# Package 'MSstatsTMT'

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**Title** Protein Significance Analysis in shotgun mass spectrometry-based proteomic experiments with tandem mass tag (TMT) labeling

**Version** 1.8.2 **Date** 2020-12-17

```
Description
      The package provides statistical tools for detecting differentially abundant proteins in shot-
     gun mass spectrometry-based proteomic experiments with tandem mass tag (TMT) labeling.
License Artistic-2.0
Depends R (>= 4.0)
Imports limma, lme4, lmerTest, dplyr, tidyr, statmod, methods,
     reshape2, data.table, matrixStats, stats, utils, ggplot2,
     grDevices, graphics, MSstats
Suggests BiocStyle, knitr, rmarkdown, testthat
VignetteBuilder knitr
biocViews ImmunoOncology, MassSpectrometry, Proteomics, Software
Encoding UTF-8
LazyData true
URL http://msstats.org/msstatstmt/
BugReports https://groups.google.com/forum/#!forum/msstats
RoxygenNote 7.1.1
git_url https://git.bioconductor.org/packages/MSstatsTMT
git_branch RELEASE_3_12
git_last_commit 5c9e714
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```

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# **R** topics documented:

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## **Description**

Annotation of example data, raw.mine, in this package. It should be prepared by users. The variables are as follows:

## Usage

annotation.mine

# **Format**

A data frame with 72 rows and 7 variables.

# **Details**

- Run: MS run ID. It should be the same as R.FileName info in raw.mine
- Channel: Labeling information (TMT6\_126, ..., TMT6\_131). The channels should be consistent with the channel columns in raw.mine.
- Condition: Condition (ex. Healthy, Cancer, Time0). If the channal doesn't have sample, please add 'Empty' under Condition.
- Mixture : Mixture of samples labeled with different TMT reagents, which can be analyzed in a single mass spectrometry experiment.

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• TechRepMixture: Technical replicate of one mixture. One mixture may have multiple technical replicates. For example, if 'TechRepMixture' = 1, 2 are the two technical replicates of one mixture, then they should match with same 'Mixture' value.

- Fraction: Fraction ID. One technical replicate of one mixture may be fractionated into multiple fractions to increase the analytical depth. Then one technical replicate of one mixture should correspond to multuple fractions. For example, if 'Fraction' = 1, 2, 3 are three fractions of the first technical replicate of one TMT mixture of biological subjects, then they should have same 'TechRepMixture' and 'Mixture' value.
- BioReplicate: Unique ID for biological subject. If the channal doesn't have sample, please add 'Empty' under BioReplicate

#### **Examples**

head(annotation.mine)

annotation.mq

Example of annotation file for evidence, which is the output of MaxQuant.

#### **Description**

Annotation of example data, evidence, in this package. It should be prepared by users. The variables are as follows:

# Usage

annotation.mq

#### **Format**

A data frame with 150 rows and 7 variables.

# **Details**

- Run: MS run ID. It should be the same as Raw.file info in raw.mq
- Channel: Labeling information (channel.0, ..., channel.9). The channel index should be consistent with the channel columns in raw.mq.
- Condition : Condition (ex. Healthy, Cancer, Time0)
- Mixture: Mixture of samples labeled with different TMT reagents, which can be analyzed in a single mass spectrometry experiment. If the channal doesn't have sample, please add 'Empty' under Condition.
- TechRepMixture: Technical replicate of one mixture. One mixture may have multiple technical replicates. For example, if 'TechRepMixture' = 1, 2 are the two technical replicates of one mixture, then they should match with same 'Mixture' value.
- Fraction: Fraction ID. One technical replicate of one mixture may be fractionated into multiple fractions to increase the analytical depth. Then one technical replicate of one mixture should correspond to multuple fractions. For example, if 'Fraction' = 1, 2, 3 are three fractions of the first technical replicate of one TMT mixture of biological subjects, then they should have same 'TechRepMixture' and 'Mixture' value.
- BioReplicate: Unique ID for biological subject. If the channal doesn't have sample, please add 'Empty' under BioReplicate.

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### **Examples**

head(annotation.mq)

annotation.pd

Example of annotation file for raw.pd, which is the PSM output of Proteome Discoverer

#### **Description**

Annotation of example data, raw.pd, in this package. It should be prepared by users. The variables are as follows:

# Usage

annotation.pd

#### **Format**

A data frame with 150 rows and 7 variables.

#### **Details**

- Run: MS run ID. It should be the same as Spectrum. File info in raw.pd.
- Channel: Labeling information (126, ... 131). It should be consistent with the channel columns in raw.pd.
- Condition: Condition (ex. Healthy, Cancer, Time0)
- Mixture: Mixture of samples labeled with different TMT reagents, which can be analyzed in a single mass spectrometry experiment. If the channal doesn't have sample, please add 'Empty' under Condition.
- TechRepMixture: Technical replicate of one mixture. One mixture may have multiple technical replicates. For example, if 'TechRepMixture' = 1, 2 are the two technical replicates of one mixture, then they should match with same 'Mixture' value.
- Fraction: Fraction ID. One technical replicate of one mixture may be fractionated into multiple fractions to increase the analytical depth. Then one technical replicate of one mixture should correspond to multuple fractions. For example, if 'Fraction' = 1, 2, 3 are three fractions of the first technical replicate of one TMT mixture of biological subjects, then they should have same 'TechRepMixture' and 'Mixture' value.
- BioReplicate: Unique ID for biological subject. If the channal doesn't have sample, please add 'Empty' under BioReplicate.

# **Examples**

head(annotation.pd)

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dataProcessPlotsTMT

Visualization for explanatory data analysis - TMT experiment

# **Description**

To illustrate the quantitative data and quality control of MS runs, dataProcessPlotsTMT takes the quantitative data from converter functions (PDtoMSstatsTMTFormat, MaxQtoMSstatsTMTFormat, SpectroMinetoMSstatsTMTFormat) as input and generate two 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.

# Usage

```
dataProcessPlotsTMT(
  data.peptide,
  data.summarization,
  type,
 ylimUp = FALSE,
 ylimDown = FALSE,
 x.axis.size = 10,
  y.axis.size = 10,
  text.size = 4,
  text.angle = 90,
  legend.size = 7,
  dot.size.profile = 2,
 ncol.guide = 5,
 width = 10,
 height = 10,
 which.Protein = "all",
  originalPlot = TRUE,
  summaryPlot = TRUE,
  address = ""
)
```

#### **Arguments**

data.peptide name of the data with peptide level, which can be the output of converter functions(PDtoMSstatsTMTFormat, MaxQtoMSstatsTMTFormat, SpectroMinetoMSstatsTMTFormat).

data.summarization

name of the data with protein-level, which can be the output of proteinSummarization

function.

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

across MS runs. "QCPlot" represents box plots of log intensities across channels

and MS runs.

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

QC Plot uses the upper limit as rounded off maximum of log2(intensities) after

normalization + 3..

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

Plot uses 0..

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

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

text.size size of labels represented each condition at the top of Profile plot and QC plot.

Default is 4.

text.angle angle of labels represented each condition at the top of Profile plot and QC plot.

Default is 0.

legend.size size of legend above Profile plot. Default is 7.

dot.size.profile

size of dots in Profile plot. Default is 2.

ncol.guide number of columns for legends at the top of plot. Default is 5.

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

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

Proteins. 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, without normalization.

summaryPlot TRUE(default) draws profile plots with protein summarization for each channel

and MS run.

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". 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.

# Value

plot or pdf

```
data(input.pd)
quant.msstats <- proteinSummarization(input.pd,</pre>
                                       method="msstats",
                                       global_norm=TRUE,
                                       reference_norm=TRUE)
## Profile plot
dataProcessPlotsTMT(data.peptide=input.pd,
                   data.summarization=quant.msstats,
                   type='ProfilePlot',
                   width = 21,
                   height = 7)
## NottoRun: QC plot
# dataProcessPlotsTMT(data.peptide=input.pd,
                    # data.summarization=quant.msstats,
                    # type='QCPlot',
                    # width = 21,
                    # height = 7)
```

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evidence

Example of output from MaxQuant for TMT-10plex experiments.

## **Description**

Example of evidence.txt from MaxQuant. It is the input for MaxQtoMSstatsTMTFormat function, with proteinGroups.txt and annotation file. Annotation file should be made by users. It includes peak intensities for 10 proteins among 15 MS runs with TMT10. The important variables are as follows:

## Usage

evidence

#### **Format**

A data frame with 1075 rows and 105 variables.

#### **Details**

- Proteins
- · Protein.group.IDs
- Modified.sequence
- Charge
- · Raw.file
- Score
- Potential.contaminant
- Reverse
- Channels: Reporter.intensity.corrected.0, ..., Reporter.intensity.corrected.9

# **Examples**

head(evidence)

 ${\tt groupComparisonTMT}$ 

Finding differentially abundant proteins across conditions in TMT experiment

# Description

Tests for significant changes in protein abundance across conditions based on a family of linear mixed-effects models in TMT experiment. Experimental design of case-control study (patients are not repeatedly measured) is automatically determined based on proper statistical model.

#### Usage

```
groupComparisonTMT(
  data,
  contrast.matrix = "pairwise",
  moderated = FALSE,
  adj.method = "BH",
  remove_norm_channel = TRUE,
  remove_empty_channel = TRUE)
```

#### **Arguments**

data

Name of the output of proteinSummarization function. It should have columns named Protein, Mixture, TechRepMixture, Run, Channel, Condition, BioReplicate, Abundance.

contrast.matrix

Comparison between conditions of interests. 1) default is 'pairwise', which compare all possible pairs between two conditions. 2) Otherwise, users can specify the comparisons 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.

 ${\tt moderated}$ 

TRUE will moderate t statistic; FALSE (default) uses ordinary t statistic.

adj.method

adjusted method for multiple comparison. "BH" is default.

remove\_norm\_channel

TRUE(default) removes 'Norm' channels from protein level data.

remove\_empty\_channel

TRUE(default) removes 'Empty' channels from protein level data.

#### Value

data.frame with result of inference

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```
colnames(comparison)<- c("0.125", "0.5", "0.667", "1") test.contrast <- groupComparisonTMT(data = quant.pd.msstats, contrast.matrix = comparison, moderated = TRUE)
```

input.pd

Example of output from PDtoMSstatsTMTFormat function

# Description

It is made from raw.pd and annotation.pd, which is the output of PDtoMSstatsTMTFormat function. It should include the required columns as below. The variables are as follows:

## Usage

input.pd

#### **Format**

A data frame with 20110 rows and 11 variables.

## **Details**

• ProteinName: Protein ID

• PeptideSequence : peptide sequence

• Charge: peptide charge

• PSM: peptide ion and spectra match

• Channel: Labeling information (126, ... 131)

• Condition: Condition (ex. Healthy, Cancer, Time0)

• BioReplicate: Unique ID for biological subject.

• Run: MS run ID

• Mixture : Unique ID for TMT mixture.

• TechRepMixture : Unique ID for technical replicate of one TMT mixture.

• Intensity: Protein Abundance

```
head(input.pd)
```

MaxQtoMSstatsTMTFormat

Generate MSstatsTMT required input format from MaxQuant output

## **Description**

Convert MaxQuant output into the required input format for MSstatsTMT.

## Usage

```
MaxQtoMSstatsTMTFormat(
  evidence,
  proteinGroups,
  annotation,
  which.proteinid = "Proteins",
  rmProt_Only.identified.by.site = FALSE,
  useUniquePeptide = TRUE,
  rmPSM_withMissing_withinRun = FALSE,
  rmPSM_withfewMea_withinRun = TRUE,
  rmProtein_with1Feature = FALSE,
  summaryforMultipleRows = sum
)
```

## **Arguments**

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

proteinGroups name of 'proteinGroups.txt' data.

annotation data frame which contains column Run, Fraction, TechRepMixture, Mixture,

Channel, BioReplicate, Condition. Refer to the example 'annotation.mq' for the

meaning of each column.

which.proteinid

Use 'Proteins' (default) column for protein name. 'Leading.proteins' or 'Leading.razor.protein' or 'Gene.names' can be used instead to get the protein ID with single protein. However, those can potentially have the shared peptides.

rmProt\_Only.identified.by.site

TRUE will remove proteins with '+' in 'Only.identified.by.site' column from proteinGroups.txt, which was identified only by a modification site. FALSE is the default.

use Unique Peptide

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

rmPSM\_withMissing\_withinRun

TRUE will remove PSM with any missing value within each Run. Defaut is FALSE.

rmPSM\_withfewMea\_withinRun

only for rmPSM\_withMissing\_withinRun = FALSE. TRUE(default) will remove the features that have 1 or 2 measurements within each Run.

rmProtein\_with1Feature

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

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```
summary for Multiple Rows
```

sum(default) or max - when there are multiple measurements for certain feature in certain run, select the feature with the largest summation or maximal value.

#### Value

input for proteinSummarization function

## **Examples**

```
head(evidence)
head(proteinGroups)
head(annotation.mq)
input.mq <- MaxQtoMSstatsTMTFormat(evidence, proteinGroups, annotation.mq)
head(input.mq)</pre>
```

MSstatsTMT

MSstatsTMT: A package for protein significance analysis in shotgun mass spectrometry-based proteomic experiments with tandem mass tag (TMT) labeling

## **Description**

A set of tools for detecting differentially abundant peptides and proteins in shotgun mass spectrometry-based proteomic experiments with tandem mass tag (TMT) labeling.

#### **functions**

- PDtoMSstatsTMTFormat : generates MSstatsTMT required input format for Proteome discoverer output.
- MaxQtoMSstatsTMTFormat: generates MSstatsTMT required input format for MaxQuant output.
- SpectroMinetoMSstatsTMTFormat : generates MSstatsTMT required input format for SpectroMine output.
- proteinSummarization: summarizes PSM level quantification to protein level quantification
- dataProcessPlotsTMT : visualizes for explanatory data analysis.
- groupComparisonTMT: tests for significant changes in protein abundance across conditions.

OpenMStoMSstatsTMTFormat

Generate MSstatsTMT required input format for OpenMS output

# Description

Convert OpenMS MSstatsTMT report into the required input format for MSstatsTMT.

#### Usage

```
OpenMStoMSstatsTMTFormat(
  input,
  useUniquePeptide = TRUE,
  rmPSM_withMissing_withinRun = FALSE,
  rmPSM_withfewMea_withinRun = TRUE,
  rmProtein_with1Feature = FALSE,
  summaryforMultiplePSMs = sum
)
```

## **Arguments**

input MSstatsTMT report from OpenMS

useUniquePeptide

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

rmPSM\_withMissing\_withinRun

TRUE will remove PSM with any missing value within each Run. Defaut is FALSE.

rmPSM\_withfewMea\_withinRun

only for rmPSM\_withMissing\_withinRun = FALSE. TRUE(default) will remove the features that have 1 or 2 measurements within each Run.

rmProtein\_with1Feature

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

 $\verb|summary| for \verb|MultiplePSMs|$ 

sum(default) or max - when there are multiple measurements for certain feature in certain run, select the feature with the largest summation or maximal value.

## Value

input for proteinSummarization function

```
head(raw.om)
input.om <- OpenMStoMSstatsTMTFormat(raw.om)
head(input.om)</pre>
```

PDtoMSstatsTMTFormat 13

 ${\it PDtoMSstatsTMTFormat} \quad \textit{Generate MSstatsTMT required input format from Proteome discoverer output}$ 

#### **Description**

Convert Proteome discoverer output into the required input format for MSstatsTMT.

# Usage

```
PDtoMSstatsTMTFormat(
   input,
   annotation,
   which.proteinid = "Protein.Accessions",
   useNumProteinsColumn = TRUE,
   useUniquePeptide = TRUE,
   rmPSM_withMissing_withinRun = FALSE,
   rmPSM_withfewMea_withinRun = TRUE,
   rmProtein_with1Feature = FALSE,
   summaryforMultipleRows = sum
)
```

#### **Arguments**

input data name of Proteome discover PSM output.

annotation data frame which contains column Run, Fraction, TechRepMixture, Mixture,

Channel, BioReplicate, Condition. Refer to the example 'annotation.pd' for the

meaning of each column.

which.proteinid

Use 'Protein.Accessions' (default) column for protein name. 'Master.Protein.Accessions' can be used instead to get the protein name with single protein.

useNumProteinsColumn

TURE(default) remove shared peptides by information of # Proteins column in PSM sheet.

useUniquePeptide

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

rmPSM\_withMissing\_withinRun

TRUE will remove PSM with any missing value within each Run. Defaut is FALSE.

rmPSM\_withfewMea\_withinRun

only for rmPSM\_withMissing\_withinRun = FALSE. TRUE(default) will remove the features that have 1 or 2 measurements within each Run.

rmProtein\_with1Feature

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

summaryforMultipleRows

sum(default) or max - when there are multiple measurements for certain feature in certain run, select the feature with the largest summation or maximal value.

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#### Value

input for proteinSummarization function

# **Examples**

```
head(raw.pd)
head(annotation.pd)
input.pd <- PDtoMSstatsTMTFormat(raw.pd, annotation.pd)
head(input.pd)</pre>
```

proteinGroups

Example of proteinGroups file from MaxQuant for TMT-10plex experiments.

# Description

Example of proteinGroup.txt file from MaxQuant, which is identified protein group information file. It is the input for MaxQtoMSstatsTMTFormat function, with evidence.txt and annotation file. It includes identified protein groups for 10 proteins among 15 MS runs with TMT10. The important variables are as follows:

# Usage

proteinGroups

#### **Format**

A data frame with 1075 rows and 105 variables.

# **Details**

- id
- Protein.IDs
- · Only.identified.by.site
- Potential.contaminant
- Reverse

# Examples

head(proteinGroups)

proteinSummarization 15

proteinSummarization Summarizing peptide level quantification to protein level quantifica-

#### **Description**

We assume missing values are censored and then impute the missing values. Protein-level summarization from peptide level quantification are performed. After all, global median normalization on peptide level data and normalization between MS runs using reference channels will be implemented.

# Usage

```
proteinSummarization(
  data,
  method = "msstats",
  global_norm = TRUE,
  reference_norm = TRUE,
  remove_norm_channel = TRUE,
  remove_empty_channel = TRUE,
  MBimpute = TRUE,
  maxQuantileforCensored = NULL
)
```

#### Arguments

data Name of the output of PDtoMSstatsTMTFormat function or peptide-level quan-

tified data from other tools. It should have columns ProteinName, PeptideSequence, Charge, PSM, Mixture, TechRepMixture, Run, Channel, Condition,

BioReplicate, Intensity

method Four different summarization methods to protein-level can be performed: "msstats" (default),

"MedianPolish", "Median", "LogSum".

global\_norm Global median normalization on peptide level data (equalizing the medians across

all the channels and MS runs). Default is TRUE. It will be performed before

protein-level summarization.

reference\_norm Reference channel based normalization between MS runs on protein level data.

TRUE(default) needs at least one reference channel in each MS run, annotated by 'Norm' in Condtion column. It will be performed after protein-level summarization. FALSE will not perform this normalization step. If data only has one

run, then reference\_norm=FALSE.

remove\_norm\_channel

TRUE(default) removes 'Norm' channels from protein level data.

remove\_empty\_channel

TRUE(default) removes 'Empty' channels from protein level data.

MBimpute only for method="msstats". TRUE (default) imputes missing va

only for method="msstats". TRUE (default) imputes missing values by Accelated failure model. FALSE uses minimum value to impute the missing value

for each peptide precursor ion.

 ${\tt maxQuantileforCensored}$ 

We assume missing values are censored. maxQuantileforCensored is Maximum quantile for deciding censored missing value, for instance, 0.999. Default is Null.

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#### Value

data.frame with protein-level summarization for each run and channel

# **Examples**

quant.pd.msstats

Example of output from proteinSummarizaiton function

# Description

It is made from input.pd. It is the output of proteinSummarization function. It should include the required columns as below. The variables are as follows:

# Usage

```
quant.pd.msstats
```

#### **Format**

A data frame with 100 rows and 8 variables.

## **Details**

Run : MS run IDProtein : Protein ID

• Abundance: Protein-level summarized abundance

• Channel: Labeling information (126, ... 131)

• Condition: Condition (ex. Healthy, Cancer, Time0)

• BioReplicate: Unique ID for biological subject.

• TechRepMixture : Unique ID for technical replicate of one TMT mixture.

• Mixture: Unique ID for TMT mixture.

```
head(quant.pd.msstats)
```

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raw.mine

Example of output from SpectroMine for TMT-6plex experiments.

#### **Description**

Example of SpectroMine PSM sheet. It is the output of SpectroMine and the input for SpectroMine-toMSstatsTMTFormat function, with annotation file. Annotation file should be made by users. It includes peak intensities for 10 proteins among 12 MS runs with TMT-6plex. The important variables are as follows:

# Usage

raw.mine

#### **Format**

A data frame with 170 rows and 28 variables.

#### **Details**

- PG.ProteinAccessions
- P.MoleculeID
- PP.Charge
- R.FileName
- PG.QValue
- PSM.Qvalue
- Channels: PSM.TMT6\_126..Raw., ..., PSM.TMT6\_131..Raw.

# **Examples**

head(raw.mine)

raw.om

Example of MSstatsTMT report from OpenMS for TMT-10plex experiments.

## **Description**

Example of MSstatsTMT PSM sheet from MaxQuant. It is the input for OpenMStoMSstatsTMT-Format function. It includes peak intensities for 10 proteins among 27 MS runs from three TMT10 mixtures. The important variables are as follows:

# Usage

raw.om

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#### **Format**

A data frame with 860 rows and 13 variables.

## **Details**

- RetentionTime
- ProteinName
- PeptideSequence
- Charge
- Channel
- Condition
- BioReplicate
- Run
- Mixture
- TechRepMixture
- Fraction
- Intensity
- Reference

# **Examples**

head(raw.om)

raw.pd

Example of output from Proteome Discoverer 2.2 for TMT-10plex experiments.

# Description

Example of Proteome discover PSM sheet. It is the input for PDtoMSstatsTMTFormat function, with annotation file. Annotation file should be made by users. It includes peak intensities for 10 proteins among 15 MS runs with TMT-10plex. The variables are as follows:

# Usage

raw.pd

# **Format**

A data frame with 2858 rows and 50 variables.

#### **Details**

- Master.Protein.Accessions
- · Protein.Accessions
- Annotated.Sequence
- Charge
- · Ions.Score
- Spectrum.File
- Quan.Info
- Channels: 126, ..., 131

#### **Examples**

```
head(raw.pd)
```

 ${\tt SpectroMinetoMSstatsTMTFormat}$ 

Generate MSstatsTMT required input format for SpectroMine output

## **Description**

Convert SpectroMine output into the required input format for MSstatsTMT.

# Usage

```
SpectroMinetoMSstatsTMTFormat(
  input,
  annotation,
  filter_with_Qvalue = TRUE,
  qvalue_cutoff = 0.01,
  useUniquePeptide = TRUE,
  rmPSM_withMissing_withinRun = FALSE,
  rmPSM_withfewMea_withinRun = TRUE,
  rmProtein_with1Feature = FALSE,
  summaryforMultipleRows = sum
)
```

# **Arguments**

input data name of SpectroMine PSM output. Read PSM sheet.

annotation data frame which contains column Run, Fraction, TechRepMixture, Mixture,

Channel, BioReplicate, Condition. Refer to the example 'annotation.mine' for

the meaning of each column.

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 NA and will be

considered as censored missing values for imputation purpose.

qvalue\_cutoff Cutoff for EG.Qvalue. default is 0.01.

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useUniquePeptide

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

rmPSM\_withMissing\_withinRun

TRUE will remove PSM with any missing value within each Run. Defaut is FALSE.

rmPSM\_withfewMea\_withinRun

only for rmPSM\_withMissing\_withinRun = FALSE. TRUE(default) will remove the features that have 1 or 2 measurements within each Run.

rmProtein\_with1Feature

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

summary for Multiple Rows

sum(default) or max - when there are multiple measurements for certain feature in certain run, select the feature with the largest summation or maximal value.

## Value

input for proteinSummarization function

#### **Examples**

```
head(raw.mine)
head(annotation.mine)
input.mine <- SpectroMinetoMSstatsTMTFormat(raw.mine, annotation.mine)
head(input.mine)</pre>
```

test.pairwise

Example of output from groupComparisonTMT function

# **Description**

It is the output of groupComparisonTMT function, which is the result of group comparions with the output of proteinSummarization function. It should include the columns as below. The variables are as follows:

#### Usage

test.pairwise

#### **Format**

A data frame with 60 rows and 7 variables.

# Details

• Protein: Protein ID

• Label: Label of the pairwise comparision or contrast

• log2FC: Log2 fold change

• SE: Standard error of the comparsion of contrast results

• DF: Degree of freedom

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- pvalue: Value of p statistic of the test
- adj.pvalue: adjusted p value

• issue: used for indicating the reason why a comparison is not testable. NA means the comparison is testable. 'oneConditionMissing' means the protein has no measurements in one conndition of the comparison. Furtherone, when 'issue = oneConditionMissing', 'log2FC = Inf' means the negative condition (with coefficient -1 in the Label column) is missing and 'log2FC = -Inf' means the positive condition (with coefficient 1 in the Label column) is missing. completeMissing' means the protein has no measurements in all the connditions of the comparison. unfittableModel' means there is no enough measurements to fit the linear model. In other words, each condition has only one measurement.

# **Examples**

head(test.pairwise)

# **Index**

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