Package ‘SmoothHazard’

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Title Estimation of Smooth Hazard Models for Interval-Censored Data with Applications to Survival and Illness-Death Models

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Description Estimation of two-state (survival) models and irreversible illness-death models with possibly interval-censored, left-truncated and right-censored data. Proportional intensities regression models can be specified to allow for covariates effects separately for each transition. We use either a parametric approach with Weibull baseline intensities or a semi-parametric approach with M-splines approximation of baseline intensities in order to obtain smooth estimates of the hazard functions. Parameter estimates are obtained by maximum likelihood in the parametric approach and by penalized maximum likelihood in the semi-parametric approach.

Encoding UTF-8

Depends R (>= 1.9.1), prodlim (>= 1.4.9)

Imports lava (>= 1.4.1), mvtnorm (>= 1.0-3)

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Fit an illness-death model

Description

Fit an illness-death model using either a semi-parametric approach (penalized likelihood with an approximation of the transition intensity functions by linear combination of M-splines) or a parametric approach (specifying Weibull distributions on the transition intensities). Left-truncated, right-censored, and interval-censored data are allowed. State 0 corresponds to the initial state, state 1 to the transient one, state 2 to the absorbant one. The allowed transitions are: 0 -> 1, 0 -> 2 and 1 -> 2.

Usage

```r
idm(
    formula01,
    formula02,
    formula12,
    data,
    maxiter = 200,
    eps = c(5, 5, 3),
    n.knots = c(7, 7, 7),
    knots = "equidistant",
    CV = FALSE,
    kappa = c(100000, 500000, 20000),
    method = "Weib",
    conf.int = 0.95,
    print.iter = FALSE,
    subset = NULL,
    na.action = na.fail
)
```
Arguments

**formula01**
A formula specifying a regression model for the 0 --> 1 transition from the initial state to the transient state of the illness-death model. The right hand side of the formula specifies the covariate terms, and the left hand side must be equal to the left hand side of `formula01`. If missing it is set to `formula01`.

**formula02**
A formula specifying a regression model for the 0 --> 2 transition from the initial state to the absorbing state. The left hand side must be equal to the left hand side of `formula01`. If missing it is set to `formula01`.

**formula12**
A formula specifying a regression model for the 1 --> 2 transition from the transient state to the absorbing state. If missing it is set to `formula01`.

**data**
A data frame in which to interpret the variables of `formula01`, `formula02` and `formula12`.

**maxiter**
Maximum number of iterations. The default is 200.

**eps**
A vector of 3 integers >0 used to define the power of three convergence criteria: 1. for the regression parameters, 2. for the likelihood, 3. for the second derivatives. The default is c(5,5,3) which is translated into convergence if the respective values change less than 10^{-5} (for regression parameters and likelihood) and 10^{-3} for the second derivatives between two iterations.

**n.knots**
For `method="Splines"` only, a vector of length 3 specifying the number of knots, one for each transition, for the M-splines estimate of the baseline intensities in the order 0 --> 1, 0 --> 2, 1 --> 2. The default is c(7,7,7). When knots are specified as a list this argument is ignored. The algorithm needs least 5 knots and at most 20 knots.

**knots**
Argument only active for the penalized likelihood approach `method="Splines"`. There are three ways to control the placement of the knots between the smallest and the largest of all time points:

- knots="equidistant" Knots are placed with same distance on the time scale.
- knots="quantiles" Knots are placed such that the number of observations is roughly the same between knots.
- knots=list() List of 1 or 2 or three vectors. The list elements are the actual placements (timepoints) of the knots for the M-spline. The list may contain one vector of placements for each transition in the order 0 --> 1, 0 --> 2, 1 --> 2. If only vector is specified the knots are used for all transitions. If only 2 vectors are specified, the knots for the 0 --> 1 transition are also used for the 1 --> 2 transition.

The algorithm needs at least 5 knots and allows no more than 20 knots.

**CV**
Binary variable equals to 1 when search (by approximated cross validation) of the smoothing parameters `kappa` and 0 otherwise. Argument for the penalized likelihood approach. The default is 0.

**kappa**
Argument only active for the penalized likelihood approach `method="Splines"`. A vector with 3 positive values (smoothing parameters), one for each transition, in the order 0 --> 1, 0 --> 2 and 1 --> 2. If CV=1 these are used as starting values for a cross validation search to optimize kappa.
method : type of estimation method: "Splines" for a penalized likelihood approach with approximation of the transition intensities by M-splines, "Weib" for a parametric approach with a Weibull distribution on the transition intensities. Default is "Weib".

conf.int : Level of confidence pointwise confidence intervals of the transition intensities, i.e., a value between 0 and 1, the default is 0.95. The default is also used when conf.int=TRUE. To avoid computation of confidence intervals, set conf.int to FALSE or NULL.

print.iter : boolean parameter. Equals to TRUE to print the likelihood during the iteration process, FALSE otherwise. Default is FALSE. This option is not running on Windows.

subset : expression indicating the subset of the rows of data to be used in the fit. All observations are included by default.

na.action : how NAs are treated. The default is first, any na.action attribute of data, second a na.action setting of options, and third 'na.fail' if that is unset. The 'factory-fresh' default is na.omit. Another possible value is NULL.

Details

The estimated parameters are obtained using the robust Marquardt algorithm (Marquardt, 1963) which is a combination between a Newton-Raphson algorithm and a steepest descent algorithm.

Value

call : the call that produced the result.
coef : regression parameters.
loglik : vector containing the log-likelihood without and with covariate.
cv : vector containing the convergence criteria.
niter : number of iterations.
converged : integer equal to 1 when the model converged, 2, 3 or 4 otherwise.
modelPar : Weibull parameters.
N : number of subjects.
events1 : number of events 0 -> 1.
events2 : number of events 0 -> 2 or 0 -> 1 -> 2.
NC : vector containing the number of covariates on transitions 0 -> 1, 0 -> 2, 1 -> 2.
responseTrans : model response for the 0 -> 1 transition. Hist or Surv object.
responseAbs : model response for the 0 -> 2 transition. Hist or Surv object.
time : times for which transition intensities have been evaluated for plotting. Vector in the Weibull approach. Matrix in the penalized likelihood approach for which the columns corresponds to the transitions 0 -> 1, 1 -> 2, 0 -> 2.
intensity01 : matched values of the intensities for transition 0 -> 1.
lowerIntensity01 : lower confidence intervals for the values of the intensities for transition 0 -> 1.
upperIntensity01
upper confidence intervals for the values of the intensities for transition 0 -> 1.

intensity02
matched values of the intensities for transition 0 -> 2.

lowerIntensity02
lower confidence intervals for the values of the intensities for transition 0 -> 2.

upperIntensity02
upper confidence intervals for the values of the intensities for transition 0 -> 2.

intensity12
matched values of the intensities for transition 1 -> 2.

lowerIntensity12
lower confidence intervals for the values of the intensities for transition 1 -> 2.

upperIntensity12
upper confidence intervals for the values of the intensities for transition 1 -> 2.

RR
vector of relative risks.

V
variance-covariance matrix derived from the Hessian of the log-likelihood if using method="Weib" or, from the Hessian of the penalized log-likelihood if using method="Splines".

se
standard errors of the regression parameters.

Xnames01
names of covariates on 0 -> 1.

Xnames02
names of covariates on 0 -> 2.

Xnames12
names of covariates on 1 -> 2.

knots01
knots to approximate by M-splines the intensity of the 0 -> 1 transition.

knots02
knots to approximate by M-splines the intensity of the 0 -> 2 transition.

knots12
knots to approximate by M-splines the intensity of the 1 -> 2 transition.

nknots01
number of knots on transition 0 -> 1.

nknots02
number of knots on transition 0 -> 2.

nknots12
number of knots on transition 1 -> 2.

theta01
square root of splines coefficients for transition 0 -> 1.

theta02
square root of splines coefficients for transition 0 -> 2.

theta12
square root of splines coefficients for transition 1 -> 2.

CV
a binary variable equals to 1 when search of the smoothing parameters kappa by approximated cross-validation, 1 otherwise. The default is 0.

kappa
vector containing the smoothing parameters for transition 0 -> 1, 0 -> 2, 1 -> 2 used to estimate the model by the penalized likelihood approach.

CVcrit
cross validation criteria.

DoF
degrees of freedom of the model.

na.action
observations deleted if missing values.

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References


See Also

`print.idm` `summary.idm` `predict.idm`

Examples

```r
library(lava)
library(prodlim)
set.seed(17)
d <- simulateIDM(100)
# right censored data
fitRC <- idm(formula01=Hist(time=observed.illtime,event=seen.ill)~X1+X2,
             formula02=Hist(time=observed.lifetime,event=seen.exit)~X1+X2,
             formula12=Hist(time=observed.lifetime,event=seen.exit)~X1+X2,data=d,
             conf.int=FALSE)
fitRC

## Not run:
set.seed(17)
d <- simulateIDM(300)
fitRC.splines <- idm(formula01=Hist(time=observed.illtime,event=seen.ill)~X1+X2,
                     formula02=Hist(time=observed.lifetime,event=seen.exit)~X1+X2,
                     formula12=Hist(time=observed.lifetime,event=seen.exit)~1,data=d,
                     conf.int=FALSE,method="splines")
## End(Not run)

# interval censored data
fitIC <- idm(formula01=Hist(time=list(L,R),event=seen.ill)~X1+X2,
             formula02=Hist(time=observed.lifetime,event=seen.exit)~X1+X2,
             formula12=Hist(time=observed.lifetime,event=seen.exit)~X1+X2,data=d,
             conf.int=FALSE)
fitIC

## Not run:
data(Paq1000)
# Illness-death model with certif on the 3 transitions
# Weibull parametrization and likelihood maximization
fit.weib <- idm(formula02=Hist(time=t,event=death,entry=e)~certif,
                formula01=Hist(time=list(l,r),event=dementia)~certif,
                data=Paq1000)

# Illness-death model with certif on transitions 01 and 02
# Splines parametrization and penalized likelihood maximization
fit.splines <- idm(formula02=Hist(time=t,event=death,entry=e)~certif,
                    formula01=Hist(time=list(l,r),event=dementia)~certif,
                    method="splines")
```
idmModel

Generate illness-death model objects

Description

Function to generate an illness-death model for simulation.

Usage

idmModel(
  scale.illtime = 1/100,
  shape.illtime = 1,
  scale.lifetime = 1/100,
  shape.lifetime = 1,
  scale.waittime = 1/100,
  shape.waittime = 1,
  scale.censtime = 1/100,
  shape.censtime = 1,
  n.inspections = 5,
  schedule = 10,
  punctuality = 5
)

Arguments

scale.illtime  Weibull scale for latent illness time
shape.illtime  Weibull shape for latent illness time
scale.lifetime  Weibull scale for latent life time
shape.lifetime  Weibull shape for latent life time
scale.waittime  Weibull scale for latent life time
shape.waittime  Weibull shape for latent life time
scale.censtime  Weibull scale for censoring time
shape.censtime  Weibull shape for censoring time
n.inspections  Number of inspection times
schedule  Mean of the waiting time between adjacent inspections.
punctuality  Standard deviation of waiting time between inspections.
Details

Based on the functionality of the lava PACKAGE the function generates a latent variable model (latent illtime, waittime, and lifetime) and censoring mechanism (censtime, inspection1, inspection2, ..., inspectionK). The function `sim.idmModel` then simulates right censored lifetimes and interval censored illness times.

Value

A latent variable model object `lvm`

Author(s)

Thomas Alexander Gerds

Examples

```r
# Not run:
library(lava)
library(prodlim)
# generate illness-death model based on exponentially
distributed times
m <- idmModel(scale.illtime=1/70,
  shape.illtime=1.8,
  scale.lifetime=1/50,
  shape.lifetime=0.7,
  scale.waittime=1/30,
  shape.waittime=0.7)
round(sim(m,6),1)

# Estimate the parameters of the Weibull models
# based on the uncensored exact event times
# and the uncensored illstatus.
set.seed(18)
d <- sim(m,100,latent=FALSE)
d$uncensored.status <- 1
f <- idm(formula01=Hist(time=illtime,event=illstatus)-1,
  formula02=Hist(time=lifetime,event=uncensored.status)-1,
data=d,
  conf.int=FALSE)
print(f)

# Change the rate of the 0->2 and 0->1 transitions
# also the rate of the 1->2 transition
# and also lower the censoring rate
m <- idmModel(scale.lifetime=1/2000,
  scale.waittime=1/30,
  scale.illtime=1/1000,
  scale.censtime=1/1000)
set.seed(18)
d <- sim(m,50,latent=TRUE)
d$uncensored.status <- 1
```
f <- idm(formula01=Hist(time=observed.illtime,event=illstatus)~1,
         formula02=Hist(time=observed.lifetime,event=uncensored.status)~1,
         data=d,
         conf.int=FALSE)
print(f)

# Estimate based on the right censored observations
fc <- idm(formula01=Hist(time=illtime,event=seen.ill)~1,
         formula02=Hist(time=observed.lifetime,event=seen.exit)~1,
         data=d,
         conf.int=FALSE)
print(fc)

# Estimate based on interval censored and right censored observations
fi <- idm(formula01=Hist(time=list(L,R),event=seen.ill)~1,
         formula02=Hist(time=observed.lifetime,event=seen.exit)~1,
         data=d,
         conf.int=FALSE)
print(fi)

# Estimation of covariate effects:
# X1, X2, X3
m <- idmModel(shape.waittime=2,
               scale.lifetime=1/2000,
               scale.waittime=1/300,
               scale.illtime=1/10000,
               scale.censtime=1/10000)
distribution(m,"X1") <- binomial.lvm(p=0.3)
distribution(m,"X2") <- normal.lvm(mean=120,sd=20)
distribution(m,"X3") <- normal.lvm(mean=50,sd=20)
regression(m,to="latent.illtime",from="X1") <- 1.7
regression(m,to="latent.illtime",from="X2") <- 0.07
regression(m,to="latent.illtime",from="X3") <- -0.1
regression(m,to="latent.waittime",from="X1") <- 1.8
set.seed(28)
d <- sim(m,100,latent=TRUE)
head(d)
table(ill=d$seen.ill,death=d$seen.exit)

# Estimation based on uncensored data
d$uncensored.status <- 1
# uncensored data
F1 <- idm(formula01=Hist(time=illtime,event=illstatus)~X1+X2+X3,
          formula02=Hist(time=lifetime,event=uncensored.status)~X1+X2+X3,
          data=d,conf.int=FALSE)
print(F1)

# Estimation based on right censored data
F2 <- idm(formula01=Hist(time=illtime,event=seen.ill)~X1+X2+X3,
          formula02=Hist(time=observed.lifetime,event=seen.exit)~X1+X2+X3,
          data=d,conf.int=FALSE)
intensity

M-spline estimate of the transition intensity function

Description

M-spline estimate of the transition intensity function and the cumulative transition intensity function for survival and illness-death models

Usage

intensity(times, knots, number.knots, theta, linear.predictor = 0)

Arguments

times Time points at which to estimate the intensity function
knots Knots for the M-spline
number.knots Number of knots for the M-splines (and I-splines see details)
theta The coefficients for the linear combination of M-splines (and I-splines see details)
linear.predictor Linear predictor beta*Z. When it is non-zero, transition and cumulative transition are multiplied by \( \exp(\text{linear.predictor}) \). Default is zero.

Details

The estimate of the transition intensity function is a linear combination of M-splines and the estimate of the cumulative transition intensity function is a linear combination of I-splines (the integral of a M-spline is called I-spline). The coefficients theta are the same for the M-splines and I-splines.

Important: the theta parameters returned by idm and shr are in fact the square root of the splines coefficients. See examples.

This function is a R-translation of a corresponding Fortran function called susp. susp is used internally by idm and shr.
Value

- **times**: The time points at which the following estimates are evaluated.
- **intensity**: The transition intensity function evaluated at times.
- **cumulative.intensity**: The cumulative transition intensity function evaluated at times.
- **survival**: The "survival" function, i.e., \( \exp(-\text{cumulative.intensity}) \)

Author(s)

R: Celia Touraine <Celia.Touraine@isped.u-bordeaux2.fr> and Thomas Alexander Gerds <tag@biostat.ku.dk>

See Also

- shr, idm

Examples

data(testdata)
fitted.su <- shr(Hist(time=list(l, r), id) ~ cov,
  data = testdata, method = "Splines", CV = TRUE)
intensity(times = fitted.su$time, knots = fitted.su$knots,
  number.knots = fitted.su$nknots, theta = fitted.su$\theta^2)

```r
# Not run:
data(Paq1000)
fitted.idm <- idm(formula02 = Hist(time = t, event = death, entry = e) ~ certif,
  formula01 = Hist(time = list(l, r), event = dementia) ~ certif,
  formula12 = ~ certif, method = "Splines", data = Paq1000)
# Probability of survival in state 0 at age 80 for a subject with no cep given
# that he is in state 0 at 70
su0 <- (intensity(times = 80, knots = fitted.idm$knots01,
  number.knots = fitted.idm$nknots01,
  theta = fitted.idm$\theta01^2)$survival
  * intensity(times = 80, knots = fitted.idm$knots02,
    number.knots = fitted.idm$nknots02,
    theta = fitted.idm$\theta02^2)$survival)/
  (intensity(times = 70, knots = fitted.idm$knots01,
    number.knots = fitted.idm$nknots01,
    theta = fitted.idm$\theta01^2)$survival
  * intensity(times = 70, knots = fitted.idm$knots02,
    number.knots = fitted.idm$nknots02,
    theta = fitted.idm$\theta02^2)$survival)
# Same result as:
predict(fitted.idm, s = 70, t = 80, conf.int = FALSE) # see first element
```

## End(Not run)
Paq1000 Paquid data set

**Description**

Paquid data set composed of 1000 subjects selected randomly from the Paquid data set of 3675 subjects.

**Format**

A data frame with 1000 rows and the following 8 columns.

- **dementia** dementia status, 0=non-demented, 1=demented
- **death** death status, 0=alive, 1=dead
- **e** age at entry in the study
- **l** for demented subjects: age at the visit before the diagnostic visit; for non-demented subjects: age at the last visit (censoring age)
- **r** for demented subjects: age at the diagnostic visit; for non-demented subjects: age at the last visit (censoring age)
- **t** for dead subjects: age at death; for alive subject: age at the latest news
- **certif** primary school certificate: 0=with certificate, 1=without certificate
- **gender** gender: 0=female, 1=male

**Examples**

```r
data(Paq1000)
```

**plot.idm**

Plot method for an illness-death model

**Description**

Plot estimated baseline transition intensities from an object of class idm optionally with confidence limits.
Usage

```r
## S3 method for class 'idm'
plot(
  x,
  conf.int = FALSE,
  ctype = "shadow",
  add = FALSE,
  axes = TRUE,
  col,
  lwd,
  lty,
  xlim,
  ylim,
  xlab,
  ylab,
  legend = TRUE,
  transition = c("01", "02", "12"),
  ...
)
```

Arguments

- `x`: a `idmWeib` class object (output from calling `idm` with the (default) option `intensities="Weib"`).
- `conf.int`: If TRUE show confidence limits
- `ctype`: Type of confidence limits, can be "shadow" or "bars"
- `add`: If TRUE add to existing plot
- `axes`: If TRUE axes are drawn
- `col`: Color of the lines
- `lwd`: Width of the lines
- `lty`: Type of the lines
- `xlim`: Limits for x-axis
- `ylim`: Limits for y-axis
- `xlab`: Label for x-axis
- `ylab`: Label for y-axis
- `legend`: If TRUE a legend is drawn, which can be further controlled via `SmartControl`.
- `transition`: Choose one of the transition intensities: c("01", "02", "12").
- `...`: Passed to `SmartControl`

Value

Print a plot of the baseline transition intensities of an illness-death model estimated using a Weibull approach.
See Also

print.idm, summary.idm, idm

Examples

library(lava)
library(prodlim)
m <- idmModel(scale.lifetime=1/10, scale.illtime=1/8)
distribution(m,"X") <- binomial.lvm()
regression(m, latent.lifetime-X) <- 0.7
set.seed(30)
d <- sim(m,100)
fit.weib <- idm(formula02=Hist(observed.lifetime, event=seen.exit)-1,
                formula01=Hist(time=list(L,R), event=seen.ill)-1,
data=d, conf.int=FALSE)
plot(fit.weib)

## Not run:
## FIXME: the limits for the 01 transition are a bit wide!?
## with bootstrap confidence limits
fit.weib <- idm(formula02=Hist(observed.lifetime, event=seen.exit)-1,
                formula01=Hist(time=list(L,R), event=seen.ill)-1,
data=d, conf.int=TRUE)
plot(fit.weib)

## End(Not run)
plot.shr

lwd,
ylim,
xlim,
xlab = "Time",
ylab,
legend = TRUE,
confint = TRUE,
timeOrigin = 0,
axes = TRUE,
percent = TRUE,
...
)

Arguments

x a shrWeib or a shrSplines class object (output from calling shr function).

type type of function to plot. The default is "shr".

add boolean.

cause newdata.

col newdata.

lty lty.

lwd lwd.

ylim ylim.

xlim xlim.

xlab xlab.

ylab ylab.

legend legend.

confint confint.

timeOrigin timeOrigin.

axes axes.

percent percent.

... other graphical parameters.

Value

Print a plot of a survival model.

Author(s)

R: Celia Touraine <Celia.Touraine@isped.u-bordeaux2.fr> Fortran: Pierre Joly <Pierre.Joly@isped.u-bordeaux2.fr>
predict.idm

Predictions for an illness-death model using either a penalized likelihood approach or a Weibull parametrization.

Description

Predict transition probabilities and cumulative probabilities from an object of class idmSplines with confidence intervals are calculated.

Usage

```r
# S3 method for class 'idm'
predict(
  object,
  s,
  t,
  newdata,
  nsim = 200,
  seed = 21,
  conf.int = 0.95,
  lifeExpect = FALSE,
  maxtime,
  ...
)
```
predict.idm

Arguments

- **object**
  - an idm class objects returned by a call to the `idm` function
- **s**
  - time point at which prediction is made.
- **t**
  - time horizon for prediction.
- **newdata**
  - A data frame with covariate values for prediction.
- **nsim**
  - number of simulations for the confidence intervals calculations. The default is 200.
- **seed**
  - Seed passed to `set.seed` for Monte Carlo simulation of confidence intervals.
- **conf.int**
  - Level of confidence, i.e., a value between 0 and 1, the default is 0.95. The default is also used when `conf.int=TRUE`. To avoid computation of confidence intervals, set `conf.int` to FALSE or NULL.
- **lifeExpect**
  - Logical. If `TRUE` compute life expectancies, i.e., `t=Inf`.
- **maxtime**
  - The upper limit of integration for calculations of life expectancies from Weibull parametrizations.
- **...**
  - other parameters.

Value

a list containing the following predictions with pointwise confidence intervals:

- **p00**
  - the transition probability \( p_{00} \).
- **p01**
  - the transition probability \( p_{01} \).
- **p11**
  - the transition probability \( p_{11} \).
- **p12**
  - the transition probability \( p_{12} \).
- **p02_0**
  - the probability of direct transition from state 0 to state 2.
- **p02_1**
  - the probability of transition from state 0 to state 2 via state 1.
- **p02**
  - transition probability \( p_{02} \). Note that \( p_{02}=p_{02_0}+p_{02_1} \).
- **F01**
  - the lifetime risk of disease. \( F_{01}=p_{01}+p_{02_1} \).
- **F0.**
  - the probability of exit from state 0. \( F_{0.}=p_{02_0}+p_{01}+p_{02_1} \).

Author(s)

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See Also

`idm`
Examples

```r
## Not run:
set.seed(100)
d=simulateIDM(n = 100)
fit <- idm(formula01=Hist(time=list(L,R),event=seen.ill)~X1+X2+X3,
           formula02=Hist(time=observed.lifetime,event=seen.exit)~X1+X2+X3,
           data=d,conf.int=FALSE)
predict(fit,s=0,t=80,conf.int=FALSE,lifeExpect=FALSE)
predict(fit,s=0,t=80,nsim=4,conf.int=TRUE,lifeExpect=FALSE)
predict(fit,s=0,t=80,nsim=4,conf.int=FALSE,lifeExpect=TRUE)

data(Paq1000)
library(prodlim)
fit.paq <- idm(formula02=Hist(time=t,event=death,entry=e)~certif,
                formula01=Hist(time=list(l,r),event=dementia)~certif,data=Paq1000)
predict(fit.paq,s=70,t=80,newdata=data.frame(certif=1))
predict(fit.paq,s=70,lifeExpect=TRUE,newdata=data.frame(certif=1))

fit.splines <- idm(formula02=Hist(time=t,event=death,entry=e)~certif,
                    formula01=Hist(time=list(l,r),event=dementia)~certif,
                    formula12=-1,
                    method="Splines",
                    data=Paq1000)
predict(fit.splines,s=70,t=80,newdata=data.frame(certif=1))
predict(fit.splines,s=70,t=80,lifeExpect=TRUE,newdata=data.frame(certif=1),nsim=20)

## End(Not run)
```

print.idm

Print method for `idm` objects

**Description**

Print a summary of a fitted illness-death model

**Usage**

```r
## S3 method for class 'idm'
print(x, conf.int = 0.95, digits = 4, pvalDigits = 4, eps = 0.0001, ...)
```

**Arguments**

- `x`: Class `idm` object, i.e. the result of a call to the `idm` function with intensities="Weib". 
Print method for `shrSplines` objects

Description

Print a summary of a fitted illness-death model using the penalized likelihood approach.

Usage

```r
## S3 method for class 'shr'
print(x, conf.int = 0.95, digits = 4, pvalDigits = 4, eps = 0.0001, ...)
```
### Arguments

- **x**: a `shr` object, i.e., the result of a call to the `shr` function with `hazard="Splines"`.
- **conf.int**: The level of confidence for the hazard ratios. The default is 0.95.
- **digits**: number of digits to print.
- **pvalDigits**: number of digits to print for p-values.
- **eps**: convergence criterion used for p-values.
- **...**: other unused arguments.

### Author(s)

R: Celia Touraine <Celia.Touraine@isped.u-bordeaux2.fr> Fortran: Pierre Joly <Pierre.Joly@isped.u-
bordeaux2.fr>

### See Also

- `summary.shr`
- `plot.shr`

### Examples

```r
## Not run:
# a penalized survival model
library(prodlim)
data(testdata)
fit.su <- shr(Hist(time=list(l,r),id)~cov,data=testdata,method="Splines")
print(fit.su)
## End(Not run)
```

---

**shr**

*Fit a survival model*

### Description

Fit a survival model using either a semi-parametric approach (penalized likelihood with an approximation of the hazard function by linear combination of M-splines) or a parametric approach (specifying a Weibull distribution on the hazard function). Left-truncated, right-censored, and interval-censored data are allowed.

### Usage

```r
shr(
    formula,  # a formula
    data,     # a data frame
    eps = c(5, 5, 3),  # convergence criteria
    n.knots = 7,     # number of knots
)
```
\begin{verbatim}
knots = "equidistant",
CV = FALSE,
kappa = 10000,
conf.int = 0.95,
maxiter = 200,
method = "Weib",
print.iter = FALSE,
na.action = na.omit
)
\end{verbatim}

**Arguments**

- **formula**
  a formula object with the response on the left hand side and the terms on the right hand side. The response must be a survival object or Hist object as returned by the 'Surv' or 'Hist' function.

- **data**
  a data frame in which to interpret the variables named in the formula.

- **eps**
  a vector of length 3 for the convergence criteria (criterion for parameters, criterion for likelihood, criterion for second derivatives). The default is \(c(5,5,3)\) and corresponds to criteria equals to \(10^{-5}\), \(10^{-5}\) and \(10^{-3}\).

- **n.knots**
  Argument only active for the penalized likelihood approach method="splines". Number of knots for the splines to use to approximate the hazard function. The default is 7. If knots are given as a vector this argument is ignored. The algorithm needs at least 5 knots and at most 20 knots.

- **knots**
  Argument only active for the penalized likelihood approach method="splines". There are three ways to control the placement of the knots between the smallest and the largest of all time points:

  - knots=":equidistant" Knots are placed with same distance on the time scale.
  - knots=":quantiles" Knots are placed such that the number of observations is roughly the same between knots.
  - knots=list() List of length 3. The list elements are the actual placements (timepoints) of the knots for the M-spline.

  The algorithm requires at least 5 knots and allows no more than 20 knots.

- **CV**
  binary variable equals to 1 when search (by approximated cross validation) of the smoothing parameter kappa and 0 otherwise. Argument for the penalized likelihood approach. The default is 0.

- **kappa**
  Argument only active for the penalized likelihood approach method=":splines". A positive number (smoothing parameter) If CV=1 the value is used as a starting value for a cross validation search to optimize kappa.

- **conf.int**
  Level of confidence pointwise confidence intervals of the survival and hazard functions, i.e., a value between 0 and 1, the default is 0.95. The default is also used when conf.int=TRUE. To avoid computation of confidence intervals, set conf.int to FALSE or NULL.

- **maxiter**
  maximum number of iterations. The default is 200.
method: type of estimation method: "Splines" for a penalized likelihood approach with approximation of the hazard function by M-splines, "Weib" for a parametric approach with a Weibull distribution on the hazard function. Default is "Weib".

print.iter: boolean parameter. Equals to TRUE to print the likelihood during the iteration process, FALSE otherwise. Default is FALSE. This option is not running on Windows.

na.action: how NAs are treated. The default is first, any na.action attribute of data, second a na.action setting of options, and third 'na.fail' if that is unset. The 'factory-fresh' default is na.omit. Another possible value is NULL.

Details
The estimated parameters are obtained using the robust Marquardt algorithm (Marquardt, 1963) which is a combination between a Newton-Raphson algorithm and a steepest descent algorithm.

Value
• call
• coef: regression parameters.
• loglik: vector containing the log-likelihood without and with covariate.
• modelPar: Weibull parameters.
• N: number of subjects.
• NC: number of covariates.
• nevents: number of events.
• modelResponse: model response: Hist or Surv object.
• converged: integer equal to 1 when the model converged, 2, 3 or 4 otherwise.
• times: integer equal to 1 when the model converged, 2, 3 or 4 otherwise.
• hazard: matched values of the hazard function.
• lowerHazard: lower confidence limits for hazard function.
• upperHazard: upper confidence limits for hazard function.
• surv: matched values of the survival function.
• lowerSurv: lower confidence limits for survival function.
• upperSurv: upper confidence limits for survival function.
• RR: vector of relative risks.
• V: variance-covariance matrix.
• sestandard errors.
• knots: knots of the M-splines estimate of the hazard function.
• nknots: number of knots.
• CV: a binary variable equals to 1 when search of the smoothing parameter kappa by approximated cross-validation, 1 otherwise. The default is 0.
• niter: number of iterations.
• cv: vector containing the convergence criteria.
• na.action: observations deleted if missing values.
sim.idmModel

Simulate illness-death model data

Description

Function to simulate illness-death model data

Usage

## S3 method for class 'idmModel'
sim(
x, n, illness.known.at.death = TRUE, compliance = 1, latent = FALSE,
)
Arguments

x  An idmModel object as obtained with idmModel

n  Number of observations

illness.known.at.death  Affects the value of variable seen.ill

compliance  Probability of missing an inspection time.

latent  if TRUE keep the latent event times

keep.inspectiontimes  if TRUE keep the inspection times.

...  Extra arguments given to sim

Details

Based on the functionality of the lava PACKAGE

Value

A data set with interval censored observations from an illness-death model

Author(s)

Thomas Alexander Gerds

Examples

example(idmModel)
help(idmModel)

Description

Function to simulate interval censored survival data

Usage

## S3 method for class 'survIC'
sim(x, n, compliance = 1, latent = TRUE, keep.inspectiontimes = FALSE, ...)

Simulate interval censored survival data
**simulateIDM**

**Arguments**

- `x` An `survIC` object as obtained with `survIC`
- `n` Number of observations
- `compliance` Probability of missing an inspection time.
- `latent` if TRUE keep the latent event times
- `keep.inspectiontimes` if TRUE keep the inspection times.
- `...` Extra arguments given to `sim`

**Details**

Based on the functionality of the `lava` PACKAGE

**Value**

A data set with interval censored observations

**Author(s)**

Thomas Alexander Gerds

**Examples**

```r
library(lava)
example(survIC)
help(survIC)
ol <- survIC()
dat.ol <- sim(ol,10)
```

---

**simulateIDM**  
*Sample illness-death model data*

**Description**

Simulate data from an illness-death model with interval censored event times and covariates

**Usage**

```r
simulateIDM(n = 100)
```

**Arguments**

- `n` number of observations
Details

Simulate data from an illness-death model with interval censored event times and covariates for the purpose of illustrating the help pages of the SmoothHazard package. See the body of the function for details, i.e., evaluate simulateIDM

See Also

idmModel sim.idmModel

Examples

# simulateIDM
simulateIDM(100)

summary.idm

Summary of a fitted illness-death model

Description

Summarize the event history data of an illness-death regression model and show regression coefficients for transition intensities

Usage

## S3 method for class 'idm'
summary(object, conf.int = 0.95, digits = 4, pvalDigits = 4, eps = 0.0001, ...)

Arguments

object a idmSplines object, i.e., the result of a call to the idm function with intensities="Splines".
conf.int The level of confidence for the hazard ratios. The default is 0.95.
digits number of digits to print.
pvalDigits number of digits to print for p-values.
eps convergence criterion used for p-values.
... other unused arguments.

Author(s)

R: Celia Touraine <Celia.Touraine@isped.u-bordeaux2.fr> Fortran: Pierre Joly <Pierre.Joly@isped.u-bordeaux2.fr>

See Also

idm, print.idm, plot.idm
Examples

```r
## Not run:
library(prodlim)
data(Paq1000)
fit.splines <- idm(formula02=Hist(time=t,event=death,entry=e)~certif,
                   formula01=Hist(time=list(l,r),event=dementia)~certif,
                   formula12=~1,
                   method="Splines",
                   data=Paq1000)
summary(fit.splines)

## End(Not run)
```

### summary.shr

#### Summary of a fitted survival model using a penalized likelihood approach

**Description**

Print a short summary of a fitted illness-death model using the penalized likelihood approach.

**Usage**

```r
## S3 method for class 'shr'
summary(object, conf.int = 0.95, digits = 4, pvalDigits = 4, eps = 0.0001, ...)
```

**Arguments**

- `object`:
  a `shr` object, i.e., the result of a call to the `shr` function.
- `conf.int`:
  The level of confidence for the hazard ratios. The default is 0.95.
- `digits`:
  number of digits to print.
- `pvalDigits`:
  number of digits to print for p-values.
- `eps`:
  convergence criterion used for p-values.
- `...`:
  other unused arguments.

**Author(s)**

Celia Touraine <Celia.Touraine@isped.u-bordeaux2.fr>

**See Also**

`shr`, `print.shr`, `plot.shr`
## Examples

```
## Not run:
# a penalized survival model
data(testdata)
library(prodlim)
fit.su <- shr(Hist(time=list(l,r),id)~cov,data=testdata,method="Splines")
summary(fit.su)

# Weibull survival model
data(testdata)
fit.su <- shr(Hist(time=list(l,r),id)~cov,data=testdata)
summary(fit.su)

## End(Not run)
```

---

**survIC**

*Generate survival model objects*

**Description**

Function to generate a latent variable model for interval censored survival times.

**Usage**

```
survIC(
  scale.time = 1/100,
  shape.time = 1,
  n.inspections = 5,
  schedule = 10,
  punctuality = 5
)
```

**Arguments**

- **scale.time**: Weibull scale for latent time
- **shape.time**: Weibull shape for latent time
- **n.inspections**: Number of inspection times
- **schedule**: Mean of the waiting time between adjacent inspections.
- **punctuality**: Standard deviation of waiting time between inspections.

**Details**

Based on the functionality of the lava PACKAGE the function generates a latent variable model with a latent time and a censoring mechanism (censtime, inspection1,inspection2,...,inspectionK). The function `sim.survIC` then simulates interval censored times.
Value

A latent variable model object `lvm`

Author(s)

Thomas Alexander Gerds

Examples

```r
## Not run:
library(lava)
library(prodlim)
# generate survival model based on exponentially
# distributed times
m <- survIC(scale.time=1/50, shape.time=0.7)
round(sim(m,6),1)

# Estimate the parameters of the Weibull models
# based on the uncensored exact event times
# and the uncensored illstatus.
set.seed(18)
d <- sim(m,100,latent=FALSE)
d$uncensored.status <- 1
f <- shr(Hist(time=list(L,R),event=uncensored.status)-1,
data=d,
conf.int=FALSE)
print(f)

## End(Not run)
```

testdata

Data set for survival models: right-censored and interval-censored data.

Description

A simulated data frame for survival models composed of right-censored and interval-censored data.

Format

A data frame with 936 observations on the following 4 variables.

- `l` for diseased subjects: left endpoint of censoring interval; for non-diseased subjects: right censoring time
- `r` for diseased subjects: right endpoint of censoring interval; for non-diseased subjects: right censoring time for the disease event
- `id` disease status
- `cov` covariate
Examples

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