

Package ‘caretSDM’

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Type Package

Title Build Species Distribution Modeling using 'caret'

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Description Use machine learning algorithms and advanced geographic information system tools to build Species Distribution Modeling in a extensible and modern fashion.

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BugReports <https://github.com/luizesser/caretSDM/issues>

Imports blockCV, caret, checkmate, cli, CoordinateCleaner, data.table, dismo, dplyr, ecospat, fs, furrr, future, ggplot2, ggspatial, glue, gtools, httr2, lwgeom, mapview, maxnet, methods, parallelly, pdp, pROC, progressr, purrr, raster, rgbif, Rtsne, sf, stars, stats, stringdist, stringr, terra, tidyr, usdm, utils

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Contents

add_predictors	3
add_scenarios	4
algorithms	6
bioc	6
buffer_sdm	7
correlate_sdm	8
data_clean	9
GBIF_data	11
gcms_ensembles	12
input_sdm	13
is_input_sdm	15
join_area	16
occ	17
occurrences_sdm	17
parana	19
pca_predictors	19
pdp_sdm	20
plot_occurrences	22
prediction_change_sdm	24
predictors	25
predict_sdm	27
print.input_sdm	29
print.models	30
print.occurrences	30
print.predictions	31
pseudoabsences	31
rivs	33
salm	34
scen	34
scen_rs	35
sdm_area	36
sdm_as_stars	37
select_predictors	39
summary_sdm	40
train_sdm	41
tsne_sdm	44

add_predictors 3

tuneGrid_sdm	45
use_esm	46
use_mem	47
varImp_sdm	48
vif_predictors	49
WorldClim_data	51
write_ensembles	52

Index 54

<code>add_predictors</code>	<i>Add predictors to sdm_area</i>
-----------------------------	-----------------------------------

Description

This function includes new predictors to the `sdm_area` object.

Usage

```
add_predictors(sa, pred, variables_selected = NULL, gdal = TRUE)
```

```
get_predictors(i)
```

Arguments

<code>sa</code>	A <code>sdm_area</code> object.
<code>pred</code>	RasterStack, SpatRaster, stars or sf object with predictors data.
<code>variables_selected</code>	character vector with variables names in <code>pred</code> to be used as predictors. If NULL adds all variables.
<code>gdal</code>	Boolean. Force the use or not of GDAL when available. See details.
<code>i</code>	<code>input_sdm</code> or <code>sdm_area</code> object to retrieve data from.

Details

`add_predictors` returns a `sdm_area` object with a grid built upon the `x` parameter. There are two ways to make the grid and resample the variables in `sdm_area`: with and without `gdal`. As standard, if `gdal` is available in you machine it will be used (`gdal = TRUE`), otherwise `sf/stars` will be used.

Value

For `add_predictors` the same input `sdm_area` object is returned including the `pred` data binded to the previous grid. `get_predictors` retrieves the grid from the `i` object.

Author(s)

Luíz Fernando Esser (luizesser@gmail.com) and Reginaldo Ré. <https://luizfesser.wordpress.com>

See Also

[sdm_area predictors bioc](#)

Examples

```
# Create sdm_area object:
sa <- sdm_area(parana, cell_size = 25000, crs = 6933)

# Include predictors:
sa <- add_predictors(sa, bioc)

# Retrieve predictors data:
get_predictors(sa)
```

add_scenarios	<i>Add scenarios to sdm_area</i>
---------------	----------------------------------

Description

This function includes scenarios in the `sdm_area` object.

Usage

```
add_scenarios(sa, scen = NULL, scenarios_names = NULL, pred_as_scen = TRUE,
              variables_selected = NULL, stationary = NULL, crop_area = NULL)

set_scenarios_names(i, scenarios_names = NULL)

scenarios_names(i)

get_scenarios_data(i)

select_scenarios(i, scenarios_names = NULL)
```

Arguments

<code>sa</code>	A <code>sdm_area</code> or <code>input_sdm</code> object.
<code>scen</code>	RasterStack, SpatRaster or stars object. If NULL adds predictors as a scenario.
<code>scenarios_names</code>	Character vector with names of scenarios.
<code>pred_as_scen</code>	Logical. If TRUE adds the current predictors as a scenario.
<code>variables_selected</code>	Character vector with variables names in <code>scen</code> to be used as variables. If NULL adds all variables.

stationary	Names of variables from sa that should be used in scenarios as stationary variables.
crop_area	A sf object to crop the scen object if necessary.
i	A sdm_area or input_sdm object.

Details

The function `add_scenarios` adds scenarios to the `sdm_area` or `input_sdm` object. If `scen` has variables that are not present as predictors the function will use only variables present in both objects. `stationary` variables are those that don't change through the scenarios. It is useful for hidrological variables in fish habitat modeling, for example (see examples below). When adding multiple scenarios in multiple runs, the function will always add a new "current" scenario. To avoid that, set `pred_as_scen = FALSE`.

Value

`add_scenarios` returns the input `sdm_area` or `input_sdm` object with a new slot called `scenarios` with scen data as a list, where each slot of the list holds a scenario and each scenario is a `sf` object. `set_scenarios_names` sets new names for scenarios in `sdm_area/input_sdm` object. `scenarios_names` returns scenarios' names. `get_scenarios_data` retrieves scenarios data as a list of `sf` objects. `select_scenarios` selects scenarios from `sdm_area/input_sdm` object.

Author(s)

Luíz Fernando Esser (luizesser@gmail.com) <https://luizfesser.wordpress.com>

See Also

[sdm_area](#) [input_sdm](#)

Examples

```
# Create sdm_area object:
sa <- sdm_area(parana, cell_size = 100000, crs = 6933)

# Include predictors:
sa <- add_predictors(sa, bioc)

# Include scenarios:
sa <- add_scenarios(sa, scen[1:2]) |> select_predictors(c("bio1", "bio12"))

# Set scenarios names:
sa <- set_scenarios_names(sa, scenarios_names = c("future_1", "future_2",
                                                "current"))

scenarios_names(sa)

# Get scenarios data:
scenarios_grid <- get_scenarios_data(sa)
scenarios_grid

# Select scenarios:
```

```
sa <- select_scenarios(sa, scenarios_names = c("future_1"))

# Setting stationary variables in scenarios:
sa <- sdm_area(rivs[c(1:200)],, cell_size = 100000, crs = 6933, lines_as_sdm_area = TRUE) |>
  add_predictors(bioc) |>
  add_scenarios(scen, stationary = c("LENGTH_KM", "DIST_DN_KM"))
```

algorithms

Caret Algorithms

Description

A `data.frame` with characteristics of each algorithm available in `caretSDM`. Each column is a different characteristic. This can be helpful for more experienced modelers select algorithms. See the source for a selection method using this data.

Usage

```
algorithms
```

Format

'algorithms' A `data.frame` with 230 rows and 60 columns:

X Algorithms names

Further columns Algorithms attributes

Source

<<https://topepo.github.io/caret/models-clustered-by-tag-similarity.html>>

bioc

Bioclimatic Variables

Description

A `stars` object with bioclimatic variables (bio1, bio4 and bio12) for the Parana state in Brazil. Data obtained from WorldClim 2.1 at 10 arc-min resolution.

Usage

```
bioc
```

Format

'bioc' A starts with 1 attribute and 3 bands:

bio1 Annual Mean Temperature

bio4 Temperature Seasonality

bio12 Annual Precipitation

Source

<<https://www.worldclim.org/>>

buffer_sdm	Create buffer around occurrences
------------	----------------------------------

Description

Create buffer around records in occ_data to be used as study area

Usage

```
buffer_sdm(occ_data, size = NULL, crs = NULL, mcp = FALSE)
```

Arguments

occ_data	A data.frame object with species, decimalLongitude and decimalLatitude columns. Usually the output from GBIF_data.
size	numeric. The distance between the record and the margin of the buffer (i.e. buffer radius).
crs	numeric. Indicates which EPSG it the occ_data in.
mcp	boolean. Should the buffer be applied in each record (FALSE) or in a minimum convex polygon/convex hull (TRUE)? Standard is FALSE.

Value

A sf buffer around occ_data records.

Author(s)

Luíz Fernando Esser (luizesser@gmail.com) <https://luizfesser.wordpress.com>

See Also

[GBIF_data](#)

Examples

```
# Create sdm_area object:
study_area <- buffer_sdm(occ, size=50000, crs=6933)
plot(study_area)
```

correlate_sdm	<i>Correlation between projections</i>
---------------	--

Description

This function aims to unveil the correlation of different algorithms outputs. For that, it uses the predictions on current scenario, but other scenarios can be tested.

Usage

```
correlate_sdm(i, scenario = "current")
```

Arguments

i	A input_sdm object containing predictions.
scenario	A character containing scenario to be tested. Standard is "current". Value must match scenarios_names(i).

Value

A data.frame with pearson correlation between projections.

Author(s)

Luíz Fernando Esser (luizesser@gmail.com) <https://luizfesser.wordpress.com>

Examples

```
# Create sdm_area object:
set.seed(1)
sa <- sdm_area(parana, cell_size = 100000, crs = 6933)

# Include predictors:
sa <- add_predictors(sa, bioc) |> select_predictors(c("bio1", "bio12"))

# Include scenarios:
sa <- add_scenarios(sa)

# Create occurrences:
oc <- occurrences_sdm(occ, crs = 6933) |> join_area(sa)

# Create input_sdm:
```



```
i <- input_sdm(oc, sa)

# Pseudoabsence generation:
i <- pseudoabsences(i, method="random", n_set=2)

# Custom trainControl:
ctrl_sdm <- caret::trainControl(method = "boot",
                                number = 1,
                                repeats = 1,
                                classProbs = TRUE,
                                returnResamp = "all",
                                summaryFunction = summary_sdm,
                                savePredictions = "all")

# Train models:
i <- train_sdm(i, algo = c("naive_bayes"), ctrl=ctrl_sdm) |>
  suppressWarnings()

# Predict models:
i <- predict_sdm(i, th = 0.8)

# Check correlations:
correlate_sdm(i)
```

pred	A sdm_area object. If occ is a input_sdm object with predictors data, than pred is obtained from it.
species	A character stating the name of the column with species names in occ (see details).
lon	A character stating the name of the column with longitude in occ (see details).
lat	A character stating the name of the column with latitude in occ (see details).
capitals	Boolean to turn on/off the exclusion from countries capitals coordinates (see ?cc_cap)
centroids	Boolean to turn on/off the exclusion from countries centroids coordinates (see ?cc_cen)
uplicated	Boolean to turn on/off the exclusion from duplicated records (see ?cc_dupl)
identical	Boolean to turn on/off the exclusion from records with identical lat/long values (see ?cc_equ)
institutions	Boolean to turn on/off the exclusion from biodiversity institutions coordinates (see ?cc_inst)
invalid	Boolean to turn on/off the exclusion from invalid coordinates (see ?cc_val)
terrestrial	Boolean to turn on/off the exclusion from coordinates falling on sea (see ?cc_sea)
independent_test	Boolean. If occ has independent test data, the data cleaning routine is also applied on it.

Details

If the user does not used GBIF_data function to obtain species records, the function may have problems to find which column from the presences table has species, longitude and latitude information. In this regard, we implemented the parameters species, lon and lat so the use can explicitly inform which columns should be used. If they remain as NA (standard) the function will try to guess which columns are the correct one.

Value

A occurrences_sdm object or input_sdm with cleaned presence data.

Author(s)

Luiz Fernando Esser (luizesser@gmail.com) <https://luizfesser.wordpress.com>

See Also

[GBIF_data](#) [occurrences_sdm](#) [sdm_area](#) [input_sdm](#) [predictors](#)

Examples

```
# Create sdm_area object:
sa <- sdm_area(parana, cell_size = 50000, crs = 6933)

# Include predictors:
sa <- add_predictors(sa, bioc) |> select_predictors(c("bio1", "bio12"))

# Create occurrences:
oc <- occurrences_sdm(occ, crs = 6933) |> join_area(sa)

# Create input_sdm:
i <- input_sdm(oc, sa)

# Clean coordinates (terrestrial is set to false to make the run quicker):
i <- data_clean(i, terrestrial = FALSE)
```

GBIF_data

*Retrieve Species data from GBIF***Description**

This function is a wrapper to get records from GBIF using `rgbif` and return a `data.frame` ready to be used in `caretSDM`.

Usage

```
GBIF_data(s, file = NULL, as_df = FALSE, ...)
```

Arguments

<code>s</code>	character vector of species names.
<code>file</code>	character with file to save the output. If not informed, data will not be saved on folder.
<code>as_df</code>	Should the output be a dataframe? Default is FALSE, returning a occurrences object.
<code>...</code>	Arguments to pass on <code>rgbif::occ_data()</code> .

Value

A `data.frame` with species occurrences data, or an occurrences object if `as_df = FALSE`.

Author(s)

Luíz Fernando Esser (luizesser@gmail.com) <https://luizfesser.wordpress.com>

References

<https://www.gbif.org>

Examples

```
# Select species names:
s <- c("Araucaria angustifolia", "Salminus brasiliensis")

# Run function:
oc <- GBIF_data(s)
```

gcms_ensembles

Ensemble GCMs into one scenario

Description

An ensembling method to group different GCMs into one SSP scenario

Usage

```
gcms_ensembles(i, gcms = NULL)
```

Arguments

i	A input_sdm object.
gcms	GCM codes in scenarios_names(i) to group scenarios.

Value

A input_sdm object with grouped GCMs.

Author(s)

Luiz Fernando Esser (luizesser@gmail.com) <https://luizfesser.wordpress.com>

See Also

[GBIF_data](#) [occurrences_sdm](#) [sdm_area](#) [input_sdm](#) [predictors](#)

Examples

```

# Create sdm_area object:
set.seed(1)
sa <- sdm_area(parana, cell_size = 100000, crs = 6933)

# Include predictors:
sa <- add_predictors(sa, bioc)

# Include scenarios:
sa <- add_scenarios(sa, scen) |> select_predictors(c("bio1", "bio12"))

# Create occurrences:
oc <- occurrences_sdm(occ, crs = 6933) |> join_area(sa)

# Create input_sdm:
i <- input_sdm(oc, sa)

# Pseudoabsence generation:
i <- pseudoabsences(i, method="random", n_set = 2)

# Custom trainControl:
ctrl_sdm <- caret::trainControl(method = "boot",
                                number = 1,
                                classProbs = TRUE,
                                returnResamp = "all",
                                summaryFunction = summary_sdm,
                                savePredictions = "all")

# Train models:
i <- train_sdm(i,
              algo = c("naive_bayes"),
              ctrl=ctrl_sdm,
              variables_selected = c("bio1", "bio12")) |>
  suppressWarnings()

# Predict models:
i <- predict_sdm(i, th=0.8)

# Ensemble GCMs:
i <- gcms_ensembles(i, gcms = c("ca", "mi"))
i

```

input_sdm

input_sdm

Description

This function creates a new input_sdm object.

Usage

```
input_sdm(...)

add_input_sdm(i1, i2)
```

Arguments

...	Data to be used in SDMs. Can be a occurrences and/or a sdm_area object.
i1	A input_sdm object.
i2	A input_sdm object.

Details

If sdm_area is used, it can include predictors and scenarios. In this case, input_sdm will detect and include as scenarios and predictors in the input_sdm output. Objects can be included in any order, since the function will work by detecting their classes. The returned object is used throughout the whole workflow to apply functions.

Value

A input_sdm object containing:

grid	sf with POLYGON geometry representing the grid for the study area or LINESTRING if sdm_area was built with a LINESTRING sf.
bbox	Four corners for the bounding box (class bbox): minimum value of X, minimum value of Y, maximum value of X, maximum value of Y
cell_size	numeric information regarding the size of the cell used to rescale variables to the study area, representing also the cell size in the grid.
epsg	character information about the EPSG used in all slots from sdm_area.
predictors	character vector with predictors names included in sdm_area.

Author(s)

Luiz Fernando Esser (luizesser@gmail.com) <https://luizfesser.wordpress.com>

See Also

[occurrences_sdm](#) [sdm_area](#)

Examples

```
# Create sdm_area object:
sa <- sdm_area(parana, cell_size = 50000, crs = 6933)

# Include predictors:
sa <- add_predictors(sa, bioc) |> select_predictors(c("bio1", "bio4", "bio12"))

# Include scenarios:
```

```
sa <- add_scenarios(sa, scen)

# Create occurrences:
oc <- occurrences_sdm(occ, crs = 6933) |> join_area(sa)

# Create input_sdm:
i <- input_sdm(oc, sa)
```

is_input_sdm	is_class <i>functions to check caretSDM data classes.</i>
--------------	---

Description

This functions returns a boolean to check caretSDM object classes.

Usage

```
is_input_sdm(x)

is_sdm_area(x)

is_occurrences(x)

is_models(x)

is_predictions(x)
```

Arguments

x Object to be tested.

Value

Boolean.

Author(s)

Luíz Fernando Esser (luizesser@gmail.com) <https://luizfesser.wordpress.com>

Examples

```
# Create sdm_area object:
sa <- sdm_area(parana, cell_size = 25000, crs = 6933)

is_sdm_area(sa)

is_input_sdm(sa)
```

join_area

Join Area

Description

Join cell_id data from sdm_area to a occurrences

Usage

```
join_area(occ, pred)
```

Arguments

occ	A occurrences object or input_sdm.
pred	A sdm_area object to retrieve cell_id from.

Details

This function is key in this SDM workflow. It attaches cell_id values to occ, deletes records outside pred and allows the use of pseudoabsences. This function also tests if CRS from both occ and pred are equal, otherwise the CRS of pred is used to convert occ.

Value

A occurrences object with cell_id to each record.

Author(s)

Luíz Fernando Esser (luizesser@gmail.com) <https://luizfesser.wordpress.com>

See Also

[occurrences_sdm](#) [sdm_area](#) [input_sdm](#) [pseudoabsences](#)

Examples

```
# Create sdm_area object:
sa <- sdm_area(parana, cell_size = 50000, crs = 6933)

# Include predictors:
sa <- add_predictors(sa, bioc) |> select_predictors(c("bio1", "bio4", "bio12"))

# Include scenarios:
sa <- add_scenarios(sa, scen)

# Create occurrences:
oc <- occurrences_sdm(occ, crs = 6933) |> join_area(sa)
```

occ	<i>Araucaria angustifolia</i> occurrence data
-----	---

Description

A data.frame object with *Araucaria angustifolia* occurrence data obtained from GBIF and filtered with Parana state sf.

Usage

```
occ
```

Format

'occ' A data.frame with 420 rows and 3 columns (EPSG:6933):

species Species name

decimalLongitude Longitude in meters

decimalLatitude Latitude in meters

Source

<<https://www.gbif.org>>

occurrences_sdm	<i>Occurrences Managing</i>
-----------------	-----------------------------

Description

This function creates and manage occurrences objects.

Usage

```
occurrences_sdm(x,
  independent_test = NULL,
  p = 0.1,
  crs = NULL,
  independent_test_crs = NULL,
  ...)
```

```
n_records(i)
```

```
species_names(i)
```

```
get_coords(i)
```

```
occurrences_as_df(i)
```

```
add_occurrences(oc1, oc2)
```

Arguments

<code>x</code>	A <code>data.frame</code> , <code>tibble</code> or <code>sf</code> with species records.
<code>independent_test</code>	Boolean. If <code>independent_test</code> is <code>TRUE</code> , a fraction of the data is kept for independent testing. Otherwise, the whole dataset <code>x</code> is used. It can also be a <code>data.frame</code> or a <code>sf</code> , with species records to be used as independent test. Structure and names should be identical to those in <code>x</code> .
<code>p</code>	Numeric. Fraction of data to be used as independent test. Standard is 0.1.
<code>crs</code>	Numeric. CRS of <code>x</code> .
<code>independent_test_crs</code>	Numeric. CRS of <code>independent_test</code> if it is a <code>data.frame</code> .
<code>...</code>	A vector with column names addressing the columns with species names, longitude and latitude, respectively, in <code>x</code> .
<code>i</code>	<code>input_sdm</code> or <code>occurrences</code> object.
<code>oc1</code>	A <code>occurrences</code> object to be summed with.
<code>oc2</code>	A <code>occurrences</code> object to be summed with.

Details

`x` must have three columns: `species`, `decimalLongitude` and `decimalLatitude`. When `sf` it is only necessary a `species` column. `n_records` return the number of presence records to each species. `species_names` return the species names. `get_coords` return a `data.frame` with coordinates of species records. `add_occurrences` return a `occurrences`. This function sums two `occurrences` objects. It can also sum a `occurrences` object with a `data.frame` object. `occurrences_as_df` returns a `data.frame` with species names and coordinates.

Value

A `occurrences` object.

Author(s)

Luíz Fernando Esser (luizesser@gmail.com) <https://luizfesser.wordpress.com>

See Also

[input_sdm GBIF_data occ](#)

Examples

```
# Create occurrences:
oc <- occurrences_sdm(occ, crs = 6933)
```

parana	<i>Paraná State</i>
--------	---------------------

Description

A sf object with a polygon for the Paraná state in Brazil. This is a subset of the brazilian map provided by official government agency (IBGE)

Usage

```
parana
```

Format

'parana' A sf with 1 row and 5 columns:

GID0 State code

CODIGOIB1 State's phone code

NOMEUF2 Name of the state

SIGLAUF3 Abbreviation of the state's name

geom Geometry column of the sf

Source

<<https://www.ibge.gov.br/geociencias/cartas-e-mapas/bases-cartograficas-continuas/15759-brasil.html>>

pca_predictors	<i>Predictors as PCA-axes</i>
----------------	-------------------------------

Description

Transform predictors data into PCA-axes.

Usage

```
pca_predictors(i, cumulative_proportion = 0.99)
```

```
pca_summary(i)
```

```
get_pca_model(i)
```

Arguments

i A input_sdm object.

cumulative_proportion

A numeric with the threshold for cumulative proportion. Standard is 0.99, meaning that axes returned as predictors sum up more than 99 variance.

Details

`pca_predictors` Transform predictors data into PCA-axes. If the user wants to use PCA-axes as future scenarios, then scenarios should be added after the PCA transformation (see examples).
`pca_summary` Returns the summary of `prcomp` function. See `?stats::prcomp`.
`get_pca_model` Returns the model built to calculate PCA-axes.

Value

`input_sdm` object with variables from both predictors and scenarios transformed in PCA-axes.

Author(s)

Luíz Fernando Esser (luizesser@gmail.com) <https://luizfesser.wordpress.com>

See Also

[vif_predictors](#) [sdm_area](#) [add_scenarios](#) [add_predictors](#)

Examples

```
# Create sdm_area object:
sa <- sdm_area(parana, cell_size = 50000, crs = 6933)

# Include predictors:
sa <- add_predictors(sa, bioc) |> select_predictors(c("bio1", "bio12"))

# Create occurrences:
oc <- occurrences_sdm(occ, crs = 6933) |> join_area(sa)

# Create input_sdm:
i <- input_sdm(oc, sa)

# PCA transformation:
i <- pca_predictors(i)
```

pdp_sdm

Model Response to Variables

Description

Obtain the Partial Dependence Plots (PDP) to each variable.

Usage

```
pdp_sdm(i, spp = NULL, algo = NULL, variables_selected = NULL, mean.only = FALSE)

get_pdp_sdm(i, spp = NULL, algo = NULL, variables_selected = NULL)
```

Arguments

<code>i</code>	A <code>input_sdm</code> object.
<code>spp</code>	A character vector with species names to obtain the PDPs. If NULL (standard), the first species in <code>species_names(i)</code> is used.
<code>algo</code>	A character containing the algorithm to obtain the PDP. If NULL (standard) all algorithms are mixed.
<code>variables_selected</code>	A character. If there is a subset of predictors that should be plotted in this, it can be informed using this parameter.
<code>mean.only</code>	Boolean. Should only the mean curve be plotted or a curve to each run should be included? Standard is FALSE.

Value

A plot (for `pdp_sdm`) or a `data.frame` (for `get_pdp_sdm`) with PDP values.

Author(s)

Luíz Fernando Esser (luizesser@gmail.com) <https://luizfesser.wordpress.com>

See Also

[varImp_sdm](#)

Examples

```
# Create sdm_area object:
sa <- sdm_area(parana, cell_size = 100000, crs = 6933)

# Include predictors:
sa <- add_predictors(sa, bioc) |> select_predictors(c("bio1", "bio12"))

# Include scenarios:
sa <- add_scenarios(sa)

# Create occurrences:
oc <- occurrences_sdm(occ, crs = 6933) |> join_area(sa)

# Create input_sdm:
i <- input_sdm(oc, sa)

# Pseudoabsence generation:
i <- pseudoabsences(i, method="random", n_set=3)

# Custom trainControl:
ctrl_sdm <- caret::trainControl(method = "repeatedcv",
                                number = 2,
                                repeats = 1,
                                classProbs = TRUE,
                                returnResamp = "all",
```

```

summaryFunction = summary_sdm,
savePredictions = "all")

# Train models:
i <- train_sdm(i, algo = c("naive_bayes"), ctrl=ctrl_sdm)
# PDP plots:
pdp_sdm(i)
get_pdp_sdm(i)

```

plot_occurrences	<i>S3 Methods for plot and mapview</i>
------------------	--

Description

This function creates different plots depending on the input.

Usage

```

plot_occurrences(i, spp_name = NULL, pa = TRUE, pa_id = 1)

plot_grid(i)

plot_predictors(i, variables_selected = NULL)

plot_scenarios(i, variables_selected = NULL, scenario = NULL)

plot_predictions(
  i,
  spp_name = NULL,
  scenario = NULL,
  id = NULL,
  ensemble = TRUE,
  ensemble_type = "mean_occ_prob"
)

mapview_grid(i)

mapview_occurrences(i, spp_name = NULL, pa = TRUE)

mapview_predictors(i, variables_selected = NULL)

mapview_scenarios(i, variables_selected = NULL, scenario = NULL)

mapview_predictions(
  i,
  spp_name = NULL,
  scenario = NULL,
  id = NULL,

```

```

    ensemble = TRUE,
    ensemble_type = "mean_occ_prob"
  )

plot_background(i, variables_selected = NULL)

plot_niche(
  i,
  spp_name = NULL,
  variables_selected = NULL,
  scenario = NULL,
  id = NULL,
  ensemble = TRUE,
  ensemble_type = "mean_occ_prob",
  raster = FALSE
)

```

Arguments

<code>i</code>	Object to be plotted. Can be a <code>input_sdm</code> , but also occurrences or <code>sdm_area</code> .
<code>spp_name</code>	A character with species to be plotted. If <code>NULL</code> , the first species is plotted.
<code>pa</code>	Boolean. Should pseudoabsences be plotted together? (not implemented yet.)
<code>pa_id</code>	The id of pseudoabsences to be plotted (only used when <code>pa = TRUE</code>). Possible values are numeric values from 1 to number of PA sets.
<code>variables_selected</code>	A character vector with names of variables to be plotted.
<code>scenario</code>	description
<code>id</code>	The id of models to be plotted (only used when <code>ensemble = FALSE</code>). Possible values are row names of <code>get_validation_metrics(i)</code> .
<code>ensemble</code>	Boolean. Should the ensemble be plotted (<code>TRUE</code>)? Otherwise a prediction will be plotted
<code>ensemble_type</code>	Character of the type of ensemble to be plotted. One of: "mean_occ_prob", "wmean_AUC" or "committee_avg"
<code>raster</code>	Should the niche be extrapolated to a raster covering all possible values in the environmental space?

Details

We implemented a bestiary of plots to help visualizing the process and results. If you are not familiar with `mapview`, consider using it to better visualize maps.

Value

The plot or `mapview` desired.

Author(s)

Luíz Fernando Esser (luizesser@gmail.com) <https://luizfesser.wordpress.com>

See Also

[WorldClim_data](#)

prediction_change_sdm *Prediction Change Analysis*

Description

Provides an automate way for the visualization of projections gain, loss, and stability between different scenarios.

Usage

```
prediction_change_sdm(i, scenario = NULL, ensemble_type = NULL, species = NULL, th = 0.5)
```

Arguments

i	A input_sdm object with projections.
scenario	Character. One of the scenarios that were projected. Can be ensembles as well.
ensemble_type	Character. Type of ensemble to be used. Standard is NULL, but will return the mean_occ_prob
species	Character. Species to be analyzed. Standard is NULL.
th	Numeric. Threshold to binarize the ensemble.

Value

A plot with comparison between current and other scenario.

Author(s)

Luiz Fernando Esser (luizesser@gmail.com) <https://luizfesser.wordpress.com>

See Also

[species_names](#) [scenarios_names](#)

Examples

```
# Create sdm_area object:
set.seed(1)
sa <- sdm_area(parana, cell_size = 100000, crs = 6933)

# Include predictors:
sa <- add_predictors(sa, bioc)

# Include scenarios:
sa <- add_scenarios(sa, scen) |> select_predictors(c("bio1", "bio12"))
```



```

# Create occurrences:
oc <- occurrences_sdm(occ, crs = 6933) |> join_area(sa)

# Create input_sdm:
i <- input_sdm(oc, sa)

# Pseudoabsence generation:
i <- pseudoabsences(i, method="random", n_set = 2)

# Custom trainControl:
ctrl_sdm <- caret::trainControl(method = "boot",
                                number = 1,
                                classProbs = TRUE,
                                returnResamp = "all",
                                summaryFunction = summary_sdm,
                                savePredictions = "all")

# Train models:
i <- train_sdm(i,
              algo = c("naive_bayes"),
              ctrl=ctrl_sdm,
              variables_selected = c("bio1", "bio12")) |>
  suppressWarnings()

# Predict models:
i <- predict_sdm(i, th=0.8)

# Ensemble GCMs:
i <- gcms_ensembles(i, gcms = c("ca", "mi"))
i

# Change Analysis
prediction_change_sdm(i, scenario = "_ssp585_2090", ensemble_type = "mean_occ_prob")

```

predictors

Predictors Names Managing

Description

This function manage predictors names in `sdm_area` objects.

Usage

```

predictors(x)

## S3 method for class 'sdm_area'
predictors(x)

```

```

## S3 method for class 'input_sdm'
predictors(x)

set_predictor_names(x, new_names)

## S3 method for class 'input_sdm'
set_predictor_names(x, new_names)

## S3 method for class 'sdm_area'
set_predictor_names(x, new_names)

get_predictor_names(x)

## S3 method for class 'sdm_area'
get_predictor_names(x)

## S3 method for class 'input_sdm'
get_predictor_names(x)

test_variables_names(sa, scen)

set_variables_names(s1 = NULL, s2 = NULL, new_names = NULL)

```

Arguments

x	A sdm_area or input_sdm object to get/set predictors names.
new_names	A character vector from size length(get_predictor_names(x))
sa	A sdm_area object.
scen	A stars object with scenarios.
s1	A stars object with scenarios.
s2	A stars object with scenarios or a sdm_area object.

Details

This functions is available so users can modify predictors names to better represent them. Use carefully to avoid giving wrong names to the predictors. Useful to make sure the predictors names are equal the names in scenarios. `test_variables_names` Tests if variables in a stars object (scen argument) matches the given sdm_area object (sa argument). `set_variables_names` will set s1 object variables names as the s2 object variables names OR assign new names to it.

Value

`predictors` and `get_predictor_names` return a character vector with predictors names. `test_variables_names` returns a logical informing if all variables are equal in both objects (TRUE) or not (FALSE). `set_variables_names` returns the s1 object with new names provided by s2 or new_names.

Author(s)

Luíz Fernando Esser (luizesser@gmail.com) <https://luizfesser.wordpress.com>

See Also[parana sdm_area](#)**Examples**

```
# Create sdm_area object:
sa <- sdm_area(parana, cell_size = 50000, crs = 6933)

# Include predictors:
sa <- add_predictors(sa, bioc)

# Check predictors' names:
get_predictor_names(sa)
```

predict_sdm

Predict SDM models in new data

Description

This function projects SDM models to new scenarios

Usage

```
predict_sdm(m,
            scen = NULL,
            metric = "ROC",
            th = 0.9,
            tp = "prob",
            ensembles = TRUE,
            file = NULL,
            add.current = TRUE)
```

```
get_predictions(i)
```

```
get_ensembles(i)
```

```
add_predictions(p1, p2)
```

Arguments

m	A input_sdm or a models object.
scen	A scenarios object or NULL. If NULL and m is a input_sdm with a scenarios slot, it will be used.
metric	A character containing the metric in which the th will be calculated/applied. Default is ROC. See ?mean_validation_metrics for the metrics available.
th	Thresholds for metrics. Can be numeric or a function.

tp	Type of output to be retrieved. See details.
ensembles	Boolean. Should ensembles be calculated? If TRUE a series of ensembles are obtained. See details.
file	File to save predictions.
add.current	If current scenario is not available, predictors will be used as the current scenario.
i	A input_sdm or a predictions object.
p1	A predictions object.
p2	A predictions object.

Details

tp is a parameter to be passed on caret to retrieve either the probabilities of classes (tp="prob") or the raw output (tp="raw"), which could vary depending on the algorithm used, but usually would be on of the classes (factor vector with presences and pseudoabsences).

When ensembles is set to TRUE, three ensembles are currently implemented. mean_occ_prob is the mean occurrence probability, which is a simple mean of predictions, wmean_AUC is the same mean_occ_prob, but weighted by AUC, and committee_avg is the committee average, as known as majority rule, where predictions are binarized and then a mean is obtained.

get_predictions returns the list of all predictions to all scenarios, all species, all algorithms and all repetitions. Useful for those who wish to implement their own ensemble methods.

get_ensembles returns a matrix of data.frames, where each column is a scenario and each row is a species.

scenarios_names returns the scenarios names in a sdm_area or input_sdm object.

get_scenarios_data returns the data from scenarios in a sdm_area or input_sdm object.

Value

A input_sdm or a predictions object.

Author(s)

Luíz Fernando Esser (luizesser@gmail.com) <https://luizfesser.wordpress.com>

See Also

[sdm_area](#) [input_sdm](#) [mean_validation_metrics](#)

Examples

```
# Create sdm_area object:
set.seed(1)
sa <- sdm_area(parana, cell_size = 100000, crs = 6933)

# Include predictors:
sa <- add_predictors(sa, bioc) |> select_predictors(c("bio1", "bio12"))
```

```

# Include scenarios:
sa <- add_scenarios(sa)

# Create occurrences:
oc <- occurrences_sdm(occ, crs = 6933) |> join_area(sa)

# Create input_sdm:
i <- input_sdm(oc, sa)

# Pseudoabsence generation:
i <- pseudoabsences(i, method="random", n_set=2)

# Custom trainControl:
ctrl_sdm <- caret::trainControl(method = "boot",
                                number = 1,
                                repeats = 1,
                                classProbs = TRUE,
                                returnResamp = "all",
                                summaryFunction = summary_sdm,
                                savePredictions = "all")

# Train models:
i <- train_sdm(i, algo = c("naive_bayes"), ctrl=ctrl_sdm) |>
  suppressWarnings()

# Predict models:
i <- predict_sdm(i, th = 0.8)
i

```

print.input_sdm	<i>Print method for input_sdm</i>
-----------------	-----------------------------------

Description

Print method for input_sdm

Usage

```
## S3 method for class 'input_sdm'
print(x, ...)
```

Arguments

x	input_sdm object
...	passed to other methods

Value

Concatenate structured characters to showcase what is stored in the object.

<code>print.models</code>	<i>Print method for models</i>
---------------------------	--------------------------------

Description

Print method for models

Usage

```
## S3 method for class 'models'
print(x, ...)
```

Arguments

- `x` models object
- `...` passed to other methods

Value

Concatenate structured characters to showcase what is stored in the object.

<code>print.occurrences</code>	<i>Print method for occurrences</i>
--------------------------------	-------------------------------------

Description

Print method for occurrences

Usage

```
## S3 method for class 'occurrences'
print(x, ...)
```

Arguments

- `x` occurrences object
- `...` passed to other methods

Value

Concatenate structured characters to showcase what is stored in the object.

print.predictions	<i>Print method for predictions</i>
-------------------	-------------------------------------

Description

Print method for predictions

Usage

```
## S3 method for class 'predictions'
print(x, ...)
```

Arguments

x	predictions object
...	passed to other methods

Value

Concatenate structured characters to showcase what is stored in the object.

pseudoabsences	<i>Obtain Pseudoabsences</i>
----------------	------------------------------

Description

This function obtains pseudoabsences given a set of predictors.

Usage

```
pseudoabsences(occ,
  pred = NULL,
  method = "random",
  n_set = 10,
  n_pa = NULL,
  variables_selected = NULL,
  th = 0,
  size = 1,
  crs = 4326)
```

```
n_pseudoabsences(i)
```

```
pseudoabsence_method(i)
```

```
pseudoabsence_data(i)
```

Arguments

<code>occ</code>	A <code>occurrences_sdm</code> or <code>input_sdm</code> object.
<code>pred</code>	A <code>sdm_area</code> object. If <code>NULL</code> and <code>occ</code> is a <code>input_sdm</code> , <code>pred</code> will be retrieved from <code>occ</code> .
<code>method</code>	Method to create pseudoabsences. One of: "random", "bioclim", "mahal.dist" or "buffer_sdm".
<code>n_set</code>	numeric. Number of datasets of pseudoabsence to create.
<code>n_pa</code>	numeric. Number of pseudoabsences to be generated in each dataset created. If <code>NULL</code> then the function prevents imbalance by using the same number of presence records (<code>n_records(occ)</code>). If you want to address different sizes to each species, you must provide a named vector (as in <code>n_records(occ)</code>).
<code>variables_selected</code>	A vector with variables names to be used while building pseudoabsences. Only used when method is not "random".
<code>th</code>	numeric Threshold to be applied in bioclim/mahal.dist projections. See details.
<code>size</code>	numeric The distance between the record and the margin of the buffer (i.e. buffer radius).
<code>crs</code>	numeric Indicates which EPSG it the size in.
<code>i</code>	A <code>input_sdm</code> object.

Details

`pseudoabsences` is used in the SDM workflow to obtain pseudoabsences, a step necessary for most of the algorithms to run. We implemented three methods so far: "random", which is self-explanatory, "bioclim", "mahal.dist" and "buffer_sdm". The two last are built with the idea that pseudoabsences should be environmentally different from presences. Thus, we implemented two presence-only methods to infer the distribution of the species. "bioclim" uses an envelope approach (bioclimatic envelope), while "mahal.dist" uses a distance approach (mahalanobis distance). `th` parameter enters here as a threshold to binarize those results. Pseudoabsences are retrieved outside the projected distribution of the species.

`n_pseudoabsences` returns the number of pseudoabsences obtained per species.

`pseudoabsence_method` returns the method used to obtain pseudoabsences.

`pseudoabsence_data` returns a list of species names. Each species name will have a lists with pseudoabsences data from class `sf`.

Value

A `occurrences_sdm` or `input_sdm` object with pseudoabsence data.

Author(s)

Luiz Fernando Esser (luizesser@gmail.com) <https://luizfesser.wordpress.com>

See Also

`link{input_sdm}` [sdm_area](#) [occurrences_sdm](#)

Examples

```
# Create sdm_area object:
sa <- sdm_area(parana, cell_size = 25000, crs = 6933)

# Include predictors:
sa <- add_predictors(sa, bioc) |> select_predictors(c("bio1", "bio4", "bio12"))

# Include scenarios:
sa <- add_scenarios(sa)

# Create occurrences:
oc <- occurrences_sdm(occ, crs = 6933) |> join_area(sa)

# Create input_sdm:
i <- input_sdm(oc, sa)

# Pseudoabsence generation:
i <- pseudoabsences(i, method="random")
```

rivs

Hydrologic Variables

Description

A sf LINESTRING object with hydrologic variables (LENGTH_KM and DIST_DN_KM) for the Paraná state in Brazil. Data obtained from HydroSHEDS for river flows $\geq 10\text{m}^3/\text{s}$.

Usage

```
rivs
```

Format

'rivs' A sf with 1031 attributes and 2 fiels:

LENGTH_KM Length of the river reach segment, in kilometers.

DIST_DN_KM Distance from the reach outlet, i.e., the most downstream pixel of the reach, to the final downstream location along the river network, in kilometers. This downstream location is either the pour point into the ocean or an endorheic sink.

Source

<<https://www.hydrosheds.org/>>

salm	<i>Salminus brasiliensis occurrence data</i>
------	--

Description

A data.frame object with Salminus brasiliensis occurrence data obtained from GBIF and filtered with Parana state sf.

Usage

```
salm
```

Format

```
## 'salm' A data.frame with 46 rows and 3 columns (EPSG:6933):
```

```
species Species name
```

```
decimalLongitude Longitude in meters
```

```
decimalLatitude Latitude in meters
```

Source

```
<https://www.gbif.org>
```

scen	<i>Bioclimatic Variables</i>
------	------------------------------

Description

A stars object with bioclimatic variables (bio1, bio4 and bio12) and four future scenarios for the Parana state in Brazil. Data from MIROC6 GCM from WorldClim 2.1 at 10 arc-min resolution.

Usage

```
scen
```

Format

```
## 'scen' A stars with 4 attribute and 3 bands:
```

```
ca_ssp245_2090 Intermediate scenario for the year 2090 and GCM CanESM5
```

```
ca_ssp585_2090 Extreme scenario for the year 2090 and GCM CanESM5
```

```
mi_ssp245_2090 Intermediate scenario for the year 2090 and GCM MIROC6
```

```
mi_ssp585_2090 Extreme scenario for the year 2090 and GCM MIROC6
```

```
bio1 Annual Mean Temperature
```

```
bio4 Temperature Seasonality
```

```
bio12 Annual Precipitation
```

Source

<<https://www.worldclim.org/>>

scen_rs	<i>Bioclimatic Variables</i>
---------	------------------------------

Description

A stars object with bioclimatic variables (bio1, bio4 and bio12) and four future scenarios for the Rio Grande do Sul state in Brazil. Data from MIROC6 GCM from WorldClim 2.1 at 10 arc-min resolution.

Usage

```
scen_rs
```

Format

'scen_rs' A stars with 5 attribute and 3 bands:

current Current scenario with the average values for the years 1970-2000

ca_ssp245_2090 Intermediate scenario for the year 2090 and GCM CanESM5

ca_ssp585_2090 Extreme scenario for the year 2090 and GCM CanESM5

mi_ssp245_2090 Intermediate scenario for the year 2090 and GCM MIROC6

mi_ssp585_2090 Extreme scenario for the year 2090 and GCM MIROC6

bio1 Annual Mean Temperature

bio4 Temperature Seasonality

bio12 Annual Precipitation

Source

<<https://www.worldclim.org/>>

sdm_area	<i>Create a sdm_area object</i>
----------	---------------------------------

Description

This function creates a new sdm_area object.

Usage

```
sdm_area(x, cell_size = NULL, crs = NULL, variables_selected = NULL,
         gdal = TRUE, crop_by = NULL, lines_as_sdm_area = FALSE)
```

```
get_sdm_area(i)
```

```
add_sdm_area(sa1, sa2)
```

Arguments

x	A shape or a raster. Usually a shape from sf class, but rasters from stars, rasterStack or SpatRaster class are also allowed.
cell_size	numeric. The cell size to be used in models.
crs	numeric. Indicates which EPSG should the output grid be in. If NULL, epsg from x is used.
variables_selected	A character vector with variables in x to be used in models. If NULL (standard), all variables in x are used.
gdal	Boolean. Force the use or not of GDAL when available. See details.
crop_by	A shape from sf to crop x.
lines_as_sdm_area	Boolean. If x is a sf with LINESTRING geometry, it can be used to model species distribution in lines and not grid cells.
i	A sdm_area or a input_sdm object.
sa1	A sdm_area object.
sa2	A sdm_area object.

Details

The function returns a sdm_area object with a grid built upon the x parameter. There are two ways to make the grid and resample the variables in sdm_area: with and without gdal. As standard, if gdal is available in you machine it will be used (gdal = TRUE), otherwise sf/stars will be used. get_sdm_area will return the grid built by sdm_area. add_sdm_area will sum two sdm_area objects. As geoprocessing in caretSDM is performed using sf objects, add_sdm_area simply applies a rbind in the two different areas.

Value

A sdm_area object containing:

grid	sf with POLYGON geometry representing the grid for the study area.
cell_size	numeric information regarding the size of the cell used to rescale variables to the study area, representing also the cell size in the grid.

Author(s)

Luíz Fernando Esser (luizesser@gmail.com) and Reginaldo Ré. <https://luizfesser.wordpress.com>

See Also

[WorldClim_data parana input_sdm, add_predictors](#)

Examples

```
# Create sdm_area object:
sa_area <- sdm_area(parana, cell_size = 50000, crs = 6933)

# Create sdm_area using a subset of rivs (lines):
sa_rivers <- sdm_area(rivs[c(1:100)], cell_size = 100000, crs = 6933, lines_as_sdm_area = TRUE)
```

sdm_as_stars	<i>sdm_as_X functions to transform caretSDM data into other classes.</i>
--------------	--

Description

This functions transform data from a caretSDM object to be used in other packages.

Usage

```
sdm_as_stars(x,
             what = NULL,
             spp = NULL,
             scen = NULL,
             id = NULL,
             ens = NULL)

sdm_as_raster(x, what = NULL, spp = NULL, scen = NULL, id = NULL, ens = NULL)

sdm_as_terra(x, what = NULL, spp = NULL, scen = NULL, id = NULL, ens = NULL)
```

Arguments

x	A caretSDM object.
what	Sometimes multiple data inside x could be transformed. This parameter allows users to specify what needs to be converted. It can be one of: "predictors", "scenarios", "predictions" or "ensembles".
spp	character. Which species should be converted?
scen	character. Which scenario should be converted?
id	character. Which id should be converted?
ens	character. Which ensemble should be converted?

Value

The output is the desired class.

Author(s)

Luiz Fernando Esser (luizesser@gmail.com) <https://luizfesser.wordpress.com>

Examples

```
# Create sdm_area object:
sa <- sdm_area(parana, cell_size = 100000, crs = 6933)

# Include predictors:
sa <- add_predictors(sa, bioc) |> select_predictors(c("bio1", "bio12"))

# Include scenarios:
sa <- add_scenarios(sa)

# Create occurrences:
oc <- occurrences_sdm(occ, crs = 6933) |> join_area(sa)

# Create input_sdm:
i <- input_sdm(oc, sa)

# Pseudoabsence generation:
i <- pseudoabsences(i, method = "random", n_set = 2)

# Custom trainControl:
ctrl_sdm <- caret::trainControl(method = "boot",
                                number = 1,
                                classProbs = TRUE,
                                returnResamp = "all",
                                summaryFunction = summary_sdm,
                                savePredictions = "all")

# Train models:
i <- train_sdm(i, algo = c("naive_bayes"), ctrl = ctrl_sdm) |>
  suppressWarnings()
```

```
# Predict models:
i <- predict_sdm(i, th = 0.8)

# Transform in stars:
sdm_as_stars(i)
```

select_predictors	<i>Tidyverse methods for caretSDM objects</i>
-------------------	---

Description

Set of functions to facilitate the use of caretSDM through tidyverse grammatics.

Usage

```
select_predictors(x, ...)

## S3 method for class 'sdm_area'
select(.data, ...)

## S3 method for class 'input_sdm'
select(.data, ...)

## S3 method for class 'sdm_area'
mutate(.data, ...)

## S3 method for class 'input_sdm'
mutate(.data, ...)

## S3 method for class 'sdm_area'
filter(.data, ..., .by, .preserve)

## S3 method for class 'input_sdm'
filter(.data, ..., .by, .preserve)

## S3 method for class 'occurrences'
filter(.data, ..., .by, .preserve)

filter_species(x, spp = NULL, ...)
```

Arguments

x	sdm_area or input_sdm object.
...	character arguments to pass to the given function.
.data	Data to pass to tidyr function.
.by	See ?dplyr::filter.

.preserve See ?dplyr::filter.
 spp Species to be filtered.

Value

The transformed sdm_area/input_sdm object.

Examples

```
# Create sdm_area object:
sa <- sdm_area(parana, cell_size = 25000, crs = 6933)

# Include predictors:
sa <- add_predictors(sa, bioc) |> select_predictors(c("bio1", "bio4", "bio12"))
```

summary_sdm	<i>Calculates performance across resamples</i>
-------------	--

Description

This function is used in `caret::trainControl(summaryFunction=summary_sdm)` to calculate performance metrics across resamples.

Usage

```
summary_sdm(data, lev = NULL, model = NULL, custom_fun=NULL)

summary_sdm_presence_only(data, lev, threshold)

validate_on_independent_data(model, data_independent, obs_col_name)
```

Arguments

data A data.frame with observed and predicted values.
 lev A character vector of factors levels for the response.
 model Models names taken from train object.
 custom_fun A custom function to be applied in models (not yet implemented).
 threshold Threshold for presence-only models.
 data_independent independent data.frame to calculate metrics.
 obs_col_name The name of the column with observed values.

Details

See ?`caret::defaultSummary` for more details and options to pass on `caret::trainControl`.

Value

A input_sdm or a predictions object.

Author(s)

Luíz Fernando Esser (luizesser@gmail.com) <https://luizfesser.wordpress.com>

See Also

[train_sdm](#)

Examples

```
# Create sdm_area object:
sa <- sdm_area(parana, cell_size = 100000, crs = 6933)

# Include predictors:
sa <- add_predictors(sa, bioc) |> select_predictors(c("bio1", "bio12"))

# Include scenarios:
sa <- add_scenarios(sa)

# Create occurrences:
oc <- occurrences_sdm(occ, crs = 6933) |> join_area(sa)

# Create input_sdm:
i <- input_sdm(oc, sa)

# Pseudoabsence generation:
i <- pseudoabsences(i, method = "random")

# Custom trainControl:
ctrl_sdm <- caret::trainControl(method = "repeatedcv",
                                number = 2,
                                repeats = 1,
                                classProbs = TRUE,
                                returnResamp = "all",
                                summaryFunction = summary_sdm,
                                savePredictions = "all")

# Train models:
i <- train_sdm(i, algo = c("naive_bayes"), ctrl=ctrl_sdm) |>
suppressWarnings()
```

Description

This function is a wrapper to fit models in caret using caretSDM data.

Usage

```
train_sdm(occ,
          pred = NULL,
          algo,
          ctrl = NULL,
          variables_selected = NULL,
          parallel = FALSE,
          ...)
```

```
get_tune_length(i)
```

```
algorithms_used(i)
```

```
get_models(i)
```

```
get_validation_metrics(i)
```

```
mean_validation_metrics(i)
```

```
add_models(m1, m2)
```

Arguments

occ	A occurrences or a input_sdm object.
pred	A predictors object. If occ is a input_sdm object, then pred is obtained from it.
algo	A character vector. Algorithms to be used. For a complete list see (https://topepo.github.io/caret/available_models.html) or in caretSDM::algorithms.
ctrl	A trainControl object to be used to build models. See ?caret::trainControl and details.
variables_selected	A vector of variables to be used as predictors. If NULL, predictors names from pred will be used. Can also be a selection method (e.g. 'vif').
parallel	Should a paralelization method be used (not yet implemented)?
...	Additional arguments to be passed to caret::train function.
i	A models or a input_sdm object.
m1	A models object.
m2	A models object.

Details

The object `algorithms` has a table comparing algorithms available. If the function detects that the necessary packages are not available it will ask for installation. This will happen just in the first time you use the algorithm. `caret::trainControl` holds multiple resources for validation and model tuning. Make sure to understand its parameters beforehand. As it is a key function in the modeling process, we also implemented spatial crossvalidation on it. You can set methods to be `cv_spatial` or `cv_cluster` and `train_sdm` will detect that and apply the method according to `blockCV` package.

`get_tune_length` return the length used in grid-search for tuning.

`algorithms_used` return the names of the algorithms used in the modeling process.

`get_models` returns a list with trained models (class `train`) to each species.

`get_validation_metrics` return a list with a data.frame to each species with complete values for ROC, Sensitivity, Specificity, with their respective Standard Deviations (SD) and TSS to each of the algorithms and pseudoabsence datasets used.

`mean_validation_metrics` return a list with a tibble to each species summarizing values for ROC, Sensitivity, Specificity and TSS to each of the algorithms used.

Value

A models or a `input_sdm` object.

Author(s)

Luíz Fernando Esser (luizesser@gmail.com) <https://luizfesser.wordpress.com>

See Also

[input_sdm](#) [sdm_area](#) [algorithms](#)

Examples

```
# Create sdm_area object:
sa <- sdm_area(parana, cell_size = 100000, crs = 6933)

# Include predictors:
sa <- add_predictors(sa, bioc) |> select_predictors(c("bio1", "bio12"))

# Include scenarios:
sa <- add_scenarios(sa)

# Create occurrences:
oc <- occurrences_sdm(occ, crs = 6933) |> join_area(sa)

# Create input_sdm:
i <- input_sdm(oc, sa)

# Pseudoabsence generation:
i <- pseudoabsences(i, method = "random")
```

```
# Custom trainControl:
ctrl_sdm <- caret::trainControl(method = "repeatedcv",
                                number = 2,
                                repeats = 1,
                                classProbs = TRUE,
                                returnResamp = "all",
                                summaryFunction = summary_sdm,
                                savePredictions = "all")

# Train models:
i <- train_sdm(i, algo = c("naive_bayes"), ctrl = ctrl_sdm) |>
suppressWarnings()
```

tsne_sdm

tSNE

Description

This function calculates tSNE with presences and pseudoabsences data and returns a list of plots.

Usage

```
tsne_sdm(occ, pred = NULL, variables_selected = NULL)
```

Arguments

occ A occurrences or input_sdm object.

pred A predictors object. If occ is of class input_sdm, then pred is retrieved from it.

variables_selected Variable to be used in t-SNE. It can also be 'vif', if previously calculated.

Value

A list of plots, where each plot is a tSNE for a given pseudoabsence dataset.

Author(s)

Luíz Fernando Esser (luizesser@gmail.com) <https://luizfesser.wordpress.com>

tuneGrid_sdm	<i>Retrieve tuneGrid from models</i>
--------------	--------------------------------------

Description

This function aims to retrieve the tune grid used to build models.

Usage

```
tuneGrid_sdm(i)
```

Arguments

i A input_sdm object containing models.

Value

A list with data.frames each one representing the table of a given model.

Author(s)

Luíz Fernando Esser (luizesser@gmail.com) <https://luizfesser.wordpress.com>

Examples

```
# Create sdm_area object:
set.seed(1)
sa <- sdm_area(parana, cell_size = 100000, crs = 6933)

# Include predictors:
sa <- add_predictors(sa, bioc) |> select_predictors(c("bio1", "bio12"))

# Include scenarios:
sa <- add_scenarios(sa)

# Create occurrences:
oc <- occurrences_sdm(occ, crs = 6933) |> join_area(sa)

# Create input_sdm:
i <- input_sdm(oc, sa)

# Pseudoabsence generation:
i <- pseudoabsences(i, method="random", n_set=2)

# Custom trainControl:
ctrl_sdm <- caret::trainControl(method = "boot",
                                number = 1,
                                repeats = 1,
                                classProbs = TRUE,
                                returnResamp = "all",
```

```

summaryFunction = summary_sdm,
savePredictions = "all")

# Train models:
i <- train_sdm(i, algo = c("naive_bayes"), ctrl=ctrl_sdm) |>
  suppressWarnings()

# Retrieve tuneGrid from model:
tuneGrid_sdm(i)

```

use_esm

Ensemble of Small Models (ESM) in caretSDM

Description

This functions set parameters to run a ESM when running train_sdm.

Usage

```
use_esm(x, spp = NULL, n_records = 20)
```

Arguments

x	A occurrences or input_sdm object containing occurrences.
spp	A vector of species names containing the species which the ESM must be applied. Standard is NULL.
n_records	Numeric. Number of species records to apply the ESM. Standard is 20.

Details

We supply two different ways to apply the ESM. If species names are provided, then ESM will be applied only in given species. If a number of species records is provided, then ESM will be applied in every species with number of records bellow the given threshold. As standard, use_esm will be apply to every species with less then 20 records.

Value

A input_sdm or occurrences object with ESM parameters.

Author(s)

Luíz Fernando Esser (luizesser@gmail.com) <https://luizfesser.wordpress.com>

Examples

```
# Create sdm_area object:
sa <- sdm_area(parana, cell_size = 100000, crs = 6933)

# Include predictors:
sa <- add_predictors(sa, bioc) |> select_predictors(c("bio1", "bio4", "bio12"))

# Include scenarios:
sa <- add_scenarios(sa)

# Create occurrences:
oc <- occurrences_sdm(occ, crs = 6933) |> join_area(sa)

# Create input_sdm:
i <- input_sdm(oc, sa)

# Use MEM:
i <- use_esm(i, n_records = 999)
```

use_mem

MacroEcological Models (MEM) in caretSDM

Description

This function sums all species records into one. Should be used before the data cleaning routine.

Usage

```
use_mem(x, add = TRUE, name = "MEM")
```

Arguments

x	A occurrences or input_sdm object containing occurrences.
add	Logical. Should the new MEM records be added to the pool (TRUE) of species or the output should have only the summed records (FALSE)? Standard is TRUE.
name	How should the new records be named? Standard is "MEM".

Value

A input_sdm or occurrences object with MEM data.

Author(s)

Luíz Fernando Esser (luizesser@gmail.com) <https://luizfesser.wordpress.com>

Examples

```
# Create sdm_area object:
sa <- sdm_area(parana, cell_size = 25000, crs = 6933)

# Include predictors:
sa <- add_predictors(sa, bioc) |> select_predictors(c("bio1", "bio4", "bio12"))

# Include scenarios:
sa <- add_scenarios(sa)

# Create occurrences:
oc <- occurrences_sdm(occ, crs = 6933) |> join_area(sa)

# Create input_sdm:
i <- input_sdm(oc, sa)

# Use MEM:
i <- use_mem(i)
```

varImp_sdm

*Calculation of variable importance for models***Description**

This function retrieves variable importance as a function of ROC curves to each predictor.

Usage

```
varImp_sdm(m, id = NULL, ...)
```

Arguments

<code>m</code>	A models or input_sdm object.
<code>id</code>	Vector of model ids to filter varImp calculation.
<code>...</code>	Parameters passing to caret::varImp().

Value

A data.frame with variable importance data.

Author(s)

Luíz Fernando Esser (luizesser@gmail.com) <https://luizfesser.wordpress.com>

Examples

```
# Create sdm_area object:
sa <- sdm_area(parana, cell_size = 100000, crs = 6933)

# Include predictors:
sa <- add_predictors(sa, bioc) |> select_predictors(c("bio1", "bio12"))

# Include scenarios:
sa <- add_scenarios(sa)

# Create occurrences:
oc <- occurrences_sdm(occ, crs = 6933) |> join_area(sa)

# Create input_sdm:
i <- input_sdm(oc, sa)

# Pseudoabsence generation:
i <- pseudoabsences(i, method = "random")

# Custom trainControl:
ctrl_sdm <- caret::trainControl(method = "repeatedcv",
                                number = 2,
                                repeats = 1,
                                classProbs = TRUE,
                                returnResamp = "all",
                                summaryFunction = summary_sdm,
                                savePredictions = "all")

# Train models:
i <- train_sdm(i, algo = c("naive_bayes"), ctrl = ctrl_sdm) |>
suppressWarnings()

# Variable importance:
varImp_sdm(i)
```

vif_predictors

Calculate VIF

Description

Apply Variance Inflation Factor (VIF) calculation.

Usage

```
vif_predictors(pred, area = "all", th = 0.5, maxobservations = 5000, variables_selected =
NULL)

vif_summary(i)
```

```
selected_variables(i)
```

Arguments

pred	A input_sdm or predictors object.
area	Character. Which area should be used in vif selection? Standard is "all".
th	Threshold to be applied in VIF routine. See ?usdm::vifcor.
maxobservations	Max observations to use to calculate the VIF.
variables_selected	If there is a subset of predictors that should be used in this function, it can be informed using this parameter. If set to NULL (standard) all variables are used.
i	A input_sdm to retrieve information from.

Details

vif_predictors is a wrapper function to run usdm::vifcor in caretSDM.

Value

A input_sdm or predictors object with VIF data.

Author(s)

Luíz Fernando Esser (luizesser@gmail.com) <https://luizfesser.wordpress.com>

See Also

[get_predictor_names](#)

Examples

```
# Create sdm_area object:
sa <- sdm_area(parana, cell_size = 25000, crs = 6933)

# Include predictors:
sa <- add_predictors(sa, bioc) |> select_predictors(c("bio1", "bio4", "bio12"))

# Include scenarios:
sa <- add_scenarios(sa, scen)

# Create occurrences:
oc <- occurrences_sdm(occ, crs = 6933) |> join_area(sa)

# Create input_sdm:
i <- input_sdm(oc, sa)

# VIF calculation:
i <- vif_predictors(i)
```

```
i

# Retrieve information about vif:
vif_summary(i)
selected_variables(i)
```

WorldClim_data

Download WorldClim v.2.1 bioclimatic data

Description

This function allows to download data from WorldClim v.2.1 (<https://www.worldclim.org/data/index.html>) considering multiple GCMs, time periods and SSPs.

Usage

```
WorldClim_data(path = NULL,
               period = "current",
               variable = "bioc",
               year = "2090",
               gcm = "mi",
               ssp = "585",
               resolution = 10)
```

Arguments

path	Directory path to save downloads.
period	Can be "current" or "future".
variable	Allows to specify which variables you want to retrieve Possible entries are: "tmax", "tmin", "prec" and/or "bioc".
year	Specify the year you want to retrieve data. Possible entries are: "2030", "2050", "2070" and/or "2090". You can use a vector to provide more than one entry.
gcm	GCMs to be considered in future scenarios. You can use a vector to provide more than one entry.
ssp	SSPs for future data. Possible entries are: "126", "245", "370" and/or "585". You can use a vector to provide more than one entry.
resolution	You can select one resolution from the following alternatives: 10, 5, 2.5 OR 30.

Details

This function will create a folder. All the data downloaded will be stored in this folder. Note that, despite being possible to retrieve a lot of data at once, it is not recommended to do so, since the data is very heavy.

Value

If data is not downloaded, the function downloads the data and has no return value.

Author(s)

Luíz Fernando Esser (luizesser@gmail.com) <https://luizfesser.wordpress.com>

References

<https://www.worldclim.org/data/index.html>

Examples

```
# download data from multiple periods:
year <- c("2050", "2090")
WorldClim_data(period = "future",
               variable = "bioc",
               year = year,
               gcm = "mi",
               ssp = "126",
               resolution = 10)

# download data from one specific period
WorldClim_data(period = "future",
               variable = "bioc",
               year = "2070",
               gcm = "mi",
               ssp = "585",
               resolution = 10)
```

write_ensembles	<i>Write caretSDM data</i>
-----------------	----------------------------

Description

This function exports caretSDM data.

Usage

```
write_ensembles(x, path = NULL, ext = ".tif", centroid = FALSE)

write_predictions(x, path = NULL, ext = ".tif", centroid = FALSE)

write_predictors(x, path = NULL, ext = ".tif", centroid = FALSE)

write_models(x, path = NULL)
```

```

write_gpkg(x, file_path, file_name)

## S3 method for class 'sdm_area'
write_gpkg(x, file_path, file_name)

write_occurrences(x, path = NULL, grid = FALSE, ...)

write_pseudoabsences(x, path = NULL, ext = ".csv", centroid = FALSE)

write_grid(x, path = NULL, centroid = FALSE)

write_validation_metrics(x, path = NULL)

```

Arguments

x	Object to be written. Can be of class input_sdm, occurrences, predictions or models.
path	A path with filename and the proper extension (see details) or the directory to save files in.
ext	How it should be saved?
centroid	Should coordinates for the centroids of each cell be included? Standard is FALSE.
file_path	A path to save the sdm_area GeoPackage file.
file_name	The name of the sdm_area GeoPackage file to be saved without extension.
grid	Boolean. Return a grid.
...	Arguments to pass to sf::st_write or write.csv.

Details

ext can be set accordingly to the desired output. Possible values are .tif and .asc for rasters, .csv for a spreadsheet, but also one of: c("bna", "csv", "e00", "gdb", "geojson", "gml", "gmt", "gpkg", "gps", "gtm", "gxt", "jml", "map", "mdb", "nc", "ods", "osm", "pbf", "shp", "sqlite", "vdt", "xls", "xlsx"). path ideally should only provide the folder. We recommend using: results/what_are_you_writing. So for writing ensembles users are advised to run: path = "results/ensembles"

Value

No return value, called for side effects.

Author(s)

Luiz Fernando Esser (luizesser@gmail.com) <https://luizfesser.wordpress.com>

Index

* datasets

- algorithms, 6
- bioc, 6
- occ, 17
- parana, 19
- rivs, 33
- salm, 34
- scen, 34
- scen_rs, 35

add_input_sdm(input_sdm), 13

add_models(train_sdm), 41

add_occurrences(occurrences_sdm), 17

add_predictions(predict_sdm), 27

add_predictors, 3, 20, 37

add_scenarios, 4, 20

add_sdm_area(sdm_area), 36

algorithms, 6, 43

algorithms_used(train_sdm), 41

bioc, 4, 6

buffer_sdm, 7

correlate_sdm, 8

data_clean, 9

filter.input_sdm(select_predictors), 39

filter.occurrences(select_predictors), 39

filter.sdm_area(select_predictors), 39

filter.species(select_predictors), 39

GBIF_data, 7, 10, 11, 12, 18

gcms_ensembles, 12

get_coords(occurrences_sdm), 17

get_ensembles(predict_sdm), 27

get_models(train_sdm), 41

get_pca_model(pca_predictors), 19

get_pdp_sdm(pdp_sdm), 20

get_predictions(predict_sdm), 27

get_predictor_names, 50

get_predictor_names(predictors), 25

get_predictors(add_predictors), 3

get_scenarios_data(add_scenarios), 4

get_sdm_area(sdm_area), 36

get_tune_length(train_sdm), 41

get_validation_metrics(train_sdm), 41

input_sdm, 5, 10, 12, 13, 16, 18, 28, 37, 43

is_input_sdm, 15

is_models(is_input_sdm), 15

is_occurrences(is_input_sdm), 15

is_predictions(is_input_sdm), 15

is_sdm_area(is_input_sdm), 15

join_area, 16

mapview_grid(plot_occurrences), 22

mapview_occurrences(plot_occurrences), 22

mapview_predictions(plot_occurrences), 22

mapview_predictors(plot_occurrences), 22

mapview_scenarios(plot_occurrences), 22

mean_validation_metrics, 28

mean_validation_metrics(train_sdm), 41

mutate.input_sdm(select_predictors), 39

mutate.sdm_area(select_predictors), 39

n_pseudoabsences(pseudoabsences), 31

n_records(occurrences_sdm), 17

occ, 17, 18

occurrences_as_df(occurrences_sdm), 17

occurrences_sdm, 10, 12, 14, 16, 17, 32

parana, 19, 27, 37

pca_predictors, 19

pca_summary(pca_predictors), 19

pdp_sdm, 20

plot_background (plot_occurrences), 22
 plot_grid (plot_occurrences), 22
 plot_niche (plot_occurrences), 22
 plot_occurrences, 22
 plot_predictions (plot_occurrences), 22
 plot_predictors (plot_occurrences), 22
 plot_scenarios (plot_occurrences), 22
 predict_sdm, 27
 prediction_change_sdm, 24
 predictors, 4, 10, 12, 25
 print.input_sdm, 29
 print.models, 30
 print.occurrences, 30
 print.predictions, 31
 pseudoabsence_data (pseudoabsences), 31
 pseudoabsence_method (pseudoabsences), 31
 pseudoabsences, 16, 31

 rivs, 33

 salm, 34
 scen, 34
 scen_rs, 35
 scenarios_names, 24
 scenarios_names (add_scenarios), 4
 sdm_area, 4, 5, 10, 12, 14, 16, 20, 27, 28, 32, 36, 43
 sdm_as_raster (sdm_as_stars), 37
 sdm_as_stars, 37
 sdm_as_terra (sdm_as_stars), 37
 select.input_sdm (select_predictors), 39
 select.sdm_area (select_predictors), 39
 select_predictors, 39
 select_scenarios (add_scenarios), 4
 selected_variables (vif_predictors), 49
 set_predictor_names (predictors), 25
 set_scenarios_names (add_scenarios), 4
 set_variables_names (predictors), 25
 species_names, 24
 species_names (occurrences_sdm), 17
 summary_sdm, 40
 summary_sdm_presence_only (summary_sdm), 40

 test_variables_names (predictors), 25
 train_sdm, 41, 41
 tsne_sdm, 44
 tuneGrid_sdm, 45

 use_esm, 46
 use_mem, 47

 validate_on_independent_data (summary_sdm), 40
 varImp_sdm, 21, 48
 vif_predictors, 20, 49
 vif_summary (vif_predictors), 49

 WorldClim_data, 24, 37, 51
 write_ensembles, 52
 write_gpkg (write_ensembles), 52
 write_grid (write_ensembles), 52
 write_models (write_ensembles), 52
 write_occurrences (write_ensembles), 52
 write_predictions (write_ensembles), 52
 write_predictors (write_ensembles), 52
 write_pseudoabsences (write_ensembles), 52
 write_validation_metrics (write_ensembles), 52