# Package 'speciesgeocodeR'

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Title Prepare Species Distributions for the Use in Phylogenetic

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Analyses

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<b>Description</b> Preparation of species occurrences and distribution data for the use in phylogenetic analyses. SpeciesgeocodeR is built for data cleaning, data exploration and data analysis and especially suited for biogeographical and ecological questions on large datasets. The package includes the easy creation of summary-tables and -graphs and geographical maps, the automatic cleaning of geographic occurrence data, the calculating of coexistence matrices and species ranges (EOO) as well as mapping diversity in geographic areas.
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speciesgeocodeR-package
CalcRange
CoExClass
GeoClean
GetElevation
IUCNest
landmass
lemurs
lemurs_in
lemurs_test
MapGrid
MapRichness

mdg_poly	. 14
plot.spgeoIN	. 15
plot.spgeoOUT	
PlotHull	17
RangeRichness	18
ReadPoints	18
RichnessGrid	19
SpeciesGeoCoder	20
Spgc2BioGeoBEARS	. 22
SpGeoCod	23
SpGeoCodH	24
summary.spgeoIN	25
summary.spgeoOUT	26
WriteOut	26
WwfLoad	27
	29

speciesgeocodeR-package

Prepare Species Distributions for the Use in Phylogenetic Analyses

## **Description**

Index

Preparation of species occurrences and distribution data for the use in phylogenetic analyses. Species-geocodeR is built for data cleaning, data exploration and data analysis and especially suited for biogeographical and ecological questions on large datasets. The package includes the easy creation of summary-tables and -graphs and geographical maps, the automatic cleaning of geographic occurrence data, the calculating of coexistence matrices and species ranges (EOO) as well as mapping diversity in geographic areas.

#### Details

Package: speciesgeocodeR

Type: Package Version: 1.04 Date: 2015-10-17 License: GPL-3

The core functions of the package can be used via the wrapper function SpeciesGeoCoder, with two input text files and one single command. In addition to this the package includes a set of functions to visualize geographic distribution data and diversity.

#### Author(s)

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CalcRange 3

#### **Examples**

```
## Not run:
data(lemurs)
data(lemurs)
data(mdg_poly)

SpeciesGeoCoder(lemurs, mdg_poly)

## End(Not run)

data(lemurs)
data(mdg_poly)
inp <- ReadPoints(lemurs, mdg_poly)

outp <- SpGeoCodH(inp)
e <- c(42, 52, -27, -10)
ivesia_abu <- RichnessGrid(outp, e, reso = 60, type = "abu")

outp <- SpGeoCodH(inp)
e <- c(42, 52, -27, -10)
lemurs_div <- RichnessGrid(outp, e, reso = 60, type = "spnum")

MapGrid(lemurs_div)</pre>
```

CalcRange

Range polygons, AOO and EOO from occurrence points

#### **Description**

Calculate the Area of Occupancy based on a user-defined grid as well as range polygons and the Extent of Occurrence based on the areaPolygon function of the geosphere, which can be used for criterion B of the IUCN red list classification using IUCNest.

#### Usage

```
CalcRange(x, index = c("A00", "E00"), eoo.value = c("area", "shape"),
eoo.terrestrial = TRUE, aoo.gridsize = 4, aoo.reps = 3,
aoo.xmin = NULL, aoo.xmax = NULL, aoo.ymin = NULL,
aoo.ymax = NULL, verbose = FALSE)
```

#### **Arguments**

x data.frame of species occurrence records, with 3 columns: 1 = species name, 2 = longitude coordinates, y = latitude coordinates.

index character string, setting the mode of calculation. Default is both.

eoo.value character string, setting the output value. If value = "area": A data.frame with

the EOO for each species in square kilometers; if value = "shape", a list of

convex hull polygons.

eoo.terrestrial

aoo.gridsize

logial. If TRUE, the shapes are clipped to continent borders. Only relevant, if

EOO is calculated.

The area of the grid cells used for the AOO calculation in square kilometers.

Folloeing the IUCN recommendations the default is to 4.

4 CoExClass

aoo.reps	The number of replicates for the AOO calculation. See details. Default = $3$ .
aoo.xmin	The minimum longitude for the AOO grid in degrees. If not specified it will be estimated from the minimum in the data.
aoo.xmax	The maximum longitude for the AOO grid in degrees. If not specified it will be estimated from the minimum in the data.
aoo.ymin	The minimum latitude for the AOO grid in degrees. If not specified it will be estimated from the minimum in the data.
aoo.ymax	The minimum latitude for the AOO grid in degrees. If not specified it will be estimated from the minimum in the data.
verbose	If TRUE, status will be reported.

#### **Details**

As the starting point of the AOO grid can influence the result (depending on ehere the boarders of the gridcells fall, occurrence points might or might not be in the same cell by chance), a replication with different grids is recommended. The starting position of each grid is determined by aoo.xmin and aoo.ymin plus (sqrt(aoo.gridsize) \* 1000) / aoo.reps. SO the higher the number of replicates the smaller the difference between the replicates will be.

#### Value

If value is not "shape" an object of the class "range.sizes" with a data.frame of AOO or EOO values (or both) in square kilometers, if value = "shape", a list of convex hull polygons.

## Note

See the speciesgeocodeR documentation for further information and examples.

# **Examples**

```
data(lemurs)
CalcRange(lemurs, index = "E00", eoo.value = "shape", eoo.terrestrial = F)
## Not run:
CalcRange(lemurs, index = "E00", eoo.value = "area", eoo.terrestrial = F)
CalcRange(lemurs, index = "A00", eoo.value = "area")
CalcRange(lemurs, index = c("A00", "E00"), eoo.value = "area", eoo.terrestrial = F)
## End(Not run)
```

CoExClass

Species Coexistence Matrices in Given Areas.

# Description

Add a coexistence matrix to an object of the class spgeoOUT created by SpGeoCod.

# Usage

```
CoExClass(x)
```

countryref 5

#### **Arguments**

Χ

An object of the class spgeoOUT, where the coexistence matrix shall be added

#### Value

An object of the class spgeoOUT, including a coexistence matrix which can be addressed with via the "coexistence\_classified" slot. The matrix will have the same dimensions as there are input species and will give the percent of occurrence points of each species (row) with all other species (columns) given the input polygons.

## Note

See the speciesgeocodeR tutorial for further information and examples.

## **Examples**

```
data(lemurs)
data(mdg_poly)

inp <- ReadPoints(lemurs, mdg_poly)
outp <- SpGeoCodH(inp)
outpcoex <- CoExClass(outp)
outpcoex$coexistence_classified</pre>
```

countryref

Country Centroids and Country Capitals

## **Description**

A data.frame with coordinates of country centroids and country capitals as reference for the GeoClean function. Coordinates are based on the Central Intelligence Agency World Factbook as provided at http://opengeocode.org/download/cow.php.

# Usage

```
data("countryref")
```

#### **Format**

A data frame with 249 observations on 7 variables.

#### **Source**

CENTRAL INTELLIGENCE AGENCY (2014) The World Factbook, Washington, DC.

```
http://opengeocode.org/download/cow.php
```

```
data(countryref)
```

GeoClean

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Automated Cleaning of Geographic Coordinates

#### **Description**

Provides a number of different tests to clean datasets with geographic coordinates. Each function argument represents a different cleaning step.

# Usage

# **Arguments**

a data. frame with at least three columns: "identifier" (species name), "XCOOR"

(longitude) and "YCOOR" (latitude). Column names might also be "species", "longitude" and "latitude". If the arguments "countrycentroid", "capitalcoords" or "countrycheck" should be used, a fourth column named "country" is needed with the country names in ISO2 or ISO3. Alternatively, a data.frame as down-

loaded from GBIF.

isna logical. If TRUE, checks for missing values in the coordinates. Default = TRUE.

isnumeric logical. If TRUE, checks for non-numeric values in the coordinates. Default =

TRUE.

coordinatevalidity

logical. If TRUE, checks for non-valid coordinates (XCOOR > 180 and < -180;

YCOOR >90 and <-90). Default = TRUE.

containszero logical. If TRUE, checks for coordinates that are exactly zero. Default = TRUE.

zerozero logical. If TRUE, checks if the coordinate fall within a rectangle around the

point 0/0. Default = TRUE.

zerozerothresh numeric. The size of the rectangle around 0/0 in decimal degrees. Default = 0.5.

latequallong logical. If TRUE, checks for rows where XCOOR = YCOOR. Default = TRUE.

GBIFhead logical. If TRUE, checks if the coordinate fall within a 0.5 degree rectangle

around the GBIF headquarters in Copenhagen. Default = FALSE.

countrycentroid

logical. If TRUE checks if the coordinate fall within a rectangle around the centroid of the country specified in x\$country. The size of the rectangle can be

controlled using the "countthresh" argument. Default = FALSE.

contthresh numeric. The size of the rectangle around the country centroid (in degrees). The

number is half the length of one rectangle side. Default = 0.5.

GeoClean 7

capitalcoords logical. If TRUE, checks if the coordinate fall within a rectangle around the capital of the country specified in x\$country. The size of the rectangle can be controlled using the "countthresh" argument. Default = FALSE.

capthresh numeric. The size of the rectangle around the capital (in degrees). The number

is half the length of one rectangle side. Default = 0.5.

countrycheck logical. If TRUE, checks if the coordinates fall within the country borders of the

country indicated in x\$country. Default = FALSE.

polygons The reference polygons for the countrycheck function. By default the wrld\_simpl

dataset from the maptools package. The maptools package must be loaded to

use countrycheck = T.

referencecountries

The reference coordinates for the country centroids and capitals. By default

from the countryref data.

outp character defining the output values. See value section.

#### **Details**

The capital and country centroids in the country ref dataset are from the CIA World Factbook. The check for country borders is based on the world\_simpl data from the maptools package. Please note that the ISO2 code for Namibia ("NA") might cause problems with the countrycheck argument. If possible use ISO3 country codes.

#### Value

if outp = 'summary', a vector of the same length as the input data.frame with TRUE = clean coordinates, FALSE = suspicious coordinates. If outp = 'detailed', a data.frame with one column for each check that was performed: TRUE = clean coordinates, FALSE = suspicious coordinates. If outp = 'cleaned', a cleaned version of the input data.

# Note

See the speciesgeocodeR documentation for further information and examples.

## References

CENTRAL INTELLIGENCE AGENCY (2014) The World Factbook, Washington, DC.

http://opengeocode.org/download/cow.php

8 GetElevation

GetElevation

Elevation Data for Multiple Species

# **Description**

A wrapper using the getData function of the raster package to get elevation information for each occurrence point in x.

# Usage

GetElevation(x)

#### **Arguments**

Х

an object of the class data.frame, spgeoIN, spgeoOUT or a character string.

#### **Details**

If x is a data.frame, it must have 3 columns: species name, longitude and latitude. If x is a character string it can be the path to a tab delimited .txt file with three columns: species name, longitude and latitude. If x is a character string of species names, the function will use rgbif to download occurrence information from GBIF. In this case no data cleaning is performed, except from the 'removeZeros' option of rgbif.

#### Value

A vector of elevation values.

# See Also

ReadPoints, SpGeoCod.

```
data(lemurs)
## Not run:
GetElevation(lemurs)
## End(Not run)
```

IUCNest 9

IUCNest	Convert Range Sizes to IUCN Assessment	

#### **Description**

The function converts range sizes (Area of Occupancy and Extent of Occurrence), as calculated by CalcRange to conservation assessments based on IUCN criterion B and extinction probabilities following Mooers et al. (2008).

## Usage

```
\label{eq:locality} \begin{split} &\text{IUCNest(x, value = c("all", "A00", "E00", "IUCN50", "IUCN100", "IUCN500"),} \\ &\text{NT.thresh = c(30000, 3000), VU.thresh = c(20000, 2000),} \\ &\text{EN.thresh = c(5000, 500), CR.thresh = c(100, 10))} \end{split}
```

#### **Arguments**

X	an object of the class range.sizes, as produced by CalcRange.
value	character string. DEfining the output value, see details. Default = "all".
NT.thresh	numeric. Thresholds for categorization to near threatend, in the form (EOO, AOO). Defaults from (IUCN Standards and Petitions Subcommittee 2014).
VU.thresh	numeric. Thresholds for categorization to vulnerable, in the form (EOO, AOO). Defaults from (IUCN Standards and Petitions Subcommittee 2014).
EN.thresh	numeric. Thresholds for categorization to endangered, in the form (EOO, AOO). Defaults from (IUCN Standards and Petitions Subcommittee 2014).
CR.thresh	numeric. Thresholds for categorization to critically endangered, in the form (EOO, AOO). Defaults from (IUCN Standards and Petitions Subcommittee 2014).

#### Details

The value argument defines the type of assessment in the output: "AOO" = assessment based on the AOO, "EOO" = assessment based on the EOO, "both" = assessment based on AOO and EOO, "IUCN50" = a extra column where the assessment is converted into a probability of extinction within the next 50 years, "IUCN100" = a extra column where the assessment is converted into a probability of extinction within the next 100 years, "IUCN500" = a extra column where the assessment is converted into a probability of extinction within the next 500 years. See Mooers et al. 2008 for reference on the conversion.

#### Value

a data.frame, the number of columns depend on the value argument (see details).

#### References

IUCN Standards and Petitions Subcommittee 2014 Guidelines for using the IUCN Red List categories and criteria. Version 11. Prepared by the Standards and Petitions Subcommittee. Downloadable from: http://www.iucnredlist.org/documents/RedListGuidelines.pdf.

Mooers, A.O., Faith D.P. & MAddison W. P. 2008 Converting Endangered Species Categories to Probabilities if Extinction for Phylogenetic Conservation Prioritization. *PLOSone*, **3**(11)

10 lemurs

## **Examples**

```
data(lemurs)
rang <- CalcRange(lemurs, terrestrial = F)
IUCNest(rang)</pre>
```

landmass

Global Coastlines

# Description

A SpatialPolygonsDataFrame with global coastlines.

## Usage

```
data("landmass")
```

#### **Source**

```
http://www.naturalearthdata.com/downloads/10m-physical-vectors/
```

## **Examples**

```
data("landmass")
plot(landmass)
```

lemurs

Distribution of Lemur Species

# Description

A data.frame with point occurrences of 39 Lemur species from Madagascar as example for the input format for the GeoClean function. The data is from GBIF.

## Usage

```
data("lemurs")
```

## **Format**

A data frame with 403 observations on the following 3 variables.

```
identifier species name
XCOOR longitude
YCOOR latitude
```

#### **Source**

```
www.gbif.org.
```

```
data(lemurs)
str(lemurs)
```

lemurs\_in 11

lemurs\_in

Example for an spgeoIN Object

# **Description**

This is an example dataset for an object of the class spgeoIN (created by ReadPoints). It contains occurrence data for 39 species of Lemurs in Madagascar. The data has 3 slots: identifier = a vector of character strings containing the species name for each occurrence points species\_coordinates = a data.frame with the longitude and latitude positions polygons = a SpatialPolygons object with a set of 3 polygons.

# Usage

```
data("lemurs_in")
```

#### See Also

ReadPoints

#### **Examples**

```
data(lemurs_in)
summary(lemurs_in)
plot(lemurs_in)
```

lemurs\_test

Distribution of Lemur Species with Problematic Records

## **Description**

A data. frame with point occurrences of 42 Lemur species in the right input format for the GeoClean. The data is downloaded from GBIF but has some added erroneous coordinates.

## Usage

```
data("lemurs_test")
```

# Format

A data frame with 627 observations on the following 4 variables.

```
identifier species name

XCOOR longitude

YCOOR latitude

country a factor with levels MDG
```

#### Source

```
www.gbif.org
```

12 MapGrid

# **Examples**

```
data(lemurs_test)
str(lemurs_test)
```

MapGrid

Plotting Rasters in the Geographical Context

# Description

A wrapper function to map a raster with country borders. Can be used with the results of RichnessGrid.

# Usage

```
MapGrid(rast, ...)
```

# Arguments

rast A raster object.

. . . Additional arguments past to plot.

# Note

See the speciesgeocodeR tutorial for further information and examples.

## See Also

RichnessGrid

```
data(lemurs)
data(mdg_poly)

inp <- ReadPoints(lemurs, mdg_poly)
outp <- SpGeoCodH(inp)
e <- c(42, 52, -27, -10)
ras <- RichnessGrid(outp, e, reso = 60, "abu")
MapGrid(ras)</pre>
```

MapRichness 13

MapRichness	Plotting Species Richness in a Set of Polygons

# **Description**

Illustrates the species number per polygon, based on and object of the class spgeoOUT (created by SpGeoCod).

# Usage

# Arguments

x	an object of the class "spgeoOUT".
areanames	a character string. If the polygons in the spgeoIN object were derived from a shape file, this column indicates the name of the column used for area names. If the polygons in the spgeoIN object were derived from a text file leave as default (Default = "").
leg	Controls the plot legend and coloring scheme. Must be either "continuous" or "discrete", (default = continuous).
show.occ	logical. Defines if the occurrence points should be added to the plot (default = F).
lin.col	character string. Defines the colour of polygon borders. If NA, the same colour as for the polygon is used (default = NA).
lwd	numeric. Line width of polygon lines.
•••	Arguments to be passed to other methods, such as graphical parameters (see par).

## Value

A plotted map.

# Note

See the speciesgeocodeR tutorial for further information and examples.

# See Also

RichnessGrid, MapGrid, RangeRichness.

```
data(lemurs)
data(mdg_poly)

inp <- ReadPoints(lemurs, mdg_poly)
outp <- SpGeoCodH(inp)
MapRichness(outp)</pre>
```

14 mdg\_poly

mdg\_biomes

WWF Biomes of Madagascar

## **Description**

A SpatialPolygonsDataFrame of a simplified version of the WWF biomes of Madagascar.

#### Usage

```
data("mdg_biomes")
```

#### References

Olson, D. M., Dinerstein, E., Wikramanayake, E. D., Burgess, N. D., Powel, G. V. N., Underwood, E. C., Damico, J. A., Itoua, I., Strand, H. E., Morrison, J. C., Loucks, C. J., Ricketts, T. H., Kura, Y., Lamoreux, J. F., Wettengel, W. W., Hedao, P., and Kassem, K.R. 2001 Terrestrial ecoregions of the world: A new map of life on earth. *BioScience*, **51**(11):933–938.

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

#### **Examples**

```
data(mdg_biomes)
str(mdg_biomes)
```

mdg\_poly

WWF Biomes of Madagascar in Table Format

# Description

A data. frame of a simplified version of the WWF biomes of Madagascar (simplified).

# Usage

```
data("mdg_poly")
```

#### **Format**

A data frame with 87 observations on the following 3 variables.

identifier a factor with levels Dry Forest Moist Forest Shrublands XCOOR a numeric vector

YCOOR a numeric vector

## References

Olson, D. M., Dinerstein, E., Wikramanayake, E. D., Burgess, N. D., Powel, G. V. N., Underwood, E. C., Damico, J. A., Itoua, I., Strand, H. E., Morrison, J. C., Loucks, C. J., Ricketts, T. H., Kura, Y., Lamoreux, J. F., Wettengel, W. W., Hedao, P., and Kassem, K.R. 2001 Terrestrial ecoregions of the world: A new map of life on earth. *BioScience*, **51**(11):933–938.

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

plot.spgeoIN 15

## **Examples**

```
data(mdg_poly)
str(mdg_poly)
```

plot.spgeoIN

Plot Method for SpgeoIN Objects

# Description

plot.spgeoIN is the plotting method for objects of the class spgeoIN. It is designed to give an overview of the data and plots the input points and polygons on a world map.

## Usage

```
## S3 method for class 'spgeoIN'
plot(x, ...)
```

# **Arguments**

x an object of the class spgeoIN

... arguments to be passed to methods, such as graphical parameters (see par).

## See Also

ReadPoints

# **Examples**

```
data(lemurs)
data(mdg_poly)
inp <- ReadPoints(lemurs, mdg_poly)
plot(inp)</pre>
```

plot.spgeoOUT

Plot Method for SpgeoOUT Objects

# **Description**

plot.spgeoOUT is the plotting method for objects of the class spgeoOUT. Depending on the plottype argument the function can be used to get an overview over the data or plot all elements of the object.

# Usage

16 plot.spgeoOUT

## **Arguments**

Х		an object of the class spgeoOUT
р	olottype	a character string. Controls which data shall be plotted. Default is to "summary". See details for more information.
р	olotout	logical. If TRUE, no user-input is needed between the plots (par(ask = FALSE)), if FALSE, user input is needed to browse between plots (par(ask = TRUE)).
m	ode	a character string. Can be "percent" or "total" controls if percentage or total is shown for plottype = "species". Default is "percent".
m	oreborders	logical. If TRUE additional borders are added to the maps. Default is FALSE.
а	reanames	character. character string. If the polygons of the spgeoOUT object were derived from a shape file, this column indicates the name of the column used for area names. See the SpeciesgeocodeR documentation for more information, if the polygons of the spgeoOUT object were derived from a text file leave as default (Default = "").
		arguments to be passed to methods, such as graphical parameters (see par).

#### **Details**

The output will be different depending on the plottype argument. plottype = "summary": a map with all occurrence points and polygons (the red points were not classified to any polygon), the species number for all polygons, and if available, a heatplot visualizing the coexistence matrix; plottype = "species": a bar chart for each input species showing the number of occurrences in each polygon; plottype = "polygons": a bar chart for each input-polygon showing the number of occurrences for each species in the polygon; plottype = "speciesrichness": a bar chart showing species number per polygon; plottype = "coexistence": a heatplot visualizing the coexistence matrix. The heatcolors code for the percent of occurrences that are shared (per rows); plottype = "mapspecies": one map per species showing all occurrence points in all polygons; plottype = "mappolygons": a series of maps, showing each polygon and its close environment, with all samples classified to this polygon. Species are color-coded; plottype = "mapunclassified": a map with all sample points that could not be classified to any polygon; plottype = "mapall": a map of all occurrence points in the object.

## See Also

```
summary.spgeoOUT, SpGeoCodH
```

```
data(lemurs)
data(mdg_poly)

inp <- ReadPoints(lemurs, mdg_poly)
outp <- SpGeoCodH(inp)
plot(outp,
plot(outp, plottype = "species")
plot(outp, plottype = "polygons")
plot(outp, plottype = "speciesrichness")
plot(outp, plottype = "coexistence")
plot(outp, plottype = "mapspecies")
plot(outp, plottype = "mappolygons")
plot(outp, plottype = "mapunclassified")
plot(outp, plottype = "mapall")</pre>
```

PlotHull 17

PlotHull Plotting Species Ranges
----------------------------------

## **Description**

Plots species distribution ranges derived with CalcRange in the geographic context.

# Usage

```
PlotHull(dat, xlim = c(-180, 180), ylim = c(-90, 90),

col = rgb(255, 0, 0, 50, maxColorValue = 255),

border = rgb(255, 0, 0, 50, maxColorValue = 255),

type = "overlay", select = "all", ...)
```

# **Arguments**

dat	a list of 'SpatialPolygons' objects created with CalcRange.
xlim	vector indicating the x-axis plotting limits. Default is c(-180, 180).
ylim	a vector indicating the y-axis plotting limits. Default = $c(-90, 90)$ .
col	character string indicating the colour of the polygons. Default is to transparent red.
border	a character string indicating the color of the polygon borders. Default is to transparent red.
type	a character string indicating if all range polygons in the input file shall be overlaid or plotted on separate plots. Must be "seperate" or "all". Default is to "all" (overlay).
select	a character string indicating which elements of the list to plot. Can be one or multiple species names. Default = "all".
	additional arguments passed to plot.

# **Details**

If type is set to "separate", set par(ask = T) to see all maps.

# Value

A plot or a set of plots.

18 ReadPoints

Species Richness Raster from Range Polygons

# Description

Calculate species richness for the specified grid based on an list of range polygons, as derived from CalcRange.

## Usage

```
RangeRichness(ra, limits = c(-180, 180, -90, 90), reso = 60, terrestrial = FALSE)
```

## **Arguments**

ra A list of SpatialPolygons with species ranges.

limits numeric. A vector with the limits of the grid in decimal degrees of the format

c(xmin, xmax, ymin, ymax). Default is worldwide c(-180, 180, -90, 90).

reso numeric. Resolution of the grid in minutes.

terrestrial logical. If TRUE, oceans are omitted.

#### Value

```
a raster object.
```

# **Examples**

```
data(lemurs)
rang <- CalcRange(lemurs, value = "shape")
sprich <- RangeRichness(rang)
MapGrid(sprich)</pre>
```

ReadPoints

Loading Data into SpeciesgeocodeR

# Description

Create an object of the class spgeoIN from the input arguments.

# Usage

```
ReadPoints(x, y, areanames = NA, verbose = FALSE, cleaning = FALSE)
```

RichnessGrid 19

#### **Arguments**

У

a data.frame with three columns named identifier, XCOOR, YCOOR or a character string. If x is a character string, it can either be the path to a tab-delimited text file containing the coordinates of points of interest (tab delimited, three columns with header: identifier, XCOOR, YCOOR) OR a vector of species names. In the latter cases GBIF is searched for occurrence data using the rgbif function of the rgbif package.

character string giving the path to a text file containing the polygons of points of interest (tab delimited, three columns with header: identifier, XCOOR, YCOOR)

OR an object of the class SpatialPolygons OR an object of the class SpatialPolygonsDataFrame.

areanames character string. If the polygons in the spgeoIN object were derived from a

shape file or a SpatialPolygonsDataframe, this column indicates the name of the column used for area names. See the Speciesgeocoder tutorial for more information. If the polygons in the spgeoIN object were derived from a text file

leave as default (Default = NA).

verbose logical. If TRUE, the functions report the progress of the analyses. This is

advisable for large datasets.

cleaning logical. If TRUE, GeoClean is applied to the input coordinates. Default =

FALSE.

... Further arguments to be passed to GeoClean.

#### Value

An object of class spgeoIN.

## Note

See the speciesgeocodeR tutorial for further information and examples.

## See Also

```
lemurs_in
```

# Examples

```
data(lemurs)
data(mdg_poly)
inp <- ReadPoints(lemurs, mdg_poly)</pre>
```

RichnessGrid

Species Richness and Abundance Grids from Point Records

# Description

Create a species richness or abundance grid, from a spgeoOUT object or a data.frame with species names and occurrence coordinates.

20 SpeciesGeoCoder

#### Usage

```
RichnessGrid(x, limits = c(-180, 180, -90, 90), reso,
type = c("spnum", "abu"))
```

#### **Arguments**

x an object of the class data. frame or spgeoIN or spgeoOUT or character string. If x is a data. frame, it must consist of three columns, indicating species name, longitude and latitude. If x is a character string, it can be the name of a tab delimited text file, with three columns indicating species name, longitude and latitude or a vector of species names. In the latter case the GBIF database is

search for geo-referenced occurrences.

limits numerical. A vector of the form (xmin, xmax, ymin, ymax) givin the raster

limits in decimal degrees.

reso numeric. resolution of the grid in minutes.

type character string indicating the type of data displayed: "spnum" = number of

species per grid cell, "abu" = the number of occurrence points per grid cell.

#### Value

a raster object.

#### See Also

MapRichness, MapGrid, RangeRichness.

## **Examples**

```
# for x = data.frame
data(lemurs)
e <- c(-125, -105, 30, 50)
dat <- RichnessGrid(lemurs, e, reso = 60, type = "spnum")

# for x = character string
e <- c(-125, -105, 30, 50)

## Not run:
dat <- RichnessGrid("Indri indri", e, reso = 60, type = "spnum")
## End(Not run)</pre>
```

 ${\tt SpeciesGeoCoder}$ 

A Complete SpeciesgeocodeR Analysis

# Description

A complete occurrence to area classification with a set of standard output files saved to the working directory.

SpeciesGeoCoder 21

#### Usage

#### **Arguments**

х	a data.frame with three columns named identifier, XCOOR, YCOOR or a character string. If x is a character string it can be the path to a tab-delimited text file containing the coordinates of points of interest (tab delimited, three columns with header: identifier, XCOOR, YCOOR) OR a shape file OR a vector of species names. In the latter cases GBIF is search for occurrence data using the rgbif function.
У	a character string giving the path to a text file containing the polygons of points of interest (tab delimited, three columns with header: identifier, XCOOR, YCOOR) OR an object of the class SpatialPolygonsOataframe.
coex	logical. Indicating if a coexistence matrix should be calculated and shown as a heat plot (default = FALSE). Only for small to medium-sized datasets.
graphs	logical. Defines if the summary plots and maps are created as pdf files in the working directory (default = TRUE).
areanames	a character string. If the input polygon file is a shape and not a text file, this column indicates the name of the column used for area names. See the SpeciesgeocodeR documentation for more information, if the polygon input file is a text file leave as default (default = "").
occ.thresh	numerical. Threshold for the minimum number of occurrences necessary in a polygon to be counted as present (in percent of the total occurrences of the species). Default $= 0$ .
elevation	logical. Should the results be split by elevation class? (default = FALSE)
threshold	numeric vector. Indicating the elevation thresholds.
verbose	logical. If TRUE, reports the progress of the analyses.
cleaning	logical. If TRUE, GeoClean is applied to the input coordinates.
	Further arguments to be passed to GeoClean.

## **Details**

This performs a complete speciesgeocodeR occurrences to area classification analysis and produces the standard set of output files: 1. classification of all samples to a polygon, 2. Summary of species (identifier) occurrence per polygon, 3. a table of samples that could not be classified to any of the input polygons, 4. a nexus-file, including the species classification, 5. a coexistence matrix, showing to which percentage species to co-occur in the input polygons, 5. a table giving species numbers per polygon. Furthermore produces a set of .pdf files in the output directory: 1. a bar chart showing the number of species per polygon, 2. a bar chart summarizing numbers of each species for each polygon, 3. a bar chart summarizing the occurrences in each polygon per species, 4. a map of all polygons with the points classified to them, colored for species (identifier), 5. a map of the occurrences of all species, a map showing all points used and all unclassified points in the geographic context, 6. a heat plot showing the co-occurrence patterns of all species in the polygons. If elevation = T, the input dataset is split at the elevation(s) provided by threshold, and separate output files are generated for each elevation class.

#### Value

A set of output tables (.txt) and graphics (.pdf).

#### Note

See the speciesgeocodeR documentation for further information and examples.

# **Examples**

```
## Not run:
data(lemurs)
data(mdg_poly)
SpeciesGeoCoder(lemurs, mdg_poly
## End(Not run)
```

Spgc2BioGeoBEARS

Geography Input for BioGeoBEARS

## **Description**

Convert an object of the class spgeoOUT in the right format to be used as geography input for BioGeoBEARS.

## Usage

```
Spgc2BioGeoBEARS(x, phyl = NULL, file = NULL, true.areas = T, true.species = T)
```

## **Arguments**

x an object of the clas 'spgeoOUT'

phyl an object of the class phylo. An optional phylogeny to be used with Bio-

GeoBEARS, to align species.

file character. The complete path and filename to the outputfile.

true.areas logical. If TRUE, areas with 0 species are removed from the matrix. Default =

TRUE.

true.species logical. If TRUE, species not occurring in any area are removed from the matrix.

Default = TRUE.

## **Details**

This will create a file in the working directory.

#### Value

A file in the working directory and a list.

#### Author(s)

Alexander Zizka, Ruud Scharn

SpGeoCod 23

## **Examples**

```
## Not run:
data(lemurs)
data(mdg_poly)

outp <- SpGeoCod(lemurs, mdg_poly)
conv <- Spgc2BioGeoBEARS(outp)

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

SpGeoCod

Point to Polygon Classification from Text Files

# Description

Wrapper around ReadPoints and SpGeoCodH, runs a complete species geocodeR occurrences to area classification analysis from text files as input.

# Usage

# Arguments

x	a character string giving the path to a text file containing the coordinates of points of interest (tab delimited, three columns with header: identifier, XCOOR, YCOOR) OR a data.frame with three columns named: identifier, XCOOR, YCOOR.
У	a character string giving the path to a shape file containing the polygons of interet OR a text file containing the polygons of interest (tab delimited, three columns with header: identifier, XCOOR, YCOOR) OR an object of the class SpatialPolygonsOR an object of the class SpatialPolygonsDataframe.
areanames	a character string. If the input polygon file is a shape and not a text file, this column indicates the name of the column used for area names. See the SpeciesgeocodeR tutorial for more information, if the polygon input file is a text file leave as default (Default = "").
occ.thresh	numerical. Threshold for the minimum number of occurrences necessary in a polygon to be counted as present (in percent of the total occurrences of the species). Default $= 0$ .
elevation	logical. Should the results be split by elevation class? (default = FALSE)
threshold	numeric vector. Indicating the elevation thresholds.
cleaning	logical. If TRUE, GeoClean is applied to the input coordinates.
	Further arguments to be passed to GeoClean.

## Value

An object of the class spgeoOUT.

24 SpGeoCodH

#### Note

See the speciesgeocodeR tutorial for further information and examples.

#### See Also

```
ReadPoints, SpGeoCodH.
```

## **Examples**

```
data(lemurs)
data(mdg_poly)
outp <- SpGeoCod(lemurs, mdg_poly)</pre>
```

SpGeoCodH

A Standard a SpeciesgeocodeR Area Classification

## **Description**

Run a standard speciesgeocodeR occurrence point to area classification analysis on an object of the class spgeoIN (created by ReadPoints). The results are stored in an object of the class spgeoOUT. See details for more information.

# Usage

```
SpGeoCodH(x, areanames = NULL, occ.thresh = 0)
```

## **Arguments**

x an object of the class spgeoIN

areanames a character string. If the polygons in the spgeoIN object were derived from

a shape file, this column indicates the name of the column used for area names. See the Speciesgeocoder tutorial for more information. If the polygons in the spgeoIN object were derived from a text file leave as default (Default = NA).

occ.thresh numerical. The threshold for the minimum number of occurrences necessary

in an area to be counted as present (in percent of the total occurrences of the

species). Default = 0.

#### **Details**

This function uses an object of the class spgeoIN and performs a point in polygon test classifying each species to a polygon, summarizes the information per samples that could not be classified and calculates a coexistence matrix. These objects are then put together with the input information to an object of the class spgeoOUT.

## Value

An object of the class spgeoOUT.

#### Note

See the speciesgeocodeR tutorial for further information and examples.

summary.spgeoIN 25

## **Examples**

```
data(lemurs)
data(mdg_poly)

inp <- ReadPoints(lemurs, mdg_poly)
outp <- SpGeoCodH(inp)
names(outp)</pre>
```

summary.spgeoIN

Summary Method for SpgeoIN

# Description

Summarizes the content of a spgeoIN object.

#### Usage

```
## S3 method for class 'spgeoIN'
summary(object, areanames = NA, ...)
```

# Arguments

object of class spgeoIN

areanames character. Indicating the column with the area names, if not specified, the ID

slot is displayed.

... arguments to be passed to methods, such as graphical parameters (see par).

## **Details**

The summary gives an overview of the number of species, occurrence points and number of polygons in the object, and gives a summary of the input coordinates as well as the polygon names.

## See Also

ReadPoints

```
data(lemurs)
data(mdg_poly)
inp <- ReadPoints(lemurs, mdg_poly)
summary(inp)</pre>
```

26 WriteOut

summary.spgeoOUT

Summary Method for SpgeoOUT

## **Description**

summary method for objects of the class spgeoOUT

#### Usage

```
## S3 method for class 'spgeoOUT'
summary(object, areanames = NA, ...)
```

## **Arguments**

object an object of the class spgeoOUT

areanames a character string. Indicating the column containing the area names, if not spec-

ified the ID slot is displayed.

arguments to be passed to methods, such as graphical parameters (see par).

#### **Details**

The summary gives an overview of the number of species, occurrence points and number of polygons in the object, and gives a summary of the input coordinates as well as the polygon names. Additionally it shows a summary of species diversity per polygon and the number of species not classified to any polygon.

#### See Also

SpGeoCodH

#### **Examples**

```
data(lemurs)
data(mdg_poly)

inp <- ReadPoints(lemurs, mdg_poly)
outp <- SpGeoCodH(inp)
summary(outp)</pre>
```

WriteOut

Write SpeciesgeocodeR results to the Working Directory

## **Description**

This function writes the results of a speciesgeocodeR analysis to the working directory. Output statistics as .txt files, maps and plots as .pdf and a nexus file for use in phylogenetic analyses.

# Usage

WwfLoad 27

## **Arguments**

x an object of the class spgeoOUT

writetype a character string. Defines which results should be written to disk. Default is to

"all", see details for other options

areanames a character string. If the polygons of the spgeoOUT object were derived from

a shape file, this column indicates the name of the column used for area names. See the SpeciesgeocodeR documentation for more information, if the polygons

of the spgeoOUT object were derived from a text file leave as default.

#### **Details**

There are different options for the writetype argument: "all" = all summary tables, plots, maps and a heatplot of species-coexistence if calculated (see CoExClass); "BioGeoBEARS" = a text file in the geography input format of BioGeoBEARS; "coexistence" = the coexistence matrix as table and heatplot "graphs" = summary graphs as pdf files; "maps" = summary maps as .pdf file; "nexus" = write out the species to area classification as nexus file; "statistics" = the summary tables as .txt files.

#### Value

a set of files in the working directory

#### See Also

SpeciesGeoCoder, SpGeoCod, SpGeoCodH

## **Examples**

```
## Not run:
data(lemurs)
data(mdg_poly)

inp <- ReadPoints(lemurs, mdg_poly)
outp <- SpGeoCodH(inp)
WriteOut(outp)
WriteOut(outp, writetype = "graphs")
WriteOut(outp, writetype = "statistics")
WriteOut(outp, writetype = "nexus")
WriteOut(outp, writetype = "coexistence")
## End(Not run)</pre>
```

WwfLoad

Load the WWF Terrestrial Ecoregions

# **Description**

Downloads a shape-file containing the terrestrial ecoregions as defined by Olson et al. 2001 from worldwildlife.org/publications/terrestrial-ecoregions-of-theworld, unpacks them and loads the into R.

28 WwfLoad

# Usage

```
WwfLoad(x = "")
```

#### **Arguments**

Х

the path to the folder where the shapes should be saved. If set to "" the files are saved to the working directory.

#### Value

An object of the class SpatialPolygonsDataframe

#### Note

See the speciesgeocodeR documentation for further information and examples.

#### References

Olson, D. M., Dinerstein, E., Wikramanayake, E. D., Burgess, N. D., Powel, G. V. N., Underwood, E. C., Damico, J. A., Itoua, I., Strand, H. E., Morrison, J. C., Loucks, C. J., Ricketts, T. H., Kura, Y., Lamoreux, J. F., Wettengel, W. W., Hedao, P., and Kassem, K.R. 2001 Terrestrial ecoregions of the world: A new map of life on earth. *BioScience*, **51**(11):933–938.

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

```
## Not run:
wwf_eco <- WwfLoad()
plot(wwf_eco)
names(wwf_eco)
## End(Not run)</pre>
```

# Index

*Topic <b>IO</b>	CoExClass, 4, 27	
GetElevation, 8	countryref, 5, 7	
ReadPoints, 18	0 0] 5 ( 10 11	
SpeciesGeoCoder, 20	GeoClean, 5, 6, 10, 11	
WriteOut, 26	getData, 8	
*Topic datasets	GetElevation, 8	
countryref, 5	IUCNest, 3, 9	
landmass, 10	TOCNEST, 3, 9	
lemurs, 10	landmass, 10	
lemurs_in, 11	lemurs, 10	
lemurs_test, 11	lemurs_in, 11, <i>19</i>	
mdg_biomes, 14	lemurs_test, 11	
mdg_poly, 14	16mar 5_0000, 11	
*Topic datgen	MapGrid, 12, 13, 20	
Spgc2BioGeoBEARS, 22	MapRichness, 13, 20	
*Topic <b>hplot</b>	mdg_biomes, 14	
MapGrid, 12	mdg_poly, 14	
MapRichness, 13		
PlotHull, 17	par, <i>13</i> , <i>16</i>	
*Topic <b>manipl</b>	plot, <i>17</i>	
GeoClean, 6	plot.spgeoIN, 15	
*Topic <b>methods</b>	plot.spgeoOUT, 15	
plot.spgeoIN, 15	PlotHull, 17	
plot.spgeoOUT, 15		
summary.spgeoIN, 25	RangeRichness, <i>13</i> , 18, <i>20</i>	
summary.spgeoOUT, 26	raster, <i>18</i> , <i>20</i>	
*Topic <b>package</b>	ReadPoints, 8, 11, 15, 18, 23–25	
speciesgeocodeR-package, 2	RichnessGrid, <i>12</i> , <i>13</i> , 19	
*Topic <b>spatial</b>		
CalcRange, 3	SpatialPolygons, 19	
CoExClass, 4	SpatialPolygonsDataFrame, 19	
GeoClean, 6	SpeciesGeoCoder, 2, 20, 27	
GetElevation, 8	speciesgeocodeR	
IUCNest, 9	(speciesgeocodeR-package), 2	
MapGrid, 12	speciesgeocodeR-package, 2	
MapRichness, 13	Spgc2BioGeoBEARS, 22	
PlotHull, 17	SpGeoCod, 4, 8, 13, 23, 27	
RangeRichness, 18	SpGeoCodH, 16, 23, 24, 24, 26, 27	
RichnessGrid, 19	summary.spgeoIN, 25	
SpeciesGeoCoder, 20	summary.spgeoOUT, $16$ , $26$	
SpGeoCod, 23	West 20 d 20	
SpGeoCodH, 24	WriteOut, 26	
	WwfLoad, 27	
CalcRange, 3, 9, 17, 18		