Package 'MGMS2'

October 12, 2022

Title 'MGMS2' for Polymicrobial Samples

Version 1.0.2

Author So Young Ryu [aut] (https://orcid.org/0000-0003-2347-7015), George Wendt [cre] (https://orcid.org/0000-0003-3608-9601)

Maintainer George Wendt < gwendt@unr.edu>

Description

A glycolipid mass spectrometry technology has the potential to accurately identify individual bacterial species from polymicrobial samples. To develop bacterial identification algorithms (e.g. machine learning) using this glycolipid technology, it is necessary to generate a large number of various in-silico polymicrobial mass spectra that are similar to real mass spectra. 'MGMS2' (Membrane Glycolipid Mass Spectrum Simulator) generates such in-silico mass spectra, considering errors in m/z (mass-to-charge ratio) and variances of intensity values, occasions of missing signature ions, and noise peaks. It estimates summary statistics of monomicrobial mass spectra for each strain or species and simulates polymicrobial glycolipid mass spectra using the summary statistics of monomicrobial mass spectra. References: Ryu, S.Y., Wendt, G.A., Chandler, C.E., Ernst, R.K. and Goodlett, D.R. (2019) <doi:10.1021/acs.analchem.9b03340> ``Modelbased Spectral Library Approach for Bacterial Identification via Membrane Glycolipids." Gibb, S. and Strim-

mer, K. (2012) <doi:10.1093/bioinformatics/bts447> ``MALDIquant: a versatile R package for the analysis of mass spectrometry data."

License GPL-3

Encoding UTF-8

LazyData true

Depends MALDIquant, MALDIquantForeign

RoxygenNote 7.0.2

Suggests testthat

NeedsCompilation no

Repository CRAN

Date/Publication 2020-04-24 04:10:02 UTC

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Description

This function characterizes peaks by species/strain in a simulated spectrum after taking the highest peak or merging peaks in each bin.

Usage

```
characterize_peak(spec, option = 1, bin.size = 1, min.mz = 1000, max.mz = 2200)
```

Arguments

spec	A data frame that contains m/z values of peaks, normalized intensities of peaks, species names, and strain names. Either an output of simulate_poly_spectra or one elements of a list output from simulate_many_poly_spectra.
option	An option on how to merge peaks. There are two options: 1) no merge, thus take the highest intensity peak in each bin after binning a spectrum by bin.size, or 2) take a sum of intensity within each bin after binning a spectrum by bin.size.
bin.size	An integer. A bin size. (1 by default)
min.mz	A real number. Minimum mass-to-charge ratio. (1000 by default)
max.mz	A real number. Maximum mass-to-charge ratio. (2200 by default)

Value

A data frame that contains m/z values of peaks (mz), intensities of peaks (int), species names (species), and strain names (strain). Species and strain columns may contain more than one species/strain if an option 2 is chosen.

Examples

```
spectra.processed.A <- process_monospectra(</pre>
   file=system.file("extdata", "listA.txt", package="MGMS2"),
   mass.range=c(1000,2200))
spectra.processed.B <- process_monospectra(</pre>
   file=system.file("extdata", "listB.txt", package="MGMS2"),
   mass.range=c(1000, 2200))
spectra.processed.C <- process_monospectra(</pre>
   file=system.file("extdata", "listC.txt", package="MGMS2"),
   mass.range=c(1000,2200))
spectra.mono.summary.A <- summarize_monospectra(</pre>
   processed.obj=spectra.processed.A,
   species='A', directory=tempdir())
spectra.mono.summary.B <- summarize_monospectra(</pre>
   processed.obj=spectra.processed.B,
   species='B', directory=tempdir())
spectra.mono.summary.C <- summarize_monospectra(</pre>
   processed.obj=spectra.processed.C,
   species='C', directory=tempdir())
mono.info=gather_summary(c(spectra.mono.summary.A, spectra.mono.summary.B, spectra.mono.summary.C))
mixture.ratio <- list()</pre>
mixture.ratio['A']=1
mixture.ratio['B']=0.5
mixture.ratio['C']=0
sim.template <- create_insilico_mixture_template(mono.info)</pre>
insilico.spectrum <- simulate_poly_spectra(sim.template, mixture.ratio)</pre>
merged.spectrum <- characterize_peak(insilico.spectrum, option=2)</pre>
```

```
create_insilico_mixture_template

create insilico mixture template
```

Description

This function generates an intial template for simulated mass spectra.

Usage

```
create_insilico_mixture_template(mono.info, mz.tol = 0.5)
```

Arguments

```
mono.info An output of gather_summary.

mz.tol A m/z tolerance in Da. (Default: 0.5)
```

Value

A data frame which contains simulated m/z, log intensity, and normalized intensity values of peaks.

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Examples

```
spectra.processed.A <- process_monospectra(</pre>
   file=system.file("extdata", "listA.txt", package="MGMS2"),
   mass.range=c(1000,2200))
spectra.processed.B <- process_monospectra(</pre>
   file=system.file("extdata", "listB.txt", package="MGMS2"),
   mass.range=c(1000,2200))
spectra.processed.C <- process_monospectra(</pre>
   file=system.file("extdata", "listC.txt", package="MGMS2"),
   mass.range=c(1000,2200))
spectra.mono.summary.A <- summarize_monospectra(</pre>
   processed.obj=spectra.processed.A,
   species='A', directory=tempdir())
spectra.mono.summary.B <- summarize_monospectra(</pre>
   processed.obj=spectra.processed.B,
   species='B', directory=tempdir())
spectra.mono.summary.C <- summarize_monospectra(</pre>
   processed.obj=spectra.processed.C,
   species='C', directory=tempdir())
mono.info=gather_summary(c(spectra.mono.summary.A, spectra.mono.summary.B, spectra.mono.summary.C))
template <- create_insilico_mixture_template(mono.info)</pre>
```

filtermass

filtermass

Description

Internal function. This function removes peaks with their mass values (m/z values) outside a given mass range. This function is used in process_monospectra.

Usage

```
filtermass(spectra, mass.range)
```

Arguments

spectra Mass Spectra (A MALDIquant MassSpectrum (S4) object). An output of importMzXml.

mass.range Mass (m/z) range $(a \ vector)$. For exmaple, c(1000,2200).

Value

A list of filtered mass spectra (MALDIquant MassSpectrum (S4) objects) which contains mass, intensity, and metaData.

gather_summary 5

gather_summary gather_summary

Description

This function combines outputs from summarize_monospectra.

Usage

```
gather_summary(x)
```

Arguments

Х

A list of multiple monomicrobial mass spectra information from summarize_monospectra.

Value

A list of combined summaries (data frames) of mass spectra from summarize_monospectra and the corresponding species (a vector).

Examples

```
spectra.processed.A <- process_monospectra(</pre>
   file=system.file("extdata", "listA.txt", package="MGMS2"),
   mass.range=c(1000,2200))
spectra.processed.B <- process_monospectra(</pre>
   file=system.file("extdata", "listB.txt", package="MGMS2"),
   mass.range=c(1000,2200))
spectra.processed.C <- process_monospectra(</pre>
   file=system.file("extdata", "listC.txt", package="MGMS2"),
   mass.range=c(1000,2200))
spectra.mono.summary.A <- summarize_monospectra(</pre>
   processed.obj=spectra.processed.A,
   species='A', directory=tempdir())
spectra.mono.summary.B <- summarize_monospectra(</pre>
   processed.obj=spectra.processed.B,
   species='B', directory=tempdir())
spectra.mono.summary.C <- summarize_monospectra(</pre>
   processed.obj=spectra.processed.C,
   species='C', directory=tempdir())
mono.info=gather_summary(c(spectra.mono.summary.A, spectra.mono.summary.B, spectra.mono.summary.C))
```

gather_summary_file

```
gather_summary_file gather_summary_file
```

Description

This function combines output files from summarize_monospectra.

Usage

```
gather_summary_file(directory)
```

Arguments

directory

A directory that contains summary files from summarize_monospectra.

Value

A list of combined summaries of mass spectra (data frames) from summarize_monospectra and the corresponding species (a vector).

Examples

```
spectra.processed.A <- process_monospectra(</pre>
   file=system.file("extdata", "listA.txt", package="MGMS2"),
   mass.range=c(1000,2200))
spectra.processed.B <- process_monospectra(</pre>
   file=system.file("extdata", "listB.txt", package="MGMS2"),
   mass.range=c(1000,2200))
spectra.processed.C <- process_monospectra(</pre>
   file=system.file("extdata", "listC.txt", package="MGMS2"),
   mass.range=c(1000,2200))
spectra.mono.summary.A <- summarize_monospectra(</pre>
   processed.obj=spectra.processed.A,
   species='A', directory=tempdir())
spectra.mono.summary.B <- summarize_monospectra(</pre>
   processed.obj=spectra.processed.B,
   species='B', directory=tempdir())
spectra.mono.summary.C <- summarize_monospectra(</pre>
   processed.obj=spectra.processed.C,
   species='C', directory=tempdir())
summary <- gather_summary_file(directory=tempdir())</pre>
```

preprocessMS 7

Description

Internal function. This function preprocesses spectra by transforming/smoothing intensity, removing baseline, and calibrating intensities.

Usage

```
preprocessMS(spectra, halfWindowSize = 20, SNIP.iteration = 60)
```

Arguments

```
spectra Spectra. A MALDIquant object. An output of either importMzXml or filtermass.

halfWindowSize halfWindowSize The highest peaks in the given window (+/-halfWindowSize) will be recognized as peaks. (Default: 20). See detectPeaks for details.

SNIP.iteration An iteration used to remove the baseline of an spectrum. (Default: 60). See removeBaseline for details.
```

Value

The processed mass spectra. A list of MALDIquant MassSpectrum objects (S4 objects).

```
process_monospectra process_monospectra
```

Description

This function processes multiple mzXML files which are listed in the file that an user specifies.

Usage

```
process_monospectra(
   file,
   mass.range = c(1000, 2200),
   halfWindowSize = 20,
   SNIP.iteration = 60
)
```

Arguments

file A file name. This file is a tab-delimited file which contains the following columns:

file names, strain.no, and strain. See below for details.

mass.range The m/z range that users want to consider for the analysis. (Default: c(1000,2200)).

halfWindowSize A half window size used for the smoothing the intensity values. (Default: 20).

See smoothIntensity for details.

SNIP. iteration An iteration used to remove the baseline of an spectrum. (Default: 60). See

removeBaseline for details.

Value

A list of processed monobacterial mass spectra (S4 objects, MALDIquant MassSpectrum objects), and their strain numbers (a vector), unique strains (a vector), and strain names (a vector).

Examples

```
spectra.processed.A <- process_monospectra(
   file=system.file("extdata", "listA.txt", package="MGMS2"),
   mass.range=c(1000,2200))</pre>
```

Description

Internal function. The function simulates m/z and intensity values using given summary statistics.

Usage

```
simulate_ind_spec_single(interest, mz.tol, species, strain)
```

Arguments

interest Summary statistics of spectra.

mz.tol The tolerance of m/z. This is used to generate m/z values of peaks.

species Species. strain Strain name.

Value

A data frame that contains m/z, (normalized) intensity values, missing rates of peaks, species name, and strain name.

Description

The function creates simulated mass spectra in pdf file and returns simulated mass spectra (m/z and intensity values of peaks).

Usage

```
simulate_many_poly_spectra(
  mono.info,
  nsim = 10000,
  file = NULL,
  mixture.ratio,
  mixture.missing.prob.peak = 0.05,
  noise.peak.ratio = 0.05,
  snr.basepeak = 500,
  noise.cv = 0.25,
  mz.range = c(1000, 2200),
  mz.tol = 0.5
)
```

Arguments

```
mono.info
                  A list output of gather_summary or gather_summary_file.
                  The number of simulated spectra. (Default: 10000)
nsim
                  An output file name. (By default, file=NULL. No pdf file will be generated.)
file
                  A list of bacterial mixture ratios for given bacterial species in sim.template.
mixture.ratio
mixture.missing.prob.peak
                  A real value. The missing probability caused by mixing multiple bacteria species.
                  (Default: 0.05)
noise.peak.ratio
                  A ratio between the numbers of noise and signal peaks. (Default: 0.05)
snr.basepeak
                  A (base peak) signal to noise ratio. (Default: 5000)
noise.cv
                  A coefficient of variation of noise peaks. (Default: 0.25)
                  A range of m/z values. (Default: c(1000,2200))
mz.range
mz.tol
                  m/z tolerance. (Default: 0.5)
```

Value

A list of data frames. A list of simulated mass spectra (data frames) that contains m/z values of peaks, normalized intensities of peaks, species names, and strain names. This function also creates pdf files which contain simulated spectra.

Examples

```
spectra.processed.A <- process_monospectra(</pre>
   file=system.file("extdata", "listA.txt", package="MGMS2"),
   mass.range=c(1000,2200))
spectra.processed.B <- process_monospectra(</pre>
   file=system.file("extdata", "listB.txt", package="MGMS2"),
   mass.range=c(1000,2200))
spectra.processed.C <- process_monospectra(</pre>
   file=system.file("extdata", "listC.txt", package="MGMS2"),
   mass.range=c(1000,2200))
spectra.mono.summary.A <- summarize_monospectra(</pre>
   processed.obj=spectra.processed.A,
   species='A', directory=tempdir())
spectra.mono.summary.B <- summarize_monospectra(</pre>
   processed.obj=spectra.processed.B,
   species='B', directory=tempdir())
spectra.mono.summary.C <- summarize_monospectra(</pre>
   processed.obj=spectra.processed.C,
   species='C', directory=tempdir())
mono.info=gather_summary(c(spectra.mono.summary.A, spectra.mono.summary.B, spectra.mono.summary.C))
mixture.ratio <- list()</pre>
mixture.ratio['A']=1
mixture.ratio['B']=0.5
mixture.ratio['C']=0
insilico.spectra <- simulate_many_poly_spectra(mono.info, mixture.ratio=mixture.ratio, nsim=10)
```

simulate_poly_spectra simulate_poly_spectra

Description

This function takes simulated m/z and intensities of peaks from create_insilico_mixture_template and modifies them based on given parameters.

Usage

```
simulate_poly_spectra(
    sim.template,
    mixture.ratio,
    spectrum.name = "Spectrum",
    mixture.missing.prob.peak = 0.05,
    noise.peak.ratio = 0.05,
    snr.basepeak = 500,
    noise.cv = 0.25,
    mz.range = c(1000, 2200)
)
```

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Arguments

sim.template A data frame which contains m/z, log intensity, normalized intensity values and missing rates of peaks. There are also species and strain information. An object of create_insilico_mixture_template. mixture.ratio A list of bacterial mixture ratios for given bacterial species in sim.template. spectrum.name A character. An user can define the spectrum name. (Default: 'Spectrum'). mixture.missing.prob.peak A real value. The missing probability caused by mixing multiple bacteria species. (Default: 0.05) noise.peak.ratio A ratio between the numbers of noise and signal peaks. (Default: 0.05) snr.basepeak A (base peak) signal to noise ratio. (Default: 500) A coefficient of variation of noise peaks. (Default: 0.25) noise.cv A range of m/z values. (Default: c(1000,2200)) mz.range

Value

A data frame that contains m/z values of peaks, normalized intensities of peaks, species names, and strain names. A modified version of sim.template.

Examples

```
spectra.processed.A <- process_monospectra(</pre>
   file=system.file("extdata", "listA.txt", package="MGMS2"),
   mass.range=c(1000,2200))
spectra.processed.B <- process_monospectra(</pre>
   file=system.file("extdata", "listB.txt", package="MGMS2"),
   mass.range=c(1000,2200))
spectra.processed.C <- process_monospectra(</pre>
   file=system.file("extdata", "listC.txt", package="MGMS2"),
   mass.range=c(1000,2200))
spectra.mono.summary.A <- summarize_monospectra(</pre>
   processed.obj=spectra.processed.A,
   species='A', directory=tempdir())
spectra.mono.summary.B <- summarize_monospectra(</pre>
   processed.obj=spectra.processed.B,
   species='B', directory=tempdir())
spectra.mono.summary.C <- summarize_monospectra(</pre>
   processed.obj=spectra.processed.C,
   species='C', directory=tempdir())
mono.info=gather_summary(c(spectra.mono.summary.A, spectra.mono.summary.B, spectra.mono.summary.C))
mixture.ratio <- list()</pre>
mixture.ratio['A']=1
mixture.ratio['B']=0.5
mixture.ratio['C']=0
sim.template <- create_insilico_mixture_template(mono.info)</pre>
insilico.spectrum <- simulate_poly_spectra(sim.template, mixture.ratio)</pre>
```

```
summarize_monospectra summarize monospectra
```

Description

This function summarizes monomicrobial spectra and writes summary in the specified directory.

Usage

```
summarize_monospectra(
  processed.obj,
  species,
  directory = NULL,
  minFrequency = 0.5,
  align.tolerance = 5e-04,
  snr = 3,
  halfWindowSize = 20,
  top.N = 50
)
```

Arguments

 $\verb|processed.obj| A list from \verb|process_monospectra| which contains peaks information for each$

strain.

species Species name.

directory Directory. (By default, no summary file will be generated.)

minFrequency Percentage value. A minimum occurrence proportion required for building a ref-

erence peaks. All peaks with their occurence proportion less than minFrequency will be moved. (Default: 0.50). See filterPeaks and referencePeaks for de-

tails.

align.tolerance

Mass tolerance. Must be multiplied by 10^-6 for ppm. (Default: 0.0005).

snr Signal-to-noise ratio. (Default: 3).

halfWindowSize The highest peaks in the given window (+/-halfWindowSize) will be recognized

as peaks. (Default: 20). See detectPeaks for details.

top. N The top N peaks will be chosen for the analysis. An integer value. (Default: 50).

Value

A data frame that contains the peaks informations: m/z, mean log intensity, standard deviation of log intensity, missing rate of peaks. In addition, it also contains species and strain information.

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Examples

```
spectra.processed.A <- process_monospectra(</pre>
   file=system.file("extdata", "listA.txt", package="MGMS2"),
   mass.range=c(1000,2200))
spectra.mono.summary.A <- summarize_monospectra(</pre>
   processed.obj=spectra.processed.A, species='A',
   directory=tempdir())
```

summary_mono

summary_mono

Description

Internal function. This function calculates summary statistics for peaks afterling aligning spectra of interest.

Usage

```
summary_mono(
  spectra.interest,
 minFrequency = 0.5,
  align.tolerance = 5e-04,
  snr = 3,
 halfWindowSize = 20,
  top.N = 50
)
```

Arguments

spectra.interest

A list which contains peaks information for a strain of interest.

minFrequency

Percentage value. A minimum occurrence proportion required for building a reference peaks. All peaks with their occurence proportion less than minFrequency will be moved. (Default: 0.50). See filterPeaks and referencePeaks for de-

tails.

align.tolerance

Mass tolerance. Must be multiplied by 10^-6 for ppm. (Default: 0.0005).

snr

Signal-to-noise ratio. (Default: 3).

halfWindowSize The highest peaks in the given window (+/-halfWindowSize) will be recognized as peaks. (Default: 20). See detectPeaks for details.

top.N

The top N peaks will be chosen for the analysis. An integer value. (Default: 50).

Value

Summary information (Data frame) of spectra of interest.

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