# Package 'MSstatsPTM'

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Type Package

Title Statistical Characterization of Post-translational Modifications

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**Description** MSstatsPTM provides general statistical methods for quantitative characterization of post-translational modifications (PTMs). Supports DDA, DIA, SRM, and tandem mass tag (TMT) labeling. Typically, the analysis involves the quantification of PTM sites (i.e., modified residues) and their corresponding proteins, as well as the integration of the quantification results. MSstatsPTM provides functions for summarization, estimation of PTM site abundance, and detection of changes in PTMs across experimental conditions.

License Artistic-2.0

**Depends** R (>= 4.3)

Imports dplyr, gridExtra, stringr, stats, ggplot2, stringi, grDevices, MSstatsTMT, MSstatsConvert, MSstats, data.table, Rcpp, Biostrings, checkmate, ggrepel

**Suggests** knitr, rmarkdown, tinytest, covr, mockery, testthat (>= 3.0.0)

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BugReports https://github.com/Vitek-Lab/MSstatsPTM/issues

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Author Devon Kohler [aut, cre], Tsung-Heng Tsai [aut], Ting Huang [aut], Mateusz Staniak [aut], Meena Choi [aut], Olga Vitek [aut]

 ${\bf Maintainer}\ \ {\bf Devon}\ Kohler\, \verb|<kohler.d@northeastern.edu>|$ 

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# Description

Power calculation for PTM experiment

# Usage

```
.calculatePowerPTM(
  desiredFC,
  FDR,
  delta,
  ptm_median_sigma_error,
  protein_median_sigma_error,
  ptm_median_sigma_subject,
  protein_median_sigma_subject,
  numSample
)
```

# Arguments

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.

# Value

float of power

.fixTerminus

Add site location and aa

# Description

Add site location and aa

# Usage

```
.fixTerminus(data, terminus_id, unmod_pep_col)
```

# Arguments

data data.table

fasta\_file string or data.table

#### Value

data.table

.getNumSamplePTM

Get sample size for PTM experiment

# Description

Get sample size for PTM experiment

# Usage

```
.getNumSamplePTM(
  desiredFC,
  power,
  alpha,
  delta,
  ptm_median_sigma_error,
  protein_median_sigma_error,
  ptm_median_sigma_subject,
  protein_median_sigma_subject
```

.joinFasta 5

# Arguments

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

the desired fold change.

power 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

#### Value

int of samples

.joinFasta

Add FASTA data into dataframe

# Description

Add FASTA data into dataframe

# Usage

```
.joinFasta(
  data,
  fasta_file,
  fasta_protein_name,
  protein_name_col,
  unmod_pep_col,
  mod_pep_col
)
```

# **Arguments**

data data.table

fasta\_file string or data.table

# Value

data.table

6 .removeCutoffSites

.locateSites

Add site location and aa

# **Description**

Add site location and aa

# Usage

```
.locateSites(
  data,
  mod_id,
  protein_name_col,
  unmod_pep_col,
  mod_pep_col,
  mod_id_is_numeric,
  replace_text = FALSE
)
```

# **Arguments**

data data.table mod\_id string

#### Value

data.table

 $. \\ remove Cut off Sites$ 

Remove sites below cutoff probability

# Description

Remove sites below cutoff probability

# Usage

```
.removeCutoffSites(data, mod_pep_col, cutoff, remove_unlocalized_peptides)
```

# **Arguments**

data data.table

mod\_pep\_col column in data with modified sites cutoff numeric cutoff. Default is .75.

remove\_unlocalized\_peptides

Boolean if to remove peptides that could not be fully localized.

annotSite 7

# Value

data.table with modifications below cutoff removed

annotSite

Annotate modification site

# **Description**

annotSite annotates modified sites as their residues and locations.

# Usage

```
annotSite(aaIndex, residue, lenIndex = NULL)
```

# Arguments

aaIndex An integer vector. Location of the sites.
residue A string vector. Amino acid residue.

lenIndex An integer. Default is NULL

# Value

A string.

# **Examples**

```
annotSite(10, "K")
annotSite(10, "K", 3L)
```

dataProcessPlotsPTM

Visualization for explanatory data analysis

# **Description**

To illustrate the quantitative data and quality control of MS runs, dataProcessPlotsPTM takes the quantitative data from dataSummarizationPTM or dataSummarizationPTM\_TMT to plot the following: (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.

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# Usage

```
dataProcessPlotsPTM(
  data,
  type = "PROFILEPLOT",
 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 = 12,
 ptm.title = "All PTMs",
 protein.title = "All Proteins",
 which.PTM = "all",
 which.Protein = NULL,
 originalPlot = TRUE,
  summaryPlot = TRUE,
 address = ""
)
```

# Arguments

data	name of the list with PTM and (optionally) Protein data, which can be the output of the MSstatsPTM dataSummarizationPTM or dataSummarizationPTM_TMT functions.
type	choice of visualization. "ProfilePlot" represents profile plot of log intensities 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
ylimDown	lower limit for y-axis in the log scale. FALSE(Default) for Profile Plot and QC Plot uses 0
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.profi	le
	size of dots in Profile plot. Default is 2.

dataSummarizationPTM 9

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.

ptm.title title of overall PTM QC plot protein.title title of overall Protein QC plot

which.PTM PTM list to draw plots. List can be names of PTMs or order numbers of PTMs.

Default is "all", which generates all plots for each protein. For QC plot, "allonly"

will generate one QC plot with all proteins.

which.Protein List of proteins to plot. Will plot all PTMs associated with listed Proteins. De-

fault is NULL which will default to which.PTM.

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

#### **Examples**

dataSummarizationPTM Data summarization function for label-free MS experiments targeting PTMs.

10 dataSummarizationPTM

### **Description**

Utilizes functionality from MSstats to clean, summarize, and normalize PTM and protein level data. Imputes missing values, performs normalization, and summarizes data. PTM data is summarized up to the modification and protein data is summarized up to the protein level. Takes as input the output of the included converters (see included raw.input data object for required input format).

### Usage

```
dataSummarizationPTM(
  data,
  logTrans = 2,
  normalization = "equalizeMedians",
  normalization.PTM = "equalizeMedians",
  nameStandards = NULL,
  nameStandards.PTM = NULL,
  featureSubset = "all",
  featureSubset.PTM = "all",
  remove_uninformative_feature_outlier = FALSE,
  remove_uninformative_feature_outlier.PTM = FALSE,
 min_feature_count = 2,
 min_feature_count.PTM = 1,
  n_{top_feature} = 3,
  n_{top_feature.PTM} = 3,
  summaryMethod = "TMP",
  equalFeatureVar = TRUE,
  censoredInt = "NA",
 MBimpute = TRUE,
 MBimpute.PTM = TRUE,
  remove50missing = FALSE,
  fix_missing = NULL,
  maxQuantileforCensored = 0.999,
  use_log_file = TRUE,
  append = TRUE,
  verbose = TRUE,
  log_file_path = NULL,
  base = "MSstatsPTM_log_"
)
```

#### **Arguments**

data name of the list with PTM and (optionally) unmodified protein data.tables, which

can be the output of the MSstatsPTM converter functions

logTrans logarithm transformation with base 2(default) or 10

normalization normalization for the protein level dataset, 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

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based on reference signals is performed. 'globalStandards' represents normalization with global standards proteins. FALSE represents no normalization is performed

normalization.PTM

normalization for PTM level dataset. Default is "equalizeMedians" Can be adjusted to any of the options described above.

nameStandards

vector of global standard peptide names for protein dataset. only for normalization with global standard peptides.

nameStandards.PTM

Same as above for PTM dataset.

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.

featureSubset.PTM

For PTM dataset only. Options same as above.

remove\_uninformative\_feature\_outlier

For protein dataset only. It only works after users used featureSubset="highQuality" in dataProcess. 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, for run-level summarization. FALSE (default) uses all features and intensities for run-level summarization.

remove\_uninformative\_feature\_outlier.PTM

For PTM dataset only. Options same as above.

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.

min\_feature\_count.PTM

For PTM dataset only. Options the same as above. Default is 1 due to low average feature count for PTMs.

n\_top\_feature For protein dataset only. The number of top features for featureSubset='topN'. Default is 3, which means to use top 3 features.

n\_top\_feature.PTM

For PTM dataset only. Options same as above.

summaryMethod

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

MBimpute For protein dataset only. 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.

MBimpute.PTM For PTM dataset only. Options same as above.

remove50missing

only for summaryMethod="TMP". TRUE removes the runs which have more

than 50% missing values. FALSE is default.

fix\_missing Default is Null. Optional, same as the 'fix\_missing' parameter in MSstatsCon-

vert::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.

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

log file.

verbose logical. If TRUE, information about data processing will 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.

base start of the file name.

### Value

list of summarized PTM and Protein results. These results contain the reformatted input to the summarization function, as well as run-level summarization results.

#### **Examples**

head(raw.input\$PTM)
head(raw.input\$PROTEIN)

 ${\tt quant.lf.msstatsptm = dataSummarizationPTM(raw.input, verbose = FALSE)}$ 

head(quant.lf.msstatsptm\$PTM\$ProteinLevelData)

#### dataSummarizationPTM\_TMT

Data summarization function for TMT labelled MS experiments targeting PTMs.

#### **Description**

Utilizes functionality from MSstatsTMT to clean, summarize, and normalize PTM and protein level data. Imputes missing values, performs normalization, and summarizes data. PTM data is summarized up to the modification and protein data is summarized up to the protein level. Takes as input the output of the included converters (see included raw.input.tmt data object for required input format).

#### Usage

```
dataSummarizationPTM_TMT(
  data,
  method = "msstats",
  global_norm = TRUE,
  global_norm.PTM = TRUE,
  reference_norm = TRUE,
  reference_norm.PTM = TRUE,
  remove_norm_channel = TRUE,
  remove_empty_channel = TRUE,
 MBimpute = TRUE,
 MBimpute.PTM = TRUE,
  maxQuantileforCensored = NULL,
  use_log_file = TRUE,
  append = FALSE,
  verbose = TRUE,
  log_file_path = NULL
)
```

### Arguments

data Name of the output of MSstatsPTM converter function or peptide-level quan-

tified data from other tools. It should be a list containing one or two data tables, named PTM and PROTEIN for modified and unmodified datasets. The list must at least contain the PTM dataset. The data should have columns Protein-Name, PeptideSequence, Charge, PSM, Mixture, TechRepMixture, Run, Chan-

nel, 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 for unmodified peptide level data (equalizing

the medians across all the channels and MS runs). Default is TRUE. It will be

performed before protein-level summarization.

global\_norm.PTM

Same as above for modified peptide level data. Default is TRUE.

Same as above for modified peptide level data. Belaute is TROL

reference\_norm Reference channel based normalization between MS runs on unmodified 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.

reference\_norm.PTM

Same as above for modified peptide level data. Default is TRUE.

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 values by Acce-

lated failure model. FALSE uses minimum value to impute the missing value

for each peptide precursor ion.

MBimpute.PTM Same as above for modified peptide level data. Default is TRUE

maxQuantileforCensored

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

Null.

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 will 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.

#### Value

list of two data.tables

#### **Examples**

designSampleSizePTM

Planning future experimental designs of PTM experiments in sample size calculation

#### **Description**

Calculate sample size for future experiments of a PTM experiment based on intensity-based linear model. Calculation is only available for group comparison experimental designs (not including time series). Two options of the calculation: (1) number of biological replicates per condition, (2) power.

#### Usage

```
designSampleSizePTM(
  data,
  desiredFC,
  FDR = 0.05,
  numSample = TRUE,
  power = 0.8,
  use_log_file = TRUE,
  append = FALSE,
  verbose = TRUE,
  log_file_path = NULL,
  base = "MSstatsPTM_log_"
)
```

#### **Arguments**

data	output of the group(	ComparisonPTM function.

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.

power 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 will 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.

base start of the file name.

#### 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.

#### **Examples**

DIANNtoMSstatsPTMFormat

Convert the output of DIA-NN PSM file into MSstatsPTM format

### Description

Takes as input the report.tsv file from DIA-NN and converts it into MSstatsPTM format. Requires PSM and an annotation file. Optionally an additional report.tsv file for a corresponding global profiling run can be included.

#### Usage

```
DIANNtoMSstatsPTMFormat(
  input,
  annotation,
  input_protein = NULL,
  annotation_protein = NULL,
  fasta_path = NULL,
  use_unmod_peptides = FALSE,
  protein_id_col = "Protein.Group";
  fasta_protein_name = "uniprot_ac",
  global_qvalue_cutoff = 0.01,
  qvalue_cutoff = 0.01,
  pg_qvalue_cutoff = 0.01,
  useUniquePeptide = TRUE,
  removeFewMeasurements = TRUE,
  removeOxidationMpeptides = TRUE,
  removeProtein_with1Feature = FALSE,
  MBR = TRUE,
```

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

#### **Arguments**

input data.frame of report.tsv file produced by Philosopher

annotation annotation with Run, Fraction, TechRepMixture, Mixture, Channel, BioRepli-

cate, Condition columns or a path to file. Refer to the example 'annotation' for

the meaning of each column.

input\_protein same as input for global profiling run. Default is NULL.

annotation\_protein

same as annotation for global profiling run. Default is NULL.

fasta\_path A string of path to a FASTA file, used to match PTM peptides.

use\_unmod\_peptides

Boolean if the unmodified peptides in the input file should be used to construct the unmodified protein output. Only used if input\_protein is not provided. Default is FALSE.

protein\_id\_col Use 'Protein.Groups'(default) column for protein name.

fasta\_protein\_name

Name of column that matches with the protein names in protein\_id\_col. The protein names in these two columns must match in order to join the FASTA file with the DIA-NN output.

global\_qvalue\_cutoff

The global qualue cutoff. Default is 0.01.

gvalue\_cutoff local gvalue cutoff for library. Default is 0.01.

pg\_qvalue\_cutoff

local qvalue cutoff for protein groups Run should be the same as filename. Default is 0.01.

useUniquePeptide

logical, if 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 within each Run.

removeOxidationMpeptides

TRUE (default) will remove the peptides including oxidation (M) sequence.

removeProtein\_with1Feature

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

MBR If analoysis was done with match between runs or not. Default is TRUE.

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.

#### Value

list of one or two data. frame of class MSstatsTMT, named PTM and PROTEIN

#### **Examples**

FragPipetoMSstatsPTMFormat

Convert output of TMT labeled Fragpipe data into MSstatsPTM format.

#### **Description**

Takes as input TMT experiments which are the output of Fragpipe and converts into MSstatsPTM format. Requires msstats.csv file and an annotation file. Optionally an additional msstats.csv file can be uploaded if a corresponding global profiling run was performed. Site localization is performed and only high probability localizations are kept.

#### Usage

```
FragPipetoMSstatsPTMFormat(
   input,
   annotation = NULL,
   input_protein = NULL,
   annotation_protein = NULL,
   use_unmod_peptides = FALSE,
   label_type = "TMT",
   protein_id_col = "Protein",
   peptide_id_col = "Peptide.Sequence",
   mod_id_col = "STY",
```

```
localization_cutoff = 0.75,
  remove_unlocalized_peptides = TRUE,
 Purity_cutoff = 0.6,
 PeptideProphet_prob_cutoff = 0.7,
  useUniquePeptide = TRUE,
  rmPSM_withfewMea_withinRun = FALSE,
  rmPeptide_OxidationM = TRUE,
  rmProtein_with1Feature = FALSE,
  summaryforMultipleRows = sum,
  use_log_file = TRUE,
  append = FALSE,
  verbose = TRUE,
  log_file_path = NULL
)
```

#### **Arguments**

input data.frame of msstats.csv file produced by Philosopher

annotation

annotation with Run, Fraction, TechRepMixture, Mixture, Channel, BioReplicate, Condition columns or a path to file. Refer to the example 'annotation' for the meaning of each column. Channel column should be consistent with the channel columns (Ignore the prefix "Channel") in msstats.csv file. Run column should be consistent with the Spectrum. File columns in msstats.csv file.

same as input for global profiling run. Default is NULL. input\_protein annotation\_protein

same as annotation for global profiling run. Default is NULL.

use\_unmod\_peptides

Boolean if the unmodified peptides in the input file should be used to construct the unmodified protein output. Only used if input\_protein is not provided. Default is FALSE.

label\_type

Type of labeling used for experiment. Must be one of "LF" or "TMT". Default is "TMT".

protein\_id\_col Use 'Protein'(default) column for TMT. This needs to be changed to "Protein-Name" for label free. For TMT, 'Master.Protein.Accessions' can be used instead to get the protein ID with single protein.

peptide\_id\_col Use 'Peptide.Sequence'(default) column for TMT. Must be changed to "PeptideSequence" for label free. "Modified.Peptide.Sequence" can be used instead to get the modified peptide sequence.

mod\_id\_col

Column containing the modified Amino Acids. For example, a Phosphorylation experiment may pass STY. The corresponding column with STY combined with the mass (e.x. STY. 79.9663) will be selected. Default is STY.

localization\_cutoff

Minimum localization score required to keep modification. Default is .75.

remove\_unlocalized\_peptides

Boolean indicating if peptides without all sites localized should be kept. Default is TRUE (non-localized sites will be removed).

Purity\_cutoff

Cutoff for purity. Default is 0.6. Purity refers to how much of the detected ion signal within a specific inclusion window belongs to the target molecule or its closely related forms, compared to any other unwanted signals or noise. Higher values indicate greater purity.

PeptideProphet\_prob\_cutoff

Cutoff for the peptide identification probability. Default is 0.7. The probability is confidence score determined by PeptideProphet and higher values indicate greater confidence.

useUniquePeptide

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

rmPSM\_withfewMea\_withinRun

TRUE (default) will remove the features that have 1 or 2 measurements within each Run.

rmPeptide\_OxidationM

TRUE (default) will remove the peptides including oxidation (M) sequence.

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.

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.

#### Value

list of one or two data. frame of class MSstatsTMT, named PTM and PROTEIN

# **Examples**

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```
mod_id_col = "STY",
                                          localization_cutoff=.75,
                                          remove_unlocalized_peptides=TRUE)
head(msstats_data$PTM)
head(msstats_data$PROTEIN)
# LFQ Example (w/out global profiling run)
input = system.file("tinytest/raw_data/Fragpipe/MSstats.csv",
                                        package = "MSstatsPTM")
input = data.table::fread(input)
annot = system.file("tinytest/raw_data/Fragpipe/experiment_annotation.tsv",
                                        package = "MSstatsPTM")
annot = data.table::fread(annot)
msstats_data = FragPipetoMSstatsPTMFormat(input,
                                          label_type="LF",
                                          mod_id_col = "STY"
                                          localization_cutoff=.75,
                                          protein_id_col = "ProteinName",
                                          peptide_id_col = "PeptideSequence")
head(msstats_data$PTM)
# Note that this is NULL because we did not include a global profiling run.
# Ideally, you should include an independent global profiling run.
head(msstats_data$PROTEIN)
```

 $fragpipe\_annotation$ 

Example annotation file for a TMT FragPipe experiment.

# Description

Automatically created by FragPipe, manually checked by the user and input into the FragPipetoMSstatsPTMFormat converter. Requires the correct columns and maps the experimental desing into the MSstats format. Specify unique bioreplicates for group comparison designs, and the same bioreplicate for repeated measure designs. The columns and descriptions are below.

# Usage

fragpipe\_annotation

#### **Format**

A data.table with 7 columns.

#### **Details**

- Run: Run name that matches exactly with FragPipe run. Used to join evidence and metadata in annotation file.
- Fraction: If multiple fractions were used (i.e. the same mixture split into multiple fractions) enter that here. TechRepMixture: Multiple runs using the same bioreplicate
- · Channel: Mixture channel used
- Condition: Name of condition that was used for each run.
- Mixture: The unique mixture (plex) name
- BioReplicate: Name of biological replicate. Repeating the same name here will tell MSstat-sPTM that the experiment is a repeated measure design.

# **Examples**

head(fragpipe\_annotation)

fragpipe\_annotation\_protein

Example annotation file for a global profiling run TMT FragPipe experiment.

#### **Description**

Automatically created by FragPipe, manually checked by the user and input into the FragPipetoMSstatsPTMFormat converter. Requires the correct columns and maps the experimental desing into the MSstats format. Specify unique bioreplicates for group comparison designs, and the same bioreplicate for repeated measure designs. The columns and descriptions are below.

# Usage

fragpipe\_annotation\_protein

#### **Format**

A data.table with 7 columns.

#### **Details**

- Run: Run name that matches exactly with FragPipe run. Used to join evidence and metadata
  in annotation file.
- Fraction: If multiple fractions were used (i.e. the same mixture split into multiple fractions) enter that here. TechRepMixture: Multiple runs using the same bioreplicate
- Channel: Mixture channel used
- Condition: Name of condition that was used for each run.
- Mixture: The unique mixture (plex) name
- BioReplicate: Name of biological replicate. Repeating the same name here will tell MSstat-sPTM that the experiment is a repeated measure design.

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# **Examples**

head(fragpipe\_annotation\_protein)

fragpipe\_input

Output of FragPipe TMT PTM experiment

# **Description**

This dataset was provided by the FragPipe team at the Nesvilab. It was processed using Philosopher and targeted Phosphorylation.

# Usage

fragpipe\_input

#### **Format**

A data.table with 29 columns and 246 rows.

#### **Examples**

head(fragpipe\_input)

fragpipe\_input\_protein

Output of FragPipe TMT global profiling experiment

# Description

This dataset was provided by the FragPipe team at the Nesvilab. It was processed using Philosopher and targeted Phosphorylation.

# Usage

```
fragpipe_input_protein
```

#### **Format**

A data.table with 27 columns and 47 rows.

#### **Examples**

head(fragpipe\_input\_protein)

groupComparisonPlotsPTM

Visualization for model-based analysis and summarization

# Description

To analyze the results of modeling changes in abundance of modified peptides and overall protein, groupComparisonPlotsPTM takes as input the results of the groupComparisonPTM function. It asses the results of three models: unadjusted PTM, adjusted PTM, and overall protein. To asses the results of the model, the following visualizations can be created: (1) VolcanoPlot (specify "VolcanoPlot" in option type), to plot peptides or proteins and their significance for each model. (2) Heatmap (specify "Heatmap" in option type), to evaluate the fold change between conditions and peptides/proteins

# Usage

```
groupComparisonPlotsPTM(
  data = 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 = 50,
 width = 10,
  height = 10,
 which.Comparison = "all",
 which.PTM = "all",
  address = ""
)
```

# **Arguments**

name of the list with models, which can be the output of the MSstatsPTM groupComparisonPTM function

type choice of visualization, one of VolcanoPlot or Heatmap

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.

y.axis.size size of axes labels, e.g. name of targeted proteins in heatmap. Default is 10.

dot.size size of dots in volcano plot and comparison plot. Default is 3.

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.

colorkey TRUE(default) shows colorkey.

numProtein The number of proteins which will be presented in each heatmap. Default is 50.

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

 $numbers\ of\ comparisons\ from\ levels (data\$Label)\ ,\ such\ as\ levels (testResultMultiComparisons\$Comparisons§Comparisons$ 

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

which.PTM 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 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 "VolcanoPlot.pdf" or "Heatmap.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

### **Examples**

groupComparisonPTM

Perform differential analysis on MS-based proteomics experiments targeting PTMs

#### **Description**

Takes summarized PTM and protein data from dataSummarizationPTM or dataSummarizationPTM\_TMT functions and performs differential analysis. Leverages unmodified protein data to perform adjustment and deconvolute the effect of the PTM and unmodified protein. If protein data is unavailable, PTM data can still be passed into the function, however adjustment can not be performed. All model results are returned for completeness.

# Usage

```
groupComparisonPTM(
  data,
  data.type,
  contrast.matrix = "pairwise",
  moderated = FALSE,
  adj.method = "BH",
  log_base = 2,
  use_log_file = TRUE,
  append = FALSE,
  verbose = TRUE,
  log_file_path = NULL,
  base = "MSstatsPTM_log_"
)
```

locateMod 27

# Arguments

data	list of summarized datasets. Output of MSstatsPTM summarization function dataSummarizationPTM or dataSummarizationPTM_TMT depending on acquisition type.
data.type	string indicating experimental acquisition type. "TMT" is used for TMT labeled experiments. For all other experiments (Label Free/ DDA/ DIA) use "Label-Free".
contrast.matri	X
	comparison between conditions of interests. Default models full pairwise comparison between all conditions
moderated	For TMT experiments only. TRUE will moderate t statistic; FALSE (default) uses ordinary t statistic. Default is FALSE.
adj.method	For TMT experiemnts only. Adjusted method for multiple comparison. "BH" is default. "BH" is used for all other experiment types
log_base	For non-TMT experiments only. The base of the logarithm used in summarization.
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 will 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.
base	start of the file name.

# Value

list of modeling results. Includes PTM, PROTEIN, and ADJUSTED data.tables with their corresponding model results.

# **Examples**

locateMod	Locate modified sites with a peptide	

# Description

locateMod locates modified sites with a peptide.

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#### Usage

```
locateMod(peptide, aaStart, residueSymbol)
```

# Arguments

peptide A string. Peptide sequence.

aaStart An integer. Starting index of the peptide.

residueSymbol A string. Modification residue and denoted symbol.

#### Value

A string.

#### **Examples**

```
locateMod("P*EP*TIDE", 3, "\\*")
```

locatePTM Annotate modified sites with associated peptides

# **Description**

PTMlocate annotates modified sites with associated peptides.

# Usage

locatePTM(peptide, uniprot, fasta, modResidue, modSymbol, rmConfound = FALSE)

# **Arguments**

peptide A string vector of peptide sequences. The peptide sequence does not include its

preceding and following AAs.

uniprot A string vector of Uniprot identifiers of the peptides' originating proteins. UniPro-

tKB entry isoform sequence is used.

fasta A data.table with FASTA information. Output of tidyFasta.

modResidue A string. Modifiable amino acid residues.

modSymbol A string. Symbol of a modified site.

rmConfound A logical. TRUE removes confounded unmodified sites, FALSE otherwise. De-

fault is FALSE.

#### Value

A data frame with three columns: uniprot\_iso, peptide, site.

### **Examples**

```
fasta = tidyFasta(system.file("extdata", "013297.fasta", package="MSstatsPTM"))
locatePTM("DRVSYIHNDSC*TR", "013297", fasta, "C", "\\*")
```

MaxQtoMSstatsPTMFormat

Convert output of label-free or TMT MaxQuant experiments into MSstatsPTM format

### Description

Takes as input LF/TMT experiments from MaxQ and converts the data into the format needed for MSstatsPTM. Requires modified evidence.txt file from MaxQ and an annotation file for PTM data. To adjust modified peptides for changes in global protein level, unmodified TMT experimental data must also be returned. Optionally can use Phospho(STY)Sites.txt (or other PTM specific files) from MaxQuant, but this is not recommended. If PTM specific file provided, the raw intensities must be provided, not a ratio.

### Usage

```
MaxQtoMSstatsPTMFormat(
  evidence = NULL,
  annotation = NULL,
  fasta_path,
  fasta_protein_name = "uniprot_ac",
  mod_id = "\(Phospho \(STY\))",
  sites_data = NULL,
  evidence_prot = NULL,
  proteinGroups = NULL,
  annotation_protein = NULL,
  use_unmod_peptides = FALSE,
  labeling_type = "LF",
  mod_num = "Single"
  TMT_keyword = "TMT"
  ptm_keyword = "phos",
  which_proteinid_ptm = "Proteins",
  which_proteinid_protein = "Proteins",
  remove_other_mods = 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 for enriched (PTM)

data.

annotation data frame annotation file for the ptm level data. Contains column Run, Fraction,

TechRepMixture, Mixture, Channel, BioReplicate, Condition.

fasta\_path A string of path to a FASTA file, used to match PTM peptides.

fasta\_protein\_name

Name of fasta column that matches with protein name in evidence file. Default

is uniprot\_ac.

mod\_id Character that indicates the modification of interest. Default is \\(Phospho\\\).

Note \\ must be included before special characters.

sites\_data (Not recommended. Only used if evidence file not provided. Only works for

TMT labeled data) Modified peptide output from MaxQuant. For example, a phosphorylation experiment would require the Phospho(STY)Sites.txt file

evidence\_prot name of 'evidence.txt' data, which includes feature-level data for global profil-

ing (unmodified) data.

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

evidence\_prot.

annotation\_protein

data frame annotation file for the protein level data. Contains column Run, Frac-

tion, TechRepMixture, Mixture, Channel, BioReplicate, Condition.

use\_unmod\_peptides

Boolean if the unmodified peptides in the input file should be used to construct the unmodified protein output. Only used if input\_protein is not provided.

Default is FALSE.

labeling\_type Either TMT or LF (Label-Free) depending on experimental design. Default is LF.

mod\_num (Only if sites.data is used) For modified peptide dataset. The number modifi-

cations per peptide to be used. If "Single", only peptides with one modification will be used. Otherwise "Total" can be selected which does not cap the number of modifications per peptide. "Single" is the default. Selecting "Total" may

confound the effect of different modifications.

TMT\_keyword (Only if sites.data is used) the sub-name of columns in sites.data file. Default

is TMT. This corresponds to the columns in the format Reporter.intensity.corrected.1.TMT1phos\_\_\_

Specifically, this parameter indicates the first section of the string TMT1phos (Before the mixture number). If TMT is present in the string, set this value to TMT. Else if TMT is not there (ie string is in the format 1phos) leave this parameter as

an empty string (").

ptm\_keyword (Only if sites.data is used) the sub-name of columns in the sites.data file. De-

fault is phos. This corresponds to the columns in the format Reporter.intensity.corrected.1.TMT1pl

Specifically, this parameter indicates the second section of the string TMT1phos (After the mixture number). If the string is present, set this parameter. Else if this part of the string is empty (ie string is in the format TMT1) leave this parameter.

eter as an empty string (").

which\_proteinid\_ptm

For PTM dataset, which column to use for protein name. 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.

which\_proteinid\_protein

For Protein dataset, which column to use for protein name. Same options as above.

remove\_other\_mods

Remove peptides which include modifications other than the one listed in mod\_id. Default is TRUE. For example, in an experiment targeting Phosphorylation, setting this parameter to TRUE would remove peptides like (Acetyl (Protein Nterm))AAAAPDSRVS(Phospho (STY))EEENLK. Set this parameter to FALSE to keep peptides with extraneous modifications.

removeMpeptides

If Oxidation (M) modifications should be removed. Default is TRUE.

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.

### Value

a list of two data.tables named 'PTM' and 'PROTEIN' in the format required by MSstatsPTM.

# **Examples**

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```
which_proteinid_ptm = "Proteins")
head(msstats_format_tmt$PTM)
head(msstats_format_tmt$PROTEIN)
# LF experiment
head(maxq_lf_evidence)
head(maxq_lf_annotation)
msstats_format_lf = MaxQtoMSstatsPTMFormat(evidence=maxq_lf_evidence,
                        annotation=maxq_lf_annotation,
                fasta=system.file("extdata", "maxq_lf_fasta.fasta", package="MSstatsPTM"),
                        fasta_protein_name="uniprot_ac",
                        mod_id="\\(Phospho \\(STY\\)\\)",
                        use_unmod_peptides=TRUE,
                        labeling_type = "LF",
                        which_proteinid_ptm = "Proteins")
head(msstats_format_lf$PTM)
head(msstats_format_lf$PROTEIN)
```

maxq\_lf\_annotation

Example annotation file for a label-free MaxQuant experiment.

# Description

Must be manually created by the user and input into the MaxQtoMSstatsPTMFormat converter. Requires the correct columns and maps the experimental desing into the MSstats format. Specify unique bioreplicates for group comparison designs, and the same bioreplicate for repeated measure designs. The columns and descriptions are below.

#### Usage

 ${\tt maxq\_lf\_annotation}$ 

### Format

A data.table with 5 columns.

#### **Details**

- Run: Run name that matches exactly with MaxQuant run. Used to join evidence and metadata
  in annotation file.
- Condition: Name of condition that was used for each run.
- BioReplicate: Name of biological replicate. Repeating the same name here will tell MSstat-sPTM that the experiment is a repeated measure design.
- Raw.file: Run name that matches exactly with MaxQuant run. Used to join evidence and metadata in annotation file.
- IsotopeLabelType: Name of isotope label. May be all L or unique depending on experimental design.

maxq\_lf\_evidence 33

### **Examples**

head(maxq\_lf\_annotation)

maxq\_lf\_evidence

Example MaxQuant evidence file from the output of a label free experiment

#### **Description**

Experiment was performed by the Olsen lab and published on Nat. Commun. (citation below).

#### Usage

maxq\_lf\_evidence

#### **Format**

a data.table with 63 columns and 511 rows, the output of MaxQuant

#### **Details**

Bekker-Jensen, D.B., Bernhardt, O.M., Hogrebe, A. et al. Rapid and site-specific deep phosphoproteome profiling by data-independent acquisition without the need for spectral libraries. Nat Commun 11, 787 (2020). https://doi.org/10.1038/s41467-020-14609-1

The experiment was processed using MaxQuant by the computational proteomics team at Pfizer (Liang Xue and Pierre Jean).

The experiment did not contain a global profiling run, but we show an example of extracting the unmodified peptides and using them in place of the profiling run.

#### **Examples**

head(maxq\_lf\_evidence)

maxq\_tmt\_annotation

Example annotation file for a TMT MaxQuant experiment.

# **Description**

Must be manually created by the user and input into the MaxQtoMSstatsPTMFormat converter. Requires the correct columns and maps the experimental desing into the MSstats format. Specify unique bioreplicates for group comparison designs, and the same bioreplicate for repeated measure designs. The columns and descriptions are below.

### Usage

maxq\_tmt\_annotation

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#### **Format**

A data.table with 7 columns.

#### **Details**

- Run: Run name that matches exactly with MaxQuant run. Used to join evidence and metadata in annotation file.
- Fraction: If multiple fractions were used (i.e. the same mixture split into multiple fractions) enter that here. TechRepMixture: Multiple runs using the same bioreplicate
- Channel: Mixture channel used
- Condition: Name of condition that was used for each run.
- Mixture: The unique mixture (plex) name
- BioReplicate: Name of biological replicate. Repeating the same name here will tell MSstat-sPTM that the experiment is a repeated measure design.

# **Examples**

head(maxq\_tmt\_annotation)

maxq\_tmt\_evidence

Example MaxQuant evidence file from the output of a TMT experiment

### **Description**

Experiment was performed by the Olsen lab and published on Nat. Commun. (citation below).

#### Usage

maxq\_tmt\_evidence

#### **Format**

a data.table with 96 columns and 199 rows, the output of MaxQuant

# **Details**

Hogrebe, A., von Stechow, L., Bekker-Jensen, D.B. et al. Benchmarking common quantification strategies for large-scale phosphoproteomics. Nat Commun 9, 1045 (2018). https://doi.org/10.1038/s41467-018-03309-6

The experiment was processed using MaxQuant by the computational proteomics team at Pfizer (Liang Xue and Pierre Jean).

The experiment did not contain a global profiling run, but we show an example of extracting the unmodified peptides and using them in place of the profiling run.

# **Examples**

head(maxq\_tmt\_evidence)

 ${\tt MetamorpheusToMSstatsPTMFormat}$ 

Import Metamorpheus files into PTM format

### Description

Import Metamorpheus files into PTM format

### Usage

```
MetamorpheusToMSstatsPTMFormat(
  input,
  annotation,
  fasta_path,
  input_protein = NULL,
  annotation_protein = NULL,
  use_unmod_peptides = FALSE,
 mod_ids = c("\local{Common Biological:Phosphorylation on S}"),
  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 Metamorpheus output file, which is tabular format. Use the AllQuanti-

fiedPeaks.tsv file from the Metamorpheus output.

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

fasta\_path string containing path to the corresponding fasta file for the modified peptide

dataset.

input\_protein same as input for global profiling run. Default is NULL.

annotation\_protein

same as annotation for global profiling run. Default is NULL.

use\_unmod\_peptides

mod\_ids

If protein\_input is not provided, unmodified peptides can be extracted from

input to be used in place of a global profiling run. Default is FALSE.

Please note that the 'mod\_ids' parameter currently supports lists of size 1 only. Future updates aim to extend its functionality to accommodate lists of greater

List of modifications of interest. Default is a list with only Common Biological: Phosphorylation on S.

sizes. Note \\ must be included before special characters.

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 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.

#### Value

a list of two data.tables named 'PTM' and 'PROTEIN' in the format required by MSstatsPTM.

#### Author(s)

Anthony Wu

### **Examples**

```
input = system.file("tinytest/raw_data/Metamorpheus/AllQuantifiedPeaks.tsv",
                                package = "MSstatsPTM")
input = data.table::fread(input)
annot = system.file("tinytest/raw_data/Metamorpheus/ExperimentalDesign.tsv",
                                package = "MSstatsPTM")
annot = data.table::fread(annot)
input_protein = system.file("tinytest/raw_data/Metamorpheus/AllQuantifiedPeaksGlobalProteome.tsv",
                                package = "MSstatsPTM")
input_protein = data.table::fread(input_protein)
annot_protein = system.file("tinytest/raw_data/Metamorpheus/ExperimentalDesignGlobalProteome.tsv",
                                package = "MSstatsPTM")
annot_protein = data.table::fread(annot_protein)
fasta_path=system.file("extdata", "metamorpheus_fasta.fasta",
                                package="MSstatsPTM")
metamorpheus_imported = MetamorpheusToMSstatsPTMFormat(
   input,
   annot,
   fasta_path=fasta_path,
```

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```
input_protein=input_protein,
  annotation_protein=annot_protein,
  use_unmod_peptides=FALSE,
  mod_ids = c("\\[Common Fixed:Carbamidomethyl on C\\]")
)
head(metamorpheus_imported$PTM)
head(metamorpheus_imported$PROTEIN)
```

**MSstatsPTM** 

MSstatsPTM: A package for detecting differentially abundant posttranslational modifications (PTM) in mass spectrometry-based proteomic experiments.

# Description

A set of tools for detecting differentially abundant PTMs and proteins in shotgun mass spectrometry-based proteomic experiments. The package can handle a variety of acquisition types, including label free and TMT experiments, acquired with DDA, DIA, SRM or PRM acquisition methods. The package includes tools to convert raw data from different spectral processing tools, summarize feature intensities, and fit a linear mixed effects model. A major advantage of the package is to leverage a separate global profiling run and adjust the PTM fold change for changes in the unmodified protein, showing the unconvoluted PTM fold change. Finally, the package includes functionality to plot a variety of data visualizations.

#### **functions**

- FragPipetoMSstatsPTMFormat : Generates MSstatsPTM required input format for TMT FragePipe outputs.
- MaxQtoMSstatsPTMFormat : Generates MSstatsPTM required input format for label-free and TMT MaxQuant outputs.
- ProgenesistoMSstatsPTMFormat: Generates MSstatsPTM required input format for label-free Progenesis outputs.
- SpectronauttoMSstatsPTMFormat : Generates MSstatsPTM required input format for label-free Spectronaut outputs.
- SkylinetoMSstatsPTMFormat: Generates MSstatsPTM required input format for Skyline outputs.
- PStoMSstatsPTMFormat: Generates MSstatsPTM required input format for PEAKS outputs.
- PDtoMSstatsPTMFormat : Generates MSstatsPTM required input format for Proteome Discoverer outputs.
- dataSummarizationPTM: Summarizes PSM level quantification to peptide (modification) and protein level quantification. For use in label-free analysis
- dataSummarizationPTM\_TMT : Summarizes PSM level quantification to peptide (modification) and protein level quantification. For use in TMT analysis.
- dataProcessPlotsPTM: Visualization for explanatory data analysis. Specifically gives ability to plot Profile and Quality Control plots.

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- groupComparisonPTM: Tests for significant changes in PTM and protein abundance across conditions. Adjusts PTM fold change for changes in protein abundance.
- groupComparisonPlotsPTM: Visualization for model-based analysis and summarization

# Author(s)

**Maintainer**: Devon Kohler <kohler.d@northeastern.edu> Authors:

- Tsung-Heng Tsai <tsai.tsungheng@gmail.com>
- Ting Huang <thuang0703@gmail.com>
- Mateusz Staniak <mtst@mstaniak.pl>
- Meena Choi <mnchoi67@gmail.com>
- Olga Vitek <o.vitek@northeastern.edu>

#### See Also

Useful links:

• Report bugs at https://github.com/Vitek-Lab/MSstatsPTM/issues

MSstatsPTMSiteLocator Locate modification site number and amino acid

## Description

Locate modification site number and amino acid

```
MSstatsPTMSiteLocator(
  data,
  protein_name_col = "ProteinName",
  unmod_pep_col = "PeptideSequence",
  mod_pep_col = "PeptideModifiedSequence",
  clean_mod = FALSE,
  fasta_file = NULL,
  fasta_protein_name = "header",
  mod_id = " \ *",
  localization_scores = FALSE,
  localization_cutoff = 0.75,
  remove_unlocalized_peptides = TRUE,
  terminus_included = FALSE,
  terminus_id = "\\.",
  mod_id_is_numeric = FALSE,
  remove_underscores = FALSE,
```

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```
remove_other_mods = FALSE,
bracket = FALSE,
replace_text = FALSE
)
```

#### **Arguments**

data

data.table of enriched experimental run. Must include ProteinName, PeptideSequence, PeptideModifiedSequence, and (optionally) Start columns.

protein\_name\_col

Name of column indicating protein. Default is ProteinName.

unmod\_pep\_col Name of column indicating unmodified peptide sequence. Default is PeptideSequence.

mod\_pep\_col Name of column indicating modified peptide sequence. Default is PeptideModifiedSequence.

clean\_mod Remove special characters and numbers around modification name. Default is FALSE

fasta\_file File path to FASTA file that matches with proteins in data. Can be either string or data. table processed with tidyFasta() function. Default to NULL if pep-

tide number included in data.

fasta\_protein\_name

Name of fasta file column that matches with protein\_name\_col. Default is header

mod\_id String that indicates what amino acid was modified in PeptideSequence.

localization\_scores

Boolean indicating if mod id is a localization score. If TRUE, mod\_id will be ignored and localization cutoff will be used to determine sites. Default is FALSE.

localization\_cutoff

Default is .75. Localization probabilities below cutoffs will be removed. localization\_scores must be TRUE.

remove\_unlocalized\_peptides

Default is TRUE. If localization\_scores is TRUE and probabilities are below localization\_cutoff, the modification site will not be able to be determined. These unlocalized peptides can be kept or removed. If FALSE the unlocalized peptides will still be used in modeling the sites that could be localized.

terminus\_included

Boolean indicating if the PeptideSequence includes the terminus amino acid.

terminus\_id String that indicates what the terminus amino acid is. Default is '.'.

mod\_id\_is\_numeric

Boolean indicating if modification identifier is a number instead of a character (i.e. +80 vs \*).

remove\_underscores

Boolean indicating if underscores around peptide exist. These should be removed to properly count where in sequence the modification occurred.

remove\_other\_mods

keeping mods that are not of interest can mess up the amino acid count. Remove them if they are causing issues.

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bracket type that encompasses PTM (usually [ or (). Always pass opening

bracket (there is a function to grab the close bracket). Default is FALSE (i.e. no

bracket).

replace\_text If PTM is noted by text (i.e. Phospho) and needs to be replaced by an indicator

(\*)

#### Value

data.table with site location added into Protein column.

# **Examples**

##TODO

PDtoMSstatsPTMFormat Convert Proteome Discoverer output into MSstatsPTM format

# **Description**

Import Proteome Discoverer files, identify modification site location.

```
PDtoMSstatsPTMFormat(
  input,
  annotation,
  fasta_path,
  protein_input = NULL,
  annotation_protein = NULL,
  labeling_type = "LF",
  mod_id = "\(Phospho\)",
  use_localization_cutoff = FALSE,
  keep_all_mods = FALSE,
  use_unmod_peptides = FALSE,
  fasta_protein_name = "uniprot_iso",
  localization_cutoff = 75,
  remove_unlocalized_peptides = TRUE,
  useNumProteinsColumn = FALSE,
  useUniquePeptide = TRUE,
  summaryforMultipleRows = max,
  removeFewMeasurements = TRUE,
  removeOxidationMpeptides = FALSE,
  removeProtein_with1Peptide = FALSE,
  which_quantification = "Precursor.Area",
  which_proteinid = "Protein.Group.Accessions",
  use_log_file = TRUE,
```

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

## **Arguments**

input PD report corresponding with enriched experimental data.

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

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

fasta\_path string containing path to the corresponding fasta file for the modified peptide

dataset.

protein\_input PD report corresponding with unmodified experimental data.

annotation\_protein

Same format as annotation corresponding to unmodified data.

labeling\_type type of experimental design, must be one of LF for label free or TMT for tandem

mass tag.

mod\_id Character that indicates the modification of interest. Default is \\((Phospho\\)).

Note \\ must be included before special characters.

use\_localization\_cutoff

Boolean indicating whether to use a custom localization cutoff or rely on PD's modifications column. TRUE is default and apply custom cutoff localization\_cutoff.

keep\_all\_mods Boolean indicating whether to keep or remove peptides not in mod\_id. Default

is FALSE.

use\_unmod\_peptides

If protein\_input is not provided, unmodified peptides can be extracted from input to be used in place of a global profiling run. Default is EALSE

input to be used in place of a global profiling run. Default is FALSE.

fasta\_protein\_name

Name of fasta column that matches with protein name in evidence file. Default is uniprot\_iso.

localization\_cutoff

Minimum localization score required to keep modification. Default is .75.

remove\_unlocalized\_peptides

Boolean indicating if peptides without all sites localized should be kept. Default is TRUE (non-localized sites will be removed).

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.

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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.

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 will 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.

#### Value

list of data.table

#### **Examples**

```
# Global profiling example
input = system.file("tinytest/raw_data/PD/pd-ptm-input.csv",
package = "MSstatsPTM")
input = data.table::fread(input)
annot = system.file("tinytest/raw_data/PD/pd-ptm-annot.csv",
                    package = "MSstatsPTM")
annot = data.table::fread(annot)
input_protein = system.file("tinytest/raw_data/PD/protein-input.csv",
                            package = "MSstatsPTM")
input_protein = data.table::fread(input_protein)
annot_protein = system.file("tinytest/raw_data/PD/protein-annot.csv",
                            package = "MSstatsPTM")
annot_protein = data.table::fread(annot_protein)
fasta_path=system.file("extdata", "pd_with_proteome.fasta",
                       package="MSstatsPTM")
pd_imported = PDtoMSstatsPTMFormat(
    input,
    annotation = annot,
   protein_input = input_protein,
```

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```
annotation_protein = annot_protein,
     fasta_path = fasta_path,
     mod_id = "\\(GG\\)",
     labeling_type = "TMT",
     use_localization_cutoff = FALSE,
     which_proteinid = "Master.Protein.Accessions")
 head(pd_imported$PTM)
 head(pd_imported$PROTEIN)
 # No global profiling example
 head(pd_psm_input)
 head(pd_annotation)
 msstats_format = PDtoMSstatsPTMFormat(pd_psm_input,
                                        pd_annotation,
                            system.file("extdata", "pd_fasta.fasta", package="MSstatsPTM"),
                                        use_unmod_peptides=TRUE,
                                        which_proteinid = "Master.Protein.Accessions")
 head(msstats_format$PTM)
 head(msstats_format$PROTEIN)
pd_annotation
                         Example annotation file for a label-free Proteome Discoverer experi-
```

# **Description**

Must be manually created by the user and input into the PDtoMSstatsPTMFormat converter. Requires the correct columns and maps the experimental desing into the MSstats format. Specify unique bioreplicates for group comparison designs, and the same bioreplicate for repeated measure designs. The columns and descriptions are below.

## Usage

pd\_annotation

#### **Format**

A data table with 3 columns.

#### **Details**

- Run: Run name that matches exactly with PD run. Used to join evidence and metadata in annotation file.
- Condition: Name of condition that was used for each run.
- BioReplicate: Name of biological replicate. Repeating the same name here will tell MSstat-sPTM that the experiment is a repeated measure design.

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#### **Examples**

head(pd\_annotation)

pd\_psm\_input

Example Proteome Discoverer evidence file from the output of a label free experiment

# **Description**

Experiment was performed by the Olsen lab and published on Nat. Commun. (citation below).

#### Usage

```
pd_psm_input
```

#### **Format**

a data.table with 60 columns and 1657 rows, the output of PD

#### **Details**

Bekker-Jensen, D.B., Bernhardt, O.M., Hogrebe, A. et al. Rapid and site-specific deep phosphoproteome profiling by data-independent acquisition without the need for spectral libraries. Nat Commun 11, 787 (2020). https://doi.org/10.1038/s41467-020-14609-1

The experiment was processed using Proteome Discoverer by the computational proteomics team at Pfizer (Liang Xue and Pierre Jean).

The experiment did not contain a global profiling run, but we show an example of extracting the unmodified peptides and using them in place of the profiling run.

#### **Examples**

head(pd\_psm\_input)

pd\_testing\_output

Example output of Proteome Discoverer converter

# Description

output using example data provided in package

```
pd_testing_output
```

#### **Format**

a list with 2 data.frames

#### **Details**

The experiment did not contain a global profiling run, but we show an example of extracting the unmodified peptides and using them in place of the profiling run.

## **Examples**

```
head(pd_testing_output)
```

 ${\tt ProgenesistoMSstatsPTMFormat}$ 

Converts non-TMT Progenesis output into the format needed for MSstatsPTM

#### **Description**

Converts non-TMT Progenesis output into the format needed for MSstatsPTM

#### Usage

```
ProgenesistoMSstatsPTMFormat(
   ptm_input,
   annotation,
   global_protein_input = FALSE,
   fasta_path = FALSE,
   useUniquePeptide = TRUE,
   summaryforMultipleRows = max,
   removeFewMeasurements = TRUE,
   removeOxidationMpeptides = FALSE,
   removeProtein_with1Peptide = FALSE,
   mod.num = "Single"
)
```

#### **Arguments**

ptm\_input

name of Progenesis output with modified peptides, 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, BioReplicate, Run, and Type (PTM or Protein) information. It will be matched with the column name of input for MS runs. Please note PTM and global Protein run names are often different, which is why an additional Type column indicating Protein or PTM is required.

global\_protein\_input

name of Progenesis output with unmodified peptides, which is wide-format. 'Accession', Sequence', 'Modification', 'Charge' and one column for each run are required

fasta\_path

string containing path to the corresponding fasta file for the modified peptide dataset.

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.

remove Oxidation Mpeptides

TRUE will remove the modified peptides including 'Oxidation (M)' sequence. FALSE is default.

removeProtein\_with1Peptide

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

mod.num

For modified peptide dataset, must be one of Single or Total. The default is Single. The number modifications per peptide to be used. If "Single", only peptides with one modification will be used. Otherwise "Total" includes peptides with more than one modification. Selecting "Total" may confound the effect of different modifications.

## Value

a list of two data.tables named 'PTM' and 'PROTEIN' in the format required by MSstatsPTM.

#### **Examples**

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PStoMSstatsPTMFormat Convert Peaks Studio output into MSstatsPTM format

#### Description

Currently only supports label-free quantification.

# Usage

```
PStoMSstatsPTMFormat(
   input,
   annotation,
   input_protein = NULL,
   annotation_protein = NULL,
   use_unmod_peptides = FALSE,
   target_modification = NULL,
   remove_oxidation_peptides = FALSE,
   remove_multi_mod_types = FALSE,
   summaryforMultipleRows = max,
   use_log_file = TRUE,
   append = FALSE,
   verbose = TRUE,
   log_file_path = NULL
)
```

#### **Arguments**

input name of Peaks Studio PTM output

annotation name of annotation file which includes Raw.file, Condition, BioReplicate, Run.

For example annotation see example below.

input\_protein name of Peaks Studio unmodified protein output (optional)

annotation\_protein

name of annotation file which includes Raw.file, Condition, BioReplicate, Run for unmodified protein output.

use\_unmod\_peptides

Boolean if the unmodified peptides in the input file should be used to construct the unmodified protein output. Only used if input\_protein is not provided. Default is FALSE

target\_modification

Character name of modification of interest. To use all mod types, leave as NULL. Default is NULL. Note that if the name includes special characters, you must include "\" before the characters. Ex. "Phosphorylation \(STY\)"

remove\_oxidation\_peptides

Boolean if Oxidation (M) modifications should be removed. Default is FALSE

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remove\_multi\_mod\_types

Used if target\_modification is not NULL. TRUE will remove peptides with multiple types of modifications (ie acetylation and phosphorylation). FALSE will keep these peptides and summarize them seperately.

summary for Multiple Rows

 $\mbox{max}(\mbox{default})$  or  $\mbox{sum}$  - 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.

#### Value

list of data.table

# **Examples**

# The output should be in the following format.

head(raw.input\$PTM)
head(raw.input\$PROTEIN)

raw.input

Example of input PTM dataset for LabelFree/DDA/DIA experiments.

# Description

It can be the output of MSstatsPTM converter ProgenesistoMSstatsPTMFormat or other MSstats converter functions (Please see MSstatsPTM\_LabelFree\_Workflow vignette). The dataset is formatted as a list with two data.tables named PTM and PROTEIN. In each data.table the variables are as follows:

# Usage

raw.input

#### **Format**

A list of two data.tables named PTM and PROTEIN with 1745 and 478 rows respectively.

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## **Details**

#'

ProteinName : Name of protein with modification site mapped in with an underscore. ie  $"Protein\_4\_Y474"$ 

- PeptideSequence
- Condition : Condition (ex. Healthy, Cancer, Time0)
- BioReplicate: Unique ID for biological subject.
- Run: MS run ID.
- Intensity
- PrecursorCharge
- FragmentIon
- ProductCharge
- IsotopeLabelType

# **Examples**

head(raw.input\$PTM)
head(raw.input\$PROTEIN)

raw.input.tmt

Example of input PTM dataset for TMT experiments.

# **Description**

It can be the output of MSstatsPTM converter MaxQtoMSstatsPTMFormat or other MSstatsTMT converter functions (Please see MSstatsPTM\_TMT\_Workflow vignette). The dataset is formatted as a list with two data.tables named PTM and PROTEIN. In each data.table the variables are as follows:

#### Usage

raw.input.tmt

#### **Format**

A list of two data.tables named PTM and PROTEIN with 1716 and 29221 rows respectively.

#### **Details**

- ProteinName: Name of protein with modification site mapped in with an underscore. ie "Protein\_4\_Y474"
- PeptideSequence
- Charge
- PSM
- 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. \item TechRepMixture: Technical replicate of one mixture. One mixture may Mixture' = 1, 2 are the two technical replicates of one mixture, then they should match with same Mixture' value. \item Run: MS run ID. \item Channel: Labeling information (126, ... 131). \ite under BioReplicate.
- Intensity

## **Examples**

```
head(raw.input.tmt$PTM)
head(raw.input.tmt$PROTEIN)
```

SkylinetoMSstatsPTMFormat

Convert Skyline output into MSstatsPTM format

# Description

Currently only supports label-free quantification.

```
SkylinetoMSstatsPTMFormat(
   input,
   fasta_path,
   fasta_protein_name = "uniprot_iso",
   annotation = NULL,
   input_protein = NULL,
   annotation_protein = NULL,
   use_unmod_peptides = FALSE,
   removeiRT = TRUE,
   filter_with_Qvalue = TRUE,
   qvalue_cutoff = 0.01,
   use_unique_peptide = TRUE,
   remove_few_measurements = FALSE,
   remove_oxidation_peptides = FALSE,
   removeProtein_with1Feature = FALSE,
```

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

#### **Arguments**

input name of Skyline PTM output

fasta\_path A string of path to a FASTA file, used to match PTM peptides.

fasta\_protein\_name

Name of fasta column that matches with protein name in evidence file. Default

is uniprot\_iso.

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.

input\_protein name of Skyline unmodified protein output (optional)

annotation\_protein

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

unmodified protein output. This can be the same as annotation.

use\_unmod\_peptides

Boolean if the unmodified peptides in the input file should be used to construct the unmodified protein output. Only used if input\_protein is not provided.

Default is FALSE.

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.

use\_unique\_peptide

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

We assume to use unique peptide for each protein.

remove\_few\_measurements

TRUE will remove the features that have 1 or 2 measurements across runs. FALSE is default.

remove\_oxidation\_peptides

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.

#### Value

list of data.table

# **Examples**

```
# The output should be in the following format.
head(raw.input$PTM)
head(raw.input$PROTEIN)
```

 ${\tt SpectronauttoMSstatsPTMFormat}$ 

Convert Spectronaut output into MSstatsPTM format

## **Description**

Converters label-free Spectronaut data into MSstatsPTM format. Requires PSM output from Spectronaut and a custom made annotation file, mapping the run name to the condition and bioreplicate. Can optionally take a seperate PSM file for a global profiling run. If no global profiling run provided, the function can extract the unmodified peptides from the PTM PSM file and use them as a global profiling run (not recommended).

```
SpectronauttoMSstatsPTMFormat(
  input,
  annotation = NULL,
  fasta_path = NULL,
  protein_input = NULL,
  annotation_protein = NULL,
  use_unmod_peptides = FALSE,
  intensity = "PeakArea",
  mod_id = "\\[Phospho \\(STY\\)\\]",
  fasta_protein_name = "uniprot_iso",
  remove_other_mods = TRUE,
  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 PTM output, which is long-format. ProteinName, Pep-

tideSequence, 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.

fasta\_path string containing path to the corresponding fasta file for the modified peptide

dataset.

protein\_input name of Spectronaut global protein output, which is as in the same format as

input parameter.

annotation\_protein

name of annotation file for global protein data, in the same format as above.

use\_unmod\_peptides

If protein\_input is not provided, unmodified peptides can be extracted from

input to be used in place of a global profiling run. Default is FALSE.

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

peak area normalized by Spectronaut. Default is NULL

mod\_id Character that indicates the modification of interest. Default is \\(Phospho\\).

Note \\ must be included before special characters.

fasta\_protein\_name

Name of fasta column that matches with protein name in evidence file. Default

is uniprot\_iso.

remove\_other\_mods

Remove peptides which include modfications other than the one listed in mod\_id. Default is TRUE. For example, in an experiment targeting Phosphorylation, setting this parameter to TRUE would remove peptides like (Acetyl (Protein Nterm))AAAAPDSRVS(Phospho (STY))EEENLK. Set this parameter to FALSE

to keep peptides with extraneous modifications.

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.

 ${\tt qvalue\_cutoff} \quad {\tt Cutoff} \ \ {\tt for} \ {\tt EG.Qvalue}. \ {\tt Default} \ {\tt is} \ 0.01.$ 

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

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

 $\mbox{max}(\mbox{default})$  or  $\mbox{sum}$  - 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.

#### Value

a list of two data.tables named 'PTM' and 'PROTEIN' in the format required by MSstatsPTM.

## **Examples**

spectronaut\_annotation

Example annotation file for a label-free Spectronaut experiment.

#### **Description**

Must be manually created by the user and input into the SpectronauttoMSstatsPTMFormat converter. Requires the correct columns and maps the experimental desing into the MSstats format. Specify unique bioreplicates for group comparison designs, and the same bioreplicate for repeated measure designs. The columns and descriptions are below.

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#### Usage

spectronaut\_annotation

#### **Format**

A data.table with 5 columns.

#### **Details**

- Run: Run name that matches exactly with Spectronaut run. Used to join evidence and metadata in annotation file.
- Condition: Name of condition that was used for each run.
- BioReplicate: Name of biological replicate. Repeating the same name here will tell MSstat-sPTM that the experiment is a repeated measure design.
- Raw.file: Run name that matches exactly with Spectronaut run. Used to join evidence and metadata in annotation file.

## **Examples**

head(spectronaut\_annotation)

 $spectronaut\_input$ 

Example Spectronaut evidence file from the output of a label free experiment

## **Description**

Experiment was performed by the Olsen lab and published on Nat. Commun. (citation below).

## Usage

spectronaut\_input

#### **Format**

a data.table with 23 columns and 2683 rows, the output of Spectronaut

#### **Details**

Bekker-Jensen, D.B., Bernhardt, O.M., Hogrebe, A. et al. Rapid and site-specific deep phosphoproteome profiling by data-independent acquisition without the need for spectral libraries. Nat Commun 11, 787 (2020). https://doi.org/10.1038/s41467-020-14609-1

The experiment was processed using Spectronaut by the computational proteomics team at Pfizer (Liang Xue and Pierre Jean).

The experiment did not contain a global profiling run, but we show an example of extracting the unmodified peptides and using them in place of the profiling run.

56 summary.data

#### **Examples**

head(spectronaut\_input)

summary.data

Example of output from dataSummarizationPTM function for non-TMT data

## Description

It is made from raw.input. It is the output of dataSummarizationPTM function from MSstatsPTM. It should include a list with two names PTM and PROTEIN. Each of these list values is also a list with two names ProteinLevelData and FeatureLevelData, which correspond to two data.tables. The columns in these two data.tables are listed below. The variables are as follows:

- FeatureLevelData:
  - PROTEIN: Protein ID with modification site mapped in. Ex. Protein\_1002\_S836
  - PEPTIDE: Full peptide with charge
  - TRANSITION: Charge
  - FEATURE: Combination of Protien, Peptide, and Transition Columns
  - LABEL:
  - GROUP: Condition (ex. Healthy, Cancer, Time0)
  - RUN: Unique ID for technical replicate of one TMT mixture.
  - SUBJECT: Unique ID for biological subject.
  - FRACTION: Unique Fraction ID
  - originalRUN: Run name
  - censored:
  - INTENSITY: Unique ID for TMT mixture.
  - ABUNDANCE: Unique ID for TMT mixture.
  - newABUNDANCE : Unique ID for TMT mixture.
  - predicted: Unique ID for TMT mixture.
- ProteinLevelData:
  - RUN: MS run ID
  - Protein: Protein ID with modification site mapped in. Ex. Protein 1002 S836
  - LogIntensities: Protein-level summarized abundance
  - original RUN: Labeling information (126, ... 131)
  - GROUP: Condition (ex. Healthy, Cancer, Time0)
  - SUBJECT : Unique ID for biological subject.
  - TotalGroupMeasurements : Unique ID for technical replicate of one TMT mixture.
  - NumMeasuredFeature : Unique ID for TMT mixture.
  - MissingPercentage: Unique ID for TMT mixture.
  - more50missing: Unique ID for TMT mixture.
  - NumImputedFeature : Unique ID for TMT mixture.

summary.data.tmt 57

## Usage

summary.data

#### **Format**

A list of two lists with four data.tables.

#### **Examples**

head(summary.data)

summary.data.tmt

Example of output from dataSummarizationPTM\_TMT function for TMT data

## **Description**

It is made from raw.input.tmt. It is the output of dataSummarizationPTM\_TMT function from MSstatsPTM. It should include a list with two names PTM and PROTEIN. Each of these list values is also a list with two names ProteinLevelData and FeatureLevelData, which correspond to two data.tables.The columns in these two data.tables are listed below. The variables are as follows:

- FeatureLevelData:
  - ProteinName: MS run ID
  - PSM: Protein ID with modification site mapped in. Ex. Protein\_1002\_S836
  - censored: Protein-level summarized abundance
  - predicted : Labeling information (126, ... 131)
  - log2Intensity : Condition (ex. Healthy, Cancer, Time0)
  - Run: Unique ID for biological subject.
  - Channel: Unique ID for technical replicate of one TMT mixture.
  - BioReplicate: Unique ID for TMT mixture.
  - Condition : Unique ID for TMT mixture.
  - Mixture: Unique ID for TMT mixture.
  - TechRepMixture : Unique ID for TMT mixture.
  - PeptideSequence : Unique ID for TMT mixture.
  - Charge: Unique ID for TMT mixture.
- ProteinLevelData:
  - Mixture: MS run ID
  - TechRepMixture: Protein ID with modification site mapped in. Ex. Protein\_1002\_S836
  - Run: Protein-level summarized abundance
  - Channel: Labeling information (126, ... 131)
  - Protein: Condition (ex. Healthy, Cancer, Time0)
  - Abundance: Unique ID for biological subject.
  - BioReplicate: Unique ID for technical replicate of one TMT mixture.
  - Condition: Unique ID for TMT mixture.

58 tidyFasta

# Usage

```
summary.data.tmt
```

#### **Format**

A list of two lists with four data.tables.

# **Examples**

```
head(summary.data.tmt)
```

tidyFasta

Read and tidy a FASTA file

# Description

tidyFasta reads and tidys FASTA file. Use this function as the first step in identifying modification sites.

## Usage

```
tidyFasta(path)
```

## **Arguments**

path

A string of path to a FASTA file.

# Value

A data.table with columns named header, sequence, uniprot\_ac, uniprot\_iso, entry\_name.

# **Examples**

```
tidyFasta(system.file("extdata", "013297.fasta", package="MSstatsPTM"))
```

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