

Package ‘tidysdm’

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Title Species Distribution Models with Tidymodels

Version 1.0.3

Description Fit species distribution models (SDMs) using the 'tidymodels' framework, which provides a standardised interface to define models and process their outputs. 'tidysdm' expands 'tidymodels' by providing methods for spatial objects, models and metrics specific to SDMs, as well as a number of specialised functions to process occurrences for contemporary and palaeo datasets. The full functionalities of the package are described in Leonardi et al. (2023) <[doi:10.1101/2023.07.24.550358](https://doi.org/10.1101/2023.07.24.550358)>.

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add_member	<i>Add best member of workflow to a simple ensemble</i>
------------	---

Description

This function adds member(s) to a [simple_ensemble\(\)](#) object, taking the best member from each workflow provided. It is possible to pass individual `tune_results` objects from a tuned workflow, or a `workflowsets::workflow_set()`.

Usage

```
add_member(x, member, ...)

## Default S3 method:
add_member(x, member, ...)

## S3 method for class 'tune_results'
add_member(x, member, metric = NULL, id = NULL, ...)

## S3 method for class 'workflow_set'
add_member(x, member, metric = NULL, ...)
```

Arguments

x	a simple_ensemble to which member(s) will be added
member	a tune_results , or a workflowsets::workflow_set
...	not used at the moment.
metric	A character string (or NULL) for which metric to optimize. If NULL, the first metric is used.
id	the name to be given to this workflow in the wflow_id column.

Value

a [simple_ensemble](#) with additional member(s)

add_repeat	<i>Add repeat(s) to a repeated ensemble</i>
------------	---

Description

This function adds repeat(s) to a [repeat_ensemble](#) object, where each repeat is a [simple_ensemble](#). All repeats must contain the same members, selected using the same metric.

Usage

```
add_repeat(x, rep, ...)

## Default S3 method:
add_repeat(x, rep, ...)

## S3 method for class 'simple_ensemble'
add_repeat(x, rep, ...)

## S3 method for class 'list'
add_repeat(x, rep, ...)
```

Arguments

x	a repeat_ensemble to which repeat(s) will be added
rep	a repeat, as a single simple_ensemble , or a list of simple_ensemble objects
...	not used at the moment.

Value

a [repeat_ensemble](#) with additional repeat(s)

`autoplot.simple_ensemble`*Plot the results of a simple ensemble*

Description

This `autoplot()` method plots performance metrics that have been ranked using a metric.

Usage

```
## S3 method for class 'simple_ensemble'
autoplot(
  object,
  rank_metric = NULL,
  metric = NULL,
  std_errs = stats::qnorm(0.95),
  ...
)
```

Arguments

<code>object</code>	A simple_ensemble whose elements have results.
<code>rank_metric</code>	A character string for which metric should be used to rank the results. If none is given, the first metric in the metric set is used (after filtering by the metric option).
<code>metric</code>	A character vector for which metrics (apart from <code>rank_metric</code>) to be included in the visualization. If <code>NULL</code> (the default), all available metrics will be plotted
<code>std_errs</code>	The number of standard errors to plot (if the standard error exists).
<code>...</code>	Other options to pass to <code>autoplot()</code> . Currently unused.

Details

This function is intended to produce a default plot to visualize helpful information across all possible applications of a [simple_ensemble](#). More sophisticated plots can be produced using standard `ggplot2` code for plotting.

The x-axis is the workflow rank in the set (a value of one being the best) versus the performance metric(s) on the y-axis. With multiple metrics, there will be facets for each metric, with the `rank_metric` first (if any was provided; otherwise the metric used to create the [simple_ensemble](#) will be used).

If multiple resamples are used, confidence bounds are shown for each result (95% confidence, by default).

Value

A `ggplot` object.

Examples

```
# we use the two_class_example from `workflowsets`
two_class_ens <- simple_ensemble() %>%
  add_member(two_class_res, metric = "roc_auc")
autoplot(two_class_ens)
```

```
autoplot.spatial_initial_split
```

Create a ggplot for a spatial initial rsplit.

Description

This method provides a good visualization method for a spatial initial rsplit.

Usage

```
## S3 method for class 'spatial_initial_split'
autoplot(object, ..., alpha = 0.6)
```

Arguments

object	A spatial_initial_rsplit object. Note that only resamples made from sf objects create spatial_initial_rsplit objects; this function will not work for resamples made with non-spatial tibbles or data.frames.
...	Options passed to <code>ggplot2::geom_sf()</code> .
alpha	Opacity, passed to <code>ggplot2::geom_sf()</code> . Values of alpha range from 0 to 1, with lower values corresponding to more transparent colors.

Details

This plot method is a wrapper around the standard spatial_rsplit method, but it re-labels the folds as *Testing* and *Training* following the convention for a standard initial_split object

Value

A ggplot object with each fold assigned a color, made using `ggplot2::geom_sf()`.

Examples

```
set.seed(123)
block_initial <- spatial_initial_split(boston_canopy,
  prop = 1 / 5, spatial_block_cv
)
autoplot(block_initial)
```

blockcv2rsample*Convert an object created with blockCV to an rsample object*

Description

This function converts objects created with blockCV to rsample objects that can be used by tidysdm. BlockCV provides more sophisticated sampling options than the spatialsample library. For example, it is possible to stratify the sampling to ensure that presences and absences are evenly distributed among the folds (see the example below).

Usage

```
blockcv2rsample(x, data)
```

Arguments

x	a object created with a blockCV function
data	the sf object used to create x

Details

Note that currently only objects of type `cv_spatial` and `cv_cluster` are supported.

Value

an rsample object

Examples

```
library(blockCV)
points <- read.csv(system.file("extdata/", "species.csv",
  package = "blockCV"
))
pa_data <- sf::st_as_sf(points, coords = c("x", "y"), crs = 7845)
sb1 <- cv_spatial(
  x = pa_data,
  column = "occ", # the response column to balance the folds
  k = 5, # number of folds
  size = 350000, # size of the blocks in metres
  selection = "random", # random blocks-to-fold
  iteration = 10
) # find evenly dispersed folds
sb1_rsample <- blockcv2rsample(sb1, pa_data)
class(sb1_rsample)
autoplot(sb1_rsample)
```

boyce_cont	<i>Boyce continuous index (BCI)</i>
------------	-------------------------------------

Description

This function the Boyce Continuous Index, a measure of model accuracy appropriate for Species Distribution Models with presence only data (i.e. using pseudoabsences or background). The algorithm used here comes from the package `enmSdm`, and uses multiple overlapping windows.

Usage

```
boyce_cont(data, ...)

## S3 method for class 'data.frame'
boyce_cont(
  data,
  truth,
  ...,
  estimator = NULL,
  na_rm = TRUE,
  event_level = "first",
  case_weights = NULL
)

## S3 method for class 'sf'
boyce_cont(data, ...)

boyce_cont_vec(
  truth,
  estimate,
  estimator = NULL,
  na_rm = TRUE,
  event_level = "first",
  case_weights = NULL,
  ...
)
```

Arguments

<code>data</code>	Either a <code>data.frame</code> containing the columns specified by the <code>truth</code> and <code>estimate</code> arguments, or a table/matrix where the true class results should be in the columns of the table.
<code>...</code>	A set of unquoted column names or one or more <code>dplyr</code> selector functions to choose which variables contain the class probabilities. If <code>truth</code> is binary, only 1 column should be selected, and it should correspond to the value of <code>event_level</code> . Otherwise, there should be as many columns as factor levels of <code>truth</code> and the ordering of the columns should be the same as the factor levels of <code>truth</code> .

truth	The column identifier for the true class results (that is a factor). This should be an unquoted column name although this argument is passed by expression and supports quasiquotation (you can unquote column names). For <code>_vec()</code> functions, a factor vector.
estimator	One of "binary", "hand_till", "macro", or "macro_weighted" to specify the type of averaging to be done. "binary" is only relevant for the two class case. The others are general methods for calculating multiclass metrics. The default will automatically choose "binary" if truth is binary, "hand_till" if truth has >2 levels and case_weights isn't specified, or "macro" if truth has >2 levels and case_weights is specified (in which case "hand_till" isn't well-defined).
na_rm	A logical value indicating whether NA values should be stripped before the computation proceeds.
event_level	A single string. Either "first" or "second" to specify which level of truth to consider as the "event". This argument is only applicable when estimator = "binary". The default uses an internal helper that generally defaults to "first"
case_weights	The optional column identifier for case weights. This should be an unquoted column name that evaluates to a numeric column in data. For <code>_vec()</code> functions, a numeric vector.
estimate	If truth is binary, a numeric vector of class probabilities corresponding to the "relevant" class. Otherwise, a matrix with as many columns as factor levels of truth. It is assumed that these are in the same order as the levels of truth.

Details

There is no multiclass version of this function, it only operates on binary predictions (e.g. presences and absences in SDMs).

Value

A tibble with columns `.metric`, `.estimator`, and `.estimate` and 1 row of values. For grouped data frames, the number of rows returned will be the same as the number of groups.

References

- Boyce, M.S., P.R. Vernier, S.E. Nielsen and F.K.A. Schmiegelow. 2002. Evaluating resource selection functions. *Ecol. Model.*, 157, 281-300.
- Hirzel, A.H., G. Le Lay, V. Helfer, C. Randin and A. Guisan. 2006. Evaluating the ability of habitat suitability models to predict species presences. *Ecol. Model.*, 199, 142-152.

See Also

Other class probability metrics: [kap_max\(\)](#), [tss_max\(\)](#)

Examples

```
boyce_cont(two_class_example, truth, Class1)
```

calib_class_thresh	<i>Calibrate class thresholds</i>
--------------------	-----------------------------------

Description

Predict for a new dataset by using a simple ensemble. Predictions from individual models are combined according to fun

Usage

```
calib_class_thresh(object, class_thresh, metric_thresh = NULL)
```

Arguments

object	an <code>simple_ensemble</code> object
class_thresh	probability threshold used to convert probabilities into classes. It can be a number (between 0 and 1), or a character metric (currently "tss_max", "kap_max" or "sensitivity"). For sensitivity, an additional target value is passed along as a second element of a vector, e.g. <code>c("sensitivity",0.8)</code> .
metric_thresh	a vector of length 2 giving a metric and its threshold, which will be used to prune which models in the ensemble will be used for the prediction. The 'metrics' need to have been computed when the workflow was tuned. The metric's threshold needs to match the value used during prediction. Examples are <code>c("accuracy",0.8)</code> or <code>c("boyce_cont",0.7)</code> .

Value

a `simple_ensemble` object

Examples

```
test_ens <- simple_ensemble() %>%
  add_member(two_class_res[1:3, ], metric = "roc_auc")
test_ens <- calib_class_thresh(test_ens, class_thresh = "tss_max")
test_ens <- calib_class_thresh(test_ens, class_thresh = "kap_max")
test_ens <- calib_class_thresh(test_ens, class_thresh = c("sens", 0.9))
```

check_sdm_presence	<i>Check that the column with presences is correctly formatted</i>
--------------------	--

Description

In `tidysdm`, the string defining presences should be the first level of the response factor. This function checks that the column is correctly formatted.

Usage

```
check_sdm_presence(.data, .col, presence_level = "presence")
```

Arguments

<code>.data</code>	a <code>data.frame</code> or <code>tibble</code> , or a derived object such as an <code>sf data.frame</code> , or a factor (e.g. the column with the response variable)
<code>.col</code>	the column containing the presences
<code>presence_level</code>	the string used to define the presence level of <code>.col</code>

Value

TRUE if correctly formatted

check_splits_balance	<i>Check the balance of presences vs pseudoabsences among splits</i>
----------------------	--

Description

Check the balance of presences vs pseudoabsences among splits

Usage

```
check_splits_balance(splits, .col)
```

Arguments

<code>splits</code>	the data splits (an <code>rset</code> or <code>split</code> object), generated by a function such as <code>spatialsample::spatial_block</code>
<code>.col</code>	the column containing the presences

Value

a `tibble` of the number of presences and pseudoabsences in the assessment and analysis set of each split (or training and testing in an initial split)

Examples

```
lacerta_thin <- readRDS(system.file("extdata/lacerta_thin_all_vars.rds",
  package = "tidysdm"
))
lacerta_cv <- spatial_block_cv(lacerta_thin, v = 5)
check_splits_balance(lacerta_cv, class)
```

clamp_predictors	<i>Clamp the predictors to match values in training set</i>
------------------	---

Description

This function clamps the environmental variables in a [terra::SpatRaster](#) or [terra::SpatRasterDataset](#) so that their minimum and maximum values do not exceed the range in the training dataset.

Usage

```
clamp_predictors(x, training, .col, use_na)

## Default S3 method:
clamp_predictors(x, training, .col, use_na)

## S3 method for class 'stars'
clamp_predictors(x, ...)

## S3 method for class 'SpatRaster'
clamp_predictors(x, training, .col, use_na = FALSE)

## S3 method for class 'SpatRasterDataset'
clamp_predictors(x, training, .col, use_na = FALSE)
```

Arguments

x	a terra::SpatRaster , stars or terra::SpatRasterDataset to clamp.
training	the training dataset (a data.frame or a sf::sf object).
.col	the column containing the presences (optional). If specified, it is excluded from the clamping.
use_na	a boolean determining whether values outside the range of the training dataset are removed (set to NA). If FALSE (the default), values outside the training range are replaced with the extremes of the training range.
...	additional arguments specific to a given object type

Value

a [terra::SpatRaster](#) or [terra::SpatRasterDataset](#) clamped to the ranges in training

```
collect_metrics.simple_ensemble
```

Obtain and format results produced by tuning functions for ensemble objects

Description

Return a tibble of performance metrics for all models.

Usage

```
## S3 method for class 'simple_ensemble'  
collect_metrics(x, ...)  
  
## S3 method for class 'repeat_ensemble'  
collect_metrics(x, ...)
```

Arguments

x	A simple_ensemble or repeat_ensemble object
...	Not currently used.

Details

When applied to a ensemble, the metrics that are returned do not contain the actual tuning parameter columns and values (unlike when these collect functions are run on other objects). The reason is that ensembles contain different types of models or models with different tuning parameters.

Value

A tibble.

See Also

[tune::collect_metrics\(\)](#)

Examples

```
collect_metrics(lacerta_ensemble)  
collect_metrics(lacerta_rep_ens)
```

control_ensemble_grid *Control wrappers*

Description

Supply these light wrappers as the `control` argument in a `tune::tune_grid()`, `tune::tune_bayes()`, or `tune::fit_resamples()` call to return the needed elements for use in an ensemble. These functions will return the appropriate control grid to ensure that assessment set predictions and information on model specifications and preprocessors, is supplied in the resampling results object!

To integrate ensemble settings with your existing control settings, note that these functions just call the appropriate `tune::control_*` function with the arguments `save_pred = TRUE`, `save_workflow = TRUE`.

These wrappers are equivalent to the ones used in the `stacks` package.

Usage

```
control_ensemble_grid()
```

```
control_ensemble_resamples()
```

```
control_ensemble_bayes()
```

Value

A `tune::control_grid`, `tune::control_bayes`, or `tune::control_resamples` object.

See Also

See the vignettes for examples of these functions used in context.

dist_pres_vs_bg	<i>Distance between the distribution of climate values for presences vs background</i>
-----------------	--

Description

For each environmental variable, this function computes the density functions of presences and absences and returns (1-overlap), which is a measure of the distance between the two distributions. Variables with a high distance are good candidates for SDMs, as species occurrences are confined to a subset of the available background.

Usage

```
dist_pres_vs_bg(.data, .col)
```

Arguments

<code>.data</code>	a <code>data.frame</code> (or derived object, such as <code>tibble</code> , or <code>sf</code>) with values for the bioclimate variables for presences and background
<code>.col</code>	the column containing the presences; it assumes presences to be the first level of this factor

Value

a name vector of distances

Examples

```
# This should be updated to use a dataset from tidysdm
data("bradypus", package = "maxnet")
bradypus_tb <- tibble::as_tibble(bradypus) %>%
  dplyr::mutate(presence = relevel(
    factor(
      dplyr::case_match(presence, 1 ~ "presence", 0 ~ "absence")
    ),
    ref = "presence"
  )) %>%
  select(-ecoreg)

bradypus_tb %>% dist_pres_vs_bg(presence)
```

explain_tidysdm

Create explainer from your tidysdm ensembles.

Description

DALEX is designed to explore and explain the behaviour of Machine Learning methods. This function creates a DALEX explainer (see [DALEX::explain\(\)](#)), which can then be queried by multiple functions from the DALEX package to create explanations of the model.

Usage

```
explain_tidysdm(
  model,
  data,
  y,
  predict_function,
  predict_function_target_column,
  residual_function,
  ...,
  label,
  verbose,
  precalculate,
```

```
    colorize,  
    model_info,  
    type,  
    by_workflow  
  )  
  
## Default S3 method:  
explain_tidysdm(  
  model,  
  data = NULL,  
  y = NULL,  
  predict_function = NULL,  
  predict_function_target_column = NULL,  
  residual_function = NULL,  
  ...,  
  label = NULL,  
  verbose = TRUE,  
  precalculate = TRUE,  
  colorize = !isTRUE(getOption("knitr.in.progress")),  
  model_info = NULL,  
  type = "classification",  
  by_workflow = FALSE  
)  
  
## S3 method for class 'simple_ensemble'  
explain_tidysdm(  
  model,  
  data = NULL,  
  y = NULL,  
  predict_function = NULL,  
  predict_function_target_column = NULL,  
  residual_function = NULL,  
  ...,  
  label = NULL,  
  verbose = TRUE,  
  precalculate = TRUE,  
  colorize = !isTRUE(getOption("knitr.in.progress")),  
  model_info = NULL,  
  type = "classification",  
  by_workflow = FALSE  
)  
  
## S3 method for class 'repeat_ensemble'  
explain_tidysdm(  
  model,  
  data = NULL,  
  y = NULL,  
  predict_function = NULL,
```



```

predict_function_target_column = NULL,
residual_function = NULL,
...,
label = NULL,
verbose = TRUE,
precalculate = TRUE,
colorize = !isTRUE(getOption("knitr.in.progress")),
model_info = NULL,
type = "classification",
by_workflow = FALSE
)

```

Arguments

model	object - a model to be explained
data	data.frame or matrix - data which will be used to calculate the explanations. If not provided, then it will be extracted from the model. Data should be passed without a target column (this shall be provided as the y argument). NOTE: If the target variable is present in the data, some of the functionalities may not work properly.
y	numeric vector with outputs/scores. If provided, then it shall have the same size as data
predict_function	function that takes two arguments: model and new data and returns a numeric vector with predictions. By default it is yhat.
predict_function_target_column	Character or numeric containing either column name or column number in the model prediction object of the class that should be considered as positive (i.e. the class that is associated with probability 1). If NULL, the second column of the output will be taken for binary classification. For a multiclass classification setting, that parameter cause switch to binary classification mode with one vs others probabilities.
residual_function	function that takes four arguments: model, data, target vector y and predict function (optionally). It should return a numeric vector with model residuals for given data. If not provided, response residuals ($y - \hat{y}$) are calculated. By default it is residual_function_default.
...	other parameters
label	character - the name of the model. By default it's extracted from the 'class' attribute of the model
verbose	logical. If TRUE (default) then diagnostic messages will be printed
precalculate	logical. If TRUE (default) then predicted_values and residual are calculated when explainer is created. This will happen also if verbose is TRUE. Set both verbose and precalculate to FALSE to omit calculations.
colorize	logical. If TRUE (default) then WARNINGS, ERRORS and NOTES are colored. Will work only in the R console. Now by default it is FALSE while knitting and TRUE otherwise.

model_info	a named list (package, version, type) containing information about model. If NULL, DALEX will seek for information on it's own.
type	type of a model, either classification or regression. If not specified then type will be extracted from model_info.
by_workflow	boolean determining whether a list of explainer, one per model, should be returned instead of a single explainer for the ensemble

Details

By default, the response variable is extracted from the ensemble object. Note that, if the response variable is passed directly, y should be a factor with presence as a reference level. To check that y is formatted correctly, use `check_sdm_presence()`.

Value

explainer object `DALEX::explain` ready to work with DALEX

Examples

```
# using the whole ensemble
lacerta_explainer <- explain_tidysdm(tidysdm::lacerta_ensemble)
# by workflow
explainer_list <- explain_tidysdm(tidysdm::lacerta_ensemble,
  by_workflow = TRUE
)
```

extrapol_mess	<i>Multivariate environmental similarity surfaces (MESS)</i>
---------------	--

Description

Compute multivariate environmental similarity surfaces (MESS), as described by Elith et al., 2010.

Usage

```
extrapol_mess(x, training, .col, ...)

## Default S3 method:
extrapol_mess(x, training, ...)

## S3 method for class 'stars'
extrapol_mess(x, ...)

## S3 method for class 'SpatRaster'
```

```
extrapol_mess(x, training, .col, filename = "", ...)
```

```
## S3 method for class 'data.frame'
```

```
extrapol_mess(x, training, .col, ...)
```

```
## S3 method for class 'SpatRasterDataset'
```

```
extrapol_mess(x, training, .col, ...)
```

Arguments

x	<code>terra::SpatRaster</code> , stars, <code>terra::SpatRasterDataset</code> or <code>data.frame</code>
training	matrix or data.frame or sf object containing the reference values; each column should correspond to one layer of the <code>terra::SpatRaster</code> object, with the exception of the presences column defined in <code>.col</code> (optional).
.col	the column containing the presences (optional). If specified, it is excluded when computing the MESS scores.
...	additional arguments as for <code>terra::writeRaster()</code>
filename	character. Output filename (optional)

Details

This function is a modified version of `mess` in package `predicts`, with a method added to work on `terra::SpatRasterDataset`. Note that the method for `terra::SpatRasterDataset` assumes that each variables is stored as a `terra::SpatRaster` with time information within `x`. Time is also assumed to be in years. If these conditions are not met, it is possible to manually extract a `terra::SpatRaster` for each time step, and use `extrapol_mess` on those `terra::SpatRasters`

Value

a `terra::SpatRaster` (data.frame) with the MESS values.

Author(s)

Jean-Pierre Rossi, Robert Hijmans, Paulo van Breugel, Andrea Manica

References

Elith J., M. Kearney M., and S. Phillips, 2010. The art of modelling range-shifting species. *Methods in Ecology and Evolution* 1:330-342.

filter_collinear	<i>Filter to retain only variables that have low collinearity</i>
------------------	---

Description

This method finds a subset of variables that have low collinearity. It provides three methods: `cor_caret`, a stepwise approach to remove variables with a pairwise correlation above a given cutoff, choosing the variable with the greatest mean correlation (based on the algorithm in `caret::findCorrelation`); `vif_step`, a stepwise approach to remove variables with an variance inflation factor above a given cutoff (based on the algorithm in `usdm::vifstep`), and `vif_cor`, a stepwise approach that, at each step, find the pair of variables with the highest correlation above the cutoff and removes the one with the largest vif. such that all have a correlation below a certain cutoff. There are methods for `terra::SpatRaster`, `data.frame` and `matrix`. For `terra::SpatRaster` and `data.frame`, only numeric variables will be considered.

Usage

```
filter_collinear(
  x,
  cutoff = NULL,
  verbose = FALSE,
  names = TRUE,
  to_keep = NULL,
  method = "cor_caret",
  cor_type = "pearson",
  max_cells = Inf,
  ...
)

## Default S3 method:
filter_collinear(
  x,
  cutoff = NULL,
  verbose = FALSE,
  names = TRUE,
  to_keep = NULL,
  method = "cor_caret",
  cor_type = "pearson",
  max_cells = Inf,
  ...
)

## S3 method for class 'stars'
filter_collinear(
  x,
  cutoff = NULL,
  verbose = FALSE,
```

```
names = TRUE,
to_keep = NULL,
method = "cor_caret",
cor_type = "pearson",
max_cells = Inf,
exhaustive = FALSE,
...
)

## S3 method for class 'SpatRaster'
filter_collinear(
  x,
  cutoff = NULL,
  verbose = FALSE,
  names = TRUE,
  to_keep = NULL,
  method = "cor_caret",
  cor_type = "pearson",
  max_cells = Inf,
  exhaustive = FALSE,
  ...
)

## S3 method for class 'data.frame'
filter_collinear(
  x,
  cutoff = NULL,
  verbose = FALSE,
  names = TRUE,
  to_keep = NULL,
  method = "cor_caret",
  cor_type = "pearson",
  max_cells = Inf,
  ...
)

## S3 method for class 'matrix'
filter_collinear(
  x,
  cutoff = NULL,
  verbose = FALSE,
  names = TRUE,
  to_keep = NULL,
  method = "cor_caret",
  cor_type = "pearson",
  max_cells = Inf,
  ...
)
```

Arguments

x	A terra::SpatRaster or stars object, a data.frame (with only numeric variables)
cutoff	A numeric value used as a threshold to remove variables. For, "cor_caret" and "vif_cor", it is the pair-wise absolute correlation cutoff, which defaults to 0.7. For "vif_step", it is the variable inflation factor, which defaults to 10
verbose	A boolean whether additional information should be provided on the screen
names	a logical; should the column names be returned TRUE or the column index FALSE)?
to_keep	A vector of variable names that we want to force in the set (note that the function will return an error if the correlation among any of those variables is higher than the cutoff).
method	character. One of "cor_caret", "vif_cor" or "vif_step".
cor_type	character. For methods that use correlation, which type of correlation: "pearson", "kendall", or "spearman". Defaults to "pearson"
max_cells	positive integer. The maximum number of cells to be used. If this is smaller than ncell(x), a regular sample of x is used
...	additional arguments specific to a given object type
exhaustive	boolean. Used only for terra::SpatRaster when downsampling to max_cells, if we require the exhaustive approach in terra::spatSample() . This is only needed for rasters that are very sparse and not too large, see the help page of terra::spatSample() for details.

Value

A vector of names of columns that are below the correlation threshold (when names = TRUE), otherwise a vector of indices. Note that the indices are only for numeric variables (i.e. if factors are present, the indices do not take them into account).

Author(s)

for cor_caret: Original R code by Dong Li, modified by Max Kuhn and Andrea Manica; for vif_step and vif_cor, original algorithm by Babak Naimi, rewritten by Andrea Manica for tidysdm

References

Naimi, B., Hamm, N.A.S., Groen, T.A., Skidmore, A.K., and Toxopeus, A.G. 2014. Where is positional uncertainty a problem for species distribution modelling?, *Ecography* 37 (2): 191-203.

filter_high_cor	<i>Deprecated: Filter to retain only variables below a given correlation threshold</i>
-----------------	--

Description

THIS FUNCTION IS DEPRECATED. USE `filter_collinear` with `method=cor_caret` instead

Usage

```
filter_high_cor(x, cutoff = 0.7, verbose = FALSE, names = TRUE, to_keep = NULL)

## Default S3 method:
filter_high_cor(x, cutoff = 0.7, verbose = FALSE, names = TRUE, to_keep = NULL)

## S3 method for class 'SpatRaster'
filter_high_cor(x, cutoff = 0.7, verbose = FALSE, names = TRUE, to_keep = NULL)

## S3 method for class 'data.frame'
filter_high_cor(x, cutoff = 0.7, verbose = FALSE, names = TRUE, to_keep = NULL)

## S3 method for class 'matrix'
filter_high_cor(x, cutoff = 0.7, verbose = FALSE, names = TRUE, to_keep = NULL)
```

Arguments

x	A <code>terra::SpatRaster</code> object, a <code>data.frame</code> (with only numeric variables), or a correlation matrix
cutoff	A numeric value for the pair-wise absolute correlation cutoff
verbose	A boolean for printing the details
names	a logical; should the column names be returned TRUE or the column index FALSE)?
to_keep	A vector of variable names that we want to force in the set (note that the function will return an error if the correlation among any of those variables is higher than the cutoff).

Details

This method finds a subset of variable such that all have a correlation below a certain cutoff. There are methods for `terra::SpatRaster`, `data.frame`, and to work directly on a correlation matrix that was previously estimated. For `data.frame`, only numeric variables will be considered. The algorithm is based on `caret::findCorrelation`, using the `exact` option. The absolute values of pair-wise correlations are considered. If two variables have a high correlation, the function looks at the mean absolute correlation of each variable and removes the variable with the largest mean absolute correlation.

There are several function in the package `subselect` that can also be used to accomplish the same goal but tend to retain more predictors.

Value

A vector of names of columns that are below the correlation threshold (when `names = TRUE`), otherwise a vector of indices. Note that the indices are only for numeric variables (i.e. if factors are present, the indices do not take them into account).

gam_formula	<i>Create a formula for gam</i>
-------------	---------------------------------

Description

This function takes the formula from a recipe, and turns numeric predictors into smooths with a given `k`. This formula can be passed to a workflow or workflow set when fitting a gam.

Usage

```
gam_formula(object, k = 10)
```

Arguments

object	a recipes::recipe , already trained
k	the k value for the smooth

Value

a formula

geom_split_violin	<i>Split violin geometry for ggplots</i>
-------------------	--

Description

This geometry displays the density distribution of two groups side by side, as two halves of a violin. Note that an `emptyx` aesthetic has to be provided even if you want to plot a single variable (see example below).

Usage

```
geom_split_violin(
  mapping = NULL,
  data = NULL,
  stat = "ydensity",
  position = "identity",
  nudge = 0,
  ...,
  draw_quantiles = NULL,
```



```

    trim = TRUE,
    scale = "area",
    na.rm = FALSE,
    show.legend = NA,
    inherit.aes = TRUE
  )

```

Arguments

mapping	Set of aesthetic mappings created by aes() . If specified and <code>inherit.aes = TRUE</code> (the default), it is combined with the default mapping at the top level of the plot. You must supply mapping if there is no plot mapping.
data	The data to be displayed in this layer. There are three options: If <code>NULL</code>, the default, the data is inherited from the plot data as specified in the call to ggplot(). A <code>data.frame</code>, or other object, will override the plot data. All objects will be fortified to produce a data frame. See fortify() for which variables will be created. A function will be called with a single argument, the plot data. The return value must be a <code>data.frame</code>, and will be used as the layer data. A function can be created from a formula (e.g. <code>~ head(.x, 10)</code>).
stat	Use to override the default connection between ggplot2::geom_violin() and ggplot2::stat_ydensity() .
position	A position adjustment to use on the data for this layer. This can be used in various ways, including to prevent overplotting and improving the display. The position argument accepts the following: • The result of calling a position function, such as position_jitter(). This method allows for passing extra arguments to the position. • A string naming the position adjustment. To give the position as a string, strip the function name of the <code>position_</code> prefix. For example, to use position_jitter(), give the position as "jitter". • For more information and other ways to specify the position, see the layer position documentation.
nudge	Add space between the half-violin and the middle of the space allotted to a given factor on the x-axis.
...	Other arguments passed on to layer()'s <code>params</code> argument. These arguments broadly fall into one of 4 categories below. Notably, further arguments to the position argument, or aesthetics that are required can <i>not</i> be passed through Unknown arguments that are not part of the 4 categories below are ignored. • Static aesthetics that are not mapped to a scale, but are at a fixed value and apply to the layer as a whole. For example, <code>colour = "red"</code> or <code>linewidth = 3</code>. The geom's documentation has an Aesthetics section that lists the available options. The 'required' aesthetics cannot be passed on to the <code>params</code>. Please note that while passing unmapped aesthetics as vectors is technically possible, the order and required length is not guaranteed to be parallel to the input data.

- When constructing a layer using a `stat_*()` function, the `...` argument can be used to pass on parameters to the `geom` part of the layer. An example of this is `stat_density(geom = "area", outline.type = "both")`. The `geom`'s documentation lists which parameters it can accept.
- Inversely, when constructing a layer using a `geom_*()` function, the `...` argument can be used to pass on parameters to the `stat` part of the layer. An example of this is `geom_area(stat = "density", adjust = 0.5)`. The `stat`'s documentation lists which parameters it can accept.
- The `key_glyph` argument of `layer()` may also be passed on through `...`. This can be one of the functions described as [key glyphs](#), to change the display of the layer in the legend.

<code>draw_quantiles</code>	[Deprecated] Previous specification of drawing quantiles.
<code>trim</code>	If TRUE (default), trim the tails of the violins to the range of the data. If FALSE, don't trim the tails.
<code>scale</code>	if "area" (default), all violins have the same area (before trimming the tails). If "count", areas are scaled proportionally to the number of observations. If "width", all violins have the same maximum width.
<code>na.rm</code>	If FALSE, the default, missing values are removed with a warning. If TRUE, missing values are silently removed.
<code>show.legend</code>	logical. Should this layer be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes. It can also be a named logical vector to finely select the aesthetics to display. To include legend keys for all levels, even when no data exists, use TRUE. If NA, all levels are shown in legend, but unobserved levels are omitted.
<code>inherit.aes</code>	If FALSE, overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn't inherit behaviour from the default plot specification, e.g. <code>ggplot2::annotation_borders</code> .

Details

The implementation is based on <https://stackoverflow.com/questions/35717353/split-violin-plot-with-ggplot2>. Credit goes to @jan-jlx for providing a complete implementation on StackOverflow, and to Trang Q. Nguyen for adding the `nudge` parameter.

Value

a `ggplot2::layer` object

Examples

```
data("bradypus", package = "maxnet")
bradypus_tb <- tibble::as_tibble(bradypus) %>%
  dplyr::mutate(presence = relevel(
    factor(
      dplyr::case_match(
        presence, 1 ~ "presence",
        0 ~ "absence"
      )
    )
  )
```

```

    ),
    ref = "presence"
  ))

ggplot(bradypus_tb, aes(
  x = "",
  y = cld6190_ann,
  fill = presence
)) +
  geom_split_violin(nudge = 0.01)

```

grid_cellsize	<i>Get default grid cellsize for a given dataset</i>
---------------	--

Description

This function facilitates using `spatialsample::spatial_block_cv` multiple times in an analysis. `spatialsample::spatial_block_cv` creates a grid based on the object in data. However, if spatial blocks are generated multiple times in an analysis (e.g. for a `spatial_initial_split()`, and then subsequently for cross-validation on the training dataset), it might be desirable to keep the same grid). By applying this function to the largest dataset, usually the full dataset before `spatial_initial_split()`. The resulting cellsize can be used as an option in `spatialsample::spatial_block_cv`.

Usage

```
grid_cellsize(data, n = c(10, 10))
```

Arguments

data	a <code>sf::sf</code> dataset used to size the grid
n	the number of cells in the grid, defaults to <code>c(10,10)</code> , which is also the default for <code>sf::st_make_grid()</code>

Value

the cell size

grid_offset	<i>Get default grid cellsize for a given dataset</i>
-------------	--

Description

This function facilitates using `spatialsample::spatial_block_cv` multiple times in an analysis. `spatialsample::spatial_block_cv` creates a grid based on the object in data. However, if spatial blocks are generated multiple times in an analysis (e.g. for a `spatial_initial_split()`, and then subsequently for cross-validation on the training dataset), it might be desirable to keep the same grid). By applying this function to the largest dataset, usually the full dataset before `spatial_initial_split()`. The resulting cellsize can be used as an option in `spatialsample::spatial_block_cv`.

Usage

```
grid_offset(data)
```

Arguments

data a `sf::sf` dataset used to size the grid

Value

the grid offset

horses	<i>Coordinates of radiocarbon dates for horses</i>
--------	--

Description

Coordinates for presences of horses from 22k to 8k YBP.

Usage

```
horses
```

Format

An tibble with 1,297 rows and 3 variables:

latitude latitudes in degrees

longitude longitudes in degrees

time_bp time in years before present

kap_max	<i>Maximum Cohen's Kappa</i>
---------	------------------------------

Description

Cohen's Kappa (`yardstick::kap()`) is a measure similar to `yardstick::accuracy()`, but it normalises the observed accuracy by the value that would be expected by chance (this helps for unbalanced cases when one class is predominant).

Usage

```
kap_max(data, ...)

## S3 method for class 'data.frame'
kap_max(
  data,
  truth,
  ...,
  estimator = NULL,
  na_rm = TRUE,
  event_level = "first",
  case_weights = NULL
)

## S3 method for class 'sf'
kap_max(data, ...)

kap_max_vec(
  truth,
  estimate,
  estimator = NULL,
  na_rm = TRUE,
  event_level = "first",
  case_weights = NULL,
  ...
)
```

Arguments

<code>data</code>	Either a <code>data.frame</code> containing the columns specified by the <code>truth</code> and <code>estimate</code> arguments, or a table/matrix where the true class results should be in the columns of the table.
<code>...</code>	A set of unquoted column names or one or more <code>dplyr</code> selector functions to choose which variables contain the class probabilities. If <code>truth</code> is binary, only 1 column should be selected, and it should correspond to the value of <code>event_level</code> . Otherwise, there should be as many columns as factor levels of <code>truth</code> and the ordering of the columns should be the same as the factor levels of <code>truth</code> .

truth	The column identifier for the true class results (that is a factor). This should be an unquoted column name although this argument is passed by expression and supports quasiquotation (you can unquote column names). For <code>_vec()</code> functions, a factor vector.
estimator	One of "binary", "hand_till", "macro", or "macro_weighted" to specify the type of averaging to be done. "binary" is only relevant for the two class case. The others are general methods for calculating multiclass metrics. The default will automatically choose "binary" if truth is binary, "hand_till" if truth has >2 levels and case_weights isn't specified, or "macro" if truth has >2 levels and case_weights is specified (in which case "hand_till" isn't well-defined).
na_rm	A logical value indicating whether NA values should be stripped before the computation proceeds.
event_level	A single string. Either "first" or "second" to specify which level of truth to consider as the "event". This argument is only applicable when estimator = "binary". The default uses an internal helper that generally defaults to "first"
case_weights	The optional column identifier for case weights. This should be an unquoted column name that evaluates to a numeric column in data. For <code>_vec()</code> functions, a numeric vector.
estimate	If truth is binary, a numeric vector of class probabilities corresponding to the "relevant" class. Otherwise, a matrix with as many columns as factor levels of truth. It is assumed that these are in the same order as the levels of truth.

Details

This function calibrates the probability threshold to classify presences to maximises kappa.

There is no multiclass version of this function, it only operates on binary predictions (e.g. presences and absences in SDMs).

Value

A tibble with columns `.metric`, `.estimator`, and `.estimate` and 1 row of values. For grouped data frames, the number of rows returned will be the same as the number of groups.

References

Cohen, J. (1960). "A coefficient of agreement for nominal scales". *Educational and Psychological Measurement*. 20 (1): 37-46.

Cohen, J. (1968). "Weighted kappa: Nominal scale agreement provision for scaled disagreement or partial credit". *Psychological Bulletin*. 70 (4): 213-220.

See Also

Other class probability metrics: [boyce_cont\(\)](#), [tss_max\(\)](#)

Examples

```
kap_max(two_class_example, truth, Class1)
```

km2m

Convert a geographic distance from km to m

Description

This function takes distance in km and converts it into meters, the units generally used by geographic operations in R. This is a trivial conversion, but this functions ensures that no zeroes are lost along the way!

Usage

```
km2m(x)
```

Arguments

x the number of km

Value

the number of meters

Examples

```
km2m(10000)
km2m(1)
```

lacerta

Coordinates of presences for Iberian emerald lizard

Description

Coordinates for presences of *Lacerta schreiberi*. The variables are as follows:

Usage

```
lacerta
```

Format

An tibble with 1,297 rows and 3 variables:

ID ids from GBIF

latitude latitudes in degrees

longitude longitudes in degrees

lacerta_ensemble	<i>A simple ensemble for the lacerta data</i>
------------------	---

Description

Ensemble SDM for *Lacerta schreiberi*, as generated in the vignette.

Usage

```
lacerta_ensemble
```

Format

A `simple_ensemble` object

lacerta_rep_ens	<i>A repeat ensemble for the lacerta data</i>
-----------------	---

Description

Ensemble SDM for *Lacerta schreiberi*, as generated in the vignette.

Usage

```
lacerta_rep_ens
```

Format

A `repeat_ensemble` object

lacertidae_background	<i>Coordinates of presences for lacertidae in the Iberian peninsula</i>
-----------------------	---

Description

Coordinates for presences of lacertidae, used as background for the `lacerta` dataset.. The variables are as follows:

Usage

```
lacertidae_background
```


Format

An tibble with 1,297 rows and 3 variables:

ID ids from GBIF

latitude latitudes in degrees

longitude longitudes in degrees

```
make_mask_from_presence
```

Make a mask from presence data

Description

This functions uses the presence column to create a mask to apply to the raster to define the area of interest. Two methods are available: one that uses a buffer around each presence, and one that create a convex hull around all presences (with the possibility of further adding a buffer around the hull).

Usage

```
make_mask_from_presence(data, method = "buffer", buffer = 0, return_sf = FALSE)
```

Arguments

data	An <code>sf::sf</code> data frame of presences..
method	the method to use to create the mask. Either 'buffer' or 'convex_hull'
buffer	the buffer to add around each presence (in the units of the crs of the data; for lat/lon, the buffer will be in meters), or around the convex hull (if method is 'convex_hull')
return_sf	whether to return the mask as an sf object (if TRUE) or as a <code>terra::SpatVector</code> object (if FALSE, default)

Details

To use `terra::mask()` on a raster, use `return_sf = FALSE` to get a `terra::SpatVector` object that can be used for masking.

Value

a `terra::SpatVector` or an `sf` object (depending on the value of `return_sf`) with the mask

Examples

```
lacerta_sf <- lacerta %>%
  sf::st_as_sf(coords = c("longitude", "latitude")) %>%
  sf::st_set_crs(4326)
land_mask <- terra::readRDS(system.file("extdata/lacerta_land_mask.rds",
  package = "tidysdm"
))
mask_buffer <- make_mask_from_presence(lacerta_sf,
  method = "buffer",
  buffer = 60000
)
terra::plot(terra::mask(land_mask, mask_buffer))
mask_ch <- make_mask_from_presence(lacerta_sf, method = "convex_hull")
terra::plot(terra::mask(land_mask, mask_ch))
```

maxent

MaxEnt model

Description

maxent defines the MaxEnt model as used in Species Distribution Models. A good guide to how options of a MaxEnt model work can be found in <https://onlinelibrary.wiley.com/doi/full/10.1111/j.1600-0587.2013.07872.x>

Usage

```
maxent(
  mode = "classification",
  engine = "maxnet",
  feature_classes = NULL,
  regularization_multiplier = NULL
)
```

Arguments

mode	A single character string for the type of model. The only possible value for this model is "classification".
engine	A single character string specifying what computational engine to use for fitting. Currently only "maxnet" is available.
feature_classes	character, continuous feature classes desired, either "default" or any subset of "lqpht" (for example, "lh")
regularization_multiplier	numeric, a constant to adjust regularization

Value

a `parsnip::model_spec` for a maxent model

Examples

```
# format the data
data("bradypus", package = "maxnet")
bradypus_tb <- tibble::as_tibble(bradypus) %>%
  dplyr::mutate(presence = relevel(
    factor(
      dplyr::case_match(presence, 1 ~ "presence", 0 ~ "absence")
    ),
    ref = "presence"
  )) %>%
  select(-ecoreg)

# fit the model, and make some predictions
maxent_spec <- maxent(feature_classes = "lq")
maxent_fitted <- maxent_spec %>%
  fit(presence ~ ., data = bradypus_tb)
pred_prob <- predict(maxent_fitted,
  new_data = bradypus[, -1],
  type = "prob"
)
pred_class <- predict(maxent_fitted,
  new_data = bradypus[, -1],
  type = "class"
)

# Now with tuning
maxent_spec <- maxent(
  regularization_multiplier = tune::tune(),
  feature_classes = tune::tune()
)
set.seed(452)
cv <- vfold_cv(bradypus_tb, v = 2)
maxent_tune_res <- maxent_spec %>%
  tune_grid(presence ~ ., cv, grid = 3)
show_best(maxent_tune_res, metric = "roc_auc")
```

maxent_params

*Parameters for maxent models***Description**

These parameters are auxiliary to MaxEnt models using the "maxnet" engine. These functions are used by the tuning functions, and the user will rarely access them directly.

Usage

```
regularization_multiplier(range = c(0.5, 3), trans = NULL)
```

```
feature_classes(values = c("l", "lq", "lqp", "lqph", "lqpht"))
```

Arguments

range	A two-element vector holding the defaults for the smallest and largest possible values, respectively. If a transformation is specified, these values should be in the transformed units.
trans	A trans object from the scales package, such as <code>scales::log10_trans()</code> or <code>scales::reciprocal_trans()</code> . If not provided, the default is used which matches the units used in range. If no transformation, NULL.
values	For <code>feature_classes()</code> , a character string of any subset of "lqpht" (for example, "lh")

Value

a param object that can be used for tuning.

Examples

```
regularization_multiplier()
feature_classes()
```

niche_overlap

Compute overlap metrics of the two niches

Description

This function computes overlap metrics between two rasters. It currently implements Schoener's D and the inverse I of Hellinger's distance.

Usage

```
niche_overlap(x, y, method = c("Schoener", "Hellinger"))
```

Arguments

x	a terra::SpatRaster with a single layer
y	a terra::SpatRaster with a single layer
method	a string (or vector of strings) taking values "Schoener" and "Hellinger"

Details

Note that Hellinger's distance is normalised by dividing by square root of 2 (which is the correct asymptote for Hellinger's D), rather than the incorrect 2 used originally in Warren et al (2008), based on the Erratum for that paper.

Value

a list of overlap metrics, with slots *D* and *I* (depending on method)

References

Warren, D.L., Glor, R.E. & Turelli M. (2008) Environmental niche equivalency versus conservatism: quantitative approaches to niche evolution. *Evolution* 62: 2868-2883

optim_thresh	<i>Find threshold that optimises a given metric</i>
--------------	---

Description

This function returns the threshold to turn probabilities into binary classes whilst optimising a given metric. Currently available for `tss_max`, `kap_max` and `sensitivity` (for which a target sensitivity is required).

Usage

```
optim_thresh(truth, estimate, metric, event_level = "first")
```

Arguments

<code>truth</code>	The column identifier for the true class results (that is a factor). This should be an unquoted column name although this argument is passed by expression and supports quasiquotation (you can unquote column names). For <code>_vec()</code> functions, a factor vector.
<code>estimate</code>	the predicted probability for the event
<code>metric</code>	character of metric to be optimised. Currently only "tss_max", "kap_max", and "sensitivity" with a given target (e.g. <code>c("sensitivity",0.8)</code>)
<code>event_level</code>	A single string. Either "first" or "second" to specify which level of truth to consider as the "event". This argument is only applicable when estimator = "binary". The default uses an internal helper that generally defaults to "first"

Value

the probability threshold for the event

Examples

```
optim_thresh(two_class_example$truth, two_class_example$Class1,
  metric = c("tss_max")
)
optim_thresh(two_class_example$truth, two_class_example$Class1,
  metric = c("sens", 0.9)
)
```

pairs,stars-method *Pairwise matrix of scatterplot for stars objects*

Description

Pairs plot of attributes for stars objects. This is equivalent to `terra::pairs()` but works with stars objects.

Usage

```
## S4 method for signature 'stars'
pairs(
  x,
  hist = TRUE,
  cor = TRUE,
  use = "pairwise.complete.obs",
  maxcells = 1e+05,
  ...
)
```

Arguments

<code>x</code>	SpatRaster
<code>hist</code>	logical. If TRUE a histogram of the values is shown on the diagonal
<code>cor</code>	logical. If TRUE the correlation coefficient is shown in the upper panels
<code>use</code>	argument passed to the <code>cor</code> function
<code>maxcells</code>	integer. Number of pixels to sample from each layer of a large SpatRaster
<code>...</code>	additional arguments (graphical parameters)

Value

a pairs plot of the attributes of the stars object.

Examples

```
r <- terra::rast(system.file("ex/elev.tif", package = "terra"))
s <- c(r, 1 / r, sqrt(r))
names(s) <- c("elevation", "inverse", "sqrt")
terra::pairs(s)
s_stars <- stars::st_as_stars(s, as_attributes = TRUE)
pairs(s_stars)
```

plot_pres_vs_bg	<i>Plot presences vs background</i>
-----------------	-------------------------------------

Description

Create a composite plots contrasting the distribution of multiple variables for presences vs the background.

Usage

```
plot_pres_vs_bg(.data, .col)
```

Arguments

.data	a <code>data.frame</code> (or derived object, such as <code>tibble::tibble</code> , or <code>sf::st_sf</code>) with values for the bioclimate variables for presences and background
.col	the column containing the presences; it assumes presences to be the first level of this factor

Value

a patchwork composite plot

Examples

```
data("bradypus", package = "maxnet")
bradypus_tb <- tibble::as_tibble(bradypus) %>%
  dplyr::mutate(presence = relevel(
    factor(
      dplyr::case_match(presence, 1 ~ "presence", 0 ~ "absence")
    ),
    ref = "presence"
  )) %>%
  select(-ecoreg)

bradypus_tb %>% plot_pres_vs_bg(presence)
```

predict.repeat_ensemble	<i>Predict for a repeat ensemble set</i>
-------------------------	--

Description

Predict for a new dataset by using a repeat ensemble. Predictions from individual models are combined according to fun

Usage

```
## S3 method for class 'repeat_ensemble'
predict(
  object,
  new_data,
  type = "prob",
  fun = "mean",
  metric_thresh = NULL,
  class_thresh = NULL,
  members = FALSE,
  ...
)
```

Arguments

<code>object</code>	an <code>repeat_ensemble</code> object
<code>new_data</code>	a data frame in which to look for variables with which to predict.
<code>type</code>	the type of prediction, "prob" or "class".
<code>fun</code>	string defining the aggregating function. It can take values <code>mean</code> , <code>median</code> , <code>weighted_mean</code> , <code>weighted_median</code> and <code>none</code> . It is possible to combine multiple functions, except for "none". If it is set to "none", only the individual member predictions are returned (this automatically sets <code>member</code> to <code>TRUE</code>)
<code>metric_thresh</code>	a vector of length 2 giving a metric and its threshold, which will be used to prune which models in the ensemble will be used for the prediction. The 'metrics' need to have been computed when the workflow was tuned. Examples are <code>c("accuracy",0.8)</code> or <code>c("boyce_cont",0.7)</code>
<code>class_thresh</code>	probability threshold used to convert probabilities into classes. It can be a number (between 0 and 1), or a character metric (currently "tss_max" or "sensitivity"). For sensitivity, an additional target value is passed along as a second element of a vector, e.g. <code>c("sensitivity",0.8)</code> .
<code>members</code>	boolean defining whether individual predictions for each member should be added to the ensemble prediction. The columns for individual members have the name of the workflow a a prefix, separated by "." from the usual column names of the predictions.
<code>...</code>	not used in this method.

Value

a tibble of predictions

predict.simple_ensemble

Predict for a simple ensemble set

Description

Predict for a new dataset by using a simple ensemble. Predictions from individual models (i.e. workflows) are combined according to fun

Usage

```
## S3 method for class 'simple_ensemble'
predict(
  object,
  new_data = NULL,
  type = "prob",
  fun = "mean",
  metric_thresh = NULL,
  class_thresh = NULL,
  members = FALSE,
  ...
)
```

Arguments

object	an simple_ensemble object
new_data	a data frame in which to look for variables with which to predict. If NULL, the predictors from the first workflow in the ensemble are used; note that this only makes sense if all workflows have the same predictors.
type	the type of prediction, "prob" or "class".
fun	string defining the aggregating function. It can take values mean, median, weighted_mean, weighted_median and none. It is possible to combine multiple functions, except for "none". If it is set to "none", only the individual member predictions are returned (this automatically sets member to TRUE)
metric_thresh	a vector of length 2 giving a metric and its threshold, which will be used to prune which models in the ensemble will be used for the prediction. The 'metrics' need to have been computed when the workflow was tuned. Examples are c("accuracy",0.8) or c("boyce_cont",0.7)
class_thresh	probability threshold used to convert probabilities into classes. It can be a number (between 0 and 1), or a character metric (currently "tss_max" or "sensitivity"). For sensitivity, an additional target value is passed along as a second element of a vector, e.g. c("sensitivity",0.8).
members	boolean defining whether individual predictions for each member should be added to the ensemble prediction. The columns for individual members have the name of the workflow a prefix, separated by "." from the usual column names of the predictions.

... not used in this method.

Value

a tibble of predictions

predict_raster	<i>Make predictions for a whole raster</i>
----------------	--

Description

This function allows to use a raster as data to make predictions from a variety of tidymodels objects, such as [simple_ensemble](#) or `stacks::stacks`

Usage

```
predict_raster(object, raster, ...)

## Default S3 method:
predict_raster(object, raster, filename = "", n = 4, ...)
```

Arguments

object	the tidymodels object of interest
raster	the terra::SpatRaster or stars with the input data. It has to include levels with the same names as the variables used in object
...	parameters to be passed to the standard <code>predict()</code> function for the appropriate object type (e.g. <code>metric_thresh</code> or <code>class_thresh</code>).
filename	the name of the output file raster file; this is only needed to save large rasters, and can be left blank for rasters that can be kept in memory.
n	positive integer indicating how many copies the data may be in memory at any point in time (it defaults to 4). This is used to determine whether rasters can be processed in one go, or in chunks. If you get an out of memory error, increase n. See terra::writeStart() for more details.

Value

a [terra::SpatRaster](#) (or stars if that is the input) with the predictions

prob_metrics_sf	<i>Probability metrics for sf objects</i>
-----------------	---

Description

tidysdm provides specialised metrics for SDMs, which have their own help pages([boyce_cont\(\)](#), [kap_max\(\)](#), and [tss_max\(\)](#)). Additionally, it also provides methods to handle [sf::sf](#) objects for the following standard yardstick metrics:

```
yardstick::average_precision()  
yardstick::brier_class()  
yardstick::classification_cost()  
yardstick::gain_capture()  
yardstick::mn_log_loss()  
yardstick::pr_auc()  
yardstick::roc_auc()  
yardstick::roc_aunp()  
yardstick::roc_aunu()
```

Usage

```
## S3 method for class 'sf'  
average_precision(data, ...)  
  
## S3 method for class 'sf'  
brier_class(data, ...)  
  
## S3 method for class 'sf'  
classification_cost(data, ...)  
  
## S3 method for class 'sf'  
gain_capture(data, ...)  
  
## S3 method for class 'sf'  
mn_log_loss(data, ...)  
  
## S3 method for class 'sf'  
pr_auc(data, ...)  
  
## S3 method for class 'sf'  
roc_auc(data, ...)  
  
## S3 method for class 'sf'  
roc_aunp(data, ...)
```

```
## S3 method for class 'sf'
roc_aunu(data, ...)
```

Arguments

data	an <code>sf::sf</code> object
...	any other parameters to pass to the <code>data.frame</code> version of the metric. See the specific man page for the metric of interest.

Details

Note that `roc_aunp` and `roc_aunu` are multiclass metrics, and as such are not relevant for SDMs (which work on a binary response). They are included for completeness, so that all class probability metrics from `yardstick` have an `sf` method, for applications other than SDMs.

Value

A tibble with columns `.metric`, `.estimator`, and `.estimate` and 1 row of values.

recipe.sf	<i>Recipe for sf objects</i>
-----------	------------------------------

Description

This method for `recipes::recipe()` handles the case when `x` is an `sf::sf` object, as commonly used in Species Distribution Model, and generates a `spatial_recipe`.

Usage

```
## S3 method for class 'sf'
recipe(x, ...)

spatial_recipe(x, ...)
```

Arguments

x	An <code>sf::sf</code> data frame.
...	parameters to be passed to <code>recipes::recipe()</code>

Details

`recipes::recipe()` are not natively compatible with `sf::sf` objects. The problem is that the geometry column of `sf::sf` objects is a list, which is incompatible with the translation of formulae in `recipes::recipe()`. This method strips the geometry column from the `data.frame` and replaces it with a simple X and Y columns before any further operations, thus allowing the usual processing by `recipes::recipe()` to succeed (X and Y are give the role of coords in a spatial recipe). When prepping and baking a `spatial_recipe`, if a `data.frame` or tibble without coordinates is used as `training` or `new_data`,

dummy X and Y columns are generated and filled with NAs. NOTE that order matters! You need to use the syntax `recipe(x=sf_obj, formula=class~.)` for the method to successfully detect the `sf::sf` object. Starting with `formula` will fail.

Value

An object of class `spatial_recipe`, which is a derived version of `recipes::recipe()`, see the manpage for `recipes::recipe()` for details.

repeat_ensemble	<i>Repeat ensemble</i>
-----------------	------------------------

Description

An ensemble based multiple sets of pseudoabsences/background. This object is a collection (list) of `simple_ensemble` objects for which predictions will be combined in a simple way (e.g. by taking either the mean or median). Each `simple_ensemble` contains the best version of a each given model type following turning; all simple ensembles will need to have the same metric estimated during the cv process.

Usage

```
repeat_ensemble(...)
```

Arguments

... not used, this function just creates an empty `repeat_ensemble` object. Members are added with `add_best_candidates()`

Value

an empty `repeat_ensemble`

sample_background	<i>Sample background points for SDM analysis</i>
-------------------	--

Description

This function samples background points from a raster given a set of presences. The locations returned as the center points of the sampled cells, which can overlap with the presences (in contrast to pseudo-absences, see `sample_pseudoabs`). The following methods are implemented:

- 'random': background randomly sampled from the region covered by the raster (i.e. not NAs).
- 'dist_max': background randomly sampled from the unioned buffers of 'dist_max' from presences (distances in 'm' for lonlat rasters, and in map units for projected rasters). Using the union of buffers means that areas that are in multiple buffers are not oversampled. This is also referred to as "thickening".
- 'bias': background points are sampled according to a surface representing the biased sampling effort.

Usage

```
sample_background(
  data,
  raster,
  n,
  coords = NULL,
  method = "random",
  class_label = "background",
  return_pres = TRUE
)
```

Arguments

data	An <code>sf::sf</code> data frame, or a data frame with coordinate variables. These can be defined in <code>coords</code> , unless they have standard names (see details below).
raster	the <code>terra::SpatRaster</code> or stars from which cells will be sampled (the first layer will be used to determine which cells are NAs, and thus can not be sampled). If sampling is "bias", then the sampling probability will be proportional to the values on the first layer (i.e. band) of the raster.
n	number of background points to sample.
coords	a vector of length two giving the names of the "x" and "y" coordinates, as found in data. If left to <code>NULL</code> , the function will try to guess the columns based on standard names <code>c("x", "y")</code> , <code>c("X", "Y")</code> , <code>c("longitude", "latitude")</code> , or <code>c("lon", "lat")</code> .
method	sampling method. One of 'random', 'dist_max', and 'bias'. For <code>dist_max</code> , the maximum distance is set as an additional element of a vector, e.g <code>c('dist_max', 70000)</code> .
class_label	the label given to the sampled points. Defaults to background
return_pres	return presences together with background in a single tibble.

Details

Note that the units of distance depend on the projection of the raster.

Value

An object of class `tibble::tibble`. If presences are returned, the presence level is set as the reference (to match the expectations in the `yardstick` package that considers the first level to be the event).

sample_background_time

Sample background points for SDM analysis for points with a time point.

Description

This function samples background points from a raster given a set of presences. The locations returned are the center points of the sampled cells, which can overlap with the presences (in contrast to pseudo-absences, see [sample_pseudoabs_time](#)). The following methods are implemented:

- 'random': background points randomly sampled from the region covered by the raster (i.e. not NAs).
- 'dist_max': background points randomly sampled from the unioned buffers of 'dist_max' from presences (distances in 'm' for lonlat rasters, and in map units for projected rasters). Using the union of buffers means that areas that are in multiple buffers are not oversampled. This is also referred to as "thickening".
- 'bias': background points are sampled according to a surface representing the biased sampling effort. Note that the surface for each time step is normalised to sum to 1; use `n_per_time_step` to affect sampling effort within each time step.

Usage

```
sample_background_time(
  data,
  raster,
  n_per_time_step,
  coords = NULL,
  time_col = "time",
  lubridate_fun = c,
  method = "random",
  class_label = "background",
  return_pres = TRUE,
  time_buffer = 0
)
```

Arguments

<code>data</code>	An <code>sf::sf</code> data frame, or a data frame with coordinate variables. These can be defined in <code>coords</code> , unless they have standard names (see details below).
<code>raster</code>	the <code>terra::SpatRaster</code> , stars or <code>terra::SpatRasterDataset</code> from which cells will be sampled. If a <code>terra::SpatRasterDataset</code> , the first dataset will be used to define which cells are valid, and which are NAs.
<code>n_per_time_step</code>	number of background points to sample for each time step (i.e. a vector of length equal to the number of time steps in raster)
<code>coords</code>	a vector of length two giving the names of the "x" and "y" coordinates, as found in data. If left to NULL, the function will try to guess the columns based on standard names <code>c("x", "y")</code> , <code>c("X", "Y")</code> , <code>c("longitude", "latitude")</code> , or <code>c("lon", "lat")</code>
<code>time_col</code>	The name of the column with time; if time is not a lubridate object, use <code>lubridate_fun</code> to provide a function that can be used to convert appropriately
<code>lubridate_fun</code>	function to convert the time column into a lubridate object

method	sampling method. One of 'random', 'dist_max', or 'bias'.
class_label	the label given to the sampled points. Defaults to background
return_pres	return presences together with background in a single tibble
time_buffer	the buffer on the time axis around presences that defines their effect when sampling background with method 'max_dist'. If set to zero, presences have an effect only on the time step to which they are assigned in raster; if a positive value, it defines the number of days before and after the date provided in the time column for which the presence should be considered (e.g. 20 days means that a presence is considered in all time steps equivalent to plus and minus twenty days from its date).

Details

Note that the time axis of the raster should be in POSIXct or Date format, or use 'tstep="years"'. See [terra::time\(\)](#) for details on how to set the time axis.

Value

An object of class [tibble::tibble](#). If presences are returned, the presence level is set as the reference (to match the expectations in the [yardstick](#) package that considers the first level to be the event)

sample_pseudoabs	<i>Sample pseudo-absence points for SDM analysis</i>
------------------	--

Description

This function samples pseudo-absence points from a raster given a set of presences. The locations returned as the center points of the sampled cells, which can not overlap with the presences (in contrast to background points, see [sample_background](#)). The following methods are implemented:

- 'random': pseudo-absences randomly sampled from the region covered by the raster (i.e. not NAs).
- 'dist_min': pseudo-absences randomly sampled from the region excluding a buffer of 'dist_min' from presences (distances in 'm' for lonlat rasters, and in map units for projected rasters).
- 'dist_max': pseudo-absences randomly sampled from the unioned buffers of 'dist_max' from presences (distances in 'm' for lonlat rasters, and in map units for projected rasters). Using the union of buffers means that areas that are in multiple buffers are not oversampled. This is also referred to as "thickening".
- 'dist_disc': pseudo-absences randomly sampled from the unioned discs around presences with the two values of 'dist_disc' defining the minimum and maximum distance from presences.

Usage

```
sample_pseudoabs(
  data,
  raster,
  n,
  coords = NULL,
  method = "random",
  class_label = "pseudoabs",
  return_pres = TRUE
)
```

Arguments

data	An <code>sf::sf</code> data frame, or a data frame with coordinate variables. These can be defined in <code>coords</code> , unless they have standard names (see details below).
raster	the <code>terra::SpatRaster</code> or stars from which cells will be sampled
n	number of pseudoabsence points to sample
coords	a vector of length two giving the names of the "x" and "y" coordinates, as found in data. If left to <code>NULL</code> , the function will try to guess the columns based on standard names <code>c("x", "y")</code> , <code>c("X", "Y")</code> , <code>c("longitude", "latitude")</code> , or <code>c("lon", "lat")</code>
method	sampling method. One of 'random', 'dist_min', 'dist_max', or 'dist_disc'. Threshold distances are set as additional elements of a vector, e.g <code>c('dist_min', 70000)</code> or <code>c('dist_disc', 50000, 200000)</code> .
class_label	the label given to the sampled points. Defaults to <code>pseudoabs</code>
return_pres	return presences together with pseudoabsences in a single tibble

Value

An object of class `tibble::tibble`. If presences are returned, the presence level is set as the reference (to match the expectations in the `yardstick` package that considers the first level to be the event)

`sample_pseudoabs_time` *Sample pseudo-absence points for SDM analysis for points with a time point.*

Description

This function samples pseudo-absence points from a raster given a set of presences. The locations returned as the center points of the sampled cells, which can not overlap with the presences (in contrast to background points, see [sample_background_time](#)). The following methods are implemented:

- 'random': pseudo-absences randomly sampled from the region covered by the raster (i.e. not NAs).

- `'dist_min'`: pseudo-absences randomly sampled from the region excluding a buffer of `'dist_min'` from presences (distances in 'm' for lonlat rasters, and in map units for projected rasters).
- `'dist_max'`: pseudo-absences randomly sampled from the unioned buffers of `'dist_max'` from presences (distances in 'm' for lonlat rasters, and in map units for projected rasters). Using the union of buffers means that areas that are in multiple buffers are not oversampled. This is also referred to as "thickening".
- `'dist_disc'`: pseudo-absences randomly sampled from the unioned discs around presences with the two values of `'dist_disc'` defining the minimum and maximum distance from presences.

Usage

```
sample_pseudoabs_time(
  data,
  raster,
  n_per_presence,
  coords = NULL,
  time_col = "time",
  lubridate_fun = c,
  method = "random",
  class_label = "pseudoabs",
  return_pres = TRUE,
  time_buffer = 0
)
```

Arguments

<code>data</code>	An <code>sf::sf</code> data frame, or a data frame with coordinate variables. These can be defined in <code>coords</code> , unless they have standard names (see details below).
<code>raster</code>	the <code>terra::SpatRaster</code> , stars or <code>terra::SpatRasterDataset</code> from which cells will be sampled. If a <code>terra::SpatRasterDataset</code> , the first dataset will be used to define which cells are valid, and which are NAs.
<code>n_per_presence</code>	number of pseudoabsence points to sample for each presence
<code>coords</code>	a vector of length two giving the names of the "x" and "y" coordinates, as found in data. If left to <code>NULL</code> , the function will try to guess the columns based on standard names <code>c("x", "y")</code> , <code>c("X", "Y")</code> , <code>c("longitude", "latitude")</code> , or <code>c("lon", "lat")</code>
<code>time_col</code>	The name of the column with time; if time is not a lubridate object, use <code>lubridate_fun</code> to provide a function that can be used to convert appropriately
<code>lubridate_fun</code>	function to convert the time column into a lubridate object
<code>method</code>	sampling method. One of <code>'random'</code> , <code>'dist_min'</code> , <code>'dist_max'</code> , or <code>'dist_disc'</code> .
<code>class_label</code>	the label given to the sampled points. Defaults to <code>pseudoabs</code>
<code>return_pres</code>	return presences together with pseudoabsences in a single tibble
<code>time_buffer</code>	the buffer on the time axis around presences that defines their effect when sampling pseudoabsences. If set to zero, presences have an effect only on the time step to which they are assigned in raster; if a positive value, it defines the number of days before and after the date provided in the <code>time</code> column for which the

presence should be considered (e.g. 20 days means that a presence is considered in all time steps equivalent to plus and minus twenty days from its date).

Details

#' @details Note that the time axis of the raster should be in POSIXct or Date format, or use 'tstep="years"'. See [terra::time\(\)](#) for details on how to set the time axis.

Value

An object of class [tibble::tibble](#). If presences are returned, the presence level is set as the reference (to match the expectations in the [yardstick](#) package that considers the first level to be the event)

sdm_metric_set	<i>Metric set for SDM</i>
----------------	---------------------------

Description

This function returns a [yardstick::metric_set](#) that includes [boyce_cont\(\)](#), [yardstick::roc_auc\(\)](#) and [tss_max\(\)](#), the most commonly used metrics for SDM.

Usage

```
sdm_metric_set(...)
```

Arguments

... additional metrics to be added to the [yardstick::metric_set](#). See the help to [yardstick::metric_set\(\)](#) for constraints on the type of metrics that can be mixed.

Value

a [yardstick::metric_set](#) object.

Examples

```
sdm_metric_set()
sdm_metric_set(accuracy)
```

sdm_spec_boost_tree *Model specification for a Boosted Trees model for SDM*

Description

This function returns a [parsnip::model_spec](#) for a Boosted Trees model to be used as a classifier of presences and absences in Species Distribution Model. It uses the library `xgboost` to fit boosted trees; to use another library, simply build the [parsnip::model_spec](#) directly.

Usage

```
sdm_spec_boost_tree(..., tune = c("sdm", "all", "custom", "none"))
```

Arguments

- | | |
|------|---|
| ... | parameters to be passed to parsnip::boost_tree() to customise the model. See the help of that function for details. |
| tune | character defining the tuning strategy. Valid strategies are: <ul style="list-style-type: none"> • "sdm" chooses hyperparameters that are most important to tune for an sdm (for <i>boost_tree</i>: 'mtry', 'trees', 'tree_depth', 'learn_rate', 'loss_reduction', and 'stop_iter') • "all" tunes all hyperparameters (for <i>boost_tree</i>: 'mtry', 'trees', 'tree_depth', 'learn_rate', 'loss_reduction', 'stop_iter', 'min_n' and 'sample_size') • "custom" passes the options from '...' • "none" does not tune any hyperparameter |

Value

a [parsnip::model_spec](#) of the model.

See Also

Other "sdm model specifications": [sdm_spec_gam\(\)](#), [sdm_spec_glm\(\)](#), [sdm_spec_maxent\(\)](#), [sdm_spec_rand_forest\(\)](#)

Examples

```
standard_bt_spec <- sdm_spec_boost_tree()
full_bt_spec <- sdm_spec_boost_tree(tune = "all")
custom_bt_spec <- sdm_spec_boost_tree(tune = "custom", mtry = tune::tune())
```

sdm_spec_gam

*Model specification for a GAM for SDM***Description**

This function returns a [parsnip::model_spec](#) for a General Additive Model to be used as a classifier of presences and absences in Species Distribution Model.

Usage

```
sdm_spec_gam(..., tune = "none")
```

Arguments

`...` parameters to be passed to [parsnip::gen_additive_mod\(\)](#) to customise the model. See the help of that function for details.

`tune` character defining the tuning strategy. As there are no hyperparameters to tune in a *gam*, the only valid option is "none". This parameter is present for consistency with other `sdm_spec_*` functions, but it does nothing in this case.

Details

Note that, when using GAMs in a `workflow_set()`, it is necessary to update the model with [gam_formula\(\)](#) (see [parsnip::model_formula](#) for a discussion of formulas with special terms in `tidymodels`):

```
workflow_set(
  preproc = list(default = my_recipe),
  models = list(gam = sdm_spec_gam()),
  cross = TRUE
) %>% update_workflow_model("default_gam",
  spec = sdm_spec_gam(),
  formula = gam_formula(my_recipe))
```

Value

a [parsnip::model_spec](#) of the model.

See Also

[parsnip::gen_additive_mod\(\)](#) [gam_formula\(\)](#)

Other "sdm model specifications": [sdm_spec_boost_tree\(\)](#), [sdm_spec_glm\(\)](#), [sdm_spec_maxent\(\)](#), [sdm_spec_rand_forest\(\)](#)

Examples

```
my_gam_spec <- sdm_spec_gam()
```

sdm_spec_glm

Model specification for a GLM for SDM

Description

This function returns a [parsnip::model_spec](#) for a Generalised Linear Model to be used as a classifier of presences and absences in Species Distribution Model.

Usage

```
sdm_spec_glm(..., tune = "none")
```

Arguments

`...` parameters to be passed to [parsnip::logistic_reg\(\)](#) to customise the model. See the help of that function for details.

`tune` character defining the tuning strategy. As there are no hyperparameters to tune in a *glm*, the only valid option is "none". This parameter is present for consistency with other `sdm_spec_*` functions, but it does nothing in this case.

Value

a [parsnip::model_spec](#) of the model.

See Also

Other "sdm model specifications": [sdm_spec_boost_tree\(\)](#), [sdm_spec_gam\(\)](#), [sdm_spec_maxent\(\)](#), [sdm_spec_rand_forest\(\)](#)

Examples

```
my_spec_glm <- sdm_spec_glm()
```

sdm_spec_maxent

Model specification for a MaxEnt for SDM

Description

This function returns a [parsnip::model_spec](#) for a MaxEnt model to be used in Species Distribution Models.

Usage

```
sdm_spec_maxent(..., tune = c("sdm", "all", "custom", "none"))
```

Arguments

- ... parameters to be passed to `maxent()` to customise the model. See the help of that function for details.
- tune character defining the tuning strategy. Valid strategies are:
- "sdm" chooses hyper-parameters that are most important to tune for an sdm (for *maxent*, 'feature_classes' and 'regularization_multiplier')
 - "all" tunes all hyperparameters (for *maxent*, 'feature_classes' and 'regularization_multiplier', the same as with tune = "sdm")
 - "custom" passes the options from '...'
 - "none" does not tune any hyperparameter

Value

a `parsnip::model_spec` of the model.

See Also

Other "sdm model specifications": `sdm_spec_boost_tree()`, `sdm_spec_gam()`, `sdm_spec_glm()`, `sdm_spec_rand_forest()`

Examples

```
test_maxent_spec <- sdm_spec_maxent(tune = "sdm")
test_maxent_spec
# setting specific values
sdm_spec_maxent(tune = "custom", feature_classes = "lq")
```

sdm_spec_rand_forest *Model specification for a Random Forest for SDM*

Description

This function returns a `parsnip::model_spec` for a Random Forest to be used as a classifier of presences and absences in Species Distribution Models. It uses the library `ranger` to fit boosted trees; to use another library, simply build the `parsnip::model_spec` directly.

Usage

```
sdm_spec_rand_forest(..., tune = c("sdm", "all", "custom", "none"))

sdm_spec_rf(..., tune = c("sdm", "all", "custom", "none"))
```

Arguments

- ... parameters to be passed to `parsnip::rand_forest()` to customise the model. See the help of that function for details.
- tune character defining the tuning strategy. Valid strategies are:
- "sdm" chooses hyperparameters that are most important to tune for an sdm (for *rf*, 'mtry')
 - "all" tunes all hyperparameters (for *rf*, 'mtry', 'trees' and 'min')
 - "custom" passes the options from '...'
 - "none" does not tune any hyperparameter

Details

`sdm_spec_rf()` is simply a short form for `sm_spec_rand_forest()`.

Value

a `parsnip::model_spec` of the model.

See Also

Other "sdm model specifications": `sdm_spec_boost_tree()`, `sdm_spec_gam()`, `sdm_spec_glm()`, `sdm_spec_maxent()`

Examples

```
test_rf_spec <- sdm_spec_rf(tune = "sdm")
test_rf_spec
# combining tuning with specific values for other hyperparameters
sdm_spec_rf(tune = "sdm", trees = 100)
```

simple_ensemble

Simple ensemble

Description

A simple ensemble is a collection of workflows for which predictions will be combined in a simple way (e.g. by taking either the mean or median). Usually these workflows will consists each of the best version of a given model algorithm following tuning. The workflows are fitted to the full training dataset before making predictions.

Usage

```
simple_ensemble(...)
```

Arguments

- ... not used, this function just creates an empty `simple_ensemble` object. Members are added with `add_best_candidates()`

Value

an empty `simple_ensemble`. This is a tibble with columns:

- `wflow_id`: the name of the workflows for which the best model was chosen
- `workflow`: the trained workflow objects
- `metrics`: metrics based on the crossvalidation resampling used to tune the models

`spatial_initial_split` *Simple Training/Test Set Splitting for spatial data*

Description

`spatial_initial_split` creates a single binary split of the data into a training set and testing set. All strategies from the package `spatialsample` are available; a random split from that strategy will be used to generate the initial split.

Usage

```
spatial_initial_split(data, prop, strategy, ...)
```

Arguments

<code>data</code>	A dataset (data.frame or tibble)
<code>prop</code>	The proportion of data to be retained for modelling/analysis.
<code>strategy</code>	A sampling strategy from <code>spatialsample</code>
<code>...</code>	parameters to be passed to the strategy

Value

An `rsplit` object that can be used with the `rsample::training` and `rsample::testing` functions to extract the data in each split.

Examples

```
set.seed(123)
block_initial <- spatial_initial_split(boston_canopy,
  prop = 1 / 5, spatial_block_cv
)
testing(block_initial)
training(block_initial)
```

thin_by_cell	<i>Thin point dataset to have 1 observation per raster cell</i>
--------------	---

Description

This function thins a dataset so that only one observation per cell is retained.

Usage

```
thin_by_cell(data, raster, coords = NULL, drop_na = TRUE, agg_fact = NULL)
```

Arguments

data	An <code>sf::sf</code> data frame, or a data frame with coordinate variables. These can be defined in <code>coords</code> , unless they have standard names (see details below).
raster	A <code>terra::SpatRaster</code> or stars object that defined the grid
coords	a vector of length two giving the names of the "x" and "y" coordinates, as found in data. If left to <code>NULL</code> , the function will try to guess the columns based on standard names <code>c("x", "y")</code> , <code>c("X", "Y")</code> , <code>c("longitude", "latitude")</code> , or <code>c("lon", "lat")</code>
drop_na	boolean on whether locations that are NA in the raster should be dropped.
agg_fact	positive integer. Aggregation factor expressed as number of cells in each direction (horizontally and vertically). Or two integers (horizontal and vertical aggregation factor) or three integers (when also aggregating over layers). Defaults to <code>NULL</code> , which implies no aggregation (i.e. thinning is done on the grid of raster)

Details

Further thinning can be achieved by aggregating cells in the raster before thinning, as achieved by setting `agg_fact > 1` (aggregation works in a manner equivalent to `terra::aggregate()`). Note that if data is an `sf` object, the function will transform the coordinates to the same projection as the raster (recommended); if data is a `data.frame`, it is up to the user to ensure that the coordinates are in the correct units.

Value

An object of class `sf::sf` or `data.frame`, the same as "data".

thin_by_cell_time	<i>Thin point dataset to have 1 observation per raster cell per time slice</i>
-------------------	--

Description

This function thins a dataset so that only one observation per cell per time slice is retained. We use a raster with layers as time slices to define the data cube on which thinning is enforced (see details below on how time should be formatted).

Usage

```
thin_by_cell_time(
  data,
  raster,
  coords = NULL,
  time_col = "time",
  lubridate_fun = c,
  drop_na = TRUE,
  agg_fact = NULL
)
```

Arguments

data	An <code>sf::sf</code> data frame, or a data frame with coordinate variables. These can be defined in <code>coords</code> , unless they have standard names (see details below).
raster	A <code>terra::SpatRaster</code> or stars object that defined the grid with layers corresponding to the time slices (times should be set as either <code>POSIXlt</code> or "years", see <code>terra::time()</code> for details), or a <code>terra::SpatRasterDataset</code> where the first dataset will be used (again, times for that dataset should be set as either <code>POSIXlt</code> or "years") <code>terra::time()</code>
coords	a vector of length two giving the names of the "x" and "y" coordinates, as found in data. If left to <code>NULL</code> , the function will try to guess the columns based on standard names <code>c("x", "y")</code> , <code>c("X", "Y")</code> , <code>c("longitude", "latitude")</code> , or <code>c("lon", "lat")</code>
time_col	The name of the column with time; if time is not a lubridate object, use <code>lubridate_fun</code> to provide a function that can be used to convert appropriately
lubridate_fun	function to convert the time column into a lubridate object
drop_na	boolean on whether locations that are NA in the raster should be dropped.
agg_fact	positive integer. Aggregation factor expressed as number of cells in each direction (horizontally and vertically). Or two integers (horizontal and vertical aggregation factor) or three integers (when also aggregating over layers). Defaults to <code>NULL</code> , which implies no aggregation (i.e. thinning is done on the grid of raster)

Details

Further spatial thinning can be achieved by aggregating cells in the raster before thinning, as achieved by setting `agg_fact > 1` (aggregation works in a manner equivalent to `terra::aggregate()`). Note that if data is an `sf` object, the function will transform the coordinates to the same projection as the raster (recommended); if data is a `data.frame`, it is up to the user to ensure that the coordinates are in the correct units.

Value

An object of class `sf::sf` or `data.frame`, the same as "data".

thin_by_dist	<i>Thin points dataset based on geographic distance</i>
--------------	---

Description

This function thins a dataset so that only observations that have a distance from each other greater than "dist_min" are retained.

Usage

```
thin_by_dist(
  data,
  dist_min,
  coords = NULL,
  dist_method = c("great_circle", "euclidean")
)
```

Arguments

data	An <code>sf::sf</code> data frame, or a data frame with coordinate variables. These can be defined in <code>coords</code> , unless they have standard names (see details below).
dist_min	Minimum distance between points (in units appropriate for the projection, or meters for lonlat data).
coords	A vector of length two giving the names of the "x" and "y" coordinates, as found in data. If left to <code>NULL</code> , the function will try to guess the columns based on standard names <code>c("x", "y")</code> , <code>c("X", "Y")</code> , <code>c("longitude", "latitude")</code> , or <code>c("lon", "lat")</code> .
dist_method	method to compute distance, either "euclidean" or "great_circle". Defaults to "great_circle", which is more accurate but takes slightly longer.

Details

Distances are measured in the appropriate units for the projection used. In case of raw latitude and longitude (e.g. as provided in a `data.frame`), the crs is set to WGS84, and units are set to meters.

This function is a modified version of the algorithm in `spThin`, adapted to work on `sf` objects.

Value

An object of class `sf::sf` or `data.frame`, the same as "data".

thin_by_dist_time	<i>Thin points dataset based on geographic and temporal distance</i>
-------------------	--

Description

This function thins a dataset so that only observations that have a distance from each other greater than "dist_min" in space and "interval_min" in time are retained.

Usage

```
thin_by_dist_time(
  data,
  dist_min,
  interval_min,
  coords = NULL,
  time_col = "time",
  lubridate_fun = c,
  dist_method = c("great_circle", "euclidean")
)
```

Arguments

data	An <code>sf::sf</code> data frame, or a data frame with coordinate variables. These can be defined in <code>coords</code> , unless they have standard names (see details below).
dist_min	Minimum distance between points (in units appropriate for the projection, or meters for lonlat data).
interval_min	Minimum time interval between points, in days.
coords	A vector of length two giving the names of the "x" and "y" coordinates, as found in data. If left to <code>NULL</code> , the function will try to guess the columns based on standard names <code>c("x", "y")</code> , <code>c("X", "Y")</code> , <code>c("longitude", "latitude")</code> , or <code>c("lon", "lat")</code> .
time_col	The name of the column with time; if time is not a lubridate object, use <code>lubridate_fun</code> to provide a function that can be used to convert appropriately.
lubridate_fun	function to convert the time column into a lubridate object
dist_method	method to compute distance, either "euclidean" or "great_circle". Defaults to "great_circle", which is more accurate but takes slightly longer.

Details

Geographic distances are measured in the appropriate units for the projection used. In case of raw latitude and longitude (e.g. as provided in a data.frame), the crs is set to WGS84, and units are set to meters. Time interval are estimated in days. Note that for very long time period, the simple conversion $x \text{ years} = 365 * x \text{ days}$ might lead to slightly shorter intervals than expected, as it ignores leap years. The function `y2d()` provides a closer approximation.

This function an algorithm analogous to `spThin`, with the exception that neighbours are defined in terms of both space and time.

Value

An object of class `sf::sf` or `data.frame`, the same as "data".

tss	<i>TSS - True Skill Statistics</i>
-----	------------------------------------

Description

The True Skills Statistic, which is defined as

Usage

```
tss(data, ...)

## S3 method for class 'data.frame'
tss(
  data,
  truth,
  estimate,
  estimator = NULL,
  na_rm = TRUE,
  case_weights = NULL,
  event_level = "first",
  ...
)
```

Arguments

data	Either a data.frame containing the columns specified by the truth and estimate arguments, or a table/matrix where the true class results should be in the columns of the table.
...	Not currently used.
truth	The column identifier for the true class results (that is a factor). This should be an unquoted column name although this argument is passed by expression and supports quasiquotation (you can unquote column names). For <code>_vec()</code> functions, a factor vector.

<code>estimate</code>	The column identifier for the predicted class results (that is also factor). As with truth this can be specified different ways but the primary method is to use an unquoted variable name. For <code>_vec()</code> functions, a factor vector.
<code>estimator</code>	One of: "binary", "macro", "macro_weighted", or "micro" to specify the type of averaging to be done. "binary" is only relevant for the two class case. The other three are general methods for calculating multiclass metrics. The default will automatically choose "binary" or "macro" based on estimate.
<code>na_rm</code>	A logical value indicating whether NA values should be stripped before the computation proceeds.
<code>case_weights</code>	The optional column identifier for case weights. This should be an unquoted column name that evaluates to a numeric column in data. For <code>_vec()</code> functions, a numeric vector.
<code>event_level</code>	A single string. Either "first" or "second" to specify which level of truth to consider as the "event". This argument is only applicable when estimator = "binary". The default is "first".

Details

sensitivity+specificity +1

This function is a wrapper around `yardstick::j_index()`, another name for the same quantity. Note that this function takes the classes as predicted by the model without any calibration (i.e. making a split at 0.5 probability). This is usually not the metric used for Species Distribution Models, where the threshold is recalibrated to maximise TSS; for that purpose, use `tss_max()`.

Value

A tibble with columns `.metric`, `.estimator`, and `.estimate` and 1 row of values. For grouped data frames, the number of rows returned will be the same as the number of groups.

Examples

```
# Two class
data("two_class_example")
tss(two_class_example, truth, predicted)
# Multiclass
library(dplyr)
data(hpc_cv)
# Groups are respected
hpc_cv %>%
  group_by(Resample) %>%
  tss(obs, pred)
```

tss_max

*Maximum TSS - True Skill Statistics***Description**

The True Skills Statistic, which is defined as

Usage

```
tss_max(data, ...)

## S3 method for class 'data.frame'
tss_max(
  data,
  truth,
  ...,
  estimator = NULL,
  na_rm = TRUE,
  event_level = "first",
  case_weights = NULL
)

## S3 method for class 'sf'
tss_max(data, ...)

tss_max_vec(
  truth,
  estimate,
  estimator = NULL,
  na_rm = TRUE,
  event_level = "first",
  case_weights = NULL,
  ...
)
```

Arguments

data	Either a data.frame containing the columns specified by the truth and estimate arguments, or a table/matrix where the true class results should be in the columns of the table.
...	A set of unquoted column names or one or more dplyr selector functions to choose which variables contain the class probabilities. If truth is binary, only 1 column should be selected, and it should correspond to the value of event_level. Otherwise, there should be as many columns as factor levels of truth and the ordering of the columns should be the same as the factor levels of truth.

truth	The column identifier for the true class results (that is a factor). This should be an unquoted column name although this argument is passed by expression and supports quasiquotation (you can unquote column names). For <code>_vec()</code> functions, a factor vector.
estimator	One of "binary", "hand_till", "macro", or "macro_weighted" to specify the type of averaging to be done. "binary" is only relevant for the two class case. The others are general methods for calculating multiclass metrics. The default will automatically choose "binary" if truth is binary, "hand_till" if truth has >2 levels and case_weights isn't specified, or "macro" if truth has >2 levels and case_weights is specified (in which case "hand_till" isn't well-defined).
na_rm	A logical value indicating whether NA values should be stripped before the computation proceeds.
event_level	A single string. Either "first" or "second" to specify which level of truth to consider as the "event". This argument is only applicable when estimator = "binary". The default uses an internal helper that generally defaults to "first"
case_weights	The optional column identifier for case weights. This should be an unquoted column name that evaluates to a numeric column in data. For <code>_vec()</code> functions, a numeric vector.
estimate	If truth is binary, a numeric vector of class probabilities corresponding to the "relevant" class. Otherwise, a matrix with as many columns as factor levels of truth. It is assumed that these are in the same order as the levels of truth.

Details

sensitivity+specificity +1

This function calibrates the probability threshold to classify presences to maximise the TSS.

There is no multiclass version of this function, it only operates on binary predictions (e.g. presences and absences in SDMs).

Value

A tibble with columns `.metric`, `.estimator`, and `.estimate` and 1 row of values. For grouped data frames, the number of rows returned will be the same as the number of groups.

See Also

Other class probability metrics: [boyce_cont\(\)](#), [kap_max\(\)](#)

Examples

```
tss_max(two_class_example, truth, Class1)
```

`y2d`*Convert a time interval from years to days*

Description

This function takes a time interval in years and converts into days, the unit commonly used in time operations in R. The simple conversion $x * 365$ does not work for large number of years, due to the presence of leap years.

Usage`y2d(x)`**Arguments**

`x` the number of years of the interval

Value

a `difftime` object (in days)

Examples

```
y2d(1)
y2d(1000)
```

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