# Package 'stablelearner'

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accuracy

Prediction Accuracy from Stability Assessment Results

### Description

Function to compute the prediction accuracy from an object of class "stablelearner" or "stablelearnerList" as a parallel to the similarity values estimated by stability in each iteration of the stability assessment procedure.

### Usage

```
accuracy(x, measure = "kappa", na.action = na.exclude,
  applyfun = NULL, cores = NULL)
```

### Arguments

Χ	an object of class "stablelearner" or "stablelearnerList".
measure	a character string (or a vector of character strings). Name(s) of the measure(s) used to compute accuracy. Currently implemented measures are "diag" = percentage of observations on the main diagonal of a confusion matrix, "kappa" = "diag" corrected for agreement by chance (default), "rand" = Rand index, and "crand" = Rand index corrected for agreemend by chance (see also classAgreement).
na.action	a function which indicates what should happen to the predictions of the results containing NAs. The default function is na.exclude.
applyfun	a lapply-like function. The default is to use lapply unless cores is specified in which case mclapply is used (for multicore computations on platforms that support these).
cores	integer. The number of cores to use in multicore computations using mclapply (see above).

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#### **Details**

This function can be used to compute prediction accuracy after the stability was estimated using stability.

#### Value

A matrix of size 2\*B times length(measure) containing prediction accuracy values of the learners trained during the stability assessment procedure.

#### See Also

```
stability
```

### **Examples**

```
library("partykit")
res <- ctree(Species ~ ., data = iris)
stab <- stability(res)
accuracy(stab)</pre>
```

addLearner

Add Learners to LearnerList

### Description

The function can be used to add new learner to LearnerList in the current R session.

### Usage

```
addLearner(x)
```

### Arguments

Х

a list containing all required information to define a new learner (see Details below).

### Details

The function can be used to add new learners to LearnerList in the current R session. The function expects a list of four elements including the name of the learners object class, the name of the package where the class and the fitting method is implemented, the name of the method and a prediction function that predicts class probabilities (in the classification case) or numeric values (in the regression case) and takes the arguments x (the fitted model object), newdata a data. frame containing the predictions of the observations in the evaluation sample and yclass a character string specifying the type of the response variable ("numeric", "factor", etc.). The elements in the list should be named class, package, method and predfun.

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### See Also

LearnerList

### **Examples**

```
newlearner <- list(
  class = "svm",
  package = "e1071",
  method = "Support Vector Machine",
  predict = function(x, newdata, yclass = NULL) {
    if(match(yclass, c("ordered", "factor"))) {
      attr(predict(x, newdata = newdata, probability = TRUE), "probabilities")
    } else {
      predict(x, newdata = newdata)
    }
})
addLearner(newlearner)</pre>
```

bootstrap

Sampler Infrastructure for Stability Assessment

### Description

Sampler objects that provide objects with functionality used by stabletree to generate resampled datasets.

### Usage

```
bootstrap(B = 500, v = 1)
subsampling(B = 500, v = 0.632)
samplesplitting(k = 5)
jackknife(d = 1, maxrep = 5000)
splithalf(B = 500)
```

### **Arguments**

В	An integer value specifying the number of resampled datasets.
k	An integer value specifying the number of folds in sample-splitting.
d	An integer value specifying the number of observations left out in jackknife.
maxrep	An integer value specifying the maximum number of resampled datasets allowed, when using jackknife.
V	A numeric value between 0 and 1 specifying the fraction of observations in each subsample.

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### **Details**

The sampler functions provide objects that include functionality to generate resampled datasets used by stabletree.

The bootstrap function provides an object that can be used to generate B bootstrap samples by sampling from n observations with replacement.

The subsampling function provides an object that can be used to generate B subsamples by sampling from floor(v\*n) observations without replacement.

The samplesplitting function provides an object that can be used to generate k-folds from n observations.

The jackknife function provides an object that can be used to generate all datasets necessary to perform leave-k-out jackknife sampling from n observations. The number of datasets is limited by maxrep to prevent unintended CPU or memory overload by accidently choosing too large values for k.

The splithalf function provides an object that can be used to generate B subsamples by sampling from floor(0.5\*n) observations without replacement. When used to implement the "splithalf" resampling strategy for measuring the stability of a result via the stability function, the matrix containing the complement learning samples is generated automatically by stability.

#### See Also

```
stabletree, stability
```

```
set.seed(0)
## bootstrap sampler
s <- bootstrap(3)</pre>
s$sampler(10)
## subsampling
s \leftarrow subsampling(3, v = 0.6)
s$sampler(10)
## 5-fold sample-splitting
s <- samplesplitting(5)</pre>
s$sampler(10)
## jackknife
s <- jackknife(d = 1)
s$sampler(10)
## splithaf
s <- splithalf(3)
s$sampler(10)
```

```
boxplot.stablelearnerList
```

Illustrate Results from Stability Assessment

### **Description**

Illustrates the results from stability assessments performed by stability using boxplots.

### Usage

```
## S3 method for class 'stablelearnerList'
boxplot(x, ..., main = NULL, xlab = NULL, ylab = NULL, reverse = TRUE)
## S3 method for class 'stablelearner'
boxplot(x, ...)
```

### **Arguments**

Х	an object of class "stablelearnerList" to be illustrated.
	Arguments passed to boxplot.
main	a character specifying the title. By default set to NULL.
xlab	a character specifying the title for the x axis. By default set to "Learner".
ylab	a character specifying the title for the y axis. By default set to NULL.
reverse	logical. If reverse = TRUE (default), the similarity values are transformed (reversed) such that higher values indicate a higher stability.

### See Also

```
stability, summary.stablelearnerList
```

```
library("partykit")
r1 <- ctree(Species ~ ., data = iris)
library("rpart")
r2 <- rpart(Species ~ ., data = iris)
stab <- stability(r1, r2, names = c("ctree", "rpart"))
boxplot(stab)</pre>
```

dgp\_twoclass 7

dgp_twoclass Data-Ggnerating Function for Two-Class Problem
---

### Description

Data-generating function to generate artificial data sets of a classification problem with two response classes, denoted as "A" and "B".

### Usage

```
dgp_twoclass(n = 100, p = 4, noise = 16, rho = 0,
b0 = 0, b = rep(1, p), fx = identity)
```

### Arguments

n	integer. Number of observations. The default is 100.
p	integer. Number of signal predictors. The default is 4.
noise	integer. Number of noise predictors. The default is 16.
rho	numeric value between -1 and 1 specifying the correlation between the signal predictors. The correlation is given by rho^k, where k is an integer value given by toeplitz structure. The default is 0 (no correlation between predictors).
b0	numeric value. Baseline probability for class "B" on the logit scale. The default is $0$ .
b	numeric value. Slope parameter for the predictors on the logit scale. The default is 1 for all predictors.
fx	a function that is used to transform the predictors. The default is identity (equivalent to no transformation).

### Value

A data. frame including a column denoted as class that is a factor with two levels "A" and "B". All other columns represent the predictor variables (signal predictors followed by noise predictors) and are named by "x1", "x2", etc..

### See Also

```
stability
```

```
dgp_twoclass(n = 200, p = 6, noise = 4)
```

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getLearner

Get Learner Details from LearnerList

### Description

Function to get information available about a specific learner in LearnerList of the current R session.

### Usage

```
getLearner(x)
```

### **Arguments**

Х

a fitted model object.

### **Details**

The function returns the entry in LearnerList found for the class of the object submitted to the function.

### See Also

```
LearnerList, addLearner
```

### **Examples**

```
library("partykit")
m <- ctree(Species ~ ., data = iris)
getLearner(m)</pre>
```

LearnerList

List of Predefined Learners for Assessing Stability

### Description

The list contains details about several predefined learners that are required to assess the stability of results from statistical learning.

### Usage

LearnerList

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#### **Details**

```
Currently implemented learners are:
```

ctree conditional inference trees using ctree from partykit.

rpart recursive partitioning using rpart from rpart.

J48 recursive partitioning using J48 from RWeka.

**C5.0** recursive partitioning using C5.0 from **C50**.

tree recursive partitioning using tree from tree.

lda linear discriminant analysis using 1da from MASS.

lm linear models using 1m from stats.

**glm** generalized linear models using glm from **stats**.

Users can add new learners to LearnerList for the current R session, see addLearner.

#### See Also

addLearner

plot.stabletree

Visualizing Tree Stability Assessments

### **Description**

Visualizations of tree stability assessments carried out via stabletree.

#### Usage

```
## S3 method for class 'stabletree'
plot(x, select = order(colMeans(x$vs), decreasing = TRUE),
  type.breaks = "levels", col.breaks = "red", lty.breaks = "dashed",
  cex.breaks = 0.7, col.main = c("black", "gray50"), main.uline = TRUE,
 args.numeric = NULL, args.factor = NULL, args.ordered = NULL, main = NULL,
 original = TRUE, ...)
## S3 method for class 'stabletree'
barplot(height, main = "Variable selection frequencies",
 xlab = "", ylab = "", horiz = FALSE, col = gray.colors(2),
 names.arg = NULL, names.uline = TRUE, names.diag = TRUE,
 cex.names = 0.9,
 ylim = if (horiz) NULL else c(0, 100), xlim = if (horiz) c(0, 100) else NULL,
 original = TRUE, ...)
## S3 method for class 'stabletree'
image(x, main = "Variable selections",
 ylab = "Repetitions", xlab = "", col = gray.colors(2),
```

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```
names.arg = NULL, names.uline = TRUE, names.diag = TRUE,
cex.names = 0.9, xaxs = "i", yaxs = "i",
col.tree = 2, lty.tree = 2, xlim = c(0, length(x$vs0)), ylim = c(0, x$B),
original = TRUE, ...)
```

### **Arguments**

x, height an object of class stabletree.

original logical. Should the original tree information be highlighted?

An vector of integer or character values representing the number(s) or the name(s) of the variable(s) to be plotted. By default all variables are plotted. The numbers correspond to the ordering of all partitioning variables used in the call of

the fitted model object that was passed to stabletree.

type.breaks A character specifying the type of information added to the lines that repre-

sent the splits in the complete data tree. type.breaks = "levels" adds the level of the splits. type.breaks = "nodeids" adds the nodeid of the splits. type.breaks = "breaks" adds the cutpoint of the splits. type.breaks = "none"

suppresses any labeling.

col.breaks Coloring of the lines and the texts that represent the splits in the complete data

tree.

1ty.breaks Type of the lines that represent the splits in the complete data tree.

cex.breaks Size of the texts that represent the splits in the complete data tree.

col.main A vector of colors of length two. The first color is used for titles of variables

that are selected in the complete data tree. The second color is used for titles of

variables that are not selected in the complete data tree.

main.uline A logical value. If TRUE, variables selected in the complete data tree are under-

lined. If FALSE, underlining is suppressed.

args.numeric A list of arguments passed to the internal function that is used for plotting a

histogram of the cutpoints in numerical splits. breaks is passed to hist (see hist for details). Further arguments in the list are passed to plot.histogram

such as col, border etc.

args.factor A list of arguments passed to the internal function that is used for plotting an

image plot of the cutpoints in categorical splits. col (a vector of two colors) is used to illustrate categorical splits. Please note that the default colors are optimized for color vision deficiency. col.na defines the color used to highlight missing categories. Further arguments in the list are passed to the function

plot.default.

args.ordered A list of arguments passed to the internal function that is used for plotting a

barplot of the cutpoints in ordered categorical splits. All arguments in the list are passed to the function barplot.default, such as col, border etc.

... further arguments passed to plotting functions, especially for labeling and anno-

tation.

main, xlab, ylab

character. Annotations of axes and main title, respectively.

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horiz	A logical value. If FALSE, the bars are drawn vertically with the first bar to the left. If TRUE, bars are drawn horizontally with the first at the bottom.
col	A vector of colors of length two used for coloring in the barplot and image methods. The first color represents selected and the second color represents not selected partitioning variables.
names.arg	A vector of labels to be plotted below each bar (in case of barplot) or each column (in case of image). If the argument is omitted, then the variable names are taken as labels.
names.uline	A logical value. If TRUE, the labels representing variables that are used for splitting in the complete data tree, are underlined. If FALSE, labels are not underlined.
names.diag	A logical value (omitted if horiz = TRUE). If TRUE, the labels are drawn diagonally. If FALSE, labels are drawn horizontally.
cex.names	Expansion factor for labels.
xlim, ylim	The limits of the plot.
xaxs, yaxs	The style of axis interval calculation to be used (see par for details).
col.tree, lty.t	tree
	color and line type to indicate differences from the original tree that was resam-

### **Details**

• plot visualizes the variability of the cutpoints.

pled.

- barplot visualizes the variable selection frequency.
- image visualizes the combinations of variables selected.

### See Also

stabletree

```
## build a tree
library("partykit")
m <- ctree(Species ~ ., data = iris)
plot(m)

## investigate stability
set.seed(0)
s <- stabletree(m, B = 500)

## show variable selection proportions
## with different labels and different ordering
barplot(s)
barplot(s, cex.names = 0.8)
barplot(s, names.diag = FALSE)</pre>
```

```
barplot(s, names.arg = c("a", "b", "c", "d"))
barplot(s, names.uline = FALSE)
barplot(s, col = c("lightgreen", "darkred"))
barplot(s, horiz = TRUE)
## illustrate variable selections of replications
## with different labels and different ordering
image(s, cex.names = 0.8)
image(s, names.diag = FALSE)
image(s, names.arg = c("a", "b", "c", "d"))
image(s, names.uline = FALSE)
image(s, col = c("lightgreen", "darkred"))
## graphical cutpoint analysis, selecting variable by number and name
## with different numerical of break points
plot(s)
plot(s, select = 3)
plot(s, select = "Petal.Width")
plot(s, args.numeric = list(breaks = 40))
# change labels of splits in complete data tree
plot(s, select = 3, type.breaks = "levels")
plot(s, select = 3, type.breaks = "nodeids")
plot(s, select = 3, type.breaks = "breaks")
plot(s, select = 3, type.breaks = "none")
```

```
similarity_measures_classification
```

Similarity Measure Infrastructure for Stability Assessment with Ordinal Responses

### **Description**

Functions that provide objects with functionality used by stability to measure the similarity between the predictions of two results in classification problems.

### Usage

```
clagree()
ckappa()

bdist()
tvdist()
hdist()
jsdiv(base = 2)
```

#### **Arguments**

base

A positive or complex number: the base with respect to which logarithms are computed. Defaults to 2.

### **Details**

The similarity measure functions provide objects that include functionality used by stability to measure the similarity between the probability predictions of two results in classification problems.

The clagree and ckappa functions provide an object that can be used to assess the similarity based on the predicted classes of two results. The predicted classes are selected by the class with the highest probability.

The bdist (Bhattacharayya distance), tvdist (Total variation distance), hdist (Hellinger distance) and jsdist (Jenson-Shannon divergence) functions provide an object that can be used to assess the similarity based on the predicted class probabilities of two results.

#### See Also

```
stability
```

```
set.seed(0)
## build trees
library("partykit")
m1 <- ctree(Species ~ ., data = iris[sample(1:nrow(iris), replace = TRUE),])</pre>
m2 <- ctree(Species ~ ., data = iris[sample(1:nrow(iris), replace = TRUE),])</pre>
p1 <- predict(m1, type = "prob")</pre>
p2 <- predict(m2, type = "prob")
## class agreement
m <- clagree()</pre>
m$measure(p1, p2)
## cohen's kappa
m <- ckappa()
m$measure(p1, p2)
## bhattacharayya distance
m <- bdist()
m$measure(p1, p2)
## total variation distance
m <- tvdist()</pre>
m$measure(p1, p2)
```

```
## hellinger distance
m <- hdist()
m$measure(p1, p2)

## jenson-shannon divergence
m <- jsdiv()
m$measure(p1, p2)

## jenson-shannon divergence (base = exp(1))
m <- jsdiv(base = exp(1))
m$measure(p1, p2)</pre>
```

similarity\_measures\_regression

Similarity Measure Infrastructure for Stability Assessment with Numerical Responses

### **Description**

Functions that provide objects with functionality used by stability to measure the similarity between numeric predictions of two results in regression problems.

### Usage

```
edist()
msdist()
rmsdist()
madist()
qadist(p = 0.95)

cprob(kappa = 0.1)
rbfkernel()
tanimoto()
cosine()

ccc()
pcc()
```

### **Arguments**

kappa

p A numeric value between 0 and 1 specifying the probability to which the sample

quantile of the absolute distance between the predictions is computed.

A positive numeric value specifying the upper limit of the absolute distance between the predictions to which the coverage probability is computed.

#### **Details**

The similarity measure functions provide objects that include functionality used by stability to measure the similarity between numeric predictions of two results in regression problems.

The edist (euclidean distance), msdist (mean squared distance), rmsdist (root mean squared distance), madist (mean absolute distance) and qadist (quantile of absolute distance) functions implement scale-variant distance measures that are unbounded.

The cprob (coverage probability), rbfkernel (gaussian radial basis function kernel), tanimoto (tanimoto coefficient) and cosine (cosine similarity) functions implement scale-variant distance measures that are bounded.

The ccc (concordance correlation coefficient) and pcc (pearson correlation coefficient) functions implement scale-invariant distance measures that are bounded between 0 and 1.

#### See Also

```
stability
```

```
set.seed(0)
library("partykit")
airq <- subset(airquality, !is.na(Ozone))</pre>
m1 <- ctree(Ozone ~ ., data = airq[sample(1:nrow(airq), replace = TRUE),])</pre>
m2 <- ctree(Ozone ~ ., data = airq[sample(1:nrow(airq), replace = TRUE),])</pre>
p1 <- predict(m1)</pre>
p2 <- predict(m2)</pre>
## euclidean distance
m <- edist()
m$measure(p1, p2)
## mean squared distance
m <- msdist()</pre>
m$measure(p1, p2)
## root mean squared distance
m <- rmsdist()</pre>
m$measure(p1, p2)
## mean absolute istance
m <- madist()</pre>
m$measure(p1, p2)
## quantile of absolute distance
m <- qadist()</pre>
```

similarity\_values

```
m$measure(p1, p2)
## coverage probability
m <- cprob()</pre>
m$measure(p1, p2)
## gaussian radial basis function kernel
m <- rbfkernel()</pre>
m$measure(p1, p2)
## tanimoto coefficient
m <- tanimoto()</pre>
m$measure(p1, p2)
## cosine similarity
m <- cosine()</pre>
m$measure(p1, p2)
## concordance correlation coefficient
m <- ccc()
m$measure(p1, p2)
## pearson correlation coefficient
m <- pcc()
m$measure(p1, p2)
```

similarity\_values

Extracting Similarity Values

### Description

Extract similarity values from object returned by stability for further illustration or analysis.

### Usage

```
similarity_values(x, reverse = TRUE)
```

### **Arguments**

x an object of class "stablelearner" or "stablelearnerList" from which the

similarity values are extracted.

reverse logical. If reverse = TRUE (default), the similarity values are transformed (re-

versed) such that higher values indicate a higher stability.

### Value

A numeric array of dimension 3 containing similarity values. The dimensions represent repetitions, results (fitted model objects) and similarity measures.

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### See Also

```
stability, summary.stablelearnerList
```

### **Examples**

```
library("partykit")
res <- ctree(Species ~ ., data = iris)
stab <- stability(res)
similarity_values(stab)</pre>
```

stability

Stability Assessment for Results from Supervised Statistical Learning

### **Description**

Stability assessment of results from supervised statistical learning (i.e., recursive partitioning, support vector machines, neural networks, etc.). The procedure involves the pairwise comparison of results generated from learning samples randomly drawn from the original data set or directly from the data-generating process (if available).

### Usage

```
stability(x, ..., data = NULL, control = stab_control(), weights = NULL,
applyfun = NULL, cores = NULL, names = NULL)
```

### **Arguments**

X	fitted model object. Any model object can be used whose class is registered in LearnerList. Users can add classes for the current R session to LearnerList, see addLearner.
	additional fitted model objects.
data	an optional data.frame or a data-generating function. By default the learning data from $x$ is used (if this can be inferred from the getCall of $x$ .
control	a list with control parameters, see stab_control.
weights	an optional matrix of dimension n * B that can be used to weight the observations from the original learning data when the models are refitted. If weights = true, the weights are computed internally according to the sampler defined in control. If weight = NULL (default), no case-weights are used and the sampler defined in control will be applied to the original data set.
applyfun	a lapply-like function. The default is to use lapply unless cores is specified in which case mclapply is used (for multicore computations on platforms that support these).
cores	integer. The number of cores to use in multicore computations using mclapply (see above).
names	a vector of characters to specify a name for each fitted model object. By default,

the objects are named by their class.

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#### **Details**

See Philipp et al. (2018).

#### Value

For a single fitted model object, stability returns an object of class "stablelearner" with the following components:

call the call from the model object x,

learner the information about the learner retrieved from LearnerList,

B the number of repetitions,

sval a matrix containing the estimated similarity values for each similarity measure

specified in control,

sampstat a list containing information on the size of the learning samples (1s), the size of

the overlap between the learning samples (10), the size of the evaluation sample (es) and the size of the overlap between the evaluation and the learning samples

(eo) in each repetition.

data a language object referring to the data.frame or the data-generating function

used for assessing the stability,

control a list with control parameters used for assessing the stability,

For several fitted model objects, stability returns an object of class "stablelearnerList" which is a list of objects of class "stablelearner".

#### References

Philipp M, Rusch T, Hornik K, Strobl C (2018). "Measuring the Stability of Results from Supervised Statistical Learning". *Journal of Computational and Graphical Statistics*, **27**(4), 685–700. doi: 10.1080/10618600.2018.1473779

#### See Also

boxplot.stablelearnerList, summary.stablelearner

```
## assessing the stability of a single result
library("partykit")
r1 <- ctree(Species ~ ., data = iris)
stab <- stability(r1)
summary(stab)

## assessing the stability of several results
library("rpart")
r2 <- rpart(Species ~ ., data = iris)
stab <- stability(r1, r2, control = stab_control(seed = 0))</pre>
```

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```
summary(stab, names = c("ctree", "rpart"))
## using case-weights instead of resampling
stability(r1, weights = TRUE)

## using self-defined case-weights
n <- nrow(iris)
B <- 500
w <- array(sample(c(0, 1), size = n * B * 3, replace = TRUE), dim = c(n, B, 3))
stability(r1, weights = w)

## assessing stability for a given data-generating process
my_dgp <- function() dgp_twoclass(n = 100, p = 2, noise = 4, rho = 0.2)
res <- ctree(class ~ ., data = my_dgp())
stability(res, data = my_dgp)</pre>
```

stabletree

Stability Assessment for Tree Learners

### **Description**

Stability assessment of variable and cutpoint selection in tree learners (i.e., recursive partitioning). By refitting trees to resampled versions of the learning data, the stability of the trees structure is assessed and can be summarized and visualized.

### Usage

```
stabletree(x, data = NULL, sampler = subsampling, weights = NULL,
applyfun = NULL, cores = NULL, savetrees = FALSE, ...)
```

### **Arguments**

X	fitted model object. Any tree-based model object that can be coerced by as.party can be used provided that suitable methods are provided.
data	an optional data. frame. By default the learning data from $x$ is used (if this can be inferred from the getCall of $x$ .
sampler	a resampling (generating) function. Either this should be a function of n that returns a matrix or a sampler generator like subsampling.
weights	an optional matrix of dimension $n * B$ that can be used to weight the observations from the original learning data when the trees are refitted. If weight = NULL, the sampler will be used.
applyfun	a lapply-like function. The default is to use lapply unless cores is specified in which case mclapply is used (for multicore computations on platforms that support these).

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cores integer. The number of cores to use in multicore computations using mclapply

(see above).

savetrees logical. If TRUE, trees based on resampled data sets are returned.

... further arguments passed to sampler.

#### **Details**

The function stabletree assesses the stability of tree learners (i.e., recursive partitioning methods) by refitting the tree to resampled versions of the learning data. By default, if data = NULL, the fitting call is extracted by getCall to infer the learning data. Subsequently, the sampler generates B resampled versions of the learning data, the tree is regrown with update, and (if necessary) coerced by as.party. For each of the resampled trees it is queried and stored which variables are selected for splitting and what the selected cutpoints are.

The resulting object of class "stabletree" comes with a set of standard methods to generic functions including print, summary for numerical summaries and plot, barplot, and image for graphical representations. See plot.stabletree for more details. In most methods, the argument original can be set to TRUE or FALSE, turning highlighting of the original tree information on and off.

#### Value

stabletree returns an object of class "stabletree" which is a list with the following components:

call the call from the model object x,

B the number of resampled datasets,

sampler the sampler function,

vs0 numeric vector of the variable selections of the original tree,

br0 list of the break points (list of nodeids, levels, and breaks) for each variable

of the original tree,

vs numeric matrix of the variable selections for each resampled dataset,

br list of the break points (only the breaks for each variable over all resampled

datasets,

classes character vector indicating the classes of all partitioning variables,

trees a list of tree objects of class "party", or NULL.

### References

Hothorn T, Zeileis A (2015). partykit: A Modular Toolkit for Recursive Partytioning in R. *Journal of Machine Learning Research*, **16**(118), 3905–3909.

Philipp M, Zeileis A, Strobl C (2016). "A Toolkit for Stability Assessment of Tree-Based Learners". In A. Colubi, A. Blanco, and C. Gatu (Eds.), Proceedings of COMPSTAT 2016 – 22nd International Conference on Computational Statistics (pp. 315–325). The International Statistical Institute/International Association for Statistical Computing. Preprint available at https://EconPapers.RePEc.org/RePEc:inn:wpaper:2016-11

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### See Also

```
plot.stabletree, as.stabletree, as.party
```

### **Examples**

```
## build a simple tree
library("partykit")
m <- ctree(Species ~ ., data = iris)</pre>
plot(m)
## investigate stability
set.seed(0)
s <- stabletree(m, B = 500)
print(s)
## variable selection statistics
summary(s)
## show variable selection proportions
barplot(s)
## illustrate variable selections of replications
image(s)
## graphical cutpoint analysis
plot(s)
```

stabletree-coercion Coercion Functions

### **Description**

Functions coercing various forest objects to objects of class "stabletree".

### Usage

```
as.stabletree(x, ...)
## S3 method for class 'randomForest'
as.stabletree(x, applyfun = NULL, cores = NULL, ...)
## S3 method for class 'RandomForest'
as.stabletree(x, applyfun = NULL, cores = NULL, ...)
## S3 method for class 'cforest'
as.stabletree(x, applyfun = NULL, cores = NULL, savetrees = FALSE, ...)
## S3 method for class 'ranger'
as.stabletree(x, applyfun = NULL, cores = NULL, ...)
```

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### **Arguments**

x an object of class randomForest, RandomForest-class, cforest, or ranger.

applyfun a lapply-like function. The default is to use lapply unless cores is specified

in which case mclapply is used (for multicore computations on platforms that

support these).

cores integer. The number of cores to use in multicore computations using mclapply

(see above).

savetrees logical. If TRUE, trees of the forests are returned.

... additional arguments (currently not used).

#### **Details**

Random forests fitted using randomForest, cforest, cforest or ranger are coerced to "stabletree" objects.

Note that when plotting a randomForest or ranger, the gray areas of levels of a nominal variable do not mimic exactly the same behavior as for classical "stabletree" objects, due to randomForest and ranger, not storing any information whether any individuals were left fulfilling the splitting criterion in the subsample. Therefore, gray areas only indicate that this level of this variable has already been used in a split before in such a way that it could not be used for any further splits.

For ranger, interaction terms are (currently) not supported.

#### Value

as.stabletree returns an object of class "stabletree" which is a list with the following components:

call the call from the model object x,

B the number of trees of the random forest,

sampler the random forest fitting function,

vs0 numeric vector of the variable selections of the original tree, here always a vector

of zeros because there is no original tree,

br0 always NULL (only included for consistency),

vs numeric matrix of the variable selections for each tree of the random forest,

br list of the break points (only the breaks for each variable over all trees of the

random forest,

classes character vector indicating the classes of all partitioning variables,

trees a list of tree objects of class "party", or NULL.

#### See Also

stabletree, plot.stabletree

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```
## build a randomForest using randomForest
library(randomForest)
set.seed(1)
rf <- randomForest(Species ~ ., data = iris)</pre>
## coerce to a stabletree
srf <- as.stabletree(rf)</pre>
print(srf)
summary(srf, original = FALSE) # there is no original tree
barplot(srf)
image(srf)
plot(srf)
## build a RandomForest using party
library("party")
set.seed(2)
cf_party <- cforest(Species ~ ., data = iris,</pre>
  control = cforest_unbiased(mtry = 2))
## coerce to a stabletree
scf_party <- as.stabletree(cf_party)</pre>
print(scf_party)
summary(scf_party, original = FALSE)
barplot(scf_party)
image(scf_party)
plot(scf_party)
## build a cforest using partykit
library("partykit")
set.seed(3)
cf_partykit <- cforest(Species ~ ., data = iris)</pre>
## coerce to a stabletree
scf_partykit <- as.stabletree(cf_partykit)</pre>
print(scf_partykit)
summary(scf_partykit, original = FALSE)
barplot(scf_partykit)
image(scf_partykit)
plot(scf_partykit)
## build a random forest using ranger
library("ranger")
set.seed(4)
rf_ranger <- ranger(Species ~ ., data = iris)</pre>
## coerce to a stabletree
srf_ranger <- as.stabletree(rf_ranger)</pre>
print(srf_ranger)
summary(srf_ranger, original = FALSE)
```

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```
barplot(srf_ranger)
image(srf_ranger)
plot(srf_ranger)
```

stab\_control

Control for Supervised Stability Assessments

### Description

Various parameters that control aspects of the stability assessment performed via stability.

### Usage

```
stab_control(B = 500, measure = list(tvdist, ccc), sampler = "bootstrap",
  evaluate = "00B", holdout = 0.25, seed = NULL, na.action = na.exclude,
  savepred = TRUE, silent = TRUE, ...)
```

### **Arguments**

В	an integer value specifying the number of repetitions. The default is B = 500.
measure	a list of similarity measure (generating) functions. Those should either be functions of p1 and p2 (the predictions of two results in a repetition, see Details below) or similarity measure generator functions, see similarity_measures_classification and similarity_measures_regression. The defaults are tvdist for the classification and ccc for the regression case.
sampler	a resampling (generating) function. Either this should be a function of n that returns a matrix or a sampler generator like bootstrap. The default is "bootstrap".
evaluate	a character specifying the evaluation strategy to be applied (see Details below). The default is "00B".
holdout	a numeric value between zero and one that specifies the proportion of observations hold out for evaluation over all repetitions, only if evaluate = "00B". The default is $0.25$ .
seed	a single value, interpreted as an integer, see set. seed and Details section below. The default is NULL, which indicates that no seed has been set.
na.action	a function which indicates what should happen when the predictions of the results contain NAs. The default function is na.exclude.
savepred	logical. Should the predictions from each iteration be saved? If savepred = FALSE, the resulting object will be smaller. However, predictions are required to subsequently compute measures of accuracy via accuracy.
silent	logical. If TRUE, error messages, generated by the learner while training or predicting, are suppressed and NA is returned for the corresponding iteration of the stability assessment procedure.
	arguments passed to sampler function.

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#### **Details**

With the argument measure one or more measures can be defined that are used to assess the stability of a result from supervised statistical learning by stability. Predefined similarity measures for the regression and the classification case are listed in similarity\_measures\_classification and similarity\_measures\_regression.

Users can define their own similarity functions f(p1,p2) that must return a single numeric value for the similarity between two results trained on resampled data sets. Such a function must take the arguments p1 and p2. In the classification case, p1 and p2 are probability matrices of size m \* K, where m is the number of predicted observations (size of the evaluation sample) and K is the number of classes. In the regression case, p1 and p2 are numeric vectors of length m.

A different way to implement new similarity functions for the current R session is to define a similarity measure generator function, which is a function without arguments that generates a list of five elements including the name of the similarity measure, the function to compute the similarity between the predictions as described above, a vector of character values specifying the response types for which the similarity measure can be used, a list containing two numeric elements lower and upper that specify the range of values of the similarity measure and the function to invert (or reverse) the similarity values such that higher values indicate higher stability. The latter can be set to NULL, if higher similarity values already indicate higher stability. Those elements should be named name, measure, classes, range and reverse.

The argument evaluate can be used to specify the evaluation strategy. If set to "ALL", all observations in the original data set are used for evaluation. If set to "OOB", only the pairwise out-of-bag observations are used for evaluation within each repetition. If set to "OOS", a fraction (defined by holdout) of the observations in the original data set are randomly sampled and used for evaluation, but not for training, over all repetitions.

The argument seed can be used to make similarity assessments comparable when comparing the stability of different results that were trained on the same data set. By default, seed is set to NULL and the learning samples are sampled independently for each fitted model object passed to stability. If seed is set to a specific number, the seed will be set for each fitted model object before the learning samples are generated using "L'Ecuyer-CMRG" (see set.seed) which guarantees identical learning samples for each stability assessment and, thus, comparability of the stability assessments between the results.

#### See Also

```
stability
```

```
library("partykit")
res <- ctree(Species ~ ., data = iris)
## less repetitions
stability(res, control = stab_control(B = 100))</pre>
```

```
## Not run:
## change similarity measure
stability(res, control = stab_control(measure = list(bdist)))

## change evaluation strategy
stability(res, control = stab_control(evaluate = "ALL"))
stability(res, control = stab_control(evaluate = "00S"))

## change resampling strategy to subsampling
stability(res, control = stab_control(sampler = subsampling))
stability(res, control = stab_control(sampler = subsampling, evaluate = "ALL"))
stability(res, control = stab_control(sampler = subsampling, evaluate = "00S"))

## change resampling strategy to splithalf
stability(res, control = stab_control(sampler = splithalf, evaluate = "ALL"))
stability(res, control = stab_control(sampler = splithalf, evaluate = "00S"))

## End(Not run)
```

summary.stablelearnerList

Summarize Results from Stability Assessment

### **Description**

Summarizes and prints the results from stability assessments performed by stability.

### Usage

```
## S3 method for class 'stablelearnerList'
summary(object, ..., reverse = TRUE,
   probs = c(0.05, 0.25, 0.5, 0.75, 0.95), digits = 3, names = NULL)
```

### **Arguments**

object	a object of class "stablelearner" or "stablelearnerList" to be summarized.
	Arguments passed from or to other functions (currently ignored).
reverse	logical. If reverse = TRUE (default), the similarity values summarized in the summary output are transformed (reversed) such that higher values indicate a higher stability.
digits	integer. The number of digits used to summarize the similarity distribution in

the summary output.

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probs a vector of probabilities used tosummarize the similarity distribution in the

summary output.

names a vector of characters to specify a name for each result from statistical learning

in the summary output.

#### See Also

```
stability
```

### **Examples**

```
library("partykit")

rval <- ctree(Species ~ ., data = iris)
stab <- stability(rval)

summary(stab)
summary(stab, reverse = FALSE)
summary(stab, probs = c(0.25, 0.5, 0.75))
summary(stab, names = "conditional inference tree")</pre>
```

titanic

Passengers and Crew on the RMS Titanic

### **Description**

the titanic data is a complete list of passengers and crew members on the RMS Titanic. It includes a variable indicating whether a person did survive the sinking of the RMS Titanic on April 15, 1912.

### Usage

```
data("titanic")
```

### **Format**

A data frame containing 2207 observations on 11 variables.

name a string with the name of the passenger.

gender a factor with levels male and female.

**age** a numeric value with the persons age on the day of the sinking. The age of babies (under 12 months) is given as a fraction of one year (1/month).

class a factor specifying the class for passengers or the type of service aboard for crew members.

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**embarked** a factor with the persons place of of embarkment.

**country** a factor with the persons home country.

ticketno a numeric value specifying the persons ticket number (NA for crew members).

**fare** a numeric value with the ticket price (NA for crew members, musicians and employees of the shipyard company).

**sibsp** an ordered factor specifying the number if siblings/spouses aboard; adopted from Vanderbild data set (see below).

**parch** an ordered factor specifying the number of parents/children aboard; adopted from Vander-bild data set (see below).

**survived** a factor with two levels (no and yes) specifying whether the person has survived the sinking.

#### **Details**

The website <a href="https://www.encyclopedia-titanica.org/">https://www.encyclopedia-titanica.org/</a> offers detailed information about passengers and crew members on the RMS Titanic. According to the website 1317 passengers and 890 crew member were abord.

8 musicians and 9 employees of the shipyard company are listed as passengers, but travelled with a free ticket, which is why they have NA values in fare. In addition to that, fare is truely missing for a few regular passengers.

#### Source

The complete list of persons on the RMS titanic was downloaded from https://www.encyclopedia-titanica.org/ on April 5, 2016. The information given in sibsp and parch was adopted from a data set obtained from https://hbiostat.org/data/.

### References

https://www.encyclopedia-titanica.org/ and https://hbiostat.org/data/.

### **Examples**

```
data("titanic", package = "stablelearner")
summary(titanic)
```

tuner

**Tuning Wrapper Function** 

### **Description**

Convenience function to train a method using different tuning parameters.

### Usage

```
tuner(method, tunerange, ...)
```

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### Arguments

method a character string. Name of the R function to train the method.

tunerange a list. A list that specifies the range of values to be used for each tuning pa-

rameter. Each element of the list should be a vector that specifies the values to be tested for the tuning parameter. The element must be named after the

corresponding tuning parameter of the method (see examples).

... additional information passed to method (such as formula, data, subset, etc.).

### **Details**

This function can be used to train any method using different values for its tuning parameter(s). The result can be passed directly to stability to compare the stability of results based on different values of the tuning parameter.

#### Value

A list that contains all fitted model objects.

Additional information about the range of values used for the tuning parameters is attached to the resulting object as an attribute.

#### See Also

```
stability
```

```
library("partykit")

## tuning cforest using different values of its tuning parameter mtry
r <- tuner("cforest", tunerange = list(mtry = 1:4), formula = Species ~ ., data = iris)

## assess stability (with B = 10 for illustration to avoid excessive computation times)
stability(r, control = stab_control(seed = 1234, B = 10))

## receive information about the range of tuning parameters
attr(r, "range")</pre>
```

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