Package 'MALDIquantForeign'

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     writing (tab, csv, mMass MSD, mzML, imzML) different file
     formats of mass spectrometry data into/from 'MALDIquant'
     objects.
License GPL (>= 3)
URL https://strimmerlab.github.io/software/maldiquant/,
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MALDIquantForeign-package

Import/Export routines for 'MALDIquant'

Description

This package reads and writes different file formats of mass spectrometry data into/from 'MALDIquant' objects.

Details

Package: MALDIquantForeign

License: GPL (>= 3)

URL: https://strimmerlab.github.io/software/maldiquant/

Author(s)

Sebastian Gibb <mail@sebastiangibb.de>

References

https://strimmerlab.github.io/software/maldiquant/

See Also

Useful links:

- https://strimmerlab.github.io/software/maldiquant/
- https://github.com/sgibb/MALDIquantForeign/
- Report bugs at https://github.com/sgibb/MALDIquantForeign/issues/

```
export, AbstractMassObject-method Export\ files
```

Description

This function provides a general interface to export AbstractMassObject-class objects (e.g. MassSpectrum-class, MassPeaks-class) into different file formats.

Usage

```
## S4 method for signature 'AbstractMassObject'
export(x, file, type="auto", force=FALSE, ...)
## S4 method for signature 'list'
export(x, path, type, force=FALSE, ...)
```

Arguments

X	a AbstractMassObject-class object or a list of AbstractMassObject-class objects.
file	character, file name.
type	character, file format. If type is set to "auto" the file extension is used.
force	logical, If TRUE the file would be overwritten or path would be created.
path	character, path to directory in which the list of AbstractMassObject-class would be exported.
	arguments to be passed to specific export functions.

Details

Specific export functions:

```
tab exportTab
csv exportCsv
imzML exportImzMl
msd exportMsd
mzML exportMzMl
```

Author(s)

Sebastian Gibb

References

```
https://strimmerlab.github.io/software/maldiquant/
```

See Also

```
MassPeaks-class, MassSpectrum-class
```

Examples

```
exportImzMl, MassSpectrum-method 
  Export \ to \ imzML \ files
```

Description

This function exports MassSpectrum-class objects into imzML files.

Usage

```
## S4 method for signature 'MassSpectrum'
exportImzMl(x, file, force=FALSE, processed=TRUE,
coordinates=NULL, pixelSize=c(100, 100), ...)
## S4 method for signature 'list'
exportImzMl(x, path, force=FALSE, processed=TRUE,
coordinates=NULL, pixelSize=c(100, 100), ...)
```

Arguments

X	a ${\tt AbstractMassObject-class}$ object or a list of ${\tt AbstractMassObject-class}$ objects.
file	character, file name.
force	logical, If TRUE the file would be overwritten or path would be created.
processed	logical, If TRUE (default) the spectra will be saved in processed mode (means mass and intensity is stored for each spectra separately in contrast to continuous mode where the mass is stored only for one spectrum).
coordinates	matrix, 2 column matrix that contains the x- and y-coordinates for the spectra.
pixelSize	numeric, a vector of length 2 that contains the x and y pixel size in micrometers (default: c(100, 100)).
path	character, path to directory in which the list of MassSpectrum-class would be exported. If path is a single filename all spectra will be exported to a single imzML file.
	arguments to be passed to internal functions.

Author(s)

Sebastian Gibb

References

```
https://strimmerlab.github.io/software/maldiquant/
```

Schramm T, Hester A, Klinkert I, Both J-P, Heeren RMA, Brunelle A, Laprevote O, Desbenoit N, Robbe M-F, Stoeckli M, Spengler B, Roempp A (2012)

imzML - A common data format for the flexible exchange and processing of mass spectrometry imaging data.

Journal of Proteomics 75 (16):5106-5110. doi:10.1016/j.jprot.2012.07.026

See Also

MassSpectrum-class

```
## End(Not run)
```

```
{\it export Msd}, {\it MassSpectrum-method} \\ {\it Export to MSD files}
```

Description

This function exports AbstractMassObject-class objects (e.g. MassSpectrum-class, MassPeaks-class) into mMass MSD files.

Usage

```
## S4 method for signature 'MassSpectrum'
exportMsd(x, file, force=FALSE, peaks, ...)
## S4 method for signature 'list'
exportMsd(x, path, force=FALSE, peaks, ...)
```

Arguments

X	a MassSpectrum-class object or a list of MassSpectrum-class objects.
file	character, file name.
force	logical, If TRUE the file would be overwritten or path would be created.
peaks	a MassPeaks-class object or a list of MassPeaks-class objects.
path	character, path to directory in which the list of ${\tt AbstractMassObject-class}$ would be exported.
	arguments to be passed to write.table.

Author(s)

Sebastian Gibb

References

```
https://strimmerlab.github.io/software/maldiquant/,
mMass homepage: http://mmass.org/
```

See Also

```
MassPeaks-class, MassSpectrum-class
```

Examples

 ${\tt exportMzMl,MassSpectrum-method}$

Export to mzML files

Description

This function exports MassSpectrum-class objects into mzML files.

Usage

```
## S4 method for signature 'MassSpectrum'
exportMzMl(x, file, force=FALSE, ...)
## S4 method for signature 'list'
exportMzMl(x, path, force=FALSE, ...)
```

Arguments

X	a AbstractMassObject-class object or a list of AbstractMassObject-class objects.
file	character, file name.
force	logical, If TRUE the file would be overwritten or path would be created.
path	character, path to directory in which the list of MassSpectrum-class would be exported. If path is a single filename all spectra will be exported to a single mzML file.
	arguments to be passed to internal functions.

Author(s)

Sebastian Gibb

References

```
https://strimmerlab.github.io/software/maldiquant/,
HUPO Proteomics Standards Inititative mzML 1.1.0 Specification: https://www.psidev.info/mzML
```

See Also

MassSpectrum-class

Examples

```
exportTab, AbstractMassObject-method  Export\ to\ text\ files
```

Description

This function exports AbstractMassObject-class objects (e.g. MassSpectrum-class, MassPeaks-class) into different text file formats.

Usage

```
## S4 method for signature 'AbstractMassObject'
exportTab(x, file, force=FALSE, ...)
## S4 method for signature 'list'
exportTab(x, path, force=FALSE, ...)
```

```
## S4 method for signature 'AbstractMassObject'
exportCsv(x, file, force=FALSE, ...)
## S4 method for signature 'list'
exportCsv(x, path, force=FALSE, ...)
```

Arguments

Х	a AbstractMassObject-class object or a list of AbstractMassObject-class objects.
file	character, file name.
force	logical, If TRUE the file would be overwritten or path would be created.
path	character, path to directory in which the list of AbstractMassObject-class would be exported.
	arguments to be passed to write.table.

Details

exportTab and exportCsv use write.table with different defaults ($sep="\t"$ in exportTab and sep="," in exportCsv).

Author(s)

Sebastian Gibb

References

```
https://strimmerlab.github.io/software/maldiquant/
```

See Also

MassPeaks-class, MassSpectrum-class, write.table

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```
## End(Not run)
```

import

Import files

Description

This function provides a general interface to import different file formats into MassSpectrum-class or MassPeaks-class objects.

Usage

```
import(
  path,
  type = "auto",
  pattern,
  excludePattern = NULL,
  removeEmptySpectra = TRUE,
  centroided = FALSE,
  massRange = c(0, Inf),
  minIntensity = 0,
  mc.cores = 1L,
  verbose = interactive(),
  ...
)
```

Arguments

path character, path to directory or file which should be read in.

type character, file format. If type is set to "auto" MALDIquant tries to detect the

correct file type automatically. It often depends on the file extension (if path is a directory the most represented file extension is used; pattern argument is

ignored).

pattern character, a regular expression to find files in a directory (see details). excludePattern character, a regular expression to exclude files in a directory (see details).

removeEmptySpectra

logical, should empty spectra excluded?

centroided logical, if centroided=FALSE (default) the data are treated as not centroided

and a list of MassSpectrum-class objects is returned. Use centroided=TRUE

to assume centroided data and get a list of MassPeaks-class objects.

massRange double, limits of mass import (left/minimal mass, right/maximal mass).

minIntensity double, minimal intensity to import.

mc.cores number of cores to use (default 1; only unix-based platforms are supported, see

MALDIquantForeign-parallel for details).

verbose logical, verbose output?

... arguments to be passed to specific import functions.

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Details

Specific import functions:

txt importTxt
tab importTab
csv importCsv
fid importBruk

fid importBrukerFlex
ciphergen importCiphergenXml

mzXML importMzXml
mzML importMzMl
imzML importImzMl
analyze importAnalyze
cdf importCdf
msd importMsd

path: In addition to the above mentioned file types the following (compressed) archives are supported, too: zip, tar, tar.gz, tar.bz2, tar.xz. The archives are uncompressed in a temporary directory. Afterwards the import function is called (with type="auto").

pattern: Sometimes unusual file extensions are used (e.g. "*.xml" for mzXML files). In this case a specific pattern could be defined to import files with an unusual file extension (e.g. pattern="^.*\.xml\$" to read all *.xml files in a directory; see regexp for details).

excludePattern: Sometimes some files should be excluded. E.g. to ignore additional aquired Bruker LIFT spectra (MALDI-TOF/TOF; which are not support, yet) you could use excludePattern="([[:digit:]\.]+)L

Value

a list of MassSpectrum-class or MassPeaks-class objects (depending on the centroided argument).

Author(s)

Sebastian Gibb

References

https://strimmerlab.github.io/software/maldiquant/

See Also

MassSpectrum-class, MassPeaks-class MALDIquantForeign-parallel

```
library("MALDIquant")
library("MALDIquantForeign")
## get example directory
```

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```
exampleDirectory <- system.file("exampledata", package="MALDIquantForeign")
## import mzXML files
s <- import(exampleDirectory, type="mzXML")
## import tab delimited file with different file extension (default: *.tab)
s <- import(exampleDirectory, type="tab", pattern="^.*\\.txt")
## import single mzML file
s <- import(file.path(exampleDirectory, "tiny1.mzML1.1.mzML"))
## import gzipped csv file
s <- import(file.path(exampleDirectory, "compressed", "csv1.csv.gz"))</pre>
```

importAnalyze

Import Analyze 7.5 files

Description

This function imports files in Analyze 7.5 file format into MassSpectrum-class or MassPeaks-class objects.

Usage

```
importAnalyze(path, ...)
```

Arguments

path character, path to directory or file which should be read in.
... arguments to be passed to import.

Value

a list of MassSpectrum-class or MassPeaks-class objects (depending on the centroided argument).

Author(s)

Sebastian Gibb

References

```
https://strimmerlab.github.io/software/maldiquant/
http://www.grahamwideman.com/gw/brain/analyze/formatdoc.htm, https://eeg.sourceforge.
net/ANALYZE75.pdf
```

See Also

MassSpectrum-class, MassPeaks-class

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importBrukerFlex

Import Bruker Daltonics *flex files

Description

This function imports files in Bruker Daltonics *flex-series file format into MassSpectrum-class or MassPeaks-class objects.

Usage

```
importBrukerFlex(path, ...)
```

Arguments

```
path character, path to directory or file which should be read in.
... arguments to be passed to readBrukerFlexFile.
```

Value

a list of MassSpectrum-class or MassPeaks-class objects (depending on the centroided argument).

Author(s)

Sebastian Gibb

References

```
https://strimmerlab.github.io/software/maldiquant/
```

See Also

MassSpectrum-class, MassPeaks-class, readBrukerFlexFile

```
library("MALDIquant")
library("MALDIquantForeign")

## get example directory
exampleDirectory <- system.file("exampledata", package="MALDIquantForeign")
s <- importBrukerFlex(exampleDirectory)</pre>
```

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importCdf

Import CDF files

Description

This function imports files in NetCDF file format into MassSpectrum-class or MassPeaks-class objects.

Please note that the *RNetCDF* is needed.

Usage

```
importCdf(path, ...)
```

Arguments

```
path character, path to directory or file which should be read in.
... arguments to be passed to import.
```

Value

a list of MassSpectrum-class or MassPeaks-class objects (depending on the centroided argument).

Author(s)

Sebastian Gibb

References

```
https://strimmerlab.github.io/software/maldiquant/
```

See Also

MassSpectrum-class, MassPeaks-class

```
library("MALDIquant")
library("MALDIquantForeign")

## get example directory
exampleDirectory <- system.file("exampledata", package="MALDIquantForeign")

## import
if (requireNamespace("RNetCDF", quietly=TRUE)) {
    s <- importCdf(exampleDirectory)
} else {
    message("You have to install the RNetCDF package to use importCdf.")</pre>
```

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}

Import Ciphergen XML files

Description

This function imports files in Ciphergen XML file format into MassSpectrum-class or MassPeaks-class objects.

Usage

```
importCiphergenXml(path, ...)
```

Arguments

path character, path to directory or file which should be read in.
... arguments to be passed to import.

Value

a list of MassSpectrum-class or MassPeaks-class objects (depending on the centroided argument).

Author(s)

Sebastian Gibb

References

https://strimmerlab.github.io/software/maldiquant/

See Also

MassSpectrum-class, MassPeaks-class

```
library("MALDIquant")
library("MALDIquantForeign")

## get example directory
exampleDirectory <- system.file("exampledata", package="MALDIquantForeign")

## import
s <- importCiphergenXml(exampleDirectory)</pre>
```

importImzMl

importImzMl

Import imzML files

Description

This function imports files in imzML file format into MassSpectrum-class or MassPeaks-class objects.

Usage

```
importImzMl(path, coordinates = NULL, ...)
```

Arguments

path character, path to directory or file which should be read in.

coordinates matrix, 2 column matrix that contains the x- and y-coordinates for spectra that

should be imported. Other spectra would be ignored.

... arguments to be passed to import.

Value

a list of MassSpectrum-class or MassPeaks-class objects (depending on the centroided argument).

Author(s)

Sebastian Gibb

References

```
https://strimmerlab.github.io/software/maldiquant/,
Definition of imzML format: https://ms-imaging.org/imzml/
```

See Also

MassSpectrum-class, MassPeaks-class

```
library("MALDIquant")
library("MALDIquantForeign")

## get example directory
exampleDirectory <- system.file("exampledata", package="MALDIquantForeign")

## import
s <- importImzMl(file.path(exampleDirectory, "tiny_continuous.imzML"))</pre>
```

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importMsd

Import MSD files

Description

This function imports files in mMass MSD file format into MassSpectrum-class or MassPeaks-class objects.

Usage

```
importMsd(path, ...)
```

Arguments

path character, path to directory or file which should be read in.
... arguments to be passed to import.

Value

a list of MassSpectrum-class or MassPeaks-class objects (depending on the centroided argument).

Author(s)

Sebastian Gibb

References

```
https://strimmerlab.github.io/software/maldiquant/,
mMass homepage: http://mmass.org/
```

See Also

MassSpectrum-class, MassPeaks-class

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Examples

```
library("MALDIquant")
library("MALDIquantForeign")

## get example directory
exampleDirectory <- system.file("exampledata", package="MALDIquantForeign")

## import
s <- importMsd(exampleDirectory)</pre>
```

importMzMl

Import mzML files

Description

This function imports files in mzML file format into MassSpectrum-class or MassPeaks-class objects.

Usage

```
importMzMl(path, ...)
```

Arguments

path character, path to directory or file which should be read in.
... arguments to be passed to import.

Value

a list of MassSpectrum-class or MassPeaks-class objects (depending on the centroided argument).

Author(s)

Sebastian Gibb

References

```
https://strimmerlab.github.io/software/maldiquant/,
Definition of mzML format: https://www.psidev.info/mzML
```

See Also

MassSpectrum-class, MassPeaks-class

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Examples

```
library("MALDIquant")
library("MALDIquantForeign")

## get example directory
exampleDirectory <- system.file("exampledata", package="MALDIquantForeign")

## import
s <- importMzMl(exampleDirectory)</pre>
```

importMzXml

Import mzXML files

Description

This function imports files in mzXML file format into MassSpectrum-class or MassPeaks-class objects.

Usage

```
importMzXml(path, ...)
```

Arguments

path character, path to directory or file which should be read in.
... arguments to be passed to readMzXmlFile.

Value

a list of MassSpectrum-class or MassPeaks-class objects (depending on the centroided argument).

Author(s)

Sebastian Gibb

References

```
https://strimmerlab.github.io/software/maldiquant/,
Definition of mzXML format: http://tools.proteomecenter.org/wiki/index.php?title=Formats:
mzXML
```

See Also

MassSpectrum-class, MassPeaks-class, readMzXmlFile

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Examples

```
library("MALDIquant")
library("MALDIquantForeign")

## get example directory
exampleDirectory <- system.file("exampledata", package="MALDIquantForeign")

## import
s <- importMzXml(exampleDirectory)</pre>
```

importTxt

Import text files

Description

This function imports different text file formats into MassSpectrum-class or MassPeaks-class objects.

Usage

```
importTxt(path, ...)
importTab(path, ...)
importCsv(path, ...)
```

Arguments

path character, path to directory or file which should be read in.
... arguments to be passed to read.table.

Details

importTab, importTxt and importCsv use read.table with different defaults.

Value

a list of MassSpectrum-class or MassPeaks-class objects (depending on the centroided argument).

Author(s)

Sebastian Gibb

References

https://strimmerlab.github.io/software/maldiquant/

See Also

MassSpectrum-class, MassPeaks-class, read.table

Examples

```
library("MALDIquant")
library("MALDIquantForeign")

## get example directory
exampleDirectory <- system.file("exampledata", package="MALDIquantForeign")

## import txt files
s <- importTxt(exampleDirectory)

## import csv files
s <- importCsv(exampleDirectory)</pre>
```

MALDIquantForeign-parallel

Parallel Support in Package MALDIquantForeign

Description

MALDIquantForeign-package offers multi-core support using mclapply and mcmapply. This approach is limited to unix-based platforms.

Details

Please note that not all import functions benfit from parallelisation. The current implementation is limited to run the parallelisation over different files. That's why only imports of multiple files could be run on multiple cores. E.g. a single mzML file containing 4 spectra would always be read on a single core. In contrast 4 mzML files each containing just one spectra could be read in using 4 cores.

The improvement in the runtime depends on the amount of data to read, the proportion of parsing/decoding of the data, the amount of memory and the speed of the hard disk.

Please note: It is possible that using parallelisation results in a worse runtime!

Author(s)

Sebastian Gibb <mail@sebastiangibb.de>

References

https://strimmerlab.github.io/software/maldiquant/

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See Also

```
MALDIquant-parallel, mclapply, mcmapply
```

Examples

```
## load packages
library("MALDIquant")
library("MALDIquantForeign")
exampleDirectory <- system.file("exampledata", package="MALDIquantForeign")</pre>
## run single-core import
print(system.time(
  s1 <- importMzMl(exampleDirectory, centroided=TRUE, verbose=FALSE)</pre>
))
if(.Platform$OS.type == "unix") {
  ## run multi-core import
  ## (because the example spectra are very small (just 5 data points) the
  ## multi-core solution is slower on most systems)
  print(system.time(
    s2 <- importMzMl(exampleDirectory, centroided=TRUE, mc.cores=2,</pre>
                     verbose=FALSE)
 ))
  stopifnot(all.equal(s1, s2))
```

supportedFileFormats Supported file formats

Description

This function prints all file formats supported by MALDIquantForeign-package.

Usage

```
supportedFileFormats()
```

Details

Import:

```
txt importTxt
tab importCsv
fid importBrukerFlex
ciphergen importCiphergenXml
mzXML importMzXml
mzML importMzMl
```

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imzML importImzMl
analyze importAnalyze
cdf importCdf
msd importMsd

Export:

tab exportTab
csv exportCsv
imzML exportImzMl
msd exportMsd
mzML exportMzMl

Value

a list with two named elements (import and export) containing a character vector of supported file types.

Author(s)

Sebastian Gibb

References

```
https://strimmerlab.github.io/software/maldiquant/
```

See Also

```
export, import
```

```
library("MALDIquantForeign")
supportedFileFormats()
```

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