Package ‘survival’
March 3, 2022

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LazyDataCompression xz
ByteCompile Yes
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Kaplan-Meier and Aalen-Johansen (multi-state) curves, Cox models,
and parametric accelerated failure time models.
License LGPL (>= 2)
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NeedsCompilation yes
Author Terry M Therneau [aut, cre],
Thomas Lumley [ctb, trl] (original S->R port and R maintainer until
2009),
Atkinson Elizabeth [ctb],
Crowson Cynthia [ctb]
Maintainer Terry M Therneau <therneau.terry@mayo.edu>
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aareg

Aalen's additive regression model for censored data

Description

Returns an object of class "aareg" that represents an Aalen model.

Usage

aareg(formula, data, weights, subset, na.action, 
qrtol=1e-07, nmin, dfbeta=FALSE, taper=1, 
test = c('aalen', 'variance', 'nrisk'), cluster, 
model=FALSE, x=FALSE, y=FALSE)
Arguments

formula: a formula object, with the response on the left of a `~` operator and the terms, separated by `+` operators, on the right. The response must be a `Surv` object. Due to a particular computational approach that is used, the model MUST include an intercept term. If `"-1"` is used in the model formula the program will ignore it.

data: data frame in which to interpret the variables named in the formula, subset, and weights arguments. This may also be a single number to handle some special cases – see below for details. If data is missing, the variables in the model formula should be in the search path.

weights: vector of observation weights. If supplied, the fitting algorithm minimizes the sum of the weights multiplied by the squared residuals (see below for additional technical details). The length of weights must be the same as the number of observations. The weights must be nonnegative and it is recommended that they be strictly positive, since zero weights are ambiguous. To exclude particular observations from the model, use the subset argument instead of zero weights.

subset: expression specifying which subset of observations should be used in the fit. This can be a logical vector (which is replicated to have length equal to the number of observations), a numeric vector indicating the observation numbers to be included, or a character vector of the observation names that should be included. All observations are included by default.

na.action: a function to filter missing data. This is applied to the model frame after any subset argument has been applied. The default is `na.fail`, which returns an error if any missing values are found. An alternative is `na.exclude`, which deletes observations that contain one or more missing values.

qrtol: tolerance for detection of singularity in the QR decomposition

nmin: minimum number of observations for an estimate; defaults to 3 times the number of covariates. This essentially truncates the computations near the tail of the data set, when n is small and the calculations can become numerically unstable.

dfbeta: should the array of dfbeta residuals be computed. This implies computation of the sandwich variance estimate. The residuals will always be computed if there is a cluster term in the model formula.

taper: allows for a smoothed variance estimate. $\text{Var}(x)$, where x is the set of covariates, is an important component of the calculations for the Aalen regression model. At any given time point t, it is computed over all subjects who are still at risk at time t. The tape argument allows smoothing these estimates, for example `taper=(1:4)/4` would cause the variance estimate used at any event time to be a weighted average of the estimated variance matrices at the last 4 death times, with a weight of 1 for the current death time and decreasing to 1/4 for prior event times. The default value gives the standard Aalen model.

test: selects the weighting to be used, for computing an overall “average” coefficient vector over time and the subsequent test for equality to zero.

cluster: the clustering group, optional. The variable will be searched for in the data argument.

model, x, y: should copies of the model frame, the x matrix of predictors, or the response vector y be included in the saved result.
Details

The Aalen model assumes that the cumulative hazard $H(t)$ for a subject can be expressed as $a(t) + X B(t)$, where $a(t)$ is a time-dependent intercept term, $X$ is the vector of covariates for the subject (possibly time-dependent), and $B(t)$ is a time-dependent matrix of coefficients. The estimates are inherently non-parametric; a fit of the model will normally be followed by one or more plots of the estimates.

The estimates may become unstable near the tail of a data set, since the increment to $B$ at time $t$ is based on the subjects still at risk at time $t$. The tolerance and/or $\text{nmin}$ parameters may act to truncate the estimate before the last death. The taper argument can also be used to smooth out the tail of the curve. In practice, the addition of a taper such as 1:10 appears to have little effect on death times when $n$ is still reasonably large, but can considerably dampen wild oscillations in the tail of the plot.

Value

an object of class "aareg" representing the fit, with the following components:

- **n**: vector containing the number of observations in the data set, the number of event times, and the number of event times used in the computation
- **times**: vector of sorted event times, which may contain duplicates
- **nrisk**: vector containing the number of subjects at risk, of the same length as times
- **coefficient**: matrix of coefficients, with one row per event and one column per covariate
- **test.statistic**: the value of the test statistic, a vector with one element per covariate
- **test.var**: variance-covariance matrix for the test
- **test**: the type of test; a copy of the test argument above
- **tweight**: matrix of weights used in the computation, one row per event
- **call**: a copy of the call that produced this result

References


See Also

print.aareg, summary.aareg, plot.aareg

Examples

# Fit a model to the lung cancer data set
lfit <- aareg(Surv(time, status) ~ age + sex + ph.ecog, data=lung, nmin=1)

## Not run:
lfit
call:
aareg(formula = Surv(time, status) ~ age + sex + ph.ecog, data = lung, nmin = 1)
n=227 (1 observations deleted due to missing values)
138 out of 138 unique event times used

<table>
<thead>
<tr>
<th>slope</th>
<th>coef</th>
<th>se(coef)</th>
<th>z</th>
<th>p</th>
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<td>3.80e-03</td>
<td>1.03e-03</td>
<td>3.70</td>
</tr>
</tbody>
</table>

Chisq=26.73 on 3 df, p=6.7e-06; test weights=aalen

plot(lfit[4], ylim=c(-4,4))  # Draw a plot of the function for ph.ecog

## End(Not run)

lfit2 <- aareg(Surv(time, status) ~ age + sex + ph.ecog, data=lung, nmin=1, taper=1:10)
## Not run: lines(lfit2[4], col=2)  # Nearly the same, until the last point

# A fit to the multiple-infection data set of children with
# Chronic Granulomatous Disease. See section 8.5 of Therneau and Grambsch.
fita2 <- aareg(Surv(tstart, tstop, status) ~ treat + age + inherit + steroids + cluster(id), data=cgd)
## Not run:

n= 203
69 out of 70 unique event times used

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<td>0.013200</td>
<td>0.010600</td>
<td>0.009700</td>
<td>1.36</td>
</tr>
</tbody>
</table>

Chisq=16.74 on 4 df, p=0.0022; test weights=aalen

## End(Not run)

---

**Description**

The check for tied survival times can fail due to floating point imprecision, which can make actual ties appear to be distinct values. Routines that depend on correct identification of ties pairs will then give incorrect results, e.g., a Cox model. This function rectifies these.

**Usage**

```r
aeqSurv(x, tolerance = sqrt(.Machine$double.eps))
```
Arguments

- `x` a Surv object
- `tolerance` the tolerance used to detect values that will be considered equal

Details

This routine is called by both `survfit` and `coxph` to deal with the issue of ties that get incorrectly broken due to floating point imprecision. See the short vignette on tied times for a simple example. Use the `timefix` argument of `survfit` or `coxph.control` to control the option if desired.

The rule for ‘equality’ is identical to that used by the `all.equal` routine. Pairs of values that are within round off error of each other are replaced by the smaller value. An error message is generated if this process causes a 0 length time interval to be created.

Value

a Surv object identical to the original, but with ties restored.

Author(s)

Terry Therneau

See Also

`survfit`, `coxph.control`

---

**aggregate.survfit**  
*Average survival curves*

Description

For a survfit object containing multiple curves, create average curves over a grouping.

Usage

```r
## S3 method for class 'survfit'
aggregate(x, by = NULL, FUN = mean, ...)
```

Arguments

- `x` a survfit object which has a data dimension.
- `by` an optional list or vector of grouping elements, each as long as `dim(x)[data]`.
- `FUN` a function to compute the summary statistic of interest.
- `...` optional further arguments to `FUN`.  

---
agreg.fit

Details

The primary use of this is to take an average over multiple survival curves that were created from a modeling function. That is, a marginal estimate of the survival. It is primarily used to average over multiple predicted curves from a Cox model.

Value

a survfit object of lower dimension.

See Also

survfit

Examples

cfit <- coxph(Surv(futime, death) ~ sex + age*hgb, data=mgus2)
# marginal effect of sex, after adjusting for the others
dummy <- rbind(mgus2, mgus2)
dummy$sex <- rep(c("F", "M"), each=nrow(mgus2)) # population data set
dummy <- na.omit(dummy)  # don't count missing hgb in our "population
csurv <- survfit(cfit, newdata=dummy)
dim(csurv) # 2 * 1384 survival curves
csurv2 <- aggregate(csurv, dummy$sex)

agreg.fit

Cox model fitting functions

Description

These are the the functions called by coxph that do the actual computation. In certain situations, e.g. a simulation, it may be advantageous to call these directly rather than the usual coxph call using a model formula.

Usage

agreg.fit(x, y, strata, offset, init, control, weights, method, rownames, resid=TRUE, nocenter=NULL)
coxph.fit(x, y, strata, offset, init, control, weights, method, rownames, resid=TRUE, nocenter=NULL)

Arguments

x  Matix of predictors. This should not include an intercept.
y  a Surv object containing either 2 columns (coxph.fit) or 3 columns (agreg.fit).
strata  a vector containing the stratification, or NULL
offset  optional offset vector
init  initial values for the coefficients
function.

control the result of a call to coxph.control
weights optional vector of weights
method method for handling ties, one of "breslow" or "efron"
rownames this is only needed for a NULL model, in which case it contains the rownames (if any) of the original data.
resid compute and return residuals.
nocenter an optional list of values. Any column of the X matrix whose values lie strictly within that set will not be recentered. Note that the coxph function has (-1, 0, 1) as the default.

Details
This routine does no checking that arguments are the proper length or type. Only use it if you know what you are doing!
The resid and concordance arguments will save some compute time for calling routines that only need the likelihood, the generation of a permutation distribution for instance.

Value
a list containing results of the fit

Author(s)
Terry Therneau

See Also
coxph

---

**aml**

Acute Myelogenous Leukemia survival data

Description
Survival in patients with Acute Myelogenous Leukemia. The question at the time was whether the standard course of chemotherapy should be extended ("maintainance") for additional cycles.

Usage
```r
aml
leukemia
data(cancer, package="survival")
```
Format

time: survival or censoring time
status: censoring status
x: maintenance chemotherapy given? (factor)

Source


## S3 method for class 'coxph'

`anova(object, ..., test = 'Chisq')`

Arguments

- `object`: An object of class `coxph`
- `...`: Further `coxph` objects
- `test`: a character string. The appropriate test is a chisquare, all other choices result in no test being done.

Details

Specifying a single object gives a sequential analysis of deviance table for that fit. That is, the reductions in the model Cox log-partial-likelihood as each term of the formula is added in turn are given in as the rows of a table, plus the log-likelihoods themselves. A robust variance estimate is normally used in situations where the model may be mis-specified, e.g., multiple events per subject. In this case a comparison of likelihood values does not make sense (differences no longer have a chi-square distribution), and `anova` will refuse to print results.

If more than one object is specified, the table has a row for the degrees of freedom and loglikelihood for each model. For all but the first model, the change in degrees of freedom and loglik is also given. (This only make statistical sense if the models are nested.) It is conventional to list the models from smallest to largest, but this is up to the user.

The table will optionally contain test statistics (and P values) comparing the reduction in loglik for each row.
Value

An object of class "anova" inheriting from class "data.frame".

Warning

The comparison between two or more models by anova will only be valid if they are fitted to the same dataset. This may be a problem if there are missing values.

See Also

coxph, anova.

Examples

```r
fit <- coxph(Surv(futime, fustat) ~ resid.ds * rx + ecog.ps, data = ovarian)
anova(fit)
fit2 <- coxph(Surv(futime, fustat) ~ resid.ds + rx + ecog.ps, data=ovarian)
anova(fit2,fit)
```

---

attrassign

Create new-style "assign" attribute

Description

The "assign" attribute on model matrices describes which columns come from which terms in the model formula. It has two versions. R uses the original version, but the alternate version found in S-plus is sometimes useful.

Usage

```r
## Default S3 method:
attrassign(object, tt,...)
## S3 method for class 'lm'
attrassign(object,...)
```

Arguments

- **object**: model matrix or linear model object
- **tt**: terms object
- **...**: ignored
Details

For instance consider the following

\[
\text{survreg}(\text{Surv}(\text{time, status}) \sim \text{age + sex + factor(ph.ecog)}, \text{lung})
\]

R gives the compact for for assign, a vector (0, 1, 2, 3, 3, 3); which can be read as “the first column of the X matrix (intercept) goes with none of the terms, the second column of X goes with term 1 of the model equation, the third column of X with term 2, and columns 4-6 with term 3”.

The alternate (S-Plus default) form is a list

\[
\begin{align*}
$(\text{Intercept}) & : 1 \\
$\text{age} & : 2 \\
$\text{sex} & : 3 \\
$\text{factor(ph.ecog)} & : 4, 5, 6
\end{align*}
\]

Value

A list with names corresponding to the term names and elements that are vectors indicating which columns come from which terms

See Also

\text{terms, model.matrix}

Examples

\begin{verbatim}
formula <- Surv(time,status)~factor(ph.ecog)
tt <- terms(formula)
mf <- model.frame(tt,data=lung)
mm <- model.matrix(tt,mf)
## a few rows of data
mm[1:3,]
## old-style assign attribute
attr(mm,"assign")
## alternate style assign attribute
attrassign(mm,tt)
\end{verbatim}

basehaz

Alias for the survfit function

Description

Compute the predicted survival curve for a Cox model.

Usage

\begin{verbatim}
basehaz(fit, centered=TRUE)
\end{verbatim}
Arguments

fit  a coxph fit
centered  if TRUE return data from a predicted survival curve at the mean values of the covariates fit$mean, if FALSE return a prediction for all covariates equal to zero.

Details

This function is simply an alias for survfit, which does the actual work and has a richer set of options. The alias exists only because some users look for predicted survival estimates under this name.

The function returns a data frame containing the time, cumhaz and optionally the strata (if the fitted Cox model used a strata statement), which are copied the survfit result. If there are factor variables in the model, then the default predictions at the "mean" are meaningless since they do not correspond to any possible subject; correct results require use of the newdata argument of survfit. Results for all covariates =0 are normally only of use as a building block for further calculations.

Value

a data frame with variable names of hazard, time and optionally strata. The first is actually the cumulative hazard.

See Also

survfit.coxph

---

Bladder Cancer Recurrences

Description

Data on recurrences of bladder cancer, used by many people to demonstrate methodology for recurrent event modelling.

Bladder1 is the full data set from the study. It contains all three treatment arms and all recurrences for 118 subjects; the maximum observed number of recurrences is 9.

Bladder is the data set that appears most commonly in the literature. It uses only the 85 subjects with nonzero follow-up who were assigned to either thiotepa or placebo, and only the first four recurrences for any patient. The status variable is 1 for recurrence and 0 for everything else (including death for any reason). The data set is laid out in the competing risks format of the paper by Wei, Lin, and Weissfeld.

Bladder2 uses the same subset of subjects as bladder, but formatted in the (start, stop] or Anderson-Gill style. Note that in transforming from the WLW to the AG style data set there is a quite common programming mistake that leads to extra follow-up time for 12 subjects: all those with follow-up beyond their 4th recurrence. This "follow-up" is a side effect of throwing away all events after the fourth while retaining the last follow-up time variable from the original data. The bladder2 data set
found here does not make this mistake, but some analyses in the literature have done so; it results in the addition of a small amount of immortal time bias and shrinks the fitted coefficients towards zero.

**Usage**

```r
bladder1
bladder
bladder2
data(cancer, package="survival")
```

**Format**

**bladder1**

```
id: Patient id
treatment: Placebo, pyridoxine (vitamin B6), or thiotepa
number: Initial number of tumours (8=8 or more)
size: Size (cm) of largest initial tumour
recur: Number of recurrences
start, stop: The start and end time of each time interval
status: End of interval code, 0=censored, 1=recurrence,
        2=death from bladder disease, 3=death other/unknown cause
rtumor: Number of tumors found at the time of a recurrence
rsize: Size of largest tumor at a recurrence
enum: Event number (observation number within patient)
```

**bladder**

```
id: Patient id
rx: Treatment 1=placebo 2=thiotepa
number: Initial number of tumours (8=8 or more)
size: size (cm) of largest initial tumour
stop: recurrence or censoring time
enum: which recurrence (up to 4)
```

**bladder2**

```
id: Patient id
rx: Treatment 1=placebo 2=thiotepa
number: Initial number of tumours (8=8 or more)
size: size (cm) of largest initial tumour
start: start of interval (0 or previous recurrence time)
stop: recurrence or censoring time
enum: which recurrence (up to 4)
```
**Source**

---

**blogit**

### Bounded link functions

**Description**
Alternate link functions that impose bounds on the input of their link function

**Usage**
```r
blogit(edge = 0.05)
bprobit(edge = 0.05)
bcloglog(edge = 0.05)
blog(edge = 0.05)
```

**Arguments**
- `edge`: input values less than the cutpoint are replaced with the cutpoint. For all be `blog` input values greater than (1-edge) are replaced with (1-edge)

**Details**
When using survival psuedovalues for binomial regression, the raw data can be outside the range (0,1), yet we want to restrict the predicted values to lie within that range. A natural way to deal with this is to use `glm` with `family = gaussian(link = "logit")`. But this will fail. The reason is that the `family` object has a component `linkfun` that does not accept values outside of (0,1).

This function is only used to create initial values for the iteration step, however. Mapping the offending input argument into the range of (edge, 1-edge) before computing the link results in starting estimates that are good enough. The final result of the fit will be no different than if explicit starting estimates were given using the `etastart` or `mustart` arguments. These functions create copies of the logit, probit, and complimentary log-log families that differ from the standard ones only in this use of a bounded input argument, and are called a "bounded logit" = `blogit`, etc.

The same argument hold when using RMST (area under the curve) pseudovalues along with a log link to ensure positive predictions, though in this case only the lower boundary needs to be mapped.

**Value**
a family object of the same form as `make.family`.

**See Also**
```r
stats[make.family]
```
Examples

```r
py <- pseudo(survfit(Surv(time, status) ~ 1, lung), time=730) # 2 year survival
range(py)
pfit <- glm(py ~ ph.ecog, data=lung, family=gaussian(link=logit()))
# For each +1 change in performance score, the odds of 2 year survival
# are multiplied by 1/2 = exp of the coefficient.
```

**cch**  
*Fits proportional hazards regression model to case-cohort data*

**Description**

Returns estimates and standard errors from relative risk regression fit to data from case-cohort studies. A choice is available among the Prentice, Self-Prentice and Lin-Ying methods for unstratified data. For stratified data the choice is between Borgan I, a generalization of the Self-Prentice estimator for unstratified case-cohort data, and Borgan II, a generalization of the Lin-Ying estimator.

**Usage**

```r
cch(formula, data, subcoh, id, stratum=NULL, cohort.size, 
    robust=FALSE)
```

**Arguments**

- `formula`: A formula object that must have a `Surv` object as the response. The `Surv` object must be of type "right", or of type "counting".
- `subcoh`: Vector of indicators for subjects sampled as part of the sub-cohort. Code 1 or TRUE for members of the sub-cohort, 0 or FALSE for others. If `data` is a data frame then `subcoh` may be a one-sided formula.
- `id`: Vector of unique identifiers, or formula specifying such a vector.
- `stratum`: A vector of stratum indicators or a formula specifying such a vector.
- `cohort.size`: Vector with size of each stratum original cohort from which subcohort was sampled.
- `data`: An optional data frame in which to interpret the variables occurring in the formula.
- `method`: Three procedures are available. The default method is "Prentice", with options for "SelfPrentice" or "LinYing".
- `robust`: For "LinYing" only, if robust=TRUE, use design-based standard errors even for phase I.
Details

Implements methods for case-cohort data analysis described by Therneau and Li (1999). The three methods differ in the choice of "risk sets" used to compare the covariate values of the failure with those of others at risk at the time of failure. "Prentice" uses the sub-cohort members "at risk" plus the failure if that occurs outside the sub-cohort and is score unbiased. "SelfPren" (Self-Prentice) uses just the sub-cohort members "at risk". These two have the same asymptotic variance-covariance matrix. "LinYing" (Lin-Ying) uses the all members of the sub-cohort and all failures outside the sub-cohort who are "at risk". The methods also differ in the weights given to different score contributions.

The data argument must not have missing values for any variables in the model. There must not be any censored observations outside the subcohort.

Value

An object of class "cch" incorporating a list of estimated regression coefficients and two estimates of their asymptotic variance-covariance matrix.

- **coef**: regression coefficients.
- **naive.var**: Self-Prentice model based variance-covariance matrix.
- **var**: Lin-Ying empirical variance-covariance matrix.

Author(s)

Norman Breslow, modified by Thomas Lumley

References


See Also
twophase and svycoxph in the "survey" package for more general two-phase designs. [http://faculty.washington.edu/tlumley/survey/](http://faculty.washington.edu/tlumley/survey/)
Examples

## The complete Wilms Tumor Data
## (Breslow and Chatterjee, Applied Statistics, 1999)
## subcohort selected by simple random sampling.
##
## subcoh <- nwtco$in.subcohort
selccoh <- with(nwtco, rel==1|subcoh==1)
ccoh.data <- nwtco[selccoh,]
ccoh.data$subcohort <- subcoh[selccoh]
## central-lab histology
ccoh.data$histol <- factor(ccoh.data$histol,labels=c("FH","UH"))
## tumour stage
ccoh.data$stage <- factor(ccoh.data$stage,labels=c("I","II","III","IV"))
ccoh.data$age <- ccoh.data$age/12 # Age in years

## Standard case-cohort analysis: simple random subcohort
##
fit.ccP <- cch(Surv(edrel, rel) ~ stage + histol + age, data = ccoh.data, 
               subcoh = ~subcohort, id=~seqno, cohort.size=4028)

fit.ccP

fit.ccSP <- cch(Surv(edrel, rel) ~ stage + histol + age, data = ccoh.data, 
               subcoh = ~subcohort, id=~seqno, cohort.size=4028, method="SelfPren")

summary(fit.ccSP)

## (post-)stratified on instit
##
stratsizes<-table(nwtco$instit)
fit.BI<- cch(Surv(edrel, rel) ~ stage + histol + age, data = ccoh.data, 
             subcoh = ~subcohort, id=~seqno, stratum=~instit, cohort.size=stratsizes, 
             method="I.Borgan")

summary(fit.BI)

---

cgd

Chronic Granulotamous Disease data

Description

Data are from a placebo controlled trial of gamma interfere in chronic granulotomous disease (CGD). Contains the data on time to serious infections observed through end of study for each patient.
Usage

cgd
data(cgd)

Format

<table>
<thead>
<tr>
<th>Variable</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>id</td>
<td>subject identification number</td>
</tr>
<tr>
<td>center</td>
<td>enrolling center</td>
</tr>
<tr>
<td>random</td>
<td>date of randomization</td>
</tr>
<tr>
<td>treatment</td>
<td>placebo or gamma interferon</td>
</tr>
<tr>
<td>sex</td>
<td>sex</td>
</tr>
<tr>
<td>age</td>
<td>age in years, at study entry</td>
</tr>
<tr>
<td>height</td>
<td>height in cm at study entry</td>
</tr>
<tr>
<td>weight</td>
<td>weight in kg at study entry</td>
</tr>
<tr>
<td>inherit</td>
<td>pattern of inheritance</td>
</tr>
<tr>
<td>steroids</td>
<td>use of steroids at study entry, 1=yes</td>
</tr>
<tr>
<td>propylac</td>
<td>use of prophylactic antibiotics at study entry</td>
</tr>
<tr>
<td>hos.cat</td>
<td>a categorization of the centers into 4 groups</td>
</tr>
<tr>
<td>tstart, tstop</td>
<td>start and end of each time interval</td>
</tr>
<tr>
<td>status</td>
<td>1=the interval ends with an infection</td>
</tr>
<tr>
<td>enum</td>
<td>observation number within subject</td>
</tr>
</tbody>
</table>

Details

The cgd0 data set is in the form found in the references, with one line per patient and no recoding of the variables. The cgd data set (this one) has been cast into (start, stop] format with one line per event, and covariates such as center recoded as factors to include meaningful labels.

Source

Fleming and Harrington, Counting Processes and Survival Analysis, appendix D.2.

See Also

link{cgd0}
**Chronic Granulotomous Disease data**

**Description**
Data are from a placebo controlled trial of gamma interferon in chronic granulotomous disease (CGD). Contains the data on time to serious infections observed through end of study for each patient.

**Usage**
cgd0

**Format**

<table>
<thead>
<tr>
<th>Variable</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>id</td>
<td>subject identification number</td>
</tr>
<tr>
<td>center</td>
<td>enrolling center</td>
</tr>
<tr>
<td>random</td>
<td>date of randomization</td>
</tr>
<tr>
<td>treatment</td>
<td>placebo or gamma interferon</td>
</tr>
<tr>
<td>sex</td>
<td>sex</td>
</tr>
<tr>
<td>age</td>
<td>age in years, at study entry</td>
</tr>
<tr>
<td>height</td>
<td>height in cm at study entry</td>
</tr>
<tr>
<td>weight</td>
<td>weight in kg at study entry</td>
</tr>
<tr>
<td>inherit</td>
<td>pattern of inheritance</td>
</tr>
<tr>
<td>steroids</td>
<td>use of steroids at study entry, 1=yes</td>
</tr>
<tr>
<td>propylac</td>
<td>use of prophylactic antibiotics at study entry</td>
</tr>
<tr>
<td>hos.cat</td>
<td>a categorization of the centers into 4 groups</td>
</tr>
<tr>
<td>futime</td>
<td>days to last follow-up</td>
</tr>
<tr>
<td>etime1-etime7</td>
<td>up to 7 infection times for the subject</td>
</tr>
</tbody>
</table>

**Details**
The cgd raw data set (this one) is in the form found in the references, with one line per patient and no recoding of the variables.
The cgd data set has been further processed so as to have one line per event, with covariates such as center recoded as factors to include meaningful labels.

**Source**
Fleming and Harrington, Counting Processes and Survival Analysis, appendix D.2.

**See Also**
cgd
cipoisson  

**Description**  
Confidence interval calculation for Poisson rates.

**Usage**  
cipoisson(k, time = 1, p = 0.95, method = c("exact", "anscombe"))

**Arguments**  
- **k**  
  Number of successes  
- **time**  
  Total time on trial  
- **p**  
  Probability level for the (two-sided) interval  
- **method**  
  The method for computing the interval.

**Details**  
The likelihood method is based on equation 10.10 of Feller, which relates poisson probabilities to tail area of the gamma distribution. The Anscombe approximation is based on the fact that sqrt(k + 3/8) has a nearly constant variance of 1/4, along with a continuity correction.

There are many other proposed intervals: Patil and Kulkarni list and evaluate 19 different suggestions from the literature!. The exact intervals can be overly broad for very small values of k, many of the other approaches try to shrink the lengths, with varying success.

**Value**  
a vector, matrix, or array. If both k and time are single values the result is a vector of length 2 containing the lower an upper limits. If either or both are vectors the result is a matrix with two columns. If k is a matrix or array, the result will be an array with one more dimension; in this case the dimensions and dimnames (if any) of k are preserved.

**References**  


**See Also**  
ppois, qpois
clogit

Examples

cipoisson(4) # 95\% confidence limit
# lower   upper
# 1.089865 10.24153
ppois(4, 10.24153) # chance of seeing 4 or fewer events with large rate
# [1] 0.02500096
1-ppois(3, 1.08986) # chance of seeing 4 or more, with a small rate
# [1] 0.02499961

clogit

Conditional logistic regression

Description

Estimates a logistic regression model by maximising the conditional likelihood. Uses a model
formula of the form case.status~exposure+strata(matched.set). The default is to use the
exact conditional likelihood, a commonly used approximate conditional likelihood is provided for
compatibility with older software.

Usage

clogit(formula, data, weights, subset, na.action,
method=c("exact", "approximate", "efron", "breslow"),
...)

Arguments

formula  Model formula
data     data frame
weights  optional, names the variable containing case weights
subset   optional, subset the data
na.action optional na.action argument. By default the global option na.action is used.
method   use the correct (exact) calculation in the conditional likelihood or one of the
approximations
...      optional arguments, which will be passed to coxph.control

Details

It turns out that the loglikelihood for a conditional logistic regression model = loglik from a Cox
model with a particular data structure. Proving this is a nice homework exercise for a PhD statistics
class; not too hard, but the fact that it is true is surprising.

When a well tested Cox model routine is available many packages use this 'trick' rather than writing
a new software routine from scratch, and this is what the clogit routine does. In detail, a stratified
Cox model with each case/control group assigned to its own stratum, time set to a constant, status
of 1=case 0=control, and using the exact partial likelihood has the same likelihood formula as a conditional logistic regression. The clogit routine creates the necessary dummy variable of times (all 1) and the strata, then calls coxph.

The computation of the exact partial likelihood can be very slow, however. If a particular strata had say 10 events out of 20 subjects we have to add up a denominator that involves all possible ways of choosing 10 out of 20, which is \(20!/(10!\cdot10!) = 184756\) terms. Gail et al describe a fast recursion method which partly ameliorates this; it was incorporated into version 2.36-11 of the survival package. The computation remains infeasible for very large groups of ties, say 100 ties out of 500 subjects, and may even lead to integer overflow for the subscripts – in this latter case the routine will refuse to undertake the task. The Efron approximation is normally a sufficiently accurate substitute.

Most of the time conditional logistic modeling is applied data with 1 case + k controls per set, in which case all of the approximations for ties lead to exactly the same result. The ’approximate’ option maps to the Breslow approximation for the Cox model, for historical reasons. Case weights are not allowed when the exact option is used, as the likelihood is not defined for fractional weights. Even with integer case weights it is not clear how they should be handled. For instance if there are two deaths in a strata, one with weight=1 and one with weight=2, should the likelihood calculation consider all subsets of size 2 or all subsets of size 3? Consequently, case weights are ignored by the routine in this case.

Value

An object of class "clogit", which is a wrapper for a "coxph" object.

References


Author(s)

Thomas Lumley

See Also

strata, coxph, glm

Examples

```r
## Not run: clogit(case ~ spontaneous + induced + strata(stratum), data=infert)
# A multinomial response recoded to use clogit
# The revised data set has one copy per possible outcome level, with new
# variable tocc = target occupation for this copy, and case = whether
# that is the actual outcome for each subject.
# See the reference below for the data.
resp <- levels(logan$occupation)
n <- nrow(logan)
```
cluster <- rep(1:n, length(resp))
logan2 <- data.frame(logan[indx,],
    id = indx,
    tocc = factor(rep(resp, each=n)))
logan2$case <- (logan2$occupation == logan2$tocc)
clogit(case ~ tocc + tocc:education + strata(id), logan2)

cluster
Identify clusters.

Description
This is a special function used in the context of survival models. It identifies correlated groups of observations, and is used on the right hand side of a formula. This style is now discouraged, use the cluster option instead.

Usage
cluster(x)

Arguments
x
A character, factor, or numeric variable.

Details
The function’s only action is semantic, to mark a variable as the cluster indicator. The resulting variance is what is known as the “working independence” variance in a GEE model. Note that one cannot use both a frailty term and a cluster term in the same model, the first is a mixed-effects approach to correlation and the second a GEE approach, and these don’t mix.

Value
x

See Also
coxph, survreg

Examples
marginal.model <- coxph(Surv(time, status) ~ rx, data= rats, cluster=litter, subset=(sex=='f'))
frailty.model <- coxph(Surv(time, status) ~ rx + frailty(litter), rats, subset=(sex=='f'))
Chemotherapy for Stage B/C colon cancer

Description

These are data from one of the first successful trials of adjuvant chemotherapy for colon cancer. Levamisole is a low-toxicity compound previously used to treat worm infestations in animals; 5-FU is a moderately toxic (as these things go) chemotherapy agent. There are two records per person, one for recurrence and one for death.

Usage

colon
data(cancer, package="survival")

Format

id: id
study: 1 for all patients
rx: Treatment - Observation, Lev(amisole), Lev(amisole)+5-FU
sex: 1=male
age: in years
obstruct: obstruction of colon by tumour
perfor: perforation of colon
adhere: adherence to nearby organs
nodes: number of lymph nodes with detectable cancer
time: days until event or censoring
status: censoring status
differ: differentiation of tumour (1=well, 2=moderate, 3=poor)
extent: Extent of local spread (1=submucosa, 2=muscle, 3=serosa, 4=contiguous structures)
surg: time from surgery to registration (0=short, 1=long)
node4: more than 4 positive lymph nodes
etype: event type: 1=recurrence, 2=death

Note

The study is originally described in Laurie (1989). The main report is found in Moertel (1990). This data set is closest to that of the final report in Moertel (1991). A version of the data with less follow-up time was used in the paper by Lin (1994).

Peter Higgins has pointed out a data inconsistency, revealed by `table(colon$nodes, colon$node4)`.

We don’t know which of the two variables is actually correct so have elected not to ’fix’ it. (Real data has warts, why not have some in the example data too?)
References


concordance  
Compute the concordance statistic for data or a model

Description

The concordance statistic compute the agreement between an observed response and a predictor. It is closely related to Kendall’s tau-a and tau-b, Goodman’s gamma, and Somers’ d, all of which can also be calculated from the results of this function.

Usage

concordance(object, ...)  
## S3 method for class 'formula'
concordance(object, data, weights, subset, na.action,  
cluster, ymin, ymax, timewt= c("n", "S", "S/G", "n/G", "n/G2", "I"),  
influence=0, ranks = FALSE, reverse=FALSE, timefix=TRUE, keepstrata=10, ...)
## S3 method for class 'lm'
concordance(object, ..., newdata, cluster, ymin, ymax,  
influence=0, ranks=FALSE, timefix=TRUE, keepstrata=10)
## S3 method for class 'coxph'
concordance(object, ..., newdata, cluster, ymin, ymax,  
timewt= c("n", "S", "S/G", "n/G", "n/G2", "I"), influence=0,  
ranks=FALSE, timefix=FALSE, keepstrata=10)
## S3 method for class 'survreg'
concordance(object, ..., newdata, cluster, ymin, ymax,  
timewt= c("n", "S", "S/G", "n/G", "n/G2", "I"), influence=0,  
ranks=FALSE, timefix=FALSE, keepstrata=10)
Arguments

- **object**: a fitted model or a formula. The formula should be of the form $y \sim x$ or $y \sim x + \text{strata}(z)$ with a single numeric or survival response and a single predictor. Counts of concordant, discordant and tied pairs are computed separately per stratum, and then added.

- **data**: a data.frame in which to interpret the variables named in the formula, or in the subset and the weights argument. Only applicable if object is a formula.

- **weights**: optional vector of case weights. Only applicable if object is a formula.

- **subset**: expression indicating which subset of the rows of data should be used in the fit. Only applicable if object is a formula.

- **na.action**: a missing-data filter function. This is applied to the model.frame after any subset argument has been used. Default is `options()$na.action`. Only applicable if object is a formula.

- **...**: multiple fitted models are allowed. Only applicable if object is a model object.

- **newdata**: optional, a new data frame in which to evaluate (but not refit) the models

- **cluster**: optional grouping vector for calculating the robust variance

- **ymin**, **ymax**: compute the concordance over the restricted range $ymin \leq y \leq ymax$. (For survival data this is a time range.)

- **timewt**: the weighting to be applied. The overall statistic is a weighted mean over event times.

- **influence**: 1= return the dfbeta vector, 2= return the full influence matrix, 3 = return both

- **ranks**: if TRUE, return a data frame containing the individual ranks that make up the overall score.

- **reverse**: if TRUE then assume that larger x values predict smaller response values y; a proportional hazards model is the common example of this.

- **timefix**: if the response is a Surv object, correct for possible rounding error; otherwise this argument has no effect. See the vignette on tied times for more explanation. For the coxph and survreg methods this issue will have already been addressed in the parent routine, so should not be revisited.

- **keepstrata**: either TRUE, FALSE, or an integer value. Computations are always done within stratum, then added. If the total number of strata greater than `keepstrata`, or `keepstrata`=FALSE, those subtotals are not kept in the output.

Details

At each event time, compute the rank of the subject who had the event as compared to all others with a longer survival, where the rank is value between 0 and 1. The concordance is a weighted mean of these values, determined by the timewt option. For uncensored data each unique response value is compared to all those which are larger.

Using the default value for timewt gives the area under the receiver operating curve (AUC) for a binary response, and $(d+1)/2$ when $y$ is continuous, where $d$ is Somers’ d. For a survival time, timewt of n gives Harrell’s c-statistic, which is closely related to the Gehan-Wilcoxon test, $S$ corresponds to the Peto-Wilcoxon, $n/G2$ is the weighted advocated by Umo, and $S/G$ the weighting proposed by Schemper.
When the number of strata is very large, such as in a conditional logistic regression for instance (clogit function), a much faster computation is available when the individual strata results are not retained. In the more general case the keepstrata = 10 default simply keeps the printout manageable.

**Value**

An object of class concordance containing the following components:

- **concordance**: the estimated concordance value or values
- **count**: a vector containing the number of concordant pairs, discordant, tied on x but not y, tied on y but not x, and tied on both x and y
- **n**: the number of observations
- **var**: a vector containing the estimated variance of the concordance based on the infinitesimal jackknife (IJ) method. If there are multiple models it contains the estimated variance/covariance matrix.
- **cvar**: a vector containing the estimated variance(s) of the concordance values, based on the variance formula for the associated score test from a proportional hazards model. (This was the primary variance used in the survConcordance function.)
- **dfbeta**: optional, the vector of leverage estimates for the concordance
- **influence**: optional, the matrix of leverage values for each of the counts, one row per observation
- **ranks**: optional, a data frame containing the Somers’ d rank at each event time, along with the time weight, case weight of the observation with an event, and variance (contribution to the proportional hazards model information matrix). A weighted mean of the ranks equals Somer’s d.

**Author(s)**

Terry Therneau

**See Also**

coxph

**Examples**

```r
fit1 <- coxph(Surv(ptime, pstat) ~ age + sex + mspike, mgus2)
concordance(fit1, timewt="n")

# logistic regression
fit2 <- glm(pstat ~ age + sex + mspike, binomial, data= mgus2)
concordance(fit2) # equal to the AUC
```
**Description**

This is the working routine behind the `concordance` function. It is not meant to be called by users, but is available for other packages to use. Input arguments, for instance, are assumed to all be the correct length and type, and missing values are not allowed: the calling routine is responsible for these things.

**Usage**

```r
cordancefit(y, x, strata, weights, ymin = NULL, ymax = NULL, timewt = c("n", "S", "S/G", "n/G", "n/G2", "I"), cluster, influence = 0, ranks = FALSE, reverse = FALSE, timefix = TRUE, keepstrata=10, robustse = TRUE)
```

**Arguments**

- `y`: the response. It can be numeric, factor, or a Surv object
- `x`: the predictor, a numeric vector
- `strata`: optional numeric vector that stratifies the data
- `weights`: options vector of case weights
- `ymin, ymax`: restrict the comparison to response values in this range
- `timewt`: the time weighting to be used
- `cluster, influence, ranks, reverse, timefix`: see the help for the `concordance` function
- `keepstrata`: either TRUE, FALSE, or an integer value. Computations are always done within stratum, then added. If the total number of strata greater than keepstrata, or keepstrata=FALSE, those subtotals are not kept in the output.
- `robustse`: compute the robust standard error

**Value**

A list containing the results

**Author(s)**

Terry Therneau

**See Also**

`concordance`
Test the Proportional Hazards Assumption of a Cox Regression

**Description**

Test the proportional hazards assumption for a Cox regression model fit (coxph).

**Usage**

```r
cox.zph(fit, transform="km", terms=TRUE, singledf=FALSE, global=TRUE)
```

**Arguments**

- `fit`  
  the result of fitting a Cox regression model, using the `coxph` or `coxme` functions.

- `transform`  
  a character string specifying how the survival times should be transformed before the test is performed. Possible values are "km", "rank", "identity" or a function of one argument.

- `terms`  
  if TRUE, do a test for each term in the model rather than for each separate covariate. For a factor variable with k levels, for instance, this would lead to a k-1 degree of freedom test. The plot for such variables will be a single curve evaluating the linear predictor over time.

- `singledf`  
  use a single degree of freedom test for terms that have multiple coefficients, i.e., the test that corresponds most closely to the plot. If `terms=FALSE` this argument has no effect.

- `global`  
  should a global chi-square test be done, in addition to the per-variable or per-term tests tests.

**Details**

The computations require the original x matrix of the Cox model fit. Thus it saves time if the `x=TRUE` option is used in `coxph`. This function would usually be followed by both a plot and a print of the result. The plot gives an estimate of the time-dependent coefficient $\beta(t)$. If the proportional hazards assumption holds then the true $\beta(t)$ function would be a horizontal line. The table component provides the results of a formal score test for slope=0, a linear fit to the plot would approximate the test.

Random effects terms such a frailty or random effects in a `coxme` model are not checked for proportional hazards, rather they are treated as a fixed offset in model.

If the model contains strata by covariate interactions, then the y matrix may contain structural zeros, i.e., deaths (rows) that had no role in estimation of a given coefficient (column). These are marked as NA. If an entire row is NA, for instance after subscripting a `cox.zph` object, that row is removed.
**Value**

an object of class "cox.zph", with components:

- **table**: a matrix with one row for each variable, and optionally a last row for the global test. Columns of the matrix contain a score test of for addition of the time-dependent term, the degrees of freedom, and the two-sided p-value.
- **x**: the transformed time axis.
- **time**: the untransformed time values; there is one entry for each event time in the data
- **strata**: for a stratified coxph model, the stratum of each of the events
- **y**: the matrix of scaled Schoenfeld residuals. There will be one column per term or per variable (depending on the terms option above), and one row per event. The row labels are a rounded form of the original times.
- **var**: a variance matrix for the covariates, used to create an approximate standard error band for plots
- **transform**: the transform of time that was used
- **call**: the calling sequence for the routine.

**Note**

In versions of the package before survival3.0 the function computed a fast approximation to the score test. Later versions compute the actual score test.

**References**


**See Also**

- coxph, Surv.

**Examples**

```r
fit <- coxph(Surv(futime, fustat) ~ age + ecog.ps, data=ovarian)
temp <- cox.zph(fit)
print(temp)  # display the results
plot(temp)   # plot curves
```
**coxph**  

*Fit Proportional Hazards Regression Model*

**Description**

Fits a Cox proportional hazards regression model. Time dependent variables, time dependent strata, multiple events per subject, and other extensions are incorporated using the counting process formulation of Andersen and Gill.

**Usage**

```r
coxph(formula, data=, weights, subset,  
    na.action, init, control,  
    ties=c("efron","breslow","exact"),  
    singular.ok=TRUE, robust,  
    model=FALSE, x=FALSE, y=TRUE, tt, method=ties,  
    id, cluster, istate, statedata, nocenter=c(-1, 0, 1), ...)
```

**Arguments**

- `formula` 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.
- `data` a data.frame in which to interpret the variables named in the `formula`, or in the `subset` and the `weights` argument.
- `weights` vector of case weights, see the note below. For a thorough discussion of these see the book by Therneau and Grambsch.
- `subset` expression indicating which subset of the rows of `data` should be used in the fit. All observations are included by default.
- `na.action` a missing-data filter function. This is applied to the model.frame after any subset argument has been used. Default is `options()$na.action`.
- `init` vector of initial values of the iteration. Default initial value is zero for all variables.
- `control` Object of class `coxph.control` specifying iteration limit and other control options. Default is `coxph.control(...)`.  
- `ties` a character string specifying the method for tie handling. If there are no tied death times all the methods are equivalent. Nearly all Cox regression programs use the Breslow method by default, but not this one. The Efron regression programs use the Breslow method by default, but not this one. The Efron approximation is used as the default here, it is more accurate when dealing with tied death times, and is as efficient computationally. The “exact partial likelihood” is equivalent to a conditional logistic model, and is appropriate when the times are a small set of discrete values. See further below.
logical value indicating how to handle collinearity in the model matrix. If TRUE, the program will automatically skip over columns of the X matrix that are linear combinations of earlier columns. In this case the coefficients for such columns will be NA, and the variance matrix will contain zeros. For ancillary calculations, such as the linear predictor, the missing coefficients are treated as zeros.

robust should a robust variance be computed. The default is TRUE if: there is a cluster argument, there are case weights that are not 0 or 1, or there are id values with more than one event.

id optional variable name that identifies subjects. Only necessary when a subject can have multiple rows in the data, and there is more than one event type. This variable will normally be found in data.

cluster optional variable which clusters the observations, for the purposes of a robust variance. If present, it implies robust. This variable will normally be found in data.

istate optional variable giving the current state at the start each interval. This variable will normally be found in data.

statedata optional data set used to describe multistate models.

model logical value: if TRUE, the model frame is returned in component model.

x logical value: if TRUE, the x matrix is returned in component x.

y logical value: if TRUE, the response vector is returned in component y.

tt optional list of time-transform functions.

method alternate name for the ties argument.

nocenter columns of the X matrix whose values lie strictly within this set are not recentered. Remember that a factor variable becomes a set of 0/1 columns.

... Other arguments will be passed to coxph.control

Details

The proportional hazards model is usually expressed in terms of a single survival time value for each person, with possible censoring. Andersen and Gill reformulated the same problem as a counting process; as time marches onward we observe the events for a subject, rather like watching a Geiger counter. The data for a subject is presented as multiple rows or "observations", each of which applies to an interval of observation (start, stop].

The routine internally scales and centers data to avoid overflow in the argument to the exponential function. These actions do not change the result, but lead to more numerical stability. Any column of the X matrix whose values lie within nocenter list are not recentered. The practical consequence of the default is to not recenter dummy variables corresponding to factors. However, arguments to offset are not scaled since there are situations where a large offset value is a purposefully used. In general, however, users should not avoid very large numeric values for an offset due to possible loss of precision in the estimates.

Value

an object of class coxph representing the fit. See coxph.object for details.
Side Effects

Depending on the call, the predict, residuals, and survfit routines may need to reconstruct the x matrix created by coxph. It is possible for this to fail, as in the example below in which the predict function is unable to find tform.

```r
tfun <- function(tform) coxph(tform, data=lung)
  fit <- tfun(Surv(time, status) ~ age)
  predict(fit)
```

In such a case add the model=TRUE option to the coxph call to obviate the need for reconstruction, at the expense of a larger fit object.

Case weights

Case weights are treated as replication weights, i.e., a case weight of 2 is equivalent to having 2 copies of that subject’s observation. When computers were much smaller grouping like subjects together was a common trick to used to conserve memory. Setting all weights to 2 for instance will give the same coefficient estimate but halve the variance. When the Efron approximation for ties (default) is employed replication of the data will not give exactly the same coefficients as the weights option, and in this case the weighted fit is arguably the correct one.

When the model includes a cluster term or the robust=TRUE option the computed variance treats any weights as sampling weights; setting all weights to 2 will in this case give the same variance as weights of 1.

Special terms

There are three special terms that may be used in the model equation. A strata term identifies a stratified Cox model; separate baseline hazard functions are fit for each strata. The cluster term is used to compute a robust variance for the model. The term + cluster(id) where each value of id is unique is equivalent to specifying the robust=TRUE argument. If the id variable is not unique, it is assumed that it identifies clusters of correlated observations. The robust estimate arises from many different arguments and thus has had many labels. It is variously known as the Huber sandwich estimator, White’s estimate (linear models/econometrics), the Horvitz-Thompson estimate (survey sampling), the working independence variance (generalized estimating equations), the infinitesimal jackknife, and the Wei, Lin, Weissfeld (WLW) estimate.

A time-transform term allows variables to vary dynamically in time. In this case the tt argument will be a function or a list of functions (if there are more than one tt() term in the model) giving the appropriate transform. See the examples below.

One user mistake that has recently arisen is to slavishly follow the advice of some coding guides and prepend survival:: onto everything, including the special terms, e.g., survival::coxph(survival:Surv(time,status) ~ age + survival::cluster(inst),data=lung) First, this is unnecessary: arguments within the coxph call will be evaluated within the survival namespace, so another package’s Surv or cluster function would not be noticed. (Full qualification of the coxph call itself may be protective, however.) Second, and more importantly, the call just above will not give the correct answer. The specials are recognized by their name, and survival::cluster is not the same as cluster; the above model would treat inst as an ordinary variable. A similar issue arises from using stats::offset as a term, in either survival or glm models.
Convergence

In certain data cases the actual MLE estimate of a coefficient is infinity, e.g., a dichotomous variable where one of the groups has no events. When this happens the associated coefficient grows at a steady pace and a race condition will exist in the fitting routine: either the log likelihood converges, the information matrix becomes effectively singular, an argument to exp becomes too large for the computer hardware, or the maximum number of interactions is exceeded. (Most often number 1 is the first to occur.) The routine attempts to detect when this has happened, not always successfully. The primary consequence for the user is that the Wald statistic = coefficient/se(coefficient) is not valid in this case and should be ignored; the likelihood ratio and score tests remain valid however.

Ties

There are three possible choices for handling tied event times. The Breslow approximation is the easiest to program and hence became the first option coded for almost all computer routines. It then ended up as the default option when other options were added in order to "maintain backwards compatibility". The Efron option is more accurate if there are a large number of ties, and it is the default option here. In practice the number of ties is usually small, in which case all the methods are statistically indistinguishable.

Using the "exact partial likelihood" approach the Cox partial likelihood is equivalent to that for matched logistic regression. (The clogit function uses the coxph code to do the fit.) It is technically appropriate when the time scale is discrete and has only a few unique values, and some packages refer to this as the "discrete" option. There is also an "exact marginal likelihood" due to Prentice which is not implemented here.

The calculation of the exact partial likelihood is numerically intense. Say for instance 180 subjects are at risk on day 7 of which 15 had an event; then the code needs to compute sums over all 180-choose-15 > 10^43 different possible subsets of size 15. There is an efficient recursive algorithm for this task, but even with this the computation can be insufferably long. With (start, stop) data it is much worse since the recursion needs to start anew for each unique start time.

A separate issue is that of artificial ties due to floating-point imprecision. See the vignette on this topic for a full explanation or the timefix option in coxph.control. Users may need to add timefix=FALSE for simulated data sets.

Penalized regression

coxph can maximise a penalised partial likelihood with arbitrary user-defined penalty. Supplied penalty functions include ridge regression (ridge), smoothing splines (pspline), and frailty models (frailty).

References


See Also

coxph.object, coxph.control, cluster, strata, Surv, survfit, pspline, ridge.
Examples

# Create the simplest test data set
test1 <- list(time=c(4,3,1,1,2,2,3),
    status=c(1,1,0,1,1,0),
    x=c(0,2,1,1,1,0,0),
    sex=c(0,0,0,0,1,1,1))
# Fit a stratified model
coxph(Surv(time, status) ~ x + strata(sex), test1)

# Create a simple data set for a time-dependent model
test2 <- list(start=c(1,2,5,2,1,7,3,4,8,8),
    stop=c(2,3,6,7,8,9,9,9,14,17),
    event=c(1,1,1,1,1,1,1,0,0,0),
    x=c(1,0,0,1,0,1,1,1,0,0))
summary(coxph(Surv(start, stop, event) ~ x, test2))

# Create a simple data set for a time-dependent model
#
test2 <- list(start=c(1, 2, 5, 2, 1, 7, 3, 4, 8, 8),
    stop =c(2, 3, 6, 7, 8, 9, 9, 9,14,17),
    event=c(1, 1, 1, 1, 1, 1, 1, 0, 0, 0),
    x =c(1, 0, 0, 1, 0, 1, 1, 1, 0, 0) )

summary(coxph( Surv(start, stop, event) ~ x, test2))

# Fit a stratified model, clustered on patients
bladder1 <- bladder[bladder$enum < 5, ]
coxph(Surv(stop, event) ~ (rx + size + number) * strata(enum),
    cluster = id, bladder1)

# Fit a time transform model using current age
coxph(Surv(time, status) ~ ph.ecog + tt(age), data=lung,
    tt=function(x,t,...) pspline(x + t/365.25))

---

**coxph.control**  
Ancillary arguments for controlling coxph fits

**Description**

This is used to set various numeric parameters controlling a Cox model fit. Typically it would only be used in a call to coxph.

**Usage**

```r
coxph.control(eps = 1e-09, toler.chol = .Machine$double.eps^0.75,
    iter.max = 20, toler.inf = sqrt(eps), outer.max = 10, timefix=TRUE)
```
Arguments

eps
Iteration continues until the relative change in the log partial likelihood is less than eps, or the absolute change is less than sqrt(eps). Must be positive.

toler.chol
Tolerance for detection of singularity during a Cholesky decomposition of the variance matrix, i.e., for detecting a redundant predictor variable.

iter.max
Maximum number of iterations to attempt for convergence.

toler.inf
Tolerance criteria for the warning message about a possible infinite coefficient value.

outer.max
For a penalized coxph model, e.g. with pspline terms, there is an outer loop of iteration to determine the penalty parameters; maximum number of iterations for this outer loop.

timefix
Resolve any near ties in the time variables.

Details

The convergence tolerances are a balance. Users think they want THE maximum point of the likelihood surface, and for well behaved data sets where this is quadratic near the max a high accuracy is fairly inexpensive: the number of correct digits approximately doubles with each iteration. Conversely, a drop of .0001 from the maximum in any given direction will be correspond to only about 1/20 of a standard error change in the coefficient. Statistically, more precision than this is straining at a gnat. Based on this the author originally had set the tolerance to 1e-5, but relented in the face of multiple "why is the answer different than package X" queries.

Asking for results that are too close to machine precision (double.eps) is a fool's errand; a reasonable criteria is often the square root of that precision. The Cholesky decompostion needs to be held to a higher standard than the overall convergence criterion, however. The toler.inf value controls a warning message; if it is too small incorrect warnings can appear, if too large some actual cases of an infinite coefficient will not be detected.

The most difficult cases are data sets where the MLE coefficient is infinite; an example is a data set where at each death time, it was the subject with the largest covariate value who perished. In that situation the coefficient increases at each iteration while the log-likelihood asymptotes to a maximum. As iteration proceeds there is a race condition condition for three endpoint: exp(coef) overflows, the Hessian matrix become singular, or the change in loglik is small enough to satisfy the convergence criterion. The first two are difficult to anticipate and lead to numeric difficulties, which is another argument for moderation in the choice of eps.

See the vignette "Roundoff error and tied times" for a more detailed explanation of the timefix option. In short, when time intervals are created via subtraction then two time intervals that are actually identical can appear to be different due to floating point round off error, which in turn can make coxph and survfit results dependent on things such as the order in which operations were done or the particular computer that they were run on. Such cases are unfortunately not rare in practice. The timefix=TRUE option adds logic similar to all.equal to ensure reliable results. In analysis of simulated data sets, however, where often by defintion there can be no duplicates, the option will often need to be set to FALSE to avoid spurious merging of close numeric values.

Value

a list containing the values of each of the above constants
**Description**

Returns the individual contributions to the first and second derivative matrix, at each unique event time.

**Usage**

```r
coxph.detail(object, riskmat=FALSE, rorder=c("data", "time"))
```

**Arguments**

- `object`: a Cox model object, i.e., the result of `coxph`.
- `riskmat`: include the at-risk indicator matrix in the output?
- `rorder`: only applicable if `riskmat=TRUE`. Should the rows of riskmat be in the original data order, or sorted by time within strata.

**Details**

This function may be useful for those who wish to investigate new methods or extensions to the Cox model. The example below shows one way to calculate the Schoenfeld residuals.

**Value**

a list with components

- `time`: the vector of unique event times
- `nevent`: the number of events at each of these time points.
- `means`: a matrix with one row for each event time and one column for each variable in the Cox model, containing the weighted mean of the variable at that time, over all subjects still at risk at that time. The weights are the risk weights \( \exp(x \%*% fit$coef) \).
- `nrisk`: number of subjects at risk.
- `score`: the contribution to the score vector (first derivative of the log partial likelihood) at each time point.
- `imat`: the contribution to the information matrix (second derivative of the log partial likelihood) at each time point.
- `hazard`: the hazard increment. Note that the hazard and variance of the hazard are always for some particular future subject. This routine uses `object$mean` as the future subject.
varhaz: the variance of the hazard increment.

x, y: copies of the input data.

strata: only present for a stratified Cox model, this is a table giving the number of time points of component time that were contributed by each of the strata.

riskmat: a matrix with one row for each observation and one column for each unique event time, containing a 0/1 value to indicate whether that observation was (1) or was not (0) at risk at the given time point. Rows are in the order of the original data (after removal of any missings by coxph), or in time order.

See Also

coxph, residuals.coxph

Examples

```r
fit <- coxph(Surv(futime, fustat) ~ age + rx + ecog.ps, ovarian, x=TRUE)
fitd <- coxph.detail(fit)
# There is one Schoenfeld residual for each unique death. It is a
# vector (covariates for the subject who died) - (weighted mean covariate
# vector at that time). The weighted mean is defined over the subjects
# still at risk, with exp(X beta) as the weight.

events <- fit$y[,2]==1
etime <- fit$y[events,1] # the event times --- may have duplicates
indx <- match(etime, fitd$time)
schoen <- fit$x[events,] - fitd$means[indx,]
```

Description

This class of objects is returned by the coxph class of functions to represent a fitted proportional hazards model. Objects of this class have methods for the functions print, summary, residuals, predict and survfit.

Arguments

coefficients: the vector of coefficients. If the model is over-determined there will be missing values in the vector corresponding to the redundant columns in the model matrix.

var: the variance matrix of the coefficients. Rows and columns corresponding to any missing coefficients are set to zero.

naive.var: this component will be present only if the robust option was true. If so, the var component will contain the robust estimate of variance, and this component will contain the ordinary estimate.

loglik: a vector of length 2 containing the log-likelihood with the initial values and with the final values of the coefficients.
score value of the efficient score test, at the initial value of the coefficients.
rscore the robust log-rank statistic, if a robust variance was requested.
wald.test the Wald test of whether the final coefficients differ from the initial values.
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 values used as the reference for each covariate. For instance, a later
call to predict(fit,type='risk') will give the hazard ratio between an ob-
servation and this reference. (For most covariates this will contain the mean.)
n the number of observations used in the fit.
event the number of events (usually deaths) used in the fit.
concordance a vector of length 6, containing the number of pairs that are concordant, discor-
dant, tied on x, tied on y, and tied on both, followed by the standard error of the
concordance statistic.
first the first derivative vector at the solution.
weights the vector of case weights, if one was used.
method the method used for handling tied survival times.
na.action the na.action attribute, if any, that was returned by the na.action routine.
timefix the value of the timefix option used in the fit
cmp the coefficient map, present for multi-state coxph fits. There a column for each
transition and a row for each coefficient, the value maps that transition/coefficient
pair to a position in the coefficient vector. If a particular covariate is not used by
the transition the matrix will contain a zero, if two transitions share a coefficient
the matrix will contain repeats.
stratum_map stratum mapping, present for multi-state coxph fits. The row labeled ‘(Baseline)’
identifies transitions that do or do not share a baseline stratum. Further rows
correspond to strata() terms in the model, each of which may apply to some
transitions and not others.
...
The object will also contain the following, for documentation see the lm object:
terms, assign, formula, call, and, optionally, x, y, and/or frame.

Components

The following components must be included in a legitimate coxph object.

See Also

coxph, coxph.detail, cox.zph, residuals.coxph, survfit, survreg.
**coxph.wtest**  
*Compute a quadratic form*

**Description**
This function is used internally by several survival routines. It computes a simple quadratic form, while properly dealing with missings.

**Usage**
```r
coxph.wtest(var, b, toler.chol = 1e-09)
```

**Arguments**
- `var`: variance matrix
- `b`: vector
- `toler.chol`: tolerance for the internal cholesky decomposition

**Details**
Compute b' V-inverse b. Equivalent to sum(b * solve(V,b)), except for the case of redundant covariates in the original model, which lead to NA values in V and b.

**Value**
a real number

**Author(s)**
Terry Therneau

---

**coxsurv.fit**  
*A direct interface to the ‘computational engine’ of survfit.coxph*

**Description**
This program is mainly supplied to allow other packages to invoke the survfit.coxph function at a ‘data’ level rather than a ‘user’ level. It does no checks on the input data that is provided, which can lead to unexpected errors if that data is wrong.

**Usage**
```r
coxsurv.fit(ctype, stype, se.fit, varmat, cluster,  
y, x, wt, risk, position, strata, oldid,  
y2, x2, risk2, strata2, id2, unlist=TRUE)
```
Arguments

- **stype**: survival curve computation: 1=direct, 2=exp(-cumulative hazard)
- **ctype**: cumulative hazard computation: 1=Breslow, 2=Efron
- **se.fit**: if TRUE, compute standard errors
- **varmat**: the variance matrix of the coefficients
- **cluster**: vector to control robust variance
- **y**: the response variable used in the Cox model. (Missing values removed of course.)
- **x**: covariate matrix used in the Cox model
- **wt**: weight vector for the Cox model. If the model was unweighted use a vector of 1s.
- **risk**: the risk score exp(X beta + offset) from the fitted Cox model.
- **position**: optional argument controlling what is counted as ‘censored’. Due to time dependent covariates, for instance, a subject might have start, stop times of (1,5),(5,30),(30,100). Times 5 and 30 are not ‘real’ censorings. Position is 1 for a real start, 2 for an actual end, 3 for both, 0 for neither.
- **strata**: strata variable used in the Cox model. This will be a factor.
- **oldid**: identifier for subjects with multiple rows in the original data.
- **y2, x2, risk2, strata2**: variables for the hypothetical subjects, for which prediction is desired
- **id2**: optional; if present and not NULL this should be a vector of identifiers of length nrow(x2). A non-null value signifies that x2 contains time dependent covariates, in which case this identifies which rows of x2 go with each subject.
- **unlist**: if FALSE the result will be a list with one element for each strata. Otherwise the strata are “unpacked” into the form found in a survfit object.

Value

A list containing nearly all the components of a survfit object. All that is missing is to add the confidence intervals, the type of the original model’s response (as in a coxph object), and the class.

Note

The source code for both this function and survfit.coxph is written using noweb. For complete documentation see the inst/sourcecode.pdf file.

Author(s)

Terry Therneau

See Also

survfit.coxph
**Description**

Partial results from a trial of laser coagulation for the treatment of diabetic retinopathy.

**Usage**

```r
diabetic
data(diabetic, package="survival")
```

**Format**

A data frame with 394 observations on the following 8 variables.

- `id`: subject id
- `laser`: laser type: xenon or argon
- `age`: age at diagnosis
- `eye`: a factor with levels of `left` `right`
- `trt`: treatment: 0 = no treatment, 1 = laser
- `risk`: risk group of 6-12
- `time`: time to event or last follow-up
- `status`: status of 0 = censored or 1 = visual loss

**Details**

The 197 patients in this dataset were a 50% random sample of the patients with "high-risk" diabetic retinopathy as defined by the Diabetic Retinopathy Study (DRS). Each patient had one eye randomized to laser treatment and the other eye received no treatment. For each eye, the event of interest was the time from initiation of treatment to the time when visual acuity dropped below 5/200 two visits in a row. Thus there is a built-in lag time of approximately 6 months (visits were every 3 months). Survival times in this dataset are therefore the actual time to blindness in months, minus the minimum possible time to event (6.5 months). Censoring was caused by death, dropout, or end of the study.

**References**

Huster, Brookmeyer and Self, Biometrics, 1989.

**Examples**

```r
# juvenile diabetes is defined as and age less than 20
juvenile <- 1*(diabetic$age < 20)
coxph(Surv(time, status) ~ trt + juvenile, cluster= id, 
    data= diabetic)
```
Distributions available in \texttt{survreg}.

\textbf{Description}

Density, cumulative distribution function, quantile function and random generation for the set of distributions supported by the \texttt{survreg} function.

\textbf{Usage}

\begin{verbatim}
\texttt{dsurvreg(x, mean, scale=1, distribution='weibull', parms)}
\texttt{psurvreg(q, mean, scale=1, distribution='weibull', parms)}
\texttt{qsurvreg(p, mean, scale=1, distribution='weibull', parms)}
\texttt{rsurvreg(n, mean, scale=1, distribution='weibull', parms)}
\end{verbatim}

\textbf{Arguments}

\begin{itemize}
  \item \texttt{x} \hspace{1cm} vector of quantiles. Missing values (\texttt{NA}s) are allowed.
  \item \texttt{q} \hspace{1cm} vector of quantiles. Missing values (\texttt{NA}s) are allowed.
  \item \texttt{p} \hspace{1cm} vector of probabilities. Missing values (\texttt{NA}s) are allowed.
  \item \texttt{n} \hspace{1cm} number of random deviates to produce
  \item \texttt{mean} \hspace{1cm} vector of linear predictors for the model. This is replicated to be the same length as \texttt{p}, \texttt{q} or \texttt{n}.
  \item \texttt{scale} \hspace{1cm} vector of (positive) scale factors. This is replicated to be the same length as \texttt{p}, \texttt{q} or \texttt{n}.
  \item \texttt{distribution} \hspace{1cm} character string giving the name of the distribution. This must be one of the elements of \texttt{survreg.distributions}
  \item \texttt{parms} \hspace{1cm} optional parameters, if any, of the distribution. For the t-distribution this is the degrees of freedom.
\end{itemize}

\textbf{Details}

Elements of \texttt{q} or \texttt{p} that are missing will cause the corresponding elements of the result to be missing.

The location and scale values are as they would be for \texttt{survreg}. The label "mean" was an unfortunate choice (made in mimicry of \texttt{qnorm}); since almost none of these distributions are symmetric it will not actually be a mean, but corresponds instead to the linear predictor of a fitted model. Translation to the usual parameterization found in a textbook is not always obvious. For example, the Weibull distribution is fit using the Extreme value distribution along with a log transformation. Letting $F(t) = 1 - \exp[-(at)^p]$ be the cumulative distribution of the Weibull using a standard parameterization in terms of $a$ and $p$, the \texttt{survreg} location corresponds to $-\log(a)$ and the scale to $1/p$ (Kalbfleisch and Prentice, section 2.2.2).

\textbf{Value}

density (\texttt{dsurvreg}), probability (\texttt{psurvreg}), quantile (\texttt{qsurvreg}), or for the requested distribution with mean and scale parameters \texttt{mean} and \texttt{sd}. 

References

See Also
survreg, Normal

Examples

```r
# List of distributions available
names(survreg.distributions)
## Not run:
[1] "extreme"  "logistic"  "gaussian"  "weibull"  "exponential"
[6] "rayleigh"  "loggaussian" "lognormal"  "loglogistic"  "t"
## End(Not run)
# Compare results
all.equal(dsurvreg(1:10, 2, 5, dist="lognormal"), dlnorm(1:10, 2, 5))

# Hazard function for a Weibull distribution
x <- seq(.1, 3, length=30)
haz <- dsurvreg(x, 2, 3)/(1-psurvreg(x, 2, 3))
## Not run:
plot(x, haz, log="xy", ylab="Hazard") #line with slope (1/scale -1)
## End(Not run)
```

finegray

*Create data for a Fine-Gray model*

Description

The Fine-Gray model can be fit by first creating a special data set, and then fitting a weighted Cox model to the result. This routine creates the data set.

Usage

```r
finegray(formula, data, weights, subset, na.action= na.pass, etype,
prefix="fg", count, id, timefix=TRUE)
```

Arguments

- `formula`: a standard model formula, with survival on the left and covariates on the right.
- `data`: an optional data frame, list or environment (or object coercible by `as.data.frame to a data frame) containing the variables in the model.
- `weights`: optional vector of observation weights
subset an optional vector specifying a subset of observations to be used in the fitting process.

na.action a function which indicates what should happen when the data contain NAs. The default is set by the na.action setting of options.

etype the event type for which a data set will be generated. The default is to use whichever is listed first in the multi-state survival object.

prefix the routine will add 4 variables to the data set: a start and end time for each interval, status, and a weight for the interval. The default names of these are "fgstart", "fgstop", "fgstatus", and "fgwt"; the prefix argument determines the initial portion of the new names.

count a variable name in the output data set for an optional variable that will contain the the replication count for each row of the input data. If a row is expanded into multiple lines it will contain 1, 2, etc.

id optional, the variable name in the data set which identifies subjects.

timfix process times through the aeqSurv function to eliminate potential roundoff issues.

Details

The function expects a multi-state survival expression or variable as the left hand side of the formula, e.g. Surv(atime, astat) where astat is a factor whose first level represents censoring and remaining levels are states. The output data set will contain simple survival data (status = 0 or 1) for a single endpoint of interest. In the output data set subjects who did not experience the event of interest become censored subjects whose times are artificially extended over multiple intervals, with a decreasing case weight from interval to interval. The output data set will normally contain many more rows than the input.

Time dependent covariates are allowed, but not (currently) delayed entry. If there are time dependent covariates, e.g., the input data set had Surv(entry, exit, stat) as the left hand side, then an id statement is required. The program does data checks in this case, and needs to know which rows belong to each subject.

The output data set will often have gaps. Say that there were events at time 50 and 100 (and none between) and censoring at 60, 70, and 80. Formally, a non event subjects at risk from 50 to 100 will have different weights in each of the 3 intervals 50-60, 60-70, and 80-100, but because the middle interval does not span any event times the subsequent Cox model will never use that row. The finegray output omits such rows.

See the competing risks vignette for more details.

Value

a data frame

Author(s)

Terry Therneau
References


See Also

coxph, aeqSurv

Examples

# Treat time to death and plasma cell malignancy as competing risks
etime <- with(mgus2, ifelse(pstat==0, futime, ptime))
event <- with(mgus2, ifelse(pstat==0, 2*death, 1))
event <- factor(event, 0:2, labels=c("censor", "pcm", "death"))

# FG model for PCM
pdata <- finegray(Surv(etime, event) ~ ., data=mgus2)
fgfit <- coxph(Surv(fgstart, fgstop, fgstatus) ~ age + sex,
weight=fgwt, data=pdata)

# Compute the weights separately by sex
adata <- finegray(Surv(etime, event) ~ . + strata(sex),
data=mgus2, na.action=na.pass)

flchain

Assay of serum free light chain for 7874 subjects.

Description

This is a stratified random sample containing 1/2 of the subjects from a study of the relationship between serum free light chain (FLC) and mortality. The original sample contains samples on approximately 2/3 of the residents of Olmsted County aged 50 or greater.

Usage

flchain
data(flchain, package="survival")

Format

A data frame with 7874 persons containing the following variables.
age age in years
sex F=female, M=male
sample.yr the calendar year in which a blood sample was obtained
flchain

kappa  serum free light chain, kappa portion
lambda  serum free light chain, lambda portion
flc.grp  the FLC group for the subject, as used in the original analysis
creatinine  serum creatinine
mgus  1 if the subject had been diagnosed with monoclonal gammopathy (MGUS)
futime  days from enrollment until death. Note that there are 3 subjects whose sample was obtained on their death date.
death  0=alive at last contact date, 1=dead
chapter  for those who died, a grouping of their primary cause of death by chapter headings of the International Code of Diseases ICD-9

Details

In 1995 Dr. Robert Kyle embarked on a study to determine the prevalence of monoclonal gammopathy of undetermined significance (MGUS) in Olmsted County, Minnesota, a condition which is normally only found by chance from a test (serum electrophoresis) which is ordered for other causes. Later work suggested that one component of immunoglobulin production, the serum free light chain, might be a possible marker for immune disregulation. In 2010 Dr. Angela Dispenzieri and colleagues assayed FLC levels on those samples from the original study for which they had patient permission and from which sufficient material remained for further testing. They found that elevated FLC levels were indeed associated with higher death rates.

Patients were recruited when they came to the clinic for other appointments, with a final random sample of those who had not yet had a visit since the study began. An interesting side question is whether there are differences between early, mid, and late recruits.

This data set contains an age and sex stratified random sample that includes 7874 of the original 15759 subjects. The original subject identifiers and dates have been removed to protect patient identity. Subsampling was done to further protect this information.

Source

The primary investigator (A Dispenzieri) and statistician (T Therneau) for the study.

References


Examples

data(flchain)
age.grp <- cut(flchain$age, c(49,54, 59,64, 69,74,79, 89, 110),
labels= paste(c(50,55,60,65,70,75,80,90),
  c(54,59,64,69,74,79,89,109), sep='-''))
table(flchain$sex, age.grp)
Description

The frailty function allows one to add a simple random effects term to a Cox model.

Usage

```r
frailty(x, distribution="gamma", ...) 
frailty.gamma(x, sparse = (nclass > 5), theta, df, eps = 1e-05, 
method = c("em","aic", "df", "fixed"), ...) 
frailty.gaussian(x, sparse = (nclass > 5), theta, df, 
method =c("reml","aic", "df", "fixed"), ...) 
frailty.t(x, sparse = (nclass > 5), theta, df, eps = 1e-05, tdf = 5, 
method = c("aic", "df", "fixed"), ...)
```

Arguments

- `x`: the variable to be entered as a random effect. It is always treated as a factor.
- `distribution`: either the gamma, gaussian or t distribution may be specified. The routines `frailty.gamma`, `frailty.gaussian` and `frailty.t` do the actual work.
- `...`: Arguments for specific distribution, including (but not limited to)
- `sparse`: cutoff for using a sparse coding of the data matrix. If the total number of levels of `x` is larger than this value, then a sparse matrix approximation is used. The correct cutoff is still a matter of exploration: if the number of levels is very large (thousands) then the non-sparse calculation may not be feasible in terms of both memory and compute time. Likewise, the accuracy of the sparse approximation appears to be related to the maximum proportion of subjects in any one class, being best when no one class has a large membership.
- `theta`: if specified, this fixes the variance of the random effect. If not, the variance is a parameter, and a best solution is sought. Specifying this implies `method='fixed'`.
- `df`: if specified, this fixes the degrees of freedom for the random effect. Specifying this implies `method='df'`. Only one of `theta` or `df` should be specified.
- `method`: the method used to select a solution for `theta`, the variance of the random effect. The `fixed` corresponds to a user-specified value, and no iteration is done. The `df` selects the variance such that the degrees of freedom for the random effect matches a user specified value. The `aic` method seeks to maximize Akaike’s information criteria $2^*(\text{partial likelihood} - \text{df})$. The `em` and `reml` methods are specific to Cox models with gamma and gaussian random effects, respectively. Please see further discussion below.
- `tdf`: the degrees of freedom for the t-distribution.
- `eps`: convergence criteria for the iteration on `theta`.
Details

The frailty plugs into the general penalized modeling framework provided by the coxph and survreg routines. This framework deals with likelihood, penalties, and degrees of freedom; these aspects work well with either parent routine.

Therneau, Grambsch, and Pankratz show how maximum likelihood estimation for the Cox model with a gamma frailty can be accomplished using a general penalized routine, and Ripatti and Palmgren work through a similar argument for the Cox model with a gaussian frailty. Both of these are specific to the Cox model. Use of gamma/ml or gaussian/reml with survreg does not lead to valid results.

The extensible structure of the penalized methods is such that the penalty function, such as frailty or pspline, is completely separate from the modeling routine. The strength of this is that a user can plug in any penalization routine they choose. A weakness is that it is very difficult for the modeling routine to know whether a sensible penalty routine has been supplied.

Note that use of a frailty term implies a mixed effects model and use of a cluster term implies a GEE approach; these cannot be mixed.

The coxme package has superseded this method. It is faster, more stable, and more flexible.

Value

this function is used in the model statement of either coxph or survreg. It's results are used internally.

References


See Also

coxph, survreg

Examples

# Random institutional effect
coxph(Surv(time, status) ~ age + frailty(inst, df=4), lung)

# Litter effects for the rats data
rfit2a <- coxph(Surv(time, status) ~ rx +
frailty.gaussian(litter, df=13, sparse=FALSE), rats,
subset= (sex=='f'))
rfit2b <- coxph(Surv(time, status) ~ rx +
frailty.gaussian(litter, df=13, sparse=TRUE), rats,
subset= (sex=='f'))
Breast cancer data sets used in Royston and Altman (2013)

Description
The gbsg data set contains patient records from a 1984-1989 trial conducted by the German Breast Cancer Study Group (GBSG) of 720 patients with node positive breast cancer; it retains the 686 patients with complete data for the prognostic variables.

Usage

gbsg
data(cancer, package="survival")

Format
A data set with 686 observations and 11 variables.

pid  patient identifier
age  age, years
meno  menopausal status (0= premenopausal, 1= postmenopausal)
size  tumor size, mm
grade  tumor grade
nodes  number of positive lymph nodes
pgr  progesterone receptors (fmol/l)
er  estrogen receptors (fmol/l)
hormon  hormonal therapy, 0= no, 1= yes
rfstime  recurrence free survival time; days to first of recurrence, death or last follow-up
status  0= alive without recurrence, 1= recurrence or death

Details
These data sets are used in the paper by Royston and Altman. The Rotterdam data is used to create a fitted model, and the GBSG data for validation of the model. The paper gives references for the data source.

References

See Also
roterdam
**heart**  

**Stanford Heart Transplant data**

**Description**

Survival of patients on the waiting list for the Stanford heart transplant program.

**Usage**

```r
heart
data(heart, package="survival")
```

**Format**

- `jasa`: original data
  - `birth.dt`: birth date
  - `accept.dt`: acceptance into program
  - `tx.date`: transplant date
  - `fu.date`: end of followup
  - `fustat`: dead or alive
  - `surgery`: prior bypass surgery
  - `age`: age (in years)
  - `futime`: followup time
  - `wait.time`: time before transplant
  - `transplant`: transplant indicator
  - `mismatch`: mismatch score
  - `hla.a2`: particular type of mismatch
  - `mscore`: another mismatch score
  - `reject`: rejection occurred

- `jasa1`, `heart`: processed data
  - `start`, `stop`, `event`: Entry and exit time and status for this interval of time
  - `age`: age-48 years
  - `year`: year of acceptance (in years after 1 Nov 1967)
  - `surgery`: prior bypass surgery 1=yes
  - `transplant`: received transplant 1=yes
  - `id`: patient id

**Source**

is.ratetable

Verify that an object is of class ratetable.

Description

The function verifies not only the class attribute, but the structure of the object.

Usage

is.ratetable(x, verbose=FALSE)

Arguments

x the object to be verified.
verbose if TRUE and the object is not a ratetable, then return a character string describing the way(s) in which x fails to be a proper ratetable object.

Details

Rate tables are used by the pyears and survexp functions, and normally contain death rates for some population, categorized by age, sex, or other variables. They have a fairly rigid structure, and the verbose option can help in creating a new rate table.

Value

returns TRUE if x is a ratetable, and FALSE or a description if it is not.

See Also

pyears, survexp.

Examples

is.ratetable(survexp.us)  # True
is.ratetable(lung)       # False
Kidney catheter data

Description

Data on the recurrence times to infection, at the point of insertion of the catheter, for kidney patients using portable dialysis equipment. Catheters may be removed for reasons other than infection, in which case the observation is censored. Each patient has exactly 2 observations.

This data has often been used to illustrate the use of random effects (frailty) in a survival model. However, one of the males (id 21) is a large outlier, with much longer survival than his peers. If this observation is removed no evidence remains for a random subject effect.

Usage

```r
data(cancer, package="survival")
```

Format

- **patient**: id
- **time**: time
- **status**: event status
- **age**: in years
- **sex**: 1=male, 2=female
- **disease**: disease type (0=GN, 1=AN, 2=PKD, 3=Other)
- **frail**: frailty estimate from original paper

Note

The original paper ignored the issue of tied times and so is not exactly reproduced by the survival package.

Source


Examples

```r
kfit <- coxph(Surv(time, status) ~ age + sex + disease + frailty(id), kidney)
kfit0 <- coxph(Surv(time, status) ~ age + sex + disease, kidney)
kfitm1 <- coxph(Surv(time, status) ~ age + sex + disease + frailty(id, dist='gauss'), kidney)
```
levels.Surv  \hspace{1cm} \textit{Return the states of a multi-state Surv object}

\begin{description}
\item[Description] For a multi-state Surv object, this will return the names of the states.
\item[Usage] \begin{verbatim}
## S3 method for class 'Surv'
levels(x)
\end{verbatim}
\item[Arguments] \begin{verbatim}
x \hspace{1cm} \text{a Surv object}
\end{verbatim}
\item[Value] for a multi-state Surv object, the vector of state names (excluding censoring); or NULL for an ordinary Surv object
\item[Examples] \begin{verbatim}
y1 <- Surv(c(1,5, 9, 17,21, 30),
          factor(c(0, 1, 2,1,0,2), 0:2, c("censored", "progression", "death")))
levels(y1)
y2 <- Surv(1:6, rep(0:1, 3))
y2
levels(y2)
\end{verbatim}
\end{description}

\begin{description}
\item[lines.survfit] \hspace{1cm} \textit{Add Lines or Points to a Survival Plot}
\item[Description] Often used to add the expected survival curve(s) to a Kaplan-Meier plot generated with \texttt{plot.survfit}.
\item[Usage] \begin{verbatim}
## S3 method for class 'survfit'
lines(x, type="s", pch=3, col=1, lty=1,
      lwd=1, cex=1, mark.time=FALSE, xmax,
      fun, conf.int=FALSE,
      conf.times, conf.cap=.005, conf.offset=.012,
      conf.type = c("log", "log-log", "plain", "logit", "arcsin"),
\end{verbatim}
\end{description}
Arguments

x

a survival object, generated from the `survfit` or `survexp` functions.

type

the line type, as described in `lines`. The default is a step function for `survfit` objects, and a connected line for `survexp` objects. All other arguments for `lines.survexp` are identical to those for `lines.survfit`.

col, lty, lwd, cex

vectors giving the mark symbol, color, line type, line width and character size for the added curves. Of this set only color is applicable to `points`.

pch

plotting characters for points, in the style of `matplot`, i.e., either a single string of characters of which the first will be used for the first curve, etc; or a vector of characters or integers, one element per curve.

mark

a historical alias for `pch`

censor

should censoring times be displayed for the `points` function?

mark.time

controls the labeling of the curves. If FALSE, no labeling is done. If TRUE, then curves are marked at each censoring time. If `mark.time` is a numeric vector, then curves are marked at the specified time points.

xmax

optional cutoff for the right hand of the curves.

fun

an arbitrary function defining a transformation of the survival curve. For example `fun=log` is an alternative way to draw a log-survival curve (but with the axis labeled with log(S) values). Four often used transformations can be specified with a character argument instead: "log" is the same as using the `log=T` option, "event" plots cumulative events (f(y) = 1-y), "cumhaz" plots the cumulative hazard function (f(y) = -log(y)) and "cloglog" creates a complimentary log-log survival plot (f(y) = log(-log(y))) along with log scale for the x-axis.

conf.int

if TRUE, confidence bands for the curves are also plotted. If set to "only", then only the CI bands are plotted, and the curve itself is left off. This can be useful for fine control over the colors or line types of a plot. A numeric value, e.g. `conf.int = .90`, can be used to

conf.times

optional vector of times at which to place a confidence bar on the curve(s). If present, these will be used instead of confidence bands.

conf.cap

width of the horizontal cap on top of the confidence bars; only used if `conf.times` is used. A value of 1 is the width of the plot region.

conf.offset

the offset for confidence bars, when there are multiple curves on the plot. A value of 1 is the width of the plot region. If this is a single number then each curve’s bars are offset by this amount from the prior curve’s bars, if it is a vector the values are used directly.
conf.type
One of "plain", "log" (the default), "log-log", "logit", or "none". Only enough of the string to uniquely identify it is necessary. The first option causes confidence intervals not to be generated. The second causes the standard intervals curve +/- k * se(curve), where k is determined from conf.int. The log option calculates intervals based on the cumulative hazard or log(survival). The log-log option bases the intervals on the log hazard or log(-log(survival)), and the logit option on log(survival/(1-survival)).

noplot
for multi-state models, curves with this label will not be plotted. The default corresponds to an unspecified state.

cumhaz
plot the cumulative hazard, rather than the survival or probability in state.

... other graphical parameters

Details
When the survfit function creates a multi-state survival curve the resulting object has class 'survfits'. The only difference in the plots is that that it defaults to a curve that goes from lower left to upper right (starting at 0), where survival curves default to starting at 1 and going down. All other options are identical.

If the user set an explicit range in an earlier plot.survfit call, e.g. via xlim or xmax, subsequent calls to this function remember the right hand cutoff. This memory can be erased by options(plot.survfit) <- NULL.

Value
a list with components x and y, containing the coordinates of the last point on each of the curves (but not of the confidence limits). This may be useful for labeling.

Side Effects
one or more curves are added to the current plot.

See Also
lines, par, plot.survfit, survfit, survexp.

Examples
fit <- survfit(Surv(time, status==2) ~ sex, pbc, subset=1:312)
plot(fit, mark.time=FALSE, xscale=365.25, xlab='Years', ylab='Survival')
lines(fit[1], lwd=2)  # darken the first curve and add marks

# Add expected survival curves for the two groups,
# based on the US census data
# The data set does not have entry date, use the midpoint of the study
efit <- survexp(~sex, data=pbc, times= (0:24)*182, ratetable=survexp.us,
         rmap=list(sex=sex, age=age*365.35, year=as.Date('1979/01/01')))
temp <- lines(efit, lty=2, lwd=2:1)
Data from the 1972-78 GSS data used by Logan

Description

Intergenerational occupational mobility data with covariates.

Usage

```r
logan
data(logan, package="survival")
```

Format

A data frame with 838 observations on the following 4 variables.

- **occupation**: subject’s occupation, a factor with levels farm, operatives, craftsmen, sales, and professional
- **focc**: father’s occupation
- **education**: total years of schooling, 0 to 20
- **race**: levels of non-black and black

Source

General Social Survey data, see the web site for detailed information on the variables. [https://gss.norc.org/](https://gss.norc.org/).

References

logLik.coxph

logLik method for a Cox model

Description
The logLik function for survival models

Usage
## S3 method for class 'coxph'
logLik(object, ...)
## S3 method for class 'survreg'
logLik(object, ...)

Arguments
object the result of a coxph or survreg fit
... optional arguments for other instances of the method

Details
The logLik function is used by summary functions in R such as AIC. For a Cox model, this method returns the partial likelihood. The number of degrees of freedom (df) used by the fit and the effective number of observations (nobs) are added as attributes. Per Raftery and others, the effective number of observations is the taken to be the number of events in the data set.

For a survreg model the proper value for the effective number of observations is still an open question (at least to this author). For right censored data the approach of logLik.coxph is the possible the most sensible, but for interval censored observations the result is unclear. The code currently does not add a nobs attribute.

Value
an object of class logLik

Author(s)
Terry Therneau

References

See Also
logLik
**lung**  

*NCCTG Lung Cancer Data*

**Description**

Survival in patients with advanced lung cancer from the North Central Cancer Treatment Group. Performance scores rate how well the patient can perform usual daily activities.

**Usage**

lung
data(cancer, package="survival")

**Format**

- **inst:** Institution code  
- **time:** Survival time in days  
- **status:** censoring status 1=censored, 2=dead  
- **age:** Age in years  
- **sex:** Male=1 Female=2  
- **ph.ecog:** ECOG performance score as rated by the physician. 0=asymptomatic, 1= symptomatic but completely ambulatory, 2= in bed <50% of the day, 3= in bed > 50% of the day but not bedbound, 4 = bedbound  
- **ph.karno:** Karnofsky performance score (bad=0-good=100) rated by physician  
- **pat.karno:** Karnofsky performance score as rated by patient  
- **meal.cal:** Calories consumed at meals  
- **wt.loss:** Weight loss in last six months (pounds)

**Note**

The use of 1/2 for alive/dead instead of the usual 0/1 is a historical footnote. For data contained on punch cards, IBM 360 Fortran treated blank as a zero, which led to a policy within the section of Biostatistics to never use "0" as a data value since one could not distinguish it from a missing value. The policy became a habit, as is often the case; and the 1/2 coding endured long beyond the demise of punch cards and Fortran.

**Source**

Terry Therneau

**References**

Description

Natural history of 241 subjects with monoclonal gammopathy of undetermined significance (MGUS).

Usage

```r
mgus
mgus1
data(cancer, package="survival")
```

Format

`mgus`: A data frame with 241 observations on the following 12 variables.

- **id**: subject id
- **age**: age in years at the detection of MGUS
- **sex**: male or female
- **dxyr**: year of diagnosis
- **pcdx**: for subjects who progress to a plasma cell malignancy
  - the subtype of malignancy: multiple myeloma (MM) is the most common, followed by amyloidosis (AM), macroglobulinemia (MA), and other lymphoproliferative disorders (LP)
- **pctime**: days from MGUS until diagnosis of a plasma cell malignancy
- **futime**: days from diagnosis to last follow-up
- **death**: 1= follow-up is until death
- **alb**: albumin level at MGUS diagnosis
- **creat**: creatinine at MGUS diagnosis
- **hgb**: hemoglobin at MGUS diagnosis
- **mspike**: size of the monoclonal protein spike at diagnosis

`mgus1`: The same data set in start,stop format. Contains the id, age, sex, and laboratory variable described above along with

- **start, stop**: sequential intervals of time for each subject
- **status**: =1 if the interval ends in an event
- **event**: a factor containing the event type: censor, death, or plasma cell malignancy
- **enum**: event number for each subject: 1 or 2

Details

Plasma cells are responsible for manufacturing immunoglobulins, an important part of the immune defense. At any given time there are estimated to be about $10^6$ different immunoglobulins in the circulation at any one time. When a patient has a plasma cell malignancy the distribution will
become dominated by a single isotype, the product of the malignant clone, visible as a spike on a
serum protein electrophoresis. Monoclonal gammopathy of undetermined significance (MGUS)
is the presence of such a spike, but in a patient with no evidence of overt malignancy. This data
set of 241 sequential subjects at Mayo Clinic was the groundbreaking study defining the natural
history of such subjects. Due to the diligence of the principle investigator 0 subjects have been lost
to follow-up.

Three subjects had MGUS detected on the day of death. In data set mgus1 these subjects have the
time to MGUS coded as .5 day before the death in order to avoid tied times.

These data sets were updated in Jan 2015 to correct some small errors.

Source

Mayo Clinic data courtesy of Dr. Robert Kyle.

References

R Kyle, Benign monoclonal gammopathy – after 20 to 35 years of follow-up, Mayo Clinic Proc
1993; 68:26-36.

Examples

# Create the competing risk curves for time to first of death or PCM
sfit <- survfit(Surv(start, stop, event) ~ sex, mgus1, id=id,
subset=(enum==1))
print(sfit)  # the order of printout is the order in which they plot

plot(sfit, xscale=365.25, lty=c(2,2,1,1), col=c(1,2,1,2),
xlab="Years after MGUS detection", ylab="Proportion")
legend(0, .8, c("Death/male", "Death/female", "PCM/male", "PCM/female"),
     lty=c(1,1,2,2), col=c(2,1,2,1), bty="n")

title("Curves for the first of plasma cell malignancy or death")
# The plot shows that males have a higher death rate than females (no
# surprise) but their rates of conversion to PCM are essentially the same.
Format

A data frame with 1384 observations on the following 10 variables.

- id  subject identifier
- age  age at diagnosis, in years
- sex  a factor with levels F M
- dxyr  year of diagnosis
- hgb  hemoglobin
- creat  creatinine
- mspike  size of the monoclonal serum spike
- ptime  time until progression to a plasma cell malignancy (PCM) or last contact, in months
- pstat  occurrence of PCM: 0=no, 1=yes
- futime  time until death or last contact, in months
- death  occurrence of death: 0=no, 1=yes

Details

This is an extension of the study found in the mgus data set, containing enrollment through 1994 and follow-up through 1999.

Source

Mayo Clinic data courtesy of Dr. Robert Kyle. All patient identifiers have been removed, age rounded to the nearest year, and follow-up times rounded to the nearest month.

References


---

**model.frame.coxph**

*Model.frame method for coxph objects*

### Description

Recreate the model frame of a coxph fit.

### Usage

```r
## S3 method for class 'coxph'
model.frame(formula, ...)
```
Reconstruct the model matrix for a cox model.

Usage

```r
## S3 method for class 'coxph'
model.matrix(object, data=NULL, contrast.arg = object$contrasts, ...)
```

Arguments

- `object` the result of a coxph model
- `data` optional, a data frame from which to obtain the data
- `contrast.arg` optional, a contrasts object describing how factors should be coded
- `...` other possible argument to `model.frame`

Details

When there is a `data` argument this function differs from most of the other `model.matrix` methods in that the response variable for the original formula is *not* required to be in the data.

If the data frame contains a `terms` attribute then it is assumed to be the result of a call to `model.frame`, otherwise a call to `model.frame` is applied with the data as an argument.
Value
The model matrix for the fit

Author(s)
Terry Therneau

See Also
model.matrix

Examples
fit1 <- coxph(Surv(time, status) ~ age + factor(ph.ecog), data=lung)
xfit <- model.matrix(fit1)

fit2 <- coxph(Surv(time, status) ~ age + factor(ph.ecog), data=lung,
x=TRUE)
all.equal(model.matrix(fit1), fit2$x)

myeloid  Acute myeloid leukemia

Description
This simulated data set is based on a trial in acute myeloid leukemia.

Usage
myeloid
data(cancer, package="survival")

Format
A data frame with 646 observations on the following 9 variables.

id  subject identifier, 1-646
trt  treatment arm A or B
sex  f=female, m=male
futime  time to death or last follow-up
death  1 if futime is a death, 0 for censoring
txtime  time to hematropetic stem cell transplant
crt ime  time to complete response
r1time  time to relapse of disease
Details
This data set is used to illustrate multi-state survival curves. The correlation between within-subject event times strongly resembles that from an actual trial, but none of the actual data values are from that source.

References

Examples
coxph(Surv(futime, death) ~ trt, data=myeloid)
# See the mstate vignette for a more complete analysis

myeloma Survival times of patients with multiple myeloma

Description
Survival times of 3882 subjects with multiple myeloma, seen at Mayo Clinic from 1947–1996.

Usage
myeloma
data("cancer", package="survival")

Format
A data frame with 3882 observations on the following 5 variables.

id  subject identifier
year  year of entry into the study
entry  time from diagnosis of MM until entry (days)
futime  follow up time (days)
death  status at last follow-up: 0 = alive, 1 = death

Details
Subjects who were diagnosed at Mayo will have entry =0, those who were diagnosed elsewhere and later referred will have positive values.

References
Examples

# Incorrect survival curve, which ignores left truncation
fit1 <- survfit(Surv(futime, death) ~ 1, myeloma)
# Correct curve
fit2 <- survfit(Surv(entry, futime, death) ~1, myeloma)

nafld

Non-alcohol fatty liver disease

Description

Data sets containing the data from a population study of non-alcoholic fatty liver disease (NAFLD). Subjects with the condition and a set of matched control subjects were followed forward for metabolic conditions, cardiac endpoints, and death.

Usage

nafld1

nafld2

nafld3
data(nafld, package="survival")

Format

nafld1 is a data frame with 17549 observations on the following 10 variables.

id  subject identifier
age  age at entry to the study
male 0=female, 1=male
weight weight in kg
height height in cm
bmi  body mass index
case.id the id of the NAFLD case to whom this subject is matched
futime time to death or last follow-up
status 0= alive at last follow-up, 1=dead

nafld2 is a data frame with 400123 observations and 4 variables containing laboratory data

id  subject identifier
days  days since index date
test the type of value recorded
value the numeric value

nafld3 is a data frame with 34340 observations and 3 variables containing outcomes

id  subject identifier
days  days since index date
event the endpoint that occurred
Details

The primary reference for the NAFLD study is Allen (2018). The incidence of non-alcoholic fatty liver disease (NAFLD) has been rising rapidly in the last decade and it is now one of the main drivers of hepatology practice Tapper2018. It is essentially the presence of excess fat in the liver, and parallels the ongoing obesity epidemic. Approximately 20-25% of NAFLD patients will develop the inflammatory state of non-alcoholic steatohepatitis (NASH), leading to fibrosis and eventual end-stage liver disease. NAFLD can be accurately diagnosed by MRI methods, but NASH diagnosis currently requires a biopsy.

The current study constructed a population cohort of all adult NAFLD subjects from 1997 to 2014 along with 4 potential controls for each case. To protect patient confidentiality all time intervals are in days since the index date; none of the dates from the original data were retained. Subject age is their integer age at the index date, and the subject identifier is an arbitrary integer. As a final protection, we include only a 90% random sample of the data. As a consequence analyses results will not exactly match the original paper.

There are 3 data sets: nafl1d1 contains baseline data and has one observation per subject, nafl1d2 has one observation for each (time dependent) continuous measurement, and nafl1d3 has one observation for each yes/no outcome that occurred.

Source

Data obtained from the author.

References


Description

A common task in medical work is to find the closest lab value to some index date, for each subject.

Usage

neardate(id1, id2, y1, y2, best = c("after", "prior"), nomatch = NA_integer_)

Arguments

id1 vector of subject identifiers for the index group
id2 vector of identifiers for the reference group
y1 normally a vector of dates for the index group, but any orderable data type is allowed
y2 reference set of dates
best if best='prior' find the index of the first y2 value less than or equal to the target y1 value, for each subject. If best='after' find the first y2 value which is greater than or equal to the target y1 value, for each subject.
nomatch the value to return for items without a match

Details
This routine is closely related to match and to findInterval, the first of which finds exact matches and the second closest matches. This finds the closest matching date within sets of exactly matching identifiers. Closest date matching is often needed in clinical studies. For example data set 1 might contain the subject identifier and the date of some procedure and data set 2 has the dates and values for laboratory tests, and the query is to find the first test value after the intervention but no closer than 7 days.

The id1 and id2 arguments are similar to match in that we are searching for instances of id1 that will be found in id2, and the result is the same length as id1. However, instead of returning the first match with id2 this routine returns the one that best matches with respect to y1.

The y1 and y2 arguments need not be dates, the function works for any data type such that the expression c(y1,y2) gives a sensible, sortable result. Be careful about matching Date and DateTime values and the impact of time zones, however, see as.POSIXct. If y1 and y2 are not of the same class the user is on their own. Since there exist pairs of unmatched data types where the result could be sensible, the routine will in this case proceed under the assumption that "the user knows what they are doing". Caveat emptor.

Value the index of the matching observations in the second data set, or the nomatch value for no successful match

Author(s)
Terry Therneau

See Also
match, findInterval

Examples
data1 <- data.frame(id = 1:10,
                     entry.dt = as.Date(paste("2011", 1:10, "5", sep='-')))
temp1 <- c(1,4,5,1,3,6,9, 2,7,8,12,4,6,7,10,12,3)
data2 <- data.frame(id = c(1,1,1,2,2,4,4,5,5,6,8,9,10,10,12),
                     lab.dt = as.Date(paste("2011", temp1, "1", sep='-')),
                     chol = round(runif(17, 130, 280)))

#first cholesterol on or after enrollment
indx1 <- neardate(data1$id, data2$id, data1$entry.dt, data2$lab.dt)
data2[indx1, "chol"]

# Closest one, either before or after.
#
indx2 <- neardate(data1$id, data2$id, data1$entry.dt, data2$lab.dt,
                   best="prior")
ifelse(is.na(indx1), indx2, # none after, take before
       ifelse(is.na(indx2), indx1, #none before
                ifelse(abs(data2$lab.dt[indx2] - data1$entry.dt) <
                        abs(data2$lab.dt[indx1] - data1$entry.dt), indx2, indx1)))

# closest date before or after, but no more than 21 days prior to index
indx2 <- ifelse((data1$entry.dt - data2$lab.dt[indx2]) >21, NA, indx2)
ifelse(is.na(indx1), indx2, # none after, take before
       ifelse(is.na(indx2), indx1, #none before
                ifelse(abs(data2$lab.dt[indx2] - data1$entry.dt) <
                        abs(data2$lab.dt[indx1] - data1$entry.dt), indx2, indx1)))

**Description**

Natural splines with knot heights as the basis.

Create the design matrix for a natural spline, such that the coefficient of the resulting fit are the values of the function at the knots.

**Usage**

```r
nsk(x, df = NULL, knots = NULL, intercept = FALSE, b = 0.05,
    Boundary.knots = quantile(x, c(b, 1 - b), na.rm = TRUE))
```

**Arguments**

- `x`: the predictor variable. Missing values are allowed.
- `df`: degrees of freedom. One can supply df rather than knots; ns() then chooses df - 1 - intercept knots at suitably chosen quantiles of x (which will ignore missing values). The default, df = NULL, sets the number of inner knots as length(knots).
- `knots`: breakpoints that define the spline. The default is no knots; together with the natural boundary conditions this results in a basis for linear regression on x. Typical values are the mean or median for one knot, quantiles for more knots. See also Boundary.knots.
- `intercept`: if TRUE, an intercept is included in the basis; default is FALSE
- `b`: default placement of the boundary knots. A value of bs=0 will replicate the default behavior of ns.
- `Boundary.knots`: boundary points at which to impose the natural boundary conditions and anchor the B-spline basis. Beyond these points the function is assumed to be linear. If both knots and Boundary.knots are supplied, the basis parameters do not depend on x. Data can extend beyond Boundary.knots.
Details

The nsk function behaves identically to the ns function, with two exceptions. The primary one is that the returned basis is such that coefficients correspond to the value of the fitted function at the knot points. If intercept = FALSE, there will be k-1 coefficients corresponding to the k knots, and they will be the difference in predicted value between knots 2-k and knot 1. The primary advantage to the basis is that the coefficients are directly interpretable. A second is that tests for the linear and non-linear components are simple contrasts.

The second difference with ns is one of opinion with respect to the default position for the boundary knots. The default here is closer to that found in the rms::rcs function.

This function is a trial if a new idea, it’s future inclusion in the package is not yet guaranteed.

Value

A matrix of dimension length(x) * df where either df was supplied or, if knots were supplied, df = length(knots) + 1 + intercept. Attributes are returned that correspond to the arguments to kns, and explicitly give the knots, Boundary.knots etc for use by predict.kns().

Note

A thin flexible metal or wooden strip is called a spline, and is the traditional method for laying out a smooth curve, e.g., for a ship’s hull or an airplane wing. Pins are put into a board and the strip is passed through them, each pin is a ‘knot’.

A mathematical spline is a piecewise function between each knot. A linear spline will be a set of connected line segments, a quadratic spline is a set of connected local quadratic functions, constrained to have a continuous first derivative, a cubic spline is cubic between each knot, constrained to have continuous first and second derivatives, and etc. Mathematical splines are not an exact representation of natural splines: being a physical object the wood or metal strip will have continuous derivatives of all orders. Cubic splines are commonly used because they are sufficiently smooth to look natural to the human eye.

If the mathematical spline is further constrained to be linear beyond the end knots, this is often called a ‘natural spline’, due to the fact that a wooden or metal spline will also be linear beyond the last knots. Another name for the same object is a ‘restricted cubic spline’, since it is achieved in code by adding further constraints. Given a vector of data points and a set of knots, it is possible to create a basis matrix X with one column per knot, such that ordinary regression of X on y will fit the cubic spline function, hence these are also called ‘regression splines’. (One of these three labels is no better or worse than another, in our opinion).

Given a basis matrix X with k columns, the matrix Z= XT for any k by k nonsingular matrix T is is also a basis matrix, and will result in identical predicted values, but a new set of coefficients gamma = (T-inverse) beta in place of beta. One can choose the basis functions so that X is easy to construct, to make the regression numerically stable, to make tests easier, or based on other considerations. It seems as though every spline library returns a different basis set, which unfortunately makes fits difficult to compare between packages. This is yet one more basis set, chosen to make the coefficients more interpretable.

See Also

ns
Examples

# make some dummy data
tdata <- data.frame(x= lung$age, y = 10*log(lung$age-35) + rnorm(228, 0, 2))
fit1 <- lm(y ~ -1 + nsk(x, df=4, intercept=TRUE) , data=tdata)
fit2 <- lm(y ~ nsk(x, df=3), data=tdata)

# the knots (same for both fits)
knots <- unlist(attributes(fit1$model)[[2]])[c('Boundary.knots', 'knots')]
sort(unname(knots))
unname(coef(fit1)) # predictions at the knot points
unname(coef(fit1)[-1] - coef(fit1)[1]) # differences: yhat[2:4] - yhat[1]
unname(coef(fit2)[-1]) # ditto

## Not run:
plot(y ~ x, data=tdata)
points(sort(knots), coef(fit1), col=2, pch=19)
print(coef(fit)[1] + c(0, coef(fit)[-1]))
## End(Not run)

nwtco

Data from the National Wilm’s Tumor Study

Description

Measurement error example. Tumor histology predicts survival, but prediction is stronger with central lab histology than with the local institution determination.

Usage

nwtco
data(nwtco, package="survival")

Format

A data frame with 4028 observations on the following 9 variables.

seqno  id number
instit  Histology from local institution
histol  Histology from central lab
stage  Disease stage
study  study
rel  indicator for relapse
edrel  time to relapse
age  age in months
in.subcohort  Included in the subcohort for the example in the paper
References


Examples

```r
with(nwtco, table(instit,histol))
anova(coxph(Surv(edrel,rel)~histol+instit,data=nwtco))
anova(coxph(Surv(edrel,rel)~instit+histol,data=nwtco))
```

---

**ovarian**  
*Ovarian Cancer Survival Data*

Description

Survival in a randomised trial comparing two treatments for ovarian cancer

Usage

```r
ovarian
data(cancer, package="survival")
```

Format

- `futime`: survival or censoring time  
- `fustat`: censoring status  
- `age`: in years  
- `resid.ds`: residual disease present (1=no, 2=yes)  
- `rx`: treatment group  
- `ecog.ps`: ECOG performance status (1 is better, see reference)

Source

Terry Therneau

References

Description

Primary sclerosing cholangitis is an autoimmune disease leading to destruction of the small bile ducts in the liver. Progression is slow but inexorable, eventually leading to cirrhosis and liver decompensation. The condition has been recognised since at least 1851 and was named "primary biliary cirrhosis" in 1949. Because cirrhosis is a feature only of advanced disease, a change of its name to "primary biliary cholangitis" was proposed by patient advocacy groups in 2014.

This data is from the Mayo Clinic trial in PBC conducted between 1974 and 1984. A total of 424 PBC patients, referred to Mayo Clinic during that ten-year interval, met eligibility criteria for the randomized placebo controlled trial of the drug D-penicillamine. The first 312 cases in the data set participated in the randomized trial and contain largely complete data. The additional 112 cases did not participate in the clinical trial, but consented to have basic measurements recorded and to be followed for survival. Six of those cases were lost to follow-up shortly after diagnosis, so the data here are on an additional 106 cases as well as the 312 randomized participants.

A nearly identical data set found in appendix D of Fleming and Harrington; this version has fewer missing values.

Usage

\[
\text{pbc} \\
\text{data(pbc, package="survival")}
\]

Format

- **age**: in years
- **albumin**: serum albumin (g/dl)
- **alk.phos**: alkaline phosphotase (U/liter)
- **ascites**: presence of ascites
- **ast**: aspartate aminotransferase, once called SGOT (U/ml)
- **bili**: serum bilirunbin (mg/dl)
- **chol**: serum cholesterol (mg/dl)
- **copper**: urine copper (ug/day)
- **edema**: 0 no edema, 0.5 untreated or successfully treated 
  1 edema despite diuretic therapy
- **hepato**: presence of hepatomegaly or enlarged liver
- **id**: case number
- **platelet**: platelet count
- **protime**: standardised blood clotting time
- **sex**: m/f
- **spiders**: blood vessel malformations in the skin
- **stage**: histologic stage of disease (needs biopsy)
- **status**: status at endpoint, 0/1/2 for censored, transplant, dead
- **time**: number of days between registration and the earlier of death,
transplantion, or study analysis in July, 1986

trt: 1/2/NA for D-penicillmain, placebo, not randomised

trig: triglycerides (mg/dl)

Source


See Also

pbcseq

---

**pbcseq**  
*Mayo Clinic Primary Biliary Cirrhosis, sequential data*

Description

This data is a continuation of the PBC data set, and contains the follow-up laboratory data for each study patient. An analysis based on the data can be found in Murtagh, et. al.

The primary PBC data set contains only baseline measurements of the laboratory parameters. This data set contains multiple laboratory results, but only on the 312 randomized patients. Some baseline data values in this file differ from the original PBC file, for instance, the data errors in prothrombin time and age which were discovered after the original analysis (see Fleming and Harrington, figure 4.6.7).

One "feature" of the data deserves special comment. The last observation before death or liver transplant often has many more missing covariates than other data rows. The original clinical protocol for these patients specified visits at 6 months, 1 year, and annually thereafter. At these protocol visits lab values were obtained for a large pre-specified battery of tests. "Extra" visits, often undertaken because of worsening medical condition, did not necessarily have all this lab work. The missing values are thus potentially informative.

Usage

pbcseq
data(pbc, package="survival")

Format

<table>
<thead>
<tr>
<th>id</th>
<th>case number</th>
</tr>
</thead>
<tbody>
<tr>
<td>age</td>
<td>in years</td>
</tr>
<tr>
<td>sex</td>
<td>m/f</td>
</tr>
<tr>
<td>trt</td>
<td>1/2/NA for D-penicillmain, placebo, not randomised</td>
</tr>
</tbody>
</table>
time: number of days between registration and the earlier of death, transplantation, or study analysis in July, 1986
status: status at endpoint, 0/1/2 for censored, transplant, dead
day: number of days between enrollment and this visit date
all measurements below refer to this date
albumin: serum albumin (mg/dl)
alk.phos: alkaline phosphotase (U/liter)
ascites: presence of ascites
ast: aspartate aminotransferase, once called SGOT (U/ml)
bili: serum bilirubin (mg/dl)
chol: serum cholesterol (mg/dl)
copper: urine copper (ug/day)
edema: 0 no edema, 0.5 untreated or successfully treated
1 edema despite diuretic therapy
hepato: presence of hepatomegaly or enlarged liver
platelet: platelet count
protime: standardised blood clotting time
spiders: blood vessel malformations in the skin
stage: histologic stage of disease (needs biopsy)
trig: triglycerides (mg/dl)

Source


 References


See Also

pbc

Examples

# Create the start-stop-event triplet needed for coxph
first <- with(pbcseq, c(TRUE, diff(id) !=0)) #first id for each subject
last <- c(first[-1], TRUE) #last id
time1 <- with(pbcseq, ifelse(first, 0, day))
time2 <- with(pbcseq, ifelse(last, futime, c(day[-1], 0)))
event <- with(pbcseq, ifelse(last, status, 0))
fit1 <- coxph(Surv(time1, time2, event) ~ age + sex + log(bili), pbcseq)

### S3 method for class 'aareg'
plot(x, se=TRUE, maxtime, type='s', ...)
Graphical Test of Proportional Hazards

Description

Displays a graph of the scaled Schoenfeld residuals, along with a smooth curve.

Usage

```r
## S3 method for class 'cox.zph'
plot(x, resid=TRUE, se=TRUE, df=4, nsmo=40, var,
     xlab="Time", ylab, lty=1:2, col=1, lwd=1, hr=FALSE, ...)
```

Arguments

- `x` result of the `cox.zph` function.
- `resid` a logical value, if TRUE the residuals are included on the plot, as well as the smooth fit.
- `se` a logical value, if TRUE, confidence bands at two standard errors will be added.
- `df` the degrees of freedom for the fitted natural spline, df=2 leads to a linear fit.
- `nsmo` number of points to use for the lines
- `var` the set of variables for which plots are desired. By default, plots are produced in turn for each variable of a model. Selection of a single variable allows other features to be added to the plot, e.g., a horizontal line at zero or a main title. This has been superseded by a subcripting method; see the example below.
- `hr` if TRUE, label the y-axis using the estimated hazard ratio rather than the estimated coefficient. (The plot does not change, only the axis label.)
- `xlab` label for the x-axis of the plot
- `ylab` optional label for the y-axis of the plot. If missing a default label is provided. This can be a vector of labels.
- `lty, col, lwd` line type, color, and line width for the overlaid curve. Each of these can be vector of length 2, in which case the second element is used for the confidence interval.
- `...` additional graphical arguments passed to the `plot` function.

Side Effects

a plot is produced on the current graphics device.

See Also

coxph, cox.zph.
Examples

```r
vfit <- coxph(Surv(time,status) ~ trt + factor(celltype) +
               karno + age, data=veteran, x=TRUE)
temp <- cox.zph(vfit)
plot(temp, var=3) # Look at Karnofsy score, old way of doing plot
plot(temp[3]) # New way with subscripting
abline(0, 0, lty=3)
# Add the linear fit as well
abline(lm(temp$y[3] ~ temp$x)$coefficients, lty=4, col=3)
title(main="VA Lung Study")
```

plot.survfit

Plot method for survfit objects

Description

A plot of survival curves is produced, one curve for each strata. The log=T option does extra work to avoid log(0), and to try to create a pleasing result. If there are zeros, they are plotted by default at 0.8 times the smallest non-zero value on the curve(s).

Curves are plotted in the same order as they are listed by print (which gives a 1 line summary of each). This will be the order in which col, lty, etc are used.

Usage

```r
## S3 method for class 'survfit'
plot(x, conf.int=, mark.time=FALSE,
pch=3, col=1, lty=1, lwd=1, cex=1, log=FALSE, xscale=1, yscale=1,
xlim, ylim, xmax, fun,
lab="", ylab="", xaxs="r", conf.times, conf.cap=.005,
conf.offset=.012,
conf.type = c("log", "log-log", "plain", "logit", "arcsin"),
mark, noplot="(s0)", cumhaz=FALSE,
firstx, ymin, ...)```

Arguments

- `x` an object of class survfit, usually returned by the survfit function.
- `conf.int` determines whether pointwise confidence intervals will be plotted. The default is to do so if there is only 1 curve, i.e., no strata, using 95% confidence intervals. Alternatively, this can be a numeric value giving the desired confidence level.
- `mark.time` controls the labeling of the curves. If set to FALSE, no labeling is done. If TRUE, then curves are marked at each censoring time. If mark is a numeric vector then curves are marked at the specified time points.
- `pch` vector of characters which will be used to label the curves. The points help file contains examples of the possible marks. A single string such as "abcd" is treated as a vector c("a","b","c","d"). The vector is reused cyclically if it
is shorter than the number of curves. If it is present this implies mark.time = TRUE.

- `col` is a vector of integers specifying colors for each curve. The default value is 1.
- `lty` is a vector of integers specifying line types for each curve. The default value is 1.
- `lwd` is a vector of numeric values for line widths. The default value is 1.
- `cex` is a numeric value specifying the size of the marks. This is not treated as a vector; all marks have the same size.
- `log` is a logical value, if TRUE the y axis will be on a log scale. Alternately, one of the standard character strings "x", "y", or "xy" can be given to specify logarithmic horizontal and/or vertical axes.

- `xlim, ylim` are optional limits for the plotting region.
- `xmax` is the maximum horizontal plot coordinate. This can be used to shrink the range of a plot. It shortens the curve before plotting it, so that unlike using the `xlim` graphical parameter, warning messages about out of bounds points are not generated.

- `fun` is an arbitrary function defining a transformation of the survival (or probability in state, or cumulative hazard) curves. For example `fun=log` is an alternative way to draw a log-survival curve (but with the axis labeled with log(S) values), and `fun=sqrt` would generate a curve on square root scale. Four often used transformations can be specified with a character argument instead: "S" gives the usual survival curve, "log" is the same as using the log=T option, "event" or "F" plots the empirical CDF \( F(t) = 1 - S(t) \) \( (f(y) = 1 - y) \), and "cloglog" creates a complimentary log-log survival plot \( f(y) = \log(-\log(y)) \) along with log scale for the x-axis). The terms "identity" and "surv" are allowed as synonyms for type="S". The argument "cumhaz" causes the cumulative hazard function to be plotted.

- `xlab` is label given to the x-axis.
- `ylab` is label given to the y-axis.
- `xaxs` is either "S" for a survival curve or a standard x axis style as listed in par; "r" (regular) is the R default. Survival curves have historically been displayed with the curve touching the y-axis, but not touching the bounding box of the plot on the other 3 sides. Type "S" accomplishes this by manipulating the plot range and then using the "i" style internally. The "S" style is becoming increasingly less common, however.

- `conf.times` is optional vector of times at which to place a confidence bar on the curve(s). If present, these will be used instead of confidence bands.

- `conf.cap` is the width of the horizontal cap on top of the confidence bars; only used if conf.times is used. A value of 1 is the width of the plot region.
conf.offset the offset for confidence bars, when there are multiple curves on the plot. A value of 1 is the width of the plot region. If this is a single number then each curve’s bars are offset by this amount from the prior curve’s bars, if it is a vector the values are used directly.

conf.type One of "plain", "log" (the default), "log-log" or "logit". Only enough of the string to uniquely identify it is necessary. The first option causes confidence intervals not to be generated. The second causes the standard intervals curve \( +k \times \text{se(curve)} \), where \( k \) is determined from \text{conf.int}. The log option calculates intervals based on the cumulative hazard or log(survival). The log-log option bases the intervals on the log hazard or log(-log(survival)), and the logit option on log(survival/(1-survival)).

mark a historical alias for pch

noplot for multi-state models, curves with this label will not be plotted. (Also see the istate0 argument in survcheck.)

cumhaz plot the cumulative hazard rather than the probability in state or survival. Optionally, this can be a numeric vector specifying which columns of the cumhaz component to plot.

ymin this will normally be given as part of the ylim argument

firstx this will normally be given as part of the xlim argument.

... other arguments that will be passed forward to the underlying plot method, such as xlab or ylab.

Details

If the object contains a cumulative hazard curve, then \( \text{fun} = \text{`cumhaz'} \) will plot that curve, otherwise it will plot \(-\log(S)\) as an approximation. Theoretically, \( S = \exp(-\Lambda) \) where \( S \) is the survival and \( \Lambda \) is the cumulative hazard. The same relationship holds for estimates of \( S \) and \( \Lambda \) only in special cases, but the approximation is often close.

When the survfit function creates a multi-state survival curve the resulting object also has class ‘survfitsms’. Competing risk curves are a common case. In this situation the \text{fun} argument is ignored.

When the conf.times argument is used, the confidence bars are offset by \text{conf.offset} units to avoid overlap. The bar on each curve are the confidence interval for the time point at which the bar is drawn, i.e., different time points for each curve. If curves are steep at that point, the visual impact can sometimes substantially differ for positive and negative values of \text{conf.offset}.

Value

a list with components x and y, containing the coordinates of the last point on each of the curves (but not the confidence limits). This may be useful for labeling.

Note

In prior versions the behavior of xscale and yscale differed: the first changed the scale both for the plot and for all subsequent actions such as adding a legend, whereas yscale affected only the axis label. This was normalized in version 2-36.4, and both parameters now only affect the labeling.
In versions prior to approximately 2.36 a \texttt{survfit} object did not contain the cumulative hazard as a separate result, and the use of \texttt{fun="cumhaz"} would plot the approximation \(-\log(\text{surv})\) to the cumulative hazard. When cumulative hazards were added to the object, the \texttt{cumhaz=TRUE} argument to the plotting function was added. In version 2.3-8 the use of \texttt{fun="cumhaz"} became a synonym for \texttt{cumhaz=TRUE}.

\textbf{See Also}

\texttt{points.survfit}, \texttt{lines.survfit}, \texttt{par}, \texttt{survfit}

\textbf{Examples}

```r
leukemia.surv <- survfit(Surv(time, status) ~ x, data = aml)
plot(leukemia.surv, lty = 2:3)
legend(100, .9, c("Maintenance", "No Maintenance"), lty = 2:3)
title("Kaplan-Meier Curves for AML Maintenance Study")
lsurv2 <- survfit(Surv(time, status) ~ x, aml, type = "fleming")
plot(lsurv2, lty = 2:3, fun = "cumhaz",
     xlab = "Months", ylab = "Cumulative Hazard")
```

\textbf{predict.coxph} \hfill \textit{Predictions for a Cox model}

\textbf{Description}

Compute fitted values and regression terms for a model fitted by \texttt{coxph}

\textbf{Usage}

```r
## S3 method for class 'coxph'
predict(object, newdata,
         type = c("lp", "risk", "expected", "terms", "survival"),
         se.fit = FALSE, na.action = na.pass, terms = names(object$assign), collapse,
         reference = c("strata", "sample", "zero"), ...)  
```

\textbf{Arguments}

- \texttt{object} the results of a coxph fit.
- \texttt{newdata} Optional new data at which to do predictions. If absent predictions are for the data frame used in the original fit. When coxph has been called with a formula argument created in another context, i.e., coxph has been called within another function and the formula was passed as an argument to that function, there can be problems finding the data set. See the note below.
- \texttt{type} the type of predicted value. Choices are the linear predictor ("lp"), the risk score \(\exp(lp)\) ("risk"), the expected number of events given the covariates and follow-up time ("expected"), and the terms of the linear predictor ("terms"). The survival probability for a subject is equal to \(\exp(-\text{expected})\).
predict.coxph

se.fit if TRUE, pointwise standard errors are produced for the predictions.

na.action applies only when the newdata argument is present, and defines the missing value action for the new data. The default is to include all observations. When there is no newdata, then the behavior of missing is dictated by the na.action option of the original fit.

terms if type="terms", this argument can be used to specify which terms should be included; the default is all.

collapse optional vector of subject identifiers. If specified, the output will contain one entry per subject rather than one entry per observation.

reference reference for centering predictions, see details below

... For future methods

Details

The Cox model is a relative risk model; predictions of type "linear predictor", "risk", and "terms" are all relative to the sample from which they came. By default, the reference value for each of these is the mean covariate within strata. The underlying reason is both statistical and practical. First, a Cox model only predicts relative risks between pairs of subjects within the same strata, and hence the addition of a constant to any covariate, either overall or only within a particular stratum, has no effect on the fitted results. Second, downstream calculations depend on the risk score exp(linear predictor), which will fall prey to numeric overflow for a linear predictor greater than .Machine\$double.max.exp. The coxph routines try to approximately center the predictors out of self protection. Using the reference="strata" option is the safest centering, since strata occasionally have different means. When the results of predict are used in further calculations it may be desirable to use a single reference level for all observations. Use of reference="sample" will use the overall means, and agrees with the linear.predictors component of the coxph object (which uses the overall mean for backwards compatability with older code). Predictions of type="terms" are almost invariably passed forward to further calculation, so for these we default to using the sample as the reference. A reference of "zero" causes no centering to be done.

Predictions of type "expected" incorporate the baseline hazard and are thus absolute instead of relative; the reference option has no effect on these. These values depend on the follow-up time for the future subjects as well as covariates so the newdata argument needs to include both the right and left hand side variables from the formula. (The status variable will not be used, but is required since the underlying code needs to reconstruct the entire formula.)

Models that contain a frailty term are a special case: due to the technical difficulty, when there is a newdata argument the predictions will always be for a random effect of zero.

Value

a vector or matrix of predictions, or a list containing the predictions (element "fit") and their standard errors (element "se.fit") if the se.fit option is TRUE.

Note

Some predictions can be obtained directly from the coxph object, and for others it is necessary for the routine to have the entirety of the original data set, e.g., for type = terms or if standard errors are requested. This extra information is saved in the coxph object if model=TRUE, if not the original
data is reconstructed. If it is known that such residuals will be required overall execution will be slightly faster if the model information is saved.

In some cases the reconstruction can fail. The most common is when coxph has been called inside another function and the formula was passed as one of the arguments to that enclosing function. Another is when the data set has changed between the original call and the time of the prediction call. In each of these the simple solution is to add model=TRUE to the original coxph call.

See Also

predict, coxph, termplot

Examples

options(na.action=na.exclude) # retain NA in predictions
fit <- coxph(Surv(time, status) ~ age + ph.ecog + strata(inst), lung)
#lung data set has status coded as 1/2
mresid <- (lung$status-1) - predict(fit, type='expected') #Martingale resid
predict(fit, type='lp')
predict(fit, type='expected')
predict(fit, type='risk', se.fit=TRUE)
predict(fit, type='terms', se.fit=TRUE)

# For someone who demands reference='zero'
pzero <- function(fit)
  predict(fit, reference="sample") + sum(coef(fit) * fit$means, na.rm=TRUE)

predict.survreg

Predicted Values for a 'survreg' Object

Description

Predicted values for a survreg object

Usage

## S3 method for class 'survreg'
predict(object, newdata,
  type=c("response", "link", "lp", "linear", "terms", "quantile", "uquantile"),
  se.fit=FALSE, terms=NULL, p=c(0.1, 0.9), na.action=na.pass, ...)

Arguments

object result of a model fit using the survreg function.
newdata data for prediction. If absent predictions are for the subjects used in the original fit.
predict.survreg

The predict function for survreg returns the type of predicted value. This can be on the original scale of the data (response), the linear predictor ("linear", with "lp" as an allowed abbreviation), a predicted quantile on the original scale of the data ("quantile"), a quantile on the linear predictor scale ("uquantile"), or the matrix of terms for the linear predictor ("terms"). At this time "link" and linear predictor ("lp") are identical.

se.fit

If TRUE, include the standard errors of the prediction in the result.

terms

Subset of terms. The default for residual type "terms" is a matrix with one column for every term (excluding the intercept) in the model.

p

Vector of percentiles. This is used only for quantile predictions.

na.action

Applies only when the newdata argument is present, and defines the missing value action for the new data. The default is to include all observations.

... for future methods

Value

A vector or matrix of predicted values.

References


See Also

survreg, residuals.survreg

Examples

# Draw figure 1 from Escobar and Meeker, 1992.
fit <- survreg(Surv(time,status) ~ age + I(age^2), data=stanford2, dist="lognormal")
with(stanford2, plot(age, time, xlab='Age', ylab='Days', ylim=c(0,65), ylim=c(.1, 10^5), log='y', type='n'))
with(stanford2, points(age, time, pch=c(2,4)[status+1], cex=.7))
pred <- predict(fit, newdata=list(age=1:65), type="quantile", p=c(.1, .5, .9))
matlines(1:65, pred, lty=c(2,1,2), col=1)

# Predicted Weibull survival curve for a lung cancer subject with ECOG score of 2
lfit <- survreg(Surv(time, status) ~ ph.ecog, data=lung)
pct <- 1:98/100 # The 100th percentile of predicted survival is at +infinity
ptime <- predict(lfit, newdata=data.frame(ph.ecog=2), type="quantile", p=pct, se=TRUE)
matplot(cbind(ptime$fit, ptime$fit + 2*ptime$se.fit, ptime$fit - 2*ptime$se.fit)/30.5, 1-pct, xlab="Months", ylab="Survival", type='l', lty=c(1,2,2), col=1)
Print an aareg object

Description

Print out a fit of Aalen's additive regression model

Usage

## S3 method for class 'aareg'
print(x, maxtime, test=c("aalen", "nrisk"),scale=1,...)

Arguments

- **x**: the result of a call to the aareg function
- **maxtime**: the upper time point to be used in the test for non-zero slope
- **test**: the weighting to be used in the test for non-zero slope. The default weights are based on the variance of each coefficient, as a function of time. The alternative weight is proportional to the number of subjects still at risk at each time point.
- **scale**: scales the coefficients. For some data sets, the coefficients of the Aalen model will be very small (10^-4); this simply multiplies the printed values by a constant, say 1e6, to make the printout easier to read.
- **...**: for future methods

Details

The estimated increments in the coefficient estimates can become quite unstable near the end of follow-up, due to the small number of observations still at risk in a data set. Thus, the test for slope will sometimes be more powerful if this last 'tail' is excluded.

Value

the calling argument is returned.

Side Effects

the results of the fit are displayed.

References


See Also

aareg
print.summary.coxph  
*Print method for summary.coxph objects*

**Description**

Produces a printed summary of a fitted coxph model

**Usage**

```r
## S3 method for class 'summary.coxph'
print(x, digits=max(getOption("digits") - 3, 3),
    signif.stars = getOption("show.signif.stars"), expand=FALSE, ...)
```

**Arguments**

- `x`: the result of a call to `summary.coxph`
- `digits`: significant digits to print
- `signif.stars`: Show stars to highlight small p-values
- `expand`: if the summary is for a multi-state coxph fit, print the results in an expanded format.
- `...`: For future methods

---

print.summary.survexp  
*Print Survexp Summary*

**Description**

Prints the results of `summary.survexp`

**Usage**

```r
## S3 method for class 'summary.survexp'
print(x, digits = max(options()$digits - 4, 3), ...)
```

**Arguments**

- `x`: an object of class `summary.survexp`
- `digits`: the number of digits to use in printing the result.
- `...`: for future methods

**Value**

- `x`, with the invisible flag set to prevent further printing.
Description

Prints the result of `summary.survfit`.

Usage

```r
## S3 method for class 'summary.survfit'
print(x, digits = max(options() $digits-4, 3), ...)
```

Arguments

- `x`: an object of class "summary.survfit", which is the result of the `summary.survfit` function.
- `digits`: the number of digits to use in printing the numbers.
- `...`: for future methods

Value

`x`, with the invisible flag set to prevent printing.

Side Effects

prints the summary created by `summary.survfit`.

See Also

`options`, `print`, `summary.survfit`. 

Author(s)

Terry Therneau

See Also

`link{summary.survexp}`, `survexp`
print.survfit

Print a Short Summary of a Survival Curve

Description
Print number of observations, number of events, the restricted mean survival and its standard error, and the median survival with confidence limits for the median.

Usage
## S3 method for class 'survfit'
print(x, scale=1, digits = max(options()$digits - 4,3),
      print.rmean=getOption("survfit.print.rmean"),
      rmean = getOption('survfit.rmean'),...)

Arguments
x the result of a call to the survfit function.
scale a numeric value to rescale the survival time, e.g., if the input data to survfit were in days, scale=365 would scale the printout to years.
digits Number of digits to print
print.rmean,rmean Options for computation and display of the restricted mean.
... for future results

Details
The mean and its variance are based on a truncated estimator. That is, if the last observation(s) is not a death, then the survival curve estimate does not go to zero and the mean is undefined. There are four possible approaches to resolve this, which are selected by the rmean option. The first is to set the upper limit to a constant, e.g., rmean=365. In this case the reported mean would be the expected number of days, out of the first 365, that would be experienced by each group. This is useful if interest focuses on a fixed period. Other options are "none" (no estimate), "common" and "individual". The "common" option uses the maximum time for all curves in the object as a common upper limit for the auc calculation. For the "individual" options the mean is computed as the area under each curve, over the range from 0 to the maximum observed time for that curve. Since the end point is random, values for different curves are not comparable and the printed standard errors are an underestimate as they do not take into account this random variation. This option is provided mainly for backwards compatibility, as this estimate was the default (only) one in earlier releases of the code. Note that SAS (as of version 9.3) uses the integral up to the last event time of each individual curve; we consider this the worst of the choices and do not provide an option for that calculation.

The median and its confidence interval are defined by drawing a horizontal line at 0.5 on the plot of the survival curve and its confidence bands. If that line does not intersect the curve, then the median is undefined. The intersection of the line with the lower CI band defines the lower limit for the median’s interval, and similarly for the upper band. If any of the intersections is not a point then
we use the center of the intersection interval, e.g., if the survival curve were exactly equal to 0.5 over an interval. When data is uncensored this agrees with the usual definition of a median.

Value

x, with the invisible flag set to prevent printing. (The default for all print functions in R is to return the object passed to them; print.survfit complies with this pattern. If you want to capture these printed results for further processing, see the table component of summary.survfit.)

Side Effects

The number of observations, the number of events, the median survival with its confidence interval, and optionally the restricted mean survival (rmean) and its standard error, are printed. If there are multiple curves, there is one line of output for each.

References


See Also

summary.survfit, quantile.survfit

Description

Produce pseudo values from a survival curve.

Usage

pseudo(fit, times, type, addNA=TRUE, data.frame=FALSE, minus1=FALSE, ...)

Arguments

fit a survfit object, or one that inherits that class.
times a vector of time points, at which to evaluate the pseudo values.
type the type of value, either the probability in state pstate, the cumulative hazard cumhaz or the expected sojourn time in the state sojourn.
addNA If any observations were removed due to missing values in the fit object, add those rows (as NA) into the return. This causes the result of pseudo to match the original dataframe.
data.frame if TRUE, return the data in "long" form as a data.frame with id, time, and pseudo as variables.
minus1 use n-1 as the multiplier rather than n.
... other arguments to the residuals.survfit function, which does the majority of the work, e.g., collapse and weighted.
Details

This function computes pseudo values based on a first order Taylor series, also known as the "infinitesimal jackknife" (IJ) or "dfbeta" residuals. To be completely correct these results could perhaps be called 'IJ pseudo values' or even pseudo pseudo-values. For moderate to large data, however, the resulting values will be almost identical, numerically, to the ordinary jackknife.

A primary advantage of this approach is computational speed. Other features, neither good nor bad, are that they will agree with robust standard errors of other survival package estimates, which are based on the IJ, and that the mean of the estimates, over subjects, is exactly the underlying survival estimate.

For the type variable, surv is an acceptable synonym for pstate, and rmst, rmts are equivalent to sojourn. All of these are case insensitive.

Value

A vector, matrix, or array. The first dimension is always the number of observations in fit object, in the same order as the original data set (less any missing values that were removed when creating the survfit object); the second, if applicable, corresponds to fit$states, e.g., multi-state survival, and the last dimension to the selected time points.

For the data.frame option, a data frame containing values for id, time, and pseudo. If the original survfit call contained an id statement, then the values in the id column will be taken from that variable. If the id statement has a simple form, e.g., id = patno, then the name of the id column will be 'patno', otherwise it will be named '(id)'.

References


See Also

residuals.survfit

Examples

fit1 <- survfit(Surv(time, status) ~ 1, data=lung)
yhat <- pseudo(fit1, times=c(365, 730))
dim(yhat)

# Restricted Mean Time in State (RMST)
rms <- pseudo(fit1, times= 730, type='RMST') # 2 years
rfit <- lm(rms ~ ph.ecog + sex, data=lung)
rhat <- predict(rfit, newdata=expand.grid(ph.ecog=0:3, sex=1:2), se.fit=TRUE)
# print it out nicely
temp1 <- cbind(matrix(rhat$fit, 4,2))
temp2 <- cbind(matrix(rhat$se.fit, 4, 2))
temp3 <- cbind(temp1[,1], temp2[,1], temp1[,2], temp2[,2])
dimnames(temp3) <- list(paste("ph.ecog", 0:3),
c("Male RMST", "(se)", "Female RMST", "(se)"))
round(temp3, 1)
# compare this to the fully non-parametric estimate
fit2 <- survfit(Surv(time, status) ~ ph.ecog, data=lung)
print(fit2, rmean=730)
# the estimate for ph.ecog=3 is very unstable (n=1), pseudovalues smooth it.
#
# In all the above we should be using the robust variance, e.g., svyglm, but
# a recommended package can't depend on external libraries.
# See the vignette for a more complete exposition.

---

**pspline**  
*Smoothing splines using a pspline basis*

**Description**

Specifies a penalised spline basis for the predictor. This is done by fitting a comparatively small set of splines and penalising the integrated second derivative. Traditional smoothing splines use one basis per observation, but several authors have pointed out that the final results of the fit are indistinguishable for any number of basis functions greater than about 2-3 times the degrees of freedom. Eilers and Marx point out that if the basis functions are evenly spaced, this leads to significant computational simplification, they refer to the result as a p-spline.

**Usage**

```r
pspline(x, df=4, theta, nterm=2.5 * df, degree=3, eps=0.1, method,
        Boundary.knots=range(x), intercept=FALSE, penalty=TRUE, combine, ...)
```

**Arguments**

- **x** for psline: a covariate vector. The function does not apply to factor variables. For psplineinverse x will be the result of a psline call.
- **df** the desired degrees of freedom. One of the arguments df or theta’ must be given, but not both. If df=0, then the AIC = (loglik -df) is used to choose an "optimal" degrees of freedom. If AIC is chosen, then an optional argument ‘caic=T’ can be used to specify the corrected AIC of Hurvich et. al.
- **theta** roughness penalty for the fit. It is a monotone function of the degrees of freedom, with theta=1 corresponding to a linear fit and theta=0 to an unconstrained fit of nterm degrees of freedom.
- **nterm** number of splines in the basis
- **degree** degree of splines
- **eps** accuracy for df
the method for choosing the tuning parameter theta. If theta is given, then
'fixed' is assumed. If the degrees of freedom is given, then 'df' is assumed. If
method='aic' then the degrees of freedom is chosen automatically using Akaike's
information criterion.

... optional arguments to the control function

the spline is linear beyond the boundary knots. These default to the range of the
data.

if TRUE, the basis functions include the intercept.

if FALSE a large number of attributes having to do with penalized fits are ex-
cluded. This is useful to create a pspline basis matrix for other uses.

an optional vector of increasing integers. If two adjacent values of combine are
equal, then the corresponding coefficients of the fit are forced to be equal. This
is useful for monotone fits, see the vignette for more details.

Object of class pspline, coxph.penalty containing the spline basis, with the appropriate attributes
to be recognized as a penalized term by the coxph or survreg functions.

For psplineinverse the original x vector is reconstructed.

Eilers, Paul H. and Marx, Brian D. (1996). Flexible smoothing with B-splines and penalties. Sta-
tistical Science, 11, 89-121.

in nonparametric regression using an improved Akaike information criterion, JRSSB, volume 60,
271–293.

See Also

coxph, survreg, ridge, frailty

Examples

lfit6 <- survreg(Surv(time, status)~pspline(age, df=2), lung)
plot(lung$age, predict(lfit6), xlab='Age', ylab="Spline prediction")
title("Cancer Data")
fit0 <- coxph(Surv(time, status) ~ ph.ecog + age, lung)
fit1 <- coxph(Surv(time, status) ~ ph.ecog + pspline(age,3), lung)
fit3 <- coxph(Surv(time, status) ~ ph.ecog + pspline(age,8), lung)
fit0
fit1
fit3
Description

This function computes the person-years of follow-up time contributed by a cohort of subjects, stratified into subgroups. It also computes the number of subjects who contribute to each cell of the output table, and optionally the number of events and/or expected number of events in each cell.

Usage

```r
pyears(formula, data, weights, subset, na.action, rmap, ratetable, scale=365.25, expect=c('event', 'pyears'), model=FALSE, x=FALSE, y=FALSE, data.frame=FALSE)
```

Arguments

- **formula**: a formula object. The response variable will be a vector of follow-up times for each subject, or a `Surv` object containing the survival time and an event indicator. The predictors consist of optional grouping variables separated by `+` operators (exactly as in `survfit`), time-dependent grouping variables such as age (specified with `tcut`), and optionally a `ratetable` term. This latter matches each subject to his/her expected cohort.
- **data**: a data frame in which to interpret the variables named in the `formula`, or in the `subset` and the `weights` argument.
- **weights**: case weights.
- **subset**: expression saying that only a subset of the rows of the data should be used in the fit.
- **na.action**: a missing-data filter function, applied to the model.frame, after any `subset` argument has been used. Default is `options()$na.action`.
- **rmap**: an optional list that maps data set names to the ratetable names. See the details section below.
- **ratetable**: a table of event rates, such as `survexp.uswhite`.
- **scale**: a scaling for the results. As most rate tables are in units/day, the default value of 365.25 causes the output to be reported in years.
- **expect**: should the output table include the expected number of events, or the expected number of person-years of observation. This is only valid with a rate table.
- **data.frame**: return a data frame rather than a set of arrays.
- **model, x, y**: If any of these is true, then the model frame, the model matrix, and/or the vector of response times will be returned as components of the final result.
Details

Because `pyyears` may have several time variables, it is necessary that all of them be in the same units. For instance, in the call

```r
py <- pyyears(futime ~ rx, rmap=list(age=age, sex=sex, year=entry.dt),
             ratetable=survexp.us)
```

the natural unit of the ratetable is hazard per day, it is important that `futime`, `age` and `entry.dt` all be in days. Given the wide range of possible inputs, it is difficult for the routine to do sanity checks of this aspect.

The ratetable being used may have different variable names than the user’s data set, this is dealt with by the `rmap` argument. The rate table for the above calculation was `survexp.us`, a call to `summary(survexp.us)` reveals that it expects to have variables `age` = age in days, `sex`, and `year` = the date of study entry, we create them in the `rmap` line. The sex variable is not mapped, therefore the code assumes that it exists in `mydata` in the correct format. (Note: for factors such as `sex`, the program will match on any unique abbreviation, ignoring case.)

A special function `tcut` is needed to specify time-dependent cutpoints. For instance, assume that age is in years, and that the desired final arrays have as one of their margins the age groups 0-2, 2-10, 10-25, and 25+. A subject who enters the study at age 4 and remains under observation for 10 years will contribute follow-up time to both the 2-10 and 10-25 subsets. If `cut(age, c(0,2,10,25,100))` were used in the formula, the subject would be classified according to his starting age only. The `tcut` function has the same arguments as `cut`, but produces a different output object which allows the `pyyears` function to correctly track the subject.

The results of `pyyears` are normally used as input to further calculations. The `print` routine, therefore, is designed to give only a summary of the table.

Value

A list with components:

- `pyyears`: an array containing the person-years of exposure. (Or other units, depending on the rate table and the scale). The dimension and dimnames of the array correspond to the variables on the right hand side of the model equation.
- `n`: an array containing the number of subjects who contribute time to each cell of the `pyyears` array.
- `event`: an array containing the observed number of events. This will be present only if the response variable is a `Surv` object.
- `expected`: an array containing the expected number of events (or person years if `expect = "pyyears"`). This will be present only if there was a `ratetable` term.
- `data`: if the `data.frame` option was set, a data frame containing the variables `n`, `event`, `pyyears` and `event` that supplants the four arrays listed above, along with variables corresponding to each dimension. There will be one row for each cell in the arrays.
- `offtable`: the number of person-years of exposure in the cohort that was not part of any cell in the `pyyears` array. This is often useful as an error check; if there is a mismatch of units between two variables, nearly all the person years may be off table.
**tcut** whether the call included any time-dependent cutpoints.

**summary** a summary of the rate-table matching. This is also useful as an error check.

**call** an image of the call to the function.

**observations** the number of observations in the input data set, after any missings were removed.

**na.action** the na.action attribute contributed by an na.action routine, if any.

**See Also**

*ratetable, survexp, Surv.*

**Examples**

```r
# Look at progression rates jointly by calendar date and age
#
temp.yr <- tcut(mgus$dxyr, 55:92, labels=as.character(55:91))
temp.age <- tcut(mgus$age, 34:101, labels=as.character(34:100))
ptime <- ifelse(is.na(mgus$pctime), mgus$futime, mgus$pctime)
pstat <- ifelse(is.na(mgus$pctime), 0, 1)
pfit <- pyears(Surv(ptime/365.25, pstat) ~ temp.yr + temp.age + sex, mgus, data.frame=TRUE)

# Turn the factor back into numerics for regression
ndata <- pfit$data
ndata$age <- as.numeric(as.character(ndata$temp.age))
data$year<- as.numeric(as.character(ndata$temp.yr))
fit1 <- glm(event ~ year + age+ sex +offset(log(pyears)), data=ndata, family=poisson)

# Not run:
# fit a gam model
gfit.m <- gam(y ~ s(age) + s(year) + offset(log(time)),
family = poisson, data = tdata)

# Example #2 Create the hearta data frame:
hearta <- by(heart, heart$id,
function(x)x[x$stop == max(x$stop),])
hearta <- do.call("rbind", hearta)

# Produce pyears table of death rates on the surgical arm
# The first is by age at randomization, the second by current age
fit1 <- pyears(Surv(stop/365.25, event) ~ cut(age + 48, c(0,50,60,70,100)) + surgery, data = hearta, scale = 1)
fit2 <- pyears(Surv(stop/365.25, event) ~ tcut(age + 48, c(0,50,60,70,100)) + surgery, data = hearta, scale = 1)
fit1$event/fit1$pyears #death rates on the surgery and non-surg arm
fit2$event/fit2$pyears #death rates on the surgery and non-surg arm
```
quantile.survfit  
Quantiles from a survfit object

Description

Retrieve quantiles and confidence intervals for them from a survfit or Surv object.

Usage

```r
## S3 method for class 'survfit'
quantile(x, probs = c(0.25, 0.5, 0.75), conf.int = TRUE,
        scale, tolerance = sqrt(.Machine$double.eps), ...)

## S3 method for class 'survfitsms'
quantile(x,_probs = c(0.25, 0.5, 0.75), conf.int = TRUE,
        scale, tolerance = sqrt(.Machine$double.eps), ...)

## S3 method for class 'survfit'
median(x, ...)
```

Arguments

- `x`  
a result of the survfit function
- `probs`  
numeric vector of probabilities with values in [0,1]
- `conf.int`  
should lower and upper confidence limits be returned?
- `scale`  
optional scale factor, e.g., `scale=365.25` would return results in years if the fit object were in days.
- `tolerance`  
tolerance for checking that the survival curve exactly equals one of the quantiles
- `...`  
optional arguments for other methods

Details

The kth quantile for a survival curve \( S(t) \) is the location at which a horizontal line at height \( p = 1-k \) intersects the plot of \( S(t) \). Since \( S(t) \) is a step function, it is possible for the curve to have a horizontal segment at exactly 1-k, in which case the midpoint of the horizontal segment is returned. This mirrors the standard behavior of the median when data is uncensored. If the survival curve does not fall to 1-k, then that quantile is undefined.

In order to be consistent with other quantile functions, the argument `prob` of this function applies to the cumulative distribution function \( F(t) = 1-S(t) \).

Confidence limits for the values are based on the intersection of the horizontal line at 1-k with the upper and lower limits for the survival curve. Hence confidence limits use the same \( p \)-value as was in effect when the curve was created, and will differ depending on the `conf.type` option of `survfit`. If the survival curves have no confidence bands, confidence limits for the quantiles are not available.

When a horizontal segment of the survival curve exactly matches one of the requested quantiles the returned value will be the midpoint of the horizontal segment; this agrees with the usual definition of a median for uncensored data. Since the survival curve is computed as a series of products,
however, there may be round off error. Assume for instance a sample of size 20 with no tied times and no censoring. The survival curve after the 10th death is \((19/20)(18/19)(17/18) \ldots (10/11) = 10/20\), but the computed result will not be exactly 0.5. Any horizontal segment whose absolute difference with a requested percentile is less than `tolerance` is considered to be an exact match.

### Value

The quantiles will be a vector if the `survfit` object contains only a single curve, otherwise it will be a matrix or array. In this case the last dimension will index the quantiles.

If confidence limits are requested, then result will be a list with components `quantile`, `lower`, and `upper`, otherwise it is the vector or matrix of quantiles.

### Author(s)

Terry Therneau

### See Also

`survfit`, `print.survfit`, `qsurvreg`

### Examples

```r
fit <- survfit(Surv(time, status) ~ ph.ecog, data=lung)
quantile(fit)

cfit <- coxph(Surv(time, status) ~ age + strata(ph.ecog), data=lung)
csurv<- survfit(cfit, newdata=data.frame(age=c(40, 60, 80)),
               conf.type ="none")
temp <- quantile(csurv, 1:5/10)
temp[2,3,] # quantiles for second level of ph.ecog, age=80
quantile(csurv[2,3], 1:5/10) # quantiles of a single curve, same result
```

---

<table>
<thead>
<tr>
<th><code>ratetable</code></th>
<th>Rate table structure</th>
</tr>
</thead>
</table>

### Description

Description of the rate tables used by expected survival routines.

### Details

A rate table contains event rates per unit time for some particular endpoint. Death rates are the most common use, the `survexp.us` table, for instance, contains death rates for the United States by year of age, sex, and calendar year.

A rate table is structured as a multi-way array with the following attributes:

- `dim` the dimensions of the array
dimnames a named list of dimnames. The names are used to match user data to the dimensions, e.g., see the `rmap` argument in the `pyears` example. If a dimension is categorical, such as `sex` in `survexp.us`, then the dimname itself is matched against user’s data values. The matching ignores case and allows abbreviations, e.g., "M", "Male", and "m" all successfully match the `survexp.us` dimname of `sex=c("male","female")`.

type a vector giving the type of each dimension, which will be 1= categorical, 2= continuous, 3= date, 4= calendar year of a US rate table. If type is 3 or 4, then the corresponding cutpoints must be one of the calendar date types: Date, POSIXt, date, or chron. This allows the code to properly match user data to the ratetable. (The published US decennial rate tables’ definition is that a subject does not begin to experience a new years’ death rate on Jan 1, but rather on their next birthday. The actual impact of this delay on any given subjects’ calculation is negligible, but the code has always tried to be correct.)

cutpoints a list with one element per dimension. If type=1 then the corresponding list element should be NULL, otherwise it should be a vector of length `dim[i]` containing the starting point of the interval to which the corresponding row/col of the array applies. Cutpoints must be in the same units as the underlying table, e.g., the `survexp.us` table contains death rates per day, so the age cutpoint vector contains age in days while `year` contains a vector of Dates. Cutpoints do not need to be evenly spaced: the `survexp.us` table, for instance, originally had age divided up as 0-1 days, 1-7 days, 7-28 days, 28 days - 1 year, 2, 3, . . . 119 years. (Changes in the source of the tables made it difficult to continue splitting out the first year.)

summary an optional summarization function. If present, it will be called with a numeric matrix that has one column per dimension and one row per observation. The function returns a character string giving a summary of the data. This is used by some routines to print an informative message, and provides one way to inform users of a data mistake, e.g., if the printout states that all subjects are between 0.14 and 0.23 years old it is likely that the user’s age variable was in years when it should have been in days.

dimid optional attribute containing the names of the dimnames. If the dimnames list itself has names, this attribute will be ignored.

See Also

`survexp`, `pyears`, `survexp.us`

---

`ratetableDate` Convert date objects to ratetable form

---

Description

This method converts dates from various forms into the internal form used in `ratetable` objects.

Usage

`ratetableDate(x)`
Arguments

x a date. The function currently has methods for Date, date, POSIXt, timeDate, and chron objects.

Details

This function is useful for those who create new ratetables, but is normally invisible to users. It is used internally by the `survexp` and `pyears` functions to map the various date formats; if a new method is added then those routines will automatically be adapted to the new date type.

Value

a numeric vector, the number of days since 1/1/1960.

Author(s)

Terry Therneau

See Also

`pyears`, `survexp`

---

**Description**

Census data sets for the expected survival and person years functions.

**Usage**

data(survexp, package="survival")

**Details**

- **survexp.us** total United States population, by age and sex, 1940 to 2012.
- **survexp.usr** United States population, by age, sex and race, 1940 to 2014. Race is white or black. For 1960 and 1970 the black population values were not reported separately, so the nonwhite values were used. (Over the years, the reported tables have differed wrt reporting non-white and/or black.)
- **survexp.mn** total Minnesota population, by age and sex, 1970 to 2013.

Each of these tables contains the daily hazard rate for a matched subject from the population, defined as $-\log(1 - q)/365.25$ where $q$ is the 1 year probability of death as reported in the original tables from the US Census. For age 25 in 1970, for instance, $p = 1 - q$ is the probability that a subject who becomes 25 years of age in 1970 will achieve his/her 26th birthday. The tables are recast in terms of hazard per day for computational convenience.
Rat treatment data from Mantel et al. Three rats were chosen from each of 100 litters, one of which was treated with a drug, and then all followed for tumor incidence.

Usage

```r
rats
data(cancer, package="survival")
```

Format

- **litter**: litter number from 1 to 100
- **rx**: treatment, (1=drug, 0=control)
- **time**: time to tumor or last follow-up
- **status**: event status, 1=tumor and 0=censored
- **sex**: male or female

Note

Since only 2/150 of the male rats have a tumor, most analyses use only females (odd numbered litters), e.g. Lee et al.

Source

References
E. W. Lee, L. J. Wei, and D. Amato, Cox-type regression analysis for large number of small groups

rats2 Rat data from Gail et al.

Description
48 rats were injected with a carcinogen, and then randomized to either drug or placebo. The number
of tumors ranges from 0 to 13; all rats were censored at 6 months after randomization.

Usage
rats2
data(cancer, package="survival")

Format
rat: id
trt: treatment,(1=drug, 0=control)
observation: within rat
start: entry time
stop: exit time
status: event status, 1=tumor, 0=censored

Source
MH Gail, TJ Santner, and CC Brown (1980), An analysis of comparative carcinogenesis experi-
ments based on multiple times to tumor. Biometrics 36, 255–266.

reliability Reliability data sets

Description
A set of data for simple reliability analyses, taken from the book by Meeker and Escobar.

Usage
data(reliability, package="survival")
Details

- capacitor: Data from a factorial experiment on the life of glass capacitors as a function of voltage and operating temperature. There were 8 capacitors at each combination of temperature and voltage. Testing at each combination was terminated after the fourth failure.
  - temperature: temperature in degrees celcius
  - voltage: applied voltage
  - time: time to failure
  - status: 1=failed, 0=censored

- cracks: Data on the time until the development of cracks in a set of 167 identical turbine parts. The parts were inspected at 8 selected times.
  - day: time of inspection
  - fail: number of fans found to have cracks, at this inspection

- Data set genfan: Time to failure of 70 diesel engine fans.
  - hours: hours of service
  - status: 1=failure, 0=censored

Data set ifluid: A data frame with two variables describing the time to electrical breakdown of an insulating fluid.
  - time: hours to breakdown
  - voltage: test voltage in kV

- Data set imotor: Breakdown of motor insulation as a function of temperature.
  - temp: temperature of the test
  - time: time to failure or censoring
  - status: 0=censored, 1=failed

- Data set turbine: Each of 432 turbine wheels was inspected once to determine whether a crack had developed in the wheel or not.
  - hours: time of inspection (100s of hours)
  - inspected: number that were inspected
  - failed: number that failed

Data set valveSeat: Time to replacement of valve seats for 41 diesel engines. More than one seat may be replaced at a particular service, leading to duplicate times in the data set. The final inspection time for each engine will have status=0.
  - id: engine identifier
  - time: time of the inspection, in days
  - status: 1=replacement occured, 0= not

References


Examples

```
survreg(Surv(time, status) ~ temperature + voltage, capacitor)
```
residuals.coxph  Calculate Residuals for a ‘coxph’ Fit

Description

Calculates martingale, deviance, score or Schoenfeld residuals for a Cox proportional hazards model.

Usage

## S3 method for class 'coxph'
residuals(object,
  type=c("martingale", "deviance", "score", "schoenfeld",
         "dfbeta", "dfbetas", "scaledsch","partial"),
  collapse=FALSE, weighted= (type %in% c("dfbeta", "dfbetas")), ...)

## S3 method for class 'coxphms'
residuals(object,
  type=c("martingale", "score", "schoenfeld",
         "dfbeta", "dfbetas", "scaledsch"),
  collapse=FALSE, weighted= FALSE, ...)

## S3 method for class 'coxph.null'
residuals(object,
  type=c("martingale", "deviance","score","schoenfeld"),
  collapse=FALSE, weighted= FALSE, ...)

Arguments

object
  an object inheriting from class coxph, representing a fitted Cox regression model. Typically this is the output from the coxph function.

type
  character string indicating the type of residual desired. Possible values are "martingale", "deviance", "score", "schoenfeld", "dfbeta", "dfbetas", "scaledsch" and "partial". Only enough of the string to determine a unique match is required.

collapse
  vector indicating which rows to collapse (sum) over. In time-dependent models more than one row data can pertain to a single individual. If there were 4 individuals represented by 3, 1, 2 and 4 rows of data respectively, then collapse=c(1,1,1,2,3,3,4,4,4,4) could be used to obtain per subject rather than per observation residuals.

weighted
  if TRUE and the model was fit with case weights, then the weighted residuals are returned.

...
  other unused arguments

Value

For martingale and deviance residuals, the returned object is a vector with one element for each subject (without collapse). For score residuals it is a matrix with one row per subject and one column per variable. The row order will match the input data for the original fit. For Schoenfeld
residuals, the returned object is a matrix with one row for each event and one column per variable. The rows are ordered by time within strata, and an attribute strata is attached that contains the number of observations in each strata. The scaled Schoenfeld residuals are used in the \texttt{cox.zph} function.

The score residuals are each individual’s contribution to the score vector. Two transformations of this are often more useful: \texttt{dfbeta} is the approximate change in the coefficient vector if that observation were dropped, and \texttt{dfbetas} is the approximate change in the coefficients, scaled by the standard error for the coefficients.

\textbf{NOTE}

For deviance residuals, the status variable may need to be reconstructed. For score and Schoenfeld residuals, the \texttt{X} matrix will need to be reconstructed.

\textbf{References}


\textbf{See Also}

\texttt{coxph}

\textbf{Examples}

```r
fit <- coxph(Surv(start, stop, event) ~ (age + surgery)* transplant, 
              data=heart)
mresid <- resid(fit, collapse=heart$id)
```

\texttt{residuals.survfit} \hspace{1cm} \textit{IJ residuals from a survfit object.}

\textbf{Description}

Return infinitesimal jackknife residuals from a survfit object, for the survival, cumulative hazard, or restricted mean time in state (RMTS).

\textbf{Usage}

```r
## S3 method for class 'survfit'
residuals(object, times, 
          type="pstate", collapse, weighted=FALSE, 
          method=1, ...)
```
residuals.survfit

Arguments

- **object**: a `survfit` object
- **times**: a vector of times at which the residuals are desired
- **type**: the type of residual, see below
- **collapse**: add the residuals for all subjects in a cluster
- **weighted**: weight the residuals by each observation’s weight
- **method**: controls a choice of algorithm. Current an internal debugging option.
- ... arguments for other methods

Details

This function is designed to efficiently compute the leverage residuals at a small number of time points; a primary use is the creation of pseudo-values. If the residuals at all time points are needed, e.g. to compute a robust pointwise confidence interval for the survival curve, then this can be done more efficiently using the influence argument of the underlying `survfit` function. But be aware that such matrices can get very large.

The residuals are the impact of each observation or cluster on the resulting probability in state curves at the given time points, the cumulative hazard curve at those time points, or the expected sojourn time in each state up to the given time points. For a simple Kaplan-Meier the `survfit` object contains only the probability in the "initial" state, i.e., the survival fraction. For the KM case the sojourn time, the expected amount of time spent in the initial state, up to the specified endpoint, is more commonly known as the restricted mean survival time (RMST). For a multistate model this same quantity is also referred to as the restricted mean time in state (RMTS). It can be computed as the area under the respective probability in state curve. The program allows any of `pstate`, `surv`, `cumhaz`, `chaz`, `sojourn`, `rmst`, `rmts` or `auc` for the type argument, ignoring upper/lowercase, so users can choose whichever abbreviation they like best.

When `collapse=TRUE` the result has the cluster identifier (which defaults to the id variable) as the dimname for the first dimension. If the fit object contains more than one curve, and the same identifier is reused in two different curves this approach does not work and the routine will stop with an error. In principle this is not necessary, e.g., the result could contain two rows with the same label, showing the separate effect on each curve, but this was deemed too confusing.

Value

A matrix or array with one row per observation or cluster, and one column for each value in `times`. For a multi-state model the three dimensions are observation, time and state. For cumulative hazard, the last dimension is the set of transitions. (A competing risks model for instance has 3 states and 2 transitions.)

See Also

- `survfit`, `survfit.formula`

Examples

```r
fit <- survfit(Surv(time, status) ~ x, aml)
resid(fit, times=c(24, 48), type="RMTS")
```
residuals.survreg  
Compute Residuals for 'survreg' Objects

Description

This is a method for the function residuals for objects inheriting from class survreg.

Usage

```r
## S3 method for class 'survreg'
residuals(object, type=c("response", "deviance", "dfbeta", "dfbetas", "working", "ldcase", "ldresp", "ldshape", "matrix"), rsigma=TRUE, collapse=FALSE, weighted=FALSE, ...)
```

Arguments

- **object**: an object inheriting from class survreg.
- **type**: type of residuals, with choices of "response", "deviance", "dfbeta", "dfbetas", "working", "ldcase", "ldresp", "ldshape", and "matrix".
- **rsigma**: include the scale parameters in the variance matrix, when doing computations. (I can think of no good reason not to).
- **collapse**: optional vector of subject groups. If given, this must be of the same length as the residuals, and causes the result to be per group residuals.
- **weighted**: give weighted residuals? Normally residuals are unweighted.
- **...**: other unused arguments

Value

A vector or matrix of residuals is returned. Response residuals are on the scale of the original data, working residuals are on the scale of the linear predictor, and deviance residuals are on log-likelihood scale. The dfbeta residuals are a matrix, where the ith row gives the approximate change in the coefficients due to the addition of subject i. The dfbetas matrix contains the dfbeta residuals, with each column scaled by the standard deviation of that coefficient.

The matrix type produces a matrix based on derivatives of the log-likelihood function. Let $L$ be the log-likelihood, $p$ be the linear predictor $X\beta$, and $s$ be $\log(\sigma)$. Then the 6 columns of the matrix are $L$, $dL/dp$, $\partial^2L/\partial p^2$, $dL/ds$, $\partial^2L/\partial s^2$, and $\partial^2L/\partial p\partial s$. Diagnostics based on these quantities are discussed in the book and article by Escobar and Meeker. The main ones are the likelihood displacement residuals for perturbation of a case weight (ldcase), the response value (ldresp), and the shape.

For a transformed distribution such as the log-normal or Weibull, matrix residuals are based on the log-likelihood of the transformed data $\log(y)$. For a monotone function $f$ the density of $f(X)$ is the density of $X$ divided by the derivative of $f$ (the Jacobian), so subtract $\log(\text{derivative})$ from each uncensored observation’s loglik value in order to match the loglik component of the result. The other columns of the matrix residual are unchanged by the transformation.
References


See Also

*predict.survreg*

Examples

```r
fit <- survreg(Surv(futime, death) ~ age + sex, mgus2)
summary(fit)  # age and sex are both important

rr <- residuals(fit, type='matrix')
sum(rr[,1]) - with(mgus2, sum(log(futime[death==1]))) # loglik

plot(mgus2$age, rr[,2], col= (1+mgus2$death)) # ldresp
```

---

**retinopathy**

*Diabetic Retinopathy*

Description

A trial of laser coagulation as a treatment to delay diabetic retinopathy.

Usage

```r
retinopathy
data(retinopathy, package="survival")
```

Format

A data frame with 394 observations on the following 9 variables.

- **id**: numeric subject id
- **laser**: type of laser used: xenon argon
- **eye**: which eye was treated: right left
- **age**: age at diagnosis of diabetes
- **type**: type of diabetes: juvenile adult, (diagnosis before age 20)
- **trt**: 0 = control eye, 1 = treated eye
- **futime**: time to loss of vision or last follow-up
- **status**: 0 = censored, 1 = loss of vision in this eye
- **risk**: a risk score for the eye. This high risk subset is defined as a score of 6 or greater in at least one eye.
Details

The 197 patients in this dataset were a 50% random sample of the patients with "high-risk" diabetic retinopathy as defined by the Diabetic Retinopathy Study (DRS). Each patient had one eye randomized to laser treatment and the other eye received no treatment, and has two observations in the data set. For each eye, the event of interest was the time from initiation of treatment to the time when visual acuity dropped below 5/200 two visits in a row. Thus there is a built-in lag time of approximately 6 months (visits were every 3 months). Survival times in this dataset are the actual time to vision loss in months, minus the minimum possible time to event (6.5 months). Censoring was caused by death, dropout, or end of the study.

References


Examples

```r
coxph(Surv(futime, status) ~ type + trt, cluster= id, retinopathy)
```

---

Description

Results of a randomized trial of rhDNase for the treatment of cystic fibrosis.

Usage

```r
rhDNase
data(rhDNase, package="survival")
```

Format

A data frame with 767 observations on the following 8 variables.

- `id`  subject id
- `inst` enrolling institution
- `trt` treatment arm: 0=placebo, 1= rhDNase
- `entry.dt` date of entry into the study
- `end.dt` date of last follow-up
- `fev` forced expiratory volume at enrollment, a measure of lung capacity
- `ivstart` days from enrollment to the start of IV antibiotics
- `ivstop` days from enrollment to the cessation of IV antibiotics
Details

In patients with cystic fibrosis, extracellular DNA is released by leukocytes that accumulate in the airways in response to chronic bacterial infection. This excess DNA thickens the mucus, which then cannot be cleared from the lung by the cilia. The accumulation leads to exacerbations of respiratory symptoms and progressive deterioration of lung function. At the time of this study more than 90% of cystic fibrosis patients eventually died of lung disease.

Deoxyribonuclease I (DNase I) is a human enzyme normally present in the mucus of human lungs that digests extracellular DNA. Genentech, Inc. cloned a highly purified recombinant DNase I (rhDNase or Pulmozyme) which when delivered to the lungs in an aerosolized form cuts extracellular DNA, reducing the viscoelasticity of airway secretions and improving clearance. In 1992 the company conducted a randomized double-blind trial comparing rhDNase to placebo. Patients were then monitored for pulmonary exacerbations, along with measures of lung volume and flow. The primary endpoint was the time until first pulmonary exacerbation; however, data on all exacerbations were collected for 169 days.

The definition of an exacerbation was an infection that required the use of intravenous (IV) antibiotics. Subjects had 0-5 such episodes during the trial, those with more than one have multiple rows in the data set, those with none have NA for the IV start and end times. A few subjects were infected at the time of enrollment, subject 173 for instance has a first infection interval of -21 to 7. We do not count this first infection as an "event", and the subject first enters the risk set at day 7. Subjects who have an event are not considered to be at risk for another event during the course of antibiotics, nor for an additional 6 days after they end. (If the symptoms reappear immediately after cessation then from a medical standpoint this would not be a new infection.)

This data set reproduces the data in Therneau and Grambsch, is does not exactly reproduce those in Therneau and Hamilton due to data set updates.

References


Examples

# Build the start-stop data set for analysis, and
# replicate line 2 of table 8.13
first <- subset(rhDNase, !duplicated(id)) # first row for each subject
dnase <- tmerge(first, first, id=id, tstop=as.numeric(end.dt-entry.dt))

# Subjects whose fu ended during the 6 day window are the reason for
# this next line
temp.end <- with(rhDNase, pmin(ivstop+6, end.dt-entry.dt))
dnase <- tmerge(dnase, rhDNase, id=id,
               infect=event(ivstart),
               end= event(temp.end))
# toss out the non-at-risk intervals, and extra variables
# 3 subjects had an event on their last day of fu, infect=1 and end=1
dnase <- subset(dnase, (infect==1 | end==0), c(id:trt, fev:infect))
ridge <- coxph(Surv(tstart, tstop, infect) ~ trt + fev, cluster=id, 
               data=dnase)

ridge

Ridge regression

Description

When used in a `coxph` or `survreg` model formula, specifies a ridge regression term. The likelihood is penalised by $\theta/2$ time the sum of squared coefficients. If `scale=TRUE` the penalty is calculated for coefficients based on rescaling the predictors to have unit variance. If `df` is specified then $\theta$ is chosen based on an approximate degrees of freedom.

Usage

ridge(..., theta, df=nvar/2, eps=0.1, scale=TRUE)

Arguments

... predictors to be ridged
theta penalty is $\theta/2$ time sum of squared coefficients
df Approximate degrees of freedom
eps Accuracy required for df
scale Scale variables before applying penalty?

Value

An object of class `coxph.penalty` containing the data and control functions.

Note

If the expression `ridge(x1,x2,x3,...)` is too many characters long then the internal `terms()` function will add newlines to the variable name and then the `coxph` routine simply gets lost. (Some labels will have the newline and some won’t.) One solution is to bundle all of the variables into a single matrix and use that matrix as the argument to `ridge` so as to shorten the call, e.g. `mdata$many <- as.matrix(mydata[,5:53])`.

References

Gray (1992) "Flexible methods of analysing survival data using splines, with applications to breast cancer prognosis" JASA 87:942–951

See Also

coxph, survreg, pspline, frailty
Examples

coxph(Surv(futime, fustat) ~ rx + ridge(age, ecog.ps, theta=1), ovarian)

lfit0 <- survreg(Surv(time, status) ~ 1, lung)
lfit1 <- survreg(Surv(time, status) ~ age + ridge(ph.ecog, theta=5), lung)
lfit2 <- survreg(Surv(time, status) ~ sex + ridge(age, ph.ecog, theta=1), lung)
lfit3 <- survreg(Surv(time, status) ~ sex + age + ph.ecog, lung)

Description

The rotterdam data set includes 2982 primary breast cancers patients whose data whose records were included in the Rotterdam tumor bank.

Usage

rotterdam
data(cancer, package="survival")

Format

A data frame with 2982 observations on the following 15 variables.

- pid: patient identifier
- year: year of surgery
- age: age at surgery
- meno: menopausal status (0= premenopausal, 1= postmenopausal)
- size: tumor size, a factor with levels <=20 20-50 >50
- grade: differentiation grade
- nodes: number of positive lymph nodes
- pgr: progesterone receptors (fmol/l)
- er: estrogen receptors (fmol/l)
- hormon: hormonal treatment (0= no, 1= yes)
- chemo: chemotherapy
- rtime: days to relapse or last follow-up
- recur: 0= no relapse, 1= relapse
- dtime: days to death or last follow-up
- death: 0= alive, 1= dead
Details

These data sets are used in the paper by Royston and Altman that is referenced below. The Rotterdam data is used to create a fitted model, and the GBSG data for validation of the model. The paper gives references for the data source.

There are 43 subjects who have died without recurrence, but whose death time is greater than the censoring time for recurrence. A common way that this happens is that a death date is updated in the health record sometime after the research study ended, and said value is then picked up when a study data set is created. But it raises serious questions about censoring. For instance subject 40 is censored for recurrence at 4.2 years and died at 6.6 years; when creating the endpoint of recurrence free survival (earlier of recurrence or death), treating them as a death at 6.6 years implicitly assumes that they were recurrence free just before death. For this to be true we would have to assume that if they had progressed in the 2.4 year interval before death (while off study), that this information would also have been noted in their general medical record, and would also be captured in the study data set. However, that may be unlikely. Death information is often in a centralized location in electronic health records, easily accessed by a programmer and merged with the study data, while recurrence may require manual review. How best to address this is an open issue.

References


See Also

gbsg

Examples

```r
status <- pmax(rotterdam$recur, rotterdam$death)
rfstime <- with(rotterdam, ifelse(recur==1, rtime, dtime))
fit1 <- coxph(Surv(rfstime, status) ~ pspline(age) + meno + size +
              pspline(nodes) + er,
              data=rotterdam, subset = (nodes > 0))
# Royston and Altman used fractional polynomials for the nonlinear terms
```

---

**royston**

*Compute Royston’s D for a Cox model*

**Description**

Compute the D statistic proposed by Royston and Sauerbrei along with several pseudo-R square values.

**Usage**

```r
royston(fit, newdata, ties = TRUE, adjust = FALSE)
```
Arguments

fit      a coxph fit
newdata  optional validation data set
ties     make a correction for ties in the risk score
adjust   adjust for possible overfitting

Details

These values are called pseudo R-squared since they involve only the linear predictor, and not the outcome. $R_D$ is the value that corresponds the Royston and Sauerbrei $D$ statistic. $R_KO$ is the value proposed by Kent and O’Quigley, $R_N$ is the value proposed by Nagelkerke, and $C_{GH}$ corresponds to Goen and Heller’s concordance measure.

An adjustment for $D$ is based on the ratio $r = \frac{\text{number of events}}{\text{number of coefficients}}$. For models which have sufficient sample size ($r > 20$) the adjustment will be small.

The Nagelkerke value is the Cox-Snell R-squared divided by a scaling constant. The two separate values are present in the result of `summary.coxph` as a 2 element vector `rsq`, and were listed as "Rsquare" and "max possible" in older versions of the print routine. (Since superseded in the default printout by the concordance.) The Nagelkerke estimate is not returned when `newdata` is present.

Value

a vector containing the value of $D$, the estimated standard error of $D$, and three or four pseudo R-squared values.

References


Examples

# An example used in Royston and Sauerbrei
cbc2 <- na.omit(pbc)  # no missing values
cfit <- coxph(Surv(time, status==2) ~ age + log(bili) + edema + albumin + 
               stage + copper, data= CBC2, ties="breslow")
royston(cfit)
rttright

**Compute redistribute-to-the-right weights**

**Description**

For many survival estimands, one approach is to redistribute each censored observation’s weight to those other observations with a longer survival time (think of distributing an estate to the heirs). Then compute on the remaining, uncensored data.

**Usage**

rttright(formula, data, weights, subset, na.action, times, id, timefix = TRUE)

**Arguments**

- **formula**: a formula object, which must have a `Surv` object as the response on the left of the `~` operator and, if desired, terms separated by `+` operators on the right. Each unique combination of predictors will define a separate strata.
- **data**: a data frame in which to interpret the variables named in the formula, subset and weights arguments.
- **weights**: The weights must be nonnegative and it is strongly recommended that they be strictly positive, since zero weights are ambiguous, compared to use of the subset argument.
- **subset**: expression saying that only a subset of the rows of the data should be used in the fit.
- **na.action**: a missing-data filter function, applied to the model frame, after any subset argument has been used. Default is `options()$na.action`.
- **times**: a vector of time points, for which to return updated weights. If missing, a time after the largest time in the data is assumed.
- **id**: optional: if the data set has multiple rows per subject, a variable containing the subject identifier of each row.
- **timefix**: correct for possible round-off error

**Details**

The `formula` argument is treated exactly the same as in the `survfit` function.

Redistribution is recursive: redistribute the weight of the first censored observation to all those with longer time, which may include other censored observations. Then redistribute the next smallest and etc. up to the specified time value. After re-distributing the weight for a censored observation to other observations that are not censored, ordinary non-censored methods can often be applied. For example, redistribution of the weights, followed by computation of the weighted cumulative distribution function, reprises the Kaplan-Meier estimator.

A primary use of this routine is illustration of methods or exploration of new methods. Methods that use RTTR directly, such as the Brier score, will normally do these computations internally.
Value

a vector or matrix of weights, with one column for each requested time

See Also

survfit

Examples

afit <- survfit(Surv(time, status) ~1, data=aml)
rwt <- rttright(Surv(time, status) ~1, data=aml)

index <- order(aml$time)
cdf <- cumsum(rwt[index]) # weighted CDF
cdf <- cdf![duplicated(aml$time[index], fromLast=TRUE)] # remove duplicates
cbind(time=afit$time, KM= afit$surv, RTTR= 1-cdf)

Data from a soldering experiment

Description

In 1988 an experiment was designed and implemented at one of AT&T's factories to investigate alternatives in the "wave soldering" procedure for mounting electronic components to printed circuit boards. The experiment varied a number of factors relevant to the process. The response, measured by eye, is the number of visible solder skips.

Usage

solder
data(solder, package="survival")

Format

A data frame with 900 observations on the following 6 variables.

- Opening  the amount of clearance around the mounting pad (3 levels)
- Solder  the amount of solder (Thick or Thin)
- Mask  type and thickness of the material used for the solder mask (A1.5, A3, A6, B3, B6)
- PadType  the geometry and size of the mounting pad (10 levels)
- Panel  each board was divided into 3 panels
- skips  the number of skips
Details

This data set is used as a detailed example in chapter 1 of Chambers and Hastie. Observations 1-360 and 541-900 form a balanced design of $3 \times 2 \times 10 \times 3 = 180$ observations for four of the pad types (A1.5, A3, B3, B6), while rows 361-540 match 3 of the 6 Solder*Opening combinations with pad type A6 and the other 3 with pad type A3.

References


Examples

# The balanced subset used by Chambers and Hastie
# contains the first 180 of each mask and deletes mask A6.
index <- 1 + (1:nrow(solder)) - match(solder$Mask, solder$Mask)
solder.balance <- droplevels(subset(solder, Mask != "A6" & index <= 180))

Description

This contains the Stanford Heart Transplant data in a different format. The main data set is in `heart`.

Usage

stanford2

Format

- id: ID number
- time: survival or censoring time
- status: censoring status
- age: in years
- t5: T5 mismatch score

Source


See Also

`predict.survreg`, `heart`
Draw a state space figure.

Description

For multi-state survival models it is useful to have a figure that shows the states and the possible transitions between them. This function creates a simple "box and arrows" figure. It's goal was simplicity.

Usage

```
statefig(layout, connect, margin = 0.03, box = TRUE, cex = 1, col = 1, lwd=1, lty=1, bcol=col, acol=col, alwd=lwd, alty=lty, offset=0)
```

Arguments

- `layout` describes the layout of the boxes on the page. See the detailed description below.
- `connect` a square matrix with one row for each state. If `connect[i,j] != 0` then an arrow is drawn from state i to state j. The row names of the matrix are used as the labels for the states.
- `margin` the fraction of white space between the label and the surrounding box, and between the box and the arrows, as a function of the plot region size.
- `box` should boxes be drawn? TRUE or FALSE.
- `cex, col, lty, lwd` default graphical parameters used for the text and boxes. The last 3 can be a vector of values.
- `bcol` color for the box, if it differs from that used for the text.
- `acol, alwd, alty` color, line type and line width for the arrows.
- `offset` used to slight offset the arrows between two boxes x and y if there is a transition in both directions. The default of 0 leads to a double headed arrow in this case – to arrows are drawn but they coincide. A positive value causes each arrow to shift to the left, from the view of someone standing at the foot of a arrow and looking towards the arrowhead, a negative offset shifts to the right. A value of 1 corresponds to the size of the plotting region.

Details

The arguments for color, line type and line width can all be vectors, in which case they are recycled as needed. Boxes and text are drawn in the order of the rownames of `connect`, and arrows are drawn in the usual R matrix order.

The `layout` argument is normally a vector of integers, e.g., the vector (1, 3, 2) describes a layout with 3 columns. The first has a single state, the second column has 3 states and the third has 2. The coordinates of the plotting region are 0 to 1 for both x and y. Within a column the centers of the boxes are evenly spaced, with 1/2 a space between the boxes and the margin, e.g., 4 boxes would be
at 1/8, 3/8, 5/8 and 7/8. If layout were a 1 column matrix with values of (1, 3, 2) then the layout will have three rows with 1, 3, and 2 boxes per row, respectively. Alternatively, the user can supply a 2 column matrix that directly gives the centers.

The values of the connect matrix should be 0 for pairs of states that do not have a transition and values between 0 and 2 for those that do. States are connected by an arc that passes through the centers of the two boxes and a third point that is between them. Specifically, consider a line segment joining the two centers and erect a second segment at right angles to the midpoint of length \(d\) times the distance from center to midpoint. The arc passes through this point. A value of \(d=0\) gives a straight line, \(d=1\) a right hand half circle centered on the midpoint and \(d=-1\) a left hand half circle. The connect matrix contains values of \(d+1\) with \(-1 < d < 1\).

The connecting arrow are drawn from (center of box 1 + offset) to (center of box 2 + offset), where the the amount of offset (white space) is determined by the box and margin parameters. If a pair of states are too close together this can result in an arrow that points the wrong way.

Value

a matrix containing the centers of the boxes, with the invisible attribute set.

Note

The goal of this function is to make “good enough” figures as simply as possible, and thereby to encourage users to draw them. The layout argument was inspired by the diagram package, which can draw more complex and well decorated figures, e.g., many different shapes, shading, multiple types of connecting lines, etc., but at the price of greater complexity.

Because curved lines are drawn as a set of short line segments, line types have almost no effect for that case.

Author(s)

Terry Therneau

Examples

```r
# Draw a simple competing risks figure
states <- c("Entry", "Complete response", "Relapse", "Death")
connect <- matrix(0, 4, 4, dimnames=list(states, states))
connect[1, -1] <- c(1.1, 1, 0.9)
statefig(c(1, 3), connect)
```

---

**strata**

Identify Stratification Variables

Description

This is a special function used in the context of the Cox survival model. It identifies stratification variables when they appear on the right hand side of a formula.
**Usage**

```r
strata(..., na.group=FALSE, shortlabel, sep=',')
```

**Arguments**

- `...`: any number of variables. All must be the same length.
- `na.group`: a logical variable, if `TRUE`, then missing values are treated as a distinct level of each variable.
- `shortlabel`: if `TRUE` omit variable names from resulting factor labels. The default action is to omit the names if all of the arguments are factors, and none of them was named.
- `sep`: the character used to separate groups, in the created label

**Details**

When used outside of a `coxph` formula the result of the function is essentially identical to the `interaction` function, though the labels from `strata` are often more verbose.

**Value**

a new factor, whose levels are all possible combinations of the factors supplied as arguments.

**See Also**

`coxph`, `interaction`

**Examples**

```r
a <- factor(rep(1:3,4), labels=c("low", "medium", "high"))
b <- factor(rep(1:4,3))
levels(strata(b))
levels(strata(a,b,shortlabel=TRUE))

coxph(Surv(futime, fustat) ~ age + strata(rx), data=ovarian)
```

---

**summary.aareg**

**Summarize an aareg fit**

**Description**

Creates the overall test statistics for an Aalen additive regression model

**Usage**

```r
## S3 method for class 'aareg'
summary(object, maxtime, test=c("aalen", "nrisk"), scale=1,...)
```
Arguments

- **object**: the result of a call to the `aareg` function
- **maxtime**: truncate the input to the model at time "maxtime"
- **test**: the relative time weights that will be used to compute the test
- **scale**: scales the coefficients. For some data sets, the coefficients of the Aalen model will be very small (10^-4); this simply multiplies the printed values by a constant, say 1e6, to make the printout easier to read.

... for future methods

Details

It is not uncommon for the very right-hand tail of the plot to have large outlying values, particularly for the standard error. The maxtime parameter can then be used to truncate the range so as to avoid these. This gives an updated value for the test statistics, without refitting the model.

The slope is based on a weighted linear regression to the cumulative coefficient plot, and may be a useful measure of the overall size of the effect. For instance when two models include a common variable, "age" for instance, this may help to assess how much the fit changed due to the other variables, in lieu of overlaying the two plots. (Of course the plots are often highly non-linear, so it is only a rough substitute). The slope is not directly related to the test statistic, as the latter is invariant to any monotone transformation of time.

Value

A list is returned with the following components:

- **table**: a matrix with rows for the intercept and each covariate, and columns giving a slope estimate, the test statistic, it's standard error, the z-score and a p-value
- **test**: the time weighting used for computing the test statistics
- **test.statistic**: the vector of test statistics
- **test.var**: the model based variance matrix for the test statistic
- **test.var2**: optionally, a robust variance matrix for the test statistic
- **chisq**: the overall test (ignoring the intercept term) for significance of any variable
- **n**: a vector containing the number of observations, the number of unique death times used in the computation, and the total number of unique death times

See Also

- `aareg`, `plot.aareg`

Examples

```r
afit <- aareg(Surv(time, status) ~ age + sex + ph.ecog, data=lung, dfbeta=TRUE)
summary(afit)
## Not run:
```
### summary.coxph

**Summary method for Cox models**

**Description**

Produces a summary of a fitted `coxph` model

**Usage**

```r
## S3 method for class 'coxph'
summary(object, conf.int=0.95, scale=1,...)
```

**Arguments**

- **object**
  - the result of a `coxph` fit
- **conf.int**
  - level for computation of the confidence intervals. If set to FALSE no confidence intervals are printed
- **scale**
  - vector of scale factors for the coefficients, defaults to 1. The printed coefficients, se, and confidence intervals will be associated with one scale unit.
- **...**
  - for future methods

**Value**

An object of class `summary.coxph`, with components:

- **n, nevent**
  - number of observations and number of events, respectively, in the fit

---

```
summary.coxph

<table>
<thead>
<tr>
<th>slope</th>
<th>test se(test)</th>
<th>robust se</th>
<th>z</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intercept</td>
<td>5.05e-03</td>
<td>1.9</td>
<td>1.54</td>
<td>1.55</td>
</tr>
<tr>
<td>age</td>
<td>4.01e-05</td>
<td>108.0</td>
<td>109.00</td>
<td>106.00</td>
</tr>
<tr>
<td>sex</td>
<td>-3.16e-03</td>
<td>-19.5</td>
<td>5.90</td>
<td>5.95</td>
</tr>
<tr>
<td>ph.ecog</td>
<td>3.01e-03</td>
<td>33.2</td>
<td>9.18</td>
<td>9.17</td>
</tr>
</tbody>
</table>

Chisq=22.84 on 3 df, p=4.4e-05; test weights=aalen

## End(Not run)

```

```
summary(afit, maxtime=600)

## Not run:

<table>
<thead>
<tr>
<th>slope</th>
<th>test se(test)</th>
<th>robust se</th>
<th>z</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intercept</td>
<td>4.16e-03</td>
<td>2.13</td>
<td>1.48</td>
<td>1.47</td>
</tr>
<tr>
<td>age</td>
<td>2.82e-05</td>
<td>85.80</td>
<td>106.00</td>
<td>100.00</td>
</tr>
<tr>
<td>sex</td>
<td>-2.54e-03</td>
<td>-20.60</td>
<td>5.61</td>
<td>5.63</td>
</tr>
<tr>
<td>ph.ecog</td>
<td>2.47e-03</td>
<td>31.60</td>
<td>8.91</td>
<td>8.67</td>
</tr>
</tbody>
</table>

Chisq=27.08 on 3 df, p=5.7e-06; test weights=aalen

## End(Not run)

---

```
loglik          the log partial likelihood at the initial and final values
coefficients   a matrix with one row for each coefficient, and columns containing the coefficient, the hazard ratio \(\exp(\text{coef})\), standard error, Wald statistic, and P value.
conf.int       a matrix with one row for each coefficient, containing the confidence limits for \(\exp(\text{coef})\)
logtest, sctest, waldtest
               the overall likelihood ratio, score, and Wald test statistics for the model
concordance    the concordance statistic and its standard error
used.robust    whether an asymptotic or robust variance was used
rsq            an approximate R^2 based on Nagelkirke (Biometrika 1991).
fail           a message, if the underlying coxph call failed
call           a copy of the call
na.action      information on missing values

Note

The pseudo r-squared of Nagelkirke is attractive because it is simple, but further work has shown that it has poor properties and it is now deprecated. The value is no longer printed by default, and will eventually be removed from the object.

See Also

coxph, print.coxph

Examples

```r
fit <- coxph(Surv(time, status) ~ age + sex, lung)
summary(fit)
```

---

**summary.pyears**  
**Summary function for pyears objects**

**Description**

Create a printable table of a person-years result.

**Usage**

```r
## S3 method for class 'pyears'
summary(object, header = TRUE, call = header, n = TRUE, event = TRUE, pyears = TRUE, expected = TRUE, rate = FALSE, rr = expected, ci.r = FALSE, ci.rr = FALSE, totals=FALSE, legend = TRUE, vline = FALSE, vertical= TRUE, nastring=".", conf.level = 0.95, scale = 1, ...)
```
Arguments

object a pyears object
header print out a header giving the total number of observations, events, person-years, and total time (if any) omitted from the table
call print out a copy of the call
n, event, pyears, expected logical arguments: should these elements be printed in the table?
rate, ci.r logical arguments: should the incidence rate and/or its confidence interval be given in the table?
rr, ci.rr logical arguments: should the hazard ratio and/or its confidence interval be given in the table?
totals should row and column totals be added?
legend should a legend be included in the printout?
vline should vertical lines be included in the printed tables?
vertical when there is only a single predictor, should the table be printed with the predictor on the left (vertical=TRUE) or across the top (vertical=FALSE)?
nasttring what to use for missing values in the table. Some of these are structural, e.g., risk ratios for a cell with no follow-up time.
conf.level confidence level for any confidence intervals
scale a scaling factor for printed rates
... optional arguments which will be passed to the format function; common choices would be digits=2 or nsmall=1.

Details

The pyears function is often used to create initial descriptions of a survival or time-to-event variable; the type of material that is often found in “table 1” of a paper. The summary routine prints this information out using one of pandoc table styles. A primary reason for choosing this style is that Rstudio is then able to automatically render the results in multiple formats: html, rtf, latex, etc.

If the pyears call has only a single covariate then the table will have that covariate as one margin and the statistics of interest as the other. If the pyears call has two predictors then those two predictors are used as margins of the table, while each cell of the table contains the statistics of interest as multiple rows within the cell. If there are more than two predictors then multiple tables are produced, in the same order as the standard R printout for an array.

The "N" entry of a pyears object is the number of observations which contributed to a particular cell. When the original call includes tcut objects then a single observation may contribute to multiple cells.

Value

a copy of the object
Notes
The pandoc system has four table types: with or without vertical bars, and with single or multiple rows of data in each cell. This routine produces all 4 styles depending on options, but currently not all of them are recognized by the Rstudio-pandoc pipeline. (And we don’t yet see why.)

Author(s)
Terry Therneau and Elizabeth Atkinson

See Also
cipoisson, pyears, format

summary.survexp  Summary function for a survexp object

Description
Returns a list containing the values of the survival at specified times.

Usage
## S3 method for class 'survexp'
summary(object, times, scale = 1, ...)

Arguments
object the result of a call to the survexp function
times vector of times; the returned matrix will contain 1 row for each time. Missing values are not allowed.
scale numeric value to rescale the survival time, e.g., if the input data to survfit were in days, scale = 365.25 would scale the output to years.
... For future methods

Details
A primary use of this function is to retrieve survival at fixed time points, which will be properly interpolated by the function.

Value
a list with the following components:
surv the estimate of survival at time t.
time the timepoints on the curve.
n.risk In expected survival each subject from the data set is matched to a hypothetical person from the parent population, matched on the characteristics of the parent population. The number at risk is the number of those hypothetical subject who are still part of the calculation.
summary.survfit

Author(s)
Terry Therneau

See Also
survexp

summary.survfit  Summary of a Survival Curve

Description
Returns a list containing the survival curve, confidence limits for the curve, and other information.

Usage
## S3 method for class 'survfit'
summary(object, times, censored=FALSE, scale=1,
  extend=FALSE, rmean=getOption('survfit.rmean'), ...)

Arguments

object  the result of a call to the survfit function.
times  vector of times; the returned matrix will contain 1 row for each time. The vector will be sorted into increasing order; missing values are not allowed. If censored=T, the default times vector contains all the unique times in fit, otherwise the default times vector uses only the event (death) times.
censored  logical value: should the censoring times be included in the output? This is ignored if the times argument is present.
scale  numeric value to rescale the survival time, e.g., if the input data to survfit were in days, scale = 365.25 would scale the output to years.
extend  logical value: if TRUE, prints information for all specified times, even if there are no subjects left at the end of the specified times. This is only used if the times argument is present.
rmean  Show restricted mean: see print.survfit for details
...
  for future methods

Value

a list with the following components:
surv  the estimate of survival at time t+0.
time  the timepoints on the curve.
n.risk  the number of subjects at risk at time t-0 (but see the comments on weights in
the survfit help file).

n.event if the times argument is missing, then this column is the number of events that
occurred at time t. Otherwise, it is the cumulative number of events that have
occurred since the last time listed until time t+0.

n.entered This is present only for counting process survival data. If the times argument is
missing, this column is the number of subjects that entered at time t. Otherwise,
it is the cumulative number of subjects that have entered since the last time listed
until time t.

n.exit.censored if the times argument is missing, this column is the number of subjects that left
without an event at time t. Otherwise, it is the cumulative number of subjects
that have left without an event since the last time listed until time t+0. This is
only present for counting process survival data.

std.err  the standard error of the survival value.

conf.int level of confidence for the confidence intervals of survival.

lower   lower confidence limits for the curve.

upper   upper confidence limits for the curve.

strata  indicates stratification of curve estimation. If strata is not NULL, there are mul-
tiple curves in the result and the surv, time, n.risk, etc. vectors will contain
multiple curves, pasted end to end. The levels of strata (a factor) are the labels
for the curves.

call    the statement used to create the fit object.

na.action same as for fit, if present.

table  table of information that is returned from print.survfit function.

type    type of data censoring. Passed through from the fit object.

Details

This routine has two uses: printing out a survival curve at specified time points (often yearly), or
extracting the values at specified time points for further processing. In the first case we normally
want extend=FALSE, i.e., don’t print out data past the end of the curve. If the times option only
contains values beyond the last point in the curve then there is nothing to print and an error message
will result. For the second usage we almost always want extend=TRUE, so that the results will have
a predictable length.

The survfit object itself will have a row of information at each censoring or event time, it does
not save information on each unique entry time. For printout at two time points t1, t2, this function
will give the the number at risk at the smallest event times that are >= t1 and >= t2, respectively,
the survival curve at the largest recorded times <= t1 and <= t2, and the number of events and
censorings in the interval t1 < t <= t2.

When the routine is called with counting process data many users are confused by counts that are
too large. For example, Surv(c(0,0,5,5),c(2,3,8,10),c(1,0,1,0)) followed by a request for
the values at time 4. The survfit object has entries only at times 2, 3, 8, and 10; there are 2 subjects
at risk at time 8, so that is what will be printed.
Surv

See Also

`survfit`, `print.summary.survfit`

Examples

```r
summary( survfit( Surv(futime, fustat)~1, data=ovarian))
summary( survfit( Surv(futime, fustat)~rx, data=ovarian))
```

---

**surv**  
*Create a Survival Object*

**Description**

Create a survival object, usually used as a response variable in a model formula. Argument matching is special for this function, see Details below.

**Usage**

```r
Surv(time, time2, event, 
   type=c("right", "left", "interval", "counting", "interval2", "mstate"), 
   origin=0)
```

```r
is.Surv(x)
```

**Arguments**

- **time**
  for right censored data, this is the follow up time. For interval data, the first argument is the starting time for the interval.

- **event**
  The status indicator, normally 0=alive, 1=dead. Other choices are TRUE/FALSE (TRUE = death) or 1/2 (2=death). For interval censored data, the status indicator is 0=right censored, 1=event at time, 2=left censored, 3=interval censored. For multiple endpoint data the event variable will be a factor, whose first level is treated as censoring. Although unusual, the event indicator can be omitted, in which case all subjects are assumed to have an event.

- **time2**
  ending time of the interval for interval censored or counting process data only. Intervals are assumed to be open on the left and closed on the right, (start, end]. For counting process data, event indicates whether an event occurred at the end of the interval.

- **type**
  character string specifying the type of censoring. Possible values are "right", "left", "counting", "interval", "interval2" or "mstate".

- **origin**
  for counting process data, the hazard function origin. This option was intended to be used in conjunction with a model containing time dependent strata in order to align the subjects properly when they cross over from one strata to another, but it has rarely proven useful.

- **x**
  any R object.
Details

When the type argument is missing the code assumes a type based on the following rules:

- If there are two unnamed arguments, they will match time and event in that order. If there are three unnamed arguments they match time, time2 and event.
- If the event variable is a factor then type mstate is assumed. Otherwise type right if there is no time2 argument, and type counting if there is.

As a consequence the type argument will normally be omitted.

When the survival type is "mstate" then the status variable will be treated as a factor. The first level of the factor is taken to represent censoring and remaining ones a transition to the given state. (If the status variable is a factor then mstate is assumed.)

Interval censored data can be represented in two ways. For the first use type = "interval" and the codes shown above. In that usage the value of the time2 argument is ignored unless event=3. The second approach is to think of each observation as a time interval with (-infinity, t2) for left censored, (t1, infinity) for right censored, (t,t) for exact and (t1, t2) for an interval. This is the approach used for type = interval2. Infinite values can be represented either by actual infinity (Inf) or NA. The second form has proven to be the more useful one.

Presently, the only methods allowing interval censored data are the parametric models computed by survreg and survival curves computed by survfit; for both of these, the distinction between open and closed intervals is unimportant. The distinction is important for counting process data and the Cox model.

The function tries to distinguish between the use of 0/1 and 1/2 coding for censored data via the condition if (max(status)==2). If 1/2 coding is used and all the subjects are censored, it will guess wrong. In any questionable case it is safer to use logical coding, e.g., Surv(time, status==3) would indicate that ‘3’ is the code for an event. For multi-state survival the status variable will be a factor, whose first level is assumed to correspond to censoring.

Surv objects can be subscripted either as a vector, e.g. x[1:3] using a single subscript, in which case the drop argument is ignored and the result will be a survival object; or as a matrix by using two subscripts. If the second subscript is missing and drop=F (the default), the result of the subscripting will be a Surv object, e.g., x[1:3,drop=F], otherwise the result will be a matrix (or vector), in accordance with the default behavior for subscripting matrices.

Value

An object of class Surv. There are methods for print, is.na, and subscripting survival objects. Surv objects are implemented as a matrix of 2 or 3 columns that has further attributes. These include the type (left censored, right censored, counting process, etc.) and labels for the states for multi-state objects. Any attributes of the input arguments are also preserved in inputAttributes. This may be useful for other packages that have attached further information to data items such as labels; none of the routines in the survival package make use of these values, however.

In the case of is.Surv, a logical value TRUE if x inherits from class "Surv", otherwise an FALSE.

Note

The use of 1/2 coding for status is an interesting historical artifact. For data contained on punch cards, IBM 360 Fortran treated blank as a zero, which led to a policy within the Mayo Clinic section
of Biostatistics to never use "0" as a data value since one could not distinguish it from a missing value. Policy became habit, as is often the case, and the use of 1/2 coding for alive/dead endured long after the demise of the punch cards that had sired the practice. At the time Surv was written many Mayo data sets still used this obsolete convention, e.g., the lung data set found in the package.

See Also

coxph, survfit, survreg, lung.

Examples

```r
with(aml, Surv(time, status))
survfit(Surv(time, status) ~ ph.ecog, data=lung)
Surv(heart$start, heart$stop, heart$event)
```

---

Surv-methods

Methods for Surv objects

Description

The list of methods that apply to Surv objects

Usage

```r
## S3 method for class 'Surv'
anyDuplicated(x, ...)
## S3 method for class 'Surv'
as.character(x, ...)
## S3 method for class 'Surv'
as.data.frame(x, ...)
## S3 method for class 'Surv'
as.integer(x, ...)
## S3 method for class 'Surv'
as.matrix(x, ...)
## S3 method for class 'Surv'
as.numeric(x, ...)
## S3 method for class 'Surv'
c(...)
## S3 method for class 'Surv'
duplicated(x, ...)
## S3 method for class 'Surv'
format(x, ...)
## S3 method for class 'Surv'
head(x, ...)
## S3 method for class 'Surv'
is.na(x)
## S3 method for class 'Surv'
length(x)
```
## S3 method for class 'Surv'

### mean(x, ...)

## S3 method for class 'Surv'

### median(x, na.rm=FALSE, ...)

### names(x)

## S3 replacement method for class 'Surv'

names(x) <- value

## S3 method for class 'Surv'

### quantile(x, probs, na.rm=FALSE, ...)

## S3 method for class 'Surv'

### plot(x, ...)

## S3 method for class 'Surv'

### rep(x, ...)

## S3 method for class 'Surv'

### rep.int(x, ...)

## S3 method for class 'Surv'

### rep_len(x, ...)

### rev(x)

## S3 method for class 'Surv'

### t(x)

## S3 method for class 'Surv'

### tail(x, ...)

## S3 method for class 'Surv'

### unique(x, ...)

#### Arguments

- **x**: a Surv object
- **probs**: a vector of probabilities
- **na.rm**: remove missing values from the calculation
- **value**: a character vector of up to the same length as x, or NULL
- **...**: other arguments to the method

#### Details

These functions extend the standard methods to Surv objects. The arguments and results from these are mostly as expected, with the following further details:

- The `as.character` function uses "5+" for right censored at time 5, "5-" for left censored at time 5, "[2,7]" for an observation that was interval censored between 2 and 7, "(1,6]" for a counting process data denoting an observation which was at risk from time 1 to 6, with an event at time 6, and "(1,6+]" for an observation over the same interval but not ending with an event. For a multi-state survival object the type of event is appended to the event time using ":type".
- The print and format methods make use of `as.character`. 
• The `as.numeric` and `as.integer` methods perform these actions on the survival times, but do not affect the censoring indicator.

• The `as.matrix` and `t` methods return a matrix.

• The length of a `Surv` object is the number of survival times it contains, not the number of items required to encode it, e.g., \( x \leftarrow \text{Surv}(1:4,5:9,c(1,0,1,0)) \); \( \text{length}(x) \) has a value of 4. Likewise \( \text{names}(x) \) will be NULL or a vector of length 4. (For technical reasons, any names are actually stored in the \( \text{rownames} \) attribute of the object.)

• For a multi-state survival object `levels` returns the names of the endpoints, otherwise it is NULL.

• The `median`, `quantile` and `plot` methods first construct a survival curve using `survfit`, then apply the appropriate method to that curve.

• The concatenation method `c()` is asymmetric, its first argument determines the execution path. For instance `c(Surv(1:4), Surv(5:6))` will concatenate the two objects, `c(Surv(1:4), 5:6)` will give an error, and `c(5:6, Surv(1:4))` is equivalent to `c(5:6, as.vector(Surv(1:4)))`.

See Also

`Surv`
Details

This function is still experimental.
When used in a coxph or survfit model, Surv2 acts as a trigger to internally convert a timeline style data set into counting process style data, which is then acted on by the routine. The repeated argument controls how repeated instances of the same event code are treated. If TRUE, they are treated as new events, an example where this might be desired is repeated infections in a subject. If FALSE, then repeats are not a new event. An example would be a data set where we wanted to use diabetes, say, as an endpoint, but this is repeated at each medical visit.

Value

An object of class Surv2. There are methods for print, is.na and subscripting.

See Also

Surv2data, coxph, survfit

Surv2data Convert data from timecourse to (time1,time2) style

Description

The multi-state survival functions coxph and survfit allow for two forms of input data. This routine converts between them. The function is normally called behind the scenes when Surv2 is as the response.

Usage

Surv2data(formula, data, subset, id)

Arguments

formula a model formula
data a data frame
subset optional, selects rows of the data to be retained
id a variable that identified multiple rows for the same subject, normally found in the referenced data set

Details

For timeline style data, each row is uniquely identified by an (identifier, time) pair. The time could be a date, time from entry to a study, age, etc, (there may often be more than one time variable). The identifier and time cannot be missing. The remaining covariates represent values that were observed at that time point. Often, a given covariate is observed at only a subset of times and is missing at others. At the time of death, in particular, often only the identifier, time, and status indicator are known.

In the resulting data set missing covariates are replaced by their last known value, and the response y will be a Surv(time1, time2, endpoint) object.
**Value**

a list with elements

*mf* an updated model frame (fewer rows, unchanged columns)

*S2.y* the constructed response variable

*S2.state* the current state for each of the rows

---

**survcheck**

Checks of a survival data set

---

**Description**

Perform a set of consistency checks on survival data

**Usage**

```r
survcheck(formula, data, subset, na.action, id, istate, istate0="(s0)",
timefix=TRUE,...)
```

**Arguments**

- **formula** a model formula with a `Surv` object as the response
- **data** data frame in which to find the id, istate and formula variables
- **subset** expression indicating which subset of the rows of data should be used in the fit. All observations are included by default.
- **na.action** a missing-data filter function. This is applied to the model.frame after any subset argument has been used. Default is `options()$na.action`.
- **id** an identifier that labels unique subjects
- **istate** an optional vector giving the current state at the start of each interval
- **istate0** default label for the initial state of each subject (at their first interval) when istate is missing
- **timefix** process times through the `aeqSurv` function to eliminate potential roundoff issues.
- **...** other arguments, which are ignored (but won’t give an error if someone added weights for instance)

**Details**

This routine will examine a multi-state data set for consistency of the data. The basic rules are that if a subject is at risk they have to be somewhere, can not be two states at once, and should make sensible transitions from state to state. It reports the number of instances of the following conditions:
**overlap** two observations for the same subject that overlap in time, e.g. intervals of (0,100) and (90,120). If \( y \) is simple (time, status) survival observation intervals implicitly start at 0, so in that case any duplicate identifiers will generate an overlap.

**jump** a hole in a subject’s timeline, where they are in one state at the end of the prior interval, but a new state in the at the start subsequent interval.

**gap** one or more gaps in a subject’s timeline; they are presumably in the same state at their return as when they left.

**teleport** two adjacent intervals for a subject, with the first interval ending in one state and the subsequent interval starting in another. They have instantaneously changed states with experiencing a transition.

The total number of occurrences of each is present in the `flags` vector. Optional components give the location and identifiers of the flagged observations.

**Value**

a list with components

- **states** the vector of possible states
- **transitions** a matrix giving the count of transitions from one state to another
- **statecount** table of the number of visits per state, e.g., 18 subjects had 2 visits to the "infection" state
- **flags** a vector giving the counts of each check
- **istate** a copy of the istate vector, if it was supplied; otherwise a constructed istate that satisfies all the checks
- **overlap** a list with the row number and id of overlaps (not present if there are no overlaps)
- **gaps** a list with the row number and id of gaps (not present if there are no gaps)
- **teleport** a list with the row number and id of inconsistent rows (not present if there are none)
- **jumps** a list with the row number and id of jumps (not present if there are no jumps)

**Note**

For data sets with time-dependent covariates, a given subject will often have intermediate rows with a status of ‘no event at this time’. (numeric value of 0). For instance a subject who started in state 1 at time 0, transitioned to state 2 at time 10, had a covariate \( x \) change from 135 to 156 at time 20, and a final transition to state 3 at time 30. The response would be `Surv(c(0,10,20),c(10,20,30),c(2,0,3))`: the status variable records changes in state, and there was no change at time 20. The istate variable would be \((1,2,2)\); it contains the current state, and so the value is unchanged when status = censored.

Thus, when there are intermediate observations `istate` is not simply a lagged version of the status, and may be more challenging to create. One approach is to let `survcheck` do the work: call it with an `istate` argument that is correct for the first row of each subject, or no `istate` argument at all, and then insert the returned value into ones data frame.
Test Survival Curve Differences

Description
Tests if there is a difference between two or more survival curves using the $G^p$ family of tests, or for a single curve against a known alternative.

Usage
survdiff(formula, data, subset, na.action, rho=0, timefix=TRUE)

Arguments

- **formula**: a formula expression as for other survival models, of the form `Surv(time,status) ~ predictors`. For a one-sample test, the predictors must consist of a single `offset(sp)` term, where `sp` is a vector giving the survival probability of each subject. For a k-sample test, each unique combination of predictors defines a subgroup. A `strata` term may be used to produce a stratified test. To cause missing values in the predictors to be treated as a separate group, rather than being omitted, use the `strata` function with its `na.group=TRUE` argument.

- **data**: an optional data frame in which to interpret the variables occurring in the formula.

- **subset**: expression indicating which subset of the rows of data should be used in the fit. This can be a logical vector (which is replicated to have length equal to the number of observations), a numeric vector indicating which observation numbers are to be included (or excluded if negative), or a character vector of row names to be included. All observations are included by default.

- **na.action**: a missing-data filter function. This is applied to the `model.frame` after any subset argument has been used. Default is `options()$na.action`.

- **rho**: a scalar parameter that controls the type of test.

- **timefix**: process times through the `aeqSurv` function to eliminate potential roundoff issues.

Value

A list with components:

- **n**: the number of subjects in each group.
- **obs**: the weighted observed number of events in each group. If there are strata, this will be a matrix with one column per stratum.
- **exp**: the weighted expected number of events in each group. If there are strata, this will be a matrix with one column per stratum.
- **chisq**: the chisquare statistic for a test of equality.
- **var**: the variance matrix of the test.
- **strata**: optionally, the number of subjects contained in each stratum.
Description

This function implements the G-rho family of Harrington and Fleming (1982), with weights on each death of $S(t)^\rho$, where $S(t)$ is the Kaplan-Meier estimate of survival. With $\rho = 0$ this is the log-rank or Mantel-Haenszel test, and with $\rho = 1$ it is equivalent to the Peto & Peto modification of the Gehan-Wilcoxon test.

Peto and Peto show that the Gehan-Wilcoxon test can be badly biased if the two groups have different censoring patterns, and proposed an alternative. Prentice and Marek later showed an actual example where this issue occurs. For most data sets the Gehan-Wilcoxon and Peto-Peto-Prentice variant will hardly differ, however.

If the right hand side of the formula consists only of an offset term, then a one sample test is done. To cause missing values in the predictors to be treated as a separate group, rather than being omitted, use the factor function with its exclude argument to recode the righ-hand-side covariate.

References


Examples

```r
## Two-sample test
survdiff(Surv(futime, fustat) ~ rx, data=ovarian)

## Stratified 7-sample test
survdiff(Surv(time, status) ~ pat.karno + strata(inst), data=lung)

## Expected survival for heart transplant patients based on
## US mortality tables
expect <- survexp(futime ~ 1, data=jasa, cohort=FALSE,
                 rmap= list(age=(accept.dt - birth.dt), sex=1, year=accept.dt),
                 ratetable=survexp.us)

## actual survival is much worse (no surprise)
survdiff(Surv(jasa$futime, jasa$fustat) ~ offset(expect))
```

survexp

Compute Expected Survival

Description

Returns either the expected survival of a cohort of subjects, or the individual expected survival for each subject.
survexp

Usage

survexp(formula, data, weights, subset, na.action, rmap, times,
method=c("ederer", "hakulinen", "conditional", "individual.h",
"individual.s"),
cohoot=TRUE, conditional=FALSE,
ratetable=survival::survexp.us, scale=1,
se.fit, model=FALSE, x=FALSE, y=FALSE)

Arguments

formula formula object. The response variable is a vector of follow-up times and is optional. The predictors consist of optional grouping variables separated by the + operator (as in survfit), and is often ~1, i.e., expected survival for the entire group.
data data frame in which to interpret the variables named in the formula, subset and weights arguments.
weights case weights. This is most useful when conditional survival for a known population is desired, e.g., the data set would contain all unique age/sex combinations and the weights would be the proportion of each.
subset expression indicating a subset of the rows of data to be used in the fit.
na.action function to filter missing data. This is applied to the model frame after subset has been applied. Default is options()$na.action.
rmap an optional list that maps data set names to the ratetable names. See the details section below.
times vector of follow-up times at which the resulting survival curve is evaluated. If absent, the result will be reported for each unique value of the vector of times supplied in the response value of the formula.
method computational method for the creating the survival curves. The individual option does not create a curve, rather it retrieves the predicted survival individual.s or cumulative hazard individual.h for each subject. The default is to use method='ederer' if the formula has no response, and method='hakulinen' otherwise.
cohort logical value. This argument has been superseded by the method argument. To maintain backwards compatibility, if is present and FALSE, it implies method='individual.s'.
conditional logical value. This argument has been superseded by the method argument. To maintain backwards compatibility, if it is present and TRUE it implies method='conditional'.
ratetable a table of event rates, such as survexp.mn, or a fitted Cox model. Note the survival:: prefix in the default argument is present to avoid the (rare) case of a user who expects the default table but just happens to have an object named "survexp.us" in their own directory.
scale numeric value to scale the results. If ratetable is in units/day, scale = 365.25 causes the output to be reported in years.
se.fit compute the standard error of the predicted survival. This argument is currently ignored. Standard errors are not a defined concept for population rate tables (they are treated as coming from a complete census), and for Cox models the
calculation is hard. Despite good intentions standard errors for this latter case have not been coded and validated.

model,x,y flags to control what is returned. If any of these is true, then the model frame, the model matrix, and/or the vector of response times will be returned as components of the final result, with the same names as the flag arguments.

Details

Individual expected survival is usually used in models or testing, to ‘correct’ for the age and sex composition of a group of subjects. For instance, assume that birth date, entry date into the study, sex and actual survival time are all known for a group of subjects. The survexp.us population tables contain expected death rates based on calendar year, sex and age. Then

```r
haz <- survexp(fu.time ~ 1, data=mydata,
               rmap = list(year=entry.dt, age=(birth.dt-entry.dt)),
               method='individual.h'))
```

gives for each subject the total hazard experienced up to their observed death time or last follow-up time (variable fu.time) This probability can be used as a rescaled time value in models:

```r
glm(status ~ 1 + offset(log(haz)), family=poisson)
glm(status ~ x + offset(log(haz)), family=poisson)
```

In the first model, a test for intercept=0 is the one sample log-rank test of whether the observed group of subjects has equivalent survival to the baseline population. The second model tests for an effect of variable x after adjustment for age and sex.

The ratetable being used may have different variable names than the user’s data set, this is dealt with by the `rmap` argument. The rate table for the above calculation was `survexp.us`, a call to `summary(survexp.us)` reveals that it expects to have variables `age` = age in days, `sex`, and `year` = the date of study entry, we create them in the `rmap` line. The sex variable was not mapped, therefore the function assumes that it exists in `mydata` in the correct format. (Note: for factors such as sex, the program will match on any unique abbreviation, ignoring case.)

Cohort survival is used to produce an overall survival curve. This is then added to the Kaplan-Meier plot of the study group for visual comparison between these subjects and the population at large. There are three common methods of computing cohort survival. In the "exact method" of Ederer the cohort is not censored, for this case no response variable is required in the formula. Hakulinen recommends censoring the cohort at the anticipated censoring time of each patient, and Verheul recommends censoring the cohort at the actual observation time of each patient. The last of these is the conditional method. These are obtained by using the respective time values as the follow-up time or response in the formula.

Value

if `cohort=TRUE` an object of class `survexp`, otherwise a vector of per-subject expected survival values. The former contains the number of subjects at risk and the expected survival for the cohort at each requested time. The cohort survival is the hypothetical survival for a cohort of subjects enrolled from the population at large, but matching the data set on the factors found in the rate table.
survexp.fit

References


38:933-942.

Chapter 10.


See Also

survfit, pyears, survexp.us, ratetable, survexp.fit.

Examples

# Stanford heart transplant data
# We don't have sex in the data set, but know it to be nearly all males.
# Estimate of conditional survival
fit1 <- survexp(futime ~ 1, rmap=list(sex="male", year=accept.dt, 
age=(accept.dt-birth.dt)), method='conditional', data=jasa)
survexp(fit1, times=1:1:10*182.5, scale=365) #expected survival by 1/2 years

# Estimate of expected survival stratified by prior surgery
survexp(~ surgery, rmap= list(sex="male", year=accept.dt, 
age=(accept.dt-birth.dt)), method='ederer', data=jasa, 
times=1:10 * 182.5)

## Compare the survival curves for the Mayo PBC data to Cox model fit
##
#fit <- coxph(Surv(time,status>0) ~ trt + log(bili) + log(protime) + age + 
#platelet, data=pbc)
#plot(survfit(Surv(time, status>0) ~ trt, data=pbc), mark.time=FALSE)
#lines(survexp(~ trt, ratetable=fit, data=pbc), col='purple')

survexp.fit

Description

Compute expected survival times.

Usage

survexp.fit(group, x, y, times, death, ratetable)
survexp.object

Arguments

- **group**: if there are multiple survival curves this identifies the group, otherwise it is a constant. Must be an integer.
- **x**: A matrix whose columns match the dimensions of the `ratetable`, in the correct order.
- **y**: the follow up time for each subject.
- **times**: the vector of times at which a result will be computed.
- **death**: a logical value, if TRUE the conditional survival is computed, if FALSE the cohort survival is computed. See `survexp` for more details.
- **ratetable**: a rate table, such as `survexp.uswhite`.

Details

For conditional survival `y` must be the time of last follow-up or death for each subject. For cohort survival it must be the potential censoring time for each subject, ignoring death.

For an exact estimate `times` should be a superset of `y`, so that each subject at risk is at risk for the entire sub-interval of time. For a large data set, however, this can use an inordinate amount of storage and/or compute time. If the `times` spacing is more coarse than this, an actuarial approximation is used which should, however, be extremely accurate as long as all of the returned values are > .99.

For a subgroup of size 1 and `times > y`, the conditional method reduces to \( \exp(-h) \) where \( h \) is the expected cumulative hazard for the subject over his/her observation time. This is used to compute individual expected survival.

Value

A list containing the number of subjects and the expected survival(s) at each time point. If there are multiple groups, these will be matrices with one column per group.

Warning

Most users will call the higher level routine `survexp`. Consequently, this function has very few error checks on its input arguments.

See Also

- `survexp`, `survexp.us`.

---

**survexp.object**

*Expected Survival Curve Object*

Description

This class of objects is returned by the `survexp` class of functions to represent a fitted survival curve.

Objects of this class have methods for summary, and inherit the `print`, `plot`, `points` and `lines` methods from `survfit`. 
Arguments

- **surv**: the estimate of survival at time $t+0$. This may be a vector or a matrix.
- **n.risk**: the number of subjects who contribute at this time.
- **time**: the time points at which the curve has a step.
- **std.err**: the standard error of the cumulative hazard or $-\log(survival)$.
- **strata**: if there are multiple curves, this component gives the number of elements of the `time` etc. vectors corresponding to the first curve, the second curve, and so on. The names of the elements are labels for the curves.
- **method**: the estimation method used. One of "Ederer", "Hakulinen", or "conditional".
- **na.action**: the returned value from the na.action function, if any. It will be used in the printout of the curve, e.g., the number of observations deleted due to missing values.
- **call**: an image of the call that produced the object.

Structure

The following components must be included in a legitimate `survfit` object.

Subscripts

Survexp objects that contain multiple survival curves can be subscripted. This is most often used to plot a subset of the curves.

Details

In expected survival each subject from the data set is matched to a hypothetical person from the parent population, matched on the characteristics of the parent population. The number at risk printed here is the number of those hypothetical subject who are still part of the calculation. In particular, for the Ederer method all hypotheticals are retained for all time, so `n.risk` will be a constant.

See Also

- `plot.survfit`, `summary.survexp`, `print.survfit`, `survexp`.

---

**survfit**

Create survival curves

Description

This function creates survival curves from either a formula (e.g. the Kaplan-Meier), a previously fitted Cox model, or a previously fitted accelerated failure time model.

Usage

`survfit(formula, ...)`
Arguments

formula either a formula or a previously fitted model
... other arguments to the specific method

Details

A survival curve is based on a tabulation of the number at risk and number of events at each unique death time. When time is a floating point number the definition of "unique" is subject to interpretation. The code uses factor() to define the set. For further details see the documentation for the appropriate method, i.e., ?survfit.formula or ?survfit.coxph.

A survfit object may contain a single curve, a set of curves (vector), a matrix of curves, or even a 3 way array: dim(fit) will reveal the dimensions. Predicted curves from a coxph model have one row for each stratum in the Cox model fit and one column for each specified covariate set. Curves from a multi-state model have one row for each stratum and a column for each state, the strata correspond to predictors on the right hand side of the equation. The default printing and plotting order for curves is by column, as with other matrices.

Value

An object of class survfit containing one or more survival curves.

Note

Older releases of the code also allowed the specification for a single curve to omit the right hand of the formula, i.e., survfit(Surv(time,status)), in which case the formula argument is not actually a formula. Handling this case required some non-standard and fairly fragile manipulations, and this case is no longer supported.

Author(s)

Terry Therneau

See Also

survfit.formula, survfit.coxph, survfit.object, print.survfit, plot.survfit, quantile.survfit, residuals.survfit, summary.survfit

survfit.coxph

Compute a Survival Curve from a Cox model

Description

Computes the predicted survivor function for a Cox proportional hazards model.
Usage

```r
## S3 method for class 'coxph'
survfit(formula, newdata,
        se.fit=TRUE, conf.int=.95, individual=FALSE, stype=2, ctype,
        conf.type=c("log", "log-log", "plain", "none", "logit", "arcsin"),
        censor=TRUE, start.time, id, influence=FALSE,
        na.action=na.pass, type, ...)
```

```r
## S3 method for class 'coxphms'
survfit(formula, newdata,
        se.fit=FALSE, conf.int=.95, individual=FALSE, stype=2, ctype,
        conf.type=c("log", "log-log", "plain", "none", "logit", "arcsin"),
        censor=TRUE, start.time, id, influence=FALSE,
        na.action=na.pass, type, p0=NULL, ...)
```

Arguments

- **formula**: A coxph object.
- **newdata**: a data frame with the same variable names as those that appear in the coxph formula. It is also valid to use a vector, if the data frame would consist of a single row.
  The curve(s) produced will be representative of a cohort whose covariates correspond to the values in newdata. Default is the mean of the covariates used in the coxph fit.
- **se.fit**: a logical value indicating whether standard errors should be computed. Default is TRUE for standard models, FALSE for multi-state (code not yet present for that case.)
- **conf.int**: the level for a two-sided confidence interval on the survival curve(s). Default is 0.95.
- **individual**: deprecated argument, replaced by the general id
- **stype**: computation of the survival curve, 1=direct, 2=exponential of the cumulative hazard.
- **ctype**: whether the cumulative hazard computation should have a correction for ties, 1=no, 2=yes.
- **conf.type**: One of "none", "plain", "log" (the default), "log-log" or "logit". Only enough of the string to uniquely identify it is necessary. The first option causes confidence intervals not to be generated. The second causes the standard intervals \( \pm k \times \text{se}(\text{curve}) \), where \( k \) is determined from conf.int. The log option calculates intervals based on the cumulative hazard or log(survival). The log-log option uses the log hazard or log(-log(survival)), and the logit log(survival/(1-survival)).
- **censor**: if FALSE time points at which there are no events (only censoring) are not included in the result.
- **id**: optional variable name of subject identifiers. If this is present, it will be searched for in the newdata data frame. Each group of rows in newdata with the same subject id represents the covariate path through time of a single subject, and the
result will contain one curve per subject. If the coxph fit had strata then that must also be specified in newdata. If newid is not present, then each individual row of newdata is presumed to represent a distinct subject.

**start.time** optional starting time, a single numeric value. If present the returned curve contains survival after start.time conditional on surviving to start.time.

**influence** option to return the influence values

**na.action** the na.action to be used on the newdata argument

**type** older argument that encompassed stype and ctype, now deprecated

**p0** optional, a vector of probabilities. The returned curve will be for a cohort with this mixture of starting states. Most often a single state is chosen

... for future methods

**Details**

This routine produces Pr(state) curves based on a coxph model fit. For single state models it produces the single curve for S(t) = Pr(remain in initial state at time t), known as the survival curve; for multi-state models a matrix giving probabilities for all states. The stype argument states the type of estimate, and defaults to the exponential of the cumulative hazard, better known as the Breslow estimate. For a multi-state Cox model this involves the exponential of a matrix. The argument stype=1 uses a non-exponential or ‘direct’ estimate. For a single endpoint coxph model the code evaluates the Kalbfleich-Prentice estimate, and for a multi-state model it uses an analog of the Aalen-Johansen estimator. The latter approach is the default in the mstate package.

The ctype option affects the estimated cumulative hazard, and if stype=2 the estimated P(state) curves as well. If not present it is chosen so as to be concordant with the ties option in the coxph call. (For multistate coxphms objects only ctype=1 is currently implemented.) Likewise the choice between a model based and robust variance estimate for the curve will mirror the choice made in the coxph call, any clustering is also inherited from the parent model.

If the newdata argument is missing, then a curve is produced for a single "pseudo" subject with covariate values equal to the means of the data set. The resulting curve(s) almost never make sense, but The default remains due to an unwarranted attachment to the option shown by some users and by other packages. Two particularly egregious examples are factor variables and interactions. Suppose one were studying interspecies transmission of a virus, and the data set has a factor variable with levels ("pig", "chicken") and about equal numbers of observations for each. The “mean” covariate level will be 0.5 – is this a flying pig? As to interactions assume data with sex coded as 0/1, ages ranging from 50 to 80, and a model with age*sex. The “mean” value for the age:sex interaction term will be about 30, a value that does not occur in the data. Users are strongly advised to use the newdata argument. For these reasons predictions from a multistate coxph model require the newdata argument.

When the original model contains time-dependent covariates, then the path of that covariate through time needs to be specified in order to obtain a predicted curve. This requires newdata to contain multiple lines for each hypothetical subject which gives the covariate values, time interval, and strata for each line (a subject can change strata), along with an id variable which demarks which rows belong to each subject. The time interval must have the same (start, stop, status) variables as the original model: although the status variable is not used and thus can be set to a dummy value of 0 or 1, it is necessary for the response to be recognized as a Surv object. Last, although predictions
with a time-dependent covariate path can be useful, it is very easy to create a prediction that is senseless. Users are encouraged to seek out a text that discusses the issue in detail.

When a model contains strata but no time-dependent covariates the user of this routine has a choice. If newdata argument does not contain strata variables then the returned object will be a matrix of survival curves with one row for each strata in the model and one column for each row in newdata. (This is the historical behavior of the routine.) If newdata does contain strata variables, then the result will contain one curve per row of newdata, based on the indicated stratum of the original model. In the rare case of a model with strata by covariate interactions the strata variable must be included in newdata, the routine does not allow it to be omitted (predictions become too confusing). (Note that the model Surv(time, status) ~ age*strata(sex) expands internally to strata(sex) + age:sex; the sex variable is needed for the second term of the model.)

See `survfit` for more details about the counts (number of events, number at risk, etc.)

**Value**

an object of class "survfit". See `survfit.object` for details. Methods defined for survfit objects are `print`, `plot`, `lines`, and `points`.

**Notes**

If the following pair of lines is used inside of another function then the `model=TRUE` argument must be added to the coxph call: `fit <- coxph(...); survfit(fit)`. This is a consequence of the non-standard evaluation process used by the `model.frame` function when a formula is involved.

**References**


**See Also**

`print.survfit`, `plot.survfit`, `lines.survfit`, `coxph`, `Surv`, `strata`
survfit.formula

Compute a Survival Curve for Censored Data

Description

Computes an estimate of a survival curve for censored data using the Aalen-Johansen estimator. For ordinary (single event) survival this reduces to the Kaplan-Meier estimate.

Usage

```r
## S3 method for class 'formula'
survfit(formula, data, weights, subset, na.action,
    stype=1, ctype=1, id, cluster, robust, istate, timefix=TRUE,
    etype, error, ...)
```

Arguments

- `formula` a formula object, which must have a `Surv` object as the response on the left of the `~` operator and, if desired, terms separated by `+` operators on the right. One of the terms may be a `strata` object. For a single survival curve the right hand side should be `~ 1`.
- `data` a data frame in which to interpret the variables named in the formula, `subset` and `weights` arguments.
- `weights` The weights must be nonnegative and it is strongly recommended that they be strictly positive, since zero weights are ambiguous, compared to use of the `subset` argument.
- `subset` expression saying that only a subset of the rows of the data should be used in the fit.
- `na.action` a missing-data filter function, applied to the model frame, after any `subset` argument has been used. Default is `options()$na.action`.
- `stype` the method to be used estimation of the survival curve: `1` = direct, `2` = exp(cumulative hazard).
- `ctype` the method to be used for estimation of the cumulative hazard: `1` = Nelson-Aalen formula, `2` = Fleming-Harrington correction for tied events.
- `id` identifies individual subjects, when a given person can have multiple lines of data.
- `cluster` used to group observations for the infinitesimal jackknife variance estimate, defaults to the value of `id`.
- `robust` logical, should the function compute a robust variance. For multi-state survival curves this is true by default. For single state data see details, below.
- `istate` for multi-state models, identifies the initial state of each subject or observation.
- `timefix` process times through the `aeqSurv` function to eliminate potential roundoff issues.


etype  a variable giving the type of event. This has been superseded by multi-state Surv objects and is deprecated; see example below.

error  this argument is no longer used

...  The following additional arguments are passed to internal functions called by survfit.

se.fit  logical value, default is TRUE. If FALSE then standard error computations are omitted.

conf.type  One of "none", "plain", "log" (the default), "log-log", "logit" or "arcsin". Only enough of the string to uniquely identify it is necessary. The first option causes confidence intervals not to be generated. The second causes the standard intervals curve +-k *se(curve), where k is determined from conf.int. The log option calculates intervals based on the cumulative hazard or log(survival). The log-log option bases the intervals on the log hazard or log(-log(survival)), the logit option on log(survival/(1-survival)) and arcsin on arcsin(survival).

conf.lower  a character string to specify modified lower limits to the curve, the upper limit remains unchanged. Possible values are "usual" (unmodified), "peto", and "modified". The modified lower limit is based on an "effective n" argument. The confidence bands will agree with the usual calculation at each death time, but unlike the usual bands the confidence interval becomes wider at each censored observation. The extra width is obtained by multiplying the usual variance by a factor m/n, where n is the number currently at risk and m is the number at risk at the last death time. (The bands thus agree with the un-modified bands at each death time.) This is especially useful for survival curves with a long flat tail.

The Peto lower limit is based on the same "effective n" argument as the modified limit, but also replaces the usual Greenwood variance term with a simple approximation. It is known to be conservative.

start.time  numeric value specifying a time to start calculating survival information. The resulting curve is the survival conditional on surviving to start.time.

conf.int  the level for a two-sided confidence interval on the survival curve(s). Default is 0.95.

se.fit  a logical value indicating whether standard errors should be computed. Default is TRUE.

influence  a logical value indicating whether to return the infinitesimal jackknife (influence) values for each subject. These contain the values of the derivative of each value with respect to the case weights of each subject i: \( \partial p/\partial \omega_i \), evaluated at the vector of weights. The resulting object will contain influence.surv and influence.chaz components. Alternatively, options of influence=1 or influence=2 will return values for only the survival or hazard curves, respectively.

p0  this applies only to multi-state curves. An optional vector giving the initial probability across the states. If this is missing, then p0 is estimated using the frequency of the starting states of all observations at risk at start.time, or if that is not specified, at the time of the first event.
type an older argument that combined stype and ctype, now deprecated. Legal values were "kaplan-meier" which is equivalent to stype=1,ctype=1, "fleming-harrington" which is equivalent to stype=2,ctype=1, and "fh2" which is equivalent to stype=2,ctype=2.

Details

If there is a data argument, then variables in the formula, codeweights, subset, id, cluster and istate arguments will be searched for in that data set.

The routine returns both an estimated probability in state and an estimated cumulative hazard estimate. The cumulative hazard estimate is the Nelson-Aalen (NA) estimate or the Fleming-Harrington (FH) estimate, the latter includes a correction for tied event times. The estimated probability in state can estimated either using the exponential of the cumulative hazard, or as a direct estimate using the Aalen-Johansen approach. For single state data the AJ estimate reduces to the Kaplan-Meier and the probability in state to the survival curve; for competing risks data the AJ reduces to the cumulative incidence (CI) estimator. For backward compatibility the type argument can be used instead.

When the data set includes left censored or interval censored data (or both), then the EM approach of Turnbull is used to compute the overall curve. Currently this algorithm is very slow, only a survival curve is produced, and it does not support a robust variance.

Robust variance: If a robust is TRUE, or for multi-state curves, then the standard errors of the results will be based on an infinitesimal jackknife (IJ) estimate, otherwise the standard model based estimate will be used. For single state curves, the default for robust will be TRUE if one of: there is a cluster argument, there are non-integer weights, or there is a id statement and at least one of the id values has multiple events, and FALSE otherwise. The default represents our best guess about when one would most often desire a robust variance. When there are non-integer case weights and (time1, time2) survival data the routine is at an impasse: a robust variance likely is called for, but requires either id or cluster information to be done correctly; it will default to robust=FALSE.

One unanticipated side effect of defaulting to a robust variance when there are non integer case weights is that a fit using interval censored data, which iterates an underlying KM using an updated weight vector, now returns a robust variance by default. Based on Sun (2001) this may be preferred, as the naive estimate ignores the estimation of the weights. The prior behavior can be obtained with robust= FALSE.

With the IJ estimate, the leverage values themselves can be returned as arrays with dimensions: number of subjects, number of unique times, and for a multi-state model, the number of unique states. Be forwarned that these arrays can be huge. If there is a cluster argument this first dimension will be the number of clusters and the variance will be a grouped IJ estimate; this can be an important tool for reducing the size. A numeric value for the influence argument allows finer control: 0= return neither (same as FALSE), 1= return the influence array for probability in state, 2= return the influence array for the cumulative hazard, 3= return both (same as TRUE).

Value

an object of class "survfit". See survfit.object for details. Methods defined for survfit objects are print, plot, lines, and points.
References


See Also

survfit.coxph for survival curves from Cox models, survfit.object for a description of the components of a survfit object, print.survfit, plot.survfit, lines.survfit, coxph, Surv.

Examples

# fit a Kaplan-Meier and plot it
fit <- survfit(Surv(time, status) ~ x, data = aml)
plot(fit, lty = 2:3)
legend(100, .8, c("Maintained", "Nonmaintained"), lty = 2:3)

# fit a Cox proportional hazards model and plot the predicted survival for a 60 year old
fit <- coxph(Surv(futime, fustat) ~ age, data = ovarian)
plot(survfit(fit, newdata=data.frame(age=60)),
     xscale=365.25, xlab = "Years", ylab="Survival")

# Here is the data set from Turnbull
# There are no interval censored subjects, only left-censored (status=3),
# right-censored (status 0) and observed events (status 1)
#
# Time
# Type of observation
# death 12 6 2 3
# losses 3 2 0 3
# late entry 2 4 2 5
#
tdata <- data.frame(time =c(1,1,2,2,2,3,3,3,4,4,4,4),
                    status=rep(c(1,0,2),4),
                    n =c(12,3,2,6,2,4,2,0,2,3,3,5))
```
fit <- survfit(Surv(time, time, status, type='interval') ~ 1,
data=tdata, weight=n)

# Three curves for patients with monoclonal gammopathy.
# 1. KM of time to PCM, ignoring death (statistically incorrect)
# 2. Competing risk curves (also known as "cumulative incidence")
# 3. Multi-state, showing Pr(in each state, at time t)

fitKM <- survfit(Surv(stop, event=='pcm') ~ 1, data=mgus1,
                 subset=(start==0))
fitCR <- survfit(Surv(stop, event) ~ 1,
data=mgus1, subset=(start==0))
fitMS <- survfit(Surv(start, stop, event) ~ 1, id=id, data=mgus1)

## Not run:
# CR curves show the competing risks
plot(fitCR, xscale=365.25, xmax=7300, mark.time=FALSE,
col=2:3, xlab="Years post diagnosis of MGUS",
ylab="P(state)"
lines(fitKM, fun='event', xmax=7300, mark.time=FALSE,
conf.int=FALSE)
text(3652, .4, "Competing risk: death", col=3)
text(5840, .15,"Competing risk: progression", col=2)
text(5480, .30,"KM:prog")

## End(Not run)
```

---

**survfit.matrix**

Create Aalen-Johansen estimates of multi-state survival from a matrix of hazards.

### Description

This allows one to create the Aalen-Johansen estimate of P, a matrix with one column per state and one row per time, starting with the individual hazard estimates. Each row of P will sum to 1. Note that this routine has been superseded by the use of multi-state Cox models, and will eventually be removed.

### Usage

```
## S3 method for class 'matrix'
survfit(formula, p0, method = c("discrete", "matexp"),
        start.time, ...)
```

### Arguments

- **formula**
  a matrix of lists, each element of which is either NULL or a survival curve object.
### survfit.matrix

- **p0**
  
  the initial state vector. The names of this vector are used as the names of the states in the output object. If there are multiple curves then p0 can be a matrix with one row per curve.

- **method**
  
  use a product of discrete hazards, or a product of matrix exponentials. See details below.

- **start.time**
  
  optional; start the calculations at a given starting point

- **...**
  
  further arguments used by other survfit methods

### Details

On input the matrix should contain a set of predicted curves for each possible transition, and NULL in other positions. Each of the predictions will have been obtained from the relevant Cox model. This approach for multistate curves is easy to use but has some caveats. First, the input curves must be consistent. The routine checks as best it can, but can easily be fooled. For instance, if one were to fit two Cox models, obtain predictions for males and females from one, and for treatment A and B from the other, this routine will create two curves but they are not meaningful. A second issue is that standard errors are not produced.

The names of the resulting states are taken from the names of the vector of initial state probabilities. If they are missing, then the dimnames of the input matrix are used, and lacking that the labels '1', '2', etc. are used.

For the usual Aalen-Johansen estimator the multiplier at each event time is the matrix of hazards H (also written as I + dA). When using predicted survival curves from a Cox model, however, it is possible to get predicted hazards that are greater than 1, which leads to probabilities less than 0. If the `method` argument is not supplied and the input curves are derived from a Cox model this routine instead uses the approximation expm(H-I) as the multiplier, which always gives valid probabilities. (This is also the standard approach for ordinary survival curves from a Cox model.)

### Value

- a survfitms object

### Note

The R syntax for creating a matrix of lists is very fussy.

### Author(s)

Terry Therneau

### See Also

- `survfit`

### Examples

```r
etime <- with(mgus2, ifelse(pstat==0, futime, ptime))
event <- with(mgus2, ifelse(pstat==0, 2*death, 1))
event <- factor(event, 0:2, labels=c("censor", "pcm", "death"))
```
```r
cfit1 <- coxph(Surv(etime, event="pcm") ~ age + sex, mgus2)
cfit2 <- coxph(Surv(etime, event="death") ~ age + sex, mgus2)

# predicted competing risk curves for a 72 year old with mspike of 1.2
# (median values), male and female.
# The survfit call is a bit faster without standard errors.
newdata <- expand.grid(sex=c("F", "M"), age=72, mspike=1.2)
AJmat <- matrix(list(), 3,3)
AJmat[1,2] <- list(survfit(cfit1, newdata, std.err=FALSE))
AJmat[1,3] <- list(survfit(cfit2, newdata, std.err=FALSE))
csurv <- survfit(AJmat, p0 =c(entry=1, PCM=0, death=0))
```

---

### survfit.object

**Survival Curve Object**

**Description**

This class of objects is returned by the `survfit` class of functions to represent a fitted survival curve. For a multi-state model the object has class `c('survfitms', 'survfit')`.

Objects of this class have methods for the functions `print`, `summary`, `plot`, `points` and `lines`. The `print.survfit` method does more computation than is typical for a print method and is documented on a separate page.

**Arguments**

- `n` total number of subjects in each curve.
- `time` the time points at which the curve has a step.
- `n.risk` the number of subjects at risk at t.
- `n.event` the number of events that occur at time t.
- `n.enter` for counting process data only, the number of subjects that enter at time t.
- `n.censor` for counting process data only, the number of subjects who exit the risk set, without an event, at time t. (For right censored data, this number can be computed from the successive values of the number at risk).
- `surv` the estimate of survival at time t+0. This may be a vector or a matrix. The latter occurs when a set of survival curves is created from a single Cox model, in which case there is one column for each covariate set.
- `pstate` a multi-state survival will have the `pstate` component instead of `surv`. It will be a matrix containing the estimated probability of each state at each time, one column per state.
- `std.err` for a survival curve this contains standard error of the cumulative hazard or `-log(survival)`, for a multi-state curve it contains the standard error of prev. This difference is a reflection of the fact that each is the natural calculation for that case.
- `cumhaz` hazard optional. Contains the cumulative hazard for each possible transition.
survfit.object

- **strata**: if there are multiple curves, this component gives the number of elements of the time vector corresponding to the first curve, the second curve, and so on. The names of the elements are labels for the curves.
- **upper**: optional upper confidence limit for the survival curve or pstate
- **lower**: options lower confidence limit for the survival curve or pstate
- **start.time**: optional, the starting time for the curve if other than 0
- **p0, sp0**: for a multistate object, the distribution of starting states. If the curve has a strata dimension, this will be a matrix one row per stratum. The sp0 element has the standard error of p0, if p0 was estimated.
- **newdata**: for survival curves from a fitted model, this contains the covariate values for the curves
- **n.all**: the total number of observations that were available. For counting process data, and any time that the start.time argument was used, not all may have been used in creating the curve, in which case this value will be larger than n above. The print and plot routines in the package do no use this value, it is for information only.
- **conf.type**: the approximation used to compute the confidence limits.
- **conf.int**: the level of the confidence limits, e.g., 90 or 95%.
- **transitions**: for multi-state data, the total number of transitions of each type.
- **na.action**: the returned value from the na.action function, if any. It will be used in the printout of the curve, e.g., the number of observations deleted due to missing values.
- **call**: an image of the call that produced the object.
- **type**: type of survival censoring.
- **influence.p, influence.c**: optional influence matrices for the pstate (or surv) and for the cumhaz estimates. A list with one element per stratum, each element of the list is an array indexed by subject, time, state.
- **version**: the version of the object. Will be missing, 2, or 3

**Structure**

The following components must be included in a legitimate survfit or survfits object.

**Subscripts**

Survfit objects can be subscripted. This is often used to plot a subset of the curves, for instance. From the user’s point of view the survfit object appears to be a vector, matrix, or array of curves. The first dimension is always the underlying number of curves or “strata”; for multi-state models the state is always the last dimension. Predicted curves from a Cox model can have a second dimension which is the number of different covariate prediction vectors.
Details

The `survfit` object has evolved over time: when first created there was no thought of multi-state models for instance. This evolution has almost entirely been accomplished by the addition of new elements. One change in survival version 3 is the addition of a `survfitconf` routine which will compute confidence intervals for a `survfit` object. This allows the computation of CI intervals to be deferred until later, if desired, rather than making them a permanent part of the object. Later iterations of the base routines may omit the confidence intervals.

The `survfit` object starts at the first observation time, but survival curves are normally plotted from time 0. A helper routine `survfit0` can be used to add this first time point and align the data.

See Also

`plot.survfit`, `summary.survfit`, `print.survfit`, `survfit`, `survfit0`

```r
survfit0

Convert the format of a survfit object.
```

Description

Add the point for a starting time (time 0) to a survfit object's elements. This is useful for plotting.

Usage

```
survfit0(x, start.time=0)
```

Arguments

- `x`: a survfit object
- `start.time`: the desired starting time; see details below.

Details

Survival curves are traditionally plotted forward from time 0, but since the true starting time is not known as a part of the data, the `survfit` routine does not include a time 0 value in the resulting object. Someone might look at cumulative mortgage defaults versus calendar year, for instance, with the ‘time’ value a Date object. The plotted curve probably should not start at 0 = 1970/01/01. Due to this uncertainty, it was decided not to include a "time 0" as part of a survfit object. If the original `survfit` call included a `start.time` argument, that value is of course retained.

Whether that (1989) decision was wise or foolish, it is now far too late to change it. (We tried it once as a trial, resulting in over 20 errors in the survival test suite. We extrapolate that it might break 1/2 - 2/3 of the other CRAN packages that depend on survival, if made a default.) If the original `survfit` call included a `start.time` argument, that value is of course retained.

One problem with this choice is that some functions must choose a starting point, plots for example. This utility function is used by `plot.survfit` and `summary.survfit` to do so, adding a new time point at the front of each curve in a consistent way: the optional argument to the `survfit0` function as the first choice (if supplied), then the user's `start.time` if present, otherwise `min(0,x$time)`.
The resulting object is not guaranteed to work with functions that further manipulate a `survfit` object such as subscripting, aggregation, pseudovalues, etc. (remember the 20 errors). Rather it is intended as a penultimate step, most often when creating a plot.

**Value**

a reformulated version of the object with an initial data point at `start.time` added. The time, `surv`, `pstate`, `cumhaz`, `std.err`, `std.cumhaz` and other components will all be aligned, so as to make plots and summaries easier to produce.

---

**Description**

This program is mainly supplied to allow other packages to invoke the `survfit.coxph` function at a ‘data’ level rather than a ‘user’ level. It does no checks on the input data that is provided, which can lead to unexpected errors if that data is wrong.

**Usage**

```r
survfitcoxph.fit(y, x, wt, x2, risk, newrisk, strata, se.fit, survtype, vartype, varmat, id, y2, strata2, unlist=TRUE)
```

**Arguments**

- `y`: the response variable used in the Cox model. (Missing values removed of course.)
- `x`: covariate matrix used in the Cox model
- `wt`: weight vector for the Cox model. If the model was unweighted use a vector of 1s.
- `x2`: matrix describing the hypothetical subjects for which a curve is desired. Must have the same number of columns as `x`.
- `risk`: the risk score \( \exp(X \beta) \) from the fitted Cox model. If the model had an offset, include it in the argument to `exp`.
- `newrisk`: risk scores for the hypothetical subjects
- `strata`: strata variable used in the Cox model. This will be a factor.
- `se.fit`: if TRUE the standard errors of the curve(s) are returned
- `survtype`: 1=Kalbfleisch-Prentice, 2=Nelson-Aalen, 3=Efron. It is usual to match this to the approximation for ties used in the coxph model: KP for ‘exact’, N-A for ‘breslow’ and Efron for ‘efron’.
- `vartype`: 1=Greenwood, 2=Aalen, 3=Efron
- `varmat`: the variance matrix of the coefficients
id optional; if present and not NULL this should be a vector of identifiers of length nrow(x2). A mon-null value signifies that x2 contains time dependent covariates, in which case this identifies which rows of x2 go with each subject.

y2 survival times, for time dependent prediction. It gives the time range (time1,time2] for each row of x2. Note: this must be a Surv object and thus contains a status indicator, which is never used in the routine, however.

strata2 vector of strata indicators for x2. This must be a factor.

unlist if FALSE the result will be a list with one element for each strata. Otherwise the strata are “unpacked” into the form found in a survfit object.

Value

a list containing nearly all the components of a survfit object. All that is missing is to add the confidence intervals, the type of the original model’s response (as in a coxph object), and the class.

Note

The source code for both this function and survfit.coxph is written using noweb. For complete documentation see the inst/sourcecode.pdf file.

Author(s)

Terry Therneau

See Also

survfit.coxph

survConcordance(formula, data, weights, subset, na.action) # use concordance
survConcordance.fit(y, x, strata, weight) # use concordancefit
Argument

**formula**
- 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.

**data**
- a data frame

**weights, subset, na.action**
- as for `coxph`

**x, y, strata, weight**
- predictor, response, strata, and weight vectors for the direct call

See Also

**Deprecated**

**survobrien**

O’Brien’s Test for Association of a Single Variable with Survival

Description

Peter O’Brien’s test for association of a single variable with survival This test is proposed in Biometrics, June 1978.

Usage

```
survobrien(formula, data, subset, na.action, transform)
```

Arguments

**formula**
- a valid formula for a cox model.

**data**
- a data.frame in which to interpret the variables named in the formula, or in the subset and the weights argument.

**subset**
- expression indicating which subset of the rows of data should be used in the fit. All observations are included by default.

**na.action**
- a missing-data filter function. This is applied to the model.frame after any subset argument has been used. Default is `options()$na.action`.

**transform**
- the transformation function to be applied at each time point. The default is O’Brien’s suggestion `logit(tr)` where `tr = (rank(x)- 1/2)/ length(x)` is the rank shifted to the range 0-1 and `logit(x) = log(x/(1-x))` is the logit transform.

Value

a new data frame. The response variables will be column names returned by the `Surv` function, i.e., "time" and "status" for simple survival data, or "start", "stop", "status" for counting process data. Each individual event time is identified by the value of the variable .strata.. Other variables retain their original names. If a predictor variable is a factor or is protected with I (), it is retained as is. Other predictor variables have been replaced with time-dependent logit scores.

The new data frame will have many more rows that the original data, approximately the original number of rows * number of deaths/2.
Method

A time-dependent cox model can now be fit to the new data. The univariate statistic, as originally proposed, is equivalent to single variable score tests from the time-dependent model. This equivalence is the rationale for using the time dependent model as a multivariate extension of the original paper.

In O’Brien’s method, the x variables are re-ranked at each death time. A simpler method, proposed by Prentice, ranks the data only once at the start. The results are usually similar.

Note

A prior version of the routine returned new time variables rather than a strata. Unfortunately, that strategy does not work if the original formula has a strata statement. This new data set will be the same size, but the coxph routine will process it slightly faster.

References


See Also

survdiff

Examples

```r
xx <- survobrien(Surv(futime, fustat) ~ age + factor(rx) + I(ecog.ps),
                  data=ovarian)
coxph(Surv(time, status) ~ age + strata(.strata.), data=xx)
```

survreg

Regression for a Parametric Survival Model

Description

Fit a parametric survival regression model. These are location-scale models for an arbitrary transform of the time variable; the most common cases use a log transformation, leading to accelerated failure time models.

Usage

```r
survreg(formula, data, weights, subset,
         na.action, dist="weibull", init=NULL, scale=0,
         control, parms=NULL, model=FALSE, x=FALSE,
         y=TRUE, robust=FALSE, cluster, score=FALSE, ...)
```
Arguments

formula a formula expression as for other regression models. The response is usually a survival object as returned by the Surv function. See the documentation for Surv, lm and formula for details.
data a data frame in which to interpret the variables named in the formula, weights or the subset arguments.weights optional vector of case weights.subset subset of the observations to be used in the fit.na.action a missing-data filter function, applied to the model.frame, after any subset argument has been used. Default is options()$na.action.dist assumed distribution for y variable. If the argument is a character string, then it is assumed to name an element from survreg.distributions. These include "weibull", "exponential", "gaussian", "logistic", "lognormal" and "loglogistic". Otherwise, it is assumed to be a user defined list conforming to the format described in survreg.distributions.parms a list of fixed parameters. For the t-distribution for instance this is the degrees of freedom; most of the distributions have no parameters.init optional vector of initial values for the parameters.scale optional fixed value for the scale. If set to <=0 then the scale is estimated.control a list of control values, in the format produced by survreg.control. The default value is survreg.control().model,x,y flags to control what is returned. If any of these is true, then the model frame, the model matrix, and/or the vector of response times will be returned as components of the final result, with the same names as the flag arguments.score return the score vector. (This is expected to be zero upon successful convergence.)robust Use robust sandwich error instead of the asymptotic formula. Defaults to TRUE if there is a cluster argument.cluster Optional variable that identifies groups of subjects, used in computing the robust variance. Like model variables, this is searched for in the dataset pointed to by the data argument.... other arguments which will be passed to survreg.control.

Details

All the distributions are cast into a location-scale framework, based on chapter 2.2 of Kalbfleisch and Prentice. The resulting parameterization of the distributions is sometimes (e.g. gaussian) identical to the usual form found in statistics textbooks, but other times (e.g. Weibull) it is not. See the book for detailed formulas.

When using weights be aware of the difference between replication weights and sampling weights. In the former, a weight of ‘2’ means that there are two identical observations, which have been combined into a single row of data. With sampling weights there is a single observed value, with a weight used to achieve balance with respect to some population. To get proper variance with replication weights use the default variance, for sampling weights use the robust variance. Replication weights were once common (when computer memory was much smaller) but are now rare.
survreg.control

Package options for survreg and coxph

Value

an object of class survreg is returned.

References


See Also

survreg.object, survreg.distributions, pspline, frailty, ridge

Examples

# Fit an exponential model: the two fits are the same
survreg(Surv(futime, fustat) ~ ecog.ps + rx, ovarian, dist='weibull', scale=1)
survreg(Surv(futime, fustat) ~ ecog.ps + rx, ovarian, dist="exponential")

# A model with different baseline survival shapes for two groups, i.e.,
# two different scale parameters
survreg(Surv(time, status) ~ ph.ecog + age + strata(sex), lung)

# There are multiple ways to parameterize a Weibull distribution. The survreg
# function embeds it in a general location-scale family, which is a
# different parameterization than the rweibull function, and often leads
# to confusion.
# survreg's scale = 1/(rweibull shape)
# survreg's intercept = log(rweibull scale)
# For the log-likelihood all parameterizations lead to the same value.
y <- rweibull(1000, shape=2, scale=5)
survreg(Surv(y)~1, dist="weibull")

# Economists fit a model called 'tobit regression', which is a standard
# linear regression with Gaussian errors, and left censored data.
tobinf <- survreg(Surv(durable, durable>0, type='left') ~ age + quant,
  data=tobin, dist='gaussian')
Usage

survreg.control(maxiter=30, rel.tolerance=1e-09,
toler.chol=1e-10, iter.max, debug=0, outer.max=10)

Arguments

maxiter maximum number of iterations
rel.tolerance relative tolerance to declare convergence
toler.chol Tolerance to declare Cholesky decomposition singular
iter.max same as maxiter
debug print debugging information
outer.max maximum number of outer iterations for choosing penalty parameters

Value

A list with the same elements as the input

See Also

survreg

survreg.distributions  Parametric Survival Distributions

Description

List of distributions for accelerated failure models. These are location-scale families for some transformation of time. The entry describes the cdf $F$ and density $f$ of a canonical member of the family.

Usage

survreg.distributions

Format

There are two basic formats, the first defines a distribution de novo, the second defines a new distribution in terms of an old one.

name: name of distribution
variance: function(parms) returning the variance (currently unused)
init(x,weights,...): Function returning an initial estimate of the mean and variance (used for initial values in the iteration)
density(x,parms): Function returning a matrix with columns $F$, $1 - F$, $f$, $f'/f$, and $f''/f$
quantile(p,parms): Quantile function
scale: Optional fixed value for the scale parameter
parms: Vector of default values and names for any additional parameters
deviance(y,scale,parms): Function returning the deviance for a
saturated model; used only for deviance residuals.

and to define one distribution in terms of another

name: name of distribution
dist: name of parent distribution
trans: transformation (e.g., log)
dtrans: derivative of transformation
itrans: inverse of transformation
scale: Optional fixed value for scale parameter

Details

There are four basic distributions: extreme, gaussian, logistic, and t. The last three are parametrised in the same way as the distributions already present in R. The extreme value cdf is

\[ F = 1 - e^{-e^t}. \]

When the logarithm of survival time has one of the first three distributions we obtain respectively weibull, lognormal, and loglogistic. The location-scale parameterization of a Weibull distribution found in survreg is not the same as the parameterization of rweibull.

The other predefined distributions are defined in terms of these. The exponential and rayleigh distributions are Weibull distributions with fixed scale of 1 and 0.5, respectively, and loggaussian is a synonym for lognormal.

For speed, parts of the three most commonly used distributions are hardcoded in C; for this reason the elements of survreg.distributions with names of "Extreme value", "Logistic", and "Gaussian" should not be modified. (The order of these in the list is not important, recognition is by name.) As an alternative to modifying survreg.distributions a new distribution can be specified as a separate list. This is the preferred method of addition and is illustrated below.

See Also

survreg, pweibull, pnorm, plogis, pt, survregDtest

Examples

# time transformation
survreg(Surv(time, status) ~ ph.ecog + sex, dist='weibull', data=lung)
# change the transformation to work in years
# intercept changes by log(365), everything else stays the same
my.weibull <- survreg.distributions$weibull
my.weibull$trans <- function(y) log(y/365)
my.weibull$itrans <- function(y) 365*exp(y)
survreg(Surv(time, status) ~ ph.ecog + sex, lung, dist=my.weibull)
# Weibull parametrization
y <- rweibull(1000, shape=2, scale=5)
survreg(Surv(y)-1, dist="weibull")
# survreg scale parameter maps to 1/shape, linear predictor to log(scale)

# Cauchy fit
mycauchy <- list(name = 'Cauchy',
  init = function(x, weights, ...)
    c(median(x), mad(x)),
  density = function(x, parms) {
    temp <- 1/(1 + x^2)
    cbind(.5 + atan(x)/pi, .5+ atan(-x)/pi,
      temp/pi, -2 *x*temp, 2*temp*(4*x^2*temp -1))
  },
  quantile = function(p, parms) tan((p-.5)*pi),
  deviance = function(...) stop('deviance residuals not defined')
)
survreg(Surv(log(time), status) ~ ph.ecog + sex, lung, dist=mycauchy)

---

survreg.object  Parametric Survival Model Object

Description

This class of objects is returned by the `survreg` function to represent a fitted parametric survival model. Objects of this class have methods for the functions `print`, `summary`, `predict`, and `residuals`.

COMPONENTS

The following components must be included in a legitimate `survreg` object.

- **coefficients**: the coefficients of the `linear.predictors`, which multiply the columns of the model matrix. It does not include the estimate of error (sigma). The names of the coefficients are the names of the single-degree-of-freedom effects (the columns of the model matrix). If the model is over-determined there will be missing values in the coefficients corresponding to non-estimable coefficients.

- **icoef**: coefficients of the baseline model, which will contain the intercept and log(scale), or multiple scale factors for a stratified model.

- **var**: the variance-covariance matrix for the parameters, including the log(scale) parameter(s).

- **loglik**: a vector of length 2, containing the log-likelihood for the baseline and full models.

- **iter**: the number of iterations required

- **linear.predictors**: the linear predictor for each subject.

- **df**: the degrees of freedom for the final model. For a penalized model this will be a vector with one element per term.

- **scale**: the scale factor(s), with length equal to the number of strata.
**survregDtest**

```
  degrees of freedom for the initial model.
  a vector of the column means of the coefficient matrix.
  the distribution used in the fit.
  included for a weighted fit.

The object will also have the following components found in other model results (some are optional): linear predictors, weights, x, y, model, call, terms and formula. See `lm`.

**See Also**

`survreg`, `lm`

---

**survregDtest**  Verify a survreg distribution

**Description**

This routine is called by `survreg` to verify that a distribution object is valid.

**Usage**

```
survregDtest(dlist, verbose = F)
```

**Arguments**

- `dlist` the list describing a survival distribution
- `verbose` return a simple TRUE/FALSE from the test for validity (the default), or a verbose description of any flaws.

**Details**

If the `survreg` function rejects your user-supplied distribution as invalid, this routine will tell you why it did so.

**Value**

TRUE if the distribution object passes the tests, and either FALSE or a vector of character strings if not.

**Author(s)**

Terry Therneau

**See Also**

`survreg.distributions`, `survreg`
Examples

```r
# An invalid distribution (it should have "init =" on line 2)
# surveg would give an error message
mycauchy <- list(name="Cauchy",
                   init<- function(x, weights, ...)
                   c(median(x), mad(x)),
                   density= function(x, parms) {
                       temp <- 1/(1 + x^2)
                       cbind(.5 + atan(temp)/pi, .5+ atan(-temp)/pi,
                             temp/pi, -2 *x*temp, 2*temp*2*(4*x^2*temp -1))
                   },
                   quantile= function(p, parms) tan((p-.5)*pi),
                   deviance= function(...) stop('deviance residuals not defined')
)

survregDtest(mycauchy, TRUE)
```

**survSplit**

*Split a survival data set at specified times*

**Description**

Given a survival data set and a set of specified cut times, split each record into multiple subrecords at each cut time. The new data set will be in `counting process` format, with a start time, stop time, and event status for each record.

**Usage**

```r
survSplit(formula, data, subset, na.action=na.pass,
           cut, start="tstart", id, zero=0, episode,
           end="tstop", event="event")
```

**Arguments**

- `formula`: a model formula
- `data`: a data frame
- `subset, na.action`: rows of the data to be retained
- `cut`: the vector of timepoints to cut at
- `start`: character string with the name of a start time variable (will be created if needed)
- `id`: character string with the name of new id variable to create (optional). This can be useful if the data set does not already contain an identifier.
- `zero`: If `start` doesn’t already exist, this is the time that the original records start.
- `episode`: character string with the name of new episode variable (optional)
- `end`: character string with the name of event time variable
- `event`: character string with the name of censoring indicator
Details

Each interval in the original data is cut at the given points; if an original row were (15, 60] with a
cut vector of (10,30, 40) the resulting data set would have intervals of (15,30], (30,40] and (40, 60].

Each row in the final data set will lie completely within one of the cut intervals. Which interval for
each row of the output is shown by the episode variable, where 1= less than the first cutpoint, 2= between the first and the second, etc. For the example above the values would be 2, 3, and 4.

The routine is called with a formula as the first argument. The right hand side of the formula can
be used to delimit variables that should be retained; normally one will use ~ . as a shorthand to
retain them all. The routine will try to retain variable names, e.g. Surv(adam, joe, fred)~. will
result in a data set with those same variable names for tstart, end, and event options rather than
the defaults. Any user specified values for these options will be used if they are present, of course.
However, the routine is not sophisticated; it only does this substitution for simple names. A call of
Surv(time, stat==2) for instance will not retain "stat" as the name of the event variable.

Rows of data with a missing time or status are copied across unchanged, unless the na.action argu-
ment is changed from its default value of na.pass. But in the latter case any row that is missing for
any variable will be removed, which is rarely what is desired.

Value

New, longer, data frame.

See Also

Surv, cut, reshape

Examples

```r
fit1 <- coxph(Surv(time, status) ~ karno + age + trt, veteran)
plot(cox.zph(fit1)[1])
# a cox.zph plot of the data suggests that the effect of Karnofsky score
# begins to diminish by 60 days and has faded away by 120 days.
# Fit a model with separate coefficients for the three intervals.
#
vet2 <- survSplit(Surv(time, status) ~., veteran,
cut=c(60, 120), episode ="timegroup")
fit2 <- coxph(Surv(tstart, time, status) ~ karno* strata(timegroup) +
age + trt, data= vet2)
c(overall= coef(fit1)[1],
t0_60 = coef(fit2)[1],
t60_120= sum(coef(fit2)[c(1,4)]),
t120 = sum(coef(fit2)[c(1,5)]))
```
tcut

Factors for person-year calculations

Description

Attaches categories for person-year calculations to a variable without losing the underlying continuous representation.

Usage

tcut(x, breaks, labels, scale=1)
## S3 method for class 'tcut'

levels(x)

Arguments

- **x**: numeric/date variable
- **breaks**: breaks between categories, which are right-continuous
- **labels**: labels for categories
- **scale**: Multiply x and breaks by this.

Value

An object of class tcut

See Also

cut, pyyears

Examples

```r
mdy.date <- function(m,d,y)
  as.Date(paste(ifelse(y<100, y+1900, y), m, d, sep='/'))
temp1 <- mdy.date(6,6,36)
temp2 <- mdy.date(6,6,55)# Now compare the results from person-years
# temp.age <- tcut(temp2-temp1, floor(c(-1, (18:31 * 365.24))),
labels=c('0-18', paste(18:30, 19:31, sep='-', sep='')))
temp.yr <- tcut(temp2, mdy.date(1,1,1954:1965), labels=1954:1964)
temp.time <- 3700 #total days of fu
py1 <- pyyears(temp.time ~ temp.age + temp.yr, scale=1) #output in days
py1
```
**tmerge**

*Time based merge for survival data*

**Description**

A common task in survival analysis is the creation of start, stop data sets which have multiple intervals for each subject, along with the covariate values that apply over that interval. This function aids in the creation of such data sets.

**Usage**

`tmerge(data1, data2, id,..., tstart, tstop, options)`

**Arguments**

- `data1`: the primary data set, to which new variables and/or observation will be added
- `data2`: second data set in which all the other arguments will be found
- `id`: subject identifier
- `...`: operations that add new variables or intervals, see below
- `tstart`: optional variable to define the valid time range for each subject, only used on an initial call
- `tstop`: optional variable to define the valid time range for each subject, only used on an initial call
- `options`: a list of options. Valid ones are idname, tstartname, tstopname, delay, na.rm, and tdcstart. See the explanation below.

**Details**

The program is often run in multiple passes, the first of which defines the basic structure, and subsequent ones that add new variables to that structure. For a more complete explanation of how this routine works refer to the vignette on time-dependent variables.

There are 4 types of operational arguments: a time dependent covariate (tdc), cumulative count (cumtdc), event (event) or cumulative event (cumevent). Time dependent covariates change their values before an event, events are outcomes.

- `newname = tdc(y, x, init)` A new time dependent covariate variable will created. The argument `y` is assumed to be on the scale of the start and end time, and each instance describes the occurrence of a "condition" at that time. The second argument `x` is optional. In the case where `x` is missing the count variable starts at 0 for each subject and becomes 1 at the time of the event. If `x` is present the value of the time dependent covariate is initialized to value of `init`, if present, or the tdcstart option otherwise, and is updated to the value of `x` at each observation. If the option `na.rm=TRUE` missing values of `x` are first removed, i.e., the update will not create missing values.

- `newname = cumtdc(y,x, init)` Similar to tdc, except that the event count is accumulated over time for each subject. The variable `x` must be numeric.
newname = event(y, x) Mark an event at time y. In the usual case that x is missing the new 0/1 variable will be similar to the 0/1 status variable of a survival time.

newname = cumevent(y, x) Cumulative events.

The function adds three new variables to the output data set: tstart, tstop, and id. The options argument can be used to change these names. If, in the first call, the id argument is a simple name, that variable name will be used as the default for the idname option. If data1 contains the tstart variable then that is used as the starting point for the created time intervals, otherwise the initial interval for each id will begin at 0 by default. This will lead to an invalid interval and subsequent error if say a death time were <= 0.

The na.rm option affects creation of time-dependent covariates. Should a data row in data2 that has a missing value for the variable be ignored or should it generate an observation with a value of NA? The default of TRUE causes the last non-missing value to be carried forward. The delay option causes a time-dependent covariate’s new value to be delayed, see the vignette for an example.

Value

a data frame with two extra attributes tm.retain and tcount. The first contains the names of the key variables, and which names correspond to tdc or event variables. The tcount variable contains counts of the match types. New time values that occur before the first interval for a subject are "early", those after the last interval for a subject are "late", and those that fall into a gap are of type "gap". All these are are considered to be outside the specified time frame for the given subject. An event of this type will be discarded. An observation in data2 whose identifier matches no rows in data1 is of type "missid" and is also discarded. A time-dependent covariate value will be applied to later intervals but will not generate a new time point in the output.

The most common type will usually be "within", corresponding to those new times that fall inside an existing interval and cause it to be split into two. Observations that fall exactly on the edge of an interval but within the (min, max] time for a subject are counted as being on a "leading" edge, "trailing" edge or "boundary". The first corresponds for instance to an occurrence at 17 for someone with an intervals of (0,15] and (17, 35]. A tdc at time 17 will affect this interval but an event at 17 would be ignored. An event occurrence at 15 would count in the (0,15] interval. The last case is where the main data set has touching intervals for a subject, e.g. (17, 28] and (28,35] and a new occurrence lands at the join. Events will go to the earlier interval and counts to the latter one. A last column shows the number of additions where the id and time point were identical. When this occurs, the tdc and event operators will use the final value in the data (last edit wins), but ignoring missing, while cumtdc and cumevent operators add up the values.

These extra attributes are ephemeral and will be discarded if the dataframe is modified. This is intentional, since they will become invalid if for instance a subset were selected.

Author(s)

Terry Therneau

See Also

neardate
Examples

# The pbc data set contains baseline data and follow-up status
# for a set of subjects with primary biliary cirrhosis, while the
# pbcseq data set contains repeated laboratory values for those
# subjects.
# The first data set contains data on 312 subjects in a clinical trial plus
# 106 that agreed to be followed off protocol, the second data set has data
# only on the trial subjects.
temp <- subset(pbc, id <= 312, select=c(id:sex, stage)) # baseline data
pbc2 <- tmerge(temp, temp, id=id, endpt = event(time, status))
pbc2 <- tmerge(pbc2, pbcseq, id=id, ascites = tdc(day, ascites),
            bili = tdc(day, bili), albumin = tdc(day, albumin),
            protime = tdc(day, protime), alk.phos = tdc(day, alk.phos))
fit <- coxph(Surv(tstart, tstop, endpt==2) ~ protime + log(bili), data=pbc2)

---

tobin

Tobin’s Tobit data

Description

Economists fit a parametric censored data model called the ‘tobit’. These data are from Tobin’s
original paper.

Usage

tobin
data(tobin, package="survival")

Format

A data frame with 20 observations on the following 3 variables.

- **durable**  Durable goods purchase
- **age**    Age in years
- **quant** Liquidty ratio (x 1000)

Source


Examples

tfit <- survreg(Surv(durable, durable>0, type='left') ~ age + quant,
               data=tobin, dist='gaussian')
predict(tfit, type="response")
Description
Subjects on a liver transplant waiting list from 1990-1999, and their disposition: received a transplant, died while waiting, withdrew from the list, or censored.

Usage
transplant
data(transplant, package="survival")

Format
A data frame with 815 (transplant) observations on the following 6 variables.

- age  age at addition to the waiting list
- sex  m or f
- abo  blood type: A, B, AB or 0
- year  year in which they entered the waiting list
- futime  time from entry to final disposition
- event  final disposition: censored, death, ltx or withdraw

Details
This represents the transplant experience in a particular region, over a time period in which liver transplant became much more widely recognized as a viable treatment modality. The number of liver transplants rises over the period, but the number of subjects added to the liver transplant waiting list grew much faster. Important questions addressed by the data are the change in waiting time, who waits, and whether there was an consequent increase in deaths while on the list.

Blood type is an important consideration. Donor livers from subjects with blood type O can be used by patients with A, B, AB or 0 blood types, whereas an AB liver can only be used by an AB recipient. Thus type O subjects on the waiting list are at a disadvantage, since the pool of competitors is larger for type O donor livers.

This data is of historical interest and provides a useful example of competing risks, but it has little relevance to current practice. Liver allocation policies have evolved and now depend directly on each individual patient’s risk and need, assessments of which are regularly updated while a patient is on the waiting list. The overall organ shortage remains acute, however.

The transplant data set was a version used early in the analysis, transplant2 has several additions and corrections, and was the final data set and matches the paper.

References
Examples

```r
# since event is a factor, survfit creates competing risk curves
pfit <- survfit(Surv(futime, event) ~ abo, transplant)
pfit[,2] # time to liver transplant, by blood type
plot(pfit[,2], mark.time=FALSE, col=1:4, lwd=2, xmax=735,
     xscale=30.5, xlab="Months", ylab="Fraction transplanted",
     xaxt = 'n')
temp <- c(0, 6, 12, 18, 24)
axis(1, temp*30.5, temp)
legend(450, .35, levels(transplant$abo), lty=1, col=1:4, lwd=2)

# competing risks for type O
plot(pfit[4,], xscale=30.5, xmax=735, col=1:3, lwd=2)
legend(450, .4, c("Death", "Transplant", "Withdrawal"), col=1:3, lwd=2)
```

udca

Data from a trial of ursodeoxycholic acid

Description

Data from a trial of ursodeoxycholic acid (UDCA) in patients with primary biliary cirrhosis (PBC).

Usage

```r
udca
data(udca, package="survival")
```

Format

A data frame with 170 observations on the following 15 variables.

- `id` subject identifier
- `trt` treatment of 0=placebo, 1=UDCA
- `entry.dt` date of entry into the study
- `last.dt` date of last on-study visit
- `stage` stage of disease
- `bili` bilirubin value at entry
- `riskscore` the Mayo PBC risk score at entry
- `death.dt` date of death
- `tx.dt` date of liver transplant
- `hprogress.dt` date of histologic progression
- `varices.dt` appearance of esophageal varices
- `ascites.dt` appearance of ascites
- `enceph.dt` appearance of encephalopathy
- `double.dt` doubling of initial bilirubin
- `worsen.dt` worsening of symptoms by two stages
Details

This data set is used in the Therneau and Grambsch. The udca1 data set contains the baseline variables along with the time until the first endpoint (any of death, transplant, . . . , worsening). The udca2 data set treats all of the endpoints as parallel events and has a stratum for each.

References


Examples

# values found in table 8.3 of the book
fit1 <- coxph(Surv(futime, status) ~ trt + log(bili) + stage,
cluster =id, data=udca1)
fit2 <- coxph(Surv(futime, status) ~ trt + log(bili) + stage +
strata(endpoint), cluster=id, data=udca2)

untangle.specials  
Help Process the 'specials' Argument of the 'terms' Function.

Description

Given a terms structure and a desired special name, this returns an index appropriate for subscripting the terms structure and another appropriate for the data frame.

Usage

untangle.specials(tt, special, order=1)

Arguments

tt a terms object.
special the name of a special function, presumably used in the terms object.
order the order of the desired terms. If set to 2, interactions with the special function will be included.

Value

a list with two components:

vars a vector of variable names, as would be found in the data frame, of the specials.
terms a numeric vector, suitable for subscripting the terms structure, that indexes the terms in the expanded model formula which involve the special.
Examples

```
formula <- Surv(tt, ss) ~ x + z*strata(id)
tms <- terms(formula, specials="strata")
## the specials attribute
attr(tms, "specials")
## main effects
untangle.specials(tms, "strata")
## and interactions
untangle.specials(tms, "strata", order=1:2)
```

---

**uspop2**

**Projected US Population**

**Description**

US population by age and sex, for 2000 through 2020

**Format**

The data is a matrix with dimensions age, sex, and calendar year. Age goes from 0 through 100, where the value for age 100 is the total for all ages of 100 or greater.

**Details**

This data is often used as a "standardized" population for epidemiology studies.

**Source**


**See Also**

uspop

**Examples**

```
us50 <- uspop2[51:101,, "2000"]  # US 2000 population, 50 and over
age <- as.integer(dimnames(us50)[[1]])
smat <- model.matrix(~ factor(floor(age/5)) -1)
ustot <- t(smat) %*% us50  # totals by 5 year age groups
temp <- c(50, 55, 60, 65, 70, 75, 80, 85, 90, 95)
dimnames(ustot) <- list(c(paste(temp, temp+4, sep="-"), "100+"), c("male", "female"))
```
vcov.coxph

Variance-covariance matrix

**Description**

Extract and return the variance-covariance matrix.

**Usage**

```r
## S3 method for class 'coxph'
vcov(object, complete=TRUE, ...)
## S3 method for class 'survreg'
vcov(object, complete=TRUE, ...)
```

**Arguments**

- `object`: a fitted model object
- `complete`: logical indicating if the full variance-covariance matrix should be returned. This has an effect only for an over-determined fit where some of the coefficients are undefined, and `coef(object)` contains corresponding NA values. If `complete=TRUE` the returned matrix will have row/column for each coefficient, if FALSE it will contain rows/columns corresponding to the non-missing coefficients. The `coef()` function has a similar `complete` argument.
- `...`: additional arguments for method functions

**Details**

For the `coxph` and `survreg` functions the returned matrix is a particular generalized inverse: the row and column corresponding to any NA coefficients will be zero. This is a side effect of the generalized cholesky decomposition used in the underlying computation.

**Value**

a matrix

---

**veteran**

*Veterans' Administration Lung Cancer study*

**Description**

Randomised trial of two treatment regimens for lung cancer. This is a standard survival analysis data set.
xtfrm.Surv

Sorting order for Surv objects

Description
Sort survival objects into a partial order, which is the same one used internally for many of the calculations.

Usage

## S3 method for class 'Surv'
xtfrm(x)

Arguments

x a Surv object

Details
This creates a partial ordering of survival objects. The result is sorted in time order, for tied pairs of times right censored events come after observed events (censor after death), and left censored events are sorted before observed events. For counting process data (tstart, tstop, status) the ordering is by stop time, status, and start time, again with censoring last. Interval censored data is sorted using the midpoint of each interval.

The xtfrm routine is used internally by order and sort, so these results carry over to those routines.

Source

Usage

veteran
data(cancer, package="survival")

Format

trt: 1=standard 2=test
celltype: 1=squamous, 2=smallcell, 3=adeno, 4=large
time: survival time
status: censoring status
karno: Karnofsky performance score (100=good)
diagtime: months from diagnosis to randomisation
age: in years
prior: prior therapy 0=no, 10=yes
Value

a vector of integers which will have the same sort order as \( x \).

Author(s)

Terry Therneau

See Also

\texttt{sort}, \texttt{order}

Examples

```r
test <- c(Surv(c(10, 9, 9, 8, 8, 8, 7, 5, 5, 4), rep(1:0, 5)), Surv(6.2, NA))

test

sort(test)
```

---

**yates**

*Population prediction*

---

**Description**

Compute population marginal means (PMM) from a model fit, for a chosen population and statistic.

**Usage**

```r
yates(fit, term, population = c("data", "factorial", "sas"),
      levels, test = c("global", "trend", "pairwise"), predict = "linear",
      options, nsim = 200, method = c("direct", "sgtt"))
```

**Arguments**

- **fit**: a model fit. Examples using \texttt{lm}, \texttt{glm}, and \texttt{coxph} objects are given in the vignette.
- **term**: the term from the model which is to be evaluated. This can be written as a character string or as a formula.
- **population**: the population to be used for the adjusting variables. User can supply their own data frame or select one of the built-in choices. The argument also allows "empirical" and "yates" as aliases for data and factorial, respectively, and ignores case.
- **levels**: optional, what values for \texttt{term} should be used.
- **test**: the test for comparing the population predictions.
- **predict**: what to predict. For a \texttt{glm} model this might be the 'link' or 'response'. For a \texttt{coxph} model it can be linear, risk, or survival. User-written functions are allowed.
- **options**: optional arguments for the prediction method.
number of simulations used to compute a variance for the predictions. This is not needed for the linear predictor.

- **method**: the computational approach for testing equality of the population predictions. Either the direct approach or the algorithm used by the SAS glim procedure for "type 3" tests.

**Details**

The many options and details of this function are best described in a vignette on population prediction.

**Value**

an object of class `yates` with components of

- **estimate**: a data frame with one row for each level of the term, and columns containing the level, the mean population predicted value (mppv) and its standard deviation.
- **tests**: a matrix giving the test statistics
- **mvar**: the full variance-covariance matrix of the mppv values
- **summary**: optional: any further summary if the values provided by the prediction method.

**Author(s)**

Terry Therneau

**Examples**

```r
fit1 <- lm(skips ~ Solder*Opening + Mask, data = solder)
yates(fit1, ~Opening, population = "factorial")

fit2 <- coxph(Surv(time, status) ~ factor(ph.ecog)*sex + age, lung)
yates(fit2, ~ ph.ecog, predict="risk") # hazard ratio
```

**Description**

This is a method which is called by the `yates` function, in order to setup the code to handle a particular model type. Methods for glm, coxph, and default are part of the survival package.

**Usage**

`yates_setup(fit, ...)`
Arguments

fit          a fitted model object
...         optional arguments for some methods

Details

If the predicted value should be the linear predictor, the function should return NULL. The yates routine has particularly efficient code for this case. Otherwise it should return a prediction function or a list of two elements containing the prediction function and a summary function. The prediction function will be passed the linear predictor as a single argument and should return a vector of predicted values.

Note

See the vignette on population prediction for more details.

Author(s)

Terry Therneau

See Also

yates
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