# Package 'PAA'

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Title PAA (Protein Array Analyzer)

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LinkingTo Rcpp

Suggests BiocStyle, RUnit, BiocGenerics, vsn

Description PAA imports single color (protein) microarray data that has been saved in gpr file format - esp. ProtoArray data. After preprocessing (background correction, batch filtering, normalization) univariate feature preselection is performed (e.g., using the ``minimum M statistic" approach - hereinafter referred to as ``mMs"). Subsequently, a multivariate feature selection is conducted to discover biomarker candidates. Therefore, either a frequency-based backwards elimination aproach or ensemble feature selection can be used. PAA provides a complete toolbox of analysis tools including several different plots for results examination and evaluation.

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2 batchAdjust

# **Contents**

	oatchAdjust	2
	patchFilter	3
	oatchFilter.anova	4
	liffAnalysis	5
	oadGPR	7
	nMsMatrix	9
	normalizeArrays	C
	olotArray	2
	olotFeatures	4
	olotFeaturesHeatmap	6
	olotFeaturesHeatmap.2	7
	olotMAPlots	8
	olotNormMethods	C
	preselect	1
	printFeatures	3
	ovaluePlot	4
	electFeatures	
	huffleData	9
	volcanoPlot	C
Index		1
ınaex	3	12
hatck	djust Adjust microarray data for batch effects.	
batti	ajust merourray and for batch effects.	

# Description

Adjusts EListRaw or EList data for batch/lot effects.

# Usage

batchAdjust(elist=NULL, log=NULL)

## **Arguments**

elist EList or EListRaw object containing the data to be adjusted (mandatory).

log logical indicating whether the data is in log scale (mandatory; note: if TRUE

log2 scale is expected).

# **Details**

This is a wrapper to sva's function ComBat() for batch adjustment using the empirical Bayes approach. To use batchAdjust the targets information of the EList or EListRaw object must contain the columns "Batch" (containing batch/lot information for each particular array) and "Group" (containing experimental group information for each particular array).

batchFilter 3

### Value

An EListRaw or EList object with the adjusted data in log scale is returned.

#### Note

The targets information of the EListRaw or EList object must contain the columns "Batch" and "Group".

#### Author(s)

Michael Turewicz, <michael.turewicz@rub.de>

### References

The package sva by Jeffrey T. Leek et al. can be downloaded from Bioconductor (http://www.bioconductor.org/).

Johnson WE, Li C, and Rabinovic A (2007) Adjusting batch effects in microarray expression data using empirical Bayes methods. Biostatistics 8:118-27.

## **Examples**

```
cwd <- system.file(package="PAA")
load(paste(cwd, "/extdata/Alzheimer.RData", sep=""))
elist <- elist[elist$genes$Block < 10,]
elist <- batchAdjust(elist=elist, log=FALSE)</pre>
```

batchFilter

Remove differential features regarding array batches/lots.

# Description

Finds differential features regarding array batches/lots and removes them.

## Usage

```
batchFilter(elist=NULL, lot1=NULL, lot2=NULL, log=NULL, p.thresh=0.05,
fold.thresh=1.5, output.path=NULL)
```

## **Arguments**

elist	EList or EListRaw object (mandatory).
lot1	vector of column names for group 1 (mandatory).
lot2	vector of column names for group 2 (mandatory).
log	logical indicating whether the data is in log scale (mandatory; note: if TRUE log2 scale is expected).

4 batchFilter.anova

p.thresh	positive float number between 0 and 1 indicating the maximum Student's t-test p-value for features to be considered as differential (e.g., "0.5").
fold.thresh	float number indicating the minimum fold change for features to be considered as differential (e.g., "1.5").
output.path	string indicating a path for saving results (optional).

#### **Details**

This function takes an EList or EListRaw object (see limma documentation) and the batch-specific column name vectors lot1 and lot2 to find differential features regarding batches/lots. For this purpose, thresholds for p-values (Student's t-test) and fold changes can be defined. To visualize the differential features a volcano plot is drawn. Then, differential features are removed and the remaining data are returned. When an output path is defined (via output.path) volcano plots and result files are saved on the hard disk.

#### Value

An EList or EListRaw object without differential features regarding array batches/lots.

### Author(s)

Michael Turewicz, <michael.turewicz@rub.de>

## **Examples**

```
cwd <- system.file(package="PAA")
load(paste(cwd, "/extdata/Alzheimer.RData", sep=""))
elist <- elist[elist$genes$Block < 10,]
lot1 <- elist$targets[elist$targets$Batch=='Batch1','ArrayID']
lot2 <- elist$targets[elist$targets$Batch=='Batch2','ArrayID']
elist <- batchFilter(elist=elist, lot1=lot1, lot2=lot2, log=FALSE, p.thresh=0.001, fold.thresh=3)</pre>
```

batchFilter.anova Remove features which are differential regarding microarray batches / lots in a multi-batch scenario.

## Description

Finds features which are differential regarding at least two microarray batches / lots in a multi-batch scenario (i.e., > 2 batches) via one-way analysis of variance (ANOVA) and removes them.

#### Usage

```
batchFilter.anova(elist=NULL, log=NULL, p.thresh=0.05, fold.thresh=1.5,
output.path=NULL)
```

diffAnalysis 5

### **Arguments**

elist	EList or EListRaw object (mandatory).
log	logical indicating whether the data is in log scale (mandatory; note: if TRUE log2 scale is expected).
p.thresh	positive float number between 0 and 1 indicating the maximum Student's t-test p-value for features to be considered as differential (e.g., "0.5").
fold.thresh	float number indicating the minimum fold change for features to be considered as differential (e.g., "1.5").
output.path	string indicating a path for saving results (optional).

#### **Details**

This function takes an EList or EListRaw object (see limma documentation) to find features which are differential regarding at least two microarray batches / lots in a multi-batch scenario (i.e., more than two batches). For this purpose, thresholds for p-values obtained from an one-way analysis of variance (ANOVA) and fold changes can be defined. To visualize the differential features a volcano plot is drawn. Then, differential features are removed and the remaining data are returned. When an output path is defined (via output.path) volcano plots and result files are saved on the hard disk.

## Value

An EList or EListRaw object without differential features regarding at least two microarray batches / lots.

### Author(s)

Ivan Grishagin (Rancho BioSciences LLC, San Diego, CA, USA), John Obenauer (Rancho Bio-Sciences LLC, San Diego, CA, USA) and Michael Turewicz (Ruhr-University Bochum, Bochum, Germany), <michael.turewicz@rub.de>

#### **Examples**

```
cwd <- system.file(package="PAA")
load(paste(cwd, "/extdata/Alzheimer.RData", sep=""))
elist <- elist[elist$genes$Block < 10,]
elist <- batchFilter.anova(elist=elist, log=FALSE, p.thresh=0.001,
    fold.thresh=3)</pre>
```

diffAnalysis Differential analysis.

# **Description**

Performs a univariate differential analysis.

6 diffAnalysis

## Usage

```
diffAnalysis(input=NULL, label1=NULL, label2=NULL, class1=NULL, class2=NULL,
output.path=NULL, mMs.matrix1=NULL, mMs.matrix2=NULL, above=1500,
between=400, features=NULL, feature.names=NULL)
```

# Arguments

input	EList\$E- or EListRaw\$E-matrix extended by row names comprising BRC-IDs of the corresponding features (mandatory; note: it is expected that this matrix is in original scale and not in log2 scale).
label1	vector of column names for group 1 (mandatory).
label2	vector of column names for group 2 (mandatory).
class1	label of group 1 (mandatory).
class2	label of group 2 (mandatory).
output.path	string indicating a path for saving the results (optionally).
mMs.matrix1	precomputed mMs reference matrix (see ${\tt mMsMatrix()})$ for group 1 (mandatory).
mMs.matrix2	precomputed mMs reference matrix (see ${\tt mMsMatrix()})$ for group 2 (mandatory).
above	mMs above parameter (integer). Default is "1500".
between	mMs between parameter (integer). Default is "400".
features	vector of row indices (optional).
feature.names	vector of corresponding feature names (additionally to features).

#### **Details**

This function takes an EList\$E- or EListRaw\$E-matrix (e.g., temp <- elist\$E) extended by row names comprising BRC-IDs of the corresponding features. The BRC-IDs can be created via: brc <- paste(elist\$genes[,1], elist\$genes[,3], elist.\$genes[,2]).

The BRC-row names can be defined as follows: rownames(temp) <- brc. Furthermore, the corresponding column name vectors, group labels and mMs-parameters are needed to perform the univariate differential analysis. This analysis covers inter alia p-value computation, p-value adjustment (method: Benjamini & Hochberg, 1995), and fold change computation. Since the results table is usually large, a path for saving the results can be defined via output.path. Optionally, a vector of row indices (features) and additionally (not mandatory for subset analysis) a vector of corresponding feature names (feature.names) can be forwarded to perform the analysis for a feature subset.

## Value

A matrix containing the analysis results is returned.

## Author(s)

Michael Turewicz, <michael.turewicz@rub.de>

IoadGPR 7

## **Examples**

```
cwd <- system.file(package="PAA")
load(paste(cwd, "/extdata/Alzheimer.RData", sep=""))
elist <- elist[elist$genes$Block < 10,]
c1 <- paste(rep("AD",20), 1:20, sep="")
c2 <- paste(rep("NDC",20), 1:20, sep="")
mMs.matrix1 <- mMs.matrix2 <- mMsMatrix(x=20, y=20)
temp <- elist$E
rownames(temp) <- paste(elist$genes[,1], elist$genes[,3], elist$genes[,2])
diffAnalysis(input=temp, label1=c1, label2=c2, class1="AD", class2="NDC",
mMs.matrix1=mMs.matrix1, mMs.matrix2=mMs.matrix2, above=1500,
between=400)</pre>
```

loadGPR

Importing raw data from gpr files.

### **Description**

Constructs an EListRaw object from a set of gpr files containing ProtoArray data or other protein microarray data.

## Usage

```
loadGPR(gpr.path = NULL, targets.path = NULL, array.type = NULL,
aggregation = "none", array.columns = list(E = "F635 Median",
Eb = "B635 Median"),
array.annotation = c("Block", "Column", "Row", "Description", "Name", "ID"),
description = NULL, description.features = NULL, description.discard = NULL)
```

#### **Arguments**

gpr.path	string indicating the path to a folder containing gpr files (mandatory).
targets.path	string indicating the path to targets file (see limma, mandatory).
array.type	string indicating the microarray type of the imported gpr files. Only for ProtoArrays duplicate aggregation will be performed. The possible options are: "ProtoArray", "HuProt" and "other" (mandatory).
aggregation	string indicating which type of ProtoArray spot duplicate aggregation should be performed. If "min" is chosen, the value for the corresponding feature will be the minimum of both duplicate values. If "mean" is chosen, the arithmetic mean will be computed. Alternatively, no aggregation will be performed, if "none" is chosen. The default is "min" (optional).
array.columns	list containing the column names for foreground intensities $(E)$ and background intensities $(Eb)$ in the gpr files that is passed to limma's "read.maimages" function (optional).
array.annotatio	n

rray.annotation

string vector containing further mandatory column names that are passed to limma (optional).

8 loadGPR

description

string indicating the column name of an alternative column containing the information which spot is a feature, control or to be discarded for gpr files not providing the column "Description" (optional).

description.features

string containing a regular expression identifying feature spots. Mandatory when description has been defined.

description.discard

string containing a regular expression identifying spots to be discarded (e.g., empty spots). Mandatory when description has been defined.

#### **Details**

This function is partially a wrapper to limma's function read.maimages() featuring optional duplicate aggregation for ProtoArray data. Paths to a targets file and to a folder containing gpr files (all gpr files in that folder that are listed in the targets file will be read) are mandatory. The folder "R\_HOME/library/PAA/extdata" contains an exemplary targets file that can be used as a template. If array.type (also mandatory) is set to "ProtoArray", duplicate spots can be aggregated. The corresponding method ("min", "mean" or "none") can be specified via the argument aggregation. As another ProtoArray-specific feature, control spot data and information will be stored in additional components of the returned object (see below). Arguments array.columns and array.annotation define the columns where read.maimages() will find foreground and background intensity values as well as other important columns. For array.annotation the default columns "Block", "Column", "Row", "Description", "Name" and "ID" are mandatory.

If the column "Description" is not provided by the gpr files for ProtoArrays a makeshift column will be constructed from the column "Name" automatically. For other microarrays the arguments description, description.features and description.discard can be used to provide the mandatory information (see the example below).

#### Value

An extended object of class EListRaw (see the documentation of limma for details) is returned. If array.type is set to "ProtoArray" (default), the object provides additional components for control spot data: C, Cb and cgenes which are analogous to the probe spot data E, Eb and genes. Moreover, the returned object always provides the additional component array.type indicating the type of the imported protein microarray data (e.g., "ProtoArray").

#### Note

Don't forget to check column names in your gpr files. They may differ from the default settings of loadGPR() and should be renamed to the default column names (see also the exemplary gpr files accompanying PAA as a reference for the default column names). At worst, important columns in your gpr files may be completely missing and should be added in order to provide all information needed by PAA.

Note that if array.type is not "ProtoArray", neither aggregation will be done nor controls components will be added to the returned object of class EListRaw.

# Author(s)

Michael Turewicz, <michael.turewicz@rub.de>

mMsMatrix 9

#### References

The package limma by Gordon Smyth et al. can be downloaded from Bioconductor (http://www.bioconductor.org/).

Smyth, G. K. (2005). Limma: linear models for microarray data. In: Bioinformatics and Computational Biology Solutions using R and Bioconductor, R. Gentleman, V. Carey, S. Dudoit, R. Irizarry, W. Huber (eds.), Springer, New York, pages 397-420.

## **Examples**

```
gpr <- system.file("extdata", package="PAA")
targets <- list.files(system.file("extdata", package="PAA"),
  pattern = "dummy_targets", full.names=TRUE)
elist <- loadGPR(gpr.path=gpr, targets.path=targets, array.type="ProtoArray")

# Example showing how to use the arguments description, description.features and
# description.discard in order to construct a makeshift column 'Description'
# for gpr files without this column. Please see also the exemplary gpr files
# coming with PAA.
targets2 <- list.files(system.file("extdata", package="PAA"),
  pattern = "dummy_no_descr_targets", full.names=TRUE)
elist2 <- loadGPR(gpr.path=gpr, targets.path=targets2, array.type="other",
  description="Name", description.features="^Hs~", description.discard="Empty")</pre>
```

mMsMatrix

Compute a reference minimum M statistic (n1 x n2)-matrix.

## **Description**

Computes a reference minimum M statistic (n1 x n2)-matrix (mMs matrix).

# Usage

```
mMsMatrix(x, y)
```

#### **Arguments**

- x integer, first dimension (i.e., number of samples in group 1) of the mMs matrix to be computed (mandatory).
- y integer, second dimension (i.e., number of samples in group 2) of the mMs matrix to be computed (mandatory).

### **Details**

For feature preselection the "minimum M Statistic" (mMs) proposed by Love B. can be used. The mMs is a univariate measure that is sensitive to population subgroups. To avoid redundant mMs computations for a large number of features (e.g., ca. 9500 features on ProtoArray v5) a reference matrix containing all relevant mMs values can be precomputed. For this purpose, only two parameters are needed: the number of samples in group 1 (n1) and the number of samples in group 2 (n2).

10 normalizeArrays

According to mMs definition for each matrix element (i,m) a mMs value (= the probability of) for having m values in group 1 larger than the i-th largest value in group 2 is computed.

#### Value

A (n1 x n2)-matrix containing all mMs values for group 1 and group 2.

#### Note

To check whether a feature is more prevalent in group 1 or in group 2, PAA needs both the mMs for having m values in group 1 larger than the i-th largest element in group 2 as well as the mMs for having m values in group 2 larger than the i-th largest element in group 1. Hence, always both must be computed: mMsMatrix(n1,n2) and mMsMatrix(n2,n1).

#### Author(s)

Michael Turewicz, <michael.turewicz@rub.de>

#### References

Love B: The Analysis of Protein Arrays. In: Functional Protein Microarrays in Drug Discovery. CRC Press; 2007: 381-402.

# **Examples**

```
#exemplary computation for a group 1 comprising 10 arrays and a group 2
#comprising 12 arrays
mMs.matrix1 <- mMsMatrix(x=10, y=12)
mMs.matrix2 <- mMsMatrix(x=12, y=10)</pre>
```

normalizeArrays

Normalize microarray data.

# **Description**

Normalizes EListRaw data and returns an EList object containing normalized data in log2 scale.

# Usage

```
normalizeArrays(elist = NULL, method = "quantile", cyclicloess.method = "pairs",
controls="internal", group1 = NULL, group2 = NULL, output.path=NULL)
```

normalizeArrays 11

#### **Arguments**

elist EListRaw object containing raw data to be normalized (mandatory).

method string indicating the normalization method ("cyclicloess", "quantile", "vsn"

or "rlm") to be used (mandatory).

cyclicloess.method

string indicating which type of cyclicloess normalization ("pairs", "fast",

"affy") should be performed (optional).

controls sring indicating the ProtoArray controls for rlm normalization (optional). Valid

options are "internal" (default), "external", "both" or a regular expression

defining a specific control or a specific set of controls.

group1 vector of integers (column indices) indicating all group 1 samples (optional).

group2 vector of integers (column indices) indicating all group 2 samples (optional).

output.path output.path for ProtoArray rlm normalization (optional).

#### **Details**

This function is partially a wrapper to limma's function normalizeBetweenArrays() for interarray normalization featuring optional groupwise normalization when the arguments group1 AND group2 are assigned. For more information on "cyclicloess", "quantile" or "vsn" see the documentation of the limma package. Furthermore, for ProtoArrays robust linear normalization ("rlm", see Sboner A. et al.) is provided.

For rlm normalization (method = "rlm") the additional argument controls needs to be specified in order to select a set of controls used for normalization. Valid options are "internal" (default), "external" and "both" which refer to the following sets of ProtoArray controls:

- internal: The set of all internal controls spotted on the ProtoArray. The human-IgG series and anti-human-IgG series, which respond to serum and secondary antibodies.
- external: The V5-CMK1 series spotted on the ProtoArray which responds to exogenously added anti-V5 antibody (external control).
- both: The combined set of both the internal and the external controls (i.e., the human-IgG and anti-human-IgG series and the V5-CMK1 series).

Moreover, via controls a regular expression can be passed in order to select a more specific group of controls. Please check the column "Name" in your gpr files in order to obtain the complete list of names of all controls spotted on the ProtoArray. In the following some examples of valid regular expressions are given:

- "^HumanIg" Only human IgGs and IgAs are selected (esp., no anti-human Igs).
- "Anti-HumanIgA" Only anti-human-IgAs are selected (esp., no human IgGs and IgAs).
- "(Anti-HumanIg|^V5control|BSA|ERa)" Only anti-human IgGs and anti-human IgAs, the V5-CMK1 series, BSA and ERa are selected.
- "HumanIgG" Only human IgGs and anti-human IgGs are selected.
- "V5control" Only the V5-CMK1 series is selected.

## Value

An EList object with the normalized data in log2 scale is returned.

12 plotArray

#### Author(s)

Michael Turewicz, <michael.turewicz@rub.de>

#### References

The package limma by Gordon Smyth et al. can be downloaded from Bioconductor (http://www.bioconductor.org/).

Smyth, G. K. (2005). Limma: linear models for microarray data. In: Bioinformatics and Computational Biology Solutions using R and Bioconductor, R. Gentleman, V. Carey, S. Dudoit, R. Irizarry, W. Huber (eds.), Springer, New York, pages 397-420.

Sboner A. et al., Robust-linear-model normalization to reduce technical variability in functional protein microarrays. J Proteome Res 2009, 8(12):5451-5464.

## **Examples**

```
cwd <- system.file(package="PAA")
load(paste(cwd, "/extdata/Alzheimer.RData", sep=""))
elist <- elist[elist$genes$Block < 10,]
normalized.elist <- normalizeArrays(elist=elist, method="quantile")</pre>
```

plotArray

Plot ProtoArray expression intensities in the original arrangement mimicking the original scan image.

## **Description**

Uses the "Block", "Row" and "Column" information of an EList or EListRaw object to resemble the original positions on the array(s). The resulting plot is similar to the original scan image of the considered array(s). Thus, this function is a visualization tool that can be used to visualize protein microarrays for which the original scan image is not available. Visual inspection of the spatial expression pattern can then identify possible local tendencies and strong spatial biases. Moreover, the array can be inspected at all stages of the preprocessing workflow in order to check the impact of the particular methods that have been applied.

## Usage

```
plotArray(elist=NULL, idx=NULL, data.type="fg", log=NULL, normalized=NULL,
   aggregation=NULL, colpal="heat.colors", graphics.device="tiff",
   output.path=NULL)
```

## Arguments

elist EList or EListRaw object (mandatory).

idx integer, vector of integers or the string "all" indicating the column indices of

the sample(s) for drawing the plot(s) (mandatory).

plotArray 13

be plotted. The default is "fg" (optional).

logical indicating whether the input data is logarithmized. If TRUE the log2 scale is expected. If FALSE a log2-transformation will be performed (mandatory).

normalized logical indicating whether elist was normalized (mandatory).

string indicating whether the data stored in elist has been aggregated and, if this is the case, which method has been used by the function loadGPR(). Possible values are "min", "mean" and "none" (mandatory).

colpal string indicating the color palette for the plot(s). The default is "heat.colors"

(optional).

graphics.device

data.type

string indicating the file format for the plot(s) saved in output.path. Accepted

string indicating whether the foreground ("fg") or background ("bg") data should

values are "tiff" and "png". The default is "tiff" (optional).

output.path string indicating the output path for the plots (optional).

### **Details**

This function allows plotting of protein microarray data using the gplots function heatmap.2() for visual quality control. The data obtained from an EList or EListRaw object is re-ordered and represented in the same way the spots are ordered on the actual microarray. Consequently, the resulting plot is similar to the original scan image of the considered array. This allows for visual control and assessment of possible patterns in spatial distribution.

Mandatory arguments are elist, idx, log, normalized and aggregation. While elist specifies the EList or EListRaw object to be used, idx designates the array column index in elist to plot a single array from the EList object. Alternatively, a vector (e.g., 1:5) or the string "all" can be designated to include multiple, respectively, all arrays that were imported.

Furthermore, data.type allows for plotting of "fg", foreground data (i.e., elist\$E and elist\$C), which is the default or "bg", background data (i.e., elist\$Eb and elist\$Cb).

The normalization approaches of PAA which comprise also data logarithmization do not include control data. With normalized=TRUE it is indicated that the input data was normalized, so the control data will be logarithmized (log2) before plotting as well. However, since the complete data (foreground and background values of protein features and control spots) can be logarithmized regardless of normalization the argument log states whether the designated data is already logarithmized (note: log2 scale is always expected).

The parameter aggregation indicates whether the protein microarray data has been aggregated by loadGPR() and, if so, which method has been used.

Moreover, the parameter colpal defines the color palette that will be used for the plot. Some exemplary values are "heat.colors" (default), "terrain.colors", "topo.colors", "greenred" and "bluered".

Finally, the output path optionally can be specified with the argument output.path to save the plot(s). Then, one or more tiff or png file(s) containing the corresponding plot(s) are saved into the subfolder "array\_plots".

14 plotFeatures

#### Value

No value is returned.

#### Note

Please note the instructions of the PAA function loadGPR(). Note that the data has to be imported including controls to avoid annoying gaps in the plot (for ProtoArrays this is done automatically and for other types of arrays the arguments description, description. features and description.discard must be defined). Note that the data can be imported without aggregation by loadGPR() (when aggregation="none") in order to inspect the array visually with plotArray() before duplicate aggregation.

#### Author(s)

Daniel Bemmerl and Michael Turewicz <michael.turewicz@rub.de>

#### References

The package gplots by Gregory R. Warnes et al. can be downloaded from CRAN (http://CRAN. R-project.org/package=gplots).

Gregory R. Warnes, Ben Bolker, Lodewijk Bonebakker, Robert Gentleman, Wolfgang Huber, Andy Liaw, Thomas Lumley, Martin Maechler, Arni Magnusson, Steffen Moeller, Marc Schwartz and Bill Venables (2015). gplots: Various R Programming Tools for Plotting Data. R package version 2.17.0. http://CRAN.R-project.org/package=gplots

### **Examples**

```
cwd <- system.file(package="PAA")
load(paste(cwd, "/extdata/BadData.RData", sep=""))
plotArray(elist=bad.elist, idx=1, data.type="bg", log=FALSE, normalized=FALSE,
    aggregation="none")</pre>
```

plotFeatures

Plot intensities of features.

### **Description**

Plots intensities of all given features (one sub-plot per feature) in group- specific colors.

# Usage

```
plotFeatures(features = NULL, elist = NULL, n1 = NULL, n2 = NULL,
group1 = "group1", group2 = "group2", output.path = NULL)
```

plotFeatures 15

# Arguments

features	vector containing "BRC"-IDs (mandatory).
elist	EListRaw or EList object containing all intensity data in log2 scale (mandatory).
n1	integer indicating the sample size of group 1 (mandatory).
n2	integer indicating the sample size of group 2 (mandatory).
group1	class label of group 1.
group2	class label of group 2.
output.path	string indicating the folder where the figure will be saved (optional).

### **Details**

Plots intensities of given features (e.g., selected by the function selectFeatures()) in group-specific colors (one sub-plot per feature). All sub-plots are aggregated to one figure. When the argument output.path is not NULL this figure will be saved in a tiff file in output.path. This function can be used to check whether the selected features are differential.

### Value

No value is returned.

#### Author(s)

Michael Turewicz, <michael.turewicz@rub.de>

16 plotFeaturesHeatmap

plotFeaturesHeatmap

Plot feature intensities as a heatmap.

## **Description**

Plots intensities of given features as a heatmap.

#### Usage

```
plotFeaturesHeatmap(features = NULL, elist = NULL, n1 = NULL, n2 = NULL, output.path = NULL, description=FALSE)
```

## **Arguments**

features	vector containing "BRC"-IDs (mandatory).
elist	${\tt EListRaw}$ or ${\tt EList}$ object containing all intensity data in log2 scale (mandatory).
n1	integer indicating the sample size of group 1 (mandatory).
n2	integer indicating the sample size of group 2 (mandatory).
output.path	path for saving the heatmap as a tiff file (default: NULL).
description	if TRUE, features will be described via protein names instead of UniProtKB

accessions (default: FALSE).

### **Details**

Plots intensities of all features given in the vector features via their corresponding "BRC"-IDs as a heatmap. If description is TRUE (default: FALSE), features will be described via protein names instead of UniProtKB accessions. Furthermore, if output.path is not NULL, the heatmap will be saved as a tiff file in output.path. This function can be used to check whether the selected features are differential.

#### Value

No value is returned.

## Author(s)

Michael Turewicz, <michael.turewicz@rub.de>

```
cwd <- system.file(package="PAA")
load(paste(cwd, "/extdata/Alzheimer.RData", sep=""))
#elist <- elist[elist$genes$Block < 10,]

#c1 <- paste(rep("AD",20), 1:20, sep="")
#c2 <- paste(rep("NDC",20), 1:20, sep="")</pre>
```

plotFeaturesHeatmap. 2 Alternative function to plot feature intensities as a heatmap.

## **Description**

This function is an alternative to plotFeaturesHeatmap() and is based on the function heatmap. 2() provided by the package gplots.

## Usage

```
plotFeaturesHeatmap.2(features = NULL, elist = NULL, n1 = NULL, n2 = NULL,
  output.path = NULL, description=FALSE)
```

## **Arguments**

features	vector containing the selected features as "BRC"-IDs (mandatory).
elist	EListRaw or EList object containing all intensity data in log2 scale (mandatory).
n1	integer indicating the sample size of group 1 (mandatory).
n2	integer indicating the sample size of group 2 (mandatory).
output.path	path for saving the heatmap as a png file (default: NULL).
description	if TRUE, features will be described via protein names instead of UniProtKB accessions (default: FALSE).

#### **Details**

Plots intensities of all features given in the vector features via their corresponding "BRC"-IDs as a heatmap. If description is TRUE (default: FALSE), features will be described via protein names instead of UniProtKB accessions. Furthermore, if output.path is not NULL, the heatmap will be saved as a png file in output.path. This function can be used to check whether the selected features are differential.

plotFeaturesHeatmap.2() is an alternative to plotFeaturesHeatmap() and is based on the function heatmap.2() provided by the package gplots.

18 plotMAPlots

#### Value

No value is returned.

### Author(s)

Ivan Grishagin (Rancho BioSciences LLC, San Diego, CA, USA), John Obenauer (Rancho Bio-Sciences LLC, San Diego, CA, USA) and Michael Turewicz (Ruhr-University Bochum, Bochum, Germany), <michael.turewicz@rub.de>

#### References

The package gplots by Gregory R. Warnes et al. can be downloaded from CRAN (http://CRAN. R-project.org/package=gplots).

Gregory R. Warnes, Ben Bolker, Lodewijk Bonebakker, Robert Gentleman, Wolfgang Huber, Andy Liaw, Thomas Lumley, Martin Maechler, Arni Magnusson, Steffen Moeller, Marc Schwartz and Bill Venables (2015). gplots: Various R Programming Tools for Plotting Data. R package version 2.17.0. http://CRAN.R-project.org/package=gplots

### **Examples**

```
cwd <- system.file(package="PAA")
load(paste(cwd, "/extdata/Alzheimer.RData", sep=""))
#elist <- elist[elist$genes$Block < 10,]

#c1 <- paste(rep("AD",20), 1:20, sep="")
#c2 <- paste(rep("NDC",20), 1:20, sep="")

#pre.sel.results <- preselect(elist=elist, columns1=c1, columns2=c2, label1="AD",
# label2="NDC", discard.threshold=0.1, fold.thresh=1.9, discard.features=TRUE,
# method="tTest")
#elist <- elist[-pre.sel.results$discard,]

#selectFeatures.results <- selectFeatures(elist,n1=20,n2=20,label1="AD",
# label2="NDC",selection.method="rf.rfe",preselection.method="none",subruns=2,
# k=2,candidate.number=20,method="frequency")

load(paste(cwd, "/extdata/selectFeaturesResultsFreq.RData", sep=""))
plotFeaturesHeatmap.2(features=selectFeatures.results$features, elist=elist,
n1=20, n2=20, description=TRUE)</pre>
```

plotMAPlots

Check normalization results with MA plots.

## Description

Draws MA plots of raw data and data after all kinds of normalization provided by PAA.

plotMAPlots 19

### Usage

```
plotMAPlots(elist = NULL, idx="all", include.rlm=FALSE, controls="internal",
output.path = NULL)
```

## **Arguments**

elist	EListRaw object containing raw data (mandatory).
idx	integer indicating the column index of the sample for drawing MA plots or the string 'all' for drawing MA plots for all samples (default: all).
include.rlm	logical indicating whether RLM normalization should be included (for ProtoArrays only; deafault: FALSE).
controls	string indicating the ProtoArray controls for rlm normalization (optional). Valid options are "internal" (default), "external", "both" or a regular expression defining a specific control or a specific set of controls.
output.path	string indicating the folder where the tiff files will be saved (mandatory when

### **Details**

When idx="all" (default) for each microarray a tiff file containing MA plots for raw data, cyclicoess normalized data, quantile normalized data and vsn normalized data (and, optionally, for ProtoArrays, rlm normalized data) will be created. When idx is an integer indicating the column index of a particular sample, MA plots only for this sample will be created. For A and M value computation the artificial median array is used as reference signal. All figures can be saved in output.path (mandatory when idx="all"). The resulting MA plots can be used to compare the results of the different normalization methods.

## Value

No value is returned.

### Author(s)

Michael Turewicz, <michael.turewicz@rub.de>

idx='all').

```
cwd <- system.file(package="PAA")
load(paste(cwd, "/extdata/Alzheimer.RData", sep=""))
elist <- elist[elist$genes$Block == 1,]
plotMAPlots(elist=elist, idx=1)</pre>
```

20 plotNormMethods

plotNormMethods Check normalization results with boxplots.
--

### **Description**

Draws sample-wise boxplots of raw data and data after all kinds of normalization provided by PAA.

## Usage

```
plotNormMethods(elist = NULL, include.rlm=FALSE, controls="internal",
output.path = NULL)
```

## **Arguments**

elist EListRaw object containing raw data (mandatory).

include.rlm logical indicating whether RLM normalization should be included (for ProtoAr-

rays only, deafault: FALSE).

controls string indicating the ProtoArray controls for rlm normalization (optional). Valid

options are "internal" (default), "external", "both" or a regular expression

defining a specific control or a specific set of controls.

output.path string indicating a folder for saving the boxplots as tiff files (optional).

#### **Details**

For each normalization approach sample-wise boxplots are created. All boxplots can be saved as high-quality tiff files (when an output path has been specified via the argument output.path). The resulting boxplots can be used to compare the results of different normalization methods.

#### Value

No value is returned.

#### Author(s)

Michael Turewicz, <michael.turewicz@rub.de>

```
cwd <- system.file(package="PAA")
load(paste(cwd, "/extdata/Alzheimer.RData", sep=""))
elist <- elist[elist$genes$Block == 1,]
plotNormMethods(elist=elist)</pre>
```

preselect 21

preselect	Score and preselect features.	

# Description

Iterates all features to score them via mMs, Student's t-test, or mRMR. Optionally, a list of not informative features can be obtained (for discarding them).

# Usage

```
preselect(elist=NULL, columns1=NULL, columns2=NULL, label1="A", label2="B",
    log=NULL, discard.threshold=0.5, fold.thresh=1.5, discard.features=TRUE,
    mMs.above=1500, mMs.between=400, mMs.matrix1=NULL,
    mMs.matrix2=NULL, method=NULL)
```

## **Arguments**

elist	EListRaw or EList object (mandatory).
columns1	column name vector (string vector) of group 1 (mandatory).
columns2	column name vector (string vector) of group 2 (mandatory).
label1	class label of group 1.
label2	class label of group 2.
log	indicates whether the data is in log scale (mandatory; note: if TRUE log2 scale is expected).
discard.thresh	
	positive numeric between 0 and 1 indicating the maximum mMs or, respectively, the maximum t-test p-value for features to be included for further analysis. Default is "0.5".
fold.thresh	numeric indicating the minimum fold change for features to be included for further analysis. Default is "1.5".
discard.features	
	boolean indicating whether merely feature scores (i.e., mMs or t-test p-values) (="FALSE") or feature scores and a discard list (="TRUE") should be returned. Default is "TRUE".
mMs.above	mMs above parameter (integer). Default is "1500".
mMs.between	mMs between parameter (integer). Default is "400".
mMs.matrix1	precomputed mMs reference matrix (see mMsMatrix()) for group 1 (mandatory).
mMs.matrix2	precomputed mMs reference matrix (see $mMsMatrix()$ ) for group 2 (mandatory).
method	preselection method ( "mMs", "tTest", "mrmr"). Default is "mMs".

22 preselect

#### **Details**

This function takes an EListRaw or EList object and group-specific column vectors. Furthermore, the class labels of group 1 and group 2 are needed. If discard.features is "TRUE" (default), all features that are considered as not differential will be collected and returned for discarding.

If method = "mMs", additionally precomputed mMs reference matrices (see mMsMatrix()) for group 1 and group 2 will be needed to compute mMs values (see Love B.) as scoring method. All mMs parameters (mMs.above and mMs.between) can be set. The defaults are "1500" for mMs.above and "400" for mMs.between. Features having an mMs value larger than discard.threshold (here: numeric between 0.0 and 1.0) or do not satisfy the minimal absolute fold change fold.thresh are considered as not differential.

If method = "tTest", Student's t-test will be used as scoring method. Features having a p-value larger than discard. threshold (here: numeric between 0.0 and 1.0) or do not satisfy the minimal absolute fold change fold. thresh are considered as not differential.

If method = "mrmr", mRMR scores for all features will be computed as scoring method (using the function mRMR.classic() of the CRAN R package mRMRe). Features that are not the discard. threshold (here: integer indicating a number of features) best features regarding their mRMR score are considered as not differential.

#### Value

If discard.features is "FALSE": matrix containing metadata, feature scores and intensity values for the whole data set.

If discard. features is "TRUE", a list containing:

results matrix containing metadata, feature scores and intensity values for the whole

data set.

discard vector containing row indices (= features) for discarding features considered as

not differential.

#### Author(s)

Michael Turewicz, <michael.turewicz@rub.de>

# References

Love B: The Analysis of Protein Arrays. In: Functional Protein Microarrays in Drug Discovery. CRC Press; 2007: 381-402.

The software "Prospector" for ProtoArray analysis can be downloaded from the Thermo Fisher Scientific web page (https://www.thermofisher.com).

The R package mRMRe can be downloaded from CRAN. See also: De Jay N, Papillon-Cavanagh S, Olsen C, El-Hachem N, Bontempi G, Haibe-Kains B. mRMRe: an R package for parallelized mRMR ensemble feature selection. Bioinformatics 2013.

The package limma by Gordon Smyth et al. can be downloaded from Bioconductor (https://www.bioconductor.org).

Smyth, G. K. (2005). Limma: linear models for microarray data. In: Bioinformatics and Computational Biology Solutions using R and Bioconductor, R. Gentleman, V. Carey, S. Dudoit, R. Irizarry, W. Huber (eds.), Springer, New York, pages 397-420.

printFeatures 23

### **Examples**

```
cwd <- system.file(package="PAA")
load(paste(cwd, "/extdata/Alzheimer.RData", sep=""))
elist <- elist[elist$genes$Block < 10,]
c1 <- paste(rep("AD",20), 1:20, sep="")
c2 <- paste(rep("NDC",20), 1:20, sep="")
preselect(elist, columns1=c1, columns2=c2, label1="AD", label2="NDC", log=FALSE,
    discard.threshold=0.5, fold.thresh=1.5, discard.features=TRUE, method="tTest")</pre>
```

printFeatures

Print features into a table.

### **Description**

Creates a table containing the given features (e.g., the selected biomarker candidate panel).

#### Usage

```
printFeatures(features = NULL, elist = NULL, output.path = NULL)
```

### **Arguments**

features vector containing "BRC"-IDs (mandatory).

elist EListRaw or EList object containing all intensity data (mandatory).

output.path string indicating the folder where the table will be saved as a txt file (optional).

#### **Details**

Creates a table containing the given features (e.g., the selected biomarker candidate panel) as well as additional information. When output.path is defined this table will be saved in a txt file ("candidates.txt").

## Value

Table containing the given features.

## Author(s)

Michael Turewicz, <michael.turewicz@rub.de>

24 pvaluePlot

### **Examples**

```
cwd <- system.file(package="PAA")
load(paste(cwd, "/extdata/Alzheimer.RData", sep=""))
#elist <- elist[elist$genes$Block < 10,]

#c1 <- paste(rep("AD",20), 1:20, sep="")
#c2 <- paste(rep("NDC",20), 1:20, sep="")

#pre.sel.results <- preselect(elist=elist, columns1=c1, columns2=c2, label1="AD",
# label2="NDC", discard.threshold=0.1, fold.thresh=1.9, discard.features=TRUE,
# method="tTest")
#elist <- elist[-pre.sel.results$discard,]

#selectFeatures.results <- selectFeatures(elist,n1=20,n2=20,label1="AD",
# label2="NDC",selection.method="rf.rfe",preselection.method="none",subruns=2,
# k=2,candidate.number=20,method="frequency")

load(paste(cwd, "/extdata/selectFeaturesResultsFreq.RData", sep=""))
printFeatures(features=selectFeatures.results$features, elist=elist)</pre>
```

pvaluePlot

Draw a p-value plot.

### **Description**

Draws a p-value plot to visualize the p-values for all features stored in a EList or EListRaw object.

### Usage

```
pvaluePlot(elist=NULL, group1=NULL, group2=NULL, log=NULL, method="tTest",
output.path=NULL, tag="", mMs.matrix1=NULL, mMs.matrix2=NULL, above=1500,
between=400, adjust=FALSE)
```

#### Arguments

elist	EList or EListRaw object (mandatory).
group1	vector of column names for group 1 (mandatory).
group2	vector of column names for group 2 (mandatory).
log	indicates whether the data is in log scale (mandatory; note: if TRUE $\log 2$ scale is expected).
method	method for p-value computation: "tTest" or "mMs". Default is "tTest".
output.path	string indicating a path for saving the plot (optional).
tag	string that can be used for tagging the saved plot (optional).
mMs.matrix1	precomputed M score reference matrix (see $mMsMatrix()$ ) for group 1 (mandatory when method = $"mMs"$ ).

mMs.matrix2 precomputed M score reference matrix (see mMsMatrix()) for group 2 (manda-

tory when method = "mMs").

above M score above parameter (integer). Default is "1500".

between M score between parameter (integer). Default is "400".

adjust logical indicating whether p-values should be adjusted. Default is FALSE.

#### **Details**

This function takes an EList or EListRaw object and the corresponding column name vectors to draw a plot of p-values for all features stored in elist (sorted in increasing order and in log2 scale). The p-value computation method ("tTest" or "mMs") can be set via the argument method. Furthermore, when adjust=TRUE adjusted p-values (method: Benjamini & Hochberg, 1995, computed via p.adjust()) will be used. When an output path is defined (via output.path) the plot will be saved as a tiff file.

#### Value

No value is returned.

### Author(s)

Michael Turewicz, <michael.turewicz@rub.de>

### **Examples**

```
cwd <- system.file(package="PAA")
load(paste(cwd, "/extdata/Alzheimer.RData", sep=""))
elist <- elist[elist$genes$Block < 10,]
c1 <- paste(rep("AD",20), 1:20, sep="")
c2 <- paste(rep("NDC",20), 1:20, sep="")
pvaluePlot(elist=elist, group1=c1, group2=c2, log=FALSE, method="tTest", tag="_tTest", adjust=FALSE)</pre>
```

selectFeatures

Select features using frequency-based or ensemble feature selection.

### **Description**

Performs a multivariate feature selection using frequency-based feature selection (based on RF-RFE, RJ-RFE or SVM-RFE) or ensemble feature selection (based on SVM-RFE).

#### Usage

```
selectFeatures(elist = NULL, n1 = NULL, n2 = NULL, label1 = "A", label2 = "B",
log=NULL, cutoff = 10, selection.method = "rf.rfe",
preselection.method = "mMs", subruns = 100, k = 10, subsamples = 10,
bootstraps = 10, candidate.number = 300, above=1500, between=400,
panel.selection.criterion="accuracy", importance.measure="MDA", ntree = 500,
mtry = NULL, plot = FALSE, output.path = NULL, verbose = FALSE,
method = "frequency")
```

#### **Arguments**

elist EListRaw or EList object containing all microarray data (mandatory). n1 integer indicating the sample number in group 1 (mandatory). integer indicating the sample number in group 2 (mandatory). n2 label1 class label of group 1 (default: "A"). class label of group 2 (default: "B"). label2 indicates whether the data is in log scale (mandatory; note: if TRUE log2 scale log is expected). cutoff integer indicating how many features will be selected (default: 10). selection.method string indicating the feature selection method: "rf.rfe" (default), "svm.rfe" or "rj.rfe". Has no effect when method="ensemble". preselection.method string indicating the feature preselection method: "mMs" (default), "tTest", "mrmr" or "none". Has no effect when method="ensemble". integer indicating the number of resampling repeats to be performed (default: subruns 100). Has no effect when method="ensemble". k integer indicating the number of k-fold cross validation subsets (default: 10, i.e., 10-fold CV). subsamples integer indicating the number of subsamples for ensemble feature selection (default: 10). Has no effect when method="frequency". integer indicating the number of bootstrap samples for ensemble feature selecbootstraps tion (default: 10). Has no effect when method="frequency" only. candidate.number

integer indicating how many features shall be preselected. Default is "300". Has no effect when method="ensemble".

mMs above parameter (integer). Default is "1500". There will be no effect when method="ensemble".

mMs between parameter (integer). Default is "400". There will be no effect when method="ensemble".

panel.selection.criterion

above

between

indicating the panel selection criterion: "accuracy" (default), "sensitivity" or "specificity". No effect for method="ensemble".

importance.measure

string indicating the random forest importance measure: "MDA" (default) or

"MDG". Has no effect when method="ensemble".

ntree random forest parameter ntree (default: "500"). There will be no effect when

method="ensemble".

mtry random forest parameter mtry (default: sqrt(p) where p is the number of pre-

dictors). Has no effect when method="ensemble".

plot logical indicating whether performance plots shall be plotted (default: FALSE).

output.path string indicating the results output folder (optional).

verbose logical indicating whether additional information shall be printed to the console

(default: FALSE).

method the feature selection method: "frequency" (default) for frequency-based or "en-

semble" for ensemble feature selection.

#### **Details**

This function takes an EListRaw or EList object, group-specific sample numbers, group labels and parameters choosing and configuring a multivariate feature selection method (frequency-based or ensemble feature selection) to select a panel of differential features. When an output path is defined (via output.path) results will be saved on the hard disk and when verbose is TRUE additional information will be printed to the console.

Frequency-based feature selection (method="frequency"): The whole data is splitted in k cross validation training and test set pairs. For each training set a multivariate feature selection procedure is performed. The resulting k feature subsets are tested using the corresponding test sets (via classification). As a result, selectFeatures() returns the average k-fold cross validation classification accuracy as well as the selected feature panel (i.e., the union set of the k particular feature subsets). As multivariate feature selection methods random forest recursive feature elimination (RF-RFE), random jungle recursive feature elimination (RJ-RFE) and support vector machine recursive feature elimination (SVM-RFE) are supported. To reduce running times, optionally, univariate feature preselection can be performed (control via preselection.method). As univariate preselection methods mMs ("mMs"), Student's t-test ("tTest") and mRMR ("mrmr") are supported. Alternatively, no preselection can be chosen ("none"). This approach is similar to the method proposed in Baek et al.

Ensemble feature selection (method="ensemble"): From the whole data the previously defined number of subsamples is drawn defining pairs of training and test sets. Moreover, for each training set a previously defined number of bootstrap samples is drawn. Then, for each bootstrap sample SVM-RFE is performed and a feature ranking is obtained. To obtain a final ranking for a particular training set, all associated bootstrap rankings are aggregated to a single ranking. To score the cutoff best features, for each subsample a classification of the test set is performed (using a sym trained with the cutoff best features from the training set) and the classification accuracy is determined. Finally, the stability of the subsample-specific panels is assessed (via Kuncheva index, Kuncheva LI, 2007), all subsample-specific rankings are aggregated, the top n features (defined by cutoff) are selected, the average classification accuracy is computed, and all these results are returned in a list. This approach has been proposed in Abeel et al.

#### Value

If method is "frequency", the results list contains the following elements:

accuracy average k-fold cross validation accuracy.
sensitivity average k-fold cross validation sensitivity.
specificity average k-fold cross validation specificity.

features selected feature panel.

all.results complete cross validation results.

If method is "ensemble", the results list contains the following elements:

accuracy average accuracy regarding all subsamples. sensitivity average sensitivity regarding all subsamples. specificity average specificity regarding all subsamples.

features selected feature panel.
all.results all feature ranking results.

stability stability of the feature panel (i.e., Kuncheva index for the subrun-specific pan-

els).

## Author(s)

Michael Turewicz, <michael.turewicz@rub.de>

#### References

Baek S, Tsai CA, Chen JJ.: Development of biomarker classifiers from high-dimensional data. Brief Bioinform. 2009 Sep;10(5):537-46.

Abeel T, Helleputte T, Van de Peer Y, Dupont P, Saeys Y: Robust biomarker identification for cancer diagnosis with ensemble feature selection methods. Bioinformatics. 2010 Feb 1;26(3):392-8.

Kuncheva, LI: A stability index for feature selection. Proceedings of the IASTED International Conference on Artificial Intelligence and Applications. February 12-14, 2007. Pages: 390-395.

```
cwd <- system.file(package="PAA")
load(paste(cwd, "/extdata/Alzheimer.RData", sep=""))
elist <- elist[elist$genes$Block < 10,]

c1 <- paste(rep("AD",20), 1:20, sep="")
c2 <- paste(rep("NDC",20), 1:20, sep=""))

pre.sel.results <- preselect(elist=elist, columns1=c1, columns2=c2, label1="AD",
    label2="NDC", log=FALSE, discard.threshold=0.1, fold.thresh=1.9,
    discard.features=TRUE, method="tTest")
elist <- elist[-pre.sel.results$discard,]

selectFeatures.results <- selectFeatures(elist, n1=20, n2=20, label1="AD",
    label2="NDC", log=FALSE, subsamples=2, bootstraps=1, candidate.number=20,
    method="ensemble")</pre>
```

shuffleData 29

shuffleData	Shuffles class labels to obtain random groups.	

# Description

Shuffles class labels of an EList or EListRaw object randomly to obtain two random groups (e.g. "A" and "B").

## Usage

```
shuffleData(elist=NULL, n1=NULL, n2=NULL, label1="A", label2="B")
```

# Arguments

elist	EList or EListRaw object (mandatory).
n1	sample size of random group 1 (mandatory).
n2	sample size of random group 2 (mandatory).
label1	class label of random group 1 (default: "A").
label2	class label of random group 2 (default: "B").

### **Details**

Shuffles class labels of an EList or EListRaw object randomly to obtain two random groups (e.g. "A" and "B").

## Value

EList or EListRaw object with random groups.

## Author(s)

Michael Turewicz, <michael.turewicz@rub.de>

```
cwd <- system.file(package="PAA")
load(paste(cwd, "/extdata/Alzheimer.RData", sep=""))
shuffleData(elist=elist, n1=20, n2=20, label1="A", label2="B")</pre>
```

30 volcanoPlot

# **Description**

Draws a volcano plot to visualize differential features.

## Usage

```
volcanoPlot(elist=NULL, group1=NULL, group2=NULL, log=NULL, method="tTest",
p.thresh=NULL, fold.thresh=NULL, output.path=NULL, tag="", mMs.matrix1=NULL,
mMs.matrix2=NULL, above=1500, between=400)
```

### **Arguments**

elist	EList or EListRaw object (mandatory).
group1	vector of column names for group 1 (mandatory).
group2	vector of column names for group 2 (mandatory).
log	indicates whether the data is in log scale (mandatory; note: if TRUE log2 scale is expected; mandatory).
method	method for p-value computation: "tTest" or "mMs". Default is "tTest".
p.thresh	positive float number between 0 and 1 indicating the maximum p-value for features to be considered as differential (e.g., "0.5"). This argument is optional.
fold.thresh	float number indicating the minimum fold change for features to be considered as differential (e.g., "1.5"). This argument is optional.
output.path	string indicating a path for saving the plot (optional).
tag	string that can be used for tagging the saved plot (optional).
mMs.matrix1	a precomputed M score reference matrix (see $mMsMatrix()$ ) for group 1 (mandatory when method = $mMs$ ).
mMs.matrix2	a precomputed M score reference matrix (see $mMsMatrix()$ ) for group 2 (mandatory when method = $mMs$ ).
above	M score above parameter (integer). Default is "1500".
between	M score between parameter (integer). Default is "400".

# **Details**

This function takes an EList or EListRaw object and the corresponding column name vectors to draw a volcano plot. To visualize differential features, thresholds for p-values and fold changes can be defined. Furthermore, the p-value computation method ("mMs" or "tTest") can be set. When an output path is defined (via output.path) the plot will be saved as a tiff file.

## Value

No value is returned.

volcanoPlot 31

## Author(s)

Michael Turewicz, <michael.turewicz@rub.de>

```
cwd <- system.file(package="PAA")
load(paste(cwd, "/extdata/Alzheimer.RData", sep=""))
elist <- elist[elist$genes$Block < 10,]
c1 <- paste(rep("AD",20), 1:20, sep="")
c2 <- paste(rep("NDC",20), 1:20, sep="")
volcanoPlot(elist=elist, group1=c1, group2=c2, log=FALSE, method="tTest", p.thresh=0.01, fold.thresh=2)</pre>
```

# **Index**

* <b>Differential analysis</b> diffAnalysis, 5	plotMAPlots, 18 plotNormMethods, 20
pvaluePlot, 24	preselect, 21
volcanoPlot, 30	printFeatures, 23
* Feature selection	pvaluePlot, 24
plotFeatures, 14	p ( a 2 a c , 2 )
plotFeaturesHeatmap, 16	selectFeatures, 25
plotFeaturesHeatmap.2,17	shuffleData, 29
preselect, 21	
printFeatures, 23	volcanoPlot, 30
selectFeatures, 25	
* Input/output	
loadGPR, 7	
* Preprocessing	
batchAdjust, 2	
batchAdjust, 2 batchFilter, 3	
batchFilter.anova,4	
normalizeArrays, 10	
plotArray, 12	
plotMAPlots, 18	
plotNormMethods, 20	
shuffleData, 29	
* mMs	
mMsMatrix, 9	
batchAdjust, 2	
batchFilter, 3	
batchFilter.anova,4	
diffAnalysis, 5	
loadGPR, 7	
mMsMatrix,9	
normalizeArrays, 10	
plotArray, 12 plotFeatures, 14 plotFeaturesHeatmap, 16 plotFeaturesHeatmap.2, 17	