypothesis test of "K" or "lambda".

nsim for method="K", number of simulations in randomization test.

se named vector containing the standard errors for each species.

start vector of starting values for optimization of (respectively) sigma^2 and lambda.

Only used in method="lambda" and se!=NULL.

control list of control parameters for multidimensional optimization, implemented in

optim. Only used in method="lambda" and se!=NULL.

Details

 λ optimization is performed using optimize with the range of lambda set between 0 and the theoretical upper limit of lambda (determined by the relative height of the most recent internal node on the tree).

Value

If (method="K"), a list with the following components:

K value of the K-statistic.

sig2 rate of evolution, σ^2 , for estimation with sampling error.

logL log-likelihood, for estimation with sampling error.

P optionally, the P-value from the randomization test.

If (method="lambda"), a list with the following components:

lambda fitted value of lambda.

sig2 rate of evolution, for estimation with sampling error.

logL log-likelihood.

logL0 log-likelihood for lambda=0.0.

P P-value of the likelihood ratio test.

convergence value for convergence, for estimation with sampling error only. (See optim).

message from optim, for estimation with sampling error only.

Author(s)

120 plot.backbonePhylo

References

Blomberg, S. P., T. Garland Jr., A. R. Ives (2003) Testing for phylogenetic signal in comparative data: Behavioral traits are more labile. *Evolution*, **57**, 717-745.

Ives, A. R., P. E. Midford, T. Garland Jr. (2007) Within-species variation and measurement error in phylogenetic comparative biology. *Systematic Biology*, **56**, 252-270.

Pagel, M. (1999) Inferring the historical patterns of biological evolution. *Nature*, **401**, 877-884.

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

Examples

```
tree<-pbtree(n=100)
x<-fastBM(tree)
phylosig(tree,x,method="lambda",test=TRUE)</pre>
```

plot.backbonePhylo

Plots backbone tree with triangles as clades

Description

Function plots a backbone tree (stored as an object of class "backbonePhylo") with triangles as subtrees.

Usage

```
## S3 method for class 'backbonePhylo' plot(x, ...)
```

Arguments

x an object of class "backbonePhylo".

optional arguments. So far includes only vscale, which is used to rescale the vertical dimension in plotting.

Value

Plots a tree.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

plotBranchbyTrait 121

See Also

```
phylo.toBackbone
```

Examples

```
## first create our backbone tree with
## random subtree diversities
tree<-phytools:::lambdaTree(pbtree(n=10),lambda=0.5)</pre>
## for old versions of geiger, use lambdaTree
## create a translation table
## leaving a couple of single-taxon clades for fun
tip.label<-sample(tree$tip.label,8)</pre>
clade.label<-LETTERS[1:8]</pre>
N<-ceiling(runif(n=8,min=1,max=20))
## set crown node depth to 1/2 the maximum depth
depth<-sapply(tip.label,function(x,y)</pre>
0.5*y$edge.length[which(tree$edge[,2]==
which(y$tip.label==x))],y=tree)
trans<-data.frame(tip.label,clade.label,N,depth)</pre>
rownames(trans)<-NULL</pre>
rm(tip.label,clade.label,N,depth)
## here's what trans looks like
trans
## convert
tt<-phylo.toBackbone(tree,trans)
## plot
plot(tt)
```

plotBranchbyTrait

Plot branch colors by a quantitative trait or value

Description

Function plots a tree with branches colored by the value for a quantitative trait or probability, by various methods. Unlike most other tree plotting functions in phytools, this function calls plot.phylo (not plotSimmap) internally.

Usage

```
plotBranchbyTrait(tree, x, mode=c("edges","tips","nodes"), palette="rainbow",
legend=TRUE, xlims=NULL, ...)
```

Arguments

```
tree an object of class "phylo".

x either a vector of states for the edges, tips, or nodes of the tree (for mode="edges", "tips", and "nodes", respectively).
```

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string indicating plotting mode. mode="edges", the default, requires that the mapping state of each edge in the tree should be provided. mode="tips" takes the tip values and estimates the state at each internal node. The mapped character value along each branch is the average of the nodes subtending that branch. mode="nodes" similar to "tips", except that the node values are provided instead of estimated.

color palette to translate character values to color. Options are presently "rainbow"

(the default), "heat.colors", and "gray".

legend can be a logical value (TRUE or FALSE) or a numeric value greater than 0. In the

latter case the numeric value gives the length of the plotted legend, which also

acts as a scale bar for the branch lengths of the tree.

xlims range for the translation map between trait values and the color map. Should be

inclusive of all the values in x.

... other optional arguments to be passed to plot.phylo - pretty much all arguments are qualible. In addition, there also provide her the follows:

ments are available. In addition, there plotBranchbyTrait has the following additional optional arguments: tol a small tolerance value to be added to the range of x; prompt for legend=TRUE, a logical value indicating whether to prompt for the position of the legend (or not) - the default is to put the legend in the lower left hand size of the plot; title for legend=TRUE, the title of the legend; and digits for legend=TRUE, the number of digits in the quantitative

scale of the legend.

Details

palette

Note that if prompt=TRUE, the function will prompt for the position of the legend.

Value

Plots a phylogeny.

Author(s)

Liam Revell < liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

plotSimmap	Plot stochastic character mapped tree	

Description

Function plots one or multiple stochastic character mapped trees.

plotSimmap 123

Usage

```
plotSimmap(tree, colors=NULL, fsize=1.0, ftype="reg", lwd=2, pts=FALSE,
node.numbers=FALSE, mar=NULL, add=FALSE, offset=NULL, direction="rightwards",
type="phylogram", setEnv=TRUE, part=1.0, xlim=NULL, ylim=NULL,
nodes="intermediate", tips=NULL, maxY=NULL, hold=TRUE, split.vertical=FALSE)
## S3 method for class 'simmap'
plot(x, ...)
## S3 method for class 'multiSimmap'
plot(x, ...)
```

Arguments

a modified object of class "phylo" or "multiPhylo" containing a stochastic mapping or set of mappings (e.g., see read.simmap & make.simmap).
a vector with names translating the mapped states to colors - see Examples.
relative font size for tip labels.
font type - options are "reg", "i" (italics), "b" (bold), or "bi" (bold-italics).
line width for plotting.
logical value indicating whether or not to plot filled circles at each vertex of the tree, as well as at transition points between mapped states. Default is FALSE.
a logical value indicating whether or not node numbers should be plotted.
vector containing the margins for the plot to be passed to par. If not specified, the default margins are [0.1,0.1,0.1].
a logical value indicating whether or not to add the plotted tree to the current plot (TRUE) or create a new plot (FALSE, the default).
offset for the tip labels. Primarily to be used internally by densityMap.
plotting direction. Options are "rightwards" (the default) and "leftwards". Note that for some reason that is not totally clear, node.numbers=TRUE does not work for direction="leftwards".
plot type. Can be "phylogram" or "fan". Only a subset of options are presently available for type="fan".
logical value indicating whether or not to set the environment .PlotPhyloEnv. Setting this to TRUE (the default) will allow compatibility with ape labeling functions such as nodelabels.
value between 0 and 1 for type="fan" indicating what fraction of the full circular tree to use as plotting area. For instance, part=0.5 will plot a half fan phylogeny. It also affects the axis scaling used.
x-limits for the plot.
y-limits for the plot.
node placement following Felsenstein (2004; pp. 574-576). Can be "intermediate", "centered", "weighted", or "inner". So far only works for type="phylogram".
labeled vector containing the vertical position of tips. Normally this will be 1:N for N tips in the tree.

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maxY	maximum value of y to use before rotating a tree into fan configuration. This will only make a difference if different from Ntip(tree).
hold	logical argument indicating whether or not to hold the output to the graphical device before plotting. Defaults to hold=TRUE.
split.vertical	split the color of the vertically plotted edges by the state of the daughter edges. Only applies if the edge state changes exactly at a node.
x	for S3 plotting method, object of class "simmap" or "multiSimmap".
	for S3 plotting method, other arguments to be passed to plotSimmap.

Details

The underscore character "_" is automatically swapped for a space in tip labels, as in plot.phylo.

Value

Plots a tree.

Author(s)

Liam Revell < liam.revell@umb.edu>

References

Bollback, J. P. (2006) Stochastic character mapping of discrete traits on phylogenies. *BMC Bioinformatics*, 7, 88.

Felsenstein, J. (2004) Inferring Phylogenies. Sinauer.

Huelsenbeck, J. P., R. Neilsen, and J. P. Bollback. (2003) Stochastic mapping of morphological characters. *Systematic Biology*, **52**, 131-138.

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

```
densityMap, make.simmap, read.simmap
```

Examples

```
# simulate a mapped tree
Q<-matrix(c(-2,1,1,1,-2,1,1,1,-2),3,3)
rownames(Q)<-colnames(Q)<-letters[1:3]
tree<-sim.history(pbtree(n=100,scale=1),Q)
cols<-setNames(c("blue","red","green"),letters[1:3])
# plot the mapping
plot(tree,cols,ftype="i",fsize=0.7)</pre>
```

plotThresh 125

plotThresh	Tree plotting with posterior probabilities of ancestral states from the threshold model

Description

This function uses the object returned by ancThresh to plot the posterior probabilities of ancestral states under the threshold model. It is also called internally by ancThresh.

Usage

```
plotThresh(tree, x, mcmc, burnin=NULL, piecol, tipcol="input", legend=TRUE, ...)
```

Arguments

tree	phylogenetic tree.
X	a named vector containing discrete character states; or a matrix containing the tip species, in rows, and probabilities of being in each state, in columns.
mcmc	list object returned by ancThresh.
burnin	number of generations (not samples) to exclude as burn in; if NULL then 20 percent of generations are excluded as burn-in.
piecol	a named vector containing the colors for the posterior probabilities plotted as pie charts at internal nodes.
tipcol	a string indicating whether the tip colors should be based on the input data ("input") or sampled tip liabilities ("estimated"). These will only differ if there is uncertainty in the tip states.
legend	logical value or text to be plotted in the legend.
	other arguments to be passed to plot.phylo - label.offset should be >0 so that tip labels and species names do not overlap.

Value

Plots a tree.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

Revell, L. J. (2014) Ancestral character estimation under the threshold model from quantitative genetics. *Evolution*, bold68, 743-759.

plotTree

See Also

```
ancThresh, plot.phylo
```

plotTree

Plots rooted phylogenetic tree

Description

This function plots a rooted phylogram. Arguments in ... are passed to plotSimmap, with the exception of optional argument color which is used to determine the plotted color of the branch lengths of the tree.

Usage

```
plotTree(tree, ...)
```

Arguments

tree a phylogenetic tree in "phylo" format; or multiple trees as an object of class

"multiPhylo".

... optional arguments.

Value

This function plots a rooted phylogram.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

```
plot.phylo, plotSimmap
```

Examples

```
tree<-pbtree(n=25)
plotTree(tree,color="blue",ftype="i")</pre>
```

plotTree.wBars 127

plotTree.wBars	Plot a tree with bars at the tips
----------------	-----------------------------------

Description

Function plots a phylogeny in phylogram or fan style with bars at the tips representing the values for a phenotypic trait.

Usage

```
plotTree.wBars(tree, x, scale=1, width=NULL, type="phylogram", method="plotTree",
tip.labels=FALSE, col="grey", border=NULL, ...)
```

Arguments

tree	an object of class "phylo".
X	a named vector of trait values (normally > 0).
scale	scaling factor for the tip bars (relative to the total tree height).
width	width of the tip bars.
type	plot type. Can be "phylogram" or "fan".
method	plotting method to use. Can be "plotTree" (for plotTree) or "plotSimmap" (for plotSimmap).
tip.labels	argument indicating whether or not tip labels should be plotted. Defaults to tip.labels=FALSE.
col	colors of the plotted bars. Can be a single value or a vector with length equal to the number of tips in the tree.
border	single value specifying the color of the border for the plotted bars. Defaults to border=NULL, which means that black borders will be plotted.
	optional arguments to be passed to codeplotTree or plotSimmap.

Value

Plots a tree.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

```
plotSimmap, plotTree
```

128 posterior.evolrate

posterior.evolrate

Analysis of the posterior sample from evol.rate.mcmc

Description

This function takes a phylogenetic tree, an average split position, and a raw MCMC output from evol.rate.mcmc and returns a posterior sample of evolutionary rates rootward $(\sigma(1)^2)$ and tipward $(\sigma(2)^2)$ from the average split.

Usage

```
posterior.evolrate(tree, ave.shift, mcmc, tips, showTree=FALSE)
```

Arguments

tree a phylogenetic tree in "phylo" format.

ave.shift mean or median shift-point from the posterior sample (see minSplit.

mcmc matrix \$mcmc from evol.rate.mcmc (probably with burnin excluded).

tips list of stips in state $sig(1)^2$ for each sampled generation of MCMC.

showTree optional logical value indicating whether or not to plot the stretched and shrunken

tree generated by the pre-processing algorithm implemented in this function (de-

fault is FALSE).

Value

A matrix containing the posterior sample of evolutionary rates and shift-points between rates.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J., D. L. Mahler, P. Peres-Neto, and B. D. Redelings (2012) A new method for identifying exceptional phenotypic diversification. *Evolution*, **66**, 135-146.

See Also

```
evol.rate.mcmc, minSplit
```

print.backbonePhylo 129

print.backbonePhylo

Print method for backbone phylogeny

Description

Print method for an object of class "backbonePhylo".

Usage

```
## S3 method for class 'backbonePhylo' print(x, ...)
```

Arguments

x an object of class "backbonePhylo".

... optional arguments.

Value

Prints to screen.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

```
phylo.toBackbone
```

ratebystate

Method for investigating the rate of one trait as a function of the state of another

Description

This function attempts to ask if the rate of a continuous character, y, depends on the state of a separate continuous trait, x. This is accomplished by regressing the squared contrasts in y on the branch or node ancestral estimates of x.

ratebystate

Usage

```
ratebystate(tree, x, y, nsim=100, corr=c("pearson", "spearman"), ...)
```

Arguments

tree	phylogenetic tree.

x a continuous character - the dependent variable in the model.

y a second continuous trait - the response variable.

nsim number of simulations for hypothesis testing.

corr correlation method to use. Same as in cor.

optional arguments which include sim.method ("fastBM" or "sim.corrs";

see fastBM and sim.corrs); method ("by.node" or "by.branch" indicating whether to assume the rate varies as a function of the node state or the mean branch state); message - a logical value indicating whether or not to return corr and method; finally logarithm - indicating whether or not to fit a model in which the variance of Brownian evolution in y changes as a multiplicative func-

tion of x. The default is logarithm=FALSE.

Value

This function returns a list with up to the following four elements:

beta value of the regression coefficient for square of the contrasts in y regressed on

the ancestral or branch-wise estimated states for x.

r correlation coefficient for corr=corr.

corr string giving the value of corr.
method string giving the value of method.

Author(s)

Liam Revell < liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

fastAnc, pic

rateshift 131

rateshift	Find the temporal position of one or more rate shifts
Tateshirt	Tina the temporal position of one of more rate shifts

Description

Function finds the location of one or more rate shifts.

Usage

```
rateshift(tree, x, nrates=1, niter=10, method="ML", ...)
## S3 method for class 'rateshift'
plot(x, ...)
```

Arguments

tree object of class "phylo".

x vector of phenotypic trait values for species. names(x) should contain the

species names and match tree\$tip.label.

nrates number of rates.

niter number of iterations of optimization routine to ensure convergence.

method optimization method. Can be "ML" (maximum likelihood) or "REML" (restricted

maximum likelihood).

... optional arguments. In the case of the plot method, these will be passed to

plotSimmap. For rateshift, optional arguments include: tol, tolerance; plot & print, logical values indicating whether to plot or print the progress of the optimization (default to FALSE); quiet, logical argument indicating whether to suppress all notifications (defaults to FALSE); minL, numeric value; and fixed.shift,

either a vector of fixed shift points, or a logical value.

Value

An object of class "rateshift".

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

```
brownie.lite
```

132 read.newick

read	. r	ıew	1 C	k

Robust Newick style tree reader

Description

This function reads a Newick style tree from file.

Usage

```
read.newick(file="", text, ...)
```

Arguments

file name of text file with single Newick style tree or multiple trees, one per line.

text character string containing tree.

optional arguments to be passed to scan. Note that if the arguments sep

optional arguments to be passed to scan. Note that if the arguments sep or what are supplied this could generate an error. Useful optional arguments might include skip (number of lines to skip) and nlines (number of lines to read).

Details

This function is almost completely redundant with read. tree; however it is 'robust' in that it does not fail if the tree contains so-called 'singles' (nodes with only one descendant).

Value

An object of class "phylo", possibly containing singles (see collapse.singles).

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

```
read.tree, read.nexus
```

Examples

```
tree<-"((Human,Chimp),Gorilla),Monkey);"
phy<-read.newick(text=tree)</pre>
```

read.simmap 133

	Read SIMMAP style trees from file	read.simmap
--	-----------------------------------	-------------

Description

This reads one or multiple SIMMAP style trees from file.

Usage

```
read.simmap(file="", text, format="nexus", rev.order=TRUE, version=1)
```

Arguments

-8	
file	name of text file with one or multiple SIMMAP v1.0 or v1.5 style trees.
text	character string containing the tree. If version=1.5 this argument is ignored. (This format tree can only be read from file in the present version.)
format	format of the trees: either "phylip" or "nexus" - the latter is the default output from SIMMAP. If version=1.5 this argument is ignored.
rev.order	a logical value indicating whether the states and times along each branch is given (from root to tip) in right-to-left order (if TRUE) or in left-to-right order. If version=1.5 this argument is ignored.
version	version of SIMMAP for input tree. If the tree(s) was/were simulated in SIMMAP v1.0 or written to file by link{make.simmap} then version=1.0; if the tree(s) was/were simulated using SIMMAP v1.5 then version=1.5.

Details

This function now accepts trees in both SIMMAP v1.0 and SIMMAP v1.5 format. In addition, it can read a more flexible format than is produced by SIMMAP (for instance, multi-character mapped states and more than 7 mapped states). Uses some modified code from read.nexus from the "ape" package to read the NEXUS block created by SIMMAP. Also creates the attribute "map.order" which indicates whether the stochastic map was read in from left to right or right to left. This attribute is used by default by write.simmap to write the tree in the same order.

Value

A modified object of class "phylo" (or list of class "multiPhylo") with the following additional elements:

maps a list of named vectors containing the times spent in each state on each branch,

in the order in which they occur.

mapped.edge a matrix containing the total time spent in each state along each edge of the tree.

Author(s)

References

Bollback, J. P. (2006) Stochastic character mapping of discrete traits on phylogenies. *BMC Bioinformatics*, **7**, 88.

Paradis, E., J. Claude, and K. Strimmer (2004) APE: Analyses of phylogenetics and evolution in R language. *Bioinformatics*, **20**, 289-290.

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

```
brownie.lite, evol.vcv, read.tree, read.nexus
```

reorder.backbonePhylo Reorders a backbone phylogeny

Description

Function reorders an object of class "backbonePhylo".

Usage

```
## S3 method for class 'backbonePhylo'
reorder(x, order="cladewise", ...)
```

Arguments

```
x an object of class "backbonePhylo".order order. See reorder.phylo for possible orderings.optional arguments.
```

Value

An object of class "backbonePhylo".

Author(s)

Liam Revell < liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

```
phylo.toBackbone
```

reorderSimmap 135

ree	
-----	--

Description

Function returns a reordered modified "phylo" object by using reorder.phylo but then sorting the additional elements \$mapped.edge and \$maps to have the same order as \$edge.

Usage

```
reorderSimmap(tree, order="cladewise", index.only=FALSE, ...)
```

Arguments

```
tree a modified object of class "phylo".

order either "cladewise" or "pruningwise" (see reorder.phylo).

index.only logical value indicating whether only an index should be returned.

other arguments.
```

Value

A modified object of class "phylo".

Author(s)

```
Liam Revell liam.revell@umb.edu>
```

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

```
reorder.phylo, plotSimmap
```

rep.phylo

rep.phylo

Replicate a tree or set of trees

Description

S3 method rep for object of class "phylo" or "multiPhylo". repPhylo is just an alias for rep.phylo and rep.multiPhylo.

Usage

```
## S3 method for class 'phylo'
rep(x, ...)
## S3 method for class 'multiPhylo'
rep(x, ...)
repPhylo(tree, times)
```

Arguments

tree object of class "phylo".
times number of times to replicate tree.
x for S3 method an object of class "phylo" or "multiPhylo".
... other arguments for rep (specifically, times).

Value

An object of class "multiPhylo".

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

```
c.phylo, rep
```

Examples

```
tree<-pbtree(n=100)
trees<-rep(tree,100)</pre>
```

reroot 137

reroot	Re-root a tree along an edge	
101000	ne root a tree atong an eage	

Description

This function re-roots a phylogenetic tree at an arbitrary position along an edge.

Usage

```
reroot(tree, node.number, position=NULL, interactive=FALSE, ...)
```

Arguments

tree a phylogenetic tree in "phylo" format.

node.number number of the node descending from the target branch in tree\$edge - this can also be a tip in which case the node number is the index number of the tip in tree\$tip.label.

position position along the target edge at which to re-root the tree. If not supplied, then the tree will be re-rooted at the node or tip.

interactive logical value indicating whether to use interactive mode (defaults to interactive=FALSE).

arguments to be passed to plotTree for interactive=TRUE only.

Details

This function had an error for rootings along edges descended from the root node for phytools<=0.2-47. This should be fixed in the present version. Now uses paste.tree, root, and splitTree internally. Earlier versions also had an error related to node labels. This should be fixed in phytools>=0.4-47.

Value

A phylogenetic tree in "phylo" format.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Paradis, E., J. Claude, and K. Strimmer (2004) APE: Analyses of phylogenetics and evolution in R language. *Bioinformatics*, **20**, 289-290.

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

```
splitTree, paste.tree, root
```

138 rerootingMethod

reroot	inc	Met	hod
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Get marginal ancestral state reconstructions by re-rooting

Description

This function uses the re-rooting method of Yang et al. (1995) to get the marginal ancestral state estimates for each internal node of the tree using likelihood. This method get the conditional scaled likelihoods for the root node (which is the same as the marginal ancestral state reconstruction for that node) and successively moves the root to each node in the tree. The function can also return the posterior probabilities for the tip nodes of the tree.

Usage

```
rerootingMethod(tree, x, model=c("ER","SYM"), ...)
```

Arguments

tree an object of class "phylo".

x a vector of tip values for species, or a matrix containing the prior probability

that the tip is in each state. If x is a vector, then names(x) should be the species names. If x is a matrix of prior probabilities, then rownames should be species names, column names should be states for the discrete character, and rows of

the matrix should sum to 1.0.

model any revsersible model. model=c("ER", "SYM") recommended.

... optional arguments. Presently the logical argument tips. If tips=TRUE, then

the function will also compute the empirical Bayes posterior probabilities of the

tips following Yang (2006).

Details

This function calls fitMk internally. fitMk uses some code adapted from ace in the ape package.

Value

A list containing the following elements:

loglik the log-likelihood.

Q the fitted transition matrix between states.

marginal.anc the marginal ancestral state reconstructions for each node (and, optionally, each

tip).

Author(s)

rescaleSimmap 139

References

Paradis, E., J. Claude, and K. Strimmer (2004) APE: Analyses of phylogenetics and evolution in R language. *Bioinformatics*, **20**, 289-290.

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

Yang, Z., Kumar, S., Nei, M. (1995) A new method of inference of ancestral nucleotide and amino acid sequences. *Genetics*, **141**, 1641-1650.

See Also

```
ace, fitMk, make.simmap
```

rescaleSimmap

Rescale SIMMAP style tree

Description

This function scales a tree with a mapped discrete character to an arbitrary total height, preserving the relative time spent in each state along each edge.

Usage

```
rescaleSimmap(tree, ...)
```

Arguments

tree a phylogenetic tree in modified "phylo" format with a discrete character map-

ping (e.g., see read.simmap or make.simmap).

... other arguments, such as depth.

Details

Replaces rescaleTree (now rescale.phylo) in the 'geiger' package for SIMMAP style trees.

Value

A phylogenetic tree in modified "phylo" format.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

140 rotateNodes

See Also

```
make.simmap, read.simmap
```

rotateNodes

Rotates a node or set of nodes in a phylogenetic tree

Description

This function is a wrapper for rotate which rotates a set of nodes or all nodes.

Usage

```
rotateNodes(tree, nodes, polytom=c(1,2), ...)
```

Arguments

tree object of class "phylo".

nodes either a single node number to rotate, a vector of node numbers, or the string

"all".

polytom a vector of mode numeric and length two specifying the two clades that should

be exchanged in a polytomy (see rotate).

... optional arguments.

Details

Also addresses the problem that the product of multiple rotations from rotate can be non-compliant with the implicit "phylo" standard because the tip numbers in tree\$edge are not in numerical order 1:n for n tips.

Value

An object of class "phylo" (i.e., a phylogenetic tree).

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

roundBranches 141

Rounds the branch tengths of a tree	roundBranches	Rounds the branch lengths of a tree
-------------------------------------	---------------	-------------------------------------

Description

This function rounds the branch lengths of a tree or trees and reconciles any mappings (as in read.simmap) with the rounded branch lengths.

Usage

```
roundBranches(tree, digits)
```

Arguments

```
tree an object of class "phylo" or "multiPhylo".
digits number of digits for rounding. Passed to round.
```

Value

A tree with branch lengths, or modified "phylo" or "multiPhylo" object with a mapped discrete character.

Author(s)

Liam Revell < liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

```
roundPhylogram Plot a round phylogram
```

Description

Function plots one or multiple round phylograms.

Usage

```
roundPhylogram(tree, fsize=1.0, ftype="reg", lwd=2, mar=NULL, offset=NULL,
direction="rightwards", type="phylogram", xlim=NULL, ylim=NULL, ...)
```

142 roundPhylogram

Arguments

tree	an object of class "phylo" or "multiPhylo" containing one or multiple phylogenies.
fsize	relative font size for tip labels.
ftype	font type - options are "reg", "i" (italics), "b" (bold), or "bi" (bold-italics).
lwd	line width for plotting.
mar	vector containing the margins for the plot to be passed to par. If not specified, the default margins are [0.1,0.1,0.1,0.1].
offset	offset for the tip labels.
direction	plotting direction. Only the option direction="rightwards" is presently supported.
type	plot type. Can be "phylogram" or "cladogram". If type="cladogram" then the branch lengths are not necessary (and, indeed, are not used).
xlim	x-limits for the plot.
ylim	y-limits for the plot.

Details

The underscore character "_" is automatically swapped for a space in tip labels, as in plotSimmap.

Value

Plots a tree.

Author(s)

Liam Revell liam.revell@umb.edu>

optional arguments.

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). $Methods\ Ecol.\ Evol.,\ 3,\ 217-223.$

See Also

plotSimmap, plotTree

rstate 143

rstate

Pick a random state according to a vector of probabilities

Description

Primarily an internal function for make.simmap.

Usage

```
rstate(y)
```

Arguments

У

vector of probabilities. Must have names & should probably add to 1.0.

Details

This function picks a random element in a vector according to the probability assigned that element. It returns the name. Uses rmultinom.

Value

A character or string.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

sampleFrom

Sample from a set of distributions

Description

Function samples from a set of normal distributions with parameters given in xbar and xvar.

Usage

```
sampleFrom(xbar=0, xvar=1, n=1, randn=NULL, type="norm")
```

144 setMap

Arguments

xbar a named vector of means. xvar a named vector of variances.

n a vector containing the sample sizes of each species.

randn a range of sample sizes are to be random.

type "norm" is the only distribution implemented so far.

Value

A vector, with labels.

Author(s)

Liam Revell < liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

setMap

Set color map for objects of class "contMap" or "densityMap"

Description

Function to change the color map (ramp) in an object of class "contMap" or "densityMap".

Usage

```
setMap(x, ...)
```

Arguments

x an object of class "contMap" or "densityMap".

arguments to be passed to colorRampPalette. Also, the argument invert which (if invert=TRUE) will just flip the current color ramp.

Value

An object of class "contMap" or "densityMap".

Author(s)

sim.corrs 145

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

Revell, L. J. 2013. Two new graphical methods for mapping trait evolution on phylogenies. *Methods in Ecology and Evolution*, **4**, 754-759.

See Also

contMap, densityMap

sim.corrs

Multivariate Brownian simulation with multiple correlations and rates

Description

This function conducts BM simulation on a tree with multiple rates and/or multiple evolutionary correlations between characters. If vcv is a single matrix, instead of a list of matrices, sim.corrs will simulate multivariate BM with a single rate matrix.

Usage

```
sim.corrs(tree, vcv, anc=NULL, internal=FALSE)
```

Arguments

	is a phylogenetic tree in '		
tree			

discrete character.

vcv is a square covariance matrix or named list of matrices (one for each mapped

state on the tree).

anc optional vector of values for the root state.

internal logical value indicating whether to return states at internal nodes.

Value

A matrix containing the multivariate tip states for the n species in the tree (and nodes if internal=TRUE).

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J., and D. C. Collar (2009) Phylogenetic analysis of the evolutionary correlation using likelihood. *Evolution*, **63**, 1090-1100.

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

sim.history

See Also

fastBM, make.simmap, read.simmap, sim.history, sim.rates

sim.history

Simulate stochastic character history under some model

Description

This function simulates a stochastic character history for a discretely valued character trait on the tree. The resultant tree is stored as a modified "phylo" object in stochastic character map (e.g., make.simmap) format.

Usage

```
sim.history(tree, Q, anc=NULL, nsim=1, ...)
```

Arguments

tree	a phylogenetic tree as an	object of class "	phylo".

a matrix containing the instantaneous transition rates between states. Note that normally this is the *transpose* of the matrix produced by fitDiscrete in the geiger package or make.simmap in phytools; that is to say the transition rate from i -> j should be given by Q[j,i]. However, if your matrix is properly conformed (i.e., rows *or* columns sum to 0), then sim.history will attempt to transpose your matrix correctly & will return an informative message

(if message=TRUE, see below).

anc an optional value for the state at the root node; if NULL, a random state will be

assigned. anc can be a vector of states, in which one of the states will be chosen randomly for each simulation. anc can be a vector of probabilities with names, in which case a state will be chosen in proportion to the given probabilities.

nsim number of simulations.

... other optional arguments. Currently only message, a logical value indicating

whether or not to turn on informational messages (defaults to message=TRUE).

Value

A modified phylogenetic tree of class "phylo" (or a modified "multiPhylo" object, for nsim > 1) with the following additional elements:

maps a list of named vectors containing the times spent in each state on each branch,

in the order in which they occur.

mapped.edge a matrix containing the total time spent in each state along each edge of the tree.

states a vector containing the tip states.

node.states a matrix containing the states at internal & terminal nodes (according to the

dimensions of edge).

sim.ratebystate 147

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

```
make.simmap, read.simmap, plotSimmap, sim.rates
```

sim.ratebystate

Conduct simulation of state dependent rate variation

Description

This function attempts to simulate two characters under a model in which the rate of evolution for the second (y) depends on the states for the first x. See ratebystate for more details.

Usage

```
sim.ratebystate(tree, sig2x=1, sig2y=1, beta=c(0,1), ...)
```

Arguments

tree	phylogenetic tree.
sig2x	variance of the Brownian process of evolution for x.
sig2y	variance of the Brownian process of evolution for y when x-min(x)==1 (for logarithm=FALSE) or x==0 (for logarithm=TRUE).
beta	intercept and slope of the relationship between the value of x and the Brownian rate in y.
	optional arguments which include method ("by.node" or "by.branch" indicating whether to assume the rate varies as a function of the node state or the mean branch state); plot, a logical value indicating whether or not to plot a phenogram with the branches used for simulation of y after rescaling by the state of x; and logarithm, a logical value indicating whether or not simulate changes in the variance of Brownian evolution for y as an additive logarithm=FALSE or multiplicative function of x. The default is logarithm=FALSE.

Value

This function returns a matrix.

Author(s)

Liam Revell liam.revell@umb.edu>

148 sim.rates

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

fastBM, ratebystate

sim.rates

Brownian simulation with multiple evolutionary rates

Description

This function conducts BM simulation on a tree with multiple rates.

Usage

```
sim.rates(tree, sig2, anc=0, nsim=1, internal=F, plot=F)
```

Arguments

tree	is a stochastic map format phylogenetic tree in modified "phylo" format (e.g., see make.simmap).
sig2	a named vector containing the rates for each state; names should be states in mtree.
anc	optional value for the root state.
nsim	number of simulations.
internal	logical value indicating whether to return states at internal nodes.
plot	logical value indicating whether or not to visual the rate heterogeneity (default value is FALSE.

Value

A vector (for nsim=1) or matrix containing the tip states for the n species in the tree.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

fastBM, make.simmap, read.simmap, sim.history

skewers 149

skewers

Matrix comparison using the method of random skewers

Description

This function performs the random skewers matrix comparison method of Cheverud (1996; also see Cheverud & Marroig 2007 for more details). In addition, it includes a more robust hypothesis test in which random covariance matrices are simulated under a variety of models, and then the mean correlation between response vectors to random skewers are computed.

Usage

```
skewers(X, Y, nsim=100, method=NULL)
```

Arguments

X covariance matrix.
Y covariance matrix.

nsim number of random vectors.

method method to generate a null d

method to generate a null distribution of the random skewers correlation between matrices. If method=NULL then the correlation will be compared to the correlation between random vectors; however this test has type I error substantially above the nominal level for ostensibly random matrices. Other values of method will be passed as covMethod to genPositiveDefMat for a more robust hypothesis test (see below). Recommended values include "unifcorrmat".

Value

A list with the following components:

r mean random skewers correlation.

p p-value from simulation.

Author(s)

Liam Revell < liam.revell@umb.edu>

References

Cheverud, J. M. (1996) Quantitative genetic analysis of cranial morphology in the cotton-top (Saguinus oedipus) and saddle-back (S. fuscicollis) tamarins. *J. Evol. Biol.*, **9**, 5–42.

Cheverud, J. M. & Marroig, G. (2007) Comparing covariance matrices: Random skewers method compared to the common principal components model. *Genetics & Molecular Biology*, **30**, 461–469.

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

splitplotTree

splitEdgeColor

Split edge colors when descendant edges have different mapped states

Description

This function splits the vertical line colors to match the daughter edges when the daughters have different states. Mostly to be used with trees generated using paintBranches or paintSubTree. Also used internally by plotSimmap.

Usage

```
splitEdgeColor(tree,colors,lwd=2)
```

Arguments

tree object of class "simmap".

colors named vector of colors to be used for plotting.

1wd width of the plotted lines.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

plotSimmap

splitplotTree

Plots a phylogeny in two columns

Description

Function plots a tree in two columns or windows.

Usage

```
splitplotTree(tree, fsize=1.0, ftype="reg", lwd=2, split=NULL, new.window=FALSE)
plotTree.splits(tree, splits=NULL, file=NULL, fn=NULL, ...)
```

splitTree 151

Arguments

tree	an object of class "phylo".
fsize	relative font size for tip labels.
ftype	font type - options are "reg", "i" (italics), "b" (bold), or "bi" (bold-italics).
lwd	line width for plotting.
split	relative vertical position for splitting the tree (between 0 & 1).
new.window	whether or not to plot the split tree in a new window. If FALSE then the tree will be plotted in two columns within the same plotting window.
splits	for plotTree.splits relative positions (from 0 to 1) to split the tree across pages or devices.
file	filename if saving to a PDF file is desired. Otherwise will plot to the default plotting device.
fn	function to be executed on each plotted page. For instance, might be: function() cladelabels() if clade labels are desired.

Value

. . .

Plots a tree.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

other arguments to be passed to plotTree.

See Also

plotTree, plotSimmap

lit tree at a point	plitTree	Split tree at a point
1		1

Description

Primarily an internal function for posterior.evolrate, this function splits the tree at a given point, and returns the two subtrees as an object of class "multiPhylo".

Usage

```
splitTree(tree, split)
```

starTree

Arguments

tree phylogenetic tree.

split split encoded as a list with two elements: node: the node number tipward of the

split; and bp: the position along the branch to break the tree, measured from the

rootward end of the edge.

Details

Probably do not use this unless you can figure out what you are doing.

Value

Two trees in a list.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

starTree

Create star phylogeny

Description

This function creates a star phylogeny.

Usage

```
starTree(species, branch.lengths=NULL)
```

Arguments

species a list of species.

branch.lengths an optional list of branch lengths in the same order as species.

Details

Creates a star phylogeny with (optionally) user specified branch lengths.

Value

An object of class "phylo".

strahlerNumber 153

Author(s)

Liam Revell < liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

stree

strahlerNumber

Computes Strahler number for trees and nodes

Description

The function strahlerNumber computes the Strahler number of all nodes and tips in the tree. For more information about Strahler numbers see http://en.wikipedia.org/wiki/Strahler_number. The function extract.strahlerNumber extracts all of the most inclusive clades of Strahler number i.

Usage

```
strahlerNumber(tree, plot=TRUE)
extract.strahlerNumber(tree, i, plot=TRUE)
```

Arguments

tree an object of class "phylo".

i order of Strahler number to extract for extract.strahlerNumber.

plot logical value indicating whether to plot the tree with Strahler numbers for node

labels.

Value

Either a vector with the Strahler number for each tip and internal node; or (for extract.strahlerNumber the set of (most inclusive) subtrees with Strahler number i as an object of class "multiPhylo".

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

154 threshBayes

threshBayes	Threshold model using	g Bayesian MCMC
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Description

This function uses Bayesian MCMC to fit the quantitative genetics threshold model (Felsenstein 2012) to data for two discrete characters or one discrete and one continuous character.

Usage

```
threshBayes(tree, X, types=NULL, ngen=1000, control=list())
```

Arguments

tree an object of class "phylo".

X a numeric matrix containing values for a numerically coded discrete character

and a continuous character; or two discrete characters. The row names of X

should be species names.

types a vector of length ncol(X) containing the data types for each column of X, for

instance c("discrete", "continuous").

ngen a integer indicating the number of generations for the MCMC.

control a list of control parameters for the MCMC. Control parameters include: sample,

the sampling interval for the MCMC; propvar, a vector containing (in this order) proposal variances for the two rates (if the type is "discrete" this will be ignored), the two ancestral states, and the correlation; propliab, a single proposal variance for the liabilities; pr.mean, a vector for the mean of the prior probability distributions for each parameter, in the same order as propvar; pr.liab, currently ignored; pr.var, a vector with variances for the prior densities for each parameter, in the same order as pr.mean - note that for the rates we use an exponential distribution so the first two means are currently ignored; and

pr.vliab currently ignored.

Value

This function returns a list with two elements: par a matrix containing the posterior sample for the model parameters (evolutionary rates, ancestral states, and correlation); liab a matrix containing the posterior sample of the liabilities. For continuous characters, the liabilities are treated as known and so the posterior samples are just the observed values.

Author(s)

Liam Revell liam.revell@umb.edu>

threshDIC 155

References

Felsenstein, J. (2012) A comparative method for both discrete and continuous characters using the threshold model. *American Naturalist*, **179**, 145-156.

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

Revell, L. J. (2014) Ancestral character estimation under the threshold model from quantitative genetics. *Evolution*, bold68, 743-759.

See Also

```
anc.Bayes, bmPlot, evol.rate.mcmc
```

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Deviance Information Criterion from the threshold model

Description

This function computes the Deviance Information Criterion from the MCMC object returned by ancThresh.

Usage

```
threshDIC(tree, x, mcmc, burnin=NULL, sequence=NULL, method="pD")
```

Arguments

tree	phylogenetic tree.
X	a named vector containing discrete character states; or a matrix containing the tip species, in rows, and probabilities of being in each state, in columns.
mcmc	list object returned by ancThresh.
burnin	number of generations (not samples) to exclude as burn in; if not supplied then 20 percent of generations are excluded.
sequence	assumed ordering of the discrete character state. If not supplied and x is a vector then numerical-alphabetical order is assumed; if not supplied and x is a matrix, then the column order of x is used.
method	method for computing the effective number of parameters (options are "pD" and "pV").

Value

A vector containing the mean deviance and deviance for the parameter means, the effective number of parameters, and the DIC.

Author(s)

Liam Revell liam.revell@umb.edu>

156 threshState

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

Revell, L. J. (2014) Ancestral character estimation under the threshold model from quantitative genetics. *Evolution*, bold68, 743-759.

See Also

ancThresh

threshState

Computes value for a threshold character from a liability and thresholds

Description

Primarily to be used internally by ancThresh; can also be used to simulate threshold traits.

Usage

```
threshState(x, thresholds)
```

Arguments

x liability.

thresholds a named vector containing the thresholds.

Value

A discrete character value.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

Revell, L. J. (2014) Ancestral character estimation under the threshold model from quantitative genetics. *Evolution*, bold68, 743-759.

See Also

ancThresh, threshDIC

to.matrix 157

to.matrix

Convert a character vector to a binary matrix

Description

This function takes a vector of characters and computes a binary matrix. Primarily to be used internally by make.simmap and rerootingMethod.

Usage

```
to.matrix(x, seq)
```

Arguments

x a vector of characters.

seq the sequence for the columns in the output matrix.

Value

A binary matrix of dimensions length(x) by length(seq).

Author(s)

Liam Revell < liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

treeSlice

Slices the tree at a particular point and returns all subtrees

Description

This function slices a tree at a particular height above the root and returns all subtrees or all non-trivial subtrees (i.e., subtrees with more than 1 taxon). Uses extract.clade in the "ape" package.

Usage

```
treeSlice(tree, slice, trivial=FALSE, prompt=FALSE, ...)
```

158 untangle

Arguments

tree	is a phylogenetic tree in "phylo" format.
slice	a real number indicating the height above the root at which to slice the tree.
trivial	a logical value indicating whether or not to return subtrees with a number of tips less than two (default is FALSE).
prompt	logical value indicating whether or not the height of the slice should be given interactively.
	for prompt=TRUE, other arguments to be passed to plotTree.

Value

An object of class "phylo" or "multiPhylo".

Author(s)

Liam Revell < liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

```
extract.clade
```

untangle

Attempts to untangle crossing branches for plotting

Description

This function attempts to untangle the branches of a tree that are tangled in plotting with plot.phylo, plotTree, or plotSimmap. Note that method="read.tree" does not presently work for SIMMAP style trees

Usage

```
untangle(tree, method=c("reorder","read.tree"))
```

Arguments

tree as an object of class "phylo". Can be a SIMMAP style tree (e.g., read.simmap.

method method to use to attempt to untangle branches. method="reorder" uses two
calls of reorder.phylo or reorderSimmap; method="read.tree" writes the
tree to a text string and then reads it back into memory using read.tree.

vcvPhylo 159

Value

A tree with branch lengths, or modified "phylo" object with a mapped discrete character.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

vcvPhylo

Calculates cophenetic (i.e., phylogenetic VCV) matrix

Description

This function returns a so-called *phylogenetic variance covariance matrix* (e.g., see vcv.phylo), but (optionally) including ancestral nodes and under different evolutionary models.

Usage

```
vcvPhylo(tree, anc.nodes=TRUE, ...)
```

Arguments

tree object of class "phylo".

anc.nodes logical value indicating whether or not to include ancestral nodes.

optional arguments including internal (synonym of anc.nodes) and model

(can be "BM", "OU", or "lambda".

Value

A matrix.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

160 write.simmap

write.simmap	Write a stochastic character mapped tree to file	
--------------	--	--

Description

This function writes stochastic character mapped trees to file using the Newick style format of SIMMAP v1.0 (Bollback 2006). Note, can only write one tree at a time to file (hence the append option).

Usage

```
write.simmap(tree, file=NULL, append=FALSE, map.order=NULL, quiet=FALSE)
```

Arguments

tree a phylogenetic tree as a modified object of class "phylo". See make.simmap

and read.simmap.

file an optional filename.

append a logical value indicating whether to append to file.

map.order a optional value specifying whether to write the map in left-to-right or right-to-

left order. Acceptable values are "left-to-right" or "right-to-left" or some abbreviation of either. If not provided, write.simmap will use attr(tree, "map.order")

if available.

quiet logical value indicating whether or not to print a warning message when map.order

is neither specified by a function argument or as an attribute of tree.

Value

A file or string (if file=NULL).

Author(s)

Liam Revell liam.revell@umb.edu>

References

Bollback, J. P. (2006) Stochastic character mapping of discrete traits on phylogenies. *BMC Bioinformatics*, **7**, 88.

Huelsenbeck, J. P., R. Neilsen, and J. P. Bollback (2003) Stochastic mapping of morphological characters. *Systematic Biology*, **52**, 131-138.

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

make.simmap, read.simmap, plotSimmap

writeAncestors 161

writeAncestors	Write a tree to file with ancestral states and (optionally) CIs at nodes

Description

This function writes a tree to file with ancestral character states and (optionally) 95-percent confidence intervals stored as node value..

Usage

```
writeAncestors(tree, Anc=NULL, file="", digits=6, format=c("phylip", "nexus"), ...)
```

Arguments

tree	a phylogenetic tree or set of trees as an object of class "phylo" or "multiPhylo".
Anc	a vector of ancestral states, a list containing the ancestral states and 95-percent confidence intervals (as from fastAnc or ace, or a list of such results.
file	an optional string with the filename for output.
digits	an integer indicating the number of digits to print for branch lengths and ancestral character values.
format	a string indicating whether to output the result in simple Newick (i.e., "phylip") or Nexus format.
	additional arguments including x: a vector of character values, in which case ancestral states are estimated internally using fastAnc; and CI: a logical value indicating whether or not to estimate 95-percent confidence intervals.

Value

A file, string, or vector of strings.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

```
ace, fastAnc, write.tree
```

162 writeNexus

writeNexus

Write a tree to file in Nexus format

Description

This function writes one or multiple phylogenetic trees to file in NEXUS format. Redundant with ape::write.nexus.

Usage

```
writeNexus(tree, file="")
```

Arguments

tree object of class "phylo" or "multiPhylo".

file file name for output.

Value

Trees written to file.

Author(s)

Liam Revell liam.revell@umb.edu>

References

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.*, **3**, 217-223.

See Also

```
write.simmap, write.nexus
```

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