gbif.range

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Title A toolbox to efficiently download and filter large GBIF observational datasets for sound spatial analyses
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Description Workflow to generate taxa range maps from scratch based on ecoregions and an user-friendly GBIF wrapper (no hard-limit of < 100,000 species observations, filetring parameters to easily flag records, access to the GBIF backbone taxonomy)
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R topics documented:
conv_function 2 get_gbif 2 get_range 5 make_tiles 6 wsl_doi 7 wsl_obs_filter 9 wsl_taxonomy 10
Index 11

2 get_gbif

conv_function

Create polygon objects in different bioregions

Description

Not to be called directly by the user

Usage

```
conv_function(sp_coord, bwp, bipl, bwpo, temp_dir, g = NULL)
```

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 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 succesive 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_search() (implemented rgbif function) is set to FALSE.

Usage

```
get_gbif(
  sp_name = NULL,
  conf_match = 90,
  geo = NULL,
  grain = 1000,
  duplicates = FALSE,
  absences = FALSE,
 no_xy = FALSE,
 basis = c("OBSERVATION", "HUMAN_OBSERVATION", "MACHINE_OBSERVATION",
  "MATERIAL_SAMPLE", "PRESERVED_SPECIMEN", "FOSSIL_SPECIMEN", "LIVING_SPECIMEN",
    "LITERATURE", "UNKNOWN"),
  add_infos = NULL,
  time_period = c(1000, 3000),
  identic_xy = FALSE,
 wConverted_xy = FALSE,
  centroids = FALSE,
```

get_gbif 3

```
ntries = 10,
error.skip = TRUE,
...
)
```

Arguments

sp_name Character. Scientific name to run an online search (i.e. with GBIF-API) for

species observations. Works also for genus and higher taxa levels.

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 'SpaVector' (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 (x2)

according to their uncertainties and number of coordinate decimals. Records with no information on coordinate uncertainties (column 'coordinateUncertain-

tyInMeters') are be 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, only records with no coordinates are downloaded.

basis Character. Which basis of records should be selected? Default is all i.e. c("OBSERVATION",

"HUMAN_OBSERVATION", "MACHINE_OBSERVATION", "MATERIAL_SAMPLE",

 $"PRESERVED_SPECIMEN", "FOSSIL_SPECIMEN", "LIVING_SPECIMEN", \\$

"LITERATURE", "UNKNOWN"). Description may be found here: https://docs.gbif.org/course-

data-use/en/basis-of-record.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', '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 lon/lat converted xy 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 attempts failed, an empty data frame is return

by default.

error.skip Logical. Should the search process continues if ntries failed?

. . . Additional parameters for the function cd_round() of CoordinateCleaner.

4 get_gbif

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 110 km < grain <= 11 km, lon/lat with >= 1 decimal are kept, if 110 km < grain <= 1100 m, lon/lat with >= 2 decimals kept; if 1100 m < grain <= 110 m, lon/lat with >= 4 decimals are kept; if 11 m < grain <= 1.1 m, 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, addtional 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) CoordinateCelaner packages for additional and more general approaches on (1) downloading GBIF observations and (2) post-filetering those.

```
# Load maptools for the map world
library(maptools)
data(wrld_simpl)

# Load the Alps Extend
data(geo_dat)

# Downloading worldwide the observations of Panthera tigris
test1 = get_gbif("Panthera tigris", basis=c("OBSERVATION", "HUMAN_OBSERVATION"))
plot(wrld_simpl)
points(test1[,c("decimalLongitude", "decimalLatitude")],pch=20,col="#238b4550",cex=4)

# Downloading in the Alps the observations of Cypripedium calceolus (with a 100m grain and
# by adding the 'issues' column)
test2 = get_gbif("Cypripedium calceolus", geo = shp.lonlat, grain = 100, add_infos = c("issue"))
plot(shp.lonlat)
```

get_range 5

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 bioregions. 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 bioregion. If there there is only one observation in a bioregion, a buffer around this point will be created. If all points in a bioregion 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.

Usage

```
get_range(
   species_name = NULL,
   occ_coord = NULL,
   Bioreg = NULL,
   Bioreg_name = NULL,
   degrees_outlier = 10,
   clustered_points_outlier = 3,
   buffer_width_point = 0.5,
   buffer_increment_point_line = 0.5,
   buffer_width_polygon = 0.1,
   dir_temp = paste0("temp", sample(1:99999999, 1))
)
```

Arguments

species_name character string of the species name. E.g. "Anemone nemorosa".

occ_coord a SpatialPoints object.

Bioreg shapefile containg different bioregions (convex hulls will be classified on a bioreg

basis).

Bioreg_name how is the slot containing the bioregion names called?

6 make_tiles

degrees_outlier

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

maximum number of points which are closer to each other than the degrees_outlier, but should still be considered as outliers.

buffer_width_point

buffer (in degrees) which will be applied around single observations.

buffer_increment_point_line

how much should the buffer be increased for each point on a line.

buffer_width_polygon

buffer (in degrees) which will be applied around distribution polygons (for each bioregion).

dir_temp where should the temporary text file for the convex hull be saved? (text file will be deleted again).

Details

...

Value

a shapefile.

References

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 10.1111/nph.18158

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

See Also

...

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)
```

wsl_doi 7

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 POLY-GON()?

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
data(geo_dat)

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

wsl_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 wsl_gbif() outputs.

Usage

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

8 wsl_doi

Arguments

wsl_gbif data.frame or list. One wsl_gbif() output or a list of several.

title The title for your derived dataset.

source_url A link to where the dataset is stored.

user Your GBIF username.

pwd Your GBIF password.

... Additional parameters for derived_dataset() in rgbif.

descritpion A description of the dataset.

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

wsl_obs_filter 9

wsl_obs_filter

Filter a set of GBIF observations according to a defined grain

Description

Whereas the 'grain' parameter in wsl_gbif() allows GBIF observations to be filtered according to a certain spatial precision, wsl_obs_filter() accepts as input a wsl_gbif() output (one or several species) and filter the observations according to a specific 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

```
wsl_obs_filter(wsl_gbif, grid)
```

Arguments

wsl_gbif one wsl_gbif() output including one or several species. Note that if GBIF ab-

sences are kept in the output(s), the function should be used distinctively for

observations and absences.

grid Object of class 'SpatRaster', 'RasterLayer', 'RasterBrick' or 'RasterStack' of

desired resolution and extent (WGS84)

Value

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

```
# Load the European Alps extent and a raster of a random resolution
data(geo_dat)
data(exrst)
# Downloading in the European Alps the observations of two plant species
obs.arcto = wsl_gbif("Arctostaphylos alpinus",geo=shp.lonlat)
obs.saxi = wsl_gbif("Saxifraga cernua",geo=shp.lonlat)
plot(vect(shp.lonlat))
points(obs.arcto[,c("decimalLongitude","decimalLatitude")],pch=20,col="#238b4550",cex=1)
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 = wsl_obs_filter(both.sp,rst)
# Check new points
x11();plot(vect(shp.lonlat))
points(obs.filt[obs.filt$Species%in%"Arctostaphylos alpinus",c("x","y")],pch=20,col="#238b4550",cex=1)
points(obs.filt[obs.filt$Species%in%"Saxifraga cernua",c("x","y")],pch=20,col="#99000d50",cex=1)
```

10 wsl_taxonomy

wsl_taxonomy	Retrieve from GRI	F all scientific names	of a specific Taxa
WSI_caxonomy	Rentere from ODI	1 an scientific names	of a specific rana

Description

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

Usage

```
wsl_taxonomy(sp_name = NULL, conf_match = 90, all = FALSE)
```

Arguments

sp_name	Character. Species name from which the user wants to retrieve all existing GBIF names
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

A data.frame with two columns: (1) Names and (2) Backbone Taxonomy 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.

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

Index

```
conv_function, 2
get_gbif, 2
get_range, 5
make_tiles, 6
wsl_doi, 7
wsl_obs_filter, 9
wsl_taxonomy, 10
```