

Package ‘artMS’

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Description artMS provides a set of tools for the analysis of proteomics label-free datasets. It takes as input the MaxQuant search result output (evidence.txt file) and performs quality control, relative quantification using MSstats, downstream analysis and integration. artMS also provides a set of functions to re-format and make it compatible with other analytical tools, including, SAINTq, SAINTexpress, Phosphate, and PHOTON. Check <http://artms.org> for details.

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URL <https://github.com/biodavidjm/artMS>

BugReports <https://github.com/biodavidjm/artMS/issues>

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artmsAnalysisQuantifications
Analysis of the Relative Quantifications

Description

Analysis of relative quantifications, including:

- Annotations
- Summary files in different format (xls, txt) and shapes (long, wide)
- Numerous summary plots
- Enrichment analysis using Gprofiler
- PCA of quantifications
- Clustering analysis
- Basic imputation of missing values

Usage

```
artmsAnalysisQuantifications(  
  log2fc_file,  
  modelqc_file,  
  species,  
  output_dir = "analysis_quant",  
  outliers = c("keep", "iqr", "std"),  
  enrich = TRUE,  
  l2fc_thres = 1,  
  choosePvalue = c("adjpvalue", "pvalue"),  
  isBackground = "nobackground",  
  isPtm = "global",  
  mnbr = 2,  
  isFluomics = FALSE,  
  pathogen = "nopathogen",  
  plotPvaluesLog2fcDist = TRUE,  
  plotAbundanceStats = TRUE,  
  plotReproAbundance = TRUE,  
  plotCorrConditions = TRUE,  
  plotCorrQuant = TRUE,  
  plotPCAabundance = TRUE,  
  plotFinalDistributions = TRUE,  
  plotPropImputation = TRUE,  
  plotHeatmapsChanges = TRUE,  
  plotTotalQuant = TRUE,  
  plotClusteringAnalysis = TRUE,  
  verbose = TRUE  
)
```

Arguments

log2fc_file	(char) MSstats results file location
modelqc_file	(char) MSstats modelqc file location
species	(char) Select one species. Species currently supported for a full analysis (including enrichment analysis): <ul style="list-style-type: none"> • HUMAN • MOUSE <p>To find out species supported only for annotation check ?artmsIsSpeciesSupported()</p>
output_dir	(char) Name for the folder to output the results from the function. Default is current directory (recommended to provide a new folder name).
outliers	(char) It allows to keep or remove outliers. Options: <ul style="list-style-type: none"> • keep (default): it keeps outliers 'keep', 'iqr', 'std' • iqr (recommended): remove outliers +/- 6 x Interquartile Range (IQR) • std : 6 x standard deviation
enrich	(logical) Performed enrichment analysis using GprofileR? Only available for species HUMAN and MOUSE. TRUE (default if "human" or "mouse" are the species) or FALSE
l2fc_thres	(int) log2fc cutoff for enrichment analysis (default, l2fc_thres = 1.5)
choosePvalue	(char) specify whether pvalue or adjpvalue should use for the analysis. The default option is adjpvalue (multiple testing correction). But if the number of biological replicates for a given experiment is too low (for example n = 2), then choosePvalue = pvalue is recommended.
isBackground	(char) background of gene names for enrichment analysis. nobackground (default) will use the total number of genes detected. Alternatively provided the file path name to the background gene list.
isPtm	(char) Is a ptm-site quantification? <ul style="list-style-type: none"> • global (default), • ptmsites (for site specific analysis), • ptmph (Jeff Johnson script output evidence file)
mnbr	(int) minimal number of biological replicates for imputation and filtering. Default: mnbr = 2 (Proteins must be found in one of the conditions in at least 2 of the biological replicates)
isFluomics	(logical) Does this data belong to the FluOMICs project? TRUE or FALSE (default)
pathogen	(char) Is there a pathogen in the dataset as well? if it does not, then use pathogen = nopathogen (default). Pathogens available: tb (Tuberculosis), lpn (Legionella)
plotPvaluesLog2fcDist	(logical) If TRUE (default) plots pvalues and log2fc distributions
plotAbundanceStats	(logical) If TRUE (default) plots stats graphs about abundance values
plotReproAbundance	(logical) If TRUE plots reproducibility based on normalized abundance values
plotCorrConditions	(logical) If TRUE plots correlation between the different conditions
plotCorrQuant	(logical) if TRUE plots correlation between the available quantifications (comparisons)

plotPCAabundance	(logical) if TRUE performs PCA analysis of conditions using normalized abundance values
plotFinalDistributions	(logical) if TRUE plots distribution of both log2fc and pvalues
plotPropImputation	(logical) if TRUE plots proportion of overall imputation
plotHeatmapsChanges	(logical) if TRUE plots heatmaps of quantified changes (both all and significant only)
plotTotalQuant	(logical) if TRUE plots barplot of total number of quantifications per comparison
plotClusteringAnalysis	(logical) if TRUE performs clustering analysis between quantified comparisons (more than 1 comparison required)
verbose	(logical) TRUE (default) shows function messages

Value

(data.frame) summary of quantifications, including annotations, enrichments, etc

Examples

```
# Testing that the files cannot be empty
artmsAnalysisQuantifications(log2fc_file = NULL,
                             modelqc_file = NULL,
                             species = NULL,
                             output_dir = NULL)
```

artmsAnnotateSpecie *Adding a column with the species name*

Description

Adding the species name to every protein. This makes more sense if there are more than one species in the dataset, which must be specified in the pathogen option. Influenza is a special case that it does not need to be specified, as far as the proteins were originally annotated as INFLUENZAGENE_STRAIN (strains covered H1N1, H3N2, H5N1), as for example, NS1_H1N1

Usage

```
artmsAnnotateSpecie(df, pathogen = "nopathogen", species)
```

Arguments

df	(data.frame) with a Protein column (of uniprot ids)
pathogen	(char) Is there a pathogen in the dataset as well? if it does not, then use pathogen = nopathogen (default). Supported tb (Tuberculosis), lpn (Legionella)
species	(char) Host organism (supported for now: human or mouse)

artmsAvgIntensityRT *Summarize average intensity and retention time per protein*

Description

Input an evidence file from MaxQuant and a file containing a list of proteins of interest (optional). The function will summarize from the evidence file and report back the average intensity, average retention time, and the average calibrated retention time. If a list of proteins is provided, then only those proteins will be summarized and returned.

Usage

```
artmsAvgIntensityRT(  
  evidence_file,  
  protein_file = NULL,  
  output_file = FALSE,  
  species,  
  verbose = TRUE  
)
```

Arguments

evidence_file	(char) The filepath to the MaxQuant searched data (evidence) file (txt tab delimited file).
protein_file	(char) The filepath to a file or vector containing a list of proteins of interest.
output_file	(char) The file name for the results (must have the extension .txt). If empty, then the results will be returned as an R object.
species	(char) The species name. Check ?artmsMapUniprot2Entrez for supported species
verbose	(logical) TRUE (default) shows function messages

Value

An R object with the results and a file with the results (if the output_file argument is provided). It contains averages of Intensity, Retention Time, Calibrated Retention Time

Examples

```
ave_int <- artmsAvgIntensityRT(evidence_file = artms_data_ph_evidence,  
                              species = "human")
```

artmsChangeColumnName *Change a specific column name in a given data.frame*

Description

Making easier to change a column name in any data.frame

Usage

```
artmsChangeColumnName(dataset, oldname, newname)
```

Arguments

dataset	(data.frame) with the column name you want to change
oldname	(char) the old column name
newname	(char) the new name for that column

Value

(data.frame) with the new specified column name

Examples

```
artms_data_ph_evidence <- artmsChangeColumnName(  
  dataset = artms_data_ph_evidence,  
  oldname = "Phospho..STY.",  
  newname = "PH_STY")
```

artmsConvertMetabolomics

Convert Markview Metabolomics file (alignment table) into a artMS compatible format

Description

artMS enables the relative quantification of untargeted polar metabolites using the alignment table generated by Markview. MarkerView is an ABSciex software that supports the files generated by Analyst software (.wiff) used to run our specific mass spectrometer (ABSciex Triple TOF 5600+). It also supports .t2d files generated by the Applied Biosystems 4700/4800 MALDI-TOF. MarkerView software is used to align mass spectrometry data from several samples for comparison. Using the import feature in the software, .wiff files (also .t2d MALDI-TOF files and tab-delimited .txt mass spectra data in mass-intensity format) are loaded for retention time alignment. Once the data files are selected, a series of windows will appear wherein peak finding, alignment, and filtering options can be entered and selected. These options include minimum spectral peak width, minimum retention time peak width, retention time and mass tolerance, and the ability to filter out peaks that do not appear in more than a user selected number of samples.

‘artmsConvertMetabolomics’ processes the markview file to enable QC analysis and relative quantification using the artMS functions

Usage

```
artmsConvertMetabolomics(input_file, out_file, id_file = NULL, verbose = TRUE)
```

Arguments

```
input_file      (char) Markview input file
out_file        (char) Output file name
id_file         (char) KEGG database
verbose         (logical) TRUE (default) shows function messages
```

Value

(text file) Outputs the converted output name

Examples

```
# Testing that the arguments cannot be null
artmsConvertMetabolomics(input_file = NULL,
                          out_file = NULL)
```

artmsDataPlots

Individual Normalized abundance dot plots for every protein

Description

Protein abundance dot plots for each unique uniprot id. It can take a long time

Usage

```
artmsDataPlots(input_file, output_file, verbose = TRUE)
```

Arguments

```
input_file      (char) File path and name to the -normalized.txt output file from MSstats
output_file     (char) Output file (path) name (add the .pdf extension)
verbose         (logical) TRUE (default) shows function messages
```

Value

(pdf) file with each individual protein abundance plot for each conditions

Examples

```
artmsDataPlots(input_file = "results/ab-results-mss-normalized.txt",
                output_file = "results/ab-results-mss-normalized.pdf")
```

artmsEnrichLog2fc *Enrichment of changes in protein abundance or PTMs*

Description

Enrichment analysis of the selected proteins

Usage

```
artmsEnrichLog2fc(  
  dataset,  
  species,  
  background,  
  heatmaps = FALSE,  
  output_name = "enrichment.txt",  
  verbose = TRUE  
)
```

Arguments

dataset	(data.frame) with a Gene and Comparison or Label (with the name of the comparisons specified in the contrast file) columns
species	(char) Specie, only supported "human" or "mouse"
background	(vector) Background genes for the enrichment analysis.
heatmaps	(logical) if TRUE generates heatmaps (pdf), FALSE (default) otherwise.
output_name	(char) Name of the annotation files, which will be used as well for the heatmaps (if heatmaps is selected) Default output_name = "enrichment.txt"
verbose	(logical) TRUE (default) shows function messages

Value

(data.frame) Results from the enrichment analysis using Gprofiler and heatmaps (if selected)

Examples

```
# The data must be annotated (Protein and Gene columns)  
data_annotated <- artmsAnnotationUniprot(  
  x = artms_data_ph_msstats_results,  
  columnid = "Protein",  
  species = "human")  
  
# And then the enrichment  
enrich_set <- artmsEnrichLog2fc(  
  dataset = data_annotated,  
  species = "human",  
  background = unique(data_annotated$Gene))
```

 artmsEnrichProfiler *Enrichment analysis using GprofileR*

Description

This function simplifies the enrichment analysis performed by the excellent tool GprofileR.

Usage

```
artmsEnrichProfiler(
  x,
  categorySource = c("GO"),
  species,
  background = NA,
  verbose = TRUE
)
```

Arguments

- x** (list, data.frame) List of protein ids. It can be anything: either a list of ids, or you could also send a data.frame and it will find the columns with the IDs. Is not cool? Multiple list can be also sent simultaneously, as for example running: `tmp <- split(enrichment$Gene, enrichment$cl_number, drop= TRUE)`
- categorySource** (vector) Resources providing the terms on which the enrichment will be performed. The supported resources by gprofiler are:
- GO (GO:BP, GO:MF, GO:CC): Gene Ontology (see more below)
 - KEGG: Biological pathways
 - REAC: Biological pathways (Reactome)
 - TF: Regulatory motifs in DNA (TRANSFAC TFBS)
 - MI: Regulatory motifs in DNA (miRBase microRNAs)
 - CORUM: protein complexes database
 - HP: Human Phenotype Ontology
 - HPA: Protein databases (Human Protein Atlas)
 - OMIM: Online Mendelian Inheritance in Man annotations:
 - BIOGRID: BioGRID protein-protein interactions The type of annotations for Gene Ontology:
 - Inferred from experiment (IDA, IPI, IMP, IGI, IEP)
 - Direct assay (IDA) / Mutant phenotype (IMP)
 - Genetic interaction (IGI) / Physical interaction (IPI)
 - Traceable author (TAS) / Non-traceable author (NAS) / Inferred by curator (IC)
 - Expression pattern (IEP) / Sequence or structural similarity (ISS) / Genomic context (IGC)
 - Biological aspect of ancestor (IBA) / Rapid divergence (IRD)
 - Reviewed computational analysis (RCA) / Electronic annotation (IEA)
 - No biological data (ND) / Not annotated or not in background (NA)

species	(char) Specie code: Organism names are constructed by concatenating the first letter of the name and the family name. Example: human - 'hsapiens', mouse - 'mmusculus'. Check gProfileR to find out more about supported species.
background	(vector) gene list to use as background for the enrichment analysis. Default: NA
verbose	(logical) TRUE (default) shows function messages

Details

This function uses the following gprofiler arguments as default:

- ordered_query = FALSE
- significant = TRUE
- exclude_iea = TRUE
- underrep = FALSE
- evcodes = FALSE
- region_query = FALSE
- max_p_value = 0.05
- min_set_size = 0
- max_set_size = 0
- min_isect_size = 0
- correction_method = "analytical" #Options: "gSCS", "fdr", "bonferroni"
- hier_filtering = "none"
- domain_size = "known" # annotated or known
- numeric_ns = ""
- png_fn = NULL
- include_graph = TRUE

Value

The enrichment results as provided by gprofiler

Examples

```
# annotate the MSstats results to get the Gene name
data_annotated <- artmsAnnotationUniprot(
  x = artms_data_ph_msstats_results,
  columnid = "Protein",
  species = "human")

# Filter the list of genes with a log2fc > 2
filtered_data <-
unique(data_annotated$Gene[which(data_annotated$log2FC > 2)])

# And perform enrichment analysis
data_annotated_enrich <- artmsEnrichProfiler(
  x = filtered_data,
  categorySource = c('KEGG'),
  species = "hsapiens",
  background = unique(data_annotated$Gene))
```

`artmsEvidenceToSaintExpress`*MaxQuant evidence file to SAINTexpress format*

Description

Converts the MaxQuant evidence file to the 3 required files by SAINTexpress. One can choose to either use the spectral counts (use `msspc`) or the intensities (use `msint`) for the analysis.

Usage

```
artmsEvidenceToSaintExpress(  
  evidence_file,  
  keys_file,  
  ref_proteome_file,  
  quant_variable = c("msspc", "msint"),  
  output_file,  
  verbose = TRUE  
)
```

Arguments

<code>evidence_file</code>	(char) The evidence file path and name
<code>keys_file</code>	(char) Keys file with a SAINT column specifying test (T) and control (C) conditions
<code>ref_proteome_file</code>	(char) Reference proteome path file name in fasta format
<code>quant_variable</code>	(char) choose either <ul style="list-style-type: none">• <code>msspc</code> (spectral counts, default) or• <code>msint</code> (MS Intensity)
<code>output_file</code>	(char) Output file name (must have extension <code>.txt</code>)
<code>verbose</code>	(logical) TRUE (default) shows function messages

Value

The 3 required files by SAINTexpress:

- `interactions.txt`
- `preys.txt`
- `baits.txt`

Examples

```
# Testing that the files cannot be empty  
artmsEvidenceToSaintExpress(evidence_file = NULL,  
  keys_file = NULL, ref_proteome_file = NULL)
```

artmsEvidenceToSAINTq *MaxQuant evidence file to SAINTq format*

Description

Converts the MaxQuant evidence file to the required files by SAINTq. The user can choose to use either peptides with spectral counts (use `msspc`) or the all the peptides (use `all`) for the analysis. The quantitative can be also chosen (either MS Intensity or Spectral Counts)

Usage

```
artmsEvidenceToSAINTq(
  evidence_file,
  keys_file,
  output_dir = ".",
  sc_option = c("all", "msspc"),
  fractions = FALSE,
  quant_variable = c("msint", "msspc"),
  verbose = TRUE
)
```

Arguments

<code>evidence_file</code>	(char or data.frame) The evidence file path and name, or data.frame
<code>keys_file</code>	(char) Keys file with a SAINT column specifying test (T) and control (C) conditions
<code>output_dir</code>	(char) New directory to create and save files. Default is current directory (recommended to provide a new folder name).
<code>sc_option</code>	(char). Filter peptides with spectral counts only. Two options: <ul style="list-style-type: none"> • <code>msspc</code>: use only peptides with <code>spectral_counts</code> • <code>all</code> (default): all peptides detected (including the one resulting from the MaxQuant 'Match between run' algorithm)
<code>fractions</code>	(logical) TRUE for 2D proteomics (fractions). Default: FALSE
<code>quant_variable</code>	(char) Select the quantitative variable. Two options available: <ul style="list-style-type: none"> • <code>msint</code>: MS Intensity (default) • <code>msspc</code>: MS.MS.count (Spectral Counts)
<code>verbose</code>	(logical) TRUE (default) shows function messages

Details

After running the script, the new specified folder should contain the following files:

- `saintq-config-peptides`
- `saintq-config-proteins`
- `saintq_input_peptides.txt`
- `saintq_input_proteins.txt`

Then cd into the new folder and run either of the following two options (assuming that saintq is installed in your linux/unix/mac os x system):

```
> saintq config-saintq-peptides
```

or

```
> saintq config-saintq-proteins
```

Value

The input files requires to run SAINTq

Examples

```
# Testing that the files cannot be empty
artmsEvidenceToSAINTq (evidence_file = NULL,
                      keys_file = NULL,
                      output_dir = NULL)
```

artmsFilterEvidenceContaminants

Remove contaminants and empty proteins from the MaxQuant evidence file

Description

Remove contaminants and erroneously identified 'reverse' sequences by MaxQuant, in addition to empty protein ids

Usage

```
artmsFilterEvidenceContaminants(x, verbose = TRUE)
```

Arguments

x	(data.frame) of the Evidence file
verbose	(logical) TRUE (default) shows function messages

Value

(data.frame) without REV__ and CON__ Protein ids

Examples

```
ef <- artmsFilterEvidenceContaminants(x = artms_data_ph_evidence)
```

artmsIsEvidenceNewVersion

Check if a given evidence file was generated by a new version of MaxQuant (v>1)

Description

MaxQuant introduced changes in the column names and number of columns for the evidence file in version 1 (we think). This function check whether the evidence comes from the latest version of MaxQuant.

Usage

```
artmsIsEvidenceNewVersion(evidence_file)
```

Arguments

evidence_file the evidence file name

Value

(logical) TRUE if it is a newer version of MaxQuant, FALSE otherwise

Examples

```
artmsIsEvidenceNewVersion(evidence_file = artms_data_ph_evidence)
```

artmsIsSpeciesSupported

Check if a species is supported and available

Description

Given a species name, it checks whether is supported, and if supported, check whether the annotation package is installed.

Usage

```
artmsIsSpeciesSupported(species, verbose = TRUE)
```

Arguments

species (char) The species name. Species currently supported as part of artMS:

- HUMAN
- MOUSE

And the following species can be used as well, but the user needs to install the corresponding org.db package:

- ANOPHELES (install.packages(org.Ag.eg.db))
- BOVINE (install.packages(org.Bt.eg.db))

- WORM (install.packages(org.Ce.eg.db))
- CANINE (install.packages(org.Cf.eg.db))
- FLY (install.packages(org.Dm.eg.db))
- ZEBRAFISH (install.packages(org.Dr.eg.db))
- CHICKEN (install.packages(org.Gg.eg.db))
- RHESUS (install.packages(org.Mmu.eg.db))
- CHIMP (install.packages(org.Pt.eg.db))
- RAT (install.packages(org.Rn.eg.db))
- YEAST (install.packages(org.Sc.sgd.db))
- PIG (install.packages(org.Ss.eg.db))
- XENOPUS (install.packages(org.Xl.eg.db))

verbose (logical) TRUE (default) shows function messages

Value

(string) Name of the package for the given species

Examples

```
# Should return TRUE
artmsIsSpeciesSupported(species = "HUMAN")
artmsIsSpeciesSupported(species = "CHIMP")
```

artmsLeaveOnlyUniprotEntryID

Leave only the Entry ID from a typical full Uniprot IDs in a given column

Description

Downloading a Reference Uniprot fasta database includes several Uniprot IDs for every protein. If the regular expression available in Maxquant is not activated, the full id will be used in the Proteins, Lead Protein, and Leading Razor Protein columns. This script leaves only the Entry ID.

For example, values in a Protein column like this:

```
sp|P12345|Entry_name;sp|P54321|Entry_name2
```

will be replace by

```
'P12345;P54321'
```

Usage

```
artmsLeaveOnlyUniprotEntryID(x, columnid)
```

Arguments

x (data.frame) that contains the columnid
 columnid (char) Column name with the full uniprot ids

Value

(data.frame) with only Entry IDs.

Examples

```
# Example of data frame with full uniprot ids and sequences
p <- c("sp|A6NIE6|RN3P2_HUMAN;sp|Q9NYV6|RRN3_HUMAN",
      "sp|A7E2V4|ZSWM8_HUMAN",
      "sp|A5A6H4|ROA1_PANTR;sp|P09651|ROA1_HUMAN;sp|Q32P51|RA1L2_HUMAN",
      "sp|A0FGR8|ESYT2_HUMAN")
s <- c("ALENDFNSPPRK", "GWGSPGRPK", "SSGPYGGGGQYFAK", "VLVALASEELAK")
evidence <- data.frame(Proteins = p, Sequences = s, stringsAsFactors = FALSE)

# Replace the Proteins column with only Entry ids
evidence <- artmsLeaveOnlyUniprotEntryID(x = evidence, columnid = "Proteins")
```

```
artmsMapUniprot2Entrez
```

Map GENE SYMBOL, NAME, AND ENTREZID to a vector of Uniprot IDS

Description

Map GENE SYMBOL, NAME, AND ENTREZID to a vector of Uniprot IDS

Usage

```
artmsMapUniprot2Entrez(uniprotkb, species)
```

Arguments

uniprotkb	(vector) Vector of UniprotKB IDs
species	(char) The species name. Species currently supported as part of artMS: check ?artmsIsSpeciesSupported() to find out the list of supported species'

Value

(data.frame) with ENTREZID and GENENAMES mapped on UniprotKB ids

Examples

```
# Load an example with human proteins
exampleID <- c("Q6P996", "B1N8M6")
artmsMapUniprot2Entrez(uniprotkb = exampleID,
                      species = "HUMAN")
```

```
artmsMergeEvidenceAndKeys
```

Merge evidence.txt (or summary.txt) with keys.txt files

Description

Merge the evidence and keys files on the given columns

Usage

```
artmsMergeEvidenceAndKeys(
  x,
  keys,
  by = c("RawFile"),
  isSummary = FALSE,
  verbose = TRUE
)
```

Arguments

x	(data.frame or char) The evidence data, either as data.frame or the file name (and path). It also works for the summary.txt file
keys	The keys data, either as a data.frame or file name (and path)
by	(vector) specifying the columns use to merge the evidence and keys. Default: by=c('RawFile')
isSummary	(logical) TRUE or FALSE (default)
verbose	(logical) TRUE (default) shows function messages

Value

(data.frame) with the evidence and keys merged

Examples

```
evidenceKeys <- artmsMergeEvidenceAndKeys(x = artms_data_ph_evidence,
                                          keys = artms_data_ph_keys)
```

```
artmsMsstatsSummary
```

Summarize the MSStats results and data quantification

Description

Converts the MSStats results file to wide format (unique Protein ID and columns are the comparisons), as well as adds BioReplicate information about

- the Number of Unique Peptides,
- Spectral Counts
- Intensities for each protein. In cases where there are multiple values for a Protein-BioReplicate pair due to minute changes in sequence, the maximum value is taken for the pair. Any pairs without a value are assigned a value of NA.

Usage

```
artmsMsstatsSummary(
  evidence_file,
  prot_group_file,
  keys_file,
  results_file,
  return_df = FALSE,
  verbose = TRUE
)
```

Arguments

`evidence_file` (char or data.frame) The filepath to the MaxQuant searched data (evidence) file (txt tab delimited file). Only works for the newer versions of the evidence file.

`prot_group_file` (char) The filepath to the MaxQuant proteinGroups.txt file (txt tab delimited file) or data.frame

`keys_file` (char) The filepath to the keys file used with MSSstats (txt tab delimited file).

`results_file` (char) The filepath to the MSSstats results file in the default long format (txt tab delimited file or data.frame).

`return_df` (data.frame) Whether or not to return the results to the R environment upon completion. This is useful if this is being used in an R pipeline and you want to feed the results directly into the next stage of analysis via an R environment/terminal. Regardless, the results will be written to file. Default = FALSE

`verbose` (logical) TRUE (default) shows function messages

Value

(data.frame or txt file) with the summary

Examples

```
# Testing warning if files are not submitted
test <- artmsMsstatsSummary(evidence_file = NULL,
  prot_group_file = NULL,
  keys_file = NULL,
  results_file = NULL)
```

artmsPhosphateOutput *Generate Phosphate Input file*

Description

It takes as input the imputedL2fcExtended.txt results generated by the artmsAnalysisQuantifications() function and generates the **Phosphate** input file (or data.frame) Please, notice that the only species supported by Phosphate is humans.

Usage

```
artmsPhosphateOutput(inputFile, output_dir = ".", verbose = TRUE)
```

Arguments

inputFile	(char) the imputedL2fcExtended.txt file name and location
output_dir	(char) Name of the folder to output results (Default: current directory. Recommended: phosphate_input)
verbose	(logical) TRUE (default) to show function messages

Value

Multiple output files (inputs of phosphate)

Examples

```
artmsPhosphateOutput(inputFile)
```

artmsPhotonOutput	<i>Generate PHOTON Input file</i>
-------------------	-----------------------------------

Description

It takes as input the imputedL2fcExtended.txt results generated by the artmsAnalysisQuantifications() function and generates the **PHOTON** input file. Please, notice that the only species supported by PHOTON is humans.

Usage

```
artmsPhotonOutput(inputFile, output_dir = ".", verbose = TRUE)
```

Arguments

inputFile	(char) the imputedL2fcExtended.txt file name and location
output_dir	(char) Name of the folder to output results (Default: current. Recommended: "photon_input_files" or similar)
verbose	(logical) TRUE (default) to show function messages

Value

Multiple output files (inputs of phosphate)

Examples

```
artmsPhotonOutput(inputFile)
```

artmsPlotHeatmapQuant *Outputs a heatmap of the MSStats results created using the log2fold changes*

Description

Heatmap of the Relative Quantifications (MSStats results)

Usage

```
artmsPlotHeatmapQuant(
  input_file,
  output_file = "quantifications_heatmap.pdf",
  species,
  labels = "*",
  cluster_cols = FALSE,
  display = "log2FC",
  lfc_lower = -2,
  lfc_upper = 2,
  whatPvalue = "adj.pvalue",
  FDR = 0.05,
  verbose = TRUE
)
```

Arguments

input_file	(char) MSstats results.txt file and location (or data.frame of results)
output_file	(char) Output file name (pdf format) and location. Default: "quantifications_heatmap.pdf"
species	(char). Specie name to be able to add the Gene name. To find out more about the supported species check ?artmsMapUniprot2Entrez
labels	(vector) of uniprot ids if only specific labes would like to be plotted. Default: all labels
cluster_cols	(boolean) True or False to cluster columns. Default: FALSE
display	Metric to be displayed. Options: <ul style="list-style-type: none"> • log2fc (default) • adj.pvalue • pvalue
lfc_lower	(int) Lower limit for the log2fc. Default: -2
lfc_upper	(int) Upper limit for the log2fc. Default: +2
whatPvalue	(char) pvalue or adj.pvalue (default)
FDR	(int) Upper limit false discovery rate (or pvalue). Default: 0.05
verbose	(logical) TRUE (default) shows function messages

Value

(pdf or ggplot2 object) heatmap of the MSStats results using the selected metric

Examples

```
artmsPlotHeatmapQuant(input_file = artms_data_ph_msstats_results,
                      species = "human",
                      output_file = NULL,
                      whatPvalue = "pvalue",
                      lfc_lower = -1,
                      lfc_upper = 1)
```

```
artmsProtein2SiteConversion
```

Converts the Protein ID column of the evidence file selected by the user to mod-site-specific notation: ProteinID to ProteinID_AAnumber notation

Description

It enables the modified-peptide specific quantification by converting the Protein column of the evidence file selected by the user to an ProteinID_AAnumber notation. In this way, each of the modified peptides can be quantified independently across conditions.

!!

WARNING: we have detected a version of MaxQuant (>1.6.3.0) outputs a "Modified sequence" column of the evidence file that has two important changes for the annotation of phosphorylation:

- Uses p instead of (ph)
- The modified residue (i.e. STY) is the residue on the right of the p, instead of the residue to the left of (ph), as usual. We have introduced a modification to detect and address this issue, but we advice the user to double check both the new evidence file with the introduce new notation and the -mapping.txt file and check that there are no NA values for the notation of phophopeptides.

!!

Usage

```
artmsProtein2SiteConversion(
  evidence_file,
  ref_proteome_file,
  column_name = c("Leading razor protein", "Leading proteins", "Proteins"),
  output_file,
  mod_type,
  overwrite_evidence = FALSE,
  verbose = TRUE
)
```

Arguments

evidence_file (char) The evidence file name and location

ref_proteome_file

(char) The reference proteome used as database to search the evidence.txt file with MaxQuant. It will be used to map the modified peptide to the protein sequence and find the site location. Therefore, it does not use the MaxQuant's Phospho (STY)Sites.txt

column_name	(char) The Protein Column Name to map. Options: <ul style="list-style-type: none"> • Leadind razor protein (default) • Leading protein • Proteins It only supports Uniprot Entry IDs and RefSeq, but it might work for other database IDs
output_file	(char) Output file name (ptmsites-evidence.txt recommended)
mod_type	(char) The posttranslational modification. Options: <ul style="list-style-type: none"> • UB: Protein Ubiquitination • PH: Protein Phosphorylation • AC: Protein Acetylation
overwrite_evidence	(logical) if <output_file> is the same as <evidence_file>, overwrite_evidence = FALSE (default) doesn't allow to overwrite the evidence file. Otherwise, overwrite_evidence = TRUE allows to overwrite the evidence_file (this option might be activated if the user allows to use the same ptm-sites-evidence.txt file to re-annotate all the Protein IDs columns)
verbose	(logical) TRUE (default) shows function messages

Value

(file) Return a new evidence file with the specified Protein id column modified by adding the sequence site location(s) + postranslational modification(s) to the uniprot entry / refseq id.

Output ID examples: A34890_ph3; Q64890_ph24_ph456; Q64890_ub34_ub129_ub234; Q64890_ac35.

Examples

```
# Testing warning if files are not submitted.
artmsProtein2SiteConversion(evidence_file = NULL, ref_proteome_file = NULL,
output_file = NULL)
```

```
artmsQualityControlEvidenceBasic
```

Quality Control analysis of the MaxQuant evidence file

Description

Quality Control analysis of the MaxQuant evidence file

Usage

```
artmsQualityControlEvidenceBasic(
  evidence_file,
  keys_file,
  prot_exp = c("AB", "PH", "UB", "APMS"),
  fractions = 0,
  output_name = "qcPlots_evidence",
  isSILAC = FALSE,
  plotINTDIST = TRUE,
  plotREPRO = TRUE,
```

```

plotCORMAT = TRUE,
plotINTMISC = TRUE,
plotPTMSTATS = TRUE,
printPDF = TRUE,
verbose = TRUE
)

```

Arguments

evidence_file	(char or data.frame) The evidence file path and name, or data.frame
keys_file	(char or data.frame) The keys file path and name or data.frame
prot_exp	(char) Proteomics experiment. 4 options available: <ul style="list-style-type: none"> • APMS: affinity purification mass spectrometry • AB: protein abundance • PH: protein phosphorylation • UB: protein ubiquitination (aka ubiquitylation)
fractions	(binary) Is a fractionated experiment? <ul style="list-style-type: none"> • 1 yes • 0 no (default)
output_name	(char) prefix output name (no extension). Default: "qcPlots_evidence"
isSILAC	if TRUE processes SILAC input files. Default is FALSE
plotINTDIST	if TRUE (default) plots both <i>Box-dot plot</i> and <i>Jitter plot</i> of biological replicates based on MS (raw) intensity values.
plotREPRO	if TRUE (default) plots a correlation dotplot for all the combinations of biological replicates of conditions, based on MS Intensity values using features (peptide+charge)
plotCORMAT	if TRUE (default) plots a <ul style="list-style-type: none"> • <i>Correlation matrix</i> for all the biological replicates using MS Intensity values, • <i>Clustering matrix</i> of the MS Intensities and correlation distribution • <i>histogram</i> of the distribution of correlations
plotINTMISC	if TRUE (default) plots several pages, including bar plots of <i>Total Sum of Intensities in BioReplicates</i> , <i>Total Sum of Intensities in Conditions</i> , <i>Total Peptide Counts in BioReplicates</i> , <i>Total Peptide Counts in conditions</i> separated by categories: CON: contaminants, PROT peptides, REV reversed sequences used by MaxQuant to estimate the FDR; <i>Box plots</i> of MS Intensity values per biological replicates and conditions; <i>bar plots</i> of total intensity (excluding contaminants) by bioreplicates and conditions; Barplots of <i>total feature counts</i> by bioreplicates and conditions.
plotPTMSTATS	IF TRUE (default) plots stats related to the selected modification, including: <i>bar plot of peptide counts and intensities</i> , broken by PTM/other categories; bar plots of <i>total sum-up of MS intensity values</i> by other/PTM categories.
printPDF	If TRUE (default) prints out the pdfs. Warning: plot objects are not returned due to the large number of them.
verbose	(logical) TRUE (default) shows function messages

Value

Quality control files and plots

Examples

```
artmsQualityControlEvidenceBasic(evidence_file = artms_data_ph_evidence,
                                  keys_file = artms_data_ph_keys,
                                  prot_exp = "PH",
                                  isSILAC = FALSE,
                                  plotINTDIST = FALSE,
                                  plotREPRO = TRUE,
                                  plotCORMAT = FALSE,
                                  plotINTMISC = FALSE,
                                  plotPTMSTATS = FALSE,
                                  printPDF = FALSE,
                                  verbose = FALSE)

# But we recommend the following test:
# 1. Go to a working directory:
# setwd("/path/to/your/working/directory/")
# 2. Run the following command to print out all the pdf files
# artmsQualityControlEvidenceBasic(evidence_file = artms_data_ph_evidence,
#                                   keys_file = artms_data_ph_keys,
#                                   prot_exp = "PH")
# 3. Check your working directory and you should find pdf files with
# all the QC plots
```

artmsQualityControlEvidenceExtended

Extended Quality Control of the MaxQuant evidence.txt file

Description

Performs quality control based on the information available in the MaxQuant evidence.txt file.

Usage

```
artmsQualityControlEvidenceExtended(
  evidence_file,
  keys_file,
  isSILAC = FALSE,
  plotPSM = TRUE,
  plotIONS = TRUE,
  plotTYPE = TRUE,
  plotPEPTIDES = TRUE,
  plotPROTEINS = TRUE,
  plotPIO = TRUE,
  plotCS = TRUE,
  plotME = TRUE,
  plotMOCD = TRUE,
  plotPEPICV = TRUE,
  plotPEPDETECT = TRUE,
  plotPROTICV = TRUE,
  plotPROTDETECT = TRUE,
  plotIDoverlap = TRUE,
  plotIC = TRUE,
```

```

plotSP = TRUE,
printPDF = TRUE,
verbose = TRUE
)

```

Arguments

evidence_file	(char or data.frame) The evidence file path and name, or data.frame
keys_file	(char or data.frame) The keys file path and name or data.frame
isSILAC	if TRUE processes SILAC input files. Default is FALSE
plotPSM	(logical) TRUE generates peptide-spectrum-matches (PSMs) statistics plot: Page 1 shows the number of PSMs confidently identified in each BioReplicate. If replicates are present, Page 2 shows the mean number of PSMs per condition with error bar showing the standard error of the mean. Note that potential contaminant proteins are plotted separately.
plotIONS	(logical) TRUE generates peptide ion statistics plot: A peptide ion is defined in the context of m/z, in other words, an unique peptide sequence may give rise to multiple ions with different charge state and/or amino acid modification. Page 1 shows the number of ions confidently identified in each BioReplicate. If replicates are present, Page 2 shows the mean number of peptide ions per condition with error bar showing the standard error of the mean. Note that potential contaminant proteins are plotted separately.
plotTYPE	(logical) TRUE generates identification type statistics plot: MaxQuant classifies each peptide identification into different categories (e.g., MSMS, MULTI-MSMS, MULTI-SECPEP). Page 1 shows the distribution of identification type in each BioReplicate
plotPEPTIDES	(logical) TRUE generates peptide statistics plot: Page 1 shows the number of unique peptide sequences (disregard the charge state or amino acid modifications) confidently identified in each BioReplicate. If replicates are present, Page 2 shows the mean number of peptides per condition with error bar showing the standard error of the mean. Note that potential contaminant proteins are plotted separately. Pages 3 and 4 show peptide identification intersection between BioReplicates (the bars are ordered by degree or frequency, respectively), and Page 4 shows the intersections across conditions instead of BioReplicates.
plotPROTEINS	(logical) TRUE generates protein statistics plot: Page 1 shows the number of protein groups confidently identified in each BioReplicate. If replicates are present, Page 2 shows the mean number of protein groups per condition with error bar showing the standard error of the mean. Note that potential contaminant proteins are plotted separately. Pages 3 and 4 show peptide identification intersection between BioReplicates (the bars are ordered by degree or frequency, respectively), and Page 4 shows the intersections across conditions instead of BioReplicates.
plotPIO	(logical) TRUE generates oversampling statistics plot: Page 1 shows the proportion of all peptide ions (including peptides matched across runs) fragmented once, twice and thrice or more. Page 2 shows the proportion of peptide ions (with intensity detected) fragmented once, twice and thrice or more. Page 3 shows the proportion of peptide ions (with intensity detected and MS/MS identification) fragmented once, twice and thrice or more
plotCS	(logical) TRUE generates charge state plot: Page 1 shows the charge state distribution of PSMs confidently identified in each BioReplicate.

plotME	(logical) TRUE generates precursor mass error plot: Page 1 shows the distribution of precursor error for all PSMs confidently identified in each BioReplicate.
plotMOCD	(logical) TRUE generates precursor mass-over-charge plot: Page 1 shows the distribution of precursor mass-over-charge for all PSMs confidently identified in each BioReplicate.
plotPEPICV	(logical) TRUE generates peptide intensity coefficient of variance (CV) plot: The CV is calculated for each feature (peptide ion) identified in more than one replicate. Page 1 shows the distribution of CV's for each condition, while Page 2 shows the distribution of CV's within 4 bins of intensity (i.e., 4 quantiles of average intensity).
plotPEPDETECT	(logical) TRUE generates peptide detection frequency plot: Page 1 summarizes the frequency that each peptide is detected across BioReplicates of each condition, showing the percentage of peptides detected once, twice, thrice, and so on (for whatever number of replicates each condition has).
plotPROTICV	(logical) TRUE generates protein intensity coefficient of variance (CV) plot: The CV is calculated for each protein (after summing the peptide intensities) identified in more than one replicate. Page 1 shows the distribution of CV's for each condition, while Page 2 shows the distribution of CV's within 4 bins of intensity (i.e., 4 quantiles of average intensity).
plotPROTDETECT	(logical) TRUE generates protein detection frequency plot: Page 1 summarizes the frequency that each protein group is detected across BioReplicates of each condition, showing the percentage of proteins detected once, twice, thrice, and so on (for whatever number of replicates each condition has). Page 2 shows the feature (peptide ion) intensity distribution within each BioReplicate (potential contaminant proteins are plot separately). Page 3 shows the density of feature intensity for different feature types (i.e., MULTI-MSMS, MULTI-SECPEP).
plotIDoverlap	(logical) TRUE generates pairwise identification heatmap overlap: Pages 1 and 2 show pairwise peptide and protein overlap between any 2 BioReplicates, respectively.
plotIC	(logical) TRUE generates pairwise intensity correlation: Page 1 and 3 show pairwise peptide and protein intensity correlation and scatter plot between any 2 BioReplicates, respectively. Page 2 and 4 show principal component analysis at the intensity level for both peptide and proteins, respectively.
plotSP	(logical) TRUE generates sample quality metrics: Page 1 shows missing cleavage distribution of all peptides confidently identified in each BioReplicate. Page 2 shows the fraction of peptides with at least one methionine oxidized in each BioReplicate.
printPDF	If TRUE (default) prints out the pdfs. Warning: plot objects are not returned due to the large number of them.
verbose	(logical) TRUE (default) shows function messages

Details

all the plots are generated by default

Value

A number of QC plots based on the evidence file

Examples

```
# Testing warning if files are not submitted
test <- artmsQualityControlEvidenceExtended(evidence_file = NULL,
keys_file = NULL)
```

```
artmsQualityControlMetabolomics
```

Quality Control analysis of the evidence-like metabolomics dataset

Description

Quality Control analysis of the evidence-like metabolomics dataset

Usage

```
artmsQualityControlMetabolomics(
  evidence_file,
  keys_file,
  met_exp = c("MV"),
  output_name = "qcPlots_metab",
  plotINTDIST = TRUE,
  plotREPRO = TRUE,
  plotCORMAT = TRUE,
  plotINTMISC = TRUE,
  printPDF = TRUE,
  verbose = TRUE
)
```

Arguments

evidence_file	(char or data.frame) The evidence file path and name, or data.frame
keys_file	(char or data.frame) The keys file path and name or data.frame
met_exp	(char) Proteomics experiment. Only one option available (so far): <ul style="list-style-type: none"> • MV: Markview output
output_name	(char) prefix output name (no extension). Default: "qcPlots_metab"
plotINTDIST	if TRUE (default) plots both <i>Box-dot plot</i> and <i>Jitter plot</i> of biological replicates based on MS (raw) intensity values.
plotREPRO	if TRUE (default) plots a correlation dotplot for all the combinations of biological replicates of conditions, based on MS Intensity values using features (mz_rt+charge)
plotCORMAT	if TRUE (default) generates up to 3 pdf files for technical replicates, biological replicates, and conditions. Each pdf file contains: <ul style="list-style-type: none"> • <i>Correlation matrix</i> for all the biological replicates using MS Intensity values, • <i>Clustering matrix</i> of the MS Intensities and correlation distribution • <i>histogram</i> of the distribution of correlations

plotINTMISC	if TRUE (default) plots several pages, including bar plots of <i>Total Sum of Intensities in BioReplicates</i> , <i>Total Sum of Intensities in Conditions</i> , <i>Total Feature Counts in BioReplicates</i> , <i>Total Feature Counts in conditions</i> separated by categories (INT: has a intensity value NOINT: no intensity value) <i>Box plots</i> of MS Intensity values per biological replicates and conditions; <i>bar plots</i> of total intensity by bioreplicates and conditions; <i>Barplots of total feature counts</i> by bioreplicates and conditions.
printPDF	If TRUE (default) prints out the pdfs. Warning: plot objects are not returned due to the large number of them.
verbose	(logical) TRUE (default) shows function messages

Value

Quality control files and plots for metabolomics

Examples

```
# Testing that input arguments cannot be null
artmsQualityControlMetabolomics(evidence_file = NULL,
                                keys_file = NULL,
                                met_exp = "MV")
```

artmsQualityControlSummaryExtended

Quality Control of the MaxQuant summary.txt file

Description

Performs quality control based on the information available in the MaxQuant summary.txt file.

Usage

```
artmsQualityControlSummaryExtended(
  summary_file,
  keys_file,
  isFractions = FALSE,
  plotMS1SCANS = TRUE,
  plotMS2 = TRUE,
  plotMSMS = TRUE,
  plotISOTOPE = TRUE,
  verbose = TRUE
)
```

Arguments

summary_file	(char or data.frame) The evidence file path and name, or data.frame
keys_file	(char or data.frame) The keys file path and name or data.frame
isFractions	(logical) TRUE if it is a 2D experiment (fractions). Default: FALSE

plotMS1SCANS	(logical) TRUE generates MS1 scan counts plot: Page 1 shows the number of MS1 scans in each BioReplicate. If replicates are present, Page 2 shows the mean number of MS1 scans per condition with error bar showing the standard error of the mean. If isFractions TRUE, each fraction is a stack on the individual bar graphs.
plotMS2	(logical) TRUE generates MS2 scan counts plot: Page 1 shows the number of MSs scans in each BioReplicate. If replicates are present, Page 2 shows the mean number of MS1 scans per condition with error bar showing the standard error of the mean. If isFractions TRUE, each fraction is a stack on the individual bar graphs.
plotMSMS	(logical) TRUE generates MS2 identification rate (%) plot: Page 1 shows the fraction of MS2 scans confidently identified in each BioReplicate. If replicates are present, Page 2 shows the mean rate of MS2 scans confidently identified per condition with error bar showing the standard error of the mean. If isFractions TRUE, each fraction is a stack on the individual bar graphs.
plotISOTOPE	(logical) TRUE generates Isotope Pattern counts plot: Page 1 shows the number of Isotope Patterns with charge greater than 1 in each BioReplicate. If replicates are present, Page 2 shows the mean number of Isotope Patterns with charge greater than 1 per condition with error bar showing the standard error of the mean. If isFractions TRUE, each fraction is a stack on the individual bar graphs.
verbose	(logical) TRUE (default) shows function messages

Value

A number of plots from the summary file

Examples

```
# Testing warning if files are not submitted
test <- artmsQualityControlSummaryExtended(summary_file = NULL,
keys_file = NULL)
```

artmsQuantification *Relative quantification using MSstats*

Description

Relative quantification using MSstats including:

- plots
- quantifications (log2fc, pvalues, etc)
- normalized abundance values

Usage

```
artmsQuantification(yaml_config_file, verbose = TRUE)
```

Arguments

```
yaml_config_file
      (char) The yaml file name and location
verbose
      (logical) TRUE (default) shows function messages
```


Value

The relative quantification of the conditions and comparisons specified in the keys/contrast file resulting from running MSstats, in addition to quality control plots (if selected)

Examples

```
artmsQuantification("artms-ab-config.yaml")
```

artmsResultsWide	<i>Reshape the MSstats results file from long to wide format</i>
------------------	--

Description

Converts the normal MSStats results.txt file into "wide" format where each row represents a unique protein's results, and each column represents the comparison made by MSStats. The fold change and p-value of each comparison will be its own column.

Usage

```
artmsResultsWide(
  results_msstats,
  output_file = NULL,
  select_pvalues = c("adjpvalue", "pvalue"),
  species,
  verbose = TRUE
)
```

Arguments

results_msstats	(char) Input file name and location (MSstats results.txt file)
output_file	(char) Output file name and location (e.g. results-wide.txt). If NULL (default) returns an R object (data.frame)
select_pvalues	(char) Either <ul style="list-style-type: none"> • pvalue or • adjpvalue (default)
species	(char) Specie name for annotation purposes. Check ?artmsMapUniprot2Entrez to find out more about the supported species (e.g species = "human")
verbose	(logical) TRUE (default) shows function messages

Value

(output file tab delimited) reshaped file with unique protein ids and as many columns log2fc and adj.pvalues as comparisons available

Examples

```
ph_results_wide <- artmsResultsWide(
  results_msstats = artms_data_ph_msstats_results,
  output_file = NULL,
  species = "human")
```

artmsSILACtoLong	<i>Convert the SILAC evidence file to MSstats format</i>
------------------	--

Description

Converting the evidence file from a SILAC search to a format compatible with MSstats. It basically modifies the Raw.files adding the Heavy and Light label

Usage

```
artmsSILACtoLong(evidence_file, output = NULL, verbose = TRUE)
```

Arguments

evidence_file (char) Text filepath to the evidence file
output (char) Text filepath of the output name. If NULL it does not write the output
verbose (logical) TRUE (default) shows function messages

Value

(data.frame) with SILAC data processed for MSstats (and output file)

Examples

```
evidence2silac <- artmsSILACtoLong(evidence_file = "silac.evidence.txt",
  output = "silac-evidence.txt")
```

artmsSpectralCounts	<i>Outputs the spectral counts from the MaxQuant evidence file.</i>
---------------------	---

Description

Outputs the spectral counts from the MaxQuant evidence file.

Usage

```
artmsSpectralCounts(
  evidence_file,
  keys_file,
  output_file = NULL,
  verbose = TRUE
)
```

Arguments

evidence_file (char) Maxquant evidence file or data object
 keys_file (char) Keys file with the experimental design or data object
 output_file (char) Output file name (add .txt extension). If NULL (default) it returns a data.frame object
 verbose (logical) TRUE (default) shows function messages

Value

A txt file with biological replicates, protein id, and spectral count columns

Examples

```
summary_spectral_counts <- artmsSpectralCounts(
  evidence_file = artms_data_ph_evidence,
  keys_file = artms_data_ph_keys)
```

artmsVolcanoPlot	<i>Volcano plot (log2fc / pvalues)</i>
------------------	--

Description

It generates a scatter-plot used to quickly identify changes

Usage

```
artmsVolcanoPlot(
  mss_results,
  output_name = "volcano_plot.pdf",
  lfc_upper = 1,
  lfc_lower = -1,
  whatPvalue = "adj.pvalue",
  FDR = 0.05,
  PDF = TRUE,
  decimal_threshold = 16,
  verbose = TRUE
)
```

Arguments

mss_results (data.frame or file) Selected MSstats results
 output_name (char) Name for the output file (don't forget the .pdf extension)
 lfc_upper (numeric) log2fc upper threshold (positive value)
 lfc_lower (numeric) log2fc lower threshold (negative value)
 whatPvalue (char) pvalue or adj.pvalue (default)
 FDR (numeric) False Discovery Rate threshold
 PDF (logical) Option to generate pdf format. Default: T
 decimal_threshold (numeric) Decimal threshold for the pvalue. Default: 16 (10⁻¹⁶)
 verbose (logical) TRUE (default) shows function messages

Value

(pdf) of a volcano plot

Examples

```
artmsVolcanoPlot(mss_results = artms_data_ph_msstats_results,  
                 whatPvalue = "pvalue",  
                 PDF = FALSE)
```

artmsWriteConfigYamlFile

Write out a template file of the artMS configuration file (yaml)

Description

Creates a template file of the artMS configuration file, which is required to run `artmsQuantification`. Check `?artms_config` and the vignettes to find out more about the details of the structure of the file and how to fill it up

Usage

```
artmsWriteConfigYamlFile(  
  config_file_name = "artms_config_file.yaml",  
  verbose = TRUE  
)
```

Arguments

`config_file_name` (char) The name for the configuration file. It must have a `.yaml` extension. If `NULL`, it returns the config as a yaml object

`verbose` (logical) `TRUE` (default) shows function messages

Value

A file (or yaml data object) of the artMS configuration file

Examples

```
config_empty <- artmsWriteConfigYamlFile(config_file_name = NULL)
```

artms_config	<i>artMS configuration template</i>
--------------	-------------------------------------

Description

The configuration file in yaml format contains the configuration details required to run `artmsQuantification()`, which includes quality control functions

Usage

artms_config

Format

The configuration (yaml) file contains the following sections:

- files**
 - evidence : /path/to/the/evidence.txt
 - keys : /path/to/the/keys.txt
 - contrasts : /path/to/the/contrast.txt
 - summary : /path/to/the/summary.txt
 - output : /path/to/the/output/results/results.txt
- qc**
 - basic: 1 # 1 = yes; 0 = no
 - extended: 1 # 1 = yes; 0 = no
 - extendedSummary: 0 # 1 = yes; 0 = no
- data**
 - enabled : 1 # 1 = yes; 0 = no
 - fractions:
 - enabled : 0 # 1 for protein fractionation
 - silac:
 - enabled : 0 # 1 for SILAC experiments
 - filters:
 - enabled : 1
 - contaminants : 1
 - protein_groups : remove #remove, keep
 - modifications : ab # PH, UB, AB, APMS
 - sample_plots : 1 # correlation plots
- msstats**
 - enabled : 1
 - msstats_input : # blank if not previous msstats input file is available
 - profilePlots : none # before, after, before-after, none
 - normalization_method : equalizeMedians # globalStandards (include a reference protein(s)), equalizeMedians, quantile, 0
 - normalization_reference : #should be a value in the Protein column
 - summaryMethod : TMP # "TMP"(default) means Tukey's median polish, which is robust estimation method. "linear" uses linear mixed model. "logOfSum" conducts log2 (sum of intensities) per run.
 - censoredInt : NA # 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.

- cutoffCensored : minFeature # Cutoff value for censoring. only with censoredInt='NA' or '0'. Default is 'minFeature', which uses minimum value for each feature.'minFeatureNRun' uses the smallest between minimum value of corresponding feature and minimum value of corresponding run. 'minRun' uses minimum value for each run.
- MBimpute : 1 # only for summaryMethod="TMP" and censoredInt='NA' or '0'. TRUE (default) imputes 'NA' or '0' (depending on censoredInt option) by Accelerated failure model. FALSE uses the values assigned by cutoffCensored.
- feature_subset: all # allhighQuality : highQuality seems to be buggy right now

output_extras • output_extras :

- enabled : 1 # if 0, it wont do anything in this section
- annotate :
 - enabled: 1 # 1|0 whether to annotate the proteins in the results or not
- species : HUMAN # Supported species: HUMAN, MOUSE, ANOPHELES, ARABIDOPSIS, BOVINE, WORM, CANINE, FLY, ZEBRAFISH, ECOLI_STRAIN_K12, ECOLI_STRAIN_SAKAI, CHICKEN, RHESUS, MALARIA, CHIMP, RAT, YEAST, PIG, XENOPUS
- plots:
 - volcano: 1
 - heatmap: 1
 - LFC : -1.5 1.5 # Range of minimal log2fc
 - FDR : 0.05
 - heatmap_cluster_cols : 0
 - heatmap_display : log2FC # log2FC or pvalue

artms_data_corum_mito_database

CORUM Protein Complexes database use for complex enrichment analysis

Description

The list of protein complexes has been enriched with mitochondria proteins from mouse, as described in this paper:

2018 - Ruchi Masand, Esther Paulo, Dongmei Wu , Yangmeng Wang, Danielle L. Swaney, David Jimenez-Morales, Nevan J. Krogan, and Biao Wang Proteome Imbalance of Mitochondrial Electron Transport Chain in Brown Adipocytes Leads to Metabolic Benefits. Cell Metab. 2018 Mar 06; 27(3):616-629.e4

Usage

artms_data_corum_mito_database

Format

Tab delimited file.

To find out more about the format and columns available at CORUM, please visit this [link](#)

Details

LAST CORUM DOWNLOAD DATE: 2017-08-01

artms_data_pathogen_LPN

*LPN PATHOGEN: Legionella pneumophila subsp. pneumophila
(strain Philadelphia 1 / ATCC 33152 / DSM 7513) UNIPROT IDS*

Description

LPN PATHOGEN: Legionella pneumophila subsp. pneumophila (strain Philadelphia 1 / ATCC 33152 / DSM 7513) UNIPROT IDS

Usage

artms_data_pathogen_LPN

Format

A data.frame of Entry IDs

artms_data_pathogen_TB

*TB PATHOGEN: Mycobacterium tuberculosis (strain ATCC 35801 /
TMC 107 / Erdman) UNIPROTS IDS*

Description

TB PATHOGEN: Mycobacterium tuberculosis (strain ATCC 35801 / TMC 107 / Erdman) UNIPROTS IDS

Usage

artms_data_pathogen_TB

Format

A data.frame of Entry IDs

artms_data_ph_evidence

Evidence file example

Description

Evidence file from a PH experiment consisting of two head and neck cancer cell lines ("Conditions" "Ca133" and "HSC6").

Unfortunately, the number of lines was reduced to 1/8 due to bioconductor limitations on data size, which means that this data is not very representative of a real evidence file. However, both the full evidence.txt and keys.txt file are available at: <http://kroganlab.ucsf.edu/artms/ph/evidence.txt>
<http://kroganlab.ucsf.edu/artms/ph/keys.txt>

Usage

artms_data_ph_evidence

Format

A data frame with all the columns available in an evidence file generated with MaxQuant version 1.6.2.3

artms_data_ph_keys

Keys File Example

Description

the artMS keys file provides the details of the experimental design for any given proteomics experiment.

This particular example belongs to a PH experiment consisting of two head and neck cancer cell lines ("Conditions" "Ca133" and "HSC6"), with 2 biological replicates each (in this reduced version)

Usage

artms_data_ph_keys

Format

Tab delimited file with the following columns:

Raw.file Raw file processed. Each one should be a unique biological (or technical) replicate

IsotopeLabelType Type of labeling. L is used for label free experiments

Condition Label for conditions. VERY IMPORTANT: Only alpha-numeric characters and underscore (_) are allowed

BioReplicate Label for the Biological replicates. VERY IMPORTANT: Use the same labeling for bioreplicate as the Condition, but adding a dash (-) corresponding to the number of biological replicate. For example, for Condition "Ca1", use Ca1-1, Ca1-2, Ca1-3, etc for the bioreplicates

Run The MS run number

`artms_data_ph_msstats_results`*MSstats results file example*

Description

Relative quantification results obtained running MSstats on a PH datasets (global analysis). Changes in protein phosphorylation were quantified between two conditions

Usage`artms_data_ph_msstats_results`**Format**

A data frame resulting from running the latest version of MSstats

`artms_data_randomDF` *Random data set*

Description

Dataset randomly generated for testing purposes

Usage`artms_data_randomDF`**Format**

A data frame with 100 rows and 10 variables:

Dataset generated using this code

```
data.frame(replicate(10, sample(0:1, 100, rep=TRUE)))
```

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