Package 'MSstatsLiP'

July 10, 2025

Type Package

Title LiP Significance Analysis in shotgun mass spectrometry-based proteomic experiments

Version 1.14.1

Date 2024-3-19

Description Tools for LiP peptide and protein significance analysis. Provides functions for summarization, estimation of LiP peptide abundance, and detection of changes across conditions. Utilizes functionality across the MSstats family of packages.

License Artistic-2.0

Depends R (>= 4.1)

Imports dplyr, gridExtra, stringr, ggplot2, grDevices, MSstats, MSstatsConvert, data.table, Biostrings, MSstatsPTM, Rcpp, checkmate, factoextra, ggpubr, purrr, tibble, tidyr, tidyverse, scales, stats

Suggests BiocStyle, knitr, rmarkdown, covr, tinytest, gghighlight

LinkingTo Rcpp

VignetteBuilder knitr

biocViews ImmunoOncology, MassSpectrometry, Proteomics, Software, DifferentialExpression, OneChannel, TwoChannel, Normalization, QualityControl

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

Encoding UTF-8

LazyData TRUE

Roxygen list(markdown = TRUE)

RoxygenNote 7.2.3

git_url https://git.bioconductor.org/packages/MSstatsLiP

git_branch RELEASE_3_21

git_last_commit 0915f78

git_last_commit_date 2025-05-27

Repository Bioconductor 3.21

Date/Publication 2025-07-09

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

Maintainer Devon Kohler <kohler.d@northeastern.edu>

Contents

annotSite
calculateProteolyticResistance
calculateTrypticity
correlationPlotLiP
dataProcessPlotsLiP
dataSummarizationLiP
DIANNtoMSstatsLiPFormat
groupComparisonLiP
groupComparisonPlotsLiP
LiPRawData
locateMod
locatePTM
MSstatsLiP
MSstatsLiP_data 20
MSstatsLiP_model
MSstatsLiP_Summarized
PCAPlotLiP
raw_lip 25
raw_prot
ResistanceBarcodePlotLiP
SkylineTest
SkylinetoMSstatsLiPFormat 30
SpectronauttoMSstatsLiPFormat
StructuralBarcodePlotLiP
tidyFasta
TrPRawData
trypticHistogramLiP

Index

annotSite

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)

calculateProteolyticResistance

Calcutates proteolytic resistance for provided data. Requires input from dataSummarizationLiP function. Can optionally calculate differential analysis using proteolytic resistance. In order for this function to work, Conditions and run numbers must match between the LiP and TrP data.

Description

Calcutates proteolytic resistance for provided data. Requires input from dataSummarizationLiP function. Can optionally calculate differential analysis using proteolytic resistance. In order for this function to work, Conditions and run numbers must match between the LiP and TrP data.

Usage

```
calculateProteolyticResistance(
  LiP_data,
  fasta_file,
  differential_analysis = FALSE,
  contrast.matrix = "pairwise"
)
```

Arguments

LiP_data	name of variable containing LiP data. Must be output of dataSummarizationLiP function.	
fasta_file	name of variable containing FASTA data. If FASTA file has not been processed please run the tidyFasta() function on it before inputting into this function. Protein names in file must match those in LiP_data.	
differential_analysis		
	logical indicating whether to run differential analysis. Default is FALSE. Con- ditions and run numbers must match between the LiP and TrP data.	
contrast.matrix		
	either a string of "pairwise" or a matrix including what comparisons to make in the differential analysis. Only required if differential_analysis=TRUE. Default is "pairwise".	

Value

a data.frame including either the summarized Proteolytic data or differential analysis depending on parameter selection.

Examples

```
fasta <- tidyFasta(system.file("extdata", "ExampleFastaFile.fasta", package="MSstatsLiP"))
#calculateProteolyticResistance(MSstatsLiP_data, fasta)</pre>
```

calculateTrypticity Calculates level of trypticity for a list of LiP Peptides.

Description

Takes as as input LiP data and a fasta file. These can be the outputs of MSstatsLiP functions.

Usage

```
calculateTrypticity(LiP_data, fasta_file)
```

4

Arguments

LiP_data	name of variable containing LiP data. Must contain at least two columns named 'PeptideSequence' and 'ProteinName'. The values in these column must match with what is in the corresponding FASTA file.
fasta_file	name of variable containing FASTA data. If FASTA file has not been processed please run the tidyFasta() function on it before inputting into this function.

Value

a data.frame including protein, peptide, and trypticity metrics.

Examples

```
fasta <- tidyFasta(system.file("extdata", "ExampleFastaFile.fasta", package="MSstatsLiP"))
calculateTrypticity(MSstatsLiP_data$LiP, fasta)</pre>
```

correlationPlotLiP Plot run correlation for provided LiP and TrP experiment.

Description

Plot run correlation for provided LiP and TrP experiment.

Usage

```
correlationPlotLiP(
  data,
  method = "pearson",
  value_columns = "INTENSITY",
  x.axis.size = 10,
  y.axis.size = 10,
  legend.size = 10,
  width = 10,
  height = 10,
  address = ""
)
```

data	output of MSstatsLiP converter function. Must include at least ProteinName, Run, and Intensity columns
method	one of "pearson", "kendall", "spearman". Default is pearson.
value_columns	one of "INTENSITY" or "ABUNDANCE". INTENSITY is the raw data, whereas ABUNDANCE is the log transformed INTENSITY column. INTENSITY is default.

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.
legend.size	size of legend for color at the bottom of volcano plot. Default is 10.
width	width of the saved file. Default is 10.
height	height of the saved file. Default is 10.
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

Use output of dataSummarizationLiP function correlationPlotLiP(MSstatsLiP_Summarized, address = FALSE)

dataProcessPlotsLiP Visualization for explanatory data analysis

Description

To illustrate the quantitative data and quality control of MS runs, dataProcessPlotsLiP takes the quantitative data from MSstatsLiP converter functions 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

```
dataProcessPlotsLiP(
   data,
   type = "PROFILEPLOT",
   ylimUp = FALSE,
   ylimDown = FALSE,
   x.axis.size = 10,
   y.axis.size = 10,
   text.size = 4,
   text.angle = 90,
   legend.size = 7,
```

dataProcessPlotsLiP

```
dot.size.profile = 2,
ncol.guide = 5,
width = 10,
height = 12,
lip.title = "All Peptides",
protein.title = "All Proteins",
which.Peptide = "all",
which.Protein = NULL,
originalPlot = TRUE,
summaryPlot = TRUE,
address = ""
```

data	name of the list with LiP and (optionally) Protein data, which can be the output of the MSstatsLiP. dataSummarizationLiP function.	
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.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.	
lip.title	title of all LiP QC plot	
protein.title	title of all Protein QC plot	
which.Peptide	LiP peptide list to draw plots. List can be names of LiP peptides or order num- bers of LiPs. Default is "all", which generates all plots for each protein. For QC plot, "allonly" will generate one QC plot with all proteins.	
which.Protein	String of protein's to plot if the user would like to plot all Peptides associated with a given Protein. Default is NULL. Please do not include "all" or "allonly" here.	

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

```
# Use the output of the MSstatsLiP_Summarized function
# Profile Plot
dataProcessPlotsLiP(MSstatsLiP_Summarized, type = "ProfilePlot")
# QCPlot Plot
dataProcessPlotsLiP(MSstatsLiP_Summarized, type = "QCPlot")
```

dataSummarizationLiP	Summarizes LiP and TrP datasets seperately using methods from
	MSstats.

Description

Utilizes functionality from MSstats and MSstatsPTM to clean, summarize, and normalize LiP peptide and TrP global protein data. Imputes missing values, protein and LiP peptide level summarization from peptide level quantification. Applies global median normalization on peptide level data and normalizes between runs. Returns list of two summarized datasets.

Usage

```
dataSummarizationLiP(
   data,
   logTrans = 2,
   normalization = "equalizeMedians",
   normalization.LiP = "equalizeMedians",
   nameStandards = NULL,
   nameStandards.LiP = NULL,
   featureSubset = "all",
   featureSubset.LiP = "all",
```

```
remove_uninformative_feature_outlier = FALSE,
remove_uninformative_feature_outlier.LiP = FALSE,
min_feature_count = 2,
min_feature_count.LiP = 1,
n_top_feature = 3,
n_top_feature.LiP = 3,
summaryMethod = "TMP",
equalFeatureVar = TRUE,
censoredInt = "NA",
MBimpute = TRUE,
MBimpute.LiP = FALSE,
remove50missing = FALSE,
fix_missing = NULL,
maxQuantileforCensored = 0.999,
use_log_file = FALSE,
append = FALSE,
verbose = TRUE,
log_file_path = NULL,
base = "MSstatsLiP_log_"
```

```
)
```

data	name of the list with LiP and TrP data.tables, which can be the output of the MSstatsPTM converter functions $% \mathcal{M} = \mathcal{M} = \mathcal{M} + \mathcal{M} + \mathcal{M}$	
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. 'equalizeMedi- ans'(default) represents constant normalization (equalizing the medians) based on reference signals is performed. 'quantile' represents quantile normalization based on reference signals is performed. 'globalStandards' represents normal- ization with global standards proteins. FALSE represents no normalization is performed	
normalization.LiP		
	normalization for LiP 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 normaliza- tion with global standard peptides.	
nameStandards.LiP		
	Same as above for LiP dataset.	
featureSubset	For protein dataset only. "all"(default) uses all features that the data set has. "top3" uses top 3 features which have highest average of log2(intensity) across runs. "topN" uses top N features which has highest average of log2(intensity) across runs. It needs the input for n_top_feature option. "highQuality" flags uninformative feature and outliers	
featureSubset.LiP		
	For LiP dataset only. Options same as above.	

remove_uninform	native_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 col- umn, feature_quality="Uninformative" which are features with bad quality, 2) outliers that are flagged in the column, is_outlier=TRUE, for run-level summa- rization. FALSE (default) uses all features and intensities for run-level summa- rization.
remove_uninform	native_feature_outlier.LiP
min_feature_cou	For LiP dataset only. Options same as above.
	optional. Only required if featureSubset = "highQuality". Defines a minimum number of informative features a protein needs to be considered in the feature selection algorithm.
<pre>min_feature_cou</pre>	
	For LiP dataset only. Options the same as above.
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.L	
	For LiP 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 inten- sities 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 intensities 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.LiP	For LiP dataset only. Options same as above. Default is FALSE.
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
maxQuantilefor(-
	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 LiP and TrP results. These results contain the reformatted input to the summarization function, as well as run-level summarization results.

Examples

```
# Use output of converter
head(MSstatsLiP_data[["LiP"]])
head(MSstatsLiP_data[["TrP"]])
```

Run summarization
MSstatsLiP_model <- dataSummarizationLiP(MSstatsLiP_data)</pre>

DIANNtoMSstatsLiPFormat

Converts raw LiP MS data from DIA-NN into the format needed for MSstatsLiP.

Description

Takes as as input both raw LiP and Trp outputs from DIA-NN

Usage

```
DIANNtoMSstatsLiPFormat(
  lip_data,
  trp_data = NULL,
  annotation = NULL,
  global_qvalue_cutoff = 0.01,
  qvalue_cutoff = 0.01,
  pg_qvalue_cutoff = 0.01,
  useUniquePeptide = TRUE,
  removeFewMeasurements = TRUE,
  removeOxidationMpeptides = FALSE,
  removeProtein_with1Feature = FALSE,
  use_log_file = FALSE,
  append = FALSE,
  verbose = TRUE,
  log_file_path = NULL
)
```

Arguments

lip_data	name of LiP Skyline output, which is long-format.
trp_data	name of TrP Skyline output, which is long-format.
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.
global_qvalue_c	cutoff
	The global qvalue cutoff. Default is 0.01.
qvalue_cutoff	Cutoff for DetectionQValue. Default is 0.01.
pg_qvalue_cutof	f
	local qvalue cutoff for protein groups Run should be the same as filename. De- fault is .01.
useUniquePeptic	le
	TRUE (default) removes peptides that are assigned for more than one proteins. We assume to use unique peptide for each protein.
removeFewMeasur	rements
	TRUE (default) will remove the features that have 1 or 2 measurements across runs.
removeOxidation	Mpeptides
	TRUE will remove the peptides including 'oxidation (M)' in modification. FALSE is default.
<pre>removeProtein_w</pre>	vith1Feature
	TRUE will remove the proteins which have only 1 feature, which is the combi- nation of peptide, precursor charge, fragment and charge. FALSE is default.
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 saved to a 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.frames in MSstatsLiP format

Examples

```
## Output will be in format
head(MSstatsLiP_data[["LiP"]])
head(MSstatsLiP_data[["TrP"]])
```

groupComparisonLiP

Model LiP and TrP data and make adjustments if needed Returns list of three modeled datasets

Description

Takes summarized LiP peptide and TrP protein data from dataSummarizationLiP If global protein data is unavailable, LiP data only can be passed into the function. Including protein data allows for adjusting LiP Fold Change by the change in global protein abundance..

Usage

```
groupComparisonLiP(
   data,
   contrast.matrix = "pairwise",
   fasta.path = NULL,
   log_base = 2,
   use_log_file = FALSE,
   append = FALSE,
   verbose = TRUE,
   log_file_path = NULL,
   base = "MSstatsLiP_log_"
)
```

data	list of summarized datasets. Can be output of MSstatsLiP summarization func- tion dataSummarizationLiP. Must include dataset named "LiP" as minimum.
contrast.matrix	
	comparison between conditions of interests. Default models full pairwise comparison between all conditions
fasta.path	a file path to a fasta file that includes the proteins listed in the data. Default is NULL. Include this parameter to determine trypticity of peptides in LiP models.
log_base	base of the logarithm used in dataProcess.
use_log_file	logical. If TRUE, information about data processing will be saved to a file.
append	logical. If TRUE, information about data processing will be added to an existing log file.
verbose	logical. If TRUE, information about data processing 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 LiP, PROTEIN, and ADJUSTED LiP data.tables with their corresponding model results.

Examples

groupComparisonPlotsLiP

Visualization for model-based analysis and summarization

Description

To analyze the results of modeling changes in abundance of LiP peptides and overall protein, group-ComparisonPlotsLiP takes as input the results of the groupComparisonLiP function. It asses the results of three models: unadjusted LiP, adjusted LiP, 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

```
groupComparisonPlotsLiP(
   data = data,
   type = type,
   sig = 0.05,
   FCcutoff = 1,
   logBase.pvalue = 10,
   ylimUp = FALSE,
   ylimDown = FALSE,
   xlimUp = FALSE,
   x.axis.size = 10,
   y.axis.size = 10,
   dot.size = 3,
```

```
text.size = 4,
text.angle = 0,
legend.size = 13,
ProteinName = TRUE,
colorkey = TRUE,
numProtein = 100,
width = 10,
height = 10,
which.Comparison = "all",
which.Peptide = "all",
which.Protein = NULL,
address = ""
```

data	name of the list with models, which can be the output of the MSstatsLiP groupComparisonLiP function
type	choice of visualization, one of VolcanoPlot or Heatmap
sig	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	or 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.

	for volcano plot only, whether display protein/peptide 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.Compariso	n
	list of comparisons to draw plots. List can be labels of comparisons or order numbers of comparisons from levels(data\$Label), such as levels(testResultMultiComparisons\$Comparison Default is "all", which generates all plots for each protein.
	Peptide list to draw comparison plots. List can be names of Peptides or order numbers of Peptides from levels. Default is "all", which generates all compari- son plots for each protein.
	Protein list to draw comparison plots. Will draw all peptide plots for listed Proteins. List must be names of Proteins. Default is "all", which generates all comparison plots for each protein.
	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

Use output of the groupComparisonLiP function
Volcano Plot
groupComparisonPlotsLiP(MSstatsLiP_model, type = "VOLCANOPLOT")
Heatmap Plot
groupComparisonPlotsLiP(MSstatsLiP_model, type = "HEATMAP")

LiPRawData

LiPRawData

Description

Example of input LiP dataset.

LiPRawData

Usage

LiPRawData

Format

A data.table consisting of 546 rows and 29 columns. Raw LiP data for use in testing and examples.

Details

Input to MSstatsLiP converter SpectronauttoMSstatsLiPFormat. Contains the following columns:

- R.Condition : Label of conditions (EG Disease/Control)
- R.FileName : Name of spectral processing run
- R.Replicate : Name of biological replicate
- PG.ProteinAccessions : Protein name
- PG.ProteinGroups : Protein name, can be multiple
- PG.Quantity : Protein Quantity
- PEP.GroupingKey : Peptide grouping
- PEP.StrippedSequence : Peptide sequence
- PEP.Quantity : Peptide quantity
- EG.iRTPredicted : Predicted value
- EG.Library : Name of library
- · EG.ModifiedSequence : Peptide sequence including any post-translational modifications
- EG.PrecursorId : Peptide sequence wiht modifications including charge
- EG.Qvalue : Qvalue
- FG.Charge : Identified Ion charge
- FG.Id : Peptide sequence with charge
- FG.PrecMz : Prec Mz reading
- FG.Quantity : Initial quantity reading
- F.Charge : F.Charge
- F.FrgIon : Fragment ion
- F.FrgLossType : Label for loss type
- F.FrgMz : Mz reading
- F.FrgNum : numeric Frg
- F.FrgType : character label for Frg
- F.ExcludedFromQuantification : True/False boolean for if to exclude
- F.NormalizedPeakArea : Normalized peak intensity
- F.NormalizedPeakHeight : Normalized peak height
- F.PeakArea : Unnormalized peak area
- F.PeakHeight : Unnormalized peak height

Examples

head(LiPRawData)

locateMod

Locate modified sites with a peptide

Description

locateMod locates modified sites with a peptide.

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)

18

MSstatsLiP

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 tibble 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. Default is FALSE.

Value

A data frame with three columns: uniprot_iso, peptide, site.

Examples

```
fasta <- tidyFasta(system.file("extdata", "013297.fasta", package="MSstatsLiP"))
locatePTM("DRVSYIHNDSC*TR", "013297", fasta, "C", "\\*")</pre>
```

MSstatsLiP	
------------	--

MSstatsLiP: A package for identifying and analyzing changes in protein structures caused by compound binding in cellur lysates.

Description

A set of tools for detecting differentially abundant LiP peptides in shotgun mass spectrometry-based proteomic experiments. The package includes tools to convert raw data from different spectral processing tools, summarize feature intensities, and fit a linear mixed effects model. If overall protein abundance changes are included, the package will also adjust the LiP peptide fold change for changes in overall protein abundace. Additionally the package includes functionality to plot a variety of data visualizations.

functions

- SpectronauttoMSstatsLiPFormat : Generates MSstatsLiP required input format for Spectronaut outputs.
- trypticHistogramLiP : Histogram of Half vs Fully tryptic peptides. Calculates proteotypicity, and then uses calculations in histogram.
- correlationPlotLiP : Plot run correlation for provided LiP and TrP experiment.
- dataSummarizationLiP : Summarizes PSM level quantification to peptide (LiP) and protein level quantification.
- dataProcessPlotsLiP: Visualization for explanatory data analysis. Specifically gives ability to plot Profile and Quality Control plots.

- PCAPlotLiP : Visualize PCA analysis for LiP and TrP datasets. Specifically gives ability to plot explanined variance per component, Protein/Peptide PCA, and Condition PCA.
- groupComparisonLiP : Tests for significant changes in LiP and protein abundance across conditions. Adjusts LiP fold change for changes in protein abundance.
- groupComparisonPlotsLiP : Visualization for model-based analysis and summarization.
- PCAPlotLiP : Runs PCA on the summarized data. Can visualize the PCA analysis in three different plots.
- StructuralBarcodePlotLiP : Shows protein coverage of LiP modified peptides. Shows significant, insignificant, and missing coverage.

MSstatsLiP_data MSstatsLiP_data

Description

Example output of MSstatsLiP converter functions.

Usage

MSstatsLiP_data

Format

A data.table consisting of 546 rows and 29 columns. Raw TrP data for use in testing and examples.

Details

Example output of MSstatsLiP converter functions. (Eg. SpectronauttoMSstatsLiPFormat). A list containing two data.tables named LiP and TrP corresponding to the processed LiP and TrP data now in MSstatsLiP format. The data.tables contain the following columns:

- · ProteinName : Character column of protein names
- PeptideSequence : Character column of peptide sequence name
- PrecursorCharge : Numeric charge feature
- FragmentIon : Character fragment ion feature
- ProductCharge : Numeric charge of product
- IsotopeLabelType : Character label type
- Condition : Character label for condition (Eg. Disease/Control)
- BioReplicate : Name of biological replicate
- Run : Name of run
- · Fraction : Fraction number if fractionation is present
- Intensity : Unnormalized feature intensity
- FULL_PEPTIDE(LiP data only) : Combined protein name and peptide sequence. Used for LiP data only because LiP is summarized to peptide level (not protein)

MSstatsLiP_model

Examples

```
head(MSstatsLiP_data$LiP)
head(MSstatsLiP_data$TrP)
```

MSstatsLiP_model MSstatsLiP_model

Description

Example output of groupComparisonLiP converter functions.

Usage

MSstatsLiP_model

Format

A data.table consisting of 546 rows and 29 columns. Raw TrP data for use in testing and examples.

Details

Example output of MSstatsLiP groupComparisonLiP function. A list containing three data.tables corresponding to unadjusted LiP, TrP, and adjusted LiP models. The data.tables contain the following columns:

- ProteinName : Character column of protein names
- PeptideSequence : Character column of peptide sequence name
- Label : Condition comparison (Eg. Disease vs Control)
- log2FC : Fold Change output results of model
- SE : Standard error output of model
- · Tvalue : Tvalue output of model
- DF : Degrees of Freedom output of model
- pvalue : Pvalue result of model (unadjusted)
- · adj.pvalue : Adjusted Pvalue, generally BH adjustement is used
- issue : Issue in model if any is reported
- MissingPercentage : Percent of missing values in specific model
- ImputationPercentage : Percent of values that needed to be imputed
- fully_TRI: Boolean indicating if Peptide is fully tryptic
- NSEMI_TRI: Boolean indicating if Peptide is NSEMI tryptic
- CSEMI_TRI: Boolean indicating if Peptide is CSEMI tryptic
- CTERMINUS: Boolean indicating if Peptide is CTERMINUS tryptic

- NTERMINUS: Boolean indicating if Peptide is NTERMINUS tryptic
- StartPos: Start position of peptide sequence
- EndPos: End position of peptide sequence
- FULL_PEPTIDE(LiP data only) : Combined protein name and peptide sequence. Used for LiP data only because LiP is summarized to peptide level (not protein)

Examples

```
head(MSstatsLiP_model$LiP.Model)
head(MSstatsLiP_model$TrP.Model)
head(MSstatsLiP_model$Adjusted.LiP.Model)
```

MSstatsLiP_Summarized MSstatsLiP_Summarized

Description

Example output of MSstatsLiP summarization function dataSummarizationLiP.

Usage

MSstatsLiP_Summarized

Format

A list containing two lists of summarization information for LiP and TrP data.

Details

Example output of MSstatsLiP summarization function dataSummarizationLiP. A list containing two lists named LiP and TrP containing summarization information for LiP and TrP data. Each of LiP and TrP contain data named: FeatureLevelData, ProteinLevelData, SummaryMethod, ModelQC, PredictBySurvival. The two main data.tables (FeatureLevelData and ProteinLevelData are shown below):

- FeatureLevelData :
 - PROTEIN : Protein ID with modification site mapped in. Ex. Protein_1002_S836
 - FULL_PEPTIDE (LiP Only) : Combined name of protein and peptide sequence
 - 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 : Original intensity value
- ABUNDANCE : Log adjusted intensity value
- newABUNDANCE : Normalized abundance column
- ProteinLevelData :
 - RUN : MS run ID
 - FULL_PEPTIDE (LiP Only) : Combined name of protein and peptide sequence
 - Protein : Protein ID with modification site mapped in. Ex. Protein_1002_S836
 - LogIntensities: Protein-level summarized abundance
 - originalRUN : 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.

Examples

head(MSstatsLiP_Summarized\$LiP\$FeatureLevelData) head(MSstatsLiP_Summarized\$LiP\$ProteinLevelData)

head(MSstatsLiP_Summarized\$TrP\$FeatureLevelData) head(MSstatsLiP_Summarized\$TrP\$ProteinLevelData)

PCAPlotLiP

Visualize PCA analysis for LiP and TrP datasets.

Description

Takes as input LiP and TrP data from summarization function dataSummarizationLiP. Runs PCA on the summarized data. Can visualize the PCA analysis in three different plots: (1) BarPlot (specify "bar.plot=TRUE" in option bar.plot), to plot a bar plot showing the explained variance per PCA component (2) Peptide/Protein PCA (specify "protein.pca = TRUE" in option protein.pca), to create a dot plot with PCA component 1 and 2 on the axis, for different peptides and proteins. (3) Comparison PCA (specify "comparison.pca = TRUE" in option comparison.pca), to create a arrow plot with PCA component 1 and 2 on the axis, for different comparison.pca) to create a arrow plot with PCA component 1 and 2 on the axis, for different comparison.pca) to create a arrow plot with PCA component 1 and 2 on the axis, for different comparison.pca arrow plot with PCA component 1 and 2 on the axis, for different comparison.pca arrow plot with PCA component 1 and 2 on the axis, for different comparison.pca arrow plot with PCA component 1 and 2 on the axis, for different comparison.pca arrow plot with PCA component 1 and 2 on the axis, for different comparison.pca arrow plot with PCA component 1 and 2 on the axis, for different comparison.pca arrow plot with PCA component 1 and 2 on the axis, for different comparison.pca arrow plot with PCA component 1 and 2 on the axis, for different comparison.pca arrow plot with PCA component 1 and 2 on the axis, for different comparison.pca arrow plot with PCA component 1 and 2 on the axis, for different comparison place arrow plot arrow plot place arrow plot place arrow plot place arrow place

Usage

```
PCAPlotLiP(
   data,
   center.pca = TRUE,
   scale.pca = TRUE,
   n.components = 10,
   bar.plot = TRUE,
   protein.pca = TRUE,
   comparison.pca = FALSE,
   which.pep = "all",
   which.comparison = "all",
   width = 10,
   height = 10,
   address = ""
)
```

Arguments

data	data name of the list with LiP and (optionally) Protein data, which can be the output of the MSstatsLiP. dataSummarizationLiP function.
center.pca	a logical value indicating whether the variables should be shifted to be zero centered. Alternately, a vector of length equal the number of columns of x can be supplied. The value is passed to scale
scale.pca	a logical value indicating whether the variables should be scaled to have unit variance before the analysis takes place. The default is FALSE for consistency with S, but in general scaling is advisable. Alternatively, a vector of length equal the number of columns of x can be supplied. The value is passed to scale.
n.components	an integer of PCA components to be returned. Default is 10.
bar.plot	a logical value indicating if to visualize PCA bar plot
protein.pca	a logical value indicating if to visualize PCA peptide plot
comparison.pca	a logical value indicating if to visualize PCA comparison plot
which.pep	a list of peptides to be visualized. Default is "all". If too many peptides are plotted the names can overlap.
which.compariso	n
	a list of comparisons to be visualized. Default is "all".
width	width of the saved file. Default is 10.
height	height of the saved file. Default is 10.
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

24

raw_lip

Value

plot or pdf

Examples

```
# Use output of dataSummarizationLiP function
# BarPlot
PCAPlotLiP(MSstatsLiP_Summarized, bar.plot = TRUE, protein.pca = FALSE)
# Protein/Peptide PCA Plot
PCAPlotLiP(MSstatsLiP_Summarized, bar.plot = FALSE, protein.pca = TRUE)
# Condition PCA Plot
PCAPlotLiP(MSstatsLiP_Summarized, bar.plot = FALSE, protein.pca = FALSE, comparison.pca = TRUE)
```

raw_lip

raw_lip

Description

A different example of input LiP dataset.

Usage

raw_lip

Format

A data.table consisting of 6,944 rows and 29 columns. Raw LiP data for use in testing and examples.

Details

Input to MSstatsLiP converter SpectronauttoMSstatsLiPFormat. Contains the following columns:

- R.Condition : Label of conditions (EG Disease/Control)
- R.FileName : Name of spectral processing run
- R.Replicate : Name of biological replicate
- PG.ProteinAccessions : Protein name
- PG.ProteinGroups : Protein name, can be multiple
- PG.Quantity : Protein Quantity
- PEP.GroupingKey : Peptide grouping
- PEP.StrippedSequence : Peptide sequence
- PEP.Quantity : Peptide quantity

- EG.iRTPredicted : Predicted value
- EG.Library : Name of library
- EG.ModifiedSequence : Peptide sequence including any post-translational modifications
- EG.PrecursorId : Peptide sequence wiht modifications including charge
- EG.Qvalue : Qvalue
- FG.Charge : Identified Ion charge
- FG.Id : Peptide sequence with charge
- FG.PrecMz : Prec Mz reading
- FG.Quantity : Initial quantity reading
- F.Charge : F.Charge
- F.FrgIon : Fragment ion
- F.FrgLossType : Label for loss type
- F.FrgMz : Mz reading
- F.FrgNum : numeric Frg
- F.FrgType : character label for Frg
- F.ExcludedFromQuantification : True/False boolean for if to exclude
- F.NormalizedPeakArea : Normalized peak intensity
- F.NormalizedPeakHeight : Normalized peak height
- F.PeakArea : Unnormalized peak area
- F.PeakHeight : Unnormalized peak height

Examples

head(raw_lip)

raw_prot

raw_prot

Description

Example of input TrP dataset.

Usage

raw_prot

Format

A data.table consisting of 9,120 rows and 29 columns. Raw TrP data for use in testing and examples.

raw_prot

Details

Input to MSstatsLiP converter SpectronauttoMSstatsLiPFormat. Contains the following columns:

- R.Condition : Label of conditions (EG Disease/Control)
- R.FileName : Name of spectral processing run
- R.Replicate : Name of biological replicate
- PG.ProteinAccessions : Protein name
- PG.ProteinGroups : Protein name, can be multiple
- PG.Quantity : Protein Quantity
- PEP.GroupingKey : Peptide grouping
- PEP.StrippedSequence : Peptide sequence
- PEP.Quantity : Peptide quantity
- EG.iRTPredicted : Predicted value
- EG.Library : Name of library
- EG.ModifiedSequence : Peptide sequence including any post-translational modifications
- EG.PrecursorId : Peptide sequence wiht modifications including charge
- EG.Qvalue : Qvalue
- FG.Charge : Identified Ion charge
- FG.Id : Peptide sequence with charge
- FG.PrecMz : Prec Mz reading
- FG.Quantity : Initial quantity reading
- F.Charge : F.Charge
- F.FrgIon : Fragment ion
- F.FrgLossType : Label for loss type
- F.FrgMz : Mz reading
- F.FrgNum : numeric Frg
- F.FrgType : character label for Frg
- F.ExcludedFromQuantification : True/False boolean for if to exclude
- F.NormalizedPeakArea : Normalized peak intensity
- F.NormalizedPeakHeight : Normalized peak height
- F.PeakArea : Unnormalized peak area
- F.PeakHeight : Unnormalized peak height

Examples

head(raw_prot)

```
ResistanceBarcodePlotLiP
```

Proteolytic Resistance Barcode plot. Shows accessibility score of different fully tryptic peptides in a protein.

Description

Proteolytic Resistance Barcode plot. Shows accessibility score of different fully tryptic peptides in a protein.

Usage

```
ResistanceBarcodePlotLiP(
  data,
  fasta_file,
  which.prot = "all",
  which.condition = "all",
  differential_analysis = FALSE,
  which.comp = "all",
  adj.pvalue.cutoff = 0.05,
  FC.cutoff = 0,
  width = 12,
  height = 4,
  address = ""
)
```

data	list of data.tables containing LiP and TrP data in MSstatsLiP format. Should be output of summarization function as dataSummarizationLiP.	
fasta_file	A string of path to a FASTA file	
which.prot	a list of peptides to be visualized. Default is "all" which will plot a separate barcode plot for each protein.	
which.condition		
	a list of conditions to be visualized. Default is "all" which will plot all conditions for a single protein in the same barcode plot.	
differential_analysis		
	a boolean indicating if a barcode plot showing the differential analysis should be plotted. If this is selected you must have performed differential analysis on the proteoltic data in the calculateProteolyticResistance function. Default is FALSE.	
which.comp	a list of comparisons to be visualized, if differential analysis is passed to plot_differential variable. Default is "all" which will plot a separate barcode plot for each comparison and protein.	
adj.pvalue.cuto	ff	
	Default is .05. Alpha value for testing significance of model output.	

SkylineTest

FC.cutoff	Default is 0. Minimum absolute FC before a comparison will be considered significant.
width	width of the saved file. Default is 10.
height	height of the saved file. Default is 10.
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

```
# Specify Fasta path
fasta_path = system.file("extdata", "ExampleFastaFile.fasta", package="MSstatsLiP")
# Use model data to create Barcode Plot
#ResistanceBarcodePlotLiP(MSstatsLiP_model, fasta_path)
```

SkylineTest SkylineTest

Description

Example of input data from Skylinet.

Usage

SkylineTest

Format

A data.table consisting of 2115 rows and 13 columns. Raw data for use in testing and examples.

Details

Input to MSstatsLiP converter SkylinetoMSstatsLiPFormat Contains the following columns:

- Protein.Name : Name of Proteins identified by Skyline
- Peptide.Modified.Sequence : Peptide sequence
- Precursor.Charge : Charge of ion

- Fragment.Ion : Fragment ion
- Product.Charge : Identified Ion charge
- Isotope.Label.Type : Label Type
- Condition : Name of condition
- · BioReplicate : name of bioreplicate annotated to data
- File.Name : Name of spectral processing run
- Area : Abudance area
- Standard.Type : Type name for row
- Truncated : Boolean if row was truncated

Examples

head(SkylineTest)

SkylinetoMSstatsLiPFormat

Converts raw LiP MS data from Skyline into the format needed for MSstatsLiP.

Description

Takes as as input both raw LiP and Trp outputs from Skyline.

Usage

```
SkylinetoMSstatsLiPFormat(
 LiP.data,
 TrP.data = NULL,
  annotation = NULL,
 msstats_format = FALSE,
  removeiRT = TRUE,
  filter_with_Qvalue = TRUE,
  qvalue_cutoff = 0.01,
  useUniquePeptide = TRUE,
  removeFewMeasurements = TRUE,
  removeOxidationMpeptides = FALSE,
  removeProtein_with1Feature = FALSE,
  use_log_file = FALSE,
  append = FALSE,
  verbose = TRUE,
  log_file_path = NULL
)
```

Arguments

LiP.data	name of LiP Skyline output, which is long-format.
TrP.data	name of TrP Skyline output, which is long-format.
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.
msstats_format	logical indicating how the data was output from Skyline. FALSE (default) indi- cates that standard Skyline output was selected. TRUE should be selected if the Skyline data was output using the MSstats format option in Skyline.
removeiRT	TRUE (default) will remove the proteins or peptides which are labeld 'iRT' in 'StandardType' column. FALSE will keep them.
filter_with_Qva	alue
	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.
useUniquePeptic	de
	TRUE (default) removes peptides that are assigned for more than one proteins. We assume to use unique peptide for each protein.
removeFewMeasu	rements
	TRUE (default) will remove the features that have 1 or 2 measurements across runs.
removeOxidation	nMpeptides
	TRUE will remove the peptides including 'oxidation (M)' in modification. FALSE is default.
<pre>removeProtein_w</pre>	
	TRUE will remove the proteins which have only 1 feature, which is the combi- nation of peptide, precursor charge, fragment and charge. FALSE is default.
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 saved to a 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.frames in MSstatsLiP format

Examples

Output will be in format head(MSstatsLiP_data[["LiP"]]) head(MSstatsLiP_data[["TrP"]]) ${\tt Spectronautto} {\tt MSstatsLiPFormat}$

Converts raw LiP MS data from Spectronautt into the format needed for MSstatsLiP.

Description

Takes as as input both raw LiP and Trp outputs from Spectronautt.

Usage

```
SpectronauttoMSstatsLiPFormat(
 LiP.data,
  fasta,
 Trp.data = NULL,
  annotation = NULL,
  intensity = "PeakArea",
  filter_with_Qvalue = TRUE,
  qvalue_cutoff = 0.01,
  useUniquePeptide = TRUE,
  removeFewMeasurements = TRUE,
  removeProtein_with1Feature = FALSE,
  removeNonUniqueProteins = TRUE,
  removeModifications = TRUE,
  removeiRT = TRUE,
  summaryforMultipleRows = max,
 which.Conditions = "all",
  use_log_file = FALSE,
  append = FALSE,
  verbose = TRUE,
  log_file_path = NULL,
  base = "MSstatsLiP_log_"
)
```

LiP.data	name of LiP Spectronaut output, which is long-format.
fasta	A string of path to a FASTA file, used to match LiP peptides.
Trp.data	name of TrP Spectronaut output, which is long-format.
annotation	name of 'annotation.txt' data which includes Condition, BioReplicate, Run. If annotation is already complete in Spectronaut, use annotation=NULL (default). It will use the annotation information from input.
intensity	'PeakArea' (default) uses not normalized peak area. 'NormalizedPeakArea' uses peak area normalized by Spectronaut

filter_with_Qva	alue	
	TRUE(default) will filter out the intensities that have greater than qvalue_cutoff in EG.Qvalue column. Those intensities will be replaced with zero and will be considered as censored missing values for imputation purpose.	
qvalue_cutoff	Cutoff for EG.Qvalue. default is 0.01.	
useUniquePeptide		
	TRUE(default) removes peptides that are assigned for more than one proteins. We assume to use unique peptide for each protein.	
removeFewMeasurements		
	TRUE (default) will remove the features that have 1 or 2 measurements across runs.	
removeProtein_w		
	TRUE will remove the proteins which have only 1 feature, which is the combination of peptide, precursor charge, fragment and charge. FALSE is default.	
removeNonUnique		
	TRUE will remove proteins that were not uniquely identified. IE if the protein column contains multiple proteins separated by ";". TRUE is default	
removeModificat	ions	
	TRUE will remove peptide that contain a modification. Modification must be indicated by "[". TRUE is default	
removeiRT	TRUE will remove proteins that contain iRT. True is default	
summaryforMulti	pleRows	
	max(default) or sum - when there are multiple measurements for certain feature and certain run, use highest or sum of multiple intensities.	
which.Condition	IS	
	list of conditions to format into MSstatsLiP format. If "all" all conditions will be used. Default is "all".	
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

a list of two data.frames in MSstatsLiP format

Examples

Output datasets of Spectronaut head(LiPRawData) head(TrPRawData)

StructuralBarcodePlotLiP

Barcode plot. Shows protein coverge of LiP modified peptides.

Description

Barcode plot. Shows protein coverge of LiP modified peptides.

Usage

```
StructuralBarcodePlotLiP(
   data,
   fasta,
   model_type = "Adjusted",
   which.prot = "all",
   which.comp = "all",
   adj.pvalue.cutoff = 0.05,
   FC.cutoff = 0,
   FT.only = FALSE,
   width = 12,
   height = 4,
   address = ""
```

)

data	list of data.tables containing LiP and TrP data in MSstatsLiP format. Should be output of modeling function such as groupComparisonLiP.	
fasta	A string of path to a FASTA file	
<pre>model_type</pre>	A string of either "Adjusted" or "Unadjusted", indicating whether to plot the adjusted or unadjusted models. Default is "Adjusted".	
which.prot	a list of peptides to be visualized. Default is "all" which will plot a separate barcode plot for each protein.	
which.comp	a list of comparisons to be visualized. Default is "all" which will plot a separate barcode plot for each comparison and protein.	
adj.pvalue.cutoff		
	Default is .05. Alpha value for testing significance of model output.	

tidyFasta

FC.cutoff	Default is 0. Minimum absolute FC before a comparison will be considered significant.
FT.only	FALSE plots all FT and HT peptides, TRUE plots FT peptides only. Default is FALSE.
width	width of the saved file. Default is 10.
height	height of the saved file. Default is 10.
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

tidyFasta

Read and tidy a FASTA file

Description

reads and tidys FASTA file.

Usage

tidyFasta(path)

Arguments

path a string of path pointing towards a fasta file

Value

a tibble of formatted FASTA information

Examples

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

TrPRawData TrPRawData

Description

Example of input TrP dataset.

Usage

TrPRawData

Format

A data.table consisting of 4692 rows and 29 columns. Raw TrP data for use in testing and examples.

Details

Input to MSstatsLiP converter SpectronauttoMSstatsLiPFormat. Contains the following columns:

- R.Condition : Label of conditions (EG Disease/Control)
- R.FileName : Name of spectral processing run
- R.Replicate : Name of biological replicate
- PG.ProteinAccessions : Protein name
- PG.ProteinGroups : Protein name, can be multiple
- PG.Quantity : Protein Quantity
- PEP.GroupingKey : Peptide grouping
- PEP.StrippedSequence : Peptide sequence
- PEP.Quantity : Peptide quantity
- EG.iRTPredicted : Predicted value
- EG.Library : Name of library
- EG.ModifiedSequence : Peptide sequence including any post-translational modifications
- EG.PrecursorId : Peptide sequence wiht modifications including charge
- EG.Qvalue : Qvalue
- FG.Charge : Identified Ion charge
- FG.Id : Peptide sequence with charge
- FG.PrecMz : Prec Mz reading
- FG.Quantity : Initial quantity reading
- F.Charge : F.Charge

36

trypticHistogramLiP

- F.FrgIon : Fragment ion
- F.FrgLossType : Label for loss type
- F.FrgMz : Mz reading
- F.FrgNum : numeric Frg
- F.FrgType : character label for Frg
- F.ExcludedFromQuantification : True/False boolean for if to exclude
- F.NormalizedPeakArea : Normalized peak intensity
- F.NormalizedPeakHeight : Normalized peak height
- F.PeakArea : Unnormalized peak area
- F.PeakHeight : Unnormalized peak height

Examples

head(TrPRawData)

trypticHistogramLiP	Histogram of Half vs Fully tryptic peptides.	Calculates proteotypicity,
	and then uses calcualtions in histogram.	

Description

Histogram of Half vs Fully tryptic peptides. Calculates proteotypicity, and then uses calcualtions in histogram.

Usage

```
trypticHistogramLiP(
  data,
  fasta,
  x.axis.size = 10,
  y.axis.size = 10,
  legend.size = 10,
  width = 12,
  height = 4,
  color_scale = "bright",
  address = ""
)
```

Arguments

data	output of MSstatsLiP converter function. Must include at least ProteinName, PeptideSequence, BioReplicate, and Condition columns
fasta	A string of path to a FASTA file, used to match LiP peptides.
x.axis.size	size of x-axis labeling for plot. Default is 10.
y.axis.size	size of y-axis labeling for plot. Default is 10.
legend.size	size of feature legend for half vs fully tryptic peptides below graph. Default is 7.
width	Width of final pdf to be plotted
height	Height of final pdf to be plotted
color_scale	colors of bar chart. Must be one of "bright" or "grey". Default is "bright".
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 "TyrpticPlot.pdf". If address=FALSE, plot will be not saved as pdf file but shown in window

Value

plot or pdf

Examples

Index

* datasets LiPRawData, 16 MSstatsLiP_data, 20 MSstatsLiP_model, 21 MSstatsLiP_Summarized, 22 raw_lip, 25 raw_prot, 26 SkylineTest, 29 TrPRawData, 36 annotSite, 3 calculateProteolyticResistance, 3 calculateTrypticity, 4 correlationPlotLiP, 5, 19 dataProcessPlotsLiP, 6, 19 dataSummarizationLiP, 7, 8, 13, 19, 24, 28 DIANNtoMSstatsLiPFormat, 11 groupComparisonLiP, 13, 15, 20, 34 groupComparisonPlotsLiP, 14, 20 LiPRawData, 16 locateMod. 18 locatePTM, 18 MSstatsLiP, 19 $\texttt{MSstatsLiP_data, } 20$ MSstatsLiP_model, 21 MSstatsLiP_Summarized, 22 PCAPlotLiP, 20, 23 raw_lip, 25 raw_prot, 26 ResistanceBarcodePlotLiP, 28 SkylineTest, 29 SkylinetoMSstatsLiPFormat, 30 SpectronauttoMSstatsLiPFormat, 19, 32

StructuralBarcodePlotLiP, 20, 34

tidyFasta, 35 TrPRawData, 36 trypticHistogramLiP, *19*, 37