

Package ‘HQM’

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Type Package

Title Superefficient Estimation of Future Conditional Hazards Based on Marker Information

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Description Provides a nonparametric smoothed kernel estimator for the future conditional hazard rate function when time-dependent covariates are present, a bandwidth selector for the estimator's implementation and pointwise and uniform confidence bands. Methods used in the package refer to Bagkavos, Isakson, Mammen, Nielsen and Proust-Lima (2025) <[doi:10.1093/biomet/asaf008](https://doi.org/10.1093/biomet/asaf008)>.

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auc.hqm

AUC for the High Quality Marker estimator

Description

Calculates the AUC for the HQM estimator.

Usage

```
auc.hqm(xin, est, landm, th, event_time_name, status_name)
```

Arguments

xin	A data frame containing event times and the patient status.
est	The HQM estimator values, typically the output of get_h_x .
landm	Landmark time.
th	Time horizon.
event_time_name	The column name of the event times in the xin data frame.
status_name	The column name of the status variable in the xin frame.

Details

The function `auc.hqm` implements the AUC calculation for the HQM estimator estimator.

Value

A vector of two values: the landmark time of the calculation and the AUC value.

See Also

`bs.hqm`

Examples

```
library(timeROC)
library(survival)
Landmark <- 2
pbcT1 <- pbc2[which(pbc2$year< Landmark & pbc2$years> Landmark),]
timesS2 <- seq(Landmark,14,by=0.5)
b=0.9
arg1<- get_h_x(pbcT1, 'albumin', event_time_name = 'years',
               time_name = 'year', event_name = 'status2', 2, 0.9)
br_s2 = seq(Landmark, 14, length=99)
sfalb2<- make_sf( (br_s2[2]-br_s2[1])/4 , arg1)
tHor <- 1.5
auc.hq.use<-auc.hqm(pbcT1, sfalb2, Landmark,tHor,
                    event_time_name = 'years', status_name = 'status2')
auc.hq.use
```

bs.hqm

Brier score for the High Quality Marker estimator

Description

Calculates the Brier score for the HQM estimator.

Usage

```
bs.hqm(xin, est, landm, th, event_time_name, status_name)
```

Arguments

<code>xin</code>	A data frame containing event times and the patient status.
<code>est</code>	The HQM estimator values, typically the output of <code>get_h_x</code> .
<code>landm</code>	Landmark time.
<code>th</code>	Time horizon.
<code>event_time_name</code>	The column name of the event times in the data frame <code>xin</code> .
<code>status_name</code>	The column name of the status variable in the data frame <code>xin</code> .

Details

The function `bs.hqm` implements the Brier score calculation for the HQM estimator estimator.

Value

Scalar: the Brier score of the HQM estimator.

See Also

[auc.hqm](#)

Examples

```
library(pec)
library(survival)
Landmark <- 2

pbcT1 <- pbc2[which(pbc2$year< Landmark & pbc2$years> Landmark),]
timesS2 <- seq(Landmark,14,by=0.5)

b=0.9
arg1<- get_h_x(pbcT1, 'albumin', event_time_name = 'years',
               time_name = 'year', event_name = 'status2', 2, 0.9)
br_s2 = seq(Landmark, 14, length=99)
sfalb2<- make_sf( (br_s2[2]-br_s2[1])/4 , arg1)

tHor <- 1.5
bs.use<-bs.hqm(pbcT1, sfalb2, Landmark,tHor,
               event_time_name = 'years', status_name = 'status2')
bs.use
```

b_selection

Cross validation bandwidth selection

Description

Implements the bandwidth selection for the future conditional hazard rate $\hat{h}_x(t)$ based on K-fold cross validation.

Usage

```
b_selection(data, marker_name, event_time_name = 'years',
            time_name = 'year', event_name = 'status2', I, b_list)
```

Arguments

data	A data frame of time dependent data points. Missing values are allowed.
marker_name	The column name of the marker values in the data frame data .
event_time_name	The column name of the event times in the data frame data .
time_name	The column name of the times the marker values were observed in the data frame data .
event_name	The column name of the events in the data frame data .
I	Number of observations leave out for a K cross validation.
b_list	Vector of bandwidths that need to be tested.

Details

The function [b_selection](#) implements the cross validation bandwidth selection for the future conditional hazard rate $\hat{h}_x(t)$ given by

$$b_{CV} = \operatorname{argmin}_b \sum_{i=1}^N \int_0^T \int_s^T Z_i(t) Z_i(s) (\hat{h}_{X_i(s)}(t-s) - h_{X_i(s)}(t-s))^2 dt ds,$$

where $\hat{h}_x(t)$ is a smoothed kernel density estimator of $h_x(t)$ and Z_i the exposure process of individual i . Note that $\hat{h}_x(t)$ is dependent on b .

Value

A list with the tested bandwidths and its cross validation scores.

See Also

[b_selection_prep_g](#), [Q1](#), [R_K](#), [prep_cv](#), [dataset_split](#)

Examples

```
I = 26
b_list = seq(0.9, 1.3, 0.1)

b_scores_alb = b_selection(pbc2, 'albumin', 'years', 'year', 'status2', I, b_list)
b_scores_alb[[2]][which.min(b_scores_alb[[1]])]
```

b_selection_prep_g	<i>Preparations for bandwidth selection</i>
--------------------	---

Description

Calculates an intermediate part for the K-fold cross validation.

Usage

```
b_selection_prep_g(h_mat, int_X, size_X_grid, n, Yi)
```

Arguments

h_mat	A matrix of the estimator for the future conditional hazard rate for all values x and t.
int_X	Vector of the position of the observed marker values in the grid for marker values.
size_X_grid	Numeric value indicating the number of grid points for marker values.
n	Number of individuals.
Yi	A matrix made by make_Yi indicating the exposure.

Details

The function [b_selection_prep_g](#) calculates a key component for the bandwidth selection

$$\hat{g}_i^{-I_j}(t) = \int_0^t Z_i(s) \hat{h}_{X_i(s)}^{-I_j}(t-s) ds,$$

where $\hat{h}^{-I_j}$ is estimated without information from all counting processes i with $i \in I_j$ and Z is the exposure.

Value

A matrix with $\hat{g}_i^{-I_j}(t)$ for all individuals i and time grid points t .

See Also

[b_selection](#)

Examples

```
pb2_id = to_id(pbc2)
size_s_grid <- size_X_grid <- 100
n = max(as.numeric(pbc2$id))
s = pbc2$year
X = pbc2$serBilir
XX = pb2_id$serBilir
```

```

ss <- pbc2_id$years
delta <- pbc2_id$status2
br_s = seq(0, max(s), max(s)/( size_s_grid-1))
br_X = seq(min(X), max(X), (max(X)-min(X))/( size_X_grid-1))

X_lin = lin_interpolate(br_s, pbc2_id$id, pbc2$id, X, s)

int_X <- findInterval(X_lin, br_X)
int_s = rep(1:length(br_s), n)

N <- make_N(pbc2, pbc2_id, breaks_X=br_X, breaks_s=br_s, ss, XX, delta)
Y <- make_Y(pbc2, pbc2_id, X_lin, br_X, br_s, size_s_grid, size_X_grid, int_s, int_X, 'years', n)

b = 1.7
alpha<-get_alpha(N, Y, b, br_X, K=Epan )

Yi <- make_Yi(pbc2, pbc2_id, X_lin, br_X, br_s, size_s_grid, size_X_grid, int_s, int_X, 'years', n)
Ni <- make_Ni(breaks_s=br_s, size_s_grid, ss, delta, n)

t = 2

h_xt_mat = t(sapply(br_s[1:99], function(si){
  h_xt_vec(br_X, br_s, size_s_grid, alpha, t, b, Yi, int_X, n)}))

b_selection_prep_g(h_xt_mat, int_X, size_X_grid, n, Yi)

```

Conf_bands

*Confidence bands***Description**

Implements the uniform and pointwise confidence bands for the future conditional hazard rate based on the last observed marker measure.

Usage

```

Conf_bands(data, marker_name, event_time_name = 'years',
           time_name = 'year', event_name = 'status2', x, b)

```

Arguments

data	A data frame of time dependent data points. Missing values are allowed.
marker_name	The column name of the marker values in the data frame data .
event_time_name	The column name of the event times in the data frame data .
time_name	The column name of the times the marker values were observed in the data frame data .
event_name	The column name of the events in the data frame data .
x	Numeric value of the last observed marker value.
b	Bandwidth.

Details

The function `Conf_bands` implements the pointwise and uniform confidence bands for the estimator of the future conditional hazard rate $\hat{h}_x(t)$. The confidence bands are based on a wild bootstrap approach $h_{x_*,B}^*(t)$.

Pointwise: For a given $t \in (0, T)$ generate $h_{x_*,B}^{*(1)}(t), \dots, h_{x_*,B}^{*(N)}(t)$ for $N = 1000$ and order it $h_{x_*,B}^{*[1]}(t) \leq \dots \leq h_{x_*,B}^{*[N]}(t)$. Then

$$\hat{I}_{n,N}^1 = \left[\hat{h}_{x_*}(t) - \hat{\sigma}_{G_{x_*}}(t) \frac{h_{x_*,B}^{*[N(1-\frac{\alpha}{2})]}(t)}{\sqrt{n}}, \hat{h}_{x_*}(t) + \hat{\sigma}_{G_{x_*}}(t) \frac{h_{x_*,B}^{*[N\frac{\alpha}{2}]}(t)}{\sqrt{n}} \right]$$

is a $1 - \alpha$ pointwise confidence band for $h_{x_*}(t)$, where $\hat{\sigma}_{G_{x_*}}(t)$ is a bootstrap estimate of the variance. For more details on the wild bootstrap approach, please see `prep_boot` and `g_xt`.

Uniform: Generate $\bar{h}_{x_*,B}^{(1)}(t), \dots, \bar{h}_{x_*,B}^{(N)}(t)$ for $N = 1000$ for all $t \in [\delta_T, T - \delta_T]$ and define $W^{(i)} = \sup_{t \in [0, T]} |\bar{h}_{x_*,B}^{(i)}(t)|$ for $i = 1, \dots, N$. Order $W^{[1]} \leq \dots \leq W^{[N]}$. Then

$$\hat{I}_{n,N}^2 = \left[\hat{h}_{x_*}(t) \pm \hat{\sigma}_{G_{x_*}}(t) \frac{W^{[N(1-\alpha)]}}{\sqrt{n}} \right]$$

is a $1 - \alpha$ uniform confidence band for $h_{x_*}(t)$.

Value

A list with pointwise, uniform confidence bands and the estimator $\hat{h}_x(t)$ for all possible time points t .

See Also

`g_xt`, `prep_boot`

Examples

```
b = 10
x = 3
size_s_grid <- 100
s = pbc2$year
br_s = seq(0, max(s), max(s)/(size_s_grid-1))

c_bands = Conf_bands(pbc2, 'serBilir', event_time_name = 'years',
                    time_name = 'year', event_name = 'status2', x, b)

J = 60
plot(br_s[1:J], c_bands$h_hat[1:J], type = "l", ylim = c(0,1), ylab = 'Hazard', xlab = 'Years')

lines(br_s[1:J], c_bands$I_p_up[1:J], col = "red")
lines(br_s[1:J], c_bands$I_p_do[1:J], col = "red")
lines(br_s[1:J], c_bands$I_nu[1:J], col = "blue")
lines(br_s[1:J], c_bands$I_nd[1:J], col = "blue")
```

dataset_split	<i>Split dataset for K-fold cross validation</i>
---------------	--

Description

Creates multiple splits of a dataset which is then used in the bandwidth selection with K-fold cross validation.

Usage

```
dataset_split(I, data)
```

Arguments

data	A data frame of time dependent data points. Missing values are allowed.
I	The number of individuals that should be left out. Optimally, $K = n/I$ should be an integer, where n is the number of individuals.

Details

The function `dataset_split` takes a data frame and transforms it into $K = n/I$ data frames with I individuals missing from each data frame. Let I_j be sets of indices with $\cup_{j=1}^K I_j = \{1, \dots, n\}$, $I_k \cap I_j = \emptyset$ and $|I_j| = |I_k| = I$ for all $j, k \in \{1, \dots, K\}$. Then data frames with $\{1, \dots, n\}/I_j$ individuals are created.

Value

A list of data frames with `I` individuals missing in the above way.

See Also

[b_selection](#)

Examples

```
splitted_dataset = dataset_split(26, pbc2)
```

dij	<i>D matrix entries, used for the implementation of the local linear kernel</i>
-----	---

Description

Calculates the entries of the D matrix in the definition of the local linear kernel

Usage

dij(b,x,y, K)

Arguments

x	A vector of design points where the kernel will be evaluated.
y	A vector of sample data points.
b	The bandwidth to use (a scalar).
K	The kernel function to use.

Details

Implements the calculation of all $d \times d$ entries of matrix D , which is part of the definition of the local linear kernel. The actual calculation is performed by

$$d_{jk} = \sum_{i=1}^n \int_0^T K_b(x - X_i(s)) \{x - X_{ij}(s)\} \{x - X_{ik}(s)\} Z_i(s) ds,$$

Value

scalar value, the result of d_{jk} .

Epan	<i>Epanechnikov kernel</i>
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Description

Implements the Epanechnikov kernel function

Usage

Epan(x)

Arguments

x	A vector of design points where the kernel will be evaluated.
---	---

Details

Implements the Epanechnikov kernel function

$$K(x) = \frac{3}{4}(1 - x^2) * (|x| < 1),$$

Value

Scalar, the value of the Epanechnikov kernel at x .

get_alpha	Marker-only hazard rate
-----------	-------------------------

Description

Calculates the marker-only hazard rate for time dependent data.

Usage

```
get_alpha(N, Y, b, br_X, K=Epan )
```

Arguments

N	A matrix made by <code>make_N</code> indicating the occurrences of events.
Y	A matrix made by <code>make_Y</code> indicating the exposure.
b	Bandwidth.
br_X	Vector of grid points for the marker values X .
K	Used kernel function.

Details

The function `get_alpha` implements the marker-only hazard estimator

$$\hat{\alpha}_i(z) = \frac{\sum_{k \neq i} \int_0^T K_{b_1}(z - X_k(s)) dN_k(s)}{\sum_{k \neq i} \int_0^T K_{b_1}(z - X_k(s)) Z_k(s) ds},$$

where X is the marker and Z is the exposure. The marker-only hazard is defined as the underlying hazard which is not dependent on time

$$\alpha(X(t), t) = \alpha(X(t))$$

Value

A vector of marker-only values for `br_X`.

See Also[h_xt](#)**Examples**

```

pbc2_id = to_id(pbc2)
size_s_grid <- size_X_grid <- 100
n = max(as.numeric(pbc2$id))
s = pbc2$year
X = pbc2$serBilir
XX = pbc2_id$serBilir
ss <- pbc2_id$years
delta <- pbc2_id$status2
br_s = seq(0, max(s), max(s)/( size_s_grid-1))
br_X = seq(min(X), max(X), (max(X)-min(X))/( size_X_grid-1))

X_lin = lin_interpolate(br_s, pbc2_id$id, pbc2$id, X, s)

int_X <- findInterval(X_lin, br_X)
int_s = rep(1:length(br_s), n)

N <- make_N(pbc2, pbc2_id, breaks_X=br_X, breaks_s=br_s, ss, XX, delta)
Y <- make_Y(pbc2, pbc2_id, X_lin, br_X, br_s, size_s_grid,
            size_X_grid, int_s, int_X, event_time = 'years', n)

b = 1.7
alpha<-get_alpha(N, Y, b, br_X, K=Epan )

```

get_h_x

Local constant future conditional hazard rate estimator

Description

Calculates the local constant future hazard rate function, conditional on a marker value x , across across a set of time values t .

Usage

```
get_h_x(data, marker_name, event_time_name, time_name, event_name, x, b)
```

Arguments

data	A data frame of time dependent data points. Missing values are allowed.
marker_name	The column name of the marker values in the data frame data .
event_time_name	The column name of the event times in the data frame data .
time_name	The column name of the times the marker values were observed in the data frame data .

event_name	The column name of the events in the data frame data .
x	Numeric value of the last observed marker value.
b	Bandwidth parameter.

Details

The function [get_h_x](#) implements the future local constant conditional hazard estimator

$$\hat{h}_x(t) = \frac{\sum_{i=1}^n \int_0^T \hat{\alpha}_i(X_i(t+s)) Z_i(t+s) Z_i(s) K_b(x - X_i(s)) ds}{\sum_{i=1}^n \int_0^T Z_i(t+s) Z_i(s) K_b(x - X_i(s)) ds},$$

across a grid of possible time values t , where X is the marker, Z is the exposure and $\alpha(z)$ is the marker-only hazard, see [get_alpha](#) for more details.

Value

A vector of $\hat{h}_x(t)$ for a grid of possible time values t .

See Also

[get_alpha](#), [h_xt](#)

Examples

```
library(survival)
b = 10
x = 3
Landmark <- 2
pbcT1 <- pbc2[which(pbc2$year< Landmark & pbc2$years> Landmark),]
b=0.9

arg1l1<-get_h_x1l(pbcT1,'albumin',event_time_name='years',
                  time_name='year',event_name='status2',2,0.9)
arg1lc<-get_h_x(pbcT1,'albumin',event_time_name='years',
                time_name='year',event_name='status2',2,0.9)

#Caclulate the local contant and local linear survival functions
br_s = seq(Landmark, 14, length=99)
sfalb2l1<- make_sf( (br_s[2]-br_s[1])/4 , arg1l1)
sfalb2lc<- make_sf( (br_s[2]-br_s[1])/4 , arg1lc)

#For comparison, also calculate the Kaplan-Meier
kma2<- survfit(Surv(years , status2) ~ 1, data = pbcT1)

#Plot the survival functions:
plot(br_s, sfalb2l1, type="l", col=1, lwd=2, ylab="Survival probability", xlab="Marker level")
lines(br_s, sfalb2lc, lty=2, lwd=2, col=2)
lines(kma2$time, kma2$surv, type="s", lty=2, lwd=2, col=3)

legend("topright", c( "Local linear HQM", "Local constant HQM",
                     "Kaplan-Meier"), lty=c(1, 2, 2), col=1:3, lwd=2, cex=1.7)
```

get_h_xll

*Local linear future conditional hazard rate estimator***Description**

Calculates the local linear future hazard rate function, conditional on a marker value x , across a set of time values t .

Usage

```
get_h_xll(data, marker_name, event_time_name, time_name, event_name, x, b)
```

Arguments

data	A data frame of time dependent data points. Missing values are allowed.
marker_name	The column name of the marker values in the data frame data .
event_time_name	The column name of the event times in the data frame data .
time_name	The column name of the times the marker values were observed in the data frame data .
event_name	The column name of the events in the data frame data .
x	Numeric value of the last observed marker value.
b	Bandwidth parameter.

Details

The function [get_h_xll](#) implements the local linear future conditional hazard estimator

$$\hat{h}_x(t) = \frac{\sum_{i=1}^n \int_0^T \hat{\alpha}_i(X_i(t+s)) Z_i(t+s) Z_i(s) K_b(x - X_i(s)) ds}{\sum_{i=1}^n \int_0^T Z_i(t+s) Z_i(s) K_b(x - X_i(s)) ds},$$

across a grid of possible time values t , where X is the marker, Z is the exposure and $\alpha(z)$ is the marker-only hazard, see [get_alpha](#) for more details.

Value

A vector of $\hat{h}_x(t)$ for a grid of possible time values t .

See Also

[get_alpha](#), [h_xt](#)

Examples

```

library(survival)
library(JM)

# Compare Local constant and local linear estimator, use KM for reference
# Albumin marker, use landmarking
Landmark <- 2
pbcT1 <- pbc2[which(pbc2$year< Landmark & pbc2$years> Landmark),]
b=0.9

arg1ll<-get_h_xll(pbcT1, 'albumin', event_time_name = 'years', time_name = 'year',
                 event_name = 'status2', 2, 0.9)
arg1lc<-get_h_x(pbcT1, 'albumin', event_time_name = 'years', time_name = 'year',
                 event_name = 'status2', 2, 0.9)

#Calculate the local constant and local linear survival functions
br_s = seq(Landmark, 14, length=99)
sfalb2ll<- make_sf( (br_s[2]-br_s[1])/4 , arg1ll)
sfalb2lc<- make_sf( (br_s[2]-br_s[1])/4 , arg1lc)

#For comparison, also calculate the Kaplan-Meier
kma2<- survfit(Surv(years , status2) ~ 1, data = pbcT1)

#Plot the survival functions:
plot(br_s, sfalb2ll, type="l", col=1, lwd=2, ylab="Survival probability", xlab="Marker level")
lines(br_s, sfalb2lc, lty=2, lwd=2, col=2)
lines(kma2$time, kma2$surv, type="s", lty=2, lwd=2, col=3)

legend("topright", c( "Local linear HQM", "Local constant HQM", "Kaplan-Meier"),
      lty=c(1, 2, 2), col=1:3, lwd=2, cex=1.7)

## Not run:
#Compare JM, HQM and KM for Bilirubin
b = 10
Landmark <- 1
lmeFit <- lme(serBilir ~ year, random = ~ year | id, data = pbc2)
coxFit <- coxph(Surv(years, status2) ~ serBilir, data = pbc2.id, x = TRUE)

jointFit0 <- jointModel(lmeFit, coxFit, timeVar = "year",
                      method = "piecewise-PH-aGH")
pbcT1 <- pbc2[which(pbc2$year< Landmark & pbc2$years> Landmark),]

timesS1 <- seq(1,14,by=0.5)
predT1 <- survfitJM(jointFit0, newdata = pbcT1,survTimes = timesS1)
nm<-length(predT1$summaries)

mat.out1<-matrix(nrow=length(timesS1), ncol=nm)
for(r in 1:nm)
{

```

```

SurvLand <- predT1$summaries[[r]][,"Mean"][1]
mat.out1[,r] <- predT1$summaries[[r]][,"Mean"]/SurvLand
}
sfit1y<-rowMeans(mat.out1, na.rm=TRUE)

arg1<- get_h_x(pbcT1, 'serBilir', event_time_name = 'years',
               time_name = 'year', event_name = 'status2', 1, 10)
br_s1 = seq(Landmark, 14, length=99)
sfbil1<- make_sf( (br_s1[2]-br_s1[1])/5.4 , arg1)
kma1<- survfit(Surv(years , status2) ~ 1, data = pbcT1)

plot(br_s1, sfbil1, type="l", ylim=c(0,1), xlim=c(Landmark,14),
      ylab="Survival probability", xlab="years",lwd=2)
lines(timesS1, sfit1y, col=2, lwd=2, lty=2)
lines(kma1$time, kma1$surv, type="s", lty=2, lwd=2, col=3 )
legend("bottomleft", c("HQM est.", "Joint Model est.", "Kaplan-Meier"),
      lty=c(1,2,2), col=1:3, lwd=2, cex=1.7)

## End(Not run)

```

g_xt

*Computation of a key component for wild bootstrap***Description**

Implements a key part for the wild bootstrap of the hqm estimator.

Usage

```
g_xt(br_X, br_s, size_s_grid, int_X, x, t, b, Yi, Y, n)
```

Arguments

br_X	Marker value grid points that will be used in the evaluation.
br_s	Time value grid points that will be used in the evaluation.
size_s_grid	Size of the time grid.
int_X	Position of the linear interpolated marker values on the marker grid.
x	Numeric value of the last observed marker value.
t	Numeric value of the time the function should be evaluated.
b	Bandwidth.
Yi	A matrix made by <code>make_Yi</code> indicating the exposure.
Y	A matrix made by <code>make_Y</code> indicating the exposure.
n	Number of individuals.

Details

The function implements

$$\hat{g}_{t,x}(z) = \frac{1}{n} \sum_{j=1}^n \int_0^{T-t} \hat{E}(X_j(t+s))^{-1} K_b(z, X_j(t+s)) Z_j(t+s) Z_j(s) K_b(x, X_j(s)) ds,$$

for every value z on the marker grid, where $\hat{E}(x) = \frac{1}{n} \sum_{j=1}^n \int_0^T K_b(x, X_j(s)) Z_j(s) ds$, Z the exposure and X the marker.

Value

A vector of $\hat{g}_{t,x}(z)$ for all values z on the marker grid.

Examples

```

pbc2_id = to_id(pbc2)
size_s_grid <- size_X_grid <- 100
n = max(as.numeric(pbc2$id))
X = pbc2$serBilir
s = pbc2$year
br_s = seq(0, max(s), max(s)/(size_s_grid-1))
br_X = seq(min(X), max(X), (max(X)-min(X))/(size_X_grid-1))

X_lin = lin_interpolate(br_s, pbc2_id$id, pbc2$id, X, s)

int_X <- findInterval(X_lin, br_X)
int_s = rep(1:length(br_s), n)

Yi<-make_Yi(pbc2, pbc2_id, X_lin, br_X, br_s, size_s_grid, size_X_grid, int_s, int_X, 'years', n)
Y<-make_Y(pbc2, pbc2_id, X_lin, br_X, br_s, size_s_grid, size_X_grid, int_s, int_X, 'years', n)

t = 2
x = 2
b = 10

g_xt(br_X, br_s, size_s_grid, int_X, x, t, b, Yi, Y, n)

```

h_xt

Local constant future conditional hazard rate estimation at a single time point

Description

Calculates the future conditional hazard rate for a marker value x and a time value t .

Usage

```
h_xt(br_X, br_s, int_X, size_s_grid, alpha, x,t, b, Yi,n)
```

Arguments

<code>br_X</code>	Vector of grid points for the marker values X .
<code>br_s</code>	Vector of grid points for the time values s .
<code>int_X</code>	Position of the linear interpolated marker values on the marker grid.
<code>size_s_grid</code>	Size of the time grid.
<code>alpha</code>	Marker-hazard obtained from get_alpha .
<code>x</code>	Numeric value of the last observed marker value.
<code>t</code>	Numeric time value.
<code>b</code>	Bandwidth.
<code>Yi</code>	A matrix made by make_Yi indicating the exposure.
<code>n</code>	Number of individuals.

Details

Function [h_xt](#) implements the future conditional hazard estimator

$$\hat{h}_x(t) = \frac{\sum_{i=1}^n \int_0^T \hat{\alpha}_i(X_i(t+s)) Z_i(t+s) Z_i(s) K_b(x - X_i(s)) ds}{\sum_{i=1}^n \int_0^T Z_i(t+s) Z_i(s) K_b(x - X_i(s)) ds},$$

where X is the marker, Z is the exposure and $\alpha(z)$ is the marker-only hazard, see [get_alpha](#) for more details. The future conditional hazard is defined as

$$h_{x,T}(t) = P(T_i \in (t+T, t+T+dt) | X_i(T) = x, T_i > t+T),$$

where T_i is the survival time and X_i the marker of individual i observed in the time frame $[0, T]$. Function [h_xt](#) uses an classic (unmodified) kernel function $K_b()$, e.g. the Epanechnikov kernel.

Value

A single numeric value of $\hat{h}_x(t)$.

References

[doi:10.1080/03461238.1998.10413997](https://doi.org/10.1080/03461238.1998.10413997)

See Also

[get_alpha](#)

Examples

```

pbc2_id = to_id(pbc2)
size_s_grid <- size_X_grid <- 100
n = max(as.numeric(pbc2$id))
s = pbc2$year
X = pbc2$serBilir
XX = pbc2_id$serBilir

```

```

ss <- pbc2_id$years
delta <- pbc2_id$status2
br_s = seq(0, max(s), max(s)/( size_s_grid-1))
br_X = seq(min(X), max(X), (max(X)-min(X))/( size_X_grid-1))

X_lin = lin_interpolate(br_s, pbc2_id$id, pbc2$id, X, s)

int_X <- findInterval(X_lin, br_X)
int_s = rep(1:length(br_s), n)

N <- make_N(pbc2, pbc2_id, breaks_X=br_X, breaks_s=br_s, ss, XX, delta)
Y <- make_Y(pbc2, pbc2_id, X_lin, br_X, br_s, size_s_grid, size_X_grid, int_s, int_X, 'years', n)

b = 1.7
alpha<-get_alpha(N, Y, b, br_X, K=Epan )

Yi <- make_Yi(pbc2, pbc2_id, X_lin, br_X, br_s, size_s_grid, size_X_grid, int_s, int_X, 'years', n)

x = 2
t = 2
h_hat = h_xt(br_X, br_s, int_X, size_s_grid, alpha, x, t, b, Yi, n)

```

h_xtll	<i>Local linear future conditional hazard rate estimation at a single time point</i>
--------	--

Description

Calculates the local linear future conditional hazard rate for a marker value x and a time value t .

Usage

```
h_xtll(br_X, br_s, int_X, size_s_grid, alpha, x,t, b, Yi,n, Y)
```

Arguments

br_X	Vector of grid points for the marker values X .
br_s	Vector of grid points for the time values s .
int_X	Position of the linear interpolated marker values on the marker grid.
size_s_grid	Size of the time grid.
alpha	Marker-hazard obtained from get_alpha .
x	Numeric value of the last observed marker value.
t	Numeric time value.
b	Bandwidth.
Yi	A matrix made by make_Yi indicating the exposure.
n	Number of individuals.
Y	A matrix made by make_Y indicating the exposure.

Details

Function `h_xtll` implements the future conditional hazard estimator

$$\hat{h}_x(t) = \frac{\sum_{i=1}^n \int_0^T \hat{\alpha}_i(X_i(t+s)) Z_i(t+s) Z_i(s) K_b(x - X_i(s)) ds}{\sum_{i=1}^n \int_0^T Z_i(t+s) Z_i(s) K_b(x - X_i(s)) ds},$$

where X is the marker, Z is the exposure and $\alpha(z)$ is the marker-only hazard, see `get_alpha` for more details. The future conditional hazard is defined as

$$h_{x,T}(t) = P(T_i \in (t+T, t+T+dt) | X_i(T) = x, T_i > t+T),$$

where T_i is the survival time and X_i the marker of individual i observed in the time frame $[0, T]$.

The function `h_xtll`, in the place of $K_b()$ uses the kernel

$$K_{x,b}(u) = \frac{K_b(u) - K_b(u)u^T D^{-1}c_1}{c_0 - c_1^T D^{-1}c_1},$$

where $c_1 = (c_{11}, \dots, c_{1d})^T$, $D = (d_{ij})_{(d+1) \times (d+1)}$ with

$$c_0 = \sum_{i=1}^n \int_0^T K_b(x - X_i(s)) Z_i(s) ds, c_{ij} = \sum_{i=1}^n \int_0^T K_b(x - X_i(s)) \{x - X_{ij}(s)\} Z_i(s) ds, d_{jk} = \sum_{i=1}^n \int_0^T K_b(x - X_i(s)) \{x - X_{ij}(s)\} \{x - X_{ik}(s)\} Z_i(s) ds,$$

see also Nielsen (1998).

Value

A single numeric value of $\hat{h}_x(t)$.

References

[doi:10.1080/03461238.1998.10413997](https://doi.org/10.1080/03461238.1998.10413997)

See Also

`get_alpha`, `dij`

Examples

```

pbc2_id = to_id(pbc2)
size_s_grid <- size_X_grid <- 100
n = max(as.numeric(pbc2$id))
s = pbc2$year
X = pbc2$serBilir
XX = pbc2_id$serBilir
ss <- pbc2_id$years
delta <- pbc2_id$status2
br_s = seq(0, max(s), max(s)/(size_s_grid-1))
br_X = seq(min(X), max(X), (max(X)-min(X))/(size_X_grid-1))

X_lin = lin_interpolate(br_s, pbc2_id$id, pbc2$id, X, s)

```

```

int_X <- findInterval(X_lin, br_X)
int_s = rep(1:length(br_s), n)

N <- make_N(pbc2, pbc2_id, breaks_X=br_X, breaks_s=br_s, ss, XX, delta)
Y <- make_Y(pbc2, pbc2_id, X_lin, br_X, br_s, size_s_grid, size_X_grid, int_s, int_X, 'years', n)

b = 1.7
alpha<-get_alpha(N, Y, b, br_X, K=Epan )

Yi <- make_Yi(pbc2, pbc2_id, X_lin, br_X, br_s, size_s_grid, size_X_grid, int_s, int_X, 'years', n)

x = 2
t = 2
h_hat = h_xtll(br_X, br_s, int_X, size_s_grid, alpha, x, t, b, Yi, n, Y)

```

h_xt_vec

Hqm estimator on the marker grid

Description

Computes the hqm estimator on the marker grid.

Usage

```
h_xt_vec(br_X, br_s, size_s_grid, alpha, t, b, Yi, int_X, n)
```

Arguments

br_X	Marker value grid points that will be used in the evaluation.
br_s	Time value grid points that will be used in the evaluation.
size_s_grid	Size of the time grid.
alpha	Marker-hazard obtained from get_alpha .
t	Numeric value of the time the function should be evaluated.
b	Bandwidth.
Yi	A matrix made by make_Yi indicating the exposure.
int_X	Position of the linear interpolated marker values on the marker grid.
n	Number of individuals.

Details

The function implements the future conditional hazard estimator

$$\hat{h}_x(t) = \frac{\sum_{i=1}^n \int_0^T \hat{\alpha}_i(X_i(t+s)) Z_i(t+s) Z_i(s) K_b(x - X_i(s)) ds}{\sum_{i=1}^n \int_0^T Z_i(t+s) Z_i(s) K_b(x - X_i(s)) ds},$$

for every x on the marker grid where X is the marker, Z is the exposure and $\alpha(z)$ is the marker-only hazard, see [get_alpha](#) for more details.

Value

A vector of $\hat{h}_x(t)$ for all values x on the marker grid.

Examples

```
# Longitudinal data example

pbc2_id = to_id(pbc2)
size_s_grid <- size_X_grid <- 100
n = max(as.numeric(pbc2$id))
s = pbc2$year
X = pbc2$serBilir
XX = pbc2_id$serBilir
ss <- pbc2_id$years
delta <- pbc2_id$status2
br_s = seq(0, max(s), max(s)/( size_s_grid-1))
br_X = seq(min(X), max(X), (max(X)-min(X))/( size_X_grid-1))

X_lin = lin_interpolate(br_s, pbc2_id$id, pbc2$id, X, s)

int_X <- findInterval(X_lin, br_X)
int_s = rep(1:length(br_s), n)

N <- make_N(pbc2, pbc2_id, br_X, br_s, ss, XX, delta)
Y <- make_Y(pbc2, pbc2_id, X_lin, br_X, br_s,
            size_s_grid, size_X_grid, int_s, int_X, event_time = 'years', n)

b = 1.7
alpha<-get_alpha(N, Y, b, br_X, K=Epan )

Yi <- make_Yi(pbc2, pbc2_id, X_lin, br_X, br_s,
              size_s_grid, size_X_grid, int_s, int_X, event_time = 'years', n)

t = 2

h_xt_vec(br_X, br_s, size_s_grid, alpha, t, b, Yi, int_X, n)

# Time-invariant data example:
# pbc2 dataset, single event per individual version:
# 312 observations, most recent event per individual.
# Use landmarking to produce comparable curve with KM.
library(survival)
Landmark <- 3 #set the landmark to 3 years
pbc2.use<- to_id(pbc2) # keep only the most recent row per patient
pbcT1 <- pbc2.use[which(pbc2.use$year< Landmark & pbc2.use$years> Landmark),]

timesS2 <- seq(Landmark,14,by=0.5)
b=0.9
arg1<- get_h_x(pbcT1, 'albumin', event_time_name = 'years', time_name = 'year',
               event_name = 'status2', 2, b)

br_s2 = seq(Landmark, 14, length=99)
sfalb2<- make_sf( (br_s2[2]-br_s2[1])/1.35 , arg1)
```

```

kma2<- survfit(Surv(years , status2) ~ 1, data = pbcT1)

#Plot the survival functions:
plot(br_s2, sfalb2, type="l", ylim=c(0,1), xlim=c(Landmark,14), ylab="Survival probability",
      xlab="years",lwd=2, main="HQM and KM survival functions, conditional on albumin=2,
      for the time-invariant pbc dataset")
lines(kma2$time, kma2$surv, type="s",lty=2, lwd=2, col=2)
legend("bottomleft", c("HQM est.", "Kaplan-Meier"), lty=c(1,2), col=1:2, lwd=2, cex=1.7)

```

Kernels

Classical (unmodified) kernel and related functionals

Description

Implements the classical kernel function and related functionals

Usage

```

K_b(b,x,y, K)
xK_b(b,x,y, K)
K_b_mat(b,x,y, K)

```

Arguments

x	A vector of design points where the kernel will be evaluated.
y	A vector of sample data points.
b	The bandwidth to use (a scalar).
K	The kernel function to use.

Details

The function `K_b` implements the classical kernel function calculation

$$h^{-1}K\left(\frac{x-y}{h}\right)$$

for scalars x and y while `xK_b` implements the functional

$$h^{-1}K\left(\frac{x-y}{h}\right)(x-y)$$

again for for scalars x and y . The function `K_b_mat` is the vectorized version of `K_b`. It uses as inputs the vectors $(X_1, \dots, X_n)$ and $(Y_1, \dots, Y_n)$ and returns a $n \times n$ matrix with entries

$$h^{-1}K\left(\frac{X_i - Y_j}{h}\right)$$

Value

Scalar values for K_b and xK_b and matrix outputs for K_b_mat.

lin_interpolate	<i>Linear interpolation</i>
-----------------	-----------------------------

Description

Implements a linear interpolation between observed marker values.

Usage

```
lin_interpolate(t, i, data_id, data_marker, data_time)
```

Arguments

t	A vector of time values where the function should be evaluated.
i	A vector of ids of individuals for whom the marker values should be interpolated.
data_id	The vector of ids from a data frame of time dependent variables.
data_marker	The vector of marker values from a data frame of time dependent variables.
data_time	The vector of time values from a data frame of time dependent variables.

Details

Given time points $t_1, \dots, t_K$ and marker values $m_1, \dots, m_J$ at different time points $t_1^m, \dots, t_J^m$, the function calculates a linear interpolation f with $f(t_i^m) = m_i$ at the time points $t_1, \dots, t_K$ for all indicated individuals. Returned are then $(f(t_1), \dots, f(t_K))$. Note that the first value is always observed at time point 0 and the function f is extrapolated constantly after the last observed marker value.

Value

A matrix with columns $(f(t_1), \dots, f(t_K))$ as described above for every individual in the vector i.

Examples

```
size_s_grid <- 100
X = pbc2$serBilir
s = pbc2$year
br_s = seq(0, max(s), max(s)/(size_s_grid-1))
pbc2_id = to_id(pbc2)

X_lin = lin_interpolate(br_s, pbc2_id$id, pbc2$id, X, s)
```

11K_b

Local linear kernel

Description

Implements the local linear kernel function.

Usage

```
11K_b(b,x,y, K)
```

Arguments

x	A vector of design points where the kernel will be evaluated.
y	A vector of sample data points.
b	The bandwidth to use (a scalar).
K	The kernel function to use.

Details

Implements the local linear kernel

$$K_{x,b}(u) = \frac{K_b(u) - K_b(u)u^T D^{-1}c_1}{c_0 - c_1^T D^{-1}c_1},$$

where $c_1 = (c_{11}, \dots, c_{1d})^T$, $D = (d_{ij})_{(d+1) \times (d+1)}$ with

$$c_0 = \sum_{i=1}^n \int_0^T K_b(x - X_i(s)) Z_i(s) ds, c_{ij} = \sum_{i=1}^n \int_0^T K_b(x - X_i(s)) \{x - X_{ij}(s)\} Z_i(s) ds, d_{jk} = \sum_{i=1}^n \int_0^T K_b(x - X_i(s)) \{x - X_{ij}(s)\} \{x - X_{ik}(s)\} Z_i(s) ds,$$

see also Nielsen (1998).

Value

Matrix output with entries the values of the kernel function at each point.

References

[doi:10.1080/03461238.1998.10413997](https://doi.org/10.1080/03461238.1998.10413997)

llweights

Local linear weight functions

Description

Implements the weights to be used in the local linear HQM estimator.

Usage

```
sn.0(xin, xout, h, kfun)
sn.1(xin, xout, h, kfun)
sn.2(xin, xout, h, kfun)
```

Arguments

xin	Sample values.
xout	Grid points where the estimator will be evaluated.
h	Bandwidth parameter.
kfun	Kernel function.

Details

The function implements the local linear weights in the definition of the estimator $\hat{h}_x(t)$, see also [h_xt](#)

Value

A vector of $s_n(x)$ for all values x on the marker grid.

make_N, make_Ni, make_Y, make_Yi

Occurance and Exposure on grids

Description

Auxiliary functions that help automate the process of calculating integrals with occurances or exposure processes.

Usage

```
make_N(data, data.id, breaks_X, breaks_s, ss, XX, delta)
make_Ni(breaks_s, size_s_grid, ss, delta, n)
make_Y(data, data.id, X_lin, breaks_X, breaks_s,
        size_s_grid, size_X_grid, int_s, int_X, event_time = 'years', n)
make_Yi(data, data.id, X_lin, breaks_X, breaks_s,
        size_s_grid, size_X_grid, int_s, int_X, event_time = 'years', n)
```

Arguments

data	A data frame of time dependent data points. Missing values are allowed.
data.id	An id data frame obtained from to_id .
breaks_X	Marker value grid points where the function will be evaluated.
breaks_s	Time value grid points where the function will be evaluated.
ss	Vector with event times.
XX	Vector of last observed marker values.
delta	0-1 vector of whether events happened.
size_s_grid	Size of the time grid.
size_X_grid	Size of the marker grid.
n	Number of individuals.
X_lin	Linear interpolation of observed marker values evaluated on the marker grid.
int_s	Position of the observed time values on the time grid.
int_X	Position of the linear interpolated marker values on the marker grid.
event_time	String of the column name with the event times.

Details

Implements matrices for the computation of integrals with occurrences and exposures of the form

$$\int f(s)Z(s)Z(s+t)ds, \int f(s)Z(s)ds, \int f(s)dN(s).$$

where N is a 0-1 counting process, Z the exposure and f an arbitrary function.

Value

The functions [make_N](#) and [make_Y](#) return a matrix on the time grid and marker grid for occurrence and exposure, respectively, while [make_Ni](#) and [make_Yi](#) return a matrix on the time grid for every individual again for occurrence and exposure, respectively.

See Also

[h_xt](#), [g_xt](#), [get_alpha](#)

Examples

```

pbc2_id = to_id(pbc2)
size_s_grid <- size_X_grid <- 100
n = max(as.numeric(pbc2$id))
s = pbc2$year
X = pbc2$serBilir
XX = pbc2_id$serBilir
ss <- pbc2_id$years
delta <- pbc2_id$status2
br_s = seq(0, max(s), max(s)/(size_s_grid-1))

```

```

br_X = seq(min(X), max(X), (max(X)-min(X))/( size_X_grid-1))

X_lin = lin_interpolate(br_s, pbc2_id$id, pbc2$id, X, s)

int_X <- findInterval(X_lin, br_X)
int_s = rep(1:length(br_s), n)

N <- make_N(pbc2, pbc2_id, br_X, br_s, ss, XX, delta)
Y <- make_Y(pbc2, pbc2_id, X_lin, br_X, br_s,
            size_s_grid, size_X_grid, int_s, int_X, event_time = 'years', n)
Yi <- make_Yi(pbc2, pbc2_id, X_lin, br_X, br_s,
              size_s_grid, size_X_grid, int_s, int_X, event_time = 'years', n)
Ni <- make_Ni(br_s, size_s_grid, ss, delta, n)

```

make_sf

Survival function from a hazard

Description

Creates a survival function from a hazard rate which was calculated on a grid.

Usage

```
make_sf(step_size_s_grid, haz)
```

Arguments

`step_size_s_grid` Numeric value indicating the distance between two grid continuous grid points.

`haz` Vector of hazard values. Hazard rate must have been calculated on a time grid.

Details

The function `make_sf` calculates the survival function

$$S(t) = \exp\left(-\int_0^t h(t)dt\right),$$

where h is the hazard rate. Here, a discretisation via an equidistant grid $\{t_i\}$ on $[0, t]$ is used to calculate the integral and it is assumed that h has been calculated for exactly these time points t_i .

Value

A vector of values $S(t_i)$.

Examples

```
make_sf(0.1, rep(0.1,10))
```

pbc2

*Mayo Clinic Primary Biliary Cirrhosis Data***Description**

Followup of 312 randomised patients with primary biliary cirrhosis, a rare autoimmune liver disease, at Mayo Clinic.

Usage

pbc2

Format

A data frame with 1945 observations on the following 20 variables.

`id` patients identifier; in total there are 312 patients.

`years` number of years between registration and the earlier of death, transplantation, or study analysis time.

`status` a factor with levels alive, transplanted and dead.

`drug` a factor with levels placebo and D-penicil.

`age` at registration in years.

`sex` a factor with levels male and female.

`year` number of years between enrollment and this visit date, remaining values on the line of data refer to this visit.

`ascites` a factor with levels No and Yes.

`hepatomegaly` a factor with levels No and Yes.

`spiders` a factor with levels No and Yes.

`edema` a factor with levels No edema (i.e., no edema and no diuretic therapy for edema), edema no diuretics (i.e., edema present without diuretics, or edema resolved by diuretics), and edema despite diuretics (i.e., edema despite diuretic therapy).

`serBilir` serum bilirubin in mg/dl.

`serChol` serum cholesterol in mg/dl.

`albumin` albumin in gm/dl.

`alkaline` alkaline phosphatase in U/liter.

`SGOT` SGOT in U/ml.

`platelets` platelets per cubic ml / 1000.

`prothrombin` prothrombin time in seconds.

`histologic` histologic stage of disease.

`status2` a numeric vector with the value 1 denoting if the patient was dead, and 0 if the patient was alive or transplanted.

References

- Fleming, T. and Harrington, D. (1991) *Counting Processes and Survival Analysis*. Wiley, New York.
- Therneau, T. and Grambsch, P. (2000) *Modeling Survival Data: Extending the Cox Model*. Springer-Verlag, New York.

Examples

```
summary(pbc2)
```

```
prep_boot
```

```
Precomputation for wild bootstrap
```

Description

Implements key components for the wild bootstrap of the hqm estimator in preparation for obtaining confidence bands.

Usage

```
prep_boot(g_xt, alpha, Ni, Yi, size_s_grid, br_X, br_s, t, b, int_X, x, n)
```

Arguments

<code>g_xt</code>	A vector obtained by <code>g_xt</code> .
<code>alpha</code>	A vector of the marker only hazard on the marker grid obtained by <code>get_alpha</code> .
<code>Ni</code>	A matrix made by <code>make_Ni</code> indicating the occurrence.
<code>Yi</code>	A matrix made by <code>make_Yi</code> indicating the exposure.
<code>size_s_grid</code>	Size of the time grid.
<code>br_X</code>	Vector of grid points for the marker values.
<code>br_s</code>	Time value grid points that will be used in the evaluation.
<code>t</code>	Numeric value of the time the function should be evaluated.
<code>b</code>	Bandwidth.
<code>int_X</code>	Position of the linear interpolated marker values on the marker grid.
<code>x</code>	Numeric value of the last observed marker value.
<code>n</code>	Number of individuals.

Details

The function implements

$$A_B(t) = \frac{1}{\sqrt{n}} \sum_{i=1}^n \int_0^T \hat{g}_{i,t,x_*}(X_i(s)) V_i \{dN_i(s) - \hat{\alpha}_i(X_i(s)) Z_i(s) ds\},$$

and

$$B_B(t) = \frac{1}{\sqrt{n}} \sum_{i=1}^n V_i \{ \hat{\Gamma}(t, x_*)^{-1} W_i(t, x_*) - \hat{h}_{x_*}(t) \},$$

where $V \sim N(0, 1)$,

$$W_i(t) = \int_0^T \hat{\alpha}_i(X_i(t+s)) Z_i(t+s) Z_i(s) K_b(x_*, X_i(s)) ds,$$

and

$$\hat{\Gamma}(t, x) = \frac{1}{n} \sum_{i=1}^n \int_0^{T-t} Z_i(t+s) Z_i(s) K_b(x, X_i(s)) ds,$$

with Z being the exposure and X the marker.

Value

A list of 5 items. The first two are vectors for calculating A_B and the third one a vector for B_B . The 4th one is the value of the hqm estimator that can also be obtained by [h_xt](#) and the last one is the value of Γ .

See Also

[Conf_bands](#)

Examples

```

pbc2_id = to_id(pbc2)
size_s_grid <- size_X_grid <- 100
n = max(as.numeric(pbc2$id))
s = pbc2$year
X = pbc2$serBilir
XX = pbc2_id$serBilir
ss <- pbc2_id$years
delta <- pbc2_id$status2
br_s = seq(0, max(s), max(s)/(size_s_grid-1))
br_X = seq(min(X), max(X), (max(X)-min(X))/(size_X_grid-1))

X_lin = lin_interpolate(br_s, pbc2_id$id, pbc2$id, X, s)

int_X <- findInterval(X_lin, br_X)
int_s = rep(1:length(br_s), n)

N <- make_N(pbc2, pbc2_id, br_X, br_s, ss, XX, delta)
Y <- make_Y(pbc2, pbc2_id, X_lin, br_X, br_s,
```

```

size_s_grid, size_X_grid, int_s, int_X, event_time = 'years', n)

b = 1.7
alpha<-get_alpha(N, Y, b, br_X, K=Epan )

Yi <- make_Yi(pbc2, pbc2_id, X_lin, br_X, br_s,
              size_s_grid, size_X_grid, int_s, int_X, event_time = 'years', n)
Ni  <- make_Ni(br_s, size_s_grid, ss, delta, n)

t = 2
x = 2

g = g_xt(br_X, br_s, size_s_grid, int_X, x, t, b, Yi, Y, n)

Boot_all = prep_boot(g, alpha, Ni, Yi, size_s_grid, br_X, br_s, t, b, int_X, x, n)
Boot_all

```

prep_cv

Prepare for Cross validation bandwidth selection

Description

Implements the calculation of the hqm estimator on cross validation data sets. This is a preparation for the cross validation bandwidth selection technique for future conditional hazard rate estimation based on marker information data.

Usage

```

prep_cv(data, data.id, marker_name, event_time_name = 'years',
        time_name = 'year', event_name = 'status2', n, I, b)

```

Arguments

data	A data frame of time dependent data points. Missing values are allowed.
data.id	An id data frame obtained from to_id .
marker_name	The column name of the marker values in the data frame data .
event_time_name	The column name of the event times in the data frame data .
time_name	The column name of the times the marker values were observed in the data frame data .
event_name	The column name of the events in the data frame data .
n	Number of individuals.
I	Number of observations leave out for a K cross validation.
b	Bandwidth.

Details

The function splits the data set via [dataset_split](#) and calculates for every splitted data set the hqm estimator

$$\hat{h}_x(t) = \frac{\sum_{i=1}^n \int_0^T \hat{\alpha}_i(X_i(t+s)) Z_i(t+s) Z_i(s) K_b(x - X_i(s)) ds}{\sum_{i=1}^n \int_0^T Z_i(t+s) Z_i(s) K_b(x - X_i(s)) ds},$$

for all x on the marker grid and t on the time grid, where X is the marker, Z is the exposure and $\alpha(z)$ is the marker-only hazard, see [get_alpha](#) for more details.

Value

A list of matrices for every cross validation data set with $\hat{h}_x(t)$ for all x on the marker grid and t on the time grid.

See Also

[b_selection](#)

Examples

```

pbc2_id = to_id(pbc2)
n = max(as.numeric(pbc2$id))
b = 1.5
I = 26
h_xt_mat_list = prep_cv(pbc2, pbc2_id, 'serBilir', 'years', 'year', 'status2', n, I, b)

```

Q1

*Bandwidth selection score Q1***Description**

Calculates a part for the K-fold cross validation score.

Usage

```
Q1(h_xt_mat, int_X, size_X_grid, n, Yi)
```

Arguments

<code>h_xt_mat</code>	A matrix of the estimator for the future conditional hazard rate for all values x and t .
<code>int_X</code>	Vector of the position of the observed marker values in the grid for marker values.
<code>size_X_grid</code>	Numeric value indicating the number of grid points for marker values.
<code>n</code>	Number of individuals.
<code>Yi</code>	A matrix made by make_Yi indicating the exposure.

Details

The function implements

$$Q_1 = \sum_{i=1}^N \int_0^T \int_s^T Z_i(t) Z_i(s) \hat{h}_{X_i(s)}^2(t-s) dt ds,$$

where $\hat{h}$ is the hqm estimator, Z the exposure and X the marker.

Value

A value of the score Q1.

See Also

[b_selection](#)

Examples

```

pbc2_id = to_id(pbc2)
size_s_grid <- size_X_grid <- 100
n = max(as.numeric(pbc2$id))
s = pbc2$year
X = pbc2$serBilir
XX = pbc2_id$serBilir
ss <- pbc2_id$years
delta <- pbc2_id$status2
br_s = seq(0, max(s), max(s)/(size_s_grid-1))
br_X = seq(min(X), max(X), (max(X)-min(X))/(size_X_grid-1))

X_lin = lin_interpolate(br_s, pbc2_id$id, pbc2$id, X, s)

int_X <- findInterval(X_lin, br_X)
int_s = rep(1:length(br_s), n)

N <- make_N(pbc2, pbc2_id, br_X, br_s, ss, XX, delta)
Y <- make_Y(pbc2, pbc2_id, X_lin, br_X, br_s,
            size_s_grid, size_X_grid, int_s, int_X, event_time = 'years', n)

b = 1.7
alpha <- get_alpha(N, Y, b, br_X, K=Epan)

Yi <- make_Yi(pbc2, pbc2_id, X_lin, br_X, br_s,
              size_s_grid, size_X_grid, int_s, int_X, event_time = 'years', n)
Ni <- make_Ni(br_s, size_s_grid, ss, delta, n)

t = 2

h_xt_mat = t(sapply(br_s[1:99],
                    function(si){h_xt_vec(br_X, br_s, size_s_grid, alpha, t, b, Yi, int_X, n)}))

Q = Q1(h_xt_mat, int_X, size_X_grid, n, Yi)

```

R_K

*Bandwidth selection score R***Description**

Calculates a part for the K-fold cross validation score.

Usage

```
R_K(h_xt_mat_list, int_X, size_X_grid, Yi, Ni, n)
```

Arguments

h_xt_mat_list	A list of matrices for all cross validation data sets. Each matrix contains the estimator with the future conditional hazard rate for all values x and t and the respected data set.
int_X	Vector of the position of the observed marker values in the grid for marker values.
size_X_grid	Numeric value indicating the number of grid points for marker values.
Yi	A matrix made by make_Yi indicating the exposure.
Ni	A matrix made by make_Ni indicating the occurrence.
n	Number of individuals.

Details

The function implements the estimator

$$\hat{R}_K = \sum_{j=1}^K \sum_{i \in I_j} \int_0^T g_i^{-I_j}(t) dN_i(t),$$

where $\hat{g}_i^{-I_j}(t) = \int_0^t Z_i(s) \hat{h}_{X_i(s)}^{-I_j}(t-s) ds$, and $\hat{h}^{-I_j}$ is estimated without information from all counting processes i with $i \in I_j$. This function estimates

$$R = \sum_{i=1}^N \int_0^T \int_s^T Z_i(t) Z_i(s) \hat{h}_{X_i(s)}(t-s) h_{X_i(s)}(t-s) dt ds.$$

where $\hat{h}$ is the hqm estimator, Z the exposure and X the marker.

Value

A matrix with $\hat{g}_i^{-I_j}(t)$ for all individuals i and time grid points t .

See Also

[b_selection](#)

Examples

```

pbc2_id = to_id(pbc2)
n = max(as.numeric(pbc2$id))
b = 1.5
I = 104
h_xt_mat_list = prep_cv(pbc2, pbc2_id, 'serBilir', 'years', 'year', 'status2', n, I, b)

size_s_grid <- size_X_grid <- 100
s = pbc2$year
X = pbc2$serBilir
br_s = seq(0, max(s), max(s)/( size_s_grid-1))
br_X = seq(min(X), max(X), (max(X)-min(X))/( size_X_grid-1))

ss <- pbc2_id$years
delta <- pbc2_id$status2

X_lin = lin_interpolate(br_s, pbc2_id$id, pbc2$id, X, s)
int_X <- findInterval(X_lin, br_X)
int_s = rep(1:length(br_s), n)

Yi <- make_Yi(pbc2, pbc2_id, X_lin, br_X, br_s,
              size_s_grid, size_X_grid, int_s, int_X, 'years', n)
Ni <- make_Ni(br_s, size_s_grid, ss, delta, n)

R = R_K(h_xt_mat_list, int_X, size_X_grid, Yi, Ni, n)
R

```

to_id

Event data frame

Description

Creates a data frame with only one entry per individual from a data frame with time dependent data. The resulting data frame focusses on the event time and the last observed marker value.

Usage

```
to_id(data_set)
```

Arguments

data_set A data frame of time dependent data points. Missing values are allowed.

Details

The function `to_id` uses a data frame of time dependent marker data to create a smaller data frame with only one entry per individual, the last observed marker value and the event time. Note that the column indicating the individuals must have the name `id`. Note also that this data frame is similar

to `pbcb2.id` from the JM package with the difference that the last observed marker value instead of the first one is captured.

Value

A data frame with only one entry per individual.

Examples

```
data_set.id = to_id(pbc2)
```

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