# Package 'MSbox'

## December 8, 2022

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| Description Common mass spectrometry tools described in John Roboz (2013) <doi:10.1201 b15436="">. It allows checking element isotopes, calculating (isotope labelled) exact monoisitopic mass, m/z values and mass accuracy, and inspecting possible contaminant mass peaks, examining possible adducts in electrospray ionization (ESI) and matrix-assisted laser desorption ionization (MALDI) ion sources.</doi:10.1201> |
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adduct

Common adducts

## **Description**

calculate common adduct ions in positive or negative ion mode

## Usage

```
adduct(F, mode = c("+", "-"))
```

## **Arguments**

F chemical formula, case insensitive

mode ionization mode, either positive '+' or negative '-'

#### Author(s)

Yonghui Dong

```
adduct('C1H4', mode = '-')
adduct('C1h4', mode = '+')
```

contam 3

contam

Contaminants in MS

## **Description**

check the possible contaminants

## Usage

```
contam(mz, mode = NULL, ppm = 10)
```

#### **Arguments**

mz suspected m/z value

mode ionization mode, either positive '+' or negative '-'

ppm mass tolerance, default value = 10

#### Author(s)

Yonghui Dong

## **Examples**

```
contam(33.0335, ppm = 10, mode = '+')
contam(44.998, ppm = 10, mode = '-')
```

describe

Get the compound information

## **Description**

get compound formula and structure from https://cactus.nci.nih.gov/chemical/structure

#### Usage

```
describe(chem, representation = "formula", info = FALSE)
```

## **Arguments**

chem, chemical name of the compound

representation,

representation methods, formula is default

info, extra molecular information that users can query

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#### Author(s)

Yonghui Dong

## **Examples**

```
## Not run:
describe('malic acid', "formula")
describe(c('malic acid', 'citric acid', 'tartaric acid'), "smiles")
## End(Not run)
```

doStat

Performing statistics

## Description

performing statistics, including calculating fold change, p-values and VIP values

## Usage

```
doStat(x, Group = NULL)
```

## **Arguments**

x sample ion intensity matrix, row sample, column feature.Group sample group information

#### Value

a dataframe with statistical information

```
dat <- matrix(runif(2*300), ncol = 2, nrow = 300)
rownames(dat) <- 1:dim(dat)[1]
myGroup <- rep_len(LETTERS[1:3], 300)
ret <- doStat(dat, Group = myGroup)</pre>
```

E\_iso 5

E\_iso

Element isotopes

## **Description**

check element isotope information

## Usage

E\_iso(S)

## Arguments

S

element, can be element symbol (i.e. C) or full name (i.e. Carbon). Both Element symbol and full name are case insensitive.

#### Author(s)

Yonghui Dong

## **Examples**

```
E_iso('Na') # element symbol
E_iso('nA') # element symbol, case insensitive
E_iso('Carbon') # element full name
E_iso('carBon') # element full name, case insensitive
```

getCV

Calculate coefficient of variation (CV)

## Description

Calculate coefficient of variation (CV), also known as relative standard deviation (RSD) among different sample groups

## Usage

```
getCV(x, Group = NULL)
```

## Arguments

x sample ion intensity matrix, row sample, column feature.

Group sample group information

#### Value

a dataframe with mean values and cv

6 getMax

#### **Examples**

```
dat <- matrix(runif(2*300), ncol = 2, nrow = 300)
myGroup <- rep_len(LETTERS[1:2], 300)
ret <- getCV(dat, Group = myGroup)</pre>
```

getFC

calculate fold change

## Description

calculate fold change among different samples.

## Usage

```
getFC(x, Group = NULL)
```

## **Arguments**

x sample ion intensity matrix, row sample, column feature.

Group sample group information

#### Value

a dataframe with mean values and fold changes

## **Examples**

```
dat <- matrix(runif(2*300), ncol = 2, nrow = 300)
myGroup <- rep_len(LETTERS[1:2], 300)
ret <- getFC(dat, Group = myGroup)</pre>
```

getMax

Get the sample name which has the max ion intensity

## Description

get the sample name which has the max ion intensity

#### Usage

```
getMax(x)
```

#### **Arguments**

x sample ion intensity matrix, row sample, column feature.

getP 7

## Value

a data frame

## **Examples**

```
dat <- cbind.data.frame(mz = c(100, 101, 300), mz2 = c(0, 0 , 1), mz3 = c(1, 9, 1)) rownames(dat) <- c("A", "B", "C") out <- getMax(dat)
```

getP

get p-values

## Description

get p-values from Post Hoc analysis

## Usage

```
getP(x, Group = NULL)
```

## **Arguments**

x sample ion intensity matrix, row sample, column feature.

Group sample group information

#### Value

a data frame

```
dat <- matrix(runif(2*300), ncol = 2, nrow = 300)
myGroup <- rep_len(LETTERS[1:3], 300)
out <- getP(dat, Group = myGroup)</pre>
```

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Iso\_mass

Isotope labelled molecular mass

## **Description**

Calculate isotope labelled molecular mass

## Usage

```
Iso_mass(F, iso)
```

## Arguments

F, chemical formula, case insensitive iso, labelled elements, case insensitive

#### Author(s)

Yonghui Dong

#### **Examples**

```
Iso_mass(F = 'C7H604', iso = '[13]C2[2]H3') # Two 13C and three 2H are labled
```

Iso\_mz

Isotope labelled molecular mass

## Description

Calculate isotope labelled m/z

## Usage

```
Iso_mz(F, iso, z)
```

## **Arguments**

F, chemical formula, case insensitive iso, labelled elements, case insensitive z charge

## Author(s)

Yonghui Dong

```
Iso_mz(F = 'C7H604', iso = '[13]C2[2]H3', z = -1) \# Two 13C and three 2H are labled
```

mass 9

mass

molecular mass

## Description

calculate accurate molecular mass

#### Usage

```
mass(F, caseSensitive = FALSE)
```

#### **Arguments**

F

chemical formula, case insensitive

caseSensitive

if case sensitive is 'FALSE' (default), the elements are seperated by numbers. for instance, Carbon dioxyde can be written as 'c1o2' or any combination of the two elements in lower or upper cases. However, the number of elements should be clearly stated in the chemical formula. if case sensitive is 'TRUE', the elements are seperated by upper case letters. For instance, Carbon dioxyde must be written as 'C1O2' or 'CO2'. You don't meed to write the number of the element if it is 1.

#### Author(s)

Yonghui Dong

## **Examples**

```
\label{eq:mass('C7h701')} $mass('C7H70', caseSensitive = TRUE)$ $mass(c('C7H704', 'C'), caseSensitive = TRUE) $\#$ vector input $mass(c('c7h704', 'c1'))$ $$
```

mz

Calculate accurate mass-to-charge ratio

## **Description**

Calculate accurate mass-to-charge ratio (m/z)

## Usage

```
mz(m, z, caseSensitive = FALSE)
```

10 ppm

#### **Arguments**

m chemical formula of an ion, case insensitive

z charge

caseSensitive if case sensitive is 'FALSE' (default), the elements are seperated by numbers.

for instance, Carbon dioxyde can be written as 'c1o2' or any combination of the two elements in lower or upper cases. However, the number of elements should be clearly stated in the chemical formula. if case sensitive is 'TRUE', the elements are seperated by upper case letters. For instance, Carbon dioxyde must be written as 'C1O2' or 'CO2'. You don't meed to write the number of the

element if it is 1.

#### Author(s)

Yonghui Dong

## **Examples**

```
mz('C7h7o1', z=1)
mz('C7H7O', z=1, caseSensitive = TRUE)
mz(c('C7H7O4', 'C'), z=-1, caseSensitive = TRUE) # vector input
mz(c('c7h7O4', 'c1'), z=-1)
```

ppm

mass accuracy

#### **Description**

calculate the mass accuracy of measured m/z. lazy input allowed

## Usage

```
ppm(m, t, lazy = TRUE)
```

## **Arguments**

 $\begin{array}{ll} m & measured \ m/z \\ t & theoretical \ m/z \end{array}$ 

lazy if lazy input is allowed

#### Author(s)

Yonghui Dong

```
ppm(155.03383, 155.03388) # with m/z value ppm(155.03383, .03388) # lazy input when the integer parts of m and t are the same ppm(155.03384, mz('C7H7O4', z = 1)) # with ion formula
```

searchDB 11

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Search in customized database

#### **Description**

search in customized database based on accurate m/z and RT

#### Usage

```
searchDB(DF, DB, ppm = 5, RT = 0.2, useRT = FALSE)
```

#### **Arguments**

DF input file, should contain at least a column named mz
DB database, should contain at least a column named mz

ppm mass tolerance, default 5ppm

RT retention time tolerance, default 0.2min

useRT should RT be considered during database search?

#### Author(s)

Yonghui Dong

## **Examples**

```
DF <- cbind.data.frame(mz = c(100.001, 100.1), RT = c(10, 11))
DB <- cbind.data.frame(mz = c(100.001, 100.1), RT = c(10, 12.1))
searchDB(DF, DB, ppm = 5, RT = 0.2, useRT = TRUE)
```

what

search for m/z in from the idiom metabolomics database

## Description

tentative metabolite identification based on m/z value search

## Usage

```
what(myMZ, mode = NULL, ppm = 5, useDB = "HMDB")
```

## **Arguments**

myMZ m/z values

mode ionization mode, either positive '+' or negative '-'

ppm mass tolerance, default value = 10

useDB which database to use, HMDB or KEGG? default is HMDB

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## Author(s)

Yonghui Dong

```
a = what(133.014, mode = '-', ppm = 10)
```

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