

Package ‘LargeMetabo’

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Type Package

Version 0.1.0

Title an out-of-the-box tool for processing and analyzing large-scale metabolomic data

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Depends R (≥ 3.0), dynamicTreeCut (≥ 1.62), fastcluster

Imports genefilter, d3heatmap, ggplot2, ggfortify, factoextra, igraph, FSelector, varSelRF, corrplot, mixOmics, ropls, CluMSID, siggenes, e1071, SOMbrero

Suggests rmarkdown, knitr, hexbin, shinycssloaders, shinyBS, shinyjs, shinyjqui

Description This package for processing and analyzing large-scale metabolomic data, including data combination for metabolomics data, biomarker selection, enhanced metabolite annotation and enrichment analysis.

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URL <https://github.com/LargeMetabo/LargeMetabo>

RoxygenNote 7.1.2

NeedsCompilation no

LazyData true

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Integrate_Data	Integrating multiple datasets
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Description

Integrating multiple datasets from different analytical experiments

Usage

Integrate_Data(mutile_Group, RT_Tolerance_1 = 10, mz_Tolerance_1 = 0.1, RT_Tolerance_2 = 10, mz_Tolerance_2 = 0.1)

Arguments

mutile_Group	Multiple datasets from multiple groups for integrating as a comprehensive dataset.
RT_Tolerance_1	Set the tolerance of retention time for data integration in the primary phase.
mz_Tolerance_1	Set the tolerance of mass-to-charge ratio for data integration in the primary phase.

RT_Tolerance_2	Set the tolerance of retention time for data integration in the secondary phase.
mz_Tolerance_2	Set the tolerance of mass-to-charge ratio for data integration in the secondary phase.

Value

A matrix of a comprehensive dataset integrated from multiple groups is returned.

Author(s)

Qingxia Yang <yangqx@cqu.edu.cn>

Examples

```
Integrate_Data(mutile_Group, RT_Tolerance_1 = 10, mz_Tolerance_1 = 0.1, RT_Tolerance_2 = 10, mz_Tolerance_2 = 0.1)
```

Removal_Batch	Remove batch effects for the integrated dataset
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Description

Remove batch effects among different analytical experiments for the integrated dataset

Usage

```
Removal_Batch(mutile_align, n = 3, algorithm = "BMC/PAMR")
```

Arguments

mutile_align	The comprehensive dataset integrated from multiple groups.
n	The number of multiple groups.
algorithm	The algorithm for removing batch effects in multiple groups. The algorithm could be BMC/PAMR, ComBat/EB, GlobalNorm, Log2 or None.

Value

A matrix of a dataset after removing batch effects for the integrated dataset.

Author(s)

Qingxia Yang <yangqx@cqu.edu.cn>

Examples

```
Removal_Batch(mutile_align, n = 3, algorithm = "BMC/PAMR")
```

Sample_Separation	Plot of sample separation
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Description

Plot of sample separation for visualizing the clustering and separation of different samples

Usage

```
Sample_Separation(finalData, finalLabel, clusters=2, method = "HCA")
```

Arguments

finalData	The matrix of dataset for sample separation.
finalLabel	The label of dataset for sample separation.
clusters	The number of cluster for sample separation.
method	The method for sample separation. The method could be HCA, KMC, PCA and SOM.

Value

The plot of sample separation.

Author(s)

Qingxia Yang <yangqx@cqu.edu.cn>

Examples

```
finalData <- MarkerData$finalData
```

```
finalLabel <- MarkerData$finalLabel
```

```
Sample_Separation(finalData, finalLabel, clusters=2, method = "HCA")
```

Marker_Identify	Marker identification in large-scale metabolomics data
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Description

Marker identification in large-scale metabolomics data

Usage

```
Marker_Identify(finalData, finalLabel, method = "FC")
```

Arguments

finalData	The matrix of dataset for marker identification.
finalLabel	The label of dataset for marker identification.
method	The method for marker identification. The method could be FC, PLS-DA, OPLS-DA, t-test, CHIS, CFS, ENTROPY, LMEB, RELIEF, RF, SAM, SVMRFE or WRST.

Value

The result of marker identification for metabolomics data.

Author(s)

Qingxia Yang <yangqx@cqu.edu.cn>

Examples

```
finalData <- MarkerData$finalData
```

```
finalLabel <- MarkerData$finalLabel
```

```
Marker_Identify(finalData, finalLabel, method = "FC")
```

MetaboAnnotation	Metabolite annotation for MS1 spectra
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Description

Metabolite annotation for MS1 spectra based on a metabolite database

Usage

MetaboAnnotation(AnnotaData, masstole = 10, toleUnit = 1, annotaDB = "metlin", ionMode = "pos")

Arguments

AnnotaData	The value of mass-to-charge ratio for metabolite annotation.
masstole	The tolerance of m/z for metabolite annotation.
toleUnit	The unit of m/z for metabolite annotation. The unit could be 1 for Da, or 2 for ppm.
annotaDB	The database of m/z for metabolite annotation. The database could be metlin or hmdb.
ionMode	The mode of m/z for annotation. The mode could be pos, neg or neu.

Value

The list of results of metabolite annotation for different charges for primary mass spectrometry.

Author(s)

Qingxia Yang <yangqx@cqu.edu.cn>

Examples

```
AnnotaMS <- AnnotaData$AnnotaMS
MetaboA_res <- MetaboAnnotation(AnnotaMS)
MetaboA_res$`M+H-2H2O`
```

AnnotaTandem

Parameters of metabolite annotation for MS/MS

Description

Parameters of metabolite annotation of tandem mass spectrum (MS/MS)

Usage

```
AnnotaTandem(Parent_mass = 181.04, TandemData, massTandem = 0.1, toleUnitTandem = 1,
massmzTandem = 0.5, toleUnitmzTandem = 1, ModeTandem = "Positive", ionEnergy =
"low(10V)")
```

Arguments

Parent_mass	The value of parent ion mass for metabolite annotation of tandem mass spectrum.
TandemData	The value of MS/MS peak list (m/z & Intensity) for metabolite annotation of tandem mass spectrum.
massTandem	The tolerance of parent ion mass for metabolite annotation of tandem mass spectrum.
toleUnitTandem	The unit of parent ion mass for metabolite annotation of tandem mass spectrum. The unit could be 1 for Da, or 2 for ppm.
massmzTandem	The tolerance of MS/MS peak mass for metabolite annotation of tandem mass spectrum.
toleUnitmzTandem	The unit of MS/MS peak mass for metabolite annotation of tandem mass spectrum. The unit could be 1 for Da, or 2 for ppm.
ModeTandem	The ionization mode for metabolite annotation of tandem mass spectrum. The mode could be Positive or Negative.
ionEnergy	The CID energy for metabolite annotation of tandem mass spectrum. The mode could be low(10V) Medium(25V) High(40V) or All.

Value

The parameters of metabolite annotation of tandem mass spectrum (MS/MS).

Author(s)

Qingxia Yang <yangqx@cqu.edu.cn>

Examples

```
Parent_mass <- AnnotaData$Parent_mass
```

```
TandemData <- AnnotaData$TandomData
```

```
AnnotaParamTandem <- AnnotaTandem(Parent_mass, TandemData)
```

annotaDataTandem	Table of metabolite annotation for MS/MS
------------------	--

Description

The table of results for metabolite annotation of tandem mass spectrum (MS/MS).

Usage

```
annotaDataTandem(AnnotaParamTandem)
```

Arguments

AnnotaParamTandem The parameters of metabolite annotation of tandem mass spectrum.

Value

The table of results for metabolite annotation of tandem mass spectrum.

Author(s)

Qingxia Yang <yangqx@cqu.edu.cn>

Examples

```
Parent_mass <- AnnotaData$Parent_mass
```

```
TandemData <- AnnotaData$TandomData
```

```
AnnotaParamTandem <- AnnotaTandem(Parent_mass, TandemData)
```

```
annotaDataTandem(AnnotaParamTandem)
```

AnnotaTandem_plot	Plot of metabolite annotation of MS/MS
-------------------	--

Description

The plot of results for metabolite annotation of tandem mass spectrum (MS/MS).

Usage

```
AnnotaTandem_plot(AnnotaParamTandem, TandemData)
```

Arguments

AnnotaParamTandem The parameters of metabolite annotation of tandem mass spectrum.

TandemData The value of MS/MS peak list (m/z & Intensity) for metabolite annotation of tandem mass spectrum.

Value

The plot of results for metabolite annotation of tandem mass spectrum.

Author(s)

Qingxia Yang <yangqx@cqu.edu.cn>

Examples

```
Parent_mass <- AnnotaData$Parent_mass
```

```
TandemData <- AnnotaData$TandomData
```

```
AnnotaParamTandem <- AnnotaTandem(Parent_mass, TandemData)
```

```
AnnotaTandem_plot(AnnotaParamTandem, TandemData)
```

KEGGENrichPlotPanel	Parameters for metabolite enrichment
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Description

The parameters for metabolite enrichment based on the metabolite database.

Usage

```
KEGGENrichPlotPanel(sampleData, enrichDB = "kegg", pvalcutoff = 0.05, IDtype = 1, cateIdx = 1)
```

Arguments

sampleData The charactor of input for metabolite enrichment.

enrichDB The database of the input data. The database could be kegg, smpdb, cfam, foodb, biofunc, tcm, spectax or toxin.

pvalcutoff The cutoff of p value for metabolite enrichment.

IDtype The number for name type for metabolite enrichment, such as KEGG ID, CAS ID, PubChem ID, Name or HMDB ID.

cateIdx The number of category for metabolite enrichment.

Value

The parameters for metabolite enrichment.

Author(s)

Qingxia Yang <yangqx@cqu.edu.cn>

Examples

```
sampleDatakegg <- EnrichData$sampleDatakegg
```

```
KEGGenrichPlotPanel(sampleDatakegg, enrichDB = "kegg", pvalcutoff = 0.05, IDtype = 1,  
cateIdx = 1)
```

Enrichment	Table for metabolite enrichment
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Description

The table for metabolite enrichment.

Usage

```
Enrichment(param)
```

Arguments

param The parameters for metabolite enrichment.

Value

The table of results for metabolite enrichment.

Author(s)

Qingxia Yang <yangqx@cqu.edu.cn>

Examples

```
sampleDatakegg <- EnrichData$sampleDatakegg
```

```
EnrichParam <- KEGGenrichPlotPanel(sampleDatakegg, enrichDB = "kegg", pvalcutoff = 0.05,  
IDtype = 1, cateIdx = 1)
```

```
EnrichResultList <- Enrichment(EnrichParam); EnrichResultList$Table.Result
```

KEGGENrichPlot	Plot of metabolite enrichment using KEGG database
----------------	---

Description

The plot of metabolite enrichment using KEGG database.

Usage

```
KEGGENrichPlot(EnrichResultList, cpdID, cpdFC)
```

Arguments

EnrichResultList The table of results for metabolite enrichment using KEGG database.

cpdID The character of input for metabolite enrichment.

cpdFC The distance in the metabolite enrichment plot using KEGG database.

Value

The plot of results for metabolite enrichment using KEGG database.

Author(s)

Qingxia Yang <yangqx@cqu.edu.cn>

Examples

```
sampleDatakegg <- EnrichData$sampleDatakegg
```

```
EnrichParam <- KEGGENrichPlotPanel(sampleDatakegg, enrichDB = "kegg", pvalcutoff = 0.05,  
IDtype = 1, cateIdx = 1)
```

```
EnrichResultList <- Enrichment(EnrichParam)
```

```
EnrichFC <- seq(from = -2, to = 2, length.out = 24)
```

```
KEGGENrichPlot(EnrichResultList = EnrichResultList, cpdID = sampleDatakegg, cpdFC =  
EnrichFC)
```

EnrichPlot	Plot of results for metabolite enrichment
------------	---

Description

The plot of results for metabolite enrichment.

Usage

`EnrichPlot(db, EnrichResultList)`

Arguments

`db` The database of input for metabolite enrichment.

`EnrichResultList` The table of results for metabolite enrichment.

Value

The plot of results for metabolite enrichment.

Author(s)

Qingxia Yang <yangqx@cqu.edu.cn>

Examples

```
sampleDataacas <- EnrichData$sampleDataacas  
enrichDB <- EnrichData$enrichDB  
  
EnrichParam <- KEGGEnrichPlotPanel(sampleDataacas, enrichDB = enrichDB, pvalcutoff =  
0.05, IDtype = 2, cateIdx = 1)  
  
EnrichResultList <- Enrichment(EnrichParam)  
  
dbChoice <- enrichDB  
  
EnrichPlot(dbChoice, EnrichResultList)
```