Package 'crestr'

January 22, 2025

```
Title A Probabilistic Approach to Reconstruct Past Climates Using Palaeoecological Datasets
Version 1.4.0
Description Applies the CREST climate reconstruction
      method. It can be used using the calibration data that can be obtained
     through the package or by importing private data. An ensemble of
     graphical outputs were designed to facilitate the use of the
     package and the interpretation of the results. More information can
     be obtained from Chevalier (2022) <doi:10.5194/cp-18-821-2022>.
License MIT + file LICENSE
Encoding UTF-8
LazyData true
Roxygen list(markdown = TRUE)
RoxygenNote 7.3.1
Imports clipr,
     DBI,
     data.table,
     methods,
     openxlsx,
     plot3D,
     plyr,
     RPostgres,
     RSQLite,
     scales,
     stringr,
     terra,
     viridis
Suggests knitr,
     rmarkdown,
     pals
Depends R (>= 2.10)
Language en-us
VignetteBuilder knitr
URL https://github.com/mchevalier2/crestr, https:
      //www.manuelchevalier.com/crestr/index.html
```

BugReports https://github.com/mchevalier2/crestr/issues

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Description

Return the list of oceans and seas.

Usage

```
accBasinNames(basin = NA)
```

Arguments

basin

A name of basin. Default is NA and returns a list of all the accepted names.

Value

A list of accepted names.

See Also

https://www.marineregions.org/downloads.php

```
accBasinNames()
accBasinNames('Indian Ocean')
```

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accClimateVariables

Describes all the variables available in the database.

Description

Provides the index and the short and full names of all the variables available in the database.

Usage

```
accClimateVariables(v = NA, domain = NA)
```

Arguments

v The name of a variable to quickly access its description and ID (default NA re-

turns all possible values).

domain ('Terrestrial' or 'Marine') of the variables. Default value is NA and

both terrestrial and marine climate variable names are returned.

Value

A data frame descriptive of the climate variables available in the database (if v=NA) or the description of variable v.

See Also

https://www.worldclim.org/data/bioclim.html for details on the 'bio' data, https://csidotinfo.wordpress.com/2019/01/24/global-aridity-index-and-potential-evapotranspiration-climate-database for details on 'ai', https://www.ncei.noaa.gov/products/world-ocean-atlas for details on the sea surface temperature, sea surface salinity and nutrient Concentration data, or https://psl.noaa.gov/data/gridded/data.noaa.oisst.v2.highres.html for the sea ice concentration data.

Examples

```
## Not run:
    accClimateVariables()
    accClimateVariables(v='bio12')
## End(Not run)
```

accCountryNames

Return the list of the continents and associated countries.

Description

Return the list of the continents and associated countries.

```
accCountryNames(continent = NA)
```

accRealmNames 5

Arguments

continent A name of continent. Default is NA and returns a list of all the country names

sorted by continent.

Value

A list where each element is a vector of corresponding country names.

See Also

https://www.naturalearthdata.com/downloads/10m-cultural-vectors/10m-admin-0-countries/

Examples

```
accCountryNames()
accCountryNames('Europe')
```

accRealmNames

Return the list of the realms and associated biomes and ecoregions.

Description

Return the list of the realms and associated biomes and ecoregions.

Usage

```
accRealmNames(realm = NA, ecoregion = TRUE)
```

Arguments

realm A name of accepted realm. Default is NA and returns a list of all the biome and

ecoregion names sorted by realm.

ecoregion A boolean to choose whether to get the ecoregions names.

Value

A list with elements that correspond to the biomes (and possibly ecoregions) of each realm.

See Also

```
https://www.worldwildlife.org/publications/terrestrial-ecoregions-of-the-world
```

```
accRealmNames(realm='Africotropical')
accRealmNames(realm='Africotropical', ecoregion=FALSE)
```

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calib_clim_space

Calibrate the distribution of the modern climate space.

Description

Calibrate the distribution of the modern climate space.

Usage

```
calib_clim_space(climate, bin_width)
```

Arguments

climate A vector of climatic values where the species is present.

bin_width The width of the bins used to correct for unbalanced climate state. Use values

that split the studied climate gradient in 15-25 classes (e.g. 2°C for temperature

variables). Default is 1.

Value

A ccs object that will be used by fit_pdfsp.

Examples

```
calib_clim_space(sample(0:300 / 10, 4000, replace = TRUE), 2)
```

check_coordinates

Check if the coordinates are correct.

Description

Check if the coordinates are correct.

Usage

```
check_coordinates(xmn, xmx, ymn, ymx)
```

Arguments

```
xmn, xmx, ymn, ymx
```

The coordinates defining the study area.

Value

Return a set of valid coordinates.

```
check_coordinates(NA, NA, NA, NA)
check_coordinates(-200, 0, 0, 90)
check_coordinates(20, 0, 90, 0)
```

cite_climate_data 7

cite_climate_data	cite_climate_data	
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Description

Returns the references associated with the climate data used to fit the pdfs.

Usage

```
cite_climate_data(x, verbose = TRUE)
```

Arguments

x A crestObj produced by one of the crest, crest.get_modern_data, crest.calibrate,

crest.reconstruct or loo functions.

verbose A boolean to print non-essential comments on the terminal (default TRUE).

Value

A list of references to add if the data generated by crestr are published.

cite_crest Returns the list of references associated to the reconstruction.

Description

Returns the list of references associated to the reconstruction.

Usage

```
cite_crest(x, verbose = TRUE)
```

Arguments

verbose

 $x \\ A \verb|crestObj| produced by one of the \verb|crest|, \verb|crest.get_modern_data|, \verb|crest.calibrate|, \\$

A boolean to print non-essential comments on the terminal (default TRUE).

crest.reconstruct or loo functions.

Value

A list of references to add if the data generated by crestr are published.

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cite_distrib_data Returns the references associated with the GBIF data used to fit the pdfs.	!
--	---

Description

Returns the references associated with the GBIF data used to fit the pdfs.

Usage

```
cite_distrib_data(x, verbose = TRUE)
```

Arguments

x A crestObj produced by one of the crest, crest.get_modern_data, crest.calibrate,

crest.reconstruct or loo functions.

verbose A boolean to print non-essential comments on the terminal (default TRUE).

Value

A list of references to add if the data generated by crestr are published.

cite_method Returns the references associated with the development of CREST.

Description

Returns the references associated with the development of CREST.

Usage

```
cite_method(x, verbose = TRUE)
```

Arguments

verbose

x A crestObj produced by one of the crest, crest.get_modern_data, crest.calibrate, crest.reconstruct or loo functions.

A boolean to print non-essential comments on the terminal (default TRUE).

Value

A list of references to add if the data generated by crestr are published.

climate_from_xy 9

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Extract The climate values associated to a set of coordinates.

Description

Extract The climate values associated to a set of coordinates.

Usage

```
climate_from_xy(
  long,
  lat,
  climate = accClimateVariables()[, 2],
  resol = 0.25,
  dbname = "gbif4crest_02"
)
```

Arguments

long The longitude of the site.lat The latitude of the site.

climate The climate variables to extract the values from. Returns all possible values by

default.

resol The resolution of the target climatology (default 0.25 degrees).

dbname The name of the data source database.

Value

A data frame containing the climate values.

Examples

```
## Not run:
   climate_from_xy(50, 10, c('bio1', 'ai'))
   climate_from_xy(50, 10)
## End(Not run)
```

close_db_connection

Disconnect the database connection.

Description

Disconnect the database connection.

```
close_db_connection(db)
```

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Arguments

db

An active database connection

Value

No return value, function called to close the connection to the database.

Examples

```
## Not run:
   db <- connect_online()
   close_db_connection(db)
## End(Not run)</pre>
```

colour_theme

Returns a vector of colours

Description

Returns a vector of colours

Usage

```
colour_theme(n)
```

Arguments

n

An index to select the colour theme

Value

A vector of colours.

```
colour_theme(1)
```

connect_local_sqlite3 11

connect_local_sqlite3 Connect to the gbif4crest calibration database

Description

Connect to the gbif4crest_02 database using a local SQLite3 copy.

Usage

```
connect_local_sqlite3(dbname = "gbif4crest_02.sqlite3")
```

Arguments

dbname

The complete path to the SQLite3 file. The name should end by '.sqlite3'

Value

An active connection to a database

See Also

The SQLite3 database can be downloaded from https://figshare.com/articles/dataset/GBIF_for_CREST_database/6743207.

Examples

```
## Not run:
   db <- connect_online()
## End(Not run)</pre>
```

connect_online

Connect to the gbif4crest calibration database

Description

Connect to the gbif4crest_02 database by accessing the server on Amazon.

```
connect_online(
  dbname = "gbif4crest_02",
  port = 5432,
  host = "gbif4crest.cvqgy2mnjwtg.eu-west-3.rds.amazonaws.com",
  user = "guestuser",
  password = "pwd12345"
)
```

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Arguments

dbname The name of the database. Default is 'gbif4crest_02'.

port The port to connect to the server. Default is 5432.

host The host of the database server. Default is 'gbif4crest.cvqgy2mnjwtg.eu-west-3.rds.amazonaw

user The user name to use to connect. Default is 'guestuser'.

password The password associated with the user name. Default is 'pwd12345'.

Value

An active connection to a database

Examples

```
## Not run:
   db <- connect_online()
## End(Not run)</pre>
```

convert2percentages

Convert abundance data into percentage data.

Description

Convert abundance data into percentage data.

Usage

```
convert2percentages(df, col2convert = 2:ncol(df))
```

Arguments

df The dataframe containing the data to convert.

col2convert A vector of the columns to convert. Default is all the columns but the first, which

contains an age, a depth or a sampleID.

Value

A vector of unique taxonIDs.

```
df <- data.frame(matrix(1:25, ncol = 5))
colnames(df) <- paste(rep("col", 5), 1:5, sep = "")
convert2percentages(df)
convert2percentages(df, col2convert = 3:5)</pre>
```

convert2presenceAbsence

Convert data into presence/absence data.

Description

Convert data into presence/absence data.

Usage

```
convert2presenceAbsence(df, threshold = 2, col2convert = 2:ncol(df))
```

Arguments

df The dataframe containing the data to convert.

threshold The threshold that defines presence (presence if >= threshold)

col2convert A vector of the columns to convert. Default is all the columns but the first, which

contains an age, a depth or a sampleID.

Value

A vector of unique taxonIDs.

Examples

```
df <- data.frame(matrix(1:25, ncol = 5))
colnames(df) <- paste(rep("col", 5), 1:5, sep = "")
convert2presenceAbsence(df, threshold = 15)
convert2presenceAbsence(df, col2convert = 3:5)</pre>
```

copy_crest

Copy crest data to the clipboard.

Description

Copy crest data to the clipboard for an easy extraction of the data from the R environment.

```
copy_crest(
   x,
   climate = x$parameters$climate,
   optima = TRUE,
   mean = FALSE,
   uncertainties = FALSE
)
```

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Arguments

A crestObj produced by the crest.reconstruct or crest functions.

A vector of the climate variables to extract. See accClimateVariables for the list of accepted values.

A boolean value to indicate if the optima should be copied to the clipboard.

mean A boolean value to indicate if the means should be copied to the clipboard.

uncertainties A boolean value to indicate if the uncertainties should be copied to the clipboard.

Value

No return value. This function is called to copy the crest data to the clipboard.

Examples

```
## Not run:
if(requireNamespace('clipr', quietly=TRUE)) {
    reconstr <- crest(
      df = crest_ex, pse = crest_ex_pse, taxaType = 0,
            climate = c("bio1", "bio12"), bin_width = c(2, 20),
            shape = c("normal", "lognormal"),
            selectedTaxa = crest_ex_selection, dbname = "crest_example",
            leave_one_out = TRUE
    )
            copy_crest(reconstr, uncertainties=TRUE)
            ## You can now paste the values in a spreadsheet.
}

## End(Not run)</pre>
```

createPSE

Creates a spreadsheet with the format required for a PSE.

Description

Creates a spreadsheet with the format required for a PSE from a list of taxa.

Usage

```
createPSE(taxa, loc = "proxy_species_equivalency.xlsx")
```

Arguments

taxa A list of taxa to include in the PSE file.

An absolute or relative path that indicates where the spreadsheet should be saved. Also used to specify the name of the file. Default: the file is saved in

the working directory under the name proxy_species_equivalency.xlsx.

Value

No return value, called to create a PSE file.

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Examples

crest

A wrapper for all the crest functions.

Description

Runs all the different steps of a CREST reconstruction in one function.

```
crest(
  df,
  climate,
  pse = NA,
  taxaType = 0,
  distributions = NA,
  site_info = rep(NA, length(climate)),
  site_name = NA,
  xmn = NA,
  xmx = NA,
  ymn = NA,
  ymx = NA,
  continents = NA,
  countries = NA,
  realms = NA,
  biomes = NA,
  ecoregions = NA,
  minGridCells = 20,
  selectedTaxa = NA,
  bin_width = rep(1, length(x$parameters$climate)),
  shape = rep("normal", length(x$parameters$climate)),
  npoints = 500,
  ai.sqrt = FALSE,
  geoWeighting = TRUE,
  climateSpaceWeighting = TRUE,
  climateSpaceWeighting.type = "linear",
  presenceThreshold = 0,
  taxWeight = "normalisation",
  uncertainties = c(0.5, 0.95),
  leave_one_out = FALSE,
  verbose = TRUE,
  dbname = "gbif4crest_02"
)
```

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Arguments

df A data frame containing the data to reconstruct (counts, percentages or pres-

ence/absence data).

climate A vector of the climate variables to extract. See accClimateVariables for the

list of accepted values.

pse A pollen-Species equivalency table. See createPSE for details.

taxaType A numerical index (between 1 and 6) to define the type of palaeoproxy used: 1

for plants, 2 for beetles, 3 for chironomids, 4 for foraminifers, 5 for diatoms and 6 for rodents. The example dataset uses taxaType=0 (pseudo-data). Default is 1.

distributions A dataframe containing the presence records of the studied proxies and their

associated climate values.

site_info A vector containing the coordinates of the study site. Default c(NA, NA).

site_name The name of the dataset (default NA).

xmn, xmx, ymn, ymx

The coordinates defining the study area.

 $continents \qquad \quad A \ vector \ of \ the \ continent \ names \ defining \ the \ study \ area.$

countries A vector of the country names defining the study area.

realms A vector of the studied botanical realms defining the study area.

biomes A vector of the studied botanical biomes defining the study area.

ecoregions A vector of the studied botanical ecoregions defining the study area.

minGridCells The minimum number of unique presence data necessary to estimate a species'

climate response. Default is 20.

selectedTaxa A data frame assigns which taxa should be used for each variable (1 if the taxon

should be used, 0 otherwise). The colnames should be the climate variables' names and the rownames the taxa names. Default is 1 for all taxa and all vari-

ables.

bin_width The width of the bins used to correct for unbalanced climate state. Use values

that split the studied climate gradient in 15-25 classes (e.g. 2°C for temperature

variables). Default is 1.

shape The imposed shape of the species pdfs. We recommend using 'normal' for tem-

perature variables and 'lognormal' for the variables that can only take positive

values, such as precipitation or aridity. Default is 'normal' for all.

npoints The number of points to be used to fit the pdfs. Default 200.

ai.sqrt A boolean to indicate whether ai values should be square-root transformed (de-

fault FALSE).

geoWeighting A boolean to indicate if the species should be weighting by the square root

of their extension when estimating a genus/family level taxon-climate relation-

ships.

climateSpaceWeighting

A boolean to indicate if the species pdfs should be corrected for the modern

distribution of the climate space (default TRUE).

climateSpaceWeighting.type

A correction factor for the clame space weighting correction to limit the edge

effects. Either 'linear' (default), 'sqrt' or 'log'.

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presenceThreshold

All values above that threshold will be used in the reconstruction (e.g. if set at 1, all percentages below 1 will be set to 0 and the associated presences discarded).

Default is 0.

taxWeight One value among the following: 'originalData', 'presence/absence', 'percent-

ages' or 'normalisation' (default).

uncertainties A (vector of) threshold value(s) indicating the error bars that should be calcu-

lated (default both 50 and 95% ranges).

leave_one_out A boolean to indicate whether the leave one out (loo) reconstructions should be

computed (default FALSE).

verbose A boolean to print non-essential comments on the terminal (default TRUE).

dbname The name of the database. Default is 'gbif4crest_02'.

Value

A crestObj containing the reconstructions.

Examples

```
## Not run:
    data(crest_ex)
    data(crest_ex_pse)
    data(crest_ex_selection)
    reconstr <- crest(
        df = crest_ex, pse = crest_ex_pse, taxaType = 0,
            site_info = c(7.5, 7.5), site_name = 'crest_example',
        climate = c("bio1", "bio12"), bin_width = c(2, 50),
        shape = c("normal", "lognormal"),
        selectedTaxa = crest_ex_selection, dbname = "crest_example",
        leave_one_out = TRUE,
        verbose = FALSE
    )
    plot(reconstr)
    plot_loo(reconstr)

## End(Not run)</pre>
```

crest.calibrate

Fit the species and proxy pdfs

Description

This function fits the climate response of the selected taxa to the selected climate variables.

```
crest.calibrate(
   x,
   bin_width = x$parameters$bin_width,
   shape = x$parameters$shape,
```

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```
npoints = x$parameters$npoints,
  geoWeighting = x$parameters$geoWeighting,
  climateSpaceWeighting = x$parameters$climateSpaceWeighting,
  climateSpaceWeighting.type = x$parameters$climateSpaceWeighting.type,
  verbose = TRUE
)
```

Arguments

x A crestObj produced by the crest.get_modern_data function.

bin_width The width of the bins used to correct for unbalanced climate state. Use values

that split the studied climate gradient in 15-25 classes (e.g. 2°C for temperature

variables). Default is 1.

shape The imposed shape of the species pdfs. We recommend using 'normal' for tem-

perature variables and 'lognormal' for the variables that can only take positive

values, such as precipitation or aridity. Default is 'normal' for all.

npoints The number of points to be used to fit the pdfs. Default 200.

geoWeighting A boolean to indicate if the species should be weighting by the square root

of their extension when estimating a genus/family level taxon-climate relation-

ships.

climateSpaceWeighting

A boolean to indicate if the species pdfs should be corrected for the modern

distribution of the climate space (default TRUE).

climateSpaceWeighting.type

A correction factor for the clame space weighting correction to limit the edge

effects. Either 'linear' (default), 'sqrt' or 'log'.

verbose A boolean to print non-essential comments on the terminal (default TRUE).

Value

A crest0bj object containing the spatial distributions and the climate space.

```
## Not run:
  data(crest_ex_pse)
  data(crest_ex_selection)
  data(crest_ex)
  x <- crest.get_modern_data( df = crest_ex,</pre>
    pse = crest_ex_pse, taxaType = 0,
    climate = c("bio1", "bio12"),
    selectedTaxa = crest_ex_selection, dbname = "crest_example",
    verbose = FALSE
  )
  x <- crest.calibrate(x,</pre>
    geoWeighting = TRUE, climateSpaceWeighting = TRUE,
    bin_width = c(2, 50), shape = c("normal", "lognormal"),
    verbose = FALSE
  )
## End(Not run)
```

crest.get_modern_data Extract distributions from the database

Description

This function will extract the distributions of all the species composing each taxon and return them as a list.

Usage

```
crest.get_modern_data(
  pse,
  taxaType,
  climate,
  df = NA,
  ai.sqrt = FALSE,
  xmn = NA,
  xmx = NA,
  ymn = NA,
  ymx = NA,
  continents = NA,
  countries = NA,
  basins = NA,
  sectors = NA,
  realms = NA,
  biomes = NA,
  ecoregions = NA,
  minGridCells = 20,
  climateWithObs = FALSE,
  elev_min = NA,
  elev_max = NA,
  elev_range = NA,
  year_min = 1900,
  year_max = 2021,
  nodate = TRUE,
  type_of_obs = c(1, 2, 3, 8, 9),
  selectedTaxa = NA,
  site_info = c(NA, NA),
  site_name = NA,
  dbname = "gbif4crest_02",
  verbose = TRUE
)
```

Arguments

A pollen-Species equivalency table. See createPSE for details.

A numerical index (between 1 and 6) to define the type of palaeoproxy used: 1 for plants, 2 for beetles, 3 for chironomids, 4 for foraminifers, 5 for diatoms and 6 for rodents. The example dataset uses taxaType=0 (pseudo-data). Default is 1.

Climate A vector of the climate variables to extract. See accClimateVariables for the list of accepted values.

df A data frame containing the data to reconstruct (counts, percentages or pres-

ence/absence data).

ai.sqrt A boolean to indicate whether ai values should be square-root transformed (de-

fault FALSE).

xmn, xmx, ymn, ymx

The coordinates defining the study area.

continents A vector of the continent names defining the study area.

A vector of the country names defining the study area.

A vector of the ocean names defining the study area.

sectors A vector of the marine sector names defining the study area.

realms A vector of the studied botanical realms defining the study area.

biomes A vector of the studied botanical biomes defining the study area.

ecoregions A vector of the studied botanical ecoregions defining the study area.

minGridCells The minimum number of unique presence data necessary to estimate a species'

climate response. Default is 20.

 ${\tt climateWithObs} \quad A \ boolean \ to \ indicate \ whether \ all \ climate \ values \ from \ the \ calibration \ dataset \ be$

included (FALSE, default) or only the climate values that corresponds to proxy observations (TRUE). Only useful in if the climate space weighting is activated.

elev_min, elev_max

Parameters to only selected grid cells with an elevation higher than elev_min or

lower than elev_max (default is 'NA).

elev_range Parameters discard the grid cell with a high elevation range (default is NA).

year_min, year_max

The oldest and youngest occurrences accepted (default is 1900-2021).

nodate A boolean to accept occurrences without a date (can overlap with occurrences

with a date; default TRUE).

type_of_obs The type of observation to use in the study. 1: human observation, 2: observa-

tion, 3: preserved specimen, 4: living specimen, 5: fossil specimen, 6: material sample, 7: machine observation, 8: literature, 9: unknown (Default c(1, 2, 3,

8,9))

selectedTaxa A data frame assigns which taxa should be used for each variable (1 if the taxon

should be used, 0 otherwise). The colnames should be the climate variables' names and the rownames the taxa names. Default is 1 for all taxa and all vari-

ables.

site_info A vector containing the coordinates of the study site. Default c(NA, NA).

site_name The name of the dataset (default NA).

dbname The name of the database. Default is 'gbif4crest_02' and data will be ex-

tracted from the online database. The SQLite3 version of the database can also be used here by providing the complete path to a file ending by .sqlite3, e.g.

/path/to/file/gbif4crest_02.sqlite3

verbose A boolean to print non-essential comments on the terminal (default TRUE).

Value

A crest0bj object containing the spatial distributions.

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See Also

The SQLite3 database can be downloaded from https://figshare.com/articles/dataset/GBIF_for_CREST_database/6743207.

Examples

crest.reconstruct

Reconstruct climate from fossil data

Description

This function fits the climate response of the selected taxa to the selected climate variables.

Usage

```
crest.reconstruct(
   x,
   presenceThreshold = 0,
   taxWeight = "normalisation",
   uncertainties = c(0.5, 0.95),
   skip_for_loo = FALSE,
   verbose = TRUE
)
```

Arguments

x A crestObj produced by the crest.calibrate function.

presenceThreshold

All values above that threshold will be used in the reconstruction (e.g. if set at 1, all percentages below 1 will be set to 0 and the associated presences discarded). Default is 0.

One value among the following: 'originalData', 'presence/absence', 'percentages' or 'normalisation' (default).

uncertainties A (vector of) threshold value(s) indicating the error bars that should be calculated (default healt 50 and 05% reason)

lated (default both 50 and 95% ranges).

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skip_for_loo A boolean that tells the loo function to skip parts and fasten the process. Not for users, always leave to FALSE.

verbose A boolean to print non-essential comments on the terminal (default TRUE).

Value

A crest0bj object containing the reconstructions and all the associated data.

Examples

```
data(crest_ex_pse)
data(crest_ex_selection)
data(crest_ex)
## Not run:
  x <- crest.get_modern_data( df = crest_ex,</pre>
    pse = crest_ex_pse, taxaType = 0,
    climate = c("bio1", "bio12"),
    selectedTaxa = crest_ex_selection, dbname = "crest_example",
    verbose = FALSE
  )
  x <- crest.calibrate(x,</pre>
    geoWeighting = TRUE, climateSpaceWeighting = TRUE,
    bin_width = c(2, 50), shape = c("normal", "lognormal"),
    verbose = FALSE
  x <- crest.reconstruct(x,</pre>
    verbose = FALSE)
  plot(x)
## End(Not run)
```

crest.set_modern_data Format a crestObj with private data.

Description

Format a crestObj with private data.

```
crest.set_modern_data(
    distributions,
    climate,
    df = NA,
    climate_space = NA,
    weight = FALSE,
    minGridCells = 20,
    selectedTaxa = NA,
    site_info = c(NA, NA),
    site_name = NA,
    site_climate = rep(NA, length(climate)),
    verbose = TRUE
)
```

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Arguments

distributions	A dataframe containing the presence records of the studied proxies and their associated climate values.
climate	A vector of the climate variables to extract. See accClimateVariables for the list of accepted values.
df	A data frame containing the data to reconstruct (counts, percentages or presence/absence data).
climate_space	A dataframe of climate values across the study area useful to correct for the imbalance of the sampling data (see 'crest.calibrate for more details). Default is NA.
weight	The records in the distributions can be weighted using the percentages by setting weight=TRUE. Include a column called 'weight' in the distributions table.
minGridCells	The minimum number of unique presence data necessary to estimate a species' climate response. Default is 20.
selectedTaxa	A data frame assigns which taxa should be used for each variable (1 if the taxon should be used, 0 otherwise). The colnames should be the climate variables' names and the rownames the taxa names. Default is 1 for all taxa and all variables.
site_info	A vector containing the coordinates of the study site. Default c(NA, NA).
site_name	The name of the dataset (default NA).
site_climate	The climate values at the location of the dataset '(default NA).
verbose	A boolean to print non-essential comments on the terminal (default TRUE).

Value

A crest0bj object containing the spatial distributions.

```
#> Reformating the example dataset to fit this function
distributions <- cbind('ProxyName' = rep('Taxon1', nrow(reconstr$modelling$distributions[[1]])),</pre>
                        reconstr$modelling$distributions[[1]],
                        stringsAsFactors = FALSE)
for(tax in names(reconstr$modelling$distributions)[-1]) {
  distributions <- rbind(distributions,</pre>
              cbind('ProxyName'= rep(tax, nrow(reconstr$modelling$distributions[[tax]])),
                            reconstr$modelling$distributions[[tax]],
                            stringsAsFactors = FALSE)
distributions <- distributions[, c(2,1,3:6)]</pre>
print(head(distributions))
climate_space <- reconstr$modelling$climate_space</pre>
print(head(climate_space))
x <- crest.set_modern_data(distributions, df=crest_ex,</pre>
                            climate = c("bio1", "bio12"))
x <- crest.set_modern_data(distributions, df=crest_ex,</pre>
                            climate_space=climate_space,
                            climate = c("bio1", "bio12"))
```

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crest.simplify

Simplify a crestObj into a dataframe.

Description

Simplify a crestObj with reconstructed values into a dataframe.

Usage

```
crest.simplify(x, optima = TRUE)
```

Arguments

```
x A crestObj produced by the crest, crest.reconstruct or loo functions.

Optima A boolean to indicate whether to plot the optimum (TRUE) or the mean (FALSE) estimates.
```

Value

A dataframe with the age/depth of each sample and all the best reconstructed values.

Examples

```
head(crest.simplify(reconstr))
```

crestObj

Create a crestObj object.

Description

Creates a crestObj object with all default parameters.

```
crestObj(
  taxa.name,
  taxaType,
  climate,
  pse = NA,
  dbname = NA,
  continents = NA,
  countries = NA,
  basins = NA,
  sectors = NA,
  realms = NA,
  biomes = NA,
```

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```
ecoregions = NA,
  xmn = NA,
  xmx = NA,
  ymn = NA,
  ymx = NA,
  elev_min = NA,
  elev_max = NA,
  elev_range = NA,
  year_min = 1900,
  year_max = 2021,
  nodate = TRUE,
  type_of_obs = c(1, 2, 3, 8, 9),
  df = NA,
  x = NA,
  x.name = "",
  minGridCells = 20,
  weightedPresences = FALSE,
  bin_width = NA,
  shape = NA,
  npoints = 200,
  geoWeighting = TRUE,
  climateSpaceWeighting = TRUE,
  climateSpaceWeighting.type = "linear",
  climateWithObs = FALSE,
  selectedTaxa = NA,
  distributions = NA,
  presenceThreshold = 0,
  taxWeight = "normalisation",
  uncertainties = c(0.5, 0.95)
)
```

Arguments

xmn, xmx, ymn, ymx

taxa.name	A vector that contains the names of the taxa to study.
taxaType	A numerical index (between 1 and 6) to define the type of palaeoproxy used: 1 for plants, 2 for beetles, 3 for chironomids, 4 for foraminifers, 5 for diatoms and 6 for rodents. The example dataset uses taxaType=0 (pseudo-data). Default is 1.
climate	A vector of the climate variables to extract. See accClimateVariables for the list of accepted values.
pse	A pollen-Species equivalency table. See createPSE for details.
dbname	The name of the data source database.
continents	A vector of the continent names defining the study area.
countries	A vector of the country names defining the study area.
basins	A vector of the ocean names defining the study area.
sectors	A vector of the marine sector names defining the study area.
realms	A vector of the studied botanical realms defining the study area.
biomes	A vector of the studied botanical biomes defining the study area.
ecoregions	A vector of the studied botanical ecoregions defining the study area.

The coordinates defining the study area.

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elev_min, elev_max

Parameters to only selected grid cells with an elevation higher than elev_min or

lower than elev_max (default is 'NA).

elev_range Parameters discard the grid cell with a high elevation range (default is NA).

year_min, year_max

The oldest and youngest occurrences accepted (default is 1900-2021).

nodate A boolean to accept occurrences without a date (can overlap with occurrences

with a date; default TRUE).

type_of_obs The type of observation to use in the study. 1: human observation, 2: observa-

tion, 3: preserved specimen, 4: living specimen, 5: fossil specimen, 6: material sample, 7: machine observation, 8: literature, 9: unknown (Default c(1, 2, 3,

8,9))

df A data frame containing the data to reconstruct (counts, percentages or pres-

ence/absence data).

x The name, age or depth of the rows of df (the samples).

x.name A string describing the x axis (e.g. 'Sample Name', 'Age', 'Depth').

minGridCells The minimum number of unique presence data necessary to estimate a species'

climate response. Default is 20.

weightedPresences

A boolean to indicate whether the presence records should be weighted. Default

is FALSE.

bin_width The width of the bins used to correct for unbalanced climate state. Use values

that split the studied climate gradient in 15-25 classes (e.g. 2°C for temperature

variables). Default is 1.

shape The imposed shape of the species pdfs. We recommend using 'normal' for tem-

perature variables and 'lognormal' for the variables that can only take positive

values, such as precipitation or aridity. Default is 'normal' for all.

npoints The number of points to be used to fit the pdfs. Default 200.

geoWeighting A boolean to indicate if the species should be weighting by the square root

of their extension when estimating a genus/family level taxon-climate relation-

ships.

climateSpaceWeighting

A boolean to indicate if the species pdfs should be corrected for the modern

distribution of the climate space (default TRUE).

climateSpaceWeighting.type

A correction factor for the clame space weighting correction to limit the edge

effects. Either 'linear' (default), 'sqrt' or 'log'.

climateWithObs A boolean to indicate whether all climate values from the calibration dataset be

included (FALSE, default) or only the climate values that corresponds to proxy observations (TRUE). Only useful in if the climate space weighting is activated.

selectedTaxa A data frame assigns which taxa should be used for each variable (1 if the taxon

should be used, 0 otherwise). The colnames should be the climate variables' names and the rownames the taxa names. Default is 1 for all taxa and all vari-

ables.

distributions A dataframe containing the presence records of the studied proxies and their

associated climate values.

crest_ex 27

presenceThreshold

All values above that threshold will be used in the reconstruction (e.g. if set at 1, all percentages below 1 will be set to 0 and the associated presences discarded).

Default is 0.

taxWeight One value among the following: 'originalData', 'presence/absence', 'percent-

ages' or 'normalisation' (default).

uncertainties A (vector of) threshold value(s) indicating the error bars that should be calcu-

lated (default both 50 and 95% ranges).

Value

A crestObj object that is used to store data and information for reconstructing climate

See Also

See vignette('technicalities') for details about the structure of the object. See also https://gbif.github.io/parsers/apidocs/org/gbif/api/vocabulary/BasisOfRecord.html for a detailed explanation of the types of observation.

crest_ex

Example dataset to run the CREST method for the first time.

Description

A dataset containing 20 randomly generated pollen samples for 7 pollen taxa.

Usage

crest_ex

Format

A data frame with 20 rows (samples) and 8 columns (1 column for the age and one for each of the 7 taxa):

Age: Age of each sample

Taxon1: Percentage of Taxon1 in each sample.

Taxon2: Percentage of Taxon2 in each sample.

Taxon3: Percentage of Taxon3 in each sample.

Taxon4: Percentage of Taxon4 in each sample.

Taxon5: Percentage of Taxon5 in each sample.

Taxon6: Percentage of Taxon6 in each sample.

Taxon7: Percentage of Taxon7 in each sample.

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crest_ex_pse

Example dataset to Extract data from the example database.

Description

A database indicating the taxonomy of the example proxies.

Usage

```
crest_ex_pse
```

Format

A data frame with 7 rows (taxa) and 5 columns (taxonomy description):

Level: An integer indicating the taxonomic resolution (1 family, 2 genus, 3 species, 4 or higher

ignore taxon)

Family: The family corresponding to the ProxyName **Genus:** The genus corresponding to the ProxyName **Species:** The species corresponding to the ProxyName

ProxyName: The names of the observed proxies, as reported in the main data file

crest_ex_selection

Example dataset to associate taxa with climate variables.

Description

A data frame indicating the taxa that should be used to reconstruct each climate variable (1s in the matrix) and those who should be excluded (0s).

Usage

```
crest_ex_selection
```

Format

A data frame with 7 rows (taxa) and 2 columns (climate variables):

bio1: The first variable to reconstruct (mean annual temperature)

bio12: The second variable to reconstruct (annual precipitation)

crop 29

crop

Crop the dataset obtained from crest.get_modern_data

Description

 $Crop\ the\ dataset\ obtained\ from\ \verb|crest.get_modern_data|\ according\ to\ an\ object\ of\ the\ class\ Spatial Polygons Data Frames and the class Spatial Polygons Data Frames and the class Spatial Polygons Data Frames Data Frames and the class Spatial Polygons Data Frames Data$

Usage

```
crop(x, shp)
```

Arguments

x A crestObj produced by the crest.get_modern_data function.

shp A shapefile (spatVect) to crop the data. Data points will be kept if their centroid

is within the shape.

Value

An cropped version of the input crestObj.

Examples

```
## Not run:
  data(M1)
 M1 <- terra::unwrap(M1)
  ## We want only the data covering Nigeria
  M2 <- M1[M1$COUNTRY == 'Nigeria', ]
  data(reconstr)
  reconstr.cropped <- crop(reconstr, M2)</pre>
  data1 <- terra::rast(reconstr$modelling$climate_space[, 1:3],</pre>
                                  crs=terra::crs(M1), type='xyz')
  data2 <- terra::rast(reconstr.cropped$modelling$climate_space[, 1:3],</pre>
                                  crs=terra::crs(M1), type='xyz')
  layout(matrix(c(1,2,3,4), byrow=FALSE, ncol=2), width=1, height=c(0.2, 0.8))
  plot_map_eqearth(data1, brks.pos=seq(13,29,2), colour_scale=TRUE,
                   title='Full dataset', zlim=c(13, 29))
  plot_map_eqearth(data2, brks.pos=seq(13,29,2), colour_scale=TRUE,
                   title='Cropped dataset', zlim=c(13, 29))
## End(Not run)
```

dbDownload

Download the gbif4crest_02 dataset from figShare.

Description

Download the gbif4crest_02 dataset from figShare.

30 dbRequest

Usage

```
dbDownload(
  filename = "gbif4crest_03.zip",
  version = 3,
  lite = TRUE,
  res = "5min",
  timeout = 10000
)
```

Arguments

filename The path and name of the file where the database should be saved.

version The version of the gbif4crest dataset desired. Either 2 or 3.

lite Deprecated since v3. A boolean (default TRUE) to indicate if the full database

should be downloaded (including the raw presences from GBIF; lite = FALSE)

or only the curated data (lite = TRUE).

res The spatial resolution of the dataset. Either 15min or 5min. Since v3, 5min is

the only version available.

timeout Maximum duration in seconds of the download. If the file is not fully down-

loaded after 'timeout' seconds, it will be interrupted.

See Also

The full SQLite3 database can be downloaded from https://figshare.com/articles/dataset/GBIF_for_CREST_database/6743207.

Examples

```
## Not run:
   dbDownload() ## This will download the latest version of the database in
   your working directory.
## End(Not run)
```

dbRequest

Connect to the gbif4crest database

Description

Connect to the gbif4crest_02 database by accessing the server on Amazon.

Usage

```
dbRequest(request, dbname = "gbif4crest_02")
```

Arguments

request A SQL request to be executed.

dbname The name of the data source database.

dbSubset 31

Value

The result of the request.

Examples

```
## Not run:
    # Extracting the number of taxa recorded in the database
    dbRequest("SELECT count(*) FROM taxa")

# Extracting all the taxa that have at least one occurrence in South Africa.
southAfricaTaxa <- dbRequest(paste0(
        "SELECT DISTINCT taxa.* ",
        "FROM taxa, distrib_qdgc, geo_qdgc ",
        "WHERE taxa.taxonid=distrib_qdgc.taxonid ",
        "AND distrib_qdgc.latitude=geo_qdgc.latitude ",
        "AND distrib_qdgc.longitude=geo_qdgc.longitude ",
        "AND geo_qdgc.countryname='South Africa'"
))
head(southAfricaTaxa)

## End(Not run)</pre>
```

dbSubset

Create a subset of the global calibration dataset

Description

Create a subset of the global calibration dataset

Usage

```
dbSubset(
  taxaType,
  xmn = NA,
  xmx = NA,
  ymn = NA,
  ymx = NA,
  dbname = "gbif4crest_02",
  out = "gbif4crest_reduced",
  verbose = TRUE
)
```

Arguments

taxaType

A numerical index (between 1 and 6) to define the type of palaeoproxy used: 1 for plants, 2 for beetles, 3 for chironomids, 4 for foraminifers, 5 for diatoms and 6 for rodents. The example dataset uses taxaType=0 (pseudo-data). Default is 1.

```
xmn, xmx, ymn, ymx
```

The coordinates defining the study area.

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dbname The name of the database. Default is 'gbif4crest_02' and data will be ex-

tracted from the online database. The SQLite3 version of the database can also be used here by providing the complete path to a file ending by .sqlite3, e.g.

/path/to/file/gbif4crest_02.sqlite3

out The name or path of the new dataset

verbose A boolean to print non-essential comments on the terminal (default TRUE).

See Also

The full SQLite3 database can be downloaded from https://figshare.com/articles/dataset/GBIF_for_CREST_database/6743207.

Examples

```
## Not run:
  dbSubset(2, xmn=0, xmx=15, ymn=0, ymx=15, out='example.sqlite3')
## End(Not run)
```

eqearth_get_ext

Calculates the extent of the plot in the equal earth projection.

Description

Calculates the extent of the plot in the equal earth projection.

Usage

```
eqearth_get_ext(ext, npoints = 15)
```

Arguments

ext A set of coordinates.

npoints The number of points used to draw the polygon along each dimension.

Value

The set of coordinates ext projected in equal earth.

```
## Not run:
    eqearth_get_ext(c(-15, 50, 30, 70))
## End(Not run)
```

excludeTaxa 33

excludeTaxa	Excludes the list of taxa from the reconstructions.
-------------	---

Description

Excludes the list of taxa from the reconstructions.

Usage

```
excludeTaxa(x, taxa, climate)
```

Arguments

x A crestObj produced by one of the crest, crest.get_modern_data, crest.calibrate, crest.reconstruct or loo functions.

taxa A vector of taxa to exclude.

climate A vector of climate variables to unlink the taxa with.

Value

Return the updated crestObj.

Examples

```
data(reconstr)
print(reconstr$inputs$selectedTaxa)
reconstr <- excludeTaxa(reconstr, 'Taxon3', 'bio1')
## 'Taxon3' is now excluded from the reconstruction of 'bio1'.
print(reconstr$inputs$selectedTaxa)</pre>
```

```
explore_calibration_dataset
```

Extract distributions from the database

Description

This function will extract the distributions of all the studied climate proxy and plot the data on a map.

```
explore_calibration_dataset(
  taxaType,
  save = FALSE,
  filename = "calibrationDataset.pdf",
  col = viridis::viridis(22)[3:22],
  width = 7.48,
  height = 7.48,
  as.png = FALSE,
```

```
png.res = 300,
xmn = NA,
xmx = NA,
ymn = NA,
ymx = NA,
continents = NA,
countries = NA,
realms = NA,
biomes = NA,
ecoregions = NA,
dbname = "gbif4crest_02"
)
```

Arguments

taxaType A numerical index (between 1 and 6) to define the type of palaeoproxy used: 1

for plants, 2 for beetles, 3 for chironomids, 4 for foraminifers, 5 for diatoms and 6 for rodents. The example dataset uses taxaType=0 (pseudo-data). Default is 1.

save A boolean to indicate if the diagram should be saved as a pdf file. Default is

FALSE.

filename An absolute or relative path that indicates where the diagram should be saved.

Also used to specify the name of the file. Default: the file is saved in the working

 $directory\ under\ the\ name\ 'Reconstruction_climate.pdf'.$

col A colour gradient.

width, height The dimensions of the pdf file (default 7.48in ~19cm).

as.png A boolean to indicate if the output should be saved as a png. Default is FALSE

and the figure is saved as a pdf file.

png.res The resolution of the png file (default 300 pixels per inch).

xmn, xmx, ymn, ymx

The coordinates defining the study area.

continents A vector of the continent names defining the study area.

A vector of the country names defining the study area.

realms A vector of the studied botanical realms defining the study area.

biomes A vector of the studied botanical biomes defining the study area.

ecoregions A vector of the studied botanical ecoregions defining the study area.

dbname The name of the database. Default is 'gbif4crest_02'.

Value

The distribution data

export 35

```
## End(Not run)
```

export

Export the results

Description

Export the results generated by the reconstruction

Usage

```
export(
    x,
    dataname = x$misc$site_info$site_name,
    climate = x$parameters$climate,
    loc = getwd(),
    as.csv = FALSE,
    fullUncertainties = FALSE,
    loo = FALSE,
    weights = FALSE,
    pdfs = FALSE
```

Arguments

x	The name, age or depth of the rows of df (the samples).	
dataname	The name of the site (default: x\$misc\$site_info\$site_name)	
climate	The climate data to export. Data for all climate variables are saved by default.	
loc	The path where to export the data (default: working directory)	
as.csv	Boolean to indicate if the data should be exported as csv (TRUE) or xlsx (FALSE, default)	
fullUncertainties		
	A boolean to export the full climate uncertainty distribution (default FALSE)	
100	A boolean to export the leave-one-out data if they exist (default FALSE)	
weights	A boolean to export the weights derived from the percentages (default FALSE)	
pdfs	A boolean to export the taxa's pdfs (default FALSE)	

Value

No return value, function called to export the results.

36 export_pdfs

Examples

export_pdfs

Export the pdfs fitted for the different taxa.

Description

Export the pdfs fitted for the different taxa.

Usage

```
export_pdfs(
   x,
   dataname = x$misc$site_info$site_name,
   climate = x$parameters$climate,
   taxa = x$inputs$taxa.name,
   loc = getwd(),
   as.csv = FALSE
)
```

Arguments

X	The name, age or depth of the rows of df (the samples).
dataname	The name of the site (default: x\$misc\$site_info\$site_name)
climate	A vector of the climate variables to extract. See accClimateVariables for the list of accepted values.
taxa	The names of the taxa of interest. All the pdfs are saved by default.
loc	The path where to export the data (default: working directory)
as.csv	Boolean to indicate if the data should be exported as csv (TRUE) or xlsx (FALSE, default)

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Value

No return value, function called to export the PDFs as files.

Examples

```
## Not run:
  data(crest_ex)
  data(crest_ex_pse)
  data(crest_ex_selection)
  reconstr <- crest(</pre>
    df = crest_ex, pse = crest_ex_pse, taxaType = 0,
    site_info = c(7.5, 7.5),
    climate = c("bio1", "bio12"), bin_width = c(2, 20), shape = c("normal", "lognormal"),
    selectedTaxa = crest_ex_selection, dbname = "crest_example",
    leave_one_out = TRUE
  #> Replace 'tempdir()' by the location where the sample should be saved (e.g. 'getwd()')
  export_pdfs(reconstr,
               dataname='crest_example',
               loc=tempdir()
  )
## End(Not run)
```

find.original.name

Returns the name of the function argument in the global environment

Description

Returns the name of the function argument in the global environment

Usage

```
find.original.name(x)
```

Arguments

Х

The function argument

Value

The name of the function argument in the global environment.

38 fit_pdfsp

fit_pdfsp

Fit the species pdfs.

Description

Fit the species pdfs.

Usage

```
fit_pdfsp(
  climate,
  ccs,
  bin_width,
  shape,
   xrange,
  use_ccs = TRUE,
  climateSpaceWeighting.type = "linear"
)
```

Arguments

climate A vector of climatic values where the species is present.

ccs A ccs object returned by calib_clim_space.

bin_width The width of the bins used to correct for unbalanced climate state. Use values

that split the studied climate gradient in 15-25 classes (e.g. 2°C for temperature

variables). Default is 1.

shape The imposed shape of the species pdfs. We recommend using 'normal' for tem-

perature variables and 'lognormal' for the variables that can only take positive

values, such as precipitation or aridity. Default is 'normal' for all.

xrange The climate gradient upon which the pdf with be defined.

use_ccs Boolean to indicate if the pdfsp should be corrected by the distribution of the

modern climate space

climateSpaceWeighting.type

A correction factor for the clame space weighting correction to limit the edge

effects. Either 'linear' (default), 'sqrt' or 'log'.

Value

The pdf of the species.

```
# Creating one randomised species
climate_species <- round(stats::rnorm(50, 15, 2), 1)
# Creating one randomised climate space
climate_space <- base::sample(0:300 / 10, 4000, replace = TRUE)

ccs <- calib_clim_space(climate_space, 2)
xrange <- fit_xrange(ccs, "normal", 2)</pre>
```

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```
pdfsp <- fit_pdfsp(climate_species, ccs, 2, "normal", xrange)
plot(xrange, pdfsp, type = "1")

# Testing that the area under the curve is equal to 1.
all.equal(sum(pdfsp * (xrange[2] - xrange[1])), 1)</pre>
```

fit_xrange

Define the climate gradient to fit the pdfs.

Description

Define the climate gradient to fit the pdfs.

Usage

```
fit_xrange(ccs, shape, bin_width, npoints = 500)
```

Arguments

ccs	A ccs object returned by calib_clim_space.
shape	The imposed shape of the species pdfs. We recommend using 'normal' for temperature variables and 'lognormal' for the variables that can only take positive values, such as precipitation or aridity. Default is 'normal' for all.
bin_width	The width of the bins used to correct for unbalanced climate state. Use values that split the studied climate gradient in 15-25 classes (e.g. 2°C for temperature variables). Default is 1.
npoints	The number of points to be used to fit the pdfs. Default 200.

Value

A regularly spaced climate gradient with npoints points.

```
# Creating one randomised climate space
climate_space <- sample(0:300 / 10, 4000, replace = TRUE)
ccs <- calib_clim_space(climate_space, 2)
xrange <- fit_xrange(ccs, "normal", 2)
head(xrange)</pre>
```

40 getClimateSpace

f_locid

Calculates a unique identifier from coordinates.

Description

Calculates a unique identifier from coordinates.

Usage

```
f_locid(long, lat, resol)
```

Arguments

long The longitude of the grid cell of interest

The latitude of the grid cell of interest

resol The spatial resolution of the database to target (1/12 for gbif4crest_03)

Value

A unique integer describing the cell.

Examples

```
f_locid(0, 0, 1/12)
```

getClimateSpace

Extract the distribution of the studied climate gradient(s) across the study area.

Description

Extract the distribution of the studied climate gradient(s) across the study area.

Usage

```
getClimateSpace(
  climate,
  xmn = NA,
  xmx = NA,
  ymn = NA,
  ymx = NA,
  continents = NA,
  countries = NA,
  basins = NA,
  sectors = NA,
  realms = NA,
  biomes = NA,
  ecoregions = NA,
```

getClimateSpace 41

```
elev_min = NA,
  elev_max = NA,
  elev_range = NA,
  dbname = "gbif4crest_02"
)
```

Arguments

climate A vector of the climate variables to extract. See accClimateVariables for the

list of accepted values.

xmn, xmx, ymn, ymx

The coordinates defining the study area.

continents A vector of the continent names defining the study area.

Countries A vector of the country names defining the study area.

A vector of the ocean names defining the study area.

sectors $\mbox{\ \ A vector of the marine sector names defining the study area.}$

realms A vector of the studied botanical realms defining the study area.

biomes A vector of the studied botanical biomes defining the study area.

ecoregions A vector of the studied botanical ecoregions defining the study area.

elev_min, elev_max

Parameters to only selected grid cells with an elevation higher than elev_min or

lower than elev_max (default is 'NA).

elev_range Parameters discard the grid cell with a high elevation range (default is NA).

dbname The name of the data source database.

Value

A matrix of occurrence records with the associated climate.

See Also

accClimateVariables for a list of accepted climate variable names, accCountryNames for a list of accepted continent and country names, accBasinNames for a list of accepted basin and sector names, accRealmNames for a list of accepted realm, biome and ecoregion names.

```
## Not run:
    climate <- getClimateSpace("bio1", -90, 90, -90, 90,
        continents = "Europe",
        countries = c("Germany", "Netherlands", "Sweden"),
        realms = "Palaearctic"
)
    head(climate)
    terra::plot(terra::rast(climate, type='xyz'), asp=1)
## End(Not run)</pre>
```

42 getDistribTaxa

getDistribTaxa

Extract taxonID(s) corresponding to the taxonomic description

Description

Extract all possible TaxonIDs corresponding to the provided taxonomical description, which can be at the family, the genus or the species levels.

Usage

```
getDistribTaxa(
  taxIDs,
  climate = NA,
  xmn = NA,
  xmx = NA,
  ymn = NA,
  ymx = NA,
  continents = NA,
  countries = NA,
  basins = NA,
  sectors = NA,
  realms = NA,
  biomes = NA,
  ecoregions = NA,
  elev_min = NA,
  elev_max = NA,
  elev_range = NA,
  year_min = 1900,
  year_max = 2021,
  nodate = TRUE,
  type_of_obs = c(1, 2, 7, 8),
  dbname = "gbif4crest_02"
)
```

Arguments

taxIDs	A vector of accepted	Taxa IDs (as returned	by getTaxonID).
--------	----------------------	-----------------------	-----------------

climate A vector of the climate variables to extract. See accClimateVariables for the

list of accepted values.

xmn, xmx, ymn, ymx

The coordinates defining the study area.

continents A vector of the continent names defining the study area.

A vector of the country names defining the study area.

A vector of the ocean names defining the study area.

sectors A vector of the marine sector names defining the study area.

realms A vector of the studied botanical realms defining the study area.

biomes A vector of the studied botanical biomes defining the study area.

ecoregions A vector of the studied botanical ecoregions defining the study area.

getResol 43

elev_min, elev_max

Parameters to only selected grid cells with an elevation higher than elev_min or

lower than elev_max (default is 'NA).

elev_range Parameters discard the grid cell with a high elevation range (default is NA).

year_min, year_max

The oldest and youngest occurrences accepted (default is 1900-2021).

nodate A boolean to accept occurrences without a date (can overlap with occurrences

with a date; default TRUE).

type_of_obs The type of observation to use in the study. 1: human observation, 2: observa-

tion, 3: preserved specimen, 4: living specimen, 5: fossil specimen, 6: material sample, 7: machine observation, 8: literature, 9: unknown (Default c(1, 2, 3,

8, 9))

dbname The name of the data source database.

Value

A matrix of occurrence records with the associated climate.

See Also

getTaxonID for taxIDs, accClimateVariables for a list of accepted climate variable names, accCountryNames for a list of accepted continent and country names, accRealmNames for a list of accepted realm, biome and ecoregion names.

Examples

```
## Not run:
    taxIDs <- getTaxonID("Zamiaceae", "Ceratozamia")
    distrib <- getDistribTaxa(taxIDs, "bio1", -90, 90, -90, 90,
        continents = "Europe",
        countries = c("Germany", "Netherlands", "Sweden"),
        realms = "Palaearctic"
    )
    distrib
## End(Not run)</pre>
```

getResol

Calculates a unique identifier from coordinates.

Description

Calculates a unique identifier from coordinates.

Usage

```
getResol(crest)
```

Arguments

crest

A crestObj object.

Value

A unique integer describing the cell.

Examples

```
getResol(reconstr)
```

getSpeciesdiversity

Calculates how many species compose the response of each taxon.

Description

Calculates how many species compose the response of each taxon.

Usage

```
getSpeciesdiversity(x)
```

Arguments

Χ

A crest0bj generated by the crest.reconstruct, loo or crest functions.

Value

Return the number of composing species of each taxon.

Examples

```
data(reconstr)
getSpeciesdiversity(reconstr)
```

getTaxaTypeFromTaxID

Returns the taxa type corresponding to the taxID.

Description

Returns the taxa type corresponding to the taxID.

Usage

```
getTaxaTypeFromTaxID(taxID)
```

Arguments

taxID

An integer between 0 and 6

Value

Returns the taxa type ID corresponding to the taxon ID.

getTaxonID 45

get]	 T D	۱

Extract taxonID(s) corresponding to the taxonomic description

Description

Extract all possible TaxonIDs corresponding to the provided taxonomical description, which can be at the family, the genus or the species levels.

Usage

```
getTaxonID(
  family = "",
  genus = "",
  species = "",
  taxaType = 1,
  dbname = "gbif4crest_02"
)
```

Arguments

family	The name of the family.
genus	The name of the genus.
species	The name of the species.
taxaType	A numerical index (between 1 and 6) to define the type of palaeoproxy used: 1 for plants, 2 for beetles, 3 for chironomids, 4 for foraminifers, 5 for diatoms and 6 for rodents. The example dataset uses taxaType=0 (pseudo-data). Default is 1.
dbname	The name of the data source database.

Value

A vector of unique taxonIDs.

```
## Not run:
    getTaxonID("Zamiaceae")
    getTaxonID("Zamiaceae", "Ceratozamia")

## \code{taxaType = 2} searches for beetles and not plants, so the next line returns nothing.
    getTaxonID("Zamiaceae", "Ceratozamia", taxaType = 2)

## End(Not run)
```

46 getTaxonomy

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Extract taxonID(s) corresponding to the taxonomic description

Description

Extract all possible TaxonIDs corresponding to the provided taxonomical description, which can be at the family, the genus or the species levels.

Usage

```
getTaxonomy(
  family = "",
  genus = "",
  species = "",
  taxaType = 1,
  depth.out = 8,
  dbname = "gbif4crest_02",
  crest = NA
)
```

Arguments

family The name of the family.
genus The name of the genus.
species The name of the species.

taxaType A numerical index (between 1 and 5) to define the type of palaeoproxy used: 1

for plants, 2 for beetles, 3 for chironomids, 4 for foraminifers, 5 for diatoms and

6 for rodents.

depth.out The taxonomic resolution of the output table. 1 for Kingdom, 2 for phylum, 3

for class_name, 4 for order_name, 5 for family, 6 for genus, 7 for species and 8

to also include the taxonID.

dbname The name of the data source database.

crest A crestObj to be used to refine the selection to a specific study area. Set to NA

by default (global search).

Value

A vector of unique taxonIDs.

```
## Not run:
    getTaxonomy("Zamiaceae ")
    getTaxonomy(genus="Ceratozamia", depth.out=6)
## \code{taxaType = 2} searches for beetles and not plants, so the next line returns nothing.
    getTaxonomy("Zamiaceae", "Ceratozamia", taxaType = 2)
## End(Not run)
```

get_taxa_type 47

get_taxa_type

Returns the taxa type corresponding to the index.

Description

Returns the taxa type corresponding to the index.

Usage

```
get_taxa_type(taxaType)
```

Arguments

taxaType

An integer between 0 and 6

Value

Returns the taxa type corresponding to the index.

 $identify {\tt Database}$

Identify the calibration database used.

Description

Identify the calibration database used.

Usage

```
identifyDatabase(dbname)
```

Arguments

dbname

A functional crestObj or a database name.

Value

A string uniquely characterising the database used. Possible values are 'privateDB', 'exampleDB', 'gbif4crest_02', and 'gbif4crest_03'.

```
identifyDatabase(reconstr)
```

48 is.crestObj

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Includes the list of taxa into the reconstructions.

Description

Includes the list of taxa into the reconstructions.

Usage

```
includeTaxa(x, taxa, climate)
```

Arguments

x A crestObj produced by one of the crest, crest.get_modern_data, crest.calibrate,

crest.reconstruct or loo functions.

taxa A vector of taxa to include.

climate A vector of climate variables to link the taxa with.

Value

Return the updated crestObj.

Examples

```
data(reconstr)
print(reconstr$inputs$selectedTaxa)
reconstr <- includeTaxa(reconstr, reconstr$inputs$taxa.name, 'bio12')
## All the taxa are not selected for 'bio12', except for 'Taxon7' for which
## data are unavailable.
print(reconstr$inputs$selectedTaxa)</pre>
```

is.crestObj

Test if x *is a crestObj.*

Description

```
Test if x is a crestObj.
```

Usage

```
is.crestObj(x)
```

Arguments

Χ

The object to be tested

Value

```
TRUE (x is a crestObj) or FALSE (not a crestObj).
```

isColourStr 49

isColourStr

Test if R can interpret a string as a colour

Description

Test if R can interpret a string as a colour

Usage

```
isColourStr(col)
```

Arguments

col

The string to be tested.

Value

A boolean value, TRUE if col is a valid colour, FALSE otherwise

Examples

```
isColourStr('black')
isColourStr('blakc')
```

100

Performs the leave-one-out analysis

Description

Repeat the reconstructions by removing one taxon at a time.

Usage

```
loo(x, climate = x$parameters$climate, verbose = TRUE)
```

Arguments

x a crest0bj produced by the crest.reconstruct or crest functions.

climate A vector of the climate variables to extract. See accClimateVariables for the

list of accepted values.

verbose A boolean to print non-essential comments on the terminal (default TRUE).

Value

A crest0bj object containing the reconstructions and all the associated data.

50 makeTransparent

Examples

```
## Not run:
    data(crest_ex)
    data(crest_ex_pse)
    data(crest_ex_selection)
    reconstr <- crest(
        df = crest_ex, pse = crest_ex_pse, taxaType = 0,
            climate = c("bio1", "bio12"), bin_width = c(2, 20),
        shape = c("normal", "lognormal"),
        selectedTaxa = crest_ex_selection, dbname = "crest_example"
    )
    reconstr <- loo(reconstr)

## End(Not run)
## example using pre-saved reconstruction obtained with the previous command.
data(reconstr)
lapply(reconstr$reconstructions$bio12$loo, head)
plot_loo(reconstr)</pre>
```

M1

A shapefile of the world's country borders. Use 'M1 \leftarrow terra::unwrap(M1)' to render the object usable in R.

Description

A shapefile of the world's country borders. Use 'M1 \leftarrow terra::unwrap(M1)' to render the object usable in R.

Usage

M1

Format

An object of class PackedSpatVector of length 1.

makeTransparent

Wrapper function of to add transparency to a colour.

Description

Add transparency to the selected colours.

Usage

```
makeTransparent(colour, alpha)
```

meanPositiveValues 51

Arguments

colour A R colour

alpha A value between 0 and 1 that defines the transparency 0 for full transparency

and 1 for no transparency

Value

Return a colour with the provided level of transparency.

Examples

```
makeTransparent('black',0.5)
makeTransparent('black',1:10/10)
makeTransparent(rainbow(10), 1:10/10)
```

meanPositiveValues

Calculate the mean of all strictly positive values.

Description

Calculate the mean of all strictly positive values.

Usage

```
meanPositiveValues(x)
```

Arguments

Х

A vector of values.

Value

The average of all the positive values. Returns NaN is no strictly positive values are found.

```
meanPositiveValues(-10:10)
```

52 pdf_ranges

normalise

Normalises the percentages

Description

Normalises the percentages

Usage

```
normalise(df, col2convert = 2:ncol(df))
```

Arguments

df

The dataframe containing the data to convert.

col2convert

A vector of the columns to convert. Default is all the columns but the first, which contains an age, a depth or a sampleID.

Value

A vector of unique taxonIDs.

Examples

```
df <- data.frame(matrix(1:25, ncol = 5))
colnames(df) <- paste(rep("col", 5), 1:5, sep = "")
normalise(df)
normalise(df, col2convert = 3:5)</pre>
```

pdf_ranges

Calculate the climate tolerance of the taxa from their pdfs.

Description

Calculate the climate tolerance of the taxa from their pdfs.

Usage

```
pdf_ranges(
    x,
    climate = x$parameters$climate,
    taxanames = x$input$taxa.name,
    uncertainties = x$parameters$uncertainties,
    orderby = NA
)
```

plot.crestObj 53

Arguments

A crestObj generated by either the crest.calibrate, crest.reconstruct or crest functions.

Climate Climate variables to be used to generate the plot. By default all the variables are included.

taxanames A list of taxa to use for the plot (default is all the recorded taxa).

uncertainties A (vector of) threshold value(s) indicating the error bars that should be calculated (default are the values stored in x).

orderby A string ('name', or one of the climate variables) to sort the output table

Value

The set of coordinates ext projected in equal earth.

Examples

```
data(reconstr)
pdf_ranges(reconstr, climate='bio1')
pdf_ranges(reconstr, climate='bio12', orderby='bio1', uncertainties=c(0.2, 0.6, 0.95))
```

plot.crestObj

Plot the reconstructions.

Description

Plot the reconstructions and their uncertainties if they exist.

Usage

```
## S3 method for class 'crestObj'
plot(
  Х,
  climate = x$parameters$climate[1],
  uncertainties = x$parameters$uncertainties,
  optima = TRUE,
  add_modern = FALSE,
  simplify = FALSE,
  as.anomaly = FALSE,
  anomaly.base = x$misc$site_info$climate[climate],
  xlim = NA,
  ylim = NA,
  pt.cex = 0.8,
  pt.lwd = 0.8,
  pt.col = ifelse(simplify, "black", "white"),
  col.hiatus = "white",
  save = FALSE,
  width = 5.51,
  height = 5.51,
  as.png = FALSE,
```

54 plot.crestObj

```
png.res = 300,
filename = "Reconstruction.pdf",
col = viridis::viridis(125)[26:125],
...
)
```

Arguments

x	A crestObj produced by the crest, crest.reconstruct or loo functions.
climate	The climate variables to plot (default is all the reconstructed variables from x)
uncertainties	A (vector of) threshold value(s) indicating the error bars that should be calculated (default are the values stored in \mathbf{x}).
optima	A boolean to indicate whether to plot the optimum (TRUE) or the mean (FALSE) estimates.
add_modern	Adds the modern climate values to the plot.
simplify	A boolean to indicate if the full distribution of uncertainties should be plotted (FALSE, default) or if they should be simplified to the uncertainty range(s).
as.anomaly	A boolean to indicate if the reconstructions should be plotted as absolute values (Default, FALSE) or anomalies '(TRUE).
anomaly.base	The anomaly value. Should be a vector with the same length as climate. Default values are the climate values correpsonding to the location of the record (site_info in crest.get_modern_data).
xlim	the x limits $(x1, x2)$ of the plot. Note that $x1 > x2$ is allowed and leads to a 'reversed axis'.
	The default value, NULL, indicates that the range of the finite values to be plotted should be used.
ylim	the y limits of the plot.
pt.cex	The size of the points (default 0.8).
pt.lwd	The thickness of the lines (default 0.8).
pt.col	The colour of the points and lines.
col.hiatus	A colour for the hiatus(es) of the record (default white)
save	A boolean to indicate if the diagram should be saved as a pdf file. Default is FALSE.
width, height	The dimensions of the pdf file (default 5.51in ~14cm).
as.png	A boolean to indicate if the output should be saved as a png. Default is FALSE and the figure is saved as a pdf file.
png.res	The resolution of the png file (default 300 pixels per inch).
filename	An absolute or relative path that indicates where the diagram should be saved. Also used to specify the name of the file. Default: the file is saved in the working directory under the name 'Reconstruction_climate.pdf'.
col	A colour gradient.
	other graphical parameters (see par and section 'Details' below).

Value

No return value, this function is used to plot.

plot_climateSpace 55

Examples

```
## Not run:
    data(crest_ex)
    data(crest_ex_pse)
    data(crest_ex_selection)
    reconstr <- crest(
        df = crest_ex, pse = crest_ex_pse, taxaType = 0,
            climate = c("bio1", "bio12"), bin_width = c(2, 20),
            shape = c("normal", "lognormal"),
            selectedTaxa = crest_ex_selection, dbname = "crest_example"
    )
    reconstr <- loo(reconstr)

## End(Not run)
## example using pre-saved reconstruction obtained with the previous command.
data(reconstr)
plot(reconstr, climate='bio1', simplify = TRUE, as.anomaly=TRUE)</pre>
```

plot_climateSpace

Plot the studied climate space.

Description

Plot the studied climate space.

Usage

```
plot_climateSpace(
    x,
    climate = x$parameters$climate,
    bin_width = x$parameters$bin_width[x$parameters$climate, ],
    save = FALSE,
    filename = "Climate_space.pdf",
    as.png = FALSE,
    png.res = 300,
    width = 7.48,
    height = min(9, 3.5 * length(climate)),
    y0 = 0.4,
    add_modern = FALSE,
    resol = getResol(x)
)
```

Arguments

x A crestObj generated by either the crest.calibrate, crest.reconstruct or crest functions.

climate

Climate variables to be used to generate the plot. By default all the variables are included.

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bin_width	The width of the bins used to correct for unbalanced climate state. Use values that split the studied climate gradient in 15-25 classes (e.g. 2°C for temperature variables). Default is 1.
save	A boolean to indicate if the diagram should be saved as a pdf file. Default is FALSE.
filename	An absolute or relative path that indicates where the diagram should be saved. Also used to specify the name of the file. Default: the file is saved in the working directory under the name 'Climate_space.pdf'.
as.png	A boolean to indicate if the output should be saved as a png. Default is FALSE and the figure is saved as a pdf file.
png.res	The resolution of the png file (default 300 pixels per inch).
width	The width of the output file in inches (default 7.48in ~ 19cm).
height	The height of the output file in inches (default 3in ~ 7.6cm per variables).
y0	The space to allocate to each title (default 0.4in ~ 1 cm.
add_modern	A boolean to add the location and the modern climate values to the plot (default FALSE).
resol	For advanced users only: if higher resolution data are used to estimate the pdfs, use this parameter to define the resolution of the maps maps on the figures. (default is 0.25 degrees to match with the default database).

Value

No return value, this function is used to plot.

Examples

```
## Not run:
    data(crest_ex_pse)
    data(crest_ex_selection)
    reconstr <- crest.get_modern_data(
        pse = crest_ex_pse, taxaType = 0,
        climate = c("bio1", "bio12"),
        selectedTaxa = crest_ex_selection, dbname = "crest_example"
)
    reconstr <- crest.calibrate(reconstr,
        geoWeighting = TRUE, climateSpaceWeighting = TRUE,
        bin_width = c(2, 20), shape = c("normal", "lognormal")
)
    plot_climateSpace(reconstr)
## End(Not run)</pre>
```

plot_combinedPDFs

Plot representing how the pdfs combine to produce the reconstruction.

Description

Plot representing how the pdfs combine to produce the reconstruction.

plot_combinedPDFs 57

Usage

```
plot_combinedPDFs(
  х,
  ages = range(x$inputs$x),
  samples = which(x$inputs$x >= min(ages) & x$inputs$x <= max(ages)),</pre>
  climate = x$parameters$climate[1],
  optima = TRUE,
  xlim = NA,
  only.present = FALSE,
  only.selected = FALSE,
  col = crestr::colour_theme(1),
  save = FALSE,
  filename = "samplePDFs.pdf",
  as.png = FALSE,
  png.res = 300,
  width = 7.48,
  height = 5
)
```

Arguments

save

x	A crest0bj generated by the crest.reconstruct or crest functions.
ages	An age range to subset the samples to plot. By default, all the samples will be selected.
samples	The list of sample indexes for which the plot should be plotted. All samples will be plotted by default.
climate	The climate variable to use to plot the variable. Default is first variable (x\$parameters\$climate\[1\
optima	A boolean to indicate whether to plot the optimum (TRUE) or the mean (FALSE) estimates.
xlim	The climate range to plot the pdfs on. Default is the full range used to fit the pdfs (x\$modelling\$xrange).
only.present	A boolean to only add the names of the taxa recorded in the sample (default FALSE).
only.selected	A boolean to only add the names of the selected taxa (default FALSE).
col	A range of colour values to colour the pdfs. Colours will be recycled to match the number of taxa.

A boolean to indicate if the diagram should be saved as a pdf file. Default is

FALSE.

filename An absolute or relative path that indicates where the diagram should be saved.

Also used to specify the name of the file. Default: the file is saved in the working

directory under the name 'samplePDFs.pdf'.

A boolean to indicate if the output should be saved as a png. Default is FALSE as.png

and the figure is saved as a pdf file.

The resolution of the png file (default 300 pixels per inch). png.res

The width of the output file in inches (default 7.48in ~ 19cm). width

height The height of the output file in inches (default 5in ~ 12.7cm). 58 plot_diagram

Value

No return value, this function is used to plot.

Examples

```
## Not run:
    data(crest_ex)
    data(crest_ex_pse)
    data(crest_ex_selection)
    reconstr <- crest(
        df = crest_ex, pse = crest_ex_pse, taxaType = 0,
            climate = c("bio1", "bio12"), bin_width = c(2, 20),
            shape = c("normal", "lognormal"),
            selectedTaxa = crest_ex_selection, dbname = "crest_example",
            leave_one_out = FALSE
        )

## End(Not run)
## example using pre-saved reconstruction obtained with the previous command.
data(reconstr)
plot_combinedPDFs(reconstr, ages=c(6,9), climate='bio12')</pre>
```

plot_diagram

Plot stratigraphic data as polygons or barplots.

Description

This function plots stratigraphic data either as polygons or bars.

Usage

```
plot_diagram(
  х,
  bars = FALSE,
  col = "black",
  amplif = 5,
  save = FALSE,
  filename = "Diagram.pdf",
  width = 3.54,
  height = 9,
  as.png = FALSE,
  png.res = 300,
  yax_incr = 5,
  bar_width = diff(range(x$inputs$x))/length(x$inputs$x),
  xlim = NA,
  tickAtSample = TRUE,
  col_pos = "black",
  col_neg = "grey80",
  title = NA,
  src = NA
)
```

plot_diagram 59

Arguments

bars A boolean that indicates if the data should be plotted as polygons (default: bars=FALSE) or vertical bars (bars=TRUE).
Colours to be used for the polygons. If the number of colours does not match the number of taxa, colors will be recycled.
amplif A factor the show exaggeration on the diagram. Only for polygon plot. Default 5.
A boolean to indicate if the diagram should be saved as a pdf file. Default is FALSE.
An absolute or relative path that indicates where the diagram should be saved. Also used to specify the name of the file. Default: the file is saved in the working directory under the name 'Diagram.pdf'.
width The width of the output file in inches (default 3.54in ~ 9cm).
height The height of the output file in inches (default 9in ~ 23cm).
as.png A boolean to indicate if the output should be saved as a png. Default is FALSE and the figure is saved as a pdf file.
png.res The resolution of the png file (default 300 pixels per inch).
yax_incr Graphical parameters describing the increment size on the y-axis (default 5).
bar_width Width of the bars of the barplot (default 1/50th of the x range).
xlim The range covered by the x-axis. Can be adjusted to get round numbers on the x-axis. If smaller than the range covered by the data, the data will be truncated (default: range of the data).
tickAtSample Boolean that indicates whether a tick mark should be added on the x-axis at the location of each sample (default TRUE).
col_pos Graphical parameter for the barplot. Colour of all the positive values (default black).
col_neg Graphical parameter for the barplot. Colour of all the negative values (default light grey).
title Name to be added on top of the plot (default NA).
src A graphical parameter used by the plot_loo() function.

Value

No return value, this function is used to plot.

plot_loo

plot_loo

Plot the results of the leave-one-out analysis.

Description

Plot the results of the leave-one-out analysis.

Usage

```
plot_loo(
  х,
  optima = TRUE,
  climate = x$parameters$climate[unlist(lapply(x$reconstructions, function(y)
    return("loo" %in% names(y))))],
  taxanames = x$inputs$taxa.name,
  save = FALSE,
  filename = "Diagram_loo.pdf",
  as.png = FALSE,
  png.res = 300,
  width = 3.54,
  height = 9,
  yax_incr = NA,
  bar_width = diff(range(x$inputs$x))/length(x$inputs$x),
  xlim = NA,
  tickAtSample = FALSE,
  sort = NA,
  filter = 0,
  col_pos = "black",
  col_neg = "grey80",
  title = NA
)
```

Arguments

X	A data frame of the data to plot (first column with age or depth) and the taxa in the following columns. x can also be a crestObj.
optima	A boolean to indicate whether to plot the optimum (TRUE) or the mean (FALSE) estimates.
climate	Climate variables to be used to generate the plot. By default all the variables are included.
taxanames	A list of taxa to use for the plot (default is all the recorded taxa).
save	A boolean to indicate if the diagram should be saved as a pdf file. Default is FALSE.
filename	An absolute or relative path that indicates where the diagram should be saved. Also used to specify the name of the file. Default: the file is saved in the working directory under the name 'Diagram_loo_climate.pdf'.
as.png	A boolean to indicate if the output should be saved as a png. Default is FALSE and the figure is saved as a pdf file.
png.res	The resolution of the png file (default 300 pixels per inch).

plot_loo 61

width	The width of the output file in inches (default 3.54in ~ 9cm).
height	The height of the output file in inches (default 9in ~ 23cm).
yax_incr	Graphical parameters describing the increment size on the y-axis (default 5).
bar_width	Width of the bars of the barplot (default 1).
xlim	The range covered by the x-axis. Can be adjusted to get round numbers on the x-axis. If smaller than the range covered by the data, the data will be truncated (default: range of the data).
tickAtSample	Boolean that indicates whether a tick mark should be added on the x-axis at the location of each sample (default TRUE).
sort	A string to sort the order of the taxa from the highest to lowest anomalies (sort='incr') or from the lowest to highest (sort='decr'). Use the default value NA to keep the taxa unsorted.
filter	A threshold value that determines the mean absolute anomaly value required for the taxon to be plotted (default 0 means that all taxa are plotted)
col_pos	Graphical parameter for the barplot. Colour of all the positive values (default black).
col_neg	Graphical parameter for the barplot. Colour of all the negative values (default grey80).
title	Name to be added on top of the plot (default NA).

Value

When used with a crestObj, it returns the average leave-one-out values for each selected taxa

62 plot_map_eqearth

Description

Plots raster data in equal earth projection.

Usage

```
plot_map_eqearth(
  dat,
  ext = as.vector(terra::ext(dat)),
  zlim = range(terra::values(dat), na.rm = TRUE),
  col = viridis::viridis(20),
  brks.pos = c(0, 1),
  brks.lab = brks.pos,
  npoints = 15,
  nlines = 9,
  title = "",
  colour_scale = TRUE,
  top_layer = NA,
  top_layer.col = "ghostwhite",
  site_xy = NA,
  dim = NA
)
```

Arguments

dat	The raster data to plot.
ext	The extent to use to plot the data. (default is extent of dat)
zlim	The range of the values to plot. (default is estimated from dat)
col	The color gradient to use. (default is viridis)
brks.pos	The position where to draw tick marks on the legend
brks.lab	The labels to add where the tickmarks are draw (default is tickmarks position)
npoints	The number of points used to draw the polygons and lines along each dimension. (default is 15 for a smooth result)
nlines	The number of coordinate lines to add in the background (default is 9)
title	A description title (default is empty).
colour_scale	A boolean to add the colour scale to the plot (default TRUE).
top_layer	A raster to overlay on top of the map (e.g. a distribution).
top_layer.col	A colour for plotting top_layer (default 'ghostwhite').
site_xy	Coordinates of a location to add to the plot.
dim	The dimension of the plotting window in inches (default dev.size()).

Value

The set of coordinates ext projected in equal earth.

plot_scatterPDFs 63

plot_scatterPDFs

Plot the pdf optima and uncertainty ranges in a climate biplot

Description

Plot the pdf optima and uncertainty ranges in a climate biplot

Usage

```
plot_scatterPDFs(
    x,
    climate = x$parameters$climate[1:2],
    taxanames = x$input$taxa.name,
    uncertainties = x$parameters$uncertainties,
    xlim = range(x$modelling$climate_space[, climate[1]]),
    ylim = range(x$modelling$climate_space[, climate[2]]),
    add_modern = FALSE,
    save = FALSE,
    filename = "scatterPDFs.pdf",
    width = 5.51,
    height = 5.51,
    as.png = FALSE,
    png.res = 300
)
```

Arguments

х	A crestObj generated by either the crest.calibrate, crest.reconstruct or crest functions.
climate	Names of the two climate variables to be used to generate the plot. By default plot. By default the first two variables are included.
taxanames	A list of taxa to use for the plot (default is all the recorded taxa).
uncertainties	A (vector of) threshold value(s) indicating the error bars that should be calculated (default are the values stored in \mathbf{x}).

xlim ylim The climate range to plot the data. Default is the full range of the observed climate space.

ylim the y limits of the plot.

add_modern A boolean to add the location and the modern climate values to the plot (default

FALSE).

save A boolean to indicate if the diagram should be saved as a pdf file. Default is

FALSE.

filename An absolute or relative path that indicates where the diagram should be saved.

Also used to specify the name of the file. Default: the file is saved in the working

directory under the name 'violinPDFs.pdf'.

width The width of the output file in inches (default 7.48in ~ 19cm).

height The height of the output file in inches (default 3in ~ 7.6cm per variables).

as.png A boolean to indicate if the output should be saved as a png. Default is FALSE

and the figure is saved as a pdf file.

png.res The resolution of the png file (default 300 pixels per inch).

Value

A table with the climate tolerances of all the taxa

Examples

```
## Not run:
  data(crest_ex_pse)
  data(crest_ex_selection)
  reconstr <- crest.get_modern_data(</pre>
    pse = crest_ex_pse, taxaType = 0,
    climate = c("bio1", "bio12"),
    selectedTaxa = crest_ex_selection, dbname = "crest_example"
  reconstr <- crest.calibrate(reconstr,</pre>
    geoWeighting = TRUE, climateSpaceWeighting = TRUE,
    bin_width = c(2, 20), shape = c("normal", "lognormal")
## End(Not run)
## example using pre-saved reconstruction obtained with the previous command.
data(reconstr)
dat <- plot_scatterPDFs(reconstr, save=FALSE,</pre>
                 taxanames=c(reconstr$inputs$taxa.name[c(2,4,5,1)]))
dat
```

plot_taxaCharacteristics

Plot the distribution and responses of the studied taxa

Description

Plot the distribution and responses of the studied taxa

Usage

```
plot_taxaCharacteristics(
    x,
    taxanames = x$inputs$taxa.name,
    climate = x$parameters$climate,
    col.density = viridis::plasma(20)[5:20],
    col.climate = viridis::viridis(22)[3:20],
    save = FALSE,
    filename = "taxaCharacteristics.pdf",
    as.png = FALSE,
    png.res = 300,
    width = 7.48,
    w0 = 0.2,
    height = 3 * length(climate) + h0,
    h0 = 0.4,
    add_modern = FALSE,
    resol = getResol(x)
```

Arguments

A crestObj generated by either the crest.calibrate, crest.reconstruct, loo or crest functions.
A list of taxa to use for the plot (default is all the recorded taxa).
Climate variables to be used to generate the plot. By default all the variables are included.
The colour gradient to use to map the density of species (top left map).
The colour gradient to use to map the climate gradients (left column).
A boolean to indicate if the diagram should be saved as a pdf file. Default is FALSE.
An absolute or relative path that indicates where the diagram should be saved. Also used to specify the name of the file. Default: the file is saved in the working directory under the name 'taxaCharacteristics.pdf'.
A boolean to indicate if the output should be saved as a png. Default is FALSE and the figure is saved as a pdf file.
The resolution of the png file (default 300 pixels per inch).
The width of the output file in inches (default 7.48in ~ 19cm).
The width of the left column with the names.
The height of the output file in inches (default 3in ~ 7.6cm per variables).
The vertical space used for the x-axes.
A boolean to add the location and the modern climate values to the plot (default $\mbox{\sf FALSE}).$
For advanced users only: if higher resolution data are used to estimate the pdfs, use this parameter to define the resolution of the maps on the figures. (default is 0.25 degrees to match with the default database)

Value

No return value, this function is used to plot.

```
## Not run:
    data(crest_ex_pse)
    data(crest_ex_selection)
    reconstr <- crest.get_modern_data(
        pse = crest_ex_pse, taxaType = 0, df = crest_ex,
        climate = c("bio1", "bio12"),
        selectedTaxa = crest_ex_selection, dbname = "crest_example"
)
    reconstr <- crest.calibrate(reconstr,
        geoWeighting = TRUE, climateSpaceWeighting = TRUE,
        bin_width = c(2, 20), shape = c("normal", "lognormal")
)
    plot_taxaCharacteristics(reconstr, taxanames='Taxon1')
## End(Not run)</pre>
```

66 plot_violinPDFs

plot_violinPDFs	Plot the pdfs as violins
-----------------	--------------------------

Description

Plot the pdfs as violins

Usage

```
plot_violinPDFs(
    x,
    climate = x$parameters$climate[1],
    taxanames = x$input$taxa.name,
    col = viridis::viridis(20),
    ylim = range(x$modelling$xrange[[climate]]),
    save = FALSE,
    filename = "violinPDFs.pdf",
    width = 7.48,
    height = 5,
    as.png = FALSE,
    png.res = 300
)
```

Arguments

	or crest functions.
climate	Climate variables to be used to generate the plot. By default all the variables are included.
taxanames	A list of taxa to use for the plot (default is all the recorded taxa).
col	A vector of colours that will be linearly interpolated to give a unique colour to each taxon.
ylim	The climate range to plot the pdfs on. Default is the full range used to fit the pdfs (x =modelling\$xrange)
save	A boolean to indicate if the diagram should be saved as a pdf file. Default is FALSE.
filename	An absolute or relative path that indicates where the diagram should be saved. Also used to specify the name of the file. Default: the file is saved in the working directory under the name 'violinPDFs.pdf'.
width	The width of the output file in inches (default 7.48in ~ 19cm).
height	The height of the output file in inches (default 3in ~ 7.6cm per variables).
as.png	A boolean to indicate if the output should be saved as a png. Default is FALSE and the figure is saved as a pdf file.
png.res	The resolution of the png file (default 300 pixels per inch).

A crestObj generated by either the crest.calibrate, crest.reconstruct

Value

A table with the climate tolerances of all the taxa

PSE_log 67

Examples

```
## Not run:
  data(crest_ex_pse)
  data(crest_ex_selection)
  reconstr <- crest.get_modern_data(</pre>
    pse = crest_ex_pse, taxaType = 0,
    climate = c("bio1", "bio12"),
    selectedTaxa = crest_ex_selection, dbname = "crest_example"
  reconstr <- crest.calibrate(reconstr,</pre>
    geoWeighting = TRUE, climateSpaceWeighting = TRUE,
    bin_width = c(2, 20), shape = c("normal", "lognormal")
  )
## End(Not run)
## example using pre-saved reconstruction obtained with the previous command.
data(reconstr)
ranges <- plot_violinPDFs(reconstr, save=FALSE, ylim=c(5,35),</pre>
                taxanames=c(reconstr$inputs$taxa.name[c(2,4,5,1)]),
                col=c('darkblue', 'firebrick3'))
lapply(ranges, head)
```

PSE_log

Displays the log of the proxy-species association

Description

Displays the log of the proxy-species association performed by the crest.get_modern_data() or crest.set_modern_data()

Usage

```
PSE_{log}(x)
```

Arguments

Χ

A crestObj generated by one of the following functions: crest.get_modern_data, crest.set_modern_data, crest.calibrate, crest.reconstruct, loo, or crest functions.

```
PSE_log(reconstr)
```

68 taxonComposition

reconstr

A crestObj ran with the example dataset.

Description

A crest0bj ran with the example dataset. Useful to illustrate many functions of the package.

Usage

reconstr

Format

An object of class crestObj of length 5.

taxonComposition

Return the size of the distribution of each composing species of each taxon

Description

Return the size of the distribution of each composing species of each taxon

Usage

```
taxonComposition(x, taxanames = x$input$taxa.name)
```

Arguments

x A crestObj generated by either the crest.calibrate, crest.reconstruct

or crest functions.

taxanames A list of taxa to use (default is all the recorded taxa).

Value

A list with the number of unique occurrences for each composing species

```
## Not run:
   data(reconstr)
   taxonComposition(reconstr)
## End(Not run)
```

testConnection 69

testConnection

Test if a connection can be established with the calibration data

Description

Test if a connection can be established with the calibration data

Usage

```
testConnection(dbname = "gbif4crest_02")
```

Arguments

dbname

The name of the data source database.

Value

TRUE if the connection can be established, else FALSE.

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