

# Package ‘survcomp’

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**Type** Package

**Title** Performance Assessment and Comparison for Survival Analysis

**Version** 1.8.1

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**Description** R package providing functions to assess and to compare the performance of risk prediction (survival) models.

**biocViews** GeneExpression, DifferentialExpression, Visualization

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**Depends** R (>= 2.10), survival, prodlim

**Suggests** Hmisc, CPE, clinfun, survJamda, Biobase, xtable

**Imports** ipred, SuppDists, KernSmooth, survivalROC, bootstrap, grid,rmeta

**License** Artistic-2.0

**URL** <http://compbio.dfc.harvard.edu>

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survcomp-package

*Performance Assessment and Comparison for Survival Analysis*

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## Description

Functions to perform the performance assessment and comparison of risk prediction (survival) models.

## Details

Package: survcomp  
 Type: Package  
 Version: 1.8.1  
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 License: Artistic-2.0

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**See Also**

survival, Hmisc, Design, prodlim, survivalROC, ipred, bootstrap, CPE, clinfun

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breastCancerData

*Sample data containing six datasets for gene expression, annotations and clinical data.*

---

**Description**

This dataset contains a subset of the gene expression, annotations and clinical data from 6 different datasets (see section details). The subsets contain the seven gene signature introduced by Desmedt et al. 2008.

**Usage**

```
data(breastCancerData)
```

## Format

Six ExpressionSets. Example for 'mainz7g': eSet with 7 features and 200 samples, containing:

- `exprs(mainz7g)`: Matrix containing gene expressions as measured by Affymetrix hgu133a technology (single-channel, oligonucleotides).
- `fData(mainz7g)`: AnnotatedDataFrame containing annotations of Affy microarray platform hgu133a.
- `pData(mainz7g)`: AnnotatedDataFrame containing Clinical information of the breast cancer patients whose tumors were hybridized.
- `experimentalData(mainz7g)`: MIAME object containing information about the dataset.
- `annotation(mainz7g)`: Name of the affy chip.

## Details

This dataset represents subsets of the studies published by Schmidt et al. 2008 [mainz7g], Wang, et al. 2005 and Minn et al. 2007 [vdx7g], Miller et al. 2005 [upp7g], Sotiriou et al. 2006 [unt7g], Desmedt et al. 2007 and TRANSBIG [transbig7g], van't Veer et al. 2002 and van de Vijver et al. 2002 [nki7g]. Each subset contains the genes AURKA (also known as STK6, STK7, or STK15), PLAU (also known as uPA), STAT1, VEGF, CASP3, ESR1, and ERBB2, as introduced by Desmedt et al. 2008. The seven genes represent the proliferation, tumor invasion/metastasis, immune response, angiogenesis, apoptosis phenotypes, and the ER and HER2 signaling, respectively.

## Source

**mainz:** <http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE11121>

**transbig:** <http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE7390>

**upp:** <http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE3494>

**unt:** <http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE2990>

<http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE6532>

**vdx:** <http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE2034>

<http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE5327>

**nki:** <http://www.rii.com/publications/2002/vantveer.html>

## References

Marcus Schmidt, Daniel Boehm, Christian von Toerne, Eric Steiner, Alexander Puhl, Heryk Pilch, Hans-Anton Lehr, Jan G. Hengstler, Hainz Koelbl and Mathias Gehrman (2008) "The Humoral Immune System Has a Key Prognostic Impact in Node-Negative Breast Cancer", *Cancer Research*, **68**(13):5405-5413

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patient survival" *Proceedings of the National Academy of Sciences of the United States of America* **102**(38):13550-13555

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Andy J. Minn, Gaorav P. Gupta, David Padua, Paula Bos, Don X. Nguyen, Dimitry Nuyten, Bas Kreike, Yi Zhang, Yixin Wang, Hemant Ishwaran, John A. Foekens, Marc van de Vijver and Joan Massague (2007) "Lung metastasis genes couple breast tumor size and metastatic spread", *Proceedings of the National Academy of Sciences*, **104**(16):6740-6745

Laura J. van't Veer, Hongyue Dai, Marc J. van de Vijver, Yudong D. He, Augustinus A.M. Hart, Mao Mao, Hans L. Peterse, Karin van der Kooy, Matthew J. Marton, Anke T. Witteveen, George J. Schreiber, Ron M. Kerkhoven, Chris Roberts, Peter S. Linsley, Rene Bernards and Stephen H. Friend (2002) "Gene expression profiling predicts clinical outcome of breast cancer", *Nature*, **415**:530-536

M. J. van de Vijver, Y. D. He, L. van't Veer, H. Dai, A. M. Hart, D. W. Voskuil, G. J. Schreiber, J. L. Peterse, C. Roberts, M. J. Marton, M. Parrish, D. Atsma, A. Witteveen, A. Glas, L. Delahaye, T. van der Velde, H. Bartelink, S. Rodenhuis, E. T. Rutgers, S. H. Friend and R. Bernards (2002) "A Gene Expression Signature as a Predictor of Survival in Breast Cancer", *New England Journal of Medicine*, **347**(25):1999-2009

## Examples

```
## load Biobase package
library(Biobase)
## load the dataset
data(breastCancerData)
#####
## Example for the mainz7g dataset
#####
## show the first 5 columns of the expression data
exprs(mainz7g)[,1:5]
## show the first 6 rows of the phenotype data
head(pData(mainz7g))
## show first 20 feature names
featureNames(mainz7g)
## show the experiment data summary
experimentData(mainz7g)
## show the used platform
annotation(mainz7g)
## show the abstract for this dataset
abstract(mainz7g)
```

censor.time

*Function to artificially censor survival data*

---

**Description**

The function censors the survival data at a specific point in time. This is useful if you used datasets having different follow-up periods.

**Usage**

```
censor.time(surv.time, surv.event, time.cens = 0)
```

**Arguments**

surv.time	vector of times to event occurrence
surv.event	vector of indicators for event occurrence
time.cens	point in time at which the survival data must be censored

**Value**

surv.time.cens	vector of censored times to event occurrence
surv.event.cens	vector of censored indicators for event occurrence

**Author(s)**

Benjamin Haibe-Kains

**Examples**

```
set.seed(12345)
stime <- rexp(30)
cens <- runif(30,0.5,2)
sevent <- as.numeric(stime <= cens)
stime <- pmin(stime, cens)
censor.time(surv.time=stime, surv.event=sevent, time.cens=1)
```

---

cindex.comp*Function to compare two concordance indices*

---

**Description**

This function compares two concordance indices computed from the same data by using the function concordance.index. The statistical test is a Student t test for dependent samples.

**Usage**

```
cindex.comp(cindex1, cindex2)
```

**Arguments**

cindex1	first concordance index as returned by the concordance.index function.
cindex2	second concordance index as returned by the concordance.index function.

**Details**

The two concordance indices must be computed from the same samples (and corresponding survival data). The function uses a Student t test for dependent samples.

**Value**

p.value	p-value from the Student t test for the comparison $cindex1 > cindex2$ .
cindex1	value of the first concordance index.
cindex2	value of the second concordance index.

**Author(s)**

Benjamin Haibe-Kains

**References**

Haibe-Kains, B. and Desmedt, C. and Sotiriou, C. and Bontempi, G. (2008) "A comparative study of survival models for breast cancer prognostication based on microarray data: does a single gene beat them all?", *Bioinformatics*, **24**(19), pages 2200–2208.

**See Also**

[concordance.index](#).

**Examples**

```
set.seed(12345)
age <- rnorm(100, 50, 10)
size <- rexp(100,1)
stime <- rexp(100)
cens <- runif(100,.5,2)
sevent <- as.numeric(stime <= cens)
stime <- pmin(stime, cens)
c1 <- concordance.index(x=age, surv.time=stime, surv.event=sevent,
  method="noether")
c2 <- concordance.index(x=size, surv.time=stime, surv.event=sevent,
  method="noether")
cindex.comp(c1, c2)
```

---

cindex.comp.meta      *Function to compare two concordance indices*

---

### Description

This function compares two lists of concordance indices computed from the same survival data by using the function `concordance.index`. The statistical test is a Student t test for dependent samples.

### Usage

```
cindex.comp.meta(list.cindex1, list.cindex2, hetero = FALSE)
```

### Arguments

<code>list.cindex1</code>	first list of concordance indices as returned by the <code>concordance.index</code> function.
<code>list.cindex2</code>	second list of concordance indices as returned by the <code>concordance.index</code> function.
<code>hetero</code>	if TRUE, a random effect model is used to compute the meta-estimators. Otherwise a fixed effect model is used.

### Details

In meta-analysis, we estimate the statistic of interest in several independent datasets. It results a list of estimates such as list of concordance indices. The two lists of concordance indices must be computed from the same samples (and corresponding survival data). The function computes a meta-estimator for the correlations between the two scores and uses a Student t test for dependent samples.

### Value

<code>p.value</code>	p-value from the Student t test for the comparison <code>cindex1 &gt; cindex2</code> .
<code>cindex1</code>	meta-estimator of the first concordance index.
<code>cindex2</code>	meta-estimator of the second concordance index.

### Author(s)

Benjamin Haibe-Kains

### References

Cochrane, W. G. (1954) "The combination of estimates from different experiments", *Biometrics*, **10**, pages 101–129.

Haibe-Kains, B. and Desmedt, C. and Sotiriou, C. and Bontempi, G. (2008) "A comparative study of survival models for breast cancer prognostication based on microarray data: does a single gene beat them all?", *Bioinformatics*, **24**, 19, pages 2200–2208.

### See Also

[concordance.index](#).



**Examples**

```

#first dataset
set.seed(12345)
age <- rnorm(100, 50, 10)
size <- rexp(100,1)
stime <- rexp(100)
cens <- runif(100,.5,2)
sevent <- as.numeric(stime <= cens)
stime <- pmin(stime, cens)
c1.1 <- concordance.index(x=age, surv.time=stime, surv.event=sevent,
  method="noether")
c2.1 <- concordance.index(x=size, surv.time=stime, surv.event=sevent,
  method="noether")
#second dataset
set.seed(54321)
age <- rnorm(110, 53, 10)
size <- rexp(110,1.1)
stime <- rexp(110)
cens <- runif(110,.55,2)
sevent <- as.numeric(stime <= cens)
stime <- pmin(stime, cens)
c1.2 <- concordance.index(x=age, surv.time=stime, surv.event=sevent,
  method="noether")
c2.2 <- concordance.index(x=size, surv.time=stime, surv.event=sevent,
  method="noether")
cindex.comp.meta(list.cindex1=list("cindex.age1"=c1.1, "cindex.age2"=c1.2),
  list.cindex2=list("cindex.size1"=c2.1, "cindex.size2"=c2.2))

```

combine.est

*Function to combine estimates***Description**

The function combines several estimators using meta-analytical formula to compute a meta-estimate.

**Usage**

```
combine.est(x, x.se, hetero = FALSE, na.rm = FALSE)
```

**Arguments**

x	vector of estimates
x.se	vector of standard errors of the corresponding estimates
hetero	TRUE is the heterogeneity should be taken into account (random effect model), FALSE otherwise (fixed effect model)
na.rm	TRUE if the missing values should be removed from the data, FALSE otherwise

**Value**

estimate	meta-estimate
se	standard error of the meta-estimate

**Author(s)**

Benjamin Haibe-Kains

**References**

Cochrane, W. G. (1954) "The combination of estimates from different experiments", *Biometrics*, **10**, pages 101–129.

**See Also**

test.hetero.est

**Examples**

```
set.seed(12345)
x1 <- rnorm(100, 50, 10) + rnorm(100, 0, 2)
m1 <- mean(x1)
se1 <- sqrt(var(x1))
x2 <- rnorm(100, 75, 15) + rnorm(100, 0, 5)
m2 <- mean(x2)
se2 <- sqrt(var(x2))

#fixed effect model
combine.est(x=c(m1, m2), x.se=c(se1, se2), hetero=FALSE)
#random effect model
combine.est(x=c(m1, m2), x.se=c(se1, se2), hetero=TRUE)
```

---

combine.test

*Function to combine probabilities*

---

**Description**

The function combines several p-value estimated from the same null hypothesis in different studies involving independent data.

**Usage**

```
combine.test(p, weight, method = c("fisher", "z.transform", "logit"),
  hetero = FALSE, na.rm = FALSE)
```

**Arguments**

p	vector of p-values
weight	vector of weights (e.g. sample size of each study)
method	fisher for the Fisher's combined probability test, z.transform for the Z-transformed test, logit for the weighted Z-method
hetero	TRUE is the heterogeneity should be taken into account, FALSE otherwise
na.rm	TRUE if the missing values should be removed from the data, FALSE otherwise

**Details**

The p-values must be one-sided and computed from the same null hypothesis.

**Value**

p-value

**Author(s)**

Benjamin Haibe-Kains

**References**

Whitlock, M. C. (2005) "Combining probability from independent tests: the weighted Z-method is superior to Fisher's approach", *J. Evol. Biol.*, **18**, pages 1368–1373.

**See Also**

test.hetero.test

**Examples**

```
p <- c(0.01, 0.13, 0.07, 0.2)
w <- c(100, 50, 200, 30)

#with equal weights
combine.test(p=p, method="z.transform")
#with p-values weighted by the sample size of the studies
combine.test(p=p, weight=w, method="z.transform")
```

---

concordance.index	<i>Function to compute the concordance index for survival or binary class prediction</i>
-------------------	--

---

**Description**

Function to compute the concordance index for a risk prediction, i.e. the probability that, for a pair of randomly chosen comparable samples, the sample with the higher risk prediction will experience an event before the other sample or belongs to a higher binary class.

**Usage**

```
concordance.index(x, surv.time, surv.event, cl, weights, comppairs=10, strat, alpha = 0.05, outx = TRUE, meth
```

**Arguments**

x	a vector of risk predictions.
surv.time	a vector of event times.
surv.event	a vector of event occurrence indicators.
cl	a vector of binary class indicators.
weights	weight of each sample.

comppairs	threshold for comparable patients.
strat	stratification indicator.
alpha	alpha level to compute confidence interval.
outx	set to TRUE to not count pairs of observations tied on x as a relevant pair. This results in a Goodman-Kruskal gamma type rank correlation.
method	can take the value conservative, noether or name (see paper Pencina et al. for details).
alternative	a character string specifying the alternative hypothesis, must be one of "two.sided" (default), "greater" (concordance index is greater than 0.5) or "less" (concordance index is less than 0.5). You can specify just the initial letter.
na.rm	TRUE if missing values should be removed.

### Value

c.index	concordance index estimate.
se	standard error of the estimate.
lower	lower bound for the confidence interval.
upper	upper bound for the confidence interval.
p.value	p-value for the statistical test if the estimate is different from 0.5.
n	number of samples used for the estimation.
data	list of data used to compute the index (x, surv.time and surv.event, or cl).
comppairs	number of comparable pairs.

### Note

The "direction" of the concordance index ( $< 0.5$  or  $> 0.5$ ) is the opposite than the [rcorr.cens](#) function in the Hmisc package. So you can easily get the same results than [rcorr.cens](#) by changing the sign of x.

### Author(s)

Benjamin Haibe-Kains, Markus Schroeder

### References

Harrel Jr, F. E. and Lee, K. L. and Mark, D. B. (1996) "Tutorial in biostatistics: multivariable prognostic models: issues in developing models, evaluating assumptions and adequacy, and measuring and reducing error", *Statistics in Medicine*, **15**, pages 361–387.

Pencina, M. J. and D'Agostino, R. B. (2004) "Overall C as a measure of discrimination in survival analysis: model specific population value and confidence interval estimation", *Statistics in Medicine*, **23**, pages 2109–2123, 2004.

### See Also

[rcorr.cens](#), [phcpe](#), [coxphCPE](#)

**Examples**

```

set.seed(12345)
age <- rnorm(100, 50, 10)
sex <- sample(0:1, 100, replace=TRUE)
stime <- rexp(100)
cens <- runif(100,.5,2)
sevent <- as.numeric(stime <= cens)
stime <- pmin(stime, cens)
strat <- sample(1:3, 100, replace=TRUE)
weight <- runif(100, min=0, max=1)
comppairs <- 10
cat("survival prediction:\n")
concordance.index(x=age, surv.time=stime, surv.event=sevent, strat=strat,
  weights=weight, method="noether", comppairs=comppairs)
cat("binary class prediction:\n")
## is age predictive of sex?
concordance.index(x=age, cl=sex, strat=strat, method="noether")

```

---

cvpl

*Function to compute the CVPL*


---

**Description**

The function computes the cross-validated partial likelihood (CVPL) for the Cox model.

**Usage**

```

cvpl(x, surv.time, surv.event, strata, nfold = 1, setseed,
  na.rm = FALSE, verbose = FALSE)

```

**Arguments**

x	data matrix
surv.time	vector of times to event occurrence
surv.event	vector of indicators for event occurrence
strata	stratification variable
nfold	number of folds for the cross-validation
setseed	seed for the random generator
na.rm	TRUE if the missing values should be removed from the data, FALSE otherwise
verbose	verbosity of the function

**Value**

cvpl	mean cross-validated partial likelihood (lower is better)
pl	vector of cross-validated partial likelihoods
convergence	vector of booleans reporting the convergence of the Cox model in each fold
n	number of observations used to estimate the cross-validated partial likelihood

**Author(s)**

Benjamin Haibe-Kains

**References**

Verweij PJM. and van Houwelingen H (1993) "Cross-validation in survival analysis", *Statistics in Medicine*, **12**, pages 2305–2314

van Houwelingen H, Bruinsma T, Hart AA, van't Veer LJ, and Wessels LFA (2006) "Cross-validated Cox regression on microarray gene expression data", *Statistics in Medicine*, **25**, pages 3201–3216.

**See Also**

[logpl](#), [coxph](#)

**Examples**

```
set.seed(12345)
age <- rnorm(100, 50, 10)
stime <- rexp(100)
cens <- runif(100,.5,2)
sevent <- as.numeric(stime <= cens)
stime <- pmin(stime, cens)
strat <- sample(1:3, 100, replace=TRUE)
cvpl(x=age, surv.time=stime, surv.event=sevent, strata=strat,
     nfold=10, setseed=54321)
```

---

D.index

*Function to compute the D index*

---

**Description**

Function to compute the D index for a risk prediction, i.e. an estimate of the log hazard ratio comparing two equal-sized prognostic groups. This is a natural measure of separation between two independent survival distributions under the proportional hazards assumption.

**Usage**

```
D.index(x, surv.time, surv.event, weights, strat, alpha = 0.05, method.test = c("logrank", "likelihood.ratio", "w"))
```

**Arguments**

x	a vector of risk predictions.
surv.time	a vector of event times.
surv.event	a vector of event occurrence indicators.
weights	weight of each sample.
strat	stratification indicator.
alpha	alpha level to compute confidence interval.
method.test	Statistical test to use in order to compute the p-values related to a D. index, see <a href="#">summary.coxph</a> for more details.
na.rm	TRUE if missing values should be removed.
...	additional parameters to be passed to the <a href="#">coxph</a> function.

**Details**

The D index is computed using the Cox model fitted on the scaled rankits of the risk scores instead of the risk scores themselves. The scaled rankits are the expected standard Normal order statistics scaled by  $\kappa = \sqrt{8/\pi}$ . See (Royston and Sauerbrei, 2004) for details.

Note that the value D reported in (Royston and Sauerbrei, 2004) is given

**Value**

d.index	D index estimate.
se	standard error of the estimate.
lower	lower bound for the confidence interval.
upper	upper bound for the confidence interval.
p.value	p-value for the statistical test if the estimate is different from 0.5.
n	number of samples used for the estimation.
coxmodel	<a href="#">coxph.object</a> fitted on the survival data and z (see below).
data	list of data used to compute the index (x, z, surv.time and surv.event). The item z contains the scaled rankits which are the expected standard Normal order statistics scaled by kappa.

**Author(s)**

Benjamin Haibe-Kains

**References**

Royston, P. and Sauerbrei, W. (2004) "A new measure of prognostic separation in survival data", *Statistics in Medicine*, **23**, pages 723–748.

**See Also**

[coxph](#), [coxph.object](#), [normOrder](#)

**Examples**

```
set.seed(12345)
age <- rnorm(100, 50, 10)
stime <- rexp(100)
cens <- runif(100,.5,2)
sevent <- as.numeric(stime <= cens)
stime <- pmin(stime, cens)
strat <- sample(1:3, 100, replace=TRUE)
weight <- runif(100, min=0, max=1)
D.index(x=age, surv.time=stime, surv.event=sevent, weights=weight, strat=strat)
```

---

`dindex.comp`*Function to compare two D indices*

---

### Description

This function compares two D indices from their betas and standard errors as computed by a Cox model for instance. The statistical test is a Student t test for dependent samples. The two D indices must be computed using the [D.index](#) function from the same survival data.

### Usage

```
dindex.comp(dindex1, dindex2)
```

### Arguments

<code>dindex1</code>	first D index
<code>dindex2</code>	second D index

### Details

The two D indices must be computed using the [D.index](#) function from the same samples (and corresponding survival data). The function uses a Student t test for dependent samples.

### Value

<code>p.value</code>	p-value from the Wilcoxon rank sum test for the comparison <code>dindex1 &gt; dindex2</code>
<code>dindex1</code>	value of the first D index
<code>dindex2</code>	value of the second D index

### Author(s)

Benjamin Haibe-Kains

### References

Haibe-Kains, B. and Desmedt, C. and Sotiriou, C. and Bontempi, G. (2008) "A comparative study of survival models for breast cancer prognostication based on microarray data: does a single gene beat them all?", *Bioinformatics*, **24**, 19, pages 2200–2208.

### See Also

[D.index](#), [t.test](#)

### Examples

```
set.seed(12345)
age <- rnorm(100, 50, 10)
size <- rexp(100,1)
stime <- rexp(100)
cens <- runif(100,.5,2)
sevent <- as.numeric(stime <= cens)
stime <- pmin(stime, cens)
```



```
d1 <- D.index(x=age, surv.time=stime, surv.event=sevent)
d2 <- D.index(x=size, surv.time=stime, surv.event=sevent)
dindex.comp(d1, d2)
```

---

dindex.comp.meta      *Function to compare two D indices*

---

### Description

This function compares two lists of D indices computed from the same survival data by using the function `D.index`. The statistical test is a Student t test for dependent samples.

### Usage

```
dindex.comp.meta(list.dindex1, list.dindex2, hetero = FALSE)
```

### Arguments

<code>list.dindex1</code>	first list of D indices as returned by the <code>D.index</code> function.
<code>list.dindex2</code>	second list of D indices as returned by the <code>D.index</code> function.
<code>hetero</code>	if TRUE, a random effect model is use to compute the meta-estimators. Otherwise a fixed effect model is used.

### Details

In meta-analysis, we estimate the statistic of interest in several independent datasets. It results a list of estimates such as list of D indices. The two lists of D indices must be computed from the same samples (and corresponding survival data). The function computes a meta-estimator for the correlations between the two scores and uses a Student t test for dependent samples.

### Value

<code>p.value</code>	p-value from the Student t test for the comparison <code>dindex1 &gt; dindex2</code> .
<code>dindex1</code>	meta-estimator of the first D index.
<code>dindex2</code>	meta-estimator of the second D index.

### Author(s)

Benjamin Haibe-Kains

### References

Cochrane, W. G. (1954) "The combination of estimates from different experiments", *Biometrics*, **10**, pages 101–129.

Haibe-Kains, B. and Desmedt, C. and Sotiriou, C. and Bontempi, G. (2008) "A comparative study of survival models for breast cancer prognostication based on microarray data: does a single gene beat them all?", *Bioinformatics*, **24**, 19, pages 2200–2208.

### See Also

[concordance.index](#).

**Examples**

```

#first dataset
set.seed(12345)
age <- rnorm(100, 50, 10)
size <- rexp(100,1)
stime <- rexp(100)
cens <- runif(100,.5,2)
sevent <- as.numeric(stime <= cens)
stime <- pmin(stime, cens)
d1.1 <- D.index(x=age, surv.time=stime, surv.event=sevent)
d2.1 <- D.index(x=size, surv.time=stime, surv.event=sevent)
#second dataset
set.seed(54321)
age <- rnorm(110, 53, 10)
size <- rexp(110,1.1)
stime <- rexp(110)
cens <- runif(110,.55,2)
sevent <- as.numeric(stime <= cens)
stime <- pmin(stime, cens)
d1.2 <- D.index(x=age, surv.time=stime, surv.event=sevent)
d2.2 <- D.index(x=size, surv.time=stime, surv.event=sevent)
dindex.comp.meta(list.dindex1=list("dindex.age1"=d1.1, "dindex.age2"=d1.2),
  list.dindex2=list("dindex.size1"=d2.1, "dindex.size2"=d2.2))

```

fisherz

*Function to compute Fisher z transformation***Description**

The function computes the Fisher z transformation useful to calculate the confidence interval of Pearson's correlation coefficient.

**Usage**

```
fisherz(x, inv = FALSE, eps = 1e-16)
```

**Arguments**

x	value, e.g. Pearson's correlation coefficient
inv	TRUE for inverse Fisher z transformation, FALSE otherwise
eps	tolerance for extreme cases, i.e.

*latex*

when inv = FALSE and

*latex*

when inv = TRUE

**Details**

The sampling distribution of Pearson's *latex* is not normally distributed. R. A. Fisher developed a transformation now called "Fisher's z transformation" that converts Pearson's *latex* to the normally distributed variable z. The formula for the transformation is

$$latex$$

Two attributes of the distribution of the z statistic: (1) It is normally distributed and (2) it has a known standard error of

$$latex$$

where *latex* is the number of samples.

Fisher's z is used for computing confidence intervals on Pearson's correlation and for confidence intervals on the difference between correlations.

**Value**

Fisher's z statistic

**Author(s)**

Benjamin Haibe-Kains

**References**

R. A. Fisher (1915) "Frequency distribution of the values of the correlation coefficient in samples of an indefinitely large population". *Biometrika*, **10**, pages 507–521.

**See Also**

[cor](#)

**Examples**

```
set.seed(12345)
x1 <- rnorm(100, 50, 10)
x2 <- runif(100,.5,2)
cc <- cor(x1, x2)
z <- fisherz(x=cc, inv=FALSE)
z.se <- 1 / sqrt(100 - 3)
fisherz(z, inv=TRUE)
```

---

forestplot.surv

*Forest plots enables to display performance estimates of survival models*

---

**Description**

Draw a forest plot together with a table of text.

**Usage**

```
forestplot.surv(labeltext, mean, lower, upper, align = NULL,
  is.summary = FALSE, clip = c(-Inf, Inf), xlab = "", zero = 0,
  graphwidth = unit(2, "inches"), col, xlog = FALSE,
  box.size = NULL, x.ticks = NULL, ...)
```

**Arguments**

labeltext	Matrix of strings or NAs for blank spaces
mean	Vector of centers of confidence intervals (or NAs for blank space)
lower	Vector of lower ends of confidence intervals
upper	Vector of upper ends of confidence intervals
align	Vector giving alignment (l,r,c) for columns of table
is.summary	Vector of logicals. Summary lines have bold text and diamond confidence intervals.
clip	Lower and upper limits for clipping confidence intervals to arrows
xlab	x-axis label
zero	x-axis coordinate for zero line
graphwidth	Width of confidence interval graph
col	See <a href="#">meta.colors</a>
xlog	If TRUE, x-axis tick marks are exponentiated
box.size	Override the default box size based on precision
x.ticks	Optional user-specified x-axis tick marks. Specify NULL to use the defaults, numeric(0) to omit the x-axis.
...	Not used.

**Details**

This function is more flexible than [metaplot](#) and the plot methods for meta-analysis objects, but requires more work by the user.

In particular, it allows for a table of text, and clips confidence intervals to arrows when they exceed specified limits.

**Value**

None

**References**

rmeta package, CRAN, Thomas Lumley <tlumley@u.washington.edu>. Functions for simple fixed and random effects meta-analysis for two-sample comparisons and cumulative meta-analyses. Draws standard summary plots, funnel plots, and computes summaries and tests for association and heterogeneity.

**See Also**

[metaplot](#), [forestplot](#)

**Examples**

```

require(rmeta)
myspace <- "  "
labeltext <- cbind(c("Gene Symbol", "AAA", "BBB", "CCC"),c(rep(myspace,4)))
bs <- rep(0.5, nrow(labeltext))
r.mean <- c(NA, 0.35, 0.5, 0.65)
r.lower <- c(NA, 0.33, 0.4, 0.6)
r.upper <- c(NA, 0.37, 0.6, 0.7)

forestplot.surv(labeltext=labeltext, mean=r.mean, lower=r.lower, upper=r.upper, zero=0.5,
  align=c("l"), graphwidth=unit(2, "inches"), x.ticks=seq(0.3,0.8,0.1), xlab=paste( "Forestplot Example", myspace, sep=" ",
  col=meta.colors(box="royalblue", line="darkblue", zero="darkred"), box.size=bs, clip=c(0.3,0.8))

```

---

getsurv2	<i>Function to retrieve the survival probabilities at a specific point in time</i>
----------	--

---

**Description**

The function retrieves the survival probabilities from a survfit object, for a specific point in time.

**Usage**

```
getsurv2(sf, time, which.est = c("point", "lower", "upper"))
```

**Arguments**

sf	survfit object
time	time at which the survival probabilities must be retrieved
which.est	which estimation to be returned? point for the point estimate, lower for the lower bound and upper for the upper bound

**Details**

The survival probabilities are estimated through the [survfit](#) function.

**Value**

vector of survival probabilities

**Author(s)**

Benjamin Haibe-Kains

**See Also**

[survfit](#)

**Examples**

```
require(survival)
set.seed(12345)
age <- rnorm(30, 50, 10)
stime <- rexp(30)
cens <- runif(30,.5,2)
sevent <- as.numeric(stime <= cens)
stime <- pmin(stime, cens)
sf <- survfit(Surv(stime, sevent) ~ 1)
getsurv2(sf, time=1)
```

hazard.ratio

*Function to estimate the hazard ratio through Cox regression***Description**

Function to compute the hazard ratio for a risk prediction.

**Usage**

```
hazard.ratio(x, surv.time, surv.event, weights, strat, alpha = 0.05, method.test = c("logrank", "likelihood.ratio"))
```

**Arguments**

x	a vector of risk predictions.
surv.time	a vector of event times.
surv.event	a vector of event occurrence indicators.
weights	weight of each sample.
strat	stratification indicator.
alpha	alpha level to compute confidence interval.
method.test	Statistical test to use in order to compute the p-values related to a D. index, see <a href="#">summary.coxph</a> for more details.
na.rm	TRUE if missing values should be removed.
...	additional parameters to be passed to the <a href="#">coxph</a> function.

**Details**

The hazard ratio is computed using the Cox model.

**Value**

hazard.ratio	hazard ratio estimate.
coef	coefficient (beta) estimated in the cox regression model.
se	standard error of the coefficient (beta) estimate.
lower	lower bound for the confidence interval.
upper	upper bound for the confidence interval.
p.value	p-value computed using the likelihood ratio test whether the hazard ratio is different from 1.

n                    number of samples used for the estimation.  
coxnm                [coxph.object](#) fitted on the survival data and x (see below).  
data                 list of data used to compute the hazard ratio (x, surv.time and surv.event).

**Author(s)**

Benjamin Haibe-Kains

**References**

Cox, D. R. (1972) "Regression Models and Life Tables", *Journal of the Royal Statistical Society Series B*, **34**, pages 187–220.

**See Also**

[coxph](#), [coxph.object](#)

**Examples**

```
set.seed(12345)
age <- rnorm(100, 50, 10)
stime <- rexp(100)
cens <- runif(100,.5,2)
sevent <- as.numeric(stime <= cens)
stime <- pmin(stime, cens)
strat <- sample(1:3, 100, replace=TRUE)
weight <- runif(100, min=0, max=1)
hazard.ratio(x=age, surv.time=stime, surv.event=sevent, weights=weight,
             strat=strat)
```

---

hr.comp

*Function to statistically compare two hazard ratios*

---

**Description**

This function compares two hazard ratios from their betas and standard errors as computed by a Cox model for instance. The statistical test is a Student t test for dependent samples. The two hazard ratios must be computed from the same survival data.

**Usage**

```
hr.comp(hr1, hr2)
```

**Arguments**

hr1                 first hazard ratio.  
hr2                 second hazard ratio.

**Details**

The two hazard ratios must be computed from the same samples (and corresponding survival data). The function uses a Student t test for dependent samples.

**Value**

p.value	p-value from the Student t test for the comparison $\beta_1 > \beta_2$ (equivalently $hr_1 > hr_2$ )
hr1	value of the first hazard ratio
hr2	value of the second hazard ratio

**Author(s)**

Benjamin Haibe-Kains

**References**

Student 1908) "The Probable Error of a Mean", *Biometrika*, **6**, 1, pages 1–25.

Haibe-Kains, B. and Desmedt, C. and Sotiriou, C. and Bontempi, G. (2008) "A comparative study of survival models for breast cancer prognostication based on microarray data: does a single gene beat them all?", *Bioinformatics*, **24**, 19, pages 2200–2208.

**See Also**

[coxph](#), [t.test](#)

**Examples**

```
require(survival)
set.seed(12345)
age <- as.numeric(rnorm(100, 50, 10) >= 50)
size <- as.numeric(rexp(100,1) > 1)
stime <- rexp(100)
cens <- runif(100,.5,2)
sevent <- as.numeric(stime <= cens)
stime <- pmin(stime, cens)
hr1 <- hazard.ratio(x=age, surv.time=stime, surv.event=sevent)
hr2 <- hazard.ratio(x=size, surv.time=stime, surv.event=sevent)
hr.comp(hr1=hr1, hr2=hr2)
```

---

hr.comp.meta

*Function to compare two concordance indices*

---

**Description**

This function compares two lists of hazard ratios computed from the same survival data by using the function `hazard.ratio`. The statistical test is a Student t test for dependent samples.

**Usage**

```
hr.comp.meta(list.hr1, list.hr2, hetero = FALSE)
```

**Arguments**

list.hr1	first list of D indices as returned by the <code>hazard.ratio</code> function.
list.hr2	second list of D indices as returned by the <code>hazard.ratio</code> function.
hetero	if TRUE, a random effect model is use to compute the meta-estimators. Otherwise a fixed effect model is used.



## Details

In meta-analysis, we estimate the statistic of interest in several independent datasets. It results a list of estimates such as list of hazard ratios. The two lists of hazard ratios must be computed from the same samples (and corresponding survival data). The function computes a meta-estimator for the correlations between the two scores and uses a Student t test for dependent samples.

## Value

p.value	p-value from the Student t test for the comparison $hr1 > hr2$ .
hr1	meta-estimator of the first D index.
hr2	meta-estimator of the second D index.

## Author(s)

Benjamin Haibe-Kains

## References

Cochrane, W. G. (1954) "The combination of estimates from different experiments", *Biometrics*, **10**, pages 101–129.

Haibe-Kains, B. and Desmedt, C. and Sotiriou, C. and Bontempi, G. (2008) "A comparative study of survival models for breast cancer prognostication based on microarray data: does a single gene beat them all?", *Bioinformatics*, **24**, 19, pages 2200–2208.

## See Also

[concordance.index](#).

## Examples

```
#first dataset
set.seed(12345)
age <- rnorm(100, 50, 10)
size <- rexp(100,1)
stime <- rexp(100)
cens <- runif(100,.5,2)
sevent <- as.numeric(stime <= cens)
stime <- pmin(stime, cens)
h1.1 <- hazard.ratio(x=age, surv.time=stime, surv.event=sevent)
h2.1 <- hazard.ratio(x=size, surv.time=stime, surv.event=sevent)
#second dataset
set.seed(54321)
age <- rnorm(110, 53, 10)
size <- rexp(110,1.1)
stime <- rexp(110)
cens <- runif(110,.55,2)
sevent <- as.numeric(stime <= cens)
stime <- pmin(stime, cens)
h1.2 <- hazard.ratio(x=age, surv.time=stime, surv.event=sevent)
h2.2 <- hazard.ratio(x=size, surv.time=stime, surv.event=sevent)
hr.comp.meta(list.hr1=list("hr.age1"=h1.1, "hr.age2"=h1.2),
  list.hr2=list("hr.size1"=h2.1, "hr.size2"=h2.2))
```

hr.comp2

*Function to statistically compare two hazard ratios (alternative interface)***Description**

This function compares two hazard ratios from their betas and standard errors as computed by a Cox model for instance. The statistical test is a Student t test for dependent samples. The two hazard ratios must be computed from the same survival data.

**Usage**

```
hr.comp2(x1, beta1, se1, x2, beta2, se2, n)
```

**Arguments**

x1	risk score used to estimate the first hazard ratio.
beta1	beta estimate for the first hazard ratio.
se1	standard error of beta estimate for the first hazard ratio.
x2	risk score used to estimate the second hazard ratio.
beta2	beta estimate for the second hazard ratio.
se2	standard error of beta estimate for the first hazard ratio.
n	number of samples from which the hazard ratios were estimated.

**Details**

The two hazard ratios must be computed from the same samples (and corresponding survival data). The function uses a Student t test for dependent samples.

**Value**

p.value	p-value from the Student t test for the comparison $\beta_1 > \beta_2$ (equivalently $hr_1 > hr_2$ )
hr1	value of the first hazard ratio
hr2	value of the second hazard ratio

**Author(s)**

Benjamin Haibe-Kains

**References**

Student 1908) "The Probable Error of a Mean", *Biometrika*, **6**, 1, pages 1–25.  
 Haibe-Kains, B. and Desmedt, C. and Sotiriou, C. and Bontempi, G. (2008) "A comparative study of survival models for breast cancer prognostication based on microarray data: does a single gene beat them all?", *Bioinformatics*, **24**, 19, pages 2200–2208.

**See Also**

[coxph](#), [t.test](#)

**Examples**

```

require(survival)
set.seed(12345)
age <- as.numeric(rnorm(100, 50, 10) >= 50)
size <- as.numeric(rexp(100,1) > 1)
stime <- rexp(100)
cens <- runif(100,.5,2)
sevent <- as.numeric(stime <= cens)
stime <- pmin(stime, cens)
cox1 <- coxph(Surv(stime, sevent) ~ age)
cox2 <- coxph(Surv(stime, sevent) ~ size)
hr.comp2(x1=age, beta1=cox1$coefficients, se1=drop(sqrt(cox1$var)),
         x2=size, beta2=cox2$coefficients, se2=drop(sqrt(cox2$var)), n=100)

```

iauc.comp

*Function to compare two IAUCs through time-dependent ROC curves***Description**

This function compares two integrated areas under the curves (IAUC) through the results of time-dependent ROC curves at some points in time. The statistical test is a Wilcoxon rank sum test for dependent samples.

**Usage**

```
iauc.comp(auc1, auc2, time)
```

**Arguments**

auc1	vector of AUCs computed from the first time-dependent ROC curves for some points in time
auc2	vector of AUCs computed from the second time-dependent ROC curves for some points in time
time	vector of points in time for which the AUCs are computed

**Details**

The two vectors of AUCs must be computed from the same samples (and corresponding survival data) and for the same points in time. The function uses a Wilcoxon rank sum test for dependent samples.

**Value**

p.value	p-value from the Wilcoxon rank sum test for the comparison <code>iauc1 &gt; iauc2</code>
iauc1	value of the IAUC for the first set of time-depdent ROC curves
iauc2	value of the IAUC for the second set of time-depdent ROC curves

**Author(s)**

Benjamin Haibe-Kains

## References

Wilcoxon, F. (1945) "Individual comparisons by ranking methods", *Biometrics Bulletin*, **1**, pages 80–83.

Haibe-Kains, B. and Desmedt, C. and Sotiriou, C. and Bontempi, G. (2008) "A comparative study of survival models for breast cancer prognostication based on microarray data: does a single gene beat them all?", *Bioinformatics*, **24**, 19, pages 2200–2208.

## See Also

[tdrocc](#), [wilcox.test](#)

## Examples

```
set.seed(12345)
age <- rnorm(30, 50, 10)
size <- rexp(30,1)
stime <- rexp(30)
cens <- runif(30,.5,2)
sevent <- as.numeric(stime <= cens)
stime <- pmin(stime, cens)
##time-dependent ROC curves
tt <- unique(sort(stime[sevent == 1]))
##size
mytdroc1 <- NULL
for(i in 1:length(tt)) {
  rr <- tdrocc(x=size, surv.time=stime, surv.event=sevent, time=tt[i],
    na.rm=TRUE, verbose=FALSE)
  mytdroc1 <- c(mytdroc1, list(rr))
}
auc1 <- unlist(lapply(mytdroc1, function(x) { return(x$AUC) }))
##age
mytdroc2 <- NULL
for(i in 1:length(tt)) {
  rr <- tdrocc(x=age, surv.time=stime, surv.event=sevent, time=tt[i],
    na.rm=TRUE, verbose=FALSE)
  mytdroc2 <- c(mytdroc2, list(rr))
}
auc2 <- unlist(lapply(mytdroc2, function(x) { return(x$AUC) }))
iauc.comp(auc1=auc1, auc2=auc2, time=tt)
```

---

ibsc.comp

*Function to compare two IBSCs*

---

## Description

This function compares two integrated Briers scores (IBSC) through the estimation of the Brier scores (BSC) at some points in time. The statistical test is a Wilcoxon rank sum test for dependent samples.

## Usage

```
ibsc.comp(bsc1, bsc2, time)
```

**Arguments**

bsc1	vector of BSCs computed from the first predicted probabilities for some points in time
bsc2	vector of BSCs computed from the second predicted probabilities for some points in time
time	vector of points in time for which the BSCs are computed

**Details**

The two vectors of BSCs must be computed from the same samples (and corresponding survival data) and for the same points in time. The function uses a Wilcoxon rank sum test for dependent samples.

**Value**

p.value	p-value from the Wilcoxon rank sum test for the comparison $\text{ibsc1} < \text{ibsc2}$
ibsc1	value of the IBSC for the first set of BSCs
ibsc2	value of the IBSC for the second set of BSCs

**Author(s)**

Benjamin Haibe-Kains

**References**

Wilcoxon, F. (1945) "Individual comparisons by ranking methods", *Biometrics Bulletin*, **1**, pages 80–83.

Haibe-Kains, B. and Desmedt, C. and Sotiriou, C. and Bontempi, G. (2008) "A comparative study of survival models for breast cancer prognostication based on microarray data: does a single gene beat them all?", *Bioinformatics*, **24**, 19, pages 2200–2208.

**See Also**

[sbrier.score2proba](#), [sbrier](#)

**Examples**

```
set.seed(12345)
age <- rnorm(30, 50, 10)
size <- rexp(30,1)
stime <- rexp(30)
cens <- runif(30,.5,2)
sevent <- as.numeric(stime <= cens)
stime <- pmin(stime, cens)
##Brier scores
##size
dd <- data.frame("time"=stime, "event"=sevent, "score"=size)
bsc1 <- sbrier.score2proba(data.tr=dd, data.ts=dd, method="cox")
##size
dd <- data.frame("time"=stime, "event"=sevent, "score"=age)
bsc2 <- sbrier.score2proba(data.tr=dd, data.ts=dd, method="cox")
if(!all(bsc1$time == bsc2$time)) {
  stop("the two vector of BSCs must be computed for the same points in time!") }
ibsc.comp(bsc1=bsc1$bsc, bsc2=bsc2$bsc, time=bsc1$time)
```

km.coxph.plot

*Function to plot several Kaplan-Meier survival curves***Description**

Function to plot several Kaplan-Meier survival curves with number of individuals at risk at some time points.

**Usage**

```
km.coxph.plot(formula.s, data.s, sub.s = "all", x.label, y.label, main.title, sub.title, leg.text, leg.pos = "bottomright")
```

**Arguments**

formula.s	formula composed of a Surv object and a strata variable (i.e. stratification).
data.s	data frame composed of the variables used in the formula.
sub.s	vector of booleans specifying if only a subset of the data should be considered. Default value is "all".
x.label	label for the y-axis.
y.label	label for the x-axis.
main.title	main title at the top of the plot.
sub.title	subtitle at the bottom of the plot.
leg.text	text in the legend.
leg.pos	the location may also be specified by setting 'x' to a single keyword from the list "bottomright", "bottom", "bottomleft", "left", "topleft", "top", "topright", "right" and "center". This places the legend on the inside of the plot frame at the given location.
leg.bty	the type of box to be drawn around the legend. The allowed values are "o" (the default) and "n".
leg.inset	inset distance from the margins as a fraction of the plot region. Default value is 0.05.
o.text	plot the logrank p-value.
v.line	x coordinate(s) for vertical line(s).
h.line	y coordinate(s) for horizontal line(s).
.col	vector of colors for the different survival curves.
.lty	vector of line types for the different survival curves
.lwd	vector of line widths for the different survival curves.
show.n.risk	if TRUE, show the numbers of samples at risk for each time step.
n.risk.step	vector specifying the time to be the steps for displaying the number of individuals at risk.
n.risk.cex	size of the number of individuals at risk. Default value is 0.85.
verbose	verbosity level (TRUE or FALSE). Default value is TRUE.
...	additional parameters to be passed to the <a href="#">plot</a> function.

**Details**

The original version of this function was kindly provided by Dr Christos Hatzis (January, 17th 2006).

**Author(s)**

Christos Hatzis, Benjamin Haibe-Kains

**See Also**

[survfit](#), [coxph](#)

**Examples**

```
set.seed(12345)
stime <- rexp(100) * 10
cens <- runif(100,.5,2) * 10
sevent <- as.numeric(stime <= cens)
stime <- pmin(stime, cens)
strat <- sample(1:3, 100, replace=TRUE)
dd <- data.frame("surv.time"=stime, "surv.event"=sevent, "strat"=strat)

km.coxph.plot(formula.s=Surv(surv.time, surv.event) ~ strat, data.s=dd,
  sub.s="all", x.label="Time (years)", y.label="Probability of survival",
  main.title="", leg.text=paste(c("Low", "Intermediate", "High"), " ", sep=""),
  leg.pos="topright", leg.inset=0, .col=c("darkblue", "darkgreen", "darkred"),
  .lty=c(1,1,1), show.n.risk=TRUE, n.risk.step=2, n.risk.cex=0.85, verbose=FALSE)
```

---

logpl

---

*Function to compute the log partial likelihood of a Cox model*


---

**Description**

The function computes the log partial likelihood of a set of coefficients given some survival data.

**Usage**

```
logpl(pred, surv.time, surv.event, strata, na.rm = FALSE, verbose = FALSE)
```

**Arguments**

surv.time	vector of times to event occurrence
surv.event	vector of indicators for event occurrence
pred	linear predictors computed using the Cox model
strata	stratification variable
na.rm	TRUE if the missing values should be removed from the data, FALSE otherwise
verbose	verbosity of the function

**Value**

vector of two elements: logpl and event for the estimation of the log partial likelihood and the number of events, respectively

**Author(s)**

Benjamin Haibe-Kains

**References**

Cox, D. R. (1972) "Regression Models and Life Tables", *Journal of the Royal Statistical Society Series B*, **34**, pages 187–220.

**See Also**

[coxph](#), [cvpl](#)

**Examples**

```
require(survival)
set.seed(12345)
age <- rnorm(100, 50, 10)
stime <- rexp(100)
cens <- runif(100,.5,2)
sevent <- as.numeric(stime <= cens)
stime <- pmin(stime, cens)
dd <- data.frame("stime"=stime, "sevent"=sevent, "age"=age)
##Cox model
coxM <- coxph(Surv(stime, sevent) ~ age, data=dd)
##log partial likelihood of the null model
logpl(pred=rep(0, nrow(dd)), surv.time=stime, surv.event=sevent)
##log partial likelihood of the Cox model
logpl(pred=predict(object=coxM, newdata=dd), surv.time=stime, surv.event=sevent)
##equivalent to
coxM$loglik
```

---

mainz7g

*Subset of MAINZ dataset containing gene expression, annotations and clinical data.*

---

**Description**

This dataset contains a subset of the gene expression, annotations and clinical data from the MAINZ datasets (see section details). The subset contains the seven genes introduced by Desmedt et al. 2008

**Format**

ExpressionSet with 7 features and 200 samples, containing:

- `exprs(mainz7g)`: Matrix containing gene expressions as measured by Affymetrix hgu133a technology (single-channel, oligonucleotides).
- `fData(mainz7g)`: AnnotatedDataFrame containing annotations of Affy microarray platform hgu133a.



- `pData(mainz7g)`: AnnotatedDataFrame containing Clinical information of the breast cancer patients whose tumors were hybridized.
- `experimentalData(mainz7g)`: MIAME object containing information about the dataset.
- `annotation(mainz7g)`: Name of the affy chip.

### Details

This dataset represents a subset of the study published by Schmidt et al. 2008. The subset contains the genes AURKA (also known as STK6, STK7, or STK15), PLAU (also known as UPA), STAT1, VEGF, CASP3, ESR1, and ERBB2, as introduced by Desmedt et al. 2008. The seven genes represent the proliferation, tumor invasion/metastasis, immune response, angiogenesis, apoptosis phenotypes, and the ER and HER2 signaling, respectively.

### Source

#### mainz:

<http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE11121>

### References

Marcus Schmidt, Daniel Boehm, Christian von Toerne, Eric Steiner, Alexander Puhl, Heryk Pilch, Hans-Anton Lehr, Jan G. Hengstler, Hainz Koelbl and Mathias Gehrman (2008) "The Humoral Immune System Has a Key Prognostic Impact in Node-Negative Breast Cancer", *Cancer Research*, **68**(13):5405-5413

### Examples

```
## load Biobase package
library(Biobase)
## load the dataset
data(breastCancerData)
## show the first 5 columns of the expression data
exprs(mainz7g)[,1:5]
## show the first 6 rows of the phenotype data
head(pData(mainz7g))
## show first 20 feature names
featureNames(mainz7g)
## show the experiment data summary
experimentData(mainz7g)
## show the used platform
annotation(mainz7g)
## show the abstract for this dataset
abstract(mainz7g)
```

---

metaplot.surv

*Meta-analysis plot (forest plot)*

---

### Description

Plot confidence intervals with boxes indicating the sample size/precision and optionally a diamond indicating a summary confidence interval. This function is usually called by plot methods for meta-analysis objects. Additionally, you can specify your own lower and upper boarder from the confidence interval.

**Usage**

```
metaplot.surv(mn, se=NULL, lower=NULL, upper=NULL, nn=NULL,
  labels=NULL, conf.level = .95, xlab = "", ylab = "", xlim = NULL,
  summn = NULL, sumse = NULL, sumlower = NULL, sumupper = NULL,
  sumnn = NULL, summlabel = "Summary", logeffect = FALSE,
  lwd = 2, boxsize = 1, zero = as.numeric(logeffect),
  colors, xaxt="s", logticks=TRUE, ... )
```

**Arguments**

mn	point estimates from studies
se	standard errors of mn
lower	Vector of lower ends of confidence intervals
upper	Vector of upper ends of confidence intervals
nn	precision: box area is proportional to this. $1/se^2$ is the default
labels	labels for each interval
conf.level	Confidence level for confidence intervals
xlab	label for the point estimate axis
ylab	label for the axis indexing the different studies
xlim	the range for the x axis.
summn	summary estimate
sumse	standard error of summary estimate
sumlower	lower end of confidence intervals of summary estimate
sumupper	upper end of confidence intervals of summary estimate
sumnn	precision of summary estimate
summlabel	label for summary estimate
logeffect	TRUE to display on a log scale
lwd	line width
boxsize	Scale factor for box size
zero	"Null" effect value
xaxt	use "n" for no x-axis (to add a customised one)
logticks	if TRUE and logscale, have tick values approximately equally spaced on a log scale.
colors	see <a href="#">meta.colors</a>
...	Other graphical parameters

**Value**

This function is used for its side-effect.

**See Also**

[forestplot.surv](#) for more flexible plots

**Examples**

```
metaplot.surv(mn=c(0.4,0.5,0.6), lower=c(0.35,0.4,0.57), upper=c(0.45,0.6,0.63), labels=c("A","B","C"), xlim=c(0.2,0.8))
```

---

mrmr.cindex	<i>Function to compute the concordance index for survival or binary class prediction</i>
-------------	--

---

### Description

Function to compute the minimum redundancy - maximum relevance (mRMR) ranking for a risk prediction or a binary classification task based on the concordance index. The mRMR feature selection has been adapted to use the concordance index to estimate the correlation between a variable and the output (binary or survival) data.

### Usage

```
mrmr.cindex(x, surv.time, surv.event, cl, weights, comppairs=10, strat, alpha = 0.05, outx = TRUE, method = c
```

### Arguments

x	a vector of risk predictions.
surv.time	a vector of event times.
surv.event	a vector of event occurrence indicators.
cl	a vector of binary class indicators.
weights	weight of each sample.
comppairs	threshold for comparable patients.
strat	stratification indicator.
alpha	alpha level to compute confidence interval.
outx	set to TRUE to not count pairs of observations tied on x as a relevant pair. This results in a Goodman-Kruskal gamma type rank correlation.
method	can take the value conservative, noether or name (see paper Pencina et al. for details).
alternative	a character string specifying the alternative hypothesis, must be one of "two.sided" (default), "greater" (concordance index is greater than 0.5) or "less" (concordance index is less than 0.5). You can specify just the initial letter.
na.rm	TRUE if missing values should be removed.

### Value

A mRMR ranking

### Note

The "direction" of the concordance index ( $< 0.5$  or  $> 0.5$ ) is the opposite than the [rcorr.cens](#) function in the Hmisc package. So you can easily get the same results than [rcorr.cens](#) by changing the sign of x.

### Author(s)

Benjamin Haibe-Kains, Markus Schroeder

## References

Harrel Jr, F. E. and Lee, K. L. and Mark, D. B. (1996) "Tutorial in biostatistics: multivariable prognostic models: issues in developing models, evaluating assumptions and adequacy, and measuring and reducing error", *Statistics in Medicine*, **15**, pages 361–387.

Pencina, M. J. and D'Agostino, R. B. (2004) "Overall C as a measure of discrimination in survival analysis: model specific population value and confidence interval estimation", *Statistics in Medicine*, **23**, pages 2109–2123, 2004.

## See Also

[rcorr.cens](#), [phepe](#), [coxphCPE](#)

## Examples

```
set.seed(12345)
age <- rnorm(100, 50, 10)
sex <- sample(0:1, 100, replace=TRUE)
stime <- rexp(100)
cens <- runif(100,.5,2)
sevent <- as.numeric(stime <= cens)
stime <- pmin(stime, cens)
strat <- sample(1:3, 100, replace=TRUE)
weight <- runif(100, min=0, max=1)
comppairs <- 10
xx <- data.frame("age"=age, "sex"=sex)
cat("survival prediction:\n")
mrmr.cindex(x=xx, surv.time=stime, surv.event=sevent, strat=strat, weights=weight, method="noether", comppairs=
```

---

mrmr.cindex.ensemble    *Function to compute the concordance index for survival or binary class prediction*

---

## Description

Function to compute the minimum redundancy - maximum relevance (mRMR) ranking for a risk prediction or a binary classification task based on the concordance index. The mRMR feature selection has been adapted to use the concordance index to estimate the correlation between a variable and the output (binary or survival) data.

## Usage

```
mrmr.cindex.ensemble(x, surv.time, surv.event, cl, weights, comppairs=10, strat, alpha = 0.05, outx = TRUE, m
```

## Arguments

x	a vector of risk predictions.
surv.time	a vector of event times.
surv.event	a vector of event occurrence indicators.
cl	a vector of binary class indicators.
weights	weight of each sample.

comppairs	threshold for comparable patients.
strat	stratification indicator.
alpha	alpha level to compute confidence interval.
outx	set to TRUE to not count pairs of observations tied on x as a relevant pair. This results in a Goodman-Kruskal gamma type rank correlation.
method	can take the value conservative, noether or name (see paper Pencina et al. for details).
alternative	a character string specifying the alternative hypothesis, must be one of "two.sided" (default), "greater" (concordance index is greater than 0.5) or "less" (concordance index is less than 0.5). You can specify just the initial letter.
maxparents	maximum number of candidate variables to be added in the ranking solutions tree.
maxnsol	maximum number of ranking solutions to be considered.
nboot	number of bootstraps to compute standard error of a ranking solution.
na.rm	TRUE if missing values should be removed.

**Value**

A mRMR ranking

**Note**

The "direction" of the concordance index ( $< 0.5$  or  $> 0.5$ ) is the opposite than the [rcorr.cens](#) function in the Hmisc package. So you can easily get the same results than [rcorr.cens](#) by changing the sign of x.

**Author(s)**

Benjamin Haibe-Kains, Markus Schroeder

**References**

Harrel Jr, F. E. and Lee, K. L. and Mark, D. B. (1996) "Tutorial in biostatistics: multivariable prognostic models: issues in developing models, evaluating assumptions and adequacy, and measuring and reducing error", *Statistics in Medicine*, **15**, pages 361–387.

Pencina, M. J. and D'Agostino, R. B. (2004) "Overall C as a measure of discrimination in survival analysis: model specific population value and confidence interval estimation", *Statistics in Medicine*, **23**, pages 2109–2123, 2004.

**See Also**

[rcorr.cens](#), [phcpe](#), [coxphCPE](#)

---

nki7g	<i>Subset of NKI dataset containing gene expression, annotations and clinical data.</i>
-------	---

---

### Description

This dataset contains a subset of the gene expression, annotations and clinical data from the NKI datasets (see section details). The subset contains the seven genes introduced by Desmedt et al. 2008

### Format

ExpressionSet with 7 features and 337 samples, containing:

- `exprs(nki7g)`: Matrix containing gene expressions as measured by Agilent technology (dual-channel, oligonucleotides).
- `fData(nki7g)`: AnnotatedDataFrame containing annotations of Agilent microarray platform.
- `pData(nki7g)`: AnnotatedDataFrame containing Clinical information of the breast cancer patients whose tumors were hybridized.
- `experimentalData(nki7g)`: MIAME object containing information about the dataset.
- `annotation(nki7g)`: Name of the agilent chip.

### Details

This dataset represents a subset of the study published by van't Veer et al. 2002 and van de Vijver et al. 2002. The subset contains the genes AURKA (also known as STK6, STK7, or STK15), PLAU (also known as UPA), STAT1, VEGF, CASP3, ESR1, and ERBB2, as introduced by Desmedt et al. 2008. The seven genes represent the proliferation, tumor invasion/metastasis, immune response, angiogenesis, apoptosis phenotypes, and the ER and HER2 signaling, respectively.

### Source

**nki:**

<http://www.rii.com/publications/2002/vantveer.html>

### References

Laura J. van't Veer, Hongyue Dai, Marc J. van de Vijver, Yudong D. He, Augustinus A.M. Hart, Mao Mao, Hans L. Peterse, Karin van der Kooy, Matthew J. Marton, Anke T. Witteveen, George J. Schreiber, Ron M. Kerkhoven, Chris Roberts, Peter S. Linsley, Rene Bernards and Stephen H. Friend (2002) "Gene expression profiling predicts clinical outcome of breast cancer", *Nature*, **415**:530-536

M. J. van de Vijver, Y. D. He, L. van't Veer, H. Dai, A. M. Hart, D. W. Voskuil, G. J. Schreiber, J. L. Peterse, C. Roberts, M. J. Marton, M. Parrish, D. Atsma, A. Witteveen, A. Glas, L. Delahaye, T. van der Velde, H. Bartelink, S. Rodenhuis, E. T. Rutgers, S. H. Friend and R. Bernards (2002) "A Gene Expression Signature as a Predictor of Survival in Breast Cancer", *New England Journal of Medicine*, **347**(25):1999-2009

**Examples**

```
## load Biobase package
library(Biobase)
## load the dataset
data(breastCancerData)
## show the first 5 columns of the expression data
exprs(nki7g)[ ,1:5]
## show the first 6 rows of the phenotype data
head(pData(nki7g))
## show first 20 feature names
featureNames(nki7g)
## show the experiment data summary
experimentData(nki7g)
## show the used platform
annotation(nki7g)
## show the abstract for this dataset
abstract(nki7g)
```

---

no.at.risk

---

*Function to compute the number of individuals at risk*


---

**Description**

Function to compute the number of individuals at risk at certain time points, as used in the Kaplan-Meier estimator for instance, depending on stratification.

**Usage**

```
no.at.risk(formula.s, data.s, sub.s = "all", t.step, t.end)
```

**Arguments**

formula.s	formula composed of a Surv object and a strata variable (i.e. stratification).
data.s	data frame composed of the variables used in the formula.
sub.s	vector of booleans specifying if only a subset of the data should be considered.
t.step	time step at which the number of individuals at risk is computed.
t.end	maximum time to be considered.

**Details**

The original version of this function was kindly provided by Dr Christos Hatzis (January, 17th 2006).

**Value**

number of individuals at risk at each time step specified in t.step up to t.end.

**Author(s)**

Christos Hatzis, Benjamin Haibe-Kains

**See Also**

[survfit](#), [km.coxph.plot](#)

**Examples**

```
require(survival)
set.seed(12345)
stime <- rexp(100)
cens <- runif(100,.5,2)
sevent <- as.numeric(stime <= cens)
stime <- pmin(stime, cens)
strat <- sample(1:3, 100, replace=TRUE)
dd <- data.frame("surv.time"=stime, "surv.event"=sevent, "strat"=strat)
no.at.risk(formula.s=Surv(surv.time,surv.event) ~ strat, data.s=dd,
  sub.s="all", t.step=0.05, t.end=1)
```

---

sbrier.score2proba	<i>Function to compute the BSCs from a risk score, for all the times of event occurrence</i>
--------------------	--

---

**Description**

The function computes all the Brier scores (BSC) and the corresponding integrated Briser score (IBSC) from a risk score, for all the times of event occurrence. The risk score is first transformed in survival probabilities using either a Cox model or the product-limit estimator.

**Usage**

```
sbrier.score2proba(data.tr, data.ts, method = c("cox", "prodlim"))
```

**Arguments**

data.tr	the data frame for the training set. This data frame must contain three columns for the times, the event occurrence and the risk score. These columns are called "time", "event" and "score" respectively.
data.ts	the data frame for the test set. This data frame must contain three columns for the times, the event occurrence and the risk score. These columns are called "time", "event" and "score" respectively.
method	method for survival probabilities estimation using either a Cox model or the product-limit estimator

**Value**

time	vector of points in time
bsc	vector of Brier scores (BSC) at ome points in time
bsc.integrated	value of the integrated Brier score (IBSC)

**Author(s)**

Benjamin Haibe-Kains



## References

- Brier, G. W. (1950) "Verification of forecasts expressed in terms of probabilities", *Monthly Weather Review*, **78**, pages 1–3.
- Graf, E. and Schmoor, C. and Sauerbrei, W. and Schumacher, M. (1999) "Assessment and comparison of prognostic classification schemes for survival data ", *Statistics in Medicine*, **18**, pages 2529–2545.
- Cox, D. R. (1972) "Regression Models and Life Tables", *Journal of the Royal Statistical Society Series B*, **34**, pages 187–220.
- Andersen, P. K. and Borgan, O. and Gill, R. D. and Keiding, N. (1993) "Statistical Models Based on Counting Processes", *Springer*.

## See Also

[sbrier](#), [coxph](#), [prodlim](#)

## Examples

```
set.seed(12345)
age <- rnorm(30, 50, 10)
stime <- rexp(30)
cens <- runif(30,.5,2)
sevent <- as.numeric(stime <= cens)
stime <- pmin(stime, cens)
dd <- data.frame("time"=stime, "event"=sevent, "score"=age)

#Cox's model
sbrier.score2proba(data.tr=dd, data.ts=dd, method="cox")
#product-limit estimator
sbrier.score2proba(data.tr=dd, data.ts=dd, method="prodlim")
```

---

score2proba

*Function to compute the survival probabilities from a risk score*

---

## Description

the function uses either a Cox model or the product-limit estimator to compute the survival probabilities from a risk score for a specific point in time.

## Usage

```
score2proba(data.tr, score, yr, method = c("cox", "prodlim"),
  conf.int = 0.95, which.est = c("point", "lower", "upper"))
```

## Arguments

data.tr	the data frame for the training set. This data frame must contain three columns for the times, the event occurrence and the risk score. These columns are called "time", "event" and "score" respectively
score	risk score for the test set
yr	a point in time for which the survival probabilities must be computed

method	method for survival probabilities estimation, either cox or prodlim for the Cox model or the product-limit estimator, respectively
conf.int	value in [0,1]. Default at 0.95
which.est	which estimation to be returned? point for the point estimate, lower for the lower bound and upper for the upper bound

**Value**

vector of predicted survival probabilities

**Author(s)**

Benjamin Haibe-Kains

**References**

Cox, D. R. (1972) "Regression Models and Life Tables", *Journal of the Royal Statistical Society Series B*, **34**, pages 187–220.

Andersen, P. K. and Borgan, O. and Gill, R. D. and Keiding, N. (1993) "Statistical Models Based on Counting Processes", *Springer*.

**See Also**

[coxph](#), [prodlim](#)

**Examples**

```
set.seed(12345)
age <- rnorm(30, 50, 10)
stime <- rexp(30)
cens <- runif(30,.5,2)
sevent <- as.numeric(stime <= cens)
stime <- pmin(stime, cens)
dd <- data.frame("time"=stime, "event"=sevent, "score"=age)

#Cox's model
score2proba(data.tr=dd, score=dd$score, yr=1, method="cox")
#product-limit estimator
score2proba(data.tr=dd, score=dd$score, yr=1, method="prodlim")
```

---

td.sens.spec	<i>Function to compute sensitivity and specificity for a binary classification of survival data</i>
--------------	---

---

**Description**

The function is a wrapper for the [survivalROC.C](#) function in order to compute sensitivity and specificity for a binary classification of survival data.

**Usage**

```
td.sens.spec(cl, surv.time, surv.event, time, span = 0, sampling = FALSE,
na.rm = FALSE, ...)
```

**Arguments**

cl	vector of binary classes.
surv.time	vector of times to event occurrence.
surv.event	vector of event occurrence indicators.
time	time point for sensitivity and specificity estimations.
span	Span for the NNE. Default value is 0.
sampling	jackknife procedure to estimate the standard error of sensitivity and specificity estimations.
na.rm	TRUE if the missing values should be removed from the data, FALSE otherwise.
...	additional arguments to be passed to the <a href="#">survivalROC</a> function.

**Details**

Only NNE method is used to estimate sensitivity and specificity (see [survivalROC.C](#)). The standard error for sensitivity and specificity is estimated through jackknife procedure (see [jackknife](#)).

**Value**

sens	sensitivity estimate
sens.se	standard error for sensitivity estimate
spec	specificity estimate
spec.se	standard error for specificity estimate

**Author(s)**

Benjamin Haibe-Kains

**References**

Heagerty, P. J. and Lumley, T. L. and Pepe, M. S. (2000) "Time-Dependent ROC Curves for Censored Survival Data and a Diagnostic Marker", *Biometrics*, **56**, pages 337–344.

Efron, B. and Tibshirani, R. (1986). "The Bootstrap Method for standard errors, confidence intervals, and other measures of statistical accuracy", *Statistical Science*, **1** (1), pages 1–35.

**See Also**

[survivalROC](#)

**Examples**

```
set.seed(12345)
gender <- sample(c(0,1), 100, replace=TRUE)
stime <- rexp(100)
cens <- runif(100,.5,2)
sevent <- as.numeric(stime <= cens)
stime <- pmin(stime, cens)
mysenspec <- td.sens.spec(cl=gender, surv.time=stime, surv.event=sevent,
  time=1, span=0, na.rm=FALSE)
```

tdrocc

*Function to compute time-dependent ROC curves***Description**

The function is a wrapper for the [survivalROC](#) function in order to compute the time-dependent ROC curves.

**Usage**

```
tdrocc(x, surv.time, surv.event, surv.entry = NULL, time, cutpts = NA,
       na.rm = FALSE, verbose = FALSE, span = 0, lambda = 0, ...)
```

**Arguments**

x	vector of risk scores.
surv.time	vector of times to event occurrence.
surv.event	vector of event occurrence indicators.
surv.entry	entry time for the subjects.
time	time point for the ROC curve.
cutpts	cut points for the risk score.
na.rm	TRUE if the missing values should be removed from the data, FALSE otherwise.
verbose	verbosity of the function.
span	Span for the NNE, need either lambda or span for NNE.
lambda	smoothing parameter for NNE.
...	additional arguments to be passed to the <a href="#">survivalROC</a> function.

**Value**

spec	specificity estimates
sens	sensitivity estimates
rule	rule to compute the predictions at each cutoff
cuts	cutoffs
time	time point at which the time-dependent ROC is computed
survival	overall survival at the time point
AUC	Area Under the Curve (AUC) of the time-dependent ROC curve
data	survival data and risk score used to compute the time-dependent ROC curve

**Author(s)**

Benjamin Haibe-Kains

**References**

Heagerty, P. J. and Lumley, T. L. and Pepe, M. S. (2000) "Time-Dependent ROC Curves for Censored Survival Data and a Diagnostic Marker", *Biometrics*, **56**, pages 337–344.

**See Also**[survivalROC](#)**Examples**

```

set.seed(12345)
age <- rnorm(100, 50, 10)
stime <- rexp(100)
cens <- runif(100,.5,2)
sevent <- as.numeric(stime <= cens)
stime <- pmin(stime, cens)
tdroc <- tdrocc(x=age, surv.time=stime, surv.event=sevent, time=1,
  na.rm=TRUE, verbose=FALSE)
##plot the time-dependent ROC curve
plot(x=1-tdroc$spec, y=tdroc$sens, type="l", xlab="1 - specificity",
  ylab="sensitivity", xlim=c(0, 1), ylim=c(0, 1))
lines(x=c(0,1), y=c(0,1), lty=3, col="red")

```

test.hetero.est

*Function to test the heterogeneity of set of probabilities***Description**

The function tests whether a set of p-values are heterogeneous.

**Usage**

```
test.hetero.est(x, x.se, na.rm = FALSE)
```

**Arguments**

x	vector of estimates
x.se	vector of standard errors of the corresponding estimates
na.rm	TRUE if the missing values should be removed from the data, FALSE otherwise

**Details**

The heterogeneity test is known to be very conservative. Consider a p-value < 0.1 as significant.

**Value**

Q	Q statistic
p.value	p-value of the heterogeneity test

**Author(s)**

Benjamin Haibe-Kains

**References**

Cochrane, W. G. (1954) "The combination of estimates from different experiments", *Biometrics*, **10**, pages 101–129.

**See Also**

combine.test

**Examples**

```
set.seed(12345)
x1 <- rnorm(100, 50, 10) + rnorm(100, 0, 2)
m1 <- mean(x1)
se1 <- sqrt(var(x1))
x2 <- rnorm(100, 75, 15) + rnorm(100, 0, 5)
m2 <- mean(x2)
se2 <- sqrt(var(x2))

test.hetero.est(x=c(m1, m2), x.se=c(se1, se2))
```

---

test.hetero.test

---

*Function to test the heterogeneity of set of probabilities*


---

**Description**

The function tests whether a set of p-values are heterogeneous.

**Usage**

```
test.hetero.test(p, weight, na.rm = FALSE)
```

**Arguments**

p	vector of p-values
weight	vector of weights (e.g. sample size of each study)
na.rm	TRUE if the missing values should be removed from the data, FALSE otherwise

**Details**

The p-values should be one-sided and computed from the same null hypothesis.

**Value**

Q	Q statistic
p.value	p-value of the heterogeneity test

**Author(s)**

Benjamin Haibe-Kains

## References

Cochrane, W. G. (1954) "The combination of estimates from different experiments", *Biometrics*, **10**, pages 101–129.

Whitlock, M. C. (2005) "Combining probability from independent tests: the weighted Z-method is superior to Fisher's approach", *J. Evol. Biol.*, **18**, pages 1368–1373.

## See Also

combine.test

## Examples

```
p <- c(0.01, 0.13, 0.07, 0.2)
w <- c(100, 50, 200, 30)

#with equal weights
test.hetero.test(p=p)
#with p-values weighted by the sample size of the studies
test.hetero.test(p=p, weight=w)
```

---

transbig7g

*Subset of the TRANSBIG dataset containing gene expression, annotations and clinical data.*

---

## Description

This dataset contains a subset of the gene expression, annotations and clinical data from the TRANSBIG dataset (see section details). The subset contains the seven genes introduced by Desmedt et al. 2008

## Format

ExpressionSet with 7 features and 198 samples, containing:

- `exprs(transbig7g)`: Matrix containing gene expressions as measured by Affymetrix hgu133a technology (single-channel, oligonucleotides).
- `fData(transbig7g)`: AnnotatedDataFrame containing annotations of Affy microarray platform hgu133a.
- `pData(transbig7g)`: AnnotatedDataFrame containing Clinical information of the breast cancer patients whose tumors were hybridized.
- `experimentalData(transbig7g)`: MIAME object containing information about the dataset.
- `annotation(transbig7g)`: Name of the affy chip.

## Details

This dataset represents a subset of the study published by Desmedt et al. 2007. The subset contains the genes AURKA (also known as STK6, STK7, or STK15), PLAU (also known as UPA), STAT1, VEGF, CASP3, ESR1, and ERBB2, as introduced by Desmedt et al. 2008. The seven genes represent the proliferation, tumor invasion/metastasis, immune response, angiogenesis, apoptosis phenotypes, and the ER and HER2 signaling, respectively.

**Source****transbig:**

<http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE7390>

**References**

Christine Desmedt, Fanny Piette, Sherene Loi, Yixin Wang, Françoise Lallemand, Benjamin Haibe-Kains, Giuseppe Viale, Mauro Delorenzi, Yi Zhang, Mahasti Saghatchian d Assignies, Jonas Bergh, Rosette Lidereau, Paul Ellis, Adrian L. Harris, Jan G. M. Klijn, John A. Foekens, Fatima Cardoso, Martine J. Piccart, Marc Buyse and Christos Sotiriou (2007) "Strong Time Dependence of the 76- Gene Prognostic Signature for Node-Negative Breast Cancer Patients in the TRANSBIG Multicenter Independent Validation Series", *Clinical Cancer Research*, **13**(11):3207-3214

**Examples**

```
## load Biobase package
library(Biobase)
## load the dataset
data(breastCancerData)
## show the first 5 columns of the expression data
exprs(transbig7g)[ ,1:5]
## show the first 6 rows of the phenotype data
head(pData(transbig7g))
## show first 20 feature names
featureNames(transbig7g)
## show the experiment data summary
experimentData(transbig7g)
## show the used platform
annotation(transbig7g)
## show the abstract for this dataset
abstract(transbig7g)
```

---

unt7g

*Subset of UNT dataset containing gene expression, annotations and clinical data.*

---

**Description**

This dataset contains a subset of the gene expression, annotations and clinical data from the UNT datasets (see section details). The subset contains the seven genes introduced by Desmedt et al. 2008

**Format**

ExpressionSet with 7 features and 137 samples, containing:

- `exprs(unt7g)`: Matrix containing gene expressions as measured by Affymetrix hgu133a and hgu133b technology (single-channel, oligonucleotides).
- `fData(unt7g)`: AnnotatedDataFrame containing annotations of Affy microarray platform hgu133a and hgu133b.
- `pData(unt7g)`: AnnotatedDataFrame containing Clinical information of the breast cancer patients whose tumors were hybridized.



- `experimentalData(unt7g)`: MIAME object containing information about the dataset.
- `annotation(unt7g)`: Name of the affy chip.

## Details

This dataset represents a subset of the study published by Sotiriou et al. 2006. The subset contains the genes AURKA (also known as STK6, STK7, or STK15), PLAU (also known as UPA), STAT1, VEGF, CASP3, ESR1, and ERBB2, as introduced by Desmedt et al. 2008. The seven genes represent the proliferation, tumor invasion/metastasis, immune response, angiogenesis, apoptosis phenotypes, and the ER and HER2 signaling, respectively.

## Source

**unt:**

<http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE2990>

<http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE6532>

## References

Christos Sotiriou, Pratyaksha Wirapati, Sherene Loi, Adrian Harris, Steve Fox, Johanna Smeds, Hans Nordgren, Pierre Farmer, Viviane Praz, Benjamin Haibe-Kains, Christine Desmedt, Denis Larsimont, Fatima Cardoso, Hans Peterse, Dimitry Nuyten, Marc Buyse, Marc J. Van de Vijver, Jonas Bergh, Martine Piccart and Mauro Delorenzi (2006) "Gene Expression Profiling in Breast Cancer: Understanding the Molecular Basis of Histologic Grade To Improve Prognosis", *Journal of the National Cancer Institute*, **98**(4):262-272

## Examples

```
## load Biobase package
library(Biobase)
## load the dataset
data(breastCancerData)
## show the first 5 columns of the expression data
exprs(unt7g)[,1:5]
## show the first 6 rows of the phenotype data
head(pData(unt7g))
## show first 20 feature names
featureNames(unt7g)
## show the experiment data summary
experimentData(unt7g)
## show the used platform
annotation(unt7g)
## show the abstract for this dataset
abstract(unt7g)
```

## Description

This dataset contains a subset of the gene expression, annotations and clinical data from the UPP datasets (see section details). The subset contains the seven genes introduced by Desmedt et al. 2008

## Format

ExpressionSet with 7 features and 251 samples, containing:

- `exprs(upp7g)`: Matrix containing gene expressions as measured by Affymetrix hgu133a and hgu133b technology (single-channel, oligonucleotides).
- `fData(upp7g)`: AnnotatedDataFrame containing annotations of Affy microarray platform hgu133a and hgu133b.
- `pData(upp7g)`: AnnotatedDataFrame containing Clinical information of the breast cancer patients whose tumors were hybridized.
- `experimentalData(upp7g)`: MIAME object containing information about the dataset.
- `annotation(upp7g)`: Name of the affy chip.

## Details

This dataset represents a subset of the study published by Miller et al. 2005. The subset contains the genes AURKA (also known as STK6, STK7, or STK15), PLAU (also known as UPA), STAT1, VEGF, CASP3, ESR1, and ERBB2, as introduced by Desmedt et al. 2008. The seven genes represent the proliferation, tumor invasion/metastasis, immune response, angiogenesis, apoptosis phenotypes, and the ER and HER2 signaling, respectively.

## Source

**upp:**

<http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE3494>

## References

Lance D. Miller, Johanna Smeds, Joshy George, Vinsensius B. Vega, Liza Vergara, Alexander Ploner, Yudi Pawitan, Per Hall, Sigrid Klaar, Edison T. Liu and Jonas Bergh (2005) "An expression signature for p53 status in human breast cancer predicts mutation status, transcriptional effects, and patient survival" *Proceedings of the National Academy of Sciences of the United States of America* **102**(38):13550-13555

## Examples

```
## load Biobase package
library(Biobase)
## load the dataset
data(breastCancerData)
## show the first 5 columns of the expression data
exprs(upp7g)[,1:5]
## show the first 6 rows of the phenotype data
head(pData(upp7g))
## show first 20 feature names
featureNames(upp7g)
## show the experiment data summary
experimentData(upp7g)
```

```
## show the used platform
annotation(upp7g)
## show the abstract for this dataset
abstract(upp7g)
```

---

vdx7g	<i>Subset of VDX dataset containing gene expression, annotations and clinical data.</i>
-------	---

---

## Description

This dataset contains a subset of the gene expression, annotations and clinical data from the VDX datasets (see section details). The subset contains the seven genes introduced by Desmedt et al. 2008

## Format

ExpressionSet with 7 features and 344 samples, containing:

- `exprs(vdx7g)`: Matrix containing gene expressions as measured by Affymetrix hgu133a technology (single-channel, oligonucleotides).
- `fData(vdx7g)`: AnnotatedDataFrame containing annotations of Affy microarray platform hgu133a.
- `pData(vdx7g)`: AnnotatedDataFrame containing Clinical information of the breast cancer patients whose tumors were hybridized.
- `experimentalData(vdx7g)`: MIAME object containing information about the dataset.
- `annotation(vdx7g)`: Name of the affy chip.

## Details

This dataset represents a subset of the study published by Wang. et al. 2005 and Minn et al. 2007 . The subset contains the genes AURKA (also known as STK6, STK7, or STK15), PLAU (also known as UPA), STAT1, VEGF, CASP3, ESR1, and ERBB2, as introduced by Desmedt et al. 2008. The seven genes represent the proliferation, tumor invasion/metastasis, immune response, angiogenesis, apoptosis phenotypes, and the ER and HER2 signaling, respectively.

## Source

**vdx:**

<http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE2034>

## References

Y. Wang, J. G. Klijn, Y. Zhang, A. M. Sieuwerts, M. P. Look, F. Yang, D. Talantov, M. Timmermans, M. E. Meijer-van Gelder, J. Yu, T. Jatkoe, E. M. Berns, D. Atkins and J. A. Foekens (2005) "Gene-Expression Profiles to Predict Distant Metastasis of Lymph-Node-Negative Primary Breast Cancer", *Lancet*, **365**:671-679

Andy J. Minn, Gaorav P. Gupta, David Padua, Paula Bos, Don X. Nguyen, Dimitry Nuyten, Bas Kreike, Yi Zhang, Yixin Wang, Hemant Ishwaran, John A. Foekens, Marc van de Vijver and Joan Massague (2007) "Lung metastasis genes couple breast tumor size and metastatic spread", *Proceedings of the National Academy of Sciences*, **104**(16):6740-6745

**Examples**

```
## load Biobase package
library(Biobase)
## load the dataset
data(breastCancerData)
## show the first 5 columns of the expression data
exprs(vdx7g)[,1:5]
## show the first 6 rows of the phenotype data
head(pData(vdx7g))
## show first 20 feature names
featureNames(vdx7g)
## show the experiment data summary
experimentData(vdx7g)
## show the used platform
annotation(vdx7g)
## show the abstract for this dataset
abstract(vdx7g)
```

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