Package 'phylospatial'

January 24, 2025

Title Spatial Phylogenetic Analysis

Version 1.0.0

Description Conduct various analyses on spatial phylogenetics. Use your data on an evolutionary tree and geographic distributions of the terminal taxa to compute diversity and endemism metrics, test significance with null model randomization, analyze community turnover and biotic regionalization, and perform spatial conservation prioritizations. All functions support quantitative community data in addition to binary data.

License MIT + file LICENSE

Encoding UTF-8

RoxygenNote 7.3.2

Imports ape, sf, stats, terra, vegan

Suggests canaper, furrr, future, test that $(>= 3.0.0)$, betapart, knitr, rmarkdown, tmap, magrittr

URL <https://matthewkling.github.io/phylospatial/>

Config/testthat/edition 3

Depends R ($>= 3.5$)

VignetteBuilder knitr

NeedsCompilation no

Author Matthew Kling [aut, cre, cph] (<<https://orcid.org/0000-0001-9073-4240>>)

Maintainer Matthew Kling <mattkling@berkeley.edu>

Repository CRAN

Date/Publication 2025-01-24 10:40:02 UTC

Contents

2 benefit

Index [27](#page-26-0)

benefit *Calculate taxon conservation benefit*

Description

Nonlinear function that converts proportion of range conserved into conservation "benefit."

Usage

benefit(x , lambda = 1)

Arguments

Value

Value between 0 and 1.

Description

Get example phylospatial data set based on a phylogeny and modeled distributions of 443 moss species across California. This data set is a coarser version of data from Kling et al. (2024). It contains occurrence probabilities, and is available in raster or polygon spatial formats.

Usage

moss(format = "raster")

Arguments

format Either "raster" (default) or "polygon"

Value

a phylospatial object

Source

Kling, Gonzalez-Ramirez, Carter, Borokini, and Mishler (2024) bioRxiv, https://doi.org/10.1101/2024.12.16.628580.

Examples

moss()

phylospatial *Create a spatial phylogenetic object*

Description

This function creates a phylospatial object. This is the core data type in the phylospatial library, and is a required input to most other functions in the package. The two essential components of a spatial phylogenetic object are a phylogenetic tree and an community data set.

Usage

```
phylospatial(
  comm,
  tree = NULL,
  spatial = NULL,data_type = c("auto", "probability", "binary", "abundance", "other"),
  clade_fun = NULL,
  build = TRUE,check = TRUE,area_to1 = 0.01\mathcal{L}
```
Arguments

Details

This function formats the input data as a phylospatial object. Beyond validating, cleaning, and restructing the data, the main operation it performs is to compute community occurrence data for every internal clade on the tree. For a given clade and site, community data for all the terminals in the clade are used to calculate the clade's occurrence value in the site. As described above, this calculation can happen in various ways, depending on what type of community data you have (e.g. binary, probability, or abundance) and how you want to summarize them. By default, the function tries to detect your data_type and use it to automatically select an appropriate summary function as described above, but you can override this by providing your own function to clade_fun.

You can also disable construction of the clade community matrix columns altogether by setting build = FALSE). This is atypical, but you might want to use this option if you have your own distribution data data on all clades (e.g. from modeling occurrence probabilities for clades in addition to terminal species), or if your community data comes from a previously-constructed phylospatial object.

Value

A phylospatial object, which is a list containing the following elements:

"data type": Character indicating the community data type

"tree": Phylogeny of class phylo

- "comm": Community matrix, including a column for every terminal taxon and every larger clade. Column order corresponds to tree edge order.
- "spatial": A SpatRaster or sf providing spatial coordinates for the rows in comm. May be missing if no spatial data was supplied.
- "dissim": A community dissimilary matrix of class dist indicating pairwise phylogenetic dissimilarity between sites. Missing unless ps_dissim ..., add = TRUE) is called.

Examples

```
# load species distribution data and phylogeny
comm <- terra::rast(system.file("extdata", "moss_comm.tif", package = "phylospatial"))
tree <- ape::read.tree(system.file("extdata", "moss_tree.nex", package = "phylospatial"))
# construct `phylospatial` object
ps <- phylospatial(comm, tree)
ps
```
plot.phylospatial *Plot a* phylospatial *object*

Description

Plot a phylospatial object

Usage

S3 method for class 'phylospatial' $plot(x, y = c("tree", "comm"), max_taxa = 12, ...)$

Arguments

Value

A plot of the tree or community data.

Examples

```
ps <- ps_simulate()
plot(ps, "tree")
plot(ps, "comm")
```
plot_lambda *Plot alternative lambda values*

Description

Show a plot illustrating alternative values for the lambda parameter in [ps_prioritize.](#page-14-1) Lambda determines the shape of the "benefit" function that determines the conservation value of protecting a given proportion of the geographic range of a species or clade. Positive values place a higher priority on protecting additional populations of largely unprotected taxa, whereas negative values place a higher priority on protecting additional populations of relatively well-protected taxa. The default value used by [ps_prioritize](#page-14-1) is 1.

Usage

```
plot_lambda(lambda = c(-1, -0.5, 0, 0.5, 2, 1))
```
ps_add_dissim 7

Arguments

lambda A vector of lambda values to plot

Value

Plots a figure

Examples

```
plot_lambda()
plot_lambda(seq(0, 3, .1))
```
ps_add_dissim *Add community dissimilarity data to a* phylospatial *object*

Description

This function calculates pairwise phylogenetic dissimilarity between communities and returns the phylospatial object with the dissimilarity data added as an element called dissim. See [ps_dissim](#page-9-1) for details.

Usage

```
ps_add_dissim(ps, method = "sorensen", ...)
```
Arguments

Value

ps with a new dissim element added.

Examples

```
ps <- ps_simulate(data_type = "prob")
ps_add_dissim(ps)
ps_add_dissim(ps, fun = "vegdist", method = "jaccard", endemism = TRUE)
```


Description

This function classifies sites into areas of significant endemism according to the scheme of Mishler et al. (2014). Categorization is based on randomization quantile values for PE, RPE, and CE (which Mishler et al. call "PE on the comparison tree").

Usage

 $ps_canape(rand, alpha = 0.05)$

Arguments

Details

Endemism significance categories are defined as follows:

- Endemism not significant: neither PE nor CE are significantly high at alpha.
- Significant neoendemism: PE or CE are significantly high at alpha; RPE significantly low at alpha / 2 (two-tailed test).
- Significant paleoendemism: PE or CE are significantly high at alpha; RPE significantly high at alpha / 2 (two-tailed test)..
- Significant mixed-endemism: PE or CE are significantly high at alpha; RPE not significant.
- Significant super-endemism: PE or CE are significantly high at alpha / 5; RPE not significant.

Value

An object of the same class as rand containing a variable called "canape", with values 0-4 corresponding to not-significant, mixed-, super-, neo-, and paleo-endemism, respectively.

References

Mishler, B. D., Knerr, N., González-Orozco, C. E., Thornhill, A. H., Laffan, S. W., & Miller, J. T. (2014). Phylogenetic measures of biodiversity and neo-and paleo-endemism in Australian Acacia. Nature Communications, 5(1), 4473.

ps_canaper 9

Examples

```
# classic CANAPE using binary data and the curveball algorithm
# (note that a real analysis would require a much higher `n_rand`)
set.seed(123456)
ps <- moss()
rand <- ps_rand(ps, metric = c("PE", "RPE", "CE"),
                fun = "nullmodel", method = "curveball",
                nrand = 25, burnin = 10000, progress = FALSE)
canape <- ps_canape(rand)
terra::plot(canape)
```
ps_canaper *Binary randomization tests including CANAPE*

Description

This function is a wrapper around canaper::cpr_rand_test(). It only works with binary community data. It is largely redundant with ps_rand() and ps_canape(), which are more flexible in supporting data sets with non-binary community data. However, this function runs faster, and supports custom null models via [make.commsim.](#page-0-0)

Usage

```
ps\_canaper(ps, null_model = "curveball", spatial = TRUE, ...)
```
Arguments

Details

This function runs canaper::cpr_rand_test(); see the help for that function for details.

It also runs canaper::cpr_classify_endem() on the result, and includes the resulting classification as an additional variable, 'endem_type', in the output. 'endem_type' values 0-4 correspond to not-significant, neo, paleo, mixed, and super endemesim, respectively.

Value

A matrix or SpatRaster, or sf with a column or layer for each metric.

References

Mishler, B. D., Knerr, N., González-Orozco, C. E., Thornhill, A. H., Laffan, S. W., & Miller, J. T. (2014). Phylogenetic measures of biodiversity and neo-and paleo-endemism in Australian Acacia. Nature Communications, 5(1), 4473.

Nitta, J. H., Laffan, S. W., Mishler, B. D., & Iwasaki, W. (2023). canaper: categorical analysis of neo-and paleo-endemism in R. Ecography, 2023(9), e06638.

See Also

```
ps_canape(), ps_rand()
```
Examples

```
if(requireNamespace("canaper")){
     ps <- ps_simulate(data_type = "binary")
      terra::plot(ps_canaper(ps)$pd_obs_p_upper)
}
```


```
ps_dissim Quantitative phylogenetic dissimilarity
```
Description

This function calculates pairwise phylogenetic dissimilarity between communities. It works with both binary and quantitative community data sets. A wide range of phylogentic community dissimilarity metrics are supported, including phylogenetic Sorensen's and Jaccard's distances, turnover and nestedness components of Sorensen's distance (Baselga & Orme, 2012), and phylogenetic versions of all community distance indices provided through the vegan library. The function also includes options to scale the community matrix in order to focus the analysis on endemism and/or on proportional differences in community composition. The results from this function can be visualized using [ps_rgb](#page-21-1) or [ps_regions,](#page-19-1) or used in a variety of statistical analyses.

Usage

```
ps_dissim(
  ps,
 method = "sorensen",
  fun = c("vegdist", "designdist", "chaodist"),
  endemism = FALSE,
 normalize = FALSE,
  ...
)
```


ps_dissim 11

Arguments

Value

A pairwise phylogenetic dissimilarity matrix of class dist.

References

Graham, C. H., & Fine, P. V. (2008). Phylogenetic beta diversity: linking ecological and evolutionary processes across space in time. Ecology Letters, 11(12), 1265-1277.

Baselga, A., & Orme, C. D. L. (2012). betapart: an R package for the study of beta diversity. Methods in Ecology and Evolution, 3(5), 808-812.

Pavoine, S. (2016). A guide through a family of phylogenetic dissimilarity measures among sites. Oikos, 125(12), 1719-1732.

See Also

[ps_add_dissim\(\)](#page-6-1)

Examples

```
# example data set:
ps <- ps_simulate(n_tips = 50)
```

```
# The default arguments give Sorensen's quantitative dissimilarity index
# (a.k.a. Bray-Curtis distance):
d <- ps_dissim(ps)
# Specifying a custom formula explicitly via `designdist`;
# (this is the Bray-Curtis formula, so it's equivalent to the prior example)
d \leq p s\_dissim(ps, method = "(b+c)/(2*a+b+c)",fun = "designdist", terms = "minimum", abcd = TRUE)
# Alternative arguments can specify a wide range of dissimilarity measures;
# here's endemism-weighted Jaccard's dissimilarity:
d <- ps_dissim(ps, method = "jaccard", endemism = TRUE)
```


ps_diversity *Calculate spatial phylogenetic diversity and endemism metrics*

Description

This function calculates a variety of diversity and endemism metrics including Faith's phylogenetic diversity, Shannon phylogenetic entropy, Simpson phylogentic diversity, relative phylogentic diversity, richness of clades, richness of terminals (typically species), and versions of all these metrics weighted by endemism (i.e. rarity). If continuous community data (probabilities or abundances) are provided, they are used in calculations, giving quantitative versions of the classic binary metrics.

Usage

ps_diversity(ps, metric = "all", spatial = TRUE)

Arguments

Details

The function calculates the following metrics:

- TD—Terminal Diversity, i.e. richness of terminal taxa (in many cases these are species): $\sum_t p_t$
- TE—Terminal Endemism, i.e. total endemism-weighted diversity of terminal taxa, a.k.a. "weighted endemism": $\sum_t p_t r_t^{-1}$
- CD—Clade Diversity, i.e. richness of taxa at all levels (equivalent to PD on a cladogram): $\sum_b p_b$
- CE—Clade Endemism, i.e. total endemism-weighted diversity of taxa at all levels (equivalent to PE on a cladrogram): $\sum_b p_b r_b^{-1}$
- PD—Phylogenetic Diversity: $\sum_b L_b p_b$
- PE—Phylogenetic Endemism, i.e. endemism-weighted PD: $\sum_b L_b p_b r_b^{-1}$
- RPD—Relative Phylogenetic Diversity, i.e. branch length of mean resident (equivalent to PD / CR): $\sum_b L_b p_b / \sum_b p_b$
- RPE—Relative Phylogenetic Endemism, i.e. mean endemism-weighted branch length (equivalent to PE / CE): $\sum_{b} L_b p_b r_b^{-1} / \sum_{b} p_b r_b^{-1}$
- ShPD—Shannon Phylogenetic Diversity, a.k.a. "phylogenetic entropy": $-\sum_{b} L_b n_b log(n_b)$
- ShPE–Shannon phylogenetic Endemism, an endemism-weighted version of ShPD: $-\sum_b L_b n_b log(e_b) r_b^{-1}$
- SiPD—Simpson Phylogenetic Diversity: $1/\sum_b L_b n_b^2$
- SiPE—Simpson Phylogenetic Endemism, an endemism-weighted version of SiPD: $1/\sum_b L_b r_b^{-1} e_b^2$

where b indexes all taxa including terminals and larger clades; t indexes terminals only; p_i is the occurrence value (binary, probability, or abundance) of clade/terminal i in a given community; L_b is the length of the phylogenetic branch segment unique to clade b; and r_i is the sum of p_i across all sites.

For Shannon and Simpson indices, only nonzero elements of p_b are used, $n_b = p_b / \sum_b p_b L_b$, and $e_b = p_b / \sum_b p_b L_b r_b^{-1}.$

Value

A matrix, sf data frame, or SpatRaster with a column or layer for each requested diversity metric.

References

Faith, D. P. (1992). Conservation evaluation and phylogenetic diversity. Biological Conservation, 61(1), 1-10.

Laffan, S. W., & Crisp, M. D. (2003). Assessing endemism at multiple spatial scales, with an example from the Australian vascular flora. Journal of Biogeography, 30(4), 511-520.

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.

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(2), 236-243.

Chao, A., Chiu, C. H., & Jost, L. (2010). Phylogenetic diversity measures based on Hill numbers. Philosophical Transactions of the Royal Society B: Biological Sciences, 365(1558), 3599-3609.

Mishler, B. D., Knerr, N., González-Orozco, C. E., Thornhill, A. H., Laffan, S. W., & Miller, J. T. (2014). Phylogenetic measures of biodiversity and neo-and paleo-endemism in Australian Acacia. Nature Communications, 5(1), 4473.

Kling, M. M., Mishler, B. D., Thornhill, A. H., Baldwin, B. G., & Ackerly, D. D. (2019). Facets of phylodiversity: evolutionary diversification, divergence and survival as conservation targets. Philosophical Transactions of the Royal Society B, 374(1763), 20170397.

Examples

```
ps <- ps_simulate()
div <- ps_diversity(ps)
terra::plot(div)
```
ps_get_comm *Get* phylospatial *community data*

Description

Get phylospatial community data

Usage

```
ps_get_comm(ps, tips_only = TRUE, spatial = TRUE)
```
Arguments

Value

Either a SpatRaster with a layer for every taxon, or an sf data frame with a variable for every taxon, depending on which data type was used to create ps.

Examples

```
ps <- ps_simulate()
# the defaults return a spatial object of terminal taxa distributions:
ps_get_comm(ps)
# get distributions for all taxa, as a matrix
pcomm <- ps_get_comm(ps, tips_only = FALSE, spatial = FALSE)
```


Description

Perform an ordination that reduces a spatial phylogenetic data set into k dimensions, using one of several alternative ordination algorithms.

Usage

```
ps\_ordinate(ps, method = c("nmds", "cmds", "pca"), k = 3, spatial = TRUE)
```
Arguments

Value

A matrix or spatial object with k variables.

Examples

```
ps <- ps_add_dissim(ps_simulate(50, 5, 5))
ord \leq ps_ordinate(ps, method = "cmds", k = 4)
terra::plot(ord)
```
ps_prioritize *Phylogenetic conservation prioritization*

Description

Create a ranking of conservation priorities using optimal or probabilistic forward stepwise selection. Prioritization accounts for the occurrence quantities for all lineages present in the site, including terminal taxa and larger clades; the evolutionary branch lengths of these lineages on the phylogeny, which represent their unique evolutionary heritage; the impact that protecting the site would have on these lineages' range-wide protection levels; the compositional complementarity between the site, other high-priority sites, and existing protected areas; the site's initial protection level; the relative cost of protecting the site; and a free parameter "lambda" determining the shape of the conservation benefit function.

Usage

```
ps_prioritize(
  ps,
  init = NULL,
  cost = NULL,lambda = 1,
  protection = 1,
  max_iter = NULL,
  method = c("optimal", "probable"),
  trans = function(x) replace(x, which(rank(-x) > 25), 0),
  n_reps = 100,
  n\_cores = 1,summarize = TRUE,
  spatial = TRUE,progress = interactive()
\mathcal{L}
```
Arguments

n_reps optimizations are performed, and ranks for each site are summarized across repetitions. trans A function that transforms marginal values into relative selection probabilities; only used if method = "probable". The function should take a vector of positive numbers representing marginal values and return an equal-length vector of positive numbers representing a site's relative likelihood of being selected. The default function returns the marginal value if a site is in the top 25 highest-value sites, and zero otherwise. n_reps Number of random repetitions to do; only used if method = "probable". Depending on the data set, a large number of reps (more than the default of 100) may be needed in order to achieve a stable result. This may be a computational barrier for large data sets; multicore processing via n_cores can help. n_cores Number of compute cores to use for parallel processing; only used if method = "probable". summarize Logical: should summary statistics across reps (TRUE, default) or the reps themselves (FALSE) be returned? Only relevant if method = "probable". spatial Logical: should the function return a spatial object (TRUE, default) or a matrix (FALSE)? progress Logical: should a progress bar be displayed?

Details

This function uses the forward stepwise selection algorithm of Kling et al. (2019) to generate a ranked conservation prioritization. Prioritization begins with the starting protected lands network identified in init, if provided. At each iteration, the marginal conservation value of fully protecting each site is calculated, and a site is selected to be added to the reserve network. Selection can happen either in an "optimal" or "probable" fashion as described under the method argument. This process is repeated until all sites are fully protected or until max_iter has been reached, with sites selected early in the process considered higher conservation priorities.

The benefit of the probabilistic approach is that it relaxes the potentially unrealistic assumption that protected land will actually be added in the optimal order. Since the algorithm avoids compositional redundancy between high-priority sites, the optimal approach will never place high priority on a site that has high marginal value but is redundant with a slightly higher-value site, whereas the probabilistic approach will select them at similar frequencies (though never in the same randomized run).

Every time a new site is protected as the algorithm progresses, it changes the marginal conservation value of the other sites. Marginal value is the increase in conservation benefit that would arise from fully protecting a given site, divided by the cost of protecting the site. This is calculated as a function of the site's current protection level, the quantitative presence probability or abundance of all terminal taxa and larger clades present in the site, their evolutionary branch lengths on the phylogeny, the impact that protecting the site would have on their range-wide protection levels, and the free parameter lambda. lambda determines the relative importance of protecting a small portion of every taxon's range, versus fully protecting the ranges of more valuable taxa (those with longer evolutionary branches and smaller geographic ranges).

Matrix or spatial object containing a ranking of conservation priorities. Lower rank values represent higher conservation priorities. All sites with a lower priority than max _iter have a rank value equal to the number of sites in the input data set (i.e. the lowest possible priority).

If method = "optimal". the result contains a single variable "priority" containing the ranking.

- If method = "probable" and summarize = TRUE, the "priority" variable gives the average rank across reps, variables labeled "pctX" give the Xth percentile of the rank distribution for each site, variables labeled "topX" give the proportion of reps in which a site was in the top X highest-priority sites, and variables labeled "treX" give a ratio representing "topX" relative to the null expectation of how often "top X " should occur by chance alone.
- If method = "probable" and summarize = FALSE, the result contains the full set of n -rep solutions, each representing the the ranking, with low values representing higher priorities..

References

Kling, M. M., Mishler, B. D., Thornhill, A. H., Baldwin, B. G., & Ackerly, D. D. (2019). Facets of phylodiversity: evolutionary diversification, divergence and survival as conservation targets. Philosophical Transactions of the Royal Society B, 374(1763), 20170397.

See Also

[benefit\(\)](#page-1-1), [plot_lambda\(\)](#page-5-1)

terra::plot(p\$top10)

Examples

```
# simulate a toy `phylospatial` data set
set.seed(123)
ps <- ps_simulate()
# basic prioritization
p <- ps_prioritize(ps)
# specifying locations of initial protected areas
# (can be binary, or can be continuous values between 0 and 1)
# here we'll create an `init` raster with arbitrary values ranging from 0-1,
# using the reference raster layer that's part of our `phylospatial` object
protected <- terra::setValues(ps$spatial, seq(0, 1, length.out = 400))
cost <- terra::setValues(ps$spatial, rep(seq(100, 20, length.out = 20), 20))
p \leq ps_prioritize(ps, init = protected, cost = cost)
# using probabilistic prioritization
p <- ps_prioritize(ps, init = protected, cost = cost,
      method = "prob", n_{res} = 1000, max_{iter} = 10)
```
Value

Description

This function compares to diversity metrics calculated in [ps_diversity](#page-11-1) to their null distributions computed by randomizing the community matrix. Randomization is done using the [quantize](#page-23-1) method for community matrices containing continuous quantities such as occurrence probabilities or abundances.

Usage

```
ps_rand(
  ps,
  metric = "all",
  fun = "quantize",
  method = "curveball",
  n<sub>r</sub>and = 100,
  spatial = TRUE,
  n\_cores = 1,progress = interactive(),
  ...
\mathcal{L}
```
Arguments

Value

A matrix with a row for every row of x, a column for every metric specified in metric, and values indicating the proportion of randomizations in which the observed diversity metric was greater than the randomized metric. Or if spatial = TRUE, a sf or SpatRaster object containing these data.

See Also

[ps_diversity\(\)](#page-11-1)

Examples

```
# simulate a `phylospatial` data set and run randomization with default settings
ps <- ps_simulate(data_type = "prob")
rand \leq ps_rand(ps)
# using the default `quantize` function, but with alternative arguments
rand \leq ps_rand(ps, transform = sqrt, n_strata = 4, priority = "rows")
# using binary data
ps2 <- ps_simulate(data_type = "binary")
rand \leq ps_rand(ps2, fun = "nullmodel", method = "r2")
# using abundance data, and demonstrating alternative metric choices
ps3 <- ps_simulate(data_type = "abund")
rand <- ps_rand(ps3, metric = c("ShPD", "SiPD"), fun = "nullmodel", method = "abuswap_c")
rand
```
ps_regions *Cluster analysis to identify phylogenetic regions*

Description

Perform a clustering analysis that categorizes sites into biogeographic regions based on phylogenetic community compositional similarity.

Usage

```
ps_regions(ps, k = 5, method = "average", endemism = FALSE, normalize = TRUE)
```
Arguments

Value

A raster or matrix with an integer indicating which of the k regions each site belongs to.

References

Daru, B. H., Elliott, T. L., Park, D. S., & Davies, T. J. (2017). Understanding the processes underpinning patterns of phylogenetic regionalization. Trends in Ecology & Evolution, 32(11), 845-860.

Examples

```
ps <- ps_simulate()
# using kmeans clustering algorithm
terra::plot(ps_regions(ps, method = "kmeans"))
# to use a hierarchical clustering method, first we have to `ps_add_dissim()`
terra::plot(ps_regions(ps_add_dissim(ps), k = 7, method = "average"))
```
ps_regions_eval *Evaluate region numbers*

Description

This function compares multiple potential values for k, the number of clusters in to use in ps_regions(), to help you decide how well different numbers of regions fit your data set. For each value of k, it performs a cluster analysis and calculates the proportion of total variance explained (SSE, the sum

of squared pairwise distances explained). It also calculates second-order metrics of the relationship between k and SSE. While many data sets have no optimal value of k and the choice is often highly subjective, these evaluation metrics can help you identify potential points where the variance explained stops increasing quickly as k increases.

Usage

 $ps_{\text{regions_eval}(ps, k = 1:20, plot = TRUE, ...)$

Arguments

Value

The function generates a data frame with the following columns. If plot = FALSE the data frame is returned, otherwise the function prints a plot of the latter variables as a function of k:

- "k": The number of clusters.
- "sse": The proportion of total variance explained, with variance defined as squared pairwise community phylogenetic dissimilarity between sites.
- "curvature": The local second derivative. Lower (more negative) values indicate more attractive break-point values of k.
- "dist11": The distance from the point to the 1:1 line on a plot of k vs sse in which k values over the interval from 1 to the number of sites are rescaled to the unit interval. Higher values indicate more attractive values for k.

Examples

```
ps <- ps_add_dissim(ps_simulate())
ps_{\text{regions\_eval}(ps, k = 1:15, plot = TRUE)}
```
ps_rgb *Map phylospatial data onto RGB color bands*

Description

Perform an ordination that reduces a spatial phylogenetic data set into three dimensions that can be plotted as the RGB bands of color space to visualize spatial patterns of community phylogenetic composition. This function is a wrapper around ps_ordinate().

ps_simulate 23

Usage

```
ps_rgb(ps, method = c("nmds", "cmds", "pca"), trans = identity, spatial = TRUE)
```
Arguments

Value

A matrix or spatial object with three variables containing RGB color values in the range 0-1.

Examples

```
ps <- ps_add_dissim(moss())
RGB < -ps\_rgb(ps, method = "cmds")terra::plotRGB(RGB * 255, smooth = FALSE)
```

```
ps_simulate Simulate a toy spatial phylogenetic data set
```
Description

This function generates a simple phylospatial object that can be used for testing other functions in the package. It is not intended to be realistic.

Usage

```
ps_simulate(
  n\_tips = 10,
  n_{-}x = 20,
  n_y = 20,
  data_type = c("probability", "binary", "abundance"),
  spatial_type = c("raster", "none"),
  seed = NULL
\mathcal{E}
```
Arguments

Value

phylospatial object, comprising a random phylogeny and community matrix in which each terminal has a circular geographic range with a random radius and location. The spatial reference data is a SpatRaster.

Examples

```
# using all the defaults
ps_simulate()
# specifying some arguments
plot(ps_simulate(n_tips = 50, n_x = 30, n_y = 40, data_type = "abundance"), "comm")
```
quantize *Stratified randomization of community matrix*

Description

This is a community null model method for quantitative community data (e.g. abundance or occurrence probability). It is designed to adapt binary null model algorithms for use with quantitative data, which can be useful if there is not a quantitative-specific algorithm available that has the desired properties. For example, use with the binary "curveball" algorithm preserves row and column totals, and also approximately preserves the marginal distributions of rows and columns. For each randomization, the data set is split into strata representing numerical ranges of the input quantities, a separate binary randomization is done for each stratum, and the results are combined to produce a randomized, quantitative community matrix. See vegan::commsim() for details about other binary and quantitative null models.

Usage

quantize(x, method = "curveball", \ldots)

to_spatial 25

Arguments

Value

A randomized version of x.

Examples

```
# example quantitative community matrix
comm <- ps_get_comm(moss("polygon"), tips_only = TRUE, spatial = FALSE)[1:50, 1:50]
# examples of different quantize usage
rand <- quantize(comm)
rand <- quantize(comm, n_strata = 4, transform = sqrt, priority = "rows")
rand <- quantize(comm, method = "swap", burnin = 10)
# (note: this `burnin` value is far too small for a real analysis)
```
to_spatial *Convert a site-by-variable matrix into a SpatRaster or sf object*

Description

Convert a site-by-variable matrix into a SpatRaster or sf object

26 to_spatial control of the control of th

Usage

to_spatial(m, template)

Arguments

Value

SpatRaster with a layer for every column in m, or sf data frame with a variable for every column in m, depending on the data type of template.

Examples

```
ps <- moss()
to_spatial(ps$comm[, 1:5], ps$spatial)
```
Index

benefit, [2](#page-1-0) benefit(), *[18](#page-17-0)* chaodist, *[11](#page-10-0)* commsim, *[19](#page-18-0)* designdist, *[11](#page-10-0)* hclust, *[21](#page-20-0)* make.commsim, *[9](#page-8-0)* moss, [3](#page-2-0) nullmodel, *[19](#page-18-0)* phylo, *[4](#page-3-0)* phylospatial, [3](#page-2-0) plot, *[6](#page-5-0)* plot.phylo, *[6](#page-5-0)* plot.phylospatial, [6](#page-5-0) plot.sf, *[6](#page-5-0)* plot_lambda, [6,](#page-5-0) *[16](#page-15-0)* plot_lambda(), *[18](#page-17-0)* ps_add_dissim, [7,](#page-6-0) *[15](#page-14-0)*, *[21–](#page-20-0)[23](#page-22-0)* ps_add_dissim(), *[11](#page-10-0)* ps_canape, [8](#page-7-0) ps_canape(), *[10](#page-9-0)* ps_canaper, [9](#page-8-0) ps_dissim, *[7](#page-6-0)*, [10](#page-9-0) ps_diversity, [12,](#page-11-0) *[19](#page-18-0)* ps_diversity(), *[20](#page-19-0)* ps_get_comm, [14](#page-13-0) ps_ordinate, [15](#page-14-0) ps_prioritize, *[6](#page-5-0)*, [15](#page-14-0) ps_rand, [19](#page-18-0) ps_rand(), *[10](#page-9-0)* ps_regions, *[10](#page-9-0)*, [20,](#page-19-0) *[22](#page-21-0)* ps_regions_eval, *[21](#page-20-0)*, [21](#page-20-0) ps_rgb, *[10](#page-9-0)*, [22](#page-21-0) ps_simulate, [23](#page-22-0)

quantize, *[19,](#page-18-0) [20](#page-19-0)*, [24](#page-23-0) simulate.nullmodel, *[19,](#page-18-0) [20](#page-19-0)*, *[25](#page-24-0)* SpatRaster, *[4](#page-3-0)* to_spatial, [25](#page-24-0) vegdist, *[11](#page-10-0)*