

Package ‘MsBackendMsp’

April 3, 2023

Title Mass Spectrometry Data Backend for NIST msp Files

Version 1.2.0

Description Mass spectrometry (MS) data backend supporting import and handling of MS/MS spectra from NIST MSP Format (msp) files. Import of data from files with different MSP *flavours* is supported. Objects from this package add support for MSP files to Bioconductor's Spectra package. This package is thus not supposed to be used without the Spectra package that provides a complete infrastructure for MS data handling.

Depends R (>= 4.1.0), Spectra (>= 1.5.14)

Imports BiocParallel, S4Vectors, IRanges, MsCoreUtils, methods, stats

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

License Artistic-2.0

Encoding UTF-8

VignetteBuilder knitr

BugReports <https://github.com/RforMassSpectrometry/MsBackendMsp/issues>

URL <https://github.com/RforMassSpectrometry/MsBackendMsp>

biocViews Infrastructure, Proteomics, MassSpectrometry, Metabolomics, DataImport

Roxygen list(markdown=TRUE)

RoxygenNote 7.1.2

Collate 'hidden_aliases.R' 'MsBackendMsp.R' 'functions-msp.R'

git_url <https://git.bioconductor.org/packages/MsBackendMsp>

git_branch RELEASE_3_16

git_last_commit 3cdc114

git_last_commit_date 2022-11-01

Date/Publication 2023-04-03

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MsBackendMsp	<i>MS data backend for msp files</i>
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Description

The MsBackendMsp class supports import of MS/MS spectra data from files in NIST MSP file format. MsBackendMsp extends the `MsBackendDataFrame()` backend directly and supports thus the `applyProcessing()` function to make data manipulations persistent.

New objects are created with the MsBackendMsp function. The backendInitialize method has to be subsequently called to initialize the object and import MS/MS data from (one or more) msp files.

The MsBackendMsp backend provides an export method that allows to export the data from the Spectra object (parameter x) to a file in msp format. Parameters to this function are:

- x: the Spectra object that should be exported.
- file: character(1) with the desired file name.
- mapping: named character providing the mapping between spectra variables and MSP data fields. Defaults to mapping = spectraVariableMapping(MsBackendMsp()).
- allVariables: logical(1) whether all spectra variables in x should be exported or only those defined with mapping.
- exportName: logical(1) whether a NAME field should always be exported even if not provided in x.

See the package vignette for details and examples.

The spectraVariableMapping function allows to provide the mapping between spectra variable names (i.e. the names that will be used for the spectra variables in the `Spectra()` object) and the data field names of the MSP file. Parameter format allows to select pre-defined mappings. Currently supported mapping flavors are:

- format = "msp": default MSP field names. Should work with standard NIST MSP files or MSP files exported from MS-DIAL.
- format = "mona": MSP file format from MoNA including LipidBlast.

Usage

```
## S4 method for signature 'MsBackendMsp'
backendInitialize(
  object,
  file,
  mapping = spectraVariableMapping(object),
  ...,
  BPPARAM = bpparam()
)

MsBackendMsp()

## S4 method for signature 'MsBackendMsp'
spectraVariableMapping(object, format = c("msp", "mona"))

## S4 method for signature 'MsBackendMsp'
export(
  object,
  x,
  file = tempfile(),
  mapping = spectraVariableMapping(MsBackendMsp()),
  allVariables = TRUE,
  exportName = TRUE,
  ...
)
```

Arguments

object	Instance of MsBackendMsp class.
file	character with the (full) file name(s) of the msp file(s) from which MS/MS data should be imported or exported.
mapping	named character vector to rename MSP fields to spectra variables (see spectraVariableMapping()). This allows to correctly import also custom fields or data from files with different MSP <i>flavors</i> .
...	Currently ignored.
BPPARAM	Parameter object defining the parallel processing setup to import data in parallel. Defaults to BPPARAM = bpparam() . See bpparam() for more information.
format	For spectraVariableMapping: character(1) specifying for which MSP <i>flavour</i> the mapping should be returned. Currently supported are: format = "msp" (generic MSP format, for example for MS-DIAL MSP files) and format = "mona" (MSP files in MoNA flavour).
x	For export: a Spectra() object that should be exported to the specified MSP file.
allVariables	logical(1) whether all spectra variables in x should be exported or only those defined with mapping.
exportName	logical(1) whether a NAME field should always be exported even if not provided in x.

Value

MsBackendMsp and backendInitialize return an instance of a MsBackendMsp class. spectraVariableMapping a named character vector with the mapping between spectra variables and MSP data fields.

Author(s)

Steffen Neumann, Michael Witting, Laurent Gatto and Johannes Rainer

Examples

```
## Import spectra from a MSP file from LipidBlast
f <- system.file("extdata", "small-export-LipidBlast.msp",
  package = "MsBackendMsp")
be <- backendInitialize(MsBackendMsp(), f)
be

be$msLevel
be$intensity
be$mz

## precursor m/z are however all missing
be$precursorMz

## Default spectra variable mapping
spectraVariableMapping(MsBackendMsp())

## In fact, to read MSP files in "LipidBlast flavour" (same as MoNA) we
## should use a different spectra variable mapping
spectraVariableMapping(MsBackendMsp(), "mona")

## Importing the data with this will correctly retrieve data
be <- backendInitialize(MsBackendMsp(), f,
  mapping = spectraVariableMapping(MsBackendMsp(), "mona"))
be$precursorMz

## Other fields are also correctly mapped, but might need to be converted
## to e.g. numeric, such as "exactmass"
be$exactmass

be$exactmass <- as.numeric(be$exactmass)

be$adduct
be$formula

## Exporting Spectra objects in MSP format.

sps <- Spectra(be)
export(MsBackendMsp(), sps, file = stdout())
```

readMsp	<i>Reading MSP files</i> The readMsp function imports the data from a file in MGF format reading all specified fields and returning the data as a DataFrame() .
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Description

Reading MSP files

The readMsp function imports the data from a file in MGF format reading all specified fields and returning the data as a [DataFrame\(\)](#).

Usage

```
readMsp(f, msLevel = 2L, mapping = spectraVariableMapping(MsBackendMsp()), ...)
```

Arguments

f	character(1) with the path to an MSP file.
msLevel	numeric(1) with the MS level. Default is 2. This value will be reported as the spectra's MS level unless the source MSP file defines the MS level.
mapping	named character vector to rename MSP fields to spectra variables (see spectraVariableMapping()). This allows to correctly import also custom fields or data from files with different MSP <i>flavors</i> .
...	Additional parameters, currently ignored.

Value

A DataFrame with each row containing the data from one spectrum in the MSP file. m/z and intensity values are available in columns "mz" and "intensity" in a list representation.

Author(s)

Laurent Gatto, Steffen Neumann, Johannes Rainer

Examples

```
fls <- dir(system.file("extdata", package = "MsBackendMsp"),
  full.names = TRUE, pattern = "msp$")[1L]

readMsp(fls)
```

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