

Package ‘gbif.range’

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Version 1.0

Title A toolbox to generate species range maps based on ecoregions and a user-friendly GBIF wrapper

Depends R (>= 4.0.0)

Imports terra, rgbif, CoordinateCleaner, sf, ClusterR, FNN, geometry, cluster, rnatu-
ralearth, mclust, methods

Description This package provides a workflow to gener-
ate species range maps from scratch based on ecoregions and a user-friendly GBIF wrapper. It in-
cludes functions to create taxa range maps using ecoregions and to interact with the Global Bio-
diversity Information Facility (GBIF) database (retrieving species records, backbone taxon-
omy and IUCN conservation status). The package facilitates the process of access-
ing GBIF data and integrating it with ecoregion information for species distribution analysis.

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BugReports <https://github.com/8Ginette8/gbif.range/issues>

Encoding UTF-8

LazyData true

RoxygenNote 7.3.1

Collate 'make_tiles.R'
'get_gbif.R'
'get_status.R'
'get_doi.R'
'obs_filter.R'
'get_range.R'
'conv_function.R'
'make_ecoregion.R'

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`get_doi`*Get a custom DOI for a GBIF filtered dataset*

Description

A small user friendly wrapper of the `derived_dataset()` function of the `rgbif` R package, compatible with one or several `get_gbif()` outputs.

Usage

```
get_doi(  
  gbifs = NULL,  
  title = NULL,  
  description = NULL,  
  source_url = "https://example.com/",  
  user = "",  
  pwd = "",  
  ...  
)
```

Arguments

<code>gbifs</code>	data.frame or list. One <code>get_gbif()</code> output or a list of several.
<code>title</code>	Title for your derived dataset.
<code>description</code>	Description of the dataset.
<code>source_url</code>	Link to where the dataset is stored.
<code>user</code>	Your GBIF username.
<code>pwd</code>	Your GBIF password.
<code>...</code>	Additonnal parameters for <code>derived_dataset()</code> in <code>rgbif</code> .

Details

see `derived_dataset()` function from the `rgbif` R package.

Value

One citable DOI and its information.

References

Chamberlain, S., Oldoni, D., & Waller, J. (2022). `rgbif`: interface to the global biodiversity information facility API. 10.5281/zenodo.6023735

See Also

The `rgbif` package for additional and more general approaches to get GBIF DOI.

Examples

```
# Download worldwide the observations of Panthera tigris and Ailuropoda melanoleuca
obs.pt <- get_gbif("Panthera tigris")
obs.am <- get_gbif("Ailuropoda melanoleuca")
## Not run:
# Retrieve DOI for only one get_gbif() output
get_doi(obs.pt,title="GBIF_test1",description="A small example 1",
  source_url="https://example.com/",user="",pwd="") # Use your own GBIF credentials here

# Retrieve DOIs for several get_gbif() outputs
get_doi(list(obs.pt,obs.am),title="GBIF_test2",description="A small example 2",
  source_url="https://example.com/",user="",pwd="") # Use your own GBIF credentials here

## End(Not run)
```

get_gbif

Massively download and filter GBIF observations for sound spatial analyses

Description

Implement an user-friendly workflow to download and clean gbif taxa observations. The function uses the rgbif R package but (1) implements the same search result found if www.gbif.org is employed i.e., based on the input taxa name, all species records related to its accepted name and synonyms are extracted. The function also (2) bypasses the rgbif hard limit on the number of records (100'000 max). For this purpose, a dynamic moving window is created and used across the geographic extent defined by the user. This window automatically fragments the specified study area in successive tiles of different sizes, until all tiles include < 100'000 observations. The function also (3) automatically applies a post-filtering of observations based on the chosen resolution of the study/analysis and by partly employing the CoordinateCleaner R package. Filtering options may be chosen and involve several choices: study's extent, removal of duplicates, removal of absences, basis of records selection, removal of invalid/uncertain xy coordinates (WGS84), time period selection and removal of raster centroids. By default, the argument `hasGeospatialIssue` in `occ_data()` (implemented `rgbif` function) is set to `FALSE`.

Usage

```
get_gbif(
  sp_name = NULL,
  search = TRUE,
  rank = NULL,
  phylum = NULL,
  class = NULL,
  order = NULL,
  family = NULL,
  conf_match = 80,
  geo = NULL,
  grain = 1000,
  duplicates = FALSE,
  absences = FALSE,
  no_xy = FALSE,
```

```

basis = c("OBSERVATION", "HUMAN_OBSERVATION", "MACHINE_OBSERVATION", "OCCURRENCE",
          "MATERIAL_CITATION", "MATERIAL_SAMPLE", "LITERATURE"),
add_infos = NULL,
time_period = c(1000, 3000),
identic_xy = FALSE,
wConverted_xy = FALSE,
centroids = FALSE,
ntries = 10,
error.skip = TRUE,
occ_samp = 99000,
...
)

```

Arguments

sp_name	Character. Species name from which the user wants to retrieve all existing GBIF names with associated taxonomy and IUCN status
search	Logical. If TRUE, the function will strictly look for the most relevant result, based on a list of names given by rgbif, and give an error if name matching was impeded by synonym duplicates. If FALSE, the function will simply pick the first most relevant name from the list. Also, unlike search=TRUE, fuzzy search (~approximative name match) is here allowed, and the 'rank', 'phylum', 'class', 'order' and 'family' parameters are optionally used only if no convincing name match is found.
rank	Character. "SPECIES", "SUBSPECIES" or "VARIETY". If NULL (default), the order of priority is (1) species, (2) subspecies and (3) variety unless "subsp." or "var." is found in 'sp_name'.
phylum	Character. Optional. What is the species' Phylum? Adds a criteria to deal with alternative name matches and select the right synonym. Available options are the GBIF Phylums (listed per Kingdom → https://www.gbif.org/species/1). If search = FALSE, used only if no direct match is found.
class	Character. Optional. What is the species' Class? Same as above but at the finer class level. Available options are the GBIF Classes (same url). If search = FALSE, used only if no direct match is found.
order	Character. Optional. What is the species' Order? Same as above but at the finer order level. Available options are the GBIF Orders (same url). If search = FALSE, used only if no direct match is found.
family	Character. Optional. What is the species' Family? Same as above but at the finer family level. Available options are the GBIF Orders (same url). If search = FALSE, used only if no direct match is found.
conf_match	Numeric from 0 to 100. Determine the confidence threshold of match of 'sp_name' with the GBIF backbone taxonomy. Default is 90.
geo	Object of class Extent, SpatExtent, SpatialPolygon, SpatialPolygonDataframe, or SpatVector (WGS84) to define the study's area extent. Default is NULL i.e. the whole globe.
grain	Numeric. Specify in meters the study resolution. Used to filter gbif records according to their (1) spatial uncertainties and (2) number of coordinate decimals. Records with no information on coordinate uncertainties (column 'coordinateUncertaintyInMeters') are kept by default. See details.
duplicates	Logical. Should duplicated records be kept?

absences	Logical. Should absence records be kept?
no_xy	Logical. Default is FALSE i.e., only records with coordinates are downloaded. If TRUE, records with no coordinates are also downloaded.
basis	Character. Which basis of records should be selected? Available (old and new) are 'OBSERVATION', 'HUMAN_OBSERVATION', 'MACHINE_OBSERVATION', 'MATERIAL_CITATION', 'MATERIAL_SAMPLE', 'PRESERVED_SPECIMEN', 'FOSSIL_SPECIMEN', 'LIVING_SPECIMEN', 'LITERATURE', 'UNKNOWN' and 'OCCURRENCE'. Default setting removes specimens and unknown observations. Description may be found here: https://docs.gbif.org/course-data-use/en/basis-of-record.html , https://gbif.github.io/parsers/apidocs/org/gbif/api/vocabulary/BasisOfRecord.html
add_infos	Character. Infos that may be added to the default output information. List of IDs may be found at: https://www.gbif.org/developer/occurrence . Default IDs contain 'taxonKey', 'scientificName', 'acceptedTaxonKey', 'acceptedScientificName', 'individualCount', 'decimalLatitude', 'decimalLongitude', 'basisOfRecord', 'coordinateUncertaintyInMeters', 'countryCode', 'country', 'year', 'datasetKey', 'institutionCode', 'publishingOrgKey', 'taxonomicStatus' and 'taxonRank'.
time_period	Numerical vector. Observations will be downloaded according to the chosen year range. Default is c(1000,3000). Observations with year = NA are kept by default.
identic_xy	Logical. Should records with identical xy be kept?
wConverted_xy	Logical. Should incorrectly converted lon/lat be kept? Uses cd_ddmm() from 'CoordinateCleaner' R package.
centroids	Logical. Should species records from raster centroids be kept? Uses cd_round() from 'CoordinateCleaner' R package.
ntries	Numeric. In case of failure from GBIF server or within the rgbif package, how many download attempts should the function request? Default is '10' with a 2 seconds interval between tries. If the attempts failed, an empty data.frame is return by default.
error.skip	Logical. Should the search process continues if ntries failed ?
occ_samp	Numeric. Determine how many GBIF occurrences will be sampled per geographic tiles of the fragmented study area. Default is the maximum number of GBIF observations found in a tile (i.e. ~100'000 records). A lower number may be set (<99'000) if the user only wants a sample of the species GBIF observations, hence increasing the download process and the generation of its range map if get_range() is employed afterwards.
...	Additonnal parameters for the function cd_round() of CoordinateCleaner.

Details

Argument 'grain' used for two distinct gbif records filtering. (1) Records filtering according to gbif 'coordinateUncertaintyInMeters'; every records uncertainty > grain/2 are removed. Note: Records with no information on coordinate uncertainties are kept by default. (2) Records filtering according to the number of longitude/latitude decimals; if 110km < grain <= 11km, lon/lat with >= 1 decimal are kept; if 11km < grain <= 1100m, lon/lat with >= 2 decimals kept; if 1100m < grain <= 110m, lon/lat with >= 3 decimals are kept; if 110m < grain <= 11m, lon/lat with >= 4 decimals are kept; if 11m < grain <= 1.1m, lon/lat with >= 5 decimals are kept etc...

Value

Object of class `data.frame` with requested GBIF information. Although the function works accurately, error outputs might still occur depending on the `'sp_name'` used. Therefore, default information detailed in `'add_infos'` is stored so that sanity checks may still be applied afterwards. Although crucial preliminary checks of species records are done by the function, additional post exploration with the `CoordinateCleaner` R package is still highly recommended.

References

- Chauvier, Y., Thuiller, W., Brun, P., Lavergne, S., Descombes, P., Karger, D. N., ... & Zimmermann, N. E. (2021). Influence of climate, soil, and land cover on plant species distribution in the European Alps. *Ecological monographs*, 91(2), e01433. 10.1002/ecm.1433
- Chamberlain, S., Oldoni, D., & Waller, J. (2022). `rgbif`: interface to the global biodiversity information facility API. 10.5281/zenodo.6023735
- Zizka, A., Silvestro, D., Andermann, T., Azevedo, J., Duarte Ritter, C., Edler, D., ... & Antonelli, A. (2019). `CoordinateCleaner`: Standardized cleaning of occurrence records from biological collection databases. *Methods in Ecology and Evolution*, 10(5), 744-751. 10.1111/2041-210X.13152
- Hijmans, Robert J. "terra: Spatial Data Analysis. R Package Version 1.6-7." (2022). Terra - CRAN

See Also

The (1) `rgbif` and (2) `CoordinateCleaner` packages for additional and more general approaches on (1) downloading GBIF observations and (2) post-filtering those.

Examples

```
# Downloading worldwide the observations of Panthera tigris
obs.pt <- get_gbif("Panthera tigris",
  basis=c("OBSERVATION", "HUMAN_OBSERVATION", "MACHINE_OBSERVATION"))
countries <- terra::vect(rnaturalearth::ne_countries(type = "countries", returnclass = "sf"))
terra::plot(countries, col="#bcbddc")
points(obs.pt[,c("decimalLongitude", "decimalLatitude")], pch=20, col="#238b4550", cex=4)

## Not run:

# Downloading worldwide the observations of Ailuropoda melanoleuca (with a 100km grain, after 1990
# and by keeping duplicates and by adding the name of the person who collected the panda records)
obs.am <- get_gbif("Ailuropoda melanoleuca", grain = 100000 , duplicates = TRUE,
  time_period = c(1990,3000), add_infos = c("recordedBy", "issue"))
terra::plot(countries, col="#bcbddc")
graphics::points(obs.am[,c("decimalLongitude", "decimalLatitude")], pch=20, col="#238b4550", cex=4)

# Downloading worldwide the observations of Phascolarctos cinereus (with a 1km grain, after 1980,
# and keeping raster centroids)
obs.pc <- get_gbif("Phascolarctos cinereus", grain = 1000,
  time_period = c(1990,3000), centroids = TRUE)

## End(Not run)
```

get_range

*Create a species range map based on a get_gbif() output***Description**

Estimates species ranges based on occurrence data (GBIF or not) and ecoregions (may be an external, in-house or make_ecoregion() input). It first deletes outliers from the observation dataset and then creates a polygon (convex hull) with a user specified buffer around all the observations of one ecoregion. If there is only one observation in an ecoregion, a buffer around this point will be created. If all points in an ecoregion are on a line, the function will also create a buffer around these points, however, the buffer size increases with the number of points in the line. Finally, also note that in case of too many records, get_range can be used with a sub-sample of species observations to ensure a faster polygon process and/or to overcome potential RAM crash of the function.

Usage

```
get_range(
  sp_name = NULL,
  occ_coord = NULL,
  Bioreg = eco.earth,
  Bioreg_name = "ECO_NAME",
  degrees_outlier = 3,
  clustered_points_outlier = 2,
  buffer_width_point = 4,
  buffer_increment_point_line = 0.5,
  buffer_width_polygon = 4,
  dir_temp = paste0("temp", sample(1:99999999, 1)),
  raster = TRUE,
  res = 100
)
```

Arguments

sp_name	Character. Species name e.g., 'Anemone nemorosa'.
occ_coord	a get_gbif() output or a data.frame containing two columns named "decimal-Longitude" and "decimalLatitude".
Bioreg	'SpatialPolygonsDataFrame', 'SpatVector' or 'sf' object containing different ecoregions (convex hulls will be classified on a bioreg basis) and of CRS WGS84. Note that this parameter may be fed with an external, generated (function make_ecoregion) or in-house ecoregion shapefile. Three in-house shapefiles are already included in the library: 'eco.earth' (for terrestrial species; Nature conservancy version adapted from Olson & al. 2001), 'eco.marine' (for marine species; Spalding & al. 2007, 2012) and 'eco.fresh' (for freshwater species; Abell & al. 2008). For marine species, eco.earth may also be used if the user wants to represent the terrestrial range of species that also partially settle on mainland. For fresh water species, same may be done if the user considers that terrestrial ecoregions should be more representative of the species ecology.
Bioreg_name	Character. How is the shapefile attribute containing the ecoregion names called? Default is the very detailed level of 'eco.earth' (aka 'ECO_NAME'). Note that 'EcoRegion' must always be used when using a make_ecoregion() output. See details.

degrees_outlier	Numeric. Distance threshold (degrees) for outlier classification. If the nearest minimal distance to the next point is larger than this threshold, it will be considered as an outlier.
clustered_points_outlier	Numeric. Maximum number of points which are closer to each other than the degrees_outlier, but should still be considered as outliers.
buffer_width_point	Numeric. Buffer (in degrees) which will be applied around single observations.
buffer_increment_point_line	Numeric. How much should the buffer be increased for each point on a line.
buffer_width_polygon	Numeric. Buffer (in degrees) which will be applied around distribution polygons (for each ecoregion).
dir_temp	Character. Where should the temporary text file for the convex hull be saved? (text file will be deleted again).
raster	Logical. Should the output be a unified raster? Default is TRUE
res	Numeric. If raster = TRUE, which resolution? Final resolution in $^{\circ} = 1^{\circ}/\text{res}$ e.g., $= 0.1^{\circ}$ (i.e. $\sim 10\text{km}$) if $\text{res} = 10$. Default is 100 ($\sim 1\text{km}$). It is important to note that the highest achievable resolution of the output will depend on its 'Bioreg' precision, e.g., a species range output can reach the same resolution of the rasters used to create a 'make_ecoregion' object.

Details

Ecoregions cover relatively large areas of land or water, and contain characteristic, geographically distinct assemblages of natural communities sharing a large majority of species, dynamics, and environmental conditions. The biodiversity of flora, fauna and ecosystems that characterise an ecoregion tends to be distinct from that of other ecoregions (<https://en.wikipedia.org/wiki/Ecoregion>).

Each ecoregion shapefile has one or more categories, which describe more or less precisely the ecoregion world distribution (from the more to the less detailed):

- 'eco.earth' has three different levels: 'ECO_NAME', 'WWF_MHTNAM' and 'WWF_REALM2'.
- 'eco.fresh' has only one: 'FEOW_ID'.

- 'eco.marine' contains a mix of two types of marine ecoregions. Either common ('PROVINC' and 'REALM') or distinct levels:

—> For PPOW (Pelagic provinces of the world): 'BIOME'.

—> For MEOW (Marine ecoregions of the world): 'ECOREGION'.

Value

A 'SpatVector' or 'SpatRaster'.

References

Oskar Hagen, Lisa Vaterlaus, Camille Albouy, Andrew Brown, Flurin Leugger, Renske E. Onstein, Charles Novaes de Santana, Christopher R. Scotese, Loïc Pellissier. (2019) Mountain building, climate cooling and the richness of cold-adapted plants in the Northern Hemisphere. *Journal of Biogeography*. doi: 10.1111/jbi.13653

Lyu, L., Leugger, F., Hagen, O., Fopp, F., Boschman, L. M., Strijk, J. S., ... & Pellissier, L. (2022). An integrated high resolution mapping shows congruent biodiversity patterns of Fagales and Pinales. *New Phytologist*, 235(2), 759-772. doi: 10.1111/nph.18158

Olson, D. M., Dinerstein, E., Wikramanayake, E. D., Burgess, N. D., Powell, G. V. N., Underwood, E. C., D'Amico, J. A., Itoua, I., Strand, H. E., Morrison, J. C., Loucks, C. J., Allnutt, T. F., Ricketts, T. H., Kura, Y., Lamoreux, J. F., Wettengel, W. W., Hedao, P., Kassem, K. R. 2001. Terrestrial ecoregions of the world: a new map of life on Earth. *Bioscience* 51(11):933-938. doi: 10.1641/0006-3568(2001)051

The Nature Conservancy (2009). Global Ecoregions, Major Habitat Types, Biogeographical Realms and The Nature Conservancy Terrestrial Assessment Units. GIS layers developed by The Nature Conservancy with multiple partners, combined from Olson et al. (2001), Bailey 1995 and Wiken 1986. Cambridge (UK): The Nature Conservancy.

Mark D. Spalding, Helen E. Fox, Gerald R. Allen, Nick Davidson, Zach A. Ferdaña, Max Finlayson, Benjamin S. Halpern, Miguel A. Jorge, Al Lombana, Sara A. Lourie, Kirsten D. Martin, Edmund McManus, Jennifer Molnar, Cheri A. Recchia, James Robertson, Marine Ecoregions of the World: A Bioregionalization of Coastal and Shelf Areas, *BioScience*, Volume 57, Issue 7, July 2007, Pages 573–583. doi: 10.1641/B570707

Spalding, M. D., Agostini, V. N., Rice, J., & Grant, S. M. (2012). Pelagic provinces of the world: a biogeographic classification of the world's surface pelagic waters. *Ocean & Coastal Management*, 60, 19-30. doi: 10.1016/j.ocecoaman.2011.12.016

The Nature Conservancy (2012). Marine Ecoregions and Pelagic Provinces of the World. GIS layers developed by The Nature Conservancy with multiple partners, combined from Spalding et al. (2007) and Spalding et al. (2012). Cambridge (UK): The Nature Conservancy.

Robin Abell, Michele L. Thieme, Carmen Revenga, Mark Bryer, Maurice Kottelat, Nina Bogutskaya, Brian Coad, Nick Mandrak, Salvador Contreras Balderas, William Bussing, Melanie L. J. Stiassny, Paul Skelton, Gerald R. Allen, Peter Unmack, Alexander Naseka, Rebecca Ng, Nikolai Sindorf, James Robertson, Eric Armijo, Jonathan V. Higgins, Thomas J. Heibel, Eric Wikramanayake, David Olson, Hugo L. López, Roberto E. Reis, John G. Lundberg, Mark H. Sabaj Pérez, Paulo Petry, *Freshwater Ecoregions of the World: A New Map of Biogeographic Units for Freshwater Biodiversity Conservation*, *BioScience*, Volume 58, Issue 5, May 2008, Pages 403–414. doi: 10.1641/B580507

Hijmans, Robert J. "terra: Spatial Data Analysis. R Package Version 1.6-7." (2022). Terra - CRAN

See Also

For more information on the original code and methods, check Hagen, Oskar et al. (2019), Data from: Mountain building, climate cooling and the richness of cold-adapted plants in the northern hemisphere, Dryad, Dataset, <https://doi.org/10.5061/dryad.0ff6b04>.

Examples

```
## Not run:
# Load available ecoregions
data(ecoregions)

# First download the worldwide observations of Panthera tigris and convert to SpatialPoints
# get occurrence points from GBIF
occ <- get_gbif("Panthera tigris",
               basis=c("OBSERVATION", "HUMAN_OBSERVATION", "MACHINE_OBSERVATION"))
# make range from occurrence points
range <- get_range("Panthera tigris", occ, eco.earth, "ECO_NAME") #TODO change eco.earth to eco.land
## TODO download=TRUE by default, check presence of file.
```

```

# Plot
terra::plot(range, axes = FALSE, box = FALSE, legend=FALSE, col="chartreuse4")
# plot political world boundaries
terra::plot(rnaturalearth::ne_countries(returnclass = "sf")[1], add=T, col=NA)
graphics::points(occ[,c("decimalLongitude", "decimalLatitude")], pch=20, col="#99340470", cex=1.5)

# plot(countries,col="#bcbddc")
# plot(ne_countries(returnclass = "sf")[4] , col="antiquewhite")
# TODO FIX THIS
# points(occ[,c("decimalLongitude", "decimalLatitude")], pch=20, col="#99340470", cex=1.5)

## End(Not run)

```

get_status

Retrieve from GBIF the IUCN and taxonomy status of a specific Taxa

Description

Generates, based on a given species name, its IUCN red list status and a list of all scientific names (accepted, synonyms) found in the GBIF backbone taxonomy and used to download the data in `get_gbif()`. Children and related doubtful names not used to download the data may also be extracted.

Usage

```

get_status(
  sp_name = NULL,
  search = TRUE,
  rank = NULL,
  phylum = NULL,
  class = NULL,
  order = NULL,
  family = NULL,
  conf_match = 80,
  all = FALSE
)

```

Arguments

sp_name	Character. Species name from which the user wants to retrieve all existing GBIF names with associated taxonomy and IUCN status
search	Logical. If TRUE, the function will strictly look for the most relevant result, based on a list of names given by <code>rgbif</code> (only species, subspecies and variety allowed here), and give an error if name matching was impeded by synonym duplicates. If FALSE, the function will simply pick the first most relevant name from the list (higher taxa level than species allowed here). Also, unlike <code>search=TRUE</code> , fuzzy search (~approximative name match) is here allowed, and the 'rank', 'phylum', 'class', 'order' and 'family' parameters are optionally used only if no convincing name match is found.
rank	Character. "SPECIES", "SUBSPECIES" or "VARIETY". If NULL (default), the order of priority is (1) species, (2) subspecies and (3) variety unless "subsp." or "var." is found in 'sp_name'.

phylum	Character. Optional. What is the species' Phylum? Adds a criteria to deal with alternative name matches and select the right synonym. Available options are the GBIF Phylums (listed per Kingdom -> https://www.gbif.org/species/1). If search = FALSE, used only if no direct match is found.
class	Character. Optional. What is the species' Class? Same as above but at the finer class level. Available options are the GBIF Classes (same url). If search = FALSE, used only if no direct match is found.
order	Character. Optional. What is the species' Order? Same as above but at the finer order level. Available options are the GBIF Orders (same url). If search = FALSE, used only if no direct match is found.
family	Character. Optional. What is the species' Family? Same as above but at the finer family level. Available options are the GBIF Orders (same url). If search = FALSE, used only if no direct match is found.
conf_match	Numeric. From 0 to 100. Determine the confidence threshold of match of 'sp_name' with the GBIF backbone taxonomy. Default is 90.
all	Logical. Default is FALSE. Should all species names be retrieved or only the accepted name and its synonyms?

Value

Data.frame with nine columns: (0) Simplified name, (1) GBIF taxonomic key, (2) scientificName, (3) Backbone Taxonomy Status, (4) Genus, (5) Family, (6) Order, (7) Phylum and (8) IUCN status

References

Chamberlain, S., Oldoni, D., & Waller, J. (2022). rgbif: interface to the global biodiversity information facility API. 10.5281/zenodo.6023735

See Also

The rgbif package for additional and more general approaches on how to retrieve scientific names from the GBIF backbone taxonomy.

Examples

```
get_status("Cypripedium calceolus",all=FALSE)
get_status("Cypripedium calceolus",all=TRUE)
```

make_ecoregion

Make an ecoregion map based on input environmental variables

Description

This function may be used if the in-house ecoregion shapefiles are too coarse for a given geographic region (e.g., for local studies) or a shapefile of finer environmental details is needed. Based on several environmental layers (e.g. climate, soil and land cover), this function can generate a map of environmental regions containing n categories/classes. The classes are calculated with the 'clustering large applications' method (CLARA), which recognize patterns and relationships existing in spatial data, and classify it in clusters.

Usage

```
make_ecoregion(
  env = NULL,
  nclass = NULL,
  path = "",
  name = "",
  raster = FALSE,
  ...
)
```

Arguments

<code>env</code>	Object of class <code>SpatRaster</code> , <code>RasterBrick</code> or <code>RasterStack</code> of desired resolution, crs and extent defining the study area. Used to generate a map of clusters summarizing the environmental space of the study area.
<code>nclass</code>	Numeric, How many number of environmental classes should have the output?
<code>path</code>	Character. Folder path where the output should be saved. Default is none.
<code>name</code>	Character. If 'path' is used, should include the name of the output file (without file extension)
<code>raster</code>	Logical. Whether the output should be a raster layer. Default is FALSE.
<code>...</code>	Additonnal parameters for the function <code>clara()</code> of the <code>clutser</code> R package.

Value

A TIFF or SHP file

References

- Chauvier, Y., Zimmermann, N. E., Poggiato, G., Bystrova, D., Brun, P., & Thuiller, W. (2021). Novel methods to correct for observer and sampling bias in presence-only species distribution models. *Global Ecology and Biogeography*, 30(11), 2312-2325.
- Maechler, M., Rousseeuw, P., Struyf, A., Hubert, M., & Hornik, K. (2021). *cluster: Cluster Analysis Basics and Extensions*. R package version 2.1.2 — For new features, see the 'Changelog' file (in the package source). <https://CRAN.R-project.org/package=cluster>
- Reynolds, A. P., Richards, G., de la Iglesia, B., & Rayward-Smith, V. J. (2006). Clustering rules: A comparison of partitioning and hierarchical clustering algorithms. *Journal of Mathematical Modelling and Algorithms*, 5(4), 475–504. doi: 10.1007/s10852-005-9022-1
- Schubert, E., & Rousseeuw, P. J. (2019). Faster k-Medoids clustering: Improving the PAM, CLARA, and CLARANS algorithms. In G. Amato, C. Gennaro, V. Oria, & M. Radovanović (Eds.), *Similarity search and applications. SISAP 2019. Lecture Notes in Computer Science* (Vol. 11807, pp. 171–187). Springer.

Examples

```
#TODO transfer to data()
rst.path <- paste0(system.file(package = "gbif.range"), "/extdata/rst_enl.tif")
rst <- terra::rast(rst.path)
shp.path <- paste0(system.file(package = "gbif.range"), "/extdata/shp_lonlat.shp")
shp.lonlat <- terra::vect(shp.path)
rst <- terra::crop(rst, shp.lonlat)
#plot(crp)
```

```
# Apply the function by infering 50 classes of environments
my.eco <- make_ecoregion(rst,50)
terra::plot(my.eco)

# Downloading in the European Alps the observations of one plant species
obs.arcto <- get_gbif("Arctostaphylos alpinus",geo=shp.lonlat)

# Create the range map based on our custom ecoregion
range.arcto <- get_range("Arctostaphylos alpinus",obs.arcto,my.eco,"EcoRegion",res=20)

# Plot
terra::plot(shp.lonlat, col="grey")
terra::plot(range.arcto,add=TRUE,col=rgb(0.2,1,0.2,0.5,1))
graphics::points(obs.arcto[,c("decimalLongitude","decimalLatitude")],pch=20,col="orange1",cex=1)
```

make_tiles

Create a specific number of tiles based on a raster extent

Description

Based on a specific extent, one or several tiles are generated. Tiles can be smaller raster extents or geometry arguments POLYGON(). The original extent is therefore either converted into a POLYGON() argument, or divided into Ntiles of regular fragments which are converted into POLYGON() arguments and smaller SpatExtent.

Usage

```
make_tiles(geo, Ntiles, sext = TRUE)
```

Arguments

geo	Object of class Extent, SpatExtent, SpatialPolygon, SpatialPolygonDataframe, or SpaVector (WGS84 or planar) to define the study's area extent. Default is NULL i.e. the whole globe.
Ntiles	Numeric. In how many tiles/fragments should geo be divided approximately?
sext	Logical. Should a list of SpatExtent also be returned for each generated POLYGON()?

Value

A list of geometry arguments POLYGON() of length Ntiles (and of SpatExtent if sext=TRUE).

References

Chauvier, Y., Thuiller, W., Brun, P., Lavergne, S., Descombes, P., Karger, D. N., ... & Zimmermann, N. E. (2021). Influence of climate, soil, and land cover on plant species distribution in the European Alps. *Ecological monographs*, 91(2), e01433. 10.1002/ecm.1433

Examples

```
# Load the European Alps Extent
shp.path <- paste0(system.file(package = "gbif.range"), "/extdata/shp_lonlat.shp")
shp.lonlat <- terra::vect(shp.path)

# Apply the function to divide the extent in ~20 fragments
mt = make_tiles(geo=shp.lonlat, Ntiles=20, sext=TRUE); mt
```

obs_filter

Filter a set of GBIF observations according to a defined grain

Description

Whereas the 'grain' parameter in `get_gbif()` allows GBIF observations to be filtered according to a certain spatial precision, `obs_filter()` accepts as input a `get_gbif()` output (one or several species) and filter the observations according to a given grid resolution (one observation per pixel grid kept). This function allows the user to refine the density of GBIF observations according to a defined analysis/study's resolution.

Usage

```
obs_filter(gbifs, grid)
```

Arguments

gbifs	One <code>get_gbif()</code> output including one or several species. Note that if GBIF absences are kept in the output(s), the function should be used distinctively for observations and absences.
grid	Object of class <code>SpatRaster</code> , <code>RasterLayer</code> , <code>RasterBrick</code> or <code>RasterStack</code> of desired resolution and extent (WGS84).

Value

Data frame with two columns named 'x' and 'y' comprising the new set of observations filtered at grid resolution.

Examples

```
#read example shapefile
shp.path <- paste0(system.file(package = "gbif.range"), "/extdata/shp_lonlat.shp")
shp.lonlat <- terra::vect(shp.path)
rst.path <- paste0(system.file(package = "gbif.range"), "/extdata/rst_enl.tif")
rst <- terra::rast(rst.path)

# Downloading in the European Alps the observations of two plant species
obs.arcto <- get_gbif("Arctostaphylos alpinus", geo=shp.lonlat)
obs.saxi <- get_gbif("Saxifraga cernua", geo=shp.lonlat)
terra::plot(shp.lonlat)
graphics::points(obs.arcto[, c("decimalLongitude", "decimalLatitude")], pch=20, col="#238b4550", cex=1)
graphics::points(obs.saxi[, c("decimalLongitude", "decimalLatitude")], pch=20, col="#99000d50", cex=1)

# rbind both datasets
```

```
both.sp <- rbind(obs.arcto,obs.saxi)

# Run function
obs.filt <- obs_filter(both.sp,rst)

# Check new points
terra::plot(shp.lonlat)

graphics::points(obs.filt[obs.filt$Species%in%"Arctostaphylos alpinus",c("x","y")],
pch=20,col="#238b4550",cex=1)

graphics::points(obs.filt[obs.filt$Species%in%"Saxifraga cernua",c("x","y")],
pch=20,col="#99000d50",cex=1)
```

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