

Package: canaper (via r-universe)

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Title Categorical Analysis of Neo- And Paleo-Endemism

Version 1.0.1.9000

Description Provides functions to analyze the spatial distribution of biodiversity, in particular categorical analysis of neo- and paleo-endemism (CANAPE) as described in Mishler et al (2014) <[doi:10.1038/ncomms5473](https://doi.org/10.1038/ncomms5473)>. 'canaper' conducts statistical tests to determine the types of endemism that occur in a study area while accounting for the evolutionary relationships of species.

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<https://docs.ropensci.org/canaper/>

BugReports <https://github.com/ropensci/canaper/issues>

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acacia	<i>Acacia example data</i>
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Description

Dataset of Australian *Acacia* from Mishler et al. 2014 (Nat. Comm.)

Usage

acacia

Format

A list with two elements:

phy Phylogeny of Australian *Acacia* (list of class "phylo"). Tip labels are specific epithet, except for the outgroup, which includes genus and specific epithet. Includes 508 ingroup taxa (genus *Acacia*) and two outgroup taxa.

comm Community matrix of Australian *Acacia* (dataframe). Column names are specific epithet of each species. Row names are centroids of 50km grid cells in Australian Albers equal area EPSG:3577 projection. 3037 rows (sites) x 506 columns (species). Data are counts, i.e., the number of times a species was observed in a grid cell.

References

Mishler, B., Knerr, N., González-Orozco, C. et al. Phylogenetic measures of biodiversity and neo- and paleo-endemism in Australian *Acacia*. *Nat Commun* 5, 4473 (2014). doi:10.1038/ncomms5473

Examples

```
# Example phylogeny
acacia$phy
# Example community
acacia$comm[1:5, 1:5]
```

biod_example	<i>Biodiverse example data</i>
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Description

Fictional data for testing purposes from **Biodiverse**.

Usage

```
biod_example
```

Format

A list with two elements:

phy Phylogeny with 31 tips

comm Community matrix with 127 sites and 31 species. Data are counts, i.e., the number of times a species was observed in a grid cell.

Details

Corresponds to the the community matrix ("groups" object) and phylogeny from the Biodiverse **example_project.bps**. # nolint

References

Laffan, S.W., Lubarsky, E. & Rosauer, D.F. (2010) Biodiverse, a tool for the spatial analysis of biological and related diversity. *Ecography*. Vol 33, 643-647 (Version 3.1). doi:10.1111/j.1600-0587.2010.06237.x

Examples

```
# Example phylogeny
biod_example$phy
# Example community
biod_example$comm[1:5, 1:5]
```

biod_results	<i>Output from Biodiverse</i>
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Description

Output of analyzing test data with **Biodiverse**.

Usage

```
biod_results
```

Format

A tibble (dataframe) with 127 rows and 7 columns. Columns include:

site Site name; corresponds to row names of `biod_example$comm`
pd_biodiv Phylogenetic diversity (PD; PD_P in Biodiverse)
pd_alt_biodiv Alternative PD (PD measured on tree with all branchlengths equal; PHYLO_RPD_NULL2 in Biodiverse)
rpd_biodiv Relative PD (PD divided by alternative PD; PHYLO_RPD_NULL2 in Biodiverse)
pe_biodiv Phylogenetic endemism (PE; PE_WE_P in Biodiverse)
pe_alt_biodiv Alternative PE (PE measured on tree with all branchlengths equal; PHYLO_RPE_NULL2 in Biodiverse)
rpe_biodiv Relative PE (PE divided by alternative PD; PHYLO_RPE2 in Biodiverse)

Details

The `example_project.bps` # nolint example data set was used as input, which corresponds to the `biod_example` dataset in this package.

For a description of all Biodiverse metrics, see the **Biodiverse wiki**. # nolint

References

Laffan, S.W., Lubarsky, E. & Rosauer, D.F. (2010) Biodiverse, a tool for the spatial analysis of biological and related diversity. *Ecography*. Vol 33, 643-647 (Version 3.1). doi:10.1111/j.1600-0587.2010.06237.x

Examples

```
biod_results
```

cpr_classify_endem *Classify phylogenetic endemism*

Description

Given the results of `cpr_rand_test()`, classifies phylogenetic endemism according to CANAPE scheme of Mishler et al. 2014.

Usage

```
cpr_classify_endem(df)
```

Arguments

df Input data frame. Must have the following columns:

- `pe_obs_p_upper`: Upper p -value comparing observed phylogenetic endemism to random values
- `pe_alt_obs_p_upper`: Upper p -value comparing observed phylogenetic endemism on alternate tree to random values
- `rpe_obs_p_upper`: Upper p -value comparing observed relative phylogenetic endemism to random values

Details

For a summary of the classification scheme, see: <http://biodiverse-analysis-software.blogspot.com/2014/11/canape-categorical-analysis-of-palaeo.html> # nolint

Value

Object of class `data.frame` with column `endem_type` (character) added. Values of `endem_type` type include `paleo` (paleoendemic), `neo` (neoendemic), `not_significant` (what it says), `mixed` (mixed endemism), and `super` (super-endemic; both `pe_obs` and `pe_obs_alt` are highly significant).

References

Mishler, B., Knerr, N., González-Orozco, C. et al. (2014) Phylogenetic measures of biodiversity and neo- and paleo-endemism in Australian Acacia. *Nat Commun*, 5: 4473. doi:10.1038/ncomms5473

Examples

```
set.seed(12345)
data(phylocom)
rand_test <- cpr_rand_test(
  phylocom$comm, phylocom$phy,
  null_model = "curveball", metrics = c("pe", "rpe"), n_reps = 10
)
cpr_classify_endem(rand_test)
```

cpr_classify_signif *Classify statistical significance*

Description

Given the results of `cpr_rand_test()`, classifies statistical significance of a biodiversity metric. The null hypothesis is that observed value does not lie in the extreme of the random values.

Usage

```
cpr_classify_signif(df, metric, one_sided = FALSE, upper = FALSE)
```

Arguments

df	Input data frame.
metric	Character vector of length 1; selected metric to classify significance. May choose from pd (phylogenetic diversity), rpd (relative phylogenetic diversity), pe (phylogenetic endemism), rpe (relative phylogenetic endemism) (case-sensitive).
one_sided	Logical vector of length 1; is the null hypothesis one-sided? If TRUE, values will be classified as significant if they are in either the top 5% or bottom 5%. If FALSE, values will be classified as significant if they are in the top 2.5% or bottom 2.5%, combined.
upper	Logical vector of length 1; only applies if one_sided is TRUE. If TRUE, values in the top 5% will be classified as significant. If FALSE, values in the bottom 5% will be classified as significant.

Details

For metrics like pe, you probably want to consider a one-sided hypothesis testing values in the upper extreme (i.e., we are interested in areas that have higher than expected endemism). For this, you would set `one_sided = TRUE`, `upper = TRUE`. For metrics like pd, you probably want to consider a two-sided hypothesis (i.e., we are interested in areas that are either more diverse or less than diverse than expected at random). For this, set `one_sided = FALSE`.

Value

Object of class `data.frame` with column added for statistical significance of the selected metric. The new column name is the name of the metric with `_signif` appended. The new column is a character that may contain the following values, depending on the null hypothesis:

- `< 0.01, < 0.025, > 0.975, > 0.99`, not significant (two-sided)
- `< 0.01, < 0.05, > 0.99, > 0.95`, not significant (one-sided)

Examples

```
set.seed(12345)
data(phylocom)
rand_test <- cpr_rand_test(
  phylocom$comm, phylocom$phy,
  null_model = "curveball", metrics = "pd", n_reps = 50
)
cpr_classify_signif(rand_test, "pd")
```

cpr_endem_cols

CVD-friendly color palette for plotting results of CANAPE

Description

Character vector with names corresponding to endemism types and values corresponding to color codes. Should be distinguishable to people with color vision deficiency (CVD).

Usage

```
cpr_endem_cols
```

Format

An object of class character of length 5.

Details

Color scheme:

- paleo: dark blue
- neo: red
- not significant: beige
- mixed: green
- super: yellow

See Also

Other colors: [cpr_endem_cols_2](#), [cpr_endem_cols_3](#), [cpr_endem_cols_4](#), [cpr_make_pal\(\)](#), [cpr_signif_cols_2](#), [cpr_signif_cols](#), [mishler_endem_cols](#), [mishler_signif_cols](#)

Examples

```
cpr_endem_cols
scales::show_col(cpr_endem_cols)
```

cpr_endem_cols_2	<i>CVD-friendly color palette for plotting results of CANAPE, version 2</i>
------------------	---

Description

Character vector with names corresponding to endemism types and values corresponding to color codes. Should be distinguishable to people with color vision deficiency (CVD).

Usage

```
cpr_endem_cols_2
```

Format

An object of class character of length 5.

Details

Color scheme:

- paleo: dark blue
- neo: red
- not significant: light grey
- mixed: green
- super: yellow

See Also

Other colors: [cpr_endem_cols_3](#), [cpr_endem_cols_4](#), [cpr_endem_cols](#), [cpr_make_pal\(\)](#), [cpr_signif_cols_2](#), [cpr_signif_cols](#), [mishler_endem_cols](#), [mishler_signif_cols](#)

Examples

```
cpr_endem_cols_2  
scales::show_col(cpr_endem_cols_2)
```

cpr_endem_cols_3	<i>CVD-friendly color palette for plotting results of CANAPE, version 3</i>
------------------	---

Description

Character vector with names corresponding to endemism types and values corresponding to color codes. Should be distinguishable to people with color vision deficiency (CVD).

Usage

```
cpr_endem_cols_3
```

Format

An object of class character of length 5.

Details

Color scheme:

- paleo: dark blue
- neo: red
- not significant: light grey
- mixed: green
- super: orange

See Also

Other colors: [cpr_endem_cols_2](#), [cpr_endem_cols_4](#), [cpr_endem_cols](#), [cpr_make_pal\(\)](#), [cpr_signif_cols_2](#), [cpr_signif_cols](#), [mishler_endem_cols](#), [mishler_signif_cols](#)

Examples

```
cpr_endem_cols_3  
scales::show_col(cpr_endem_cols_3)
```

cpr_endem_cols_4	<i>CVD-friendly color palette for plotting results of CANAPE, version 4</i>
------------------	---

Description

Character vector with names corresponding to endemism types and values corresponding to color codes. Should be distinguishable to people with color vision deficiency (CVD).

Usage

```
cpr_endem_cols_4
```

Format

An object of class character of length 5.

Details

Color scheme:

- paleo: light blue
- neo: red
- not significant: light grey
- mixed: green
- super: yellow

See Also

Other colors: [cpr_endem_cols_2](#), [cpr_endem_cols_3](#), [cpr_endem_cols](#), [cpr_make_pal\(\)](#), [cpr_signif_cols_2](#), [cpr_signif_cols](#), [mishler_endem_cols](#), [mishler_signif_cols](#)

Examples

```
cpr_endem_cols_4  
scales::show_col(cpr_endem_cols_4)
```

cpr_iter_sim	<i>Simulate the number of iterations needed to generate a random community that is sufficiently different from the original community</i>
--------------	---

Description

For randomization algorithms that involve swapping (iterations), there is no way to know *a-priori* how many iterations are needed to sufficiently "mix" the community data matrix. [cpr_iter_sim\(\)](#) records the percentage similarity between the original matrix and a matrix that has been randomized with successive swapping iterations, at each iteration.

Usage

```
cpr_iter_sim(
  comm,
  null_model = "curveball",
  n_iterations = 100,
  thin = 1,
  seed = NULL
)
```

Arguments

comm	Dataframe or matrix; input community data with sites (communities) as rows and species as columns. Values of each cell are the presence/absence (0 or 1) or number of individuals (abundance) of each species in each site.
null_model	Character vector of length 1 or object of class <code>commsim</code> ; either the name of the model to use for generating random communities (null model), or a custom null model. For full list of available predefined null models, see the help file of vegan::commsim() , or run vegan::make.commsim() . An object of class <code>commsim</code> can be generated with vegan::commsim() .
n_iterations	Numeric vector of length 1; maximum number of iterations to conduct.
thin	Numeric vector of length 1; frequency to record percentage similarity between original matrix and randomized matrix. Results will be recorded every <code>thin</code> iterations (see Details).
seed	Integer vector of length 1 or <code>NULL</code> ; random seed that will be used in a call to <code>set.seed()</code> before randomizing the matrix. Default (<code>NULL</code>) will not change the random generator state.

Details

The user should inspect the results to determine at what number of iterations the original matrix and randomized matrix reach maximum dissimilarity (see Examples). This number will strongly depend on the size and structure of the original matrix. Large matrices with many zeros will likely take more iterations, and even then still retain relatively high similarity between the original matrix and the randomized matrix.

Available memory may be quickly exhausted if many (e.g., tens or hundreds of thousands, or more) of iterations are used with no thinning on large matrices; use `thin` to only record a portion of the results and save on memory.

Of course, `cpr_iter_sim()` only makes sense for randomization algorithms that use iterations.

Only presence/absence information is used to calculate percentage similarity between community matrices.

Value

Tibble (dataframe) with the following columns:

- `iteration`: Number of iterations used to generate random community
- `similarity`: Percentage similarity between original community and random community

Examples

```
# Simulate generation of a random community with maximum of 10,000
# iterations, recording similarity every 100 iterations
(res <- cpr_iter_sim(
  comm = biod_example$comm,
  null_model = "swap",
  n_iterations = 10000,
  thin = 100,
  seed = 123
))

# Plot reveals that ca. 1000 iterations are sufficient to
# completely mix random community
plot(res$iteration, res$similarity, type = "l")
```

cpr_make_pal

Make a palette of colors for plotting CANAPE results

Description

Palettes can also be loaded by name directly (see "Other colors" below).

Usage

```
cpr_make_pal(name, type)
```

Arguments

<code>name</code>	Character vector of length 1; name of palette to select. Must be one of 'mishler2014', 'canaper1', 'canaper2', 'canaper3', or 'canaper4'
<code>type</code>	Character vector of length 1; type of palette to select. Must be one of 'endem' (endemism) or 'signif' (<i>p</i> -rank significance)

Details

Palettes with the name 'mishler2014' correspond to colors used in Mishler et al. 2014. This color scheme has been widely used to show CANAPE results, but some colors may not be distinguishable to people with color vision deficiency (CVD).

Palettes with the name 'canaper1' or 'canaper2' are based on the Okabe-Ito palette (endemism; Okabe & Ito 2002) or RColorBrewer (significance) and should be CVD-friendly.

Names of colors correspond to either endemism type or p -rank output by `cpr_classify_endem()` or `cpr_classify_signif()`, respectively. Not all names are all available for both types.

Value

Named character vector; color hex codes

References

Mishler, B., Knerr, N., González-Orozco, C. et al. (2014) Phylogenetic measures of biodiversity and neo- and paleo-endemism in Australian Acacia. Nat Commun, 5: 4473. doi:10.1038/ncomms5473

Okabe, M., & Ito, K. (2002) Color Universal Design (CUD) - How to make figures and presentations that are friendly to Colorblind people. J*FLY Data Depository for Drosophila Researchers. <https://jfly.uni-koeln.de/color/>

See Also

Other colors: `cpr_endem_cols_2`, `cpr_endem_cols_3`, `cpr_endem_cols_4`, `cpr_endem_cols`, `cpr_signif_cols_2`, `cpr_signif_cols`, `mishler_endem_cols`, `mishler_signif_cols`

Examples

```
cpr_make_pal("mishler2014", "endem")
cpr_make_pal("canaper1", "endem")
scales::show_col(cpr_make_pal("mishler2014", "endem"))
```

cpr_rand_comm

Randomize a single community matrix

Description

Note that binary null models return a binary matrix, even if an abundance matrix was used as input.

Usage

```
cpr_rand_comm(comm, null_model, n_iterations = 1, thin = 1, seed = NULL)
```

Arguments

<code>comm</code>	Dataframe or matrix; input community data with sites (communities) as rows and species as columns. Values of each cell are the presence/absence (0 or 1) or number of individuals (abundance) of each species in each site.
<code>null_model</code>	Character vector of length 1 or object of class <code>commsim</code> ; either the name of the model to use for generating random communities (null model), or a custom null model. For full list of available predefined null models, see the help file of <code>vegan::commsim()</code> , or run <code>vegan::make.commsim()</code> . An object of class <code>commsim</code> can be generated with <code>vegan::commsim()</code> (see Examples).
<code>n_iterations</code>	Numeric vector of length 1; number of iterations for sequential null models. Ignored by non-sequential null models.
<code>thin</code>	Numeric vector of length 1; thinning parameter used by some null models in <code>vegan</code> (e.g., <code>quasiswap</code>); ignored for other models.
<code>seed</code>	Integer vector of length 1 or <code>NULL</code> ; random seed that will be used in a call to <code>set.seed()</code> before randomizing the matrix. Default (<code>NULL</code>) will not change the random generator state.

Value

Matrix

Examples

```
set.seed(12345)

# Check list of available pre-defined null models in vegan
vegan::make.commsim()

# Binary null model produces binary output
data(phylocom)
cpr_rand_comm(phylocom$comm, "swap", 100)

# Quantitative null model produces quantitative output
cpr_rand_comm(phylocom$comm, "swap_count", 100)

# How to use a custom null model
# 1. Define a randomizing function, e.g. re-sample the matrix while
# preserving total number of presences (same as the "r00" model)
randomizer <- function(x, n, ...) {
  array(replicate(n, sample(x)), c(dim(x), n))
}

# 2. Generate a commsim object
cs_object <- vegan::commsim(
  "r00_model",
  fun = randomizer, binary = TRUE,
  isSeq = FALSE, mode = "integer"
)
```

```
# 3. Generate the null community
cpr_rand_comm(phylocom$comm, cs_object, 100)
```

cpr_rand_test

Run a randomization analysis for one or more biodiversity metrics

Description

The observed value of the biodiversity metric(s) will be calculated for the input community data, then compared against a set of random communities. Various statistics are calculated from the comparison (see **Value** below).

Usage

```
cpr_rand_test(
  comm,
  phy,
  null_model,
  n_reps = 100,
  n_iterations = 10000,
  thin = 1,
  metrics = c("pd", "rpd", "pe", "rpe"),
  site_col = "site",
  tbl_out = tibble::is_tibble(comm),
  quiet = FALSE
)
```

Arguments

comm	Dataframe, tibble, or matrix; input community data with sites (communities) as rows and species as columns. Either presence-absence data (values only 0s or 1s) or abundance data (values ≥ 0) accepted, but calculations do not use abundance-weighting, so results from abundance data will be the same as if converted to presence-absence before analysis.
phy	List of class phylo; input phylogeny.
null_model	Character vector of length 1 or object of class commsim; either the name of the model to use for generating random communities (null model), or a custom null model. For full list of available predefined null models, see the help file of <code>vegan::commsim()</code> , or run <code>vegan::make.commsim()</code> . An object of class commsim can be generated with <code>vegan::commsim()</code> (see Examples in <code>cpr_rand_comm()</code>).
n_reps	Numeric vector of length 1; number of random communities to replicate.
n_iterations	Numeric vector of length 1; number of iterations to use for sequential null models; ignored for non-sequential models.
thin	Numeric vector of length 1; thinning parameter used by some null models in vegan (e.g., quasibswap); ignored for other models.

<code>metrics</code>	Character vector; names of biodiversity metrics to calculate. May include one or more of: <code>pd</code> , <code>rpd</code> , <code>pe</code> , <code>rpe</code> (case-sensitive).
<code>site_col</code>	Character vector of length 1; name of column in <code>comm</code> that contains the site names; only used if <code>comm</code> is a tibble (object of class <code>tbl_df</code>).
<code>tbl_out</code>	Logical vector of length 1; should the output be returned as a tibble? If <code>FALSE</code> , will return a dataframe. Defaults to <code>TRUE</code> if <code>comm</code> is a tibble.
<code>quiet</code>	Logical vector of length 1; if <code>TRUE</code> , suppress all warnings and messages that would be emitted by this function.

Details

The biodiversity metrics (`metrics`) available for analysis include:

- `pd`: Phylogenetic diversity (Faith 1992)
- `rpd`: Relative phylogenetic diversity (Mishler et al 2014)
- `pe`: Phylogenetic endemism (Rosauer et al 2009)
- `rpe`: Relative phylogenetic endemism (Mishler et al 2014)

(`pe` and `rpe` are needed for CANAPE with `cpr_classify_endem()`)

The choice of a randomization algorithm (`null_model`) is not trivial, and may strongly affect results. `cpr_rand_test()` uses null models provided by `vegan`; for a complete list, see the help file of `vegan::commsim()` or run `vegan::make.commsim()`. One frequently used null model is swap (Gotelli & Entsminger 2003), which randomizes the community matrix while preserving column and row sums (marginal sums). For a review of various null models, see Strona et al. (2018); swap is an "FF" model in the sense of Strona et al. (2018).

Instead of using one of the predefined null models in `vegan::commsim()`, it is also possible to define a custom null model; see **Examples** in `cpr_rand_comm()`

Note that the pre-defined models in `vegan` include binary models (designed for presence-absence data) and quantitative models (designed for abundance data). Although the binary models will accept abundance data, they treat it as binary and always return a binary (presence-absence) matrix. The PD and PE calculations in `canaper` are not abundance-weighted, so they return the same result regardless of whether the input is presence-absence or abundance. In that sense, binary null models are appropriate for `cpr_rand_test()`. The quantitative models could also be used for abundance data, but the output will be treated as binary anyways when calculating PD and PE. The effects of using binary vs. quantitative null models for `cpr_rand_test()` have not been investigated.

A minimum of 5 species and sites are required as input; fewer than that is likely cause the some randomization algorithms (e.g., swap) to enter an infinite loop. Besides, inferences on very small numbers of species and/or sites is not recommended generally.

The following rules apply to `comm` input:

- If dataframe or matrix, must include row names (site names) and column names (species names).
- If tibble, a single column (default, `site`) must be included with site names, and other columns must correspond to species names.
- Column names cannot start with a number and must be unique.
- Row names (site names) must be unique.

- Values (other than site names) should only include integers ≥ 0 ; non-integer input will be converted to integer.

The results are identical regardless of whether the input for comm is abundance or presence-absence data (i.e., abundance weighting is not used).

Value

Dataframe. For each of the biodiversity metrics, the following 9 columns will be produced:

- *_obs: Observed value
- *_obs_c_lower: Count of times observed value was lower than random values
- *_obs_c_upper: Count of times observed value was higher than random values
- *_obs_p_lower: Percentage of times observed value was lower than random values
- *_obs_p_upper: Percentage of times observed value was higher than random values
- *_obs_q: Count of the non-NA random values used for comparison
- *_obs_z: Standard effect size (z-score)
- *_rand_mean: Mean of the random values
- *_rand_sd: Standard deviation of the random values

So if you included pd in metrics, the output columns would include pd_obs, pd_obs_c_lower, etc...

References

- Faith DP (1992) Conservation evaluation and phylogenetic diversity. *Biological Conservation*, 61:1–10. doi:[10.1016/00063207\(92\)912013](https://doi.org/10.1016/00063207(92)912013)
- Gotelli, N.J. and Entsminger, N.J. (2003). Swap algorithms in null model analysis. *Ecology* 84, 532–535.
- Mishler, B., Knerr, N., González-Orozco, C. et al. (2014) Phylogenetic measures of biodiversity and neo- and paleo-endemism in Australian Acacia. *Nat Commun*, 5: 4473. doi:[10.1038/ncomms5473](https://doi.org/10.1038/ncomms5473)
- Rosauer, D., Laffan, S.W., Crisp, M.D., Donnellan, S.C. and Cook, L.G. (2009) Phylogenetic endemism: a new approach for identifying geographical concentrations of evolutionary history. *Molecular Ecology*, 18: 4061–4072. doi:[10.1111/j.1365294X.2009.04311.x](https://doi.org/10.1111/j.1365294X.2009.04311.x)
- Strona, G., Ulrich, W. and Gotelli, N.J. (2018), Bi-dimensional null model analysis of presence-absence binary matrices. *Ecology*, 99: 103–115. doi:[10.1002/ecy.2043](https://doi.org/10.1002/ecy.2043)

Examples

```
set.seed(12345)
data(phylocom)
# Returns a dataframe by default
cpr_rand_test(
  phylocom$comm, phylocom$phy,
  null_model = "curveball", metrics = "pd", n_reps = 10
)

# Tibbles may be preferable because of the large number of columns
cpr_rand_test(
```

```
phylocom$comm, phylocom$phy,  
null_model = "curveball", tbl_out = TRUE, n_reps = 10  
)
```

cpr_signif_cols	<i>CVD-friendly color palette for plotting results of randomization test</i>
-----------------	--

Description

Character vector with names corresponding to significance levels and values corresponding to color codes.

Usage

```
cpr_signif_cols
```

Format

An object of class character of length 5.

See Also

Other colors: [cpr_endem_cols_2](#), [cpr_endem_cols_3](#), [cpr_endem_cols_4](#), [cpr_endem_cols](#), [cpr_make_pal\(\)](#), [cpr_signif_cols_2](#), [mishler_endem_cols](#), [mishler_signif_cols](#)

Examples

```
cpr_signif_cols  
scales::show_col(cpr_signif_cols)
```

cpr_signif_cols_2	<i>CVD-friendly color palette for plotting results of randomization test, version 2</i>
-------------------	---

Description

Character vector with names corresponding to significance levels and values corresponding to color codes, with "not significant" colored grey.

Usage

```
cpr_signif_cols_2
```

Format

An object of class character of length 5.

See Also

Other colors: [cpr_endem_cols_2](#), [cpr_endem_cols_3](#), [cpr_endem_cols_4](#), [cpr_endem_cols](#), [cpr_make_pal\(\)](#), [cpr_signif_cols](#), [mishler_endem_cols](#), [mishler_signif_cols](#)

Examples

```
cpr_signif_cols_2
scales::show_col(cpr_signif_cols_2)
```

mishler_endem_cols	<i>Original color palette for plotting results of CANAPE</i>
--------------------	--

Description

Character vector with names corresponding to endemism types and values corresponding to color codes. Original palette used by Mishler et al. (2014). May not be distinguishable to people with color vision deficiency (CVD).

Usage

```
mishler_endem_cols
```

Format

An object of class character of length 5.

Details

Color scheme:

- paleo: blue
- neo: red
- not significant: beige
- mixed: light purple
- super: dark purple

References

Mishler, B., Knerr, N., González-Orozco, C. et al. (2014) Phylogenetic measures of biodiversity and neo- and paleo-endemism in Australian Acacia. Nat Commun, 5: 4473. doi:[10.1038/ncomms5473](#)

See Also

Other colors: [cpr_endem_cols_2](#), [cpr_endem_cols_3](#), [cpr_endem_cols_4](#), [cpr_endem_cols](#), [cpr_make_pal\(\)](#), [cpr_signif_cols_2](#), [cpr_signif_cols](#), [mishler_signif_cols](#)

Examples

```
mishler_endem_cols  
scales::show_col(mishler_endem_cols)
```

mishler_signif_cols	<i>Original color palette for plotting results of CANAPE</i>
---------------------	--

Description

Character vector with names corresponding to endemism types and values corresponding to color codes. Original palette used by Mishler et al. (2014). May not be distinguishable to people with color vision deficiency (CVD).

Usage

```
mishler_signif_cols
```

Format

An object of class character of length 5.

References

Mishler, B., Knerr, N., González-Orozco, C. et al. (2014) Phylogenetic measures of biodiversity and neo- and paleo-endemism in Australian Acacia. Nat Commun, 5: 4473. doi:[10.1038/ncomms5473](https://doi.org/10.1038/ncomms5473)

See Also

Other colors: [cpr_endem_cols_2](#), [cpr_endem_cols_3](#), [cpr_endem_cols_4](#), [cpr_endem_cols](#), [cpr_make_pal\(\)](#), [cpr_signif_cols_2](#), [cpr_signif_cols](#), [mishler_endem_cols](#)

Examples

```
mishler_signif_cols  
scales::show_col(mishler_signif_cols)
```

phylocom

Phylocom example data

Description

Fictional data for testing purposes from **Phylocom** (Webb et al. 2008)

Usage

phylocom

Format

A list with three elements:

phylo Phylogeny with 32 tips

sample Community matrix with 6 sites (rows) and 25 species (columns).

traits Trait data; a data.frame with 32 species (rows) and 4 traits (columns)

Details

Obtained via the *picante* package (Kembel et al. 2010)

References

Webb, C.O., Ackerly, D.D., and Kembel, S.W. 2008. Phylocom: software for the analysis of phylogenetic community structure and trait evolution. Version 4.0.1. <http://www.phylodiversity.net/phylocom/>.

Kembel, et al. *Picante: R tools for integrating phylogenies and ecology*, *Bioinformatics* 26: 1463–1464
[doi:10.1093/bioinformatics/btq166](https://doi.org/10.1093/bioinformatics/btq166)

Examples

```
# Example phylogeny
phylocom$phy
# Example community
phylocom$comm
```

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