Package 'MSstats'

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Description A set of tools for statistical relative protein significance analysis in DDA, SRM and DIA experiments.

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Description

Check if data represents repeated measurements design

Usage

checkRepeatedDesign(summarization_output)

Arguments

```
summarization_output of the dataProcess function
```

Details

This extracts information required by the group comparison workflow

Value

logical, TRUE if data represent repeated measurements design

```
QuantData1 <- dataProcess(SRMRawData, use_log_file = FALSE)
checkRepeatedDesign(QuantData1)</pre>
```

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dataProcess

Process MS data: clean, normalize and summarize before differential analysis

Description

Process MS data: clean, normalize and summarize before differential analysis

Usage

```
dataProcess(
  raw,
  logTrans = 2,
  normalization = "equalizeMedians",
  nameStandards = NULL,
  featureSubset = "all"
  remove_uninformative_feature_outlier = FALSE,
 min_feature_count = 2,
  n_{top_feature} = 3,
  summaryMethod = "TMP".
  equalFeatureVar = TRUE,
  censoredInt = "NA",
  MBimpute = TRUE,
  remove50missing = FALSE,
  fix_missing = NULL,
  maxQuantileforCensored = 0.999,
  use_log_file = TRUE,
  append = FALSE,
  verbose = TRUE,
  log_file_path = NULL
)
```

Arguments

raw name of the raw (input) data set.

logTrans base of logarithm transformation: 2 (default) or 10.

normalization normalization to remove systematic bias between MS runs. There are three

different normalizations supported: 'equalizeMedians' (default) represents constant normalization (equalizing the medians) based on reference signals is performed. 'quantile' represents quantile normalization based on reference signals 'globalStandards' represents normalization with global standards proteins. If

FALSE, no normalization is performed.

nameStandards optional vector of global standard peptide names. Required only for normaliza-

tion with global standard peptides.

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featureSubset

"all" (default) uses all features that the data set has. "top3" uses top 3 features which have highest average of log-intensity across runs. "topN" uses top N features which has highest average of log-intensity across runs. It needs the input for n_top_feature option. "highQuality" flags uninformative feature and outliers.

remove_uninformative_feature_outlier

optional. Only required if featureSubset = "highQuality". TRUE allows to remove 1) noisy features (flagged in the column feature quality with "Uninformative"), 2) outliers (flagged in the column, is_outlier with TRUE, before run-level summarization. FALSE (default) uses all features and intensities for run-level summarization.

min_feature_count

optional. Only required if featureSubset = "highQuality". Defines a minimum number of informative features a protein needs to be considered in the feature selection algorithm.

optional. Only required if featureSubset = 'topN'. It that case, it specifies numn_top_feature ber of top features that will be used. Default is 3, which means to use top 3

features.

"TMP" (default) means Tukey's median polish, which is robust estimation method. summaryMethod

"linear" uses linear mixed model.

equalFeatureVar

only for summaryMethod = "linear". default is TRUE. Logical variable for whether the model should account for heterogeneous variation among intensities from different features. Default is TRUE, which assume equal variance among intensities from features. FALSE means that we cannot assume equal variance among intensities from features, then we will account for heterogeneous varia-

tion from different features.

censoredInt Missing values are censored or at random. 'NA' (default) assumes that all 'NA's

> in 'Intensity' column are censored. '0' uses zero intensities as censored intensity. In this case, NA intensities are missing at random. The output from Skyline should use '0'. Null assumes that all NA intensites are randomly missing.

only for summaryMethod = "TMP" and censoredInt = 'NA' or '0'. TRUE (de-**MBimpute**

fault) imputes 'NA' or '0' (depending on censoredInt option) by Accelated fail-

ure model. FALSE uses the values assigned by cutoffCensored.

remove50missing

only for summaryMethod = "TMP". TRUE removes the runs which have more than 50% missing values. FALSE is default.

fix_missing Optional, same as the 'fix missing' parameter in MSstatsConvert::MSstatsBalancedDesign

function

maxQuantileforCensored

Maximum quantile for deciding censored missing values, default is 0.999

use_log_file logical. If TRUE, information about data processing will be saved to a file.

logical. If TRUE, information about data processing will be added to an existing append

log file.

verbose logical. If TRUE, information about data processing wil be printed to the con-

sole.

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log_file_path character. Path to a file to which information about data processing will be saved. If not provided, such a file will be created automatically. If 'append = TRUE', has to be a valid path to a file.

Examples

```
# Consider a raw data (i.e. SRMRawData) for a label-based SRM experiment from a yeast study
# with ten time points (T1-T10) of interests and three biological replicates.
# It is a time course experiment. The goal is to detect protein abundance changes
# across time points.
head(SRMRawData)
# Log2 transformation and normalization are applied (default)
QuantData<-dataProcess(SRMRawData, use_log_file = FALSE)
head(QuantData$FeatureLevelData)
# Log10 transformation and normalization are applied
QuantData1<-dataProcess(SRMRawData, logTrans=10, use_log_file = FALSE)
head(QuantData1$FeatureLevelData)
# Log2 transformation and no normalization are applied
QuantData2<-dataProcess(SRMRawData,normalization=FALSE, use_log_file = FALSE)
head(QuantData2$FeatureLevelData)</pre>
```

dataProcessPlots

Visualization for explanatory data analysis

Description

To illustrate the quantitative data after data-preprocessing and quality control of MS runs, data-aProcessPlots takes the quantitative data from function (dataProcess) as input and automatically generate three types of figures in pdf files as output: (1) profile plot (specify "ProfilePlot" in option type), to identify the potential sources of variation for each protein; (2) quality control plot (specify "QCPlot" in option type), to evaluate the systematic bias between MS runs; (3) mean plot for conditions (specify "ConditionPlot" in option type), to illustrate mean and variability of each condition per protein.

Usage

```
dataProcessPlots(
  data,
  type,
  featureName = "Transition",
  ylimUp = FALSE,
  ylimDown = FALSE,
  scale = FALSE,
  interval = "CI",
  x.axis.size = 10,
  y.axis.size = 10,
  text.size = 4,
```

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```
text.angle = 0,
legend.size = 7,
dot.size.profile = 2,
dot.size.condition = 3,
width = 10,
height = 10,
which.Protein = "all",
originalPlot = TRUE,
summaryPlot = TRUE,
save_condition_plot_result = FALSE,
remove_uninformative_feature_outlier = FALSE,
address = ""
```

Arguments

data name of the (output of dataProcess function) data set.

type choice of visualization. "ProfilePlot" represents profile plot of log intensities

across MS runs. "QCPlot" represents quality control plot of log intensities across MS runs. "ConditionPlot" represents mean plot of log ratios (Light/Heavy)

across conditions.

featureName for "ProfilePlot" only, "Transition" (default) means printing feature legend in

transition-level; "Peptide" means printing feature legend in peptide-level; "NA"

means no feature legend printing.

ylimUp upper limit for y-axis in the log scale. FALSE(Default) for Profile Plot and

QC Plot use the upper limit as rounded off maximum of log2(intensities) after normalization + 3. FALSE(Default) for Condition Plot is maximum of log ratio

+ SD or CI.

ylimDown lower limit for y-axis in the log scale. FALSE(Default) for Profile Plot and QC

Plot is 0. FALSE(Default) for Condition Plot is minumum of log ratio - SD or

CI.

scale for "ConditionPlot" only, FALSE(default) means each conditional level is not

scaled at x-axis according to its actual value (equal space at x-axis). TRUE means each conditional level is scaled at x-axis according to its actual value

(unequal space at x-axis).

interval for "ConditionPlot" only, "CI"(default) uses confidence interval with 0.95 sig-

nificant level for the width of error bar. "SD" uses standard deviation for the

width of error bar.

x.axis.size size of x-axis labeling for "Run" in Profile Plot and QC Plot, and "Condition" in

Condition Plot. Default is 10.

y.axis.size size of y-axis labels. Default is 10.

text.size size of labels represented each condition at the top of graph in Profile Plot and

QC plot. Default is 4.

text.angle angle of labels represented each condition at the top of graph in Profile Plot and

QC plot or x-axis labeling in Condition plot. Default is 0.

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legend.size size of feature legend (transition-level or peptide-level) above graph in Profile

Plot. Default is 7.

dot.size.profile

size of dots in profile plot. Default is 2.

dot.size.condition

size of dots in condition plot. Default is 3.

width width of the saved file. Default is 10. height height of the saved file. Default is 10.

which.Protein Protein list to draw plots. List can be names of Proteins or order numbers of

Proteins from levels(data\$FeatureLevelData\$PROTEIN). Default is "all", which generates all plots for each protein. For QC plot, "allonly" will generate one QC

plot with all proteins.

originalPlot TRUE(default) draws original profile plots.

summaryPlot TRUE(default) draws profile plots with summarization for run levels.

save_condition_plot_result

TRUE saves the table with values using condition plots. Default is FALSE.

remove_uninformative_feature_outlier

 $It only works \ after users \ used \ feature Subset="highQuality" in \ data Process. \ TRUE$

allows to remove 1) the features are flagged in the column, feature_quality="Uninformative"

which are features with bad quality, 2) outliers that are flagged in the column, is_outlier=TRUE in Profile plots. FALSE (default) shows all features and inten-

sities in profile plots.

address the name of folder that will store the results. Default folder is the current work-

ing directory. The other assigned folder has to be existed under the current working directory. An output pdf file is automatically created with the default name of "ProfilePlot.pdf" or "QCplot.pdf" or "ConditionPlot.pdf" or "ConditionPlot_value.csv". The command address can help to specify where to store the file as well as how to modify the beginning of the file name. If address=FALSE,

plot will be not saved as pdf file but showed in window.

Details

- Profile Plot: identify the potential sources of variation of each protein. QuantData\$FeatureLevelData is used for plots. X-axis is run. Y-axis is log-intensities of transitions. Reference/endogenous signals are in the left/right panel. Line colors indicate peptides and line types indicate transitions. In summarization plots, gray dots and lines are the same as original profile plots with QuantData\$FeatureLevelData. Dark dots and lines are for summarized intensities from QuantData\$ProteinLevelData.
- QC Plot: illustrate the systematic bias between MS runs. After normalization, the reference signals for all proteins should be stable across MS runs. QuantData\$FeatureLevelData is used for plots. X-axis is run. Y-axis is log-intensities of transition. Reference/endogenous signals are in the left/right panel. The pdf file contains (1) QC plot for all proteins and (2) QC plots for each protein separately.
- Condition Plot: illustrate the systematic difference between conditions. Summarized intensities from QuantData\$ProteinLevelData are used for plots. X-axis is condition. Y-axis is summarized log transformed intensity. If scale is TRUE, the levels of conditions is scaled

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according to its actual values at x-axis. Red points indicate the mean for each condition. If interval is "CI", blue error bars indicate the confidence interval with 0.95 significant level for each condition. If interval is "SD", blue error bars indicate the standard deviation for each condition. The interval is not related with model-based analysis.

The input of this function is the quantitative data from function dataProcess.

Examples

```
# Consider quantitative data (i.e. QuantData) from a yeast study with ten time points of interests,
# three biological replicates, and no technical replicates which is a time-course experiment.
# The goal is to provide pre-analysis visualization by automatically generate two types of figures
# in two separate pdf files.
# Protein IDHC (gene name IDP2) is differentially expressed in time point 1 and time point 7,
# whereas, Protein PMG2 (gene name GPM2) is not.

QuantData<-dataProcess(SRMRawData, use_log_file = FALSE)
head(QuantData$FeatureLevelData)
# Profile plot
dataProcessPlots(data=QuantData,type="ProfilePlot")
# Quality control plot
dataProcessPlots(data=QuantData,type="QCPlot")
# Quantification plot for conditions
dataProcessPlots(data=QuantData,type="ConditionPlot")</pre>
```

DDARawData

Example dataset from a label-free DDA, a controlled spike-in experiment.

Description

This is a data set obtained from a published study (Mueller, et. al, 2007). A controlled spike-in experiment, where 6 proteins, (horse myoglobin, bovine carbonic anhydrase, horse Cytochrome C, chicken lysozyme, yeast alcohol dehydrogenase, rabbit aldolase A) were spiked into a complex background in known concentrations in a latin square design. The experiment contained 6 mixtures, and each mixture was analyzed in label-free LC-MS mode with 3 technical replicates (resulting in the total of 18 runs). Each protein was represented by 7-21 peptides, and each peptide was represented by 1-5 transition.

Usage

DDARawData

Format

data.frame

Details

The raw data (input data for MSstats) is required to contain variable of ProteinName, PeptideSequence, PrecursorCharge, FragmentIon, ProductCharge, IsotopeLabelType, Condition, BioReplicate, Run, Intensity. The variable names should be fixed.

If the information of one or more columns is not available for the original raw data, please retain the column variables and type in fixed value. For example, the original raw data does not contain the information of PrecursorCharge and ProductCharge, we retain the column PrecursorCharge and ProductCharge and then type in NA for all transitions in RawData.

Variable Intensity is required to be original signal without any log transformation and can be specified as the peak of height or the peak of area under curve.

Value

data.frame with the required format of MSstats.

Author(s)

Meena Choi, Olga Vitek.

Maintainer: Meena Choi (<mnchoi67@gmail.com>)

References

Meena Choi, Ching-Yun Chang, Timothy Clough, Daniel Broudy, Trevor Killeen, Brendan MacLean and Olga Vitek. "MSstats: an R package for statistical analysis of quantitative mass spectrometry-based proteomic experiments" *Bioinformatics*, 30(17):1514-1526, 2014.

Timothy Clough, Safia Thaminy, Susanne Ragg, Ruedi Aebersold, Olga Vitek. "Statistical protein quantification and significance analysis in label-free LC-M experiments with complex designs" *BMC Bioinformatics*, 13:S16, 2012.

Mueller, L. N., Rinner, O., Schmidt, A., Letarte, S., Bodenmiller, B., Brusniak, M., Vitek, O., Aebersold, R., and Muller, M. (2007). SuperHirn - a novel tool for high resolution LC-MS based peptide/protein profiling. Proteomics, 7, 3470-3480. 3, 34

Examples

head(DDARawData)

DDARawData.Skyline

Example dataset from a label-free DDA, a controlled spike-in experiment, processed by Skyline.

Description

This is a data set obtained from a published study (Mueller, et. al, 2007). A controlled spike-in experiment, where 6 proteins, (horse myoglobin, bovine carbonic anhydrase, horse Cytochrome C, chicken lysozyme, yeast alcohol dehydrogenase, rabbit aldolase A) were spiked into a complex background in known concentrations in a latin square design. The experiment contained 6 mixtures, and each mixture was analyzed in label-free LC-MS mode with 3 technical replicates (resulting in the total of 18 runs). Each protein was represented by 7-21 peptides, and each peptide was represented by 1-5 transition. Skyline is used for processing.

Usage

DDARawData.Skyline

Format

data.frame

Details

The raw data (input data for MSstats) is required to contain variable of ProteinName, PeptideSequence, PrecursorCharge, FragmentIon, ProductCharge, IsotopeLabelType, Condition, BioReplicate, Run, Intensity. The variable names should be fixed.

This is 'MSstats input' format from Skyline used by 'MSstats_report.skyr'. The column names, 'FileName' and 'Area', should be changed to 'Run' and 'Intensity'. There are two extra columns called 'StandardType' and 'Truncated'.'StandardType' column can be used for normalization='globalStandard' in dataProcess. 'Truncated' columns can be used to remove the truncated peaks with skylineReport=TRUE in dataProcess.

If the information of one or more columns is not available for the original raw data, please retain the column variables and type in fixed value. For example, the original raw data does not contain the information of PrecursorCharge and ProductCharge, we retain the column PrecursorCharge and ProductCharge and then type in NA for all transitions in RawData.

Variable Intensity is required to be original signal without any log transformation and can be specified as the peak of height or the peak of area under curve.

Value

data.frame with the required format of MSstats.

Author(s)

Meena Choi, Olga Vitek.

Maintainer: Meena Choi (<mnchoi67@gmail.com>)

References

Meena Choi, Ching-Yun Chang, Timothy Clough, Daniel Broudy, Trevor Killeen, Brendan MacLean and Olga Vitek. "MSstats: an R package for statistical analysis of quantitative mass spectrometry-based proteomic experiments" *Bioinformatics*, 30(17):1514-1526, 2014.

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Timothy Clough, Safia Thaminy, Susanne Ragg, Ruedi Aebersold, Olga Vitek. "Statistical protein quantification and significance analysis in label-free LC-M experiments with complex designs" *BMC Bioinformatics*, 13:S16, 2012.

Examples

```
head(DDARawData.Skyline)
```

designSampleSize

Planning future experimental designs of Selected Reaction Monitoring (SRM), Data-Dependent Acquisition (DDA or shotgun), and Data-Independent Acquisition (DIA or SWATH-MS) experiments in sample size calculation

Description

Calculate sample size for future experiments of a Selected Reaction Monitoring (SRM), Data-Dependent Acquisition (DDA or shotgun), and Data-Independent Acquisition (DIA or SWATH-MS) experiment based on intensity-based linear model. Two options of the calculation: (1) number of biological replicates per condition, (2) power.

Usage

```
designSampleSize(
  data,
  desiredFC,
  FDR = 0.05,
  numSample = TRUE,
  power = 0.9,
  use_log_file = TRUE,
  append = FALSE,
  verbose = TRUE,
  log_file_path = NULL
)
```

Arguments

data FittedModel	in testing output from	function groupComparison.
------------------	------------------------	---------------------------

desiredFC the range of a desired fold change which includes the lower and upper values of

the desired fold change.

FDR a pre-specified false discovery ratio (FDR) to control the overall false positive

rate. Default is 0.05

numSample minimal number of biological replicates per condition. TRUE represents you

require to calculate the sample size for this category, else you should input the

exact number of biological replicates.

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a pre-specified statistical power which defined as the probability of detecting a true fold change. TRUE represent you require to calculate the power for this category, else you should input the average of power you expect. Default is 0.9 use_log_file logical. If TRUE, information about data processing will be saved to a file.

append logical. If TRUE, information about data processing will be added to an existing log file.

verbose logical. If TRUE, information about data processing wil be printed to the console.

log_file_path character. Path to a file to which information about data processing will be saved. If not provided, such a file will be created automatically. If 'append = TRUE', has to be a valid path to a file.

TRUE, has to be a valid path to a me

Details

The function fits the model and uses variance components to calculate sample size. The underlying model fitting with intensity-based linear model with technical MS run replication. Estimated sample size is rounded to 0 decimal. The function can only obtain either one of the categories of the sample size calculation (numSample, numPep, numTran, power) at the same time.

Value

data.frame - sample size calculation results including varibles: desiredFC, numSample, FDR, and power.

Author(s)

Meena Choi, Ching-Yun Chang, Olga Vitek.

```
# Consider quantitative data (i.e. QuantData) from yeast study.
# A time course study with ten time points of interests and three biological replicates.
QuantData <- dataProcess(SRMRawData)</pre>
head(QuantData$FeatureLevelData)
## based on multiple comparisons (T1 vs T3; T1 vs T7; T1 vs T9)
comparison1<-matrix(c(-1,0,1,0,0,0,0,0,0,0), nrow=1)
comparison2<-matrix(c(-1,0,0,0,0,0,1,0,0,0), nrow=1)
comparison3<-matrix(c(-1,0,0,0,0,0,0,0,1,0), nrow=1)
comparison<-rbind(comparison1,comparison2, comparison3)</pre>
row.names(comparison)<-c("T3-T1","T7-T1","T9-T1")
colnames(comparison)<-unique(QuantData$ProteinLevelData$GROUP)</pre>
testResultMultiComparisons<-groupComparison(contrast.matrix=comparison,data=QuantData)
## Calculate sample size for future experiments:
#(1) Minimal number of biological replicates per condition
designSampleSize(data=testResultMultiComparisons$FittedModel, numSample=TRUE,
                 desiredFC=c(1.25,1.75), FDR=0.05, power=0.8)
#(2) Power calculation
designSampleSize(data=testResultMultiComparisons$FittedModel, numSample=2,
```

```
desiredFC=c(1.25,1.75), FDR=0.05, power=TRUE)
```

designSampleSizePlots Visualization for sample size calculation

Description

To illustrate the relationship of desired fold change and the calculated minimal number sample size which are (1) number of biological replicates per condition, (2) number of peptides per protein, (3) number of transitions per peptide, and (4) power. The input is the result from function (designSampleSize.

Usage

```
designSampleSizePlots(data)
```

Arguments

data

output from function designSampleSize.

Details

Data in the example is based on the results of sample size calculation from function designSampleSize

Value

Plot for estimated sample size with assigned variable.

Author(s)

Meena Choi, Ching-Yun Chang, Olga Vitek.

Examples

```
# Based on the results of sample size calculation from function designSampleSize,
# we generate a series of sample size plots for number of biological replicates, or peptides,
# or transitions or power plot.
QuantData<-dataProcess(SRMRawData)
head(QuantData$ProcessedData)
## based on multiple comparisons (T1 vs T3; T1 vs T7; T1 vs T9)
comparison1<-matrix(c(-1,0,1,0,0,0,0,0,0,0,0,0,nrow=1)
comparison2<-matrix(c(-1,0,0,0,0,0,1,0,0,0,nrow=1)
comparison3<-matrix(c(-1,0,0,0,0,0,0,0,1,0),nrow=1)
comparison<-rbind(comparison1,comparison2, comparison3)
row.names(comparison)<-c("T3-T1","T7-T1","T9-T1")
colnames(comparison)<-unique(QuantData$ProteinLevelData$GROUP)
```

testResultMultiComparisons<-groupComparison(contrast.matrix=comparison, data=QuantData)

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DIARawData

Example dataset from a label-free DIA, a group comparison study of S.Pyogenes.

Description

This example dataset was obtained from a group comparison study of S. Pyogenes. Two conditions, S. Pyogenes with 0% and 10% of human plasma added (denoted Strep 0% and Strep 10%), were profiled in two replicates, in the label-free mode, with a SWATH-MS-enabled AB SCIEX TripleTOF 5600 System. The identification and quantification of spectral peaks was assisted by a spectral library, and was performed using OpenSWATH software (http://proteomics.ethz.ch/openswath.html). For reasons of space, the example dataset only contains two proteins from this study. Protein FabG shows strong evidence of differential abundance, while protein Probable RNA helicase exp9 only shows moderate evidence of dif- ferential abundance between conditions.

Usage

DTARawData

Format

data.frame

Details

The raw data (input data for MSstats) is required to contain variable of ProteinName, PeptideSequence, PrecursorCharge, FragmentIon, ProductCharge, IsotopeLabelType, Condition, BioReplicate, Run, Intensity. The variable names should be fixed.

If the information of one or more columns is not available for the original raw data, please retain the column variables and type in fixed value. For example, the original raw data does not contain the information of PrecursorCharge and ProductCharge, we retain the column PrecursorCharge and ProductCharge and then type in NA for all transitions in RawData.

Variable Intensity is required to be original signal without any log transformation and can be specified as the peak of height or the peak of area under curve.

Value

data.frame with the required format of MSstats.

Author(s)

```
Meena Choi, Olga Vitek.

Maintainer: Meena Choi (<mnchoi67@gmail.com>)
```

Examples

```
head(DIARawData)
```

 ${\tt DIAUmpiretoMSstatsFormat}$

Import DIA-Umpire files

Description

Import DIA-Umpire files

Usage

```
DIAUmpiretoMSstatsFormat(
    raw.frag,
    raw.pep,
    raw.pro,
    annotation,
    useSelectedFrag = TRUE,
    useSelectedPep = TRUE,
    removeFewMeasurements = TRUE,
    removeProtein_with1Feature = FALSE,
    summaryforMultipleRows = max,
    use_log_file = TRUE,
    append = FALSE,
    verbose = TRUE,
    log_file_path = NULL,
    ...
)
```

Arguments

raw. frag name of FragSummary_date.xls data, which includes feature-level data.

raw. pep name of PeptideSummary_date.xls data, which includes selected fragments information.

raw. pro name of ProteinSummary_date.xls data, which includes selected peptides information.

annotation name of annotation data which includes Condition, BioReplicate, Run informa-

useSelectedFrag

TRUE will use the selected fragment for each peptide. 'Selected_fragments' column is required.

useSelectedPep TRUE will use the selected peptide for each protein. 'Selected_peptides' column is required.

removeFewMeasurements

TRUE (default) will remove the features that have 1 or 2 measurements across runs.

removeProtein_with1Feature

TRUE will remove the proteins which have only 1 feature, which is the combination of peptide, precursor charge, fragment and charge. FALSE is default.

summaryforMultipleRows

max(default) or sum - when there are multiple measurements for certain feature and certain run, use highest or sum of multiple intensities.

use_log_file logical. If TRUE, information about data processing will be saved to a file.

append logical. If TRUE, information about data processing will be added to an existing

log file.

verbose logical. If TRUE, information about data processing wil be printed to the con-

sole.

log_file_path character. Path to a file to which information about data processing will be

saved. If not provided, such a file will be created automatically. If 'append =

TRUE', has to be a valid path to a file.

... additional parameters to 'data.table::fread'.

Value

data.frame in the MSstats required format.

Author(s)

Meena Choi, Olga Vitek

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getProcessed

Get feature-level data to be used in the MSstatsSummarizationOutput function

Description

Get feature-level data to be used in the MSstatsSummarizationOutput function

Usage

```
getProcessed(input)
```

Arguments

input

data.table processed by dataProcess subfunctions

Value

data.table processed by dataProcess subfunctions

```
raw = DDARawData
method = "TMP"
cens = "NA"
impute = TRUE
MSstatsConvert::MSstatsLogsSettings(FALSE)
input = MSstatsPrepareForDataProcess(raw, 2, NULL)
input = MSstatsNormalize(input, "EQUALIZEMEDIANS")
input = MSstatsMergeFractions(input)
input = MSstatsHandleMissing(input, "TMP", TRUE, "NA", 0.999)
input_all = MSstatsSelectFeatures(input, "all") # all features
input_5 = MSstatsSelectFeatures(data.table::copy(input),
"topN", top_n = 5) # top 5 features
proc1 = getProcessed(input_all)
proc2 = getProcessed(input_5)
proc1
proc2
```

getSamplesInfo 19

getSamplesInfo

Get information about number of measurements for each group

Description

Get information about number of measurements for each group

Usage

```
getSamplesInfo(summarization_output)
```

Arguments

```
summarization_output of the dataProcess function
```

Details

This function extracts information required to compute percentages of missing and imputed values in group comparison.

Value

data.table

Examples

```
QuantData <- dataProcess(DDARawData, use_log_file = FALSE)
samples_info <- getSamplesInfo(QuantData)
samples_info</pre>
```

getSelectedProteins

Get proteins based on names or integer IDs

Description

Get proteins based on names or integer IDs

Usage

```
getSelectedProteins(chosen_proteins, all_proteins)
```

Arguments

```
chosen_proteins
```

protein names or integers IDs

all_proteins all unique proteins

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Value

character

groupComparison

Whole plot testing

Description

Whole plot testing

Usage

```
groupComparison(
  contrast.matrix,
  data,
  save_fitted_models = TRUE,
  log_base = 2,
  use_log_file = TRUE,
  append = FALSE,
  verbose = TRUE,
  log_file_path = NULL
)
```

Arguments

contrast.matrix

comparison between conditions of interests.

data name of the (output of dataProcess function) data set.

save_fitted_models

logical, if TRUE, fitted models will be added to the output.

log_base base of the logarithm used in dataProcess.

use_log_file logical. If TRUE, information about data processing will be saved to a file.

append logical. If TRUE, information about data processing will be added to an existing

log file.

verbose logical. If TRUE, information about data processing wil be printed to the con-

sole.

log_file_path character. Path to a file to which information about data processing will be

saved. If not provided, such a file will be created automatically. If 'append =

TRUE', has to be a valid path to a file.

Details

contrast.matrix: comparison of interest. Based on the levels of conditions, specify 1 or -1 to the conditions of interests and 0 otherwise. The levels of conditions are sorted alphabetically. Command levels(QuantData\$FeatureLevelData\$GROUP_ORIGINAL) can illustrate the actual order of the levels of conditions. The underlying model fitting functions are lm and lmer for the fixed effects model and mixed effects model, respectively. The input of this function is the quantitative data from function (dataProcess).

Value

list that consists of three elements: "ComparisonResult" - data.frame with results of statistical testing, "ModelQC" - data.frame with data used to fit models for group comparison and "FittedModel" - list of fitted models.

Examples

```
# Consider quantitative data (i.e. QuantData) from yeast study with ten time points of interests,
# three biological replicates, and no technical replicates.
# It is a time-course experiment and we attempt to compare differential abundance
# between time 1 and 7 in a set of targeted proteins.
# In this label-based SRM experiment, MSstats uses the fitted model with expanded scope of
# Biological replication.
QuantData <- dataProcess(SRMRawData, use_log_file = FALSE)
head(QuantData$FeatureLevelData)
levels(QuantData$ProteinLevelData$GROUP)
comparison <- matrix(c(-1,0,0,0,0,0,1,0,0,0),nrow=1)
row.names(comparison) <- "T7-T1"</pre>
groups = levels(QuantData$ProteinLevelData$GROUP)
colnames(comparison) <- groups[order(as.numeric(groups))]</pre>
# Tests for differentially abundant proteins with models:
# label-based SRM experiment with expanded scope of biological replication.
testResultOneComparison <- groupComparison(contrast.matrix=comparison, data=QuantData,
                                            use_log_file = FALSE)
# table for result
testResultOneComparison$ComparisonResult
```

groupComparisonPlots Visualization for model-based analysis and summarizing differentially abundant proteins

Description

To summarize the results of log-fold changes and adjusted p-values for differentially abundant proteins, groupComparisonPlots takes testing results from function (groupComparison) as input and automatically generate three types of figures in pdf files as output: (1) volcano plot (specify "VolcanoPlot" in option type) for each comparison separately; (2) heatmap (specify "Heatmap" in option type) for multiple comparisons; (3) comparison plot (specify "ComparisonPlot" in option type) for multiple comparisons per protein.

Usage

```
groupComparisonPlots(
  data,
  type,
  sig = 0.05,
  FCcutoff = FALSE,
  logBase.pvalue = 10,
 ylimUp = FALSE,
 ylimDown = FALSE,
 xlimUp = FALSE,
  x.axis.size = 10,
 y.axis.size = 10,
  dot.size = 3,
  text.size = 4,
  text.angle = 0,
  legend.size = 13,
  ProteinName = TRUE,
  colorkey = TRUE,
  numProtein = 100,
  clustering = "both",
 width = 10,
  height = 10,
 which.Comparison = "all",
 which.Protein = "all",
  address = ""
)
```

Arguments

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data	ComparisonResilif	in festing official from	n function groupComparison.
aaca	Companisonitesant	in testing output nor	in ranction group comparison.

type choice of visualization. "VolcanoPlot" represents volcano plot of log fold changes

and adjusted p-values for each comparison separately. "Heatmap" represents heatmap of adjusted p-values for multiple comparisons. "ComparisonPlot" represents comparison plot of log fold changes for multiple comparisons per pro-

tein.

FDR cutoff for the adjusted p-values in heatmap and volcano plot. level of sig-

nificance for comparison plot. 100(1-sig)% confidence interval will be drawn.

sig=0.05 is default.

FCcutoff for volcano plot or heatmap, whether involve fold change cutoff or not. FALSE

(default) means no fold change cutoff is applied for significance analysis. FC-

cutoff = specific value means specific fold change cutoff is applied.

logBase.pvalue for volcano plot or heatmap, (-) logarithm transformation of adjusted p-value

with base 2 or 10(default).

ylimUp for all three plots, upper limit for y-axis. FALSE (default) for volcano plot/heatmap

use maximum of -log2 (adjusted p-value) or -log10 (adjusted p-value). FALSE

(default) for comparison plot uses maximum of log-fold change + CI.

ylimDown for all three plots, lower limit for y-axis. FALSE (default) for volcano plot/heatmap use minimum of -log2 (adjusted p-value) or -log10 (adjusted p-value). FALSE (default) for comparison plot uses minimum of log-fold change - CI. xlimUp for Volcano plot, the limit for x-axis. FALSE (default) for use maximum for absolute value of log-fold change or 3 as default if maximum for absolute value of log-fold change is less than 3. x.axis.size size of axes labels, e.g. name of the comparisons in heatmap, and in comparison plot. Default is 10. size of axes labels, e.g. name of targeted proteins in heatmap. Default is 10. y.axis.size size of dots in volcano plot and comparison plot. Default is 3. dot.size text.size size of ProteinName label in the graph for Volcano Plot. Default is 4. text.angle angle of x-axis labels represented each comparison at the bottom of graph in comparison plot. Default is 0. legend.size size of legend for color at the bottom of volcano plot. Default is 7. ProteinName for volcano plot only, whether display protein names or not. TRUE (default) means protein names, which are significant, are displayed next to the points. FALSE means no protein names are displayed. colorkev TRUE(default) shows colorkey. The number of proteins which will be presented in each heatmap. Default is numProtein 100. Maximum possible number of protein for one heatmap is 180. clustering Determines how to order proteins and comparisons. Hierarchical cluster analysis with Ward method(minimum variance) is performed. 'protein' means that protein dendrogram is computed and reordered based on protein means (the order of row is changed). 'comparison' means comparison dendrogram is computed and reordered based on comparison means (the order of comparison is changed). 'both' means to reorder both protein and comparison. Default is 'protein'. width width of the saved file. Default is 10. height height of the saved file. Default is 10. which.Comparison list of comparisons to draw plots. List can be labels of comparisons or order numbers of comparisons from levels(data\$Label), such as levels(testResultMultiComparisons\$Comparison Default is "all", which generates all plots for each protein. which.Protein Protein list to draw comparison plots. List can be names of Proteins or order numbers of Proteins from levels(testResultMultiComparisons\$ComparisonResult\$Protein). Default is "all", which generates all comparison plots for each protein. address the name of folder that will store the results. Default folder is the current working directory. The other assigned folder has to be existed under the current working directory. An output pdf file is automatically created with the default name of "VolcanoPlot.pdf" or "Heatmap.pdf" or "ComparisonPlot.pdf". The command address can help to specify where to store the file as well as how to

modify the beginning of the file name. If address=FALSE, plot will be not saved

as pdf file but showed in window.

Details

- Volcano plot: illustrate actual log-fold changes and adjusted p-values for each comparison separately with all proteins. The x-axis is the log fold change. The base of logarithm transformation is the same as specified in "logTrans" from dataProcess. The y-axis is the negative log2 or log10 adjusted p-values. The horizontal dashed line represents the FDR cutoff. The points below the FDR cutoff line are non-significantly abundant proteins (colored in black). The points above the FDR cutoff line are significantly abundant proteins (colored in red/blue for up-/down-regulated). If fold change cutoff is specified (FCcutoff = specific value), the points above the FDR cutoff line but within the FC cutoff line are non-significantly abundant proteins (colored in black)/
- Heatmap: illustrate up-/down-regulated proteins for multiple comparisons with all proteins.
 Each column represents each comparison of interest. Each row represents each protein. Color red/blue represents proteins in that specific comparison are significantly up-regulated/down-regulated proteins with FDR cutoff and/or FC cutoff. The color scheme shows the evidences of significance. The darker color it is, the stronger evidence of significance it has. Color gold represents proteins are not significantly different in abundance.
- Comparison plot: illustrate log-fold change and its variation of multiple comparisons for single protein. X-axis is comparison of interest. Y-axis is the log fold change. The red points are the estimated log fold change from the model. The blue error bars are the confidence interval with 0.95 significant level for log fold change. This interval is only based on the standard error, which is estimated from the model.

```
QuantData<-dataProcess(SRMRawData, use_log_file = FALSE)
head(QuantData$FeatureLevelData)
## based on multiple comparisons (T1 vs T3; T1 vs T7; T1 vs T9)
comparison1<-matrix(c(-1,0,1,0,0,0,0,0,0,0), nrow=1)
comparison2<-matrix(c(-1,0,0,0,0,0,1,0,0,0), nrow=1)
comparison3<-matrix(c(-1,0,0,0,0,0,0,0,1,0), nrow=1)
comparison<-rbind(comparison1, comparison2, comparison3)</pre>
row.names(comparison)<-c("T3-T1","T7-T1","T9-T1")</pre>
groups = levels(QuantData$ProteinLevelData$GROUP)
colnames(comparison) <- groups[order(as.numeric(groups))]</pre>
testResultMultiComparisons<-groupComparison(contrast.matrix=comparison,
data=QuantData,
use_log_file = FALSE)
testResultMultiComparisons$ComparisonResult
# Volcano plot with FDR cutoff = 0.05 and no FC cutoff
groupComparisonPlots(data=testResultMultiComparisons$ComparisonResult, type="VolcanoPlot",
logBase.pvalue=2, address="Ex1_")
# Volcano plot with FDR cutoff = 0.05, FC cutoff = 70, upper y-axis limit = 100,
# and no protein name displayed
# FCcutoff=70 is for demonstration purpose
groupComparisonPlots(data=testResultMultiComparisons$ComparisonResult, type="VolcanoPlot",
FCcutoff=70, logBase.pvalue=2, ylimUp=100, ProteinName=FALSE,address="Ex2_")
# Heatmap with FDR cutoff = 0.05
groupComparisonPlots(data=testResultMultiComparisons$ComparisonResult, type="Heatmap",
logBase.pvalue=2, address="Ex1_")
# Heatmap with FDR cutoff = 0.05 and FC cutoff = 70
```

makePeptidesDictionary

```
# FCcutoff=70 is for demonstration purpose
groupComparisonPlots(data=testResultMultiComparisons$ComparisonResult, type="Heatmap",
FCcutoff=70, logBase.pvalue=2, address="Ex2_")
# Comparison Plot
groupComparisonPlots(data=testResultMultiComparisons$ComparisonResult, type="ComparisonPlot",
address="Ex1_")
# Comparison Plot
groupComparisonPlots(data=testResultMultiComparisons$ComparisonResult, type="ComparisonPlot",
ylimUp=8, ylimDown=-1, address="Ex2_")
```

makePeptidesDictionary

Prepare a peptides dictionary for global standards normalization

Description

Prepare a peptides dictionary for global standards normalization

Usage

```
makePeptidesDictionary(input, normalization)
```

Arguments

input 'data.table' in MSstats standard format normalization normalization method

Details

This function extracts information required to perform normalization with global standards. It is useful for running the summarization workflow outside of the dataProcess function.

```
input = data.table::as.data.table(DDARawData)
peptides_dict = makePeptidesDictionary(input, "GLOBALSTANDARDS")
head(peptides_dict) # ready to be passed to the MSstatsNormalize function
```

MaxQtoMSstatsFormat Import MaxQuant files

Description

Import MaxQuant files

Usage

```
MaxQtoMSstatsFormat(
  evidence,
  annotation,
  proteinGroups,
  proteinID = "Proteins",
  useUniquePeptide = TRUE,
  summaryforMultipleRows = max,
  removeFewMeasurements = TRUE,
  removeMpeptides = FALSE,
  removeOxidationMpeptides = FALSE,
  removeProtein_with1Peptide = FALSE,
  use_log_file = TRUE,
  append = FALSE,
  verbose = TRUE,
  log_file_path = NULL,
)
```

Arguments

evidence name of 'evidence.txt' data, which includes feature-level data.

annotation name of 'annotation.txt' data which includes Raw.file, Condition, BioReplicate,

Run, IsotopeLabelType information.

proteinGroups name of 'proteinGroups.txt' data. It needs to matching protein group ID. If

proteinGroups=NULL, use 'Proteins' column in 'evidence.txt'.

proteinID 'Proteins' (default) or 'Leading.razor.protein' for Protein ID.

useUniquePeptide

TRUE (default) removes peptides that are assigned for more than one proteins. We assume to use unique peptide for each protein.

summaryforMultipleRows

max(default) or sum - when there are multiple measurements for certain feature and certain run, use highest or sum of multiple intensities.

removeFewMeasurements

TRUE (default) will remove the features that have 1 or 2 measurements across runs.

removeMpeptides

TRUE will remove the peptides including 'M' sequence. FALSE is default.

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removeOxidationMpeptides

TRUE will remove the peptides including 'oxidation (M)' in modification. FALSE

is default.

removeProtein_with1Peptide

TRUE will remove the proteins which have only 1 peptide and charge. FALSE

is default.

use_log_file logical. If TRUE, information about data processing will be saved to a file.

append logical. If TRUE, information about data processing will be added to an existing

log file.

verbose logical. If TRUE, information about data processing wil be printed to the con-

sole.

log_file_path character. Path to a file to which information about data processing will be

saved. If not provided, such a file will be created automatically. If 'append =

TRUE', has to be a valid path to a file.

... additional parameters to 'data.table::fread'.

Value

data.frame in the MSstats required format.

Note

Warning: MSstats does not support for metabolic labeling or iTRAQ experiments.

Author(s)

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28 modelBasedQCPlots

modelBasedQCPlots

Visualization for model-based quality control in fitting model

Description

To check the assumption of linear model for whole plot inference, modelBasedQCPlots takes the results after fitting models from function (groupComparison) as input and automatically generate two types of figures in pdf files as output: (1) normal quantile-quantile plot (specify "QQPlot" in option type) for checking normally distributed errors.; (2) residual plot (specify "ResidualPlot" in option type).

Usage

```
modelBasedQCPlots(
  data,
  type,
  axis.size = 10,
  dot.size = 3,
  width = 10,
  height = 10,
  which.Protein = "all",
  address = ""
)
```

Arguments

data output from function groupComparison.

type choice of visualization. "QQPlots" represents normal quantile-quantile plot for

each protein after fitting models. "ResidualPlots" represents a plot of residuals

versus fitted values for each protein in the dataset.

axis.size size of axes labels. Default is 10.

dot.size size of points in the graph for residual plots and QQ plots. Default is 3.

width width of the saved file. Default is 10. height height of the saved file. Default is 10.

which.Protein Protein list to draw plots. List can be names of Proteins or order numbers of Pro-

teins from levels(testResultOneComparison\$ComparisonResult\$Protein). De-

fault is "all", which generates all plots for each protein.

address name that will serve as a prefix to the name of output file.

Details

Results based on statistical models for whole plot level inference are accurate as long as the assumptions of the model are met. The model assumes that the measurement errors are normally distributed with mean 0 and constant variance. The assumption of a constant variance can be checked by examining the residuals from the model.

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• QQPlots: a normal quantile-quantile plot for each protein is generated in order to check whether the errors are well approximated by a normal distribution. If points fall approximately along a straight line, then the assumption is appropriate for that protein. Only large deviations from the line are problematic.

 ResidualPlots: The plots of residuals against predicted(fitted) values. If it shows a random scatter, then the assumption is appropriate.

Value

produce a pdf file

Examples

```
QuantData <- dataProcess(SRMRawData, use_log_file = FALSE)
head(QuantData$FeatureLevelData)
levels(QuantData$FeatureLevelData$GROUP)
comparison <- matrix(c(-1,0,0,0,0,1,0,0,0),nrow=1)
row.names(comparison) <- "T7-T1"
colnames(comparison) <- unique(QuantData$ProteinLevelData$GROUP)
# Tests for differentially abundant proteins with models:
# label-based SRM experiment with expanded scope of biological replication.
testResultOneComparison <- groupComparison(contrast.matrix=comparison, data=QuantData, use_log_file = FALSE)
# normal quantile-quantile plots
modelBasedQCPlots(data=testResultOneComparison, type="QQPlots", address="")
# residual plots
modelBasedQCPlots(data=testResultOneComparison, type="ResidualPlots", address="")
```

MSstatsContrastMatrix Create a contrast matrix for groupComparison function

Description

Create a contrast matrix for groupComparison function

Usage

```
MSstatsContrastMatrix(contrasts, conditions, labels = NULL)
```

Arguments

contrasts One of the following: i) list of lists. Each sub-list consists of two vectors that

name conditions that will be compared. See the details section for more information ii) matrix. In this case, it's correctness will be checked iii) "pairwise". In this case, pairwise comparison matrix will be generated iv) data.frame. In this

case, input will be converted to matrix

conditions unique condition labels

labels labels for contrasts (row.names of the contrast matrix)

MSstatsGroupComparison

Group comparison

Description

Group comparison

Usage

```
MSstatsGroupComparison(
   summarized_list,
   contrast_matrix,
   save_fitted_models,
   repeated,
   samples_info
)
```

Arguments

MSstatsGroupComparisonOutput

Create output of group comparison based on results for individual proteins

Description

Create output of group comparison based on results for individual proteins

Usage

```
MSstatsGroupComparisonOutput(input, summarization_output, log_base = 2)
```

Arguments

```
input output of MSstatsGroupComparison function
summarization_output
output of dataProcess function
log_base base of the logarithm used in fold-change calculation
```

Value

list, same as the output of 'groupComparison'

MSstatsGroupComparisonSingleProtein

Group comparison for a single protein

Description

Group comparison for a single protein

Usage

```
MSstatsGroupComparisonSingleProtein(
    single_protein,
    contrast_matrix,
    repeated,
    groups,
    samples_info,
    save_fitted_models,
    has_imputed
)
```

Arguments

```
single_protein data.table with summarized data for a single protein contrast_matrix

contrast matrix

repeated if TRUE, repeated measurements will be modeled groups unique labels of experimental conditions samples_info number of runs per group save_fitted_models if TRUE, fitted model will be saved. If not, it will be replaced with NULL has_imputed TRUE if missing values have been imputed
```

```
QuantData <- dataProcess(SRMRawData, use_log_file = FALSE)
group_comparison_input <- MSstatsPrepareForGroupComparison(QuantData)
levels(QuantData$ProteinLevelData$GROUP)
comparison <- matrix(c(-1,0,0,0,0,0,1,0,0,0),nrow=1)
row.names(comparison) <- "T7-T1"
groups = levels(QuantData$ProteinLevelData$GROUP)
colnames(comparison) <- groups[order(as.numeric(groups))]
samples_info <- getSamplesInfo(QuantData)
repeated <- checkRepeatedDesign(QuantData)
single_output <- MSstatsGroupComparisonSingleProtein(
   group_comparison_input[[1]], comparison, repeated, groups, samples_info,
   FALSE, TRUE)
single_output # same as a single element of MSstatsGroupComparison output</pre>
```

MSstatsHandleMissing

MSstatsHandleMissing Handle censored missing values

Description

Handle censored missing values

Usage

```
MSstatsHandleMissing(
  input,
  summary_method,
  impute,
  missing_symbol,
  censored_cutoff
)
```

Arguments

```
input 'data.table' in MSstats data format
summary_method summarization method ('summaryMethod' parameter to 'dataProcess')
impute if TRUE, missing values are supposed to be imputed ('MBimpute' parameter to 'dataProcess')
missing_symbol 'censoredInt' parameter to 'dataProcess'
censored_cutoff
'maxQuantileforCensored' parameter to 'dataProcess'
```

Value

data.table

```
raw = DDARawData
method = "TMP"
cens = "NA"
impute = TRUE
MSstatsConvert::MSstatsLogsSettings(FALSE)
input = MSstatsPrepareForDataProcess(raw, 2, NULL)
input = MSstatsNormalize(input, "EQUALIZEMEDIANS")
input = MSstatsMergeFractions(input)
input = MSstatsHandleMissing(input, "TMP", TRUE, "NA", 0.999)
head(input)
```

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 ${\tt MSstatsMergeFractions} \ \ \textit{Re-format the data before feature selection}$

Description

Re-format the data before feature selection

Usage

```
MSstatsMergeFractions(input)
```

Arguments

input 'data.table' in MSstats format

Value

data.table

Examples

```
raw = DDARawData
method = "TMP"
cens = "NA"
impute = TRUE
MSstatsConvert::MSstatsLogsSettings(FALSE)
input = MSstatsPrepareForDataProcess(raw, 2, NULL)
input = MSstatsNormalize(input, "EQUALIZEMEDIANS")
input = MSstatsMergeFractions(input)
head(input)
```

MSstatsNormalize

Normalize MS data

Description

Normalize MS data

Usage

```
MSstatsNormalize(
  input,
  normalization_method,
  peptides_dict = NULL,
  standards = NULL
)
```

Arguments

input data.table in MSstats format

normalization_method

name of a chosen normalization method: "NONE" or "FALSE" for no normalization, "EQUALIZEMEDIANS" for median normalization, "QUANTILE" normalization for quantile normalization from 'preprocessCore' package, "GLOB-ALSTANDARDS" for normalization based on selected peptides or proteins.

peptides_dict 'data.table' of names of peptides and their corresponding features.

standards character vector with names of standards, required if "GLOBALSTANDARDS"

method was selected.

Value

data.table

Examples

```
raw = DDARawData
method = "TMP"
cens = "NA"
impute = TRUE
MSstatsConvert::MSstatsLogsSettings(FALSE)
input = MSstatsPrepareForDataProcess(raw, 2, NULL)
input = MSstatsNormalize(input, "EQUALIZEMEDIANS") # median normalization
head(input)
```

MSstatsPrepareForDataProcess

Prepare data for processing by 'dataProcess' function

Description

Prepare data for processing by 'dataProcess' function

Usage

MSstatsPrepareForDataProcess(input, log_base, fix_missing)

Arguments

input 'data.table' in MSstats format

log_base base of the logarithm to transform intensities

fix_missing str, optional. Defaults to NULL, which means no action. If not NULL, must

be one of the options: "zero_to_na" or "na_to_zero". If "zero_to_na", Intensity values equal exactly to 0 will be converted to NA. If "na_to_zero", missing

values will be replaced by zeros.

Value

data.table

Examples

```
raw = DDARawData
method = "TMP"
cens = "NA"
impute = TRUE
MSstatsConvert::MSstatsLogsSettings(FALSE)
input = MSstatsPrepareForDataProcess(raw, 2, NULL)
head(input)
```

 ${\it MSstatsPrepareForGroupComparison}$

Prepare output for dataProcess for group comparison

Description

Prepare output for dataProcess for group comparison

Usage

MSstatsPrepareForGroupComparison(summarization_output)

Arguments

```
summarization_output of dataProcess
```

Value

list of run-level data for each protein in the input. This list has a "has_imputed" attribute that indicates if missing values were imputed in the input dataset.

```
QuantData <- dataProcess(SRMRawData, use_log_file = FALSE)
group_comparison_input = MSstatsPrepareForGroupComparison(QuantData)
length(group_comparison_input) # list of length equal to number of proteins
# in protein-level data of QuantData
head(group_comparison_input[[1]])</pre>
```

 ${\it MSstatsPrepareForSummarization}$

Prepare feature-level data for protein-level summarization

Description

Prepare feature-level data for protein-level summarization

Usage

```
MSstatsPrepareForSummarization(
  input,
  method,
  impute,
  censored_symbol,
  remove_uninformative_feature_outlier
)
```

Arguments

input feature-level data processed by dataProcess subfunctions

method summarization method - 'summaryMethod' parameter of the dataProcess func-

tion

impute if TRUE, censored missing values will be imputed - 'MBimpute' parameter of

the dataProcess function

censored_symbol

censored missing value indicator - 'censoredInt' parameter of the dataProcess

function

remove_uninformative_feature_outlier

if TRUE, features labeled as outlier of uninformative by the MSstatsSelectFea-

tures function will not be used in summarization

Value

data.table

```
raw = DDARawData
method = "TMP"
cens = "NA"
impute = TRUE
MSstatsConvert::MSstatsLogsSettings(FALSE)
input = MSstatsPrepareForDataProcess(raw, 2, NULL)
head(input)
```

38 MSstatsSelectFeatures

MSstatsSelectFeatures Feature selection before feature-level data summarization

Description

Feature selection before feature-level data summarization

Usage

```
MSstatsSelectFeatures(input, method, top_n = 3, min_feature_count = 2)
```

Arguments

Value

data.table

```
raw = DDARawData
method = "TMP"
cens = "NA"
impute = TRUE
MSstatsConvert::MSstatsLogsSettings(FALSE)
input = MSstatsPrepareForDataProcess(raw, 2, NULL)
input = MSstatsNormalize(input, "EQUALIZEMEDIANS")
input = MSstatsMergeFractions(input)
input = MSstatsHandleMissing(input, "TMP", TRUE, "NA", 0.999)
input_all = MSstatsSelectFeatures(input, "all") # all features
input_5 = MSstatsSelectFeatures(data.table::copy(input), "topN", top_n = 5) # top 5 features
input_informative = MSstatsSelectFeatures(input, "highQuality") # feature selection
head(input_all)
head(input_all)
head(input_informative)
```

MSstatsSummarizationOutput

Post-processing output from MSstats summarization

Description

Post-processing output from MSstats summarization

Usage

```
MSstatsSummarizationOutput(
  input,
  summarized,
  processed,
  method,
  impute,
  censored_symbol
)
```

Arguments

```
input 'data.table' in MSstats format

summarized output of the 'MSstatsSummarize' function

processed output of MSstatsSelectFeatures

method name of the summarization method ('summaryMethod' parameter to 'dataProcess')

impute if TRUE, censored missing values were imputed ('MBimpute' parameter to 'dataProcess')

censored_symbol

censored missing value indicator ('censoredInt' parameter to 'dataProcess')
```

Value

list that consists of the following elements:

- FeatureLevelData feature-level data after processing
- ProteinLevelData protein-level (summarized) data
- SummaryMethod (string) name of summarization method that was used

```
raw = DDARawData
method = "TMP"
cens = "NA"
impute = TRUE
MSstatsConvert::MSstatsLogsSettings(FALSE)
```

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```
input = MSstatsPrepareForDataProcess(raw, 2, NULL)
input = MSstatsNormalize(input, "EQUALIZEMEDIANS")
input = MSstatsMergeFractions(input)
input = MSstatsHandleMissing(input, "TMP", TRUE, "NA", 0.999)
input = MSstatsSelectFeatures(input, "all")
processed = getProcessed(input)
input = MSstatsPrepareForSummarization(input, method, impute, cens, FALSE)
input_split = split(input, input$PROTEIN)
summarized = MSstatsSummarize(input_split, method, impute, cens, FALSE, TRUE)
output = output = MSstatsSummarizationOutput(input, summarized, processed, method, impute, cens)
```

MSstatsSummarize

Feature-level data summarization

Description

Feature-level data summarization

Usage

```
MSstatsSummarize(
   proteins_list,
   method,
   impute,
   censored_symbol,
   remove50missing,
   equal_variance
)
```

Arguments

proteins_list list of processed feature-level data

method summarization method: "linear" or "TMP"

impute only for summaryMethod = "TMP" and censoredInt = 'NA' or '0'. TRUE (de-

fault) imputes 'NA' or '0' (depending on censoredInt option) by Accelated fail-

ure model. FALSE uses the values assigned by cutoffCensored

censored_symbol

Missing values are censored or at random. 'NA' (default) assumes that all 'NA's in 'Intensity' column are censored. '0' uses zero intensities as censored intensity. In this case, NA intensities are missing at random. The output from Skyline should use '0'. Null assumes that all NA intensites are randomly missing.

remove50missing

only for summaryMethod = "TMP". TRUE removes the runs which have more than 50% missing values. FALSE is default.

equal_variance only for summaryMethod = "linear". Default is TRUE. Logical variable for whether the model should account for heterogeneous variation among intensities from different features. Default is TRUE, which assume equal variance among intensities from features. FALSE means that we cannot assume equal variance

among intensities from features, then we will account for heterogeneous variation from different features.

Value

list of length one with run-level data.

Examples

```
raw = DDARawData
method = "TMP"
cens = "NA"
impute = TRUE
MSstatsConvert::MSstatsLogsSettings(FALSE)
input = MSstatsPrepareForDataProcess(raw, 2, NULL)
input = MSstatsNormalize(input, "EQUALIZEMEDIANS")
input = MSstatsMergeFractions(input)
input = MSstatsHandleMissing(input, "TMP", TRUE, "NA", 0.999)
input = MSstatsSelectFeatures(input, "all")
processed = getProcessed(input)
input = MSstatsPrepareForSummarization(input, method, impute, cens, FALSE)
input_split = split(input, input$PROTEIN)
summarized = MSstatsSummarize(input_split, method, impute, cens, FALSE, TRUE)
length(summarized) # list of summarization outputs for each protein
head(summarized[[1]][[1]]) # run-level summary
```

 ${\tt MSstatsSummarizeSingleLinear}$

Linear model-based summarization for a single protein

Description

Linear model-based summarization for a single protein

Usage

```
MSstatsSummarizeSingleLinear(single_protein, equal_variances = TRUE)
```

Arguments

```
single_protein feature-level data for a single protein equal_variances
```

if TRUE, observation are assumed to be homoskedastic

Value

list with protein-level data

Examples

```
raw = DDARawData
method = "linear"
cens = NULL
impute = FALSE
# currently, MSstats only supports MBimpute = FALSE for linear summarization
MSstatsConvert::MSstatsLogsSettings(FALSE)
input = MSstatsPrepareForDataProcess(raw, 2, NULL)
input = MSstatsNormalize(input, "EQUALIZEMEDIANS")
input = MSstatsMergeFractions(input)
input = MSstatsHandleMissing(input, "TMP", TRUE, "NA", 0.999)
input = MSstatsSelectFeatures(input, "all")
input = MSstatsPrepareForSummarization(input, method, impute, cens, FALSE)
input_split = split(input, input$PROTEIN)
single_protein_summary = MSstatsSummarizeSingleLinear(input_split[[1]])
head(single_protein_summary[[1]])
```

MSstatsSummarizeSingleTMP

Tukey Median Polish summarization for a single protein

Description

Tukey Median Polish summarization for a single protein

Usage

```
MSstatsSummarizeSingleTMP(
    single_protein,
    impute,
    censored_symbol,
    remove50missing
)
```

Arguments

```
single_protein feature-level data for a single protein

only for summaryMethod = "TMP" and censoredInt = 'NA' or '0'. TRUE (default) imputes 'NA' or '0' (depending on censoredInt option) by Accelated failure model. FALSE uses the values assigned by cutoffCensored
```

```
censored_symbol
```

Missing values are censored or at random. 'NA' (default) assumes that all 'NA's in 'Intensity' column are censored. '0' uses zero intensities as censored intensity. In this case, NA intensities are missing at random. The output from Skyline should use '0'. Null assumes that all NA intensities are randomly missing.

remove50missing

only for summaryMethod = "TMP". TRUE removes the runs which have more than 50% missing values. FALSE is default.

Value

list of two data.tables: one with fitted survival model, the other with protein-level data

Examples

```
raw = DDARawData
method = "TMP"
cens = "NA"
impute = TRUE
# currently, MSstats only supports MBimpute = FALSE for linear summarization
MSstatsConvert::MSstatsLogsSettings(FALSE)
input = MSstatsPrepareForDataProcess(raw, 2, NULL)
input = MSstatsNormalize(input, "EQUALIZEMEDIANS")
input = MSstatsMergeFractions(input)
input = MSstatsHandleMissing(input, "TMP", TRUE, "NA", 0.999)
input = MSstatsSelectFeatures(input, "all")
input = MSstatsPrepareForSummarization(input, method, impute, cens, FALSE)
input_split = split(input, input$PROTEIN)
single_protein_summary = MSstatsSummarizeSingleTMP(input_split[[1]],
                                                    impute, cens, FALSE)
head(single_protein_summary[[1]])
```

OpenMStoMSstatsFormat Import OpenMS files

Description

Import OpenMS files

```
OpenMStoMSstatsFormat(
  input,
  annotation = NULL,
  useUniquePeptide = TRUE,
  removeFewMeasurements = TRUE,
  removeProtein_with1Feature = FALSE,
  summaryforMultipleRows = max,
```

```
use_log_file = TRUE,
append = FALSE,
verbose = TRUE,
log_file_path = NULL,
...
)
```

Arguments

input name of MSstats input report from OpenMS, which includes feature(peptide

ion)-level data.

annotation name of 'annotation.txt' data which includes Condition, BioReplicate, Run. Run

should be the same as filename.

useUniquePeptide

TRUE (default) removes peptides that are assigned for more than one proteins.

We assume to use unique peptide for each protein.

removeFewMeasurements

TRUE (default) will remove the features that have 1 or 2 measurements across

runs.

removeProtein_with1Feature

TRUE will remove the proteins which have only 1 feature, which is the combi-

nation of peptide, precursor charge, fragment and charge. FALSE is default.

summaryforMultipleRows

max(default) or sum - when there are multiple measurements for certain feature

and certain run, use highest or sum of multiple intensities.

use_log_file logical. If TRUE, information about data processing will be saved to a file.

append logical. If TRUE, information about data processing will be added to an existing

log file.

verbose logical. If TRUE, information about data processing wil be printed to the con-

sole.

log_file_path character. Path to a file to which information about data processing will be

saved. If not provided, such a file will be created automatically. If 'append =

TRUE', has to be a valid path to a file.

... additional parameters to 'data.table::fread'.

Value

data.frame in the MSstats required format.

Author(s)

Meena Choi, Olga Vitek.

Examples

OpenSWATHtoMSstatsFormat

Import OpenSWATH files

Description

Import OpenSWATH files

Usage

```
OpenSWATHtoMSstatsFormat(
   input,
   annotation,
   filter_with_mscore = TRUE,
   mscore_cutoff = 0.01,
   useUniquePeptide = TRUE,
   removeFewMeasurements = TRUE,
   removeProtein_with1Feature = FALSE,
   summaryforMultipleRows = max,
   use_log_file = TRUE,
   append = FALSE,
   verbose = TRUE,
   log_file_path = NULL,
   ...
)
```

Arguments

input name of MSstats input report from OpenSWATH, which includes feature-level

 $annotation \qquad \quad name\ of\ `annotation.txt'\ data\ which\ includes\ Condition,\ BioReplicate,\ Run.\ Run$

should be the same as filename.

filter_with_mscore

TRUE(default) will filter out the features that have greater than mscore_cutoff in m_score column. Those features will be removed.

mscore_cutoff Cutoff for m_score. Default is 0.01. useUniquePeptide

TRUE (default) removes peptides that are assigned for more than one proteins. We assume to use unique peptide for each protein.

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removeFewMeasurements

TRUE (default) will remove the features that have 1 or 2 measurements across

removeProtein_with1Feature

TRUE will remove the proteins which have only 1 feature, which is the combination of peptide, precursor charge, fragment and charge. FALSE is default.

summaryforMultipleRows

max(default) or sum - when there are multiple measurements for certain feature

and certain run, use highest or sum of multiple intensities.

use_log_file logical. If TRUE, information about data processing will be saved to a file.

append logical. If TRUE, information about data processing will be added to an existing

log file.

verbose logical. If TRUE, information about data processing wil be printed to the con-

sole.

log_file_path character. Path to a file to which information about data processing will be

saved. If not provided, such a file will be created automatically. If 'append =

TRUE', has to be a valid path to a file.

... additional parameters to 'data.table::fread'.

Value

data.frame in the MSstats required format.

Author(s)

Meena Choi, Olga Vitek.

Examples

PDtoMSstatsFormat

Import Proteome Discoverer files

Description

Import Proteome Discoverer files

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Usage

```
PDtoMSstatsFormat(
  input,
  annotation,
  useNumProteinsColumn = FALSE,
  useUniquePeptide = TRUE,
  summaryforMultipleRows = max,
  removeFewMeasurements = TRUE,
  removeOxidationMpeptides = FALSE,
  removeProtein_with1Peptide = FALSE,
 which.quantification = "Precursor.Area",
 which.proteinid = "Protein.Group.Accessions",
 which.sequence = "Sequence",
  use_log_file = TRUE,
  append = FALSE,
  verbose = TRUE,
  log_file_path = NULL,
)
```

Arguments

input PD report or a path to it.

annotation name of 'annotation.txt' or 'annotation.csv' data which includes Condition, BioRepli-

cate, Run information. 'Run' will be matched with 'Spectrum.File'.

useNumProteinsColumn

TRUE removes peptides which have more than 1 in # Proteins column of PD output.

useUniquePeptide

TRUE (default) removes peptides that are assigned for more than one proteins. We assume to use unique peptide for each protein.

summaryforMultipleRows

max(default) or sum - when there are multiple measurements for certain feature and certain run, use highest or sum of multiple intensities.

removeFewMeasurements

TRUE (default) will remove the features that have 1 or 2 measurements across runs.

removeOxidationMpeptides

TRUE will remove the peptides including 'oxidation (M)' in modification. FALSE is default.

removeProtein_with1Peptide

TRUE will remove the proteins which have only 1 peptide and charge. FALSE is default.

which.quantification

Use 'Precursor.Area' (default) column for quantified intensities. 'Intensity' or 'Area' can be used instead.

which.proteinid

Use 'Protein. Accessions' (default) column for protein name. 'Master. Protein. Accessions'

can be used instead.

which.sequence Use 'Sequence' (default) column for peptide sequence. 'Annotated.Sequence'

can be used instead.

use_log_file logical. If TRUE, information about data processing will be saved to a file.

append logical. If TRUE, information about data processing will be added to an existing

log file.

verbose logical. If TRUE, information about data processing wil be printed to the con-

sole.

log_file_path character. Path to a file to which information about data processing will be

saved. If not provided, such a file will be created automatically. If 'append =

TRUE', has to be a valid path to a file.

.. additional parameters to 'data.table::fread'.

Value

data.frame in the MSstats required format.

Author(s)

Meena Choi, Olga Vitek

Examples

 ${\tt ProgenesistoMSstatsFormat}$

Import Progenesis files

Description

Import Progenesis files

Usage

```
ProgenesistoMSstatsFormat(
   input,
   annotation,
   useUniquePeptide = TRUE,
   summaryforMultipleRows = max,
   removeFewMeasurements = TRUE,
   removeOxidationMpeptides = FALSE,
   removeProtein_with1Peptide = FALSE,
   use_log_file = TRUE,
   append = FALSE,
   verbose = TRUE,
   log_file_path = NULL,
   ...
)
```

Arguments

input name of Progenesis output, which is wide-format. 'Accession', 'Sequence',

'Modification', 'Charge' and one column for each run are required.

annotation name of 'annotation.txt' or 'annotation.csv' data which includes Condition, BioRepli-

cate, Run information. It will be matched with the column name of input for MS

runs.

runs.

useUniquePeptide

TRUE (default) removes peptides that are assigned for more than one proteins.

We assume to use unique peptide for each protein.

summaryforMultipleRows

max(default) or sum - when there are multiple measurements for certain feature

and certain run, use highest or sum of multiple intensities.

removeFewMeasurements

TRUE (default) will remove the features that have 1 or 2 measurements across

removeOxidationMpeptides

TRUE will remove the peptides including 'oxidation (M)' in modification. FALSE

removeProtein_with1Peptide

TRUE will remove the proteins which have only 1 peptide and charge. FALSE

is default.

use_log_file logical. If TRUE, information about data processing will be saved to a file.

append logical. If TRUE, information about data processing will be added to an existing

log file.

verbose logical. If TRUE, information about data processing wil be printed to the con-

sole.

log_file_path character. Path to a file to which information about data processing will be

saved. If not provided, such a file will be created automatically. If 'append =

TRUE', has to be a valid path to a file.

.. additional parameters to 'data.table::fread'.

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Value

data.frame in the MSstats required format.

Author(s)

Meena Choi, Olga Vitek, Ulrich Omasits

Examples

quantification

Protein sample quantification or group quantification

Description

Model-based quantification for each condition or for each biological sample per protein in a targeted Selected Reaction Monitoring (SRM), Data-Dependent Acquisition (DDA or shotgun), and Data-Independent Acquisition (DIA or SWATH-MS) experiment. Quantification takes the processed data set by dataProcess as input and automatically generate the quantification results (data.frame) in a long or matrix format.

```
quantification(
  data,
  type = "Sample",
  format = "matrix",
  use_log_file = TRUE,
  append = FALSE,
  verbose = TRUE,
  log_file_path = NULL
)
```

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Arguments

data name of the (processed) data set.

type choice of quantification. "Sample" or "Group" for protein sample quantification

or group quantification.

format choice of returned format. "long" for long format which has the columns named

Protein, Condition, LogIntensities (and BioReplicate if it is subject quantification), NumFeature for number of transitions for a protein, and NumPeaks for number of observed peak intensities for a protein. "matrix" for data matrix format which has the rows for Protein and the columns, which are Groups(or Conditions) for group quantification or the combinations of BioReplicate and Condition (labeled by "BioReplicate"_"Condition") for sample quantification.

Default is "matrix"

use_log_file logical. If TRUE, information about data processing will be saved to a file.

append logical. If TRUE, information about data processing will be added to an existing

log file.

verbose logical. If TRUE, information about data processing wil be printed to the con-

sole.

log_file_path character. Path to a file to which information about data processing will be

saved. If not provided, such a file will be created automatically. If 'append =

TRUE', has to be a valid path to a file.

Details

Sample quantification: individual biological sample quantification for each protein. The label
of each biological sample is a combination of the corresponding group and the sample ID. If
there are no technical replicates or experimental replicates per sample, sample quantification
is the same as run summarization from dataProcess. If there are technical replicates or experimental replicates, sample quantification is median among run quantification corresponding
MS runs.

- Group quantification : quantification for individual group or individual condition per protein. It is median among sample quantification.
- The quantification for endogenous samples is based on run summarization from subplot model, with TMP robust estimation.

Value

data.frame as described in details.

- # Consider quantitative data (i.e. QuantData) from a yeast study with ten time points of
- # interests, three biological replicates, and no technical replicates which is
- # a time-course experiment.
- # Sample quantification shows model-based estimation of protein abundance in each biological
- # replicate within each time point.
- # Group quantification shows model-based estimation of protein abundance in each time point. QuantData<-dataProcess(SRMRawData, use_log_file = FALSE)

```
head(QuantData$FeatureLevelData)
# Sample quantification
sampleQuant<-quantification(QuantData, use_log_file = FALSE)
head(sampleQuant)
# Group quantification
groupQuant<-quantification(QuantData, type="Group", use_log_file = FALSE)
head(groupQuant)</pre>
```

savePlot

Save a plot to pdf file

Description

Save a plot to pdf file

Usage

```
savePlot(name_base, file_name, width, height)
```

Arguments

name_base path to a folder (or "" for working directory)

file_name name of a file to save. If this file already exists, an integer will be appended to

this name

width width of a plot height height of a plot

 ${\bf SkylinetoMSstatsFormat}$

Import Skyline files

Description

Import Skyline files

```
SkylinetoMSstatsFormat(
  input,
  annotation = NULL,
  removeiRT = TRUE,
  filter_with_Qvalue = TRUE,
  qvalue_cutoff = 0.01,
  useUniquePeptide = TRUE,
```

```
removeFewMeasurements = TRUE,
removeOxidationMpeptides = FALSE,
removeProtein_with1Feature = FALSE,
use_log_file = TRUE,
append = FALSE,
verbose = TRUE,
log_file_path = NULL,
...
)
```

Arguments

input name of MSstats input report from Skyline, which includes feature-level data.

annotation name of 'annotation.txt' data which includes Condition, BioReplicate, Run. If

annotation is already complete in Skyline, use annotation=NULL (default). It

will use the annotation information from input.

removeiRT TRUE (default) will remove the proteins or peptides which are labeld 'iRT' in

'StandardType' column. FALSE will keep them.

filter_with_Qvalue

TRUE(default) will filter out the intensities that have greater than qvalue_cutoff in DetectionQValue column. Those intensities will be replaced with zero and will be considered as censored missing values for imputation purpose.

qvalue_cutoff Cutoff for DetectionQValue. default is 0.01.

useUniquePeptide

TRUE (default) removes peptides that are assigned for more than one proteins. We assume to use unique peptide for each protein.

removeFewMeasurements

TRUE (default) will remove the features that have 1 or 2 measurements across runs

removeOxidationMpeptides

TRUE will remove the peptides including 'oxidation (M)' in modification. FALSE is default.

removeProtein_with1Feature

TRUE will remove the proteins which have only 1 feature, which is the combination of peptide, precursor charge, fragment and charge. FALSE is default.

use_log_file logical. If TRUE, information about data processing will be saved to a file.

append logical. If TRUE, information about data processing will be added to an existing

log file.

verbose logical. If TRUE, information about data processing wil be printed to the con-

sole.

log_file_path character. Path to a file to which information about data processing will be

saved. If not provided, such a file will be created automatically. If 'append =

TRUE', has to be a valid path to a file.

... additional parameters to 'data.table::fread'.

Value

data.frame in the MSstats required format.

Author(s)

Meena Choi, Olga Vitek

Examples

 ${\tt SpectronauttoMSstatsFormat}$

Import Spectronaut files

Description

Import Spectronaut files

```
SpectronauttoMSstatsFormat(
   input,
   annotation = NULL,
   intensity = "PeakArea",
   filter_with_Qvalue = TRUE,
   qvalue_cutoff = 0.01,
   useUniquePeptide = TRUE,
   removeFewMeasurements = TRUE,
   removeProtein_with1Feature = FALSE,
   summaryforMultipleRows = max,
   use_log_file = TRUE,
   append = FALSE,
   verbose = TRUE,
   log_file_path = NULL,
   ...
)
```

Arguments

input name of Spectronaut output, which is long-format. ProteinName, PeptideSe-

quence, PrecursorCharge, FragmentIon, ProductCharge, IsotopeLabelType, Condition, BioReplicate, Run, Intensity, F.ExcludedFromQuantification are required.

Rows with F.ExcludedFromQuantification=True will be removed.

annotation name of 'annotation.txt' data which includes Condition, BioReplicate, Run. If

annotation is already complete in Spectronaut, use annotation=NULL (default).

It will use the annotation information from input.

intensity 'PeakArea' (default) uses not normalized peak area. 'NormalizedPeakArea' uses

peak area normalized by Spectronaut.

filter_with_Qvalue

TRUE(default) will filter out the intensities that have greater than qvalue_cutoff in EG.Qvalue column. Those intensities will be replaced with zero and will be

considered as censored missing values for imputation purpose.

qvalue_cutoff Cutoff for EG.Qvalue. default is 0.01.

useUniquePeptide

TRUE (default) removes peptides that are assigned for more than one proteins. We assume to use unique peptide for each protein.

removeFewMeasurements

TRUE (default) will remove the features that have 1 or 2 measurements across

removeProtein_with1Feature

TRUE will remove the proteins which have only 1 feature, which is the combi-

nation of peptide, precursor charge, fragment and charge. FALSE is default.

summaryforMultipleRows

max(default) or sum - when there are multiple measurements for certain feature

and certain run, use highest or sum of multiple intensities.

use_log_file logical. If TRUE, information about data processing will be saved to a file.

append logical. If TRUE, information about data processing will be added to an existing

log file.

verbose logical. If TRUE, information about data processing wil be printed to the con-

sole.

log_file_path character. Path to a file to which information about data processing will be

saved. If not provided, such a file will be created automatically. If 'append =

TRUE', has to be a valid path to a file.

... additional parameters to 'data.table::fread'.

Value

data.frame in the MSstats required format.

Author(s)

Meena Choi, Olga Vitek

56 SRMRawData

Examples

SRMRawData

Example dataset from a SRM experiment with stable isotope labeled reference of a time course yeast study

Description

This is a partial data set obtained from a published study (Picotti, et. al, 2009). The experiment targeted 45 proteins in the glycolysis/gluconeogenesis/TCA cycle/glyoxylate cycle network, which spans the range of protein abundance from less than 128 to 10E6 copies per cell. Three biological replicates were analyzed at ten time points (T1-T10), while yeasts transited through exponential growth in a glucose-rich medium (T1-T4), diauxic shift (T5-T6), post-diauxic phase (T7-T9), and stationary phase (T10). Prior to trypsinization, the samples were mixed with an equal amount of proteins from the same N15-labeled yeast sample, which was used as a reference. Each sample was profiled in a single mass spectrometry run, where each protein was represented by up to two peptides and each peptide by up to three transitions. The goal of this study is to detect significantly change in protein abundance across time points. Transcriptional activity under the same experimental conditions has been previously investigated by (DeRisi et. al., 1997). Genes coding for 29 of the proteins are differentially expressed between conditions similar to those represented by T7 and T1 and could be treated as external sources to validate the proteomics analysis. In this exampled data set, two of the targeted proteins are selected and validated with gene expression study: Protein IDHC (gene name IDP2) is differentially expressed in time point 1 and time point 7, whereas, Protein PMG2 (gene name GPM2) is not. The protein names are based on Swiss Prot Name.

Usage

SRMRawData

Format

data.frame

Details

The raw data (input data for MSstats) is required to contain variable of ProteinName, PeptideSequence, PrecursorCharge, FragmentIon, ProductCharge, IsotopeLabelType, Condition, BioReplicate, Run, Intensity. The variable names should be fixed.

If the information of one or more columns is not available for the original raw data, please retain the column variables and type in fixed value. For example, the original raw data does not contain theme_msstats 57

the information of ProductCharge, we retain the column ProductCharge and type in NA for all transitions in RawData.

The column BioReplicate should label with unique patient ID (i.e., same patients should label with the same ID).

Variable Intensity is required to be original signal without any log transformation and can be specified as the peak of height or the peak of area under curve.

Value

data.frame with the required format of MSstats.

Author(s)

```
Meena Choi, Olga Vitek.

Maintainer: Meena Choi (<mnchoi67@gmail.com>)
```

References

Ching-Yun Chang, Paola Picotti, Ruth Huttenhain, Viola Heinzelmann-Schwarz, Marko Jovanovic, Ruedi Aebersold, Olga Vitek. Protein significance analysis in selected reaction monitoring (SRM) measurements. *Molecular & Cellular Proteomics*, 11:M111.014662, 2012.

Examples

```
head(SRMRawData)
```

theme_msstats

Theme for MSstats plots

Description

Theme for MSstats plots

```
theme_msstats(
  type,
  x.axis.size = 10,
  y.axis.size = 10,
  legend_size = 13,
  strip_background = element_rect(fill = "gray95"),
  strip_text_x = element_text(colour = c("black"), size = 14),
  legend_position = "top",
  legend_box = "vertical",
  text_angle = 0,
  text_hjust = NULL,
  text_vjust = NULL,
  ...
)
```

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Arguments

type of a plot

x.axis.size size of text on the x axis y.axis.size size of text on the y axis

legend_size size of the legend

strip_background

background of facet

strip_text_x size of text on facets

 ${\tt legend_position}$

position of the legend

legend_box legend.box

text_angle angle of text on the x axis (for condition and comparison plots)
text_hjust hjust parameter for x axis text (for condition and comparison plots)
text_vjust vjust parameter for x axis text (for condition and comparison plots)

... additional parameters passed on to ggplot2::theme()

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