

Package: riskRegression (via r-universe)

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Type Package

Title Risk Regression Models and Prediction Scores for Survival
Analysis with Competing Risks

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testthat, R.rsp

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Description Implementation of the following methods for event history analysis. Risk regression models for survival endpoints also in the presence of competing risks are fitted using binomial regression based on a time sequence of binary event status variables. A formula interface for the Fine-Gray regression model and an interface for the combination of cause-specific Cox regression models. A toolbox for assessing and comparing performance of risk predictions (risk markers and risk prediction models). Prediction performance is measured by the Brier score and the area under the ROC curve for binary possibly time-dependent outcome. Inverse probability of censoring weighting and pseudo values are used to deal with right censored data. Lists of risk markers and lists of risk models are assessed simultaneously. Cross-validation repeatedly splits the data, trains the risk prediction models on one part of each split and then summarizes and compares the performance across splits.

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BugReports <https://github.com/tagteam/riskRegression/issues>

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 anova.ate

Risk Comparison Over Time

Description

Comparison of risk differences or risk ratios over all timepoints.

Usage

```
## S3 method for class 'ate'
anova(
  object,
  allContrast = NULL,
  type = "diff",
  estimator = object$estimator[1],
  test = "CvM",
  transform = NULL,
  alternative = "two.sided",
  n.sim = 10000,
  print = TRUE,
  ...
)
```

Arguments

object	A ate object, i.e. output of the ate function.
allContrast	[matrix] contrast for which the risks should be compared. Matrix with two rows, the first being the sequence of reference treatments and the second the sequence of alternative treatments.
type	[character vector] the functional used to compare the risks: "diffRisk" or "ratioRisk".
estimator	[character] The type of estimator relative to which the comparison should be performed.
test	[character] The type of statistic used to compare the risks over times: "KM" (extremum risk), "CvM" (sum of squares of the risk), or "sum" (sum of the risks).
transform	[character] Should a transformation be used, e.g. the test is performed after log-transformation of the estimate, standard error, and influence function.
alternative	[character] a character string specifying the alternative hypothesis, must be one of "two.sided", "greater" or "less".
n.sim	[integer, >0] the number of simulations used to compute the p-values.
print	[logical] should the results be displayed?
...	Not used.

Details

Experimental!!!

Examples

```

library(survival)
library(data.table)
library(ggplot2)

## Not run:
## simulate data
set.seed(12)
n <- 200
dtS <- sampleData(n,outcome="survival")
dtS$X12 <- LETTERS[as.numeric(as.factor(paste0(dtS$X1,dtS$X2)))]
dtS <- dtS[dtS$X12!="D"]

## model fit
fit <- coxph(formula = Surv(time,event)~ X1+X6,data=dtS,y=TRUE,x=TRUE)
seqTime <- 1:10
ateFit <- ate(fit, data = dtS, treatment = "X1", contrasts = NULL,
             times = seqTime, B = 0, iid = TRUE, se = TRUE, verbose = TRUE, band = TRUE)

## display
autoplot(ateFit)

## inference (two sided)
statistic <- ateFit$diffRisk$estimate/ateFit$diffRisk$se
confint(ateFit, p.value = TRUE, method.band = "bonferroni")$diffRisk
confint(ateFit, p.value = TRUE, method.band = "maxT-simulation")$diffRisk

anova(ateFit, test = "KS")
anova(ateFit, test = "CVM")
anova(ateFit, test = "sum")

## manual calculation (one sided)
n.sim <- 1e4
statistic <- ateFit$diffRisk[, estimate/se]
iid.norm <- scale(ateFit$iid$GFORMULA[["1"]]-ateFit$iid$GFORMULA[["0"]],
                 scale = ateFit$diffRisk$se)

ls.out <- lapply(1:n.sim, function(iSim){
  iG <- rnorm(NROW(iid.norm))
  iCurve <- t(iid.norm) %*% iG
  data.table(max = max(iCurve), L2 = sum(iCurve^2), sum = sum(iCurve),
             maxC = max(iCurve) - max(statistic),
             L2C = sum(iCurve^2) - sum(statistic^2),
             sumC = sum(iCurve) - sum(statistic),
             sim = iSim)
})

dt.out <- do.call(rbind,ls.out)

```

```

dt.out[,.(max = mean(.SD$maxC>=0),
           L2 = mean(.SD$L2C>=0),
           sum = mean(.SD$sumC>=0))]

## permutation
n.sim <- 250
stats.perm <- vector(mode = "list", length = n.sim)
pb <- txtProgressBar(max = n.sim, style=3)
treatVar <- ateFit$variables["treatment"]

for(iSim in 1:n.sim){ ## iSim <- 1
  iData <- copy(dtS)
  iIndex <- sample.int(NROW(iData), replace = FALSE)
  iData[, c(treatVar) := .SD[[treatVar]][iIndex]]

  iFit <- update(fit, data = iData)
  iAteSim <- ate(iFit, data = iData, treatment = unname(treatVar),
                times = seqTime, verbose = FALSE)
  iStatistic <- iAteSim$diffRisk[,estimate/se]
  stats.perm[[iSim]] <- cbind(iAteSim$diffRisk[,.(max = max(iStatistic),
                                                         L2 = sum(iStatistic^2),
                                                         sum = sum(iStatistic))],
                             sim = iSim)
  stats.perm[[iSim]]$maxC <- stats.perm[[iSim]]$max - max(statistic)
  stats.perm[[iSim]]$L2C <- stats.perm[[iSim]]$L2 - sum(statistic^2)
  stats.perm[[iSim]]$sumC <- stats.perm[[iSim]]$sum - sum(statistic)
  setTxtProgressBar(pb, iSim)
}

dtstats.perm <- do.call(rbind,stats.perm)
dtstats.perm[,.(max = mean(.SD$maxC>=0),
                L2 = mean(.SD$L2C>=0),
                sum = mean(.SD$sumC>=0))]

## End(Not run)

```

as.data.table.ate *Turn ate Object Into a data.table*

Description

Turn ate object into a data.table.

Usage

```

## S3 method for class 'ate'
as.data.table(
  x,
  keep.rownames = FALSE,
  estimator = x$estimator,

```

```

  type = c("meanRisk", "diffRisk", "ratioRisk"),
  ...
)

```

Arguments

x	object obtained with function <code>ate</code>
keep.rownames	Not used.
estimator	[character] The type of estimator relative to which the estimates should be output.
type	[character vector] The type of risk to export. Can be "meanRisk" to export the risks specific to each treatment group, "diffRisk" to export the difference in risks between treatment groups, or "ratioRisk" to export the ratio of risks between treatment groups.
...	Not used.

```
as.data.table.influenceTest
```

Turn influenceTest Object Into a data.table

Description

Turn `influenceTest` object into a `data.table`.

Usage

```

## S3 method for class 'influenceTest'
as.data.table(x, keep.rownames = FALSE, se = TRUE, ...)

```

Arguments

x	object obtained with function <code>influenceTest</code>
keep.rownames	Not used.
se	[logical] Should standard errors/quantile for confidence bands be displayed?
...	Not used.

`as.data.table.predictCox`*Turn predictCox Object Into a data.table*

Description

Turn predictCox object into a data.table.

Usage

```
## S3 method for class 'predictCox'  
as.data.table(x, keep.rownames = FALSE, se = TRUE, ...)
```

Arguments

x	object obtained with function predictCox
keep.rownames	Not used.
se	[logical] Should standard errors/quantile for confidence bands be displayed?
...	Not used.

`as.data.table.predictCSC`*Turn predictCSC Object Into a data.table*

Description

Turn predictCSC object into a data.table.

Usage

```
## S3 method for class 'predictCSC'  
as.data.table(x, keep.rownames = FALSE, se = TRUE, ...)
```

Arguments

x	object obtained with function predictCSC
keep.rownames	not used
se	should standard errors/quantile for confidence bands be displayed?
...	not used

ate *Average Treatment Effects Computation*

Description

Use the g-formula or the IPW or the double robust estimator to estimate the average treatment effect (absolute risk difference or ratio) based on Cox regression with or without competing risks.

Usage

```
ate(
  event,
  treatment,
  censor = NULL,
  data,
  data.index = NULL,
  formula = NULL,
  estimator = NULL,
  strata = NULL,
  contrasts = NULL,
  allContrasts = NULL,
  times,
  cause = NA,
  landmark = NULL,
  se = TRUE,
  iid = (B == 0) && (se || band),
  known.nuisance = FALSE,
  band = FALSE,
  B = 0,
  seed,
  handler = "foreach",
  mc.cores = 1,
  cl = NULL,
  verbose = TRUE,
  ...
)
```

Arguments

event	Outcome model which describes how the probability of experiencing a terminal event depends on treatment and covariates. The object carry its own call and have a predictRisk method. See details section and examples section.
treatment	Name of the treatment variable or treatment model which describes how the probability of being allocated to a treatment group depends on covariates. See details section and examples section.
censor	Censoring model which describes how the probability of being censored depends on treatment and covariates. See details section and examples section.

data	[data.frame or data.table] Data set in which to evaluate risk predictions based on the outcome model
data.index	[numeric vector] Position of the observation in argument data relative to the dataset used to obtain the argument event, treatment, censor. Only necessary for the standard errors when computing the Average Treatment Effects on a subset of the data set.
formula	For analyses with time-dependent covariates, the response formula. See examples.
estimator	[character] The type of estimator used to compute the average treatment effect. Can be "G-formula", "IPTW", or "AIPTW". When using <code>estimator="G-formula"</code> , a model for the outcome should be provided (argument event). When using <code>estimator="IPTW"</code> , a model for the treatment should be provided (argument treatment), as well as for the censoring (if any, argument censor). When using <code>estimator="AIPTW"</code> (double robust estimator), a model for the outcome and the treatment should be provided (argument event and treatment), as well as for the censoring (if any, argument censor).
strata	[character] Strata variable on which to compute the average risk. Incompatible with treatment. Experimental.
contrasts	[character vector] levels of the treatment variable for which the risks should be assessed and compared. Default is to consider all levels.
allContrasts	[2-row character matrix] levels of the treatment variable to be compared. Default is to consider all pairwise comparisons.
times	[numeric vector] Time points at which to evaluate average treatment effects.
cause	[integer/character] the cause of interest.
landmark	for models with time-dependent covariates the landmark time(s) of evaluation. In this case, argument time may only be one value and for the prediction of risks it is assumed that the covariates do not change between landmark and landmark+time.
se	[logical] If TRUE compute and add the standard errors to the output.
iid	[logical] If TRUE compute and add the influence function to the output.
known.nuisance	[logical] If FALSE the uncertainty related to the estimation of the nuisance parameters is ignored. This greatly simplifies computations but requires to use a double robust estimator. The resulting standard error is known to be consistent when all event, treatment, and censoring models are valid.
band	[logical] If TRUE compute and add the quantiles for the confidence bands to the output.
B	[integer, >0] the number of bootstrap replications used to compute the confidence intervals. If it equals 0, then the influence function is used to compute Wald-type confidence intervals/bands.
seed	[integer, >0] seed number used to generate seeds for bootstrap and to achieve reproducible results.
handler	[character] Parallel handler for bootstrap. "foreach" is the default and the only option on Windows. It uses <code>parallel</code> to create a cluster. Other operating systems can use <code>"mclapply"</code> . This argument is ignored when <code>mc.cores=1</code> and <code>cl=NULL</code> .

<code>mc.cores</code>	[integer, >0] The number of cores to use, i.e., the upper limit for the number of child processes that run simultaneously. Passed to <code>parallel::mclapply</code> or <code>parallel::makeCluster</code> . The option is initialized from environment variable <code>mc_cores</code> if set.
<code>cl</code>	A parallel socket cluster used to perform cluster calculation in parallel (output by <code>parallel::makeCluster</code>). The packages necessary to run the computations (e.g. <code>riskRegression</code>) must already be loaded on each worker. Only used when <code>handler="foreach"</code> .
<code>verbose</code>	[logical] If TRUE inform about estimated run time.
<code>...</code>	passed to <code>predictRisk</code>

Details

Argument **event**:

- IPTW estimator: a character vector indicating the event and status variables.
- G-formula or AIPTW estimator: a survival model (e.g. Cox `"survival::coxph"`, Kaplan Meier `"prodlm::prodlm"`), a competing risk model (e.g. cause-specific Cox `"riskRegression::CSC"`), a logistic model (e.g. `"stats::glm"` when argument `tensor` is NULL otherwise `"riskRegression::wglm"`). Other models can be specified, provided a suitable `predictRisk` method exists, but the standard error should be computed using non-parametric bootstrap, as the influence function of the estimator will typically not be available.

Argument **treatment**:

- G-formula estimator: a character indicating the treatment variable.
- IPTW or AIPTW estimator: a `"stats::glm"` model with family `"binomial"` (two treatment options) or a `"nnet::multinom"` (more than two treatment options).

Argument **tensor**:

- G-formula estimator: NULL
- IPTW or AIPTW estimator: NULL if no censoring and otherwise a survival model (e.g. Cox `"survival::coxph"`, Kaplan Meier `"prodlm::prodlm"`)

Argument **estimator**: when set to `"AIPTW"` with argument `event` being IPCW logistic model (`"riskRegression::wglm"`), the integral term w.r.t. to the martingale of the censoring process is not computed, i.e. using the notation of Ozenne et al. (2020) an AIPTW,IPCW estimator instead of an AIPTW,AIPCW is evaluated.

In presence of censoring, the computation time and memory usage for the evaluation of the AIPTW estimator and its uncertainty do not scale well with the number of observations (n) or the number of unique timepoints (T). Point estimation involves n by T matrices, influence function involves n by T by n arrays.

- for large datasets (e.g. $n > 5000$), bootstrap is recommended as the memory need for the influence function is likely prohibitive.

- it is possible to decrease the memory usage for the point estimation by setting the (hidden) argument `store=c(size.split=1000)`. The integral term of the AIPTW estimator is then evaluated for 1000 observations at a time, i.e. involving matrices of size 1000 by T instead of n by T. This may lead to increased computation time.
- reducing the number of unique timepoints (e.g. by rounding them) will lead to an approximate estimation procedure that is less demanding both in term of computation and memory. The resulting estimator will be more variable than the one based on the original timepoints (i.e. wider confidence intervals).

Author(s)

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References

Brice Maxime Hugues Ozenne, Thomas Harder Scheike, Laila Staerk, and Thomas Alexander Gerds. On the estimation of average treatment effects with right-censored time to event outcome and competing risks. *Biometrical Journal*, 62 (3):751–763, 2020.

See Also

[autoplot.ate](#) for a graphical representation the standardized risks.

[coef.ate](#) to output estimates for the the average risk, or difference in average risks, or ratio between average risks.

[confint.ate](#) to output a list containing all estimates (average & difference & ratio) with their confidence intervals and p-values.

[model.tables.ate](#) to output a data.frame containing one type of estimates (average or difference or ratio) with its confidence intervals and p-values.

[summary.ate](#) for displaying in the console a summary of the results.

Examples

```
library(survival)
library(prodlim)
library(data.table)

#####
#### Survival settings ####
#### ATE with Cox model ####
#####

#### generate data ####
n <- 100
set.seed(10)
dtS <- sampleData(n, outcome="survival")
dtS$time <- round(dtS$time,1)
dtS$X1 <- factor(rbinom(n, prob = c(0.3,0.4) , size = 2), labels = paste0("T",0:2))

##### estimate the survival model #####
## Cox proportional hazard model
fit.Cox <- coxph(Surv(time,event)~ X1+X2, data=dtS, y=TRUE, x=TRUE)
```

```

## Kaplan Meier - same as copxh(Surv(time,event)~ strata(X1)+strata(X2), ties = "breslow")
fit.KM <- prodlim(Hist(time,event)~ X1+X2, data=dtS)

##### compute the ATE #####
## at times 5, 6, 7, and 8 using X1 as the treatment variable
## standard error computed using the influence function
## confidence intervals / p-values based on asymptotic results
ateFit1a <- ate(fit.Cox, treatment = "X1", times = 5:8, data = dtS)
summary(ateFit1a)
model.tables(ateFit1a, type = "meanRisk")
model.tables(ateFit1a, type = "diffRisk")
model.tables(ateFit1a, type = "ratioRisk")

## relaxing PH assumption using a stratified model
ateFit1a.NPH <- ate(fit.KM, treatment = "X1", times = 5:8, data = dtS,
                  product.limit = FALSE)
model.tables(ateFit1a.NPH, times = 5) ## no PH assumption
model.tables(ateFit1a, times = 5) ## PH assumption

## Not run:
##### ATE with confidence bands #####
## same as before with in addition the confidence bands / adjusted p-values
## (argument band = TRUE)
ateFit1b <- ate(fit.Cox, treatment = "X1", times = 5:8, data = dtS, band = TRUE)
summary(ateFit1b)

## by default bands/adjusted p-values computed separately for each treatment modality
summary(ateFit1b, band = 1,
       se = FALSE, type = "diffRisk", short = TRUE, quantile = TRUE)
## adjustment over treatment and time using the band argument of confint
summary(ateFit1b, band = 2,
       se = FALSE, type = "diffRisk", short = TRUE, quantile = TRUE)

## End(Not run)

## Not run:
##### ATE with non-parametric bootstrap #####
## confidence intervals / p-values computed using 1000 bootstrap samples
## (argument se = TRUE and B = 1000)
ateFit1c <- ate(fit.Cox, treatment = "X1", times = 5:8, data = dtS,
              se = TRUE, B = 50, handler = "mclapply")
## NOTE: for real applications 50 bootstrap samples is not enough

## same but using 2 cpus for generating and analyzing the bootstrap samples
## (parallel computation, argument mc.cores = 2)
ateFit1d <- ate(fit.Cox, treatment = "X1", times = 5:8, data = dtS,
              se = TRUE, B = 50, mc.cores = 2)

## manually defining the cluster to be used
## useful when specific packages need to be loaded in each cluster
fit.CoxNL <- coxph(Surv(time,event)~ X1+X2+rcs(X6),data=dtS,y=TRUE,x=TRUE)

cl <- parallel::makeCluster(2)

```

```

parallel::clusterEvalQ(cl, library(rms))

ateFit1e <- ate(fit.CoxNL, treatment = "X1", times = 5:8, data = dtS,
              se = TRUE, B = 50, handler = "foreach", cl = cl)
parallel::stopCluster(cl)

## End(Not run)

#####
#### Competing risks settings      ####
#### ATE with cause specific Cox regression ####
#####

#### generate data ####
n <- 500
set.seed(10)
dt <- sampleData(n, outcome="competing.risks")
dt$X1 <- factor(rbinom(n, prob = c(0.2,0.3) , size = 2), labels = paste0("T",0:2))

#### estimate cause specific Cox model ####
fit.CR <- CSC(Hist(time,event)~ X1+X8,data=dt,cause=1)

#### compute the ATE ####
## at times 1, 5, 10
## using X1 as the treatment variable
ateFit2a <- ate(fit.CR, treatment = "X1", times = c(1,5,10), data = dt,
              cause = 1, se = TRUE, band = TRUE)
summary(ateFit2a)
as.data.table(ateFit2a)

#####
#### Survival settings without censoring ####
#### ATE with glm      ####
#####

#### generate data ####
n <- 100
dtB <- sampleData(n, outcome="binary")
dtB[, X2 := as.numeric(X2)]

##### estimate a logistic regression model ####
fit.glm <- glm(formula = Y ~ X1+X2, data=dtB, family = "binomial")

#### compute the ATE ####
## using X1 as the treatment variable
## only point estimate (argument se = FALSE)
ateFit1a <- ate(fit.glm, treatment = "X1", data = dtB, se = FALSE)
ateFit1a

## Not run:
## with confidence intervals
ateFit1b <- ate(fit.glm, data = dtB, treatment = "X1",

```

```

        times = 5) ## just for having a nice output not used in computations
summary(ateFit1b, short = TRUE)

## using the lava package
library(lava)
ateLava <- estimate(fit.glm, function(p, data){
  a <- p["(Intercept)"]; b <- p["X11"]; c <- p["X2"];
  R.X11 <- expit(a + b + c * data[["X2"]])
  R.X10 <- expit(a + c * data[["X2"]])
  list(risk0=R.X10,risk1=R.X11,riskdiff=R.X11-R.X10)},
  average=TRUE)
ateLava

## End(Not run)

#####
#### Survival settings ####
#### ATE with glm      ####
#####

## see wglm for handling right-censoring with glm

#####
#### Double robust estimator ####
#####

## Not run:
## generate data
n <- 500
set.seed(10)
dt <- sampleData(n, outcome="competing.risks")
dt$X1 <- factor(rbinom(n, prob = c(0.4) , size = 1), labels = paste0("T",0:1))

## working models
m.event <- CSC(Hist(time,event)~ X1+X2+X3+X5+X8,data=dt)
m.censor <- coxph(Surv(time,event==0)~ X1+X2+X3+X5+X8,data=dt, x = TRUE, y = TRUE)
m.treatment <- glm(X1~X2+X3+X5+X8,data=dt,family=binomial(link="logit"))

## prediction + average
ateRobust <- ate(event = m.event,
  treatment = m.treatment,
  censor = m.censor,
  data = dt, times = 5:10,
  cause = 1)
summary(ateRobust)

## compare various estimators
system.time( ## about 1.5s
ateRobust2 <- ate(event = m.event,
  treatment = m.treatment,
  censor = m.censor,
  estimator = c("GFORMULA","IPTW","AIPTW"),
  data = dt, times = c(5:10),

```



```

        cause = 1, se = TRUE)
)
as.data.table(ateRobust2, type = "meanRisk")
as.data.table(ateRobust2, type = "diffRisk")

## reduce memory load by computing the integral term
## on only 100 observations at a time (only relevant when iid = FALSE)
ateRobust3 <- ate(event = m.event,
  treatment = m.treatment,
  censor = m.censor,
  data = dt, times = 5:10,
  cause = 1, se = FALSE,
  store = c("size.split" = 100), verbose = 2)
coef(ateRobust3) - coef(ateRobust) ## same

## approximation to speed up calculations
dt$time.round <- round(dt$time/0.5)*0.5 ## round to the nearest half
dt$time.round[dt$time.round==0] <- 0.1 ## ensure strictly positive event times
mRound.event <- CSC(Hist(time.round,event)~ X1+X2+X3+X5+X8,data=dt)
mRound.censor <- coxph(Surv(time.round,event==0)~ X1+X2+X3+X5+X8,data=dt, x = TRUE, y = TRUE)
system.time( ## about 0.4s
ateRobustFast <- ate(event = mRound.event, treatment = m.treatment,
  censor = mRound.censor,
  estimator = c("GFORMULA","IPTW","AIPTW"), product.limit = FALSE,
  data = dt, times = c(5:10), cause = 1, se = TRUE)
)
coef(ateRobustFast) - coef(ateRobust) ## inaccuracy

## End(Not run)

#####
#### time-dependent covariates ####
#####

## Not run:
#### example 1 (survival)
## data management: split trajectories
vet2 <- survSplit(Surv(time, status) ~., data = veteran,
  cut=c(60, 120), episode ="timegroup")

## fit Cox model
fit.TD <- coxph(Surv(tstart, time, status) ~ celltype + karno + age + trt,
  data= vet2, x = TRUE)

## run ATE with bootstrap for uncertainty quantification
set.seed(16)
resVet <- ate(fit.TD, treatment = "celltype", times = 5, data = vet2,
  formula=Hist(entry=tstart,time=time,event=status)~1,
  landmark = c(0,30,60,90), B = 50)
summary(resVet)

## for reference
fit.Ref <- coxph(Surv(time, status) ~ celltype + karno + age + trt,

```

```

      data= veteran, x = TRUE)
ateRef <- ate(fit.Ref, treatment = "celltype", times = c(5,35), data = veteran)
## exactly the same when landmark = 0
confint(ateRef)$meanRisk[1]
confint(resVet)$meanRisk[1]
## very different with landmark != 0
## since one condition on being alive at the landmark time
confint(ateRef)$meanRisk[2]
suppressWarnings(confint(resVet)$meanRisk[2])

## End(Not run)

## Not run:
#### example 2 (competing risks)
## generate data
set.seed(137)
d <- sampleDataTD(127)
d[,status:=1*(event==1)]
d[,X3:=as.factor(X3)]
## fit cause specific Cox model
cscTD <- CSC(Hist(time=time, event=event,entry=start) ~ X3+X5+X6+X8, data=d)
## run ATE
set.seed(16)
resTD <- ate(cscTD,formula=Hist(entry=start,time=time,event=event)~1,
            data = d, treatment = "X3", contrasts = NULL,
            times=.5,verbose=1,
            landmark = c(0,0.5,1), cause = 1, B = 20, se = 1,
            band = FALSE, mc.cores=1)
resTD

## End(Not run)

```

autoplot.ate

Plot Average Risks

Description

Plot average risks.

Usage

```

## S3 method for class 'ate'
autoplot(
  object,
  type = "meanRisk",
  first.derivative = FALSE,
  estimator = object$estimator[1],
  ci = object$inference$ci,
  band = object$inference$band,

```

```

plot.type = "1",
plot = TRUE,
smooth = FALSE,
digits = 2,
alpha = NA,
ylab = NULL,
...
)

```

Arguments

object	Object obtained with the function <code>ate</code> .
type	[character vector] what to displayed. Can be "meanRisk" to display the risks specific to each treatment group, "diffRisk" to display the difference in risks between treatment groups, or "ratioRisk" to display the ratio of risks between treatment groups,.
first.derivative	[logical] If TRUE, display the first derivative over time of the risks/risk differences/risk ratios. (confidence intervals are obtained via simulation).
estimator	[character] The type of estimator relative to which the risks should be displayed.
ci	[logical] If TRUE display the confidence intervals for the average risks.
band	[logical] If TRUE display the confidence bands for the average risks.
plot.type	[character] Type of plot to be used. <code>plot.type="2"</code> is useful when looking simultaneous at all eventtimes. Otherwise use <code>plot.type="1"</code> .
plot	[logical] Should the graphic be plotted.
smooth	[logical] Should a smooth version of the risk function be plotted instead of a simple function?
digits	[integer, >0] Number of decimal places.
alpha	[numeric, 0-1] Transparency of the confidence bands. Argument passed to <code>ggplot2::geom_ribbon</code> .
ylab	[character] Label for the y axis.
...	Additional parameters to customize the display.

Value

Invisible. A list containing:

- `plot`: the `ggplot` object.
- `data`: the data used to create the plot.

See Also

[ate](#) to compute average risks.

Examples

```

library(survival)
library(ggplot2)

#### simulate data ####
n <- 1e2
set.seed(10)
dtS <- sampleData(n,outcome="survival")
seqTimes <- c(0,sort(dtS$time[dtS$event==1]),max(dtS$time))

#### Cox model ####
fit <- coxph(formula = Surv(time,event)~ X1+X2,data=dtS,y=TRUE,x=TRUE)

#### plot.type = 1: for few timepoints ####
ateFit <- ate(fit, data = dtS, treatment = "X1",
             times = c(1,2,5,10), se = TRUE, band = TRUE)
ggplot2::autoplot(ateFit)
## Not run:
ggplot2::autoplot(ateFit, band = FALSE)
ggplot2::autoplot(ateFit, type = "diffRisk")
ggplot2::autoplot(ateFit, type = "ratioRisk")

## End(Not run)

#### plot.type = 2: when looking at all jump times ####
## Not run:
ateFit <- ate(fit, data = dtS, treatment = "X1",
             times = seqTimes, se = TRUE, band = TRUE)

ggplot2::autoplot(ateFit, plot.type = "2")

## customize plot
outGG <- ggplot2::autoplot(ateFit, plot.type = "2", alpha = 0.25)
outGG$plot + facet_wrap(~X1, labeller = label_both)

## Looking at the difference after smoothing
outGGS <- ggplot2::autoplot(ateFit, plot.type = "2", alpha = NA, smooth = TRUE)
outGGS$plot + facet_wrap(~X1, labeller = label_both)

## first derivative
## (computation of the confidence intervals takes time)
## (based on simulation - n.sim parameter)
ggplot2::autoplot(ateFit, plot.type = "2", smooth = TRUE,
                 band = FALSE, type = "diffRisk")
ggplot2::autoplot(ateFit, plot.type = "2", smooth = TRUE, first.derivative = TRUE,
                 band = FALSE, type = "diffRisk")

## End(Not run)

```

 autoplot.predictCox *Plot Predictions From a Cox Model*

Description

Plot predictions from a Cox model.

Usage

```
## S3 method for class 'predictCox'
autoplot(
  object,
  type = NULL,
  ci = object$se,
  band = object$band,
  plot = TRUE,
  smooth = NULL,
  digits = 2,
  alpha = NA,
  group.by = "row",
  reduce.data = FALSE,
  ylab = NULL,
  first.derivative = FALSE,
  ...
)
```

Arguments

object	Object obtained with the function predictCox.
type	[character] The type of predicted value to display. Choices are: "hazard" the hazard function, "cumhazard" the cumulative hazard function, or "survival" the survival function.
ci	[logical] If TRUE display the confidence intervals for the predictions.
band	[logical] If TRUE display the confidence bands for the predictions.
plot	[logical] Should the graphic be plotted.
smooth	[logical] Should a smooth version of the risk function be plotted instead of a simple function?
digits	[integer] Number of decimal places when displaying the values of the covariates in the caption.
alpha	[numeric, 0-1] Transparency of the confidence bands. Argument passed to ggplot2::geom_ribbon.
group.by	[character] The grouping factor used to color the prediction curves. Can be "row", "strata", or "covariates".
reduce.data	[logical] If TRUE only the covariates that does take identical values for all observations are displayed.

ylab [character] Label for the y axis.
 first.derivative [logical] If TRUE, display the first derivative over time of the risks/risk differences/risk ratios. (confidence intervals are obtained via simulation).
 ... Additional parameters to customize the display.

Value

Invisible. A list containing:

- plot: the ggplot object.
- data: the data used to create the plot.

See Also

[predictCox](#) to compute cumulative hazard and survival based on a Cox model.

Examples

```
library(survival)
library(ggplot2)

#### simulate data ####
set.seed(10)
d <- sampleData(1e2, outcome = "survival")
seqTau <- c(0,sort(unique(d$time[d$event==1])), max(d$time))

#### Cox model ####
m.cox <- coxph(Surv(time,event)~ X1 + X2 + X3,
               data = d, x = TRUE, y = TRUE)

## display baseline hazard
e.basehaz <- predictCox(m.cox)
autoplot(e.basehaz, type = "cumhazard")
## Not run:
autoplot(e.basehaz, type = "cumhazard", size.point = 0) ## without points
autoplot(e.basehaz, type = "cumhazard", smooth = TRUE)
autoplot(e.basehaz, type = "cumhazard", smooth = TRUE, first.derivative = TRUE)

## End(Not run)

## display baseline hazard with type of event
## Not run:
e.basehaz <- predictCox(m.cox, keep.newdata = TRUE)
autoplot(e.basehaz, type = "cumhazard")
autoplot(e.basehaz, type = "cumhazard", shape.point = c(3,NA))

## End(Not run)

## display predicted survival
## Not run:
```

```

pred.cox <- predictCox(m.cox, newdata = d[1:2,],
  times = seqTau, type = "survival", keep.newdata = TRUE)
autoplot(pred.cox)
autoplot(pred.cox, smooth = TRUE)
autoplot(pred.cox, group.by = "covariates")
autoplot(pred.cox, group.by = "covariates", reduce.data = TRUE)
autoplot(pred.cox, group.by = "X1", reduce.data = TRUE)

## End(Not run)

## predictions with confidence interval/bands
## Not run:
pred.cox <- predictCox(m.cox, newdata = d[1:2,drop=FALSE],
  times = seqTau, type = "survival", band = TRUE, se = TRUE, keep.newdata = TRUE)
res <- autoplot(pred.cox, ci = TRUE, band = TRUE, plot = FALSE)
res$plot + facet_wrap(~row)
res2 <- autoplot(pred.cox, ci = TRUE, band = TRUE, alpha = 0.1, plot = FALSE)
res2$plot + facet_wrap(~row)

## End(Not run)

#### Stratified Cox model ####
## Not run:
m.cox.strata <- coxph(Surv(time,event)~ strata(X1) + strata(X2) + X3 + X4,
  data = d, x = TRUE, y = TRUE)

## baseline hazard
pred.baseline <- predictCox(m.cox.strata, keep.newdata = TRUE, type = "survival")
res <- autoplot(pred.baseline)
res$plot + facet_wrap(~strata, labeller = label_both)

## predictions
pred.cox.strata <- predictCox(m.cox.strata, newdata = d[1:3,drop=FALSE],
  time = seqTau, keep.newdata = TRUE, se = TRUE)

res2 <- autoplot(pred.cox.strata, type = "survival", group.by = "strata", plot = FALSE)
res2$plot + facet_wrap(~strata, labeller = label_both) + theme(legend.position="bottom")

## smooth version
autoplot(pred.cox.strata, type = "survival", group.by = "strata", smooth = TRUE, ci = FALSE)

## End(Not run)

#### Cox model with splines ####
## Not run:
require(splines)
m.cox.spline <- coxph(Surv(time,event)~ X1 + X2 + ns(X6,4),
  data = d, x = TRUE, y = TRUE)
grid <- data.frame(X1 = factor(0,0:1), X2 = factor(0,0:1),
  X6 = seq(min(d$X6),max(d$X6), length.out = 100))
pred.spline <- predictCox(m.cox.spline, newdata = grid, keep.newdata = TRUE,
  se = TRUE, band = TRUE, centered = TRUE, type = "lp")
autoplot(pred.spline, group.by = "X6")

```

```

autoplot(pred.spline, group.by = "X6", alpha = 0.5)

grid2 <- data.frame(X1 = factor(1,0:1), X2 = factor(0,0:1),
                   X6 = seq(min(d$X6),max(d$X6), length.out = 100))
pred.spline <- predictCox(m.cox.spline, newdata = rbind(grid,grid2), keep.newdata = TRUE,
                        se = TRUE, band = TRUE, centered = TRUE, type = "lp")
autoplot(pred.spline, group.by = c("X6","X1"), alpha = 0.5, plot = FALSE)$plot + facet_wrap(~X1)

## End(Not run)

```

autoplot.predictCSC *Plot Predictions From a Cause-specific Cox Proportional Hazard Regression*

Description

Plot predictions from a Cause-specific Cox proportional hazard regression.

Usage

```

## S3 method for class 'predictCSC'
autoplot(
  object,
  ci = object$se,
  band = object$band,
  plot = TRUE,
  smooth = FALSE,
  digits = 2,
  alpha = NA,
  group.by = "row",
  reduce.data = FALSE,
  ...
)

```

Arguments

object	Object obtained with the function predictCox.
ci	[logical] If TRUE display the confidence intervals for the predictions.
band	[logical] If TRUE display the confidence bands for the predictions.
plot	[logical] Should the graphic be plotted.
smooth	[logical] Should a smooth version of the risk function be plotted instead of a simple function?
digits	[integer] Number of decimal places.
alpha	[numeric, 0-1] Transparency of the confidence bands. Argument passed to ggplot2::geom_ribbon.

group.by	[character] The grouping factor used to color the prediction curves. Can be "row", "strata", or "covariates".
reduce.data	[logical] If TRUE only the covariates that does take indential values for all observations are displayed.
...	Additional parameters to cutomize the display.

Value

Invisible. A list containing:

- plot: the ggplot object.
- data: the data used to create the plot.

See Also

[predict.CauseSpecificCox](#) to compute risks based on a CSC model.

Examples

```
library(survival)
library(rms)
library(ggplot2)
library(proplim)

#### simulate data ####
set.seed(10)
d <- sampleData(1e2, outcome = "competing.risks")
seqTau <- c(0,unique(sort(d[d$event==1,time])), max(d$time))

#### CSC model ####
m.CSC <- CSC(Hist(time,event)~ X1 + X2 + X6, data = d)

pred.CSC <- predict(m.CSC, newdata = d[1:2,], time = seqTau, cause = 1, band = TRUE)
autoplot(pred.CSC, alpha = 0.2)

#### stratified CSC model ####
m.SCSC <- CSC(Hist(time,event)~ strata(X1) + strata(X2) + X6,
             data = d)
pred.SCSC <- predict(m.SCSC, time = seqTau, newdata = d[1:4,],
                    cause = 1, keep.newdata = TRUE, keep.strata = TRUE)
autoplot(pred.SCSC, group.by = "strata")
```

autoplot.Score

ggplot AUC curve

Description

ggplot AUC curves

Usage

```
## S3 method for class 'Score'
autoplot(
  object,
  models,
  type = "score",
  lwd = 2,
  xlim,
  ylim,
  axes = TRUE,
  conf.int = FALSE,
  ...
)
```

Arguments

object	Object obtained with <code>Score.list</code>
models	Choice of models to plot
type	Character. Either "score" to show AUC or "contrasts" to show differences between AUC.
lwd	Line width
xlim	Limits for x-axis
ylim	Limits for y-axis
axes	Logical. If TRUE draw axes.
conf.int	Logical. If TRUE draw confidence shadows.
...	Not yet used

Examples

```
library(survival)
library(ggplot2)
set.seed(10)
d=sampleData(100,outcome="survival")
nd=sampleData(100,outcome="survival")
f1=coxph(Surv(time,event)~X1+X6+X8,data=d,x=TRUE,y=TRUE)
f2=coxph(Surv(time,event)~X2+X5+X9,data=d,x=TRUE,y=TRUE)
xx=Score(list(f1,f2), formula=Surv(time,event)~1,
data=nd, metrics="auc", null.model=FALSE, times=seq(3:10))
g <- autoplot(xx)
print(g)
aucgraph <- plotAUC(xx)
plotAUC(xx,conf.int=TRUE)
plotAUC(xx,which="contrasts")
plotAUC(xx,which="contrasts",conf.int=TRUE)
```

Description

C++ function to estimate the baseline hazard from a Cox Model

Usage

```
baseHaz_cpp(  
    starttimes,  
    stoptimes,  
    status,  
    eXb,  
    strata,  
    predtimes,  
    emaxtimes,  
    nPatients,  
    nStrata,  
    cause,  
    Efron,  
    reverse  
)
```

Arguments

starttimes	a vector of times (begin at risk period).
stoptimes	a vector of times (end at risk period).
status	a vector indicating censoring or event.
eXb	a numeric vector (exponential of the linear predictor).
strata	a vector of integers (index of the strata for each observation).
predtimes	a vector of times (time at which to evaluate the hazard). Must be sorted.
emaxtimes	another vector of times, one per strata (last observation time in each strata).
nPatients	number of observations.
nStrata	number of strata
cause	the status value corresponding to event.
Efron	whether Efron or Breslow estimator should be used in presence of ties.
reverse	whether censoring occurs before events in presence of ties.

Details

WARNING stoptimes status eXb and strata must be sorted by strata, stoptimes, and status

boot2pvalue

*Compute the p.value from the distribution under H1***Description**

Compute the p.value associated with the estimated statistic using a bootstrap sample of its distribution under H1.

Usage

```
boot2pvalue(
  x,
  null,
  estimate = NULL,
  alternative = "two.sided",
  FUN.ci = quantileCI,
  tol = .Machine$double.eps^0.5
)
```

Arguments

x	[numeric vector] a vector of bootstrap estimates of the statistic.
null	[numeric] value of the statistic under the null hypothesis.
estimate	[numeric] the estimated statistic.
alternative	[character] a character string specifying the alternative hypothesis, must be one of "two.sided" (default), "greater" or "less".
FUN.ci	[function] the function used to compute the confidence interval. Must take x, alternative, conf.level and sign.estimate as arguments and only return the relevant limit (either upper or lower) of the confidence interval.
tol	[numeric] the absolute convergence tolerance.

Details

For test statistic close to 0, this function returns 1.

For positive test statistic, this function search the quantile alpha such that:

- $\text{quantile}(x, \text{probs} = \alpha) = 0$ when the argument alternative is set to "greater".
- $\text{quantile}(x, \text{probs} = 0.5 * \alpha) = 0$ when the argument alternative is set to "two.sided".

If the argument alternative is set to "less", it returns 1.

For negative test statistic, this function search the quantile alpha such that:

- `quantile(x, probs = 1-alpha=0` when the argument `alternative` is set to `"less"`.
- `quantile(x, probs = 1-0.5*alpha=0` when the argument `alternative` is set to `"two.sided"`.

If the argument `alternative` is set to `"greater"`, it returns 1.

Examples

```
set.seed(10)

#### no effect ####
x <- rnorm(1e3)
boot2pvalue(x, null = 0, estimate = mean(x), alternative = "two.sided")
## expected value of 1
boot2pvalue(x, null = 0, estimate = mean(x), alternative = "greater")
## expected value of 0.5
boot2pvalue(x, null = 0, estimate = mean(x), alternative = "less")
## expected value of 0.5

#### positive effect ####
x <- rnorm(1e3, mean = 1)
boot2pvalue(x, null = 0, estimate = 1, alternative = "two.sided")
## expected value of 0.32 = 2*pnorm(q = 0, mean = -1) = 2*mean(x<=0)
boot2pvalue(x, null = 0, estimate = 1, alternative = "greater")
## expected value of 0.16 = pnorm(q = 0, mean = 1) = mean(x<=0)
boot2pvalue(x, null = 0, estimate = 1, alternative = "less")
## expected value of 0.84 = 1-pnorm(q = 0, mean = 1) = mean(x>=0)

#### negative effect ####
x <- rnorm(1e3, mean = -1)
boot2pvalue(x, null = 0, estimate = -1, alternative = "two.sided")
## expected value of 0.32 = 2*(1-pnorm(q = 0, mean = -1)) = 2*mean(x>=0)
boot2pvalue(x, null = 0, estimate = -1, alternative = "greater")
## expected value of 0.84 = pnorm(q = 0, mean = -1) = mean(x<=0)
boot2pvalue(x, null = 0, estimate = -1, alternative = "less") # pnorm(q = 0, mean = -1)
## expected value of 0.16 = 1-pnorm(q = 0, mean = -1) = mean(x>=0)
```

boxplot.Score

Boxplot risk quantiles

Description

Retrospective boxplots of risk quantiles conditional on outcome

Usage

```
## S3 method for class 'Score'
boxplot(
  x,
  model,
```

```

reference,
type = "risk",
timepoint,
overall = 1L,
lwd = 3,
xlim,
xlab = "",
main,
outcome.label,
outcome.label.offset = 0,
event.labels,
refline = (type != "risk"),
add = FALSE,
...
)

```

Arguments

x	Score object obtained by calling function Score.
model	Choice of risk prediction model
reference	Choice of reference risk prediction model for calculation of risk differences.
type	Either "risk" for predicted risks or "diff" for differences between predicted risks.
timepoint	time point specifying the prediction horizon
overall	Logical. Tag to be documented.
lwd	line width
xlim	x-axis limits
xlab	x-axis label
main	title of plot
outcome.label	Title label for column which shows the outcome status
outcome.label.offset	Vertical offset for outcome.label
event.labels	Labels for the different events (causes).
refline	Logical, for type="diff" only. If TRUE draw a red vertical line at 0.
add	Logical. Tag to be documented.
...	not used

Examples

```

# binary outcome
library(data.table)
library(prodlim)
set.seed(10)
db=sampleData(40,outcome="binary")
fitconv=glm(Y~X3+X5,data=db,family=binomial)

```

```

fitnew=glm(Y~X1+X3+X5+X6+X7,data=db,family=binomial)
x=Score(list(new=fitnew,conv=fitconv),
         formula=Y~1,contrasts=list(c(2,1)),
         data=db,plots="box",null.model=FALSE)
boxplot(x)

# survival outcome
library(survival)
ds=sampleData(40,outcome="survival")
fit=coxph(Surv(time,event)~X6+X9,data=ds,x=TRUE,y=TRUE)
## Not run:
scoreobj=Score(list("Cox"=fit),
               formula=Hist(time,event)~1, data=ds,
               metrics=NULL, plots="box",
               times=c(1,5),null.model=FALSE)
boxplot(scoreobj,timepoint=5)
boxplot(scoreobj,timepoint=1)

## End(Not run)

# competing risks outcome
library(survival)
data(Melanoma, package = "riskRegression")
fit = CSC(Hist(time,event,cens.code="censored")~invasion+age+sex,data=Melanoma)
scoreobj=Score(list("CSC"=fit),
               formula=Hist(time,event,cens.code="censored")~1,
               data=Melanoma,plots="box",times=5*365.25,null.model=FALSE)
par(mar=c(4,12,4,4))
boxplot(scoreobj,timepoint=5*365.25)

# more than 2 competing risks
m=lava::lvm(~X1+X2+X3)
lava::distribution(m, "eventtime1") <- lava::coxWeibull.lvm(scale = 1/100)
lava::distribution(m, "eventtime2") <- lava::coxWeibull.lvm(scale = 1/100)
lava::distribution(m, "eventtime3") <- lava::coxWeibull.lvm(scale = 1/100)
lava::distribution(m, "censtime") <- lava::coxWeibull.lvm(scale = 1/100)
lava::regression(m,eventtime2~X3)=1.3
m <- lava::eventTime(m,
time ~ min(eventtime1 = 1, eventtime2 = 2, eventtime3 = 3, censtime = 0), "event")
set.seed(101)
dcr=as.data.table(lava::sim(m,101))
fit = CSC(Hist(time,event)~X1+X2+X3,data=dcr)
scoreobj=Score(list("my model"=fit),
               formula=Hist(time,event)~1,
               data=dcr,plots="box",times=5,null.model=FALSE)
boxplot(scoreobj)

```

Description

Compute the standard error associated to the predictions from Cox regression model using a first order von Mises expansion of the functional (cumulative hazard or survival).

Usage

```
calcSeCox(
  object,
  times,
  nTimes,
  type,
  diag,
  Lambda0,
  object.n,
  object.time,
  object.eXb,
  object.strata,
  nStrata,
  new.n,
  new.eXb,
  new.LPdata,
  new.strata,
  new.survival,
  nVar.lp,
  export,
  store.iid
)
```

Arguments

<code>object</code>	The fitted Cox regression model object either obtained with <code>coxph</code> (survival package) or <code>cph</code> (rms package).
<code>times</code>	Vector of times at which to return the estimated hazard/survival.
<code>nTimes</code>	the length of the argument <code>times</code> .
<code>type</code>	One or several strings that match (either in lower or upper case or mixtures) one or several of the strings "hazard", "cumhazard", "survival".
<code>diag</code>	[logical] when FALSE the hazard/cumulative hazard/survival for all observations at all times is computed, otherwise it is only computed for the <i>i</i> -th observation at the <i>i</i> -th time.
<code>Lambda0</code>	the baseline hazard estimate returned by <code>BaseHazStrata_cpp</code> .
<code>object.n</code>	the number of observations in the dataset used to estimate the object.
<code>object.time</code>	the time to event of the observations used to estimate the object.
<code>object.eXb</code>	the exponential of the linear predictor relative to the observations used to estimate the object.
<code>object.strata</code>	the strata index of the observations used to estimate the object.

nStrata	the number of strata.
new.n	the number of observations for which the prediction was performed.
new.eXb	the linear predictor evaluated for the new observations.
new.LPdata	the variables involved in the linear predictor for the new observations.
new.strata	the strata indicator for the new observations.
new.survival	the survival evaluated for the new observations.
nVar.lp	the number of variables that form the linear predictor.
export	can be "iid" to return the value of the influence function for each observation. "se" to return the standard error for a given timepoint. "average.iid" to return the value of the average influence function over the observations for which the prediction was performed.
store.iid	Implementation used to estimate the influence function and the standard error. Can be "full" or "minimal". See the details section.

Details

store.iid="full" compute the influence function for each observation at each time in the argument times before computing the standard error / influence functions. store.iid="minimal" recompute for each subject specific prediction the influence function for the baseline hazard. This avoid to store all the influence functions but may lead to repeated evaluation of the influence function. This solution is therefore more efficient in memory usage but may not be in terms of computation time.

Value

A list optionally containing the standard error for the survival, cumulative hazard and hazard.

Author(s)

Brice Ozenne broz@sund.ku.dk, Thomas A. Gerds tag@biostat.ku.dk

calcSeCSC	<i>Standard error of the absolute risk predicted from cause-specific Cox models</i>
-----------	---

Description

Standard error of the absolute risk predicted from cause-specific Cox models using a first order von Mises expansion of the absolute risk functional.

Usage

```
calcSeCSC(
  object,
  cif,
  hazard,
  cumhazard,
  survival,
  object.time,
  object.maxtime,
  eXb,
  new.LPdata,
  new.strata,
  times,
  surv.type,
  ls.infoVar,
  new.n,
  cause,
  nCause,
  nVar.lp,
  export,
  store.iid,
  diag
)
```

Arguments

object	The fitted cause specific Cox model
cif	the cumulative incidence function at each prediction time for each individual.
hazard	list containing the baseline hazard for each cause in a matrix form. Columns correspond to the strata.
cumhazard	list containing the cumulative baseline hazard for each cause in a matrix form. Columns correspond to the strata.
survival	list containing the (all cause) survival in a matrix form at t-. Columns correspond to event times.
object.time	a vector containing all the events regardless to the cause.
object.maxtime	a matrix containing the latest event in the strata of the observation for each cause.
eXb	a matrix containing the exponential of the linear predictor evaluated for the new observations (rows) for each cause (columns)
new.LPdata	a list of design matrices for the new observations for each cause.
new.strata	a matrix containing the strata indicator for each observation and each cause.
times	the time points at which to evaluate the predictions.
surv.type	see the surv.type argument of CSC .
ls.infoVar	A list containing the output of <code>coxVariableName</code> for each Cox model.
new.n	the number of new observations.

cause	the cause of interest.
nCause	the number of causes.
nVar.lp	the number of variables that form the linear predictor in each Cox model
export	can be "iid" to return the value of the influence function for each observation "se" to return the standard error for a given timepoint
store.iid	the method used to compute the influence function and the standard error. Can be "full" or "minimal". See the details section.
diag	[logical] when FALSE the absolute risk/survival for all observations at all times is computed, otherwise it is only computed for the i-th observation at the i-th time.

Details

Can also return the empirical influence function of the functionals cumulative hazard or survival or the sum over the observations of the empirical influence function.

`store.iid="full"` compute the influence function for each observation at each time in the argument `times` before computing the standard error / influence functions. `store.iid="minimal"` recompute for each subject specific prediction the influence function for the baseline hazard. This avoid to store all the influence functions but may lead to repeated evaluation of the influence function. This solution is therefore efficient more efficient in memory usage but may not be in term of computation time.

Cforest	<i>S3-wrapper function for cforest from the party package</i>
---------	---

Description

S3-wrapper function for cforest from the party package

Usage

```
Cforest(formula, data, ...)
```

Arguments

formula	Passed on as is. See cforest of the party package
data	Passed on as is. See cforest of the party package
...	Passed on as they are. See cforest of the party package

Details

See cforest of the party package.

Value

list with two elements: cforest and call

References

Ulla B. Mogensen, Hemant Ishwaran, Thomas A. Gerds (2012). Evaluating Random Forests for Survival Analysis Using Prediction Error Curves. *Journal of Statistical Software*, 50(11), 1-23. URL <http://www.jstatsoft.org/v50/i11/>.

coef.ate

*Estimated Average Treatment Effect.***Description**

Estimated average treatment effect.

Usage

```
## S3 method for class 'ate'
coef(
  object,
  contrasts = NULL,
  times = NULL,
  estimator = NULL,
  type = NULL,
  ...
)
```

Arguments

object	A ate object, i.e. output of the ate function.
contrasts	[character vector] levels of the treatment variable for which the estimates should be assessed or compared. Default is to consider all levels.
times	[numeric vector] The timepoints at which the estimates should be displayed. Default is to consider all timepoints.
estimator	[character] The type of estimator relative to which the estimates should be displayed.
type	[character] should the average risk per treatment be displayed ("meanRisk"), or the difference in average risk between any two pairs of treatments ("diffRisk"), or the ratio in average risk between any two pairs of treatments ("ratioRisk").
...	Not used. For compatibility with the generic method.

Value

A numeric vector.

Author(s)

Brice Ozenne <broz@sund.ku.dk>

coef.CauseSpecificCox *Extract coefficients from a Cause-Specific Cox regression model*

Description

Extract coefficients from a Cause-Specific Cox regression model

Usage

```
## S3 method for class 'CauseSpecificCox'  
coef(object, ...)
```

Arguments

object	Object obtained with CSC
...	not used

coef.riskRegression *Extract coefficients from riskRegression model*

Description

Extract coefficients from riskRegression model

Usage

```
## S3 method for class 'riskRegression'  
coef(object, digits = 3, eps = 10^-4, ...)
```

Arguments

object	Object obtained with ARR or LRR or riskRegression
digits	Number of digits
eps	P-values below this number are shown as <eps
...	not used

 coef.wglm

Estimates from IPCW Logistic Regressions

Description

Display the estimated regression parameters from logistic regressions.

Usage

```
## S3 method for class 'wglm'
coef(object, times = NULL, simplify = TRUE, ...)
```

Arguments

object	a wglm object.
times	[numeric vector] time points at which the estimates should be output.
simplify	[logical] should the output be converted to a vector when only one timepoint is requested. Otherwise will always return a matrix.
...	Not used.

 colCenter_cpp

Apply - by column

Description

Fast computation of sweep(X, MARGIN = 1, FUN = "-", STATS = center)

Usage

```
colCenter_cpp(X, center)
```

Arguments

X	A matrix.
center	a numeric vector of length equal to the number of rows of x

Value

A matrix of same size as X.

Author(s)

Brice Ozenne <broz@sund.ku.dk>

Examples

```
x <- matrix(1,6,5)
sweep(x, MARGIN = 1, FUN = "-", STATS = 1:6)
colCenter_cpp(x, 1:6 )
```

colCumSum	<i>Apply cumsum in each column</i>
-----------	------------------------------------

Description

Fast computation of `apply(x,2,cumsum)`

Usage

```
colCumSum(x)
```

Arguments

x A matrix.

Value

A matrix of same size as x.

Author(s)

Thomas Alexander Gerds <tag@biostat.ku.dk>

Examples

```
x <- matrix(1:8,ncol=2)
colCumSum(x)
```

colMultiply_cpp	<i>Apply * by column</i>
-----------------	--------------------------

Description

Fast computation of `sweep(X, MARGIN = 1, FUN = "*", STATS = scale)`

Usage

```
colMultiply_cpp(X, scale)
```

Arguments

`X` A matrix.
`scale` a numeric vector of length equal to the number of rows of `x`

Value

A matrix of same size as `X`.

Author(s)

Brice Ozenne <broz@sund.ku.dk>

Examples

```
x <- matrix(1,6,5)
sweep(x, MARGIN = 1, FUN = "*", STATS = 1:6)
colMultiply_cpp(x, 1:6 )
```

colScale_cpp	<i>Apply / by column</i>
--------------	--------------------------

Description

Fast computation of `sweep(X, MARGIN = 1, FUN = "/", STATS = scale)`

Usage

```
colScale_cpp(X, scale)
```

Arguments

`X` A matrix.
`scale` a numeric vector of length equal to the number of rows of `x`

Value

A matrix of same size as `X`.

Author(s)

Brice Ozenne <broz@sund.ku.dk>

Examples

```
x <- matrix(1,6,5)
sweep(x, MARGIN = 1, FUN = "/", STATS = 1:6)
colScale_cpp(x, 1:6 )
```

confint.ate	<i>Confidence Intervals and Confidence Bands for the average treatment effect.</i>
-------------	--

Description

Confidence intervals and confidence Bands for the average treatment effect.

Usage

```
## S3 method for class 'ate'
confint(
  object,
  parm = NULL,
  level = 0.95,
  n.sim = 10000,
  estimator = object$estimator,
  contrasts = object$contrasts,
  allContrasts = object$allContrasts,
  meanRisk.transform = "none",
  diffRisk.transform = "none",
  ratioRisk.transform = "none",
  seed = NA,
  ci = object$inference$se,
  band = object$inference$band,
  p.value = TRUE,
  method.band = "maxT-simulation",
  alternative = "two.sided",
  bootci.method = "perc",
  ...
)
```

Arguments

object	A ate object, i.e. output of the ate function.
parm	not used. For compatibility with the generic method.
level	[numeric, 0-1] Level of confidence.
n.sim	[integer, >0] the number of simulations used to compute the quantiles for the confidence bands and/or perform adjustment for multiple comparisons.
estimator	[character] The type of estimator relative to which the estimates should be displayed.
contrasts	[character vector] levels of the treatment variable for which the risks should be assessed and compared. Default is to consider all levels.
allContrasts	[2-row character matrix] levels of the treatment variable to be compared. Default is to consider all pairwise comparisons.

<code>meanRisk.transform</code>	[character] the transformation used to improve coverage of the confidence intervals for the mean risk in small samples. Can be "none", "log", "loglog", "cloglog".
<code>diffRisk.transform</code>	[character] the transformation used to improve coverage of the confidence intervals for the risk difference in small samples. Can be "none", "atanh".
<code>ratioRisk.transform</code>	[character] the transformation used to improve coverage of the confidence intervals for the risk ratio in small samples. Can be "none", "log".
<code>seed</code>	[integer, >0] seed number set when performing simulation for the confidence bands. If not given or NA no seed is set.
<code>ci</code>	[logical] should the confidence intervals be computed?
<code>band</code>	[logical] should the confidence bands be computed?
<code>p.value</code>	[logical] should the p-values/adjusted p-values be computed? Requires argument <code>ci</code> and/or <code>band</code> to be TRUE.
<code>method.band</code>	[character] method used to adjust for multiple comparisons. Can be any element of <code>p.adjust.methods</code> (e.g. "holm"), "maxT-integration", or "maxT-simulation".
<code>alternative</code>	[character] a character string specifying the alternative hypothesis, must be one of "two.sided" (default), "greater" or "less".
<code>bootci.method</code>	[character] Method for constructing bootstrap confidence intervals. Either "perc" (the default), "norm", "basic", "stud", or "bca".
<code>...</code>	Not used. For compatibility with the generic method.

Details

Argument `ci`, `band`, `p.value`, `method.band`, `alternative`, `meanRisk.transform`, `diffRisk.transform`, `ratioRisk.transform` are only active when the `ate` object contains the influence function. Argument `bootci.method` is only active when the `ate` object contains bootstrap samples.

Influence function: confidence bands and confidence intervals computed via the influence function are automatically restricted to the interval of definition of the parameter (e.g. [0;1] for the average risk). Single step max adjustment for multiple comparisons, i.e. accounting for the correlation between the test statistics but not for the ordering of the tests, can be performed setting the argument `method.band` to "maxT-integration" or "maxT-simulation". The former uses numerical integration (`pmvnorm` and `qmvnorm` to perform the adjustment while the latter using simulation. Both assume that the test statistics are jointly normally distributed.

Bootstrap: confidence intervals obtained via bootstrap are computed using the `boot.ci` function of the `boot` package. p-value are obtained using test inversion method (finding the smallest confidence level such that the interval contain the null hypothesis).

Value

A list with elements

- `meanRisk`: estimated average risk (i.e. had everybody received the same treatment).
- `diffRisk`: difference in estimated average risk
- `ratioRisk`: ratio between estimated average risk

Author(s)

Brice Ozenne

Examples

```

library(survival)
library(data.table)

## ## generate data ####
set.seed(10)
d <- sampleData(70,outcome="survival")
d[, X1 := paste0("T",rbinom(.N, size = 2, prob = c(0.51)))]
## table(d$X1)

#### stratified Cox model ####
fit <- coxph(Surv(time,event)~X1 + strata(X2) + X6,
             data=d, ties="breslow", x = TRUE, y = TRUE)

#### average treatment effect ####
fit.ate <- ate(fit, treatment = "X1", times = 1:3, data = d,
              se = TRUE, iid = TRUE, band = TRUE)
summary(fit.ate)
dt.ate <- as.data.table(fit.ate)

## manual calculation of se
dd <- copy(d)
dd$X1 <- rep(factor("T0", levels = paste0("T",0:2)), NROW(dd))
out <- predictCox(fit, newdata = dd, se = TRUE, times = 1:3, average.iid = TRUE)
term1 <- -out$survival.average.iid
term2 <- sweep(1-out$survival, MARGIN = 2, FUN = "-", STATS = colMeans(1-out$survival))
sqrt(colSums((term1 + term2/NROW(d))^2))
## fit.ate$meanRisk[treatment=="T0",se]

## note
out2 <- predictCox(fit, newdata = dd, se = TRUE, times = 1:3, iid = TRUE)
mean(out2$survival.iid[1,1,])
out$survival.average.iid[1,1]

## check confidence intervals (no transformation)
dt.ate[,.(lower = pmax(0,estimate + qnorm(0.025) * se),
           lower2 = lower,
           upper = estimate + qnorm(0.975) * se,
           upper2 = upper)]

## add confidence intervals computed on the log-log scale
## and backtransformed
outCI <- confint(fit.ate,
                 meanRisk.transform = "loglog", diffRisk.transform = "atanh",
                 ratioRisk.transform = "log")
summary(outCI, type = "risk", short = TRUE)

dt.ate[type == "meanRisk", newse := se/(estimate*log(estimate))]

```

```
dt.ate[type == "meanRisk", .(lower = exp(-exp(log(-log(estimate)) - 1.96 * newse)),
                             upper = exp(-exp(log(-log(estimate)) + 1.96 * newse)))]
```

confint.influenceTest *Confidence Intervals and Confidence Bands for the Difference Between Two Estimates*

Description

Confidence intervals and confidence Bands for the difference between two estimates.

Usage

```
## S3 method for class 'influenceTest'
confint(
  object,
  parm = NULL,
  level = 0.95,
  n.sim = 10000,
  transform = "none",
  seed = NA,
  ...
)
```

Arguments

object	A influenceTest object, i.e. output of the influenceTest function.
parm	not used. For compatibility with the generic method.
level	[numeric, 0-1] Level of confidence.
n.sim	[integer, >0] the number of simulations used to compute the quantiles for the confidence bands.
transform	[character] the transformation used to improve coverage of the confidence intervals. Can be "none" or "atanh".
seed	[integer, >0] seed number set before performing simulations for the confidence bands. If not given or NA no seed is set.
...	not used.

Details

Except for the cumulative hazard, the confidence bands and confidence intervals are automatically restricted to the interval [-1;1].

Author(s)

Brice Ozenne

confint.predictCox	<i>Confidence Intervals and Confidence Bands for the predicted Survival/Cumulative Hazard</i>
--------------------	---

Description

Confidence intervals and confidence Bands for the predicted survival/cumulative Hazard.

Usage

```
## S3 method for class 'predictCox'
confint(
  object,
  parm = NULL,
  level = 0.95,
  n.sim = 10000,
  cumhazard.transform = "log",
  survival.transform = "loglog",
  seed = NA,
  ...
)
```

Arguments

object	A predictCox object, i.e. output of the predictCox function.
parm	[character] the type of predicted value for which the confidence intervals should be output. Can be "survival" or "cumhazard".
level	[numeric, 0-1] Level of confidence.
n.sim	[integer, >0] the number of simulations used to compute the quantiles for the confidence bands.
cumhazard.transform	[character] the transformation used to improve coverage of the confidence intervals for the cumulative hazard in small samples. Can be "none", "log".
survival.transform	[character] the transformation used to improve coverage of the confidence intervals for the survival in small samples. Can be "none", "log", "loglog", "cloglog".
seed	[integer, >0] seed number set before performing simulations for the confidence bands. If not given or NA no seed is set.
...	not used.

Details

The confidence bands and confidence intervals are automatically restricted to the interval of definition of the statistic, i.e. a confidence interval for the survival of [0.5;1.2] will become [0.5;1].

Author(s)

Brice Ozenne

Examples

```

library(survival)

#### generate data ####
set.seed(10)
d <- sampleData(40,outcome="survival")

#### estimate a stratified Cox model ####
fit <- coxph(Surv(time,event)~X1 + strata(X2) + X6,
             data=d, ties="breslow", x = TRUE, y = TRUE)

#### compute individual specific survival probabilities
fit.pred <- predictCox(fit, newdata=d[1:3], times=c(3,8), type = "survival",
                      se = TRUE, iid = TRUE, band = TRUE)

fit.pred

## check standard error
sqrt(rowSums(fit.pred$survival.iid[,1]^2)) ## se for individual 1

## check confidence interval
newse <- fit.pred$survival.se/(-fit.pred$survival*log(fit.pred$survival))
cbind(lower = as.double(exp(-exp(log(-log(fit.pred$survival)) + 1.96 * newse))),
      upper = as.double(exp(-exp(log(-log(fit.pred$survival)) - 1.96 * newse)))
)

#### compute confidence intervals without transformation
confint(fit.pred, survival.transform = "none")
cbind(lower = as.double(fit.pred$survival - 1.96 * fit.pred$survival.se),
      upper = as.double(fit.pred$survival + 1.96 * fit.pred$survival.se)
)

```

confint.predictCSC	<i>Confidence Intervals and Confidence Bands for the Predicted Absolute Risk (Cumulative Incidence Function)</i>
--------------------	--

Description

Confidence intervals and confidence Bands for the predicted absolute risk (cumulative incidence function).

Usage

```

## S3 method for class 'predictCSC'
confint(

```

```

    object,
    parm = NULL,
    level = 0.95,
    n.sim = 10000,
    absRisk.transform = "loglog",
    seed = NA,
    ...
)

```

Arguments

object	A predictCSC object, i.e. output of the predictCSC function.
parm	not used. For compatibility with the generic method.
level	[numeric, 0-1] Level of confidence.
n.sim	[integer, >0] the number of simulations used to compute the quantiles for the confidence bands.
absRisk.transform	[character] the transformation used to improve coverage of the confidence intervals for the predicted absolute risk in small samples. Can be "none", "log", "loglog", "cloglog".
seed	[integer, >0] seed number set before performing simulations for the confidence bands. If not given or NA no seed is set.
...	not used.

Details

The confidence bands and confidence intervals are automatically restricted to the interval [0;1].

Author(s)

Brice Ozenne

Examples

```

library(survival)
library(prodlim)
#### generate data ####
set.seed(10)
d <- sampleData(100)

#### estimate a stratified CSC model ###
fit <- CSC(Hist(time,event)~ X1 + strata(X2) + X6, data=d)

#### compute individual specific risks
fit.pred <- predict(fit, newdata=d[1:3], times=c(3,8), cause = 1,
                  se = TRUE, iid = TRUE, band = TRUE)
fit.pred

## check confidence intervals

```

```

newse <- fit.pred$absRisk.se/(-fit.pred$absRisk*log(fit.pred$absRisk))
cbind(lower = as.double(exp(-exp(log(-log(fit.pred$absRisk)) + 1.96 * newse))),
      upper = as.double(exp(-exp(log(-log(fit.pred$absRisk)) - 1.96 * newse)))
)

#### compute confidence intervals without transformation
confint(fit.pred, absRisk.transform = "none")
cbind(lower = as.double(fit.pred$absRisk - 1.96 * fit.pred$absRisk.se),
      upper = as.double(fit.pred$absRisk + 1.96 * fit.pred$absRisk.se)
)

```

confint.wglm

Confidence intervals for Estimate from IPCW Logistic Regressions

Description

Display the confidence intervals w.r.t. the estimated regression parameters from IPCW logistic regressions.

Usage

```

## S3 method for class 'wglm'
confint(object, parm = NULL, level = 0.95, times = NULL, type = "robust", ...)

```

Arguments

object	a wglm object.
parm	not used. For compatibility with the generic method.
level	[numeric, 0-1] Level of confidence.
times	[numeric vector] time points at which the estimates should be output.
type	[character] should the robust variance-covariance matrix be computing accounting for the uncertainty of the IPCW ("robust") or ignoring the uncertainty of the IPCW ("robust-wknown"), or model-based ignoring the uncertainty of the IPCW ("model-wknown")?
...	Not used.

coxBaseEstimator	<i>Extract the type of estimator for the baseline hazard</i>
------------------	--

Description

Extract the type of estimator for the baseline hazard

Usage

```
coxBaseEstimator(object)

## S3 method for class 'coxph'
coxBaseEstimator(object)

## S3 method for class 'phreg'
coxBaseEstimator(object)

## S3 method for class 'prodlm'
coxBaseEstimator(object)
```

Arguments

object	The fitted Cox regression model object either obtained with coxph (survival package), cph (rms package), or phreg (mets package).
--------	---

Author(s)

Brice Ozenne broz@sund.ku.dk

coxCenter	<i>Extract the mean value of the covariates</i>
-----------	---

Description

Extract the mean value of the covariates

Usage

```
coxCenter(object)

## S3 method for class 'cph'
coxCenter(object)

## S3 method for class 'coxph'
coxCenter(object)

## S3 method for class 'phreg'
coxCenter(object)
```

Arguments

object The fitted Cox regression model object either obtained with coxph (survival package), cph (rms package), or phreg (mets package).

Author(s)

Brice Ozenne broz@sund.ku.dk

coxFormula

Extract the formula from a Cox model

Description

Extract the formula from a Cox model

Usage

```
coxFormula(object)

## S3 method for class 'cph'
coxFormula(object)

## S3 method for class 'coxph'
coxFormula(object)

## S3 method for class 'phreg'
coxFormula(object)

## S3 method for class 'glm'
coxFormula(object)

## S3 method for class 'prodlm'
coxFormula(object)
```

Arguments

object The fitted Cox regression model object either obtained with coxph (survival package), cph (rms package), or phreg (mets package).

Author(s)

Brice Ozenne broz@sund.ku.dk

coxLP	<i>Compute the linear predictor of a Cox model</i>
-------	--

Description

Compute the linear predictor of a Cox model

Usage

```
coxLP(object, data, center)

## S3 method for class 'cph'
coxLP(object, data, center)

## S3 method for class 'coxph'
coxLP(object, data, center)

## S3 method for class 'phreg'
coxLP(object, data, center)

## S3 method for class 'prodlm'
coxLP(object, data, center)
```

Arguments

object	The fitted Cox regression model object either obtained with coxph (survival package), cph (rms package), or phreg (mets package).
data	a data.frame or a data.table
center	should the linear predictor be computed after centering the covariates

Details

In case of empty linear predictor returns a vector of 0 with the same length as the number of rows of the dataset

Author(s)

Brice Ozenne broz@sund.ku.dk

coxModelFrame	<i>Extract the design matrix used to train a Cox model</i>
---------------	--

Description

Extract the design matrix used to train a Cox model. Should contain the time of event, the type of event, the variable for the linear predictor, the strata variables and the date of entry (in case of delayed entry).

Usage

```
coxModelFrame(object, center)

## S3 method for class 'coxph'
coxModelFrame(object, center = FALSE)

## S3 method for class 'cph'
coxModelFrame(object, center = FALSE)

## S3 method for class 'phreg'
coxModelFrame(object, center = FALSE)

## S3 method for class 'prodlm'
coxModelFrame(object, center = FALSE)
```

Arguments

object	The fitted Cox regression model object either obtained with coxph (survival package), cph (rms package), or phreg (mets package).
center	[logical] Should the variables of the linear predictor be added ?

Author(s)

Brice Ozenne broz@sund.ku.dk

coxN	<i>Extract the number of observations from a Cox model</i>
------	--

Description

Extract the number of observations from a Cox model

Usage

```
coxN(object)

## S3 method for class 'cph'
coxN(object)

## S3 method for class 'coxph'
coxN(object)

## Default S3 method:
coxN(object)

## S3 method for class 'phreg'
coxN(object)

## S3 method for class 'CauseSpecificCox'
coxN(object)

## S3 method for class 'glm'
coxN(object)

## S3 method for class 'prodlm'
coxN(object)
```

Arguments

object	The fitted Cox regression model object either obtained with coxph (survival package), cph (rms package), or phreg (mets package).
--------	---

Author(s)

Brice Ozenne broz@sund.ku.dk

coxSpecial

Special characters in Cox model

Description

Return the special character(s) of the Cox model, e.g. used to indicate the strata variables.

Usage

```
coxSpecial(object)

## S3 method for class 'coxph'
coxSpecial(object)
```

```
## S3 method for class 'cph'
coxSpecial(object)

## S3 method for class 'phreg'
coxSpecial(object)

## S3 method for class 'prodlm'
coxSpecial(object)
```

Arguments

object The fitted Cox regression model object either obtained with `coxph` (survival package), `cph` (rms package), or `phreg` (mets package).

Details

Must return a list with at least one element `strata` indicating the character in the formula marking the variable(s) defining the strata.

Author(s)

Brice Ozenne broz@sund.ku.dk

coxStrata	<i>Define the strata for a new dataset</i>
-----------	--

Description

Define the strata in a dataset to match those of a stratified Cox model

Usage

```
coxStrata(object, data, sterms, strata.vars, strata.levels)

## S3 method for class 'cph'
coxStrata(object, data, sterms, strata.vars, strata.levels)

## S3 method for class 'coxph'
coxStrata(object, data, sterms, strata.vars, strata.levels)

## S3 method for class 'phreg'
coxStrata(object, data, sterms, strata.vars, strata.levels)

## S3 method for class 'prodlm'
coxStrata(object, data, sterms, strata.vars, strata.levels)
```

Arguments

object	The fitted Cox regression model object either obtained with coxph (survival package), cph (rms package), or phreg (mets package).
data	a data.frame or a data.table
sterms	terms in the formula corresponding to the strata variables
strata.vars	the name of the variables used to define the strata
strata.levels	a named list containing for each variable used to form the strata all its possible levels
levels	the strata levels that have been used to fit the Cox model

Details

if no strata variables returns a vector of "1" (factor).

Author(s)

Brice Ozenne broz@sund.ku.dk

coxStrataLevel	<i>Returns the name of the strata in Cox model</i>
----------------	--

Description

Return the name of the strata in Cox model

Usage

```
coxStrataLevel(object)

## S3 method for class 'coxph'
coxStrataLevel(object)

## S3 method for class 'cph'
coxStrataLevel(object)

## S3 method for class 'phreg'
coxStrataLevel(object)

## S3 method for class 'prodlm'
coxStrataLevel(object)
```

Arguments

object	The fitted Cox regression model object either obtained with coxph (survival package), cph (rms package), or phreg (mets package).
--------	---

Author(s)

Brice Ozenne broz@sund.ku.dk

coxVarCov

Extract the variance covariance matrix of the beta from a Cox model

Description

Extract the variance covariance matrix of the beta from a Cox model

Usage

```
coxVarCov(object)

## S3 method for class 'cph'
coxVarCov(object)

## S3 method for class 'coxph'
coxVarCov(object)

## S3 method for class 'phreg'
coxVarCov(object)

## S3 method for class 'prodlm'
coxVarCov(object)
```

Arguments

object The fitted Cox regression model object either obtained with coxph (survival package), cph (rms package), or phreg (mets package).

Details

Should return NULL if the Cox model has no covariate. The rows and columns of the variance covariance matrix must be named with the names used in the design matrix.

Author(s)

Brice Ozenne broz@sund.ku.dk

coxVariableName	<i>Extract variable names from a model</i>
-----------------	--

Description

Extract the name of the variables belonging to the linear predictor or used to form the strata

Usage

```
coxVariableName(object, model.frame)
```

Arguments

object	The fitted Cox regression model object either obtained with <code>coxph</code> (survival package) or <code>cph</code> (rms package).
model.frame	[data.frame] dataset containing all the relevant variables (entry, time to event, type of event, variables in the linear predictor, strata). Output from <code>coxModelFrame</code> .

Author(s)

Brice Ozenne broz@sund.ku.dk

CSC	<i>Cause-specific Cox proportional hazard regression</i>
-----	--

Description

Interface for fitting cause-specific Cox proportional hazard regression models in competing risk.

Usage

```
CSC(formula, data, cause, surv.type = "hazard", fitter = "coxph", ...)
```

Arguments

formula	Either a single Hist formula or a list of formulas. If it is a list it must contain as many Hist formulas as there are causes when <code>surv.type="hazard"</code> and exactly two formulas when <code>surv.type="survival"</code> . If it is a list the first formula is used for the cause of interest specific Cox regression and the other formula(s) either for the other cause specific Cox regression(s) or for the Cox regression of the combined event where each cause counts as event. Note that when only one formula is given the covariates enter in exactly the same way into all Cox regression analyses.
data	A data in which to fit the models.
cause	The cause of interest. Defaults to the first cause (see Details).

<code>surv.type</code>	Either "hazard" (the default) or "survival". If "hazard" fit cause-specific Cox regression models for all causes. If "survival" fit one cause-specific Cox regression model for the cause of interest and also a Cox regression model for event-free survival.
<code>fitter</code>	Routine to fit the Cox regression models. If <code>coxph</code> use <code>survival::coxph</code> else use <code>rms::cph</code> .
<code>...</code>	Arguments given to <code>fitter</code> , e.g., <code>coxph</code> .

Details

The causes and their order are determined by `prodlim::getStates()` applied to the `Hist` object.

Value

<code>models</code>	a list with the fitted (cause-specific) Cox regression objects
<code>response</code>	the event history response
<code>eventTimes</code>	the sorted (unique) event times
<code>surv.type</code>	the value of <code>surv.type</code>
<code>theCause</code>	the cause of interest. see <code>cause</code>
<code>causes</code>	the other causes

Author(s)

Thomas A. Gerds <tag@biostat.ku.dk> and Ulla B. Mogensen

References

B. Ozenne, A. L. Soerensen, T.H. Scheike, C.T. Torp-Pedersen, and T.A. Gerds. `riskregression`: Predicting the risk of an event using Cox regression models. *R Journal*, 9(2):440–460, 2017.

J Benichou and Mitchell H Gail. Estimates of absolute cause-specific risk in cohort studies. *Biometrics*, pages 813–826, 1990.

T.A. Gerds, T.H. Scheike, and P.K. Andersen. Absolute risk regression for competing risks: Interpretation, link functions, and prediction. *Statistics in Medicine*, 31(29):3921–3930, 2012.

See Also

[coxph](#)

Examples

```
library(prodlim)
library(survival)
data(Melanoma)
## fit two cause-specific Cox models
## different formula for the two causes
fit1 <- CSC(list(Hist(time,status)~sex+age,Hist(time,status)~invasion+epicel+log(thick)),
             data=Melanoma)
print(fit1)
```

```

## Not run:
library(Publish)
publish(fit1)

## End(Not run)

## model hazard of all cause mortality instead of hazard of type 2
fit1a <- CSC(list(Hist(time,status)~sex+age,Hist(time,status)~invasion+epicel+log(thick)),
              data=Melanoma,
              surv.type="surv")

## the predicted probabilities are similar
plot(predictRisk(fit1,times=500,cause=1,newdata=Melanoma),
      predictRisk(fit1a,times=500,cause=1,newdata=Melanoma))

## special case where cause 2 has no covariates
fit1b <- CSC(list(Hist(time,status)~sex+age,Hist(time,status)~1),
              data=Melanoma)
print(fit1b)
predict(fit1b,cause=1,times=100,newdata=Melanoma)

## same formula for both causes
fit2 <- CSC(Hist(time,status)~invasion+epicel+age,
            data=Melanoma)
print(fit2)

## combine a cause-specific Cox regression model for cause 2
## and a Cox regression model for the event-free survival:
## different formula for cause 2 and event-free survival
fit3 <- CSC(list(Hist(time,status)~sex+invasion+epicel+age,
                Hist(time,status)~invasion+epicel+age),
            surv.type="surv",
            data=Melanoma)
print(fit3)

## same formula for both causes
fit4 <- CSC(Hist(time,status)~invasion+epicel+age,
            data=Melanoma,
            surv.type="surv")
print(fit4)

## strata
fit5 <- CSC(Hist(time,status)~invasion+epicel+age+strata(sex),
            data=Melanoma,
            surv.type="surv")
print(fit5)

## sanity checks

cox1 <- coxph(Surv(time,status==1)~invasion+epicel+age+strata(sex),data=Melanoma)
cox2 <- coxph(Surv(time,status!=0)~invasion+epicel+age+strata(sex),data=Melanoma)
all.equal(coef(cox1),coef(fit5$models[[1]]))

```

```

all.equal(coef(cox2),coef(fit5$models[[2]]))

## predictions
##
## surv.type = "hazard": predictions for both causes can be extracted
## from the same fit
fit2 <- CSC(Hist(time,status)~invasion+epicel+age, data=Melanoma)
predict(fit2,cause=1,newdata=Melanoma[c(17,99,108),],times=c(100,1000,10000))
predictRisk(fit2,cause=1,newdata=Melanoma[c(17,99,108),],times=c(100,1000,10000))
predictRisk(fit2,cause=2,newdata=Melanoma[c(17,99,108),],times=c(100,1000,10000))
predict(fit2,cause=1,newdata=Melanoma[c(17,99,108),],times=c(100,1000,10000))
predict(fit2,cause=2,newdata=Melanoma[c(17,99,108),],times=c(100,1000,10000))

## surv.type = "surv" we need to change the cause of interest
library(survival)
fit5.2 <- CSC(Hist(time,status)~invasion+epicel+age+strata(sex),
             data=Melanoma,
             surv.type="surv",cause=2)
## now this does not work because the object was fitted with surv.type='surv'
try(predictRisk(fit5.2,cause=1,newdata=Melanoma,times=4000))

## but this does
predictRisk(fit5.2,cause=2,newdata=Melanoma,times=100)
predict(fit5.2,cause=2,newdata=Melanoma,times=100)
predict(fit5.2,cause=2,newdata=Melanoma[4,],times=100)

fit5.2 <- CSC(Hist(time,status)~invasion+epicel+age+strata(sex),
             data=Melanoma,
             surv.type="hazard",cause=2)

```

Ctree

S3-Wrapper for ctree.

Description

The call is added to an ctree object

Usage

```
Ctree(...)
```

Arguments

... passed to ctree

Value

list with two elements: ctree and call

Author(s)

Thomas A. Gerds <tag@biostat.ku.dk>

See Also

Cforest

Examples

```
if (require("party",quietly=TRUE)){
  library(proclim)
  library(party)
  library(survival)
  set.seed(50)
  d <- SimSurv(50)
  nd <- data.frame(X1=c(0,1,0),X2=c(-1,0,1))
  f <- Ctree(Surv(time,status)~X1+X2,data=d)
  predictRisk(f,newdata=nd,times=c(3,8))
}
```

discreteRoot

Dichotomic search for monotone function

Description

Find the root of a monotone function on a discrete grid of value using dichotomic search

Usage

```
discreteRoot(
  fn,
  grid,
  increasing = TRUE,
  check = TRUE,
  tol = .Machine$double.eps^0.5
)
```

Arguments

fn	[function] objective function to minimize in absolute value.
grid	[vector] possible minimizers.
increasing	[logical] is the function fn increasing?
check	[logical] should the program check that fn takes a different sign for the first vs. the last value of the grid?
tol	[numeric] the absolute convergence tolerance.

FGR*Formula wrapper for crr from cmprsk*

Description

Formula interface for Fine-Gray regression competing risk models.

Usage

```
FGR(formula, data, cause = 1, y = TRUE, ...)
```

Arguments

formula	A formula whose left hand side is a <code>Hist</code> object – see Hist . The right hand side specifies (a linear combination of) the covariates. See examples below.
data	A <code>data.frame</code> in which all the variables of <code>formula</code> can be interpreted.
cause	The failure type of interest. Defaults to 1.
y	logical value: if <code>TRUE</code> , the response vector is returned in component response.
...	...

Details

Formula interface for the function `crr` from the `cmprsk` package.

The function `crr` allows to multiply some covariates by time before they enter the linear predictor. This can be achieved with the formula interface, however, the code becomes a little cumbersome. See the examples. Note that `FGR` does not allow for delayed entry (left-truncation). The assumed value for indicating censored observations in the event variable is `0`. The function `Hist` has an argument `cens.code` which can change this (if you do not want to change the event variable).

Value

See `crr`.

Author(s)

Thomas Alexander Gerds <tag@biostat.ku.dk>

References

Gerds, TA and Scheike, T and Andersen, PK (2011) Absolute risk regression for competing risks: interpretation, link functions and prediction Research report 11/7. Department of Biostatistics, University of Copenhagen

See Also

[riskRegression](#)

Examples

```

library(prodlim)
library(survival)
library(cmprsk)
library(lava)
d <- prodlim::SimCompRisk(100)
f1 <- FGR(Hist(time,cause)~X1+X2,data=d)
print(f1)

## crr allows that some covariates are multiplied by
## a function of time (see argument tf of crr)
## by FGR uses the identity matrix
f2 <- FGR(Hist(time,cause)~cov2(X1)+X2,data=d)
print(f2)

## same thing, but more explicit:
f3 <- FGR(Hist(time,cause)~cov2(X1)+cov1(X2),data=d)
print(f3)

## both variables can enter cov2:
f4 <- FGR(Hist(time,cause)~cov2(X1)+cov2(X2),data=d)
print(f4)

## change the function of time
qFun <- function(x){x^2}
noFun <- function(x){x}
sqFun <- function(x){x^0.5}

## multiply X1 by time^2 and X2 by time:
f5 <- FGR(Hist(time,cause)~cov2(X1,tf=qFun)+cov2(X2),data=d)
print(f5)
print(f5$crrFit)
## same results as crr
with(d,crr(ftime=time,
           fstatus=cause,
           cov2=d[,c("X1","X2")],
           tf=function(time){cbind(qFun(time),time)}))

## still same result, but more explicit
f5a <- FGR(Hist(time,cause)~cov2(X1,tf=qFun)+cov2(X2,tf=noFun),data=d)
f5a$crrFit

## multiply X1 by time^2 and X2 by sqrt(time)
f5b <- FGR(Hist(time,cause)~cov2(X1,tf=qFun)+cov2(X2,tf=sqFun),data=d,cause=1)

## additional arguments for crr
f6<- FGR(Hist(time,cause)~X1+X2,data=d, cause=1,gtol=1e-5)
f6
f6a<- FGR(Hist(time,cause)~X1+X2,data=d, cause=1,gtol=0.1)
f6a

```

getSplitMethod *Input for data splitting algorithms*

Description

Parse hyperparameters for data splitting algorithm

Usage

```
getSplitMethod(split.method, B, N, M, seed)
```

Arguments

split.method	A character string specifying the algorithm for data splitting: <ul style="list-style-type: none">• "loob" leave one out bootstrap• "bootcv" bootstrap cross validation• "cv5" 5-fold cross validation• "loocv" leave one out cross validation aka N-1 fold cross validation• "632plus" Efron's .632+ bootstrap
B	Number of repetitions of bootstrap or k-fold cross-validation
N	Sample size
M	Subsample size. Default is N (no subsampling).
seed	Integer passed to set.seed. If not given or NA no seed is set.

Value

A list with the following elements:

- split.methodName: the print name of the algorithm
- split.method: the internal name of the algorithm
- index: the index for data splitting. For bootstrap splitting this is a matrix with B columns and M rows identifying the in-bag subjects. For k-fold cross-validation this is a matrix with B columns identifying the membership to the k groups.
- k: the k of k-fold cross-validation
- N: the sample size
- M: the subsample size

Author(s)

Thomas A. Gerds <tag@biostat.ku.dk>

See Also

Score

Examples

```
# 3-fold crossvalidation
getSplitMethod("cv3",B=4,N=37)

# bootstrap with replacement
getSplitMethod("loob",B=4,N=37)

# bootstrap without replacement
getSplitMethod("loob",B=4,N=37,M=20)
```

GLMnet

*Fitting GLMnet for use with predictRisk***Description**

Fit GLMnet models via a formula and a data set for use with [predictRisk](#).

Usage

```
GLMnet(
  formula,
  data,
  lambda = NULL,
  cv = TRUE,
  alpha = 1,
  nfolds = 10,
  type.measure = "deviance",
  family,
  ...
)
```

Arguments

formula	A formula.
data	The data on which to fit the model.
lambda	The tuning parameters for GLMnet. If set to NULL, then it the parameters are chosen for you.
cv	Whether to use cross-validation or not. Default is TRUE.
alpha	The elasticnet mixing parameter. See the <code>?glmnet</code> for more details.
nfolds	Number of folds for cross-validation. Default is 10.
type.measure	loss to use for cross-validation. Default is deviance.
family	passed to <code>glmnet</code> . Defaults for binary outcome to "binomial" and for survival to "cox".
...	Additional arguments that are passed on to the <code>glmnet</code> .

Hal9001

*Fitting HAL for use with predictRisk***Description**

Fit HAL models via a formula and a data set for use with `predictRisk`.

Usage

```
Hal9001(formula, data, lambda = NULL, ...)
```

Arguments

formula	A formula.
data	The data on which to fit the model.
lambda	The tuning parameters for HAL. If set to NULL, then it the parameters are chosen for you.
...	Additional arguments that are passed on to the <code>hal_fit</code> .

iid.wglm

*IID for IPCW Logistic Regressions***Description**

Compute the decomposition in iid elements of the ML estimator of IPCW logistic regressions.

Usage

```
## S3 method for class 'wglm'
iid(x, times = NULL, simplify = TRUE, store = NULL, ...)
```

Arguments

x	a wglm object.
times	[numeric vector] time points at which the iid should be output.
simplify	[logical] should the output be converted to a matrix when only one timepoint is requested. Otherwise will always return a list.
store	[vector] when evaluating the iid, should prediction be only computed for unique covariate sets and mapped back to the original dataset (<code>data="minimal"</code>). Otherwise use <code>data="full"</code> .
...	Not used.

`iidCox`*Extract iid decomposition from a Cox model*

Description

Compute the influence function for each observation used to estimate the model

Usage

```
iidCox(  
  object,  
  newdata,  
  baseline.iid,  
  tau.hazard,  
  tau.max,  
  store.iid,  
  keep.times,  
  return.object  
)  
  
## S3 method for class 'coxph'  
iidCox(  
  object,  
  newdata = NULL,  
  baseline.iid = TRUE,  
  tau.hazard = NULL,  
  tau.max = NULL,  
  store.iid = "full",  
  keep.times = TRUE,  
  return.object = TRUE  
)  
  
## S3 method for class 'cph'  
iidCox(  
  object,  
  newdata = NULL,  
  baseline.iid = TRUE,  
  tau.hazard = NULL,  
  tau.max = NULL,  
  store.iid = "full",  
  keep.times = TRUE,  
  return.object = TRUE  
)  
  
## S3 method for class 'phreg'  
iidCox(  
  object,
```

```

    newdata = NULL,
    baseline.iid = TRUE,
    tau.hazard = NULL,
    tau.max = NULL,
    store.iid = "full",
    keep.times = TRUE,
    return.object = TRUE
)

## S3 method for class 'prodlim'
iidCox(
  object,
  newdata = NULL,
  baseline.iid = TRUE,
  tau.hazard = NULL,
  tau.max = NULL,
  store.iid = "full",
  keep.times = TRUE,
  return.object = TRUE
)

## S3 method for class 'CauseSpecificCox'
iidCox(
  object,
  newdata = NULL,
  baseline.iid = TRUE,
  tau.hazard = NULL,
  tau.max = NULL,
  store.iid = "full",
  keep.times = TRUE,
  return.object = TRUE
)

```

Arguments

<code>object</code>	object The fitted Cox regression model object either obtained with <code>coxph</code> (survival package) or <code>cph</code> (rms package).
<code>newdata</code>	[data.frame] Optional new data at which to do iid decomposition
<code>baseline.iid</code>	[logical] Should the influence function for the baseline hazard be computed.
<code>tau.hazard</code>	[numeric vector] the vector of times at which the i.i.d decomposition of the baseline hazard will be computed
<code>tau.max</code>	[numeric] latest time at which the i.i.d decomposition of the baseline hazard will be computed. Alternative to <code>tau.hazard</code> .
<code>store.iid</code>	[character] the method used to compute the influence function and the standard error. Can be "full" or "minimal". See the details section.
<code>keep.times</code>	[logical] If TRUE add the evaluation times to the output.

`return.object` [logical] If TRUE return the object where the iid decomposition has been added. Otherwise return a list (see the return section)

Details

This function implements the first three formula (no number,10,11) of the subsection "Empirical estimates" in Ozenne et al. (2017). If there is no event in a strata, the influence function for the baseline hazard is set to 0.

Argument `store.iid`: If `n` denotes the sample size, `J` the number of jump times, and `p` the number of coefficients:

- `store.iid="full"` exports the influence function for the coefficients and the baseline hazard at each event time.
- `store.iid="minimal"` exports the influence function for the coefficients. For the baseline hazard it only computes the quantities necessary to compute the influence function in order to save memory.

More details can be found in appendix B of Ozenne et al. (2017).

Value

For Cox models, it returns the object with an additional iid slot (i.e. `object$iid`). It is a list containing:

- `IFbeta`: Influence function for the regression coefficient.
- `IFhazard`: Time differential of the influence function of the hazard.
- `IFcumhazard`: Influence function of the cumulative hazard.
- `calcIFhazard`: Elements used to compute the influence function at a given time.
- `time`: Times at which the influence function has been evaluated.
- `etime1.min`: Time of first event (i.e. jump) in each strata.
- `etime.max`: Last observation time (i.e. jump or censoring) in each strata.
- `indexObs`: Index of the observation in the original dataset.

For Cause-Specific Cox models, it returns the object with an additional iid slot for each model (e.g. `object$models[[1]]iid`).

References

Brice Ozenne, Anne Lyngholm Sorensen, Thomas Scheike, Christian Torp-Pedersen and Thomas Alexander Gerds. `riskRegression`: Predicting the Risk of an Event using Cox Regression Models. *The R Journal* (2017) 9:2, pages 440-460.

Examples

```
library(survival)
library(data.table)
library(prodlim)
set.seed(10)
```

```
d <- sampleData(100, outcome = "survival")[,.(eventtime,event,X1,X6)]
setkey(d, eventtime)

m.cox <- coxph(Surv(eventtime, event) ~ X1+X6, data = d, y = TRUE, x = TRUE)
system.time(IF.cox <- iidCox(m.cox))

IF.cox.all <- iidCox(m.cox, tau.hazard = sort(unique(c(7,d$eventtime))))
IF.cox.beta <- iidCox(m.cox, baseline.iid = FALSE)
```

influenceTest	<i>Influence test [Experimental!!]</i>
---------------	--

Description

Compare two estimates using their influence function

Usage

```
influenceTest(object, ...)

## S3 method for class 'list'
influenceTest(
  object,
  newdata,
  times,
  type,
  cause,
  keep.newdata = TRUE,
  keep.strata = FALSE,
  ...
)

## Default S3 method:
influenceTest(object, object2, band = TRUE, ...)
```

Arguments

object	either a list of models or an object of class predictCox or predictCSC.
...	additional arguments to be passed to lower level functions.
newdata	[data.frame or data.table] Contain the values of the predictor variables defining subject specific predictions.
times	[numeric vector] Time points at which to return the estimated absolute risk.
type	[character]the type of predicted value.
cause	[integer/character] Identifies the cause of interest among the competing events.

keep.newdata	[logical] If TRUE add the value of the covariates used to make the prediction in the output.
keep.strata	[logical] If TRUE add the value of the strata used to make the prediction in the output.
object2	same as predict1 but for another model.
band	[logical] If TRUE add the influence function to the output such that confint will be able to compute the confidence bands.

Examples

```

library(lava)
library(survival)
library(prodlm)
library(data.table)
n <- 100

#### Under H1
set.seed(1)
newdata <- data.frame(X1=0:1)

## simulate non proportional hazard using lava
m <- lvm()
regression(m) <- y ~ 1
regression(m) <- s ~ exp(-2*X1)
distribution(m,~X1) <- binomial.lvm()
distribution(m,~cens) <- coxWeibull.lvm(scale=1)
distribution(m,~y) <- coxWeibull.lvm(scale=1,shape=~s)
eventTime(m) <- eventtime ~ min(y=1,cens=0)
d <- as.data.table(sim(m,n))
setkey(d, eventtime)

## fit cox models
m.cox <- coxph(Surv(eventtime, status) ~ X1,
               data = d, y = TRUE, x = TRUE)

mStrata.cox <- coxph(Surv(eventtime, status) ~ strata(X1),
                   data = d, y = TRUE, x = TRUE)

## compare models
# one time point
outIF <- influenceTest(list(m.cox, mStrata.cox),
                       type = "survival", newdata = newdata, times = 0.5)
confint(outIF)

# several timepoints
outIF <- influenceTest(list(m.cox, mStrata.cox),
                       type = "survival", newdata = newdata, times = c(0.5,1,1.5))
confint(outIF)

#### Under H0 (Cox) ####
set.seed(1)

```

```

## simulate proportional hazard using lava
m <- lvm()
regression(m) <- y ~ 1
distribution(m,~X1) <- binomial.lvm()
distribution(m,~cens) <- coxWeibull.lvm()
distribution(m,~y) <- coxWeibull.lvm()
eventTime(m) <- eventtime ~ min(y=1,cens=0)
d <- as.data.table(sim(m,n))
setkey(d, eventtime)

## fit cox models
Utime <- sort(unique(d$eventtime))
m.cox <- coxph(Surv(eventtime, status) ~ X1,
              data = d, y = TRUE, x = TRUE)

mStrata.cox <- coxph(Surv(eventtime, status) ~ strata(X1),
                   data = d, y = TRUE, x = TRUE)

p.cox <- predictCox(m.cox, newdata = newdata, time = Utime, type = "survival")
p.coxStrata <- predictCox(mStrata.cox, newdata = newdata, time = Utime, type = "survival")

## display
library(ggplot2)
autoplot(p.cox)
autoplot(p.coxStrata)

## compare models
outIF <- influenceTest(list(m.cox, mStrata.cox),
                      type = "survival", newdata = newdata, times = Utime[1:6])
confint(outIF)

#### Under H0 (CSC) ####
set.seed(1)
ff <- ~ f(X1,2) + f(X2,-0.033)
ff <- update(ff, ~ .+ f(X3,0) + f(X4,0) + f(X5,0))
ff <- update(ff, ~ .+ f(X6,0) + f(X7,0) + f(X8,0) + f(X9,0))
d <- sampleData(n, outcome = "competing.risk", formula = ff)
d[,X1:=as.numeric(as.character(X1))]
d[,X2:=as.numeric(as.character(X2))]
d[,X3:=as.numeric(as.character(X3))]
d[,X4:=as.numeric(as.character(X4))]
d[,X5:=as.numeric(as.character(X5))]
setkey(d, time)

Utime <- sort(unique(d$time))

## fit cox models
m.CSC <- CSC(Hist(time, event) ~ X1 + X2, data = d)
mStrata.CSC <- CSC(Hist(time, event) ~ strata(X1) + X2 + X3, data = d)

## compare models
outIF <- influenceTest(list(m.CSC, mStrata.CSC),
                      cause = 1, newdata = unique(d[,.(X1,X2,X3)]), times = Utime[1:5])

```



```
confint(outIF)
```

```
information.wglm      Information for IPCW Logistic Regressions
```

Description

Compute the information (i.e. opposite of the expectation of the second derivative of the log-likelihood) for IPCW logistic regressions.

Usage

```
## S3 method for class 'wglm'
information(x, times = NULL, simplify = TRUE, ...)
```

Arguments

x	a wglm object.
times	[numeric vector] time points at which the score should be output.
simplify	[logical] should the output be converted to a matrix when only one timepoint is requested. Otherwise will always return a list.
...	Not used.

IPA	<i>Explained variation for settings with binary, survival and competing risk outcome</i>
-----	--

Description

Index of Prediction Accuracy: General R^2 for binary outcome and right censored time to event (survival) outcome also with competing risks

Usage

```
rsquared(object,...)
IPA(object,...)
## Default S3 method:
rsquared(object,formula,newdata,times,cause,...)
## S3 method for class 'glm'
rsquared(object,formula,newdata,...)
## S3 method for class 'coxph'
rsquared(object,formula,newdata,times,...)
## S3 method for class 'CauseSpecificCox'
rsquared(object,formula,newdata,times,cause,...)
## Default S3 method:
```

```

IPA(object,formula,newdata,times,cause,...)
## S3 method for class 'glm'
IPA(object,formula,newdata,...)
## S3 method for class 'coxph'
IPA(object,formula,newdata,times,...)
## S3 method for class 'CauseSpecificCox'
IPA(object,formula,newdata,times,cause,...)

```

Arguments

object	Model for which we want IPA.
...	passed to <code>riskRegression::Score</code>
newdata	Optional validation data set in which to compute IPA
formula	Formula passed to <code>Score</code> . If not provided, try to use the formula of the call of object, if any.
cause	For competing risk models the event of interest
times	Vector of time points used as prediction horizon for the computation of Brier scores.

Details

IPA (R^2) is calculated based on the model's predicted risks. The Brier score of the model is compared to the Brier score of the null model.

Value

Data frame with explained variation values for the full model.

Author(s)

Thomas A. Gerds <tag@biostat.ku.dk>

See Also

`Score`

Examples

```

library(prodlm)
library(data.table)
# binary outcome
library(lava)
set.seed(18)
learndat <- sampleData(48,outcome="binary")
lr1 = glm(Y~X1+X2+X7+X9,data=learndat,family=binomial)
IPA(lr1)

## validation data
valdat=sampleData(94,outcome="binary")

```

```

IPA(lr1,newdata=valdat)

## predicted risks externally given
p1=predictRisk(lr1,newdata=valdat)
IPA(p1,formula=Y~1,valdat)

# survival
library(survival)
data(pbc)
pbc=na.omit(pbc)
pbctest=(1:NROW(pbc)) %in% sample(1:NROW(pbc),size=.632*NROW(pbc))
pbclearn=pbc[pbctest,]
cox1= coxph(Surv(time,status!=0)~age+sex+log(bili)+log(albumin)+log(protime),
            data=pbclearn,x=TRUE)

## same data
IPA(cox1,formula=Surv(time,status!=0)~1,times=1000)

## validation data
pbcval=pbc[!pbctest,]
IPA(cox1,formula=Surv(time,status!=0)~1,newdata=pbcval,times=1000)

## predicted risks externally given
p2=predictRisk(cox1,newdata=pbcval,times=1000)
IPA(cox1,formula=Surv(time,status!=0)~1,newdata=pbcval,times=1000)

# competing risks
data(Melanoma)
Melanomatest=(1:NROW(Melanoma)) %in% sample(1:NROW(Melanoma),size=.632*NROW(Melanoma))
Melanomalearn=Melanoma[Melanomatest,]
fit1 <- CSC(list(Hist(time,status)~sex,
                Hist(time,status)~invasion+epicel+age),
            data=Melanoma)
IPA(fit1,times=1000,cause=2)

## validation data
Melanomaval=Melanoma[!Melanomatest,]
IPA(fit1,formula=Hist(time,status)~1,newdata=Melanomaval,times=1000)

## predicted risks externally given
p3= predictRisk(fit1,cause=1,newdata=Melanomaval,times=1000)
IPA(p3,formula=Hist(time,status)~1,cause=1,newdata=Melanomaval,times=1000)

```

Description

This function is used internally to obtain inverse of the probability of censoring weights.

Usage

```
ipcw(
  formula,
  data,
  method,
  args,
  times,
  subject.times,
  lag = 1,
  what,
  keep = NULL
)
```

Arguments

formula	A survival formula like, $\text{Surv}(\text{time}, \text{status}) \sim 1$, where as usual $\text{status}=0$ means censored. The status variable is internally reversed for estimation of censoring rather than survival probabilities. Some of the available models (see argument <code>method</code>) will use predictors on the right hand side of the formula.
data	The data used for fitting the censoring model
method	Censoring model used for estimation of the (conditional) censoring distribution.
args	A list of arguments which is passed to <code>method</code>
times	For <code>what="IPCW.times"</code> a vector of times at which to compute the probabilities of not being censored.
subject.times	For <code>what="IPCW.subject.times"</code> a vector of individual times at which the probabilities of not being censored are computed.
lag	If equal to 1 then obtain $G(T_i X_i)$, if equal to 0 estimate the conditional censoring distribution at the <code>subject.times</code> , i.e. $(G(T_i X_i))$.
what	Decide about what to do: If equal to <code>"IPCW.times"</code> then weights are estimated at given times. If equal to <code>"IPCW.subject.times"</code> then weights are estimated at individual <code>subject.times</code> . If missing then produce both.
keep	Which elements to add to the output. Any subset of the vector <code>c("times", "fit", "call")</code> .

Details

Inverse of the probability of censoring weights (IPCW) usually refer to the probabilities of not being censored at certain time points. These probabilities are also the values of the conditional survival function of the censoring time given covariates. The function `ipcw` estimates the conditional survival function of the censoring times and derives the weights.

IMPORTANT: the data set should be ordered, `order(time, -status)` in order to get the values `IPCW.subject.times` in the right order for some choices of `method`.

Value

A list with elements depending on argument `keep`.

times	The times at which weights are estimated
IPCW.times	Estimated weights at times
IPCW.subject.times	Estimated weights at individual time values subject.times
fit	The fitted censoring model
method	The method for modelling the censoring distribution
call	The call

Author(s)

Thomas A. Gerds <tag@biostat.ku.dk>

Examples

```
library(prodlim)
library(rms)
dat=SimSurv(30)

dat <- dat[order(dat$time),]

# using the marginal Kaplan-Meier for the censoring times

WKM=ipcw(Hist(time,status)~X2,
  data=dat,
  method="marginal",
  times=sort(unique(dat$time)),
  subject.times=dat$time,keep=c("fit"))
plot(WKM$fit)
WKM$fit

# using the Cox model for the censoring times given X2
library(survival)
WCox=ipcw(Hist(time=time,event=status)~X2,
  data=dat,
  method="cox",
  times=sort(unique(dat$time)),
  subject.times=dat$time,keep=c("fit"))
WCox$fit

plot(WKM$fit)
lines(sort(unique(dat$time)),
  1-WCox$IPCW.times[1,],
  type="l",
  col=2,
  lty=3,
  lwd=3)
lines(sort(unique(dat$time)),
  1-WCox$IPCW.times[5,],
  type="l",
  col=3,
  lty=3,
```

```

      lwd=3)

# using the stratified Kaplan-Meier
# for the censoring times given X2

WKM2=ipcw(Hist(time,status)~X2,
  data=dat,
  method="nonpar",
  times=sort(unique(dat$time)),
  subject.times=dat$time,keep=c("fit"))
plot(WKM2$fit,add=FALSE)

```

 is.iidCox

Check Computation of the Influence Function in a Cox Model

Description

Check whether the influence function of the Cox model or cause specific Cox models has been stored in the object.

Usage

```
is.iidCox(object)
```

Arguments

object	fitted Cox regression model object either obtained with coxph (survival package), cph (rms package), CSC (riskRegression package).
--------	--

 Melanoma

Malignant melanoma data

Description

In the period 1962-77, 205 patients with malignant melanoma (cancer of the skin) had a radical operation performed at Odense University Hospital, Denmark. All patients were followed until the end of 1977 by which time 134 were still alive while 71 had died (of out whom 57 had died from cancer and 14 from other causes).

Format

A data frame with 205 observations on the following 12 variables.

time time in days from operation

status a numeric with values 0=censored 1=death.malignant.melanoma 2=death.other.causes

event a factor with levels censored death.malignant.melanoma death.other.causes

invasion a factor with levels level.0, level.1, level.2

ici inflammatory cell infiltration (IFI): 0, 1, 2 or 3

epicel a factor with levels not present present

ulcer a factor with levels not present present

thick tumour thickness (in 1/100 mm)

sex a factor with levels Female Male

age age at operation (years)

logthick tumour thickness on log-scale

Details

The object of the study was to assess the effect of risk factors on survival. Among such risk factors were the sex and age of the patients and the histological variables tumor thickness and ulceration (absent vs. present).

References

Regression with linear predictors (2010)

Andersen, P.K. and Skovgaard, L.T.

Springer Verlag

Examples

```
data(Melanoma)
```

model.matrix.cph

Extract design matrix for cph objects

Description

Extract design matrix for cph objects

Usage

```
## S3 method for class 'cph'  
model.matrix(object, data, ...)
```

Arguments

object	a cph object.
data	a dataset.
...	not used

model.matrix.phreg	<i>Extract design matrix for phreg objects</i>
--------------------	--

Description

Extract design matrix for phreg objects

Usage

```
## S3 method for class 'phreg'
model.matrix(object, data, ...)
```

Arguments

object	a phreg object.
data	a dataset.
...	not used

Details

mainly a copy paste of the begining of the phreg function.

model.tables.ate	<i>Statistical Inference for the Average Treatment Effect</i>
------------------	---

Description

Export estimated average treatment effects with their uncertainty (standard errors, confidence intervals and p-values).

Usage

```
## S3 method for class 'ate'
model.tables(
  x,
  contrasts = NULL,
  times = NULL,
  estimator = NULL,
  type = NULL,
  ...
)
```


Arguments

x	A ate object, i.e. output of the ate function.
contrasts	[character vector] levels of the treatment variable for which the estimates should be assessed or compared. Default is to consider all levels.
times	[numeric vector] The timepoints at which the estimates should be displayed. Default is to consider all timepoints.
estimator	[character] The type of estimator relative to which the estimates should be displayed.
type	[character] should the average risk per treatment be displayed ("meanRisk"), or the difference in average risk between any two pairs of treatments ("diffRisk"), or the ratio in average risk between any two pairs of treatments ("ratioRisk").
...	Additional arguments (meanRisk.transform, ..., method.band, ...) passed to <code>confint.ate</code> .

Value

a data.frame.

Author(s)

Brice Ozenne <broz@sund.ku.dk>

model.tables.wglm

Statistical Inference for Estimate from IPCW Logistic Regressions

Description

Export estimated regression parameters from IPCW logistic regressions with their uncertainty (standard errors, confidence intervals and p-values).

Usage

```
## S3 method for class 'wglm'
model.tables(x, times = NULL, type = "robust", level = 0.95, ...)
```

Arguments

x	a wglm object.
times	[numeric vector] time points at which the estimates should be output.
type	[character] should the robust variance-covariance matrix be computing accounting for the uncertainty of the IPCW ("robust") or ignoring the uncertainty of the IPCW ("robust-wknown"), or model-based ignoring the uncertainty of the IPCW ("model-wknown")?
level	[numeric, 0-1] Level of confidence.
...	Not used.

Paquid

Paquid sample

Description

PAQUID is a prospective cohort study initiated in 1988 in South Western France to explore functional and cerebral ageing. This sample includes $n=2561$ subjects. Data contains a time-to-event, a type of event and two cognitive scores measured at baseline.

Format

A data frame with 2561 observations on the following 4 variables.

`time` the time-to-event (in years).

`status` the type of event $0 =$ censored, $1 =$ dementia onset and $2 =$ death without dementia.

`DSST` score at the Digit Symbol Substitution Score Test. This test explores attention and psychomotor speed.

`MMSE` score at the Mini Mental State Examination. This test is often used as an index of global cognitive performance.

Source

The data have been first made publicly available via the package `timeROC`.

References

Dartigues, J., Gagnon, M., Barberger-Gateau, P., Letenneur, L., Commenges, D., Sauvel, C., Michel, P., and Salamon, R. (1992). The paquid epidemiological program on brain ageing. *Neuroepidemiology*, 11(1):14–18.

Blanche, P., Dartigues, J. F., & Jacqmin-Gadda, H. (2013). Estimating and comparing time-dependent areas under receiver operating characteristic curves for censored event times with competing risks. *Statistics in Medicine*, 32(30), 5381-5397.

Examples

```
data(Paquid)
```

penalizedS3

*S3-wrapper for S4 function penalized***Description**

S3-wrapper for S4 function penalized

Usage

```
penalizedS3(formula, data, type = "elastic.net", lambda1, lambda2, fold, ...)
```

Arguments

formula	Communicated outcome and explanatory variables. See examples.
data	Data set in which formula is to be interpreted
type	String specifying the type of penalization. Should match one of the following values: "ridge", "lasso", "elastic.net".
lambda1	Lasso penalty
lambda2	ridge penalty
fold	passed to penalized::profL1
...	Arguments passed to penalized

Examples

```
library(proclim)
## Not run:
## too slow
if (require("penalized",quietly=TRUE)){
  library(penalized)
  set.seed(8)
  d <- sampleData(200,outcome="binary")
  newd <- sampleData(80,outcome="binary")
  fitridge <- penalizedS3(Y~X1+X2+pen(7:8), data=d, type="ridge",
    standardize=TRUE, model="logistic",trace=FALSE)
  fitlasso <- penalizedS3(Y~X1+X2+pen(7:8), data=d, type="lasso",
    standardize=TRUE, model="logistic",trace=FALSE)
  # fitnet <- penalizedS3(Y~X1+X2+pen(7:8), data=d, type="elastic.net",
  # standardize=TRUE, model="logistic",trace=FALSE)
  predictRisk(fitridge,newdata=newd)
  predictRisk(fitlasso,newdata=newd)
  # predictRisk(fitnet,newdata=newd)
  Score(list(fitridge),data=newd,formula=Y~1)
  Score(list(fitridge),data=newd,formula=Y~1,split.method="bootcv",B=2)
  data(nki70) ## S4 fit
  fitS4 <- penalized(Surv(time, event), penalized = nki70[,8:77],
    unpenalized = ~ER+Age+Diam+N+Grade, data = nki70,
    lambda1 = 1)
```

```

fitS3 <- penalizedS3(Surv(time,event)~ER+Age+Diam+pen(8:77)+N+Grade,
                    data=nki70, lambda1=1)
## or
penS3 <- penalizedS3(Surv(time,event)~ER+pen(TSPYL5,Contig63649_RC)+pen(10:77)+N+Grade,
                    data=nki70, lambda1=1)
## also this works
penS3 <- penalizedS3(Surv(time,event)~ER+Age+pen(8:33)+Diam+pen(34:77)+N+Grade,
                    data=nki70, lambda1=1)
}
## End(Not run)

```

plot.riskRegression *Plotting predicted risk*

Description

Show predicted risk obtained by a risk prediction model as a function of time.

Usage

```

## S3 method for class 'riskRegression'
plot(x,
     cause,
     newdata,
     xlab,
     ylab,
     xlim,
     ylim,
     lwd,
     col,
     lty,
     axes=TRUE,
     percent=TRUE,
     legend=TRUE,
     add=FALSE,
     ...)

```

Arguments

x	Fitted object obtained with one of ARR, LRR, riskRegression.
cause	For CauseSpecificCox models the cause of interest.
newdata	A data frame containing predictor variable combinations for which to compute predicted risk.
xlab	See plot
ylab	See plot
xlim	See plot

ylim	See plot
lwd	A vector of line thicknesses for the regression coefficients.
col	A vector of colors for the regression coefficients.
lty	A vector of line types for the regression coefficients.
axes	Logical. If FALSE then do not draw axes.
percent	If true the y-axis is labeled in percent.
legend	If true draw a legend.
add	Logical. If TRUE then add lines to an existing plot.
...	Used for transclusion of smart arguments for plot, lines, axis and background. See function SmartControl from prodlim.

Author(s)

Thomas Alexander Gerds <tag@biostat.ku.dk>

Examples

```
library(survival)
library(prodlim)
data(Melanoma)
fit.arr <- ARR(Hist(time,status)~invasion+age+strata(sex),data=Melanoma,cause=1)
plot(fit.arr,xlim=c(500,3000))
```

plotAUC

Plot of time-dependent AUC curves

Description

Plot of time-dependent AUC curves

Usage

```
plotAUC(
  x,
  models,
  which = "score",
  xlim,
  ylim,
  xlab,
  ylab,
  col,
  lwd,
  lty = 1,
```

```

    cex = 1,
    pch = 1,
    type = "l",
    axes = 1L,
    percent = 1L,
    conf.int = 0L,
    legend = 1L,
    ...
)

```

Arguments

x	Object obtained with <code>Score.list</code>
models	Choice of models to plot
which	Character. Either "score" to show AUC or "contrasts" to show differences between AUC.
xlim	Limits for x-axis
ylim	Limits for y-axis
xlab	Label for x-axis
ylab	Label for y-axis
col	line color
lwd	line width
lty	line style
cex	point size
pch	point style
type	line type
axes	Logical. If TRUE draw axes.
percent	Logical. If TRUE scale y-axis in percent.
conf.int	Logical. If TRUE draw confidence shadows.
legend	Logical. If TRUE draw legend.
...	Used for additional control of the subroutines: plot,

Examples

```

set.seed(9)
library(survival)
library(prodlim)
set.seed(10)
d=sampleData(100,outcome="survival")
nd=sampleData(100,outcome="survival")
f1=coxph(Surv(time,event)~X1+X6+X8,data=d,x=TRUE,y=TRUE)
f2=coxph(Surv(time,event)~X2+X5+X9,data=d,x=TRUE,y=TRUE)
xx=Score(list("X1+X6+X8"=f1,"X2+X5+X9"=f2), formula=Surv(time,event)~1,
data=nd, metrics="auc", null.model=FALSE, times=seq(3:10))
aucgraph <- plotAUC(xx)

```

```

plotAUC(xx,conf.int=TRUE)
## difference between
plotAUC(xx,which="contrasts",conf.int=TRUE)

```

plotBrier

Plot Brier curve

Description

Plot Brier score curves

Usage

```

plotBrier(
  x,
  models,
  which = "score",
  xlim,
  ylim,
  xlab,
  ylab,
  col,
  lwd,
  lty = 1,
  cex = 1,
  pch = 1,
  type = "l",
  axes = 1L,
  percent = 1L,
  conf.int = 0L,
  legend = 1L,
  ...
)

```

Arguments

x	Object obtained with Score
models	Choice of models to plot
which	Character. Either "score" to show AUC or "contrasts" to show differences between AUC.
xlim	Limits for x-axis
ylim	Limits for y-axis
xlab	Label for x-axis
ylab	Label for y-axis

col	line color
lwd	line width
lty	line style
cex	point size
pch	point style
type	line type
axes	Logical. If TRUE draw axes.
percent	Logical. If TRUE scale y-axis in percent.
conf.int	Logical. If TRUE draw confidence shadows.
legend	Logical. If TRUE draw legend.
...	Used for additional control of the subroutines: plot, axis, lines, legend. See SmartControl .

Examples

```
# survival
library(survival)
library(proplim)
set.seed(7)
ds1=sampleData(40,outcome="survival")
ds2=sampleData(40,outcome="survival")
f1 <- coxph(Surv(time,event)~X1+X3+X5+X7+X9,data=ds1,x=TRUE)
f2 <- coxph(Surv(time,event)~X2+X4+X6+X8+X10,data=ds1,x=TRUE)
xscore <- Score(list(f1,f2),formula=Hist(time,event)~1,data=ds2,times=0:12,metrics="brier")
plotBrier(xscore)
```

plotCalibration	<i>Plot Calibration curve</i>
-----------------	-------------------------------

Description

Plot Calibration curves for risk prediction models

Usage

```
plotCalibration(
  x,
  models,
  times,
  method = "nne",
  cens.method = "local",
  round = 2,
  bandwidth = NULL,
  q = 10,
  bars = FALSE,
```



```

    hanging = FALSE,
    names = "quantiles",
    pseudo = FALSE,
    rug,
    show.frequencies = FALSE,
    plot = TRUE,
    add = FALSE,
    diag = !add,
    legend = !add,
    auc.in.legend,
    brier.in.legend,
    axes = !add,
    xlim = c(0, 1),
    ylim = c(0, 1),
    xlab = ifelse(bars, "Risk groups", "Predicted risk"),
    ylab,
    col,
    lwd,
    lty,
    pch,
    type,
    percent = TRUE,
    na.action = na.fail,
    cex = 1,
    ...
)

```

Arguments

x	Object obtained with function Score
models	Choice of models to plot
times	Time point specifying the prediction horizon.
method	The method for estimating the calibration curve(s): <ul style="list-style-type: none"> • "quantile": The observed proportion at predicted risk value 'p' is obtained in groups defined by quantiles of the predicted event probabilities of all subjects. The number of groups is controlled by argument q. • "nne": The observed proportion at predicted risk value 'p' is obtained based on the subjects whose predicted risk is inside a nearest neighborhood around the value 'p'. The larger the bandwidth the more subjects are included in the current neighborhood.
cens.method	For right censored data only. How observed proportions are calculated. Either "jackknife" or "local": <ul style="list-style-type: none"> • "jackknife": Compute a running mean of the jackknife pseudovalues across neighborhoods/groups of the predicted risks. Here we rely on the assumption that censoring is independent of the event time and the covariates, see References.

	<ul style="list-style-type: none"> • "local": Compute the Kaplan-Meier estimator in absence of competing risks and the Aalen-Johansen estimator in presence of competing risks locally like a running mean in neighborhoods of the predicted risks. The widths of the neighborhoods are defined according to method.
round	If TRUE predicted probabilities are rounded to two digits before smoothing. This may have a considerable effect on computing efficiency in large data sets.
bandwidth	The bandwidth for method="nne"
q	The number of quantiles for method="quantile" and bars=TRUE.
bars	If TRUE, use barplots to show calibration.
hanging	Barplots only. If TRUE, hang bars corresponding to observed frequencies (estimated actual risk) at the value of the corresponding prediction.
names	Barplots only. Names argument passed to names.arg of barplot.
pseudo	If TRUE show pseudo values (only for right censored data).
rug	If TRUE show rug plot at the predictions
show.frequencies	Barplots only. If TRUE, show frequencies above the bars.
plot	If FALSE, do not plot the results, just return a plottable object.
add	If TRUE the line(s) are added to an existing plot.
diag	If FALSE no diagonal line is drawn.
legend	Logical. If TRUE draw legend.
auc.in.legend	Logical. If TRUE add AUC to legend.
brier.in.legend	Logical. If TRUE add Brier score to legend.
axes	If FALSE no axes are drawn.
xlim	Limits of x-axis.
ylim	Limits of y-axis.
xlab	Label for y-axis.
ylab	Label for x-axis.
col	Vector with colors, one for each element of object. Passed to lines .
lwd	Vector with line widths, one for each element of object. Passed to lines .
lty	lwd Vector with line style, one for each element of object. Passed to lines .
pch	Passed to lines .
type	Passed to lines .
percent	If TRUE axes labels are multiplied by 100 and thus interpretable on a percent scale.
na.action	what to do with NA values. Passed to model.frame
cex	Default cex used for legend and labels.
...	Used to control the subroutines: plot, axis, lines, barplot, legend, addtable2plot, points (pseudo values), rug. See SmartControl .

Details

In uncensored data, the observed frequency of the outcome event is calculated locally at the predicted risk. In right censored data with and without competing risks, the actual risk is calculated using the Kaplan-Meier and the Aalen-Johansen method, respectively, locally at the predicted risk.

Examples

```
library(prodlm)
# binary
set.seed(10)
db=sampleData(100,outcome="binary")
fb1=glm(Y~X1+X5+X7,data=db,family="binomial")
fb2=glm(Y~X1+X3+X6+X7,data=db,family="binomial")
xb=Score(list(model1=fb1,model2=fb2),Y~1,data=db,
         plots="cal")
plotCalibration(xb,brier.in.legend=TRUE)
plotCalibration(xb,bars=TRUE,model="model1")
plotCalibration(xb,models=1,bars=TRUE,names.cex=1.3)

# survival
library(survival)
library(prodlm)
dslearn=sampleData(56,outcome="survival")
dstest=sampleData(100,outcome="survival")
fs1=coxph(Surv(time,event)~X1+X5+X7,data=dslearn,x=1)
fs2=coxph(Surv(time,event)~strata(X1)+X3+X6+X7,data=dslearn,x=1)
xs=Score(list(Cox1=fs1,Cox2=fs2),Surv(time,event)~1,data=dstest,
         plots="cal",metrics=NULL)
plotCalibration(xs)
plotCalibration(xs,cens.method="local",pseudo=1)
plotCalibration(xs,method="quantile")

# competing risks

## Not run:
data(Melanoma)
f1 <- CSC(Hist(time,status)~age+sex+epicel+ulcer,data=Melanoma)
f2 <- CSC(Hist(time,status)~age+sex+logthick+epicel+ulcer,data=Melanoma)
x <- Score(list(model1=f1,model2=f2),Hist(time,status)~1,data=Melanoma,
         cause= 2,times=5*365.25,plots="cal")
plotCalibration(x)

## End(Not run)
```

Description

Plot time-varying effects from a risk regression model.

Usage

```
plotEffects(
  x,
  formula,
  level,
  ref.line = TRUE,
  conf.int = 0.95,
  xlim,
  ylim,
  xlab = "Time",
  ylab = "Cumulative coefficient",
  col,
  lty,
  lwd,
  add = FALSE,
  legend,
  axes = TRUE,
  ...
)
```

Arguments

x	Fitted object obtained with one of ARR, LRR, riskRegression.
formula	A formula to specify the variable(s) whose regression coefficients should be plotted.
level	For categorical variables the level (group) whose contrast to the reference level (group) should be plotted.
ref.line	Logical. If TRUE then add a horizontal line at zero.
conf.int	Logical. If TRUE then add confidence limits. Can be controlled using smart arguments. See examples
xlim	See plot
ylim	See plot
xlab	See plot
ylab	See plot
col	A vector of colors for the regression coefficients.
lty	A vector of line types for the regression coefficients.
lwd	A vector of line thicknesses for the regression coefficients.
add	Logical. If TRUE then add lines to an existing plot.
legend	Logical. If TRUE then add a legend. Can be controlled using smart arguments. See examples.

axes Logical. If FALSE then do not draw axes.
 ... Used for transclusion of smart arguments for plot, axis. See function [SmartControl](#) from prodlim.

Author(s)

Thomas H. Scheike <ts@biostat.ku.dk>

Thomas A. Gerds <>tag@biostat.ku.dk>

Examples

```
library(survival)
library(prodlim)
data(Melanoma)

fit.tarr <- ARR(Hist(time,status)~strata(sex),
               data=Melanoma,
               cause=1)
plotEffects(fit.tarr)

fit.tarr <- ARR(Hist(time,status)~strata(sex)+strata(invasion),
               data=Melanoma,
               cause=1,
               times=seq(800,3000,20))
plotEffects(fit.tarr,formula=~sex)
plotEffects(fit.tarr,formula=~invasion)
plotEffects(fit.tarr,
            formula=~invasion,
            level="invasionlevel.1")

## legend arguments are transcluded:
plotEffects(fit.tarr,
            formula=~invasion,
            legend.bty="b",
            legend.cex=1)

## and other smart arguments too:
plotEffects(fit.tarr,
            formula=~invasion,
            legend.bty="b",
            axis2.las=2,
            legend.cex=1)
```

Description

Time-dependent event risk predictions.

Usage

```
plotPredictRisk(
  x,
  newdata,
  times,
  cause = 1,
  xlim,
  ylim,
  xlab,
  ylab,
  axes = TRUE,
  col,
  density,
  lty,
  lwd,
  add = FALSE,
  legend = TRUE,
  percent = FALSE,
  ...
)
```

Arguments

x	Object specifying an event risk prediction model.
newdata	A data frame with the same variable names as those that were used to fit the model x.
times	Vector of times at which to return the estimated probabilities.
cause	Show predicted risk of events of this cause
xlim	Plotting range on the x-axis.
ylim	Plotting range on the y-axis.
xlab	Label given to the x-axis.
ylab	Label given to the y-axis.
axes	Logical. If FALSE no axes are drawn.
col	Vector of colors given to the survival curve.
density	Density of the color – useful for showing many (overlapping) curves.
lty	Vector of lty's given to the survival curve.
lwd	Vector of lwd's given to the survival curve.
add	Logical. If TRUE only lines are added to an existing device
legend	Logical. If TRUE a legend is plotted by calling the function legend. Optional arguments of the function legend can be given in the form legend.x=val where x is the name of the argument and val the desired value. See also Details.

percent Logical. If TRUE the y-axis is labeled in percent.
 ... Parameters that are filtered by `SmartControl` and then passed to the functions: `plot`, `axis`, `legend`.

Details

Arguments for the invoked functions `legend` and `axis` can be specified as `legend.lty=2`. The specification is not case sensitive, thus `Legend.lty=2` or `LEGEND.lty=2` will have the same effect. The function `axis` is called twice, and arguments of the form `axis1.labels`, `axis1.at` are used for the time axis whereas `axis2.pos`, `axis1.labels`, etc., are used for the y-axis.

These arguments are processed via `...{}` of `plotPredictRisk` and inside by using the function `SmartControl`.

Value

The (invisible) object.

Author(s)

Ulla B. Mogensen and Thomas A. Gerds <tag@biostat.ku.dk>

References

Ulla B. Mogensen, Hemant Ishwaran, Thomas A. Gerds (2012). Evaluating Random Forests for Survival Analysis Using Prediction Error Curves. *Journal of Statistical Software*, 50(11), 1-23. URL <http://www.jstatsoft.org/v50/i11/>.

See Also

[plotRisk](#)

Examples

```
library(survival)
# generate survival data
# no effect
set.seed(8)
d <- sampleData(80,outcome="survival",formula = ~f(X6, 0) + f(X7, 0))
d[,table(event)]
f <- coxph(Surv(time,event)~X6+X7,data=d,x=1)
plotPredictRisk(f)

# large effect
set.seed(8)
d <- sampleData(80,outcome="survival",formula = ~f(X6, 0.1) + f(X7, -0.1))
d[,table(event)]
f <- coxph(Surv(time,event)~X6+X7,data=d,x=1)
plotPredictRisk(f)

# generate competing risk data
# small effect
```

```

set.seed(8)
d <- sampleData(40, formula = ~f(X6, 0.01) + f(X7, -0.01))
d[, table(event)]
f <- CSC(Hist(time, event)~X5+X6, data=d)
plotPredictRisk(f)

# large effect
set.seed(8)
d <- sampleData(40, formula = ~f(X6, 0.1) + f(X7, -0.1))
d[, table(event)]
f <- CSC(Hist(time, event)~X5+X6, data=d)
plotPredictRisk(f)

```

plotRisk

plot predicted risks

Description

plot predicted risks

Usage

```

plotRisk(
  x,
  models,
  times,
  xlim = c(0, 1),
  ylim = c(0, 1),
  xlab,
  ylab,
  col,
  pch,
  cex = 1,
  preclipse = 0,
  preclipse.shade = FALSE,
  ...
)

```

Arguments

x	Object obtained with function Score
models	Choice of two models to plot. The predicted risks of the first (second) are shown along the x-axis (y-axis).
times	Time point specifying the prediction horizon.
xlim	x-axis limits
ylim	y-axis limits
xlab	x-axis labels

ylab	y-axis labels
col	Colors used according to the outcome. binary outcome (two colors: no event, event), survival outcome (three colors: censored, event, no event) competing risk outcome (4 or more colors: event, competing risk 1, ..., competing risk k, censored, no event)
pch	Symbols used according to the outcome binary outcome (two symbols: no event, event), survival outcome (three symbols: censored, event, no event) competing risk outcome (4 or more symbols: event, competing risk 1, ..., competing risk k, censored, no event)
cex	point size
preclipse	Value between 0 and 1 defining the preclipse area
preclipse.shade	Logical. If TRUE shade the area of clinically meaningful change.
...	Used to control the subroutines: plot, axis, lines, barplot, legend. See SmartControl .

Details

Two rival prediction models are applied to the same data.

Value

a nice graph

Author(s)

Thomas A. Gerds <tag@biostat.ku.dk>

Examples

```
library(prodlim)
## uncensored
set.seed(10)
learndat = sampleData(40,outcome="binary")
testdat = sampleData(40,outcome="binary")
lr1 = glm(Y~X1+X2+X7+X9,data=learndat,family="binomial")
lr2 = glm(Y~X3+X5+X6,data=learndat,family="binomial")
xb=Score(list("LR(X1+X2+X7+X9)"=lr1,"LR(X3+X5+X6)"=lr2),formula=Y~1,
          data=testdat,summary="risks",null.model=0L)
plotRisk(xb)
## survival
library(survival)
set.seed(10)
learndat = sampleData(40,outcome="survival")
testdat = sampleData(40,outcome="survival")
cox1 = coxph(Surv(time,event)~X1+X2+X7+X9,data=learndat,x=TRUE)
cox2 = coxph(Surv(time,event)~X3+X5+X6,data=learndat,x=TRUE)
xs=Score(list("Cox(X1+X2+X7+X9)"=cox1,"Cox(X3+X5+X6)"=cox2),formula=Surv(time,event)~1,
          data=testdat,summary="risks",null.model=0L,times=c(3,5,6))
plotRisk(xs,times=5)
```

```
## competing risk
## Not run:
library(prodlim)
library(survival)
set.seed(8)
learndat = sampleData(80,outcome="competing.risk")
testdat = sampleData(140,outcome="competing.risk")
m1 = FGR(Hist(time,event)~X2+X7+X9,data=learndat,cause=1)
m2 = CSC(Hist(time,event)~X2+X7+X9,data=learndat,cause=1)
xcr=Score(list("FGR"=m1,"CSC"=m2),formula=Hist(time,event)~1,
            data=testdat,summary="risks",null.model=0L,times=c(3,5))
plotRisk(xcr,times=3)

## End(Not run)
```

plotROC

Plot ROC curves

Description

Plot ROC curve

Usage

```
plotROC(
  x,
  models,
  times,
  xlab = "1-Specificity",
  ylab = "Sensitivity",
  col,
  lwd,
  lty = 1,
  cex = 1,
  pch = 1,
  legend = !add,
  auc.in.legend = TRUE,
  brier.in.legend = FALSE,
  add = FALSE,
  ...
)
```

Arguments

x	Object obtained with function Score
models	Choice of models to plot
times	Time point(s) specifying the prediction horizon

xlab	Label for x-axis
ylab	Label for y-axis
col	line color
lwd	line width
lty	line style
cex	point size
pch	point style
legend	logical. If 1L draw a legend with the values of AUC.
auc.in.legend	Logical. If TRUE add AUC to legend.
brier.in.legend	Logical. If TRUE add Brier score to legend.
add	logical. If 1L add lines to an existing plot.
...	Used for additional control of the subroutines: plot, axis, lines, legend, addtable2plot. See SmartControl .

Examples

```
## binary
set.seed(18)
if (require("randomForest", quietly=TRUE)){
  library(randomForest)
  library(proclim)
  bd1 <- sampleData(40, outcome="binary")
  bdt <- sampleData(58, outcome="binary")
  bd1[, y:=factor(Y)]
  bdt[, y:=factor(Y)]
  fb1 <- glm(y~X1+X2+X3+X4+X5+X6+X7+X8+X9+X10, data=bd1, family="binomial")
  fb2 <- randomForest(y~X1+X2+X3+X4+X5+X6+X7+X8+X9+X10, data=bd1)
  xb <- Score(list("glm"=fb1, "rf"=fb2), y~1, data=bdt,
              plots="roc", metrics=c("auc", "brier"))
  plotROC(xb, brier.in.legend=1L)

# with cross-validation
## Not run:
xb3 <- Score(list("glm"=fb1, "rf"=fb2), y~1, data=bd1,
              plots="roc", B=3, split.method="bootcv",
              metrics=c("auc"))

## End(Not run)
}
## survival
set.seed(18)
library(survival)
sd1 <- sampleData(40, outcome="survival")
sd2 <- sampleData(58, outcome="survival")
fs1 <- coxph(Surv(time, event)~X3+X5+X6+X7+X8+X10, data=sd1, x=TRUE)
fs2 <- coxph(Surv(time, event)~X1+X2+X9, data=sd1, x=TRUE)
xs <- Score(list(model1=fs1, model2=fs2), Hist(time, event)~1, data=sd2,
```

```

      times=5,plots="roc",metrics="auc")
plotROC(xs)
## competing risks
data(Melanoma)
f1 <- CSC(Hist(time,status)~age+sex+epicel+ulcer,data=Melanoma)
f2 <- CSC(Hist(time,status)~age+sex+logthick+epicel+ulcer,data=Melanoma)
x <- Score(list(model1=f1,model2=f2),Hist(time,status)~1,data=Melanoma,
             cause=1,times=5*365.25,plots="roc",metrics="auc")
plotROC(x)

```

predict.CauseSpecificCox

Predicting Absolute Risk from Cause-Specific Cox Models

Description

Apply formula to combine two or more Cox models into absolute risk (cumulative incidence function).

Usage

```

## S3 method for class 'CauseSpecificCox'
predict(
  object,
  newdata,
  times,
  cause,
  type = "absRisk",
  landmark = NA,
  keep.times = 1L,
  keep.newdata = 1L,
  keep.strata = 1L,
  se = FALSE,
  band = FALSE,
  iid = FALSE,
  confint = (se + band) > 0,
  average.iid = FALSE,
  product.limit = TRUE,
  store = NULL,
  diag = FALSE,
  max.time = NULL,
  ...
)

```

Arguments

object The fitted cause specific Cox model

<code>newdata</code>	[data.frame or data.table] Contain the values of the predictor variables defining subject specific predictions relative to each cause. Should have the same structure as the data set used to fit the object.
<code>times</code>	[numeric vector] Time points at which to return the estimated absolute risk.
<code>cause</code>	[integer/character] Identifies the cause of interest among the competing events.
<code>type</code>	[character] Can be changed to "survival" if the event free survival should be output instead of the absolute risk.
<code>landmark</code>	[integer] The starting time for the computation of the cumulative risk.
<code>keep.times</code>	[logical] If TRUE add the evaluation times to the output.
<code>keep.newdata</code>	[logical] If TRUE add the value of the covariates used to make the prediction in the output list.
<code>keep.strata</code>	[logical] If TRUE add the value of the strata used to make the prediction in the output list.
<code>se</code>	[logical] If TRUE compute and add the standard errors to the output.
<code>band</code>	[logical] If TRUE compute and add the quantiles for the confidence bands to the output.
<code>iid</code>	[logical] If TRUE compute and add the influence function to the output.
<code>confint</code>	[logical] If TRUE compute and add the confidence intervals/bands to the output. They are computed applying the <code>confint</code> function to the output.
<code>average.iid</code>	[logical]. If TRUE add the average of the influence function over <code>newdata</code> to the output.
<code>product.limit</code>	[logical]. If TRUE the survival is computed using the product limit estimator. Otherwise the exponential approximation is used (i.e. $\exp(-\text{cumulative hazard})$).
<code>store</code>	[vector of length 2] Whether prediction should only be computed for unique covariate sets and mapped back to the original dataset (<code>data="minimal"</code>) and whether the influence function should be stored in a memory efficient way (<code>iid="minimal"</code>). Otherwise use <code>data="full"</code> or <code>iid="full"</code> .
<code>diag</code>	[logical] when FALSE the absolute risk/survival for all observations at all times is computed, otherwise it is only computed for the <i>i</i> -th observation at the <i>i</i> -th time.
<code>max.time</code>	[numeric] maximum time of the response of the fitted data. Only relevant if model response element has been removed
<code>...</code>	not used.

Details

This function computes the absolute risk as given by formula 2 of (Ozenne et al., 2017). Confidence intervals and confidence bands can be computed using a first order von Mises expansion. See the section "Construction of the confidence intervals" in (Ozenne et al., 2017).

A detailed explanation about the meaning of the argument `store = c(iid="full")` can be found in (Ozenne et al., 2017) Appendix B "Saving the influence functions".

Note: for Cox regression models with time varying covariates it does not make sense to use this function, because the predicted risk has to be a measurable function of the data available at the time origin.

The iid decomposition is output using an array containing the value of the influence of each subject used to fit the object (dim 1), for each subject in newdata (dim 3), and each time (dim 2).

Author(s)

Brice Ozenne broz@sund.ku.dk, Thomas A. Gerds tag@biostat.ku.dk

References

Brice Ozenne, Anne Lyngholm Sorensen, Thomas Scheike, Christian Torp-Pedersen and Thomas Alexander Gerds. *riskRegression: Predicting the Risk of an Event using Cox Regression Models*. *The R Journal* (2017) 9:2, pages 440-460.

See Also

[confint.predictCSC](#) to compute confidence intervals/bands. [autoplot.predictCSC](#) to display the predictions.

Examples

```
library(survival)
library(prodlim)
library(data.table)

#####
#### generate data ####
#####

set.seed(5)
d <- sampleData(80,outcome="comp") ## training dataset
nd <- sampleData(4,outcome="comp") ## validation dataset
d$time.round <- round(d$time,1) ## create tied events
ttt <- sort(sample(x = unique(d$time), size = 10))

#####
#### Aalen-Johansen ####
#####

#### no ties
fit.AE <- CSC(Hist(time, event) ~ 1, data = d)
GS.AE <- prodlim(Hist(time, event) ~ 1, data = d)

## product limit
ePL.AE <- predict(fit.AE, newdata = d[1], times = GS.AE$time,
                  product.limit = TRUE, cause = 1)
range(ePL.AE$absRisk - GS.AE$cuminc[[1]]) ## same

## exponential approximation
predict(fit.AE, newdata = d[1], times = GS.AE$time,
        product.limit = FALSE, cause = 1)

#### with ties (use Breslow estimator instead of Efron to match prodlim)
```

```

fit.AE.ties <- CSC(Hist(time.round, event) ~ 1, ties = "breslow",
                  data = as.data.frame(d)[c("time.round","event")])
GS.AE.ties <- prodlim(Hist(time.round, event) ~ 1,
                     data = as.data.frame(d)[c("time.round","event")])

## product limit
ePL.AE.ties <- predict(fit.AE.ties, newdata = d[1], times = GS.AE.ties$time,
                      product.limit = TRUE, cause = 1)
range(ePL.AE.ties$absRisk - GS.AE.ties$cuminc[[1]]) ## same

#####
#### Stratified Aalen-Johansen ####
#####

#### no ties
fit.SAE <- CSC(Hist(time, event) ~ strata(X1), data = d)
GS.SAE <- prodlim(Hist(time, event) ~ X1, data = d)

## product limit
index.strata1 <- GS.SAE$first.strata[1]:(GS.SAE$first.strata[2]-1)
ePL.SAE <- predict(fit.SAE, newdata = d[1],
                  times = GS.SAE$time[index.strata1],
                  product.limit = TRUE, cause = 1)
range(ePL.SAE$absRisk - GS.SAE$cuminc[[1]][index.strata1]) ## same

## exponential approximation
predict(fit.SAE, newdata = d[1:10], times = 1:10, product.limit = FALSE)

#####
#### Cause-specific Cox ####
#####

## estimate a CSC model based on the coxph function
CSC.fit <- CSC(Hist(time,event)~ X3+X8, data=d, method = "breslow")

## compute the absolute risk of cause 1, in the validation dataset
## at time 1:10
CSC.risk <- predict(CSC.fit, newdata=nd, times=1:10, cause=1)
CSC.risk

## compute absolute risks with CI for cause 2
## (without displaying the value of the covariates)
predict(CSC.fit,newdata=nd,times=1:10,cause=2,se=TRUE,
        keep.newdata = FALSE)

## other example
library(survival)
CSC.fit.s <- CSC(list(Hist(time,event)~ strata(X1)+X2+X9,
                    Hist(time,event)~ X2+strata(X4)+X8+X7),data=d, method = "breslow")
predict(CSC.fit.s,cause=1,times=ttt,se=1L) ## note: absRisk>1 due to small number of observations

## using the cph function instead of coxph
CSC.cph <- CSC(Hist(time,event)~ X1+X2,data=d, method = "breslow", fitter = "cph")#'
```

```

predict(CSC.cph, newdata = d, cause = 2, times = ttt)

## landmark analysis
T0 <- 1
predCSC.afterT0 <- predict(CSC.fit, newdata = d, cause = 2, times = ttt[ttt>T0], landmark = T0)
predCSC.afterT0

```

predict.FGR	<i>Predict subject specific risks (cumulative incidence) based on Fine-Gray regression model</i>
-------------	--

Description

Predict subject specific risks (cumulative incidence) based on Fine-Gray regression model

Usage

```

## S3 method for class 'FGR'
predict(object, newdata, times, ...)

```

Arguments

object	Result of call to FGR
newdata	Predictor values of subjects for who to predict risks
times	Time points at which to evaluate the risks
...	passed to predict.crr

Examples

```

library(prodlm)
library(survival)
set.seed(10)
d <- sampleData(101, outcome = "competing.risk")
tFun<-function(t) {t}
fgr<-FGR(Hist(time, event)~X1+strata(X2)+X6+cov2(X7, tf=tFun),
         data=d, cause=1)
predictRisk(fgr, times=5, newdata=d[1:10])

```

```
predict.riskRegression
```

Predict individual risk.

Description

Extract predictions from a risk prediction model.

Usage

```
## S3 method for class 'riskRegression'  
predict(object, newdata, ...)
```

Arguments

object	Fitted object obtained with one of ARR, LRR, riskRegression.
newdata	A data frame containing predictor variable combinations for which to compute predicted risk.
...	not used

Author(s)

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Thomas A. Gerds <>tag@biostat.ku.dk>

References

Gerds, TA and Scheike, T and Andersen, PK (2011) Absolute risk regression for competing risks: interpretation, link functions and prediction Research report 11/8. Department of Biostatistics, University of Copenhagen

Examples

```
data(Melanoma)  
library(proplim)  
library(survival)  
  
fit.tarr <- ARR(Hist(time,status)~age+invasion+strata(sex),data=Melanoma,cause=1)  
predict(fit.tarr,newdata=data.frame(age=48,  
  invasion=factor("level.1",  
    levels=levels(Melanoma$invasion)),  
  sex=factor("Female",levels=levels(Melanoma$sex))))  
predict(fit.tarr,newdata=data.frame(age=48,  
  invasion=factor("level.1",  
    levels=levels(Melanoma$invasion)),  
  sex=factor("Male",levels=levels(Melanoma$sex))))  
predict(fit.tarr,newdata=data.frame(age=c(48,58,68),  
  invasion=factor("level.1",
```

```

      levels=levels(Melanoma$invasion)),
      sex=factor("Male", levels=levels(Melanoma$sex))))
predict(fit.tarr, newdata=Melanoma[1:4,])

```

predictCox	<i>Survival probabilities, hazards and cumulative hazards from Cox regression models</i>
------------	--

Description

Routine to get baseline hazards and subject specific hazards as well as survival probabilities from a `survival::coxph` or `rms::cph` object.

Usage

```

predictCox(
  object,
  times,
  newdata = NULL,
  centered = TRUE,
  type = c("cumhazard", "survival"),
  keep.strata = TRUE,
  keep.times = TRUE,
  keep.newdata = FALSE,
  keep.infoVar = FALSE,
  se = FALSE,
  band = FALSE,
  iid = FALSE,
  confint = (se + band) > 0,
  diag = FALSE,
  average.iid = FALSE,
  product.limit = FALSE,
  store = NULL
)

```

Arguments

object	The fitted Cox regression model object either obtained with <code>coxph</code> (survival package) or <code>cph</code> (rms package).
times	[numeric vector] Time points at which to return the estimated hazard/cumulative hazard/survival.
newdata	[data.frame or data.table] Contain the values of the predictor variables defining subject specific predictions. Should have the same structure as the data set used to fit the object.

centered	[logical] If TRUE return prediction at the mean values of the covariates <code>fit\$mean</code> , if FALSE return a prediction for all covariates equal to zero. in the linear predictor. Will be ignored if argument <code>newdata</code> is used. For internal use.
type	[character vector] the type of predicted value. Choices are <ul style="list-style-type: none"> • "hazard" the baseline hazard function when argument <code>newdata</code> is not used and the hazard function when argument <code>newdata</code> is used. • "cumhazard" the cumulative baseline hazard function when argument <code>newdata</code> is not used and the cumulative hazard function when argument <code>newdata</code> is used. • "survival" the survival baseline hazard function when argument <code>newdata</code> is not used and the cumulative hazard function when argument <code>newdata</code> is used. <p>Several choices can be combined in a vector of strings that match (no matter the case) strings "hazard", "cumhazard", "survival".</p>
keep.strata	[logical] If TRUE add the (<code>newdata</code>) strata to the output. Only if there any.
keep.times	[logical] If TRUE add the evaluation times to the output.
keep.newdata	[logical] If TRUE add the value of the covariates used to make the prediction in the output list.
keep.infoVar	[logical] For internal use.
se	[logical] If TRUE compute and add the standard errors to the output.
band	[logical] If TRUE compute and add the quantiles for the confidence bands to the output.
iid	[logical] If TRUE compute and add the influence function to the output.
confint	[logical] If TRUE compute and add the confidence intervals/bands to the output. They are computed applying the <code>confint</code> function to the output.
diag	[logical] when FALSE the hazard/cumulative hazard/survival for all observations at all times is computed, otherwise it is only computed for the <i>i</i> -th observation at the <i>i</i> -th time.
average.iid	[logical] If TRUE add the average of the influence function over <code>newdata</code> to the output.
product.limit	[logical]. If TRUE the survival is computed using the product limit estimator. Otherwise the exponential approximation is used (i.e. $\exp(-\text{cumulative hazard})$).
store	[vector of length 2] Whether prediction should only be computed for unique covariate sets and mapped back to the original dataset (<code>data="minimal"</code>) and whether the influence function should be stored in a memory efficient way (<code>iid="minimal"</code>). Otherwise use <code>data="full"</code> and/or <code>iid="full"</code> .

Details

When the argument `newdata` is not specified, the function computes the baseline hazard estimate. See (Ozenne et al., 2017) section "Handling of tied event times".

Otherwise the function computes survival probabilities with confidence intervals/bands. See (Ozenne et al., 2017) section "Confidence intervals and confidence bands for survival probabilities". The survival is computed using the exponential approximation (equation 3).

A detailed explanation about the meaning of the argument `store = c(iid="full")` can be found in (Ozenne et al., 2017) Appendix B "Saving the influence functions".

The function is not compatible with time varying predictor variables.

The centered argument enables us to reproduce the results obtained with the `basehaz` function from the `survival` package but should not be modified by the user.

Value

The iid decomposition is output in an attribute called `"iid"`, using an array containing the value of the influence of each subject used to fit the object (dim 1), for each subject in `newdata` (dim 3), and each time (dim 2).

Author(s)

Brice Ozenne broz@sund.ku.dk, Thomas A. Gerds tag@biostat.ku.dk

References

Brice Ozenne, Anne Lyngholm Sorensen, Thomas Scheike, Christian Torp-Pedersen and Thomas Alexander Gerds. `riskRegression`: Predicting the Risk of an Event using Cox Regression Models. *The R Journal* (2017) 9:2, pages 440-460.

See Also

[confint.predictCox](#) to compute confidence intervals/bands. [autoplot.predictCox](#) to display the predictions.

Examples

```
library(survival)
library(data.table)

#####
#### generate data ####
#####

set.seed(10)
d <- sampleData(50,outcome="survival") ## training dataset
nd <- sampleData(5,outcome="survival") ## validation dataset

## add ties
d$time.round <- round(d$time,1)
any(duplicated(d$time.round))

## add categorical variables
d$U <- sample(letters[1:3],replace=TRUE,size=NROW(d))
d$V <- sample(letters[5:8],replace=TRUE,size=NROW(d))
nd <- cbind(nd,d[sample(NROW(d),replace=TRUE,size=NROW(nd)),c("U","V")])

#####
#### Kaplan Meier ####
```

```
#####

#### no ties
fit.KM <- coxph(Surv(time,event)~ 1, data=d, x = TRUE, y = TRUE)
predictCox(fit.KM) ## exponential approximation
ePL.KM <- predictCox(fit.KM, product.limit = TRUE) ## product-limit
as.data.table(ePL.KM)

range(survfit(Surv(time,event)~1, data = d)$surv - ePL.KM$survival) ## same

#### with ties (exponential approximation, Efron estimator for the baseline hazard)
fit.KM.ties <- coxph(Surv(time.round,event)~ 1, data=d, x = TRUE, y = TRUE)
predictCox(fit.KM)

## retrieving survfit results with ties
fit.KM.ties <- coxph(Surv(time.round,event)~ 1, data=d,
                    x = TRUE, y = TRUE, ties = "breslow")
ePL.KM.ties <- predictCox(fit.KM, product.limit = TRUE)
range(survfit(Surv(time,event)~1, data = d)$surv - ePL.KM.ties$survival) ## same

#####
#### Stratified Kaplan Meier ####
#####

fit.SKM <- coxph(Surv(time,event)~strata(X2), data=d, x = TRUE, y = TRUE)
ePL.SKM <- predictCox(fit.SKM, product.limit = TRUE)
ePL.SKM

range(survfit(Surv(time,event)~X2, data = d)$surv - ePL.SKM$survival) ## same

#####
#### Cox model ####
#####

fit.Cox <- coxph(Surv(time,event)~X1 + X2 + X6, data=d, x = TRUE, y = TRUE)

#### compute the baseline cumulative hazard
Cox.haz0 <- predictCox(fit.Cox)
Cox.haz0

range(survival::basehaz(fit.Cox)$hazard-Cox.haz0$cumhazard) ## same

#### compute individual specific cumulative hazard and survival probabilities
## exponential approximation
fit.predCox <- predictCox(fit.Cox, newdata=nd, times=c(3,8),
                        se = TRUE, band = TRUE)

fit.predCox

## product-limit
fitPL.predCox <- predictCox(fit.Cox, newdata=nd, times=c(3,8),
                          product.limit = TRUE, se = TRUE)

fitPL.predCox
```

```

## Note: product limit vs. exponential does not affect uncertainty quantification
range(fitPL.predCox$cumhazard.se - fit.predCox$cumhazard.se) ## same
range(fitPL.predCox$survival.se - fit.predCox$survival.se) ## different
## except through a multiplicative factor (multiply by survival in the delta method)
fitPL.predCox$survival.se - fitPL.predCox$cumhazard.se * fitPL.predCox$survival

#### left truncation
test2 <- list(start=c(1,2,5,2,1,7,3,4,8,8),
              stop=c(2,3,6,7,8,9,9,9,14,17),
              event=c(1,1,1,1,1,1,1,1,0,0),
              x=c(1,0,0,1,0,1,1,1,0,0))
m.cph <- coxph(Surv(start, stop, event) ~ 1, test2, x = TRUE)
as.data.table(predictCox(m.cph))

basehaz(m.cph)

#####
#### Stratified Cox model ####
#####

#### one strata variable
fit.SCox <- coxph(Surv(time,event)~X1+strata(X2)+X6, data=d, x = TRUE, y = TRUE)
predictCox(fit.SCox, newdata=nd, times = c(3,12))
predictCox(fit.SCox, newdata=nd, times = c(3,12), product.limit = TRUE)
## NA: timepoint is beyond the last observation in the strata and censoring
tapply(d$time,d$X2,max)

#### two strata variables
fit.S2Cox <- coxph(Surv(time,event)~X1+strata(U)+strata(V)+X2,
                  data=d, x = TRUE, y = TRUE)

base.S2Cox <- predictCox(fit.S2Cox)
range(survival::basehaz(fit.S2Cox)$hazard - base.S2Cox$cumhazard) ## same
predictCox(fit.S2Cox, newdata=nd, times = 3)

```

predictCoxPL

Deprecated Function for Product Limit Estimation of Survival Probabilities .

Description

Deprecated function for Product Limit Estimation of Survival Probabilities from a Cox model. Use the [predictCox](#) function instead with argument `product.limit=TRUE`.

Usage

```
predictCoxPL(object, ...)
```

Arguments

object The fitted Cox regression model object either obtained with `coxph` (survival package) or `cph` (rms package).
... additional arguments to be passed to `predictCox`.

predictRisk *Extracting predicting risks from regression models*

Description

Extract event probabilities from fitted regression models and machine learning objects. The function `predictRisk` is a generic function, meaning that it invokes specifically designed functions depending on the 'class' of the first argument. See `predictRisk`.

Usage

```
predictRisk(object, newdata, ...)  
  
## Default S3 method:  
predictRisk(object, newdata, times, cause, ...)  
  
## S3 method for class 'double'  
predictRisk(object, newdata, times, cause, ...)  
  
## S3 method for class 'integer'  
predictRisk(object, newdata, times, cause, ...)  
  
## S3 method for class 'factor'  
predictRisk(object, newdata, times, cause, ...)  
  
## S3 method for class 'numeric'  
predictRisk(object, newdata, times, cause, ...)  
  
## S3 method for class 'glm'  
predictRisk(object, newdata, iid = FALSE, average.iid = FALSE, ...)  
  
## S3 method for class 'multinom'  
predictRisk(  
  object,  
  newdata,  
  iid = FALSE,  
  average.iid = FALSE,  
  cause = NULL,  
  ...  
)
```

```
## S3 method for class 'formula'
predictRisk(object, newdata, ...)

## S3 method for class 'BinaryTree'
predictRisk(object, newdata, ...)

## S3 method for class 'lrm'
predictRisk(object, newdata, ...)

## S3 method for class 'rpart'
predictRisk(object, newdata, ...)

## S3 method for class 'randomForest'
predictRisk(object, newdata, ...)

## S3 method for class 'matrix'
predictRisk(object, newdata, times, cause, ...)

## S3 method for class 'aalen'
predictRisk(object, newdata, times, ...)

## S3 method for class 'cox.aalen'
predictRisk(object, newdata, times, ...)

## S3 method for class 'comprisk'
predictRisk(object, newdata, times, ...)

## S3 method for class 'coxph'
predictRisk(
  object,
  newdata,
  times,
  product.limit = FALSE,
  diag = FALSE,
  iid = FALSE,
  average.iid = FALSE,
  ...
)

## S3 method for class 'coxphTD'
predictRisk(object, newdata, times, landmark, ...)

## S3 method for class 'CSCTD'
predictRisk(object, newdata, times, cause, landmark, ...)

## S3 method for class 'coxph.penall'
predictRisk(object, newdata, times, ...)
```



```
## S3 method for class 'cph'
predictRisk(
  object,
  newdata,
  times,
  product.limit = FALSE,
  diag = FALSE,
  iid = FALSE,
  average.iid = FALSE,
  ...
)

## S3 method for class 'selectCox'
predictRisk(object, newdata, times, ...)

## S3 method for class 'prodlm'
predictRisk(
  object,
  newdata,
  times,
  cause,
  diag = FALSE,
  iid = FALSE,
  average.iid = FALSE,
  ...
)

## S3 method for class 'survfit'
predictRisk(object, newdata, times, ...)

## S3 method for class 'psm'
predictRisk(object, newdata, times, ...)

## S3 method for class 'ranger'
predictRisk(object, newdata, times, cause, ...)

## S3 method for class 'rfsrc'
predictRisk(object, newdata, times, cause, ...)

## S3 method for class 'FGR'
predictRisk(object, newdata, times, cause, ...)

## S3 method for class 'riskRegression'
predictRisk(object, newdata, times, cause, ...)

## S3 method for class 'ARR'
predictRisk(object, newdata, times, cause, ...)
```

```
## S3 method for class 'CauseSpecificCox'
predictRisk(
  object,
  newdata,
  times,
  cause,
  product.limit = TRUE,
  diag = FALSE,
  iid = FALSE,
  average.iid = FALSE,
  truncate = FALSE,
  ...
)

## S3 method for class 'penfitS3'
predictRisk(object, newdata, times, ...)

## S3 method for class 'SuperPredictor'
predictRisk(object, newdata, ...)

## S3 method for class 'gbm'
predictRisk(object, newdata, times, ...)

## S3 method for class 'flexsurvreg'
predictRisk(object, newdata, times, ...)

## S3 method for class 'Hal9001'
predictRisk(object, newdata, times, cause, ...)

## S3 method for class 'GLMnet'
predictRisk(object, newdata, times = NA, ...)

## S3 method for class 'singleEventCB'
predictRisk(object, newdata, times, cause, ...)

## S3 method for class 'wglm'
predictRisk(
  object,
  newdata,
  times = NULL,
  product.limit = NULL,
  diag = FALSE,
  iid = FALSE,
  average.iid = FALSE,
  ...
)

## S3 method for class 'CoxConfidential'
```

```
predictRisk(object, newdata, ...)
```

Arguments

object	A fitted model from which to extract predicted event probabilities.
newdata	A data frame containing predictor variable combinations for which to compute predicted event probabilities.
...	Additional arguments that are passed on to the current method.
times	A vector of times in the range of the response variable, for which the cumulative incidences event probabilities are computed.
cause	Identifies the cause of interest among the competing events.
iid	Should the iid decomposition be output using an attribute?
average.iid	Should the average iid decomposition be output using an attribute?
product.limit	If TRUE the survival is computed using the product limit estimator. Otherwise the exponential approximation is used (i.e. $\exp(-\text{cumulative hazard})$).
diag	when FALSE the hazard/cumulative hazard/survival for all observations at all times is computed, otherwise it is only computed for the i-th observation at the i-th time.
landmark	The starting time for the computation of the cumulative risk.
truncate	If TRUE truncates the predicted risks to be in the range [0, 1]. For now only implemented for the Cause Specific Cox model.

Details

In uncensored binary outcome data there is no need to choose a time point.

When operating on models for survival analysis (without competing risks) the function still predicts the risk, as $1 - S(t|X)$ where $S(t|X)$ is survival chance of a subject characterized by X .

When there are competing risks (and the data are right censored) one needs to specify both the time horizon for prediction (can be a vector) and the cause of the event. The function then extracts the absolute risks $F_c(t|X)$ aka the cumulative incidence of an event of type/cause c until time t for a subject characterized by X . Depending on the model it may or not be possible to predict the risk of all causes in a competing risks setting. For example, a cause-specific Cox (CSC) object allows to predict both cases whereas a Fine-Gray regression model (FGR) is specific to one of the causes.

Value

For binary outcome a vector with predicted risks. For survival outcome with and without competing risks a matrix with as many rows as `NROW(newdata)` and as many columns as `length(times)`. Each entry is a probability and in rows the values should be increasing.

Author(s)

Thomas A. Gerds <tag@biostat.ku.dk>

Examples

```

## binary outcome
library(rms)
set.seed(7)
d <- sampleData(80,outcome="binary")
nd <- sampleData(80,outcome="binary")
fit <- lrm(Y~X1+X8,data=d)
predictRisk(fit,newdata=nd)

## survival outcome
# generate survival data
library(prodlim)
set.seed(100)
d <- sampleData(100,outcome="survival")
d[,X1:=as.numeric(as.character(X1))]
d[,X2:=as.numeric(as.character(X2))]
# then fit a Cox model
library(rms)
cphmodel <- cph(Surv(time,event)~X1+X2,data=d,surv=TRUE,x=TRUE,y=TRUE)
# or via survival
library(survival)
coxphmodel <- coxph(Surv(time,event)~X1+X2,data=d,x=TRUE,y=TRUE)

# Extract predicted survival probabilities
# at selected time-points:
ttt <- quantile(d$time)
# for selected predictor values:
ndat <- data.frame(X1=c(0.25,0.25,-0.05,0.05),X2=c(0,1,0,1))
# as follows
predictRisk(cphmodel,newdata=ndat,times=ttt)
predictRisk(coxphmodel,newdata=ndat,times=ttt)

## simulate learning and validation data
set.seed(10)
learndat <- sampleData(80,outcome="survival")
valdat <- sampleData(10,outcome="survival")
## use the learning data to fit a Cox model
library(survival)
fitCox <- coxph(Surv(time,event)~X6+X2,data=learndat,x=TRUE,y=TRUE)
## suppose we want to predict the survival probabilities for all subjects
## in the validation data at the following time points:
## 0, 1, 2, 3, 4
psurv <- predictRisk(fitCox,newdata=valdat,times=seq(0,4,1))
## This is a matrix with event probabilities (1-survival)
## one column for each of the 5 time points
## one row for each validation set individual

## competing risks
library(survival)
library(riskRegression)
library(prodlim)
set.seed(8)

```

```

train <- sampleData(80)
test <- sampleData(10)
cox.fit <- CSC(Hist(time,event)~X1+X6,data=train,cause=1)
predictRisk(cox.fit,newdata=test,times=seq(1:10),cause=1)

## with strata
cox.fit2 <- CSC(list(Hist(time,event)~strata(X1)+X6,
                    Hist(time,cause)~X1+X6),data=train)
predictRisk(cox.fit2,newdata=test,times=seq(1:10),cause=1)

```

print.ate *Print Average Treatment Effects*

Description

Print average treatment effects.

Usage

```

## S3 method for class 'ate'
print(x, estimator = x$estimator, ...)

```

Arguments

x	object obtained with function ate
estimator	[character] The type of estimator relative to which the risks should be output.
...	for internal use

See Also

[summary.ate](#) to obtain a more detailed output [confint.ate](#) to compute confidence intervals/bands.
[ate](#) to compute the average treatment effects.

print.CauseSpecificCox
Print of a Cause-Specific Cox regression model

Description

Print of a Cause-Specific Cox regression model

Usage

```

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

```

Arguments

x	Object obtained with CSC
...	Passed to print

print.FGR	<i>Print of a Fine-Gray regression model</i>
-----------	--

Description

Print of a Fine-Gray regression model

Usage

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

Arguments

x	Object fitted with function FGR
...	passed to cmprsk::summary.crr

print.influenceTest	<i>Output of the Difference Between Two Estimates</i>
---------------------	---

Description

Output of the difference between two estimates.

Usage

```
## S3 method for class 'influenceTest'
print(x, digits = 3, ...)
```

Arguments

x	object obtained with the function <code>influenceTest</code> .
digits	[integer, >0] indicating the number of decimal places.
...	Passed to print.

Details

to display confidence intervals/bands, the `confint` method needs to be applied on the object.

See Also

[confint.influenceTest](#) to compute confidence intervals/bands. [influenceTest](#) to perform the comparison.

print.IPA	<i>Print IPA object</i>
-----------	-------------------------

Description

Print method for IPA

Usage

```
## S3 method for class 'IPA'
print(x, digits = 2, ...)
```

Arguments

x	Object obtained with IPA
digits	Number of digits
...	passed to print

Author(s)

Thomas A. Gerds <tag@biostat.ku.dk>

print.predictCox	<i>Print Predictions From a Cox Model</i>
------------------	---

Description

Print predictions from a Cox model.

Usage

```
## S3 method for class 'predictCox'
print(x, digits = 3, ...)
```

Arguments

x	object obtained with the function predictCox.
digits	[integer, >0] indicating the number of decimal places.
...	Passed to print.

Details

to display confidence intervals/bands, the confint method needs to be applied on the object.

See Also

[confint.predictCox](#) to compute confidence intervals/bands. [predictCox](#) to compute the predicted cumulative hazard/survival.

<code>print.predictCSC</code>	<i>Print Predictions From a Cause-specific Cox Proportional Hazard Regression</i>
-------------------------------	---

Description

Print predictions from a Cause-specific Cox proportional hazard regression.

Usage

```
## S3 method for class 'predictCSC'
print(x, digits = 3, ...)
```

Arguments

<code>x</code>	object obtained with the function <code>predictCox</code> .
<code>digits</code>	[integer, >0] indicating the number of decimal places.
<code>...</code>	Passed to <code>print</code> .

Details

to display confidence intervals/bands, the `confint` method needs to be applied on the object.

See Also

[confint.predictCSC](#) to compute confidence intervals/bands. [predict.CauseSpecificCox](#) to compute the predicted risks.

<code>print.riskRegression</code>	<i>Print function for riskRegression models</i>
-----------------------------------	---

Description

Print function for riskRegression models

Usage

```
## S3 method for class 'riskRegression'
print(x, times, digits = 3, eps = 10^-4, verbose = TRUE, conf.int = 0.95, ...)
```


Arguments

x	Object obtained with ARR, LRR or riskRegression
times	Time points at which to show time-dependent coefficients
digits	Number of digits for all numbers but p-values
eps	p-values smaller than this number are shown as such
verbose	Level of verbosity
conf.int	level of confidence. default is 0.95
...	not used

print.Score *Print Score object*

Description

Print method for risk prediction scores

Usage

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

Arguments

x	Object obtained with Score.list
digits	Number of digits
...	passed to print

print.subjectWeights *Print subject weights*

Description

Print subject weights

Usage

```
## S3 method for class 'subjectWeights'
print(x, digits = 3, ...)
```

Arguments

x	Subject weights
digits	Digits
...	not used

reconstructData	<i>Reconstruct the original dataset</i>
-----------------	---

Description

Reconstruct the original dataset from the elements stored in the coxph object

Usage

```
reconstructData(object)
```

Arguments

object a coxph object.

Author(s)

Brice Ozenne broz@sund.ku.dk and Thomas A. Gerds tag@biostat.ku.dk

riskLevelPlot	<i>Level plots for risk prediction models</i>
---------------	---

Description

Level plots for predicted risks

Usage

```
riskLevelPlot(
  object,
  formula,
  data = parent.frame(),
  horizon = NULL,
  cause = 1,
  ...
)
```

Arguments

object	risk prediction model object
formula	formula
data	data
horizon	time point
cause	cause of interst
...	passed to lattice::levelplot

Details

Level plots for predicted risks

Author(s)

Thomas A. Gerds <tag@biostat.ku.dk>

Examples

```
# ----- logistic regression -----
expit <- function(x){exp(x)/(1+exp(x))}
partyData <- function(N){
  Age <- runif(N,.5,15)
  Parasites <- rnorm(N,mean=3.5-0.03*Age)
  Fever <- factor(rbinom(N,1,expit(-3.5-.3*Age+.55*Parasites+0.15*Age*Parasites)))
  data.frame(Fever, Age, Parasites)
}
d <- partyData(100)
f <- glm(Fever~Age+Parasites,data=d,family="binomial")
riskLevelPlot(f,Fever~Age+Parasites,d)
## Not run:
if (require("randomForest",quietly=TRUE)){
  rf <- randomForest::randomForest(Fever~Age+Parasites,data=d)
  riskLevelPlot(f,Fever~Age+Parasites,d)
  riskLevelPlot(rf,Fever~Age+Parasites,d)
}

## End(Not run)

# ----- survival analysis -----

# --simulate an artificial data frame
# with survival response and three predictors

library(survival)
library(prodlim)
set.seed(140515)
sdat <- sampleData(43,outcome="survival")
# -- fit a Cox regression model
survForm = Surv(time,event) ~ X8 + X9
cox <- coxph(survForm, data = sdat,x=TRUE)

# --choose a time horizon for the predictions and plot the risks
timeHorizon <- floor(median(sdat$time))
riskLevelPlot(cox, survForm, data = sdat, horizon = timeHorizon)

# ----- competing risks -----

# -- simulate an artificial data frame
# with competing cause response and three predictors
library(cmprsk)
library(riskRegression)
```

```

set.seed(140515)
crdat <- sampleData(49)

# -- fit a cause-specific Cox regression model
crForm <- Hist(time,event)~X8+X9
csCox <- CSC(crForm, data=crdat)

# -- choose a time horizon and plot the risk for a given cause
timeHorizon <- floor(median(crdat$time))
riskLevelPlot(csCox, crForm, data = crdat, horizon = timeHorizon, cause = 1)

```

riskRegression	<i>Risk Regression Fits a regression model for the risk of an event – allowing for competing risks.</i>
----------------	---

Description

This is a wrapper for the function `comp.risk` from the `timereg` package. The main difference is one marks variables in the formula that should have a time-dependent effect whereas in `comp.risk` one marks variables that should have a time constant (proportional) effect.

Usage

```

riskRegression(
  formula,
  data,
  times,
  link = "relative",
  cause,
  conf.int = TRUE,
  cens.model,
  cens.formula,
  max.iter = 50,
  conservative = TRUE,
  ...
)

```

Arguments

formula	Formula where the left hand side specifies the event history <code>event.history</code> and the right hand side the linear predictor. See examples.
data	The data for fitting the model in which includes all the variables included in formula.
times	Vector of times. For each time point in <code>times</code> estimate the baseline risk and the timevarying coefficients.

link	"relative" for the absolute risk regression model. "logistic" for the logistic risk regression model. "prop" for the Fine-Gray regression model.
cause	The cause of interest.
conf.int	If TRUE return the iid decomposition, that can be used to construct confidence bands for predictions.
cens.model	Specified the model for the (conditional) censoring distribution used for deriving weights (IFPW). Defaults to "KM" (the Kaplan-Meier method ignoring covariates) alternatively it may be "Cox" (Cox regression).
cens.formula	Right hand side of the formula used for fitting the censoring model. If not specified the right hand side of formula is used.
max.iter	Maximal number of iterations.
conservative	If TRUE use variance formula that ignores the contribution by the estimate of the inverse of the probability of censoring weights
...	Further arguments passed to <code>comp.risk</code>

Author(s)

Thomas A. Gerds <tag@biostat.ku.dk>, Thomas H. Scheike <ts@biostat.ku.dk>

References

Thomas A Gerds, Thomas H Scheike, and Per K Andersen. Absolute risk regression for competing risks: interpretation, link functions, and prediction. *Statistics in medicine*, 31(29):3921–3930, 2012.

Scheike, Zhang and Gerds (2008), Predicting cumulative incidence probability by direct binomial regression, *Biometrika*, 95, 205-220.

Scheike and Zhang (2007), Flexible competing risks regression modelling and goodness of fit, *LIDA*, 14, 464-483.

Martinussen and Scheike (2006), *Dynamic regression models for survival data*, Springer.

Examples

```
library(prodlim)
data(Melanoma,package="riskRegression")
## tumor thickness on the log-scale
Melanoma$logthick <- log(Melanoma$thick)

# Single binary factor

## absolute risk regression
library(survival)
library(prodlim)
fit.arr <- ARR(Hist(time,status)~sex,data=Melanoma,cause=1)
print(fit.arr)
# show predicted cumulative incidences
plot(fit.arr,col=3:4,newdata=data.frame(sex=c("Female","Male")))

## compare with non-parametric Aalen-Johansen estimate
```

```

library(prodlim)
fit.aj <- prodlim(Hist(time,status)~sex,data=Melanoma)
plot(fit.aj,conf.int=FALSE)
plot(fit.arr,add=TRUE,col=3:4,newdata=data.frame(sex=c("Female","Male")))

## with time-dependent effect
fit.tarr <- ARR(Hist(time,status)~strata(sex),data=Melanoma,cause=1)
plot(fit.tarr,newdata=data.frame(sex=c("Female","Male")))

## logistic risk regression
fit.lrr <- LRR(Hist(time,status)~sex,data=Melanoma,cause=1)
summary(fit.lrr)

# Single continuous factor

## tumor thickness on the log-scale
Melanoma$logthick <- log(Melanoma$thick)

## absolute risk regression
fit2.arr <- ARR(Hist(time,status)~logthick,data=Melanoma,cause=1)
print(fit2.arr)
# show predicted cumulative incidences
plot(fit2.arr,col=1:5,newdata=data.frame(logthick=quantile(Melanoma$logthick)))

## comparison with nearest neighbor non-parametric Aalen-Johansen estimate
library(prodlim)
fit2.aj <- prodlim(Hist(time,status)~logthick,data=Melanoma)
plot(fit2.aj,conf.int=FALSE,newdata=data.frame(logthick=quantile(Melanoma$logthick)))
plot(fit2.arr,add=TRUE,col=1:5,lty=3,newdata=data.frame(logthick=quantile(Melanoma$logthick)))

## logistic risk regression
fit2.lrr <- LRR(Hist(time,status)~logthick,data=Melanoma,cause=1)
summary(fit2.lrr)

## change model for censoring weights
library(rms)
fit2a.lrr <- LRR(Hist(time,status)~logthick,
                data=Melanoma,
                cause=1,
                cens.model="cox",
                cens.formula=~sex+epicel+ulcer+age+logthick)
summary(fit2a.lrr)

## compare prediction performance
Score(list(ARR=fit2.arr,AJ=fit2.aj,LRR=fit2.lrr),formula=Hist(time,status)~1,data=Melanoma)

# multiple regression
library(riskRegression)
library(prodlim)
# absolute risk model
multi.arr <- ARR(Hist(time,status)~logthick+sex+age+ulcer,data=Melanoma,cause=1)

```

```
# stratified model allowing different baseline risk for the two gender
multi.arr <- ARR(Hist(time,status)~thick+strata(sex)+age+ulcer,data=Melanoma,cause=1)

# stratify by a continuous variable: strata(age)
multi.arr <- ARR(Hist(time,status)~tp(thick,power=0)+strata(age)+sex+ulcer,
                 data=Melanoma,
                 cause=1)

fit.arr2a <- ARR(Hist(time,status)~tp(thick,power=1),data=Melanoma,cause=1)
summary(fit.arr2a)
fit.arr2b <- ARR(Hist(time,status)~timevar(thick),data=Melanoma,cause=1)
summary(fit.arr2b)

## logistic risk model
fit.lrr <- LRR(Hist(time,status)~thick,data=Melanoma,cause=1)
summary(fit.lrr)

## nearest neighbor non-parametric Aalen-Johansen estimate
library(prodlim)
fit.aj <- prodlim(Hist(time,status)~thick,data=Melanoma)
plot(fit.aj,conf.int=FALSE)

# prediction performance
x <- Score(list(fit.arr2a,fit.arr2b,fit.lrr),
           data=Melanoma,
           formula=Hist(time,status)~1,
           cause=1,
           split.method="none")
```

riskRegression.options

Global options for riskRegression

Description

Output and set global options for the riskRegression package.

Usage

```
riskRegression.options(...)
```

Arguments

... for now limited to method.predictRisk and mehtod.predictRiskIID.

Details

only used by the ate function.

Examples

```
options <- riskRegression.options()

## add new method.predictRiskIID
riskRegression.options(method.predictRiskIID = c(options$method.predictRiskIID,"xx"))

riskRegression.options()
```

rowCenter_cpp	<i>Apply - by row</i>
---------------	-----------------------

Description

Fast computation of `sweep(X, MARGIN = 2, FUN = "-", STATS = center)`

Usage

```
rowCenter_cpp(X, center)
```

Arguments

X	A matrix.
center	a numeric vector of length equal to the number of rows of x

Value

A matrix of same size as X.

Author(s)

Brice Ozenne <broz@sund.ku.dk>

Examples

```
x <- matrix(1,6,5)
sweep(x, MARGIN = 2, FUN = "-", STATS = 1:5)
rowCenter_cpp(x, 1:5 )

rowCenter_cpp(x, colMeans(x) )
```

rowCumSum	<i>Apply cumsum in each row</i>
-----------	---------------------------------

Description

Fast computation of `t(apply(x,1,cumsum))`

Usage

```
rowCumSum(x)
```

Arguments

`x` A matrix.

Value

A matrix of same size as `x`.

Author(s)

Thomas Alexander Gerds <tag@biostat.ku.dk>

Examples

```
x <- matrix(1:8,ncol=2)
rowCumSum(x)
```

rowMultiply_cpp	<i>Apply * by row</i>
-----------------	-----------------------

Description

Fast computation of `sweep(X, MARGIN = 2, FUN = "*", STATS = scale)`

Usage

```
rowMultiply_cpp(X, scale)
```

Arguments

`X` A matrix.

`scale` a numeric vector of length equal to the number of rows of `x`

Value

A matrix of same size as `X`.

Author(s)

Brice Ozenne <broz@sund.ku.dk>

Examples

```
x <- matrix(1,6,5)
sweep(x, MARGIN = 2, FUN = "*", STATS = 1:5)
rowMultiply_cpp(x, 1:5 )

rowMultiply_cpp(x, 1/colMeans(x) )
```

rowPaste

Collapse Rows of Characters.

Description

Collapse rows of characters. Fast alternative to `apply(x,1,paste0,collapse="")`

Usage

```
rowPaste(object)
```

Arguments

object A matrix/data.frame/list containing the characters.

Examples

```
## Not run:
M <- matrix(letters,nrow = 26, ncol = 2)
rowPaste(M)

## End(Not run)
```

rowScale_cpp

Apply / by row

Description

Fast computation of `sweep(X, MARGIN = 2, FUN = "/", STATS = scale)`

Usage

```
rowScale_cpp(X, scale)
```

Arguments

`X` A matrix.
`scale` a numeric vector of length equal to the number of rows of `x`

Value

A matrix of same size as `X`.

Author(s)

Brice Ozenne <broz@sund.ku.dk>

Examples

```
x <- matrix(1,6,5)
sweep(x, MARGIN = 2, FUN = "/", STATS = 1:5)
rowScale_cpp(x, 1:5 )

rowScale_cpp(x, colMeans(x) )
```

rowSumsCrossprod *Apply crossprod and rowSums*

Description

Fast computation of `crossprod(rowSums(X),Y)`

Usage

```
rowSumsCrossprod(X, Y, transposeY)
```

Arguments

`X` A matrix with dimensions $n*k$. Hence the result of `rowSums(X)` has length n .
`Y` A matrix with dimenions $n*m$. Can be a matrix with dimension $m*n$ but then `transposeY` should be `TRUE`.
`transposeY` Logical. If `TRUE` transpose `Y` before matrix multiplication.

Value

A vector of length m .

Author(s)

Thomas Alexander Gerds <tag@biostat.ku.dk>

Examples

```
x <- matrix(1:10,nrow=5)
y <- matrix(1:20,ncol=4)
rowSumsCrossprod(x,y,0)
```

```
x <- matrix(1:10,nrow=5)
y <- matrix(1:20,ncol=5)
rowSumsCrossprod(x,y,1)
```

sampleData

Simulate data with binary or time-to-event outcome

Description

Simulate data with binary outcome and 10 covariates.

Usage

```
sampleData(n,outcome="competing.risks",
formula= ~ f(X1,2)+f(X2,-0.033)+f(X3,0.4)+f(X6,.1)+f(X7,-.1)+f(X8,.5)+f(X9,-1),
intercept=0)
sampleDataTD(n,n.intervals=5,outcome="competing.risks",
formula= ~ f(X1,2)+f(X2,-0.033)+f(X3,0.4)+f(X6,.1)+f(X7,-.1)+f(X8,.5)+f(X9,-1))
```

Arguments

n	Sample size
outcome	Character vector. Response variables are generated according to keywords: "binary" = binary response, "survival" = survival response, "competing.risks" = competing risks response
formula	Specify regression coefficients
intercept	For binary outcome the intercept of the logistic regression.
n.intervals	sampleDataTD only: the maximum number of episodes in which the covariates are updated.

Details

For the actual lava::regression parameters see the function definition.

Value

Simulated data as data.table with n rows and the following columns: Y (binary outcome), time (non-binary outcome), event (non-binary outcome), X1-X5 (binary predictors), X6-X10 (continuous predictors)

Author(s)

Thomas A. Gerds <tag@biostat.ku.dk>

See Also

lvm

Examples

```
set.seed(10)
sampleData(10,outcome="binary")
sampleData(10,outcome="survival")
sampleData(10,outcome="competing.risks")
```

saveCoxConfidential *Save confidential Cox objects*

Description

Save confidential Cox objects

Usage

```
saveCoxConfidential(object, times)
```

Arguments

object	An object of class coxph.
times	The times at which we want to predict risk.

Details

This function can save coxph objects such that we do not need to export the data on which it was fitted at given times.

Examples

```
library(survival)
library(lava)
set.seed(18)
trainSurv <- sampleData(300,outcome="survival")
testSurv <- sampleData(40,outcome="survival")
fit = coxph(Surv(time,event)~X1+X2+X3+X7+X9,data=trainSurv, y=TRUE, x = TRUE)
u=saveCoxConfidential(fit,times=3)
## Not run:
# write object as plain text file
sink("~/tmp/u.R")
cat("U <- ")
```

```

dput(u)
sink(NULL)
# reload object
source("~/tmp/u.R")
class(u) <- "CoxConfidential"

## End(Not run)
predictRisk(u,newdata=testSurv)
cox1 = coxph(Surv(time,event)~strata(X1)+X2+X3+X7+X9,data=trainSurv, y=TRUE, x = TRUE)
z<-saveCoxConfidential(cox1,c(2,5))

dput(z) ## get output to copy object

all.equal(predictRisk(z,newdata=testSurv),
           predictRisk(cox1,newdata=testSurv,times=c(2,5)))

cox2 = coxph(Surv(time,event)~X1+X2+X7+X9,data=trainSurv, y=TRUE, x = TRUE)
z<-saveCoxConfidential(cox2,c(2,5))

all.equal(predictRisk(z, testSurv),predictRisk(z, testSurv,c(2,5)))

```

saveSynth

Export a synth object.

Description

Function for exporting objects of type synth via dput.

Usage

```
saveSynth(object, file = "")
```

Arguments

object	An object of type synth.
file	File to print to.

Value

Either a string or nothing if printed to a file.

See Also

lvm

Examples

```
# See the documentation for the \code{synthesize} function.
```

Score	<i>Score risk predictions</i>
-------	-------------------------------

Description

Methods to score the predictive performance of risk markers and risk prediction models

Usage

```
Score(object, ...)
```

```
## S3 method for class 'list'
```

```
Score(  
  object,  
  formula,  
  data,  
  metrics = c("auc", "brier"),  
  summary = NULL,  
  plots = NULL,  
  cause,  
  times,  
  landmarks,  
  use.event.times = FALSE,  
  null.model = TRUE,  
  se.fit = TRUE,  
  conservative = FALSE,  
  conf.int = 0.95,  
  contrasts = TRUE,  
  probs = c(0, 0.25, 0.5, 0.75, 1),  
  cens.method = "ipcw",  
  cens.model = "cox",  
  split.method,  
  B,  
  M,  
  seed,  
  trainseeds,  
  parallel = c("no", "multicore", "snow", "as.registered"),  
  ncpus = 1,  
  cl = NULL,  
  progress.bar = 3,  
  errorhandling = "pass",  
  keep,  
  predictRisk.args,  
  censoring.save.memory = FALSE,  
  breaks = seq(0, 1, 0.01),  
  roc.method = "vertical",  
  roc.grid = switch(roc.method, vertical = seq(0, 1, 0.01), horizontal = seq(1, 0,
```

```

    -0.01)),
  cutpoints = NULL,
  verbose = 2,
  ...
)

```

Arguments

<code>object</code>	List of risk predictions (see details and examples).
<code>...</code>	Named list containing additional arguments that are passed on to the <code>predictRisk</code> methods corresponding to <code>object</code> . See examples.
<code>formula</code>	A formula which identifies the outcome (left hand side). E.g., $Y \sim 1$ for binary and <code>Hist(time, status) ~ 1</code> for time-to-event outcome. In right censored data, the right hand side of the formula is used to estimate the inverse probability of censoring weights (IPCW) model.
<code>data</code>	<code>data.frame</code> or <code>data.table</code> in which the formula can be interpreted.
<code>metrics</code>	Character vector specifying which metrics to apply. Case does not matter. Choices are "AUC" and "Brier".
<code>summary</code>	<p>Character vector specifying which summary statistics to apply to the predicted risks. Choices are "risks", "IPA", "riskQuantile" and "ibs". Can be all <code>c("risks", "IPA", "riskQuantile", "ibs")</code> or a subset thereof.</p> <ul style="list-style-type: none"> • "risks" adds the predicted risks to the output. • "ipa" computes the index of prediction accuracy (AKA R-squared) based on Brier scores for model vs null model • "riskQuantile" calculates time-point specific boxplots for the predicted risks (or biomarker values) conditional on the outcome at the time-point. • "ibs" calculates integrated Brier scores across the time points at which the Brier score is computed. This works only with time-to-event outcome and the results depend on the argument <code>times</code>. <p>Set to <code>NULL</code> to avoid estimation of summary statistics.</p>
<code>plots</code>	Character vector specifying for which plots to put data into the result. Currently implemented are "ROC", "Calibration" and "boxplot". In addition, one can plot AUC and Brier score as function of time as soon as <code>times</code> has at least two different values.
<code>cause</code>	Event of interest. Used for binary outcome Y to specify that risks are risks of the event $Y=\text{event}$ and for competing risks outcome to specify the cause of interest.
<code>times</code>	For survival and competing risks outcome: list of prediction horizons. All times which are greater than the maximal observed time in the data set are automatically removed. Note that the object returned by the function may become huge when the prediction performance is estimated at many prediction horizons.
<code>landmarks</code>	Not yet implemented.
<code>use.event.times</code>	If <code>TRUE</code> merge all unique event times with the vector given by argument <code>times</code> .

<code>null.model</code>	If TRUE fit a risk prediction model which ignores the covariates and predicts the same value for all subjects. The model is fitted using <code>data</code> and the left hand side of <code>formula</code> . For binary outcome this is just the empirical prevalence. For (right censored) time to event outcome, the null models are equal to the Kaplan-Meier estimator (no competing risks) and the Aalen-Johansen estimator (with competing risks).
<code>se.fit</code>	Logical or 0 or 1. If FALSE or 0 do not calculate standard errors.
<code>conservative</code>	Logical, only relevant in right censored data. If TRUE ignore variability of the estimate of the inverse probability of censoring weights when calculating standard errors for prediction performance parameters. This can potentially reduce computation time and memory usage at a usually very small expense of a slightly higher standard error.
<code>conf.int</code>	Either logical or a numeric value between 0 and 1. In right censored data, confidence intervals are based on Blanche et al (see references). Setting FALSE prevents the computation of confidence intervals. TRUE computes 95 percent confidence intervals and corresponding p-values for AUC and Brier score. If set to 0.87, the level of significance is 13 percent. So, do not set it to 0.87.
<code>contrasts</code>	Either logical or a list of contrasts. A list of contrasts defines which risk prediction models (markers) should be contrasted with respect to their prediction performance. If TRUE do all possible comparisons. For example, when object is a list with two risk prediction models and <code>null.model=TRUE</code> setting TRUE is equivalent to <code>list(c(0, 1, 2), c(1, 2))</code> where <code>c(0, 1, 2)</code> codes for the two comparisons: 1 vs 0 and 2 vs 0 (positive integers refer to elements of object, 0 refers to the benchmark null model which ignores the covariates). This again is equivalent to explicitly setting <code>list(c(0, 1), c(0, 2), c(1, 2))</code> . A more complex example: Suppose object has 7 elements and you want to do the following 3 comparisons: 6 vs 3, 2 vs 5 and 2 vs 3, you should set <code>contrasts=c(6, 3), c(2, 5, 3)</code> .
<code>probs</code>	Quantiles for retrospective summary statistics of the predicted risks. This affects the result of the function <code>boxplot.Score</code> .
<code>cens.method</code>	Method for dealing with right censored data. Either "ipcw" or "pseudo". Here IPCW refers to inverse probability of censoring weights and pseudo for jackknife pseudo values. Right now pseudo values are only used for calibration curves.
<code>cens.model</code>	Model for estimating inverse probability of censored weights (IPCW). Implemented are the Kaplan-Meier method ("km") and Cox regression ("cox") both applied to the censored times. If the right hand side of <code>formula</code> does not specify covariates, the Kaplan-Meier method is used even if this argument is set to "cox". Also implemented is a template for users specifying other models to estimate the IPCW. Here the user should be supply a function, taking as input a "formula" and "data". This does come at the cost of only being able to calculate conservative confidence intervals.
<code>split.method</code>	Method for cross-validation. Right now the available choices are <ul style="list-style-type: none"> • "bootcv" bootstrap cross-validation. Argument B controls the number of bootstraps. • "cv5" 5-fold cross-validation. Argument B controls how many times to repeat 5-fold crossvalidation.

	<ul style="list-style-type: none"> • "cv10" 10-fold cross-validation. Argument B controls how many times to repeat 10-fold crossvalidation. • "cvk" k-fold cross-validation for any value of k between 2 and N-1 where N is the sample size. Argument B controls how many times to repeat k-fold crossvalidation. • "1oob" leave-one-out bootstrap cross-validation is performed for the Brier score and leave-pair-out bootstrap cross-validation is performed for the AUC. Argument B controls the number of bootstraps. • "none" no data splitting
B	Number of bootstrap sets for cross-validation. B should be set to 1, when k-fold cross-validation is used.
M	Size of subsamples for bootstrap cross-validation. If specified it has to be an integer smaller than the size of data.
seed	Super seed for setting training data seeds when randomly splitting (bootstrapping) the data during cross-validation.
trainseeds	Seeds for training models during cross-validation.
parallel	The type of parallel operation to be used (if any). If missing, the default is "no".
ncpus	integer: number of processes to be used in parallel operation.
cl	An optional parallel or snow cluster for use if parallel = "snow". If not supplied, a cluster on the local machine is created for the duration of the Score call.
progress.bar	Style for txtProgressBar. Can be 1,2,3 see help(txtProgressBar) or NULL to avoid the progress bar.
errorhandling	Argument passed as .errorhandling to foreach. Default is "pass".
keep	list of characters (not case sensitive) which determines additional output. "residuals" provides Brier score residuals and "splitindex" provides sampling index used to split the data into training and validation sets. It is a function, whose argument is the bootstrap sample, which one wishes to look at. "vcov" provides the variance-covariance matrix for the estimates. "iid" provides the estimated influence function of the estimates.
predictRisk.args	<p>A list of argument-lists to control how risks are predicted. The names of the lists should be the S3-classes of the object. The argument-lists are then passed on to the S3-class specific predictRisk method. For example, if your object contains one or several random forest model fitted with the function randomForestSRC::rfsrc then you can specify additional arguments for the function riskRegression::predictRisk.rfsrc which will pass these on to the function randomForestSRC::predict.rfsrc. A specific example in this case would be list(rfsrc=list(na.action="na.i</p> <p>A more flexible approach is to write a new predictRisk S3-method. See Details.</p>
censoring.save.memory	Only relevant in censored data where censoring weights are obtained with Cox regression and argument conservative is set to FALSE. If TRUE, save memory by not storing the influence function of the cumulative hazard of the censoring as a matrix when calculating standard errors with Cox censoring. This can allow one to use Score on larger test data sets, but may be slower.

breaks	Break points for computing the Roc curve. Defaults to <code>seq(0,1,.01)</code> when some form of crossvalidation is applied, otherwise to all unique values of the predictive marker.
roc.method	Method for averaging ROC curves across data splits. If 'horizontal' average crossvalidated specificities for fixed sensitivity values, specified in <code>roc.grid</code> , otherwise, if 'vertical', average crossvalidated specificities for fixed sensitivity values. See Fawcett, T. (2006) for details.
roc.grid	Grid points for the averaging of ROC curves. A sequence of values at which to compute averages across the ROC curves obtained for different data splits during crossvalidation.
cutpoints	If not NULL, estimates and standard errors of the TPR (True Positive Rate), FPR (False Positive Rate), PPV (Positive Predictive Value), and NPV (Negative Predictive Value) are given at the cutpoints. These values are saved in <code>object\$AUC\$cutpoints</code> .
verbose	Verbosity level. Set to '-1' or 'FALSE' to get complete silence. Set to '0' to quiet all messages and warnings but keep the progress.bar. Set to '1' to see warnings and to '2' (the default) to see both warnings and messages. If 3 or higher, the program produces debug messages.

Details

The function implements a toolbox for the risk prediction modeller: all tools work for the three outcomes: (1) binary (uncensored), (2) right censored time to event without competing risks, (3) right censored time to event with competing risks

Computed are the (time-dependent) Brier score and the (time-dependent) area under the ROC curve for a list of risk prediction models either in external validation data or in the learning data using bootstrap cross-validation. The function optionally provides results for plotting (time-point specific) ROC curves, for (time-point specific) calibration curves and for (time-point specific) retrospective boxplots.

For uncensored binary outcome the DeLong-DeLong test is used to contrast AUC of rival models. In right censored survival data (with and without competing risks) the p-values correspond to Wald tests based on standard errors obtained with an estimate of the influence function as described in detail in the appendix of Blanche et al. (2015).

This function works with one or multiple models that predict the risk of an event $R(t|X)$ for a subject characterized by predictors X at time t . With binary endpoints (outcome 0/1 without time component) the risk is simply $R(X)$. In case of a survival object without competing risks the function still works with predicted event probabilities, i.e., $R(t|X)=1-S(t|X)$ where $S(t|X)$ is the predicted survival chance for subject X at time t .

The already existing `predictRisk` methods (see `methods(predictRisk)`) may not cover all models and methods for predicting risks. But users can quickly extend the package as explained in detail in Mogensen et al. (2012) for the predecessors `pec::predictSurvProb` and `pec::predictEventProb` which have been unified as `riskRegression::predictRisk`.

Bootstrap Crossvalidation (see also Gerds & Schumacher 2007 and Mogensen et al. 2012)

$B=10$, M (not specified or $M=NROW(data)$) Training of each of the models in each of 10 bootstrap data sets (learning data sets). Learning data sets are obtained by sampling $NROW(data)$ subjects of

the data set with replacement. There are roughly $.632 \times \text{NROW}(\text{data})$ subjects in the learning data (inbag) and $.368 \times \text{NROW}(\text{data})$ subjects not in the validation data sets (out-of-bag).

These are used to estimate the scores: AUC, Brier, etc. Reported are averages across the 10 splits.

```
## Bootstrap with replacement set.seed(13) N=17 data = data.frame(id=1:N, y=rbinom(N,1,.3), x=rnorm(N))
boot.index = sample(1:N, size=N, replace=TRUE) boot.index inbag = 1:N outofbag = !inbag
learn.data = data[inbag] val.data = data[outofbag] riskRegression::getSplitMethod("bootcv", B=10, N=17)
NOTE: the number .632 is the expected probability to draw one subject (for example subject
1) with replacement from the data, which does not depend on the sample size: B=10000 N=137
mean(sapply(1:B, function(b){match(1, sample(1:N, size=N, replace=TRUE), nomatch=0)}))
N=30 mean(sapply(1:B, function(b){match(1, sample(1:N, size=N, replace=TRUE), nomatch=0)}))
N=300 mean(sapply(1:B, function(b){match(1, sample(1:N, size=N, replace=TRUE), nomatch=0)}))

## Bootstrap without replacement (training size set to be 70 percent of data) B=10, M=.7
```

Training of each of the models in each of 10 bootstrap data sets (learning data sets). Learning data sets are obtained by sampling $\text{round}(.8 \times \text{NROW}(\text{data}))$ subjects of the data set without replacement. There are $\text{NROW}(\text{data}) - \text{round}(.8 \times \text{NROW}(\text{data}))$ subjects not in the learning data sets. These are used to estimate the scores: AUC, Brier, etc. Reported are averages across the 10 splits. `set.seed(13) N=17 data = data.frame(id=1:N, y=rbinom(N,1,.3), x=rnorm(N))`
`boot.index = sample(1:N, size=M, replace=FALSE) boot.index inbag = 1:N outofbag = !inbag`
`learn.data = data[inbag] val.data = data[outofbag] riskRegression::getSplitMethod("bootcv", B=10, N=17,`

Value

List with scores and assessments of contrasts, i.e., tests and confidence limits for performance and difference in performance (AUC and Brier), summaries and plots. Most elements are in `data.table` format.

Author(s)

Thomas A Gerds <tag@biostat.ku.dk> and Paul Blanche <paul.blanche@univ-ubs.fr>

References

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Thomas A. Gerds, Martin Schumacher (2007) Efron-Type Measures of Prediction Error for Survival Analysis *Biometrics*, 63(4), 1283–1287 doi:10.1111/j.1541-0420.2007.00832.x

Martin Schumacher, Harald Binder, and Thomas Gerds. Assessment of survival prediction models based on microarray data. *Bioinformatics*, 23(14):1768-74, 2007.

Mark A. van de Wiel, Johannes Berkhof, and Wessel N. van Wieringen Testing the prediction error difference between 2 predictors *Biostatistics* (2009) 10(3): 550-560 doi:10.1093/biostatistics/kxp011

Michael W Kattan and Thomas A Gerds. The index of prediction accuracy: an intuitive measure useful for evaluating risk prediction models. *Diagnostic and Prognostic Research*, 2(1):7, 2018.

Fawcett, T. (2006). An introduction to ROC analysis. *Pattern Recognition Letters*, 27, 861-874.

Examples

```
# binary outcome
library(lava)
set.seed(18)
learndat <- sampleData(48,outcome="binary")
testdat <- sampleData(40,outcome="binary")

## score logistic regression models
lr1 = glm(Y~X1+X2+X7+X9,data=learndat,family=binomial)
lr2 = glm(Y~X3+X5,data=learndat,family=binomial)
x=Score(list("LR(X1+X2+X7+X9)"=lr1,"LR(X3+X5)"=lr2),formula=Y~1,data=testdat)
print(x)
summary(x)

## ROC curve and calibration plot
xb=Score(list("LR(X1+X2+X7+X9)"=lr1,"LR(X3+X5+X6)"=lr2),formula=Y~1,
          data=testdat,plots=c("calibration","ROC"))
## Not run: plotROC(xb)
plotCalibration(xb)

## End(Not run)

## compute AUC for a list of continuous markers
markers = as.list(testdat[,.(X6,X7,X8,X9,X10)])
Score(markers,formula=Y~1,data=testdat,metrics=c("auc"))

# cross-validation
## Not run:
  set.seed(10)
  learndat=sampleData(400,outcome="binary")
  lr1a = glm(Y~X6,data=learndat,family=binomial)
  lr2a = glm(Y~X7+X8+X9,data=learndat,family=binomial)
  ## bootstrap cross-validation
  x1=Score(list("LR1"=lr1a,"LR2"=lr2a),formula=Y~1,data=learndat,split.method="bootcv",B=100)
  x1
```

```

## leave-one-out and leave-pair-out bootstrap
x2=Score(list("LR1"=lr1a,"LR2"=lr2a),formula=Y~1,data=learndat,
          split.method="loob",
          B=100,plots="calibration")

x2
## 5-fold cross-validation
x3=Score(list("LR1"=lr1a,"LR2"=lr2a),formula=Y~1,data=learndat,
          split.method="cv5",
          B=1,plots="calibration")

x3

## End(Not run)
# survival outcome

# Score Cox regression models
## Not run: library(survival)
library(rms)
library(proplim)
set.seed(18)
trainSurv <- sampleData(100,outcome="survival")
testSurv <- sampleData(40,outcome="survival")
cox1 = coxph(Surv(time,event)~X1+X2+X7+X9,data=trainSurv, y=TRUE, x = TRUE)
cox2 = coxph(Surv(time,event)~X3+X5+X6,data=trainSurv, y=TRUE, x = TRUE)
x=Score(list("Cox(X1+X2+X7+X9)"=cox1,"Cox(X3+X5+X6)"=cox2),
         formula=Surv(time,event)~1,data=testSurv,conf.int=FALSE,times=c(5,8))
## Use Cox to estimate censoring weights
y=Score(list("Cox(X1+X2+X7+X9)"=cox1,"Cox(X3+X5+X6)"=cox2),
         formula=Surv(time,event)~X1+X8,data=testSurv,conf.int=FALSE,times=c(5,8))
## Use GLMnet to estimate censoring weights
z=Score(list("Cox(X1+X2+X7+X9)"=cox1,"Cox(X3+X5+X6)"=cox2),
         formula=Surv(time,event)~X1+X8,cens.model = "GLMnet",data=testSurv,
         conf.int=FALSE,times=c(5,8))
## Use hal9001 to estimate censoring weights
w=Score(list("Cox(X1+X2+X7+X9)"=cox1,"Cox(X3+X5+X6)"=cox2),
         formula=Surv(time,event)~X1+X8,cens.model = "Hal9001",data=testSurv,
         conf.int=FALSE,times=c(5,8))

x
y
z
w

## End(Not run)

## Not run: library(survival)
library(rms)
library(proplim)
set.seed(18)
trainSurv <- sampleData(100,outcome="survival")
testSurv <- sampleData(40,outcome="survival")
cox1 = coxph(Surv(time,event)~X1+X2+X7+X9,data=trainSurv, y=TRUE, x = TRUE)
cox2 = coxph(Surv(time,event)~X3+X5+X6,data=trainSurv, y=TRUE, x = TRUE)
xs=Score(list("Cox(X1+X2+X7+X9)"=cox1,"Cox(X3+X5+X6)"=cox2),
          formula=Surv(time,event)~1,data=testSurv,conf.int=FALSE,times=c(5,8))

```

```

xs

## End(Not run)
# Integrated Brier score
## Not run:
xs=Score(list("Cox(X1+X2+X7+X9)"=cox1,"Cox(X3+X5+X6)"=cox2),
          formula=Surv(time,event)~1,data=testSurv,conf.int=FALSE,
          summary="ibs",
          times=sort(unique(testSurv$time)))

## End(Not run)

# time-dependent AUC for list of markers
## Not run: survmarkers = as.list(testSurv[,.(X6,X7,X8,X9,X10)])
Score(survmarkers,
      formula=Surv(time,event)~1,metrics="auc",data=testSurv,
      conf.int=TRUE,times=c(5,8))

# compare models on test data
Score(list("Cox(X1+X2+X7+X9)"=cox1,"Cox(X3+X5+X6)"=cox2),
      formula=Surv(time,event)~1,data=testSurv,conf.int=TRUE,times=c(5,8))

## End(Not run)
# crossvalidation models in traindata
## Not run:
library(survival)
set.seed(18)
trainSurv <- sampleData(400,outcome="survival")
cox1 = coxph(Surv(time,event)~X1+X2+X7+X9,data=trainSurv, y=TRUE, x = TRUE)
cox2 = coxph(Surv(time,event)~X3+X5+X6,data=trainSurv, y=TRUE, x = TRUE)
x1 = Score(list("Cox(X1+X2+X7+X9)"=cox1,"Cox(X3+X5+X6)"=cox2),
           formula=Surv(time,event)~1,data=trainSurv,conf.int=TRUE,times=c(5,8),
           split.method="loob",B=100,plots="calibration")

x2= Score(list("Cox(X1+X2+X7+X9)"=cox1,"Cox(X3+X5+X6)"=cox2),
          formula=Surv(time,event)~1,data=trainSurv,conf.int=TRUE,times=c(5,8),
          split.method="bootcv",B=100)

## End(Not run)

# restrict number of comparisons
## Not run:
Score(list("Cox(X1+X2+X7+X9)"=cox1,"Cox(X3+X5+X6)"=cox2),
      formula=Surv(time,event)~1,data=trainSurv,contrasts=TRUE,
      null.model=FALSE,conf.int=TRUE,times=c(5,8),split.method="bootcv",B=3)

# competing risks outcome
set.seed(18)
trainCR <- sampleData(400,outcome="competing.risks")
testCR <- sampleData(400,outcome="competing.risks")
library(riskRegression)
library(cmprsk)
# Cause-specific Cox regression

```

```

csc1 = CSC(Hist(time,event)~X1+X2+X7+X9,data=trainCR)
csc2 = CSC(Hist(time,event)~X3+X5+X6,data=trainCR)
# Fine-Gray regression
fgr1 = FGR(Hist(time,event)~X1+X2+X7+X9,data=trainCR,cause=1)
fgr2 = FGR(Hist(time,event)~X3+X5+X6,data=trainCR,cause=1)
Score(list("CSC(X1+X2+X7+X9)"=csc1,"CSC(X3+X5+X6)"=csc2,
          "FGR(X1+X2+X7+X9)"=fgr1,"FGR(X3+X5+X6)"=fgr2),
       formula=Hist(time,event)~1,data=testCR,se.fit=1L,times=c(5,8))

## End(Not run)

## Not run:
# reproduce some results of Table IV of Blanche et al. Stat Med 2013
data(Paquid)
ResPaquid <- Score(list("DSST"--Paquid$DSST,"MMSE"--Paquid$MMSE),
                  formula=Hist(time,status)~1,
                  data=Paquid,
                  null.model = FALSE,
                  conf.int=TRUE,
                  metrics=c("auc"),
                  times=c(3,5,10),
                  plots="ROC")

ResPaquid
plotROC(ResPaquid,time=5)

## End(Not run)
## Not run:
# parallel options
# by erikvona: Here is a generic example of using future
# and doFuture, works great with the current version:
library(riskRegression)
library(future)
library(foreach)
library(doFuture)
library(survival)
# Register all available cores for parallel operation
plan(multiprocess, workers = availableCores())
registerDoFuture()
set.seed(10)
trainSurv <- sampleData(400,outcome="survival")
cox1 = coxph(Surv(time,event)~X1+X2+X7+X9,data=trainSurv,
             y=TRUE, x = TRUE)
# Bootstrapping on multiple cores
x1 = Score(list("Cox(X1+X2+X7+X9)"=cox1),
           formula=Surv(time,event)~1,data=trainSurv, times=c(5,8),
           parallel = "as.registered", split.method="bootcv",B=100)

## End(Not run)

```

score.wglm	<i>Score for IPCW Logistic Regressions</i>
------------	--

Description

Compute the first derivative of the log-likelihood for IPCW logistic regressions.

Usage

```
## S3 method for class 'wglm'
score(x, indiv = FALSE, times = NULL, simplify = TRUE, ...)
```

Arguments

x	a wglm object.
indiv	[logical] should the individual score be output? Otherwise the total score (i.e. summed over all individuals will be output).
times	[numeric vector] time points at which the score should be output.
simplify	[logical] should the output be converted to a matrix when only one timepoint is requested. Otherwise will always return a list.
...	Not used.

selectCox	<i>Backward variable selection in the Cox regression model</i>
-----------	--

Description

This is a wrapper function which first selects variables in the Cox regression model using `fastbw` from the `rms` package and then returns a fitted Cox regression model with the selected variables.

Usage

```
selectCox(formula, data, rule = "aic")
```

Arguments

formula	A formula object with a <code>Surv</code> object on the left-hand side and all the variables on the right-hand side.
data	Name of a data frame containing all needed variables.
rule	The method for selecting variables. See fastbw for details.

Details

This function first calls `cph` then `fastbw` and finally `cph` again.

References

Ulla B. Mogensen, Hemant Ishwaran, Thomas A. Gerds (2012). Evaluating Random Forests for Survival Analysis Using Prediction Error Curves. *Journal of Statistical Software*, 50(11), 1-23. URL <http://www.jstatsoft.org/v50/i11/>.

Examples

```
library(survival)
set.seed(74)
d <- sampleData(89, outcome="survival")
f <- selectCox(Surv(time, event)~X1+X2+X3+X4+X6+X7+X8+X9, data=d)
```

selectJump	<i>Evaluate the influence function at selected times</i>
------------	--

Description

Evaluate the influence function at selected times

Usage

```
selectJump(IF, times, type)
```

Arguments

IF	influence function returned by iidCox
times	the times at which the influence function should be assessed
type	can be "hazard" or/and "cumhazard".

Value

An object with the same dimensions as IF

Author(s)

Brice Ozenne broz@sund.ku.dk

simActiveSurveillance *Simulate data of a hypothetical active surveillance prostate cancer study*

Description

Simulate data of a hypothetical active surveillance prostate cancer study

Usage

```
simActiveSurveillance(n)
```

Arguments

n sample size

Details

This is based on the functionality of `library(lava)`.

Value

data table of size n

Author(s)

Thomas A. Gerds <tag@biostat.ku.dk>

Examples

```
set.seed(71)
simActiveSurveillance(3)
```

simMelanoma *Simulate data alike the Melanoma data*

Description

Simulate data alike the Melanoma data

Usage

```
simMelanoma(n)
```

Arguments

n sample size

Details

This is based on the functionality of `library(lava)`.

Value

data table of size n

Author(s)

Thomas A. Gerds <tag@biostat.ku.dk>

Examples

```
set.seed(71)
simMelanoma(3)
```

simPBC

simulating data alike the pbc data

Description

This function can be used to simulate data alike the pbc data from the survival package.

Usage

```
simPBC(n)
```

Arguments

n Sample size

Details

using lava to synthesize data

Value

The simulated data.

Author(s)

Thomas A. Gerds <tag@biostat.ku.dk>

Examples

```

library(survival)
library(lava)
# simulate data alike pbc data
set.seed(98)
d=simPBC(847)
d$protimegrp1 <- d$protimegrp=="10-11"
d$protimegrp2 <- d$protimegrp==">11"
d$sex <- factor(d$sex,levels=0:1,labels=c("m","f"))
sF1 <- survreg(Surv(time,status==1)~sex+age+logbili+protimegrp1+protimegrp2+stage3+stage4,
data=d)
coxF1 <- coxph(Surv(time,status==1)~sex+age+logbili+protimegrp1+protimegrp2+stage3+stage4,
data=d)
# load real pbc data
data(pbc,package="survival")
pbc <- na.omit(pbc[,c("time","status","age","sex","stage","bili","protime","trt")])
pbc$stage <- factor(pbc$stage)
levels(pbc$stage) <- list("1/2"=c(1,2),"3"=3,"4"=4)
pbc$logbili <- log(pbc$bili)
pbc$logprotime <- log(pbc$protime)
pbc$stage3 <- 1*(pbc$stage=="3")
pbc$stage4 <- 1*(pbc$stage=="4")
pbc$protimegrp <- cut(pbc$protime,c(-Inf,10,11,Inf),labels=c("<=10","10-11",">11"))
pbc$protimegrp1 <- pbc$protimegrp=="10-11"
pbc$protimegrp2 <- pbc$protimegrp==">11"
form1=Surv(time,status==1)~sex+age+logbili+protimegrp1+protimegrp2+stage3+stage4
F1 <- survival::survreg(form1,data=pbc)
form2=Surv(time,status==2)~sex+age+logbili+protimegrp1+protimegrp2+stage3+stage4
F2 <- survival::survreg(form1,data=pbc)
sF2 <- survreg(Surv(time,status==2)~sex+age+logbili+protimegrp1+protimegrp2+stage3+stage4,
data=d)
G <- survreg(Surv(time,status==0)~sex+age+logbili+protimegrp1+protimegrp2+stage3+stage4,
data=pbc)
sG <- survreg(Surv(time,status==0)~sex+age+logbili+protimegrp1+protimegrp2+stage3+stage4,
data=d)
# compare fits in real and simulated pbc data
cbind(coef(F1),coef(sF1))
cbind(coef(F2),coef(sF2))
cbind(coef(G),coef(sG))
cbind(coef(glm(protimegrp1~age+sex+logbili,data=pbc,family="binomial")),
coef(glm(protimegrp1~age+sex+logbili,data=d,family="binomial")))
cbind(coef(lm(logbili~age+sex,data=pbc)),coef(lm(logbili~age+sex,data=d)))

```

Description

Simulating from a synthesized object

Usage

```
simsynth(object, n = 200, drop.latent = FALSE, ...)
```

Arguments

object	generated with <code>synthesize</code>
n	sample size
drop.latent	if TRUE remove the latent event times from the resulting data set.
...	additional arguments passed on to <code>lava::sim</code>

Examples

```
library(survival)
m=synthesize(Surv(time,status)~sex+age+bili,data=pbcc)
simsynth(m,10,drop.latent=TRUE)
```

SmcFcs

SmcFcs

Description

TODO

Usage

```
SmcFcs(formula, data, m = 5, method, fitter = "glm", fit.formula, ...)
```

Arguments

formula	TODO
data	TODO
m	TODO
method	TODO
fitter	TODO
fit.formula	TODO
...	TODO
	# @export

splitStrataVar	<i>Reconstruct each of the strata variables</i>
----------------	---

Description

Reconstruct each of the strata variables from the strata variable stored in the coxph object.

Usage

```
splitStrataVar(object)
```

Arguments

object a coxph object.

Author(s)

Brice Ozenne broz@sund.ku.dk and Thomas A. Gerds tag@biostat.ku.dk

subjectWeights	<i>Estimation of censoring probabilities at subject specific times</i>
----------------	--

Description

This function is used internally to construct pseudo values by inverse of the probability of censoring weights.

Usage

```
subjectWeights(
  formula,
  data,
  method = c("cox", "marginal", "km", "nonpar", "forest", "none"),
  args,
  lag = 1
)
```

Arguments

formula	A survival formula like, Surv(time,status)~1 or Hist(time,status)~1 where status=0 means censored. The status variable is internally reversed for estimation of censoring rather than survival probabilities. Some of the available models, see argument model, will use predictors on the right hand side of the formula.
data	The data used for fitting the censoring model
method	Censoring model used for estimation of the (conditional) censoring distribution.

args	Arguments passed to the fitter of the method.
lag	If equal to 1 then obtain $G(T_i - X_i)$, if equal to 0 estimate the conditional censoring distribution at the subject.times, i.e. $(G(T_i X_i))$.

Details

Inverse of the probability of censoring weights usually refer to the probabilities of not being censored at certain time points. These probabilities are also the values of the conditional survival function of the censoring time given covariates. The function `subjectWeights` estimates the conditional survival function of the censoring times and derives the weights.

IMPORTANT: the data set should be ordered, `order(time, -status)` in order to get the weights in the right order for some choices of method.

Value

times	The times at which weights are estimated
weights	Estimated weights at individual time values <code>subject.times</code>
lag	The time lag.
fit	The fitted censoring model
method	The method for modelling the censoring distribution
call	The call

Author(s)

Thomas A. Gerds <tag@biostat.ku.dk>

Examples

```
library(prodlm)
library(survival)
dat=SimSurv(300)

dat <- dat[order(dat$time,-dat$status),]

# using the marginal Kaplan-Meier for the censoring times

WKM=subjectWeights(Hist(time,status)~X2,data=dat,method="marginal")
plot(WKM$fit)
WKM$fit
WKM$weights

# using the Cox model for the censoring times given X2

WCox=subjectWeights(Surv(time,status)~X2,data=dat,method="cox")
WCox
plot(WCox$weights,WKM$weights)

# using the stratified Kaplan-Meier for the censoring times given X2
```



```
WKM2 <- subjectWeights(Surv(time,status)~X2,data=dat,method="nonpar")
plot(WKM2$fit,add=FALSE)
```

subsetIndex

Extract Specific Elements From An Object

Description

Extract specific elements from an object.

Usage

```
subsetIndex(object, index, default, ...)

## Default S3 method:
subsetIndex(object, index, default, ...)

## S3 method for class 'matrix'
subsetIndex(object, index, default, col = TRUE, ...)
```

Arguments

object	A vector or a matrix.
index	index of the elements to be extracted. 0 indicates that the column should be set to the default value. NA indicates that the column should be set to NA.
default	the default value.
...	Only used by the generic method.
col	If object is a matrix, TRUE lead to extract the columns and FALSE the rows.

Examples

```
M <- matrix(rnorm(50),5,10)
subsetIndex(M, index = c(0,0,1), default = 0)
subsetIndex(M, index = c(0,2,3,NA), default = 0)
subsetIndex(M, index = c(0,NA,2,3,NA), default = 0)

C <- 1:10
subsetIndex(C, index = c(0,0,1,5,NA), default = 0)
```

summary.ate

*Summary Average Treatment Effects***Description**

Summary average treatment effects.

Usage

```
## S3 method for class 'ate'
summary(
  object,
  estimator = object$estimator[1],
  short = FALSE,
  type = c("meanRisk", "diffRisk"),
  se = FALSE,
  quantile = FALSE,
  estimate.boot = TRUE,
  digits = 3,
  ...
)
```

Arguments

object	object obtained with function ate
estimator	[character] The type of estimator relative to which the estimates should be displayed.
short	[logical] If TRUE, only displays the estimated risks.
type	[character vector] what to displayed. Can be "meanRisk" to display the risks specific to each treatment group, "diffRisk" to display the difference in risks between treatment groups, or "ratioRisk" to display the ratio of risks between treatment groups,.
se	[logical] should the standard error of the risks be displayed?
quantile	[logical] should the quantile of the confidence bands be displayed?
estimate.boot	[logical] should the average estimate on the bootstrap samples be displayed?
digits	[integer, >0] Number of digits.
...	passed to confint

Details

to display confidence intervals/bands and p.value, the confint method needs to be applied on the object.

See Also

[as.data.table](#) to extract the estimates in a `data.table` object. [autoplot.ate](#) for a graphical representation the standardized risks. [confint.ate](#) to compute p-values and adjusted p-values or perform statistical inference using a transformation. [confint.ate](#) to compute (pointwise/simultaneous) confidence intervals and (unadjusted/adjusted) p-values, possibly using a transformation.

summary.FGR

Summary of a Fine-Gray regression model

Description

Summary of a Fine-Gray regression model

Usage

```
## S3 method for class 'FGR'
summary(object, ...)
```

Arguments

object	Object fitted with function FGR
...	passed to <code>cmprsk::summary.crr</code>

summary.riskRegression

Summary of a risk regression model

Description

Summary of a risk regression model

Usage

```
## S3 method for class 'riskRegression'
summary(
  object,
  times,
  digits = 3,
  pvalue.digits = 4,
  eps = 10^-4,
  verbose = TRUE,
  ...
)
```

Arguments

object	Object obtained with ARR, LRR or riskRegression
times	Time points at which to show time-dependent coefficients
digits	Number of digits for all numbers but p-values
pvalue.digits	Number of digits for p-values
eps	p-values smaller than this number are shown as such
verbose	Level of verbosity
...	not used

summary.Score	<i>Summary of prediction performance metrics</i>
---------------	--

Description

Summarizing a Score object

Usage

```
## S3 method for class 'Score'
summary(
  object,
  times,
  what = c("score", "contrasts"),
  models,
  digits = 1,
  pvalue.digits = 4,
  ...
)
```

Arguments

object	Object obtained with Score.
times	Select time points
what	Either "score", "contrasts" or both, i.e., c("score", "contrasts")
models	Select which models to summarize. Need to be a subset of object\$models
digits	For rounding everything but p-values
pvalue.digits	For rounding p-values
...	not used

Details

The AUC and the Brier score are put into tables

Value

List of tables

Author(s)

Thomas A. Gerds <tag@biostat.ku.dk>

See Also

Score

SuperPredictor *Formula interface for SuperLearner::SuperLearner*

Description

Formula interface for SuperLearner::SuperLearner

Usage

```
SuperPredictor(
  formula,
  data,
  family = "binomial",
  SL.library = c("SL.glm", "SL.glm.interaction", "SL.ranger"),
  ...
)
```

Arguments

formula	where the left hand side specifies the outcome and the right hand side the predictors
data	data set in which formula can be evaluated
family	the outcome family. default is binomial
SL.library	the SuperLearner libraries
...	passed to SuperLearner::SuperLearner

Details

Formula interface for SuperLearner::SuperLearner ##' @param formula

Examples

```
## Not run:
if(require("SuperLearner",quietly=TRUE)){
  library(SuperLearner)
  library(data.table)
  set.seed(10)
  d = sampleData(338, outcome="binary")
  spfit = SuperPredictor(Y~X1+X2+X3+X4+X5+X6+X7+X8+X9+X10,data=d)
  predictRisk(spfit)
  x <- Score(list(spfit),data=d,formula=Y~1)
}

## End(Not run)
```

SurvResponseVar

Extract the time and event variable from a Cox model

Description

Extract the time and event variable from a Cox model

Usage

```
SurvResponseVar(formula)
```

Arguments

```
formula      a formula
```

Author(s)

```
Brice Ozenne broz@sund.ku.dk
```

Examples

```
## Not run:
SurvResponseVar(Surv(time,event)~X1+X2)
SurvResponseVar(Hist(time,event==0)~X1+X2)
SurvResponseVar(Surv(start,time, status,type="counting") ~ X3+X5)
SurvResponseVar(Surv(start,event=status, time2=time,type="counting") ~ X3+X5)

SurvResponseVar(survival::Surv(start,event=status, time2=time,type="counting") ~ X3+X5)
SurvResponseVar(status ~ X3+X5)
SurvResponseVar(I(status == 1) ~ X3+X5)
SurvResponseVar(list(Hist(time, event) ~ X1+X6,Hist(time, event) ~ X6))

## End(Not run)
```

 synthesize

Cooking and synthesizing survival data

Description

Fit parametric regression models to the outcome distribution and optionally also parametric regression models for the joint distribution of the predictors structural equation models. Then the function `sim.synth` can be called on the resulting object to simulate from the parametric model based on the machinery of the `lava` package

Usage

```

synthesize(object, data, ...)

## S3 method for class 'formula'
synthesize(
  object,
  data,
  recursive = FALSE,
  max.levels = 10,
  verbose = FALSE,
  ...
)

## S3 method for class 'lvm'
synthesize(
  object,
  data,
  max.levels = 10,
  logtrans = NULL,
  verbose = FALSE,
  fix.names = FALSE,
  ...
)

```

Arguments

<code>object</code>	Specification of the synthesizing model structures. Either a formula or a lvm object. See examples.
<code>data</code>	Data to be synthesized.
<code>...</code>	Not used yet.
<code>recursive</code>	Let covariates recursively depend on each other.
<code>max.levels</code>	Integer used to guess which variables are categorical. When set to 10, the default, variables with less than 10 unique values in data are treated as categorical.
<code>verbose</code>	Logical. If TRUE then more messages and warnings are provided.

logtrans	Vector of covariate names that should be log-transformed. This is primarily for internal use.
fix.names	Fix possible problematic covariate names.

Details

Synthesizes survival data (also works for linear models and generalized linear models). The idea is to be able to simulate new data sets that mimic the original data. See the vignette `vignette("synthesize", package = "riskRegression")` for more details.

The simulation engine is: lava.

Value

lava object

Author(s)

Thomas A. Gerds <tag@biostat.ku.dk>

See Also

lvm

Examples

```
# pbc data
library(survival)
library(lava)
data(pbc)
pbc <- na.omit(pbc[,c("time", "status", "sex", "age", "bili")])
pbc$logbili <- log(pbc$bili)
v_synt <- synthesize(object=Surv(time, status)~sex+age+logbili, data=pbc)
d <- simsynth(v_synt, 1000)
fit_sim <- coxph(Surv(time, status==1)~age+sex+logbili, data=d)
fit_real <- coxph(Surv(time, status==1)~age+sex+logbili, data=pbc)
# compare estimated log-hazard ratios between simulated and real data
cbind(coef(fit_sim), coef(fit_real))

u <- lvm()
distribution(u, ~sex) <- binomial.lvm()
distribution(u, ~age) <- normal.lvm()
distribution(u, ~trt) <- binomial.lvm()
distribution(u, ~logbili) <- normal.lvm()
u <- eventTime(u, time~min(time.cens=0, time.transplant=1, time.death=2), "status")
lava::regression(u, logbili~age+sex) <- 1
lava::regression(u, time.transplant~sex+age+logbili) <- 1
lava::regression(u, time.death~sex+age+logbili) <- 1
lava::regression(u, time.cens~1) <- 1
transform(u, logbili~bili) <- function(x){log(x)}
u_synt <- synthesize(object=u, data=na.omit(pbc))
saveSynth(u_synt)
```



```

set.seed(8)
d <- simsynth(u_synt,n=1000)
# note: synthesizer may relabel status variable
fit_sim <- coxph(Surv(time,status==1)~age+sex+logbili,data=d)
fit_real <- coxph(Surv(time,status==1)~age+sex+log(bili),data=pbcc)
# compare estimated log-hazard ratios between simulated and real data
cbind(coef(fit_sim),coef(fit_real))

#
# Cancer data
#
data(cancer)
b <- lvm()
distribution(b,~rx) <- binomial.lvm()
distribution(b,~age) <- normal.lvm()
distribution(b,~resid.ds) <- binomial.lvm()
distribution(b,~ecog.ps) <- binomial.lvm()
lava::regression(b,time.death~age+rx+resid.ds) <- 1
b<-eventTime(b,futime~min(time.cens=0,time.death=1), "fustat")
b_synt <- synthesizer(object = b, data = ovarian)
D <- simsynth(b_synt,1000)
fit_real <- coxph(Surv(futime,fustat)~age+rx+resid.ds, data=ovarian)
fit_sim <- coxph(Surv(futime,fustat)~age+rx+resid.ds, data=D)
cbind(coef(fit_sim),coef(fit_real))
w_synt <- synthesizer(object=Surv(futime,fustat)~age+rx+resid.ds, data=ovarian)
D <- simsynth(w_synt,1000)
fit_sim <- coxph(Surv(futime,fustat==1)~age+rx+resid.ds,data=D)
fit_real <- coxph(Surv(futime,fustat==1)~age+rx+resid.ds,data=ovarian)
# compare estimated log-hazard ratios between simulated and real data
cbind(coef(fit_sim),coef(fit_real))

```

terms.phreg

Extract terms for phreg objects

Description

Extract terms for phreg objects

Usage

```
## S3 method for class 'phreg'
terms(x, ...)
```

Arguments

x	a phreg object.
...	not used.

transformCIBP	<i>Compute Confidence Intervals/Bands and P-values After a Transformation</i>
---------------	---

Description

Compute confidence intervals/bands and p-values after a transformation

Usage

```
transformCIBP(
  estimate,
  se,
  iid,
  null,
  conf.level,
  alternative,
  ci,
  type,
  min.value,
  max.value,
  band,
  method.band,
  n.sim,
  seed,
  p.value,
  df = NULL
)
```

Arguments

estimate	[numeric matrix] the estimate value before transformation.
se	[numeric matrix] the standard error before transformation.
iid	[numeric array] the iid decomposition before transformation.
null	[numeric] the value of the estimate (before transformation) under the null hypothesis.
conf.level	[numeric, 0-1] Level of confidence.
alternative	[character] a character string specifying the alternative hypothesis, must be one of "two.sided" (default), "greater" or "less".
ci	[logical] should confidence intervals be computed.
type	[character] the transformation. Can be "log", "loglog", "cloglog", or "atanh" (Fisher transform), or "atanh2" (modified Fisher transform for [0-1] variable).
min.value	[numeric] if not NULL and the lower bound of the confidence interval is below min, it will be set at min.

max.value	[numeric] if not NULL and the lower bound of the confidence interval is below max, it will be set at max.
band	[integer 0,1,2] When non-0, the confidence bands are computed for each contrasts (band=1) or over all contrasts (band=2).
method.band	[character] method used to adjust for multiple comparisons. Can be any element of p.adjust.methods (e.g. "holm"), "maxT-integration", or "maxT-simulation".
n.sim	[integer, >0] the number of simulations used to compute the quantiles for the confidence bands.
seed	[integer, >0] seed number set before performing simulations for the confidence bands.
p.value	[logical] should p-values and adjusted p-values be computed. Only active if ci=TRUE or band>0.
df	[integer, >0] optional. Degrees of freedom used for the student distribution of the test statistic. If not specified, use a normal distribution instead.

Details

The iid decomposition must have dimensions [n.obs,time,n.prediction] while estimate and se must have dimensions [n.prediction,time].

Single step max adjustment for multiple comparisons, i.e. accounting for the correlation between the test statistics but not for the ordering of the tests, can be performed setting the arguemnt method.band to "maxT-integration" or "maxT-simulation". The former uses numerical integration (pmvnorm and qmvnorm to perform the adjustment while the latter using simulation. Both assume that the test statistics are jointly normally distributed.

Examples

```
set.seed(10)
n <- 100
X <- rnorm(n)

res2sided <- transformCIBP(estimate = mean(X), se = cbind(sd(X)/sqrt(n)), null = 0,
  type = "none", ci = TRUE, conf.level = 0.95, alternative = "two.sided",
  min.value = NULL, max.value = NULL, band = FALSE,
  p.value = TRUE, seed = 10, df = n-1)

resLess <- transformCIBP(estimate = mean(X), se = cbind(sd(X)/sqrt(n)), null = 0,
  type = "none", ci = TRUE, conf.level = 0.95, alternative = "less",
  min.value = NULL, max.value = NULL, band = FALSE,
  p.value = TRUE, seed = 10, df = n-1)

resGreater <- transformCIBP(estimate = mean(X), se = cbind(sd(X)/sqrt(n)), null = 0,
  type = "none", ci = TRUE, conf.level = 0.95, alternative = "greater",
  min.value = NULL, max.value = NULL, band = FALSE,
  p.value = TRUE, seed = 10, df = n-1)

## comparison with t-test
GS <- t.test(X, alternative = "two.sided")
```

```

res2sided$p.value - GS$p.value
unlist(res2sided[c("lower", "upper")]) - GS$conf.int

GS <- t.test(X, alternative = "less")
resLess$p.value - GS$p.value
unlist(resLess[c("lower", "upper")]) - GS$conf.int

GS <- t.test(X, alternative = "greater")
resGreater$p.value - GS$p.value
unlist(resGreater[c("lower", "upper")]) - GS$conf.int

```

vcov.ate

*Variance-Covariance Matrix for the Average Treatment Effect.***Description**

Variance covariance matrix for the estimated average treatment effect.

Usage

```

## S3 method for class 'ate'
vcov(
  object,
  contrasts = NULL,
  times = NULL,
  estimator = NULL,
  type = NULL,
  ...
)

```

Arguments

object	A ate object, i.e. output of the ate function.
contrasts	[character vector] levels of the treatment variable for which the variance-covariance matrix should be assessed. Default is to consider all levels.
times	[numeric vector] The timepoints at which the variance-covariance matrix should be displayed. Default is to consider all timepoints.
estimator	[character] The type of estimator relative to which the variance-covariance matrix should be displayed.
type	[character] should the variance-covariance matrix w.r.t. the average risk per treatment be displayed ("meanRisk"), or the difference in average risk between any two pairs of treatments ("diffRisk"), or the ratio in average risk between any two pairs of treatments ("ratioRisk").
...	Not used. For compatibility with the generic method.

Value

A numeric matrix.

Author(s)

Brice Ozenne <broz@sund.ku.dk>

vcov.wglm

Variance-covariance for IPCW Logistic Regressions

Description

Compute the variance-covariance matrix of the estimated model parameters of IPCW logistic regressions.

Usage

```
## S3 method for class 'wglm'
vcov(object, times = NULL, type = "robust", simplify = TRUE, ...)
```

Arguments

object	a wglm object.
times	[numeric vector] time points at which the variance-covariance matrix should be output.
type	[character] should the robust variance-covariance matrix be computing accounting for the uncertainty of the IPCW ("robust") or ignoring the uncertainty of the IPCW ("robust-wknown"), or model-based ignoring the uncertainty of the IPCW ("model-wknown")?
simplify	[logical] should the output be converted to a matrix when only one timepoint is requested. Otherwise will always return a list.
...	Not used.

weights.wglm

Extract IPCW Weights

Description

Extract IPCW weights of IPCW logistic regressions.

Usage

```
## S3 method for class 'wglm'
weights(object, times = NULL, prefix = "t", simplify = TRUE, ...)
```

Arguments

object	a wglm object.
times	[numeric vector] time points at which the weights should be output.
prefix	[character] used to name the columns. Can be NA to keep the original names.
simplify	[logical] should the output be converted to a vector when only one timepoint is requested. Otherwise will always return a matrix.
...	Not used.

wglm

*Logistic Regression Using IPCW***Description**

Logistic regression over multiple timepoints where right-censoring is handled using inverse probability of censoring weighting (IPCW).

Usage

```
wglm(
  formula.event,
  times,
  data,
  formula.censor = ~1,
  cause = NA,
  fitter = NULL,
  ties = NULL,
  product.limit = NULL,
  iid = FALSE,
  se = TRUE,
  store = NULL
)
```

Arguments

formula.event	[formula] a formula with a Surv object on the left hand side and the covariates for the logistic regression on the right hand side.
times	[numeric vector] time points at which to model the probability of experiencing an event.
data	[data.frame] dataset containing the time at which the event occurred, the type of event, and regressors used to fit the censoring and logistic models.
formula.censor	[formula] an optional formula indicating on its right hand side the covariates for the censoring model.
cause	[character or numeric] the cause of interest. Defaults to the first cause.
fitter	[character] routine to fit the Cox regression models.

<code>ties</code>	[character] method used to handle ties when using a Cox model ("breslow" or "efron"). Ignored if <code>fitter</code> equals to "prodlm".
<code>product.limit</code>	[logical] if TRUE the survival is computed using the product limit estimator.
<code>iid</code>	[logical] should the influence function of the logistic regression parameters be computed, accounting for the uncertainty of the weights. This can be computationally and memory intensive.
<code>se</code>	[logical] should the variance-covariance matrix of the logistic regression parameters be stored, accounting for the uncertainty of the weights. This can be computationally and memory intensive.
<code>store</code>	[vector] when evaluating the iid, should prediction be only computed for unique covariate sets and mapped back to the original dataset (<code>data="minimal"</code>). Otherwise use <code>data="full"</code> .

Details

First, a Cox model is fitted (argument `formula.censor`) and the censoring probabilities are computed relative to each timepoint (argument `times`) to obtain the censoring weights. Then, for each timepoint, a logistic regression is fitted with the appropriate censoring weights and where the outcome is the indicator of having experience the event of interest (argument `cause`) at or before the timepoint.

Value

an object of class "wglm".

See Also

[coef.wglm](#) to output the estimated parameters from the logistic regression.
[confint.wglm](#) to output the estimated parameters from the logistic regression with their confidence interval.
[model.tables.wglm](#) to output a data.frame containing the estimated parameters from the logistic regression with its confidence intervals and p-values.
[predictRisk.wglm](#) to evaluate event probabilities (e.g. survival probabilities) conditional on covariates.
[summary.wglm](#) for displaying in the console a summary of the results.
[weights.wglm](#) to extract the IPCW weights.

Examples

```
library(survival)

#### simulate data ####
set.seed(10)
n <- 250
tau <- 1:5
d <- sampleData(n, outcome = "competing.risks")
dFull <- d[event!=0] ## (artificially) remove censoring
dSurv <- d[event!=2] ## (artificially) remove competing risk

##### no censoring #####
```

```
e.wglm <- wglm(Surv(time,event) ~ X1,
              times = tau, data = dFull, product.limit = TRUE)
e.wglm ## same as a logistic regression at each timepoint

coef(e.wglm)
confint(e.wglm)
model.tables(e.wglm)

summary(ate(e.wglm, data = dFull, times = tau, treatment = "X1", verbose = FALSE))

#### right-censoring ####
## no covariate in the censoring model (independent censoring)
eC.wglm <- wglm(Surv(time,event) ~ X1,
               times = tau, data = dSurv, product.limit = TRUE)
summary(eC.wglm)

weights(eC.wglm)

## with covariates in the censoring model
eC2.wglm <- wglm(Surv(time,event) ~ X1 + X8, formula.censor = ~ X1*X8,
                times = tau, data = dSurv)
eC2.wglm

#### Competing risks ####
## here Kaplan-Meier as censoring model
eCR.wglm <- wglm(Surv(time,event) ~ X1, formula.censor = ~X1,
                times = tau, data = d)
eCR.wglm
summary(eCR.wglm)
eCR.wglm <- wglm(Surv(time,event) ~ X1, formula.censor = ~X1,
                times = tau, data = d)
```


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