

Package ‘simulatorZ’

October 9, 2015

Type Package

Title Simulator for Collections of Independent Genomic Data Sets

Version 1.2.0

Date 2014-08-03

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Description simulatorZ is a package intended primarily to simulate collections of independent genomic data sets, as well as performing training and validation with predicting algorithms. It supports ExpressionSets and SummarizedExperiment objects.

License Artistic-2.0

Depends R (>= 3.1), Biobase, survival, CoxBoost

Imports GenomicRanges, gbm, Hmisc, stats, graphics

Suggests BiocGenerics, RUnit, BiocStyle, curatedOvarianData, parathyroidSE, superpc

URL <https://github.com/zhangyuqing/simulatorZ>

BugReports <https://github.com/zhangyuqing/simulatorZ>

biocViews Survival

NeedsCompilation yes

R topics documented:

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 simulatorZ-package *Simulator for Collections of Independent Genomic Data Sets*

Description

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Details

Package:	simulatorZ
Type:	Package
Title:	Simulator for Collections of Independent Genomic Data Sets
Version:	0.99.1
Date:	2014-08-03
Author:	Yuqing Zhang, Christoph Bernau, Levi Waldron
Maintainer:	Yuqing Zhang <zhangyuqing.pkusms@gmail.com>
License:	Artistic-2.0
Depends:	R (>= 3.1), Biobase, survival, CoxBoost
Imports:	GenomicRanges, gbm, Hmisc
Suggests:	BiocStyle, curatedOvarianData, parathyroidSE, superpc
URL:	https://github.com/zhangyuqing/simulatorZ
BugReports:	https://github.com/zhangyuqing/simulatorZ
biocViews:	Survival

Author(s)

Yuqing Zhang, Christoph Bernau, Levi Waldron

 cvSubsets

cvSubsets

Description

To generate a list of subsets(indices of observations) from one set

Usage

```
cvSubsets(obj, fold)
```

Arguments

obj	a ExpressionSet, matrix or SummarizedExperiment object. If it is a matrix, columns represent samples
fold	the number of folds in cross validation. Number of observations in the set does not need to be a multiple of fold

Value

returns the list of indices of subsets

Author(s)

Yuqing Zhang

Examples

```
library(curatedOvarianData)

data(E.MTAB.386_eset)

set.seed(8)

id <- cvSubsets(E.MTAB.386_eset, 3)

subset1 <- E.MTAB.386_eset[, id[[1]]]

subset2 <- E.MTAB.386_eset[, id[[2]]]

subset3 <- E.MTAB.386_eset[, id[[3]]]
```

```

## Number of observations in the set does not need to be a multiple of

## the fold parameter

id2 <- cvSubsets(E.MTAB.386_eset, 5)

subsets <- list()

subsets[[1]] <- E.MTAB.386_eset[, id2[[1]]]

subsets[[2]] <- E.MTAB.386_eset[, id2[[2]]]

subsets[[3]] <- E.MTAB.386_eset[, id2[[3]]]

subsets[[4]] <- E.MTAB.386_eset[, id2[[4]]]

subsets[[5]] <- E.MTAB.386_eset[, id2[[5]]]

```

funcV

*funCV***Description**

Cross validation function

Usage

```
funCV(obj, fold, y.var, trainFun = masomenos, funCvSubset = cvSubsets)
```

Arguments

obj	a ExpressionSet, matrix or SummarizedExperiment object. If it is a matrix, columns represent samples
fold	the number of folds in cross validation
y.var	response variable, matrix, data.frame(with 2 columns) or Surv object
trainFun	training function, which takes gene expression matrix X and response variable y as input, the coefficients as output
funCvSubset	function to divide one Expression Set into subsets for cross validation

Value

returns the c statistics of cross validation(CV)

Author(s)

Yuqing Zhang

Examples

```
library(curatedOvarianData)

library(GenomicRanges)

set.seed(8)

data( E.MTAB.386_eset )

eset <- E.MTAB.386_eset[1:100, 1:30]

time <- eset$days_to_death

cens.chr <- eset$vital_status

cens <- c()

for(i in 1:length(cens.chr)){

  if(cens.chr=="living") cens[i] <- 1

  else cens[i] <- 0

}

y <- Surv(time, cens)

y1 <- cbind(time, cens)
```

```
nrows <- 200; ncols <- 6

counts <- matrix(runif(nrows * ncols, 1, 1e4), nrows)

rowData <- GRanges(rep(c("chr1", "chr2"), c(50, 150)),
                     IRanges(floor(runif(200, 1e5, 1e6)), width=100),
                     strand=sample(c("+", "-"), 200, TRUE))

colData <- DataFrame(Treatment=rep(c("ChIP", "Input"), 3),
                      row.names=LETTERS[1:6])

sset <- SummarizedExperiment(assays=SimpleList(counts=counts),
                             rowData=rowData, colData=colData)

time <- sample(4500:4700, 6, replace=TRUE)

cens <- sample(0:1, 6, replace=TRUE)

y.vars <- Surv(time, cens)

funCV(eset, 3, y)

funCV(eset, 3, y1, trainFun=plusMinus)

funCV(exprs(eset), 3, y)
```

```
funCV(sset, 3, y.vars)

## any training function will do as long as it takes the gene expression matrix X

## and response variable y(matrix, data.frame or Surv object) as parameters, and

## return the coefficients as its value
```

*geneFilter**geneFilter*

Description

the function to filter genes by Intergrative Correlation

Usage

```
geneFilter(obj, cor.cutoff = 0.5)
```

Arguments

obj	a list of ExpressionSet, matrix or SummarizedExperiment objects. If its elements are matrices, columns represent samples, rows represent genes
cor.cutoff	the cutoff threshold for filtering genes. Only when the integrative correlation between every pair of sets is larger than the cutoff value, will the gene be selected.

Value

returns a list of ExpressionSets matrix or SummarizedExperiment objects with genes filtered

Author(s)

Yuqing Zhang

References

Garrett-Mayer, E., Parmigiani, G., Zhong, X., Cope, L.,
Gabrielson, E., Cross-study validation and combined analysis of gene
expression microarray data. Biostatistics. 2008 Apr;9(2):333-354.

Examples

```
set.seed(8)

library(curatedOvarianData)

library(GenomicRanges)

data(GSE17260_eset)

data(E.MTAB.386_eset)

data(GSE14764_eset)

esets <- list(GSE17260=GSE17260_eset, E.MTAB.386=E.MTAB.386_eset, GSE14764=GSE14764_eset)

esets.list <- lapply(esets, function(eset){

  return(eset[1:1500, 1:10])

})

result.set <- geneFilter(esets.list, 0)

result.set

### as we cannot calculate correlation with one set, this function just

### delivers the same set if esets has length 1

result.oneset <- geneFilter(esets.list[1])

result.oneset
```

```
## Support matrices

X.list <- lapply(esets.list, function(eset){

  return(exprs(eset)) ## Columns represent samples!

})

result.set <- geneFilter(X.list, 0)

dim(result.set[[1]])

## Support SummarizedExperiment

nrows <- 200; ncols <- 6

counts <- matrix(runif(nrows * ncols, 1, 1e4), nrows)

rowData <- GRanges(rep(c("chr1", "chr2"), c(50, 150)),

  IRanges(floor(runif(200, 1e5, 1e6)), width=100),

  strand=sample(c("+", "-"), 200, TRUE))

colData <- DataFrame(Treatment=rep(c("ChIP", "Input"), 3),

  row.names=LETTERS[1:6])

sset <- SummarizedExperiment(assays=SimpleList(counts=counts),
```

```

    rowData=rowData, colData=colData)

s.list <- list(sset, sset)

result.set <- geneFilter(s.list, 0.9)

## the same set should resemble each other, no genes filtered

dim(assay(result.set[[1]]))

```

getTrueModel*getTrueModel***Description**

The parametric bootstrap simulation depends on the true model of original sets.
 This function is to generate useful values from the true models for further analysis.
 We fit CoxBoost to the original sets and use the coefficients to simulate
 the survival and censoring time. grid, survH, censH, which are useful for this purpose.
 grid=grid corresponding to hazard estimations censH and survH
 survH=cumulative hazard for survival times distribution
 censH=cumulative hazard for censoring times distribution

Usage

```
getTrueModel(obj, y.vars, parstep)
```

Arguments

<i>obj</i>	a list of ExpressionSets, matrix or SummarizedExperiment
<i>y.vars</i>	a list of response variables, Surv, matrix or data.frame object
<i>parstep</i>	number of steps in CoxBoost

Value

returns a list of values:

beta: True coefficients obtained by fitting CoxBoost to the original ExpressionSets
 grid: timeline grid corresponding to hazard estimations censH and survH
 survH: cumulative hazard for survival times distribution
 censH: cumulative hazard for censoring times distribution
 lp: true linear predictors

Author(s)

Yuqing Zhang, Christoph Bernau

Examples

```
library(curatedOvarianData)

data(GSE17260_eset)

data(E.MTAB.386_eset)

data(GSE14764_eset)

esets <- list(GSE17260=GSE17260_eset, E.MTAB.386=EMTAB.386_eset, GSE14764=GSE14764_eset)

esets.list <- lapply(esets, function(eset){

  return(eset[1:500, 1:20])

})

## simulate on multiple ExpressionSets

set.seed(8)

y.list <- lapply(esets.list, function(eset){

  time <- eset$days_to_death

  cens.chr <- eset$vital_status
```

```

cens <- c()

for(i in 1:length(cens.chr)){
  if(cens.chr[i] == "living") cens[i] <- 1
  else cens[i] <- 0
}

y <- Surv(time, cens)

return(y)
}

res1 <- getTrueModel(esets.list, y.list, 100)

## Get true model from one set

res2 <- getTrueModel(esets.list[1], y.list[1], 100)

names(res2)

res2$lp

## note that y.list[1] cannot be replaced by y.list[[1]]

```

Description

function for Mas-o-menos algorithm

Usage

```
masomenos(X, y)
```

Arguments

- X gene expression matrix, with rows corresponding to patients and columns corresponding to genes
- y response variable, a data.frame, matrix or Surv object, with the first column as time to event and the second column censoring status

Value

return betas

Author(s)

Yuqing Zhang

References

- Zhao, S., Huttenhower, G. P. C., and Waldron, L. (2013). Mas-o-menos: a simple sign averaging method for discrimination in genomic data analysis. <http://biostats.bepress.com/harvardbiostat/paper158/>. Accessed: 2013-10-24.

Examples

```
set.seed(8)

library(curatedOvarianData)

data( E.MTAB.386_eset )

eset <- E.MTAB.386_eset[1:100, 1:30]

X <- t(exprs(eset))
```

```

time <- eset$days_to_death

cens <- sample(0:1, 30, replace=TRUE)

y <- Surv(time, cens)

beta <- masomenos(X, y)

beta

```

plusMinus*plusMinus***Description**

function for plusMinus algorithm

Usage

```

plusMinus(X, y, lambda = NULL, tuningpar = "nfeatures", standardize = FALSE,
          directionality = "posneg", ties.method = "average", votingthresholdquantile = 0.5,
          modeltype = "plusminus")

```

Arguments

X	gene expression matrix
y	response variables
lambda	
tuningpar	
standardize	
directionality	
ties.method	
votingthresholdquantile	
modeltype	

Value

returns regression coefficients

Author(s)

Levi Waldron

References

Zhao, S., Huttenhower, G. P. C., and Waldron, L. (2013). Mas-o-menos: a simple sign averaging method for discrimination in genomic data analysis. <http://biostats.bepress.com/harvardbiostat/paper158/>. Accessed: 2013-10-24.

Examples

```
set.seed(8)

library(curatedOvarianData)

data( E.MTAB.386_eset )

eset <- E.MTAB.386_eset[1:100, 1:30]

X <- t(exprs(eset))

time <- eset$days_to_death

cens <- sample(0:1, 30, replace=TRUE)

y <- Surv(time, cens)

beta <- plusMinus(X, y)
```

beta

rowCoxTests

rowCoxTests

Description

method for performing Cox regression

Usage

```
rowCoxTests(X, y, option = c("fast", "slow"), ...)
```

Arguments

- | | |
|---------------|--|
| X | Gene expression data. The following formats are available:
matrix Rows correspond to observations, columns to variables.
data.frame Rows correspond to observations, columns to variables.
ExpressionSet rowCoxTests will extract the expressions using exprs(). |
| y | Survival Response, an object of class:
Surv if X is of type data.frame or matrix
character if X is of type ExpressionSet.
In this case y is the name of the survival
response in the phenoData of X. If survival
time and indicator are stored separately
in the phenoData one can specify a two-element
character vector the first element representing
the survival time variable. |
| option | "fast" loops over rows in C, "slow" calls coxph
directly in R. The latter method may be used if
something goes wrong with the "fast" method. |
| ... | currently unused |

Value

dataframe with two columns: coef = Cox regression
coefficients, p.value =
Wald Test p-values. Rows correspond to the rows of X.

Author(s)

Christoph Bernau

Examples

```
#test

##regressor-matrix (gene expressions)

X<-matrix(rnorm(1e6),nrow=10000)

#seed

set.seed(123)

#times

time<-rnorm(n=ncol(X),mean=100)

#censoring(1->death)

status<-rbinom(n=ncol(X),size=1, prob=0.8)

##survival object

y<-Surv(time,status)

## Do 10,000 Cox regressions:

system.time(output <- rowCoxTests(X=X,y=y, option="fast"))
```

<code>simBootstrap</code>	<i>simBootstrap</i>
---------------------------	---------------------

Description

the driver function to perform three-step bootstrap resampling
to get independent genomic data sets

Usage

```
simBootstrap(obj, y.vars, n.samples, parstep, type = "two-steps",
            balance.variables = NULL, funSimData = simData, funTrueModel = getTrueModel,
            funSurvTime = simTime)
```

Arguments

<code>obj</code>	a list of ExpressionSet, matrix or SummarizedExperiment
<code>y.vars</code>	a list of reponse variables, elements can be class Surv, matrix or data.frame
<code>n.samples</code>	number of samples to resample in each set
<code>parstep</code>	step number to fit CoxBoost
<code>type</code>	whether to include resampling set labels
<code>balance.variables</code>	covariate names to balance in the simulated sets
<code>funSimData</code>	function to perform non-parametric bootstrap
<code>funTrueModel</code>	function to construct true models in original sets
<code>funSurvTime</code>	function to perform parametric bootstrap

Value

a list of values including:
`obj.list` = a list of simulated objects the same type as input
`indices.list` = a list of indices indicating which sample the simulated sample is in the original set
`setsID` = a vector to indicate the original ID of simulated sets, if `type=="original"`, `setsID` should be 1,2,3,...
`lp.list` = a list of true linear predictor of each original data sets
`beta.list` = a list of true coefficients used for simulating observations
`survH.list` = list of cumulative survival hazard
`censH.list` = list of cumulative censoring hazard
`grid.list` = list of timeline grid corresponding to `survH` and `censH` respectivley

Author(s)

Yuqing Zhang, Levi Waldron

Examples

```
library(curatedOvarianData)

library(GenomicRanges)

data(GSE17260_eset)

data(E.MTAB.386_eset)

data(GSE14764_eset)

esets <- list(GSE17260=GSE17260_eset, E.MTAB.386=E.MTAB.386_eset, GSE14764=GSE14764_eset)

esets.list <- lapply(esets, function(eset){

  return(eset[1:500, 1:20])

})

## simulate on multiple ExpressionSets

set.seed(8)

y.list <- lapply(esets.list, function(eset){

  time <- eset$days_to_death
```

```
cens.chr <- eset$vital_status

cens <- c()

for(i in 1:length(cens.chr)){

  if(cens.chr[i] == "living") cens[i] <- 1

  else cens[i] <- 0

}

y <- Surv(time, cens)

return(y)

})

simmodels <- simBootstrap(obj=esets.list, y.vars=y.list, 10, 100)

simmodels$obj.list[[1]]


# balance covariates

simmodels <- simBootstrap(obj=esets.list, y.vars=y.list, 10, 100,

                           balance.variables="tumorstage")

## Support SummarizedExperiment

nrows <- 200; ncols <- 10
```

```
counts <- matrix(runif(nrows * ncols, 1, 1e4), nrows)

rowData <- GRanges(rep(c("chr1", "chr2"), c(50, 150)),
                     IRanges(floor(runif(200, 1e5, 1e6)), width=100),
                     strand=sample(c("+", "-"), 200, TRUE))

colData <- DataFrame(Treatment=rep(c("ChIP", "Input"), 5),
                      row.names=LETTERS[1:10])

sset <- SummarizedExperiment(assays=SimpleList(counts=counts),
                             rowData=rowData, colData=colData)

s.list <- list(sset[,1:5], sset[,6:10])

time <- c(540, 527, 668, 587, 620, 540, 527, 668, 587, 620)

cens <- c(1, 0, 0, 1, 0, 1, 0, 0, 1, 0)

y.vars <- Surv(time, cens)

y.vars <- list(y.vars[1:5,],y.vars[1:5,])

simmodels <- simBootstrap(obj=s.list, y.vars=y.vars, 100, 100)
```

Description

simData is a function to perform non-parametric bootstrap resampling on a list of (original) data sets, both on set level and patient level, in order to simulate independent genomic sets.

Usage

```
simData(obj, n.samples, y.vars = list(), type = "two-steps",
        balance.variables = NULL)
```

Arguments

<code>obj</code>	a list of ExpressionSets, matrices or SummarizedExperiments. If elements are matrices, columns represent samples
<code>n.samples</code>	an integer indicating how many samples should be resampled from each set
<code>y.vars</code>	a list of response variables, can be Surv object, or matrix or data.frame with two columns
<code>type</code>	string "one-step" or "two-steps". If type="one-step", the function will skip resampling the datasets, and directly resample from the original list of <code>obj</code>
<code>balance.variables</code>	<code>balance.variables</code> will be a vector of covariate names that should be balanced in the simulation. After balancing, the prevalence of covariate in each result set should be the same as the overall distribution across all original data sets. Default is set as NULL, when it will not balance over any covariate. if isn't NULL, <code>esets</code> parameter should only be of class ExpressionSet

Value

returns a list of simulated ExpressionSets, with names indicating its original set, and indices of the original patients.

`prob.desired` and `prob.real` are only useful when `balance.variables` is set.

`prob.desired` shows overall distribution of the specified covariate. `prob.list`

shows the sampling probability in each set after balancing

Author(s)

Yuqing Zhang, Levi Waldron

Examples

```
library(curatedOvarianData)

library(GenomicRanges)

data(GSE17260_eset)

data(E.MTAB.386_eset)

data(GSE14764_eset)

esets <- list(GSE17260=GSE17260_eset, E.MTAB.386=E.MTAB.386_eset, GSE14764=GSE14764_eset)

esets.list <- lapply(esets, function(eset){

  return(eset[1:1000, 1:10])

})

## simulate on multiple ExpressionSets

set.seed(8)

# one-step bootstrap: skip resampling set labels

simmodels <- simData(esets.list, 20, type="one-step")

# two-step-non-parametric bootstrap
```

```
simmodels <- simData(esets.list, 10, type="two-steps")

## simulate one set

simmodels <- simData(list(esets.list[[1]]), 10, type="two-steps")

## balancing covariates

# single covariate

simmodels <- simData(list(esets.list[[1]]), 5, balance.variables="tumorstage")

# multiple covariates

simmodels <- simData(list(esets.list[[1]]), 5,
                      balance.variables=c("tumorstage", "age_at_initial_pathologic_diagnosis"))

## Support matrices

X.list <- lapply(esets.list, function(eset){

  return(exprs(eset))

})
```

```
simmodels <- simData(X.list, 20, type="two-steps")

## Support SummarizedExperiment

nrows <- 200; ncols <- 6

counts <- matrix(runif(nrows * ncols, 1, 1e4), nrows)

rowData <- GRanges(rep(c("chr1", "chr2"), c(50, 150)),

IRanges(floor(runif(200, 1e5, 1e6)), width=100),

strand=sample(c("+", "-"), 200, TRUE))

colData <- DataFrame(Treatment=rep(c("ChIP", "Input"), 3),

row.names=LETTERS[1:6])

sset <- SummarizedExperiment(assays=SimpleList(counts=counts),

rowData=rowData, colData=colData)

s.list <- list(sset[,1:3], sset[,4:6])

simmodels <- simData(s.list, 20, type="two-steps")
```

Description

simTime is a function to perform the parametric-bootstrap step, where we use the true coefficients and cumulative hazard to simulate survival and censoring.

Usage

```
simTime(simmodels, result)
```

Arguments

<i>simmodels</i>	a list in the form of the return value of <i>simData()</i> which consists of three lists: <i>obj</i> : a list of ExpressionSets, matrices or SummarizedExperiments <i>setsID</i> : a list of set labels indicating which original set the simulated one is from <i>indices</i> : a list of patient labels to tell which patient in the original set is drawn
<i>result</i>	a list in the form of return of <i>getTrueModel()</i> which consists of five lists: <i>Beta</i> : a list of coefficients obtained by <i>grid</i> : timeline grid corresponding to hazard estimations <i>censH</i> and <i>survH</i> <i>survH</i> : cumulative hazard for survival times distribution <i>censH</i> : cumulative hazard for censoring times distribution <i>lp</i> : true linear predictors

Value

survival time is saved in phenodata, here the function still returns the ExpressionSets

Author(s)

Yuqing Zhang, Christoph Bernau

Examples

```
library(curatedOvarianData)

data(GSE17260_eset)

data(E.MTAB.386_eset)

data(GSE14764_eset)

esets <- list(GSE17260=GSE17260_eset, E.MTAB.386=E.MTAB.386_eset, GSE14764=GSE14764_eset)
```

```
esets.list <- lapply(esets, function(eset){

  return(eset[1:500, 1:20])

})

## simulate on multiple ExpressionSets

set.seed(8)

y.list <- lapply(esets.list, function(eset){

  time <- eset$days_to_death

  cens.chr <- eset$vital_status

  cens <- c()

  for(i in 1:length(cens.chr)){

    if(cens.chr[i] == "living") cens[i] <- 1

    else cens[i] <- 0

  }

  y <- Surv(time, cens)

  return(y)

})
```

```
})
```

```
# To perform both parametric and non-parametric bootstrap, you can call simBootstrap()
```

```
# or, you can divide the steps into:
```

```
res <- getTrueModel(esets.list, y.list, 100)
```

```
simmodels <- simData(obj=esets.list, y.vars=y.list, n.samples=10)
```

```
# Then, use this function
```

```
simmodels <- simTime(simmodels=simmodels, result=res)
```

```
# it also supports performing only the parametric bootstrap step on a list of expressionsets
```

```
# but you need to construct the parameter by scratch
```

```
res <- getTrueModel(esets.list, y.list, 100)
```

```
setsID <- 1:length(esets.list)
```

```
indices <- list()
```

```
for(i in setsID){
```

```
    indices[[i]] <- 1:length(sampleNames(esets.list[[i]]))
```

```
}
```

```
simmodels <- list(obj=esets.list, y.vars=y.list, indices=indices, setsID=setsID)
```

```
new.simmodels <- simTime(simmodels=simmodels, result=res)
```

zmatrix

zmatrix

Description

generate a matrix of c statistics

Usage

```
zmatrix(obj, y.vars, fold, trainingFun = masomenos, cvFun = funCV,  
cvSubsetFun = cvSubsets)
```

Arguments

<code>obj</code>	a list of ExpressionSet, matrix or SummarizedExperiment objects, if its elements are matrices, columns represent samples
<code>y.vars</code>	a list of response variables, all the response variables should be matrix, data.frame(with 2 columns) or Surv object
<code>fold</code>	cvFun parameter, in this case passes to funCV()
<code>trainingFun</code>	training function
<code>cvFun</code>	function to perform cross study within one set
<code>cvSubsetFun</code>	function to divide the expression sets into subsets for cross validation

Value

outputs one matrix of validation statistics

Author(s)

Yuqing Zhang, Christoph Bernau, Levi Waldron

Examples

```
library(curatedOvarianData)

library(GenomicRanges)

data(GSE17260_eset)

data(E.MTAB.386_eset)

data(GSE14764_eset)

esets <- list(GSE17260=GSE17260_eset, E.MTAB.386=E.MTAB.386_eset, GSE14764=GSE14764_eset)

esets.list <- lapply(esets, function(eset){

  return(eset[1:500, 1:20])

})

## simulate on multiple ExpressionSets

set.seed(8)

y.list <- lapply(esets.list, function(eset){

  time <- eset$days_to_death

  cens.chr <- eset$vital_status

  cens <- c()
```

```
for(i in 1:length(cens.chr)){
  if(cens.chr[i] == "living") cens[i] <- 1
  else cens[i] <- 0
}

y <- Surv(time, cens)

return(y)
}

# generate on original ExpressionSets

z <- zmatrix(esets.list, y.list, 3)

# generate on simulated ExpressionSets

simmodels <- simBootstrap(esets.list, y.list, 10, 100)

z <- zmatrix(simmodels$obj.list, simmodels$y.vars.list, 3)

# support matrix

X.list <- lapply(esets.list, function(eset){
```

```
newx <- exprs(eset) ### columns represent samples !!

return(newx)

})

z <- zmatrix(X.list, y.list, 3)

# support SummarizedExperiment

nrows <- 200; ncols <- 6

counts <- matrix(runif(nrows * ncols, 1, 1e4), nrows)

rowData <- GRanges(rep(c("chr1", "chr2"), c(50, 150)),

IRanges(floor(runif(200, 1e5, 1e6)), width=100),

strand=sample(c("+", "-"), 200, TRUE))

colData <- DataFrame(Treatment=rep(c("ChIP", "Input"), 3),

row.names=LETTERS[1:6])

sset <- SummarizedExperiment(assays=SimpleList(counts=counts),

rowData=rowData, colData=colData)

time <- sample(4500:4700, 6, replace=TRUE)

cens <- sample(0:1, 6, replace=TRUE)
```

```
y.vars <- Surv(time, cens)
```

```
z <- zmatrix(list(sset[,1:3], sset[,4:6]), list(y.vars[1:3,],y.vars[4:6,]), 3)
```

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