Package 'MSstatsBig'

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Type Package

Title MSstats Preprocessing for Larger than Memory Data

Version 1.7.0

Description MSstats package provide tools for preprocessing, summarization and differential analysis of mass spectrometry (MS) proteomics data. Recently, some MS protocols enable acquisition of data sets that result in larger than memory quantitative data. MSstats functions are not able to process such data. MSstatsBig package provides additional converter functions that enable processing larger than memory data sets.

License Artistic-2.0

Encoding UTF-8

utils

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Imports arrow, DBI, dplyr, MSstats, MSstatsConvert, readr, sparklyr,

Suggests knitr, rmarkdown

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```
bigFragPipetoMSstatsFormat
```

Convert out-of-memory FragPipe files to MSstats format.

Description

Convert out-of-memory FragPipe files to MSstats format.

Usage

```
bigFragPipetoMSstatsFormat(
    input_file,
    output_file_name,
    backend,
    max_feature_count = 20,
    filter_unique_peptides = FALSE,
    aggregate_psms = FALSE,
    filter_few_obs = FALSE,
    remove_annotation = FALSE,
    connection = NULL
)
```

<pre>input_file</pre>	name of the input text file in 10-column MSstats format.	
output_file_name	ne	
	name of an output file which will be saved after pre-processing	
backend	"arrow" or "sparklyr". Option "sparklyr" requires a spark installation and con- nection to spark instance provided in the 'connection' parameter.	
<pre>max_feature_com</pre>	unt	
	maximum number of features per protein. Features will be selected based on highest average intensity.	
filter_unique_peptides		
	If TRUE, shared peptides will be removed. Please refer to the 'Details' section for additional information.	
aggregate_psms	If TRUE, multiple measurements per PSM in a Run will be aggregated (by tak- ing maximum value). Please refer to the 'Details' section for additional infor- mation.	
filter_few_obs	If TRUE, feature with less than 3 observations across runs will be removed. Please refer to the 'Details' section for additional information.	
remove_annotation		
	If TRUE, columns BioReplicate and Condition will be removed to reduce output file size. These will need to be added manually later before using dataProcess function. Only applicable to sparklyr backend.	
connection	Connection to a spark instance created with the 'spark_connect' function from 'sparklyr' package.	

bigSpectronauttoMSstatsFormat

Value

either arrow object or sparklyr table that can be optionally collected into memory by using dplyr::collect function.

Examples

```
converted_data <- bigFragPipetoMSstatsFormat(
   system.file("extdata", "fgexample.csv", package = "MSstatsBig"),
   "output_file.csv",
   backend = "arrow")
converted_data <- dplyr::collect(converted_data)
head(converted_data)</pre>
```

```
bigSpectronauttoMSstatsFormat
```

Convert out-of-memory Spectronaut files to MSstats format.

Description

Convert out-of-memory Spectronaut files to MSstats format.

Usage

```
bigSpectronauttoMSstatsFormat(
    input_file,
    output_file_name,
    backend,
    filter_by_excluded = FALSE,
    filter_by_identified = FALSE,
    filter_by_qvalue = TRUE,
    qvalue_cutoff = 0.01,
    max_feature_count = 20,
    filter_unique_peptides = FALSE,
    aggregate_psms = FALSE,
    filter_few_obs = FALSE,
    remove_annotation = FALSE,
    connection = NULL
)
```

input_file	name of the input text file in 10-column MSstats format.	
output_file_name		
	name of an output file which will be saved after pre-processing	
backend	"arrow" or "sparklyr". Option "sparklyr" requires a spark installation and con- nection to spark instance provided in the 'connection' parameter.	
filter_by_excluded		
	if TRUE, will filter by the 'F.ExcludedFromQuantification' column.	
filter_by_ident	ified	
	if TRUE, will filter by the 'EG.Identified' column.	

filter_by_qvalue		
	if TRUE, will filter by EG.Qvalue and PG.Qvalue columns.	
qvalue_cutoff	cutoff which will be used for q-value filtering.	
max_feature_count		
	maximum number of features per protein. Features will be selected based on highest average intensity.	
filter_unique_peptides		
	If TRUE, shared peptides will be removed. Please refer to the 'Details' section for additional information.	
aggregate_psms	If TRUE, multiple measurements per PSM in a Run will be aggregated (by tak- ing maximum value). Please refer to the 'Details' section for additional infor- mation.	
filter_few_obs	If TRUE, feature with less than 3 observations across runs will be removed. Please refer to the 'Details' section for additional information.	
remove_annotation		
	If TRUE, columns BioReplicate and Condition will be removed to reduce output file size. These will need to be added manually later before using dataProcess function. Only applicable to sparklyr backend.	
connection	Connection to a spark instance created with the 'spark_connect' function from 'sparklyr' package.	

Value

either arrow object or sparklyr table that can be optionally collected into memory by using dplyr::collect function.

Examples

```
converted_data <- bigSpectronauttoMSstatsFormat(
   system.file("extdata", "spectronaut_input.csv", package = "MSstatsBig"),
   "output_file.csv",
   backend="arrow")
converted_data <- dplyr::collect(converted_data)
head(converted_data)</pre>
```

MSstatsAddAnnotationBig

Merge annotation to output of MSstatsPreprocessBig

Description

Merge annotation to output of MSstatsPreprocessBig

Usage

MSstatsAddAnnotationBig(input, annotation)

input	output of MSstatsPreprocessBig
annotation	run annotation

MSstatsPreprocessBig

Value

table of 'input' and 'annotation' merged by Run column.

Examples

```
converted_data <- bigFragPipetoMSstatsFormat(
  system.file("extdata", "fgexample.csv", package = "MSstatsBig"),
  "output_file.csv",
  backend = "arrow")
converted_data <- dplyr::collect(converted_data)
head(converted_data)
# Change annotation as an example:
converted_data$Condition <- NULL
converted_data$BioReplicate <- NULL
annot <- data.frame(Run = unique(converted_data[["Run"]]))
annot$BioReplicate <- rep(1:53, times = 2)
annot$Condition <- rep(1:2, each = 53)
head(MSstatsAddAnnotationBig(converted_data, annot))
```

MSstatsPreprocessBig General converter for larger-than-memory csv files in MSstats format 10-column format

Description

General converter for larger-than-memory csv files in MSstats format 10-column format

Usage

```
MSstatsPreprocessBig(
    input_file,
    output_file_name,
    backend,
    max_feature_count = 20,
    filter_unique_peptides = FALSE,
    aggregate_psms = FALSE,
    filter_few_obs = FALSE,
    remove_annotation = FALSE,
    connection = NULL
)
```

input_file	name of the input text file in 10-column MSstats format.	
output_file_name		
	name of an output file which will be saved after pre-processing	
backend	"arrow" or "sparklyr". Option "sparklyr" requires a spark installation and connection to spark instance provided in the 'connection' parameter.	
max_feature_count		
	maximum number of features per protein. Features will be selected based on highest average intensity.	

filter_unique_peptides		
		If TRUE, shared peptides will be removed. Please refer to the 'Details' section for additional information.
	aggregate_psms	If TRUE, multiple measurements per PSM in a Run will be aggregated (by tak- ing maximum value). Please refer to the 'Details' section for additional infor- mation.
	filter_few_obs	If TRUE, feature with less than 3 observations across runs will be removed. Please refer to the 'Details' section for additional information.
remove_annotation		
		If TRUE, columns BioReplicate and Condition will be removed to reduce output file size. These will need to be added manually later before using dataProcess function. Only applicable to sparklyr backend.
	connection	Connection to a spark instance created with the 'spark_connect' function from 'sparklyr' package.

Details

Filtering and aggregation may be very time consuming and the ability to perform them in a given R session depends on available memory, settings of external packages, etc. Hence, all value of related parameters ('filter_unique_peptides', 'aggregate_psms', 'filter_few_obs') are set to FALSE by default and only feature selection is performed, which saves both computation time and memory. Appropriately configured spark backend provides the most consistent way to perform these operations.

Value

either arrow object or sparklyr table that can be optionally collected into memory by using dplyr::collect function.

Examples

```
converted_data <- bigFragPipetoMSstatsFormat(
   system.file("extdata", "fgexample.csv", package = "MSstatsBig"),
   "tencol_format.csv",
   backend="arrow")
procd <- MSstatsPreprocessBig("tencol_format.csv", "proc_out.csv", backend = "arrow")
head(dplyr::collect(procd))</pre>
```

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