Package 'xmsPANDA'

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Type Package

for biomarker discovery and biomarker-driven network analysis for studying disease mechanisms using metabolomics
Version 1.0.7.2
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Description This package includes functions to perform biomarker discovery for classification and regression scenarios and to perform correlation based network analysis (all metabolites and/or targeted metabolites eg: glucose metabolism pathway).
License GPL2.0
LazyLoad yes
corpcor, earth, pROC, multcomp, RColorBrewer, nlme, pls, plsgenomics, igraph, ROCR, flashClust, data.table, dplyr,plyr,mclust,parallel,RankAggreg,pcaMethods R topics documented:
xmsPANDA-package
data_preprocess
diffexp
do_wgcna
get_boxplots
get_hca
get_manhattanplots
get_pca
get_roc
metabnet
Index 21

2 data_preprocess

xmsPANDA-package xmsPANDA

Description

R pacakge for biomarker discovery, supervised and unsupervised learning, and network analysis.

Details

Package: xmsPANDA
Type: Package
Version: 1.0.7.1
Date: 2018-03-23
License: gpl2.0
LazyLoad: yes

Author(s)

Karan Uppal Maintainer: <kuppal2@emory.edu>

data_preprocess data_preprocess

Description

This function performs data transformation, normalization

Usage

```
data_preprocess(Xmat=NA,Ymat=NA,feature_table_file, parentoutput_dir, class_labels_file,
num_replicates = 3,
feat.filt.thresh = NA, summarize.replicates = TRUE, summary.method = "mean",
all.missing.thresh=0.5,
group.missing.thresh = 0.7, log2transform = TRUE,
medcenter = TRUE, znormtransform = FALSE, quantile_norm = TRUE,
lowess_norm = FALSE,madscaling = FALSE, missing.val = 0, samplermindex = NA,
rep.max.missing.thresh = 0.5,summary.na.replacement = "zeros",featselmethod=NA)
```

Arguments

Xmat R object for feature table. If this is given, then feature table can be set to NA.

Ymat R object for response/class labels matrix. If this is given, then class can be set

to NA.

data_preprocess 3

feature_table_file

Feature table that includes the mz, retention time, and measured intensity in each sample for each analyte. The first 2 columns should be the mz and time. The remaining columns should correspond to the samples in the class labels file with each column including the intensity profile of a sample. Full path required. Eg: C:/My Documents/test.txt The feature table should be in a tab-delimited format. An example of the input file is provided under the "example" folder.

parentoutput_dir

Provide full path of the folder where you want the results to be written. Eg: C:/My Documents/ProjectA/results/

class_labels_file

File with class labels information for each sample. Samples should be in the same order as in the feature table. Please use the same format as in the example folder.

num_replicates Number of technical replicates

feat.filt.thresh

Percent Intensity Difference or Coefficient of variation threshold; feature filtering Use NA to skip this step.

summarize.replicates

Do the technical replicates per sample need to be averaged or median summarized?

summary.method Method for summarizing the replicates. Options: "mean" or "median" summary.na.replacement

How should the missing values be represented? Options: "zeros", "halffeaturemin", "halfsamplemin", "halfdatamin", "none" "zeros": replaces missing values by 0 "halfsamplemin": replaces missing value by one-half of the lowest signal intensity in the corresponding sample "halfdatamin": replaces missing value by one-half of the lowest signal intensity in the complete dataset "halffeaturemin": replaces missing value by one-half of the lowest signal intensity for the current feature "none": keeps missing values as NAs

Users are recommended to perform imputation prior to performing biomarker discovery.

missing.val How are the missing values represented in the input data? Options: "0" or "NA"

samplermindex Column index of any additional or irrelevant columns to be deleted. Options: "NA" or list of column numbers. eg: c(1,3,4) Default=NA

rep.max.missing.thresh

What propotion of replicates are allowed to have missing values during the averaging or median summarization step of each biological sample? If the number of replicates with missing values is greater than the defined threshold, then the summarized value is represented by the "missing.val" parameter. If the number of replicates with missing values is less than or equal to the defined threshold, then the summarized value is equal to the mean or the median of the non-missing values. Default: 0.5

all.missing.thresh

What propotion of total number of samples should have an intensity? Default: 0.5

group.missing.thresh

What propotion of samples in either of the two groups should have an intensity? If at least x for further analysis. Default: 0.7

log2transform Data transformation: Please refer to http://www.biomedcentral.com/1471-2164/7/142

Try different combinations; such as log2transform=TRUE, znormtransfrom=FALSE

or log2transform=FALSE, znormtransfrom=TRUE

medcenter Median centering of metabolites

znormtransform Auto scaling; each metabolite will have a mean of 0 and unit variance

quantile_norm Performs quantile normalization. Normalization options: Please set only one of

the options to be TRUE

lowess_norm Performs lowess normalization. Normalization options: Please set only one of

the options to be TRUE

madscaling Performs median adjusted scale normalization. Normalization options: Please

set only one of the options to be TRUE

Value

Pre-processed data matrix.

Author(s)

Karan Uppal < kuppal2@emory.edu>

diffexp diffexp

Description

This function performs biomarker discovery and generates a correlation network based on the metabolome-wide (and targeted) correlation analysis of the differentially expressed features. The "featselmethod" allows users to select the method for selecting discriminatory features. The function evaluates the k-fold cross-validation accuracy using Support Vector Machine, performs hierarchical clustering analysis, PCA analysis (R2/Q2 diagnostics), and generates boxplots for discriminatory features identified at each relative standard deviation (coefficient of variation) threshold across all samples (if one feature selection method is used). An optimization score that minimizes the number of false positives and increases the classification accuracy is used to select the best set of features. The best set is then used for correlation (complete or partial) based metabolome-wide network analysis. Additionally, users have the option to provide a list of mzs corresponding to chemicals of interest such as (phenylalanine, choline, etc). The function using the getVenn function in xMSanalyzer to find the mzs matching the target list based on a user defined mass search threshold (+/- ppm).

Usage

```
diffexp(Xmat = NA, Ymat = NA, feature_table_file, parentoutput_dir, class_labels_file,
num_replicates = 3, summarize.replicates = TRUE, summary.method = "mean",
summary.na.replacement = "zeros", missing.val = 0, rep.max.missing.thresh = 0.3,
all.missing.thresh = 0.5, group.missing.thresh = 0.8, input.intensity.scale = "raw",
log2transform = TRUE, medcenter = FALSE, znormtransform = FALSE, quantile_norm = TRUE,
lowess_norm = FALSE, madscaling = FALSE, rsd.filt.list = seq(0, 75, 5), pairedanalysis = FALSE,
featselmethod = "limma", pvalue.thresh = 0.05, fdrthresh = 0.05, fdrmethod = "BH",
cor.method = "spearman", networktype = "complete", abs.cor.thresh = 0.4, cor.fdrthresh = 0.05,
kfold = 10, pred.eval.method = "BER", globalcor = TRUE, target.metab.file = NA,
```

```
target.mzmatch.diff = 10, target.rtmatch.diff = NA, max.cor.num = 100,
numtrees = 20000, analysismode = "classification", net_node_colors = c("green", "red"),
net_legend = TRUE, svm_kernel = "radial", heatmap.col.opt = "redblue",
manhattanplot.col.opt = c("darkblue", "red3"), boxplot.col.opt = c("grey57"),
sample.col.opt = "rainbow", rf_selmethod = "absVIMthresh", pls_vip_thresh = 1,
num_nodes = 2, max_varsel = 100, pls_ncomp = 5, pca.stage2.eval = TRUE,
scoreplot_legend = TRUE, pca.global.eval = TRUE, rocfeatlist = seq(2, 11, 1),
rocfeatincrement = TRUE, rocclassifier = "svm", foldchangethresh = 2, wgcnarsdthresh = 20,
WGCNAmodules = TRUE, optselect = TRUE, max_comp_sel = 1, saveRda = TRUE,
legendlocation = "topleft", pca.cex.val = 4, pca.ellipse = FALSE, ellipse.conf.level = 0.95,
    permutations.count = 1000, svm.acc.tolerance = 5, limmadecideTests = TRUE,
pls.vip.selection = "max", globalclustering = TRUE, plots.res = 600, plots.width = 8,
    plots.height = 8, plots.type = "cairo", output.device.type = "pdf", ,
    individualsampleplot.col.opt = NA, pamr.threshold.select.max = FALSE,
    aggregation.method = "RankAggreg", ...)
```

Arguments

Xmat R object for feature table. If this is given, then feature table can be set to NA.

Ymat R object for response/class labels matrix. If this is given, then class can be set to NA.

feature_table_file

Feature table that includes the mz, retention time, and measured intensity in each sample for each analyte. The first 2 columns should be the mz and time. The remaining columns should correspond to the samples in the class labels file with each column including the intensity profile of a sample. Full path required. Eg: C:/My Documents/test.txt The feature table should be in a tab-delimited format. An example of the input file is provided under the "example" folder.

parentoutput_dir

Provide full path of the folder where you want the results to be written. Eg: C:/My Documents/ProjectA/results/

class_labels_file

File with class labels information for each sample. Samples should be in the same order as in the feature table. Please use the same format as in the example folder. If you want to adjust for covariates in "Imreg" option, then you can add additional columns, one per covariate. Categorical variables should be strings (eg: "male", "female"). Please see "classlabels_gender.txt" file as an example.

num_replicates Number of technical replicates

feat.filt.thresh

Percent Intensity Difference or Coefficient of variation threshold; feature filtering Use NA to skip this step.

summarize.replicates

Do the technical replicates per sample need to be averaged or median summarized?

summary.method Method for summarizing the replicates. Options: "mean" or "median" summary.na.replacement

How should the missing values be represented? Options: "zeros", "halffeaturemin", "halfsamplemin", "halfdatamin", "none" "zeros": replaces missing values by 0 "halfsamplemin": replaces missing value by one-half of the lowest signal intensity in the corresponding sample "halfdatamin": replaces missing

value by one-half of the lowest signal intensity in the complete dataset "halffeaturemin": replaces missing value by one-half of the lowest signal intensity for the current feature "none": keeps missing values as NAs

Users are recommended to perform imputation prior to performing biomarker discovery.

missing.val How are the missing values represented in the input data? Options: "0" or "NA" rep.max.missing.thresh

What propotion of replicates are allowed to have missing values during the averaging or median summarization step of each biological sample? If the number of replicates with missing values is greater than the defined threshold, then the summarized value is represented by the "missing.val" parameter. If the number of replicates with missing values is less than or equal to the defined threshold, then the summarized value is equal to the mean or the median of the non-missing values. Default: 0.5

all.missing.thresh

What propotion of total number of samples should have an intensity? Default: 0.5

input.intensity.scale

Are the intensities in the input feature table at raw scale or log2 scale? eg: "raw" or "log2" Default: "raw"

group.missing.thresh

What propotion of samples in either of the two groups should have an intensity? If at least x for further analysis. Default: 0.7

log2transform Data transformation: Please refer to http://www.biomedcentral.com/1471-2164/7/142

Try different combinations; such as log2transform=TRUE, znormtransfrom=FALSE

or log2transform=FALSE, znormtransfrom=TRUE

medcenter Median centering of metabolites

 ${\tt znormtransform} \ \ Auto \ scaling; \ each \ metabolite \ will \ have \ a \ mean \ of \ 0 \ and \ unit \ variance$

quantile_norm Performs quantile normalization. Normalization options: Please set only one of

the options to be TRUE

lowess_norm Performs lowess normalization. Normalization options: Please set only one of

the options to be TRUE

madscaling Performs median adjusted scale normalization. Normalization options: Please

set only one of the options to be TRUE

rsd.filt.list This parameter allows to perform feature filtering based on overall variance

(across all samples) prior to performing hypothesis testing. Eg: seq(0,30,5).

pairedanalysis Is this a paired-study design? TRUE or FALSE If samples are paired, then

the feature table and the class labels file should be organized so that the paired samples are arranged in the same order in each group. For example, the first

sample in group A and the first sample in group B should be paired.

featselmethod Options: "limma": for one-way ANOVA using LIMMA (mode=classification)

"limma2way": for two-way ANOVA using LIMMA (mode=classification) "limma1wayrepeat":

for one-way ANOVA repeated measures using LIMMA (mode=classification) "limma2wayrepeat": for two-way ANOVA repeated measures using LIMMA (mode=classification) "lm1wayanova": for one-way ANOVA using linear model (mode=classification) "lm2wayanova": for two-way ANOVA using linear model (mode=classification) "lm1wayanovarepeat": for one-way ANOVA repeated measures using linear model (mode=classification) "lm2wayanovarepeat": for two-way ANOVA repeated measures using linear model (mode=classification) "lm-reg": variable selection based on p-values calculated using a linear regression

> model; allows adjustment for covariates (mode= regression or classification) "logitreg": variable selection based on p-values calculated using a logistic regression model; allows adjustment for covariates (mode= classification) "rfesvm": uses recursive feature elimination SVM algorithm for variable selection; (mode=classification) "RF": for random forest based feature selection (mode= regression or classification) "RFconditional": for conditional random forest based feature selection (mode= regression or classification) "pamr": for prediction analysis for microarrays algorithm based on the nearest shrunken centroid method (mode= classification) "MARS": for multiple adaptive regression splines (MARS) based feature selection (mode= regression or classification) "pls": for partial least squares (PLS) based feature selection (mode= regression or classification) "spls": for sparse partial least squares (PLS) based feature selection (mode= regression or classification) "o1pls": for orthogonal partial least squares (OPLS) based feature selection (mode= regression or classification)

fdrthresh

False discovery rate threshold. Eg: 0.05

fdrmethod

Options: "BH", "ST", "Strimmer", "none" "BH": Benjamini-Hochberg (1995) (Default: more conservative than "ST" and "Strimmer") "ST": Storey & Tibshirani (Storey 2001, PNAS) algorithm implemented in the qvalue package "Strimmer": (Strimmer 2008, Bioinformatics) algorithm implemented in the fdrtool package "none": No FDR correction will be performed. fdrthresh will be treated as raw p-value cutoff

cor.method

Correlation method. Options: "pearson" or "spearman". Default: "spearman"

networktype

Options: "complete" or "GGM" "complete": performs network analysis using ordinary Pearson or Spearman correlation statistic "GGM": generates network based on partial correlation analysis using the GeneNet package

abs.cor.thresh Absolute Pearson correlation coefficient for network analysis. Default: 0.4

cor.fdrthresh

False discovery rate threshold for correlation analysis. Default: 0.05

kfold

Number of subsets in which the data should be divided for cross-validation. If kfold=10, then the data set will be divided into 10 subsets of size (N/10), where N is the total number of samples. 9 subsets are used for training and the remaining 1 is used for testing. This process is repeated 10 times and the CVaccuracy would be the mean of the classification accuracy from the 10 iterations. The same will be true for any other value of k. WARNING: The kfold value should be less than or equal to the total number of samples.

pred.eval.method

Criteria for evaluating the performance of the model. CV: Overall Cross-validation classification accuracy, balanced error rate (BER): (sum of accuracy in each class)/(number of classes) area under the curve (AUC) Eg: "CV", "BER", or "AUC". Default: "BER"

globalcor

Do you want to perform correlation analysis after biomarker discovery? Options: "TRUE" or "FALSE"

target.metab.file

File that includes the mz and/or retention time of the targeted metabolites. See

target.mzmatch.diff

+/- ppm mass tolerance for searching the target m/z in the current feature table

target.rtmatch.diff

+/- retention time tolerance for searching the target m/z in the current feature

Maximum number of correlated metabolites to be included in the network figure. max.cor.num Default: 100 pcacenter Data centering for PCA. Options: "TRUE" or "FALSE". Default=TRUE Data scaling for PCA. Options: "TRUE" or "FALSE". Default=TRUE pcascale Column index of any additional or irrelevant columns to be deleted. Options: samplermindex "NA" or list of column numbers. eg: c(1,3,4) Default=NA numtrees Number of trees to be used for random forest method. Default=500 "classification" for group-wise comparison (case vs control) or "regression" for analysismode continuous response variables. Default: "classification" net_node_colors Colors of nodes in the correlation networks. Eg: c("pink", "skyblue"), or ("red", "green") Should the network be displayed for the correlation network? eg: TRUE or net_legend **FALSE** Max number of variables to be used for sPLS, rfesvm, and Random Forest? max_var eg:150 SVM kernel eg: "radial" or "linear" svm_kernel heatmap.col.opt Color scheme for HCA hetmap eg: "RdBu", "topo", "heat", or "terrain" sample.col.opt Color scheme for PCA and heatmap sample axis eg: "rainbow", "heat" or "topo" Random forest VIP based selection method. If rankbased option is selected, rf_selmethod variables are ranked based on the Variable Importance Measure. Only the top "max_varsel" variables are selected. If absVIMthresh is selected, then all features with VIM greater than the absolute value of the lowest VIM are selected. eg: "absVIMthresh" or "rankbased" pls_vip_thresh Threshold for VIP score from PLS/O1PLS. eg: 1 max_varsel Maximum number of variables to keep if "rankbased" RF or spls is used. eg: Maximum number of components to be considered during the PLS optimal numpls_ncomp ber of components selection step. eg: 2 pca.stage2.eval Should PCA diagnostics be performed in stage 2? eg: TRUE or FALSE scoreplot_legend Should legends be included in score plots? eg: TRUE or FALSE pca.global.eval Should global PCA evaluation be performed? Default:TRUE eg: TRUE or **FALSE** rocfeatlist Vector indicating number of features to be used for ROC evaluation: eg: c(2,4,6)will generate ROC for top 2, top 4, and top 6 feautres. Default: seq(2,10,1) Classifier to be used for ROC evaluation. Options: "svm" or "logitreg". Default: rocclassifier "svm" foldchangethresh Secondary feature selection criteria based on fold change threshold. This is performed after statistical significance or importance evaluation. wgcnarsdthresh Relative standard deviation or coefficient of variation (across all samples) based

filtering threshold before performing WGCNA module preservation analysis.

Default: 20

WGCNAmodules Perform WGCNA module preservation analysis. TRUE or FALSE Default:

TRUE

optselect Determine optimal number of PLS components. Default: TRUE

max_comp_sel Number of PLS components to use for VIP or sparse loading selection (sPLS).

Default=1

saveRda Should the results be saved in a binary R object. Default: TRUE

legendlocation Legend location for PLS or PCA plots pca.cex.val Size of points on PCA plots. eg: 4

pca.ellipse Should ellipse be plotted on PCA plots? eg: TRUE or FALSE

ellipse.conf.level

Confidence interval for PCA ellipses eg: 0.95

pls.permut.count

Number of permutations for calculating p-values for PLS or sPLS models. eg:

1000

svm.acc.tolerance

Stopping criteria for forward feature selection using "rfeSVM" method. If the difference between best accuracy and current accuracy based on the newly added feature drops below the tolerance level, the forward selection process is termi-

nated. eg: 5

pamr.threshold.select.max

If two or more thresholds for shrinking the d statistic in the PAM algorithm (Tibshirani et al. Statistical Science 2003) have equal accuracy, should the maximum

value (lowest number of features) be used? Default: FALSE

aggregation.method

Method for combining the results from mutliple feature selection methods Options: Consensus: will only keep features that are selected in all methods Rank-Aggreg: will use the cross entropy algorithm with Spearman footrule distance as the distance measure (RankAggreg; Pihur et al. BMC Bioinformatics 2009)

Details

This function performs data transformation, normalization, FDR analysis using LIMMA, variable selection using random forests, evaluates the predictive accuracy of the FDR significant features using k-fold cross-validation with a Support Vector Machine classifier, performs two-way hierarchical clustering analysis, and principal component analysis. An optimization scheme is used to select the best set of features from different log2 fold change filtering thresholds. Finally, metabolome-wide and targeted correlation based network analysis of the FDR significant features is performed.

Value

diffexp_metabs Best set of discriminatory features.

all_metabs Results for all features.

mw.an.fdr Metabolome-wide significant correlation network of differentially expressed metabo-

ites.

targeted.an.fdr

Correlation network of differentially expressed metabolites with targeted metabo-

lites.

Following files are generated in the parent output location: Manhattan plots: showing metabolome wide p-values; Heatmap from Two-way hierarchical clustering analysis; Pairwise score plots from

10 do_wgcna

Principal Component Analysis; PCA score distribution plots; ROC plots; List of differentially expressed metabolites; Boxplots of differentially expressed metabolites; Correlation network figure and matrix; Pairwise correlation matrix CIRCOS format ready to be uploaded to: http://mkweb.bcgsc.ca/tableviewer/visu Or uploaded to Cytoscape gml format

Author(s)

Karan Uppal < kuppal2@emory.edu>

do_wgcna

do_wgcna

Description

This function performs module preservation analysis using WGCNA.

Usage

```
do_wgcna(feature_table_file = NA, class_labels_file = NA, X = NA,
Y = NA, sigfeats = NA)
```

Arguments

feature_table_file

Path and name of feature table that includes the mz, retention time, and measured intensity in each sample for each analyte. The first 2 columns should be the mz and time. The remaining columns should correspond to the samples in the class labels file with each column including the intensity profile of a sample. Full path required. Eg: C:/My Documents/test.txt The feature table should be in a tab-delimited format. An example of the input file is provided under the "example" folder.

class_labels_file

File with class labels information for each sample. Samples should be in the same order as in the feature table. Please use the same format as in the example folder. If you want to adjust for covariates in "Imreg" option, then you can add additional columns, one per covariate. Categorical variables should be strings (eg: "male", "female"). Please see "classlabels_gender.txt" file as an example.

X R object for feature table. If this is given, then feature table can be set to NA.

R object for response/class labels matrix. If this is given, then class labels file can be set to NA.

sigfeats List of differentially expressed features. Default: NA

Details

Υ

This function calls WGCNA to perform module preservation analysis between different classes or groups.

Value

PDF plots for module preservation from WGCNA and preservation matrix

get_boxplots 11

Author(s)

Karan Uppal

References

WGCNA (Horvath 2007)

get_boxplots

get_boxplots

Description

This function generates boxplots for m/z features. The input intensity matrix could be transformed or non-transformed intensities. Sample labels in the class labels file should be in the same order as the intensity matrix or feature table.

Usage

```
get_boxplots(feature_table_file, parentoutput_dir, class_labels_file, sample.col.opt = "rainbow",
```

Arguments

feature_table_file

Feature table that includes the mz, retention time, and measured intensity in each sample for each analyte. The first 2 columns should be the mz and time. The remaining columns should correspond to the samples in the class labels file with each column including the intensity profile of a sample. Full path required. Eg: C:/My Documents/test.txt The feature table should be in a tab-delimited format. An example of the input file is provided under the "example" folder.

parentoutput_dir

Provide full path of the folder where you want the results to be written. Eg: C:/My Documents/ProjectA/results/

class_labels_file

File with class labels information for each sample. Samples should be in the same order as in the feature table. Please use the same format as in the example folder.

sample.col.opt Color scheme for PCA and heatmap sample axis eg: "heat" or "topo"

alphacol=0.3 Color scaling parameter eg:0.3

Value

Creates a PDF with boxplots for each m/z feature.

Author(s)

Karan Uppal kuppal2@emory.edu

12 get_hca

Description

This function performs two-way hierarchical clustering analysis and generates a heatmap showing the clustering results. The input intensity matrix could be transformed or non-transformed intensities. Sample labels in the class labels file should be in the same order as the intensity matrix or feature table.

Usage

```
get_hca(feature_table_file, parentoutput_dir, class_labels_file, heatmap.col.opt = "RdBu", cor.met
is.data.znorm = FALSE, analysismode = "classification", sample.col.opt = "rainbow", plots.width = 20
plots.height = 2000, plots.res = 300, alphacol = 0.3,hca_type="two-way")
```

Arguments

feature_table_file

Feature table that includes the mz, retention time, and measured intensity in each sample for each analyte. The first 2 columns should be the mz and time. The remaining columns should correspond to the samples in the class labels file with each column including the intensity profile of a sample. Full path required. Eg: C:/My Documents/test.txt The feature table should be in a tab-delimited format. An example of the input file is provided under the "example" folder.

parentoutput_dir

Provide full path of the folder where you want the results to be written. Eg: C:/My Documents/ProjectA/results/

class_labels_file

File with class labels information for each sample. Samples should be in the same order as in the feature table. Please use the same format as in the example folder.

heatmap.col.opt

Color scheme for HCA hetmap eg: "RdBu", "topo", "heat", or "terrain"

cor.method Correlation method. Options: "person" or "spearman". Default: "spearman"

is.data.znorm Is the data already auto-scaled or z-scaled? eg: TRUE or FALSE

analysismode "classification" for group-wise comparison (case vs control) or "regression" for

continuous response variables. Default: "classification"

sample.col.opt Color scheme for PCA and heatmap sample axis eg: "heat" or "topo" or "rain-

how"

plots.width Width of the tiff file. eg: 2000
plots.height Height of the tiff file. eg: 2000
plots.res Resolution of the tiff file. eg: 300
alphacol Color scaling parameter eg:0.3

hcatype Color scaling parameter eg:"two-way" or "one-way"

get_manhattanplots 13

Value

Heatmap from Two-way hierarchical clustering analysis; Intensity matrix in the same order as the dendrograms in heatmap; Sample cluster labels

Author(s)

Karan Uppal < kuppal2@emory.edu>

get_manhattanplots get_manhattanplots

Description

Function to generate Manhattan plots.

Usage

```
get_manhattanplots(xvec, yvec, up_or_down, maintext = "", ythresh = 0.05,
ylab, xlab, colorvec = c("darkgreen", "firebrick1"),
col_seq = c("brown", "chocolate3", "orange3", "coral", "pink", "skyblue",
"blue", "darkblue", "purple", "violet"), xincrement = 150, yincrement = 1)
```

Arguments

xvec	Vector with values for the x-axis. eg: m/z or retention time values
yvec	Vector with values for the y-axis. eg: (-)Log10 of p-values, VIP, loadings, regression coefficients, etc.
up_or_down	Vector indicating directionality of change. eg: Fold change values
maintext	Text for the plot title
ythresh	Y-axis threshold for significance or differential expression. eg: 3 for p=0.001; $y=(-1)*log10(0.001)$ or 2 for VIP from PLS
ylab	Y-axis label
xlab	X-axis label
colorvec	Vector of colors for representing up-regulation and down-regulation. eg: c("darkgreen", "firebrick1") In this case, features that are up-regulated in class A will have "darkgreen" color, and features that are up-regulated in class B will have "firebrick1" color.
col_seq	Vector of colors for plotting different segments of the x-axis
xincrement	Window size for breaking the x-axis into different segments for visualization purposes. eg: 150
yincrement	Window size for breaking the y-axis into different segments for visualization purposes. eg: 1

Details

This function can be used to generate Type 1 Manhattan plots: significance vs m/z Type 2 Manhattan plots: significance vs retention time Type 3 Manhattan plots: significance vs intensity

14 get_pca

Value

Manhattan plots

Note

#Example pdf("Manhattanplot.pdf") get_manhattanplots(...) #pass arguments dev.off()

Author(s)

Karan Uppal

get_pca

Perfors PCA analysis

Description

This function uses the pca function implemented in the mixOmics package for PCA analysis

Usage

```
get_pca(X, samplelabels, legendlocation = "topright", filename = NA,
ncomp = 5, center = TRUE, scale = TRUE, legendcex = 0.5,
outloc = getwd(), col_vec = NA, sample.col.opt = "default",
alphacol = 0.3, class_levels = NA, pca.cex.val = 3,
pca.ellipse = TRUE, ellipse.conf.level = 0.5, samplenames = FALSE)
```

Arguments

samplenames

Χ	Data matrix without m/z and time.
samplelabels	Vector with class label for each sample.
legendlocation	Location of the legend on PCA score plots
filename	eg: "all", "signficantfeats"
ncomp	Number of components; please use ?pca for more information
center	Should the data be centered?; please use ?pca for more information
scale	Should the data be scaled?; please use ?pca for more information
legendcex	Size of the legend text in the PCA score plots. e.g.: 0.5 or 0.7
outloc	Output folder location
col_vec	Provide vector of colors for each group. eg: NA or c("red", "green") for cases and controls, respectively. This argument is ignored if sample.col.opt is provided
sample.col.opt	Select R color palette. eg: "rainbow", "terrain", "topo". "heat", "default"
alphacol	Semi-transparent colors eg: 0.2
class_levels	Vector with names of different sample groups. eg: c("case", "control") or NA
pca.cex.val	Size of dots in PCA score plots. eg: 0.4
pca.ellipse	Should the score confidence interval for each group be drawn? eg: TRUE or FALSE
ellipse.conf.le	evel
	Confidence interval level eg: 0.95

Should the sample names be included in PCA plots? eg: TRUE or FALSE

get_pcascoredistplots 15

Details

This function performs PCA analysis. The results are saved in a RDA file.

Value

The function returns PCA results as an object and generates pairwise score plots for the first three components

Author(s)

Karan Uppal

References

mixOmics

```
get_pcascoredistplots get pcascoredistplots
```

Description

PCA score distribution (25th percentile, median, 75th percentile) plots

Usage

```
get_pcascoredistplots(X = NA, Y = NA, feature_table_file, parentoutput_dir,
class_labels_file, sample.col.opt = "rainbow", plots.width = 2000,
plots.height = 2000, plots.res = 300, alphacol = 0.3, col_vec,
pairedanalysis = FALSE, pca.cex.val = 3, legendlocation = "topright",
pca.ellipse = TRUE, ellipse.conf.level = 0.5, filename = "all")
```

Arguments

X R object for feature table. If this is given, then feature table can be set to NA.

Y R object for response/class labels matrix. If this is given, then class labels file can be set to NA.

feature_table_file

Path and name of feature table that includes the mz, retention time, and measured intensity in each sample for each analyte. The first 2 columns should be the mz and time. The remaining columns should correspond to the samples in the class labels file with each column including the intensity profile of a sample. Full path required. Eg: C:/My Documents/test.txt The feature table should be in a tab-delimited format. An example of the input file is provided under the "example" folder.

parentoutput_dir

Provide full path of the folder where you want the results to be written. Eg: C:/My Documents/ProjectA/results/

16 get_roc

class_labels_file

File with class labels information for each sample. Samples should be in the same order as in the feature table. Please use the same format as in the example folder. If you want to adjust for covariates in "Imreg" option, then you can add additional columns, one per covariate. Categorical variables should be strings (eg: "male", "female"). Please see "classlabels_gender.txt" file as an example.

sample.col.opt Color scheme for PCA and heatmap sample axis eg: "rainbow", "heat" or "topo"

alphacol Color scaling parameter eg:0.3 col_vec Vector of colors for each sample.

pairedanalysis Is this a paired-study design? TRUE or FALSE If samples are paired, then

the feature table and the class labels file should be organized so that the paired samples are arranged in the same order in each group. For example, the first

sample in group A and the first sample in group B should be paired.

pca.cex.val Size of points on PCA plots. eg: 4

legendlocation Legend location on PCA plots

pca.ellipse Should ellipse be plotted on PCA plots? eg: TRUE or FALSE

ellipse.conf.level

Confidence interval for PCA ellipses eg: 0.95

filename Name of output PDF file

Details

This function performs PCA and generates pariwise score plots as well as score distribution plots (per group). It uses the Y vector and classlabels for color coding the samples in the pairwise score plots.

Value

The output includes: Pairwise PCA score plots, PCA score distribution plots, PCA scores and loadings text files.

Note

The plots can be sent to an external device by running the following commands: pdf("get_pcascoredistplots.pdf") get_pcascoredistplots(...) dev.off()

Author(s)

Karan Uppal

get_roc get_roc

Description

This function generates Receiver Operating Characteristic curves using SVM and Logistic Regression as classifiers.

Usage

```
get_roc(dataA, classlabels, classifier = "svm", kname = "radial",
rocfeatlist = seq(2, 10, 1), rocfeatincrement = TRUE,
testset = NA, testclasslabels = NA, mainlabel = NA)
```

Arguments

dataA R object for feature table with only differentially expressed features. This is the

training set.

classlabels Class labels vector.

classifier Classification algorithm to be used for ROC analysis. svm: Support Vector

Machine logitreg: Logistic Regression eg: "svm" or "logitreg"

kname Kernel for SVM. eg: "radial"

rocfeatlist Vector indicating number of features to be used for ROC evaluation: eg: c(2,4,6)

will generate ROC for top 2, top 4, and top 6 feautres. Default: seq(2,10,1)

rocfeatincrement

Turns on or off forward selection. eg: TRUE or FALSE

testset R object for test feature table with only differentially expressed features. This is

the test set.

testclasslabels

Class labels vector for samples in the test set.

mainlabel Main text label for the ROC plot. eg: "Group A vs B ROC curve"

Details

Function to perform ROC curve analysis using only training set or using both training and test set.

Value

PDF file with ROC plot

Description

Function for correlation (complete or partial) based metabolome-wide network analysis. Additionally, users have the option to provide a matrix of m/z features corresponding to chemicals of interest such as (phenylalanine, choline, etc) and/or a matrix of m/z features corresponding to discriminatory metabolites.

Usage

```
metabnet(feature_table_file, target.metab.file, sig.metab.file, class_labels_file=NA, parentoutput_conum_replicates=3, cor.method="spearman", abs.cor.thresh=0.4,
cor.fdrthresh=0.05, target.mzmatch.diff=10, target.rtmatch.diff=NA,
max.cor.num=100, feat.filt.thresh=NA, summarize.replicates=TRUE,
summary.method="mean", all.missing.thresh=0.5,
group.missing.thresh=0.7,
```

```
log2transform=TRUE,medcenter=TRUE,znormtransform=FALSE,
quantile_norm=TRUE,lowess_norm=FALSE,madscaling=FALSE,
missing.val=0, networktype="complete", samplermindex=NA,
rep.max.missing.thresh=0.3,summary.na.replacement="zeros",
net_node_colors = c("pink", "skyblue"),
net_legend = FALSE)
```

Arguments

feature_table_file

Feature table that includes the mz, retention time, and measured intensity in each sample for each analyte. The first 2 columns should be the mz and time. The remaining columns should correspond to the samples in the class labels file with each column including the intensity profile of a sample. Full path required. Eg: C:/My Documents/test.txt The feature table should be in a tab-delimited format. An example of the input file is provided under the "example" folder.

target.metab.file

File that includes the mz and/or retention time of the targeted metabolites corresponding to pathways or chemicals of interest. See example.

sig.metab.file File that includes the mz and/or retention time of the discriminatory metabolites. See example.

class_labels_file

File with class labels information for each sample. Samples should be in the same order as in the feature table. Please use the same format as in the example folder.

parentoutput_dir

Provide full path of the folder where you want the results to be written. Eg: C:/My Documents/ProjectA/results/

num_replicates Number of technical replicates

cor.method Correlation method. Options: "pearson" or "spearman". Default: "spearman"

abs.cor.thresh Absolute Pearson correlation coefficient for network analysis. Eg: 0.5

cor.fdrthresh False discovery rate threshold for correlation analysis. Eg: 0.05

target.mzmatch.diff

+/- ppm mass tolerance for searching the target m/z in the current feature table

target.rtmatch.diff

+/- retention time tolerance for searching the target m/z in the current feature table

max.cor.num Maximum number of correlated metabolites to be included in the network figure.

Default: 100

feat.filt.thresh

Percent Intensity Difference or Coefficient of variation threshold; feature filtering Use NA to skip this step.

summarize.replicates

Do the technical replicates per sample need to be averaged or median summarized?

summary.method Method for summarizing the replicates. Options: "mean" or "median" summary.na.replacement

How should the missing values be represented? Options: "zeros", "halfsamplemin", "halfdatamin", "none" "zeros": replaces missing values by 0 "halfsamplemin": replaces missing value by one-half of the lowest signal intensity in the

corresponding sample "halfdatamin": replaces missing value by one-half of the lowest signal intensity in the complete dataset "none": keeps missing values as NAs

Users are recommended to perform imputation prior to performing biomarker discovery.

all.missing.thresh

What propotion of total number of samples should have an intensity? Default: 0.5

group.missing.thresh

What propotion of samples in either of the two groups should have an intensity? If at least x for further analysis. Default: 0.7

log2transform Data transformation: Please refer to http://www.biomedcentral.com/1471-2164/7/142

 $Try\ different\ combinations; such\ as\ log2 transform = TRUE,\ znorm transfrom = FALSE$

or log2transform=FALSE, znormtransfrom=TRUE

medcenter Median centering of metabolites

znormtransform Auto scaling; each metabolite will have a mean of 0 and unit variance

quantile_norm Performs quantile normalization. Normalization options: Please set only one of

the options to be TRUE

lowess_norm Performs lowess normalization. Normalization options: Please set only one of

the options to be TRUE

madscaling Performs median adjusted scale normalization. Normalization options: Please

set only one of the options to be TRUE

missing.val How are the missing values represented in the input data? Options: "0" or "NA"

networktype Options: "complete" or "GGM" "complete": performs network analysis using

ordinary Pearson or Spearman correlation statistic "GGM": generates network

based on partial correlation analysis using the GeneNet package

samplermindex Column index of any additional or irrelevant columns to be deleted. Options:

"NA" or list of column numbers. eg: c(1,3,4) Default=NA

rep.max.missing.thresh

What propotion of replicates are allowed to have missing values during the averaging or median summarization step of each biological sample? If the number of replicates with missing values is greater than the defined threshold, then the summarized value is represented by the "missing.val" parameter. If the number of replicates with missing values is less than or equal to the defined threshold, then the summarized value is equal to the mean or the median of the non-missing

values. Default: 0.5

net_node_colors

Colors of nodes in the correlation networks. Eg: c("pink", "skyblue"), or ("red", "green")

net_legend Should the network be displayed for the correlation network? eg: TRUE or

FALSE

Details

Function for metabolomic network analysis

Value

Correlation matrix and network of metabolites.

Author(s)

Karan Uppal <kuppal2@emory.edu>

Index

```
*Topic PCA
    {\tt get\_pcascoredistplots}, 15
*Topic \textasciitildePCA
    get_pca, 14
*Topic \textasciitildeWGCNA
    do\_wgcna,\, \textcolor{red}{10}
*Topic xmsPANDA
    xmsPANDA-package, 2
data_preprocess, 2
diffexp, 4
do\_wgcna,\, \textcolor{red}{10}
get\_boxplots, 11
get_hca, 12
get_manhattanplots, 13
get_pca, 14
get\_pcascoredistplots, 15
get_roc, 16
metabnet, 17
xmsPANDA (xmsPANDA-package), 2
xmsPANDA-package, 2
```