# Package 'spatialRF'

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Title Easy Spatial Modeling with Random Forest

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#### BugReports https://github.com/BlasBenito/spatialRF/issues/

**Description** Automatic generation and selection of spatial predictors for spatial regression with Random Forest. Spatial predictors are surrogates of variables driving the spatial structure of a response variable. The package offers two methods to generate spatial predictors from a distance matrix among training cases: 1) Moran's Eigenvector Maps (MEMs; Dray, Legendre, and Peres-Neto 2006 <DOI:10.1016/j.ecolmodel.2006.02.015>): computed as the eigenvectors of a weighted matrix of dis-

tances; 2) RFsp (Hengl et al. <DOI:10.7717/peerj.5518>): columns of the distance matrix used as spatial predictors. Spatial predictors help minimize the spatial autocorrelation of the model residuals and facilitate an honest assessment of the importance scores of the nonspatial predictors. Additionally, functions to reduce multicollinearity, identify relevant variable interactions, tune random forest hyperparameters, assess model transferability via spatial cross-validation, and explore model results via partial dependence curves and interaction surfaces are included in the package. The modelling functions are built around the highly efficient 'ranger' package (Wright and Ziegler 2017 <DOI:10.18637/jss.v077.i01>).

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auc

Area under the ROC curve

# Description

Computes the area under the ROC curve in models with binary responses.

### Usage

auc(o, p)

# Arguments

0	Numeric vector with observations, must have the same length as p.
р	Numeric vector with predictions, must have the same length as o.

# Value

Numeric, AUC value.

### Examples

```
if(interactive()){
  out <- auc(
    o = c(0, 0, 1, 1),
    p = c(0.1, 0.6, 0.4, 0.8)
  )
}</pre>
```

auto\_cor

Multicollinearity reduction via Pearson correlation

### Description

Computes the correlation matrix among a set of predictors, orders the correlation matrix according to a user-defined preference order, and removes variables one by one, taking into account the preference order, until the remaining ones are below a given Pearson correlation threshold. **Warning**: variables in preference.order not in colnames(x), and non-numeric columns are removed silently from x and preference.order. The same happens with rows having NA values (na.omit() is applied). The function issues a warning if zero-variance columns are found.

#### Usage

```
auto_cor(
  x = NULL,
  preference.order = NULL,
  cor.threshold = 0.5,
  verbose = TRUE
)
```

#### Arguments

х	A data frame with predictors, or the result of auto_vif() Default: NULL.	
preference.order		
	Character vector indicating the user's order of preference to keep variables. Doesn't need to contain If not provided, variables in $x$ are prioritised by their column order. Default: NULL.	
cor.threshold	Numeric between 0 and 1, with recommended values between 0.5 and 0.9. Max- imum Pearson correlation between any pair of the selected variables. Default: 0.50	
verbose	Logical. if TRUE, describes the function operations to the user. Default:: TRUE	

### Details

Can be chained together with auto\_vif() through pipes, see the examples below.

### auto\_vif

### Value

List with three slots:

- cor: correlation matrix of the selected variables.
- selected.variables: character vector with the names of the selected variables.
- selected.variables.df: data frame with the selected variables.

### See Also

auto\_vif()

### Examples

```
if(interactive()){
#load data
data(plant_richness_df)
#on a data frame
out <- auto_cor(x = plant_richness_df[, 5:21])</pre>
#getting the correlation matrix
 out$cor
#getting the names of the selected variables
out$selected.variables
#getting the data frame of selected variables
out$selected.variables.df
#on the result of auto_vif
out <- auto_vif(x = plant_richness_df[, 5:21])</pre>
out <- auto_cor(x = out)</pre>
#with pipes
out <- plant_richness_df[, 5:21] %>%
auto_vif() %>%
auto_cor()
```

```
}
```

auto\_vif

### Description

Selects predictors that are not linear combinations of other predictors by using computing their variance inflation factors (VIF). Allows the user to define an order of preference for the selection of predictors. **Warning**: variables in preference.order not in colnames(x), and non-numeric columns are removed silently from x and preference.order. The same happens with rows having NA values (na.omit() is applied). The function issues a warning if zero-variance columns are found.

#### Usage

```
auto_vif(
  x = NULL,
  preference.order = NULL,
  vif.threshold = 5,
  verbose = TRUE
)
```

#### Arguments

х	A data frame with predictors or the result of auto_cor(). Default: NULL.	
preference.order		
	a character vector with columns names of x ordered by the user preference, Default: NULL.	
vif.threshold	Numeric between 2.5 and 10 defining the selection threshold for the VIF analysis. Higher numbers result in a more relaxed variable selection. Default: 5.	
verbose	Logical. if TRUE, describes the function operations to the user. Default:: TRUE	

### Details

This function has two modes of operation:

- 1. When the argument preference.order is NULL, the function removes on each iteration the variable with the highest VIF until all VIF values are lower than vif.threshold.
- 2. When preference.order is provided, the variables are selected by giving them priority according to their order in preference.order. If there are variables not in preference.order, these are selected as in option 1. Once both groups of variables have been processed, all variables are put together and selected by giving priority to the ones in preference.order. This method preserves the variables desired by the user as much as possible.

Can be chained together with auto\_cor() through pipes, see the examples below.

#### Value

List with three slots:

- vif: data frame with the names of the selected variables and their respective VIF scores.
- selected.variables: character vector with the names of the selected variables.
- selected.variables.df: data frame with the selected variables.

beowulf\_cluster

#### See Also

auto\_cor()

#### Examples

```
if(interactive()){
```

#loading data
data(plant\_richness\_df)

```
#on a data frame
out <- auto_vif(x = plant_richness_df[, 5:21])</pre>
```

#getting out the vif data frame
out\$vif

#getting the names of the selected variables
out\$selected.variables

#getting the data frame of selected variables
out\$selected.variables.df

```
#on the result of auto_cor
out <- auto_cor(x = plant_richness_df[, 5:21])
out <- auto_vif(x = out)</pre>
```

```
#with pipes
out <- plant_richness_df[, 5:21] %>%
auto_cor() %>%
auto_vif()
```

### }

beowulf\_cluster Defines a beowulf cluster

#### Description

Defines a Beowulf cluster from the IPs of the machines in the cluster, the number of cores of each machine, and the user name. The returned cluster has to be registered with doParallel::registerDoParallel().

#### Usage

```
beowulf_cluster(
   cluster.ips = NULL,
   cluster.cores = NULL,
   cluster.user = Sys.info()[["user"]],
   cluster.port = "11000",
   outfile = NULL
)
```

### Arguments

cluster.ips	Character vector with the IPs of the machines in the cluster. The first machine will be considered the main node of the cluster, and will generally be the machine on which the R code is being executed. Default: NULL.
cluster.cores	Numeric integer vector, number of cores on each machine. Default: NULL.
cluster.user	Character string, name of the user (should be the same throughout machines), Defaults to the current system user.
cluster.port	Character, port used by the machines in the cluster to communicate. The firewall in all computers must allow traffic from and to such port. Default: "11000"
outfile	Where to direct the messages provided by the workers. When working on a local computer, "" prints the worker's messages in the console. A text file path will append worker's messages on the given file. Default: /dev/null en Linux and nul: on windows.

### Value

A list ready to be used as input for the spec argument of the function makeCluster.

### Examples

```
if(interactive()){
beowulf.cluster <- beowulf_cluster(
  cluster.ips = c(
    "10.42.0.1",
    "10.42.0.34",
    "10.42.0.104"
    ),
  cluster.cores = c(7, 4, 4),
  cluster.user = "blas",
  cluster.port = "11000"
)

doParallel::registerDoParallel(cl = beowulf.cluster)
#PARALLELIZED foreach LOOP HERE
parallel::stopCluster(cl = beowulf.cluster)
}</pre>
```

case\_weights

### Description

When the data is binary, setting the ranager argument case.weights helps to minimize the issues produced by class imbalance. This function takes a binary response variable and returns a vector of weights populated with the values 1/#zeros and 1/#ones. It is used internally by the function rf().

#### Usage

```
case_weights(data = NULL, dependent.variable.name = NULL)
```

### Arguments

data Data frame with a response variable and a set of predictors. Default: NULL

dependent.variable.name

Character string with the name of the response variable. Must be in the column names of data. Default: NULL

### Value

A vector with a length equal to nrow(data) with the respective weights of the cases.

#### Examples

```
if(interactive()){
  data <- data.frame(
    response = c(0, 0, 0, 1, 1)
  )
  case_weights(
    data = data,
    dependent.variable.name = "response"
  )
}</pre>
```

default\_distance\_thresholds

Default distance thresholds to generate spatial predictors

### Description

Generates four distance thresholds, from 0 to max(distance.matrix)/2.

### Usage

```
default_distance_thresholds(distance.matrix = NULL)
```

### Arguments

distance.matrix

Distance matrix. Default: NULL.

### Value

A numeric vector with distance thresholds.

### Examples

if(interactive()){
 #loading example distance matrix
 data(distance\_matrix)
 #computing set of default distance thresholds

default\_distance\_thresholds(distance\_matrix)

}

distance\_matrix Matrix of distances among ecoregion edges.

### Description

Distance matrix (in km) among the edges of the American ecoregions described in the plant\_richness\_df dataset.

#### Usage

data(distance\_matrix)

### Format

A numeric matrix with 227 rows and columns.

#### See Also

plant\_richness\_df

double\_center\_distance\_matrix

Double centers a distance matrix

### Description

Generates a double-centered matrix (row and column means are zero) from the weights of a distance matrix (see weights\_from\_distance\_matrix()) and a distance threshold. This is a required step before the computation of Moran's Eigenvector Maps.

#### Usage

```
double_center_distance_matrix (
   distance.matrix = NULL,
   distance.threshold = 0
)
```

### Arguments

distance.matrix Distance matrix. Default: NULL. distance.threshold Numeric, positive, in the range of values of x. Distances below this value in the distance matrix are set to 0. Default: 0.

### Value

A double-centered matrix of the same dimensions as x.

### See Also

weights\_from\_distance\_matrix(), mem(), mem\_multithreshold()

#### Examples

```
if(interactive()){
```

#loading the distance matrix
data(distance\_matrix)

x <- double\_center\_distance\_matrix(</pre>

```
distance.matrix = distance_matrix
)
x
}
```

filter\_spatial\_predictors

Removes redundant spatial predictors

### Description

Removes spatial predictors that are pair-wise correlated with other spatial predictors (which happens when there are several close distance thresholds), and spatial predictors correlated with non-spatial predictors.

### Usage

```
filter_spatial_predictors(
  data = NULL,
  predictor.variable.names = NULL,
  spatial.predictors.df = NULL,
  cor.threshold = 0.5
)
```

### Arguments

data	Data frame with a response variable and a set of predictors. Default: NULL
predictor.vari	able.names
	Character vector with the names of the predictive variables. Every element of
	this vector must be in the column names of data. Default: NULL
spatial.predictors.df	
	Data frame of spatial predictors.
cor.threshold	Numeric between 0 and 1, maximum Pearson correlation between any pair of the selected variables. Default: 0.50

#### Value

A data frame with non-redundant spatial predictors.

### Examples

```
if(interactive()){
#loading data
data("distance_matrix")
```

```
data("plant_richness_df")
```

#### get\_evaluation

```
#computing Moran's Eigenvector Maps
spatial.predictors.df <- mem_multithreshold(
    distance_matrix = distance_matrix,
    distance.thresholds = c(0, 1000)
    )
#filtering spatial predictors
spatial.predictors.df <- filter_spatial_predictors(
    data = plant_richness_df,
    predictor.variable.names = colnames(plant_richness_df)[5:21],
    spatial.predictors.df = spatial.predictors.df,
    cor.threshold = 0.50
    )
```

}

get\_evaluation Gets performa

Gets performance data frame from a cross-validated model

#### Description

Returns performance metrics produced by rf\_evaluate().

#### Usage

```
get_evaluation(model)
```

### Arguments

model

A model fitted with rf\_evaluate().

#### Value

A data frame with evaluation scores. The following columns are shown:

- model: Identifies the given model. The values are "Full", (original model introduced into rf\_evaluate()), "Training" (model trained on an independent training spatial fold), and "Testing" (predictive performance of the training model on an independent testing spatial fold). The performance values of the "Testing" model represent the model performance on unseen data, and hence its ability to generalize.
- metric: Four values representing different evaluation metrics, "rmse", "nrmse", "r.squared", and "pseudo.r.squared".
- mean, sd, min, and max: Average, standard deviation, minimum, and maximum of each metric across the evaluation (cross-validation) iterations.

#### See Also

rf\_evaluate(), plot\_evaluation(), print\_evaluation()

### Examples

```
if(interactive()){
#loading data
data(plant_richness_df)
data(distance_matrix)
#fitting a random forest model
rf.model <- rf(</pre>
  data = plant_richness_df,
  dependent.variable.name = "richness_species_vascular",
  predictor.variable.names = colnames(plant_richness_df)[5:21],
  distance.matrix = distance_matrix,
  distance.thresholds = 0,
  n.cores = 1,
  verbose = FALSE
)
#evaluating the model with spatial cross-validation
rf.model <- rf_evaluate(</pre>
  model = rf.model,
  xy = plant_richness_df[, c("x", "y")],
  n.cores = 1,
  verbose = FALSE
)
#getting evaluation results from the model
x <- get_evaluation(rf.model)</pre>
х
}
```

get\_importance Gets the global importance data frame from a model

#### Description

Gets variable importance scores from rf(), rf\_repeat(), and rf\_spatial() models.

#### Usage

get\_importance(model)

### Arguments

model A model fitted with rf(), rf\_repeat(), or rf\_spatial(). Default: NULL

### Value

A data frame with variable names and importance scores.

get\_importance\_local

### See Also

rf(), rf\_repeat(), rf\_spatial(), plot\_importance(), print\_importance().

#### Examples

```
if(interactive()){
data(plant_richness_df)
data(distance_matrix)

rf.model <- rf(
   data = plant_richness_df,
   dependent.variable.name = "richness_species_vascular",
   predictor.variable.names = colnames(plant_richness_df)[5:21],
   distance.matrix = distance_matrix,
   distance.thresholds = 0,
   n.cores = 1,
   verbose = FALSE
)

x <- get_importance(rf.model)
x
}</pre>
```

get\_importance\_local Gets the local importance data frame from a model

### Description

```
Gets local importance scores from rf(), rf_repeat(), and rf_spatial() models.
```

#### Usage

```
get_importance_local(model)
```

#### Arguments

model A model fitted with rf(), rf\_repeat(), or rf\_spatial(). Default: NULL

### Value

A data frame with variable names and local importance scores.

### See Also

rf(), rf\_repeat(), rf\_spatial(), plot\_importance(), print\_importance().

### Examples

```
if(interactive()){
#loading example data
data(plant_richness_df)
data(distance_matrix)
#fittinga random forest model
rf.model <- rf(</pre>
  data = plant_richness_df,
  dependent.variable.name = "richness_species_vascular",
  predictor.variable.names = colnames(plant_richness_df)[5:21],
  distance.matrix = distance_matrix,
  distance.thresholds = 0,
  n.cores = 1,
  verbose = FALSE
)
#getting importance scores
x <- get_importance_local(rf.model)</pre>
х
}
```

get\_moran

#### Gets Moran's I test of model residuals

#### Description

Returns the Moran's I test on the residuals of a model produced by rf(), rf\_repeat(), or rf\_spatial().

### Usage

```
get_moran(model)
```

#### Arguments

```
model A model fitted with rf(), rf_repeat(), or rf_spatial()
```

#### Value

A data frame with Moran's I test results produced by moran\_multithreshold().

### See Also

```
moran(), moran_multithreshold(), plot_moran(), print_moran().
```

### get\_performance

if(interactive()){

### Examples

```
#loading example data
 data(plant_richness_df)
 data(distance_matrix)
 #fitting a random forest model
 rf.model <- rf(</pre>
  data = plant_richness_df,
  dependent.variable.name = "richness_species_vascular",
  predictor.variable.names = colnames(plant_richness_df)[5:21],
  distance.matrix = distance_matrix,
  distance.thresholds = c(0, 1000, 2000),
  n.cores = 1,
  verbose = FALSE
)
#getting Moran's I of the residuals
x <- get_moran(rf.model)</pre>
}
```

get\_performance Gets out-of-bag performance scores from a model

#### Description

Returns the performance slot of an rf(), rf\_repeat(), or rf\_spatial() model computed on the out-of-bag data.

### Usage

```
get_performance(model)
```

#### Arguments

```
model Model fitted with rf(), rf_repeat(), or rf_spatial().
```

#### Value

A data frame with four columns:

- metric Name of the performance metric.
- median Value of the performance metric. Truly a median only if the model is fitted with rf\_repeat().
- median\_absolute\_deviation median absolute deviation (MAD), only if the model is fitted with rf\_repeat(), and NA otherwise.

### See Also

print\_performance()

### Examples

```
if(interactive()){
#loading example data
data(plant_richness_df)
data(distance.matrix)
#fitting random forest model
rf.model <- rf(</pre>
  data = plant_richness_df,
  dependent.variable.name = "richness_species_vascular",
  predictor.variable.names = colnames(plant_richness_df)[5:21],
  distance.matrix = distance_matrix,
  distance.thresholds = 0,
  n.cores = 1,
  verbose = FALSE
)
#getting model performance
x <- get_performance(rf.model)</pre>
х
}
```

get\_predictions Gets model predictions

### Description

Returns model predictions from a model fitted with rf(), rf\_repeat(), or rf\_spatial().

### Usage

```
get_predictions(model)
```

### Arguments

model A model produced by rf(), rf\_repeat(), or rf\_spatial().

### Value

A vector with predictions, or median of the predictions across repetitions if the model was fitted with rf\_repeat().

### get\_residuals

### Examples

```
if(interactive()){
#loading example data
data(plant_richness_df)
#fitting a random forest model
rf.model <- rf(
   data = plant_richness_df,
   dependent.variable.name = "richness_species_vascular",
   predictor.variable.names = colnames(plant_richness_df)[5:21],
   n.cores = 1,
   verbose = FALSE
)
#get vector of predictions
x <- get_predictions(rf.model)
X
}</pre>
```

get\_residuals Gets model residuals

### Description

Returns the residuals of models fitted with rf(), rf\_repeat(), or rf\_spatial().

#### Usage

get\_residuals(model)

### Arguments

model

A model fitted with rf(), rf\_repeat(), or rf\_spatial().

### Value

A vector with model residuals, or the median of model residuals across repetitions if the model was fitted with rf\_repeat().

#### Examples

```
if(interactive()){
```

#load example data
data(plant\_richness\_df)

#fit random forest model

```
rf.model <- rf(
   data = plant_richness_df,
   dependent.variable.name = "richness_species_vascular",
   predictor.variable.names = colnames(plant_richness_df)[5:21],
   n.cores = 1,
   verbose = FALSE
)
#getting vector with residuals
x <- get_residuals(rf.model)
x
}</pre>
```

get\_response\_curves Gets data to allow custom plotting of response curves

#### Description

Generates and returns the data required to plot the response curves of a model fitted with rf(), rf\_repeat(), or rf\_spatial().

### Usage

```
get_response_curves(
  model = NULL,
  variables = NULL,
  quantiles = c(0.1, 0.5, 0.9),
  grid.resolution = 200,
  verbose = TRUE
)
```

### Arguments

model	A model fitted with rf(), rf_repeat(), or rf_spatial().	
variables	Character vector, names of predictors to plot. If NULL, the most important variables (importance higher than the median) in model are selected. Default: NULL.	
quantiles	Numeric vector with values between 0 and 1, argument probs of quantile. Quantiles to set the other variables to. Default: $c(0.1, 0.5, 0.9)$	
grid.resolution		
	Integer between 20 and 500. Resolution of the plotted curve Default: 100	
verbose	Logical, if TRUE the plot is printed. Default: TRUE	

#### Details

All variables that are not plotted in a particular response curve are set to the values of their respective quantiles, and the response curve for each one of these quantiles is shown in the plot.

### Value

A data frame with the following columns:

- response: Predicted values of the response, obtained with stats::predict().
- predictor: Values of the given predictor.
- quantile: Grouping column, values of the quantiles at which the other predictors are set to generate the response curve.
- model: Model number, only relevant if the model was produced with rf\_repeat().
- predictor.name: Grouping variable, name of the predictor.
- response.name: Grouping variable, name of the response variable.

### See Also

plot\_response\_curves()

### Examples

```
if(interactive()){
```

#loading example data
data(plant\_richness\_df)

```
#fitting random forest model
out <- rf(
    data = plant_richness_df,
    dependent.variable.name = "richness_species_vascular",
    predictor.variable.names = colnames(plant_richness_df)[5:21],
    n.cores = 1,
    verbose = FALSE
)
#getting data frame with response curves
p <- get_response_curves(out)
head(p)
}
```

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get\_spatial\_predictors

Gets the spatial predictors of a spatial model

### Description

Returns spatial predictors from a model fitted with rf\_spatial() in order of importance.

#### Usage

get\_spatial\_predictors(model)

### Arguments

model

A model fitted with rf\_spatial().

### Value

A data frame with the spatial predictors included in the model.

### Examples

```
if(interactive()){
 #loading example data
 data(distance_matrix)
 data(plant_richness_df)
 #fittind spatial model
 model <- rf_spatial(</pre>
  data = plant_richness_df,
  dependent.variable.name = "richness_species_vascular",
  predictor.variable.names = colnames(plant_richness_df)[5:21],
  distance.matrix = distance_matrix,
  distance.thresholds = c(0, 1000),
  n.cores = 1,
  method = "mem.moran.sequential"
)
#getting data frame with the selected spatial predictors
spatial.predictors <- get_spatial_predictors(model)</pre>
head(spatial.predictors)
}
```

is\_binary

Checks if dependent variable is binary with values 1 and 0

#### Description

Returns TRUE if dependent.variable.name is a binary variable with the values 1 and 0.

#### Usage

is\_binary(data = NULL, dependent.variable.name = NULL)

#### Arguments

data Data frame with a response variable and a set of predictors. Default: NULL dependent.variable.name

Character string with the name of the response variable. Must be in the column names of data. Default: NULL

make\_spatial\_fold

#### Value

Logical.

### Examples

```
if(interactive()){
    #dummy data frame
    data <- data.frame(
      response = c(0, 0, 0, 1, 1)
)
    #checking if response is binary
    is_binary(
      data = data,
      dependent.variable.name = "response"
)
}</pre>
```

make\_spatial\_fold Makes one training and one testing spatial folds

### Description

Used internally by make\_spatial\_folds() and rf\_evaluate(). Uses the coordinates of a point xy.i to generate two spatially independent data folds from the data frame xy. It does so by growing a rectangular buffer from xy.i until a number of records defined by training.fraction is inside the buffer. The indices of these records are then stored as "training" in the output list. The indices of the remaining records outside of the buffer are stored as "testing". These training and testing records can be then used to evaluate a model on independent data via cross-validation.

### Usage

```
make_spatial_fold(
    data = NULL,
    dependent.variable.name = NULL,
    xy.i = NULL,
    xy = NULL,
    distance.step.x = NULL,
    distance.step.y = NULL,
    training.fraction = 0.8
)
```

### Arguments

data	Data frame with a response variable and a set of predictors. Default: NULL	
dependent.variable.name		
	Character string with the name of the response variable. Must be in the column names of data. Default: NULL $\ensuremath{NULL}$	
xy.i	One row data frame with at least three columns: "x" (longitude), "y" (latitude), and "id" (integer, id of the record). Can be a row of xy. Default: NULL.	
ху	A data frame with at least three columns: "x" (longitude), "y" (latitude), and "id" (integer, index of the record). Default: NULL.	
distance.step.x		
	Numeric, distance step used during the growth in the x axis of the buffers defin- ing the training folds. Default: NULL (1/1000th the range of the x coordinates).	
distance.step.y		
	Numeric, distance step used during the growth in the y axis of the buffers defin- ing the training folds. Default: NULL (1/1000th the range of the y coordinates).	
training.fraction		
	Numeric, fraction of the data to be included in the training fold, Default: 0.8.	

### Value

A list with two slots named training and testing with the former having the indices of the training records selected from xy, and the latter having the indices of the testing records.

### See Also

make\_spatial\_folds(), rf\_evaluate()

### Examples

```
if(interactive()){
    #loading example data
    data(plant_richness_df)
    #getting case coordinates
    xy <- plant_richness_df[, 1:3]
    colnames(xy) <- c("id", "x", "y")
    #building a spatial fold centered in the first pair of coordinates
    out <- make_spatial_fold(
        xy.i = xy[1, ],
        xy = xy,
        training.fraction = 0.6
    )
    #indices of the training and testing folds
    out$training
    out$testing</pre>
```

#### make\_spatial\_folds

```
#plotting the data
plot(xy[ c("x", "y")], type = "n", xlab = "", ylab = "")
#plots training points
points(xy[out$training, c("x", "y")], col = "red4", pch = 15)
#plots testing points
points(xy[out$testing, c("x", "y")], col = "blue4", pch = 15)
#plots xy.i
points(xy[1, c("x", "y")], col = "black", pch = 15, cex = 2)
}
```

make\_spatial\_folds Makes training and testing spatial folds

### Description

Applies make\_spatial\_fold() to every record in a data frame xy.selected to generate as many spatially independent folds over the dataset xy as rows are in xy.selected.

### Usage

```
make_spatial_folds(
    data = NULL,
    dependent.variable.name = NULL,
    xy.selected = NULL,
    xy = NULL,
    distance.step.x = NULL,
    training.fraction = 0.75,
    n.cores = parallel::detectCores() - 1,
    cluster = NULL
)
```

### Arguments

data	Data frame with a response variable and a set of predictors. Default: NULL	
dependent.variable.name		
	Character string with the name of the response variable. Must be in the column names of data. Default: NULL	
xy.selected	Data frame with at least three columns: "x" (longitude), "y" (latitude), and "id" (integer, id of the record). Usually a subset of xy. Usually the result of applying thinning() or thinning_til_n() to 'xy' Default: NULL.	
ху	data frame with at least three columns: "x" (longitude), "y" (latitude), and "id" (integer, index of the record). Default: NULL.	
distance.step.x		
	Numeric, distance step used during the growth in the x axis of the buffers defin- ing the training folds. Default: NULL (1/1000th the range of the x coordinates).	

distance.step.y	
	Numeric, distance step used during the growth in the y axis of the buffers defin- ing the training folds. Default: NULL (1/1000th the range of the y coordinates).
training.fracti	on
	numeric, fraction of the data to be included in the growing buffer as training data, Default: $0.75$
n.cores	Integer, number of cores to use for parallel execution. Creates a socket cluster with parallel::makeCluster(), runs operations in parallel with foreach and %dopar%, and stops the cluster with parallel::clusterStop() when the job is done. Default: parallel::detectCores() - 1
cluster	A cluster definition generated with parallel::makeCluster(). If provided, overrides n.cores. When cluster = NULL (default value), and model is provided, the cluster in model, if any, is used instead. If this cluster is NULL, then the function uses n.cores instead. The function does not stop a provided cluster, so it should be stopped with parallel::stopCluster() afterwards. The cluster definition is stored in the output list under the name "cluster" so it can be passed to other functions via the model argument, or using the %>% pipe. Default: NULL

### Value

A list with as many slots as rows are in xy.selected. Each slot has two slots named training and testing, with the former having the indices of the training records selected from xy, and the latter having the indices of the testing records.

#### See Also

make\_spatial\_fold(), rf\_evaluate()

### Examples

if(interactive()){

```
#loading example data
data(plant_richness_df)
#getting case coordinates
xy <- plant_richness_df[, 1:3]
colnames(xy) <- c("id", "x", "y")
#thining til 20 cases
xy.selected <- thinning_til_n(
  xy = xy,
  n = 20
  )
#making spatial folds centered on these 20 cases
out <- make_spatial_folds(
  xy.selected = xy.selected,
  xy = xy,
```

```
distance.step = 0.05, #degrees
training.fraction = 0.6,
n.cores = 1
)
#plotting training and testing folds
plot(xy[ c("x", "y")], type = "n", xlab = "", ylab = "")
#plots training points
points(xy[out[[10]]$training, c("x", "y")], col = "red4", pch = 15)
#plots testing points
points(xy[out[[10]]$testing, c("x", "y")], col = "blue4", pch = 15)
#plots xy.i
points(xy[10, c("x", "y")], col = "black", pch = 15, cex = 2)
}
```

mem

Moran's Eigenvector Maps of a distance matrix

#### Description

Computes the positive Moran's Eigenvector Maps of a distance matrix.

#### Usage

```
mem(
  distance.matrix = NULL,
  distance.threshold = 0,
  colnames.prefix = "mem"
)
```

#### Arguments

distance.matrix

Distance matrix. Default: NULL.

```
distance.threshold
```

Numeric vector with distance thresholds defining different neighborhood extents within the distance matrix, Default: 0

colnames.prefix

Character, name prefix for the output columns. Default: "mem"

### Details

Takes the distance matrix x, double-centers it with double\_center\_distance\_matrix(), applies eigen, and returns eigenvectors with positive normalized eigenvalues (a.k.a Moran's Eigenvector Maps, or MEMs). These MEMs are later used as spatial predictors by rf\_spatial().

#### Value

A data frame with positive Moran's Eigenvector Maps.

#### See Also

```
mem_multithreshold(), rf_spatial()
```

#### Examples

```
if(interactive()){
```

```
#loading example distance matrix
data(distance_matrix)
#Moran's Eigenvector Maps of the distance matrix
mem <- mem(x = distance_matrix)</pre>
```

}

mem\_multithreshold Moran's Eigenvector Maps for different distance thresholds

#### Description

Computes Moran's Eigenvector Maps of a distance matrix (using mem()) over different distance thresholds.

#### Usage

```
mem_multithreshold(
   distance.matrix = NULL,
   distance.thresholds = NULL,
   max.spatial.predictors = NULL
)
```

### Arguments

distance.matrix

Distance matrix. Default: NULL.

distance.thresholds

Numeric vector with distance thresholds defining neighborhood in the distance matrix, Default: NULL.

max.spatial.predictors

Maximum number of spatial predictors to generate. Only useful to save memory when the distance matrix x is very large. Default: NULL.

#### **Details**

The function takes the distance matrix x, computes its weights at difference distance thresholds, double-centers the resulting weight matrices with double\_center\_distance\_matrix(), applies eigen to each double-centered matrix, and returns eigenvectors with positive normalized eigenvalues for different distance thresholds.

#### moran

### Value

A data frame with as many rows as the distance matrix x containing positive Moran's Eigenvector Maps. The data frame columns are named "spatial\_predictor\_DISTANCE\_COLUMN", where DISTANCE is the given distance threshold, and COLUMN is the column index of the given spatial predictor.

### Examples

```
if(interactive()){
    #loading example data
    data(distance_matrix)
    #computing Moran's eigenvector maps for 0, 1000, and 2000 km
    mem.df <- mem_multithreshold(
        distance.matrix = distance_matrix,
        distance.thresholds = c(0, 1000, 2000)
        )
    head(mem.df)
}</pre>
```

moran

Moran's I test

#### Description

Computes the spatial correlation coefficient (Moran's I) of a vector given a distance matrix, and a distance threshold used to define "neighborhood".

### Usage

```
moran(
  x = NULL,
  distance.matrix = NULL,
  distance.threshold = NULL,
  verbose = TRUE
)
```

#### Arguments

```
x Numeric vector, generally model residuals, Default: NULL
distance.matrix
Distance matrix among cases in x. The number of rows of this matrix must be
equal to the length of x. Default: NULL
distance.threshold
numeric value in the range of values available in distance.matrix. Distances
below such threshold are set to 0. Default: NULL (which defaults to 0).
verbose
Logical, if TRUE, prints a Moran's I plot. Default: TRUE
```

#### Details

Inspired in the Moran. I() function of the ape package.

### Value

A list with three named slots:

- test: Data frame with observed and expected Moran's I values, p-value, and interpretation.
- plot: Moran's plot of the vector x against the spatial lags of x.
- plot.df: Data used in the Moran's plot.

### See Also

moran\_multithreshold()

### Examples

```
if(interactive()){
    #loading example data
    data(distance_matrix)
    data(plant_richness)

    #Moran's I of the response variable
    out <- moran(
        x = plant_richness$richness_species_vascular,
        distance.matrix = distance_matrix
        )
    out
}</pre>
```

moran\_multithreshold Moran's I test on a numeric vector for different neighborhoods

### Description

Applies moran() to different distance thresholds at the same time.

### Usage

```
moran_multithreshold(
    x = NULL,
    distance.matrix = NULL,
    distance.thresholds = NULL,
    verbose = TRUE
)
```

#### Arguments

х	Numeric vector, generally model residuals, Default: NULL	
distance.matri	Х	
	Distance matrix among cases in x. The number of rows of this matrix must be equal to the length of x. Default: NULL	
distance.thresholds		
	Numeric vector, distances below each value are set to 0 on separated copies of the distance matrix for the computation of Moran's I at different neigh- borhood distances. If NULL, it defaults to seq(0, max(distance.matrix)/4, length.out = 2). Default: NULL	
verbose	Logical, if TRUE, plots Moran's I values for each distance threshold. Default: TRUE	

### Details

Using different distance thresholds helps to take into account the uncertainty about what "neighborhood" means in ecological systems (1000km in geological time means little, but 100m might be quite a long distance for a tree to disperse seeds over), and allows to explore spatial autocorrelation of model residuals for several minimum-distance criteria at once.

### Value

A named list with the slots:

- df: Data frame with the results of moran per distance threshold.
- plot: A plot of Moran's I across distance thresholds.
- max.moran: Maximum value of Moran's I across thresholds.
- max.moran.distance.threshold: Distance threshold with the maximum Moran's I value.

#### See Also

moran()

#### Examples

```
if(interactive()){
```

```
#loading example data
data(distance_matrix)
data(plant_richness)
```

```
#computing Moran's I for the response variable at several reference distances
out <- moran_multithreshold(
    x = plant_richness$richness_species_vascular,
    distance.matrix = distance_matrix,
    distance.thresholds = c(0, 100, 1000, 10000),
    plot = TRUE
    )</pre>
```

```
out
```

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objects\_size

Shows size of objects in the R environment

### Description

Shows the size of the objects currently in the R environment. Helps to locate large objects cluttering the R environment and/or causing memory problems during the execution of large workflows.

#### Usage

 $objects_size(n = 10)$ 

### Arguments

n

Number of objects to show, Default: 10

### Value

A data frame with the row names indicating the object name, the field 'Type' indicating the object type, 'Size' indicating the object size, and the columns 'Length/Rows' and 'Columns' indicating the object dimensions if applicable.

#### Examples

```
if(interactive()){
    #creating dummy objects
    x <- matrix(runif(100), 10, 10)
    y <- matrix(runif(10000), 100, 100)
    #reading their in-memory size
    objects_size()</pre>
```

}

optimization\_function Optimization equation to select spatial predictors

### Description

Optimizes the selection of spatial predictors using two different methods: "moran.i", and "p.value".

### Usage

```
optimization_function(
  x = NULL,
  weight.r.squared = NULL,
  weight.penalization.n.predictors = NULL,
  optimization.method = "moran.i"
)
```

### Arguments

x	Optimization data frame generated internally by select_spatial_predictors_sequential() or select_spatial_predictors_recursive(). Default: NULL
weight.r.squared	
	Numeric between 0 and 1, weight of R-squared in the optimization process.
	Default: NULL
<pre>weight.penalization.n.predictors</pre>	
	Numeric between 0 and 1, weight of the penalization on the number of added
	spatial predictors. Default: NULL
optimization.method	
	Character, one of "moran.i", and "p.value". Default: "moran.i"

### Details

The method "moran.i" tries to maximize 1 - Moran's I while taking into account the R-squared of the model and a penalization on the number of introduced spatial predictors through the expression

(1 - Moran's I) + w1 \* r.squared - w2 \* penalization

The method "p.value" uses a binary version of the p-values of Moran's I (1 if  $\geq 0.05$ , 0 otherwise), and uses the expression

max(1 - Moran's I, binary p-value) + w1 \* r.squared - w2 \* penalization

The "moran.i" method generally selects more spatial predictors than the "p.value" method.

#### Value

A numeric vector with the optimization criteria.

### See Also

select\_spatial\_predictors\_recursive(), select\_spatial\_predictors\_sequential()

### Description

Extracts all factors of a principal component analysis of a matrix or data frame. Just a convenient wrapper for prcomp.

### Usage

```
pca(
  x = NULL,
  colnames.prefix = "pca_factor"
)
```

### Arguments

x numeric matrix or data frame, Default: NULL colnames.prefix

character, name prefix for the output columns, Default: 'pca\_factor'

### Details

Columns in x with zero variance are removed before computing the PCA.

### Value

A data frame with the PCA factors of x.

### See Also

pca\_multithreshold()

### Examples

```
if(interactive()){
```

#load example distance matrix
data(distance\_matrix)

#PCA of the distance matrix
out <- pca(x = distance\_matrix)
out</pre>

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### рса

pca\_multithreshold PCA of a distance matrix over distance thresholds

#### Description

Computes PCA factors of a distance matrix over different distance thresholds to generate spatial predictors for a model fitted with rf\_spatial().

### Usage

```
pca_multithreshold(
  distance.matrix = NULL,
  distance.thresholds = NULL,
  max.spatial.predictors = NULL
)
```

### Arguments

```
distance.matrix
```

Distance matrix. Default: NULL

distance.thresholds

Numeric vector with distance thresholds defining neighborhood in the distance matrix, Default: 0

max.spatial.predictors

Integer, maximum number of spatial predictors to generate. Only useful when the distance matrix x is very large. Default: NULL

#### Details

The distance matrix is converted into weights with weights\_from\_distance\_matrix() before computing the PCA. This produces more meaningful spatial predictors than using the distance matrix as is.

### Value

A data frame with the PCA factors of the thresholded matrix. The data frame columns are named "spatial\_predictor\_DISTANCE\_COLUMN", where DISTANCE is the given distance threshold, and COLUMN is the column index of the given predictor.

### See Also

pca()

### Examples

```
if(interactive()){
    #loading example distance matrix
    load(distance_matrix)
    #PCA factors of the distance matrix for two reference distances
    x <- pca_multithreshold(
        distance.matrix = distance_matrix,
        distance.thresholds = c(0, 1000)
        )
    head(x)
}</pre>
```

plant\_richness\_df Plant richness and predictors of American ecoregions

### Description

Richness of vascular plants of the American ecoregions as defined in Ecoregions 2017.

#### Usage

```
data(plant_richness_df)
```

### Format

A data frame with 227 rows and 22 columns:

- ecoregion\_id: Id of the ecoregion).
- x: Longitude in degrees (WGS84).
- y: Latitude in degrees (WGS84).
- richness\_species\_vascular: Number of vascular species found in the ecoregion. Response variable.
- bias\_area\_km2: Area of the ecoregion in squared kilometers.
- bias\_species\_per\_record: Number of species divided by the number of spatial GBIF records available in the ecoregion as a measure of sampling bias.
- climate\_aridity\_index\_average: Average of the ecoregion.
- climate\_hypervolume: Volume of the climatic envelope of the ecoregion, computed with the hypervolume package.
- climate\_velocity\_lgm\_average: Average climate velocity of the ecoregion since the Last Glacial Maximum.
- neighbors\_count: Number of immediate neighbors of the ecoregion as a measure of connectivity/isolation.

#### plot\_evaluation

- neighbors\_percent\_shared\_edge: Percentage of shared edge with the neighbors as a measure of connectivity/isolation.
- human\_population\_density: Population density of the ecoregion.
- topography\_elevation\_average: Average elevation of the ecoregion.
- landcover\_herbs\_percent\_average: Average cover percentage of herbs extracted from MODIS Vegetation Continuous Fields.
- fragmentation\_cohesion: Geographic fragmentation index of the ecoregion as computed with the R package landscapemetrics.
- fragmentation\_division: Another fragmentation index.
- neighbors\_area: Total area of the ecoregions's immediate neighbors.
- human\_population: Human population in the ecoregion.
- human\_footprint\_average: Average human footprint in the ecoregion.
- climate\_bio1\_average: Average mean annual temperature.
- climate\_bio15\_minimum: Average precipitation seasonality.

## See Also

distance\_matrix

plot\_evaluation *Plots the results of a spatial cross-validation* 

#### Description

Plots the results of an spatial cross-validation performed with rf\_evaluate().

#### Usage

```
plot_evaluation(
  model,
  fill.color = viridis::viridis(
    3,
    option = "F",
    alpha = 0.8,
    direction = -1
    ),
  line.color = "gray30",
  verbose = TRUE,
  notch = TRUE
)
```

#### Arguments

model	A model resulting from rf_evaluate().
fill.color	Character vector with three hexadecimal codes (e.g. "#440154FF" "#21908CFF" "#FDE725FF"), or function generating a palette (e.g. viridis::viridis(3)). Default: viridis::viridis(3, option = "F", alpha = 0.8, direction = -1)
line.color	Character string, color of the line produced by ggplot2::geom_smooth(). De-fault: "gray30"
verbose	Logical, if TRUE the plot is printed. Default: TRUE
notch	Logical, if TRUE, boxplot notches are plotted. Default: TRUE

#### Value

A ggplot.

## See Also

rf\_evaluate(), get\_evaluation(), print\_evaluation().

#### Examples

```
if(interactive()){
```

```
#loading example data
data(plant_richness_df)
data(distance_matrix)
#fitting a random forest model
rf.model <- rf(</pre>
  data = plant_richness_df,
  dependent.variable.name = "richness_species_vascular",
  predictor.variable.names = colnames(plant_richness_df)[5:21],
  distance.matrix = distance_matrix,
  distance.thresholds = 0,
  n.cores = 1,
  verbose = FALSE
)
#evaluating the model with spatial cross-validation
rf.model <- rf_evaluate(</pre>
 model = rf.model,
  xy = plant_richness_df[, c("x", "y")],
  n.cores = 1
)
#plotting the evaluation results
plot_evaluation(rf.model)
```

}

#### Description

Plots variable importance scores of rf(), rf\_repeat(), and rf\_spatial() models. Distributions of importance scores produced with rf\_repeat() are plotted using ggplot2::geom\_violin, which shows the median of the density estimate rather than the actual median of the data. However, the violin plots are ordered from top to bottom by the real median of the data to make small differences in median importance easier to spot. Ths function does not plot the result of rf\_importance() yet, but you can find it under model\$importance\$cv.per.variable.plot.

# Usage

```
plot_importance(
  model,
  fill.color = viridis::viridis(
    100,
    option = "F",
    direction = -1,
    alpha = 1,
    end = 0.9
  ),
  line.color = "white",
   verbose = TRUE
)
```

## Arguments

model	A model fitted with rf(), rf_repeat(), or rf_spatial(), or a data frame with variable importance scores (only for internal use within the package functions).
	Character vector with hexadecimal codes (e.g. "#440154FF" "#21908CFF" "#FDE725FF"), or function generating a palette (e.g. viridis::viridis(100)). Default: viridis::viridis(100, option = "F", direction = -1, alpha = 0.8, end = 0.9)
line.color	Character string, color of the line produced by ggplot2::geom_smooth(). De-fault: "white"
verbose	Logical, if TRUE, the plot is printed. Default: TRUE

## Value

A ggplot.

#### See Also

print\_importance(), get\_importance()

## Examples

```
if(interactive()){
```

```
#loading example data
data(plant_richness_df)
data(distance_matrix)
#fitting a random forest model
rf.model <- rf(</pre>
 data = plant_richness_df,
 dependent.variable.name = "richness_species_vascular",
 predictor.variable.names = colnames(plant_richness_df)[5:21],
 distance.matrix = distance_matrix,
 distance.thresholds = 0,
 n.cores = 1,
 verbose = FALSE
)
#plotting variable importance scores
plot_importance(model = rf.model)
}
```

plot\_moran

#### Plots a Moran's I test of model residuals

#### Description

Plots the results of spatial autocorrelation tests for a variety of functions within the package. The x axis represents the Moran's I estimate, the y axis contains the values of the distance thresholds, the dot sizes represent the p-values of the Moran's I estimate, and the red dashed line represents the theoretical null value of the Moran's I estimate.

#### Usage

```
plot_moran(
   model,
   point.color = viridis::viridis(
      100,
      option = "F",
      direction = -1
    ),
   line.color = "gray30",
   option = 1,
   ncol = 1,
   verbose = TRUE
)
```

## plot\_moran

## Arguments

model	A model fitted with rf(), rf_repeat(), or rf_spatial(), or a data frame generated by moran(). Default: NULL
point.color	Colors of the plotted points. Can be a single color name (e.g. "red4"), a character vector with hexadecimal codes (e.g. "#440154FF" "#21908CFF" "#FDE725FF"), or function generating a palette (e.g. viridis::viridis(100)). Default: viridis::viridis(100, option = "F")
line.color	Character string, color of the line produced by ggplot2::geom_smooth(). De-fault: "gray30"
option	Integer, type of plot. If 1 (default) a line plot with Moran's I and p-values across distance thresholds is returned. If 2, scatterplots of residuals versus lagged residuals per distance threshold and their corresponding slopes are returned. In models fitted with rf_repeat(), the residuals and lags of the residuals are computed from the median residuals across repetitions. Option 2 is disabled if x is a data frame generated by moran().
ncol	Number of columns of the plot. Only relevant when option = 2. Argument ncol of wrap_plots.
verbose	Logical, if TRUE, the resulting plot is printed, Default: TRUE

## Value

A ggplot.

# See Also

moran(), moran\_multithreshold()

```
if(interactive()){
#loading example data
data(plant_richness_df)
data(distance.matrix)
#fitting a random forest model
rf.model <- rf(</pre>
  data = plant_richness_df,
  dependent.variable.name = "richness_species_vascular",
  predictor.variable.names = colnames(plant_richness_df)[5:21],
  distance.matrix = distance_matrix,
  distance.thresholds = c(0, 1000, 2000),
  n.cores = 1,
  verbose = FALSE
)
#Incremental/multiscale Moran's I
plot_moran(rf.model)
```

```
#Moran's scatterplot
plot_moran(rf.model, option = 2)
}
```

plot\_optimization Optimization plot of a selection of spatial predictors

# Description

Plots optimization data frames produced by select\_spatial\_predictors\_sequential() and select\_spatial\_predictors\_recursive().

#### Usage

```
plot_optimization(
   model,
   point.color = viridis::viridis(
       100,
       option = "F",
       direction = -1
   ),
   verbose = TRUE
)
```

## Arguments

model	A model produced by rf_spatial(), or an optimization data frame produced by select_spatial_predictors_sequential() or select_spatial_predictors_recursive().
point.color	Colors of the plotted points. Can be a single color name (e.g. "red4"), a character vector with hexadecimal codes (e.g. "#440154FF" "#21908CFF" "#FDE725FF"), or function generating a palette (e.g. viridis::viridis(100)). Default: viridis::viridis(100, option = "F", direction = -1)
verbose	Logical, if TRUE the plot is printed. Default: TRUE

# Details

If the method used to fit a model with rf\_spatial() is "hengl", the function returns nothing, as this method does not require optimization.

## Value

A ggplot.

#### Examples

```
if(interactive()){
#loading example data
data(distance_matrix)
 data(plant_richness_df)
 #names of the response and predictors
 dependent.variable.name <- "richness_species_vascular"</pre>
 predictor.variable.names <- colnames(plant_richness_df)[5:21]</pre>
#spatial model
model <- rf_spatial(</pre>
  data = plant_richness_df,
  dependent.variable.name = dependent.variable.name,
  predictor.variable.names = predictor.variable.names,
  distance.matrix = distance_matrix,
  distance.thresholds = 0,
  method = "mem.moran.sequential",
  n.cores = 1,
  seed = 1
```

```
#plotting selection of spatial predictors
plot_optimization(model = model)
```

```
}
```

)

plot\_residuals\_diagnostics *Plot residuals diagnostics* 

#### Description

Plots normality and autocorrelation tests of model residuals.

#### Usage

```
plot_residuals_diagnostics(
    model,
    point.color = viridis::viridis(100, option = "F"),
    line.color = "gray10",
    fill.color = viridis::viridis(4, option = "F", alpha = 0.95)[2],
    option = 1,
    ncol = 1,
    verbose = TRUE
)
```

## Arguments

model	A model produced by rf(), rf_repeat(), or rf_spatial().
point.color	Colors of the plotted points. Can be a single color name (e.g. "red4"), a character vector with hexadecimal codes (e.g. "#440154FF" "#21908CFF" "#FDE725FF"), or function generating a palette (e.g. viridis::viridis(100)). Default: viridis::viridis(100, option = "F")
line.color	Character string, color of the line produced by ggplot2::geom_smooth(). De-fault: "gray30"
fill.color	Character string, fill color of the bars produced by ggplot2::geom_histogram(). Default: viridis::viridis(4, option = "F", alpha = 0.95)[2]
option	(argument of plot_moran()) Integer, type of plot. If 1 (default) a line plot with Moran's I and p-values across distance thresholds is returned. If 2, scatterplots of residuals versus lagged residuals per distance threshold and their correspond- ing slopes are returned. In models fitted with rf_repeat(), the residuals and lags of the residuals are computed from the median residuals across repetitions. Option 2 is disabled if x is a data frame generated by moran().
ncol	(argument of plot_moran()) Number of columns of the Moran's I plot if option = 2.
verbose	Logical, if TRUE, the resulting plot is printed, Default: TRUE

#### Value

A patchwork object.

```
if(interactive()){
    #load example data
    data(plant_richness_df)
    data(distance_matrix)

    #fit a random forest model
    x <- rf(
        data = plant_richness_df,
        dependent.variable.name = "richness_species_vascular",
        predictor.variable.names = colnames(plant_richness_df)[5:21],
        distance.matrix = distance_matrix,
        n.cores = 1
    )

    #residuals diagnostics
    plot_residuals_diagnostics(x)
}</pre>
```

plot\_response\_curves *Plots the response curves of a model.* 

## Description

Plots the response curves of models fitted with rf(), rf\_repeat(), or rf\_spatial().

#### Usage

```
plot_response_curves(
  model = NULL,
  variables = NULL,
  quantiles = c(0.1, 0.5, 0.9),
  grid.resolution = 200,
  line.color = viridis::viridis(length(quantiles), option = "F", end = 0.9),
  ncol = 2,
  show.data = FALSE,
  verbose = TRUE
)
```

#### Arguments

model	A model fitted with rf(), rf_repeat(), or rf_spatial().
variables	Character vector, names of predictors to plot. If NULL, the most important variables (importance higher than the median) in x are selected. Default: NULL.
quantiles	Numeric vector with values between 0 and 1, argument probs of quantile. Quantiles to set the other variables to. Default: $c(0.1, 0.5, 0.9)$
grid.resolutio	n
	Integer between 20 and 500. Resolution of the plotted curve Default: 100
line.color	Character vector with colors, or function to generate colors for the lines repre- senting quantiles. Must have the same number of colors as quantiles are de- fined. Default: viridis::viridis(length(quantiles), option = "F", end = 0.9)
ncol	Integer, argument of wrap_plots. Defaults to the rounded squared root of the number of plots. Default: 2
show.data	Logical, if TRUE, the observed data is plotted along with the response curves. Default: FALSE
verbose	Logical, if TRUE the plot is printed. Default: TRUE

#### Details

All variables that are not plotted in a particular response curve are set to the values of their respective quantiles, and the response curve for each one of these quantiles is shown in the plot. When the input model was fitted with rf\_repeat() with keep.models = TRUE, then the plot shows the median of all model runs, and each model run separately as a thinner line. The output list can be plotted all at once with patchwork::wrap\_plots(p) or cowplot::plot\_grid(plotlist = p), or one by one by extracting each plot from the list.

A list with slots named after the selected variables, with one ggplot each.

#### See Also

plot\_response\_surface()

# Examples

```
if(interactive()){
#loading example data
data(plant_richness_df)
#fitting a random forest model
m <- rf(
   data = plant_richness_df,
   dependent.variable.name = "richness_species_vascular",
   predictor.variable.names = colnames(plant_richness_df)[5:21],
   n.cores = 1,
   verbose = FALSE
)
#response curves of most important predictors
plot_response_curves(model = m)
}</pre>
```

plot\_response\_surface *Plots the response surfaces of a random forest model* 

## Description

Plots response surfaces for any given pair of predictors in a rf(), rf\_repeat(), or rf\_spatial() model.

## Usage

```
plot_response_surface(
  model = NULL,
  a = NULL,
  b = NULL,
  quantiles = 0.5,
  grid.resolution = 100,
  point.size.range = c(0.5, 2.5),
  point.alpha = 1,
  fill.color = viridis::viridis(100, option = "F", direction = -1, alpha = 0.9),
  point.color = "gray30",
  verbose = TRUE
)
```

#### Arguments

model	A model fitted with rf(), rf_repeat(), or rf_spatial(). Default NULL
а	Character string, name of a model predictor. If NULL, the most important variable in model is selected. Default: NULL
b	Character string, name of a model predictor. If NULL, the second most important variable in model is selected. Default: NULL
quantiles	Numeric vector between 0 and 1. Argument probs of the function quantile. Quantiles to set the other variables to. Default: 0.5
grid.resolutio	on
	Integer between 20 and 500. Resolution of the plotted surface Default: 100
point.size.range	
	Numeric vector of length 2 with the range of point sizes used by geom_point. Using $c(-1, -1)$ removes the points. Default: $c(0.5, 2.5)$
point.alpha	Numeric between 0 and 1, transparency of the points. Setting it to 0 removes all points. Default: 1.
fill.color	Character vector with hexadecimal codes (e.g. "#440154FF" "#21908CFF" "#FDE725FF"), or function generating a palette (e.g. viridis::viridis(100)). Default: viridis::viridis(100, option = "F", direction = -1, alpha = 0.9)
point.color	Character vector with a color name (e.g. "red4"). Default: gray30
verbose	Logical, if TRUE the plot is printed. Default: TRUE

# Details

All variables that are not a or b in a response curve are set to the values of their respective quantiles to plot the response surfaces. The output list can be plotted all at once with patchwork::wrap\_plots(p) or cowplot::plot\_grid(plotlist = p), or one by one by extracting each plot from the list.

# Value

A list with slots named after the selected quantiles, each one with a ggplot.

#### See Also

plot\_response\_curves()

```
if(interactive()){
    #load example data
    data(plant_richness_df)
    #fit random forest model
    out <- rf(
        data = plant_richness_df,
        dependent.variable.name = "richness_species_vascular",</pre>
```

```
n.cores = 1,
verbose = FALSE
)
#plot interactions between most important predictors
plot_response_surfaces(x = out)
}
```

plot\_training\_df Scatterplots of a training data frame

## Description

Plots the dependent variable against each predictor.

## Usage

```
plot_training_df(
    data = NULL,
    dependent.variable.name = NULL,
    predictor.variable.names = NULL,
    ncol = 4,
    method = "loess",
    point.color = viridis::viridis(100, option = "F"),
    line.color = "gray30"
)
```

# Arguments

data	Data frame with a response variable and a set of predictors. Default: NULL
dependent.vari	able.name
	Character string with the name of the response variable. Must be in the column names of data. If the dependent variable is binary with values 1 and 0, the argument case.weights of ranger is populated by the function case_weights().
predictor.vari	Default: NULL able_names
p. cu cu	Character vector with the names of the predictive variables. Every element of this vector must be in the column names of data. Optionally, the result of auto_cor() or auto_vif() Default: NULL
ncol	Number of columns of the plot. Argument ncol of wrap_plots.
method	Method for geom_smooth, one of: "lm", "glm", "gam", "loess", or a function, for example mgcv::gam Default: 'loess'
point.color	Colors of the plotted points. Can be a single color name (e.g. "red4"), a character vector with hexadecimal codes (e.g. "#440154FF" "#21908CFF" "#FDE725FF"), or function generating a palette (e.g. viridis::viridis(100)). Default: viridis::viridis(100, option = "F")

line.color Character string, color of the line produced by ggplot2::geom\_smooth(). Default: "gray30"

#### Value

A wrap\_plots object.

## Examples

```
if(interactive()){
    #load example data
    data(plant_richness_df)
    #scatterplot of the training data
    plot_training_data(
        data = plant_richness_df,
        dependent.variable.name = "richness_species_vascular",
        predictor.variable.names = colnames(plant_richness_df)[5:21]
        )
}
```

plot\_training\_df\_moran

```
Moran's I plots of a training data frame
```

#### Description

Plots the the Moran's I test of the response and the predictors in a training data frame.

# Usage

```
plot_training_df_moran(
  data = NULL,
  dependent.variable.name = NULL,
  predictor.variable.names = NULL,
  distance.matrix = NULL,
  distance.thresholds = NULL,
  fill.color = viridis::viridis(100, option = "F", direction = -1),
  point.color = "gray30"
)
```

#### Arguments

data Data frame with a response variable and a set of predictors. Default: NULL

	dependent.variable.name	
		Character string with the name of the response variable. Must be in the column names of data. If the dependent variable is binary with values 1 and 0, the argument case.weights of ranger is populated by the function case_weights(). Default: NULL
predictor.variable.names		able.names
		Character vector with the names of the predictive variables. Every element of this vector must be in the column names of data. Optionally, the result of auto_cor() or auto_vif() Default: NULL
distance.matrix		< compared by the second se
		Squared matrix with the distances among the records in data. The number of rows of distance.matrix and data must be the same. If not provided, the computation of the Moran's I of the residuals is omitted. Default: NULL
distance.thresholds		nolds
		Numeric vector, distances below each value are set to 0 on separated copies of the distance matrix for the computation of Moran's I at different neigh- borhood distances. If NULL, it defaults to seq(0, max(distance.matrix)/4, length.out = 2). Default: NULL
	fill.color	Character vector with hexadecimal codes (e.g. "#440154FF" "#21908CFF" "#FDE725FF"), or function generating a palette (e.g. viridis::viridis(100)). Default: viridis::viridis(100, option = "F", direction = -1)
	point.color	Character vector with a color name (e.g. "red4"). Default: gray30

# Value

A ggplot2 object.

# Examples

}

```
if(interactive()){
  #load example data
  data(plant_richness_df)
  data(distance_matrix)
  #plot Moran's I of training data
  plot_moran_training_data(
    data = plant_richness_df,
    dependent.variable.name = "richness_species_vascular",
    predictor.variable.names = colnames(plant_richness_df)[5:21],
    distance.matrix = distance_matrix,
    distance.thresholds = c(
      0,
      2000,
      4000,
      6000,
      8000
      )
    )
```

plot\_tuning

#### Description

Plots the tuning of the hyperparameters num.trees, mtry, and min.node.size performed by rf\_tuning().

#### Usage

```
plot_tuning(
  model,
  point.color = viridis::viridis(
    100,
    option = "F"
  ),
  verbose = TRUE
)
```

#### Arguments

model	A model fitted with rf_tuning(). Default: NULL
point.color	Colors of the plotted points. Can be a single color name (e.g. "red4"), a character vector with hexadecimal codes (e.g. "#440154FF" "#21908CFF" "#FDE725FF"), or function generating a palette (e.g. viridis::viridis(100)). Default: viridis::viridis(100, option = "F")
verbose	Logical, if TRUE, the plot is printed. Default: TRUE

### Value

A ggplot.

# See Also

rf\_tuning()

```
if(interactive()){
```

```
#load example data
data(plant_richness_df)
#fit random forest model
rf.model <- rf(
   data = plant_richness_df,
   dependent.variable.name = "richness_species_vascular",
   predictor.variable.names = colnames(plant_richness_df)[5:21],</pre>
```

```
distance.matrix = distance_matrix,
distance.thresholds = 0,
n.cores = 1,
verbose = FALSE
)
#tune random forest model
rf.model <- rf_tuning(
model = rf.model,
xy = plant_richness_df[, c("x", "y")],
n.cores = 1,
verbose = FALSE
)
#generate tuning plot
plot_tuning(model = rf.model)
```

}

prepare\_importance\_spatial
Prepares variable importance objects for spatial models

#### Description

Prepares variable importance data frames and plots for models fitted with rf\_spatial().

#### Usage

```
prepare_importance_spatial(model)
```

#### Arguments

model

An importance data frame with spatial predictors, or a model fitted with rf\_spatial().

#### Value

A list with importance data frames in different formats depending on whether the model was fitted with rf() or  $rf_repeat()$ .

## Examples

```
if(interactive()){
    #loading example data
    data(distance_matrix)
    data(plant_richness_df)
    #fittind spatial model
    model <- rf_spatial(</pre>
```

## print.rf

```
data = plant_richness_df,
  dependent.variable.name = "richness_species_vascular",
    predictor.variable.names = colnames(plant_richness_df)[5:21],
    distance.matrix = distance_matrix,
    distance.thresholds = 0,
    n.cores = 1
)
#preparing the importance data frame
importance <- prepare_importance_spatial(model)
names(importance)
```

print.rf

}

#### *Custom print method for random forest models*

#### Description

Custom print method for models fitted with rf(), rf\_repeat(), and rf\_spatial().

#### Usage

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

#### Arguments

Х	A model fitted with rf(), rf_repeat(), or rf_spatial().
	Additional arguments for print methods.

#### Value

Prints model details to the console.

# See Also

print\_evaluation(), print\_importance(), print\_moran(), print\_performance()

```
if(interactive()){
    #loading example data
```

```
data("plant_richness_df")
data("distance_matrix")
#fitting random forest model
```

```
data = plant_richness_df,
  dependent.variable.name = "richness_species_vascular",
    predictor.variable.names = colnames(plant_richness_df)[5:21],
    distance.matrix = distance_matrix,
    distance.thresholds = 0,
    n.cores = 1
)
#printing model summary
print(rf.model)
}
```

print\_evaluation Prints cross-validation results

#### Description

Prints the results of an spatial cross-validation performed with rf\_evaluate().

## Usage

```
print_evaluation(model)
```

#### Arguments

model A model resulting from rf\_evaluate().

#### Value

A table printed to the standard output.

#### See Also

plot\_evaluation(), get\_evaluation()

## Examples

```
if(interactive()){
```

```
#loading example data
data(plant_richness_df)
data(distance_matrix)
```

```
#fitting random forest model
rf.model <- rf(
   data = plant_richness_df,
   dependent.variable.name = "richness_species_vascular",
   predictor.variable.names = colnames(plant_richness_df)[5:21],
   distance.matrix = distance_matrix,
   distance.thresholds = 0,</pre>
```

## print\_importance

```
n.cores = 1,
verbose = FALSE
)
#evaluation with spatial cross-validation
rf.model <- rf_evaluate(
  model = rf.model,
  xy = plant_richness_df[, c("x", "y")],
  n.cores = 1
)
#checking evaluation results
print_evaluation(rf.model)
}
```

print\_importance Prints variable importance

# Description

Prints variable importance scores from rf, rf\_repeat, and rf\_spatial models.

#### Usage

```
print_importance(
   model,
   verbose = TRUE
)
```

## Arguments

model	A model fitted with rf, rf_repeat, or rf_spatial.
verbose	Logical, if TRUE, variable importance is returned. Default: TRUE

# Value

A table printed to the standard output.

# See Also

plot\_importance(), get\_importance()

## Examples

```
if(interactive()){
```

```
#loading example data
data(plant_richness_df)
data(distance.matrix)
#fitting a random forest model
rf.model <- rf(</pre>
  data = plant_richness_df,
  dependent.variable.name = "richness_species_vascular",
  predictor.variable.names = colnames(plant_richness_df)[5:21],
  distance.matrix = distance_matrix,
  distance.thresholds = 0,
  n.cores = 1,
  verbose = FALSE
)
#printing variable importance scores
print_importance(model = rf.model)
}
```

print\_moran

#### Prints results of a Moran's I test

## Description

Prints the results of a Moran's I test on the residuals of a model.

## Usage

```
print_moran(
   model,
   caption = NULL,
   verbose = TRUE
)
```

# Arguments

model	A model fitted with rf(), rf_repeat(), or rf_spatial().
caption	Character, caption of the output table, Default: NULL
verbose	Logical, if TRUE, the resulting table is printed into the console, Default: $\ensuremath{TRUE}$

## Value

Prints a table in the console using the huxtable package.

#### print\_performance

#### See Also

```
moran(), moran_multithreshold(), get_moran(), plot_moran()
```

#### Examples

```
if(interactive()){
 #loading example data
data(plant_richness_df)
 data(distance.matrix)
 #fitting random forest model
rf.model <- rf(</pre>
  data = plant_richness_df,
  dependent.variable.name = "richness_species_vascular",
  predictor.variable.names = colnames(plant_richness_df)[5:21],
  distance.matrix = distance_matrix,
  distance.thresholds = c(0, 1000, 2000),
  n.cores = 1,
  verbose = FALSE
)
 #printing Moran's I of model's residuals
print_moran(rf.model)
}
```

print\_performance print\_performance

## Description

Prints the performance slot of a model fitted with rf(), rf\_repeat(), or rf\_spatial(). For models fitted with rf\_repeat() it shows the median and the median absolute deviation of each performance measure.

#### Usage

```
print_performance(model)
```

## Arguments

model Model fitted with rf(), rf\_repeat(), or rf\_spatial().

#### Value

Prints model performance scores to the console.

#### See Also

print\_performance(), get\_performance()

#### Examples

```
if(interactive()){
#loading example data
data(plant_richness_df)
data(distance.matrix)
#fitting a random forest model
rf.model <- rf(</pre>
  data = plant_richness_df,
  dependent.variable.name = "richness_species_vascular",
  predictor.variable.names = colnames(plant_richness_df)[5:21],
  distance.matrix = distance_matrix,
  distance.thresholds = 0,
  n.cores = 1,
  verbose = FALSE
)
#printing performance scores
print_performance(rf.model)
}
```

rank\_spatial\_predictors

Ranks spatial predictors

#### Description

Ranks spatial predictors generated by mem\_multithreshold() or pca\_multithreshold() by their effect in reducing the Moran's I of the model residuals (ranking.method = "effect"), or by their own Moran's I (ranking.method = "moran").

In the former case, one model of the type  $y \sim \text{predictors} + \text{spatial_predictor_X}$  is fitted per spatial predictor, and the Moran's I of this model's residuals is compared with the one of the model without spatial predictors ( $y \sim \text{predictors}$ ), to finally rank the spatial predictor from maximum to minimum difference in Moran's I.

In the latter case, the spatial predictors are ordered by their Moran's I alone (this is the faster option).

In both cases, spatial predictors that are redundant with others at a Pearson correlation > 0.5 and spatial predictors with no effect (no reduction of Moran's I or Moran's I of the spatial predictor equal or lower than 0) are removed.

This function has been designed to be used internally by rf\_spatial() rather than directly by a user.

rank\_spatial\_predictors

# Usage

```
rank_spatial_predictors(
    data = NULL,
    dependent.variable.name = NULL,
    predictor.variable.names = NULL,
    distance.matrix = NULL,
    distance.thresholds = NULL,
    ranger.arguments = NULL,
    spatial.predictors.df = NULL,
    ranking.method = c("moran", "effect"),
    reference.moran.i = 1,
    verbose = FALSE,
    n.cores = parallel::detectCores() - 1,
    cluster = NULL
)
```

# Arguments

data	Data frame with a response variable and a set of predictors. Default: NULL
dependent.varia	able.name
	Character string with the name of the response variable. Must be in the column names of data. Default: NULL
predictor.varia	able.names
	Character vector with the names of the predictive variables. Every element of this vector must be in the column names of data. Default: NULL
distance.matrix	K
	Squared matrix with the distances among the records in data. The number of rows of distance.matrix and data must be the same. If not provided, the computation of the Moran's I of the residuals is omitted. Default: NULL
distance.thresh	nolds
	Numeric vector with neighborhood distances. All distances in the distance ma- trix below each value in dustance.thresholds are set to 0 for the computation of Moran's I. If NULL, it defaults to seq(0, max(distance.matrix), length.out = 4). Default: NULL
ranger.argument	ts
	List with ranger arguments. See rf or rf_repeat for further details.
spatial.predict	tors.df
	Data frame of spatial predictors.
ranking.method	Character, method used by to rank spatial predictors. The method "effect" ranks spatial predictors according how much each predictor reduces Moran's I of the model residuals, while the method "moran" ranks them by their own Moran's I. Default: "moran".
reference.moran.i	
	Moran's I of the residuals of the model without spatial predictors. Default: 1
verbose	Logical, ff TRUE, messages and plots generated during the execution of the function are displayed, Default: TRUE

n.cores	Integer, number of cores to use for parallel execution. Creates a socket cluster with parallel::makeCluster(), runs operations in parallel with foreach and %dopar%, and stops the cluster with parallel::clusterStop() when the job is done. Default: parallel::detectCores() - 1
cluster	A cluster definition generated with parallel::makeCluster(). If provided, overrides n.cores. When cluster = NULL (default value), and model is pro- vided, the cluster in model, if any, is used instead. If this cluster is NULL, then the function uses n.cores instead. The function does not stop a provided clus- ter, so it should be stopped with parallel::stopCluster() afterwards. The cluster definition is stored in the output list under the name "cluster" so it can be passed to other functions via the model argument, or using the %>% pipe. Default: NULL

#### Value

A list with four slots:

- method: Character, name of the method used to rank the spatial predictors.
- criteria: Data frame with two different configurations depending on the ranking method. If ranking.method = "effect", the columns contain the names of the spatial predictors, the rsquared of the model, the Moran's I of the model residuals, the difference between the Moran's I of the model including the given spatial predictor, and the Moran's I of the model fitted without spatial predictors, and the interpretation of the Moran's I value. If ranking.method = "moran", only the name of the spatial predictor and it's Moran's I are in the output data frame.
- ranking: Ordered character vector with the names of the spatial predictors selected.
- spatial.predictors.df: data frame with the selected spatial predictors in the order of the ranking.

```
if(interactive()){
#loading distance matrix
data(distance_matrix)
#computing Moran's Eigenvector Maps
mem.df <- mem(</pre>
 distance.matrix = distance_matrix[1:50, 1:50],
 distance.threshold = 0
)
#ranking by the Moran's I of the spatial predictor
rank <- rank_spatial_predictors(</pre>
 distance.matrix = distance_matrix[1:50, 1:50],
 distance.thresholds = 0,
 spatial.predictors.df = mem.df,
 ranking.method = "moran",
 n.cores = 1
)
```

## rescale\_vector

```
#checking Moran's I of MEMs
rank$criteria
#checking rank of MEMs
rank$ranking
}
```

rescale\_vector Rescales a numeric vector into a new range

## Description

Rescales a numeric vector to a new range.

#### Usage

```
rescale_vector(
  x = NULL,
  new.min = 0,
  new.max = 1,
  integer = FALSE
)
```

# Arguments

Х	Numeric vector. Default: NULL
new.min	New minimum value. Default: 0
new.max	New maximum value. Default: 1
integer	Logical, if TRUE, coerces the output to integer. Default: FALSE

## Value

A numeric vector of the same length as x, but with its values rescaled between new.min and new.max.

```
if(interactive()){
  out <- rescale_vector(
    x = rnorm(100),
    new.min = 0,
    new.max = 100,
    integer = TRUE
    )
    out</pre>
```

residuals\_diagnostics Normality test of a numeric vector

#### Description

Applies a Shapiro-Wilks test to a numeric vector, and plots the qq plot and the histogram.

## Usage

residuals\_diagnostics(residuals, predictions)

#### Arguments

residuals	Numeric vector, model residuals.
predictions	Numeric vector, model predictions.

#### Details

The function shapiro.test() has a hard limit of 5000 cases. If the model residuals have more than 5000 cases, then sample(x = residuals, size = 5000) is applied to the model residuals before the test.

## Value

A list with four slots:

/item w W statistic returned by shapiro.test(). /item p.value p-value of the Shapiro test. /item interpretation Character vector, one of "x is normal", "x is not normal". /item plot A patchwork plot with the qq plot and the histogram of x.

# See Also

ggplot,aes,geom\_qq\_line,ggtheme,labs,geom\_freqpoly,geom\_abline plot\_annotation

```
if(interactive()){
  residuals_diagnostics(
    residuals = runif(100),
    predictions = runif(100)
)
}
```

residuals\_test

#### Description

Applies a Shapiro-Wilks test to a numeric vector, and returns a list with the statistic W, its p-value, and a character string with the interpretation.

#### Usage

```
residuals_test(residuals)
```

#### Arguments

residuals Numeric vector, model residuals.

#### Value

A list with four slots:

/item w W statistic returned by shapiro.test(). /item p.value p-value of the Shapiro test. /item interpretation Character vector, one of "x is normal", "x is not normal". /item plot A patchwork plot with the qq plot and the histogram of x.

#### See Also

ggplot,aes,geom\_qq\_line,ggtheme,labs,geom\_freqpoly,geom\_abline plot\_annotation

#### Examples

```
if(interactive()){
  residuals_test(residuals = runif(100))
}
```

rf

Random forest models with Moran's I test of the residuals

## Description

A convenient wrapper for ranger that completes its output by providing the Moran's I of the residuals for different distance thresholds, the rmse and nrmse (as computed by root\_mean\_squared\_error()), and variable importance scores based on a scaled version of the data generated by scale.

## Usage

```
rf(
   data = NULL,
   dependent.variable.name = NULL,
   predictor.variable.names = NULL,
   distance.matrix = NULL,
   distance.thresholds = NULL,
   xy = NULL,
   ranger.arguments = NULL,
   scaled.importance = FALSE,
   seed = 1,
   verbose = TRUE,
   n.cores = parallel::detectCores() - 1,
   cluster = NULL
)
```

# Arguments data

Data frame with a response variable and a set of predictors. Default: NULL

dependent.variable.name

Character string with the name of the response variable. Must be in the column names of data. If the dependent variable is binary with values 1 and 0, the argument case.weights of ranger is populated by the function case\_weights(). Default: NULL

#### predictor.variable.names

Character vector with the names of the predictive variables. Every element of this vector must be in the column names of data. Optionally, the result of auto\_cor() or auto\_vif(). Default: NULL

#### distance.matrix

Squared matrix with the distances among the records in data. The number of rows of distance.matrix and data must be the same. If not provided, the computation of the Moran's I of the residuals is omitted. Default: NULL

#### distance.thresholds

Numeric vector with neighborhood distances. All distances in the distance matrix below each value in dustance.thresholds are set to 0 for the computation of Moran's I. If NULL, it defaults to seq(0, max(distance.matrix), length.out = 4). Default: NULL

xy (optional) Data frame or matrix with two columns containing coordinates and named "x" and "y". It is not used by this function, but it is stored in the slot ranger.arguments\$xy of the model, so it can be used by rf\_evaluate() and rf\_tuning(). Default: NULL

ranger.arguments

Named list with ranger arguments (other arguments of this function can also go here). All ranger arguments are set to their default values except for 'importance', that is set to 'permutation' rather than 'none'. The ranger arguments x, y, and formula are disabled. Please, consult the help file of ranger if you are not familiar with the arguments of this function.

#### scaled.importance

	Logical, if TRUE, the function scales data with scale and fits a new model to compute scaled variable importance scores. This makes variable importance scores of different models somewhat comparable. Default: FALSE
seed	Integer, random seed to facilitate reproducibility. If set to a given number, the returned model is always the same. Default: 1
verbose	Boolean. If TRUE, messages and plots generated during the execution of the function are displayed. Default: TRUE
n.cores	Integer, number of cores to use. Default: parallel::detectCores() - 1
cluster	A cluster definition generated with parallel::makeCluster(). This function does not use the cluster, but can pass it on to other functions when using the %>% pipe. It will be stored in the slot cluster of the output list. Default: NULL

# Details

Please read the help file of ranger for further details. Notice that the formula interface of ranger is supported through ranger.arguments, but variable interactions are not allowed (but check the\_feature\_engineer()).

#### Value

A ranger model with several extra slots:

- ranger.arguments: Stores the values of the arguments used to fit the ranger model.
- importance: A list containing a data frame with the predictors ordered by their importance, a ggplot showing the importance values, and local importance scores (difference in accuracy between permuted and non permuted variables for every case, computed on the out-of-bag data).
- performance: performance scores: R squared on out-of-bag data, R squared (cor(observed, predicted) ^ 2), pseudo R squared (cor(observed, predicted)), RMSE, and normalized RMSE (NRMSE).
- residuals: residuals, normality test of the residuals computed with residuals\_test(), and spatial autocorrelation of the residuals computed with moran\_multithreshold().

```
if(interactive()){
    #loading example data
    data("plant_richness_df")
    data("distance_matrix")
    #fittind random forest model
    out <- rf(
        data = plant_richness_df,
        dependent.variable.name = "richness_species_vascular",
        predictor.variable.names = colnames(plant_richness_df)[5:21],
        distance.matrix = distance_matrix,
        distance.thresholds = 0,</pre>
```

## rf\_compare

```
n.cores = 1
)
 class(out)
 #data frame with ordered variable importance
 out$importance$per.variable
 #variable importance plot
 out$importance$per.variable.plot
 #performance
 out$performance
 #spatial correlation of the residuals
 out$spatial.correlation.residuals$per.distance
 #plot of the Moran's I of the residuals for different distance thresholds
 out$spatial.correlation.residuals$plot
 #predictions for new data as done with ranger models:
predicted <- stats::predict(</pre>
  object = out,
  data = plant_richness_df,
  type = "response"
 )$predictions
 #alternative data input methods
 #ranger.arguments can contain ranger arguments and any other rf argument
 my.ranger.arguments <- list(</pre>
 data = plant_richness_df,
 dependent.variable.name = "richness_species_vascular",
predictor.variable.names = colnames(plant_richness_df)[8:21],
distance.matrix = distance_matrix,
distance.thresholds = c(0, 1000)
)
 #fitting model with these ranger arguments
 out <- rf(</pre>
  ranger.arguments = my.ranger.arguments,
  n.cores = 1
  )
}
```

rf\_compare

## rf\_compare

## Description

Uses rf\_evaluate() to compare the performance of several models on independent spatial folds via spatial cross-validation.

# Usage

```
rf_compare(
 models = NULL,
 xy = NULL,
 repetitions = 30,
  training.fraction = 0.75,
 metrics = c("r.squared", "pseudo.r.squared", "rmse", "nrmse", "auc"),
 distance.step = NULL,
 distance.step.x = NULL,
 distance.step.y = NULL,
 fill.color = viridis::viridis(100, option = "F", direction = -1, alpha = 0.8),
 line.color = "gray30",
 seed = 1,
 verbose = TRUE,
 n.cores = parallel::detectCores() - 1,
 cluster = NULL
)
```

# Arguments

models	Named list with models resulting from rf(), rf_spatial(), rf_tuning(), or rf_evaluate(). Example: models = list(a = model.a, b = model.b). Default: NULL
ху	Data frame or matrix with two columns containing coordinates and named "x" and "y". Default: NULL
repetitions	Integer, number of spatial folds to use during cross-validation. Must be lower than the total number of rows available in the model's data. Default: 30
training.fract	ion
	Proportion between 0.5 and 0.9 indicating the proportion of records to be used as training set during spatial cross-validation. Default: 0.75
metrics	Character vector, names of the performance metrics selected. The possible values are: "r.squared" (cor(obs, pred) ^ 2), "pseudo.r.squared" (cor(obs, pred)), "rmse" (sqrt(sum((obs - pred)^2)/length(obs))), "nrmse" (rmse/(quantile(obs, 0.75) - quantile(obs, 0.25))). Default: c("r.squared", "pseudo.r.squared", "rmse", "nrmse")
distance.step	Numeric, argument distance.step of thinning_til_n(). distance step used during the selection of the centers of the training folds. These fold centers are selected by thinning the data until a number of folds equal or lower than repetitions is reached. Its default value is 1/1000th the maximum distance within records in xy. Reduce it if the number of training folds is lower than expected.

distance.step.x	
	Numeric, argument distance.step.x of make_spatial_folds(). Distance step used during the growth in the x axis of the buffers defining the training folds. Default: NULL (1/1000th the range of the x coordinates).
distance.step.y	
	Numeric, argument distance.step.x of make_spatial_folds(). Distance step used during the growth in the y axis of the buffers defining the training folds. Default: NULL (1/1000th the range of the y coordinates).
fill.color	Character vector with hexadecimal codes (e.g. "#440154FF" "#21908CFF" "#FDE725FF"), or function generating a palette (e.g. viridis::viridis(100)). Default: viridis::viridis(100, option = "F", direction = -1)
line.color	Character string, color of the line produced by ggplot2::geom_smooth(). De-fault: "gray30"
seed	Integer, random seed to facilitate reproduciblity. If set to a given number, the results of the function are always the same. Default: 1.
verbose	Logical. If TRUE, messages and plots generated during the execution of the func- tion are displayed, Default: TRUE
n.cores	Integer, number of cores to use for parallel execution. Creates a socket cluster with parallel::makeCluster(), runs operations in parallel with foreach and %dopar%, and stops the cluster with parallel::clusterStop() when the job is done. Default: parallel::detectCores() - 1
cluster	A cluster definition generated with parallel::makeCluster(). If provided, overrides n.cores. When cluster = NULL (default value), and model is pro- vided, the cluster in model, if any, is used instead. If this cluster is NULL, then the function uses n.cores instead. The function does not stop a provided clus- ter, so it should be stopped with parallel::stopCluster() afterwards. The cluster definition is stored in the output list under the name "cluster" so it can be passed to other functions via the model argument, or using the %>% pipe. Default: NULL

# Value

A list with three slots:

- comparison.df: Data frame with one performance value per spatial fold, metric, and model.
- spatial.folds: List with the indices of the training and testing records for each evaluation repetition.
- plot: Violin-plot of comparison.df.

## See Also

rf\_evaluate()

# Examples

if(interactive()){

## rf\_evaluate

```
#loading example data
 data(distance_matrix)
 data(plant_richness_df)
 #fitting random forest model
 rf.model <- rf(</pre>
  data = plant_richness_df,
  dependent.variable.name = "richness_species_vascular",
  predictor.variable.names = colnames(plant_richness_df)[5:21],
  distance.matrix = distance_matrix,
  distance.thresholds = 0,
  n.cores = 1
)
 #fitting a spatial model with Moran's Eigenvector Maps
 rf.spatial <- rf_spatial(</pre>
model = rf.model,
n.cores = 1
)
 #comparing the spatial and non spatial models
comparison <- rf_compare(</pre>
models = list(
   `Non spatial` = rf.model,
  Spatial = rf.spatial
),
xy = plant_richness_df[, c("x", "y")],
metrics = c("r.squared", "rmse"),
n.cores = 1
)
}
```

rf\_evaluate Evaluates random forest models with spatial cross-validation

#### Description

Evaluates the performance of random forest on unseen data over independent spatial folds.

## Usage

```
rf_evaluate(
  model = NULL,
  xy = NULL,
  repetitions = 30,
  training.fraction = 0.75,
  metrics = c("r.squared", "pseudo.r.squared", "rmse", "nrmse", "auc"),
  distance.step = NULL,
  distance.step.x = NULL,
```

```
distance.step.y = NULL,
grow.testing.folds = FALSE,
seed = 1,
verbose = TRUE,
n.cores = parallel::detectCores() - 1,
cluster = NULL
)
```

# Arguments

model	Model fitted with rf(), rf_repeat(), or rf_spatial().
ху	Data frame or matrix with two columns containing coordinates and named "x" and "y". If NULL, the function will throw an error. Default: NULL
repetitions	Integer, number of spatial folds to use during cross-validation. Must be lower than the total number of rows available in the model's data. Default: 30
training.fract:	ion
	Proportion between 0.5 and 0.9 indicating the proportion of records to be used as training set during spatial cross-validation. Default: 0.75
metrics	Character vector, names of the performance metrics selected. The possible values are: "r.squared" (cor(obs, pred) ^ 2), "pseudo.r.squared" (cor(obs, pred)), "rmse" (sqrt(sum((obs - pred)^2)/length(obs))), "nrmse" (rmse/(quantile(obs, 0.75) - quantile(obs, 0.25))), and "auc" (only for binary responses with values 1 and 0). Default: c("r.squared", "pseudo.r.squared", "rmse", "nrmse")
distance.step	Numeric, argument distance.step of thinning_til_n(). distance step used during the selection of the centers of the training folds. These fold centers are selected by thinning the data until a number of folds equal or lower than repetitions is reached. Its default value is 1/1000th the maximum distance within records in xy. Reduce it if the number of training folds is lower than expected.
distance.step.:	-
	Numeric, argument distance.step.x of make_spatial_folds(). Distance step used during the growth in the x axis of the buffers defining the training folds. Default: NULL (1/1000th the range of the x coordinates).
distance.step.	y .
	Numeric, argument distance.step.x of make_spatial_folds(). Distance step used during the growth in the y axis of the buffers defining the training folds. Default: NULL (1/1000th the range of the y coordinates).
grow.testing.folds	
	Logic. By default, this function grows contiguous training folds to keep the spa- tial structure of the data as intact as possible. However, when setting grow.testing.folds = TRUE, the argument training.fraction is set to 1 - training.fraction, and the training and testing folds are switched. This option might be useful when the training data has a spatial structure that does not match well with the default behavior of the function. Default: FALSE
seed	Integer, random seed to facilitate reproduciblity. If set to a given number, the results of the function are always the same. Default: 1.

verbose	Logical. If TRUE, messages and plots generated during the execution of the function are displayed, Default: $\ensuremath{TRUE}$
n.cores	Integer, number of cores to use for parallel execution. Creates a socket cluster with parallel::makeCluster(), runs operations in parallel with foreach and %dopar%, and stops the cluster with parallel::clusterStop() when the job is done. Default: parallel::detectCores() - 1
cluster	A cluster definition generated with parallel::makeCluster(). If provided, overrides n.cores. When cluster = NULL (default value), and model is pro- vided, the cluster in model, if any, is used instead. If this cluster is NULL, then the function uses n.cores instead. The function does not stop a provided clus- ter, so it should be stopped with parallel::stopCluster() afterwards. The cluster definition is stored in the output list under the name "cluster" so it can be passed to other functions via the model argument, or using the %>% pipe. Default: NULL

## Details

The evaluation algorithm works as follows: the number of repetitions and the input dataset (stored in model\$ranger.arguments\$data) are used as inputs for the function thinning\_til\_n(), that applies thinning() to the input data until as many cases as repetitions are left, and as separated as possible. Each of these remaining records will be used as a "fold center". From that point, the fold grows, until a number of points equal (or close) to training.fraction is reached. The indices of the records within the grown spatial fold are stored as "training" in the output list, and the remaining ones as "testing". Then, for each spatial fold, a "training model" is fitted using the cases corresponding with the training indices, and predicted over the cases corresponding with the testing indices. The model predictions on the "unseen" data are compared with the observations, and the performance measures (R squared, pseudo R squared, RMSE and NRMSE) computed.

## Value

A model of the class "rf\_evaluate" with a new slot named "evaluation", that is a list with the following slots:

- training.fraction: Value of the argument training.fraction.
- spatial.folds: Result of applying make\_spatial\_folds() on the data coordinates. It is a list with as many slots as repetitions are indicated by the user. Each slot has two slots named "training" and "testing", each one having the indices of the cases used on the training and testing models.
- per.fold: Data frame with the evaluation results per spatial fold (or repetition). It contains the ID of each fold, it's central coordinates, the number of training and testing cases, and the training and testing performance measures: R squared, pseudo R squared (cor(observed, predicted)), rmse, and normalized rmse.
- per.model: Same data as above, but organized per fold and model ("Training", "Testing", and "Full").
- aggregated: Same data, but aggregated by model and performance measure.

## Examples

```
if(interactive()){
#loading example data
data(plant_richness_df)
data(distance_matrix)
#fitting random forest model
rf.model <- rf(
  data = plant_richness_df,
  dependent.variable.name = "richness_species_vascular",
  predictor.variable.names = colnames(plant_richness_df)[5:21],
  distance.matrix = distance_matrix,
  distance.thresholds = 0,
  n.cores = 1,
  verbose = FALSE
)
#evaluation with spatial cross-validation
rf.model <- rf_evaluate(</pre>
  model = rf.model,
  xy = plant_richness_df[, c("x", "y")],
  n.cores = 1
)
#checking evaluation results
plot_evaluation(rf.model)
print_evaluation(rf.model)
x <- get_evaluation(rf.model)</pre>
}
```

rf\_importance Contribution of each predictor to model transferability

#### Description

Evaluates the contribution of the predictors to model transferability via spatial cross-validation. The function returns the median increase or decrease in a given evaluation metric (R2, pseudo R2, RMSE, nRMSE, or AUC) when a variable is introduced in a model, by comparing and evaluating via spatial cross-validation models with and without the given variable. This function was devised to provide importance scores that would be less sensitive to spatial autocorrelation than those computed internally by random forest on the out-of-bag data. This function is experimental.

#### Usage

```
rf_importance(
    model = NULL,
    xy = NULL,
```

# rf\_importance

```
repetitions = 30,
training.fraction = 0.75,
metric = c("r.squared", "pseudo.r.squared", "rmse", "nrmse", "auc"),
distance.step = NULL,
distance.step.x = NULL,
fill.color = viridis::viridis(100, option = "F", direction = -1, alpha = 1, end = 0.9),
line.color = "white",
seed = 1,
verbose = TRUE,
n.cores = parallel::detectCores() - 1,
cluster = NULL
)
```

# Arguments

model	Model fitted with rf() and/or rf_spatial(). The function doesn't work with models fitted with rf_repeat(). Default: NULL	
ху	Data frame or matrix with two columns containing coordinates and named "x" and "y". If NULL, the function will throw an error. Default: NULL	
repetitions	Integer, number of spatial folds to use during cross-validation. Must be lower than the total number of rows available in the model's data. Default: 30	
training.fracti	on	
	Proportion between 0.5 and 0.9 indicating the proportion of records to be used as training set during spatial cross-validation. Default: 0.75	
metric	Character, nams of the performance metric to use. The possible values are: "r.squared" (cor(obs, pred) ^ 2), "pseudo.r.squared" (cor(obs, pred)), "rmse" (sqrt(sum((obs - pred)^2)/length(obs))), "nrmse" (rmse/(quantile(obs, 0.75) - quantile(obs, 0.25))), and "auc" (only for binary responses with values 1 and 0). Default: "r.squared"	
distance.step	Numeric, argument distance.step of thinning_til_n(). distance step used during the selection of the centers of the training folds. These fold centers are selected by thinning the data until a number of folds equal or lower than repetitions is reached. Its default value is 1/1000th the maximum distance within records in xy. Reduce it if the number of training folds is lower than expected.	
distance.step.x		
	Numeric, argument distance.step.x of make_spatial_folds(). Distance step used during the growth in the x axis of the buffers defining the training folds. Default: NULL (1/1000th the range of the x coordinates).	
distance.step.y		
	Numeric, argument distance.step.x of make_spatial_folds(). Distance step used during the growth in the y axis of the buffers defining the training folds. Default: NULL (1/1000th the range of the y coordinates).	
fill.color	Character vector with hexadecimal codes (e.g. "#440154FF" "#21908CFF" "#FDE725FF"), or function generating a palette (e.g. viridis::viridis(100)). Default: viridis::viridis(100, option = "F", direction = -1, alpha = 0.8, end = 0.9)	

line.color	Character string, color of the line produced by ggplot2::geom_smooth(). Default: "white"
seed	Integer, random seed to facilitate reproduciblity. If set to a given number, the results of the function are always the same. Default: 1.
verbose	Logical. If TRUE, messages and plots generated during the execution of the function are displayed, Default: TRUE
n.cores	Integer, number of cores to use for parallel execution. Creates a socket cluster with parallel::makeCluster(), runs operations in parallel with foreach and %dopar%, and stops the cluster with parallel::clusterStop() when the job is done. Default: parallel::detectCores() - 1
cluster	A cluster definition generated with parallel::makeCluster(). If provided, overrides n.cores. When cluster = NULL (default value), and model is provided, the cluster in model, if any, is used instead. If this cluster is NULL, then the function uses n.cores instead. The function does not stop a provided cluster, so it should be stopped with parallel::stopCluster() afterwards. The cluster definition is stored in the output list under the name "cluster" so it can be passed to other functions via the model argument, or using the %>% pipe. Default: NULL

#### Value

The input model with new data in its "importance" slot. The new importance scores are included in the data frame model\$importance\$per.variable, under the column names "importance.cv" (median contribution to transferability over spatial cross-validation repetitions), "importance.cv.mad" (median absolute deviation of the performance scores over spatial cross-validation repetitions), "importance.cv.percent" ("importance.cv" expressed as a percent, taking the full model's performance as baseline), and "importance.cv.mad" (median absolute deviation of "importance.cv"). The plot is stored as "cv.per.variable.plot".

#### Examples

```
if(interactive()){
#loading example data
data(plant_richness_df)
data(distance_matrix)
xy <- plant_richness_df[, c("x", "y")]</pre>
#fitting random forest model
rf.model <- rf(</pre>
 data = plant_richness_df,
 dependent.variable.name = "richness_species_vascular",
 predictor.variable.names = colnames(plant_richness_df)[5:21],
 distance.matrix = distance_matrix,
 distance.thresholds = 0,
 xy = xy,
 n.cores = 1,
 verbose = FALSE
)
```

```
#computing predictor contribution to model transferability
rf.model <- rf_importance(rf.model)
}</pre>
```

rf\_repeat

Fits several random forest models on the same data

## Description

Fits several random forest models on the same data in order to capture the effect of the algorithm's stochasticity on the variable importance scores, predictions, residuals, and performance measures. The function relies on the median to aggregate performance and importance values across repetitions. It is recommended to use it after a model is fitted  $(rf() \text{ or } rf_spatial())$ , tuned  $(rf_tuning())$ , and/or evaluated  $(rf_evaluate())$ . This function is designed to be used after fitting a model with rf() or  $rf_spatial()$ , tuning it with  $rf_tuning()$  and evaluating it with  $rf_evaluate()$ .

#### Usage

```
rf_repeat(
 model = NULL,
 data = NULL,
  dependent.variable.name = NULL,
 predictor.variable.names = NULL,
  distance.matrix = NULL,
  distance.thresholds = NULL,
  xy = NULL,
  ranger.arguments = NULL,
  scaled.importance = FALSE,
  repetitions = 10,
  keep.models = TRUE,
  seed = 1,
  verbose = TRUE,
  n.cores = parallel::detectCores() - 1,
 cluster = NULL
)
```

#### Arguments

model	A model fitted with rf(). If provided, the data and ranger arguments are taken directly from the model definition (stored in model\$ranger.arguments). Default: NULL	
data	Data frame with a response variable and a set of predictors. Default: NULL	

Character string with the name of the response variable. Must be in the column names of data. If the dependent variable is binary with values 1 and 0, the argument case.weights of ranger is populated by the function case\_weights(). Default: NULL

predictor.variable.names

Character vector with the names of the predictive variables. Every element of this vector must be in the column names of data. Default: NULL

dis	tance	.matrix	

Squared matrix with the distances among the records in data. The number of rows of distance.matrix and data must be the same. If not provided, the computation of the Moran's I of the residuals is omitted. Default: NULL

#### distance.thresholds

Numeric vector with neighborhood distances. All distances in the distance matrix below each value in dustance.thresholds are set to 0 for the computation of Moran's I. If NULL, it defaults to seq(0, max(distance.matrix), length.out = 4). Default: NULL

xy (optional) Data frame or matrix with two columns containing coordinates and named "x" and "y". It is not used by this function, but it is stored in the slot ranger.arguments\$xy of the model, so it can be used by rf\_evaluate() and rf\_tuning(). Default: NULL

ranger.arguments

Named list with ranger arguments (other arguments of this function can also go here). All ranger arguments are set to their default values except for 'importance', that is set to 'permutation' rather than 'none'. Please, consult the help file of ranger if you are not familiar with the arguments of this function.

scaled.importance

Logical. If TRUE, and 'importance = "permutation', the function scales 'data' with scale and fits a new model to compute scaled variable importance scores. Default: FALSE

repetitions Integer, number of random forest models to fit. Default: 10

- keep.models Logical, if TRUE, the fitted models are returned in the models slot. Set to FALSE if the accumulation of models is creating issues with the RAM memory available. Default: TRUE.
- seed Integer, random seed to facilitate reproducibility. If set to a given number, the results of the function are always the same. Default: 1.
- verbose Logical, ff TRUE, messages and plots generated during the execution of the function are displayed, Default: TRUE
- n.cores Integer, number of cores to use for parallel execution. Creates a socket cluster with parallel::makeCluster(), runs operations in parallel with foreach and %dopar%, and stops the cluster with parallel::clusterStop() when the job is done. Default: parallel::detectCores() - 1
- cluster A cluster definition generated with parallel::makeCluster(). If provided, overrides n.cores. When cluster = NULL (default value), and model is provided, the cluster in model, if any, is used instead. If this cluster is NULL, then the function uses n.cores instead. The function does not stop a provided cluster, so it should be stopped with parallel::stopCluster() afterwards. The

cluster definition is stored in the output list under the name "cluster" so it can be passed to other functions via the model argument, or using the %>% pipe. Default: NULL

#### Value

A ranger model with several new slots:

- ranger.arguments: Stores the values of the arguments used to fit the ranger model.
- importance: A list containing a data frame with the predictors ordered by their importance, a ggplot showing the importance values, and local importance scores.
- performance: out-of-bag performance scores: R squared, pseudo R squared, RMSE, and normalized RMSE (NRMSE).
- pseudo.r.squared: computed as the correlation between the observations and the predictions.
- residuals: residuals, normality test of the residuals computed with residuals\_test(), and spatial autocorrelation of the residuals computed with moran\_multithreshold().

## Examples

```
if(interactive()){
```

```
#loading example data
data(plant_richness_df)
data(distance_matrix)
#fitting 5 random forest models
out <- rf_repeat(</pre>
 data = plant_richness_df,
 dependent.variable.name = "richness_species_vascular",
 predictor.variable.names = colnames(plant_richness_df)[5:21],
 distance.matrix = distance_matrix,
 distance.thresholds = 0,
 repetitions = 5,
 n.cores = 1
)
#data frame with ordered variable importance
out$importance$per.variable
#per repetition
out$importance$per.repetition
#variable importance plot
out$importance$per.repetition.plot
#performance
```

out\$performance

#spatial correlation of the residuals for different distance thresholds

```
out$spatial.correlation.residuals$per.distance
 #plot of the Moran's I of the residuals for different distance thresholds
 out$spatial.correlation.residuals$plot
 #using a model as an input for rf_repeat()
 rf.model <- rf(</pre>
  data = plant_richness_df,
  dependent.variable.name = "richness_species_vascular",
  predictor.variable.names = colnames(plant_richness_df)[8:21],
  distance.matrix = distance_matrix,
  distance.thresholds = 0,
  n.cores = 1
  )
 #repeating the model 5 times
 rf.repeat <- rf_repeat(</pre>
  model = rf.model,
  n.cores = 1
  )
 rf.repeat$performance
 rf.repeat$importance$per.repetition.plot
}
```

## Description

rf\_spatial

Fits spatial random forest models using different methods to generate, rank, and select spatial predictors acting as proxies of spatial processes not considered by the non-spatial predictors. The end goal is providing the model with information about the spatial structure of the data to minimize the spatial correlation (Moran's I) of the model residuals and generate honest variable importance scores.

Fits spatial random forest models

## Usage

```
rf_spatial(
   model = NULL,
   data = NULL,
   dependent.variable.name = NULL,
   predictor.variable.names = NULL,
   distance.matrix = NULL,
   distance.thresholds = NULL,
   xy = NULL,
   ranger.arguments = NULL,
```

## rf\_spatial

```
scaled.importance = TRUE,
method = c("mem.moran.sequential", "mem.effect.sequential", "mem.effect.recursive",
    "hengl", "hengl.moran.sequential", "hengl.effect.sequential",
    "hengl.effect.recursive", "pca.moran.sequential", "pca.effect.sequential",
    "pca.effect.recursive"),
    max.spatial.predictors = NULL,
    weight.r.squared = NULL,
    weight.penalization.n.predictors = NULL,
    seed = 1,
    verbose = TRUE,
    n.cores = parallel::detectCores() - 1,
    cluster = NULL
)
```

## Arguments

model	A model fitted with rf(). If used, the arguments data, dependent.variable.name, predictor.variable.names, distance.matrix, distance.thresholds, ranger.arguments, and scaled.importance are taken directly from the model definition. Default: NULL
data	Data frame with a response variable and a set of predictors. Default: NULL
dependent.vari	able.name
	Character string with the name of the response variable. Must be in the column
	names of data. If the dependent variable is binary with values 1 and 0, the argu-
	ment case.weights of ranger is populated by the function case_weights().
nnadiatan yani	Default: NULL
predictor.vari	
	Character vector with the names of the predictive variables. Every element of this vector must be in the column names of data. Default: NULL
distance.matri	
4100411001114011	Squared matrix with the distances among the records in data. The number of
	rows of distance.matrix and data must be the same. If not provided, the
	computation of the Moran's I of the residuals is omitted. Default: NULL
distance.thres	holds
	Numeric vector with distances in the same units as distance.matrix Distances
	below each distance threshold are set to 0 on separated copies of the distance
	matrix to compute Moran's I at different neighborhood distances. If NULL, it
	defaults to seq(0, max(distance.matrix)/2, length.out = 4) (defined by
	<pre>default_distance_thresholds()). Default: NULL</pre>
ху	(optional) Data frame or matrix with two columns containing coordinates and
	named "x" and "y". It is not used by this function, but it is stored in the slot
	ranger.arguments\$xy of the model, so it can be used by rf_evaluate() and
ranger.argumen	rf_tuning(). Default: NULL
ranger . ar gumen	Named list with ranger arguments (other arguments of this function can also go
	here). All ranger arguments are set to their default values except for 'impor-
	tance', that is set to 'permutation' rather than 'none'. Please, consult the help
	file of ranger if you are not familiar with the arguments of this function.

scaled.importance		
	Logical. If TRUE, and 'importance = "permutation', the function scales 'data' with scale and fits a new model to compute scaled variable importance scores. Default: TRUE	
method	Character, method to build, rank, and select spatial predictors. One of:	
	• "hengl"	
	• "hengl.moran.sequential" (experimental)	
	• "hengl.effect.sequential" (experimental)	
	• "hengl.effect.recursive" (experimental)	
	• "pca.moran.sequential" (experimental)	
	• "pca.effect.sequential" (experimental)	
	• "pca.effect.recursive" (experimental)	
	• "mem.moran.sequential"	
	• "mem.effect.sequential"	
	• "mem.effect.recursive"	
max.spatial.pr		
	Integer, maximum number of spatial predictors to generate. Useful when mem- ory problems arise due to a large number of spatial predictors, Default: NULL	
weight.r.squar		
	Numeric between 0 and 1, weight of R-squared in the selection of spatial components. See Details, Default: NULL	
weight.penalization.n.predictors		
	Numeric between 0 and 1, weight of the penalization for adding an increasing number of spatial predictors during selection. Default: NULL	
seed	Integer, random seed to facilitate reproducibility. Default: 1.	
verbose	Logical. If TRUE, messages and plots generated during the execution of the function are displayed, Default: TRUE	
n.cores	Integer, number of cores to use for parallel execution. Creates a socket cluster with parallel::makeCluster(), runs operations in parallel with foreach and %dopar%, and stops the cluster with parallel::clusterStop() when the job is done. Default: parallel::detectCores() - 1	
cluster	A cluster definition generated with parallel::makeCluster(). If provided, overrides n.cores. When cluster = NULL (default value), and model is pro- vided, the cluster in model, if any, is used instead. If this cluster is NULL, then the function uses n.cores instead. The function does not stop a provided clus- ter, so it should be stopped with parallel::stopCluster() afterwards. The cluster definition is stored in the output list under the name "cluster" so it can be passed to other functions via the model argument, or using the %>% pipe. Default: NULL	

## Details

The function uses three different methods to generate spatial predictors ("hengl", "pca", and "mem"), two methods to rank them in order to define in what order they are introduced in the model ("effect" and "moran), and two methods to select the spatial predictors that minimize the spatial correlation

of the model residuals ("sequential" and "recursive"). All method names but "hengl" (that uses the complete distance matrix as predictors in the spatial model) are named by combining a method to generate the spatial predictors, a method to rank them, and a method to select them, separated by a point. Examples are "mem.moran.sequential" or "mem.effect.recursive". All combinations are not possible, since the ranking method "moran" cannot be used with the selection method "recursive" (because the logics behind them are very different, see below). Methods to generate spatial predictors:

- "hengl": named after the method RFsp presented in the paper "Random forest as a generic framework for predictive modeling of spatial and spatio-temporal variables", by Hengl et al. (2018), where the authors propose to use the distance matrix among records as predictors in spatial random forest models (RFsp method). In this function, all methods starting with "hengl" use either the complete distance matrix, or select columns of the distance matrix as spatial predictors.
- "mem": Generates Moran's Eigenvector Maps, that is, the eigenvectors of the double-centered weights of the distance matrix. The method is described in "Spatial modelling: a comprehensive framework for principal coordinate analysis of neighbour matrices (PCNM)", by Dray et al. (2006), and "Statistical methods for temporal and space-time analysis of community composition data", by Legendre and Gauthier (2014).
- "pca": Computes spatial predictors from the principal component analysis of a weighted distance matrix (see weights\_from\_distance\_matrix()). This is an experimental method, use with caution.

Methods to rank spatial predictors (see rank\_spatial\_predictors()):

- "moran": Computes the Moran's I of each spatial predictor, selects the ones with positive values, and ranks them from higher to lower Moran's I.
- "effect": If a given non-spatial random forest model is defined as y = p1 + ... + pn, being p1 + ... + pn the set of predictors, for every spatial predictor generated (spX) a spatial model y = p1 + ... + pn + spX is fitted, and the Moran's I of its residuals is computed. The spatial predictors are then ranked by how much they help to reduce spatial autocorrelation between the non-spatial and the spatial model.

Methods to select spatial predictors:

- "sequential" (see select\_spatial\_predictors\_sequential()): The spatial predictors are added one by one in the order they were ranked, and once all spatial predictors are introduced, the best first n predictors are selected. This method is similar to the one employed in the MEM methodology (Moran's Eigenvector Maps) described in the paper "Spatial modelling: a comprehensive framework for principal coordinate analysis of neighbour matrices (PCNM)", by Dray et al. (2006), and "Statistical methods for temporal and space-time analysis of community composition data", by Legendre and Gauthier (2014). This method generally introduces tens of predictors into the model, but usually offers good results.
- "recursive" (see select\_spatial\_predictors\_recursive()): This method tries to find the smallest combination of spatial predictors that reduce the spatial correlation of the model's residuals the most. The algorithm goes as follows: 1. The first ranked spatial predictor is introduced into the model; 2. the remaining predictors are ranked again using the "effect" method, using the model in 1. as reference. The first spatial predictor in the resulting ranking is then introduced into the model, and the steps 1. and 2. are repeated until spatial predictors

stop having an effect in reducing the Moran's I of the model residuals. This method takes longer to compute, but generates smaller sets of spatial predictors. This is an experimental method, use with caution.

Once ranking procedure is completed, an algorithm is used to select the minimal subset of spatial predictors that reduce the most the Moran's I of the residuals: for each new spatial predictor introduced in the model, the Moran's I of the residuals, it's p-value, a binary version of the p-value (0 if < 0.05 and 1 if >= 0.05), the R-squared of the model, and a penalization linear with the number of spatial predictors introduced (computed as (1 / total spatial predictors) \* introduced spatial predictors) are rescaled between 0 and 1. Then, the optimization criteria is computed as max(1 - Moran's I, p-value binary) + (weig The predictors from the first one to the one with the highest optimization criteria are then selected as the best ones in reducing the spatial correlation of the model residuals, and used along with data to fit the final spatial model.

#### Value

A ranger model with several new slots:

- ranger.arguments: Values of the arguments used to fit the ranger model.
- importance: A list containing the vector of variable importance as originally returned by ranger (scaled or not depending on the value of 'scaled.importance'), a data frame with the predictors ordered by their importance, and a ggplot showing the importance values.
- performance: With the out-of-bag R squared, pseudo R squared, RMSE and NRMSE of the model.
- residuals: residuals, normality test of the residuals computed with residuals\_test(), and spatial autocorrelation of the residuals computed with moran\_multithreshold().
- spatial: A list with four slots:
  - method: Character, method used to generate, rank, and select spatial predictors.
  - names: Character vector with the names of the selected spatial predictors. Not returned if the method is "hengl".
  - optimization: Criteria used to select the spatial predictors. Not returned if the method is "hengl".
  - plot: Plot of the criteria used to select the spatial predictors. Not returned if the method is "hengl".

#### Examples

```
if(interactive()){
```

```
#loading example data
data(distance_matrix)
data(plant_richness_df)
```

```
#names of the response and predictors
dependent.variable.name <- "richness_species_vascular"
predictor.variable.names <- colnames(plant_richness_df)[5:21]</pre>
```

#hengl
model <- rf\_spatial(</pre>

```
data = plant_richness_df,
  dependent.variable.name = dependent.variable.name,
  predictor.variable.names = predictor.variable.names,
  distance.matrix = distance_matrix,
  distance.thresholds = 0,
  method = "hengl",
  n.cores = 1
)
 #mem.moran.sequential
 model <- rf_spatial(</pre>
  data = plant_richness_df,
  dependent.variable.name = dependent.variable.name,
  predictor.variable.names = predictor.variable.names,
  distance.matrix = distance_matrix,
  distance.thresholds = 0,
  method = "mem.moran.sequential",
  n.cores = 1
)
 #fitting an rf_spatial model from an rf model
 rf.model <- rf(</pre>
  data = plant_richness_df,
  dependent.variable.name = "richness_species_vascular",
  predictor.variable.names = colnames(plant_richness_df)[5:21],
  distance.matrix = distance_matrix,
  distance.thresholds = 0,
  n.cores = 1,
  verbose = FALSE
 )
 rf.model$spatial.correlation.residuals$plot
 #spatial version of the rf model
 rf.spatial <- rf_spatial(model = rf.model)</pre>
 rf.spatial$spatial.correlation.residuals$plot
}
```

rf\_tuning

*Tuning of random forest hyperparameters via spatial cross-validation* 

## Description

Finds the optimal set of random forest hyperparameters num.trees, mtry, and min.node.size via grid search by maximizing the model's R squared, or AUC, if the response variable is binomial, via spatial cross-validation performed with rf\_evaluate().

# Usage

```
rf_tuning(
  model = NULL,
  num.trees = NULL,
  mtry = NULL,
  min.node.size = NULL,
  xy = NULL,
  repetitions = 30,
  training.fraction = 0.75,
  seed = 1,
  verbose = TRUE,
  n.cores = parallel::detectCores() - 1,
  cluster = NULL
)
```

# Arguments

model	A model fitted with rf(). If provided, the training data is taken directly from the model definition (stored in model\$ranger.arguments). Default: NULL
num.trees	Numeric integer vector with the number of trees to fit on each model repetition. Default: c(500, 1000, 2000).
mtry	Numeric integer vector, number of predictors to randomly select from the com- plete pool of predictors on each tree split. Default: floor(seq(1, length(predictor.variable.names length.out = 4))
<pre>min.node.size</pre>	Numeric integer, minimal number of cases in a terminal node. Default: c(5, 10, 20, 40)
ху	Data frame or matrix with two columns containing coordinates and named "x" and "y". If NULL, the function will throw an error. Default: NULL
repetitions	Integer, number of independent spatial folds to use during the cross-validation. Default: 30.
training.fract	ion
	Proportion between 0.2 and 0.9 indicating the number of records to be used in model training. Default: 0.75
seed	Integer, random seed to facilitate reproduciblity. If set to a given number, the results of the function are always the same. Default: 1.
verbose	Logical. If TRUE, messages and plots generated during the execution of the function are displayed, Default: TRUE
n.cores	Integer, number of cores to use for parallel execution. Creates a socket cluster with parallel::makeCluster(), runs operations in parallel with foreach and %dopar%, and stops the cluster with parallel::clusterStop() when the job is done. Default: parallel::detectCores() - 1
cluster	A cluster definition generated with parallel::makeCluster(). If provided, overrides n.cores. When cluster = NULL (default value), and model is pro- vided, the cluster in model, if any, is used instead. If this cluster is NULL, then the function uses n.cores instead. The function does not stop a provided clus- ter, so it should be stopped with parallel::stopCluster() afterwards. The

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cluster definition is stored in the output list under the name "cluster" so it can be passed to other functions via the model argument, or using the %>% pipe. Default: NULL

## Value

A model with a new slot named tuning, with a data frame with the results of the tuning analysis.

#### See Also

rf\_evaluate()

## Examples

```
if(interactive()){
```

```
#loading example data
data(plant_richness_df)
data(distance_matrix)
#fitting model to tune
out <- rf(
    data = plant_richness_df,
    dependent variable name = "richness"</pre>
```

```
dependent.variable.name = "richness_species_vascular",
 predictor.variable.names = colnames(plant_richness_df)[5:21],
 distance.matrix = distance_matrix,
 distance.thresholds = 0,
 n.cores = 1
)
#model tuning
tuning <- rf_tuning(</pre>
 model = out,
 num.trees = c(100, 500),
 mtry = c(2, 8),
 min.node.size = c(5, 10),
 xy = plant_richness_df[, c("x", "y")],
 n.cores = 1
)
}
```

root\_mean\_squared\_error

RMSE and normalized RMSE

#### Description

Computes the rmse or normalized rmse (nrmse) between two numeric vectors of the same length representing observations and model predictions.

#### Usage

```
root_mean_squared_error(
    o,
    p,
    normalization = c("rmse", "all", "mean", "sd", "maxmin", "iq")
)
```

#### Arguments

0	Numeric vector with observations, must have the same length as p.
р	Numeric vector with predictions, must have the same length as o.
normalization	character, normalization method, Default: "rmse" (see Details).

# Details

The normalization methods go as follows:

- "rmse": RMSE with no normalization.
- "mean": RMSE dividied by the mean of the observations (rmse/mean(o)).
- "sd": RMSE dividied by the standard deviation of the observations (rmse/sd(o)).
- "maxmin": RMSE divided by the range of the observations (rmse/(max(o) min(o))).
- "iq": RMSE divided by the interquartile range of the observations (rmse/(quantile(o, 0.75) quantile(o, 0.25)))

#### Value

Named numeric vector with either one or 5 values, as selected by the user.

## Examples

```
if(interactive()){
  root_mean_squared_error(
    o = runif(10),
    p = runif(10)
    )
}
```

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### Description

Selects spatial predictors following these steps:

- 1. Gets the spatial predictors ranked by rank\_spatial\_predictors() and fits a model of the form y ~ predictors + best\_spatial\_predictor\_1. The Moran's I of the residuals of this model is used as reference value for the next step.
- The remaining spatial predictors are introduced again into rank\_spatial\_predictors(), and the spatial predictor with the highest ranking is introduced in a new model of the form y ~predictors + best\_spatial\_predictor\_1 + best\_spatial\_predictor\_2.
- 3. Steps 1 and 2 are repeated until the Moran's I doesn't improve for a number of repetitions equal to the 20 percent of the total number of spatial predictors introduced in the function.

This method allows to select the smallest set of spatial predictors that have the largest joint effect in reducing the spatial correlation of the model residuals, while maintaining the model's R-squared as high as possible. As a consequence of running rank\_spatial\_predictors() on each iteration, this method includes in the final model less spatial predictors than the sequential method implemented in select\_spatial\_predictors\_sequential() would do, while minimizing spatial correlation and maximizing the R squared of the model as much as possible.

#### Usage

```
select_spatial_predictors_recursive(
    data = NULL,
    dependent.variable.name = NULL,
    predictor.variable.names = NULL,
    distance.matrix = NULL,
    distance.thresholds = NULL,
    ranger.arguments = NULL,
    spatial.predictors.df = NULL,
    spatial.predictors.ranking = NULL,
    weight.r.squared = 0.25,
    weight.penalization.n.predictors = 0,
    n.cores = parallel::detectCores() - 1,
    cluster = NULL
)
```

#### /

## Arguments

data Data frame with a response variable and a set of predictors. Default: NULL dependent.variable.name

Character string with the name of the response variable. Must be in the column names of data. Default: NULL

predictor.variable.names

Character vector with the names of the predictive variables. Every element of this vector must be in the column names of data. Default: NULL

#### distance.matrix

Squared matrix with the distances among the records in data. The number of rows of distance.matrix and data must be the same. If not provided, the computation of the Moran's I of the residuals is omitted. Default: NULL

#### distance.thresholds

Numeric vector with neighborhood distances. All distances in the distance matrix below each value in dustance. thresholds are set to 0 for the computation of Moran's I. If NULL, it defaults to seq(0, max(distance.matrix), length.out = 4). Default: NULL

#### ranger.arguments

Named list with ranger arguments (other arguments of this function can also go here). All ranger arguments are set to their default values except for 'importance', that is set to 'permutation' rather than 'none'. Please, consult the help file of ranger if you are not familiar with the arguments of this function.

```
spatial.predictors.df
```

Data frame of spatial predictors.

```
spatial.predictors.ranking
```

Ranking of predictors returned by rank\_spatial\_predictors().

weight.r.squared

Numeric between 0 and 1, weight of R-squared in the optimization index. Default: 0.25

weight.penalization.n.predictors

Numeric between 0 and 1, weight of the penalization for the number of spatial predictors added in the optimization index. Default: 0

- n.cores Integer, number of cores to use. Default: parallel::detectCores() 1
- cluster A cluster definition generated by parallel::makeCluster(). Default: NULL

#### Details

The algorithm works as follows. If the function rank\_spatial\_predictors() returns 10 ranked spatial predictors (sp1 to sp10, being sp7 the best one), select\_spatial\_predictors\_recursive() is going to first fit the model y ~ predictors + sp7. Then, the spatial predictors sp2 to sp9 are again ranked with rank\_spatial\_predictors() using the model y ~ predictors + sp7 as reference (at this stage, some of the spatial predictors might be dropped due to lack of effect). When the new ranking of spatial predictors is ready (let's say they are sp5, sp3, and sp4), the best one (sp5) is included in the model y ~ predictors + sp7 + sp5, and the remaining ones go again to rank\_spatial\_predictors() to repeat the process until spatial predictors are depleted.

## Value

A list with two slots: optimization, a data frame with the index of the spatial predictor added on each iteration, the spatial correlation of the model residuals, and the R-squared of the model, and best.spatial.predictors, that is a character vector with the names of the spatial predictors that minimize the Moran's I of the residuals and maximize the R-squared of the model.

#### Examples

```
if(interactive()){
#loading example data
data(distance_matrix)
data(plant_richness_df)
#response and preditor names
dependent.variable.name = "richness_species_vascular"
predictor.variable.names = colnames(plant_richness_df)[5:21]
#non-spatial model
model <- rf(
 data = plant_richness_df,
 dependent.variable.name = dependent.variable.name,
 predictor.variable.names = predictor.variable.names,
 distance.matrix = distance_matrix,
 distance.thresholds = 0,
 n.cores = 1
)
#preparing spatial predictors
spatial.predictors <- mem_multithreshold(</pre>
 distance.matrix = distance_matrix,
 distance.thresholds = 0
)
#ranking spatial predictors
spatial.predictors.ranking <- rank_spatial_predictors(</pre>
 data = plant_richness_df,
 dependent.variable.name = dependent.variable.name,
 predictor.variable.names = predictor.variable.names,
 spatial.predictors.df = spatial.predictors,
 ranking.method = "moran",
 reference.moran.i = model$spatial.correlation.residuals$max.moran,
 distance.matrix = distance_matrix,
 distance.thresholds = 0,
 n.cores = 1
)
#selecting the best subset of predictors
selection <- select_spatial_predictors_recursive(</pre>
 data = plant_richness_df,
 dependent.variable.name = dependent.variable.name,
 predictor.variable.names = predictor.variable.names,
 distance.matrix = distance_matrix,
 distance.thresholds = 0,
 spatial.predictors.df = spatial.predictors,
 spatial.predictors.ranking = spatial.predictors.ranking,
 n.cores = 1
)
```

```
selection$optimization
selection$best.spatial.predictors
plot_optimization(selection$optimization)
```

}

## Description

Selects spatial predictors by adding them sequentially into a model while monitoring the Moran's I of the model residuals and the model's R-squared. Once all the available spatial predictors have been added to the model, the function identifies the first n predictors that minimize the spatial correlation of the residuals and maximize R-squared, and returns the names of the selected spatial predictors and a data frame with the selection criteria.

## Usage

```
select_spatial_predictors_sequential(
    data = NULL,
    dependent.variable.name = NULL,
    predictor.variable.names = NULL,
    distance.matrix = NULL,
    distance.thresholds = NULL,
    ranger.arguments = NULL,
    spatial.predictors.df = NULL,
    spatial.predictors.ranking = NULL,
    weight.r.squared = 0.75,
    weight.penalization.n.predictors = 0.25,
    verbose = FALSE,
    n.cores = parallel::detectCores() - 1,
    cluster = NULL
)
```

#### Arguments

data Data frame with a response variable and a set of predictors. Default: NULL dependent.variable.name

Character string with the name of the response variable. Must be in the column names of data. Default: NULL

predictor.variable.names

Character vector with the names of the predictive variables. Every element of this vector must be in the column names of data. Default: NULL

distance.matrix	
	Squared matrix with the distances among the records in data. The number of rows of distance.matrix and data must be the same. If not provided, the computation of the Moran's I of the residuals is omitted. Default: NULL
distance.thresh	olds
	Numeric vector with neighborhood distances. All distances in the distance ma- trix below each value in dustance.thresholds are set to 0 for the computation of Moran's I. If NULL, it defaults to seq(0, max(distance.matrix), length.out = 4). Default: NULL
ranger.argument	S
	Named list with ranger arguments (other arguments of this function can also go here). All ranger arguments are set to their default values except for 'impor- tance', that is set to 'permutation' rather than 'none'. Please, consult the help file of ranger if you are not familiar with the arguments of this function.
spatial.predict	ors.df
	Data frame of spatial predictors.
spatial.predict	ors.ranking
	Ranking of the spatial predictors returned by rank_spatial_predictors().
weight.r.square	d
	Numeric between 0 and 1, weight of R-squared in the optimization index. Default: $0.75$
weight.penaliza	tion.n.predictors
	Numeric between 0 and 1, weight of the penalization for the number of spatial predictors added in the optimization index. Default: 0.25
verbose	Logical, ff TRUE, messages and plots generated during the execution of the function are displayed, Default: FALSE
n.cores	Integer, number of cores to use. Default: parallel::detectCores() - 1
cluster	A cluster definition generated by parallel::makeCluster(). Default: NULL

# Details

The algorithm works as follows: If the function rank\_spatial\_predictors returns 10 spatial predictors (sp1 to sp10, ordered from best to worst), select\_spatial\_predictors\_sequential is going to fit the models  $y \sim \text{predictors} + \text{sp1}$ ,  $y \sim \text{predictors} + \text{sp1} + \text{sp2}$ , until all spatial predictors are used in  $y \sim \text{predictors} + \text{sp1} \dots \text{sp10}$ . The model with lower Moran's I of the residuals and higher R-squared (computed on the out-of-bag data) is selected, and its spatial predictors returned.

## Value

A list with two slots: optimization, a data frame with the index of the spatial predictor added on each iteration, the spatial correlation of the model residuals, and the R-squared of the model, and best.spatial.predictors, that is a character vector with the names of the spatial predictors that minimize the Moran's I of the residuals and maximize the R-squared of the model.

## Examples

if(interactive()){

```
#loading example data
data(distance_matrix)
data(plant_richness_df)
#common arguments
dependent.variable.name = "richness_species_vascular"
predictor.variable.names = colnames(plant_richness_df)[5:21]
#non-spatial model
model <- rf(</pre>
 data = plant_richness_df,
 dependent.variable.name = dependent.variable.name,
 predictor.variable.names = predictor.variable.names,
 distance.matrix = distance_matrix,
 distance.thresholds = 0,
 n.cores = 1
)
#preparing spatial predictors
spatial.predictors <- mem_multithreshold(</pre>
 distance.matrix = distance.matrix,
 distance.thresholds = 0
)
#ranking spatial predictors by their Moran's I (faster option)
spatial.predictors.ranking <- rank_spatial_predictors(</pre>
 ranking.method = "moran",
 spatial.predictors.df = spatial.predictors,
 reference.moran.i = model$spatial.correlation.residuals$max.moran,
 distance.matrix = distance.matrix,
 distance.thresholds = 0,
 n.cores = 1
)
#selecting the best subset of predictors
selection <- select_spatial_predictors_sequential(</pre>
 data = plant_richness_df,
 dependent.variable.name = dependent.variable.name,
 predictor.variable.names = predictor.variable.names,
 distance.matrix = distance_matrix,
 distance.thresholds = 0,
 spatial.predictors.df = spatial.predictors,
 spatial.predictors.ranking = spatial.predictors.ranking,
 n.cores = 1
)
selection$optimization
selection$best.spatial.predictors
plot_optimization(selection$optimization)
}
```

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standard\_error

## Description

Computes the standard error of the mean of a numeric vector as round(sqrt(var(x)/length(x)), 3)

## Usage

standard\_error(x)

# Arguments ×

A numeric vector.

#### Details

The function removes NA values before computing the standard error, and rounds the result to 3 decimal places.

## Value

A numeric value.

## Examples

```
if(interactive()){
   standard_error(runif(10))
```

}

statistical\_mode Statistical mode of a vector

## Description

Computes the mode of a numeric or character vector

## Usage

```
statistical_mode(x)
```

## Arguments

x Numeric or character vector.

#### Value

Statistical mode of x.

#### Examples

```
if(interactive()){
   statistical_mode(c(10, 9, 10, 8))
}
```

the\_feature\_engineer Suggest variable interactions and composite features for random forest models

#### Description

Suggests candidate variable interactions and composite features able to improve predictive accuracy over data not used to train the model via spatial cross-validation with rf\_evaluate(). For a pair of predictors a and b, interactions are build via multiplication (a \* b), while composite features are built by extracting the first factor of a principal component analysis performed with pca(), after rescaling a and b between 1 and 100. Interactions and composite features are named a..x..b and a..pca..b respectively.

Candidate variables a and b are selected from those predictors in predictor.variable.names with a variable importance above importance.threshold (set by default to the median of the importance scores).

For each interaction and composite feature, a model including all the predictors plus the interaction or composite feature is fitted, and it's R squared (or AUC if the response is binary) computed via spatial cross-validation (see rf\_evaluate()) is compared with the R squared of the model without interactions or composite features.

From all the potential interactions screened, only those with a positive increase in R squared (or AUC when the response is binomial) of the model, a variable importance above the median, and a maximum correlation among themselves and with the predictors in predictor.variable.names not higher than cor.threshold (set to 0.5 by default) are selected. Such a restrictive set of rules ensures that the selected interactions can be used right away for modeling purposes without increasing model complexity unnecessarily. However, the suggested variable interactions might not make sense from a domain expertise standpoint, so please, examine them with care.

The function returns the criteria used to select the interactions, and the data required to use these interactions a model.

#### Usage

```
the_feature_engineer(
   data = NULL,
   dependent.variable.name = NULL,
   predictor.variable.names = NULL,
```

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## the\_feature\_engineer

```
xy = NULL,
ranger.arguments = NULL,
repetitions = 30,
training.fraction = 0.75,
importance.threshold = 0.75,
cor.threshold = 0.75,
point.color = viridis::viridis(100, option = "F", alpha = 0.8),
seed = NULL,
verbose = TRUE,
n.cores = parallel::detectCores() - 1,
cluster = NULL
)
```

# Arguments

data	Data frame with a response variable and a set of predictors. Default: NULL
dependent.varia	able.name
	Character string with the name of the response variable. Must be in the column names of data. If the dependent variable is binary with values 1 and 0, the argument case.weights of ranger is populated by the function case_weights(). Default: NULL
predictor.varia	
	Character vector with the names of the predictive variables, or object of class "variable_selection" produced by auto_vif() and/or auto_cor(). Every element of this vector must be in the column names of data. Default: NULL
ху	Data frame or matrix with two columns containing coordinates and named "x" and "y". If not provided, the comparison between models with and without variable interactions is not done.
ranger.argument	LS .
	Named list with ranger arguments (other arguments of this function can also go here). All ranger arguments are set to their default values except for 'impor- tance', that is set to 'permutation' rather than 'none'. Please, consult the help file of ranger if you are not familiar with the arguments of this function.
repetitions	Integer, number of spatial folds to use during cross-validation. Must be lower than the total number of rows available in the model's data. Default: 30
training.fracti	ion
	Proportion between 0.5 and 0.9 indicating the proportion of records to be used as training set during spatial cross-validation. Default: 0.75
importance.thre	eshold
	Numeric between 0 and 1, quantile of variable importance scores over which to select individual predictors to explore interactions among them. Larger values reduce the number of potential interactions explored. Default: 0.75
cor.threshold	Numeric, maximum Pearson correlation between any pair of the selected inter- actions, and between any interaction and the predictors in predictor.variable.names. Default: 0.75
point.color	Colors of the plotted points. Can be a single color name (e.g. "red4"), a character vector with hexadecimal codes (e.g. "#440154FF" "#21908CFF" "#FDE725FF"),

	or function generating a palette (e.g. viridis::viridis(100)). Default: viridis::viridis(100, option = "F", alpha = 0.8)
seed	Integer, random seed to facilitate reproduciblity. If set to a given number, the results of the function are always the same. Default: NULL
verbose	Logical. If TRUE, messages and plots generated during the execution of the func- tion are displayed. Default: TRUE
n.cores	Integer, number of cores to use for parallel execution. Creates a socket cluster with parallel::makeCluster(), runs operations in parallel with foreach and %dopar%, and stops the cluster with parallel::clusterStop() when the job is done. Default: parallel::detectCores() - 1
cluster	A cluster definition generated with parallel::makeCluster(). If provided, overrides n.cores. When cluster = NULL (default value), and model is pro- vided, the cluster in model, if any, is used instead. If this cluster is NULL, then the function uses n.cores instead. The function does not stop a provided clus- ter, so it should be stopped with parallel::stopCluster() afterwards. The cluster definition is stored in the output list under the name "cluster" so it can be passed to other functions via the model argument, or using the %>% pipe. Default: NULL

#### Value

A list with seven slots:

- screening: Data frame with selection scores of all the interactions considered.
- selected: Data frame with selection scores of the selected interactions.
- df: Data frame with the computed interactions.
- plot: List of plots of the selected interactions versus the response variable. The output list can be plotted all at once with patchwork::wrap\_plots(p) or cowplot::plot\_grid(plotlist = p), or one by one by extracting each plot from the list.
- data: Data frame with the response variable, the predictors, and the selected interactions, ready to be used as data argument in the package functions.
- dependent.variable.name: Character, name of the response.
- predictor.variable.names: Character vector with the names of the predictors and the selected interactions.

#### Examples

if(interactive()){

```
#load example data
data(plant_richness_df)
new.features <- the_feature_engineer(
   data = plant_richness_df,
   dependent.variable.name = "richness_species_vascular",
   predictor.variable.names = colnames(plant_richness_df)[5:21],
   n.cores = 1,</pre>
```

## thinning

```
verbose = TRUE
)
new.features$screening
new.features$selected
new.features$columns
```

}

thinning

#### Applies thinning to pairs of coordinates

## Description

Resamples a set of points with x and y coordinates to impose a minimum distance among nearby points.

# Usage

thinning(xy, minimum.distance = NULL)

#### Arguments

#### ху

A data frame with columns named "x" and "y" representing geographic coordinates.

minimum.distance

Numeric, minimum distance to be set between nearby points, in the same units as the coordinates of xy.

## Details

Generally used to remove redundant points that could produce pseudo-replication, and to limit sampling bias by disaggregating clusters of points.

## Value

A data frame with the same columns as xy with points separated by the defined minimum distance.

## See Also

thinning\_til\_n()

# Examples

```
if(interactive()){
    #load example data
    data(plant_richness_df)
    #thinning to points separated by 5 degrees
    plant_richness.thin <- thinning(
        x = plant_richness_df,
        minimum.distance = 5 #points separated by at least 5 degrees
        )
    plant_richness.thin
}</pre>
```

thinning\_til\_n Applies thinning to pairs of coordinates until reaching a given n

#### Description

Resamples a set of points with x and y coordinates by increasing the distance step by step until a given sample size is obtained.

## Usage

```
thinning_til_n(
   xy,
   n = 30,
   distance.step = NULL
)
```

## Arguments

ху	A data frame with columns named "x" and "y" representing geographic coordinates. Default: NULL
n	Integer, number of samples to obtain. Must be lower than nrow(xy). Default: 30
distance.step	Numeric, distance step used during the thinning iterations. If NULL, the one percent of the maximum distance among points in xy is used. Default: NULL

## Value

A data frame with the same columns as xy with a row number close to n.

## See Also

thinning()

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vif

#### Examples

```
if(interactive()){
    #loading example data
    data(plant_richness_df)
    #thinning to ~20 records
    plant_richness.thin <- thinning_til_n(
        x = plant_richness_df,
        n = 20
        )
    plant_richness.thin
}</pre>
```

vif

#### Variance Inflation Factor of a data frame

## Description

Computes the variance inflation factor (VIF) of the colums in a data frame. **Warning**: variables in preference.order not in colnames(x), and non-numeric columns are removed silently from x and preference.order. The same happens with rows having NA values (na.omit()) is applied). The function issues a warning if zero-variance columns are found.

## Usage

vif(x)

## Arguments

```
Х
```

Data frame with numeric columns, typically containing a set of model predictors.

#### Value

A data frame with two columns having the name of the variables in 'x' and their respective VIF values.

## See Also

auto\_vif(), auto\_cor()

#### Examples

```
if(interactive()){
  data(plant_richness_df)
  vif(plant_richness_df[, 5:21])
}
```

weights\_from\_distance\_matrix

Transforms a distance matrix into a matrix of weights

## Description

Transforms a distance matrix into weights (1/distance.matrix) normalized by the row sums. Used to compute Moran's I values and Moran's Eigenvector Maps. Allows to apply a threshold to the distance matrix before computing the weights.

## Usage

```
weights_from_distance_matrix(
   distance.matrix = NULL,
   distance.threshold = 0
)
```

#### Arguments

distance.matrix Distance matrix. Default: NULL. distance.threshold Numeric, positive, in the range of values of distance.matrix. Distances below this value in the distance matrix are set to 0., Default: 0.

## Value

A weighted distance matrix.

# Examples

```
if(interactive()){
    #loading example distance matrix
    data(distance_matrix)
    #computing matrix of weights
    distance.matrix.weights <- weights_from_distance_matrix(
        distance.matrix = distance_matrix</pre>
```

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) distance.matrix.weights

}

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