

Package ‘ggRandomForests’

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Description Graphic elements for exploring Random Forests using the 'randomForest' or 'randomForestSRC' package for survival, regression and classification forests and 'ggplot2' package plotting.

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ggRandomForests-package

ggRandomForests: Visually Exploring Random Forests

Description

ggRandomForests is a utility package for randomForestSRC (Ishwaran et.al. 2014, 2008, 2007) for survival, regression and classification forests and uses the ggplot2 (Wickham 2009) package for plotting results. ggRandomForests is structured to extract data objects from the random forest and provides S3 functions for printing and plotting these objects.

The randomForestSRC package provides a unified treatment of Breiman's (2001) random forests for a variety of data settings. Regression and classification forests are grown when the response is numeric or categorical (factor) while survival and competing risk forests (Ishwaran et al. 2008, 2012) are grown for right-censored survival data.

Many of the figures created by the `ggRandomForests` package are also available directly from within the `randomForestSRC` package. However, `ggRandomForests` offers the following advantages:

- Separation of data and figures: `ggRandomForest` contains functions that operate on either the `rfsrc` forest object directly, or on the output from `randomForestSRC` post processing functions (i.e. `plot.variable`, `var.select`, `find.interaction`) to generate intermediate `ggRandomForests` data objects. S3 functions are provided to further process these objects and plot results using the `ggplot2` graphics package. Alternatively, users can use these data objects for additional custom plotting or analysis operations.
- Each data object/figure is a single, self contained object. This allows simple modification and manipulation of the data or `ggplot2` objects to meet users specific needs and requirements.
- The use of `ggplot2` for plotting. We chose to use the `ggplot2` package for our figures to allow users flexibility in modifying the figures to their liking. Each S3 plot function returns either a single `ggplot2` object, or a list of `ggplot2` objects, allowing users to use additional `ggplot2` functions or themes to modify and customize the figures to their liking.

The `ggRandomForests` package contains the following data functions:

- `gg_rfsrc`: `randomForest[SRC]` predictions.
- `gg_error`: `randomForest[SRC]` convergence rate based on the OOB error rate.
- `gg_roc`: ROC curves for `randomForest` classification models.
- `gg_vimp`: Variable Importance ranking for variable selection.
- `gg_minimal_depth`: Minimal Depth ranking for variable selection (Ishwaran et.al. 2010).
- `gg_minimal_vimp`: Comparing Minimal Depth and VIMP rankings for variable selection.
- `gg_interaction`: Minimal Depth interaction detection (Ishwaran et.al. 2010)
- `gg_variable`: Marginal variable dependence.
- `gg_partial`: Partial (risk adjusted) variable dependence.
- `gg_partial_coplot`: Partial variable conditional dependence (computationally expensive).
- `gg_survival`: Kaplan-Meier/Nelson-Aalen hazard analysis.

Each of these data functions has an associated S3 plot function that returns `ggplot2` objects, either individually or as a list, which can be further customized using standard `ggplot2` commands.

References

- Breiman, L. (2001). Random forests, *Machine Learning*, 45:5-32.
- Ishwaran H. and Kogalur U.B. (2014). Random Forests for Survival, Regression and Classification (RF-SRC), R package version 1.5.5.12.
- Ishwaran H. and Kogalur U.B. (2007). Random survival forests for R. *R News* 7(2), 25–31.
- Ishwaran H., Kogalur U.B., Blackstone E.H. and Lauer M.S. (2008). Random survival forests. *Ann. Appl. Statist.* 2(3), 841–860.
- Ishwaran, H., U. B. Kogalur, E. Z. Gorodeski, A. J. Minn, and M. S. Lauer (2010). High-dimensional variable selection for survival data. *J. Amer. Statist. Assoc.* 105, 205-217.
- Ishwaran, H. (2007). Variable importance in binary regression trees and forests. *Electronic J. Statist.*, 1, 519-537.
- Wickham, H. *ggplot2: elegant graphics for data analysis*. Springer New York, 2009.

`calc_auc`*Area Under the ROC Curve calculator*

Description

Area Under the ROC Curve calculator

Usage

```
calc_auc(x)
```

Arguments

x [gg_roc](#) object

Details

`calc_auc` uses the trapezoidal rule to calculate the area under the ROC curve. This is a helper function for the [gg_roc](#) functions.

Value

AUC. 50% is random guessing, higher is better.

See Also

[calc_roc](#) [gg_roc](#)

[plot.gg_roc](#)

Examples

```
##  
## Taken from the gg_roc example  
rfsrc_iris <- rfsrc(Species ~ ., data = iris)  
  
## Not run:  
gg_dta <- gg_roc(rfsrc_iris, which_outcome=1)  
  
calc_auc(gg_dta)  
  
## End(Not run)  
  
gg_dta <- gg_roc(rfsrc_iris, which_outcome=2)  
  
calc_auc(gg_dta)  
  
## randomForest tests  
rf_iris <- randomForest::randomForest(Species ~ ., data = iris)  
gg_dta <- gg_roc(rfsrc_iris, which_outcome=2)
```

```
calc_auc(gg_dta)
```

calc_roc.rfsrc	<i>Receiver Operator Characteristic calculator</i>
----------------	--

Description

Receiver Operator Characteristic calculator

Usage

```
## S3 method for class 'rfsrc'  
calc_roc(object, dta, which_outcome = "all", oob = TRUE, ...)
```

Arguments

object	rfsrc or predict.rfsrc object containing predicted response
dta	True response variable
which_outcome	If defined, only show ROC for this response.
oob	Use OOB estimates, the normal validation method (TRUE)
...	extra arguments passed to helper functions

Details

For a randomForestSRC prediction and the actual response value, calculate the specificity (1-False Positive Rate) and sensitivity (True Positive Rate) of a predictor.

This is a helper function for the [gg_roc](#) functions, and not intended for use by the end user.

Value

A [gg_roc](#) object

See Also

[calc_auc](#) [gg_roc](#)

[plot.gg_roc](#)

Examples

```
## Taken from the gg_roc example
rfsrc_iris <- rfsrc(Species ~ ., data = iris)

gg_dta <- calc_roc(rfsrc_iris, rfsrc_iris$yvar,
  which_outcome=1, oob=TRUE)
gg_dta <- calc_roc(rfsrc_iris, rfsrc_iris$yvar,
  which_outcome=1, oob=FALSE)

rf_iris <- randomForest(Species ~ ., data = iris)
gg_dta <- calc_roc(rf_iris, rf_iris$yvar,
  which_outcome=1)
gg_dta <- calc_roc(rf_iris, rf_iris$yvar,
  which_outcome=2)
```

combine.gg_partial *combine two gg_partial objects*

Description

The `combine.gg_partial` function assumes the two `gg_partial` objects were generated from the same `rfsrc` object. So, the function joins along the `gg_partial` list item names (one per partial plot variable). Further, we combine the two `gg_partial` objects along the group variable.

Hence, to join three `gg_partial` objects together (i.e. for three different time points from a survival random forest) would require two `combine.gg_partial` calls: One to join the first two `gg_partial` object, and one to append the third `gg_partial` object to the output from the first call. The second call will append a single `lbls` label to the `gg_partial` object.

Usage

```
combine.gg_partial(x, y, lbls, ...)
```

Arguments

<code>x</code>	<code>gg_partial</code> object
<code>y</code>	<code>gg_partial</code> object
<code>lbls</code>	vector of 2 strings to label the combined data.
<code>...</code>	not used

Value

`gg_partial` or `gg_partial_list` based on class of `x` and `y`.

Examples

```

## Not run:
# We need to create this dataset
data(pbc, package = "randomForestSRC",)
# For whatever reason, the age variable is in days... makes no sense to me
for (ind in seq_len(dim(pbc)[2])) {
  if (!is.factor(pbc[, ind])) {
    if (length(unique(pbc[which(!is.na(pbc[, ind])), ind])) <= 2) {
      if (sum(range(pbc[, ind], na.rm = TRUE) == c(0, 1)) == 2) {
        pbc[, ind] <- as.logical(pbc[, ind])
      }
    }
  } else {
    if (length(unique(pbc[which(!is.na(pbc[, ind])), ind])) <= 2) {
      if (sum(sort(unique(pbc[, ind])) == c(0, 1)) == 2) {
        pbc[, ind] <- as.logical(pbc[, ind])
      }
      if (sum(sort(unique(pbc[, ind])) == c(FALSE, TRUE)) == 2) {
        pbc[, ind] <- as.logical(pbc[, ind])
      }
    }
  }
  if (!is.logical(pbc[, ind]) &
      length(unique(pbc[which(!is.na(pbc[, ind])), ind])) <= 5) {
    pbc[, ind] <- factor(pbc[, ind])
  }
}
#Convert age to years
pbc$age <- pbc$age / 364.24

pbc$years <- pbc$days / 364.24
pbc <- pbc[, -which(colnames(pbc) == "days")]
pbc$treatment <- as.numeric(pbc$treatment)
pbc$treatment[which(pbc$treatment == 1)] <- "DPCA"
pbc$treatment[which(pbc$treatment == 2)] <- "placebo"
pbc$treatment <- factor(pbc$treatment)
dta_train <- pbc[-which(is.na(pbc$treatment)), ]
# Create a test set from the remaining patients
pbc_test <- pbc[which(is.na(pbc$treatment)), ]

#=====
# build the forest:
rfsrc_pbc <- randomForestSRC::rfsrc(
  Surv(years, status) ~ .,
  dta_train,
  nsplit = 10,
  na.action = "na.impute",
  forest = TRUE,
  importance = TRUE,
  save.memory = TRUE
)
xvar <- c("bili", "albumin", "copper", "prothrombin", "age")

```

```

xvar_cat <- c("edema")
xvar <- c(xvar, xvar_cat)

time_index <- c(which(rfsrc_pbc$time.interest > 1)[1] - 1,
               which(rfsrc_pbc$time.interest > 3)[1] - 1,
               which(rfsrc_pbc$time.interest > 5)[1] - 1)

partial_pbc <- mclapply(rfsrc_pbc$time.interest[time_index],
                       function(tm) {
                         plot.variable(rfsrc_pbc, surv.type = "surv",
                                       time = tm, xvar.names = xvar,
                                       partial = TRUE,
                                       show.plots = FALSE)
                       })

# A list of 2 plot.variable objects
length(partial_pbc)
class(partial_pbc)

class(partial_pbc[[1]])
class(partial_pbc[[2]])

# Create gg_partial objects
ggPrtl.1 <- gg_partial(partial_pbc[[1]])
ggPrtl.2 <- gg_partial(partial_pbc[[2]])

# Combine the objects to get multiple time curves
# along variables on a single figure.
ggpart <- combine.gg_partial(ggPrtl.1, ggPrtl.2,
                             lbls = c("1 year", "3 years"))

# Plot each figure separately
plot(ggpart)

# Get the continuous data for a panel of continuous plots.
ggcont <- ggpart
ggcont$edema <- ggcont$ascites <- ggcont$stage <- NULL
plot(ggcont, panel=TRUE)

# And the categorical for a panel of categorical plots.
nms <- colnames(sapply(ggcont, function(st) {st}))
for(ind in nms) {
  ggpart[[ind]] <- NULL
}
plot(ggpart, panel=TRUE)

## End(Not run)

```


Description

Extract the cumulative (OOB) randomForestSRC error rate as a function of number of trees.

Usage

```
gg_error(object, ...)
```

Arguments

object [rfsrc](#) object.
 ... optional arguments (not used).

Details

The `gg_error` function simply returns the `rfsrc$err.rate` object as a data.frame, and assigns the class for connecting to the S3 `plot.gg_error` function.

Value

`gg_error` data.frame with one column indicating the tree number, and the remaining columns from the `rfsrc$err.rate` return value.

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.
 Ishwaran H. and Kogalur U.B. (2013). Random Forests for Survival, Regression and Classification (RF-SRC), R package version 1.4.

See Also

[plot.gg_error](#) `rfsrc` `plot.rfsrc`

Examples

```
## Examples from RFSRC package...
## -----
## classification example
## -----
## ----- iris data
## You can build a randomForest
rfsrc_iris <- rfsrc(Species ~ ., data = iris, tree.err = TRUE)

# Get a data.frame containing error rates
gg_dta <- gg_error(rfsrc_iris)

# Plot the gg_error object
plot(gg_dta)

## RandomForest example
```

```

rf_iris <- randomForest::randomForest(Species ~ ., data = iris,
                                     tree.err = TRUE, )
gg_dta <- gg_error(rf_iris)
plot(gg_dta)

gg_dta <- gg_error(rf_iris, training=TRUE)
plot(gg_dta)
## -----
## Regression example
## -----
## Not run:
## ----- airq data
rfsrc_airq <- rfsrc(Ozone ~ ., data = airquality,
                  na.action = "na.impute", tree.err = TRUE, )

# Get a data.frame containing error rates
gg_dta <- gg_error(rfsrc_airq)

# Plot the gg_error object
plot(gg_dta)

## End(Not run)

## ----- Boston data
data(Boston, package = "MASS")
Boston$chas <- as.logical(Boston$chas)
rfsrc_boston <- rfsrc(medv ~ .,
                    data = Boston,
                    forest = TRUE,
                    importance = TRUE,
                    tree.err = TRUE,
                    save.memory = TRUE)

# Get a data.frame containing error rates
gg_dta<- gg_error(rfsrc_boston)

# Plot the gg_error object
plot(gg_dta)

## Not run:
## ----- mtcars data
rfsrc_mtcars <- rfsrc(mpg ~ ., data = mtcars, tree.err = TRUE)
# Get a data.frame containing error rates
gg_dta<- gg_error(rfsrc_mtcars)

# Plot the gg_error object
plot(gg_dta)

## End(Not run)

## -----
## Survival example
## -----

```

```

## Not run:
## ----- veteran data
## randomized trial of two treatment regimens for lung cancer
data(veteran, package = "randomForestSRC")
rfsrc_veteran <- rfsrc(Surv(time, status) ~ ., data = veteran,
                      tree.err = TRUE)

gg_dta <- gg_error(rfsrc_veteran)
plot(gg_dta)

## ----- pbc data
# Load a cached randomForestSRC object
# We need to create this dataset
data(pbc, package = "randomForestSRC",)
# For whatever reason, the age variable is in days... makes no sense to me
for (ind in seq_len(dim(pbc)[2])) {
  if (!is.factor(pbc[, ind])) {
    if (length(unique(pbc[which(!is.na(pbc[, ind])), ind])) <= 2) {
      if (sum(range(pbc[, ind], na.rm = TRUE) == c(0, 1)) == 2) {
        pbc[, ind] <- as.logical(pbc[, ind])
      }
    }
  } else {
    if (length(unique(pbc[which(!is.na(pbc[, ind])), ind])) <= 2) {
      if (sum(sort(unique(pbc[, ind])) == c(0, 1)) == 2) {
        pbc[, ind] <- as.logical(pbc[, ind])
      }
      if (sum(sort(unique(pbc[, ind])) == c(FALSE, TRUE)) == 2) {
        pbc[, ind] <- as.logical(pbc[, ind])
      }
    }
  }
  if (!is.logical(pbc[, ind]) &
      length(unique(pbc[which(!is.na(pbc[, ind])), ind])) <= 5) {
    pbc[, ind] <- factor(pbc[, ind])
  }
}
#Convert age to years
pbc$age <- pbc$age / 364.24

pbc$years <- pbc$days / 364.24
pbc <- pbc[, -which(colnames(pbc) == "days")]
pbc$treatment <- as.numeric(pbc$treatment)
pbc$treatment[which(pbc$treatment == 1)] <- "DPCA"
pbc$treatment[which(pbc$treatment == 2)] <- "placebo"
pbc$treatment <- factor(pbc$treatment)
dta_train <- pbc[-which(is.na(pbc$treatment)), ]
# Create a test set from the remaining patients
pbc_test <- pbc[which(is.na(pbc$treatment)), ]

#####
# build the forest:
rfsrc_pbc <- randomForestSRC::rfsrc(

```

```

    Surv(years, status) ~ .,
    dta_train,
    nsplit = 10,
    na.action = "na.impute",
    tree.err = TRUE,
    forest = TRUE,
    importance = TRUE,
    save.memory = TRUE
)

gg_dta <- gg_error(rfsrc_pbc)
plot(gg_dta)

## End(Not run)

```

gg_interaction	<i>Minimal</i>	<i>Depth</i>	<i>Variable</i>	<i>Interaction</i>	<i>data</i>	<i>object</i>
	(find.interaction).					

Description

Converts the matrix returned from [find.interaction](#) to a `data.frame` and add attributes for S3 identification. If passed a `rfsrc` object, `gg_interaction` first runs the [find.interaction](#) function with all optional arguments.

Usage

```
gg_interaction(object, ...)
```

Arguments

`object` a `rfsrc` object or the output from the [find.interaction](#) function call.
`...` optional extra arguments passed to [find.interaction](#).

Value

`gg_interaction` object

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 high-dimensional data. *Statist. Anal. Data Mining*, 4:115-132.

See Also

[rfsrc.find.interaction.max.subtree.var.select.vimp.plot.gg_interaction](#)

Examples

```
## Examples from randomForestSRC package...
## -----
## find interactions, classification setting
## -----
## ----- iris data
iris.obj <- rfsrc(Species ~., data = iris)
## TODO: VIMP interactions not handled yet...
## randomForestSRC::find.interaction(iris.obj, method = "vimp", nrep = 3)

interaction_iris <- randomForestSRC::find.interaction(iris.obj)
gg_dta <- gg_interaction(interaction_iris)

plot(gg_dta, xvar="Petal.Width")
plot(gg_dta, panel=TRUE)

## -----
## find interactions, regression setting
## -----
## Not run:
## ----- air quality data
airq.obj <- rfsrc(Ozone ~ ., data = airquality)
##
## TODO: VIMP interactions not handled yet...
## randomForestSRC::find.interaction(airq.obj, method = "vimp", nrep = 3)
interaction_airq <- randomForestSRC::find.interaction(airq.obj)

gg_dta <- gg_interaction(interaction_airq)

plot(gg_dta, xvar="Temp")
plot(gg_dta, xvar="Solar.R")

plot(gg_dta, panel=TRUE)

## End(Not run)
## Not run:
## ----- Boston data
data(Boston, package = "MASS")
Boston$chas <- as.logical(Boston$chas)
rfsrc_boston <- rfsrc(medv ~ .,
  data = Boston,
  forest = TRUE,
  importance = TRUE,
  tree.err = TRUE,
  save.memory = TRUE)

interaction_boston <- find.interaction(rfsrc_boston)
```

```

gg_dta <- gg_interaction(interaction_boston)

plot(gg_dta, panel=TRUE)

## End(Not run)
## Not run:
## ----- mtcars data
rfsrc_mtcars <- rfsrc(mpg ~ ., data = mtcars)

interaction_mtcars <- find.interaction(rfsrc_mtcars)

gg_dta <- gg_interaction(interaction_mtcars)

plot(gg_dta, panel=TRUE)

## End(Not run)
## Not run:
## -----
## find interactions, survival setting
## ----- veteran data
## randomized trial of two treatment regimens for lung cancer
data(veteran, package = "randomForestSRC")
rfsrc_veteran <- rfsrc(Surv(time, status) ~ ., data = veteran)

interaction_vet <- find.interaction(rfsrc_veteran)

gg_dta <- gg_interaction(interaction_vet)

plot(gg_dta, panel = True)

## -----
## ----- pbc data
# We need to create this dataset
data(pbc, package = "randomForestSRC",)
# For whatever reason, the age variable is in days... makes no sense to me
for (ind in seq_len(dim(pbc)[2])) {
  if (!is.factor(pbc[, ind])) {
    if (length(unique(pbc[which(!is.na(pbc[, ind])), ind])) <= 2) {
      if (sum(range(pbc[, ind], na.rm = TRUE) == c(0, 1)) == 2) {
        pbc[, ind] <- as.logical(pbc[, ind])
      }
    }
  } else {
    if (length(unique(pbc[which(!is.na(pbc[, ind])), ind])) <= 2) {
      if (sum(sort(unique(pbc[, ind])) == c(0, 1)) == 2) {
        pbc[, ind] <- as.logical(pbc[, ind])
      }
      if (sum(sort(unique(pbc[, ind])) == c(FALSE, TRUE)) == 2) {
        pbc[, ind] <- as.logical(pbc[, ind])
      }
    }
  }
}
if (!is.logical(pbc[, ind]) &

```

```

    length(unique(pbc[which(!is.na(pbc[, ind])), ind])) <= 5) {
      pbc[, ind] <- factor(pbc[, ind])
    }
  }
  # Convert age to years
  pbc$age <- pbc$age / 364.24

  pbc$years <- pbc$days / 364.24
  pbc <- pbc[, -which(colnames(pbc) == "days")]
  pbc$treatment <- as.numeric(pbc$treatment)
  pbc$treatment[which(pbc$treatment == 1)] <- "DPCA"
  pbc$treatment[which(pbc$treatment == 2)] <- "placebo"
  pbc$treatment <- factor(pbc$treatment)
  dta_train <- pbc[-which(is.na(pbc$treatment)), ]
  # Create a test set from the remaining patients
  pbc_test <- pbc[which(is.na(pbc$treatment)), ]

  #=====
  # build the forest:
  rfsrc_pbc <- randomForestSRC::rfsrc(
    Surv(years, status) ~ .,
    dta_train,
    nsplit = 10,
    na.action = "na.impute",
    forest = TRUE,
    importance = TRUE,
    save.memory = TRUE
  )

  interaction_pbc <- find.interaction(rfsrc_pbc, nvar = 9)
  gg_dta <- gg_interaction(interaction_pbc)

  plot(gg_dta, xvar="bili")
  plot(gg_dta, panel=TRUE)

  ## End(Not run)

```

gg_minimal_depth

Minimal depth data object ([randomForestSRC]{var.select})

Description

the [randomForestSRC]{var.select} function implements random forest variable selection using tree minimal depth methodology. The gg_minimal_depth function takes the output from [randomForestSRC]{var.select} and creates a data.frame formatted for the [plot.gg_minimal_depth](#) function.

Usage

```
gg_minimal_depth(object, ...)
```

Arguments

object	A [randomForestSRC]{rfsrc} object, [randomForestSRC]{predict} object or the list from the [randomForestSRC]{var.select.rfsrc} function.
...	optional arguments passed to the [randomForestSRC]{var.select} function if operating on an [randomForestSRC]{rfsrc} object.

Value

gg_minimal_depth object, A modified list of variables from the [randomForestSRC]{var.select} function, ordered by minimal depth rank.

See Also

[randomForestSRC]{var.select}
[plot.gg_minimal_depth](#)

Examples

```
## Examples from RFSRC package...
## -----
## classification example
## -----
## ----- iris data
## You can build a randomForest
rfsrc_iris <- rfsrc(Species ~ ., data = iris)
varsel_iris <- randomForestSRC::var.select(rfsrc_iris)

# Get a data.frame containing minimaldepth measures
gg_dta <- gg_minimal_depth(varsel_iris)

# Plot the gg_minimal_depth object
plot(gg_dta)

## -----
## Regression example
## -----
## Not run:
## ----- air quality data
rfsrc_airq <- rfsrc(Ozone ~ ., data = airquality,
                  na.action = "na.impute")
varsel_airq <- var.select(rfsrc_airq)

# Get a data.frame containing error rates
gg_dta <- gg_minimal_depth(varsel_airq)

# Plot the gg_minimal_depth object
plot(gg_dta)
```



```

## End(Not run)
## Not run:

## ----- Boston data
data(Boston, package = "MASS")
Boston$chas <- as.logical(Boston$chas)
rfsrc_boston <- rfsrc(medv ~ .,
  data = Boston,
  forest = TRUE,
  importance = TRUE,
  tree.err = TRUE,
  save.memory = TRUE)

varsel_boston <- var.select(rfsrc_boston)

# Get a data.frame containing error rates
gg_dta <- gg_minimal_depth(varsel_boston)
print(gg_dta)
plot(gg_dta)

## End(Not run)
## Not run:
## ----- mtcars data
rfsrc_mtcars <- rfsrc(mpg ~ ., data = mtcars)
varsel_mtcars <- var.select(rfsrc_mtcars)

# Get a data.frame containing error rates
plot.gg_minimal_depth(varsel_mtcars)

## End(Not run)

## -----
## Survival example
## -----
## Not run:
## ----- veteran data
## randomized trial of two treatment regimens for lung cancer
data(veteran, package = "randomForestSRC")
rfsrc_veteran <- rfsrc(Surv(time, status) ~ ., data = veteran, ntree = 100)
varsel_veteran <- randomForestSRC::var.select(rfsrc_veteran)

gg_dta <- gg_minimal_depth(varsel_veteran)
plot(gg_dta)

## -----
## ----- pbc data
# We need to create this dataset
data(pbc, package = "randomForestSRC",)
# For whatever reason, the age variable is in days... makes no sense to me
for (ind in seq_len(dim(pbc)[2])) {
  if (!is.factor(pbc[, ind])) {
    if (length(unique(pbc[which(!is.na(pbc[, ind])), ind])) <= 2) {

```

```

    if (sum(range(pbc[, ind], na.rm = TRUE) == c(0, 1)) == 2) {
      pbc[, ind] <- as.logical(pbc[, ind])
    }
  } else {
    if (length(unique(pbc[which(!is.na(pbc[, ind])), ind])) <= 2) {
      if (sum(sort(unique(pbc[, ind])) == c(0, 1)) == 2) {
        pbc[, ind] <- as.logical(pbc[, ind])
      }
      if (sum(sort(unique(pbc[, ind])) == c(FALSE, TRUE)) == 2) {
        pbc[, ind] <- as.logical(pbc[, ind])
      }
    }
  }
}
if (!is.logical(pbc[, ind]) &
    length(unique(pbc[which(!is.na(pbc[, ind])), ind])) <= 5) {
  pbc[, ind] <- factor(pbc[, ind])
}
}
#Convert age to years
pbc$age <- pbc$age / 364.24

pbc$years <- pbc$days / 364.24
pbc <- pbc[, -which(colnames(pbc) == "days")]
pbc$treatment <- as.numeric(pbc$treatment)
pbc$treatment[which(pbc$treatment == 1)] <- "DPCA"
pbc$treatment[which(pbc$treatment == 2)] <- "placebo"
pbc$treatment <- factor(pbc$treatment)
dta_train <- pbc[-which(is.na(pbc$treatment)), ]
# Create a test set from the remaining patients
pbc_test <- pbc[which(is.na(pbc$treatment)), ]

#=====
# build the forest:
rfsrc_pbc <- randomForestSRC::rfsrc(
  Surv(years, status) ~ .,
  dta_train,
  nsplit = 10,
  na.action = "na.impute",
  forest = TRUE,
  importance = TRUE,
  save.memory = TRUE
)

varsel_pbc <- var.select(rfsrc_pbc)

gg_dta <- gg_minimal_depth(varsel_pbc)
plot(gg_dta)

## End(Not run)

```

gg_minimal_vimp	<i>Minimal depth vs VIMP comparison by variable rankings.</i>
-----------------	---

Description

Minimal depth vs VIMP comparison by variable rankings.

Usage

```
gg_minimal_vimp(object, ...)
```

Arguments

object	A <code>rfsrc</code> object, <code>predict.rfsrc</code> object or the list from the <code>var.select.rfsrc</code> function.
...	optional arguments passed to the <code>var.select</code> function if operating on an <code>rfsrc</code> object.

Value

`gg_minimal_vimp` comparison object.

See Also

[plot.gg_minimal_vimp](#) [var.select](#)

Examples

```
## Examples from RFSRC package...
## -----
## classification example
## -----
## ----- iris data
## You can build a randomForest
rfsrc_iris <- rfsrc(Species ~ ., data = iris)
varsel_iris <- randomForestSRC::var.select(rfsrc_iris)

# Get a data.frame containing minimaldepth measures
gg_dta<- gg_minimal_vimp(varsel_iris)

# Plot the gg_minimal_depth object
plot(gg_dta)

## -----
## Regression example
## -----
## Not run:
## ----- air quality data
rfsrc_airq <- rfsrc(Ozone ~ ., data = airquality,
```

```

na.action = "na.impute")
varsel_airq <- randomForestSRC::var.select(rfsrc_airq)

# Get a data.frame containing error rates
gg_dta<- gg_minimal_vimp(varsel_airq)

# Plot the gg_minimal_vimp object
plot(gg_dta)

## End(Not run)
## Not run:
## ----- Boston data
data(Boston, package = "MASS")
Boston$chas <- as.logical(Boston$chas)
rfsrc_boston <- rfsrc(medv ~ .,
  data = Boston,
  forest = TRUE,
  importance = TRUE,
  tree.err = TRUE,
  save.memory = TRUE)

varsel_boston <- var.select(rfsrc_boston)

# Get a data.frame containing error rates
gg_dta<- gg_minimal_vimp(varsel_boston)

# Plot the gg_minimal_vimp object
plot(gg_dta)

## End(Not run)
## Not run:
## ----- mtcars data

rfsrc_mtcars <- rfsrc(mpg ~ ., data = mtcars)
varsel_mtcars <- var.select(rfsrc_mtcars)

# Get a data.frame containing error rates
gg_dta <- gg_minimal_vimp(varsel_mtcars)

# Plot the gg_minimal_vimp object
plot(gg_dta)

## End(Not run)
## -----
## Survival example
## -----
## Not run:
## ----- veteran data
## randomized trial of two treatment regimens for lung cancer
data(veteran, package = "randomForestSRC")
rfsrc_veteran <- rfsrc(Surv(time, status) ~ ., data = veteran,
  ntree = 100)
varsel_veteran <- randomForestSRC::var.select(rfsrc_veteran)

```

```

gg_dta <- gg_minimal_vimp(varsel_veteran)
plot(gg_dta)

## -----
## ----- pbc data
# We need to create this dataset
data(pbc, package = "randomForestSRC")
# For whatever reason, the age variable is in days... makes no sense to me
for (ind in seq_len(dim(pbc)[2])) {
  if (!is.factor(pbc[, ind])) {
    if (length(unique(pbc[which(!is.na(pbc[, ind])), ind])) <= 2) {
      if (sum(range(pbc[, ind], na.rm = TRUE) == c(0, 1)) == 2) {
        pbc[, ind] <- as.logical(pbc[, ind])
      }
    }
  } else {
    if (length(unique(pbc[which(!is.na(pbc[, ind])), ind])) <= 2) {
      if (sum(sort(unique(pbc[, ind])) == c(0, 1)) == 2) {
        pbc[, ind] <- as.logical(pbc[, ind])
      }
      if (sum(sort(unique(pbc[, ind])) == c(FALSE, TRUE)) == 2) {
        pbc[, ind] <- as.logical(pbc[, ind])
      }
    }
  }
  if (!is.logical(pbc[, ind]) &
      length(unique(pbc[which(!is.na(pbc[, ind])), ind])) <= 5) {
    pbc[, ind] <- factor(pbc[, ind])
  }
}
#Convert age to years
pbc$age <- pbc$age / 364.24

pbc$years <- pbc$days / 364.24
pbc <- pbc[, -which(colnames(pbc) == "days")]
pbc$treatment <- as.numeric(pbc$treatment)
pbc$treatment[which(pbc$treatment == 1)] <- "DPCA"
pbc$treatment[which(pbc$treatment == 2)] <- "placebo"
pbc$treatment <- factor(pbc$treatment)
dta_train <- pbc[-which(is.na(pbc$treatment)), ]
# Create a test set from the remaining patients
pbc_test <- pbc[which(is.na(pbc$treatment)), ]

#=====
# build the forest:
rfsrc_pbc <- randomForestSRC::rfsrc(
  Surv(years, status) ~ .,
  dta_train,
  nsplit = 10,
  na.action = "na.impute",
  forest = TRUE,

```

```
importance = TRUE,  
save.memory = TRUE  
)  
  
vare1_pbc <- var.select(rfsrc_pbc)  
  
gg_dta <- gg_minimal_vimp(vare1_pbc)  
plot(gg_dta)  
  
## End(Not run)
```

gg_partial

Partial variable dependence object

Description

The `plot.variable` function returns a list of either marginal variable dependence or partial variable dependence data from a `rfsrc` object. The `gg_partial` function formulates the `plot.variable` output for partial plots (where `partial=TRUE`) into a data object for creation of partial dependence plots using the `plot.gg_partial` function.

Partial variable dependence plots are the risk adjusted estimates of the specified response as a function of a single covariate, possibly subsetted on other covariates.

An option named `argument` can name a column for merging multiple plots together

Usage

```
gg_partial(object, ...)
```

Arguments

<code>object</code>	the partial variable dependence data object from <code>plot.variable</code> function
<code>...</code>	optional arguments

Value

`gg_partial` object. A data.frame or list of data.frames corresponding the variables contained within the `plot.variable` output.

References

Friedman, Jerome H. 2000. "Greedy Function Approximation: A Gradient Boosting Machine." *Annals of Statistics* 29: 1189-1232."

See Also

[plot.gg_partial](#)
[plot.variable](#)

Examples

```

## -----
## classification
## -----
## ----- iris data
## iris "Petal.Width" partial dependence plot
##
rfsrc_iris <- rfsrc(Species ~., data = iris)
partial_iris <- plot.variable(rfsrc_iris,
                             xvar.names = "Petal.Width",
                             partial=TRUE)

gg_dta <- gg_partial(partial_iris)
plot(gg_dta)

## -----
## regression
## -----
## Not run:
## ----- air quality data
## airquality "Wind" partial dependence plot
##
rfsrc_airq <- rfsrc(Ozone ~ ., data = airquality)
partial_airq <- plot.variable(rfsrc_airq,
                              xvar.names = "Wind",
                              partial=TRUE, show.plot=FALSE)

gg_dta <- gg_partial(partial_airq)
plot(gg_dta)

## End(Not run)
## Not run:
## ----- Boston data
data(Boston, package = "MASS")
Boston$chas <- as.logical(Boston$chas)
rfsrc_boston <- rfsrc(medv ~ .,
                     data = Boston,
                     forest = TRUE,
                     importance = TRUE,
                     tree.err = TRUE,
                     save.memory = TRUE)

varsel_boston <- var.select(rfsrc_boston)

partial_boston <- plot.variable(rfsrc_boston,
                                xvar.names = varsel_boston$topvars,
                                sorted = FALSE,
                                partial = TRUE,
                                show.plots = FALSE)
gg_dta <- gg_partial(partial_boston)
plot(gg_dta, panel=TRUE)

```

```

## End(Not run)
## Not run:
## ----- mtcars data
rfsrc_mtcars <- rfsrc(mpg ~ ., data = mtcars)
varel_mtcars <- var.select(rfsrc_mtcars)

partial_mtcars <- plot.variable(rfsrc_mtcars,
  xvar.names = varel_mtcars$topvars,
  sorted = FALSE,
  partial = TRUE,
  show.plots = FALSE)

gg_dta <- gg_partial(partial_mtcars)

gg_dta.cat <- gg_dta
gg_dta.cat[["disp"]] <- gg_dta.cat[["wt"]] <- gg_dta.cat[["hp"]] <- NULL
gg_dta.cat[["drat"]] <- gg_dta.cat[["carb"]] <- gg_dta.cat[["qsec"]] <- NULL

plot(gg_dta.cat, panel=TRUE, notch=TRUE)

gg_dta[["cyl"]] <- gg_dta[["vs"]] <- gg_dta[["am"]] <- NULL
gg_dta[["gear"]] <- NULL
plot(gg_dta, panel=TRUE)

## End(Not run)

## -----
## survival examples
## -----
## Not run:
## ----- veteran data
## survival "age" partial variable dependence plot
##
data(veteran, package = "randomForestSRC")
rfsrc_veteran <- rfsrc(Surv(time,status)~., veteran, nsplit = 10,
  ntree = 100)

varel_rfsrc <- var.select(rfsrc_veteran)

## 30 day partial plot for age
partial_veteran <- plot.variable(rfsrc_veteran, surv.type = "surv",
  partial = TRUE, time=30,
  show.plots=FALSE)

gg_dta <- gg_partial(partial_veteran)
plot(gg_dta, panel=TRUE)

gg_dta.cat <- gg_dta
gg_dta.cat[["celltype"]] <- gg_dta.cat[["trt"]] <- gg_dta.cat[["prior"]] <- NULL
plot(gg_dta, panel=TRUE)

gg_dta.cat[["karno"]] <- gg_dta.cat[["diagtime"]] <-

```



```

    gg_dta.cat[["age"]] <- NULL
    plot(gg_dta.cat, panel=TRUE, notch=TRUE)

    gg_dta <- lapply(partial_veteran, gg_partial)
    gg_dta <- combine.gg_partial(gg_dta[[1]], gg_dta[[2]] )

    plot(gg_dta[["karno"]])
    plot(gg_dta[["celltype"]])

    gg_dta.cat <- gg_dta
    gg_dta[["celltype"]] <- gg_dta[["trt"]] <- gg_dta[["prior"]] <- NULL
    plot(gg_dta, panel=TRUE)

    gg_dta.cat[["karno"]] <- gg_dta.cat[["diagtime"]] <-
      gg_dta.cat[["age"]] <- NULL
    plot(gg_dta.cat, panel=TRUE, notch=TRUE)

    ## -----
    ## ----- pbc data
    # We need to create this dataset
    data(pbc, package = "randomForestSRC",)
    # For whatever reason, the age variable is in days... makes no sense to me
    for (ind in seq_len(dim(pbc)[2])) {
      if (!is.factor(pbc[, ind])) {
        if (length(unique(pbc[which(!is.na(pbc[, ind])), ind])) <= 2) {
          if (sum(range(pbc[, ind], na.rm = TRUE) == c(0, 1)) == 2) {
            pbc[, ind] <- as.logical(pbc[, ind])
          }
        }
      } else {
        if (length(unique(pbc[which(!is.na(pbc[, ind])), ind])) <= 2) {
          if (sum(sort(unique(pbc[, ind])) == c(0, 1)) == 2) {
            pbc[, ind] <- as.logical(pbc[, ind])
          }
          if (sum(sort(unique(pbc[, ind])) == c(FALSE, TRUE)) == 2) {
            pbc[, ind] <- as.logical(pbc[, ind])
          }
        }
      }
    }
    if (!is.logical(pbc[, ind]) &
        length(unique(pbc[which(!is.na(pbc[, ind])), ind])) <= 5) {
      pbc[, ind] <- factor(pbc[, ind])
    }
  }
  #Convert age to years
  pbc$age <- pbc$age / 364.24

  pbc$years <- pbc$days / 364.24
  pbc <- pbc[, -which(colnames(pbc) == "days")]
  pbc$treatment <- as.numeric(pbc$treatment)
  pbc$treatment[which(pbc$treatment == 1)] <- "DPCA"
  pbc$treatment[which(pbc$treatment == 2)] <- "placebo"
  pbc$treatment <- factor(pbc$treatment)

```

```

dta_train <- pbc[-which(is.na(pbc$treatment)), ]
# Create a test set from the remaining patients
pbc_test <- pbc[which(is.na(pbc$treatment)), ]

#=====
# build the forest:
rfsrc_pbc <- randomForestSRC::rfsrc(
  Surv(years, status) ~ .,
  dta_train,
  nsplit = 10,
  na.action = "na.impute",
  forest = TRUE,
  importance = TRUE,
  save.memory = TRUE
)

varsel_pbc <- var.select(rfsrc_pbc)

xvar <- varsel_pbc$topvars

# Convert all partial plots to gg_partial objects
gg_dta <- lapply(partial_pbc, gg_partial)

# Combine the objects to get multiple time curves
# along variables on a single figure.
pbc_ggpart <- combine.gg_partial(gg_dta[[1]], gg_dta[[2]],
                                lbls = c("1 Year", "3 Years"))

summary(pbc_ggpart)
class(pbc_ggpart[["bili"]])

# Plot the highest ranked variable, by name.
#plot(pbc_ggpart[["bili"]])

# Create a temporary holder and remove the stage and edema data
ggpart <- pbc_ggpart
ggpart$edema <- NULL

# Panel plot the remainder.
plot(ggpart, panel = TRUE)

plot(pbc_ggpart[["edema"]])

## End(Not run)

```

Description

Data structures for stratified partial coplots

Usage

```
## S3 method for class 'rfsrc'
gg_partial_coplot(
  object,
  xvar,
  groups,
  surv_type = c("mort", "rel.freq", "surv", "years.lost", "cif", "chf"),
  time,
  show_plots = FALSE,
  ...
)
```

Arguments

object	rfsrc object
xvar	list of partial plot variables
groups	vector of stratification variable.
surv_type	for survival random forests, c("mort", "rel.freq", "surv", "years.lost", "cif", "chf")
time	vector of time points for survival random forests partial plots.
show_plots	boolean passed to plot.variable show.plots argument.
...	extra arguments passed to plot.variable function

Value

gg_partial_coplot object. An subclass of a [gg_partial_list](#) object

Examples

```
## Not run:

## -----
## ----- pbc data
## We need to create this dataset
data(pbc, package = "randomForestSRC")
# For whatever reason, the age variable is in days... makes no sense to me
for (ind in seq_len(dim(pbc)[2])) {
  if (!is.factor(pbc[, ind])) {
    if (length(unique(pbc[which(!is.na(pbc[, ind])), ind])) <= 2) {
      if (sum(range(pbc[, ind], na.rm = TRUE) == c(0, 1)) == 2) {
        pbc[, ind] <- as.logical(pbc[, ind])
      }
    }
  } else {
    if (length(unique(pbc[which(!is.na(pbc[, ind])), ind])) <= 2) {
```



```
# Partial coplot
plot(partial_coplot_pbc) #, se = FALSE)

## End(Not run)
```

```
gg_rfsrc.rfsrc      Predicted response data object
```

Description

Extracts the predicted response values from the `rfsrc` object, and formats data for plotting the response using `plot.gg_rfsrc`.

Usage

```
## S3 method for class 'rfsrc'
gg_rfsrc(object, oob = TRUE, by, ...)
```

Arguments

<code>object</code>	<code>rfsrc</code> object
<code>oob</code>	boolean, should we return the oob prediction , or the full forest prediction.
<code>by</code>	stratifying variable in the training dataset, defaults to NULL
<code>...</code>	extra arguments

Details

`surv_type` ("surv", "chf", "mortality", "hazard") for survival forests
`oob` boolean, should we return the oob prediction , or the full forest prediction.

Value

`gg_rfsrc` object

See Also

[plot.gg_rfsrc](#) [rfsrc](#) [plot.rfsrc](#) [gg_survival](#)

Examples

```
## -----
## classification example
## -----
## ----- iris data
rfsrc_iris <- rfsrc(Species ~ ., data = iris)
gg_dta<- gg_rfsrc(rfsrc_iris)
```

```

plot(gg_dta)

## -----
## Regression example
## -----
## Not run:
## ----- air quality data
rfsrc_airq <- rfsrc(Ozone ~ ., data = airquality, na.action = "na.impute")
gg_dta<- gg_rfsrc(rfsrc_airq)

plot(gg_dta)

## End(Not run)

## ----- Boston data
data(Boston, package = "MASS")
Boston$chas <- as.logical(Boston$chas)
rfsrc_boston <- rfsrc(medv ~ .,
  data = Boston,
  forest = TRUE,
  importance = TRUE,
  tree.err = TRUE,
  save.memory = TRUE)

plot(gg_rfsrc(rfsrc_boston))

### randomForest example
data(Boston, package="MASS")
rf_boston <- randomForest::randomForest(medv ~ ., data = Boston)
plot(gg_rfsrc(rf_boston))

## Not run:
## ----- mtcars data
rfsrc_mtcars <- rfsrc(mpg ~ ., data = mtcars)
gg_dta<- gg_rfsrc(rfsrc_mtcars)

plot(gg_dta)

## End(Not run)
## -----
## Survival example
## -----
## Not run:
## ----- veteran data
## randomized trial of two treatment regimens for lung cancer
data(veteran, package = "randomForestSRC")
rfsrc_veteran <- rfsrc(Surv(time, status) ~ ., data = veteran, ntree = 100)

gg_dta <- gg_rfsrc(rfsrc_veteran)
plot(gg_dta)

gg_dta <- gg_rfsrc(rfsrc_veteran, conf.int=.95)
plot(gg_dta)

```

```

gg_dta <- gg_rfsrc(rfsrc_veteran, by="trt")
plot(gg_dta)

## ----- pbc data
## We don't run this because of bootstrap confidence limits
# We need to create this dataset
data(pbc, package = "randomForestSRC",)
# For whatever reason, the age variable is in days... makes no sense to me
for (ind in seq_len(dim(pbc)[2])) {
  if (!is.factor(pbc[, ind])) {
    if (length(unique(pbc[which(!is.na(pbc[, ind])), ind])) <= 2) {
      if (sum(range(pbc[, ind], na.rm = TRUE) == c(0, 1)) == 2) {
        pbc[, ind] <- as.logical(pbc[, ind])
      }
    }
  } else {
    if (length(unique(pbc[which(!is.na(pbc[, ind])), ind])) <= 2) {
      if (sum(sort(unique(pbc[, ind])) == c(0, 1)) == 2) {
        pbc[, ind] <- as.logical(pbc[, ind])
      }
      if (sum(sort(unique(pbc[, ind])) == c(FALSE, TRUE)) == 2) {
        pbc[, ind] <- as.logical(pbc[, ind])
      }
    }
  }
  if (!is.logical(pbc[, ind]) &
      length(unique(pbc[which(!is.na(pbc[, ind])), ind])) <= 5) {
    pbc[, ind] <- factor(pbc[, ind])
  }
}
#Convert age to years
pbc$age <- pbc$age / 364.24

pbc$years <- pbc$days / 364.24
pbc <- pbc[, -which(colnames(pbc) == "days")]
pbc$treatment <- as.numeric(pbc$treatment)
pbc$treatment[which(pbc$treatment == 1)] <- "DPCA"
pbc$treatment[which(pbc$treatment == 2)] <- "placebo"
pbc$treatment <- factor(pbc$treatment)
dta_train <- pbc[-which(is.na(pbc$treatment)), ]
# Create a test set from the remaining patients
pbc_test <- pbc[which(is.na(pbc$treatment)), ]

#=====
# build the forest:
rfsrc_pbc <- randomForestSRC::rfsrc(
  Surv(years, status) ~ .,
  dta_train,
  nsplit = 10,
  na.action = "na.impute",
  forest = TRUE,

```

```

importance = TRUE,
save.memory = TRUE
)
gg_dta <- gg_rfsrc(rfsrc_pbc)
plot(gg_dta)

gg_dta <- gg_rfsrc(rfsrc_pbc, conf.int=.95)
plot(gg_dta)

gg_dta <- gg_rfsrc(rfsrc_pbc, by="treatment")
plot(gg_dta)

## End(Not run)

```

gg_roc.rfsrc	<i>ROC (Receiver operator curve) data from a classification random forest.</i>
--------------	--

Description

The sensitivity and specificity of a randomForest classification object.

Usage

```

## S3 method for class 'rfsrc'
gg_roc(object, which_outcome, oob, ...)

```

Arguments

object	an <code>rfsrc</code> classification object
which_outcome	select the classification outcome of interest.
oob	use oob estimates (default TRUE)
...	extra arguments (not used)

Value

gg_roc data.frame for plotting ROC curves.

See Also

[plot.gg_roc rfsrc randomForest](#)

Examples

```
## -----  
## classification example  
## -----  
## ----- iris data  
rfsrc_iris <- rfsrc(Species ~ ., data = iris)  
  
# ROC for setosa  
gg_dta <- gg_roc(rfsrc_iris, which_outcome=1)  
plot(gg_dta)  
  
# ROC for versicolor  
gg_dta <- gg_roc(rfsrc_iris, which_outcome=2)  
plot(gg_dta)  
  
# ROC for virginica  
gg_dta <- gg_roc(rfsrc_iris, which_outcome=3)  
plot(gg_dta)  
  
## ----- iris data  
rf_iris <- randomForest::randomForest(Species ~ ., data = iris)  
  
# ROC for setosa  
gg_dta <- gg_roc(rf_iris, which_outcome=1)  
plot(gg_dta)  
  
# ROC for versicolor  
gg_dta <- gg_roc(rf_iris, which_outcome=2)  
plot(gg_dta)  
  
# ROC for virginica  
gg_dta <- gg_roc(rf_iris, which_outcome=3)  
plot(gg_dta)
```

gg_survival

Nonparametric survival estimates.

Description

Nonparametric survival estimates.

Usage

```
gg_survival(  
  interval = NULL,  
  censor = NULL,  
  by = NULL,
```

```

    data,
    type = c("kaplan", "nelson"),
    ...
  )

```

Arguments

interval	name of the interval variable in the training dataset.
censor	name of the censoring variable in the training dataset.
by	stratifying variable in the training dataset, defaults to NULL
data	name of the training data.frame
type	one of ("kaplan","nelson"), defaults to Kaplan-Meier
...	extra arguments passed to Kaplan or Nelson functions.

Details

gg_survival is a wrapper function for generating nonparametric survival estimates using either [nelson-Aalen](#) or [kaplan-Meier](#) estimates.

Value

A gg_survival object created using the non-parametric Kaplan-Meier or Nelson-Aalen estimators.

See Also

[kaplan nelson](#)
[plot.gg_survival](#)

Examples

```

## Not run:
## ----- pbc data
data(pbc, package="randomForestSRC")
pbc$time <- pbc$days/364.25

# This is the same as kaplan
gg_dta <- gg_survival(interval="time", censor="status",
                     data=pbc)

plot(gg_dta, error="none")
plot(gg_dta)

# Stratified on treatment variable.
gg_dta <- gg_survival(interval="time", censor="status",
                     data=pbc, by="treatment")

plot(gg_dta, error="none")
plot(gg_dta)

```

```
# ...with smaller confidence limits.
gg_dta <- gg_survival(interval="time", censor="status",
                     data=pbcc, by="treatment", conf.int=.68)

plot(gg_dta, error="lines")

## End(Not run)
```

gg_variable

Marginal variable dependence data object.

Description

`plot.variable` generates a `data.frame` containing the marginal variable dependence or the partial variable dependence. The `gg_variable` function creates a `data.frame` of containing the full set of covariate data (predictor variables) and the predicted response for each observation. Marginal dependence figures are created using the `plot.gg_variable` function.

Optional arguments `time point` (or vector of points) of interest (for survival forests only) `time_labels` If more than one time is specified, a vector of time labels for differentiating the time points (for survival forests only) `oob` indicate if predicted results should include oob or full data set.

Usage

```
gg_variable(object, ...)
```

Arguments

<code>object</code>	a <code>rfsrc</code> object
<code>...</code>	optional arguments

Details

The marginal variable dependence is determined by comparing relation between the predicted response from the `randomForest` and a covariate of interest.

The `gg_variable` function operates on a `rfsrc` object, or the output from the `plot.variable` function.

Value

`gg_variable` object

See Also

[plot.gg_variable](#)
[plot.variable](#)

Examples

```

## -----
## classification
## -----
## ----- iris data
## iris
rfsrc_iris <- rfsrc(Species ~., data = iris)

gg_dta <- gg_variable(rfsrc_iris)
plot(gg_dta, xvar="Sepal.Width")
plot(gg_dta, xvar="Sepal.Length")

plot(gg_dta, xvar=rfsrc_iris$xvar.names,
      panel=TRUE) # , se=FALSE)

## -----
## regression
## -----
## Not run:
## ----- air quality data
rfsrc_airq <- rfsrc(Ozone ~ ., data = airquality)
gg_dta <- gg_variable(rfsrc_airq)

# an ordinal variable
gg_dta[, "Month"] <- factor(gg_dta[, "Month"])

plot(gg_dta, xvar="Wind")
plot(gg_dta, xvar="Temp")
plot(gg_dta, xvar="Solar.R")

plot(gg_dta, xvar=c("Solar.R", "Wind", "Temp", "Day"), panel=TRUE)

plot(gg_dta, xvar="Month", notch=TRUE)

## End(Not run)
## Not run:
## ----- motor trend cars data
rfsrc_mtcars <- rfsrc(mpg ~ ., data = mtcars)

gg_dta <- gg_variable(rfsrc_mtcars)

# mtcars$cyl is an ordinal variable
gg_dta$cyl <- factor(gg_dta$cyl)
gg_dta$am <- factor(gg_dta$am)
gg_dta$vs <- factor(gg_dta$vs)
gg_dta$gear <- factor(gg_dta$gear)
gg_dta$carb <- factor(gg_dta$carb)

plot(gg_dta, xvar="cyl")

# Others are continuous

```

```

plot(gg_dta, xvar="disp")
plot(gg_dta, xvar="hp")
plot(gg_dta, xvar="wt")

# panels
plot(gg_dta,xvar=c("disp","hp", "drat", "wt", "qsec"), panel=TRUE)
plot(gg_dta, xvar=c("cyl", "vs", "am", "gear", "carb"), panel=TRUE,
      notch=TRUE)

## End(Not run)
## ----- Boston data
data(Boston, package="MASS")

rf_boston <- randomForest::randomForest(medv~., data=Boston)
gg_dta <- gg_variable(rf_boston)
plot(gg_dta)
plot(gg_dta, panel = TRUE)
## -----
## survival examples
## -----
## Not run:
## ----- veteran data
## survival
data(veteran, package = "randomForestSRC")
rfsrc_veteran <- rfsrc(Surv(time,status)~., veteran, nsplit = 10,
                      ntree = 100)

# get the 1 year survival time.
gg_dta <- gg_variable(rfsrc_veteran, time=90)

# Generate variable dependence plots for age and diagtime
plot(gg_dta, xvar = "age")
plot(gg_dta, xvar = "diagtime", )

# Generate coplots
plot(gg_dta, xvar = c("age", "diagtime"), panel=TRUE, se=FALSE)

# If we want to compare survival at different time points, say 30, 90 day
# and 1 year
gg_dta <- gg_variable(rfsrc_veteran, time=c(30, 90, 365))

# Generate variable dependence plots for age and diagtime
plot(gg_dta, xvar = "age")

## End(Not run)
## Not run:
## ----- pbc data
## We don't run this because of bootstrap confidence limits
# We need to create this dataset
data(pbc, package = "randomForestSRC",)
# For whatever reason, the age variable is in days... makes no sense to me
for (ind in seq_len(dim(pbc)[2])) {
  if (!is.factor(pbc[, ind])) {

```

```

    if (length(unique(pbc[which(!is.na(pbc[, ind])), ind])) <= 2) {
      if (sum(range(pbc[, ind], na.rm = TRUE) == c(0, 1)) == 2) {
        pbc[, ind] <- as.logical(pbc[, ind])
      }
    }
  } else {
    if (length(unique(pbc[which(!is.na(pbc[, ind])), ind])) <= 2) {
      if (sum(sort(unique(pbc[, ind])) == c(0, 1)) == 2) {
        pbc[, ind] <- as.logical(pbc[, ind])
      }
      if (sum(sort(unique(pbc[, ind])) == c(FALSE, TRUE)) == 2) {
        pbc[, ind] <- as.logical(pbc[, ind])
      }
    }
  }
}
if (!is.logical(pbc[, ind]) &
    length(unique(pbc[which(!is.na(pbc[, ind])), ind])) <= 5) {
  pbc[, ind] <- factor(pbc[, ind])
}
}
#Convert age to years
pbc$age <- pbc$age / 364.24

pbc$years <- pbc$days / 364.24
pbc <- pbc[, -which(colnames(pbc) == "days")]
pbc$treatment <- as.numeric(pbc$treatment)
pbc$treatment[which(pbc$treatment == 1)] <- "DPCA"
pbc$treatment[which(pbc$treatment == 2)] <- "placebo"
pbc$treatment <- factor(pbc$treatment)
dta_train <- pbc[-which(is.na(pbc$treatment)), ]
# Create a test set from the remaining patients
pbc_test <- pbc[which(is.na(pbc$treatment)), ]

#=====
# build the forest:
rfsrc_pbc <- randomForestSRC::rfsrc(
  Surv(years, status) ~ .,
  dta_train,
  nsplit = 10,
  na.action = "na.impute",
  forest = TRUE,
  importance = TRUE,
  save.memory = TRUE
)

gg_dta <- gg_variable(rfsrc_pbc, time=c(.5, 1, 3))
plot(gg_dta, xvar = "age")
plot(gg_dta, xvar = "trig")

# Generate coplots
plot(gg_dta, xvar = c("age", "trig"), panel=TRUE, se=FALSE)

```

```
## End(Not run)
```

```
gg_vimp          Variable Importance (VIMP) data object
```

Description

gg_vimp Extracts the variable importance (VIMP) information from a `rfsrc` object.

Usage

```
gg_vimp(object, nvar, ...)
```

Arguments

object	A <code>rfsrc</code> object or output from <code>vimp</code>
nvar	argument to control the number of variables included in the output.
...	arguments passed to the <code>vimp.rfsrc</code> function if the <code>rfsrc</code> object does not contain importance information.

Value

gg_vimp object. A `data.frame` of VIMP measures, in rank order.

References

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

See Also

`plot.gg_vimp.rfsrc`
`vimp`

Examples

```
## -----
## classification example
## -----
## ----- iris data
rfsrc_iris <- rfsrc(Species ~ ., data = iris,
                  importance = TRUE)
gg_dta <- gg_vimp(rfsrc_iris)
plot(gg_dta)

## -----
## regression example
```

```

## -----
## Not run:
## ----- air quality data
rfsrc_airq <- rfsrc(Ozone ~ ., airquality,
                  importance = TRUE)
gg_dta <- gg_vimp(rfsrc_airq)
plot(gg_dta)

## End(Not run)

## ----- Boston data
data(Boston, package="MASS")
rfsrc_boston <- randomForestSRC::rfsrc(medv~., Boston,
                                       importance = TRUE)
gg_dta <- gg_vimp(rfsrc_boston)
plot(gg_dta)

## ----- Boston data
rf_boston <- randomForest::randomForest(medv~., Boston)
gg_dta <- gg_vimp(rf_boston)
plot(gg_dta)

## Not run:
## ----- mtcars data
rfsrc_mtcars <- rfsrc(mpg ~ ., data = mtcars,
                    importance = TRUE)
gg_dta <- gg_vimp(rfsrc_mtcars)
plot(gg_dta)

## End(Not run)
## -----
## survival example
## -----
## Not run:
## ----- veteran data
data(veteran, package = "randomForestSRC")
rfsrc_veteran <- rfsrc(Surv(time, status) ~ .,
                      data = veteran,
                      ntree = 100,
                      importance = TRUE)

gg_dta <- gg_vimp(rfsrc_veteran)
plot(gg_dta)

## ----- pbc data
# We need to create this dataset
data(pbc, package = "randomForestSRC",)
# For whatever reason, the age variable is in days...
# makes no sense to me
for (ind in seq_len(dim(pbc)[2])) {
  if (!is.factor(pbc[, ind])) {
    if (length(unique(pbc[which(!is.na(pbc[, ind])), ind])) <= 2) {
      if (sum(range(pbc[, ind], na.rm = TRUE) == c(0, 1)) == 2) {

```



```

    pbc[, ind] <- as.logical(pbc[, ind])
  }
} else {
  if (length(unique(pbc[which(!is.na(pbc[, ind])), ind])) <= 2) {
    if (sum(sort(unique(pbc[, ind])) == c(0, 1)) == 2) {
      pbc[, ind] <- as.logical(pbc[, ind])
    }
    if (sum(sort(unique(pbc[, ind])) == c(FALSE, TRUE)) == 2) {
      pbc[, ind] <- as.logical(pbc[, ind])
    }
  }
}
if (!is.logical(pbc[, ind]) &
    length(unique(pbc[which(!is.na(pbc[, ind])), ind])) <= 5) {
  pbc[, ind] <- factor(pbc[, ind])
}
}
#Convert age to years
pbc$age <- pbc$age / 364.24

pbc$years <- pbc$days / 364.24
pbc <- pbc[, -which(colnames(pbc) == "days")]
pbc$treatment <- as.numeric(pbc$treatment)
pbc$treatment[which(pbc$treatment == 1)] <- "DPCA"
pbc$treatment[which(pbc$treatment == 2)] <- "placebo"
pbc$treatment <- factor(pbc$treatment)
dta_train <- pbc[-which(is.na(pbc$treatment)), ]
# Create a test set from the remaining patients
pbc_test <- pbc[which(is.na(pbc$treatment)), ]

#####
# build the forest:
rfsrc_pbc <- randomForestSRC::rfsrc(
  Surv(years, status) ~ .,
  dta_train,
  nsplit = 10,
  na.action = "na.impute",
  forest = TRUE,
  importance = TRUE,
  save.memory = TRUE
)

gg_dta <- gg_vimp(rfsrc_pbc)
plot(gg_dta)

# Restrict to only the top 10.
gg_dta <- gg_vimp(rfsrc_pbc, nvar=10)
plot(gg_dta)

## End(Not run)

```

kaplan	<i>nonparametric Kaplan-Meier estimates</i>
--------	---

Description

nonparametric Kaplan-Meier estimates

Usage

```
kaplan(interval, censor, data, by = NULL, ...)
```

Arguments

interval	name of the interval variable in the training dataset.
censor	name of the censoring variable in the training dataset.
data	name of the training set data. frame
by	stratifying variable in the training dataset, defaults to NULL
...	arguments passed to the survfit function

Value

[gg_survival](#) object

See Also

[gg_survival](#) [nelson](#) [plot.gg_survival](#)

Examples

```
## Not run:
# These get run through the gg_survival examples.
data(pbc, package="randomForestSRC")
pbc$time <- pbc$days/364.25

# This is the same as gg_survival
gg_dta <- kaplan(interval="time", censor="status",
                 data=pbc)

plot(gg_dta, error="none")
plot(gg_dta)

# Stratified on treatment variable.
gg_dta <- gg_survival(interval="time", censor="status",
                     data=pbc, by="treatment")

plot(gg_dta, error="none")
plot(gg_dta)

## End(Not run)
```

nelson	<i>nonparametric Nelson-Aalen estimates</i>
--------	---

Description

nonparametric Nelson-Aalen estimates

Usage

```
nelson(interval, censor, data, by = NULL, weight = NULL, ...)
```

Arguments

interval	name of the interval variable in the training dataset.
censor	name of the censoring variable in the training dataset.
data	name of the survival training data.frame
by	stratifying variable in the training dataset, defaults to NULL
weight	for each observation (default=NULL)
...	arguments passed to the survfit function

Value

[gg_survival](#) object

See Also

[gg_survival nelson plot.gg_survival](#)

Examples

```
## Not run:
# These get run through the gg_survival examples.
data(pbc, package="randomForestSRC")
pbc$time <- pbc$days/364.25

# This is the same as gg_survival
gg_dta <- nelson(interval="time", censor="status",
                 data=pbc)

plot(gg_dta, error="none")
plot(gg_dta)

# Stratified on treatment variable.
gg_dta <- gg_survival(interval="time", censor="status",
                     data=pbc, by="treatment")

plot(gg_dta, error="none")
```

```
plot(gg_dta, error="lines")
plot(gg_dta)

gg_dta <- gg_survival(interval="time", censor="status",
                     data=pbcc, by="treatment",
                     type="nelson")

plot(gg_dta, error="bars")
plot(gg_dta)

## End(Not run)
```

plot.gg_error

Plot a gg_error object

Description

A plot of the cumulative OOB error rates of the random forest as a function of number of trees.

Usage

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

Arguments

x	gg_error object created from a rfsrc object
...	extra arguments passed to ggplot functions

Details

The `gg_error` plot is used to track the convergence of the randomForest. This figure is a reproduction of the error plot from the [plot.rfsrc](#) function.

Value

ggplot object

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.

Ishwaran H. and Kogalur U.B. (2013). Random Forests for Survival, Regression and Classification (RF-SRC), R package version 1.4.

See Also

[gg_error](#) [rfsrc](#) [plot.rfsrc](#)

Examples

```

## Not run:
## Examples from RFSRC package...
## -----
## classification example
## -----
## ----- iris data
## You can build a randomForest
rfsrc_iris <- rfsrc(Species ~ ., data = iris, tree.err = TRUE)

# Get a data.frame containing error rates
gg_dta <- gg_error(rfsrc_iris)

# Plot the gg_error object
plot(gg_dta)

## RandomForest example
rf_iris <- randomForest::randomForest(Species ~ ., data = iris,
                                     tree.err = TRUE, )
gg_dta <- gg_error(rf_iris)
plot(gg_dta)

gg_dta <- gg_error(rf_iris, training=TRUE)
plot(gg_dta)
## -----
## Regression example
## -----
## ----- airq data
rfsrc_airq <- rfsrc(Ozone ~ ., data = airquality,
                  na.action = "na.impute", tree.err = TRUE, )

# Get a data.frame containing error rates
gg_dta <- gg_error(rfsrc_airq)

# Plot the gg_error object
plot(gg_dta)

## ----- Boston data
data(Boston, package = "MASS")
Boston$chas <- as.logical(Boston$chas)
rfsrc_boston <- rfsrc(medv ~ .,
                    data = Boston,
                    forest = TRUE,
                    importance = TRUE,
                    tree.err = TRUE,
                    save.memory = TRUE)

# Get a data.frame containing error rates
gg_dta <- gg_error(rfsrc_boston)

# Plot the gg_error object

```

```

plot(gg_dta)

## ----- mtcars data
rfsrc_mtcars <- rfsrc(mpg ~ ., data = mtcars, tree.err = TRUE)
# Get a data.frame containing error rates
gg_dta<- gg_error(rfsrc_mtcars)

# Plot the gg_error object
plot(gg_dta)

## -----
## Survival example
## -----
## ----- veteran data
## randomized trial of two treatment regimens for lung cancer
data(veteran, package = "randomForestSRC")
rfsrc_veteran <- rfsrc(Surv(time, status) ~ ., data = veteran,
                      tree.err = TRUE)

gg_dta <- gg_error(rfsrc_veteran)
plot(gg_dta)

## ----- pbc data
# Load a cached randomForestSRC object
# We need to create this dataset
data(pbc, package = "randomForestSRC",)
# For whatever reason, the age variable is in days... makes no sense to me
for (ind in seq_len(dim(pbc)[2])) {
  if (!is.factor(pbc[, ind])) {
    if (length(unique(pbc[which(!is.na(pbc[, ind])), ind])) <= 2) {
      if (sum(range(pbc[, ind], na.rm = TRUE) == c(0, 1)) == 2) {
        pbc[, ind] <- as.logical(pbc[, ind])
      }
    }
  } else {
    if (length(unique(pbc[which(!is.na(pbc[, ind])), ind])) <= 2) {
      if (sum(sort(unique(pbc[, ind])) == c(0, 1)) == 2) {
        pbc[, ind] <- as.logical(pbc[, ind])
      }
      if (sum(sort(unique(pbc[, ind])) == c(FALSE, TRUE)) == 2) {
        pbc[, ind] <- as.logical(pbc[, ind])
      }
    }
  }
}
if (!is.logical(pbc[, ind]) &
    length(unique(pbc[which(!is.na(pbc[, ind])), ind])) <= 5) {
  pbc[, ind] <- factor(pbc[, ind])
}
}
#Convert age to years
pbc$age <- pbc$age / 364.24

```

```

pbc$years <- pbc$days / 364.24
pbc <- pbc[, -which(colnames(pbc) == "days")]
pbc$treatment <- as.numeric(pbc$treatment)
pbc$treatment[which(pbc$treatment == 1)] <- "DPCA"
pbc$treatment[which(pbc$treatment == 2)] <- "placebo"
pbc$treatment <- factor(pbc$treatment)
dta_train <- pbc[-which(is.na(pbc$treatment)), ]
# Create a test set from the remaining patients
pbc_test <- pbc[which(is.na(pbc$treatment)), ]

#=====
# build the forest:
rfsrc_pbc <- randomForestSRC::rfsrc(
  Surv(years, status) ~ .,
  dta_train,
  nsplit = 10,
  na.action = "na.impute",
  tree.err = TRUE,
  forest = TRUE,
  importance = TRUE,
  save.memory = TRUE
)

gg_dta <- gg_error(rfsrc_pbc)
plot(gg_dta)

## End(Not run)

```

plot.gg_interaction *plot.gg_interaction* Plot a [gg_interaction](#) object,

Description

plot.gg_interaction Plot a [gg_interaction](#) object,

Usage

```

## S3 method for class 'gg_interaction'
plot(x, xvar, lbls, ...)

```

Arguments

x	gg_interaction object created from a rfsrc object
xvar	variable (or list of variables) of interest.
lbls	A vector of alternative variable names.
...	arguments passed to the gg_interaction function.

Value

ggplot object

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.

Ishwaran H. and Kogalur U.B. (2013). Random Forests for Survival, Regression and Classification (RF-SRC), R package version 1.4.

See Also

[rfsrc](#) [find.interaction](#) [max.subtree](#) [var.select](#) [vimp](#) [plot.gg_interaction](#)

Examples

```
## Not run:
## Examples from randomForestSRC package...
## -----
## find interactions, classification setting
## -----
## ----- iris data
## iris.obj <- rfsrc(Species ~., data = iris)
## TODO: VIMP interactions not handled yet...
## find.interaction(iris.obj, method = "vimp", nrep = 3)
## interaction_iris <- find.interaction(iris.obj)
data(interaction_iris, package="ggRandomForests")
gg_dta <- gg_interaction(interaction_iris)

plot(gg_dta, xvar="Petal.Width")
plot(gg_dta, xvar="Petal.Length")
plot(gg_dta, panel=TRUE)

## -----
## find interactions, regression setting
## -----
## ----- air quality data
## airq.obj <- rfsrc(Ozone ~ ., data = airquality)
##
## TODO: VIMP interactions not handled yet...
## find.interaction(airq.obj, method = "vimp", nrep = 3)
## interaction_airq <- find.interaction(airq.obj)
data(interaction_airq, package="ggRandomForests")
gg_dta <- gg_interaction(interaction_airq)

plot(gg_dta, xvar="Temp")
plot(gg_dta, xvar="Solar.R")
plot(gg_dta, panel=TRUE)

## ----- Boston data
data(interaction_boston, package="ggRandomForests")
```



```

gg_dta <- gg_interaction(interaction_boston)

plot(gg_dta, panel=TRUE)

## ----- mtcars data
data(interaction_mtcars, package="ggRandomForests")
gg_dta <- gg_interaction(interaction_mtcars)

plot(gg_dta, panel=TRUE)

## -----
## find interactions, survival setting
## -----
## ----- pbc data
## data(pbc, package = "randomForestSRC")
## pbc.obj <- rfsrc(Surv(days,status) ~ ., pbc, nsplit = 10)
## interaction_pbc <- find.interaction(pbc.obj, nvar = 8)
data(interaction_pbc, package="ggRandomForests")
gg_dta <- gg_interaction(interaction_pbc)

plot(gg_dta, xvar="bili")
plot(gg_dta, xvar="copper")
plot(gg_dta, panel=TRUE)

## ----- veteran data
data(interaction_veteran, package="ggRandomForests")
gg_dta <- gg_interaction(interaction_veteran)

plot(gg_dta, panel=TRUE)

## End(Not run)

```

plot.gg_minimal_depth *Plot a [gg_minimal_depth](#) object for random forest variable ranking.*

Description

Plot a [gg_minimal_depth](#) object for random forest variable ranking.

Usage

```
## S3 method for class 'gg_minimal_depth'
plot(x, selection = FALSE, type = c("named", "rank"), lbls, ...)
```

Arguments

x [gg_minimal_depth](#) object created from a [rfsrc](#) object

selection	should we restrict the plot to only include variables selected by the minimal depth criteria (boolean).
type	select type of y axis labels c("named","rank")
lbls	a vector of alternative variable names.
...	optional arguments passed to gg_minimal_depth

Value

ggplot object

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.

Ishwaran H. and Kogalur U.B. (2014). Random Forests for Survival, Regression and Classification (RF-SRC), R package version 1.5.

See Also

[var.select](#) [gg_minimal_depth](#)

Examples

```
## Not run:
## Examples from RFSRC package...
## -----
## classification example
## -----
## ----- iris data
## You can build a randomForest
rfsrc_iris <- rfsrc(Species ~ ., data = iris)
varsel_iris <- var.select(rfsrc_iris)

# Get a data.frame containing minimaldepth measures
gg_dta<- gg_minimal_depth(varsel_iris)

# Plot the gg_minimal_depth object
plot(gg_dta)

## -----
## Regression example
## -----
## ----- air quality data
rfsrc_airq <- rfsrc(Ozone ~ ., data = airquality, na.action = "na.impute")
varsel_airq <- var.select(rfsrc_airq)

# Get a data.frame containing error rates
gg_dta<- gg_minimal_depth(varsel_airq)

# Plot the gg_minimal_depth object
```

```

plot(gg_dta)

## ----- Boston data
data(Boston, package="MASS")
rfsrc_boston <- randomForestSRC::rfsrc(medv~., Boston)
# Get a data.frame containing error rates
plot(gg_minimal_depth(varsel_boston))

## ----- mtcars data
rfsrc_mtcars <- rfsrc(mpg ~ ., data = mtcars)
varsel_mtcars <- var.select(rfsrc_mtcars)

# Get a data.frame containing error rates
plot.gg_minimal_depth(varsel_mtcars)

## -----
## Survival example
## -----
## ----- veteran data
## randomized trial of two treatment regimens for lung cancer
data(veteran, package = "randomForestSRC")
rfsrc_veteran <- rfsrc(Surv(time, status) ~ ., data = veteran, ntree = 100)
varsel_veteran <- var.select(rfsrc_veteran)

gg_dta <- gg_minimal_depth(varsel_veteran)
plot(gg_dta)

## ----- pbc data
#' # We need to create this dataset
data(pbc, package = "randomForestSRC",)
# For whatever reason, the age variable is in days... makes no sense to me
for (ind in seq_len(dim(pbc)[2])) {
  if (!is.factor(pbc[, ind])) {
    if (length(unique(pbc[which(!is.na(pbc[, ind])), ind])) <= 2) {
      if (sum(range(pbc[, ind], na.rm = TRUE) == c(0, 1)) == 2) {
        pbc[, ind] <- as.logical(pbc[, ind])
      }
    }
  } else {
    if (length(unique(pbc[which(!is.na(pbc[, ind])), ind])) <= 2) {
      if (sum(sort(unique(pbc[, ind])) == c(0, 1)) == 2) {
        pbc[, ind] <- as.logical(pbc[, ind])
      }
      if (sum(sort(unique(pbc[, ind])) == c(FALSE, TRUE)) == 2) {
        pbc[, ind] <- as.logical(pbc[, ind])
      }
    }
  }
}
if (!is.logical(pbc[, ind]) &
    length(unique(pbc[which(!is.na(pbc[, ind])), ind])) <= 5) {
  pbc[, ind] <- factor(pbc[, ind])
}

```

```

}
#Convert age to years
pbc$age <- pbc$age / 364.24

pbc$years <- pbc$days / 364.24
pbc <- pbc[, -which(colnames(pbc) == "days")]
pbc$treatment <- as.numeric(pbc$treatment)
pbc$treatment[which(pbc$treatment == 1)] <- "DPCA"
pbc$treatment[which(pbc$treatment == 2)] <- "placebo"
pbc$treatment <- factor(pbc$treatment)
dta_train <- pbc[-which(is.na(pbc$treatment)), ]
# Create a test set from the remaining patients
pbc_test <- pbc[which(is.na(pbc$treatment)), ]

#=====
# build the forest:
rfsrc_pbc <- randomForestSRC::rfsrc(
  Surv(years, status) ~ .,
  dta_train,
  nsplit = 10,
  na.action = "na.impute",
  forest = TRUE,
  importance = TRUE,
  save.memory = TRUE
)

varsel_pbc <- var.select(rfsrc_pbc)

gg_dta <- gg_minimal_depth(varsel_pbc)
plot(gg_dta)

## End(Not run)

```

plot.gg_minimal_vimp *Plot a gg_minimal_vimp object for comparing the Minimal Depth and VIMP variable rankings.*

Description

Plot a `gg_minimal_vimp` object for comparing the Minimal Depth and VIMP variable rankings.

Usage

```

## S3 method for class 'gg_minimal_vimp'
plot(x, nvar, lbls, ...)

```

Arguments

x [gg_minimal_depth](#) object created from a `var.select` object
 nvar should the figure be restricted to a subset of the points.
 lbls a vector of alternative variable names.
 ... optional arguments (not used)

Value

ggplot object

See Also

[gg_minimal_vimp](#) [var.select](#)

Examples

```
## Not run:
## Examples from RFSRC package...
## -----
## classification example
## -----
## ----- iris data
## You can build a randomForest
rfsrc_iris <- rfsrc(Species ~ ., data = iris)
varsel_iris <- var.select(rfsrc_iris)

# Get a data.frame containing minimaldepth measures
gg_dta<- gg_minimal_vimp(varsel_iris)

# Plot the gg_minimal_depth object
plot(gg_dta)

## -----
## Regression example
## -----
## ----- air quality data
rfsrc_airq <- rfsrc(Ozone ~ ., data = airquality, na.action = "na.impute")
varsel_airq <- var.select(rfsrc_airq)

# Get a data.frame containing error rates
gg_dta<- gg_minimal_vimp(varsel_airq)

# Plot the gg_minimal_vimp object
plot(gg_dta)

## ----- Boston data
data(Boston, package="MASS")
rfsrc_boston <- randomForestSRC::rfsrc(medv~., Boston)

varsel_boston <- var.select(rfsrc_boston)
```

```

# Get a data.frame containing error rates
gg_dta<- gg_minimal_vimp(varsel_boston)

# Plot the gg_minimal_vimp object
plot(gg_dta)

## ----- mtcars data
rfsrc_mtcars <- rfsrc(mpg ~ ., data = mtcars)
varsel_mtcars <- var.select(rfsrc_mtcars)

# Get a data.frame containing error rates
gg_dta<- gg_minimal_vimp(varsel_mtcars)

# Plot the gg_minimal_vimp object
plot(gg_dta)

## -----
## Survival example
## -----
## ----- veteran data
## randomized trial of two treatment regimens for lung cancer
data(veteran, package = "randomForestSRC")
rfsrc_veteran <- rfsrc(Surv(time, status) ~ ., data = veteran, ntree = 100)
varsel_veteran <- var.select(rfsrc_veteran)

gg_dta <- gg_minimal_vimp(varsel_veteran)
plot(gg_dta)

## ----- pbc data
# We need to create this dataset
data(pbc, package = "randomForestSRC",)
# For whatever reason, the age variable is in days... makes no sense to me
for (ind in seq_len(dim(pbc)[2])) {
  if (!is.factor(pbc[, ind])) {
    if (length(unique(pbc[which(!is.na(pbc[, ind])), ind])) <= 2) {
      if (sum(range(pbc[, ind], na.rm = TRUE) == c(0, 1)) == 2) {
        pbc[, ind] <- as.logical(pbc[, ind])
      }
    }
  } else {
    if (length(unique(pbc[which(!is.na(pbc[, ind])), ind])) <= 2) {
      if (sum(sort(unique(pbc[, ind])) == c(0, 1)) == 2) {
        pbc[, ind] <- as.logical(pbc[, ind])
      }
      if (sum(sort(unique(pbc[, ind])) == c(FALSE, TRUE)) == 2) {
        pbc[, ind] <- as.logical(pbc[, ind])
      }
    }
  }
}
if (!is.logical(pbc[, ind]) &
    length(unique(pbc[which(!is.na(pbc[, ind])), ind])) <= 5) {
  pbc[, ind] <- factor(pbc[, ind])
}

```

```

}
#Convert age to years
pbc$age <- pbc$age / 364.24

pbc$years <- pbc$days / 364.24
pbc <- pbc[, -which(colnames(pbc) == "days")]
pbc$treatment <- as.numeric(pbc$treatment)
pbc$treatment[which(pbc$treatment == 1)] <- "DPCA"
pbc$treatment[which(pbc$treatment == 2)] <- "placebo"
pbc$treatment <- factor(pbc$treatment)
dta_train <- pbc[-which(is.na(pbc$treatment)), ]
# Create a test set from the remaining patients
pbc_test <- pbc[which(is.na(pbc$treatment)), ]

#=====
# build the forest:
rfsrc_pbc <- randomForestSRC::rfsrc(
  Surv(years, status) ~ .,
  dta_train,
  nsplit = 10,
  na.action = "na.impute",
  forest = TRUE,
  importance = TRUE,
  save.memory = TRUE
)

varsel_pbc <- var.select(rfsrc_pbc)

gg_dta <- gg_minimal_vimp(varsel_pbc)
plot(gg_dta)

## End(Not run)

```

plot.gg_partial

Partial variable dependence plot, operates on a [gg_partial](#) object.

Description

Generate a risk adjusted (partial) variable dependence plot. The function plots the [rfsrc](#) response variable (y-axis) against the covariate of interest (specified when creating the [gg_partial](#) object).

Usage

```

## S3 method for class 'gg_partial'
plot(x, points = TRUE, error = c("none", "shade", "bars", "lines"), ...)

```

Arguments

x	<code>gg_partial</code> object created from a <code>rfsrc</code> forest object
points	plot points (boolean) or a smooth line.
error	"shade", "bars", "lines" or "none"
...	extra arguments passed to <code>ggplot2</code> functions.

Value

ggplot object

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.

Ishwaran H. and Kogalur U.B. (2013). Random Forests for Survival, Regression and Classification (RF-SRC), R package version 1.4.

See Also

[plot.variable](#) [gg_partial](#) [plot.gg_partial_list](#) [gg_variable](#) [plot.gg_variable](#)

Examples

```
## Not run:
## -----
## classification
## -----
## ----- iris data

## iris "Petal.Width" partial dependence plot
##
# rfsrc_iris <- rfsrc(Species ~ ., data = iris)
# partial_iris <- plot.variable(rfsrc_iris, xvar.names = "Petal.Width",
#                             partial=TRUE)
# data(partial_iris, package="ggRandomForests")

gg_dta <- gg_partial(partial_iris)
plot(gg_dta)

## -----
## regression
## -----
## ----- air quality data
## airquality "Wind" partial dependence plot
##
# rfsrc_airq <- rfsrc(Ozone ~ ., data = airquality)
# partial_airq <- plot.variable(rfsrc_airq, xvar.names = "Wind",
#                              partial=TRUE, show.plot=FALSE)
# data(partial_airq, package="ggRandomForests")
```



```

gg_dta <- gg_partial(partial_airq)
plot(gg_dta)

gg_dta.m <- gg_dta[["Month"]]
plot(gg_dta.m, notch=TRUE)

gg_dta[["Month"]] <- NULL
plot(gg_dta, panel=TRUE)

## ----- Boston data
data(partial_boston, package="ggRandomForests")

gg_dta <- gg_partial(partial_boston)
plot(gg_dta)
plot(gg_dta, panel=TRUE)

## ----- mtcars data
data(partial_mtcars, package="ggRandomForests")

gg_dta <- gg_partial(partial_mtcars)

plot(gg_dta)

gg_dta.cat <- gg_dta
gg_dta.cat[["disp"]] <- gg_dta.cat[["wt"]] <- gg_dta.cat[["hp"]] <- NULL
gg_dta.cat[["drat"]] <- gg_dta.cat[["carb"]] <-
  gg_dta.cat[["qsec"]] <- NULL

plot(gg_dta.cat, panel=TRUE)

gg_dta[["cyl"]] <- gg_dta[["vs"]] <- gg_dta[["am"]] <- NULL
gg_dta[["gear"]] <- NULL
plot(gg_dta, panel=TRUE)

## -----
## survival examples
## -----
## ----- veteran data
## survival "age" partial variable dependence plot
##
# data(veteran, package = "randomForestSRC")
# rfsrc_veteran <- rfsrc(Surv(time,status)~., veteran, nsplit = 10,
#                       ntree = 100)
#
## 30 day partial plot for age
# partial_veteran <- plot.variable(rfsrc_veteran, surv.type = "surv",
#                                 partial = TRUE, time=30,
#                                 xvar.names = "age",
#                                 show.plots=FALSE)
data(partial_veteran, package="ggRandomForests")

gg_dta <- gg_partial(partial_veteran[[1]])

```

```

plot(gg_dta)

gg_dta.cat <- gg_dta
gg_dta[["celltype"]] <- gg_dta[["trt"]] <- gg_dta[["prior"]] <- NULL
plot(gg_dta, panel=TRUE)

gg_dta.cat[["karno"]] <- gg_dta.cat[["diagtime"]] <-
  gg_dta.cat[["age"]] <- NULL
plot(gg_dta.cat, panel=TRUE, notch=TRUE)

gg_dta <- lapply(partial_veteran, gg_partial)
length(gg_dta)
gg_dta <- combine.gg_partial(gg_dta[[1]], gg_dta[[2]] )

plot(gg_dta[["karno"]])
plot(gg_dta[["celltype"]])

gg_dta.cat <- gg_dta
gg_dta[["celltype"]] <- gg_dta[["trt"]] <- gg_dta[["prior"]] <- NULL
plot(gg_dta, panel=TRUE)

gg_dta.cat[["karno"]] <- gg_dta.cat[["diagtime"]] <-
  gg_dta.cat[["age"]] <- NULL
plot(gg_dta.cat, panel=TRUE, notch=TRUE)

## ----- pbc data

## End(Not run)

```

plot.gg_partial_list *Partial variable dependence plot, operates on a gg_partial_list object.*

Description

Generate a risk adjusted (partial) variable dependence plot. The function plots the `rfsrc` response variable (y-axis) against the covariate of interest (specified when creating the `gg_partial_list` object).

Usage

```

## S3 method for class 'gg_partial_list'
plot(x, points = TRUE, panel = FALSE, ...)

```

Arguments

<code>x</code>	<code>gg_partial_list</code> object created from a <code>gg_partial</code> forest object
<code>points</code>	plot points (boolean) or a smooth line.

panel should the entire list be plotted together?
 ... extra arguments

Value

list of ggplot objects, or a single faceted ggplot object

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.

Ishwaran H. and Kogalur U.B. (2013). Random Forests for Survival, Regression and Classification (RF-SRC), R package version 1.4.

See Also

[plot.variable](#) [gg_partial](#) [plot.gg_partial](#) [gg_variable](#) [plot.gg_variable](#)

Examples

```
## Not run:
## -----
## classification
## -----
## ----- iris data

## iris "Petal.Width" partial dependence plot
##
# rfsrc_iris <- rfsrc(Species ~., data = iris)
# partial_iris <- plot.variable(rfsrc_iris, xvar.names = "Petal.Width",
#                             partial=TRUE)
data(partial_iris, package="ggRandomForests")

gg_dta <- gg_partial(partial_iris)
plot(gg_dta)

## -----
## regression
## -----
## ----- air quality data
## airquality "Wind" partial dependence plot
##
# rfsrc_airq <- rfsrc(Ozone ~ ., data = airquality)
# partial_airq <- plot.variable(rfsrc_airq, xvar.names = "Wind",
#                             partial=TRUE, show.plot=FALSE)
data(partial_airq, package="ggRandomForests")

gg_dta <- gg_partial(partial_airq)
plot(gg_dta)

gg_dta.m <- gg_dta[["Month"]]
```

```

plot(gg_dta.m, notch=TRUE)

gg_dta[["Month"]] <- NULL
plot(gg_dta, panel=TRUE)

## ----- Boston data
data(partial_boston, package="ggRandomForests")

gg_dta <- gg_partial(partial_boston)
plot(gg_dta)
plot(gg_dta, panel=TRUE)

## ----- mtcars data
data(partial_mtcars, package="ggRandomForests")

gg_dta <- gg_partial(partial_mtcars)

plot(gg_dta)

gg_dta.cat <- gg_dta
gg_dta.cat[["disp"]] <- gg_dta.cat[["wt"]] <- gg_dta.cat[["hp"]] <- NULL
gg_dta.cat[["drat"]] <- gg_dta.cat[["carb"]] <- gg_dta.cat[["qsec"]] <- NULL

plot(gg_dta.cat, panel=TRUE)

gg_dta[["cyl"]] <- gg_dta[["vs"]] <- gg_dta[["am"]] <- NULL
gg_dta[["gear"]] <- NULL
plot(gg_dta, panel=TRUE)

## -----
## survival examples
## -----
## ----- veteran data
## survival "age" partial variable dependence plot
##
# data(veteran, package = "randomForestSRC")
# rfsrc_veteran <- rfsrc(Surv(time,status)~., veteran, nsplit = 10,
#                       ntree = 100)
#
## 30 day partial plot for age
# partial_veteran <- plot.variable(rfsrc_veteran, surv.type = "surv",
#                                 partial = TRUE, time=30,
#                                 xvar.names = "age",
#                                 show.plots=FALSE)
data(partial_veteran, package="ggRandomForests")

gg_dta <- gg_partial(partial_veteran[[1]])
plot(gg_dta)

gg_dta.cat <- gg_dta
gg_dta[["celltype"]] <- gg_dta[["trt"]] <- gg_dta[["prior"]] <- NULL
plot(gg_dta, panel=TRUE)

```

```

gg_dta.cat[["karno"]] <- gg_dta.cat[["diagtime"]] <-
  gg_dta.cat[["age"]] <- NULL
plot(gg_dta.cat, panel=TRUE, notch=TRUE)

gg_dta <- lapply(partial_veteran, gg_partial)
length(gg_dta)
gg_dta <- combine.gg_partial(gg_dta[[1]], gg_dta[[2]] )

plot(gg_dta[["karno"]])
plot(gg_dta[["celltype"]])

gg_dta.cat <- gg_dta
gg_dta.cat[["celltype"]] <- gg_dta[["trt"]] <- gg_dta[["prior"]] <- NULL
plot(gg_dta, panel=TRUE)

gg_dta.cat[["karno"]] <- gg_dta.cat[["diagtime"]] <-
  gg_dta.cat[["age"]] <- NULL
plot(gg_dta.cat, panel=TRUE, notch=TRUE)

## ----- pbc data

## End(Not run)

```

plot.gg_rfsrc

Predicted response plot from a gg_rfsrc object.

Description

Plot the predicted response from a `gg_rfsrc` object, the `rfsrc` prediction, using the OOB prediction from the forest.

Usage

```

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

```

Arguments

`x` `gg_rfsrc` object created from a `rfsrc` object

`...` arguments passed to `gg_rfsrc`.

Value

ggplot object

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.
- Ishwaran H. and Kogalur U.B. (2013). Random Forests for Survival, Regression and Classification (RF-SRC), R package version 1.4.

See Also

[gg_rfsrc](#) [rfsrc](#)

Examples

```
## Not run:
## -----
## classification example
## -----
## ----- iris data
# rfsrc_iris <- rfsrc(Species ~ ., data = iris)
data(rfsrc_iris, package="ggRandomForests")
gg_dta<- gg_rfsrc(rfsrc_iris)

plot(gg_dta)

## -----
## Regression example
## -----
## ----- air quality data
rfsrc_airq <- rfsrc(Ozone ~ ., data = airquality, na.action = "na.impute")
gg_dta<- gg_rfsrc(rfsrc_airq)

plot(gg_dta)

## ----- Boston data
data(Boston, package = "MASS")
rfsrc_boston <- randomForestSRC::rfsrc(medv~., Boston)

plot(rfsrc_boston)

## ----- mtcars data
rfsrc_mtcars <- rfsrc(mpg ~ ., data = mtcars)
gg_dta<- gg_rfsrc(rfsrc_mtcars)

plot(gg_dta)

## -----
## Survival example
## -----
## ----- veteran data
## randomized trial of two treatment regimens for lung cancer
data(veteran, package = "randomForestSRC")
rfsrc_veteran <- rfsrc(Surv(time, status) ~ ., data = veteran, ntree = 100)
```

```

gg_dta <- gg_rfsrc(rfsrc_veteran)
plot(gg_dta)

gg_dta <- gg_rfsrc(rfsrc_veteran, conf.int=.95)
plot(gg_dta)

gg_dta <- gg_rfsrc(rfsrc_veteran, by="trt")
plot(gg_dta)

## ----- pbc data
#' # We need to create this dataset
data(pbc, package = "randomForestSRC",)
# For whatever reason, the age variable is in days... makes no sense to me
for (ind in seq_len(dim(pbc)[2])) {
  if (!is.factor(pbc[, ind])) {
    if (length(unique(pbc[which(!is.na(pbc[, ind])), ind])) <= 2) {
      if (sum(range(pbc[, ind], na.rm = TRUE) == c(0, 1)) == 2) {
        pbc[, ind] <- as.logical(pbc[, ind])
      }
    }
  } else {
    if (length(unique(pbc[which(!is.na(pbc[, ind])), ind])) <= 2) {
      if (sum(sort(unique(pbc[, ind])) == c(0, 1)) == 2) {
        pbc[, ind] <- as.logical(pbc[, ind])
      }
      if (sum(sort(unique(pbc[, ind])) == c(FALSE, TRUE)) == 2) {
        pbc[, ind] <- as.logical(pbc[, ind])
      }
    }
  }
  if (!is.logical(pbc[, ind]) &
      length(unique(pbc[which(!is.na(pbc[, ind])), ind])) <= 5) {
    pbc[, ind] <- factor(pbc[, ind])
  }
}
#Convert age to years
pbc$age <- pbc$age / 364.24

pbc$years <- pbc$days / 364.24
pbc <- pbc[, -which(colnames(pbc) == "days")]
pbc$treatment <- as.numeric(pbc$treatment)
pbc$treatment[which(pbc$treatment == 1)] <- "DPCA"
pbc$treatment[which(pbc$treatment == 2)] <- "placebo"
pbc$treatment <- factor(pbc$treatment)
dta_train <- pbc[-which(is.na(pbc$treatment)), ]
# Create a test set from the remaining patients
pbc_test <- pbc[which(is.na(pbc$treatment)), ]

#=====
# build the forest:
rfsrc_pbc <- randomForestSRC::rfsrc(
  Surv(years, status) ~ .,
  dta_train,

```

```
nsplit = 10,  
na.action = "na.impute",  
forest = TRUE,  
importance = TRUE,  
save.memory = TRUE  
)  
  
gg_dta <- gg_rfsrc(rfsrc_pbc)  
plot(gg_dta)  
  
gg_dta <- gg_rfsrc(rfsrc_pbc, conf.int=.95)  
plot(gg_dta)  
  
gg_dta <- gg_rfsrc(rfsrc_pbc, by="treatment")  
plot(gg_dta)  
  
## End(Not run)
```

plot.gg_roc

ROC plot generic function for a [gg_roc](#) object.

Description

ROC plot generic function for a [gg_roc](#) object.

Usage

```
## S3 method for class 'gg_roc'  
plot(x, which_outcome = NULL, ...)
```

Arguments

x [gg_roc](#) object created from a classification forest
which_outcome for multiclass problems, choose the class for plotting
... arguments passed to the [gg_roc](#) function

Value

ggplot object of the ROC curve

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.
Ishwaran H. and Kogalur U.B. (2013). Random Forests for Survival, Regression and Classification (RF-SRC), R package version 1.4.

See Also[gg_roc](#) rfsrc**Examples**

```
## Not run:
## -----
## classification example
## -----
## ----- iris data
#rfsrc_iris <- rfsrc(Species ~ ., data = iris)
data(rfsrc_iris, package="ggRandomForests")

# ROC for setosa
gg_dta <- gg_roc(rfsrc_iris, which_outcome=1)
plot.gg_roc(gg_dta)

# ROC for versicolor
gg_dta <- gg_roc(rfsrc_iris, which_outcome=2)
plot.gg_roc(gg_dta)

# ROC for virginica
gg_dta <- gg_roc(rfsrc_iris, which_outcome=3)
plot.gg_roc(gg_dta)

# Alternatively, you can plot all three outcomes in one go
# by calling the plot function on the forest object.
plot.gg_roc(rfsrc_iris)

## End(Not run)
```

plot.gg_survival *Plot a [gg_survival](#) object.*

Description

Plot a [gg_survival](#) object.

Usage

```
## S3 method for class 'gg_survival'
plot(
  x,
  type = c("surv", "cum_haz", "hazard", "density", "mid_int", "life", "proplife"),
  error = c("shade", "bars", "lines", "none"),
  label = NULL,
  ...
)
```

Arguments

x	gg_survival or a survival gg_rfsrc object created from a rfsrc object
type	"surv", "cum_haz", "hazard", "density", "mid_int", "life", "proplife"
error	"shade", "bars", "lines" or "none"
label	Modify the legend label when gg_survival has stratified samples
...	not used

Value

ggplot object

Examples

```
## Not run:
## ----- pbc data
data(pbc, package="randomForestSRC")
pbc$time <- pbc$days/364.25

# This is the same as kaplan
gg_dta <- gg_survival(interval="time", censor="status",
                      data=pbc)

plot(gg_dta, error="none")
plot(gg_dta)

# Stratified on treatment variable.
gg_dta <- gg_survival(interval="time", censor="status",
                      data=pbc, by="treatment")

plot(gg_dta, error="none")
plot(gg_dta)
plot(gg_dta, label="treatment")

# ...with smaller confidence limits.
gg_dta <- gg_survival(interval="time", censor="status",
                      data=pbc, by="treatment", conf.int=.68)

plot(gg_dta, error="lines")
plot(gg_dta, label="treatment", error="lines")

# ...with smaller confidence limits.
gg_dta <- gg_survival(interval="time", censor="status",
                      data=pbc, by="sex", conf.int=.68)

plot(gg_dta, error="lines")
plot(gg_dta, label="sex", error="lines")

## End(Not run)
```

plot.gg_variable *Plot a gg_variable object,*

Description

Plot a `gg_variable` object,

Usage

```
## S3 method for class 'gg_variable'
plot(
  x,
  xvar,
  time,
  time_labels,
  panel = FALSE,
  oob = TRUE,
  points = TRUE,
  smooth = TRUE,
  ...
)
```

Arguments

<code>x</code>	<code>gg_variable</code> object created from a <code>rfsrc</code> object
<code>xvar</code>	variable (or list of variables) of interest.
<code>time</code>	For survival, one or more times of interest
<code>time_labels</code>	string labels for times
<code>panel</code>	Should plots be faceted along multiple <code>xvar</code> ?
<code>oob</code>	oob estimates (boolean)
<code>points</code>	plot the raw data points (boolean)
<code>smooth</code>	include a smooth curve (boolean)
<code>...</code>	arguments passed to the <code>ggplot2</code> functions.

Value

A single `ggplot` object, or list of `ggplot` objects

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.
- Ishwaran H. and Kogalur U.B. (2013). Random Forests for Survival, Regression and Classification (RF-SRC), R package version 1.4.

Examples

```

## Not run:
## -----
## classification
## -----
## ----- iris data
## iris
#rfsrc_iris <- rfsrc(Species ~., data = iris)
data(rfsrc_iris, package="ggRandomForests")

gg_dta <- gg_variable(rfsrc_iris)
plot(gg_dta, xvar="Sepal.Width")
plot(gg_dta, xvar="Sepal.Length")

## !! TODO !! this needs to be corrected
plot(gg_dta, xvar=rfsrc_iris$xvar.names,
      panel=TRUE, se=FALSE)

## -----
## regression
## -----
## ----- air quality data
#rfsrc_airq <- rfsrc(Ozone ~ ., data = airquality)
data(rfsrc_airq, package="ggRandomForests")
gg_dta <- gg_variable(rfsrc_airq)

# an ordinal variable
gg_dta[, "Month"] <- factor(gg_dta[, "Month"])

plot(gg_dta, xvar="Wind")
plot(gg_dta, xvar="Temp")
plot(gg_dta, xvar="Solar.R")

plot(gg_dta, xvar=c("Solar.R", "Wind", "Temp", "Day"), panel=TRUE)

plot(gg_dta, xvar="Month", notch=TRUE)

## ----- motor trend cars data
#rfsrc_mtcars <- rfsrc(mpg ~ ., data = mtcars)
data(rfsrc_mtcars, package="ggRandomForests")
gg_dta <- gg_variable(rfsrc_mtcars)

# mtcars$cyl is an ordinal variable
gg_dta$cyl <- factor(gg_dta$cyl)
gg_dta$am <- factor(gg_dta$am)
gg_dta$vs <- factor(gg_dta$vs)
gg_dta$gear <- factor(gg_dta$gear)
gg_dta$carb <- factor(gg_dta$carb)

plot(gg_dta, xvar="cyl")

# Others are continuous

```

```

plot(gg_dta, xvar="disp")
plot(gg_dta, xvar="hp")
plot(gg_dta, xvar="wt")

# panel
plot(gg_dta,xvar=c("disp","hp", "drat", "wt", "qsec"), panel=TRUE)
plot(gg_dta, xvar=c("cyl", "vs", "am", "gear", "carb") ,panel=TRUE)

## ----- Boston data

## -----
## survival examples
## -----
## ----- veteran data
## survival
data(veteran, package = "randomForestSRC")
rfsrc_veteran <- rfsrc(Surv(time,status)~., veteran, nsplit = 10,
                      ntree = 100)

# get the 1 year survival time.
gg_dta <- gg_variable(rfsrc_veteran, time=90)

# Generate variable dependance plots for age and diagtime
plot(gg_dta, xvar = "age")
plot(gg_dta, xvar = "diagtime")

# Generate coplots
plot(gg_dta, xvar = c("age", "diagtime"), panel=TRUE)

# If we want to compare survival at different time points, say 30, 90 day
# and 1 year
gg_dta <- gg_variable(rfsrc_veteran, time=c(30, 90, 365))

# Generate variable dependance plots for age and diagtime
plot(gg_dta, xvar = "age")
plot(gg_dta, xvar = "diagtime")

# Generate coplots
plot(gg_dta, xvar = c("age", "diagtime"), panel=TRUE)

## ----- pbc data

## End(Not run)

```

plot.gg_vimp

Plot a `gg_vimp` object, extracted variable importance of a `rfsrc` object

Description

Plot a `gg_vimp` object, extracted variable importance of a `rfsrc` object

Usage

```
## S3 method for class 'gg_vimp'
plot(x, relative, lbls, ...)
```

Arguments

<code>x</code>	<code>gg_vimp</code> object created from a <code>rfsrc</code> object
<code>relative</code>	should we plot vimp or relative vimp. Defaults to vimp.
<code>lbls</code>	A vector of alternative variable labels. Item names should be the same as the variable names.
<code>...</code>	optional arguments passed to <code>gg_vimp</code> if necessary

Value

ggplot object

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.

Ishwaran H. and Kogalur U.B. (2013). Random Forests for Survival, Regression and Classification (RF-SRC), R package version 1.4.

See Also

[gg_vimp](#)

Examples

```
## Not run:
## -----
## classification example
## -----
## ----- iris data
# rfsrc_iris <- rfsrc(Species ~ ., data = iris)
data(rfsrc_iris, package="ggRandomForests")
gg_dta <- gg_vimp(rfsrc_iris)
plot(gg_dta)

## -----
## regression example
## -----
## ----- air quality data
# rfsrc_airq <- rfsrc(Ozone ~ ., airquality)
data(rfsrc_airq, package="ggRandomForests")
```

```
gg_dta <- gg_vimp(rfsrc_airq)
plot(gg_dta)

## ----- Boston data
data(rfsrc_boston, package="ggRandomForests")
gg_dta <- gg_vimp(rfsrc_boston)
plot(gg_dta)

## ----- mtcars data
data(rfsrc_mtcars, package="ggRandomForests")
gg_dta <- gg_vimp(rfsrc_mtcars)
plot(gg_dta)

## -----
## survival example
## -----
## ----- veteran data
data(rfsrc_veteran, package="ggRandomForests")
gg_dta <- gg_vimp(rfsrc_veteran)
plot(gg_dta)

## ----- pbc data
data(rfsrc_pbc, package="ggRandomForests")
gg_dta <- gg_vimp(rfsrc_pbc)
plot(gg_dta)

## End(Not run)
```

print.gg_minimal_depth

Print a [gg_minimal_depth](#) object.

Description

Print a [gg_minimal_depth](#) object.

Usage

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

Arguments

x a [gg_minimal_depth](#) object.
... optional arguments

Examples

```

## Not run:
## -----
## classification example
## -----
## You can build a randomForest
rfsrc_iris <- rfsrc(Species ~ ., data = iris)
varsel_iris <- var.select(rfsrc_iris)

# Get a data.frame containing minimaldepth measures
gg_dta <- gg_minimal_depth(varsel_iris)
print(gg_dta)

## -----
## regression example
## -----
# ... or load a cached randomForestSRC object
rfsrc_airq <- rfsrc(Ozone ~ ., data = airquality, na.action = "na.impute")
varsel_airq <- var.select(rfsrc_airq)

# Get a data.frame containing minimaldepth measures
gg_dta<- gg_minimal_depth(varsel_airq)
print(gg_dta)

# To nicely print a rfsrc::var.select output...
print(varsel_airq)

# ... or load a cached randomForestSRC object
data(Boston, package="MASS")
rfsrc_boston <- randomForestSRC::rfsrc(medv~., Boston)

varsel_boston <- var.select(rfsrc_boston)

# Get a data.frame containing minimaldepth measures
gg_dta<- gg_minimal_depth(varsel_boston)
print(gg_dta)

# To nicely print a rfsrc::var.select output...
print(varsel_boston)

## End(Not run)

```

quantile_pts

Find points evenly distributed along the vectors values.

Description

This function finds point values from a vector argument to produce groups intervals. Setting groups=2 will return three values, the two end points, and one mid point (at the median value

of the vector).

The output can be passed directly into the breaks argument of the cut function for creating groups for coplots.

Usage

```
quantile_pts(object, groups, intervals = FALSE)
```

Arguments

object	vector object of values.
groups	how many points do we want
intervals	should we return the raw points or intervals to be passed to the cut function

Value

vector of groups+1 cut point values.

See Also

[cut](#) [gg_partial_coplot](#)

Examples

```
data(Boston, package="MASS")
rfsrc_boston <- randomForestSRC::rfsrc(medv~., Boston)

# To create 6 intervals, we want 7 points.
# quantile_pts will find balanced intervals
rm_pts <- quantile_pts(rfsrc_boston$xvar$rm, groups=6, intervals=TRUE)

# Use cut to create the intervals
rm_grp <- cut(rfsrc_boston$xvar$rm, breaks=rm_pts)

summary(rm_grp)
```

shift	<i>lead function to shift by one (or more).</i>
-------	---

Description

lead function to shift by one (or more).

Usage

```
shift(x, shift_by = 1)
```

Arguments

x	a vector of values
shift_by	an integer of length 1, giving the number of positions to lead (positive) or lag (negative) by

Details

Lead and lag are useful for comparing values offset by a constant (e.g. the previous or next value)

Taken from: <http://ctszkin.com/2012/03/11/generating-a-laglead-variables/>

This function allows me to remove the `dplyr::lead` depends. Still suggest for vignettes though.

Examples

```
d<-data.frame(x=1:15)
#generate lead variable
d$df_lead2<-ggRandomForests:::shift(d$x,2)
#generate lag variable
d$df_lag2<-ggRandomForests:::shift(d$x,-2)
```

surface_matrix	<i>Construct a set of (x, y, z) matrices for surface plotting a gg_partial_coplot object</i>
----------------	--

Description

Construct a set of (x, y, z) matrices for surface plotting a `gg_partial_coplot` object

Usage

```
surface_matrix(dta, xvar)
```

Arguments

dta	a <code>gg_partial_coplot</code> object containing at least 3 numeric columns of data
xvar	a vector of 3 column names from the data object, in (x, y, z) order

Details

To create a surface plot, the `plot3D::surf3D` function expects 3 matrices of `n.x` by `n.y`. Take the `p+1` by `n` `gg_partial_coplot` object, and extract and construct the x, y and z matrices from the provided `xvar` column names.

Examples

```

## Not run:
## From vignette(randomForestRegression, package="ggRandomForests")
data(Boston, package="MASS")
rfsrc_boston <- randomForestSRC::rfsrc(medv~., Boston)

varsel_boston <- var.select(rfsrc_boston)

rm_pts <- quantile_pts(rfsrc_boston$xvar$rm,
  groups = 9,
  intervals = TRUE)

partial_boston_surf <- lapply(rm_pts, function(ct) {
  rfsrc_boston$xvar$rm <- ct
  randomForestSRC::plot.variable(
    rfsrc_boston,
    xvar.names = "lstat",
    time = 1,
    npts = 10,
    show.plots = FALSE,
    partial = TRUE
  )
})

# Instead of groups, we want the raw rm point values,
# To make the dimensions match, we need to repeat the values
# for each of the 50 points in the lstat direction
rm.tmp <- do.call(c,lapply(rm_pts,
  function(grp) {rep(grp,
    length(partial_boston_surf))}))

# Convert the list of plot.variable output to
partial_surf <- do.call(rbind,lapply(partial_boston_surf, gg_partial))

# attach the data to the gg_partial_coplot
partial_surf$rm <- rm.tmp

# Transform the gg_partial_coplot object into a list of three named matrices
# for surface plotting with plot3D::surf3D
srf <- surface_matrix(partial_surf, c("lstat", "rm", "yhat"))

## End(Not run)

## Not run:
# surf3D is in the plot3D package.
library(plot3D)
# Generate the figure.
surf3D(x=srf$x, y=srf$y, z=srf$z, col=topo.colors(10),
  colkey=FALSE, border = "black", bty="b2",
  shade = 0.5, expand = 0.5,
  lighting = TRUE, lphi = -50,
  xlab="Lower Status", ylab="Average Rooms", zlab="Median Value"

```

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surface_matrix

)

End(Not run)

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