Package ‘LTRCforests’

November 30, 2023

Version 0.7.0
Date 2023-11-29
Title Ensemble Methods for Survival Data with Time-Varying Covariates
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Depends R (>= 3.5.0)
Imports stats, utils, survival, ipred, parallel, prodlim, partykit
Suggests randomForestSRC, LTRCtrees
Description Implements the conditional inference forest and relative risk forest
algorithm to modeling left-truncated right-censored data with time-invariant
co covariates, and (left-truncated) right-censored survival data with time-varying
covariates. It also provides functions to tune the parameters and evaluate the
NeedsCompilation yes
License GPL (>= 2)
Encoding UTF-8
LazyData true
RoxygenNote 7.2.3
Repository CRAN
Date/Publication 2023-11-30 03:50:02 UTC

R topics documented:

LTRCforests-package .................................................. 2
ltrccif ................................................................. 3
ltrcrf ................................................................. 5
pbcsample ............................................................. 8
predictProb ............................................................ 9
Constructs forest methods for left-truncated and right-censored (LTRC) survival data

Description

Constructs a LTRC conditional inference forest (LTRCCIF) or a LTRC relative risk forest (LTRCRRF) for left-truncated and right-censored data, it also allows for (left-truncated) right-censored survival data with time-varying covariates (Yao et al. 2022). The main functions of this package are ltrccif and ltrcrrf.

Details

**Problem setup and existing methods:** Continuous-time survival data with time-varying covariates are common in practice. Methods like the Cox proportional hazards model rely on restrictive assumptions such as proportional hazards and a log-linear relationship between the hazard function and covariates. Furthermore, because these methods are often parametric, nonlinear effects of variables must be modeled by transformations or expanding the design matrix to include specialized basis functions for more complex data structures in real world applications. The functions LTRCIT and LTRCART provide a conditional inference tree method and a relative risk tree method for left-truncated right-censored survival data, which also allows for right-censored survival data with time-varying covariates. Tree estimators are nonparametric and as such often exhibit low bias and high variance. Ensemble methods like bagging and random forest can reduce variance while preserving low bias. The most popular survival forest methods, including conditional inference forest (see cforest), relative risk forest, and random survival forest method (see rfsrC) can only be applied to right-censored survival data with time-invariant covariates.

**LTRC forests:** This package implements ltrccif and ltrcrrf. ltrccif extends the conditional inference forest (see cforest) to LTRC survival data. It uses LTRC conditional inference survival trees as base learners. ltrcrrf extends the relative risk forest (Ishwaran et al. 2004) to left-truncated right-censored survival data. It uses LTRC risk relative tree as base learners. The main functions ltrccif and ltrcrrf fit a corresponding LTRC forest for LTRC data, with parameter mtry tuned by tune.ltrccif or tune.ltrcrrf. This tuning procedure relies on the evaluation of the out-of-bag errors, which is performed by the function sbrier_ltrc. print prints summary output for ltrccif objects and ltrcrrf objects. predictProb constructs survival function estimates for ltrccif objects and ltrcrrf objects. For (left-truncated) right-censored survival data with time-varying covariates, one can first reformat the data structure to one with LTRC observations, where the multiple records of a subject become a list of pseudo-subjects and are treated independently. This procedure is usually referred to as the Andersen-Gill method (Andersen and Gill, 1982). Then LTRC forest methods can be applied on this reformatted dataset.
Overall, the methods in this package can handle all combinations of left truncation, right censoring, time-invariant covariates, and time-varying covariates. If one is in the traditional case with right-censored data and time-invariant covariates, however, then it is recommended to use the functions `cforest` and `rfsrc` directly to construct conditional inference forests and random survival forests, respectively.

References


See Also

`ltrccif`, `ltrcrrf`, `predictProb`, `print`, `tune.ltrccif`, `tune.ltrcrrf`, `sbrier_ltrc`.

ltrccif

*Fit a LTRC conditional inference forest*

**Description**

An implementation of the random forest and bagging ensemble algorithms utilizing LTRC conditional inference trees `LTRCIT` as base learners for left-truncated right-censored survival data with time-invariant covariates. It also allows for (left-truncated) right-censored survival data with time-varying covariates.

**Usage**

```r
ltrccif(formula, data, id, mtry = NULL, ntree = 100L, bootstrap = c("by.sub", "by.root", "by.user", "none"), samptype = c("swor", "swr"), sampfrac = 0.632, samp = NULL, na.action = "na.omit",)
```
stepFactor = 2,
trace = TRUE,
applyfun = NULL,
cores = NULL,
control = partykit::ctree_control(teststat = "quad", testtype = "Univ", minsplit =
  max(ceiling(sqrt(nrow(data))), 20), minbucket = max(ceiling(sqrt(nrow(data))), 7),
  minprob = 0.01, mincriterion = 0, saveinfo = FALSE)
)

Arguments

formula a formula object, with the response being a Surv object, with form Surv(tleft, tright, event).

data a data frame containing n rows of left-truncated right-censored observations. For time-varying data, this should be a data frame containing pseudo-subject observations based on the Andersen-Gill reformulation.
id variable name of subject identifiers. If this is present, it will be searched for in the data data frame. Each group of rows in data with the same subject id represents the covariate path through time of a single subject. If not specified, the algorithm then assumes data contains left-truncated and right-censored survival data with time-invariant covariates.
mtry number of input variables randomly sampled as candidates at each node for random forest algorithms. The default mtry is tuned by tune.ltrccif.
ntree an integer, the number of the trees to grow for the forest. ntree = 100L is set by default.
bootstrap bootstrap protocol. (1) If id is present, the choices are: "by.sub" (by default) which bootstraps subjects, "by.root" which bootstraps pseudo-subjects. Both can be with or without replacement (by default sampling is without replacement; see the option perturb below); (2) If id is not specified, it bootstraps the data by sampling with or without replacement. Regardless of the presence of id, if "none" is chosen, data is not bootstrapped at all, and is used in every individual tree. If "by.user" is chosen, the bootstrap specified by samp is used.
samptype choices are swor (sampling without replacement) and swr (sampling with replacement). The default action here is sampling without replacement.
sampfrac a fraction, determining the proportion of subjects to draw without replacement when samptype = "swor". The default value is 0.632. To be more specific, if id is present, 0.632 * N of subjects with their pseudo-subject observations are drawn without replacement (N denotes the number of subjects); otherwise, 0.632 * n is the requested size of the sample.
samp Bootstrap specification when bootstype = "by.user". Array of dim n x ntree specifying how many times each record appears in each bootstrap sample.
na.action action taken if the data contains NA's. The default "na.omit" removes the entire record if any of its entries is NA (for x-variables this applies only to those specifically listed in formula). See function cforest for other available options.
stepFactor at each iteration, mtry is inflated (or deflated) by this value, used when mtry is not specified (see tune.ltrccif). The default value is 2.
trace whether to print the progress of the search of the optimal value of mtry, when mtry is not specified (see tune.ltrccif). trace = TRUE is set by default.

applyfun an optional lapply-style function with arguments function(X, FUN, ...). It is used for computing the variable selection criterion. The default is to use the basic lapply function unless the cores argument is specified (see below). See ctree_control.

cores numeric. See ctree_control.

control a list of control parameters, see ctree_control. control parameters minsplit, minbucket have been adjusted from the cforest defaults. Other default values correspond to those of the default values used by ctree_control.

Details

This function extends the conditional inference survival forest algorithm in cforest to fit left-truncated and right-censored data, which allow for time-varying covariates.

Value

An object belongs to the class ltrccif, as a subclass of cforest.

References


See Also

predictProb for prediction and tune.ltrccif for mtry tuning.

Examples

#### Example with time-varying data pbcsample
library(survival)
Formula = Surv(Start, Stop, Event) ~ age + alk.phos + ast + chol + edema
## Fit an LTRCCIF on the time-invariant data, with mtry tuned with stepFactor = 3.
LTRCCIFobj = ltrccif(formula = Formula, data = pbcsample, ntree = 20L, stepFactor = 3)

### Fit a LTRC relative risk forest

Description

An implementation of the random forest algorithms utilizing LTRC rpart trees LTRCART as base learners for left-truncated right-censored survival data with time-invariant covariates. It also allows for (left-truncated) right-censored survival data with time-varying covariates.
ltcrcrf(
    formula,
    data,
    id,
    ntree = 100L,
    mtry = NULL,
    nodesize = max(ceiling(sqrt(nrow(data))), 15),
    bootstrap = c("by.sub", "by.root", "by.node", "by.user", "none"),
    samptype = c("swor", "swr"),
    sampfrac = 0.632,
    samp = NULL,
    na.action = "na.omit",
    stepFactor = 2,
    trace = TRUE,
    nodedepth = NULL,
    nsplit = 10L,
    ntime
)

Arguments

formula a formula object, with the response being a Surv object, with form Surv(tleft, tright, event).
data a data frame containing n rows of left-truncated right-censored observations. For time-varying data, this should be a data frame containing pseudo-subject observations based on the Andersen-Gill reformulation.
id variable name of subject identifiers. If this is present, it will be searched for in the data data frame. Each group of rows in data with the same subject id represents the covariate path through time of a single subject. If not specified, the algorithm then assumes data contains left-truncated and right-censored survival data with time-invariant covariates.
ntree an integer, the number of the trees to grow for the forest. ntree = 100L is set by default.
mtry number of input variables randomly sampled as candidates at each node for random forest like algorithms. The default mtry is tuned by tune.ltcrrf.
nodesize an integer, forest average terminal node size.
bootstrap bootstrap protocol. (1) If id is present, the choices are: "by.sub" (by default) which bootstraps subjects, "by.root" which bootstraps pseudo-subjects. Both can be with or without replacement (by default sampling is without replacement; see the option samptype below). (2) If id is not specified, the default is "by.root" which bootstraps the data by sampling with or without replacement; if "by.node" is choosen, data is bootstrapped with replacement at each node while growing the tree. Regardless of the presence of id, if "none" is chosen, data is not bootstrapped at all, and is used in every individual tree. If "by.user" is choosen, the bootstrap specified by samp is used.
samptype choices are swor (sampling without replacement) and swr (sampling with replacement). The default action here is sampling without replacement.

sampfrac a fraction, determining the proportion of subjects to draw without replacement when samptype = "swor". The default value is 0.632. To be more specific, if id is present, 0.632 * N of subjects with their pseudo-subject observations are drawn without replacement (N denotes the number of subjects); otherwise, 0.632 * n is the requested size of the sample.

samp Bootstrap specification when bootstype = "by.user". Array of dim n x ntree specifying how many times each record appears in each bootstrap sample.

na.action action taken if the data contains NA's. The default "na.omit" removes the entire record if any of its entries is NA (for x-variables this applies only to those specifically listed in formula). See function rfsr for other available options.

stepFactor at each iteration, mtry is inflated (or deflated) by this value, used when mtry is not specified (see tune.ltrcrrf). The default value is 2.

trace whether to print the progress of the search of the optimal value of mtry if mtry is not specified (see tune.ltrcrrf). trace = TRUE is set by default.

nodedepth maximum depth to which a tree should be grown. The default behaviour is that this parameter is ignored.

nsplit an non-negative integer value for number of random splits to consider for each candidate splitting variable. This significantly increases speed. When zero or NULL, the algorithm uses much slower deterministic splitting where all possible splits are considered. nsplit = 10L by default.

ntime an integer value used for survival to constrain ensemble calculations to a grid of ntime time points. Alternatively if a vector of values of length greater than one is supplied, it is assumed these are the time points to be used to constrain the calculations (note that the constrained time points used will be the observed event times closest to the user supplied time points). If no value is specified, the default action is to use all observed event times. Further details can be found in rfsr.

Details
This function extends the relative risk forest algorithm (Ishwaran et al. 2004) to fit left-truncated and right-censored data, which allows for time-varying covariates. The algorithm is built based on employing the fast C code from rfsr.

Value
An object belongs to the class ltrcrrf, as a subclass of rfsr.

References


See Also

`predictProb` for prediction and `tune.ltrcrrf` for `mtry` tuning.

Examples

```r
### Example with time-varying data pbsample
library(survival)
Formula = Surv(Start, Stop, Event) ~ age + alk.phos + ast + chol + edema
# Built a LTRCRRF forest (based on bootstrapping subjects without replacement)
# on the time-varying data by specifying id:
LTRCRRFobj = ltrcrrf(formula = Formula, data = pbsample, id = ID, stepFactor = 3,
                      ntree = 10L)
```

### pbsample

**Sample Mayo Clinic Primary Biliary Cirrhosis Data**

**Description**

A sample real dataset with time varying covariates. It contains multiple records of measurements of risk factors at multiple time points from 10 patients with primary biliary cirrhosis (PBC), constructed from `pbcseq` in the R package `survival`. The data structure has been reformatted with left-truncated right-censored pseudo-subject observations based on the Andersen-Gill reformulation method.

**Value**

A data frame with 57 rows and 9 variables:

- **ID**: patient id.
- **Start**: the left truncation time point for the corresponding pseudo-subject observation.
- **Stop**: the right censoring time point for the corresponding pseudo-subject observation.
- **Event**: a binary value, with 1 indicating the event occurring at the corresponding Stop, 0 indicating right-censored.
- **age**: patient’s age at entry, in years.
- **alk.phos**: alkaline phosphotase (U/liter).
- **ast**: aspartate aminotransferase (U/ml).
- **chol**: serum cholesterol (mg/dl).
- **edema**: 0–no edema, 0.5–untreated or successfully treated, 1–edema despite diuretic therapy.
**predictProb**

**Source**


**References**


---

**predictProb**

Compute a Survival Curve from a LTRCCIF model or a LTRCRRF model

**Description**

Constructs a monotone nonincreasing estimated survival curve from a LTRCCIF model or a LTRCRRF model for any given (left-truncated) right-censored survival data with time-varying covariates. It can also compute survival function estimates for left-truncated right-censored data with time-invariant covariates.

**Usage**

```
predictProb(
  object,
  newdata = NULL,
  newdata.id,
  OOB = FALSE,
  time.eval,
  time.tau = NULL
)
```

**Arguments**

- **object**: an object as returned by `ltrccif` or by `ltrcrrf`.
- **newdata**: an optional data frame containing the test data (with the names of the variables the same as those in data from object).
- **newdata.id**: optional variable name of subject identifiers for newdata. If this is present, it will be searched for in the newdata data frame. Each group of rows in newdata with the same subject id represents the covariate path through time of a single subject, and the result will contain one curve per subject. If it is not specified, then an estimated survival curve is returned for each row of newdata.
- **OOB**: a logical specifying whether out-of-bag predictions are desired (only if newdata = NULL).
- **time.eval**: a vector of time points, at which the estimated survival probabilities will be computed.
predictProb

time.tau an optional vector, with the i-th entry giving the upper time limit for the computed survival probabilities for the i-th data of interest (i.e., only computes survival probabilities at time.eval[time.eval <= time.tau[i]] for the i-th data of interest). If OOB = TRUE, the length of time.tau is equal to the size of data used to train the object; if OOB = FALSE, the length of time.tau is equal to the size of newdata, or equal to the size of data if newdata is not given. The default NULL is simply to set all entries of time.tau equal to the maximum value of time.eval, so that all estimated survival probabilities are computed at the same time.eval.

Value

A list containing:

- survival.id subject identifiers.
- survival.obj an object of class Surv.
- survival.probs the estimated survival probabilities for each data of interest. It is a list if the length of the estimated values differs from one to another; otherwise, it is a matrix with the number of columns equal to the number of the data of interest, number of rows equal to the number of the time points at which the estimated survival probabilities are computed.
- survival.tau the input value time.tau.
- survival.times the input value time.eval.

See Also

sbrier_ltrc for evaluation of model fit

Examples

```r
### Example with data pbcsample
library(survival)
Formula <- Surv(Start, Stop, Event) ~ age + alk.phos + ast + chol + edema
## Fit an LTRC conditional inference forest on time-varying data
LTRCCIFobj <- ltrccif(formula = Formula, data = pbcsample, id = ID,
                      mtry = 3, ntree = 50L)

## Construct an estimated survival estimate for the second subject
tpnt <- seq(0, max(pbcsample$Stop), length.out = 50)
newData <- pbcsample[pbcsample$ID == 2, ]
Pred <- predictProb(object = LTRCCIFobj, newdata = newData, newdata.id = ID,
                    time.eval = tpnt)
## Since time.tau = NULL, Pred$survival.probs is in the matrix format, with dimensions:
dim(Pred$survival.probs) # length(time.eval) x nrow(newdata)
## Plot the estimated survival curve
plot(Pred$survival.times, Pred$survival.probs, type = "l", col = "red",
     xlab = "Time", ylab = "Survival probabilities")
```
**Description**

Print summary output after a LTRCCIF or a LTRCRRF model is built. This is the default print method for objects in the class of `ltrccif` or `ltrcrrf`.

**Usage**

```r
print(x)
```

**Arguments**

- `x` an object of class `ltrccif` or `ltrcrrf`.

**Value**

A printout object containing the following components:
- **Number of (pseudo-subject) observations**
  number of left-truncated right-censored pseudo-subject observations based on the Andersen-Gill reformulation.
- **Number of subjects**
  number of independent subject observations.
- **Number of deaths**
  number of times that an event occurs in the whole dataset.
- **Number of trees**
  the value set for argument `ntree`, see `ltrccif` and `ltrcrrf`.
- **minsplit**
  the value set for argument `minsplit` that controls the growth of individual trees; see `ctree_control`.
- **minbucket**
  the value set for argument `minbucket` that controls the growth of individual trees; see `ctree_control`.
- **minprob**
  the value set for argument `minprob` that controls the growth of individual trees; see `ctree_control`.
- **maxdepth**
  the value set for argument `maxdepth` that controls the maximum depth of individual trees; see `ctree_control`.
- **No. of variables tried at each split**
  number of input variables randomly sampled as candidates at each node for random forest algorithms, which is either set as an argument `mtry` in `ltrccif` and `ltrcrrf`, or tuned by `tune.ltrccif` or `tune.ltrcrrf`, respectively.
- **Total no. of variables**
  the number of features provided in data.
- **Bootstrap type to grow trees**
  the values set for argument `bootstrap`, see `ltrccif` and `ltrcrrf`. 
Resampling used to grow trees
  the value set for argument samptype, see `ltrccif` and `ltrcrrf`.
Resampling rate used to grow trees
  the values set for argument sampfrac, see `ltrccif` and `ltrcrrf`.
Analysis
  LTRCCIF for a `ltrccif` object or LTRCRRF for `ltrcrrf`.
Family
  the family used in the analysis, `surv`.
Splitting rule
  the splitting rule that is implemented, conditional inference framework for a `ltrccif` object or Poisson splitting for `ltrcrrf`.
Number of random split points
  the values set for argument nsplit in `ltrcrrf`.

See Also
  `ltrccif`, `ltrcrrf`

Examples

```r
library(survival)
Formula = Surv(Start, Stop, Event) ~ age + alk.phos + ast + chol + edema
# Built a LTRCCIF forest on the time-varying data by specifying id, with mtry specified:
LTRCCIFobj = ltrccif(formula = Formula, data = pbcsample, id = ID, mtry = 3, ntree = 50L)
print(LTRCCIFobj)

# Built a LTRCCIF forest on the time-invariant data, with resampling, with mtry specified:
LTRCCIFobj = ltrccif(formula = Formula, data = pbcsample, samptype = "swr",
                     mtry = 3, ntree = 50L)
print(LTRCCIFobj)
```

---

**sbrier_ltrc**  
*Model fit evaluation for LTRC forests.*

**Description**

Compute the (integrated) Brier score to evaluate the model fit for (left-truncated) right-censored survival data with time-varying covariates, as well as left-truncated right-censored data with time-invariant covariates.

**Usage**

```r
sbrier_ltrc(obj, id = NULL, pred, type = c("IBS", "BS"))
```
Arguments

- **obj**: an object of class `Surv`, formed on left-truncated right-censored observations (which are pseudo-subject observations in the time-varying case).
- **id**: an optional vector as subject identifiers for `obj`.
- **pred**: a list. This should contain 1) either a matrix or a list of survival probabilities named `survival.probs`; 2) a sequence of time points `survival.times`; 3) a vector of upper time limits `survival.tau`. See the values returned by `predictProb`.
- **type**: a character string denoting the type of scores returned. If `type = "IBS"`, the integrated Brier score up to the last time point in `pred$surv.times` that is not larger than the minimum value of `pred$surv.tau` is returned. If `type = "BS"`, the Brier score at every time point in `pred$surv.times` up to the minimum value of `pred$surv.tau` is returned. `type = "IBS"` is set by default.

Value

If `type = "IBS"`, this returns the integrated Brier score.

If `type = "BS"`, this returns `BScore`, the Brier scores and `Time`, the time points at which the scores are computed.

Examples

```r
### Example with dataset pbcsample
library(survival)
Formula = Surv(Start, Stop, Event) ~ age + alk.phos + ast + chol + edema
## Fit an LTRC conditional inference forest on time-varying data
LTRCCIFobj = ltrccif(formula = Formula, data = pbcsample, id = ID, mtry = 3, ntree = 50L)

# Time points
tpnt = seq(0, 6000, by = 100)
# Set different upper time limits for each of the subjects
tau = seq(4001, 6200, length.out = length(unique(pbcsample$ID)))
## Obtain estimation at time points tpnt
Predobj = predictProb(object = LTRCCIFobj, time.eval = tpnt, time.tau = tau)

## Compute the integrated Brier score:
pbcobj = Surv(pbcsample$Start, pbcsample$Stop, pbcsample$Event)
IBS = sbrier_ltrc(obj = pbcobj, id = pbcsample$ID, pred = Predobj, type = "IBS")

## Compute the Brier score at each value of tpnt
BS = sbrier_ltrc(obj = pbcobj, id = pbcsample$ID, pred = Predobj, type = "BS")
## Plot the Brier scores
plot(BS$Time, BS$BScore, pch = 20, xlab = "Time", ylab = "Brier score", col = 2)
## As one can see, the Brier scores are returned at all tpnt up to 4000,
## this is because the algorithm set the last evaluation time point
## to be 4000 based on the value of time.eval and time.tau
## (max(tpnt[tpnt <= min(tau)]) == 4000).
```
tune.ltrccif

Tune mtry to the optimal value with respect to out-of-bag error for a LTRCCIF model

Description

Starting with the default value of mtry, search for the optimal value (with respect to out-of-bag error estimate) of mtry for ltrccif.

Usage

tune.ltrccif(
  formula, data, id, mtryStart = NULL, stepFactor = 2, time.eval = NULL, time.tau = NULL, ntreeTry = 100L, bootstrap = c("by.sub", "by.root", "none", "by.user"), samptype = c("swor", "swr"), sampfrac = 0.632, samp = NULL, na.action = "na.omit", trace = TRUE, doBest = FALSE, plot = FALSE, applyfun = NULL, cores = NULL, control = partykit::ctree_control(teststat = "quad", testtype = "Univ", mincriterion = 0, saveinfo = FALSE, minsplit = max(ceiling(sqrt(nrow(data))), 20), minbucket = max(ceiling(sqrt(nrow(data))), 7), minprob = 0.01)
)

Arguments

formula a formula object, with the response being a Surv object, with form Surv(tleft, tright, event).

data a data frame containing n rows of left-truncated right-censored observations.
id variable name of subject identifiers. If this is present, it will be searched for in the data data frame. Each group of rows in data with the same subject id represents the covariate path through time of a single subject. If not specified, the algorithm then assumes data contains left-truncated and right-censored survival data with time-invariant covariates.
mtryStart starting value of mtry; default is sqrt(nvar).
### `tune.ltrccif`

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
<th>Default Value</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>stepFactor</code></td>
<td>at each iteration, mtry is inflated (or deflated) by this value. The default value is 2.</td>
<td></td>
<td></td>
</tr>
<tr>
<td><code>time.eval</code></td>
<td>a vector of time points, at which the estimated survival probabilities are evaluated.</td>
<td></td>
<td></td>
</tr>
<tr>
<td><code>time.tau</code></td>
<td>an optional vector, with the (i)-th entry giving the upper time limit for the computed survival probabilities for the (i)-th data (i.e., only computes survival probabilities at (time.eval[time.eval \leq time.tau[i]]) for the (i)-th data of interest).</td>
<td></td>
<td></td>
</tr>
<tr>
<td><code>ntreeTry</code></td>
<td>number of trees used at the tuning step.</td>
<td></td>
<td></td>
</tr>
<tr>
<td><code>bootstrap</code></td>
<td>bootstrap protocol. (1) If id is present, the choices are: &quot;by.sub&quot; (by default) which bootstraps subjects, &quot;by.root&quot; which bootstraps pseudo-subjects. Both can be with or without replacement (by default sampling is without replacement; see the option <code>perturb</code> below); (2) If id is not specified, it bootstraps the data by sampling with or without replacement. Regardless of the presence of id, if &quot;none&quot; is chosen, data is not bootstrapped at all, and is used in every individual tree. If &quot;by.user&quot; is chosen, the bootstrap specified by <code>samp</code> is used.</td>
<td></td>
<td></td>
</tr>
<tr>
<td><code>samptype</code></td>
<td>choices are <code>swor</code> (sampling without replacement) and <code>swr</code> (sampling with replacement). The default action here is sampling without replacement.</td>
<td></td>
<td></td>
</tr>
<tr>
<td><code>sampfrac</code></td>
<td>a fraction, determining the proportion of subjects to draw without replacement when <code>samptype = &quot;swor&quot;</code>. The default value is 0.632. To be more specific, if id is present, 0.632 * N of subjects with their pseudo-subject observations are drawn without replacement (N denotes the number of subjects); otherwise, 0.632 * n is the requested size of the sample.</td>
<td></td>
<td></td>
</tr>
<tr>
<td><code>samp</code></td>
<td>Bootstrap specification when <code>bootstype = &quot;by.user&quot;</code>. Array of dim n x ntree specifying how many times each record appears in each bootstrap sample.</td>
<td></td>
<td></td>
</tr>
<tr>
<td><code>na.action</code></td>
<td>action taken if the data contains NA's. The default &quot;na.omit&quot; removes the entire record if any of its entries is NA (for x-variables this applies only to those specifically listed in formula). See function <code>cforest</code> for other available options.</td>
<td></td>
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</tr>
<tr>
<td><code>trace</code></td>
<td>whether to print the progress of the search. <code>trace = TRUE</code> is set by default.</td>
<td></td>
<td></td>
</tr>
<tr>
<td><code>doBest</code></td>
<td>whether to run a <code>ltrccif</code> object using the optimal mtry found. <code>doBest = FALSE</code> is set by default.</td>
<td></td>
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<tr>
<td><code>plot</code></td>
<td>whether to plot the out-of-bag error as a function of mtry. <code>plot = FALSE</code> is set by default.</td>
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</tr>
<tr>
<td><code>applyfun</code></td>
<td>an optional <code>lapply</code>-style function with arguments <code>function(X, FUN, ...)</code>. It is used for computing the variable selection criterion. The default is to use the basic <code>lapply</code> function unless the <code>cores</code> argument is specified (see below). See <code>ctree_control</code>.</td>
<td></td>
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</tr>
<tr>
<td><code>cores</code></td>
<td>numeric. See <code>ctree_control</code>.</td>
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<tr>
<td><code>control</code></td>
<td>a list with control parameters, see <code>cforest</code>. The default values correspond to those of the default values used by <code>ltrccif</code>.</td>
<td></td>
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</tr>
</tbody>
</table>

### Value

If `doBest = FALSE` (default), this returns the optimal mtry value of those searched. If `doBest = TRUE`, this returns the `ltrccif` object produced with the optimal mtry.
See Also

`sbrier_ltrc` for evaluation of model fit when searching for the optimal value of `mtry`.

Examples

```r
### Example with data pbcsample
library(survival)
Formula = Surv(Start, Stop, Event) ~ age + alk.phos + ast + chol + edema
## mtry tuned by the OOB procedure with stepFactor 3, number of trees built 10.
mtryT = tune.ltrccif(formula = Formula, data = pbcsample, id = ID, stepFactor = 3,
                      ntreeTry = 10L)
```

---

### Description

Starting with the default value of `mtry`, search for the optimal value (with respect to out-of-bag error estimate) of `mtry` for `ltrcrrf`.

### Usage

```r
tune.ltrcrrf(
  formula, 
  data, 
  id, 
  mtryStart = NULL, 
  stepFactor = 2, 
  time.eval = NULL, 
  time.tau = NULL, 
  ntreeTry = 100L, 
  bootstrap = c("by.sub", "by.root", "by.node", "by.user", "none"), 
  samptype = c("swor", "swr"), 
  sampfrac = 0.632, 
  samp = NULL, 
  na.action = "na.omit", 
  trace = TRUE, 
  doBest = FALSE, 
  plot = FALSE, 
  ntime, 
  nsplit = 10L, 
  nodesizeTry = max(ceiling(sqrt(nrow(data))), 15), 
  nodedepth = NULL
)
```
Arguments

- **formula**: a formula object, with the response being a `Surv` object, with form `Surv(tleft, tright, event)`.
- **data**: a data frame containing n rows of left-truncated right-censored observations.
- **id**: variable name of subject identifiers. If this is present, it will be searched for in the data data frame. Each group of rows in data with the same subject id represents the covariate path through time of a single subject. If not specified, the algorithm then assumes data contains left-truncated and right-censored survival data with time-invariant covariates.
- **mtryStart**: starting value of mtry; default is `sqrt(nvar)`.
- **stepFactor**: at each iteration, mtry is inflated (or deflated) by this value, used when mtry is not specified (see `ltrcrf`). The default value is 2.
- **time.eval**: a vector of time points, at which the estimated survival probabilities are evaluated.
- **time.tau**: an optional vector, with the i-th entry giving the upper time limit for the computed survival probabilities for the i-th data (i.e., only computes survival probabilities at `time.eval[time.eval <= time.tau[i]]` for the i-th data of interest).
- **ntreeTry**: number of trees used at the tuning step.
- **bootstrap**: bootstrap protocol. (1) If id is present, the choices are: "by.sub" (by default) which bootstraps subjects, "by.root" which bootstraps pseudo-subjects. Both can be with or without replacement (by default sampling is without replacement; see the option `samptype` below). (2) If id is not specified, the default is "by.root" which bootstraps the data by sampling with or without replacement; if "by.node" is choosen, data is bootstrapped with replacement at each node while growing the tree. Regardless of the presence of id, if "none" is chosen, the data is not bootstrapped at all. If "by.user" is choosen, the bootstrap specified by samp is used.
- **samptype**: choices are swor (sampling without replacement) and swr (sampling with replacement). The default action here is sampling without replacement.
- **sampfrac**: a fraction, determining the proportion of subjects to draw without replacement when `samptype = "swor"`. The default value is 0.632. To be more specific, if id is present, 0.632 * N of subjects with their pseudo-subject observations are drawn without replacement (N denotes the number of subjects); otherwise, 0.632 * n is the requested size of the sample.
- **samp**: Bootstrap specification when `bootstype = "by.user"`. Array of dim n x ntree specifying how many times each record appears in each bootstrap sample.
- **na.action**: action taken if the data contains NA's. The default "na.omit" removes the entire record if any of its entries is NA (for x-variables this applies only to those specifically listed in formula). See function `rfsr` for other available options.
- **trace**: whether to print the progress of the search. `trace = TRUE` is set by default.
- **doBest**: whether to run a `ltrcrf` object using the optimal mtry found. `doBest = FALSE` is set by default.
- **plot**: whether to plot the out-of-bag error as a function of mtry. `plot = FALSE` is set by default.
ntime an integer value used for survival to constrain ensemble calculations to a grid of ntime time points. Alternatively if a vector of values of length greater than one is supplied, it is assumed these are the time points to be used to constrain the calculations (note that the constrained time points used will be the observed event times closest to the user supplied time points). If no value is specified, the default action is to use all observed event times.

nsplit an non-negative integer value for number of random splits to consider for each candidate splitting variable. This significantly increases speed. When zero or NULL, the algorithm uses much slower deterministic splitting where all possible splits are considered. nsplit = 10L by default.

nodesizeTry forest average terminal node size used at the tuning step.

nodedepth maximum depth to which a tree should be grown. The default behaviour is that this parameter is ignored.

Value
If doBest = FALSE (default), this returns the optimal mtry value of those searched.
If doBest = TRUE, this returns the ltrcrrf object produced with the optimal mtry.

See Also
sbrier_ltrc for evaluation of model fit for the optimal value of mtry.

Examples
### Example with data pbcsample
library(survival)
Formula = Surv(Start, Stop, Event) ~ age + alk.phos + ast + chol + edema
## mtry tuned by the OOB procedure with stepFactor 3, number of trees built 10.
mtryT = tune.ltrcrrf(formula = Formula, data = pbcsample, stepFactor = 3, ntreeTry = 10L)
Index

cforest, 2–5, 15
ctree_control, 5, 11, 15

LTRCART, 2, 5
ltrccif, 2, 3, 3, 9, 11, 12, 14, 15
LTRCforests-package, 2
LTRCIT, 2, 3
ltrcrrf, 2, 3, 5, 9, 11, 12, 16–18

pbcsample, 8
pbcseq, 8
predictProb, 2, 3, 5, 8, 9, 13
print, 2, 3, 11

rfsrc, 2, 3, 7, 17

sbrier_ltrc, 2, 3, 10, 12, 16, 18
Surv, 4, 6, 10, 13, 14, 17
survival, 8

tune.ltrccif, 2–5, 11, 14
tune.ltrcrrf, 2, 3, 6–8, 11, 16