Package: funbiogeo (via r-universe)

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Type Package

Title Functional Biogeography Analyses

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Description Helps users with analyses in functional biogeogeography by loading and combining data, computing trait coverage, as well as computing functional diversity indices, drawing maps, correlating them with the environment, and upscaling assemblages.

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https://github.com/frbcesab/funbiogeo

BugReports https://github.com/frbcesab/funbiogeo/issues

License GPL (≥ 2)

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Contents

Contents

fb_aggregate_site_data 3
fb_count_sites_by_species
fb_count_species_by_site
fb_count_species_by_trait
fb_count_traits_by_species
fb_cwm
fb_filter_sites_by_species_coverage
fb_filter_sites_by_trait_coverage
fb_filter_species_by_site_coverage
fb_filter_species_by_trait_coverage 11
fb_filter_traits_by_species_coverage
fb_format_site_locations
fb_format_site_species
fb_format_species_categories
fb_format_species_traits
fb_get_all_trait_coverages_by_site
fb_get_environment
fb_get_trait_combination_coverage 18
fb_get_trait_coverage_by_site 20
fb_make_report
fb_map_raster
fb_map_site_data
fb_map_site_traits_completeness
fb_plot_distribution_site_trait_coverage
fb_plot_number_sites_by_species
fb_plot_number_species_by_trait
fb_plot_number_traits_by_species
fb_plot_site_environment
fb_plot_site_traits_completeness
fb_plot_species_traits_completeness
fb_plot_species_traits_missingness
fb_plot_trait_combination_frequencies
fb_plot_trait_correlation
fb_table_trait_summary
site_locations
site_species
species_categories
species_traits

Index

fb_aggregate_site_data

Aggregate Site Data Along Coarser Spatial Grid

Description

This function helps aggregating site data along a coarser grid. The idea is that you have any type of data at the site scale (diversity metrics, environmental data, etc.) but you would like to work on it at a coarser scale, or you want to visualize it at that scale. This function helps you do exactly that.

Usage

```
fb_aggregate_site_data(site_locations, site_data, agg_grid, fun = mean)
```

Arguments

site_locations	an sf object with the spatial geometries of sites. NOTE : the first column should be named "site" and indicate site names.
site_data	a matrix or data.frame containing values per sites to aggregate along the pro- vided grid. Can have one or several columns (variables to aggregate). The first column must contain sites names as provided in the example dataset site_locations.
agg_grid	a SpatRaster object (package terra). A raster of one single layer, that defines the grid along which to aggregate.
fun	the function used to aggregate points values when there are multiple points in one cell. Default is mean.

Value

A SpatRaster object with as many layers as columns in site_data.

```
library("funbiogeo")
```

```
data("site_locations")
data("site_species")
## Import grid ----
tavg <- system.file("extdata", "annual_mean_temp.tif", package = "funbiogeo")
tavg <- terra::rast(tavg)
## Rasterize 3 first species counts ----
fb_aggregate_site_data(site_locations, site_species[, 1:4], tavg, fun = sum)</pre>
```

fb_count_sites_by_species

Count Number of Sites Occupied by Species

Description

For each species computes the percentage of sites where the species is present (distribution value higher than 0 and non-NA).

Usage

fb_count_sites_by_species(site_species)

Arguments

```
site_species a data.frame with sites in rows and species in columns. NOTE: the first col-
umn should be named "site" and indicate site names. The other columns
should be named according to species names.
```

Value

A three-column data.frame with:

- species: the name of the species;
- n_sites: the number of sites where the species is present;
- coverage: the percentage of sites where the species is present.

Examples

```
library("funbiogeo")
```

```
data("site_species")
```

```
site_coverage_by_species <- fb_count_sites_by_species(site_species)
head(site_coverage_by_species)</pre>
```

fb_count_species_by_site

Count Number of Species per Site

Description

For each site computes the proportion of species present (distribution value higher than 0 and non-NA) compared to all species provided. For example, a site could contain only 20% of all species provided.

fb_count_species_by_trait

Usage

fb_count_species_by_site(site_species)

Arguments

```
site_species a data.frame with sites in rows and species in columns. NOTE: the first col-
umn should be named "site" and indicate site names. The other columns
should be named according to species names.
```

Value

A three-column data.frame with:

- site: the name of the site;
- n_species: the number of present species;
- coverage: the percentage of present species.

Examples

library("funbiogeo")

data("site_species")

species_coverage_by_site <- fb_count_species_by_site(site_species)
head(species_coverage_by_site)</pre>

fb_count_species_by_trait

Count Number of Species for Each Trait

Description

For each trait computes the percentage of species without NA (missing trait values).

Usage

```
fb_count_species_by_trait(species_traits)
```

Arguments

species_traits a data.frame with species in rows and traits as columns. **NOTE**: The first column should be named "species" and contain species names. The other columns should be named according to trait names.

Value

A three-column data.frame with:

- trait: the name of the trait;
- n_species: the number of species with non-missing value for the trait;
- coverage: the percentage of species with non-missing value for the trait.

Examples

library("funbiogeo")

```
data("species_traits")
```

```
species_coverage_by_trait <- fb_count_species_by_trait(species_traits)
head(species_coverage_by_trait)</pre>
```

fb_count_traits_by_species

Compute Number of Known Trait(s) per Species

Description

For each species computes the percentage of traits without NA (missing trait values).

Usage

```
fb_count_traits_by_species(species_traits)
```

Arguments

species_traits a data.frame with species in rows and traits as columns. **NOTE**: The first column should be named "species" and contain species names. The other columns should be named according to trait names.

Value

A three-column data.frame with:

- species: the name of the species;
- n_traits: the number of traits with non-missing value for the species;
- coverage: the percentage of traits with non-missing value for the species.

fb_cwm

Examples

library("funbiogeo")

```
data("species_traits")
```

```
trait_coverage_by_species <- fb_count_traits_by_species(species_traits)
head(trait_coverage_by_species)</pre>
```

```
fb_cwm
```

Compute community-weighted means (CWM) of trait values

Description

This function returns the community-weighted mean of provided trait values. It only works with quantitative traits and will warn you otherwise. It will remove species that either have NA values in the site_species input or NA values as their trait.

Usage

fb_cwm(site_species, species_traits)

Arguments

site_species	a data.frame with sites in rows and species in columns. NOTE: the first col-
	umn should be named "site" and indicate site names. The other columns
	should be named according to species names.
<pre>species_traits</pre>	a data.frame with species in rows and traits as columns. $\ensuremath{\textbf{NOTE}}$: The first
	column should be named "species" and contain species names. The other
	columns should be named according to trait names.

Value

A data.frame with sites in rows and the following variables:

- site, the site label,
- trait, the trait label as provided in species_traits,
- and cwm, the community-weighted means of quantitative traits values.

```
library("funbiogeo")
```

```
data("site_species")
data("species_traits")
site_cwm <- fb_cwm(head(site_species), species_traits)</pre>
```

```
head(site_cwm)
```

```
fb_filter_sites_by_species_coverage
```

Filter sites with a given species coverage threshold

Description

Selects sites (rows) for which the percentage of present species (distribution value higher than 0 and non-NA) is higher than a threshold.

Usage

```
fb_filter_sites_by_species_coverage(
   site_species,
   threshold_species_proportion = 0
)
```

Arguments

site_species	a data.frame with sites in rows and species in columns. NOTE: the first col-
	umn should be named "site" and indicate site names. The other columns
	should be named according to species names.

threshold_species_proportion

a numeric of length 1 between 0 and 1. The percentage of species coverage threshold.

Value

A subset of site_species with sites covered by X% of species.

Examples

```
library("funbiogeo")
data("site_species")
# Get sites with more than 40% of the species
new_site_species <- fb_filter_sites_by_species_coverage(
    site_species,
    threshold_species_proportion = 0.4
)</pre>
```

There are now only 148 sites
dim(new_site_species)
new_site_species[1:3, 1:4]

Description

•••

Usage

```
fb_filter_sites_by_trait_coverage(
    site_species,
    species_traits,
    threshold_traits_proportion = 1
)
```

Arguments

site_species	a data.frame with sites in rows and species in columns. NOTE : the first column should be named "site" and indicate site names. The other columns should be named according to species names.
species_traits	a data.frame with species in rows and traits as columns. NOTE : The first column should be named "species" and contain species names. The other columns should be named according to trait names.
threshold_traits_proportion	
	a numeric of length 1 between 0 and 1. The percentage trait coverage threshold

Value

A subset of site_species with sites covered by X% of abundance/coverage considering all provided traits.

Examples

```
library("funbiogeo")
```

```
data("site_species")
data("species_traits")
```

```
# Filter all the sites where all species have known traits
new_site_species <- fb_filter_sites_by_trait_coverage(
    site_species, species_traits
)
# There is only one such site
nrow(new_site_species)
```

Filter sites where at least 80% of species have known traits

```
new_site_species_2 <- fb_filter_sites_by_trait_coverage(
    site_species, species_traits, threshold_traits_proportion = 0.8
)
# There are now four sites
nrow(new_site_species_2)</pre>
```

fb_filter_species_by_site_coverage

Filter species with a given sites coverage threshold

Description

Selects species (columns) for which the percentage of sites where the species is present (distribution value higher than 0 and non-NA) is higher than a threshold.

Usage

```
fb_filter_species_by_site_coverage(
   site_species,
   threshold_sites_proportion = 0
)
```

Arguments

site_species	a data.frame with sites in rows and species in columns. NOTE: the first col-
	umn should be named "site" and indicate site names. The other columns
	should be named according to species names.
threshold_site	s_proportion
	a numeric of length 1 between 0 and 1. The percentage of sites coverage thresh-
	old.

Value

A subset of site_species with species with a prevalence higher than threshold_sites_proportion.

Examples

library("funbiogeo")

```
data("site_species")
# Filter species present in at least 10% of the sites
new_site_species <- fb_filter_species_by_site_coverage(
    site_species,
    threshold_sites_proportion = 0.1
)
new_site_species[1:3, 1:4]</pre>
```

10

```
# There are now only 84 species (instead of 149)
ncol(new_site_species)
```

fb_filter_species_by_trait_coverage

Filter species with a given traits coverage threshold

Description

Selects species (rows) for which the percentage of traits without NA (missing trait values) is higher than a threshold.

Usage

```
fb_filter_species_by_trait_coverage(
    species_traits,
    threshold_traits_proportion = 0
)
```

Arguments

species_traits a data.frame with species in rows and traits as columns. **NOTE**: The first column should be named "species" and contain species names. The other columns should be named according to trait names.

threshold_traits_proportion

a numeric of length 1 between 0 and 1. The percentage of traits coverage threshold.

Value

A subset of species_traits with species covered by X% of traits.

```
library("funbiogeo")
```

```
data("species_traits")
```

```
# Filter species that have at least 60% of the traits described
new_species_traits <- fb_filter_species_by_trait_coverage(
    species_traits,
    threshold_traits_proportion = 0.6
)
```

```
# There are now only 93 species
nrow(new_species_traits)
```

```
fb_filter_traits_by_species_coverage
```

Filter traits with a given species coverage threshold

Description

Selects traits (columns) for which the percentage of species without NA (missing trait values) is higher than a threshold.

Usage

```
fb_filter_traits_by_species_coverage(
    species_traits,
    threshold_species_proportion = 0
)
```

Arguments

species_traits a data.frame with species in rows and traits as columns. **NOTE**: The first column should be named "species" and contain species names. The other columns should be named according to trait names.

threshold_species_proportion

numeric(1) [default = NULL]
between 0 and 1. The percentage of species coverage threshold.

Value

A subset of species_traits with traits for the specified proportion of species.

```
library("funbiogeo")
data("species_traits")
# Filter traits that have at least 60% non-missing values
new_species_traits <- fb_filter_traits_by_species_coverage(
    species_traits,
    threshold_species_proportion = 0.6
)
# There are now only 2 traits
ncol(new_species_traits)</pre>
```

fb_format_site_locations

Extract site x locations information from long format data

Description

Convert a flat data.frame with site coordinates into a proper sf object that can then be used by other functions. This function assumes that the coordinates are given in WGS84 (longitude vs. latitude). The function automatically removes repeated coordinates from the input dataset.

Usage

```
fb_format_site_locations(
    data,
    site,
    longitude,
    latitude,
    crs = sf::st_crs(4326),
    na_rm = FALSE
)
```

Arguments

data	a data.frame in a long format (see example).
site	a character of length 1. Name of the column with site labels.
longitude	a character of length 1. Name of the column with longitude. The function assumes coordinates are WGS84 (EPSG:4326).
latitude	a character of length 1. Name of the column with latitude. The function assumes coordinates are WGS84 (EPSG:4326).
crs	a character of length 1 or an object of class crs. Coordinate Reference System (CRS) of the specified coordinates. The CRS should be a valid CRS in R. It can either be a character like "+proj=longlat +datum=WGS84 +no_defs" or as specified using sf::st_crs() like sf::st_crs(4326) the default value.
na_rm	a logical value. If TRUE remove sites with incomplete coordinates. Default is FALSE.

Value

An sf object with a site column specifying site coordinates.

Examples

library("funbiogeo")

fb_format_site_species

Extract site x species information from long format data

Description

Convert a flat data.frame with species occurrence/abundance at site level into a proper data.frame object that can then be used by other functions. The final output contains sites in rows and species in columns.

Usage

```
fb_format_site_species(data, site, species, value, na_to_zero = TRUE)
```

Arguments

data	a data.frame in a long format (see example).
site	a character of length 1. Name of the column with site labels.
species	a character of length 1. Name of the column with species names.
value	a character of length 1. Name of the column with species occurrence/abundance.
na_to_zero	a logical value. If TRUE (default) NA are replaced by 0.

Value

A data.frame with sites in rows and species in columns. The first column is named "site" and contains the name of the sites.

Examples

```
library("funbiogeo")
```

14

fb_format_species_categories

Extract species x categories information from long format data

Description

Convert a flat data.frame with species names and species (supra-)category (e.g. family, order, endemism status, etc.) into a proper data.frame object that can then be used by other functions. The final output contains species in rows and two columns (species name and species category).

Usage

fb_format_species_categories(data, species, category)

Arguments

data	a data.frame in a long format (see example).
species	a character of length 1. Name of the column with species names.
category	a character of length 1. Name of the column with species category.

Value

A data.frame with species in rows and two columns (species name and species category).

fb_format_species_traits

Extract species x traits information from long format data

Description

Convert a flat data.frame with traits values for different species into a proper data.frame object that can then be used by other functions. The final output contains species in rows and traits in columns.

Usage

fb_format_species_traits(data, species, traits)

Arguments

data	a data.frame in a long format (see example).
species	a character of length 1. Name of the column with species names.
traits	a character of length >= 1. Name(s) of trait column(s).

Value

A data.frame with species in rows and traits in columns, with the first column names "species" containing the species names.

Description

Compute trait coverage for all sites, i.e., the percentage of total abundance/presence of species that have traits data compared to total species. This function assumes that all species provided in the traits dataset have all their traits specified (meaning that all species have either known or NA values reported as their traits). The coverage of each trait separately is returned as well as all traits taken together if wanted.

Usage

```
fb_get_all_trait_coverages_by_site(
   site_species,
   species_traits,
   all_traits = TRUE
)
```

Arguments

site_species	a data.frame with sites in rows and species in columns. NOTE : the first col- umn should be named "site" and indicate site names. The other columns should be named according to species names.
species_traits	a data.frame with species in rows and traits as columns. NOTE : The first column should be named "species" and contain species names. The other columns should be named according to trait names.
all_traits	a logical (default = TRUE) which tell if the coverage considering all provided traits should be provided in an additional column all_traits

Value

a data.frame with a column with sites and one column per provided trait giving its coverage (percent species per site, weighted by abundance that have trait data), and, when argument all_traits = TRUE, an additional column named all_traits considering the coverage of all traits taken together.

```
site_trait_cov <- fb_get_all_trait_coverages_by_site(
    site_species, species_traits
)
head(site_trait_cov)</pre>
```

fb_get_environment Extract Raster Values at Location of Sites

Description

Extract Raster Values at Location of Sites

raster.

Usage

```
fb_get_environment(site_locations, environment_raster)
```

Arguments

Value

A data.frame with average environmental values (columns) per site (rows), with the first column being "site" indicating site names.

Examples

```
library("funbiogeo")
data("site_locations")
## Import climate rasters ----
prec <- system.file("extdata", "annual_tot_prec.tif", package = "funbiogeo")
tavg <- system.file("extdata", "annual_mean_temp.tif", package = "funbiogeo")
layers <- terra::rast(c(tavg, prec))
fb_get_environment(head(site_locations), layers)</pre>
```

fb_get_trait_combination_coverage

Compute site trait coverage for each trait combination

Description

This function computes trait coverage for each site for different trait combinations. If not provided, consider all possible trait combinations. The function will not run if the total number of combinations given is over 10,000.

fb_get_trait_combination_coverage

Usage

```
fb_get_trait_combination_coverage(
   site_species,
   species_traits,
   comb_size = NULL
)
```

Arguments

site_species	a data.frame with sites in rows and species in columns. NOTE : the first col- umn should be named "site" and indicate site names. The other columns should be named according to species names.
species_traits	a data.frame with species in rows and traits as columns. NOTE : The first column should be named "species" and contain species names. The other columns should be named according to trait names.
comb_size	an integer vector defining one or more sizes of combinations (default: NULL)

Value

a data.frame with the following columns:

- site with the site indices from site_species,
- combination_length with the number of traits in given combinations,
- combination_name with the name of the trait combination (concatenated trait names with __),
- trait_coverage the corresponding trait coverage for the given trait combination and site.

```
# Compute Coverages using All Trait Combinations
all_combinations = fb_get_trait_combination_coverage(
site_species, species_traits
)
# Get only combinations of 3 traits
three_traits = fb_get_trait_combination_coverage(
site_species, species_traits, 3
)
# Combinations of 2, 3, and 4 traits
two_to_four = fb_get_trait_combination_coverage(
site_species, species_traits, c(2, 3, 4)
)
```

```
fb_get_trait_coverage_by_site
```

Compute Trait Coverage For Each Site Weighted by Abundance

Description

Compute trait coverage for all sites, i.e., the percentage of total abundance/presence of species that have traits data compared to total species. This function assumes that all species provided in the traits dataset have all their traits specified (meaning that all species have either known or NA values reported as their traits). **NB**: this function returns trait coverage using all traits provided in the input species_traits data.frame.

Usage

fb_get_trait_coverage_by_site(site_species, species_traits)

Arguments

site_species	a data.frame with sites in rows and species in columns. NOTE : the first column should be named "site" and indicate site names. The other columns should be named according to species names.
species_traits	a data.frame with species in rows and traits as columns. NOTE : The first column should be named "species" and contain species names. The other columns should be named according to trait names.

Value

A data.frame with n rows (where n is the number of sites) and two columns: site, the site label, and trait_coverage, the percent of total abundance/presence of species that have traits data.

Examples

```
library("funbiogeo")
```

```
data("site_species")
data("species_traits")
```

site_trait_cov <- fb_get_trait_coverage_by_site(site_species, species_traits)</pre>

```
head(site_trait_cov)
```

fb_make_report

Description

Creates an R Markdown (.Rmd) report from a template to explore and summarize users data. User can modify this report and use the function rmarkdown::render() to convert this .Rmd in different formats:

- HTML document (output_format = "bookdown::html_document2");
- PDF document (output_format = "bookdown::pdf_document2");
- Word document (output_format = "bookdown::word_document2");
- HTML, PDF and Word documents (output_format = "all").

Usage

```
fb_make_report(
   path = ".",
   filename = NULL,
   title = NULL,
   author = NULL,
   species_traits_name,
   site_species_name,
   site_locations_name,
   overwrite = FALSE,
   open = TRUE
)
```

Arguments

path	a character of length 1. The directory in which the .Rmd file will be created. This directory must exist.	
filename	a character of length 1. The name of the .Rmd file to be created. If NULL (default) the .Rmd file will be named funbiogeo_report.Rmd.	
title	a character of length 1. The title of the report. If NULL (default) the title will be funbiogeo Report.	
author	a character of length 1. The $author(s)$ of the report. If NULL (default) no author will be added.	
<pre>species_traits_name</pre>		
	a character of length 1. The name of the species x traits dataset (not the object). Note that before rendering the report this dataset must be loaded.	
site_species_name		
	a character of length 1. The name of the sites x species dataset (not the object). Note that before rendering the report this dataset must be loaded.	

site_locations_name		
	a character of length 1. The name of the sites x locations dataset (not the object). Note that before rendering the report this dataset must be loaded.	
overwrite	a logical. If this file is already present and overwrite = TRUE, it will be erased and replaced by the template. Default is FALSE.	
open	a logical. If TRUE (default), this file will be opened on the text editor.	

Value

No return value.

Examples

```
# Create temporary folder
temp_path <- tempdir()</pre>
# Load data
data("species_traits")
data("site_species")
data("site_locations")
# Create report
fb_make_report(
  path
                      = temp_path,
  author
                     = "Casajus N. and Grenié M.",
  species_traits_name = "species_traits",
  site_species_name = "site_species",
  site_locations_name = "site_locations",
  open = FALSE
)
## Not run:
# Open Rmd file ----
utils::file.edit(file.path(temp_path, "funbiogeo_report.Rmd"))
# Render Rmd file ----
rmarkdown::render(file.path(temp_path, "funbiogeo_report.Rmd"),
                  output_format = "all")
## End(Not run)
```

fb_map_raster Map a Single Raster Layer

Description

Map a Single Raster Layer

fb_map_site_data

Usage

fb_map_raster(x, ...)

Arguments

х	a SpatRaster object (package terra). A raster of one single layer
	other parameters passed to theme()

Value

A ggplot object.

Examples

```
library(ggplot2)
## Load raster ----
tavg <- system.file("extdata", "annual_mean_temp.tif", package = "funbiogeo")</pre>
tavg <- terra::rast(tavg)</pre>
## Default map ----
fb_map_raster(tavg)
## Map with custom theme ----
fb_map_raster(tavg, legend.position = "bottom")
## Advanced customization ----
my_map <- fb_map_raster(tavg) +</pre>
  scale_fill_distiller("Temperature", palette = "Spectral") +
  theme(legend.position = "bottom") +
  ggtitle("Mean annual temperature in Western Europe")
my_map
## Map w/o annotation ----
fb_map_raster(tavg) +
```

fb_map_site_data Map Arbitrary Site Data

theme(legend.position = "none")

Description

theme_void() +

From the site-locations data and a dataset organized by site, plot a map of this information. The returned plot is as little customized as possible to let the user do the customization.

```
fb_map_site_data(site_locations, site_data, selected_col)
```

site_locations	an sf object with the spatial geometries of sites. NOTE : the first column should be named "site" and indicate site names.
site_data	<pre>data.frame() of additional site information containing the column "site" to merge with the site_locations argument</pre>
selected_col	character(1) name of the column to plot

Value

a ggplot object.

Examples

site_rich <- fb_count_species_by_site(site_species)</pre>

```
# Map of Species Richness
rich_map <- fb_map_site_data(site_locations, site_rich, "n_species")
rich_map
# Customize the map
rich_map +
ggplot2::scale_fill_viridis_c("Species Richness")</pre>
```

Description

Returns a ggplot2 map of sites colored by trait coverage (proportion of species having a known trait value). By default shows one plot for each trait and add an additional facet named "all_traits" considering the trait coverage with all traits taken together.

```
fb_map_site_traits_completeness(
    site_locations,
    site_species,
    species_traits,
    all_traits = TRUE
)
```

site_locations	an sf object with the spatial geometries of sites. NOTE : the first column should be named "site" and indicate site names.
site_species	a data.frame with sites in rows and species in columns. NOTE : the first col- umn should be named "site" and indicate site names. The other columns should be named according to species names.
species_traits	a data.frame with species in rows and traits as columns. NOTE : The first column should be named "species" and contain species names. The other columns should be named according to trait names.
all_traits	a logical (default = TRUE) which tell if the coverage considering all provided traits should be provided in an additional column all_traits

Value

a 'ggplot2' object

Examples

fb_map_site_traits_completeness(site_locations, site_species, species_traits)

Description

Plots the distributions of trait coverage across site, i.e. the proportion of species weighted by abundance with known trait values, for each trait separately and all traits taken together. The trait distributions are ordered from the lowest to the highest average trait coverage (top to bottom). The top always displays a distribution named "all_traits" that contains the distribution of coverage all traits taken together.

```
fb_plot_distribution_site_trait_coverage(
    site_species,
    species_traits,
    species_categories = NULL,
    all_traits = TRUE
)
```

site_species	a data.frame with sites in rows and species in columns. NOTE : the first col- umn should be named "site" and indicate site names. The other columns should be named according to species names.	
species_traits	a data.frame with species in rows and traits as columns. NOTE : The first column should be named "species" and contain species names. The other columns should be named according to trait names.	
species_categories		
	(default = NULL) 2-columns data.frame giving species categories, with the first column describing the species name, and the second column giving their corresponding categories	
all_traits	a logical (default = TRUE) which tell if the coverage considering all provided traits should be provided in an additional column all_traits	

Value

a 'ggplot2' object

Examples

fb_plot_distribution_site_trait_coverage(site_species, species_traits)

Description

Represent all species in each function of the number of sites they occupy. The species are ordered from the ones that occupy the least number of sites from the ones that occupy the most. The number of site is indicated at the bottom x-axis, while the top x-axis represents the proportion of occupied sites. The left y-axis label species names and their rank by increasing prevalence. The user can supplied a threshold of sites to see how many species occupy more or less than the given proportion of sites.

```
fb_plot_number_sites_by_species(
    site_species,
    threshold_sites_proportion = NULL
)
```

site_species a data.frame with sites in rows and species in columns. **NOTE**: the first column should be named "site" and indicate site names. The other columns should be named according to species names. threshold_sites_proportion

a numeric of length 1 between 0 and 1. The percentage of sites coverage threshold.

Value

a ggplot2 object

Examples

fb_plot_number_sites_by_species(site_species)

```
# Add a vertical cutoff line (40% of sites)
fb_plot_number_sites_by_species(site_species, 0.4)
```

Description

Display a lollipop graph showing the number and proportion of species with non-NA trait for each trait ranked in decreasing order.

Usage

```
fb_plot_number_species_by_trait(
    species_traits,
    species_categories = NULL,
    threshold_species_proportion = NULL
)
```

Arguments

```
species_traits a data.frame with species in rows and traits as columns. NOTE: The first column should be named "species" and contain species names. The other columns should be named according to trait names.
```

```
species_categories
```

(default = NULL) 2-columns data.frame giving species categories, with the first column describing the species name, and the second column giving their corresponding categories

threshold_species_proportion

numeric(1) [default = NULL]

between 0 and 1. The percentage of species coverage threshold.

Value

a ggplot2 object

Examples

```
data(species_traits)
```

```
fb_plot_number_species_by_trait(species_traits)
```

```
# Add a vertical cutoff line (12.5% of species)
fb_plot_number_species_by_trait(species_traits, NULL, 1/8)
```

Description

Display a graph showing the number (and proportion) of species having at least 0, 1, etc. number of traits. It provides a lollipop graph to examine which number of trait cover a certain proportion of the species. This plot doesn't show which traits are concerned.

Usage

```
fb_plot_number_traits_by_species(
    species_traits,
    species_categories = NULL,
    threshold_species_proportion = NULL
)
```

Arguments

```
species_traits a data.frame with species in rows and traits as columns. NOTE: The first column should be named "species" and contain species names. The other columns should be named according to trait names.
```

species_categories

(default = NULL) 2-columns data.frame giving species categories, with the first column describing the species name, and the second column giving their corresponding categories

```
threshold_species_proportion
```

numeric(1) [default = NULL]
between 0 and 1. The percentage of species coverage threshold.

Value

a ggplot2 object

28

fb_plot_site_environment

Examples

```
data(species_traits)
fb_plot_number_traits_by_species(species_traits)
# Add a vertical cutoff line (33% of the species)
fb_plot_number_traits_by_species(
   species_traits, threshold_species_proportion = 1/3
)
```

fb_plot_site_environment

Plot Position of Sites in Environmental Space

Description

Plot a figure showing the average environmental space of given sites compared to a full environmental vector. For the sake of simplicity only represents the figure along two environmental axes. The average environmental value are extracted for each site.

Usage

```
fb_plot_site_environment(
   site_locations,
   environment_raster,
   first_layer = names(environment_raster)[1],
   second_layer = names(environment_raster)[2]
)
```

Arguments

site_locations	an sf object with the spatial geometries of sites. NOTE : the first column should be named "site" and indicate site names.
environment_ras	ster
	a SpatRaster object (package terra). A single or multi-layers environmental raster.
first_layer	character(1) the name of the first layer to use, by default uses the first layer of environment_raster
second_layer	character(1) the name of the second layer to use, by default uses the second layer of environment_raster

Value

a ggplot object

Examples

```
data("site_locations")
# Import climate rasters
prec <- system.file("extdata", "annual_tot_prec.tif", package = "funbiogeo")
tavg <- system.file("extdata", "annual_mean_temp.tif", package = "funbiogeo")
layers <- terra::rast(c(tavg, prec))
# Make plot (show environmental position of 6 first sites)
fb_plot_site_environment(head(site_locations), layers)</pre>
```

Description

Display a binary heatmap visualizing the site x traits matrix with colors displaying the proportion of occurring species with known trait values. Traits are ordered from the most to the least known (left to right). Sites are ordered from the ones with highest to lowest overall trait coverage (bottom to top). The site average proportion of species with known trait for each trait (across all sites) is shown in the x-axis labels. An additional column at the very right of the plot named "all_traits" shows a summary considering traits together.

Usage

```
fb_plot_site_traits_completeness(
    site_species,
    species_traits,
    species_categories = NULL,
    all_traits = TRUE
)
```

Arguments

site_species	a data.frame with sites in rows and species in columns. NOTE : the first column should be named "site" and indicate site names. The other columns should be named according to species names.
species_traits	a data.frame with species in rows and traits as columns. NOTE : The first column should be named "species" and contain species names. The other columns should be named according to trait names.
species_categor	ries
	(default = NULL) 2-columns data.frame giving species categories, with the first column describing the species name, and the second column giving their corresponding categories
all_traits	a logical (default = TRUE) which tell if the coverage considering all provided traits should be provided in an additional column all_traits

30

Value

a ggplot2 object

Examples

fb_plot_site_traits_completeness(site_species, species_traits)

fb_plot_species_traits_completeness

Plot Trait Coverage per Species for each Trait

Description

Display a binary heatmap visualizing the species x traits matrix with colors displaying present and missing traits. Traits are ordered from the most to the least known (left to right). Species are ordered from the ones with most to the ones with least traits (bottom to top). The proportion of species with non-missing traits is shown on the x-axis labels. An additional column at the very right of the plot named "all_traits" shows a summary considering if all other traits are known.

Usage

```
fb_plot_species_traits_completeness(
    species_traits,
    species_categories = NULL,
    all_traits = TRUE
)
```

Arguments

species_traits	a data.frame with species in rows and traits as columns. NOTE : The first column should be named "species" and contain species names. The other columns should be named according to trait names.	
species_categories		
	(default = NULL) 2-columns data.frame giving species categories, with the first column describing the species name, and the second column giving their corresponding categories	
all_traits	a logical (default = TRUE) which tell if the coverage considering all provided traits should be provided in an additional column all_traits	

Value

a ggplot2 object

Examples

```
data(species_traits)
fb_plot_species_traits_completeness(species_traits)
```

fb_plot_species_traits_missingness

Plot Trait Coverage per Species for each Trait

Description

Display a binary heatmap visualizing the species x traits matrix with colors displaying present and missing traits. Traits are ordered from the most to the least known (left to right). Species are ordered from the ones with most to the ones with least traits (bottom to top). The proportion of species with non-missing traits is shown on the x-axis labels. An additional column at the very right of the plot named "all_traits" shows a summary considering if all other traits are known.

Usage

```
fb_plot_species_traits_missingness(
    species_traits,
    species_categories = NULL,
    all_traits = TRUE
)
```

Arguments

species_traits	a data.frame with species in rows and traits as columns. NOTE : The first column should be named "species" and contain species names. The other columns should be named according to trait names.
species_categor	ries
	2-columns data.frame giving species categories NULL by default, with the first column describing the species name, and the second column giving their corresponding categories
all_traits	a logical (default = TRUE) which tell if the coverage considering all provided traits should be provided in an additional column all_traits

Value

a ggplot2 object

Examples

```
data(species_traits)
fb_plot_species_traits_missingness(species_traits)
```

32

fb_plot_trait_combination_frequencies

Plot Present and Missing Trait Combinations Frequencies

Description

Display a figure of Present/Missing Trait Combinations where one square is represents one trait. The y-axis gives the frequency of the row (as well as its proportions compared to the total number of rows).

Usage

```
fb_plot_trait_combination_frequencies(
   species_traits,
   species_categories = NULL,
   order_by = c("number", "complete")
)
```

Arguments

species_traits a data.frame with species in rows and traits as columns. **NOTE**: The first column should be named "species" and contain species names. The other columns should be named according to trait names.

species_categories

(default = NULL) 2-columns data.frame giving species categories, with the first column describing the species name, and the second column giving their corresponding categories

order_by character(1) either "number" or "complete If "number" order rows by frequency so that most frequent rows are at the bottom. Otherwise order rows to order table by the number of non-missing traits then by the frequency of combinations

Value

a ggplot2 object

Examples

fb_plot_trait_combination_frequencies(species_traits, NULL, "complete")

fb_plot_trait_correlation

Plot Trait Correlation Matrix

Description

This functions outputs a ggplot2 figure that outputs a trait correlation matrix. It only works on numerical traits and will output messages if non-numerical traits are found and errors if no numerical traits are found. Internally it uses the stats::cor() function and only works on complete observation (it removes any row that contains any NA). Use the ... argument to pass options to the cor() function.

Usage

```
fb_plot_trait_correlation(species_traits, species_categories = NULL, ...)
```

Arguments

<pre>species_traits</pre>	a data.frame with species in rows and traits as columns. NOTE: The first	
	column should be named "species" and contain species names. The other	
	columns should be named according to trait names.	
species_categories		
	$(default = \text{NULL}) \ 2\text{-}columns \ data. frame giving species categories, with the first column describing the species name, and the second column giving their corresponding categories$	
	Additional options passed to stats::cor()	

Value

a ggplot object

```
fb_plot_trait_correlation(species_traits)
```

```
# Plot Spearman's correlation
fb_plot_trait_correlation(species_traits, method = "spearman")
```

```
fb_table_trait_summary
```

Summary Table on Traits (Missingness, Range, etc.)

Description

This function outputs a data.frame that summarises the species by trait table to have many information in a glance. This can then return a data.frame or a nicely formatted knitr::kable() for inclusion in an Rmarkdown document.

Usage

```
fb_table_trait_summary(species_traits, kable = FALSE)
```

Arguments

<pre>species_traits</pre>	a data.frame with species in rows and traits as columns. NOTE: The first
	column should be named "species" and contain species names. The other
	columns should be named according to trait names.
kable	TRUE or FALSE Should function returns a knitr::kable()? defaults FALSE

Value

a data.frame with the following columns:

- trait_name: a character column with the trait name as indicated in species_traits
- trait_type: the nature of the trait (numeric, categorical, or ordered)
- number_non_missing: the total number of non-NA trait values
- proportion_non_missing: the proportion of non-NA trait values
- trait_range: for numerical traits, the range of values
- trait_mean_sd: for numerical traits, the mean plus-minus the standard deviation
- number_distinct: for non-numerical traits, the number of categories
- list_distinct: for non-numerical traits, the list of categories

```
# Get a data.frame back
fb_table_trait_summary(species_traits)
```

```
# Get a kable (to use in Rmd documents)
fb_table_trait_summary(species_traits, TRUE)
```

site_locations

Description

This dataset contains spatial coordinates (grid cells of $0.5^{\circ} \times 0.5^{\circ}$ horizontal resolution) of sites of Western Europe where mammals species presence/absence (see site_species dataset) has been sampled.

Usage

site_locations

Format

An sf object with 1,505 rows (grid cells) with a WGS84 (EPSG:4236) Coordinate Reference System.

Details

See https://github.com/frbcesab/eumammals for a complete description of the workflow used to create this dataset.

This dataset shows the format of the argument site_locations used in several functions of funbiogeo.

The variable site corresponds to the sites labels.

Examples

```
data("site_locations")
class(site_locations)
dim(site_locations)
head(site_locations)
```

site_species

Sites x Species of Western Europe Mammals

Description

This dataset contains the presence/absence of mammals species in Western Europe. This dataset is derived from IUCN range maps downloaded at: https://www.iucnredlist.org/resources/spatial-data-download. In respect to the license, species names have been anonymized.

Usage

site_species

Format

A data.frame with 1,505 rows (sites) and 150 columns (1 column for site label and 149 for species occurrence).

Details

See https://github.com/frbcesab/eumammals for a complete description of the workflow used to create this dataset.

This dataset shows the format of the argument site_species used in several functions of funbiogeo. Note that sites labels (site) is the first column of this dataset.

References

IUCN (2021) The IUCN Red List of Threatened Species. Version 2021-2. Downloaded on 2021/10/11.

Examples

```
data("site_species")
class(site_species)
dim(site_species)
site_species[1:6, 1:6]
```

species_categories Species x Categories of Western Europe Mammals

Description

This dataset contains the order of mammals species in Western Europe. This dataset is derived from IUCN range maps downloaded at: https://www.iucnredlist.org/resources/spatial-data-download. In respect to the license, species names have been anonymized.

Usage

species_categories

Format

A data.frame with 149 rows (species) and 2 columns (1 column for species name and 1 for the order).

Details

See https://github.com/frbcesab/eumammals for a complete description of the workflow used to create this dataset.

This dataset shows the format of the argument species_categories used in several functions of funbiogeo.

Note that species names (species) is the first column of this dataset.

References

IUCN (2021) The IUCN Red List of Threatened Species. Version 2021-2. Downloaded on 2021/10/11.

Examples

```
data("species_categories")
class(species_categories)
dim(species_categories)
species_categories[1:6, ]
```

species_traits

Species x Traits of Western Europe Mammals

Description

This dataset contains values for six functional traits of Western Europe mammals species. These values were extracted from the **PanTHERIA** database (Jones 2009), a database on World mammals traits available at: https://esapubs.org/archive/ecol/E090/184/metadata.htm. In respect to the IUCN license, species names have been anonymized.

Usage

species_traits

Format

A matrix with 149 rows (species) and the following seven columns:

species species name corresponding to the columns of site_species

adult_body_mass adult body mass (in grams)

gestation_length length of time of non-inactive fetal growth (in days)

litter_size number of offspring born per litter per female

max_longevity maximum adult age (in months)

sexual_maturity_age age when individuals are first physically capable of reproducing (in days)

diet_breadth number of dietary categories eaten

Details

See https://github.com/frbcesab/eumammals for a complete description of the workflow used to create this dataset.

This dataset shows the format of the argument species_traits used in several functions of funbiogeo. Note that species names (species) is the first column of this dataset.

38

species_traits

References

Jones KE *et al.* (2009) PanTHERIA: A species-level database of life history, ecology, and geography of extant and recently extinct mammals. *Ecology*, **90**, 2648. DOI: 10.1890/08-1494.1.

```
data("species_traits")
class(species_traits)
dim(species_traits)
head(species_traits)
```

Index

* datasets site_locations, 36 site_species, 36 species_categories, 37 species_traits, 38 fb_aggregate_site_data, 3 fb_count_sites_by_species, 4 fb_count_species_by_site, 4 fb_count_species_by_trait, 5 fb_count_traits_by_species, 6 fb_cwm, 7 fb_filter_sites_by_species_coverage, 8 fb_filter_sites_by_trait_coverage, 9 fb_filter_species_by_site_coverage, 10 fb_filter_species_by_trait_coverage, 11 fb_filter_traits_by_species_coverage, 12 fb_format_site_locations, 13 fb_format_site_species, 14 fb_format_species_categories, 15 fb_format_species_traits, 16 fb_get_all_trait_coverages_by_site, 17 fb_get_environment, 18 fb_get_trait_combination_coverage, 18 fb_get_trait_coverage_by_site, 20 fb_make_report, 21 fb_map_raster, 22 fb_map_site_data, 23 fb_map_site_traits_completeness, 24 fb_plot_distribution_site_trait_coverage, 25 fb_plot_number_sites_by_species, 26 fb_plot_number_species_by_trait, 27 fb_plot_number_traits_by_species, 28 fb_plot_site_environment, 29 fb_plot_site_traits_completeness, 30 fb_plot_species_traits_completeness, 31

knitr::kable(), 35

rmarkdown::render(), 21

site_locations, 36
site_species, 36
species_categories, 37
species_traits, 38
stats::cor(), 34

theme(), 23