

Package ‘readMzXmlData’

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Title Reads Mass Spectrometry Data in mzXML Format

Depends R (>= 4.2.0)

Imports base64enc, digest, XML

Description Functions for reading mass spectrometry data in mzXML format.

License GPL (>= 3)

URL <https://strimmerlab.github.io/software/maldisquant/>

<https://github.com/sgibb/readMzXmlData/>

BugReports <https://github.com/sgibb/readMzXmlData/issues/>

LazyLoad yes

RoxygenNote 7.2.3

NeedsCompilation no

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readMzXmlData-package *The readMzXmlData Package*

Description

The package reads mass spectrometry data in mzXML format.

Details

Main functions:

[readMzXmlFile](#): Reads mass spectrometry data in mzXML format.

[readMzXmlDir](#): Reads recursively mass spectrometry data in mzXML format in a specific directory.

[mqReadMzXml](#): Reads mass spectrometry data into MALDIquant.

Author(s)

Sebastian Gibb <mail@sebastiangibb.de>

References

See website: <https://strimmerlab.github.io/software/maldiquest/>

See Also

[readMzXmlDir](#), [readMzXmlFile](#)

readMzXmlDir

Reads recursively mass spectrometry data in mzXML format.

Description

Reads recursively all mass spectrometry data in mzXML format in a specified directory.

Usage

```
readMzXmlDir(  
  mzXmlDir,  
  removeCalibrationScans = TRUE,  
  removeMetaData = FALSE,  
  rewriteNames = TRUE,  
  fileExtension = "mzXML",  
  verbose = FALSE  
)
```

Arguments

mzXmlDir character, path to *directory* which should be read recursively.
 removeCalibrationScans logical, if TRUE all scans in directories called “[Cc]alibration” will be ignored.
 removeMetaData logical, to save memory metadata could be deleted.
 rewriteNames logical, if TRUE all list elements get an unique name from metadata otherwise file path is used.
 fileExtension character, file extension of mzXML formatted files. The directory is only searched for *fileExtension* files. In most cases it would be ““mzXML”” but sometimes you have to use ““xml””.
 verbose logical, verbose output?

Details

See [readMzXmlFile](#).

Value

A list of spectra.

- [[1]]\$spectrum\$mass: A vector of calculated mass.
- [[1]]\$spectrum\$intensity: A vector of intensity values.
- [[1]]\$metaData: A list of metaData depending on read spectrum.

Author(s)

Sebastian Gibb <mail@sebastiangibb.de>

See Also

[readMzXmlFile](#), [importMzXml](#)

Examples

```

## load library
library("readMzXmlData")

## get examples directory
exampleDirectory <- system.file("Examples", package="readMzXmlData")

## read example spectra
spec <- readMzXmlDir(exampleDirectory)

## plot spectra
plot(spec[[1]]$spectrum$mass, spec[[1]]$spectrum$intensity, type="n")

l <- length(spec)
legendStr <- character(l)

```

```

for (i in seq(along=spec)) {
  lines(spec[[i]]$spectrum$mass, spec[[i]]$spectrum$intensity, type="l",
        col=rainbow(l)[i])
  legendStr[i] <- basename(spec[[i]]$metaData$file)
}

## draw legend
legend(x="topright", legend=legendStr, col=rainbow(l), lwd=1)

```

readMzXmlFile*Reads mass spectrometry data in mzXML format.***Description**

Reads mass spectrometry data in mzXML format defined in <http://tools.proteomecenter.org/wiki/index.php?title=Formats:mzXML>

Usage

```
readMzXmlFile(mzXmlFile, removeMetaData = FALSE, verbose = FALSE)
```

Arguments

<code>mzXmlFile</code>	character, path to <i>mzXML</i> file which should be read.
<code>removeMetaData</code>	logical, to save memory metadata could be deleted.
<code>verbose</code>	logical, verbose output?

Value

A list of spectra and metadata.

- `spectrum$mass`: A vector of calculated mass.
- `spectrum$intensity`: A vector of intensity values.
- `metaData`: A list of metaData depending on read spectrum.

Author(s)

Sebastian Gibb <mail@sebastiangibb.de>

References

Definition of mzXML format: <http://tools.proteomecenter.org/wiki/index.php?title=Formats:mzXML>

See Also

[readMzXmlDir](#), [importMzXml](#)

Examples

```
## load library
library("readMzXmlData")

## get examples directory
exampleDirectory <- system.file("Examples", package="readMzXmlData")

## read example spectrum
spec <- readMzXmlFile(file.path(exampleDirectory, "A1-0_A1.mzXML"))

## print metaData
print(spec$metaData)

## plot spectrum
plot(spec$spectrum$mass, spec$spectrum$intensity, type="l")
```

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