Package ‘greyzoneSurv’

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Title   Fit a Grey-Zone Model with Survival Data
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Description Allows one to classify patients into low, intermediate, and high risk groups for disease progression based on a continuous marker that is associated with progression-free survival. It uses a latent class model to link the marker and survival outcome and produces two cut-offs for the marker to divide patients into three groups. See the References section for more details.
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greyzoneSurv-package  Fit a Grey-Zone Model with Survival Data

Description

Allows one to classify patients into low, intermediate, and high risk groups for disease progression based on a continuous marker that is associated with progression-free survival. It uses a latent class model to link the marker and survival outcome and produces two cutoffs for the marker to divide patients into three groups. See the References section for more details.

Details

To fit the grey-zone model, one would need to call the functions in the order of `em.func`, `cov.func`, and `greyzone.func`.

The package also provides a function `bestcut2` to fit a 2-group model, that is, it will find an optimal cutoff of the marker to divide patients into high and low risk groups. Plus there is a function `gensurvdata` to generate survival data with a fixed censoring rate.

Author(s)

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References

Pingping Qu, Bart Barlogie and John Crowley (2015) "Using a Latent Class Model to Refine Risk Stratification in Multiple Myeloma" (under review)

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bestcut2  Find an Optimal Cutoff for the 2-group Model

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Description

This function uses a brute force method to search for the best cutoff value for a marker based on the log rank test to divide patients into high and low risk groups given survival data.

Usage

```r
bestcut2(data, stime, sind, var, leave = 20)
```
**cox.summary**

**Summarize after Fitting Cox regression**

**Description**

A wrapper function to summarize 2-group and grey-zone (3-group) models with $R^2$ and c-index after fitting Cox regression.

**Usage**

```r
cox.summary(stime, sind, var)
```

**Arguments**

- **stime**: A nx1 vector of survival time
- **sind**: A nx1 vector of censoring indicator
- **var**: A nx1 vector of risk groupings to correlate with survival. For the 2-group and 3-group models, it is a categorical vector

**Value**

It returns a data frame with the input data as well as the final optimal high and low risk groupings saved in the column `bestcut2` (1=high risk and 0=low risk). Additionally, it has columns such as the cutoff value for the marker, the chi-square statistics and the log rank p values for testing equality of survival in the resulting high and low risk groups from using each possible marker value as cutoff.

**Examples**

```r
## Use the package data "mydata" to fit the 2-group model
data(mydata)
res=bestcut2(data=mydata, stime='time', sind='event', var='x')
table(res[, 'bestcut2'])

#compare the true groupings and that from the 2-group model
table(res[, c('xhigh', 'bestcut2')])
```
Value

It returns a vector with components such as the p value from fitting a Cox regression model, AIC (Akaike information criterion), and the c-index [1, 2], $R^2$ from the XO method [3], the OXS method [4] and the Nagelkerke method.

References


See Also

`greyzone.func`

Examples

```r
# See Examples for greyzone.func in this package.
```

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**genSurvData**

*Simulate Survival Outcome Given Marker Values*

Description

This function simulates survival outcome with a fixed censoring rate based on a Weibull distribution given input values such as study recruitment period, patient marker values, their true risk groupings (1=high risk and 0=low risk), and true regression coefficients.

Usage

```r
genSurvData(n, recruitment.yrs=2, baseline.hazard=365.25*5, shape=1, censoring.rate=0, beta.continuous, beta.binary=0, x, xhigh, ran.seed)
```

Arguments

- `n` Sample size
- `recruitment.yrs` Patient recruitment period in years (default=2)
- `baseline.hazard` Baseline hazard, which is the mean survival time (in days) when covariates=0 (default=365.25*5 days)
shape  The shape parameter for the Weibull distribution (it is exponential when shape=1)
censoring.rate  Censoring rate
beta.continuous  A true regression coefficient that links the continuous marker values to the survival outcome
beta.binary  A true regression coefficient that links the high risk group to the survival outcome
x  A nx1 vector for the marker values
xhigh  A nx1 vector of 1s and 0s indicating patient true risk identities (1=high risk and 0=low risk)
ran.seed  Seed number for random number generation

Details
The function can be used to generate survival data if you do not have any to try the grey-zone model.

Value
It returns a list with two components: the simulated survival data in days and the final censoring rate (which should be the same as the input censoring rate).

See Also
em.func, cov.func, greyzone.func

Examples
```r
## Generate package data called “mydata”
## Simulate high/low risk groupings, continuous marker values for each group, and survival data
## so that the higher marker values correspond to shorter survival.

n=300
censoring.rate=0.3
rate.lrisk=0.7  #rate of low risk
n.lrisk=n*rate.lrisk
n.hrisk=n-n.lrisk
mu=3
beta.continuous=0.5
beta.binary=0.5
ran.seed=1000
set.seed(ran.seed)
x0=rnorm(n.lrisk, 0, 1)  #low risk patients have marker values distributed as Normal(0,1)
set.seed(ran.seed)
x1=rnorm(n.hrisk, mu, 1)  #high risk patients have maker values distributed as Normal(mu,1)
score=c(x0, x1)
score.high=c(rep(0, n.lrisk), rep(1, n.hrisk))
mydata=genSurvData(n=n, censoring.rate=censoring.rate, beta.continuous=beta.continuous, beta.binary=beta.binary, x=score, xhigh=score.high, ran.seed=ran.seed)$data

dim(mydata)
head(mydata)
```
greyzone.funcs  
*Fit a Grey-Zone Model with Survival Data*

**Description**

Find two cutoffs for a marker by fitting a grey-zone model that will define high, intermediate, and low risk groups when the outcome is survival.

**Usage**

em.func(error = 0.001, max.iter = 300, initial.values=NULL, y, delta, x)

cov.func(em.results, y, delta, x)

greyzone.func(cov.results, y, delta, x, plot.logistic=T)

**Arguments**

- **error**  
  Convergence criterion as largest difference in parameter estimates between two consecutive iterations

- **max.iter**  
  Maximum number of iterations

- **initial.values**  
  A list of initial parameters with such components as z, lamda, beta, and gamma with the default being NULL.
  
z is an initial guess of dimension nx1 regarding whether each subject or patient is at high risk (=1) or low risk (=0); lamda is an initial guess about the scale parameter for a Weibull distribution (lamda=1 for an exponential distribution); beta is an initial guess on the regression coefficients which include an intercept and a slope linking the latent class variable and the survival outcome, and gamma is an initial guess on the regression coefficients which include an intercept and a slope linking the latent class variable and the marker values.

- **y**  
  A nx1 vector of survival time in years

- **delta**  
  A nx1 vector of censoring indicator

- **x**  
  A nx2 matrix, with the 1st column being all 1s and the 2nd column being the marker values

- **em.results**  
  Results from calling em.func

- **cov.results**  
  Results from calling cov.func

- **plot.logistic**  
  A logical variable. If T, it will plot the fitted logistic function between the marker values and the fitted probability of high risk for each patient

**Details**

The package assumes higher marker values correspond to shorter survival times, so it is important to make sure this is the case with your data. If initial.values is NULL, the function will generate initial values automatically based on this assumption.
Value

The function `em.func` returns a list with components such as parameter estimates and number of iterations from the EM algorithm as well as whether the EM algorithm fitting generated an error.

The function `cov.func` returns a list with components such as the variance-covariance matrix, standard errors and 95% confidence limits for the parameter estimates estimated from calling `em.func`. An additional component it returns is the estimated probability of being at high risk for each patient given patient survival and marker data.

The function `greyzone.func` returns a list with components such as the grey zone cutoffs `greyzone.ll` and `greyzone.ul` so that patient will be classified as low risk if marker value is < `greyzone.ll`, high risk if marker value is >= `greyzone.ul`, and intermediate risk (or in the grey zone) if marker value is >= `greyzone.ll` and < `greyzone.ul`.

See Also

`genSurvData`, `cox.summary`

Examples

```r
# use the package data "mydata" to fit the grey-zone model in 3 steps
data(mydata)
dim(mydata)
head(mydata)

# step 1: extract information needed for fitting the model and
# make some initial guesses of some parameter values
n=nrow(mydata)
y=mydata$time/365.25
delta=mydata$event
score=mydata$x
x=cbind(rep(1L, n), score)

# step 2: get EM estimates and variance-covariance matrix
results.em=em.func(initial.values=NULL, y=y, delta=delta, x=x)
is.na(results.em$error)
# only if above is true, you should proceed; otherwise try a different set
# of intial values and try calling em.func again
names(results.em)
# after you successfully get an EM solution proceed to get the variance-covariance matrix
results.cov=cov.func(results.em, y=y, delta=delta, x=x)
names(results.cov)

# step 3: when there are no errors above proceed to calculate the grey zone
results=greyzone.func(results.cov, y, delta, x, plot.logistic=FALSE)
names(results)
!is.na(results$greyzone.ll) & !is.na(results$greyzone.ul)
# only when above is true, you have a grey-zone solution and you can proceed
# sometimes there is not a grey-zone solution even if the EM fitting is successful.
# In that case, it means the grey-zone model is not a good fit for the data.
score3=rep(0, n)
score3[score3>=results$greyzone.ll & score3<results$greyzone.ul]=1
```
mydata

score3[score>results$greyzone.ul]=2
# if you get through steps 1-3, you are done fitting the grey-zone model!

# now you may want to compare with a 2-group model, so fit a 2-group model
res=bestcut2(data=mydata, stime='time', sind='event', var='x')
score2=res['bestcut2']

# then compare the 2-group and grey-zone models, but note here we compare on the training data
# ideally we want to compare them on a test data set
cox.2group=cox.summary(stime=mydata$time, sind=mydata$event, var=score2)
cox.3group=cox.summary(stime=mydata$time, sind=mydata$event, var=score3)

cox.2group
cox.3group
fit.2group = survfit(Surv(mydata$time, mydata$event)-score2)
fit.3group = survfit(Surv(mydata$time, mydata$event)-score3)

par(mfrow=c(1,2))
plot(fit.2group, lty=1:2, main='2-group model', las=1,
ylab='Probability of Survival', xlab='Days from Time 0')
plot(fit.3group, lty=1:3, main='grey-zone model', las=1,
ylab='Probability of Survival', xlab='Days from Time 0')

---

mydata

Package Data

Description

The data set can be used to test the grey-zone and 2-group models.

Usage

data(mydata)

Format

The data set is a data frame with 300 rows and 4 columns for patient survival time (time), censoring event (event), marker values (x), and true risk groups (xhigh=1 for high risk and 0 for low risk).

Details

The data were generated based on a Weibull distribution for survival times and normal distributions for marker values in both the high and low risk patients.

Source

The data were generated with the code in the Examples of genSurvData in this package.

Examples

data(mydata)
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