Package 'multipleNCC'

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Type Package

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Description Fit Cox proportional hazard models with a weighted partial likelihood. It handles one or multiple endpoints, additional matching and makes it possible to reuse controls for other endpoints.
Depends survival, mgcv
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R topics documented:
multipleNCC-package
Chenprob
CVD_Accidents
GAMprob
GLMprob
KMprob
ModelbasedVar
multipleNCC-internal
pChen
nGAM

 pKM
 11

 PoststratVar
 12

 print.wpl
 12

 summary.wpl
 13

 wpl
 13

Index 17

multipleNCC-package Weighted partial likelihood for nested case-control data

Description

Fits Cox proportional hazards models with a weighted partial likelihood. It handles competing risks (with one endpoint being a special situation). It uses cases and controls from other endpoints as additional controls for each endpoint. See wpl for help.

Four weight estimators are implemented; Kaplan-Meier type KMprob, GAM (GAMprob), GLM (GLMprob) and local averaging (Chenprob)

Details

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Author(s)

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References

Samuelsen, SO. (1997) A pseudolikelihood approach to analysis of nested case-control studies. *Biometrika* **84(2)**, 379-394

Samuelsen, SO., et al. (2007) Stratified case-cohort analysis of general cohort sampling designs. *Scand J Stat* **34(1)**, 103-119

Chen, KN. (2001) Generalized case-cohort sampling. *J Roy Stat Soc Ser B* **63(4)**, 791 - 809 Stoer NC and Samuelsen SO (2012): Comparison of estimators in nested case-control studies with multiple outcomes. Lifetime Data Analysis, 18(3), 261-283.

See Also

wpl, coxph, Chenprob, GLMprob, GAMprob, KMprob

Chenprob 3

Description

Estimates sampling probabilities with local averaging (Chen, 2001). The weights included in the Cox-regressions (wpl) and which could be used for other procedures are inverse sampling probabilities i.e. the inverse of these probabilities. The probabilities are estimated for all subjects in the cohort.

Usage

```
Chenprob(survtime, samplestat, no.intervals = 10, left.time = 0, no.intervals.left = c(3,4))
```

Arguments

survtime Follow-up time for all cohort subjects

samplestat A vector containing sampling and status information: 0 represents non-sampled

subjects in the cohort, 1: sampled controls, 2,3,... indicate different events.

Cohort dimension.

soring

left.time Entry time if the survival times are left-truncated. Cohort dimension.

no.intervals.left

Number of intervals for Chen-weights with left-truncation. A vector on the form [number of intervals for left truncated time, number of intervals for survival

time].

Value

A vector of cohort dimension of sampling probabilities.

Author(s)

Nathalie C. Stoer

References

Chen KN (2001) Generalized case-cohort sampling. J Roy Stat Soc Ser B 63(4):791-809 Stoer NC and Samuelsen SO (2012): Comparison of estimators in nested case-control studies with multiple outcomes. Lifetime Data Analysis, 18(3), 261-283.

See Also

```
wpl, coxph, GAMprob, GLMprob, KMprob
```

4 CVD_Accidents

Examples

CVD_Accidents

Causes of death in three counties in Norway in 1974-2000

Description

Causes of death from 1974-2000 for all men and women participating in a cardiovascular health screening in 1974-1978 in three counties in Norway. All variables are know for all cohort members and it is thus a synthetic nested case-control stidy. One control per case is sampled for cardiovascular disease cases and subjects who died from alcohol abuse, liver disease, and accidents and violence. The controls are matched sex and BMI plus/minus 2 in addition to being alive at the time the case died.

Usage

```
data(CVD_Accidents)
```

Format

A data frame with 3933 observations on the following 23 variables.

```
agestart Age at health survey, inclusion time
agestop Age at censoring
dead Indicator for death from any cause (0=censored, 1=dead)
```

dead1 Indicator for cancer death (0=censored or dead from other cause than cancer, 1=dead from cancer)

dead2 Indicator for death from cardiovascular disease, including sudden death (0=censored or dead from other causes than cardiovascular diseas, 1=dead from cardiovascular diseas)

GAMprob 5

dead3 Indicator for death from other medical causes (0=censored or dead from cancer, cardiovascular diseas, alcohol abuse, liver disease, violence or accidents, 1=dead from other medical causes)

dead4 Indicator for death from alcohol abuse, liver disease, violence and accidents (0=censored or death from other medical causes than alcohol abuse, liver disase, violence or accidents, 1=death from alcohol abuse, liver disease, violence and accidents)

sex sex (1=male, 2=female)

county county in Norway (5=Oppland, 14=Sogn og Fjordane, 20=Finnmark)

sbp Systolic blood pressure at health screening

bmi Body mass index at helth screening

smkstart Age started smoking

smkgr Smoking group (1=never smoked, 2=former smoker, 3=1-9 cigaretts per day, 4=10-19 cigaretts per day, 5=20+ cigaretts per day, 6=pipe or cigar)

smoking3gr Smoking 3 groups (1=never smoked, 2=former smoker, 3=smoker)

samplestat Indicator for sampling and events (0=non-sampled subjects in the cohort, 1=sampled controls, 2=dead from cardiovascular disease, 3=dead from alcohol abuse, liver disease, violence or accidents

dead24 Indicator for death from either cardiovascular disease or alcohol abuse, liver disease, violence or accidents (0=censored or dead from other causes than cardiovascular disease, alcohol abuse, liver disease, violence or accidents, 1=death from cardiovascular disease, alcohol abuse, liver disease, violence or accidents)

Source

http://folk.uio.no/borgan/abg-2008/data/data.html

GAMprob

Sampling probabilities estimated with generalized additive models.

Description

Estimates sampling probabilities with generalized additive models. The weights included in the Cox-regressions (wpl) and which could be used for other procedures are inverse sampling probabilities i.e. the inverse of these probabilities. The probabilities are estimated for all subjects in the cohort.

survtime, left.time and continuous matching variables will be smoothed on while categorical matching variables are taken as factors.

Usage

```
GAMprob(survtime, samplestat, left.time = 0, match.var = 0, match.int = 0)
```

GAMprob

Arguments

survtime	Follow-up time for all cohort subjects
samplestat	A vector containing sampling and status information: 0 represents non-sampled subjects in the cohort, 1: sampled controls, 2,3, indicate different events. Cohort dimension.
left.time	Entry time if the survival times are left-truncated. Cohort dimension.
match.var	If the controls are matched to the cases (on other variables than time), match.var is the vector of matching variables. Cohort dimension.
match.int	A vector of length $2*$ number of matching variables. For caliper matching (matched on value pluss/minus epsilon) match.int should consist of c(-epsilon,epsilon). For exact matching match.int should consist of c(0,0).

Value

A vector of cohort dimension of sampling probabilities.

Author(s)

Nathalie C. Stoer

References

Stoer NC and Samuelsen SO (2013): Inverse probability weighting in nested case-control studies with additional matching - a simulation study. Statistics in Medicine, 32(30), 5328-5339.

See Also

```
wpl, coxph, Chenprob, GLMprob, KMprob, gam
```

Examples

```
data(CVD_Accidents)
attach(CVD_Accidents)
GAMprob(agestop,samplestat,agestart)
GAMprob(agestop,samplestat,agestop,match.var=cbind(sex,bmi),match.int=c(0,0,-2,2))
## The function is currently defined as
function (survtime, samplestat, left.time = 0, match.var = 0, match.int = 0)
{
    n.cohort = length(survtime)
    status = rep(0, n.cohort)
    status[samplestat > 1] = 1
    pgam = pGAM(status, survtime, samplestat, n.cohort, left.time)
    p = rep(1, n.cohort)
    p[status == 0] = pgam
    p
}
```

GLMprob 7

	GLMprob	Sampling probabilities estimated with logistic regression.
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Description

Estimates sampling probabilities with logistic regression. The weights included in the Cox-regressions (wpl) and which could be used for other procedures are inverse sampling probabilities i.e. the inverse of these probabilities. The probabilities are estimated for all subjects in the cohort.

survtime, left.time and continuous matching variables are included in the logistic regression as continuous variables while categorical matching variables are taken as factors.

Usage

```
GLMprob(survtime, samplestat, left.time = 0, match.var = 0, match.int = 0)
```

Arguments

survtime	Follow-up time for all cohort subjects
samplestat	A vector containing sampling and status information: 0 represents non-sampled subjects in the cohort, 1: sampled controls, 2,3, indicate different events. Cohort dimension.
left.time	Entry time if the survival times are left-truncated. Cohort dimension.
match.var	If the controls are matched to the cases (on other variables than time), match.var is the vector of matching variables. Cohort dimension.
match.int	A vector of length 2*number of matching variables. For caliper matching (matched on value pluss/minus epsilon) match.int should consist of c(-epsilon,epsilon). For exact matching match.int should consist of c(0,0).

Value

A vector of cohort dimension of sampling probabilities.

Author(s)

Nathalie C. Stoer

References

Stoer NC and Samuelsen SO (2013): Inverse probability weighting in nested case-control studies with additional matching - a simulation study. Statistics in Medicine, 32(30), 5328-5339.

See Also

```
wpl, coxph, Chenprob, GAMprob, KMprob
```

8 KMprob

Examples

```
data(CVD_Accidents)
attach(CVD_Accidents)
GLMprob(agestop, samplestat, agestart)
GLMprob(agestop, samplestat, agestart, match.var=cbind(sex, bmi), match.int=c(0,0,-2,2))
## The function is currently defined as
function (survtime, samplestat, left.time = 0, match.var = 0,
   match.int = 0)
{
   n.cohort = length(survtime)
   status = rep(0, n.cohort)
   status[samplestat > 1] = 1
   samplestat[samplestat > 1] = 1
   pglm = pGLM(status, survtime, samplestat, n.cohort, left.time,
       match.var, match.int)
   p = rep(1, n.cohort)
   p[status == 0] = pglm
 }
```

KMprob

Sampling probabilities estimated with a Kaplan-Meier type formula

Description

Estimates sampling probabilities with a Kaplan-Meier type formula. The weights included in the Cox-regressions (wpl) and which could be used for other procedures are inverse sampling probabilities i.e. the inverse of these probabilities. The probabilities are estimated for all subjects in the cohort.

Usage

```
KMprob(survtime, samplestat, m, left.time = 0, match.var = 0, match.int = 0)
```

Arguments

survtime	Follow-up time for all cohort subjects
samplestat	A vector containing sampling and status information: 0 represents non-sampled subjects in the cohort, 1: sampled controls, 2,3, indicate different events. Cohort dimension.
m	Number of sampled controls. A scalar if equal number of controls for all case. If unequal number of controls per case: A vector of length number of cases. The vector must be in the same order as the cases in the samplestat-vector.
left.time	Entry time if the survival times are left-truncated. Cohort dimension.
match.var	If the controls are matched to the cases (on other variables than time), match.var is the vector of matching variables. Cohort dimension.

KMprob 9

match.int

A vector of length 2*number of matching variables. For caliper matching (matched on value pluss/minus epsilon) match.int should consist of c(-epsilon,epsilon). For exact matching match.int should consist of c(0,0).

Value

A vector of cohort dimension of sampling probabilities.

Author(s)

Nathalie C. Stoer

References

Samuelsen SO. A pseudolikelihood approach to analysis of nested case-control studies. Biometrika, 84(2):379-394, 1997.

Stoer NC and Samuelsen SO (2013): Inverse probability weighting in nested case-control studies with additional matching - a simulation study. Statistics in Medicine, 32(30), 5328-5339.

See Also

```
wpl, coxph, Chenprob, GLMprob, GAMprob
```

Examples

```
data(CVD_Accidents)
attach(CVD_Accidents)
KMprob(agestop, samplestat, m=1, agestart)
KMprob(agestop,samplestat,m=1,agestart,match.var=cbind(bmi),match.int=c(-2,2))
## The function is currently defined as
function (survtime, samplestat, m, left.time = 0, match.var = 0, match.int = 0)
   n.cohort = length(survtime)
   status = rep(0, n.cohort)
   status[samplestat > 1] = 1
   o = order(survtime)
   status = status[o]
    survtime = survtime[o]
   if (length(left.time) == n.cohort) {
        left.time = left.time[o]
   if (length(match.var) == n.cohort) {
        match.var = match.var[o]
    if (length(match.var) > n.cohort) {
        match.var = match.var[o, ]
   tilbakestill = (1:n.cohort)[o]
   p = pKM(status, survtime, m, n.cohort, left.time, match.var,
        match.int)
   p[status > 0] = 1
```

pChen

```
p = p[order(tilbakestill)]
p
}
```

ModelbasedVar

Modelbased variance using Kaplan-Meier weights

Description

For internal use only

Author(s)

Nathalie C. Stoer and Sven Ove Samuelsen

 $\verb|multipleNCC-internal| Internal function|$

Description

Internal function

Author(s)

Nathalie C. Stoer

pChen

Chen-weights

Description

Estimates Chen-weights. For internal use only. Users should use the wrapper Chenprob.

Author(s)

Nathalie C. Stoer

See Also

wpl, Chenprob

pGAM

pGAM

Generlaized additive model weights

Description

Estimates GAM-weights. For internal use only. Users should use the wrapper GAMprob.

Author(s)

Nathalie C. Stoer

See Also

wpl, GAMprob, gam

pGLM

Logistic regression weights

Description

Estimates GLM-weights. For internal use only. Users should use the wrapper GLMprob.

Author(s)

Nathalie C. Stoer

See Also

wpl, GLMprob, glm

pKM

Kaplan-Meier weights

Description

Estimates Kaplan-Meier weights. For internal use only. Users should use the wrapper KMprob.

Author(s)

Nathalie C. Stoer

See Also

wpl, KMprob

12 print.wpl

PoststratVar

Modelbased variance using Chen-weights

Description

For internal use only.

Author(s)

Nathalie C. Stoer and Sven Ove Samuelsen

print.wpl

Print a wpl object

Description

Prints the fit of (each) weighted Cox-regression

Usage

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

Arguments

x The result of a call to wpl

... For future methods

Author(s)

Nathalie C. Stoer

See Also

wpl

summary.wpl 13

summary.wpl

Summary method for wpl

Description

produces a summary of a fitted wpl object

Usage

```
## S3 method for class 'wpl'
summary(object,...)
```

Arguments

object the result of a wpl fit
... for future methods

Author(s)

Nathalie C. Stoer

See Also

wpl

wpl

Weighted partial likelihood for nested case-control data

Description

Fits Cox proportional hazards models for nested case-control data with a weighted partial likelihood. Matching between cases and controls is broken which enables the controls to be reused for other endpoints. It handles competing risks (with simple survival data with one endpoint being a special case) and cases and controls from one endpoint are being used as additional controls for another endpoint. There are four choices of weights; Samuelsen (1997) KM, estimated with logistic regression (glm), logistic genralized additive model (gam) and local averaging (Chen, 2001) (Chen). KM, glm and gam handle additional matching, while all of them handle left-truncation.

Usage

```
wpl(x, data, samplestat, m = 1, weight.method = "KM", no.intervals = 10,
variance = "robust", no.intervals.left = c(3, 4), match.var = 0, match.int = 0)
```

14 wpl

Arguments

x A formula object, with the response on the left of a ~ operator, and the terms

on the right. The response must be a survival object as returned by the Surv function. The status variable going in to Surv is not actually used but should have 1 for cases and zero for controls and non-sampled subjects. All elements going into the formula should have length equal to the number of subjects in the cohort. Generally some of the covariates are not known for all subjects in the cohort (due to the NCC-sampling). The covariate values for those subjects should just be given some value e.g. 0 (not NA). Which value choosen is not

important as the values are never used.

data.frame in which to interpret the variables named in the formula.

samplestat A vector containing sampling and status information: 0 represents non-sampled

subjects in the cohort, 1: sampled controls, 2,3,... indicate different events.

Cohort dimension.

m Number of sampled controls. A scalar if equal number of controls for all cases.

If unequal number of controls per case: A vector of length number of cases. The

vector must be in the same order as the cases in the samplestat-vector.

weight.method Which weigths should be used, possibilities "KM", "gam", "glm", "Chen"

soring

variance Default is robust variances, but model based variance (only for KM-weights),

"Modelbased" and variance based on stratified case-cohort "Poststrat" (only for Chen-weights) is also possible. Pseudo-variance and Strat-variance will

appear under "est.se(coef)" in the output.

no.intervals.left

Number of intervals for Chen-weights with left-truncation. A vector on the form [number of intervals for left truncated time, number of intervals for survival

timel.

match.var If the controls are matched to the cases (on other variables than time), match.var

is the vector or matrix of matching variables. Cohort dimension.

match.int A vector of length 2*number of matching variables. For caliper matching (matched

on value pluss/minus epsilon) match.int should consist of c(-epsilon,epsilon).

For exact matching match int should consist of c(0,0).

Value

An object of class wpl representing the fit. Objects of this class have methods for the functions print and summary. The wpl-object consists of the following elements which are repeated for each endpoint. Unfortunately only the values for the first endpoint can be reached by \$-operator(ex. fit\$coefficients only return the coefficients for the first endpoint)

coefficients The vector of coefficients.

var Robust or estimated variance

weighted.loglik

A vector of length 2 containing the log-likelihood with the initial values and with the final values of the coefficients.

wpl 15

iter Number of iterations used

linear.predictors

The vector of linear predictors, one per subject. Note that this vector has been

centered, see predict.coxph for more details

residuals The martingale residuals

means Vector of column means of the X matrix

method The computation method used

The number of observations used in the fit

nevent The number of events (usually deaths) used in the fit

naive.var naive.var

rscore The robust log-rank statistic

wald.test The Wald test of whether the final coefficients differ from the initial values

y Inclusion time and event/censoring time

weights The vector of weights, which are inverse sampling probabilities

est.var Estimated variance (T) or robust variance (F)

.

Author(s)

Nathalie C. Stoer

References

Samuelsen SO. A pseudolikelihood approach to analysis of nested case-control studies. Biometrika, 84(2):379-394, 1997.

Stoer NC and Samuelsen SO (2013): Inverse probability weighting in nested case-control studies with additional matching - a simulation study. Statistics in Medicine, 32(30), 5328-5339.

See Also

coxph, Chenprob, GLMprob, GAMprob

Examples

```
data(CVD_Accidents)
wpl(Surv(agestart,agestop,dead24)~factor(smoking3gr)+bmi+factor(sex),data=CVD_Accidents,
samplestat=CVD_Accidents$samplestat,weight.method="gam")
wpl(Surv(agestart,agestop,dead24)~factor(smoking3gr)+bmi+factor(sex),data=CVD_Accidents,
samplestat=CVD_Accidents$samplestat,m=1,match.var=cbind(CVD_Accidents$sex,
```

 $CVD_Accidents\$bmi)$, match.int=c(0,0,-2,2), weight.method="glm")

16 wpl

```
## The function is currently defined as
function (x, data, samplestat, m = 1, weight.method = "KM", no.intervals = 10,
    variance = "robust", no.intervals.left = c(3, 4), match.var = 0,
    match.int = 0)
{
    UseMethod("wpl")
}
```

Index

```
* package
    multipleNCC-package, 2
Chenprob, 2, 3, 6, 7, 9, 10, 15
coxph, 2, 3, 6, 7, 9, 15
CVD_Accidents, 4
gam, 6, 11
GAMprob, 2, 3, 5, 7, 9, 11, 15
glm, 11
GLMprob, 2, 3, 6, 7, 9, 11, 15
KMprob, 2, 3, 6, 7, 8, 11
ModelbasedVar, 10
multipleNCC (multipleNCC-package), 2
{\tt multipleNCC-internal}, {\tt 10}
multipleNCC-package, 2
pChen, 10
pChen (multipleNCC-internal), 10
pGAM, 11
pGAM (multipleNCC-internal), 10
pGLM, 11
{\tt pGLM}\,({\tt multipleNCC-internal}),\, {\tt 10}
pKM (multipleNCC-internal), 10
PoststratVar, 12
print.wpl, 12
summary.wpl, 13
wpl, 2, 3, 6, 7, 9–13, 13
wpl.default (multipleNCC-internal), 10
wpl.formula(multipleNCC-internal), 10
wplEst(multipleNCC-internal), 10
```