Package ‘ranger’

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Type Package

Title A Fast Implementation of Random Forests

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Description A fast implementation of Random Forests, particularly suited for high
dimensional data. Ensembles of classification, regression, survival and
probability prediction trees are supported. Data from genome-wide association
studies can be analyzed efficiently. In addition to data frames, datasets of
class 'gwaa.data' (R package 'GenABEL') and 'dgCMatrix' (R package 'Matrix')
can be directly analyzed.

License GPL-3

Imports Rcpp (>= 0.11.2), Matrix

LinkingTo Rcpp, RcppEigen

Depends R (>= 3.1)

Suggests covr, survival, testthat

Encoding UTF-8

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R topics documented:

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In case-specific random forests (CSRF), random forests are built specific to the cases of interest. Instead of using equal probabilities, the cases are weighted according to their difference to the case of interest.

```r
csf(formula, training_data, test_data, params1 = list(), params2 = list(), verbose = FALSE)
```

**formula** Object of class formula or character describing the model to fit.

**training_data** Training data of class data.frame.

**test_data** Test data of class data.frame.

**params1** Parameters for the proximity random forest grown in the first step.

**params2** Parameters for the prediction random forests grown in the second step.

**verbose** Logical indicating whether or not to print computation progress.
Details

The algorithm consists of 3 steps:

1. Grow a random forest on the training data
2. For each observation of interest (test data), the weights of all training observations are computed by counting the number of trees in which both observations are in the same terminal node.
3. For each test observation, grow a weighted random forest on the training data, using the weights obtained in step 2. Predict the outcome of the test observation as usual.

In total, n+1 random forests are grown, where n is the number observations in the test dataset. For details, see Xu et al. (2014).

Value

Predictions for the test dataset.

Author(s)

Marvin N. Wright

References


Examples

```r
## Split in training and test data
train.idx <- sample(nrow(iris), 2/3 * nrow(iris))
iris.train <- iris[train.idx, ]
iris.test <- iris[-train.idx, ]

## Run case-specific RF
csf(Species ~ ., training_data = iris.train, test_data = iris.test,
   params1 = list(num.trees = 50, mtry = 4),
   params2 = list(num.trees = 5))
```

---

**getTerminalNodeIDs**  
*Get terminal node IDs (deprecated)*

Description

This function is deprecated. Please use predict() with type = "terminalNodes" instead. This function calls predict() now.
Usage

getTerminalNodeIDs(rf, dat)

Arguments

rf ranger object.
dat New dataset. Terminal node IDs for this dataset are obtained.

Value

Matrix with terminal node IDs for all observations in dataset and trees.

Examples

rf <- ranger(Species ~ ., data = iris, num.trees = 5, write.forest = TRUE)
getTerminalNodeIDs(rf, iris)

Description

Grow two random forests on two cross-validation folds. Instead of out-of-bag data, the other fold
is used to compute permutation importance. Related to the novel permutation variable importance
by Janitza et al. (2015).

Usage

holdoutRF(...)  

Arguments

... All arguments are passed to ranger() (except importance, case.weights, 
replace and holdout.).

Value

Hold-out random forests with variable importance.

Author(s)

Marvin N. Wright

References

importance.ranger

See Also

ranger

---

importance.ranger  
*ranger variable importance*

Description

Extract variable importance of ranger object.

Usage

```r
## S3 method for class 'ranger'
importance(x, ...)
```

Arguments

- `x`  
ranger object.

- `...`  
Further arguments passed to or from other methods.

Value

Variable importance measures.

Author(s)

Marvin N. Wright

See Also

ranger

---

importance_pvalues  
*ranger variable importance p-values*

Description

Compute variable importance with p-values. For high dimensional data, the fast method of Janitza et al. (2016) can be used. The permutation approach of Altmann et al. (2010) is computationally intensive but can be used with all kinds of data. See below for details.
Usage

importance_pvalues(
  x,
  method = c("janitza", "altmann"),
  num.permutations = 100,
  formula = NULL,
  data = NULL,
  ...
)

Arguments

x ranger or holdoutRF object.
method Method to compute p-values. Use "janitza" for the method by Janitza et al. (2016) or "altmann" for the non-parametric method by Altmann et al. (2010).
num.permutations Number of permutations. Used in the "altmann" method only.
formula Object of class formula or character describing the model to fit. Used in the "altmann" method only.
data Training data of class data.frame or matrix. Used in the "altmann" method only.
...
Further arguments passed to ranger(). Used in the "altmann" method only.

Details

The method of Janitza et al. (2016) uses a clever trick: With an unbiased variable importance measure, the importance values of non-associated variables vary randomly around zero. Thus, all non-positive importance values are assumed to correspond to these non-associated variables and they are used to construct a distribution of the importance under the null hypothesis of no association to the response. Since only the non-positive values of this distribution can be observed, the positive values are created by mirroring the negative distribution. See Janitza et al. (2016) for details.

The method of Altmann et al. (2010) uses a simple permutation test: The distribution of the importance under the null hypothesis of no association to the response is created by several replications of permuting the response, growing an RF and computing the variable importance. The authors recommend 50-100 permutations. However, much larger numbers have to be used to estimate more precise p-values. We add 1 to the numerator and denominator to avoid zero p-values.

Value

Variable importance and p-value for each variable.

Author(s)

Marvin N. Wright
parse.formula

References


See Also

ranger

Examples

## Janitzia's p-values with corrected Gini importance
n <- 50
p <- 400
dat <- data.frame(y = factor(rbinom(n, 1, .5)), replicate(p, runif(n)))
rf.sim <- ranger(y ~ ., dat, importance = "impurity_corrected")
importance_pvalues(rf.sim, method = "janitza")

## Permutation p-values
## Not run:
rf.iris <- ranger(Species ~ ., data = iris, importance = 'permutation')
importance_pvalues(rf.iris, method = "altmann", formula = Species ~ ., data = iris)
## End(Not run)

parse.formula Parse formula

Description

Parse formula and return dataset containing selected columns. Interactions are supported for numerical columns only. An interaction column is the product of all interacting columns.

Usage

parse.formula(formula, data, env = parent.frame())

Arguments

  formula Object of class formula or character describing the model to fit.
  data Training data of class data.frame.
  env The environment in which the left hand side of formula is evaluated.

Value

Dataset including selected columns and interactions.
**predict.ranger**

**Ranger prediction**

**Description**

Prediction with new data and a saved forest from Ranger.

**Usage**

```r
## S3 method for class 'ranger'
predict(
  object,
  data = NULL,
  predict.all = FALSE,
  num.trees = object$num.trees,
  type = "response",
  se.method = "infjack",
  quantiles = c(0.1, 0.5, 0.9),
  what = NULL,
  seed = NULL,
  num.threads = NULL,
  verbose = TRUE,
  ...)
```

**Arguments**

- `object` Ranger ranger object.
- `data` New test data of class `data.frame` or `gwaa.data` (GenABEL).
- `predict.all` Return individual predictions for each tree instead of aggregated predictions for all trees. Return a matrix (sample x tree) for classification and regression, a 3d array for probability estimation (sample x class x tree) and survival (sample x time x tree).
- `num.trees` Number of trees used for prediction. The first `num.trees` in the forest are used.
- `type` Type of prediction. One of 'response', 'se', 'terminalNodes', 'quantiles' with default 'response'. See below for details.
- `se.method` Method to compute standard errors. One of 'jack', 'infjack' with default 'infjack'. Only applicable if type = 'se'. See below for details.
- `quantiles` Vector of quantiles for quantile prediction. Set type = 'quantiles' to use.
- `what` User specified function for quantile prediction used instead of `quantile`. Must return numeric vector, see examples.
- `seed` Random seed. Default is `NULL`, which generates the seed from R. Set to 0 to ignore the R seed. The seed is used in case of ties in classification mode.
- `num.threads` Number of threads. Default is number of CPUs available.
- `verbose` Verbosity output on or off.
- `...` further arguments passed to or from other methods.
predict.ranger

Details

For type = 'response' (the default), the predicted classes (classification), predicted numeric values (regression), predicted probabilities (probability estimation) or survival probabilities (survival) are returned. For type = 'se', the standard error of the predictions are returned (regression only). The jackknife-after-bootstrap or infinitesimal jackknife for bagging is used to estimate the standard errors based on out-of-bag predictions. See Wager et al. (2014) for details. For type = 'terminalNodes', the IDs of the terminal node in each tree for each observation in the given dataset are returned. For type = 'quantiles', the selected quantiles for each observation are estimated. See Meinshausen (2006) for details.

If type = 'se' is selected, the method to estimate the variances can be chosen with se.method. Set se.method = 'jack' for jackknife-after-bootstrap and se.method = 'infjack' for the infinitesimal jackknife for bagging.

For classification and predict.all = TRUE, a factor levels are returned as numerics. To retrieve the corresponding factor levels, use rf$forest$levels, if rf is the ranger object.

Value

Object of class ranger.prediction with elements

- predictions: Predicted classes/values (only for classification and regression)
- unique.death.times: Unique death times (only for survival).
- chf: Estimated cumulative hazard function for each sample (only for survival).
- survival: Estimated survival function for each sample (only for survival).
- num.trees: Number of trees.
- num.independent.variables: Number of independent variables.
- treetype: Type of forest/tree. Classification, regression or survival.
- num.samples: Number of samples.

Author(s)

Marvin N. Wright

References


See Also

ranger
Examples

```r
## Classification forest
ranger(Species ~ ., data = iris)
train.idx <- sample(nrow(iris), 2/3 * nrow(iris))
iris.train <- iris[train.idx, ]
iris.test <- iris[-train.idx, ]
rg.iris <- ranger(Species ~ ., data = iris.train)
pred.iris <- predict(rg.iris, data = iris.test)
table(iris.test$Species, pred.iris$predictions)

## Quantile regression forest
rf <- ranger(mpg ~ ., mtcars[1:26, ], quantreg = TRUE)
pred <- predict(rf, mtcars[27:32, ], type = "quantiles", quantiles = c(0.1, 0.5, 0.9))
pred$predictions

## Quantile regression forest with user-specified function
rf <- ranger(mpg ~ ., mtcars[1:26, ], quantreg = TRUE)
pred <- predict(rf, mtcars[27:32, ], type = "quantiles",
                  what = function(x) sample(x, 10, replace = TRUE))
pred$predictions
```

predict.ranger.forest  Ranger prediction

Description

Prediction with new data and a saved forest from Ranger.

Usage

```r
## S3 method for class 'ranger.forest'
predict(
  object,
  data,
  predict.all = FALSE,
  num.trees = object$num.trees,
  type = "response",
  se.method = "infjack",
  seed = NULL,
  num.threads = NULL,
  verbose = TRUE,
  inbag.counts = NULL,
  ...
)
```
predict.ranger.forest

Arguments

- **object**: Ranger ranger.forest object.
- **data**: New test data of class data.frame or gwaa.data (GenABEL).
- **predict.all**: Return individual predictions for each tree instead of aggregated predictions for all trees. Return a matrix (sample x tree) for classification and regression, a 3d array for probability estimation (sample x class x tree) and survival (sample x time x tree).
- **num.trees**: Number of trees used for prediction. The first num.trees in the forest are used.
- **type**: Type of prediction. One of 'response', 'se', 'terminalNodes', 'quantiles' with default 'response'. See below for details.
- **se.method**: Method to compute standard errors. One of 'jack', 'infjack' with default 'infjack'. Only applicable if type = 'se'. See below for details.
- **seed**: Random seed. Default is NULL, which generates the seed from R. Set to 0 to ignore the R seed. The seed is used in case of ties in classification mode.
- **num.threads**: Number of threads. Default is number of CPUs available.
- **verbose**: Verbose output on or off.
- **inbag.counts**: Number of times the observations are in-bag in the trees.
- **...**: further arguments passed to or from other methods.

Details

For type = 'response' (the default), the predicted classes (classification), predicted numeric values (regression), predicted probabilities (probability estimation) or survival probabilities (survival) are returned. For type = 'se', the standard error of the predictions are returned (regression only). The jackknife-after-bootstrap or infinitesimal jackknife for bagging is used to estimate the standard errors based on out-of-bag predictions. See Wager et al. (2014) for details. For type = 'terminalNodes', the IDs of the terminal node in each tree for each observation in the given dataset are returned.

If type = 'se' is selected, the method to estimate the variances can be chosen with se.method. Set se.method = 'jack' for jackknife after bootstrap and se.method = 'infjack' for the infinitesimal jackknife for bagging.

For classification and predict.all = TRUE, a factor levels are returned as numerics. To retrieve the corresponding factor levels, use rf$forest$levels, if rf is the ranger object.

Value

Object of class ranger.prediction with elements

- **predictions**: Predicted classes/values (only for classification and regression)
- **unique.death.times**: Unique death times (only for survival).
- **chf**: Estimated cumulative hazard function for each sample (only for survival).
- **survival**: Estimated survival function for each sample (only for survival).
- **num.trees**: Number of trees.
- **num.independent.variables**: Number of independent variables.
- **treetype**: Type of forest/tree. Classification, regression or survival.
- **num.samples**: Number of samples.
Author(s)
Marvin N. Wright

References


See Also

ranger

Description

Extract training data predictions of Ranger object.

Usage

```r
## S3 method for class 'ranger'
predictions(x, ...)
```

Arguments

- `x` Ranger object.
- `...` Further arguments passed to or from other methods.

Value

Predictions: Classes for Classification forests, Numerical values for Regressions forests and the estimated survival functions for all individuals for Survival forests.

Author(s)
Marvin N. Wright

See Also

ranger
predictions.ranger.prediction

**Ranger predictions**

**Description**

Extract predictions of Ranger prediction object.

**Usage**

```r
## S3 method for class 'ranger.prediction'
predictions(x, ...)
```

**Arguments**

- `x`: Ranger prediction object.
- `...`: Further arguments passed to or from other methods.

**Value**

Predictions: Classes for Classification forests, Numerical values for Regressions forests and the estimated survival functions for all individuals for Survival forests.

**Author(s)**

Marvin N. Wright

**See Also**

- `ranger`

print.ranger

**Print Ranger**

**Description**

Print contents of Ranger object.

**Usage**

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

**Arguments**

- `x`: Object of class 'ranger'.
- `...`: Further arguments passed to or from other methods.
Author(s)
Marvin N. Wright

See Also
ranger

print.ranger.forest  
Print Ranger forest

Description
Print contents of Ranger forest object.

Usage
## S3 method for class 'ranger.forest'
print(x, ...)

Arguments
x  Object of class 'ranger.forest'.
...  further arguments passed to or from other methods.

Author(s)
Marvin N. Wright

print.ranger.prediction  
Print Ranger prediction

Description
Print contents of Ranger prediction object.

Usage
## S3 method for class 'ranger.prediction'
print(x, ...)

Arguments
x  Object of class 'ranger.prediction'.
...  further arguments passed to or from other methods.
Author(s)

Marvin N. Wright

Description

Ranger is a fast implementation of random forests (Breiman 2001) or recursive partitioning, particularly suited for high dimensional data. Classification, regression, and survival forests are supported. Classification and regression forests are implemented as in the original Random Forest (Breiman 2001), survival forests as in Random Survival Forests (Ishwaran et al. 2008). Includes implementations of extremely randomized trees (Geurts et al. 2006) and quantile regression forests (Meinshausen 2006).

Usage

ranger(
    formula = NULL,
    data = NULL,
    num.trees = 500,
    mtry = NULL,
    importance = "none",
    write.forest = TRUE,
    probability = FALSE,
    min.node.size = NULL,
    max.depth = NULL,
    replace = TRUE,
    sample.fraction = ifelse(replace, 1, 0.632),
    case.weights = NULL,
    class.weights = NULL,
    splitrule = NULL,
    num.random.splits = 1,
    alpha = 0.5,
    minprop = 0.1,
    split.select.weights = NULL,
    always.split.variables = NULL,
    respect.unordered.factors = NULL,
    scale.permutation.importance = FALSE,
    local.importance = FALSE,
    regularization.factor = 1,
    regularization.usedepth = FALSE,
    keep.inbag = FALSE,
    inbag = NULL,
    holdout = FALSE,
    quantreg = FALSE,
Arguments

formula  Object of class formula or character describing the model to fit. Interaction
terms supported only for numerical variables.

data  Training data of class data.frame, matrix, dgCMatrix (Matrix) or gwaa.data
(GenABEL).

num.trees  Number of trees.
mtry  Number of variables to possibly split at in each node. Default is the (rounded
down) square root of the number variables. Alternatively, a single argument
function returning an integer, given the number of independent variables.

importance  Variable importance mode, one of ’none’, ’impurity’, ’impurity_corrected’, ’per-
mutation’. The ’impurity’ measure is the Gini index for classification, the vari-
ance of the responses for regression and the sum of test statistics (see splitrule)
for survival.

write.forest  Save ranger.forest object, required for prediction. Set to FALSE to reduce
memory usage if no prediction intended.

probability  Grow a probability forest as in Malley et al. (2012).

min.node.size  Minimal node size. Default 1 for classification, 5 for regression, 3 for survival,
and 10 for probability.

max.depth  Maximal tree depth. A value of NULL or 0 (the default) corresponds to unlim-
ited depth, 1 to tree stumps (1 split per tree).

replace  Sample with replacement.

sample.fraction  Fraction of observations to sample. Default is 1 for sampling with replacement
and 0.632 for sampling without replacement. For classification, this can be a
vector of class-specific values.

case.weights  Weights for sampling of training observations. Observations with larger weights
will be selected with higher probability in the bootstrap (or subsampled) samples
for the trees.

class.weights  Weights for the outcome classes (in order of the factor levels) in the splitting
rule (cost sensitive learning). Classification and probability prediction only. For
classification the weights are also applied in the majority vote in terminal nodes.
splitrule

Splitting rule. For classification and probability estimation "gini", "extratrees" or "hellinger" with default "gini". For regression "variance", "extratrees", "maxstat" or "beta" with default "variance". For survival "logrank", "extratrees", "C" or "maxstat" with default "logrank".

num.random.splits

For "extratrees" splitrule.: Number of random splits to consider for each candidate splitting variable.

alpha

For "maxstat" splitrule: Significance threshold to allow splitting.

minprop

For "maxstat" splitrule: Lower quantile of covariate distribution to be considered for splitting.

split.select.weights

Numeric vector with weights between 0 and 1, representing the probability to select variables for splitting. Alternatively, a list of size num.trees, containing split select weight vectors for each tree can be used.

always.split.variables

Character vector with variable names to be always selected in addition to the mtry variables tried for splitting.

respect.unordered.factors

Handling of unordered factor covariates. One of 'ignore', 'order' and 'partition'. For the "extratrees" splitrule the default is 'partition' for all other splitrules 'ignore'. Alternatively TRUE (='order') or FALSE (='ignore') can be used. See below for details.

scale.permutation.importance

Scale permutation importance by standard error as in (Breiman 2001). Only applicable if permutation variable importance mode selected.

local.importance

Calculate and return local importance values as in (Breiman 2001). Only applicable if importance is set to 'permutation'.

regularization.factor

Regularization factor (gain penalization), either a vector of length p or one value for all variables.

regularization.usedepth

Consider the depth in regularization.

keep.inbag

Save how often observations are in-bag in each tree.

inbag

Manually set observations per tree. List of size num.trees, containing inbag counts for each observation. Can be used for stratified sampling.

holdout

Hold-out mode. Hold-out all samples with case weight 0 and use these for variable importance and prediction error.

quantreg

Prepare quantile prediction as in quantile regression forests (Meinshausen 2006). Regression only. Set keep.inbag = TRUE to prepare out-of-bag quantile prediction.

oob.error

Compute OOB prediction error. Set to FALSE to save computation time, e.g. for large survival forests.

num.threads

Number of threads. Default is number of CPUs available.
save.memory: Use memory saving (but slower) splitting mode. No effect for survival and GWAS data. Warning: This option slows down the tree growing, use only if you encounter memory problems.

verbose: Show computation status and estimated runtime.

seed: Random seed. Default is NULL, which generates the seed from R. Set to 0 to ignore the R seed.

dependent.variable.name: Name of dependent variable, needed if no formula given. For survival forests this is the time variable.

status.variable.name: Name of status variable, only applicable to survival data and needed if no formula given. Use 1 for event and 0 for censoring.

classification: Set to TRUE to grow a classification forest. Only needed if the data is a matrix or the response numeric.

x: Predictor data (independent variables), alternative interface to data with formula or dependent.variable.name.

y: Response vector (dependent variable), alternative interface to data with formula or dependent.variable.name. For survival use a Surv() object or a matrix with time and status.

...: Further arguments passed to or from other methods (currently ignored).

Details

The tree type is determined by the type of the dependent variable. For factors classification trees are grown, for numeric values regression trees and for survival objects survival trees. The Gini index is used as default splitting rule for classification. For regression, the estimated response variances or maximally selected rank statistics (Wright et al. 2016) can be used. For Survival the log-rank test, a C-index based splitting rule (Schmid et al. 2015) and maximally selected rank statistics (Wright et al. 2016) are available. For all tree types, forests of extremely randomized trees (Geurts et al. 2006) can be grown.

With the probability option and factor dependent variable a probability forest is grown. Here, the node impurity is used for splitting, as in classification forests. Predictions are class probabilities for each sample. In contrast to other implementations, each tree returns a probability estimate and these estimates are averaged for the forest probability estimate. For details see Malley et al. (2012). Note that for classification and regression nodes with size smaller than min.node.size can occur, as in original Random Forests. For survival all nodes contain at least min.node.size samples. Variables selected with always.split.variables are tried additionally to the mtry variables randomly selected. In split.select.weights, weights do not need to sum up to 1, they will be normalized later. The weights are assigned to the variables in the order they appear in the formula or in the data if no formula is used. Names of the split.select.weights vector are ignored. The usage of split.select.weights can increase the computation times for large forests.

Unordered factor covariates can be handled in 3 different ways by using respect.unordered.factors: For ‘ignore’ all factors are regarded ordered, for ‘partition’ all possible 2-partitions are considered for splitting. For ‘order’ and 2-class classification the factor levels are ordered by their proportion falling in the second class, for regression by their mean response, as described in Hastie et al. (2009), chapter 9.2.4. For multiclass classification the factor levels are ordered by the first principal
component of the weighted covariance matrix of the contingency table (Coppersmith et al. 1999), for survival by the median survival (or the largest available quantile if the median is not available). The use of 'order' is recommended, as it computationally fast and can handle an unlimited number of factor levels. Note that the factors are only reordered once and not again in each split.

The 'impurity_corrected' importance measure is unbiased in terms of the number of categories and category frequencies and is almost as fast as the standard impurity importance. It is a modified version of the method by Sandri & Zuccolotto (2008), which is faster and more memory efficient. See Nembrini et al. (2018) for details. This importance measure can be combined with the methods to estimate p-values in `importance_pvalues`.

Regularization works by penalizing new variables by multiplying the splitting criterion by a factor, see Deng & Runger (2012) for details. If `regularization.usedepth=TRUE`, \( f^d \) is used, where \( f \) is the regularization factor and \( d \) the depth of the node. If regularization is used, multithreading is deactivated because all trees need access to the list of variables that are already included in the model.

For a large number of variables and data frames as input data the formula interface can be slow or impossible to use. Alternatively dependent `variable.name` (and status `variable.name` for survival) or `x` and `y` can be used. Use `x` and `y` with a matrix for `x` to avoid conversions and save memory. Consider setting `save.memory = TRUE` if you encounter memory problems for very large datasets, but be aware that this option slows down the tree growing.

For GWAS data consider combining `ranger` with the GenABEL package. See the Examples section below for a demonstration using Plink data. All SNPs in the GenABEL object will be used for splitting. To use only the SNPs without sex or other covariates from the phenotype file, use 0 on the right hand side of the formula. Note that missing values are treated as an extra category while splitting.

See https://github.com/imbs-hl/ranger for the development version.

With recent R versions, multithreading on Windows platforms should just work. If you compile yourself, the new RTools toolchain is required.

**Value**

Object of class `ranger` with elements

- **forest**: Saved forest (If `write.forest` set to `TRUE`). Note that the variable IDs in the split.varIDs object do not necessarily represent the column number in R.
- **predictions**: Predicted classes/values, based on out of bag samples (classification and regression only).
- **variable.importance**: Variable importance for each independent variable.
- **variable.importance.local**: Variable importance for each independent variable and each sample, if `local.importance` is set to `TRUE` and `importance` is set to 'permutation'.
- **prediction.error**: Overall out of bag prediction error. For classification this is the fraction of misclassified samples, for probability estimation the Brier score, for regression the mean squared error and for survival one minus Harrell’s C-index.
r.squared  R squared. Also called explained variance or coefficient of determination (regression only). Computed on out of bag data.

confusion.matrix
Contingency table for classes and predictions based on out of bag samples (classification only).

unique.death.times
Unique death times (survival only).

chf
Estimated cumulative hazard function for each sample (survival only).

survival
Estimated survival function for each sample (survival only).

call
Function call.

num.trees
Number of trees.

num.independent.variables
Number of independent variables.

mtry
Value of mtry used.

min.node.size
Value of minimal node size used.

treetype
Type of forest/tree. classification, regression or survival.

importance.mode
Importance mode used.

num.samples
Number of samples.

inbag.counts
Number of times the observations are in-bag in the trees.

Author(s)
Marvin N. Wright

References


- Deng & Runger (2012). Feature selection via regularized trees. The 2012 International Joint Conference on Neural Networks (IJCNN), Brisbane, Australia. doi: 10.1109/IJCNN.2012.6252640.

See Also
dpredict.ranger

Examples

### Classification forest with default settings
ranger(Species ~ ., data = iris)

```r
train.idx <- sample(nrow(iris), 2/3 * nrow(iris))
iris.train <- iris[train.idx, ]
iris.test <- iris[-train.idx, ]
rg.iris <- ranger(Species ~ ., data = iris.train)
pred.iris <- predict(rg.iris, data = iris.test)
table(iris.test$Species, pred.iris$predictions)
```

### Quantile regression forest

```r
rf <- ranger(mpg ~ ., mtcars[1:26, ], quantreg = TRUE)
pred <- predict(rf, mtcars[27:32, ], type = "quantiles")
pred$predictions
```

### Variable importance

```r
rg.iris <- ranger(Species ~ ., data = iris, importance = "impurity")
rg.iris$variable.importance
```

### Survival forest

```r
require(survival)
rg.veteran <- ranger(Surv(time, status) ~ ., data = veteran)
plot(rg.veteran$unique.death.times, rg.veteran$survival[1,])
```

### Alternative interfaces (same results)
ranger(dependent.variable.name = "Species", data = iris)
ranger(y = iris[, 5], x = iris[, -5])

### Not run:
## Use GenABEL interface to read Plink data into R and grow a classification forest
## The ped and map files are not included
library(GenABEL)
convert.snp.ped("data.ped", "data.map", "data.raw")
dat.gwaa <- load.gwaa.data("data.pheno", "data.raw")
phdata(dat.gwaa)$trait <- factor(phdata(dat.gwaa)$trait)
ranger(trait ~ ., data = dat.gwaa)

## End(Not run)

---

**timepoints.ranger**

*Ranger timepoints*

### Description

Extract unique death times of Ranger Survival forest

### Usage

```r
## S3 method for class 'ranger'
timepoints(x, ...)
```

### Arguments

- `x` Ranger Survival forest object.
- `...` Further arguments passed to or from other methods.

### Value

Unique death times

### Author(s)

Marvin N. Wright

### See Also

`ranger`
timepoints.ranger.prediction

Ranger timepoints

Description

Extract unique death times of Ranger Survival prediction object.

Usage

## S3 method for class 'ranger.prediction'
timepoints(x, ...)

Arguments

x Ranger Survival prediction object.
...
Further arguments passed to or from other methods.

Value

Unique death times

Author(s)

Marvin N. Wright

See Also

ranger

treeInfo

Tree information in human readable format

Description

Extract tree information of a ranger object.

Usage

treeInfo(object, tree = 1)

Arguments

object ranger object.
tree Number of the tree of interest.
Details

Node and variable ID's are 0-indexed, i.e., node 0 is the root node. If the formula interface is used in the `ranger` call, the variable ID's are usually different to the original data used to grow the tree. Refer to the variable name instead to be sure.

Splitting at unordered factors (nominal variables) depends on the option `respect.unordered.factors` in the `ranger` call. For the "ignore" and "order" approaches, all values smaller or equal the `splitval` value go to the left and all values larger go to the right, as usual. However, with "order" the values correspond to the order in `object$forest$covariate.levels` instead of the original order (usually alphabetical). In the "partition" mode, the `splitval` values for unordered factor are comma separated lists of values, representing the factor levels (in the original order) going to the right.

Value

A data.frame with the columns

- `nodeID` The nodeID, 0-indexed.
- `leftChild` ID of the left child node, 0-indexed.
- `rightChild` ID of the right child node, 0-indexed.
- `splitvarID` ID of the splitting variable, 0-indexed. Caution, the variable order changes if the formula interface is used.
- `splitvarName` Name of the splitting variable.
- `splitval` The splitting value. For numeric or ordinal variables, all values smaller or equal go to the left, larger values to the right.
- `terminal` Logical, TRUE for terminal nodes.
- `prediction` One column with the predicted class (factor) for classification and the predicted numerical value for regression.

Author(s)

Marvin N. Wright

See Also

`ranger`

Examples

```
rf <- ranger(Species ~ ., data = iris)
treeInfo(rf, 1)
```
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