Package 'enmpa'

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```
Title Ecological Niche Modeling using Presence-Absence Data
Version 0.1.8
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Description A set of tools to perform Ecological Niche Modeling with
      presence-absence data. It includes algorithms for data partitioning,
      model fitting, calibration, evaluation, selection, and prediction.
      Other functions help to explore signals of ecological niche using univariate
      and multivariate analyses, and model features such as variable response
      curves and variable importance. Unique characteristics of this package are
      the ability to exclude models with concave quadratic responses, and the
      option to clamp model predictions to specific variables. These tools are
      implemented following principles proposed in
      Cobos et al., (2022) <doi:10.17161/bi.v17i.15985>,
      Cobos et al., (2019) <doi:10.7717/peerj.6281>,
      and Peterson et al., (2008) <doi:10.1016/j.ecolmodel.2007.11.008>.
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```

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Description

Creates candidate models based on distinct parameter settings, evaluates models, and selects the ones that perform the best.

calibration_glm 3

Usage

Arguments

data frame or matrix of data to be used in model calibration. Columns represent

dependent and independent variables.

dependent (character) name of dependent variable.

independent (character) vector of name(s) of independent variable(s).

weights (numeric) a vector with the weights for observations.

response_type (character) a character string that must contain "l", "p", "q" or a combination of

them. l = lineal, q = quadratic, p = interaction between two variables. Default =

"1".

formula_mode (character) a character string to indicate the strategy to create the formulas for

candidate models. Options are: "light", "moderate", "intensive", or "complex". Default = "moderate". "complex" returns only the most complex formula de-

fined in response_type.

minvar (numeric) minimum number of independent variables in formulas.

maxvar (numeric) maximum number of independent variables in formulas.

 $user_formulas$ (character) vector with formula(s) to test. Default = NULL.

cv_kfolds (numeric) number of folds to use for k-fold cross-validation exercises. Default

= 5. Ignored if partition_index is defined.

partition_index

list of indices for cross-validation in k-fold. The default, NULL, uses the func-

tion kfold_partition.

seed (numeric) a seed for k-fold partitioning.

n_threshold (logical) number of threshold values to produce evaluation metrics. Default =

100.

selection_criterion

(character) criterion used to select best models, options are "TSS" and "ESS".

Default = "TSS".

exclude_bimodal

(logical) whether to filter out models with one or more variables presenting con-

cave responses. Default = FALSE.

tolerance (numeric) value to modify the limit value of the metric used to filter models

during model selection if none of the models meet initial considerations. Default

= 0.01

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out_dir	(character) output directory name to save the main calibration results. Default = NULL.
parallel	(logical) whether to run on parallel or sequential. Default = FALSE.
n_cores	(numeric) number of cores to use. Default = number of free processors - 1.
verbose	(logical) whether to print messages and show progress bar. Default = TRUE

Details

Model evaluation is done considering the ability to predict presences and absences, as well as model fitting and complexity. Model selection consists of three steps: 1) a first filter to keep the models with ROC AUC >= 0.5 (statistically significant models), 2) a second filter to maintain only models that meet the selection_criterion ("TSS": TSS >= 0.4; or "ESS": maximum Accuracy tolerance), and 3) from those, pick the ones with delta AIC <= 2.

formula_mode options determine what strategy to iterate the predictors defined in type for creating models:

- light.- returns simple iterations of complex formulas.
- moderate. returns a comprehensive number of iterations.
- intensive.— returns all possible combination. Very time-consuming for 6 or more independent variables.
- complex. returns only the most complex formula.

Value

An object of the class enmpa_calibration containing: selected models, a summary of statistics for all models, results obtained in cross-validation for all models, original data used, weights, and data-partition indices used.

cal_res 5

cal_res

Example of results obtained from GLM calibration using enmpa

Description

An object of the class enmpa_calibration storing the results from GLM calibration.

Usage

```
cal_res
```

Format

An object of class enmpa_calibration with results from the function \link{calibration_glm}.

Examples

```
data("cal_res", package = "enmpa")
str(cal_res)
```

enmpa

enmpa: Ecological Niche Modeling using Presence-Absence Data

Description

enmpa contains a set of tools to perform detailed Ecological Niche Modeling using presence-absence data.

Details

It includes algorithms for data partitioning, model fitting, calibration, evaluation, selection, and prediction. Other functions help to explore model features as such variable response curves and variable importance.

Main functions in enmpa

```
calibration_glm, evaluation_stats, fit_glms, fit_selected, get_formulas, independent_eval1, kfold_partition, model_selection model_validation, niche_signal, optimize_metrics, predict_glm, predict_selected, response_curve, var_importance
```

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See Also

Useful links:

- https://github.com/Luisagi/enmpa
- Report bugs at https://github.com/Luisagi/enmpa/issues

enmpa_calibration

Constructor of S3 objects of class enmpa_calibration

Description

Constructor of S3 objects of class enmpa_calibration

Usage

Arguments

selected date.frame with information about selected models.

summary data.frame a summary of statistics for all models.

calibration_results

data.frame with results obtained from cross-validation for all models.

data data.frame or matrix with the input data used for calibration.

partitioned_data

a list of partition indices.

weights (numeric) a vector with the weights for observations. Default = NULL.

Value

An S3 object of class enmpa_calibration.

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enmpa_fitted_models

Constructor of S3 objects of class enmpa_fitted_models

Description

Constructor of S3 objects of class enmpa_fitted_models

Usage

```
new_enmpa_fitted_models(glms_fitted, selected, data, weights = NULL)
```

Arguments

glms_fitted a list of fitted GLMs.

selected date.frame with information about selected models.

data data.frame or matrix with the input data used for calibration.

weights (numeric) a vector with the weights for observations. Default = NULL.

Value

An S3 object of class enmpa_fitted_models.

enm_data

Example data used to run model calibration exercises

Description

A dataset containing information on presence and absence, and independent variables used to fit GLM models.

Usage

enm_data

Format

A data frame with 5627 rows and 3 columns.

Sp numeric, values of 0 = absence and 1 = presence.

bio_1 numeric, temperature values.

bio_12 numeric, precipitation values.

```
data("enm_data", package = "enmpa")
head(enm_data)
```

fit_selected

evaluation_stats

Summary of evaluation statistics for candidate models

Description

Calculate median and standard deviation of evaluation results for all candidate models considering cross-validation kfolds.

Usage

```
evaluation_stats(evaluation_results, bimodal_toexclude = FALSE)
```

Arguments

evaluation_results

data.frame model evaluation results. These results are the output of the function model_validation.

bimodal_toexclude

(logical) whether models in which binomial variable response curves were detected will be excluded during selection processes.

Value

A data frame with the mean and standard deviation for all metrics considering cross-validation kfolds.

Examples

```
# data
data("cal_res", package = "enmpa")
all_res <- cal_res$calibration_results[, -1]

# statistics for all evaluation results
evaluation_stats(all_res, bimodal_toexclude = TRUE)</pre>
```

fit_selected

Fitting selected GLMs models

Description

Functions to facilitate fitting multiple GLMs.

```
fit_selected(glm_calibration)
fit_glms(formulas, data, weights = NULL, id = NULL)
```

get_formulas 9

Arguments

```
a list resulting from calibration_glm. Models fitted are those in the slot "selected".

formulas (character) a vector containing the formula(s) for GLM(s).

data data.frame with the dependent and independent variables.

weights (numeric) a vector with the weights for observations. Default = NULL.

id (character) id code for models fitted. Default = NULL.
```

Value

A list of fitted GLMs.

For fit_selected, an enmpa fitted models object.

Examples

get_formulas

Get GLM formulas according to defined response types

Description

Generate GLM formulas for independent variables predicting a dependent variable, taking into account response types required. All possible combinations of variables can be created using arguments of the function.

10 get_formulas

Usage

Arguments

dependent (character) name of dependent variable.

independent (character) vector of name(s) of independent variable(s).

type (character) a character string that must contain "l", "p", "q" or a combination of

them. l = lineal, q = quadratic, p = interaction between two variables. Default =

"1".

mode (character) (character) a character string to indicate the strategy to create the

formulas for candidate models. Options are: "light", "moderate", "intensive", or

"complex". Default = "moderate".

minvar (numeric) minimum number of independent variables in formulas.

maxvar (numeric) maximum number of independent variables in formulas.

complex (logical) whether to return the most complex formula.

var_names sames as independent.

string same as type.

Details

mode options determine what strategy to iterate the predictors defined in type for creating models:

- **light**.– returns simple iterations of complex formulas.
- moderate. returns a comprehensive number of iterations.
- intensive.— returns all possible combination. Very time-consuming for 6 or more independent variables.
- **complex**.– returns only the most complex formula.

Value

A character vector containing the resulting formula(s).

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Examples

```
# example variables
dep <- "sp"
ind <- c("temp", "rain", "slope")

# The most complex formula according to "type"
get_formulas(dep, ind, type = "lqp", mode = "complex")

# mode = 'light', combinations according to type
get_formulas(dep, ind, type = "lqp", mode = "light")

# mode = 'light', combinations according to type
get_formulas(dep, ind, type = "lqp", mode = "intensive")</pre>
```

independent_eval1

Evaluate final models using independent data

Description

Final evaluation steps for model predictions using an independent dataset (not used in model calibration).

Usage

Arguments

prediction (numeric) vector or SpatRaster object. If numeric, predicted values in independent data (for independent_eval01), or the entire area of prediction (for independent_eval1). If SpatRaster prediction over the area of interest. threshold (numeric) the lowest predicted probability value for an occurrence point. This value must be defined for presences-only data. Default = NULL. test_prediction (numeric) vector of predictions for independent data. Default = NULL. lon_lat matrix or data frame of coordinates (longitude and latitude, in that order) of independent data. Points must be located within the valid area of prediction. For independent_eval01 they must correspond with values in observation. observation (numeric) vector of observed (known) values of presence or absence to test against prediction (if numeric) or values of prediction in lon_lat.

Value

A data.frame or list containing evaluation results.

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Examples

```
# Independent test data based on coordinates (lon/lat WGS 84) from presence
# and absences records
data("test", package = "enmpa")
head(test)
# Loading a model prediction
pred <- terra::rast(system.file("extdata", "proj_out_wmean.tif",</pre>
                                 package = "enmpa"))
terra::plot(pred)
# Evaluation using presence-absence data
independent_eval01(prediction = pred, observation = test$Sp,
                   lon_lat = test[, 2:3])
# Evaluation using presence-only data
test_p_only <- test[test$Sp == 1, ]</pre>
th_maxTSS <- 0.1274123
                                     # threshold based on the maxTSS
independent_eval1(prediction = pred, threshold = th_maxTSS,
                  lon_lat = test_p_only[, 2:3])
```

kfold_partition

K-fold data partitioning

Description

Creates indices to partition available data into k equal-sized subsets or folds, maintaining the global proportion of presence-absences in each fold.

Usage

```
kfold_partition(data, dependent, k = 2, seed = 1)
```

Arguments

data data.frame or matrix containing at least two columns.

dependent (character) name of column that contains the presence-absence records (1-0).

k (numeric) the number of groups that the given data is to be split into.

seed (numeric) integer value to specify an initial seed. Default = 1.

Value

A list of vectors with the indices of rows corresponding to each fold.

model_selection 13

Examples

model_selection

Selection of best candidate models considering various criteria

Description

Applies a series of criteria to select best candidate models.

Usage

Arguments

evaluation_stats

data.frame with the statistics of model evaluation results. These results are the output of the function evaluation_stats.

criterion

(character) metric used as the predictive criterion for model selection.

exclude_bimodal

(logical) whether to exclude models in which binomial variable response curves were detected.

tolerance (numeric)

Value

A data.frame with one or more selected models.

```
# data
data("cal_res", package = "enmpa")
eval_stats <- cal_res$summary[, -1]

# selecting best model
selected_mod <- model_selection(eval_stats, exclude_bimodal = TRUE)</pre>
```

model_validation

_validation
tion Model validation options

Description

Model evaluation using entire set of data and a k-fold cross validation approach. Models are assessed based on discrimination power (ROC-AUC), classification ability (accuracy, sensitivity, specificity, TSS, etc.), and the balance between fitting and complexity (AIC).

Usage

Arguments

formula	(character) expression to be used as a glm formula.			
data	data.frame with dependent and independent variables.			
family	a family object for models used by functions such as glm. Default = binomial(link = "logit").			
weights	(numeric) vector with weights for observations. Default = NULL.			
cv	(logical) whether to use a k-fold cross validation for evaluation. Default = FALSE.			
partition_index				
	list of indices for cross validation in k-fold. Obtained with the function $kfold_partition$. Default = NULL.			
k	(numeric) number of folds for a new k-fold index preparation. Ignored if $partition_index$ is defined or if $cv = FALSE$. Default = $NULL$.			
dependent	(character) name of dependent variable. Ignore if cv = FALSE. Default = NULL.			
n_threshold	(numeric) number of threshold values to be used for ROC. Default = 100.			
keep_coefficients				
	(logical) whether to keep model coefficients. Default = FALSE.			
seed	(numeric) a seed number. Default = 1.			

Value

A data.frame with results from evaluation.

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Examples

```
# Load species occurrences and environmental data.
data("enm_data", package = "enmpa")
head(enm_data)

# Custom formula
form <- c("Sp ~ bio_1 + I(bio_1^2) + I(bio_12^2)")

# Model evaluation using the entire set of records
model_validation(form, data = enm_data)

# Model evaluation using a k-fold cross-validation (k = 3)
model_validation(form, data = enm_data, cv = TRUE, k = 3, dependent = "Sp")</pre>
```

niche_signal

Niche Signal detection using one or multiple variables

Description

Identifies whether a signal of niche can be detected using one or multiple variables. This is an implementation of the methods developed by Cobos & Peterson (2022) doi:10.17161/bi.v17i. 15985 that focuses on identifying niche signals in presence-absence data.

Usage

Arguments

data	matrix or data.frame containing at least the following information: a column representing condition (positive = 1 or negative = 0), and one or more columns representing environmental variables.
condition	(character) name of the column with numeric information about detection (positive $= 1$ or negative $= 0$).
variables	(character) vector of one or more names of columns to be used as environmental variables. If method = "univariate", only one variable is used; for method = "permanova", multiple variables can be used.
method	(character) name of the method to be used for niche comparison. Default = "univariate".

optimize_metrics

permanova_method

(character) name of the dissimilarity index to be used as method in adonis2.

See all options in vegdist. Default = "mahalanobis".

iterations (numeric) number of iterations to be used in analysis. Default = 1000. If method

= "permanova", permutations = iterations - 1.

set_seed (numeric) integer value to specify a initial seed. Default = 1.

verbose (logical) whether or not to print messages about the process. Default = TRUE.

... other arguments to be passed to adonis2.

variable (character) name of the column containing data to be used as environmental

variable.

permutations number of permutations to be performed.

Value

A list with results from analysis depending on method.

Examples

optimize_metrics

Find threshold values to produce three optimal metrics

Description

The metrics true skill statistic (TSS), sensitivity, specificity are explored by comparing actual vs predicted values to find threshold values that produce sensitivity = specificity, maximum TSS, and a sensitivity value of 0.9.

```
optimize_metrics(actual, predicted, n_threshold = 100)
```

plot_importance 17

Arguments

actual	(numeric) vector of actual values $(0, 1)$ to be compared to predicted values after applying a threshold.
predicted	(numeric) vector of predicted probability values to be thresholded and compared to actual.
n_threshold	(numeric) number of threshold values to be used. Default = 100.

Value

A list containing a data frame with the resulting metrics for all threshold values tested, and a second data frame with the results for the threshold values that produce sensitivity = specificity (ESS), maximum TSS (maxTSS), and a sensitivity value of 0.9 (SEN90).

Examples

```
# example data
act <- c(rep(1, 20), rep(0, 80))
pred <- c(runif(20, min = 0.4, max = 0.7), runif(80, min = 0, max = 0.5))
# run example
om <- optimize_metrics(actual = act, predicted = pred)
om$optimized</pre>
```

plot_importance

Plot variable importance

Description

Visualization of the results obtained with the function var_importance.

Usage

Arguments

х	data.frame output from var_importance.
xlab	(character) a label for the x axis.
ylab	(character) a label for the y axis.
main	(character) main title for the plot.
extra_info	(logical) when results are from more than one model, it adds information about the number of models using each predictor and the mean contribution found.
	additional arguments passed to barplot or boxplot.

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Value

A plot

Examples

plot_niche_signal

Plot Niche Signal results

Description

Plots to interpret results from niche_signal tests (Cobos & Peterson (2022) doi:10.17161/bi.v17i.15985).

plot_niche_signal 19

Arguments

lwd

niche_signal_list list of results from niche_signal. (character) name of the statistic for which results will be explored when results statistic come for univariate analysis. Default = "mean". Options are: "mean", "median", "SD", and "range". variables (character) name of variables to used in plots when results come from analysis using the permanova method. The default, NULL, uses the first two variables. ellipses (logical) whether to use ellipses to represent all and positive data when results come from PERMANOVA. The default, FALSE, plots points instead. level (numeric) value from 0 to 1 representing the limit of the ellipse to be plotted. Default = 0.99. breaks breaks in the histogram as in hist. Default = "Sturges". (character) title for plot. Default = "". main xlab (character) x axis label. Default = NULL. For results from PERMANOVA, appropriate variable names are used. ylab (character) y axis label. Default = NULL. For univariate results, the default turn into "Frequency". For results from PERMANOVA, appropriate variable names are used. h_col a color to be used to fill the bars of histograms. Default = "lightgray". (numeric) value by which plotting text and symbols should be magnified relative h_cex to the default in histograms. Default = 0.8. lty (numeric) line type. See options in par. Default = 2. A vector of length = 2 can be used, in order, all and positive.

line color for observed value of positives and confidence intervals. Default = c("blue", "black").
 color of ellipse lines for all and positive data. Default = c("black", "red").

(numeric) line width. See options in par. Default = 1. A vector of length = 2

pch point type. See options in points. Default = 19. A vector of length = 2 can be

used, in order, all and positive.

can be used, in order, all and positive.

pt_cex (numeric) value by which points will be magnified. Values for all and positive points are recommended. Default = c(1.3, 0.8).

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Value

A plot.

Examples

predict_glm

Extension of glm predict to generate predictions of different types

Description

Obtains predictions from a fitted generalized linear model objects. It also allows the clamping option to restrict extrapolation in areas outside the calibration area.

```
predict_glm(
  model,
  newdata,
  data = NULL,
  extrapolation_type = "E",
```

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```
restricted_vars = NULL,
type = "response"
)
```

Arguments

model a glm object.

newdata a data.frame or matrix with the new data to project the predictions.

data.frame or matrix of data used in the model calibration step. Default = NULL.

extrapolation_type

(character) to indicate extrapolation type of model. Models can be transferred with three options: free extrapolation ('E'), extrapolation with clamping ('EC'), and no extrapolation ('NE'). Default = 'E'.

restricted_vars

(character) a vector containing the names of the variables that will undergo clamping or no extrapolation. For clamping, these variables are set to minimum and maximum values established for the max and min values within calibration values. For no extrapolation, the variables outside calibration limits became NA. If no specific names are provided, the value is set to NULL by default, indicating that clamping (EC) or no extrapolation (NE) will be applied to all variables. Ignore if extrapolation_type = 'E'.

type

(character) the type of prediction required. For a default binomial model the default predictions are of log-odds (probabilities on logit scale). The default, "response", returns predicted probabilities.

Value

A SpatRaster object or a vector with predictions.

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predict_selected

Predictions for the models selected after calibration

Description

Wrapper function that facilitates the prediction of those models selected as the most robust. In addition, it allows the calculation of consensus models, when more than one model are selected.

Usage

Arguments

fitted

an enmpa-classfitted models object obtained using the functions fit_selected.

newdata

a SpatRaster, data.frame, or matrix with the new data on which to predict.

extrapolation_type

(character) to indicate extrapolation type of model. Models can be transferred with three options: free extrapolation ('E'), extrapolation with clamping ('EC'), and no extrapolation ('NE'). Default = 'E'.

restricted_vars

(character) a vector containing the names of the variables that will undergo clamping or no extrapolation. For clamping, these variables are set to minimum and maximum values established for the max and min values within calibration values. For no extrapolation, the variables outside calibration limits became NA. If no specific names are provided, the value is set to NULL by default, indicating that clamping (EC) or no extrapolation (NE) will be applied to all variables. Ignore if extrapolation_type = 'E'.

type

(character) the type of prediction required. For a default binomial model the default predictions are of log-odds (probabilities on logit scale). The default, "response", returns predicted probabilities.

consensus

(logical) valid if newdata is a SpatRaster, whether to produce consensus results obtained by combining the predictions from the collection of selected models. By default consensuses are calculated using the mean, median, variance, and weighted average using the AIC weights. Default = TRUE.

Value

A list with predictions of selected models on the newdata and fitted selected model(s). Consensus predictions are added if multiple selected models exits and if newdata is a SpatRaster object.

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Examples

```
# Load a fitted selected model
data(sel_fit, package = "enmpa")

# Load raster layers to be projected
env_vars <- terra::rast(system.file("extdata", "vars.tif", package = "enmpa"))

# Predictions (only one selected mode, no consensus required)
preds <- predict_selected(sel_fit, newdata = env_vars, consensus = FALSE)

# Plot prediction
terra::plot(preds$predictions)</pre>
```

print

Print a short version of elements in 'calibration' and 'fitted models' objects

Description

Print a short version of elements in 'calibration' and 'fitted models' objects

Usage

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

Arguments

x object of enmpa_fitted_models or enmpa_calibration
 ... additional arguments affecting the summary produced. Ignored in these functions.

proc_enm

Partial ROC calculation

Description

proc applies partial ROC tests to model predictions.

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Arguments

test_prediction

(numeric) vector of model predictions for testing data.

prediction SpatRaster or numeric vector of model predictions to be evaluated.

threshold (numeric) value from 0 to 100 to represent the percentage of potential error (E)

that the data could have due to any source of uncertainty. Default = 5.

sample_percentage

(numeric) percentage of testing data to be used in each bootstrapped process for

calculating the partial ROC. Default = 50.

iterations (numeric) number of bootstrap iterations to be performed; default = 500.

Details

Partial ROC is calculated following Peterson et al. (2008) doi:10.1016/j.ecolmodel.2007.11.008.

Value

A list with the summary of the results and a data.frame containing the AUC values and AUC ratios calculated for all iterations.

Examples

response_curve

Variable response curves for GLMs

Description

A view of variable responses in models. Responses based on single or multiple models can be provided.

response_curve 25

Arguments

fitted	an object of class glm, a list of GLMs obtained using the function fit_glms , or an object enmpa_fitted_models from $fit_selected$.
variable	(character) name of the variables to be plotted.
data	data.frame or matrix of data used in the model calibration step. Default = $NULL$.
modelID	(character) vector of ModelID(s) to be considered when the fitted models is an enmpa_fitted_object. By default all models are included. Default = NULL.
n	(numeric) an integer guiding the number of breaks. Default = 100
new_data	a SpatRaster, data.frame, or matrix of variables representing the range of variable values in an area of interest. Default = NULL. It must be defined in case the model entered does not explicitly include a data component.
extrapolate	(logical) whether to allow extrapolation to study the behavior of the response outside the calibration limits. Ignored if new_data is defined. Default = TRUE.
xlab	(character) a label for the x axis. The default, NULL, uses the name defined in variable.
ylab	(character) a label for the y axis. Default = "Probability".
col	(character) color for lines. Default = "red".
	additional arguments passed to plot.

Details

The function calculates these probabilities by focusing on a single environmental variable while keeping all other variables constant at their mean values.

When responses for multiple models are to be plotted, the mean and confidence intervals for the set of responses are calculated using a GAM.

Value

A plot with the response curve for a variable.

```
# Load a fitted selected model
data(sel_fit, package = "enmpa")

# Response curve for single models
response_curve(sel_fit$ModelID_7, variable = "bio_1")

# Response curve when model(s) are in a list (only one model in this one)
response_curve(sel_fit, variable = "bio_12")
```

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sel_fit

Example of selected models fitted

Description

An object of the class enmpa_fitted_models containing fitted selected model(s) and the information from model evaluation for such model(s).

Usage

```
sel_fit
```

Format

A list with four elements.

```
glms_fitted a fitted glm (ModelID_7).
```

selected a data.frame with results from evaluation of ModelID_7

data a data.frame containing information on presence and absence, and independent variables used to fit GLM models.

weights a vector with the weights for observations.

Examples

```
data("sel_fit", package = "enmpa")
```

summary

Summary of 'calibration' and 'fitted models'

Description

Summary of 'calibration' and 'fitted models'

Usage

```
## $3 method for class 'enmpa_calibration'
summary(object, ...)
## $3 method for class 'enmpa_fitted_models'
summary(object, ...)
```

Arguments

```
object of class enmpa_calibration or enmpa_fitted_models
```

... additional arguments affecting the summary produced. Ignored in these func-

tions.

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Value

A printed summary.

test

Example data used to test models

Description

A dataset containing information on presence and absence, and independent variables used to fit GLM models.

Usage

test

Format

A data frame with 100 rows and 3 columns.

Sp numeric, values of 0 = absence and 1 = presence.

lon numeric, longitude values.

lat numeric, latitude values.

Examples

```
data("test", package = "enmpa")
head(test)
```

var_importance

Variable importance for GLMs

Description

Calculates the relative importance of predictor variables based on the concept of explained deviance. This is achieved by fitting a GLMs multiple times, each time leaving out a different predictor variable to observe its impact on the model's performance.

```
var_importance(fitted, modelID = NULL, data = NULL)
```

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Arguments

fitted	an object of class glm, a list of GLMs obtained using the function fit_glms, or an object enmpa_fitted_models from fit_selected.
modelID	(character) vector of $ModelID(s)$ to be considered when the fitted input is of the class enmpa_fitted_models. By default all models are included. Default = $NULL$.
data	data.frame or matrix of data used in the model calibration step. It must be de-

fined in case the model entered does not explicitly include a data component.

Default = NULL.

Details

The process begins by fitting the full GLM model, which includes all predictor variables. Subsequently, separate GLM models are fitted, excluding one variable at a time to assess the influence of its absence on the model's performance. By systematically evaluating the effect of removing each predictor variable, the function provides valuable insights into their individual contributions to the model's overall performance and explanatory power.

Value

A data.frame containing the relative contribution of each variable. An identification for distinct models is added if fitted contains multiple models.

```
# Load a fitted selected model
data(sel_fit, package = "enmpa")

# Variable importance for single models
var_importance(sel_fit, modelID = "ModelID_7")

# Variable importance for multiple models (only one model in this list)
var_importance(sel_fit)
```

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