Package 'survPen'

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Description Fits (excess) hazard, relative mortality ratio or marginal intensity models with multidimensional penalized splines allowing for time-dependent effects, non-linear effects and interactions between several continuous covariates. In survival and net survival analysis, in addition to modelling the effect of time (via the baseline hazard), one has often to deal with several continuous covariates and model their functional forms, their time-dependent effects, and their interactions. Model specification becomes therefore a complex problem and penalized regression splines represent an appealing solution to that problem as splines offer the required flexibility while penalization limits overfitting issues. Current implementations of penalized survival models can be slow or unstable and sometimes lack some key features like taking into account expected mortality to provide net survival and excess hazard estimates. In contrast, survPen provides an automated, fast, and stable implementation (thanks to explicit calculation of the derivatives of the likelihood) and offers a unified framework for multidimensional penalized hazard and excess hazard models. Later versions (>2.0.0) include penalized models for relative mortality ratio, and marginal intensity in recurrent event setting. survPen may be of interest to those who 1) analyse any kind of time-to-event data: mortality, disease relapse, machinery breakdown, unemployment, etc 2) wish to describe the associated hazard and to understand which predictors impact its dynamics, 3) wish to model the relative mortality ratio between a cohort and a reference population, 4) wish to describe the marginal intensity for recurrent event data. See Fauvernier et al. (2019a) [<doi:10.21105/joss.01434>](https://doi.org/10.21105/joss.01434) for an overview of the pack-age and Fauvernier et al. (2019b) [<doi:10.1111/rssc.12368>](https://doi.org/10.1111/rssc.12368) for the method.

Depends R $(>= 4.0.0)$

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Imports statmod, stats, $\text{Rcpp} (> = 1.0.2)$

LinkingTo Rcpp, RcppEigen

URL <https://github.com/fauvernierma/survPen>

BugReports <https://github.com/fauvernierma/survPen/issues> Encoding UTF-8 LazyData true

Title Multidimensional Penalized Splines for (Excess) Hazard Models, Relative Mortality Ratio Models and Marginal Intensity Models

2 Contents

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colSums2 *colSums of a matrix*

Description

colSums of a matrix

Usage

colSums2(Mat)

Arguments

Mat a matrix.

Value

colSums(Mat)

Description

Applies the sum-to-zero constraints to design and penalty matrices.

Usage

 $constraint(X, S, Z = NULL)$

Arguments

Value

List of objects with the following items:

Examples

library(survPen)

set.seed(15)

 $X \leftarrow matrix(rnorm(10*3), nrow=10, ncol=3)$ S <- matrix(rnorm(3*3),nrow=3,ncol=3) ; S <- 0.5*(S + t(S))

applying sum-to-zero constraint to a desgin matrix and a penalty matrix constr <- constraint(X,S)

Description

Takes the model at convergence and calculates the variance matrix corrected for smoothing parameter uncertainty

Usage

cor.var(model)

Arguments

model survPen object, see [survPen.fit](#page-59-1) for details

Value

survPen object with corrected variance Vc

crs *Bases for cubic regression splines (equivalent to "cr" in* mgcv*)*

Description

Builds the design matrix and the penalty matrix for cubic regression splines.

Usage

 $crs(x, knots = NULL, df = 10, intercept = TRUE)$

Arguments

Details

See package mgcv and section 4.1.2 of Wood (2006) for more details about this basis

Value

List of three elements

References

Wood, S. N. (2006), Generalized additive models: an introduction with R. London: Chapman & Hall/CRC.

Examples

x <- seq(1,10,length=100) # natural cubic spline with 3 knots crs(x,knots=c(1,5,10))

crs.FP *Penalty matrix constructor for cubic regression splines*

Description

constructs the penalty matrix associated with cubic regression splines basis. This function is called inside [crs](#page-4-1).

Usage

```
crs.FP(knots, h)
```
Arguments

Value

List of two elements:

CumulHazard 7

Examples

library(survPen)

```
# construction of the penalty matrix using a sequence of knots
knots <- c(0,0.25,0.5,0.75,1)
diff.knots <- diff(knots)
```
crs.FP(knots,diff.knots)

CumulHazard *Cumulative hazard (integral of hazard) only*

Description

Cumulative hazard (integral of hazard) only

Usage

```
CumulHazard(X_GL, weights, tm, n_legendre, n, beta, is_pwcst, pwcst_weights)
```
Arguments

Value

cumulative hazard (integral of hazard)

Description

A simulated dataset containing the follow-up times of 2000 patients diagnosed with cervical cancer between 1990 and 2010. End of follow-up is June 30th 2013. The variables are as follows:

- begin. beginning of follow-up. For illustration purposes about left truncation only (0–1)
- fu. follow-up time in years $(0-5)$
- age. age at diagnosis in years, from 21.39 to 99.33
- yod. decimal year of diagnosis, from 1990.023 to 2010.999
- dead. censoring indicator (1 for dead, 0 for censored)
- rate. expected mortality rate (from overall mortality of the general population) (0–0.38)

Usage

data(datCancer)

Format

A data frame with 2000 rows and 6 variables

Description

Cumulative hazard (integral of hazard) and its first and second derivatives wrt regression parameters beta

Usage

```
DerivCumulHazard(
  X_GL,
  weights,
  tm,
  n_legendre,
  n,
  p,
  beta,
  expected,
  type,
  is_pwcst,
  pwcst_weights
)
```
deriv_R 9

Arguments

Value

List of objects with the following items:

Description

Derivative of a Choleski factor

Usage

deriv_R(deriv_Vp, p, R1)

Arguments

Value

a list containing the derivatives of R1 wrt rho (log smoothing parameters)

Description

Builds the design matrix for the whole model when the sum-to-zero constraints are specified. The function is called inside [model.cons](#page-21-1) for Gauss-Legendre quadrature.

Usage

```
design.matrix(
  formula,
  data.spec,
  t1.name,
  Z.smf,
  Z.tensor,
  Z.tint,
  list.smf,
  list.tensor,
  list.tint,
  list.rd
\mathcal{L}
```
Arguments

Value

design matrix for the model

expected.table 11

Examples

```
library(survPen)
# standard spline of time with 4 knots
data <- data.frame(time=seq(0,5,length=100),event=1,t0=0)
form \leq \sim smf(time, knots=c(0,1,3,5))
t1 <- eval(substitute(time), data)
t0 <- eval(substitute(t0), data)
event <- eval(substitute(event), data)
# Setting up the model
model.c <- model.cons(form,lambda=0,data.spec=data,t1=t1,t1.name="time",
t0=rep(0,100),t0.name="t0",event=event,event.name="event",
expected=rep(0,100),expected.name=NULL,type="overall",n.legendre=20,
cl="survPen(form,data,t1=time,event=event)",beta.ini=NULL)
# Retrieving the sum-to-zero constraint matrices and the list of knots
Z.smf <- model.c$Z.smf ; list.smf <- model.c$list.smf
# Calculating the design matrix
design.M <- design.matrix(form,data.spec=data,t1.name="time",Z.smf=Z.smf,list.smf=list.smf,
Z.tensor=NULL,Z.tint=NULL,list.tensor=NULL,list.tint=NULL,list.rd=NULL)
```
expected.table *French women mortality table*

Description

French women mortality table to serve as example of reference/expected mortality in excess hazard and relative mortality ratio models The data come from the human mortality databse website: https://www.mortality.org/Country/Country?cntr=FRATNP

- Age. Age group for 1-year interval from exact age x to just before exact age x+1 (0-110+)
- Year. Calendar Year (1816-2021)
- mx. Central death rate between ages x and x+1

Usage

```
data(expected.table)
```
Format

A data frame with 22866 rows and 3 variables

Description

Gradient vector of LCV and LAML wrt rho (log smoothing parameters)

Usage

grad_rho(X_GL, GL_temp, haz_GL, deriv_rho_beta, weights, tm, nb_smooth, p, n_legendre, S_list, temp_LAML, Vp, S_beta, beta, inverse_new_S, X, temp_deriv3, event, expected, type, Ve, mat_temp, method

Arguments

)

Value

List of objects with the following items:

Description

Gradient vector of LCV and LAML wrt rho (log smoothing parameters). Version for multiplicative decomposition : relative mortality ratio model

Usage

```
grad_rho_mult(
  X_GL,
  GL_temp,
  haz_GL,
  deriv_rho_beta,
  weights,
  tm,
  nb_smooth,
  p,
  n_legendre,
  S_list,
  temp_LAML,
  Vp,
  S_beta,
  beta,
  inverse_new_S,
  X,
  event,
  expected,
  Ve,
  mat_temp,
  method
)
```


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Value

List of objects with the following items:

grad_rho gradient vector of LCV or LAML deriv_rho_inv_Hess_beta List of first derivatives of Vp wrt rho deriv_rho_Hess_unpen_beta List of first derivatives of the Hessian of the unpenalized log-likelihood wrt rho

HazGL *Gauss-Legendre evaluations*

Description

Gauss-Legendre evaluations

Usage

HazGL(X_GL, n_legendre, beta)

Arguments

Value

list of all the matrix-vector multiplications X.GL[[i]]%*%beta for Gauss Legendre integration in order to save computation time

Description

A simulated dataset containing 3 068 observations (2 268 events) in 800 patients with heart failure. The dataset is based on hfaction_cpx12 dataset from package WA. The variables are as follows:

- id. patient identifcation number
- treatment. treatment=0 for control and treatment=1 for exercise training
- t0. beginning of follow-up for a given event
- t1. end of follow-up for a given event (up to 3.27 years)
- enum. event identification number for a given patient (between 1 and 6 events per patient)
- event. event indicator (1 for hospitalization, 0 for censored)

Usage

data(HeartFailure)

Format

A data frame with 3 068 rows and 6 variables

Hess_rho *Hessian matrix of LCV and LAML wrt rho (log smoothing parameters)*

Description

Hessian matrix of LCV and LAML wrt rho (log smoothing parameters)

Usage

Hess_rho(X_GL, X_GL_Q, GL_temp, haz_GL, deriv2_rho_beta, deriv_rho_beta, weights, tm, nb_smooth, p, n_legendre,

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```
deriv_rho_inv_Hess_beta,
deriv_rho_Hess_unpen_beta,
S_list,
minus_eigen_inv_Hess_beta,
temp_LAML,
temp_LAML2,
Vp,
S_beta,
beta,
inverse_new_S,
X,
X_Q,
temp_deriv3,
temp_deriv4,
event,
expected,
type,
Ve,
deriv_rho_Ve,
mat_temp,
deriv_mat_temp,
eigen_mat_temp,
method
```
Arguments

 \mathcal{L}

Value

Hessian matrix of LCV or LAML wrt rho

Description

Hessian matrix of LCV and LAML wrt rho (log smoothing parameters). Version for multiplicative decomposition : relative mortality ratio model

Usage

```
Hess_rho_mult(
 X_GL,
  X_GL_Q,
 GL_temp,
 haz_GL,
  deriv2_rho_beta,
  deriv_rho_beta,
  weights,
  tm,
  nb_smooth,
 p,
  n_legendre,
  deriv_rho_inv_Hess_beta,
  deriv_rho_Hess_unpen_beta,
  S_list,
  minus_eigen_inv_Hess_beta,
  temp_LAML,
  temp_LAML2,
  Vp,
  S_beta,
 beta,
  inverse_new_S,
 X,
 X_Q,
  event,
  expected,
  Ve,
  deriv_rho_Ve,
 mat_temp,
 deriv_mat_temp,
  eigen_mat_temp,
 method
```
Arguments

 $\overline{}$

Value

Hessian matrix of LCV or LAML wrt rho

Description

Returns the position of the nth occurrence of str2 in str1. Returns 0 if str2 is not found. This code was first suggested by Abdelmonem Mahmoud Amer in https://stackoverflow.com/a/33005653/5421090

Usage

```
instr(str1, str2, startpos = 1, n = 1)
```
Arguments

Value

number representing the nth position of str2 in str1

Examples

library(survPen)

```
instr("character test to find the position of the third letter r","r",n=3)
```


Reverses the initial reparameterization for stable evaluation of the log *determinant of the penalty matrix*

Description

Transforms the final model by reversing the initial reparameterization performed by [repam](#page-37-1). Derives the corrected version of the Bayesian covariance matrix

Usage

inv.repam(model, X.ini, S.pen.ini)

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Arguments

Value

survPen object with standard parameterization

list.wicss *List of ICSS standards for age-standardization of cancer (net) survival*

Description

Four data frames are available in the list : 1, 2, 3 and "prostate". Each one corresponds to certain types of cancer. Details can be found in Corazzieri et al. (2004) (10.1016/j.ejca.2004.07.002) or at (in French) : https://www.santepubliquefrance.fr/docs/survie-des-personnes-atteintes-de-canceren-france-metropolitaine-1989-2018-materiel-et-methodes For each data frame, the variables are as follows:

- AgeClass. Age classes considered. Closed on the left and open on the right.
- AgeWeights. Weights associated with each age class

Usage

data(list.wicss)

Format

A list containing four data frames of 5 rows and 2 variables each

model.cons *Design and penalty matrices for the model*

Description

Sets up the model before optimization. Builds the design matrix, the penalty matrix and all the design matrices needed for Gauss-Legendre quadrature.

model.cons 23

Usage

```
model.cons(
  formula,
  lambda,
  data.spec,
  t1,
  t1.name,
  t0,
  t0.name,
  event,
  event.name,
  expected,
  expected.name,
  type,
  n.legendre,
  cl,
  beta.ini
\mathcal{L}
```
Arguments

Value

List of objects with the following items:

NR.beta 25

Examples

library(survPen)

standard spline of time with 4 knots data <- data.frame(time=seq(0,5,length=100),event=1,t0=0) form \leq \sim smf(time, knots=c(0,1,3,5)) t1 <- eval(substitute(time), data) t0 <- eval(substitute(t0), data) event <- eval(substitute(event), data) # The following code sets up everything we need in order to fit the model model.c <- model.cons(form,lambda=0,data.spec=data,t1=t1,t1.name="time", t0=rep(0,100),t0.name="t0",event=event,event.name="event", expected=rep(0,100),expected.name=NULL,type="overall",n.legendre=20, cl="survPen(form,data,t1=time,event=event)",beta.ini=NULL)

NR.beta *Inner Newton-Raphson algorithm for regression parameters estimation*

Description

Applies Newton-Raphson algorithm for beta estimation. Two specific modifications aims at guaranteeing convergence : first the hessian is perturbed whenever it is not positive definite and second, at each step, if the penalized log-likelihood is not maximized, the step is halved until it is.

Usage

```
NR.beta(build, beta.ini, detail.beta, max.it.beta = 200, tol.beta = 1e-04)
```
Arguments

Details

If we note 11.pen and beta respectively the current penalized log-likelihood and estimated parameters and 11. pen. old and betaold the previous ones, the algorithm goes on while (abs(ll.penll.pen.old)>tol.beta) or any(abs((beta-betaold)/betaold)>tol.beta)

Value

List of objects:

Examples

library(survPen)

standard spline of time with 4 knots

data <- data.frame(time=seq(0,5,length=100),event=1,t0=0)

```
form \leq \sim smf(time, knots=c(0,1,3,5))
```

```
t1 <- eval(substitute(time), data)
t0 <- eval(substitute(t0), data)
event <- eval(substitute(event), data)
```

```
# Setting up the model before fitting
model.c <- model.cons(form,lambda=0,data.spec=data,t1=t1,t1.name="time",
t0=rep(0,100),t0.name="t0",event=event,event.name="event",
```
NR.rho 27

```
expected=rep(0,100),expected.name=NULL,type="overall",n.legendre=20,
cl="survPen(form,data,t1=time,event=event)",beta.ini=NULL)
```

```
# Estimating the regression parameters at given smoothing parameter (here lambda=0)
Newton1 <- NR.beta(model.c,beta.ini=rep(0,4),detail.beta=TRUE)
```
NR.rho *Outer Newton-Raphson algorithm for smoothing parameters estimation via LCV or LAML optimization*

Description

Applies Newton-Raphson algorithm for smoothing parameters estimation. Two specific modifications aims at guaranteeing convergence : first the hessian is perturbed whenever it is not positive definite and second, at each step, if LCV or -LAML is not minimized, the step is halved until it is.

Usage

```
NR.rho(
  build,
  rho.ini,
  data,
  formula,
  max.it.beta = 200,max.it.rho = 30,
  beta.ini = NULL,
  detail.rho = FALSE,
  detail.beta = FALSE,
  nb.smooth,
  tol.beta = 1e-04,
  tol.rho = 1e-04,step.max = 5,
  method = "LAML")
```


Details

If we note val the current LCV or LAML value, val. old the previous one and grad the gradient vector of LCV or LAML with respect to the log smoothing parameters, the algorithm goes on while(abs(val-val.old)>tol.rho|any(abs(grad)>tol.rho))

Value

object of class survPen (see [survPen.fit](#page-59-1) for details)

Examples

```
library(survPen)
# standard spline of time with 4 knots
data <- data.frame(time=seq(0,5,length=100),event=1,t0=0)
form \leq \sim smf(time, knots=c(0,1,3,5))
t1 <- eval(substitute(time), data)
t0 <- eval(substitute(t0), data)
event <- eval(substitute(event), data)
# Setting up the model before fitting
model.c <- model.cons(form,lambda=0,data.spec=data,t1=t1,t1.name="time",
t0=rep(0,100),t0.name="t0",event=event,event.name="event",
expected=0,expected.name=NULL,type="overall",n.legendre=20,
cl="survPen(form,data,t1=time,event=event)",beta.ini=NULL)
```
Estimating the smoothing parameter and the regression parameters

```
# we need to apply a reparameterization to model.c before fitting
constructor <- repam(model.c)$build # model constructor
constructor$optim.rho <- 1 # we tell it we want to estimate the log smoothing parameters (rho)
Newton2 <- NR.rho(constructor,rho.ini=-1,data,form,nb.smooth=1,detail.rho=TRUE)
```
predict.survPen *Hazard and Survival prediction from fitted* survPen *model*

Description

Takes a fitted survPen object and produces hazard and survival predictions given a new set of values for the model covariates.

Usage

```
## S3 method for class 'survPen'
predict(
 object,
  newdata,
  newdata.ref = NULL,
  n.legendre = 50,
  conf.int = 0.95,
  do.surv = TRUE,type = "standard",
  exclude.random = FALSE,
  get.deriv.H = FALSE,
  ...
)
```


Details

The confidence intervals noted CI.U are built on the log cumulative hazard scale U=log(H) (efficient scale in terms of respect towards the normality assumption) using Delta method. The confidence intervals on the survival scale are then CI.surv = exp(-exp(CI.U))

Value

List of objects:

References

Wood, S.N., Pya, N. and Saefken, B. (2016), Smoothing parameter and model selection for general smooth models (with discussion). Journal of the American Statistical Association 111, 1548-1575

Examples

```
library(survPen)
data(datCancer) # simulated dataset with 2000 individuals diagnosed with cervical cancer
f1 <- \simtensor(fu,age,df=c(5,5))
# hazard model
mod1 <- survPen(f1,data=datCancer,t1=fu,event=dead,expected=NULL,method="LAML")
# predicting hazard and survival curves for age 60
nt <- seq(0,5,le=50)
pred <- predict(mod1,data.frame(fu=nt,age=60))
pred$haz
pred$surv
# predicting hazard ratio at 1 year according to age (with reference age of 50)
newdata1 <- data.frame(fu=1,age=seq(30,90,by=1))
newdata.ref1 <- data.frame(fu=1,age=rep(50,times=61))
predHR_1 <- predict(mod1,newdata=newdata1,newdata.ref=newdata.ref1,type="HR")
predHR_1$HR
predHR_1$HR.inf
predHR_1$HR.sup
# predicting hazard ratio at 3 years according to age (with reference age of 50)
# and difference of survival at 3 years
```
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```
newdata3 <- data.frame(fu=3,age=seq(30,90,by=1))
newdata.ref3 <- data.frame(fu=3,age=rep(50,times=61))
predHR_3 <- predict(mod1,newdata=newdata3,newdata.ref=newdata.ref3,type="HR")
# Hazard ratio
predHR_3$HR
predHR_3$HR.inf
predHR_3$HR.sup
# Difference of survival
predHR_3$diff.surv
predHR_3$diff.surv.inf
predHR_3$diff.surv.sup
```
predSNS *Prediction of grouped indicators : population (net) survival (PNS) and age-standardized (net) survival (SNS)*

Description

Allows the prediction of population and age-standardized (net) survival as well as associated confidence intervals

Usage

```
predSNS(
  model,
  time.points,
  newdata,
  weight.table,
  var.name,
  var.model,
  conf.int = 0.95,
  method = "exact",
 n.legendre = 50
)
```


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Details

The weight table used should always be in the same format as elements of [list.wicss](#page-21-2). Only agestandardization is possible for now. All other variables necessary for model predictions should be fixed to a single value. For simplicity, in what follows we will consider that survival only depends on time and age.

Value

List of nine elements

Population Net Survival (PNS)

For a given group of individuals, PNS at time t is defined as

$$
PNS(t) = \sum_{i} 1/n * S_i(t, a_i)
$$

where a_i is the age of individual i

Standardized Net Survival (SNS)

SNS at time t is defined as

$$
SNS(t) = \sum_{i} w_i * S_i(t, a_i)
$$

where a_i is the age of individual i and $w_i = w_{refj(i)}/n_{j(i)}$. $w_{refj(i)}$ is the weigth of age class j in the reference population (it corresponds to weight.table\$AgeWeights). Where $n_{j(i)}$ is the total number of individuals present in age class $j(i)$: the age class of individual i.

Standardized Net Survival (SNS) with method="approx"

For large datasets, SNS calculation is quite heavy. To reduce computational cost, the idea is to regroup individuals who have similar age values. By using floor(age) $+0.5$ instead of age, the gain will be substantial while the prediction error will be minimal (method="approx" will give slightly different predictions compared to method="exact"). Of course, if the provided age values are whole numbers then said provided age values will be used directly for grouping and there will be no prediction error (method="approx" and method="exact" will give the exact same predictions).

$$
SNS(t) = \sum_{a} \tilde{w}_a * S(t, a)
$$

The sum is here calculated over all possible values of age instead of all individuals. We have $\tilde{w}_a = n_a * w_{refj(a)}/n_{j(a)}$. Where $j(a)$ is the age class of age a while n_a is the number of individuals with age a .

Variance and Confidence Intervals

Confidence intervals for SNS are derived assuming normality of log(log(-SNS)) Lower and upper bound are given by

$$
IC_{95\%}(SNS) = [SNS^{1.96*}\sqrt(Var(Log(Deltas_{NS})))}; SNS^{-1.96*}\sqrt(Var(Log(Deltas_{NS})))]
$$

with

$$
Delta_{SNS} = -log(SNS)
$$

 $Var(Log(Delta_{SNS}))$ is derived by Delta method.

Confidence intervals for PNS are derived in the exact same way.

References

Corazziari, I., Quinn, M., & Capocaccia, R. (2004). Standard cancer patient population for age standardising survival ratios. European journal of cancer (Oxford, England : 1990), 40(15), 2307–2316. https://doi.org/10.1016/j.ejca.2004.07.002.

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Examples

```
data(datCancer)
data(list.wicss)
don <- datCancer
don$agec <- don$age - 50 # using centered age for modelling
#-------------------- model with time and age
knots.t<-quantile(don$fu[don$dead==1],probs=seq(0,1,length=6)) # knots for time
knots.agec<-quantile(don$agec[don$dead==1],probs=seq(0,1,length=5)) # knots for age
formula <- as.formula(~tensor(fu,agec,df=c(length(knots.t),length(knots.agec)),
knots=list(fu=knots.t,age=knots.agec)))
mod <- survPen(formula,data=don,t1=fu,event=dead,n.legendre=20, expected=rate)
#-------------------- Age classes and associated weights for age-standardized
# net survival prediction
# weights of type 1
wicss <- list.wicss[["1"]]
# to estimate population net survival, prediction dataframe
# is needed. It should contain original data for age
pred.pop <- data.frame(age=don$age)
#-------------------- prediction : age-standardized net survival and population net survival
pred <- predSNS(mod,time.points=seq(0,5,by=0.1),newdata=pred.pop,
weight.table=wicss,var.name=list(agec="age"),
var.model=list(agec=function(age) age - 50), method="approx")
```
print.summary.survPen *print summary for a* survPen *fit*

Description

print summary for a survPen fit

Usage

```
## S3 method for class 'summary.survPen'
print(
```

```
x,
  digits = max(3, getOption("digits") - 2),signif.stars = getOption("show.signif.stars"),
  ...
)
```
Arguments

Value

print of summary

pwcst *Defining piecewise constant (excess) hazard in survPen formulae*

Description

Used inside a formula object to define a piecewise constant (excess) hazard. This is useful since it triggers an explicit calculation of cumulative hazard calculation (much more efficient and more precise than Gauss-Legendre quadrature when hazard is constant). The breaks given are used to defined sub-intervals that are left-open (except the first interval which is always left-closed) and right-closed. Internally, this constructor uses the cut function on the follow-up time with options include.lowest=TRUE and right=TRUE Important : this function must not be used with other timedependent effect functions because the Gauss-Legendre quadrature will not operate correctly. If you really want to fit such a model, please use the cut function with the time variable as an argument to fit a piecewise constant hazard (and do not forget to use a huge number of Gauss-Legendre quadrature nodes, typically n.legendre=500)

Usage

pwcst(breaks)

Arguments

breaks numeric vector that specifies the boundaries of each sub-interval on which the hazard is constant

Value

object of class pwcst.spec

pwcst.breaks numeric vector that specifies the boundaries of each sub-interval on which the hazard is constant

Examples

library(survPen)

data(datCancer)

```
# piece constant hazard on 6 sub-intervals : [0;0.5]; ]0.5;1]; ]1;2]; ]2;3]; ]3;4]; ]4;5]
formula <- ~pwcst(breaks=c(0,0.5,1,2,3,4,5))
mod <- survPen(formula,t1=fu,event=dead,data=datCancer)
```

```
# The same but in an inefficient way
formula2 <- ~cut(fu,breaks=c(0,0.5,1,2,3,4,5),include.lowest=TRUE,right=TRUE)
mod.inefficient <- survPen(formula2,t1=fu,event=dead,data=datCancer,n.legendre=500)
```
rd *Defining random effects in survPen formulae*

Description

Used inside a formula object to define a random effect.

Usage

rd(...)

Arguments

... Any number of covariates separated by ","

Value

object of class rd.smooth.spec

Examples

cubic regression spline of time with 10 unspecified knots + random effect at the cluster level formula.test <- ~smf(time,df=10) + rd(cluster)

rd 37

38 repamate the contract of th

repam *Applies initial reparameterization for stable evaluation of the log determinant of the penalty matrix*

Description

Transforms the object from [model.cons](#page-21-1) by applying the matrix reparameterization (matrix U.F). The reparameterization is reversed at convergence by inv. repam.

Usage

repam(build)

Arguments

Value

Examples

re.model.c <- repam(model.c)

```
library(survPen)
# standard spline of time with 4 knots
data <- data.frame(time=seq(0,5,length=100),event=1,t0=0)
form \leq \sim smf(time, knots=c(0,1,3,5))
t1 <- eval(substitute(time), data)
t0 <- eval(substitute(t0), data)
event <- eval(substitute(event), data)
# Setting up the model before fitting
model.c <- model.cons(form,lambda=0,data.spec=data,t1=t1,t1.name="time",
t0=rep(0,100),t0.name="t0",event=event,event.name="event",
expected=rep(0,100),expected.name=NULL,type="overall",n.legendre=20,
cl="survPen(form,data,t1=time,event=event)",beta.ini=NULL)
# Reparameterization allows separating the parameters into unpenalized and
# penalized ones for maximum numerical stability
```


Description

Takes the model at convergence and calculates the robust variance matrix accounting for correlated survival times

Usage

robust.var(model, data, cluster.name, n.legendre = 50)

Arguments

Value

survPen object with robust variance Vr

smf *Defining smooths in survPen formulae*

Description

Used inside a formula object to define a smooth, a tensor product smooth or a tensor product interaction. Natural cubic regression splines (linear beyond the knots, equivalent to ns from package splines) are used as marginal bases. While tensor builds a tensor product of marginal bases including the intercepts, tint applies a tensor product of the marginal bases without their intercepts. Unlike tensor, the marginal effects of the covariates should also be present in the formula when using tint. For a conceptual difference between tensor products and tensor product interactions see Section 5.6.3 from Wood (2017)

Usage

```
smf(\ldots, knots = NULL, df = NULL, by = NULL, same. rho = FALSE)tensor(..., knots = NULL, df = NULL, by = NULL, same.rho = FALSE)tint(..., knots = NULL, df = NULL, by = NULL, same.rho = FALSE)
```
Arguments

Value

object of class smf.smooth.spec, tensor.smooth.spec or tint.smooth.spec (see [smooth.spec](#page-43-1) for details)

References

Wood, S. N. (2017), Generalized additive models: an introduction with R. Second Edition. London: Chapman & Hall/CRC.

Examples

penalized cubic regression spline of time with 5 unspecified knots formula.test <- ~smf(time,df=5)

suppose that we want to fit a model from formula.test library(survPen) data(datCancer)

mod.test <- survPen(~smf(fu,df=5) ,data=datCancer,t1=fu,event=dead)

then the knots can be retrieved like this: mod.test\$list.smf[[1]]\$knots # or calculated like this quantile(unique(datCancer\$fu),seq(0,1,length=5))

penalized cubic regression splines of time and age with respectively 5 and 7 unspecified knots formula.test2 <- ~smf(time,df=5)+smf(age,df=7)

penalized cubic regression splines of time and age with respectively 3 and 4 specified knots formula.test3 <- $\text{~smp}(\text{time},\text{knots}=(0,3,5))$ +smf(age,knots=c(30,50,70,90))

penalized tensor product for time and age with respectively 5 and 4 unspecified knots leading # to 5*4 = 20 regression parameters formula.test <- ~tensor(time,age,df=c(5,4))

penalized tensor product for time and age with respectively 3 and 4 specified knots

smooth.cons 41

formula.test3 <- \sim tensor(time,agec,knots=list(c(0,3,5),c(30,50,70,90)))

penalized tensor product for time, age and year with respectively 6, 5 and 4 unspecified knots formula.test <- ~tensor(time,age,year,df=c(6,5,4))

penalized tensor product interaction for time and age with respectively 5 and 4 unspecified knots # main effects are specified as penalized cubic regression splines formula.test <- ~smf(time,df=5)+smf(age,df=4)+tint(time,age,df=c(5,4))

Description

Builds the design and penalty matrices from the result of [smooth.spec](#page-43-1).

Usage

```
smooth.cons(
  term,
  knots,
  df,
  by = NULL,
 option,
  data.spec,
  same.rho = FALSE,
  name
)
```


Value

List of objects with the following items:

Examples

library(survPen)

```
# standard spline of time with 4 knots (so we get a design matrix with 3 columns
# because of centering constraint)
```

```
data <- data.frame(time=seq(0,5,length=100))
smooth.c <- smooth.cons("time",knots=list(c(0,1,3,5)),df=4,option="smf",
data.spec=data,name="smf(time)")
```


Description

Almost identical to [smooth.cons](#page-40-1). This version is dedicated to Gauss-Legendre quadrature. Here, the sum-to-zero constraints must be specified so that they correspond to the ones that were calculated with the initial dataset.

smooth.cons.integral 43

Usage

```
smooth.cons.integral(
  term,
  knots,
  df,
  by = NULL,
  option,
  data.spec,
  Z.smf,
  Z.tensor,
  Z.tint,
  name
\mathcal{L}
```
Arguments

Value

design matrix

Examples

library(survPen)

standard spline of time with 4 knots (so we get a design matrix with 3 columns # because of centering constraint)

```
data <- data.frame(time=seq(0,5,length=100))
# retrieving sum-to-zero constraint matrices
Z.smf \leq smooth.cons("time", knots=list(c(0,1,3,5)),df=4,option="smf",
data.spec=data,name="smf(time)")$Z.smf
# constructing the design matrices for Gauss-Legendre quadrature
smooth.c.int <- smooth.cons.integral("time",knots=list(c(0,1,3,5)),df=4,option="smf",data.spec=data,
name="smf(time)",Z.smf=Z.smf,Z.tensor=NULL,Z.tint=NULL)
```
smooth.spec *Covariates specified as penalized splines*

Description

Specifies the covariates to be considered as penalized splines.

Usage

```
smooth.spec(
  ...,
 knots = NULL,
 df = NULL,by = NULL,
 option = NULL,
 same.rho = FALSE
)
```


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Value

object of class smooth.spec

Examples

library(survPen)

standard spline of time with 10 unspecified knots smooth.spec(time)

```
# tensor of time and age with 5*5 specified knots
smooth.s <- smooth.spec(time,age,knots=list(time=seq(0,5,length=5),age=seq(20,80,length=5)),
option="tensor")
```


Description

This function allows splitting the original dataset in order to retrieve all the expected mortality rates available according to each individual's follow-up time. Typically, the expected mortality rates come from national mortality tables and values are available for every combination of age and year (often with 1-year increment).

Usage

```
splitmult(data, cut, start = NULL, end, event)
```


Details

This function is close to the survsplit function proposed in relsurv package, but it is simpler since fewer features are needed.

Value

split dataset with follow-up time split at specified times. An 'id_row' column is added to identify original row numbers

Examples

```
library(survPen)
data(datCancer)
data(expected.table)
#-------------------- creating split dataset for multiplicative model
splitdat <- splitmult(datCancer, cut = (1:5), end = "fu",
event = "dead")#-------------------- merging with expected mortality table
# deriving current age and year (closest whole number)
splitdat$age_current <- floor(splitdat$age + splitdat$fu + 0.5)
splitdat$year_current <- floor(splitdat$yod + splitdat$fu + 0.5)
splitdat <- merge(splitdat, expected.table,
                by.x=c("age_current","year_current"), by.y=c("Age","Year"),all.x=TRUE)
```
summary.survPen *Summary for a* survPen *fit*

Description

Takes a fitted survPen object and produces various useful summaries from it.

Usage

S3 method for class 'survPen' summary(object, ...)

Value

List of objects:

Examples

library(survPen)

data(datCancer) # simulated dataset with 2000 individuals diagnosed with cervical cancer

model : unidimensional penalized spline for time since diagnosis with 5 knots f1 \leq \sim smf(fu,df=5)

```
# fitting hazard model
mod1 <- survPen(f1,data=datCancer,t1=fu,event=dead,expected=NULL,method="LAML")
```
summary summary(mod1)

survPen *(Excess) hazard model with (multidimensional) penalized splines and integrated smoothness estimation*

Description

Fits an (excess) hazard model with (multidimensional) penalized splines allowing for time-dependent effects, non-linear effects and interactions between several continuous covariates. The linear predictor is specified on the logarithm of the (excess) hazard. Smooth terms are represented using cubic regression splines with associated quadratic penalties. For multidimensional smooths, tensor product splines or tensor product interactions are available. Smoothness is estimated automatically by optimizing one of two criteria: Laplace approximate marginal likelihood (LAML) or likelihood cross-validation (LCV). When specifying the model's formula, no distinction is made between the part relative to the form of the baseline hazard and the one relative to the effects of the covariates. Thus, time-dependent effects are naturally specified as interactions with some function of time via "*" or ":". See the examples below for more details. The main functions of the survPen package are [survPen](#page-47-1), [smf](#page-38-1), [tensor](#page-38-2), [tint](#page-38-2) and [rd](#page-36-1). The first one fits the model while the other four are constructors for penalized splines.

The user must be aware that the survPen package does not depend on mgcv. Thus, all the functionalities available in mgcv in terms of types of splines (such as thin plate regression splines or P-splines) are not available in survPen (yet).

Usage

```
survPen(
  formula,
  data,
  t1,
  t0 = NULL,event,
  expected = NULL,lambda = NULL,rho.ini = NULL,
 max.it.beta = 200,max.it.rho = 30,
 beta.ini = NULL,
  detail.rho = FALSE,
 detail.beta = FALSE,
  n.legendre = NULL,
 method = "LAML",tol.beta = 1e-04.
  tol.rho = 1e-04,step.max = 5.
  type = "overall",cluster = NULL
)
```
survPen 2008 and 200

Details

In time-to-event analysis, we may deal with one or several continuous covariates whose functional forms, time-dependent effects and interaction structure are challenging. One possible way to deal with these effects and interactions is to use the classical approximation of the survival likelihood by a Poisson likelihood. Thus, by artificially splitting the data, the package mgcv can then be used to fit penalized hazard models (Remontet et al. 2018). The problem with this option is that the setup is rather complex and the method can fail with huge datasets (before splitting). Wood et al. (2016) provided a general penalized framework that made available smooth function estimation to a wide variety of models. They proposed to estimate smoothing parameters by maximizing a Laplace approximate marginal likelihood (LAML) criterion and demonstrate how statistical consistency is maintained by doing so. The [survPen](#page-47-1) function implements the framework described by Wood et al. (2016) for modelling time-to-event data without requiring data splitting and Poisson likelihood approximation. The effects of continuous covariates are represented using low rank spline bases with associated quadratic penalties. The [survPen](#page-47-1) function allows to account simultaneously for time-dependent effects, non-linear effects and interactions between several continuous covariates without the need to build a possibly demanding model-selection procedure. Besides LAML, a likelihood cross-validation (LCV) criterion (O Sullivan 1988) can be used for smoothing parameter estimation. First and second derivatives of LCV with respect to the smoothing parameters are implemented so that LCV optimization is computationally equivalent to the LAML optimization proposed by Wood et al. (2016). In practice, LAML optimization is generally both a bit faster and a bit more stable so it is used as default. For m covariates (x_1, \ldots, x_m) , if we note $h(t, x_1, \ldots, x_m)$ the hazard at time t , the hazard model is the following :

$$
log[h(t, x_1, \ldots, x_m)] = \sum_j g_j(t, x_1, \ldots, x_m)
$$

where each g_i is either the marginal basis of a specific covariate or a tensor product smooth of any number of covariates. The marginal bases of the covariates are represented as natural (or restricted) cubic splines (as in function ns from library splines) with associated quadratic penalties. Full parametric (unpenalized) terms for the effects of covariates are also possible (see the examples below). Each g_i is then associated with zero, one or several smoothing parameters. The estimation procedure is based on outer Newton-Raphson iterations for the smoothing parameters and on inner Newton-Raphson iterations for the regression parameters (see Wood et al. 2016). Estimation of the regression parameters in the inner algorithm is by direct maximization of the penalized likelihood of the survival model, therefore avoiding data augmentation and Poisson likelihood approximation. The cumulative hazard included in the log-likelihood is approximated by Gauss-Legendre quadrature for numerical stability.

Value

Object of class "survPen" (see [survPenObject](#page-60-1) for details)

by variables

The [smf](#page-38-1), [tensor](#page-38-2) and [tint](#page-38-2) terms used to specify smooths accept an argument by. This by argument allows for building varying-coefficient models i.e. for letting smooths interact with factors or parametric terms. If a by variable is numeric, then its ith element multiples the ith row of the model matrix corresponding to the smooth term concerned. If a by variable is a factor then it generates an indicator vector for each level of the factor, unless it is an ordered factor. In the non-ordered case, the model matrix for the smooth term is then replicated for each factor level, and each copy has its rows multiplied by the corresponding rows of its indicator variable. The smoothness penalties are also duplicated for each factor level. In short a different smooth is generated for each factor level. The main interest of by variables over separated models is the same.rho argument (for [smf](#page-38-1), [tensor](#page-38-2) and [tint](#page-38-2)) which allows forcing all smooths to have the same smoothing parameter(s). Ordered by variables are handled in the same way, except that no smooth is generated for the first level of the ordered factor. This is useful if you are interested in differences from a reference level.

See the [survival_analysis_with_survPen vignette](../doc/survival_analysis_with_survPen.html) for more details.

Random effects

i.i.d random effects can be specified using penalization. Indeed, the ridge penalty is equivalent to an assumption that the regression parameters are i.i.d. normal random effects. Thus, it is easy to fit a frailty hazard model. For example, consider the model term rd(clust) which will result in a model matrix component corresponding to model.matrix(~clust-1) being added to the model matrix for the whole model. The associated regression parameters are assumed i.i.d. normal, with unknown variance (to be estimated). This assumption is equivalent to an identity penalty matrix (i.e. a ridge penalty) on the regression parameters. The unknown smoothing parameter λ associated with the term rd(clust) is directly linked to the unknown variance σ^2 : $\sigma^2 = \frac{1}{\lambda * S. scale}$. Then, the estimated log standard deviation is: $log(\hat{\sigma}) = -0.5 * log(\hat{\lambda}) - 0.5 * log(S-scale)$. And the estimated variance of the log standard deviation is: $Var[log(\hat{\sigma})] = 0.25 * Var[log(\hat{\lambda})] = 0.25 * inv. Hess.rho.$ See the [survival_analysis_with_survPen vignette](../doc/survival_analysis_with_survPen.html) for more details.

This approach allows implementing commonly used random effect structures. For example if g is a factor then $rd(g)$ produces a random parameter for each level of g , the random parameters being i.i.d. normal. If g is a factor and x is numeric, then $rd(g, x)$ produces an i.i.d. normal random slope relating the response to x for each level of g. Thus, random effects treated as penalized splines allow specifying frailty (excess) hazard models (Charvat et al. 2016). For each individual i from cluster (usually geographical unit) j, a possible model would be:

$$
log[h(t_{ij}, x_{ij1}, \dots, x_{ijm})] = \sum_{k} g_k(t_{ij}, x_{ij1}, \dots, x_{ijm}) + w_j
$$

where w_{-j} follows a normal distribution with mean 0. The random effect associated with the cluster variable is specified with the model term rd(cluster). We could also specify a random effect depending on age for example with the model term $rd(cluster, age)$. $u_j = exp(w_j)$ is known as the shared frailty.

See the [survival_analysis_with_survPen vignette](../doc/survival_analysis_with_survPen.html) for more details.

Excess hazard model

When studying the survival of patients who suffer from a common pathology we may be interested in the concept of excess mortality that represents the mortality due to that pathology. For example, in cancer epidemiology, individuals may die from cancer or from another cause. The problem is that the cause of death is often either unavailable or unreliable. Supposing that the mortality due to other causes may be obtained from the total mortality of the general population (called expected mortality for cancer patients), we can define the concept of excess mortality. The excess mortality is directly linked to the concept of net survival, which would be the observed survival if patients could not die from other causes. Therefore, when such competing events are present, one may choose to fit an excess hazard model instead of a classical hazard model. Flexible excess hazard models have already been proposed (for examples see Remontet et al. 2007, Charvat et al. 2016) but none of them deals with a penalized framework (in a non-fully Bayesian setting). Excess mortality can be estimated supposing that, in patients suffering from a common pathology, mortality due to others causes than the pathology can be obtained from the (all cause) mortality of the general population; the latter is referred to as the expected mortality h_P . The mortality observed in the patients (h_O) is actually decomposed as the sum of h_P and the excess mortality due to the pathology (h_E). This may be written as:

$$
h_O(t, x) = h_E(t, x) + h_P(a + t, z)
$$

In that equation, t is the time since cancer diagnosis, a is the age at diagnosis, h_P is the mortality of the general population at age $a + t$ given demographical characteristics z (h_P is considered known and available from national statistics), and x a vector of variables that may have an effect on h_E . Including the age in the model is necessary in order to deal with the informative censoring due to other causes of death. Thus, for m covariates (x_1, \ldots, x_m) , if we note $h_E(t, x_1, \ldots, x_m)$ the excess hazard at time t , the excess hazard model is the following:

$$
log[h_E(t, x_1, \ldots, x_m)] = \sum_j g_j(t, x_1, \ldots, x_m)
$$

Relative mortality ratio model

Another important feature of the survPen package is that it allows fitting penalized relative mortality ratio models.

As we discussed above, the excess mortality setting considers that the mortality (all causes) observed in the patients (h_O) is actually decomposed as the sum of the expected mortality h_P and the excess mortality due to the pathology (h_E) .

This may be written as:

$$
h_O(t, x) = h_E(t, x) + h_P(a + t, z)
$$

One limitation of such a decomposition is that h_E is considered positive. Indeed, sometimes this assumption is not met. For example, in prostate cancer patients with low stages at diagnosis, we observe an 'undermortality' due to selection effects and better overall medical care. In that case, the excess mortality is actually neagtive and the net survival setting fails to describe the reality of those patients. Besides, the excess mortality setting considers the studied disease as an independent cause of death (conditionally on the covariates) compared to the other causes. This point of view is not usely considered in multiple sclerosis epidemiology for example, where the disease is seen as a comorbidity impacting all pre- existing causes of death. In that case, the observed hazard is decomposed as product of population hazard and a relative mortality ratio r

This may be written as:

$$
h_O(t, x) = r(t, x) * h_P(a + t, z)
$$

This decomposition was first proposed in a modelling framework by Andersen et al. (1985). However Andersen's model was a non-flexible semi-parametric model.

The survPen package allows modelling the relative mortality ratio r as a multidimensional function of time and covariates. For m covariates (x_1, \ldots, x_m) , if we note $r(t, x_1, \ldots, x_m)$ the relative mortality ratio at time t , the model is as follows:

$$
log[r(t, x_1, \dots, x_m)] = \sum_j g_j(t, x_1, \dots, x_m)
$$

Where the g_j functions may be penalized unidimensional or penalized tensor product splines. All features described for the (excess) hazard setting still apply when fitting a relative mortality ratio model. One difference lies in the predictions. With a fitted relative mortality ratio model, you can only retrieve the relative mortality ratio and cumulative relative mortality ratio predictions (with CIs), as well as the ratios of realtive mortality ratio (with type='HR'). No survival prediction (let alone survival difference) will be directly available because its calculation depends on expected mortality rates.

Finally, one important difference between an excess hazard model and relative mortality ratio model is data preparation. For an excess hazard model we only need individual data with expected mortality rate at the time of death. Whereas in a relative mortality ratio model, the contribution to an individual to the likelihood requires all possible expected mortality rate values during the entire follow-up. Therefore, since the expected mortality rates come from national mortality tables usually available in 1-year intervals, we need to split the original dataset as many times as there are 1-year intervals during each individual's follow-up. The function [splitmult](#page-44-1) will help you getting the splitdataset from the original one.

See the [survival_analysis_with_survPen vignette](../doc/survival_analysis_with_survPen.html) for more details and an example of analysis.

Marginal hazard (intensity) models with robust standard errors

In presence of correlated time-to-event data (for example recurrent event data), robust standard errors accounting for said correlation need to be derived. The 'survPen' package allows deriving such robust standard errors based on sandwich estimators (often called Huber sandwich estimator, see also Coz et al. submitted to Biostatistics, for an example in the recurrent event setting).

The user only needs to specify the 'cluster' variable defining the statistical units for which repeated observations are available. This specification is performed via the 'cluster' argument.

See the survival analysis with survPen vignette for more details and an example of analysis.

Convergence

No convergence indicator is given. If the function returns an object of class survPen, it means that the algorithm has converged. If convergence issues occur, an error message is displayed. If convergence issues occur, do not refrain to use detail.rho and/or detail.beta to see exactly what is going on in the optimization process. To achieve convergence, consider lowering step.max and/or changing rho.ini and beta.ini. If your excess hazard model fails to converge, consider fitting a hazard model and use its estimated parameters as initial values for the excess hazard model. Finally, do not refrain to change the "method" argument (LCV or LAML) if convergence issues occur.

Other

Be aware that all character variables are transformed to factors before fitting.

References

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Coz, E., Charvat, H., Maucort-Boulch, D., and Fauvernier, M. (submitted to Biostatistics). Flexible penalized marginal intensity models for recurrent event data. Fauvernier, M., Roche, L., Uhry, Z., Tron, L., Bossard, N., Remontet, L. and the CENSUR Working Survival Group. Multidimensional penalized hazard model with continuous covariates: applications for studying trends and social inequalities in cancer survival, in revision in the Journal of the Royal Statistical Society, series C.

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Remontet, L., Bossard, N., Belot, A., & Esteve, J. (2007), An overall strategy based on regression models to estimate relative survival and model the effects of prognostic factors in cancer survival studies. Statistics in medicine, 26(10), 2214-2228.

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Wood, S.N., Pya, N. and Saefken, B. (2016), Smoothing parameter and model selection for general smooth models (with discussion). Journal of the American Statistical Association 111, 1548-1575

Examples

```
library(survPen)
data(datCancer) # simulated dataset with 2000 individuals diagnosed with cervical cancer
#-------------------------------------------------------- example 0
# Comparison between restricted cubic splines and penalized restricted cubic splines
library(splines)
# unpenalized
f \leq -\gammans(fu, knots=c(0.25, 0.5, 1, 2, 4), Boundary.knots=c(0,5))
mod <- survPen(f,data=datCancer,t1=fu,event=dead)
```

```
# penalized
f.pen \leq ~ smf(fu, knots=c(0,0.25, 0.5, 1, 2, 4,5)) # careful here: the boundary knots are included
mod.pen <- survPen(f.pen,data=datCancer,t1=fu,event=dead)
# predictions
new.time \leq seq(0,5,length=100)
pred <- predict(mod,data.frame(fu=new.time))
pred.pen <- predict(mod.pen,data.frame(fu=new.time))
par(mfrow=c(1,1))
plot(new.time,pred$haz,type="l",ylim=c(0,0.2),main="hazard vs time",
xlab="time since diagnosis (years)",ylab="hazard",col="red")
lines(new.time,pred.pen$haz,col="blue3")
legend("topright",legend=c("unpenalized","penalized"),
col=c("red","blue3"),lty=rep(1,2))
#-------------------------------------------------------- example 1
# hazard models with unpenalized formulas compared to a penalized tensor product smooth
library(survPen)
data(datCancer) # simulated dataset with 2000 individuals diagnosed with cervical cancer
# constant hazard model
f.cst \leq -1mod.cst <- survPen(f.cst,data=datCancer,t1=fu,event=dead)
# piecewise constant hazard model
f.pwcst \leq \sim \text{pwest}(\text{breaks}=\text{seq}(0,5,\text{by}=0.5))mod.pwcst <- survPen(f.pwcst,data=datCancer,t1=fu,event=dead)
# linear effect of time
f.lin \leq \sim \leqmod.lin <- survPen(f.lin,data=datCancer,t1=fu,event=dead)
# linear effect of time and age with proportional effect of age
f.lin.age <- ~fu+age
mod.lin.age <- survPen(f.lin.age,data=datCancer,t1=fu,event=dead)
# linear effect of time and age with time-dependent effect of age (linear)
f.lin.inter.age <- ~fu*age
mod.lin.inter.age <- survPen(f.lin.inter.age,data=datCancer,t1=fu,event=dead)
# cubic B-spline of time with a knot at 1 year, linear effect of age and time-dependent effect
# of age with a quadratic B-spline of time with a knot at 1 year
library(splines)
f.spline.inter.age <- ~bs(fu,knots=c(1),Boundary.knots=c(0,5))+age+
age:bs(fu,knots=c(1),Boundary.knots=c(0,5),degree=2)
# here, bs indicates an unpenalized cubic spline
```

```
mod.spline.inter.age <- survPen(f.spline.inter.age,data=datCancer,t1=fu,event=dead)
# tensor of time and age
f.tensor <- ~tensor(fu,age)
mod.tensor <- survPen(f.tensor,data=datCancer,t1=fu,event=dead)
# predictions of the models at age 60
new.time <- seq(0,5,length=100)
pred.cst <- predict(mod.cst,data.frame(fu=new.time))
pred.pwcst <- predict(mod.pwcst,data.frame(fu=new.time))
pred.lin <- predict(mod.lin,data.frame(fu=new.time))
pred.lin.age <- predict(mod.lin.age,data.frame(fu=new.time,age=60))
pred.lin.inter.age <- predict(mod.lin.inter.age,data.frame(fu=new.time,age=60))
pred.spline.inter.age <- predict(mod.spline.inter.age,data.frame(fu=new.time,age=60))
pred.tensor <- predict(mod.tensor,data.frame(fu=new.time,age=60))
1wd1 < -2par(mfrow=c(1,1))
plot(new.time,pred.cst$haz,type="l",ylim=c(0,0.2),main="hazard vs time",
xlab="time since diagnosis (years)",ylab="hazard",col="blue3",lwd=lwd1)
segments(x0=new.time[1:99],x1=new.time[2:100],y0=pred.pwcst$haz[1:99],col="lightblue2",lwd=lwd1)
lines(new.time,pred.lin$haz,col="green3",lwd=lwd1)
lines(new.time,pred.lin.age$haz,col="yellow",lwd=lwd1)
lines(new.time,pred.lin.inter.age$haz,col="orange",lwd=lwd1)
lines(new.time,pred.spline.inter.age$haz,col="red",lwd=lwd1)
lines(new.time,pred.tensor$haz,col="black",lwd=lwd1)
legend("topright",
legend=c("cst","pwcst","lin","lin.age","lin.inter.age","spline.inter.age","tensor"),
col=c("blue3","lightblue2","green3","yellow","orange","red","black"),
lty=rep(1,7),lwd=rep(lwd1,7))
# you can also calculate the hazard yourself with the lpmatrix option.
# For example, compare the following predictions:
haz.tensor <- pred.tensor$haz
X.tensor <- predict(mod.tensor,data.frame(fu=new.time,age=60),type="lpmatrix")
haz.tensor.lpmatrix <- exp(X.tensor%mult%mod.tensor$coefficients)
summary(haz.tensor.lpmatrix - haz.tensor)
#---------------- The 95% confidence intervals can be calculated like this:
# standard errors from the Bayesian covariance matrix Vp
std <- sqrt(rowSums((X.tensor%mult%mod.tensor$Vp)*X.tensor))
qt.norm <- stats::qnorm(1-(1-0.95)/2)
haz.inf <- as.vector(exp(X.tensor%mult%mod.tensor$coefficients-qt.norm*std))
haz.sup <- as.vector(exp(X.tensor%mult%mod.tensor$coefficients+qt.norm*std))
```

```
# checking that they are similar to the ones given by the predict function
summary(haz.inf - pred.tensor$haz.inf)
summary(haz.sup - pred.tensor$haz.sup)
#-------------------------------------------------------- example 2
library(survPen)
data(datCancer) # simulated dataset with 2000 individuals diagnosed with cervical cancer
# model : unidimensional penalized spline for time since diagnosis with 5 knots
f1 \leq \sim smf(fu,df=5)
# when knots are not specified, quantiles are used. For example, for the term "smf(x,df=df1)",
# the vector of knots will be: quantile(unique(x),seq(0,1,length=df1))
# you can specify your own knots if you want
# f1 <- ~smf(fu,knots=c(0,1,3,6,8))
# hazard model
mod1 <- survPen(f1,data=datCancer,t1=fu,event=dead,expected=NULL,method="LAML")
summary(mod1)
# to see where the knots were placed
mod1$list.smf
# with LCV instead of LAML
mod1bis <- survPen(f1,data=datCancer,t1=fu,event=dead,expected=NULL,method="LCV")
summary(mod1bis)
# hazard model taking into account left truncation (not representative of cancer data,
# the begin variable was simulated for illustration purposes only)
mod2 <- survPen(f1,data=datCancer,t0=begin,t1=fu,event=dead,expected=NULL,method="LAML")
summary(mod2)
# excess hazard model
mod3 <- survPen(f1,data=datCancer,t1=fu,event=dead,expected=rate,method="LAML")
summary(mod3)
# compare the predictions of the models
new.time \leq seq(0,5,length=50)
pred1 <- predict(mod1,data.frame(fu=new.time))
pred1bis <- predict(mod1bis,data.frame(fu=new.time))
pred2 <- predict(mod2,data.frame(fu=new.time))
pred3 <- predict(mod3,data.frame(fu=new.time))
# LAML vs LCV
par(mfrow=c(1,2))
plot(new.time,pred1$haz,type="l",ylim=c(0,0.2),main="LCV vs LAML",
xlab="time since diagnosis (years)",ylab="hazard")
lines(new.time,pred1bis$haz,col="blue3")
legend("topright",legend=c("LAML","LCV"),col=c("black","blue3"),lty=c(1,1))
```

```
plot(new.time,pred1$surv,type="l",ylim=c(0,1),main="LCV vs LAML",
xlab="time since diagnosis (years)",ylab="survival")
lines(new.time,pred1bis$surv,col="blue3")
# hazard vs excess hazard
par(mfrow=c(1,2))
plot(new.time,pred1$haz,type="l",ylim=c(0,0.2),main="hazard vs excess hazard",
xlab="time since diagnosis (years)",ylab="hazard")
lines(new.time,pred3$haz,col="green3")
legend("topright",legend=c("overall","excess"),col=c("black","green3"),lty=c(1,1))
plot(new.time,pred1$surv,type="l",ylim=c(0,1),main="survival vs net survival",
xlab="time",ylab="survival")
lines(new.time,pred3$surv,col="green3")
legend("topright",legend=c("overall survival","net survival"), col=c("black","green3"), lty=c(1,1))
# hazard vs excess hazard with 95% Bayesian confidence intervals (based on Vp matrix,
# see predict.survPen)
par(mfrow=c(1,1))
plot(new.time,pred1$haz,type="l",ylim=c(0,0.2),main="hazard vs excess hazard",
xlab="time since diagnosis (years)",ylab="hazard")
lines(new.time,pred3$haz,col="green3")
legend("topright",legend=c("overall","excess"),col=c("black","green3"),lty=c(1,1))
lines(new.time,pred1$haz.inf,lty=2)
lines(new.time,pred1$haz.sup,lty=2)
lines(new.time,pred3$haz.inf,lty=2,col="green3")
lines(new.time,pred3$haz.sup,lty=2,col="green3")
#-------------------------------------------------------- example 3
library(survPen)
data(datCancer) # simulated dataset with 2000 individuals diagnosed with cervical cancer
# models: tensor product smooth vs tensor product interaction of time since diagnosis and
# age at diagnosis. Smoothing parameters are estimated via LAML maximization
f2 <- \simtensor(fu,age,df=c(5,5))
f3 \le \le \le tint(fu,df=5)+tint(age,df=5)+tint(fu,age,df=c(5,5))
# hazard model
mod4 <- survPen(f2,data=datCancer,t1=fu,event=dead)
summary(mod4)
mod5 <- survPen(f3,data=datCancer,t1=fu,event=dead)
summary(mod5)
# predictions
```

```
new.age <- seq(50,90,length=50)
new.time \leq seq(0,7,length=50)
Z4 <- outer(new.time,new.age,function(t,a) predict(mod4,data.frame(fu=t,age=a))$haz)
Z5 <- outer(new.time,new.age,function(t,a) predict(mod5,data.frame(fu=t,age=a))$haz)
# color settings
col.pal <- colorRampPalette(c("white", "red"))
colors <- col.pal(100)
facet \leq function(z){
facet.center <- (z[-1, -1] + z[-1, -nco1(z)] + z[-nrow(z), -1] + z[-nrow(z), -nco1(z)])/4cut(facet.center, 100)
}
# plot the hazard surfaces for both models
par(mfrow=c(1,2))
persp(new.time,new.age,Z4,col=colors[facet(Z4)],main="tensor",theta=30,
xlab="time since diagnosis",ylab="age at diagnosis",zlab="excess hazard",ticktype="detailed")
persp(new.time,new.age,Z5,col=colors[facet(Z5)],main="tint",theta=30,
xlab="time since diagnosis",ylab="age at diagnosis",zlab="excess hazard",ticktype="detailed")
#-------------------------------------------------------- example 4
library(survPen)
data(datCancer) # simulated dataset with 2000 individuals diagnosed with cervical cancer
# model : tensor product spline for time, age and yod (year of diagnosis)
# yod is not centered here since it does not create unstability but be careful in practice
# and consider centering your covariates if you encounter convergence issues
f4 <- ~tensor(fu,age,yod,df=c(5,5,5))
# excess hazard model
mod6 <- survPen(f4,data=datCancer,t1=fu,event=dead,expected=rate)
summary(mod6)
# predictions of the surfaces for ages 50, 60, 70 and 80
new.year <- seq(1990,2010,length=30)
new.time <- seq(0,5,length=50)
Z_50 <- outer(new.time,new.year,function(t,y) predict(mod6,data.frame(fu=t,yod=y,age=50))$haz)
Z_60 <- outer(new.time,new.year,function(t,y) predict(mod6,data.frame(fu=t,yod=y,age=60))$haz)
Z_70 <- outer(new.time,new.year,function(t,y) predict(mod6,data.frame(fu=t,yod=y,age=70))$haz)
Z_80 <- outer(new.time,new.year,function(t,y) predict(mod6,data.frame(fu=t,yod=y,age=80))$haz)
# plot the hazard surfaces for a given age
par(mfrow=c(2,2))
persp(new.time,new.year,Z_50,col=colors[facet(Z_50)],main="age 50",theta=20,
```
xlab="time since diagnosis",ylab="yod",zlab="excess hazard",ticktype="detailed")

persp(new.time,new.year,Z_60,col=colors[facet(Z_60)],main="age 60",theta=20, xlab="time since diagnosis",ylab="yod",zlab="excess hazard",ticktype="detailed") persp(new.time,new.year,Z_70,col=colors[facet(Z_70)],main="age 70",theta=20, xlab="time since diagnosis",ylab="yod",zlab="excess hazard",ticktype="detailed") persp(new.time,new.year,Z_80,col=colors[facet(Z_80)],main="age 80",theta=20, xlab="time since diagnosis",ylab="yod",zlab="excess hazard",ticktype="detailed")

```
########################################
```
survPen.fit *(Excess) hazard model with multidimensional penalized splines for given smoothing parameters*

Description

Fits an (excess) hazard model. If penalized splines are present, the smoothing parameters are specified.

Usage

```
survPen.fit(
 build,
  data,
  formula,
 max.it.beta = 200,
 beta.ini = NULL,detail.beta = FALSE,
 method = "LAML",tol.beta = 1e-04\mathcal{E}
```


survPenObject 61

Value

Object of class "survPen" (see [survPenObject](#page-60-1) for details)

Examples

library(survPen)

```
# standard spline of time with 4 knots
data <- data.frame(time=seq(0,5,length=100),event=1,t0=0)
form \leq \sim smf(time, knots=c(0,1,3,5))
t1 <- eval(substitute(time), data)
t0 <- eval(substitute(t0), data)
event <- eval(substitute(event), data)
# Setting up the model before fitting
model.c <- model.cons(form,lambda=0,data.spec=data,t1=t1,t1.name="time",
t0=rep(0,100),t0.name="t0",event=event,event.name="event",
expected=rep(0,100),expected.name=NULL,type="overall",n.legendre=20,
cl="survPen(form,data,t1=time,event=event)",beta.ini=NULL)
# fitting
```
mod <- survPen.fit(model.c,data,form)

survPenObject *Fitted survPen object*

Description

A fitted survPen object returned by function [survPen](#page-47-1) and of class "survPen". Method functions predict and summary are available for this class.

Value

A survPen object has the following elements:

References

Wood, S.N., Pya, N. and Saefken, B. (2016), Smoothing parameter and model selection for general smooth models (with discussion). Journal of the American Statistical Association 111, 1548-1575

tensor.in *tensor model matrix for two marginal bases*

Description

Function called recursively inside [tensor.prod.X](#page-65-1).

Usage

```
tensor.in(X1, X2)
```
Arguments

Value

Matrix of dimensions $n*(p1*p2)$ representing the row tensor product of the matrices X1 and X2

tensor.prod.S 65

Examples

library(survPen)

row-wise tensor product between two design matrices set.seed(15) X1 <- matrix(rnorm(10*3),nrow=10,ncol=3) X2 <- matrix(rnorm(10*2),nrow=10,ncol=2)

tensor.in(X1,X2)

tensor.prod.S *Tensor product for penalty matrices*

Description

Computes the penalty matrices of a tensor product smooth from the marginal penalty matrices. The code is from function tensor.prod.penalties in mgcv package.

Usage

tensor.prod.S(S)

Arguments

Value

TS List of the penalty matrices associated with the tensor product smooth

Examples

library(survPen)

tensor product between three penalty matrices set.seed(15)

S1 <- matrix(rnorm(3*3),nrow=3,ncol=3) S2 <- matrix(rnorm(2*2),nrow=2,ncol=2)

 $S1 \leftarrow 0.5*(S1 + t(S1))$; $S2 \leftarrow 0.5*(S2 + t(S2))$

tensor.prod.S(list(S1,S2))

tensor.prod.X *tensor model matrix*

Description

Computes the model matrix of tensor product smooth from the marginal bases.

Usage

tensor.prod.X(X)

Arguments

X list of m design matrices with n rows and p1, p2, ... pm columns respectively

Value

T Matrix of dimensions $n*(p1*p2*x...*pm)$ representing the row tensor product of the matrices in X

Examples

library(survPen)

row-wise tensor product between three design matrices set.seed(15)

 $X1 \leftarrow \text{matrix}(rnorm(10*3), nrow=10, ncol=3)$ X2 <- matrix(rnorm(10*2),nrow=10,ncol=2) X3 <- matrix(rnorm(10*2),nrow=10,ncol=2) tensor.prod.X(list(X1,X2,X3))

%cross% *Matrix cross-multiplication between two matrices*

Description

Matrix cross-multiplication between two matrices

Usage

Mat1 %cross% Mat2

% mult% 67

Value

prod the product t(Mat1)

%mult% *Matrix multiplication between two matrices*

Description

Matrix multiplication between two matrices

Usage

Mat1 %mult% Mat2

Arguments

Value

prod the product Mat1

%vec% *Matrix multiplication between a matrix and a vector*

Description

Matrix multiplication between a matrix and a vector

Usage

Mat %vec% vec

Arguments

Value

prod the product Mat

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