Package 'IrregLong'

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Title Analysis of Longitudinal Data with Irregular Observation Times

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Description Analysis of longitudinal data for which the times of observation are random variables that are potentially associated with the outcome process. The package includes inverseintensity weighting methods (Lin H, Scharfstein DO, Rosenheck RA (2004) <doi:10.1111/j.1467-9868.2004.b5543.x>) and multiple outputation (Pullenayegum EM (2016) <doi:10.1002/sim.6829>).

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Description

Create an abacus plot Creates an abacus plot, depicting visits per subject over time

Usage

```
abacus.plot(
    n,
    time,
    id,
    data,
    tmin,
    tmax,
    xlab.abacus = "Time",
    ylab.abacus = "Subject",
    pch.abacus = 16,
    col.abacus = 1
```

n	the number of subjects to randomly sample. Subjects are sampled without re- placement and therefore n must be smaller than the total number of subjects in the dataset
time	character string indicating which column of the data contains the time at which the visit occurred
id	character string indicating which column of the data identifies subjects
data	data frame containing the variables in the model
tmin	the smallest time to include on the x-axis
tmax	the largest time to include on the x-axis
xlab.abacus	the label for the x-axis
ylab.abacus	the label for the y-axis
pch.abacus	the plotting character for the points on the abacus plot
col.abacus	the colour of the rails on the abacus plot

addcensoredrows

Details

This function creates a plot for n randomly sampled individuals from the supplied dataset, with one row per subject and one point per visit. This can be useful for visualising the extent of irregularity in the visit process. For example, with perfect repeated measures data (i.e., no irregularity), the points will line up vertically. With greater irregularity, the points will be randomly scattered over time.

Value

produces a plot depicting observation times for each subject. No values are returned

Examples

```
library(MEMSS)
data(Phenobarb)
Phenobarb$event <- 1-as.numeric(is.na(Phenobarb$conc))
data <- Phenobarb[Phenobarb$event==1,]
abacus.plot(n=20,time="time",id="Subject",data=data,tmin=0,tmax=16*24,
xlab.abacus="Time in hours",pch=16,col.abacus=gray(0.8))
```

addcensoredrows Add rows corresponding to censoring times to a longitudinal dataset

Description

Add rows corresponding to censoring times to a longitudinal dataset

Usage

```
addcensoredrows(data, maxfu, tinvarcols, id, time, event)
```

data	The dataset to which rows are to be added. The data should have one row per observation
maxfu	The maximum follow-up time per subject. If all subjects have the same follow- up time, this can be supplied as a single number. Otherwise, maxfu should be a dataframe with the first column specifying subject identifiers and the second giving the follow-up time for each subject.
tinvarcols	A vector of column numbers corresponding to variables in data that are time-invariant.
id	character string indicating which column of the data identifies subjects
time	character string indicating which column of the data contains the time at which the visit occurred
event	character string indicating which column of the data indicates whether or not a visit occurred. If every row corresponds to a visit, then this column will consist entirely of ones

Value

The original dataset with extra rows corresponding to censoring times

Examples

```
x <- c(1:3,1:2,1:5)
x0 <- c(rep(2,3),rep(0,2),rep(1,5))
id <- c(rep(1,3),rep(2,2),rep(3,5))
time <- c(0,4,6,2,3,1,3,5,6,7)
event <- c(1,1,1,0,1,0,1,1,1,1)
data <- as.data.frame(cbind(x,id,time,event,x0))
addcensoredrows(data,maxfu=8,id="id",time="time",tinvarcols=5,event="event")
```

```
x <- c(1:3,1:2,1:5)
x0 <- c(rep(2,3),rep(0,2),rep(1,5))
id <- c(rep(1,3),rep(2,2),rep(3,5))
time <- c(0,4,6,2,3,1,3,5,6,7)
event <- c(1,1,1,0,1,0,1,1,1,1)
data <- as.data.frame(cbind(x,id,time,event,x0))
maxfu.id <- 1:3
maxfu.time <- c(6,5,8)
maxfu <- cbind(maxfu.id,maxfu.time)
maxfu <- as.data.frame(maxfu)
addcensoredrows(data,maxfu=maxfu,id="id",time="time",tinvarcols=5,event="event")
```

```
extent.of.irregularity
```

Measures of extent of visit irregularity Provides visual and numeric measures of the extent of irregularity in observation times in a longitudinal dataset

Description

Measures of extent of visit irregularity Provides visual and numeric measures of the extent of irregularity in observation times in a longitudinal dataset

Usage

```
extent.of.irregularity(
   data,
   time = "time",
   id = "id",
   scheduledtimes = NULL,
   cutpoints = NULL,
   ncutpts = NULL,
   maxfu = NULL,
   plot = FALSE,
```

extent.of.irregularity

```
legendx = NULL,
legendy = NULL,
formula = NULL,
tau = NULL
```

Arguments

)

The data containing information on subject identifiers and visit times
A character indicating which column of the data contains the times at which each of the observations in data was made
A character indicating which column of the data contains subject identifiers. ids are assumed to be consecutive integers, with the first subject having id 1
For studies with protocol-specified visit times, a vector of these times. Defaults to NULL, in which case it is assumed that there are no protocolized visit times
For studies with scheduled visit times, an array of dimension ncutpts by length(scheduledtimes) by 2 giving, for ncutpts sets of left and right cutpoints for each protocolized scheduled visit times. The left-hand cutpoints correspond to cutpoints[,,1] and the right-hand cutpoints to cutpoints[,,2]. Defaults to NULL, in which case cutpoints are computed as described below.
The number of sets of cutpoints to consider
The maximum follow-up time per subject. If all subjects have the same follow- up time, this can be supplied as a single number. Otherwise, maxfu should be a dataframe with the first column specifying subject identifiers and the second giving the follow-up time for each subject.
logical parameter indicating whether plots should be produced.
The x-coordinate for the position of the legend in the plot of mean proportion of individuals with 0, 1 and \$>\$ 1 visit per bin.
The y-coordinate for the position of the legend in the plot of mean proportion of individuals with 0, 1 and \$>\$ 1 visit per bin.
For studies without protocolized visit times, the formula for the null counting process model for the visit times
The maximum time of interest

Details

This function provides plots and a numerical summary of the extent of irregularity in visit times. For any given set of cutpoints, it computes the proportion of individuals with 0, 1 and >1 observation(s) in each bin, then takes the mean over bins. The sizes of the bins are varied and these proportions are plotted against bin size. In addition, then mean proportion of individuals with >1 visit per bin is plotted vs. the mean proportion of individuals with 0 visits per bin, and the area under the curve is calculated (AUC). An AUC of 0 represents perfect repeated measures while a Poisson Process has an AUC of 0. If cutpoints are not supplied, they are computed as follows: (a) for studies with protocolized visit times, the left- and right-hand cutpoints are positioned at the protocolized time minus (or plus, for right-hand cutpoints) (1,...,ncutpts)/ncutpts times the gap to the previous (or next, respectively) protocolized visit time; (b) for studies with no protocolized visit times, cutpoints

are calculated by finding, for each j in 1,...,ncutpts the largest times for which the cumulative hazard is less than j divided by the cumulative hazard evaluated at the maximum time of interest. This corresponds to choosing cutpoints such that the expected number of visits per bin is roughly equal within each set.

Value

a list with counts equal to a 3-dimensional by neutpts matrix giving, for each set of cutpoints, the mean proportion of individuals with zero, 1 and >1 visits per bin, and AUC, the area under the curve of the plot of the proportion of individuals with >1 visit per bin vs. the proportion of individuals with 0 visits per bin.

Examples

```
library(nlme)
library(survival)
data(Phenobarb)
Phenobarb$event <- 1-as.numeric(is.na(Phenobarb$conc))
data <- Phenobarb
data <- data[data$event==1,]
data$id <- as.numeric(data$Subject)
counts <- extent.of.irregularity(data,time="time",id="id",scheduledtimes=NULL,
cutpoints=NULL,ncutpts=10,
maxfu=16*24,plot=TRUE,legendx=NULL,legendy=NULL,formula=Surv(time.lag,time,event)~1,tau=16*24)
counts$counts
counts$auc</pre>
```

Given a	proportional	hazards	model fo	r visit	intensities,	compute
inverse-in	ntensity weigh	its.				

Description

iiw

For a longitudinal dataset subject to irregular observation, use a Cox proportional hazards model for visit intensities to compute inverse intensity weights

Usage

iiw(phfit, data, id, time, first)

Arguments

phfit	coxph object for the visit process
data	The dataset featuring longitudinal data subject to irregular observation for which inverse-intensity weights are desired
id	character string indicating which column of the data identifies subjects
time	character string indicating which column of the data contains the time at which the visit occurred

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iiw.weights

first logical variable. If TRUE, the first observation for each individual is assigned an intensity of 1. This is appropriate if the first visit is a baseline visit at which recruitment to the study occurred; in this case the baseline visit is observed with probability 1.

Value

A vector of inverse-intensity weights for each row of the dataset. The first observation for each subject is assumed to have an intensity of 1.

See Also

```
Other iiw: iiw.weights(), iiwgee()
```

Examples

```
library(nlme)
data(Phenobarb)
library(survival)
library(geepack)
data(Phenobarb)
Phenobarb$event <- 1-as.numeric(is.na(Phenobarb$conc))</pre>
data <- Phenobarb
data <- data[data$event==1,]</pre>
data$id <- as.numeric(data$Subject)</pre>
data <- data[data$time<16*24,]</pre>
data <- lagfn(data, lagvars=c("time","conc"), id="Subject", time="time", lagfirst = NA)</pre>
head(data)
mph <- coxph(Surv(time.lag,time,event)~I(conc.lag>0 & conc.lag<=20) + I(conc.lag>20 & conc.lag<=30)</pre>
 + I(conc.lag>30)+ cluster(id),,data=data)
summary(mph)
data$weight <- iiw(mph,data,"id","time",TRUE)</pre>
head(data)
```

iiw.weights Compute inverse-intensity weights.

Description

Since the vector of weights is ordered on id and time, if you intend to merge these weights onto your original dataset it is highly recommended that you sort the data before running iiw.weights

Usage

```
iiw.weights(
  formulaph,
  formulanull = NULL,
  data,
```

```
id,
time,
event,
lagvars,
invariant = NULL,
maxfu,
lagfirst = lagfirst,
first,
frailty = FALSE
)
```

Arguments

formulaph	the formula for the proportional hazards model for the visit intensity that will be used to derive inverse-intensity weights. The formula should usually use the counting process format (i.e. Surv(start,stop,event)). If a frailty model is used, the cluster(id) term should appear before other covariates
formulanull	if stabilised weights are to be used, the formula for the null model used to sta- bilise the weights
data	data frame containing the variables in the model
id	character string indicating which column of the data identifies subjects
time	character string indicating which column of the data contains the time at which the visit occurred
event	character string indicating which column of the data indicates whether or not a visit occurred. If every row corresponds to a visit, then this column will consist entirely of ones
lagvars	a vector of variable names corresponding to variables which need to be lagged by one visit to fit the visit intensity model. Typically time will be one of these variables. The function will internally add columns to the data containing the values of the lagged variables from the previous visit. Values of lagged variables for a subject's first visit will be set to NA. To access these variables in specifying the proportional hazards formulae, add ".lag" to the variable you wish to lag. For example, if time is the variable for time, time.lag is the time of the previous visit
invariant	a vector of variable names corresponding to variables in data that are time- invariant. It is not necessary to list every such variable, just those that are invari- ant and also included in the proportional hazards model
maxfu	the maximum follow-up time(s). If everyone is followed for the same length of time, this can be given as a single value. If individuals have different follow-up times, maxfu should have the same number of elements as there are rows of data
lagfirst	A vector giving the value of each lagged variable for the first time within each subject. This is helpful if, for example, time is the variable to be lagged and you know that all subjects entered the study at time zero
first	logical variable. If TRUE, the first observation for each individual is assigned an intensity of 1. This is appropriate if the first visit is a baseline visit at which recruitment to the study occurred; in this case the baseline visit is observed with probability 1.

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iiw.weights

frailty	logical variable. If TRUE, a frailty model is fit to calculate the inverse intensity
	weights. If FALSE, a marginal semi-parametric model is fit. Frailty models are
	helpful when fitting semi-parametric joint models.

Details

Given longitudinal data with irregular visit times, fit a Cox proportional hazards model for the visit intensity, then use it to compute inverse-intensity weights

Value

a vector of inverse-intensity weights, ordered on id then time

References

- Lin H, Scharfstein DO, Rosenheck RA. Analysis of Longitudinal data with Irregular, Informative Follow-up. Journal of the Royal Statistical Society, Series B (2004), 66:791-813
- Buzkova P, Lumley T. Longitudinal data analysis for generalized linear models with follow-up dependent on outcome-related variables. The Canadian Journal of Statistics 2007; 35:485-500.

See Also

Other iiw: iiwgee(), iiw() Other iiw: iiwgee(), iiw()

```
library(nlme)
data(Phenobarb)
library(survival)
library(geepack)
Phenobarb$event <- 1-as.numeric(is.na(Phenobarb$conc))</pre>
data <- Phenobarb
data <- data[data$event==1,]</pre>
data$id <- as.numeric(data$Subject)</pre>
data <- data[data$time<16*24,]</pre>
i <- iiw.weights(Surv(time.lag,time,event)~I(conc.lag>0 & conc.lag<=20) +
I(conc.lag>20 & conc.lag<=30) + I(conc.lag>30)+ cluster(id),
id="id",time="time",event="event",data=data,
invariant="id",lagvars=c("time","conc"),maxfu=16*24,lagfirst=0,first=TRUE)
data$weight <- i$iiw.weight</pre>
summary(i$m)
# can use to fit a weighted GEE
mw <- geeglm(conc ~ I(time^3) + log(time) , id=Subject, data=data, weights=weight)</pre>
summary(mw)
# agrees with results through the single command iiwgee
miiwgee <- iiwgee(conc ~ I(time^3) + log(time),</pre>
Surv(time.lag,time,event)~I(conc.lag>0 & conc.lag<=20) +</pre>
I(conc.lag>20 & conc.lag<=30) + I(conc.lag>30)+ cluster(id),
id="id",time="time",event="event",data=data,
```

iiwgee

```
invariant="id",lagvars=c("time","conc"),maxfu=16*24,lagfirst=0,first=TRUE)
summary(miiwgee$geefit)
```

iiwgee

Fit an inverse-intensity weighted GEE.

Description

Implements inverse-intensity weighted GEEs as first described by Lin, Scharfstein and Rosenheck (2004). A Cox proportional hazards model is applied to the visit intensities, and the hazard multipliers are used to compute inverse-intensity weights. Using the approach described by Buzkova and Lumley (2007) avoids the need to compute the baseline hazard.

Usage

```
iiwgee(
  formulagee,
  formulaph,
  formulanull = NULL,
  data,
  id,
  time,
 event,
  family = gaussian,
 lagvars,
  invariant = NULL,
 maxfu,
 lagfirst,
  first
```

Arguments

)

formulagee	the formula for the GEE model to be fit. The syntax used is the same as in geeglm
formulaph	the formula for the proportional hazards model for the visit intensity that will be used to derive inverse-intensity weights. The formula should usually use the counting process format (i.e. Surv(start,stop,event))
formulanull	if stabilised weights are to be used, the formula for the null model used to stabilise the weights
data	data frame containing the variables in the model
id	character string indicating which column of the data identifies subjects
time	character string indicating which column of the data contains the time at which the visit occurred

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iiwgee

event	character string indicating which column of the data indicates whether or not a visit occurred. If every row corresponds to a visit, then this column will consist entirely of ones
family	family to be used in the GEE fit. See geeglm for documentation
lagvars	a vector of variable names corresponding to variables which need to be lagged by one visit to fit the visit intensity model. Typically time will be one of these variables. The function will internally add columns to the data containing the values of the lagged variables from the previous visit. Values of lagged variables for a subject's first visit will be set to NA. To access these variables in specifying the proportional hazards formulae, add ".lag" to the variable you wish to lag. For example, if time is the variable for time, time.lag is the time of the previous visit
invariant	a vector of variable names corresponding to variables in data that are time- invariant. It is not necessary to list every such variable, just those that are invari- ant and also included in the proportional hazards model
maxfu	the maximum follow-up time(s). If everyone is followed for the same length of time, this can be given as a single value. If individuals have different follow-up times, maxfu should have the same number of elements as there are rows of data
lagfirst	A vector giving the value of each lagged variable for the first time within each subject. This is helpful if, for example, time is the variable to be lagged and you know that all subjects entered the study at time zero
first	logical variable. If TRUE, the first observation for each individual is assigned an intensity of 1. This is appropriate if the first visit is a baseline visit at which recruitment to the study occurred; in this case the baseline visit is observed with probability 1.

Details

Let the outcome of interest be Y and suppose that subject i has j^{th} observation at T_{ij} . Let $N_i(t)$ be a counting process for the number of observations for subject i up to and including time t. Suppose that N_i has intensity λ given by

$$\lambda_i(t) = \lambda 0(t) \exp(Z_i(t)\gamma).$$

Then the inverse-intensity weights are

$$exp(-Z_i(t)\gamma).$$

If Y_i is the vector of observations for subject *i*, to be regressed onto X_i (i.e. $E(Y_i|X_i) = \mu(X_i;\beta)$ with $g(\mu(X_i; beta) = X_i\beta$, then the inverse-intensity weighted GEE equations are

$$\sum_{i} \frac{\partial \mu_i}{\partial \beta} V_i^{-1} \Delta_i (Y_i X_i \beta) = 0$$

, where Δ_i is a diagonal matrix with j^{th} entry equal to $\exp(-Z_i(T_{ij})\gamma)$ and V_i is the working variance matrix. Warning: Due to the way some gee functions incorporate weights, if using inverse-intensity weighting you should use working independence.

Value

a list, with the following elements:

geefit	the fitted GEE, see documentation for geeglm for details
phfit	the fitted proportional hazards model, see documentation for coxph for details

References

- Lin H, Scharfstein DO, Rosenheck RA. Analysis of Longitudinal data with Irregular, Informative Follow-up. Journal of the Royal Statistical Society, Series B (2004), 66:791-813
- Buzkova P, Lumley T. Longitudinal data analysis for generalized linear models with follow-up dependent on outcome-related variables. The Canadian Journal of Statistics 2007; 35:485-500.

See Also

Other iiw: iiw.weights(), iiw()

```
library(nlme)
data(Phenobarb)
library(survival)
library(geepack)
Phenobarb$event <- 1-as.numeric(is.na(Phenobarb$conc))</pre>
data <- Phenobarb
data <- data[data$event==1,]</pre>
data$id <- as.numeric(data$Subject)</pre>
data <- data[data$time<16*24,]</pre>
miiwgee <- iiwgee(conc ~ I(time^3) + log(time),</pre>
Surv(time.lag,time,event)~I(conc.lag>0 & conc.lag<=20) +</pre>
I(conc.lag>20 & conc.lag<=30) + I(conc.lag>30)+ cluster(id),
id="id",time="time",event="event",data=data,
invariant="id",lagvars=c("time","conc"),maxfu=16*24,lagfirst=0,first=TRUE)
summary(miiwgee$geefit)
summary(miiwgee$phfit)
# compare to results without weighting
m <- geeglm(conc ~ I(time^3) + log(time) , id=Subject, data=data); print(summary(m))</pre>
time <- (1:200)
unweighted <- cbind(rep(1,200),time^3,log(time))%*%m$coefficients</pre>
weighted <- cbind(rep(1,200),time^3,log(time))%*%miiwgee$geefit$coefficients</pre>
plot(data$time,data$conc,xlim=c(0,200),pch=16)
lines(time,unweighted,type="1")
lines(time,weighted,col=2)
legend (0,60,legend=c("Unweighted","Inverse-intensity weighted"),col=1:2,bty="n",lty=1)
```

lagfn

Description

Create lagged versions the variables in data

Usage

```
lagfn(data, lagvars, id, time, lagfirst = NA)
```

Arguments

data	The data to be lagged
lagvars	The names of the columns in the data to be lagged
id	A character indicating which column of the data contains subject identifiers. ids are assumed to be consecutive integers, with the first subject having id 1
time	A character indicating which column of the data contains the times at which each of the observations in data was made
lagfirst	A vector giving the value of each lagged variable for the first time within each subject. This is helpful if, for example, time is the variable to be lagged and you know that all subjects entered the study at time zero

Value

The original data frame with lagged variables added on as columns. For example, if the data frame contains a variable named x giving the value of x for each subject i at each visit j, the returned data frame will contain a column named x.lag containing the value of x for subject i at visit j-1. If j is the first visit for subject i, the lagged value is set to NA

```
library(nlme)
data(Phenobarb)
head(Phenobarb)
data <- lagfn(Phenobarb, "time", "Subject", "time")
head(data)</pre>
```

Liang

Description

Fits a semi-parametric joint model as described by Liang et al. (2009).

Usage

```
Liang(
  data,
  Yname,
  Xnames,
  Wnames,
  Znames = NULL,
  formulaobs = NULL,
  id,
  time,
  invariant = NULL,
  lagvars = NULL,
  lagfirst = NULL,
  maxfu,
  baseline,
  n.knots = NULL,
  kappa = NULL,
  Xfn = NULL,
  Wfn = NULL,
  . . .
)
```

data	data frame containing the variables in the model
Yname	character string indicating the column containing the outcome variable
Xnames	vector of character strings indicating the names of the columns of the fixed effects in the outcome regression model
Wnames	vector of character strings indicating the names of the columns of the random effects in the outcome regression model
Znames	vector of character strings indicating the names of the columns of the covariates in the visit intensity model
formulaobs	formula for the observation intensity model
id	character string indicating which column of the data identifies subjects
time	character string indicating which column of the data contains the time at which the visit occurred

Liang

invariant	a vector of variable names corresponding to variables in data that are time- invariant. It is not necessary to list every such variable, just those that are invari- ant and also included in the visit intensity model
lagvars	a vector of variable names corresponding to variables which need to be lagged by one visit to fit the visit intensity model. Typically time will be one of these variables. The function will internally add columns to the data containing the values of the lagged variables from the previous visit. Values of lagged variables for a subject's first visit will be set to NA. To access these variables in specifying the proportional hazards formulae, add ".lag" to the variable you wish to lag. For example, if time is the variable for time, time.lag is the time of the previous visit
lagfirst	A vector giving the value of each lagged variable for the first time within each subject. This is helpful if, for example, time is the variable to be lagged and you know that all subjects entered the study at time zero
maxfu	The maximum follow-up time per subject. If all subjects have the same follow- up time, this can be supplied as a single number. Otherwise, maxfu should be a dataframe with the first column specifying subject identifiers and the second giving the follow-up time for each subject.
baseline	An indicator for whether baseline (time=0) measurements are included by design. Equal to 1 if yes, 0 if no.
n.knots	integer giving the number of knots to use in fitting the frailty model. See documentation for frailtyPenal for more details
kappa	positive smoothing parameter in the penalized likelihood estimation. See documentation for frailtyPenal for more details
Xfn	A function that takes as its first argument the subject identifier and has time as its second argument, and returns the value of X for the specified subject at the specified time.
Wfn	A function that takes as its first argument the subject identifier and has time as its second argument, and returns the value of W for the specified subject at the specified time
	other arguments to Xfn and Yfn

Details

This function fits a semi-parametric joint model as described in Liang (2009), using a frailty model to estimate the parameters in the visit intensity model

The Liang method requires a value of X and W for every time over the observation period. If Xfn is left as NULL, then the Liang function will use, for each subject and for each time t, the values of X and W at the observation time closest to t.

Value

the regression coefficients corresponding to the fixed effects in the outcome regression model. Closed form expressions for standard errors of the regression coefficients are not available, and Liang et al (2009) recommend obtaining these through bootstrapping.

References

Liang Y, Lu W, Ying Z. Joint modelling and analysis of longitudinal data with informative observation times. Biometrics 2009; 65:377-384.

```
# replicate simulation in Liang et al.
## Not run:
library(data.table)
library(survival)
datasimi <- function(id){</pre>
X1 <- runif(1,0,1)
X2 <- rbinom(1,1,0.5)
Z <- rgamma(1,1,1)
Z1 <- rnorm(1, Z-1, 1)
gamma < - c(0.5, -0.5)
beta <- c(1,-1)
hazard <- Z*exp(X1/2 - X2/2)
C <- runif(1,0,5.8)
t <- 0
tlast <- t
y <- t + X1-X2 + Z1*X2 + rnorm(1,0,1)</pre>
wait <- rexp(1,hazard)</pre>
while(tlast+wait<C){</pre>
  tnew <- tlast+wait</pre>
    y <- c(y,tnew + X1-X2 + Z1*X2 + rnorm(1,0,1))</pre>
    t <- c(t,tnew)</pre>
    tlast <- tnew</pre>
    wait <- rexp(1,hazard)</pre>
 }
 datai <- list(id=rep(id,length(t)),t=t,y=y,</pre>
      X1=rep(X1,length(t)),X2=rep(X2,length(t)),C=rep(C,length(t)))
 return(datai)
 }
 sim1 <- function(it,nsubj){</pre>
 data <- lapply(1:nsubj,datasimi)</pre>
 data <- as.data.frame(rbindlist(data))</pre>
 data$event <- 1
 C <- tapply(data$C,data$id,mean)</pre>
 tapply(data$C,data$id,sd)
 maxfu <- cbind(1:nsubj,C)</pre>
 maxfu <- as.data.frame(maxfu)</pre>
 res <- Liang(data=data, id="id",time="t",Yname="y",</pre>
             Xnames=c("X1","X2"),
             Wnames=c("X2"),Znames=c("X1","X2"), formulaobs=Surv(t.lag,t,event)~X1
             + X2+ cluster(id), invariant=c
             ("id", "X1", "X2"), lagvars="t", lagfirst=NA, maxfu=maxfu,
             baseline=1,n.knots=6, kappa=10000)
 return(res)
 }
 # change n to 500 to replicate results of Liang et al.
 n <- 10
```

то

```
s <- lapply(1:n,sim1,nsubj=200)
smat <- matrix(unlist(s),byrow=TRUE,ncol=2)
apply(smat,2,mean)</pre>
```

End(Not run)

mo

Multiple outputation for longitudinal data subject to irregular observation.

Description

Multiple outputation is a procedure whereby excess observations are repeatedly randomly sampled and discarded. The method was originally developed to handle clustered data where cluster size is informative, for example when studying pups in a litter. In this case, analysis that ignores cluster size results in larger litters being over-represented in a marginal analysis. Multiple outputation circumvents this problem by randomly selecting one observation per cluster. Multiple outputation has been further adapted to handle longitudinal data subject to irregular observation; here the probability of being retained on any given outputation is inversely proportional to the visit intensity. This function creates multiply outputted datasets, analyses each separately, and combines the results to produce a single estimate.

Usage

```
mo(
    noutput,
    fn,
    data,
    weights,
    singleobs,
    id,
    time,
    keep.first,
    var = TRUE,
    ...
)
```

noutput	the number of outputations to be used
fn	the function to be applied to the outputted datasets. fn should return a vector or scalar; if var=TRUE the second column of fn should be an estimate of standard error.
data	the original dataset on which multiple outputation is to be performed
weights	the weights to be used in the outputation, i.e. the inverse of the probability that a given observation will be selected in creating an outputted dataset. Ignored if singleobs=TRUE

singleobs

logical variable indicating whether a single observation should be retained for
each subject
character string indicating which column of the data identifies subjects

id	character string indicating which column of the data identifies subjects
time	character string indicating which column of the data contains the time at which the visit occurred
keep.first	logical variable indicating whether the first observation should be retained with probability 1. This is useful if the data consists of an observation at baseline followed by follow-up at stochastic time points.
var	logical variable indicating whether fn returns variances in addition to point esti- mates
	other arguments to fn.

Value

18

a list containing the multiple outputation estimate of the function fn applied to the data, its standard error, and the relative efficiency of using noutput outputations as opposed to an infinite number

References

- Hoffman E, Sen P, Weinberg C. Within-cluster resampling. Biometrika 2001; 88:1121-1134
- Follmann D, Proschan M, Leifer E. Multiple outputation: inference for complex clustered data by averaging analyses from independent data. Biometrics 2003; 59:420-429
- Pullenayegum EM. Multiple outputation for the analysis of longitudinal data subject to irregular observation. Statistics in Medicine (in press)

See Also

.

Other mo: outputation()

```
library(nlme)
data(Phenobarb)
library(survival)
library(geepack)
Phenobarb$event <- 1-as.numeric(is.na(Phenobarb$conc))</pre>
data <- Phenobarb
data <- data[data$event==1,]</pre>
data$id <- as.numeric(data$Subject)</pre>
data <- data[data$time<16*24,]</pre>
i <- iiw.weights(Surv(time.lag,time,event)~I(conc.lag>0 & conc.lag<=20) +
I(conc.lag>20 & conc.lag<=30) + I(conc.lag>30)+ cluster(id),
id="id",time="time",event="event",data=data, invariant="id",lagvars=c("time","conc"),maxfu=16*24,
         lagfirst=c(0,0),first=TRUE)
wt <- i$iiw.weight
wt[wt>quantile(i$iiw.weight,0.95)] <- quantile(i$iiw.weight,0.95)</pre>
```

outputation

```
data$wt <- wt
reg <- function(data){
est <- summary(geeglm(conc~I(time^3) + log(time), id=id,data=data))$coefficients[,1:2]
est <- data.matrix(est)
return(est)
}
mo(20,reg,data,wt,singleobs=FALSE,id="id",time="time",keep.first=FALSE)
# On outputation, the dataset contains small numbers of observations per subject
# and hence the GEE sandwich variance estimate underestimates variance; this is why
# the outputation-based variance estimate fails. This can be remedied by using a
# sandwich variance error correction (e.g. Fay-Graubard, Mancl-DeRouen).</pre>
```

outputation

Create an outputted dataset for use with multiple outputation.

Description

Multiple outputation is a procedure whereby excess observations are repeatedly randomly sampled and discarded. The method was originally developed to handle clustered data where cluster size is informative, for example when studying pups in a litter. In this case, analysis that ignores cluster size results in larger litters being over-represented in a marginal analysis. Multiple outputation circumvents this problem by randomly selecting one observation per cluster. Multiple outputation has been further adapted to handle longitudinal data subject to irregular observation; here the probability of being retained on any given outputation is inversely proportional to the visit intensity. This function creates a single outputted dataset.

Usage

```
outputation(data, weights, singleobs, id, time, keep.first)
```

data	the original dataset on which multiple outputation is to be performed
weights	the weights to be used in the outputation, i.e. the inverse of the probability that a given observation will be selected in creating an outputted dataset. Ignored if singleobs=TRUE
singleobs	logical variable indicating whether a single observation should be retained for each subject
id	character string indicating which column of the data identifies subjects
time	character string indicating which column of the data contains the time at which the visit occurred
keep.first	logical variable indicating whether the first observation should be retained with probability 1. This is useful if the data consists of an observation at baseline followed by follow-up at stochastic time points.

Value

the outputted dataset.

References

- Hoffman E, Sen P, Weinberg C. Within-cluster resampling. Biometrika 2001; 88:1121-1134
- Follmann D, Proschan M, Leifer E. Multiple outputation: inference for complex clustered data by averaging analyses from independent data. Biometrics 2003; 59:420-429
- Pullenayegum EM. Multiple outputation for the analysis of longitudinal data subject to irregular observation. Statistics in Medicine (in press).

See Also

Other mo: mo()

```
library(nlme)
data(Phenobarb)
library(survival)
library(geepack)
Phenobarb$event <- 1-as.numeric(is.na(Phenobarb$conc))</pre>
data <- Phenobarb
data <- data[data$event==1,]</pre>
data$id <- as.numeric(data$Subject)</pre>
data <- data[data$time<16*24,]</pre>
i <- iiw.weights(Surv(time.lag,time,event)~I(conc.lag>0 & conc.lag<=20) +</pre>
I(conc.lag>20 & conc.lag<=30) + I(conc.lag>30)+ cluster(id),
id="id",time="time",event="event",data=data,
invariant="id",lagvars=c("time","conc"),maxfu=16*24,lagfirst=0,first=TRUE)
data$weight <- i$iiw.weight</pre>
head(data)
data.output1 <- outputation(data,data$weight,singleobs=FALSE,</pre>
id="id",time="time",keep.first=FALSE)
head(data.output1)
data.output2 <- outputation(data,data$weight,singleobs=FALSE,</pre>
id="id",time="time",keep.first=FALSE)
head(data.output2)
data.output3 <- outputation(data,data$weight,singleobs=FALSE,</pre>
id="id",time="time",keep.first=FALSE)
head(data.output3)
# Note that the outputted dataset varies with each command run; outputation is done at random
```

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