

Package ‘parsemsf’

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Title Parse Thermo MSF Files and Estimate Protein Abundances

Version 0.1.0

Description Provides functions for parsing Thermo MSF files produced by Proteome Discoverer 1.4.x (see <<https://thermofisher.com>> for more information). This package makes it easy to view individual peptide information, including peak areas, and to map peptides to locations within the parent protein sequence. This package also estimates protein abundances from peak areas and across multiple technical replicates. The author of this package is not affiliated with ThermoFisher Scientific in any way.

URL <https://github.com/benjaminjack/parsemsf/>

Depends R (>= 3.2.4)

License GPL-2

Encoding UTF-8

LazyData true

RoxygenNote 5.0.1

Imports dplyr (>= 0.5.0), lazyeval, RSQLite (>= 1.0.0), stats, stringr (>= 1.1.0), tidyr (>= 0.6.0)

Suggests testthat (>= 1.0.2), knitr, rmarkdown, ggplot2

VignetteBuilder knitr

BugReports <https://github.com/benjaminjack/parsemsf/issues>

NeedsCompilation no

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make_area_table	<i>Make a table of peptide areas</i>
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Description

Areas under each peptide peak that can be used downstream for quantitation. See [quantitate](#) for protein quantitation.

Usage

```
make_area_table(msf_file, min_conf = "High",
  prot_regex = "^>([a-zA-Z0-9._]+)\b", collapse = TRUE)
```

Arguments

msf_file	A file path to a Thermo MSF file.
min_conf	"High", "Medium", or "Low". The minimum peptide confidence level to retrieve from MSF file.
prot_regex	Regular expression where the first group matches a protein name or ID from the protein description. Regex must contain ONE group. The protein description is typically generated from a fasta reference file that was used for the database search.
collapse	If TRUE, peptides that match to multiple protein sequences are collapsed into a single row with multiple protein descriptions and protein IDs in the Proteins and ProteinID columns separated by semi-colons (;).

Value

A data frame containing peptide areas for peptides at or above the minimum confidence level.

peptide_id	a unique peptide ID
spectrum_id	a unique spectrum ID
protein_desc	protein description from reference database used to assign peptides to protein groups, parsed according to prot_regex
sequence	amino acid sequence (does not show post-translational modifications)
area	area under peptide peak
mass	peptide mass
m_z	mass-to-charge ratio
charge	peptide charge
intensity	peak intensity; useful if no area is available
first_scan	first scan in which peptide appears

Examples

```
make_area_table(parsemsf_example("test_db.msf"))
```

make_pep_table	<i>Make a data frame of peptides</i>
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Description

Extracts amino acid sequences (without post-translational modifications), assigned protein groups, and quality scores.

Usage

```
make_pep_table(msf_file, min_conf = "High",
  prot_regex = "^>([a-zA-Z0-9._]+)\\b", collapse = TRUE)
```

Arguments

msf_file	A file path to a Thermo MSF file.
min_conf	"High", "Medium", or "Low". The minimum peptide confidence level to retrieve from MSF file.
prot_regex	Regular expression where the first group matches a protein name or ID from the protein description. Regex must contain ONE group. The protein description is typically generated from a fasta reference file that was used for the database search.
collapse	If TRUE, peptides that match to multiple protein sequences are collapsed into a single row with multiple protein descriptions and protein IDs in the Proteins and ProteinID columns separated by semi-colons (";").

Value

A data frame of all peptides above the confidence cut-off from a thermo MSF file.

peptide_id	a unique peptide ID
spectrum_id	a unique spectrum ID
protein_id	unique protein group ID to which this peptide maps
protein_desc	protein description from reference database used to assign peptides to protein groups, parsed according to prot_regex
sequence	amino acid sequence (does not show post-translational modifications)
pep_score	PEP score
q_value	Q-value score

Examples

```
# Read from a path

make_pep_table(parsemsf_example("test_db.msf"))

# Retrieve full protein description

make_pep_table(parsemsf_example("test_db.msf"), prot_regex = "")

# ...which is also equivalent to...

make_pep_table(parsemsf_example("test_db.msf"), prot_regex = "^(.+)$")
```

map_peptides

Map peptides to their locations within a protein

Description

Takes a Thermo MSF file and finds the location of each peptide within its corresponding protein sequence. In cases where a single peptide maps to multiple locations within a protein sequence, only the first location is reported.

Usage

```
map_peptides(msf_file, min_conf = "High", prot_regex = "")
```

Arguments

msf_file	A file path to a Thermo MSF file.
min_conf	"High", "Medium", or "Low". The minimum peptide confidence level to retrieve from MSF file.
prot_regex	Regular expression where the first group matches a protein name or ID from the protein description. Regex must contain ONE group. The protein description is typically generated from a fasta reference file that was used for the database search.

Value

A dataframe containing start and stop positions (relative to the parent protein sequence) for each peptide in the database.

peptide_id	a unique peptide ID
spectrum_id	a unique spectrum ID
protein_id	unique protein group ID to which this peptide maps
protein_desc	protein description from reference database used to assign peptides to protein groups, parsed according to prot_regex

peptide_sequence	amino acid sequence (does not show post-translational modifications)
pep_score	PEP score
q_value	Q-value score
protein_sequence	parent protein sequence
start	start position of peptide within protein sequence
end	end position of peptide within protein sequence

Examples

```
map_peptides(parsemsf_example("test_db.msf"))
```

parsemsf	<i>parsemsf: Parse Thermo MSF files and estimate protein abundances.</i>
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Description

parsemsf: Parse Thermo MSF files and estimate protein abundances.

quantitate	<i>Combine technical replicates and quantitate proteins</i>
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Description

Takes a list of thermo MSF files, parses and combines them into a single data frame, and computes areas for each protein group based on the top 3 method of quantitation.

Usage

```
quantitate(reps, normalize = T, match_peps = T, relabel = c())
```

Arguments

reps	Vector. List of thermo MSF file names
normalize	Boolean. Should we normalize peptide areas for technical replicate to the total areas in a given replicate?
match_peps	Boolean. Should we quantitate only on matching peptides across technical replicates?
relabel	Named vector for relabeling protein groups. Names correspond to a pattern or string to match (i.e. the name or ID of a protein group), and values correspond to the new value (i.e. new protein group name).

Value

A data frame containing area information for all proteins.

protein_desc	protein description
area_mean	average peptide area
area_sd	peptide area standard deviation
peps_per_rep	Number of peptides per technical replicate used to calculate area_mean and area_sd. This is typically 3 peptides, but may be less.

Examples

```
quantitate(c(parsemsf_example("test_db.ms"),
             parsemsf_example("test_db2.ms")),
           relabel = c("NP_12345.1" = "NP_1000.1"))
```

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