

# Package ‘POMA’

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**Title** User-friendly Workflow for Metabolomics and Proteomics Data Analysis

**Version** 1.4.0

**Description** A structured, reproducible and easy-to-use workflow for the visualization, pre-processing, exploratory data analysis, and statistical analysis of metabolomics and proteomics data. The main aim of POMA is to enable a flexible data cleaning and statistical analysis processes in one comprehensible and user-friendly R package. This package also has a Shiny app version that implements all POMA functions. See <https://github.com/pcastellanoescuder/POMAShiny>.

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PomaBoxplots

*Classical Boxplots*

---

### Description

PomaBoxplots() generates a boxplot for subjects or features. This boxplot can help in the comparison between pre and post normalized data and in the "validation" of the normalization process.

**Usage**

```
PomaBoxplots(  
  data,  
  group = "samples",  
  jitter = TRUE,  
  feature_name = NULL,  
  label_size = 10,  
  legend_position = "bottom"  
)
```

**Arguments**

|                              |  |
|------------------------------|--|
| <code>data</code>            | A MSnSet object. First pData column must be the subject group/type.  |
| <code>group</code>           | Grouping factor for the plot. Options are "samples" and "features". Option "samples" (default) will create a boxplot for each sample and option "features" will create a boxplot of each variable. |
| <code>jitter</code>          | Logical. If it's TRUE (default), the boxplot will show all points.   |
| <code>feature_name</code>    | A vector with the name/s of feature/s to plot. If it's NULL (default) a boxplot of all features will be created.   |
| <code>label_size</code>      | Numeric indicating the size of x-axis labels.  |
| <code>legend_position</code> | Character indicating the legend position. Options are "none", "top", "bottom", "left", and "right".  |

**Value**

A ggplot2 object.

**Author(s)**

Pol Castellano-Escuder

**Examples**

```
data("st000284")  
  
# samples  
PomaBoxplots(st000284)  
  
# features  
PomaBoxplots(st000284, group = "features")  
  
# concrete features  
PomaBoxplots(st000284, group = "features",  
              feature_name = c("ornithine", "orotate"))
```

**Description**

This function performs a classical multidimensional scaling (MDS) using all features in the data and computes a cluster analysis for k clusters. Then, the calculated clusters will be represented on a MDS plot.

**Usage**

```
PomaClust(  
  data,  
  method = "euclidean",  
  k = NA,  
  k_max = 15,  
  show_clusters = TRUE,  
  labels = FALSE,  
  show_group = FALSE  
)
```

**Arguments**

|               |   |
|---------------|---|
| data          | A MSnSet object. First pData column must be the subject group/type.   |
| method        | Distance measure method to perform MDS. Options are "euclidean", "maximum", "manhattan", "canberra" and "minkowski". See ?dist().                       |
| k             | Number of clusters (default is NA). The optimum number of clusters will be used by default.   |
| k_max         | Number of clusters among which the optimal one will be selected.  |
| show_clusters | Logical indicating if clusters should be plotted or not. If this parameter is set to FALSE the resultant plot will be a classical 2-dimension MDS plot. |
| labels        | Logical indicating if sample names should be plotted or not.  |
| show_group    | Logical indicating if the original sample group from pData should be plotted instead of sample ID or not. Only works if labels is set to TRUE.          |

**Value**

A list with the results.

**Author(s)**

Pol Castellano-Escuder

**Examples**

```
data("st000284")

PomaClust(st000284)
```

---

PomaCorr

*Correlation Analysis*


---

**Description**

This function returns different correlation plots (correlogram and network plots) and a table with all pairwise correlations in the data.

**Usage**

```
PomaCorr(
  data,
  method = "pearson",
  shape = "square",
  type = "full",
  show_corr = FALSE,
  low = "#336B87",
  outline = "white",
  high = "#EA8620",
  label_size = 12,
  corr_type = "cor",
  coeff = 0.7
)
```

**Arguments**

|           |   |
|-----------|---|
| data      | A MSnSet object. First pData column must be the subject group/type.   |
| method    | Character indicating which correlation coefficient has to be computed. Options are "pearson" (default), "kendall" and "spearman".   |
| shape     | Character indicating shape of correlogram. Options are "square" (default) and "circle".   |
| type      | Character indicating type of correlogram. Options are "full" (default), "lower" or "upper".   |
| show_corr | Logical indicating if correlation coefficient for each pair of features should be plotted in correlogram or not (default = FALSE). Only recommended for a low number of features. |
| low       | Colour for low end of the gradient in correlogram.  |
| outline   | Colour for the outline of the gradient in correlogram.  |
| high      | Colour for high end of the gradient in correlogram.   |

|            |  |
|------------|--|
| label_size | Numeric indicating label size in correlogram.  |
| corr_type  | Type of network to be made with correlation matrix. Options are "cor" (for global correlations) and "glasso" (for gaussian graphical model). Default is "cor". See gLasso R package for the second option.   |
| coeff      | Numeric indicatin correlation coefficient. Edges with absolute weight below this value will be removed from the network. If "corr_type" is set to "glasso", this parameter indicates the regularization parameter for lasso (rho = 0 means no regularization). See gLasso::glasso(). |

**Value**

A list with the results.

**Author(s)**

Pol Castellano-Escuder

**References**

Jerome Friedman, Trevor Hastie and Rob Tibshirani (2019). glasso: Graphical Lasso: Estimation of Gaussian Graphical Models. R package version 1.11. <https://CRAN.R-project.org/package=glasso>

**Examples**

```
data("st000284")

# pearson correlation
PomaCorr(st000284)$correlations
PomaCorr(st000284)$corrplot

# gaussian graphical model
# library(ggraph)
# PomaCorr(st000284, corr_type = "glasso")
```

---

PomaDensity

*Distribution Plot*

---

**Description**

PomaDensity() generates a density plot of not normalized and normalized MS data. This plot can help in the comparison between pre and post normalized data and in the "validation" of the normalization process.

**Usage**

```
PomaDensity(  
  data,  
  group = "samples",  
  feature_name = NULL,  
  legend_position = "bottom"  
)
```

**Arguments**

|                              |   |
|------------------------------|---|
| <code>data</code>            | A MSnSet object. First pData column must be the subject group/type.   |
| <code>group</code>           | Grouping factor for the plot. Options are "samples" and "features". Option "samples" (default) will create a density plot for each group and option "features" will create a density plot of each variable. |
| <code>feature_name</code>    | A vector with the name/s of feature/s to plot. If it's NULL (default) a density plot of all variables will be created.  |
| <code>legend_position</code> | Character indicating the legend position. Options are "none", "top", "bottom", "left", and "right".   |

**Value**

A ggplot2 object.

**Author(s)**

Pol Castellano-Escuder

**Examples**

```
data("st000284")  
  
# samples  
PomaDensity(st000284)  
  
# features  
PomaDensity(st000284, group = "features")  
  
# concrete features  
PomaDensity(st000284, group = "features",  
            feature_name = c("ornithine", "orotate"))
```

## Description

This function automatically generates a PDF report with different exploratory plots and tables from an MSnSet object.

## Usage

```
PomaEDA(  
  data,  
  imputation = "knn",  
  normalization = "log_pareto",  
  clean_outliers = TRUE,  
  coeff_outliers = 1.5,  
  username = "Username"  
)
```

## Arguments

|                |  |
|----------------|--|
| data           | A MSnSet object. First pData column must be the subject group/type.  |
| imputation     | Imputation method. Options are "none", "half_min", "median", "mean", "min" and "knn" (default). If "none", all missing values will be replaced by zero.                    |
| normalization  | Normalization method. Options are "none", "auto_scaling", "level_scaling", "log_scaling", "log_transformation", "vast_scaling" and "log_pareto" (default).                 |
| clean_outliers | Logical. If it's set to TRUE, outliers will be removed from EDA.   |
| coeff_outliers | This value corresponds to the classical $1.5$ in $Q3 + 1.5 * IQR$ formula to detect outliers. By changing this value, the permissiveness in outlier detection will change. |
| username       | This name will be included as a report subtitle.   |

## Value

An exploratory data analysis PDF report.

## Author(s)

Pol Castellano-Escuder



---

|             |                          |
|-------------|--------------------------|
| PomaHeatmap | <i>Classical Heatmap</i> |
|-------------|--------------------------|

---

### Description

This function returns a basic heatmap plot made with base R.

### Usage

```
PomaHeatmap(  
  data,  
  sample_names = TRUE,  
  feature_names = FALSE,  
  show_legend = TRUE  
)
```

### Arguments

|               |   |
|---------------|---|
| data          | A MSnSet object. First pData column must be the subject group/type.             |
| sample_names  | Logical indicating if sample names should be plotted or not. Default is TRUE.   |
| feature_names | Logical indicating if feature names should be plotted or not. Default is FALSE. |
| show_legend   | Logical indicating if legend should be plotted or not. Default is TRUE.         |

### Value

A heatmap.

### Author(s)

Pol Castellano-Escuder

### Examples

```
data("st000284")  
  
st000284 %>%  
  PomaNorm() %>%  
  PomaHeatmap()
```

**Description**

PomaImpute() offers different methods to impute missing values in MS data.

**Usage**

```
PomaImpute(  
  data,  
  ZerosAsNA = FALSE,  
  RemoveNA = TRUE,  
  cutoff = 20,  
  method = "knn"  
)
```

**Arguments**

|           |   |
|-----------|---|
| data      | A MSnSet object. First pData column must be the subject group/type.   |
| ZerosAsNA | Logical that indicates if the zeros in the data are missing values. Default is FALSE.   |
| RemoveNA  | Logical that indicates if those features with more than selected cutoff missing values in each group have to be removed. Default is TRUE.   |
| cutoff    | Numeric that indicates the percentage of missing values allowed in each group. If one of the groups have less missing values than selected cutoff value, these feature will not be removed. |
| method    | Imputation method. Options are: "none", "half_min", "median", "mean", "min", "knn" and "rf". If "none", all missing values will be replaced by zero.  |

**Value**

A MSnSet object with cleaned data.

**Author(s)**

Pol Castellano-Escuder

**References**

Armitage, E. G., Godzien, J., Alonso-Herranz, V., López-González, Á., & Barbas, C. (2015). Missing value imputation strategies for metabolomics data. *Electrophoresis*, 36(24), 3050-3060.

**Examples**

```
data("st000336")  
  
PomaImpute(st000336, method = "knn")
```

---

|           |  |
|-----------|--|
| PomaLasso | <i>Lasso, Ridge and Elasticnet Regularized Generalized Linear Models for Binary Outcomes</i> |
|-----------|--|

---

## Description

PomaLasso() is an implementation of the lasso, ridge and elasticnet regression from glmnet package for binary outcomes.

## Usage

```
PomaLasso(
  data,
  alpha = 1,
  ntest = NULL,
  nfolds = 10,
  lambda = NULL,
  labels = FALSE
)
```

## Arguments

|        |  |
|--------|--|
| data   | A MSnSet object. First pData column must be the subject group/type.  |
| alpha  | Elasticnet mixing parameter. alpha = 1 is the lasso penalty and alpha = 0 is the ridge penalty. This value must be between 0 and 1.  |
| ntest  | Numeric indicating the percentage of observations that will be used as test set. Default is NULL (no test set).  |
| nfolds | Number of folds for CV (default is 10). Although nfolds can be as large as the sample size (leave-one-out CV), it is not recommended for large datasets. Smallest value allowable is nfolds = 3. |
| lambda | A user supplied lambda sequence. Typical usage is to have the program compute its own lambda sequence based on nlambda and lambda.min.ratio. See ?glmnet::glmnet().                              |
| labels | Logical indicating if feature names should be plotted in coefficient plot or not. Default is FALSE.  |

## Value

A list with all results including plots, data frames and the resulting prediction model.

## Author(s)

Pol Castellano-Escuder

## References

Jerome Friedman, Trevor Hastie, Robert Tibshirani (2010). Regularization Paths for Generalized Linear Models via Coordinate Descent. *Journal of Statistical Software*, 33(1), 1-22. URL <http://www.jstatsoft.org/v33/i01/>.

## Examples

```
data("st000336")

# lasso
st000336 %>%
  PomaImpute() %>%
  PomaNorm() %>%
  PomaOutliers() %>%
  PomaLasso()

# elasticnet
st000336 %>%
  PomaImpute() %>%
  PomaNorm() %>%
  PomaOutliers() %>%
  PomaLasso(alpha = 0.5)

# ridge
st000336 %>%
  PomaImpute() %>%
  PomaNorm() %>%
  PomaOutliers() %>%
  PomaLasso(alpha = 0)
```

---

PomaLimma

*Implementation of limma R Package on Mass Spectrometry Data*

---

## Description

PomaLimma() uses the classical limma package for MS data.

## Usage

```
PomaLimma(
  data,
  contrast = NULL,
  covariates = FALSE,
  adjust = "fdr",
  cutoff = NULL
)
```

**Arguments**

|            |  |
|------------|--|
| data       | A MSnSet object. First pData column must be the subject group/type.  |
| contrast   | A character with the limma comparison. For example, "Group1-Group2" or "control-intervention".   |
| covariates | Logical. If it's set to TRUE all metadata variables stored in pData will be used as covariables. Default = FALSE.  |
| adjust     | Multiple comparisons correction method. Options are: "fdr", "holm", "hochberg", "hommel", "bonferroni", "BH" and "BY".   |
| cutoff     | Default is NULL. If this value is replaced for a numeric value, the resultant table will contains only those features with an adjusted p-value below selected value. |

**Value**

A data frame with limma results.

**Author(s)**

Pol Castellano-Escuder

**References**

Matthew E. Ritchie, Belinda Phipson, Di Wu, Yifang Hu, Charity W. Law, Wei Shi, Gordon K. Smyth, limma powers differential expression analyses for RNA-sequencing and microarray studies, Nucleic Acids Research, Volume 43, Issue 7, 20 April 2015, Page e47, <https://doi.org/10.1093/nar/gkv007>

**Examples**

```
data("st000284")

st000284 %>%
  PomaNorm() %>%
  PomaLimma(contrast = "Healthy-CRC", adjust = "fdr")
```

---

|                 |   |
|-----------------|---|
| PomaMSnSetClass | <i>Convert data frames to a MSnSet Object</i> |
|-----------------|---|

---

**Description**

This function converts data frame objects to a MSnSet object.

**Usage**

```
PomaMSnSetClass(target, features)
```

**Arguments**

|          |   |
|----------|---|
| target   | Metadata variables structured in columns. Sample ID must be the first column and group/type/treatment of the study must be the second column. |
| features | Table of features. Each feature in one column.  |

**Value**

A MSnSet object.

**Author(s)**

Pol Castellano-Escuder

**References**

Laurent Gatto and Kathryn S. Lilley. MSnbase - an R/Bioconductor package for isobaric tagged mass spectrometry data visualization, processing and quantitation. *Bioinformatics* 28, 288-289 (2012).

**Examples**

```
data(iris)

# create target: two column (or more) data frame with IDs and Group factor
target <- data.frame(ID = 1:150, Group = iris$Species)

# create features: p column data frame (or matrix) with features
features <- iris[,1:4]

# create an MSnSet object with POMA
object <- PomaMSnSetClass(target = target, features = features)
```

---

PomaMultivariate

*Multivariate Statistical Methods for Mass Spectrometry Data*

---

**Description**

PomaMultivariate() allows users to perform different multivariate statistical analysis on MS data.

**Usage**

```
PomaMultivariate(  
  data,  
  method = "pca",  
  components = 5,  
  center = FALSE,  
  scale = FALSE,  
  labels = FALSE,
```

```

    load_length = 1,
    ellipse = TRUE,
    validation = "Mfold",
    folds = 5,
    nrepeat = 10,
    vip = 1.5,
    num_features = 10,
    legend_position = "bottom"
)

```

### Arguments

|                 |   |
|-----------------|---|
| data            | A MSnSet object. First pData column must be the subject group/type.   |
| method          | A multivariate method. Options are: "pca", "plsda" and "splstda".   |
| components      | Numeric. Number of components to include in the model. Default is 5.  |
| center          | Logical that indicates whether the variables should be shifted to be zero centered. Default is FALSE.   |
| scale           | Logical that indicates whether the variables should be scaled to have unit variance before the analysis takes place. Default is FALSE.                                      |
| labels          | Logical indicating if sample names should be plotted or not.  |
| load_length     | Numeric between 1 and 2. Define the length of biplot loadings. Default is 1.  |
| ellipse         | Logical that indicates whether a 95%CI ellipse should be plotted in scores plot. Default is TRUE.   |
| validation      | (Only for "plsda" and "splstda" methods) Validation method. Options are "Mfold" and "loo".  |
| folds           | (Only for "plsda" and "splstda" methods) Numeric. Number of folds for Mfold validation method (default is 5). If the validation method is loo, this value will become to 1. |
| nrepeat         | (Only for "plsda" and "splstda" methods) Numeric. Number of iterations for the validation method selected.  |
| vip             | (Only for "plsda" method) Numeric indicating VIP cutoff to select features that will be displayed in vip plot.  |
| num_features    | (Only for "splstda" method) Numeric. Number of variables selected to discriminate groups.   |
| legend_position | Character indicating the legend position. Options are "none", "top", "bottom", "left", and "right".   |

### Value

A list with all results for multivariate statistical analysis including plots and data frames.

### Author(s)

Pol Castellano-Escuder

## Examples

```
data("st000336")

# PCA
st000336 %>%
  PomaImpute() %>%
  PomaNorm() %>%
  PomaOutliers() %>%
  PomaMultivariate(method = "pca")

# PLSDA
st000336 %>%
  PomaImpute() %>%
  PomaNorm() %>%
  PomaOutliers() %>%
  PomaMultivariate(method = "plsda", vip = 1)

# sPLSDA
st000336 %>%
  PomaImpute() %>%
  PomaNorm() %>%
  PomaOutliers() %>%
  PomaMultivariate(method = "splstda")
```

---

PomaNorm

*Collection of Normalization Methods for Mass Spectrometry Data*

---

## Description

PomaNorm() offers different methods to normalize MS data. This function contains both centering and scaling functions to normalize the data.

## Usage

```
PomaNorm(data, method = "log_pareto", round = 3)
```

## Arguments

|        |   |
|--------|---|
| data   | A MSnSet object. First pData column must be the subject group/type.   |
| method | Normalization method. Options are: "none", "auto_scaling", "level_scaling", "log_scaling", "log_transformation", "vast_scaling" and "log_pareto". |
| round  | Numeric. Number of decimal places (Default is 3).   |

## Value

A MSnSet object with normalized data.



**Author(s)**

Pol Castellano-Escuder

**References**

van den Berg, R. A., Hoefsloot, H. C., Westerhuis, J. A., Smilde, A. K., & van der Werf, M. J. (2006). Centering, scaling, and transformations: improving the biological information content of metabolomics data. *BMC genomics*, 7(1), 142.

**Examples**

```
data("st000284")  
  
PomaNorm(st000284, method = "log_pareto")
```

---

|               |  |
|---------------|--|
| PomaOddsRatio | <i>Logistic Regression Model Odds Ratios</i> |
|---------------|--|

---

**Description**

PomaOddsRatio() calculates the Odds Ratios for each feature from a logistic regression model using the binary outcome (group/type must be a binary factor) as a dependent variable.

**Usage**

```
PomaOddsRatio(data, feature_name = NULL, covariates = FALSE, showCI = TRUE)
```

**Arguments**

|              |  |
|--------------|--|
| data         | A MSnSet object. First pData column must be the subject group/type.  |
| feature_name | A vector with the name/s of feature/s that will be used to fit the model. If it's NULL (default), all variables will be included in the model. |
| covariates   | Logical that indicates if covariates will be included in logistic regression model. Default is FALSE.  |
| showCI       | Logical that indicates if the 95% confidence intervals will be plotted. Default is TRUE.   |

**Value**

A data frame with the Odds Ratios for all features with their 95% confidence intervals and a ggplot2 object.

**Author(s)**

Pol Castellano-Escuder

**Examples**

```

data("st000336")

st000336 %>%
  PomaImpute() %>%
  PomaNorm() %>%
  PomaOddsRatio(feature_name = c("glutamic_acid", "glutamine",
                                "glycine", "histidine"))

```

---

PomaOutliers

*Remove and Analyze Outliers*


---

**Description**

This function allows users to analyze outliers by different plots and remove them from an MSnSet object.

**Usage**

```

PomaOutliers(
  data,
  do = "clean",
  method = "euclidean",
  type = "median",
  coef = 1.5,
  labels = FALSE
)

```

**Arguments**

|        |  |
|--------|--|
| data   | A MSnSet object. First pData column must be the subject group/type.  |
| do     | Action to do. Options are "clean" (to remove detected outliers) and "analyze" (to analyze data outliers). Note that the output of this function will be different depending on this parameter. |
| method | Distance measure method to perform MDS. Options are "euclidean", "maximum", "manhattan", "canberra" and "minkowski". See <code>?dist()</code> .  |
| type   | Type of outliers analysis to perform. Options are "median" (default) and "centroid". See <code>vegan::betadisper</code> .  |
| coef   | This value corresponds to the classical 1.5 in $Q3 + 1.5 * IQR$ formula to detect outliers. By changing this value, the permissiveness in outlier detection will change.                       |
| labels | Logical indicating if sample IDs should to be plotted or not.  |

**Value**

A MSnSet object with cleaned data or different exploratory plots for the detailed analysis of outliers (depending on "do" parameter).

**Author(s)**

Pol Castellano-Escuder

**Examples**

```
data("st000336")

# clean outliers
st000336 %>%
  PomaImpute() %>%
  PomaNorm() %>%
  PomaOutliers()

# analyze outliers
st000336 %>%
  PomaImpute() %>%
  PomaNorm() %>%
  PomaOutliers(do = "analyze")
```

---

PomaRandForest

*Classification Random Forest for Mass Spectrometry Data*

---

**Description**

PomaRandForest() allows users to perform a classification Random Forest with a MS data matrix using the classical randomForest R package.

**Usage**

```
PomaRandForest(
  data,
  ntest = 20,
  ntree = 500,
  mtry = floor(sqrt(ncol(t(MSnbase::exprs(data))))),
  nodesize = 1,
  nvar = 20
)
```

**Arguments**

|          |   |
|----------|---|
| data     | A MSnSet object. First pData column must be the subject group/type.   |
| ntest    | Numeric indicating the percentage of observations that will be used as test set. Default is 20% of observations.                                      |
| ntree    | Number of trees to grow.  |
| mtry     | Number of variables randomly sampled as candidates at each split. This value is set $\sqrt{p}$ (where $p$ is number of variables in data) by default. |
| nodesize | Minimum size of terminal nodes. By default is equal to 1.   |
| nvar     | Number of variables to show in the Gini plot.   |

**Value**

A list with all results for Random Forest including plots and data frames.

**Author(s)**

Pol Castellano-Escuder

**References**

A. Liaw and M. Wiener (2002). Classification and Regression by randomForest. R News 2(3), 18–22.

**Examples**

```
data("st000336")

st000336 %>%
  PomaImpute() %>%
  PomaRandForest()
```

---

PomaRankProd

*Rank Product/Rank Sum Analysis for Mass Spectrometry Data*

---

**Description**

PomaRankProd() performs the Rank Product method to identify differential feature concentration/intensity.

**Usage**

```
PomaRankProd(
  data,
  logged = TRUE,
  logbase = 2,
  paired = NA,
  cutoff = 0.05,
  method = "pfp"
)
```

**Arguments**

|         |   |
|---------|---|
| data    | A MSnSet object. First pData column must be the subject group/type.   |
| logged  | If "TRUE" (default) data have been previously log transformed.  |
| logbase | Numerical. Base for log transformation.   |
| paired  | Number of random pairs generated in the function, if set to NA (default), the odd integer closer to the square of the number of replicates is used. |

|        |  |
|--------|--|
| cutoff | The pfp/pvalue threshold value used to select features.  |
| method | If cutoff is provided, the method needs to be selected to identify features. "pfp" uses percentage of false prediction, which is a default setting. "pval" uses p-values which is less stringent than pfp. |

**Value**

A list with all results for Rank Product analysis including tables and plots.

**Author(s)**

Pol Castellano-Escuder

**References**

Breitling, R., Armengaud, P., Amtmann, A., and Herzyk, P.(2004) Rank Products: A simple, yet powerful, new method to detect differentially regulated genes in replicated microarray experiments, FEBS Letter, 57383-92

Hong, F., Breitling, R., McEntee, W.C., Wittner, B.S., Nemhauser, J.L., Chory, J. (2006). RankProd: a bioconductor package for detecting differentially expressed genes in meta-analysis *Bioinformatics*. 22(22):2825-2827

Del Carratore, F., Jankevics, A., Eisinga, R., Heskes, T., Hong, F. & Breitling, R. (2017). RankProd 2.0: a refactored Bioconductor package for detecting differentially expressed features in molecular profiling datasets. *Bioinformatics*. 33(17):2774-2775

---

PomaUnivariate

*Univariate Statistical Methods for Mass Spectrometry Data*

---

**Description**

PomaUnivariate() allows users to perform different univariate statistical analysis on MS data.

**Usage**

```
PomaUnivariate(  
  data,  
  covariates = FALSE,  
  method = "ttest",  
  paired = FALSE,  
  var_equal = FALSE,  
  adjust = "fdr"  
)
```

**Arguments**

|            |  |
|------------|--|
| data       | A MSnSet object. First pData column must be the subject group/type.  |
| covariates | Logical. If it's set to TRUE all metadata variables stored in pData will be used as covariables. Default = FALSE.      |
| method     | Univariate statistical method. Options are: "ttest", "anova", "mann" and "kruskal".                                    |
| paired     | Logical that indicates if the data is paired or not.   |
| var_equal  | Logical that indicates if the data variance is equal or not.   |
| adjust     | Multiple comparisons correction method. Options are: "fdr", "holm", "hochberg", "hommel", "bonferroni", "BH" and "BY". |

**Value**

A data frame with results.

**Author(s)**

Pol Castellano-Escuder

**Examples**

```

data("st000336")
data("st000284")

# ttest
st000336 %>%
  PomaImpute() %>%
  PomaNorm() %>%
  PomaOutliers() %>%
  PomaUnivariate(method = "ttest")

# ANOVA
st000284 %>%
  PomaImpute() %>%
  PomaNorm() %>%
  PomaOutliers() %>%
  PomaUnivariate(method = "anova")

```

---

PomaVolcano

*Volcano Plot*

---

**Description**

PomaVolcano() generates a volcano plot from the PomaUnivariate(method = "ttest") result. The data can't have negative values.

**Usage**

```
PomaVolcano(  
  data,  
  pval = "raw",  
  pval_cutoff = 0.05,  
  adjust = "fdr",  
  log2FC = 0.6,  
  xlim = 2,  
  labels = FALSE,  
  paired = FALSE,  
  var_equal = FALSE,  
  interactive = FALSE,  
  plot_title = TRUE  
)
```

**Arguments**

|                          |  |
|--------------------------|--|
| <code>data</code>        | A MSnSet object. First pData column must be the subject group/type. Only for two group data!   |
| <code>pval</code>        | Select a pvalue type to generate the volcano plot. Options are: "raw" and "adjusted".  |
| <code>pval_cutoff</code> | Numeric. Define the pvalue cutoff (horizontal line).   |
| <code>adjust</code>      | Multiple comparisons correction method for t test result. Options are: "fdr", "holm", "hochberg", "hommel", "bonferroni", "BH" and "BY". |
| <code>log2FC</code>      | Numeric. Define the log2 fold change cutoff (vertical lines).  |
| <code>xlim</code>        | Numeric. Define the limits for x axis.   |
| <code>labels</code>      | Logical that indicates if selected labels will be plotted or not. Default is FALSE.  |
| <code>paired</code>      | Logical that indicates if the data is paired or not.   |
| <code>var_equal</code>   | Logical that indicates if the data variance is equal or not.   |
| <code>interactive</code> | Logical that indicates if an interactive plot will be plotted or not. Default is FALSE.  |
| <code>plot_title</code>  | Logical that indicates if title will be plotted or not. Default is TRUE.   |

**Value**

A ggplot2 object.

**Author(s)**

Pol Castellano-Escuder

**Examples**

```
data("st000336")  
  
st000336 %>%
```

```
PomaImpute() %>%  
PomaVolcano()
```

st000284

---

*Colorectal Cancer Detection Using Targeted Serum Metabolic Profiling*

---

**Description**

Colorectal cancer (CRC) is one of the most prevalent and deadly cancers in the world. Despite an expanding knowledge of its molecular pathogenesis during the past two decades, robust biomarkers to enable screening, surveillance, and therapy monitoring of CRC are still lacking. In this study, we present a targeted liquid chromatography-tandem mass spectrometry-based metabolic profiling approach for identifying biomarker candidates that could enable highly sensitive and specific CRC detection using human serum samples. In this targeted approach, 158 metabolites from 25 metabolic pathways of potential significance were monitored in 234 serum samples from three groups of patients (66 CRC patients, 76 polyp patients, and 92 healthy controls). Partial least squares-discriminant analysis (PLS-DA) models were established, which proved to be powerful for distinguishing CRC patients from both healthy controls and polyp patients. Receiver operating characteristic curves generated based on these PLS-DA models showed high sensitivities (0.96 and 0.89, respectively, for differentiating CRC patients from healthy controls or polyp patients); good specificities (0.80 and 0.88), and excellent areas under the curve (0.93 and 0.95) were also obtained. Monte Carlo cross validation (MCCV) was also applied, demonstrating the robust diagnostic power of this metabolic profiling approach.

**Usage**

st000284

**Format**

A MSnSet object: 224 samples, 113 metabolites, 4 covariables and 3 groups (CRC, Healthy and Polyp).

**metabolites** 113 serum metabolites.

**covariables** Age at consent, Gender, Smoking Condition and Alcohol Consumption.

**Source**

[https://www.metabolomicsworkbench.org/data/DRCCMetadata.php?Mode=Study&StudyID=ST000284&StudyType=MS&ResultType=1%20target=\\_blank](https://www.metabolomicsworkbench.org/data/DRCCMetadata.php?Mode=Study&StudyID=ST000284&StudyType=MS&ResultType=1%20target=_blank)

**References**

Colorectal Cancer Detection Using Targeted Serum Metabolic Profiling, J. Proteome. Res., 2014, 13, 4120-4130.



---

st000336

*Targeted LC/MS of urine from boys with DMD and controls*

---

### Description

Duchenne Muscular Dystrophy (DMD) is an X-linked recessive form of muscular dystrophy that affects males via a mutation in the gene for the muscle protein, dystrophin. Progression of the disease results in severe muscle loss, ultimately leading to paralysis and death. Steroid therapy has been a commonly employed method for reducing the severity of symptoms. This study aims to quantify the urine levels of amino acids and organic acids in patients with DMD both with and without steroid treatment. Track the progression of DMD in patients who have provided multiple urine samples.

### Usage

st000336

### Format

A MSnSet object: 57 samples, 31 metabolites, 1 covariable and 2 groups (Controls and DMD).

**metabolites** 31 urine metabolites.

**covariables** Steroid status.

### Source

<https://www.metabolomicsworkbench.org/data/DRCCMetadata.php?Mode=Study&DataMode=AllData&StudyID=ST000336&StudyType=MS&ResultType=1#DataTabs>

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