Package 'genefu'

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Description This package contains functions implementing various tasks usually required by gene expression analysis, especially in breast cancer studies: gene mapping between different microarray platforms, identification of molecular subtypes, implementation of published gene signatures, gene selection, and survival analysis.

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Description

This package contains functions implementing various tasks usually required by gene expression analysis, especially in breast cancer studies: gene mapping between different microarray platforms, identification of molecular subtypes, implementation of published gene signatures, gene selection, and survival analysis.

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See Also

Useful links:

• http://www.pmgenomics.ca/bhklab/software/genefu

bimod

Function to identify bimodality for gene expression or signature score

Description

This function fits a mixture of two Gaussians to identify bimodality. Useful to identify ER of HER2 status of breast tumors using ESR1 and ERBB2 expressions respectively.

Usage

```
bimod(x, data, annot, do.mapping = FALSE, mapping, model = c("E", "V"), do.scale = TRUE, verbose = FALSE, ...)
```

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Arguments

x	Matrix containing the gene(s) in the gene list in rows and at least three columns: "probe", "EntrezGene.ID" and "coefficient" standing for the name of the probe, the NCBI Entrez Gene id and the coefficient giving the direction and the strength of the association of each gene in the gene list.
data	Matrix of gene expressions with samples in rows and probes in columns, dimnames being properly defined.
annot	Matrix of annotations with at least one column named "EntrezGene.ID", dimnames being properly defined.
do.mapping	TRUE if the mapping through Entrez Gene ids must be performed (in case of ambiguities, the most variant probe is kept for each gene), FALSE otherwise.
mapping	Matrix with columns "EntrezGene.ID" and "probe" used to force the mapping such that the probes are not selected based on their variance.
model	Model name used in Mclust.
do.scale	TRUE if the gene expressions or signature scores must be rescaled (see rescale), FALSE otherwise.
verbose	TRUE to print informative messages, FALSE otherwise.
	Additional parameters to pass to sig.score.

Value

A list with items:

- status: Status being 0 or 1.
- status 1. proba: Probability p to be of status 1, the probability to be of status 0 being 1-p.
- gaussians: Matrix of parameters fitted in the mixture of two Gaussians. Matrix of NA values if EM algorithm did not converge.
- BIC: Values (gene expressions or signature scores) used to identify bimodality.
- BI: Bimodality Index (BI) as defined by Wang et al., 2009.
- x: Values (gene expressions or signature scores) used to identify bimodality

References

Desmedt C, Haibe-Kains B, Wirapati P, Buyse M, Larsimont D, Bontempi G, Delorenzi M, Piccart M, and Sotiriou C (2008) "Biological processes associated with breast cancer clinical outcome depend on the molecular subtypes", Clinical Cancer Research, 14(16):5158–5165. Wirapati P, Sotiriou C, Kunkel S, Farmer P, Pradervand S, Haibe-Kains B, Desmedt C, Ignatiadis M, Sengstag T, Schutz F, Goldstein DR, Piccart MJ and Delorenzi M (2008) "Meta-analysis of Gene-Expression Profiles in Breast Cancer: Toward a Unified Understanding of Breast Cancer Sub-typing and Prognosis Signatures", Breast Cancer Research, 10(4):R65. Fraley C and Raftery E (2002) "Model-Based Clustering, Discriminant Analysis, and Density Estimation", Journal of American Statistical Association, 97(458):611–631. Wang J, Wen S, Symmans FW, Pusztai L and Coombes KR (2009) "The bimodality index: a criterion for discovering and ranking bimodal signatures from cancer gene expression profiling data", Cancer Informatics, 7:199–216.

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See Also

mclust::Mclust

Examples

```
# load NKI data
data(nkis)
# load gene modules from Desmedt et al. 2008
data(mod1)
# retrieve esr1 affy probe and Entrez Gene id
esr1 <- mod1$ESR1[1, ,drop=FALSE]
# computation of signature scores
esr1.bimod <- bimod(x=esr1, data=data.nkis, annot=annot.nkis, do.mapping=TRUE,
    model="V", verbose=TRUE)
table("ER.IHC"=demo.nkis[ ,"er"], "ER.GE"=esr1.bimod$status)</pre>
```

boxplotplus2

Box plot of group of values with corresponding jittered points

Description

This function allows for display a boxplot with jittered points.

Usage

```
boxplotplus2(x, .jit = 0.25, .las = 1, .ylim, box.col = "lightgrey",
pt.col = "blue", pt.cex = 0.5, pt.pch = 16, med.line = FALSE,
med.col = "goldenrod", ...)
```

Arguments

x	could be a list of group values or a matrix (each group is a row).
.jit	Amount of jittering noise.
.las	Numeric in 0,1,2,3; the style of axis labels.
.ylim	Range for y axis.
box.col	Color for boxes.
pt.col	Color for groups (jittered points).
pt.cex	A numerical value giving the amount by which plotting jittered points should be magnified relative to the default.
pt.pch	Either an integer specifying a symbol or a single character to be used as the default in plotting jittered points. See points for possible values and their interpretation.
med.line	TRUE if a line should link the median of each group, FALSE otherwise.
med.col	Color of med.line.
	Additional parameters for boxplot function.

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Value

Number of samples in each group.

Note

2.21.2006 - Christos Hatzis, Nuvera Biosciences

See Also

```
graphics::boxplot, base::jitter
```

Examples

```
dd <- list("G1"=runif(20), "G2"=rexp(30) * -1.1, "G3"=rnorm(15) * 1.3)
boxplotplus2(x=dd, .las=3, .jit=0.75, .ylim=c(-3,3), pt.cex=0.75,
  pt.col=c(rep("darkred", 20), rep("darkgreen", 30), rep("darkblue", 15)),
  pt.pch=c(0, 9, 17))</pre>
```

claudinLow

Claudin-low classification for Breast Cancer Data

Description

Subtyping method for identifying Claudin-Low Breast Cancer Samples. Code generously provided by Aleix Prat.

Usage

```
claudinLow(x, classes="", y, nGenes="", priors="equal",
    std=FALSE, distm="euclidean", centroids=FALSE)
```

Arguments

X	the data matrix of training samples, or pre-calculated centroids.
classes	a list labels for use in coloring the points.
У	the data matrix of test samples.
nGenes	the number of genes selected when training the model.
priors	'equal' assumes equal class priors, 'class' calculates them based on proportion in the data.
std	when true, the training and testing samples are standardized to mean= 0 and var= 1 .
distm	the distance metric for determining the nearest centroid, can be one of euclidean, pearson, or spearman.
centroids	when true, it is assumed that x consists of pre-calculated centroids.

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Value

A list with items:

- predictions
- testData
- distances
- · centroids

References

Aleix Prat, Joel S Parker, Olga Karginova, Cheng Fan, Chad Livasy, Jason I Herschkowitz, Xiaping He, and Charles M. Perou (2010) "Phenotypic and molecular characterization of the claudin-low intrinsic subtype of breast cancer", Breast Cancer Research, 12(5):R68

See Also

```
medianCtr(), q
```

Examples

```
data(claudinLowData)
#Training Set
train <- claudinLowData
train$xd <- medianCtr(train$xd)
# Testing Set
test <- claudinLowData
test$xd <- medianCtr(test$xd)

# Generate Predictions
predout <- claudinLow(x=train$xd, classes=as.matrix(train$classes$Group,ncol=1), y=test$xd)

# Obtain results
results <- cbind(predout$predictions, predout$distances)
#write.table(results, "T.E.9CELL.LINE_results.txt", sep="\t", col=T, row=FALSE)</pre>
```

claudinLowData

claudinLowData for use in the claudinLow classifier. Data generously provided by Aleix Prat.

Description

Training and Testing Data for use with the Claudin-Low Classifier

Usage

```
data(claudinLowData)
```

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Format

• xd: Matrix of 807 features and 52 samples

• classes: factor to split samples

• nfeatures: number of features

• nsamples: number of samples

• fnames: names of features

• snames: names of samples

Source

http://jnci.oxfordjournals.org/cgi/content/full/98/4/262/DC1

References

Aleix Prat, Joel S Parker, Olga Karginova, Cheng Fan, Chad Livasy, Jason I Herschkowitz, Xiaping He, and Charles M. Perou (2010) "Phenotypic and molecular characterization of the claudin-low intrinsic subtype of breast cancer", Breast Cancer Research, 12(5):R68

See Also

claudinLow()

collapseIDs

Utility function to collapse IDs

Description

Utility function called within the claudinLow classifier

Usage

```
collapseIDs(x,allids=row.names(x),method="mean")
```

Arguments

x Matrix of numbers.

allids Defaults to rownames of matrix.

method Default method is "mean".

Value

A matrix

References

```
citation("claudinLow")
```

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See Also

claudinLow

Description

This function performs a statistical comparison of the correlation coefficients as computed between each probe and prototype.

Usage

```
compareProtoCor(gene.cor, proto.cor, nn,
p.adjust.m = c("none", "holm", "hochberg", "hommel", "bonferroni", "BH", "BY", "fdr"))
```

Arguments

gene.cor Correlation coefficients between the probes and each of the prototypes.

Pairwise correlation coefficients of the prototypes.

Number of samples used to compute the correlation coefficients between the probes and each of the prototypes.

p.adjust.m Correction method as defined in p.adjust.

Value

Data frame with probes in rows and with three columns: "proto" is the prototype to which the probe is the most correlated, "cor" is the actual correlation, and "signif" is the (corrected) p-value for the superiority of the correlation to this prototype compared to the second highest correlation.

See Also

compute.proto.cor.meta, compute.pairw.cor.meta

```
# load VDX dataset
data(vdxs)
# load NKI dataset
data(nkis)
# reduce datasets
ginter <- intersect(annot.vdxs[ ,"EntrezGene.ID"], annot.nkis[ ,"EntrezGene.ID"])
ginter <- ginter[!is.na(ginter)][1:30]
myx <- unique(c(match(ginter, annot.vdxs[ ,"EntrezGene.ID"]),
    sample(x=1:nrow(annot.vdxs), size=20)))
data2.vdxs <- data.vdxs[ ,myx]
annot2.vdxs <- annot.vdxs[myx, ]</pre>
```

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```
myx <- unique(c(match(ginter, annot.nkis[ ,"EntrezGene.ID"]),</pre>
sample(x=1:nrow(annot.nkis), size=20)))
data2.nkis <- data.nkis[ ,myx]</pre>
annot2.nkis <- annot.nkis[myx, ]</pre>
# mapping of datasets
datas <- list("VDX"=data2.vdxs,"NKI"=data2.nkis)</pre>
annots <- list("VDX"=annot2.vdxs, "NKI"=annot2.nkis)</pre>
datas.mapped <- map.datasets(datas=datas, annots=annots, do.mapping=TRUE)</pre>
# define some prototypes
protos <- paste("geneid", ginter[1:3], sep=".")</pre>
# compute meta-estimate of correlation coefficients to the three prototype genes
probecor <- compute.proto.cor.meta(datas=datas.mapped$datas, proto=protos,</pre>
 method="pearson")
# compute meta-estimate of pairwise correlation coefficients between prototypes
datas.proto <- lapply(X=datas.mapped$datas, FUN=function(x, p) {</pre>
  return(x[ ,p,drop=FALSE]) }, p=protos)
protocor <- compute.pairw.cor.meta(datas=datas.proto, method="pearson")</pre>
# compare correlation coefficients to each prototype
res <- compareProtoCor(gene.cor=probecor$cor, proto.cor=protocor$cor,</pre>
nn=probecor$cor.n, p.adjust.m="fdr")
head(res)
```

compute.pairw.cor.meta

Function to compute pairwise correlations in a meta-analytical framework

Description

This function computes meta-estimate of pairwise correlation coefficients for a set of genes from a list of gene expression datasets.

Usage

```
compute.pairw.cor.meta(datas, method = c("pearson", "spearman"))
```

Arguments

datas List of datasets. Each dataset is a matrix of gene expressions with samples in

rows and probes in columns, dimnames being properly defined. All the datasets

must have the same probes.

method Estimator for correlation coefficient, can be either pearson or spearman.

Value

A list with items:

- cor Matrix of meta-estimate of correlation coefficients with probes in rows and prototypes in columns
- cor.n Number of samples used to compute meta-estimate of correlation coefficients.

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See Also

map.datasets, compute.proto.cor.meta

Examples

```
# load VDX dataset
data(vdxs)
# load NKI dataset
data(nkis)
# reduce datasets
ginter <- intersect(annot.vdxs[ ,"EntrezGene.ID"], annot.nkis[ ,"EntrezGene.ID"])</pre>
ginter <- ginter[!is.na(ginter)][1:30]</pre>
myx <- unique(c(match(ginter, annot.vdxs[ ,"EntrezGene.ID"]),</pre>
  sample(x=1:nrow(annot.vdxs), size=20)))
data2.vdxs <- data.vdxs[ ,myx]</pre>
annot2.vdxs <- annot.vdxs[myx, ]</pre>
myx <- unique(c(match(ginter, annot.nkis[ ,"EntrezGene.ID"]),</pre>
  sample(x=1:nrow(annot.nkis), size=20)))
data2.nkis <- data.nkis[ ,myx]</pre>
annot2.nkis <- annot.nkis[myx, ]</pre>
# mapping of datasets
datas <- list("VDX"=data2.vdxs,"NKI"=data2.nkis)</pre>
annots <- list("VDX"=annot2.vdxs, "NKI"=annot2.nkis)</pre>
datas.mapped <- map.datasets(datas=datas, annots=annots, do.mapping=TRUE)</pre>
# compute meta-estimate of pairwise correlation coefficients
pairwcor <- compute.pairw.cor.meta(datas=datas.mapped$datas, method="pearson")</pre>
str(pairwcor)
```

 ${\it compute.pairw.cor.z} \qquad \textit{Function to compute the Z transformation of the pairwise correlations} \\ \textit{for a list of datasets}$

Description

This function computes the Z transformation of the meta-estimate of pairwise correlation coefficients for a set of genes from a list of gene expression datasets.

Usage

```
compute.pairw.cor.z(datas, method = c("pearson"))
```

Arguments

datas List of datasets. Each dataset is a matrix of gene expressions with samples in

rows and probes in columns, dimnames being properly defined. All the datasets

must have the same probes.

method Estimator for correlation coefficient, can be either pearson or spearman.

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Value

A list with items: -z Z transformation of the meta-estimate of correlation coefficients. -se Standard error of the Z transformation of the meta-estimate of correlation coefficients. -nn Number of samples used to compute the meta-estimate of correlation coefficients.

See Also

map.datasets, compute.pairw.cor.meta, compute.proto.cor.meta

compute.proto.cor.meta

Function to compute correlations to prototypes in a meta-analytical framework

Description

This function computes meta-estimate of correlation coefficients between a set of genes and a set of prototypes from a list of gene expression datasets.

Usage

```
compute.proto.cor.meta(datas, proto, method = c("pearson", "spearman"))
```

Arguments

.1 . 4	T '-4 - C 1-44-	Tr1. 1.4	·		1
datas	List of datasets	- Hach dataset	is a matrix of get	ne expressions with	samples in
aacas	List of datasets.	Lacii dataset	is a madify of goi	ic expressions with	builipies in

rows and probes in columns, dimnames being properly defined. All the datasets

must have the same probes.

proto Names of prototypes (e.g. their EntrezGene ID).

method Estimator for correlation coefficient, can be either pearson or spearman

Value

A list with items: -cor Matrix of meta-estimate of correlation coefficients with probes in rows and prototypes in columns. -cor.n Number of samples used to compute meta-estimate of correlation coefficients.

See Also

map.datasets

14 cordiff.dep

Examples

```
# load VDX dataset
data(vdxs)
# load NKI dataset
data(nkis)
# reduce datasets
ginter <- intersect(annot.vdxs[ ,"EntrezGene.ID"], annot.nkis[ ,"EntrezGene.ID"])</pre>
ginter <- ginter[!is.na(ginter)][1:30]</pre>
myx <- unique(c(match(ginter, annot.vdxs[ ,"EntrezGene.ID"]),</pre>
  sample(x=1:nrow(annot.vdxs), size=20)))
data2.vdxs <- data.vdxs[ ,myx]</pre>
annot2.vdxs <- annot.vdxs[myx, ]</pre>
myx <- unique(c(match(ginter, annot.nkis[ ,"EntrezGene.ID"]),</pre>
  sample(x=1:nrow(annot.nkis), size=20)))
data2.nkis <- data.nkis[ ,myx]</pre>
annot2.nkis <- annot.nkis[myx, ]</pre>
# mapping of datasets
datas <- list("VDX"=data2.vdxs,"NKI"=data2.nkis)</pre>
annots <- list("VDX"=annot2.vdxs, "NKI"=annot2.nkis)</pre>
datas.mapped <- map.datasets(datas=datas, annots=annots, do.mapping=TRUE)</pre>
# define some prototypes
protos <- paste("geneid", ginter[1:3], sep=".")</pre>
# compute meta-estimate of correlation coefficients to the three prototype genes
probecor <- compute.proto.cor.meta(datas=datas.mapped$datas, proto=protos,</pre>
  method="pearson")
str(probecor)
```

cordiff.dep

Function to estimate whether two dependent correlations differ

Description

This function tests for statistical differences between two dependent correlations using the formula provided on page 56 of Cohen & Cohen (1983). The function returns a t-value, the DF and the p-value.

Usage

```
cordiff.dep(r.x1y, r.x2y, r.x1x2, n,
  alternative = c("two.sided", "less", "greater"))
```

tors).

Arguments

r.x1y	The correlation between x1 and y where y is typically your outcome variable.
r.x2y	The correlation between x2 and y where y is typically your outcome variable.
r.x1x2	The correlation between x1 and x2 (the correlation between your two predic-

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n The sample size.

alternative A character string specifying the alternative hypothesis, must be one of "two.sided"

default), "greater" or "less". You can specify just the initial letter.

Details

This function is inspired from the cordif.dep.

Value

Vector of three values: t statistics, degree of freedom, and p-value.

References

Cohen, J. & Cohen, P. (1983) "Applied multiple regression/correlation analysis for the behavioral sciences (2nd Ed.)" Hillsdale, nJ: Lawrence Erlbaum Associates.

See Also

stats::cor, stats::t.test, compareProtoCor

Examples

```
# load VDX dataset
data(vdxs)
# retrieve ESR1, AURKA and MKI67 gene expressions
x1 <- data.vdxs[ ,"208079_s_at"]
x2 <- data.vdxs[ ,"205225_at"]
y <- data.vdxs[ ,"212022_s_at"]
# is MKI67 significantly more correlated to AURKA than ESR1?
cc.ix <- complete.cases(x1, x2, y)
cordiff.dep(r.x1y=abs(cor(x=x1[cc.ix], y=y[cc.ix], use="everything", method="pearson")), r.x2y=abs(cor(x=x2[cc.ix], y=y[cc.ix], use="everything", method="pearson")), r.x1x2=abs(cor(x=x1[cc.ix], y=x2[cc.ix], use="everything", method="pearson")), n=sum(cc.ix), alternative="greater")</pre>
```

endoPredict

Function to compute the endoPredict signature as published by Filipits et al 2011

Description

This function computes signature scores and risk classifications from gene expression values following the algorithm used for the endoPredict signature as published by Filipits et al 2011.

Usage

```
endoPredict(data, annot, do.mapping = FALSE, mapping, verbose = FALSE)
```

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Arguments

data	Matrix of gene expressions with samples in rows and probes in columns, dimnames being properly defined.
annot	Matrix of annotations with at least one column named "EntrezGene.ID", dimnames being properly defined.
do.mapping	TRUE if the mapping through Entrez Gene ids must be performed (in case of ambiguities, the most variant probe is kept for each gene), FALSE otherwise. Note that for Affymetrix HGU datasets, the mapping is not necessary.
mapping	Matrix with columns "EntrezGene.ID" and "probe" used to force the mapping such that the probes are not selected based on their variance.
verbose	TRUE to print informative messages, FALSE otherwise.

Details

The function works best if data have been noralized with MAS5. Note that for Affymetrix HGU datasets, the mapping is not necessary.

Value

A list with items: -score Continuous signature scores -risk Binary risk classification, 1 being high risk and 0 being low risk. -mapping Mapping used if necessary. -probe If mapping is performed, this matrix contains the correspondence between the gene list (aka signature) and gene expression data.

References

Filipits, M., Rudas, M., Jakesz, R., Dubsky, P., Fitzal, F., Singer, C. F., et al. (2011). "A new molecular predictor of distant recurrence in ER-positive, HER2-negative breast cancer adds independent information to conventional clinical risk factors." Clinical Cancer Research, 17(18):6012–6020.

```
# load GENE70 signature
data(sig.endoPredict)
# load NKI dataset
data(vdxs)
# compute relapse score
rs.vdxs <- endoPredict(data=data.vdxs, annot=annot.vdxs, do.mapping=FALSE)</pre>
```

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expos	Gene expression, annotations and clinical data from the International Genomics Consortium
	Contonues Consortium

Description

This dataset contains (part of) the gene expression, annotations and clinical data from the expO dataset collected by the International Genomics Consortium ().

Usage

```
data(expos)
```

Format

expos is a dataset containing three matrices

- data.expos: Matrix containing gene expressions as measured by Affymetrix hgu133plus2 technology (single-channel, oligonucleotides)
- annot.expos: Matrix containing annotations of ffymetrix hgu133plus2 microarray platform
- · demo.expos: Clinical information of the breast cancer patients whose tumors were hybridized

Source

```
http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE2109
```

References

International Genomics Consortium, http://www.intgen.org/research-services/biobanking-experience/expo/McCall MN, Bolstad BM, Irizarry RA. (2010) "Frozen robust multiarray analysis (fRMA)", Biostatistics, 11(2):242-253.

fuzzy.ttest	Function to compute the fuzzy Student t test based on weighted mean and weighted variance

Description

This function allows for computing the weighted mean and weighted variance of a vector of continuous values.

Usage

```
fuzzy.ttest(x, w1, w2, alternative=c("two.sided", "less", "greater"),
  check.w = TRUE, na.rm = FALSE)
```

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Arguments

Х	an object containing the observed values.
w1	a numerical vector of weights of the same length as x giving the weights to use for elements of x in the first class.
w2	a numerical vector of weights of the same length as x giving the weights to use for elements of x in the second class.
alternative	a character string specifying the alternative hypothesis, must be one of "two.sided" (default), "greater" or "less". You can specify just the initial letter.
check.w	TRUE if weights should be checked such that $0 \le w \le 1$ and $w1[i] + w2[i]) \le 1$ for $1 \le i \le length(x)$, FALSE otherwise. Beware that weights greater than one may inflate over-optimistically resulting p-values, use with caution.
na.rm	TRUE if missing values should be removed, FALSE otherwise.

Details

The weights w1 and w2 should represent the likelihood for each observation stored in x to belong to the first and second class, respectively. Therefore the values contained in w1 and w2 should lay in [0,1] and $\emptyset \le (w1[i] + w2[i]) \le 1$ for i in 0,1,...,n where n is the length of x. The Welch's version of the t test is implemented in this function, therefore assuming unequal sample size and unequal variance. The sample size of the first and second class are calculated as the sum(w1) and sum(w2), respectively.

Value

A numeric vector of six values that are the difference between the two weighted means, the value of the t statistic, the sample size of class 1, the sample size of class 2, the degree of freedom and the corresponding p-value.

References

http://en.wikipedia.org/wiki/T_test

See Also

stats::weighted.mean

```
set.seed(54321) # random generation of 50 normally distributed values for each of the two classes xx \leftarrow c(rnorm(50), rnorm(50)+1) # fuzzy membership to class 1 ww1 \leftarrow runif(50) + 0.3 ww1[ww1 > 1] \leftarrow 1 ww1 \leftarrow c(ww1, 1 - ww1) # fuzzy membership to class 2 ww2 \leftarrow 1 - ww1 # Welch's t test weighted by fuzzy membership to class 1 and 2
```

gene70 19

```
wt <- fuzzy.ttest(x=xx, w1=ww1, w2=ww2)
print(wt)
# Not run:
# permutation test to compute the null distribution of the weighted t statistic
wt <- wt[2]
rands <- t(sapply(1:1000, function(x,y) { return(sample(1:y)) }, y=length(xx)))
randst <- apply(rands, 1, function(x, xx, ww1, ww2)
{ return(fuzzy.ttest(x=xx, w1=ww1[x], w2=ww2[x])[2]) }, xx=xx, ww1=ww1, ww2=ww2)
ifelse(wt < 0, sum(randst <= wt), sum(randst >= wt)) / length(randst)
# End(Not run)
```

gene70

Function to compute the 70 genes prognosis profile (GENE70) as published by van't Veer et al. 2002

Description

This function computes signature scores and risk classifications from gene expression values following the algorithm used for the 70 genes prognosis profile (GENE70) as published by van't Veer et al. 2002.

Usage

```
gene70(data, annot, do.mapping = FALSE, mapping,
  std = c("none", "scale", "robust"), verbose = FALSE)
```

Arguments

da	ta	Matrix of gene expressions with samples in rows and probes in columns, dimnames being properly defined.
an	not	Matrix of annotations with at least one column named "EntrezGene.ID", dimnames being properly defined.
do	.mapping	TRUE if the mapping through Entrez Gene ids must be performed (in case of ambiguities, the most variant probe is kept for each gene), FALSE otherwise.
ma	pping	Matrix with columns "EntrezGene.ID" and "probe" used to force the mapping such that the probes are not selected based on their variance.
st	d	Standardization of gene expressions: scale for traditional standardization based on mean and standard deviation, robust for standardization based on the 0.025 and 0.975 quantiles, none to keep gene expressions unchanged.
ve	rbose	TRUE to print informative messages, FALSE otherwise.

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Value

A list with items:

- score Continuous signature scores
- risk Binary risk classification, 1 being high risk and 0 being low risk.
- mapping Mapping used if necessary.
- probe If mapping is performed, this matrix contains the correspondence between the gene list (aka signature) and gene expression data

References

L. J. van't Veer and H. Dai and M. J. van de Vijver and Y. D. He and A. A. Hart and M. Mao and H. L. Peterse and K. van der Kooy and M. J. Marton and A. T. Witteveen and G. J. Schreiber and R. M. Kerkhiven and C. Roberts and P. S. Linsley and R. Bernards and S. H. Friend (2002) "Gene Expression Profiling Predicts Clinical Outcome of Breast Cancer", Nature, 415:530–536.

See Also

nkis

Examples

```
# load GENE70 signature
data(sig.gene70)
# load NKI dataset
data(nkis)
# compute relapse score
rs.nkis <- gene70(data=data.nkis)
table(rs.nkis$risk)
# note that the discrepancies compared to the original publication
# are closed to the official cutoff, raising doubts on its exact value.
# computation of the signature scores on a different microarray platform
# load VDX dataset
data(vdxs)
# compute relapse score
rs.vdxs <- gene70(data=data.vdxs, annot=annot.vdxs, do.mapping=TRUE)
table(rs.vdxs$risk)</pre>
```

gene76

Function to compute the Relapse Score as published by Wang et al. 2005

Description

This function computes signature scores and risk classifications from gene expression values following the algorithm used for the Relapse Score (GENE76) as published by Wang et al. 2005.

geneid.map 21

Usage

```
gene76(data, er)
```

Arguments

data Matrix of gene expressions with samples in rows and probes in columns, dim-

names being properly defined.

er Vector containing the estrogen receptor (ER) status of breast cancer patients in

the dataset.

Value

A list with items:

- score Continuous signature scores
- risk Binary risk classification, 1 being high risk and 0 being low risk.

References

Y. Wang and J. G. Klijn and Y. Zhang and A. M. Sieuwerts and M. P. Look and F. Yang and D. Talantov and M. Timmermans and M. E. Meijer-van Gelder and J. Yu and T. Jatkoe and E. M. Berns and D. Atkins and J. A. Foekens (2005) "Gene-Expression Profiles to Predict Distant Metastasis of Lymph-Node-Negative Primary Breast Cancer", Lancet, 365(9460):671–679.

See Also

ggi

Examples

```
# load GENE76 signature
data(sig.gene76)
# load VDX dataset
data(vdxs)
# compute relapse score
rs.vdxs <- gene76(data=data.vdxs, er=demo.vdxs[ ,"er"])
table(rs.vdxs$risk)</pre>
```

geneid.map

Function to find the common genes between two datasets or a dataset and a gene list

Description

This function allows for fast mapping between two datasets or a dataset and a gene list. The mapping process is performed using Entrez Gene id as reference. In case of ambiguities (several probes representing the same gene), the most variant probe is selected.

22 geneid.map

Usage

```
geneid.map(geneid1, data1, geneid2, data2, verbose = FALSE)
```

Arguments

geneid1	First vector of Entrez Gene ids. The name of the vector cells must be the name of the probes in the dataset data1.
data1	First dataset with samples in rows and probes in columns. The dimnames must be properly defined.
geneid2	Second vector of Entrez Gene ids. The name of the vector cells must be the name of the probes in the dataset data1 if it is not missing, proper names must be assigned otherwise.
data2	First dataset with samples in rows and probes in columns. The dimnames must be properly defined. It may be missing.
verbose	TRUE to print informative messages, FALSE otherwise.

Value

A list with items:

- geneid1 Mapped gene list from geneid1.
- data1 Mapped dataset from data1.
- geneid2 Mapped gene list from geneid2.
- data2 Mapped dataset from data2.

Note

It is mandatory that the names of geneid1 and geneid2 must be the probe names of the microarray platform.

genius 23

genius	Function to compute the Gene Expression progNostic Index Using Subtypes (GENIUS) as published by Haibe-Kains et al. 2010

Description

This function computes the Gene Expression progNostic Index Using Subtypes (GENIUS) as published by Haibe-Kains et al. 2010. Subtype-specific risk scores are computed for each subtype signature separately and an overall risk score is computed by combining these scores with the posterior probability to belong to each of the breast cancer molecular subtypes.

Usage

```
genius(data, annot, do.mapping = FALSE, mapping, do.scale = TRUE)
```

Arguments

data	Matrix of gene expressions with samples in rows and probes in columns, dimnames being properly defined.
annot	Matrix of annotations with at least one column named "EntrezGene.ID", dimnames being properly defined.
do.mapping	TRUE if the mapping through Entrez Gene ids must be performed (in case of ambiguities, the most variant probe is kept for each gene), FALSE otherwise.
mapping	Matrix with columns "EntrezGene.ID" and "probe" used to force the mapping such that the probes are not selected based on their variance.
do.scale	TRUE if the ESR1, ERBB2 and AURKA (module) scores must be rescaled (see rescale), FALSE otherwise.

Value

A list with items:

- GENIUSM1: Risk score from the ER-/HER2- subtype signature in GENIUS model.
- GENIUSM2: Risk score from the HER2+ subtype signature in GENIUS model.
- GENIUSM3: Risk score from the ER+/HER2- subtype signature in GENIUS model.
- score: Overall risk prediction as computed by the GENIUS model.a.

References

Haibe-Kains B, Desmedt C, Rothe F, Sotiriou C and Bontempi G (2010) "A fuzzy gene expression-based computational approach improves breast cancer prognostication", Genome Biology, 11(2):R18

See Also

subtype.cluster.predict,sig.score

24 ggi

Examples

```
# load NKI dataset
data(nkis)
data(scmod1.robust)
data(sig.genius)

# compute GENIUS risk scores based on GENIUS model fitted on VDX dataset
genius.nkis <- genius(data=data.nkis, annot=annot.nkis, do.mapping=TRUE)
str(genius.nkis)
# the performance of GENIUS overall risk score predictions are not optimal
# since only part of the NKI dataset was used</pre>
```

ggi Function to compute the raw and scaled Gene expression Grade Index (GGI)

Description

This function computes signature scores and risk classifications from gene expression values following the algorithm used for the Gene expression Grade Index (GGI).

Usage

```
ggi(data, annot, do.mapping = FALSE, mapping, hg, verbose = FALSE)
```

Arguments

data	Matrix of gene expressions with samples in rows and probes in columns, dimnames being properly defined.
annot	Matrix of annotations with at least one column named "EntrezGene.ID", dimnames being properly defined.
do.mapping	TRUE if the mapping through Entrez Gene ids must be performed (in case of ambiguities, the most variant probe is kept for each gene), FALSE otherwise.
mapping	Matrix with columns "EntrezGene.ID" and "probe" used to force the mapping such that the probes are not selected based on their variance.
hg	Vector containing the histological grade (HG) status of breast cancer patients in the dataset.
verbose	TRUE to print informative messages, FALSE otherwise.

Value

A list with items:

- score: Continuous signature scores
- risk: Binary risk classification, 1 being high risk and 0 being low risk.

ihc4 25

- · mapping: Mapping used if necessary.
- probe: If mapping is performed, this matrix contains the correspondence between the gene list (aka signature) and gene expression data.

References

Sotiriou C, Wirapati P, Loi S, Harris A, Bergh J, Smeds J, Farmer P, Praz V, Haibe-Kains B, Lallemand F, Buyse M, Piccart MJ and Delorenzi M (2006) "Gene expression profiling in breast cancer: Understanding the molecular basis of histologic grade to improve prognosis", Journal of National Cancer Institute, 98:262–272

See Also

gene76

Examples

```
# load GGI signature
data(sig.ggi)
# load NKI dataset
data(nkis)
# compute relapse score
ggi.nkis <- ggi(data=data.nkis, annot=annot.nkis, do.mapping=TRUE,
    hg=demo.nkis[ ,"grade"])
table(ggi.nkis$risk)</pre>
```

ihc4

Function to compute the IHC4 prognostic score as published by Paik et al. in 2004.

Description

This function computes the prognostic score based on four measured IHC markers (ER, PGR, HER2, Ki-67), following the algorithm as published by Cuzick et al. 2011. The user has the option to either obtain just the shrinkage-adjusted IHC4 score (IHC4) or the overall score htat also combines the clinical score (IHC4+C)

Usage

```
ihc4(ER, PGR, HER2, Ki67,age,size,grade,node,ana,scoreWithClinical=FALSE, na.rm = FALSE)
```

Arguments

ER score between 0-10, calculated as (H-score/30).

PGR Progesterone Receptor score between 0-10.

HER2 Her2/neu status (0 or 1).

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Ki67 score based on percentage of positively staining malignant cells.

age patient age.

size tumor size in cm.

grade Histological grade, i.e. low (1), intermediate (2) and high (3) grade.

node Nodal status.

ana treatment with anastrozole.

scoreWithClinical

TRUE to get IHC4+C score, FALSE to get just the IHC4 score.

na.rm TRUE if missing values should be removed, FALSE otherwise.

Value

Shrinkage-adjusted IHC4 score or the Overall Prognostic Score based on IHC4+C (IHC4+Clinical Score)

References

Jack Cuzick, Mitch Dowsett, Silvia Pineda, Christopher Wale, Janine Salter, Emma Quinn, Lila Zabaglo, Elizabeth Mallon, Andrew R. Green, Ian O. Ellis, Anthony Howell, Aman U. Buzdar, and John F. Forbes (2011) "Prognostic Value of a Combined Estrogen Receptor, Progesterone Receptor, Ki-67, and Human Epidermal Growth Factor Receptor 2 Immunohistochemical Score and Comparison with the Genomic Health Recurrence Score in Early Breast Cancer", Journal of Clinical Oncologoy, 29(32):4273–4278.

```
# load NKI dataset
data(nkis)
# compute shrinkage-adjusted IHC4 score
count<-nrow(demo.nkis)
ihc4(ER=sample(x=1:10, size=count,replace=TRUE),PGR=sample(x=1:10, size=count,replace=TRUE),
HER2=sample(x=0:1,size=count,replace=TRUE),Ki67=sample(x=1:100, size=count,replace=TRUE),
scoreWithClinical=FALSE, na.rm=TRUE)
# compute IHC4+C score
ihc4(ER=sample(x=1:10, size=count,replace=TRUE),PGR=sample(x=1:10, size=count,replace=TRUE),
HER2=sample(x=0:1,size=count,replace=TRUE),Ki67=sample(x=1:100, size=count,replace=TRUE),
age=demo.nkis[,"age"],size=demo.nkis[,"size"],grade=demo.nkis[,"grade"],node=demo.nkis[,"node"],
ana=sample(x=0:1,size=count,replace=TRUE), scoreWithClinical=TRUE, na.rm=TRUE)</pre>
```

intrinsic.cluster 27

intrinsic.cluster	Function to fit a Single Sample Predictor (SSP) as in Perou, Sorlie,
The Thate. Cluster	1 inclion to fit a strigge sample I redictor (331) as in Ferou, Sortie,
	Hu, and Parker publications

Description

This function fits the Single Sample Predictor (SSP) as published in Sorlie et al 2003, Hu et al 2006 and Parker et al 2009. This model is actually a nearest centroid classifier where the centroids representing the breast cancer molecular subtypes are identified through hierarchical clustering using an "intrinsic gene list".

Usage

```
intrinsic.cluster(data, annot, do.mapping = FALSE, mapping,
  std = c("none", "scale", "robust"), rescale.q = 0.05, intrinsicg,
  number.cluster = 3, mins = 5, method.cor = c("spearman", "pearson"),
  method.centroids = c("mean", "median", "tukey"), filen, verbose = FALSE)
```

Arguments

6	
data	Matrix of gene expressions with samples in rows and probes in columns, dimnames being properly defined.
annot	Matrix of annotations with at least one column named "EntrezGene.ID", dimnames being properly defined.
do.mapping	TRUE if the mapping through Entrez Gene ids must be performed (in case of ambiguities, the most variant probe is kept for each gene), FALSE otherwise.
mapping	Matrix with columns "EntrezGene.ID" and "probe" used to force the mapping such that the probes are not selected based on their variance.
std	Standardization of gene expressions: scale for traditional standardization based on mean and standard deviation, robust for standardization based on the 0.025 and 0.975 quantiles, none to keep gene expressions unchanged.
rescale.q	Proportion of expected outliers for (robust) rescaling the gene expressions.
intrinsicg	Intrinsic gene lists. May be specified by the user as a matrix wit hat least 2 columns named probe and EntrezGene.ID for the probe names and the corresponding Entrez Gene ids. The intrinsic gene lists published by Sorlie et al. 2003, Hu et al. 2006 and Parker et al. 2009 are stored in ssp2003, ssp2006 and pam50 respectively.
number.cluster	The number of main clusters to be identified by hierarchical clustering.
mins	The minimum number of samples to be in a main cluster.
method.cor	Correlation coefficient used to identified the nearest centroid. May be spearman or pearson.
method.centroids	
	LMethod to compute a centroid from gene expressions of a cluster of samples: mean, median or tukey (Tukey's Biweight Robust Mean).
filen	Name of the csv file where the subtype clustering model must be stored.
verbose	TRUE to print informative messages, FALSE otherwise.

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Value

A list with items:

- model:Single Sample Predictor
- subtype: Subtypes identified by the SSP. For published intrinsic gene lists, subtypes can be either "Basal", "Her2", "LumA", "LumB" or "Normal".
- subtype.proba: Probabilities to belong to each subtype estimated from the correlations to each centroid.
- cor: Correlation coefficient to each centroid.

References

T. Sorlie and R. Tibshirani and J. Parker and T. Hastie and J. S. Marron and A. Nobel and S. Deng and H. Johnsen and R. Pesich and S. Geister and J. Demeter and C. Perou and P. E. Lonning and P. O. Brown and A. L. Borresen-Dale and D. Botstein (2003) "Repeated Observation of Breast Tumor Subtypes in Independent Gene Expression Data Sets", Proceedings of the National Academy of Sciences, 1(14):8418-8423 Hu, Zhiyuan and Fan, Cheng and Oh, Daniel and Marron, JS and He, Xiaping and Qaqish, Bahjat and Livasy, Chad and Carey, Lisa and Reynolds, Evangeline and ressler, Lynn and Nobel, Andrew and Parker, Joel and Ewend, Matthew and Sawyer, Lynda and Wu, Junyuan and Liu, Yudong and Nanda, Rita and Tretiakova, Maria and Orrico, Alejandra and Dreher, Donna and Palazzo, Juan and Perreard, Laurent and Nelson, Edward and Mone, Mary and Hansen, Heidi and Mullins, Michael and Quackenbush, John and Ellis, Matthew and Olopade, Olufunmilayo and Bernard, Philip and Perou, Charles (2006) "The molecular portraits of breast tumors are conserved across microarray platforms", BMC Genomics, 7(96) Parker, Joel S. and Mullins, Michael and Cheang, Maggie C.U. and Leung, Samuel and Voduc, David and Vickery, Tammi and Davies, Sherri and Fauron, Christiane and He, Xiaping and Hu, Zhiyuan and Quackenbush, John F. and Stijleman, Inge J. and Palazzo, Juan and Marron, J.S. and Nobel, Andrew B. and Mardis, Elaine and Nielsen, Torsten O. and Ellis, Matthew J. and Perou, Charles M. and Bernard, Philip S. (2009) "Supervised Risk Predictor of Breast Cancer Based on Intrinsic Subtypes", Journal of Clinical Oncology, 27(8):1160-1167

See Also

subtype.cluster, intrinsic.cluster.predict, ssp2003, ssp2006, pam50

```
# load SSP signature published in Sorlie et al. 2003
data(ssp2003)
# load NKI data
data(nkis)
# load VDX data
data(vdxs)
ssp2003.nkis <- intrinsic.cluster(data=data.nkis, annot=annot.nkis,
    do.mapping=TRUE, std="robust",
    intrinsicg=ssp2003$centroids.map[ ,c("probe", "EntrezGene.ID")],
    number.cluster=5, mins=5, method.cor="spearman",
    method.centroids="mean", verbose=TRUE)
str(ssp2003.nkis, max.level=1)</pre>
```

intrinsic.cluster.predict 29

intrinsic.cluster.predict

Function to identify breast cancer molecular subtypes using the Single Sample Predictor (SSP)

Description

This function identifies the breast cancer molecular subtypes using a Single Sample Predictor (SSP) fitted by intrinsic.cluster.

Usage

```
intrinsic.cluster.predict(sbt.model, data, annot, do.mapping = FALSE,
   mapping, do.prediction.strength = FALSE, verbose = FALSE)
```

Arguments

sbt.model	Subtype Clustering Model as returned by intrinsic.cluster.
data	Matrix of gene expressions with samples in rows and probes in columns, dimnames being properly defined.
annot	Matrix of annotations with at least one column named "EntrezGene.ID", dimnames being properly defined.
do.mapping	TRUE if the mapping through Entrez Gene ids must be performed (in case of ambiguities, the most variant probe is kept for each gene), FALSE otherwise.
mapping	Matrix with columns "EntrezGene.ID" and "probe" used to force the
do.prediction.strength	
	TRUE if the prediction strength must be computed (Tibshirani and Walther 2005), FALSE otherwise.
verbose	TRUE to print informative messages, FALSE otherwise.

Value

A list with items:

- subtype: Subtypes identified by the SSP. For published intrinsic gene lists, subtypes can be either "Basal", "Her2", "LumA", "LumB" or "Normal".
- subtype.proba: Probabilities to belong to each subtype estimated from the correlations to each centroid.
- cor: Correlation coefficient to each centroid.
- prediction.strength: Prediction strength for subtypes.
- subtype.train: Classification (similar to subtypes) computed during fitting of the model for prediction strength.
- centroids.map: Mapped probes from the intrinsic gene list used to compute the centroids.
- profiles: Intrinsic gene expression profiles for each sample.

30 map.datasets

References

T. Sorlie and R. Tibshirani and J. Parker and T. Hastie and J. S. Marron and A. Nobel and S. Deng and H. Johnsen and R. Pesich and S. Geister and J. Demeter and C. Perou and P. E. Lonning and P. O. Brown and A. L. Borresen-Dale and D. Botstein (2003) "Repeated Observation of Breast Tumor Subtypes in Independent Gene Expression Data Sets", Proceedings of the National Academy of Sciences, 1(14):8418-8423 Hu, Zhiyuan and Fan, Cheng and Oh, Daniel and Marron, JS and He, Xiaping and Qaqish, Bahjat and Livasy, Chad and Carey, Lisa and Reynolds, Evangeline and Dressler, Lynn and Nobel, Andrew and Parker, Joel and Ewend, Matthew and Sawyer, Lynda and Wu, Junyuan and Liu, Yudong and Nanda, Rita and Tretiakova, Maria and Orrico, Alejandra and Dreher, Donna and Palazzo, Juan and Perreard, Laurent and Nelson, Edward and Mone, Mary and Hansen, Heidi and Mullins, Michael and Quackenbush, John and Ellis, Matthew and Olopade, Olufunmilayo and Bernard, Philip and Perou, Charles (2006) "The molecular portraits of breast tumors are conserved across microarray platforms", BMC Genomics, 7(96) Parker, Joel S. and Mullins, Michael and Cheang, Maggie C.U. and Leung, Samuel and Voduc, David and Vickery, Tammi and Davies, Sherri and Fauron, Christiane and He, Xiaping and Hu, Zhiyuan and Quackenbush, John F. and Stijleman, Inge J. and Palazzo, Juan and Marron, J.S. and Nobel, Andrew B. and Mardis, Elaine and Nielsen, Torsten O. and Ellis, Matthew J. and Perou, Charles M. and Bernard, Philip S. (2009) "Supervised Risk Predictor of Breast Cancer Based on Intrinsic Subtypes", Journal of Clinical Oncology, 27(8):1160-1167 Tibshirani R and Walther G (2005) "Cluster Validation by Prediction Strength", Journal of Computational and Graphical Statistics, 14(3):511–528

See Also

intrinsic.cluster, ssp2003, ssp2006, pam50

Examples

```
# load SSP fitted in Sorlie et al. 2003
data(ssp2003)
# load NKI data
data(nkis)
# SSP2003 applied on NKI
ssp2003.nkis <- intrinsic.cluster.predict(sbt.model=ssp2003,
    data=data.nkis, annot=annot.nkis, do.mapping=TRUE,
    do.prediction.strength=FALSE, verbose=TRUE)
table(ssp2003.nkis$subtype)</pre>
```

map.datasets

Function to map a list of datasets through EntrezGene IDs in order to get the union of the genes

Description

This function maps a list of datasets through EntrezGene IDs in order to get the union of the genes.

map.datasets 31

Usage

```
map.datasets(datas, annots, do.mapping = FALSE,
   mapping.coln = "EntrezGene.ID", mapping, verbose = FALSE)
```

Arguments

datas	List of matrices of gene expressions with samples in rows and probes in columns, dimnames being properly defined.
annots	List of matrices of annotations with at least one column named "EntrezGene.ID", dimnames being properly defined.
do.mapping	TRUE if the mapping through Entrez Gene ids must be performed (in case of ambiguities, the most variant probe is kept for each gene), FALSE otherwise.
mapping.coln	Name of the column containing the biological annotation to be used to map the different datasets, default is "EntrezGene.ID".
mapping	Matrix with columns "EntrezGene.ID" and "probe.x" used to force the mapping such that the probes of platform x are not selected based on their variance.
verbose	TRUE to print informative messages, FALSE otherwise.

Details

In case of several probes representing the same EntrezGene ID, the most variant is selected if mapping is not specified. When a EntrezGene ID does not exist in a specific dataset, NA values are introduced.

Value

A list with items:

- datas: List of datasets (gene expression matrices)
- annots: List of annotations (annotation matrices)

```
# load VDX dataset
data(vdxs)
# load NKI dataset
data(nkis)
# reduce datasets
ginter <- intersect(annot.vdxs[ ,"EntrezGene.ID"], annot.nkis[ ,"EntrezGene.ID"])</pre>
ginter <- ginter[!is.na(ginter)][1:30]</pre>
myx <- unique(c(match(ginter, annot.vdxs[ ,"EntrezGene.ID"]),</pre>
  sample(x=1:nrow(annot.vdxs), size=20)))
data2.vdxs <- data.vdxs[ ,myx]</pre>
annot2.vdxs <- annot.vdxs[myx, ]</pre>
myx <- unique(c(match(ginter, annot.nkis[ ,"EntrezGene.ID"]),</pre>
  sample(x=1:nrow(annot.nkis), size=20)))
data2.nkis <- data.nkis[ ,myx]</pre>
annot2.nkis <- annot.nkis[myx, ]</pre>
# mapping of datasets
```

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```
datas <- list("VDX"=data2.vdxs,"NKI"=data2.nkis)
annots <- list("VDX"=annot2.vdxs, "NKI"=annot2.nkis)
datas.mapped <- map.datasets(datas=datas, annots=annots, do.mapping=TRUE)
str(datas.mapped, max.level=2)</pre>
```

medianCtr

Center around the median

Description

Utility function called within the claudinLow classifier

Usage

```
medianCtr(x)
```

Arguments

Х

Matrix of numbers

Value

A matrix of median-centered numbers

References

```
citation("claudinLow")
```

See Also

claudinLow

mod1

Gene modules published in Desmedt et al. 2008

Description

List of seven gene modules published in Desmedt et a. 2008, i.e. ESR1 (estrogen receptor pathway), ERBB2 (her2/neu receptor pathway), AURKA (proliferation), STAT1 (immune response), PLAU (tumor invasion), VEGF (angogenesis) and CASP3 (apoptosis).

Usage

```
data(mod1)
```

mod2 33

Details

mod1 is a list of seven gene signatures, i.e. matrices with 3 columns containing the annotations and information related to the signatures themselves.

References

Desmedt C, Haibe-Kains B, Wirapati P, Buyse M, Larsimont D, Bontempi G, Delorenzi M, Piccart M, and Sotiriou C (2008) "Biological processes associated with breast cancer clinical outcome depend on the molecular subtypes", Clinical Cancer Research, 14(16):5158–5165.

mod2

Gene modules published in Wirapati et al. 2008

Description

List of seven gene modules published in Wirapati et a. 2008, i.e. ESR1 (estrogen receptor pathway), ERBB2 (her2/neu receptor pathway) and AURKA (proliferation).

Usage

data(mod2)

Details

mod2 is a list of three gene signatures, i.e. matrices with 3 columns containing the annotations and information related to the signatures themselves.

Source

http://breast-cancer-research.com/content/10/4/R65

References

Wirapati P, Sotiriou C, Kunkel S, Farmer P, Pradervand S, Haibe-Kains B, Desmedt C, Ignatiadis M, Sengstag T, Schutz F, Goldstein DR, Piccart MJ and Delorenzi M (2008) "Meta-analysis of Gene-Expression Profiles in Breast Cancer: Toward a Unified Understanding of Breast Cancer Sub-typing and Prognosis Signatures", Breast Cancer Research, 10(4):R65.

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modelOvcAngiogenic Model used to classify ovarian tumors into A genic subtypes.	Angiogenic and NonAngio-
---	--------------------------

Description

Object containing the set of parameters for the mixture of Gaussians used as a model to classify ovarian tumors into Angiogenic and NonAngiogenic subtypes.

Usage

```
data(modelOvcAngiogenic)
```

Source

http://jnci.oxfordjournals.org/cgi/content/full/98/4/262/DC1

References

Bentink S, Haibe-Kains B, Risch T, Fan J-B, Hirsch MS, Holton K, Rubio R, April C, Chen J, Wickham-Garcia E, Liu J, Culhane AC, Drapkin R, Quackenbush JF, Matulonis UA (2012) "Angiogenic mRNA and microRNA Gene Expression Signature Predicts a Novel Subtype of Serous Ovarian Cancer", PloS one, 7(2):e30269

```
{\it molecular.subtyping} \qquad {\it Function\ to\ identify\ breast\ cancer\ molecular\ subtypes\ using\ the\ Subtype\ Clustering\ Model}
```

Description

This function identifies the breast cancer molecular subtypes using a Subtype Clustering Model fitted by subtype.cluster.

Usage

```
molecular.subtyping(sbt.model = c("scmgene", "scmod1", "scmod2",
   "pam50", "ssp2006", "ssp2003", "intClust", "AIMS","claudinLow"),
   data, annot, do.mapping = FALSE, verbose = FALSE)
```

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Arguments

Subtyping classification model, can be either "scmgene", "scmod1", "scmod2", "pam50", "ssp2006", "ssp2003", "intClust", "AIMS", or "claudinLow".

Matrix of gene expressions with samples in rows and probes in columns, dimnames being properly defined.

Matrix of annotations with at least one column named "EntrezGene.ID" (for ssp, scm, AIMS, and claudinLow models) or "Gene.Symbol" (for the intClust model), dimnames being properly defined.

do.mapping TRUE if the mapping through Entrez Gene ids must be performed (in case of

ambiguities, the most variant probe is kept for each gene), FALSE otherwise.

verbose TRUE if informative messages should be displayed, FALSE otherwise.

Value

A list with items:

• subtype: Subtypes identified by the subtyping classification model.

- subtype.proba: Probabilities to belong to each subtype estimated by the subtyping classification model.
- subtype.crisp: Crisp classes identified by the subtyping classification model.

References

T. Sorlie and R. Tibshirani and J. Parker and T. Hastie and J. S. Marron and A. Nobel and S. Deng and H. Johnsen and R. Pesich and S. Geister and J. Demeter and C. Perou and P. E. Lonning and P. O. Brown and A. L. Borresen-Dale and D. Botstein (2003) "Repeated Observation of Breast Tumor Subtypes in Independent Gene Expression Data Sets", Proceedings of the National Academy of Sciences, 1(14):8418-8423 Hu, Zhiyuan and Fan, Cheng and Oh, Daniel and Marron, JS and He, Xiaping and Qaqish, Bahjat and Livasy, Chad and Carey, Lisa and Reynolds, Evangeline and Dressler, Lynn and Nobel, Andrew and Parker, Joel and Ewend, Matthew and Sawyer, Lynda and Wu, Junyuan and Liu, Yudong and Nanda, Rita and Tretiakova, Maria and Orrico, Alejandra and Dreher, Donna and Palazzo, Juan and Perreard, Laurent and Nelson, Edward and Mone, Mary and Hansen, Heidi and Mullins, Michael and Quackenbush, John and Ellis, Matthew and Olopade, Olufunmilayo and Bernard, Philip and Perou, Charles (2006) "The molecular portraits of breast tumors are conserved across microarray platforms", BMC Genomics, 7(96) Parker, Joel S. and Mullins, Michael and Cheang, Maggie C.U. and Leung, Samuel and Voduc, David and Vickery, Tammi and Davies, Sherri and Fauron, Christiane and He, Xiaping and Hu, Zhiyuan and Quackenbush, John F. and Stijleman, Inge J. and Palazzo, Juan and Marron, J.S. and Nobel, Andrew B. and Mardis, Elaine and Nielsen, Torsten O. and Ellis, Matthew J. and Perou, Charles M. and Bernard, Philip S. (2009) "Supervised Risk Predictor of Breast Cancer Based on Intrinsic Subtypes", Journal of Clinical Oncology, 27(8):1160-1167 Desmedt C, Haibe-Kains B, Wirapati P, Buyse M, Larsimont D, Bontempi G, Delorenzi M, Piccart M, and Sotiriou C (2008) "Biological processes associated with breast cancer clinical outcome depend on the molecular subtypes", Clinical Cancer Research, 14(16):5158-5165. Wirapati P, Sotiriou C, Kunkel S, Farmer P, Pradervand S, Haibe-Kains B, Desmedt C, Ignatiadis M, Sengstag T, Schutz F, Goldstein DR, Piccart MJ and Delorenzi M (2008) "Metaanalysis of Gene-Expression Profiles in Breast Cancer: Toward a Unified Understanding of Breast Cancer Sub-typing and Prognosis Signatures", Breast Cancer Research, 10(4):R65. Haibe-Kains 36 molecular.subtyping

B, Desmedt C, Loi S, Culhane AC, Bontempi G, Quackenbush J, Sotiriou C. (2012) "A three-gene model to robustly identify breast cancer molecular subtypes.", J Natl Cancer Inst., 104(4):311-325. Curtis C, Shah SP, Chin SF, Turashvili G, Rueda OM, Dunning MJ, Speed D, Lynch AG, Samarajiwa S, Yuan Y, Graf S, Ha G, Haffari G, Bashashati A, Russell R, McKinney S; METABRIC Group, Langerod A, Green A, Provenzano E, Wishart G, Pinder S, Watson P, Markowetz F, Murphy L, Ellis I, Purushotham A, Borresen-Dale AL, Brenton JD, Tavare S, Caldas C, Aparicio S. (2012) "The genomic and transcriptomic architecture of 2,000 breast tumours reveals novel subgroups.", Nature, 486(7403):346-352. Paquet ER, Hallett MT. (2015) "Absolute assignment of breast cancer intrinsic molecular subtype.", J Natl Cancer Inst., 107(1):357. Aleix Prat, Joel S Parker, Olga Karginova, Cheng Fan, Chad Livasy, Jason I Herschkowitz, Xiaping He, and Charles M. Perou (2010) "Phenotypic and molecular characterization of the claudin-low intrinsic subtype of breast cancer", Breast Cancer Research, 12(5):R68

See Also

subtype.cluster.predict, intrinsic.cluster.predict

```
##### without mapping (affy hgu133a or plus2 only)
# load VDX data
require(iC10TrainingData)
data(vdxs)
data(AIMSmodel)
data(scmgene.robust)
# Subtype Clustering Model fitted on EXPO and applied on VDX
sbt.vdx.SCMGENE <- molecular.subtyping(sbt.model="scmgene",</pre>
 data=data.vdxs, annot=annot.vdxs, do.mapping=FALSE)
table(sbt.vdx.SCMGENE$subtype)
# Using the AIMS molecular subtyping algorithm
sbt.vdxs.AIMS <- molecular.subtyping(sbt.model="AIMS", data=data.vdxs,</pre>
                                      annot=annot.vdxs, do.mapping=FALSE)
table(sbt.vdxs.AIMS$subtype)
# Using the IntClust molecular subtyping algorithm
colnames(annot.vdxs)[3]<-"Gene.Symbol"</pre>
sbt.vdxs.intClust <- molecular.subtyping(sbt.model="intClust", data=data.vdxs,</pre>
 annot=annot.vdxs, do.mapping=FALSE)
table(sbt.vdxs.intClust$subtype)
##### with mapping
# load NKI data
data(nkis)
# Subtype Clustering Model fitted on EXPO and applied on NKI
sbt.nkis <- molecular.subtyping(sbt.model="scmgene", data=data.nkis,</pre>
 annot=annot.nkis, do.mapping=TRUE)
table(sbt.nkis$subtype)
##### with mapping
```

nkis 37

```
## load vdxs data
data(vdxs)
data(claudinLowData)

## Claudin-Low classification of 150 VDXS samples
sbt.vdxs.CL <- molecular.subtyping(sbt.model="claudinLow", data=data.vdxs,
    annot=annot.vdxs, do.mapping=TRUE)
table(sbt.vdxs.CL$subtype)</pre>
```

nkis

Gene expression, annotations and clinical data from van de Vijver et al. 2002

Description

This dataset contains (part of) the gene expression, annotations and clinical data as published in van de Vijver et al. 2002.

Usage

data(nkis)

Format

nkis is a dataset containing three matrices:

- data.nkis: Matrix containing gene expressions as measured by Agilent technology (dual-channel, oligonucleotides)
- annot.nkis: Matrix containing annotations of Agilent microarray platform
- · demon.nkis: Clinical information of the breast cancer patients whose tumors were hybridized

Details

This dataset represent only partially the one published by van de Vijver et al. in 2008. Indeed, only part of the patients (150) and gene expressions (922) in data.nkis.

Source

http://www.nature.com/nature/journal/v415/n6871/full/415530a.html

References

M. J. van de Vijver and Y. D. He and L. van't Veer and H. Dai and A. M. Hart and D. W. Voskuil and G. J. Schreiber and J. L. Peterse and C. Roberts and M. J. Marton and M. Parrish and D. Atsma and A. Witteveen and A. Glas and L. Delahaye and T. van der Velde and H. Bartelink and S. Rodenhuis and E. T. Rutgers and 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

npi

npi

Function to compute the Nottingham Prognostic Index

Description

This function computes the Nottingham Prognostic Index (NPI) as published in Galeat et al, 1992. NPI is a clinical index shown to be highly prognostic in breast cancer.

Usage

```
npi(size, grade, node, na.rm = FALSE)
```

Arguments

size tumor size in cm.
grade Histological grade, i.e. low (1), intermediate (2) and high (3) grade.

node Nodal status. If only binary nodal status (0/1) is available, map 0 to 1 and 1 to 3.

TRUE if missing values should be removed, FALSE otherwise.

Details

na.rm

The risk prediction is either Good if score < 3.4, Intermediate if $3.4 \le$ score < 5.4, or Poor if score > 5.4.

Value

A list with items:

- score: Continuous signature scores
- risk: Binary risk classification, 1 being high risk and 0 being low risk.

References

Galea MH, Blamey RW, Elston CE, and Ellis IO (1992) "The nottingham prognostic index in primary breast cancer", Breast Cancer Reasearch and Treatment, 22(3):207-219.

See Also

st.gallen

Examples

```
# load NKI dataset
data(nkis)
# compute NPI score and risk classification
npi(size=demo.nkis[ ,"size"], grade=demo.nkis[ ,"grade"],
    node=ifelse(demo.nkis[ ,"node"] == 0, 1, 3), na.rm=TRUE)
```

oncotypedx 39

oncotypedx Function et al. in	n to compute the OncotypeDX signature as published by Paik 2004.
-------------------------------	--

Description

This function computes signature scores and risk classifications from gene expression values following the algorithm used for the OncotypeDX signature as published by Paik et al. 2004.

Usage

```
oncotypedx(data, annot, do.mapping = FALSE, mapping, do.scaling=TRUE,
  verbose = FALSE)
```

Arguments

data	Matrix of gene expressions with samples in rows and probes in columns, dimnames being properly defined.
annot	Matrix of annotations with at least one column named "EntrezGene.ID", dimnames being properly defined.
do.mapping	TRUE if the mapping through Entrez Gene ids must be performed (in case of ambiguities, the most variant probe is kept for each gene), FALSE otherwise. Note that for Affymetrix HGU datasets, the mapping is not necessary.
mapping	Matrix with columns "EntrezGene.ID" and "probe" used to force the mapping such that the probes are not selected based on their variance.
do.scaling	Should the data be scaled?
verbose	TRUE to print informative messages, FALSE otherwise.

Details

Note that for Affymetrix HGU datasets, the mapping is not necessary.

Value

A list with items:

- score: Continuous signature scores
- risk: Binary risk classification, 1 being high risk and 0 being low risk.
- mapping: Mapping used if necessary.
- probe: If mapping is performed, this matrix contains the correspondence between the gene list (aka signature) and gene expression data.

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References

S. Paik, S. Shak, G. Tang, C. Kim, J. Bakker, M. Cronin, F. L. Baehner, M. G. Walker, D. Watson, T. Park, W. Hiller, E. R. Fisher, D. L. Wickerham, J. Bryant, and N. Wolmark (2004) "A Multigene Assay to Predict Recurrence of Tamoxifen-Treated, Node-Negative Breast Cancer", New England Journal of Medicine, 351(27):2817-2826.

Examples

```
# load GENE70 signature
data(sig.oncotypedx)
# load NKI dataset
data(nkis)
# compute relapse score
rs.nkis <- oncotypedx(data=data.nkis, annot=annot.nkis, do.mapping=TRUE)
table(rs.nkis$risk)</pre>
```

ovcAngiogenic

Function to compute the subtype scores and risk classifications for the angiogenic molecular subtype in ovarian cancer

Description

This function computes subtype scores and risk classifications from gene expression values following the algorithm developed by Bentink, Haibe-Kains et al. to identify the angiogenic molecular subtype in ovarian cancer.

Usage

```
ovcAngiogenic(data, annot, hgs,
gmap = c("entrezgene", "ensembl_gene_id", "hgnc_symbol", "unigene"),
do.mapping = FALSE, verbose = FALSE)
```

Arguments

data	Matrix of gene expressions with samples in rows and probes in columns, dimnames being properly defined.
annot	Matrix of annotations with one column named as gmap, dimnames being properly defined.
hgs	vector of booleans with TRUE represents the ovarian cancer patients who have a high grade, late stage, serous tumor, FALSE otherwise. This is particularly important for properly rescaling the data. If hgs is missing, all the patients will be used to rescale the subtype score.
gmap	character string containing the biomaRt attribute to use for mapping if do.mapping=TRUE
do.mapping	TRUE if the mapping through Entrez Gene ids must be performed (in case of ambiguities, the most variant probe is kept for each gene), FALSE otherwise.
verbose	TRUE to print informative messages, FALSE otherwise.

ovcCrijns 41

Value

A list with items:

- score: Continuous signature scores.
- risk: Binary risk classification, 1 being high risk and 0 being low risk.
- mapping: Mapping used if necessary.
- probe: If mapping is performed, this matrix contains the correspondence between the gene list (aka signature) and gene expression data.
- subtype: data frame reporting the subtype score, maximum likelihood classification and corresponding subtype probabilities.

References

Bentink S, Haibe-Kains B, Risch T, Fan J-B, Hirsch MS, Holton K, Rubio R, April C, Chen J, Wickham-Garcia E, Liu J, Culhane AC, Drapkin R, Quackenbush JF, Matulonis UA (2012) "Angiogenic mRNA and microRNA Gene Expression Signature Predicts a Novel Subtype of Serous Ovarian Cancer", PloS one, 7(2):e30269

See Also

sigOvcAngiogenic

Examples

```
# load the ovcAngiogenic signature

# load NKI dataset
data(nkis)
colnames(annot.nkis)[is.element(colnames(annot.nkis), "EntrezGene.ID")] <-
    "entrezgene"

# compute relapse score
ovcAngiogenic.nkis <- ovcAngiogenic(data=data.nkis, annot=annot.nkis,
    gmap="entrezgene", do.mapping=TRUE)
table(ovcAngiogenic.nkis$risk)</pre>
```

ovcCrijns

Function to compute the subtype scores and risk classifications for the prognostic signature published by Crinjs et al.

Description

This function computes subtype scores and risk classifications from gene expression values using the weights published by Crijns et al.

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Usage

```
ovcCrijns(data, annot, hgs,
  gmap = c("entrezgene", "ensembl_gene_id", "hgnc_symbol", "unigene"),
  do.mapping = FALSE, verbose = FALSE)
```

Arguments

data Matrix of gene expressions with samples in rows and probes in columns, dim-

names being properly defined.

annot Matrix of annotations with one column named as gmap, dimnames being prop-

erly defined.

hgs vector of booleans with TRUE represents the ovarian cancer patients who have

a high grade, late stage, serous tumor, FALSE otherwise. This is particularly important for properly rescaling the data. If hgs is missing, all the patients will

be used to rescale the subtype score.

gmap character string containing the biomaRt attribute to use for mapping if do.mapping=TRUE

do.mapping TRUE if the mapping through Entrez Gene ids must be performed (in case of

ambiguities, the most variant probe is kept for each gene), FALSE otherwise.

verbose TRUE to print informative messages, FALSE otherwise.

Details

Note that the original algorithm has not been implemented as it necessitates refitting of the model weights in each new dataset. However the current implementation should give similar results.

Value

A list with items:

- score: Continuous signature scores.
- risk: Binary risk classification, 1 being high risk and 0 being low risk.
- mapping: Mapping used if necessary.
- probe: If mapping is performed, this matrix contains the correspondence. between the gene list (aka signature) and gene expression data.

References

Crijns APG, Fehrmann RSN, de Jong S, Gerbens F, Meersma G J, Klip HG, Hollema H, Hofstra RMW, te Meerman GJ, de Vries EGE, van der Zee AGJ (2009) "Survival-Related Profile, Pathways, and Transcription Factors in Ovarian Cancer" PLoS Medicine, 6(2):e1000024.

See Also

sigOvcCrijns

ovcTCGA 43

Examples

```
# load the ovsCrijns signature
data(sigOvcCrijns)
# load NKI dataset
data(nkis)
colnames(annot.nkis)[is.element(colnames(annot.nkis), "EntrezGene.ID")] <-
    "entrezgene"
# compute relapse score
ovcCrijns.nkis <- ovcCrijns(data=data.nkis, annot=annot.nkis,
    gmap="entrezgene", do.mapping=TRUE)
table(ovcCrijns.nkis$risk)</pre>
```

ovcTCGA

Function to compute the prediction scores and risk classifications for the ovarian cancer TCGA signature

Description

This function computes signature scores and risk classifications from gene expression values following the algorithm developed by the TCGA consortium for ovarian cancer.

Usage

```
ovcTCGA(data, annot,
  gmap = c("entrezgene", "ensembl_gene_id", "hgnc_symbol", "unigene"),
  do.mapping = FALSE, verbose = FALSE)
```

Arguments

data	Matrix of gene expressions with samples in rows and probes in columns, dimnames being properly defined.
annot	Matrix of annotations with one column named as gmap, dimnames being properly defined.
gmap	character string containing the biomaRt attribute to use for mapping if do.mapping=TRUE
do.mapping	TRUE if the mapping through Entrez Gene ids must be performed (in case of ambiguities, the most variant probe is kept for each gene), FALSE otherwise.
verbose	TRUE to print informative messages, FALSE otherwise.

Value

A list with items:

- score: Continuous signature scores.
- risk: Binary risk classification, 1 being high risk and 0 being low risk.
- mapping: Mapping used if necessary.
- probe: If mapping is performed, this matrix contains the correspondence between the gene list (aka signature) and gene expression data.

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References

Bell D, Berchuck A, Birrer M et al. (2011) "Integrated genomic analyses of ovarian carcinoma", Nature, 474(7353):609-615

See Also

```
sigOvcTCGA
```

Examples

```
# load the ovcTCGA signature
data(sigOvcTCGA)
# load NKI dataset
data(nkis)
colnames(annot.nkis)[is.element(colnames(annot.nkis), "EntrezGene.ID")] <- "entrezgene"
# compute relapse score
ovcTCGA.nkis <- ovcTCGA(data=data.nkis, annot=annot.nkis, gmap="entrezgene", do.mapping=TRUE)
table(ovcTCGA.nkis$risk)</pre>
```

ovcYoshihara

Function to compute the subtype scores and risk classifications for the prognostic signature published by Yoshihara et al.

Description

This function computes subtype scores and risk classifications from gene expression values following the algorithm developed by Yoshihara et al, for prognosis in ovarian cancer.

Usage

```
ovcYoshihara(data, annot, hgs,
  gmap = c("entrezgene", "ensembl_gene_id", "hgnc_symbol", "unigene", "refseq_mrna"),
  do.mapping = FALSE, verbose = FALSE)
```

Arguments

data	Matrix of gene expressions with samples in rows and probes in columns, dimnames being properly defined.
annot	Matrix of annotations with one column named as gmap, dimnames being properly defined.
hgs	vector of booleans with TRUE represents the ovarian cancer patients who have a high grade, late stage, serous tumor, FALSE otherwise. This is particularly important for properly rescaling the data. If hgs is missing, all the patients will be used to rescale the subtype score.
gmap	character string containing the biomaRt attribute to use for mapping if do.mapping=TRUE

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do.mapping TRUE if the mapping through Entrez Gene ids must be performed (in case of

ambiguities, the most variant probe is kept for each gene), FALSE otherwise.

verbose TRUE to print informative messages, FALSE otherwise.

Value

A list with items:

• score: Continuous signature scores.

• risk: Binary risk classification, 1 being high risk and 0 being low risk.

• mapping: Mapping used if necessary.

• probe: If mapping is performed, this matrix contains the correspondence between the gene list (aka signature) and gene expression data.

References

Yoshihara K, Tajima A, Yahata T, Kodama S, Fujiwara H, Suzuki M, Onishi Y, Hatae M, Sueyoshi K, Fujiwara H, Kudo, Yoshiki, Kotera K, Masuzaki H, Tashiro H, Katabuchi H, Inoue I, Tanaka K (2010) "Gene expression profile for predicting survival in advanced-stage serous ovarian cancer across two independent datasets", PloS one, 5(3):e9615.

See Also

sigOvcYoshihara

Examples

```
# load the ovcYoshihara signature
data(sigOvcYoshihara)
# load NKI dataset
data(nkis)
colnames(annot.nkis)[is.element(colnames(annot.nkis), "EntrezGene.ID")] <- "entrezgene"
# compute relapse score
ovcYoshihara.nkis <- ovcYoshihara(data=data.nkis,
    annot=annot.nkis, gmap="entrezgene", do.mapping=TRUE)
table(ovcYoshihara.nkis$risk)</pre>
```

overlapSets

Overlap two datasets

Description

Utility function called within the claudinLow classifien.

Usage

```
overlapSets(x,y)
```

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Arguments

x Matrix1y Matrix2

Value

A list of overlapped dataset

References

```
citation("claudinLow")
```

See Also

claudinLow

pam50

PAM50 classifier for identification of breast cancer molecular subtypes (Parker et al 2009)

Description

List of parameters defining the PAM50 classifier for identification of breast cancer molecular subtypes (Parker et al 2009).

Usage

```
data(pam50)
data(pam50.scale)
data(pam50.robust)
```

Format

List of parameters for PAM50:

- centroids: Gene expression centroids for each subtype.
- centroids.map: Mapping for centroids.
- method.cor: Method of correlation used to compute distance to the centroids.
- method.centroids: Method used to compute the centroids.
- std: Method of standardization for gene expressions ("none", "scale" or "robust")
- mins: Minimum number of samples within each cluster allowed during the fitting of the model.

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Details

Three versions of the model are provided, each of ones differs by the gene expressions standardization method since it has an important impact on the subtype classification:

- pam50: Use of the official centroids without scaling of the gene expressions.
- pam50.scale: Use of the official centroids with traditional scaling of the gene expressions (see base::scale())
- pam50.robust: Use of the official centroids with robust scaling of the gene expressions (see rescale()) The model 'pam50.robust" has been shown to reach the best concordance with the traditional clinical parameters (ER IHC, HER2 IHC/FISH and histological grade). However the use of this model is recommended only when the dataset is representative of a global population of breast cancer patients (no sampling bias, the 5 subtypes should be present).

Source

http://jco.ascopubs.org/cgi/content/short/JCO.2008.18.1370v1

References

Parker, Joel S. and Mullins, Michael and Cheang, Maggie C.U. and Leung, Samuel and Voduc, David and Vickery, Tammi and Davies, Sherri and Fauron, Christiane and He, Xiaping and Hu, Zhiyuan and Quackenbush, John F. and Stijleman, Inge J. and Palazzo, Juan and Marron, J.S. and Nobel, Andrew B. and Mardis, Elaine and Nielsen, Torsten O. and Ellis, Matthew J. and Perou, Charles M. and Bernard, Philip S. (2009) "Supervised Risk Predictor of Breast Cancer Based on Intrinsic Subtypes", Journal of Clinical Oncology, 27(8):1160–1167

pik3cags	Function to compute the PIK3CA gene signature (PIK3CA-GS)

Description

This function computes signature scores from gene expression values following the algorithm used for the PIK3CA gene signature (PIK3CA-GS).

Usage

```
pik3cags(data, annot, do.mapping = FALSE, mapping, verbose = FALSE)
```

Arguments

data	Matrix of gene expressions with samples in rows and probes in columns, dimnames being properly defined.
annot	Matrix of annotations with at least one column named "EntrezGene.ID", dimnames being properly defined.
do.mapping	TRUE if the mapping through Entrez Gene ids must be performed (in case of ambiguities, the most variant probe is kept for each gene), FALSE otherwise.

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mapping Matrix with columns "EntrezGene.ID" and "probe" used to force the mapping

such that the probes are not selected based on their variance.

verbose TRUE to print informative messages, FALSE otherwise.

Value

Vector of signature scores for PIK3CA-GS

References

Loi S, Haibe-Kains B, Majjaj S, Lallemand F, Durbecq V, Larsimont D, Gonzalez-Angulo AM, Pusztai L, Symmans FW, Bardelli A, Ellis P, Tutt AN, Gillett CE, Hennessy BT., Mills GB, Phillips WA, Piccart MJ, Speed TP, McArthur GA, Sotiriou C (2010) "PIK3CA mutations associated with gene signature of low mTORC1 signaling and better outcomes in estrogen receptor-positive breast cancer", Proceedings of the National Academy of Sciences, 107(22):10208-10213

See Also

gene76

Examples

```
# load GGI signature
data(sig.pik3cags)
# load NKI dataset
data(nkis)
# compute relapse score
pik3cags.nkis <- pik3cags(data=data.nkis, annot=annot.nkis, do.mapping=TRUE)
head(pik3cags.nkis)</pre>
```

power.cor

Function for sample size calculation for correlation coefficients

Description

This function enables to compute the sample size requirements for estimating pearson, kendall and spearman correlations

Usage

```
power.cor(rho, w, alpha = 0.05, method = c("pearson", "kendall", "spearman"))
```

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Arguments

rho	Correaltion coefficients rho (Pearson, Kendall or Spearman)
W	a numerical vector of weights of the same length as x giving the weights to use for elements of x in the first class.
alpha	alpha level
method	a character string specifying the method to compute the correlation coefficient, must be one of "pearson" (default), "kendall" or "spearman". You can specify

just the initial letter.

Value

sample size requirement

References

Bonett, D. G., and Wright, T. A. (2000). Sample size requirements for estimating pearson, kendall and spearman correlations. Psychometrika, 65(1), 23-28. doi:10.1007/BF02294183

Examples

```
power.cor(rho=0.5, w=0.1, alpha=0.05, method="spearman")
```

ps.cluster Function to compute the prediction strength of a clustering model

Description

This function computes the prediction strength of a clustering model as published in R. Tibshirani and G. Walther 2005.

Usage

```
ps.cluster(cl.tr, cl.ts, na.rm = FALSE)
```

Arguments

cl.tr	Clusters membership as defined by the original clustering model, i.e. the one that was not fitted on the dataset of interest.
cl.ts	Clusters membership as defined by the clustering model fitted on the dataset of interest.
na.rm	TRUE if missing values should be removed, FALSE otherwise.

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Value

A list with items:

- ps: the overall prediction strength (minimum of the prediction strengths at cluster level).
- ps.cluster: Prediction strength for each cluster
- ps.individual: Prediction strength for each sample.

References

R. Tibshirani and G. Walther (2005) "Cluster Validation by Prediction Strength", Journal of Computational and Graphical Statistics, 14(3):511-528.

Examples

```
# load SSP signature published in Sorlie et al. 2003
data(ssp2003)
# load NKI data
data(nkis)
# SP2003 fitted on NKI
ssp2003.2nkis <- intrinsic.cluster(data=data.nkis, annot=annot.nkis,</pre>
 do.mapping=TRUE, std="robust",
 intrinsicg=ssp2003$centroids.map[ ,c("probe", "EntrezGene.ID")],
 number.cluster=5, mins=5, method.cor="spearman",
 method.centroids="mean", verbose=TRUE)
# SP2003 published in Sorlie et al 2003 and applied in VDX
ssp2003.nkis <- intrinsic.cluster.predict(sbt.model=ssp2003,</pre>
 data=data.nkis, annot=annot.nkis, do.mapping=TRUE, verbose=TRUE)
# prediction strength of sp2003 clustering model
ps.cluster(cl.tr=ssp2003.2nkis$subtype, cl.ts=ssp2003.nkis$subtype,
 na.rm = FALSE)
```

read.m.file

Function to read a 'csv' file containing gene lists (aka gene signatures)

Description

This function allows for reading a 'csv' file containing gene signatures. Each gene signature is composed of at least four columns: "gene.list" is the name of the signature on the first line and empty fields below, "probes" are the probe names, "EntrezGene.ID" are the EntrezGene IDs and "coefficient" are the coefficients of each probe.

Usage

```
read.m.file(file, ...)
```

readArray 51

Arguments

. . .

file Filename of the 'csv' file. Additional parameters for read.csv function.

Value

List of gene signatures.

See Also

```
mod1, mod2, 'extdata/desmedt2008_genemodules.csv', 'extdata/haibekains2009_sig_genius.csv'
```

Examples

```
# read the seven gene modules as published in Desmedt et al 2008
genemods <- read.m.file(system.file("extdata/desmedt2008_genemodules.csv",</pre>
 package = "genefu"))
str(genemods, max.level=1)
# read the three subtype signtaures from GENIUS
geniusm <- read.m.file(system.file("extdata/haibekains2009_sig_genius.csv",</pre>
 package = "genefu"))
str(geniusm, max.level=1)
```

readArray

Overlap two datasets

Description

Formatting function to read arrays and format for use in the claudinLow classifier.

Usage

```
readArray(dataFile,designFile=NA,hr=1,impute=TRUE,method="mean")
```

Arguments

dataFile file with matrix to be read.

designFile Design of file.

Header rows as Present (2) or Absent (1). hr

impute whether data will be imputed or not.

method Default method is "mean".

Value

A list

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References

```
citation("claudinLow")
```

See Also

claudinLow

rename.duplicate

Function to rename duplicated strings

Description

This function renames duplicated strings by adding their number of occurrences at the end.

Usage

```
rename.duplicate(x, sep = "_", verbose = FALSE)
```

Arguments

x vector of strings.

sep a character to be the separator between the number added at the end and the

string itself.

verbose TRUE to print informative messages, FALSE otherwise.

Value

A list with items:

- new.x: new strings (without duplicates).
- duplicated.x: strings which were originally duplicated.

Examples

```
nn <- sample(letters[1:10], 30, replace=TRUE)
table(nn)
rename.duplicate(x=nn, verbose=TRUE)</pre>
```

rescale 53

rescale

Function to rescale values based on quantiles

Description

This function rescales values x based on quantiles specified by the user such that x' = (x - q1) / (q2 - q1) where q is the specified quantile, q1 = q / 2, q2 = 1 - q/2) and x' are the new rescaled values.

Usage

```
rescale(x, na.rm = FALSE, q = 0)
```

Arguments

x The matrix or vector to rescale.

na.rm TRUE if missing values should be removed, FALSE otherwise.
q Quantile (must lie in [0,1]).

Details

In order to rescale gene expressions, q = 0.05 yielded comparable scales in numerous breast cancer microarray datasets (data not shown). The rational behind this is that, in general, 'extreme cases' (e.g. low and high proliferation, high and low expression of ESR1, ...) are often present in microarray datasets, making the estimation of 'extreme' quantiles quite stable. This is specially true for genes exhibiting some multi-modality like ESR1 or ERBB2.

Value

A vector of rescaled values with two attributes q1 and q1 containing the values of the lower and the upper quantiles respectively.

See Also

```
base::scale()
```

Examples

```
# load VDX dataset
data(vdxs)
# load NKI dataset
data(nkis)
# example of rescaling for ESR1 expression
par(mfrow=c(2,2))
hist(data.vdxs[ ,"205225_at"], xlab="205225_at", breaks=20,
    main="ESR1 in VDX")
hist(data.nkis[ ,"NM_000125"], xlab="NM_000125", breaks=20,
    main="ESR1 in NKI")
hist((rescale(x=data.vdxs[ ,"205225_at"], q=0.05) - 0.5) * 2,
```

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```
xlab="205225_at", breaks=20, main="ESR1 in VDX\nrescaled")
hist((rescale(x=data.nkis[ ,"NM_000125"], q=0.05) - 0.5) * 2,
xlab="NM_000125", breaks=20, main="ESR1 in NKI\nrescaled")
```

rorS Function to compute the rorS signature as published by Parker et al 2009

Description

This function computes signature scores and risk classifications from gene expression values following the algorithm used for the rorS signature as published by Parker et al 2009.

Usage

```
rorS(data, annot, do.mapping = FALSE, mapping, verbose = FALSE)
```

Arguments

data	Matrix of gene expressions with samples in rows and probes in columns, dimnames being properly defined.
annot	Matrix of annotations with at least one column named "EntrezGene.ID", dimnames being properly defined.
do.mapping	TRUE if the mapping through Entrez Gene ids must be performed (in case of ambiguities, the most variant probe is kept for each gene), FALSE otherwise. Note that for Affymetrix HGU datasets, the mapping is not necessary.
mapping	Matrix with columns "EntrezGene.ID" and "probe" used to force the mapping such that the probes are not selected based on their variance.
verbose	TRUE to print informative messages, FALSE otherwis.

Value

A list with items:

- score: Continuous signature scores
- risk: Binary risk classification, 1 being high risk and 0 being low risk.
- mapping: Mapping used if necessary.
- probe: If mapping is performed, this matrix contains the correspondence between the gene list (aka signature) and gene expression data.

References

Parker, Joel S. and Mullins, Michael and Cheang, Maggie C.U. and Leung, Samuel and Voduc, David and Vickery, Tammi and Davies, Sherri and Fauron, Christiane and He, Xiaping and Hu, Zhiyuan and Quackenbush, John F. and Stijleman, Inge J. and Palazzo, Juan and Marron, J.S. and Nobel, Andrew B. and Mardis, Elaine and Nielsen, Torsten O. and Ellis, Matthew J. and Perou, Charles M. and Bernard, Philip S. (2009) "Supervised Risk Predictor of Breast Cancer Based on Intrinsic Subtypes", Journal of Clinical Oncology, 27(8):1160-1167

scmgene.robust 55

Examples

```
# load NKI dataset
data(vdxs)
data(pam50)

# compute relapse score
rs.vdxs <- rorS(data=data.vdxs, annot=annot.vdxs, do.mapping=TRUE)</pre>
```

scmgene.robust

Subtype Clustering Model using only ESR1, ERBB2 and AURKA genes for identification of breast cancer molecular subtypes

Description

List of parameters defining the Subtype Clustering Model as published in Wirapati et al 2009 and Desmedt et al 2008 but using single genes instead of gene modules.

Usage

```
data(scmgene.robust)
```

Format

List of parameters for SCMGENE:

- parameters: List of parameters for the mixture of three Gaussians (ER-/HER2-, HER2+ and ER+/HER2-) that define the Subtype Clustering Model. The structure is the same than for an mclust::Mclust object.
- cutoff.AURKA: Cutoff for AURKA module score in order to identify ER+/HER2- High Proliferation (aka Luminal B) tumors and ER+/HER2- Low Proliferation (aka Luminal A) tumors.
- mod: ESR1, ERBB2 and AURKA modules.

Source

http://clincancerres.aacrjournals.org/content/14/16/5158.abstract?ck=nck

References

Desmedt C, Haibe-Kains B, Wirapati P, Buyse M, Larsimont D, Bontempi G, Delorenzi M, Piccart M, and Sotiriou C (2008) "Biological processes associated with breast cancer clinical outcome depend on the molecular subtypes", Clinical Cancer Research, 14(16):5158–5165.

56 scmod2.robust

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Description

List of parameters defining the Subtype Clustering Model as published in Desmedt et al 2008.

Usage

```
data(scmod1.robust)
```

Format

List of parameters for SCMOD1:

- parameters: List of parameters for the mixture of three Gaussians (ER-/HER2-, HER2+ and ER+/HER2-) that define the Subtype Clustering Model. The structure is the same than for an mclust::Mclust() object.
- cutoff.AURKA: Cutoff for AURKA module score in order to identify ER+/HER2- High Proliferation (aka Luminal B) tumors and ER+/HER2- Low Proliferation (aka Luminal A) tumors.
- mod: ESR1, ERBB2 and AURKA modules.

Source

http://clincancerres.aacrjournals.org/content/14/16/5158.abstract?ck=nck

References

Desmedt C, Haibe-Kains B, Wirapati P, Buyse M, Larsimont D, Bontempi G, Delorenzi M, Piccart M, and Sotiriou C (2008) "Biological processes associated with breast cancer clinical outcome depend on the molecular subtypes", *Clinical Cancer Research*, *14*(16):5158–5165.

scmod2.robust	Subtype Clustering Model using ESR1, ERBB2 and AURKA modules for identification of breast cancer molecular subtypes (Desmedt et al 2008)
	2000)

Description

List of parameters defining the Subtype Clustering Model as published in Desmedt et al 2008.

Usage

```
data(scmod1.robust)
```

setcolclass.df 57

Format

List of parameters for SCMOD2:

• parameters: List of parameters for the mixture of three Gaussians (ER-/HER2-, HER2+ and ER+/HER2-) that define the Subtype Clustering Model. The structure is the same than for an mclust::Mclust object.

- cutoff.AURKA: Cutoff for AURKA module score in order to identify ER+/HER2- High Proliferation (aka Luminal B) tumors and ER+/HER2- Low Proliferation (aka Luminal A) tumors.
- mod: ESR1, ERBB2 and AURKA modules.

Source

http://breast-cancer-research.com/content/10/4/R65k

References

Wirapati P, Sotiriou C, Kunkel S, Farmer P, Pradervand S, Haibe-Kains B, Desmedt C, Ignatiadis M, Sengstag T, Schutz F, Goldstein DR, Piccart MJ and Delorenzi M (2008) "Meta-analysis of Gene-Expression Profiles in Breast Cancer: Toward a Unified Understanding of Breast Cancer Sub-typing and Prognosis Signatures", Breast Cancer Research, 10(4):R65.

setcolclass.df

Function to set the class of columns in a data.frame

Description

This function enables to set the class of each column in a data.frame.

Usage

```
setcolclass.df(df, colclass, factor.levels)
```

Arguments

df data.frame for which columns' class need to be updated.

colclass class for each column of the data.frame.

factor.levels list of levels for each factor.

Value

A data.frame with columns' class and levels properly set

Examples

```
tt <- data.frame(matrix(NA, nrow=3, ncol=3, dimnames=list(1:3, paste("column", 1:3, sep="."))),
    stringsAsFactors=FALSE)
tt <- setcolclass.df(df=tt, colclass=c("numeric", "factor", "character"),
    factor.levels=list(NULL, c("F1", "F2", "F3"), NULL))</pre>
```

sig.gene70

sig.endoPredict	Signature used to compute the endoPredict signature as published by Filipits et al 2011

Description

List of 11 genes included in the endoPredict signature. The EntrezGene.ID allows for mapping and the mapping to affy probes is already provided.

Usage

```
data(sig.endoPredict)
```

Format

sig.endoPredict is a matrix with 5 columns containing the annotations and information related to the signature itself (including a mapping to Affymetrix HGU platform).

References

Filipits, M., Rudas, M., Jakesz, R., Dubsky, P., Fitzal, F., Singer, C. F., et al. (2011). "A new molecular predictor of distant recurrence in ER-positive, HER2-negative breast cancer adds independent information to conventional clinical risk factors." *Clinical Cancer Research*, **17**(18):6012–6020.

sig.gene70 Signature used to compute the 70 genes prognosis profile (GENE70 as published by van't Veer et al. 2002
--

Description

List of 70 agilent probe ids representing 56 unique genes included in the GENE70 signature. The EntrezGene.ID allows for mapping and the "average.good.prognosis.profile" values allows for signature computation.

Usage

```
data(sig.gene70)
```

Format

sig.gene70 is a matrix with 9 columns containing the annotations and information related to the signature itself.

Source

http://www.nature.com/nature/journal/v415/n6871/full/415530a.html

sig.gene76

References

L. J. van't Veer and H. Dai and M. J. van de Vijver and Y. D. He and A. A. Hart and M. Mao and H. L. Peterse and K. van der Kooy and M. J. Marton and A. T. Witteveen and G. J. Schreiber and R. M. Kerkhiven and C. Roberts and P. S. Linsley and R. Bernards and S. H. Friend (2002) "Gene Expression Profiling Predicts Clinical Outcome of Breast Cancer", Nature, 415:530–536.

sig.gene76

Signature used to compute the Relapse Score (GENE76) as published in Wang et al. 2005

Description

List of 76 affymetrix hgu133a probesets representing 60 unique genes included in the GENE76 signature. The EntrezGene.ID allows for mapping and the coefficient allows for signature computation.

Usage

data(sig.gene76)

Format

sig.gene70 is a matrix with 10 columns containing the annotations and information related to the signature itself.

Source

http://www.thelancet.com/journals/lancet/article/PIIS0140-6736(05)17947-1/abstract

References

Y. Wang and J. G. Klijn and Y. Zhang and A. M. Sieuwerts and M. P. Look and F. Yang and D. Talantov and M. Timmermans and M. E. Meijer-van Gelder and J. Yu and T. Jatkoe and E. M. Berns and D. Atkins and J. A. Foekens (2005) "Gene-Expression Profiles to Predict Distant Metastasis of Lymph-Node-Negative Primary Breast Cancer", Lancet, 365(9460):671–679.

sig.ggi

sig.genius	Gene Expression progNostic Index Using Subtypes (GENIUS) as pub-
	lished by Haibe-Kains et al. 2010.

Description

List of three gene signatures which compose the Gene Expression progNostic Index Using Subtypes (GENIUS) as published by Haibe-Kains et al. 2009. GENIUSM1, GENIUSM2 and GENIUSM3 are the ER-/HER2-, HER2+ and ER+/HER2- subtype signatures respectively.

Format

sig.genius is a list a three subtype signatures.

References

Haibe-Kains B, Desmedt C, Rothe F, Sotiriou C and Bontempi G (2010) "A fuzzy gene expression-based computational approach improves breast cancer prognostication", Genome Biology, 11(2):R18

2006	sig.ggi	Gene expression Grade Index (GGI) as published in Sotiriou et al. 2006
------	---------	--

Description

List of 128 affymetrix hgu133a probesets representing 97 unique genes included in the GGI signature. The "EntrezGene.ID" column allows for mapping and "grade" defines the up-regulation of the expressions either in histological grade 1 or 3.

Usage

```
data(sig.ggi)
```

Format

sig.ggi is a matrix with 9 columns containing the annotations and information related to the signature itself.

Source

```
http://jnci.oxfordjournals.org/cgi/content/full/98/4/262/DC1
```

References

Sotiriou C, Wirapati P, Loi S, Harris A, Bergh J, Smeds J, Farmer P, Praz V, Haibe-Kains B, Lallemand F, Buyse M, Piccart MJ and Delorenzi M (2006) "Gene expression profiling in breast cancer: Understanding the molecular basis of histologic grade to improve prognosis", Journal of National Cancer Institute, 98:262–272

sig.oncotypedx 61

sig.oncotypedx	Signature used to compute the OncotypeDX signature as published by Paik et al 2004

Description

List of 21 genes included in the OncotypeDX signature. The EntrezGene.ID allows for mapping and the mapping to affy probes is already provided.

Usage

data(sig.oncotypedx)

References

S. Paik, S. Shak, G. Tang, C. Kim, J. Bakker, M. Cronin, F. L. Baehner, M. G. Walker, D. Watson, T. Park, W. Hiller, E. R. Fisher, D. L. Wickerham, J. Bryant, and N. Wolmark (2004) "A Multigene Assay to Predict Recurrence of Tamoxifen-Treated, Node-Negative Breast Cancer", New England Journal of Medicine, 351(27):2817–2826.

sig.pik3cags	Gene expression Grade Index (GGI) as published in Sotiriou et al. 2006
--------------	--

Description

List of 278 affymetrix hgu133a probesets representing 236 unique genes included in the PIK3CA-GS signature. The "EntrezGene.ID" column allows for mapping and "coefficient" refers to to the direction of association with PIK3CA mutation.

Usage

```
data(sig.pik3cags)
```

Format

sig.pik3cags is a matrix with 3 columns containing the annotations and information related to the signature itself.

Source

http://www.pnas.org/content/107/22/10208/suppl/DCSupplemental

62 sig.score

References

Loi S, Haibe-Kains B, Majjaj S, Lallemand F, Durbecq V, Larsimont D, Gonzalez-Angulo AM, Pusztai L, Symmans FW, Bardelli A, Ellis P, Tutt AN, Gillett CE, Hennessy BT., Mills GB, Phillips WA, Piccart MJ, Speed TP, McArthur GA, Sotiriou C (2010) "PIK3CA mutations associated with gene signature of low mTORC1 signaling and better outcomes in estrogen receptor-positive breast cancer", Proceedings of the National Academy of Sciences, 107(22):10208–10213

sig.score	Function to compute signature scores as linear combination of gene expressions

Description

This function computes a signature score from a gene list (aka gene signature), i.e. a signed average as published in Sotiriou et al. 2006 and Haibe-Kains et al. 2009.

Usage

```
sig.score(x, data, annot, do.mapping = FALSE, mapping, size = 0,
  cutoff = NA, signed = TRUE, verbose = FALSE)
```

Arguments

X	Matrix containing the gene(s) in the gene list in rows and at least three columns: "probe", "EntrezGene.ID" and "coefficient" standing for the name of the probe, the NCBI Entrez Gene id and the coefficient giving the direction and the strength of the association of each gene in the gene list.
data	Matrix of gene expressions with samples in rows and probes in columns, dimnames being properly defined.
annot	Matrix of annotations with at least one column named "EntrezGene.ID", dimnames being properly defined.
do.mapping	TRUE if the mapping through Entrez Gene ids must be performed (in case of ambiguities, the most variant probe is kept for each gene), FALSE otherwise.
mapping	Matrix with columns "EntrezGene.ID" and "probe" used to force the mapping such that the probes are not selected based on their variance.
size	Integer specifying the number of probes to be considered in signature computation. The probes will be sorted by absolute value of coefficients.
cutoff	Only the probes with coefficient greater than cutoff will be considered in signature computation.
signed	TRUE if only the sign of the coefficient must be considered in signature computation, FALSE otherwise.
verbose	TRUE to print informative messages, FALSE otherwise.

sig.tamr13 63

Value

A list with items:

- score: Signature score.
- mapping: Mapping used if necessary.
- probe: If mapping is performed, this matrix contains the correspondence between the gene list (aka signature) and gene expression data.

References

Sotiriou C, Wirapati P, Loi S, Harris A, Bergh J, Smeds J, Farmer P, Praz V, Haibe-Kains B, Lallemand F, Buyse M, Piccart MJ and Delorenzi M (2006) "Gene expression profiling in breast cancer: Understanding the molecular basis of histologic grade to improve prognosis", Journal of National Cancer Institute, 98:262-272 Haibe-Kains B (2009) "Identification and Assessment of Gene Signatures in Human Breast Cancer", PhD thesis at Universite Libre de Bruxelles, http://theses.ulb.ac.be/ETD-db/collection/available/ULBetd-02182009-083101/

Examples

```
# load NKI data
data(nkis)
# load GGI signature
data(sig.ggi)
# make of ggi signature a gene list
ggi.gl <- cbind(sig.ggi[ ,c("probe", "EntrezGene.ID")],
    "coefficient"=ifelse(sig.ggi[ ,"grade"] == 1, -1, 1))
# computation of signature scores
ggi.score <- sig.score(x=ggi.gl, data=data.nkis, annot=annot.nkis,
    do.mapping=TRUE, signed=TRUE, verbose=TRUE)
str(ggi.score)</pre>
```

sig.tamr13

Tamoxifen Resistance signature composed of 13 gene clusters (TAMR13) as published by Loi et al. 2008.

Description

List of 13 clusters of genes (and annotations) and their corresponding coefficient as an additional attribute.

Usage

```
data(sig.tamr13)
```

Format

sig.tamr13 is a list a 13 clusters of genes with their corresponding coefficient.

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References

Loi S, Haibe-Kains B, Desmedt C, Wirapati P, Lallemand F, Tutt AM, Gillet C, Ellis P, Ryder K, Reid JF, Daidone MG, Pierotti MA, Berns EMJJ, Jansen MPHM, Foekens JA, Delorenzi M, Bontempi G, Piccart MJ and Sotiriou C (2008) "Predicting prognosis using molecular profiling in estrogen receptor-positive breast cancer treated with tamoxifen", BMC Genomics, 9(1):239

sigOvcAngiogenic

sigOvcAngiogenic dataset

Description

sigOvcAngiogenic dataset

Source

http://jnci.oxfordjournals.org/cgi/content/full/98/4/262/DC1

References

Bentink S, Haibe-Kains B, Risch T, Fan J-B, Hirsch MS, Holton K, Rubio R, April C, Chen J, Wickham-Garcia E, Liu J, Culhane AC, Drapkin R, Quackenbush JF, Matulonis UA (2012) "Angiogenic mRNA and microRNA Gene Expression Signature Predicts a Novel Subtype of Serous Ovarian Cancer", PloS one, 7(2):e30269

sigOvcCrijns

sigOvcCrijns dataset

Description

sigOvcCrijns dataset

Source

http://jnci.oxfordjournals.org/cgi/content/full/98/4/262/DC1

References

Crijns APG, Fehrmann RSN, de Jong S, Gerbens F, Meersma G J, Klip HG, Hollema H, Hofstra RMW, te Meerman GJ, de Vries EGE, van der Zee AGJ (2009) "Survival-Related Profile, Pathways, and Transcription Factors in Ovarian Cancer" PLoS Medicine, 6(2):e1000024.

sigOvcSpentzos 65

sigOvcSpentzos

sigOcvSpentzos dataset

Description

sigOcvSpentzos dataset

Source

http://jnci.oxfordjournals.org/cgi/content/full/98/4/262/DC1

References

Spentzos, D., Levine, D. A., Ramoni, M. F., Joseph, M., Gu, X., Boyd, J., et al. (2004). "Gene expression signature with independent prognostic significance in epithelial ovarian cancer". Journal of clinical oncology, 22(23), 4700–4710. doi:10.1200/JCO.2004.04.070

sigOvcTCGA

sigOvcTCGA dataset

Description

sigOvcTCGA dataset

Source

http://jnci.oxfordjournals.org/cgi/content/full/98/4/262/DC1

References

Bell D, Berchuck A, Birrer M et al. (2011) "Integrated genomic analyses of ovarian carcinoma", Nature, 474(7353):609–615

66 spearmanCI

sig0vcYoshihara

sigOvcYoshihara dataset

Description

sigOvcYoshihara dataset

Source

```
http://jnci.oxfordjournals.org/cgi/content/full/98/4/262/DC1
```

References

Yoshihara K, Tajima A, Yahata T, Kodama S, Fujiwara H, Suzuki M, Onishi Y, Hatae M, Sueyoshi K, Fujiwara H, Kudo, Yoshiki, Kotera K, Masuzaki H, Tashiro H, Katabuchi H, Inoue I, Tanaka K (2010) "Gene expression profile for predicting survival in advanced-stage serous ovarian cancer across two independent datasets", PloS one, 5(3):e9615.

spearmanCI

Function to compute the confidence interval for the Spearman correelation coefficient

Description

This function enables to compute the confidence interval for the Spearman correlation coefficient using the Fischer Z transformation.

Usage

```
spearmanCI(x, n, alpha = 0.05)
```

Arguments

x Spearman correlation coefficient rho.

n the sample size used to compute the Spearman rho.

alpha alpha level for confidence interval.

Value

A vector containing the lower, upper values for the confidence interval and p-value for Spearman rho

Examples

```
spearmanCI(x=0.2, n=100, alpha=0.05)
```

ssp2003 67

ssp2003	SSP2003 classifier for identification of breast cancer molecular subtypes (Sorlie et al 2003)

Description

List of parameters defining the SSP2003 classifier for identification of breast cancer molecular subtypes (Sorlie et al 2003).

Usage

```
data(ssp2003)
data(ssp2003.robust)
data(ssp2003.scale)
```

Format

List of parameters for SSP2003:

- centroids: Gene expression centroids for each subtype.
- centroids.map: Mapping for centroids.
- method.cor: Method of correlation used to compute distance to the centroids.
- method.centroids: Method used to compute the centroids.
- std: Method used to compute the centroids.
- mins: Minimum number of samples within each cluster allowed during the fitting of the model.

Source

```
http://www.pnas.org/content/100/14/8418
```

References

T. Sorlie and R. Tibshirani and J. Parker and T. Hastie and J. S. Marron and A. Nobel and S. Deng and H. Johnsen and R. Pesich and S. Geister and J. Demeter and C. Perou and P. E. Lonning and P. O. Brown and A. L. Borresen-Dale and D. Botstein (2003) "Repeated Observation of Breast Tumor Subtypes in Independent Gene Expression Data Sets", Proceedings of the National Academy of Sciences, 1(14):8418–8423

68 ssp2006

ssp2006	SSP2006 classifier for identification of breast cancer molecular sub-
	types (Hu et al 2006)

Description

List of parameters defining the SSP2006 classifier for identification of breast cancer molecular subtypes (Hu et al 2006).

Usage

```
data(ssp2006)
data(ssp2006.robust)
data(ssp2006.scale)
```

Format

List of parameters for SSP2006:

- centroids: Gene expression centroids for each subtype.
- centroids.map: Mapping for centroids.
- method.cor: Method of correlation used to compute distance to the centroids.
- method.centroids: Method used to compute the centroids.
- std: Method of standardization for gene expressions.
- mins: Minimum number of samples within each cluster allowed during the fitting of the model.

Details

Three versions of the model are provided, each of ones differs by the gene expressions standardization method since it has an important impact on the subtype classification:

- ssp2006: Use of the official centroids without scaling of the gene expressions.
- ssp2006.scale: Use of the official centroids with traditional scaling of the gene expressions (see base::scale())
- ssp2006.robust: Use of the official centroids with robust scaling of the gene expressions (see rescale()) The model ssp2006.robust has been shown to reach the best concordance with the traditional clinical parameters (ER IHC, HER2 IHC/FISH and histological grade). However the use of this model is recommended only when the dataset is representative of a global population of breast cancer patients (no sampling bias, the 5 subtypes should be present).

Source

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References

Hu, Zhiyuan and Fan, Cheng and Oh, Daniel and Marron, JS and He, Xiaping and Qaqish, Bahjat and Livasy, Chad and Carey, Lisa and Reynolds, Evangeline and Dressler, Lynn and Nobel, Andrew and Parker, Joel and Ewend, Matthew and Sawyer, Lynda and Wu, Junyuan and Liu, Yudong and Nanda, Rita and Tretiakova, Maria and Orrico, Alejandra and Dreher, Donna and Palazzo, Juan and Perreard, Laurent and Nelson, Edward and Mone, Mary and Hansen, Heidi and Mullins, Michael and Quackenbush, John and Ellis, Matthew and Olopade, Olufunmilayo and Bernard, Philip and Perou, Charles (2006) "The molecular portraits of breast tumors are conserved across microarray platforms", *BMC Genomics*, 7(96)

st.gallen Function to compute the cation	St Gallen consensus criterion for prognosti-
	si Gatten consensus criterion for prognosti-

Description

This function computes the updated St Gallen consensus criterions as published by Goldhirsh et al 2003.

Usage

```
st.gallen(size, grade, node, her2.neu, age, vascular.inv, na.rm = FALSE)
```

Arguments

size	tumor size in cm.
grade	Histological grade, i.e. low (1), intermediate (2) and high (3) grade.
node	Nodal status (0 or 1 for no lymph node invasion a,d at least 1 invaded lymph ode respectively).
her2.neu	Her2/neu status (0 or 1).
age	Age at diagnosis (in years).
vascular.inv	Peritumoral vascular invasion (0 or 1).
na.rm	TRUE if missing values should be removed, FALSE otherwise.

Value

Vector of risk predictions: "Good", "Intermediate", and "Poor".

References

Goldhirsh A, Wood WC, Gelber RD, Coates AS, Thurlimann B, and Senn HJ (2003) "Meeting highlights: Updated international expert consensus on the primary therapy of early breast cancer", Journal of Clinical Oncology, 21(17):3357-3365.

70 stab.fs

See Also

npi

Examples

```
# load nkis dataset
data(nkis)

# compute St Gallen predictions
st.gallen(size=demo.nkis[ ,"size"], grade=demo.nkis[ ,"grade"],
node=demo.nkis[ ,"node"], her2.neu=sample(x=0:1, size=nrow(demo.nkis),
replace=TRUE), age=demo.nkis[ ,"age"], vascular.inv=sample(x=0:1,
size=nrow(demo.nkis), replace=TRUE), na.rm=TRUE)
```

stab.fs

Function to quantify stability of feature selection

Description

This function computes several indexes to quantify feature selection stability. This is usually estimated through perturbation of the original dataset by generating multiple sets of selected features.

Usage

```
stab.fs(fsets, N, method = c("kuncheva", "davis"), ...)
```

Arguments

fsets list of sets of selected features, each set of selected features may have different size.

N total number of features on which feature selection is performed.

method stability index (see details section).

... additional parameters passed to stability index (penalty that is a numeric for

Davis' stability index, see details section).

Details

Stability indices may use different parameters. In this version only the Davis index requires an additional parameter that is penalty, a numeric value used as penalty term. Kuncheva index (kuncheva) lays in [-1, 1], An index of -1 means no intersection between sets of selected features, +1 means that all the same features are always selected and 0 is the expected stability of a random feature selection. Davis index (davis) lays in [0,1], With a penalty term equal to 0, an index of 0 means no intersection between sets of selected features and +1 means that all the same features are always selected. A penalty of 1 is usually used so that a feature selection performed with no or all features has a Davis stability index equals to 0. None estimate of the expected Davis stability index of a random feature selection was published.

stab.fs.ranking 71

Value

A numeric that is the stability index.

References

Davis CA, Gerick F, Hintermair V, Friedel CC, Fundel K, Kuffner R, Zimmer R (2006) "Reliable gene signatures for microarray classification: assessment of stability and performance", Bioinformatics, 22(19):356-2363. Kuncheva LI (2007) "A stability index for feature selection", AIAP'07: Proceedings of the 25th conference on Proceedings of the 25th IASTED International Multi-Conference, pages 390-395.

See Also

stab.fs.ranking

Examples

```
set.seed(54321)
# 100 random selection of 50 features from a set of 10,000 features
fsets <- lapply(as.list(1:100), function(x, size=50, N=10000) {
    return(sample(1:N, size, replace=FALSE))} )
names(fsets) <- paste("fsel", 1:length(fsets), sep=".")

# Kuncheva index
stab.fs(fsets=fsets, N=10000, method="kuncheva")
# close to 0 as expected for a random feature selection

# Davis index
stab.fs(fsets=fsets, N=10000, method="davis", penalty=1)</pre>
```

stab.fs.ranking

Function to quantify stability of feature ranking

Description

This function computes several indexes to quantify feature ranking stability for several number of selected features. This is usually estimated through perturbation of the original dataset by generating multiple sets of selected features.

Usage

```
stab.fs.ranking(fsets, sizes, N, method = c("kuncheva", "davis"), ...)
```

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Arguments

fsets	list or matrix of sets of selected features (in rows), each ranking must have the same size.
sizes	Number of top-ranked features for which the stability index must be computed.
N	total number of features on which feature selection is performed
method	stability index (see details section).
•••	additional parameters passed to stability index (penalty that is a numeric for Davis' stability index, see details section).

Details

Stability indices may use different parameters. In this version only the Davis index requires an additional parameter that is penalty, a numeric value used as penalty term. Kuncheva index (kuncheva) lays in [-1, 1], An index of -1 means no intersection between sets of selected features, +1 means that all the same features are always selected and 0 is the expected stability of a random feature selection. Davis index (davis) lays in [0,1], With a penalty term equal to 0, an index of 0 means no intersection between sets of selected features and +1 means that all the same features are always selected. A penalty of 1 is usually used so that a feature selection performed with no or all features has a Davis stability index equals to 0. None estimate of the expected Davis stability index of a random feature selection was published.

Value

A vector of numeric that are stability indices for each size of the sets of selected features given the rankings.

References

Davis CA, Gerick F, Hintermair V, Friedel CC, Fundel K, Kuffner R, Zimmer R (2006) "Reliable gene signatures for microarray classification: assessment of stability and performance", Bioinformatics, 22(19):356-2363. Kuncheva LI (2007) "A stability index for feature selection", AIAP'07: Proceedings of the 25th conference on Proceedings of the 25th IASTED International Multi-Conference, pages 390-395.

See Also

stab.fs

Examples

```
# 100 random selection of 50 features from a set of 10,000 features
fsets <- lapply(as.list(1:100), function(x, size=50, N=10000) {
    return(sample(1:N, size, replace=FALSE))} )
names(fsets) <- paste("fsel", 1:length(fsets), sep=".")

# Kuncheva index
stab.fs.ranking(fsets=fsets, sizes=c(1, 10, 20, 30, 40, 50),
    N=10000, method="kuncheva")
# close to 0 as expected for a random feature selection</pre>
```

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```
# Davis index
stab.fs.ranking(fsets=fsets, sizes=c(1, 10, 20, 30, 40, 50),
    N=10000, method="davis", penalty=1)
```

strescR

Utility function to escape LaTeX special characters present in a string

Description

This function returns a vector of strings in which LaTeX special characters are escaped, this was useful in conjunction with xtable.

Usage

```
strescR(strings)
```

Arguments

strings

A vector of strings to deal with.

Value

A vector of strings with escaped characters within each string.

References

```
citation("seqinr")
```

See Also

stresc

```
strescR("MISC_RNA")
strescR(c("BB_0001","BB_0002"))
```

74 subtype.cluster

Description

This function fits the Subtype Clustering Model as published in Desmedt et al. 2008 and Wiarapati et al. 2008. This model is actually a mixture of three Gaussians with equal shape, volume and variance (see EEI model in Mclust). This model is adapted to breast cancer and uses ESR1, ERBB2 and AURKA dimensions to identify the molecular subtypes, i.e. ER-/HER2-, HER2+ and ER+/HER2-(Low and High Prolif).

Usage

```
subtype.cluster(module.ESR1, module.ERBB2, module.AURKA, data, annot,
  do.mapping = FALSE, mapping, do.scale = TRUE, rescale.q = 0.05,
  model.name = "EEI", do.BIC = FALSE, plot = FALSE, filen, verbose = FALSE)
```

Arguments

verbose

gaments				
module.ESR1	Matrix containing the ESR1-related gene(s) in rows and at least three columns: "probe", "EntrezGene.ID" and "coefficient" standing for the name of the probe, the NCBI Entrez Gene id and the coefficient giving the direction and the strength of the association of each gene in the gene list.			
module.ERBB2	Idem for ERBB2.			
module.AURKA	Idem for AURKA.			
data	Matrix of gene expressions with samples in rows and probes in columns, dimnames being properly defined.			
annot	Matrix of annotations with at least one column named "EntrezGene.ID", dimnames being properly defined.			
do.mapping	TRUE if the mapping through Entrez Gene ids must be performed (in case of ambiguities, the most variant probe is kept for each gene), FALSE otherwise.			
mapping	DEPRECATED Matrix with columns "EntrezGene.ID" and "probe" used to force the mapping such that the probes are not selected based on their variance.			
do.scale	TRUE if the ESR1, ERBB2 and AURKA (module) scores must be rescaled (see rescale), FALSE otherwise.			
rescale.q	Proportion of expected outliers for rescaling the gene expressions.			
model.name	Name of the model used to fit the mixture of Gaussians with the Mclust from the mclust package; default is "EEI" for fitting a mixture of Gaussians with diagonal variance, equal volume, equal shape and identical orientation.			
do.BIC	TRUE if the Bayesian Information Criterion must be computed for number of clusters ranging from 1 to 10, FALSE otherwise.			
plot	TRUE if the patients and their corresponding subtypes must be plotted, FALSE otherwise.			
filen	Name of the csv file where the subtype clustering model must be stored.			

TRUE to print informative messages, FALSE otherwise.

subtype.cluster 75

Value

A list with items:

model: Subtype Clustering Model (mixture of three Gaussians), like scmgene.robust, scmod1.robust and scmod2.robust when this function is used on expO dataset (International Genomics Consortium) with the gene modules published in the two references cited below.

- BIC: Bayesian Information Criterion for the Subtype Clustering Model with number of clusters ranging from 1 to 10.
- subtype: Subtypes identified by the Subtype Clustering Model. Subtypes can be either "ER-/HER2-", "HER2+" or "ER+/HER2-".
- subtype.proba: Probabilities to belong to each subtype estimated by the Subtype Clustering Model.
- subtype2: Subtypes identified by the Subtype Clustering Model using AURKA to discriminate low and high proliferative tumors. Subtypes can be either "ER-/HER2-", "HER2+", "ER+/HER2- High Prolif" or "ER+/HER2- Low Prolif".
- subtype.proba2: Probabilities to belong to each subtype (including discrimination between lowly and highly proliferative ER+/HER2- tumors, see subtype2) estimated by the Subtype Clustering Model.
- module.scores: Matrix containing ESR1, ERBB2 and AURKA module scores.

References

Desmedt C, Haibe-Kains B, Wirapati P, Buyse M, Larsimont D, Bontempi G, Delorenzi M, Piccart M, and Sotiriou C (2008) "Biological processes associated with breast cancer clinical outcome depend on the molecular subtypes", Clinical Cancer Research, 14(16):5158-5165. Wirapati P, Sotiriou C, Kunkel S, Farmer P, Pradervand S, Haibe-Kains B, Desmedt C, Ignatiadis M, Sengstag T, Schutz F, Goldstein DR, Piccart MJ and Delorenzi M (2008) "Meta-analysis of Gene-Expression Profiles in Breast Cancer: Toward a Unified Understanding of Breast Cancer Sub-typing and Prognosis Signatures", Breast Cancer Research, 10(4):R65.

See Also

subtype.cluster.predict, intrinsic.cluster, intrinsic.cluster.predict, scmod1.robust, scmod2.robust

```
# example without gene mapping
# load exp0 data
data(expos)
# load gene modules
data(mod1)
# fit a Subtype Clustering Model
scmod1.expos <- subtype.cluster(module.ESR1=mod1$ESR1, module.ERBB2=mod1$ERBB2,
    module.AURKA=mod1$AURKA, data=data.expos, annot=annot.expos, do.mapping=FALSE,
    do.scale=TRUE, plot=TRUE, verbose=TRUE)
str(scmod1.expos, max.level=1)
table(scmod1.expos$subtype2)</pre>
```

```
# example with gene mapping
# load NKI data
data(nkis)
# load gene modules
data(mod1)
# fit a Subtype Clustering Model
scmod1.nkis <- subtype.cluster(module.ESR1=mod1$ESR1, module.ERBB2=mod1$ERBB2,
    module.AURKA=mod1$AURKA, data=data.nkis, annot=annot.nkis, do.mapping=TRUE,
    do.scale=TRUE, plot=TRUE, verbose=TRUE)
str(scmod1.nkis, max.level=1)
table(scmod1.nkis$subtype2)</pre>
```

subtype.cluster.predict

Function to identify breast cancer molecular subtypes using the Subtype Clustering Model

Description

This function identifies the breast cancer molecular subtypes using a Subtype Clustering Model fitted by subtype.cluster.

Usage

```
subtype.cluster.predict(sbt.model, data, annot, do.mapping = FALSE,
  mapping, do.prediction.strength = FALSE,
  do.BIC = FALSE, plot = FALSE, verbose = FALSE)
```

Arguments

	sbt.model	Subtype Clustering Model as returned by subtype.cluster.	
	data	Matrix of gene expressions with samples in rows and probes in columns, dimnames being properly defined.	
	annot	Matrix of annotations with at least one column named "EntrezGene.ID", dimnames being properly defined.	
	do.mapping	TRUE if the mapping through Entrez Gene ids must be performed (in case of ambiguities, the most variant probe is kept for each gene), FALSE otherwise.	
	mapping	DEPRECATED Matrix with columns "EntrezGene.ID" and "probe" used to force the mapping such that the probes are not selected based on their variance.	
do.prediction.strength			
		TRUE if the prediction strength must be computed (Tibshirani and Walther 2005), FALSE otherwise.	
	do.BIC	TRUE if the Bayesian Information Criterion must be computed for number of clusters ranging from 1 to 10, FALSE otherwise.	
	plot	TRUE if the patients and their corresponding subtypes must be plotted, FALSE otherwise.	
	verbose	TRUE to print informative messages, FALSE otherwise.	

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Value

A list with items:

• subtype: Subtypes identified by the Subtype Clustering Model. Subtypes can be either "ER-/HER2-", "HER2+" or "ER+/HER2-".

- subtype.proba: Probabilities to belong to each subtype estimated by the Subtype Clustering Model.
- prediction.strength: Prediction strength for subtypes.
- BIC: Bayesian Information Criterion for the Subtype Clustering Model with number of clusters ranging from 1 to 10.
- subtype2: Subtypes identified by the Subtype Clustering Model using AURKA to discriminate low and high proliferative tumors. Subtypes can be either "ER-/HER2-", "HER2+", "ER+/HER2- High Prolif" or "ER+/HER2- Low Prolif".
- subtype.proba2: Probabilities to belong to each subtype (including discrimination between lowly and highly proliferative ER+/HER2- tumors, see subtype2) estimated by the Subtype Clustering Model.
- prediction.strength2: Prediction strength for subtypes2.
- module.scores: Matrix containing ESR1, ERBB2 and AURKA module scores.
- mapping: Mapping if necessary (list of matrices with 3 columns: probe, EntrezGene.ID and new.probe).

References

Desmedt C, Haibe-Kains B, Wirapati P, Buyse M, Larsimont D, Bontempi G, Delorenzi M, Piccart M, and Sotiriou C (2008) "Biological processes associated with breast cancer clinical outcome depend on the molecular subtypes", Clinical Cancer Research, 14(16):5158-5165. Wirapati P, Sotiriou C, Kunkel S, Farmer P, Pradervand S, Haibe-Kains B, Desmedt C, Ignatiadis M, Sengstag T, Schutz F, Goldstein DR, Piccart MJ and Delorenzi M (2008) "Meta-analysis of Gene-Expression Profiles in Breast Cancer: Toward a Unified Understanding of Breast Cancer Sub-typing and Prognosis Signatures", Breast Cancer Research, 10(4):R65. Tibshirani R and Walther G (2005) "Cluster Validation by Prediction Strength", Journal of Computational and Graphical Statistics, 14(3):511-528

See Also

subtype.cluster, scmod1.robust, scmod2.robust

```
# without mapping (affy hgu133a or plus2 only)
# load VDX data
data(vdxs)
data(scmgene.robust)

# Subtype Clustering Model fitted on EXPO and applied on VDX
sbt.vdxs <- subtype.cluster.predict(sbt.model=scmgene.robust, data=data.vdxs,
    annot=annot.vdxs, do.mapping=FALSE, do.prediction.strength=FALSE,
    do.BIC=FALSE, plot=TRUE, verbose=TRUE)</pre>
```

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```
table(sbt.vdxs$subtype)
table(sbt.vdxs$subtype2)

# with mapping
# load NKI data
data(nkis)
# Subtype Clustering Model fitted on EXPO and applied on NKI
sbt.nkis <- subtype.cluster.predict(sbt.model=scmgene.robust, data=data.nkis,
    annot=annot.nkis, do.mapping=TRUE, do.prediction.strength=FALSE,
    do.BIC=FALSE, plot=TRUE, verbose=TRUE)
table(sbt.nkis$subtype)
table(sbt.nkis$subtype2)</pre>
```

tamr13

Function to compute the risk scores of the tamoxifen resistance signature (TAMR13)

Description

This function computes signature scores from gene expression values following the algorithm used for the Tamoxifen Resistance signature (TAMR13).

Usage

```
tamr13(data, annot, do.mapping = FALSE, mapping, verbose = FALSE)
```

Arguments

data	Matrix of gene expressions with samples in rows and probes in columns, dimnames being properly defined.
annot	Matrix of annotations with at least one column named "EntrezGene.ID", dimnames being properly defined.
do.mapping	TRUE if the mapping through Entrez Gene ids must be performed (in case of ambiguities, the most variant probe is kept for each gene), FALSE otherwise.
mapping	Matrix with columns "EntrezGene.ID" and "probe" used to force the mapping such that the probes are not selected based on their variance.
verbose	TRUE to print informative messages, FALSE otherwise.

Value

A list with items:

- score: Continuous signature scores.
- risk: Binary risk classification, 1 being high risk and 0 being low risk (not implemented, the function will return NA values).

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References

Loi S, Haibe-Kains B, Desmedt C, Wirapati P, Lallemand F, Tutt AM, Gillet C, Ellis P, Ryder K, Reid JF, Daidone MG, Pierotti MA, Berns EMJJ, Jansen MPHM, Foekens JA, Delorenzi M, Bontempi G, Piccart MJ and Sotiriou C (2008) "Predicting prognosis using molecular profiling in estrogen receptor- positive breast cancer treated with tamoxifen", BMC Genomics, 9(1):239

See Also

```
gene76
```

Examples

```
# load TAMR13 signature
data(sig.tamr13)
# load VDX dataset
data(vdxs)
# compute relapse score
tamr13.vdxs <- tamr13(data=data.vdxs, annot=annot.vdxs, do.mapping=FALSE)
summary(tamr13.vdxs$score)</pre>
```

tbrm

Function to compute Tukey's Biweight Robust Mean

Description

Computation of Tukey's Biweight Robust Mean, a robust average that is unaffected by outliers.

Usage

```
tbrm(x, C = 9)
```

Arguments

x a numeric vector

a constant. C is preassigned a value of 9 according to the Cook reference below but other values are possible.

Details

This is a one step computation that follows the Affy whitepaper below see page 22. This function is called by chron to calculate a robust mean. C determines the point at which outliers are given a weight of 0 and therefore do not contribute to the calculation of the mean. C=9 sets values roughly +/-6 standard deviations to 0. C=6 is also used in tree-ring chronology development. Cook and Kairiukstis (1990) have further details. Retrieved from tbrm.

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Value

A numeric mean.

References

Statistical Algorithms Description Document, 2002, Affymetrix. p22. Cook, E. R. and Kairiukstis, L.A. (1990) Methods of Dendrochronology: Applications in the Environmental Sciences. ISBN-13: 978-0792305866. Mosteller, F. and Tukey, J. W. (1977) Data Analysis and Regression: a second course in statistics. Addison-Wesley. ISBN-13: 978-0201048544.

See Also

chron

Examples

tbrm(rnorm(100))

vdxs

Gene expression, annotations and clinical data from Wang et al. 2005 and Minn et al 2007

Description

This dataset contains (part of) the gene expression, annotations and clinical data as published in Wang et al. 2005 and Minn et al 2007.

Format

vdxs is a dataset containing three matrices:

- data.vdxs: Matrix containing gene expressions as measured by Affymetrix hgu133a technology (single-channel, oligonucleotides)
- annot.vdxs: Matrix containing annotations of ffymetrix hgu133a microarray platform
- · demo.vdxs: Clinical information of the breast cancer patients whose tumors were hybridized

Details

This dataset represent only partially the one published by Wang et al. 2005 and Minn et al 2007. Indeed only part of the patients (150) and gene expressions (966) are contained in data.vdxs.

Source

```
http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE2034
http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE5327
```

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References

Y. Wang and J. G. Klijn and Y. Zhang and A. M. Sieuwerts and M. P. Look and F. Yang and D. Talantov and M. Timmermans and M. E. Meijer-van Gelder and J. Yu and T. Jatkoe and E. M. Berns and 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

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

weighted.meanvar

Function to compute the weighted mean and weighted variance of 'x'

Description

This function allows for computing the weighted mean and weighted variance of a vector of continuous values.

Usage

```
weighted.meanvar(x, w, na.rm = FALSE)
```

Arguments

x an object containing the values whose weighted mean is to be computed.

w a numerical vector of weights of the same length as x giving the weights to use

for elements of x.

na.rm TRUE if missing values should be removed, FALSE otherwise.

Details

If w is missing then all elements of x are given the same weight, otherwise the weights coerced to numeric by as.numeric. On the contrary of weighted.mean the weights are NOT normalized to sum to one. If the sum of the weights is zero or infinite, NAs will be returned.

Value

A numeric vector of two values that are the weighted mean and weighted variance respectively.

References

http://en.wikipedia.org/wiki/Weighted_variance#Weighted_sample_variance

See Also

stats::weighted.mean

82 write.m.file

Examples

```
set.seed(54321)
weighted.meanvar(x=rnorm(100) + 10, w=runif(100))
```

write.m.file Function to write a 'csv' file containing gene lists (aka gene signatures)

Description

This function allows for writing a 'csv' file containing gene signatures. Each gene signature is composed of at least four columns: "gene.list" is the name of the signature on the first line and empty fields below, "probes" are the probe names, "EntrezGene.ID" are the EntrezGene IDs and "coefficient" are the coefficients of each probe.

Usage

```
write.m.file(obj, file, ...)
```

Arguments

obj List of gene signatures.
file Filename of the 'csv' file.

... Additional parameters for read.csv function.

Value

None.

```
# load gene modules published by Demsedt et al 2009
data(mod1)
# write these gene modules in a 'csv' file
# Not run: write.m.file(obj=mod1, file="desmedt2009_genemodules.csv")
```

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