Package 'WEGE'

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Type Package

```
Title A Metric to Rank Locations for Biodiversity Conservation
Version 0.1.0
Maintainer Harith Farooq <a href="mailto:harithmorgadinho@gmail.com">harithmorgadinho@gmail.com</a>
Description Calculates the WEGE (Weighted Endemism including Global
      Endangerment index) index for a particular area. Additionally it also
      calculates rasters of KBA's (Key Biodiversity Area) criteria (A1a, A1b, A1e,
      and B1), Weighted endemism (WE), the EDGE (Evolutionarily Distinct and
      Globally Endangered) score, Evolutionary Distinctiveness (ED) and Extinction
      risk (ER). Farooq, H., Azevedo, J., Belluardo F., Nanvonamuquitxo, C.,
      Bennett, D., Moat, J., Soares, A., Faurby, S. & Antonelli, A. (2020)
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Author Harith Farooq [aut, cre] (<a href="https://orcid.org/0000-0001-9031-2785">https://orcid.org/0000-0001-9031-2785</a>),
      Josué Azevedo [aut],
      Francesco Belluardo [aut],
      Cristóvão Nanvonamuquitxo [aut],
      Dom Bennett [aut] (<https://orcid.org/0000-0003-2722-1359>),
      Jason Moat [aut],
      Amadeu Soares [aut],
      Søren Faurby [aut],
      Alexandre Antonelli [aut] (<a href="https://orcid.org/0000-0003-1842-9297">https://orcid.org/0000-0003-1842-9297</a>)
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R topics documented:

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```

Description

A function to get the WEGE index value for a provided polygon.

Usage

```
get_edge(
  target_area,
  input,
  x,
  y,
  species = "binomial",
  ed = "ED",
  category = "category",
  res = 1
)
```

Arguments

target_area	Either a sp or sf object to which to calculate the WEGE index.
input	Species ranges, either from a shapefile or from a georeferenced species list with a column for species, two columns for coordinates and one for the IUCN cate-
	gory.
X	name of the longitude column.
У	name of the latitude column.
species	name of the species column.
ed	name of the evolutionary distinctiveness column.
category	name of IUCN the category column. Terminology must be as follows: DD for Data Deficient, LC for Least Concern, NT for Near Threatened, EN, for Endangered, CR for Critically Endangered, EW for Extinct in the wild and EX for Extinct.
res	grid-cell size to use to calculate the range of the species in case a georeferenced species list was provided.

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Value

a value corresponding to the EDGE score of the provided target_area

Examples

```
library(WEGE)
library(sp)
library(sf)
species <- letters[1:26]</pre>
range_list <- list()</pre>
for (i in seq_along(species)) {
  temp0 <- cbind(runif(3,1,50),runif(3,1,50))
  temp <- Polygon(rbind(temp0, temp0[1,]))</pre>
  range_list[[i]] <- Polygons(list(temp), ID = c(species[i]))</pre>
}
input <- st_as_sf(SpatialPolygons(range_list))</pre>
categories <- c('LC','NT','VU','EN','CR')</pre>
input$binomial <- species</pre>
input$category <- sample(size = nrow(input),x = categories,replace = TRUE)</pre>
input$ED <- runif(nrow(input),1,30)</pre>
temp0 <- cbind(runif(3,1,50),runif(3,1,50))
target_area <- Polygon(rbind(temp0,temp0[1,]))</pre>
target_area <- Polygons(list(target_area), ID = 'Target area')</pre>
target_area <- st_as_sf(SpatialPolygons(list(target_area)))</pre>
get_edge(target_area = target_area,input = input,species = 'binomial',category = 'category')
```

get_kba_criteria

Function to get the KBA criteria

Description

A function to get the WEGE index value for a provided polygon.

Usage

```
get_kba_criteria(
  target_area,
  input,
  x,
  y,
  species = "binomial",
  category = "category",
  res = 1
)
```

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Arguments

target_area Either a sp or sf object to which to calculate the WEGE index. input Species ranges, either from a shapefile or from a georeferenced species list with a column for species, two columns for coordinates and one for the IUCN category. name of the longitude column. Х name of the latitude column. У name of the species column. species category name of IUCN the category column. Terminology must be as follows: DD for Data Deficient, LC for Least Concern, NT for Near Threatened, EN, for Endangered, CR for Critically Endangered, EW for Extinct in the wild and EX for Extinct. grid-cell size to use to calculate the range of the species in case a georeferenced res species list was provided.

Value

a data.frame containing all the species that trigger KBA status as well as the criteria they trigger.

Examples

```
library(WEGE)
library(sp)
library(sf)
species <- letters[1:26]</pre>
range_list <- list()</pre>
for (i in seq_along(species)){
  temp0 <- cbind(runif(3,1,50),runif(3,1,50))</pre>
  temp <- Polygon(rbind(temp0,temp0[1,]))</pre>
  range_list[[i]] <- Polygons(list(temp), ID = c(species[i]))</pre>
input <- st_as_sf(SpatialPolygons(range_list))</pre>
categories <- c('LC','NT','VU','EN','CR')</pre>
input$binomial <- species</pre>
input$category <- sample(size = nrow(input),x = categories,replace = TRUE)</pre>
temp0 <- cbind(runif(3,1,50),runif(3,1,50))
target_area <- Polygon(rbind(temp0,temp0[1,]))</pre>
target_area <- Polygons(list(target_area), ID = 'Target area')</pre>
target_area <- st_as_sf(SpatialPolygons(list(target_area)))</pre>
get_kba_criteria(target_area,input)
```

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Description

A function to get the WEGE index value for a provided polygon.

Usage

```
get_wege(
  target_area,
  input,
  x,
  y,
  species = "binomial",
  category = "category",
  res = 1
)
```

Arguments

target_area	Either a sp or sf object to which to calculate the WEGE index.
input	Species ranges, either from a shapefile or from a georeferenced species list with a column for species, two columns for coordinates and one for the IUCN category.
X	name of the longitude column.
у	name of the latitude column.
species	name of the species column.
category	name of IUCN the category column. Terminology must be as follows: DD for Data Deficient, LC for Least Concern, NT for Near Threatened, EN, for Endangered, CR for Critically Endangered, EW for Extinct in the wild and EX for Extinct.
res	grid-cell size to use to calculate the range of the species in case a georeferenced species list was provided.

Value

a value corresponding to the WEGE index value of the provided target_area

Examples

```
library(sp)
library(sf)
library(WEGE)
species <- letters[1:26]</pre>
```

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```
range_list <- list()
for (i in seq_along(species)){
   temp <- Polygon(cbind(runif(4,1,50),runif(4,1,50)))
   range_list[[i]] <- Polygons(list(temp), ID = c(species[i]))}
input <- st_as_sf(SpatialPolygons(range_list))
categories <- c('LC','NT','VU','EN','CR')
input$binomial <- species
input$category <- sample(size = nrow(input),x = categories,replace = TRUE)

target_area <- Polygon(cbind(runif(4,1,50),runif(4,1,50)))
target_area <- Polygons(list(target_area), ID = 'Target area')
target_area <- st_as_sf(SpatialPolygons(list(target_area)))
get_wege(target_area,input,species = 'binomial',category = 'category')</pre>
```

spat_ras

spat_ras

Description

A function to get the WEGE index value for a provided polygon.

Usage

```
spat_ras(
  target_area,
  input,
  x,
  y,
  species = "binomial",
  category = "category",
  show_progress = FALSE,
  ed,
  res = 1
)
```

Arguments

target_area Either a sp or sf object to which to calculate the WEGE index.

input Species ranges, either from a shapefile or from a georeferenced species list with a column for species, two columns for coordinates and one for the IUCN cate-

gory.

x name of the longitude column.y name of the latitude column.species name of the species column.

category name of IUCN the category column. Terminology must be as follows: DD

for Data Deficient, LC for Least Concern, NT for Near Threatened, EN, for Endangered, CR for Critically Endangered, EW for Extinct in the wild and EX

for Extinct.

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show_progress	Progress of the analysis by showing the number of the grid where the function is calculating the different indices.
ed	name of the evolutionary distinctiveness column.
res	grid-cell size to use to calculate the range of the species in case a georeferenced species list was provided.

Value

A RasterStack with rasters for each KBA criteria (A1a,A1b,A1e,B1) and indices calculated (GE,ED,EDGE,WEGE)

Examples

```
library(WEGE)
library(sp)
library(sf)
library(raster)
 species <- sample(letters, 10)</pre>
 range_list <- list()</pre>
 for (i in seq_along(species)) {
   temp0 <- cbind(runif(3,1,50),runif(3,1,50))
   temp <- Polygon(rbind(temp0,temp0[1,]))</pre>
   range_list[[i]] <- Polygons(list(temp), ID = c(species[i]))</pre>
   }
 input <- st_as_sf(SpatialPolygons(range_list))</pre>
 categories <- c('LC','NT','VU','EN','CR')</pre>
 input$binomial <- species</pre>
 input$category <- sample(size = nrow(input),x = categories,replace = TRUE)</pre>
 input$ed <- runif(runif(10,1,50))</pre>
 temp0 <- cbind(runif(3,1,50),runif(3,1,50))
 target_area <- Polygon(rbind(temp0,temp0[1,]))</pre>
 target_area <- Polygons(list(target_area), ID = 'Target area')</pre>
 target_area <- st_as_sf(SpatialPolygons(list(target_area)))</pre>
 spat_ras(target_area,input,species = 'binomial',ed='ed', res = 1)
```

WEGE

WEGE: A new metric for ranking locations for biodiversity conservation

Description

WEGE is an R package that allows the user to calculate the WEGE index for a particular area. Additionally it also calculates rasters of KBA criteria (A1a, A1b, A1e, and B1) Weighted endemism, the EDGE score, Evolutionary Distinctiveness and Extinction risk.

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