Package 'randomForestSRC'

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BugReports https://github.com/kogalur/randomForestSRC/issues/

Depends R (>= 4.3.0),

Imports parallel, data.tree, DiagrammeR

Suggests survival, pec, prodlim, mlbench, interp, caret, imbalance, cluster, fst, data.table

Description Fast OpenMP parallel computing of Breiman's random forests for univariate, multivariate, unsupervised, survival, competing risks, class imbalanced classification and quantile regression. New Mahalanobis splitting for correlated outcomes. Extreme random forests and randomized splitting. Suite of imputation methods for missing data. Fast random forests using subsampling. Confidence regions and standard errors for variable importance. New improved holdout importance. Case-specific importance. Minimal depth variable importance. Visualize trees on your Safari or Google Chrome browser. Anonymous random forests for data privacy.

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URL https://www.randomforestsrc.org/https://ishwaran.org/

NeedsCompilation yes

Repository CRAN

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randomForestSRC-package

Fast Unified Random Forests for Survival, Regression, and Classification (RF-SRC)

Description

Fast OpenMP-parallel implementation of Breiman's random forests (Breiman, 2001) for regression, classification, survival analysis (Ishwaran, 2008), competing risks (Ishwaran, 2012), multivariate outcomes (Segal and Xiao, 2011), unsupervised learning (Mantero and Ishwaran, 2020), quantile regression (Meinshausen, 2006; Zhang et al., 2019; Greenwald and Khanna, 2001), and imbalanced q-classification (O'Brien and Ishwaran, 2019).

Supports deterministic and randomized splitting rules (Geurts et al., 2006; Ishwaran, 2015) across all families. Variable importance (VIMP), holdout VIMP, and confidence regions (Ishwaran and Lu, 2019) can be computed for single and grouped variables. Includes minimal depth variable selection (Ishwaran et al., 2010, 2011) and a fast interface for missing data imputation using multiple forest-based methods (Tang and Ishwaran, 2017).

Tree structures can be visualized in Safari or Chrome for any family; see get.tree.

Package Overview

This package contains many useful functions. Users are encouraged to read the help files in full for detailed guidance. Below is a brief overview of key functions to help navigate the package.

1. rfsrc

The main entry point to the package. Builds a random forest using user-supplied training data. The returned object is of class (rfsrc, grow).

2. rfsrc.fast

A computationally efficient version of rfsrc using subsampling.

3. quantreg.rfsrc, quantreg

Univariate and multivariate quantile regression forests for training and testing. Includes methods such as the Greenwald-Khanna (2001) algorithm, ideal for large data due to its memory efficiency.

4. predict.rfsrc, predict

Predicts outcomes by dropping test data down the trained forest. Returns an object of class (rfsrc, predict).

5. sidClustering.rfsrc, sidClustering

Unsupervised clustering using SID (Staggered Interaction Data). Also includes Breiman's artificial two-class method (Breiman, 2003).

6. vimp, subsample, holdout.vimp

Functions for variable selection and importance assessment:

- (a) vimp: Computes variable importance (VIMP) by perturbing each variable (e.g., via permutation). Can also be computed directly in rfsrc and predict.rfsrc.
- (b) subsample: Computes confidence intervals for VIMP using subsampling.
- (c) holdout.vimp: Measures the effect of removing a variable from the model.
- (d) **VarPro** (**VarPro** package): For advanced model-independent variable selection using rule-based variable priority. Supports regression, classification, survival, and unsupervised data. See https://www.varprotools.org.
- 7. imbalanced.rfsrc, imbalanced

Implements q-classification and G-mean-based VIMP for class-imbalanced data.

8. impute.rfsrc, impute

A fast interface for missing data imputation. While rfsrc and predict.rfsrc can handle missing data internally, this provides a dedicated, efficient solution for imputation tasks.

9. partial.rfsrc, partial

Computes partial dependence functions to assess the marginal effect of one or more variables on the forest ensemble.

Home page, Vignettes, Discussions, Bug Reporting, Source Code, Beta Builds

- 1. The package home page, with vignettes, manuals, GitHub links, and additional documentation, is available at: https://www.randomforestsrc.org/index.html
- 2. Questions, comments, and general usage discussions (non-bug-related) can be posted at: https://github.com/kogalur/randomForestSRC/discussions/
- Bug reports should be submitted at: https://github.com/kogalur/randomForestSRC/ issues/

Please use this only for bugs, and include the following with your report:

- Output from sessionInfo().
- A minimal reproducible example including:
 - A minimal dataset required to reproduce the error.
 - The smallest runnable code needed to reproduce the issue.
 - Version details of R and all relevant packages.
 - A random seed (via set.seed()) if randomness is involved.
- 4. The latest stable release of the package is available on CRAN: https://cran.r-project.org/package=randomForestSRC/
- 5. Development builds (unstable) with bug fixes and new features are hosted on GitHub: https: //github.com/kogalur/randomForestSRC/

OpenMP Parallel Processing – Installation

This package supports OpenMP shared-memory parallel programming on systems where the architecture and operating system permit it. OpenMP is enabled by default.

Detailed instructions for configuring OpenMP parallel processing can be found at: https://www.randomforestsrc.org/articles/installation.html

Note that running the package with OpenMP (or Open MPI) may increase memory (RAM) usage. Users are advised to understand their system's hardware limits and to monitor resource consumption to avoid overtaxing CPU and memory capacity.

Reproducibility

Model reproducibility is determined by three components: the random seed, the forest topology (i.e., the structure of trees), and terminal node membership for the training data. These elements together allow the model and its terminal node statistics to be faithfully restored.

Other outputs, such as variable importance (VIMP) and performance metrics, rely on additional internal randomization and are not considered part of the model definition. As a result, such statistics are subject to Monte Carlo variability and may differ across runs, even with the same seed.

Author(s)

Hemant Ishwaran and Udaya B. Kogalur

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Segal M.R. and Xiao Y. Multivariate random forests. (2011). *Wiley Interdisciplinary Reviews: Data Mining and Knowledge Discovery*. 1(1):80-87.

Tang F. and Ishwaran H. (2017). Random forest missing data algorithms. *Statistical Analysis and Data Mining*, 10:363-377.

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See Also

find.interaction.rfsrc,

get.tree.rfsrc,

holdout.vimp.rfsrc,

imbalanced.rfsrc, impute.rfsrc,

max.subtree.rfsrc,

partial.rfsrc, plot.competing.risk.rfsrc, plot.rfsrc, plot.survival.rfsrc, plot.variable.rfsrc, predict.rfsrc, print.rfsrc,

quantreg.rfsrc,

rfsrc, rfsrc.cart, rfsrc.fast,

sidClustering.rfsrc,

stat.split.rfsrc, subsample.rfsrc, synthetic.rfsrc,

tune.rfsrc,

vimp.rfsrc

breast

Wisconsin Prognostic Breast Cancer Data

Description

Recurrence of breast cancer from 198 breast cancer patients, all of which exhibited no evidence of distant metastases at the time of diagnosis. The first 30 features of the data describe characteristics of the cell nuclei present in the digitized image of a fine needle aspirate (FNA) of the breast mass.

Source

The data were obtained from the UCI machine learning repository, see http://archive.ics.uci.edu/ml/datasets/Breast+Cancer+Wisconsin+(Prognostic).

Examples

```
## ------
## Standard analysis
## ------
data(breast, package = "randomForestSRC")
breast <- na.omit(breast)
o <- rfsrc(status ~ ., data = breast, nsplit = 10)
print(o)</pre>
```

find.interaction.rfsrc

Find Interactions Between Pairs of Variables

Description

Find pairwise interactions between variables.

Usage

```
## S3 method for class 'rfsrc'
find.interaction(object, xvar.names, cause, m.target,
    importance = c("permute", "random", "anti",
                          "permute.ensemble", "random.ensemble", "anti.ensemble"),
    method = c("maxsubtree", "vimp"), sorted = TRUE, nvar, nrep = 1,
    na.action = c("na.omit", "na.impute", "na.random"),
    seed = NULL, do.trace = FALSE, verbose = TRUE, ...)
```

Arguments

| object | An object of class (rfsrc, grow) or (rfsrc, forest). |
|------------|--|
| xvar.names | Character vector of names of target x-variables. Default is to use all variables. |
| cause | For competing risk families, integer value between 1 and J indicating the event of interest, where J is the number of event types. The default is to use the first event type. |
| m.target | Character value for multivariate families specifying the target outcome to be used. If left unspecified, the algorithm will choose a default target. |
| importance | Type of variable importance (VIMP). See rfsrc for details. |
| method | Method of analysis: maximal subtree or VIMP. See details below. |
| sorted | Should variables be sorted by VIMP? Does not apply for competing risks. |
| nvar | Number of variables to be used. |
| nrep | Number of Monte Carlo replicates when 'method="vimp"'. |
| na.action | Action to be taken if the data contains NA values. Applies only when `method="vimp"'. |
| seed | Seed for random number generator. Must be a negative integer. |
| do.trace | Number of seconds between updates to the user on approximate time to com- pletion. |
| verbose | Set to TRUE for verbose output. |
| | Further arguments passed to or from other methods. |

Details

Using a previously grown forest, identify pairwise interactions for all pairs of variables from a specified list. There are two distinct approaches specified by the option 'method'.

'method="maxsubtree"'

This invokes a maximal subtree analysis. In this case, a matrix is returned where entries [i][i] are the normalized minimal depth of variable [i] relative to the root node (normalized wrt the size of the tree) and entries [i][j] indicate the normalized minimal depth of a variable [j] wrt the maximal subtree for variable [i] (normalized wrt the size of [i]'s maximal subtree). Smaller [i][i] entries indicate predictive variables. Small [i][j] entries having small [i][i] entries are a sign of an interaction between variable i and j (note: the user should scan rows, not columns, for small entries). See Ishwaran et al. (2010, 2011) for more details.

2. 'method="vimp"

This invokes a joint-VIMP approach. Two variables are paired and their paired VIMP calculated (refered to as 'Paired' importance). The VIMP for each separate variable is also calculated. The sum of these two values is referred to as 'Additive' importance. A large positive or negative difference between 'Paired' and 'Additive' indicates an association worth pursuing if the univariate VIMP for each of the paired-variables is reasonably large. See Ishwaran (2007) for more details.

Computations might be slow depending upon the size of the data and the forest. In such cases, consider setting 'nvar' to a smaller number. If 'method="maxsubtree"', consider using a smaller number of trees in the original grow call.

If 'nrep' is greater than 1, the analysis is repeated nrep times and results averaged over the replications (applies only when 'method="vimp").

Value

Invisibly, the interaction table (a list for competing risk data) or the maximal subtree matrix.

Author(s)

Hemant Ishwaran and Udaya B. Kogalur

References

Ishwaran H. (2007). Variable importance in binary regression trees and forests, *Electronic J. Statist.*, 1:519-537.

Ishwaran H., Kogalur U.B., Gorodeski E.Z, Minn A.J. and Lauer M.S. (2010). High-dimensional variable selection for survival data. *J. Amer. Statist. Assoc.*, 105:205-217.

Ishwaran H., Kogalur U.B., Chen X. and Minn A.J. (2011). Random survival forests for highdimensional data. *Statist. Anal. Data Mining*, 4:115-132.

See Also

holdout.vimp.rfsrc,max.subtree.rfsrc,vimp.rfsrc

follic

Examples

```
## find interactions, survival setting
data(pbc, package = "randomForestSRC")
pbc.obj <- rfsrc(Surv(days,status) ~ ., pbc, importance = TRUE)</pre>
find.interaction(pbc.obj, method = "vimp", nvar = 8)
## -----
## find interactions, competing risks
data(wihs, package = "randomForestSRC")
wihs.obj <- rfsrc(Surv(time, status) ~ ., wihs, nsplit = 3, ntree = 100,
                importance = TRUE)
find.interaction(wihs.obj)
find.interaction(wihs.obj, method = "vimp")
## -----
## find interactions, regression setting
## ______
airq.obj <- rfsrc(Ozone ~ ., data = airquality, importance = TRUE)</pre>
find.interaction(airq.obj, method = "vimp", nrep = 3)
find.interaction(airq.obj)
## find interactions, classification setting
## -----
iris.obj <- rfsrc(Species ~., data = iris, importance = TRUE)</pre>
find.interaction(iris.obj, method = "vimp", nrep = 3)
find.interaction(iris.obj)
## interactions for multivariate mixed forests
## -----
mtcars2 <- mtcars</pre>
mtcars2$cyl <- factor(mtcars2$cyl)</pre>
mtcars2$carb <- factor(mtcars2$carb, ordered = TRUE)</pre>
mv.obj <- rfsrc(cbind(carb, mpg, cyl) ~., data = mtcars2, importance = TRUE)</pre>
find.interaction(mv.obj, method = "vimp", outcome.target = "carb")
find.interaction(mv.obj, method = "vimp", outcome.target = "mpg")
find.interaction(mv.obj, method = "vimp", outcome.target = "cyl")
```

follic

Follicular Cell Lymphoma

Description

Competing risk data set involving follicular cell lymphoma.

Format

A data frame containing:

| age | age |
|---------|--|
| hgb | hemoglobin (g/l) |
| clinstg | clinical stage: 1=stage I, 2=stage II |
| ch | chemotherapy |
| rt | radiotherapy |
| time | first failure time |
| status | censoring status: 0=censored, 1=relapse, 2=death |

Source

Table 1.4b, Competing Risks: A Practical Perspective.

References

Pintilie M., (2006) Competing Risks: A Practical Perspective. West Sussex: John Wiley and Sons.

Examples

```
data(follic, package = "randomForestSRC")
follic.obj <- rfsrc(Surv(time, status) ~ ., follic, nsplit = 3, ntree = 100)</pre>
```

get.tree.rfsrc Extract a Single Tree from a Forest and plot it on your browser

Description

Extracts a single tree from a forest which can then be plotted on the users browser. Works for all families. Missing data not permitted.

Usage

```
## S3 method for class 'rfsrc'
get.tree(object, tree.id, target, m.target = NULL,
    time, surv.type = c("mort", "rel.freq", "surv", "years.lost", "cif", "chf"),
    class.type = c("bayes", "rfq", "prob"),
    ensemble = FALSE, oob = TRUE, show.plots = TRUE, do.trace = FALSE)
```

get.tree.rfsrc

Arguments

| ass of interest (defaults to 1 and J (number of event rst event type). |
|--|
| ltivariate families. If un- |
| is evaluated (depends on |
| rned. See Details. |
| e. See Details. |
| ole of all trees. If FALSE |
| values (FALSE). Only ap- |
| |
| |
| |

Details

Extracts a specified tree from a forest and converts it into a hierarchical structure compatible with the **data.tree** package. Plotting the resulting object renders an interactive tree visualization in the user's web browser.

Left-hand splits are shown. For continuous variables, the left split is displayed as an inequality (e.g., x < value); the right split is the reverse. For factor variables, the left daughter node is defined by a set of levels assigned to it; the right daughter is its complement.

Terminal nodes are highlighted with color and display both sample size and predicted value. By default, the predicted value corresponds to the prediction from the selected tree, and the sample size refers to the in-bag cases reaching the terminal node. If ensemble = TRUE, the predicted value equals the forest ensemble prediction, allowing visualization of the full forest predictor over the selected tree's partition. In this case, sample sizes refer to all observations (not just in-bag cases).

Predicted values displayed in terminal nodes are defined as follows:

- 1. For regression: the mean of the response.
- 2. For classification: depends on the class.type argument and target class:
 - If class.type = "bayes", the predicted class with the most votes, or the RFQ classifier threshold in two-class problems.
 - If class.type = "prob", the class probability for the target class.
- 3. For multivariate families: the predicted value for the outcome specified by m.target, using the logic above depending on whether the outcome is continuous or categorical.
- 4. For survival:
 - mort: estimated mortality (Ishwaran et al., 2008).

- rel.freq: relative frequency of mortality.
- surv: predicted survival probability at the specified time (time).
- 5. For competing risks:
 - years.lost: expected number of life years lost.
 - cif: cumulative incidence function.
 - chf: cause-specific cumulative hazard function.

For cif and chf, predictions are evaluated at the time point given by time, and all metrics are specific to the event type indicated by target.

Value

Invisibly, returns an object with hierarchical structure formatted for use with the data.tree package.

Author(s)

Hemant Ishwaran and Udaya B. Kogalur

Many thanks to @dbarg1 on GitHub for the initial prototype of this function

Examples

```
## ______
## survival/competing risk
## -----
## survival - veteran data set but with factors
## note that diagtime has many levels
data(veteran, package = "randomForestSRC")
vd <- veteran
vd$celltype=factor(vd$celltype)
vd$diagtime=factor(vd$diagtime)
vd.obj <- rfsrc(Surv(time,status)~., vd, ntree = 100, nodesize = 5)</pre>
plot(get.tree(vd.obj, 3))
## competing risks
data(follic, package = "randomForestSRC")
follic.obj <- rfsrc(Surv(time, status) ~ ., follic, nsplit = 3, ntree = 100)</pre>
plot(get.tree(follic.obj, 2))
## -----
## regression
## ______
airq.obj <- rfsrc(Ozone ~ ., data = airquality)</pre>
plot(get.tree(airq.obj, 10))
## -----
## two-class imbalanced data (see imbalanced function)
```

data(breast, package = "randomForestSRC")

get.tree.rfsrc

```
breast <- na.omit(breast)</pre>
f <- as.formula(status ~ .)</pre>
breast.obj <- imbalanced(f, breast)</pre>
## compare RFQ to Bayes Rule
plot(get.tree(breast.obj, 1, class.type = "rfq", ensemble = TRUE))
plot(get.tree(breast.obj, 1, class.type = "bayes", ensemble = TRUE))
## ______
## classification
## ------
iris.obj <- rfsrc(Species ~., data = iris, nodesize = 10)</pre>
## equivalent
plot(get.tree(iris.obj, 25))
plot(get.tree(iris.obj, 25, class.type = "bayes"))
## predicted probability displayed for terminal nodes
plot(get.tree(iris.obj, 25, class.type = "prob", target = "setosa"))
plot(get.tree(iris.obj, 25, class.type = "prob", target = "versicolor"))
plot(get.tree(iris.obj, 25, class.type = "prob", target = "virginica"))
## multivariate regression
## -----
mtcars.mreg <- rfsrc(Multivar(mpg, cyl) ~., data = mtcars)</pre>
plot(get.tree(mtcars.mreg, 10, m.target = "mpg"))
plot(get.tree(mtcars.mreg, 10, m.target = "cyl"))
## -----
## multivariate mixed outcomes
## ------
mtcars2 <- mtcars</pre>
mtcars2$carb <- factor(mtcars2$carb)</pre>
mtcars2$cyl <- factor(mtcars2$cyl)</pre>
mtcars.mix <- rfsrc(Multivar(carb, mpg, cyl) ~ ., data = mtcars2)</pre>
plot(get.tree(mtcars.mix, 5, m.target = "cyl"))
plot(get.tree(mtcars.mix, 5, m.target = "carb"))
## -----
## unsupervised analysis
## -----
mtcars.unspv <- rfsrc(data = mtcars)</pre>
```

plot(get.tree(mtcars.unspv, 5))

Hodgkin's Disease

Description

Competing risk data set involving Hodgkin's disease.

Format

A data frame containing:

| age | age |
|----------|---|
| sex | gender |
| trtgiven | treatment: RT=radition, CMT=Chemotherapy and radiation |
| medwidsi | mediastinum involvement: N=no, S=small, L=Large |
| extranod | extranodal disease: Y=extranodal disease, N=nodal disease |
| clinstg | clinical stage: 1=stage I, 2=stage II |
| time | first failure time |
| status | censoring status: 0=censored, 1=relapse, 2=death |

Source

Table 1.6b, Competing Risks: A Practical Perspective.

References

Pintilie M., (2006) Competing Risks: A Practical Perspective. West Sussex: John Wiley and Sons.

Examples

```
data(hd, package = "randomForestSRC")
```

holdout.vimp.rfsrc *Hold out variable importance (VIMP)*

Description

Hold out VIMP is calculated from the error rate of mini ensembles of trees (blocks of trees) grown with and without a variable. Applies to all families.

hd

holdout.vimp.rfsrc

Usage

```
## S3 method for class 'rfsrc'
holdout.vimp(formula, data,
   ntree = function(p, vtry){1000 * p / vtry},
   nsplit = 10,
   ntime = 50,
   sampsize = function(x){x * .632},
   samptype = "swor",
   block.size = 10,
   vtry = 1,
   ...)
```

Arguments

| formula | A symbolic description of the model to be fit. |
|------------|--|
| data | Data frame containing the y-outcome and x-variables. |
| ntree | Specifies the number of trees used to grow the forest. Can be a function of data dimension and number of holdout variables, or a fixed numeric value. |
| nsplit | Non-negative integer specifying the number of random split points used to split a node. A value of zero corresponds to deterministic splitting, which is signifi- cantly slower. |
| ntime | Integer value used for survival settings to constrain ensemble calculations to a grid of ntime time points. |
| sampsize | Specifies the size of the subsampled data. Can be either a function or a numeric value. |
| samptype | Type of bootstrap used when subsampling. |
| vtry | Number of variables randomly selected to be held out when growing a tree. Can also be a list for targeted holdout variable importance analysis. See details for more information. |
| block.size | Specifies the number of trees in a block when calculating holdout variable importance. |
| | Further arguments passed to rfsrc. |

Details

Holdout variable importance (holdout VIMP) measures the importance of a variable by comparing prediction error between two forests (blocks of trees): one in which selected variables are held out during tree growing (the *holdout forest*) and one in which no variables are held out (the *baseline forest*).

For each variable-block combination, the bootstrap samples used to grow the trees are the same in both forests. The difference in out-of-bag (OOB) prediction error between the holdout and baseline forests gives the holdout VIMP for that variable-block pair. The final holdout VIMP for a variable is the average of these differences over all blocks in which the variable was held out.

The option vtry controls how many variables are held out per tree. The default is one, meaning a single variable is held out per tree. Larger values of vtry increase the number of times each variable

is held out, reducing the required total number of trees. However, interpretation of holdout VIMP changes when vtry exceeds one, and this option should be used cautiously.

High accuracy requires a sufficiently large number of trees. As a general guideline, we recommend using ntree = 1000 * p / vtry, where p is the number of features. Accuracy also depends on block.size, which determines how many trees comprise a block. Smaller values yield better accuracy but are computationally more demanding. The most accurate setting is block.size = 1. Ensure that block.size does not exceed ntree / p, otherwise insufficient trees may be available for certain variables.

Targeted holdout VIMP analysis can be requested by specifying vtry as a list with two components: a vector of variable indices (xvar) and a logical flag joint indicating whether to compute joint VIMP. For example, to compute holdout VIMP only for variables 1, 4, and 5 individually:

vtry = list(xvar = c(1, 4, 5), joint = FALSE)

To compute the joint effect of removing these three variables together:

vtry = list(xvar = c(1, 4, 5), joint = TRUE)

Targeted analysis is useful when the user has prior knowledge of variables of interest and can significantly reduce computation. Joint VIMP quantifies the combined importance of specific groups of variables. See the Iris example below for illustration.

Value

Invisibly a list with the following components (which themselves can be lists):

| importance | Holdout VIMP. |
|------------|---|
| baseline | Prediction error for the baseline forest. |
| holdout | Prediction error for the holdout forest. |

Author(s)

Hemant Ishwaran and Udaya B. Kogalur

References

Lu M. and Ishwaran H. (2018). Expert Opinion: A prediction-based alternative to p-values in regression models. *J. Thoracic and Cardiovascular Surgery*, 155(3), 1130–1136.

See Also

vimp.rfsrc

Examples

-----analysis ## regression analysis ## -----

new York air quality measurements

```
airq.obj <- holdout.vimp(Ozone ~ ., data = airquality, na.action = "na.impute")
print(airq.obj$importance)
## classification analysis
## -----
## iris data
iris.obj <- holdout.vimp(Species ~., data = iris)</pre>
print(iris.obj$importance)
## iris data using brier prediction error
iris.obj <- holdout.vimp(Species ~., data = iris, perf.type = "brier")</pre>
print(iris.obj$importance)
## ------
## illustration of targeted holdout vimp analysis
## ______
## iris data - only interested in variables 3 and 4
vtry <- list(xvar = c(3, 4), joint = FALSE)</pre>
print(holdout.vimp(Species ~., data = iris, vtry = vtry)$impor)
## iris data - joint importance of variables 3 and 4
vtry <- list(xvar = c(3, 4), joint = TRUE)</pre>
print(holdout.vimp(Species ~., data = iris, vtry = vtry)$impor)
## iris data - joint importance of variables 1 and 2
vtry <- list(xvar = c(1, 2), joint = TRUE)</pre>
print(holdout.vimp(Species ~., data = iris, vtry = vtry)$impor)
## -----
## imbalanced classification (using RFQ)
## -----
if (library("caret", logical.return = TRUE)) {
 ## experimental settings
 n <- 400
 q <- 20
 ir <- 6
 f <- as.formula(Class ~ .)</pre>
 ## simulate the data, create minority class data
 d <- twoClassSim(n, linearVars = 15, noiseVars = q)</pre>
 d$Class <- factor(as.numeric(d$Class) - 1)
 idx.0 <- which(d$Class == 0)</pre>
 idx.1 <- sample(which(d$Class == 1), sum(d$Class == 1) / ir , replace = FALSE)</pre>
 d <- d[c(idx.0,idx.1),, drop = FALSE]</pre>
 ## VIMP for RFQ with and without blocking
 vmp1 <- imbalanced(f, d, importance = TRUE, block.size = 1)$importance[, 1]</pre>
```

```
vmp10 <- imbalanced(f, d, importance = TRUE, block.size = 10)$importance[, 1]</pre>
 ## holdout VIMP for RFQ with and without blocking
 hvmp1 <- holdout.vimp(f, d, rfq = TRUE,</pre>
            perf.type = "g.mean", block.size = 1)$importance[, 1]
 hvmp10 <- holdout.vimp(f, d, rfq = TRUE,</pre>
            perf.type = "g.mean", block.size = 10)$importance[, 1]
 ## compare VIMP values
 imp <- 100 * cbind(vmp1, vmp10, hvmp1, hvmp10)</pre>
 legn <- c("vimp-1", "vimp-10", "hvimp-1", "hvimp-10")</pre>
 colr <- rep(4,20+q)
 colr[1:20] <- 2
 ylim <- range(c(imp))</pre>
 nms <- 1:(20+q)
 par(mfrow=c(2,2))
 barplot(imp[,1],col=colr,las=2,main=legn[1],ylim=ylim,names.arg=nms)
 barplot(imp[,2],col=colr,las=2,main=legn[2],ylim=ylim,names.arg=nms)
 barplot(imp[,3],col=colr,las=2,main=legn[3],ylim=ylim,names.arg=nms)
 barplot(imp[,4],col=colr,las=2,main=legn[4],ylim=ylim,names.arg=nms)
}
## ______
## multivariate regression analysis
## -----
mtcars.mreg <- holdout.vimp(Multivar(mpg, cyl) ~., data = mtcars,</pre>
                               vtry = 3,
                               block.size = 1,
                               samptype = "swr",
                               sampsize = dim(mtcars)[1])
print(mtcars.mreg$importance)
## ______
## mixed outcomes analysis
## ______
mtcars.new <- mtcars</pre>
mtcars.new$cyl <- factor(mtcars.new$cyl)</pre>
mtcars.new$carb <- factor(mtcars.new$carb, ordered = TRUE)</pre>
mtcars.mix <- holdout.vimp(cbind(carb, mpg, cyl) ~., data = mtcars.new,</pre>
                              ntree = 100,
                              block.size = 2,
                              vtry = 1)
print(mtcars.mix$importance)
##-----
## survival analysis
## Primary biliary cirrhosis (PBC) of the liver
data(pbc, package = "randomForestSRC")
pbc.obj <- holdout.vimp(Surv(days, status) ~ ., pbc,</pre>
```

housing

```
nsplit = 10,
ntree = 1000,
na.action = "na.impute")
print(pbc.obj$importance)
##-----
## competing risks
##-----
## WIHS analysis
## cumulative incidence function (CIF) for HAART and AIDS stratified by IDU
data(wihs, package = "randomForestSRC")
wihs.obj <- holdout.vimp(Surv(time, status) ~ ., wihs,
nsplit = 3,
ntree = 100)
print(wihs.obj$importance)
```

```
housing
```

Ames Iowa Housing Data

Description

Data from the Ames Assessor's Office used in assessing values of individual residential properties sold in Ames, Iowa from 2006 to 2010. This is a regression problem and the goal is to predict "SalePrice" which records the price of a home in thousands of dollars.

References

De Cock, D., (2011). Ames, Iowa: Alternative to the Boston housing data as an end of semester regression project. *Journal of Statistics Education*, 19(3), 1–14.

Examples

```
## load the data
data(housing, package = "randomForestSRC")
```

the original data contains lots of missing data, so impute it
use missForest, can be slow so grow trees with small training sizes
housing2 <- impute(data = housing, mf.q = 1, sampsize = function(x){x * .1})</pre>

```
## same idea ... but directly use rfsrc.fast and multivariate missForest
housing3 <- impute(data = housing, mf.q = .5, fast = TRUE)</pre>
```

```
## even faster, but potentially less acurate
housing4 <- impute(SalePrice~., housing, splitrule = "random", nimpute = 1)</pre>
```

imbalanced.rfsrc

Description

Implements various solutions to the two-class imbalanced problem, including the newly proposed quantile-classifier approach of O'Brien and Ishwaran (2017). Also includes Breiman's balanced random forests undersampling of the majority class. Performance is assessed using the G-mean, but misclassification error can be requested.

Usage

```
## S3 method for class 'rfsrc'
imbalanced(formula, data, ntree = 3000,
  method = c("rfq", "brf", "standard"), splitrule = "auc",
  perf.type = NULL, block.size = NULL, fast = FALSE,
  ratio = NULL, ...)
```

Arguments

| formula | A symbolic description of the model to be fit. |
|------------|---|
| data | A data frame containing the two-class y-outcome and x-variables. |
| ntree | Number of trees to grow. |
| method | Method used to fit the classifier. The default is "rfq", which implements the random forest quantile classifier (RFQ) of O'Brien and Ishwaran (2017). The option "brf" applies the balanced random forest (BRF) approach of Chen et al. (2004), which undersamples the majority class to match the minority class size. The option "standard" performs a standard random forest analysis. |
| splitrule | Splitting rule used to grow trees. The default is "auc", which optimizes G-mean performance. Other supported options are "gini" and "entropy". |
| perf.type | Performance metric used for evaluating the classifier and computing downstream quantities such as VIMP. Defaults depend on the method: "gmean" for RFQ and BRF; "misclass" (misclassification error) for standard random forests. Users may override this by specifying "gmean", "misclass", or "brier" (normalized Brier score). See examples for usage. |
| block.size | Controls how the cumulative error rate is computed. If NULL, it is calculated only once for the final tree. If set to an integer, cumulative error and VIMP are computed in blocks of that size. If unspecified, uses the default in rfsrc. |
| fast | Logical. If TRUE, uses the fast random forest implementation via rfsrc.fast instead of rfsrc. Improves speed at the cost of accuracy. Applies only to RFQ. |
| ratio | Optional and experimental. Specifies the proportion (between 0 and 1) of major- ity class cases to sample during RFQ training. Sampling is without replacement. Ignored for BRF. |
| | Additional arguments passed to rfsrc to control random forest behavior. |

imbalanced.rfsrc

Details

Imbalanced data, also known as the minority class problem, refers to two-class classification settings where the majority class significantly outnumbers the minority class. This function supports two approaches to address class imbalance:

- The random forests quantile classifier (RFQ) proposed by O'Brien and Ishwaran (2017).
- The balanced random forest (BRF) undersampling method of Chen et al. (2004).

By default, the performance metric used is the G-mean (Kubat et al., 1997), which balances sensitivity and specificity.

Handling of missing values: Missing data are not supported for BRF or when the ratio option is specified. In these cases, records with missing values are removed prior to analysis.

Variable importance: Permutation-based VIMP is used by default in this setting, in contrast to anti-VIMP which is the default for other families. Empirical results suggest that permutation VIMP is more reliable in highly imbalanced settings.

Tree count recommendation: We recommend using a relatively large value for ntree in imbalanced problems to ensure stable performance estimation, especially for G-mean. As a general guideline, use at least five times the usual number of trees.

Performance metrics: A helper function, get.imbalanced.performance, is provided for extracting classification performance summaries. The metric names are self-explanatory in most cases. Some key metrics include:

- F1: The harmonic mean of precision and recall.
- F1mod: The harmonic mean of sensitivity, specificity, precision, and negative predictive value.
- F1gmean: The average of F1 and G-mean.
- F1modgmean: The average of F1mod and G-mean.

Value

A two-class random forest fit under the requested method and performance value.

Author(s)

Hemant Ishwaran and Udaya B. Kogalur

References

Chen, C., Liaw, A. and Breiman, L. (2004). Using random forest to learn imbalanced data. University of California, Berkeley, Technical Report 110.

Kubat, M., Holte, R. and Matwin, S. (1997). Learning when negative examples abound. *Machine Learning*, ECML-97: 146-153.

O'Brien R. and Ishwaran H. (2019). A random forests quantile classifier for class imbalanced data. *Pattern Recognition*, 90, 232-249

See Also

rfsrc, rfsrc.fast

Examples

```
## use the breast data for illustration
## ______
data(breast, package = "randomForestSRC")
breast <- na.omit(breast)</pre>
f <- as.formula(status ~ .)</pre>
##------
## default RFQ call
o.rfq <- imbalanced(f, breast)</pre>
print(o.rfq)
## equivalent to:
## rfsrc(f, breast, rfq = TRUE, ntree = 3000,
     perf.type = "gmean", splitrule = "auc")
##
##------
## detailed output using customized performance function
print(get.imbalanced.performance(o.rfq))
## RF using misclassification error with gini splitting
o.std <- imbalanced(f, breast, method = "stand", splitrule = "gini")</pre>
##-----
## RF using G-mean performance with AUC splitting
## -----
o.std <- imbalanced(f, breast, method = "stand", perf.type = "gmean")</pre>
## equivalent to:
## rfsrc(f, breast, ntree = 3000, perf.type = "gmean", splitrule = "auc")
##-----
## default BRF call
##-----
o.brf <- imbalanced(f, breast, method = "brf")</pre>
## equivalent to:
## imbalanced(f, breast, method = "brf", perf.type = "gmean")
##-----
## BRF call with misclassification performance
```

imbalanced.rfsrc

```
##-----
o.brf <- imbalanced(f, breast, method = "brf", perf.type = "misclass")</pre>
##_____
## train/test example
##-----
trn <- sample(1:nrow(breast), size = nrow(breast) / 2)</pre>
o.trn <- imbalanced(f, breast[trn,], importance = TRUE)</pre>
o.tst <- predict(o.trn, breast[-trn,], importance = TRUE)</pre>
print(o.trn)
print(o.tst)
print(100 * cbind(o.trn$impo[, 1], o.tst$impo[, 1]))
##-----
##
## illustrates how to optimize threshold on training data
## improves Gmean for RFQ in many situations
##
##-----
if (library("caret", logical.return = TRUE)) {
 ## experimental settings
 n <- 2 * 5000
 q <- 20
 ir <- 6
 f <- as.formula(Class ~ .)</pre>
 ## simulate the data, create minority class data
 d <- twoClassSim(n, linearVars = 15, noiseVars = q)</pre>
 d$Class <- factor(as.numeric(d$Class) - 1)
 idx.0 <- which(d$Class == 0)</pre>
 idx.1 <- sample(which(d$Class == 1), sum(d$Class == 1) / ir , replace = FALSE)</pre>
 d <- d[c(idx.0,idx.1),, drop = FALSE]</pre>
 ## split data into train and test
 trn.pt <- sample(1:nrow(d), size = nrow(d) / 2)</pre>
 trn <- d[trn.pt, ]</pre>
 tst <- d[setdiff(1:nrow(d), trn.pt), ]</pre>
 ## run rfq on training data
 o <- imbalanced(f, trn)</pre>
 ## (1) default threshold (2) directly optimized gmean threshold
 th.1 <- get.imbalanced.performance(o)["threshold"]</pre>
 th.2 <- get.imbalanced.optimize(o)["threshold"]</pre>
 ## training performance
 cat("----- train performance -----\n")
 print(get.imbalanced.performance(o, thresh=th.1))
```

```
print(get.imbalanced.performance(o, thresh=th.2))
 ## test performance
 cat("----- test performance -----\n")
 pred.o <- predict(o, tst)</pre>
 print(get.imbalanced.performance(pred.o, thresh=th.1))
 print(get.imbalanced.performance(pred.o, thresh=th.2))
}
## illustrates RFQ with and without SMOTE
##
## - simulation example using the caret R-package
## - creates imbalanced data by randomly sampling the class 1 data
## - use SMOTE from "imbalance" package to oversample the minority
##
##_____
if (library("caret", logical.return = TRUE) &
   library("imbalance", logical.return = TRUE)) {
 ## experimental settings
 n <- 5000
 q <- 20
 ir <- 6
 f <- as.formula(Class ~ .)</pre>
 ## simulate the data, create minority class data
 d <- twoClassSim(n, linearVars = 15, noiseVars = q)</pre>
 d$Class <- factor(as.numeric(d$Class) - 1)
 idx.0 <- which(d$Class == 0)</pre>
 idx.1 <- sample(which(d$Class == 1), sum(d$Class == 1) / ir , replace = FALSE)
 d <- d[c(idx.0,idx.1),, drop = FALSE]</pre>
 d <- d[sample(1:nrow(d)), ]</pre>
 ## define train/test split
 trn <- sample(1:nrow(d), size = nrow(d) / 2, replace = FALSE)</pre>
 ## now make SMOTE training data
 newd.50 <- mwmote(d[trn, ], numInstances = 50, classAttr = "Class")</pre>
 newd.500 <- mwmote(d[trn, ], numInstances = 500, classAttr = "Class")</pre>
 ## fit RFQ with and without SMOTE
 o.with.50 <- imbalanced(f, rbind(d[trn, ], newd.50))</pre>
 o.with.500 <- imbalanced(f, rbind(d[trn, ], newd.500))</pre>
 o.without <- imbalanced(f, d[trn, ])</pre>
 ## compare performance on test data
 print(predict(o.with.50, d[-trn, ]))
 print(predict(o.with.500, d[-trn, ]))
 print(predict(o.without, d[-trn, ]))
```

imbalanced.rfsrc

```
}
##-----
##
## illustrates effectiveness of blocked VIMP
##
##-----
if (library("caret", logical.return = TRUE)) {
 ## experimental settings
 n <- 1000
 q <- 20
 ir <- 6
 f <- as.formula(Class ~ .)</pre>
 ## simulate the data, create minority class data
 d <- twoClassSim(n, linearVars = 15, noiseVars = q)</pre>
 d$Class <- factor(as.numeric(d$Class) - 1)
 idx.0 <- which(d$Class == 0)</pre>
 idx.1 <- sample(which(d$Class == 1), sum(d$Class == 1) / ir , replace = FALSE)</pre>
 d <- d[c(idx.0,idx.1),, drop = FALSE]</pre>
 ## permutation VIMP for BRF with and without blocking
 ## blocked VIMP is a hybrid of Breiman-Cutler/Ishwaran-Kogalur VIMP
 brf <- imbalanced(f, d, method = "brf", importance = "permute", block.size = 1)</pre>
 brfB <- imbalanced(f, d, method = "brf", importance = "permute", block.size = 10)</pre>
 ## permutation VIMP for RFQ with and without blocking
 rfq <- imbalanced(f, d, importance = "permute", block.size = 1)</pre>
 rfqB <- imbalanced(f, d, importance = "permute", block.size = 10)</pre>
 ## compare VIMP values
 imp <- 100 * cbind(brf$importance[, 1], brfB$importance[, 1],</pre>
                   rfq$importance[, 1], rfqB$importance[, 1])
 legn <- c("BRF", "BRF-block", "RFQ", "RFQ-block")</pre>
 colr <- rep(4,20+q)</pre>
 colr[1:20] <- 2
 ylim <- range(c(imp))</pre>
 nms <- 1:(20+q)
 par(mfrow=c(2,2))
 barplot(imp[,1],col=colr,las=2,main=legn[1],ylim=ylim,names.arg=nms)
 barplot(imp[,2],col=colr,las=2,main=legn[2],ylim=ylim,names.arg=nms)
 barplot(imp[,3],col=colr,las=2,main=legn[3],ylim=ylim,names.arg=nms)
 barplot(imp[,4],col=colr,las=2,main=legn[4],ylim=ylim,names.arg=nms)
}
##-----
##
## confidence intervals for G-mean permutation VIMP using subsampling
##
##_____
```

```
if (library("caret", logical.return = TRUE)) {
 ## experimental settings
 n <- 1000
 q <- 20
 ir <- 6
 f <- as.formula(Class ~ .)</pre>
 ## simulate the data, create minority class data
 d <- twoClassSim(n, linearVars = 15, noiseVars = q)</pre>
 d$Class <- factor(as.numeric(d$Class) - 1)
 idx.0 <- which(d$Class == 0)</pre>
 idx.1 <- sample(which(d$Class == 1), sum(d$Class == 1) / ir , replace = FALSE)</pre>
 d <- d[c(idx.0,idx.1),, drop = FALSE]</pre>
 ## RFQ
 o <- imbalanced(Class ~ ., d, importance = "permute", block.size = 10)</pre>
 ## subsample RFQ
 smp.o <- subsample(o, B = 100)
 plot(smp.o, cex.axis = .7)
}
```

impute.rfsrc Impute Only Mode

Description

Fast imputation mode. A random forest is grown and used to impute missing data. No ensemble estimates or error rates are calculated.

Usage

```
## S3 method for class 'rfsrc'
impute(formula, data,
    ntree = 100, nodesize = 1, nsplit = 10,
    nimpute = 2, fast = FALSE, blocks,
    mf.q, max.iter = 10, eps = 0.01,
    ytry = NULL, always.use = NULL, verbose = TRUE,
    ...)
```

Arguments

formula

A symbolic model description. Can be omitted if outcomes are unspecified or if distinction between outcomes and predictors is unnecessary. Ignored for multi-variate missForest.

impute.rfsrc

| data | A data frame containing variables to be imputed. |
|------------|--|
| ntree | Number of trees grown for each imputation. |
| nodesize | Minimum terminal node size in each tree. |
| nsplit | Non-negative integer for specifying random splitting. |
| nimpute | Number of iterations for the missing data algorithm. Ignored for multivariate missForest, which iterates to convergence unless capped by max.iter. |
| fast | If TRUE, uses rfsrcFast instead of rfsrc. Increases speed but may reduce accuracy. |
| blocks | Number of row-wise blocks to divide the data into. May improve speed for large data, but can reduce imputation accuracy. No action if unspecified. |
| mf.q | Enables missForest. Either a fraction (between 0 and 1) of variables treated as responses, or an integer indicating number of response variables. $mf.q = 1$ corresponds to standard missForest. |
| max.iter | Maximum number of iterations for multivariate missForest. |
| eps | Convergence threshold for multivariate missForest (change in imputed values). |
| ytry | Number of variables used as pseudo-responses in unsupervised forests. See Details. |
| always.use | Character vector of variables always included as responses in multivariate miss- Forest. Ignored by other methods. |
| verbose | If TRUE, prints progress during multivariate missForest imputation. |
| | Additional arguments passed to or from methods. |

Details

- 1. Before imputation, observations and variables with all values missing are removed.
- 2. A forest is grown and used solely for imputation. No ensemble statistics (e.g., error rates) are computed. Use this function when imputation is the only goal.
- 3. For standard imputation (not missForest), splits are based only on non-missing data. If a split variable has missing values, they are temporarily imputed by randomly drawing from in-bag, non-missing values to allow node assignment.
- 4. If mf.q is specified, multivariate missForest imputation is applied (Stekhoven and B\"uhlmann, 2012). A fraction (or integer count) of variables are selected as multivariate responses, predicted using the remaining variables with multivariate composite splitting. Each round imputes a disjoint set of variables, and the full cycle is repeated until convergence, controlled by max.iter and eps. Setting mf.q = 1 reverts to standard missForest. This method is typically the most accurate, but also the most computationally intensive.
- 5. If no formula is provided, unsupervised splitting is used. The default ytry is sqrt(p), where p is the number of variables. For each of mtry candidate variables, a random subset of ytry variables is selected as pseudo-responses. A multivariate composite splitting rule is applied, and the split is made on the variable yielding the best result (Tang and Ishwaran, 2017).
- 6. If no missing values remain after preprocessing, the function returns the processed data without further action.
- 7. All standard rfsrc options apply; see examples below for illustration.

Value

Invisibly, the data frame containing the orginal data with imputed data overlaid.

Author(s)

Hemant Ishwaran and Udaya B. Kogalur

References

Ishwaran H., Kogalur U.B., Blackstone E.H. and Lauer M.S. (2008). Random survival forests, *Ann. App. Statist.*, 2:841-860.

Stekhoven D.J. and Buhlmann P. (2012). MissForest-non-parametric missing value imputation for mixed-type data. *Bioinformatics*, 28(1):112-118.

Tang F. and Ishwaran H. (2017). Random forest missing data algorithms. *Statistical Analysis and Data Mining*, 10:363-377.

See Also

rfsrc, rfsrc.fast

Examples

```
## -----
## example of survival imputation
## -----
## default everything - unsupervised splitting
data(pbc, package = "randomForestSRC")
pbc1.d <- impute(data = pbc)</pre>
## imputation using outcome splitting
f <- as.formula(Surv(days, status) ~ .)</pre>
pbc2.d <- impute(f, data = pbc, nsplit = 3)</pre>
## random splitting can be reasonably good
pbc3.d <- impute(f, data = pbc, splitrule = "random", nimpute = 5)</pre>
## -----
## example of regression imputation
## -----
air1.d <- impute(data = airquality, nimpute = 5)</pre>
air2.d <- impute(Ozone ~ ., data = airquality, nimpute = 5)</pre>
air3.d <- impute(Ozone ~ ., data = airquality, fast = TRUE)
## -----
## multivariate missForest imputation
## ______
```

data(pbc, package = "randomForestSRC")

impute.rfsrc

```
## missForest algorithm - uses 1 variable at a time for the response
pbc.d <- impute(data = pbc, mf.q = 1)</pre>
## multivariate missForest - use 10 percent of variables as responses
## i.e. multivariate missForest
pbc.d <- impute(data = pbc, mf.q = .01)</pre>
## missForest but faster by using random splitting
pbc.d <- impute(data = pbc, mf.q = 1, splitrule = "random")</pre>
## missForest but faster by increasing nodesize
pbc.d <- impute(data = pbc, mf.q = 1, nodesize = 20, splitrule = "random")</pre>
## missForest but faster by using rfsrcFast
pbc.d <- impute(data = pbc, mf.q = 1, fast = TRUE)</pre>
## -----
## another example of multivariate missForest imputation
## (suggested by John Sheffield)
test_rows <- 1000</pre>
set.seed(1234)
a <- rpois(test_rows, 500)</pre>
b <- a + rnorm(test_rows, 50, 50)</pre>
c <- b + rnorm(test_rows, 50, 50)</pre>
d <- c + rnorm(test_rows, 50, 50)</pre>
e <- d + rnorm(test_rows, 50, 50)</pre>
f <- e + rnorm(test_rows, 50, 50)</pre>
g <- f + rnorm(test_rows, 50, 50)
h <- g + rnorm(test_rows, 50, 50)
i <- h + rnorm(test_rows, 50, 50)</pre>
fake_data <- data.frame(a, b, c, d, e, f, g, h, i)</pre>
fake_data_missing <- data.frame(lapply(fake_data, function(x) {</pre>
  x[runif(test_rows) <= 0.4] <- NA</pre>
  х
}))
imputed_data <- impute(</pre>
  data = fake_data_missing,
  mf.q = 0.2,
 ntree = 100,
  fast = TRUE,
  verbose = TRUE
)
par(mfrow=c(3,3))
o=lapply(1:ncol(imputed_data), function(j) {
  pt <- is.na(fake_data_missing[, j])</pre>
```

```
x <- fake_data[pt, j]
y <- imputed_data[pt, j]
plot(x, y, pch = 16, cex = 0.8, xlab = "raw data",
    ylab = "imputed data", col = 2)
points(x, y, pch = 1, cex = 0.8, col = gray(.9))
lines(supsmu(x, y, span = .25), lty = 1, col = 4, lwd = 4)
mtext(colnames(imputed_data)[j])
NULL
})</pre>
```

max.subtree.rfsrc Acquire Maximal Subtree Information

Description

Extract maximal subtree information from a RF-SRC object. Used for variable selection and identifying interactions between variables.

Usage

```
## S3 method for class 'rfsrc'
max.subtree(object,
max.order = 2, sub.order = FALSE, conservative = FALSE, ...)
```

Arguments

| object | An object of class (rfsrc, grow) or (rfsrc, forest). |
|--------------|---|
| max.order | Non-negative integer specifying the maximum interaction order for which min- imal depth is calculated. Defaults to 2. Set max.order=0 to return first-order depths only. When max.order=0, conservative is automatically set to FALSE. |
| sub.order | Logical. If TRUE, returns the minimal depth of each variable conditional on every other variable. Useful for investigating variable interdependence. See Details. |
| conservative | Logical. If TRUE, uses a conservative threshold for selecting variables based on the marginal minimal depth distribution (Ishwaran et al., 2010). If FALSE, uses the tree-averaged distribution, which is less conservative and typically identifies more variables in high-dimensional settings. |
| | Additional arguments passed to or from other methods. |

Details

The maximal subtree for a variable x is the largest subtree in which the root node splits on x. The largest possible maximal subtree is the full tree (root node), though multiple maximal subtrees may exist for a variable. A variable may also have no maximal subtree if it is never used for splitting. See Ishwaran et al. (2010, 2011) for further discussion.

max.subtree.rfsrc

The minimal depth of a maximal subtree-called the *first-order depth*-quantifies the predictive strength of a variable. It is defined as the distance from the root node to the parent of the closest maximal subtree for *x*. Smaller values indicate stronger predictive impact. A variable is flagged as strong if its minimal depth is below the mean of the minimal depth distribution.

The *second-order depth* is the distance from the root to the second-closest maximal subtree of x. To request depths beyond first order, use the max.order option (e.g., max.order = 2 returns both first and second-order depths). Set max.order = 0 to retrieve first-order depths for each variable in each tree.

Set sub.order = TRUE to obtain the relative minimal depth of each variable j within the maximal subtree of another variable i. This returns a p x p matrix (with p the number of variables) whose entry (i,j) is the normalized relative depth of j in i's subtree. Entry (i,i) gives the depth of i relative to the root. Read the matrix across rows to assess inter-variable relationships: small (i,j) entries suggest interactions between variables i and j. See find.interaction for further details.

For competing risks, all analyses are unconditional (non-event specific).

Value

Invisibly returns a list with the following components:

| order | Matrix of order depths for each variable up to max.order, averaged over trees. The matrix has p rows and max.order columns, where p is the number of variables. If max.order = 0, returns a matrix of dimension $p \times ntree$ containing first-order depths for each variable by tree. |
|-----------------|---|
| count | Average number of maximal subtrees per variable, normalized by tree size. |
| nodes.at.depth | List of vectors recording the number of non-terminal nodes at each depth level for each tree. |
| sub.order | Matrix of average minimal depths of each variable relative to others (i.e., condi- tional minimal depth matrix). NULL if sub.order = FALSE. |
| threshold | Threshold value for selecting strong variables based on the mean of the minimal depth distribution. |
| threshold.1se | Conservative threshold equal to the mean minimal depth plus one standard error. |
| topvars | Character vector of selected variable names using the threshold criterion. |
| topvars.1se | Character vector of selected variable names using the threshold.1se criterion. |
| percentile | Percentile value of minimal depth for each variable. |
| density | Estimated density of the minimal depth distribution. |
| second.order.th | reshold |
| | Threshold used for selecting strong second-order depth variables. |

Author(s)

Hemant Ishwaran and Udaya B. Kogalur

References

Ishwaran H., Kogalur U.B., Gorodeski E.Z, Minn A.J. and Lauer M.S. (2010). High-dimensional variable selection for survival data. *J. Amer. Statist. Assoc.*, 105:205-217.

Ishwaran H., Kogalur U.B., Chen X. and Minn A.J. (2011). Random survival forests for highdimensional data. *Statist. Anal. Data Mining*, 4:115-132.

See Also

holdout.vimp.rfsrc,vimp.rfsrc

Examples

```
## ------
## survival analysis
## first and second order depths for all variables
## -----
data(veteran, package = "randomForestSRC")
v.obj <- rfsrc(Surv(time, status) ~ . , data = veteran)</pre>
v.max <- max.subtree(v.obj)</pre>
# first and second order depths
print(round(v.max$order, 3))
# the minimal depth is the first order depth
print(round(v.max$order[, 1], 3))
# strong variables have minimal depth less than or equal
# to the following threshold
print(v.max$threshold)
# this corresponds to the set of variables
print(v.max$topvars)
## -----
## regression analysis
## try different levels of conservativeness
mtcars.obj <- rfsrc(mpg ~ ., data = mtcars)</pre>
max.subtree(mtcars.obj)$topvars
```

nutrigenomic

max.subtree(mtcars.obj, conservative = TRUE)\$topvars

nutrigenomic

Description

Investigates the effects of five dietary treatments on 21 liver lipids and 120 hepatic gene expressions in wild-type and PPAR-alpha deficient mice. A multivariate mixed random forest analysis is performed by regressing gene expression, diet, and genotype (x-variables) on lipid expression profiles (multivariate y-responses).

References

Martin P.G. et al. (2007). Novel aspects of PPAR-alpha-mediated regulation of lipid and xenobiotic metabolism revealed through a nutrigenomic study. *Hepatology*, 45(3), 767–777.

Examples

```
## multivariate regression forests using Mahalanobis splitting
## lipids (all real values) used as the multivariate y
## ______
## load the data
data(nutrigenomic, package = "randomForestSRC")
## parse into y and x data
ydta <- nutrigenomic$lipids</pre>
xdta <- data.frame(nutrigenomic$genes,</pre>
               diet = nutrigenomic$diet,
               genotype = nutrigenomic$genotype)
## multivariate mixed forest call
obj <- rfsrc(get.mv.formula(colnames(ydta)),</pre>
          data.frame(ydta, xdta),
          importance=TRUE, nsplit = 10,
          splitrule = "mahalanobis")
print(obj)
## ------
## plot the standarized performance and VIMP values
## ------
## acquire the error rate for each of the 21-coordinates
## standardize to allow for comparison across coordinates
serr <- get.mv.error(obj, standardize = TRUE)</pre>
## acquire standardized VIMP
svimp <- get.mv.vimp(obj, standardize = TRUE)</pre>
par(mfrow = c(1,2))
plot(serr, xlab = "Lipids", ylab = "Standardized Performance")
matplot(svimp, xlab = "Genes/Diet/Genotype", ylab = "Standardized VIMP")
## ______
## plot some trees
```

```
-----
##
plot(get.tree(obj, 1))
plot(get.tree(obj, 2))
plot(get.tree(obj, 3))
## -----
##
## Compare above to (1) user specified covariance matrix
##
                 (2) default composite (independent) splitting
##
         _____
##
## user specified sigma matrix
obj2 <- rfsrc(get.mv.formula(colnames(ydta)),</pre>
            data.frame(ydta, xdta),
            importance = TRUE, nsplit = 10,
            splitrule = "mahalanobis",
            sigma = cov(ydta))
print(obj2)
## default independence split rule
obj3 <- rfsrc(get.mv.formula(colnames(ydta)),</pre>
           data.frame(ydta, xdta),
            importance=TRUE, nsplit = 10)
print(obj3)
## compare vimp
imp <- data.frame(mahalanobis = rowMeans(get.mv.vimp(obj, standardize = TRUE)),</pre>
               mahalanobis2 = rowMeans(get.mv.vimp(obj2, standardize = TRUE)),
               default
                         = rowMeans(get.mv.vimp(obj3, standardize = TRUE)))
print(head(100 * imp[order(imp$mahalanobis, decreasing = TRUE), ], 15))
```

partial.rfsrc Acquire Partial Effect of a Variable

Description

Direct, fast inferface for partial effect of a variable. Works for all families.

Usage

```
partial.rfsrc(object, oob = TRUE,
    partial.type = NULL, partial.xvar = NULL, partial.values = NULL,
    partial.xvar2 = NULL, partial.values2 = NULL,
    partial.time = NULL, get.tree = NULL, seed = NULL, do.trace = FALSE, ...)
```

partial.rfsrc

Arguments

| object | An object of class (rfsrc, grow). |
|-----------------|---|
| oob | By default out-of-bag values are returned, but inbag values can be requested by setting this option to FALSE. |
| partial.type | Character vector specifying type of predicted value requested. See details below. |
| partial.xvar | Character value specifying the single primary partial x-variable to be used. |
| partial.values | Vector of values that the primary partialy x-variable will assume. |
| partial.xvar2 | Vector of character values specifying the second order x-variables to be used. |
| partial.values2 | |
| | Vector of values that the second order x-variables will assume. Each second order x-variable can only assume a single value. This the length of partial.xvar2 and partial.values2 will be the same. In addition, the user must do the appropriate conversion for factors, and represent a value as a numeric element. |
| partial.time | For survival families, the time at which the predicted survival value is evaluated at (depends on partial.type). |
| get.tree | Vector of integer(s) identifying trees over which the partial values are calculated over. By default, uses all trees in the forest. |
| seed | Negative integer specifying seed for the random number generator. |
| do.trace | Number of seconds between updates to the user on approximate time to completion. |
| | Further arguments passed to or from other methods. |

Details

Used for direct, efficient call to obtain partial plot effects. This function is intended primarily for experts.

Out-of-bag (OOB) values are returned by default.

For factors, the partial value should be encoded as a positive integer reflecting the level number of the factor. The actual label of the factor should not be used.

The utility function get.partial.plot.data is supplied for processing returned raw partial effects in a format more convenient for plotting. Options are specified as in plot.variable. See examples for illustration.

Raw partial plot effects data is returned either as an array or a list of length equal to the number of outcomes (length is one for univariate families) with entries depending on the underlying family:

- 1. For regression, partial plot data is returned as a list in regrOutput with dim [n] x [length(partial.values)].
- For classification, partial plot data is returned as a list in classOutput of dim [n] x [1 + yvar.nlevels[.]] x [length(partial.values)].
- 3. For mixed multivariate regression, values are returned in list format both in regrOutput and classOutput
- 4. For survival, values are returned as either a matrix or array in survOutput. Depending on partial type specified this can be:

- For partial type surv returns the survival function of dim [n] x [length(partial.time)] x [length(partial.values)].
- For partial type mort returns mortality of dim [n] x [length(partial.values)].
- For partial type chf returns the cumulative hazard function of dim [n] x [length(partial.time)] x [length(partial.values)].
- 5. For competing risks, values are returned as either a matrix or array in survOutput. Depending on the options specified this can be:
 - For partial type years.lost returns the expected number of life years lost of dim [n] x [length(event.info\$event.type)] x [length(partial.values)].
 - For partial type cif returns the cumulative incidence function of dim [n] x [length(partial.time)] x [length(event.info\$event.type)] x [length(partial.values)].
 - For partial type chf returns the cumulative hazard function of dim [n] x [length(partial.time)] x [length(event.info\$event.type)] x [length(partial.values)].

Author(s)

Hemant Ishwaran and Udaya B. Kogalur

References

Ishwaran H., Kogalur U.B. (2007). Random survival forests for R, Rnews, 7(2):25-31.

Ishwaran H., Kogalur U.B., Blackstone E.H. and Lauer M.S. (2008). Random survival forests, *Ann. App. Statist.*, 2:841-860.

See Also

plot.variable.rfsrc

Examples
```
## example where we display all the partial effects
## instead of averaging - use the granule=TRUE option
pdta <- get.partial.plot.data(partial.obj, granule = TRUE)</pre>
boxplot(pdta$yhat ~ pdta$x, xlab = "Wind", ylab = "partial effect")
##
## regression: partial effects for two variables simultaneously
##
## ------
airq.obj <- rfsrc(Ozone ~ ., data = airquality)</pre>
## specify wind and temperature values of interest
wind <- sort(unique(airq.obj$xvar$Wind))</pre>
temp <- sort(unique(airq.obj$xvar$Temp))</pre>
## partial effect for wind, for a given temp
pdta <- do.call(rbind, lapply(temp, function(x2) {</pre>
 o <- partial(airq.obj,</pre>
        partial.xvar = "Wind", partial.xvar2 = "Temp",
        partial.values = wind, partial.values2 = x2)
 cbind(wind, x2, get.partial.plot.data(o)$yhat)
}))
pdta <- data.frame(pdta)</pre>
colnames(pdta) <- c("wind", "temp", "effectSize")</pre>
## coplot of partial effect of wind and temp
coplot(effectSize ~ wind|temp, pdta, pch = 16, overlap = 0)
## -----
##
## regression: partial effects for three variables simultaneously
## (can be slow, so modify accordingly)
##
## -----
n <- 1000
x \leftarrow matrix(rnorm(n \times 3), ncol = 3)
y \le x[, 1] + x[, 1] * x[, 2] + x[, 1] * x[, 2] * x[, 3]
o <- rfsrc(y ~ ., data = data.frame(y = y, x))</pre>
## define target x values
x1 <- seq(-3, 3, length = 40)
x2 <- x3 <- seq(-3, 3, length = 10)
## extract second order partial effects
pdta <- do.call(rbind,</pre>
         lapply(x3, function(x3v) {
           cat("outer loop x3 = ", x3v, "\n")
           do.call(rbind,lapply(x2, function(x2v) {
            o <- partial(o,</pre>
```

```
partial.xvar = "X1",
                   partial.values = x1,
                   partial.xvar2 = c("X2", "X3"),
                   partial.values2 = c(x2v, x3v))
            cbind(x1, x2v, x3v, get.partial.plot.data(o)$yhat)
          }))
         }))
pdta <- data.frame(pdta)</pre>
colnames(pdta) <- c("x1", "x2", "x3", "effectSize")</pre>
## coplot of partial effects
coplot(effectSize ~ x1|x2*x3, pdta, pch = 16, overlap = 0)
## -----
##
## classification
##
## ------
iris.obj <- rfsrc(Species ~., data = iris)</pre>
## partial effect for sepal length
partial.obj <- partial(iris.obj,</pre>
                partial.xvar = "Sepal.Length",
                partial.values = iris.obj$xvar$Sepal.Length)
## extract partial effects for each species outcome
pdta1 <- get.partial.plot.data(partial.obj, target = "setosa")</pre>
pdta2 <- get.partial.plot.data(partial.obj, target = "versicolor")</pre>
pdta3 <- get.partial.plot.data(partial.obj, target = "virginica")</pre>
## plot the results
par(mfrow=c(1,1))
plot(pdta1$x, pdta1$yhat, type="b", pch = 16,
    xlab = "sepal length", ylab = "adjusted probability",
    ylim = range(pdta1$yhat,pdta2$yhat,pdta3$yhat))
points(pdta2$x, pdta2$yhat, col = 2, type = "b", pch = 16)
points(pdta3$x, pdta3$yhat, col = 4, type = "b", pch = 16)
legend("topleft", legend=levels(iris.obj$yvar), fill = c(1, 2, 4))
## ---
       _____
##
## survival
##
data(veteran, package = "randomForestSRC")
v.obj <- rfsrc(Surv(time,status)~., veteran, nsplit = 10, ntree = 100)</pre>
## partial effect of age on mortality
partial.obj <- partial(v.obj,</pre>
 partial.type = "mort",
```

partial.rfsrc

```
partial.xvar = "age",
 partial.values = v.obj$xvar$age,
 partial.time = v.obj$time.interest)
pdta <- get.partial.plot.data(partial.obj)</pre>
plot(lowess(pdta$x, pdta$yhat, f = 1/3),
  type = "l", xlab = "age", ylab = "adjusted mortality")
## example where x is discrete - partial effect of age on mortality
## we use the granule=TRUE option
partial.obj <- partial(v.obj,</pre>
      partial.type = "mort"
      partial.xvar = "trt",
      partial.values = v.obj$xvar$trt,
      partial.time = v.obj$time.interest)
pdta <- get.partial.plot.data(partial.obj, granule = TRUE)</pre>
boxplot(pdta$yhat ~ pdta$x, xlab = "treatment", ylab = "partial effect")
## partial effects of karnofsky score on survival
karno <- quantile(v.obj$xvar$karno)</pre>
partial.obj <- partial(v.obj,</pre>
 partial.type = "surv",
 partial.xvar = "karno",
 partial.values = karno,
 partial.time = v.obj$time.interest)
pdta <- get.partial.plot.data(partial.obj)</pre>
matplot(pdta$partial.time, t(pdta$yhat), type = "l", lty = 1,
    xlab = "time", ylab = "karnofsky adjusted survival")
legend("topright", legend = paste0("karnofsky = ", karno), fill = 1:5)
## -----
##
## competing risk
##
## ------
data(follic, package = "randomForestSRC")
follic.obj <- rfsrc(Surv(time, status) ~ ., follic, nsplit = 3, ntree = 100)</pre>
## partial effect of age on years lost
partial.obj <- partial(follic.obj,</pre>
 partial.type = "years.lost",
 partial.xvar = "age",
 partial.values = follic.obj$xvar$age,
 partial.time = follic.obj$time.interest)
pdta1 <- get.partial.plot.data(partial.obj, target = 1)</pre>
pdta2 <- get.partial.plot.data(partial.obj, target = 2)</pre>
par(mfrow=c(2,2))
plot(lowess(pdta1$x, pdta1$yhat),
```

```
type = "l", xlab = "age", ylab = "adjusted years lost relapse")
plot(lowess(pdta2$x, pdta2$yhat),
   type = "l", xlab = "age", ylab = "adjusted years lost death")
## partial effect of age on cif
partial.obj <- partial(follic.obj,</pre>
 partial.type = "cif",
 partial.xvar = "age",
 partial.values = quantile(follic.obj$xvar$age),
 partial.time = follic.obj$time.interest)
pdta1 <- get.partial.plot.data(partial.obj, target = 1)</pre>
pdta2 <- get.partial.plot.data(partial.obj, target = 2)</pre>
matplot(pdta1$partial.time, t(pdta1$yhat), type = "1", lty = 1,
     xlab = "time", ylab = "age adjusted cif for relapse")
matplot(pdta2$partial.time, t(pdta2$yhat), type = "1", lty = 1,
     xlab = "time", ylab = "age adjusted cif for death")
## ------
##
## multivariate mixed outcomes
##
## ______
mtcars2 <- mtcars</pre>
mtcars2$carb <- factor(mtcars2$carb)</pre>
mtcars2$cyl <- factor(mtcars2$cyl)</pre>
mtcars.mix <- rfsrc(Multivar(carb, mpg, cyl) ~ ., data = mtcars2)</pre>
## partial effect of displacement for each the three-outcomes
partial.obj <- partial(mtcars.mix,</pre>
                 partial.xvar = "disp",
                 partial.values = mtcars.mix$xvar$disp)
pdta1 <- get.partial.plot.data(partial.obj, m.target = "carb")</pre>
pdta2 <- get.partial.plot.data(partial.obj, m.target = "mpg")</pre>
pdta3 <- get.partial.plot.data(partial.obj, m.target = "cyl")</pre>
par(mfrow=c(2,2))
plot(lowess(pdta1$x, pdta1$yhat), type = "1", xlab="displacement", ylab="carb")
plot(lowess(pdta2$x, pdta2$yhat), type = "l", xlab="displacement", ylab="mpg")
plot(lowess(pdta3$x, pdta3$yhat), type = "l", xlab="displacement", ylab="cyl")
```

Primary Biliary Cirrhosis (PBC) Data

40

pbc

peakVO2

Description

Data from the Mayo Clinic trial in primary biliary cirrhosis (PBC) of the liver conducted between 1974 and 1984. A total of 424 PBC patients, referred to Mayo Clinic during that ten-year interval, met eligibility criteria for the randomized placebo controlled trial of the drug D-penicillamine. The first 312 cases in the data set participated in the randomized trial and contain largely complete data.

Source

Flemming and Harrington, 1991, Appendix D.1.

References

Flemming T.R and Harrington D.P., (1991) *Counting Processes and Survival Analysis*. New York: Wiley.

Examples

data(pbc, package = "randomForestSRC")
pbc.obj <- rfsrc(Surv(days, status) ~ ., pbc, nsplit = 3)</pre>

peakV02

Systolic Heart Failure Data

Description

The data involve 2231 patients with systolic heart failure who underwent cardiopulmonary stress testing at the Cleveland Clinic. The primary end point was all-cause death. In total, 39 variables were measured for each patient, including baseline clinical values and exercise stress test results. A key variable of interest is peak VO2 (mL/kg per min), the peak respiratory exchange ratio. More details regarding the data can be found in Hsich et al. (2011).

References

Hsich E., Gorodeski E.Z., Blackstone E.H., Ishwaran H. and Lauer M.S. (2011). Identifying important risk factors for survival in systolic heart failure patients using random survival forests. Circulation: Cardio. Qual. Outcomes, 4(1), 39-45.

Examples

```
## load the data
data(peakV02, package = "randomForestSRC")
## random survival forest analysis
o <- rfsrc(Surv(ttodead, died)~., peakV02)
print(o)
## partial effect of peak V02 on mortality</pre>
```

```
partial.o <- partial(o,</pre>
       partial.type = "mort",
       partial.xvar = "peak.vo2",
       partial.values = o$xvar$peak.vo2,
       partial.time = o$time.interest)
pdta.m <- get.partial.plot.data(partial.o)</pre>
## partial effect of peak V02 on survival
pvo2 <- quantile(o$xvar$peak.vo2)</pre>
partial.o <- partial(o,</pre>
       partial.type = "surv",
       partial.xvar = "peak.vo2",
       partial.values = pvo2,
       partial.time = o$time.interest)
pdta.s <- get.partial.plot.data(partial.o)</pre>
## compare the two plots
par(mfrow=c(1,2))
plot(lowess(pdta.m$x, pdta.m$yhat, f = 2/3),
     type = "1", xlab = "peak VO2", ylab = "adjusted mortality")
rug(o$xvar$peak.vo2)
matplot(pdta.s$partial.time, t(pdta.s$yhat), type = "l", lty = 1,
          xlab = "years", ylab = "peak VO2 adjusted survival")
legend("bottomleft", legend = paste0("peak VO2 = ", pvo2),
       bty = "n", cex = .75, fill = 1:5)
```

plot.competing.risk.rfsrc

Plots for Competing Risks

Description

Plot useful summary curves from a random survival forest competing risk analysis.

Usage

S3 method for class 'rfsrc'
plot.competing.risk(x, plots.one.page = FALSE, ...)

Arguments

| х | An object of class (rfsrc, grow) or (rfsrc, predict). |
|---------------------------|---|
| <pre>plots.one.page</pre> | Should plots be placed on one page? |
| | Further arguments passed to or from other methods. |

Details

Given a random survival forest object from a competing risk analysis (Ishwaran et al. 2014), plots from top to bottom, left to right: (1) cause-specific cumulative hazard function (CSCHF) for each event, (2) cumulative incidence function (CIF) for each event, and (3) continuous probability curves (CPC) for each event (Pepe and Mori, 1993).

Does not apply to right-censored data. Whenever possible, out-of-bag (OOB) values are displayed.

Author(s)

Hemant Ishwaran and Udaya B. Kogalur

References

Ishwaran H., Gerds T.A., Kogalur U.B., Moore R.D., Gange S.J. and Lau B.M. (2014). Random survival forests for competing risks. *Biostatistics*, 15(4):757-773.

Pepe, M.S. and Mori, M., (1993). Kaplan-Meier, marginal or conditional probability curves in summarizing competing risks failure time data? *Statistics in Medicine*, 12(8):737-751.

See Also

follic, hd, rfsrc, wihs

Examples

```
## follicular cell lymphoma
## -----
 data(follic, package = "randomForestSRC")
 follic.obj <- rfsrc(Surv(time, status) ~ ., follic, nsplit = 3, ntree = 100)</pre>
 print(follic.obj)
 plot.competing.risk(follic.obj)
## -------
## Hodgkin's Disease
## ------
 data(hd, package = "randomForestSRC")
 hd.obj <- rfsrc(Surv(time, status) ~ ., hd, nsplit = 3, ntree = 100)</pre>
 print(hd.obj)
 plot.competing.risk(hd.obj)
## -----
## competing risk analysis of pbc data from the survival package
## events are transplant (1) and death (2)
## -----
if (library("survival", logical.return = TRUE)) {
 data(pbc, package = "survival")
 pbc$id <- NULL</pre>
```

```
plot.competing.risk(rfsrc(Surv(time, status) ~ ., pbc))
}
```

plot.quantreg.rfsrc Plot Quantiles from Quantile Regression Forests

Description

Plots quantiles obtained from a quantile regression forest. Additionally insets the continuous rank probability score (crps), a useful diagnostic of accuracy.

Usage

```
## S3 method for class 'rfsrc'
plot.quantreg(x, prbL = .25, prbU = .75,
    m.target = NULL, crps = TRUE, subset = NULL, xlab = NULL, ylab = NULL, ...)
```

Arguments

| A quantile regression object returned by a call to quantreg. |
|--|
| Lower quantile level, typically less than 0.5. |
| Upper quantile level, typically greater than 0.5. |
| Character string specifying the target outcome for multivariate families. If not provided, a default target is selected automatically. |
| Logical. If TRUE, calculates the continuous ranked probability score (CRPS) and adds it to the plot. |
| Optional vector specifying a subset of the data to be plotted. Defaults to plotting all data points. |
| Label for the x-axis. |
| Label for the y-axis. |
| Additional arguments passed to or from other methods. |
| |

Author(s)

Hemant Ishwaran and Udaya B. Kogalur

See Also

quantreg.rfsrc

plot.rfsrc

Description

Plot out-of-bag (OOB) error rates and variable importance (VIMP) from a RF-SRC analysis. This is the default plot method for the package.

Usage

```
## S3 method for class 'rfsrc'
plot(x, m.target = NULL,
    plots.one.page = TRUE, sorted = TRUE, verbose = TRUE, ...)
```

Arguments

| х | An object of class (rfsrc, grow), (rfsrc, synthetic), or (rfsrc, predict) |
|----------------|--|
| m.target | Character value for multivariate families specifying the target outcome to be used. If left unspecified, the algorithm will choose a default target. |
| plots.one.page | Should plots be placed on one page? |
| sorted | Should variables be sorted by importance values? |
| verbose | Should VIMP be printed? |
| | Further arguments passed to or from other methods. |

Details

Plot cumulative OOB error rates as a function of number of trees and variable importance (VIMP) if available. Note that the default settings are now such that the error rate is no longer calculated on every tree and VIMP is only calculated if requested. To get OOB error rates for ever tree, use the option block.size = 1 when growing or restoring the forest. Likewise, to view VIMP, use the option importance when growing or restoring the forest.

Author(s)

Hemant Ishwaran and Udaya B. Kogalur

References

Breiman L. (2001). Random forests, *Machine Learning*, 45:5-32.

Ishwaran H. and Kogalur U.B. (2007). Random survival forests for R, Rnews, 7(2):25-31.

Examples

```
## ------
## classification example
## ------
iris.obj <- rfsrc(Species ~ ., data = iris,</pre>
   block.size = 1, importance = TRUE)
plot(iris.obj)
## competing risk example
## -----
## use the pbc data from the survival package
## events are transplant (1) and death (2)
if (library("survival", logical.return = TRUE)) {
 data(pbc, package = "survival")
 pbc$id <- NULL</pre>
 plot(rfsrc(Surv(time, status) ~ ., pbc, block.size = 1))
}
## -----
## multivariate mixed forests
## ______
mtcars.new <- mtcars</pre>
mtcars.new$cyl <- factor(mtcars.new$cyl)</pre>
mtcars.new$carb <- factor(mtcars.new$carb, ordered = TRUE)</pre>
mv.obj <- rfsrc(cbind(carb, mpg, cyl) ~., data = mtcars.new, block.size = 1)</pre>
plot(mv.obj, m.target = "carb")
plot(mv.obj, m.target = "mpg")
plot(mv.obj, m.target = "cyl")
```

plot.subsample.rfsrc Plot Subsampled VIMP Confidence Intervals

Description

Plots VIMP (variable importance) confidence regions obtained from subsampling a forest.

Usage

```
## S3 method for class 'rfsrc'
plot.subsample(x, alpha = .01, xvar.names,
   standardize = TRUE, normal = TRUE, jknife = FALSE, target, m.target = NULL,
   pmax = 75, main = "", sorted = TRUE, show.plots = TRUE, ...)
```

Arguments

| x | An object obtained from calling subample. |
|-------------|--|
| alpha | Desired level of significance. |
| xvar.names | Names of the x-variables to be used. If not specified all variables used. |
| standardize | Standardize VIMP? For regression families, VIMP is standardized by dividing by the variance. For all other families, VIMP is unaltered. |
| normal | Use parametric normal confidence regions or nonparametric regions? Generally, parametric regions perform better. |
| jknife | Use the delete-d jackknife variance estimator? |
| target | For classification families, an integer or character value specifying the class VIMP will be conditioned on (default is to use unconditional VIMP). For competing risk families, an integer value between 1 and J indicating the event VIMP is requested, where J is the number of event types. The default is to use the first event. |
| m.target | Character value for multivariate families specifying the target outcome to be used. If left unspecified, the algorithm will choose a default target. |
| pmax | Trims the data to this number of variables (sorted by VIMP). |
| main | Title used for plot. |
| sorted | Should variables be sorted by importance values? |
| show.plots | Should plots be displayed? Allows users to produce their own custom plots. |
| | Further arguments that can be passed to bxp. |

Details

Most of the options used by the R function bxp will work here and can be used for customization of plots. Currently the following parameters will work:

"xaxt", "yaxt", "las", "cex.axis", "col.axis", "cex.main", "col.main", "sub", "cex.sub", "col.sub", "ylab", "cex.lab", "col.lab"

Value

Invisibly, returns the boxplot data that is plotted.

Author(s)

Hemant Ishwaran and Udaya B. Kogalur

References

Ishwaran H. and Lu M. (2017). Standard errors and confidence intervals for variable importance in random forest regression, classification, and survival.

Politis, D.N. and Romano, J.P. (1994). Large sample confidence regions based on subsamples under minimal assumptions. *The Annals of Statistics*, 22(4):2031-2050.

Shao, J. and Wu, C.J. (1989). A general theory for jackknife variance estimation. *The Annals of Statistics*, 17(3):1176-1197.

See Also

subsample.rfsrc

Examples

```
o <- rfsrc(Ozone ~ ., airquality)
oo <- subsample(o)
plot.subsample(oo)
plot.subsample(oo, xvar.names = o$xvar.names[1:3])
plot.subsample(oo, jknife = FALSE)
plot.subsample(oo, alpha = .01)
plot(oo,cex.axis=.5)
```

plot.survival.rfsrc Plot of Survival Estimates

Description

Plot various survival estimates.

Usage

```
## S3 method for class 'rfsrc'
plot.survival(x, show.plots = TRUE, subset,
    collapse = FALSE, cens.model = c("km", "rfsrc"), ...)
```

Arguments

| х | An object of class (rfsrc, grow) or (rfsrc, predict). |
|------------|--|
| show.plots | Should plots be displayed? |
| subset | Vector indicating which cases from x we want estimates for. All cases used if not specified. |
| collapse | Collapse the survival function? |
| cens.model | Using the training data, specifies method for estimating the censoring distribu- tion used in the inverse probability of censoring weights (IPCW) for calculating the Brier score: |
| | km: Uses the Kaplan-Meier estimator. |
| | rfscr: Uses a censoring random survival forest estimator. |
| | Further arguments passed to or from other methods. |

Details

Produces the following plots (going from top to bottom, left to right):

- 1. Forest estimated survival function for each individual (thick red line is overall ensemble survival, thick green line is Nelson-Aalen estimator).
- 2. Brier score (0=perfect, 1=poor, and 0.25=guessing) stratified by ensemble mortality. Based on the IPCW method described in Gerds et al. (2006). Stratification is into 4 groups corresponding to the 0-25, 25-50, 50-75 and 75-100 percentile values of mortality. Red line is overall (non-stratified) Brier score.
- 3. Continuous rank probability score (CRPS) equal to the integrated Brier score divided by time.
- Plot of mortality of each individual versus observed time. Points in blue correspond to events, black points are censored observations. Not given for prediction settings lacking survival response information.

Whenever possible, out-of-bag (OOB) values are used.

Only applies to survival families. In particular, fails for competing risk analyses. Use plot.competing.risk in such cases.

Mortality (Ishwaran et al., 2008) represents estimated risk for an individual calibrated to the scale of number of events (as a specific example, if *i* has a mortality value of 100, then if all individuals had the same x-values as *i*, we would expect an average of 100 events).

The utility function get.brier.survival can be used to extract the Brier score among other useful quantities.

Value

Invisibly, the conditional and unconditional Brier scores, and the integrated Brier score.

Author(s)

Hemant Ishwaran and Udaya B. Kogalur

References

Gerds T.A and Schumacher M. (2006). Consistent estimation of the expected Brier score in general survival models with right-censored event times, *Biometrical J.*, 6:1029-1040.

Graf E., Schmoor C., Sauerbrei W. and Schumacher M. (1999). Assessment and comparison of prognostic classification schemes for survival data, *Statist. in Medicine*, 18:2529-2545.

Ishwaran H. and Kogalur U.B. (2007). Random survival forests for R, Rnews, 7(2):25-31.

Ishwaran H., Kogalur U.B., Blackstone E.H. and Lauer M.S. (2008). Random survival forests, *Ann. App. Statist.*, 2:841-860.

See Also

plot.competing.risk.rfsrc, predict.rfsrc, rfsrc

Examples

```
## veteran data
data(veteran, package = "randomForestSRC")
plot.survival(rfsrc(Surv(time, status)~ ., veteran), cens.model = "rfsrc")
## pbc data
data(pbc, package = "randomForestSRC")
pbc.obj <- rfsrc(Surv(days, status) ~ ., pbc)
## use subset to focus on specific individuals
plot.survival(pbc.obj, subset = 3)
plot.survival(pbc.obj, subset = c(3, 10))
plot.survival(pbc.obj, subset = c(3, 10), collapse = TRUE)
## get.brier.survival function does many nice things!
plot(get.brier.survival(pbc.obj, cens.model="km")$brier.score,type="s", col=2)
lines(get.brier.survival(pbc.obj, cens.model="rfsrc")$brier.score, type="s", col=4)
legend("bottomright", legend=c("cens.model = km", "cens.model = rfsrc"), fill=c(2,4))
```

plot.variable.rfsrc *Plot Marginal Effect of Variables*

Description

Plot the marginal effect of an x-variable on the class probability (classification), response (regression), mortality (survival), or the expected years lost (competing risk). Users can select between marginal (unadjusted, but fast) and partial plots (adjusted, but slower).

Usage

```
## S3 method for class 'rfsrc'
plot.variable(x, xvar.names, target,
    m.target = NULL, time, surv.type = c("mort", "rel.freq",
    "surv", "years.lost", "cif", "chf"), class.type =
    c("prob", "bayes"), partial = FALSE, oob = TRUE,
    show.plots = TRUE, plots.per.page = 4, granule = 5, sorted = TRUE,
    nvar, npts = 25, smooth.lines = FALSE, subset, ...)
```

Arguments

| х | An object of class (rfsrc, grow), (rfsrc, synthetic), or (rfsrc, plot.variable). |
|------------|---|
| xvar.names | Character vector of x-variable names to include. If not specified, all variables are used. |
| target | For classification, an integer or character specifying the class of interest (default is the first class). For competing risks, an integer between 1 and J indicating the event of interest, where J is the number of event types. Default is the first event type. |

| m.target | Character value for multivariate families specifying the target outcome. If un- specified, a default is automatically chosen. |
|-------------------------|--|
| time | (Survival only) Time point at which the predicted survival value is evaluated, depending on surv.type. |
| surv.type | (Survival only) Type of predicted survival value to compute. See plot.variable details. |
| class.type | (Classification only) Type of predicted classification value to use. See plot.variable details. |
| partial | Logical. If TRUE, partial dependence plots are generated. |
| oob | Logical. If TRUE, out-of-bag predictions are used; otherwise, in-bag predictions are used. |
| show.plots | Logical. If TRUE, plots are displayed on the screen. |
| plots.per.page | Integer controlling the number of plots displayed per page. |
| granule | Integer controlling the coercion of continuous variables to factors (used to generate boxplots). Larger values increase coercion. |
| sorted | Logical. If TRUE, variables are sorted by variable importance. |
| nvar | Number of variables to plot. Defaults to all available variables. |
| npts | Maximum number of points used when generating partial plots for continuous variables. |
| <pre>smooth.lines</pre> | Logical. If TRUE, applies lowess smoothing to partial plots. |
| subset | Vector indicating which rows of x\$xvar to use. Defaults to all rows. Important: do not define subset based on the original dataset (which may have been altered |
| | due to missing data or other processing); define it relative to x\$xvar. |

Details

The vertical axis displays the ensemble-predicted value, while x-variables are plotted along the horizontal axis.

- 1. For regression, the predicted response is plotted.
- 2. For classification, the plotted value is the predicted class probability for the class specified by target, or the most probable class (Bayes rule) depending on whether class.type is set to "prob" or "bayes".
- 3. For multivariate families, the prediction corresponds to the outcome specified by m.target. If this is a classification outcome, target may also be used to indicate the class of interest.
- 4. For survival, the vertical axis shows the predicted value determined by surv.type, with the following options:
 - mort: Mortality (Ishwaran et al., 2008), interpreted as the expected number of events for an individual with the same covariates.
 - rel.freq: Relative frequency of mortality.
 - surv: Predicted survival probability at a specified time point (default is the median follow-up time), controlled via time.

- 5. For competing risks, the vertical axis shows one of the following quantities, depending on surv.type:
 - years.lost: Expected number of life-years lost.
 - cif: Cumulative incidence function for the specified event.
 - chf: Cause-specific cumulative hazard function.

In all competing risks settings, the event of interest is specified using target, and cif and chf are evaluated at the time point given by time.

To generate partial dependence plots, set partial = TRUE. These differ from marginal plots in that they isolate the effect of a single variable X on the predicted value by averaging over all other covariates:

$$\tilde{f}(x) = \frac{1}{n} \sum_{i=1}^{n} \hat{f}(x, x_{i,o}),$$

where $x_{i,o}$ denotes the observed values of all covariates other than X for individual *i*, and \hat{f} is the prediction function. Generating partial plots can be computationally expensive; use a smaller value for npts to reduce the number of grid points evaluated for x.

Plot display conventions:

- For continuous variables: red points indicate partial values; dashed red lines represent an error band of two standard errors. Black dashed lines show the raw partial values. Use smooth.lines = TRUE to overlay a lowess smoothed line.
- For discrete (factor) variables: boxplots are used, with whiskers extending approximately two standard errors from the mean.
- Standard errors are provided only as rough indicators and should be interpreted cautiously.

Partial plots can be slow to compute. Setting npts to a small value can improve performance.

For additional flexibility and speed, consider using partial.rfsrc, which directly computes partial plot data and allows for greater customization.

Author(s)

Hemant Ishwaran and Udaya B. Kogalur

References

Friedman J.H. (2001). Greedy function approximation: a gradient boosting machine, *Ann. of Statist.*, 5:1189-1232.

Ishwaran H., Kogalur U.B. (2007). Random survival forests for R, Rnews, 7(2):25-31.

Ishwaran H., Kogalur U.B., Blackstone E.H. and Lauer M.S. (2008). Random survival forests, *Ann. App. Statist.*, 2:841-860.

Ishwaran H., Gerds T.A., Kogalur U.B., Moore R.D., Gange S.J. and Lau B.M. (2014). Random survival forests for competing risks. *Biostatistics*, 15(4):757-773.

plot.variable.rfsrc

See Also

rfsrc, synthetic.rfsrc, partial.rfsrc, predict.rfsrc

Examples

```
## ------
## survival/competing risk
## survival
data(veteran, package = "randomForestSRC")
v.obj <- rfsrc(Surv(time,status)~., veteran, ntree = 100)</pre>
plot.variable(v.obj, plots.per.page = 3)
plot.variable(v.obj, plots.per.page = 2, xvar.names = c("trt", "karno", "age"))
plot.variable(v.obj, surv.type = "surv", nvar = 1, time = 200)
plot.variable(v.obj, surv.type = "surv", partial = TRUE, smooth.lines = TRUE)
plot.variable(v.obj, surv.type = "rel.freq", partial = TRUE, nvar = 2)
## example of plot.variable calling a pre-processed plot.variable object
p.v <- plot.variable(v.obj, surv.type = "surv", partial = TRUE, smooth.lines = TRUE)</pre>
plot.variable(p.v)
p.v$plots.per.page <- 1</pre>
p.v$smooth.lines <- FALSE</pre>
plot.variable(p.v)
## example using a pre-processed plot.variable to define custom plots
p.v <- plot.variable(v.obj, surv.type = "surv", partial = TRUE, show.plots = FALSE)
plotthis <- p.v$plotthis</pre>
plot(plotthis[["age"]], xlab = "age", ylab = "partial effect", type = "b")
boxplot(yhat ~ x, plotthis[["trt"]], xlab = "treatment", ylab = "partial effect")
## competing risks
data(follic, package = "randomForestSRC")
follic.obj <- rfsrc(Surv(time, status) ~ ., follic, nsplit = 3, ntree = 100)</pre>
plot.variable(follic.obj, target = 2)
## regression
## airquality
airq.obj <- rfsrc(Ozone ~ ., data = airquality)</pre>
plot.variable(airq.obj, partial = TRUE, smooth.lines = TRUE)
plot.variable(airq.obj, partial = TRUE, subset = airq.obj$xvar$Solar.R < 200)</pre>
## motor trend cars
mtcars.obj <- rfsrc(mpg ~ ., data = mtcars)</pre>
plot.variable(mtcars.obj, partial = TRUE, smooth.lines = TRUE)
## ______
## classification
```

```
## -----
## iris
iris.obj <- rfsrc(Species ~., data = iris)</pre>
plot.variable(iris.obj, partial = TRUE)
## motor trend cars: predict number of carburetors
mtcars2 <- mtcars</pre>
mtcars2$carb <- factor(mtcars2$carb,</pre>
  labels = paste("carb", sort(unique(mtcars$carb))))
mtcars2.obj <- rfsrc(carb ~ ., data = mtcars2)</pre>
plot.variable(mtcars2.obj, partial = TRUE)
## -----
## multivariate regression
## ------
mtcars.mreg <- rfsrc(Multivar(mpg, cyl) ~., data = mtcars)</pre>
plot.variable(mtcars.mreg, m.target = "mpg", partial = TRUE, nvar = 1)
plot.variable(mtcars.mreg, m.target = "cyl", partial = TRUE, nvar = 1)
## ------
## multivariate mixed outcomes
## ------
mtcars2 <- mtcars</pre>
mtcars2$carb <- factor(mtcars2$carb)</pre>
mtcars2$cyl <- factor(mtcars2$cyl)</pre>
mtcars.mix <- rfsrc(Multivar(carb, mpg, cyl) ~ ., data = mtcars2)</pre>
plot.variable(mtcars.mix, m.target = "cyl", target = "4", partial = TRUE, nvar = 1)
plot.variable(mtcars.mix, m.target = "cyl", target = 2, partial = TRUE, nvar = 1)
```

predict.rfsrc

Prediction for Random Forests for Survival, Regression, and Classification

Description

Obtain predicted values using a forest. Also returns performance values if the test data contains y-outcomes.

Usage

```
## S3 method for class 'rfsrc'
predict(object,
    newdata,
    importance = c(FALSE, TRUE, "none", "anti", "permute", "random"),
    get.tree = NULL,
```

predict.rfsrc

Arguments

| object | An object of class (rfsrc, grow) or (rfsrc, forest). |
|------------|--|
| newdata | Test data. If omitted, the original training data is used. |
| importance | Method for computing variable importance (VIMP). See vimp for additional options including joint importance. See holdout.vimp for an alternative importance measure. |
| get.tree | Vector of integers specifying which trees to use for ensemble calculations. De- faults to all trees. Useful for extracting ensembles, VIMP, or proximity from specific trees. If specified, block.size is overridden to match the number of trees. See examples for per-tree VIMP extraction. |
| block.size | Controls the granularity of error rate and VIMP calculation. If NULL, error is reported only for the final tree. Set to an integer k to compute error every k trees. For VIMP, calculations are done in blocks of size block.size, balancing between tree-level and forest-level assessments. |
| na.action | Action to take when missing values are present. Options are "na.omit" (de-fault), "na.random" for fast random imputation, or "na.impute" to use the imputation method in rfsrc. |
| outcome | Specifies whether predicted values should be based on the outcomes from the training data ("train", default) or test data. Ignored if newdata is missing or if test outcomes are unavailable. |
| perf.type | Optional metric for prediction, VIMP, and error. Currently used for classifica- tion and multivariate classification. Choices: "misclass" (default), "brier", and "gmean". |
| proximity | Whether to compute the proximity matrix for test observations. Options include "inbag", "oob", "all", TRUE, or FALSE. Not all options are valid in all contexts; TRUE is the safest choice. |
| distance | Whether to compute the distance matrix. Options are the same as for proximity. |
| forest.wt | Whether to compute the forest weight matrix. Options are the same as for proximity. |

| ptn.count | If nonzero, each tree is pruned to have this many terminal nodes. Only the terminal node membership is returned; no prediction is made. Default is $ptn.count = 0$. |
|---------------|--|
| var.used | If TRUE, records how many times each variable was used for splitting. |
| split.depth | If TRUE, returns minimal depth of each variable per case. |
| case.depth | If TRUE, returns a matrix of the depth at which each case first splits in each tree. |
| seed | Negative integer used to set the random seed. |
| do.trace | Number of seconds between progress updates during execution. |
| membership | If TRUE, returns terminal node membership and in-bag information. |
| statistics | If TRUE, returns split statistics, which can be parsed using stat.split. |
| marginal.xvar | Vector of variable names to marginalize over when calculating weights or prox- imity. If a variable is marginalized, its split does not partition the data; all cases are passed to both daughters. When all splits involve marginalized variables, ter- minal nodes contain the full dataset. When no marginalized variables are used, membership is unchanged. Default is NULL (no marginalization). |
| | Additional arguments passed to or from other methods. |

Details

Predicted values are obtained by "dropping" the test data down the trained forest-i.e., the forest grown using the training data. If the test data includes y-outcome values, performance metrics are also returned. Variable importance (VIMP), including joint VIMP, is returned if requested.

If no test data is supplied, the function uses the original training data and enters "restore" mode. This allows users to extract outputs from the trained forest that were not requested during the original grow call.

If outcome = "test", predictions are computed using y-outcomes from the test data (which must include outcome values). Terminal node statistics are recalculated using these outcomes, while the tree topology remains fixed from training. Error rates and VIMP are then computed by bootstrapping the test set and applying out-of-bagging to maintain unbiased estimates.

Set csv = TRUE to return case-specific VIMP, and cse = TRUE to return case-specific error rates. These apply to all families except survival. Both options can also be used at training time.

Value

An object of class (rfsrc, predict), which is a list with the following components:

call The original grow call to rfsrc.

family The family used in the analysis.

n Sample size of the test data (after handling missing values).

ntree Number of trees in the trained forest.

yvar Y-outcome values from the test data or original grow data (if newdata is missing).

yvar.names Character vector of response variable names.

xvar Data frame of test set predictor variables.

predict.rfsrc

- xvar.names Character vector of predictor variable names.
- leaf.count Vector of length ntree giving the number of terminal nodes per tree.
- proximity Proximity matrix computed on the test data.
- forest The trained forest object.
- forest.wt Forest weight matrix for test cases.
- ptn.membership Matrix of pruned terminal node membership. Only returned if ptn.count > 0.
- **membership** Matrix of terminal node membership for test cases. Each column corresponds to one tree.
- **inbag** Matrix indicating how many times each case appears in the bootstrap sample for each tree. **var.used** Number of times each variable was used in splitting.
- **imputed.indv** Indices of test observations with missing values.
- **imputed.data** Imputed version of the test data. Columns are ordered with responses first, followed by predictors.
- **split.depth** Matrix or array recording minimal depth of each variable for each case, optionally by tree.

node.stats Split statistics (only if statistics = TRUE); see stat.split.

err.rate Cumulative out-of-bag (OOB) error rate, if y-outcomes are present.

importance Variable importance (VIMP) for the test data. May be NULL.

- predicted Predicted values for the test data.
- predicted.oob OOB predicted values. May be NULL depending on context.
- quantile Estimated quantile values at the requested probabilities (quantile regression only).
- quantile.oob OOB quantile values. May be NULL.
- class (Classification only) Predicted class labels.
- class.oob (Classification only) OOB predicted class labels.
- **regrOutput** (Multivariate only) List of performance measures for multivariate regression outcomes.
- **clasOutput** (Multivariate only) List of performance measures for multivariate categorical outcomes.
- chf (Survival or competing risks) Cumulative hazard function (CHF).

chf.oob (Survival or competing risks) OOB CHF. May be NULL.

- survival (Survival only) Survival function estimates.
- survival.oob (Survival only) OOB survival function. May be NULL.
- time.interest (Survival or competing risks) Sorted unique event times.
- ndead (Survival or competing risks) Number of deaths observed.

cif (Competing risks only) Cumulative incidence function (CIF) for each event type.

cif.oob (Competing risks only) OOB CIF. May be NULL.

chf (Competing risks only) Cause-specific cumulative hazard function (CSCHF).

chf.oob (Competing risks only) OOB CSCHF. May be NULL.

Note

The dimensions and contents of returned objects depend on the forest family and whether youtcomes are available in the test data. In particular, performance-related components (e.g., error rate, VIMP) will be NULL if y-outcomes are missing.

For multivariate families, predicted values, VIMP, error rates, and performance metrics are stored in the lists regrOutput and clasOutput. These can be accessed using the helper functions get.mv.predicted, get.mv.vimp, and get.mv.error.

Author(s)

Hemant Ishwaran and Udaya B. Kogalur

References

Breiman L. (2001). Random forests, Machine Learning, 45:5-32.

Ishwaran H., Kogalur U.B., Blackstone E.H. and Lauer M.S. (2008). Random survival forests, *Ann. App. Statist.*, 2:841-860.

Ishwaran H. and Kogalur U.B. (2007). Random survival forests for R, Rnews, 7(2):25-31.

See Also

holdout.vimp.rfsrc, plot.competing.risk.rfsrc, plot.rfsrc, plot.survival.rfsrc, plot.variable.rfsrc, rfsrc, rfsrc.fast, stat.split.rfsrc, synthetic.rfsrc, vimp.rfsrc

Examples

```
## -----
## typical train/testing scenario
## ______
data(veteran, package = "randomForestSRC")
train <- sample(1:nrow(veteran), round(nrow(veteran) * 0.80))</pre>
veteran.grow <- rfsrc(Surv(time, status) ~ ., veteran[train, ])</pre>
veteran.pred <- predict(veteran.grow, veteran[-train, ])</pre>
print(veteran.grow)
print(veteran.pred)
## -----
## restore mode
## - if predict is called without specifying the test data
  the original training data is used and the forest is restored
##
## first train the forest
airq.obj <- rfsrc(Ozone ~ ., data = airquality)</pre>
## now we restore it and compare it to the original call
## they are identical
predict(airq.obj)
```

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```
print(airq.obj)
## we can retrieve various outputs that were not asked for in
## in the original call
## here we extract the proximity matrix
prox <- predict(airq.obj, proximity = TRUE)$proximity</pre>
print(prox[1:10,1:10])
## here we extract the number of times a variable was used to grow
## the grow forest
var.used <- predict(airq.obj, var.used = "by.tree")$var.used</pre>
print(head(var.used))
## -----
## prediction when test data has missing values
## -----
                                       _____
data(pbc, package = "randomForestSRC")
trn <- pbc[1:312,]</pre>
tst <- pbc[-(1:312),]</pre>
o <- rfsrc(Surv(days, status) ~ ., trn)</pre>
## default imputation method used by rfsrc
print(predict(o, tst, na.action = "na.impute"))
## random imputation
print(predict(o, tst, na.action = "na.random"))
## -----
## requesting different performance for classification
## ______
## default performance is misclassification
o <- rfsrc(Species~., iris)</pre>
print(o)
## get (normalized) brier performance
print(predict(o, perf.type = "brier"))
## -----
## vimp for each tree: illustrates get.tree
## ______
## regression analysis but no VIMP
o <- rfsrc(mpg~., mtcars)</pre>
## now extract VIMP for each tree using get.tree
vimp.tree <- do.call(rbind, lapply(1:o$ntree, function(b) {</pre>
    predict(o, get.tree = b, importance = TRUE)$importance
}))
## boxplot of tree VIMP
```

```
boxplot(vimp.tree, outline = FALSE, col = "cyan")
abline(h = 0, lty = 2, col = "red")
## summary information of tree VIMP
print(summary(vimp.tree))
## extract tree-averaged VIMP using importance=TRUE
## remember to set block.size to 1
print(predict(o, importance = TRUE, block.size = 1)$importance)
## use direct call to vimp() for tree-averaged VIMP
print(vimp(o, block.size = 1)$importance)
## -----
## vimp for just a few trees
## illustrates how to get vimp if you have a large data set
## ______
## survival analysis but no VIMP
data(pbc, package = "randomForestSRC")
o <- rfsrc(Surv(days, status) ~ ., pbc, ntree = 2000)</pre>
## get vimp for a small number of trees
print(predict(o, get.tree=1:250, importance = TRUE)$importance)
## -----
## case-specific vimp
## returns VIMP for each case
o <- rfsrc(mpg~., mtcars)</pre>
op <- predict(o, importance = TRUE, csv = TRUE)</pre>
csvimp <- get.mv.csvimp(op, standardize=TRUE)</pre>
print(csvimp)
## ------
## case-specific error rate
## returns tree-averaged error rate for each case
## -----
o <- rfsrc(mpg~., mtcars)</pre>
op <- predict(o, importance = TRUE, cse = TRUE)</pre>
cserror <- get.mv.cserror(op, standardize=TRUE)</pre>
print(cserror)
## -----
## predicted probability and predicted class labels are returned
## in the predict object for classification analyses
## ______
```

data(breast, package = "randomForestSRC")

predict.rfsrc

```
breast.obj <- rfsrc(status ~ ., data = breast[(1:100), ])</pre>
breast.pred <- predict(breast.obj, breast[-(1:100), ])</pre>
print(head(breast.pred$predicted))
print(breast.pred$class)
## unique feature of randomForestSRC
## cross-validation can be used when factor labels differ over
## training and test data
## -----
## first we convert all x-variables to factors
data(veteran, package = "randomForestSRC")
veteran2 <- data.frame(lapply(veteran, factor))</pre>
veteran2$time <- veteran$time</pre>
veteran2$status <- veteran$status
## split the data into unbalanced train/test data (25/75)
## the train/test data have the same levels, but different labels
train <- sample(1:nrow(veteran2), round(nrow(veteran2) * .25))</pre>
summary(veteran2[train,])
summary(veteran2[-train,])
## train the forest and use this to predict on test data
o.grow <- rfsrc(Surv(time, status) ~ ., veteran2[train, ])</pre>
o.pred <- predict(o.grow, veteran2[-train , ])</pre>
print(o.grow)
print(o.pred)
## even harder ... factor level not previously encountered in training
veteran3 <- veteran2[1:3, ]</pre>
veteran3$celltype <- factor(c("newlevel", "1", "3"))</pre>
o2.pred <- predict(o.grow, veteran3)</pre>
print(o2.pred)
## the unusual level is treated like a missing value but is not removed
print(o2.pred$xvar)
## -----
## example illustrating the flexibility of outcome = "test"
## illustrates restoration of forest via outcome = "test"
## -----
## first we train the forest
data(pbc, package = "randomForestSRC")
pbc.grow <- rfsrc(Surv(days, status) ~ ., pbc)</pre>
## use predict with outcome = TEST
pbc.pred <- predict(pbc.grow, pbc, outcome = "test")</pre>
## notice that error rates are the same!!
print(pbc.grow)
print(pbc.pred)
```

```
## note this is equivalent to restoring the forest
pbc.pred2 <- predict(pbc.grow)</pre>
print(pbc.grow)
print(pbc.pred)
print(pbc.pred2)
## similar example, but with na.action = "na.impute"
airq.obj <- rfsrc(Ozone ~ ., data = airquality, na.action = "na.impute")</pre>
print(airq.obj)
print(predict(airq.obj))
## ... also equivalent to outcome="test" but na.action = "na.impute" required
print(predict(airq.obj, airquality, outcome = "test", na.action = "na.impute"))
## classification example
iris.obj <- rfsrc(Species ~., data = iris)</pre>
print(iris.obj)
print(predict.rfsrc(iris.obj, iris, outcome = "test"))
## another example illustrating outcome = "test"
## unique way to check reproducibility of the forest
## ______
## training step
set.seed(542899)
data(pbc, package = "randomForestSRC")
train <- sample(1:nrow(pbc), round(nrow(pbc) * 0.50))</pre>
pbc.out <- rfsrc(Surv(days, status) ~ ., data=pbc[train, ])</pre>
## standard prediction call
pbc.train <- predict(pbc.out, pbc[-train, ], outcome = "train")</pre>
##non-standard predict call: overlays the test data on the grow forest
pbc.test <- predict(pbc.out, pbc[-train, ], outcome = "test")</pre>
## check forest reproducibilility by comparing "test" predicted survival
## curves to "train" predicted survival curves for the first 3 individuals
Time <- pbc.out$time.interest</pre>
matplot(Time, t(pbc.train$survival[1:3,]), ylab = "Survival", col = 1, type = "l")
matlines(Time, t(pbc.test$survival[1:3,]), col = 2)
## ------
## multivariate forest example
## ------
## train the forest
trn <- 1:20
o <- rfsrc(cbind(mpg, disp)~.,mtcars[trn,])</pre>
## print training results for each outcome
print(o, outcome.target="mpg")
print(o, outcome.target="disp")
```

print.rfsrc

```
## print test results for each outcome
p <- predict(o, mtcars[-trn,])
print(p, outcome.target="mpg")
print(p, outcome.target="disp")
```

print.rfsrc

Print Summary Output of a RF-SRC Analysis

Description

Print summary output from a RF-SRC analysis. This is the default print method for the package.

Usage

```
## S3 method for class 'rfsrc'
print(x, outcome.target = NULL, ...)
```

Arguments

| х | An object of class (rfsrc, grow), (rfsrc, synthetic), or (rfsrc, predict). |
|---|--|
| - | Character value for multivariate families specifying the target outcome to be used. The default is to use the first coordinate from the continuous outcomes (otherwise if none, the first coordinate from the categorical outcomes). |
| | Further arguments passed to or from other methods. |

Author(s)

Hemant Ishwaran and Udaya B. Kogalur

References

Ishwaran H. and Kogalur U.B. (2007). Random survival forests for R, Rnews, 7/2:25-31.

Examples

```
options(rf.cores=2, mc.cores=2)
iris.obj <- rfsrc(Species ~., data = iris, ntree=10)
print(iris.obj)</pre>
```

quantreg.rfsrc

Description

Grows a univariate or multivariate quantile regression forest and returns its conditional quantile and density values. Can be used for both training and testing purposes.

Usage

```
## S3 method for class 'rfsrc'
quantreg(formula, data, object, newdata,
   method = "local", splitrule = NULL, prob = NULL, prob.epsilon = NULL,
   oob = TRUE, fast = FALSE, maxn = 1e3, ...)
```

Arguments

| formula | A symbolic description of the model to be fit. Must be specified unless object is given. |
|--------------|--|
| data | Data frame containing the y-outcome and x-variables in the model. Must be specified unless object is given. |
| object | (Optional) A previously grown quantile regression forest. |
| newdata | (Optional) Test data frame used for prediction. Note that prediction on test data must always be done with the quantreg function and not the predict function. See example below. |
| method | Method used to calculate quantiles. Three methods are provided: (1) A varia- tion of the method used in Meinshausen (2006) based on forest weight (method = "forest"); (2) The Greenwald-Khanna algorithm, suited for big data, and specified by any one of the following: "gk", "GK", "G-K", "g-k"; (3) The de- fault method, method = "local", which uses the local adjusted cdf approach of Zhang et al. (2019). This does not rely on forest weights and is reasonably fast. See below for further discussion. |
| splitrule | The default action is local adaptive quantile regression splitting, but this can be over-ridden by the user. Not applicable to multivariate forests. See details below. |
| prob | Target quantile probabilities when training. If left unspecified, uses percentiles (1 through 99) for method = "forest", and for Greenwald-Khanna selects equally spaced percentiles optimized for accuracy (see below). |
| prob.epsilon | Greenwald-Khanna allowable error for quantile probabilities when training. |
| oob | Return OOB (out-of-bag) quantiles? If false, in-bag values are returned. |
| fast | Use fast random forests, rfsrc.fast, in place of rfsrc? Improves speed but may be less accurate. |
| maxn | Maximum number of unique y training values used when calculating the condi- tional density. |
| | Further arguments to be passed to the rfsrc function used for fitting the quantile regression forest. |

quantreg.rfsrc

Details

The most common method for calculating RF quantiles uses the method described in Meinshausen (2006) using forest weights. The forest weights method employed here (specified using method="forest"), however differs in that quantiles are estimated using a weighted local cumulative distribution function estimator. For this reason, results may differ from Meinshausen (2006). Moreover, results may also differ as the default splitting rule uses local adaptive quantile regression splitting instead of CART regression mean squared splitting which was used by Meinshausen (2006). Note that local adaptive quantile regression splitting is not available for multivariate forests which reverts to the default multivariate composite splitting rule. In multivariate regression, users however do have the option to over-ride this using Mahalanobis splitting by setting splitrule="mahalanobis"

A second method for estimating quantiles uses the Greenwald-Khanna (2001) algorithm (invoked by method="gk", "GK", "G-K" or "g-k"). While this will not be as accurate as forest weights, the high memory efficiency of Greenwald-Khanna makes it feasible to implement in big data settings unlike forest weights.

The Greenwald-Khanna algorithm is implemented roughly as follows. To form a distribution of values for each case, from which we sample to determine quantiles, we create a chain of values for the case as we grow the forest. Every time a case lands in a terminal node, we insert all of its co-inhabitants to its chain of values.

The best case scenario is when tree node size is 1 because each case gets only one insert into its chain for that tree. The worst case scenario is when node size is so large that trees stump. This is because each case receives insertions for the entire in-bag population.

What the user needs to know is that Greenwald-Khanna can become slow in counter-intutive settings such as when node size is large. The easy fix is to change the epsilon quantile approximation that is requested. You will see a significant speed-up just by doubling prob.epsilon. This is because the chains stay a lot smaller as epsilon increases, which is exactly what you want when node sizes are large. Both time and space requirements for the algorithm are affected by epsilon.

The best results for Greenwald-Khanna come from setting the number of quantiles equal to 2 times the sample size and epsilon to 1 over 2 times the sample size which is the default values used if left unspecified. This will be slow, especially for big data, and less stringent choices should be used if computational speed is of concern.

Finally, the default method, method="local", implements the locally adjusted cdf estimator of Zhang et al. (2019). This does not use forest weights and is reasonably fast and can be used for large data. However, this relies on the assumption of homogeneity of the error distribution, i.e. that errors are iid and therefore have equal variance. While this is reasonably robust to departures of homogeneity, there are instances where this may perform poorly; see Zhang et al. (2019) for details. If hetereogeneity is suspected we recommend method="forest".

Value

Returns the object quantreg containing quantiles for each of the requested probabilities (which can be conveniently extracted using get.quantile). Also contains the conditional density (and conditional cdf) for each case in the training data (or test data if provided) evaluated at each of the unique grow y-values. The conditional density can be used to calculate conditional moments, such as the mean and standard deviation. Use get.quantile.stat as a way to conveniently obtain these quantities.

For multivariate forests, returned values will be a list of length equal to the number of target outcomes.

Author(s)

Hemant Ishwaran and Udaya B. Kogalur

References

Greenwald M. and Khanna S. (2001). Space-efficient online computation of quantile summaries. *Proceedings of ACM SIGMOD*, 30(2):58-66.

Meinshausen N. (2006) Quantile regression forests, *Journal of Machine Learning Research*, 7:983-999.

Zhang H., Zimmerman J., Nettleton D. and Nordman D.J. (2019). Random forest prediction intervals. *The American Statistician*. 4:1-5.

See Also

rfsrc

Examples

```
## -----
## regression example
## -----
## standard call
o <- quantreg(mpg ~ ., mtcars)</pre>
## extract conditional quantiles
print(get.quantile(o))
print(get.quantile(o, c(.25, .50, .75)))
## extract conditional mean and standard deviation
print(get.quantile.stat(o))
## standardized continuous rank probabiliy score (crps) performance
plot(get.quantile.crps(o), type = "1")
## -----
## train/test regression example
## -----
## train (grow) call followed by test call
o <- quantreg(mpg ~ ., mtcars[1:20,])</pre>
o.tst <- quantreg(object = o, newdata = mtcars[-(1:20),])</pre>
## extract test set quantiles and conditional statistics
print(get.quantile(o.tst))
print(get.quantile.stat(o.tst))
```

quantreg.rfsrc

```
## ______
## quantile regression for Boston Housing using forest method
##______
if (library("mlbench", logical.return = TRUE)) {
 ## quantile regression with mse splitting
 data(BostonHousing)
 o <- quantreg(medv ~ ., BostonHousing, method = "forest", nodesize = 1)</pre>
 ## standardized continuous rank probabiliy score (crps)
 plot(get.quantile.crps(o), type = "1")
 ## quantile regression plot
 plot.quantreg(o, .05, .95)
 plot.quantreg(o, .25, .75)
 ## (A) extract 25,50,75 quantiles
 quant.dat <- get.quantile(o, c(.25, .50, .75))</pre>
 ## (B) values expected under normality
 quant.stat <- get.quantile.stat(o)</pre>
 c.mean <- quant.stat$mean</pre>
 c.std <- quant.stat$std</pre>
 q.25.est <- c.mean + qnorm(.25) * c.std</pre>
 q.75.est <- c.mean + qnorm(.75) * c.std</pre>
 ## compare (A) and (B)
 print(head(data.frame(quant.dat[, -2], q.25.est, q.75.est)))
}
## -----
## multivariate mixed outcomes example
## quantiles are only returned for the continous outcomes
## -----
dta <- mtcars
dta$cyl <- factor(dta$cyl)</pre>
dta$carb <- factor(dta$carb, ordered = TRUE)</pre>
o <- quantreg(cbind(carb, mpg, cyl, disp) ~., data = dta)</pre>
plot.quantreg(o, m.target = "mpg")
plot.quantreg(o, m.target = "disp")
## ------
## multivariate regression example using Mahalanobis splitting
## -----
```

dta <- mtcars

quantreg.rfsrc

```
o <- quantreg(cbind(mpg, disp) ~., data = dta, splitrule = "mahal")</pre>
plot.quantreg(o, m.target = "mpg")
plot.quantreg(o, m.target = "disp")
## ______
## example of quantile regression for ordinal data
## ______
## use the wine data for illustration
data(wine, package = "randomForestSRC")
## run quantile regression
o <- quantreg(quality ~ ., wine, ntree = 100)</pre>
## extract "probabilities" = density values
qo.dens <- o$quantreg$density</pre>
yunq <- o$quantreg$yunq</pre>
colnames(qo.dens) <- yunq</pre>
## convert y to a factor
yvar <- factor(cut(o$yvar, c(-1, yunq), labels = yunq))</pre>
## confusion matrix
qo.confusion <- get.confusion(yvar, qo.dens)</pre>
print(qo.confusion)
## normalized Brier score
cat("Brier:", 100 * get.brier.error(yvar, qo.dens), "\n")
## ______
## example of large data using Greenwald-Khanna algorithm
## ______
## load the data and do quick and dirty imputation
data(housing, package = "randomForestSRC")
housing <- impute(SalePrice ~ ., housing,</pre>
       ntree = 50, nimpute = 1, splitrule = "random")
## Greenwald-Khanna algorithm
## request a small number of quantiles
o <- quantreg(SalePrice ~ ., housing, method = "gk",</pre>
      prob = (1:20) / 20, prob.epsilon = 1 / 20, ntree = 250)
plot.quantreg(o)
## -----
## using mse splitting with local cdf method for large data
## load the data and do quick and dirty imputation
data(housing, package = "randomForestSRC")
```

```
housing <- impute(SalePrice ~ ., housing,</pre>
```

```
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```

rfsrc

```
ntree = 50, nimpute = 1, splitrule = "random")
```

```
## use mse splitting and reduce number of trees
o <- quantreg(SalePrice ~ ., housing, splitrule = "mse", ntree = 250)
plot.quantreg(o)</pre>
```

rfsrc

Fast Unified Random Forests for Survival, Regression, and Classification (RF-SRC)

Description

Fast OpenMP-parallel implementation of random forests (Breiman, 2001) for regression, classification, survival analysis (Ishwaran et al., 2008), competing risks (Ishwaran et al., 2012), multivariate outcomes (Segal and Xiao, 2011), unsupervised learning (Mantero and Ishwaran, 2020), quantile regression (Meinshausen, 2006; Zhang et al., 2019; Greenwald and Khanna, 2001), and imbalanced q-classification (O'Brien and Ishwaran, 2019).

The package supports both deterministic and randomized splitting rules (Geurts et al., 2006; Ishwaran, 2015) across all families. Multiple types of variable importance (VIMP) are available, including holdout VIMP and confidence regions (Ishwaran and Lu, 2019), for both individual and grouped variables. Variable selection can be performed using minimal depth (Ishwaran et al., 2010, 2011). Fast interfaces for missing data imputation are provided using several forest-based algorithms (Tang and Ishwaran, 2017).

Highlighted updates:

- 1. For variable selection, we recommend using **VarPro**, an R package for model-independent variable selection using rule-based variable priority. It supports regression, classification, survival analysis, and includes a new mode for unsupervised learning. See https://www.varprotools.org for more information.
- 2. For computational speed, the default VIMP method has changed from "permute" (Breiman-Cutler permutation) to "anti" (importance = "anti" or importance = TRUE). While faster, this may be less accurate in settings such as highly imbalanced classification. To revert to permutation VIMP, use importance = "permute".

This is the main entry point to the **randomForestSRC** package. For more information on OpenMP support and the package as a whole, see package?randomForestSRC.

Usage

```
rfsrc(formula, data, ntree = 500,
mtry = NULL, ytry = NULL,
nodesize = NULL, nodedepth = NULL,
splitrule = NULL, nsplit = NULL,
importance = c(FALSE, TRUE, "none", "anti", "permute", "random"),
```

```
block.size = if (any(is.element(as.character(importance),
                   c("none", "FALSE")))) NULL else 10,
bootstrap = c("by.root", "none", "by.user"),
samptype = c("swor", "swr"), samp = NULL, membership = FALSE,
sampsize = if (samptype == "swor") function(x){x * .632} else function(x){x},
na.action = c("na.omit", "na.impute"), nimpute = 1,
ntime = 150, cause,
perf.type = NULL,
proximity = FALSE, distance = FALSE, forest.wt = FALSE,
xvar.wt = NULL, yvar.wt = NULL, split.wt = NULL, case.wt = NULL,
case.depth = FALSE,
forest = TRUE,
save.memory = FALSE,
var.used = c(FALSE, "all.trees", "by.tree"),
split.depth = c(FALSE, "all.trees", "by.tree"),
seed = NULL,
do.trace = FALSE,
statistics = FALSE,
...)
```

convenient interface for growing a CART tree
rfsrc.cart(formula, data, ntree = 1, mtry = ncol(data), bootstrap = "none", ...)

Arguments

| formula | A formula describing the model to fit. Interaction terms are not supported. If missing, unsupervised splitting is used. |
|------------|--|
| data | Data frame containing the response and predictor variables. |
| ntree | Number of trees to grow. |
| mtry | Number of candidate variables randomly selected at each split. Defaults: regression uses p/3, others use sqrt(p); rounded up. |
| ytry | Number of pseudo-response variables randomly selected for unsupervised split- ting. Default is 1. |
| nodesize | Minimum terminal node size. Defaults: survival/competing risks (15), regression (5), classification (1), mixed/unsupervised (3). |
| nodedepth | Maximum tree depth. Ignored by default. |
| splitrule | Splitting rule. See Details. |
| nsplit | Number of random split points per variable. 0 uses all values (deterministic). Default is 10. |
| importance | Variable importance (VIMP) method. Choices: FALSE, TRUE, "none", "anti", "permute", "random". Default is "none". VIMP can be computed later using vimp or predict. |
| block.size | Controls frequency of cumulative error/VIMP updates. Default is 10 if impor- tance is requested; otherwise NULL. See Details. |
| bootstrap | Bootstrap method. Options: "by.root" (default), "by.user", or "none" (no OOB error possible). |

| samptype | Sampling type for by.root bootstrap. Options: "swor" (without replacement, default), "swr" (with replacement). |
|-------------|--|
| samp | Bootstrap weights (only for bootstrap="by.user"). A matrix of size n x ntree giving in-bag counts per tree. |
| membership | Return inbag and terminal node membership? |
| sampsize | Bootstrap sample size (used when bootstrap="by.root"). Defaults: 0.632 x n for swor; n for swr. Can also be numeric. |
| na.action | Missing data handling. "na.omit" (default) removes rows with any NA; "na.impute" performs fast internal imputation. See also impute. |
| nimpute | Number of iterations for internal imputation. If >1, OOB error rates may be optimistic. |
| ntime | For survival models: number or grid of time points used in ensemble estimation. If NULL or 0, uses all event times. |
| cause | For competing risks: event of interest (1 to J), or a vector of weights over all J events. Defaults to an average over all events. |
| perf.type | Optional performance metric for prediction, VIMP, and error. Defaults to the family-specific metric. "none" disables performance. See Details. |
| proximity | Compute proximity matrix? Options: "inbag", "oob", "all", TRUE (inbag), or FALSE. |
| distance | Compute pairwise distances between cases? Similar options as proximity. See Details. |
| forest.wt | Return forest weight matrix? Same options as proximity. Default is TRUE (inbag). |
| xvar.wt | Optional weights on x-variables for sampling at splits. Does not need to sum to 1. Defaults to uniform. |
| yvar.wt | Weights on response variables (for multivariate regression). Used when y is high-dimensional. |
| split.wt | Weights applied to each variable's split statistic. Higher weight increases likeli- hood of splitting. |
| case.wt | Sampling weights for cases in the bootstrap. Higher values increase selection probability. See class imbalance example. |
| case.depth | Return matrix recording depth of first split for each case? Default is FALSE. |
| forest | Save forest object for future prediction? Set FALSE if prediction is not needed. |
| save.memory | Reduce memory usage by avoiding storage of prediction quantities. Recom- mended for large survival or competing risk forests. |
| var.used | Return variable usage statistics? Options: FALSE, "all.trees", "by.tree". |
| split.depth | Return minimal depth of splits for each variable? Options: FALSE, "all.trees", "by.tree". See Details. |
| seed | Integer seed for reproducibility (negative values only). |
| do.trace | Print progress updates every do.trace seconds. |
| statistics | Return raw split statistics? Use stat.split to extract. |
| | Additional arguments passed to or from other methods. |

Details

1. Types of forests

The type of forest is automatically inferred from the outcome and formula. Supported forest types include:

- Regression forests for continuous outcomes.
- Classification forests for factor outcomes.
- Multivariate forests for continuous, categorical, or mixed outcomes.
- Unsupervised forests when no outcome is specified.
- Survival forests for right-censored time-to-event data.
- Competing risk forests for multi-event survival settings.
- 2. Splitting
 - (a) Splitting rules are set using the splitrule option.
 - (b) Random splitting is invoked via splitrule = "random".
 - (c) Use nsplit to enable randomized splitting and improve speed; see *Improving computational speed*.
- 3. Available splitting rules
 - Regression
 - (a) "mse" (default): weighted mean squared error (Breiman et al., 1984).
 - (b) "quantile.regr": quantile regression via check-loss; see quantreg.rfsrc.
 - (c) "la.quantile.regr": local adaptive quantile regression.
 - Classification
 - (a) "gini" (default): Gini index.
 - (b) "auc": AUC-based splitting; appropriate for imbalanced data.
 - (c) "entropy": entropy-based splitting.
 - Survival
 - (a) "logrank" (default): log-rank splitting.
 - (b) "bs.gradient": Brier score gradient splitting. Uses 90th percentile of observed times by default, or set prob.
 - (c) "logrankscore": log-rank score splitting.
 - Competing risks (see Ishwaran et al., 2014)
 - (a) "logrankCR" (default): Gray's test-based weighted log-rank splitting.
 - (b) "logrank": cause-specific weighted log-rank; use cause to target specific events.
 - Multivariate
 - (a) Default: normalized composite splitting (Tang and Ishwaran, 2017).
 - (b) "mahalanobis": Mahalanobis splitting with optional covariance matrix; for multi-variate regression.
 - **Unsupervised** Splitting uses pseudo-outcomes and the composite rule. See sidClustering for advanced unsupervised analysis.
 - **Custom splitting** Custom rules can be defined using splitCustom.c. Up to 16 rules per family are allowed. Use "custom", "custom1", etc. Compilation required.
- 4. Improving computational speed

See rfsrc.fast. Strategies include:
- Increase nodesize.
- Set save.memory = TRUE for large survival or competing risk models.
- Set block.size = NULL to avoid repeated cumulative error computation.
- Use perf. type = "none" to disable VIMP and C-index calculations.
- Set nsplit to a small integer (e.g., 1-10).
- Reduce bootstrap size with sampsize, samptype.
- Set ntime to a coarse grid (e.g., 50) for survival models.
- Pre-filter variables; use max.subtree for fast variable selection.
- 5. Prediction Error

Error is computed using OOB data:

- Regression: mean squared error.
- Classification: misclassification rate, Brier score, AUC.
- Survival: C-error = 1 Harrell's concordance index.

If bootstrap = "none", OOB error is unavailable. Use predict.rfsrc for cross-validation error instead.

6. Variable Importance (VIMP)

VIMP methods:

- "permute": permutation VIMP (Breiman-Cutler).
- "random": randomized left/right assignment.
- "anti" (default): anti-split assignment.

The block.size option controls granularity. For confidence intervals, see subsampling. Also see holdout.vimp for a more conservative variant.

7. Multivariate Forests

Use:

```
rfsrc(Multivar(y1, ..., yd) ~ ., data)
```

or

rfsrc(cbind(y1, ..., yd) ~ ., data)

Use get.mv.formula, get.mv.predicted, get.mv.error for multivariate extraction.

8. Unsupervised Forests

Use:

rfsrc(data = X)

or

rfsrc(Unsupervised(ytry) ~ ., data = X)

Random subsets of ytry pseudo-responses are used for each mtry variable. No performance metrics are computed.

- 9. Survival, Competing Risks
 - Survival: use Surv(time, status) ~ .. Status must be 0 (censored) or 1 (event).
 - Competing risks: status = 0 (censored), 1-J (event types). Use cause to target specific events.

10. Missing data imputation

Use na.action = "na.impute". Iteration with nimpute > 1 replaces missing values using OOB predictions. Observations or variables with all missing values are removed.

11. Allowable data types and factors

Variables must be numeric, integer, factor, or logical. Non-factors are coerced to numeric. For unordered factors, all complementary subsets are considered for splits.

Factor levels are mapped to ensure consistency across training/test data. Consider converting factors to numeric for high-dimensional settings.

Value

An object of class (rfsrc, grow) with the following components:

call The original call to rfsrc.

family The family used in the analysis.

n Sample size after applying na.action.

ntree Number of trees grown.

mtry Number of variables randomly selected at each node.

nodesize Minimum terminal node size.

nodedepth Maximum depth allowed for each tree.

splitrule Splitting rule used.

nsplit Number of random split points.

yvar Response values.

yvar.names Character vector of response variable names.

xvar Data frame of predictor variables.

xvar.names Character vector of predictor variable names.

xvar.wt Non-negative weights specifying the selection probability of each variable.

split.wt Non-negative weights adjusting each variable's split statistic.

cause.wt Weights for composite competing risk splitting.

leaf.count Number of terminal nodes per tree. A value of 0 indicates a rejected tree (may occur with missing data); a value of 1 indicates a stump.

proximity Proximity matrix indicating how often case pairs fall in the same terminal node.

forest Forest object, returned if forest=TRUE. Required for prediction and most wrappers.

forest.wt Forest weight matrix.

membership Terminal node membership matrix (rows: trees; columns: cases).

inbag Inbag count matrix (rows: trees; columns: cases).

var.used Number of times each variable is used to split a node.

imputed.indv Indices of individuals with missing values.

imputed.data Imputed dataset with responses followed by predictors.

split.depth Matrix or array recording minimal split depth of variables by case and tree. **node.stats** Split statistics returned if statistics=TRUE, parsed with stat.split. err.rate Cumulative OOB error rate. err.block.rate Cumulative error per ensemble block (size defined by block.size). If block.size = 1, error per tree. importance Variable importance (VIMP) for each predictor. predicted In-bag predicted values. predicted.oob Out-of-bag (OOB) predicted values. class (Classification) In-bag predicted class labels. class.oob (Classification) OOB predicted class labels. regrOutput (Multivariate) List of performance results for continuous outcomes. clasOutput (Multivariate) List of performance results for categorical outcomes. survival (Survival) In-bag survival functions. survival.oob (Survival) OOB survival functions. chf (Survival or competing risks) In-bag cumulative hazard function. chf.oob (Survival or competing risks) OOB cumulative hazard function. time.interest (Survival or competing risks) Unique sorted event times. ndead (Survival or competing risks) Total number of observed events. **cif** (Competing risks) In-bag cumulative incidence function by cause. cif.oob (Competing risks) OOB cumulative incidence function by cause.

Note

Values returned by the forest depend on the family:

- Regression: predicted and predicted.oob are vectors of predicted values.
- **Classification**: predicted and predicted.oob are matrices of class probabilities. VIMP and performance metrics are returned as a matrix with J+1 columns (J = number of classes). The first column ("all") gives unconditional results; remaining columns give class-conditional results.
- **Survival**: predicted contains mortality estimates (Ishwaran et al., 2008). These are calibrated to the number of expected events under identical covariate profiles. Also returned are matrices of the survival function and CHF for each individual over time.interest.
- **Competing risks**: predicted contains expected life years lost by cause (Ishwaran et al., 2013). Also returned are three-dimensional arrays for CIF and CSCHF indexed by case, time, and event type.
- **Multivariate**: Predictions, VIMP, and error rates are returned in regrOutput and clasOutput. Use get.mv.predicted, get.mv.vimp, and get.mv.error to extract results.

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See Also

find.interaction.rfsrc,

get.tree.rfsrc,

holdout.vimp.rfsrc,

imbalanced.rfsrc, impute.rfsrc,

max.subtree.rfsrc,

partial.rfsrc, plot.competing.risk.rfsrc, plot.rfsrc, plot.survival.rfsrc, plot.variable.rfsrc, predict.rfsrc, print.rfsrc,

quantreg.rfsrc,

rfsrc, rfsrc.anonymous, rfsrc.cart, rfsrc.fast,

sidClustering.rfsrc,

stat.split.rfsrc, subsample.rfsrc, synthetic.rfsrc,

tune.rfsrc,

vimp.rfsrc

Examples

##----## survival analysis
##----## veteran data
randomized trial of two treatment regimens for lung cancer

data(veteran, package = "randomForestSRC")
v.obj <- rfsrc(Surv(time, status) ~ ., data = veteran, block.size = 1)</pre>

plot tree number 3

```
plot(get.tree(v.obj, 3))
## print results of trained forest
print(v.obj)
## plot results of trained forest
plot(v.obj)
## plot survival curves for first 10 individuals -- direct way
matplot(v.obj$time.interest, 100 * t(v.obj$survival.oob[1:10, ]),
    xlab = "Time", ylab = "Survival", type = "1", lty = 1)
## plot survival curves for first 10 individuals
## using function "plot.survival"
plot.survival(v.obj, subset = 1:10)
## obtain Brier score using KM and RSF censoring distribution estimators
bs.km <- get.brier.survival(v.obj, cens.model = "km")$brier.score</pre>
bs.rsf <- get.brier.survival(v.obj, cens.model = "rfsrc")$brier.score</pre>
## plot the brier score
plot(bs.km, type = "s", col = 2)
lines(bs.rsf, type ="s", col = 4)
legend("topright", legend = c("cens.model = km", "cens.model = rfsrc"), fill = c(2,4))
## plot CRPS (continuous rank probability score) as function of time
## here's how to calculate the CRPS for every time point
trapz <- randomForestSRC:::trapz</pre>
time <- v.obj$time.interest</pre>
crps.km <- sapply(1:length(time), function(j) {</pre>
  trapz(time[1:j], bs.km[1:j, 2] / diff(range(time[1:j])))
})
crps.rsf <- sapply(1:length(time), function(j) {</pre>
  trapz(time[1:j], bs.rsf[1:j, 2] / diff(range(time[1:j])))
})
plot(time, crps.km, ylab = "CRPS", type = "s", col = 2)
lines(time, crps.rsf, type ="s", col = 4)
legend("bottomright", legend=c("cens.model = km", "cens.model = rfsrc"), fill=c(2,4))
## fast nodesize optimization for veteran data
## optimal nodesize in survival is larger than other families
## see the function "tune" for more examples
tune.nodesize(Surv(time,status) ~ ., veteran)
## Primary biliary cirrhosis (PBC) of the liver
data(pbc, package = "randomForestSRC")
pbc.obj <- rfsrc(Surv(days, status) ~ ., pbc)</pre>
print(pbc.obj)
```

save.memory example for survival

```
## growing many deep trees creates memory issue without this option!
data(pbc, package = "randomForestSRC")
print(rfsrc(Surv(days, status) ~ ., pbc, splitrule = "random",
          ntree = 25000, nodesize = 1, save.memory = TRUE))
##-----
## trees can be plotted for any family
## see get.tree for details and more examples
##-----
## survival where factors have many levels
data(veteran, package = "randomForestSRC")
vd <- veteran
vd$celltype=factor(vd$celltype)
vd$diagtime=factor(vd$diagtime)
vd.obj <- rfsrc(Surv(time,status)~., vd, ntree = 100, nodesize = 5)
plot(get.tree(vd.obj, 3))
## classification
iris.obj <- rfsrc(Species ~., data = iris)</pre>
plot(get.tree(iris.obj, 25, class.type = "bayes"))
plot(get.tree(iris.obj, 25, target = "setosa"))
plot(get.tree(iris.obj, 25, target = "versicolor"))
plot(get.tree(iris.obj, 25, target = "virginica"))
## -----
## simple example of VIMP using iris classification
## ______
## directly from trained forest
print(rfsrc(Species~., iris, importance=TRUE)$importance)
## VIMP (and performance) use misclassification error by default
## but brier prediction error can be requested
print(rfsrc(Species~., iris, importance=TRUE, perf.type="brier")$importance)
## example using vimp function (see vimp help file for details)
iris.obj <- rfsrc(Species ~., data = iris)</pre>
print(vimp(iris.obj)$importance)
print(vimp(iris.obj,perf.type="brier")$importance)
## example using hold out vimp (see holdout.vimp help file for details)
print(holdout.vimp(Species~.,iris)$importance)
print(holdout.vimp(Species~.,iris,perf.type="brier")$importance)
## -----
## confidence interval for vimp using subsampling
## compare with holdout vimp
## ______
```

new York air quality measurements

```
o <- rfsrc(Ozone ~ ., data = airquality)</pre>
so <- subsample(o)</pre>
plot(so)
## compare with holdout vimp
print(holdout.vimp(Ozone ~ ., data = airquality)$importance)
## example of imputation in survival analysis
data(pbc, package = "randomForestSRC")
pbc.obj2 <- rfsrc(Surv(days, status) ~ ., pbc, na.action = "na.impute")</pre>
## same as above but iterate the missing data algorithm
pbc.obj3 <- rfsrc(Surv(days, status) ~ ., pbc,</pre>
        na.action = "na.impute", nimpute = 3)
## fast way to impute data (no inference is done)
## see impute for more details
pbc.imp <- impute(Surv(days, status) ~ ., pbc, splitrule = "random")</pre>
##-----
## compare RF-SRC to Cox regression
## Illustrates C-error and Brier score measures of performance
## assumes "pec" and "survival" libraries are loaded
##-----
if (library("survival", logical.return = TRUE)
   & library("pec", logical.return = TRUE)
   & library("prodlim", logical.return = TRUE))
{
 ##prediction function required for pec
 predictSurvProb.rfsrc <- function(object, newdata, times, ...){</pre>
   ptemp <- predict(object,newdata=newdata,...)$survival</pre>
   pos <- sindex(jump.times = object$time.interest, eval.times = times)</pre>
   p <- cbind(1,ptemp)[, pos + 1]</pre>
   if (NROW(p) != NROW(newdata) || NCOL(p) != length(times))
     stop("Prediction failed")
   р
 }
 ## data, formula specifications
 data(pbc, package = "randomForestSRC")
 pbc.na <- na.omit(pbc) ##remove NA's</pre>
 surv.f <- as.formula(Surv(days, status) ~ .)</pre>
 pec.f <- as.formula(Hist(days,status) ~ 1)</pre>
 ## run cox/rfsrc models
 ## for illustration we use a small number of trees
 cox.obj <- coxph(surv.f, data = pbc.na, x = TRUE)</pre>
```

}

```
rfsrc.obj <- rfsrc(surv.f, pbc.na, ntree = 150)</pre>
 ## compute bootstrap cross-validation estimate of expected Brier score
 ## see Mogensen, Ishwaran and Gerds (2012) Journal of Statistical Software
 set.seed(17743)
 prederror.pbc <- pec(list(cox.obj,rfsrc.obj), data = pbc.na, formula = pec.f,</pre>
                       splitMethod = "bootcv", B = 50)
 print(prederror.pbc)
 plot(prederror.pbc)
 ## compute out-of-bag C-error for cox regression and compare to rfsrc
 rfsrc.obj <- rfsrc(surv.f, pbc.na)</pre>
 cat("out-of-bag Cox Analysis ...", "\n")
 cox.err <- sapply(1:100, function(b) {</pre>
    if (b%%10 == 0) cat("cox bootstrap:", b, "\n")
   train <- sample(1:nrow(pbc.na), nrow(pbc.na), replace = TRUE)</pre>
   cox.obj <- tryCatch({coxph(surv.f, pbc.na[train, ])}, error=function(ex){NULL})</pre>
   if (!is.null(cox.obj)) {
   get.cindex(pbc.na$days[-train], pbc.na$status[-train], predict(cox.obj, pbc.na[-train, ]))
   } else NA
 })
 cat("\n\t00B error rates\n\n")
 cat("\tRSF : ", rfsrc.obj$err.rate[rfsrc.obj$ntree], "\n")
 cat("\tCox regression : ", mean(cox.err, na.rm = TRUE), "\n")
               -----
##-----
## competing risks
## WIHS analysis
## cumulative incidence function (CIF) for HAART and AIDS stratified by IDU
data(wihs, package = "randomForestSRC")
wihs.obj <- rfsrc(Surv(time, status) ~ ., wihs, nsplit = 3, ntree = 100)
plot.competing.risk(wihs.obj)
cif <- wihs.obj$cif.oob</pre>
Time <- wihs.obj$time.interest</pre>
idu <- wihs$idu
cif.haart <- cbind(apply(cif[,,1][idu == 0,], 2, mean),</pre>
                  apply(cif[,,1][idu == 1,], 2, mean))
cif.aids <- cbind(apply(cif[,,2][idu == 0,], 2, mean),</pre>
                  apply(cif[,,2][idu == 1,], 2, mean))
matplot(Time, cbind(cif.haart, cif.aids), type = "1",
       lty = c(1,2,1,2), col = c(4, 4, 2, 2), lwd = 3,
       ylab = "Cumulative Incidence")
legend("topleft",
      legend = c("HAART (Non-IDU)", "HAART (IDU)", "AIDS (Non-IDU)", "AIDS (IDU)"),
      lty = c(1,2,1,2), col = c(4, 4, 2, 2), lwd = 3, cex = 1.5)
```

illustrates the various splitting rules ## illustrates event specific and non-event specific variable selection

```
if (library("survival", logical.return = TRUE)) {
 ## use the pbc data from the survival package
 ## events are transplant (1) and death (2)
 data(pbc, package = "survival")
 pbc$id <- NULL</pre>
 ## modified Gray's weighted log-rank splitting
 ## (equivalent to cause=c(1,1) and splitrule="logrankCR")
 pbc.cr <- rfsrc(Surv(time, status) ~ ., pbc)</pre>
 ## log-rank cause-1 specific splitting and targeted VIMP for cause 1
 pbc.log1 <- rfsrc(Surv(time, status) ~ ., pbc,</pre>
            splitrule = "logrankCR", cause = c(1,0), importance = TRUE)
 ## log-rank cause-2 specific splitting and targeted VIMP for cause 2
 pbc.log2 <- rfsrc(Surv(time, status) ~ ., pbc,</pre>
            splitrule = "logrankCR", cause = c(0,1), importance = TRUE)
 ## extract VIMP from the log-rank forests: event-specific
 ## extract minimal depth from the Gray log-rank forest: non-event specific
 var.perf <- data.frame(md = max.subtree(pbc.cr)$order[, 1],</pre>
                      vimp1 = 100 * pbc.log1$importance[ ,1],
                      vimp2 = 100 * pbc.log2$importance[ ,2])
 print(var.perf[order(var.perf$md), ], digits = 2)
}
## ______
## regression analysis
## -----
## new York air quality measurements
airq.obj <- rfsrc(Ozone ~ ., data = airquality, na.action = "na.impute")
# partial plot of variables (see plot.variable for more details)
plot.variable(airq.obj, partial = TRUE, smooth.lines = TRUE)
## motor trend cars
mtcars.obj <- rfsrc(mpg ~ ., data = mtcars)</pre>
## ------
## regression with custom bootstrap
## ------
ntree <- 25
n <- nrow(mtcars)</pre>
s.size <- n / 2
swr <- TRUE
samp <- randomForestSRC:::make.sample(ntree, n, s.size, swr)</pre>
o <- rfsrc(mpg ~ ., mtcars, bootstrap = "by.user", samp = samp)</pre>
## -----
```

```
## classification analysis
## -----
## iris data
iris.obj <- rfsrc(Species ~., data = iris)</pre>
## wisconsin prognostic breast cancer data
data(breast, package = "randomForestSRC")
breast.obj <- rfsrc(status ~ ., data = breast, block.size=1)</pre>
plot(breast.obj)
## -----
## big data set, reduce number of variables using simple method
## ------
## use Iowa housing data set
data(housing, package = "randomForestSRC")
## original data contains lots of missing data, use fast imputation
## however see impute for other methods
housing2 <- impute(data = housing, fast = TRUE)</pre>
## run shallow trees to find variables that split any tree
xvar.used <- rfsrc(SalePrice ~., housing2, ntree = 250, nodedepth = 4,</pre>
                var.used="all.trees", mtry = Inf, nsplit = 100)$var.used
## now fit forest using filtered variables
xvar.keep <- names(xvar.used)[xvar.used >= 1]
o <- rfsrc(SalePrice~., housing2[, c("SalePrice", xvar.keep)])</pre>
print(o)
## ______
## imbalanced classification data
## see the "imbalanced" function for further details
##
## (a) use balanced random forests with undersampling of the majority class
## Specifically let n0, n1 be sample sizes for majority, minority
## cases. We sample 2 x n1 cases with majority, minority cases chosen
## with probabilities n1/n, n0/n where n=n0+n1
##
## (b) balanced random forests using "imbalanced"
##
## (c) q-classifier (RFQ) using "imbalanced"
##
## ______
## Wisconsin breast cancer example
data(breast, package = "randomForestSRC")
breast <- na.omit(breast)</pre>
## balanced random forests - brute force
y <- breast$status
obdirect <- rfsrc(status ~ ., data = breast, nsplit = 10,</pre>
```

```
case.wt = randomForestSRC:::make.wt(y),
           sampsize = randomForestSRC:::make.size(y))
print(obdirect)
print(get.imbalanced.performance(obdirect))
## balanced random forests - using "imbalanced"
ob <- imbalanced(status ~ ., data = breast, method = "brf")</pre>
print(ob)
print(get.imbalanced.performance(ob))
## q-classifier (RFQ) - using "imbalanced"
oq <- imbalanced(status ~ ., data = breast)</pre>
print(oq)
print(get.imbalanced.performance(oq))
## q-classifier (RFQ) - with auc splitting
oqauc <- imbalanced(status ~ ., data = breast, splitrule = "auc")</pre>
print(oqauc)
print(get.imbalanced.performance(oqauc))
## -----
## unsupervised analysis
## ------
## two equivalent ways to implement unsupervised forests
mtcars.unspv <- rfsrc(Unsupervised() ~., data = mtcars)</pre>
mtcars2.unspv <- rfsrc(data = mtcars)</pre>
## illustration of sidClustering for the mtcars data
## see sidClustering for more details
mtcars.sid <- sidClustering(mtcars, k = 1:10)</pre>
print(split(mtcars, mtcars.sid$cl[, 3]))
print(split(mtcars, mtcars.sid$cl[, 10]))
## -----
## bivariate regression using Mahalanobis splitting
## also illustrates user specified covariance matrix
## -----
if (library("mlbench", logical.return = TRUE)) {
 ## load boston housing data, specify the bivariate regression
 data(BostonHousing)
 f <- formula("Multivar(lstat, nox) ~.")</pre>
 ## Mahalanobis splitting
 bh.mreg <- rfsrc(f, BostonHousing, importance = TRUE, splitrule = "mahal")</pre>
 ## performance error and vimp
 vmp <- get.mv.vimp(bh.mreg)</pre>
 pred <- get.mv.predicted(bh.mreg)</pre>
```

```
## standardized error and vimp
 err.std <- get.mv.error(bh.mreg, standardize = TRUE)</pre>
 vmp.std <- get.mv.vimp(bh.mreg, standardize = TRUE)</pre>
 ## same analysis, but with user specified covariance matrix
 sigma <- cov(BostonHousing[, c("lstat","nox")])</pre>
 bh.mreg2 <- rfsrc(f, BostonHousing, splitrule = "mahal", sigma = sigma)</pre>
}
## -----
## multivariate mixed forests (nutrigenomic study)
## study effects of diet, lipids and gene expression for mice
## diet, genotype and lipids used as the multivariate y
## genes used for the x features
## ______
                            ------
## load the data (data is a list)
data(nutrigenomic, package = "randomForestSRC")
## assemble the multivariate y data
ydta <- data.frame(diet = nutrigenomic$diet,</pre>
                 genotype = nutrigenomic$genotype,
                 nutrigenomic$lipids)
## multivariate mixed forest call
## uses "get.mv.formula" for conveniently setting formula
mv.obj <- rfsrc(get.mv.formula(colnames(ydta)),</pre>
              data.frame(ydta, nutrigenomic$genes),
importance=TRUE, nsplit = 10)
## print results for diet and genotype y values
print(mv.obj, outcome.target = "diet")
print(mv.obj, outcome.target = "genotype")
## extract standardized VIMP
svimp <- get.mv.vimp(mv.obj, standardize = TRUE)</pre>
## plot standardized VIMP for diet, genotype and lipid for each gene
boxplot(t(svimp), col = "bisque", cex.axis = .7, las = 2,
       outline = FALSE,
       ylab = "standardized VIMP",
       main = "diet/genotype/lipid VIMP for each gene")
## -----
## illustrates yvar.wt which sets the probability of selecting
## the response variables in multivariate regression
## ______
## use mtcars: add fake responses
```

mult.mtcars <- cbind(mtcars, mtcars\$mpg, mtcars\$mpg)</pre>

```
names(mult.mtcars) = c(names(mtcars), "mpg2", "mpg3")
## noise up the fake responses
mult.mtcars$mpg2 <- sample(mtcars$mpg)</pre>
mult.mtcars$mpg3 <- sample(mtcars$mpg)</pre>
formula = as.formula(Multivar(mpg, mpg2, mpg3) ~ .)
## select 2 of the 3 responses randomly at each split with an associated weight vector.
## choose the noisy y responses which should degrade performance
yvar.wt = c(0.000001, 0.5, 0.5)
ytry = 2
mult.grow <- rfsrc(formula = formula, data = mult.mtcars, ytry = ytry, yvar.wt = yvar.wt)</pre>
print(mult.grow)
print(get.mv.error(mult.grow))
## Also, compare the following two results, as they should be similar:
yvar.wt = c(1.0, 00000.1, 00000.1)
ytry = 1
result1 = rfsrc(formula = formula, data = mult.mtcars, ytry = ytry, yvar.wt = yvar.wt)
result2 = rfsrc(mpg ~ ., mtcars)
print(get.mv.error(result1))
print(get.mv.error(result2))
## ------
## custom splitting using the pre-coded examples
## ______
## motor trend cars
mtcars.obj <- rfsrc(mpg ~ ., data = mtcars, splitrule = "custom")</pre>
## iris analysis
iris.obj <- rfsrc(Species ~., data = iris, splitrule = "custom1")</pre>
## WIHS analysis
wihs.obj <- rfsrc(Surv(time, status) ~ ., wihs, nsplit = 3,</pre>
                 ntree = 100, splitrule = "custom1")
```

rfsrc.anonymous Anonymous Random Forests

Description

Anonymous random forests is carefully modified to ensure that the original training data is not retained. This enables users to share the trained forest with others without disclosing the underlying

rfsrc.anonymous

data.

Usage

```
rfsrc.anonymous(formula, data, forest = TRUE, ...)
```

Arguments

| formula | A symbolic description of the model to be fit. If missing, unsupervised splitting is performed. |
|---------|---|
| data | A data frame containing the y-outcome and x-variables. |
| forest | Logical. Should the forest object be returned? Required for prediction on new data and by many other package functions. |
| | Additional arguments passed to rfsrc. See the rfsrc help file for full details. |

Details

This function calls **rfsrc** and returns a forest object with the original training data removed. This enables users to share their forest while preserving the privacy of their data.

To enable prediction on new (test) data, certain minimal information from the training data must still be retained. This includes:

- Names of the original variables.
- · For factor variables, the levels of each factor.
- Summary statistics used for imputation: the mean for continuous variables and the most frequent class for factors.
- Tree topology, including split points used to grow the trees.

For maximal privacy, users are strongly encouraged to replace variable names with non-identifiable labels and convert all variables to continuous format when possible. If factor variables are used, their levels should also be anonymized. However, the user is solely responsible for de-identifying the data and verifying that privacy is maintained. We provide NO GUARANTEES regarding data confidentiality.

Missing data handling: Anonymous forests do not support imputation of training data. The option na.action = "na.impute" is automatically downgraded to "na.omit". If training data contain missing values, we recommend pre-imputing them using impute.

Test data, however, *can* be imputed at prediction time:

- na.action = "na.impute" performs a fast imputation by replacing missing values with the training mean (for numeric variables) or most frequent class (for factors).
- na.action = "na.random" uses a fast random draw from training distributions for imputation.

Although anonymous forests are compatible with many package functions, they are only guaranteed to work with functions that do not explicitly require access to the original training data.

An object of class (rfsrc, grow, anonymous).

Author(s)

Hemant Ishwaran and Udaya B. Kogalur

See Also

rfsrc

Examples

```
## regression
## ------
print(rfsrc.anonymous(mpg ~ ., mtcars))
## -----
## plot anonymous regression tree (using get.tree)
## TBD CURRENTLY NOT IMPLEMENTED
## -----
## plot(get.tree(rfsrc.anonymous(mpg ~ ., mtcars), 10))
## -----
## classification
## ------
print(rfsrc.anonymous(Species ~ ., iris))
## -----
## survival
## -----
data(veteran, package = "randomForestSRC")
print(rfsrc.anonymous(Surv(time, status) ~ ., data = veteran))
## -----
## competing risks
## -----
data(wihs, package = "randomForestSRC")
print(rfsrc.anonymous(Surv(time, status) ~ ., wihs, ntree = 100))
## -----
## unsupervised forests
## -----
print(rfsrc.anonymous(data = iris))
## ------
## multivariate regression
## -----
print(rfsrc.anonymous(Multivar(mpg, cyl) ~., data = mtcars))
```

rfsrc.fast

```
## -----
## prediction on test data with missing values using pbc data
## cases 1 to 312 have no missing values
## cases 313 to 418 having missing values
## ------
data(pbc, package = "randomForestSRC")
pbc.obj <- rfsrc.anonymous(Surv(days, status) ~ ., pbc)</pre>
print(pbc.obj)
## mean value imputation
print(predict(pbc.obj, pbc[-(1:312),], na.action = "na.impute"))
## random imputation
print(predict(pbc.obj, pbc[-(1:312),], na.action = "na.random"))
## -----
## train/test setting but tricky because factor labels differ over
## training and test data
## ------
# first we convert all x-variables to factors
data(veteran, package = "randomForestSRC")
veteran.factor <- data.frame(lapply(veteran, factor))</pre>
veteran.factor$time <- veteran$time</pre>
veteran.factor$status <- veteran$status</pre>
# split the data into train/test data (25/75)
# the train/test data have the same levels, but different labels
train <- sample(1:nrow(veteran), round(nrow(veteran) * .5))</pre>
summary(veteran.factor[train, ])
summary(veteran.factor[-train, ])
# grow the forest on the training data and predict on the test data
v.grow <- rfsrc.anonymous(Surv(time, status) ~ ., veteran.factor[train, ])</pre>
v.pred <- predict(v.grow, veteran.factor[-train, ])</pre>
print(v.grow)
print(v.pred)
```

rfsrc.fast

Fast Random Forests

Description

Fast approximate random forests using subsampling with forest options set to encourage computational speed. Applies to all families.

rfsrc.fast

Usage

```
rfsrc.fast(formula, data,
    ntree = 500,
    nsplit = 10,
    bootstrap = "by.root",
    sampsize = function(x){min(x * .632, max(150, x ^ (3/4)))},
    samptype = "swor",
    samp = NULL,
    ntime = 50,
    forest = FALSE,
    save.memory = TRUE,
    ...)
```

Arguments

| formula | Model to be fit. If missing, unsupervised splitting is implemented. |
|-------------|---|
| data | Data frame containing the y-outcome and x-variables. |
| ntree | Number of trees. |
| nsplit | Non-negative integer value specifying number of random split points used to split a node (deterministic splitting corresponds to the value zero and can be slower). |
| bootstrap | Bootstrap protocol used in growing a tree. |
| sampsize | Function specifying size of subsampled data. Can also be a number. |
| samptype | Type of bootstrap used. |
| samp | Bootstrap specification when "by.user" is used. |
| ntime | Integer value used for survival to constrain ensemble calculations to a grid of ntime time points. |
| forest | Save key forest values? Turn this on if you want prediction on test data. |
| save.memory | Save memory? Setting this to FALSE stores terminal node quantities used for prediction on test data. This yields rapid prediction but can be memory intensive for big data, especially competing risks and survival models. |
| | Further arguments to be passed to rfsrc. |

Details

Calls rfsrc by choosing options (like subsampling) to encourage computational speeds. This will provide a good approximation but will not be as good as default settings of rfsrc.

Value

An object of class (rfsrc, grow).

Author(s)

Hemant Ishwaran and Udaya B. Kogalur

rfsrc.fast

See Also

rfsrc

Examples

```
## ------
## regression
## load the Iowa housing data
data(housing, package = "randomForestSRC")
## do quick and *dirty* imputation
housing <- impute(SalePrice ~ ., housing,</pre>
       ntree = 50, nimpute = 1, splitrule = "random")
## grow a fast forest
o1 <- rfsrc.fast(SalePrice ~ ., housing)</pre>
o2 <- rfsrc.fast(SalePrice ~ ., housing, nodesize = 1)</pre>
print(o1)
print(o2)
## grow a fast bivariate forest
o3 <- rfsrc.fast(cbind(SalePrice,Overall.Qual) ~ ., housing)</pre>
print(o3)
## classification
## -----
data(wine, package = "randomForestSRC")
wine$quality <- factor(wine$quality)</pre>
o <- rfsrc.fast(quality ~ ., wine)</pre>
print(o)
## -----
## grow fast random survival forests without C-calculation
## use brier score to assess model performance
## compare pure random splitting to logrank splitting
## -----
data(peakV02, package = "randomForestSRC")
f <- as.formula(Surv(ttodead, died)~.)</pre>
o1 <- rfsrc.fast(f, peakV02, perf.type = "none")</pre>
o2 <- rfsrc.fast(f, peakV02, perf.type = "none", splitrule = "random")</pre>
bs1 <- get.brier.survival(o1, cens.model = "km")</pre>
bs2 <- get.brier.survival(o2, cens.model = "km")</pre>
plot(bs2$brier.score, type = "s", col = 2)
lines(bs1$brier.score, type = "s", col = 4)
legend("bottomright", legend = c("random", "logrank"), fill = c(2,4))
## -----
```

```
## competing risks
## -----
data(wihs, package = "randomForestSRC")
o <- rfsrc.fast(Surv(time, status) ~ ., wihs)</pre>
print(o)
## -----
## class imbalanced data using gmean performance
## ------
data(breast, package = "randomForestSRC")
breast <- na.omit(breast)</pre>
f <- as.formula(status ~ .)</pre>
o <- rfsrc.fast(f, breast, perf.type = "gmean")</pre>
print(o)
## -----
## class imbalanced data using random forests quantile-classifer (RFQ)
## fast=TRUE => rfsrc.fast
## see imbalanced function for further details
## ------
data(breast, package = "randomForestSRC")
breast <- na.omit(breast)</pre>
f <- as.formula(status ~ .)</pre>
o <- imbalanced(f, breast, fast = TRUE)</pre>
```

```
print(o)
```

rfsrc.news Show the NEWS file

Description

Show the NEWS file of the **randomForestSRC** package.

Usage

```
rfsrc.news(...)
```

Arguments

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

Value

None.

sidClustering.rfsrc

Author(s)

Hemant Ishwaran and Udaya B. Kogalur

sidClustering.rfsrc sidClustering using SID (Staggered Interaction Data) for Unsupervised Clustering

Description

Clustering of unsupervised data using SID (Mantero and Ishwaran, 2021). Also implements the artificial two-class approach of Breiman (2003).

Usage

```
## S3 method for class 'rfsrc'
sidClustering(data,
    method = "sid",
    k = NULL,
    reduce = TRUE,
    ntree = 500,
    ntree.reduce = function(p, vtry){100 * p / vtry},
    fast = FALSE,
    x.no.sid = NULL,
    use.sid.for.x = TRUE,
    x.only = NULL, y.only = NULL,
    dist.sharpen = TRUE, ...)
```

Arguments

| data | A data frame containing the unsupervised data. |
|--------|--|
| method | Clustering method. Default is "sid", which implements SID clustering using Staggered Interaction Data (Mantero and Ishwaran, 2021). An alternative ap- proach reformulates the problem as a two-class supervised learning task using artificial data, per Breiman (2003) and Shi-Horvath (2006). Mode 1 is specified via "sh", "SH", "sh1", or "SH1"; Mode 2 via "sh2" or "SH2". A third method, "unsupv", uses a plain unsupervised forest where the data act as both features and responses, split using the multivariate rule. This is faster than SID but may be less accurate. |
| k | Requested number of clusters. Can be a single integer or a vector. If a scalar, returns a vector assigning each observation to a cluster. If a vector, returns a matrix with one column per requested value of k, each containing a clustering assignment. |
| reduce | Logical. If TRUE, applies a variable reduction step via holdout VIMP. This is conservative and computationally intensive but has strong false discovery control. Applies only when method = "sid". |

| ntree | Number of trees used in the main SID clustering analysis. |
|---------------|---|
| ntree.reduce | Number of trees used in the holdout VIMP step during variable reduction. See holdout.vimp for details. |
| fast | Logical. If TRUE, uses the fast implementation rfsrc.fast instead of rfsrc. Improves speed at the cost of accuracy. |
| x.no.sid | Variables to exclude from SID transformation. Can be either a separate data frame (not overlapping with data) or a character vector of variable names from data. These variables will be included in the final design matrix without SID processing. Applies only when method = "sid". |
| use.sid.for.x | Logical. If FALSE, reverses the roles of features and responses in the SID pro- cess. Staggered interactions are applied to the outcome rather than to features. This option is slower and generally less effective. Included for legacy compati- bility. Applies only when method = "sid". |
| x.only | Character vector specifying which variables to use as features. Applies only when method = "unsupv". |
| y.only | Character vector specifying which variables to use as multivariate responses. Applies only when method = "unsupv". |
| dist.sharpen | Logical. If TRUE (default), applies Euclidean distance to the forest distance ma- trix to improve clustering ("distance sharpening"). The resulting distance matrix will not be bounded between 0 and 1. Turning this off speeds up computation but may reduce clustering quality. Applies only when method = "sid" or "unsupv". |
| | Additional arguments passed to rfsrc to control forest construction. |

Details

Given an unsupervised dataset, random forests is used to compute a distance matrix measuring dissimilarity between all pairs of observations. By default, hierarchical clustering is applied to this distance matrix, although users may apply any other clustering algorithm. See the examples below for alternative workflows.

The default method, method = "sid", implements SID clustering (sidClustering). The algorithm begins by enhancing the original feature space using Staggered Interaction Data (SID). This transformation creates:

- SID main features: shifted and staggered versions of the original features that are made strictly positive and mutually non-overlapping in range;
- SID interaction features: pairwise multiplicative interactions formed between all SID main features.

A multivariate random forest is trained to predict SID main features using the SID interaction features as predictors. The rationale is that if a feature is informative for distinguishing clusters, it will exhibit systematic variation across the data space. Because each interaction feature is uniquely defined by the features it is formed from, node splits on interaction terms are able to capture and separate such variation, thus effectively identifying the clusters. See Mantero and Ishwaran (2021) for further details.

Since SID includes all pairwise interactions, the dimensionality of the feature space grows quadratically with the number of original variables (or worse when factor variables are present). As such, the reduction step using holdout variable importance (VIMP) is strongly recommended (enabled by default). This step can be disabled using reduce = FALSE, but only when the original feature space is of manageable size.

A second approach, proposed by Breiman (2003) and refined by Shi and Horvath (2006), transforms the unsupervised task into a two-class supervised classification problem. The first class consists of the original data, while the second class is generated artificially. The goal is to separate real data from synthetic data. A proximity matrix is constructed from this supervised model, and the proximity values for the original class are extracted and converted into a distance matrix (distance = 1 - proximity) for clustering.

Artificial data can be generated using two modes:

- mode 1 (default): draws random values from the empirical distribution of each feature;
- mode 2: draws uniformly between the observed minimum and maximum of each feature.

This method is invoked by setting method = "sh", "sh1", or "sh2". Mantero and Ishwaran (2021) found that while this approach works in certain settings, it can fail when clusters exist in lowerdimensional subspaces (e.g., when defined by interactions or involving both factors and continuous variables). Among the two modes, mode 1 is generally more robust.

The third method, method = "unsupv", trains a multivariate forest using the data both as predictors and as responses. The multivariate splitting rule is applied at each node. This method is fast and simple but may be less accurate compared to SID clustering.

The package includes a helper function sid.perf.metric for evaluating clustering performance using a normalized score; smaller values indicate better performance. See Mantero and Ishwaran (2021) for theoretical background and empirical benchmarking.

Value

A list with the following components:

| clustering | A vector or matrix assigning each observation to a cluster. If multiple values of k were specified, this is a matrix with one column per clustering solution. |
|------------|---|
| rf | The trained random forest object used in the clustering procedure. This is typi- cally a multivariate forest (for method = "sid" or "unsupv") or a classification forest (for Breiman-style methods). |
| dist | The distance matrix computed from the forest. Used for clustering. For method = "sid", this is based on the forest dissimilarity; for Breiman/SH methods, this is one minus the proximity matrix. |
| sid | The SID-transformed data used in the clustering (applies only to method = "sid"). Provided as a list with separate components for the staggered features and their interactions, corresponding to outcomes and predictors in the multivariate forest. |

Author(s)

Hemant Ishwaran and Udaya B. Kogalur

References

Breiman, L. (2003). *Manual on setting up, using and understanding random forest, V4.0.* University of California Berkeley, Statistics Department, Berkeley.

Mantero A. and Ishwaran H. (2021). Unsupervised random forests. *Statistical Analysis and Data Mining*, 14(2):144-167.

Shi, T. and Horvath, S. (2006). Unsupervised learning with random forest predictors. *Journal of Computational and Graphical Statistics*, 15(1):118-138.

See Also

rfsrc, rfsrc.fast

Examples

```
## -----
## mtcars example
## ______
## default SID method
o1 <- sidClustering(mtcars)</pre>
print(split(mtcars, o1$cl[, 10]))
## using artifical class approach
o1.sh <- sidClustering(mtcars, method = "sh")</pre>
print(split(mtcars, o1.sh$cl[, 10]))
## -----
## glass data set
## ------
if (library("mlbench", logical.return = TRUE)) {
 ## this is a supervised problem, so we first strip the class label
 data(Glass)
 glass <- Glass
 y <- Glass$Type
 glass$Type <- NULL
 ## default SID call
 o2 <- sidClustering(glass, k = 6)
 print(table(y, o2$cl))
 print(sid.perf.metric(y, o2$cl))
 ## compare with Shi-Horvath mode 1
 o2.sh <- sidClustering(glass, method = "sh1", k = 6)
 print(table(y, o2.sh$cl))
 print(sid.perf.metric(y, o2.sh$cl))
 ## plain-vanilla unsupervised analysis
 o2.un <- sidClustering(glass, method = "unsupv", k = 6)
```

```
print(table(y, o2.un$cl))
 print(sid.perf.metric(y, o2.un$cl))
}
## -----
## vowel data set
## -----
if (library("mlbench", logical.return = TRUE) &&
   library("cluster", logical.return = TRUE)) {
 ## strip the class label
 data(Vowel)
 vowel <- Vowel
 y <- Vowel$Class</pre>
 vowel$Class <- NULL</pre>
 ## SID
 o3 <- sidClustering(vowel, k = 11)
 print(table(y, o3$cl))
 print(sid.perf.metric(y, o3$cl))
 ## compare to Shi-Horvath which performs poorly in
 ## mixed variable settings
 o3.sh <- sidClustering(vowel, method = "sh1", k = 11)
 print(table(y, o3.sh$cl))
 print(sid.perf.metric(y, o3.sh$cl))
 ## Shi-Horvath improves with PAM clustering
 ## but still not as good as SID
 o3.sh.pam <- pam(o3.sh$dist, k = 11)$clustering
 print(table(y, o3.sh.pam))
 print(sid.perf.metric(y, o3.sh.pam))
 ## plain-vanilla unsupervised analysis
 o3.un <- sidClustering(vowel, method = "unsupv", k = 11)
 print(table(y, o3.un$cl))
 print(sid.perf.metric(y, o3.un$cl))
}
## -----
## two-d V-shaped cluster (y=x, y=-x) sitting in 12-dimensions
## illustrates superiority of SID to Breiman/Shi-Horvath
p <- 10
m <- 250
n <- 2 * m
std <- .2
x <- runif(n, 0, 1)</pre>
```

```
noise <- matrix(runif(n * p, 0, 1), n)</pre>
y <- rep(NA, n)
y[1:m] <- x[1:m] + rnorm(m, sd = std)</pre>
y[(m+1):n] <- -x[(m+1):n] + rnorm(m, sd = std)</pre>
vclus <- data.frame(clus = c(rep(1, m), rep(2,m)), x = x, y = y, noise)</pre>
## SID
o4 <- sidClustering(vclus[, -1], k = 2)
print(table(vclus[, 1], o4$cl))
print(sid.perf.metric(vclus[, 1], o4$cl))
## Shi-Horvath
o4.sh <- sidClustering(vclus[, -1], method = "sh1", k = 2)
print(table(vclus[, 1], o4.sh$cl))
print(sid.perf.metric(vclus[, 1], o4.sh$cl))
## plain-vanilla unsupervised analysis
o4.un <- sidClustering(vclus[, -1], method = "unsupv", k = 2)
print(table(vclus[, 1], o4.un$cl))
print(sid.perf.metric(vclus[, 1], o4.un$cl))
## -----
## two-d V-shaped cluster using fast random forests
## -----
o5 <- sidClustering(vclus[, -1], k = 2, fast = TRUE)
```

```
print(table(vclus[, 1], o5$cl))
print(sid.perf.metric(vclus[, 1], o5$cl))
```

stat.split.rfsrc Acquire Split Statistic Information

Description

Extract split statistic information from the forest. The function returns a list of length ntree, in which each element corresponds to a tree. The element [[b]] is itself a vector of length xvar.names identified by its x-variable name. Each element [[b]]\$xvar contains the complete list of splits on xvar with associated identifying information. The information is as follows:

- 1. treeID Tree identifier.
- 2. nodeID Node identifier.
- 3. parmID Variable indentifier.
- 4. contPT Value node was split in the case of a continuous variable.
- 5. mwcpSZ Size of the multi-word complementary pair in the case of a factor split.
- 6. *dpthID* Zero (0) based depth of split.

7. *spltTY* Split type for parent node:

| bit 0 | meaning |
|-------|---|
| | |
| 0 | 0 = both daughters have valid splits |
| 1 | 1 = only the right daughter is terminal |
| 0 | 2 = only the left daughter is terminal |
| 1 | 3 = both daughters are terminal |
| | 0 1 |

- 8. *spltEC* End cut statistic for real valued variables between [0,0.5] that is small when the split is towards the edge and large when the split is towards the middle. Subtracting this value from 0.5 yields the end cut statistic studied in Ishwaran (2014) and is a way to identify ECP behavior (end cut preference behavior).
- 9. *spltST* Split statistic:
 - (a) For objects of class (rfsrc, grow), this is the split statistic that resulted in the variable being choosen for the split.
 - (b) For an object of class (rfsrc, pred) this is the variance of the response within the node for the test data. This value is relevant only for real valued responses. In classification and survival, it is not relevant.

Usage

```
## S3 method for class 'rfsrc'
stat.split(object, ...)
```

Arguments

| object | An object of class (rfsrc, grow), (rfsrc, synthetic) or (rfsrc, predict) |
|--------|--|
| | Further arguments passed to or from other methods. |

Value

Invisibly, a list with the following components:

... ...

Author(s)

Hemant Ishwaran and Udaya B. Kogalur

References

Ishwaran H. (2015). The effect of splitting on random forests. Machine Learning, 99:75-118.

Examples

```
## run a forest, then make a call to stat.split
grow.obj <- rfsrc(mpg ~., data = mtcars, membership=TRUE, statistics=TRUE)
stat.obj <- stat.split(grow.obj)</pre>
## nice wrapper to extract split-statistic for desired variable
## for continuous variables plots ECP data
get.split <- function(splitObj, xvar, inches = 0.1, ...) {</pre>
  which.var <- which(names(split0bj[[1]]) == xvar)</pre>
  ntree <- length(split0bj)</pre>
  stat <- data.frame(do.call(rbind, sapply(1:ntree, function(b) {</pre>
    splitObj[[b]][which.var]})))
  dpth <- stat$dpthID</pre>
  ecp <- 1/2 - stat$spltEC</pre>
  sp <- stat$contPT</pre>
  if (!all(is.na(sp))) {
    fgC <- function(x) {
      as.numeric(as.character(cut(x, breaks = c(-1, 0.2, 0.35, 0.5),
      labels = c(1, 4, 2))))
    }
    symbols(jitter(sp), jitter(dpth), ecp, inches = inches, bg = fgC(ecp),
      xlab = xvar, ylab = "node depth", ...)
    legend("topleft", legend = c("low ecp", "med ecp", "high ecp"),
      fill = c(1, 4, 2))
   }
  invisible(stat)
}
## use get.split to investigate ECP behavior of variables
get.split(stat.obj, "disp")
```

subsample.rfsrc Subsample Forests for VIMP Confidence Intervals

Description

Use subsampling to calculate confidence intervals and standard errors for VIMP (variable importance). Applies to all families.

Usage

```
## S3 method for class 'rfsrc'
subsample(obj,
    B = 100,
    block.size = 1,
    importance,
    subratio = NULL,
    stratify = TRUE,
```

subsample.rfsrc

```
performance = FALSE,
performance.only = FALSE,
joint = FALSE,
xvar.names = NULL,
bootstrap = FALSE,
verbose = TRUE)
```

Arguments

| obj | A forest grow object of class (rfsrc, grow). | |
|------------------|---|--|
| В | Number of subsamples (or bootstrap iterations, if bootstrap = TRUE). | |
| block.size | Number of trees in each block used when calculating VIMP. If VIMP is already included in the original grow object, that setting is used instead. | |
| importance | Type of variable importance (VIMP) to compute. Choices are "anti", "permute", or "random". If not specified, the default importance setting from the original grow call is used (if available). | |
| subratio | Subsample size as a proportion of the original sample size. The default is approximately the inverse square root of the sample size. | |
| stratify | Logical. If TRUE, uses stratified subsampling to preserve class balance. See Details for more information. | |
| performance | Logical. If TRUE, calculates generalization error along with standard error and confidence intervals. | |
| performance.only | | |
| | Logical. If TRUE, only generalization error and its uncertainty are returned; VIMP is not computed. | |
| joint | Logical. If TRUE, joint VIMP is computed for all variables. To calculate joint VIMP for a subset of variables, use xvar.names. | |
| xvar.names | Character vector specifying variables to be used for joint VIMP. If omitted, all variables are included. | |
| bootstrap | Logical. If TRUE, uses the double bootstrap instead of subsampling. This is typically slower but may provide more accurate uncertainty estimates. | |
| verbose | Logical. If TRUE, prints progress updates during computation. | |

Details

This function applies subsampling (or optional double bootstrapping) to a previously trained forest to estimate standard errors and construct confidence intervals for variable importance (VIMP), as described in Ishwaran and Lu (2019). It also supports inference for the out-of-bag (OOB) prediction error via the performance = TRUE option. Joint VIMP for selected or all variables can be obtained using joint and xvar.names.

If the original forest does not include VIMP, it will be computed prior to subsampling. For repeated calls to subsample, it is recommended that VIMP be requested in the original rfsrc call. This not only avoids redundant computation, but also ensures consistency of the importance type (e.g., anti, permute, or random) and related parameters, which may otherwise be unclear. Note that permutation importance is *not* the default for most families.

Subsampled forests are constructed using the same tuning parameters as the original forest. While most settings are automatically recovered, certain advanced configurations (e.g., custom sampling schemes) may not be fully supported.

Both subsampled variance estimates (Politis and Romano, 1994) and delete-\(d\) jackknife variance estimates (Shao and Wu, 1989) are returned. The jackknife estimator tends to produce larger standard errors, offering a conservative bias correction, particularly for signal variables.

By default, stratified subsampling is used for classification, survival, and competing risk families:

- For classification, strata correspond to class labels.
- For survival and competing risks, strata include event type and censoring.

Stratification helps ensure representation of key outcome types and is especially important for small sample sizes. Overriding this behavior is discouraged. Note that stratification is *not* available for multivariate families, and caution should be exercised when subsampling in that context.

The function extract.subsample can be used to retrieve detailed information from the subsample object. By default, returned VIMP values are standardized: for regression families, VIMP is divided by the variance of the response; for other families, no transformation is applied. To obtain raw (unstandardized) values, set standardize = FALSE. For expert users, the option raw = TRUE returns detailed internal output, including VIMP from each individual subsampled forest (constructed on a smaller sample size), which is used internally by plot.subsample.rfsrc to generate confidence intervals.

Printed and plotted outputs also standardize VIMP by default. This behavior can be disabled via standardize. The alpha option controls the confidence level and is preset in wrapper functions but can be adjusted by the user.

Value

A list with the following key components:

| rf | Original forest grow object. |
|----------|---|
| vmp | Variable importance values for grow forest. |
| vmpS | Variable importance subsampled values. |
| subratio | Subratio used. |

Author(s)

Hemant Ishwaran and Udaya B. Kogalur

References

Ishwaran H. and Lu M. (2019). Standard errors and confidence intervals for variable importance in random forest regression, classification, and survival. *Statistics in Medicine*, 38, 558-582.

Politis, D.N. and Romano, J.P. (1994). Large sample confidence regions based on subsamples under minimal assumptions. *The Annals of Statistics*, 22(4):2031-2050.

Shao, J. and Wu, C.J. (1989). A general theory for jackknife variance estimation. *The Annals of Statistics*, 17(3):1176-1197.

subsample.rfsrc

See Also

holdout.vimp.rfsrc plot.subsample.rfsrc, rfsrc, vimp.rfsrc

Examples

```
## ------
## regression
## training the forest
reg.o <- rfsrc(Ozone ~ ., airquality)</pre>
## default subsample call
reg.smp.o <- subsample(reg.o)</pre>
## plot confidence regions
plot.subsample(reg.smp.o)
## summary of results
print(reg.smp.o)
## joint vimp and confidence region for generalization error
reg.smp.o2 <- subsample(reg.o, performance = TRUE,</pre>
         joint = TRUE, xvar.names = c("Day", "Month"))
plot.subsample(reg.smp.o2)
## now try the double bootstrap (slower)
reg.dbs.o <- subsample(reg.o, B = 25, bootstrap = TRUE)</pre>
print(reg.dbs.o)
plot.subsample(reg.dbs.o)
## standard error and confidence region for generalization error only
gerror <- subsample(reg.o, performance.only = TRUE)</pre>
plot.subsample(gerror)
## -----
## classification
## ______
## 3 non-linear, 15 linear, and 5 noise variables
if (library("caret", logical.return = TRUE)) {
 d <- twoClassSim(1000, linearVars = 15, noiseVars = 5)</pre>
 ## VIMP based on (default) misclassification error
 cls.o <- rfsrc(Class ~ ., d)</pre>
 cls.smp.o <- subsample(cls.o, B = 100)</pre>
 plot.subsample(cls.smp.o, cex.axis = .7)
 ## same as above, but with VIMP defined using normalized Brier score
 cls.o2 <- rfsrc(Class ~ ., d, perf.type = "brier")</pre>
 cls.smp.o2 <- subsample(cls.o2, B = 100)</pre>
 plot.subsample(cls.smp.o2, cex.axis = .7)
```

subsample.rfsrc

```
}
## ______
## class-imbalanced data using RFQ classifier with G-mean VIMP
if (library("caret", logical.return = TRUE)) {
 ## experimental settings
 n <- 1000
 q <- 20
 ir <- 6
 f <- as.formula(Class ~ .)</pre>
 ## simulate the data, create minority class data
 d <- twoClassSim(n, linearVars = 15, noiseVars = q)</pre>
 d$Class <- factor(as.numeric(d$Class) - 1)
 idx.0 <- which(d$Class == 0)</pre>
 idx.1 <- sample(which(d$Class == 1), sum(d$Class == 1) / ir , replace = FALSE)</pre>
 d <- d[c(idx.0,idx.1),, drop = FALSE]</pre>
 ## RFQ classifier
 oq <- imbalanced(Class ~ ., d, importance = TRUE, block.size = 10)</pre>
 ## subsample the RFQ-classifier
 smp.oq <- subsample(oq, B = 100)</pre>
 plot.subsample(smp.oq, cex.axis = .7)
}
## -----
## survival
## ------
data(pbc, package = "randomForestSRC")
srv.o <- rfsrc(Surv(days, status) ~ ., pbc)</pre>
srv.smp.o <- subsample(srv.o, B = 100)</pre>
plot(srv.smp.o)
## -----
## competing risks
## target event is death (event = 2)
## ______
if (library("survival", logical.return = TRUE)) {
 data(pbc, package = "survival")
 pbc$id <- NULL</pre>
 cr.o <- rfsrc(Surv(time, status) ~ ., pbc, splitrule = "logrankCR", cause = 2)</pre>
 cr.smp.o <- subsample(cr.o, B = 100)</pre>
 plot.subsample(cr.smp.o, target = 2)
}
## -----
```

synthetic

```
## multivariate
## -----
if (library("mlbench", logical.return = TRUE)) {
 ## simulate the data
 data(BostonHousing)
 bh <- BostonHousing</pre>
 bh$rm <- factor(round(bh$rm))</pre>
 o <- rfsrc(cbind(medv, rm) ~ ., bh)</pre>
 so <- subsample(o)</pre>
 plot.subsample(so)
 plot.subsample(so, m.target = "rm")
 ##generalization error
 gerror <- subsample(o, performance.only = TRUE)</pre>
 plot.subsample(gerror, m.target = "medv")
 plot.subsample(gerror, m.target = "rm")
}
## -----
## largish data example - use rfsrc.fast for fast forests
## ------
if (library("caret", logical.return = TRUE)) {
 ## largish data set
 d <- twoClassSim(1000, linearVars = 15, noiseVars = 5)</pre>
 ## use a subsampled forest with Brier score performance
 ## remember to set forest=TRUE for rfsrc.fast
 o <- rfsrc.fast(Class ~ ., d, ntree = 100,</pre>
          forest = TRUE, perf.type = "brier")
 so <- subsample(o, B = 100)</pre>
 plot.subsample(so, cex.axis = .7)
}
```

synthetic

Synthetic Random Forests

Description

Grows a synthetic random forest (RF) using RF machines as synthetic features. Applies only to regression and classification settings.

Usage

```
## S3 method for class 'rfsrc'
synthetic(formula, data, object, newdata,
    ntree = 1000, mtry = NULL, nodesize = 5, nsplit = 10,
```

```
mtrySeq = NULL, nodesizeSeq = c(1:10,20,30,50,100),
min.node = 3,
fast = TRUE,
use.org.features = TRUE,
na.action = c("na.omit", "na.impute"),
oob = TRUE,
verbose = TRUE,
...)
```

Arguments

| formula | Model to be fit. Must be specified unless object is provided. | |
|-----------------|--|--|
| data | Data frame containing the y-outcome and x-variables. Must be specified unless object is provided. | |
| object | An object of class (rfsrc, synthetic). Used to bypass the fitting step. Not required if formula and data are supplied. | |
| newdata | Optional test data for prediction. If omitted, the training data is used. | |
| ntree | Number of trees used in each RF machine. | |
| mtry | mtry value used in the final synthetic forest. | |
| nodesize | nodesize value used in the final synthetic forest. | |
| nsplit | Number of random splits used in randomized splitting. Increases speed when set to a small positive integer. | |
| mtrySeq | Sequence of mtry values used to train the ensemble of RF machines. If NULL, defaults to ceiling(p / 3), where p is the number of variables. | |
| nodesizeSeq | Sequence of nodesize values used to train the ensemble of RF machines. | |
| min.node | Minimum forest-averaged number of terminal nodes required for an RF machine to be retained as a synthetic feature. | |
| fast | Use rfsrc.fast instead of rfsrc to fit base learners? Improves speed at the cost of accuracy. | |
| use.org.feature | es | |
| | Should original features be included alongside synthetic features in the final synthetic forest? | |
| na.action | Action to be taken on missing data. The default, "na.omit", removes records with any missing values. Set to "na.impute" to pre-impute data using impute.rfsrc. | |
| oob | Preserve out-of-bag (OOB) estimation for error rates and VIMP? Defaults to TRUE. | |
| verbose | Display detailed output of the fitting process? Defaults to FALSE. | |
| | Additional arguments passed to rfsrc for training the synthetic forest. | |

Details

A collection of random forests are trained using different values of nodesize (and optionally mtry). The out-of-bag (OOB) predicted values from these forests are then used as synthetic features (referred to as RF machines) to train a final synthetic random forest. The original features can optionally be included in the final model.

synthetic

This approach is currently implemented for regression and classification settings (both univariate and multivariate).

Synthetic features are generated using OOB predictions to prevent overfitting. To ensure that performance metrics (such as error rates and VIMP) remain valid, the same bootstrap samples are reused across all trees for both the synthetic forest and its constituent RF machines. This behavior is controlled by the oob=TRUE option. Disabling this may yield misleading performance estimates and should be done with caution.

If values for mtrySeq are provided, RF machines are constructed for every combination of nodesizeSeq and mtrySeq.

Value

A list with the following components:

| rfMachines | RF machines used to construct the synthetic features. |
|-------------|---|
| rfSyn | The (grow) synthetic RF built over training data. |
| rfSynPred | The predict synthetic RF built over test data (if available). |
| synthetic | List containing the synthetic features. |
| opt.machine | Optimal machine: RF machine with smallest OOB error rate. |

Author(s)

Hemant Ishwaran and Udaya B. Kogalur

References

Ishwaran H. and Malley J.D. (2014). Synthetic learning machines. BioData Mining, 7:28.

See Also

rfsrc, rfsrc.fast

Examples

```
## -----
## compare synthetic forests to regular forest (classification)
## -----
## rfsrc and synthetic calls
if (library("mlbench", logical.return = TRUE)) {
    ## simulate the data
    ring <- data.frame(mlbench.ringnorm(250, 20))
    ## classification forests
    ringRF <- rfsrc(classes ~., ring)
    ## synthetic forests
    ## 1 = nodesize varied</pre>
```

```
## 2 = nodesize/mtry varied
 ringSyn1 <- synthetic(classes ~., ring)</pre>
 ringSyn2 <- synthetic(classes ~., ring, mtrySeq = c(1, 10, 20))</pre>
 ## test-set performance
 ring.test <- data.frame(mlbench.ringnorm(500, 20))</pre>
 pred.ringRF <- predict(ringRF, newdata = ring.test)</pre>
 pred.ringSyn1 <- synthetic(object = ringSyn1, newdata = ring.test)$rfSynPred</pre>
 pred.ringSyn2 <- synthetic(object = ringSyn2, newdata = ring.test)$rfSynPred</pre>
 print(pred.ringRF)
 print(pred.ringSyn1)
 print(pred.ringSyn2)
}
## -----
## compare synthetic forest to regular forest (regression)
## -----
## simulate the data
n <- 250
ntest <- 1000
N <- n + ntest
d <- 50
std <- 0.1
x <- matrix(runif(N * d, -1, 1), ncol = d)</pre>
y <- 1 * (x[,1] + x[,4]^3 + x[,9] + sin(x[,12]*x[,18]) + rnorm(n, sd = std)>.38)
dat <- data.frame(x = x, y = y)</pre>
test <- (n+1):N
## regression forests
regF <- rfsrc(y ~ ., dat[-test, ], )</pre>
pred.regF <- predict(regF, dat[test, ])</pre>
## synthetic forests using fast rfsrc
synF1 <- synthetic(y ~ ., dat[-test, ], newdata = dat[test, ])</pre>
synF2 <- synthetic(y ~ ., dat[-test, ],</pre>
 newdata = dat[test, ], mtrySeq = c(1, 10, 20, 30, 40, 50))
## standardized MSE performance
mse <- c(tail(pred.regF$err.rate, 1),</pre>
        tail(synF1$rfSynPred$err.rate, 1),
        tail(synF2$rfSynPred$err.rate, 1)) / var(y[-test])
names(mse) <- c("forest", "synthetic1", "synthetic2")</pre>
print(mse)
## -----
## multivariate synthetic forests
## -----
```

mtcars.new <- mtcars</pre>

```
mtcars.new$cyl <- factor(mtcars.new$cyl)
mtcars.new$carb <- factor(mtcars.new$carb, ordered = TRUE)
trn <- sample(1:nrow(mtcars.new), nrow(mtcars.new)/2)
mvSyn <- synthetic(cbind(carb, mpg, cyl) ~., mtcars.new[trn,])
mvSyn.pred <- synthetic(object = mvSyn, newdata = mtcars.new[-trn,])</pre>
```

tune.rfsrc

Tune Random Forest for the optimal mtry and nodesize parameters

Description

Finds the optimal mtry and nodesize tuning parameter for a random forest using out-of-sample error. Applies to all families.

Usage

```
## S3 method for class 'rfsrc'
tune(formula, data,
    mtryStart = ncol(data) / 2,
    nodesizeTry = c(1:9, seq(10, 100, by = 5)), ntreeTry = 100,
    sampsize = function(x){min(x * .632, max(150, x ^ (3/4)))},
    nsplit = 1, stepFactor = 1.25, improve = 1e-3, strikeout = 3, maxIter = 25,
    trace = FALSE, doBest = TRUE, ...)
```

```
## S3 method for class 'rfsrc'
tune.nodesize(formula, data,
    nodesizeTry = c(1:9, seq(10, 150, by = 5)), ntreeTry = 100,
    sampsize = function(x){min(x * .632, max(150, x ^ (4/5)))},
    nsplit = 1, trace = TRUE, ...)
```

Arguments

| formula | A symbolic formula describing the model to be fit. |
|-------------|--|
| data | A data frame containing the response variable and predictor variables. |
| mtryStart | Initial value of mtry used to start the tuning search. |
| nodesizeTry | Vector of nodesize values over which tuning is performed. |
| ntreeTry | Number of trees used during the tuning step. |
| sampsize | Function specifying the size of the subsample. Can also be a numeric value. |
| nsplit | Number of random split points considered when splitting a node. |
| stepFactor | Multiplicative factor used to adjust mtry at each iteration. |
| improve | Minimum relative improvement in out-of-sample error required to continue the search. |

| strikeout | Number of consecutive non-improving steps (negative improvement) allowed before stopping the search. Increase to allow a more exhaustive search. |
|-----------|--|
| maxIter | Maximum number of iterations allowed for the mtry bisection search. |
| trace | If TRUE, prints progress during the search. |
| doBest | If TRUE, fits and returns a forest using the optimal mtry and nodesize. |
| | Additional arguments passed to rfsrc.fast. |

Details

tune returns a matrix with three columns: the first and second columns contain the nodesize and mtry values evaluated during the tuning process, and the third column contains the corresponding out-of-sample error.

The error is standardized. For multivariate forests, it is averaged over the outcomes; for competing risks, it is averaged over the event types.

If doBest = TRUE, the function also returns a forest object fit using the optimal mtry and nodesize values.

All tuning calculations, including the final optimized forest, are performed using the fast forest interface rfsrc.fast, which relies on subsampling. This makes the procedure computationally efficient but approximate. Users seeking more accurate tuning results may wish to adjust parameters such as:

- Increasing sampsize, which controls the size of the subsample used for tuning.
- Increasing ntreeTry, which defaults to 100 for speed.

It is also helpful to visualize the out-of-sample error surface as a function of mtry and nodesize using a contour plot (see example below) to identify regions of low error.

The function tune.nodesize performs a simplified search by optimizing only over nodesize.

Author(s)

Hemant Ishwaran and Udaya B. Kogalur

See Also

rfsrc.fast

Examples

```
## -----
## White wine classification example
## -----
## load the data
data(wine, package = "randomForestSRC")
wine$quality <- factor(wine$quality)
## set the sample size manually
o <- tune(quality ~ ., wine, sampsize = 100)</pre>
```

```
## here is the optimized forest
print(o$rf)
## visualize the nodesize/mtry OOB surface
if (library("interp", logical.return = TRUE)) {
 ## nice little wrapper for plotting results
 plot.tune <- function(o, linear = TRUE) {</pre>
   x <- o$results[,1]</pre>
   y <- o$results[,2]</pre>
   z <- o$results[,3]</pre>
   so <- interp(x=x, y=y, z=z, linear = linear)</pre>
   idx <- which.min(z)</pre>
   x0 <- x[idx]
   y0 <- y[idx]
   filled.contour(x = so$x,
                y = so$y,
                z = so$z,
                xlim = range(sox, finite = TRUE) + c(-2, 2),
                ylim = range(soy, finite = TRUE) + c(-2, 2),
                color.palette =
                  colorRampPalette(c("yellow", "red")),
                 xlab = "nodesize",
                 ylab = "mtry",
                main = "error rate for nodesize and mtry",
                 key.title = title(main = "00B error", cex.main = 1),
                 plot.axes = {axis(1);axis(2);points(x0,y0,pch="x",cex=1,font=2);
                            points(x,y,pch=16,cex=.25)})
 }
 ## plot the surface
 plot.tune(o)
}
## -----
## tuning for class imbalanced data problem
## - see imbalanced function for details
## - use rfq and perf.type = "gmean"
## -----
data(breast, package = "randomForestSRC")
breast <- na.omit(breast)</pre>
o <- tune(status ~ ., data = breast, rfq = TRUE, perf.type = "gmean")</pre>
print(o)
## ------
## tune nodesize for competing risk - wihs data
## -----
data(wihs, package = "randomForestSRC")
```

```
plot(tune.nodesize(Surv(time, status) ~ ., wihs, trace = TRUE)$err)
```

vdv

van de Vijver Microarray Breast Cancer

Description

Gene expression profiling for predicting clinical outcome of breast cancer (van't Veer et al., 2002). Microarray breast cancer data set of 4707 expression values on 78 patients with survival information.

References

van't Veer L.J. et al. (2002). Gene expression profiling predicts clinical outcome of breast cancer. *Nature*, **12**, 530–536.

Examples

data(vdv, package = "randomForestSRC")

veteran

Veteran's Administration Lung Cancer Trial

Description

Randomized trial of two treatment regimens for lung cancer. This is a standard survival analysis data set.

Source

Kalbfleisch and Prentice, The Statistical Analysis of Failure Time Data.

References

Kalbfleisch J. and Prentice R, (1980) The Statistical Analysis of Failure Time Data. New York: Wiley.

Examples

data(veteran, package = "randomForestSRC")

vimp.rfsrc

Description

Calculate variable importance (VIMP) for a single variable or group of variables for training or test data.

Usage

```
## S3 method for class 'rfsrc'
vimp(object, xvar.names,
    importance = c("anti", "permute", "random"), block.size = 10,
    joint = FALSE, seed = NULL, do.trace = FALSE, ...)
```

Arguments

| object | An object of class (rfsrc, grow) or (rfsrc, forest). The original rfsrc call must have been made with forest = TRUE. |
|------------|--|
| xvar.names | Character vector of x-variable names to be evaluated. If not specified, all variables are used. |
| importance | Type of variable importance (VIMP) to compute. |
| block.size | Integer specifying the number of trees per block used for VIMP calculation. Balances between ensemble-level and tree-level estimates. |
| joint | Logical indicating whether to compute joint VIMP for the specified variables. |
| seed | Negative integer used to set the random number generator seed. |
| do.trace | Number of seconds between printed progress updates. |
| | Additional arguments passed to or from other methods. |

Details

Using a previously trained forest, this function calculates variable importance (VIMP) for the specified variables in xvar.names. By default, VIMP is computed using the original training data, but the user may supply a new test set via the newdata argument. See rfsrc for further details on how VIMP is computed.

If joint = TRUE, joint VIMP is returned. This is defined as the importance of a group of variables when the entire group is perturbed simultaneously.

Setting csv = TRUE returns case-specific VIMP, which provides VIMP estimates at the individual observation level. This applies to all families except survival. See examples below.

Value

An object of class (rfsrc, predict) containing importance values.

Author(s)

Hemant Ishwaran and Udaya B. Kogalur

References

Ishwaran H. (2007). Variable importance in binary regression trees and forests, *Electronic J. Statist.*, 1:519-537.

See Also

holdout.vimp.rfsrc, rfsrc

Examples

```
## ------
## classification example
## showcase different vimp
iris.obj <- rfsrc(Species ~ ., data = iris)</pre>
## anti vimp (default)
print(vimp(iris.obj)$importance)
## anti vimp using brier prediction error
print(vimp(iris.obj, perf.type = "brier")$importance)
## permutation vimp
print(vimp(iris.obj, importance = "permute")$importance)
## random daughter vimp
print(vimp(iris.obj, importance = "random")$importance)
## joint anti vimp
print(vimp(iris.obj, joint = TRUE)$importance)
## paired anti vimp
print(vimp(iris.obj, c("Petal.Length", "Petal.Width"), joint = TRUE)$importance)
print(vimp(iris.obj, c("Sepal.Length", "Petal.Width"), joint = TRUE)$importance)
## -----
## survival example
## anti versus permute VIMP with different block sizes
data(pbc, package = "randomForestSRC")
pbc.obj <- rfsrc(Surv(days, status) ~ ., pbc)</pre>
print(vimp(pbc.obj)$importance)
print(vimp(pbc.obj, block.size=1)$importance)
print(vimp(pbc.obj, importance="permute")$importance)
print(vimp(pbc.obj, importance="permute", block.size=1)$importance)
```

vimp.rfsrc

```
## -----
## imbalanced classification example
## see the imbalanced function for more details
data(breast, package = "randomForestSRC")
breast <- na.omit(breast)</pre>
f <- as.formula(status ~ .)</pre>
o <- rfsrc(f, breast, ntree = 2000)</pre>
## permutation vimp
print(100 * vimp(o, importance = "permute")$importance)
## anti vimp using gmean performance
print(100 * vimp(o, perf.type = "gmean")$importance[, 1])
## ------
## regression example
## -----
airq.obj <- rfsrc(Ozone ~ ., airquality)</pre>
print(vimp(airq.obj))
## ------
## regression example where vimp is calculated on test data
## -----
set.seed(100080)
train <- sample(1:nrow(airquality), size = 80)</pre>
airq.obj <- rfsrc(Ozone~., airquality[train, ])</pre>
## training data vimp
print(airq.obj$importance)
print(vimp(airq.obj)$importance)
## test data vimp
print(vimp(airq.obj, newdata = airquality[-train, ])$importance)
## ------
## case-specific vimp
## returns VIMP for each case
## ______
o <- rfsrc(mpg~., mtcars)</pre>
v <- vimp(o, csv = TRUE)
csvimp <- get.mv.csvimp(v, standardize=TRUE)</pre>
print(csvimp)
## -----
## case-specific joint vimp
## returns joint VIMP for each case
## -----
```

```
o <- rfsrc(mpg~., mtcars)
v <- vimp(o, joint = TRUE, csv = TRUE)
csvimp <- get.mv.csvimp(v, standardize=TRUE)
print(csvimp)
## ------
## case-specific joint vimp for multivariate regression
## returns joint VIMP for each case, for each outcome
## ------
o <- rfsrc(Multivar(mpg, cyl) ~., data = mtcars)
v <- vimp(o, joint = TRUE, csv = TRUE)
csvimp <- get.mv.csvimp(v, standardize=TRUE)
print(csvimp)
```

wihs

Women's Interagency HIV Study (WIHS)

Description

Competing risk data set involving AIDS in women.

Format

A data frame containing:

| time | time to event |
|----------|--|
| status | censoring status: 0=censoring, 1=HAART initiation, 2=AIDS/Death before HAART |
| ageatfda | age in years at time of FDA approval of first protease inhibitor |
| idu | history of IDU: 0=no history, 1=history |
| black | race: 0=not African-American; 1=African-American |
| cd4nadir | CD4 count (per 100 cells/ul) |

Source

Study included 1164 women enrolled in WIHS, who were alive, infected with HIV, and free of clinical AIDS on December, 1995, when the first protease inhibitor (saquinavir mesylate) was approved by the Federal Drug Administration. Women were followed until the first of the following occurred: treatment initiation, AIDS diagnosis, death, or administrative censoring (September, 2006). Variables included history of injection drug use at WIHS enrollment, whether an individual was African American, age, and CD4 nadir prior to baseline.

References

Bacon M.C, von Wyl V., Alden C., et al. (2005). The Women's Interagency HIV Study: an observational cohort brings clinical sciences to the bench, *Clin Diagn Lab Immunol*, 12(9):1013-1019.

wine

Examples

```
data(wihs, package = "randomForestSRC")
wihs.obj <- rfsrc(Surv(time, status) ~ ., wihs, nsplit = 3, ntree = 100)</pre>
```

wine

White Wine Quality Data

Description

The inputs include objective tests (e.g. PH values) and the output is based on sensory data (median of at least 3 evaluations made by wine experts) of white wine. Each expert graded the wine quality between 0 (very bad) and 10 (very excellent).

References

Cortez, P., Cerdeira, A., Almeida, F., Matos T. and Reis, J. (2009). Modeling wine preferences by data mining from physicochemical properties. In *Decision Support Systems*, Elsevier, 47(4):547-553.

Examples

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## load wine and convert to a multiclass problem
data(wine, package = "randomForestSRC")
wine$quality <- factor(wine$quality)</pre>
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