

Package ‘mlr’

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Title Machine Learning in R

Version 2.18.0

Description Interface to a large number of classification and regression techniques, including machine-readable parameter descriptions. There is also an experimental extension for survival analysis, clustering and general, example-specific cost-sensitive learning. Generic resampling, including cross-validation, bootstrapping and subsampling. Hyperparameter tuning with modern optimization techniques, for single- and multi-objective problems. Filter and wrapper methods for feature selection. Extension of basic learners with additional operations common in machine learning, also allowing for easy nested resampling. Most operations can be parallelized.

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URL <https://mlr.mlr-org.com>, <https://github.com/mlr-org/mlr>

BugReports <https://github.com/mlr-org/mlr/issues>

Depends ParamHelpers (>= 1.10), R (>= 3.0.2)

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Suggests ada, adabag, bartMachine, batchtools, brnn, bst, C50, care, caret (>= 6.0-57), class, clue, cluster, ClusterR, clusterSim (>= 0.44-5), cmaes, cowplot, CoxBoost, crs, Cubist, deepnet, DiceKriging, DiscriMiner, e1071, earth, elasticnet, emoa, evtree, extraTrees, fda.usc, FDboost, FNN, forecast (>= 8.3), fpc, frbs, FSelector, FSelectorRcpp (>= 0.2.1), gbm, GenSA, ggpubr, glmnet, GPfit, h2o (>= 3.6.0.8), Hmisc, irace (>= 2.0), kernlab, kknn, klaR, knitr, laGP, LiblineaR, lintr (>= 1.0.0.9001), MASS, mboost, mco, mda, memoise, mlbench, mldr, mlrMBO, mmpf, modeltools, mRMRe, neuralnet, nnet, nodeHarvest (>= 0.7-3), numDeriv, pamr, pander, party, penalized (>= 0.9-47), pls, PMCMR (>= 4.1), praznik (>= 5.0.0), randomForest, randomForestSRC (>= 2.7.0), ranger (>= 0.8.0), rappdirs,

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mlr-package

mlr: Machine Learning in R

Description

Interface to a large number of classification and regression techniques, including machine-readable parameter descriptions. There is also an experimental extension for survival analysis, clustering and general, example-specific cost-sensitive learning. Generic resampling, including cross-validation, bootstrapping and subsampling. Hyperparameter tuning with modern optimization techniques, for single- and multi-objective problems. Filter and wrapper methods for feature selection. Extension of basic learners with additional operations common in machine learning, also allowing for easy nested resampling. Most operations can be parallelized.

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See Also

Useful links:

- <https://mlr.mlr-org.com>
- <https://github.com/mlr-org/mlr>
- Report bugs at <https://github.com/mlr-org/mlr/issues>

addRRMeasure	<i>Compute new measures for existing ResampleResult</i>
--------------	---

Description

Adds new measures to an existing ResampleResult.

Usage

```
addRRMeasure(res, measures)
```

Arguments

res	(ResampleResult) The result of resample run with <code>keep.pred = TRUE</code> .
measures	(Measure list of Measure) Performance measure(s) to evaluate. Default is the default measure for the task, see here getDefaultMeasure .

Value

([ResampleResult](#)).

See Also

Other resample: [ResamplePrediction](#), [ResampleResult](#), [getRRPredictionList\(\)](#), [getRRPredictions\(\)](#), [getRRTaskDescription\(\)](#), [getRRTaskDesc\(\)](#), [makeResampleDesc\(\)](#), [makeResampleInstance\(\)](#), [resample\(\)](#)

Aggregation	<i>Aggregation object.</i>
-------------	----------------------------

Description

An aggregation method reduces the performance values of the test (and possibly the training sets) to a single value. To see all possible implemented aggregations look at [aggregations](#).

The aggregation can access all relevant information of the result after resampling and combine them into a single value. Though usually something very simple like taking the mean of the test set performances is done.

Object members:

id (character(1)) Name of the aggregation method.

name (character(1)) Long name of the aggregation method.

properties (**character**) Properties of the aggregation.

fun (**'function(task, perf.test, perf.train, measure, group, pred)]**) Aggregation function.

See Also[makeAggregation](#)

aggregations

Aggregation methods.

Description

- **test.mean**
Mean of performance values on test sets.
- **test.sd**
Standard deviation of performance values on test sets.
- **test.median**
Median of performance values on test sets.
- **test.min**
Minimum of performance values on test sets.
- **test.max**
Maximum of performance values on test sets.
- **test.sum**
Sum of performance values on test sets.
- **train.mean**
Mean of performance values on training sets.
- **train.sd**
Standard deviation of performance values on training sets.
- **train.median**
Median of performance values on training sets.
- **train.min**
Minimum of performance values on training sets.
- **train.max**
Maximum of performance values on training sets.
- **train.sum**
Sum of performance values on training sets.
- **b632**
Aggregation for B632 bootstrap.
- **b632plus**
Aggregation for B632+ bootstrap.
- **testgroup.mean**
Performance values on test sets are grouped according to resampling method. The mean for every group is calculated, then the mean of those means. Mainly used for repeated CV.
- **testgroup.sd**
Similar to **testgroup.mean** - after the mean for every group is calculated, the standard deviation of those means is obtained. Mainly used for repeated CV.

- **test.join**

Performance measure on joined test sets. This is especially useful for small sample sizes where unbalanced group sizes have a significant impact on the aggregation, especially for cross-validation test.join might make sense now. For the repeated CV, the performance is calculated on each repetition and then aggregated with the arithmetic mean.

See Also

[Aggregation](#)

agri.task	<i>European Union Agricultural Workforces clustering task.</i>
-----------	--

Description

Contains the task (agri.task).

References

See [cluster::agriculture](#).

analyzeFeatSelResult	<i>Show and visualize the steps of feature selection.</i>
----------------------	---

Description

This function prints the steps [selectFeatures](#) took to find its optimal set of features and the reason why it stopped. It can also print information about all calculations done in each intermediate step.

Currently only implemented for sequential feature selection.

Usage

```
analyzeFeatSelResult(res, reduce = TRUE)
```

Arguments

res	(FeatSelResult) The result of of selectFeatures .
reduce	(logical(1)) Per iteration: Print only the selected feature (or all features that were evaluated)? Default is TRUE.

Value

(invisible(NULL)).

See Also

Other featsel: [FeatSelControl](#), [getFeatSelResult\(\)](#), [makeFeatSelWrapper\(\)](#), [selectFeatures\(\)](#)

asROCRPrediction	<i>Converts predictions to a format package ROCR can handle.</i>
------------------	--

Description

Converts predictions to a format package ROCR can handle.

Usage

```
asROCRPrediction(pred)
```

Arguments

pred	(Prediction) Prediction object.
------	--

See Also

Other roc: [calculateROCMasures\(\)](#)

Other predict: [getPredictionProbabilities\(\)](#), [getPredictionResponse\(\)](#), [getPredictionTaskDesc\(\)](#), [predict.WrappedModel\(\)](#), [setPredictThreshold\(\)](#), [setPredictType\(\)](#)

batchmark	<i>Run machine learning benchmarks as distributed experiments.</i>
-----------	--

Description

This function is a very parallel version of [benchmark](#) using [batchtools](#). Experiments are created in the provided registry for each combination of learners, tasks and resamplings. The experiments are then stored in a registry and the runs can be started via [batchtools::submitJobs](#). A job is one train/test split of the outer resampling. In case of nested resampling (e.g. with [makeTuneWrapper](#)), each job is a full run of inner resampling, which can be parallelized in a second step with **ParallelMap**.

For details on the usage and support backends have a look at the batchtools tutorial page: <https://github.com/mlg/batchtools>.

The general workflow with batchmark looks like this:

1. Create an ExperimentRegistry using [batchtools::makeExperimentRegistry](#).
2. Call `batchmark(...)` which defines jobs for all learners and tasks in an [base::expand.grid](#) fashion.
3. Submit jobs using [batchtools::submitJobs](#).
4. Babysit the computation, wait for all jobs to finish using [batchtools::waitForJobs](#).

5. Call `reduceBatchmarkResult()` to reduce results into a [BenchmarkResult](#).

If you want to use this with **OpenML** datasets you can generate tasks from a vector of dataset IDs easily with `tasks = lapply(data.ids, function(x) convertOMLDataSetToMlr(getOMLDataSet(x)))`.

Usage

```
batchmark(
  learners,
  tasks,
  resamplings,
  measures,
  keep.pred = TRUE,
  keep.extract = FALSE,
  models = FALSE,
  reg = batchtools::getDefaultRegistry()
)
```

Arguments

<code>learners</code>	(list of Learner character) Learning algorithms which should be compared, can also be a single learner. If you pass strings the learners will be created via makeLearner .
<code>tasks</code>	list of Task Tasks that learners should be run on.
<code>resamplings</code>	[(list of) ResampleDesc] Resampling strategy for each tasks. If only one is provided, it will be replicated to match the number of tasks. If missing, a 10-fold cross validation is used.
<code>measures</code>	(list of Measure) Performance measures for all tasks. If missing, the default measure of the first task is used.
<code>keep.pred</code>	(logical(1)) Keep the prediction data in the <code>pred</code> slot of the result object. If you do many experiments (on larger data sets) these objects might unnecessarily increase object size / mem usage, if you do not really need them. The default is set to TRUE.
<code>keep.extract</code>	(logical(1)) Keep the <code>extract</code> slot of the result object. When creating a lot of benchmark results with extensive tuning, the resulting R objects can become very large in size. That is why the tuning results stored in the <code>extract</code> slot are removed by default (<code>keep.extract = FALSE</code>). Note that when <code>keep.extract = FALSE</code> you will not be able to conduct analysis in the tuning results.
<code>models</code>	(logical(1)) Should all fitted models be stored in the ResampleResult ? Default is FALSE.
<code>reg</code>	(batchtools::Registry) Registry, created by batchtools::makeExperimentRegistry . If not explicitly passed, uses the last created registry.

Value

([data.table](#)). Generated job ids are stored in the column “job.id”.

See Also

Other benchmark: [BenchmarkResult](#), [benchmark\(\)](#), [convertBMRTorankMatrix\(\)](#), [friedmanPostHocTestBMR\(\)](#), [friedmanTestBMR\(\)](#), [generateCritDifferencesData\(\)](#), [getBMRAggrPerformances\(\)](#), [getBMRFeatSelResults\(\)](#), [getBMRFilteredFeatures\(\)](#), [getBMRLearnerIds\(\)](#), [getBMRLearnerShortNames\(\)](#), [getBMRLearners\(\)](#), [getBMRMeasureIds\(\)](#), [getBMRMeasures\(\)](#), [getBMRModels\(\)](#), [getBMRPerformances\(\)](#), [getBMRPredictions\(\)](#), [getBMRTaskDescs\(\)](#), [getBMRTaskIds\(\)](#), [getBMRTuneResults\(\)](#), [plotBMRBoxplots\(\)](#), [plotBMRRanksAsBarChart\(\)](#), [plotBMRSummary\(\)](#), [plotCritDifferences\(\)](#), [reduceBatchmarkResults\(\)](#)

bc.task

Wisconsin Breast Cancer classification task.

Description

Contains the task (`bc.task`).

References

See [mlbench::BreastCancer](#). The column “Id” and all incomplete cases have been removed from the task.

benchmark

Benchmark experiment for multiple learners and tasks.

Description

Complete benchmark experiment to compare different learning algorithms across one or more tasks w.r.t. a given resampling strategy. Experiments are paired, meaning always the same training / test sets are used for the different learners. Furthermore, you can of course pass “enhanced” learners via wrappers, e.g., a learner can be automatically tuned using [makeTuneWrapper](#).

Usage

```
benchmark(
  learners,
  tasks,
  resamplings,
  measures,
  keep.pred = TRUE,
  keep.extract = FALSE,
  models = FALSE,
  show.info = getMlrOption("show.info")
)
```

Arguments

learners	(list of Learner character) Learning algorithms which should be compared, can also be a single learner. If you pass strings the learners will be created via makeLearner .
tasks	list of Task Tasks that learners should be run on.
resamplings	(list of ResampleDesc ResampleInstance) Resampling strategy for each tasks. If only one is provided, it will be replicated to match the number of tasks. If missing, a 10-fold cross validation is used.
measures	(list of Measure) Performance measures for all tasks. If missing, the default measure of the first task is used.
keep.pred	(logical(1)) Keep the prediction data in the pred slot of the result object. If you do many experiments (on larger data sets) these objects might unnecessarily increase object size / mem usage, if you do not really need them. The default is set to TRUE.
keep.extract	(logical(1)) Keep the extract slot of the result object. When creating a lot of benchmark results with extensive tuning, the resulting R objects can become very large in size. That is why the tuning results stored in the extract slot are removed by default (keep.extract = FALSE). Note that when keep.extract = FALSE you will not be able to conduct analysis in the tuning results.
models	(logical(1)) Should all fitted models be stored in the ResampleResult ? Default is FALSE.
show.info	(logical(1)) Print verbose output on console? Default is set via configureMlr .

Value

[BenchmarkResult](#).

See Also

Other benchmark: [BenchmarkResult](#), [batchmark\(\)](#), [convertBMRTToRankMatrix\(\)](#), [friedmanPostHocTestBMR\(\)](#), [friedmanTestBMR\(\)](#), [generateCritDifferencesData\(\)](#), [getBMRAggrPerformances\(\)](#), [getBMRFeatSelResults\(\)](#), [getBMRFilteredFeatures\(\)](#), [getBMRLearnerIds\(\)](#), [getBMRLearnerShortNames\(\)](#), [getBMRLearners\(\)](#), [getBMRMeasureIds\(\)](#), [getBMRMeasures\(\)](#), [getBMRModels\(\)](#), [getBMRPerformances\(\)](#), [getBMRPredictions\(\)](#), [getBMRTaskDescs\(\)](#), [getBMRTaskIds\(\)](#), [getBMRTuneResults\(\)](#), [plotBMRBoxplots\(\)](#), [plotBMRRanksAsBarChart\(\)](#), [plotBMRSummary\(\)](#), [plotCritDifferences\(\)](#), [reduceBatchmarkResults\(\)](#)

Examples

```
lrns = list(makeLearner("classif.lda"), makeLearner("classif.rpart"))
tasks = list(iris.task, sonar.task)
rdesc = makeResampleDesc("CV", iters = 2L)
meas = list(acc, ber)
bmr = benchmark(lrns, tasks, rdsc, measures = meas)
```



```

rmat = convertBMRTToRankMatrix(bmr)
print(rmat)
plotBMRSummary(bmr)
plotBMRBoxplots(bmr, ber, style = "violin")
plotBMRRanksAsBarChart(bmr, pos = "stack")
friedmanTestBMR(bmr)
friedmanPostHocTestBMR(bmr, p.value = 0.05)

```

BenchmarkResult	<i>BenchmarkResult object.</i>
-----------------	--------------------------------

Description

Result of a benchmark experiment conducted by [benchmark](#) with the following members:

results (list of [ResampleResult](#)): A nested [list](#) of resample results, first ordered by task id, then by learner id.

measures (list of [Measure](#)): The performance measures used in the benchmark experiment.

learners (list of [Learner](#)): The learning algorithms compared in the benchmark experiment.

The print method of this object shows aggregated performance values for all tasks and learners.

It is recommended to retrieve required information via the `getBMR*` getter functions. You can also convert the object using [as.data.frame](#).

See Also

Other benchmark: [batchmark\(\)](#), [benchmark\(\)](#), [convertBMRTToRankMatrix\(\)](#), [friedmanPostHocTestBMR\(\)](#), [friedmanTestBMR\(\)](#), [generateCritDifferencesData\(\)](#), [getBMRAggrPerformances\(\)](#), [getBMRFeatSelResults\(\)](#), [getBMRFilteredFeatures\(\)](#), [getBMRLearnerIds\(\)](#), [getBMRLearnerShortNames\(\)](#), [getBMRLearners\(\)](#), [getBMRMeasureIds\(\)](#), [getBMRMeasures\(\)](#), [getBMRModels\(\)](#), [getBMRPerformances\(\)](#), [getBMRPredictions\(\)](#), [getBMRTaskDescs\(\)](#), [getBMRTaskIds\(\)](#), [getBMRTuneResults\(\)](#), [plotBMRBoxplots\(\)](#), [plotBMRRanksAsBarChart\(\)](#), [plotBMRSummary\(\)](#), [plotCritDifferences\(\)](#), [reduceBatchmarkResults\(\)](#)

bh.task	<i>Boston Housing regression task.</i>
---------	--

Description

Contains the task (`bh.task`).

References

See [mlbench::BostonHousing](#).

cache_helpers	<i>Get or delete mlr cache directory</i>
---------------	--

Description

Helper functions to deal with mlr caching.

Usage

```
getCacheDir()
deleteCacheDir()
```

Details

getCacheDir() returns the default mlr cache directory
 deleteCacheDir() clears the default mlr cache directory. Custom cache directories must be deleted by hand.

calculateConfusionMatrix	<i>Confusion matrix.</i>
--------------------------	--------------------------

Description

Calculates the confusion matrix for a (possibly resampled) prediction. Rows indicate true classes, columns predicted classes. The marginal elements count the number of classification errors for the respective row or column, i.e., the number of errors when you condition on the corresponding true (rows) or predicted (columns) class. The last bottom right element displays the total amount of errors.

A list is returned that contains multiple matrices. If `relative = TRUE` we compute three matrices, one with absolute values and two with relative. The relative confusion matrices are normalized based on rows and columns respectively, if `FALSE` we only compute the absolute value matrix.

The `print` function returns the relative matrices in a compact way so that both row and column marginals can be seen in one matrix. For details see [ConfusionMatrix](#).

Note that for resampling no further aggregation is currently performed. All predictions on all test sets are joined to a vector `yhat`, as are all labels joined to a vector `y`. Then `yhat` is simply tabulated vs. `y`, as if both were computed on a single test set. This probably mainly makes sense when cross-validation is used for resampling.

Usage

```
calculateConfusionMatrix(pred, relative = FALSE, sums = FALSE, set = "both")

## S3 method for class 'ConfusionMatrix'
print(x, both = TRUE, digits = 2, ...)
```

Arguments

pred	(Prediction) Prediction object.
relative	(logical(1)) If TRUE two additional matrices are calculated. One is normalized by rows and one by columns.
sums	(logical(1)) If TRUE add absolute number of observations in each group.
set	(character(1)) Specifies which part(s) of the data are used for the calculation. If set equals train or test, the pred object must be the result of a resampling, otherwise an error is thrown. Defaults to “both”. Possible values are “train”, “test”, or “both”.
x	(ConfusionMatrix) Object to print.
both	(logical(1)) If TRUE both the absolute and relative confusion matrices are printed.
digits	(integer(1)) How many numbers after the decimal point should be printed, only relevant for relative confusion matrices.
...	(any) Currently not used.

Value

([ConfusionMatrix](#)).

Methods (by generic)

- print:

See Also

Other performance: [ConfusionMatrix](#), [calculateROCMeasures\(\)](#), [estimateRelativeOverfitting\(\)](#), [makeCostMeasure\(\)](#), [makeCustomResampledMeasure\(\)](#), [makeMeasure\(\)](#), [measures](#), [performance\(\)](#), [setAggregation\(\)](#), [setMeasurePars\(\)](#)

Examples

```
# get confusion matrix after simple manual prediction
allinds = 1:150
train = sample(allinds, 75)
test = setdiff(allinds, train)
mod = train("classif.lda", iris.task, subset = train)
pred = predict(mod, iris.task, subset = test)
print(calculateConfusionMatrix(pred))
print(calculateConfusionMatrix(pred, sums = TRUE))
print(calculateConfusionMatrix(pred, relative = TRUE))
```

```
# now after cross-validation
r = crossval("classif.lda", iris.task, iters = 2L)
print(calculateConfusionMatrix(r$pred))
```

calculateROCMasures *Calculate receiver operator measures.*

Description

Calculate the absolute number of correct/incorrect classifications and the following evaluation measures:

- tpr True positive rate (Sensitivity, Recall)
- fpr False positive rate (Fall-out)
- fnr False negative rate (Miss rate)
- tnr True negative rate (Specificity)
- ppv Positive predictive value (Precision)
- for False omission rate
- lrp Positive likelihood ratio (LR+)
- fdr False discovery rate
- npv Negative predictive value
- acc Accuracy
- lrm Negative likelihood ratio (LR-)
- dor Diagnostic odds ratio

For details on the used measures see [measures](#) and also https://en.wikipedia.org/wiki/Receiver_operating_characteristic.

The element for the false omission rate in the resulting object is not called for but fomr since for should never be used as a variable name in an object.

Usage

```
calculateROCMasures(pred)

## S3 method for class 'ROCMasures'
print(x, abbreviations = TRUE, digits = 2, ...)
```

Arguments

pred	(Prediction) Prediction object.
x	(ROCMeasures) Created by calculateROCMeasures .
abbreviations	(logical(1)) If TRUE a short paragraph with explanations of the used measures is printed additionally.
digits	(integer(1)) Number of digits the measures are rounded to.
...	(any) Currently not used.

Value

(ROCMeasures). A list containing two elements `confusion.matrix` which is the 2 times 2 confusion matrix of absolute frequencies and measures, a list of the above mentioned measures.

Methods (by generic)

- `print`:

See Also

Other roc: [asROCRPrediction\(\)](#)

Other performance: [ConfusionMatrix](#), [calculateConfusionMatrix\(\)](#), [estimateRelativeOverfitting\(\)](#), [makeCostMeasure\(\)](#), [makeCustomResampledMeasure\(\)](#), [makeMeasure\(\)](#), [measures](#), [performance\(\)](#), [setAggregation\(\)](#), [setMeasurePars\(\)](#)

Examples

```
lrn = makeLearner("classif.rpart", predict.type = "prob")
fit = train(lrn, sonar.task)
pred = predict(fit, task = sonar.task)
calculateROCMeasures(pred)
```

capLargeValues

Convert large/infinite numeric values in a data.frame or task.

Description

Convert numeric entries which large/infinite (absolute) values in a data.frame or task. Only numeric/integer columns are affected.

Usage

```
capLargeValues(  
  obj,  
  target = character(0L),  
  cols = NULL,  
  threshold = Inf,  
  impute = threshold,  
  what = "abs"  
)
```

Arguments

obj	(data.frame Task) Input data.
target	(character) Name of the column(s) specifying the response. Target columns will not be capped. Default is <code>character(0)</code> .
cols	(character) Which columns to convert. Default is all numeric columns.
threshold	(numeric(1)) Threshold for capping. Every entry whose absolute value is equal or larger is converted. Default is <code>Inf</code> .
impute	(numeric(1)) Replacement value for large entries. Large negative entries are converted to <code>-impute</code> . Default is <code>threshold</code> .
what	(character(1)) What kind of entries are affected? “abs” means $\text{abs}(x) > \text{threshold}$, “pos” means $\text{abs}(x) > \text{threshold} \ \&\& \ x > 0$, “neg” means $\text{abs}(x) > \text{threshold} \ \&\& \ x < 0$. Default is “abs”.

Value

([data.frame](#))

See Also

Other `eda_and_preprocess`: [createDummyFeatures\(\)](#), [dropFeatures\(\)](#), [mergeSmallFactorLevels\(\)](#), [normalizeFeatures\(\)](#), [removeConstantFeatures\(\)](#), [summarizeColumns\(\)](#), [summarizeLevels\(\)](#)

Examples

```
capLargeValues(iris, threshold = 5, impute = 5)
```

`configureMlr`*Configures the behavior of the package.*

Description

Configuration is done by setting custom [options](#).

If you do not set an option here, its current value will be kept.

If you call this function with an empty argument list, everything is set to its defaults.

Usage

```
configureMlr(  
  show.info,  
  on.learner.error,  
  on.learner.warning,  
  on.par.without.desc,  
  on.par.out.of.bounds,  
  on.measure.not.applicable,  
  show.learner.output,  
  on.error.dump  
)
```

Arguments

<code>show.info</code>	(logical(1)) Some methods of <code>mlr</code> support a <code>show.info</code> argument to enable verbose output on the console. This option sets the default value for these arguments. Setting the argument manually in one of these functions will overwrite the default value for that specific function call. Default is TRUE.
<code>on.learner.error</code>	(character(1)) What should happen if an error in an underlying learning algorithm is caught: “stop”: R exception is generated. “warn”: A <code>FailureModel</code> will be created, which predicts only NAs and a warning will be generated. “quiet”: Same as “warn” but without the warning. Default is “stop”.
<code>on.learner.warning</code>	(character(1)) What should happen if a warning in an underlying learning algorithm is generated: “warn”: The warning is generated as usual. “quiet”: The warning is suppressed. Default is “warn”.

- `on.par.without.desc`
 (character(1))
 What should happen if a parameter of a learner is set to a value, but no parameter description object exists, indicating a possibly wrong name:
 “stop”: R exception is generated.
 “warn”: Warning, but parameter is still passed along to learner.
 “quiet”: Same as “warn” but without the warning.
 Default is “stop”.
- `on.par.out.of.bounds`
 (character(1))
 What should happen if a parameter of a learner is set to an out of bounds value.
 “stop”: R exception is generated.
 “warn”: Warning, but parameter is still passed along to learner.
 “quiet”: Same as “warn” but without the warning.
 Default is “stop”.
- `on.measure.not.applicable`
 (logical(1))
 What should happen if a measure is not applicable to a learner.
 “stop”: R exception is generated.
 “warn”: Warning, but value of the measure will be NA.
 “quiet”: Same as “warn” but without the warning.
 Default is “stop”.
- `show.learner.output`
 (logical(1))
 Should the output of the learning algorithm during training and prediction be shown or captured and suppressed? Default is TRUE.
- `on.error.dump` (logical(1))
 Specify whether [FailureModel](#) models and failed predictions should contain an error dump that can be used with debugger to inspect an error. This option is only effective if `on.learner.error` is “warn” or “quiet”. If it is TRUE, the dump can be accessed using [getFailureModelDump](#) on the [FailureModel](#), [getPredictionDump](#) on the failed prediction, and [getRRDump](#) on resample predictions. Default is FALSE.

Value

(invisible(NULL)).

See Also

Other configure: [getMlrOptions\(\)](#)

Description

The result of [calculateConfusionMatrix](#).

Object members:

result (**matrix**) Confusion matrix of absolute values and marginals. Can also contain row and column sums of observations.

task.desc (**TaskDesc**) Additional information about the task.

sums (logical(1)) Flag if marginal sums of observations are calculated.

relative (logical(1)) Flag if the relative confusion matrices are calculated.

relative.row (**matrix**) Confusion matrix of relative values and marginals normalized by row.

relative.col (**matrix**) Confusion matrix of relative values and marginals normalized by column.

relative.error (numeric(1)) Relative error overall.

See Also

Other performance: [calculateConfusionMatrix\(\)](#), [calculateROCMeasures\(\)](#), [estimateRelativeOverfitting\(\)](#), [makeCostMeasure\(\)](#), [makeCustomResampledMeasure\(\)](#), [makeMeasure\(\)](#), [measures](#), [performance\(\)](#), [setAggregation\(\)](#), [setMeasurePars\(\)](#)

convertBMRTToRankMatrix

Convert BenchmarkResult to a rank-matrix.

Description

Computes a matrix of all the ranks of different algorithms over different datasets (tasks). Ranks are computed from aggregated measures. Smaller ranks imply better methods, so for measures that are minimized, small ranks imply small scores. for measures that are maximized, small ranks imply large scores.

Usage

```
convertBMRTToRankMatrix(
  bmr,
  measure = NULL,
  ties.method = "average",
  aggregation = "default"
)
```

Arguments

bmr	(BenchmarkResult) Benchmark result.
measure	(Measure) Performance measure. Default is the first measure used in the benchmark experiment.
ties.method	(character(1)) See base::rank for details.
aggregation	(character(1)) “mean” or “default”. See getBMRAggrPerformances for details on “default”.

Value

([matrix](#)) with measure ranks as entries. The matrix has one row for each learner, and one column for each task.

See Also

Other benchmark: [BenchmarkResult](#), [batchmark\(\)](#), [benchmark\(\)](#), [friedmanPostHocTestBMR\(\)](#), [friedmanTestBMR\(\)](#), [generateCritDifferencesData\(\)](#), [getBMRAggrPerformances\(\)](#), [getBMRFeatSelResults\(\)](#), [getBMRFilteredFeatures\(\)](#), [getBMRLearnerIds\(\)](#), [getBMRLearnerShortNames\(\)](#), [getBMRLearners\(\)](#), [getBMRMeasureIds\(\)](#), [getBMRMeasures\(\)](#), [getBMRModels\(\)](#), [getBMRPerformances\(\)](#), [getBMRPredictions\(\)](#), [getBMRTaskDescs\(\)](#), [getBMRTaskIds\(\)](#), [getBMRTuneResults\(\)](#), [plotBMRBoxplots\(\)](#), [plotBMRRanksAsBarChart\(\)](#), [plotBMRSummary\(\)](#), [plotCritDifferences\(\)](#), [reduceBatchmarkResults\(\)](#)

Examples

```
# see benchmark
```

```
convertMLBenchObjToTask
```

Convert a machine learning benchmark / demo object from package mlbench to a task.

Description

We auto-set the target column, drop any column which is called “Id” and convert logicals to factors.

Usage

```
convertMLBenchObjToTask(x, n = 100L, ...)
```

Arguments

x	(character(1)) Name of an mlbench function or dataset.
n	(integer(1)) Number of observations for data simul functions. Note that for a few mlbench function this setting is not exactly respected by mlbench. Default is 100.
...	(any) Passed on to data simul functions.

Examples

```
print(convertMLBenchObjToTask("Ionosphere"))
print(convertMLBenchObjToTask("mlbench.spirals", n = 100, sd = 0.1))
```

costiris.task	<i>Iris cost-sensitive classification task.</i>
---------------	---

Description

Contains the task (costiris.task).

References

See [datasets::iris](#). The cost matrix was generated artificially following Tu, H.-H. and Lin, H.-T. (2010), One-sided support vector regression for multiclass cost-sensitive classification. In ICML, J. Fürnkranz and T. Joachims, Eds., Omnipress, 1095–1102.

createDummyFeatures	<i>Generate dummy variables for factor features.</i>
---------------------	--

Description

Replace all factor features with their dummy variables. Internally [model.matrix](#) is used. Non factor features will be left untouched and passed to the result.

Usage

```
createDummyFeatures(
  obj,
  target = character(0L),
  method = "1-of-n",
  cols = NULL
)
```

Arguments

obj	(data.frame Task) Input data.
target	(character (1) character (2) character (n.classes)) Name(s) of the target variable(s). Only used when obj is a data.frame, otherwise ignored. If survival analysis is applicable, these are the names of the survival time and event columns, so it has length 2. For multilabel classification these are the names of logical columns that indicate whether a class label is present and the number of target variables corresponds to the number of classes.
method	(character (1)) Available are: "1-of-n" : For n factor levels there will be n dummy variables. "reference" : There will be n-1 dummy variables leaving out the first factor level of each variable. Default is "1-of-n".
cols	(character) Columns to create dummy features for. Default is to use all columns.

Value

[data.frame](#) | [Task](#). Same type as obj.

See Also

Other `eda_and_preprocess`: [capLargeValues\(\)](#), [dropFeatures\(\)](#), [mergeSmallFactorLevels\(\)](#), [normalizeFeatures\(\)](#), [removeConstantFeatures\(\)](#), [summarizeColumns\(\)](#), [summarizeLevels\(\)](#)

createSpatialResamplingPlots

Create (spatial) resampling plot objects.

Description

Visualize partitioning of resample objects with spatial information.

Usage

```
createSpatialResamplingPlots(
  task = NULL,
  resample = NULL,
  crs = NULL,
  datum = 4326,
  repetitions = 1,
  color.train = "#0072B5",
  color.test = "#E18727",
```

```

    point.size = 0.5,
    axis.text.size = 14,
    x.axis.breaks = waiver(),
    y.axis.breaks = waiver()
  )

```

Arguments

task	Task Task object.
resample	ResampleResult or named list with (multiple) ResampleResult As returned by resample .
crs	integer Coordinate reference system (EPSG code number) for the supplied coordinates in the Task.
datum	integer Coordinate reference system which should be used in the resulting map.
repetitions	integer Number of repetitions.
color.train	character Color for train set.
color.test	character Color for test set.
point.size	integer Point size.
axis.text.size	integer Font size of axis labels.
x.axis.breaks	numeric Custom x axis breaks
y.axis.breaks	numeric Custom y axis breaks

Details

If a named list is given to `resample`, names will appear in the title of each fold. If multiple inputs are given to `resample`, these must be named.

This function makes a hard cut at five columns of the resulting gridded plot. This means if the `resample` object consists of `foldes > 5`, these folds will be put into the new row.

For file saving, we recommend to use `cowplot::save_plot`.

When viewing the resulting plot in RStudio, margins may appear to be different than they really are. Make sure to save the file to disk and inspect the image.

When modifying axis breaks, negative values need to be used if the area is located in either the western or southern hemisphere. Use positive values for the northern and eastern hemisphere.

Value

(list of 2L containing (1) multiple ‘gg’ objects and (2) their corresponding labels.

CRS

The crs has to be suitable for the coordinates stored in the Task. For example, if the coordinates are UTM, crs should be set to a UTM projection. Due to a limited axis space in the resulting grid (especially on the x-axis), the data will by default be projected into a lat/lon projection, specifically EPSG 4326. If other projections are desired for the resulting map, please set argument datum accordingly. This argument will be passed onto `ggplot2::coord_sf`.

Author(s)

Patrick Schratz

See Also

Other plot: `plotBMRBoxplots()`, `plotBMRRanksAsBarChart()`, `plotBMRSummary()`, `plotCalibration()`, `plotCritDifferences()`, `plotLearningCurve()`, `plotPartialDependence()`, `plotROCCurves()`, `plotResiduals()`, `plotThreshVsPerf()`

Examples

```
rdesc = makeResampleDesc("SpRepCV", folds = 5, reps = 4)
r = resample(makeLearner("classif.qda"), spatial.task, rdesc)

## -----
## single unnamed resample input with 5 folds and 2 repetitions
## -----

plots = createSpatialResamplingPlots(spatial.task, r, crs = 32717,
  repetitions = 2, x.axis.breaks = c(-79.065, -79.085),
  y.axis.breaks = c(-3.970, -4))
cowplot::plot_grid(plotlist = plots[["Plots"]], ncol = 5, nrow = 2,
  labels = plots[["Labels"]])

## -----
## single named resample input with 5 folds and 1 repetition and 32717 datum
## -----

plots = createSpatialResamplingPlots(spatial.task, list("Resamp" = r),
  crs = 32717, datum = 32717, repetitions = 1)
cowplot::plot_grid(plotlist = plots[["Plots"]], ncol = 5, nrow = 1,
  labels = plots[["Labels"]])

## -----
## multiple named resample inputs with 5 folds and 1 repetition
## -----

rdesc1 = makeResampleDesc("SpRepCV", folds = 5, reps = 4)
```

```

r1 = resample(makeLearner("classif.qda"), spatial.task, rdesc1)
rdesc2 = makeResampleDesc("RepCV", folds = 5, reps = 4)
r2 = resample(makeLearner("classif.qda"), spatial.task, rdesc2)

plots = createSpatialResamplingPlots(spatial.task,
  list("SpRepCV" = r1, "RepCV" = r2), crs = 32717, repetitions = 1,
  x.axis.breaks = c(-79.055, -79.085), y.axis.breaks = c(-3.975, -4))
cowplot::plot_grid(plotlist = plots[["Plots"]], ncol = 5, nrow = 2,
  labels = plots[["Labels"]])

## -----
## Complex arrangements of multiple named resample inputs with 5 folds and 1 repetition
## -----

p1 = cowplot::plot_grid(plots[["Plots"]][[1]], plots[["Plots"]][[2]],
  plots[["Plots"]][[3]], ncol = 3, nrow = 1, labels = plots[["Labels"]][1:3],
  label_size = 18)
p12 = cowplot::plot_grid(plots[["Plots"]][[4]], plots[["Plots"]][[5]],
  ncol = 2, nrow = 1, labels = plots[["Labels"]][4:5], label_size = 18)

p2 = cowplot::plot_grid(plots[["Plots"]][[6]], plots[["Plots"]][[7]],
  plots[["Plots"]][[8]], ncol = 3, nrow = 1, labels = plots[["Labels"]][6:8],
  label_size = 18)
p22 = cowplot::plot_grid(plots[["Plots"]][[9]], plots[["Plots"]][[10]],
  ncol = 2, nrow = 1, labels = plots[["Labels"]][9:10], label_size = 18)

cowplot::plot_grid(p1, p12, p2, p22, ncol = 1)

```

crossover

Crossover.

Description

Takes two bit strings and creates a new one of the same size by selecting the items from the first string or the second, based on a given rate (the probability of choosing an element from the first string).

Arguments

x	(logical) First parent string.
y	(logical) Second parent string.
rate	(numeric(1)) A number representing the probability of selecting an element of the first string. Default is 0.5.

Value

([crossover](#)).

downsample	<i>Downsample (subsample) a task or a data.frame.</i>
------------	---

Description

Decrease the observations in a task or a ResampleInstance to a given percentage of observations.

Usage

```
downsample(obj, perc = 1, stratify = FALSE)
```

Arguments

obj	(Task ResampleInstance) Input data or a ResampleInstance.
perc	(numeric(1)) Percentage from (0, 1). Default is 1.
stratify	(logical(1)) Only for classification: Should the downsampled data be stratified according to the target classes? Default is FALSE.

Value

([data.frame] | [Task] | [ResampleInstance]). Same type as obj'.

See Also

[makeResampleInstance](#)

Other downsample: [makeDownsampleWrapper\(\)](#)

dropFeatures	<i>Drop some features of task.</i>
--------------	------------------------------------

Description

Drop some features of task.

Usage

```
dropFeatures(task, features)
```


Arguments

task	(Task) The task.
features	(character) Features to drop.

Value

[Task](#).

See Also

Other eda_and_preprocess: [capLargeValues\(\)](#), [createDummyFeatures\(\)](#), [mergeSmallFactorLevels\(\)](#), [normalizeFeatures\(\)](#), [removeConstantFeatures\(\)](#), [summarizeColumns\(\)](#), [summarizeLevels\(\)](#)

estimateRelativeOverfitting

Estimate relative overfitting.

Description

Estimates the relative overfitting of a model as the ratio of the difference in test and train performance to the difference of test performance in the no-information case and train performance. In the no-information case the features carry no information with respect to the prediction. This is simulated by permuting features and predictions.

Usage

```
estimateRelativeOverfitting(  
  predish,  
  measures,  
  task,  
  learner = NULL,  
  pred.train = NULL,  
  iter = 1  
)
```

Arguments

predish	(ResampleDesc ResamplePrediction Prediction) Resampling strategy or resampling prediction or test predictions.
measures	(Measure list of Measure) Performance measure(s) to evaluate. Default is the default measure for the task, see here getDefaultMeasure .
task	(Task) The task.

learner	(Learner character(1)) The learner. If you pass a string the learner will be created via makeLearner .
pred.train	(Prediction) Training predictions. Only needed if test predictions are passed.
iter	(integer) Iteration number. Default 1, usually you don't need to specify this. Only needed if test predictions are passed.

Details

Currently only support for classification and regression tasks is implemented.

Value

([data.frame](#)). Relative overfitting estimate(s), named by measure(s), for each resampling iteration.

References

Bradley Efron and Robert Tibshirani; Improvements on Cross-Validation: The .632+ Bootstrap Method, Journal of the American Statistical Association, Vol. 92, No. 438. (Jun., 1997), pp. 548-560.

See Also

Other performance: [ConfusionMatrix](#), [calculateConfusionMatrix\(\)](#), [calculateROCMeasures\(\)](#), [makeCostMeasure\(\)](#), [makeCustomResampledMeasure\(\)](#), [makeMeasure\(\)](#), [measures](#), [performance\(\)](#), [setAggregation\(\)](#), [setMeasurePars\(\)](#)

Examples

```
task = makeClassifTask(data = iris, target = "Species")
rdesc = makeResampleDesc("CV", iters = 2)
estimateRelativeOverfitting(rdesc, acc, task, makeLearner("classif.knn"))
estimateRelativeOverfitting(rdesc, acc, task, makeLearner("classif.lda"))
rpred = resample("classif.knn", task, rdesc)$pred
estimateRelativeOverfitting(rpred, acc, task)
```

```
estimateResidualVariance
```

Estimate the residual variance.

Description

Estimate the residual variance of a regression model on a given task. If a regression learner is provided instead of a model, the model is trained (see [train](#)) first.

Usage

```
estimateResidualVariance(x, task, data, target)
```

Arguments

x	(Learner or WrappedModel) Learner or wrapped model.
task	(RegrTask) Regression task. If missing, data and target must be supplied.
data	(data.frame) A data frame containing the features and target variable. If missing, task must be supplied.
target	(<code>character(1)</code>) Name of the target variable. If missing, task must be supplied.

extractFDABsignal	<i>Bspline mlq features</i>
-------------------	-----------------------------

Description

The function extracts features from functional data based on the Bspline fit. For more details refer to [FDboost::bsignal\(\)](#).

Usage

```
extractFDABsignal(bsignal.knots = 10L, bsignal.df = 3)
```

Arguments

bsignal.knots	(<code>integer(1)</code>) The number of knots for bspline.
bsignal.df	(<code>numeric(1)</code>) The effective degree of freedom of penalized bspline.

Value

([data.frame](#)).

See Also

Other fda_featextractor: [extractFDADTWKernel\(\)](#), [extractFDAFPCA\(\)](#), [extractFDAFourier\(\)](#), [extractFDAMultiResFeatures\(\)](#), [extractFDATsfeatures\(\)](#), [extractFDAWavelets\(\)](#)

extractFDADTWKernel *DTW kernel features*

Description

The function extracts features from functional data based on the DTW distance with a reference dataframe.

Usage

```
extractFDADTWKernel(  
  ref.method = "random",  
  n.refs = 0.05,  
  refs = NULL,  
  dtwindow = 0.05  
)
```

Arguments

ref.method	(character(1)) How should the reference curves be obtained? Method random draws n.refs random reference curves, while all uses all curves as references. In order to use user-provided reference curves, this parameter is set to fixed.
n.refs	(numeric(1)) Number of reference curves to be drawn (as a fraction of the number of observations in the training data).
refs	(matrix integer(n)) Integer vector of training set row indices or a matrix of reference curves with the same length as the functionals in the training data. Overwrites ref.method and n.refs.
dtwindow	(numeric(1)) Size of the warping window size (as a proportion of query length).

Value

([data.frame](#)).

See Also

Other fda_featextractor: [extractFDABsignal\(\)](#), [extractFDAFPCA\(\)](#), [extractFDAFourier\(\)](#), [extractFDAMultiResFeatu](#), [extractFDATsfeatures\(\)](#), [extractFDAWavelets\(\)](#)

extractFDAFeatures *Extract features from functional data.*

Description

Extract non-functional features from functional features using various methods.

The function `extractFDAFeatures` performs the extraction for all functional features via the methods specified in `feat.methods` and transforms all mentioned functional (matrix) features into regular data.frame columns. Additionally, a “extractFDAFeatDesc” object which contains learned coefficients and other helpful data for re-extraction during the predict-phase is returned. This can be used with `reextractFDAFeatures` in order to extract features during the prediction phase.

Usage

```
extractFDAFeatures(obj, target = character(0L), feat.methods = list(), ...)
```

Arguments

<code>obj</code>	(Task data.frame) Task or data.frame to extract functional features from. Must contain functional features as matrix columns.
<code>target</code>	(character(1)) Task target column. Only necessary for data.frames Default is <code>character(0)</code> .
<code>feat.methods</code>	(named list) List of functional features along with the desired methods for each functional feature. “all” applies the <code>extractFDAFeatures</code> method to each functional feature. Names of <code>feat.methods</code> must match column names of functional features. Available feature extraction methods are available under family <code>fda_featextractor</code> . Specifying a functional feature multiple times with different extraction methods allows for the extraction of different features from the same functional. Default is <code>list()</code> which does nothing.
<code>...</code>	(any) Further hyperparameters passed on to the <code>feat.methods</code> specified above.

Details

The description object contains these slots:

- target `character`: See argument.
- coln `character`: Colum names of data.
- fd.cols `character`: Functional feature names.
- extractFDAFeat `list`: Contains `feature.methods` and relevant parameters for reextraction.

Value

(list)
 data|task ([data.frame](#) | [Task](#))
 Extracted features, same type as obj.
 desc ([extractFDAFeatDesc](#))
 Description object. See description for details.

See Also

Other fda: [makeExtractFDAFeatMethod\(\)](#), [makeExtractFDAFeatsWrapper\(\)](#)

Examples

```
df = data.frame(x = matrix(rnorm(24), ncol = 8), y = factor(c("a", "a", "b")))
fdf = makeFunctionalData(df, fd.features = list(x1 = 1:4, x2 = 5:8), exclude.cols = "y")
task = makeClassifTask(data = fdf, target = "y")
extracted = extractFDAFeatures(task,
  feat.methods = list("x1" = extractFDAFourier(), "x2" = extractFDWavelets(filter = "haar")))
print(extracted$task)
reextractFDAFeatures(task, extracted$desc)
```

`extractFDAFourier` *Fast Fourier transform features.*

Description

The function extracts features from functional data based on the fast fourier transform. For more details refer to [stats::fft](#).

Usage

```
extractFDAFourier(trafo.coeff = "phase")
```

Arguments

`trafo.coeff` ([character\(1\)](#))
 Specifies which transformation of the complex frequency domain representation should be calculated as a feature representation. Must be one of “amplitude” or “phase”. Default is “phase”. The phase shift is returned in Rad, i.e. values lie in [-180, 180].

Value

([data.frame](#)).

See Also

Other fda_featextractor: [extractFDABsignal\(\)](#), [extractFDADTWKernel\(\)](#), [extractFDAFPCA\(\)](#), [extractFDAMultiResFeatures\(\)](#), [extractFDATsfeatures\(\)](#), [extractFDWavelets\(\)](#)

extractFDAFPCA	<i>Extract functional principal component analysis features.</i>
----------------	--

Description

The function extracts the functional principal components from a data.frame containing functional features. Uses stats::prcomp.

Usage

```
extractFDAFPCA(rank. = NULL, center = TRUE, scale. = FALSE)
```

Arguments

rank.	(integer(1)) Number of principal components to extract. Default is NULL
center	(logical(1)) Should data be centered before applying PCA?
scale.	(logical(1)) Should data be scaled before applying PCA?

Value

([data.frame](#)).

See Also

Other fda_featextractor: [extractFDABsignal\(\)](#), [extractFDADTWKernel\(\)](#), [extractFDAFourier\(\)](#), [extractFDAMultiResFeatures\(\)](#), [extractFDATsfeatures\(\)](#), [extractFDAWavelets\(\)](#)

extractFDAMultiResFeatures	<i>Multiresolution feature extraction.</i>
----------------------------	--

Description

The function extracts currently the mean of multiple segments of each curve and stacks them as features. The segments length are set in a hierarchy way so the features cover different resolution levels.

Usage

```
extractFDAMultiResFeatures(res.level = 3L, shift = 0.5, seg.lens = NULL)
```

Arguments

res.level	(integer(1)) The number of resolution hierachy, each length is divided by a factor of 2.
shift	(numeric(1)) The overlapping proportion when slide the window for one step.
seg.lens	(integer(1)) Curve subsequence lengths. Needs to sum up to the length of the functional.

Value

(data.frame).

See Also

Other fda_featextractor: [extractFDABsignal\(\)](#), [extractFDADTWKernel\(\)](#), [extractFDAFPCA\(\)](#), [extractFDAFourier\(\)](#), [extractFDATsfeatures\(\)](#), [extractFDAAWavelets\(\)](#)

extractFDATsfeatures *Time-Series Feature Heuristics*

Description

The function extracts features from functional data based on known Heuristics. For more details refer to [tsfeatures::tsfeatures\(\)](#). Under the hood this function uses the package [tsfeatures::tsfeatures\(\)](#). For more information see Hyndman, Wang and Laptev, Large-Scale Unusual Time Series Detection, ICDM 2015.

Note: Currently computes the following features:

"frequency", "stl_features", "entropy", "acf_features", "arch_stat", "crossing_points", "flat_spots", "hurst", "holt_parameters", "lumpiness", "max_kl_shift", "max_var_shift", "max_level_shift", "stability", "nonlinearity"

Usage

```
extractFDATsfeatures(
  scale = TRUE,
  trim = FALSE,
  trim_amount = 0.1,
  parallel = FALSE,
  na.action = na.pass,
  feats = NULL,
  ...
)
```


Arguments

scale	(logical(1)) If TRUE, time series are scaled to mean 0 and sd 1 before features are computed.
trim	(logical(1)) If TRUE, time series are trimmed by trim_amount before features are computed. Values larger than trim_amount in absolute value are set to NA.
trim_amount	(numeric(1)) Default level of trimming if trim==TRUE.
parallel	(logical(1)) If TRUE, multiple cores (or multiple sessions) will be used. This only speeds things up when there are a large number of time series.
na.action	(logical(1)) A function to handle missing values. Use na.interp to estimate missing values
feats	(character) A character vector of function names to apply to each time-series in order to extract features. Default: feats = c("frequency", "stl_features", "entropy", "acf_features", "arch_stat", "crossing_points", "flat_spots", "hurst", "holt_parameters", "lumpiness", "max_kl_shift", "max_var_shift", "max_level_shift", "stability", "nonlinearity")
...	(any) Further arguments passed on to the respective tsfeatures functions.

Value

([data.frame](#))

References

Hyndman, Wang and Laptev, Large-Scale Unusual Time Series Detection, ICDM 2015.

See Also

Other `fda_featextractor`: [extractFDABsignal\(\)](#), [extractFDADTWKernel\(\)](#), [extractFDAFPCA\(\)](#), [extractFDAFourier\(\)](#), [extractFDAMultiResFeatures\(\)](#), [extractFDAWavelets\(\)](#)

extractFDAWavelets *Discrete Wavelet transform features.*

Description

The function extracts discrete wavelet transform coefficients from the raw functional data. See [wavelets::dwt](#) for more information.

Usage

```
extractFDWavelets(filter = "la8", boundary = "periodic")
```

Arguments

filter	(character(1)) Specifies which filter should be used. Must be one of d1l1b1lc followed by an even number for the level of the filter. The level of the filter needs to be smaller or equal then the time-series length. For more information and acceptable filters see <code>help(wt.filter)</code> . Defaults to la8.
boundary	(character(1)) Boundary to be used. "periodic" assumes circular time series, for "reflection" the series is extended to twice its length. Default is "periodic".

Value

([data.frame](#)).

See Also

Other `fda_featextractor`: [extractFDABsignal\(\)](#), [extractFDADTWKernel\(\)](#), [extractFDAFPCA\(\)](#), [extractFDAFourier\(\)](#), [extractFDAMultiResFeatures\(\)](#), [extractFDATsfeatures\(\)](#)

FailureModel

Failure model.

Description

A subclass of [WrappedModel](#). It is created

- if you set the respective option in [configureMlr](#) - when a model internally crashed during training. The model always predicts NAs.

The if `mlr` option `on.error.dump` is TRUE, the `FailureModel` contains the debug trace of the error. It can be accessed with `getFailureModelDump` and inspected with debugger.

Its encapsulated learner `.model` is simply a string: The error message that was generated when the model crashed. The following code shows how to access the message.

See Also

Other debug: [ResampleResult](#), [getPredictionDump\(\)](#), [getRRDump\(\)](#)

Examples

```

configureMlr(on.learner.error = "warn")
data = iris
data$newfeat = 1 # will make LDA crash
task = makeClassifTask(data = data, target = "Species")
m = train("classif.lda", task) # LDA crashed, but mlr catches this
print(m)
print(m$learner.model) # the error message
p = predict(m, task) # this will predict NAs
print(p)
print(performance(p))
configureMlr(on.learner.error = "stop")

```

FeatSelControl

Create control structures for feature selection.

Description

Feature selection method used by [selectFeatures](#).

The methods used here follow a wrapper approach, described in Kohavi and John (1997) (see references).

The following optimization algorithms are available:

FeatSelControlExhaustive Exhaustive search. All feature sets (up to a certain number of features `max.features`) are searched.

FeatSelControlRandom Random search. Features vectors are randomly drawn, up to a certain number of features `max.features`. A feature is included in the current set with probability `prob`. So we are basically drawing (0,1)-membership-vectors, where each element is Bernoulli(`prob`) distributed.

FeatSelControlSequential Deterministic forward or backward search. That means extending (forward) or shrinking (backward) a feature set. Depending on the given method different approaches are taken.

`sfs` Sequential Forward Search: Starting from an empty model, in each step the feature increasing the performance measure the most is added to the model.

`sbs` Sequential Backward Search: Starting from a model with all features, in each step the feature decreasing the performance measure the least is removed from the model.

`sffs` Sequential Floating Forward Search: Starting from an empty model, in each step the algorithm chooses the best model from all models with one additional feature and from all models with one feature less.

`sfbs` Sequential Floating Backward Search: Similar to `sffs` but starting with a full model.

FeatSelControlGA Search via genetic algorithm. The GA is a simple (`mu`, `lambda`) or (`mu` + `lambda`) algorithm, depending on the comma setting. A comma strategy selects a new population of size `mu` out of the `lambda` > `mu` offspring. A plus strategy uses the joint pool of `mu` parents and `lambda` offspring for selecting `mu` new candidates. Out of those `mu` features, the new `lambda` features are generated by randomly choosing pairs of parents. These are crossed over and `crossover.rate` represents the probability of choosing a feature from the first parent instead of the second parent. The resulting offspring is mutated, i.e., its bits are flipped

with probability `mutation.rate`. If `max.features` is set, offspring are repeatedly generated until the setting is satisfied.

Usage

```
makeFeatSelControlExhaustive(  
  same.resampling.instance = TRUE,  
  maxit = NA_integer_,  
  max.features = NA_integer_,  
  tune.threshold = FALSE,  
  tune.threshold.args = list(),  
  log.fun = "default"  
)
```

```
makeFeatSelControlGA(  
  same.resampling.instance = TRUE,  
  impute.val = NULL,  
  maxit = NA_integer_,  
  max.features = NA_integer_,  
  comma = FALSE,  
  mu = 10L,  
  lambda,  
  crossover.rate = 0.5,  
  mutation.rate = 0.05,  
  tune.threshold = FALSE,  
  tune.threshold.args = list(),  
  log.fun = "default"  
)
```

```
makeFeatSelControlRandom(  
  same.resampling.instance = TRUE,  
  maxit = 100L,  
  max.features = NA_integer_,  
  prob = 0.5,  
  tune.threshold = FALSE,  
  tune.threshold.args = list(),  
  log.fun = "default"  
)
```

```
makeFeatSelControlSequential(  
  same.resampling.instance = TRUE,  
  impute.val = NULL,  
  method,  
  alpha = 0.01,  
  beta = -0.001,  
  maxit = NA_integer_,  
  max.features = NA_integer_,  
  tune.threshold = FALSE,  
  tune.threshold.args = list(),
```

```

    log.fun = "default"
  )

```

Arguments

same.resampling.instance	(logical(1)) Should the same resampling instance be used for all evaluations to reduce variance? Default is TRUE.
maxit	(integer(1)) Maximal number of iterations. Note, that this is usually not equal to the number of function evaluations.
max.features	(integer(1)) Maximal number of features.
tune.threshold	(logical(1)) Should the threshold be tuned for the measure at hand, after each feature set evaluation, via tuneThreshold ? Only works for classification if the predict type is "prob". Default is FALSE.
tune.threshold.args	(list) Further arguments for threshold tuning that are passed down to tuneThreshold . Default is none.
log.fun	(function character(1)) Function used for logging. If set to "default" (the default), the evaluated design points, the resulting performances, and the runtime will be reported. If set to "memory" the memory usage for each evaluation will also be displayed, with character(1) small increase in run time. Otherwise character(1) function with arguments learner, resampling, measures, par.set, control, opt.path, dob, x, y, remove.nas, stage and prev.stage is expected. The default displays the performance measures, the time needed for evaluating, the currently used memory and the max memory ever used before (the latter two both taken from gc). See the implementation for details.
impute.val	(numeric) If something goes wrong during optimization (e.g. the learner crashes), this value is fed back to the tuner, so the tuning algorithm does not abort. It is not stored in the optimization path, an NA and a corresponding error message are logged instead. Note that this value is later multiplied by -1 for maximization measures internally, so you need to enter a larger positive value for maximization here as well. Default is the worst obtainable value of the performance measure you optimize for when you aggregate by mean value, or Inf instead. For multi-criteria optimization pass a vector of imputation values, one for each of your measures, in the same order as your measures.
comma	(logical(1)) Parameter of the GA feature selection, indicating whether to use a (mu, lambda) or (mu + lambda) GA. The default is FALSE.
mu	(integer(1)) Parameter of the GA feature selection. Size of the parent population.

lambda	(integer(1)) Parameter of the GA feature selection. Size of the children population (should be smaller or equal to mu).
crossover.rate	(numeric(1)) Parameter of the GA feature selection. Probability of choosing a bit from the first parent within the crossover mutation.
mutation.rate	(numeric(1)) Parameter of the GA feature selection. Probability of flipping a feature bit, i.e. switch between selecting / deselecting a feature.
prob	(numeric(1)) Parameter of the random feature selection. Probability of choosing a feature.
method	(character(1)) Parameter of the sequential feature selection. A character representing the method. Possible values are sfs (forward search), sbs (backward search), sfbs (floating forward search) and sfbs (floating backward search).
alpha	(numeric(1)) Parameter of the sequential feature selection. Minimal required value of improvement difference for a forward / adding step. Default is 0.01.
beta	(numeric(1)) Parameter of the sequential feature selection. Minimal required value of improvement difference for a backward / removing step. Negative values imply that you allow a slight decrease for the removal of a feature. Default is -0.001.

Value

(FeatSelControl). The specific subclass is one of [FeatSelControlExhaustive](#), [FeatSelControlRandom](#), [FeatSelControlSequential](#), [FeatSelControlGA](#).

References

Ron Kohavi and George H. John, Wrappers for feature subset selection, Artificial Intelligence Volume 97, 1997, 273-324. <http://ai.stanford.edu/~ronnyk/wrappersPrint.pdf>.

See Also

Other featsel: [analyzeFeatSelResult\(\)](#), [getFeatSelResult\(\)](#), [makeFeatSelWrapper\(\)](#), [selectFeatures\(\)](#)

FeatSelResult	<i>Result of feature selection.</i>
---------------	-------------------------------------

Description

Container for results of feature selection. Contains the obtained features, their performance values and the optimization path which lead there.

You can visualize it using [analyzeFeatSelResult](#).

Details

Object members:

learner (**Learner**) Learner that was optimized.

control (**FeatSelControl**) Control object from feature selection.

x (**character**) Vector of feature names identified as optimal.

y (**numeric**) Performance values for optimal x.

threshold (**numeric**) Vector of finally found and used thresholds if `tune.threshold` was enabled in **FeatSelControl**, otherwise not present and hence `NULL`.

opt.path (**ParamHelpers::OptPath**) Optimization path which lead to x.

filterFeatures	<i>Filter features by thresholding filter values.</i>
----------------	---

Description

First, calls [generateFilterValuesData](#). Features are then selected via `select` and `val`.

Usage

```
filterFeatures(
  task,
  method = "randomForestSRC_importance",
  fval = NULL,
  perc = NULL,
  abs = NULL,
  threshold = NULL,
  fun = NULL,
  fun.args = NULL,
  mandatory.feats = NULL,
  select.method = NULL,
  base.methods = NULL,
  cache = FALSE,
  ...
)
```

Arguments

task	(Task) The task.
method	(<code>character(1)</code>) See listFilterMethods . Default is “randomForestSRC_importance”.
fval	(FilterValues) Result of generateFilterValuesData . If you pass this, the filter values in the object are used for feature filtering. <code>method</code> and <code>...</code> are ignored then. Default is <code>NULL</code> and not used.

perc	(numeric(1)) If set, select perc*100 top scoring features. perc = 1 means to select all features. Mutually exclusive with arguments abs, threshold and fun.
abs	(numeric(1)) If set, select abs top scoring features. Mutually exclusive with arguments perc, threshold and fun.
threshold	(numeric(1)) If set, select features whose score exceeds threshold. Mutually exclusive with arguments perc, abs and fun.
fun	(function) If set, select features via a custom thresholding function, which must return the number of top scoring features to select. Mutually exclusive with arguments perc, abs and threshold.
fun.args	(any) Arguments passed to the custom thresholding function.
mandatory.feats	(character) Mandatory features which are always included regardless of their scores
select.method	If multiple methods are supplied in argument method, specify the method that is used for the final subsetting.
base.methods	If method is an ensemble filter, specify the base filter methods which the ensemble method will use.
cache	(character(1) logical) Whether to use caching during filter value creation. See details.
...	(any) Passed down to selected filter method.

Value[Task.](#)**Caching**

If cache = TRUE, the default mlr cache directory is used to cache filter values. The directory is operating system dependent and can be checked with `getCacheDir()`.

The default cache can be cleared with `deleteCacheDir()`. Alternatively, a custom directory can be passed to store the cache.

Note that caching is not thread safe. It will work for parallel computation on many systems, but there is no guarantee.

Simple and ensemble filters

Besides passing (multiple) simple filter methods you can also pass an ensemble filter method (in a list). The ensemble method will use the simple methods to calculate its ranking. See `listFilterEnsembleMethods()` for available ensemble methods.

See Also

Other filter: [generateFilterValuesData\(\)](#), [getFilteredFeatures\(\)](#), [listFilterEnsembleMethods\(\)](#), [listFilterMethods\(\)](#), [makeFilterEnsemble\(\)](#), [makeFilterWrapper\(\)](#), [makeFilter\(\)](#), [plotFilterValues\(\)](#)

Examples

```
# simple filter
filterFeatures(iris.task, method = "FSelectorRcpp_gain.ratio", abs = 2)
# ensemble filter
filterFeatures(iris.task, method = "E-min",
  base.methods = c("FSelectorRcpp_gain.ratio",
    "FSelectorRcpp_information.gain"), abs = 2)
```

```
friedmanPostHocTestBMR
```

Perform a posthoc Friedman-Nemenyi test.

Description

Performs a [PMCMR::posthoc.friedman.nemenyi.test](#) for a [BenchmarkResult](#) and a selected measure.

This means *all pairwise comparisons* of learners are performed. The null hypothesis of the post hoc test is that each pair of learners is equal. If the null hypothesis of the included ad hoc [stats::friedman.test](#) can be rejected an object of class `pairwise.htest` is returned. If not, the function returns the corresponding [friedman.test](#).

Note that benchmark results for at least two learners on at least two tasks are required.

Usage

```
friedmanPostHocTestBMR(
  bmr,
  measure = NULL,
  p.value = 0.05,
  aggregation = "default"
)
```

Arguments

bmr	(BenchmarkResult) Benchmark result.
measure	(Measure) Performance measure. Default is the first measure used in the benchmark experiment.
p.value	(<code>numeric(1)</code>) p-value for the tests. Default: 0.05
aggregation	(<code>character(1)</code>) “mean” or “default”. See getBMRAggrPerformances for details on “default”.

Value

(pairwise.htest): See [PMCMR::posthoc.friedman.nemenyi.test](#) for details. Additionally two components are added to the list:

- `f.rejnull` (`logical(1)`):
Whether the according `friedman.test` rejects the Null hypothesis at the selected p.value
- `crit.difference` (`list(2)`):
Minimal difference the mean ranks of two learners need to have in order to be significantly different

See Also

Other benchmark: [BenchmarkResult](#), [batchmark\(\)](#), [benchmark\(\)](#), [convertBMRToRankMatrix\(\)](#), [friedmanTestBMR\(\)](#), [generateCritDifferencesData\(\)](#), [getBMRAggrPerformances\(\)](#), [getBMRFeatSelResults\(\)](#), [getBMRFilteredFeatures\(\)](#), [getBMRLearnerIds\(\)](#), [getBMRLearnerShortNames\(\)](#), [getBMRLearners\(\)](#), [getBMRMeasureIds\(\)](#), [getBMRMeasures\(\)](#), [getBMRModels\(\)](#), [getBMRPerformances\(\)](#), [getBMRPredictions\(\)](#), [getBMRTaskDescs\(\)](#), [getBMRTaskIds\(\)](#), [getBMRTuneResults\(\)](#), [plotBMRBoxplots\(\)](#), [plotBMRRanksAsBarChart\(\)](#), [plotBMRSummary\(\)](#), [plotCritDifferences\(\)](#), [reduceBatchmarkResults\(\)](#)

Examples

```
# see benchmark
```

<code>friedmanTestBMR</code>	<i>Perform overall Friedman test for a <code>BenchmarkResult</code>.</i>
------------------------------	--

Description

Performs a [stats::friedman.test](#) for a selected measure. The null hypothesis is that apart from an effect of the different (**Task**), the location parameter (aggregated performance measure) is the same for each **Learner**. Note that benchmark results for at least two learners on at least two tasks are required.

Usage

```
friedmanTestBMR(bmr, measure = NULL, aggregation = "default")
```

Arguments

<code>bmr</code>	(BenchmarkResult) Benchmark result.
<code>measure</code>	(Measure) Performance measure. Default is the first measure used in the benchmark experiment.
<code>aggregation</code>	(character(1)) “mean” or “default”. See getBMRAggrPerformances for details on “default”.

Value

(h_{test}): See [stats::friedman.test](#) for details.

See Also

Other benchmark: [BenchmarkResult](#), [batchmark\(\)](#), [benchmark\(\)](#), [convertBMRTToRankMatrix\(\)](#), [friedmanPostHocTestBMR\(\)](#), [generateCritDifferencesData\(\)](#), [getBMRAggrPerformances\(\)](#), [getBMRFeatSelResults\(\)](#), [getBMRFilteredFeatures\(\)](#), [getBMRLearnerIds\(\)](#), [getBMRLearnerShortNames\(\)](#), [getBMRLearners\(\)](#), [getBMRMeasureIds\(\)](#), [getBMRMeasures\(\)](#), [getBMRModels\(\)](#), [getBMRPerformances\(\)](#), [getBMRPredictions\(\)](#), [getBMRTaskDescs\(\)](#), [getBMRTaskIds\(\)](#), [getBMRTuneResults\(\)](#), [plotBMRBoxplots\(\)](#), [plotBMRRanksAsBarChart\(\)](#), [plotBMRSummary\(\)](#), [plotCritDifferences\(\)](#), [reduceBatchmarkResults\(\)](#)

Examples

```
# see benchmark
```

fuelsubset.task	<i>FuelSubset functional data regression task.</i>
-----------------	--

Description

Contains the task (`fuelsubset.task`). 2 functional covariates and 1 scalar covariate. You have to predict the heat value of some fuel based on the ultraviolet radiation spectrum and infrared ray radiation and one scalar column called `h2o`.

Details

The features and grids are scaled in the same way as in [FDboost::FDboost](#).

References

See Brockhaus, S., Scheipl, F., Hothorn, T., & Greven, S. (2015). The functional linear array model. *Statistical Modelling*, 15(3), 279–300.

<code>generateCalibrationData</code>	<i>Generate classifier calibration data.</i>
--------------------------------------	--

Description

A calibrated classifier is one where the predicted probability of a class closely matches the rate at which that class occurs, e.g. for data points which are assigned a predicted probability of class A of .8, approximately 80 percent of such points should belong to class A if the classifier is well calibrated. This is estimated empirically by grouping data points with similar predicted probabilities for each class, and plotting the rate of each class within each bin against the predicted probability bins.

Usage

```
generateCalibrationData(obj, breaks = "Sturges", groups = NULL, task.id = NULL)
```

Arguments

obj	(list of Prediction list of ResampleResult BenchmarkResult) Single prediction object, list of them, single resample result, list of them, or a benchmark result. In case of a list probably produced by different learners you want to compare, then name the list with the names you want to see in the plots, probably learner shortnames or ids.
breaks	(character(1) numeric) If character(1), the algorithm to use in generating probability bins. See hist for details. If numeric , the cut points for the bins. Default is "Sturges".
groups	(integer(1)) The number of bins to construct. If specified, breaks is ignored. Default is NULL.
task.id	(character(1)) Selected task in BenchmarkResult to do plots for, ignored otherwise. Default is first task.

Value

[CalibrationData](#). A list containing:

proportion	data.frame with columns: <ul style="list-style-type: none"> • Learner Name of learner. • bin Bins calculated according to the breaks or groups argument. • Class Class labels (for binary classification only the positive class). • Proportion Proportion of observations from class Class among all observations with posterior probabilities of class Class within the interval given in bin.
data	data.frame with columns: <ul style="list-style-type: none"> • Learner Name of learner. • truth True class label. • Class Class labels (for binary classification only the positive class). • Probability Predicted posterior probability of Class. • bin Bin corresponding to Probability.
task	(TaskDesc) Task description.

References

Vuk, Miha, and Curk, Tomaz. "ROC Curve, Lift Chart, and Calibration Plot." *Metodoloski zvezki*. Vol. 3. No. 1 (2006): 89-108.

See Also

Other generate_plot_data: [generateCritDifferencesData\(\)](#), [generateFeatureImportanceData\(\)](#), [generateFilterValuesData\(\)](#), [generateLearningCurveData\(\)](#), [generatePartialDependenceData\(\)](#), [generateThreshVsPerfData\(\)](#), [plotFilterValues\(\)](#)

Other calibration: [plotCalibration\(\)](#)

generateCritDifferencesData

Generate data for critical-differences plot.

Description

Generates data that can be used to plot a critical differences plot. Computes the critical differences according to either the "Bonferroni-Dunn" test or the "Nemenyi" test.

"Bonferroni-Dunn" usually yields higher power as it does not compare all algorithms to each other, but all algorithms to a baseline instead.

Learners are drawn on the y-axis according to their average rank.

For test = "nemenyi" a bar is drawn, connecting all groups of not significantly different learners.

For test = "bd" an interval is drawn around the algorithm selected as a baseline. All learners within this interval are not significantly different from the baseline.

Calculation:

$$CD = q_{\alpha} \sqrt{\left(\frac{k(k+1)}{6N}\right)}$$

Where q_{α} is based on the studentized range statistic. See references for details.

Usage

```
generateCritDifferencesData(
  bmr,
  measure = NULL,
  p.value = 0.05,
  baseline = NULL,
  test = "bd"
)
```

Arguments

bmr	(BenchmarkResult) Benchmark result.
measure	(Measure) Performance measure. Default is the first measure used in the benchmark experiment.
p.value	(numeric(1)) P-value for the critical difference. Default: 0.05

baseline	(character(1)): (learner.id) Select a learner.id as baseline for the test = "bd" ("Bonferroni-Dunn") critical differences diagram. The critical difference interval will then be positioned around this learner. Defaults to best performing algorithm. For test = "nemenyi", no baseline is needed as it performs <i>all pairwise comparisons</i> .
test	(character(1)) Test for which the critical differences are computed. "bd" for the Bonferroni-Dunn Test, which is comparing all classifiers to a baseline, thus performing a comparison of one classifier to all others. Algorithms not connected by a single line are statistically different from the baseline. "nemenyi" for the PMCMR::posthoc.friedman.nemenyi.test which is comparing all classifiers to each other. The null hypothesis that there is a difference between the classifiers can not be rejected for all classifiers that have a single grey bar connecting them.

Value

(critDifferencesData). List containing:

data	(data.frame) containing the info for the descriptive part of the plot
friedman.nemenyi.test	(list) of class pairwise.htest contains the calculated PMCMR::posthoc.friedman.nemenyi.test
cd.info	(list) containing info on the critical difference and its positioning
baseline	baseline chosen for plotting
p.value	p.value used for the PMCMR::posthoc.friedman.nemenyi.test and for computation of the critical difference

See Also

Other generate_plot_data: [generateCalibrationData\(\)](#), [generateFeatureImportanceData\(\)](#), [generateFilterValuesData\(\)](#), [generateLearningCurveData\(\)](#), [generatePartialDependenceData\(\)](#), [generateThreshVsPerfData\(\)](#), [plotFilterValues\(\)](#)

Other benchmark: [BenchmarkResult](#), [batchmark\(\)](#), [benchmark\(\)](#), [convertBMRTToRankMatrix\(\)](#), [friedmanPostHocTestBMR\(\)](#), [friedmanTestBMR\(\)](#), [getBMRAggrPerformances\(\)](#), [getBMRFeatSelResults\(\)](#), [getBMRFilteredFeatures\(\)](#), [getBMRLearnerIds\(\)](#), [getBMRLearnerShortNames\(\)](#), [getBMRLearners\(\)](#), [getBMRMeasureIds\(\)](#), [getBMRMeasures\(\)](#), [getBMRModels\(\)](#), [getBMRPerformances\(\)](#), [getBMRPredictions\(\)](#), [getBMRTaskDescs\(\)](#), [getBMRTaskIds\(\)](#), [getBMRTuneResults\(\)](#), [plotBMRBoxplots\(\)](#), [plotBMRRanksAsBarChart\(\)](#), [plotBMRSummary\(\)](#), [plotCritDifferences\(\)](#), [reduceBatchmarkResults\(\)](#)

```
generateFeatureImportanceData
```

Generate feature importance.

Description

Estimate how important individual features or groups of features are by contrasting prediction performances. For method “permutation.importance” compute the change in performance from permuting the values of a feature (or a group of features) and compare that to the predictions made on the unmcuted data.

Usage

```
generateFeatureImportanceData(  
  task,  
  method = "permutation.importance",  
  learner,  
  features = getTaskFeatureNames(task),  
  interaction = FALSE,  
  measure,  
  contrast = function(x, y) x - y,  
  aggregation = mean,  
  nmc = 50L,  
  replace = TRUE,  
  local = FALSE,  
  show.info = FALSE  
)
```

Arguments

task	(Task) The task.
method	(character(1)) The method used to compute the feature importance. The only method available is “permutation.importance”. Default is “permutation.importance”.
learner	(Learner character(1)) The learner. If you pass a string the learner will be created via makeLearner .
features	(character) The features to compute the importance of. The default is all of the features contained in the Task .
interaction	(logical(1)) Whether to compute the importance of the features argument jointly. For method = “permutation.importance” this entails permuting the values of all features together and then contrasting the performance with that of the performance without the features being permuted. The default is FALSE.

measure	(Measure) Performance measure. Default is the first measure used in the benchmark experiment.
contrast	(function) A difference function that takes a numeric vector and returns a numeric vector of the same length. The default is element-wise difference between the vectors.
aggregation	(function) A function which aggregates the differences. This function must take a numeric vector and return a numeric vector of length 1. The default is mean.
nmc	(integer(1)) The number of Monte-Carlo iterations to use in computing the feature importance. If nmc == -1 and method = "permutation.importance" then all permutations of the features are used. The default is 50.
replace	(logical(1)) Whether or not to sample the feature values with or without replacement. The default is TRUE.
local	(logical(1)) Whether to compute the per-observation importance. The default is FALSE.
show.info	(logical(1)) Whether progress output (feature name, time elapsed) should be displayed.

Value

(FeatureImportance). A named list which contains the computed feature importance and the input arguments.

Object members:

res	(data.frame) Has columns for each feature or combination of features (colon separated) for which the importance is computed. A row corresponds to importance of the feature specified in the column for the target.
interaction	(logical(1)) Whether or not the importance of the features was computed jointly rather than individually.
measure	(Measure)

The measure used to compute performance.

contrast	(function) The function used to compare the performance of predictions.
aggregation	(function) The function which is used to aggregate the contrast between the performance of predictions across Monte-Carlo iterations.
replace	(logical(1)) Whether or not, when method = "permutation.importance", the feature values are sampled with replacement.

nmc	(integer(1)) The number of Monte-Carlo iterations used to compute the feature importance. When nmc == -1 and method = "permutation.importance" all permutations are used.
local	(logical(1)) Whether observation-specific importance is computed for the features.

References

Jerome Friedman; Greedy Function Approximation: A Gradient Boosting Machine, Annals of Statistics, Vol. 29, No. 5 (Oct., 2001), pp. 1189-1232.

See Also

Other generate_plot_data: [generateCalibrationData\(\)](#), [generateCritDifferencesData\(\)](#), [generateFilterValuesData\(\)](#), [generateLearningCurveData\(\)](#), [generatePartialDependenceData\(\)](#), [generateThreshVsPerfData\(\)](#), [plotFilterValues\(\)](#)

Examples

```
lrn = makeLearner("classif.rpart", predict.type = "prob")
fit = train(lrn, iris.task)
imp = generateFeatureImportanceData(iris.task, "permutation.importance",
  lrn, "Petal.Width", nmc = 10L, local = TRUE)
```

generateFilterValuesData

Calculates feature filter values.

Description

Calculates numerical filter values for features. For a list of features, use [listFilterMethods](#).

Usage

```
generateFilterValuesData(
  task,
  method = "randomForestSRC_importance",
  nselect = getTaskNFeats(task),
  ...,
  more.args = list()
)
```

Arguments

task	(Task) The task.
method	(character list) Filter method(s). In case of ensemble filters the <code>list</code> notation needs to be used. See the examples for more information. Default is “randomForestSRC_importance”.
nselect	(integer(1)) Number of scores to request. Scores are getting calculated for all features per default.
...	(any) Passed down to selected method. Can only be use if method contains one element.
more.args	(named list) Extra args passed down to filter methods. List elements are named with the filter method name the args should be passed down to. A more general and flexible option than ... Default is empty list.

Value

([FilterValues](#)). A list containing:

task.desc	[TaskDesc] Task description.
data	(data.frame) with columns: <ul style="list-style-type: none"> • name(character) Name of feature. • type(character) Feature column type. • method(numeric) One column for each method with the feature importance values.

Simple and ensemble filters

Besides passing (multiple) simple filter methods you can also pass an ensemble filter method (in a list). The ensemble method will use the simple methods to calculate its ranking. See `listFilterEnsembleMethods()` for available ensemble methods.

See Also

Other generate_plot_data: [generateCalibrationData\(\)](#), [generateCritDifferencesData\(\)](#), [generateFeatureImportanceData\(\)](#), [generateLearningCurveData\(\)](#), [generatePartialDependenceData\(\)](#), [generateThreshVsPerfData\(\)](#), [plotFilterValues\(\)](#)

Other filter: [filterFeatures\(\)](#), [getFilteredFeatures\(\)](#), [listFilterEnsembleMethods\(\)](#), [listFilterMethods\(\)](#), [makeFilterEnsemble\(\)](#), [makeFilterWrapper\(\)](#), [makeFilter\(\)](#), [plotFilterValues\(\)](#)

Examples

```
# two simple filter methods
fval = generateFilterValuesData(iris.task,
  method = c("FSelectorRcpp_gain.ratio", "FSelectorRcpp_information.gain"))
# using ensemble method "E-mean"
fval = generateFilterValuesData(iris.task,
  method = list("E-mean", c("FSelectorRcpp_gain.ratio",
    "FSelectorRcpp_information.gain")))
```

```
generateHyperParsEffectData
```

Generate hyperparameter effect data.

Description

Generate cleaned hyperparameter effect data from a tuning result or from a nested cross-validation tuning result. The object returned can be used for custom visualization or passed downstream to an out of the box mlr method, [plotHyperParsEffect](#).

Usage

```
generateHyperParsEffectData(
  tune.result,
  include.diagnostics = FALSE,
  trafo = FALSE,
  partial.dep = FALSE
)
```

Arguments

tune.result	(TuneResult ResampleResult) Result of tuneParams (or resample ONLY when used for nested cross-validation). The tuning result (or results if the output is from nested cross-validation), also containing the optimizer results. If nested CV output is passed, each element in the list will be considered a separate run, and the data from each run will be included in the dataframe within the returned HyperParsEffectData .
include.diagnostics	(logical(1)) Should diagnostic info (eol and error msg) be included? Default is FALSE.
trafo	(logical(1)) Should the units of the hyperparameter path be converted to the transformed scale? This is only useful when trafo was used to create the path. Default is FALSE.
partial.dep	(logical(1)) Should partial dependence be requested based on converting to reg task? This sets a flag so that we know to use partial dependence downstream. This should

most likely be set to TRUE if 2 or more hyperparameters were tuned simultaneously. Partial dependence should always be requested when more than 2 hyperparameters were tuned simultaneously. Setting to TRUE will cause `plotHyperParsEffect` to automatically plot partial dependence when called downstream. Default is FALSE.

Value

(HyperParsEffectData) Object containing the hyperparameter effects dataframe, the tuning performance measures used, the hyperparameters used, a flag for including diagnostic info, a flag for whether nested cv was used, a flag for whether partial dependence should be generated, and the optimization algorithm used.

Examples

```
## Not run:
# 3-fold cross validation
ps = makeParamSet(makeDiscreteParam("C", values = 2^(-4:4)))
ctrl = makeTuneControlGrid()
rdesc = makeResampleDesc("CV", iters = 3L)
res = tuneParams("classif.ksvm", task = pid.task, resampling = rdesc,
  par.set = ps, control = ctrl)
data = generateHyperParsEffectData(res)
plt = plotHyperParsEffect(data, x = "C", y = "mmce.test.mean")
plt + ylab("Misclassification Error")

# nested cross validation
ps = makeParamSet(makeDiscreteParam("C", values = 2^(-4:4)))
ctrl = makeTuneControlGrid()
rdesc = makeResampleDesc("CV", iters = 3L)
lrn = makeTuneWrapper("classif.ksvm", control = ctrl,
  resampling = rdesc, par.set = ps)
res = resample(lrn, task = pid.task, resampling = cv2,
  extract = getTuneResult)
data = generateHyperParsEffectData(res)
plotHyperParsEffect(data, x = "C", y = "mmce.test.mean", plot.type = "line")

## End(Not run)
```

```
generateLearningCurveData
```

Generates a learning curve.

Description

Observe how the performance changes with an increasing number of observations.

Usage

```
generateLearningCurveData(
  learners,
  task,
  resampling = NULL,
  percs = seq(0.1, 1, by = 0.1),
  measures,
  stratify = FALSE,
  show.info = getMlrOption("show.info")
)
```

Arguments

learners	[(list of) Learner] Learning algorithms which should be compared.
task	(Task) The task.
resampling	(ResampleDesc ResampleInstance) Resampling strategy to evaluate the performance measure. If no strategy is given a default "Holdout" will be performed.
percs	(numeric) Vector of percentages to be drawn from the training split. These values represent the x-axis. Internally makeDownsampleWrapper is used in combination with benchmark . Thus for each percentage a different set of observations is drawn resulting in noisy performance measures as the quality of the sample can differ.
measures	[(list of) Measure] Performance measures to generate learning curves for, representing the y-axis.
stratify	(logical(1)) Only for classification: Should the downsampled data be stratified according to the target classes?
show.info	(logical(1)) Print verbose output on console? Default is set via configureMlr .

Value

([LearningCurveData](#)). A list containing:

- The [Task](#)
- List of [Measure](#)
Performance measures
- data ([data.frame](#)) with columns:
 - learner Names of learners.
 - percentage Percentages drawn from the training split.
 - One column for each [Measure](#) passed to [generateLearningCurveData](#).

See Also

Other generate_plot_data: [generateCalibrationData\(\)](#), [generateCritDifferencesData\(\)](#), [generateFeatureImportanceData\(\)](#), [generateFilterValuesData\(\)](#), [generatePartialDependenceData\(\)](#), [generateThreshVsPerfData\(\)](#), [plotFilterValues\(\)](#)

Other learning_curve: [plotLearningCurve\(\)](#)

Examples

```
r = generateLearningCurveData(list("classif.rpart", "classif.knn"),
  task = sonar.task, percs = seq(0.2, 1, by = 0.2),
  measures = list(tp, fp, tn, fn),
  resampling = makeResampleDesc(method = "Subsample", iters = 5),
  show.info = FALSE)
plotLearningCurve(r)
```

```
generatePartialDependenceData
```

Generate partial dependence.

Description

Estimate how the learned prediction function is affected by one or more features. For a learned function $f(x)$ where x is partitioned into x_s and x_c , the partial dependence of f on x_s can be summarized by averaging over x_c and setting x_s to a range of values of interest, estimating $E_{x_c}(f(x_s, x_c))$. The conditional expectation of f at observation i is estimated similarly. Additionally, partial derivatives of the marginalized function w.r.t. the features can be computed.

Usage

```
generatePartialDependenceData(
  obj,
  input,
  features = NULL,
  interaction = FALSE,
  derivative = FALSE,
  individual = FALSE,
  fun = mean,
  bounds = c(qnorm(0.025), qnorm(0.975)),
  uniform = TRUE,
  n = c(10, NA),
  ...
)
```

Arguments

obj	(WrappedModel) Result of <code>train</code> .
input	(data.frame Task) Input data.
features	character A vector of feature names contained in the training data. If not specified all features in the input will be used.
interaction	(<code>logical(1)</code>) Whether the features should be interacted or not. If TRUE then the Cartesian product of the prediction grid for each feature is taken, and the partial dependence at each unique combination of values of the features is estimated. Note that if the length of features is greater than two, plotPartialDependence cannot be used. If FALSE each feature is considered separately. In this case features can be much longer than two. Default is FALSE.
derivative	(<code>logical(1)</code>) Whether or not the partial derivative of the learned function with respect to the features should be estimated. If TRUE <code>interaction</code> must be FALSE. The partial derivative of individual observations may be estimated. Note that computation time increases as the learned prediction function is evaluated at <code>gridsize</code> points * the number of points required to estimate the partial derivative. Additional arguments may be passed to numDeriv::grad (for regression or survival tasks) or numDeriv::jacobian (for classification tasks). Note that functions which are not smooth may result in estimated derivatives of 0 (for points where the function does not change within +/- epsilon) or estimates trending towards +/- infinity (at discontinuities). Default is FALSE.
individual	(<code>logical(1)</code>) Whether to plot the individual conditional expectation curves rather than the aggregated curve, i.e., rather than aggregating (using <code>fun</code>) the partial dependences of features, plot the partial dependences of all observations in data across all values of the features. The algorithm is developed in Goldstein, Kapelner, Bleich, and Pitkin (2015). Default is FALSE.
fun	function A function which operates on the output on the predictions made on the input data. For regression this means a numeric vector, and, e.g., for a multiclass classification problem, this might instead be probabilities which are returned as a numeric matrix. This argument can return vectors of arbitrary length, however, if their length is greater than one, they must be named, e.g., <code>fun = mean</code> or <code>fun = function(x) c("mean" = mean(x), "variance" = var(x))</code> . The default is the mean, unless <code>obj</code> is classification with <code>predict.type = "response"</code> in which case the default is the proportion of observations predicted to be in each class.
bounds	(<code>numeric(2)</code>) The value (lower, upper) the estimated standard error is multiplied by to estimate the bound on a confidence region for a partial dependence. Ignored if

	predict.type != "se" for the learner. Default is the 2.5 and 97.5 quantiles (-1.96, 1.96) of the Gaussian distribution.
uniform	(logical(1)) Whether or not the prediction grid for the features is a uniform grid of size n[1] or sampled with replacement from the input. Default is TRUE.
n	(integer21) The first element of n gives the size of the prediction grid created for each feature. The second element of n gives the size of the sample to be drawn without replacement from the input data. Setting n[2] less than the number of rows in the input will decrease computation time. The default for n[1] is 10, and the default for n[2] is the number of rows in the input.
...	additional arguments to be passed to <code>mmpf::marginalPrediction</code> .

Value

PartialDependenceData. A named list, which contains the partial dependence, input data, target, features, task description, and other arguments controlling the type of partial dependences made.

Object members:

data	data.frame Has columns for the prediction: one column for regression and survival analysis, and a column for class and the predicted probability for classification as well as a column for each element of features. If individual = TRUE then there is an additional column idx which gives the index of the data that each prediction corresponds to.
task.desc	TaskDesc Task description.
target	Target feature for regression, target feature levels for classification, survival and event indicator for survival.
features	character Features argument input.
interaction	(logical(1)) Whether or not the features were interacted (i.e. conditioning).
derivative	(logical(1)) Whether or not the partial derivative was estimated.
individual	(logical(1)) Whether the partial dependences were aggregated or the individual curves are retained.

References

Goldstein, Alex, Adam Kapelner, Justin Bleich, and Emil Pitkin. "Peeking inside the black box: Visualizing statistical learning with plots of individual conditional expectation." *Journal of Computational and Graphical Statistics*. Vol. 24, No. 1 (2015): 44-65.

Friedman, Jerome. "Greedy Function Approximation: A Gradient Boosting Machine." *The Annals of Statistics*. Vol. 29, No. 5 (2001): 1189-1232.

See Also

Other partial_dependence: [plotPartialDependence\(\)](#)

Other generate_plot_data: [generateCalibrationData\(\)](#), [generateCritDifferencesData\(\)](#), [generateFeatureImportanceData\(\)](#), [generateFilterValuesData\(\)](#), [generateLearningCurveData\(\)](#), [generateThreshVsPerfData\(\)](#), [plotFilterValues\(\)](#)

Examples

```
lrn = makeLearner("regr.svm")
fit = train(lrn, bh.task)
pd = generatePartialDependenceData(fit, bh.task, "lstat")
plotPartialDependence(pd, data = getTaskData(bh.task))
```

```
lrn = makeLearner("classif.rpart", predict.type = "prob")
fit = train(lrn, iris.task)
pd = generatePartialDependenceData(fit, iris.task, "Petal.Width")
plotPartialDependence(pd, data = getTaskData(iris.task))
```

```
generateThreshVsPerfData
```

Generate threshold vs. performance(s) for 2-class classification.

Description

Generates data on threshold vs. performance(s) for 2-class classification that can be used for plotting.

Usage

```
generateThreshVsPerfData(
  obj,
  measures,
  gridsize = 100L,
  aggregate = TRUE,
  task.id = NULL
)
```

Arguments

obj	(list of Prediction list of ResampleResult BenchmarkResult) Single prediction object, list of them, single resample result, list of them, or a benchmark result. In case of a list probably produced by different learners you want to compare, then name the list with the names you want to see in the plots, probably learner shortnames or ids.
measures	(Measure list of Measure) Performance measure(s) to evaluate. Default is the default measure for the task, see here getDefaultMeasure .

gridsize	(integer(1)) Grid resolution for x-axis (threshold). Default is 100.
aggregate	(logical(1)) Whether to aggregate ResamplePredictions or to plot the performance of each iteration separately. Default is TRUE.
task.id	(character(1)) Selected task in BenchmarkResult to do plots for, ignored otherwise. Default is first task.

Value

([ThreshVsPerfData](#)). A named list containing the measured performance across the threshold grid, the measures, and whether the performance estimates were aggregated (only applicable for (list of) [ResampleResults](#)).

See Also

Other generate_plot_data: [generateCalibrationData\(\)](#), [generateCritDifferencesData\(\)](#), [generateFeatureImportanceData\(\)](#), [generateFilterValuesData\(\)](#), [generateLearningCurveData\(\)](#), [generatePartialDependenceData\(\)](#), [plotFilterValues\(\)](#)

Other thresh_vs_perf: [plotROCCurves\(\)](#), [plotThreshVsPerf\(\)](#)

```
getBMRAggrPerformances
```

Extract the aggregated performance values from a benchmark result.

Description

Either a list of lists of “aggr” numeric vectors, as returned by [resample](#), or these objects are rbind-ed with extra columns “task.id” and “learner.id”.

Usage

```
getBMRAggrPerformances(
  bmr,
  task.ids = NULL,
  learner.ids = NULL,
  as.df = FALSE,
  drop = FALSE
)
```

Arguments

bmr	(BenchmarkResult) Benchmark result.
task.ids	(character(1)) Restrict result to certain tasks. Default is all.

learner.ids	(character(1)) Restrict result to certain learners. Default is all.
as.df	(character(1)) Return one data.frame as result - or a list of lists of objects?. Default is FALSE.
drop	(logical(1)) If drop is FALSE (the default), a nested list with the following structure is returned: res[task.ids][learner.ids]. If drop is set to TRUE it is checked if the list structure can be simplified. If only one learner was passed, a list with entries for each task is returned. If only one task was passed, the entries are named after the corresponding learner. For an experiment with both one task and learner, the whole list structure is removed. Note that the name of the task/learner will be dropped from the return object.

Value

(list | [data.frame](#)). See above.

See Also

Other benchmark: [BenchmarkResult](#), [batchmark\(\)](#), [benchmark\(\)](#), [convertBMRTToRankMatrix\(\)](#), [friedmanPostHocTestBMR\(\)](#), [friedmanTestBMR\(\)](#), [generateCritDifferencesData\(\)](#), [getBMRFeatSelResults\(\)](#), [getBMRFilteredFeatures\(\)](#), [getBMRLearnerIds\(\)](#), [getBMRLearnerShortNames\(\)](#), [getBMRLearners\(\)](#), [getBMRMeasureIds\(\)](#), [getBMRMeasures\(\)](#), [getBMRModels\(\)](#), [getBMRPerformances\(\)](#), [getBMRPredictions\(\)](#), [getBMRTaskDescs\(\)](#), [getBMRTaskIds\(\)](#), [getBMRTuneResults\(\)](#), [plotBMRBoxplots\(\)](#), [plotBMRRanksAsBarChart\(\)](#), [plotBMRSummary\(\)](#), [plotCritDifferences\(\)](#), [reduceBatchmarkResults\(\)](#)

getBMRFeatSelResults *Extract the feature selection results from a benchmark result.*

Description

Returns a nested list of [FeatSelResults](#). The first level of nesting is by data set, the second by learner, the third for the benchmark resampling iterations. If `as.df` is TRUE, a data frame with “task.id”, “learner.id”, the resample iteration and the selected features is returned.

Note that if more than one feature is selected and a data frame is requested, there will be multiple rows for the same dataset-learner-iteration; one for each selected feature.

Usage

```
getBMRFeatSelResults(
  bmr,
  task.ids = NULL,
  learner.ids = NULL,
```

```

    as.df = FALSE,
    drop = FALSE
  )

```

Arguments

bmr	(BenchmarkResult) Benchmark result.
task.ids	(character(1)) Restrict result to certain tasks. Default is all.
learner.ids	(character(1)) Restrict result to certain learners. Default is all.
as.df	(character(1)) Return one data.frame as result - or a list of lists of objects?. Default is FALSE.
drop	(logical(1)) If drop is FALSE (the default), a nested list with the following structure is returned: res[task.ids][learner.ids]. If drop is set to TRUE it is checked if the list structure can be simplified. If only one learner was passed, a list with entries for each task is returned. If only one task was passed, the entries are named after the corresponding learner. For an experiment with both one task and learner, the whole list structure is removed. Note that the name of the task/learner will be dropped from the return object.

Value

([list](#) | [data.frame](#)). See above.

See Also

Other benchmark: [BenchmarkResult](#), [batchmark\(\)](#), [benchmark\(\)](#), [convertBMRToRankMatrix\(\)](#), [friedmanPostHocTestBMR\(\)](#), [friedmanTestBMR\(\)](#), [generateCritDifferencesData\(\)](#), [getBMRAggrPerformances\(\)](#), [getBMRFilteredFeatures\(\)](#), [getBMRLearnerIds\(\)](#), [getBMRLearnerShortNames\(\)](#), [getBMRLearners\(\)](#), [getBMRMeasureIds\(\)](#), [getBMRMeasures\(\)](#), [getBMRModels\(\)](#), [getBMRPerformances\(\)](#), [getBMRPredictions\(\)](#), [getBMRTaskDescs\(\)](#), [getBMRTaskIds\(\)](#), [getBMRTuneResults\(\)](#), [plotBMRBoxplots\(\)](#), [plotBMRRanksAsBarChart\(\)](#), [plotBMRSummary\(\)](#), [plotCritDifferences\(\)](#), [reduceBatchmarkResults\(\)](#)

getBMRFilteredFeatures

Extract the feature selection results from a benchmark result.

Description

Returns a nested list of characters. The first level of nesting is by data set, the second by learner, the third for the benchmark resampling iterations. The list at the lowest level is the list of selected features. If `as.df` is TRUE, a data frame with “task.id”, “learner.id”, the resample iteration and the selected features is returned.

Note that if more than one feature is selected and a data frame is requested, there will be multiple rows for the same dataset-learner-iteration; one for each selected feature.

Usage

```
getBMRFilteredFeatures(
  bmr,
  task.ids = NULL,
  learner.ids = NULL,
  as.df = FALSE,
  drop = FALSE
)
```

Arguments

<code>bmr</code>	(BenchmarkResult) Benchmark result.
<code>task.ids</code>	(<code>character(1)</code>) Restrict result to certain tasks. Default is all.
<code>learner.ids</code>	(<code>character(1)</code>) Restrict result to certain learners. Default is all.
<code>as.df</code>	(<code>character(1)</code>) Return one data.frame as result - or a list of lists of objects?. Default is FALSE.
<code>drop</code>	(<code>logical(1)</code>) If <code>drop</code> is FALSE (the default), a nested list with the following structure is returned: <code>res[task.ids][learner.ids]</code> . If <code>drop</code> is set to TRUE it is checked if the list structure can be simplified. If only one learner was passed, a list with entries for each task is returned. If only one task was passed, the entries are named after the corresponding learner. For an experiment with both one task and learner, the whole list structure is removed. Note that the name of the task/learner will be dropped from the return object.

Value

([list](#) | [data.frame](#)). See above.

See Also

Other benchmark: [BenchmarkResult](#), [batchmark\(\)](#), [benchmark\(\)](#), [convertBMRToRankMatrix\(\)](#), [friedmanPostHocTestBMR\(\)](#), [friedmanTestBMR\(\)](#), [generateCritDifferencesData\(\)](#), [getBMRAggrPerformances\(\)](#),

[getBMRFeatSelResults\(\)](#), [getBMRLearnerIds\(\)](#), [getBMRLearnerShortNames\(\)](#), [getBMRLearners\(\)](#), [getBMRMeasureIds\(\)](#), [getBMRMeasures\(\)](#), [getBMRModels\(\)](#), [getBMRPerformances\(\)](#), [getBMRPredictions\(\)](#), [getBMRTaskDescs\(\)](#), [getBMRTaskIds\(\)](#), [getBMRTuneResults\(\)](#), [plotBMRBoxplots\(\)](#), [plotBMRRanksAsBarChart\(\)](#), [plotBMRSummary\(\)](#), [plotCritDifferences\(\)](#), [reduceBatchmarkResults\(\)](#)

getBMRLearnerIds *Return learner ids used in benchmark.*

Description

Gets the IDs of the learners used in a benchmark experiment.

Usage

```
getBMRLearnerIds(bmr)
```

Arguments

bmr ([BenchmarkResult](#))
Benchmark result.

Value

([character](#)).

See Also

Other benchmark: [BenchmarkResult](#), [batchmark\(\)](#), [benchmark\(\)](#), [convertBMRToRankMatrix\(\)](#), [friedmanPostHocTestBMR\(\)](#), [friedmanTestBMR\(\)](#), [generateCritDifferencesData\(\)](#), [getBMRAggrPerformances\(\)](#), [getBMRFeatSelResults\(\)](#), [getBMRFilteredFeatures\(\)](#), [getBMRLearnerShortNames\(\)](#), [getBMRLearners\(\)](#), [getBMRMeasureIds\(\)](#), [getBMRMeasures\(\)](#), [getBMRModels\(\)](#), [getBMRPerformances\(\)](#), [getBMRPredictions\(\)](#), [getBMRTaskDescs\(\)](#), [getBMRTaskIds\(\)](#), [getBMRTuneResults\(\)](#), [plotBMRBoxplots\(\)](#), [plotBMRRanksAsBarChart\(\)](#), [plotBMRSummary\(\)](#), [plotCritDifferences\(\)](#), [reduceBatchmarkResults\(\)](#)

getBMRLearners *Return learners used in benchmark.*

Description

Gets the learners used in a benchmark experiment.

Usage

```
getBMRLearners(bmr)
```

Arguments

bmr [\(BenchmarkResult\)](#)
Benchmark result.

Value

(list).

See Also

Other benchmark: [BenchmarkResult](#), [batchmark\(\)](#), [benchmark\(\)](#), [convertBMRTToRankMatrix\(\)](#), [friedmanPostHocTestBMR\(\)](#), [friedmanTestBMR\(\)](#), [generateCritDifferencesData\(\)](#), [getBMRAggrPerformances\(\)](#), [getBMRFeatSelResults\(\)](#), [getBMRFilteredFeatures\(\)](#), [getBMRLearnerIds\(\)](#), [getBMRLearnerShortNames\(\)](#), [getBMRMeasureIds\(\)](#), [getBMRMeasures\(\)](#), [getBMRModels\(\)](#), [getBMRPerformances\(\)](#), [getBMRPredictions\(\)](#), [getBMRTaskDescs\(\)](#), [getBMRTaskIds\(\)](#), [getBMRTuneResults\(\)](#), [plotBMRBoxplots\(\)](#), [plotBMRRanksAsBarChart\(\)](#), [plotBMRSummary\(\)](#), [plotCritDifferences\(\)](#), [reduceBatchmarkResults\(\)](#)

getBMRLearnerShortNames

Return learner short.names used in benchmark.

Description

Gets the learner short.names of the learners used in a benchmark experiment.

Usage

```
getBMRLearnerShortNames(bmr)
```

Arguments

bmr [\(BenchmarkResult\)](#)
Benchmark result.

Value

(character).

See Also

Other benchmark: [BenchmarkResult](#), [batchmark\(\)](#), [benchmark\(\)](#), [convertBMRTToRankMatrix\(\)](#), [friedmanPostHocTestBMR\(\)](#), [friedmanTestBMR\(\)](#), [generateCritDifferencesData\(\)](#), [getBMRAggrPerformances\(\)](#), [getBMRFeatSelResults\(\)](#), [getBMRFilteredFeatures\(\)](#), [getBMRLearnerIds\(\)](#), [getBMRLearners\(\)](#), [getBMRMeasureIds\(\)](#), [getBMRMeasures\(\)](#), [getBMRModels\(\)](#), [getBMRPerformances\(\)](#), [getBMRPredictions\(\)](#), [getBMRTaskDescs\(\)](#), [getBMRTaskIds\(\)](#), [getBMRTuneResults\(\)](#), [plotBMRBoxplots\(\)](#), [plotBMRRanksAsBarChart\(\)](#), [plotBMRSummary\(\)](#), [plotCritDifferences\(\)](#), [reduceBatchmarkResults\(\)](#)

getBMRMeasureIds	<i>Return measures IDs used in benchmark.</i>
------------------	---

Description

Gets the IDs of the measures used in a benchmark experiment.

Usage

```
getBMRMeasureIds(bmr)
```

Arguments

bmr	(BenchmarkResult) Benchmark result.
-----	--

Value

([list](#)). See above.

See Also

Other benchmark: [BenchmarkResult](#), [batchmark\(\)](#), [benchmark\(\)](#), [convertBMRTToRankMatrix\(\)](#), [friedmanPostHocTestBMR\(\)](#), [friedmanTestBMR\(\)](#), [generateCritDifferencesData\(\)](#), [getBMRAggrPerformances\(\)](#), [getBMRFeatSelResults\(\)](#), [getBMRFilteredFeatures\(\)](#), [getBMRLearnerIds\(\)](#), [getBMRLearnerShortNames\(\)](#), [getBMRLearners\(\)](#), [getBMRMeasures\(\)](#), [getBMRModels\(\)](#), [getBMRPerformances\(\)](#), [getBMRPredictions\(\)](#), [getBMRTaskDescs\(\)](#), [getBMRTaskIds\(\)](#), [getBMRTuneResults\(\)](#), [plotBMRBoxplots\(\)](#), [plotBMRRanksAsBarChart\(\)](#), [plotBMRSummary\(\)](#), [plotCritDifferences\(\)](#), [reduceBatchmarkResults\(\)](#)

getBMRMeasures	<i>Return measures used in benchmark.</i>
----------------	---

Description

Gets the measures used in a benchmark experiment.

Usage

```
getBMRMeasures(bmr)
```

Arguments

bmr	(BenchmarkResult) Benchmark result.
-----	--

Value

(list). See above.

See Also

Other benchmark: [BenchmarkResult](#), [batchmark\(\)](#), [benchmark\(\)](#), [convertBMRTToRankMatrix\(\)](#), [friedmanPostHocTestBMR\(\)](#), [friedmanTestBMR\(\)](#), [generateCritDifferencesData\(\)](#), [getBMRAggrPerformances\(\)](#), [getBMRFeatSelResults\(\)](#), [getBMRFilteredFeatures\(\)](#), [getBMRLearnerIds\(\)](#), [getBMRLearnerShortNames\(\)](#), [getBMRLearners\(\)](#), [getBMRMeasureIds\(\)](#), [getBMRModels\(\)](#), [getBMRPerformances\(\)](#), [getBMRPredictions\(\)](#), [getBMRTaskDescs\(\)](#), [getBMRTaskIds\(\)](#), [getBMRTuneResults\(\)](#), [plotBMRBoxplots\(\)](#), [plotBMRRanksAsBarChart\(\)](#), [plotBMRSummary\(\)](#), [plotCritDifferences\(\)](#), [reduceBatchmarkResults\(\)](#)

getBMRModels	<i>Extract all models from benchmark result.</i>
--------------	--

Description

A list of lists containing all [WrappedModels](#) trained in the benchmark experiment.

If `models` is FALSE in the call to [benchmark](#), the function will return NULL.

Usage

```
getBMRModels(bmr, task.ids = NULL, learner.ids = NULL, drop = FALSE)
```

Arguments

<code>bmr</code>	(BenchmarkResult) Benchmark result.
<code>task.ids</code>	(<code>character(1)</code>) Restrict result to certain tasks. Default is all.
<code>learner.ids</code>	(<code>character(1)</code>) Restrict result to certain learners. Default is all.
<code>drop</code>	(<code>logical(1)</code>) If <code>drop</code> is FALSE (the default), a nested list with the following structure is returned: <code>res[task.ids][learner.ids]</code> . If <code>drop</code> is set to TRUE it is checked if the list structure can be simplified. If only one learner was passed, a list with entries for each task is returned. If only one task was passed, the entries are named after the corresponding learner. For an experiment with both one task and learner, the whole list structure is removed. Note that the name of the task/learner will be dropped from the return object.

Value

(list).

See Also

Other benchmark: [BenchmarkResult](#), [batchmark\(\)](#), [benchmark\(\)](#), [convertBMRTToRankMatrix\(\)](#), [friedmanPostHocTestBMR\(\)](#), [friedmanTestBMR\(\)](#), [generateCritDifferencesData\(\)](#), [getBMRAggrPerformances\(\)](#), [getBMRFeatSelResults\(\)](#), [getBMRFilteredFeatures\(\)](#), [getBMRLearnerIds\(\)](#), [getBMRLearnerShortNames\(\)](#), [getBMRLearners\(\)](#), [getBMRMeasureIds\(\)](#), [getBMRMeasures\(\)](#), [getBMRPerformances\(\)](#), [getBMRPredictions\(\)](#), [getBMRTaskDescs\(\)](#), [getBMRTaskIds\(\)](#), [getBMRTuneResults\(\)](#), [plotBMRBoxplots\(\)](#), [plotBMRRanksAsBarChart\(\)](#), [plotBMRSummary\(\)](#), [plotCritDifferences\(\)](#), [reduceBatchmarkResults\(\)](#)

getBMRPerformances *Extract the test performance values from a benchmark result.*

Description

Either a list of lists of “measure.test” data.frames, as returned by [resample](#), or these objects are rbind-ed with extra columns “task.id” and “learner.id”.

Usage

```
getBMRPerformances(
  bmr,
  task.ids = NULL,
  learner.ids = NULL,
  as.df = FALSE,
  drop = FALSE
)
```

Arguments

bmr	(BenchmarkResult) Benchmark result.
task.ids	(character(1)) Restrict result to certain tasks. Default is all.
learner.ids	(character(1)) Restrict result to certain learners. Default is all.
as.df	(character(1)) Return one data.frame as result - or a list of lists of objects?. Default is FALSE.
drop	(logical(1)) If drop is FALSE (the default), a nested list with the following structure is returned: res[task.ids][learner.ids]. If drop is set to TRUE it is checked if the list structure can be simplified. If only one learner was passed, a list with entries for each task is returned. If only one task was passed, the entries are named after the corresponding learner. For an experiment with both one task and learner, the whole list structure is removed. Note that the name of the task/learner will be dropped from the return object.

Value

(list | data.frame). See above.

See Also

Other benchmark: [BenchmarkResult](#), [batchmark\(\)](#), [benchmark\(\)](#), [convertBMRtoRankMatrix\(\)](#), [friedmanPostHocTestBMR\(\)](#), [friedmanTestBMR\(\)](#), [generateCritDifferencesData\(\)](#), [getBMRAggrPerformances\(\)](#), [getBMRFeatSelResults\(\)](#), [getBMRFilteredFeatures\(\)](#), [getBMRLearnerIds\(\)](#), [getBMRLearnerShortNames\(\)](#), [getBMRLearners\(\)](#), [getBMRMeasureIds\(\)](#), [getBMRMeasures\(\)](#), [getBMRModels\(\)](#), [getBMRPredictions\(\)](#), [getBMRTaskDescs\(\)](#), [getBMRTaskIds\(\)](#), [getBMRTuneResults\(\)](#), [plotBMRBoxplots\(\)](#), [plotBMRRanksAsBarChart\(\)](#), [plotBMRSummary\(\)](#), [plotCritDifferences\(\)](#), [reduceBatchmarkResults\(\)](#)

getBMRPredictions	<i>Extract the predictions from a benchmark result.</i>
-------------------	---

Description

Either a list of lists of [ResamplePrediction](#) objects, as returned by [resample](#), or these objects are rbind-ed with extra columns “task.id” and “learner.id”.

If predict.type is “prob”, the probabilities for each class are returned in addition to the response.

If keep.pred is FALSE in the call to [benchmark](#), the function will return NULL.

Usage

```
getBMRPredictions(
  bmr,
  task.ids = NULL,
  learner.ids = NULL,
  as.df = FALSE,
  drop = FALSE
)
```

Arguments

bmr	(BenchmarkResult) Benchmark result.
task.ids	(character(1)) Restrict result to certain tasks. Default is all.
learner.ids	(character(1)) Restrict result to certain learners. Default is all.
as.df	(character(1)) Return one data.frame as result - or a list of lists of objects?. Default is FALSE.

drop (logical(1))
 If drop is FALSE (the default), a nested list with the following structure is returned:
 res[task.ids][learner.ids].
 If drop is set to TRUE it is checked if the list structure can be simplified.
 If only one learner was passed, a list with entries for each task is returned.
 If only one task was passed, the entries are named after the corresponding learner.
 For an experiment with both one task and learner, the whole list structure is removed.
 Note that the name of the task/learner will be dropped from the return object.

Value

(list | data.frame). See above.

See Also

Other benchmark: [BenchmarkResult](#), [batchmark\(\)](#), [benchmark\(\)](#), [convertBMRTToRankMatrix\(\)](#), [friedmanPostHocTestBMR\(\)](#), [friedmanTestBMR\(\)](#), [generateCritDifferencesData\(\)](#), [getBMRAggrPerformances\(\)](#), [getBMRFeatSelResults\(\)](#), [getBMRFilteredFeatures\(\)](#), [getBMRLearnerIds\(\)](#), [getBMRLearnerShortNames\(\)](#), [getBMRLearners\(\)](#), [getBMRMeasureIds\(\)](#), [getBMRMeasures\(\)](#), [getBMRModels\(\)](#), [getBMRPerformances\(\)](#), [getBMRTaskDescs\(\)](#), [getBMRTaskIds\(\)](#), [getBMRTuneResults\(\)](#), [plotBMRBoxplots\(\)](#), [plotBMRRanksAsBarChart\(\)](#), [plotBMRSummary\(\)](#), [plotCritDifferences\(\)](#), [reduceBatchmarkResults\(\)](#)

getBMRTaskDescriptions

Extract all task descriptions from benchmark result (DEPRECATED).

Description

A list containing all [TaskDescs](#) for each task contained in the benchmark experiment.

Usage

```
getBMRTaskDescriptions(bmr)
```

Arguments

bmr (BenchmarkResult)
 Benchmark result.

Value

(list).

getBMRTaskDescs *Extract all task descriptions from benchmark result.*

Description

A list containing all [TaskDescs](#) for each task contained in the benchmark experiment.

Usage

```
getBMRTaskDescs(bmr)
```

Arguments

bmr ([BenchmarkResult](#))
Benchmark result.

Value

([list](#)).

See Also

Other benchmark: [BenchmarkResult](#), [batchmark\(\)](#), [benchmark\(\)](#), [convertBMRTToRankMatrix\(\)](#), [friedmanPostHocTestBMR\(\)](#), [friedmanTestBMR\(\)](#), [generateCritDifferencesData\(\)](#), [getBMRAggrPerformances\(\)](#), [getBMRFeatSelResults\(\)](#), [getBMRFilteredFeatures\(\)](#), [getBMRLearnerIds\(\)](#), [getBMRLearnerShortNames\(\)](#), [getBMRLearners\(\)](#), [getBMRMeasureIds\(\)](#), [getBMRMeasures\(\)](#), [getBMRModels\(\)](#), [getBMRPerformances\(\)](#), [getBMRPredictions\(\)](#), [getBMRTaskIds\(\)](#), [getBMRTuneResults\(\)](#), [plotBMRBoxplots\(\)](#), [plotBMRRanksAsBarChart\(\)](#), [plotBMRSummary\(\)](#), [plotCritDifferences\(\)](#), [reduceBatchmarkResults\(\)](#)

getBMRTaskIds *Return task ids used in benchmark.*

Description

Gets the task IDs used in a benchmark experiment.

Usage

```
getBMRTaskIds(bmr)
```

Arguments

bmr ([BenchmarkResult](#))
Benchmark result.

Value

(character).

See Also

Other benchmark: [BenchmarkResult](#), [batchmark\(\)](#), [benchmark\(\)](#), [convertBMRTToRankMatrix\(\)](#), [friedmanPostHocTestBMR\(\)](#), [friedmanTestBMR\(\)](#), [generateCritDifferencesData\(\)](#), [getBMRAggrPerformances\(\)](#), [getBMRFeatSelResults\(\)](#), [getBMRFilteredFeatures\(\)](#), [getBMRLearnerIds\(\)](#), [getBMRLearnerShortNames\(\)](#), [getBMRLearners\(\)](#), [getBMRMeasureIds\(\)](#), [getBMRMeasures\(\)](#), [getBMRModels\(\)](#), [getBMRPerformances\(\)](#), [getBMRPredictions\(\)](#), [getBMRTaskDescs\(\)](#), [getBMRTuneResults\(\)](#), [plotBMRBoxplots\(\)](#), [plotBMRRanksAsBarChart\(\)](#), [plotBMRSummary\(\)](#), [plotCritDifferences\(\)](#), [reduceBatchmarkResults\(\)](#)

getBMRTuneResults	<i>Extract the tuning results from a benchmark result.</i>
-------------------	--

Description

Returns a nested list of [TuneResults](#). The first level of nesting is by data set, the second by learner, the third for the benchmark resampling iterations. If `as.df` is TRUE, a data frame with the “task.id”, “learner.id”, the resample iteration, the parameter values and the performances is returned.

Usage

```
getBMRTuneResults(
  bmr,
  task.ids = NULL,
  learner.ids = NULL,
  as.df = FALSE,
  drop = FALSE
)
```

Arguments

bmr	(BenchmarkResult) Benchmark result.
task.ids	(character(1)) Restrict result to certain tasks. Default is all.
learner.ids	(character(1)) Restrict result to certain learners. Default is all.
as.df	(character(1)) Return one data.frame as result - or a list of lists of objects?. Default is FALSE.
drop	(logical(1)) If drop is FALSE (the default), a nested list with the following structure is returned: res[task.ids][learner.ids]. If drop is set to TRUE it is checked if the list structure can be simplified.

If only one learner was passed, a list with entries for each task is returned.
If only one task was passed, the entries are named after the corresponding learner.

For an experiment with both one task and learner, the whole list structure is removed.

Note that the name of the task/learner will be dropped from the return object.

Value

(list | data.frame). See above.

See Also

Other benchmark: `BenchmarkResult`, `batchmark()`, `benchmark()`, `convertBMRTToRankMatrix()`, `friedmanPostHocTestBMR()`, `friedmanTestBMR()`, `generateCritDifferencesData()`, `getBMRAggrPerformances()`, `getBMRFeatSelResults()`, `getBMRFilteredFeatures()`, `getBMRLearnerIds()`, `getBMRLearnerShortNames()`, `getBMRLearners()`, `getBMRMeasureIds()`, `getBMRMeasures()`, `getBMRModels()`, `getBMRPerformances()`, `getBMRPredictions()`, `getBMRTaskDescs()`, `getBMRTaskIds()`, `plotBMRBoxplots()`, `plotBMRRanksAsBarChart()`, `plotBMRSummary()`, `plotCritDifferences()`, `reduceBatchmarkResults()`

getCaretParamSet	<i>Get tuning parameters from a learner of the caret R-package.</i>
------------------	---

Description

Constructs a grid of tuning parameters from a learner of the caret R-package. These values are then converted into a list of non-tunable parameters (`par.vals`) and a tunable `ParamHelpers::ParamSet` (`par.set`), which can be used by `tuneParams` for tuning the learner. Numerical parameters will either be specified by their lower and upper bounds or they will be discretized into specific values.

Usage

```
getCaretParamSet(learner, length = 3L, task, discretize = TRUE)
```

Arguments

learner	(character(1)) The name of the learner from caret (cf. https://topepo.github.io/caret/available-models.html). Note that the names in caret often differ from the ones in mlr.
length	(integer(1)) A length / precision parameter which is used by caret for generating the grid of tuning parameters. caret generates either as many values per tuning parameter / dimension as defined by length or only a single value (in case of non-tunable <code>par.vals</code>).
task	(Task) Learning task, which might be requested for creating the tuning grid.

discretize (logical(1))
Should the numerical parameters be discretized? Alternatively, they will be defined by their lower and upper bounds. The default is TRUE.

Value

(list(2)). A list of parameters:

- `par.vals` contains a list of all constant tuning parameters
- `par.set` is a [ParamHelpers::ParamSet](#), containing all the configurable tuning parameters

Examples

```
if (requireNamespace("caret") && requireNamespace("mlbench")) {
  library(caret)
  classifTask = makeClassifTask(data = iris, target = "Species")

  # (1) classification (random forest) with discretized parameters
  getCaretParamSet("rf", length = 9L, task = classifTask, discretize = TRUE)

  # (2) regression (gradient boosting machine) without discretized parameters
  library(mlbench)
  data(BostonHousing)
  regrTask = makeRegrTask(data = BostonHousing, target = "medv")
  getCaretParamSet("gbm", length = 9L, task = regrTask, discretize = FALSE)
}
```

getClassWeightParam *Get the class weight parameter of a learner.*

Description

Gets the class weight parameter of a learner.

Usage

```
getClassWeightParam(learner, lrn.id = NULL)
```

Arguments

learner (Learner | character(1))
The learner. If you pass a string the learner will be created via [makeLearner](#).

lrn.id (character)
Only used for BaseEnsembles. It is possible that multiple learners in a base ensemble have a class weight param. Specify the learner from which the class weight should be extracted.

Value

[numeric LearnerParam](#): A numeric parameter object, containing the class weight parameter of the given learner.

See Also

Other learner: [LearnerProperties](#), [getHyperPars\(\)](#), [getLearnerId\(\)](#), [getLearnerNote\(\)](#), [getLearnerPackages\(\)](#), [getLearnerParVals\(\)](#), [getLearnerParamSet\(\)](#), [getLearnerPredictType\(\)](#), [getLearnerShortName\(\)](#), [getLearnerType\(\)](#), [getParamSet\(\)](#), [helpLearnerParam\(\)](#), [helpLearner\(\)](#), [makeLearners\(\)](#), [makeLearner\(\)](#), [removeHyperPars\(\)](#), [setHyperPars\(\)](#), [setId\(\)](#), [setLearnerId\(\)](#), [setPredictThreshold\(\)](#), [setPredictType\(\)](#)

getConfMatrix	<i>Confusion matrix.</i>
---------------	--------------------------

Description

getConfMatrix is deprecated. Please use [calculateConfusionMatrix](#).

Calculates confusion matrix for (possibly resampled) prediction. Rows indicate true classes, columns predicted classes.

The marginal elements count the number of classification errors for the respective row or column, i.e., the number of errors when you condition on the corresponding true (rows) or predicted (columns) class. The last element in the margin diagonal displays the total amount of errors.

Note that for resampling no further aggregation is currently performed. All predictions on all test sets are joined to a vector \hat{y} , as are all labels joined to a vector y . Then \hat{y} is simply tabulated vs y , as if both were computed on a single test set. This probably mainly makes sense when cross-validation is used for resampling.

Usage

```
getConfMatrix(pred, relative = FALSE)
```

Arguments

pred	(Prediction) Prediction object.
relative	(logical(1)) If TRUE rows are normalized to show relative frequencies. Default is FALSE.

Value

[\(matrix\)](#). A confusion matrix.

See Also

[predict.WrappedModel](#)

getDefaultMeasure *Get default measure.*

Description

Get the default measure for a task type, task, task description or a learner. Currently these are:

- classif: mmce
- regr: mse
- cluster: db
- surv: cindex
- costsen: mcp
- multilabel: multilabel.hamloss

Usage

```
getDefaultMeasure(x)
```

Arguments

x	([character(1)' Task TaskDesc Learner) Task type, task, task description, learner name, a learner, or a type of learner (e.g. "classif").
---	---

Value

([Measure](#)).

getFailureModelDump *Return the error dump of FailureModel.*

Description

Returns the error dump that can be used with `debugger()` to evaluate errors. If `configureMlr` configuration `on.error.dump` is `FALSE`, this returns `NULL`.

Usage

```
getFailureModelDump(model)
```

Arguments

model	(WrappedModel) The model.
-------	--

Value

(`last.dump`).

getFailureModelMsg *Return error message of FailureModel.*

Description

Such a model is created when one sets the corresponding option in [configureMlr](#). If no failure occurred, NA is returned.

For complex wrappers this getter returns the first error message encountered in ANY model that failed.

Usage

```
getFailureModelMsg(model)
```

Arguments

model ([WrappedModel](#))
The model.

Value

(character(1)).

getFeatSelResult *Returns the selected feature set and optimization path after training.*

Description

Returns the selected feature set and optimization path after training.

Usage

```
getFeatSelResult(object)
```

Arguments

object ([WrappedModel](#))
Trained Model created with [makeFeatSelWrapper](#).

Value

([FeatSelResult](#)).

See Also

Other featsel: [FeatSelControl](#), [analyzeFeatSelResult\(\)](#), [makeFeatSelWrapper\(\)](#), [selectFeatures\(\)](#)

getFeatureImportance *Calculates feature importance values for trained models.*

Description

For some learners it is possible to calculate a feature importance measure. `getFeatureImportance` extracts those values from trained models. See below for a list of supported learners.

Usage

```
getFeatureImportance(object, ...)
```

Arguments

object	(WrappedModel) Wrapped model, result of <code>train()</code> .
...	(any) Additional parameters, which are passed to the underlying importance value generating function.

Details

- `boosting`
Measure which accounts the gain of Gini index given by a feature in a tree and the weight of that tree.
- `cforest`
Permutation principle of the 'mean decrease in accuracy' principle in `randomForest`. If `auc=TRUE` (only for binary classification), area under the curve is used as measure. The algorithm used for the survival learner is 'extremely slow and experimental; use at your own risk'. See [party::varimp\(\)](#) for details and further parameters.
- `gbm`
Estimation of relative influence for each feature. See [gbm::relative.influence\(\)](#) for details and further parameters.
- `h2o`
Relative feature importances as returned by [h2o::h2o.varimp\(\)](#).
- `randomForest`
For `type = 2` (the default) the 'MeanDecreaseGini' is measured, which is based on the Gini impurity index used for the calculation of the nodes. Alternatively, you can set `type` to 1, then the measure is the mean decrease in accuracy calculated on OOB data. Note, that in this case the learner's parameter `importance` needs to be set to be able to compute feature importance values. See [randomForest::importance\(\)](#) for details.
- `RRF`
This is identical to `randomForest`.
- `randomForestSRC`
This method can calculate feature importance for various measures. By default the Breiman-Cutler permutation method is used. See [randomForestSRC::vimp\(\)](#) for details.

- **ranger**
Supports both measures mentioned above for the randomForest learner. Note, that you need to specifically set the learners parameter importance, to be able to compute feature importance measures. See [ranger::importance\(\)](#) and [ranger::ranger\(\)](#) for details.
- **rpart**
Sum of decrease in impurity for each of the surrogate variables at each node
- **xgboost**
The value implies the relative contribution of the corresponding feature to the model calculated by taking each feature's contribution for each tree in the model. The exact computation of the importance in xgboost is undocumented.

Value

(FeatureImportance) An object containing a data.frame of the variable importances and further information.

getFilteredFeatures *Returns the filtered features.*

Description

Returns the filtered features.

Usage

```
getFilteredFeatures(model)
```

Arguments

model ([WrappedModel](#))
Trained Model created with [makeFilterWrapper](#).

Value

([character](#)).

See Also

Other filter: [filterFeatures\(\)](#), [generateFilterValuesData\(\)](#), [listFilterEnsembleMethods\(\)](#), [listFilterMethods\(\)](#), [makeFilterEnsemble\(\)](#), [makeFilterWrapper\(\)](#), [makeFilter\(\)](#), [plotFilterValues\(\)](#)

getFunctionalFeatures *Get only functional features from a task or a data.frame.*

Description

The parameters “subset”, “features”, and “recode.target” are ignored for the data.frame method.

Usage

```
getFunctionalFeatures(object, subset = NULL, features, recode.target = "no")

## S3 method for class 'Task'
getFunctionalFeatures(object, subset = NULL, features, recode.target = "no")

## S3 method for class 'data.frame'
getFunctionalFeatures(object, subset = NULL, features, recode.target = "no")
```

Arguments

object	(Task/data.frame) Object to check on.
subset	(integer logical NULL) Selected cases. Either a logical or an index vector. By default NULL if all observations are used.
features	(character integer logical) Vector of selected inputs. You can either pass a character vector with the feature names, a vector of indices, or a logical vector. In case of an index vector each element denotes the position of the feature name returned by <code>getTaskFeatureNames</code> . Note that the target feature is always included in the resulting task, you should not pass it here. Default is to use all features.
recode.target	(character(1)) Should target classes be recoded? Supported are binary and multilabel classification and survival. Possible values for binary classification are “01”, “-1+1” and “drop.levels”. In the two latter cases the target vector is converted into a numeric vector. The positive class is coded as “+1” and the negative class either as “0” or “-1”. “drop.levels” will remove empty factor levels in the target column. In the multilabel case the logical targets can be converted to factors with “multilabel.factor”. For survival, you may choose to recode the survival times to “left”, “right” or “interval2” censored times using “lcens”, “rcens” or “icens”, respectively. See <code>survival::Surv</code> for the format specification. Default for both binary classification and survival is “no” (do nothing).

Value

Returns a data.frame containing only the functional features.

```
getHomogeneousEnsembleModels
```

Deprecated, use getLearnerModel instead.

Description

Deprecated, use getLearnerModel instead.

Usage

```
getHomogeneousEnsembleModels(model, learner.models = FALSE)
```

Arguments

model Deprecated.

learner.models Deprecated.

```
getHyperPars
```

Get current parameter settings for a learner.

Description

Retrieves the current hyperparameter settings of a learner.

Usage

```
getHyperPars(learner, for.fun = c("train", "predict", "both"))
```

Arguments

learner ([Learner](#))
The learner.

for.fun (character(1))
Restrict the returned settings to hyperparameters corresponding to when the are used (see [ParamHelpers::LearnerParam](#)). Must be a subset of: "train", "predict" or "both". Default is c("train", "predict", "both").

Details

This function only shows hyperparameters that differ from the learner default (because mlr changed the default) or if the user set hyperparameters manually during learner creation. If you want to have an overview of all available hyperparameters use [getParamSet\(\)](#).

Value

([list](#)). A named list of values.

See Also

Other learner: [LearnerProperties](#), [getClassWeightParam\(\)](#), [getLearnerId\(\)](#), [getLearnerNote\(\)](#), [getLearnerPackages\(\)](#), [getLearnerParVals\(\)](#), [getLearnerParamSet\(\)](#), [getLearnerPredictType\(\)](#), [getLearnerShortName\(\)](#), [getLearnerType\(\)](#), [getParamSet\(\)](#), [helpLearnerParam\(\)](#), [helpLearner\(\)](#), [makeLearners\(\)](#), [makeLearner\(\)](#), [removeHyperPars\(\)](#), [setHyperPars\(\)](#), [setId\(\)](#), [setLearnerId\(\)](#), [setPredictThreshold\(\)](#), [setPredictType\(\)](#)

Examples

```
getHyperPars(makeLearner("classif.ranger"))

## set learner hyperparameter `mtry` manually
getHyperPars(makeLearner("classif.ranger", mtry = 100))
```

getLearnerId	<i>Get the ID of the learner.</i>
--------------	-----------------------------------

Description

Get the ID of the learner.

Usage

```
getLearnerId(learner)
```

Arguments

learner [\(Learner | character\(1\)\)](#)
The learner. If you pass a string the learner will be created via [makeLearner](#).

Value

(character(1)).

See Also

Other learner: [LearnerProperties](#), [getClassWeightParam\(\)](#), [getHyperPars\(\)](#), [getLearnerNote\(\)](#), [getLearnerPackages\(\)](#), [getLearnerParVals\(\)](#), [getLearnerParamSet\(\)](#), [getLearnerPredictType\(\)](#), [getLearnerShortName\(\)](#), [getLearnerType\(\)](#), [getParamSet\(\)](#), [helpLearnerParam\(\)](#), [helpLearner\(\)](#), [makeLearners\(\)](#), [makeLearner\(\)](#), [removeHyperPars\(\)](#), [setHyperPars\(\)](#), [setId\(\)](#), [setLearnerId\(\)](#), [setPredictThreshold\(\)](#), [setPredictType\(\)](#)

getLearnerModel	<i>Get underlying R model of learner integrated into mlr.</i>
-----------------	---

Description

Get underlying R model of learner integrated into mlr.

Usage

```
getLearnerModel(model, more.unwrap = FALSE)
```

Arguments

model	(WrappedModel) The model, returned by e.g., train .
more.unwrap	(logical(1)) Some learners are not basic learners from R, but implemented in mlr as meta-techniques. Examples are everything that inherits from HomogeneousEnsemble . In these cases, the learner.model is often a list of mlr WrappedModels . This option allows to strip them further to basic R models. The option is simply ignored for basic learner models. Default is FALSE.

Value

(any). A fitted model, depending the learner / wrapped package. E.g., a model of class [rpart::rpart](#) for learner “[classif.rpart](#)”.

getLearnerNote	<i>Get the note for the learner.</i>
----------------	--------------------------------------

Description

Get the note for the learner.

Usage

```
getLearnerNote(learner)
```

Arguments

learner	(Learner character(1)) The learner. If you pass a string the learner will be created via makeLearner .
---------	--

Value

([character](#)).

See Also

Other learner: [LearnerProperties](#), [getClassWeightParam\(\)](#), [getHyperPars\(\)](#), [getLearnerId\(\)](#), [getLearnerPackages\(\)](#), [getLearnerParVals\(\)](#), [getLearnerParamSet\(\)](#), [getLearnerPredictType\(\)](#), [getLearnerShortName\(\)](#), [getLearnerType\(\)](#), [getParamSet\(\)](#), [helpLearnerParam\(\)](#), [helpLearner\(\)](#), [makeLearners\(\)](#), [makeLearner\(\)](#), [removeHyperPars\(\)](#), [setHyperPars\(\)](#), [setId\(\)](#), [setLearnerId\(\)](#), [setPredictThreshold\(\)](#), [setPredictType\(\)](#)

getLearnerPackages *Get the required R packages of the learner.*

Description

Get the R packages the learner requires.

Usage

```
getLearnerPackages(learner)
```

Arguments

learner ([Learner](#) | character(1))
 The learner. If you pass a string the learner will be created via [makeLearner](#).

Value

([character](#)).

See Also

Other learner: [LearnerProperties](#), [getClassWeightParam\(\)](#), [getHyperPars\(\)](#), [getLearnerId\(\)](#), [getLearnerNote\(\)](#), [getLearnerParVals\(\)](#), [getLearnerParamSet\(\)](#), [getLearnerPredictType\(\)](#), [getLearnerShortName\(\)](#), [getLearnerType\(\)](#), [getParamSet\(\)](#), [helpLearnerParam\(\)](#), [helpLearner\(\)](#), [makeLearners\(\)](#), [makeLearner\(\)](#), [removeHyperPars\(\)](#), [setHyperPars\(\)](#), [setId\(\)](#), [setLearnerId\(\)](#), [setPredictThreshold\(\)](#), [setPredictType\(\)](#)

getLearnerParamSet *Get the parameter set of the learner.*

Description

Alias for [getParamSet](#).

Usage

```
getLearnerParamSet(learner)
```

Arguments

learner (Learner | character(1))
The learner. If you pass a string the learner will be created via [makeLearner](#).

Value

ParamSet.

See Also

Other learner: [LearnerProperties](#), [getClassWeightParam\(\)](#), [getHyperPars\(\)](#), [getLearnerId\(\)](#), [getLearnerNote\(\)](#), [getLearnerPackages\(\)](#), [getLearnerParVals\(\)](#), [getLearnerPredictType\(\)](#), [getLearnerShortName\(\)](#), [getLearnerType\(\)](#), [getParamSet\(\)](#), [helpLearnerParam\(\)](#), [helpLearner\(\)](#), [makeLearners\(\)](#), [makeLearner\(\)](#), [removeHyperPars\(\)](#), [setHyperPars\(\)](#), [setId\(\)](#), [setLearnerId\(\)](#), [setPredictThreshold\(\)](#), [setPredictType\(\)](#)

getLearnerParVals *Get the parameter values of the learner.*

Description

Alias for [getHyperPars](#).

Usage

```
getLearnerParVals(learner, for.fun = c("train", "predict", "both"))
```

Arguments

learner (Learner | character(1))
The learner. If you pass a string the learner will be created via [makeLearner](#).

for.fun (character(1))
Restrict the returned settings to hyperparameters corresponding to when the are used (see [ParamHelpers::LearnerParam](#)). Must be a subset of: “train”, “predict” or “both”. Default is `c("train", "predict", "both")`.

Value

(list). A named list of values.

See Also

Other learner: [LearnerProperties](#), [getClassWeightParam\(\)](#), [getHyperPars\(\)](#), [getLearnerId\(\)](#), [getLearnerNote\(\)](#), [getLearnerPackages\(\)](#), [getLearnerParamSet\(\)](#), [getLearnerPredictType\(\)](#), [getLearnerShortName\(\)](#), [getLearnerType\(\)](#), [getParamSet\(\)](#), [helpLearnerParam\(\)](#), [helpLearner\(\)](#), [makeLearners\(\)](#), [makeLearner\(\)](#), [removeHyperPars\(\)](#), [setHyperPars\(\)](#), [setId\(\)](#), [setLearnerId\(\)](#), [setPredictThreshold\(\)](#), [setPredictType\(\)](#)

getLearnerPredictType *Get the predict type of the learner.*

Description

Get the predict type of the learner.

Usage

```
getLearnerPredictType(learner)
```

Arguments

learner ([Learner](#) | character(1))
The learner. If you pass a string the learner will be created via [makeLearner](#).

Value

(character(1)).

See Also

Other learner: [LearnerProperties](#), [getClassWeightParam\(\)](#), [getHyperPars\(\)](#), [getLearnerId\(\)](#), [getLearnerNote\(\)](#), [getLearnerPackages\(\)](#), [getLearnerParVals\(\)](#), [getLearnerParamSet\(\)](#), [getLearnerShortName\(\)](#), [getLearnerType\(\)](#), [getParamSet\(\)](#), [helpLearnerParam\(\)](#), [helpLearner\(\)](#), [makeLearners\(\)](#), [makeLearner\(\)](#), [removeHyperPars\(\)](#), [setHyperPars\(\)](#), [setId\(\)](#), [setLearnerId\(\)](#), [setPredictThreshold\(\)](#), [setPredictType\(\)](#)

getLearnerShortName *Get the short name of the learner.*

Description

For an ordinary learner simply its short name is returned. For wrapped learners, the wrapper id is successively attached to the short name of the base learner. E.g: “rf.bagged.imputed”

Usage

```
getLearnerShortName(learner)
```

Arguments

learner ([Learner](#) | character(1))
The learner. If you pass a string the learner will be created via [makeLearner](#).

Value

(character(1)).

See Also

Other learner: [LearnerProperties](#), [getClassWeightParam\(\)](#), [getHyperPars\(\)](#), [getLearnerId\(\)](#), [getLearnerNote\(\)](#), [getLearnerPackages\(\)](#), [getLearnerParVals\(\)](#), [getLearnerParamSet\(\)](#), [getLearnerPredictType\(\)](#), [getLearnerType\(\)](#), [getParamSet\(\)](#), [helpLearnerParam\(\)](#), [helpLearner\(\)](#), [makeLearners\(\)](#), [makeLearner\(\)](#), [removeHyperPars\(\)](#), [setHyperPars\(\)](#), [setId\(\)](#), [setLearnerId\(\)](#), [setPredictThreshold\(\)](#), [setPredictType\(\)](#)

getLearnerType	<i>Get the type of the learner.</i>
----------------	-------------------------------------

Description

Get the type of the learner.

Usage

```
getLearnerType(learner)
```

Arguments

learner ([Learner](#) | character(1))
The learner. If you pass a string the learner will be created via [makeLearner](#).

Value

(character(1)).

See Also

Other learner: [LearnerProperties](#), [getClassWeightParam\(\)](#), [getHyperPars\(\)](#), [getLearnerId\(\)](#), [getLearnerNote\(\)](#), [getLearnerPackages\(\)](#), [getLearnerParVals\(\)](#), [getLearnerParamSet\(\)](#), [getLearnerPredictType\(\)](#), [getLearnerShortName\(\)](#), [getParamSet\(\)](#), [helpLearnerParam\(\)](#), [helpLearner\(\)](#), [makeLearners\(\)](#), [makeLearner\(\)](#), [removeHyperPars\(\)](#), [setHyperPars\(\)](#), [setId\(\)](#), [setLearnerId\(\)](#), [setPredictThreshold\(\)](#), [setPredictType\(\)](#)

getMlrOptions *Returns a list of mlr's options.*

Description

Gets the options for mlr.

Usage

```
getMlrOptions()
```

Value

(list).

See Also

Other configure: [configureMlr\(\)](#)

getMultilabelBinaryPerformances
Retrieve binary classification measures for multilabel classification predictions.

Description

Measures the quality of each binary label prediction w.r.t. some binary classification performance measure.

Usage

```
getMultilabelBinaryPerformances(pred, measures)
```

Arguments

pred	(Prediction) Multilabel Prediction object.
measures	(Measure list of Measure) Performance measure(s) to evaluate, must be applicable to binary classification performance. Default is mmce.

Value

(named matrix). Performance value(s), column names are measure(s), row names are labels.

See Also

Other multilabel: [makeMultilabelBinaryRelevanceWrapper\(\)](#), [makeMultilabelClassifierChainsWrapper\(\)](#), [makeMultilabelDBRWrapper\(\)](#), [makeMultilabelNestedStackingWrapper\(\)](#), [makeMultilabelStackingWrapper\(\)](#)

Examples

```
# see makeMultilabelBinaryRelevanceWrapper
```

```
getNestedTuneResultsOptPathDf
```

Get the opt.paths from each tuning step from the outer resampling.

Description

After you resampled a tuning wrapper (see [makeTuneWrapper](#)) with `resample(..., extract = getTuneResult)` this helper returns a `data.frame` with with all `opt.paths` combined by `rbind`. An additional column `iter` indicates to what resampling iteration the row belongs.

Usage

```
getNestedTuneResultsOptPathDf(r, trafo = FALSE)
```

Arguments

<code>r</code>	(ResampleResult) The result of resampling of a tuning wrapper.
<code>trafo</code>	<code>(logical(1))</code> Should the units of the hyperparameter path be converted to the transformed scale? This is only necessary when <code>trafo</code> was used to create the <code>opt.paths</code> . Note that <code>opt.paths</code> are always stored on the untransformed scale. Default is <code>FALSE</code> .

Value

[\(data.frame\)](#). See above.

See Also

Other tune: [TuneControl](#), [getNestedTuneResultsX\(\)](#), [getResamplingIndices\(\)](#), [getTuneResult\(\)](#), [makeModelMultiplexerParamSet\(\)](#), [makeModelMultiplexer\(\)](#), [makeTuneControlCMAES\(\)](#), [makeTuneControlDesign\(\)](#), [makeTuneControlGenSA\(\)](#), [makeTuneControlGrid\(\)](#), [makeTuneControlIrace\(\)](#), [makeTuneControlIMBO\(\)](#), [makeTuneControlRandom\(\)](#), [makeTuneWrapper\(\)](#), [tuneParams\(\)](#), [tuneThreshold\(\)](#)

Examples

```
# see example of makeTuneWrapper
```

`getNestedTuneResultsX` *Get the tuned hyperparameter settings from a nested tuning.*

Description

After you resampled a tuning wrapper (see [makeTuneWrapper](#)) with `resample(..., extract = getTuneResult)` this helper returns a `data.frame` with the best found hyperparameter settings for each resampling iteration.

Usage

```
getNestedTuneResultsX(r)
```

Arguments

`r` ([ResampleResult](#))
The result of resampling of a tuning wrapper.

Value

([data.frame](#)). One column for each tuned hyperparameter and one row for each outer resampling iteration.

See Also

Other tune: [TuneControl](#), [getNestedTuneResultsOptPathDf\(\)](#), [getResamplingIndices\(\)](#), [getTuneResult\(\)](#), [makeModelMultiplexerParamSet\(\)](#), [makeModelMultiplexer\(\)](#), [makeTuneControlCMAES\(\)](#), [makeTuneControlDesign\(\)](#), [makeTuneControlGenSA\(\)](#), [makeTuneControlGrid\(\)](#), [makeTuneControlIrace\(\)](#), [makeTuneControlMBO\(\)](#), [makeTuneControlRandom\(\)](#), [makeTuneWrapper\(\)](#), [tuneParams\(\)](#), [tuneThreshold\(\)](#)

Examples

```
# see example of makeTuneWrapper
```

`getOOBPreds` *Extracts out-of-bag predictions from trained models.*

Description

Learners like `randomForest` produce out-of-bag predictions. `getOOBPreds` extracts this information from trained models and builds a prediction object as provided by `predict` (with prediction time set to NA). In the classification case: What is stored exactly in the ([Prediction](#)) object depends on the `predict.type` setting of the [Learner](#).

You can call `listLearners(properties = "oobpreds")` to get a list of learners which provide this.

Usage

```
getOOBPreds(model, task)
```

Arguments

model	(WrappedModel) The model.
task	(Task) The task.

Value

([Prediction](#)).

Examples

```
training.set = sample(1:150, 50)
lrn = makeLearner("classif.ranger", predict.type = "prob", predict.threshold = 0.6)
mod = train(lrn, sonar.task, subset = training.set)
oob = getOOBPreds(mod, sonar.task)
oob
performance(oob, measures = list(auc, mmce))
```

getParamSet

Get a description of all possible parameter settings for a learner.

Description

Returns the [ParamHelpers::ParamSet](#) from a [Learner](#).

Value

[ParamSet](#).

See Also

Other learner: [LearnerProperties](#), [getClassWeightParam\(\)](#), [getHyperPars\(\)](#), [getLearnerId\(\)](#), [getLearnerNote\(\)](#), [getLearnerPackages\(\)](#), [getLearnerParVals\(\)](#), [getLearnerParamSet\(\)](#), [getLearnerPredictType\(\)](#), [getLearnerShortName\(\)](#), [getLearnerType\(\)](#), [helpLearnerParam\(\)](#), [helpLearner\(\)](#), [makeLearners\(\)](#), [makeLearner\(\)](#), [removeHyperPars\(\)](#), [setHyperPars\(\)](#), [setId\(\)](#), [setLearnerId\(\)](#), [setPredictThreshold\(\)](#), [setPredictType\(\)](#)

getPredictionDump *Return the error dump of a failed Prediction.*

Description

Returns the error dump that can be used with `debugger()` to evaluate errors. If `configureMlr` configuration on `.error.dump` is `FALSE` or if the prediction did not fail, this returns `NULL`.

Usage

```
getPredictionDump(pred)
```

Arguments

pred ([Prediction](#))
Prediction object.

Value

(last.dump).

See Also

Other debug: [FailureModel](#), [ResampleResult](#), [getRRDump\(\)](#)

getPredictionProbabilities
Get probabilities for some classes.

Description

Get probabilities for some classes.

Usage

```
getPredictionProbabilities(pred, cl)
```

Arguments

pred ([Prediction](#))
Prediction object.

cl ([character](#))
Names of classes. Default is either all classes for multi-class / multilabel problems or the positive class for binary classification.

Value

([data.frame](#)) with numerical columns or a numerical vector if length of `c1` is 1. Order of columns is defined by `c1`.

See Also

Other predict: [asROCRPrediction\(\)](#), [getPredictionResponse\(\)](#), [getPredictionTaskDesc\(\)](#), [predict.WrappedModel\(\)](#), [setPredictThreshold\(\)](#), [setPredictType\(\)](#)

Examples

```
task = makeClassifTask(data = iris, target = "Species")
lrn = makeLearner("classif.lda", predict.type = "prob")
mod = train(lrn, task)
# predict probabilities
pred = predict(mod, newdata = iris)

# Get probabilities for all classes
head(getPredictionProbabilities(pred))

# Get probabilities for a subset of classes
head(getPredictionProbabilities(pred, c("setosa", "virginica")))
```

`getPredictionResponse` *Get response / truth from prediction object.*

Description

The following types are returned, depending on task type:

classif	factor
regr	numeric
se	numeric
cluster	integer
surv	numeric
multilabel	logical matrix, columns named with labels

Usage

```
getPredictionResponse(pred)
```

```
getPredictionSE(pred)
```

```
getPredictionTruth(pred)
```

Arguments

pred [\(Prediction\)](#)
Prediction object.

Value

See above.

See Also

Other predict: [asROCRPrediction\(\)](#), [getPredictionProbabilities\(\)](#), [getPredictionTaskDesc\(\)](#), [predict.WrappedModel\(\)](#), [setPredictThreshold\(\)](#), [setPredictType\(\)](#)

getPredictionTaskDesc *Get summarizing task description from prediction.*

Description

See title.

Usage

```
getPredictionTaskDesc(pred)
```

Arguments

pred [\(Prediction\)](#)
Prediction object.

Value

ret_taskdesc

See Also

Other predict: [asROCRPrediction\(\)](#), [getPredictionProbabilities\(\)](#), [getPredictionResponse\(\)](#), [predict.WrappedModel\(\)](#), [setPredictThreshold\(\)](#), [setPredictType\(\)](#)

getProbabilities *Deprecated, use getPredictionProbabilities instead.*

Description

Deprecated, use getPredictionProbabilities instead.

Usage

```
getProbabilities(pred, cl)
```

Arguments

pred	Deprecated.
cl	Deprecated.

getResamplingIndices *Get the resampling indices from a tuning or feature selection wrapper.*

Description

After you resampled a tuning or feature selection wrapper (see [makeTuneWrapper](#)) with `resample(..., extract = getTuneResult)` or `resample(..., extract = getFeatSelResult)` this helper returns a list with the resampling indices used for the respective method.

Usage

```
getResamplingIndices(object, inner = FALSE)
```

Arguments

object	(ResampleResult) The result of resampling of a tuning or feature selection wrapper.
inner	(logical) If TRUE, returns the inner indices of a nested resampling setting.

Value

[\(list\)](#). One list for each outer resampling fold.

See Also

Other tune: [TuneControl](#), [getNestedTuneResultsOptPathDf\(\)](#), [getNestedTuneResultsX\(\)](#), [getTuneResult\(\)](#), [makeModelMultiplexerParamSet\(\)](#), [makeModelMultiplexer\(\)](#), [makeTuneControlCMAES\(\)](#), [makeTuneControlDesign\(\)](#), [makeTuneControlGenSA\(\)](#), [makeTuneControlGrid\(\)](#), [makeTuneControlIrace\(\)](#), [makeTuneControlMBO\(\)](#), [makeTuneControlRandom\(\)](#), [makeTuneWrapper\(\)](#), [tuneParams\(\)](#), [tuneThreshold\(\)](#)

Examples

```

task = makeClassifTask(data = iris, target = "Species")
lrn = makeLearner("classif.rpart")
# stupid mini grid
ps = makeParamSet(
  makeDiscreteParam("cp", values = c(0.05, 0.1)),
  makeDiscreteParam("minsplit", values = c(10, 20))
)
ctrl = makeTuneControlGrid()
inner = makeResampleDesc("Holdout")
outer = makeResampleDesc("CV", iters = 2)
lrn = makeTuneWrapper(lrn, resampling = inner, par.set = ps, control = ctrl)
# nested resampling for evaluation
# we also extract tuned hyper pars in each iteration and by that the resampling indices
r = resample(lrn, task, outer, extract = getTuneResult)
# get tuning indices
getResamplingIndices(r, inner = TRUE)

```

getRRDump

Return the error dump of ResampleResult.

Description

Returns the error dumps generated during resampling, which can be used with `debugger()` to debug errors. These dumps are saved if `configureMlr` configuration `on.error.dump`, or the corresponding learner `config`, is `TRUE`.

The returned object is a list with as many entries as the resampling being used has folds. Each of these entries can have a subset of the following slots, depending on which step in the resampling iteration failed: “train” (error during training step), “predict.train” (prediction on training subset), “predict.test” (prediction on test subset).

Usage

```
getRRDump(res)
```

Arguments

res ([ResampleResult](#))
The result of `resample`.

Value

[list](#).

See Also

Other debug: [FailureModel](#), [ResampleResult](#), [getPredictionDump\(\)](#)

getRRPredictionList *Get list of predictions for train and test set of each single resample iteration.*

Description

This function creates a list with two slots train and test where each slot is again a list of [Prediction](#) objects for each single resample iteration. In case that predict = "train" was used for the resample description (see [makeResampleDesc](#)), the slot test will be NULL and in case that predict = "test" was used, the slot train will be NULL.

Usage

```
getRRPredictionList(res, ...)
```

Arguments

res ([ResampleResult](#))
The result of [resample](#) run with keep.pred = TRUE.

... (any)
Further options passed to [makePrediction](#).

Value

[list](#).

See Also

Other resample: [ResamplePrediction](#), [ResampleResult](#), [addRRMeasure\(\)](#), [getRRPredictions\(\)](#), [getRRTaskDescription\(\)](#), [getRRTaskDesc\(\)](#), [makeResampleDesc\(\)](#), [makeResampleInstance\(\)](#), [resample\(\)](#)

getRRPredictions *Get predictions from resample results.*

Description

Very simple getter.

Usage

```
getRRPredictions(res)
```

Arguments

res ([ResampleResult](#))
The result of [resample](#) run with keep.pred = TRUE.

Value

(ResamplePrediction).

See Also

Other resample: ResamplePrediction, ResampleResult, addRRMeasure(), getRRPredictionList(), getRRTaskDescription(), getRRTaskDesc(), makeResampleDesc(), makeResampleInstance(), resample()

getRRTaskDesc

Get task description from resample results (DEPRECATED).

Description

Get a summarizing task description.

Usage

```
getRRTaskDesc(res)
```

Arguments

res (ResampleResult)
The result of resample.

Value

(TaskDesc).

See Also

Other resample: ResamplePrediction, ResampleResult, addRRMeasure(), getRRPredictionList(), getRRPredictions(), getRRTaskDescription(), makeResampleDesc(), makeResampleInstance(), resample()

getRRTaskDescription *Get task description from resample results (DEPRECATED).*

Description

Get a summarizing task description.

Usage

```
getRRTaskDescription(res)
```

Arguments

res [\(ResampleResult\)](#)
The result of [resample](#).

Value

[\(TaskDesc\)](#).

See Also

Other resample: [ResamplePrediction](#), [ResampleResult](#), [addRRMeasure\(\)](#), [getRRPredictionList\(\)](#), [getRRPredictions\(\)](#), [getRRTaskDesc\(\)](#), [makeResampleDesc\(\)](#), [makeResampleInstance\(\)](#), [resample\(\)](#)

getStackedBaseLearnerPredictions

Returns the predictions for each base learner.

Description

Returns the predictions for each base learner.

Usage

```
getStackedBaseLearnerPredictions(model, newdata = NULL)
```

Arguments

model [\(WrappedModel\)](#)
Wrapped model, result of train.

newdata [\(data.frame\)](#)
New observations, for which the predictions using the specified base learners should be returned. Default is NULL and extracts the base learner predictions that were made during the training.

Details

None.

getTaskClassLevels *Get the class levels for classification and multilabel tasks.*

Description

NB: For multilabel, [getTaskTargetNames](#) and [getTaskClassLevels](#) actually return the same thing.

Usage

```
getTaskClassLevels(x)
```

Arguments

x ([Task](#) | [TaskDesc](#))
Task or its description object.

Value

([character](#)).

See Also

Other task: [getTaskCosts\(\)](#), [getTaskData\(\)](#), [getTaskDesc\(\)](#), [getTaskFeatureNames\(\)](#), [getTaskFormula\(\)](#), [getTaskId\(\)](#), [getTaskNFeats\(\)](#), [getTaskSize\(\)](#), [getTaskTargetNames\(\)](#), [getTaskTargets\(\)](#), [getTaskType\(\)](#), [subsetTask\(\)](#)

getTaskCosts *Extract costs in task.*

Description

Returns “NULL” if the task is not of type “costsens”.

Usage

```
getTaskCosts(task, subset = NULL)
```

Arguments

task ([CostSensTask](#))
The task.

subset ([integer](#) | [logical](#) | NULL)
Selected cases. Either a logical or an index vector. By default NULL if all observations are used.

Value

(matrix | NULL).

See Also

Other task: [getTaskClassLevels\(\)](#), [getTaskData\(\)](#), [getTaskDesc\(\)](#), [getTaskFeatureNames\(\)](#), [getTaskFormula\(\)](#), [getTaskId\(\)](#), [getTaskNFeats\(\)](#), [getTaskSize\(\)](#), [getTaskTargetNames\(\)](#), [getTaskTargets\(\)](#), [getTaskType\(\)](#), [subsetTask\(\)](#)

getTaskData	<i>Extract data in task.</i>
-------------	------------------------------

Description

Useful in [trainLearner](#) when you add a learning machine to the package.

Usage

```
getTaskData(
  task,
  subset = NULL,
  features,
  target.extra = FALSE,
  recode.target = "no",
  functionals.as = "dfcols"
)
```

Arguments

task	(Task) The task.
subset	(integer logical NULL) Selected cases. Either a logical or an index vector. By default NULL if all observations are used.
features	(character integer logical) Vector of selected inputs. You can either pass a character vector with the feature names, a vector of indices, or a logical vector. In case of an index vector each element denotes the position of the feature name returned by getTaskFeatureNames . Note that the target feature is always included in the resulting task, you should not pass it here. Default is to use all features.
target.extra	(logical (1)) Should target vector be returned separately? If not, a single data.frame including the target columns is returned, otherwise a list with the input data.frame and an extra vector or data.frame for the targets. Default is FALSE.

`recode.target` (character(1))
Should target classes be recoded? Supported are binary and multilabel classification and survival. Possible values for binary classification are “01”, “-1+1” and “drop.levels”. In the two latter cases the target vector is converted into a numeric vector. The positive class is coded as “+1” and the negative class either as “0” or “-1”. “drop.levels” will remove empty factor levels in the target column. In the multilabel case the logical targets can be converted to factors with “multilabel.factor”. For survival, you may choose to recode the survival times to “left”, “right” or “interval2” censored times using “lcens”, “rcens” or “icens”, respectively. See [survival::Surv](#) for the format specification. Default for both binary classification and survival is “no” (do nothing).

`functionals.as` (character(1))
How to represents functional features? Option “matrix”: Keep them as matrix columns in the data.frame. Option “dfcols”: Convert them to individual numeric data.frame columns. Default is “dfcols”.

Value

Either a data.frame or a list with data.frame data and vector target.

See Also

Other task: [getTaskClassLevels\(\)](#), [getTaskCosts\(\)](#), [getTaskDesc\(\)](#), [getTaskFeatureNames\(\)](#), [getTaskFormula\(\)](#), [getTaskId\(\)](#), [getTaskNFeats\(\)](#), [getTaskSize\(\)](#), [getTaskTargetNames\(\)](#), [getTaskTargets\(\)](#), [getTaskType\(\)](#), [subsetTask\(\)](#)

Examples

```
library("mlbench")
data(BreastCancer)

df = BreastCancer
df$Id = NULL
task = makeClassifTask(id = "BreastCancer", data = df, target = "Class", positive = "malignant")
head(getTaskData)
head(getTaskData(task, features = c("Cell.size", "Cell.shape"), recode.target = "-1+1"))
head(getTaskData(task, subset = 1:100, recode.target = "01"))
```

getTaskDesc

Get a summarizing task description.

Description

See title.

Usage

```
getTaskDesc(x)
```

Arguments

x (Task | TaskDesc)
Task or its description object.

Value

ret_taskdesc

See Also

Other task: [getTaskClassLevels\(\)](#), [getTaskCosts\(\)](#), [getTaskData\(\)](#), [getTaskFeatureNames\(\)](#), [getTaskFormula\(\)](#), [getTaskId\(\)](#), [getTaskNFeats\(\)](#), [getTaskSize\(\)](#), [getTaskTargetNames\(\)](#), [getTaskTargets\(\)](#), [getTaskType\(\)](#), [subsetTask\(\)](#)

getTaskDescription *Deprecated, use [getTaskDesc](#) instead.*

Description

Deprecated, use [getTaskDesc](#) instead.

Usage

getTaskDescription(x)

Arguments

x (Task | TaskDesc)
Task or its description object.

getTaskFeatureNames *Get feature names of task.*

Description

Target column name is not included.

Usage

getTaskFeatureNames(task)

Arguments

task (Task)
The task.

Value

([character](#)).

See Also

Other task: [getTaskClassLevels\(\)](#), [getTaskCosts\(\)](#), [getTaskData\(\)](#), [getTaskDesc\(\)](#), [getTaskFormula\(\)](#), [getTaskId\(\)](#), [getTaskNFeats\(\)](#), [getTaskSize\(\)](#), [getTaskTargetNames\(\)](#), [getTaskTargets\(\)](#), [getTaskType\(\)](#), [subsetTask\(\)](#)

getTaskFormula	<i>Get formula of a task.</i>
----------------	-------------------------------

Description

This is usually simply `<target> ~ .`. For multilabel it is `<target_1> + ... + <target_k> ~ .`

Usage

```
getTaskFormula(
  x,
  target = getTaskTargetNames(x),
  explicit.features = FALSE,
  env = parent.frame()
)
```

Arguments

<code>x</code>	(Task TaskDesc) Task or its description object.
<code>target</code>	(character (1)) Left hand side of the formula. Default is defined by task <code>x</code> .
<code>explicit.features</code>	(logical (1)) Should the features (right hand side of the formula) be explicitly listed? Default is FALSE, i.e., they will be represented as <code>" . "</code> .
<code>env</code>	(environment) Environment of the formula. Default is <code>parent.frame()</code> .

Value

([formula](#)).

See Also

Other task: [getTaskClassLevels\(\)](#), [getTaskCosts\(\)](#), [getTaskData\(\)](#), [getTaskDesc\(\)](#), [getTaskFeatureNames\(\)](#), [getTaskId\(\)](#), [getTaskNFeats\(\)](#), [getTaskSize\(\)](#), [getTaskTargetNames\(\)](#), [getTaskTargets\(\)](#), [getTaskType\(\)](#), [subsetTask\(\)](#)

getTaskId	<i>Get the id of the task.</i>
-----------	--------------------------------

Description

See title.

Usage

```
getTaskId(x)
```

Arguments

x	(Task TaskDesc) Task or its description object.
---	--

Value

(character(1)).

See Also

Other task: [getTaskClassLevels\(\)](#), [getTaskCosts\(\)](#), [getTaskData\(\)](#), [getTaskDesc\(\)](#), [getTaskFeatureNames\(\)](#), [getTaskFormula\(\)](#), [getTaskNFeats\(\)](#), [getTaskSize\(\)](#), [getTaskTargetNames\(\)](#), [getTaskTargets\(\)](#), [getTaskType\(\)](#), [subsetTask\(\)](#)

getTaskNFeats	<i>Get number of features in task.</i>
---------------	--

Description

See title.

Usage

```
getTaskNFeats(x)
```

Arguments

x	(Task TaskDesc) Task or its description object.
---	--

Value

(integer(1)).

See Also

Other task: [getTaskClassLevels\(\)](#), [getTaskCosts\(\)](#), [getTaskData\(\)](#), [getTaskDesc\(\)](#), [getTaskFeatureNames\(\)](#), [getTaskFormula\(\)](#), [getTaskId\(\)](#), [getTaskSize\(\)](#), [getTaskTargetNames\(\)](#), [getTaskTargets\(\)](#), [getTaskType\(\)](#), [subsetTask\(\)](#)

getTaskSize	<i>Get number of observations in task.</i>
-------------	--

Description

See title.

Usage

```
getTaskSize(x)
```

Arguments

x	(Task TaskDesc) Task or its description object.
---	--

Value

(integer(1)).

See Also

Other task: [getTaskClassLevels\(\)](#), [getTaskCosts\(\)](#), [getTaskData\(\)](#), [getTaskDesc\(\)](#), [getTaskFeatureNames\(\)](#), [getTaskFormula\(\)](#), [getTaskId\(\)](#), [getTaskNFeats\(\)](#), [getTaskTargetNames\(\)](#), [getTaskTargets\(\)](#), [getTaskType\(\)](#), [subsetTask\(\)](#)

getTaskTargetNames	<i>Get the name(s) of the target column(s).</i>
--------------------	---

Description

NB: For multilabel, [getTaskTargetNames](#) and [getTaskClassLevels](#) actually return the same thing.

Usage

```
getTaskTargetNames(x)
```

Arguments

x	(Task TaskDesc) Task or its description object.
---	--

Value

(character).

See Also

Other task: [getTaskClassLevels\(\)](#), [getTaskCosts\(\)](#), [getTaskData\(\)](#), [getTaskDesc\(\)](#), [getTaskFeatureNames\(\)](#), [getTaskFormula\(\)](#), [getTaskId\(\)](#), [getTaskNFeats\(\)](#), [getTaskSize\(\)](#), [getTaskTargets\(\)](#), [getTaskType\(\)](#), [subsetTask\(\)](#)

<code>getTaskTargets</code>	<i>Get target data of task.</i>
-----------------------------	---------------------------------

Description

Get target data of task.

Usage

```
getTaskTargets(task, recode.target = "no")
```

Arguments

<code>task</code>	(Task)	The task.
<code>recode.target</code>	(character(1))	Should target classes be recoded? Supported are binary and multilabel classification and survival. Possible values for binary classification are “01”, “-1+1” and “drop.levels”. In the two latter cases the target vector is converted into a numeric vector. The positive class is coded as “+1” and the negative class either as “0” or “-1”. “drop.levels” will remove empty factor levels in the target column. In the multilabel case the logical targets can be converted to factors with “multilabel.factor”. For survival, you may choose to recode the survival times to “left”, “right” or “interval2” censored times using “lcens”, “rcens” or “icens”, respectively. See survival::Surv for the format specification. Default for both binary classification and survival is “no” (do nothing).

Value

A factor for classification or a numeric for regression, a data.frame of logical columns for multilabel.

See Also

Other task: [getTaskClassLevels\(\)](#), [getTaskCosts\(\)](#), [getTaskData\(\)](#), [getTaskDesc\(\)](#), [getTaskFeatureNames\(\)](#), [getTaskFormula\(\)](#), [getTaskId\(\)](#), [getTaskNFeats\(\)](#), [getTaskSize\(\)](#), [getTaskTargetNames\(\)](#), [getTaskType\(\)](#), [subsetTask\(\)](#)

Examples

```
task = makeClassifTask(data = iris, target = "Species")
getTaskTargets(task)
```

getTaskType	<i>Get the type of the task.</i>
-------------	----------------------------------

Description

See title.

Usage

```
getTaskType(x)
```

Arguments

x [\(Task | TaskDesc\)](#)
Task or its description object.

Value

(character(1)).

See Also

Other task: [getTaskClassLevels\(\)](#), [getTaskCosts\(\)](#), [getTaskData\(\)](#), [getTaskDesc\(\)](#), [getTaskFeatureNames\(\)](#), [getTaskFormula\(\)](#), [getTaskId\(\)](#), [getTaskNFeats\(\)](#), [getTaskSize\(\)](#), [getTaskTargetNames\(\)](#), [getTaskTargets\(\)](#), [subsetTask\(\)](#)

getTuneResult	<i>Returns the optimal hyperparameters and optimization path after training.</i>
---------------	--

Description

Returns the optimal hyperparameters and optimization path after training.

Usage

```
getTuneResult(object)
```

Arguments

object [\(WrappedModel\)](#)
Trained Model created with [makeTuneWrapper](#).

Value

([TuneResult](#)).

See Also

Other tune: [TuneControl](#), [getNestedTuneResultsOptPathDf\(\)](#), [getNestedTuneResultsX\(\)](#), [getResamplingIndices\(\)](#), [makeModelMultiplexerParamSet\(\)](#), [makeModelMultiplexer\(\)](#), [makeTuneControlCMAES\(\)](#), [makeTuneControlDesign\(\)](#), [makeTuneControlGenSA\(\)](#), [makeTuneControlGrid\(\)](#), [makeTuneControlIrace\(\)](#), [makeTuneControlMBO\(\)](#), [makeTuneControlRandom\(\)](#), [makeTuneWrapper\(\)](#), [tuneParams\(\)](#), [tuneThreshold\(\)](#)

`getTuneResultOptPath` *Get the optimization path of a tuning result.*

Description

Returns the `opt.path` from a ([TuneResult](#)) object.

Usage

```
getTuneResultOptPath(tune.result, as.df = TRUE)
```

Arguments

<code>tune.result</code>	(TuneResult) A tuning result of the (tuneParams) function.
<code>as.df</code>	(<code>logical(1)</code>) Should the optimization path be returned as a data frame? Default is TRUE.

Value

([ParamHelpers::OptPath](#)) or (`data.frame`).

`gunpoint.task` *Gunpoint functional data classification task.*

Description

Contains the task (`gunpoint.task`). You have to classify whether a person raises up a gun or just an empty hand.

References

See Ratanamahatana, C. A. & Keogh. E. (2004). Everything you know about Dynamic Time Warping is Wrong. Proceedings of SIAM International Conference on Data Mining (SDM05), 506-510.

hasFunctionalFeatures *Check whether the object contains functional features.*

Description

See title.

Usage

hasFunctionalFeatures(obj)

Arguments

obj	(Task TaskDesc data.frame) Object to check.
-----	--

Value

(logical(1))

hasProperties *Deprecated, use hasLearnerProperties instead.*

Description

Deprecated, use hasLearnerProperties instead.

Usage

hasProperties(learner, props)

Arguments

learner	Deprecated.
props	Deprecated.

helpLearner	<i>Access help page of learner functions.</i>
-------------	---

Description

Interactive function that gives the user quick access to the help pages associated with various functions involved in the given learner.

Usage

```
helpLearner(learner)
```

Arguments

learner	(Learner character(1)) The learner. If you pass a string the learner will be created via makeLearner .
---------	--

See Also

Other learner: [LearnerProperties](#), [getClassWeightParam\(\)](#), [getHyperPars\(\)](#), [getLearnerId\(\)](#), [getLearnerNote\(\)](#), [getLearnerPackages\(\)](#), [getLearnerParVals\(\)](#), [getLearnerParamSet\(\)](#), [getLearnerPredictType\(\)](#), [getLearnerShortName\(\)](#), [getLearnerType\(\)](#), [getParamSet\(\)](#), [helpLearnerParam\(\)](#), [makeLearners\(\)](#), [makeLearner\(\)](#), [removeHyperPars\(\)](#), [setHyperPars\(\)](#), [setId\(\)](#), [setLearnerId\(\)](#), [setPredictThreshold\(\)](#), [setPredictType\(\)](#)

Other help: [helpLearnerParam\(\)](#)

helpLearnerParam	<i>Get specific help for a learner's parameters.</i>
------------------	--

Description

Print the description of parameters of a given learner. The description is automatically extracted from the help pages of the learner, so it may be incomplete.

Usage

```
helpLearnerParam(learner, param = NULL)
```

Arguments

learner	(Learner character(1)) The learner. If you pass a string the learner will be created via makeLearner .
param	(character NULL) Parameter(s) to describe. Defaults to NULL, which prints information on the documentation status of all parameters.

See Also

Other learner: [LearnerProperties](#), [getClassWeightParam\(\)](#), [getHyperPars\(\)](#), [getLearnerId\(\)](#), [getLearnerNote\(\)](#), [getLearnerPackages\(\)](#), [getLearnerParVals\(\)](#), [getLearnerParamSet\(\)](#), [getLearnerPredictType\(\)](#), [getLearnerShortName\(\)](#), [getLearnerType\(\)](#), [getParamSet\(\)](#), [helpLearner\(\)](#), [makeLearners\(\)](#), [makeLearner\(\)](#), [removeHyperPars\(\)](#), [setHyperPars\(\)](#), [setId\(\)](#), [setLearnerId\(\)](#), [setPredictThreshold\(\)](#), [setPredictType\(\)](#)

Other help: [helpLearner\(\)](#)

imputations

Built-in imputation methods.

Description

The built-ins are:

- `imputeConstant(const)` for imputation using a constant value,
- `imputeMedian()` for imputation using the median,
- `imputeMode()` for imputation using the mode,
- `imputeMin(multiplier)` for imputing constant values shifted below the minimum using $\min(x) - \text{multiplier} * \text{diff}(\text{range}(x))$,
- `imputeMax(multiplier)` for imputing constant values shifted above the maximum using $\max(x) + \text{multiplier} * \text{diff}(\text{range}(x))$,
- `imputeNormal(mean, sd)` for imputation using normally distributed random values. Mean and standard deviation will be calculated from the data if not provided.
- `imputeHist(breaks, use.mids)` for imputation using random values with probabilities calculated using `table` or `hist`.
- `imputeLearner(learner, features = NULL)` for imputations using the response of a classification or regression learner.

Usage

```
imputeConstant(const)
```

```
imputeMedian()
```

```
imputeMean()
```

```
imputeMode()
```

```
imputeMin(multiplier = 1)
```

```
imputeMax(multiplier = 1)
```

```
imputeUniform(min = NA_real_, max = NA_real_)
```

```
imputeNormal(mu = NA_real_, sd = NA_real_)
```

```
imputeHist(breaks, use.mids = TRUE)
```

```
imputeLearner(learner, features = NULL)
```

Arguments

const	(any) Constant valued use for imputation.
multiplier	(numeric(1)) Value that stored minimum or maximum is multiplied with when imputation is done.
min	(numeric(1)) Lower bound for uniform distribution. If NA (default), it will be estimated from the data.
max	(numeric(1)) Upper bound for uniform distribution. If NA (default), it will be estimated from the data.
mu	(numeric(1)) Mean of normal distribution. If missing it will be estimated from the data.
sd	(numeric(1)) Standard deviation of normal distribution. If missing it will be estimated from the data.
breaks	(numeric(1)) Number of breaks to use in graphics::hist . If missing, defaults to auto-detection via “Sturges”.
use.mids	(logical(1)) If x is numeric and a histogram is used, impute with bin mids (default) or instead draw uniformly distributed samples within bin range.
learner	(Learner character(1)) Supervised learner. Its predictions will be used for imputations. If you pass a string the learner will be created via makeLearner . Note that the target column is not available for this operation.
features	(character) Features to use in learner for prediction. Default is NULL which uses all available features except the target column of the original task.

See Also

Other impute: [impute\(\)](#), [makeImputeMethod\(\)](#), [makeImputeWrapper\(\)](#), [reimpute\(\)](#)

 impute

Impute and re-impute data

Description

Allows imputation of missing feature values through various techniques. Note that you have the possibility to re-impute a data set in the same way as the imputation was performed during training. This especially comes in handy during resampling when one wants to perform the same imputation on the test set as on the training set.

The function `impute` performs the imputation on a data set and returns, alongside with the imputed data set, an “ImputationDesc” object which can contain “learned” coefficients and helpful data. It can then be passed together with a new data set to [reimpute](#).

The imputation techniques can be specified for certain features or for feature classes, see function arguments.

You can either provide an arbitrary object, use a built-in imputation method listed under [imputations](#) or create one yourself using [makeImputeMethod](#).

Usage

```
impute(
  obj,
  target = character(0L),
  classes = list(),
  cols = list(),
  dummy.classes = character(0L),
  dummy.cols = character(0L),
  dummy.type = "factor",
  force.dummies = FALSE,
  impute.new.levels = TRUE,
  recode.factor.levels = TRUE
)
```

Arguments

<code>obj</code>	(data.frame Task) Input data.
<code>target</code>	(character) Name of the column(s) specifying the response. Default is <code>character(0)</code> .
<code>classes</code>	(named list) Named list containing imputation techniques for classes of columns. E.g. <code>list(numeric = imputeMedian())</code> .
<code>cols</code>	(named list) Named list containing names of imputation methods to impute missing values in the data column referenced by the list element’s name. Overrides imputation set via classes.

<code>dummy.classes</code>	(character) Classes of columns to create dummy columns for. Default is <code>character(0)</code> .
<code>dummy.cols</code>	(character) Column names to create dummy columns (containing binary missing indicator) for. Default is <code>character(0)</code> .
<code>dummy.type</code>	(character(1)) How dummy columns are encoded. Either as 0/1 with type “numeric” or as “factor”. Default is “factor”.
<code>force.dummies</code>	(logical(1)) Force dummy creation even if the respective data column does not contain any NAs. Note that (a) most learners will complain about constant columns created this way but (b) your feature set might be stochastic if you turn this off. Default is FALSE.
<code>impute.new.levels</code>	(logical(1)) If new, unencountered factor level occur during reimputation, should these be handled as NAs and then be imputed the same way? Default is TRUE.
<code>recode.factor.levels</code>	(logical(1)) Recode factor levels after reimputation, so they match the respective element of <code>lvls</code> (in the description object) and therefore match the levels of the feature factor in the training data after imputation?. Default is TRUE.

Details

The description object contains these slots

target ([character](#)) See argument.

features ([character](#)) Feature names (column names of data),.

classes ([character](#)) Feature classes (storage type of data).

lvls ([named list](#)) Mapping of column names of factor features to their levels, including newly created ones during imputation.

impute ([named list](#)) Mapping of column names to imputation functions.

dummies ([named list](#)) Mapping of column names to imputation functions.

impute.new.levels ([logical\(1\)](#)) See argument.

recode.factor.levels ([logical\(1\)](#)) See argument.

Value

([list](#))

`data` ([data.frame](#))

Imputed data.

`desc` ([ImputationDesc](#))

Description object.

See Also

Other impute: [imputations](#), [makeImputeMethod\(\)](#), [makeImputeWrapper\(\)](#), [reimpute\(\)](#)

Examples

```
df = data.frame(x = c(1, 1, NA), y = factor(c("a", "a", "b")), z = 1:3)
imputed = impute(df, target = character(0), cols = list(x = 99, y = imputeMode()))
print(imputed$data)
reimpute(data.frame(x = NA_real_), imputed$desc)
```

iris.task	<i>Iris classification task.</i>
-----------	----------------------------------

Description

Contains the task (iris.task).

References

See [datasets::iris](#).

isFailureModel	<i>Is the model a FailureModel?</i>
----------------	-------------------------------------

Description

Such a model is created when one sets the corresponding option in [configureMlr](#).

For complex wrappers this getter returns TRUE if ANY model contained in it failed.

Usage

```
isFailureModel(model)
```

Arguments

model	(WrappedModel) The model.
-------	--

Value

(logical(1)).

joinClassLevels	<i>Join some class existing levels to new, larger class levels for classification problems.</i>
-----------------	---

Description

Join some class existing levels to new, larger class levels for classification problems.

Usage

```
joinClassLevels(task, new.levels)
```

Arguments

task	(Task) The task.
new.levels	(list of character) Element names specify the new class levels to create, while the corresponding element character vector specifies the existing class levels which will be joined to the new one.

Value

[Task](#).

Examples

```
joinClassLevels(iris.task, new.levels = list(foo = c("setosa", "virginica")))
```

learnerArgsToControl	<i>Convert arguments to control structure.</i>
----------------------	--

Description

Find all elements in ... which are not missing and call control on them.

Usage

```
learnerArgsToControl(control, ...)
```

Arguments

control	(function) Function that creates control structure.
...	(any) Arguments for control structure function.

Value

Control structure for learner.

LearnerProperties *Query properties of learners.*

Description

Properties can be accessed with `getLearnerProperties(learner)`, which returns a character vector.

The learner properties are defined as follows:

numerics, factors, ordered Can numeric, factor or ordered factor features be handled?

functionals Can an arbitrary number of functional features be handled?

single.functional Can exactly one functional feature be handled?

missings Can missing values in features be handled?

weights Can observations be weighted during fitting?

oneclas, twoclass, multiclass Only for `classif`: Can one-class, two-class or multi-class classification problems be handled?

class.weights Only for `classif`: Can class weights be handled?

rcens, lcens, icens Only for `surv`: Can right, left, or interval censored data be handled?

prob For `classif`, `cluster`, `multilabel`, `surv`: Can probabilities be predicted?

se Only for `regr`: Can standard errors be predicted?

oobpreds Only for `classif`, `regr` and `surv`: Can out of bag predictions be extracted from the trained model?

featimp For `classif`, `regr`, `surv`: Does the model support extracting information on feature importance?

Usage

```
getLearnerProperties(learner)
```

```
hasLearnerProperties(learner, props)
```

Arguments

`learner` ([Learner](#) | `character(1)`)
The learner. If you pass a string the learner will be created via [makeLearner](#).

`props` ([character](#))
Vector of properties to query.

Value

`getLearnerProperties` returns a character vector with learner properties. `hasLearnerProperties` returns a logical vector of the same length as props.

See Also

Other learner: `getClassWeightParam()`, `getHyperPars()`, `getLearnerId()`, `getLearnerNote()`, `getLearnerPackages()`, `getLearnerParVals()`, `getLearnerParamSet()`, `getLearnerPredictType()`, `getLearnerShortName()`, `getLearnerType()`, `getParamSet()`, `helpLearnerParam()`, `helpLearner()`, `makeLearners()`, `makeLearner()`, `removeHyperPars()`, `setHyperPars()`, `setId()`, `setLearnerId()`, `setPredictThreshold()`, `setPredictType()`

learners

List of supported learning algorithms.

Description

All supported learners can be found by `listLearners` or as a table in the tutorial appendix: https://mlr.mlr-org.com/articles/tutorial/integrated_learners.html.

listFilterEnsembleMethods

List ensemble filter methods.

Description

Returns a subset-able dataframe with filter information.

Usage

```
listFilterEnsembleMethods(desc = TRUE)
```

Arguments

`desc` (logical(1))
Provide more detailed information about filters. Default is TRUE.

Value

([data.frame](#)).

See Also

Other filter: `filterFeatures()`, `generateFilterValuesData()`, `getFilteredFeatures()`, `listFilterMethods()`, `makeFilterEnsemble()`, `makeFilterWrapper()`, `makeFilter()`, `plotFilterValues()`

listFilterMethods	<i>List filter methods.</i>
-------------------	-----------------------------

Description

Returns a subset-able dataframe with filter information.

Usage

```
listFilterMethods(  
  desc = TRUE,  
  tasks = FALSE,  
  features = FALSE,  
  include.deprecated = FALSE  
)
```

Arguments

desc	(logical(1)) Provide more detailed information about filters. Default is TRUE.
tasks	(logical(1)) Provide information on supported tasks. Default is FALSE.
features	(logical(1)) Provide information on supported features. Default is FALSE.
include.deprecated	(logical(1)) Should deprecated filter methods be included in the list. Default is FALSE.

Value

([data.frame](#)).

See Also

Other filter: [filterFeatures\(\)](#), [generateFilterValuesData\(\)](#), [getFilteredFeatures\(\)](#), [listFilterEnsembleMethod](#), [makeFilterEnsemble\(\)](#), [makeFilterWrapper\(\)](#), [makeFilter\(\)](#), [plotFilterValues\(\)](#)

listLearnerProperties *List the supported learner properties*

Description

This is useful for determining which learner properties are available.

Usage

```
listLearnerProperties(type = "any")
```

Arguments

type (character(1))
Only return properties for a specified task type. Default is “any”.

Value

(character).

listLearners *Find matching learning algorithms.*

Description

Returns learning algorithms which have specific characteristics, e.g. whether they support missing values, case weights, etc.

Note that the packages of all learners are loaded during the search if you create them. This can be a lot. If you do not create them we only inspect properties of the S3 classes. This will be a lot faster.

Note that for general cost-sensitive learning, mlr currently supports mainly “wrapper” approaches like [CostSensWeightedPairsWrapper](#), which are not listed, as they are not basic R learning algorithms. The same applies for many multilabel methods, see, e.g., [makeMultilabelBinaryRelevanceWrapper](#).

Usage

```
listLearners(  
  obj = NA_character_,  
  properties = character(0L),  
  quiet = TRUE,  
  warn.missing.packages = TRUE,  
  check.packages = FALSE,  
  create = FALSE  
)
```

```

## Default S3 method:
listLearners(
  obj = NA_character_,
  properties = character(0L),
  quiet = TRUE,
  warn.missing.packages = TRUE,
  check.packages = FALSE,
  create = FALSE
)

## S3 method for class 'character'
listLearners(
  obj = NA_character_,
  properties = character(0L),
  quiet = TRUE,
  warn.missing.packages = TRUE,
  check.packages = FALSE,
  create = FALSE
)

## S3 method for class 'Task'
listLearners(
  obj = NA_character_,
  properties = character(0L),
  quiet = TRUE,
  warn.missing.packages = TRUE,
  check.packages = TRUE,
  create = FALSE
)

```

Arguments

obj	(character(1) Task) Either character(1) task or the type of the task, in the latter case one of: “clas- sif” “regr” “surv” “costsens” “cluster” “multilabel”. Default is NA matching all types.
properties	(character) Set of required properties to filter for. Default is character(0).
quiet	(logical(1)) Construct learners quietly to check their properties, shows no package startup messages. Turn off if you suspect errors. Default is TRUE.
warn.missing.packages	(logical(1)) If some learner cannot be constructed because its package is missing, should a warning be shown? Default is TRUE.
check.packages	(logical(1)) Check if required packages are installed. Calls find.package(). If create is TRUE, this is done implicitly and the value of this parameter is ignored. If

create is FALSE and check.packages is TRUE the returned table only contains learners whose dependencies are installed. If check.packages set to FALSE, learners that cannot actually be constructed because of missing packages may be returned. Default is FALSE.

create (logical(1))
Instantiate objects (or return info table)? Packages are loaded if and only if this option is TRUE. Default is FALSE.

Value

([data.frame|list' of [Learner](#)). Either a descriptive data.frame that allows access to all properties of the learners or a list of created learner objects (named by ids of listed learners).

Examples

```
## Not run:  
listLearners("classif", properties = c("multiclass", "prob"))  
data = iris  
task = makeClassifTask(data = data, target = "Species")  
listLearners(task)  
  
## End(Not run)
```

listMeasureProperties *List the supported measure properties.*

Description

This is useful for determining which measure properties are available.

Usage

```
listMeasureProperties()
```

Value

([character](#)).

listMeasures	<i>Find matching measures.</i>
--------------	--------------------------------

Description

Returns the matching measures which have specific characteristics, e.g. whether they supports classification or regression.

Usage

```
listMeasures(obj, properties = character(0L), create = FALSE)

## Default S3 method:
listMeasures(obj, properties = character(0L), create = FALSE)

## S3 method for class 'character'
listMeasures(obj, properties = character(0L), create = FALSE)

## S3 method for class 'Task'
listMeasures(obj, properties = character(0L), create = FALSE)
```

Arguments

obj	(character(1) Task) Either character(1) task or the type of the task, in the latter case one of: “classif” “regr” “surv” “costsens” “cluster” “multilabel”. Default is NA matching all types.
properties	(character) Set of required properties to filter for. See Measure for some standardized properties. Default is character(0).
create	(logical(1)) Instantiate objects (or return strings)? Default is FALSE.

Value

([character|list' of [Measure](#)). Class names of matching measures or instantiated objects.

listTaskTypes	<i>List the supported task types in mlr</i>
---------------	---

Description

Returns a character vector with each of the supported task types in mlr.

Usage

```
listTaskTypes()
```

Value

([character](#)).

lung.task	<i>NCCTG Lung Cancer survival task.</i>
-----------	---

Description

Contains the task (lung.task).

References

See [survival::lung](#). Incomplete cases have been removed from the task.

makeAggregation	<i>Specify your own aggregation of measures.</i>
-----------------	--

Description

This is an advanced feature of mlr. It gives access to some inner workings so the result might not be compatible with everything!

Usage

```
makeAggregation(id, name = id, properties, fun)
```

Arguments

id	(character (1)) Name of the aggregation method (preferably the same name as the generated function).
name	(character (1)) Long name of the aggregation method. Default is id.
properties	(character) Set of aggregation properties. req.train Are prediction or train sets required to calculate the aggregation? req.test Are prediction or test sets required to calculate the aggregation?
fun	(function(task, perf.test, perf.train, measure, group, pred)) Calculates the aggregated performance. In most cases you will only need the performances <code>perf.test</code> and optionally <code>perf.train</code> on the test and training data sets.

task (**Task**) The task.
 perf.test (**numeric**) performance results on the test data sets.
 perf.train (**numeric**) performance results on the training data sets.
 measure (**Measure**) Performance measure.
 group (**factor**) Grouping of resampling iterations. This encodes whether specific iterations 'belong together' (e.g. repeated CV).
 pred (**Prediction**) Prediction object.

Value

(Aggregation).

See Also

[aggregations](#), [setAggregation](#)

Examples

```

# computes the interquartile range on all performance values
test.iqr = makeAggregation(
  id = "test.iqr", name = "Test set interquartile range",
  properties = "req.test",
  fun = function(task, perf.test, perf.train, measure, group, pred) IQR(perf.test)
)

```

makeBaggingWrapper *Fuse learner with the bagging technique.*

Description

Fuses a learner with the bagging method (i.e., similar to what a randomForest does). Creates a learner object, which can be used like any other learner object. Models can easily be accessed via [getLearnerModel](#).

Bagging is implemented as follows: For each iteration a random data subset is sampled (with or without replacement) and potentially the number of features is also restricted to a random subset. Note that this is usually handled in a slightly different way in the random forest where features are sampled at each tree split).

Prediction works as follows: For classification we do majority voting to create a discrete label and probabilities are predicted by considering the proportions of all predicted labels. For regression the mean value and the standard deviations across predictions is computed.

Note that the passed base learner must always have `predict.type = 'response'`, while the BaggingWrapper can estimate probabilities and standard errors, so it can be set, e.g., to `predict.type = 'prob'`. For this reason, when you call [setPredictType](#), the type is only set for the BaggingWrapper, not passed down to the inner learner.

Usage

```
makeBaggingWrapper(
  learner,
  bw.iters = 10L,
  bw.replace = TRUE,
  bw.size,
  bw.feats = 1
)
```

Arguments

learner	(Learner character(1)) The learner. If you pass a string the learner will be created via makeLearner .
bw.iters	(integer(1)) Iterations = number of fitted models in bagging. Default is 10.
bw.replace	(logical(1)) Sample bags with replacement (bootstrapping)? Default is TRUE.
bw.size	(numeric(1)) Percentage size of sampled bags. Default is 1 for bootstrapping and 0.632 for subsampling.
bw.feats	(numeric(1)) Percentage size of randomly selected features in bags. Default is 1. At least one feature will always be selected.

Value

[Learner](#).

See Also

Other wrapper: [makeClassificationViaRegressionWrapper\(\)](#), [makeConstantClassWrapper\(\)](#), [makeCostSensClassifWrapper\(\)](#), [makeCostSensRegrWrapper\(\)](#), [makeDownsampleWrapper\(\)](#), [makeDummyFeaturesWrapper\(\)](#), [makeExtractFDAFeatsWrapper\(\)](#), [makeFeatSelWrapper\(\)](#), [makeFilterWrapper\(\)](#), [makeImputeWrapper\(\)](#), [makeMulticlassWrapper\(\)](#), [makeMultilabelBinaryRelevanceWrapper\(\)](#), [makeMultilabelClassifierChainsWrapper\(\)](#), [makeMultilabelDBRWrapper\(\)](#), [makeMultilabelNestedStackingWrapper\(\)](#), [makeMultilabelStackingWrapper\(\)](#), [makeOverBaggingWrapper\(\)](#), [makePreprocWrapperCaret\(\)](#), [makePreprocWrapper\(\)](#), [makeRemoveConstantFeaturesWrapper\(\)](#), [makeSMOTEWrapper\(\)](#), [makeTuneWrapper\(\)](#), [makeUndersampleWrapper\(\)](#), [makeWeightedClassesWrapper\(\)](#)

makeClassificationViaRegressionWrapper

Classification via regression wrapper.

Description

Builds regression models that predict for the positive class whether a particular example belongs to it (1) or not (-1).

Probabilities are generated by transforming the predictions with a softmax.

Inspired by WEKA's ClassificationViaRegression (<http://weka.sourceforge.net/doc.dev/weka/classifiers/meta/ClassificationV>)

Usage

```
makeClassificationViaRegressionWrapper(learner, predict.type = "response")
```

Arguments

learner	(Learner character(1)) The learner. If you pass a string the learner will be created via makeLearner .
predict.type	(character(1)) "response" (= labels) or "prob" (= probabilities and labels by selecting the one with maximal probability).

Value

[Learner](#).

See Also

Other wrapper: [makeBaggingWrapper\(\)](#), [makeConstantClassWrapper\(\)](#), [makeCostSensClassifWrapper\(\)](#), [makeCostSensRegrWrapper\(\)](#), [makeDownsampleWrapper\(\)](#), [makeDummyFeaturesWrapper\(\)](#), [makeExtractFDAFeatsWrap](#), [makeFeatSelWrapper\(\)](#), [makeFilterWrapper\(\)](#), [makeImputeWrapper\(\)](#), [makeMulticlassWrapper\(\)](#), [makeMultilabelBinaryRelevanceWrapper\(\)](#), [makeMultilabelClassifierChainsWrapper\(\)](#), [makeMultilabelDBRWrap](#), [makeMultilabelNestedStackingWrapper\(\)](#), [makeMultilabelStackingWrapper\(\)](#), [makeOverBaggingWrapper\(\)](#), [makePreprocWrapperCaret\(\)](#), [makePreprocWrapper\(\)](#), [makeRemoveConstantFeaturesWrapper\(\)](#), [makeSMOTEWrapper\(\)](#), [makeTuneWrapper\(\)](#), [makeUndersampleWrapper\(\)](#), [makeWeightedClassesWrapper\(\)](#)

Examples

```
lrn = makeLearner("regr.rpart")
lrn = makeClassificationViaRegressionWrapper(lrn)
mod = train(lrn, sonar.task, subset = 1:140)
predictions = predict(mod, newdata = getTaskData(sonar.task)[141:208, 1:60])
```

makeClassifTask	<i>Create a classification task.</i>
-----------------	--------------------------------------

Description

Create a classification task.

Usage

```
makeClassifTask(
  id = deparse(substitute(data)),
  data,
  target,
  weights = NULL,
  blocking = NULL,
  coordinates = NULL,
  positive = NA_character_,
  fixup.data = "warn",
  check.data = TRUE
)
```

Arguments

id	(character(1)) Id string for object. Default is the name of the R variable passed to data.
data	(data.frame) A data frame containing the features and target variable(s).
target	(character(1) character(2) character(n.classes)) Name(s) of the target variable(s). For survival analysis these are the names of the survival time and event columns, so it has length 2. For multilabel classification it contains the names of the logical columns that encode whether a label is present or not and its length corresponds to the number of classes.
weights	(numeric) Optional, non-negative case weight vector to be used during fitting. Cannot be set for cost-sensitive learning. Default is NULL which means no (= equal) weights.
blocking	(factor) An optional factor of the same length as the number of observations. Observations with the same blocking level “belong together”. Specifically, they are either put all in the training or the test set during a resampling iteration. Default is NULL which means no blocking.
coordinates	(data.frame) Coordinates of a spatial data set that will be used for spatial partitioning of the data in a spatial cross-validation resampling setting. Coordinates have to be numeric values. Provided data.frame needs to have the same number of rows as data and consist of at least two dimensions.
positive	(character(1)) Positive class for binary classification (otherwise ignored and set to NA). Default is the first factor level of the target attribute.
fixup.data	(character(1)) Should some basic cleaning up of data be performed? Currently this means removing empty factor levels for the columns. Possible choices are: “no” = Don’t do it. “warn” = Do it but warn about it. “quiet” = Do it but keep silent. Default is “warn”.

check.data (logical(1))
Should sanity of data be checked initially at task creation? You should have good reasons to turn this off (one might be speed). Default is TRUE.

See Also

[Task](#) [CostSensTask](#) [ClusterTask](#) [MultilabelTask](#) [RegrTask](#) [SurvTask](#)

makeClusterTask *Create a cluster task.*

Description

Create a cluster task.

Usage

```
makeClusterTask(
  id = deparse(substitute(data)),
  data,
  weights = NULL,
  blocking = NULL,
  coordinates = NULL,
  fixup.data = "warn",
  check.data = TRUE
)
```

Arguments

id (character(1))
Id string for object. Default is the name of the R variable passed to data.

data ([data.frame](#))
A data frame containing the features and target variable(s).

weights ([numeric](#))
Optional, non-negative case weight vector to be used during fitting. Cannot be set for cost-sensitive learning. Default is NULL which means no (= equal) weights.

blocking ([factor](#))
An optional factor of the same length as the number of observations. Observations with the same blocking level “belong together”. Specifically, they are either put all in the training or the test set during a resampling iteration. Default is NULL which means no blocking.

coordinates ([data.frame](#))
Coordinates of a spatial data set that will be used for spatial partitioning of the data in a spatial cross-validation resampling setting. Coordinates have to be numeric values. Provided [data.frame](#) needs to have the same number of rows as data and consist of at least two dimensions.

fixup.data	(character(1)) Should some basic cleaning up of data be performed? Currently this means removing empty factor levels for the columns. Possible choices are: “no” = Don’t do it. “warn” = Do it but warn about it. “quiet” = Do it but keep silent. Default is “warn”.
check.data	(logical(1)) Should sanity of data be checked initially at task creation? You should have good reasons to turn this off (one might be speed). Default is TRUE.

See Also

[Task](#) [ClassifTask](#) [CostSensTask](#) [MultilabelTask](#) [RegrTask](#) [SurvTask](#)

makeConstantClassWrapper

Wraps a classification learner to support problems where the class label is (almost) constant.

Description

If the training data contains only a single class (or almost only a single class), this wrapper creates a model that always predicts the constant class in the training data. In all other cases, the underlying learner is trained and the resulting model used for predictions.

Probabilities can be predicted and will be 1 or 0 depending on whether the label matches the majority class or not.

Usage

```
makeConstantClassWrapper(learner, frac = 0)
```

Arguments

learner	(Learner character(1)) The learner. If you pass a string the learner will be created via makeLearner .
frac	numeric(1) The fraction of labels in [0, 1) that can be different from the majority label. Default is 0, which means that constant labels are only predicted if there is exactly one label in the data.

Value

[Learner](#).

See Also

Other wrapper: [makeBaggingWrapper\(\)](#), [makeClassificationViaRegressionWrapper\(\)](#), [makeCostSensClassifWrapper\(\)](#), [makeCostSensRegrWrapper\(\)](#), [makeDownsampleWrapper\(\)](#), [makeDummyFeaturesWrapper\(\)](#), [makeExtractFDAFeatsWrapper\(\)](#), [makeFeatSelWrapper\(\)](#), [makeFilterWrapper\(\)](#), [makeImputeWrapper\(\)](#), [makeMulticlassWrapper\(\)](#), [makeMultilabelBinaryRelevanceWrapper\(\)](#), [makeMultilabelClassifierChainsWrapper\(\)](#), [makeMultilabelDBRWrapper\(\)](#), [makeMultilabelNestedStackingWrapper\(\)](#), [makeMultilabelStackingWrapper\(\)](#), [makeOverBaggingWrapper\(\)](#), [makePreprocWrapperCaret\(\)](#), [makePreprocWrapper\(\)](#), [makeRemoveConstantFeaturesWrapper\(\)](#), [makeSMOTERWrapper\(\)](#), [makeTuneWrapper\(\)](#), [makeUndersampleWrapper\(\)](#), [makeWeightedClassesWrapper\(\)](#)

makeCostMeasure	<i>Creates a measure for non-standard misclassification costs.</i>
-----------------	--

Description

Creates a cost measure for non-standard classification error costs.

Usage

```
makeCostMeasure(
  id = "costs",
  minimize = TRUE,
  costs,
  combine = mean,
  best = NULL,
  worst = NULL,
  name = id,
  note = ""
)
```

Arguments

id	(character(1)) Name of measure. Default is “costs”.
minimize	(logical(1)) Should the measure be minimized? Otherwise you are effectively specifying a benefits matrix. Default is TRUE.
costs	(matrix) Matrix of misclassification costs. Rows and columns have to be named with class labels, order does not matter. Rows indicate true classes, columns predicted classes.
combine	(function) How to combine costs over all cases for a SINGLE test set? Note this is not the same as the aggregate argument in makeMeasure . You can set this as well via setAggregation , as for any measure. Default is mean .
best	(numeric(1)) Best obtainable value for measure. Default is -Inf or Inf, depending on minimize.

worst	(numeric(1)) Worst obtainable value for measure. Default is Inf or -Inf, depending on minimize.
name	(character) Name of the measure. Default is id.
note	(character) Description and additional notes for the measure. Default is "".

Value

[Measure](#).

See Also

Other performance: [ConfusionMatrix](#), [calculateConfusionMatrix\(\)](#), [calculateROCMeasures\(\)](#), [estimateRelativeOverfitting\(\)](#), [makeCustomResampledMeasure\(\)](#), [makeMeasure\(\)](#), [measures](#), [performance\(\)](#), [setAggregation\(\)](#), [setMeasurePars\(\)](#)

makeCostSensClassifWrapper

Wraps a classification learner for use in cost-sensitive learning.

Description

Creates a wrapper, which can be used like any other learner object. The classification model can easily be accessed via [getLearnerModel](#).

This is a very naive learner, where the costs are transformed into classification labels - the label for each case is the name of class with minimal costs. (If ties occur, the label which is better on average w.r.t. costs over all training data is preferred.) Then the classifier is fitted to that data and subsequently used for prediction.

Usage

```
makeCostSensClassifWrapper(learner)
```

Arguments

learner	(Learner character(1)) The classification learner. If you pass a string the learner will be created via makeLearner .
---------	---

Value

[Learner](#).

See Also

Other costsens: [makeCostSensRegrWrapper\(\)](#), [makeCostSensTask\(\)](#), [makeCostSensWeightedPairsWrapper\(\)](#)

Other wrapper: [makeBaggingWrapper\(\)](#), [makeClassificationViaRegressionWrapper\(\)](#), [makeConstantClassWrapper\(\)](#), [makeCostSensRegrWrapper\(\)](#), [makeDownsampleWrapper\(\)](#), [makeDummyFeaturesWrapper\(\)](#), [makeExtractFDAFeatsWrapper\(\)](#), [makeFeatSelWrapper\(\)](#), [makeFilterWrapper\(\)](#), [makeImputeWrapper\(\)](#), [makeMulticlassWrapper\(\)](#), [makeMultilabelBinaryRelevanceWrapper\(\)](#), [makeMultilabelClassifierChainsWrapper\(\)](#), [makeMultilabelDBRWrapper\(\)](#), [makeMultilabelNestedStackingWrapper\(\)](#), [makeMultilabelStackingWrapper\(\)](#), [makeOverBaggingWrapper\(\)](#), [makePreprocWrapperCaret\(\)](#), [makePreprocWrapper\(\)](#), [makeRemoveConstantFeaturesWrapper\(\)](#), [makeSMOTEWrapper\(\)](#), [makeTuneWrapper\(\)](#), [makeUndersampleWrapper\(\)](#), [makeWeightedClassesWrapper\(\)](#)

makeCostSensRegrWrapper

Wraps a regression learner for use in cost-sensitive learning.

Description

Creates a wrapper, which can be used like any other learner object. Models can easily be accessed via [getLearnerModel](#).

For each class in the task, an individual regression model is fitted for the costs of that class. During prediction, the class with the lowest predicted costs is selected.

Usage

```
makeCostSensRegrWrapper(learner)
```

Arguments

learner ([Learner](#) | character(1))
The regression learner. If you pass a string the learner will be created via [makeLearner](#).

Value

[Learner](#).

See Also

Other costsens: [makeCostSensClassifWrapper\(\)](#), [makeCostSensTask\(\)](#), [makeCostSensWeightedPairsWrapper\(\)](#)

Other wrapper: [makeBaggingWrapper\(\)](#), [makeClassificationViaRegressionWrapper\(\)](#), [makeConstantClassWrapper\(\)](#), [makeCostSensClassifWrapper\(\)](#), [makeDownsampleWrapper\(\)](#), [makeDummyFeaturesWrapper\(\)](#), [makeExtractFDAFeatsWrapper\(\)](#), [makeFeatSelWrapper\(\)](#), [makeFilterWrapper\(\)](#), [makeImputeWrapper\(\)](#), [makeMulticlassWrapper\(\)](#), [makeMultilabelBinaryRelevanceWrapper\(\)](#), [makeMultilabelClassifierChainsWrapper\(\)](#), [makeMultilabelDBRWrapper\(\)](#), [makeMultilabelNestedStackingWrapper\(\)](#), [makeMultilabelStackingWrapper\(\)](#), [makeOverBaggingWrapper\(\)](#), [makePreprocWrapperCaret\(\)](#), [makePreprocWrapper\(\)](#), [makeRemoveConstantFeaturesWrapper\(\)](#), [makeSMOTEWrapper\(\)](#), [makeTuneWrapper\(\)](#), [makeUndersampleWrapper\(\)](#), [makeWeightedClassesWrapper\(\)](#)

makeCostSensTask *Create a cost-sensitive classification task.*

Description

Create a cost-sensitive classification task.

Usage

```
makeCostSensTask(  
  id = deparse(substitute(data)),  
  data,  
  costs,  
  blocking = NULL,  
  coordinates = NULL,  
  fixup.data = "warn",  
  check.data = TRUE  
)
```

Arguments

id	(character(1)) Id string for object. Default is the name of the R variable passed to data.
data	(data.frame) A data frame containing the features and target variable(s).
costs	(data.frame) A numeric matrix or data frame containing the costs of misclassification. We assume the general case of observation specific costs. This means we have n rows, corresponding to the observations, in the same order as data. The columns correspond to classes and their names are the class labels (if unnamed we use y1 to yk as labels). Each entry (i,j) of the matrix specifies the cost of predicting class j for observation i.
blocking	(factor) An optional factor of the same length as the number of observations. Observations with the same blocking level “belong together”. Specifically, they are either put all in the training or the test set during a resampling iteration. Default is NULL which means no blocking.
coordinates	(data.frame) Coordinates of a spatial data set that will be used for spatial partitioning of the data in a spatial cross-validation resampling setting. Coordinates have to be numeric values. Provided data.frame needs to have the same number of rows as data and consist of at least two dimensions.
fixup.data	(character(1)) Should some basic cleaning up of data be performed? Currently this means removing empty factor levels for the columns. Possible choices are: “no” =

Don't do it. "warn" = Do it but warn about it. "quiet" = Do it but keep silent. Default is "warn".

check.data (logical(1))
Should sanity of data be checked initially at task creation? You should have good reasons to turn this off (one might be speed). Default is TRUE.

See Also

[Task](#) [ClassifTask](#) [ClusterTask](#) [MultilabelTask](#) [RegrTask](#) [SurvTask](#)

Other costsens: [makeCostSensClassifWrapper\(\)](#), [makeCostSensRegrWrapper\(\)](#), [makeCostSensWeightedPairsWrapper\(\)](#)

makeCostSensWeightedPairsWrapper

Wraps a classifier for cost-sensitive learning to produce a weighted pairs model.

Description

Creates a wrapper, which can be used like any other learner object. Models can easily be accessed via [getLearnerModel](#).

For each pair of labels, we fit a binary classifier. For each observation we define the label to be the element of the pair with minimal costs. During fitting, we also weight the observation with the absolute difference in costs. Prediction is performed by simple voting.

This approach is sometimes called cost-sensitive one-vs-one (CS-OVO), because it is obviously very similar to the one-vs-one approach where one reduces a normal multi-class problem to multiple binary ones and aggregates by voting.

Usage

```
makeCostSensWeightedPairsWrapper(learner)
```

Arguments

learner ([Learner](#) | character(1))
The classification learner. If you pass a string the learner will be created via [makeLearner](#).

Value

([Learner](#)).

References

Lin, HT.: Reduction from Cost-sensitive Multiclass Classification to One-versus-one Binary Classification. In: Proceedings of the Sixth Asian Conference on Machine Learning. JMLR Workshop and Conference Proceedings, vol 39, pp. 371-386. JMLR W&CP (2014). <https://www.jmlr.org/proceedings/papers/v39/lin14.pdf>

See Also

Other costsens: [makeCostSensClassifWrapper\(\)](#), [makeCostSensRegrWrapper\(\)](#), [makeCostSensTask\(\)](#)

makeCustomResampledMeasure

Construct your own resampled performance measure.

Description

Construct your own performance measure, used after resampling. Note that individual training / test set performance values will be set to NA, you only calculate an aggregated value. If you can define a function that makes sense for every single training / test set, implement your own [Measure](#).

Usage

```
makeCustomResampledMeasure(
  measure.id,
  aggregation.id,
  minimize = TRUE,
  properties = character(0L),
  fun,
  extra.args = list(),
  best = NULL,
  worst = NULL,
  measure.name = measure.id,
  aggregation.name = aggregation.id,
  note = ""
)
```

Arguments

measure.id	(character(1)) Short name of measure.
aggregation.id	(character(1)) Short name of aggregation.
minimize	(logical(1)) Should the measure be minimized? Default is TRUE.
properties	(character) Set of measure properties. For a list of values see Measure . Default is character(0).
fun	(function(task, group, pred, extra.args)) Calculates performance value from ResamplePrediction object. For rare cases you can also use the task, the grouping or the extra arguments extra.args. - task (Task) The task. - group (factor) Grouping of resampling iterations. This encodes whether specific iterations 'belong together' (e.g. repeated CV). - pred (Prediction)

	Prediction object. - extra.args (list) See below.
extra.args	(list) List of extra arguments which will always be passed to fun. Default is empty list.
best	(numeric(1)) Best obtainable value for measure. Default is -Inf or Inf, depending on minimize.
worst	(numeric(1)) Worst obtainable value for measure. Default is Inf or -Inf, depending on minimize.
measure.name	(character(1)) Long name of measure. Default is measure.id.
aggregation.name	(character(1)) Long name of the aggregation. Default is aggregation.id.
note	(character) Description and additional notes for the measure. Default is "".

Value

[Measure](#).

See Also

Other performance: [ConfusionMatrix](#), [calculateConfusionMatrix\(\)](#), [calculateROCMeasures\(\)](#), [estimateRelativeOverfitting\(\)](#), [makeCostMeasure\(\)](#), [makeMeasure\(\)](#), [measures](#), [performance\(\)](#), [setAggregation\(\)](#), [setMeasurePars\(\)](#)

makeDownsampleWrapper *Fuse learner with simple downsampling (subsampling).*

Description

Creates a learner object, which can be used like any other learner object. It will only be trained on a subset of the original data to save computational time.

Usage

```
makeDownsampleWrapper(learner, dw.perc = 1, dw.stratify = FALSE)
```

Arguments

learner	(Learner character(1)) The learner. If you pass a string the learner will be created via makeLearner .
dw.perc	(numeric(1)) See downsample . Default is 1.
dw.stratify	(logical(1)) See downsample . Default is FALSE.

Value

[Learner](#).

See Also

Other downsample: [downsample\(\)](#)

Other wrapper: [makeBaggingWrapper\(\)](#), [makeClassificationViaRegressionWrapper\(\)](#), [makeConstantClassWrapper\(\)](#), [makeCostSensClassifWrapper\(\)](#), [makeCostSensRegrWrapper\(\)](#), [makeDummyFeaturesWrapper\(\)](#), [makeExtractFDAFeatsWrapper\(\)](#), [makeFeatSelWrapper\(\)](#), [makeFilterWrapper\(\)](#), [makeImputeWrapper\(\)](#), [makeMulticlassWrapper\(\)](#), [makeMultilabelBinaryRelevanceWrapper\(\)](#), [makeMultilabelClassifierChainsWrapper\(\)](#), [makeMultilabelDBRWrapper\(\)](#), [makeMultilabelNestedStackingWrapper\(\)](#), [makeMultilabelStackingWrapper\(\)](#), [makeOverBaggingWrapper\(\)](#), [makePreprocWrapperCaret\(\)](#), [makePreprocWrapper\(\)](#), [makeRemoveConstantFeaturesWrapper\(\)](#), [makeSMOTEWrapper\(\)](#), [makeTuneWrapper\(\)](#), [makeUndersampleWrapper\(\)](#), [makeWeightedClassesWrapper\(\)](#)

makeDummyFeaturesWrapper

Fuse learner with dummy feature creator.

Description

Fuses a base learner with the dummy feature creator (see [createDummyFeatures](#)). Returns a learner which can be used like any other learner.

Usage

```
makeDummyFeaturesWrapper(learner, method = "1-of-n", cols = NULL)
```

Arguments

learner	(Learner character(1)) The learner. If you pass a string the learner will be created via makeLearner .
method	(character(1)) Available are: "1-of-n" : For n factor levels there will be n dummy variables. "reference" : There will be n-1 dummy variables leaving out the first factor level of each variable. Default is "1-of-n".
cols	(character) Columns to create dummy features for. Default is to use all columns.

Value

[Learner](#).

See Also

Other wrapper: `makeBaggingWrapper()`, `makeClassificationViaRegressionWrapper()`, `makeConstantClassWrapper()`, `makeCostSensClassifWrapper()`, `makeCostSensRegrWrapper()`, `makeDownsampleWrapper()`, `makeExtractFDAFeatsWrapper()`, `makeFeatSelWrapper()`, `makeFilterWrapper()`, `makeImputeWrapper()`, `makeMulticlassWrapper()`, `makeMultilabelBinaryRelevanceWrapper()`, `makeMultilabelClassifierChainsWrapper()`, `makeMultilabelDBRWrapper()`, `makeMultilabelNestedStackingWrapper()`, `makeMultilabelStackingWrapper()`, `makeOverBaggingWrapper()`, `makePreprocWrapperCaret()`, `makePreprocWrapper()`, `makeRemoveConstantFeaturesWrapper()`, `makeSMOTEWrapper()`, `makeTuneWrapper()`, `makeUndersampleWrapper()`, `makeWeightedClassesWrapper()`

makeExtractFDAFeatMethod

Constructor for FDA feature extraction methods.

Description

This can be used to implement custom FDA feature extraction. Takes a learn and a reextract function along with some optional parameters to those as argument.

Usage

```
makeExtractFDAFeatMethod(learn, reextract, args = list(), par.set = NULL)
```

Arguments

learn	(function(data, target, col, ...)) Function to learn and extract information on functional column col. Arguments are: <ul style="list-style-type: none"> • data <code>data.frame</code> Data.frame containing matrices with one row per observation of a single functional or time series and one column per measurement at each time point. All entries need to be numeric. • target <code>character(1)</code> Name of the target variable. Default: “NULL”. The variable is only set to be consistent with the API. • col <code>character(1) numeric(1)</code> column names or indices, the extraction should be performed on. The function has to return a named list of values.
reextract	(function(data, target, col, ...)) Function used for reextracting data in predict phase. Can be equal to learn.
args	(list) Named list of arguments to pass to learn via ...
par.set	(ParamSet) Paramset added to the learner if used in conjunction with a <code>makeExtractFDAFeatsWrapper</code> . Can be NULL.

See Also

Other fda: [extractFDAFeatures\(\)](#), [makeExtractFDAFeatsWrapper\(\)](#)

makeExtractFDAFeatsWrapper

Fuse learner with an extractFDAFeatures method.

Description

Fuses a base learner with an extractFDAFeatures method. Creates a learner object, which can be used like any other learner object. Internally uses [extractFDAFeatures](#) before training the learner and [reextractFDAFeatures](#) before predicting.

Usage

```
makeExtractFDAFeatsWrapper(learner, feat.methods = list())
```

Arguments

learner	(Learner character(1)) The learner. If you pass a string the learner will be created via makeLearner .
feat.methods	(named list) List of functional features along with the desired methods for each functional feature. “all” applies the extractFDAFeatures method to each functional feature. Names of feat.methods must match column names of functional features. Available feature extraction methods are available under family fda_featextractor. Specifying a functional feature multiple times with different extraction methods allows for the extraction of different features from the same functional. Default is list() which does nothing.

Value

[Learner](#).

See Also

Other fda: [extractFDAFeatures\(\)](#), [makeExtractFDAFeatMethod\(\)](#)

Other wrapper: [makeBaggingWrapper\(\)](#), [makeClassificationViaRegressionWrapper\(\)](#), [makeConstantClassWrapper\(\)](#), [makeCostSensClassifWrapper\(\)](#), [makeCostSensRegrWrapper\(\)](#), [makeDownsampleWrapper\(\)](#), [makeDummyFeaturesWrapper\(\)](#), [makeFeatSelWrapper\(\)](#), [makeFilterWrapper\(\)](#), [makeImputeWrapper\(\)](#), [makeMulticlassWrapper\(\)](#), [makeMultilabelBinaryRelevanceWrapper\(\)](#), [makeMultilabelClassifierChainsWrapper\(\)](#), [makeMultilabelDBRWrapper\(\)](#), [makeMultilabelNestedStackingWrapper\(\)](#), [makeMultilabelStackingWrapper\(\)](#), [makeOverBaggingWrapper\(\)](#), [makePreprocWrapperCaret\(\)](#), [makePreprocWrapper\(\)](#), [makeRemoveConstantFeaturesWrapper\(\)](#), [makeSMOTEWrapper\(\)](#), [makeTuneWrapper\(\)](#), [makeUndersampleWrapper\(\)](#), [makeWeightedClassesWrapper\(\)](#)

makeFeatSelWrapper *Fuse learner with feature selection.*

Description

Fuses a base learner with a search strategy to select variables. Creates a learner object, which can be used like any other learner object, but which internally uses [selectFeatures](#). If the train function is called on it, the search strategy and resampling are invoked to select an optimal set of variables. Finally, a model is fitted on the complete training data with these variables and returned. See [selectFeatures](#) for more details.

After training, the optimal features (and other related information) can be retrieved with [getFeatSelResult](#).

Usage

```
makeFeatSelWrapper(
  learner,
  resampling,
  measures,
  bit.names,
  bits.to.features,
  control,
  show.info = getMlrOption("show.info")
)
```

Arguments

learner	(Learner character(1)) The learner. If you pass a string the learner will be created via makeLearner .
resampling	(ResampleInstance ResampleDesc) Resampling strategy for feature selection. If you pass a description, it is instantiated once at the beginning by default, so all points are evaluated on the same training/test sets. If you want to change that behavior, look at FeatSelControl .
measures	(list of Measure Measure) Performance measures to evaluate. The first measure, aggregated by the first aggregation function is optimized, others are simply evaluated. Default is the default measure for the task, see here getDefaultMeasure .
bit.names	character Names of bits encoding the solutions. Also defines the total number of bits in the encoding. Per default these are the feature names of the task. Has to be used together with <code>bits.to.features</code> .
bits.to.features	(function(x, task)) Function which transforms an integer-0-1 vector into a character vector of selected features. Per default a value of 1 in the <i>i</i> th bit selects the <i>i</i> th feature to be

	in the candidate solution. The vector <code>x</code> will correspond to the <code>bit.names</code> and has to be of the same length.
<code>control</code>	[see FeatSelControl] Control object for search method. Also selects the optimization algorithm for feature selection.
<code>show.info</code>	(logical(1)) Print verbose output on console? Default is set via configureMlr .

Value

[Learner](#).

See Also

Other featsel: [FeatSelControl](#), [analyzeFeatSelResult\(\)](#), [getFeatSelResult\(\)](#), [selectFeatures\(\)](#)

Other wrapper: [makeBaggingWrapper\(\)](#), [makeClassificationViaRegressionWrapper\(\)](#), [makeConstantClassWrapper\(\)](#), [makeCostSensClassifWrapper\(\)](#), [makeCostSensRegrWrapper\(\)](#), [makeDownsampleWrapper\(\)](#), [makeDummyFeaturesWrapper\(\)](#), [makeExtractFDAFeatsWrapper\(\)](#), [makeFilterWrapper\(\)](#), [makeImputeWrapper\(\)](#), [makeMulticlassWrapper\(\)](#), [makeMultilabelBinaryRelevanceWrapper\(\)](#), [makeMultilabelClassifierChainsWrapper\(\)](#), [makeMultilabelDBRWrapper\(\)](#), [makeMultilabelNestedStackingWrapper\(\)](#), [makeMultilabelStackingWrapper\(\)](#), [makeOverBaggingWrapper\(\)](#), [makePreprocWrapperCaret\(\)](#), [makePreprocWrapper\(\)](#), [makeRemoveConstantFeaturesWrapper\(\)](#), [makeSMOTEWrapper\(\)](#), [makeTuneWrapper\(\)](#), [makeUndersampleWrapper\(\)](#), [makeWeightedClassesWrapper\(\)](#)

Examples

```
# nested resampling with feature selection (with a nonsense algorithm for selection)
outer = makeResampleDesc("CV", iters = 2L)
inner = makeResampleDesc("Holdout")
ctrl = makeFeatSelControlRandom(maxit = 1)
lrn = makeFeatSelWrapper("classif.ksvm", resampling = inner, control = ctrl)
# we also extract the selected features for all iteration here
r = resample(lrn, iris.task, outer, extract = getFeatSelResult)
```

makeFilter

Create a feature filter.

Description

Creates and registers custom feature filters. Implemented filters can be listed with [listFilterMethods](#). Additional documentation for the fun parameter specific to each filter can be found in the description.

Usage

```
makeFilter(name, desc, pkg, supported.tasks, supported.features, fun)
```

Arguments

name	(character(1)) Identifier for the filter.
desc	(character(1)) Short description of the filter.
pkg	(character(1)) Source package where the filter is implemented.
supported.tasks	(character) Task types supported.
supported.features	(character) Feature types supported.
fun	(function(task, nselect, ...)) Function which takes a task and returns a named numeric vector of scores, one score for each feature of task. Higher scores mean higher importance of the feature. At least nselect features must be calculated, the remaining may be set to NA or omitted, and thus will not be selected. the original order will be restored if necessary.

Value

Object of class “Filter”.

References

Kira, Kenji and Rendell, Larry (1992). The Feature Selection Problem: Traditional Methods and a New Algorithm. AAAI-92 Proceedings.

Kononenko, Igor et al. Overcoming the myopia of inductive learning algorithms with RELIEFF (1997), Applied Intelligence, 7(1), p39-55.

See Also

Other filter: [filterFeatures\(\)](#), [generateFilterValuesData\(\)](#), [getFilteredFeatures\(\)](#), [listFilterEnsembleMethods\(\)](#), [listFilterMethods\(\)](#), [makeFilterEnsemble\(\)](#), [makeFilterWrapper\(\)](#), [plotFilterValues\(\)](#)

makeFilterEnsemble *Create an ensemble feature filter.*

Description

Creates and registers custom ensemble feature filters. Implemented ensemble filters can be listed with [listFilterEnsembleMethods](#). Additional documentation for the fun parameter specific to each filter can be found in the description.

Usage

```
makeFilterEnsemble(name, base.methods, desc, fun)
```

Arguments

name	(character(1)) Identifier for the filter.
base.methods	the base filter methods which the ensemble method will use.
desc	(character(1)) Short description of the filter.
fun	(function(task, nselect, ...)) Function which takes a task and returns a named numeric vector of scores, one score for each feature of task. Higher scores mean higher importance of the feature. At least nselect features must be calculated, the remaining may be set to NA or omitted, and thus will not be selected. the original order will be restored if necessary.

Value

Object of class “FilterEnsemble”.

See Also

Other filter: [filterFeatures\(\)](#), [generateFilterValuesData\(\)](#), [getFilteredFeatures\(\)](#), [listFilterEnsembleMethods\(\)](#), [listFilterMethods\(\)](#), [makeFilterWrapper\(\)](#), [makeFilter\(\)](#), [plotFilterValues\(\)](#)

makeFilterWrapper *Fuse learner with a feature filter method.*

Description

Fuses a base learner with a filter method. Creates a learner object, which can be used like any other learner object. Internally uses [filterFeatures](#) before every model fit.

Usage

```
makeFilterWrapper(
  learner,
  fw.method = "randomForestSRC_importance",
  fw.base.methods = NULL,
  fw.perc = NULL,
  fw.abs = NULL,
  fw.threshold = NULL,
  fw.fun = NULL,
  fw.fun.args = NULL,
  fw.mandatory.feats = NULL,
```

```

    cache = FALSE,
    ...
  )

```

Arguments

learner	(Learner character(1)) The learner. If you pass a string the learner will be created via makeLearner .
fw.method	(character(1)) Filter method. See listFilterMethods . Default is “randomForestSRC_importance”.
fw.base.methods	(character(1)) Simple Filter methods for ensemble filters. See listFilterMethods . Can only be used in combination with ensemble filters. See listFilterEnsembleMethods .
fw.perc	(numeric(1)) If set, select fw.perc*100 top scoring features. Mutually exclusive with arguments fw.abs, fw.threshold and ‘fw.fun’.
fw.abs	(numeric(1)) If set, select fw.abs top scoring features. Mutually exclusive with arguments fw.perc, fw.threshold and fw.fun.
fw.threshold	(numeric(1)) If set, select features whose score exceeds fw.threshold. Mutually exclusive with arguments fw.perc, fw.abs and fw.fun.
fw.fun	(function) If set, select features via a custom thresholding function, which must return the number of top scoring features to select. Mutually exclusive with arguments fw.perc, fw.abs and fw.threshold.
fw.fun.args	(any) Arguments passed to the custom thresholding function
fw.mandatory.feats	(character) Mandatory features which are always included regardless of their scores
cache	(character(1) logical) Whether to use caching during filter value creation. See details.
...	(any) Additional parameters passed down to the filter. If you are using more than one filter method, you need to pass the arguments in a named list via more.args. For example more.args = list("FSelectorRcpp_information.gain" = list(equal = TRUE)).

Details

If ensemble = TRUE, ensemble feature selection using all methods specified in fw.method is performed. At least two methods need to be selected.

After training, the selected features can be retrieved with [getFilteredFeatures](#).

Note that observation weights do not influence the filtering and are simply passed down to the next learner.

Value

[Learner](#).

Caching

If `cache = TRUE`, the default mlr cache directory is used to cache filter values. The directory is operating system dependent and can be checked with `getCacheDir()`. Alternatively a custom directory can be passed to store the cache. The cache can be cleared with `deleteCacheDir()`. Caching is disabled by default. Care should be taken when operating on large clusters due to possible write conflicts to disk if multiple workers try to write the same cache at the same time.

See Also

Other filter: [filterFeatures\(\)](#), [generateFilterValuesData\(\)](#), [getFilteredFeatures\(\)](#), [listFilterEnsembleMethods\(\)](#), [listFilterMethods\(\)](#), [makeFilterEnsemble\(\)](#), [makeFilter\(\)](#), [plotFilterValues\(\)](#)

Other wrapper: [makeBaggingWrapper\(\)](#), [makeClassificationViaRegressionWrapper\(\)](#), [makeConstantClassWrapper\(\)](#), [makeCostSensClassifWrapper\(\)](#), [makeCostSensRegrWrapper\(\)](#), [makeDownsampleWrapper\(\)](#), [makeDummyFeaturesWrapper\(\)](#), [makeExtractFDAFeatsWrapper\(\)](#), [makeFeatSelWrapper\(\)](#), [makeImputeWrapper\(\)](#), [makeMulticlassWrapper\(\)](#), [makeMultilabelBinaryRelevanceWrapper\(\)](#), [makeMultilabelClassifierChainsWrapper\(\)](#), [makeMultilabelDBRWrapper\(\)](#), [makeMultilabelNestedStackingWrapper\(\)](#), [makeMultilabelStackingWrapper\(\)](#), [makeOverBaggingWrapper\(\)](#), [makePreprocWrapperCaret\(\)](#), [makePreprocWrapper\(\)](#), [makeRemoveConstantFeaturesWrapper\(\)](#), [makeSMOTEWrapper\(\)](#), [makeTuneWrapper\(\)](#), [makeUndersampleWrapper\(\)](#), [makeWeightedClassesWrapper\(\)](#)

Examples

```
task = makeClassifTask(data = iris, target = "Species")
lrn = makeLearner("classif.lda")
inner = makeResampleDesc("Holdout")
outer = makeResampleDesc("CV", iters = 2)
lrn = makeFilterWrapper(lrn, fw.perc = 0.5)
mod = train(lrn, task)
print(getFilteredFeatures(mod))
# now nested resampling, where we extract the features that the filter method selected
r = resample(lrn, task, outer, extract = function(model) {
  getFilteredFeatures(model)
})
print(r$extract)

# usage of an ensemble filter
lrn = makeLearner("classif.lda")
lrn = makeFilterWrapper(lrn, fw.method = "E-Borda",
  fw.base.methods = c("FSelectorRcpp_gain.ratio", "FSelectorRcpp_information.gain"),
  fw.perc = 0.5)
r = resample(lrn, task, outer, extract = function(model) {
  getFilteredFeatures(model)
})
print(r$extract)

# usage of a custom thresholding function
```

```

biggest_gap = function(values, diff) {
  gap_size = 0
  gap_location = 0

  for (i in (diff + 1):length(values)) {
    gap = values[[i - diff]] - values[[i]]
    if (gap > gap_size) {
      gap_size = gap
      gap_location = i - 1
    }
  }
  return(gap_location)
}

lrn = makeLearner("classif.lda")
lrn = makeFilterWrapper(lrn, fw.method = "randomForestSRC_importance",
  fw.fun = biggest_gap, fw.fun.args = list("diff" = 1))
r = resample(lrn, task, outer, extract = function(model) {
  getFilteredFeatures(model)
})
print(r$extract)

```

makeFixedHoldoutInstance

Generate a fixed holdout instance for resampling.

Description

Generate a fixed holdout instance for resampling.

Usage

```
makeFixedHoldoutInstance(train.inds, test.inds, size)
```

Arguments

train.inds	(integer) Indices for training set.
test.inds	(integer) Indices for test set.
size	(integer(1)) Size of the data set to resample. The function needs to know the largest possible index of the whole data set.

Value

(ResampleInstance).

makeFunctionalData	<i>Create a data.frame containing functional features from a normal data.frame.</i>
--------------------	---

Description

To work with functional features, those features need to be stored as a `matrix` column in the `data.frame`, so `mlr` can automatically recognize them as functional features. This function allows for an easy conversion from a `data.frame` with numeric columns to the required format. If the data already contains matrix columns, they are left as-is if not specified otherwise in `fd.features`. See Examples for the structure of the generated output.

Usage

```
makeFunctionalData(data, fd.features = NULL, exclude.cols = NULL)
```

Arguments

<code>data</code>	(data.frame) A <code>data.frame</code> that contains the functional features as numeric columns.
<code>fd.features</code>	(list) Named list containing integer column indices or character column names. Each element defines a functional feature, in the given order of the indices or column names. The name of the list element defines the name of the functional feature. All selected columns have to correspond to numeric <code>data.frame</code> entries. The default is <code>NULL</code> , which means all numeric features are considered to be a single functional “fd1”.
<code>exclude.cols</code>	(character integer) Column names or indices to exclude from conversion to functionals, even if they are included in <code>fd.features</code> . Default is not to exclude anything.

Value

([data.frame](#)).

Examples

```
# data.frame where columns 1:6 and 8:10 belong to a functional feature
d1 = data.frame(matrix(rnorm(100), nrow = 10), "target" = seq_len(10))
# Transform to functional data
d2 = makeFunctionalData(d1, fd.features = list("fd1" = 1:6, "fd2" = 8:10))
# Create a regression task
makeRegrTask(data = d2, target = "target")
```

makeImputeMethod *Create a custom imputation method.*

Description

This is a constructor to create your own imputation methods.

Usage

```
makeImputeMethod(learn, impute, args = list())
```

Arguments

learn	(function(data, target, col, ...)) Function to learn and extract information on column col out of data frame data. Argument target specifies the target column of the learning task. The function has to return a named list of values.
impute	(function(data, target, col, ...)) Function to impute missing values in col using information returned by learn on the same column. All list elements of the return values of learn are passed to this function into
args	(list) Named list of arguments to pass to learn via

See Also

Other impute: [imputations](#), [impute\(\)](#), [makeImputeWrapper\(\)](#), [reimpute\(\)](#)

makeImputeWrapper *Fuse learner with an imputation method.*

Description

Fuses a base learner with an imputation method. Creates a learner object, which can be used like any other learner object. Internally uses [impute](#) before training the learner and [reimpute](#) before predicting.

Usage

```
makeImputeWrapper(
  learner,
  classes = list(),
  cols = list(),
  dummy.classes = character(0L),
  dummy.cols = character(0L),
```

```

dummy.type = "factor",
force.dummies = FALSE,
impute.new.levels = TRUE,
recode.factor.levels = TRUE
)

```

Arguments

learner	(Learner character(1)) The learner. If you pass a string the learner will be created via makeLearner .
classes	(named list) Named list containing imputation techniques for classes of columns. E.g. <code>list(numeric = imputeMedian())</code> .
cols	(named list) Named list containing names of imputation methods to impute missing values in the data column referenced by the list element's name. Overrides imputation set via classes.
dummy.classes	(character) Classes of columns to create dummy columns for. Default is <code>character(0)</code> .
dummy.cols	(character) Column names to create dummy columns (containing binary missing indicator) for. Default is <code>character(0)</code> .
dummy.type	(character (1)) How dummy columns are encoded. Either as 0/1 with type "numeric" or as "factor". Default is "factor".
force.dummies	(logical (1)) Force dummy creation even if the respective data column does not contain any NAs. Note that (a) most learners will complain about constant columns created this way but (b) your feature set might be stochastic if you turn this off. Default is FALSE.
impute.new.levels	(logical (1)) If new, unencountered factor level occur during reimputation, should these be handled as NAs and then be imputed the same way? Default is TRUE.
recode.factor.levels	(logical (1)) Recode factor levels after reimputation, so they match the respective element of <code>lvls</code> (in the description object) and therefore match the levels of the feature factor in the training data after imputation?. Default is TRUE.

Value

[Learner](#).

See Also

Other impute: [imputations](#), [impute\(\)](#), [makeImputeMethod\(\)](#), [reimpute\(\)](#)

Other wrapper: `makeBaggingWrapper()`, `makeClassificationViaRegressionWrapper()`, `makeConstantClassWrapper()`, `makeCostSensClassifWrapper()`, `makeCostSensRegrWrapper()`, `makeDownsampleWrapper()`, `makeDummyFeaturesWrapper()`, `makeExtractFDAFeatsWrapper()`, `makeFeatSelWrapper()`, `makeFilterWrapper()`, `makeMulticlassWrapper()`, `makeMultilabelBinaryRelevanceWrapper()`, `makeMultilabelClassifierChainsWrapper()`, `makeMultilabelDBRWrapper()`, `makeMultilabelNestedStackingWrapper()`, `makeMultilabelStackingWrapper()`, `makeOverBaggingWrapper()`, `makePreprocWrapperCaret()`, `makePreprocWrapper()`, `makeRemoveConstantFeaturesWrapper()`, `makeSMOTERWrapper()`, `makeTuneWrapper()`, `makeUndersampleWrapper()`, `makeWeightedClassesWrapper()`

makeLearner

Create learner object.

Description

For a classification learner the `predict.type` can be set to “prob” to predict probabilities and the maximum value selects the label. The threshold used to assign the label can later be changed using the `setThreshold` function.

To see all possible properties of a learner, go to: [LearnerProperties](#).

Usage

```
makeLearner(
  cl,
  id = cl,
  predict.type = "response",
  predict.threshold = NULL,
  fix.factors.prediction = FALSE,
  ...,
  par.vals = list(),
  config = list()
)
```

Arguments

<code>cl</code>	(character(1)) Class of learner. By convention, all classification learners start with “classif.” all regression learners with “regr.” all survival learners start with “surv.” all clustering learners with “cluster.” and all multilabel classification learners start with “multilabel.”. A list of all integrated learners is available on the learners help page.
<code>id</code>	(character(1)) Id string for object. Used to display object. Default is <code>cl</code> .
<code>predict.type</code>	(character(1)) Classification: “response” (= labels) or “prob” (= probabilities and labels by selecting the ones with maximal probability). Regression: “response” (= mean response) or “se” (= standard errors and mean response). Survival: “response”

(= some sort of orderable risk) or “prob” (= time dependent probabilities). Clustering: “response” (= cluster IDS) or “prob” (= fuzzy cluster membership probabilities), Multilabel: “response” (= logical matrix indicating the predicted class labels) or “prob” (= probabilities and corresponding logical matrix indicating class labels). Default is “response”.

`predict.threshold`

(numeric)

Threshold to produce class labels. Has to be a named vector, where names correspond to class labels. Only for binary classification it can be a single numerical threshold for the positive class. See [setThreshold](#) for details on how it is applied. Default is NULL which means 0.5 / an equal threshold for each class.

`fix.factors.prediction`

(logical(1))

In some cases, problems occur in underlying learners for factor features during prediction. If the new features have LESS factor levels than during training (a strict subset), the learner might produce an error like “type of predictors in new data do not match that of the training data”. In this case one can repair this problem by setting this option to TRUE. We will simply add the missing factor levels missing from the test feature (but present in training) to that feature. Default is FALSE.

...

(any)

Optional named (hyper)parameters. If you want to set specific hyperparameters for a learner during model creation, these should go here. You can get a list of available hyperparameters using `getParamSet(<learner>)`. Alternatively hyperparameters can be given using the `par.vals` argument but ... should be preferred!

`par.vals`

(list)

Optional list of named (hyper)parameters. The arguments in ... take precedence over values in this list. We strongly encourage you to use ... for passing hyperparameters.

`config`

(named list)

Named list of config option to overwrite global settings set via [configureMlr](#) for this specific learner.

Value

([Learner](#)).

`par.vals` vs. ...

The former aims at specifying default hyperparameter settings from `mlr` which differ from the actual defaults in the underlying learner. For example, `respect.unordered.factors` is set to `order` in `mlr` while the default in [ranger::ranger](#) depends on the argument `splitrule`. `getHyperPars(<learner>)` can be used to query hyperparameter defaults that differ from the underlying learner. This function also shows all hyperparameters set by the user during learner creation (if these differ from the learner defaults).

regr.randomForest

For this learner we added additional uncertainty estimation functionality (`predict.type = "se"`) for the `randomForest`, which is not provided by the underlying package.

Currently implemented methods are:

- If `se.method = "jackknife"` the standard error of a prediction is estimated by computing the jackknife-after-bootstrap, the mean-squared difference between the prediction made by only using trees which did not contain said observation and the ensemble prediction.
- If `se.method = "bootstrap"` the standard error of a prediction is estimated by bootstrapping the random forest, where the number of bootstrap replicates and the number of trees in the ensemble are controlled by `se.boot` and `se.ntree` respectively, and then taking the standard deviation of the bootstrap predictions. The "brute force" bootstrap is executed when `ntree = se.ntree`, the latter of which controls the number of trees in the individual random forests which are bootstrapped. The "noisy bootstrap" is executed when `se.ntree < ntree` which is less computationally expensive. A Monte-Carlo bias correction may make the latter option preferable in many cases. Defaults are `se.boot = 50` and `se.ntree = 100`.
- If `se.method = "sd"`, the default, the standard deviation of the predictions across trees is returned as the variance estimate. This can be computed quickly but is also a very naive estimator.

For both "jackknife" and "bootstrap", a Monte-Carlo bias correction is applied and, in the case that this results in a negative variance estimate, the values are truncated at 0.

Note that when using the "jackknife" procedure for `se` estimation, using a small number of trees can lead to training data observations that are never out-of-bag. The current implementation ignores these observations, but in the original definition, the resulting `se` estimation would be undefined.

Please note that all of the mentioned `se.method` variants do not affect the computation of the posterior mean "response" value. This is always the same as from the underlying `randomForest`.

regr.featureless

A very basic baseline method which is useful for model comparisons (if you don't beat this, you very likely have a problem). Does not consider any features of the task and only uses the target feature of the training data to make predictions. Using observation weights is currently not supported.

Methods "mean" and "median" always predict a constant value for each new observation which corresponds to the observed mean or median of the target feature in training data, respectively.

The default method is "mean" which corresponds to the ZeroR algorithm from WEKA, see <https://weka.wikispaces.com/ZeroR>.

classif.featureless

Method "majority" predicts always the majority class for each new observation. In the case of ties, one randomly sampled, constant class is predicted for all observations in the test set. This method is used as the default. It is very similar to the ZeroR classifier from WEKA (see <https://weka.wikispaces.com/ZeroR>). The only difference is that ZeroR always predicts the first class of the tied class values instead of sampling them randomly.

Method "sample-prior" always samples a random class for each individual test observation according to the prior probabilities observed in the training data.

If you opt to predict probabilities, the class probabilities always correspond to the prior probabilities observed in the training data.

See Also

Other learner: [LearnerProperties](#), [getClassWeightParam\(\)](#), [getHyperPars\(\)](#), [getLearnerId\(\)](#), [getLearnerNote\(\)](#), [getLearnerPackages\(\)](#), [getLearnerParVals\(\)](#), [getLearnerParamSet\(\)](#), [getLearnerPredictType\(\)](#), [getLearnerShortName\(\)](#), [getLearnerType\(\)](#), [getParamSet\(\)](#), [helpLearnerParam\(\)](#), [helpLearner\(\)](#), [makeLearners\(\)](#), [removeHyperPars\(\)](#), [setHyperPars\(\)](#), [setId\(\)](#), [setLearnerId\(\)](#), [setPredictThreshold\(\)](#), [setPredictType\(\)](#)

Examples

```
makeLearner("classif.rpart")
makeLearner("classif.lda", predict.type = "prob")
lrn = makeLearner("classif.lda", method = "t", nu = 10)
getHyperPars(lrn)
```

makeLearners	<i>Create multiple learners at once.</i>
--------------	--

Description

Small helper function that can save some typing when creating mutiple learner objects. Calls [makeLearner](#) multiple times internally.

Usage

```
makeLearners(cls, ids = NULL, type = NULL, ...)
```

Arguments

cls	(character) Classes of learners.
ids	(character) Id strings. Must be unique. Default is cls.
type	(character(1)) Shortcut to prepend type string to cls so one can set cls = "rpart". Default is NULL, i.e., this is not used.
...	(any) Optional named (hyper)parameters. If you want to set specific hyperparameters for a learner during model creation, these should go here. You can get a list of available hyperparameters using <code>getParamSet(<learner>)</code> . Alternatively hyperparameters can be given using the <code>par.vals</code> argument but ... should be preferred!

Value

(named list of [Learner](#)). Named by ids.

See Also

Other learner: [LearnerProperties](#), [getClassWeightParam\(\)](#), [getHyperPars\(\)](#), [getLearnerId\(\)](#), [getLearnerNote\(\)](#), [getLearnerPackages\(\)](#), [getLearnerParVals\(\)](#), [getLearnerParamSet\(\)](#), [getLearnerPredictType\(\)](#), [getLearnerShortName\(\)](#), [getLearnerType\(\)](#), [getParamSet\(\)](#), [helpLearnerParam\(\)](#), [helpLearner\(\)](#), [makeLearner\(\)](#), [removeHyperPars\(\)](#), [setHyperPars\(\)](#), [setId\(\)](#), [setLearnerId\(\)](#), [setPredictThreshold\(\)](#), [setPredictType\(\)](#)

Examples

```
makeLearners(c("rpart", "lda"), type = "classif", predict.type = "prob")
```

makeMeasure	<i>Construct performance measure.</i>
-------------	---------------------------------------

Description

A measure object encapsulates a function to evaluate the performance of a prediction. Information about already implemented measures can be obtained here: [measures](#).

A learner is trained on a training set `d1`, results in a model `m` and predicts another set `d2` (which may be a different one or the training set) resulting in the prediction. The performance measure can now be defined using all of the information of the original task, the fitted model and the prediction.

Usage

```
makeMeasure(
  id,
  minimize,
  properties = character(0L),
  fun,
  extra.args = list(),
  aggr = test.mean,
  best = NULL,
  worst = NULL,
  name = id,
  note = ""
)
```

Arguments

<code>id</code>	(<code>character(1)</code>) Name of measure.
<code>minimize</code>	(<code>logical(1)</code>) Should the measure be minimized? Default is TRUE.

properties	<p>(character)</p> <p>Set of measure properties. Some standard property names include: - <code>classif</code>: Is the measure applicable for classification? - <code>classif.multi</code>: Is the measure applicable for multi-class classification? - <code>multilabel</code>: Is the measure applicable for multilabel classification? - <code>regr</code>: Is the measure applicable for regression? - <code>surv</code>: Is the measure applicable for survival? - <code>cluster</code>: Is the measure applicable for cluster? - <code>costsens</code>: Is the measure applicable for cost-sensitive learning? - <code>req.pred</code>: Is prediction object required in calculation? Usually the case. - <code>req.truth</code>: Is truth column required in calculation? Usually the case. - <code>req.task</code>: Is task object required in calculation? Usually not the case - <code>req.model</code>: Is model object required in calculation? Usually not the case. - <code>req.feats</code>: Are feature values required in calculation? Usually not the case. - <code>req.prob</code>: Are predicted probabilities required in calculation? Usually not the case, example would be AUC.</p> <p>Default is <code>character(0)</code>.</p>
fun	<p>(<code>function(task, model, pred, feats, extra.args)</code>)</p> <p>Calculates the performance value. Usually you will only need the prediction object <code>pred</code>. - <code>task</code> (Task) The task. - <code>model</code> (WrappedModel) The fitted model. - <code>pred</code> (Prediction) Prediction object. - <code>feats</code> (data.frame) The features. - <code>extra.args</code> (list) See below.</p>
extra.args	<p>(list)</p> <p>List of extra arguments which will always be passed to <code>fun</code>. Can be changed after construction via <code>setMeasurePars()</code>. Default is empty list.</p>
aggr	<p>(Aggregation)</p> <p>Aggregation function, which is used to aggregate the values measured on test / training sets of the measure to a single value. Default is <code>test.mean</code>.</p>
best	<p>(<code>numeric(1)</code>)</p> <p>Best obtainable value for measure. Default is <code>-Inf</code> or <code>Inf</code>, depending on minimize.</p>
worst	<p>(<code>numeric(1)</code>)</p> <p>Worst obtainable value for measure. Default is <code>Inf</code> or <code>-Inf</code>, depending on minimize.</p>
name	<p>(character)</p> <p>Name of the measure. Default is <code>id</code>.</p>
note	<p>(character)</p> <p>Description and additional notes for the measure. Default is <code>""</code>.</p>

Value

[Measure](#).

See Also

Other performance: [ConfusionMatrix](#), [calculateConfusionMatrix\(\)](#), [calculateROCMeasures\(\)](#), [estimateRelativeOverfitting\(\)](#), [makeCostMeasure\(\)](#), [makeCustomResampledMeasure\(\)](#), [measures](#), [performance\(\)](#), [setAggregation\(\)](#), [setMeasurePars\(\)](#)

Examples

```
f = function(task, model, pred, extra.args) {
  sum((pred$data$response - pred$data$truth)^2)
}
makeMeasure(id = "my.sse", minimize = TRUE,
  properties = c("regr", "response"), fun = f)
```

makeModelMultiplexer *Create model multiplexer for model selection to tune over multiple possible models.*

Description

Combines multiple base learners by dispatching on the hyperparameter “selected.learner” to a specific model class. This allows to tune not only the model class (SVM, random forest, etc) but also their hyperparameters in one go. Combine this with [tuneParams](#) and [makeTuneControlIrace](#) for a very powerful approach, see example below.

The parameter set is the union of all (unique) base learners. In order to avoid name clashes all parameter names are prefixed with the base learner id, i.e. learnerId.parameterName.

The predict.type of the Multiplexer is inherited from the predict.type of the base learners.

The getter [getLearnerProperties](#) returns the properties of the selected base learner.

Usage

```
makeModelMultiplexer(base.learners)
```

Arguments

base.learners (list of [Learner](#))
List of Learners with unique IDs.

Value

([ModelMultiplexer](#)). A [Learner](#) specialized as ModelMultiplexer.

Note

Note that logging output during tuning is somewhat shortened to make it more readable. I.e., the artificial prefix before parameter names is suppressed.

See Also

Other multiplexer: [makeModelMultiplexerParamSet\(\)](#)

Other tune: [TuneControl](#), [getNestedTuneResultsOptPathDf\(\)](#), [getNestedTuneResultsX\(\)](#), [getResamplingIndices\(\)](#), [getTuneResult\(\)](#), [makeModelMultiplexerParamSet\(\)](#), [makeTuneControlCMAES\(\)](#), [makeTuneControlDesign\(\)](#), [makeTuneControlGenSA\(\)](#), [makeTuneControlGrid\(\)](#), [makeTuneControlIrace\(\)](#), [makeTuneControlMBO\(\)](#), [makeTuneControlRandom\(\)](#), [makeTuneWrapper\(\)](#), [tuneParams\(\)](#), [tuneThreshold\(\)](#)

Examples

```

set.seed(123)

library(BBmisc)
bls = list(
  makeLearner("classif.ksvm"),
  makeLearner("classif.randomForest")
)
lrn = makeModelMultiplexer(bls)
# simple way to construct param set for tuning
# parameter names are prefixed automatically and the 'requires'
# element is set, too, to make all parameters subordinate to 'selected.learner'
ps = makeModelMultiplexerParamSet(lrn,
  makeNumericParam("sigma", lower = -10, upper = 10, trafo = function(x) 2^x),
  makeIntegerParam("ntree", lower = 1L, upper = 500L)
)
print(ps)
rdesc = makeResampleDesc("CV", iters = 2L)
# to save some time we use random search. but you probably want something like this:
# ctrl = makeTuneControlIrace(maxExperiments = 500L)
ctrl = makeTuneControlRandom(maxit = 10L)
res = tuneParams(lrn, iris.task, rdesc, par.set = ps, control = ctrl)
print(res)

df = as.data.frame(res$opt.path)
print(head(df[, -ncol(df)]))

# more unique and reliable way to construct the param set
ps = makeModelMultiplexerParamSet(lrn,
  classif.ksvm = makeParamSet(
    makeNumericParam("sigma", lower = -10, upper = 10, trafo = function(x) 2^x)
  ),
  classif.randomForest = makeParamSet(
    makeIntegerParam("ntree", lower = 1L, upper = 500L)
  )
)

# this is how you would construct the param set manually, works too
ps = makeParamSet(
  makeDiscreteParam("selected.learner", values = extractSubList(bls, "id")),
  makeNumericParam("classif.ksvm.sigma", lower = -10, upper = 10, trafo = function(x) 2^x,
    requires = quote(selected.learner == "classif.ksvm")),
  makeIntegerParam("classif.randomForest.ntree", lower = 1L, upper = 500L,
    requires = quote(selected.learner == "classif.randomForst"))
)

# all three ps-objects are exactly the same internally.

```

makeModelMultiplexerParamSet

Creates a parameter set for model multiplexer tuning.

Description

Handy way to create the param set with less typing.

The following is done automatically:

- The selected.learner param is created
- Parameter names are prefixed.
- The requires field of each param is set. This makes all parameters subordinate to selected.learner

Usage

```
makeModelMultiplexerParamSet(multiplexer, ..., .check = TRUE)
```

Arguments

multiplexer	(ModelMultiplexer) The multiplexer learner.
...	(ParamHelpers::ParamSet ParamHelpers::Param) (a) First option: Named param sets. Names must correspond to base learners. You only need to enter the parameters you want to tune without reference to the selected.learner field in any way. (b) Second option. Just the params you would enter in the param sets. Even shorter to create. Only works when it can be uniquely identified to which learner each of your passed parameters belongs.
.check	(logical) Check that for each param in ... one param is found in the base learners. Default is TRUE

Value

[ParamSet](#).

See Also

Other multiplexer: [makeModelMultiplexer\(\)](#)

Other tune: [TuneControl](#), [getNestedTuneResultsOptPathDf\(\)](#), [getNestedTuneResultsX\(\)](#), [getResamplingIndices\(\)](#), [getTuneResult\(\)](#), [makeModelMultiplexer\(\)](#), [makeTuneControlCMAES\(\)](#), [makeTuneControlDesign\(\)](#), [makeTuneControlGenSA\(\)](#), [makeTuneControlGrid\(\)](#), [makeTuneControlIrace\(\)](#), [makeTuneControlMBO\(\)](#), [makeTuneControlRandom\(\)](#), [makeTuneWrapper\(\)](#), [tuneParams\(\)](#), [tuneThreshold\(\)](#)

Examples

```
# See makeModelMultiplexer
```

makeMulticlassWrapper *Fuse learner with multiclass method.*

Description

Fuses a base learner with a multi-class method. Creates a learner object, which can be used like any other learner object. This way learners which can only handle binary classification will be able to handle multi-class problems, too.

We use a multiclass-to-binary reduction principle, where multiple binary problems are created from the multiclass task. How these binary problems are generated is defined by an error-correcting-output-code (ECOC) code book. This also allows the simple and well-known one-vs-one and one-vs-rest approaches. Decoding is currently done via Hamming decoding, see e.g. here <https://jmlr.org/papers/volume11/escalera10a/escalera10a.pdf>.

Currently, the approach always operates on the discrete predicted labels of the binary base models (instead of their probabilities) and the created wrapper cannot predict posterior probabilities.

Usage

```
makeMulticlassWrapper(learner, mcw.method = "onevsrest")
```

Arguments

learner	(Learner character(1)) The learner. If you pass a string the learner will be created via makeLearner .
mcw.method	(character(1) function) “onevsone” or “onevsrest”. You can also pass a function, with signature function(task) and which returns a ECOC codematrix with entries +1,-1,0. Columns define new binary problems, rows correspond to classes (rows must be named). 0 means class is not included in binary problem. Default is “onevsrest”.

Value

[Learner](#).

See Also

Other wrapper: [makeBaggingWrapper\(\)](#), [makeClassificationViaRegressionWrapper\(\)](#), [makeConstantClassWrapper\(\)](#), [makeCostSensClassifWrapper\(\)](#), [makeCostSensRegrWrapper\(\)](#), [makeDownsampleWrapper\(\)](#), [makeDummyFeaturesWrapper\(\)](#), [makeExtractFDAFeatsWrapper\(\)](#), [makeFeatSelWrapper\(\)](#), [makeFilterWrapper\(\)](#), [makeImputeWrapper\(\)](#), [makeMultilabelBinaryRelevanceWrapper\(\)](#), [makeMultilabelClassifierChainsWrapper\(\)](#), [makeMultilabelDBRWrapper\(\)](#), [makeMultilabelNestedStackingWrapper\(\)](#), [makeMultilabelStackingWrapper\(\)](#), [makeOverBaggingWrapper\(\)](#), [makePreprocWrapperCaret\(\)](#), [makePreprocWrapper\(\)](#), [makeRemoveConstantFeaturesW](#), [makeSMOTEWrapper\(\)](#), [makeTuneWrapper\(\)](#), [makeUndersampleWrapper\(\)](#), [makeWeightedClassesWrapper\(\)](#)

makeMultilabelBinaryRelevanceWrapper

Use binary relevance method to create a multilabel learner.

Description

Every learner which is implemented in mlr and which supports binary classification can be converted to a wrapped binary relevance multilabel learner. The multilabel classification problem is converted into simple binary classifications for each label/target on which the binary learner is applied.

Models can easily be accessed via [getLearnerModel](#).

Note that it does not make sense to set a threshold in the used base learner when you predict probabilities. On the other hand, it can make a lot of sense, to call [setThreshold](#) on the MultilabelBinaryRelevanceWrapper for each label individually; Or to tune these thresholds with [tuneThreshold](#); especially when you face very unbalanced class distributions for each binary label.

Usage

```
makeMultilabelBinaryRelevanceWrapper(learner)
```

Arguments

learner ([Learner](#) | character(1))
The learner. If you pass a string the learner will be created via [makeLearner](#).

Value

[Learner](#).

References

Tsoumakas, G., & Katakis, I. (2006) *Multi-label classification: An overview*. Dept. of Informatics, Aristotle University of Thessaloniki, Greece.

See Also

Other wrapper: [makeBaggingWrapper\(\)](#), [makeClassificationViaRegressionWrapper\(\)](#), [makeConstantClassWrapper\(\)](#), [makeCostSensClassifWrapper\(\)](#), [makeCostSensRegrWrapper\(\)](#), [makeDownsampleWrapper\(\)](#), [makeDummyFeaturesWrapper\(\)](#), [makeExtractFDAFeatsWrapper\(\)](#), [makeFeatSelWrapper\(\)](#), [makeFilterWrapper\(\)](#), [makeImputeWrapper\(\)](#), [makeMulticlassWrapper\(\)](#), [makeMultilabelClassifierChainsWrapper\(\)](#), [makeMultilabelDBRWrapper\(\)](#), [makeMultilabelNestedStackingWrapper\(\)](#), [makeMultilabelStackingWrapper\(\)](#), [makeOverBaggingWrapper\(\)](#), [makePreprocWrapperCaret\(\)](#), [makePreprocWrapper\(\)](#), [makeRemoveConstantFeaturesWrapper\(\)](#), [makeSMOTEWrapper\(\)](#), [makeTuneWrapper\(\)](#), [makeUndersampleWrapper\(\)](#), [makeWeightedClassesWrapper\(\)](#)

Other multilabel: [getMultilabelBinaryPerformances\(\)](#), [makeMultilabelClassifierChainsWrapper\(\)](#), [makeMultilabelDBRWrapper\(\)](#), [makeMultilabelNestedStackingWrapper\(\)](#), [makeMultilabelStackingWrapper\(\)](#)

Examples

```
d = getTaskData(yeast.task)
# drop some labels so example runs faster
d = d[seq(1, nrow(d), by = 20), c(1:2, 15:17)]
task = makeMultilabelTask(data = d, target = c("label1", "label2"))
lrn = makeLearner("classif.rpart")
lrn = makeMultilabelBinaryRelevanceWrapper(lrn)
lrn = setPredictType(lrn, "prob")
# train, predict and evaluate
mod = train(lrn, task)
pred = predict(mod, task)
performance(pred, measure = list(multilabel.hamloss, multilabel.subset01, multilabel.f1))
# the next call basically has the same structure for any multilabel meta wrapper
getMultilabelBinaryPerformances(pred, measures = list(mmce, auc))
# above works also with predictions from resample!
```

```
makeMultilabelClassifierChainsWrapper
```

Use classifier chains method (CC) to create a multilabel learner.

Description

Every learner which is implemented in mlr and which supports binary classification can be converted to a wrapped classifier chains multilabel learner. CC trains a binary classifier for each label following a given order. In training phase, the feature space of each classifier is extended with true label information of all previous labels in the chain. During the prediction phase, when true labels are not available, they are replaced by predicted labels.

Models can easily be accessed via [getLearnerModel](#).

Usage

```
makeMultilabelClassifierChainsWrapper(learner, order = NULL)
```

Arguments

learner	(Learner character(1)) The learner. If you pass a string the learner will be created via makeLearner .
order	(character) Specifies the chain order using the names of the target labels. E.g. for m target labels, this must be a character vector of length m that contains a permutation of the target label names. Default is NULL which uses a random ordering of the target label names.

Value

[Learner](#).

References

Montanes, E. et al. (2013) *Dependent binary relevance models for multi-label classification* Artificial Intelligence Center, University of Oviedo at Gijon, Spain.

See Also

Other wrapper: [makeBaggingWrapper\(\)](#), [makeClassificationViaRegressionWrapper\(\)](#), [makeConstantClassWrapper\(\)](#), [makeCostSensClassifWrapper\(\)](#), [makeCostSensRegrWrapper\(\)](#), [makeDownsampleWrapper\(\)](#), [makeDummyFeaturesWrapper\(\)](#), [makeExtractFDAFeatsWrapper\(\)](#), [makeFeatSelWrapper\(\)](#), [makeFilterWrapper\(\)](#), [makeImputeWrapper\(\)](#), [makeMulticlassWrapper\(\)](#), [makeMultilabelBinaryRelevanceWrapper\(\)](#), [makeMultilabelDBRWrapper\(\)](#), [makeMultilabelNestedStackingWrapper\(\)](#), [makeMultilabelStackingWrapper\(\)](#), [makeOverBaggingWrapper\(\)](#), [makePreprocWrapperCaret\(\)](#), [makePreprocWrapper\(\)](#), [makeRemoveConstantFeaturesWrapper\(\)](#), [makeSMOTEWrapper\(\)](#), [makeTuneWrapper\(\)](#), [makeUndersampleWrapper\(\)](#), [makeWeightedClassesWrapper\(\)](#)

Other multilabel: [getMultilabelBinaryPerformances\(\)](#), [makeMultilabelBinaryRelevanceWrapper\(\)](#), [makeMultilabelDBRWrapper\(\)](#), [makeMultilabelNestedStackingWrapper\(\)](#), [makeMultilabelStackingWrapper\(\)](#)

Examples

```
d = getTaskData(yeast.task)
# drop some labels so example runs faster
d = d[seq(1, nrow(d), by = 20), c(1:2, 15:17)]
task = makeMultilabelTask(data = d, target = c("label1", "label2"))
lrn = makeLearner("classif.rpart")
lrn = makeMultilabelBinaryRelevanceWrapper(lrn)
lrn = setPredictType(lrn, "prob")
# train, predict and evaluate
mod = train(lrn, task)
pred = predict(mod, task)
performance(pred, measure = list(multilabel.hamloss, multilabel.subset01, multilabel.f1))
# the next call basically has the same structure for any multilabel meta wrapper
getMultilabelBinaryPerformances(pred, measures = list(mmce, auc))
# above works also with predictions from resample!
```

makeMultilabelDBRWrapper

Use dependent binary relevance method (DBR) to create a multilabel learner.

Description

Every learner which is implemented in mlr and which supports binary classification can be converted to a wrapped DBR multilabel learner. The multilabel classification problem is converted into simple binary classifications for each label/target on which the binary learner is applied. For each target, actual information of all binary labels (except the target variable) is used as additional features. During prediction these labels need are obtained by the binary relevance method using the same binary learner.

Models can easily be accessed via [getLearnerModel](#).

Usage

```
makeMultilabelDBRWrapper(learner)
```

Arguments

learner ([Learner](#) | character(1))
 The learner. If you pass a string the learner will be created via [makeLearner](#).

Value

[Learner](#).

References

Montanes, E. et al. (2013) *Dependent binary relevance models for multi-label classification* Artificial Intelligence Center, University of Oviedo at Gijon, Spain.

See Also

Other wrapper: [makeBaggingWrapper\(\)](#), [makeClassificationViaRegressionWrapper\(\)](#), [makeConstantClassWrapper\(\)](#), [makeCostSensClassifWrapper\(\)](#), [makeCostSensRegrWrapper\(\)](#), [makeDownsampleWrapper\(\)](#), [makeDummyFeaturesWrapper\(\)](#), [makeExtractFDAFeatsWrapper\(\)](#), [makeFeatSelWrapper\(\)](#), [makeFilterWrapper\(\)](#), [makeImputeWrapper\(\)](#), [makeMulticlassWrapper\(\)](#), [makeMultilabelBinaryRelevanceWrapper\(\)](#), [makeMultilabelClassifierChainsWrapper\(\)](#), [makeMultilabelNestedStackingWrapper\(\)](#), [makeMultilabelStackingWrapper\(\)](#), [makeOverBaggingWrapper\(\)](#), [makePreprocWrapperCaret\(\)](#), [makePreprocWrapper\(\)](#), [makeRemoveConstantFeaturesWrapper\(\)](#), [makeSMOTEWrapper\(\)](#), [makeTuneWrapper\(\)](#), [makeUndersampleWrapper\(\)](#), [makeWeightedClassesWrapper\(\)](#)

Other multilabel: [getMultilabelBinaryPerformances\(\)](#), [makeMultilabelBinaryRelevanceWrapper\(\)](#), [makeMultilabelClassifierChainsWrapper\(\)](#), [makeMultilabelNestedStackingWrapper\(\)](#), [makeMultilabelStackingWrapper\(\)](#)

Examples

```
d = getTaskData(yeast.task)
# drop some labels so example runs faster
d = d[seq(1, nrow(d), by = 20), c(1:2, 15:17)]
task = makeMultilabelTask(data = d, target = c("label1", "label2"))
lrn = makeLearner("classif.rpart")
lrn = makeMultilabelBinaryRelevanceWrapper(lrn)
lrn = setPredictType(lrn, "prob")
# train, predict and evaluate
mod = train(lrn, task)
pred = predict(mod, task)
performance(pred, measure = list(multilabel.hamloss, multilabel.subset01, multilabel.f1))
# the next call basically has the same structure for any multilabel meta wrapper
getMultilabelBinaryPerformances(pred, measures = list(mmce, auc))
# above works also with predictions from resample!
```

```
makeMultilabelNestedStackingWrapper
```

Use nested stacking method to create a multilabel learner.

Description

Every learner which is implemented in mlr and which supports binary classification can be converted to a wrapped nested stacking multilabel learner. Nested stacking trains a binary classifier for each label following a given order. In training phase, the feature space of each classifier is extended with predicted label information (by cross validation) of all previous labels in the chain. During the prediction phase, predicted labels are obtained by the classifiers, which have been learned on all training data.

Models can easily be accessed via [getLearnerModel](#).

Usage

```
makeMultilabelNestedStackingWrapper(learner, order = NULL, cv.folds = 2)
```

Arguments

learner	(Learner character(1)) The learner. If you pass a string the learner will be created via makeLearner .
order	(character) Specifies the chain order using the names of the target labels. E.g. for m target labels, this must be a character vector of length m that contains a permutation of the target label names. Default is NULL which uses a random ordering of the target label names.
cv.folds	(integer(1)) The number of folds for the inner cross validation method to predict labels for the augmented feature space. Default is 2.

Value

[Learner](#).

References

Montanes, E. et al. (2013), *Dependent binary relevance models for multi-label classification* Artificial Intelligence Center, University of Oviedo at Gijon, Spain.

See Also

Other wrapper: [makeBaggingWrapper\(\)](#), [makeClassificationViaRegressionWrapper\(\)](#), [makeConstantClassWrapper\(\)](#), [makeCostSensClassifWrapper\(\)](#), [makeCostSensRegrWrapper\(\)](#), [makeDownsampleWrapper\(\)](#), [makeDummyFeaturesWrapper\(\)](#), [makeExtractFDAFeatsWrapper\(\)](#), [makeFeatSelWrapper\(\)](#), [makeFilterWrapper\(\)](#), [makeImputeWrapper\(\)](#), [makeMulticlassWrapper\(\)](#), [makeMultilabelBinaryRelevanceWrapper\(\)](#),

```

makeMultilabelClassifierChainsWrapper(), makeMultilabelDBRWrapper(), makeMultilabelStackingWrapper(),
makeOverBaggingWrapper(), makePreprocWrapperCaret(), makePreprocWrapper(), makeRemoveConstantFeaturesW
makeSMOTEWrapper(), makeTuneWrapper(), makeUndersampleWrapper(), makeWeightedClassesWrapper()
Other multilabel: getMultilabelBinaryPerformances(), makeMultilabelBinaryRelevanceWrapper(),
makeMultilabelClassifierChainsWrapper(), makeMultilabelDBRWrapper(), makeMultilabelStackingWrapper()

```

Examples

```

d = getTaskData(yeast.task)
# drop some labels so example runs faster
d = d[seq(1, nrow(d), by = 20), c(1:2, 15:17)]
task = makeMultilabelTask(data = d, target = c("label1", "label2"))
lrn = makeLearner("classif.rpart")
lrn = makeMultilabelBinaryRelevanceWrapper(lrn)
lrn = setPredictType(lrn, "prob")
# train, predict and evaluate
mod = train(lrn, task)
pred = predict(mod, task)
performance(pred, measure = list(multilabel.hamloss, multilabel.subset01, multilabel.f1))
# the next call basically has the same structure for any multilabel meta wrapper
getMultilabelBinaryPerformances(pred, measures = list(mmce, auc))
# above works also with predictions from resample!

```

```
makeMultilabelStackingWrapper
```

Use stacking method (stacked generalization) to create a multilabel learner.

Description

Every learner which is implemented in mlr and which supports binary classification can be converted to a wrapped stacking multilabel learner. Stacking trains a binary classifier for each label using predicted label information of all labels (including the target label) as additional features (by cross validation). During prediction these labels need are obtained by the binary relevance method using the same binary learner.

Models can easily be accessed via [getLearnerModel](#).

Usage

```
makeMultilabelStackingWrapper(learner, cv.folds = 2)
```

Arguments

learner	(Learner character(1)) The learner. If you pass a string the learner will be created via makeLearner .
cv.folds	(integer(1)) The number of folds for the inner cross validation method to predict labels for the augmented feature space. Default is 2.

Value

[Learner](#).

References

Montanes, E. et al. (2013) *Dependent binary relevance models for multi-label classification* Artificial Intelligence Center, University of Oviedo at Gijon, Spain.

See Also

Other wrapper: [makeBaggingWrapper\(\)](#), [makeClassificationViaRegressionWrapper\(\)](#), [makeConstantClassWrapper\(\)](#), [makeCostSensClassifWrapper\(\)](#), [makeCostSensRegrWrapper\(\)](#), [makeDownsampleWrapper\(\)](#), [makeDummyFeaturesWrapper\(\)](#), [makeExtractFDAFeatsWrapper\(\)](#), [makeFeatSelWrapper\(\)](#), [makeFilterWrapper\(\)](#), [makeImputeWrapper\(\)](#), [makeMulticlassWrapper\(\)](#), [makeMultilabelBinaryRelevanceWrapper\(\)](#), [makeMultilabelClassifierChainsWrapper\(\)](#), [makeMultilabelDBRWrapper\(\)](#), [makeMultilabelNestedStackingWrapper\(\)](#), [makeOverBaggingWrapper\(\)](#), [makePreprocWrapperCaret\(\)](#), [makePreprocWrapper\(\)](#), [makeRemoveConstantFeaturesWrapper\(\)](#), [makeSMOTEWrapper\(\)](#), [makeTuneWrapper\(\)](#), [makeUndersampleWrapper\(\)](#), [makeWeightedClassesWrapper\(\)](#)

Other multilabel: [getMultilabelBinaryPerformances\(\)](#), [makeMultilabelBinaryRelevanceWrapper\(\)](#), [makeMultilabelClassifierChainsWrapper\(\)](#), [makeMultilabelDBRWrapper\(\)](#), [makeMultilabelNestedStackingWrapper\(\)](#)

Examples

```
d = getTaskData(yeast.task)
# drop some labels so example runs faster
d = d[seq(1, nrow(d), by = 20), c(1:2, 15:17)]
task = makeMultilabelTask(data = d, target = c("label1", "label2"))
lrn = makeLearner("classif.rpart")
lrn = makeMultilabelBinaryRelevanceWrapper(lrn)
lrn = setPredictType(lrn, "prob")
# train, predict and evaluate
mod = train(lrn, task)
pred = predict(mod, task)
performance(pred, measure = list(multilabel.hamloss, multilabel.subset01, multilabel.f1))
# the next call basically has the same structure for any multilabel meta wrapper
getMultilabelBinaryPerformances(pred, measures = list(mmce, auc))
# above works also with predictions from resample!
```

`makeMultilabelTask` *Create a multilabel task.*

Description

Create a multilabel task.

Usage

```
makeMultilabelTask(
  id = deparse(substitute(data)),
  data,
  target,
  weights = NULL,
  blocking = NULL,
  coordinates = NULL,
  fixup.data = "warn",
  check.data = TRUE
)
```

Arguments

id	(character(1)) Id string for object. Default is the name of the R variable passed to data.
data	(data.frame) A data frame containing the features and target variable(s).
target	(character(1) character(2) character(n.classes)) Name(s) of the target variable(s). For survival analysis these are the names of the survival time and event columns, so it has length 2. For multilabel classification it contains the names of the logical columns that encode whether a label is present or not and its length corresponds to the number of classes.
weights	(numeric) Optional, non-negative case weight vector to be used during fitting. Cannot be set for cost-sensitive learning. Default is NULL which means no (= equal) weights.
blocking	(factor) An optional factor of the same length as the number of observations. Observations with the same blocking level “belong together”. Specifically, they are either put all in the training or the test set during a resampling iteration. Default is NULL which means no blocking.
coordinates	(data.frame) Coordinates of a spatial data set that will be used for spatial partitioning of the data in a spatial cross-validation resampling setting. Coordinates have to be numeric values. Provided data.frame needs to have the same number of rows as data and consist of at least two dimensions.
fixup.data	(character(1)) Should some basic cleaning up of data be performed? Currently this means removing empty factor levels for the columns. Possible choices are: “no” = Don’t do it. “warn” = Do it but warn about it. “quiet” = Do it but keep silent. Default is “warn”.
check.data	(logical(1)) Should sanity of data be checked initially at task creation? You should have good reasons to turn this off (one might be speed). Default is TRUE.

Details

For multilabel classification we assume that the presence of labels is encoded via logical columns in data. The name of the column specifies the name of the label. `target` is then a char vector that points to these columns.

Note

For multilabel classification we assume that the presence of labels is encoded via logical columns in data. The name of the column specifies the name of the label. `target` is then a char vector that points to these columns.

See Also

[Task](#) [ClassifTask](#) [ClusterTask](#) [CostSensTask](#) [RegrTask](#) [SurvTask](#)

makeOverBaggingWrapper

Fuse learner with the bagging technique and oversampling for imbalance correction.

Description

Fuses a classification learner for binary classification with an over-bagging method for imbalance correction when we have strongly unequal class sizes. Creates a learner object, which can be used like any other learner object. Models can easily be accessed via [getLearnerModel](#).

OverBagging is implemented as follows: For each iteration a random data subset is sampled. Class examples are oversampled with replacement with a given rate. Members of the other class are either simply copied into each bag, or bootstrapped with replacement until we have as many majority class examples as in the original training data. Features are currently not changed or sampled.

Prediction works as follows: For classification we do majority voting to create a discrete label and probabilities are predicted by considering the proportions of all predicted labels.

Usage

```
makeOverBaggingWrapper(  
  learner,  
  obw.iters = 10L,  
  obw.rate = 1,  
  obw.maxcl = "boot",  
  obw.cl = NULL  
)
```


Arguments

learner	(Learner character(1)) The learner. If you pass a string the learner will be created via makeLearner .
obw.iters	(integer(1)) Number of fitted models in bagging. Default is 10.
obw.rate	(numeric(1)) Factor to upsample a class in each bag. Must be between 1 and Inf, where 1 means no oversampling and 2 would mean doubling the class size. Default is 1.
obw.maxcl	(character(1)) How should other class (usually larger class) be handled? “all” means every instance of the class gets in each bag, “boot” means the class instances are bootstrapped in each iteration. Default is “boot”.
obw.cl	(character(1)) Which class should be over- or undersampled. If NULL, makeOverBaggingWrapper will take the smaller class.

Value

[Learner](#).

See Also

Other imbalancecy: [makeUndersampleWrapper\(\)](#), [oversample\(\)](#), [smote\(\)](#)

Other wrapper: [makeBaggingWrapper\(\)](#), [makeClassificationViaRegressionWrapper\(\)](#), [makeConstantClassWrapper\(\)](#), [makeCostSensClassifWrapper\(\)](#), [makeCostSensRegrWrapper\(\)](#), [makeDownsampleWrapper\(\)](#), [makeDummyFeaturesWrapper\(\)](#), [makeExtractFDAFeatsWrapper\(\)](#), [makeFeatSelWrapper\(\)](#), [makeFilterWrapper\(\)](#), [makeImputeWrapper\(\)](#), [makeMulticlassWrapper\(\)](#), [makeMultilabelBinaryRelevanceWrapper\(\)](#), [makeMultilabelClassifierChainsWrapper\(\)](#), [makeMultilabelDBRWrapper\(\)](#), [makeMultilabelNestedStackingWrapper\(\)](#), [makeMultilabelStackingWrapper\(\)](#), [makePreprocWrapperCaret\(\)](#), [makePreprocWrapper\(\)](#), [makeRemoveConstantFeaturesWrapper\(\)](#), [makeSMOTEWrapper\(\)](#), [makeTuneWrapper\(\)](#), [makeUndersampleWrapper\(\)](#), [makeWeightedClassesWrapper\(\)](#)

makePreprocWrapper *Fuse learner with preprocessing.*

Description

Fuses a base learner with a preprocessing method. Creates a learner object, which can be used like any other learner object, but which internally preprocesses the data as requested. If the train or predict function is called on data / a task, the preprocessing is always performed automatically.

Usage

```
makePreprocWrapper(
  learner,
  train,
  predict,
  par.set = makeParamSet(),
  par.vals = list()
)
```

Arguments

learner	(Learner character(1)) The learner. If you pass a string the learner will be created via makeLearner .
train	(function(data, target, args)) Function to preprocess the data before training. target is a string and denotes the target variable in data. args is a list of further arguments and parameters to influence the preprocessing. Must return a list(data, control), where data is the preprocessed data and control stores all information necessary to do the preprocessing before predictions.
predict	(function(data, target, args, control)) Function to preprocess the data before prediction. target is a string and denotes the target variable in data. args are the args that were passed to train. control is the object you returned in train. Must return the processed data.
par.set	(ParamHelpers::ParamSet) Parameter set of ParamHelpers::LearnerParam objects to describe the parameters in args. Default is empty set.
par.vals	(list) Named list of default values for params in args respectively par.set. Default is empty list.

Value

([Learner](#)).

See Also

Other wrapper: [makeBaggingWrapper\(\)](#), [makeClassificationViaRegressionWrapper\(\)](#), [makeConstantClassWrapper\(\)](#), [makeCostSensClassifWrapper\(\)](#), [makeCostSensRegrWrapper\(\)](#), [makeDownsampleWrapper\(\)](#), [makeDummyFeaturesWrapper\(\)](#), [makeExtractFDAFeatsWrapper\(\)](#), [makeFeatSelWrapper\(\)](#), [makeFilterWrapper\(\)](#), [makeImputeWrapper\(\)](#), [makeMulticlassWrapper\(\)](#), [makeMultilabelBinaryRelevanceWrapper\(\)](#), [makeMultilabelClassifierChainsWrapper\(\)](#), [makeMultilabelDBRWrapper\(\)](#), [makeMultilabelNestedStackingWrapper\(\)](#), [makeMultilabelStackingWrapper\(\)](#), [makeOverBaggingWrapper\(\)](#), [makePreprocWrapperCaret\(\)](#), [makeRemoveConstantFeaturesWrapper\(\)](#), [makeSMOTERWrapper\(\)](#), [makeTuneWrapper\(\)](#), [makeUndersampleWrapper\(\)](#), [makeWeightedClassesWrapper\(\)](#)

 makePreprocWrapperCaret

Fuse learner with preprocessing.

Description

Fuses a learner with preprocessing methods provided by [caret::preProcess](#). Before training the preprocessing will be performed and the preprocessing model will be stored. Before prediction the preprocessing model will transform the test data according to the trained model.

After being wrapped the learner will support missing values although this will only be the case if `ppc.knnImpute`, `ppc.bagImpute` or `ppc.medianImpute` is set to TRUE.

Usage

```
makePreprocWrapperCaret(learner, ...)
```

Arguments

learner	(Learner character(1)) The learner. If you pass a string the learner will be created via makeLearner .
...	(any) See caret::preProcess for parameters not listed above. If you use them you might want to define them in the <code>add.par.set</code> so that they can be tuned.

Value

[Learner](#).

See Also

Other wrapper: [makeBaggingWrapper\(\)](#), [makeClassificationViaRegressionWrapper\(\)](#), [makeConstantClassWrapper\(\)](#), [makeCostSensClassifWrapper\(\)](#), [makeCostSensRegrWrapper\(\)](#), [makeDownsampleWrapper\(\)](#), [makeDummyFeaturesWrapper\(\)](#), [makeExtractFDAFeatsWrapper\(\)](#), [makeFeatSelWrapper\(\)](#), [makeFilterWrapper\(\)](#), [makeImputeWrapper\(\)](#), [makeMulticlassWrapper\(\)](#), [makeMultilabelBinaryRelevanceWrapper\(\)](#), [makeMultilabelClassifierChainsWrapper\(\)](#), [makeMultilabelDBRWrapper\(\)](#), [makeMultilabelNestedStackingWrapper\(\)](#), [makeMultilabelStackingWrapper\(\)](#), [makeOverBaggingWrapper\(\)](#), [makePreprocWrapper\(\)](#), [makeRemoveConstantFeatWrapper\(\)](#), [makeSMOTEWrapper\(\)](#), [makeTuneWrapper\(\)](#), [makeUndersampleWrapper\(\)](#), [makeWeightedClassesWrapper\(\)](#)

makeRegrTask	<i>Create a regression task.</i>
--------------	----------------------------------

Description

Create a regression task.

Usage

```
makeRegrTask(
  id = deparse(substitute(data)),
  data,
  target,
  weights = NULL,
  blocking = NULL,
  coordinates = NULL,
  fixup.data = "warn",
  check.data = TRUE
)
```

Arguments

id	(character(1)) Id string for object. Default is the name of the R variable passed to data.
data	(data.frame) A data frame containing the features and target variable(s).
target	(character(1) character(2) character(n.classes)) Name(s) of the target variable(s). For survival analysis these are the names of the survival time and event columns, so it has length 2. For multilabel classification it contains the names of the logical columns that encode whether a label is present or not and its length corresponds to the number of classes.
weights	(numeric) Optional, non-negative case weight vector to be used during fitting. Cannot be set for cost-sensitive learning. Default is NULL which means no (= equal) weights.
blocking	(factor) An optional factor of the same length as the number of observations. Observations with the same blocking level “belong together”. Specifically, they are either put all in the training or the test set during a resampling iteration. Default is NULL which means no blocking.
coordinates	(data.frame) Coordinates of a spatial data set that will be used for spatial partitioning of the data in a spatial cross-validation resampling setting. Coordinates have to be numeric values. Provided data.frame needs to have the same number of rows as data and consist of at least two dimensions.

fixup.data	(character(1)) Should some basic cleaning up of data be performed? Currently this means removing empty factor levels for the columns. Possible choices are: “no” = Don’t do it. “warn” = Do it but warn about it. “quiet” = Do it but keep silent. Default is “warn”.
check.data	(logical(1)) Should sanity of data be checked initially at task creation? You should have good reasons to turn this off (one might be speed). Default is TRUE.

See Also

[Task](#) [ClassifTask](#) [CostSensTask](#) [ClusterTask](#) [MultilabelTask](#) [SurvTask](#)

makeRemoveConstantFeaturesWrapper

Fuse learner with removal of constant features preprocessing.

Description

Fuses a base learner with the preprocessing implemented in [removeConstantFeatures](#).

Usage

```
makeRemoveConstantFeaturesWrapper(
  learner,
  perc = 0,
  dont.rm = character(0L),
  na.ignore = FALSE,
  tol = .Machine$double.eps^0.5
)
```

Arguments

learner	(Learner character(1)) The learner. If you pass a string the learner will be created via makeLearner .
perc	(numeric(1)) The percentage of a feature values in [0, 1) that must differ from the mode value. Default is 0, which means only constant features with exactly one observed level are removed.
dont.rm	(character) Names of the columns which must not be deleted. Default is no columns.
na.ignore	(logical(1)) Should NAs be ignored in the percentage calculation? (Or should they be treated as a single, extra level in the percentage calculation?) Note that if the feature has only missing values, it is always removed. Default is FALSE.
tol	(numeric(1)) Numerical tolerance to treat two numbers as equal. Variables stored as double will get rounded accordingly before computing the mode. Default is <code>sqrt(.Machine\$double.eps)</code> .

Value

[Learner](#).

See Also

Other wrapper: [makeBaggingWrapper\(\)](#), [makeClassificationViaRegressionWrapper\(\)](#), [makeConstantClassWrapper\(\)](#), [makeCostSensClassifWrapper\(\)](#), [makeCostSensRegrWrapper\(\)](#), [makeDownsampleWrapper\(\)](#), [makeDummyFeaturesWrapper\(\)](#), [makeExtractFDAFeatsWrapper\(\)](#), [makeFeatSelWrapper\(\)](#), [makeFilterWrapper\(\)](#), [makeImputeWrapper\(\)](#), [makeMulticlassWrapper\(\)](#), [makeMultilabelBinaryRelevanceWrapper\(\)](#), [makeMultilabelClassifierChainsWrapper\(\)](#), [makeMultilabelDBRWrapper\(\)](#), [makeMultilabelNestedStackingWrapper\(\)](#), [makeMultilabelStackingWrapper\(\)](#), [makeOverBaggingWrapper\(\)](#), [makePreprocWrapperCaret\(\)](#), [makePreprocWrapper\(\)](#), [makeSMOTEWrapper\(\)](#), [makeTuneWrapper\(\)](#), [makeUndersampleWrapper\(\)](#), [makeWeightedClassesWrapper\(\)](#)

<code>makeResampleDesc</code>	<i>Create a description object for a resampling strategy.</i>
-------------------------------	---

Description

A description of a resampling algorithm contains all necessary information to create a [ResampleInstance](#), when given the size of the data set.

Usage

```
makeResampleDesc(
  method,
  predict = "test",
  ...,
  stratify = FALSE,
  stratify.cols = NULL,
  fixed = FALSE,
  blocking.cv = FALSE
)
```

Arguments

<code>method</code>	(character(1)) “CV” for cross-validation, “LOO” for leave-one-out, “RepCV” for repeated cross-validation, “Bootstrap” for out-of-bag bootstrap, “Subsample” for subsampling, “Holdout” for holdout, “GrowingWindowCV” for growing window cross-validation, “FixedWindowCV” for fixed window cross validation.
<code>predict</code>	(character(1)) What to predict during resampling: “train”, “test” or “both” sets. Default is “test”.
<code>...</code>	(any) Further parameters for strategies.

	<p>iters (integer(1)) Number of iterations, for “CV”, “Subsample” and “Bootstrap”.</p> <p>split (numeric(1)) Proportion of training cases for “Holdout” and “Subsample” between 0 and 1. Default is 2 / 3.</p> <p>reps (integer(1)) Repeats for “RepCV”. Here <code>iters = folds * reps</code>. Default is 10.</p> <p>folds (integer(1)) Folds in the repeated CV for RepCV. Here <code>iters = folds * reps</code>. Default is 10.</p> <p>horizon (numeric(1)) Number of observations in the forecast test set for “GrowingWindowCV” and “FixedWindowCV”. When <code>horizon > 1</code> this will be treated as the number of observations to forecast, else it will be a fraction of the initial window. IE, for 100 observations, initial window of .5, and horizon of .2, the test set will have 10 observations. Default is 1.</p> <p>initial.window (numeric(1)) Fraction of observations to start with in the training set for “GrowingWindowCV” and “FixedWindowCV”. When <code>initial.window > 1</code> this will be treated as the number of observations in the initial window, else it will be treated as the fraction of observations to have in the initial window. Default is 0.5.</p> <p>skip (numeric(1)) How many resamples to skip to thin the total amount for “GrowingWindowCV” and “FixedWindowCV”. This is passed through as the “by” argument in <code>seq()</code>. When <code>skip > 1</code> this will be treated as the increment of the sequence of resampling indices, else it will be a fraction of the total training indices. IE for 100 training sets and a value of .2, the increment of the resampling indices will be 20. Default is “horizon” which gives mutually exclusive chunks of test indices.</p>
stratify	<p>(logical(1))</p> <p>Should stratification be done for the target variable? For classification tasks, this means that the resampling strategy is applied to all classes individually and the resulting index sets are joined to make sure that the proportion of observations in each training set is as in the original data set. Useful for imbalanced class sizes. For survival tasks stratification is done on the events, resulting in training sets with comparable censoring rates.</p>
stratify.cols	<p>(character)</p> <p>Stratify on specific columns referenced by name. All columns have to be factor or integer. Note that you have to ensure yourself that stratification is possible, i.e. that each strata contains enough observations. This argument and <code>stratify</code> are mutually exclusive.</p>
fixed	<p>(logical(1))</p> <p>Whether indices supplied via argument ‘blocking’ in the task should be used as fully pre-defined indices. Default is FALSE which means they will be used following the ‘blocking’ approach. <code>fixed</code> only works with ResampleDesc CV and the supplied indices must match the number of observations. When <code>fixed = TRUE</code>, the <code>iters</code> argument will be ignored and is internally set to the number of supplied factor levels in <code>blocking</code>.</p>
blocking.cv	<p>(logical(1))</p> <p>Should ‘blocking’ be used in CV? Default to FALSE. This is different to <code>fixed</code></p>

= TRUE and cannot be combined. Please check the mlr online tutorial for more details.

Details

Some notes on some special strategies:

Repeated cross-validation Use “RepCV”. Then you have to set the aggregation function for your preferred performance measure to “testgroup.mean” via [setAggregation](#).

B632 bootstrap Use “Bootstrap” for bootstrap and set predict to “both”. Then you have to set the aggregation function for your preferred performance measure to “b632” via [setAggregation](#).

B632+ bootstrap Use “Bootstrap” for bootstrap and set predict to “both”. Then you have to set the aggregation function for your preferred performance measure to “b632plus” via [setAggregation](#).

Fixed Holdout set Use [makeFixedHoldoutInstance](#).

Object slots:

id (character(1)) Name of resampling strategy.

iters (integer(1)) Number of iterations. Note that this is always the complete number of generated train/test sets, so for a 10-times repeated 5fold cross-validation it would be 50.

predict (character(1)) See argument.

stratify (logical(1)) See argument.

All parameters passed in ... under the respective argument name See arguments.

Value

([ResampleDesc](#)).

Standard ResampleDesc objects

For common resampling strategies you can save some typing by using the following description objects:

hout holdout a.k.a. test sample estimation (two-thirds training set, one-third testing set)

cv2 2-fold cross-validation

cv3 3-fold cross-validation

cv5 5-fold cross-validation

cv10 10-fold cross-validation

See Also

Other resample: [ResamplePrediction](#), [ResampleResult](#), [addRRMeasure\(\)](#), [getRRPredictionList\(\)](#), [getRRPredictions\(\)](#), [getRRTaskDescription\(\)](#), [getRRTaskDesc\(\)](#), [makeResampleInstance\(\)](#), [resample\(\)](#)

Examples

```
# Bootstrapping
makeResampleDesc("Bootstrap", iters = 10)
makeResampleDesc("Bootstrap", iters = 10, predict = "both")

# Subsampling
makeResampleDesc("Subsample", iters = 10, split = 3 / 4)
makeResampleDesc("Subsample", iters = 10)

# Holdout a.k.a. test sample estimation
makeResampleDesc("Holdout")
```

makeResampleInstance *Instantiates a resampling strategy object.*

Description

This class encapsulates training and test sets generated from the data set for a number of iterations. It mainly stores a set of integer vectors indicating the training and test examples for each iteration.

Usage

```
makeResampleInstance(desc, task, size, ...)
```

Arguments

desc	(ResampleDesc character(1)) Resampling description object or name of resampling strategy. In the latter case makeResampleDesc will be called internally on the string.
task	(Task) Data of task to resample from. Prefer to pass this instead of size.
size	(integer) Size of the data set to resample. Can be used instead of task.
...	(any) Passed down to makeResampleDesc in case you passed a string in desc. Otherwise ignored.

Details

Object slots:

desc ([ResampleDesc](#)) See argument.

size ([integer\(1\)](#)) See argument.

train.inds ([list of integer](#)) List of of training indices for all iterations.

test.inds ([list of integer](#)) List of of test indices for all iterations.

group ([factor](#)) Optional grouping of resampling iterations. This encodes whether specific iterations 'belong together' (e.g. repeated CV), and it can later be used to aggregate performance values accordingly. Default is 'factor()'.

Value

(ResampleInstance).

See Also

Other resample: ResamplePrediction, ResampleResult, addRRMeasure(), getRRPredictionList(), getRRPredictions(), getRRTaskDescription(), getRRTaskDesc(), makeResampleDesc(), resample()

Examples

```
rdesc = makeResampleDesc("Bootstrap", iters = 10)
rin = makeResampleInstance(rdesc, task = iris.task)

rdesc = makeResampleDesc("CV", iters = 50)
rin = makeResampleInstance(rdesc, size = nrow(iris))

rin = makeResampleInstance("CV", iters = 10, task = iris.task)
```

```
makeRLearner.classif.fdausc.glm
```

Classification of functional data by Generalized Linear Models.

Description

Learner for classification using Generalized Linear Models.

Usage

```
## S3 method for class 'classif.fdausc.glm'
makeRLearner()
```

```
makeRLearner.classif.fdausc.kernel
```

Learner for kernel classification for functional data.

Description

Learner for kernel Classification.

Usage

```
## S3 method for class 'classif.fdausc.kernel'
makeRLearner()
```

```
makeRLearner.classif.fdausc.np
```

Learner for nonparametric classification for functional data.

Description

Learner for Nonparametric Supervised Classification.

Usage

```
## S3 method for class 'classif.fdausc.np'
makeRLearner()
```

```
makeSMOTEWrapper
```

Fuse learner with SMOTE oversampling for imbalance correction in binary classification.

Description

Creates a learner object, which can be used like any other learner object. Internally uses [smote](#) before every model fit.

Note that observation weights do not influence the sampling and are simply passed down to the next learner.

Usage

```
makeSMOTEWrapper(
  learner,
  sw.rate = 1,
  sw.nn = 5L,
  sw.standardize = TRUE,
  sw.alt.logic = FALSE
)
```

Arguments

learner	(Learner character(1)) The learner. If you pass a string the learner will be created via makeLearner .
sw.rate	(numeric(1)) Factor to oversample the smaller class. Must be between 1 and Inf, where 1 means no oversampling and 2 would mean doubling the class size. Default is 1.
sw.nn	(integer(1)) Number of nearest neighbors to consider. Default is 5.

<code>sw.standardize</code>	<code>(logical(1))</code> Standardize input variables before calculating the nearest neighbors for data sets with numeric input variables only. For mixed variables (numeric and factor) the gower distance is used and variables are standardized anyway. Default is TRUE.
<code>sw.alt.logic</code>	<code>(logical(1))</code> Use an alternative logic for selection of minority class observations. Instead of sampling a minority class element AND one of its nearest neighbors, each minority class element is taken multiple times (depending on rate) for the interpolation and only the corresponding nearest neighbor is sampled. Default is FALSE.

Value

[Learner](#).

See Also

Other wrapper: [makeBaggingWrapper\(\)](#), [makeClassificationViaRegressionWrapper\(\)](#), [makeConstantClassWrapper\(\)](#), [makeCostSensClassifWrapper\(\)](#), [makeCostSensRegrWrapper\(\)](#), [makeDownsampleWrapper\(\)](#), [makeDummyFeaturesWrapper\(\)](#), [makeExtractFDAFeatsWrapper\(\)](#), [makeFeatSelWrapper\(\)](#), [makeFilterWrapper\(\)](#), [makeImputeWrapper\(\)](#), [makeMulticlassWrapper\(\)](#), [makeMultilabelBinaryRelevanceWrapper\(\)](#), [makeMultilabelClassifierChainsWrapper\(\)](#), [makeMultilabelDBRWrapper\(\)](#), [makeMultilabelNestedStackingWrapper\(\)](#), [makeMultilabelStackingWrapper\(\)](#), [makeOverBaggingWrapper\(\)](#), [makePreprocWrapperCaret\(\)](#), [makePreprocWrapper\(\)](#), [makeRemoveConstantFeaturesWrapper\(\)](#), [makeTuneWrapper\(\)](#), [makeUndersampleWrapper\(\)](#), [makeWeightedClassesWrapper\(\)](#)

`makeStackedLearner` *Create a stacked learner object.*

Description

A stacked learner uses predictions of several base learners and fits a super learner using these predictions as features in order to predict the outcome. The following stacking methods are available:

- `average`
Averaging of base learner predictions without weights.
- `stack.nocv`
Fits the super learner, where in-sample predictions of the base learners are used.
- `stack.cv`
Fits the super learner, where the base learner predictions are computed by cross-validated predictions (the resampling strategy can be set via the `resampling` argument).
- `hill.climb`
Select a subset of base learner predictions by hill climbing algorithm.
- `compress`
Train a neural network to compress the model from a collection of base learners.

Usage

```
makeStackedLearner(
  base.learners,
  super.learner = NULL,
  predict.type = NULL,
  method = "stack.nocv",
  use.feats = FALSE,
  resampling = NULL,
  parset = list()
)
```

Arguments

- `base.learners` ((list of) [Learner](#))
A list of learners created with `makeLearner`.
- `super.learner` ([Learner](#) | `character(1)`)
The super learner that makes the final prediction based on the base learners. If you pass a string, the super learner will be created via `makeLearner`. Not used for `method = 'average'`. Default is `NULL`.
- `predict.type` (`character(1)`)
Sets the type of the final prediction for `method = 'average'`. For other methods, the predict type should be set within `super.learner`. If the type of the base learner prediction, which is set up within `base.learners`, is
- "prob"
then `predict.type = 'prob'` will use the average of all base learner predictions and `predict.type = 'response'` will use the class with highest probability as final prediction.
 - "response"
then, for classification tasks with `predict.type = 'prob'`, the final prediction will be the relative frequency based on the predicted base learner classes and classification tasks with `predict.type = 'response'` will use majority vote of the base learner predictions to determine the final prediction. For regression tasks, the final prediction will be the average of the base learner predictions.
- `method` (`character(1)`)
"average" for averaging the predictions of the base learners, "stack.nocv" for building a super learner using the predictions of the base learners, "stack.cv" for building a super learner using cross-validated predictions of the base learners. "hill.climb" for averaging the predictions of the base learners, with the weights learned from hill climbing algorithm and "compress" for compressing the model to mimic the predictions of a collection of base learners while speeding up the predictions and reducing the size of the model. Default is "stack.nocv",
- `use.feats` (`logical(1)`)
Whether the original features should also be passed to the super learner. Not used for `method = 'average'`. Default is `FALSE`.

resampling	(ResampleDesc) Resampling strategy for method = 'stack.cv'. Currently only CV is allowed for resampling. The default NULL uses 5-fold CV.
parset	the parameters for hill.climb method, including <ul style="list-style-type: none"> • replace Whether a base learner can be selected more than once. • init Number of best models being included before the selection algorithm. • bagprob The proportion of models being considered in one round of selection. • bagtime The number of rounds of the bagging selection. • metric The result evaluation metric function taking two parameters pred and true, the smaller the score the better. <p>the parameters for compress method, including</p> <ul style="list-style-type: none"> • k the size multiplier of the generated data • prob the probability to exchange values • s the standard deviation of each numerical feature

Examples

```
# Classification
data(iris)
tsk = makeClassifTask(data = iris, target = "Species")
base = c("classif.rpart", "classif.lda", "classif.svm")
lrns = lapply(base, makeLearner)
lrns = lapply(lrns, setPredictType, "prob")
m = makeStackedLearner(base.learners = lrns,
  predict.type = "prob", method = "hill.climb")
tmp = train(m, tsk)
res = predict(tmp, tsk)

# Regression
data(BostonHousing, package = "mlbench")
tsk = makeRegrTask(data = BostonHousing, target = "medv")
base = c("regr.rpart", "regr.svm")
lrns = lapply(base, makeLearner)
m = makeStackedLearner(base.learners = lrns,
  predict.type = "response", method = "compress")
tmp = train(m, tsk)
res = predict(tmp, tsk)
```

makeSurvTask	<i>Create a survival task.</i>
--------------	--------------------------------

Description

Create a survival task.

Usage

```
makeSurvTask(  
  id = deparse(substitute(data)),  
  data,  
  target,  
  weights = NULL,  
  blocking = NULL,  
  coordinates = NULL,  
  fixup.data = "warn",  
  check.data = TRUE  
)
```

Arguments

id	(character(1)) Id string for object. Default is the name of the R variable passed to data.
data	(data.frame) A data frame containing the features and target variable(s).
target	(character(1) character(2) character(n.classes)) Name(s) of the target variable(s). For survival analysis these are the names of the survival time and event columns, so it has length 2. For multilabel classification it contains the names of the logical columns that encode whether a label is present or not and its length corresponds to the number of classes.
weights	(numeric) Optional, non-negative case weight vector to be used during fitting. Cannot be set for cost-sensitive learning. Default is NULL which means no (= equal) weights.
blocking	(factor) An optional factor of the same length as the number of observations. Observations with the same blocking level “belong together”. Specifically, they are either put all in the training or the test set during a resampling iteration. Default is NULL which means no blocking.
coordinates	(data.frame) Coordinates of a spatial data set that will be used for spatial partitioning of the data in a spatial cross-validation resampling setting. Coordinates have to be numeric values. Provided data.frame needs to have the same number of rows as data and consist of at least two dimensions.

fixup.data	(character(1)) Should some basic cleaning up of data be performed? Currently this means removing empty factor levels for the columns. Possible choices are: “no” = Don’t do it. “warn” = Do it but warn about it. “quiet” = Do it but keep silent. Default is “warn”.
check.data	(logical(1)) Should sanity of data be checked initially at task creation? You should have good reasons to turn this off (one might be speed). Default is TRUE.

See Also

[Task ClassifTask](#) [ClusterTask](#) [CostSensTask](#) [MultilabelTask](#) [RegrTask](#)

makeTuneControlCMAES *Create control object for hyperparameter tuning with CMAES.*

Description

CMA Evolution Strategy with method `cmaes::cma_es`. Can handle numeric(vector) and integer(vector) hyperparameters, but no dependencies. For integers the internally proposed numeric values are automatically rounded. The sigma variance parameter is initialized to 1/4 of the span of box-constraints per parameter dimension.

Usage

```
makeTuneControlCMAES(
  same.resampling.instance = TRUE,
  impute.val = NULL,
  start = NULL,
  tune.threshold = FALSE,
  tune.threshold.args = list(),
  log.fun = "default",
  final.dw.perc = NULL,
  budget = NULL,
  ...
)
```

Arguments

same.resampling.instance	(logical(1)) Should the same resampling instance be used for all evaluations to reduce variance? Default is TRUE.
impute.val	(numeric) If something goes wrong during optimization (e.g. the learner crashes), this value is fed back to the tuner, so the tuning algorithm does not abort. It is not stored in the optimization path, an NA and a corresponding error message are

logged instead. Note that this value is later multiplied by -1 for maximization measures internally, so you need to enter a larger positive value for maximization here as well. Default is the worst obtainable value of the performance measure you optimize for when you aggregate by mean value, or Inf instead. For multi-criteria optimization pass a vector of imputation values, one for each of your measures, in the same order as your measures.

start	(list) Named list of initial parameter values.
tune.threshold	(logical(1)) Should the threshold be tuned for the measure at hand, after each hyperparameter evaluation, via <code>tuneThreshold</code> ? Only works for classification if the predict type is “prob”. Default is FALSE.
tune.threshold.args	(list) Further arguments for threshold tuning that are passed down to <code>tuneThreshold</code> . Default is none.
log.fun	(function character(1)) Function used for logging. If set to “default” (the default), the evaluated design points, the resulting performances, and the runtime will be reported. If set to “memory” the memory usage for each evaluation will also be displayed, with character(1) small increase in run time. Otherwise character(1) function with arguments learner, resampling, measures, par.set, control, opt.path, dob, x, y, remove.nas, stage and prev.stage is expected. The default displays the performance measures, the time needed for evaluating, the currently used memory and the max memory ever used before (the latter two both taken from <code>gc</code>). See the implementation for details.
final.dw.perc	(boolean) If a Learner wrapped by a <code>makeDownsampleWrapper</code> is used, you can define the value of <code>dw.perc</code> which is used to train the Learner with the final parameter setting found by the tuning. Default is NULL which will not change anything.
budget	(integer(1)) Maximum budget for tuning. This value restricts the number of function evaluations. The budget corresponds to the product of the number of generations (<code>maxit</code>) and the number of offsprings per generation (<code>lambda</code>).
...	(any) Further control parameters passed to the control arguments of <code>cmaes::cma_es</code> or <code>GenSA::GenSA</code> , as well as towards the <code>tunerConfig</code> argument of <code>irace::irace</code> .

Value

(`TuneControlCMAES`)

See Also

Other tune: `TuneControl`, `getNestedTuneResultsOptPathDf()`, `getNestedTuneResultsX()`, `getResamplingIndices()`, `getTuneResult()`, `makeModelMultiplexerParamSet()`, `makeModelMultiplexer()`, `makeTuneControlDesign()`, `makeTuneControlGenSA()`, `makeTuneControlGrid()`, `makeTuneControlIrace()`, `makeTuneControlMBO()`, `makeTuneControlRandom()`, `makeTuneWrapper()`, `tuneParams()`, `tuneThreshold()`

`makeTuneControlDesign` *Create control object for hyperparameter tuning with predefined design.*

Description

Completely pre-specify a data.frame of design points to be evaluated during tuning. All kinds of parameter types can be handled.

Usage

```
makeTuneControlDesign(
  same.resampling.instance = TRUE,
  impute.val = NULL,
  design = NULL,
  tune.threshold = FALSE,
  tune.threshold.args = list(),
  log.fun = "default"
)
```

Arguments

<code>same.resampling.instance</code>	(logical(1)) Should the same resampling instance be used for all evaluations to reduce variance? Default is TRUE.
<code>impute.val</code>	(numeric) If something goes wrong during optimization (e.g. the learner crashes), this value is fed back to the tuner, so the tuning algorithm does not abort. It is not stored in the optimization path, an NA and a corresponding error message are logged instead. Note that this value is later multiplied by -1 for maximization measures internally, so you need to enter a larger positive value for maximization here as well. Default is the worst obtainable value of the performance measure you optimize for when you aggregate by mean value, or Inf instead. For multi-criteria optimization pass a vector of imputation values, one for each of your measures, in the same order as your measures.
<code>design</code>	(data.frame) data.frame containing the different parameter settings to be evaluated. The columns have to be named according to the ParamSet which will be used in <code>tune()</code> . Proper designs can be created with ParamHelpers::generateDesign for instance.
<code>tune.threshold</code>	(logical(1)) Should the threshold be tuned for the measure at hand, after each hyperparameter evaluation, via tuneThreshold? Only works for classification if the predict type is "prob". Default is FALSE.

tune.threshold.args	(list) Further arguments for threshold tuning that are passed down to tuneThreshold . Default is none.
log.fun	(function character(1)) Function used for logging. If set to “default” (the default), the evaluated design points, the resulting performances, and the runtime will be reported. If set to “memory” the memory usage for each evaluation will also be displayed, with character(1) small increase in run time. Otherwise character(1) function with arguments learner, resampling, measures, par.set, control, opt.path, dob, x, y, remove.nas, stage and prev.stage is expected. The default displays the performance measures, the time needed for evaluating, the currently used memory and the max memory ever used before (the latter two both taken from gc). See the implementation for details.

Value

(TuneControlDesign)

See Also

Other tune: [TuneControl](#), [getNestedTuneResultsOptPathDf\(\)](#), [getNestedTuneResultsX\(\)](#), [getResamplingIndices\(\)](#), [getTuneResult\(\)](#), [makeModelMultiplexerParamSet\(\)](#), [makeModelMultiplexer\(\)](#), [makeTuneControlCMAES\(\)](#), [makeTuneControlGenSA\(\)](#), [makeTuneControlGrid\(\)](#), [makeTuneControlIrace\(\)](#), [makeTuneControlMBO\(\)](#), [makeTuneControlRandom\(\)](#), [makeTuneWrapper\(\)](#), [tuneParams\(\)](#), [tuneThreshold\(\)](#)

makeTuneControlGenSA *Create control object for hyperparameter tuning with GenSA.*

Description

Generalized simulated annealing with method [GenSA::GenSA](#). Can handle numeric(vector) and integer(vector) hyperparameters, but no dependencies. For integers the internally proposed numeric values are automatically rounded.

Usage

```
makeTuneControlGenSA(
  same.resampling.instance = TRUE,
  impute.val = NULL,
  start = NULL,
  tune.threshold = FALSE,
  tune.threshold.args = list(),
  log.fun = "default",
  final.dw.perc = NULL,
  budget = NULL,
  ...
)
```

Arguments

same.resampling.instance	(logical(1)) Should the same resampling instance be used for all evaluations to reduce variance? Default is TRUE.
impute.val	(numeric) If something goes wrong during optimization (e.g. the learner crashes), this value is fed back to the tuner, so the tuning algorithm does not abort. It is not stored in the optimization path, an NA and a corresponding error message are logged instead. Note that this value is later multiplied by -1 for maximization measures internally, so you need to enter a larger positive value for maximization here as well. Default is the worst obtainable value of the performance measure you optimize for when you aggregate by mean value, or Inf instead. For multi-criteria optimization pass a vector of imputation values, one for each of your measures, in the same order as your measures.
start	(list) Named list of initial parameter values.
tune.threshold	(logical(1)) Should the threshold be tuned for the measure at hand, after each hyperparameter evaluation, via <code>tuneThreshold</code> ? Only works for classification if the predict type is “prob”. Default is FALSE.
tune.threshold.args	(list) Further arguments for threshold tuning that are passed down to <code>tuneThreshold</code> . Default is none.
log.fun	(function character(1)) Function used for logging. If set to “default” (the default), the evaluated design points, the resulting performances, and the runtime will be reported. If set to “memory” the memory usage for each evaluation will also be displayed, with character(1) small increase in run time. Otherwise character(1) function with arguments learner, resampling, measures, par.set, control, opt.path, dob, x, y, remove.nas, stage and prev.stage is expected. The default displays the performance measures, the time needed for evaluating, the currently used memory and the max memory ever used before (the latter two both taken from <code>gc</code>). See the implementation for details.
final.dw.perc	(boolean) If a Learner wrapped by a <code>makeDownsampleWrapper</code> is used, you can define the value of <code>dw.perc</code> which is used to train the Learner with the final parameter setting found by the tuning. Default is NULL which will not change anything.
budget	(integer(1)) Maximum budget for tuning. This value restricts the number of function evaluations. <code>GenSA::GenSA</code> defines the budget via the argument <code>max.call</code> . However, one should note that this algorithm does not stop its local search before its end. This behavior might lead to an extension of the defined budget and will result in a warning.

... (any)
 Further control parameters passed to the control arguments of `cmaes::cma_es` or `GenSA::GenSA`, as well as towards the `tunerConfig` argument of `irace::irace`.

Value

(`TuneControlGenSA`).

See Also

Other tune: `TuneControl`, `getNestedTuneResultsOptPathDf()`, `getNestedTuneResultsX()`, `getResamplingIndices()`, `getTuneResult()`, `makeModelMultiplexerParamSet()`, `makeModelMultiplexer()`, `makeTuneControlCMAES()`, `makeTuneControlDesign()`, `makeTuneControlGrid()`, `makeTuneControlIrace()`, `makeTuneControlMBO()`, `makeTuneControlRandom()`, `makeTuneWrapper()`, `tuneParams()`, `tuneThreshold()`

`makeTuneControlGrid` *Create control object for hyperparameter tuning with grid search.*

Description

A basic grid search can handle all kinds of parameter types. You can either use their correct param type and resolution, or discretize them yourself by always using `ParamHelpers::makeDiscreteParam` in the `par.set` passed to `tuneParams`.

Usage

```
makeTuneControlGrid(
  same.resampling.instance = TRUE,
  impute.val = NULL,
  resolution = 10L,
  tune.threshold = FALSE,
  tune.threshold.args = list(),
  log.fun = "default",
  final.dw.perc = NULL,
  budget = NULL
)
```

Arguments

`same.resampling.instance` (logical(1))
 Should the same resampling instance be used for all evaluations to reduce variance? Default is TRUE.

`impute.val` (numeric)
 If something goes wrong during optimization (e.g. the learner crashes), this value is fed back to the tuner, so the tuning algorithm does not abort. It is not stored in the optimization path, an NA and a corresponding error message are

logged instead. Note that this value is later multiplied by -1 for maximization measures internally, so you need to enter a larger positive value for maximization here as well. Default is the worst obtainable value of the performance measure you optimize for when you aggregate by mean value, or Inf instead. For multi-criteria optimization pass a vector of imputation values, one for each of your measures, in the same order as your measures.

resolution	(integer) Resolution of the grid for each numeric/integer parameter in <code>par.set</code> . For vector parameters, it is the resolution per dimension. Either pass one resolution for all parameters, or a named vector. See ParamHelpers::generateGridDesign . Default is 10.
tune.threshold	(logical(1)) Should the threshold be tuned for the measure at hand, after each hyperparameter evaluation, via tuneThreshold? Only works for classification if the predict type is “prob”. Default is FALSE.
tune.threshold.args	(list) Further arguments for threshold tuning that are passed down to tuneThreshold . Default is none.
log.fun	(function character(1)) Function used for logging. If set to “default” (the default), the evaluated design points, the resulting performances, and the runtime will be reported. If set to “memory” the memory usage for each evaluation will also be displayed, with <code>character(1)</code> small increase in run time. Otherwise <code>character(1)</code> function with arguments <code>learner</code> , <code>resampling</code> , <code>measures</code> , <code>par.set</code> , <code>control</code> , <code>opt.path</code> , <code>dob</code> , <code>x</code> , <code>y</code> , <code>remove.nas</code> , <code>stage</code> and <code>prev.stage</code> is expected. The default displays the performance measures, the time needed for evaluating, the currently used memory and the max memory ever used before (the latter two both taken from gc). See the implementation for details.
final.dw.perc	(boolean) If a <code>Learner</code> wrapped by a makeDownsampleWrapper is used, you can define the value of <code>dw.perc</code> which is used to train the <code>Learner</code> with the final parameter setting found by the tuning. Default is NULL which will not change anything.
budget	(integer(1)) Maximum budget for tuning. This value restricts the number of function evaluations. If set, must equal the size of the grid.

Value

([TuneControlGrid](#))

See Also

Other tune: [TuneControl](#), [getNestedTuneResultsOptPathDf\(\)](#), [getNestedTuneResultsX\(\)](#), [getResamplingIndices\(\)](#), [getTuneResult\(\)](#), [makeModelMultiplexerParamSet\(\)](#), [makeModelMultiplexer\(\)](#), [makeTuneControlCMAES\(\)](#), [makeTuneControlDesign\(\)](#), [makeTuneControlGenSA\(\)](#), [makeTuneControlIrace\(\)](#), [makeTuneControlMBO\(\)](#), [makeTuneControlRandom\(\)](#), [makeTuneWrapper\(\)](#), [tuneParams\(\)](#), [tuneThreshold\(\)](#)

makeTuneControlIrace *Create control object for hyperparameter tuning with Irace.*

Description

Tuning with iterated F-Racing with method `irace::irace`. All kinds of parameter types can be handled. We return the best of the final elite candidates found by irace in the last race. Its estimated performance is the mean of all evaluations ever done for that candidate. More information on irace can be found in package vignette: `vignette("irace-package", package = "irace")`

For resampling you have to pass a `ResampleDesc`, not a `ResampleInstance`. The resampling strategy is randomly instantiated `n.instances` times and these are the instances in the sense of irace (`instances` element of `tunerConfig` in `irace::irace`). Also note that irace will always store its tuning results in a file on disk, see the package documentation for details on this and how to change the file path.

Usage

```
makeTuneControlIrace(
  impute.val = NULL,
  n.instances = 100L,
  show.irace.output = FALSE,
  tune.threshold = FALSE,
  tune.threshold.args = list(),
  log.fun = "default",
  final.dw.perc = NULL,
  budget = NULL,
  ...
)
```

Arguments

<code>impute.val</code>	(numeric) If something goes wrong during optimization (e.g. the learner crashes), this value is fed back to the tuner, so the tuning algorithm does not abort. It is not stored in the optimization path, an NA and a corresponding error message are logged instead. Note that this value is later multiplied by -1 for maximization measures internally, so you need to enter a larger positive value for maximization here as well. Default is the worst obtainable value of the performance measure you optimize for when you aggregate by mean value, or Inf instead. For multi-criteria optimization pass a vector of imputation values, one for each of your measures, in the same order as your measures.
<code>n.instances</code>	(integer(1)) Number of random resampling instances for irace, see details. Default is 100.
<code>show.irace.output</code>	(logical(1)) Show console output of irace while tuning? Default is FALSE.

tune.threshold	(logical(1)) Should the threshold be tuned for the measure at hand, after each hyperparameter evaluation, via tuneThreshold ? Only works for classification if the predict type is “prob”. Default is FALSE.
tune.threshold.args	(list) Further arguments for threshold tuning that are passed down to tuneThreshold . Default is none.
log.fun	(function character(1)) Function used for logging. If set to “default” (the default), the evaluated design points, the resulting performances, and the runtime will be reported. If set to “memory” the memory usage for each evaluation will also be displayed, with character(1) small increase in run time. Otherwise character(1) function with arguments learner, resampling, measures, par.set, control, opt.path, dob, x, y, remove.nas, stage and prev.stage is expected. The default displays the performance measures, the time needed for evaluating, the currently used memory and the max memory ever used before (the latter two both taken from gc). See the implementation for details.
final.dw.perc	(boolean) If a Learner wrapped by a makeDownsampleWrapper is used, you can define the value of dw.perc which is used to train the Learner with the final parameter setting found by the tuning. Default is NULL which will not change anything.
budget	(integer(1)) Maximum budget for tuning. This value restricts the number of function evaluations. It is passed to <code>maxExperiments</code> .
...	(any) Further control parameters passed to the control arguments of cmaes::cma_es or GenSA::GenSA , as well as towards the <code>tunerConfig</code> argument of irace::irace .

Value

([TuneControlIrace](#))

See Also

Other tune: [TuneControl](#), [getNestedTuneResultsOptPathDf\(\)](#), [getNestedTuneResultsX\(\)](#), [getResamplingIndices\(\)](#), [getTuneResult\(\)](#), [makeModelMultiplexerParamSet\(\)](#), [makeModelMultiplexer\(\)](#), [makeTuneControlCMAES\(\)](#), [makeTuneControlDesign\(\)](#), [makeTuneControlGenSA\(\)](#), [makeTuneControlGrid\(\)](#), [makeTuneControlMBO\(\)](#), [makeTuneControlRandom\(\)](#), [makeTuneWrapper\(\)](#), [tuneParams\(\)](#), [tuneThreshold\(\)](#)

makeTuneControlMBO *Create control object for hyperparameter tuning with MBO.*

Description

Model-based / Bayesian optimization with the function [mlrMBO::mbo](#) from the [mlrMBO](#) package. Please refer to <https://github.com/mlr-org/mlrMBO> for further info.

Usage

```

makeTuneControlMBO(
  same.resampling.instance = TRUE,
  impute.val = NULL,
  learner = NULL,
  mbo.control = NULL,
  tune.threshold = FALSE,
  tune.threshold.args = list(),
  continue = FALSE,
  log.fun = "default",
  final.dw.perc = NULL,
  budget = NULL,
  mbo.design = NULL
)

```

Arguments

`same.resampling.instance` (logical(1))
Should the same resampling instance be used for all evaluations to reduce variance? Default is TRUE.

`impute.val` (numeric)
If something goes wrong during optimization (e.g. the learner crashes), this value is fed back to the tuner, so the tuning algorithm does not abort. It is not stored in the optimization path, an NA and a corresponding error message are logged instead. Note that this value is later multiplied by -1 for maximization measures internally, so you need to enter a larger positive value for maximization here as well. Default is the worst obtainable value of the performance measure you optimize for when you aggregate by mean value, or Inf instead. For multi-criteria optimization pass a vector of imputation values, one for each of your measures, in the same order as your measures.

`learner` (Learner | NULL)
The surrogate learner: A regression learner to model performance landscape. For the default, NULL, `mlrMBO` will automatically create a suitable learner based on the rules described in `mlrMBO::makeMBOLEarner`.

`mbo.control` (mlrMBO::MBOControl | NULL)
Control object for model-based optimization tuning. For the default, NULL, the control object will be created with all the defaults as described in `mlrMBO::makeMBOControl`.

`tune.threshold` (logical(1))
Should the threshold be tuned for the measure at hand, after each hyperparameter evaluation, via `tuneThreshold`? Only works for classification if the predict type is "prob". Default is FALSE.

`tune.threshold.args` (list)
Further arguments for threshold tuning that are passed down to `tuneThreshold`. Default is none.

continue	(logical(1)) Resume calculation from previous run using <code>mlrMBO::mboContinue?</code> Requires “save.file.path” to be set. Note that the <code>ParamHelpers::OptPath</code> in the <code>mlrMBO::OptResult</code> will only include the evaluations after the continuation. The complete <code>OptPath</code> will be found in the slot <code>\$mbo.result\$opt.path</code> .
log.fun	(function character(1)) Function used for logging. If set to “default” (the default), the evaluated design points, the resulting performances, and the runtime will be reported. If set to “memory” the memory usage for each evaluation will also be displayed, with character(1) small increase in run time. Otherwise character(1) function with arguments <code>learner</code> , <code>resampling</code> , <code>measures</code> , <code>par.set</code> , <code>control</code> , <code>opt.path</code> , <code>dob</code> , <code>x</code> , <code>y</code> , <code>remove.nas</code> , <code>stage</code> and <code>prev.stage</code> is expected. The default displays the performance measures, the time needed for evaluating, the currently used memory and the max memory ever used before (the latter two both taken from <code>gc</code>). See the implementation for details.
final.dw.perc	(boolean) If a Learner wrapped by a <code>makeDownsampleWrapper</code> is used, you can define the value of <code>dw.perc</code> which is used to train the Learner with the final parameter setting found by the tuning. Default is NULL which will not change anything.
budget	(integer(1)) Maximum budget for tuning. This value restricts the number of function evaluations.
mbo.design	(data.frame NULL) Initial design as data frame. If the parameters have corresponding trafo functions, the design must not be transformed before it is passed! For the default, NULL, a default design is created like described in <code>mlrMBO::mbo</code> .

Value`(TuneControlMBO)`**References**

Bernd Bischl, Jakob Richter, Jakob Bossek, Daniel Horn, Janek Thomas and Michel Lang; mlrMBO: A Modular Framework for Model-Based Optimization of Expensive Black-Box Functions, Preprint: <https://arxiv.org/abs/1703.03373> (2017).

See Also

Other tune: `TuneControl`, `getNestedTuneResultsOptPathDf()`, `getNestedTuneResultsX()`, `getResamplingIndices()`, `getTuneResult()`, `makeModelMultiplexerParamSet()`, `makeModelMultiplexer()`, `makeTuneControlCMAES()`, `makeTuneControlDesign()`, `makeTuneControlGenSA()`, `makeTuneControlGrid()`, `makeTuneControlIrace()`, `makeTuneControlRandom()`, `makeTuneWrapper()`, `tuneParams()`, `tuneThreshold()`

makeTuneControlRandom *Create control object for hyperparameter tuning with random search.*

Description

Random search. All kinds of parameter types can be handled.

Usage

```
makeTuneControlRandom(
  same.resampling.instance = TRUE,
  maxit = NULL,
  tune.threshold = FALSE,
  tune.threshold.args = list(),
  log.fun = "default",
  final.dw.perc = NULL,
  budget = NULL
)
```

Arguments

same.resampling.instance	(logical(1)) Should the same resampling instance be used for all evaluations to reduce variance? Default is TRUE.
maxit	(integer(1) NULL) Number of iterations for random search. Default is 100.
tune.threshold	(logical(1)) Should the threshold be tuned for the measure at hand, after each hyperparameter evaluation, via tuneThreshold ? Only works for classification if the predict type is "prob". Default is FALSE.
tune.threshold.args	(list) Further arguments for threshold tuning that are passed down to tuneThreshold . Default is none.
log.fun	(function character(1)) Function used for logging. If set to "default" (the default), the evaluated design points, the resulting performances, and the runtime will be reported. If set to "memory" the memory usage for each evaluation will also be displayed, with character(1) small increase in run time. Otherwise character(1) function with arguments learner, resampling, measures, par.set, control, opt.path, dob, x, y, remove.nas, stage and prev.stage is expected. The default displays the performance measures, the time needed for evaluating, the currently used memory and the max memory ever used before (the latter two both taken from gc). See the implementation for details.

final.dw.perc	(boolean) If a Learner wrapped by a makeDownsampleWrapper is used, you can define the value of dw.perc which is used to train the Learner with the final parameter setting found by the tuning. Default is NULL which will not change anything.
budget	(integer(1)) Maximum budget for tuning. This value restricts the number of function evaluations. The budget equals the number of iterations (maxit) performed by the random search algorithm.

Value[\(TuneControlRandom\)](#)**See Also**

Other tune: [TuneControl](#), [getNestedTuneResultsOptPathDf\(\)](#), [getNestedTuneResultsX\(\)](#), [getResamplingIndices\(\)](#), [getTuneResult\(\)](#), [makeModelMultiplexerParamSet\(\)](#), [makeModelMultiplexer\(\)](#), [makeTuneControlCMAES\(\)](#), [makeTuneControlDesign\(\)](#), [makeTuneControlGenSA\(\)](#), [makeTuneControlGrid\(\)](#), [makeTuneControlIrace\(\)](#), [makeTuneControlMBO\(\)](#), [makeTuneWrapper\(\)](#), [tuneParams\(\)](#), [tuneThreshold\(\)](#)

makeTuneWrapper	<i>Fuse learner with tuning.</i>
-----------------	----------------------------------

Description

Fuses a base learner with a search strategy to select its hyperparameters. Creates a learner object, which can be used like any other learner object, but which internally uses [tuneParams](#). If the train function is called on it, the search strategy and resampling are invoked to select an optimal set of hyperparameter values. Finally, a model is fitted on the complete training data with these optimal hyperparameters and returned. See [tuneParams](#) for more details.

After training, the optimal hyperparameters (and other related information) can be retrieved with [getTuneResult](#).

Usage

```
makeTuneWrapper(
  learner,
  resampling,
  measures,
  par.set,
  control,
  show.info = getMlrOption("show.info")
)
```

Arguments

learner	(Learner character(1)) The learner. If you pass a string the learner will be created via makeLearner .
resampling	(ResampleInstance ResampleDesc) Resampling strategy to evaluate points in hyperparameter space. If you pass a description, it is instantiated once at the beginning by default, so all points are evaluated on the same training/test sets. If you want to change that behavior, look at TuneControl .
measures	(list of Measure Measure) Performance measures to evaluate. The first measure, aggregated by the first aggregation function is optimized, others are simply evaluated. Default is the default measure for the task, see here getDefaultMeasure .
par.set	(ParamHelpers::ParamSet) Collection of parameters and their constraints for optimization. Dependent parameters with a requires field must use quote and not expression to define it.
control	(TuneControl) Control object for search method. Also selects the optimization algorithm for tuning.
show.info	(logical(1)) Print verbose output on console? Default is set via configureMlr .

Value

[Learner](#).

See Also

Other tune: [TuneControl](#), [getNestedTuneResultsOptPathDf\(\)](#), [getNestedTuneResultsX\(\)](#), [getResamplingIndices\(\)](#), [getTuneResult\(\)](#), [makeModelMultiplexerParamSet\(\)](#), [makeModelMultiplexer\(\)](#), [makeTuneControlCMAES\(\)](#), [makeTuneControlDesign\(\)](#), [makeTuneControlGenSA\(\)](#), [makeTuneControlGrid\(\)](#), [makeTuneControlIrace\(\)](#), [makeTuneControlMBO\(\)](#), [makeTuneControlRandom\(\)](#), [tuneParams\(\)](#), [tuneThreshold\(\)](#)

Other wrapper: [makeBaggingWrapper\(\)](#), [makeClassificationViaRegressionWrapper\(\)](#), [makeConstantClassWrapper\(\)](#), [makeCostSensClassifWrapper\(\)](#), [makeCostSensRegrWrapper\(\)](#), [makeDownsampleWrapper\(\)](#), [makeDummyFeaturesWrapper\(\)](#), [makeExtractFDAFeatsWrapper\(\)](#), [makeFeatSelWrapper\(\)](#), [makeFilterWrapper\(\)](#), [makeImputeWrapper\(\)](#), [makeMulticlassWrapper\(\)](#), [makeMultilabelBinaryRelevanceWrapper\(\)](#), [makeMultilabelClassifierChainsWrapper\(\)](#), [makeMultilabelDBRWrapper\(\)](#), [makeMultilabelNestedStackingWrapper\(\)](#), [makeMultilabelStackingWrapper\(\)](#), [makeOverBaggingWrapper\(\)](#), [makePreprocWrapperCaret\(\)](#), [makePreprocWrapper\(\)](#), [makeRemoveConstantFeaturesWrapper\(\)](#), [makeSMOTERWrapper\(\)](#), [makeUndersampleWrapper\(\)](#), [makeWeightedClassesWrapper\(\)](#)

Examples

```
task = makeClassifTask(data = iris, target = "Species")
lrn = makeLearner("classif.rpart")
```

```

# stupid mini grid
ps = makeParamSet(
  makeDiscreteParam("cp", values = c(0.05, 0.1)),
  makeDiscreteParam("minsplit", values = c(10, 20))
)
ctrl = makeTuneControlGrid()
inner = makeResampleDesc("Holdout")
outer = makeResampleDesc("CV", iters = 2)
lrn = makeTuneWrapper(lrn, resampling = inner, par.set = ps, control = ctrl)
mod = train(lrn, task)
print(getTuneResult(mod))
# nested resampling for evaluation
# we also extract tuned hyper pars in each iteration
r = resample(lrn, task, outer, extract = getTuneResult)
print(r$extract)
getNestedTuneResultsOptPathDf(r)
getNestedTuneResultsX(r)

```

makeUndersampleWrapper

Fuse learner with simple ove/underrsampling for imbalance correction in binary classification.

Description

Creates a learner object, which can be used like any other learner object. Internally uses [oversample](#) or [undersample](#) before every model fit.

Note that observation weights do not influence the sampling and are simply passed down to the next learner.

Usage

```
makeUndersampleWrapper(learner, usw.rate = 1, usw.cl = NULL)
```

```
makeOversampleWrapper(learner, osw.rate = 1, osw.cl = NULL)
```

Arguments

learner	(Learner character(1)) The learner. If you pass a string the learner will be created via makeLearner .
usw.rate	(numeric(1)) Factor to downsample a class. Must be between 0 and 1, where 1 means no downsampling, 0.5 implies reduction to 50 percent and 0 would imply reduction to 0 observations. Default is 1.
usw.cl	(character(1)) Class that should be undersampled. Default is NULL, which means the larger one.

osw.rate	(numeric(1)) Factor to oversample a class. Must be between 1 and Inf, where 1 means no oversampling and 2 would mean doubling the class size. Default is 1.
osw.cl	(character(1)) Class that should be oversampled. Default is NULL, which means the smaller one.

Value

[Learner](#).

See Also

Other imbalancecy: [makeOverBaggingWrapper\(\)](#), [oversample\(\)](#), [smote\(\)](#)

Other wrapper: [makeBaggingWrapper\(\)](#), [makeClassificationViaRegressionWrapper\(\)](#), [makeConstantClassWrapper\(\)](#), [makeCostSensClassifWrapper\(\)](#), [makeCostSensRegrWrapper\(\)](#), [makeDownsampleWrapper\(\)](#), [makeDummyFeaturesWrapper\(\)](#), [makeExtractFDAFeatsWrapper\(\)](#), [makeFeatSelWrapper\(\)](#), [makeFilterWrapper\(\)](#), [makeImputeWrapper\(\)](#), [makeMulticlassWrapper\(\)](#), [makeMultilabelBinaryRelevanceWrapper\(\)](#), [makeMultilabelClassifierChainsWrapper\(\)](#), [makeMultilabelDBRWrapper\(\)](#), [makeMultilabelNestedStackingWrapper\(\)](#), [makeMultilabelStackingWrapper\(\)](#), [makeOverBaggingWrapper\(\)](#), [makePreprocWrapperCaret\(\)](#), [makePreprocWrapper\(\)](#), [makeRemoveConstantFeaturesWrapper\(\)](#), [makeSMOTEWrapper\(\)](#), [makeTuneWrapper\(\)](#), [makeWeightedClassesWrapper\(\)](#)

makeWeightedClassesWrapper

Wraps a classifier for weighted fitting where each class receives a weight.

Description

Creates a wrapper, which can be used like any other learner object.

Fitting is performed in a weighted fashion where each observation receives a weight, depending on the class it belongs to, see `wcw.weight`. This might help to mitigate problems caused by imbalanced class distributions.

This weighted fitting can be achieved in two ways:

a) The learner already has a parameter for class weighting, so one weight can directly be defined per class. Example: “`classif.ksvm`” and parameter `class.weights`. In this case we don’t really do anything fancy. We convert `wcw.weight` a bit, but basically simply bind its value to the class weighting param. The wrapper in this case simply offers a convenient, consistent fashion for class weighting - and tuning! See example below.

b) The learner does not have a direct parameter to support class weighting, but supports observation weights, so `hasLearnerProperties(learner, 'weights')` is TRUE. This means that an individual, arbitrary weight can be set per observation during training. We set this weight depending on the class internally in the wrapper. Basically we introduce something like a new “`class.weights`” parameter for the learner via observation weights.

Usage

```
makeWeightedClassesWrapper(learner, wcw.param = NULL, wcw.weight = 1)
```

Arguments

learner	(Learner character(1)) The classification learner. If you pass a string the learner will be created via makeLearner .
wcw.param	(character(1)) Name of already existing learner parameter, which allows class weighting. The default (wcw.param = NULL) will use the parameter defined in the learner (class.weights.param). During training, the parameter must accept a named vector of class weights, where length equals the number of classes.
wcw.weight	(numeric) Weight for each class. Must be a vector of the same number of elements as classes are in task, and must also be in the same order as the class levels are in <code>getTaskDesc(task)\$class.levels</code> . For convenience, one must pass a single number in case of binary classification, which is then taken as the weight of the positive class, while the negative class receives a weight of 1. Default is 1.

Value

[Learner](#).

See Also

Other wrapper: [makeBaggingWrapper\(\)](#), [makeClassificationViaRegressionWrapper\(\)](#), [makeConstantClassWrapper\(\)](#), [makeCostSensClassifWrapper\(\)](#), [makeCostSensRegrWrapper\(\)](#), [makeDownsampleWrapper\(\)](#), [makeDummyFeaturesWrapper\(\)](#), [makeExtractFDAFeatsWrapper\(\)](#), [makeFeatSelWrapper\(\)](#), [makeFilterWrapper\(\)](#), [makeImputeWrapper\(\)](#), [makeMulticlassWrapper\(\)](#), [makeMultilabelBinaryRelevanceWrapper\(\)](#), [makeMultilabelClassifierChainsWrapper\(\)](#), [makeMultilabelDBRWrapper\(\)](#), [makeMultilabelNestedStackingWrapper\(\)](#), [makeMultilabelStackingWrapper\(\)](#), [makeOverBaggingWrapper\(\)](#), [makePreprocWrapperCaret\(\)](#), [makePreprocWrapper\(\)](#), [makeRemoveConstantFeaturesWrapper\(\)](#), [makeSMOTEWrapper\(\)](#), [makeTuneWrapper\(\)](#), [makeUndersampleWrapper\(\)](#)

Examples

```
set.seed(123)
# using the direct parameter of the SVM (which is already defined in the learner)
lrn = makeWeightedClassesWrapper("classif.ksvm", wcw.weight = 0.01)
res = holdout(lrn, sonar.task)
print(calculateConfusionMatrix(res$pred))

# using the observation weights of logreg
lrn = makeWeightedClassesWrapper("classif.logreg", wcw.weight = 0.01)
res = holdout(lrn, sonar.task)
print(calculateConfusionMatrix(res$pred))
```



```

# tuning the imbalance param and the SVM param in one go
lrn = makeWeightedClassesWrapper("classif.ksvm", wcw.param = "class.weights")
ps = makeParamSet(
  makeNumericParam("wcv.weight", lower = 1, upper = 10),
  makeNumericParam("C", lower = -12, upper = 12, trafo = function(x) 2^x),
  makeNumericParam("sigma", lower = -12, upper = 12, trafo = function(x) 2^x)
)
ctrl = makeTuneControlRandom(maxit = 3L)
rdesc = makeResampleDesc("CV", iters = 2L, stratify = TRUE)
res = tuneParams(lrn, sonar.task, rdesc, par.set = ps, control = ctrl)
print(res)
# print(res$opt.path)

```

makeWrappedModel	<i>Induced model of learner.</i>
------------------	----------------------------------

Description

Result from [train](#).

It internally stores the underlying fitted model, the subset used for training, features used for training, levels of factors in the data set and computation time that was spent for training.

Object members: See arguments.

The constructor `makeWrappedModel` is mainly for internal use.

Usage

```

makeWrappedModel(
  learner,
  learner.model,
  task.desc,
  subset,
  features,
  factor.levels,
  time
)

```

Arguments

learner	(Learner character(1)) The learner. If you pass a string the learner will be created via makeLearner .
learner.model	(any) Underlying model.
task.desc	TaskDesc Task description object.

subset	(integer logical NULL) Selected cases. Either a logical or an index vector. By default NULL if all observations are used.
features	(character) Features used for training.
factor.levels	(named list of character) Levels of factor variables (features and potentially target) in training data. Named by variable name, non-factors do not occur in the list.
time	(numeric(1)) Computation time for model fit in seconds.

Value

[WrappedModel](#).

MeasureProperties	<i>Query properties of measures.</i>
-------------------	--------------------------------------

Description

Properties can be accessed with `getMeasureProperties(measure)`, which returns a character vector.

The measure properties are defined in [Measure](#).

Usage

```
getMeasureProperties(measure)
```

```
hasMeasureProperties(measure, props)
```

Arguments

measure [\(Measure\)](#)
Performance measure. Default is the first measure used in the benchmark experiment.

props [\(character\)](#)
Vector of properties to query.

Value

`getMeasureProperties` returns a character vector with measure properties. `hasMeasureProperties` returns a logical vector of the same length as `props`.

measures

Performance measures.

Description

A performance measure is evaluated after a single train/predict step and returns a single number to assess the quality of the prediction (or maybe only the model, think AIC). The measure itself knows whether it wants to be minimized or maximized and for what tasks it is applicable.

All supported measures can be found by [listMeasures](#) or as a table in the tutorial appendix: <https://mlr.mlr-org.com/articles/tutorial/measures.html>.

If you want a measure for a misclassification cost matrix, look at [makeCostMeasure](#). If you want to implement your own measure, look at [makeMeasure](#).

Most measures can directly be accessed via the function named after the scheme measureX (e.g. [measureSSE](#)).

For clustering measures, we compact the predicted cluster IDs such that they form a continuous series starting with 1. If this is not the case, some of the measures will generate warnings.

Some measure have parameters. Their defaults are set in the constructor [makeMeasure](#) and can be overwritten using [setMeasurePars](#).

Usage

`measureSSE(truth, response)`

`measureMSE(truth, response)`

`measureRMSE(truth, response)`

`measureMEDSE(truth, response)`

`measureSAE(truth, response)`

`measureMAE(truth, response)`

`measureMEDAE(truth, response)`

`measureRSQ(truth, response)`

`measureEXPVAR(truth, response)`

`measureRRSE(truth, response)`

`measureRAE(truth, response)`

`measureMAPE(truth, response)`

measureMSLE(truth, response)
measureRMSLE(truth, response)
measureKendallTau(truth, response)
measureSpearmanRho(truth, response)
measureMMCE(truth, response)
measureACC(truth, response)
measureBER(truth, response)
measureAUNU(probabilities, truth)
measureAUNP(probabilities, truth)
measureAU1U(probabilities, truth)
measureAU1P(probabilities, truth)
measureMulticlassBrier(probabilities, truth)
measureLogloss(probabilities, truth)
measureSSR(probabilities, truth)
measureQSR(probabilities, truth)
measureLSR(probabilities, truth)
measureKAPPA(truth, response)
measureWKAPPA(truth, response)
measureAUC(probabilities, truth, negative, positive)
measureBrier(probabilities, truth, negative, positive)
measureBrierScaled(probabilities, truth, negative, positive)
measureBAC(truth, response)
measureTP(truth, response, positive)
measureTN(truth, response, negative)

```
measureFP(truth, response, positive)
measureFN(truth, response, negative)
measureTPR(truth, response, positive)
measureTNR(truth, response, negative)
measureFPR(truth, response, negative, positive)
measureFNR(truth, response, negative, positive)
measurePPV(truth, response, positive, probabilities = NULL)
measureNPV(truth, response, negative)
measureFDR(truth, response, positive)
measureMCC(truth, response, negative, positive)
measureF1(truth, response, positive)
measureGMEAN(truth, response, negative, positive)
measureGPR(truth, response, positive)
measureMultilabelHamloss(truth, response)
measureMultilabelSubset01(truth, response)
measureMultilabelF1(truth, response)
measureMultilabelACC(truth, response)
measureMultilabelPPV(truth, response)
measureMultilabelTPR(truth, response)
```

Arguments

truth	(factor) Vector of the true class.
response	(factor) Vector of the predicted class.
probabilities	(numeric matrix) a) For purely binary classification measures: The predicted probabilities for the positive class as a numeric vector. b) For multiclass classification measures: The predicted probabilities for all classes, always as a numeric matrix, where

	columns are named with class labels.
negative	(character(1)) The name of the negative class.
positive	(character(1)) The name of the positive class.

References

He, H. & Garcia, E. A. (2009) *Learning from Imbalanced Data*. IEEE Transactions on Knowledge and Data Engineering, vol. 21, no. 9. pp. 1263-1284.

H. Uno et al. *On the C-statistics for Evaluating Overall Adequacy of Risk Prediction Procedures with Censored Survival Data* Statistics in medicine. 2011;30(10):1105-1117. <https://doi.org/10.1002/sim.4154>.

H. Uno et al. *Evaluating Prediction Rules for T-Year Survivors with Censored Regression Models* Journal of the American Statistical Association 102, no. 478 (2007): 527-37.

See Also

Other performance: [ConfusionMatrix](#), [calculateConfusionMatrix\(\)](#), [calculateROCMeasures\(\)](#), [estimateRelativeOverfitting\(\)](#), [makeCostMeasure\(\)](#), [makeCustomResampledMeasure\(\)](#), [makeMeasure\(\)](#), [performance\(\)](#), [setAggregation\(\)](#), [setMeasurePars\(\)](#)

mergeBenchmarkResults *Merge different BenchmarkResult objects.*

Description

The function automatically combines a list of [BenchmarkResult](#) objects into a single [BenchmarkResult](#) object as long as the full crossproduct of all task-learner combinations are available.

Usage

```
mergeBenchmarkResults(bmrs)
```

Arguments

bmrs (list of [BenchmarkResult](#))
BenchmarkResult objects that should be merged.

Details

Note that if you want to merge several [BenchmarkResult](#) objects, you must ensure that all possible learner and task combinations will be contained in the returned object. Otherwise, the user will be notified which task-learner combinations are missing or duplicated.

When merging [BenchmarkResult](#) objects with different measures, all missing measures will automatically be recomputed.

Value

[BenchmarkResult](#)

mergeSmallFactorLevels

Merges small levels of factors into new level.

Description

Merges factor levels that occur only infrequently into combined levels with a higher frequency.

Usage

```
mergeSmallFactorLevels(  
  task,  
  cols = NULL,  
  min.perc = 0.01,  
  new.level = ".merged"  
)
```

Arguments

task	(Task) The task.
cols	(character) Which columns to convert. Default is all factor and character columns.
min.perc	(numeric(1)) The smallest levels of a factor are merged until their combined proportion w.r.t. the length of the factor exceeds min.perc. Must be between 0 and 1. Default is 0.01.
new.level	(character(1)) New name of merged level. Default is “.merged”

Value

Task, where merged levels are combined into a new level of name new.level.

See Also

Other eda_and_preprocess: [capLargeValues\(\)](#), [createDummyFeatures\(\)](#), [dropFeatures\(\)](#), [normalizeFeatures\(\)](#), [removeConstantFeatures\(\)](#), [summarizeColumns\(\)](#), [summarizeLevels\(\)](#)

mlrFamilies

*mlr documentation families***Description**

List of all mlr documentation families with members.

Arguments

benchmark	batchmark, reduceBatchmarkResults, benchmark, benchmarkParallel, getBMRTaskIds, getBMRLearners, getBMRLearnerIds, getBMRLearnerShortNames, getBMRMeasures, getBMRMeasureIds, getBMRPredictions, getBMRPerformances, getBMRAggrPerformances, getBMRTuneResults, getBMRFeatSelResults, getBMRFilteredFeatures, getBMRModels, getBMRTaskDescs, convertBMRTToRankMatrix, friedmanPostHocTestBMR, friedmanTestBMR, plotBMRBoxplots, plotBMRRanksAsBarChart, generateCritDifferencesData, plotCritDifferences
calibration	generateCalibrationData, plotCalibration
configure	configureMlr, getMlrOptions
costsens	makeCostSensTask, makeCostSensWeightedPairsWrapper
debug	predictFailureModel, getPredictionDump, getRRDump, print.ResampleResult
downsample	downsample
eda_and_preprocess	capLargeValues, createDummyFeatures, dropFeatures, mergeSmallFactorLevels, normalizeFeatures, removeConstantFeatures, summarizeColumns, summarizeLevels
extractFDAFeatures	reextractFDAFeatures
fda_featextractor	extractFDAFourier, extractFDAWavelets, extractFDAFPCA, extractFDAMultiResFeatures
fda	makeExtractFDAFeatMethod, extractFDAFeatures
featsel	analyzeFeatSelResult, makeFeatSelControl, getFeatSelResult, selectFeatures
filter	filterFeatures, makeFilter, listFilterMethods, getFilteredFeatures, generateFilterValuesData, getFilterValues
generate_plot_data	generateFeatureImportanceData, plotFilterValues, generatePartialDependenceData
help	helpLearner, helpLearnerParam
imbalance	oversample, smote
impute	makeImputeMethod, imputeConstant, impute, reimpute

learner	getClassWeightParam, getHyperPars, getParamSet.Learner, getLearnerType, getLearnerId, getLearnerPredictType, getLearnerPackages, getLearnerParamSet, getLearnerParVals, setLearnerId, getLearnerShortName, getLearnerProperties, makeLearner, makeLearners, removeHyperPars, setHyperPars, setId, setPredictThreshold, setPredictType
learning_curve	generateLearningCurveData
multilabel	getMultilabelBinaryPerformances, makeMultilabelBinaryRelevanceWrapper, makeMultilabelClassifierChainsWrapper, makeMultilabelDBRWrapper, makeMultilabelNestedStackingWrapper, makeMultilabelStackingWrapper
performance	calculateConfusionMatrix, calculateROCMeasures, makeCustomResampledMeasure, makeCostMeasure, setMeasurePars, setAggregation, makeMeasure, featperc, performance, estimateRelativeOverfitting
plot	createSpatialResamplingPlots, plotLearningCurve, plotPartialDependence, plotBMR-Summary, plotResiduals
predict	asROCRPrediction, getPredictionProbabilities, getPredictionTaskDesc, getPredictionResponse, predict.WrappedModel
resample	makeResampleDesc, makeResampleInstance, makeResamplePrediction, resample, getRRPredictions, getRRTaskDescription, getRRTaskDesc, getRRPredictionList, addRRMeasure
task	getTaskDesc, getTaskType, getTaskId, getTaskTargetNames, getTaskClassLevels, getTaskFeatureNames, getTaskNFeats, getTaskSize, getTaskFormula, getTaskTargets, getTaskData, getTaskCosts, subsetTask
thresh_vs_perf	generateThreshVsPerfData, plotThreshVsPerf, plotROCCurves
tune	getNestedTuneResultsX, getNestedTuneResultsOptPathDf, getResamplingIndices, getTuneResult, makeModelMultiplexerParamSet, makeModelMultiplexer, makeTuneControlCMAES, makeTuneControlDesign, makeTuneControlGenSA, makeTuneControlGrid, makeTuneControlIrace, makeTuneControlMBO, makeTuneControl, makeTuneControlRandom, tuneParams, tuneThreshold
tune_multicrit	plotTuneMultiCritResult, makeTuneMultiCritControl, tuneParamsMultiCrit
wrapper	makeBaggingWrapper, makeClassificationViaRegressionWrapper, makeConstantClassWrapper, makeCostSensClassifWrapper, makeCostSensRegrWrapper, makeDownsampleWrapper, makeDummyFeaturesWrapper, makeExtractFDAFeatsWrapper, makeFeatSelWrapper, makeFilterWrapper, makeImputeWrapper, makeMulticlassWrapper, makeOverBaggingWrapper, makeUndersampleWrapper, makePreprocWrapperCaret, makePreprocWrapper, makeRemoveConstantFeaturesWrapper, makeSMOTEWrapper, makeTuneWrapper, makeWeightedClassesWrapper

mtcars.task

Motor Trend Car Road Tests clustering task.

Description

Contains the task (mtcars.task).

References

See [datasets::mtcars](#).

normalizeFeatures *Normalize features.*

Description

Normalize features by different methods. Internally `BBmisc::normalize` is used for every feature column. Non numerical features will be left untouched and passed to the result. For constant features most methods fail, special behaviour for this case is implemented.

Usage

```
normalizeFeatures(
  obj,
  target = character(0L),
  method = "standardize",
  cols = NULL,
  range = c(0, 1),
  on.constant = "quiet"
)
```

Arguments

obj	(data.frame Task) Input data.
target	(character(1) character(2) character(n.classes)) Name(s) of the target variable(s). Only used when obj is a data.frame, otherwise ignored. If survival analysis is applicable, these are the names of the survival time and event columns, so it has length 2. For multilabel classification these are the names of logical columns that indicate whether a class label is present and the number of target variables corresponds to the number of classes.
method	(character(1)) Normalizing method. Available are: “center”: Subtract mean. “scale”: Divide by standard deviation. “standardize”: Center and scale. “range”: Scale to a given range.
cols	(character) Columns to normalize. Default is to use all numeric columns.
range	(numeric(2)) Range for method “range”. Default is <code>c(0, 1)</code> .

`on.constant` (character(1))
 How should constant vectors be treated? Only used, of “method != center”, since this methods does not fail for constant vectors. Possible actions are:
 “quiet”: Depending on the method, treat them quietly:
 “scale”: No division by standard deviation is done, input values. will be returned untouched.
 “standardize”: Only the mean is subtracted, no division is done.
 “range”: All values are mapped to the mean of the given range.
 “warn”: Same behaviour as “quiet”, but print a warning message.
 “stop”: Stop with an error.

Value

`data.frame` | `Task`. Same type as `obj`.

See Also

`BBmisc::normalize`

Other `eda_and_preprocess`: `capLargeValues()`, `createDummyFeatures()`, `dropFeatures()`, `mergeSmallFactorLevels()`, `removeConstantFeatures()`, `summarizeColumns()`, `summarizeLevels()`

<code>oversample</code>	<i>Over- or undersample binary classification task to handle class imbalance.</i>
-------------------------	---

Description

Oversampling: For a given class (usually the smaller one) all existing observations are taken and copied and extra observations are added by randomly sampling with replacement from this class.

Undersampling: For a given class (usually the larger one) the number of observations is reduced (downsampled) by randomly sampling without replacement from this class.

Usage

```
oversample(task, rate, cl = NULL)
```

```
undersample(task, rate, cl = NULL)
```

Arguments

`task` (Task)
 The task.

`rate` (numeric(1))
 Factor to upsample or downsample a class. For undersampling: Must be between 0 and 1, where 1 means no downsampling, 0.5 implies reduction to 50 percent and 0 would imply reduction to 0 observations. For oversampling: Must

be between 1 and Inf, where 1 means no oversampling and 2 would mean doubling the class size.

`cl` (character(1))
Which class should be over- or undersampled. If NULL, `oversample` will select the smaller and `undersample` the larger class.

Value

[Task](#).

See Also

Other imbalancy: [makeOverBaggingWrapper\(\)](#), [makeUndersampleWrapper\(\)](#), [smote\(\)](#)

parallelization

Supported parallelization methods

Description

`mlr` supports different methods to activate parallel computing capabilities through the integration of the [parallelMap::parallelMap](#) package, which supports all major parallelization backends for R. You can start parallelization with `parallelStart*`, where `*` should be replaced with the chosen backend. [parallelMap::parallelStop](#) is used to stop all parallelization backends.

Parallelization is divided into different levels and will automatically be carried out for the first level that occurs, e.g. if you call `resample()` after [parallelMap::parallelStart](#), each resampling iteration is a parallel job and possible underlying calls like parameter tuning won't be parallelized further.

The supported levels of parallelization are:

"`mlr.resample`" Each resampling iteration (a train/test step) is a parallel job.

"`mlr.benchmark`" Each experiment "run this learner on this data set" is a parallel job.

"`mlr.tuneParams`" Each evaluation in hyperparameter space "resample with these parameter settings" is a parallel job. How many of these can be run independently in parallel depends on the tuning algorithm. For grid search or random search there is no limit, but for other tuners it depends on how many points to evaluate are produced in each iteration of the optimization. If a tuner works in a purely sequential fashion, we cannot work magic and the hyperparameter evaluation will also run sequentially. But note that you can still parallelize the underlying resampling.

"`mlr.selectFeatures`" Each evaluation in feature space "resample with this feature subset" is a parallel job. The same comments as for "`mlr.tuneParams`" apply here.

"`mlr.ensemble`" For all ensemble methods, the training and prediction of each individual learner is a parallel job. Supported ensemble methods are the [makeBaggingWrapper](#), [makeCostSensitiveRegrWrapper](#), [makeMulticlassWrapper](#), [makeMultilabelBinaryRelevanceWrapper](#) and the [makeOverBaggingWrapper](#).

performance	<i>Measure performance of prediction.</i>
-------------	---

Description

Measures the quality of a prediction w.r.t. some performance measure.

Usage

```
performance(  
  pred,  
  measures,  
  task = NULL,  
  model = NULL,  
  feats = NULL,  
  simpleaggr = FALSE  
)
```

Arguments

pred	(Prediction) Prediction object.
measures	(Measure list of Measure) Performance measure(s) to evaluate. Default is the default measure for the task, see here getDefaultMeasure .
task	(Task) Learning task, might be requested by performance measure, usually not needed except for clustering or survival.
model	(WrappedModel) Model built on training data, might be requested by performance measure, usually not needed except for survival.
feats	(data.frame) Features of predicted data, usually not needed except for clustering. If the prediction was generated from a task, you can also pass this instead and the features are extracted from it.
simpleaggr	(logical) If TRUE, aggregation of ResamplePrediction objects is skipped. This is used internally for threshold tuning. Default is FALSE.

Value

(named [numeric](#)). Performance value(s), named by measure(s).

See Also

Other performance: [ConfusionMatrix](#), [calculateConfusionMatrix\(\)](#), [calculateROCMeasures\(\)](#), [estimateRelativeOverfitting\(\)](#), [makeCostMeasure\(\)](#), [makeCustomResampledMeasure\(\)](#), [makeMeasure\(\)](#), [measures](#), [setAggregation\(\)](#), [setMeasurePars\(\)](#)

Examples

```
training.set = seq(1, nrow(iris), by = 2)
test.set = seq(2, nrow(iris), by = 2)

task = makeClassifTask(data = iris, target = "Species")
lrn = makeLearner("classif.lda")
mod = train(lrn, task, subset = training.set)
pred = predict(mod, newdata = iris[test.set, ])
performance(pred, measures = mmce)

# Compute multiple performance measures at once
ms = list("mmce" = mmce, "acc" = acc, "timetrain" = timetrain)
performance(pred, measures = ms, task, mod)
```

phoneme.task

Phoneme functional data multilabel classification task.

Description

Contains the task (phoneme.task). The task contains a single functional covariate and 5 equally big classes (aa, ao, dcl, iy, sh). The aim is to predict the class of the phoneme in the functional. The dataset is contained in the package fda.usc.

References

F. Ferraty and P. Vieu (2003) "Curve discrimination: a nonparametric functional approach", *Computational Statistics and Data Analysis*, 44(1-2), 161-173. F. Ferraty and P. Vieu (2006) *Nonparametric functional data analysis*, New York: Springer. T. Hastie and R. Tibshirani and J. Friedman (2009) *The elements of statistical learning: Data mining, inference and prediction*, 2nd edn, New York: Springer.

pid.task

PimaIndiansDiabetes classification task.

Description

Contains the task (pid.task).

References

See [mlbench::PimaIndiansDiabetes](#). Note that this is the uncorrected version from mlbench.

plotBMRBoxplots *Create box or violin plots for a BenchmarkResult.*

Description

Plots box or violin plots for a selected measure across all iterations of the resampling strategy, faceted by the task.id.

Usage

```
plotBMRBoxplots(
  bmr,
  measure = NULL,
  style = "box",
  order.lrns = NULL,
  order.tsks = NULL,
  pretty.names = TRUE,
  facet.wrap.nrow = NULL,
  facet.wrap.ncol = NULL
)
```

Arguments

bmr	(BenchmarkResult) Benchmark result.
measure	(Measure) Performance measure. Default is the first measure used in the benchmark experiment.
style	(character(1)) Type of plot, can be “box” for a boxplot or “violin” for a violin plot. Default is “box”.
order.lrns	(character(n.learners)) Character vector with learner.ids in new order.
order.tsks	(character(n.tasks)) Character vector with task.ids in new order.
pretty.names	(logical(1)) Whether to use the Measure name and the Learner short name instead of the id. Default is TRUE.
facet.wrap.nrow, facet.wrap.ncol	(integer) Number of rows and columns for faceting. Default for both is NULL. In this case ggplot’s facet_wrap will choose the layout itself.

Value

ggplot2 plot object.

See Also

Other plot: `createSpatialResamplingPlots()`, `plotBMRRanksAsBarChart()`, `plotBMRSummary()`, `plotCalibration()`, `plotCritDifferences()`, `plotLearningCurve()`, `plotPartialDependence()`, `plotROCCurves()`, `plotResiduals()`, `plotThreshVsPerf()`

Other benchmark: `BenchmarkResult`, `batchmark()`, `benchmark()`, `convertBMRTToRankMatrix()`, `friedmanPostHocTestBMR()`, `friedmanTestBMR()`, `generateCritDifferencesData()`, `getBMRAggrPerformances()`, `getBMRFeatSelResults()`, `getBMRFilteredFeatures()`, `getBMRLearnerIds()`, `getBMRLearnerShortNames()`, `getBMRLearners()`, `getBMRMeasureIds()`, `getBMRMeasures()`, `getBMRModels()`, `getBMRPerformances()`, `getBMRPredictions()`, `getBMRTaskDescs()`, `getBMRTaskIds()`, `getBMRTuneResults()`, `plotBMRRanksAsBarChart()`, `plotBMRSummary()`, `plotCritDifferences()`, `reduceBatchmarkResults()`

Examples

```
# see benchmark
```

```
plotBMRRanksAsBarChart
```

Create a bar chart for ranks in a BenchmarkResult.

Description

Plots a bar chart from the ranks of algorithms. Alternatively, tiles can be plotted for every rank-task combination, see `pos` for details. In all plot variants the ranks of the learning algorithms are displayed on the x-axis. Areas are always colored according to the learner .id.

Usage

```
plotBMRRanksAsBarChart(
  bmr,
  measure = NULL,
  ties.method = "average",
  aggregation = "default",
  pos = "stack",
  order.lrns = NULL,
  order.tsks = NULL,
  pretty.names = TRUE
)
```

Arguments

<code>bmr</code>	(BenchmarkResult) Benchmark result.
<code>measure</code>	(Measure) Performance measure. Default is the first measure used in the benchmark experiment.

ties.method	(character(1)) See rank for details.
aggregation	(character(1)) “mean” or “default”. See getBMRAggrPerformances for details on “default”.
pos	(character(1)) Optionally set how the bars are positioned in ggplot2. Ranks are plotted on the x-axis. “tile” plots a heat map with task as the y-axis. Allows identification of the performance in a special task. “stack” plots a stacked bar plot. Allows for comparison of learners within and across ranks. “dodge” plots a bar plot with bars next to each other instead of stacked bars.
order.lrn	(character(n.learners)) Character vector with learner.ids in new order.
order.tsks	(character(n.tasks)) Character vector with task.ids in new order.
pretty.names	(logical(1)) Whether to use the short name of the learner instead of its ID in labels. Defaults to TRUE.

Value

ggplot2 plot object.

See Also

Other plot: [createSpatialResamplingPlots\(\)](#), [plotBMRBoxplots\(\)](#), [plotBMRSummary\(\)](#), [plotCalibration\(\)](#), [plotCritDifferences\(\)](#), [plotLearningCurve\(\)](#), [plotPartialDependence\(\)](#), [plotROCCurves\(\)](#), [plotResiduals\(\)](#), [plotThreshVsPerf\(\)](#)

Other benchmark: [BenchmarkResult](#), [batchmark\(\)](#), [benchmark\(\)](#), [convertBMRTToRankMatrix\(\)](#), [friedmanPostHocTestBMR\(\)](#), [friedmanTestBMR\(\)](#), [generateCritDifferencesData\(\)](#), [getBMRAggrPerformances\(\)](#), [getBMRFeatSelResults\(\)](#), [getBMRFilteredFeatures\(\)](#), [getBMRLearnerIds\(\)](#), [getBMRLearnerShortNames\(\)](#), [getBMRLearners\(\)](#), [getBMRMeasureIds\(\)](#), [getBMRMeasures\(\)](#), [getBMRModels\(\)](#), [getBMRPerformances\(\)](#), [getBMRPredictions\(\)](#), [getBMRTaskDescs\(\)](#), [getBMRTaskIds\(\)](#), [getBMRTuneResults\(\)](#), [plotBMRBoxplots\(\)](#), [plotBMRSummary\(\)](#), [plotCritDifferences\(\)](#), [reduceBatchmarkResults\(\)](#)

Examples

```
# see benchmark
```

plotBMRSummary *Plot a benchmark summary.*

Description

Creates a scatter plot, where each line refers to a task. On that line the aggregated scores for all learners are plotted, for that task. Optionally, you can apply a rank transformation or just use one of ggplot2’s transformations like [ggplot2::scale_x_log10](#).

Usage

```
plotBMRSummary(
  bmr,
  measure = NULL,
  trafo = "none",
  order.tsks = NULL,
  pointsize = 4L,
  jitter = 0.05,
  pretty.names = TRUE
)
```

Arguments

bmr	(BenchmarkResult) Benchmark result.
measure	(Measure) Performance measure. Default is the first measure used in the benchmark experiment.
trafo	(<code>character(1)</code>) Currently either “none” or “rank”, the latter performing a rank transformation (with average handling of ties) of the scores per task. NB: You can add always add <code>ggplot2::scale_x_log10</code> to the result to put scores on a log scale. Default is “none”.
order.tsks	(<code>character(n.tasks)</code>) Character vector with <code>task.ids</code> in new order.
pointsize	(<code>numeric(1)</code>) Point size for <code>ggplot2::geom_point</code> for data points. Default is 4.
jitter	(<code>numeric(1)</code>) Small vertical jitter to deal with overplotting in case of equal scores. Default is 0.05.
pretty.names	(<code>logical(1)</code>) Whether to use the short name of the learner instead of its ID in labels. Defaults to TRUE.

Value

ggplot2 plot object.

See Also

Other benchmark: [BenchmarkResult](#), [batchmark\(\)](#), [benchmark\(\)](#), [convertBMRTToRankMatrix\(\)](#), [friedmanPostHocTestBMR\(\)](#), [friedmanTestBMR\(\)](#), [generateCritDifferencesData\(\)](#), [getBMRAggrPerformances\(\)](#), [getBMRFeatSelResults\(\)](#), [getBMRFilteredFeatures\(\)](#), [getBMRLearnerIds\(\)](#), [getBMRLearnerShortNames\(\)](#), [getBMRLearners\(\)](#), [getBMRMeasureIds\(\)](#), [getBMRMeasures\(\)](#), [getBMRModels\(\)](#), [getBMRPerformances\(\)](#), [getBMRPredictions\(\)](#), [getBMRTaskDescs\(\)](#), [getBMRTaskIds\(\)](#), [getBMRTuneResults\(\)](#), [plotBMRBoxplots\(\)](#), [plotBMRRanksAsBarChart\(\)](#), [plotCritDifferences\(\)](#), [reduceBatchmarkResults\(\)](#)

Other plot: [createSpatialResamplingPlots\(\)](#), [plotBMRBoxplots\(\)](#), [plotBMRRanksAsBarChart\(\)](#), [plotCalibration\(\)](#), [plotCritDifferences\(\)](#), [plotLearningCurve\(\)](#), [plotPartialDependence\(\)](#), [plotROCCurves\(\)](#), [plotResiduals\(\)](#), [plotThreshVsPerf\(\)](#)

Examples

```
# see benchmark
```

plotCalibration	<i>Plot calibration data using ggplot2.</i>
-----------------	---

Description

Plots calibration data from [generateCalibrationData](#).

Usage

```
plotCalibration(
  obj,
  smooth = FALSE,
  reference = TRUE,
  rag = TRUE,
  facet.wrap.nrow = NULL,
  facet.wrap.ncol = NULL
)
```

Arguments

obj	(CalibrationData) Result of generateCalibrationData .
smooth	(logical(1)) Whether to use a loess smoother. Default is FALSE.
reference	(logical(1)) Whether to plot a reference line showing perfect calibration. Default is TRUE.
rag	(logical(1)) Whether to include a rag plot which shows a rug plot on the top which pertains to positive cases and on the bottom which pertains to negative cases. Default is TRUE.
facet.wrap.nrow, facet.wrap.ncol	(integer) Number of rows and columns for faceting. Default for both is NULL. In this case ggplot's <code>facet_wrap</code> will choose the layout itself.

Value

ggplot2 plot object.

See Also

Other plot: [createSpatialResamplingPlots\(\)](#), [plotBMRBoxplots\(\)](#), [plotBMRRanksAsBarChart\(\)](#), [plotBMRSummary\(\)](#), [plotCritDifferences\(\)](#), [plotLearningCurve\(\)](#), [plotPartialDependence\(\)](#), [plotROCCurves\(\)](#), [plotResiduals\(\)](#), [plotThreshVsPerf\(\)](#)

Other calibration: [generateCalibrationData\(\)](#)

Examples

```
## Not run:
lrns = list(makeLearner("classif.rpart", predict.type = "prob"),
  makeLearner("classif.nnet", predict.type = "prob"))
fit = lapply(lrns, train, task = iris.task)
pred = lapply(fit, predict, task = iris.task)
names(pred) = c("rpart", "nnet")
out = generateCalibrationData(pred, groups = 3)
plotCalibration(out)

fit = lapply(lrns, train, task = sonar.task)
pred = lapply(fit, predict, task = sonar.task)
names(pred) = c("rpart", "lda")
out = generateCalibrationData(pred)
plotCalibration(out)

## End(Not run)
```

plotCritDifferences *Plot critical differences for a selected measure.*

Description

Plots a critical-differences diagram for all classifiers and a selected measure. If a baseline is selected for the Bonferroni-Dunn test, the critical difference interval will be positioned around the baseline. If not, the best performing algorithm will be chosen as baseline.

The positioning of some descriptive elements can be moved by modifying the generated data.

Usage

```
plotCritDifferences(obj, baseline = NULL, pretty.names = TRUE)
```

Arguments

obj (critDifferencesData) Result of [generateCritDifferencesData\(\)](#).

baseline (character(1)): (learner.id)
Overwrites baseline from [generateCritDifferencesData\(\)](#)!
Select a learner.id as baseline for the critical difference diagram, the critical difference will be positioned around this learner. Defaults to best performing algorithm.

pretty.names (logical(1))
Whether to use the short name of the learner instead of its ID in labels. Defaults to TRUE.

Value

ggplot2 plot object.

References

Janez Demsar, Statistical Comparisons of Classifiers over Multiple Data Sets, JMLR, 2006

See Also

Other plot: `createSpatialResamplingPlots()`, `plotBMRBoxplots()`, `plotBMRRanksAsBarChart()`, `plotBMRSummary()`, `plotCalibration()`, `plotLearningCurve()`, `plotPartialDependence()`, `plotROCCurves()`, `plotResiduals()`, `plotThreshVsPerf()`

Other benchmark: `BenchmarkResult`, `batchmark()`, `benchmark()`, `convertBMRToRankMatrix()`, `friedmanPostHocTestBMR()`, `friedmanTestBMR()`, `generateCritDifferencesData()`, `getBMRAggrPerformances()`, `getBMRFeatSelResults()`, `getBMRFilteredFeatures()`, `getBMRLearnerIds()`, `getBMRLearnerShortNames()`, `getBMRLearners()`, `getBMRMeasureIds()`, `getBMRMeasures()`, `getBMRModels()`, `getBMRPerformances()`, `getBMRPredictions()`, `getBMRTaskDescs()`, `getBMRTaskIds()`, `getBMRTuneResults()`, `plotBMRBoxplots()`, `plotBMRRanksAsBarChart()`, `plotBMRSummary()`, `reduceBatchmarkResults()`

Examples

```
# see benchmark
```

plotFilterValues *Plot filter values using ggplot2.*

Description

Plot filter values using ggplot2.

Usage

```
plotFilterValues(
  fvalues,
  sort = "dec",
  n.show = nrow(fvalues$data),
  filter = NULL,
  feat.type.cols = FALSE
)
```

Arguments

fvalues	(FilterValues) Filter values.
sort	(character(1)) Available options are: <ul style="list-style-type: none"> • "dec" -> descending • "inc" -> increasing • "none" -> no sorting Default is decreasing.
n.show	(integer(1)) Number of features (maximal) to show. Default is to plot all features.
filter	(character(1)) In case fvalues contains multiple filter methods, which method should be plotted?
feat.type.cols	(logical(1)) Whether to color different feature types (e.g. numeric factor). Default is to use no colors (feat.type.cols = FALSE).

Value

ggplot2 plot object.

See Also

Other filter: [filterFeatures\(\)](#), [generateFilterValuesData\(\)](#), [getFilteredFeatures\(\)](#), [listFilterEnsembleMethods\(\)](#), [listFilterMethods\(\)](#), [makeFilterEnsemble\(\)](#), [makeFilterWrapper\(\)](#), [makeFilter\(\)](#)

Other generate_plot_data: [generateCalibrationData\(\)](#), [generateCritDifferencesData\(\)](#), [generateFeatureImportanceData\(\)](#), [generateFilterValuesData\(\)](#), [generateLearningCurveData\(\)](#), [generatePartialDependenceData\(\)](#), [generateThreshVsPerfData\(\)](#)

Examples

```
fV = generateFilterValuesData(iris.task, method = "variance")
plotFilterValues(fV)
```

plotHyperParsEffect *Plot the hyperparameter effects data*

Description

Plot hyperparameter validation path. Automated plotting method for HyperParsEffectData object. Useful for determining the importance or effect of a particular hyperparameter on some performance measure and/or optimizer.

Usage

```
plotHyperParsEffect(
  hyperpars.effect.data,
  x = NULL,
  y = NULL,
  z = NULL,
  plot.type = "scatter",
  loess.smooth = FALSE,
  facet = NULL,
  global.only = TRUE,
  interpolate = NULL,
  show.experiments = FALSE,
  show.interpolated = FALSE,
  nested.agg = mean,
  partial.dep.learn = NULL
)
```

Arguments

hyperpars.effect.data	(HyperParsEffectData) Result of generateHyperParsEffectData
x	(character(1)) Specify what should be plotted on the x axis. Must be a column from HyperParsEffectData\$data. For partial dependence, this is assumed to be a hyperparameter.
y	(character(1)) Specify what should be plotted on the y axis. Must be a column from HyperParsEffectData\$data
z	(character(1)) Specify what should be used as the extra axis for a particular geom. This could be for the fill on a heatmap or color aesthetic for a line. Must be a column from HyperParsEffectData\$data. Default is NULL.
plot.type	(character(1)) Specify the type of plot: "scatter" for a scatterplot, "heatmap" for a heatmap, "line" for a scatterplot with a connecting line, or "contour" for a contour plot layered on top of a heatmap. Default is "scatter".
loess.smooth	(logical(1)) If TRUE, will add loess smoothing line to plots where possible. Note that this is probably only useful when plot.type is set to either "scatter" or "line". Must be a column from HyperParsEffectData\$data. Not used with partial dependence. Default is FALSE.
facet	(character(1)) Specify what should be used as the facet axis for a particular geom. When using nested cross validation, set this to "nested_cv_run" to obtain a facet for each outer loop. Must be a column from HyperParsEffectData\$data. Please note that faceting is not supported with partial dependence plots! Default is NULL.
global.only	(logical(1)) If TRUE, will only plot the current global optima when setting x = "iteration" and

	<p>y as a performance measure from <code>HyperParsEffectData\$measures</code>. Set this to <code>FALSE</code> to always plot the performance of every iteration, even if it is not an improvement. Not used with partial dependence. Default is <code>TRUE</code>.</p>
<code>interpolate</code>	<p>(Learner character(1))</p> <p>If not <code>NULL</code>, will interpolate non-complete grids in order to visualize a more complete path. Only meaningful when attempting to plot a heatmap or contour. This will fill in “empty” cells in the heatmap or contour plot. Note that cases of irregular hyperparameter paths, you will most likely need to use this to have a meaningful visualization. Accepts either a regression Learner object or the learner as a string for interpolation. This cannot be used with partial dependence. Default is <code>NULL</code>.</p>
<code>show.experiments</code>	<p>(logical(1))</p> <p>If <code>TRUE</code>, will overlay the plot with points indicating where an experiment ran. This is only useful when creating a heatmap or contour plot with interpolation so that you can see which points were actually on the original path. Note: if any learner crashes occurred within the path, this will become <code>TRUE</code>. Not used with partial dependence. Default is <code>FALSE</code>.</p>
<code>show.interpolated</code>	<p>(logical(1))</p> <p>If <code>TRUE</code>, will overlay the plot with points indicating where interpolation ran. This is only useful when creating a heatmap or contour plot with interpolation so that you can see which points were interpolated. Not used with partial dependence. Default is <code>FALSE</code>.</p>
<code>nested.agg</code>	<p>(function)</p> <p>The function used to aggregate nested cross validation runs when plotting 2 hyperparameters. This is also used for nested aggregation in partial dependence. Default is <code>mean</code>.</p>
<code>partial.dep.learner</code>	<p>(Learner character(1))</p> <p>The regression learner used to learn partial dependence. Must be specified if “<code>partial.dep</code>” is set to <code>TRUE</code> in generateHyperParsEffectData. Accepts either a Learner object or the learner as a string for learning partial dependence. Default is <code>NULL</code>.</p>

Value

ggplot2 plot object.

Note

Any NAs incurred from learning algorithm crashes will be indicated in the plot (except in the case of partial dependence) and the NA values will be replaced with the column min/max depending on the optimal values for the respective measure. Execution time will be replaced with the max. Interpolation by its nature will result in predicted values for the performance measure. Use interpolation with caution. If “`partial.dep`” is set to `TRUE` in [generateHyperParsEffectData](#), only partial dependence will be plotted.

Since a ggplot2 plot object is returned, the user can change the axis labels and other aspects of the plot using the appropriate ggplot2 syntax.

Examples

```
# see generateHyperParsEffectData
```

plotLearnerPrediction *Visualizes a learning algorithm on a 1D or 2D data set.*

Description

Trains the model for 1 or 2 selected features, then displays it via [ggplot2::ggplot](#). Good for teaching or exploring models.

For classification and clustering, only 2D plots are supported. The data points, the classification and potentially through color alpha blending the posterior probabilities are shown.

For regression, 1D and 2D plots are supported. 1D shows the data, the estimated mean and potentially the estimated standard error. 2D does not show estimated standard error, but only the estimated mean via background color.

The plot title displays the model id, its parameters, the training performance and the cross-validation performance.

Usage

```
plotLearnerPrediction(
  learner,
  task,
  features = NULL,
  measures,
  cv = 10L,
  ...,
  gridsize,
  pointsize = 2,
  prob.alpha = TRUE,
  se.band = TRUE,
  err.mark = "train",
  bg.cols = c("darkblue", "green", "darkred"),
  err.col = "white",
  err.size = pointsize,
  greyscale = FALSE,
  pretty.names = TRUE
)
```

Arguments

learner	(Learner character(1)) The learner. If you pass a string the learner will be created via makeLearner .
task	(Task) The task.

features	(character) Selected features for model. By default the first 2 features are used.
measures	(Measure list of Measure) Performance measure(s) to evaluate. Default is the default measure for the task, see here getDefaultMeasure .
cv	(integer(1)) Do cross-validation and display in plot title? Number of folds. 0 means no CV. Default is 10.
...	(any) Parameters for learner.
gridsize	(integer(1)) Grid resolution per axis for background predictions. Default is 500 for 1D and 100 for 2D.
pointsize	(numeric(1)) Pointsize for ggplot2 geom_point for data points. Default is 2.
prob.alpha	(logical(1)) For classification: Set alpha value of background to probability for predicted class? Allows visualization of “confidence” for prediction. If not, only a constant color is displayed in the background for the predicted label. Default is TRUE.
se.band	(logical(1)) For regression in 1D: Show band for standard error estimation? Default is TRUE.
err.mark	(character(1)): For classification: Either mark error of the model on the training data (“train”) or during cross-validation (“cv”) or not at all with “none”. Default is “train”.
bg.cols	(character(3)) Background colors for classification and regression. Sorted from low, medium to high. Default is TRUE.
err.col	(character(1)) For classification: Color of misclassified data points. Default is “white”
err.size	(integer(1)) For classification: Size of misclassified data points. Default is pointsize.
greyscale	(logical(1)) Should the plot be greyscale completely? Default is FALSE.
pretty.names	(logical(1)) Whether to use the short name of the learner instead of its ID in labels. Defaults to TRUE.

Value

The ggplot2 object.

plotLearningCurve *Plot learning curve data using ggplot2.*

Description

Visualizes data size (percentage used for model) vs. performance measure(s).

Usage

```
plotLearningCurve(  
  obj,  
  facet = "measure",  
  pretty.names = TRUE,  
  facet.wrap.nrow = NULL,  
  facet.wrap.ncol = NULL  
)
```

Arguments

obj	(LearningCurveData) Result of generateLearningCurveData , with class LearningCurveData.
facet	(character(1)) Selects “measure” or “learner” to be the facetting variable. The variable mapped to facet must have more than one unique value, otherwise it will be ignored. The variable not chosen is mapped to color if it has more than one unique value. The default is “measure”.
pretty.names	(logical(1)) Whether to use the Measure name instead of the id in the plot. Default is TRUE.
facet.wrap.nrow, facet.wrap.ncol	(integer) Number of rows and columns for facetting. Default for both is NULL. In this case ggplot’s facet_wrap will choose the layout itself.

Value

ggplot2 plot object.

See Also

Other learning_curve: [generateLearningCurveData\(\)](#)

Other plot: [createSpatialResamplingPlots\(\)](#), [plotBMRBoxplots\(\)](#), [plotBMRRanksAsBarChart\(\)](#), [plotBMRSummary\(\)](#), [plotCalibration\(\)](#), [plotCritDifferences\(\)](#), [plotPartialDependence\(\)](#), [plotROCCurves\(\)](#), [plotResiduals\(\)](#), [plotThreshVsPerf\(\)](#)

plotPartialDependence *Plot a partial dependence with ggplot2.*

Description

Plot a partial dependence from [generatePartialDependenceData](#) using ggplot2.

Usage

```
plotPartialDependence(  
  obj,  
  geom = "line",  
  facet = NULL,  
  facet.wrap.nrow = NULL,  
  facet.wrap.ncol = NULL,  
  p = 1,  
  data = NULL  
)
```

Arguments

obj	PartialDependenceData Generated by generatePartialDependenceData .
geom	(character(1)) The type of geom to use to display the data. Can be "line" or "tile". For tiling at least two features must be used with <code>interaction = TRUE</code> in the call to generatePartialDependenceData . This may be used in conjunction with the facet argument if three features are specified in the call to generatePartialDependenceData . Default is "line".
facet	(character(1)) The name of a feature to be used for facetting. This feature must have been an element of the features argument to generatePartialDependenceData and is only applicable when said argument had length greater than 1. The feature must be a factor or an integer. If generatePartialDependenceData is called with the <code>interaction</code> argument <code>FALSE</code> (the default) with argument features of length greater than one, then facet is ignored and each feature is plotted in its own facet. Default is <code>NULL</code> .
facet.wrap.nrow, facet.wrap.ncol	(integer) Number of rows and columns for facetting. Default for both is <code>NULL</code> . In this case ggplot's <code>facet_wrap</code> will choose the layout itself.
p	(numeric(1)) If <code>individual = TRUE</code> then <code>sample</code> allows the user to sample without replacement from the output to make the display more readable. Each row is sampled with probability <code>p</code> . Default is 1.

`data` ([data.frame](#))
 Data points to plot. Usually the training data. For survival and binary classification tasks a rug plot wherein ticks represent failures or instances of the positive class are shown. For regression tasks points are shown. For multiclass classification tasks ticks are shown and colored according to their class. Both the features and the target must be included. Default is NULL.

Value

ggplot2 plot object.

See Also

Other partial_dependence: [generatePartialDependenceData\(\)](#)

Other plot: [createSpatialResamplingPlots\(\)](#), [plotBMRBoxplots\(\)](#), [plotBMRRanksAsBarChart\(\)](#), [plotBMRSummary\(\)](#), [plotCalibration\(\)](#), [plotCritDifferences\(\)](#), [plotLearningCurve\(\)](#), [plotROCCurves\(\)](#), [plotResiduals\(\)](#), [plotThreshVsPerf\(\)](#)

<code>plotResiduals</code>	<i>Create residual plots for prediction objects or benchmark results.</i>
----------------------------	---

Description

Plots for model diagnostics. Provides scatterplots of true vs. predicted values and histograms of the model's residuals.

Usage

```
plotResiduals(
  obj,
  type = "scatterplot",
  loess.smooth = TRUE,
  rug = TRUE,
  pretty.names = TRUE
)
```

Arguments

<code>obj</code>	(Prediction BenchmarkResult) Input data.
<code>type</code>	Type of plot. Can be "scatterplot", the default. Or "hist", for a histogram, or in case of classification problems a barplot, displaying the residuals.
<code>loess.smooth</code>	(logical(1)) Should a loess smoother be added to the plot? Defaults to TRUE. Only applicable for regression tasks and if type is set to scatterplot.

rug	(logical(1)) Should marginal distributions be added to the plot? Defaults to TRUE. Only applicable for regression tasks and if type is set to scatterplot.
pretty.names	(logical(1)) Whether to use the short name of the learner instead of its ID in labels. Defaults to TRUE. Only applicable if a BenchmarkResult is passed to obj in the function call, ignored otherwise.

Value

ggplot2 plot object.

See Also

Other plot: [createSpatialResamplingPlots\(\)](#), [plotBMRBoxplots\(\)](#), [plotBMRRanksAsBarChart\(\)](#), [plotBMRSummary\(\)](#), [plotCalibration\(\)](#), [plotCritDifferences\(\)](#), [plotLearningCurve\(\)](#), [plotPartialDependence\(\)](#), [plotROCCurves\(\)](#), [plotThreshVsPerf\(\)](#)

plotROCCurves	<i>Plots a ROC curve using ggplot2.</i>
---------------	---

Description

Plots a ROC curve from predictions.

Usage

```
plotROCCurves(
  obj,
  measures,
  diagonal = TRUE,
  pretty.names = TRUE,
  facet.learner = FALSE
)
```

Arguments

obj	(ThreshVsPerfData) Result of generateThreshVsPerfData .
measures	(list(2)' of Measure) Default is the first 2 measures passed to generateThreshVsPerfData .
diagonal	(logical(1)) Whether to plot a dashed diagonal line. Default is TRUE.
pretty.names	(logical(1)) Whether to use the Measure name instead of the id in the plot. Default is TRUE.

facet.learner (logical(1))
 Weather to use facetting or different colors to compare multiple learners. Default is FALSE.

Value

ggplot2 plot object.

See Also

Other plot: [createSpatialResamplingPlots\(\)](#), [plotBMRBoxplots\(\)](#), [plotBMRRanksAsBarChart\(\)](#), [plotBMRSummary\(\)](#), [plotCalibration\(\)](#), [plotCritDifferences\(\)](#), [plotLearningCurve\(\)](#), [plotPartialDependence\(\)](#), [plotResiduals\(\)](#), [plotThreshVsPerf\(\)](#)

Other thresh_vs_perf: [generateThreshVsPerfData\(\)](#), [plotThreshVsPerf\(\)](#)

Examples

```
lrn = makeLearner("classif.rpart", predict.type = "prob")
fit = train(lrn, sonar.task)
pred = predict(fit, task = sonar.task)
roc = generateThreshVsPerfData(pred, list(fpr, tpr))
plotROCCurves(roc)

r = bootstrapB632plus(lrn, sonar.task, iters = 3)
roc_r = generateThreshVsPerfData(r, list(fpr, tpr), aggregate = FALSE)
plotROCCurves(roc_r)

r2 = crossval(lrn, sonar.task, iters = 3)
roc_l = generateThreshVsPerfData(list(boot = r, cv = r2), list(fpr, tpr), aggregate = FALSE)
plotROCCurves(roc_l)
```

plotThreshVsPerf	<i>Plot threshold vs. performance(s) for 2-class classification using ggplot2.</i>
------------------	--

Description

Plots threshold vs. performance(s) data that has been generated with [generateThreshVsPerfData](#).

Usage

```
plotThreshVsPerf(
  obj,
  measures = obj$measures,
  facet = "measure",
  mark.th = NA_real_,
```

```

pretty.names = TRUE,
facet.wrap.nrow = NULL,
facet.wrap.ncol = NULL
)

```

Arguments

obj	(ThreshVsPerfData) Result of generateThreshVsPerfData .
measures	(Measure list of Measure) Performance measure(s) to plot. Must be a subset of those used in generateThreshVsPerfData . Default is all the measures stored in obj generated by generateThreshVsPerfData .
facet	(character(1)) Selects “measure” or “learner” to be the faceting variable. The variable mapped to facet must have more than one unique value, otherwise it will be ignored. The variable not chosen is mapped to color if it has more than one unique value. The default is “measure”.
mark.th	(numeric(1)) Mark given threshold with vertical line? Default is NA which means not to do it.
pretty.names	(logical(1)) Whether to use the Measure name instead of the id in the plot. Default is TRUE.
facet.wrap.nrow, facet.wrap.ncol	(integer) Number of rows and columns for faceting. Default for both is NULL. In this case ggplot’s facet_wrap will choose the layout itself.

Value

ggplot2 plot object.

See Also

Other plot: [createSpatialResamplingPlots\(\)](#), [plotBMRBoxplots\(\)](#), [plotBMRRanksAsBarChart\(\)](#), [plotBMRSummary\(\)](#), [plotCalibration\(\)](#), [plotCritDifferences\(\)](#), [plotLearningCurve\(\)](#), [plotPartialDependence\(\)](#), [plotROCCurves\(\)](#), [plotResiduals\(\)](#)

Other thresh_vs_perf: [generateThreshVsPerfData\(\)](#), [plotROCCurves\(\)](#)

Examples

```

lrn = makeLearner("classif.rpart", predict.type = "prob")
mod = train(lrn, sonar.task)
pred = predict(mod, sonar.task)
pvs = generateThreshVsPerfData(pred, list(acc, setAggregation(acc, train.mean)))
plotThreshVsPerf(pvs)

```

`plotTuneMultiCritResult`*Plots multi-criteria results after tuning using ggplot2.*

Description

Visualizes the pareto front and possibly the dominated points.

Usage

```
plotTuneMultiCritResult(  
  res,  
  path = TRUE,  
  col = NULL,  
  shape = NULL,  
  pointsize = 2,  
  pretty.names = TRUE  
)
```

Arguments

<code>res</code>	(TuneMultiCritResult) Result of tuneParamsMultiCrit .
<code>path</code>	(<code>logical(1)</code>) Visualize all evaluated points (or only the non-dominated pareto front)? For the full path, the size of the points on the front is slightly increased. Default is TRUE.
<code>col</code>	(<code>character(1)</code>) Which column of <code>res\$opt.path</code> should be mapped to ggplot2 color? Default is NULL, which means none.
<code>shape</code>	(<code>character(1)</code>) Which column of <code>res\$opt.path</code> should be mapped to ggplot2 shape? Default is NULL, which means none.
<code>pointsize</code>	(<code>numeric(1)</code>) Point size for ggplot2 ggplot2::geom_point for data points. Default is 2.
<code>pretty.names</code>	(<code>logical(1)</code>) Whether to use the ID of the measures instead of their name in labels. Defaults to TRUE.

Value

ggplot2 plot object.

See Also

Other `tune_multicrit`: [TuneMultiCritControl](#), [tuneParamsMultiCrit\(\)](#)

Examples

```
# see tuneParamsMultiCrit
```

```
predict.WrappedModel Predict new data.
```

Description

Predict the target variable of new data using a fitted model. What is stored exactly in the ([Prediction](#)) object depends on the `predict.type` setting of the [Learner](#). If `predict.type` was set to “prob” probability thresholding can be done calling the [setThreshold](#) function on the prediction object.

The row names of the input task or newdata are preserved in the output.

Usage

```
## S3 method for class 'WrappedModel'
predict(object, task, newdata, subset = NULL, ...)
```

Arguments

object	(WrappedModel) Wrapped model, result of train .
task	(Task) The task. If this is passed, data from this task is predicted.
newdata	(data.frame) New observations which should be predicted. Pass this alternatively instead of task.
subset	(integer logical NULL) Selected cases. Either a logical or an index vector. By default NULL if all observations are used.
...	(any) Currently ignored.

Value

([Prediction](#)).

See Also

Other predict: [asROCRPrediction\(\)](#), [getPredictionProbabilities\(\)](#), [getPredictionResponse\(\)](#), [getPredictionTaskDesc\(\)](#), [setPredictThreshold\(\)](#), [setPredictType\(\)](#)

Examples

```
# train and predict
train.set = seq(1, 150, 2)
test.set = seq(2, 150, 2)
model = train("classif.lda", iris.task, subset = train.set)
p = predict(model, newdata = iris, subset = test.set)
print(p)
predict(model, task = iris.task, subset = test.set)

# predict now probabilities instead of class labels
lrn = makeLearner("classif.lda", predict.type = "prob")
model = train(lrn, iris.task, subset = train.set)
p = predict(model, task = iris.task, subset = test.set)
print(p)
getPredictionProbabilities(p)
```

predictLearner

Predict new data with an R learner.

Description

Mainly for internal use. Predict new data with a fitted model. You have to implement this method if you want to add another learner to this package.

Usage

```
predictLearner(.learner, .model, .newdata, ...)
```

Arguments

.learner	(RLearner) Wrapped learner.
.model	(WrappedModel) Model produced by training.
.newdata	(data.frame) New data to predict. Does not include target column.
...	(any) Additional parameters, which need to be passed to the underlying predict function.

Details

Your implementation must adhere to the following: Predictions for the observations in `.newdata` must be made based on the fitted model (`.model$learner.model`). All parameters in `...` must be passed to the underlying predict function.

Value

- For classification: Either a factor with class labels for type “response” or, if the learner supports this, a matrix of class probabilities for type “prob”. In the latter case the columns must be named with the class labels.
- For regression: Either a numeric vector for type “response” or, if the learner supports this, a matrix with two columns for type “se”. In the latter case the first column contains the estimated response (mean value) and the second column the estimated standard errors.
- For survival: Either a numeric vector with some sort of orderable risk for type “response” or, if supported, a numeric vector with time dependent probabilities for type “prob”.
- For clustering: Either an integer with cluster IDs for type “response” or, if supported, a matrix of membership probabilities for type “prob”.
- For multilabel: A logical matrix that indicates predicted class labels for type “response” or, if supported, a matrix of class probabilities for type “prob”. The columns must be named with the class labels.

```
reduceBatchmarkResults
```

Reduce results of a batch-distributed benchmark.

Description

This creates a [BenchmarkResult](#) from a [batchtools::ExperimentRegistry](#). To setup the benchmark have a look at [benchmark](#).

Usage

```
reduceBatchmarkResults(
  ids = NULL,
  keep.pred = TRUE,
  keep.extract = FALSE,
  show.info = getMlrOption("show.info"),
  reg = batchtools::getDefaultRegistry()
)
```

Arguments

ids	(data.frame or integer) A base::data.frame (or data.table::data.table) with a column named “job.id”. Alternatively, you may also pass a vector of integerish job ids. If not set, defaults to all successfully terminated jobs (return value of batchtools::findDone).
keep.pred	(logical(1)) Keep the prediction data in the pred slot of the result object. If you do many experiments (on larger data sets) these objects might unnecessarily increase object size / mem usage, if you do not really need them. The default is set to TRUE.

keep.extract	(logical(1)) Keep the extract slot of the result object. When creating a lot of benchmark results with extensive tuning, the resulting R objects can become very large in size. That is why the tuning results stored in the extract slot are removed by default (keep.extract = FALSE). Note that when keep.extract = FALSE you will not be able to conduct analysis in the tuning results.
show.info	(logical(1)) Print verbose output on console? Default is set via configureMlr .
reg	(batchtools::ExperimentRegistry) Registry, created by batchtools::makeExperimentRegistry . If not explicitly passed, uses the last created registry.

Value

([BenchmarkResult](#)).

See Also

Other benchmark: [BenchmarkResult](#), [batchmark\(\)](#), [benchmark\(\)](#), [convertBMRTToRankMatrix\(\)](#), [friedmanPostHocTestBMR\(\)](#), [friedmanTestBMR\(\)](#), [generateCritDifferencesData\(\)](#), [getBMRAggrPerformances\(\)](#), [getBMRFeatSelResults\(\)](#), [getBMRFilteredFeatures\(\)](#), [getBMRLearnerIds\(\)](#), [getBMRLearnerShortNames\(\)](#), [getBMRLearners\(\)](#), [getBMRMeasureIds\(\)](#), [getBMRMeasures\(\)](#), [getBMRModels\(\)](#), [getBMRPerformances\(\)](#), [getBMRPredictions\(\)](#), [getBMRTaskDescs\(\)](#), [getBMRTaskIds\(\)](#), [getBMRTuneResults\(\)](#), [plotBMRBoxplots\(\)](#), [plotBMRRanksAsBarChart\(\)](#), [plotBMRSummary\(\)](#), [plotCritDifferences\(\)](#)

reextractFDAFeatures *Re-extract features from a data set*

Description

This function accepts a data frame or a task and an [extractFDAFeatDesc](#) (a FDA feature extraction description) as returned by [extractFDAFeatures](#) to extract features from previously unseen data.

Usage

```
reextractFDAFeatures(obj, desc, ...)
```

Arguments

obj	(Task data.frame) Task or data.frame to extract functional features from. Must contain functional features as matrix columns.
desc	(extractFDAFeatDesc) FDAFeature extraction description as returned by extractFDAFeatures
...	(any) Further args passed on to methods.

Value

[data.frame](#) or [Task](#) containing the extracted Features

reimpute	<i>Re-impute a data set</i>
----------	-----------------------------

Description

This function accepts a data frame or a task and an imputation description as returned by [impute](#) to perform the following actions:

1. Restore dropped columns, setting them to NA
2. Add dummy variables for columns as specified in `impute`
3. Optionally check factors for new levels to treat them as NAs
4. Reorder factor levels to ensure identical integer representation as before
5. Impute missing values using previously collected data

Usage

```
reimpute(obj, desc)
```

Arguments

obj	(data.frame Task) Input data.
desc	(ImputationDesc) Imputation description as returned by impute .

Value

Imputed `data.frame` or task with imputed data.

See Also

Other impute: [imputations](#), [impute\(\)](#), [makeImputeMethod\(\)](#), [makeImputeWrapper\(\)](#)

`removeConstantFeatures`*Remove constant features from a data set.*

Description

Constant features can lead to errors in some models and obviously provide no information in the training set that can be learned from. With the argument “perc”, there is a possibility to also remove features for which less than “perc” percent of the observations differ from the mode value.

Usage

```
removeConstantFeatures(  
  obj,  
  perc = 0,  
  dont.rm = character(0L),  
  na.ignore = FALSE,  
  tol = .Machine$double.eps^0.5,  
  show.info = getMlrOption("show.info")  
)
```

Arguments

<code>obj</code>	(data.frame Task) Input data.
<code>perc</code>	(numeric (1)) The percentage of a feature values in [0, 1) that must differ from the mode value. Default is 0, which means only constant features with exactly one observed level are removed.
<code>dont.rm</code>	(character) Names of the columns which must not be deleted. Default is no columns.
<code>na.ignore</code>	(logical (1)) Should NAs be ignored in the percentage calculation? (Or should they be treated as a single, extra level in the percentage calculation?) Note that if the feature has only missing values, it is always removed. Default is FALSE.
<code>tol</code>	(numeric (1)) Numerical tolerance to treat two numbers as equal. Variables stored as double will get rounded accordingly before computing the mode. Default is <code>sqrt(.Machine\$double.eps)</code> .
<code>show.info</code>	(logical (1)) Print verbose output on console? Default is set via configureMlr .

Value

[data.frame](#) | [Task](#). Same type as `obj`.

See Also

Other `eda_and_preprocess`: [capLargeValues\(\)](#), [createDummyFeatures\(\)](#), [dropFeatures\(\)](#), [mergeSmallFactorLevels\(\)](#), [normalizeFeatures\(\)](#), [summarizeColumns\(\)](#), [summarizeLevels\(\)](#)

removeHyperPars	<i>Remove hyperparameters settings of a learner.</i>
-----------------	--

Description

Remove settings (previously set through `mlr`) for some parameters. Which means that the default behavior for that param will now be used.

Usage

```
removeHyperPars(learner, ids = character(0L))
```

Arguments

learner	(Learner <code>character(1)</code>) The learner. If you pass a string the learner will be created via makeLearner .
ids	(<code>character</code>) Parameter names to remove settings for. Default is <code>character(0L)</code> .

Value

[Learner](#).

See Also

Other learner: [LearnerProperties](#), [getClassWeightParam\(\)](#), [getHyperPars\(\)](#), [getLearnerId\(\)](#), [getLearnerNote\(\)](#), [getLearnerPackages\(\)](#), [getLearnerParVals\(\)](#), [getLearnerParamSet\(\)](#), [getLearnerPredictType\(\)](#), [getLearnerShortName\(\)](#), [getLearnerType\(\)](#), [getParamSet\(\)](#), [helpLearnerParam\(\)](#), [helpLearner\(\)](#), [makeLearners\(\)](#), [makeLearner\(\)](#), [setHyperPars\(\)](#), [setId\(\)](#), [setLearnerId\(\)](#), [setPredictThreshold\(\)](#), [setPredictType\(\)](#)

`resample`*Fit models according to a resampling strategy.*

Description

The function `resample` fits a model specified by [Learner](#) on a [Task](#) and calculates predictions and performance [measures](#) for all training and all test sets specified by either a resampling description ([ResampleDesc](#)) or resampling instance ([ResampleInstance](#)).

You are able to return all fitted models (parameter models) or extract specific parts of the models (parameter extract) as returning all of them completely might be memory intensive.

The remaining functions on this page are convenience wrappers for the various existing resampling strategies. Note that if you need to work with precomputed training and test splits (i.e., resampling instances), you have to stick with `resample`.

Usage

```
resample(  
  learner,  
  task,  
  resampling,  
  measures,  
  weights = NULL,  
  models = FALSE,  
  extract,  
  keep.pred = TRUE,  
  ...,  
  show.info = getMlrOption("show.info")  
)
```

```
crossval(  
  learner,  
  task,  
  iters = 10L,  
  stratify = FALSE,  
  measures,  
  models = FALSE,  
  keep.pred = TRUE,  
  ...,  
  show.info = getMlrOption("show.info")  
)
```

```
repcv(  
  learner,  
  task,  
  folds = 10L,  
  reps = 10L,
```

```
    stratify = FALSE,
    measures,
    models = FALSE,
    keep.pred = TRUE,
    ...,
    show.info = getMlrOption("show.info")
)

holdout(
  learner,
  task,
  split = 2/3,
  stratify = FALSE,
  measures,
  models = FALSE,
  keep.pred = TRUE,
  ...,
  show.info = getMlrOption("show.info")
)

subsample(
  learner,
  task,
  iters = 30,
  split = 2/3,
  stratify = FALSE,
  measures,
  models = FALSE,
  keep.pred = TRUE,
  ...,
  show.info = getMlrOption("show.info")
)

bootstrapOoB(
  learner,
  task,
  iters = 30,
  stratify = FALSE,
  measures,
  models = FALSE,
  keep.pred = TRUE,
  ...,
  show.info = getMlrOption("show.info")
)

bootstrapB632(
  learner,
  task,
```

```
    iters = 30,  
    stratify = FALSE,  
    measures,  
    models = FALSE,  
    keep.pred = TRUE,  
    ...,  
    show.info = getMlrOption("show.info")  
)  
  
bootstrapB632plus(  
  learner,  
  task,  
  iters = 30,  
  stratify = FALSE,  
  measures,  
  models = FALSE,  
  keep.pred = TRUE,  
  ...,  
  show.info = getMlrOption("show.info")  
)  
  
growingcv(  
  learner,  
  task,  
  horizon = 1,  
  initial.window = 0.5,  
  skip = 0,  
  measures,  
  models = FALSE,  
  keep.pred = TRUE,  
  ...,  
  show.info = getMlrOption("show.info")  
)  
  
fixedcv(  
  learner,  
  task,  
  horizon = 1L,  
  initial.window = 0.5,  
  skip = 0,  
  measures,  
  models = FALSE,  
  keep.pred = TRUE,  
  ...,  
  show.info = getMlrOption("show.info")  
)
```

Arguments

learner	(Learner character(1)) The learner. If you pass a string the learner will be created via makeLearner .
task	(Task) The task.
resampling	(ResampleDesc or ResampleInstance) Resampling strategy. If a description is passed, it is instantiated automatically.
measures	(Measure list of Measure) Performance measure(s) to evaluate. Default is the default measure for the task, see here getDefaultMeasure .
weights	(numeric) Optional, non-negative case weight vector to be used during fitting. If given, must be of same length as observations in task and in corresponding order. Overwrites weights specified in the task. By default NULL which means no weights are used unless specified in the task.
models	(logical(1)) Should all fitted models be returned? Default is FALSE.
extract	(function) Function used to extract information from a fitted model during resampling. Is applied to every WrappedModel resulting from calls to train during resampling. Default is to extract nothing.
keep.pred	(logical(1)) Keep the prediction data in the pred slot of the result object. If you do many experiments (on larger data sets) these objects might unnecessarily increase object size / mem usage, if you do not really need them. The default is set to TRUE.
...	(any) Further hyperparameters passed to learner.
show.info	(logical(1)) Print verbose output on console? Default is set via configureMlr .
iters	(integer(1)) See ResampleDesc .
stratify	(logical(1)) See ResampleDesc .
folds	(integer(1)) See ResampleDesc .
reps	(integer(1)) See ResampleDesc .
split	(numeric(1)) See ResampleDesc .
horizon	(numeric(1)) See ResampleDesc .
initial.window	(numeric(1)) See ResampleDesc .
skip	(integer(1)) See ResampleDesc .

Value

(ResampleResult).

Note

If you would like to include results from the training data set, make sure to appropriately adjust the resampling strategy and the aggregation for the measure. See example code below.

See Also

Other resample: [ResamplePrediction](#), [ResampleResult](#), [addRRMeasure\(\)](#), [getRRPredictionList\(\)](#), [getRRPredictions\(\)](#), [getRRTaskDescription\(\)](#), [getRRTaskDesc\(\)](#), [makeResampleDesc\(\)](#), [makeResampleInstance\(\)](#)

Examples

```
task = makeClassifTask(data = iris, target = "Species")
rdesc = makeResampleDesc("CV", iters = 2)
r = resample(makeLearner("classif.qda"), task, rdesc)
print(r$aggr)
print(r$measures.test)
print(r$pred)

# include the training set performance as well
rdesc = makeResampleDesc("CV", iters = 2, predict = "both")
r = resample(makeLearner("classif.qda"), task, rdesc,
  measures = list(mmce, setAggregation(mmce, train.mean)))
print(r$aggr)
```

ResamplePrediction *Prediction from resampling.*

Description

Contains predictions from resampling, returned (among other stuff) by function [resample](#). Can basically be used in the same way as [Prediction](#), its super class. The main differences are: (a) The internal data.frame (member data) contains an additional column `iter`, specifying the iteration of the resampling strategy, and and additional columns `set`, specifying whether the prediction was from an observation in the “train” or “test” set. (b) The prediction `time` is a numeric vector, its length equals the number of iterations.

See Also

Other resample: [ResampleResult](#), [addRRMeasure\(\)](#), [getRRPredictionList\(\)](#), [getRRPredictions\(\)](#), [getRRTaskDescription\(\)](#), [getRRTaskDesc\(\)](#), [makeResampleDesc\(\)](#), [makeResampleInstance\(\)](#), [resample\(\)](#)

ResampleResult *ResampleResult object.*

Description

A container for resample results.

Details

Resample Result:

A resample result is created by `resample` and contains the following object members:

task.id (character(1)): Name of the Task.

learner.id (character(1)): Name of the Learner.

measures.test (**data.frame**): Gives you access to performance measurements on the individual test sets. Rows correspond to sets in resampling iterations, columns to performance measures.

measures.train (**data.frame**): Gives you access to performance measurements on the individual training sets. Rows correspond to sets in resampling iterations, columns to performance measures. Usually not available, only if specifically requested, see general description above.

aggr (**numeric**): Named vector of aggregated performance values. Names are coded like this `<measure>.<aggregation>`.

err.msgs (**data.frame**): Number of rows equals resampling iterations and columns are: `iter`, `train`, `predict`. Stores error messages generated during train or predict, if these were caught via `configureMlr`.

err.dumps (**list of list of dump.frames**): List with length equal to number of resampling iterations. Contains lists of `dump.frames` objects that can be fed to `debugger()` to inspect error dumps generated on learner errors. One iteration can generate more than one error dump depending on which of training, prediction on training set, or prediction on test set, operations fail. Therefore the lists have named slots `$train`, `$predict.train`, or `$predict.test` if relevant. The error dumps are only saved when option `on.error.dump` is TRUE.

pred (**ResamplePrediction**): Container for all predictions during resampling.

models [**list of WrappedModel**]: List of fitted models or NULL.

extract (**list**): List of extracted parts from fitted models or NULL.

runtime (**numeric(1)**): Time in seconds it took to execute the resampling.

The `print` method of this object gives a short overview, including task and learner ids, aggregated measures and runtime for the resampling.

See Also

Other resample: [ResamplePrediction](#), [addRRMeasure\(\)](#), [getRRPredictionList\(\)](#), [getRRPredictions\(\)](#), [getRRTaskDescription\(\)](#), [getRRTaskDesc\(\)](#), [makeResampleDesc\(\)](#), [makeResampleInstance\(\)](#), [resample\(\)](#)

Other debug: [FailureModel](#), [getPredictionDump\(\)](#), [getRRDump\(\)](#)

RLearner

Internal construction / wrapping of learner object.

Description

Wraps an already implemented learning method from R to make it accessible to mlr. Call this method in your constructor. You have to pass an id (name), the required package(s), a description object for all changeable parameters (you do not have to do this for the learner to work, but it is strongly recommended), and use property tags to define features of the learner.

For a general overview on how to integrate a learning algorithm into mlr's system, please read the section in the online tutorial: https://mlr.mlr-org.com/articles/tutorial/create_learner.html

To see all possible properties of a learner, go to: [LearnerProperties](#).

Usage

```
makeRLearner()
```

```
makeRLearnerClassif(  
  cl,  
  package,  
  par.set,  
  par.vals = list(),  
  properties = character(0L),  
  name = cl,  
  short.name = cl,  
  note = "",  
  class.weights.param = NULL,  
  callees = character(0L)  
)
```

```
makeRLearnerMultilabel(  
  cl,  
  package,  
  par.set,  
  par.vals = list(),  
  properties = character(0L),  
  name = cl,  
  short.name = cl,  
  note = "",  
  callees = character(0L)  
)
```

```
makeRLearnerRegr(  
  cl,  
  package,
```

```
    par.set,  
    par.vals = list(),  
    properties = character(0L),  
    name = cl,  
    short.name = cl,  
    note = "",  
    callees = character(0L)  
  )  
  
makeRLearnerSurv(  
  cl,  
  package,  
  par.set,  
  par.vals = list(),  
  properties = character(0L),  
  name = cl,  
  short.name = cl,  
  note = "",  
  callees = character(0L)  
)  
  
makeRLearnerCluster(  
  cl,  
  package,  
  par.set,  
  par.vals = list(),  
  properties = character(0L),  
  name = cl,  
  short.name = cl,  
  note = "",  
  callees = character(0L)  
)  
  
makeRLearnerCostSens(  
  cl,  
  package,  
  par.set,  
  par.vals = list(),  
  properties = character(0L),  
  name = cl,  
  short.name = cl,  
  note = "",  
  callees = character(0L)  
)
```

Arguments

`cl` (character(1))
Class of learner. By convention, all classification learners start with “classif.”

	all regression learners with “regr.” all survival learners start with “surv.” all clustering learners with “cluster.” and all multilabel classification learners start with “multilabel.”. A list of all integrated learners is available on the learners help page.
package	(character) Package(s) to load for the implementation of the learner.
par.set	(ParamHelpers::ParamSet) Parameter set of (hyper)parameters and their constraints. Dependent parameters with a requires field must use quote and not expression to define it.
par.vals	(list) Always set hyperparameters to these values when the object is constructed. Useful when default values are missing in the underlying function. The values can later be overwritten when the user sets hyperparameters. Default is empty list.
properties	(character) Set of learner properties. See above. Default is character(0).
name	(character(1)) Meaningful name for learner. Default is id.
short.name	(character(1)) Short name for learner. Should only be a few characters so it can be used in plots and tables. Default is id.
note	(character(1)) Additional notes regarding the learner and its integration in mlr. Default is “”.
class.weights.param	(character(1)) Name of the parameter, which can be used for providing class weights.
callees	(character) Character vector naming all functions of the learner’s package being called which have a relevant R help page. Default is character(0).

Value

(R`Learner`). The specific subclass is one of [R`LearnerClassif`](#), [R`LearnerCluster`](#), [R`LearnerMultilabel`](#), [R`LearnerRegr`](#), [R`LearnerSurv`](#).

selectFeatures	<i>Feature selection by wrapper approach.</i>
----------------	---

Description

Optimizes the features for a classification or regression problem by choosing a variable selection wrapper approach. Allows for different optimization methods, such as forward search or a genetic algorithm. You can select such an algorithm (and its settings) by passing a corresponding control object. For a complete list of implemented algorithms look at the subclasses of ([FeatSelControl](#)).

All algorithms operate on a 0-1-bit encoding of candidate solutions. Per default a single bit corresponds to a single feature, but you are able to change this by using the arguments `bit.names` and `bits.to.features`. Thus allowing you to switch on whole groups of features with a single bit.

Usage

```
selectFeatures(
  learner,
  task,
  resampling,
  measures,
  bit.names,
  bits.to.features,
  control,
  show.info = getMlrOption("show.info")
)
```

Arguments

learner	(Learner character(1)) The learner. If you pass a string the learner will be created via makeLearner .
task	(Task) The task.
resampling	(ResampleInstance ResampleDesc) Resampling strategy for feature selection. If you pass a description, it is instantiated once at the beginning by default, so all points are evaluated on the same training/test sets. If you want to change that behavior, look at FeatSelControl .
measures	(list of Measure Measure) Performance measures to evaluate. The first measure, aggregated by the first aggregation function is optimized, others are simply evaluated. Default is the default measure for the task, see here getDefaultMeasure .
bit.names	character Names of bits encoding the solutions. Also defines the total number of bits in the encoding. Per default these are the feature names of the task. Has to be used together with <code>bits.to.features</code> .
bits.to.features	(function(x, task)) Function which transforms an integer-0-1 vector into a character vector of selected features. Per default a value of 1 in the <i>i</i> th bit selects the <i>i</i> th feature to be in the candidate solution. The vector <i>x</i> will correspond to the <code>bit.names</code> and has to be of the same length.
control	[see FeatSelControl] Control object for search method. Also selects the optimization algorithm for feature selection.
show.info	(logical(1)) Print verbose output on console? Default is set via configureMlr .

Value

([FeatSelResult](#)).

See Also

Other featsel: [FeatSelControl](#), [analyzeFeatSelResult\(\)](#), [getFeatSelResult\(\)](#), [makeFeatSelWrapper\(\)](#)

Examples

```
rdesc = makeResampleDesc("Holdout")
ctrl = makeFeatSelControlSequential(method = "sfs", maxit = NA)
res = selectFeatures("classif.rpart", iris.task, rdesc, control = ctrl)
analyzeFeatSelResult(res)
```

setAggregation	<i>Set aggregation function of measure.</i>
----------------	---

Description

Set how this measure will be aggregated after resampling. To see possible aggregation functions: [aggregations](#).

Usage

```
setAggregation(measure, aggr)
```

Arguments

measure	(Measure) Performance measure.
aggr	(Aggregation) Aggregation function.

Value

([Measure](#)) with changed aggregation behaviour.

See Also

Other performance: [ConfusionMatrix](#), [calculateConfusionMatrix\(\)](#), [calculateROCMasures\(\)](#), [estimateRelativeOverfitting\(\)](#), [makeCostMeasure\(\)](#), [makeCustomResampledMeasure\(\)](#), [makeMeasure\(\)](#), [measures](#), [performance\(\)](#), [setMeasurePars\(\)](#)

setHyperPars *Set the hyperparameters of a learner object.*

Description

Set the hyperparameters of a learner object.

Usage

```
setHyperPars(learner, ..., par.vals = list())
```

Arguments

learner	(Learner character(1)) The learner. If you pass a string the learner will be created via makeLearner .
...	(any) Optional named (hyper)parameters. If you want to set specific hyperparameters for a learner during model creation, these should go here. You can get a list of available hyperparameters using <code>getParamSet(<learner>)</code> . Alternatively hyperparameters can be given using the <code>par.vals</code> argument but ... should be preferred!
par.vals	(list) Optional list of named (hyper)parameters. The arguments in ... take precedence over values in this list. We strongly encourage you to use ... for passing hyperparameters.

Value

[Learner](#).

Note

If a named (hyper)parameter can't be found for the given learner, the 3 closest (hyper)parameter names will be output in case the user mistyped.

See Also

Other learner: [LearnerProperties](#), [getClassWeightParam\(\)](#), [getHyperPars\(\)](#), [getLearnerId\(\)](#), [getLearnerNote\(\)](#), [getLearnerPackages\(\)](#), [getLearnerParVals\(\)](#), [getLearnerParamSet\(\)](#), [getLearnerPredictType\(\)](#), [getLearnerShortName\(\)](#), [getLearnerType\(\)](#), [getParamSet\(\)](#), [helpLearnerParam\(\)](#), [helpLearner\(\)](#), [makeLearners\(\)](#), [makeLearner\(\)](#), [removeHyperPars\(\)](#), [setId\(\)](#), [setLearnerId\(\)](#), [setPredictThreshold\(\)](#), [setPredictType\(\)](#)

Examples

```

c11 = makeLearner("classif.ksvm", sigma = 1)
c12 = setHyperPars(c11, sigma = 10, par.vals = list(C = 2))
print(c11)
# note the now set and altered hyperparameters:
print(c12)

```

setHyperPars2	<i>Only exported for internal use.</i>
---------------	--

Description

Only exported for internal use.

Usage

```
setHyperPars2(learner, par.vals)
```

Arguments

learner	(Learner) The learner.
par.vals	(list) List of named (hyper)parameter settings.

setId	<i>Set the id of a learner object.</i>
-------	--

Description

Deprecated, use [setLearnerId](#) instead.

Usage

```
setId(learner, id)
```

Arguments

learner	(Learner <code>character(1)</code>) The learner. If you pass a string the learner will be created via makeLearner .
id	(<code>character(1)</code>) New id for learner.

Value

[Learner](#).

See Also

Other learner: [LearnerProperties](#), [getClassWeightParam\(\)](#), [getHyperPars\(\)](#), [getLearnerId\(\)](#), [getLearnerNote\(\)](#), [getLearnerPackages\(\)](#), [getLearnerParVals\(\)](#), [getLearnerParamSet\(\)](#), [getLearnerPredictType\(\)](#), [getLearnerShortName\(\)](#), [getLearnerType\(\)](#), [getParamSet\(\)](#), [helpLearnerParam\(\)](#), [helpLearner\(\)](#), [makeLearners\(\)](#), [makeLearner\(\)](#), [removeHyperPars\(\)](#), [setHyperPars\(\)](#), [setLearnerId\(\)](#), [setPredictThreshold\(\)](#), [setPredictType\(\)](#)

setLearnerId	<i>Set the ID of a learner object.</i>
--------------	--

Description

Set the ID of the learner.

Usage

```
setLearnerId(learner, id)
```

Arguments

learner	(Learner character(1)) The learner. If you pass a string the learner will be created via makeLearner .
id	(character(1)) New ID for learner.

Value

[Learner](#).

See Also

Other learner: [LearnerProperties](#), [getClassWeightParam\(\)](#), [getHyperPars\(\)](#), [getLearnerId\(\)](#), [getLearnerNote\(\)](#), [getLearnerPackages\(\)](#), [getLearnerParVals\(\)](#), [getLearnerParamSet\(\)](#), [getLearnerPredictType\(\)](#), [getLearnerShortName\(\)](#), [getLearnerType\(\)](#), [getParamSet\(\)](#), [helpLearnerParam\(\)](#), [helpLearner\(\)](#), [makeLearners\(\)](#), [makeLearner\(\)](#), [removeHyperPars\(\)](#), [setHyperPars\(\)](#), [setId\(\)](#), [setPredictThreshold\(\)](#), [setPredictType\(\)](#)

setMeasurePars	<i>Set parameters of performance measures</i>
----------------	---

Description

Sets hyperparameters of measures.

Usage

```
setMeasurePars(measure, ..., par.vals = list())
```

Arguments

measure	(Measure) Performance measure.
...	(any) Named (hyper)parameters with new settings. Alternatively these can be passed using the <code>par.vals</code> argument.
par.vals	(list) Optional list of named (hyper)parameter settings. The arguments in <code>...</code> take precedence over values in this list.

Value

[Measure](#).

See Also

Other performance: [ConfusionMatrix](#), [calculateConfusionMatrix\(\)](#), [calculateROCMeasures\(\)](#), [estimateRelativeOverfitting\(\)](#), [makeCostMeasure\(\)](#), [makeCustomResampledMeasure\(\)](#), [makeMeasure\(\)](#), [measures](#), [performance\(\)](#), [setAggregation\(\)](#)

setPredictThreshold	<i>Set the probability threshold the learner should use.</i>
---------------------	--

Description

See `predict.threshold` in [makeLearner](#) and [setThreshold](#).

For complex wrappers only the top-level `predict.type` is currently set.

Usage

```
setPredictThreshold(learner, predict.threshold)
```

Arguments

learner (Learner | character(1))
The learner. If you pass a string the learner will be created via [makeLearner](#).

predict.threshold (numeric)
Threshold to produce class labels. Has to be a named vector, where names correspond to class labels. Only for binary classification it can be a single numerical threshold for the positive class. See [setThreshold](#) for details on how it is applied. Default is NULL which means 0.5 / an equal threshold for each class.

Value

Learner.

See Also

Other predict: [asROCRPrediction\(\)](#), [getPredictionProbabilities\(\)](#), [getPredictionResponse\(\)](#), [getPredictionTaskDesc\(\)](#), [predict.WrappedModel\(\)](#), [setPredictType\(\)](#)

Other learner: [LearnerProperties](#), [getClassWeightParam\(\)](#), [getHyperPars\(\)](#), [getLearnerId\(\)](#), [getLearnerNote\(\)](#), [getLearnerPackages\(\)](#), [getLearnerParVals\(\)](#), [getLearnerParamSet\(\)](#), [getLearnerPredictType\(\)](#), [getLearnerShortName\(\)](#), [getLearnerType\(\)](#), [getParamSet\(\)](#), [helpLearnerParam\(\)](#), [helpLearner\(\)](#), [makeLearners\(\)](#), [makeLearner\(\)](#), [removeHyperPars\(\)](#), [setHyperPars\(\)](#), [setId\(\)](#), [setLearnerId\(\)](#), [setPredictType\(\)](#)

setPredictType *Set the type of predictions the learner should return.*

Description

Possible prediction types are: Classification: Labels or class probabilities (including labels). Regression: Numeric or response or standard errors (including numeric response). Survival: Linear predictor or survival probability.

For complex wrappers the predict type is usually also passed down the encapsulated learner in a recursive fashion.

Usage

```
setPredictType(learner, predict.type)
```

Arguments

learner (Learner | character(1))
The learner. If you pass a string the learner will be created via [makeLearner](#).

predict.type (character(1))
Classification: “response” or “prob”. Regression: “response” or “se”. Survival: “response” (linear predictor) or “prob”. Clustering: “response” or “prob”. Default is “response”.

Value

[Learner](#).

See Also

Other predict: [asROCRPrediction\(\)](#), [getPredictionProbabilities\(\)](#), [getPredictionResponse\(\)](#), [getPredictionTaskDesc\(\)](#), [predict.WrappedModel\(\)](#), [setPredictThreshold\(\)](#)

Other learner: [LearnerProperties](#), [getClassWeightParam\(\)](#), [getHyperPars\(\)](#), [getLearnerId\(\)](#), [getLearnerNote\(\)](#), [getLearnerPackages\(\)](#), [getLearnerParVals\(\)](#), [getLearnerParamSet\(\)](#), [getLearnerPredictType\(\)](#), [getLearnerShortName\(\)](#), [getLearnerType\(\)](#), [getParamSet\(\)](#), [helpLearnerParam\(\)](#), [helpLearner\(\)](#), [makeLearners\(\)](#), [makeLearner\(\)](#), [removeHyperPars\(\)](#), [setHyperPars\(\)](#), [setId\(\)](#), [setLearnerId\(\)](#), [setPredictThreshold\(\)](#)

setThreshold	<i>Set threshold of prediction object.</i>
--------------	--

Description

Set threshold of prediction object for classification or multilabel classification. Creates corresponding discrete class response for the newly set threshold. For binary classification: The positive class is predicted if the probability value exceeds the threshold. For multiclass: Probabilities are divided by corresponding thresholds and the class with maximum resulting value is selected. The result of both are equivalent if in the multi-threshold case the values are greater than 0 and sum to 1. For multilabel classification: A label is predicted (with entry TRUE) if a probability matrix entry exceeds the threshold of the corresponding label.

Usage

```
setThreshold(pred, threshold)
```

Arguments

pred	(Prediction) Prediction object.
threshold	(numeric) Threshold to produce class labels. Has to be a named vector, where names correspond to class labels. Only for binary classification it can be a single numerical threshold for the positive class.

Value

[\(Prediction\)](#) with changed threshold and corresponding response.

See Also

[predict.WrappedModel](#)

Examples

```
# create task and train learner (LDA)
task = makeClassifTask(data = iris, target = "Species")
lrn = makeLearner("classif.lda", predict.type = "prob")
mod = train(lrn, task)

# predict probabilities and compute performance
pred = predict(mod, newdata = iris)
performance(pred, measures = mmce)
head(as.data.frame(pred))

# adjust threshold and predict probabilities again
threshold = c(setosa = 0.4, versicolor = 0.3, virginica = 0.3)
pred = setThreshold(pred, threshold = threshold)
performance(pred, measures = mmce)
head(as.data.frame(pred))
```

simplifyMeasureNames *Simplify measure names.*

Description

Clips aggregation names from character vector. E.g: 'mmce.test.mean' becomes 'mmce'. Elements that don't contain a measure name are ignored and returned unchanged.

Usage

```
simplifyMeasureNames(xs)
```

Arguments

xs [\(character\)](#)
Character vector that (possibly) contains aggregated measure names.

Value

[\(character\)](#).

smote	<i>Synthetic Minority Oversampling Technique to handle class imbalance in binary classification.</i>
-------	--

Description

In each iteration, samples one minority class element x_1 , then one of x_1 's nearest neighbors: x_2 . Both points are now interpolated / convex-combined, resulting in a new virtual data point x_3 for the minority class.

The method handles factor features, too. The gower distance is used for nearest neighbor calculation, see [cluster::daisy](#). For interpolation, the new factor level for x_3 is sampled from the two given levels of x_1 and x_2 per feature.

Usage

```
smote(task, rate, nn = 5L, standardize = TRUE, alt.logic = FALSE)
```

Arguments

task	(Task) The task.
rate	(numeric(1)) Factor to upsample the smaller class. Must be between 1 and Inf, where 1 means no oversampling and 2 would mean doubling the class size.
nn	(integer(1)) Number of nearest neighbors to consider. Default is 5.
standardize	(integer(1)) Standardize input variables before calculating the nearest neighbors for data sets with numeric input variables only. For mixed variables (numeric and factor) the gower distance is used and variables are standardized anyway. Default is TRUE.
alt.logic	(integer(1)) Use an alternative logic for selection of minority class observations. Instead of sampling a minority class element AND one of its nearest neighbors, each minority class element is taken multiple times (depending on rate) for the interpolation and only the corresponding nearest neighbor is sampled. Default is FALSE.

Value

[Task](#).

References

Chawla, N., Bowyer, K., Hall, L., & Kegelmeyer, P. (2000) *SMOTE: Synthetic Minority Oversampling TEchnique*. In International Conference of Knowledge Based Computer Systems, pp. 46-57. National Center for Software Technology, Mumbai, India, Allied Press.

See Also

Other imbalance: [makeOverBaggingWrapper\(\)](#), [makeUndersampleWrapper\(\)](#), [oversample\(\)](#)

sonar.task	<i>Sonar classification task.</i>
------------	-----------------------------------

Description

Contains the task (sonar.task).

References

See [mlbench::Sonar](#).

spam.task	<i>Spam classification task.</i>
-----------	----------------------------------

Description

Contains the task (spam.task).

References

See [kernlab::spam](#).

spatial.task	<i>J. Muenchow's Ecuador landslide data set</i>
--------------	---

Description

Data set created by Jannes Muenchow, University of Erlangen-Nuremberg, Germany. These data should be cited as Muenchow et al. (2012) (see reference below). This publication also contains additional information on data collection and the geomorphology of the area. The data set provided here is (a subset of) the one from the 'natural' part of the RBSF area and corresponds to landslide distribution in the year 2000.

Format

a data.frame with point samples of landslide and non-landslide locations in a study area in the Andes of southern Ecuador.

References

Muenchow, J., Brenning, A., Richter, M., 2012. Geomorphic process rates of landslides along a humidity gradient in the tropical Andes. *Geomorphology*, 139-140: 271-284.

Brenning, A., 2005. Spatial prediction models for landslide hazards: review, comparison and evaluation. *Natural Hazards and Earth System Sciences*, 5(6): 853-862.

subsetTask	<i>Subset data in task.</i>
------------	-----------------------------

Description

See title.

Usage

```
subsetTask(task, subset = NULL, features)
```

Arguments

task	(Task) The task.
subset	(integer logical NULL) Selected cases. Either a logical or an index vector. By default NULL if all observations are used.
features	(character integer logical) Vector of selected inputs. You can either pass a character vector with the feature names, a vector of indices, or a logical vector. In case of an index vector each element denotes the position of the feature name returned by getTaskFeatureNames . Note that the target feature is always included in the resulting task, you should not pass it here. Default is to use all features.

Value

([Task](#)). Task with subsetted data.

See Also

Other task: [getTaskClassLevels\(\)](#), [getTaskCosts\(\)](#), [getTaskData\(\)](#), [getTaskDesc\(\)](#), [getTaskFeatureNames\(\)](#), [getTaskFormula\(\)](#), [getTaskId\(\)](#), [getTaskNFeats\(\)](#), [getTaskSize\(\)](#), [getTaskTargetNames\(\)](#), [getTaskTargets\(\)](#), [getTaskType\(\)](#)

Examples

```
task = makeClassifTask(data = iris, target = "Species")
subsetTask(task, subset = 1:100)
```

summarizeColumns	<i>Summarize columns of data.frame or task.</i>
------------------	---

Description

Summarizes a `data.frame`, somewhat differently than the normal `summary` function of R. The function is mainly useful as a basic EDA tool on `data.frames` before they are converted to tasks, but can be used on tasks as well.

Columns can be of type numeric, integer, logical, factor, or character. Characters and logicals will be treated as factors.

Usage

```
summarizeColumns(obj)
```

Arguments

obj	(data.frame Task) Input data.
-----	--

Value

([data.frame](#)). With columns:

name	Name of column.
type	Data type of column.
na	Number of NAs in column.
disp	Measure of dispersion, for numerics and integers <code>sd</code> is used, for categorical columns the qualitative variation.
mean	Mean value of column, NA for categorical columns.
median	Median value of column, NA for categorical columns.
mad	MAD of column, NA for categorical columns.
min	Minimal value of column, for categorical columns the size of the smallest category.
max	Maximal value of column, for categorical columns the size of the largest category.
nlevs	For categorical columns, the number of factor levels, NA else.

See Also

Other `eda_and_preprocess`: [capLargeValues\(\)](#), [createDummyFeatures\(\)](#), [dropFeatures\(\)](#), [mergeSmallFactorLevels\(\)](#), [normalizeFeatures\(\)](#), [removeConstantFeatures\(\)](#), [summarizeLevels\(\)](#)

Examples

```
summarizeColumns(iris)
```

summarizeLevels	<i>Summarizes factors of a data.frame by tabling them.</i>
-----------------	--

Description

Characters and logicals will be treated as factors.

Usage

```
summarizeLevels(obj, cols = NULL)
```

Arguments

obj	(data.frame Task) Input data.
cols	(character) Restrict result to columns in cols. Default is all factor, character and logical columns of obj.

Value

([list](#)). Named list of tables.

See Also

Other `eda_and_preprocess`: [capLargeValues\(\)](#), [createDummyFeatures\(\)](#), [dropFeatures\(\)](#), [mergeSmallFactorLevels\(\)](#), [normalizeFeatures\(\)](#), [removeConstantFeatures\(\)](#), [summarizeColumns\(\)](#)

Examples

```
summarizeLevels(iris)
```

Task	<i>Create a classification, regression, survival, cluster, cost-sensitive classification or multilabel task.</i>
------	--

Description

The task encapsulates the data and specifies - through its subclasses - the type of the task. It also contains a description object detailing further aspects of the data.

Useful operators are:

- [getTaskFormula](#),
- [getTaskFeatureNames](#),
- [getTaskData](#),

- [getTaskTargets](#), and
- [subsetTask](#).

Object members:

env (environment) Environment where data for the task are stored. Use [getTaskData](#) in order to access it.

weights (**numeric**) See argument. NULL if not present.

blocking (**factor**) See argument. NULL if not present.

task.desc (**TaskDesc**) Encapsulates further information about the task.

Functional data can be added to a task via matrix columns. For more information refer to [make-FunctionalData](#).

Arguments

<code>id</code>	(character(1)) Id string for object. Default is the name of the R variable passed to data.
<code>data</code>	(data.frame) A data frame containing the features and target variable(s).
<code>target</code>	(character(1) character(2) character(n.classes)) Name(s) of the target variable(s). For survival analysis these are the names of the survival time and event columns, so it has length 2. For multilabel classification it contains the names of the logical columns that encode whether a label is present or not and its length corresponds to the number of classes.
<code>costs</code>	(data.frame) A numeric matrix or data frame containing the costs of misclassification. We assume the general case of observation specific costs. This means we have n rows, corresponding to the observations, in the same order as data. The columns correspond to classes and their names are the class labels (if unnamed we use y1 to yk as labels). Each entry (i,j) of the matrix specifies the cost of predicting class j for observation i.
<code>weights</code>	(numeric) Optional, non-negative case weight vector to be used during fitting. Cannot be set for cost-sensitive learning. Default is NULL which means no (= equal) weights.
<code>blocking</code>	(factor) An optional factor of the same length as the number of observations. Observations with the same blocking level “belong together”. Specifically, they are either put all in the training or the test set during a resampling iteration. Default is NULL which means no blocking.
<code>positive</code>	(character(1)) Positive class for binary classification (otherwise ignored and set to NA). Default is the first factor level of the target attribute.
<code>fixup.data</code>	(character(1)) Should some basic cleaning up of data be performed? Currently this means

	removing empty factor levels for the columns. Possible choices are: “no” = Don’t do it. “warn” = Do it but warn about it. “quiet” = Do it but keep silent. Default is “warn”.
check.data	(logical(1)) Should sanity of data be checked initially at task creation? You should have good reasons to turn this off (one might be speed). Default is TRUE.
coordinates	(data.frame) Coordinates of a spatial data set that will be used for spatial partitioning of the data in a spatial cross-validation resampling setting. Coordinates have to be numeric values. Provided data.frame needs to have the same number of rows as data and consist of at least two dimensions.

Value

[Task](#).

See Also

[ClassifTask](#) [ClusterTask](#) [CostSensTask](#) [MultilabelTask](#) [RegrTask](#) [SurvTask](#)

Examples

```
if (requireNamespace("mlbench")) {
  library(mlbench)
  data(BostonHousing)
  data(Ionosphere)

  makeClassifTask(data = iris, target = "Species")
  makeRegrTask(data = BostonHousing, target = "medv")
  # an example of a classification task with more than those standard arguments:
  blocking = factor(c(rep(1, 51), rep(2, 300)))
  makeClassifTask(id = "myIonosphere", data = Ionosphere, target = "Class",
    positive = "good", blocking = blocking)
  makeClusterTask(data = iris[, -5L])
}
```

TaskDesc

Description object for task.

Description

Description object for task, encapsulates basic properties of the task without having to store the complete data set.

Details

Object members:

id (character(1)) Id string of task.

type (character(1)) Type of task, “`classif`” for classification, “`regr`” for regression, “`surv`” for survival and “`cluster`” for cluster analysis, “`costsens`” for cost-sensitive classification, and “`multilabel`” for multilabel classification.

target (character(0) | character(1) | character(2) | character(n.classes)) Name(s) of the target variable(s). For “`surv`” these are the names of the survival time and event columns, so it has length 2. For “`costsens`” it has length 0, as there is no target column, but a cost matrix instead. For “`multilabel`” these are the names of logical columns that indicate whether a class label is present and the number of target variables corresponds to the number of classes.

size (integer(1)) Number of cases in data set.

n.feat (integer(2)) Number of features, named vector with entries: “`numerics`”, “`factors`”, “`ordered`”, “`functionals`”.

has.missings (logical(1)) Are missing values present?

has.weights (logical(1)) Are weights specified for each observation?

has.blocking (logical(1)) Is a blocking factor for cases available in the task?

class.levels (**character**) All possible classes. Only present for “`classif`”, “`costsens`”, and “`multilabel`”.

positive (character(1)) Positive class label for binary classification. Only present for “`classif`”, NA for multiclass.

negative (character(1)) Negative class label for binary classification. Only present for “`classif`”, NA for multiclass.

train

Train a learning algorithm.

Description

Given a [Task](#), creates a model for the learning machine which can be used for predictions on new data.

Usage

```
train(learner, task, subset = NULL, weights = NULL)
```

Arguments

learner ([Learner](#) | character(1))
The learner. If you pass a string the learner will be created via [makeLearner](#).

task ([Task](#))
The task.

subset	(integer logical NULL) Selected cases. Either a logical or an index vector. By default NULL if all observations are used.
weights	(numeric) Optional, non-negative case weight vector to be used during fitting. If given, must be of same length as subset and in corresponding order. By default NULL which means no weights are used unless specified in the task (Task). Weights from the task will be overwritten.

Value

([WrappedModel](#)).

See Also

[predict.WrappedModel](#)

Examples

```
training.set = sample(seq_len(nrow(iris)), nrow(iris) / 2)

## use linear discriminant analysis to classify iris data
task = makeClassifTask(data = iris, target = "Species")
learner = makeLearner("classif.lda", method = "mle")
mod = train(learner, task, subset = training.set)
print(mod)

## use random forest to classify iris data
task = makeClassifTask(data = iris, target = "Species")
learner = makeLearner("classif.rpart", minsplit = 7, predict.type = "prob")
mod = train(learner, task, subset = training.set)
print(mod)
```

trainLearner

Train an R learner.

Description

Mainly for internal use. Trains a wrapped learner on a given training set. You have to implement this method if you want to add another learner to this package.

Usage

```
trainLearner(.learner, .task, .subset, .weights = NULL, ...)
```

Arguments

<code>.learner</code>	(RLearner) Wrapped learner.
<code>.task</code>	(Task) Task to train learner on.
<code>.subset</code>	(integer) Subset of cases for training set, index the task with this. You probably want to use getTaskData for this purpose.
<code>.weights</code>	(numeric) Weights for each observation.
<code>...</code>	(any) Additional (hyper)parameters, which need to be passed to the underlying train function.

Details

Your implementation must adhere to the following: The model must be fitted on the subset of `.task` given by `.subset`. All parameters in `...` must be passed to the underlying training function.

Value

([any](#)). Model of the underlying learner.

TuneControl	<i>Control object for tuning</i>
-------------	----------------------------------

Description

General tune control object.

Arguments

<code>same.resampling.instance</code>	(logical(1)) Should the same resampling instance be used for all evaluations to reduce variance? Default is TRUE.
<code>impute.val</code>	(numeric) If something goes wrong during optimization (e.g. the learner crashes), this value is fed back to the tuner, so the tuning algorithm does not abort. It is not stored in the optimization path, an NA and a corresponding error message are logged instead. Note that this value is later multiplied by -1 for maximization measures internally, so you need to enter a larger positive value for maximization here as well. Default is the worst obtainable value of the performance measure you optimize for when you aggregate by mean value, or Inf instead. For multi-criteria optimization pass a vector of imputation values, one for each of your measures, in the same order as your measures.

start	(list) Named list of initial parameter values.
tune.threshold	(logical(1)) Should the threshold be tuned for the measure at hand, after each hyperparameter evaluation, via <code>tuneThreshold</code> ? Only works for classification if the predict type is “prob”. Default is FALSE.
tune.threshold.args	(list) Further arguments for threshold tuning that are passed down to <code>tuneThreshold</code> . Default is none.
log.fun	(function character(1)) Function used for logging. If set to “default” (the default), the evaluated design points, the resulting performances, and the runtime will be reported. If set to “memory” the memory usage for each evaluation will also be displayed, with character(1) small increase in run time. Otherwise character(1) function with arguments learner, resampling, measures, par.set, control, opt.path, dob, x, y, remove.nas, stage and prev.stage is expected. The default displays the performance measures, the time needed for evaluating, the currently used memory and the max memory ever used before (the latter two both taken from <code>gc</code>). See the implementation for details.
final.dw.perc	(boolean) If a Learner wrapped by a <code>makeDownsampleWrapper</code> is used, you can define the value of <code>dw.perc</code> which is used to train the Learner with the final parameter setting found by the tuning. Default is NULL which will not change anything.
...	(any) Further control parameters passed to the control arguments of <code>cmaes::cma_es</code> or <code>GenSA::GenSA</code> , as well as towards the <code>tunerConfig</code> argument of <code>irace::irace</code> .

See Also

Other tune: `getNestedTuneResultsOptPathDf()`, `getNestedTuneResultsX()`, `getResamplingIndices()`, `getTuneResult()`, `makeModelMultiplexerParamSet()`, `makeModelMultiplexer()`, `makeTuneControlCMAES()`, `makeTuneControlDesign()`, `makeTuneControlGenSA()`, `makeTuneControlGrid()`, `makeTuneControlIrace()`, `makeTuneControlMBO()`, `makeTuneControlRandom()`, `makeTuneWrapper()`, `tuneParams()`, `tuneThreshold()`

TuneMultiCritControl *Create control structures for multi-criteria tuning.*

Description

The following tuners are available:

makeTuneMultiCritControlGrid Grid search. All kinds of parameter types can be handled. You can either use their correct param type and resolution, or discretize them yourself by always using `ParamHelpers::makeDiscreteParam` in the `par.set` passed to `tuneParams`.

- makeTuneMultiCritControlRandom** Random search. All kinds of parameter types can be handled.
- makeTuneMultiCritControlNSGA2** Evolutionary method [mco::nsga2](#). Can handle numeric(vector) and integer(vector) hyperparameters, but no dependencies. For integers the internally proposed numeric values are automatically rounded.
- makeTuneMultiCritControlMBO** Model-based/ Bayesian optimization. All kinds of parameter types can be handled.

Usage

```
makeTuneMultiCritControlGrid(
  same.resampling.instance = TRUE,
  resolution = 10L,
  log.fun = "default",
  final.dw.perc = NULL,
  budget = NULL
)

makeTuneMultiCritControlMBO(
  n.objectives = mbo.control$n.objectives,
  same.resampling.instance = TRUE,
  impute.val = NULL,
  learner = NULL,
  mbo.control = NULL,
  tune.threshold = FALSE,
  tune.threshold.args = list(),
  continue = FALSE,
  log.fun = "default",
  final.dw.perc = NULL,
  budget = NULL,
  mbo.design = NULL
)

makeTuneMultiCritControlNSGA2(
  same.resampling.instance = TRUE,
  impute.val = NULL,
  log.fun = "default",
  final.dw.perc = NULL,
  budget = NULL,
  ...
)

makeTuneMultiCritControlRandom(
  same.resampling.instance = TRUE,
  maxit = 100L,
  log.fun = "default",
  final.dw.perc = NULL,
  budget = NULL
)
```

)

Arguments

same.resampling.instance	(logical(1)) Should the same resampling instance be used for all evaluations to reduce variance? Default is TRUE.
resolution	(integer) Resolution of the grid for each numeric/integer parameter in par.set. For vector parameters, it is the resolution per dimension. Either pass one resolution for all parameters, or a named vector. See ParamHelpers::generateGridDesign . Default is 10.
log.fun	(function character(1)) Function used for logging. If set to “default” (the default), the evaluated design points, the resulting performances, and the runtime will be reported. If set to “memory” the memory usage for each evaluation will also be displayed, with character(1) small increase in run time. Otherwise character(1) function with arguments learner, resampling, measures, par.set, control, opt.path, dob, x, y, remove.nas, stage and prev.stage is expected. The default displays the performance measures, the time needed for evaluating, the currently used memory and the max memory ever used before (the latter two both taken from gc). See the implementation for details.
final.dw.perc	(boolean) If a Learner wrapped by a makeDownsampleWrapper is used, you can define the value of dw.perc which is used to train the Learner with the final parameter setting found by the tuning. Default is NULL which will not change anything.
budget	(integer(1)) Maximum budget for tuning. This value restricts the number of function evaluations. In case of makeTuneMultiCritControlGrid this number must be identical to the size of the grid. For makeTuneMultiCritControlRandom the budget equals the number of iterations (maxit) performed by the random search algorithm. In case of makeTuneMultiCritControlNSGA2 the budget corresponds to the product of the maximum number of generations (max(generations)) + 1 (for the initial population) and the size of the population (popsize). For makeTuneMultiCritControlMBO the budget equals the number of objective function evaluations, i.e. the number of MBO iterations + the size of the initial design. If not NULL, this will overwrite existing stopping conditions in mbo.control.
n.objectives	(integer(1)) Number of objectives, i.e. number of Measures to optimize.
impute.val	(numeric) If something goes wrong during optimization (e.g. the learner crashes), this value is fed back to the tuner, so the tuning algorithm does not abort. It is not stored in the optimization path, an NA and a corresponding error message are logged instead. Note that this value is later multiplied by -1 for maximization measures internally, so you need to enter a larger positive value for maximization

here as well. Default is the worst obtainable value of the performance measure you optimize for when you aggregate by mean value, or Inf instead. For multi-criteria optimization pass a vector of imputation values, one for each of your measures, in the same order as your measures.

learner	(Learner NULL) The surrogate learner: A regression learner to model performance landscape. For the default, NULL, <code>mlrMBO</code> will automatically create a suitable learner based on the rules described in <code>mlrMBO::makeMBOLEarner</code> .
mbo.control	(<code>mlrMBO::MBOControl</code> NULL) Control object for model-based optimization tuning. For the default, NULL, the control object will be created with all the defaults as described in <code>mlrMBO::makeMBOControl</code> .
tune.threshold	(<code>logical(1)</code>) Should the threshold be tuned for the measure at hand, after each hyperparameter evaluation, via <code>tuneThreshold</code> ? Only works for classification if the predict type is “prob”. Default is FALSE.
tune.threshold.args	(<code>list</code>) Further arguments for threshold tuning that are passed down to <code>tuneThreshold</code> . Default is none.
continue	(<code>logical(1)</code>) Resume calculation from previous run using <code>mlrMBO::mboContinue</code> ? Requires “save.file.path” to be set. Note that the <code>ParamHelpers::OptPath</code> in the <code>mlrMBO::OptResult</code> will only include the evaluations after the continuation. The complete <code>OptPath</code> will be found in the slot <code>\$mbo.result\$opt.path</code> .
mbo.design	(<code>data.frame</code> NULL) Initial design as data frame. If the parameters have corresponding trafo functions, the design must not be transformed before it is passed! For the default, NULL, a default design is created like described in <code>mlrMBO::mbo</code> .
...	(<code>any</code>) Further control parameters passed to the control arguments of <code>cmaes::cma_es</code> or <code>GenSA::GenSA</code> , as well as towards the <code>tunerConfig</code> argument of <code>irace::irace</code> .
maxit	(<code>integer(1)</code>) Number of iterations for random search. Default is 100.

Value

(`TuneMultiCritControl`). The specific subclass is one of `TuneMultiCritControlGrid`, `TuneMultiCritControlRandom`, `TuneMultiCritControlNSGA2`, `TuneMultiCritControlMBO`.

See Also

Other `tune_multicrit`: `plotTuneMultiCritResult()`, `tuneParamsMultiCrit()`

TuneMultiCritResult *Result of multi-criteria tuning.*

Description

Container for results of hyperparameter tuning. Contains the obtained pareto set and front and the optimization path which lead there.

Object members:

learner (**Learner**) Learner that was optimized.

control (**TuneControl**) Control object from tuning.

x (**list**) List of lists of non-dominated hyperparameter settings in pareto set. Note that when you have trafos on some of your params, x will always be on the TRANSFORMED scale so you directly use it.

y (**matrix**) Pareto front for x.

threshold Currently NULL.

opt.path (**ParamHelpers::OptPath**) Optimization path which lead to x. Note that when you have trafos on some of your params, the opt.path always contains the UNTRANSFORMED values on the original scale. You can simply call `trafoOptPath(opt.path)` to transform them, or, `as.data.frame{trafoOptPath(opt.path)}`

ind (**integer(n)**) Indices of Pareto optimal params in `opt.path`.

measures [(**list of Measure**)] Performance measures.

tuneParams *Hyperparameter tuning.*

Description

Optimizes the hyperparameters of a learner. Allows for different optimization methods, such as grid search, evolutionary strategies, iterated F-race, etc. You can select such an algorithm (and its settings) by passing a corresponding control object. For a complete list of implemented algorithms look at [TuneControl](#).

Multi-criteria tuning can be done with [tuneParamsMultiCrit](#).

Usage

```
tuneParams(
  learner,
  task,
  resampling,
  measures,
  par.set,
  control,
  show.info = getMlrOption("show.info"),
  resample.fun = resample
)
```

Arguments

learner	(Learner character(1)) The learner. If you pass a string the learner will be created via makeLearner .
task	(Task) The task.
resampling	(ResampleInstance ResampleDesc) Resampling strategy to evaluate points in hyperparameter space. If you pass a description, it is instantiated once at the beginning by default, so all points are evaluated on the same training/test sets. If you want to change that behavior, look at TuneControl .
measures	(list of Measure Measure) Performance measures to evaluate. The first measure, aggregated by the first aggregation function is optimized, others are simply evaluated. Default is the default measure for the task, see here getDefaultMeasure .
par.set	(ParamHelpers::ParamSet) Collection of parameters and their constraints for optimization. Dependent parameters with a <code>requires</code> field must use quote and not expression to define it.
control	(TuneControl) Control object for search method. Also selects the optimization algorithm for tuning.
show.info	(logical(1)) Print verbose output on console? Default is set via configureMlr .
resample.fun	(closure) The function to use for resampling. Defaults to resample . If a user-given function is to be used instead, it should take the arguments “learner”, “task”, “resampling”, “measures”, and “show.info”; see resample . Within this function, it is easiest to call resample and possibly modify the result. However, it is possible to return a list with only the following essential slots: the “aggr” slot for general tuning, additionally the “pred” slot if threshold tuning is performed (see TuneControl), and the “err.msgs” and “err.dumps” slots for error reporting. This parameter must be the default when mbo tuning is performed.

Value

([TuneResult](#)).

Note

If you would like to include results from the training data set, make sure to appropriately adjust the resampling strategy and the aggregation for the measure. See example code below.

See Also

[generateHyperParsEffectData](#)

Other tune: [TuneControl](#), [getNestedTuneResultsOptPathDf\(\)](#), [getNestedTuneResultsX\(\)](#), [getResamplingIndices\(\)](#), [getTuneResult\(\)](#), [makeModelMultiplexerParamSet\(\)](#), [makeModelMultiplexer\(\)](#),

```

makeTuneControlCMAES(), makeTuneControlDesign(), makeTuneControlGenSA(), makeTuneControlGrid(),
makeTuneControlIrace(), makeTuneControlMBO(), makeTuneControlRandom(), makeTuneWrapper(),
tuneThreshold()

```

Examples

```

set.seed(123)
# a grid search for an SVM (with a tiny number of points...)
# note how easily we can optimize on a log-scale
ps = makeParamSet(
  makeNumericParam("C", lower = -12, upper = 12, trafo = function(x) 2^x),
  makeNumericParam("sigma", lower = -12, upper = 12, trafo = function(x) 2^x)
)
ctrl = makeTuneControlGrid(resolution = 2L)
rdesc = makeResampleDesc("CV", iters = 2L)
res = tuneParams("classif.ksvm", iris.task, rdesc, par.set = ps, control = ctrl)
print(res)
# access data for all evaluated points
df = as.data.frame(res$opt.path)
df1 = as.data.frame(res$opt.path, trafo = TRUE)
print(head(df[, -ncol(df)]))
print(head(df1[, -ncol(df)]))
# access data for all evaluated points - alternative
df2 = generateHyperParsEffectData(res)
df3 = generateHyperParsEffectData(res, trafo = TRUE)
print(head(df2$data[, -ncol(df2$data)]))
print(head(df3$data[, -ncol(df3$data)]))
## Not run:
# we optimize the SVM over 3 kernels simultaneously
# note how we use dependent params (requires = ...) and iterated F-racing here
ps = makeParamSet(
  makeNumericParam("C", lower = -12, upper = 12, trafo = function(x) 2^x),
  makeDiscreteParam("kernel", values = c("vanilladot", "polydot", "rbfdot")),
  makeNumericParam("sigma", lower = -12, upper = 12, trafo = function(x) 2^x,
    requires = quote(kernel == "rbfdot")),
  makeIntegerParam("degree", lower = 2L, upper = 5L,
    requires = quote(kernel == "polydot"))
)
print(ps)
ctrl = makeTuneControlIrace(maxExperiments = 5, nbIterations = 1, minNbSurvival = 1)
rdesc = makeResampleDesc("Holdout")
res = tuneParams("classif.ksvm", iris.task, rdesc, par.set = ps, control = ctrl)
print(res)
df = as.data.frame(res$opt.path)
print(head(df[, -ncol(df)]))

# include the training set performance as well
rdesc = makeResampleDesc("Holdout", predict = "both")
res = tuneParams("classif.ksvm", iris.task, rdesc, par.set = ps,
  control = ctrl, measures = list(mmce, setAggregation(mmce, train.mean)))
print(res)
df2 = as.data.frame(res$opt.path)
print(head(df2[, -ncol(df2)]))

```

```
## End(Not run)
```

```
tuneParamsMultiCrit Hyperparameter tuning for multiple measures at once.
```

Description

Optimizes the hyperparameters of a learner in a multi-criteria fashion. Allows for different optimization methods, such as grid search, evolutionary strategies, etc. You can select such an algorithm (and its settings) by passing a corresponding control object. For a complete list of implemented algorithms look at [TuneMultiCritControl](#).

Usage

```
tuneParamsMultiCrit(
  learner,
  task,
  resampling,
  measures,
  par.set,
  control,
  show.info = getMlrOption("show.info"),
  resample.fun = resample
)
```

Arguments

learner	(Learner character(1)) The learner. If you pass a string the learner will be created via makeLearner .
task	(Task) The task.
resampling	(ResampleInstance ResampleDesc) Resampling strategy to evaluate points in hyperparameter space. If you pass a description, it is instantiated once at the beginning by default, so all points are evaluated on the same training/test sets. If you want to change that behavior, look at TuneMultiCritControl .
measures	[list of Measure] Performance measures to optimize simultaneously.
par.set	(ParamHelpers::ParamSet) Collection of parameters and their constraints for optimization. Dependent parameters with a <code>requires</code> field must use quote and not expression to define it.
control	(TuneMultiCritControl) Control object for search method. Also selects the optimization algorithm for tuning.

`show.info` (logical(1))
Print verbose output on console? Default is set via [configureMlr](#).

`resample.fun` (closure)
The function to use for resampling. Defaults to [resample](#) and should take the same arguments as, and return the same result type as, [resample](#).

Value

([TuneMultiCritResult](#)).

See Also

Other `tune_multicrit`: [TuneMultiCritControl](#), [plotTuneMultiCritResult\(\)](#)

Examples

```
# multi-criteria optimization of (tpr, fpr) with NGS-II
lrn = makeLearner("classif.ksvm")
rdesc = makeResampleDesc("Holdout")
ps = makeParamSet(
  makeNumericParam("C", lower = -12, upper = 12, trafo = function(x) 2^x),
  makeNumericParam("sigma", lower = -12, upper = 12, trafo = function(x) 2^x)
)
ctrl = makeTuneMultiCritControlNSGA2(popsize = 4L, generations = 1L)
res = tuneParamsMultiCrit(lrn, sonar.task, rdesc, par.set = ps,
  measures = list(tpr, fpr), control = ctrl)
plotTuneMultiCritResult(res, path = TRUE)
```

TuneResult

Result of tuning.

Description

Container for results of hyperparameter tuning. Contains the obtained point in search space, its performance values and the optimization path which lead there.

Object members:

learner ([Learner](#)) Learner that was optimized.

control ([TuneControl](#)) Control object from tuning.

x ([list](#)) Named list of hyperparameter values identified as optimal. Note that when you have trafos on some of your params, x will always be on the TRANSFORMED scale so you directly use it.

y ([numeric](#)) Performance values for optimal x.

threshold ([numeric](#)) Vector of finally found and used thresholds if `tune.threshold` was enabled in [TuneControl](#), otherwise not present and hence NULL.

opt.path (ParamHelpers::OptPath) Optimization path which lead to x. Note that when you have trafo's on some of your params, the opt.path always contains the UNTRANSFORMED values on the original scale. You can simply call `trafoOptPath(opt.path)` to transform them, or, `as.data.frame(trafoOptPath(opt.path))`. If mlr option `on.error.dump` is TRUE, OptPath will have a `.dump` object in its extra column which contains error dump traces from failed optimization evaluations. It can be accessed by `getOptPathEl(opt.path)$extra$.dump`.

tuneThreshold	<i>Tune prediction threshold.</i>
---------------	-----------------------------------

Description

Optimizes the threshold of predictions based on probabilities. Works for classification and multi-label tasks. Uses [BBmisc::optimizeSubInts](#) for normal binary class problems and [GenSA::GenSA](#) for multiclass and multilabel problems.

Usage

```
tuneThreshold(pred, measure, task, model, nsub = 20L, control = list())
```

Arguments

pred	(Prediction) Prediction object.
measure	(Measure) Performance measure to optimize. Default is the default measure for the task.
task	(Task) Learning task. Rarely needed, only when required for the performance measure.
model	(WrappedModel) Fitted model. Rarely needed, only when required for the performance measure.
nsub	(<code>integer(1)</code>) Passed to BBmisc::optimizeSubInts for 2class problems. Default is 20.
control	(<code>list</code>) Control object for GenSA::GenSA when used. Default is empty list.

Value

(`list`). A named list with with the following components: `th` is the optimal threshold, `perf` the performance value.

See Also

Other tune: [TuneControl](#), [getNestedTuneResultsOptPathDf\(\)](#), [getNestedTuneResultsX\(\)](#), [getResamplingIndices\(\)](#), [getTuneResult\(\)](#), [makeModelMultiplexerParamSet\(\)](#), [makeModelMultiplexer\(\)](#), [makeTuneControlCMAES\(\)](#), [makeTuneControlDesign\(\)](#), [makeTuneControlGenSA\(\)](#), [makeTuneControlGrid\(\)](#), [makeTuneControlIrace\(\)](#), [makeTuneControlMBO\(\)](#), [makeTuneControlRandom\(\)](#), [makeTuneWrapper\(\)](#), [tuneParams\(\)](#)

`wpbc.task`*Wisconsin Prognostic Breast Cancer (WPBC) survival task.*

Description

Contains the task (`wpbc.task`).

References

See [TH.data::wpbc](#). Incomplete cases have been removed from the task.

`yeast.task`*Yeast multilabel classification task.*

Description

Contains the task (`yeast.task`).

Source

<https://archive.ics.uci.edu/ml/datasets/Yeast> (In long instead of wide format)

References

Elisseeff, A., & Weston, J. (2001): A kernel method for multi-labelled classification. In Advances in neural information processing systems (pp. 681-687).

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