Package 'florabr'

September 9, 2024

Title Explore Flora e Funga do Brasil Database

Version 1.3.0

Description A collection of functions designed to retrieve, filter and spatialize data from the Flora e Funga do Brasil dataset. For more information about the dataset, please visit https://floradobrasil.jbrj.gov.br/consulta/>.

Imports XML (>= 3.99.0.14), data.table (>= 1.14.8), httr (>= 1.4.6), terra (>= 1.7.39), stats (>= 4.2.3), utils(>= 4.2.3), grDevices (>= 4.2.3), doSNOW (>= 1.0.20), parallel (>= 4.3.1), foreach (>= 1.5.2)

License GPL (\geq = 3)

Encoding UTF-8

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LazyData true

Suggests knitr, rmarkdown, testthat (>= 3.0.0)

VignetteBuilder knitr

URL https://wevertonbio.github.io/florabr/

BugReports https://github.com/wevertonbio/florabr/issues

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bf_data

Flora e Funga do Brasil database - Version 393.401

Description

A dataset containing a subset of the Flora e Funga do Brasil database (version 393.401)

Usage

data(bf_data)

Format

A data.frame with 50010 rows and 23 variables:

species Species names

scientificName Complete scientific name of the species

- **acceptedName** Accepted name of the species (NA when the name in species is already an accepted name)
- kingdom Kingdom to which species belongs (Plantae or Fungi)
- **group** Major group to which species belongs (Angiosperms, Gymnosperms, Ferns and Lycophytes, Bryophytes, and Algae)
- **subgroup** Subgroup to which species belongs. Only available for Bryophytes (Mosses, Hornworts, and Liverworts)
- phylum Phylum to which species belongs

class Class to which species belongs

biomes

order Order to which species belongs family Family to which species belongs genus Genus to which species belongs **lifeForm** Life form of the species (e.g: Tree, Herb, Shrub, etc.) habitat Habitat type of the species (e.g., Terrestrial, Rupicolous, Epiphytic, etc.) biome Biomes with confirmed occurrences of the species states Federal states with confirmed occurrences of the species vegetation Vegetation types with confirmed occurrences of the species origin Indicates whether the species is Native, Naturalized, or Cultivated in Brazil endemism Indicates whether the species is Endemic or Non-endemic to Brazil taxonomicStatus Indicates the level of recognition and acceptance of the species (Accepted or Synonym) nomenclaturalStatus Indicates the legitimacy and validity of the species name (Correct, Illegitimate, Uncertain_Application, etc.) vernacularName Locally or culturally used name for the species taxonRank Taxonomic rank (Species, Genus, Family, Order, etc). This data contains only Species id Unique id for species

References

Flora e Funga do Brasil. Jardim Botânico do Rio de Janeiro. Available at: http://floradobrasil.jbrj.gov.br/

biomes

SpatVector of the biomes of Brazil

Description

A simplified and packed SpatVector of the polygons of the biomes present in Brazilian territory. The spatial data was originally obtained from geobr::read_biomes. Borders have been simplified by removing vertices of borders using terra::simplifyGeom. It's necessary unpack the Spatvectos using terra::unwrap

@usage data(biomes) biomes <- terra::unwrap(biomes)</pre>

Usage

biomes

Format

A SpatVector with 6 geometries and 1 attribute:

name_biome The name of the biome (Amazon, Caatinga, Cerrado, Atlantic_Forest, Pampa, and Pantanal) brazil

Description

A simplified and packed SpatVector of the Brazil's national borders. The spatial data was originally obtained from geobr::read_country. Borders have been simplified by removing vertices of borders using terra::simplifyGeom. It's necessary unpack the Spatvectos using terra::unwrap

@usage data(brazil) brazil <- terra::unwrap(brazil)

Usage

brazil

Format

A SpatVector with 1 geometry and 0 attribute

check_names

Check species names

Description

check_names checks if the species names are correct and searches for suggestions if the name is misspelled or not found in the Flora e Funga do Brasil database

match_names finds approximate matches to the specified pattern (species) within each element of the string x (species_to_match). It is used internally by check_names.

Usage

check_names

Arguments

data	(data.frame) the data.frame imported with the load_florabr function.						
species	(character) names of the species to be checked.						
max_distance	(numeric) Maximum distance (as a fraction) allowed for searching suggestions when the name is misspelled. It can be any value between 0 and 1. The higher the value, the more suggestions are returned. For more details, see agrep. Default = 0.1 .						
include_subspec	cies						
	(logical) whether to include subspecies. Default = FALSE						
include_variety							
	(logical) whether to include varieties. Default = FALSE						
kingdom	(character) the kingdom to which the species belong. It can be "Plantae" or "Fungi". Default = "Plantae".						
parallel	(logical) whether to run in parallel. Setting this to TRUE is recommended for improved performance when working with 100 or more species.						
ncores	(numeric) number of cores to use for parallel processing. Default is 1. This is only applicable if parallel = TRUE.						
progress_bar	(logical) whether to display a progress bar during processing. Default is FALSE						
<pre>species_to_mate</pre>	species_to_match						

(character) a vector of species names to match against the species parameter.

Value

a data.frame with the following columns:

- input_name: the species names informed in species argument
- Spelling: indicates if the species name is Correct (a perfect match with a species name in the Flora e Funga do Brasil), Probably_incorrect (partial match), or Not_found (no match with any species).
- Suggested name: If Spelling is Correct, it is the same as the input_name. If Spelling is Probably_correct, one or more suggested names are listed, found according to the maximum distance. If Spelling is "Not_found", the value is NA.
- Distance: The integer Levenshtein edit distance. It represents the number of single-character edits (insertions, deletions, or substitutions) required to transform the input_name into the Suggested_name.
- taxonomicStatus: the taxonomic status of the species name ("Accepted" or "Synonym").
- nomenclaturalStatus: the nomenclatural status of the species name. This information is not available for all species.
- acceptedName: If the species name is not accepted or incorrect, the accepted name of the specie. If the species name is accepted and correct, the same as input_name and Suggested_name.
- family: the family of the specie.

References

Flora e Funga do Brasil. Jardim Botânico do Rio de Janeiro. Available at: http://floradobrasil.jbrj.gov.br/

Examples

```
data("bf_data", package = "florabr")
spp <- c("Butia cattarinensis", "Araucaria angustifolia")
check_names(data = bf_data, species = spp)</pre>
```

check_version

Check if you have the latest version of Flora e Funga do Brasil data available

Description

This function checks if you have the latest version of the Flora e Funga do Brasil data available in a specified directory.

Usage

```
check_version(data_dir)
```

Arguments

data_dir the directory where the data should be located.

Value

A message informing whether you have the latest version of Flora e Funga do Brasil available in the data_dir

Examples

```
#Check if there is a version of Flora e Funga do Brasil data available in the
#current directory
check_version(data_dir = getwd())
```

filter_florabr	Identify records outside natural ranges according to Flora e Funga do
	Brasil

Description

This function removes or flags records outside of the species' natural ranges according to information provided by the Flora e Funga do Brasil database.

filter_florabr

Usage

data	(data.frame) the data.frame imported with the load_florabr function.
осс	(data.frame) a data.frame with the records of the species.
species	(character) column name in occ with species names. Default = "species"
long	(character) column name in occ with longitude data. Default = "x"
lat	(character) column name in occ with latitude data. Default = "y"
by_state	(logical) filter records by state? Default = TRUE
buffer_state	(numeric) buffer (in km) around the polygons of the states of occurrence of the specie. Default = 20 .
by_biome	(logical) filter records by biome? Default = TRUE
buffer_biome	(numeric) buffer (in km) around the polygons of the biomes of occurrence of the specie. Default = 20.
by_endemism	(logical) filter records by endemism? Default = TRUE
buffer_brazil	(numeric) buffer (in km) around the polygons of the brazil. Default = 20.
state_vect	(SpatVector) a SpatVector of the Brazilian states. By default, it uses the SpatVector provided by geobr::read_state(). It can be another Spatvector, but the structure must be identical to geobr::read_state().
state_column	(character) name of the column in state_vect containing state abbreviations. Only use if biome_vect is not null.
biome_vect	(SpatVector) a SpatVector of the Brazilian biomes. By default, it uses the SpatVector provided by geobr::read_biomes(). It can be another SpatVector, but the structure must be identical to geobr::read_biomes() with biome names in English.
biome_column	(character) name of the column in biome_vect containing names of brazilian biomes (in English: "Amazon", "Atlantic_Forest", "Caatinga", "Cerrado", "Pampa" and "Pantanal". Only use if biome_vect is not null.
br_vect	(SpatVector) a SpatVector of brazil. By default, it uses the SpatVector provided by geobr::read_state() after being aggregated/dissolved,
value	(character) Defines output values. See Value section. Default = "flag&clean".
keep_columns	(logical) if TRUE, keep all the original columns of the input occ. If False, keep only the columns species, long and lat. Default = TRUE
verbose	(logical) Whether to display species being filtered during function execution. Set to TRUE to enable display, or FALSE to run silently. Default = TRUE.

Details

If by_state = TRUE and/or by_biome = TRUE, the function takes polygons representing the states and/or biomes with confirmed occurrences of the specie, draws a buffer around the polygons, and tests if the records of the species fall inside it. If by_endemism = TRUE, the function checks if the species is endemic to brazil. If it is endemic, the function tests if the records of the specie fall inside a polygon representing the boundaries of brazil (with a buffer).

Value

Depending on the 'value' argument. If value = "flag", it returns the same data.frame provided in data with additional columns indicating if the record falls inside the natural range of the specie (TRUE) or outside (FALSE). If value = "clean", it returns a data.frame with only the records that passes all the tests (TRUE for all the filters). If value = "flag&clean" (Default), it returns a list with two data.frames: one with the flagged records and one with the cleaned records.

References

Flora e Funga do Brasil. Jardim Botânico do Rio de Janeiro. Available at: http://floradobrasil.jbrj.gov.br/

Examples

```
get_attributes
```

Get available attributes to filter species

Description

This function displays all the options available to filter species by its characteristics

Usage

```
get_attributes(data, attribute)
```

data	(data.frame) a data.frame imported with the load_florabr function or a data.frame
	generated with the select_species function.
attribute	(character) the type of characteristic. Accept more than one option. See detail to see the options.

get_binomial

Details

The attribute argument accepts the following options: kingdom, group, subgroup, phylum, class, order, family, lifeform, habitat, vegetation, origin, endemism, biome, states, taxonomicstatus or nomenclatural status. These options represent different characteristics of species that can be used for filtering.

Value

a list of data.frames with the available options to use in the select_species function.

References

Flora e Funga do Brasil. Jardim Botânico do Rio de Janeiro. Available at: http://floradobrasil.jbrj.gov.br/

Examples

get_binomial	Extract the binomial name (Genus + specific epithet + infraspecific
	epithet (optional)) from a full Scientific Name

Description

Extract the binomial name (Genus + specific epithet + infraspecific epithet (optional)) from a full Scientific Name

Usage

Arguments

species_names (character) Scientific names to be converted to binomial names

include_subspecies

(logical) include subspecies? If TRUE (default), the function extracts any infraspecific epithet after the pattern "subsp."

include_variety

(logical) include subspecies? If TRUE (default), the function extracts any infraspecific epithet after the pattern "var."

Value

A vector with the binomial names (Genus + specific epithet).

Examples

get_florabr

Download the latest version of Flora e Funga do Brasil database

Description

This function downloads the latest or an older version of Flora e Funga do Brasil database, merges the information into a single data.frame, and saves this data.frame in the specified directory.

Usage

output_dir	(character) a directory to save the data downloaded from Flora e Funga do Brasil.							
data_version	(character) Version of the Flora e Funga do Brasil database to download. Use "latest" to get the most recent version, updated weekly. Alternatively, specify an older version (e.g., data_version = "393.319"). Default value is "latest".							
solve_discrepancy								
	Resolve discrepancies between species and subspecies/varieties information. When set to TRUE, species information is updated based on unique data from varieties and subspecies. For example, if a subspecies occurs in a certain biome, it implies that the species also occurs in that biome. Default = FALSE.							
overwrite	(logical) If TRUE, data is overwritten. Default = TRUE.							
verbose	(logical) Whether to display messages during function execution. Set to TRUE to enable display, or FALSE to run silently. Default = TRUE.							
remove_files	(logical) Whether to remove the downloaded files used in building the final dataset. Default is TRUE.							

get_pam

Value

The function downloads the latest version of the Flora e Funga do Brasil database from the official source. It then merges the information into a single data.frame, containing details on species, taxonomy, occurrence, and other relevant data. The merged data.frame is then saved as a file in the specified output directory. The data is saved in a format that allows easy loading using the load_florabr function for further analysis in R.

References

Flora e Funga do Brasil. Jardim Botânico do Rio de Janeiro. Available at: http://floradobrasil.jbrj.gov.br/

Examples

End(Not run)

get_pam

Get a presence-absence matrix

Description

Get a presence-absence matrix of species based on its distribution (states, biomes and vegetation types) according to Flora e Funga do Brasil.

Usage

data	(data.frame) a data.frame imported with the load_florabr function or gener-
	ated by either select_species or subset_species functions
by_biome	(logical) get occurrences by biome. Default = TRUE
by_state	(logical) get occurrences by State. Default = TRUE

by_vegetation	(logical) get occurrences by vegetation type. Default = FALSE
<pre>remove_empty_si</pre>	tes
	(logical) remove empty sites (sites without any species) from final presence-absence matrix. Default = TRUE
return_richness	_summary
	(logical) return a data.frame with the number of species in each site. Default = TRUE
<pre>return_spatial_</pre>	richness
	(logical) return a SpatVector with the number of species in each site. Default = TRUE
return_plot	(logical) plot map with the number of species in each site. Only works if re- turn_spatial_richness = TRUE. Default = TRUE

Value

If return_richness_summary and/or return_spatial_richness is set to TRUE, return a list with:

- PAM: the presence-absence matrix (PAM)
- Richness_summary: a data.frame with the number of species in each site
- Spatial_richness: a SpatVector with the number of species in each site (only by State and biome)

If return_richness_summary and return_spatial_richness is set to FALSE, return a presence-absence matrix

References

Flora e Funga do Brasil. Jardim Botânico do Rio de Janeiro. Available at: http://floradobrasil.jbrj.gov.br/

Examples

```
data("bf_data") #Load Flora e Funga do Brasil data
#Select endemic and native species of trees with occurrence only in Amazon
am_trees <- select_species(data = bf_data,</pre>
                          include_subspecies = FALSE,
                          include_variety = FALSE,
                          kingdom = "Plantae",
                          group = "All", subgroup = "All",
                          family = "All", genus = "All",
                          lifeForm = "Tree", filter_lifeForm = "only",
                          habitat = "All", filter_habitat = "in",
                          biome = "Amazon",
                          filter_biome = "only",
                          state = "All", filter_state = "and",
                          vegetation = "All",
                          filter_vegetation = "in",
                          endemism = "Endemic", origin = "Native",
                          taxonomicStatus = "Accepted",
                          nomenclaturalStatus = "All")
```

#Get presence-absence matrix

get_spat_occ

Get Spatial polygons (SpatVectors) of species based on its distribution (states and biomes) according to Flora e Funga do Brasil

Description

Get Spatial polygons (SpatVectors) of species based on its distribution (states and biomes) according to Flora e Funga do Brasil

Usage

```
get_spat_occ(
   data,
   species,
   state = TRUE,
   biome = TRUE,
   intersection = TRUE,
   state_vect = NULL,
   state_column = NULL,
   biome_vect = NULL,
   biome_column = NULL,
   verbose = TRUE
)
```

data	(data.frame) the data.frame imported with the load_florabr function.
species	(character) one or more species names (only genus and specific epithet, eg. "Araucaria angustifolia")
state	(logical) get SpatVector of states with occurrence of the species? Default = TRUE
biome	(logical) get SpatVector of biomes with occurrence of the species? Default = TRUE
intersection	(character) get a Spatvector representing the intersection between states and biomes with occurrence of the specie? To use intersection = TRUE, you must define state = TRUE and biome = TRUE". Default = TRUE
state_vect	(SpatVector) a SpatVector of the Brazilian states. By default, it uses the SpatVector provided by geobr::read_state(). It can be another Spatvector, but the structure must be identical to geobr::read_state().

state_column	(character) name of the column in state_vect containing state abbreviations. Only use if biome_vect is not null.
biome_vect	(SpatVector) a SpatVector of the Brazilian biomes. By default, it uses the SpatVector provided by geobr::read_biomes(). It can be another SpatVector, but the structure must be identical to geobr::read_biomes().
biome_column	(character) name of the column in biome_vect containing names of brazilian biomes (in English: "Amazon", "Atlantic_Forest", "Caatinga", "Cerrado", "Pampa" and "Pantanal". Only use if biome_vect is not null.
verbose	(logical) Whether to display species being filtered during function execution. Set to TRUE to enable display, or FALSE to run silently. Default = TRUE.

Value

A list with SpatVectors of states and/or biomes and/or Intersections for each specie.

References

Flora e Funga do Brasil. Jardim Botânico do Rio de Janeiro. Available at: http://floradobrasil.jbrj.gov.br/

Examples

```
#Plot states of occurrence of Araucaria angustifolia
plot(spp_spt[[1]]$states, main = names(spp_spt)[[1]])
#Plot biomes of occurrence of Araucaria angustifolia
plot(spp_spt[[2]]$biomes, main = names(spp_spt)[[2]])
#Plot intersection between states and biomes of occurrence of
#Araucaria angustifolia
plot(spp_spt[[1]]$states_biomes)
```

get_synonym

Retrieve synonyms for species

Description

Retrieve synonyms for species

load_florabr

Usage

Arguments

data	(data.frame) the data.frame imported with the load_florabr function	
species	(character) names of the species	
include_subspecies		
	(logical) include subspecies that are synonyms of the species? Default = TRUE	
include_variety		
	(logical) include varieties that are synonyms of the species? Default = TRUE	

Value

A data.frame containing unique synonyms of the specified species along with relevant information on taxonomic and nomenclatural statuses.

References

Flora e Funga do Brasil. Jardim Botânico do Rio de Janeiro. Available at: http://floradobrasil.jbrj.gov.br/

Examples

spp_synonyms

load_florabr

Load Flora e Funga do Brasil database

Description

Load Flora e Funga do Brasil database

Usage

Arguments

data_dir	(character) the same directory used to save the data downloaded from Flora e Funga do Brasil using the get_florabr function.
data_version	(character) the version of Flora e Funga do Brasil database to be loaded. It can be "Latest_available", which will load the latest version available; or another specified version, for example "393.364". Default = "Latest_available".
type	(character) it determines the number of columns that will be loaded. It can be "short" or "complete". Default = "short". See details.
verbose	(logical) Whether to display messages during function execution. Set to TRUE to enable display, or FALSE to run silently. Default = TRUE.

Details

The parameter type accepts two arguments. If type = short, it will load a data.frame with the 20 columns needed to run the other functions of the package: species, scientificName, acceptedName, kingdom, Group, Subgroup, family, genus, lifeForm, habitat, Biome, States, vegetationType, Origin, Endemism, taxonomicStatus, nomenclaturalStatus, vernacularName, taxonRank, and id If type = complete, it will load a data.frame with all 39 variables available in Flora e Funga do Brasil database.

Value

A data.frame with the specified version (Default is the latest available) of the Flora e Funga do Brasil database. This data.frame is necessary to run most of the functions of the package.

References

Flora e Funga do Brasil. Jardim Botânico do Rio de Janeiro. Available at: http://floradobrasil.jbrj.gov.br/

Examples

occurrences

Description

A dataset containing records of 7 plant species downloaded from GBIF. The records were obtained with plantR::rgbif2

Usage

data(occurrences)

Format

A data.frame with 1521 rows and 3 variables:

- **species** Species names (Araucaria angustifolia, Abatia americana, Passiflora edmundoi, Myrcia hatschbachii, Serjania pernambucensis, Inga virescens, and Solanum restingae)
- x Longitude

y Latitude

References

GBIF, 2024. florabr R package: Records of plant species. https://doi.org/10.15468/DD.QPGEB7

select_by_vernacular Search for taxa using vernacular names

Description

Search for taxa using vernacular names

Usage

```
select_by_vernacular(data, names, exact = FALSE)
```

data	(data.frame) the data.frame imported with the load_florabr function or gener- ated with the function select_species.
names	(character) vernacular name ("Nome comum") of the species to be searched
exact	(logic) if TRUE, the function will search only for exact matches. For example, if names = "pinheiro" and exact = TRUE, the function will return only the species popularly known as "pinheiro". On the other hand, if names = "pinheiro" and exact = FALSE, the function will return other results as "pinheiro-do-parana". Default = FALSE

Value

a data.frame with the species with vernacular names that match the input names

References

Flora e Funga do Brasil. Jardim Botânico do Rio de Janeiro. Available at: http://floradobrasil.jbrj.gov.br/ Flora e Funga do Brasil. Jardim Botânico do Rio de Janeiro. Available at: http://floradobrasil.jbrj.gov.br/

Examples

<pre>select_species</pre>	Selection of species based on its characteristics and distribution
---------------------------	--

Description

select_species allows filter species based on its characteristics and distribution available in Flora e Funga do Brasil

Usage

select_species(data,

```
include_subspecies = FALSE, include_variety = FALSE,
kingdom = "Plantae", group = "All", subgroup = "All",
phylum = "All", class ="All", order = "All",
family = "All", genus = "All",
lifeForm = "All", filter_lifeForm = "in",
habitat = "All", filter_habitat = "in",
biome = "All", filter_biome = "in",
state = "All", filter_state = "in",
vegetation = "All", filter_vegetation = "in",
endemism = "All", origin = "All",
taxonomicStatus = "Accepted",
nomenclaturalStatus = "All")
```

select_species

data	(data.frame) the data.frame imported with the load_florabr function.
include_subspec	ies
	(logical) include subspecies? Default = FALSE
include_variety	(logical) include varieties of the species? Default = FALSE
kingdom	(character) The kingdom for filtering the dataset. It can be "Plantae" or "Fungi". Default = "Plantae". To include both, use c("Plantae", "Fungi")
group	(character) The groups for filtering the datasets. It can be "Fungi", "Angiosperms", "Gymnosperms", "Ferns and Lycophytes", "Bryophytes" and "Algae". To use more than one group, put the available items in a vector, for example: group = c(Angiosperms", "Gymnosperms"). Default = "All".
subgroup	(character) The subgroups for filtering the dataset. Only available if the group is "Fungi" or "Bryophytes". For Fungi, it can be "stricto sensu" or "lato sensu". For Bryophytes, it can be "Mosses", "Hornworts" and "Liverworts" . To use more than one group, put the available items in a vector, for example: subgroup = c("Mosses", "Hornworts"). Default = "All".
phylum	(character) The phyla for filtering the dataset. It can be included more than one phylum. Default = "All".
class	(character) The classes for filtering the dataset. It can be included more than one class. Default = "All".
order	(character) The orders for filtering the dataset. It can be included more than one order. Default = "All".
family	(character) The families for filtering the dataset. It can be included more than one family. Default = "All".
genus	(character) The genus for filtering the dataset. It can be included more than one genus. Default = "All".
lifeForm	(character) The life forms for filtering the dataset. It can be included more than one lifeForm. Default = "All"
filter_lifeForm	
	(character) The type of filtering for life forms. It can be "in", "only", "not_in" and "and". See details for more about this argument.
habitat	(character) The life habitat for filtering the dataset. It can be included more than one habitat. Default = "All"
filter_habitat	(character) The type of filtering for habitat. It can be "in", "only", "not_in" and "and". See details for more about this argument.
biome	(character) The biomes for filtering the dataset. It can be included more than one biome. Default = "All"
filter_biome	(character) The type of filtering for biome. It can be "in", "only", "not_in" and "and". See details for more about this argument.
state	(character) The states for filtering the dataset. It can be included more than one state. Default = "All".

filter_state	(character) The type of filtering for state. It can be "in", "only", "not_in" and "and". See Details for more about this argument.	
vegetation	(character) The vegetation types for filtering the dataset. It can be included more than one vegetation type. Default = "All".	
filter_vegetati	on	
	(character) The type of filtering for vegetation type. It can be "in", "only", "not_in" and "and". See details for more about this argument.	
endemism	(character) The endemism (endemic or non-endemic to Brazil) for filtering the dataset. It can be "All", "Endemic" or "Non-endemic". Default = "All".	
origin	(character) The origin for filtering the dataset. It can be "All", "Native", "Cultivated" and "Naturalized". Default = "All".	
taxonomicStatus		
	(character) The taxonomic status for filtering the dataset. It can be "All", "Accepted" or "Synonym". Default = "Accepted".	
nomenclaturalStatus		
	(character) The nomenclatural status for filtering the dataset. Default = "Accepted"	

Details

It's possible to choose 4 ways to filter by lifeForm, by habitat, by biome, by state and by vegetation type: "in": selects species that have any occurrence of the determined values. It allows multiple matches. For example, if biome = $c("Amazon", Cerrado" and filter_biome = "in", it will select all species that occur in the Amazon and Cerrado, some of which may also occur in other biomes.$

"only": selects species that have only occurrence of the determined values. It allows only single matches. For example, if biome = c("Amazon", "Cerrado") and filter_biome = "only", it will select all species that occur exclusively in both the Amazon and Cerrado biomes, without any occurrences in other biomes.

"not_in": selects species that don't have occurrence of the determined values. It allows single and multiple matches. For example, if biome = c("Amazon", "Cerrado") and filter_biome = "not_in", it will select all species without occurrences in the Amazon and Cerrado biomes.

"and": selects species that have occurrence in all determined values. It allows single and multiple matches. For example, if biome = c("Amazon", "Cerrado") and filter_biome = "and", it will select all species that occurs only in both the Amazon and Cerrado biomes, including species that occurs in other biomes too.

To get the complete list of arguments available for family, genus, lifeForm, habitat, biome, state, and nomenclaturalStatus, use the function get_attributes

Value

A new dataframe with the filtered species.

References

Flora e Funga do Brasil. Jardim Botânico do Rio de Janeiro. Available at: http://floradobrasil.jbrj.gov.br/

solve_discrepancies

Examples

```
data("bf_data") #Load Flora e Funga do Brasil data
#'Select endemic and native species of trees with disjunct occurrence in
# Atlantic Forest and Amazon
am_af_only <- select_species(data = bf_data,</pre>
                             include_subspecies = FALSE,
                             include_variety = FALSE,
                             kingdom = "Plantae",
                             group = "All", subgroup = "All",
                             phylum = "All", class ="All", order = "All",
                             family = "All", genus = "All",
                             lifeForm = "Tree", filter_lifeForm = "only",
                             habitat = "All", filter_habitat = "in",
                             biome = c("Atlantic_Forest", "Amazon"),
                             filter_biome = "only",
                             state = "All", filter_state = "and",
                             vegetation = "All",
                             filter_vegetation = "in",
                             endemism = "Endemic", origin = "Native",
                             taxonomicStatus = "All",
                             nomenclaturalStatus = "All")
```

solve_discrepancies	Resolve discrepancies between species and subspecies/varieties infor-
	mation

Description

Resolve discrepancies between species and subspecies/varieties information

Usage

```
solve_discrepancies(data)
```

Arguments

data

(data.frame) the data.frame imported with the load_florabr function.

Details

In the original dataset, discrepancies may exist between species and subspecies/varieties information. An example of a discrepancy is when a species occurs only in one biome (e.g., Amazon), but a subspecies or variety of the same species occurs in another biome (e.g., Cerrado). This function rectifies such discrepancies by considering distribution (states, biomes, and vegetation), life form, and habitat. For instance, if a subspecies is recorded in a specific biome, it implies that the species also occurs in that biome.

Value

a data.frame with the discrepancies solved

Examples

```
data("bf_data") #Load Flora e Funga do Brasil data
#Check if discrepancies were solved in the dataset
attr(bf_data, "solve_discrepancies")
#Solve discrepancies
bf_solved <- solve_discrepancies(bf_data)
#Check if discrepancies were solved in the dataset
attr(bf_solved, "solve_discrepancies")
```

states

SpatVector of the federal states of Brazil

Description

A simplified and packed SpatVector of the polygons of the federal states of Brazil. The spatial data was originally obtained from geobr::read_state. Borders have been simplified by removing vertices of borders using terra::simplifyGeom. It's necessary unpack the Spatvectos using terra::unwrap

@usage data(states) states <- terra::unwrap(states)</pre>

Usage

states

Format

A SpatVector with 27 geometries and 3 attributes:

abbrev_state State acronym

name_state State's full name

name_region The region to which the state belongs

subset_species Extract a subset of species from Flora e Funga do Brasil database

Description

Returns a data.frame with a subset of species from Flora e Funga do Brasil database

Usage

22

subset_species

Arguments

data	(data.frame) the data.frame imported with the load_florabr function.	
species	(character) names of the species to be extracted from Flora e Funga do Brasil database.	
include_subspecies		
	(logical) include subspecies? Default = FALSE	
include_variety		
	(logical) include varieties of the species? Default = FALSE	
kingdom	(character) The kingdom for filtering the dataset. It can be "Plantae" or "Fungi". Default = "Plantae". To include both, use c("Plantae", "Fungi")	

Value

A data.frame with the selected species.

References

Flora e Funga do Brasil. Jardim Botânico do Rio de Janeiro. Available at: http://floradobrasil.jbrj.gov.br/

Examples

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