Package 'specificity'

June 19, 2020

Version 0.0.0.9000		
Description The purpose of this package is to calculate phylogenetic and environmental speci-		
ficity of species. I wrote this software to analyze specificity of microbes to hosts or to environ-		

ment, but there is no reason that this software wouldn't work with macroorganisms as well.

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Title Calculate Environmental or Host Phylogenetic Specificity

R topics documented:

bl_distance_ns	2
check_pes_inputs	3
daughter_seeds	2
distcalc	2
env_spec_sim	4
geo_spec_sim	7
make_nested_set	
occ_threshold	1(
pairwise_product	11
phy_or_env_spec	12
phy_spec_sim	15
plot_grid_abunds	17
plot_pairwise_spec	18
plot_specificities	19

2 bl_distance_ns

	prop_abund	20
	randomgrid	21
	random_rep_positions	22
	rao_genetic_max	23
	rao_sort_max	24
	spec_core	25
	tips_from_node	25
	tree2mat	26
	wpd	27
	wpd_table	29
Index		31

bl_distance_ns

 $bl_distance_ns$

Description

Calculates branch-length distance between tipa and tipb in a phylogenetic tree using nested-set optomization. Requires a pre-calculated nested-set.

Usage

```
bl_distance_ns(tipa, tipb, tree, ns)
```

Arguments

tipa string. Name of a tip in tree.

tipb string. Name of another tip in tree.

tree phylo object. Tree containing all unique species in x as tips. May contain tips

that are not in x.

ns matrix. Nested-set matrix for tree; use make_nested_set(tree).

Value

Distance between tipa and tipb.

Author(s)

John L. Darcy

check_pes_inputs 3

Examples

```
library(ape)
example_tree <- ape::read.tree(text=" ((((a:1,b:1):1,c:2):1,d:3):1,(e:1,f:1):3);")
plot(example_tree); axis(side=1)
example_ns <- make_nested_set(example_tree)
bl_distance_ns("a", "c", example_tree, example_ns) # should be 4
bl_distance_ns("a", "f", example_tree, example_ns) # should be 8
bl_distance_ns("d", "c", example_tree, example_ns) # should be 6</pre>
```

check_pes_inputs

check pes inputs

Description

Function used by phy_or_env_spec. checks abunds_mat, env, hosts, and hosts_phylo inputs to phy_or_env_spec to make sure there are no problems. This could include missing species in trees, incompatible dimensions, non-numeric inputs, etc. Returns an input type, which is just a string that can be "mat", "dist", "vec", "phy", or "error".

Usage

```
check_pes_inputs(abunds_mat, env, hosts, hosts_phylo, verbose = TRUE)
```

Arguments

```
abunds_mat (required, see phy_or_env_spec)

env (required, can be NULL, see phy_or_env_spec)

hosts (required, can be NULL, see phy_or_env_spec)

hosts_phylo (required, can be NULL, see phy_or_env_spec)

verbose logical. Should status messages be displayed? (DEFAULT: TRUE).
```

Value

```
string. either "mat", "dist", "vec", "phy", or "error".
```

4 disteale

daughter_seeds

daughter_seeds

Description

Makes n daughter seeds from seed s. This is useful for processes one wishes to be deterministic, but may not be executed in the same order every time.

Usage

```
daughter_seeds(n, s = 12345)
```

Arguments

n integer. Number of daughter seeds to make.

s integer. A seed (DEFAULT: 12345).

Value

vector of length n containing integer seeds.

Author(s)

John L. Darcy

distcalc

distcalc

Description

Calculates pairwise geographic distance between locations on earth. Just a convenient wrapper for fields::rdist.earth().

Usage

```
distcalc(lat, lng, sampIDs = NULL)
```

Arguments

Numeric vector. Latitudes in decimal degree format.Numeric vector. Longitudes in decimal degree format.

sampIDs Character vector. Sample identifiers. Only required if output dist should have

names associated.

env_spec_sim 5

Value

matrix containing all pairwise geographic distances in km.

Author(s)

John L. Darcy

Examples

```
data(endophyte)
geo_dists <- distcalc(metadata$Lat, metadata$Lon, metadata$SampleID)
all(rownames(geo_dists) == metadata$SampleID)</pre>
```

env_spec_sim

env_spec_sim

Description

Simulates inputs for phy_or_env_spec, by creating a species distribution over an artificial (or real) environmental variable. That distribution has a mean at the "ideal" environmental value for the simulated species, and the standard deviation of that distribution controls the extent to which the species is specific to the variable. A high SD means less specificity, and a low SD means more specificity.

Usage

```
env_spec_sim(
    sdev,
    ideal,
    ideal2 = 0,
    ideal3 = 0,
    n_ideal = 1,
    env,
    n_obs,
    up = 0,
    oceanp = 0,
    n_cores = 2,
    seed = 1234567
)
```

Arguments

sdev

numeric vector. Standard deviation of the probability distribution P(species), in the same units of env. Low values mean that the species is found acrosss only a narrow range of env, i.e. specificity. High values mean that the species is found across a wide range of env, i.e. cosmopolitanism. Multiple values can be input 6 env_spec_sim

	in order to simulate a range of specificities simultaneously. Can be length 1 or n .
ideal	numeric vector. Value of env that is ideal for the simulated species. This is the mode of the probability distribution P(species). Can be length 1 or n.
ideal2	numeric vector. Value of env that is the second ideal for the simulated species. Only used if $n_{ideal} >= 2$. This is the second mode of the probability distribution $P(species)$. Can be length 1 or n .
ideal3	numeric vector. Value of env that is the third ideal for the simulated species. Only used if $n_{ideal} = 3$. This is the third mode of the probability distribution P(species). Can be length 1 or n.
env	numeric vector. Real or fake environmental variable.
n_obs	integer vector. Number of positive observations to make, i.e. occupancy of simulated species. Can be length 1 or n (default: 1).
up	numeric vector. up=uniform proportion. This is the proportion of the probability distribution P(species) that is composed of a uniform distribution, if desired. If set to a value above zero (and blow 1), P(species) will be a weighted sum of the normal distribution described above, and a uniform distribution. The weight for the uniform distribution will be up, and the weight for the normal distribution will be 1-up (default: 0).
oceanp	numeric vector. oceanp=ocean proportion. This is the proportion of samples in env that are "in the ocean", i.e. samples where the species would not expect to be found even if env is permissive. If aliens were calculating specificity of cows to temperature, they might look in the ocean at sites where the temperature is 17C (great for cows). But cows are not found in the ocean. This proportion is used to randomly select ocean sites within env, and then p(slenvlocean) = up. Can be length 1 or n (default: 0).
n_cores	integer. Number of CPU cores for parallel computation (DEFAULT: 2).
seed	integer. Seed for randomization. Daughter seeds will be generated for parallel computations, each with the same number of digits as seed (DEFAULT: 1234567).

Details

Since this process can result in failures (if a species is requested that's highly specific to a region of env that isn't samples), some output species will be failures. Default operation is to remove those failures from output matrix and output params data frame, but this can be changed.

Value

List object containing "matrix" and "params" objects:

matrix: matrix where each column is a vector of simulated observation frequencies (counts) corresponding to a value of env; each row represents a simulated species.

params: data.frame of parameters (columns) used to simulate each species (rows).

geo_spec_sim 7

Author(s)

John L. Darcy

Examples

none yet written.

geo_spec_sim

geo_spec_sim

Description

Simulates inputs for phy_or_env_spec, by creating a species distribution over artificial (or real) geographic space. That distribution has a bivariate mean at the "ideal" location inspace for the simulated species, and the standard deviation of that (normal) distribution controls the extent to which the species specific to geographic space. A high SD means less specificity, and a low SD means more specificity.

Usage

```
geo_spec_sim(
    sdev,
    n_obs,
    grid,
    ideal_x = 0,
    ideal_y = 0,
    ideal_y2 = 0,
    ideal_y3 = 0,
    ideal_y3 = 0,
    n_ideal = 1,
    up = 0,
    seed = 123456,
    n_cores = 2
)
```

Arguments

sdev

numeric vector. Standard deviation of the probability distribution P(species), in the same units as grid. P(species) is a function of the distance between a sample site and its closest ideal location (specified with ideal_x/2/3 and ideal_y/2/3). Low values mean that the species is found in abundance within only short distances of ideal locations, high values mean the species is found across a wider area. Multiple values can be input in order to simulate a range of specificities simultaneously. Can be length 1 or n.

geo_spec_sim

n_obs	integer vector. Number of observations to make, i.e. number of times species is observed. Will be the sum of the species' output column. Can be length 1 or n.
grid	data frame with columns x and y, representing cartesian coordinates of sample locations. Can be artificial (generate with randomgrid()) or real.
ideal_x	numeric vector. x-coordinate of the ideal spatial location for species (DEFAULT=0).
ideal_y	numeric vector. y-coordinate of the ideal spatial location for species (DEFAULT=0).
ideal_x2	numeric vector. x-coordinate for secondary ideal location. Only used if n_ideal<1 (DEFAULT=0).
ideal_y2	numeric vector. y-coordinate for secondary ideal location. Only used if n_ideal<1 (DEFAULT=0).
ideal_x3	numeric vector. x-coordinate for secondary ideal location. Only used if n_ideal<2 (DEFAULT=0).
n_ideal	integer vector. number of ideal locations to use. Must be 1, 2, or 3 (DE-FAULT=1).
up	numeric vector. up=uniform proportion. This is the proportion of the probability distribution P(species) that is composed of a uniform distribution, if desired. If set to a value above zero (and blow 1), P(species) will be a weighted sum of the normal distribution described above, and a uniform distribution. The weight for the uniform distribution will be up, and the weight for the normal distribution will be 1-up (default: 0).
seed	integer. Seed for randomization. Daughter seeds will be generated for parallel computations, each with the same number of digits as seed (DEFAULT: 1234567).
n_cores	integer. Number of CPU cores for parallel computation (DEFAULT: 2).

Value

List object containing "matrix" and "params" objects:

matrix matrix where each column is a vector of simulated observations for each row in grid; each column of matrix represents a simulated species.

params data.frame of parameters (columns) used to simulate each species (rows).

Author(s)

John L. Darcy

Examples

```
g1 <- randomgrid()
plot(g1)
a1 <- geo_spec_sim(sdev=c(30, 30, 30, 30), n_obs=1000, grid=g1, up=c(0, 0.20, 0.40, 0.60))
par(mfrow=c(2,2))
plot_grid_abunds(g1, a1$matrix[,1])
plot_grid_abunds(g1, a1$matrix[,2])
plot_grid_abunds(g1, a1$matrix[,3])
plot_grid_abunds(g1, a1$matrix[,4])</pre>
```

make_nested_set 9

```
a2 <- geo_spec_sim(sdev=c(10, 20, 30, 40), n_obs=1000, grid=g1, ideal_x=-50, ideal_x2=50, n_ideal=2)
par(mfrow=c(2,2))
plot_grid_abunds(g1, a2$matrix[,1], main="sd=10")
plot_grid_abunds(g1, a2$matrix[,2], main="sd=20")
plot_grid_abunds(g1, a2$matrix[,3], main="sd=30")
plot_grid_abunds(g1, a2$matrix[,3], main="sd=30")
plot_grid_abunds(g1, a2$matrix[,4], main="sd=40")</pre>
```

make_nested_set

make_nested_set

Description

Makes a nested set table for a phylo object. Phylo objects made by the ape package store phylogenies as an "adjacency list", which in R is a table within which any given edge is represented by the two node numbers it connects. With this data structure, it is very computationally expensive to figure out which tips are the descendents of a given node. Instead, using a "nested set" data structure, this operation is trivial. A nested set stores the minimum and maximum tip index for each node, such that the descendents of that node are given by the inclusive range between those values.

Usage

```
make_nested_set(phy, n_cores = 2)
```

Arguments

phy phylo object. Must be rooted, and sorted such that tip indices are ordered. This

is the default for rooted trees read in using ape's read.tree function.

n_cores integer. Number of CPU cores to use (DEFAULT: 2). lapply will be used instead

of mclapply if ncores is 1.

Value

Matrix object representing a nested set of nodes. Each row matches rows of the "edges" object within phy. Object has the following columns:

1 (node) Node value in the original phylo object.

2 (min) minimum tip index subtended by node.

3 (max) maximum tip index subtended by node.

4 (contig) Is min:max congiguous? 1 (true) or 0 (false).

Author(s)

John L. Darcy

References

https://en.wikipedia.org/wiki/Nested_set_model https://en.wikipedia.org/wiki/Adjacency_list

10 occ_threshold

See Also

```
ape::phylo
```

Examples

```
library(specificity)
phy <- get(data(endophyte))$supertree
# check if tree is rooted:
is.rooted(phy)
# make nested set table:
phy_ns <- make_nested_set(phy)
# show that nested set table matches up with edges table in phy:
all(phy$edge[,2] == phy_ns[,1])</pre>
```

occ_threshold

occ_threshold

Description

removes species (columns) from a matrix that don't meet a minimum occupancy, defined as the number of samples in which that species was observed.

Usage

```
occ_threshold(m, threshold, max_absent = 0)
```

Arguments

m matrix or data frame of numeric values. Columns represent species, rows are

samples.

threshold integer. Minimum number of samples a species can occupy without being re-

moved.

max_absent float. Maximum abundance value at which a species will be considered absent

(DEFAULT: 0).

Value

matrix with low-occupancy species removed.

Author(s)

John L. Darcy

pairwise_product 11

Examples

```
attach(endophyte)
dim(zotutable)
zotutable_over25 <- occ_threshold(zotutable, 25)
dim(zotutable_over25)</pre>
```

pairwise_product

pairwise_product

Description

Calculates pairwise_products from unique 2-element combinations of vector x. The output vector is the same length and same order as a lower triangle of matrix with rows and columns x.

Usage

```
pairwise_product(x)
```

Arguments

Χ

numeric vector.

Value

vector of pairwise_products, of length $(1^2-1)/2$, where l=length(x).

Author(s)

John L. Darcy

Examples

```
x <- 1:6
y_cpp <- pairwise_geo_mean(x)
y_r <- as.dist(outer(x, x, function(x,y){x*y}))
print("Calculated with R's outer() function:")
y_r
print("As a vector:")
as.vector(y_r)
print("Calculated with pairwise_product (C++):")
y_cpp</pre>
```

phy_or_env_spec

phy_or_env_spec phy_or_env_spec

Description

Calculates species' specificities to either a 1-dimensional variable (vector), 2-dimensional variable (matrix), or to a phylogeny. Transforms all variable input types into a matrix D, and calculates specificity by comparing empirical Rao's Quadratic Entropy to simulated RQE (same but with permuted abundances). By default (denom_type = "index"), an index is calculated from emp and sim values such that Spec=0 indicates random assortment (null hypothesis), and more negative values indicate stronger specificity.

Usage

```
phy_or_env_spec(
  abunds_mat,
  env = NULL,
  hosts = NULL,
  hosts_phylo = NULL,
  n_{sim} = 1000,
                              m[sample(1:nrow(m)), ] },
  sim_fun = function(m) {
 p_adj = "fdr",
  seed = 1234567,
  tails = 1,
  n_{cores} = 2,
  verbose = TRUE,
  p_method = "raw",
  center = "mean",
  denom_type = "index",
  diagnostic = F
)
```

Arguments

abunds_mat	matrix or data frame of numeric values. Columns represent species, rows are samples. For columns where the value is nonzero for two or fewer data points, environmental SES cannot be calculated, and NAs will be returned. Negative values in abunds_mat are not allowed (REQUIRED).
env	numeric vector, dist, or square matrix. Environmental variable corresponding to abunds. For example, temperature, or geographic distance. Not required for computing phylogenetic specificity (DEFAULT: NULL).
hosts	character vector. Host identities corresponding to abunds. Only required if calculating SES for phylogenetic specificity (DEFAULT: NULL).
hosts_phylo	phylo object. Tree containing all unique hosts as tips. Only required if calculating SES for phylogenetic specificity (DEFAULT: NULL).

phy_or_env_spec 13

n_sim

integer. Number of simulations of abunds_mat to do under the null hypothesis that host or environmental association is random. P-values will not be calculated if $n \le 100$ (DEFAULT: 500).

sim fun

function. A function f where f(abunds_mat) returns a matrix object with the same number of rows and columns as abunds_mat. Default is f=function(m) m[sample(1:nrow(m)),], which just permutes the order of rows in abunds_mat. Users may wish to use a null model that is able to preserve row and column totals such as the function permatswap() from the vegan package or the function vaznull() from the bipartite package. Either of these can be easily adapted to return only a single matrix (see examples). However, neither can accomodate non-integer matrices.

p_adj

string. Type of multiple hypothesis testing correction performed on P-values. Can take any valid method argument to p.adjust, including "none", "bonferroni", "holm", "fdr", and others (DEFAULT: "fdr").

seed

integer. Seed to use so that this is repeatable (DEFAULT: 1234557).

tails

integer. 1 = 1-tailed, test for specificity only. 2 = 2-tailed. 3 = 1-tailed, test for cosmopolitanism only. 0 = no test, P=1.0 (DEFAULT: 1).

n_cores

integer. Number of CPU cores to use for parallel operations. If set to 1, lapply will be used instead of mclapply (DEFAULT: 2).

verbose

logical. Should status messages be displayed? (DEFAULT: TRUE).

p_method

string. "raw" for quantile method, or "gamma_fit" for calculating P by fitting a gamma distribution (DEFAULT: "raw").

center

string. Type of central tendency to use for simulated RQE values. Options are "mean", "median", and "mode". If mode is chosen, a reversible gamma distribution is fit and mode is calculated using that distribution (DEFAULT: mean).

denom_type

string. Type of denominator (d) to use (DEFAULT: "index"). Note that denominator type does NOT affect P-values.

- "ses": d for species s is calculated as the standard deviation of RQE values calculated from permuted species weights. This makes the output specificity a standardized effect size (SES). Unfortunately, this makes SES counterintuitively sensitive to occupancy, where species with high occupancy have more extreme SES than rare species, due to their more deterministic sim specificities. Included for comparative purposes, not suggested.
- "raw": d is 1 for all species, so output specificity has units of distance, i.e. the raw difference between empirical and simulated RQE. This means that results from different variables are not comparable, since it is not scale-invariant to env or hosts_phylo. It not scale-invariant to the species weights in aunds_mat, either. Not sensitive to number of samples. Not suggested because units are strange, and isn't comparable between variables.
- "index": d is the mean of simulated (permuted) RQE values for species that have stronger specificity than expected by chance, resulting in specificity values with range [-1, 0), with 0 as the null hypothesis. In this case, -1 indicates perfect specificity, where a species is associated with zero environmental variability. In the euclidean sense, this could be a species that is always found at the exact same elevation or the exact same pH. For species

phy_or_env_spec

that have weaker specificity than expected by chance, d is x minus the center (see above) of simulated RQE values, where x is the maximum possible dissimilarity observable given species weights. This d has other useful properties: scale invariance to env/hosts_phylo, insensitivity to the number of samples, insensitivity to occupancy, and strong sensitivity to specificity (DEFAULT).

diagnostic

logical. If true, changes output to include different parts of SES. This includes Pval, SES, raw, denom, emp, and all sim values with column labels as simN where N is the number of sims (DEFAULT: FALSE)

Value

data.frame where each row is an input species. First column is P-value (\$Pval), second column is specificity (\$Spec).

Author(s)

John L. Darcy

References

- Poulin et al. (2011) Host specificity in phylogenetic and geographic space. Trends Parasitol 8:355-361. doi: 10.1016/j.pt.2011.05.003
- Rao CR (2010) Quadratic entropy and analysis of diversity. Sankhya 72:70-80. doi: 10.1007/s13171-010-0016-3
- Rao CR (1982) Diversity and dissimilarity measurements: A unified approach. Theor Popul Biol 21:24-43.

Examples

```
# phylogenetic specificity using endophyte data set
attach(endophyte)
# only analyze species with occupancy >= 20
m <- occ_threshold(prop_abund(zotutable), 20)</pre>
ses_host <- phy_or_env_spec(</pre>
    abunds_mat=m,
   hosts=metadata$PlantGenus,
   hosts_phylo=supertree,
    n_cores=12
)
# using vazquez null model from bipartite package as an alternate permutation:
# note that the "creating permuted matrices" step will be slow.
library(bipartite)
ses_host_vaz <- phy_or_env_spec(
    abunds_mat=m,
    hosts=metadata$PlantGenus,
    hosts_phylo=supertree,
   n_cores=12,
    sim_fun=function(m){bipartite::vaznull(1, m)[[1]]},
```

phy_spec_sim 15

```
)
# compare naive permutation vs. vazquez:
plot(ses_host$Spec, ses_host_vaz$Spec, ylab="bipartite::vaznull", xlab="naive")
abline(h=0);abline(v=0)
hist(ses_host_vaz$Spec)
hist(ses_host$Spec)
# environmental specificity using elevation from endophyte data set:
ses_elev <- phy_or_env_spec(</pre>
    abunds_mat=m,
    env=metadata$Elevation,
    n_cores=12
)
# geographic specificity using spatial data from endophyte data set:
ses_geo <- phy_or_env_spec(</pre>
    abunds_mat=m,
    env=distcalc(metadata$Lat, metadata$Lon),
    n_cores=12
)
```

phy_spec_sim

phy_spec_sim

Description

Simulates inputs for phy_or_env_spec, by creating a species distribution over an artificial (or real) host phylogenetic tree. For a phylogeny, the species probability distribution P(s) is based on patristic distances within the tree, such that P(s) is maximized at zero patristic distance between a tip in the tree and the ideal host species for s. This distribution is given by a a truncated normal distribution centered on zero, using only positive values. A uniform proportion (up) to that distribution may be added as well, to add a baseline probability to P(s). The standard deviation of P(s) can be raised or lowered to simulate cosmopolitanism or specificity.

Usage

```
phy_spec_sim(
   sdev,
   ideal,
   ideal2 = "",
   ideal3 = "",
   n_ideal = 1,
   hosts,
   hosts_phylo,
   n_obs,
   up = 0,
   oceanp = 0,
```

phy_spec_sim

```
n_cores = 2,
seed = 1234567
)
```

Arguments

sdev numeric vector. Standard deviation of the probability distribution P(s), in units of patristic distance in hosts_phylo. Low values mean that species s is found

with a narrow grouping of hosts, i.e. specificity. High values mean that s is found across a wider group of hosts, i.e. cosmopolitanism. Multiple values can be input in order to simulate a range of specificities, simultaneously. To get a handle on this somewhat opaque variable, consider plotting a histogram of patristic distances within hosts_phylo (see: ape::cophenetic.phylo). Can be

length 1 or n.

ideal character vector. Tip label of hosts_phylo that is ideal (or closest to ideal) for the

simulated species. Does not have to be in hosts, but MUST be in hosts_phylo.

Can be length 1 or n.

ideal2 character vector. Tip label of hosts_phylo that is secondary ideal host for the

simulated species. Does not have to be in hosts, but MUST be in hosts_phylo. Can be blank ("") if corresponding n_i deal < 2. Can be length 1 or n_i (default:

"").

ideal3 character vector. Tip label of hosts_phylo that is tertiary ideal host for the sim-

ulated species. Does not have to be in hosts, but MUST be in hosts_phylo. Can be blank ("") if corresponding n ideal < 3. Can be length 1 or n (default: "").

n_ideal integer vector. number of ideal hosts to use. Must be 1, 2, or 3 (DEFAULT=1).

hosts character vector. Real of fake host identities. All must be tips within hosts_phylo.

Analogous to env argument to env_spec_sim.

hosts_phylo phylo object. Tree containing all unique hosts as tips.

n_obs integer vector. Number of positive observations to make, i.e. occupancy of

simulated species. Can be length 1 or n.

up numeric vector. up=uniform proportion. This is the proportion of the probability

distribution P(species) that is composed of a uniform distribution, if desired. If set to a value above zero (and blow 1), P(species) will be a weighted sum of the normal distribution described above, and a uniform distribution. The weight for the uniform distribution will be up, and the weight for the normal distribution

will be 1-up (default: 0).

oceanp numeric vector. See ?env_spec_sim for help.

n_cores integer. Number of CPU cores for parallel computation (DEFAULT: 2).

seed integer. Seed for randomization. Daughter seeds will be generated for par-

allel computations, each with the same number of digits as seed (DEFAULT:

1234567).

Value

List object containing "matrix" and "params" objects:

plot_grid_abunds 17

matrix: matrix where each column is a vector of simulated observations corresponding to a value of hosts; each row represents a simulated species.

params: data.frame of parameters (columns) used to simulate each species (rows). A column called "index" is included so that simulated species can be mapped back onto original data structures when some species are ommitted due to simulation failure (see fail_rm).

Author(s)

John L. Darcy

Examples

none yet written.

plot_grid_abunds

plot_grid_abunds

Description

plots species abundances across spatial sampling locations

Usage

```
plot_grid_abunds(grid, abunds, pch = "", ...)
```

Arguments

grid data frame with columns x and y, representing cartesian coordinates of sample

locations. Can be artificial (generate with randomgrid()) or real.

abunds abundances of a species, corresponding to rows in grid.

pch pch character code to use for bottom of each abundance line (DEFAULT="")

... arguments to be passed to plot.

Value

returns nothing, just makes a plot.

Author(s)

John L. Darcy

18 plot_pairwise_spec

Examples

```
g1 <- randomgrid()
plot(g1)
a1 <- geo_spec_sim(sdev=c(30, 30, 30), n_obs=1000, grid=g1, up=c(0, 0.20, 0.40, 0.60))
par(mfrow=c(2,2))
plot_grid_abunds(g1, a1$matrix[,1])
plot_grid_abunds(g1, a1$matrix[,2])
plot_grid_abunds(g1, a1$matrix[,3])
plot_grid_abunds(g1, a1$matrix[,4])</pre>
```

plot_pairwise_spec

plot_pairwise_spec

Description

Plots pairwise correlations between specificity to multiple variables. Specificity results are supplied to this function as a list of specificity tables, i.e. a list where each object within the list is an output of phy_or_env_spec, and all were created using the same abunds_mat object (see: ?phy_or_env_spec).

Usage

```
plot_pairwise_spec(
    sl,
    label_cex = 1,
    point_cex = 1,
    cor_cex = 2,
    cor_red_lim = 0.7,
    method = "pearson"
)
```

Arguments

sl	"specs list" list of outputs from phy_or_env_spec as described above.
label_cex	float. Size of variable labels, which will be displayed along the plot's diagonal. Use cex units; see ?par (DEFAULT: 1).
point_cex	float. Size of points in the plot's lower triangle. Useful to reduce this if you are plotting lots of species. Use cex units; see ?par (DEFAULT: 1).
cor_cex	float. Size of text for correlations displayed in plot's upper triangle. Use cex units; see ?par (DEFAULT: 1).
cor_red_lim	float. Correlation coefficients will be shown in red if they are equal to or more extreme than this value (DEFAULT: 0.70).
method	string. Preferred correlation method. see ?cor for options (DEFAULT: "pearson").

plot_specificities 19

Value

Returns nothing. Plots correlations in a square matrix of subplots, where variable names are shown in the diagonal, pairwise specificities are plotted in the lower triangle, and correlation coefficients are displayed in the upper triangle. For plots in the lower triangle, each point represents a species.

Author(s)

John L. Darcy

Examples

```
library(specificity)
attach(endophyte)
otutable_over10 <- occ_threshold(otutable, threshold = 10)
specs_list <- list()
specs_list$NDVI <- phy_or_env_spec(otutable_over10, env=metadata$NDVI,
    n_cores=10, n_sim=100, p_method="gamma_fit")
specs_list$Evapotranspiration <- phy_or_env_spec(otutable_over10,
    env=metadata$Evapotranspiration, n_cores=10, n_sim=100, p_method="gamma_fit")
specs_list$Rainfall <- phy_or_env_spec(otutable_over10, env=metadata$Rainfall,
    n_cores=10, n_sim=100, p_method="gamma_fit")
plot_pairwise_spec(specs_list)</pre>
```

plot_specificities plot_specificities

Description

Visualizes results from phy_or_env_spec

Usage

```
plot_specificities(
   specs_list,
   n_bins = 20,
   col_sig = "black",
   col_nsig = "gray",
   col_bord = NA,
   alpha = 0.05,
   label_cex = 0.6
)
```

20 prop_abund

Arguments

specs_list	list of data.frames. Each data.frame must be an output from phy_or_env_spec; must have columns "Spec" and "Pval".
n_bins	integer. Number of bins for stacked violins (DEFAULT: 20).
col_sig	string. Color name or hex code for species where Pval <= alpha (DEFAULT = "black").
col_nsig	string. Color name or hex code for species where Pval > alpha (DEFAULT = "gray").
col_bord	string. Color name or hex code for border color. Use NA for no border (DE-FAULT = NA).
alpha	float. alpha value for determining statistical significance; see col_sig and col_nsig above (DEFAULT = 0.05).

Value

returns nothing (a plot is made).

Author(s)

John L. Darcy

Examples

none yet written.

prop_abund prop_abund

Description

Calculates proportional abundance of each species (columns) across samples (rows) in community data matrix m. Row sums of output matrix will all be 1.

Usage

```
prop_abund(
    m,
    to_int = FALSE,
    max_int = floor(sqrt(.Machine$integer.max)),
    speciesRows = FALSE
)
```

randomgrid 21

Arguments

m	matrix or data frame of numeric values. Columns represent species, rows are samples.
to_int	logical. Should output matrix be transformed into integers from 0 to max_int? Integers take up half as much space as doubles, and as weights are equivalent for calculating specificity. The tradeoff is a little bit of precision (DEFAULT: FALSE).
max_int	integer. Maximum integer value used for to_int. If pairwise geometric means will be calculated with these data, it is nice to keep this value as the square root of the maximum integer size, which is the default.
speciesRows	logical. Do rows represent species (instead of samples)? (DEFAULT:FALSE)

Value

matrix of proportional abundances.

Author(s)

John L. Darcy

Examples

```
attach(endophyte)
m_dbl <- prop_abund(zotutable)
m_int <- prop_abund(zotutable, to_int=TRUE)
head(rowSums(m_dbl))
head(rowSums(m_int))
# note that they are off by a little bit. This small loss in precision is OK.
object.size(m_dbl)
object.size(m_int)
random_positions <- random_rep_positions(m_dbl, 100)
plot(m_int[random_positions] ~ m_dbl[random_positions])</pre>
```

randomgrid randomgrid

Description

Generates a random spatial sampling using a bivariate random uniform distribution.

22 random_rep_positions

Usage

```
randomgrid(
  n_samp = 1000,
  xmin = -100,
  xmax = 100,
  ymin = -100,
  ymax = 100,
  seed = 123456
)
```

Arguments

```
n_samp number of sampling locations to output (DEFAULT=1000).

xmin minimum x-axis coordinate (DEFAULT=-100).

xmax maximum x-axis coordinate (DEFAULT=100).

ymin minimum y-axis coordinate (DEFAULT=-100).

ymax maximum y-axis coordinate (DEFAULT=100).

seed integer, seed for randomization.
```

Value

data.frame object with x and y columns, with n_samp rows.

Author(s)

```
John L. Darcy
```

Examples

```
g <- randomgrid()
plot(g)
g2 <- randomgrid(nsamp=50, xmin=0, ymin=0)
plot(g2)</pre>
```

```
random_rep_positions random_rep_positions
```

Description

Finds positions in a vector (or matrix) that are randomly located within n_bins evenly sized bins. This is useful for 1:1 comparisons of large vectors where plotting or comparing all points is prohibitive. Only used in an example for the prop_abund() function.

Usage

```
random_rep_positions(x, nbins = 50)
```

rao_genetic_max 23

Arguments

x vector

nbins number of bins to use

Value

integer vector of positions that were selected

Author(s)

John L. Darcy

rao_genetic_max rao_genetic_max

Description

Uses a genetic algorithm to find the optimum permutation of w to maximize Rao(w,D).

Usage

```
rao_genetic_max(w, D)
```

Arguments

w numeric vector of length n - a species weights vector.

D numeric vector of length n(n-1)/2 - i.e. a dist object whose full matrix is nxn.

term_cycles integer, number of cycles with no improvement to trigger termination (DE-

FAULT=10).

maxiters integer, maximum number of iterations to run algorithm (DEFAULT=400).

popsize integer, population size for genetic algorithm (DEFAULT=300).

keep integer, number of individuals to keep during each iteration (DEFAULT=5).

prc double, precision for calculating termination with term_cycles (DEFAULT=0.001).

Value

List object containing results of genetic algorithm:

best_rao: Maximum Rao value found.

iter_raos: Max Rao value for each iteration. If termination condition was met, rest of values after

final iteration are NA.

iterations: Iteration numbers, corresponding to iter_raos.

best_w: The best permutation of w found (corresponds to best_rao).

24 rao_sort_max

Author(s)

```
John L. Darcy
```

Examples

```
set.seed(12345)
w <- runif(100) * 20
D <- dist(sample(w))
a <- rao_genetic_max(w,D)
plot(a$iter_raos ~ a$iterations)</pre>
```

rao_sort_max

rao_sort_max

Description

Sorts pairwise_product(w) and D to approximate the maximum of Rao(w,D) under permutations of w.

Usage

```
rao_sort_max(w, D)
```

Arguments

w numeric vector of length n - a species weights vector.

D numeric vector of length n(n-1)/2 - i.e. a dist object whose full matrix is nxn.

Value

A single value, approximating maximum rao under permutations of w.

Author(s)

```
John L. Darcy
```

Examples

```
# none yet written.
```

spec_core 25

Description

C++ function used by phy_or_env_spec(). Not meant for use otherwise.

Usage

```
spec_core(w,D)
```

Arguments

w numeric matrix with m rows and n columns.

D numeric vector of length (n(n-1))/2.

Value

Vector of Rao specificities corresponding to columns of w.

Author(s)

John L. Darcy

tips_from_node

Description

Determines which tip indices in a phylogeny descend from a given node. Called by make_nested_set(), not intended for use otherwise, but some may find it handy. Data should come from a rooted phylogeny, but this function doesn't check that so be careful.

Usage

```
tips_from_node(nodes, anc, des)
```

Arguments

nodes	integer vector or scalar. The node index or indices for which tip indices are desired.
anc	integer vector. "ancestor" column vector from an adjacency matrix. For an ape::phylo object phy, anc=phy\$edge[,1].
des	integer vector. "descendant" column vector from an adjacency matrix. For an ape::phylo object phy. des=phy\$edge[.2].

26 tree2mat

Value

integer vector of tip indices, in no particular order.

Author(s)

John L. Darcy

See Also

ape::phylo

Examples

```
library(specificity)
phy <- get(data(endophyte))$supertree
# check if tree is rooted:
is.rooted(phy)
# which tips are in the Cucurbitales?
plot(phy) # need to stretch out the plot to see...
nodelabels(adj=c(0,-1), bg="yellow") # node numbers
nodelabels(phy$node.label, adj=c(0,1), bg="lightblue") # node names
# we can see that Cucurbitales is node 107
cuc_tips <- tips_from_node( nodes=107, anc=phy$edge[,1], des=phy$edge[,2] )
cuc_tips
phy$tip.label[cuc_tips]</pre>
```

tree2mat

tree2mat

Description

Transforms a phylogenetic tree into a dist object containing patristic distances between tips. Dists are just lower triangles of matrices, and the rows and columns of that matrix are defined by a user-supplied vector of tip labels, which can include duplicate values. Contrast with ape::cophenetic.phylo, which produces a distance matrix containing only unique pairwise patristic distances within the phylogeny.

Usage

```
tree2mat(tree, x, n_cores = 1, delim = ";")
```

Arguments

tree	physic object. Tree containing all unique species in x as tips. May contain tips that are not in x .
Х	character vector. Vector of species identities, each of which must be in tree as a

character vector. Vector of species identities, each of which must be in tree as a tip label. May contain any given species identity more than once.

wpd 27

be done if $n_cores = 1$. Multithreading should only be used for large trees where

x has low redundancy (DEFAULT =1).

delim string. Delimiter character or string for internal use. Must not be present in

tree\$tip.label. This is checked by the function and will return an error otherwise

(DEFAULT: ";").

Value

dist object, of vector length equal to $(1^2-1)/2$ where 1 is length(x); i.e. values are the lower triangle of a patristic distance matrix with rows=x and cols=x.

Author(s)

John L. Darcy

Examples

```
example_tree <- ape::read.tree(text=" ((((a:1,b:1):1,c:2):1,d:3):1,(e:1,f:1):3);")
example_x <- c("a", "a", "a", "b", "c", "d", "c", "a", "f")
# unique patristic distance matrix:
ape::cophenetic.phylo(example_tree)
# dist object for example_x:
tree2mat(tree=example_tree, x=example_x)
# examples with other delimiters
tree2mat(tree=example_tree, x=example_x, delim="@")
tree2mat(tree=example_tree, x=example_x, delim="i love cats")
# should fail since "a" is in a tip name:
tree2mat(tree=example_tree, x=example_x, delim="a")</pre>
```

wpd wpd

Description

Calculates weighted Phylogenetic Diversity for a vector s of species observations, weighted by the frequency of each species within s. For example, if S=a, a, b, a, b, c, a, then species a will have weight 4, species b will have weight 2, and species c will have weight 1. Unobserved species have weight zero.

Usage

```
wpd(s, s_phylo, w = NULL, nested_set = NULL, metric = "Hp")
```

28 wpd

Arguments

character vector. One species name per observation. If no species was observed s

for a given datum, use NA. s can also be provided as a vector of unique species identities, in which case counts of those species can be given as w.

s_phylo phylo object. Tree containing all unique names in s as tips. Must not contain

duplicate tip labels.

numeric vector. Optional weights for s, e.g. number of parasites observed in

each sample, or boolean weights corresponding to presence or absence of parasite species, or confidence species was observed, etc. If w is not provided but a weighted metric is specified, w will be set to 1 for each value of s. Thus, weights for each unique species in s would be equal to the number of times that species appears in s. w is not used for unweighted metrics (PD). Any NA values in w

will be pairwise removed from w and s (DEFAULT: NULL).

nested_set matrix. The output of make_nested_set(s_phylo). If not provided, will be cal-

culated on the fly. Precalculation only provides speedup with very large trees

(DEFAULT: NULL).

character. Abbreviated name of desired tree-based phylogenetic diversity met-

ric. Available metrics are:

Hp: Phylogenetic Entropy. Insensitive to 0 weights, cannot increase with removal of taxa. Allen et al. 2009.

WF: Weighted Faith's PD. Sensitive to 0 weights, i.e. a clade that was heavily sampled but has lots of zeroes will cause its sister clades to be underrepresented. Swenson 2014.

PD: Original Faith's Phylogenetic Diversity. Unweighted. Simply a sum of branch-lengths in your tree (but only for taxa in s). Faith 1992.

Details

However, one may wish to exclude observations that do not meet some criterion, such as coobservation of a symbiote or parasite. For this reason, a second set of weights w can be provided as a vector of numeric values that are paired with s. These weights are then implicitely combined with the weights discussed above depending on which weighted metric is chosen. In the case of Phylogenetic Entropy (Hw), per-tip weights are calculated as the sums of w. In the case of Weighted Faith (WF), per-tip weights are averages of w.

Value

Single WPD or PD value.

Author(s)

John L. Darcy

References

 Allen B, Kon M, Bar-Yam Y (2009) A new phylogenetic diversity measure generalizing the Shannon index and its application to Phyllostomid bats. American Naturalist 174(2).

metric

wpd_table 29

 Swenson NG (2014) Functional and Phylogenetic Ecology in R. Springer UseR! Series, Springer, New York, New York, U.S.A.

• Faith DP (1992) Conservation evaluation and phylogenetic diversity. Biological Conservation 61.

See Also

rao_quad_ent, a phylogenetic diversity measure that uses a distance matrix instead of a phylogenetic tree

Examples

```
library(specificity)
set.seed(12345)
s_phylo <- get(data(endophyte))$supertree
w <- sample(c(0, 1), replace=T, size=10)
s <- sample(s_phylo$tip.label, replace=T, size=10)
wpd(s, s_phylo, w, metric="Hp")</pre>
```

wpd_table

wpd table

Description

Calculates phylogenetic entropy (Hp) for each column vector s of species observations within matrix m, weighted by the frequency of each species within s. Can also calculate Faith's PD.

Usage

```
wpd_table(m, s_phylo, nested_set, metric = "Hp", ncores = 4)
```

Arguments

m matrix of species observation vectors (s). See s argument of wpd().

s_phylo phylo object. Tree containing all unique names in s as tips. Must not contain

duplicate tip labels.

nested_set matrix. The output of make_nested_set(s_phylo). If not provided, will be cal-

culated on the fly. Precalculation only provides speedup with very large trees

(DEFAULT: NULL).

metric character. Abbreviated name of desired tree-based phylogenetic diversity met-

ric. Available metrics are:

Hp: Phylogenetic Entropy. Insensitive to 0 weights, cannot increase with re-

moval of taxa. Allen et al. 2009.

WF: Weighted Faith's PD. Sensitive to 0 weights, i.e. a clade that was heavily sampled but has lots of zeroes will cause its sister clades to be underrepre-

sented. Swenson 2014.

PD: Original Faith's Phylogenetic Diversity. Unweighted. Simply a sum of branch-lengths in your tree (but only for taxa in s). Faith 1992.

30 wpd_table

Value

multiple WPD or PD values, one for each column of m.

Author(s)

John L. Darcy

References

- Allen B, Kon M, Bar-Yam Y (2009) A new phylogenetic diversity measure generalizing the Shannon index and its application to Phyllostomid bats. American Naturalist 174(2).
- Swenson NG (2014) Functional and Phylogenetic Ecology in R. Springer UseR! Series, Springer, New York, New York, U.S.A.
- Faith DP (1992) Conservation evaluation and phylogenetic diversity. Biological Conservation 61.

Examples

none yet written.

Index

```
bl\_distance\_ns, 2
check_pes_inputs, 3
daughter_seeds, 4
\operatorname{distcalc}, 4
env_spec_sim, 5
geo_spec_sim, 7
{\sf make\_nested\_set}, 9
occ\_threshold, 10
\verb"pairwise_product", 11
phy_or_env_spec, 12
phy_spec_sim, 15
plot_grid_abunds, 17
plot_pairwise_spec, 18
plot_specificities, 19
prop_abund, 20
random_rep_positions, 22
\texttt{randomgrid}, \textcolor{red}{21}
rao_genetic_max, 23
rao_sort_max, 24
spec_core, 25
tips_from_node, 25
tree2mat, 26
wpd, 27
wpd_table, 29
```