Package 'crestr'

November 6, 2020

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accClimateVariables

Describes all the variables available in the database.

Description

Describes all the variables available in the database.

Usage

```
accClimateVariables(v = NA)
```

Arguments

٧

The name of a variable to quickly access its description and ID.

Value

A data frame descriptive of the climate variables available in the database.

```
accClimateVariables()
```

accContinentNames 3

accContinentNames

Return the list of the continents and associated countries.

Description

Return the list of the continents and associated countries.

Usage

```
accContinentNames(dbname = "gbif4crest_02")
```

Arguments

dbname

The name of the database. Default is gbif4crest_02.

Value

A list with elements that correspond to the country names of each continent.

Examples

accContinentNames()

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(ecoregion = TRUE, dbname = "gbif4crest_02")
```

Arguments

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

dbname The name of the database. Default is gbif4crest_02.

Value

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

```
accRealmNames()
```

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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 All the climate values observed across the study area.

bin_width The width of the climate bins.

Value

A ccs object that will be used by fit_pdfsp.

Examples

```
# Extracting the number of taxa recorded in the database
calib_clim_space(sample(0:300 / 10, 4000, replace = TRUE), 2)
```

close_db_connection

Disconnect the database connection.

Description

Disconnect the database connection.

Usage

```
close_db_connection(db)
```

Arguments

db

An active database connection

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

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Connect to the gbif4crest database

Description

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

Usage

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

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.amazonaws.com

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.

Examples

```
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.

```
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>
```

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crest

Connect to the gbif4crest database

Description

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

Usage

```
crest(
  df,
  pse,
  taxaType,
  climate,
  xmn = -180,
  xmx = 180,
  ymn = -90,
  ymx = 90,
  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,
  geoWeighting = TRUE,
  climateSpaceWeighting = TRUE,
  presenceThreshold = 0,
  taxWeight = "normalisation",
  leave_one_out = FALSE,
  dbname = "gbif4crest_02"
)
```

Arguments

countries

A vector of the country names defining the study area.

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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 .

selectedTaxa .
bin_width .
shape .

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

climateSpaceWeighting

The number of points to be used to fit the pdfs.

presenceThreshold

.

taxWeight 'originalData', 'presence/absence', 'percentages' or 'normalisation'

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

computed (default FALSE).

dbname The name of the database. Default is gbif4crest_02.

Value

The parameters to be used by crest()

Examples

```
data(crest_ex)
data(crest_ex_pse)
data(crest_ex_selection)
recons <- 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
)
plot(recons)
plot_loo(recons)</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.get_modern_data 9

Usage

```
crest.calibrate(
    x,
    bin_width = rep(1, length(x$parameters$climate)),
    shape = rep("normal", length(x$parameters$climate)),
    npoints = 500,
    geoWeighting = TRUE,
    climateSpaceWeighting = TRUE
)
```

Arguments

```
x a crestObj produced by the crest.climate_space function.
bin_width .
shape .
npoints The number of points to be used to fit the pdfs.
geoWeighting The number of points to be used to fit the pdfs.
climateSpaceWeighting
The number of points to be used to fit the pdfs.
```

Value

A crest() object containing the spatial distributions and the climate space

Examples

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

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,
  taxa.name = unique(pse[, "ProxyName"]),
  xmn = -180,
  xmx = 180,
  ymn = -90,
  ymx = 90,
  continents = NA,
  countries = NA,
  realms = NA,
  biomes = NA,
  ecoregions = NA,
  minGridCells = 20,
  selectedTaxa = NA,
  dbname = "gbif4crest_02"
)
```

Arguments

pse .

taxaType .

climate A vectof of the climate variables to extract.

taxa.name .

xmn The coordinates defining the study area.
 xmx The coordinates defining the study area.
 ymn The coordinates defining the study area.
 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.

minGridCells .
selectedTaxa .

dbname The name of the database. Default is gbif4crest_02.

Value

A crest() object containing the spatial distributions

```
## Not run:
data(crest_ex_pse)
data(crest_ex_selection)
```

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```
x <- crest.get_modern_data(
  pse = crest_ex_pse, taxaType = 0,
  climate = c("bio1", "bio12"),
  selectedTaxa = crest_ex_selection, dbname = "crest_example"
)
x
lapply(x$modelling$distributions, head)
## End(Not run)</pre>
```

crest.reconstruct

Fit the species and proxy pdfs

Description

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

Usage

```
crest.reconstruct(x, df, presenceThreshold = 0, taxWeight = "normalisation")
```

Arguments

Value

A crest() object containing the reconstructions and all the associated data.

```
data(crest_ex)
data(crest_ex_pse)
data(crest_ex_selection)
x <- crest.get_modern_data(
    pse = crest_ex_pse, taxaType = 0,
    climate = c("bio1", "bio12"),
    taxa.name = colnames(crest_ex)[-1],
    selectedTaxa = crest_ex_selection, dbname = "crest_example"
)
x <- crest.calibrate(x,
    geoWeighting = TRUE, climateSpaceWeighting = TRUE,
    bin_width = c(2, 20), shape = c("normal", "lognormal")
)
x <- crest.reconstruct(x, crest_ex)
plot(x)</pre>
```

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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 7 columns (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.

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 integr indicating the taxonomic resolution (1 family, 2 genus, 3 species, 4 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 13

crest_ex_selection

Example dataset to associate taxa with climate varibles.

Description

A data frame indicating which taxa should be used to reconstruct each climate variable.

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)

crest0bj

Create a crest() object.

Description

Creates a crest() object with all default parameters.

Usage

```
crestObj(
  taxa.name,
  pse,
  taxaType,
  climate,
  xmn,
  xmx,
  ymn,
  ymx,
  continents,
  countries,
  realms,
  biomes,
  ecoregions,
  df = NA,
  x = NA,
  x.name = "",
  minGridCells = 20,
  bin_width = rep(1, length(climate)),
  shape = rep("normal", length(climate)),
  npoints = 500,
  geoWeighting = TRUE,
```

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```
climateSpaceWeighting = TRUE,
  selectedTaxa = NA,
  presenceThreshold = 0,
  taxWeight = "normalisation"
)
```

Arguments

```
taxa.name
pse
taxaType
climate
                  A vectof of the climate variables to extract.
xmn, xmx, ymn, ymx
                  The coordinates defining the study area.
                  A vector of the continent names defining the study area.
continents
                  A vector of the country names defining the study area.
countries
                  A vector of the studied botanical realms defining the study area.
realms
                  A vector of the studied botanical biomes defining the study area.
biomes
ecoregions
                  A vector of the studied botanical ecoregions defining the study area.
df
x.name
minGridCells
bin_width
shape
                  The number of points to be used to fit the pdfs.
npoints
geoWeighting
                  The number of points to be used to fit the pdfs.
climateSpaceWeighting
                  The number of points to be used to fit the pdfs.
selectedTaxa
presenceThreshold
                  'originalData', 'presence/absence', 'percentages' or 'normalisation'
taxWeight
```

Value

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

dbRequest 15

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 database. Default is gbif4crest_02.

Value

The result of the request.

Examples

```
# 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.
## Not run:
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>
```

fit_pdfsp

Fit the species pdfs.

Description

Fit the species pdfs.

Usage

```
fit_pdfsp(climate, ccs, bin_width, shape, xrange, use_ccs = TRUE)
```

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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 climate bins.

shape The shape of the species pdfs. Use 'normal' or 'lognormal'. xrange The climate gradient upon which the pdf with be defined.

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

modern climate space

Value

The pdf of the species.

Examples

```
# 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)
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.
sum(pdfsp * (xrange[2] - xrange[1])) == 1</pre>
```

fit_xrange

Define teh climate gradient to fit the pdfs.

Description

Define teh 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 shape of the species pdfs. Use 'normal' or 'lognormal'.

bin_width The width of the climate bins.

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

Value

A regularly spaced climate gradient with npoints points.

getClimateSpace 17

Examples

```
# 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>
```

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 = -180,
  xmx = 180,
  ymn = -90,
  ymx = 90,
  continents = NA,
  countries = NA,
  realms = NA,
  biomes = NA,
  ecoregions = NA,
  dbname = "gbif4crest_02"
)
```

Arguments

climate A vectof of the climate variables to extract. 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

A matrix of occurrence records with the associated climate.

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See Also

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

Examples

```
climate <- getClimateSpace("bio1", -90, 90, -90, 90,
  continents = "Europe",
  countries = c("Germany", "Netherlands", "Sweden"),
  realms = "Palaearctic"
)
head(climate)
plot(climate[, -3], asp = 1)</pre>
```

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,
  xmn = -180,
  xmx = 180,
  ymn = -90,
  ymx = 90,
  continents = NA,
  countries = NA,
  realms = NA,
  biomes = NA,
  ecoregions = NA,
  dbname = "gbif4crest_02"
)
```

Arguments

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

climate A vector of the climate variables to extract.

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.

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.

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Value

A matrix of occurrence records with the associated climate.

See Also

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

Examples

```
taxIDs <- getTaxonID("Zamiaceae", "Ceratozamia")
distrib <- getDistribTaxa(taxIDs, "bio1", -90, 90, -90, 90,
  continents = "Europe",
  countries = c("Germany", "Netherlands", "Sweden"),
  realms = "Palaearctic"
)
distrib</pre>
```

getTaxonID

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 5) to define the type of palaeoproxy used: 1

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

6 for rodents.

dbname The name of the database. Default is gbif4crest_02.

Value

A vector of unique taxonIDs.

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Examples

```
getTaxonID("Zamiaceae")
getTaxonID("Zamiaceae", "Ceratozamia")
getTaxonID("Zamiaceae", "Ceratozamia", taxaType = 2)
```

 ${\tt getTaxonomy}$

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"
)
```

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 foraminifers, 4 for diatoms, 5 for chironomids 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 database. Default is gbif4crest 02.

Value

A vector of unique taxonIDs.

```
getTaxonomy("Zamiaceae")
getTaxonomy(genus="Ceratozamia", depth.out=8)
getTaxonomy("Zamiaceae", "Ceratozamia", taxaType = 2)
```

isColourStr 21

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')
```

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Connect to the gbif4crest database

Description

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

Usage

loo(x)

Arguments

Χ

a crestObj produced by the crest.reconstruct() or crest() functions.

Value

A crest() object containing the reconstructions and all the associated data.

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Examples

```
data(crest_ex)
data(crest_ex_pse)
data(crest_ex_selection)
recons <- 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"
)
recons <- loo(recons)
recons$reconstructions$bio12$loo
plot_loo(recons)</pre>
```

meanPositiveValues

Convert data into presence/absence data.

Description

Convert data into presence/absence data.

Usage

```
meanPositiveValues(x)
```

Arguments

Х

A vector of values.

Value

The average of all the positive values.

Examples

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

normalise

Convert data into presence/absence data.

Description

Convert data into presence/absence data.

Usage

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

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

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,
  yax_incr = 5,
  bar_width = 1,
  xlim = NA,
  tickAtSample = TRUE,
  col_pos = "black",
  col_neg = "white",
  title = NA
)
```

Arguments

bars

A data frame of the data to plot (first column with age or depth) and the taxa in х the following columns. x can also be a crestObj.

A boolean that indicates if the data should be plotted as polygons (default: bars=FALSE) or vertical bars (bars=TRUE).

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col	Colours to be used for the polygons. If the number of colours does not match the number of taxa, colors will be recyled.
amplif	A factor the show exageration on the diagram. Only for polygon plot. Default 5.
save	A boolean to indicate if the diagram shoud 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.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).
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. Canbe adjusted to get round numbers on the x-ais. If smaller than the range overed 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 white).
title	Name to be added on top of the plot (default NA).

Examples

plot_loo

Plot stratigraphic data as polygons or barplots.

Description

This function plots stratigraphic data either as polygons or bars.

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Usage

```
plot_loo(
    x,
    save = FALSE,
    filename = "Diagram_loo",
    width = 3.54,
    height = 9,
    yax_incr = NA,
    bar_width = 1,
    xlim = NA,
    tickAtSample = FALSE,
    col_pos = "black",
    col_neg = "white",
    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.
save		A boolean to indicate if the diagram shoud be saved as a pdf file. Default is FALSE.
filen	ame	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).
heigh	t	The height of the output file in inches (default 9in ~ 23cm).
yax_i	ncr	Graphical parameters describing the increment size on the y-axis (default 5).
bar_w	idth	Width of the bars of the barplot (default 1).
xlim		The range covered by the x-axis. Canbe adjusted to get round numbers on the x-ais. If smaller than the range overed by the data, the data will be truncated (default: range of the data).
tickA	tSample	Boolean that indicates whether a tick mark should be added on the x-axis at the location of each sample (default TRUE).
col_p	os	Graphical parameter for the barplot. Colour of all the positive values (default black).
col_n	eg	Graphical parameter for the barplot. Colour of all the negative values (default white).
title		Name to be added on top of the plot (default NA).

```
data(crest_ex)
data(crest_ex_pse)
data(crest_ex_selection)
recons <- 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"</pre>
```

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