Package 'pez'

June 17, 2021

Type Package

Title Phylogenetics for the Environmental Sciences

Version 1.2-3

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Description Eco-phylogenetic and community phylogenetic analyses.

Keeps community ecological and phylogenetic data matched up and comparable using 'comparative.comm' objects. Wrappers for common community phylogenetic indices ('pez.shape', 'pez.evenness', 'pez.dispersion', and 'pez.dissimilarity' metrics). Implementation of Cavender-Bares (2004) correlation of phylogenetic and ecological matrices ('fingerprint.regression'). Phylogenetic Generalised Linear Mixed Models (PGLMMs; 'pglmm') following Ives & Helmus (2011) and Rafferty & Ives (2013). Simulation of null assemblages, traits, and phylogenies ('scape', 'sim.meta.comm').

License GPL-3

VignetteBuilder knitr

Suggests knitr (>= 1.6), lme4 (>= 1.1-7), formatR (>= 1.7)

Imports caper (>= 0.5-2), picante (>= 1.6-2), quantreg (>= 5.05), mvtnorm (>= 1.0-0), vegan (>= 2.0-10), ade4 (>= 1.6-2), FD (>= 1.0-12), Matrix (>= 1.1-4), methods (>= 3.1.0), animation (>= 2.4-0), phytools (>= 0.6-60)

Depends ape (>= 3.1-4)

RoxygenNote 7.1.1

NeedsCompilation no

Repository CRAN

Date/Publication 2021-06-17 12:00:06 UTC

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Description

As described in the vignette, we recommend using these wrappers to manipulate species and site data, as it guarantees that everything will be kept consistent across all parts of the comparative.comm object. With them, you can drop species, sites, and work directly with each part of your data. You can also manipulate your comparative.comm object's phy, data, env, and comm slots directly if you wish, but altering the object directly yourself runs the risk of things getting unsynchronised.

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Usage

```
## S3 method for class 'comparative.comm'
x[sites, spp, warn = FALSE]
trait.names(object)
env.names(object)
species(x)
species(x) <- value</pre>
sites(x)
sites(x) <- value</pre>
traits(x) <- value</pre>
traits(x)
env(x) \leftarrow value
env(x)
comm(x) \leftarrow value
comm(x)
tree(x)
phy(x)
tree(x) <- value</pre>
phy(x) \leftarrow value
assemblage.phylogenies(data)
## S3 method for class 'comparative.comm'
as.data.frame(
  х,
  row.names = NULL,
  optional = FALSE,
  abundance.weighted = FALSE,
)
## S3 method for class 'comparative.comm'
```

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```
within(data, expr, ...)
```

Arguments

	X	comparative.comm object
	sites	numbers of sites to be kept or dropped from x; must be given as numbers. For example, $x[1:5,]$, or $x[-1:-5,]$, but not $x[c("site a","site b"),]$.
	spp	numbers of species to be kept or dropped from x; must be given as numbers. For example, $x[,1:5]$, or $x[,-1:-5]$, but not $x[c("sp a","sp b"),]$.
	warn	whether to warn if species/sites are dropped when creating object (default: TRUE)
	object	A comparative.comm object
	value	when altering a comparative.comm object's internal structure, the thing that you're inserting into it!
	data	A comparative.comm object
	row.names	ignored
	optional	ignored presence-absence dataset (default: FALSE)
abundance.weighted		nted
		whether to create to create a
		ignored
	expr	expression to be evaluated within the scope of data

Value

Names of the traits or environmental variables

Note

As described in comparative.comm, each comparative.comm object contains a phylogeny (\$phy) and a site-by-species community matrix (as used in vegan). Optionally, it may contain a data.frame of trait data (each row a species, each column a trait) *called data* for compatibility with comparative.data.

See Also

comparative.comm plot.comaparative.comm

Examples

```
data(laja)
data <- comparative.comm(invert.tree, river.sites, invert.traits, river.env)
#Subset on species, then sites
data <- data[1:5,]
data <- data[,1:5]
#Site and species can be manipulated
species(data)
sites(data)[1:3] <- c("lovely", "invert", "sites")
#Other data can be viewed
trait.names(data)</pre>
```

comparative.comm 5

```
env.names(data)
#Get assemblage phylogenies of all sites
assemblage.phylogenies(data)
#Add some trait/env data in
traits(data)$new.trait <- sample(letters, nrow(comm(data)), replace=TRUE)
env(data)$new.env <- sample(letters, ncol(comm(data)), replace=TRUE)
#Manipulate/check phylogeny and community matrix
phy(data) #...tree(data) works too...
comm(data)[1,3] <- 3
comm(data) <- comm(data)[-3,]</pre>
```

comparative.comm

Creates a community comparative ecology object, the basis of all functions in pez

Description

Basic checking of whether the input data match up is performed; you need only supply comm and phy, nothing else is mandatory. You can manipulate the internals of comparative.comm, or use the wrappers inside pez to keep everything in order. Examples of these features are given below; they are described in detailed at cc.manip.

Usage

```
comparative.comm(
  phy,
  comm,
  traits = NULL,
  env = NULL,
  warn = TRUE,
  force.root = -1
)

## S3 method for class 'comparative.comm'
print(x, ...)
```

Arguments

phy	phylogeny (in phylo format) of species
comm	community matrix (as used in vegan) with species as columns and rows as communities. Must contain rownames and colnames; NAs are not checked for but probably unwise.
traits	data.frame of species traits, with rownames matching comm. Saved in the data slot of the resulting comparative.comm object for compatibility with comparative.data.
env	data.frame of environmental data with rownames matching comm
warn	whether to warn if species/sites are dropped when creating object (default: TRUE)

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```
force.root if phy is unrooted, a root.edge of force.root will be added (default: -1, which means this will never happen). Rarely needed, rarely advisable.

x comparative.comm object to be printed
... ignored
```

Value

comparative.comm object

Note

comparative.comm is compatible with comparative.data; this means that the slot for species' trait data is called data. I appreciate this is somewhat unwieldy, but hopefully you agree it is helpful in the long-term.

Author(s)

Will Pearse

See Also

```
plot.comparative.comm cc.manip link[caper:comparative.data]{comparative.data}
```

Examples

```
data(laja)
data <- comparative.comm(invert.tree, river.sites, invert.traits, river.env)
#Subset on species, then sites
data <- data[1:5,]
data <- data[,1:5]
#Site and species can be manipulated
species(data)
sites(data)[1:3] <- c("lovely", "invert", "sites")
#Other data can be viewed
trait.names(data)
env.names(data)
#Get assemblage phylogenies of all sites
assemblage.phylogenies(data)
#Do some manual manipulation of your objects (NOTE: $data for traits)
data$data$new.trait <- sample(letters, nrow(data$comm), replace=TRUE)</pre>
```

ConDivSim

Null models for functional-phylogenetic diversity

Description

Simulate expectations (under a null model) of mean pairwise distance for a set of communities with different species richness.

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Usage

```
ConDivSim(object, type = "traits", n.sim = 100, plot = TRUE, disp99 = FALSE)
```

Arguments

object	a comparative.comm object, with presence-absence community data.	
type	character string giving the type of distance matrix on which the mean pairw distance is based. Either "trait" or "phy" to a phylogenetic or trait-based distant matrix, or an actual matrix to use (e.g., one from funct.phylo.dist)	
n.sim	The number of permutations of the presence vector used to make the estimations.	
plot	TRUE or FALSE to make the plot of the expected average mean pairwise distance, and the 5-95% confidence interval.	
disp99	Display the 99% interval?	

Details

If plot == TRUE, then a surface is drawn giving the null distribution. Lighter shades of gray give larger intervals with categories: 0.005-0.995 = 99%, 0.025-0.975 = 95%, 0.05-0.95 = 90%, 0.25-0.75 = 50%.

Value

matrix with quantiles of mean pairwise distances for all quantiles of of mean pairwise distance, with one row for the range of species richnesses in the data (see column SpRich).

Note

No serious checking of user-provided matrices is performed; this is both useful and dangerous!

Author(s)

Steve Walker, wrappers by Will Pearse

See Also

```
sim.phy scape
```

Examples

```
data(laja)
data <- comparative.comm(invert.tree, river.sites, invert.traits)
#Must have all species present in at least one community!
#...and must be presence-absence data
data <- data[,colSums(data$comm) > 0]
data$comm[data$comm>1] <- 1
sims <- ConDivSim(data)
#...without traits...
sims.phy <- ConDivSim(data, type="phy")</pre>
```

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dist.xxx

Make co-existence matrices based on phylogeny (and/or) traits, and community or environemntal overlap

Description

Make co-existence matrices based on phylogeny (and/or) traits, and community or environemntal overlap

Usage

```
comm.dist(x)
## S3 method for class 'matrix'
comm.dist(x)
## S3 method for class 'comparative.comm'
comm.dist(x)
traits.dist(x, dist.func = dist.func.default, ...)
## S3 method for class 'comparative.comm'
traits.dist(x, dist.func = dist.func.default, alltogether = TRUE, ...)
## Default S3 method:
traits.dist(x, dist.func = dist.func.default, ...)
## S3 method for class 'data.frame'
traits.dist(x, dist.func = dist.func.default, ...)
dist.func.default(x)
phylo.dist(x, ...)
## S3 method for class 'phylo'
phylo.dist(x, ...)
## S3 method for class 'comparative.comm'
phylo.dist(x, ...)
funct.phylo.dist(x, phyloWeight, p = 2, ...)
pianka.dist(x, ...)
## S3 method for class 'matrix'
pianka.dist(x, env = NULL, ...)
```

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```
## S3 method for class 'comparative.comm'
pianka.dist(x, alltogether = TRUE, ...)
```

Arguments

x	an object	
dist.func	a function for computing distances. The default, dist.func.default, returns a Euclidean distance of the scaled and centred data.	
	not used	
alltogether	should one multivariate distance matrix be computed for all traits at once (DE-FAULT; alltogether = TRUE) or for each trait at a time (alltogether = FALSE)?	
phyloWeight	phylogenetic weighting parameter (referred to as a in Cadotte et al. (2013)	
p	exponent giving the exponent to use for combining functional and phylogenetic distances (the default, p = 2, gives a Euclidean combination).	
env	environmental variable to be used to calculate the distance matrix	

Details

comm. dist returns the 1 - co-existence of species. Look at how this is calcualted; it incorporates abundances, and if you don't want it to do so simply call it on a presence/absensence (1/0) matrix.

traits.dist returns the functional trait distance of species

phylo.dist returns the phylogenetic (cophenetic) distance of species

funct.phylo.dist returns the combined phylogenetic and trait distances of species, based on the traitgram approach of Cadotte et al. (2013).

Make functional phylogenetic distance matrix

pianka. dist returns the environemntal tolerances distance matrices of species. Based on Pianka's distance (i.e., niche overlap based on environmental variables at co-occuring sites), as defined in Cavender-Bares et al. (2004) - likely not the original reference!

References

Cadotte M.A., Albert C.H., & Walker S.C. The ecology of differences: assessing community assembly with trait and evolutionary distances. Ecology Letters 16(10): 1234–1244.

Cavender-Bares J., Ackerly D.D., Baum D.A. & Bazzaz F.A. (2004) Phylogenetic overdispersion in Floridian oak communities. The Americant Naturalist 163(6): 823–843.

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drop_tip

Trim a phylogeny

Description

This is a weak wrapper around ape's drop.tip. Importantly, if asked to drop no species from a phylogeny, it will just return the phylogeny (not an empty phylogeny, as drop.tip) will.

Usage

```
drop_tip(tree, spp)
```

Arguments

tree An phylo object

spp A vector of species (one, many, or none) to be removed from tree

Value

phylo object

See Also

drop.tip extract.clade

eco.scape

eco.space scape simulation with a macro-ecological focus

Description

eco. scape is a modified version of the Helmus et al. method implemented in scape. It produces phylogenetically structured communities. It allows phylogenetic signals in niche optima, but unlike scape, does not include the ability to specify niche optima signal type (attraction/repulsion) or phylogenetic signal in range size. Instead, the focus is on having more control over the macroecological characteristics of the resulting landscapes. In particular, eco.scape produces landscapes with fixed mean range sizes, reasonable range size and abundance distributions, and control over whether species present on a tree must be present in the landscape.

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Usage

```
eco.scape(
   tree,
   scape.size = 10,
   g.center = 1,
   wd.all = 0.2 * (scape.size + 1)^2,
   signal.center = TRUE,
   center.scale = 1,
   site.stoch.scale = 0,
   sd.center = 1,
   sd.range = 1,
   K = 100,
   extinction = FALSE,
   rho = NULL
)
```

Arguments

tree	phylo object; must have branch lengths and be ultrametric	
scape.size	edge dimension of square landscape	
g.center	strength of phylogenetic signal in species range centers. See corBlomberg, 1=brownian,<1=rates of evol accelerate, >1=rates decelerate.	
wd.all	niche width, larger values simulate broader range sizes	
signal.center	simulate with phylosignal in range centers	
center.scale	adjust strength of phylogenetic attraction in range centers independent of g.center	
site.stoch.scale		
	adjust strength of random variation in species richness across sites	
sd.center	sd in rnormrnorm for the range centers, increase to get more variation in center values across species	
sd.range	sd in rnorm for the range sizes, increase to get more variation in range sizes across gradients	
К	carrying capacity of a site in terms of maximum individuals that can be present. Currently a constant value. Used to scale the presence-absence matrix to include abundances.	
extinction	TRUE/FALSE can species on the tree go extinct on the landscape? If the number of species present on the landscape should equal the number of tips on the tree, choose FALSE. See Details.	
rho	Grafen branch adjustment of phylogenetic tree see corGrafen	

Details

Simulates a landscape with species (i.e., tree tips) distributions dependent on a supplied phylogenetic tree. The amount and type of structure is determened by the signal parameter g.center. Parameters are based on an Ornstein-Uhlenbeck model of evolution with stabilizing selection. Values of g=1 indicate no stabilizing selection and correspond to the Brownian motion model of evolution;

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01 corresponds to disruptive selection where phylogenetic signal for the supplied tree is amplified. See corBlomberg. Communities are simulated along two gradients where the positions along those gradients, g.center, can exhibit phylogenetic signal.

The function returns a landscape where the average range size is equivalent to the wd.all parameter - in the scape function, this parameter is not necessarily returned in the resulting landscape. To do this, the probability of presence (th) that returns the wd.all parameter is solved for. If there is no solution that can produce the wd.all given, the error "Error in uniroot(f, lower = 0, upper = $\max(X)$, tol = 10^{-200}): f() values at end points not of opposite sign" will occur. This seems to mostly arise for extreme or unlikely parameter values (small species pools, low carrying capacities). Try adjusting parameter values first.

The extinction parameter specifies whether all of the species on the tree should be present in the final landscape. Some species will have probabilities of presence less than those required for presence. If extinctions is TRUE, these species will not be present. If FALSE, these species will be present in 1 site, that in which they have the highest probability of presence.

Value

cc comparative.comm object with presence/absence results of simulations. The

site names are the row.columns of the cells in the original grid cells that made up the data, and these co-ordinates are also given in the env slot of the object

along with the environmental gradient information.

Y presence/absence matrix

Yab abundance matrix

index spatial coordinates for X and Y (stacked columns)

X. joint full probabilities of species at sites, used to construct Y

X1 probabilities of species along gradient 1 X2 probabilities of species along gradient 2

gradient1, gradient2

environmental gradient values

nichewd average niche width of the assemblage

K carrying capacity of each cell

environ matrix depicting environmental values over the 2D landscape

sppXs full probabilities of each species as an array arranged in a scape.size X scape.size

matr ix

V.phylo initial phylogenetic covariance matrix from tree, output of vcv.phylo(tree, corr=T)
V.phylo.rho phylogenetic covariance matrix from tree scaled by Grafen if rho is provided,

other wise just an output of vcv.phylo(tree, corr=T)

V. center scaled (by g.center) phylo covariance matrix used in the simulations

bspp1 species optima for gradient 1 bspp2 pecies optima for gradient 2

Author(s)

Matt Helmus, Caroline Tucker, cosmetic edits by Will Pearse

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See Also

```
scape sim.phy sim.meta
```

Examples

```
# Simulations
tree <- rcoal(64)
scape1 <- eco.scape(tree, scape.size=25, g.center=1,</pre>
    signal.center=FALSE, K=100, extinction=TRUE)
scape2 <- eco.scape(tree, scape.size=16, g.center=0.2,</pre>
    signal.center=TRUE, K=100, extinction=FALSE)
scape3 <- eco.scape(tree, scape.size=16, g.center=20,</pre>
    signal.center=TRUE, K=100, extinction=TRUE)
# Plotting distributions and landscape patterns
original_landscape <- scape1</pre>
abundmax <- original_landscape$K
PA_mat <- as.matrix(original_landscape$Y)</pre>
abund_mat <- original_landscape$Yab</pre>
site.size <- nrow(PA_mat)</pre>
species <- ncol(PA_mat)</pre>
mx <- original_landscape$gradient</pre>
env <- original_landscape$environ$env.gradient</pre>
par(mfrow=c(2,2), oma=c(0,0,2,0))
heatcol <- (colorRampPalette(c("yellow","red")))</pre>
image(matrix(env,sqrt(site.size),sqrt(site.size),byrow=TRUE),
    col=heatcol(max(env)),xaxt="n",yaxt="n",main="Env gradient")
image(matrix(rowSums(PA_mat),sqrt(site.size),sqrt(site.size),byrow=TRUE),
    col=heatcol(16),xaxt="n",yaxt="n",main="Species Richness")
hist(colSums(PA_mat),ylab="Number of species",xlab="Number of sites",
    main="Species Area Relationship",col="lightgrey")
hist(colSums(abund_mat),ylab="Number of species",xlab="Number of individuals",
    main="Species Abundance Relationship",col="lightgrey")
mtext("Env random, phy.signal=0.2, 32 species", outer=TRUE, side=3, cex=1.25)
```

eco.xxx.regression

eco.xxx.regression

Description

Regression species co-existence against environmental tolerance, trait similarity, or phylogenetic relatedness.

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Usage

```
eco.env.regression(
  data,
  randomisation = c("taxa.labels", "richness", "frequency", "sample.pool",
    "phylogeny.pool", "independentswap", "trialswap"),
  permute = 0,
  method = c("quantile", "lm", "mantel"),
  altogether = TRUE,
  indep.swap = 1000,
  abundance = TRUE,
)
eco.phy.regression(
  data,
  randomisation = c("taxa.labels", "richness", "frequency", "sample.pool",
    "phylogeny.pool", "independentswap", "trialswap"),
  permute = 0,
 method = c("quantile", "lm", "mantel"),
  indep.swap = 1000,
  abundance = TRUE,
)
eco.trait.regression(
  data,
  randomisation = c("taxa.labels", "richness", "frequency", "sample.pool",
    "phylogeny.pool", "independentswap", "trialswap"),
  permute = 0,
  method = c("quantile", "lm", "mantel"),
  altogether = TRUE,
  indep.swap = 1000,
  abundance = TRUE,
)
## S3 method for class 'eco.xxx.regression'
summary(object, ...)
## S3 method for class 'eco.xxx.regression'
print(x, ...)
## S3 method for class 'eco.xxx.regression'
plot(x, ...)
```

Arguments

data comparative.comm for analysis

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randomisation null distribution with which to compare your community data, one of: taxa.labels

(DEFAULT), richness, frequency, sample.pool, phylogeny.pool, independentswap,

trialswap (as implemented in picante)

permute the number of null permutations to perform (DEFAULT 0)

method how to compare distance matrices (only the lower triangle;), one of: 1m (linear

regression), quantile (DEFAULT; quantreg::rq), mantel (mantel)

altogether use distance matrix based on all traits (default TRUE), or perform separate re-

gressions for each trait (returns a list, see details)

indep.swap number of independent swap iterations to perform (if specified in randomisation;

DEFAULT 1000)

abundance whether to incorporate species' abundances (default: TRUE)
... additional parameters to pass on to model fitting functions

object eco.xxx.regression object x eco.xxx.regression object

Details

These methods are similar to those performed in Cavender-Bares et al. (2004). Each function regresses the species co-existence matrix of data (calculated using comm.dist) against either species' trait dissimilarity (eco.trait.regression), species' phylogenetic distance (eco.phy.regression), or species' shared environmental tolerances as measured by Pianka's distance (eco.env.regression).

If altogether is set to FALSE, each trait or environemntal variables in your data will have a separate eco.trait.regression or eco.env.regression applied to it. The functions will return a list of individual regressions; you can either examine/plot them as a group (see examples below), or extract an individual regression and work with that. These lists are of class eco.xxx.regression.list; a bit messy, but it does work!...

Note

Like fingerprint.regression, this is a data-hungry method. Warnings will be generated if any of the methods cannot be fitted properly (the examples below give toy examples of this). In such cases the summary and plot methods of these functions may generate errors; perhaps use traceback to examine where these are coming from, and consider whether you want to be working with the data generating these errors. I am loathe to hide these errors or gloss over them, because they represent the reality of your data!

WDP loves quantile regressions, and advises that you check different quantiles using the tau options.

Author(s)

Will Pearse, Jeannine Cavender-Bares

References

Cavender-Bares J., Ackerly D.D., Baum D.A. & Bazzaz F.A. (2004) Phylogenetic overdispersion in Floridian oak communities. The Americant Naturalist 163(6): 823–843.

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Kembel, S.W., Cowan, P.D., Helmus, M.R., Cornwell, W.K., Morlon, H., Ackerly, D.D., Blomberg, S.P. & Webb, C.O. Picante: R tools for integrating phylogenies and ecology. Bioinformatics 26(11): 1463–1464.

Pagel M. Inferring the historical patterns of biological evolution. Nature 401(6756): 877–884.

See Also

```
fingerprint.regression phy.signal
```

Examples

```
data(laja)
#We wouldn't recommend only using ten permutations - this is just for speed!
data <- comparative.comm(invert.tree, river.sites, invert.traits, river.env)
eco.trait.regression(data, permute=10)
#Specify additional options
eco.trait.regression(data, tau=c(0.25,0.5,0.75), permute=10)
plot(eco.trait.regression(data, permute=10, method="lm"))
plot(eco.trait.regression(data, permute=10, method="lm", altogether=FALSE))</pre>
```

```
eco.xxx.regression.list
```

List of eco.xxx.regressions

Description

List of eco.xxx.regressions

Usage

```
## S3 method for class 'eco.xxx.regression.list'
summary(object, ...)
## S3 method for class 'eco.xxx.regression.list'
print(x, ...)
## S3 method for class 'eco.xxx.regression.list'
plot(x, ...)
```

Arguments

```
object eco.xxx.regression.list object
... additional arguments to plotting functions
x eco.xxx.regression.list object
```

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fibre.plot

fibre.plot (fibrously) plots a phylogeny

Description

fibre.plot (fibrously) plots a phylogeny

Usage

```
fibre.plot(
   tree,
   gif,
   focal,
   frames = 60,
   colours = colorRampPalette(c("blue", "black", "red")),
   f.colours = colorRampPalette(c("darkgreen", "lightgreen")),
   pca = NULL,
   clade.mat = NULL,
   delay = 0.2,
   side.tree = TRUE,
   width = 600,
   height = 600
)
```

Arguments

tree	a phylogeny (of class phylo) you wish to plot
gif	name of GIF you would like to create. This should *not* including a folder name (this is due to the use of <code>saveGIF</code>); "plot.gif" would be fine, but "work/plot.gif" would not
focal	species numbers or clade numbers to plot differently (see examples). Note that specifying a clade will highlight the clade *before* it arises; this is by design. If not specified (the default) there will be no focal species; this is fine.
frames	number of frames for animation; this will also determine the time internals for the plot
colours	a function that will return a colour ramp for use in plotting of species on the fiber plot itself as well as the standard phylogeny to the right (e.g., rainbow)
f.colours	as colours but for the focal species
pca	PCA (of class prcomp) of phylogenetic dissimilarity matrix; NULL calculates one, I recommend you use the output from a previous run to speed things up
clade.mat	clade matrix (from $clade.matrix$ clade.matrix) of phylogeny; NULL calculates one, I recommend you use the output from a previous run to speed things up
delay	the delay between each slice's frame in the output GIF; default 0.2 seconds

side.tree whether to plot a standard phylogeny to the right of the plot to aid with interpre-

tation (default: TRUE). You almost certainly want this option

width width of animation height height of animation

Details

Probably best to just plot it out and see what happens, to be honest.

Value

The data that were plotted last, the PCA and clade.matrix to speed later plots, and the colours used.

Note

I would be grateful if you could cite the article this code was released in when using this code. I maintain this code in the package "willeard" on GitHub. I give an example of how to install this code from there below. Updates will be released through that, and I welcome code improvements!

Author(s)

Will Pearse

Examples

```
## Not run:
fibre.plot(rlineage(0.1,0), "Yule_fibre.gif")
## End(Not run)
```

fingerprint.regression

Regress trait evolution against trait ecology (following Cavender-Bares et al. 2004)

Description

Calculates traits' phylogenetic inertia and regresses this against trait similarity among co-existing species (sensu Cavender-Bares et al. 2004 Figure 6)

Usage

```
fingerprint.regression(
  data,
  eco.rnd = c("taxa.labels", "richness", "frequency", "sample.pool", "phylogeny.pool",
    "independentswap", "trialswap"),
  eco.method = c("quantile", "lm", "mantel"),
  eco.permute = 1000,
```

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```
evo.method = c("lambda", "delta", "kappa", "blom.k"),
  eco.swap = 1000,
  abundance = TRUE,
    ...
)

## S3 method for class 'fingerprint.regression'
print(x, ...)

## S3 method for class 'fingerprint.regression'
summary(object, ...)

## S3 method for class 'fingerprint.regression'
plot(
    x,
    eco = c("slope", "corrected"),
    xlab = "Community Trait Similarity",
    ylab = "Phylogenetic inertia",
    ...
)
```

Arguments

data	comparative.comm for analysis
eco.rnd	null distribution with which to compare your community data, one of: taxa.labels (DEFAULT), richness, frequency, sample.pool, phylogeny.pool, independentswap, trialswap (as implemented in picante)
eco.method	how to compare distance matrices (only the lower triangle;), one of: lm (linear regression), quantile (DEFAULT; rq), mantel (mantel)
eco.permute	number of permutations for ecological null model (eco.rnd); default 1000
evo.method	how to measure phylogenetic inertia, one of: lambda (default), delta, kappa, blom.k; see phy.signal.
eco.swap	number of independent swap iterations to perform (if specified in eco.rnd; DE-FAULT 1000)
abundance	whether to incorporate species' abundances (default: TRUE)
	additional parameters to pass on to model fitting functions and plotting functions
x	fingerprint.regression object
object	fingerprint.regression object
eco	plot the observed slopes (DEFAULT: slope), or the median difference between the simulations and the observed values (corrected)
xlab	label for x-axis (default "Ecological Trait Coexistence")
ylab	label for y-axis (default "Phylogenetic inertia")

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Details

While the term 'fingerprint regression' is new to pez, the method is very similar to that employed in Cavender-Bares et al. 2004 Figure 6. For each trait, the phylogenetic inertia of species traits is regressed against their co-occurrence in the community matrix. Note that Pagel's

, λ , and κ

, and Blomberg's K, can be used, unlike the original where a mantel test was employed. Moreover, note also that Pianka's distance (as described in the manuscript) is used to measure species overlap.

Note

Like eco.xxx.regression, this is a data-hungry method. Warnings will be generated if any of the methods cannot be fitted properly (the examples below give toy examples of this). In such cases the summary and plot methods of these functions may generate errors; perhaps using traceback to examine where these are coming from, and consider whether you want to be working with the data generating these errors. I am loathe to hide these errors or gloss over them, because they represent the reality of your data!

WDP loves quantile regressions, and advises that you check different quantiles using the tau options.

Author(s)

Will Pearse and Jeannine Cavender-Bares

References

Cavender-Bares J., Ackerly D.D., Baum D.A. & Bazzaz F.A. (2004) Phylogenetic overdispersion in Floridian oak communities. The Americant Naturalist 163(6): 823–843.

Kembel, S.W., Cowan, P.D., Helmus, M.R., Cornwell, W.K., Morlon, H., Ackerly, D.D., Blomberg, S.P. & Webb, C.O. Picante: R tools for integrating phylogenies and ecology. Bioinformatics 26(11): 1463–1464.

Pagel M. Inferring the historical patterns of biological evolution. Nature 401(6756): 877–884.

See Also

```
eco.xxx.regression phy.signal
```

Examples

```
data(laja)
data <- comparative.comm(invert.tree, river.sites, invert.traits, river.env)
fingerprint.regression(data, eco.permute=10)
plot(fingerprint.regression(data, permute=10, method="lm"))</pre>
```

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generic.metrics

Calculate any metric(s) (and compare with null distributions)

Description

Allow the calculation of any metric within pez.

Usage

```
generic.null(
  data,
 metrics,
  null.model = c("taxa.labels", "richness", "frequency", "sample.pool",
    "phylogeny.pool", "independentswap", "trialswap", "trait.asm"),
  permute = 1000,
  comp.fun = .ses,
)
.ses(observed, null)
.metric.null(
  data,
 metrics,
 null.model = c("taxa.labels", "richness", "frequency", "sample.pool",
    "phylogeny.pool", "independentswap", "trialswap", "trait.asm"),
  permute = 1000,
  trait = -1,
)
generic.metrics(data, metrics, ...)
```

Arguments

data	data comparative.comm object
metrics	vector of functions to be calculated on data; see pez.metrics for a list of them.
null.model	one of "taxa.labels", "richness", "frequency", "sample.pool", "phylogeny.pool", "independentswap", or "independentswap". These correspond to the null models available in picante
permute	number of null permutations to perform (default 1000)
comp.fun	comparison function to compare observed values with null values. Default is .ses; this is a Standard Effect Size (obs - mean)/SEmean. You may supply your own function; it should take the observed site-metric matrix as its first argument, and a site-metric-permutation array as its second. See the internals of generic.null for an example of its use.

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additional arguments (e.g, dist, abundance.weighted) to be passed to any metric functions (see generic.metrics for possible arguments)

observed observed metric values in site-metric matrix (e.g., from generic.metrics)

null distributions (e.g., from .metric.null) in a site-metric-permutation array

trait if using trait.asm null.model, which trait to use (as in trait.asm)

Details

generic.null Calculate metrics and compare with null distributions. Very light wrapper around the utility functions generic.null and generic.metrics (which is, itself, a very simple function!).

Value

generic.null Output from comp.fun, by default an array (site-metric-type), where type is the observed value, the mean of the null permutations, the Standard Error of that mean, the Standard Effect Size of the metric (obs-null.mean)/SE, and then the rank of the observed value in the permutations. The rank can be considered a bootstrapped p-value of significance, but remember that this is a rank: at the 95 would be significant.

. ses Vector of standard effect sizes

.metric.null site-metric-permutation array

generic.metrics site-metric matrix

Note

comp. fun can be *anything*; much ink has been written about the use of standard effect sizes in ecophylogenetic analyses (*e.g.*, Kembel 2009). That this function makes it easy for you to compute Standard Effect Sizes does not necessarily mean that you should (see Pearse et al. 2013).

Calculating null permutations on a dispersion metric makes little sense, since (by definition; see Pearse et al. 2014) a dispersion metric *require* the use of a null distribution to be calculated. There is nothing to stop you doing so, however! The code makes no attempt to stop you calculating null dissimilarity metrics, but I am not certain that doing so is a good idea using this code as I don't know what to do with such null models!

The pez.shape, pez.evenness, pez.dispersion, and pez.dissimilarity wrapper functions go to some trouble to stop you calculating metrics using inappropriate data (see their notes). These functions give you access to the underlying code within pez; there is nothing I can do to stop you calculating a metric that, in my opinion, doesn't make any sense. You have been warned:D

Author(s)

Will Pearse

References

Kembel S.W. (2009) Disentangling niche and neutral influences on community assembly: assessing the performance of community phylogenetic structure tests. Ecology letters, 12(9), 949-960.

Pearse W.D., Jones F.A. & Purvis A. (2013) Barro Colorado Island's phylogenetic assemblage structure across fine spatial scales and among clades of different ages. Ecology, 94(12), 2861-2872.

laja 23

Examples

```
#Setup data
data(laja)
data <- comparative.comm(invert.tree, river.sites, invert.traits)
#Calculate some metrics
generic.metrics(data, c(.mpd, .pse))
#Compare with a trait-based null model (trait.asm)
generic.null(data, c(.mpd, .pse), "trait.asm", permute=10, trait="fish.pref")
#...be patient when running large (e.g., 1000) sets of null simulations
#You can also do this in pieces, giving even more flexibility
observed <- generic.metrics(data, c(.mpd, .pse))
#null <- .metric.null(data, c(.mpd, .pse))
#ses <- .ses(observed, null)
#...this is how everything works within generic.null
#...and, as with everything in pez, all internal functions start with a "."</pre>
```

laja

Macroinvertebrate samples from the Rio Laja of Mexico

Description

This data set includes macroinvertebrate samples from the Rio Laja, a phylogenetic tree of the taxa and traits that include mean body length and fish feeding preference as in Helmus *et al.* 2013.

Usage

```
data(laja)
```

Format

laja contains a phylo object, a dataframe of sites-by-taxa, a dataframe of sites-by-environment, and a dataframe of traits

Author(s)

M.R. Helmus

References

Helmus M., Mercado-Silva N. & Vander Zanden M.J. (2013). Subsidies to predators, apparent competition and the phylogenetic structure of prey communities. Oecologia, 173, 997-1007.

Examples

```
data(laja)
data <- comparative.comm(invert.tree, river.sites, invert.traits, river.env)</pre>
```

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pez

Phylogenetics for the Environmental Sciences

Description

Analysis and manipulation of eco-phylogenetic datasets containing species phylogeny, species traits, community composition, and environmental data. Provide the comparative.comm object to ease data manipulation, and wrappers for common community phylogenetic indices grouped according to Pearse et al. 2014: pez.shape, pez.evenness, pez.dispersion, and pez.dissimilarity. Implementation of Cavender-Bares et al. (2004) correlation of phylogenetic and ecological matrices (fingerprint.regression). Simulation of null assemblages, traits, and phylogenies (scape, sim.meta.comm).

References

Pearse W.D., Purvis A., Cavender-Bares J. & Helmus M.R. (2014). Metrics and Models of Community Phylogenetics. In: Modern Phylogenetic Comparative Methods and Their Application in Evolutionary Biology. Springer Berlin Heidelberg, pp. 451-464.

Cavender-Bares J., Ackerly D.D., Baum D.A. & Bazzaz F.A. (2004) Phylogenetic overdispersion in Floridian oak communities. The Americant Naturalist 163(6): 823–843.

Examples

```
require(pez)
data(laja)
data <- comparative.comm(invert.tree, river.sites, invert.traits, river.env)
pez.shape(data)</pre>
```

pez.dispersion

Calculate (phylogenetic) dispersion: examine assemblages in the context of a source pools

Description

As described in Pearse et al. (2014), a dispersion metric is one the examines the phylogenetic structure of species present in each assemblage in the context of a source pool of potentially present species. Unlike other metrics, the value of a dispersion metric is *contingent* on the definition of source pool, and (often) randomisations used to conduct that comparison. For completeness, options are provided to calculate these metrics using species traits.

pez.dispersion 25

Usage

```
pez.dispersion(
  data,
  null.model = c("taxa.labels", "richness", "frequency", "sample.pool",
        "phylogeny.pool", "independentswap", "trialswap"),
  abundance = FALSE,
  sqrt.phy = FALSE,
  traitgram = NULL,
  traitgram.p = 2,
  ext.dist = NULL,
  permute = 1000,
  ...
)
```

Arguments

data	comparative.comm object
null.model	one of "taxa.labels", "richness", "frequency", "sample.pool", "phylogeny.pool", "independentswap", or "independentswap". These correspond to the null models available in picante; only d does not use these null models
abundance	Whether to use abundance-weighted forms of these metrics (default: FALSE). D, which is presence/absence only, and so will not be calculated when TRUE.
sqrt.phy	If TRUE (default is FALSE) your phylogenetic distance matrix will be square-rooted; specifying TRUE will force the square-root transformation on phylogenetic distance matrices (in the spirit of Leitten and Cornwell, 2014). See 'details' for details about different metric calculations when a distance matrix is used.
traitgram	If not NULL (default), a number to be passed to funct.phylo.dist (phyloWeight; the 'a' parameter), causing analysis on a distance matrix reflecting both traits and phylogeny (0 -> only phylogeny, 1 -> only traits; see funct.phylo.dist). If a vector of numbers is given, pez.dispersion iterates across them and returns a data.frame with coefficients from each iteration. See 'details' for details about different metric calculations when a distance matrix is used.
traitgram.p	$A\ value\ for\ `p'\ to\ be\ used\ in\ conjunction\ with\ \texttt{traitgram}\ when\ calling\ \texttt{funct.phylo.dist}.$
ext.dist	Supply an external species-level distance matrix for use in calculations. See 'details' for comments on the use of distance matrices in different metric calculations.
permute	number of null permutations to perform (default 1000)
	additional parameters to be passed to metrics (unlikely you will want to use this!)

Details

Most of these metrics do not involve comparison with some kind of evolutionary-derived expectation for phylogenetic shape. Those that do, however, such as D, make no sense unless applied to a phylogenetic distance matrix - their null expectation *requires* it. Using square-rooted distance matrices, or distance matrices that incorporate trait information, can be an excellent thing to do, but

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(for the above reasons), pez won't give you an answer for metrics for which WDP thinks it makes no sense. SESpd can (...up to you whether it should!...) be used with a square-rooted distance matrix, but the results *will always be wrong* if you do not have an ultrametric tree (branch lengths proportional to time) and you will be warned about this. WDP strongly feels you should only be using ultrametric phylogenies in any case, but code to fix this bug is welcome.

Value

a data. frame with metric values

Author(s)

M.R. Helmus, Will Pearse

References

Pearse W.D., Purvis A., Cavender-Bares J. & Helmus M.R. (2014). Metrics and Models of Community Phylogenetics. In: Modern Phylogenetic Comparative Methods and Their Application in Evolutionary Biology. Springer Berlin Heidelberg, pp. 451-464.

sesmpd, sesmntd Webb C.O. (2000). Exploring the phylogenetic structure of ecological communities: An example for rain forest trees. American Naturalist, 156, 145-155.

sespd Webb C.O., Ackerly D.D. & Kembel S.W. (2008). Phylocom: software for the analysis of phylogenetic community structure and trait evolution. Bioinformatics Applications Note, 24, 2098-2100.

innd, mipd Ness J.H., Rollinson E.J. & Whitney K.D. (2011). Phylogenetic distance can predict susceptibility to attack by natural enemies. Oikos, 120, 1327-1334.

d Fritz S.A. & Purvis A. (2010). Selectivity in Mammalian Extinction Risk and Threat Types: a New Measure of Phylogenetic Signal Strength in Binary Traits. Conservation Biology, 24, 1042-1051.

See Also

```
pez.shape pez.evenness pez.dissimilarity
```

Examples

```
data(laja)
data <- comparative.comm(invert.tree, river.sites, invert.traits)
## Not run: pez.dispersion(data)
pez.dispersion(data, permute = 100)</pre>
```

pez.dissimilarity 27

 $\begin{tabular}{ll} \it Pez.dissimilarity: compare assemblages to one- \\ \it another \end{tabular}$

Description

As described in Pearse et al. (2014), a dissimilarity metric compares diversity between communities. WARNING: Phylosor is presented as a distance matrix here, i.e. it is *not* the fraction of shared branch length among communities, but rather '1 - shared branch length'. This means dissimilarity always returns a *distance* object, not a similarity object; this is a different convention from other packages.

Usage

```
pez.dissimilarity(
  data,
  metric = c("all", "unifrac", "pcd", "phylosor", "comdist"),
  abundance.weighted = FALSE,
  permute = 1000,
  sqrt.phy = FALSE,
  traitgram = NULL,
  traitgram.p = 2,
  ext.dist = NULL,
  ...
)
```

Arguments

data comparative.comm object

metric default (all) calculates everything; individually call-able metrics are: unifrac,

pcd, phylosor, comdist.

abundance.weighted

If TRUE (default is FALSE) metrics are calculated incorporating species abun-

dances; only comdist can incorporate abundances

permute Number of permutations for metric (currently only for pcd)

sqrt.phy If TRUE (default is FALSE) your phylogenetic distance matrix will be square-

rooted; specifying TRUE will force the square-root transformation on phylogenetic distance matrices (in the spirit of Leitten and Cornwell, 2014). See 'details' for details about different metric calculations when a distance matrix is used.

traitgram If not NULL (default), a number to be passed to funct.phylo.dist(phyloWeight;

the 'a' parameter), causing analysis on a distance matrix reflecting both traits and phylogeny (0 -> only phylogeny, 1 -> only traits; see funct.phylo.dist). Unlike other metric wrapper functions, dissimilarity does not accept a vector of traitgram values; call the function many times to get these. This is simply because it can take so long: you're probably better off looping/apply-ing over this function yourself.

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traitgram.p A value for 'p' to be used in conjunction with traitgram when calling funct.phylo.dist.

ext.dist Supply an external species-level distance matrix for use in calculations. See

'details' for comments on the use of distance matrices in different metric calcu-

lations.

... additional parameters to be passed to 'metric function(s) you are calling

Details

Using square-rooted distance matrices, or distance matrices that incorporate trait information, can be an excellent thing to do, but (for the above reasons), pez won't give you an answer for metrics for which WDP thinks it makes no sense. All results from this other than comdist *will always be wrong* if you do not have an ultrametric tree and square-root (branch lengths proportional to time) and you will be warned about this. WDP strongly feels you should only be using ultrametric phylogenies in any case, but code to fix this bug is welcome.

Value

list object of metric values.

Author(s)

M.R. Helmus, Will Pearse

References

Pearse W.D., Purvis A., Cavender-Bares J. & Helmus M.R. (2014). Metrics and Models of Community Phylogenetics. In: Modern Phylogenetic Comparative Methods and Their Application in Evolutionary Biology. Springer Berlin Heidelberg, pp. 451-464.

unifrac Lozupone C.A. & Knight R. (2005). UniFrac: a new phylogenetic method for comparing microbial communities. Applied and Environmental Microbiology, 71, 8228-8235.

pcd Ives A.R. & Helmus M.R. (2010). Phylogenetic metrics of community similarity. The American Naturalist, 176, E128-E142.

phylosor Bryant J.A., Lamanna C., Morlon H., Kerkhoff A.J., Enquist B.J. & Green J.L. (2008). Microbes on mountainsides: Contrasting elevational patterns of bacterial and plant diversity. Proceedings of the National Academy of Sciences of the United States of America, 105, 11505-11511.

comdist C.O. Webb, D.D. Ackerly, and S.W. Kembel. 2008. Phylocom: software for the analysis of phylogenetic community structure and trait evolution. Bioinformatics 18:2098-2100.

See Also

```
pez.shape pez.evenness pez.dispersion
```

Examples

```
data(laja)
data <- comparative.comm(invert.tree, river.sites, invert.traits)
## Not run:
dissim <- pez.dissimilarity(data)</pre>
```

pez.endemism 29

End(Not run)

pez.endemism

Calculate (phylogenetic) endemism

Description

At present, only a small number of metrics, but we intend for this to grow with time. Note that metrics that incorporate abundance are mixed in with those that do not. Some of these metrics make sense when used with probabilities, for example those derived from an SDM; some do not. You will have to use your own judgement (as with everything in science!).

Usage

```
pez.endemism(data, sqrt.phy = FALSE)
```

Arguments

data comparative.comm object

sqrt.phy If TRUE (default is FALSE) your phylogenetic distance matrix will be square-

rooted; specifying TRUE will force the square-root transformation on phylogenetic distance matrices (in the spirit of Leitten and Cornwell, 2014). See 'details' for details about different metric calculations when a distance matrix is used.

Value

data. frame with metric values.

Author(s)

Will Pearse, Dan Rosauer

References

BED Cadotte, M. W., & Jonathan Davies, T. (2010). Rarest of the rare: advances in combining evolutionary distinctiveness and scarcity to inform conservation at biogeographical scales. Diversity and Distributions, 16(3), 376-385.

PE Rosauer, D. A. N., Laffan, S. W., Crisp, M. D., Donnellan, S. C., & Cook, L. G. (2009). Phylogenetic endemism: a new approach for identifying geographical concentrations of evolutionary history. Molecular Ecology, 18(19), 4061-4072.

See Also

pez.shape pez.evenness pez.dispersion pez.dissimilarity

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Examples

```
data(laja)
data <- comparative.comm(invert.tree, river.sites, invert.traits)
(output<-pez.endemism(data))</pre>
```

pez.evenness

Calculate (phylogenetic) evenness: examine assemblage composition and abundance

Description

As described in Pearse et al. (2014), an evenness metric is one the examines the phylogenetic structure of species present in each assemblage, taking into account their abundances. For completeness, options are provided to calculate these metrics using species traits.

Usage

```
pez.evenness(
  data,
  sqrt.phy = FALSE,
  traitgram = NULL,
  traitgram.p = 2,
  ext.dist = NULL,
  quick = TRUE,
  q = 1
)
```

Arguments

data	comparative.	comm object
------	--------------	-------------

sqrt.phy If TRUE (default is FALSE) your phylogenetic distance matrix will be square-

rooted; specifying TRUE will force the square-root transformation on phylogenetic distance matrices (in the spirit of Leitten and Cornwell, 2014). See 'details' for details about different metric calculations when a distance matrix is used.

traitgram If not NULL (default), a number to be passed to funct.phylo.dist(phyloWeight;

the 'a' parameter), causing analysis on a distance matrix reflecting both traits and phylogeny (0->only phylogeny, 1-> only traits; see funct.phylo.dist). If a vector of numbers is given, pez.eveness iterates across them and returns a data.frame with coefficients from each iteration. See 'details' for details about

different metric calculations when a distance matrix is used.

traitgram.p A value for 'p' to be used in conjunction with traitgram when calling funct.phylo.dist.

ext.dist Supply an external species-level distance matrix for use in calculations. See

'details' for comments on the use of distance matrices in different metric calcu-

lations.

pez.evenness 31

quick

Only calculate metrics which are quick to calculate (default: TRUE); setting to FALSE will also calculate fd.dist and the Pagel transformations (

 λ , δ , κ).

q value for q in scheiner (default 1)

Details

Most of these metrics do not involve comparison with some kind of evolutionary-derived expectation for phylogenetic shape. Those that do, however, such as PSE, make no sense unless applied to a phylogenetic distance matrix - their null expectation *requires* it. Using square-rooted distance matrices, or distance matrices that incorporate trait information, can be an excellent thing to do, but (for the above reasons), pez won't give you an answer for metrics for which WDP thinks it makes no sense. pae, iac, haead & eaed can (...up to you whether you should!...) be used with a square-rooted distance matrix, but the results *will always be wrong* if you do not have an ultrametric tree (branch lengths proportional to time) and you will be warned about this. WDP strongly feels you should only be using ultrametric phylogenies in any case, but code to fix this bug is welcome.

Value

phy. structure list object of metric values. Use coefs to extract a summary metric table, or examine each individual metric (which gives more details for each) by calling print on the output (i.e., type output in the example below).

Note

As mentioned above, dist.fd is calculated using a phylogenetic distance matrix if no trait data are available, or if you specify sqrt.phy. It is not calculated by default because it generates warning messsages (which WDP is loathe to suppress) which are related to the general tendency for a low rank of phylogenetic distance matrices. Much ink has been written about this, and in par this problem is why the eigen.sum measure came to be suggested.

Some of these metrics can cause (inconsequential) warnings if given assemblages with only one species/individual in them, and return NA/NaN values depending on the metric. I consider these 'features', not bugs.

Some of the metrics in this wrapper are also in pez.shape; such metrics can be calculated using species' abundances (making them *evenness*) metrics or simply using presence/absence of species (making them *shape* metrics).

Author(s)

M.R. Helmus, Will Pearse

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References

Pearse W.D., Purvis A., Cavender-Bares J. & Helmus M.R. (2014). Metrics and Models of Community Phylogenetics. In: Modern Phylogenetic Comparative Methods and Their Application in Evolutionary Biology. Springer Berlin Heidelberg, pp. 451-464.

pse Helmus M.R., Bland T.J., Williams C.K. & Ives A.R. (2007). Phylogenetic measures of biodiversity. American Naturalist, 169, E68-E83.

Pearse W.D., Purvis A., Cavender-Bares J. & Helmus M.R. (2014). Metrics and Models of Community Phylogenetics. In: Modern Phylogenetic Comparative Methods and Their Application in Evolutionary Biology. Springer Berlin Heidelberg, pp. 451-464.

pse Helmus M.R., Bland T.J., Williams C.K. & Ives A.R. (2007). Phylogenetic measures of biodiversity. American Naturalist, 169, E68-E83.

rao Webb C.O. (2000). Exploring the phylogenetic structure of ecological communities: An example for rain forest trees. American Naturalist, 156, 145-155.

taxon Clarke K.R. & Warwick R.M. (1998). A taxonomic distinctness index and its statistical properties. J. Appl. Ecol., 35, 523-531.

entropy Allen B., Kon M. & Bar-Yam Y. (2009). A New Phylogenetic Diversity Measure Generalizing the Shannon Index and Its Application to Phyllostomid Bats. The American Naturalist, 174, 236-243.

pae, iac, haed, eaed Cadotte M.W., Davies T.J., Regetz J., Kembel S.W., Cleland E. & Oakley T.H. (2010). Phylogenetic diversity metrics for ecological communities: integrating species richness, abundance and evolutionary history. Ecology Letters, 13, 96-105.

lambda, delta, kappa Mark Pagel (1999) Inferring the historical patterns of biological evolution. Nature 6756(401): 877–884.

innd, mipd Ness J.H., Rollinson E.J. & Whitney K.D. (2011). Phylogenetic distance can predict susceptibility to attack by natural enemies. Oikos, 120, 1327-1334.

scheiner Scheiner, S.M. (20120). A metric of biodiversity that integrates abundance, phylogeny, and function. Oikos, 121, 1191-1202.

See Also

pez.shape pez.dispersion pez.dissimilarity

Examples

```
data(laja)
data <- comparative.comm(invert.tree, river.sites, invert.traits)
pez.evenness(data)</pre>
```

pez.metrics

Phylogenetic and functional trait metrics within pez

Description

Using these functions, you can calculate any of the phylogenetic metrics within pez, using comparative.comm objects. While you can call each individually, using the pez.shape, pez.evenness, pez.dispersion, and pez.dissimilarity wrapper functions (and the more flexible generic.metrics and null model functions) are probably your best bet. Note that *all of these functions* take a common first parameter: a comparative.comm object. There are additional parameters that can be passed, which are described below.

Usage

```
.hed(x, ...)
.eed(x, na.rm = TRUE, ...)
.psv(x, ...)
.psr(x, ...)
.mpd(x, dist = NULL, abundance.weighted = FALSE, ...)
.vpd(x, dist = NULL, abundance.weighted = FALSE, ...)
.vntd(x, dist = NULL, abundance.weighted = FALSE, ...)
.pd(x, include.root = TRUE, abundance.weighted = FALSE, ...)
.mntd(x, dist = NULL, abundance.weighted = FALSE, ...)
.gamma(x, ...)
.taxon(x, dist = NULL, abundance.weighted = FALSE, ...)
.eigen.sum(x, dist = NULL, which.eigen = 1, ...)
.dist.fd(x, method = "phy", abundance.weighted = FALSE, ...)
.sqrt.phy(x)
.phylo.entropy(x, ...)
.aed(x, ...)
.haed(x, ...)
```

```
.simpson.phylogenetic(x)
.iac(x, na.rm = TRUE, ...)
.pae(x, na.rm = TRUE, ...)
.scheiner(x, q = 0, abundance.weighted = FALSE, ...)
.pse(x, ...)
.rao(x, ...)
.lambda(x, ...)
.delta(x, ...)
.kappa(x, ...)
.eaed(x, ...)
.unifrac(x, ...)
.pcd(x, permute = 1000, ...)
.comdist(x, dist = NULL, abundance.weighted = FALSE, ...)
.phylosor(x, dist = NULL, abundance.weighted = FALSE, ...)
.d(x, permute = 1000, ...)
.ses.mpd(
  х,
 dist = NULL,
 null.model = "taxa.labels",
 abundance.weighted = FALSE,
 permute = 1000,
)
.ses.mntd(
 х,
 dist = NULL,
 null.model = "taxa.labels",
 abundance.weighted = FALSE,
 permute = 1000,
)
```

```
.ses.vpd(
  Х,
 dist = NULL,
 null.model = "taxa.labels",
 abundance.weighted = FALSE,
 permute = 1000,
)
.ses.vntd(
 х,
 dist = NULL,
 null.model = "taxa.labels",
 abundance.weighted = FALSE,
 permute = 1000,
)
.ses.mipd(
 х,
 dist = NULL,
 null.model = "taxa.labels",
 abundance.weighted = FALSE,
 permute = 1000,
)
.ses.innd(
 dist = NULL,
 null.model = "taxa.labels",
 abundance.weighted = FALSE,
 permute = 1000,
)
.mipd(x, dist = NULL, abundance.weighted = FALSE, ...)
.innd(x, dist = NULL, abundance.weighted = FALSE, ...)
.innd(x, dist = NULL, abundance.weighted = FALSE, ...)
.pe(x, ...)
.bed(x, ...)
```

Arguments

x comparative.comm object
... ignored

na.rm remove NAs in calculations (altering this can obscure errors that are meaningful;

I would advise leaving alone)

dist distance matrix for use with calculations; could be generated from traits, a

square-root-transformed distance matrix (see . sqrt.phy for creating a comparative.comm

object with a square-root transformed phylogeny). Default: NULL (-> calculate

distance matrix from phylogeny)

abundance.weighted

whether to include species' abundances in metric calculation, often dictating whether you're calculating a pez.shape or pez.evenness metric. Default:

FALSE

include.root include root in PD calculations (default is TRUE, as in picante, but within

pez. shape I specify FALSE

which eigen which phylo-eigenvector to be used for PVR metric

method whether to calculate using phylogeny ("phy"; default) or trait data ("traits")

q the q parameter for . scheiner; default 0.0001

permute number of permutations of null randomisations (mostly only applies to dispersion

metrics)

null.model one of "taxa.labels", "richness", "frequency", "sample.pool", "phylogeny.pool",

"independentswap", or "independentswap". These correspond to the null models

available in picante; only d does not use these null models

Details

.pd returns two metrics: Faith's PD (which does not take into account abundance) and Faith's PD corrected for species richness or total abundance (depending on abundance.weighted). I am almost certain that I got the idea for this from somewhere, but I can't find the reference: if you published on this before 2012, please get in touch with me.

. scheiner has a different formula for the case where q is equal to 1 (check the code if interested). The nature of its definition means that values very close to, but not exactly equal to, 1 may be extremely large or extremely small. This is a feature, not a bug, and an inherent aspect of its definition. Check the formula in the code for more information!

Note

Many (but not all) of these functions are fairly trivial wrappers around functions in other packages. In the citations for each metric, * indicates a function that's essentially written in picante. The Pagel family of measures are also fairly trivial wrapper around caper code, functional dissimilarity FD code, gamma, and ape code. I can't demand it, but I would be grateful if you would cite these authors when using these wrappers.

The pez.shape, pez.evenness, pez.dispersion, and pez.dissimilarity wrapper functions go to some trouble to stop you calculating metrics using inappropriate data (see their notes). These

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functions give you access to the underlying code within pez; there is nothing I can do to stop you calculating a metric that, in my opinion, doesn't make any sense. You have been warned:D

If you're a developer hoping to make your metric(s) work in this framework, please use the argument naming convention for arguments described in this help file, and use the ... operator in your definition. That way functions that don't need particular arguments can co-exist peacefully with those that do. The first argument to one of these functions should *always* be a comparative.comm object; there is no method dispatch on any of these functions and I foresee future pain without this rule.

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Examples

```
data(laja)
data <- comparative.comm(invert.tree, river.sites)
.psv(data)</pre>
```

pez.shape

Calculate (phylogenetic) shape: examine assemblage composition

Description

As described in Pearse et al. (2014), a shape metric is one the examines the phylogenetic structure of species present in each assemblage, ignoring abundances entirely. For completeness, options are provided to calculate these metrics using species traits.

Usage

```
pez.shape(
  data,
  sqrt.phy = FALSE,
  traitgram = NULL,
  traitgram.p = 2,
  ext.dist = NULL,
  which.eigen = 1,
  quick = TRUE,
  q = 1e-04
)
```

pez.shape 39

Arguments

data comparative.comm object If TRUE (default is FALSE) your phylogenetic distance matrix will be squaresgrt.phy rooted; specifying TRUE will force the square-root transformation on phylogenetic distance matrices (in the spirit of Leitten and Cornwell, 2014). See 'details' for details about different metric calculations when a distance matrix is used. traitgram If not NULL (default), a number to be passed to funct.phylo.dist (phyloWeight; the 'a' parameter), causing analysis on a distance matrix reflecting both traits and phylogeny $(0 \rightarrow \text{only phylogeny}, 1 \rightarrow \text{only traits}; \text{ see funct.phylo.dist})$. If a vector of numbers is given, pez. shape iterates across them and returns a data. frame with coefficients from each iteration. See 'details' for details about different metric calculations when a distance matrix is used. traitgram.p A value for 'p' to be used in conjunction with traitgram when calling funct.phylo.dist. ext.dist Supply an external species-level distance matrix for use in calculations. See 'details' for comments on the use of distance matrices in different metric calculations. The eigen vector to calculate for the PhyloEigen metric (eigen.sum) which.eigen quick Only calculate metrics which are quick to calculate (default: TRUE); setting to FALSE will also calculate fd.dist.

Details

q

Most of these metrics do not involve comparison with some kind of evolutionary-derived expectation for phylogenetic shape. Those that do, however, such as PSV, make no sense unless applied to a phylogenetic distance matrix - their null expectation *requires* it. Using square-rooted distance matrices, or distance matrices that incorporate trait information, can be an excellent thing to do, but (for the above reasons), pez won't give you an answer for metrics for which WDP thinks it makes no sense. pd, eed & hed can (...up to you whether you should!...) be used with a square-rooted distance matrix, but the results *will always be wrong* if you do not have an ultrametric tree (branch lengths proportional to time) and you will be warned about this. WDP strongly feels you should only be using ultrametric phylogenies in any case, but code to fix this bug is welcome.

value for q in scheiner (default 0.0001)

Value

phy.structure list object of metric values. Use coefs to extract a summary metric table, or examine each individual metric (which gives more details for each) by calling print on the output (i.e., type output in the example below).

Some of the metrics in this wrapper are also in pez.evenness; such metrics can be calculated using species' abundances (making them *evenness*) metrics or simply using presence/absence of species (making them *shape* metrics).

Note

As mentioned above, dist.fd is calculated using a phylogenetic distance matrix if no trait data are available, or if you specify sqrt.phy. It is not calculated by default because it generates warning

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messsages (which WDP is loathe to suppress) which are related to the general tendency for a low rank of phylogenetic distance matrices. Much ink has been written about this, and in part this problem is why the eigen.sum measure came to be suggested.

Many of these metrics, (e.g., eed) will cause (inconsequential) warnings if given assemblages with only one species in them, and return NA/NaN values depending on the metric. I consider these 'features', not bugs.

Author(s)

M.R. Helmus, Will Pearse

References

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See Also

```
pez.evenness pez.dispersion pez.dissimilarity
```

Examples

```
data(laja)
data <- comparative.comm(invert.tree, river.sites, invert.traits)
(output<-pez.shape(data))</pre>
```

pglmm

Phylogenetic Generalised Linear Mixed Model for Community Data

Description

This function performs Generalized Linear Mixed Models for binary and continuous phylogenetic data, estimating regression coefficients with approximate standard errors. It is modeled after lmer but is more general by allowing correlation structure within random effects; these correlations can be phylogenetic among species, or any other correlation structure, such as geographical correlations among sites. It is, however, much more specific than lmer in that it can only analyze a subset of1 the types of model designed handled by lmer. It is also much slower than lmer and requires users to specify correlation structures as covariance matrices. communityPGLMM can analyze models in Ives and Helmus (2011). It can also analyze bipartite phylogenetic data, such as that analyzed in Rafferty and Ives (2011), by giving sites phylogenetic correlations.

Usage

```
communityPGLMM(
  formula,
  data = list(),
  family = "gaussian",
  sp = NULL,
  site = NULL,
  random.effects = list(),
 REML = TRUE,
  s2.init = NULL,
 B.init = NULL,
  reltol = 10^{-6},
 maxit = 500,
  tol.pql = 10^{-6},
 maxit.pql = 200,
  verbose = FALSE
)
communityPGLMM.gaussian(
  formula,
  data = list(),
  family = "gaussian",
  sp = NULL,
  site = NULL,
  random.effects = list(),
  REML = TRUE,
  s2.init = NULL,
 B.init = NULL,
  reltol = 10^-8,
 maxit = 500,
  verbose = FALSE
```

```
)
communityPGLMM.binary(
  formula,
  data = list(),
  family = "binomial",
  sp = NULL,
  site = NULL,
  random.effects = list(),
 REML = TRUE,
 s2.init = 0.25,
 B.init = NULL,
  reltol = 10^{-5},
 maxit = 40,
  tol.pql = 10^{-6},
 maxit.pql = 200,
  verbose = FALSE
)
communityPGLMM.binary.LRT(x, re.number = 0, ...)
communityPGLMM.matrix.structure(
  formula,
  data = list(),
  family = "binomial",
  sp = NULL,
  site = NULL,
 random.effects = list(),
 ss = 1
)
## S3 method for class 'communityPGLMM'
summary(object, digits = max(3, getOption("digits") - 3), ...)
## S3 method for class 'communityPGLMM'
plot(x, digits = max(3, getOption("digits") - 3), ...)
communityPGLMM.predicted.values(x, show.plot = TRUE, ...)
```

Arguments

data

formula a two-sided linear formula object describing the fixed-effects of the model; for example, Y ~ X.

a data. frame containing the variables named in formula. The data frame should have long format with factors specifying species and sites. communityPGLMM will reorder rows of the data frame so that species are nested within sites. Please note that calling as.data.frame.comparative.comm will return your comparative.comm object into this format for you.

family either gaussian for a Linear Mixed Model, or binomial for binary dependent

data.

sp a factor variable that identifies species site a factor variable that identifies sites

random. effects a list that contains, for non-nested random effects, lists of triplets of the form

list(X,group = group,covar = V). This is modeled after the lmer formula syntax (X | group) where X is a variable and group is a grouping factor. Note that group should be either your sp or site variable specified in sp and site. The additional term V is a covariance matrix of rank equal to the number of levels of group that specifies the covariances among groups in the random effect X. For nested variable random effects, random.effects contains lists of quadruplets of the form list(X,group1 = group1,covar = V,group2 = group2) where group1

is nested within group2.

REML whether REML or ML is used for model fitting. For the generalized linear mixed

model for binary data, these don't have standard interpretations, and there is no

log likelihood function that can be used in likelihood ratio tests.

s2.init an array of initial estimates of s2 for each random effect that scales the variance.

If s2.init is not provided for family="gaussian", these are estimated using in a clunky way using lm assuming no phylogenetic signal. A better approach is to run link[lme4:lmer]{lmer} and use the output random effects for s2.init.

If s2.init is not provided for family="binomial", these are set to 0.25.

B. init initial estimates of B, a matrix containing regression coefficients in the model

for the fixed effects. This matrix must have dim(B.init)=c(p+1,1), where p is the number of predictor (independent) variables; the first element of B corresponds to the intercept, and the remaining elements correspond in order to the predictor (independent) variables in the formula. If B.init is not provided, these are estimated using in a clunky way using lm or glm assuming no phylogenetic signal. A better approach is to run lmer and use the output fixed effects

for B. init.

reltol a control parameter dictating the relative tolerance for convergence in the opti-

mization; see optim.

maxit a control parameter dictating the maximum number of iterations in the optimiza-

tion; see optim.

tol.pql a control parameter dictating the tolerance for convergence in the PQL estimates

of the mean components of the binomial GLMM.

maxit.pql a control parameter dictating the maximum number of iterations in the PQL

estimates of the mean components of the binomial GLMM.

verbose if TRUE, the model deviance and running estimates of s2 and B are plotted each

iteration during optimization.

x communityPGLMM object

re.number which random.effect in x to be tested

... additional arguments to summary and plotting functions (currently ignored)

ss which of the random.effects to produce object communityPGLMM object to be summarised

digits minimal number of significant digits for printing, as in print.default show.plot if TRUE (default), display plot

Details

The vignette 'pez-pglmm-overview' gives a gentle introduction to using PGLMMS. For linear mixed models (family = 'gaussian'), the function estimates parameters for the model of the form, for example,

$$Y = \beta_0 + \beta_1 x + b_0 + b_1 x$$
$$b_0 Gaussian(0, \sigma_0^2 I_{sp})$$
$$b_1 Gaussian(0, \sigma_0^2 V_{sp})$$
$$\eta Gaussian(0, \sigma^2)$$

where β_0 and β_1 are fixed effects, and V_{sp} is a variance-covariance matrix derived from a phylogeny (typically under the assumption of Brownian motion evolution). Here, the variation in the mean (intercept) for each species is given by the random effect b_0 that is assumed to be independent among species. Variation in species' responses to predictor variable x is given by a random effect b_0 that is assumed to depend on the phylogenetic relatedness among species given by V_{sp} ; if species are closely related, their specific responses to x will be similar. This particular model would be specified as

```
re.1 <-list(1,sp = datsp,covar = diag(nspp)) re.2 <-list(datx,sp = datp,covar = Vsp) z <-communityPGLMM(Y ~ X,data = data,family = "gaussian",random.effects = list(re.1,re.2))
```

The covariance matrix covar is standardized to have its determinant equal to 1. This in effect standardizes the interpretation of the scalar σ^2 . Although mathematically this is not required, it is a very good idea to standardize the predictor (independent) variables to have mean 0 and variance 1. This will make the function more robust and improve the interpretation of the regression coefficients. For categorical (factor) predictor variables, you will need to construct 0-1 dummy variables, and these should not be standardized (for obvious reasons).

For binary generalized linear mixed models (family = 'binomial'), the function estimates parameters for the model of the form, for example,

$$y = \beta_0 + \beta_1 x + b_0 + b_1 x$$
$$Y = logit^{-1}(y)$$
$$b_0 Gaussian(0, \sigma_0^2 I_{sp})$$
$$b_1 Gaussian(0, \sigma_0^2 V_{sp})$$

where β_0 and β_1 are fixed effects, and V_{sp} is a variance-covariance matrix derived from a phylogeny (typically under the assumption of Brownian motion evolution).

```
z <-communityPGLMM(Y ~ X, data = data, family = 'binomial', random.effects = list(re.1,re.2))
```

As with the linear mixed model, it is a very good idea to standardize the predictor (independent) variables to have mean 0 and variance 1. This will make the function more robust and improve the interpretation of the regression coefficients. For categorical (factor) predictor variables, you will need to construct 0-1 dummy variables, and these should not be standardized (for obvious reasons).

Value

an object of class communityPGLMM

formula the formula for fixed effects

data the dataset

family either gaussian or binomial depending on the model fit

random.effects the list of random effects

B estimates of the regression coefficients

B. se approximate standard errors of the fixed effects regression coefficients
 B. cov approximate covariance matrix for the fixed effects regression coefficients

B.zscore approximate Z scores for the fixed effects regression coefficients

B. pvalue approximate tests for the fixed effects regression coefficients being different

from zero

ss random effects' standard deviations for the covariance matrix $\sigma^2 V$ for each ran-

dom effect in order. For the linear mixed model, the residual variance is listed

last

s2r random effects variances for non-nested random effects s2n random effects variances for nested random effects

s2resid for linear mixed models, the residual vairance

logLIK for linear mixed models, the log-likelihood for either the restricted likelihood

(REML=TRUE) or the overall likelihood (REML=FALSE). This is set to NULL for

generalised linear mixed models

AIC for linear mixed models, the AIC for either the restricted likelihood (REML=TRUE)

or the overall likelihood (REML=FALSE). This is set to NULL for generalised

linear mixed models

BIC for linear mixed models, the BIC for either the restricted likelihood (REML=TRUE)

or the overall likelihood (REML=FALSE). This is set to NULL for generalised

linear mixed models

REML whether or not REML is used (TRUE or FALSE)

s2.init the user-provided initial estimates of s2
B.init the user-provided initial estimates of B

Y the response (dependent) variable returned in matrix form

X the predictor (independent) variables returned in matrix form (including 1s in

the first column)

H the residuals. For the generalized linear mixed model, these are the predicted

residuals in the $logit^{-1}$ space

iV the inverse of the covariance matrix for the entire system (of dimension (nsp*nsite)

by (nsp*nsite))

mu predicted mean values for the generalized linear mixed model. Set to NULL for

linear mixed models

sp, sp matrices used to construct the nested design matrix

Zt	the design	matrix	for rando	m effects

St diagonal matrix that maps the random effects variances onto the design matrix

convcode the convergence code provided by optim
niter number of iterations performed by optim

Note

These function *do not* use a comparative.comm object, but you can use as.data.frame.comparative.comm to create a data.frame for use with these functions. The power of this method comes from deciding your own parameters parameters to be determined (the data for regression, the random effects, etc.), and it is our hope that this interface gives you more flexibility in model selection/fitting.

Author(s)

Anthony R. Ives, cosmetic changes by Will Pearse

References

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Rafferty, N. E., and A. R. Ives. 2013. Phylogenetic trait-based analyses of ecological networks. Ecology 94:2321-2333.

Examples

```
## Structure of examples:
# First, a (brief) description of model types, and how they are specified
# - these are *not* to be run 'as-is'; they show how models should be organised
# Second, a run-through of how to simulate, and then analyse, data
# - these *are* to be run 'as-is'; they show how to format and work with data
#First section; brief summary of models and their use####
## Model structures from Ives & Helmus (2011)
# dat = data set for regression (note: *not* an comparative.comm object)
# nspp = number of species
# nsite = number of sites
# Vphy = phylogenetic covariance matrix for species
# Vrepul = inverse of Vphy representing phylogenetic repulsion
# Model 1 (Eq. 1)
re.site <- list(1, site = dat$site, covar = diag(nsite))
re.sp.site <- list(1, sp = dat$sp, covar = Vphy, site = dat$site) # note: nested
z \leftarrow communityPGLMM(freq \sim sp, data = dat, family = "binomial", sp
= dat$sp, site = dat$site, random.effects = list(re.site,
re.sp.site), REML = TRUE, verbose = TRUE, s2.init=.1)
```

```
# Model 2 (Eq. 2)
re.site <- list(1, site = dat$site, covar = diag(nsite))</pre>
re.slope <- list(X, sp = dat$sp, covar = diag(nspp))</pre>
re.slopephy <- list(X, sp = dat$sp, covar = Vphy)</pre>
z <- communityPGLMM(freq ~ sp + X, data = dat, family = "binomial",</pre>
sp = dat$sp, site = dat$site, random.effects = list(re.site,
re.slope, re.slopephy), REML = TRUE, verbose = TRUE, s2.init=.1)
# Model 3 (Eq. 3)
re.site <- list(1, site = dat$site, covar = diag(nsite))</pre>
re.sp.site <- list(1, sp = dat$sp, covar = Vrepul, site = dat$site) # note: nested
z <- communityPGLMM(freq ~ sp*X, data = dat, family = "binomial",</pre>
sp = dat$sp, site = dat$site, random.effects = list(re.site,
re.sp.site), REML = TRUE, verbose = TRUE, s2.init=.1)
## Model structure from Rafferty & Ives (2013) (Eq. 3)
# dat = data set
# npp = number of pollinators (sp)
# nsite = number of plants (site)
# VphyPol = phylogenetic covariance matrix for pollinators
# VphyPlt = phylogenetic covariance matrix for plants
re.a <- list(1, sp = dat$sp, covar = diag(nspp))</pre>
re.b <- list(1, sp = dat$sp, covar = VphyPol)</pre>
re.c <- list(1, sp = dat$sp, covar = VphyPol, dat$site)</pre>
re.d <- list(1, site = dat$site, covar = diag(nsite))</pre>
re.f <- list(1, site = dat$site, covar = VphyPlt)</pre>
re.g <- list(1, site = dat$site, covar = VphyPlt, dat$sp)</pre>
#term h isn't possible in this implementation, but can be done with
available matlab code
z <- communityPGLMM(freq ~ sp*X, data = dat, family = "binomial",</pre>
sp = dat$sp, site = dat$site, random.effects = list(re.a, re.b,
re.c, re.d, re.f, re.g), REML = TRUE, verbose = TRUE, s2.init=.1)
## End(Not run)
#Second section; detailed simulation and analysis ######
#NOTE: this section is explained and annotated in ######
      detail in the vignette 'pez-pglmm-overview'#######
      run 'vignette('pez-pglmm-overview') to read######
# Generate simulated data for nspp species and nsite sites
nspp <- 15
nsite <- 10
# residual variance (set to zero for binary data)
sd.resid <- 0
# fixed effects
heta0 <- 0
beta1 <- 0
```

```
# magnitude of random effects
sd.B0 <- 1
sd.B1 <- 1
# whether or not to include phylogenetic signal in B0 and B1
signal.B0 <- TRUE
signal.B1 <- TRUE
# simulate a phylogenetic tree
phy <- rtree(n = nspp)</pre>
phy <- compute.brlen(phy, method = "Grafen", power = 0.5)</pre>
# standardize the phylogenetic covariance matrix to have determinant 1
Vphy <- vcv(phy)
Vphy <- Vphy/(det(Vphy)^(1/nspp))</pre>
# Generate environmental site variable
X <- matrix(1:nsite, nrow = 1, ncol = nsite)</pre>
X \leftarrow (X - mean(X))/sd(X)
# Perform a Cholesky decomposition of Vphy. This is used to
# generate phylogenetic signal: a vector of independent normal random
# variables, when multiplied by the transpose of the Cholesky
# deposition of Vphy will have covariance matrix equal to Vphy.
iD <- t(chol(Vphy))</pre>
# Set up species-specific regression coefficients as random effects
if (signal.B0 == TRUE) {
b0 \leftarrow beta0 + iD \% rnorm(nspp, sd = sd.B0)
} else {
b0 <- beta0 + rnorm(nspp, sd = sd.B0)
}
if (signal.B1 == TRUE) {
b1 <- beta1 + iD %*% rnorm(nspp, sd = sd.B1)
} else {
b1 <- beta1 + rnorm(nspp, sd = sd.B1)
# Simulate species abundances among sites to give matrix Y that
# contains species in rows and sites in columns
y <- rep(b0, each=nsite)
y \leftarrow y + rep(b1, each=nsite) * rep(X, nspp)
y <- y + rnorm(nspp*nsite) #add some random 'error'
Y <- rbinom(length(y), size=1, prob=exp(y)/(1+exp(y)))
y <- matrix(outer(b0, array(1, dim = c(1, nsite))), nrow = nspp,
ncol = nsite) + matrix(outer(b1, X), nrow = nspp, ncol = nsite)
e <- rnorm(nspp * nsite, sd = sd.resid)
y <- y + matrix(e, nrow = nspp, ncol = nsite)
y <- matrix(y, nrow = nspp * nsite, ncol = 1)</pre>
Y \leftarrow rbinom(n = length(y), size = 1, prob = exp(y)/(1 + exp(y)))
```

```
Y <- matrix(Y, nrow = nspp, ncol = nsite)
# name the simulated species 1:nspp and sites 1:nsites
rownames(Y) <- 1:nspp</pre>
colnames(Y) <- 1:nsite</pre>
par(mfrow = c(3, 1), las = 1, mar = c(2, 4, 2, 2) - 0.1)
matplot(t(X), type = "l", ylab = "X", main = "X among sites")
hist(b0, xlab = "b0", main = "b0 among species")
hist(b1, xlab = "b1", main = "b1 among species")
#Plot out; you get essentially this from plot(your.pglmm.model)
image(t(Y), ylab = "species", xlab = "sites", main = "abundance",
col=c("black","white"))
# Transform data matrices into "long" form, and generate a data frame
YY <- matrix(Y, nrow = nspp * nsite, ncol = 1)
XX <- matrix(kronecker(X, matrix(1, nrow = nspp, ncol = 1)), nrow =
nspp * nsite, ncol = 1)
site <- matrix(kronecker(1:nsite, matrix(1, nrow = nspp, ncol =</pre>
1)), nrow = nspp * nsite, ncol = 1)
sp <- matrix(kronecker(matrix(1, nrow = nsite, ncol = 1), 1:nspp),</pre>
nrow = nspp * nsite, ncol = 1)
dat <- data.frame(Y = YY, X = XX, site = as.factor(site), sp = as.factor(sp))</pre>
# Format input and perform communityPGLMM()
# set up random effects
# random intercept with species independent
re.1 <- list(1, sp = dat$sp, covar = diag(nspp))
# random intercept with species showing phylogenetic covariances
re.2 <- list(1, sp = dat$sp, covar = Vphy)
# random slope with species independent
re.3 <- list(dat$X, sp = dat$sp, covar = diag(nspp))</pre>
# random slope with species showing phylogenetic covariances
re.4 <- list(dat$X, sp = dat$sp, covar = Vphy)</pre>
# random effect for site
re.site <- list(1, site = dat$site, covar = diag(nsite))
simple <- communityPGLMM(Y ~ X, data = dat, family = "binomial", sp</pre>
= dat$sp, site = dat$site, random.effects = list(re.site),
REML=TRUE, verbose=FALSE)
# The rest of these tests are not run to save CRAN server time;
# - please take a look at them because they're *very* useful!
## Not run:
```

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```
z.binary <- communityPGLMM(Y \sim X, data = dat, family = "binomial",
sp = dat$sp, site = dat$site, random.effects = list(re.1, re.2,
re.3, re.4), REML = TRUE, verbose = FALSE)
# output results
z.binary
plot(z.binary)
# test statistical significance of the phylogenetic random effect
# on species slopes using a likelihood ratio test
communityPGLMM.binary.LRT(z.binary, re.number = 4)$Pr
# extract the predicted values of Y
communityPGLMM.predicted.values(z.binary, show.plot = TRUE)
# examine the structure of the overall covariance matrix
communityPGLMM.matrix.structure(Y ~ X, data = dat, family =
"binomial", sp = dat$sp, site = dat$site, random.effects =
list(re.1, re.2, re.3, re.4))
# look at the structure of re.1
communityPGLMM.matrix.structure(Y ~ X, data = dat, family =
"binomial", sp = dat$sp, site = dat$site, random.effects =
list(re.1))
# compare results to glmer() when the model contains no
# phylogenetic covariance among species; the results should be
communityPGLMM(Y \sim X, data = dat, family = "binomial", sp = datsp,
site = dat$site, random.effects = list(re.1, re.3), REML = FALSE,
verbose = FALSE)
# 1mer
if(require(lme4)){
summary(glmer(Y \sim X + (1 \mid sp) + (0 + X \mid sp), data=dat, family =
"binomial"))
# compare results to lmer() when the model contains no phylogenetic
# covariance among species; the results should be similar.
communityPGLMM(Y ~ X, data = dat, family = "gaussian", sp = dat$sp,
site = dat$site, random.effects = list(re.1, re.3), REML = FALSE,
verbose = FALSE)
# 1mer
summary(lmer(Y \sim X + (1 | sp) + (0 + X | sp), data=dat, REML = FALSE))
## End(Not run)
```

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Description

Build a novel phylogeny from existing data

congeneric.impute sequentially add species to a phylogeny to form an _imputed_ bifurcating tree. Makes use of a result from Steel & Mooers (2010) that gives the expected branch-length under a Yule model whether the rate of diversification has been estimated. The intention of this is to approximate the method by which phylogenetic structure is sampled from the prior in BEAST; i.e., to approximate the standard Kuhn et al. (2011) method for imputing a phylogeny. When using congeneric.impute you should (1) repeat your analyses across many (if in doubt, thousands) of separate runs (see Kuhn et al. 2011) and (2) check for yourself that your trees are unbiased for your purpose - I make no guarantee this is appropriate, and in many cases I think it would not be. See also 'notes' below.

Usage

```
congeneric.merge(tree, species, split = "_", ...)
bind.replace(backbone, donor, replacing.tip.label, donor.length = NA)
congeneric.impute(tree, species, split = "_", max.iter = 1000, ...)
```

Arguments

tree	phylo phylogeny to have those species inserted into it	
species	vector of species names to be bound into the tree if missing from it	
split	the character that splits genus and species names in your phylogeny. Default is _, i.e. Quercus_robur.	
	ignored	
backbone	backbone phylogeny (phylo) into which the donor is to be bound	
donor	phylogeny (phylo) to bound into the backbone phylogeny	
replacing.tip.label		
	the species in the donor phylogeny that's being replaced by the donor phylogeny	
donor.length	how deep the donor phylogeny should be cut into the backbone phylogeny. If NA (default), then the bladj algorithm is followed (or, in plain English, it's put half-way along the branch)	
max.iter	Sometimes the random draw for the new branch length to be added will be too large to allow it to be added to the tree. In such cases, congeneric.imput will randomly draw another branch length, and it will repeat this process max.iter times. See 'notes' for more on this.	

Details

congeneric.merge Binds missing species into a phylogeny by replacing all members of the clade it belongs to with a polytomy. Assumes the tip.labels represent Latin binomials, split by the split argument. This code was originally shipped with phyloGenerator - this is the merge method in that program.

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bind.replace Binds a phylogeny (donor) into a bigger phylogeny ('backbone'); useful if you're building a phylogeny a la Phylomatic. A version of this R code was shipped with phyloGenerator (Pearse & Purvis 2013). This is really an internal function for congeneric.merge, but hopefully it's of some use to you!

Value

```
phylo phylogeny
phylogeny (phylo)
```

Note

Thank you to Josep Padulles Cubino, who found that the genus name splitting in a previous version of this function could cause incorrect placement in oddly named cases. As with all phylogenetic construction tools, the output from congeneric merge should be checked for accuracy before use.

Caveats for congeneric.impute: something I noticed is that BEAST randomly picks an edge to break when adding species (starting from a null tree), and this is the behaviour I have (attempted to) replicate here. It is not clear to me that this is unbiased, since a clade undergoing rapid diversification will have many edges but these will be short (and so cannot have an edge inserted into them following the method below). My understanding is this is a known problem, and I simply cannot think of a better way of doing this that doesn't incorporate what I consider to be worse pathology. Thus this method, even if it works (which I can't guarantee), it should tend to break long branches.

Author(s)

Will Pearse

Will Pearse

References

Pearse W.D. & Purvis A. phyloGenerator: an automated phylogeny generation tool for ecologists. Methods in Ecology and Evolution 4(7): 692–698.

Steel, M., & Mooers, A. (2010). The expected length of pendant and interior edges of a Yule tree. Applied Mathematics Letters, 23(11), 1315-1319.

Kuhn, T. S., Mooers, A. O., & Thomas, G. H. (2011). A simple polytomy resolver for dated phylogenies. Methods in Ecology and Evolution, 2(5), 427-436.

Examples

```
tree <- read.tree(text="((a_a:1,b_b:1):1, c_c:2):1;")
tree <- congeneric.merge(tree, c("a_nother", "a_gain", "b_sharp"))
tree <- read.tree(text="((a_a:1,b_b:1):1, c_c:2):1;")
tree <- congeneric.impute(tree, c("a_nother", "a_gain", "b_sharp"))</pre>
```

phy.signal 53

phy.signal

Calculate phylogenetic 'signal'

Description

Calculate phylogenetic 'signal'

Usage

```
phy.signal(data, method = c("lambda", "delta", "kappa", "blom.k"))
```

Arguments

data comparative.comm object what kind of signal to calculate, one of Pagel's λ (default), δ , and κ

, or Blomberg's K.

Details

Phylogenetic 'signal' is one of those concepts that is said a lot in community ecology, but some do not full consider its meaning. Think carefully before rushing to report a value whether: (1) it makes sense to assess phylogenetic 'signal' in your datasets, and (2) what the phrase 'phylogenetic signal' actually means. This code makes use of caper::pgls to get estimates of fit; alternatives that offer more flexibility exist (see below).

Value

Named numeric vector, where each element is a trait or community.

Author(s)

Will Pearse, Jeannine Cavender-Bares

References

Blomberg S.P., Garland T. & Ives A.R. Testing for phylogenetic signal in comparative data: behavioral traits are more labile. Evolution 57(4): 717–745.

R. P. Freckleton, P. H. Harvey, and M. Pagel. Phylogenetic analysis and comparative data: A test and review of evidence. American Naturalist, 160:712-726, 2002.

Mark Pagel (1999) Inferring the historical patterns of biological evolution. Nature 6756(401): 877–884.

See Also

fitContinuous fitDiscrete pgls phylosignal

Examples

```
data(laja)
data <- comparative.comm(invert.tree, river.sites, invert.traits)
phy.signal(data, "lambda")</pre>
```

Description

Dot-plots of community presence/absence or abundance

Usage

```
## S3 method for class 'comparative.comm'
plot(
    x,
    sites = NULL,
    abundance = FALSE,
    pch = 20,
    dot.cex = NULL,
    site.col = "black",
    fraction = 3,
    x.increment = NULL,
    show.tip.label = FALSE,
    ...
)
```

Arguments

```
comparative.comm object
Х
                  names of sites to plot (default: all); see examples
sites
                  make size proportional to species abundance (default: FALSE)
abundance
                  plotting character to be used for sites (see pch)
pch
dot.cex
                  function to determine point size; see examples, this isn't as terrible-sounding as
                  it seems.
site.col
                  colours to use when plotting sites; if not same length as number of sites, only
                  the first element is used (no recycling)
fraction
                  fraction of plot window to be taken up with phylogeny; e.g., 3 (default) means
                  phylogeny is 1/3 of plot
x.increment
                  specify exact spacing of points along plot; see examples
show.tip.label whether to plot species names on phylogeney (default: FALSE)
                  additional arguments passed to plotting functions
```

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Details

Take a look at the examples: this is (hopefully!) a lot more straightforward than it might seem. Getting the right spacing of dots on the phylogeny may take some playing around with the fraction and x.increment arguments. It may seem a little strange to set point size using a function, however, this gives you much more flexibility and the ability to (usefully) transform your data.

Value

List containing plot.phylo information, as well as the used x.adj values (compare with your x.increment)

Author(s)

Will Pearse, Matt Helmus

See Also

```
comparative.comm traitgram.cc
```

Examples

```
data(laja)
data <- comparative.comm(invert.tree, river.sites, invert.traits)
plot(data)
plot(data, sites=c("AT", "BP"), fraction=1.5)
settings <- plot(data, sites=c("AT", "BP"), site.col=rainbow(2), fraction=1.5)
plot(data, sites=c("AT", "BP"), site.col=rainbow(2),
fraction=1.2, x.increment=settings$x.increment/4)
#dot.cex isn't as scary as it sounds...
plot(data, site.col=rainbow(2), fraction=1.2, abundance=TRUE, dot.cex=sqrt)
#...or other trivial variants...
abund.sqrt <- function(x) ifelse(x>0, sqrt(x), 0)
plot(data, sites=c("AT", "BP"), site.col=rainbow(2), fraction=1.2,
x.increment=settings$x.increment/4, abundance=TRUE, dot.cex=abund.sqrt)
plot(data, sites=c("AT", "BP"), site.col=rainbow(2), fraction=1.2,
x.increment=settings$x.increment/4, abundance=TRUE, dot.cex=function(x) sqrt(x))
```

scape

Simulate phylogenetic community structure across a landscape

Description

scape simulates communities that are phylogenetically structured

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Usage

```
scape(
  tree,
  scape.size = 10,
 g.center = 1,
 g.range = 1,
 g.repulse = 1,
 wd.all = 150,
 signal.center = TRUE,
  signal.range = TRUE,
  same.range = TRUE,
  repulse = TRUE,
  center.scale = 1,
  range.scale = 1,
  repulse.scale = 1,
  site.stoch.scale = 0.5,
  sd.center = 1,
  sd.range = 1,
 rho = NULL,
  th = 8
)
```

Arguments

tree	phylo object
scape.size	edge dimension of square landscape
g.center	strength of phylogenetic signal in species range centers
g.range	strength of phylogenetic signal in species range sizes
g.repulse	strength of phylogenetic repulsion
wd.all	niche width, larger values simulate broader range sizes
signal.center	simulate with phylosignal in range centers
signal.range	simulate with phylosignal in range size
same.range	make all range sizes equal
repulse	include phylogenetic repulsion in range centers
center.scale	adjust strength of phylogenetic attraction in range centers independent of g. center
range.scale	adjust strength of phylogenetic signal in range size independent of g.range
repulse.scale	adjust strength of phylogenetic repulsion independent of g.repulse
site.stoch.scal	e
	adjust strength of random variation in species richness across sites
sd.center	sd in rnorm for the range centers, increase to get more variation in center values across species
sd.range	sd rnorm for the range sizes, increase to get more variation in range sizes across gradients
rho	Grafen branch adjustment of phylogenetic tree see corGrafen
th	probability threshold 10^-th above which species are considered present at a site

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Details

Simulates a landscape with species (i.e., tree tips) distributions dependent on a supplied phylogenetic tree. The amount and type of structure is determened by the signal parameters g.center, g.range and g.repulse. Parameters are based on an Ornstein-Uhlenbeck model of evolution with stabilizing selection. Values of g=1 indicate no stabilizing selection and correspond to the Brownian motion model of evolution; 0<g<1 represents stabilizing selection; and g>1 corresponds to disruptive selection where phylogenetic signal for the supplied tree is amplified. See corBlomberg. Communities are simulated along two gradients where the positions along those gradients, g.center and range sizes g.range, can exhibit phylogenetic signal. Phylogenetic attraction is simulated in the g.center parameter, while repulsion in g.repulse. Both can be exhibited such that closly related species are generally found with similar range centers (phylogenetic attraction) but just not at the same site (phylogenetic repulsion). The function then returns probabilities of of each species across sites and the presence and absences of species based a supplied threshold, th, which can be increased to obtain more species at sites and thus increase average site species richness.

Value

cc	comparative.comm object with presence/absence results of simulations. The site names are the row.columns of the cells in the original grid cells that made up the data, and these co-ordinates are also given in the env slot of the object.
X.joint	full probabilities of species at sites, used to construct cc
X1	probabilities of species along gradient 1
X2	probabilities of species along gradient 2
sppXs	full probabilities of each species as an array arranged in a $scape.size-by-scape.size$ matrix
V.phylo	initial phylogenetic covariance matrix from tree
V.phylo.rho	phylogenetic covariance matrix from tree scaled by Grafen if rho is provided
V.center	scaled by g. center phylo covariance matrix used in the simulations
V.range	scaled by g.range phylo covariance matrix used in the simulations
V.repulse	scaled by g.repulse phylo covariance matrix used in the simulations
bspp1	species optima for gradient 1
bspp2	species optima for gradient 2
u	the env gradients values for the two gradients
wd	the denominator for species ranges

Author(s)

M.R. Helmus, cosmetic changes by Will Pearse

References

Helmus M.R. & Ives A.R. (2012). Phylogenetic diversity area curves. Ecology, 93, S31-S43.

See Also

```
eco.scape sim.phy sim.meta
```

58 sim.meta

Examples

```
#Create balanced tree with equal branch-lengths (signal in centers)
tree <- stree(8,type="balanced")</pre>
tree$edge.length <- rep(1, nrow(tree$edge))</pre>
tree$root <- 1
kk <- scape(tree, scape.size=100, g.center=100, g.range=1, g.repulse=1, wd.all=150,
  signal.center=TRUE, signal.range=FALSE, same.range=FALSE, repulse=FALSE,center.scale = 1,
    range.scale = 1, repulse.scale = 1, site.stoch.scale = 0, sd.center=3, sd.range=1,
    rho=NULL, th=20)
#Make some plots
par(mfrow=c(1,Ntip(tree)),mar=c(.1,.1,.1,.1))
for(j in seq_along(tree$tip.label))
    image(t(1 - kk\$sppXs[,,j]/max(kk\$sppXs[,,j])), xlab = "",
              ylab = "",main = "",axes=FALSE, col=grey.colors(10))
par(mfrow=c(2,1))
matplot((kk$X1), type = "l", xlab="gradient",ylab = "probability",
main = "Gradient 1",col=rainbow(dim(kk$X1)[2]),lty=1)
matplot((kk$X2), type = "1", xlab="gradient",ylab = "probability",
main = "Gradient 2",col=rainbow(dim(kk$X2)[2]),lty=1)
plot(x=seq_along(sites(kk$cc)),y = rowSums(comm(kk$cc)), main = "SR",type = "1")
cor(kk$X1)
```

sim.meta

Simulate a meta-community (and its phylogeny)

Description

sim.meta.comm simulates species moving through a metacommunity. At each time-step each cell's next abundance for each species is env.quality - current.abundance + stochastic, and a species gets as many chances to migrate in each time-step as it has cells (the same cell could migrate multiple times). I use a Poisson for everything because I don't want half-species (these are individuals), and keeping everything in Poisson makes it easier to compare the relative rates of everything.

Usage

```
sim.meta.comm(
    size = 10,
    n.spp = 8,
    timesteps = 10,
    p.migrate = 0.05,
    env.lam = 10,
    abund.lam = 5,
    stoch.lam = 1
)
```

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```
sim.meta.phy.comm(
    size = 10,
    n.spp = 8,
    timesteps = 10,
    p.migrate = 0.3,
    env.lam = 10,
    abund.lam = 5,
    stoch.lam = 1,
    p.speciate = 0.05
)
```

Arguments

size the length and width of the meta-community in grid cells

n.spp number of species

timesteps number of time-steps (each discrete)

p.migrate probability that a group of species in each grid cell will migrate to another grid

cell each timestep (i.e., 10 cells occuped by a species -> 10*p.migrate chance

of migration)

env.lam

 λ

value for Poisson distribution used to distribute environmental quality; essentially the carrying capacity (for each species separately) for that cell

abund.lam

 λ

value for Poisson distribution used to distribute initial abundances and abundance after migration

stoch.lam

 λ

value for Poisson distribution of noise added to the next step abundance calculation. With equal chance, this is taken as either a positive or a negative number (see details if you're confused as to why this is Poisson!)

p.speciate

probabilty that, at each timestep, a species will speciate. A species can only speciate, migrate, or reproduce if it has individuals!

Details

sim.meta.phy.comm As above, but with a simulation of phylogeny as well - there are no additional extinction parameters, since extinction happens as a natural consequence of ecological interactions.

sim.phy

Value

For sim.meta.comm a list with a species-site matrix as the first slot, and the environment as the second. Rownames of the site-species are the List with the x and y co-ordinates of the simulation grid pasted together; colnames are arbitrary species names. sim.meta.comm, a comparative.comm object (since we have now simulated a phylogeny), with the same naming convention for the site names. phylogeny.

sim.meta.phy.comm comparative.comm object that describes the data; note that the rownames of the community object refer to the row.column of the data in the simulated grid assemblages.

Note

scape is a much more sophisticated simulation of the biogeography, but requires you to supply a phylogeny. You pays your money, you makes your choice.

Author(s)

Will Pearse

Will Pearse

See Also

sim.phy scape

sim.phy

Simulate phylogenies

Description

Simulate phylogenies under pure birth/death or as a function of trait evolution

Usage

```
sim.bd.phy(speciate = 0.1, extinction = 0.025, time.steps = 20)

sim.bd.tr.phy(
    speciate = 0.1,
    extinction = 0.025,
    time.steps = 20,
    tr.range = c(0, 1),
    sp.tr = 2,
    ext.tr = 1,
    tr.walk = 0.2,
    tr.wrap = TRUE
)

edge2phylo(edge, s, e = numeric(0), el = NA, t = NULL)
```

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Arguments

speciate	probability each species will speciate in each time-step (0-1)
extinction	probability each species will go extinct in each time-step (0-1)
time.steps	number of time-steps for simulation
tr.range	vector of length two specifying boundaries for trait values (see notes); initial two species will be at the 25th and 75th percentiles of this space. See also tr.wrap
sp.tr	speciation rate's interaction with the minimum distance between a species and the species most similar to it (see details)
ext.tr	extinction rate's interaction with the minimum distance between a species and the species most similar to it (see details)
tr.walk	at each time-step a species not undergoing speciation or extinction has its trait value drawn from a distribution centered at its current value and with a standard deviation set by this value. I.e., this is the rate of the Brownian motion trait evolution.
tr.wrap	whether to force species' trait values to stay within the boundary defined by tr.range; default TRUE.
edge	a two-column matrix where the first column is the start node, the second the destination, as in phylo\$edge
S	which of the rows in the edge matrix represent extant species
е	which of the tips in the edge matrix are extinct (DEFAULT: empty vector, i.e., none)
el	a vector to be used to give edge.length to the phylogeny (default NA, i.e., none)
t	if given (default NA), a vector to be used for traits ($traits slot$) in the phylogeny

Details

sim.bd.tree simulates a pure birth/death speciation model. There are two important things to note: (1) speciation is randomised before extinction, and only one thing can happen to a lineage per timestep. (2) This code works well for my purposes, but absurd parameter values can cause the function to crash.

sim.bd.tr.tree is an extension of sim.bd.tree, and all its caveats apply to it. It additionally simulated the evolution of a trait under Brownain motion (tr.walk). Species' speciation/extinction rates change depending on whether they have a trait value similar to other species (sp.tr, ext.tr). When a speciation event happens, the two daughters split evenly about the ancestor's trait value, taking values half-way to whatever the nearest species' value is. To be precise:

```
p(speciate)_i = speciate_i + sp.tr \times min(traitdistance) , p(extinct)_i = exinction_i + ext.tr \times min(traitdistance) , where i
```

denotes each species.

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edge2phylo is an internal function for the sim. phy and sim. meta function families, which may be of use to you. Check those functions' code for examples of use.

These functions are closely related to sim.meta; the latter are extensions that simulate meta-community structure at the same time.

Value

phylo object with random tip.labels, and trait values if using sim.br.tr.tree.

Author(s)

Will Pearse

Will Pearse

Will Pearse

See Also

sim.meta scape

Examples

```
tree <- sim.bd.phy(0.1, 0, 10)
plot(tree)</pre>
```

trait.asm

Produces simulated communities based on species attributes

Description

trait.asm calculates phylogenetic biodiversity metrics

Usage

```
trait.asm(
   a,
   m = 1000,
   meanSR = NULL,
   interval = c(0.001, 10),
   exponential = TRUE,
   Pscale = 1
)
```

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Arguments

a species attributes (e.g., traits like body size)
m number of communities to be simulated

meanSR target mean species richness across simulated communities

interval adjust to obtain meanSR

exponential use the exponential distribution when simulating communities

Pscale adjust this value when not using the exponential distribution in order to scale the

species richnesses in the simulated communities

Details

Simulates a set of communities based on the supplied attribute (trait) where larger values make it more likely for species to be in the communities.

Value

```
Y presence/absence matrix
P probabilities
a the supplied trait
exponential if the exponential distribution was used
meanSR supplied meanSR value
std estimated sd
```

Author(s)

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References

Helmus M., Mercado-Silva N. & Vander Zanden M.J. (2013). Subsidies to predators, apparent competition and the phylogenetic structure of prey communities. Oecologia, 173, 997-1007.

Examples

```
## Not run:
  data(laja)
  trait.asm(laja$fish.pref)
## End(Not run)
```

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traitgram.cc

Traitgram for comparative community object

Description

traitgram.cc A wrapper for the traitgram function in the picante package. princompOne A very soft wrapper for princomp

Usage

```
traitgram.cc(object, trait, moreArgs = NULL, ...)
princompOne(x, ...)
```

Arguments

object A comparative.comm object.

trait Which trait to plot. If missing, use the first trait. If a positive numeric vector

of length one, use the as.integer(trait)th trait. If a numeric vector, use it instead of the trait data in object. If a character vector of length one, use the trait with that name. If a function pass the trait data frame through that function and use the result (princompOne is a wrapper). If an expression, evaluate that expression in the environment of the trait data and use the result. If a character vector, then convert to an expression and evaluate in the environment of the trait

data and use the result.

moreArgs List of more arguments to pass on to trait (if its a function).

... Additional arguments to be passed on to traitgram or prcomp for traitgram.cc

and princomOne respectively.

x A matrix-like object

Value

```
traitgram.cc: see traitgram princompOne: the first axis of a PCA
```

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