

Package ‘trtf’

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Title Transformation Trees and Forests

Version 0.3-9

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Description Recursive partytioning of transformation models with corresponding random forest for conditional transformation models as described in 'Transformation Forests' (Hothorn and Zeileis, 2021, <[doi:10.1080/10618600.2021.1872581](https://doi.org/10.1080/10618600.2021.1872581)>) and 'Top-Down Transformation Choice' (Hothorn, 2018, <[DOI:10.1177/1471082X17748081](https://doi.org/10.1177/1471082X17748081)>).

Depends mlt (>= 1.3-2), partykit (>= 1.2-1)

Imports Formula, sandwich, grid, stats, variables, libcoin, utils

Suggests survival, TH.data, coin

URL <http://ctm.R-forge.R-project.org>

License GPL-2

NeedsCompilation no

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 trtf-package

 General Information on the **trtf** Package

Description

The **trtf** package implements transformation trees and transformation forests as described in Hothorn and Zeileis (2017).

Example applications of transformation trees and forests can be replicated using `demo("applications")` and `demo("BMI")`. Figure 1 in Hothorn and Zeileis (2017) can be reproduced by `demo("QRF")`. Source code of simulation experiments is available in directory `trtf/inst/sim`.

A short talk introducing transformation trees and forests is available from <https://channel9.msdn.com/Events/useR-international-R-User-conferences/useR-International-R-User-2017-Conference/Transformation-Forests>.

Author(s)

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References

Torsten Hothorn and Achim Zeileis (2017). Transformation Forests. <https://arxiv.org/abs/1701.02110>.

 traforest

 Transformation Forests

Description

Partitioned and aggregated transformation models

Usage

```
traforest(object, parm = 1:length(coef(object)), reparam = NULL,
          update = TRUE, min_update = length(coef(object)) * 2,
          mltargs = list(), ...)
## S3 method for class 'traforest'
predict(object, newdata, mnewdata = data.frame(1), K = 20, q = NULL,
        type = c("weights", "node", "coef", "trafo", "distribution", "survivor", "density",
                 "logdensity", "hazard", "loghazard", "cumhazard", "quantile"),
        OOB = FALSE, simplify = FALSE, trace = FALSE, updatestart = FALSE,
        applyfun = NULL, cores = NULL, ...)
## S3 method for class 'traforest'
logLik(object, newdata, weights = NULL, OOB = FALSE, coef = NULL, ...)
```

Arguments

object	an object of class <code>ctm</code> or <code>mlt</code> specifying the abstract model to be partitioned.
parm	parameters of object those corresponding score is used for finding partitions.
reparm	optional matrix of contrasts for reparameterisation of the scores. <code>teststat = "quadratic"</code> is invariant to this operation but <code>teststat = "max"</code> might be more powerful for example when formulating an implicit into an explicit intercept term.
mltargs	arguments to <code>mlt</code> for fitting the transformation models.
update	logical, if TRUE, models and thus scores are updated in every node. If FALSE, the model and scores are computed once in the root node. The latter option is faster but less accurate.
min_update	number of observations necessary to refit the model in a node. If less observations are available, the parameters from the parent node will be reused.
newdata	an optional data frame of observations for the forest.
mnewdata	an optional data frame of observations for the model.
K	number of grid points to generate (in the absence of q).
q	quantiles at which to evaluate the model.
type	type of prediction or plot to generate.
OOB	compute out-of-bag predictions.
simplify	simplify predictions (if possible).
trace	a logical indicating if a progress bar shall be printed while the predictions are computed.
updatestart	try to be smart about starting values for computing predictions (experimental).
applyfun	an optional <code>lapply</code> -style function with arguments <code>function(X,FUN,...)</code> for looping over <code>newdata</code> . The default is to use the basic <code>lapply</code> function unless the <code>cores</code> argument is specified (see below).
cores	numeric. If set to an integer the <code>applyfun</code> is set to <code>mclapply</code> with the desired number of cores.
weights	an optional vector of weights.
coef	an optional matrix of precomputed coefficients for <code>newdata</code> (using <code>predict</code>). Helps to compute the coefficients once for later reuse (different weights, for example).
...	arguments to <code>cforest</code> , at least formula and data.

Details

Conditional inference trees are used for partitioning likelihood-based transformation models as described in Hothorn and Zeileis (2017). The method can be seen in action in Hothorn (2018) and the corresponding code is available as `demo("BMI")`.

Value

An object of class `traforest` with corresponding `logLik` and `predict` methods.

References

Torsten Hothorn and Achim Zeileis (2021). Predictive Distribution Modelling Using Transformation Forests. *Journal of Computational and Graphical Statistics*, doi: [10.1080/10618600.2021.1872581](https://doi.org/10.1080/10618600.2021.1872581).

Torsten Hothorn (2018). Top-Down Transformation Choice. *Statistical Modelling*, **3-4**, 274-298. doi: [10.1177/1471082X17748081](https://doi.org/10.1177/1471082X17748081).

Natalia Korepanova, Heidi Seibold, Verena Steffen and Torsten Hothorn (2019). Survival Forests under Test: Impact of the Proportional Hazards Assumption on Prognostic and Predictive Forests for ALS Survival. doi: [10.1177/0962280219862586](https://doi.org/10.1177/0962280219862586).

Examples

```
### Example: Personalised Medicine Using Partitioned and Aggregated Cox-Models
### A combination of <DOI:10.1177/0962280217693034> and <arXiv:1701.02110>
### based on infrastructure in the mlt R add-on package described in
### https://cran.r-project.org/web/packages/mlt.docreg/vignettes/mlt.pdf

library("trtf")
library("survival")
### German Breast Cancer Study Group 2 data set
data("GBSG2", package = "TH.data")

### set-up Cox model with overall treatment effect in hormonal therapy
yvar <- numeric_var("y", support = c(100, 2000), bounds = c(0, Inf))
By <- Bernstein_basis(yvar, order = 5, ui = "incre")
m <- ctm(response = By, shifting = ~ horTh, todistr = "MinExt", data = GBSG2)
GBSG2$y <- with(GBSG2, Surv(time, cens))

### overall log-hazard ratio
coef(cmod <- mlt(m, data = GBSG2))["horThyes"]
### roughly the same as
coef(coxph(y ~ horTh, data = GBSG2))

## Not run:

### estimate age-dependent Cox models (here ignoring all other covariates)
ctrl <- ctree_control(minsplit = 50, minbucket = 20, mincriterion = 0)
set.seed(290875)
tf_cmod <- traforest(m, formula = y ~ horTh | age, control = ctrl,
                    ntree = 50, mtry = 1, trace = TRUE, data = GBSG2)

### plot age-dependent treatment effects vs. overall treatment effect
nd <- data.frame(age = 30:70)
cf <- predict(tf_cmod, newdata = nd, type = "coef")
nd$logHR <- sapply(cf, function(x) x["horThyes"])
plot(logHR ~ age, data = nd, pch = 19, xlab = "Age", ylab = "log-Hazard Ratio")
abline(h = coef(cmod <- mlt(m, data = GBSG2))["horThyes"])
### treatment most beneficial in very young patients
### NOTE: scale of log-hazard ratios depends on
### corresponding baseline hazard function which _differs_
### across age; interpretation of positive / negative treatment effect is,
```

```

### however, save.

### mclapply doesn't work in Windows
if (.Platform$OS.type != "windows") {

  ### computing predictions: predicted coefficients
  cf1 <- predict(tf_cmod, newdata = nd, type = "coef")
  ### speedup with plenty of RAM and 4 cores
  cf2 <- predict(tf_cmod, newdata = nd, cores = 4, type = "coef")
  ### memory-efficient with low RAM and _one_ core
  cf3 <- predict(tf_cmod, newdata = nd, cores = 4, applyfun = lapply, type = "coef")
  all.equal(cf1, cf2)
  all.equal(cf1, cf3)

}

## End(Not run)

```

trafotree

*Transformation Trees***Description**

Partitioned transformation models

Usage

```

trafotree(object, parm = 1:length(coef(object)), reparm = NULL,
          min_update = length(coef(object)) * 2,
          mltargs = list(), ...)
## S3 method for class 'trafotree'
predict(object, newdata, K = 20, q = NULL,
        type = c("node", "coef", "trafo", "distribution", "survivor", "density",
                 "logdensity", "hazard", "loghazard", "cumhazard", "quantile"),
        perm = NULL, ...)
## S3 method for class 'trafotree'
logLik(object, newdata, weights = NULL, perm = NULL, ...)

```

Arguments

object	an object of class <code>ctm</code> or <code>mlt</code> specifying the abstract model to be partitioned. For <code>predict</code> and <code>logLik</code> , object is an object of class <code>trafotree</code> .
parm	parameters of object those corresponding score is used for finding partitions.
reparm	optional matrix of contrasts for reparameterisation of the scores. <code>teststat = "quadratic"</code> is invariant to this operation but <code>teststat = "max"</code> might be more powerful for example when formulating an implicit into an explicit intercept term.

<code>min_update</code>	number of observations necessary to refit the model in a node. If less observations are available, the parameters from the parent node will be reused.
<code>mltargs</code>	arguments to <code>mlt</code> for fitting the transformation models.
<code>newdata</code>	an optional data frame of observations.
<code>K</code>	number of grid points to generate (in the absence of <code>q</code>).
<code>q</code>	quantiles at which to evaluate the model.
<code>type</code>	type of prediction or plot to generate.
<code>weights</code>	an optional vector of weights.
<code>perm</code>	a vector of integers specifying the variables to be permuted prior before splitting (i.e., for computing permutation variable importances). The default NULL doesn't alter the data, see <code>fitted_node</code> .
<code>...</code>	arguments to <code>ctree</code> , at least formula and data.

Details

Conditional inference trees are used for partitioning likelihood-based transformation models as described in Hothorn and Zeileis (2017). The method can be seen in action in Hothorn (2018) and the corresponding code is available as `demo("BMI")`. `demo("applications")` performs transformation tree analyses for some standard benchmarking problems.

Value

An object of class `trafotree` with corresponding `plot`, `logLik` and `predict` methods.

References

- Torsten Hothorn and Achim Zeileis (2021). Predictive Distribution Modelling Using Transformation Forests. *Journal of Computational and Graphical Statistics*, doi: [10.1080/10618600.2021.1872581](https://doi.org/10.1080/10618600.2021.1872581).
- Torsten Hothorn (2018). Top-Down Transformation Choice. *Statistical Modelling*, **3-4**, 274-298. doi: [10.1177/1471082X17748081](https://doi.org/10.1177/1471082X17748081)
- Natalia Korepanova, Heidi Seibold, Verena Steffen and Torsten Hothorn (2019). Survival Forests under Test: Impact of the Proportional Hazards Assumption on Prognostic and Predictive Forests for ALS Survival. doi: [10.1177/0962280219862586](https://doi.org/10.1177/0962280219862586).

Examples

```
### Example: Stratified Medicine Using Partitioned Cox-Models
### A combination of <DOI:10.1515/ijb-2015-0032> and <arXiv:1701.02110>
### based on infrastructure in the mlt R add-on package described in
### https://cran.r-project.org/web/packages/mlt.docreg/vignettes/mlt.pdf

library("trtf")
library("survival")
### German Breast Cancer Study Group 2 data set
data("GBSG2", package = "TH.data")
```

```

### set-up Cox model with overall treatment effect in hormonal therapy
yvar <- numeric_var("y", support = c(100, 2000), bounds = c(0, Inf))
By <- Bernstein_basis(yvar, order = 5, ui = "incre")
m <- ctm(response = By, shifting = ~ horTh, todistr = "MinExt", data = GBSG2)
GBSG2$y <- with(GBSG2, Surv(time, cens))

### overall log-hazard ratio
coef(cmod <- mlt(m, data = GBSG2))["horThyes"]
### roughly the same as
coef(coxph(y ~ horTh, data = GBSG2))

### partition the model, ie both the baseline hazard function AND the
### treatment effect
(part_cmod <- trafotree(m, formula = y ~ horTh | age + menostat + tsize +
  tgrade + pnodes + progrec + estrec, data = GBSG2))

### compare the log-likelihoods
logLik(cmod)
logLik(part_cmod)

### stronger effects in nodes 2 and 4 and no effect in node 5
coef(part_cmod)[, "horThyes"]

### plot the conditional survivor functions; blue is untreated
### and green is hormonal therapy
nd <- data.frame(horTh = sort(unique(GBSG2$horTh)))
plot(part_cmod, newdata = nd,
  tp_args = list(type = "survivor", col = c("cadetblue3", "chartreuse4")))

### same model, but with explicit intercept term and max-type statistic
### for _variable_ selection
K <- diag(length(coef(m)) - 1)
K[upper.tri(K)] <- 1
K <- cbind(rbind(K, 0), 0)
K[nrow(K), nrow(K)] <- 1
### horThyes is not touched, 6th parameter is intercept
coef(cmod)
(part_cmod_max <- trafotree(m, formula = y ~ horTh | age + menostat + tsize +
  tgrade + pnodes + progrec + estrec, data = GBSG2, reparam = K,
  control = ctree_control(teststat = "max")))
logLik(part_cmod_max)

### the trees (and log-likelihoods are the same) but the
### p-values are sometimes much smaller in the latter tree
cbind(format.pval(info_node(node_party(part_cmod))$criterion["p.value",]),
  format.pval(info_node(node_party(part_cmod_max))$criterion["p.value",]))

```

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