allNA

A filter function to determine if all elements of a vector are NA.

Description

allNA evaluates to FALSE if all elements of its argument are NA. anyNA evaluates to FALSE if any of the elements of its argument are NA.

Usage

allNA(x)
anyNA(x)

Arguments

x The vector to test.

Value

FALSE if all elements of x are NA.

Author(s)

R. Gentleman

See Also

pOverA

Examples

allNA(NA)
allNA(1)
anyNA(1)
anyNA(NA)
Description

Anova returns a function of one argument with bindings for cov and p. The function, when evaluated, performs an ANOVA using cov as the covariate. It returns TRUE if the p value for a difference in means is less than p.

Usage

Anova(cov, p=0.05, na.rm=TRUE)

Arguments

cov The covariate. It must have length equal to the number of columns of the array that Anova will be applied to.
p The p-value for the test.
na.rm If set to TRUE any NA’s will be removed.

Details

The function returned by Anova uses lm to fit a linear model of the form \( lm(x \sim cov) \), where \( x \) is the set of gene expressions. The F statistic for an overall effect is computed and if it has a \( p \)-value less than \( p \) the function returns TRUE, otherwise it returns FALSE for that gene.

Value

Anova returns a function with bindings for cov and p that will perform a one-way ANOVA. The covariate can be continuous, in which case the test is for a linear effect for the covariate.

Author(s)

R. Gentleman

See Also

kOverA, lm

Examples

set.seed(123)
af <- Anova(c(rep(1,5),rep(2,5)), .01)
af(rnorm(10))
**coxfilter**

A filter function for univariate Cox regression.

**Description**

A function that performs Cox regression with bindings for `surt`, `cens`, and `p` is returned. This function filters genes according to the attained p-value from a Cox regression using `surt` as the survival times, and `cens` as the censoring indicator. It requires `survival`.

**Usage**

```r
coxfilter(surt, cens, p)
```

**Arguments**

- `surt`: Survival times.
- `cens`: Censoring indicator.
- `p`: The p-value to use in filtering.

**Value**

Calls to the `coxph` function in the `survival` library are used to fit a Cox model. The filter function returns `TRUE` if the p-value in the fit is less than `p`.

**Author(s)**

R. Gentleman

**See Also**

`Anova`

**Examples**

```r
set.seed(-5)
sfun <- coxfilter(rexp(10), ifelse(runif(10) < .7, 1, 0), .05)
ffun <- filterfun(sfun)
dat <- matrix(rnorm(1000), ncol=10)
out <- genefilter(dat, ffun)
```
A filter function for the coefficient of variation.

Description

cv returns a function with values for a and b bound. This function takes a single argument. It computes the coefficient of variation for the input vector and returns TRUE if the coefficient of variation is between a and b. Otherwise it returns FALSE.

Usage

cv(a=1, b=Inf, na.rm=TRUE)

Arguments

a The lower bound for the cv.
b The upper bound for the cv.
na.rm If set to TRUE any NA’s will be removed.

Details

The coefficient of variation is the standard deviation divided by the absolute value of the mean.

Value

It returns a function of one argument. The function has an environment with bindings for a and b.

Author(s)

R. Gentleman

See Also

pOverA, kOverA

Examples

set.seed(-3)
cvfun <- cv(1,10)
cvfun(rnorm(10,10))
cvfun(rnorm(10))
Calculate an n-by-n matrix by applying a function to pairs of columns of an m-by-n matrix.

**Usage**

```r
dist2(x, fun=function(a,b) mean(abs(a-b), na.rm=TRUE), diagonal=0)
```

**Arguments**

- `x`: A matrix, or any object for which `ncol(x)` and `x[,j]` return appropriate results.
- `fun`: A symmetric function of two arguments that may be columns of `x`.
- `diagonal`: The value to be used for the diagonal elements of the resulting matrix.

**Details**

With the default value of `fun`, this function calculates for each pair of columns of `x` the mean of the absolute values of their differences (which is proportional to the L1-norm of their difference). This is a distance metric.

The implementation assumes that `fun` is symmetric, `fun(a, b) = fun(b, a)`. Hence, the returned matrix is symmetric. `fun(a, a)` is not evaluated, instead the value of `diagonal` is used to fill the diagonal elements of the returned matrix.

A use for this function is the detection of outlier arrays in a microarray experiment. Assume that each column of `x` can be decomposed as `z + β + ϵ`, where `z` is a fixed vector (the same for all columns), `ϵ` is vector of `nrow(x)` i.i.d. random numbers, and `β` is an arbitrary vector whose majority of entries are negligibly small (i.e. close to zero). In other words, `Dz` the probe effects, `ϵ` measurement noise and `β` differential expression effects. Under this assumption, all entries of the resulting distance matrix should be the same, namely a multiple of the standard deviation of `ϵ`. Arrays whose distance matrix entries are way different give cause for suspicion.

**Value**

A symmetric matrix of size `n x n`.

**Examples**

```r
z = matrix(rnorm(15693), ncol=3)
dist2(z)
```
eSetFilter  

A function to filter an eSet object

Description

Given a Bioconductor’s ExpressionSet object, this function filters genes using a set of selected filters.

Usage

eSetFilter(eSet)
getFilterNames()
getFuncDesc(lib = "genefilter", funcs = getFilterNames())
getRdAsText(lib)
parseDesc(text)
parseArgs(text)
showESet(eSet)
setESetArgs(filter)
isESet(eSet)

Arguments

eSet: an ExpressionSet object
lib: a character string for the name of an R library where functions of interests reside
funcs: a vector of character strings for names of functions of interest
text: a character of string from a field (e.g., description, argument, ..) field of an Rd file for a function
filter: a character string for the name of a filter function

Details

A set of filters may be selected to filter genes in through each of the filters in the order the filters have been selected.

Value

A logical vector of length equal to the number of rows of `expr`. The values in that vector indicate whether the corresponding row of `expr` passed the set of filter functions.

Author(s)

Jianhua Zhang

See Also

genefilter


filtered_p

Examples

```r
if( interactive() ) {
  data(sample.ExpressionSet)
  res <- eSetFilter(sample.ExpressionSet)
}
```

filtered_p | **Compute and adjust p-values, with filtering**

Description

Given filter and test statistics in the form of unadjusted p-values, or functions able to compute these statistics from the data, filter and then correct the p-values across a range of filtering stringencies.

Usage

```r
filtered_p(filter, test, theta, data, method = "none")
filtered_R(alpha, filter, test, theta, data, method = "none")
```

Arguments

- `alpha`: A cutoff to which p-values, possibly adjusted for multiple testing, will be compared.
- `filter`: A vector of stage-one filter statistics, or a function which is able to compute this vector from `data`, if `data` is supplied.
- `test`: A vector of unadjusted p-values, or a function which is able to compute this vector from the filtered portion of `data`, if `data` is supplied. The option to supply a function is useful when the value of the test statistic depends on which hypotheses are filtered out at stage one. (The `limma` t-statistic is an example.)
- `theta`: A vector with one or more filtering fractions to consider. Actual cutoffs are then computed internally by applying `quantile` to the filter statistics contained in (or produced by) the `filter` argument.
- `data`: If `filter` and/or `test` are functions rather than vectors of statistics, they will be applied to `data`. The functions will be passed the whole `data` object, and must work over rows, etc. themselves as appropriate.
- `method`: The unadjusted p-values contained in (or produced by) `test` will be adjusted for multiple testing after filtering, using the `p.adjust` function in the `stats` package. See the `method` argument there for options.

Value

For `filtered_p`, a matrix of p-values, possible adjusted for multiple testing, with one row per null hypothesis and one column per filtering fraction given in `theta`. For a given column, entries which have been filtered out are NA.

For `filtered_R`, a count of the entries in the `filtered_p` result which are less than `alpha`.

Author(s)

Richard Bourgon <bourgon@ebi.ac.uk>
filterfun

See Also

See rejection_plot for visualization of filtered_p results.

Examples

# See the vignette: Diagnostic plots for independent filtering

definition

filterfun(...) Creates a first FALSE exiting function from the list of filter functions it is given.

Description

This function creates a function that takes a single argument. The filtering functions are bound in the environment of the returned function and are applied sequentially to the argument of the returned function. When the first filter function evaluates to FALSE the function returns FALSE otherwise it returns TRUE.

Usage

filterfun(...) Filtering functions.

Value

filterfun returns a function that takes a single argument. It binds the filter functions given to it in the environment of the returned function. These functions are applied sequentially (in the order they were given to filterfun). The function returns FALSE (and exits) when the first filter function returns FALSE otherwise it returns TRUE.

Author(s)

R. Gentleman

See Also
genefilter

Examples

set.seed(333)
x <- matrix(rnorm(100,2,1),nc=10)
cvfun <- cv(.5,2.5)
ffun <- filterfun(cvfun)
which <- genefilter(x, ffun)
Description

Generate a volcano plot contrasting p-value with fold change (on the log scale), in order to visualize the effect of filtering on overall variance and also assign significance via p-value.

Usage

```r
filter_volcano(
  d, p, S,
  n1, n2,
  alpha, S_cutoff,
  cex = 0.5, pch = 19,
  xlab = expression(paste(log[2], " fold change")),
  ylab = expression(paste("-", log[10], " p")),
  cols = c("grey80", "grey50", "black"),
  ltys = c(1, 3),
  use_legend = TRUE,
  ...
)
```

Arguments

d  Fold changes, typically on the log scale, base 2.

p  The p-values

S  The overall standard deviation filter statistics, i.e., the square roots of the overall variance filter statistics.

n1  Sample size for group 1.

n2  Sample size for group 2.

alpha  Significance cutoff used for p-values.

S_cutoff  Filter cutoff used for the overall standard deviation in S.

xlab  Label for x-axis.

ylab  Label for y-axis.

cols  A vector of three colors used for plotting. These correspond to filtered data, data which pass the filter but are insignificant, and data pass the filter and are also statistically significant.

ltys  The induced bound on log-scale fold change is plotted, as is the significance cutoff for data passing the filter. The ltys argument gives line styles for these drawing these two thresholds on the plot.

use_legend  Should a legend for point color be produced?

...  Other arguments for plot.
Author(s)
Richard Bourgon <bourgon@ebi.ac.uk>

Examples
# See the vignette: Diagnostic plots for independent filtering

```
findLargest
```

Find the Entrez Gene ID corresponding to the largest statistic

Description
Most microarrays have multiple probes per gene (Entrez). This function finds all replicates, and then selects the one with the largest value of the test statistic.

Usage
```
findLargest(gN, testStat, data = "hgu133plus2")
```

Arguments
- `gN`: A vector of probe identifiers for the chip.
- `testStat`: A vector of test statistics, of the same length as `gN` with the per probe test statistics.
- `data`: The character string identifying the chip.

Details
All the probe identifiers, `gN`, are mapped to Entrez Gene IDs and the duplicates determined. For any set of probes that map to the same Gene ID, the one with the largest test statistic is found. The return vector is the named vector of selected probe identifiers. The names are the Entrez Gene IDs.

This could be extended in different ways, such as allowing the user to use a different selection criterion. Also, matching on different identifiers seems like another alternative.

Value
A named vector of probe IDs. The names are Entrez Gene IDs.

Author(s)
R. Gentleman

See Also
- `sapply`
Examples

```r
library("hgu95av2.db")
set.seed(124)
gN <- sample(ls(hgu95av2ENTREZID), 200)
stats <- rnorm(200)
findLargest(gN, stats, "hgu95av2")
```

Description

The `gapFilter` looks for genes that might usefully discriminate between two groups (possibly unknown at the time of filtering). To do this we look for a gap in the ordered expression values. The gap must come in the central portion (we exclude jumps in the initial `Prop` values or the final `Prop` values). Alternatively, if the IQR for the gene is large that will also pass our test and the gene will be selected.

Usage

```r
gapFilter(Gap, IQR, Prop, na.rm=TRUE, neg.rm=TRUE)
```

Arguments

- **Gap**: The size of the gap required to pass the test.
- **IQR**: The size of the IQR required to pass the test.
- **Prop**: The proportion (or number) of samples to exclude at either end.
- **na.rm**: If TRUE then NA's will be removed before processing.
- **neg.rm**: If TRUE then negative values in x will be removed before processing.

Details

As stated above we are interested in

Value

A function that returns either TRUE or FALSE depending on whether the vector supplied has a gap larger than `Gap` or an IQR (inter quartile range) larger than `IQR`. For computing the gap we want to exclude a proportion, `Prop` from either end of the sorted values. The reason for this requirement is that genes which differ in expression levels only for a few samples are not likely to be interesting.

Author(s)

R. Gentleman

See Also

ttest, genefilter
Examples

```r
set.seed(256)
x <- c(rnorm(10,100,3), rnorm(10, 100, 10))
y <- x + c(rep(0,10), rep(100,10))
tmp <- rbind(x,y)
Gfilter <- gapFilter(200, 100, 5)
ffun <- filterfun(Gfilter)
genefilter(tmp, ffun)
```

Description

`genefilter` filters genes in the array `expr` using the filter functions in `flist`. It returns an array of logical values (suitable for subscripting) of the same length as there are rows in `expr`. For each row of `expr` the returned value is `TRUE` if the row passed all the filter functions. Otherwise it is set to `FALSE`.

Usage

```r
genefilter(expr, flist)
```

Arguments

- `expr` A matrix or ExpressionSet that the filter functions will be applied to.
- `flist` A list of filter functions to apply to the array.

Details

This package uses a very simple but powerful protocol for filtering genes. The user simply constructs any number of tests that they want to apply. A test is simply a function (as constructed using one of the many helper functions in this package) that returns `TRUE` if the gene of interest passes the test (or filter) and `FALSE` if the gene of interest fails.

The benefit of this approach is that each test is constructed individually (and can be tested individually). The tests are then applied sequentially to each gene. The function returns a logical vector indicating whether the gene passed all tests functions or failed at least one of them.

Users can construct their own filters. These filters should accept a vector of values, corresponding to a row of the `expr` object. The user defined function should return a length 1 logical vector, with value `TRUE` or `FALSE`. User-defined functions can be combined with `filterfun`, just as built-in filters.

Value

A logical vector of length equal to the number of rows of `expr`. The values in that vector indicate whether the corresponding row of `expr` passed the set of filter functions.

Author(s)

R. Gentleman
**genefinder**

*Finds genes that have similar patterns of expression.*

**Description**

Given an `ExpressionSet` or a matrix of gene expressions, and the indices of the genes of interest, genefinder returns a list of the `numResults` closest genes. The user can specify one of the standard distance measures listed below. The number of values to return can be specified. The return value is a list with two components: genes (measured through the desired distance method) to the genes of interest (where X is the number of desired results returned) and their distances.

**Usage**

```r
genefinder(X, ilist, numResults=25, scale="none", weights, method="euclidean")
```

**Arguments**

- **X**: A numeric matrix where columns represent patients and rows represent genes.
- **ilist**: A vector of genes of interest. Contains indices of genes in matrix X.
- **numResults**: Number of results to display, starting from the least distance to the greatest.
- **scale**: One of "none", "range", or "zscore". Scaling is carried out separately on each row.
- **weights**: A vector of weights applied across the columns of X. If no weights are supplied, no weights are applied.
- **method**: One of "euclidean", "maximum", "manhattan", "canberra", "correlation", "binary".

**Details**

If the `scale` option is "range", then the input matrix is scaled using `genescale()`(). If it is "zscore", then the input matrix is scaled using the `scale` builtin with no arguments.

The method option specifies the metric used for gene comparisons. The metric is applied, row by row, for each gene specified in `ilist`.

The "correlation" option for the distance method will return a value equal to 1-correlation(x).

See `dist` for a more detailed description of the distances.
Value

The returned value is a list containing an entry for each gene specified in ilist. Each list entry contains an array of distances for that gene of interest.

Author(s)

J. Gentry and M. Kajen

See Also

genescale

Examples

set.seed(12345)

# create some fake expression profiles
m1 <- matrix(1:12, 4, 3)
v1 <- 1
nr <- 2

# find the 2 rows of m1 that are closest to row 1
genefinder(m1, v1, nr, method="euc")

v2 <- c(1,3)
genefinder(m1, v2, nr)

ngenefinder(m1, v2, nr, scale="range")

genefinder(m1, v2, nr, method="manhattan")

m2 <- matrix(rnorm(100), 10, 10)
v3 <- c(2, 5, 6, 8)
nr2 <- 6

ngenefinder(m2, v3, nr2, scale="zscore")

---

genescale Scales a matrix or vector.

Description

genescale returns a scaled version of the input matrix m by applying the following formula to each column of the matrix:

\[ y[i] = (x[i] - min(x))/(max(x) - min(x)) \]

Usage

genescatle(m, axis=2, method=c("Z", "R"), na.rm=TRUE)
half.range.mode

Arguments

- **m**: Input a matrix or a vector with numeric elements.
- **axis**: An integer indicating which axis of m to scale.
- **method**: Either "Z" or "R", indicating whether a Z scaling or a range scaling should be performed.
- **na.rm**: A boolean indicating whether NA's should be removed.

Details

Either the rows or columns of m are scaled. This is done either by subtracting the mean and dividing by the standard deviation ("Z") or by subtracting the minimum and dividing by the range.

Value

A scaled version of the input. If m is a matrix or a dataframe then the dimensions of the returned value agree with that of m, in both cases the returned value is a matrix.

Author(s)

R. Gentleman

See Also

genefinder.scale

Examples

```r
m <- matrix(1:12, 4, 3)
genecountx(m)
```

half.range.mode  Mode estimation for continuous data

Description

For data assumed to be drawn from a unimodal, continuous distribution, the mode is estimated by the “half-range” method. Bootstrap resampling for variance reduction may optionally be used.

Usage

```r
half.range.mode(data, B, B.sample, beta = 0.5, diag = FALSE)
```

Arguments

- **data**: A numeric vector of data from which to estimate the mode.
- **B**: Optionally, the number of bootstrap resampling rounds to use. Note that B = 1 resamples 1 time, whereas omitting B uses data as is, without resampling.
- **B.sample**: If bootstrap resampling is requested, the size of the bootstrap samples drawn from data. Default is to use a sample which is the same size as data. For large data sets, this may be slow and unnecessary.
- **beta**: The fraction of the remaining range to use at each iteration.
- **diag**: Print extensive diagnostics. For internal testing only... best left FALSE.
Details

Briefly, the mode estimator is computed by iteratively identifying densest half ranges. (Other fractions of the current range can be requested by setting \( \beta \) to something other than 0.5.) A densest half range is an interval whose width equals half the current range, and which contains the maximal number of observations. The subset of observations falling in the selected densest half range is then used to compute a new range, and the procedure is iterated. See the references for details.

If bootstrapping is requested, \( B \) half-range mode estimates are computed for \( B \) bootstrap samples, and their average is returned as the final estimate.

Value

The mode estimate.

Author(s)

Richard Bourgon <bourgon@stat.berkeley.edu>

References


See Also

- `shorth`

Examples

```r
## A single normal-mixture data set
x <- c( rnorm(10000), rnorm(2000, mean = 3) )
M <- half.range.mode( x )
M.bs <- half.range.mode( x, B = 100 )

if(interactive()){  
  hist( x, breaks = 40 )
  abline( v = c( M, M.bs ), col = "red", lty = 1:2 )
  legend( 1.5, par("usr")[4],
    c( "Half-range mode", "With bootstrapping (B = 100)" ),
    lwd = 1, lty = 1:2, cex = .8, col = "red"
  )
}

# Sampling distribution, with and without bootstrapping
X <- rbind(  
  matrix( rnorm(1000 * 100), ncol = 100 ),
  matrix( rnorm(200 * 100, mean = 3), ncol = 100 )
)
M.list <- list(  
  Simple = apply( X, 2, half.range.mode )
)```
kappa_p

BS = apply( X, 2, half.range.mode, B = 100 )

if(interactive()){ 
  boxplot( M.list, main = "Effect of bootstrapping" )
  abline( h = 0, col = "red" )
}

kappa_p

Compute proportionality constant for fold change bound.

Description

Filtering on overall variance induces a lower bound on fold change. This bound depends on the significance of the evidence against the null hypothesis, an is a multiple of the cutoff used for an overall variance filter. It also depends on sample size in both of the groups being compared. These functions compute the multiplier for the supplied p-values or t-statistics.

Usage

kappa_p(p, n1, n2 = n1)
kappa_t(t, n1, n2 = n1)

Arguments

p
  The p-values at which to compute the multiplier.

t
  The t-statistics at which to compute the multiplier.

n1
  Sample size for class 1.

n2
  Sample size for class 2.

Value

A vector of multipliers: one per p-value or t-static in p or t.

Author(s)

Richard Bourgon <bourgon@ebi.ac.uk>

Examples

# See the vignette: Diagnostic plots for independent filtering
### kOverA

A filter function for \( k \) elements larger than \( A \).

**Description**

kOverA returns a filter function with bindings for \( k \) and \( A \). This function evaluates to TRUE if at least \( k \) of the arguments elements are larger than \( A \).

**Usage**

```r
kOverA(k, A=100, na.rm=TRUE)
```

**Arguments**

- **A**: The value you want to exceed.
- **k**: The number of elements that have to exceed \( A \).
- **na.rm**: If set to TRUE any NA’s will be removed.

**Value**

A function with bindings for \( A \) and \( k \).

**Author(s)**

R. Gentleman

**See Also**

pOverA

**Examples**

```r
fg <- kOverA(5, 100)
fg(90:100)
fg(98:110)
```

### maxA

A filter function to filter according to the maximum.

**Description**

maxA returns a function with the parameter \( A \) bound. The returned function evaluates to TRUE if any element of its argument is larger than \( A \).

**Usage**

```r
maxA(A=75, na.rm=TRUE)
```
Arguments

A The value that at least one element must exceed.
na.rm If TRUE then NA’s are removed.

Value

maxA returns a function with an environment containing a binding for A.

Author(s)

R. Gentleman

See Also

pOverA

Examples

ff <- maxA(30)
ff(1:10)
ff(28:31)

nsFilter

Non-Specific Filtering of Features in an ExpressionSet

Description

This various provides various options for filtering (removing) features from an ExpressionSet.

Experience with microarray data suggests that features exhibiting little variation, or a consistently low signal, across samples typically target transcripts that are simply not expressed at detectable levels in any of the samples. It can be advantageous for the subsequent data analysis to remove them beforehand (see Bourgon et al., References).

Furthermore, one may decide that there is little value in considering features with insufficient annotation.

Usage

nsFilter(eset, require.entrez=TRUE,
require.GOBP=FALSE, require.GOCC=FALSE,
require.GOMF=FALSE, require.CytoBand=FALSE,
remove.dupEntrez=TRUE, var.func=IQR,
var.cutoff=0.5, var.filter=TRUE,
filterByQuantile=TRUE, feature.exclude="^AFFX", ...)

varFilter(eset, var.func=IQR, var.cutoff=0.5, filterByQuantile=TRUE)

featureFilter(eset, require.entrez=TRUE,
require.GOBP=FALSE, require.GOCC=FALSE,
require.GOMF=FALSE, require.CytoBand=FALSE,
remove.dupEntrez=TRUE, feature.exclude="^AFFX")
Arguments

eset  an ExpressionSet object

require.entrez  If TRUE, require that all probe sets have an Entrez Gene ID annotation. Probe sets without such an annotation will be filtered out. If using a package where Entrez Gene IDs are not the central ID, then that ID will be required instead.

require.GOBP  If TRUE, require that all probe sets have an annotation to at least one GO ID in the BP ontology. Probe sets without such an annotation will be filtered out.

require.GOCC  If TRUE, require that all probe sets have an annotation to at least one GO ID in the CC ontology. Probe sets without such an annotation will be filtered out.

require.GOMF  If TRUE, require that all probe sets have an annotation to at least one GO ID in the MF ontology. Probe sets without such an annotation will be filtered out.

require.CytoBand  If TRUE, require that all probe sets (for higher organisms) have at least a mapping to cytoband locations labeled p1, p2, p3, q1, q2, q3, and etc. Probe sets without cytoband locations will be filtered out. It would be useful when performing GSEA tests within chromosome band locations.

remove.dupEntrez  If TRUE and there are multiple probe sets mapping to the same Entrez Gene ID, then the probe set with the largest value of var.func will be retained and the others removed. If using a package where Entrez Gene IDs are not the central ID, then duplicates will be filtered using this other ID type.

var.func  A function that will be used to assess the variance of a probe set across all samples. This function should return a numeric vector of length one when given a numeric vector as input. Probe sets with a var.func value less than var.cutoff will be removed. The default is IQR.

var.cutoff  A numeric value to use in filtering out probe sets with small variance across samples. See the var.func argument and the details section below.

var.filter  A logical indicating whether or not to perform variance based filtering. The default is TRUE.

filterByQuantile  Logical: whether the variance-filter cutoff threshold should be interpreted as a quantile. Defaults to TRUE; if set to FALSE the cutoff value is used directly "as is".

feature.exclude  A character vector of regular expressions. Any probe sets identifiers (return value of featureNames(eset)) that match one of the specified patterns will be filtered out. The default value is intended to filter out Affymetrix quality control probe sets.

...  Unused, but available for specializing methods.

Details

In this Section, the effect of filtering on the type I error rate estimation / control of subsequent hypothesis testing is explained. A more comprehensive discussion is given by Bourgon et al. (see Reference).

Marginal type I errors: Filtering on the basis of a statistic which is independent of the test statistic used for detecting differential gene expression can increase the detection rate at the same marginal type I error. This is clearly the case for filter criteria that do not depend on the data, such as the
annotation based criteria provided by the `nsFilter` and `featureFilter` functions. However, marginal type I error can also be controlled for certain types of data-dependent criteria. Call $U^I$ the stage 1 filter statistic, which is a function that is applied feature by feature, based on whose value the feature is or is not accepted to pass to stage 2, and which depends only on the data for that feature and not any other feature, and call $U^{II}$ the stage 2 test statistic for differential expression. Sufficient conditions for marginal type-I error control are:

- $U^I$ the overall (across all samples) variance or mean, $U^{II}$ the t-statistic (or any other scale and location invariant statistic), data normal distributed and exchangeable across samples.
- $U^I$ the overall mean, $U^{II}$ the moderated t-statistic (as in limma’s `eBayes` function), data normal distributed and exchangeable.
- $U^I$ a sample-class label independent function (e.g. overall mean, median, variance, IQR), $U^{II}$ the Wilcoxon rank sum statistic, data exchangeable.

Experiment-wide type I error: Marginal type-I error control provided by the conditions above is sufficient for control of the family wise error rate (FWER). Note, however, that common false discovery rate (FDR) methods depend not only on the marginal behaviour of the test statistics under the null hypothesis, but also on their joint distribution. The joint distribution can be affected by filtering, even when this filtering leaves the marginal distributions of true-null test statistics unchanged. Filtering might, for example, change correlation structure. The effect of this is negligible in many cases in practice, but this depends on the dataset and the filter used, and the assessment is in the responsibility of the data analyst.

Annotation Based Filtering Arguments `require.entrez`, `require.GOBP`, `require.GOCC`, and `require.GOMF` turn on a filter based on available annotation data. The annotation package is determined by calling `annotation(eset)`.

Duplicate Probe Removal If `remove.dupEntrez=TRUE`, probes determined by your annotation to be pointing to the same gene will be compared, and only the probe with the highest `var.func` value will be retained.

Variance Based Filtering The `var.filter`, `var.func`, `var.cutoff` and `varByQuantile` arguments control numerical cutoff-based filtering. Probes for which `var.func` returns NA are removed. The default `var.func` is `IQR`, which is defined as `rowQ(eset, ceiling(0.75 * ncol(eset))) - rowQ(eset, floor(0.25 * ncol(eset)))`; this choice is motivated by the observation that unexpressed genes are detected most reliably through low variability of their features across samples. Additionally, IQR is robust to outliers (see note below). The default `var.cutoff` is 0.5 and is motivated by the rule of thumb that in many tissues only 40% of genes are expressed. Of course, if you believe in a different approach to numerical filtering you can choose another function as `var.func`, or turn off numerical filtering by setting `var.filter=FALSE`.

Note that by default the numerical-filter cutoff is interpreted as a quantile, so leaving the default values intact would filter out 50% of the genes remaining at this stage. If you prefer to set the cutoff at some absolute threshold, change the value of `varByQuantile` to `FALSE`, and modify `var.cutoff` accordingly.

Note also that variance filtering is performed last, so that if `varByQuantile=TRUE` and `remove.dupEntrez=TRUE`, the final number of genes does indeed exclude precisely the `var.cutoff` fraction of unique genes remaining after all other filters were passed.

The stand-alone function `varFilter` does only numerical filtering, and returns an `ExpressionSet`. `featureFilter` does only feature based filtering and duplicate removal, and returns an expression set as well. Duplicate removal is hard-coded to retain the highest-IQR probe for each gene.

Value

For `nsFilter` a list consisting of:
eset      the filtered ExpressionSet
filter.log a list giving details of how many probe sets were removed for each filtering step performed.

For both varFilter and featureFilter the filtered ExpressionSet.

Note
IQR is a reasonable variance-filter choice when the dataset is split into two roughly equal and relatively homogeneous phenotype groups. If your dataset has important groups smaller than 25% of the overall sample size, or if you are interested in unusual individual-level patterns, then IQR may not be sensitive enough for your needs. In such cases, you should consider using less robust and more sensitive measures of variance (the simplest of which would be sd).

Author(s)
Seth Falcon (somewhat revised by Assaf Oron)

References

Examples
library("hgu95av2.db")
library("Biobase")
data(sample.ExpressionSet)
ans <- nsFilter(sample.ExpressionSet)
ansest
ans$filter.log

## skip variance-based filtering
ans <- nsFilter(sample.ExpressionSet, var.filter=FALSE)

a1 <- varFilter(sample<ExpressionSet)
a2 <- featureFilter(sample.ExpressionSet)

pOverA  A filter function to filter according to the proportion of elements larger than A.

Description
A function that returns a function with values for A, p and na.rm bound to the specified values. The function takes a single vector, x, as an argument. When the returned function is evaluated it returns TRUE if the proportion of values in x that are larger than A is at least p.

Usage
pOverA(p=0.05, A=100, na.rm=TRUE)
rejection_plot

Arguments

A       The value to be exceeded.
p       The proportion that need to exceed A for TRUE to be returned.
na.rm   If TRUE then NA's are removed.

Value

pOverA returns a function with bindings for A, p and na.rm. This function evaluates to TRUE if the proportion of values in x that are larger than A exceeds p.

Author(s)

R. Gentleman

See Also

cv

Examples

ff<- pOverA(p=.1, 10)
ff(1:20)
ff(1:5)

description

Plot the number, or fraction, of null hypotheses rejected as a function of the p-value cutoff. Multiple sets of p-values are accepted, in a list or in the columns of a matrix, in order to permit comparisons.

Usage

rejection_plot(
  p,
  col, lty = 1, lwd = 1,
  xlab = "Cutoff", ylab = "Rejections",
  xlim = c(0, 1), ylim,
  legend = names(p),
  at = c("all", "sample"),
  n_at = 100,
  probability = FALSE,
  ...
)
Arguments

p  The p-values to be used for plotting. These may be in the columns of a matrix, or in the elements of a list. One curve will be generated for each column/element, and all NA entries will be dropped. If column or element names are supplied, they are used by default for a plot legend.

col  Colors to be used for each curve plotted. Recycled if necessary. If col is omitted, rainbow is used to generate a set of colors.
lty  Line styles to be used for each curve plotted. Recycled if necessary.
lwd  Line widths to be used for each curve plotted. Recycled if necessary.
xlab  X-axis text label.
ylab  Y-axis text label.
xlim  X-axis limits.
ylim  Y-axis limits.
legend  Text for legend. Matrix column names or list element names (see p above) are used by default. If NULL, no legend is plotted.
at  Should step functions be plotted with a step at every value in p, or should linear interpolation be used at a sample of points spanning xlim? The latter looks when there are many p-values.
n_at  When at = "sample" is given, how many sample points should be used for interpolation and plotting?
probability  Should the fraction of null hypotheses rejected be reported instead of the count? See the probability argument to hist.
...  Other arguments to pass to the plot call which sets up the axes. Note that the ... argument will not be passed to the lines calls which actually generate the curves.

Value

A list of the step functions used for plotting is returned invisibly.

Author(s)

Richard Bourgon <bourgon@ebi.ac.uk>

Examples

# See the vignette: Diagnostic plots for independent filtering

rowFtests  t-tests and F-tests for rows or columns of a matrix

Description

t-tests and F-tests for rows or columns of a matrix, intended to be speed efficient.
**rowFtests**

**Usage**

```r
rowttests(x, fac, tstatOnly = FALSE)
colttests(x, fac, tstatOnly = FALSE)
fastT(x, ig1, ig2, var.equal = TRUE)
```

```r
rowFtests(x, fac, var.equal = TRUE)
colFtests(x, fac, var.equal = TRUE)
```

**Arguments**

- **x**: Numeric matrix. The matrix must not contain NA values. For `rowttests` and `colttests`, `x` can also be an `ExpressionSet`.
- **fac**: Factor which codes the grouping to be tested. There must be 1 or 2 groups for the t-tests (corresponding to one- and two-sample t-test), and 2 or more for the F-tests. If `fac` is missing, this is taken as a one-group test (i.e. is only allowed for the t-tests). The length of the factor needs to correspond to the sample size: for the `row*` functions, the length of the factor must be the same as the number of columns of `x`, for the `col*` functions, it must be the same as the number of rows of `x`.
- **tstatOnly**: A logical variable indicating whether to calculate p-values from the t-distribution with appropriate degrees of freedom. If `TRUE`, just the t-statistics are returned. This can be considerably faster.
- **ig1**: The indices of the columns of `x` that correspond to group 1.
- **ig2**: The indices of the columns of `x` that correspond to group 2.
- **var.equal**: A logical variable indicating whether to treat the variances in the samples as equal. If 'TRUE', a simple F test for the equality of means in a one-way analysis of variance is performed. If 'FALSE', an approximate method of Welch (1951) is used, which generalizes the commonly known 2-sample Welch test to the case of arbitrarily many samples.

**Details**

If `fac` is specified, `rowttests` performs for each row of `x` a two-sided, two-class t-test with equal variances. `fac` must be a factor of length `ncol(x)` with two levels, corresponding to the two groups. The sign of the resulting t-statistic corresponds to "group 1 minus group 2". If `fac` is missing, `rowttests` performs for each row of `x` a two-sided one-class t-test against the null hypothesis 'mean=0'.

`rowttests` and `colttests` are implemented in C and should be reasonably fast and memory-efficient. `fastT` is an alternative implementation, in Fortran, possibly useful for certain legacy code. `rowFtests` and `colFtests` are currently implemented using matrix algebra in R. Compared to the `rowttests` and `colttests` functions, they are slower and use more memory.

**Value**

A data.frame with columns `statistic`, `p.value` (optional in the case of the t-test functions) and `dm`, the difference of the group means (only in the case of the t-test functions). The `row.names` of the data.frame are taken from the corresponding dimension names of `x`. 
The degrees of freedom are provided in the attribute `df`. For the F-tests, if `var.equal` is 'FALSE', `nrow(x)+1` degree of freedoms are given, the first one is the first degree of freedom (it is the same for each row) and the other ones are the second degree of freedom (one for each row).

**Author(s)**

Wolfgang Huber <whuber@embl.de>

**References**


**See Also**

`mt.teststat`

**Examples**

```r
## example data
##
x = matrix(runif(40), nrow=4, ncol=10)
f2 = factor(floor(runif(ncol(x))*2))
f4 = factor(floor(runif(ncol(x))*4))
##
## one- and two group row t-test; 4-group F-test
##
r1 = rowttests(x)
r2 = rowttests(x, f2)
r4 = rowFtests(x, f4)
##
## approximate equality
about.equal = function(x,y,tol=1e-10)
  stopifnot(is.numeric(x), is.numeric(y), length(x)==length(y), all(abs(x-y) < tol))
##
## compare with the implementation in t.test
##
for (j in 1:nrow(x)) {
  s1 = t.test(x[j,])
  about.equal(s1$statistic, r1$statistic[j])
  about.equal(s1$p.value, r1$p.value[j])
  s2 = t.test(x[j,] ~ f2, var.equal=TRUE)
  about.equal(s2$statistic, r2$statistic[j])
  about.equal(s2$p.value, r2$p.value[j])
  dm = -diff(tapply(x[j,], f2, mean))
  about.equal(dm, r2$dm[j])
  s4 = summary(lm(x[j,] ~ f4))
  about.equal(s4$fstatistic"value", r4$statistic[j])
}
```
## colttests

c2 = colttests(t(x), f2)
stopifnot(identical(r2, c2))

## missing values

f2n = f2
f2n[sample(length(f2n), 3)] = NA
r2n = rowttests(x, f2n)
for(j in 1:nrow(x)) {
  s2n = t.test(x[j,] ~ f2n, var.equal=TRUE)
  about.equal(s2n$statistic, r2n$statistic[j])
  about.equal(s2n$p.value, r2n$p.value[j])
}

## larger sample size

x = matrix(runif(1000000), nrow=4, ncol=25000)
f2 = factor(floor(runif(ncol(x))*2))
r2 = rowttests(x, f2)
for (j in 1:nrow(x)) {
  s2 = t.test(x[j,] ~ f2, var.equal=TRUE)
  about.equal(s2$statistic, r2$statistic[j])
  about.equal(s2$p.value, r2$p.value[j])
}

---

**rowpAUCs-methods**  
*Rowwise ROC and pAUC computation*

**Description**

Methods for fast rowwise computation of ROC curves and (partial) area under the curve (pAUC) using the simple classification rule $x > \theta$, where $\theta$ is a value in the range of $x$.

**Usage**

```r
rowpAUCs(x, fac, p=0.1, flip=TRUE, caseNames=c("1", "2"))
```

**Arguments**

- **x**: ExpressionSet or numeric matrix. The matrix must not contain NA values.
- **fac**: A factor or numeric or character that can be coerced to a factor. If `x` is an ExpressionSet, this may also be a character vector of length 1 with the name of a covariate variable in `x`. `fac` must have exactly 2 levels. For better control over the classification, use integer values in 0 and 1, where 1 indicates the "Disease" class in the sense of the Pepe et al paper (see below).
- **p**: Numeric vector of length 1. Limit in (0,1) to integrate pAUC to.
Logical. If TRUE, both classification rules \( x > \theta \) and \( x < \theta \) are tested and the (partial) area under the curve of the better one of the two is returned. This is appropriate for the cases in which the classification is not necessarily linked to higher expression values, but instead it is symmetric and one would assume both over- and under-expressed genes for both classes. You can set flip to FALSE if you only want to screen for genes which discriminate Disease from Control with the \( x > \theta \) rule.

The class names that are used when plotting the data. If \( \text{fac} \) is the name of the covariate variable in the \text{ExpressionSet} the function will use its levels as caseNames.

Details

Rowwise calculation of Receiver Operating Characteristic (ROC) curves and the corresponding partial area under the curve (pAUC) for a given data matrix or \text{ExpressionSet}. The function is implemented in C and thus reasonably fast and memory efficient. Cutpoints (\( \theta \)) are calculated before the first, in between and after the last data value. By default, both classification rules \( x > \theta \) and \( x < \theta \) are tested and the (partial) area under the curve of the better one of the two is returned. This is only valid for symmetric cases, where the classification is independent of the magnitude of \( x \) (e.g., both over- and under-expression of different genes in the same class). For unsymmetric cases in which you expect \( x \) to be consistently higher/lower in of the two classes (e.g., presence or absence of a single biomarker) set flip=FALSE or use the functionality provided in the \text{ROC} package. For better control over the classification (i.e., the choice of "Disease" and "Control" class in the sense of the Pepe et al paper), argument \( \text{fac} \) can be an integer in \([0,1]\) where 1 indicates "Disease" and 0 indicates "Control".

Value

An object of class \text{rowROC} with the calculated specificities and sensitivities for each row and the corresponding pAUCs and AUCs values. See \text{rowROC} for details.

Methods

Methods exist for \text{rowPAUCs}:

\[
\begin{align*}
\text{signature(x="matrix", fac="factor")} \\
\text{rowPAUCs} & \quad \text{signature(x="matrix", fac="numeric")} \\
\text{rowPAUCs} & \quad \text{signature(x="ExpressionSet")} \\
\text{rowPAUCs} & \quad \text{signature(x="ExpressionSet", fac="character")}
\end{align*}
\]

Author(s)

Florian Hahne <fahne@fhcrc.org>

References


See Also

\text{rocdemo.sca}, \text{pAUC}, \text{rowROC}
Examples

library(Biobase)
data(sample.ExpressionSet)

r1 = rowttests(sample.ExpressionSet, "sex")
r2 = rowpAUCs(sample.ExpressionSet, "sex", p=0.1)

plot(area(r2, total=TRUE), r1$statistic, pch=16)

sel <- which(area(r2, total=TRUE) > 0.7)
plot(r2[sel])

## this compares performance and output of rowpAUCs to function pAUC in
## package ROC
if(require(ROC)){
  ## performance
  myRule = function(x)
    pAUC(rocdemo.sca(truth = as.integer(sample.ExpressionSet$sex)-1,
       data = x, rule = dxrule.sca), t0 = 0.1)
  nGenes = 200
  cat("computation time for ", nGenes, " genes:\n")
  cat("function pAUC: ")
  print(system.time(r3 <- esApply(sample.ExpressionSet[1:nGenes, ], 1, myRule)))
  cat("function rowpAUCs: ")
  print(system.time(r2 <- rowpAUCs(sample.ExpressionSet[1:nGenes, ],
       "sex", p=1)))

  ## compare output
  myRule2 = function(x)
    pAUC(rocdemo.sca(truth = as.integer(sample.ExpressionSet$sex)-1,
       data = x, rule = dxrule.sca), t0 = 1)
  r4 <- esApply(sample.ExpressionSet[1:nGenes, ], 1, myRule2)
  plot(r4,area(r2), xlab="function pAUC", ylab="function rowpAUCs",
       main="pAUCs")

  plot(r4, area(rowpAUCs(sample.ExpressionSet[1:nGenes, ],
       "sex", p=1, flip=FALSE)), xlab="function pAUC", ylab="function rowpAUCs",
       main="pAUCs")

  r4[r4<0.5] <- 1-r4[r4<0.5]
  plot(r4, area(r2), xlab="function pAUC", ylab="function rowpAUCs",
       main="pAUCs")
}

rowROC-class

Class “rowROC”

Description

A class to model ROC curves and corresponding area under the curve as produced by rowpAUCs.

Objects from the Class

Objects can be created by calls of the form new("rowROC", ...).
Slots

data: Object of class "matrix" The input data.
ranks: Object of class "matrix" The ranked input data.
sens: Object of class "matrix" Matrix of sensitivity values for each gene at each cutpoint.
spec: Object of class "matrix" Matrix of specificity values for each gene at each cutpoint.
pAUC: Object of class "numeric" The partial area under the curve (integrated from 0 to p).
AUC: Object of class "numeric" The total area under the curve.
factor: Object of class "factor" The factor used for classification.
cutpoints: Object of class "matrix" The values of the cutpoints at which specificity ans sensitivity was calculated. (Note: the data is ranked prior to computation of ROC curves, the cutpoints map to the ranked data.
caseNames: Object of class "character" The names of the two classification cases.
p: Object of class "numeric" The limit to which pAUC is integrated.

Methods

show signature(object="rowROC") Print nice info about the object.
[ signature(x="rowROC", j="missing") Subset the object according to rows/genes.
plot signature(x="rowROC", y="missing") Plot the ROC curve of the first row of the object along with the pAUC. To plot the curve for a specific row/gene subsetting should be done first (i.e. plot(rowROC[1])).
pAUC signature(object="rowROC", p="numeric", flip="logical") Integrate area under the curve from 0 to p. This method returns a new rowROC object.
AUC signature(object="rowROC") Integrate total area under the curve. This method returns a new rowROC object.
sens signature(object="rowROC") Accessor method for sensitivity slot.
spec signature(object="rowROC") Accessor method for specificity slot.
area signature(object="rowROC", total="logical") Accessor method for pAUC slot.

Author(s)

Florian Hahne <fhahne@fhcrc.org>

References


See Also

rowpAUCs
Examples

```r
library(Biobase)
require(genefilter)
data(sample.ExpressionSet)
roc <- rowpAUCs(sample.ExpressionSet, "sex", p=0.5)
roc
area(roc[1:3])
if(interactive()) {
  par(ask=TRUE)
  plot(roc)
  plot(1-spec(roc[1]), sens(roc[2]))
  par(ask=FALSE)
}
pAUC(roc, 0.1)
roc
```

### rowSds

**Row variance and standard deviation of a numeric array**

**Description**

Row variance and standard deviation of a numeric array

**Usage**

```r
rowVars(x, ...)
rowSds(x, ...)
```

**Arguments**

- `x` An array of two or more dimensions, containing numeric, complex, integer or logical values, or a numeric data frame.
- `...` Further arguments that get passed on to `rowMeans` and `rowSums`.

**Details**

These are very simple convenience functions, the main work is done in `rowMeans` and `rowSums`. See the function definition of `rowVars`, it is very simple.

**Value**

A numeric or complex array of suitable size, or a vector if the result is one-dimensional. The ‘dimnames’ (or ‘names’ for a vector result) are taken from the original array.

**Author(s)**

Wolfgang Huber [http://www.ebi.ac.uk/huber](http://www.ebi.ac.uk/huber)

**See Also**

`rowMeans` and `rowSums`
Examples

```r
a = matrix(rnorm(1e4), nrow=10)
rowSds(a)
```

Description

A location estimator based on the shorth

Usage

```r
shorth(x, na.rm=FALSE, tie.action="mean", tie.limit=0.05)
```

Arguments

- `x`: Numeric
- `na.rm`: Logical. If TRUE, then non-finite (according to `is.finite`) values in `x` are ignored. Otherwise, presence of non-finite or NA values will lead to an error message.
- `tie.action`: Character scalar. See details.
- `tie.limit`: Numeric scalar. See details.

Details

The shorth is the shortest interval that covers half of the values in `x`. This function calculates the mean of the `x` values that lie in the shorth. This was proposed by Andrews (1972) as a robust estimator of location.

Ties: if there are multiple shortest intervals, the action specified in `ties.action` is applied. Allowed values are mean (the default), max and min. For mean, the average value is considered; however, an error is generated if the start indices of the different shortest intervals differ by more than the fraction `tie.limit` of `length(x)`. For min and max, the left-most or right-most, respectively, of the multiple shortest intervals is considered.

Rate of convergence: as an estimator of location of a unimodal distribution, under regularity conditions, the quantity computed here has an asymptotic rate of only $n^{-1/3}$ and a complicated limiting distribution.

See `half.range.mode` for an iterative version that refines the estimate iteratively and has a builtin bootstrapping option.

Value

The mean of the `x` values that lie in the shorth.

Author(s)

Wolfgang Huber [http://www.ebi.ac.uk/huber](http://www.ebi.ac.uk/huber), Ligia Pedroso Bras
References

- G Sawitzki, “The Shorth Plot.” Available at http://lshorth.r-forge.r-project.org/TheShorthPlot.pdf

See Also

half.range.mode

Examples

```r
x = c(rnorm(500), runif(500) * 10)
methods = c("mean", "median", "shorth", "half.range.mode")
est = sapply(methods, function(m) get(m)(x))

if(interactive()) {
  colors = 1:4
  hist(x, 40, col="orange")
  abline(v=ests, col=colors, lwd=3, lty=1:2)
  legend(5, 100, names(est), col=colors, lwd=3, lty=1:2)
}
```

Description

The `tdata` data frame has 500 rows and 26 columns. The columns correspond to samples while the rows correspond to genes. The row names are Affymetrix accession numbers.

Usage

data(tdata)

Format

This data frame contains 26 columns.

Source

An unknown data set.

Examples

data(tdata)
ttest  

A filter function for a t.test

Description

ttest returns a function of one argument with bindings for cov and p. The function, when evaluated, performs a t-test using cov as the covariate. It returns TRUE if the p value for a difference in means is less than p.

Usage

ttest(m, p=0.05, na.rm=TRUE)

Arguments

m  If m is of length one then it is assumed that elements one through m of x will be one group. Otherwise m is presumed to be the same length as x and constitutes the groups.

p  The p-value for the test.

na.rm  If set to TRUE any NA’s will be removed.

Details

When the data can be split into two groups (diseased and normal for example) then we often want to select genes on their ability to distinguish those two groups. The t-test is well suited to this and can be used as a filter function.

This helper function creates a t-test (function) for the specified covariate and considers a gene to have passed the filter if the p-value for the gene is less than the prespecified p.

Value

ttest returns a function with bindings for m and p that will perform a t-test.

Author(s)

R. Gentleman

See Also

kOverA, Anova, t.test

Examples

dat <- c(rep(1,5),rep(2,5))
set.seed(5)
y <- rnorm(10)
af <- ttest(dat, .01)
af(y)
af2 <- ttest(5, .01)
af2(y)
y[8] <- NA
af(y)
af2(y)
af(y)
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