Dudoit-class

Description

An implementation of the method of Dudoit and colleagues to apply te Wetfall-Young adjustment to p-values to control the family-wise error rate when analysing microarray data

Usage

```
Dudoit(data, classes, nPerm = 1000, verbose = TRUE)
## S4 method for signature 'Dudoit, missing':
plot(x, ylab='T statistics', ...)
## S4 method for signature 'Dudoit':
cutoffSignificant(object, alpha, ...)
## S4 method for signature 'Dudoit':
selectSignificant(object, alpha, ...)
## S4 method for signature 'Dudoit':
countSignificant(object, alpha, ...)
```

Arguments

data	Either a data frame or matrix with numeric values or an exprSet as defined in the BioConductor tools for analyzing microarray data.
classes	If data is a data frame or matrix, then classes must be either a logical vector or a factor. If data is an exprSet, then classes can be a character string that names one of the factor columns in the associated phenoData subobject.
nPerm	An integer; the number of permutations to perform
verbose	A logical flag
object	A Dudoit object
alpha	A real number; the target family-wise error rate
x	A Dudoit object
ylab	Label for the y axis
	The usual extra arguments for generic or plotting routines.

Details

In 2002, Dudoit and colleagues introduced a method to adjust the p-values when performing gene-by-gene tests for differential expression. The adjustment was based on the method of Westfall and Young, with the goal of controlling the family-wise error rate.

Value

The standard method for plot returns what you would expect.

The cutoffSignificant method returns a real number (its input value alpha). The selectSignficant method returns a vector of logical values identifying the significant test results, and countSignificant returns an integer counting the number of significant test results.

Objects from the Class

As usual, objects can be created by **new**, but better methods are available in the form of the **Dudoit** function. The basic inputs to this function are the same as those used for row-by-row statistical tests throughout the ClassComparison package; a detailed description can be found in the MultiTtest class.

The additional input determines the number, **nPerm**, of permutations to perform. The accuracy of the p-value adjustment depends on this value. Since the implementation is in R (and does not call out to something compiled like C or FORTRAN), however, the computations are slow. The default value of 1000 can take a long time with modern microarrays that contain 40,000 spots.

Slots

adjusted.p: The numeric vector of adjusted p-values.

t.statistics: Object of class numeric containing the computed t-statistics.

p.values: Object of class numeric containing the computed p-values.

groups: Object of class character containing the names of the classes being compared.

call: Object of class call containing the function call that created the object.

Extends

Class MultiTtest, directly. In particular, objects of this class inherit methods for summary, hist, and plot from the base class.

Methods

cutoffSignificant(object, alpha, ...) Determine cutoffs on the adjusted p-values at the desired significance level. In other words, this function simplyt returns alpha.

selectSignificant(object, alpha, ...) Compute a logical vector for selecting significant test results.

countSignificant(object, alpha, ...) Count the number of significant test results.

plot signature(x = Dudoit, y = missing): ...

Author(s)

Kevin R. Coombes <kcoombes@mdanderson.org>

References

Dudoit, S., Y.H. Yang, M.J. Callow, and T.P. Speed. 2002. Statistical Methods for Identifying Differentially Expressed Genes in Replicated cDNA Microarray Experiments, Statistica Sinica, 12(1): 111-139.

Westfall, P.H., Young, S.S. Resampling-based multiple testing: examples and methods for p-value adjustment. Wiley series in probability and mathematics statistics. John Wiley and Sons, 1993.

See Also

MultiTtest, Bum, SmoothTtest

Examples

```
ng <- 10000
ns <- 15
nd <- 200
fake.class <- factor(rep(c('A', 'B'), each=ns))
fake.data <- matrix(rnorm(ng*ns*2), nrow=ng, ncol=2*ns)
fake.data[1:nd, 1:ns] <- fake.data[1:nd, 1:ns] + 2
fake.data[(nd+1):(2*nd), 1:ns] <- fake.data[(nd+1):(2*nd), 1:ns] - 2
# the permutation test is slow. it really needs more than
# 100 permutations, but this is just an example...
dud <- Dudoit(fake.data, fake.class, nPerm=100)
summary(dud)
plot(dud)
countSignificant(dud, 0.05)
rm(ng, ns, nd, fake.class, fake.data, dud)
```

Sam-class

The Sam Class

Description

Implements the "Significance Analysis of Microarrays" approach to detecting differentially expressed genes.

Usage

```
Sam(data, classes, nPerm = 100, verbose = TRUE)
## S4 method for signature 'Sam, missing':
plot(x, tracks=NULL, xlab='Expected T Statistics (Empirical)',
ylab='Observed t Statistics', ...)
## S4 method for signature 'Sam':
summary(object, cutoff=1, ...)
## S4 method for signature 'Sam':
```

```
selectSignificant(object, cutoff=1, ...)
## S4 method for signature 'Sam':
countSignificant(object, cutoff=1, ...)
```

Arguments

data	Either a data frame or matrix with numeric values or an exprSet as defined in the BioConductor tools for analyzing microarray data.
classes	If data is a data frame or matrix, then classes must be either a logical vector or a factor. If data is an exprSet, then classes can be a character string that names one of the factor columns in the associated phenoData subobject.
nPerm	An integer; the number of permutations
verbose	A logical flag
x	A Sam object
tracks	a numeric vector
xlab	Label for the x axis
ylab	Label for the y axis
object	A Sam object
cutoff	A numeric value
	The usual extra arguments to generic functions

Details

The SAM approach to analyzing microarray data wwas developed by Tusher and colleagues; their implementation is widely available. This is an independent implementation based on trhe description in their original paper, customized to use the same interface (and thus work with exprSet objects) used by the rest of the ClassComparison package. The fundamental idea behind SAM is that the observed distribution of row-by-row two-sample t-tests should be compared not to the theoretical null distribution but to a null distribution estimated by a permutation test. The Sam constructor performs the permutation test.

Value

summary returns a SamSummary object.
selectSignificant returns a vector of logical values.

 $\verb"countSignificant" returns an integer".$

Creating Objects

As usual, objects can be created by **new**, but better methods are available in the form of the **Sam** function. The inputs to this function are the same as those used for row-by-row statistical tests throughout the ClassComparison package; a detailed description can be found in the **MultiTtest** class.

Slots

t.statistics: A numeric vector containing the observed t-statistics.

observed: A numeric vector containing the sorted observed t-statistics.

- expected: A numeric vector of the expected distribution of t-statistics based on a permutation test.
- sim.data: A matrix containing all the t-statistics from all the permutations.
- call: The function call that created the object.

Methods

summary(object, cutoff=1, ...) Compute a summary of the object.

plot(x, tracks=NULL, xlab='Expected T Statistics (Empirical)', ylab='Observed t Statistics', ...)
Plot the observed and expected t-statistics. The tracks argument causes paralle lines
to be drawn on either side of the quantile-quantile central line, at the specified offsets.
Colors in the plot are controlled by the current values of COLOR.CENTRAL.LINE and
COLOR.CONFIDENCE.CURVE

selectSignificant(object, cutoff=1, ...) compute a vector that selects significant values
countSignificant(object, cutoff=1, ...) count the number of significant values

Author(s)

Kevin R. Coombes <kcoombes@mdanderson.org>

References

Tusher, V.G., Tibshirani, R., and Chu, G. 2001. Significance analysis of microarrays applied to the ionizing radiation response. Proc Natl Acad Sci U S A 98, 5116-5121.

See Also

MultiTtest, Bum

Examples

```
ng <- 10000
ns <- 50
nd <- 100
dat <- matrix(rnorm(ng*ns), ncol=ns)
dat[1:nd, 1:(ns/2)] <- dat[1:nd, 1:(ns/2)] + 2
dat[(nd+1):(2*nd), 1:(ns/2)] <- dat[(nd+1):(2*nd), 1:(ns/2)] - 2
cla <- factor(rep(c('A', 'B'), each=25))
res <- Sam(dat, cla)
plot(res)
plot(res, tracks=1:3)
summary(res)
```

```
summary(res, cutoff=2)
a <- summary(res)
plot(a@significant.calls)
plot(a@significant.calls[1:300])
countSignificant(res, 1)
rm(ng, ns, nd, dat, cla, res, a)</pre>
```

SingleGroup-class The SingleGroup Class

Description

Preliminary analysis of one group of samples for use in the SmoothTtest class. A key feature is the standard quality control plot.

Usage

```
SingleGroup(avg, sd, span = 0.5, name = "')
## S4 method for signature 'SingleGroup':
as.data.frame(x, row.names=NULL, optional=FALSE)
## S4 method for signature 'SingleGroup':
summary(object, ...)
## S4 method for signature 'SingleGroup':
print(x, ...)
## S4 method for signature 'SingleGroup, missing':
plot(x, multiple=3, ccl=0, main=x@name,
    xlab='Mean', ylab='Std Dev', xlim=0, ylim=0, ...)
```

Arguments

avg	A numeric vector of mean values
sd	A numeric vector of standard deviations
span	The span parameter is passed onto loess.
name	A character string; the name of this object
object	A SingleGroup object
х	A SingleGroup object
multiple	A real number; the multiple of the smoothed standard deviation to call significant.
ccl	A list containing objects of the ColorCoding class. If left at its default value of zero, colors are chosen automatically.
main	Plot title
xlab	Label for the x axis

ylab	Label for the y axis
xlim	Plotting limits for the x axis. If left at the default value of zero, then the limits are automatically generated
ylim	Plotting limits for the y axis. If left at the default value of zero, then the limits are automatically generated
row.names	See the base version of as.data.frame.default
optional	See the base version of as.data.frame.default

The usual extra parameters to generic or plotting routines

Details

In 2001 and 2002, Baggerly and Coombes developed the smooth t-test for finding differentially expressed genes in microarray data. Along with many others, they began by log-transforming the data as a reasonable step in the direction of variance stabilization. They observed, however, that The gene-by-gene standard deviations still seemed to vary in a systematic way as a function of the mean log intensity. By borrowing strength across genes and using **loess** to fit the observed standard deviations as a function of the mean, one presumably got a better estimate of the true standard deviation.

Creating Objects

Objects can be created by calls to the SingleGroup constructor. Users rarely have need to create these objects directly; they are usually created as a consequence of the construction of an object of the SmoothTtest class.

Slots

name: A character string; the name of this object

avg: The numeric vector of mean values

sd: The numeric vector of standard deviations

span: The span parameter used in the loess function to fit sd as a function of avg.

fit: A list containing components x and y resulting from the loess fit.

score: A numeric vector; the ratio of the pointwise standard deviations to their smooth (loess) estimates.

Methods

as.data.frame(x, row.names=NULL, optional=FALSE) Combine the slots containing numeric vectors into a data frame, suitable for printing or exporting.

summary(object, ...) Write out a summary of the object.

print(x, ...) Print the entire object. You never want to do this.

plot(x, multiple=3, ccl=0, main=x@name, xlab='Mean', ylab='Std Dev', xlim=0, ylim=0, ...)
Produce a scatter plot of the standard deviations (x@sd) as a function of the means
(x@avg). The appropriate multiple of the loess fit is overlaid, and points that exceed

this multiple are flaged in a different color. Colors in the plotare controlled by the current values of COLOR.CENTRAL.LINE, COLOR.CONFIDENCE.CURVE, COLOR.BORING, COLOR.BAD.REPLICATE, and COLOR.WORST.REPLICATE.

Author(s)

Kevin R. Coombes <kcoombes@mdanderson.org>

References

Baggerly, K.A., Coombes, K.R., Hess, K.R., Stivers, D.N., Abruzzo, L.V., Zhang, W. Identifying differentially expressed genes in cDNA microarray experiments. J Comp Biol. 8:639-659, 2001.

Coombes, K.R., Highsmith, W.E., Krogmann, T.A., Baggerly, K.A., Stivers, D.N., Abruzzo, L.V. Identifying and quantifying sources of variation in microarray data using high-density cDNA membrane arrays. J Comp Biol. 9:655-669, 2002.

See Also

SmoothTtest

Examples

```
m <- rnorm(1000, 8, 2.5)
v <- rnorm(1000, 0.7)
plot(m, v)
x <- SingleGroup(m, v, name='bogus')
summary(x)
plot(x)
plot(x, multiple=2)
# cleanup
rm(m, v, x)
```

TwoGroupStats-class The TwoGroupStats Class

Description

Compute row-by-row means and variances for a data matrix whose columns belong to wto different groups of interest.

Usage

```
TwoGroupStats(data, classes, name = comparison, name1 = A, name2 = B)
## S4 method for signature 'TwoGroupStats':
as.data.frame(x, row.names=NULL, optional=FALSE)
## S4 method for signature 'TwoGroupStats':
summary(object, ...)
## S4 method for signature 'TwoGroupStats':
print(x, ...)
## S4 method for signature 'TwoGroupStats, missing':
plot(x, main=x@name, useLog=FALSE, ...)
```

Arguments

data	Either a data frame or matrix with numeric values or an exprSet as defined in the BioConductor tools for analyzing microarray data.
classes	If data is a data frame or matrix, then classes must be either a logical vector or a factor. If data is an exprSet, then classes can be a character string that names one of the factor columns in the associated phenoData subobject.
name	A character string; the name of this object
name1	A character string; the name of the first group
name2	A character string; the name of the second group
x	A TwoGroupStats object
row.names	See the base version of as.data.frame.default
optional	See the base version of as.data.frame.default
object	A TwoGroupStats object
main	Plot title
useLog	a logical flag; should the values be log-transformed before plotting?
	The usual extra arguments to generic functions

Details

This class was one of the earliest developments in our suite of tools to analyze microarrays. Its main purpose is to segregate out the preliminary computation of summary statistics on a row-by-row basis, along with a set of plots that could be generated automatically and used for quality control.

Creating Objects

As usual, objects can be created by **new**, but better methods are available in the form of the **TwoGroupStats** function. The inputs to this function are the same as those used for row-by-row statistical tests throughout the ClassComparison package; a detailed description can be found in the **MultiTtest** class.

One should note that this class serves as the front end to the SmoothTtest class, providing it with an interface that accepts exprSet objects compatible with the other statistical tests in the ClassComparison package.

Slots

mean1: The numeric vector of means in the first group

mean2: The numeric vector of means in the second group

overallMean: The numeric vector of overall row means

var1: The numeric vector of variances in the first group

var2: The numeric vector of variances in teh second group

- overallVar: The numeric vector of variances assuming the two groups have the same mean
- n1: The number of items in the first group
- n2: The number of items in the first group
- name1: The name of the first group
- name2: The name of the second group
- name: The name of the object

Methods

- as.data.frame(x, row.names=NULL, optional=FALSE) Collect the numeric vectors from the object into a single dat fame, suitable for printing or exporting.
- summary(object, ...) Write out a summary of the object.
- print(x, ...) Print the object. (Actually, it only prints a summary, since the whole object is almost always more than you really want to see. If you insist on printing everything, use as.data.frame.)
- plot(x, main=x@name, useLog=FALSE, ...) This function actually produces six different plots of the data, so it is usually wrapped by a graphical layout command like par(mfrow=c(2,3)). The first two plots show the relation between the mean and standard deviation for the two groups separately; the third plot does the same for the overall mean and variance. The fourth plot is a Bland-Altman of the differnce between the means against the overall mean. (In the microarray world, this is usually called an M-vs-A plot.) A loess fit is overlaid on the scatter plot, and points outside confidence bounds based on the fit are printed in a differnt color to flag them as highly variable. The fifth plot shows a loess fit (with confidence bounds) of the difference as a function of the row index (which often is related to the geometric position of spots on a microarray). Thus, this plot gives a possible indication of regions of an array where unusual things happen. The final plot compares the overall variances to the pooled variances.

Author(s)

Kevin R. Coombes <kcoombes@mdanderson.org>

References

Altman DG, Bland JM. Measurement in Medicine: the Analysis of Method Comparison Studies. The Statistician, 1983; 32: 307-317.

See Also

SmoothTtest, MultiTtest

Examples

```
bogus <- matrix(rnorm(30*1000, 8, 3), ncol=30, nrow=1000)
splitter <- rep(FALSE, 30)
splitter[16:30] <- TRUE
x <- TwoGroupStats(bogus, splitter)
summary(x)
opar<-par(mfrow=c(2,3), pch='.')
plot(x)
par(opar)
# cleanup
rm(bogus, splitter, x, opar)</pre>
```

Bum-class

The Bum Class

Description

The Bum class is used to fit a beta-uniform mixture model to a set of p-values.

Usage

```
Bum(pvals, ...)
## S4 method for signature 'Bum':
summary(object, tau=0.01, ...)
## S4 method for signature 'Bum':
hist(x, res=100, xlab='P Values', main="', ...)
## S4 method for signature 'Bum':
image(x, ...)
## S4 method for signature 'Bum':
cutoffSignificant(object, alpha, by='FDR', ...)
## S4 method for signature 'Bum':
selectSignificant(object, alpha, by='FDR', ...)
## S4 method for signature 'Bum':
countSignificant(object, alpha, by='FDR', ...)
likelihoodBum(object)
```

Arguments

pvals	A numeric vector containing values between 0 and 1
object	A Bum object
tau	A real number between 0 and 1, representing a cutoff on the p-values.

х	A Bum object
res	A positive integer; the resolution at which to plot the fitted distribution curve.
xlab	Label for the x axis
main	Graph title
alpha	<pre>Either the false discovery rate (if by = 'FDR') or the posterior probability (if by = 'EmpiricalBayes')</pre>
by	String denoting the method to use for determining cutoffs. The chioces are 'FDR', 'FalseDiscovery', or 'EmpiricalBayes'. Since the test is implemented with match.arg, unique abbreviations also work.
	All methods are defined to accept additional arguments in order to allow flexibility in designing derived classes. The usual graphical parameters can be supplied to hist and image.

Details

The BUM method was introduced by Stan Pounds and Steve Morris, although it was simultaneously discovered by several other researchers. It is generally applicable to any analysis of microarray or proteomics data that performs a separate statistical hypothesis test for each gene or protein, where each test produces a p-value that would be valid if the analyst were only performing one statistical test. When performing thousands of statistical tests, however, those p-values no longer have the same interpretation as Type I error rates. The idea behind BUM is that, under the null hypothesis that none of the genes or proteins is interesting, the expected distribution of the set of p-values is uniform. By contrast, if some of the genes are interesting, then we should see an overabundance of small p-values (or a spike in the histogram near zero). We can model the alternative hypothesis with a beta distribution, and view the set of all p-values as a mixture distribution.

Fitting the BUM model is straightforward, using a nonlinear optimizer to compute the maximum likelihood parameters. After the model has been fit, one can easily determine cutoffs on the p-values that correspond to desired false discovery rates. Alternatively, the original Pounds and Morris paper shows that their results can be reinterpreted to recover the empirical Bayes method introduced by Efron and Tibshirani. Thus, one can also determine cutoffs by specifying a desired posterior probability of significance.

Value

Graphical functions (hist and image) invisbly return the object on which they were invoked.

The cutoffSignficant method returns a real number between zero and one. P-values below this cutoff are considered statistically significant at either the specified false discovery rate or at the specified posterior probability.

The selectSignficant method returns a vector of logical values whose length is equal to the length of the vector of p-values that was used to construct the Bum object. True values in the return vector mark the statistically signficant p-values.

The countSignificant method returns an integer, the number of statistically significant p-values.

The summary method returns an object of class BumSummary.

Creating Objects

Although objects can be created directly using **new**, the most common usage will be to pass a vector of p-values to the **Bum** function.

Slots

pvals: The vector of p-values used to construct the object.

ahat: Model parameter

lhat: Model parameter

pihat: Model parameter

Methods

- summary(object, tau=0.01, ...) For each value of the p-value cutoff tau, computes estimates of the fraction of true positives (TP), false negatives (FN), false positives (FP), and true negatives (TN).
- hist(x, res=100, xlab='P Values', main=", ...) Plots a histogram of the object, and overlays (1) a straight line to indicate the contribution of the uniform component and (2) the fitted beta-uniform distribution from the observed values. Colors in the plot are controlled by COLOR.EXPECTED and COLOR.OBSERVED.
- image(x, ...) Produces four plots in a 2x2 layout: (1) the histogram produced by hist;
 (2) a plot of cutoffs against the desired false discovery rate; (3) a plot of cutoffs against the posterior probability of coming from the beat component; and (4) an ROC curve.
- cutoffSignificant(object, alpha, by='FDR', ...) Computes the cutoff needed for significance, which in this case means aridsing from the beta component rarther than the uniform component of the mixture. Significance is specified either by the false discovery rate (when by = 'FDR' or by = 'FalseDiscovery') or by the posterior probability (when by = 'EmpiricalBayes')
- countSignificant(object, alpha, by='FDR', ...) Uses selectSignificant to count the number of significant p-values.

Author(s)

Kevin R. Coombes <kcoombes@mdanderson.org>

References

Pounds S, Morris SW. Estimating the occurrence of false positives and false negatives in microarray studies by approximating and partitioning the empirical distribution of p-values. Bioinformatics. 2003 Jul 1;19(10):1236-42.

Benjamini Y, Hochberg Y. Controlling the false discovery rate: a practical and powerful approach to multiple testing. J Roy Statist Soc B, 1995; 57: 289-300.

Efron B, Tibshirani R: Empirical bayes methods and false discovery rates for microarrays. Genet Epidemiol 2002, 23: 70-86.

See Also

Two classes that produce lists of p-values that can (and often should) be analyzed using BUM are MultiTtest and MultiLinearModel. Also see BumSummary.

Examples

```
fake.data <- c(runif(700), rbeta(300, 0.3, 1))</pre>
a <- Bum(fake.data)
hist(a, res=200)
alpha <- (1:25)/100
plot(alpha, cutoffSignificant(a, alpha, by='FDR'),
     xlab='Desired False Discovery Rate', type='1',
     main='FDR Control', ylab='Significant P Value')
GAMMA <- 5*(10:19)/100
plot(GAMMA, cutoffSignificant(a, GAMMA, by='EmpiricalBayes'),
     ylab='Significant P Value', type='1',
     main='Empirical Bayes', xlab='Posterior Probability')
b <- summary(a, (0:100)/100)
be <- b@estimates
sens <- be$TP/(be$TP+be$FN)</pre>
spec <- be$TN/(be$TN+be$FP)</pre>
plot(1-spec, sens, type='l', xlim=c(0,1), ylim=c(0,1), main='ROC Curve')
points(1-spec, sens)
abline(0,1)
image(a)
countSignificant(a, 0.05, by='FDR')
countSignificant(a, 0.99, by='Emp')
#cleanup
rm(a, b, be, alpha, GAMMA, sens, spec, fake.data)
```

BumSummary-class The BumSummary class

Description

An implementation class. Users are not expected to create these objects directly; they are produced as return objects from the summary method for Bum.

Slots

bum: Object of class Bum
estimates: Object of class data.frame
Fhat: Object of class numeric

Methods

show signature(object = "BumSummary"): Print the object, which contains a summary
 of the underlying Bum object. The summary contains a data frame with estimates
 of the fraction of true positives (TP), false postives (FP), true negatives (TN) and
 false negatives (FN) at the set of p-value cutoffs specified in the call to the summary
 method.

Author(s)

Kevin R. Coombes <kcoombes@mdanderson.org>

See Also

Bum

Examples

See the examples for the Bum class.

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Wilcoxon Density Function

Description

Computes the density function for the Wilcoxon rank-sum distribution without centering.

Usage

dwil(q, m, n)

Arguments

q	vector of quantiles
m	number of observations in the first sample
n	number of observations in the second sample

Details

Computes the density function for the Wilcoxon rank-sum distribution, using exact values when both groups have fewer than 50 items and switching to a normal approximation otherwise. It was originally written for S-Plus, which still perversely insists that **m** and **n** must be less than 50. The function was retained when the OOMPA library was ported to R, since S-Plus keeps the actual rank-sum but R centers the distribution at zero. This function encapsulated the difference, allowing everything else to continue to work as it had worked previously.

Value

A vector of the same length as ${\tt q}$ containing (approximate or exact) values of the density function.

Author(s)

Kevin R. Coombes <kcoombes@mdanderson.org>

See Also

MultiWilcoxonTest

Examples

dwil(51:60, 9, 3) dwil(51:60, 9, 51)

```
matrixT
```

Multiple T Tests by Matrix Multiplication

Description

Utility functions for computing vectors of row-by-row means, variances, and t-statistics.

Usage

```
matrixMean(x)
matrixVar(x, xmean)
matrixT(m, v)
```

Arguments

х	a matrix
xmean	a numeric vector or single-column matrix
m	a matrix
v	a logical vector of length eual to the number of columns of ${\tt m}$

Value

matrixMean returns a single-column matrix containing the row-by-row means of x.

matrixVar returns a signle-column matrix containing the row-by-row means of x, assuming that xmean contains the corresponding mean values.

matrixT returns a single-column matrix of t-statistics from a two-sample t-test comparing the columns for which v is true to those for which v is false.

Author(s)

Kevin R. Coombes <kcoombes@mdanderson.org>

See Also

MultiTtest

Examples

```
ng <- 1000
ns <- 50
dat <- matrix(rnorm(ng*ns), ncol=ns)
clas <- factor(rep(c('Good', 'Bad'), each=25))
myMean <- matrixMean(dat)
myVar <- matrixVar(dat, myMean)
plot(myMean, myVar)
myT <- matrixT(dat, clas)
hist(myT)
rm(ng, ns, dat, myMean, myVar, myT)
```

MultiLinearModel-class

The MultiLinearModel Class

Description

Class to fit multiple (row-by-row) linear (fixed-effects) models on microarray or proteomics data.

Usage

```
MultiLinearModel(form, clindata, arraydata)
## S4 method for signature 'MultiLinearModel':
summary(object, ...)
## S4 method for signature 'MultiLinearModel':
hist(x, xlab='F Statistics', main=NULL, ...)
## S4 method for signature 'MultiLinearModel, missing':
plot(x, ylab='F Statistics', ...)
## S4 method for signature 'MultiLinearModel, ANY':
plot(x, y, xlab='F Statistics',
 ylab=deparse(substitute(y)), ...)
## S4 method for signature 'MultiLinearModel':
anova(object, ob2, ...)
multiTukey(object, alpha)
```

Arguments

form	A formula object specifying the linear model
clindata	Either a data frame of "clinical" or other covariates or an exprSet.
arraydata	A matrix or data frame of values to be explained by the model. If clindata is an exprSet, then arraydata can be omitted, since it is assumed to be part of the exprSet.
object	A MultiLinearModel object
ob2	Another MultiLinearModel object
x	A MultiLinearModel object
У	A numeric vector
xlab	Label for the x-axis
ylab	Label for the y-axis
main	Graph title
	Optional graphical or other parameters to generic functions
alpha	A real number between 0 and 1; the significance level for the Tukey test.

Value

The anova method returns a data frame. The rows in the data frame correpsonds to the rows in the arraydata object that was used to construct the MultiLinearModel objects. The first column contains the F-statistics and the second column contains the p-values.

The multiTukey function returns a vector whosem length equals the number of rows in the arraydata object used to construct the MultiLinearModel. Assuming that the overall F-test was significant, differences in group means (in each data row) larger than this value are significant by Tukey's test for honoestly significant difference. (Of course, that statement is incorrect, since we haven't fully corrected for multiple testing. Our standard practice is to take the p-values from the row-by-row F-tests and evaluate themusing the beta-uniform mixture model (see Bum). For the rows that correspond to models whose p-values are smaller than the Bum cutoff, we simply use the Tukey HSD values without further modification.)

Creating Objects

Objects should be created by calling the MultiLinearModel function. The first argument is a formula specifying the linear model, in the same manner that it would be passed to lm. We will fit the linear model separately for each row in the arraydata matrix. Rows of arraydata are attached to the clindata data frame and are always referred to as "Y" in the formulas. In particular, this implies that clindata can not include a column already called "Y". Further, the implementation only works if "Y" is the response variable in the model.

Multiple linear models with "exprSet" objects

The BioConductor packages uses an exprSet to combine microarray data and clinical covariates (known in their context as phenoData objects) into a single structure. You can call MultiLinearModel using an exprSet object for the clindata argument. In this case, the function extracts the phenoData slot of the exprSet to use for the clinical covariates, and extracts the exprs slot of the exprSet object to use for the array data.

Slots

call: A call object describing how the object was constructed.

model: The formula object specifying the linear model.

F.statistics: A numeric vector of F-statistics comparing the linear model to the null model.

p.values: A numeric vector containing the p-values associated to the F-statistics.

coefficients: A matrix of the coefficients in the linear models.

predictions: A matrix of the (Y-hat) values predicted by the models.

sse: A numeric vector of the sum of squared error terms from fitting the models.

ssr: A numeric vector of the sum of squared regression terms from the model.

df: A numeric vector (of length two) containing the degrees of freedom for the F-tests.

Methods

summary(object, ...) Write out a summary of the object.

hist(x, xlab='F Statistics', main=NULL, ...) Create a histogram of the F-statistics.

- plot(x, ylab='F Statistics', ...) Plot the F-statistics as a function of the row index.
- plot(x, y, xlab='F Statistics', ylab=deparse(substitute(y)), ...) Plot the F-statistics
 against the numeric vector y.
- anova(object, ob2, ...) Perform row-by-row F-tests comparing two different linear models.

Details

The MultiLinearModel constructor computes row-by-row F-tests comparing each linear model to the null model Y 1. In many instances, one wishes to use an F-test to compare two different linear models. For instance, many standard applications of analysis of variance (ANOVA) can be described using such a compoarison between two different linear models. The anova method for the MultiLinearModel class performs row-by-row F-tests comparing two competing linear models.

The implementation of MultiLinearModel does not take the naive approach of using either apply or a for-loop to attach rows one at a time and fit separate linear models. All the models are actually fit simultaneously by a series of matrix operations, which greatly reduces the amount of time needed to compute the models. The constraint on the column names in clindata still holds, since one row is attached to allow model.matrix to determine the contrasts matrix.

Author(s)

Kevin R. Coombes <kcoombes@mdanderson.org>

See Also

MultiTtest, MultiWilcoxonTest, Bum, lm, anova.

Examples

```
ng <- 10000
ns <- 50
dat <- matrix(rnorm(ng*ns), ncol=ns)</pre>
cla <- factor(rep(c('A', 'B'), 25))</pre>
cla2 <- factor(rep(c('X', 'Y', 'Z'), times=c(15, 20, 15)))</pre>
covars <- data.frame(Grade=cla, Stage=cla2)</pre>
res <- MultiLinearModel(Y ~ Grade + Stage, covars, dat)</pre>
summary(res)
hist(res, breaks=101)
plot(res)
plot(res, res@p.values)
graded <- MultiLinearModel(Y ~ Grade, covars, dat)</pre>
summary(graded)
hist(graded@p.values, breaks=101)
hist(res@p.values, breaks=101)
oop <- anova(res, graded)</pre>
hist(oop$p.values, breaks=101)
# cleanup
rm(ng, ns, dat, cla, cla2, covars, res, graded, oop)
```

MultiTtest-class The MultiTtest Class

Description

Class to perform row-by-row t-tests on microarray or proteomics data.

Usage

```
MultiTtest(data, classes)
## S4 method for signature 'MultiTtest':
summary(object, ...)
## S4 method for signature 'MultiTtest':
hist(x, xlab='T Statistics', main=NULL, ...)
## S4 method for signature 'MultiTtest, missing':
plot(x, ylab='T statistics', ...)
## S4 method for signature 'MultiTtest, ANY':
plot(x, y, xlab='T Statistics, ylab=deparse(substitute(y)), ...)
```

Arguments

data Either a data frame or matrix with numeric values or an exprSet as defined in the BioConductor tools for analyzing microarray data.

classes	If data is a data frame or matrix, then classes must be either a logical vector or a factor. If data is an exprSet, then classes can be a character string that names one of the factor columns in the associated phenoData subobject.
object	A MultiTtest object
x	A MultiTtest object
У	A numeric vector
xlab	Label for the x axis
ylab	Label for the y axis
main	Plot title
	The usual graphical parameters can by supplied to hist and plot methods.

Value

The graphical routines invisibly return the object against which they were invoked.

Creating objects

Although objects can be created using **new**, the better method is to use the MultiTtest function. In the simplest case, you simply pass in a data matrix and a logical vector assigning classes to the columns, and the constructor performs row-by-row two-sample t-tests and computes the associated (single test) p-values. To adjust for multiple testing, you can pass the p-values on to the Bum class.

If you use a factor instead of a logical vector, then the t-test compares the first level of trhe factor to everything else. To handle the case of multiple classes, see the MultiLinearModel class.

As with other class comparison functions that are part of the OOMPA, we can also perform statistical tests on **exprSet** objects from the BioConductor libraries. In this case, we pass in an **exprSet** object along with the name of a factor to use for splitting the data.

Slots

t.statistics: Object of class numeric containing the computed t-statistics.

p.values: Object of class numeric containing the computed p-values.

groups: Object of class character containing the names of the classes being compared.

call: Object of class call containing the function call that created the object.

Methods

summary(object, ...) Write out a summary of the object.

hist(x, xlab='T Statistics', main=NULL, ...) Produce a histogram of the t-statistics.

plot(x Produces a scatter plot of the t-statistics against their index.

plot(x,y) Produces a scatter plot of the t-statistics in the object x against the numeric vector y.

Author(s)

Kevin R. Coombes <kcoombes@mdanderson.org>

See Also

matrixT, Bum, MultiLinearModel, Dudoit

Examples

```
ng <- 10000
ns <- 50
dat <- matrix(rnorm(ng*ns), ncol=ns)
cla <- factor(rep(c('A', 'B'), each=25))
res <- MultiTtest(dat, cla)
summary(res)
hist(res, breaks=101)
plot(res)
plot(res, res@p.values)
hist(res@p.values, breaks=101)</pre>
```

rm(ng, ns, dat, cla, res)

MultiWilcoxonTest-class

The MultiWilcoxonTest Class

Description

The MultiWilcoxonTest class is used to perform row-by-row Wilcoxon rank-sum tests on a data matrix. Significance cutoffs are determined by the empirical Bayes method of Efron and Tibshirani

Usage

```
MultiWilcoxonTest(data, classes, histsize = NULL)
## S4 method for signature 'MultiWilcoxonTest':
summary(object, prior=1, significance=0.9, ...)
## S4 method for signature 'MultiWilcoxonTest':
hist(x, xlab='Rank Sum',
ylab='Prob(Different | Y)', main='', ...)
## S4 method for signature 'MultiWilcoxonTest, missing':
plot(x, prior=1, significance=0.9,
ylim=c(-0.5, 1), xlab='Rank Sum', ylab='Prob(Different | Y)', ...)
## S4 method for signature 'MultiWilcoxonTest':
cutoffSignificant(object, prior, significance, ...)
## S4 method for signature 'MultiWilcoxonTest':
selectSignificant(object, prior, significance, ...)
## S4 method for signature 'MultiWilcoxonTest':
countSignificant(object, prior, significance, ...)
```

Arguments

data	Either a data frame or matrix with numeric values or an exprSet as defined in the BioConductor tools for analyzing microarray data.
classes	If data is a data frame or matrix, then classes must be either a logical vector or a factor. If data is an exprSet, then classes can be a character string that names one of the factor columns in the associated phenoData subobject.
histsize	An integer; the number of bins used for the hostogram summarizing the Wilcoxon statistics. When NULL, each discrete rank-sum value gets its own bin.
object	an object of the MultiWilcoxonTest class.
x	an object of the MultiWilcoxonTest class.
xlab	Label for the x axis
ylab	Label for the y axis
ylim	Plotting limits on the y=axis
main	Graph title
prior	Prior probability that an arbitrary gene is not differentially expressed, or that an arbitrary row does not yield a significant Wilcoxon rank-sum statistic.
significance	Desired level of posterior probability
	Additional graphical parameters.

Details

See the paper by Efron and Tibshirani.

Value

The standard methods summary, hist, and plot return what you would expect.

The cutoffSignificant method returns a list of two integers. Rank-sum values msaller than the first value or larger than the second value are statistically significant in the sense that their posterior probability exceeds the specified significance level given the assumptions about the prior probability of not being significant.

The selectSignficant method returns a vector of logical values identifying the significant test results, and countSignificant returns an integer counting the number of significant test results.

Creating Objects

As usual, objects can be created by **new**, but better methods are available in the form of the MultiWilcoxonTest function. The inputs to this function are the same as those used for row-by-row statistical tests throughout the ClassComparison package; a detailed description can be found in the MultiTtest class. The constructor computes row-by-row Wilcoxon rank-sum statistics on the input data, comparing the two groups defined by the classes argument. It also estimates the observed and theoretical (expected) density functions for the collection of rank-sum statistics.

The additional input argument, **histsize** is usually best left to its default value. In certain pathological cases, we have found it necessary to use fewer bins; one suspects that the underlying model does not adequately capture the complexity of those situations.

Slots

rank.sum.statistics: A numeric vector containing the computed rank-sum statistics.

- xvals: A numeric vector, best thought of as the vector of possible rank-sum statistics given the sizes of the two groups.
- pdf: A numeric vector containing the empirical density function computed at the points of xvals.
- unravel: A numeric vector containing a smoothed estimate (by Poisson regression using B-splines) of the empirical density function evaluated at xvals.
- groups: A vector containing the names of the groups defined by classes.

call: An object of class call representing the function call that created the object.

Methods

- summary(object, prior=1, significance=0.9, ...) Write out a summary of the object.
 For a given value of the prior probability of not being differentially expressed and a
 given significance cutoff on the posterior probability, reports the cutoffs and number
 of items in both tails of the distribution.
- hist(x, xlab='Rank Sum', ylab='Prob(Different|Y)', main=", ...) Plot a histogram of the rank-sum statistics, with overlaid curves representing the expected and observed distributions. Colors of the curves are controlled by COLOR.EXPECTED and COLOR.OBSERVED.
- plot(x, prior=1, significance=0.9, ylim=c(-0.5, 1), xlab='Rank Sum', ylab='Prob(Different | Y)', .
 Plots the posterior probability of being differentially expressed for given values of the
 prior. Horixontal lines are added at each specified significance level for the poste rior probability.
- cutoffSignificant(object, prior, significance, ...) Determine cutoffs on the rank-sum statistic at the desired significance level.
- selectSignificant(object, prior, significance, ...) Compute a logical vector for selecting significant test results.
- countSignificant(object, prior, significance, ...) Count the number of significant test
 results.

Author(s)

Kevin R. Coombes <kcoombes@mdanderson.org>

References

Efron B, Tibshirani R: Empirical bayes methods and false discovery rates for microarrays. Genet Epidemiol 2002, 23: 70-86.

Pounds S, Morris SW. Estimating the occurrence of false positives and false negatives in microarray studies by approximating and partitioning the empirical distribution of p-values. Bioinformatics. 2003 Jul 1;19(10):1236-42.

See Also

Implementation is handled in part by the functions dwil and rankSum. The empirical Bayes results for alternative tests (such as MultiTtest) can be obtained using the beta-uniform mixture model in the Bum class.

Examples

```
ng <- 10000
ns <- 15
nd <- 200
fake.class <- factor(rep(c('A', 'B'), each=ns))</pre>
fake.data <- matrix(rnorm(ng*ns*2), nrow=ng, ncol=2*ns)</pre>
fake.data[1:nd, 1:ns] <- fake.data[1:nd, 1:ns] + 2</pre>
fake.data[(nd+1):(2*nd), 1:ns] <- fake.data[(nd+1):(2*nd), 1:ns] - 2</pre>
a <- MultiWilcoxonTest(fake.data, fake.class)</pre>
hist(a)
plot(a)
plot(a, prior=0.85)
abline(h=0)
cutoffSignificant(a, prior=0.85, signif=0.95)
countSignificant(a, prior=0.85, signif=0.95)
# cleanup
rm(ng, ns, nd, fake.class, fake.data, a)
```

rankSum

Wilcoxon Rank-Sum Statistic

Description

Compute the Wilcoxon rank-sum statistic

Usage

```
rankSum(data, selector)
```

Arguments

data	A numeric vector
selector	A logical vector the same length as data.

Details

This is an efficient function to compute the value of the Wilcoxon rank-sum statistic without the extra overhead of the full wilcox.test function. It is used internally by the MultiWilcoxonTest class to perform rwo-by-row Wilcoxon tests.

Value

Returns an integer, the rank-sum of the subset of the data for which the selector is true.

Author(s)

Kevin R. Coombes <kcoombes@mdanderson.org>

See Also

dwil, MultiWilcoxonTest

Examples

```
dd <- rnorm(100)
cc <- rep(c(TRUE, FALSE), each=50)
rankSum(dd, cc)
rm(cc, dd)</pre>
```

SamSummary-class The SamSummary Class

Description

An implementation class. Users are not expected to create these objects directly; they are produced as return objects from the summary method for Sam.

Slots

fdr: A number between 0 and 1; the expected false discovery rate

hi: Upper threshold for significance

10: Lower threshold for significance

cutoff: A numeric value specified in the call to the Sam summary method.

significant.calls: vector of logical values

average.false.count: The average number of alse positives in the permuted data at this cutoff level.

Methods

show signature(object = SamSummary): Print the object, which contains a summary of the underlying Sam object. In particular, the summary reports the number of genes acheiving each possible number of misclassifications.

Author(s)

Kevin R. Coombes <kcoombes@mdanderson.org>

See Also

Sam

Examples

See the examples for the Sam class.

significant

Generic Functions for Significance

Description

In the world of multiple testing that is inhabited by most microarray or protein profiling experiments, analysts frequently perform separate statistical tests for each gene or protein in the experiment. Determining cutoffs that achieve statistical significance (in a meaningful way) is an inherent part of the procedure. It is then common to select the significant items for further processing or for preparing reports, or at least to count the number of significant items. These generic functions provide a standard set of tools for selecing and counting the significant items, which can be used with various statistical tests and various ways to account for multiple testing.

Usage

```
cutoffSignificant(object, ...)
selectSignificant(object, ...)
countSignificant(object, ...)
```

Arguments

object	Typically, an object that performs multiple statistical tests on microarray or proteomics data.
	All generic methods are designed to take optional additional argument for flexibility in creating derived classes.

Value

cutoffSignificant returns appropriate cutoff values that achieve specified significance criteria.

selectSignificant returns a logical vector, with true values indicating items that satisfy the cutoff making them statistically significant.

countSignificant returns an integer, representing the numgbe of significant items.

Author(s)

Kevin R. Coombes <kcoombes@mdanderson.org>

See Also

Classes that implement these methods include Bum, Sam, Dudoit, MultiWilcoxonTest, and TNoM.

Examples

- # Since these are generic methods, there are no examples. See the
- # classes that implement specific instances.

SmoothTtest-class The SmoothTtest Class

Description

Implements the smooth t-test for differential expression as developed by Baggely and Coombes.

Usage

```
SmoothTtest(stats, aname = Group One, bname = Group Two,
name = paste(aname, vs., bname))
## S4 method for signature 'SmoothTtest':
as.data.frame(x, row.names=NULL, optional=FALSE)
## S4 method for signature 'SmoothTtest':
summary(object, ...)
## S4 method for signature 'SmoothTtest, missing':
plot(x, folddiff=3, goodflag=2, badch=4, ccl=0,
name=x@name, pch='.', xlab='log intensity', ylab='log ratio', ...)
```

Arguments

stats	An object of the TwoGroupStats class.
aname	A character string; the name of the second group
bname	A character string; the name of the second group
name	A character string; the name of this object
object	A SmoothTtest object
x	A SmoothTtest object
row.names	See the base version of as.data.frame.default
optional	See the base version of as.data.frame.default
folddiff	A real number; the level of fold difference considered large enough to be indicated in the plots.

goodflag	A real number; the level (in standard deviation units) of the smooth t- statistic considered large enough to be indicated in the plot.
badch	A real number; the level of variability in single groups considered large enough to be worrisome. See the multiple argument to the plot method in the SingleGroup class.
ccl	A list containing objects of the $\tt ColorCoding$ class. If left at its default value of zero, colors are chosen automatically.
pch	Default plotting character
xlab	Label for the x axis
ylab	Label for the y axis
• • •	

The usual extra parameters to generic or plotting routines

Details

In 2001 and 2002, Baggerly and Coombes developed the smooth t-test for finding differentially expressed genes in microarray data. Along with many others, they began by log-transforming the data as a reasonable step in the direction of variance stabilization. They observed, however, that The gene-by-gene standard deviations still seemed to vary in a systematic way as a function of the mean log intensity. By borrowing strength across genes and using **loess** to fit the observed standard deviations as a function of the mean, one presumably got a better estimate of the true standard deviation.

These smooth estimates are computed for each of two groups of samples being compared. They are then combined (gene-by-gene using the usual univariate formulas) to compute pooled "smooth" estimates of the standard deviation. These smooth estimates are then used in gene-by-gene t-tests.

The interesting question then arises of how to compute and interpret p-values associated to these individual tests. The liberal argument asserts that, because smoothing uses data from hundreds of measurements to estiamte the standard deviation, it can effectively be treated as "known" in the t-tests, which should thus be compared against the normal distribution. A conservative argument claims that the null distribution should still be the t-distribution with the degrees of freedom determined in the usual way by the number of samples. The truth probably lies somewhere in between, and is probably best approximated by some kind of permutation test. In this implementation, we take the coward's way out and don't provide any of those alternatives. You have to extract the t-statistics (from the smooth.t.statistics slot of the object) and compute your own p-values in your favorite way. If you base the computations on a theoretical model rather than a permutation test, then the Bum class provides a convenient way to account for multiple testing.

Creating Objects

In practice, users will first use a data frame and a classification vector (or an exprSet) to construct an object of the TwoGroupStats object. This object can then be handed directly to the SmoothTtest function to perform the smooth t-test.

Slots

- one: An object of the SingleGroup class representing a loess smooth of standard deviation as a function of the mean in the first group of samples.
- two: An object of the SingleGroup class representing a loess smooth of standard deviation as a function of the mean in the second group of samples.
- smooth.t.statistics: A numeric vector containing the smooth t-statistics
- fit: A data.frame containing two columns, x and y, containing the smooth estimates of the pooled standard deviation
- dif: A numeric vector of the differences in mean values between the two groups

avg: A numeric vector of the overall mean value

aname: A character string; the name of the first group

- bname: A character string; the name of the second group
- name: A character string; the name of this object

stats: The TwoGroupStats object that was used to create this object.

Methods

as.data.frame(x, row.names=NULL, optional=FALSE) Convert the object into a data frame suitable for printing or exporting.

summary(object, ...) Write out a summary of the object.

plot(x, folddiff=3, goodflag=2, badch=4, ccl=0, name=x@name, pch='.', xlab='log intensity', ylab Create a set of six plots. The first two plots are the QC plots from the SingleGroup objects representing the two groups of samples. The third plot is a scatter plot comparing the means in the two groups. The fourth plot is Bland-Altman plot of the overall mean against the difference in means (also known colloquially as an M-vs-A plot). The fifth plot is a histogram of the smooth t-statistics. The final plot is a scatter plot of the smooth t-statistics as a function of the mean intensity. Colors in the plots are controlled by the curent values of COLOR.BORING, COLOR.SIGNIFICANT, COLOR.BAD.REPLICATE, COLOR.WORST.REPLICATE, COLOR.FOLD.DIFFERENCE, COLOR.CENTRAL.LINE, AND COLOR.CONFIDENCE.CURVE.

Author(s)

Kevin R. Coombes <kcoombes@mdanderson.org>

References

Baggerly, K.A., Coombes, K.R., Hess, K.R., Stivers, D.N., Abruzzo, L.V., Zhang, W. Identifying differentially expressed genes in cDNA microarray experiments. J Comp Biol. 8:639-659, 2001.

Coombes, K.R., Highsmith, W.E., Krogmann, T.A., Baggerly, K.A., Stivers, D.N., Abruzzo, L.V. Identifying and quantifying sources of variation in microarray data using high-density cDNA membrane arrays. J Comp Biol. 9:655-669, 2002.

Altman DG, Bland JM. Measurement in Medicine: the Analysis of Method Comparison Studies. The Statistician, 1983; 32: 307-317.

See Also

Bum, MultiTtest, SingleGroup, TwoGroupStats.

Examples

```
bogus <- matrix(rnorm(30*1000, 8, 3), ncol=30, nrow=1000)
splitter <- rep(FALSE, 30)
splitter[16:30] <- TRUE
x <- TwoGroupStats(bogus, splitter)
y <- SmoothTtest(x)
opar <- par(mfrow=c(2, 3), pch='.')
plot(y, badch=2, goodflag=1)
par(opar)
# cleanup
rm(bogus, splitter, x, y, opar)</pre>
```

TNoMSummary-class The TNoMSummary Class

Description

An implementation class. Users are not expected to create these objects directly; they are produced as return objects from the summary method for TNoM.

Slots

TNoM: Object of class TNoM

counts: Object of class numeric

Methods

show signature(object = TNoMSummary): Print the object, which contains a summary
 of the underlying TNoM object. In particular, the summary reports the number of genes
 acheiving each possible number of misclassifications.

Author(s)

Kevin R. Coombes <kcoombes@mdanderson.org>

See Also

TNoM

Examples

See the examples for the TNoM class.

TNoM-class

Description

Implements the "Total Number of Misclassifications" method for finding differentially expressed genes.

Usage

```
TNoM(data, classes, verbose = TRUE)
## S4 method for signature 'TNoM':
summary(object, ...)
## S4 method for signature 'TNoM':
update(object, nPerm, verbose=FALSE, ...)
## S4 method for signature 'TNoM':
selectSignificant(object, cutoff, ...)
## S4 method for signature 'TNoM':
countSignificant(object, cutoff, ...)
## S4 method for signature 'fullTNoM, missing':
plot(x, ...)
## S4 method for signature 'fullTNoM':
```

```
hist(x, ...)
```

Arguments

data	Either a data frame or matrix with numeric values or an exprSet as defined in the BioConductor tools for analyzing microarray data.
classes	If data is a data frame or matrix, then classes must be either a logical vector or a factor. If data is an exprSet, then classes can be a character string that names one of the factor columns in the associated phenoData subobject.
verbose	A logical flag; whether to print out intermediate results
object	A TNoM object
nPerm	An integer; the number of permutations to perform
cutoff	An integer
x	A fullTNoM object
	Additional plotting or other arguments.

Details

The TNoM method was developed by Yakhini and Ben-Dor and first applied in the melanoma microarray study by Bittner and colleagues (see references). The goal of the method is to detect genes that are differentially expressed between two groups of samples. The idea is

that each gene serve as a potential classifier to distinguish the two groups. One starts by determining an optimal cutoff on the expression of each gene and counting the number of miscalssifications that gene makes. Next, we bin genes based on the total number of misclassifications. This distribution can be compared with the expected value (by simulating normal data sets of the same size). Alternatively, one can estimate the null distribution directly by srambling the sample labels to perform a permutation test.

The TNoM constructor computes the optimal cutoffs and the misclassification rates. The update method performs the simulations and permutation tests, producing an object of the fullTNoM class.

Value

summary returns a TNoMSummary object.

update returns a fullTNoM object.

selectSignificant returns a vector of logical values.

countSignificant returns an integer.

Creating Objects

As usual, objects can be created by **new**, but better methods are available in the form of the **TNoM** function. The inputs to this function are the same as those used for row-by-row statistical tests throughout the ClassComparison package; a detailed description can be found in the **MultiTtest** class.

Slots

Objects of the TNoM class have the following slots:

data: The data matrix used to contruct the object

tnomData: A numeric vector, whose length is the number of rows in data, recording the minimum number of misclassification acheived using this data row.

nCol: The number of columns in data

nRow: The number of rows in data

classifier: The classification vector used to create the object.

call: The function call that created the object

Objects of the fullTNoM class have the following slots:

dex: Numeric vector of the different possible numbers of misclassifications

fakir: Numeric vector of expected values based on simulations

obs: Numeric vector of observed values

scr: Numeric vector of values based on a permutation test

name: A character string with a name for the object

Methods

Objects of the TNoM class have the following methods:

- summary(object, ...) Write out a summary of the object, including the number of genes acheiving each possible number of misclassifications.
- countSignificant(object, cutoff, ...) Count the number of significant genes at the given
 cutoff.
- selectSignificant(object, cutoff, ...) Get a vector for selecting the number of significant
 genes at the given cutoff.
- update(object, nPerm, verbose=FALSE, ...) Perform simulation and permutation tests on the TNoM object.

Objects of the fullTNoM class have the following methods:

plot(x, ...) Plot a summary of the TNoM object. This consists of three curves: the observed cumulative number of genes at each misclassification level, along with the corresponding numbers expected based on simulations or permutation tests. The colors of the curves are controlled by the values of COLOR.OBSERVED, COLOR.EXPECTED, and COLOR.PERMTEST

hist(x, ...) Plot a not terribly useful nor informative histogram of the results.

Author(s)

Kevin R. Coombes <kcoombes@mdanderson.org>

References

Bittner M, Meltzer P, Chen Y, Jiang Y, Seftor E, Hendrix M, Radmacher M, Simon R, Yakhini Z, Ben-Dor A, Sampas N, Dougherty E, Wang E, Marincola F, Gooden C, Lueders J, Glatfelter A, Pollock P, Carpten J, Gillanders E, Leja D, Dietrich K, Beaudry C, Berens M, Alberts D, Sondak V. Molecular classification of cutaneous malignant melanoma by gene expression profiling. Nature. 2000 Aug 3;406(6795):536-40.

See Also

MultiTtest, MultiWilcoxonTest, link{Bum}

Examples

```
n.genes <- 200
n.samples <- 10
```

```
bogus <- matrix(rnorm(n.samples*n.genes, 0, 3), ncol=n.samples)
splitter <- rep(FALSE, n.samples)
splitter[sample(1:n.samples, trunc(n.samples/2))] <- TRUE</pre>
```

tn <- TNoM(bogus, splitter)
summary(tn)</pre>

tnf <- update(tn)</pre>

plot(tnf)
hist(tnf)

rm(bogus, splitter, n.genes, n.samples, tn, tnf)