

Package ‘model4you’

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Title Stratified and Personalised Models Based on Model-Based Trees
and Forests

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Description Model-based trees for subgroup analyses in clinical trials and model-based forests for the estimation and prediction of personalised treatment effects (personalised models). Currently partitioning of linear models, `lm()`, generalised linear models, `glm()`, and Weibull models, `survreg()`, is supported. Advanced plotting functionality is supported for the trees and a test for parameter heterogeneity is provided for the personalised models. For details on model-based trees for subgroup analyses see Seibold, Zeileis and Hothorn (2016) <[doi:10.1515/ijb-2015-0032](https://doi.org/10.1515/ijb-2015-0032)>; for details on model-based forests for estimation of individual treatment effects see Seibold, Zeileis and Hothorn (2017) <[doi:10.1177/0962280217693034](https://doi.org/10.1177/0962280217693034)>.

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Imports sandwich, stats, methods, ggplot2, Formula, gridExtra,
survival

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.add_modelinfo	<i>Add model information to a personalised-model-ctree</i>
----------------	--

Description

For internal use.

Usage

```
.add_modelinfo(x, nodeids, data, model, coefffun)
```

Arguments

x	constparty object.
nodeids	node ids, usually the terminal ids.
data	data.
model	model.
coefffun	function that takes the model object and returns the coefficients. Useful when coef() does not return all coefficients (e.g. survreg).

Value

tree with added info. Class still to be added.

`.modelfit`*Fit function when model object is given*

Description

Use update function to refit model and extract info such as `coef`, `logLik` and `estfun`.

Usage

```
.modelfit(model, data, coeffun = coef, weights, control, parm = NULL)
```

Arguments

<code>model</code>	model object.
<code>data</code>	data.
<code>coeffun</code>	function that takes the model object and returns the coefficients. Useful when <code>coef()</code> does not return all coefficients (e.g. <code>survreg</code>).
<code>weights</code>	weights.
<code>control</code>	control options from <code>ctree_control</code> .
<code>parm</code>	which parameters should be used for instability test?

Value

A function returning a list of

<code>coefficients</code>	<code>coef</code> .
<code>objfun</code>	<code>logLik</code> .
<code>object</code>	the model object.
<code>converged</code>	Did the model converge?
<code>estfun</code>	<code>estfun</code> .

<code>.prepare_args</code>	<i>Prepare input for ctree/cforest from input of pmtree/pmforest</i>
----------------------------	--

Description

Prepare input for ctree/cforest from input of pmtree/pmforest

Usage

```
.prepare_args(model, data, zformula, control, ...)
```

Arguments

<code>model</code>	model.
<code>data</code>	an optional data frame.
<code>zformula</code>	ormula describing which variable should be used for partitioning.
<code>control</code>	ontrol parameters, see ctree_control .
<code>...</code>	other arguments.

Value

args to be passed to ctree/cforest.

<code>binomial_glm_plot</code>	<i>Plot for a given logistic regression model (glm with binomial family) with one binary covariate.</i>
--------------------------------	---

Description

Can be used on its own but is also useable as plotfun in [node_pmterminal](#).

Usage

```
binomial_glm_plot(
  mod,
  data = NULL,
  plot_data = FALSE,
  theme = theme_classic(),
  ...
)
```

Arguments

mod	A model of class glm with binomial family.
data	optional data frame. If NULL the data stored in mod is used.
plot_data	should the data in form of a mosaic type plot be plotted?
theme	A ggplot2 theme.
...	ignored at the moment.

Examples

[illegible]

coeftable.survreg	<i>Table of coefficients for survreg model</i>
-------------------	--

Description

This function is mostly useful for plotting a pmtree. The generic plotting does not show the estimate and confidence interval of the scale parameter. This one does.

Usage

```
coeftable.survreg(model, confint = TRUE, digits = 2, intree = FALSE)
```

Arguments

model	model of class survreg
confint	should a confidence interval be computed? Default: TRUE
digits	integer, used for formatting numbers. Default: 2
intree	is the table plotted within a tree? Default: FALSE

Value

None.

Examples

```
if(require("survival") & require("TH.data")) {
  ## Load data
  data(GBSG2, package = "TH.data")

  ## Weibull model
  bmod <- survreg(Surv(time, cens) ~ horTh, data = GBSG2, model = TRUE)

  ## Coefficient table
  grid.newpage()
  coeftable.survreg(bmod)

  ## partitioned model
  tr <- pmtree(bmod)

  ## plot
  plot(tr, terminal_panel = node_pmterminal(tr, plotfun = survreg_plot,
    confint = TRUE, coeftable = coeftable.survreg))
}
```

coxph_plot

Survival plot for a given coxph model with one binary covariate.

Description

Can be used on its own but is also useable as plotfun in [node_pmterminal](#).

Usage

```
coxph_plot(mod, data = NULL, theme = theme_classic(), yrange = NULL)
```

Arguments

mod	A model of class coxph.
data	optional data frame. If NULL the data stored in mod is used.
theme	A ggplot2 theme.
yrange	Range of the y variable to be used for plotting. If NULL it will be 0 to max(y).

Examples

```
if(require("survival")) {
  coxph_plot(coxph(Surv(futime, fustat) ~ factor(rx), ovarian))
}
```

lm_plot

Density plot for a given lm model with one binary covariate.

Description

Can be used on its own but is also useable as plotfun in [node_pmterminal](#).

Usage

```
lm_plot(
  mod,
  data = NULL,
  densest = FALSE,
  theme = theme_classic(),
  yrange = NULL
)
```

Arguments

mod	A model of class lm.
data	optional data frame. If NULL the data stored in mod is used.
densest	should additional to the model density kernel density estimates (see geom_density) be computed?
theme	A ggplot2 theme.
yrange	Range of the y variable to be used for plotting. If NULL the range in the data will be used.

Details

In case of an offset, the value of the offset variable will be set to the median of the values in the data.

Examples

```
## example taken from ?lm
ctl <- c(4.17,5.58,5.18,6.11,4.50,4.61,5.17,4.53,5.33,5.14)
trt <- c(4.81,4.17,4.41,3.59,5.87,3.83,6.03,4.89,4.32,4.69)
group <- gl(2, 10, 20, labels = c("Ctl","Trt"))
weight <- c(ctl, trt)
data <- data.frame(weight, group)
lm.D9 <- lm(weight ~ group, data = data)
lm_plot(lm.D9)

## example taken from ?glm (modified version)
data(anorexia, package = "MASS")
anorexia$treatment <- factor(anorexia$Treat != "Cont")
anorex.1 <- glm(Postwt ~ treatment + offset(Prewt),
               family = gaussian, data = anorexia)
lm_plot(anorex.1)
```

logLik.pmtree

Extract log-Likelihood

Description

Extract sum of log-Likelihood contributions of all terminal nodes. By default the degrees of freedom from the models are used but optionally degrees of freedom for splits can be incorporated.

Usage

```
## S3 method for class 'pmtree'
logLik(object, dfsplit = 0, newdata = NULL, weights = NULL, perm = NULL, ...)
```

Arguments

object	pmtree object.
dfsplitted	degrees of freedom per selected split.
newdata	an optional new data frame for which to compute the sum of objective functions.
weights	weights.
perm	the number of permutations performed (see varimp).
...	ignored.

Value

Returns an object of class [logLik](#).

See Also

[objfun.pmtree](#) for the sum of contributions to the objective function (not the same when partitioning linear models [lm](#))

node_pmterminal	<i>Panel-Generator for Visualization of pmtrees</i>
-----------------	---

Description

The plot method for party and constparty objects are rather flexible and can be extended by panel functions. The pre-defined panel-generating function of class `grapcon_generator` for pmtrees is documented here.

Usage

```
node_pmterminal(
  obj,
  coeftable = TRUE,
  digits = 2,
  confint = TRUE,
  plotfun,
  nid = function(node) paste0(nam[id_node(node)], ", n = ", node$info$nobs),
  ...
)
```

Arguments

obj	an object of class party.
coeftable	logical or function. If logical: should a table with coefficients be added to the plot (TRUE/FALSE)? If function: A function comparable to coeftable.survreg .
digits	integer, used for formatting numbers.
confint	Should a confidence interval be computed.

plotfun	Plotting function to be used. Needs to be of format function(mod, data) where mod is the model object. See examples for more details.
nid	function to retrieve info on what is plotted as node ids.
...	arguments passed on to plotfun.

Examples

```

if(require("survival")) {
  ## compute survreg model
  mod_surv <- survreg(Surv(futime, fustat) ~ factor(rx), ovarian,
    dist = 'weibull')
  survreg_plot(mod_surv)

  ## partition model and plot
  tr_surv <- pmtree(mod_surv)
  plot(tr_surv, terminal_panel = node_pmterminal(tr_surv, plotfun = survreg_plot,
    confint = TRUE))
}

if(require("survival") & require("TH.data")) {
  ## Load data
  data(GBSG2, package = "TH.data")

  ## Weibull model
  bmod <- survreg(Surv(time, cens) ~ horTh, data = GBSG2, model = TRUE)

  ## Coefficient table
  grid.newpage()
  coeftable.survreg(bmod)

  ## partitioned model
  tr <- pmtree(bmod)

  ## plot with specific coeftable
  plot(tr, terminal_panel = node_pmterminal(tr, plotfun = survreg_plot,
    confint = TRUE, coeftable = coeftable.survreg))
}

```

objfun

Objective function

Description

Get the contributions of an objective function. For [glm](#) these are the (weighted) log-likelihood contributions, for [lm](#) the negative (weighted) squared error.

Usage

```
objfun(x, ...)

## S3 method for class 'survreg'
objfun(x, newdata = NULL, weights = NULL, ...)

## S3 method for class 'lm'
objfun(x, newdata = NULL, weights = NULL, ...)

## S3 method for class 'glm'
objfun(x, newdata = NULL, weights = NULL, log = TRUE, ...)
```

Arguments

x	model object.
...	further arguments passed on to objfun methods.
newdata	optional. New data frame. Can be useful for model evaluation / benchmarking.
weights	optional. Prior weights. See glm or lm .
log	should the log-Likelihood contributions or the Likelihood contributions be returned?

Value

vector of objective function contributions.

Examples

```
## Example taken from ?stats::glm
## Dobson (1990) Page 93: Randomized Controlled Trial :
counts <- c(18,17,15,20,10,20,25,13,12)
outcome <- gl(3,1,9)
treatment <- gl(3,3)
print(d.AD <- data.frame(treatment, outcome, counts))
glm.D93 <- glm(counts ~ outcome + treatment, family = poisson())
logLik_contributions <- objfun(glm.D93)
sum(logLik_contributions)
logLik(glm.D93)

if(require("survival")) {
  x <- survreg(Surv(futime, fustat) ~ rx, ovarian, dist = "weibull")
  newdata <- ovarian[3:5, ]

  sum(objfun(x))
  x$loglik

  objfun(x, newdata = newdata)
}
```

objfun.pmodel_identity

Objective function of personalised models

Description

Get the contributions of an objective function (e.g. likelihood contributions) and the sum thereof (e.g. log-Likelihood).

Usage

```
## S3 method for class 'pmodel_identity'
objfun(x, ...)

## S3 method for class 'pmodel_identity'
logLik(object, add_df = 0, ...)
```

Arguments

x, object	object of class pmodel_identity (obtained by pmodel(..., fun = identity)).
...	additional parameters passed on to objfun .
add_df	it is not very clear what the degrees of freedom are in personalised models. With this argument you can add/subtract degrees of freedom at your convenience. Default is 0 which means adding up the degrees of freedom of all individual models. For examples see pmodel .

objfun.pmtree

Objective function of a given pmtree

Description

Returns the contributions to the objective function or the sum thereof (if sum = TRUE).

Usage

```
## S3 method for class 'pmtree'
objfun(x, newdata = NULL, weights = NULL, perm = NULL, sum = FALSE, ...)
```

Arguments

x	pmtree object.
newdata	an optional new data frame for which to compute the sum of objective functions.
weights	weights.
perm	the number of permutations performed (see varimp).
sum	should the sum of objective functions be computed.
...	passed on to predict.party .

Note that `objfun.pmtree(x, sum = TRUE)` is much faster than `sum(objfun.pmtree(x))`.

Value

objective function or the sum thereof

Examples

```
## generate data
set.seed(2)
n <- 1000
trt <- factor(rep(1:2, each = n/2))
age <- sample(40:60, size = n, replace = TRUE)
eff <- -1 + I(trt == 2) + 1 * I(trt == 2) * I(age > 50)
expit <- function(x) 1/(1 + exp(-x))
success <- rbinom(n = n, size = 1, prob = expit(eff))
dat <- data.frame(success, trt, age)

## compute base model
bmod1 <- glm(success ~ trt, data = dat, family = binomial)

## compute tree
(tr1 <- pmtree(bmod1, data = dat))

## compute log-Likelihood
logLik(tr1)
objfun(tr1, newdata = dat, sum = TRUE)
objfun(tr1, sum = TRUE)

## log-Likelihood contributions of first
## 5 observations
nd <- dat[1:5, ]
objfun(tr1, newdata = nd)
```

one_factor

Check if model has only one factor covariate.

Description

See <https://stackoverflow.com/questions/50504386/check-that-model-has-only-one-factor-covariate/50514499#50514499>

Usage

```
one_factor(object)
```

Arguments

```
object          model.
```

Value

Returns TRUE if model has a single factor covariate, FALSE otherwise.

pmforest	<i>Compute model-based forest from model.</i>
----------	---

Description

Input a parametric model and get a forest.

Usage

```
pmforest(
  model,
  data = NULL,
  zformula = ~.,
  ntree = 500L,
  perturb = list(replace = FALSE, fraction = 0.632),
  mtry = NULL,
  applyfun = NULL,
  cores = NULL,
  control = ctree_control(teststat = "quad", testtype = "Univ", mincriterion = 0,
    saveinfo = FALSE, lookahead = TRUE, ...),
  trace = FALSE,
  ...
)

## S3 method for class 'pmforest'
gettree(object, tree = 1L, saveinfo = TRUE, coeffun = coef, ...)
```

Arguments

model	a model object. The model can be a parametric model with a single binary covariate.
data	data. If NULL the data from the model object are used.
zformula	formula describing which variable should be used for partitioning. Default is to use all variables in data that are not in the model (i.e. $\sim .$).
ntree	number of trees.

perturb	a list with arguments replace and fraction determining which type of resampling with replace = TRUE referring to the n-out-of-n bootstrap and replace = FALSE to sample splitting. fraction is the number of observations to draw without replacement.
mtry	number of input variables randomly sampled as candidates at each node (Default NULL corresponds to ceiling(sqrt(nvar))). Bagging, as special case of a random forest without random input variable sampling, can be performed by setting mtry either equal to Inf or equal to the number of input variables.
applyfun	see cforest .
cores	see cforest .
control	control parameters, see ctree_control .
trace	a logical indicating if a progress bar shall be printed while the forest grows.
...	additional parameters passed on to model fit such as weights.
object	an object returned by pmforest.
tree	an integer, the number of the tree to extract from the forest.
saveinfo	logical. Should the model info be stored in terminal nodes?
coeffun	function that takes the model object and returns the coefficients. Useful when coef() does not return all coefficients (e.g. survreg).

Value

cforest object

See Also

[gettree](#)

Examples

```
library("model4you")

if(require("mvtnorm") & require("survival")) {

  ## function to simulate the data
  sim_data <- function(n = 500, p = 10, beta = 3, sd = 1){

    ## treatment
    lev <- c("C", "A")
    a <- rep(factor(lev, labels = lev, levels = lev), length = n)

    ## correlated z variables
    sigma <- diag(p)
    sigma[sigma == 0] <- 0.2
    ztemp <- rmvnorm(n, sigma = sigma)
    z <- (pnorm(ztemp) * 2 * pi) - pi
    colnames(z) <- paste0("z", 1:ncol(z))
    z1 <- z[,1]
```

```

## outcome
y <- 7 + 0.2 * (a %in% "A") + beta * cos(z1) * (a %in% "A") + rnorm(n, 0, sd)

data.frame(y = y, a = a, z)
}

## simulate data
set.seed(123)
beta <- 3
ntrain <- 500
ntest <- 50
simdata <- simdata_s <- sim_data(p = 5, beta = beta, n = ntrain)
tsimdata <- tsimdata_s <- sim_data(p = 5, beta = beta, n = ntest)
simdata_s$cens <- rep(1, ntrain)
tsimdata_s$cens <- rep(1, ntest)

## base model
basemodel_lm <- lm(y ~ a, data = simdata)

## forest
frst_lm <- pmforest(basemodel_lm, ntree = 20,
                    perturb = list(replace = FALSE, fraction = 0.632),
                    control = ctree_control(mincriterion = 0))

## personalised models
# (1) return the model objects
pmodels_lm <- pmodel(x = frst_lm, newdata = tsimdata, fun = identity)
class(pmodels_lm)
# (2) return coefficients only (default)
coefs_lm <- pmodel(x = frst_lm, newdata = tsimdata)

# compare predictive objective functions of personalised models versus
# base model
sum(objfun(pmodels_lm)) # -RSS personalised models
sum(objfun(basemodel_lm, newdata = tsimdata)) # -RSS base model

if(require("ggplot2")) {
  ## dependence plot
  dp_lm <- cbind(coefs_lm, tsimdata)
  ggplot(tsimdata) +
    stat_function(fun = function(z1) 0.2 + beta * cos(z1),
                  aes(color = "true treatment\neffect")) +
    geom_point(data = dp_lm,
               aes(y = aA, x = z1, color = "estimates lm"),
               alpha = 0.5) +
    ylab("treatment effect") +
    xlab("patient characteristic z1")
}
}

```

pmodel

*Personalised model***Description**

Compute personalised models from cforest object.

Usage

```
pmodel(
  x = NULL,
  model = NULL,
  newdata = NULL,
  OOB = TRUE,
  fun = coef,
  return_attr = c("modelcall", "data", "similarity")
)
```

Arguments

x	cforest object or matrix of weights.
model	model object. If NULL the model in x\$info\$model is used.
newdata	new data. If NULL cforest learning data is used. Ignored if x is a matrix.
OOB	In case of using the learning data, should patient similarities be computed out of bag?
fun	function to apply on the personalised model before returning. The default coef returns a matrix of personalised coefficients. For returning the model objects use identity.
return_attr	which attributes to add to the object returned. If it contains "modelcall" the call of the base model is returned, if it contains "data" the data, and if it contains "similarity" the matrix of similarity weights is added.

Value

depends on fun.

Examples

```
library("model4you")

if(require("mvtnorm") & require("survival")) {

  ## function to simulate the data
  sim_data <- function(n = 500, p = 10, beta = 3, sd = 1){

    ## treatment
```

```

lev <- c("C", "A")
a <- rep(factor(lev, labels = lev, levels = lev), length = n)

## correlated z variables
sigma <- diag(p)
sigma[sigma == 0] <- 0.2
ztemp <- rmvnorm(n, sigma = sigma)
z <- (pnorm(ztemp) * 2 * pi) - pi
colnames(z) <- paste0("z", 1:ncol(z))
z1 <- z[,1]

## outcome
y <- 7 + 0.2 * (a %in% "A") + beta * cos(z1) * (a %in% "A") + rnorm(n, 0, sd)

data.frame(y = y, a = a, z)
}

## simulate data
set.seed(123)
beta <- 3
ntrain <- 500
ntest <- 50
simdata <- simdata_s <- sim_data(p = 5, beta = beta, n = ntrain)
tsimdata <- tsimdata_s <- sim_data(p = 5, beta = beta, n = ntest)
simdata_s$cens <- rep(1, ntrain)
tsimdata_s$cens <- rep(1, ntest)

## base model
basemodel_lm <- lm(y ~ a, data = simdata)

## forest
frst_lm <- pmforest(basemodel_lm, ntree = 20,
                    perturb = list(replace = FALSE, fraction = 0.632),
                    control = ctree_control(mincriterion = 0))

## personalised models
# (1) return the model objects
pmodels_lm <- pmodel(x = frst_lm, newdata = tsimdata, fun = identity)
class(pmodels_lm)
# (2) return coefficients only (default)
coefs_lm <- pmodel(x = frst_lm, newdata = tsimdata)

# compare predictive objective functions of personalised models versus
# base model
sum(objfun(pmodels_lm)) # -RSS personalised models
sum(objfun(basemodel_lm, newdata = tsimdata)) # -RSS base model

if(require("ggplot2")) {
  ## dependence plot
  dp_lm <- cbind(coefs_lm, tsimdata)
  ggplot(tsimdata) +
    stat_function(fun = function(z1) 0.2 + beta * cos(z1),

```

```

      aes(color = "true treatment\neffect")) +
    geom_point(data = dp_lm,
      aes(y = aA, x = z1, color = "estimates lm"),
      alpha = 0.5) +
    ylab("treatment effect") +
    xlab("patient characteristic z1")
  }
}

```

pmtest

*Test if personalised models improve upon base model.***Description**

This is a rudimentary test if there is heterogeneity in the model parameters. The null-hypothesis is: the base model is the correct model.

Usage

```

pmtest(forest, pmodels = NULL, data = NULL, B = 100)

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

```

Arguments

forest	pmforest object.
pmodels	pmodel_identity object (pmodel(..., fun = identity)).
data	data.
B	number of bootstrap samples.
x	object of class heterogeneity_test.
...	ignored.

Value

list where the first element is the p-value und the second element is a data.frame with all neccessary infos to compute the p-value.

The test statistic is the difference in objective function between the base model and the personalised models. To compute the distribution under the Null we draw parametric bootstrap samples from the base model. For each bootstrap sample we again compute the difference in objective function between the base model and the personalised models. If the difference in the original data is greater than the difference in the bootstrap samples, we reject the null-hypothesis.

Examples

```
## Not run:
set.seed(123)
n <- 160
trt <- factor(rep(0:1, each = n/2))
y <- 4 + (trt == 1) + rnorm(n)
z <- matrix(rnorm(n * 2), ncol = 2)

dat <- data.frame(y, trt, z)

mod <- lm(y ~ trt, data = dat)

## Note that ntree should usually be higher
frst <- pmforest(mod, ntree = 20)
pmods <- pmodel(frst, fun = identity)

## Note that B should be at least 100
## The low B is just for demonstration
## purposes.
tst <- pmtest(forest = frst,
              pmodels = pmods,
              B = 10)

tst$pvalue
tst
plot(tst)

## End(Not run)
```

pmtree

Compute model-based tree from model.

Description

Input a parametric model and get a model-based tree.

Usage

```
pmtree(
  model,
  data = NULL,
  zformula = ~.,
  control = ctree_control(),
  coeffun = coef,
  ...
)
```



```

summary(tr2[[5]]$data$horTh)
}

if(require("psychotools")) {
  data("MathExam14W", package = "psychotools")

  ## scale points achieved to [0, 100] percent
  MathExam14W$tests <- 100 * MathExam14W$tests/26
  MathExam14W$pcorrect <- 100 * MathExam14W$nsolved/13

  ## select variables to be used
  MathExam <- MathExam14W[, c("pcorrect", "group", "tests", "study",
                              "attempt", "semester", "gender")]

  ## compute base model
  bmod_math <- lm(pcorrect ~ group, data = MathExam)
  lm_plot(bmod_math, densest = TRUE)

  ## compute tree
  (tr_math <- pmtree(bmod_math, control = ctree_control(maxdepth = 2)))
  plot(tr_math, terminal_panel = node_pmterminal(tr_math, plotfun = lm_plot,
                                                  confint = FALSE))
  plot(tr_math, terminal_panel = node_pmterminal(tr_math, plotfun = lm_plot,
                                                  densest = TRUE,
                                                  confint = TRUE))

  ## predict
  newdat <- MathExam[1:5, ]

  # terminal nodes
  (nodes <- predict(tr_math, type = "node", newdata = newdat))

  # response
  (pr <- predict(tr_math, type = "pass", newdata = newdat))

  # response including confidence intervals, see ?predict.lm
  (pr1 <- predict(tr_math, type = "pass", newdata = newdat,
                  predict_args = list(interval = "confidence")))
}

```

predict.pmtree

pmtree predictions

Description

Compute predictions from pmtree object.

Usage

```
## S3 method for class 'pmtree'
predict(
  object,
  newdata = NULL,
  type = "node",
  predict_args = list(),
  perm = NULL,
  ...
)
```

Arguments

object	pmtree object.
newdata	an optional data frame in which to look for variables with which to predict, if omitted, object\$data is used.
type	character denoting the type of predicted value. The terminal node is returned for "node". If type = "pass" the model predict method is used and arguments can be passed to it via predict_args. If type = "coef" the the model coefficients are returned.
predict_args	If type = "pass" arguments can be passed on to the model predict function.
perm	an optional character vector of variable names (or integer vector of variable location in newdata). Splits of nodes with a primary split in any of these variables will be permuted (after dealing with surrogates). Note that surrogate split in the perm variables will no be permuted.
...	passed on to predict.party (e.g. perm).

Value

predictions

Examples

```
if(require("psychotools")) {
  data("MathExam14W", package = "psychotools")

  ## scale points achieved to [0, 100] percent
  MathExam14W$tests <- 100 * MathExam14W$tests/26
  MathExam14W$pcorrect <- 100 * MathExam14W$nsolved/13

  ## select variables to be used
  MathExam <- MathExam14W[, c("pcorrect", "group", "tests", "study",
                             "attempt", "semester", "gender")]

  ## compute base model
  bmod_math <- lm(pcorrect ~ group, data = MathExam)
  lm_plot(bmod_math, densest = TRUE)
```

```

## compute tree
(tr_math <- pmtree(bmod_math, control = ctree_control(maxdepth = 2)))
plot(tr_math, terminal_panel = node_pmterminal(tr_math, plotfun = lm_plot,
                                              confint = FALSE))
plot(tr_math, terminal_panel = node_pmterminal(tr_math, plotfun = lm_plot,
                                              densest = TRUE,
                                              confint = TRUE))

## predict
newdat <- MathExam[1:5, ]

# terminal nodes
(nodes <- predict(tr_math, type = "node", newdata = newdat))

# response
(pr <- predict(tr_math, type = "pass", newdata = newdat))

# response including confidence intervals, see ?predict.lm
(pr1 <- predict(tr_math, type = "pass", newdata = newdat,
               predict_args = list(interval = "confidence")))
}

```

print.pmtree

Methods for pmtree

Description

Print and summary methods for pmtree objects.

Usage

```

## S3 method for class 'pmtree'
print(
  x,
  node = NULL,
  FUN = NULL,
  digits = getOption("digits") - 4L,
  footer = TRUE,
  ...
)

## S3 method for class 'pmtree'
summary(object, node = NULL, ...)

## S3 method for class 'summary.pmtree'
print(x, digits = 4, ...)

## S3 method for class 'pmtree'
coef(object, node = NULL, ...)

```

Arguments

x	object.
node	node number, if any.
FUN	formatinfo function.
digits	number of digits.
footer	should footer be included?
...	further arguments passed on to print.party .
object	object.

Value

print

rss	<i>Residual sum of squares</i>
-----	--------------------------------

Description

Returns the sum of the squared residuals for a given object.

Usage

```
rss(object, ...)
```

Default S3 method:

```
rss(object, ...)
```

Arguments

object	model object.
...	passed on to specific methods.

Value

sum of the squared residuals.

Examples

```
## example from ?lm
ctl <- c(4.17,5.58,5.18,6.11,4.50,4.61,5.17,4.53,5.33,5.14)
trt <- c(4.81,4.17,4.41,3.59,5.87,3.83,6.03,4.89,4.32,4.69)
group <- gl(2, 10, 20, labels = c("Ctl","Trt"))
weight <- c(ctl, trt)
lm.D9 <- lm(weight ~ group)
rss(lm.D9)
```

survreg_plot	<i>Survival plot for a given survreg model with one binary covariate.</i>
--------------	---

Description

Can be used on its own but is also useable as plotfun in [node_pmterminal](#).

Usage

```
survreg_plot(mod, data = NULL, theme = theme_classic(), yrange = NULL)
```

Arguments

mod	A model of class survreg.
data	optional data frame. If NULL the data stored in mod is used.
theme	A ggplot2 theme.
yrange	Range of the y variable to be used for plotting. If NULL it will be 0 to max(y).

Examples

```
if(require("survival")) {
  survreg_plot(survreg(Surv(futime, fustat) ~ factor(rx), ovarian))
}
```

varimp.pmforest	<i>Variable Importance for pmforest</i>
-----------------	---

Description

See [varimp.cforest](#).

Usage

```
## S3 method for class 'pmforest'
varimp(
  object,
  nperm = 1L,
  OOB = TRUE,
  risk = function(x, ...) -objfun(x, sum = TRUE, ...),
  conditional = FALSE,
  threshold = 0.2,
  ...
)
```

Arguments

object	DESCRIPTION.
nperm	the number of permutations performed.
OOB	a logical determining whether the importance is computed from the out-of-bag sample or the learning sample (not suggested).
risk	the risk to be evaluated. By default the objective function (e.g. log-Likelihood) is used.
conditional	a logical determining whether unconditional or conditional computation of the importance is performed.
threshold	the value of the test statistic or 1 - p-value of the association between the variable of interest and a covariate that must be exceeded inorder to include the covariate in the conditioning scheme for the variable of interest (only relevant if conditional = TRUE).
...	passed on to objfun .

Value

A vector of 'mean decrease in accuracy' importance scores.

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