Package ‘intSDM’

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Title Reproducible Integrated Species Distribution Models Across Norway using ‘INLA’

Description Integration of disparate datasets is needed in order to make efficient use of all available data and thereby address the issues currently threatening biodiversity. Data integration is a powerful modeling framework which allows us to combine these datasets together into a single model, yet retain the strengths of each individual dataset. We therefore introduce the package, ‘intSDM’: an R package designed to help ecologists develop a reproducible workflow of integrated species distribution models, using data both provided from the user as well as data obtained freely online. An introduction to data integration methods is discussed in Issac, Jarzyna, Keil, Dambly, Boersch-Supan, Browning, Freeman, Golding, Guillera-Arroita, Henrys, Jarvis, Lahoz-Monfort, Pagel, Pescott, Schmucki, Simmonds and O’Hara (2020) <doi:10.1016/j.tree.2019.08.006>.

Version 1.0.5

Depends R (>= 3.5), inlabru (>= 2.3.1), ggplot2, sp (>= 1.4-5), stats, PointedSDMs, methods

Imports dplyr, geodata, terra, spocc, raster, maps

Suggests fields, RANN, INLA (>= 21.08.31), rgeos, viridis, ggpolyopath, ggmap, spatstat, RColorBrewer, knitr, rmarkdown, testthat (>= 3.0.0), sf, covr

Additional_repositories https://inla.r-inla-download.org/R/testing/

License GPL (>= 3)

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Description

Outputs for structuredData class

PA_redlist A dataset containing the presences and absences of three species: "Fraxinus excelsior", "Ulmus glabra" and "Arnica montana"

Description

A dataset containing the presences and absences of three species: "Fraxinus excelsior", "Ulmus glabra" and "Arnica montana"

References

https://www.gbif.org
show.structuredData-method

Print methods for structuredData object

Description
Print methods for structuredData object

Usage
## S4 method for signature 'structuredData'
show(object)

Arguments

object A "structuredData" object.

Value
A print of the names of the datasets, as well as their lengths.

species_model

species_model: function to construct an integrated species distribution model, as well as other useful outputs from the model.

Description
This function is used to construct integrated species distribution models using data specified with structured_data as well as present only data obtained from the Global Biodiversity Information Facility (GBIF).

Usage

species_model(
  speciesNames,
  date = NULL,
  gbifOptx = list(coordinateUncertaintyInMeters = "0,1000"),
  structuredData = NULL,
  spatialCovariates = NULL,
  worldclimCovariates = NULL,
  res = 0.5,
  scale = FALSE,
  location = "Norway",
  boundary = NULL,
  return = "predictions map",
)
Arguments

speciesNames A vector of species’ names to collect from GBIF.
date Vector of length two denoting the date range to select species from. Defaults to NULL.
gbifOpts A named list of additional options to filter the GBIF records. See ?rgbif::occ_search for the filters which may be applied to the GBIF records. Defaults to list(coordinateUncertaintyInMeters = c(0, 1000)) which specifies a desired range of coordinates uncertainty (in meters) between 0 and 1000.
structuredData Additional datasets to integrate with the presence only GBIF data. See the structured_data function. Defaults to NULL.
spatialCovariates Spatial covariates to include in the model. May be a Raster or Spatial object. Cannot be non-NULL if worldclimCovariates is non-NULL.
worldclimCovariates Names of the covariates to extract from Worldclim. Defaults to NULL; cannot be non-NULL if spatialCovariates is non-NULL.
res Resolution for the WorldClim covariates. Valid values are: 0.5, 2.5, 5, 10. Defaults to 0.5.
scale Should the spatial covariates be scaled. Defaults to FALSE.
location Which area of Norway to model. Defaults to ‘Norway’ which suggests a model for the entire county.
boundary SpatialPolygons object of the study area. If NULL an object may be formed with location.
return Object to return. Has to be one of c('boundary', 'species', 'species plot', 'mesh', 'mesh plot', 'model', 'predictions', 'predictions map').
mesh An inla.mesh object to include in the model. Defaults to NULL.
meshParameters A list of inla.mesh arguments to create a mesh if mesh = NULL.
spdeModel inla.spde model used in the model. May be a named list where the name of the spde object is the name of the associated dataset. Default NULL uses inla.spde2.matern.
biasField Should a second (bias) random field be added to the GBIF data. Defaults to FALSE.
species_model

biasModel inla.spde model used for the bias field. Requires biasModel to be TRUE. Default NULL uses inla.spde2.matern.

projection CRS projection to use. Defaults to CRS('?proj=utm +zone=32 +ellps=WGS84 +datum=WGS84 +units=m +no_defs').

limit Set the number of species downloaded. Defaults to 10000.

options A list of INLA and inlabru options. Defaults to NULL.

Additional arguments used in PointedSDMs's intModel function.

Value

The return of the function is determined by the argument return. For the different values of return:

- boundary A SpatialPolygons of the boundary used in the model.
- species A data.frame object of the species' coordinates used in the model.
- species plot A ggplot plot of the species across a map.
- mesh an inla.mesh object.
- mesh plot a plot of the inla.mesh object.
- model the integrated model.
- predictions predictions from the integrated model (on the linear scale).
- predictions map A ggplot plot of the predictions across the boundary.

Examples

```r
## Not run:
if(requireNamespace('INLA')) {

#Objects required for example
data("PA_redlist")
speciesNames <- c('Fraxinus excelsior', 'Ulmus glabra')

#Set up structured dataset
dataObj <- structured_data(PA_redlist, datasetType = c('PA'), responsePA = 'individualCount',
 speciesName = 'species',
 coordinateNames = c("longitude", "latitude" ))

#Get species map
predictions <- species_model(return = 'species plot',
 boundary = boundary, speciesNames = species,
 limit = 10, structuredData = dataObj,
 meshParameters = list(cutoff=0.08, max.edge=c(1, 3), offset=c(1,1)),
 worldclimCovariates = 'Annual Mean Temperature')
}
```
### Description

**bru_sdm_data class**

**Value**

An S4 object with three slots, which contain lists of the data for the three observation models allowed in the modelling framework.

**Slots**

- `dataPO` A list of the present only datasets.
- `dataPA` A list of the present absence datasets.
- `dataCount` A list of the count datasets.

### Description

**structured_data: function to organize structured datasets.**

The function is used to convert a collection of structured datasets into an object which can be used in `speciesModel`. The three types of structured data allowed by this function are present absence (PA), present only (PO) and counts/abundance datasets, which are controlled using the `datasetType` argument. The other arguments of this function are used to specify the appropriate variable (such as response name, trial name, species name and coordinate name) names in these datasets.

**Usage**

```r
structured_data(
  ...,  
  datasetType = NULL,  
  responsePA = NULL,  
  trialsPA = NULL,  
  responseCount = NULL,  
  speciesName = "species",  
  coordinateNames = c("longitude", "latitude")
)
```
**Arguments**

... The datasets used in the model. May be either datasets or a SpatialPointsDataFrames.

datasetType A vector which gives the type of dataset. Must be either 'count', 'PO' or 'PA'. Defaults to NULL.

responsePA Name of the response variable in the PA datasets. Defaults to NULL.

trialsPA Name of the trial name variable in the PA datasets. Defaults to NULL.

responseCount Name of the response variable in the count datasets. Defaults to NULL.

speciesName Name of the species variable name. Defaults to 'species'.

coordinateNames Name of the coordinates used in the model. Defaults to c('longitude', 'latitude').

**Value**

An S4 object of class structuredData, which contains three slots for the data of each observation model allowed in the framework.

**Examples**

```{
#Objects required for example
data("PA_redlist")
speciesNames <- c("Fraxinus excelsior", "Ulmus glabra")

#Set up structured dataset
dataObj <- structured_data(PA_redlist, datasetType = c("PA"), responsePA = "individualCount", speciesName = "species",
coordinateNames = c("longitude", "latitude" ))
```
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