Package ‘emil’
July 30, 2018

Encoding UTF-8
Type Package
Title Evaluation of Modeling without Information Leakage
Version 2.2.10
Date 2018-07-30
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Description A toolbox for designing and evaluating predictive models with
resampling methods. The aim of this package is to provide a simple and
efficient general framework for working with any type of prediction
problem, be it classification, regression or survival analysis, that is
easy to extend and adapt to your specific setting. Some commonly used
methods for classification, regression and survival analysis are included.
Depends R (>= 3.0.2)
Imports data.table, dplyr, ggplot2 (>= 2.0.0), graphics, grDevices,
lazyeval, magrittr, methods, stats, tidyr, utils
Suggests caret, cmprsk, e1071, Hmisc, MASS, parallel, party,
pamr, randomForest, RColorBrewer, rpart, survival (>= 2.42-5),
testthat (>= 0.9.1)
LinkingTo Rcpp (>= 0.12.1)
License GPL (>= 2)
LazyLoad yes
LazyData yes
URL https://github.com/Molmed/emil
BugReports https://github.com/Molmed/emil/issues
RoxygenNote 6.0.1
NeedsCompilation yes
Repository CRAN
Date/Publication 2018-07-30 12:00:06 UTC
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as.modeling_procedure  *Coerce to modeling procedure*

**Description**

Coerce to modeling procedure

**Usage**

as.modeling_procedure(x, ...)

**Arguments**

- `x`  
  modeling procedure.
- `...`  
  Ignored (kept for S3 consistency).

**Value**

Modeling procedure

**Author(s)**

Christofer Bäcklin

**Examples**

as.modeling_procedure("lda")

dichotomize  *Dichotomize time-to-event data*

**Description**

Convert time-to-event data (typically created with the `Surv` function) to factor or integer.

**Usage**

dichotomize(x, time, to_factor)

**Arguments**

- `x`  
  `Surv` vector.
- `time`  
  Time point to dichotomize at.
- `to_factor`  
  Depending on the type of `x` the return value may be integer or factor. Set this argument to explicitly state the return type.
Details

If no time point is given the observation times will be stripped, leaving only the event types. If a time point is given observations with events occurring before time will be labelled by their event type, observations with events occurring after time will be labelled as “no event”, and observations censored before time will be considered as missing information.

Value

Integer vector or factor.

Author(s)

Christofer Bäcklin

See Also

Surv

Description

The emil package implements a framework for working with predictive modeling problems without information leakage. For an overview of its functionality please read the original publication included as the package’s vignette (to be added).

Central topics and functions

Setting up modeling problems:

resample Functions for generating and resampling schemes and information on how to implement custom resampling methods.
pre_process Data pre-processing functions.
modeling_procedure Manages algorithms used for fitting models, making predictions, and extracting feature importance scores.
error_fun Performance estimation functions used to tune parameters and evaluate performance of modeling procedures.

Solving modeling problems:

fit Fit a model (according to a procedure).
tune Tune parameters of a procedure.
predict Use a fitted model to predict the response of observations.
evaluate Evaluate the performance of a procedure using resampling.
learning_curve Learning curve analysis.

Managing the results of modeling problems:
get_prediction Extract predictions from resampled modeling results.
get_tuning Extract feature importance scores of a fitted model or resampled modeling results.
get_importance Extract feature importance scores of a fitted model or resampled modeling results.
subtree Extracts results from the output of evaluate. It is essentially a recursive version of lapply and sapply.
select Interface between emil and the dplyr package for data manipulation. Can be used to subset modeling results, reorganize or summarize to help interpretation or prepare for plotting.

Methods included in the package

**Resampling methods:** See resample for information on usage and implementation of custom methods.
resample_holdout Repeated holdout.
resample_crossvalidation Cross validation.

**Data pre-processing methods:** See pre_process for information on usage and implementation of custom methods. The imputation functions can also be used outside of the resampling scheme, see impute.
pre_split Only split, no transformation.
pre_center Center data to have mean 0 of each feature.
pre_scale Center and scale data to have mean 0 and standard deviation 1.
pre_impute_median Impute missing values with feature medians.
pre_impute_knn Impute missing values with k-NN, see pre_impute_knn for details on how to set parameters.

**Modeling methods:** The following modeling methods are included in the emil package. For a complete list of available methods in both the emil package and other loaded packages, please use list_method. See modeling_procedure for information on usage and extension for information on implementation of custom methods.
cforest Conditional inference forest.
coxph Cox proportional hazards model.
glmnet Elastic net.
lasso LASSO.
lda Linear discriminant.
lm Linear model.
pamr Nearest shrunken centroids.
qda Quadratic discriminant.
randomForest Random forest.
ridge_regression Ridge regression.
rpart Decision trees.
It is also possible to incorporate any method from the ‘caret’ package by using the function fit_caret.
To search for emil compatible methods in all attached packages use the list_method function.
**Performance estimation methods**: See `error_fun` for information on usage and implementation of custom methods. Since the framework is designed to minimize the error when tuning parameters, some measures are negated, e.g. `neg_auc`.

For classification problems:

- **error_rate** Fraction of predictions that were incorrect.
- **weighted_error_rate** See its own documentation.
- **neg_auc** Negative area under ROC curve. To plot the ROC curves see `roc_curve`.
- **neg_gmpa** Negative geometric mean of class-specific prediction accuracy. Good for problems with imbalanced class sizes.

For regression problems:

- **mse** Mean square error.
- **rmse** Root mean square error.

For survival analysis problem:

- **neg_harrell_c** Negative Harrell's concordance index.

**Plotting**: Plotting is not the one of the main aims of the package and the methods that do exist mainly serves as examples for how to write your own. These exists for:

- Learning curve analyses.
- Resampling schemes.
- ROC-curves.

**Author(s)**

Christofer Bäcklin

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**error_fun**

*Performance estimation functions*

**Description**

These functions determine the performance of fitted model based on its predictions. They are used both for evaluating whole modeling procedures and to tune model parameters, i.e. find the parameter values with the best performance. The parameter tuning routine is designed to minimize its error function (or optimization criteria), which is why functions that are to be maximized must have their sign changed, like `neg_auc`.

**Usage**

```r
error_rate(truth, prediction, allow_rejection = !missing(rejection_cost), 
            rejection_cost)
```

```r
neg_auc(truth, prediction)
```

```r
rmse(truth, prediction, na.rm = FALSE)
```
mse(truth, prediction, na.rm = FALSE)

neg_harrell_c(truth, prediction, na.rm = FALSE)

**Arguments**

- `truth` The true response values, be it class labels, numeric values or survival outcomes.
- `prediction` A prediction object.
- `allow_rejection` If FALSE missing prediction values will produce an error. If TRUE missing values will be given a cost specified by the `rejection_cost` argument.
- `rejection_cost` See the argument `allow_rejection`. If missing a rejection cost equivalent to the error rate obtained when assigning all test observations to the most common class will be used.
- `na.rm` Whether to remove missing values or not.

**Details**

Custom performance estimation functions should be implemented as follows:

```r
function(truth, prediction)

truth A vector of true responses.
prediction Prediction returned from the prediction function.
```

In most cases the true response and the predictions are of the same type, e.g. true and fitted values in a regression or class labels in a classification problem, but it is not a requirement. An example of different types could be if the prediction function produce class probabilities for all classes rather than one label, or the risks that the observations will experience the event of interest, to be compared to the actual outcome that it did occur or has not yet occurred at a specific time point. See `neg_harrell_c` for an example of the latter.

**Author(s)**

Christofer Bäcklin

**See Also**

`emil`, `neg_gmpa`, `modeling_procedure`, `extension`
**evaluate**  
isevaluateda modeling procedure

**Description**

This function performs the important task of evaluating the performance of a modeling procedure with resampling, including tuning and pre-processing to not bias the results by information leakage.

**Usage**

evaluate(procedure, x, y, resample, pre_process = pre_split, .save = c(model = TRUE, prediction = TRUE, error = TRUE, importance = FALSE), .cores = 1, .checkpoint_dir = NULL, .return_error = .cores > 1, .verbose = getOption("emil_verbose", TRUE))

**Arguments**

- **procedure**: Modeling procedure, or list of modeling procedures, as produced by `modeling_procedure`.
- **x**: Dataset, observations as rows and descriptors as columns.
- **y**: Response vector.
- **resample**: The test subsets used for parameter tuning. Leave blank to randomly generate a resampling scheme of the same kind as is used by `evaluate` to assess the performance of the whole `modeling_procedure`.
- **pre_process**: Function that performs pre-processing and splits dataset into fitting and test subsets.
- **.save**: What parts of the modeling results to return to the user. If `importance` is `FALSE` variable importance calculation will be skipped.
- **.cores**: Number of CPU-cores to use for parallel computation. The current implementation is based on `mcmC`, which unfortunately do not work on Windows systems. It can however be re-implemented by the user fairly easily by setting up a PSOCK cluster and calling `parLapply` as in the example below. This solution might be included in future versions of the package, after further investigation.
- **.checkpoint_dir**: Directory to save intermediate results to, after every completed fold. The directory will be created if it doesn’t exist, but not recursively.
- **.return_error**: If `FALSE` the entire modeling is aborted upon an error. If `TRUE` the modeling of the particular fold is aborted and the error message is returned instead of its results.
- **.verbose**: Whether to print an activity log.
Value

A list tree where the top level corresponds to folds (in case of multiple folds), the next level corresponds to the modeling procedures (in case of multiple procedures), and the final level is specified by the `.save` parameter. It typically contains a subset of the following elements:

- `error` Performance estimate of the fitted model. See `error_fun` for more information.
- `fit` Fitted model.
- `prediction` Predictions given by the model.
- `importance` Feature importance scores.
- `tune` Results from the parameter tuning. See `tune` for details.

Author(s)

Christofer Bäcklin

References


See Also

`emil`, `modeling_procedure`

Examples

```r
x <- iris[-5]
y <- iris$Species
cv <- resample("crossvalidation", y, nfold = 4, nrepeat = 4)
result <- evaluate("lda", x, y, resample=cv)

# Multiple procedures fitted and tested simultaneously. # This is useful when the dataset is large and the splitting takes a long time.
# If you name the elements of the list emil will also name the elements of the # results object in the same way.
result <- evaluate(c(Linear = "lda", Quadratic = "qda"), x, y, resample=cv)

# Multicore parallelization (on a single computer)
result <- evaluate("lda", x, y, resample=cv, .cores=2)

# Parallelization using a cluster (not limited to a single computer) # PSOCK is supported on windows too!
require(parallel)
c1 <- makePSOCKcluster(2)
clusterEvalQ(c1, library(emil))
```
Extending the emil framework with user-defined methods

Description

This page describes how to implement custom methods compatible with the functions of the emil framework, most notably \texttt{fit}, \texttt{tune}, and \texttt{evaluate}. Pre-processing and resampling is not covered here, but in the entries \texttt{pre_process} and \texttt{resample}.

Fitting models

To write and use custom model fitting functions with the emil framework, it must take the following inputs. Optional

\begin{verbatim}
function(x, y, p1, p2, p3, ..., .verbose)
\end{verbatim}

\textbf{x} The features (or variables) of the observations you want to train the model on. This is typically a matrix or data frame where each row corresponds to an observation. In case it is more natural to characterize your observations some other way, maybe as character vectors of varying length for some document classification method, \texttt{x} can be of any form you like as long as the fitting function knows how to handle it. In that case you will also need supply you own pre-processing function (see \texttt{pre_process} that can extract training and test sets from the entire data set.

See the functions \texttt{pre_pamr} and \texttt{fit_pamr} for an example of a function that does not take its data in the default way.

\textbf{y} A response vector. This is the outcome you want to model, e.g. the feature of interest in a regression, class label in a classification problem, or anything else that a fitted model will produce when given data to make predictions from.

\textbf{p1, p2, p3, ...} (Optional) Method-specific model parameters. These will all be tunable with the \texttt{tune} and \texttt{evaluate} functions. Note that you can give them any name you want, the names used here are just an example.

\textbf{.verbose} (Optional) Indentation level of log messages. Feed this to \texttt{log_message}.

The function must return everything necessary to make future predictions, but it can take any form you like. In the simplest case it is just a number of fitted parameter values, like in a least squares regression, but it could also be some big and complex structure holding an ensemble of multiple sub-models.
Making predictions

Once a model is fitted it can be used to make predictions with a prediction function, defined as such
\[
\text{function}(\text{object}, \ x, \ ...)
\]

- object A fitted model produced by the model fitting function described above.
- x Observations to make predictions on (describing features only).
- ... Parameters to the prediction functions. These are ignored by \texttt{tune} and \texttt{evaluate}, but could be convenient if the user wants to work with it manually.

The output of the prediction function must be an object that can be compared to the true response, by an error function (see below). It is typically a list with elements named "pred" for "predictions" or "risk" for estimated risks. It can also be on an arbitrary form as long as a compatible error function is used.

Calculating feature importance scores

Estimating the importance of each feature (or variable) can often be as important as making predictions. Functions for calculating or extracting feature importance scores from fitted models should be defined as follows:

\[
\text{function}(\text{object}, \ ...)
\]

- object A fitted model produced by the model fitting function described above.
- ... Parameters to the prediction functions. These are ignored by \texttt{tune} and \texttt{evaluate}, but could be convenient if the user wants to work with it manually.

The function should return a vector of length \(p\) or a \(p\)-by-\(c\) data frame where \(p\) is the number of features in the data set and \(c\) is the number of classes.

Calculating performance

See \texttt{error_fun}.

Resampling schemes

See \texttt{resample}.

Pre-processing functions

See \texttt{pre_process}.

Code style guidelines

Names of functions, arguments and variables should be written in underscore separated lower case, singular form, unabbreviated, and American English. Users are encouraged to also follow use this style when writing extensions. However, the guidelines may be violated in cases where they break the consistency with an incorporated well established package, see for example \texttt{fit_randomForest} which according to the guidelines should be \texttt{fit_randomforest} or \texttt{fit_random_forest}.

A few exceptions to the rule against abbreviations exists, namely ‘fun’ and ‘function’ and ‘dir’ for ‘directory’. These are only used for arguments to indicate the type of value that is accepted.
factor_to_logical

Author(s)

Christofer Bäcklin

See Also

emil, error_fun, pre_process, resample

factor_to_logical  Convert factors to logicals

Description

Factors are converted to logical vectors or matrices depending on the number of levels. Ordered factors are converted to matrices where each column represent a level, coded TRUE for observations that match the level and FALSE otherwise. Unordered factors are converted in a similar way but coded TRUE for observations that match the level or a higher level. Interpreted in words, the star rating example below returns a matrix containing a column named “3 stars” that contains TRUE for observations with at least three stars and FALSE for observations with fewer than three stars.

Usage

factor_to_logical(x, base = 1L, drop = TRUE)

Arguments

x    Factor.
base  Level to consider as the basis for comparison. Can be either integer or character. Note that base = 4 is interpreted as a level named "4", but base = 4L is interpreted as the fourth level.
drop  Whether to keep the base level. The base level column never holds any information that cannot be deduced from the remaining columns.

Author(s)

Christofer Bäcklin

Examples

# Binary factor
email <- factor(sample(2, 20, TRUE), labels=c("unverified", "verified"))
factor_to_logical(email)

# Unordered multi-level factors
wine_preferences <- factor(sample(3, 20, TRUE),
                          labels=c("red", "white", "none"))
factor_to_logical(wine_preferences, base="none")
fruit <- factor(sample(4, 20, TRUE),
    labels = c("apple", "banana", "cantaloup", "durian"))
fruit[sample(length(fruit), 3)] <- NA
factor_to_logical(fruit, drop=FALSE)

# Ordered factor
rating <- factor(1:5, labels = paste(1:5, "stars"), ordered=TRUE)
factor_to_logical(rating)

# Ordered factor with custom base
tie_break <- factor(1:5,
    ordered = TRUE)
tie_status <- as.data.frame(
    factor_to_logical(tie_break, base="Deuce", drop=FALSE)
)
print(tie_status)
tie_break[tie_status$AdvAlice]
tie_break[tie_status$SetBob]
tie_break[tie_status$Deuce]

**fill**

*Replace values with something else*

**Description**

Replace values with something else

**Usage**

fill(x, pattern, replacement, invert = FALSE)

na_fill(x, replacement)

**Arguments**

- **x**  
  Variable containing NAs.
- **pattern**  
  The values in x to be replaced. Can also be a function.
- **replacement**  
  The value which is to replace the values matching pattern.
- **invert**  
  Whether to fill all values except the ones matching pattern.

**Value**

An imputed version of x.

**Author(s)**

Christofer Bäcklin
fit

Examples

```r
fill(1:10, function(x) x %% 2 == 1, 0)
na_fill(c(1,2,NA,4,5), 3)
```

### Fit a model

**Description**

Fits a model according to a modeling procedure. If the procedure contains untuned parameters they will automatically be tuned prior to fitting.

**Usage**

```r
fit(procedure, x, y, ..., .verbose = getOption("emil_verbose", FALSE))
```

**Arguments**

- `procedure`: Modeling procedure, or list of modeling procedures, as produced by `modeling_procedure`.
- `x`: Dataset, observations as rows and descriptors as columns.
- `y`: Response vector.
- `...`: Sent to `tune`, in case tuning is required, which will pass them on to `evaluate`.
- `.verbose`: Whether to print an activity log. Set to `-1` to suppress all messages.

**Value**

A list of fitted models.

**Author(s)**

Christofer Bäcklin

**See Also**

`emil, modeling_procedure, evaluate, tune, predict, get_importance`

**Examples**

```r
mod <- fit("lda", x=iris[-5], y=iris$Species)
```
fit_caret

Fit a model using the \texttt{caret} package

\textbf{Description}

Fit a model using the \texttt{caret} package

\textbf{Usage}

\begin{verbatim}
fit_caret(x, y, ...)
\end{verbatim}

\textbf{Arguments}

\begin{itemize}
  \item \textit{x} \hspace{1cm} Descriptors.
  \item \textit{y} \hspace{1cm} Response.
  \item \text{...} \hspace{1cm} Sent to \texttt{train}.
\end{itemize}

\textbf{Author(s)}

Christofer Bäcklin

\textbf{References}


fit_cforest

\textit{Fit conditional inference forest}

\textbf{Description}

A \texttt{cforest} is a random forest based on conditional inference trees, using the implementation in the \texttt{party} package. These trees can be used for classification, regression or survival analysis, but only the survival part has been properly tested so far.

\textbf{Usage}

\begin{verbatim}
fit_cforest(x, y, formula = y ~ ., ctrl_fun = party::cforest_unbiased, ...)
\end{verbatim}

\textbf{Arguments}

\begin{itemize}
  \item \textit{x} \hspace{1cm} Dataset, observations as rows and descriptors as columns.
  \item \textit{y} \hspace{1cm} Responses.
  \item \textit{formula} \hspace{1cm} Formula linking response to descriptors.
  \item \textit{ctrl_fun} \hspace{1cm} Which control function to use, see \texttt{cforest_control}.
  \item \text{...} \hspace{1cm} Sent to the function specified by \texttt{ctrl_fun}.
\end{itemize}
Details

The parameters to `cforest` are set using a `cforest_control` object. You should read the documentation as the default values are chosen for technical reasons, not predictive performance! Pay special attention to `mtry` which is set very low by default.

Value

A fitted `cforest` model.

Author(s)

Christofer Bäcklin

References


See Also

`emil`, `predict_cforest`, `modeling_procedure`

fit_coxph

Fit Cox proportional hazards model

Description

Fit Cox proportional hazards model

Usage

`fit_coxph(x, y, formula = y ~ ., ...)`

Arguments

- `x` Dataset.
- `y` Response. Required if formula is missing.
- `formula` See `coxph`.
- `...` Sent to `coxph`.
Value
Fitted Cox proportional hazards model.

Author(s)
Christofer Bäcklin

See Also
predict_coxph

Examples

```r
if(requireNamespace("survival")){
  data("ovarian", package = "survival")
  model <- fit(
    modeling_procedure(
      method = "coxph",
      parameter = list(formula = list(survival::Surv(futime, fustat) ~ age)),
      x = ovarian, y = NULL
    )
  )
  predict(model, ovarian[11:16])
}
```

fit_glmnet

Fit elastic net, LASSO or ridge regression model

Description
Using the glmnet package implementation.

Usage

```r
fit_glmnet(x, y, family, nfolds, foldid, alpha = 1, lambda = NULL, ...)
fit_ridge_regression(...)  # for ridge regression
fit_lasso(...)             # for lasso regression
```

Arguments

- `x`: Dataset.
- `y`: Response vector. Can be of many different types for solving different problems, see glmnet.
- `family`: Determines the type of problem to solve. Auto detected if `y` is numeric or survival. See family for details.
fit_lda

```r
fit_lda
```

### Arguments

- `x`  
  Dataset, numerical matrix with observations as rows.
- `y`  
  Class labels, factor.
- `...`  
  Sent to `lda`.

### Usage

```r
fit_lda(x, y, ...)
```

### Description

Wrapper for the MASS package implementation.

### Details

The alpha parameter of `glmnet` controls the type of penalty. Use 0 (default) for lasso only, 1 for ridge only, or an intermediate for a combination. This is typically the parameter to tune on. The shrinkage, controlled by the lambda parameter, can be left unspecified for internal tuning (works the same way as `fit_glmnet`).

### Value

Fitted elastic net model.

### Author(s)

Christofer Bäcklin

### References


### See Also

- `emil`
- `predict_glmnet`
- `importance_glmnet`
Value
Fitted linear discriminant.

Author(s)
Christofer Bäcklin

References

See Also
emil, predict_lda, modeling_procedure

---

fit_lm

*Fit a linear model fitted with ordinary least squares*

Description
Based on `lm`.

Usage
`fit_lm(x, y, formula = y ~ ., ...)`

Arguments
- `x` Descriptors.
- `y` Response, numeric.
- `formula` See `lm`.
- `...` Sent to `lm`.

Value
Fitted linear model.

Author(s)
Christofer Bäcklin

See Also
emil, predict_lm, modeling_procedure
**fit_naive_bayes**

*Fit a naive Bayes classifier*

**Description**

Fit a naive Bayes classifier

**Usage**

```r
fit_naive_bayes(x, y, ...)
```

**Arguments**

- `x`: Dataset, observations as rows.
- `y`: Response vector.
- `...`: Send to `naiveBayes`.

**Author(s)**

Christofer Bäcklin

---

**fit_pamr**

*Fit nearest shrunken centroids model.*

**Description**

Wrapped version of the `pamr` package implementation. Note that this function uses internal cross-validation for determining the value of the shrinkage threshold.

**Usage**

```r
fit_pamr(x, y, error_fun, cv, nfold, threshold = NULL, ..., thres_fun = function(thr, err) median(thr[err == min(err)]), slim = FALSE)
```

**Arguments**

- `x`: Dataset, numerical matrix with observations as rows.
- `y`: Class labels, factor.
- `error_fun`: Error function for tuning.
- `cv`: Cross-validation scheme for shrinkage tuning. It should be supplied on one of the following forms:
  - Resampling scheme produced with `resample` or `resample_holdout`.
  - List with elements named `nrepeat` and `nfold`
  - `NA`, `NULL` or `FALSE` to suppress shrinkage tuning.
fit_qda

fit_qda(x, y, ...)  

Arguments

- `x`  
  Dataset, numerical matrix with observations as rows.

- `y`  
  Class labels, factor.

- `...`  
  Sent to `qda`.

Value

Fitted QDA.

Author(s)

Christofer Bäcklin

See Also

`emil`, `predict_pamr`, `importance_pamr`, `modeling_procedure`
References


See Also

emil, predict_qda, modeling_procedure

---

**fit_randomForest**

*Fit random forest.*

Description

Directly calling the `randomForest` package implementation. See `randomForest` for parameter specification.

Usage

```r
fit_randomForest(x, y, ...)
```

Arguments

- `x`: Dataset, numerical matrix with observations as rows.
- `y`: Class labels, factor.
- `...`: Sent to `randomForest`.

Value

Fitted random forest.

Author(s)

Christofer Bäcklin

References


See Also

emil, predict_randomForest, importance_randomForest, modeling_procedure
fit_rpart

Fit a decision tree

Description
Fit a decision tree

Usage
fit_rpart(x, y, ...)

Arguments
x Data set (features).
y Response.
...

Sent to rpart.

Value
A fitted decision tree.

Author(s)
Christofer Bäcklin

References

fit_svm

Fit a support vector machine

Description
Fit a support vector machine

Usage
fit_svm(x, y, probability = TRUE, ranges, ...)

References
get_color

Arguments

x  Dataset, observations as rows.
y  Response vector.
probability  If FALSE support for class probability estimation is not included in the fitted model. This may save some computation time.
ranges  Parameter ranges to tune over. Sent to tune.svm.

Author(s)

Christofer Bäcklin

get_color  

Description

Can be used to modify an existing palette, e.g. change brightness, or to generate a palette for a response vector.

Usage

get_color(x, ...)

## Default S3 method:
get_color(x, s, v, alpha, ...)

## S3 method for class 'factor'
get_color(x, levels = FALSE, col = "Set1", ...)

Arguments

x  Character vector of colors or factor of class memberships to generate colors for.
...  Sent to get_color.default.
s  Saturation. s = 0 leaves it unchanged, 0 < s <= 1 increases, and -1 <= s < 0 decreases.
v  Value. s = 0 leaves it unchanged, 0 < s <= 1 increases, and -1 <= s < 0 decreases.
alpha  Transparency.
levels  If TRUE a palette with one color per level of x is returned. If FALSE one color per element in x is returned.
col  Color palette with one color per class or the name of the color brewer palette to use, see name argument of brewer_pal for a list of possible values.
Value
A character vector of hex colors.

Author(s)
Christofer Bäcklin

get_importance
Feature (variable) importance of a fitted model

Description
Note that different methods calculates feature importance in different ways and that they are not
directly comparable.

Usage
get_importance(object, format, ...)

Arguments

object Fitted model.
format Table format of the output. See http://en.wikipedia.org/wiki/Wide_and_narrow_data for more info.
... Sent on to the procedure’s feature importance scoring function.

Details
When extending the emil framework with your own method, the importance function should return
a data frame where one column is called "feature" and the remaining columns are named after the
classes.

Value
A vector of length p or an p-x-c matrix of feature importance scores where p is the number of
descriptors and c is the number of classes.

Author(s)
Christofer Bäcklin

See Also
emil
get_performance

Examples

```r
procedure <- modeling_procedure("pamr")
model <- fit("pamr", x=iris[-5], y=iris$Species)
get_importance(model)

cv <- resample("crossvalidation", iris$Species, nrepeat=2, nfold=3)
result <- evaluate("pamr", iris[-5], iris$Species, resample=cv,
 .save=c(importance=TRUE))
get_importance(result)
```

---

get_performance Extract prediction performance

Description

Extract prediction performance

Usage

```r
get_performance(result, format = c("wide", "long"))
```

Arguments

- **result**: Modeling result, as returned by `evaluate`.
- **format**: Table format of the output. See [http://en.wikipedia.org/wiki/Wide_and_narrow_data](http://en.wikipedia.org/wiki/Wide_and_narrow_data) for more info.

Value

Data frame.

Author(s)

Christofer Bäcklin

---

get_prediction Extract predictions from modeling results

Description

Extract predictions from modeling results

Usage

```r
get_prediction(result, resample, type = "prediction", format = c("long", "wide"))
```
Arguments

result  Modeling result, as returned by `evaluate` and `evaluate`.
resample  Resampling scheme used to create the results.
type  The type of prediction to return. The possible types vary between modeling procedure.
format  Table format of the output. See [http://en.wikipedia.org/wiki/Wide_and_narrow_data](http://en.wikipedia.org/wiki/Wide_and_narrow_data) for more info.

Value

A data frame where the id column refers to the observations.

Author(s)

Christofer Bäcklin

---

**get_response**

*Extract the response from a data set*

Description

Extract the response from a data set

Usage

`get_response(x, y)`

Arguments

- `x`  Data set features.
- `y`  Response vector or any other type of objects that describe how to extract the response vector from `x`.

Value

A response vector.

Author(s)

Christofer Bäcklin

Examples

```r
identical(iris$Species, get_response(iris, "Species"))
identical(iris$Sepal.Length, get_response(iris, Sepal.Length - .))
```
**get_tuning**

*Extract parameter tuning statistics*

**Description**

Extract parameter tuning statistics

**Usage**

```r
get_tuning(object)
```

**Arguments**

- **object**: Fitted model or modeling procedure

**Value**

A data frame of tuning statistics in long format.

**Author(s)**

Christofer Bäcklin

**Examples**

```r
procedure <- modeling_procedure("randomForest",
    parameter = list(mtry = c(1, 3),
        nodesize = c(4, 10)))
model <- fit(procedure, x=iris[-5], y=iris$Species)
get_tuning(model)

options(emil_max_indent=4)
ho <- resample("holdout", iris$Species, nfold=5)
result <- evaluate(procedure, iris[-5], iris$Species, resample=ho,
    .save=c(model=TRUE))
get_tuning(result)
```

---

**image.resample**

*Visualize resampling scheme*

**Description**

Class specific extension to `image`. 

importance_glmnet

Usage

## S3 method for class 'resample'
image(x, col, ...)

## S3 method for class 'crossvalidation'
image(x, col, ...)

Arguments

x
Resampling scheme, as returned by `resample`.

col
Color palette matching the values of x. Can also be the response vector used to create the scheme for automatic coloring.

...
Sent to `plot`.

Value

Nothing, produces a plot.

Author(s)

Christofer Bäcklin

See Also

`emil`, `resample`

Examples

y <- gl(2, 30)
image(resample("crossvalidation", y, nfold=3, nrepeat=8), col=y)

---

importance_glmnet Feature importance extractor for elastic net models

Description

Feature importance extractor for elastic net models

Usage

importance_glmnet(object, s, ...)

importance_ridge_regression(object, s, ...)

importance_lasso(object, s, ...)
importance_pamr

Arguments

object  Fitted elastic net model, as produced by `fit_glmnet`

s  Regularization parameter lambda.

...  Sent to `predict_glmnet`.

Value

A feature importance data frame.

Author(s)

Christofer Bäcklin

See Also

`emil`, `fit_glmnet`, `predict_glmnet`, `modeling_procedure`

---

importance_pamr  Feature importance of nearest shrunken centroids.

Description

Calculated as the absolute difference between the overall centroid and a class-wise shrunken centroid (which is the same for both classes except sign).

Usage

`importance_pamr(object, threshold, thres_fun = max, ...)`

Arguments

object  Fitted pamr classifier

threshold  Threshold to use for classification. This argument is only needed if you want to override the value set during model fitting.

thres_fun  Threshold selection function. Only needed if you want to override the function set during model fitting.

...  Sent to `pamr.predict`.

Details

In case multiple thresholds give the same error the largest one is chosen (i.e. the one keeping the fewest features).

Value

A data frame of feature importance scores.
importance_randomForest

Feature importance of random forest.

Description

Feature importance of random forest.

Usage

importance_randomForest(object, type = 1, ...)

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>object</td>
<td>Fitted randomForest classifier</td>
</tr>
</tbody>
</table>
| type     | Importance can be assessed in two ways:  
1. Permuted out-of-bag prediction error (default). This can only be used if the classifier was fitted with argument prediction=TRUE which is default.  
2. Total decrease in node impurity.  
...     | Ignored.  |

Value

An prediction vector with elements corresponding to variables.

Author(s)

Christofer Bäcklin

See Also

eamil, fit_pamr, predict_pamr, modeling_procedure
Description

If you want to impute, build model and predict you should use `pre_impute_median` or `pre_impute_knn`. This function imputes using all observations without caring about cross-validation folds.

Usage

```r
impute_knn(x, k = 0.05, distance_matrix = "auto")
impute_median(x)
```

Arguments

- `x`: Dataset.
- `k`: Number of nearest neighbors to use.
- `distance_matrix`: Distance matrix.

Details

For additional information on the parameters see `pre_impute_knn` and `pre_impute`.

Value

An imputed matrix.

Author(s)

Christofer Bäcklin

See Also

`emil, pre_process, pre_impute_knn, pre_impute_median`

Examples

```r
x <- matrix(rnorm(36), 6, 6)
x[sample(length(x), 5)] <- NA
impute_knn(x)
imputeMedian(x)
```
**indent**  
*Increase indentation*

**Description**
Increase indentation

**Usage**
`indent(base, indent)`

**Arguments**
- `base`  
  Base indentation level of the function printing the message.
- `indent`  
  Extra indentation of this message.

**Value**
An integer that can be used to specify the indentation level of messages printed with `log_message`.

**Author(s)**
Christofer Bäcklin

---

**index_fit**  
*Convert a fold to row indexes of fitting or test set*

**Description**
Convert a fold to row indexes of fitting or test set

**Usage**
`index_fit(fold, allow_oversample = TRUE)`
`index_test(fold)`

**Arguments**
- `fold`  
  A fold of a resampling scheme.
- `allow_oversample`  
  Whether or not to allow individual observation to exist in multiple copies in the training set. This is typically not the case, but can be used when a class is underrepresented in the data set.
is_blank

Value
An integer vector of row indexes.

Author(s)
Christofer Bäcklin

See Also
emil, resample

Description
This is mainly an internal function but as other dependent packages also use it sometimes and it generally is quite handy to have it is exported for public use.

Usage
is_blank(x, false_triggers = FALSE)

Arguments
x A variable.
false_triggers Whether FALSE should be considered as empty.

Value
Logical telling if variable is blank.

Author(s)
Christofer Bäcklin

Examples
is_blank(NULL)
is_constant  
Check if an object contains more than one unique value

Description
Check if an object contains more than one unique value

Usage
is_constant(x, na.rm = FALSE)

Arguments
x  Vector or matrix.
na.rm  Whether to ignore missing values.

Value
Logical scalar that is TRUE if x contains more than one unique value and FALSE if not.
A logical scalar that is TRUE if x contains more than one unique value and FALSE otherwise. In case x contains missing values NA is returned if na.rm = FALSE. If there are no non-missing values NA is always returned.

Author(s)
Christofer Bäcklin

is_multi_procedure  
Detect if modeling results contains multiple procedures

Description
Detect if modeling results contains multiple procedures

Usage
is_multi_procedure(result)

Arguments
result  Modeling results, as returned by evaluate.

Value
Logical scalar.
**Description**

This function studies the change in performance as the sizes of the training set is varied. In case the studied modeling procedures cannot produce models on the smallest training sets, please use .return_error=TRUE (see `evaluate`).

**Usage**

```r
learning_curve(procedure, x, y, test_fraction, nfold = 100, ..., .verbose = TRUE)
```

**Arguments**

- `procedure` 
  modeling_procedure.
- `x` 
  Dataset descriptors.
- `y` 
  Response.
- `test_fraction` 
  Fraction of dataset to hold out, i.e. use as test set. Defaults 20 logarithmically distributed values ranging from all but 5 observations per class in the largest test set to only 5 observations per class in the smallest test set.
- `nfold` 
  How many holdout folds that should be calculated.
- `...` 
  Sent to `evaluate`.
- `.verbose` 
  Whether to print an activity log. Set to -1 to also suppress output generated from the procedure's functions.

**Author(s)**

Christofer Bäcklin

**References**


**Examples**

```r
options(emil_max_indent=3)
lc <- learning_curve(c(Linear="lda", Quadratic="qda"),
                    iris[-5], iris$Species, test_fraction=c(.7, .5, .3))
plot(lc)
```
list_method  List all available methods

Description
This function searches all attached packages for methods compatible with the emil framework.

Usage
list_method(pos = search())

Arguments
pos  Location to search in, see ls.

Value
A data frame.

Author(s)
Christofer Bäcklin

Examples
list_method()

log_message  Print a timestamped and indented log message

Description
To suppress messages below a given indentation level set the global option setting emil_max_indent, as in the example below.

Usage
log_message(indent = 1, ..., time = TRUE, domain = "R-emil", appendLF = TRUE)

Arguments
indent  Indentation level. Messages with indent=0 are suppressed.
...  Sent to sprintf.
time  Whether or not to print timestamp.
domain  See message.
appendLF  Whether to finish the message with a linebreak or not.
mode

Author(s)

Christofer Bäcklin

Examples

equipment <- c("flashlight", "snacks", "pick")
{
  log_message(1, "Begin descent")
  log_message(2, "Oh no, forgot the %s!", sample(equipment, 1))
  log_message(2, "Hello? Can you throw it down to me?", time=FALSE)
  log_message(1, "Aw shucks, I'm coming back up.")
}

for(verbose in c(TRUE, FALSE)){
  cat("It's", verbose, "\n")
  for(i in 0:3)
    log_message(indent(verbose, i), "Down")
}

options(emil_max_indent = 2)
for(i in 1:3)
  log_message(i, "Down")

mode Get the most common value

Description

Get the most common value

Usage

mode(x, na.rm = FALSE, allow_multiple = TRUE)

Arguments

x Vector.

na.rm Whether to ignore missing values when calculating the mode. Note that modes may be identified even if x contains missing values as long as they are too few to affect the result.

allow_multiple Controls what is returned if x contains more than one mode. If TRUE all modes are returned, if FALSE NA is returned.

Value

The most common values or values in x or NA if could not be determined.
### Description

A modeling procedure is an object containing all information necessary to carry out and evaluate the performance of a predictive modeling task with `fit`, `tune`, or `evaluate`. To use an out-of-the-box algorithm with default values, only the `method` argument needs to be set. See `emil` for a list of available methods. To deviate from the defaults, e.g. by tuning parameters or using a custom function for model fitting, set the appropriate parameters as described below. For a guide on how to implement a custom method see the documentation page `extension`.

### Usage

```r
modeling_procedure(method, parameter = list(), error_fun = NULL, fit_fun, predict_fun, importance_fun)
```

### Arguments

- **method**: The name of the modeling method. Only needed to identify plug-in functions, i.e. if you supply them yourself there is no need to set method.

- **parameter**: A list of model parameters. These will be fed to the fitting function after the dataset (x and y parameters). To tune a parameter, supply the candidate values in a vector or list.

  When tuning more than one parameter, all combinations of parameter values will be tested, if the elements of parameter are named. To manually specify
modeling_procedure

which parameter value combinations to try, leave the the elements unnamed (see example 3 and 4).
Parameters that should have vectors or lists as values, e.g. trControl when using fit_caret to train pkgcaret models, must be wrapped in an additional list. That is, to set a parameter value to a list, but not tune it, make it a list of length 1 containing the list to be used (see example 6).

\*error_fun\* Performance measure used to evaluate procedures and to tune parameters. See \*error_fun\* for details.

\*fit_fun\* The function to be used for model fitting.

\*predict_fun\* The function to be used for model prediction.

\*importance_fun\* The function to be used for calculating or extracting feature importances. See \*get_importance\* for details.

\*Value\*

An object of class modeling_procedure.

\*Author(s)\*

Christofer Bäcklin

\*See Also\*

emil, evaluate, fit, tune, predict, get_importance

\*Examples\*

\# 1: Fit linear discriminants without tuning any parameter,
\# since it has none
modeling_procedure("lda")

\# 2: Tune random forest’s `mtry` parameter, with 3 possible values
modeling_procedure("randomForest", list(mtry = list(100, 250, 1000)))

\# 3: Tune random forest’s `mtry` and `maxnodes` parameters simultaneously,
\# with 3 values each, testing all 9 possible combinations
modeling_procedure("randomForest", list(mtry = list(100, 250, 1000),
maxnodes = list(5, 10, 25)))

\# 4: Tune random forest’s `mtry` and `maxnodes` parameters simultaneously,
\# but only test 3 manually specified combinations of the two
modeling_procedure("randomForest", list(list(mtry = 100, maxnodes = 5),
list(mtry = 250, maxnodes = 10),
list(mtry = 1000, maxnodes = 25)))

\# 5: Tune elastic net’s `alpha` and `lambda` parameters. Since elastic net’s
\# fitting function can tune `lambda` internally in a more efficient way
\# than the general framework is able to do, only tune `alpha` and pass all
\# `lambda` values as a single argument.
modeling_procedure("glmnet", list(alpha = seq(0, 1, length.out=6),
lambda = list(seq(0, 5, length.out=30)))

# 6: Train elastic nets using the caret package's model fitting framework
if(requireNamespace("caret", quietly = TRUE)){
  modeling_procedure("caret", list(method = "glmnet",
    trControl = list(trainControl(verboseIter = TRUE, classProbs = TRUE))));
}

---

**name_procedure**

*Get names for modeling procedures*

**Description**

Get names for modeling procedures

**Usage**

name_procedure(procedure)

**Arguments**

- **procedure**: List of modeling procedures.

**Value**

A character vector of suitable non-duplicate names.

**Author(s)**

Christofer Bäcklin

---

**na_index**

*Support function for identifying missing values*

**Description**

Support function for identifying missing values

**Usage**

na_index(data)

**Arguments**

- **data**: Fitting and testing data sets, as returned by `pre_split`. 
**neg_gmpa**

### Value

Data frame containing row and column indices of missing values or `NULL` if the data doesn’t contain any.

### Author(s)

Christofer Bäcklin

### Examples

```r
x <- as.matrix(iris[-5])
y <- iris$Species
x[sample(length(x), 10)] <- NA
cv <- resample("crossvalidation", y)
sets <- pre_split(x, y, cv[[1]])
sets <- pre_remove(sets, 3L)
na_index(sets)
```

---

**negative_gmpa**

<table>
<thead>
<tr>
<th>negative_gmpa</th>
<th>Negative geometric mean of class specific predictive accuracy</th>
</tr>
</thead>
</table>

### Description

When dealing with imbalanced classification problem, i.e. where the class sizes are very different, small classes tend to be overlooked when tuning parameters by optimizing error rate. Blagus and Lusa (2013) suggested to remedy the problem by using this performance measure instead.

### Usage

```r
neg_gmpa(truth, prediction, na.rm = FALSE)
```

### Arguments

- **truth**
  - See `error_fun`.
- **prediction**
  - See `error_fun`.
- **na.rm**
  - Whether to remove missing values or not.

### Value

A numeric scalar.

### Author(s)

Christofer Bäcklin
References

Blagus, R., & Lusa, L. (2013). *Improved shrunken centroid classifiers for high-dimensional class-imbalanced data*. BMC bioinformatics, 14, 64. doi:10.1186/1471-2105-14-64

See Also

error_fun

---

nice_axis

*Plots an axis the way an axis should be plotted.*

Description

Plots an axis the way an axis should be plotted.

Usage

```
nice_axis(..., las = 1, lwd = 0, lwd.ticks = par("lwd"), lend = 2)
```

Arguments

- `...` Sent to `axis`.
- `las` Rotation of axis labels. Always horizontal by default.
- `lwd` Width of the line drawn along the plot area. Omitted by default since it overlaps with `box` and causes it to look thicker where the axis is.
- `lwd.ticks` Width of the tick lines. These are kept by default.
- `lend` Line endings, see `par`.

Author(s)

Christofer Bäcklin

---

nice_box

*Plots a box around a plot*

Description

Plots a box around a plot

Usage

```
nice_box(lend = 2, ljoin = 1, ...)
```
nice_require

Arguments

lend Line ending style, see par. Defaults to square.
ljoin Line joint style, see par. Defaults to mitre, i.e. 90 degree corners in this case.
... Sent to box.

Author(s)

Christofer Bäcklin

---

nice_require Load a package and offer to install if missing

Description

If running R in interactive mode, the user is prompted for installing missing packages. If running in batch mode an error is thrown.

Usage

nice_require(pkg, reason)

Arguments

pkg Package name.
reason A status message that informs the user why the package is needed.

Value

Nothing

Author(s)

Christofer Bäcklin

Examples

nice_require("base", "is required to do anything at all")
notify_once  

Print a warning message if not printed earlier

Description
To avoid flooding the user with identical warning messages, this function keeps track of which have already been shown.

Usage
```r
notify_once(id, ..., fun = log_message)
reset_notification(id, if_top_level = TRUE)
```

Arguments
- `id`  Warning message id. This is used internally to refer to the message.
- `...` Sent to `warning`
- `fun` Function to display the notification with. Typical choices are `message` or `warning`.
- `if_top_level` If `TRUE` the notifications will only be reset if `reset_notification` was called from a top-level function call. This behaviour prevents the notifications from being reset multiple times during nested calls to functions such as `fit` and `evaluate`.

Author(s)
Christofer Bäcklin

plot.learning_curve  

Plot results from learning curve analysis

Description
Plot results from learning curve analysis

Usage
```r
## S3 method for class 'learning_curve'
plot(x, ..., summaries = list(mean = mean,
   '95-percentile' = function(x) quantile(x, 0.95)))
```
**plot_Surv**

Arguments

- **x**
  
  Results from `learning_curve`.

- **...**
  
  Ignored, kept for S3 consistency.

- **summaries**
  
  Named list of summary functions that can reduce a vector of performance estimates to a single quantity.

Value

A `ggplot` object.

Author(s)

Christofer Bäcklin

---

**plot_Surv** *Plot Surv vector [DEPRECATED]*

Description

Package `survival` includes an official plot function as of version 2.42-5. This will therefore be removed in the next major update.

Usage

```r
plot_Surv(x, y, segments = TRUE, flip = FALSE, legendpos = "topright",
          ...)```

Arguments

- **x**
  
  `Surv` vector.

- **y**
  
  Y-values.

- **segments**
  
  Whether to draw horizontal segments.

- **flip**
  
  Flip the plot to show time on y.

- **legendpos**
  
  Position of legend, see `legend`. Set to NA or NULL to suppress legend.

- **...**
  
  Sent to `plot`.

Author(s)

Christofer Bäcklin
predict.model

Predict the response of unknown observations

Description

Predict the response of unknown observations

Usage

```r
## S3 method for class 'model'
predict(object, x, ..., .verbose = FALSE)
```

Arguments

- `object`: Fitted model.
- `x`: Data set with observations whose response is to be predicted.
- `...`: Sent to the procedure’s prediction function.
- `.verbose`: Whether to print an activity log.

Value

See the documentation of procedure’s method.

Author(s)

Christofer Bäcklin

See Also

- `emil`, `modeling_procedure`, `evaluate`, `fit`, `tune`, `get_importance`

Examples

```r
mod <- fit("lda", x=iris[-5], y=iris$Species)
prediction <- predict(mod, iris[-5])
```
**predict_caret**  
*Predict using a caret method*

**Description**

This is not guaranteed to work with all caret methods. If it doesn’t work for a particular method, the user will need to rewrite it.

**Usage**

`predict_caret(object, x, ...)`

**Arguments**

- `object`  
  Fitted caret model.
- `x`  
  New data to predict the response of.
- `...`  
  Sent to `predict` that forwards it to the appropriate predict function in the caret package.

**Author(s)**

Christofer Bäcklin

**predict_cforest**  
*Predict with conditional inference forest*

**Description**

Prediction function for models fitted with `fit_cforest`.

**Usage**

`predict_cforest(object, x, at, ...)`

**Arguments**

- `object`  
  Fitted cforest classifier, as returned by `fit_cforest`.
- `x`  
  New data to be used for predictions.
- `at`  
  Time point to evaluate survival curves at. If omitted it is set to the last observed time point.
- `...`  
  Sent to `treeresponse`.

**Value**

The predicted chance of survival.
**Author(s)**

Christofer Bäcklin

**See Also**

emil, fit_cforest, modeling_procedure

---

**predict_coxph**

*Predict using Cox proportional hazards model*

**Description**

Predict using Cox proportional hazards model

**Usage**

predict_coxph(object, x, ...)

**Arguments**

- **object**: Fitted model, as returned by `fit_coxph`.
- **x**: Observations whose response is to be predicted.
- **...**: Sent to `predict.coxph`.

**Author(s)**

Christofer Bäcklin

**See Also**

fit_coxph

---

**predict_glmnet**

*Predict using generalized linear model with elastic net regularization*

**Description**

Due to the way `glmnet` is implemented, the regularization alpha can not be modified after the model is fitted.

**Usage**

predict_glmnet(object, x, s, ...)

predict_ridge_regression(object, x, s, ...)

predict_lasso(object, x, s, ...)


**predict_lda**

**Arguments**

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>object</td>
<td>Fitted model.</td>
</tr>
<tr>
<td>x</td>
<td>New data to be predicted.</td>
</tr>
<tr>
<td>s</td>
<td>Regularization parameter lambda.</td>
</tr>
<tr>
<td>...</td>
<td>Sent to <code>predict.glmnet</code>.</td>
</tr>
</tbody>
</table>

**Value**

A list with a subset of the following elements:

- **prediction** The response of the modeling problem, i.e. a factor for classification, problems, a numeric for regressions, and a relative risk for survival analyses.
- **probability** Data frame of predicted class probabilities.
- **link** Link function values.

**Author(s)**

Christofer Bäcklin

**See Also**

`emil`, `fit_glmnet`, `importance_glmnet`, `modeling_procedure`

---

**predict_lda**

*Prediction using already trained prediction model*

**Description**

Wrapper for the **MASS** package implementation.

**Usage**

`predict_lda(object, x, ...)`

**Arguments**

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>object</td>
<td>Fitted classifier as produced by <code>evaluate</code>.</td>
</tr>
<tr>
<td>x</td>
<td>Dataset of observations to be classified.</td>
</tr>
<tr>
<td>...</td>
<td>Sent to <code>predict.lda</code>.</td>
</tr>
</tbody>
</table>

**Value**

A list with elements:

- **prediction**: Factor of predicted class memberships.
- **probability**: Data frame of predicted class probabilities.
predict_lm

Prediction using linear model

Description

Prediction using linear model

Usage

predict_lm(object, x, ...)

Arguments

object Fitted classifier produced by fit_lm.
x Dataset to be predicted upon.
... Sent to predict_lm

Value

A list with elements:

• prediction: Vector of predicted response.

Author(s)

Christofer Bäcklin

See Also

emil, fit_lda, modeling_procedure
**predict_naive_bayes**  
*Predict using naive Bayes model*

**Description**

Predict using naive Bayes model

**Usage**

`predict_naive_bayes(object, x)`

**Arguments**

- `object`  
  Fitted naive Bayes model.

- `x`  
  Data set to predict response for

**Author(s)**

Christofer Bäcklin

**See Also**

`predict`

---

**predict_pamr**  
*Prediction using nearest shrunken centroids.*

**Description**

In case multiple thresholds give the same error the largest one is chosen (i.e. the one keeping the fewest features).

**Usage**

`predict_pamr(object, x, threshold, thres_fun, ...)`

**Arguments**

- `object`  
  Fitted classifier.

- `x`  
  Dataset of observations to be classified.

- `threshold`  
  Threshold to use for classification. This argument is only needed if you want to override the value set during model fitting.

- `thres_fun`  
  Threshold selection function. Only needed if you want to override the function set during model fitting.

- `...`  
  Sent to `pamr.predict`. 
Value

A list with elements:

- prediction: Factor of predicted class memberships.
- probability: Data frame of predicted class probabilities.

Author(s)

Christofer Bäcklin

See Also

emil, fit_pamr, importance_pamr, modeling_procedure

Description

Wrapper for the MASS package implementation.

Usage

predict_qda(object, x, ...)

Arguments

object Fitted classifier as produced by `evaluate`.

x Dataset of observations to be classified.

... Sent to `predict.qda`.

Value

A list with elements:

- prediction: Factor of predicted class memberships.
- probability: Data frame of predicted class probabilities.

Author(s)

Christofer Bäcklin

See Also

emil, fit_qda, modeling_procedure
predict_randomForest

Description
Prediction using random forest.

Usage
predict_randomForest(object, x, ...)

Arguments
- object: Fitted model.
- x: Dataset of observations to be classified.
- ...: Ignored

Value
When used for classification, a list with elements:
- prediction: Factor of predicted class memberships.
- probability: Data frame of predicted class probabilities.

When used for regression, a list with the element:
- prediction: Vector of predicted response.

Author(s)
Christofer Bäcklin

See Also
emil, fit_randomForest, importance_randomForest, modeling_procedure
predict_rpart
_________

*Predict using a fitted decision tree*

**Description**

Predict using a fitted decision tree

**Usage**

```r
predict_rpart(object, x)
```

**Arguments**

- `object` Fitted decision tree.
- `x` New data whose response is to be predicted.

**Value**

Predictions. The exact form depends on the type of application (classification or regression)

**Author(s)**

Christofer Bäcklin

---

predict_svm
_________

*Predict using support vector machine*

**Description**

Predict using support vector machine

**Usage**

```r
predict_svm(object, x, probability = object$compprob, statistic = TRUE, ...)
```

**Arguments**

- `object` Fitted SVM, as produced by `fit_svm`.
- `x` Data set to predict response for.
- `probability` Whether to calculate class probabilities.
- `statistic` Whether to return the raw classification statistics.
- `...` Sent to `predict`.

**Author(s)**

Christofer Bäcklin
**pre_factor_to_logical**  
*Convert factors to logical columns*

**Description**

Factors will be converted to one logical column per level (or one fewer if a base level is specified).

**Usage**

```r
pre_factor_to_logical(data, feature, base = 1L, drop = TRUE)
```

**Arguments**

- `data` Pre-processed data set, as produced by `pre_split`.
- `feature` Character vector with names of features to convert. Defaults to all factors in the data set.
- `base` Sent to `factor_to_logical`. To specify different bases for different columns supply a vector or list with named elements.
- `drop` Sent to `factor_to_logical`. To specify different bases for different columns supply a vector or list with named elements.

**Author(s)**

Christofer Bäcklin

**Examples**

```r
x <- mtcars[-1]
x <- transform(x,
    cyl = factor(cyl, ordered=TRUE),
    vs = factor(vs),
    gear = factor(gear)
)
y <- mtcars$mpg
cv <- resample("crossvalidation", y)
data <- pre_split(x, y, cv[[1]]) %>%
    pre_factor_to_logical(base = c(cyl="4", vs="0"),
                           drop=c(cyl=FALSE, gear=FALSE))
data$fit$x
```
**pre_impute_df**

---

**pre_impute**  
*Basic imputation*

---

**Description**

This solution is optimized for the scenario that the dataset is very large but only contains missing values in a small number of columns.

**Usage**

```r
pre_impute(data, fun, ...)
```

```r
pre_impute_median(data)
```

```r
pre_impute_mean(data)
```

**Arguments**

- `data`: Fitting and test datasets, as returned by `pre_split` or any other standard pre-processing function.
- `fun`: Function for calculating imputation values. Should take a vector and return a scalar.
- `...`: Sent to `fun`.

**Value**

A pair of fitting and testing datasets.

**Author(s)**

Christofer Bäcklin

---

**pre_impute_df**  
*Impute a data frame*

---

**Description**

This function imputes each column of data frames univariately with different functions depending on their class.

**Usage**

```r
pre_impute_df(data, class_fun = list(numeric = function(x) median(x, na.rm = TRUE), integer = function(x) median(x, na.rm = TRUE)),
               default_fun = function(x) mode(x, na.rm = TRUE, allow_multiple = FALSE))
```
Arguments

- **data**: Pre-processed data set with features in a data frame.
- **class_fun**: List of functions to use for imputating specific feature classes.
- **default_fun**: Function to use for imputation features of classes not listed in class_fun.
- **na.rm**: Whether to remove missing values.

Author(s)

Christofer Bäcklin

Examples

```r
x <- iris
x[sample(150, 3), 1] <- NA
x[sample(150, 1), 3] <- NA
x[sample(150, 5), 5] <- NA
y <- gl(2, 75)
fold <- resample("holdout", y, nfold=1)[[1]]
data <- pre_split(x, y, fold) %>%
    pre_impute_df
```

Description

Nearest neighbor methods need to have a distance matrix of the dataset it works on. When doing repeated model fittings on subsets of the entire dataset it is unnecessary to recalculate it every time, therefore this function requires the user to manually calculate it prior to resampling and supply it in a wrapper function.

Usage

```r
pre_impute_knn(data, k = 0.05, distance_matrix)
```

Arguments

- **data**: Fitting and testing data sets, as returned by `pre_split`.
- **k**: Number of nearest neighbors to calculate mean from. Set to < 1 to specify a fraction.
- **distance_matrix**: A matrix, `dist` object or "auto". Notice that "auto" will recalculate the distance matrix in each fold, which is only meaningful in case the features of `x` vary between folds. Otherwise you are just wasting time.
Details

Features with fewer than \( k \) non-missing values will be removed automatically.

Author(s)

Christofer Bäcklin

Examples

```r
x <- iris[-5]
x[sample(nrow(x), 30), 3] <- NA
my.dist <- dist(x)
evaluate(modeling_procedure("lda"), x = x, y = iris$Species,
  pre_process = function(...){
    pre_split(...) \%\% pre_impute_knn(k = 4, distance_matrix = my.dist)
  }
)
```

---

**pre_log_message**

Print log message during pre-processing

Description

Print log message during pre-processing

Usage

```r
pre_log_message(data, ...)
```

Arguments

- **data**
  - Pre-processed data set.
- **...**
  - Sent to `log_message`

Value

The same data set as inputted. This only purpose of this function is to print a log message as a side effect.

Author(s)

Christofer Bäcklin

See Also

`pre_process`, `log_message`.  

---
pre_pamr  

PAMR adapted dataset pre-processing

Description

The predict framework is designed to work with dataset where rows correspond to observations and columns to descriptors. PAMR wants it the other way, and also to have the fitting set response vector supplied in a list with the descriptors. This function applies a standard pre-processing function and then reformats the result to satisfy PAMR.

Usage

pre_pamr(data)

Arguments

data  
Fitting and testing data sets, as returned by pre_split.

Details

pre_pamr must be run last if chained with other pre-processing functions, since it substantially reshapes the data.

Value

A list with fitting and testing sets, formatted the way pamr wants them.

Author(s)

Christofer Bäcklin

References


See Also

emil, pre_process
**Description**

These functions are run in `evaluate` just prior to model fitting, to extract fitting and test sets from the entire dataset and apply transformations to pre-process the data (for handling missing values, scaling, compression etc.). They can also be used to adapt the form of the data to a specific fitting function, e.g. `pre_pamr` that transposes the dataset to make it compatible with the `pamr` classification method.

**Usage**

```r
pre_split(x, y, fold)

pre_convert(data, x_fun, y_fun, ...)

pre_transpose(data)

pre_remove(data, feature)

pre_center(data, y = FALSE, na.rm = TRUE)

pre_scale(data, y = FALSE, na.rm = TRUE, center = TRUE)

pre_remove_constant(data, na.rm = TRUE)

pre_remove_correlated(data, cutoff)

pre_pca(data, ncomponent, scale. = TRUE, ...)
```

**Arguments**

- `x` Dataset.
- `y` Response vector.
- `fold` A logical or numeric vector with TRUE or positive numbers for fitting observations, FALSE or 0 for test observations, and NA for observations not to be included.
- `data` Fitting and testing data sets, as returned by `pre_split`.
- `x_fun` Function to apply to the descriptors of the datasets (e.g. `x`). This function will be applied independently to the fitting and testing sets.
- `y_fun` Function to be applied to the response of the training and test sets (independently).
- `...` Sent to internal methods, see the code of each function.
- `feature` The features to be removed. Can be integer, logical or character.
pre_process

- `na.rm` A logical value indicating whether NA values should be ignored.
- `center` Whether to center the data before scaling.
- `cutoff` See `findCorrelation`.
- `ncomponent` Number of PCA components to use. Missing all components are used.
- `scale` Sent to `prcomp`.

**Details**

When supplied to `evaluate`, pre-processing functions can be chained (i.e. executed sequentially) after an initiating call to `pre_split`. This can either be done using the *pipe operator* defined in the `magrittr` package or by putting all pre-processing functions in a regular list (see the examples).

Note that all transformations are defined based on the fitting data only and then applied to both fitting set and test set. It is important to not let the test data in any way be part of the model fitting, including the preprocessing, to not risk information leakage and biased results!

The imputation functions can also be used outside of `evaluate` by not supplying a fold to `pre_split`. See the code of `impute_median` for an example.

**Value**

A list with the following components

- `fit` Fitting set.
- `test` Test set.
- `feature_selection` Integer vector mapping the features of the training and test sets to the original data sets.
- `fold` The fold that was used to split the data.

**Author(s)**

Christofer Bäcklin

**See Also**

`pre_factor_to_logical`, `emil`, `pre_impute_knn`

**Examples**

```r
# Setup an example to work on
x <- as.matrix(iris[-5])
x[sample(600, 6)] <- NA
y <- iris$Species
cv <- resample("crossvalidation", y, nrepeat=3, nfold=4)
procedure <- modeling_procedure("lda")

# Simple dataset splitting
sets <- pre_split(x, y, cv[[1]])

# Chaining using the pipe operator
```
sets <- pre_split(x, y, cv[[1]])  # Integration with 'evaluate'
result <- evaluate(procedure, x, y, resample=cv,
                   pre_process = function(...){
                        pre_split(...)  
                        pre_impute_median  
                        pre_scale
                   }
)

# or analogously with a list
result <- evaluate(procedure, x, y, resample=cv,
                   pre_process = list(pre_split, pre_impute_median, pre_scale))

# Imputing without splitting
x.imputed <- impute_knn(x)

# Using a whole chain without splitting
x.processed <- pre_split(x, y=NULL)  
             pre_impute_median  
             pre_scale
             (function(data) data$fit$x)

print.preprocessed_data

Print method for pre-processed data

Description

Print method for pre-processed data

Usage

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

Arguments

x  Pre-processed data, as produced by pre_split.
...
Ignored, kept for S3 consistency.

Value

Nothing
Author(s)

Christofer Bäcklin

pvalue  

*Extraction of p-value from a statistical test*

Description

These calculations are written in such a way that they avoid rounding off errors that plague the survival and cmpsk packages.

Usage

\[
pvalue(x, \ \text{log}_p = \text{FALSE}, \ \ldots)
\]

Arguments

- **x**: Test, i.e. a fitted object of a supported type.
- **log_p**: Whether to return the logarithm of the p-value.
- **\ldots**: Sent to class method.

Value

p-value.

Author(s)

Christofer Bäcklin

See Also

- pvalue.crr, pvalue.survdiff, pvalue.cuminc

pvalue.coxph  

*Extract p-value from a Cox proportional hazards model*

Description

Based on summary.coxph.

Usage

```r
# S3 method for class 'coxph'
pvalue(x, \ \text{log}_p = \text{FALSE}, test = c("logrank", "wald", "likelihood"), \ldots)
```
Arguments

- **x**: Fitted coxph model.
- **log_p**: Whether to return the logarithm of the p-value.
- **test**: What test to calculate. "likelihood" is short for means likelihood ratio test.
- **...**: Ignored. Kept for S3 consistency.

Value

- p-value.

Author(s)

Christofer Bäcklin

References


See Also

- pvalue

---

**pvalue.crr**

*Extracts p-value from a competing risk model*

Description

Extracts p-value from a competing risk model

Usage

```r
## S3 method for class 'crr'
pvalue(x, log_p = FALSE, ...)
```

Arguments

- **x**: Fitted crr model, as returned by crr.
- **log_p**: Whether to return the logarithm of the p-value.
- **...**: Ignored. Kept for S3 consistency.

Value

- Two-sided p-value.
pvalue.cuminc

Author(s)
Christofer Bäcklin

See Also
pvalue

Examples
if(requireNamespace("cmprsk", quietly = TRUE)){
    time <- 1:20
    event <- c(rep(0, 9), rep(2, 3), rep(1, 8))
    data <- rep(0:1, each=10)
    x <- cmprsk::crr(time, event, data)
    
    # Compare p-values of implementations
    print(x)
    pvalue(x)
}

pvalue.cuminc  Extract p-value from a cumulative incidence estimation

Description
This is also known as Gray’s test.

Usage
## S3 method for class 'cuminc'
pvalue(x, log_p = FALSE, ...)

Arguments

x  Fitted cuminc estimate.
log_p  Whether to return the logarithm of the p-value.
...  Ignored. Kept for S3 consistency.

Value
p-value.

Author(s)
Christofer Bäcklin
See Also

pvalue

pvalue.survdiff

Extracts p-value from a logrank test

Description

Extracts p-value from a logrank test

Usage

## S3 method for class 'survdiff'
pvalue(x, log_p = FALSE, ...)

Arguments

x
  Logrank test result, as returned by survdiff.

log_p
  Whether to return the logarithm of the p-value.

...
  Ignored. Kept for S3 consistency.

Value

p-value. if(requireNamespace("survival", quietly = TRUE))

y <- survival::Surv(time=1:100, event=rep(1:0, each=50)) groups <- rep(1:2, each=50) x <- survival::survdiff(y ~ groups)

# Compare p-values of implementations print(x) pvalue(x)

Author(s)

Christofer Bäcklin

See Also

pvalue
Description

Performance evaluation and parameter tuning use resampling methods to estimate the performance of models. These are defined by resampling schemes, which are data frames where each column corresponds to a division of the data set into mutually exclusive training and test sets. Repeated hold out and cross-validation are two methods to create such schemes.

Usage

resample(method, y, ..., subset = TRUE)

resample_holdout(y, test_fraction = 0.5, nfold = 5,
    balanced = is.factor(y), subset)

resample_crossvalidation(y, nfold = 5, nrepeat = 5,
    balanced = is.factor(y), subset)

resample_bootstrap(y, nfold = 10, fit_fraction = if (replace) 1 else 0.632,
    replace = TRUE, balanced = is.factor(y), subset)

Arguments

method The resampling method to use, e.g. "holdout" or "crossvalidation".
y Observations to be divided.
... Sent to the method specific function, e.g. "resample_holdout".
subset Which objects in y that are to be divided and which that are not to be part of neither set. If subset is a resampling scheme, a list of inner cross-validation schemes will be returned.
test_fraction Fraction of objects to hold out (0 < test_fraction < 1).
nfold Number of folds.
balanced Whether the sets should be balanced or not, i.e. if the class ratio over the sets should be kept constant (as far as possible).
nrepeat Number of fold sets to generate.
fit_fraction The size of the training set relative to the entire data set.
replace Whether to sample with replacement.

Details

Note that when setting up analyzes, the user should not call resample_holdout or resample_crossvalidation directly, as resample performs additional necessary processing of the scheme.

Resampling scheme can be visualized in a human digestible form with the image function.
Functions for generating custom resampling schemes should be implemented as follows and then
called by `resample("myMethod", ...):

```r
resample_myMethod <- function(y, ..., subset)
```

- `y` Response vector.
- `...` Method specific attributes.
- `subset` Indexes of observations to be excluded for the resampling.

The function should return a list of the following elements:

- `folds` A data frame with the folds of the scheme that conforms to the description in the 'Value'
  section below.
- `parameter` A list with the parameters necessary to generate such a resampling scheme. These are
  needed when creating subschemes needed for parameter tuning, see `subresample`.

**Value**

A data frame defining a resampling scheme. TRUE or a positive integer codes for training set and
FALSE or 0 codes for test set. Positive integers > 1 code for multiple copies of an observation in the
training set. NA codes for neither training nor test set and is used to exclude observations from the
analysis altogether.

**Author(s)**

Christofer Bäcklin

**See Also**

`emil, subresample, image.resample, index_fit`

**Examples**

```r
resample("holdout", 1:50, test_fraction=1/3)
resample("holdout", factor(runif(60) >= .5))
y <- factor(runif(60) >= .5)
cv <- resample("crossvalidation", y)
image(cv, main="Cross-validation scheme")
```

---

**roc_curve**

*Calculate ROC curves*

**Description**

Calculate ROC curves
roc_curve

Usage

roc_curve(result, y, resample, class = levels(y), statistic = "probability")

## S3 method for class 'roc_curve'
as.data.table(x, ...)

## S3 method for class 'roc_curve'
as.data.frame(x, ...)

## S3 method for class 'roc_curve'
plot(x, ...)

Arguments

result  Modeling results, as returned by evaluate.
y      True response vector used to create result.
resample Resampling scheme used to create result.
class  The class of interest to create ROC-curves for.
statistic The name of the statistic (as returned by the prediction function of the modeling procedure).
x  Roc curve object, as returned by roc_curve.
... Sent to as.data.frame or as.data.table.

Value

A data frame of class "roc".

Author(s)

Christofer Bäcklin

Examples

# Generate some noisy data
my.data <- iris

# Train and evaluate some classifiers
procedure <- list(lda = modeling_procedure("lda"),
                 qda = modeling_procedure("qda"))
cv <- resample("crossvalidation", iris$Species, nrep=1, nfold=3)
result <- evaluate(procedure, my.data[-5], my.data$Species, resample=cv)

# Study the performance
select(result, fold=TRUE, method=TRUE, error="error")
roc <- roc_curve(result, my.data$Species, cv)
plot(roc)
select  

emil and dplyr integration

Description

Modeling results can be converted to tabular format and manipulated using dplyr and other Hadley-verse packages. This is accomplished by a class specific `select_` function that differs somewhat in syntax from the default `select_`.

Usage

```r
## S3 method for class 'list'
select_(.data, ..., .dots)
```

Arguments

- `.data` Modeling results, as returned by `evaluate`.
- `...` Not used, kept for consistency with `dplyr`.
- `.dots` Indices to select on each level of `.data`, i.e. the first index specifies which top level elements of `.data` to select, the second specifies second-level-elements etc. The last index must select elements that can be converted to a data frame. In case the desired bottom-level element is related to the observations of a modeling task, e.g. the predictions of a test set, you must supply the resampling scheme used to produce `.data` at the appropriate level (see the examples).

The names of the `...` arguments specifies the names of the resulting data frame. Non-named arguments will be used to traverse the data but not returned.

In summary the `...` indices can be on the following forms:

- **Simple indices** Anything that can be used to subset objects, e.g. integers, logicals, or characters.
- **Functions** A function that produces a data frame, vector or factor.
- **Resampling schemes** The same resampling scheme that was used to produce the modeling results.

Value

A `data.frame` in long format.

Author(s)

Christofer Bäcklin

See Also

subtree
Examples

# Produce some results
x <- iris[-5]
y <- iris$Species
names(y) <- sprintf("orchid\%03i", seq_along(y))
cv <- resample("crossvalidation", y, nfold=3, nrepeat=2)
procedures <- list(nsc = modeling_procedure("pamr"),
                   rf = modeling_procedure("randomForest"))
result <- evaluate(procedures, x, y, resample=cv)

# Get the foldwise error for the NSC method
result %>% select(fold = TRUE, "nsc", error = "error")

# Compare both methods
require(tidyverse)
result %>%
  select(fold = TRUE, method = TRUE, error = "error") %>%
  spread(method, error)
require(dplyr)
result %>%
  select(fold = TRUE, method = TRUE, error = "error") %>%
  group_by(method) %>%
  summarize(mean_error = mean(error))

# Investigate the variability in estimated class 2 probability across folds
result %>%
  select(fold = cv, "nsc", "prediction", probability = function(x) x$probability[,2]) %>%
  spread(fold, probability)

---

subresample Generate resampling subschemes

Description

A subscheme is a resampling scheme that only includes observations in the training set of a fold. This function automatically fetches the type and parameters of the prototype fold and use them to generate the subscheme.

Usage

subresample(fold, y)

Arguments

fold
A resampling scheme or fold to use to define the sub scheme(s).
y
The observations used to create the resampling scheme. See resample for details.
Value
A resampling scheme.

Author(s)
Christofer Bäcklin

See Also
emil, resample

Examples
```r
cv <- resample("holdout", y=1:12, test_fraction=1/4, nfold=3)
inner.cv <- subresample(cv, y=1:12)
```

subtree
Extract a subset of a tree of nested lists

Description
Modeling results produced by `evaluate` comes in the form of nested lists. This function can be used to subset or rearrange parts of the results into vectors, matrices or data frames. Also note the `select` function that provides an extension to the `dplyr` package for data manipulation.

Usage
```r
subtree(x, i, ..., error_value, warn, simplify = TRUE)
```

Arguments
- `x` List of lists.
- `i` Indexes to extract on the first level of the tree. Can also be a function that will be applied to the downstream result of the function.
- `...` Indexes to extract on subsequent levels.
- `error_value` A template for the return value in case it is missing or invalid. Note that `NA` is a `logical` by default, causing `subtree` to also convert existing results to logicals. To get around this, please specify it as `as.numeric(NA)`, `as.character(NA)`, or similar (see the example below).
- `warn` Specifies whether warnings should be displayed (0), ignored (-1), or break execution (1). Works like the `options` parameter `warn`.
- `simplify` Whether to collapse results into vectors or matrices when possible (TRUE) or to preserve the original tree structure as a list (FALSE).

Details
This function can only be used to extract data, not to assign.
Value
A subset of the list tree.

Author(s)
Christofer Bäcklin

See Also
select, get_prediction, get_importance, get_tuning.

Examples
l <- list(A=list(a=0:2, b=3:4, c=023-22030),
         B=list(a=5:7, b=8:9))
subtree(l, 1:2, "b")
subtree(l, TRUE, mean, "a")

# More practical examples
x <- iris[-5]
y <- iris$Species
cv <- resample("crossvalidation", y, nfold=5, nrep=3)
procedure <- modeling_procedure("pamr")

# To illustrate the error handling capacities of subtree we'll introduce some
# spurious errors in the pre-processing function. By setting .return_error=TRUE
# they won't break the execution, but will instead be return in the results.
pre_error <- function(data, risk=1){
  if(runif(1) < risk)
    stop("Oh no! Random error!")
data
}
result <- evaluate(procedure, x, y, resample=cv,
                   .save=(importance=TRUE), .return_error=TRUE,
                   pre_process = function(...)(
                     pre_split(...) %>%
                     pre_error(risk=.3) %>%
                     pre_pamr
                   )
)
message(sum(sapply(result, inherits, "error"),
           "folds did not complete successfully!")

# Extract error rates. Since some folds fail it will be an ugly list with both
# numeric estimates and NULL values (for the failed folds).
subtree(result, TRUE, "error")

# To put it on a more consistent form we can impute the missing error rates
# with NA to allow automatic simplification into a vector (since it requires
# all values to be on the same form, i.e. numeric() rather than a mix
# between numeric() and NULL as in the previous example).
subtree(result, TRUE, "error", error_value=as.numeric(NA), warn=-1)
trivial_error_rate

# Sum up feature importance for all classes within each fold and extract.
# Note that the lengths (= 4) must match between the folds for the automatic
# simplification to work.
subtree(result, TRUE, "importance", function(x){
  if(is.null(x)){
    rep(NA, 3)
  } else {
    colMeans(x[2:4])
  }
})

# The equivalent 'select' command would be ...
require(tidyverse)
imp <- result %>% select(fold = TRUE, "importance", function(x){
  if(is.null(x)) return(NULL)
  x %>% gather( Species, Importance, -feature)
})
require(ggplot2)
ggplot(imp, aes(x = Species, y = Importance)) +
  geom_abline(intercept = 0, slope = 0, color = "hotpink") +
  geom_boxplot() + facet_wrap(~feature)

---

trivial_error_rate  Calculate the trivial error rate

Description

Simply predicting the most common class for all test set observations can be a deceivingly successful strategy in terms of error rate. This function shows what error rate such a strategy would result in.

Usage

trivial_error_rate(truth)

Arguments

truth  True class labels.

Author(s)

Christofer Bäcklin
tune

Tune parameters of modeling procedures

Description
These functions are rarely needed to be called manually as they are automatically called by `fit` and `evaluate` when needed.

Usage
```r
tune(procedure, ..., .verbose = getOption("emil_verbose", FALSE))

is_tuned(procedure)

is_tunable(procedure)

detune(procedure)
```

Arguments
- `procedure` Modeling procedure, or list of modeling procedures, as produced by `modeling_procedure`.
- `...` Sent to `evaluate`.
- `verbose` Whether to print an activity log. Set to -1 to suppress all messages.

Value
- A tuned modeling procedures or a list of such.
- Logical indicating if the procedure(s) are tuned.
- Logical indicating if the has tunable parameters.
- A list of untuned modeling procedures.

Author(s)
Christofer Bäcklin

See Also
- `emil`, `modeling_procedure`, `evaluate`, `fit`, `predict`, `get_importance`

Examples
```r
procedure <- modeling_procedure("randomforest", parameter=list(mtry=1:4))
tuned.procedure <- tune(procedure, x=iris[-5], y=iris$Species)
mod <- fit(tuned.procedure, x=iris[-5], y=iris$Species)
```
validate_data  Validate a pre-processed data set

Description
While writing and debugging pre-processing functions this function can be useful to confirm that the resulting data sets fulfills the necessary requirements.

Usage
validate_data(data)

Arguments
data  Pre-processed data set.

Value
Nothing, only throws an error or prints a completion message.

Author(s)
Christofer Bäcklin

vlines  Add vertical or horizontal lines to a plot

Description
Add vertical or horizontal lines to a plot

Usage
vlines(x, lend = 1, ...)

hlines(y, lend = 1, ...)

Arguments
x  Coordinates of vertical lines.
lend  Line ending style, see par.
...  Sent to segments.
y  Coordinates of horizontal lines.
weighted_error_rate

Author(s)

Christofer Bäcklin

Examples

```r
plot(0:10, 0:10, type="n")
  hlines(0:4*2.5, col="#dddddd")
  points(0:10, 0:10)
```

---

### weighted_error_rate  Weighted error rate

Description

If different types of errors are associated with different costs a weighted error function might be more appropriate than the standard.

Usage

```r
weighted_error_rate(x)
```

Arguments

- `x` Cost matrix or factor response vector.

Details

This function is not in itself an error function, but used to generate error functions. Either supply a predefined cost matrix or a response vector for a classification problem to define it automatically.

The automatically generated cost matrix will generate an error of 0 if all predictions are correct, 1 if all predictions are incorrect and 0.5 if all predictions are the same (regardless of class, i.e. if one class is smaller it will be given a higher misclassification cost).

Value

An error function.

Author(s)

Christofer Bäcklin
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