

affycoretools

April 19, 2009

affystart

Pre-processing for Affymetrix Data

Description

This function is designed to automatically read in all cel files in a directory, make all pre-processing QC plots and compute expression measures.

Usage

```
affystart(..., filenames=NULL, groups = NULL, groupnames = NULL, plot = TRUE, pca = TRUE, squarepca = FALSE, plottype="pdf", express = c("rma", "mas5", "gcrma"), addname = NULL, output = "txt", annotate = FALSE, ann.vec = c("SYMBOL", "GENENAME", "ENTREZID", "UNIGENE", "REFSEQ"))
```

Arguments

...	Requires that all variables be named.
filenames	If not all cel files in a directory will be used, pass a vector of filenames.
groups	An integer vector indicating the group assignments for the PCA plot.
groupnames	A character vector with group names for PCA legend.
plottype	What type of plot to save. Can be "pdf", "postscript", "png", "jpeg", or "bmp". Defaults to "pdf". Note that "png" and "jpeg" may not be available on a given computer. See the help page for capabilities and png for more information.
plot	Should density and degradation plots be made? Defaults to TRUE.
pca	Should a PCA plot be made? Defaults to TRUE.
squarepca	Should the y-axis of the PCA plot be made comparable to the x-axis? This may aid in interpretation of the PCA plot. Defaults to FALSE.
express	One of either rma, mas5, gcrma. Defaults to rma. Partial matching OK.
addname	Used to append something to the name of the pca plot and the expression values output file (e.g., if function is run twice using different methods to compute expression values).
output	What format to use for the output of expression values. Currently only supports text format.
annotate	Boolean. Add annotation data to the output file?
ann.vec	A character vector of annotation data to add to the output file.

Value

Returns an `ExpressionSet`.

Author(s)

James W. MacDonald <jmacdon@med.umich.edu>

See Also

`plotHist`, `plotDeg`, `plotPCA`

annBM

Select Available Annotation from a Biomart

Description

These functions are designed to do two things that are useful for an end user. If called with no arguments, they will output a character vector of annotation sources that are typically available from a Biomart database. If called with a 'mart' connection (typically created by a call to `useMart`), they will return a character vector of annotation sources that exist for that particular Biomart and species. If called with a 'mart' connection and a character vector of annotation sources, they will return a list that is intended to be used by other functions for creating HTML pages. This last function doesn't have any real utility for the end user.

Usage

```
annBM(mart, annot, species)
linksBM(mart, annot, affyid = FALSE, ann.source = NULL)
```

Arguments

<code>mart</code>	A 'mart' connection, typically created by a call to <code>useMart</code> .
<code>annot</code>	A character vector of annotation sources. This is not typically useful for an end user to specify.
<code>affyid</code>	Boolean. Are the IDs being annotated Affymetrix IDs?
<code>ann.source</code>	Character. The filter name to use if we are annotating Affymetrix IDs.
<code>species</code>	A species name, of the form e.g., 'hsapiens'

Details

The purpose of these functions is to either give an example of typical annotation sources that may be available at a particular Biomart, or to output those sources that are known to exist at a Biomart.

`linksBM` is intended to list those annotation sources that may be turned into hyperlinks whereas `annBM` is intended to list those annotation sources that will not be linked.

These functions have only a few of the possible annotation sources, and currently there is no simple way to extend these sources. Additions to the list are possible, however. Please contact me if there is something in particular that should be included in either list.

Value

Normally called by an end user to output a character vector of annotation sources.

Author(s)

James W. MacDonald <jmacdon@med.umich.edu>

Examples

```
annBM()
```

convert.back

A Function to convert M and A values back to R and G values

Description

This function takes M and A values and converts back to log₂ R and G values.

Usage

```
convert.back(M, A)
```

Arguments

M A vector of log₂ R/G values.
A A vector of log₂ R + G values.

Value

A matrix of log₂ R and G values.

Author(s)

James W. MacDonald <jmacdon@med.umich.edu>

foldFilt

Output Fold Change Data

Description

This function is designed to take an `ExpressionSet` and some comparisons and output either HTML tables, text files, or both.

Usage

```
foldFilt(object, fold = 1, groups, comps, compnames, save = FALSE, text  
= TRUE, html = TRUE, filterfun = NULL)
```

Arguments

object	An <code>ExpressionSet</code> object
fold	The log fold change cutoff to use. Note that this is log base two.
groups	A vector of group identifiers. Probably easiest to use a numeric vector
comps	A list containing all the comparisons to be made. Each list item should be a vector of length two. See details for more information.
compnames	A character vector of the names for each of the comparisons to be made. This will be the name of the resulting HTML or text file.
save	Boolean. If <code>TRUE</code> , a list will be returned. The first item in the list will be a vector showing the number of 'significant' genes for each comparison. The second item will be a matrix of -1's, 0's and 1's indicating a significant difference, and the direction of the difference. The first item is useful for creating Sweave - based reports and the second is useful for making Venn diagrams using the <code>vennDiagram</code> from the <code>limma</code> package.
html	Boolean - if <code>TRUE</code> , output HTML tables
text	Boolean - if <code>TRUE</code> , output text tables
filterfun	A filtering function, created by genefilter to filter the data using additional criteria. See details for more information

Details

This function is useful for outputting annotated gene lists for multiple fold change comparisons. The genes will be ordered by the absolute fold change. Note that this function is essentially a wrapper to call `annaffy`, so is only useful for Affymetrix GeneChips for which there is an annotation package.

Without attaching a data file to this package, it is not possible to give a working example. Instead, here is a 'for instance'.

Say you have an `ExpressionSet` containing four Affy HG-U133Plus2 chips. There is no replication, and you simply want to output genes with a two-fold or greater difference between the first chip and each of the last three (the first chip is the control, and the other three are experimentals). The `ExpressionSet` is called `eset`.

Additionally, say we don't want any genes called significant if both of the samples have very low expression. We can set up a filter using the **genefilter** package.

```
f1 <- kOverA(1,6)
```

```
filt <- filterfun(f1)
```

```
foldFilt(eset, groups=1:4, comps=list(c(2, 1), c(3, 1), c(4, 1)), compnames=c("Expt1-Cont", "Expt2-Cont", "Expt3-Cont"), filterfun = filt)
```

This will output three HTML tables called 'Expt1-Cont.html', etc., each containing sorted genes that have two-fold or greater differences between the two samples.

Value

Returns a list; see above for the elements of the list. This function is mainly called for the side effect of outputting HTML or text files containing annotated 'significant' gene lists.

Author(s)

James W. MacDonald <jmacdon@med.umich.edu>

foldFiltBM

*Output Fold Change Data using biomaRt***Description**

This function is designed to take an `ExpressionSet` and some comparisons and output HTML tables. It is very similar to `foldFilt` except it uses the `biomaRt` package to annotate genes and the `annotate` package to create the HTML table(s).

Usage

```
foldFiltBM(object, fold = 1, groups, comps, compnames, species, links =
linksBM()[1:3], otherann = annBM()[1:3], filterfun = NULL, ann.source =
"entrezgene", affyid = FALSE, mysql = TRUE, html = TRUE, text = TRUE, save = FALSE)
```

Arguments

<code>object</code>	An <code>ExpressionSet</code> object
<code>fold</code>	The log fold change cutoff to use. Note that this is log base two.
<code>groups</code>	A vector of group identifiers. Probably easiest to use a numeric vector
<code>comps</code>	A list containing all the comparisons to be made. Each list item should be a vector of length two. See details for more information.
<code>compnames</code>	A character vector of the names for each of the comparisons to be made. This will be the name of the resulting HTML or text file.
<code>species</code>	The species name. This must be in a particular format for <code>biomaRt</code> . An example for human is "hsapiens" or for mouse is "mmusculus".
<code>links</code>	A character vector of things to annotate with hyperlinks to online databases. See <code>linksBM</code> for possible values.
<code>otherann</code>	A character vector of things to annotate with text only (i.e., no hyperlinks). See <code>annBM</code> for possible values.
<code>filterfun</code>	A filtering function created by <code>genefilter</code> to filter the data using additional criteria. See details for more information
<code>ann.source</code>	The annotation source of the IDs that will be used to annotate the genes. The default value is "entrezgene". See details for other possibilities.
<code>affyid</code>	Boolean. Are the IDs used to annotate these data Affymetrix IDs?
<code>mysql</code>	Boolean. Should <code>biomaRt</code> use <code>RMySQL</code> to connect to the Biomart? Default is <code>TRUE</code> because the <code>RCurl</code> interface of <code>biomaRt</code> is very slow for more than a few genes.
<code>html</code>	Boolean. Output HTML tables? Defaults to <code>TRUE</code>
<code>text</code>	Boolean. Output text tables? Defaults to <code>TRUE</code>
<code>save</code>	Boolean. If <code>TRUE</code> , a list will be returned. The first item in the list will be a vector showing the number of 'significant' genes for each comparison. The second item will be a matrix of -1's, 0's and 1's indicating a significant difference, and the direction of the difference. The first item is useful for creating Sweave - based reports and the second is useful for making Venn diagrams using <code>vennDiagram</code> from the <code>limma</code> package.

Details

This function is useful for outputting annotated gene lists for multiple fold change comparisons. The genes will be ordered by the absolute fold change.

This function currently only supports Affymetrix data. It is designed for Affymetrix chips that don't have an annotation package, which includes data that have been analyzed using the 're-mapped' CDFs supplied to BioC by MBNI at University of Michigan.

The IDs that will be used to annotate the genes depend on the source of the data. If, for example, one is using an Affymetrix chip that doesn't have a BioC annotation package, then the IDs will be Affymetrix IDs. To find out the correct name to use for the `ann.source` argument, one can create a connection to a Biomart database using `useMart` and then get a list of available Affy arrays using `getAffyArrays`.

If one is using one of the re-mapped CDFs from MBNI at University of Michigan, then the IDs to use depend on the mapping used to create the CDF. At this time, only three types of CDFs can be used; EntrezGene, UniGene, and RefSeq. One can determine the correct `ann.source` argument by creating a connection to a Biomart database, and then calling `linksBM(mart, linksBM()[[3]])`.

One can also protect against selecting probesets that have very small expression values for all samples (which likely have a large fold change due to noise, rather than signal) by using the `filterfun` argument. An example would be:

```
f <- kOverA(1, 6)
```

```
filt <- filterfun(f)
```

Then add `filterfun = filt` as an argument to the call to `foldFilt`.

Value

Returns a list; see above for the elements of the list. This function is mainly called for the side effect of outputting HTML or text files containing annotated 'significant' gene lists.

Author(s)

James W. MacDonald <jmacdon@med.umich.edu>

getUniqueLL

Subset a Vector of Probesets

Description

This function will take a vector of Affy IDs and return a vector of Entrez IDs that have replicated IDs removed. The resulting vector will still have the corresponding Affy IDs appended as names, which is important for some functions.

Usage

```
getUniqueLL(probes, annot)
```

Arguments

probes	A vector of probe IDs
annot	The annotation package for the chip used

Details

Subsetting a set of Affy IDs to unique Entrez Gene IDs is a common thing to do prior to doing a hypergeometric test. Functions such as `hyperGtest` can use un-named vectors of Entrez IDs (e.g., `unique(getLL(probeIDs, annot))`), but there is some functionality that requires the Entrez Gene IDs to be in a named vector, with the names being the associated Probeset IDs.

As an example, `hyperGoutput` will only work correctly if the input Entrez ID vector is named with the associated Probeset IDs.

Value

A named vector of unique Entrez IDs

Author(s)

James W. MacDonald <jmacdon@med.umich.edu>

hyperG2annaffy *HTML tables from GOIDs*

Description

Output HTML tables containing the 'enriched' genes for each GO term resulting from a call to `hyperGtable`.

Usage

```
hyperG2annaffy(probinds, lib, eset, fit = NULL, subset = NULL, comp = 1, type = "MF",
pvalue = 0.05, min.count = 10)
```

Arguments

<code>probinds</code>	A vector of Affymetrix probe IDs
<code>lib</code>	An annotation package (e.g., <code>hgu95av2</code>)
<code>eset</code>	An <code>ExpressionSet</code>
<code>fit</code>	An <code>lmFit</code> object. Only necessary if statistics are desired in the resulting table. Defaults to <code>NULL</code> .
<code>subset</code>	A numeric vector used to select GO terms to output (see description for more information). Defaults to <code>NULL</code>
<code>comp</code>	Which contrast/parameter estimate should be used to extract the relevant statistics? Only used if <code>fit</code> is not <code>NULL</code> . See description for more information.
<code>type</code>	One of "MF", "CC", "BP", indicating molecular function, cellular component, or biological process, respectively.
<code>pvalue</code>	The significance level used to choose GO terms
<code>min.count</code>	The minimum number of a given GO term that must be on the chip in order to choose that GO term. This protects against very low p-values that result from the situation where there are very few genes with a given GO term on the chip, but one or two are found in the set of significant genes.

Details

This function is used to create HTML tables based on the output of `hyperGtable`. The basic idea is as follows; as part of an analysis, say `hyperGtable` was used to create a table of 'enriched' GO terms. Unfortunately, the table only lists GO terms and the number of probesets that are annotated to those GO terms, and the client may be interested in knowing what probesets are enriched for each (or some) GO term.

The default behaviour is to output an HTML table for each GO term, containing the probesets that are annotated at that term (and that are in the set of significant genes). If only some of the GO terms are of interest, one may use the `subset` argument to select only particular rows. In addition, if the relevant t-statistics, p-values and fold changes are of interest, one can also use the `fit` argument to point to an `lmFit` object that contains these data, as well as the `comp` argument to indicate which parameter or contrast to use. Note that the `comp` argument defaults to 1, so the first parameter or contrast will be extracted by default.

Value

This function is used only for the side effect of creating HTML tables.

Author(s)

James W. MacDonald <jmacdon@med.umich.edu>

hyperGoutput *Output Tables Based on Hypergeometric Test*

Description

This function will output various tables containing probesets that are annotated to a particular GO, KEGG, or PFAM term. The tables are based on the results from a call to `hyperGtest`.

Usage

```
hyperGoutput(hyptObj, eset, pvalue, categorySize, sigProbesets, fit = NULL,
             subset = NULL, comp = 1, output = c("significant", "all", "split"),
             statistics = c("tstat", "pval", "FC"), html = TRUE, text = TRUE, ...)
```

Arguments

<code>hyptObj</code>	A <code>HyperGResult</code> object, usually produced by a call to <code>hyperGtest</code>
<code>eset</code>	An <code>ExpressionSet</code> object
<code>pvalue</code>	The p-value cutoff used for selecting significant GO terms. If not specified, it will be extracted from the <code>HyperGResult</code> object
<code>categorySize</code>	Number of terms in the universe required for a term to be significant. See details for more information
<code>sigProbesets</code>	Vector of probeset IDs that were significant in the original analysis.
<code>fit</code>	An <code>MArrayLM</code> object, produced from a call to <code>eBayes</code>
<code>subset</code>	Numeric vector used to select particular tables to output. The default is to output tables for all terms. See details for more information

<code>comp</code>	Numeric vector of length one, used to indicate which comparison in the <code>MArrayLM</code> object to use for extracting relevant statistics. See details for more information
<code>output</code>	One of 'selected', 'all', or 'split'. See details for more information
<code>statistics</code>	Which statistics to output in the resulting tables. Choices include 'tstat', 'pval', or 'FC', corresponding to t-statistics, p-values, and fold change, respectively
<code>html</code>	Boolean. Output HTML tables? Defaults to <code>TRUE</code>
<code>text</code>	Boolean. Output text tables? Defaults to <code>TRUE</code>
<code>...</code>	Allows end user to pass further arguments. The most notable would be an <code>anncols</code> argument, passed to <code>probes2table</code> to control the hyperlinked annotation columns. See <code>aaf.handler</code> for more information

Details

This function is designed to be used to output the results from a hypergeometric test for over-represented terms. This function would be used at the end of an analysis such as:

1.) Compute expression values 2.) Fit a model using `limma` 3.) Output significant probesets using `limma2annaffy` 4.) Perform hypergeometric test using `hyperGTest`

At step 4, one can output a list of the over-represented terms using `htmlReport`. One might then be interested in knowing which probesets contributed to the significance of a particular term, which is what this function is designed to do.

One argument that can be passed to `htmlReport` (and also to `hyperGoutput`) is `categorySize`, which gives a lower bound for the number of probesets with a particular term in the universe. In other words, assume that a particular GO term is annotated to three probesets on a given chip. If, after doing a t-test to detect differentially expressed probesets, one of those probesets were found to be significantly differentially expressed and was then used to do a hypergeometric test, that GO term would be significant, with a small p-value. However, this is probably not very strong evidence that the GO term is actually over-represented, since there were only three to begin with. By setting `categorySize` to a sensible value (such as 10), this situation can be avoided.

This function will output HTML and/or text tables containing annotation information about each probeset as well as the expression values. In addition, if `limma` were used to fit the model, the relevant statistics (t-statistic, p-value, fold change) can also be output in the table by passing the `MArrayLM` object that resulted from a call to `eBayes`. The `statistics` argument can be used to control which statistics are output.

By default `hyperGoutput` will output tables for all significant terms, which may end up being quite a few tables. Usually only a few terms are of interest, so there is a `subset` argument that can be used to select only those terms. This argument follows directly from the order of the table output by `htmlReport` or `summary`. For instance, if the first, third and fifth terms in the HTML table output by `htmlReport` were of interest, one would use `subset=c(1,3,5)`.

One critical step prior to the hypergeometric test is to subset the probesets to unique Entrez Gene IDs. It should be noted however, that the functions used by `hyperGoutput` will output all the probesets annotated to a particular term. The `output` argument is used to control this behavior. If `output = "significant"` (the default), then only those probesets that correspond to the original subsetting will be output. If `output = "all"`, then all probesets will be output (grouped by Entrez ID), with the 'significant' probeset first. If `output = "split"`, then all the probesets will be output, with all the 'significant' probesets first, followed by the other probesets, grouped by Entrez ID.

Note that the 'significant' probesets come from one of two sources. First, one can pass a character vector of probeset IDs corresponding to those that were significant in the original analysis (recommended). Second, if the `geneIds` slot of the `GOHyperGParams` object contains a named vector

of Entrez Gene IDs, then the names from that vector will be used. This can be accomplished by using either `findLargest` or `getUniqueLL`.

Since the `geneIds` are by definition a unique set of Entrez Gene IDs, any duplicate probeset IDs will have been removed, so the first method is to be preferred for accuracy.

Value

This function returns no value, and is called solely for the side effect of outputting HTML and/or text tables.

Author(s)

James W. MacDonald <jmacdon@med.umich.edu>

See Also

[hyperGTest](#), [htmlReport](#), [probeSetSummary](#)

limma2annaffy *Function to Create HTML Tables from limma Objects*

Description

This function is designed to take an `ExpressionSet` and an `lmFit`, `model.matrix`, and contrast object from `limma` and convert into HTML tables using `annaffy`. The alternate function `limma2annaffy.na` is designed to be run without user intervention.

Usage

```
limma2annaffy(eset, fit, design, contrast, lib, adjust = "fdr",
  anncols = aaf.handler()[c(1:3, 6:7, 9:12)], number = 30, pfilt = NULL,
  fldfilt = NULL, tstat = TRUE, pval = TRUE, FC = TRUE,
  expression = TRUE, html = TRUE, text = FALSE, save = FALSE, addname =
  NULL, interactive = TRUE)
```

Arguments

<code>eset</code>	An <code>ExpressionSet</code> containing affymetrix expression values.
<code>fit</code>	An <code>lmFit</code> object.
<code>design</code>	A <code>model.matrix</code> object.
<code>contrast</code>	A contrasts matrix from <code>limma</code> .
<code>lib</code>	An annotation package for the Affy chips used.
<code>adjust</code>	Multiplicity adjustment. Choices are "fdr", "holm", "hommel", "bonferroni", or "none". Partial matching allowed.
<code>anncols</code>	A vector of things to annotate, produced by a call to <code>aaf.handler()</code> .
<code>number</code>	Number of genes to output to table. See details for more information.
<code>pfilt</code>	A p-value to filter output. See details for more information.
<code>fldfilt</code>	A fold change to filter output. See details for more information.

tstat	Boolean: Output t-statistics in table? Defaults to FALSE.
pval	Boolean: Output (adjusted) p-values in table? Defaults to FALSE.
FC	Boolean: Output fold changes in table? Defaults to FALSE.
expression	Boolean: Output expression values in table? Defaults to TRUE.
html	Boolean: Output data in HTML tables? Defaults to TRUE.
text	Boolean: Output data in text tables? Defaults to TRUE.
save	Boolean: Save tables as R objects for further processing? Defaults to FALSE.
addname	A character vector to add to the end of the automatically generated output file names. Useful for multiple calls to eliminate over-writing of existing HTML or text tables.
interactive	Boolean: Is this an interactive call, or run as part of a script (e.g., in an Sweave document)? Defaults to TRUE

Details

This function is designed to automatically output HTML or text tables, with filenames taken from the column names of the contrast matrix. The number of genes output can be controlled several different ways. First, if `pfilt` and `fdfilt` are both `NULL`, the top genes will be output based on the `number` variable. Otherwise, the genes are filtered based on p-value, fold change, or both. If the genes are filtered this way, the number of genes to be output will be listed and the filter(s) can then be adjusted if necessary.

This function currently only supports Affymetrix data.

Value

If `save` is `TRUE`, a list of tables from `topTable` will be output.

Author(s)

James W. MacDonald <jmacdon@med.umich.edu>

See Also

[topTable](#), [aafTableAnn](#)

limma2biomaRt	<i>Function to Create HTML Tables from limma Objects using biomaRt for Annotation</i>
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Description

This function is designed to take an `ExpressionSet` and an `lmFit`, `model.matrix`, and contrast object from `limma` and convert into HTML and text tables using `biomaRt`. The alternate function `limma2biomaRt.na` is designed to be run without user intervention.

Usage

```
limma2biomaRt(eset, fit, design, contrast, species, links = linksBM()[1:3],
otherdata = annBM()[1:3], ann.source = "entrezgene", adjust = "fdr",
number = 30, pfilt = NULL, fldfilt = NULL, tstat = TRUE,
pval = TRUE, FC = TRUE, expression = TRUE, html = TRUE, text = TRUE,
save = FALSE, addname = NULL, interactive = TRUE, affyid = FALSE, mysql
= TRUE)
```

Arguments

eset	An ExpressionSet containing affymetrix expression values.
fit	An lmFit object.
design	A model.matrix object.
contrast	A contrasts matrix from limma.
species	The species name. This must be in a particular format for biomaRt. An example for human is "hsapiens", or for mouse "mmusculus".
links	A character vector of things to annotate with hyperlinks to online databases. See linksBM for possible values.
otherdata	A character vector of things to annotate with text only (i.e., no hyperlinks). See annBM for possible values.
ann.source	The annotation source of the IDs that will be used to annotate the genes. The default value is "entrezgene". See details for other possibilities.
adjust	Multiplicity adjustment. Choices are "fdr", "holm", "hommel", "bonferroni", or "none". Partial matching allowed.
number	Number of genes to output to table. See details for more information.
pfilt	A p-value to filter output. See details for more information.
fldfilt	A fold change to filter output. See details for more information.
tstat	Boolean: Output t-statistics in table? Defaults to FALSE.
pval	Boolean: Output (adjusted) p-values in table? Defaults to FALSE.
FC	Boolean: Output fold changes in table? Defaults to FALSE.
expression	Boolean: Output expression values in table? Defaults to TRUE.
html	Boolean: Output data in HTML tables? Defaults to TRUE.
text	Boolean: Output data in text tables? Defaults to TRUE
save	Boolean: Save tables as R objects for further processing? Defaults to FALSE.
addname	A character vector to add to the end of the automatically generated output file names. Useful for multiple calls to eliminate over-writing of existing HTML or text tables.
interactive	Boolean: Is this an interactive call, or run as part of a script (e.g., in an Sweave document)? Defaults to TRUE
affyid	Boolean. Are the IDs used to annotate these data Affymetrix IDs?
mysql	Boolean. Should biomaRt use RMySQL to connect to the Biomart? Default is TRUE because the RCurl interface of biomaRt is very slow for more than a few genes.

Details

This function is designed to automatically output HTML tables, with filenames taken from the column names of the contrast matrix. The number of genes output can be controlled several different ways. First, if `pfilter` and `fdfilter` are both `NULL`, the top genes will be output based on the `number` variable. Otherwise, the genes are filtered based on p-value, fold change, or both. If the genes are filtered this way, the number of genes to be output will be listed and the filter(s) can then be adjusted if necessary.

This function currently only supports Affymetrix data. It is designed for Affymetrix chips that don't have an annotation package, which includes data that have been analyzed using the 're-mapped' CDFs supplied to BioC by MBNI at University of Michigan.

The IDs that will be used to annotate the genes depend on the source of the data. If, for example, one is using an Affymetrix chip that doesn't have a BioC annotation package, then the IDs will be Affymetrix IDs. To find out the correct name to use for the `ann.source` argument, one can create a connection to a Biomart database using [useMart](#) and then get a list of available Affy arrays using [getAffyArrays](#).

If one is using one of the re-mapped CDFs from MBNI at University of Michigan, then the IDs to use depend on the mapping used to create the CDF. At this time, only three types of CDFs can be used; EntrezGene, UniGene, and RefSeq. One can determine the correct `ann.source` argument by creating a connection to a Biomart database, and then calling `linksBM(mart, linksBM())[[3]]`.

Value

If `save` is `TRUE`, a list of tables from [topTable](#) will be output.

Author(s)

James W. MacDonald <jmacdon@med.umich.edu>

See Also

[topTable](#), [aafTableAnn](#)

plotDeg

Functions to Plot Density and RNA Degradation Plots

Description

These functions make density and RNA degradation plots with automatic placement of legends.

Usage

```
plotDeg(dat, filenames = NULL)
plotHist(dat, filenames = NULL)
```

Arguments

<code>dat</code>	An <code>AffyBatch</code> object, or in the case of <code>plotHist</code> , a matrix (e.g., from a call to <code>read.probematrix</code>). Note that <code>plotDeg</code> requires an <code>AffyBatch</code> object to work correctly.
<code>filenames</code>	Filenames that will be used in the legend of the resulting plot. If <code>NULL</code> (the default), these names will be extracted from the <code>sampleNames</code> slot of the <code>AffyBatch</code> object.

Value

These functions are called only for the side effect of making the plots. Nothing else is returned.

Author(s)

James W. MacDonald <jmacdon@med.umich.edu>

Examples

```
library("affydata")
data(Dilution)
plotDeg(Dilution)
plotHist(Dilution)
```

plotma

A Function to Make an MA Plot

Description

This function makes an MA plot from either a matrix or two vectors.

Usage

```
plotma(x, y = NULL, main = "", log = TRUE, out = FALSE)
```

Arguments

x	A matrix of two columns or a vector of numbers. If a matrix with more than two columns is passed, only the first two will be used.
y	A vector of numbers. If NULL, x should be a matrix with two columns.
main	A title for the plot.
log	Boolean describing if the data are log ₂ transformed.
out	Boolean describing if the M and A values should be returned.

Value

If out is TRUE, a matrix of M and A values is returned.

Author(s)

James W. MacDonald <jmacdon@med.umich.edu>

plotPCA

*A Function to Make a PCA Plot from an ExpressionSet***Description**

This function makes a PCA plot from an ExpressionSet or matrix

Usage

```
plotPCA(eset, groups = NULL, groupnames = NULL, addtext = NULL, x.coord = NULL,
y.coord = NULL, screeplot = FALSE, squarepca = FALSE, pch = NULL, col =
NULL, pcs = c(1,2), legend = TRUE, ...)
```

Arguments

eset	An ExpressionSet or matrix.
groups	A numeric vector delineating group membership for samples. Default is NULL, in which case default plotting symbols and colors will be used.
groupnames	A character vector describing the different groups. Default is NULL, in which case the sample names will be used.
addtext	A character vector of additional text to be placed just above the plotting symbol for each sample. This is helpful if there are a lot of samples for identifying e.g., outliers.
x.coord	Pass an x-coordinate if automatic legend placement fails
y.coord	Pass a y-coordinate if automatic legend placement fails.
screeplot	Boolean: Plot a <code>screeplot</code> instead of a PCA plot? Defaults to FALSE.
squarepca	Should the y-axis of the PCA plot be made comparable to the x-axis? This may aid in interpretation of the PCA plot. Defaults to FALSE.
pch	A numeric vector indicating what plotting symbols to use. Default is NULL, in which case default plotting symbols will be used. Note that this argument will override the 'groups' argument.
col	A numeric or character vector indicating what color(s) to use for the plotting symbols. Default is NULL in which case default colors will be used. Note that this argument will override the 'groups' argument.
pcs	A character vector of length two indicating which principal components to plot. Defaults to the first two principal components.
legend	Boolean. Should a legend be added to the plot? Defaults to TRUE.
...	Further arguments to be passed to <code>plot</code> . See the help page for <code>plot</code> for further information.

Value

This function returns nothing. It is called only for the side effect of producing a PCA plot or screeplot.

Author(s)

James W. MacDonald <jmacdon@med.umich.edu>

probes2table	<i>Convert Affy Probe ids to Annotated HTML Table</i>
--------------	---

Description

A function to convert a vector of Affy ids to an annotated HTML table.

Usage

```
probes2table(eset, probids, lib, otherdata = NULL,
             anncols = aaf.handler()[c(1:3, 6:7, 9:12)], html = TRUE, text = FALSE,
             express = TRUE, save = FALSE, filename)
```

Arguments

eset	An ExpressionSet containing Affy expression values.
probids	A vector of probe ids.
lib	An annotation package for the Affy chips used.
otherdata	A *named* list of additional information to include in the resulting table. Examples would be t-statistics, p-values, fold change, etc. Each list item should be a vector the same length as the probids vector. The name associated with each list item will be used as the column name in the resulting table.
anncols	A vector of things to annotate, produced by a call to aaf.handler().
html	Output data in HTML tables? Defaults to TRUE.
text	Output data in text tables? Defaults to TRUE.
express	Output expression values in table? Defaults to TRUE.
save	Should tables be saved as R objects for further processing? Defaults to FALSE.
filename	Filename of the resulting HTML table.

Value

If save is TRUE, a `data.frame` is saved containing the data.

Author(s)

James W. MacDonald <jmacdon@med.umich.edu>

See Also

[topTable](#), [aafTableAnn](#)

probes2tableBM *Convert Affy Probe ids to Annotated HTML Table using biomaRt*

Description

A function to convert a vector of Affy ids to an annotated HTML or text table. This function is very similar to `probes2table`, except it uses the `biomaRt` package to annotate genes, and the `annotate` package to create the HTML table.

Usage

```
probes2tableBM(eset, probids, species, filename, otherdata = NULL,
  links = linksBM()[1:3], otherann = annBM()[1:3], ann.source = "entrezgene",
  express = TRUE, html = TRUE, text = TRUE, affyid = FALSE, mysql = TRUE)
```

Arguments

<code>eset</code>	An <code>ExpressionSet</code> containing Affy expression values.
<code>probids</code>	A vector of probe ids.
<code>species</code>	The species name. This must be in a particular format for <code>biomaRt</code> . An example for human is "hsapiens" or for mouse is "mmusculus".
<code>filename</code>	File name of the resulting HTML table.
<code>otherdata</code>	A *named* list of additional information to include in the resulting table. Examples would be t-statistics, p-values, fold change, etc. Each list item should be a vector the same length as the <code>probids</code> vector. The name associated with each list item will be used as the column name in the resulting table.
<code>links</code>	A character vector of things to annotate with hyperlinks to online databases. See <code>linksBM</code> for possible values.
<code>otherann</code>	A character vector of things to annotate with text only (i.e., no hyperlinks). See <code>annBM</code> for possible values.
<code>ann.source</code>	The annotation source of the IDs that will be used to annotate the genes. The default value is "entrezgene". See details for other possibilities.
<code>express</code>	Output expression values in table? Defaults to <code>TRUE</code> .
<code>html</code>	Boolean. Output HTML table? Defaults to <code>TRUE</code>
<code>text</code>	Boolean. Output text table? Defaults to <code>TRUE</code>
<code>affyid</code>	Boolean. Are the IDs used to annotate these data Affymetrix IDs?
<code>mysql</code>	Boolean. Should <code>biomaRt</code> use <code>RMySQL</code> to connect to the Biomart? Default is <code>TRUE</code> because the <code>RCurl</code> interface of <code>biomaRt</code> is very slow for more than a few genes.

Details

This function is designed to output HTML tables based on a set of IDs. This function currently only supports Affymetrix data. It is designed for Affymetrix chips that don't have an annotation package, which includes data that have been analyzed using the 're-mapped' CDFs supplied to BioC by MBNI at University of Michigan.

The IDs that will be used to annotate the genes depend on the source of the data. If, for example, one is using an Affymetrix chip that doesn't have a BioC annotation package, then the IDs will be Affymetrix IDs. To find out the correct name to use for the `ann.source` argument, one can create a connection to a Biomart database using `useMart` and then get a list of available Affy arrays using `getAffyArrays`.

If one is using one of the re-mapped CDFs from MBNI at University of Michigan, then the IDs to use depend on the mapping used to create the CDF. At this time, only three types of CDFs can be used; EntrezGene, UniGene, and RefSeq. One can determine the correct `ann.source` argument by creating a connection to a Biomart database, and then calling `linksBM(mart, linksBM()[[3]])`.

Value

This function is only used for the side effect of outputting an HTML table.

Author(s)

James W. MacDonald <jmacdon@med.umich.edu>

See Also

`topTable`

vennCounts2

Compute Counts for Venn Diagram

Description

This function is designed to compute counts for a Venn diagram. It is slightly different from `vennCounts` in the additional ability to compute counts for genes that are differentially expressed in the same direction.

Usage

```
vennCounts2(x, method = "same", fit = NULL, foldFilt = NULL)
```

Arguments

<code>x</code>	A <code>TestResults</code> object, produced by a call to <code>decideTests</code> or <code>foldFilt</code> .
<code>method</code>	One of "same", "both", "up", "down". See details for more information.
<code>fit</code>	An <code>MArrayLM</code> object, produced by a call to <code>lmFit</code> and <code>eBayes</code> . Only necessary if <code>'foldFilt' = TRUE</code> .
<code>foldFilt</code>	A fold change to filter samples. This is primarily here for consistency with the corresponding argument in <code>vennSelect</code> .

Details

The function `vennCounts` will return identical results except for the "same" method. This will only select those genes that both pass the criteria of `decideTests` as well as being differentially expressed in the same direction. Note that this is different from the "both" method, which simply requires that a given gene be differentially expressed in e.g., two different comparisons without any requirement that the direction be the same.

Value

A [VennCounts](#) object.

Author(s)

James W. MacDonald <jmacdon@med.umich.edu>

Examples

```
tstat <- matrix(rt(300,df=10),100,3)
tstat[1:33,] <- tstat[1:33,]+2
clas <- classifyTestsF(tstat,df=10,p.value=0.05)
a <- vennCounts2(clas)
print(a)
vennDiagram(a)
```

 vennSelect

Select and Output Genelists Based on Venn Diagrams

Description

This function is designed to output text and/or HTML tables based on the results of a call to [decideTests](#).

Usage

```
vennSelect(eset, design, x, contrast, fit, method = "same", adj.meth =
"BH", stat = "fstat", otherstats = c("pval", "FC"), order.by = "pval",
foldFilt = NULL, save = FALSE, ...)
```

Arguments

eset	A ExpressionSet object.
design	A design matrix, usually from a call to <code>model.matrix</code> . See details for more information.
x	A TestResults object, usually from a call to decideTests .
contrast	A contrasts matrix, produced either by hand, or by a call to makeContrasts
fit	An MArrayLM object, from a call to eBayes .
method	One of "same", "both", "up", "down", "sameup", or "samedown". See details for more information.
adj.meth	Method to use for adjusting p-values. Default is 'BH', which corresponds to 'fdr'. Ideally one would set this value to be the same as was used for decideTests .
stat	The statistic to report in the resulting HTML tables. Choices are 'fstat', 'tstat', and NULL. Ideally, the statistic chosen would correspond to the method used in decideTests . In other words, if one used methods such as 'separate' or 'hierarchical', which are based on a t-statistic, one should choose 'tstat', however, if one used 'nestedF', the logical choice would be 'fstat'.
otherstats	Other statistics to be included in the HTML tables. Choices include 'pval' and 'FC'.

<code>order.by</code>	Which statistic should be used to order the probesets? Choices include 'fstat', 'tstat', 'pval', and 'FC'. Note that if 'FC' is chosen and there are more than one set of fold changes, the first will be used.
<code>foldFilt</code>	A fold change to use for filtering. Default is <code>NULL</code> , meaning no filtering will be done.
<code>save</code>	Boolean - If <code>TRUE</code> , output a count of genes that fulfill the criteria. Useful for e.g., Sweave-type reports.
<code>...</code>	Used to pass other arguments to <code>probes2table</code> , in particular, to change the argument to <code>anncols</code> which controls the columns of hyperlinks to online databases (e.g., Entrez Gene, etc.). See aaf.handler for more information.

Details

The purpose of this function is to output HTML and text tables with lists of genes that fulfill the criteria of a call to `decideTests` as well as the direction of differential expression.

Some important things to note: First, the names of the HTML and text tables are extracted from the `colnames` of the `TestResults` object, which come from the contrasts matrix, so it is important to use something descriptive. Second, the method argument is analogous to the `include` argument from `vennCounts` or `vennDiagram`. Choosing "both" will select genes that are differentially expressed in one or more comparisons, regardless of direction. Choosing "up" or "down" will select genes that are only differentially expressed in one direction. Choosing "same" will select genes that are differentially expressed in the same direction. Choosing "sameup" or "samedown" will select genes that are differentially expressed in the same direction as well as 'up' or 'down'.

Note that this is different than sequentially choosing "up" and then "down". For instance, a gene that is upregulated in one comparison and downregulated in another comparison will be listed in the intersection of those two comparisons if "both" is chosen, it will be listed in only one comparison for both the "up" and "down" methods, and it will be listed in the union (e.g., not selected) if "same" is chosen.

Calling the function normally will result in the output of HTML and text tables:

```
vennSelect(eset, fit, design, x)
```

Calling the function with `save` set to `TRUE` will output both HTML and text tables as well as a vector of counts for each comparison. This is useful when using the function programmatically (e.g., when making reports using Sweave).

```
out <- vennSelect(eset, fit, design, x, save = TRUE)
```

An alternative would be to use `vennCounts2` and `vennDiagram` to output a Venn diagram, which is probably more reasonable since the tables being output are supposed to be based on a Venn diagram.

Value

Normally called only for the side effect of producing HTML and text tables. However, setting `save` to `TRUE` will output a vector of counts that can be used for making Sweave-style reports.

Author(s)

James W. MacDonald <jmacdon@med.umich.edu>

Description

This function is designed to output HTML and text tables based on the results of a call to `decideTests`. This function is very similar to `vennSelect`, except it uses the `biomaRt` package to annotate genes, and the `annotate` package to create the HTML table.

Usage

```
vennSelectBM(eset, design, x, contrast, fit, method = "same", adj.meth = "BH",
stat = "fstat", otherstats = c("pval", "FC"), order.by = "pval",
foldFilt = NULL, save = FALSE, species, links = linksBM()[1:3],
otherdata = annBM()[1:3], mysql = TRUE, ann.source = "entrezgene",
html = TRUE, text = TRUE, affyid = FALSE, ...)
```

Arguments

<code>eset</code>	A <code>ExpressionSet</code> object.
<code>design</code>	A design matrix, usually from a call to <code>model.matrix</code> . See details for more information.
<code>x</code>	A <code>TestResults</code> object, usually from a call to <code>decideTests</code> .
<code>contrast</code>	A contrasts matrix, produced either by hand, or by a call to <code>makeContrasts</code>
<code>fit</code>	An <code>MArrayLM</code> object, from a call to <code>eBayes</code> .
<code>method</code>	One of "same", "both", "up", "down", "sameup", or "samedown". See details for more information.
<code>adj.meth</code>	Method to use for adjusting p-values. Default is 'BH', which corresponds to 'fdr'. Ideally one would set this value to be the same as was used for <code>decideTests</code> .
<code>stat</code>	The statistic to report in the resulting HTML tables. Choices are 'fstat', 'tstat', and NULL. Ideally, the statistic chosen would correspond to the method used in <code>decideTests</code> . In other words, if one used methods such as 'separate' or 'hierarchical', which are based on a t-statistic, one should choose 'tstat', however, if one used 'nestedF', the logical choice would be 'fstat'.
<code>otherstats</code>	Other statistics to be included in the HTML tables. Choices include 'pval' and 'FC'.
<code>order.by</code>	Which statistic should be used to order the probesets? Choices include 'fstat', 'tstat', 'pval', and 'FC'. Note that if 'FC' is chosen and there are more than one set of fold changes, the first will be used.
<code>foldFilt</code>	A fold change to use for filtering. Default is NULL, meaning no filtering will be done.
<code>save</code>	Boolean - If TRUE, output a count of genes that fulfill the criteria. Useful for e.g., Sweave-type reports.
<code>species</code>	The species name. This must be in a particular format for <code>biomaRt</code> . An example for human is "hsapiens", or for mouse "mmusculus".
<code>links</code>	A character vector of things to annotate with hyperlinks to online databases. See <code>linksBM</code> for possible values.

<code>otherdata</code>	A character vector of things to annotate with text only (i.e., no hyperlinks). See <code>annBM</code> for possible values.
<code>mysql</code>	Boolean. If <code>TRUE</code> , then <code>biomaRt</code> will use the <code>RMySQL</code> package to connect to the Biomart server (note that this requires <code>RMySQL</code> to be installed). If <code>FALSE</code> , <code>biomaRt</code> will use the <code>RCurl</code> interface. WARNING: if you are planning to annotate a large number of genes, you will almost certainly need to use the <code>RMySQL</code> interface.
<code>ann.source</code>	The annotation source of the IDs that will be used to annotate the genes. The default value is "entrezgene". See details for other possibilities.
<code>html</code>	Boolean. Output HTML tables? Defaults to <code>TRUE</code>
<code>text</code>	Boolean. Output text tables? Defaultst to <code>TRUE</code>
<code>affyid</code>	Boolean. Are the IDs used to annotate these data Affymetrix IDs?
<code>...</code>	Used to pass other variables to e.g., <code>htmlpage</code> .

Details

The purpose of this function is to output HTML tables with lists of genes that fulfill the criteria of a call to `decideTests` as well as the direction of differential expression.

The IDs that will be used to annotate the genes depend on the source of the data. If, for example, one is using an Affymetrix chip that doesn't have a BioC annotation package, then the IDs will be Affymetrix IDs. To find out the correct name to use for the `ann.source` argument, one can create a connection to a Biomart database using `useMart` and then deduce the correct argument by the output from `listFilters(mart)`. It will usually be something starting with 'affy', and contain the name of the chip.

If one is using one of the re-mapped CDFs from MBNI at University of Michigan, then the IDs to use depend on the mapping used to create the CDF. At this time, only three types of CDFs can be used; EntrezGene, UniGene, and RefSeq. One can determine the correct `ann.source` argument by creating a connection to a Biomart database, and then calling `linksBM(mart, linksBM())[[3]]`.

Some important things to note: First, the names of the HTML tables are extracted from the `colnames` of the `TestResults` object, which come from the contrasts matrix, so it is important to use something descriptive. Second, the method argument is analogous to the `include` argument from `vennCounts` or `vennDiagram`. Choosing "both" will select genes that are differentially expressed in one or more comparisons, regardless of direction. Choosing "up" or "down" will select genes that are only differentially expressed in one direction. Choosing "same" will select genes that are differentially expressed in the same direction. Choosing "sameup" or "samedown" will select genes that are differentially expressed in the same direction as well as 'up' or 'down'.

Note that this is different than sequentially choosing "up" and then "down". For instance, a gene that is upregulated in one comparison and downregulated in another comparison will be listed in the intersection of those two comparisons if "both" is chosen, it will be listed in only one comparison for both the "up" and "down" methods, and it will be listed in the union (e.g., not selected) if "same" is chosen.

Calling the function normally will result in the output of HTML tables.

Calling the function with `save` set to `TRUE` will output HTML tables as well as a vector of counts for each comparison. This is useful when using the function programmatically (e.g., when making reports using Sweave).

```
out <- vennSelectBM(eset, fit, design, x, <other arguments>, save = TRUE)
```

An alternative would be to use `vennCounts2` and `vennDiagram` to output a Venn diagram, which is probably more reasonable since the tables being output are supposed to be based on a Venn diagram.

Value

Normally called only for the side effect of producing HTML tables. However, setting `save` to `TRUE` will output a vector of counts that can be used for making Sweave-style reports.

Author(s)

James W. MacDonald <jmacdon@med.umich.edu>

 vennSelectFC

Select and Output Gene Lists Based on Venn Diagrams

Description

This function is designed to output text and/or HTML tables based on the results of a call to `foldFilt`. The general idea being that one might want to create a Venn diagram showing probe-sets that are unique to particular comparisons, or consistent between comparisons, and then might want to output the probesets that are contained in each cell of the Venn diagram.

Usage

```
vennSelectFC(eset, x, comps, order.by = "sum", method =
"same", text = TRUE, html = TRUE, ...)
```

Arguments

<code>eset</code>	A ExpressionSet object.
<code>x</code>	An indicator matrix showing up or down regulation based on fold change, usually from a call to <code>foldFilt</code> . See details for more information.
<code>comps</code>	A list containing all the comparisons to be made. Each list item should be a vector of length two. This should be identical to the 'comps' argument used in the call to <code>foldFilt</code> . See details for more information.
<code>order.by</code>	One of 'sum', 'max', 'median', or 'mean'. This orders the output for those tables that have multiple fold change values based on the summary statistic chosen. Defaults to 'sum'.
<code>method</code>	One of "same", "both", "up", "down", "sameup", or "samedown". See details for more information.
<code>text</code>	Boolean. Output text tables? Defaults to <code>TRUE</code>
<code>html</code>	Boolean. Output HTML tables? Defaults to <code>TRUE</code>
<code>...</code>	Used to pass other arguments to <code>probes2table</code> , in particular, to change the argument to <code>anncols</code> which controls the columns of hyperlinks to online databases (e.g., Entrez Gene, etc.). See aaf.handler for more information.

Details

The purpose of this function is to output the probesets listed in a Venn diagram that has been produced by a call to `foldFilt`. A small example would be as follows:

Assume an `ExpressionSet` exists that contains expression values for three Affymetrix chips, say a control, and two experimentals. One might want to know what probesets are different between each of the experimentals and the control, and those that are different between both of the experimentals and the control. We first make the comparisons, based on a fold change of 2 (or a difference of 1 on the log scale).

```
comps <- list(c(1,2), c(1,3))
```

This list indicates what comparisons we want. In this case 1vs2 and 1vs3.

```
out <- foldFilt(eset, fold = 1, groups = 1:3, comps = comps, compnames=c("Control vs experimen-  
tal1", "Control vs experimental2"), save = TRUE)
```

By setting `save = TRUE`, we are saving a list, the first item being a vector of the number of probesets in each comparison, the second item being an indicator matrix showing up or down regulation based on a two-fold difference. We could make a Venn diagram using this matrix with `vennCounts2` and `vennDiagram`. If we then wanted to output the probesets in each cell of that Venn diagram, we could use `vennSelectFC` as follows:

```
vennSelectFC(eset, out[[2]], comps)
```

One thing to note here is that the names of the resulting tables as well as the columns containing the fold change values will be extracted from the column names of the indicator matrix. This matrix will get its column names from the `'compnames'` argument to `foldFilt`, so it is best to use reasonable names here. Also note that any character used in the `'compnames'` argument that is not a valid character for a file name will be stripped out.

Value

Called only for the side effect of outputting HTML and/or text tables.

Author(s)

James W. MacDonald <jmacdon@med.umich.edu>

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