Package ‘MsExperiment’

January 18, 2024

Title Infrastructure for Mass Spectrometry Experiments

Version 1.4.0

Description Infrastructure to store and manage all aspects related to a complete proteomics or metabolomics mass spectrometry (MS) experiment. The MsExperiment package provides light-weight and flexible containers for MS experiments building on the new MS infrastructure provided by the Spectra, QFeatures and related packages. Along with raw data representations, links to original data files and sample annotations, additional metadata or annotations can also be stored within the MsExperiment container. To guarantee maximum flexibility only minimal constraints are put on the type and content of the data within the containers.

Depends R (>= 4.2), ProtGenerics (>= 1.9.1),

Imports methods, S4Vectors, IRanges, Spectra, SummarizedExperiment, QFeatures

Suggests testthat, knitr (>= 1.1.0), roxygen2, BiocStyle (>= 2.5.19), rmarkdown, rpx, mzR, msdata

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LazyData no

VignetteBuilder knitr

BugReports https://github.com/RforMassSpectrometry/MsExperiment/issues

URL https://github.com/RforMassSpectrometry/MsExperiment

biocViews Infrastructure, Proteomics, MassSpectrometry, Metabolomics, ExperimentalDesign, DataImport

RoxygenNote 7.2.3

Roxygen list(markdown=TRUE)

Encoding UTF-8

Collate 'MsExperiment-functions.R' 'MsExperimentFiles.R'
'MsExperiment.R' 'existMsExperimentFiles.R'
experimentFiles

**Managing Mass Spectrometry Experiments**

**Description**

The MsExperiment class allows the storage and management of all aspects related to a complete proteomics or metabolomics mass spectrometry experiment. This includes experimental design (i.e. a table with samples), raw mass spectrometry data as spectra and chromatograms, quantitative features, and identification data or any other relevant data files.

For details, see https://rformassspectrometry.github.io/MsExperiment

This package is part of the RforMassSpectrometry initiative: https://www.rformassspectrometry.org/

**Usage**

```r
experimentFiles(object)
```

```r
experimentFiles(object) <- value
```

```r
sampleData(object)
```

```r
sampleData(object) <- value
```

```r
qdata(object)
```

```r
qdata(object) <- value
```
MsExperiment()

## S4 method for signature 'MsExperiment'
show(object)

## S4 method for signature 'MsExperiment'
length(x)

## S4 method for signature 'MsExperiment'
spectra(object)

## S4 replacement method for signature 'MsExperiment'
spectra(object) <- value

otherData(object)

otherData(object) <- value

linkSampleData(
    object,
    with = character(),
    sampleIndex = seq_len(nrow(sampleData(object))),
    withIndex = integer(),
    subsetBy = 1L
)

## S4 method for signature 'MsExperiment,ANY,ANY,ANY'
x[i, j, ..., drop = FALSE]

Arguments

object An instance of class MsExperiment.
value An object of the appropriate class for the slot to be populated.
x an MsExperiment.
with for linkSampleData: character(1) defining the data to which samples should be linked. See section Linking sample data to other experimental data for details.
sampleIndex for linkSampleData: integer with the indices of the samples in sampleData(object) that should be linked.
withIndex for linkSampleData: integer with the indices of the elements in with to which the samples (specified by sampleIndex) should be linked to.
subsetBy for linkSampleData: optional integer(1) defining the dimension on which the subsetting will occur on the linked data. Defaults to subsetBy = 1L thus subsetting will happen on the first dimension (rows or elements).
i for [: an integer, character or logical referring to the indices or names (rownames of sampleData) of the samples to subset.
experimentFiles

j for [: not supported.
...
optional additional parameters.
drop for [: ignored.

Value

See help of the individual functions.

Slots

experimentFiles An instance of class MsExperimentFiles or NULL.
spectra An instance of class Spectra or NULL.
qdata An instance of class QFeatures, SummarizedExperiment or NULL.
otherData A List to store any additional data objects.
sampleData A DataFrame documenting the experimental design.
sampleDataLinks A List with link definitions between samples and data elements. Should not be directly accessed or modified by the user.
metadata A list to store additional metadata.

General information

An experiment is typically composed of several items

- Description and information (covariates etc) of each sample from the experiment. These are stored in the sampleData slot as a DataFrame, each row describing a sample with columns containing all relevant information on that sample.
- Files to data or annotations. These are stored in the experimentFiles slot as an instance of class MsExperimentFiles.
- General metadata about the experiment, stored as a list in the metadata slot.
- Mass spectrometry data. Spectra and their metadata are stored as an Spectra() object in the spectra slot. Chromatographic data is not yet supported but will be stored as a Chromatograms() object in the chromatograms slot.
- Quantification data is stored as QFeatures or SummarizedExperiment objects in the qdata slot and can be accessed or replaced with the qdata or qdata<- functions, respectively.
- Any additional data, be it other spectra data, or proteomics identification data (i.e peptide-spectrum matches defined as PSM() objects) can be added as elements to the list stored in the otherData slot.

The length of a MsExperiment is defined by the number of samples (i.e. the number of rows of the object's sampleData). A MsExperiment with two samples will thus have a length of two, independently of the number of files or length of raw data in the object. This also defines the subsetting of the object using the [: function which will always subset by samples. See the section for filtering and subsetting below for more information.

MsExperiment objects can be created using the MsExperiment() function and by subsequently adding data and information to it (see examples below or the package vignette for more information), or using the readMsExperiment() function that creates a MsExperiment by importing directly MS spectra data from provided data files.
experimentFiles

Accessing data

Data from an MsExperiment object can be accessed with the dedicated accessor functions:

- `experimentFiles`, `experimentFiles<-`: gets or sets experiment files.
- `length`: get the length of the object which represents the number of samples available in the object’s sampleData.
- `metadata`, `metadata<-`: gets or sets the object’s metadata.
- `sampleData`, `sampleData<-`: gets or sets the object’s sample data (i.e. a DataFrame containing sample descriptions).
- `spectra`, `spectra<-`: gets or sets spectra data. spectra returns a `Spectra()` object, `spectra<-` takes a Spectra data as input and returns the updated MsExperiment.
- `qdata`, `qdata<-`: gets or sets the quantification data, which can be a QFeatures or SummarizedExperiment.
- `otherData`, `otherData<-`: gets or sets the addition data types, stored as a List in the object’s otherData slot.

Linking sample data to other experimental data

To start with, an MsExperiment is just a loose collection of files and data related to an experiment, no explicit links or associations are present between the samples and related data. Such links can however be created with the `linkSampleData()` function. This function can establish links between individual (or all) samples within the object’s sampleData to individual, or multiple, data elements or files, such as Spectra or raw data files.

The presence of such links enables a (consistent) subsetting of an MsExperiment by samples. Thus, once the link is defined, any subsetting by sample will also correctly subset the linked data. All other, not linked, data elements are always retained as in the original MsExperiment.

To be able to link different elements within an MsExperiment it is also required to identify them with a consistent naming scheme. The naming scheme of slots and data elements within follows an SQL-like scheme, in which the variable (element) is identified by the name of the database table, followed by a "." and the name of the database table column. For MsExperiment, the naming scheme is defined as "<slot name>.<element name>". A column called "sample_name" within the sampleData data frame can thus be addressed with "sampleData.sample_name", while spectra.msLevel would represent the spectra variable called msLevel within the Spectra stored in the spectra slot.

Links between sample data rows and any other data element are stored as integer matrices within the sampleDataLinks slot of the object (see also the vignette for examples and illustrations). Such links can be defined/added with the `linkSampleData` function which adds a relationship between rows in sampleData to elements in any other data within the MsExperiment that are specified with parameter with. `linkSampleData` supports two different ways to define the link:

- Parameter with defines the data to which the link should be established. To link samples to raw data files that would for example be available as a character in an element called "raw_files" within the object's experimentFiles, with = experimentFiles.raw_files would have to be used. Next it is required to specify which samples should be linked with which elements in with. This needs to be defined with the parameters sampleIndex and withIndex, both are expected to be integer vectors specifying which sample in sampleData should be linked to which element in with (see examples below or vignette for examples and details).
• As an alternative way, a link could be defined with an SQL-like syntax that relates a column in sampleData to a column/element in the data to which the link should be established. To link for example individual spectra to the corresponding samples with `"sampleData.raw_file = spectra.dataOrigin"` could be used assuming that sampleData contains a column named "raw_file" with the (full path) of the raw data file for each sample from which the spectra were imported. In this case both sampleIndex and withIndex can be omitted, but it is expected/required that the columns/elements from sampleData and the data element to which the link should be established contain matching values.

Note that linkSampleData will replace a previously existing link to the same data element.

**Subsetting and filtering**

• `[:]`: MsExperiment objects can be subset by samples with `[i]` where `i` is the index or a logical defining to which samples the data should be subset. Subsetting by sample will (correctly) subset all linked data to the respective samples. If multiple samples are linked to the same data element, subsetting might duplicate that data element. This duplication of n:m relationships between samples to elements does however not affect data consistency (see examples below for more information). Not linked data (slots) will be returned as they are. Subsetting in arbitrary order is supported. See the vignette for details and examples.

**Author(s)**

Laurent Gatto, Johannes Rainer

**Examples**

```r
## An empty MsExperiment object
msexp <- MsExperiment()
msexp

example(MsExperimentFiles)
experimentFiles(msexp) <- fls
msexp

## Linking samples to data elements

## Create a small experiment
library(S4Vectors)
mse <- MsExperiment()
sd <- DataFrame(sample_id = c("QC1", "QC2"),
                sample_name = c("QC Pool", "QC Pool"),
                injection_idx = c(1, 3))

sampleData(mse) <- sd

## define file names containing spectra data for the samples and
## add them, along with other arbitrary files to the experiment
fls <- dir(system.file("sciex", package = "msdata"), full.names = TRUE)

experimentFiles(mse) <- MsExperimentFiles(
  mzML_files = fls,
  annotations = "internal_standards.txt")
```
MsExperimentFiles

A class to store experiment files

## Link samples to data files: first sample to first file in "mzML_files",
## second sample to second file in "mzML_files"
mses <- linkSampleData(mse, with = "experimentFiles.mzML_files",
sampleIndex = c(1, 2), withIndex = c(1, 2))

## Link all samples to the one file in "annotations"
mses <- linkSampleData(mse, with = "experimentFiles.annotations",
sampleIndex = c(1, 2), withIndex = c(1, 1))
mses

## Import the spectra data and add it to the experiment
library(Spectra)
spectra(mse) <- Spectra(fls, backend = MsBackendMzR())

## Link each spectrum to the respective sample. We use the alternative
## link definition that does not require sampleIndex and withIndex but
## links elements based on matching values in the specified data elements.
## We need to add the full file name as an additional column to sampleData
## in order to allow matching this file names with the value in
## spectra(mse)$dataOrigin which contains the original file names from which
## the spectra were imported.
sampleData(mse)$raw_file <- normalizePath(fls)

## The links can be added using the short notation below
mses <- linkSampleData(mse, with = "sampleData.raw_file = spectra.dataOrigin")
mses

## With sampleData links present, any subsetting of the experiment by sample
## will ensure that all linked elements are subset accordingly
b <- mses[2]
b
sampleData(b)
experimentFiles(b)$mzML_files

## Subsetting with duplication of n:m sample to data relationships
##
## Both samples were assigned above to one "annotation" file in
## "experimentFiles":
experimentFiles(mses[1])[["annotations"]]
experimentFiles(mses[2])[["annotations"]]

## Subsetting will always keep the relationship between samples and linked
## data elements. Subsetting will however eventually duplicate data elements
## that are shared among samples. Thus, while in the original object the
## element "annotations" has a single entry, subsetting with [1:2] will
## result in an MsExperiment with duplicated entries in "annotations"
experimentFiles(mse)[["annotations"]]
experimentFiles(mse[1:2])["annotations"]

MsExperimentFiles

Description

The MsExperimentFiles class stores files that are part of a mass spectrometry experiment. The objects are created with the MsExperimentFiles() function.

The files encoded in a MsExperimentFiles instance don't need to exist on the current filesystem - sometimes, these might be created in anticipation of their creation. The existMsExperimentFiles() function can be used to verify which ones currently exist: it returns a list of logicals (formally an instance of IRanges::LogicalList) of lengths equal to the MsExperimentFiles used as input.

Usage

MsExperimentFiles(..., metadata = list())

## S4 method for signature 'MsExperimentFiles'
show(object)

existMsExperimentFiles(object)

Arguments

... Either a named list or a set of named vectors. All elements are coerced to characters.

metadata list() holding arbitrary R objects as annotations.

object The existMsExperimentFiles() function works with either an instance of MsExperimentFiles or MsExperiment.

Value

MsExperimentFiles returns an instance of MsExperimentFiles.

Author(s)

Laurent Gatto

Examples

fls <- MsExperimentFiles(mzmls = c("/path/to/f1.mzML", "/path/to/f2.mzML"),
                        mzids = "/another/path/to/id1.mzid",
                        fasta = "file.fas")

fls

## A new MsExperimentFiles containing mzML or mzid files
fls[1]
fls["mzids"]

## The actual file names
fls[[1]]
fls[[2]]
fls[["fasta"]]

fls
readMsExperiment

## None of the files used in this example actually exist
existMsExperimentFiles(fls)

---

**readMsExperiment**  
*Import MS spectra data of an experiment*

**Description**

Read/import MS spectra data of an experiment from the respective (raw) data files into an `MsExperiment()` object. Files provided with the `spectraFiles` parameter are imported as a `Spectra()` object and each file is automatically *linked* to rows (samples) of a `sampleData` data frame (if provided).

**Usage**

```r
readMsExperiment(spectraFiles = character(), sampleData = data.frame(), ...)
```

**Arguments**

- `spectraFiles` character with the (absolute) file names of the MS data files that should be imported as a `Spectra()` object.
- `sampleData` data.frame or DataFrame with the sample annotations. Each row is expected to contain annotations for one file (sample). The order of the data frame’s rows is expected to match the order of the provided files (with parameter `spectraFiles`).
- `...` additional parameters for the `Spectra()` call to import the data.

**Value**

`MsExperiment`.

**Author(s)**

Johannes Rainer

**Examples**

```r
## Define the files of the experiment to import
fls <- c(system.file("microtofq/MM14.mzML", package = "msdata"),
         system.file("microtofq/MM8.mzML", package = "msdata"))

## Define a data frame with some sample annotations
ann <- data.frame(
  injection_index = 1:2,
  sample_id = c("MM14", "MM8"))

## Import the data
library(MsExperiment)
mse <- readMsExperiment(spectraFiles = fls, ann)
mse
```
## Access the spectra data
spectra(mse)

## Access the sample annotations
sampleData(mse)

## Import the data reading all MS spectra directly into memory
mse <- readMsExperiment(spectraFiles = fls, ann,
  backend = Spectra::MsBackendMemory())
mse
Index

[,MsExperiment,ANY,ANY,ANY-method (experimentFiles), 2

existMsExperimentFiles (MsExperimentFiles), 7
experimentFiles, 2
experimentFiles<- (experimentFiles), 2

IRanges::LogicalList(), 8

length,MsExperiment-method (experimentFiles), 2
linkSampleData (experimentFiles), 2

MsExperiment (experimentFiles), 2
MsExperiment(), 9
MsExperiment-class (experimentFiles), 2
MsExperimentFiles, 7
MsExperimentFiles-class (MsExperimentFiles), 7

otherData (experimentFiles), 2
otherData<- (experimentFiles), 2

qdata (experimentFiles), 2
qdata<- (experimentFiles), 2

readMsExperiment, 9
readMsExperiment(), 4

sampleData (experimentFiles), 2
sampleData<- (experimentFiles), 2
show,MsExperiment-method (experimentFiles), 2
show,MsExperimentFiles-method (MsExperimentFiles), 7
spectra, 5, 9
spectra,MsExperiment-method (experimentFiles), 2
spectra<-,MsExperiment-method (experimentFiles), 2