

# Package ‘Rdisop’

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**Title** Decomposition of Isotopic Patterns

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**Description** Identification of metabolites using high precision mass spectrometry. MS Peaks are used to derive a ranked list of sum formulae, alternatively for a given sum formula the theoretical isotope distribution can be calculated to search in MS peak lists.

**Depends** R (>= 2.0.0), Rcpp

**LinkingTo** Rcpp

**Suggests** RUnit

**SystemRequirements** None

**License** GPL-2

**StagedInstall** no

**URL** <https://github.com/sneumann/Rdisop>

**BugReports** <https://github.com/sneumann/Rdisop/issues/new>

**biocViews** ImmunoOncology, MassSpectrometry, Metabolomics

**git\_url** <https://git.bioconductor.org/packages/Rdisop>

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## R topics documented:

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|--------------|----------------------------------|
| addMolecules | <i>Add/subtract sum formulae</i> |
|--------------|----------------------------------|

### Description

Simple arithmetic modifications of sum formulae.

### Usage

```
addMolecules(formula1, formula2, elements = NULL, maxisotopes = 10)
subMolecules(formula1, formula2, elements = NULL, maxisotopes = 10)
```

### Arguments

|             |   |
|-------------|---|
| formula1    | Sum formula   |
| formula2    | Sum formula   |
| elements    | list of allowed chemical elements, defaults to full periodic system of elements |
| maxisotopes | maximum number of isotopes shown in the resulting molecules                     |

### Details

addMolecules() adds the second argument to the first. subMolecules() subtracts the second argument from the first.

This can be useful to revert e.g. adduct/fragment formation found in ESI mass spectrometry, or to mimick simple chemical reactions. No chemical checks are performed.

### Value

A list with the elements

|          |                                     |
|----------|-------------------------------------|
| formula  | repeated sum formula                |
| mass     | exact monoisotopic mass of molecule |
| score    | dummy value, always 1.0             |
| isotopes | a list of isotopes                  |

### Author(s)

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## Examples

```
# For proton-Adduct of Ethanol:  
subMolecules("C2H7O", "H")
```

---

decomposeIsotopes      *Mass Decomposition of Isotope Patterns*

---

## Description

Calculate the elementary compositions from an exact Mass or Isotope Pattern, obtained e.g.\ by FTICR or TOF mass spectrometers

## Usage

```
decomposeMass(mass, ppm=2.0, mzabs=0.0001, elements=NULL, filter=NULL,  
z=0, maxisotopes = 10, minElements="C0", maxElements="C999999")  
decomposeIsotopes(masses, intensities, ppm=2.0, mzabs=0.0001,  
elements=NULL, filter=NULL, z=0, maxisotopes = 10, minElements="C0", maxElements="C999999")  
isotopeScore(molecule, masses, intensities, elements = NULL, filter = NULL, z = 0)
```

## Arguments

|                          |  |
|--------------------------|--|
| mass                     | A single exact mass (or m/z value)   |
| masses                   | A vector of masses (or m/z values) of an isotope cluster                                     |
| intensities              | Abolute or relative intensities of the masses peaks  |
| ppm                      | allowed deviation of hypotheses from given mass  |
| mzabs                    | absolute deviation in dalton (mzabs and ppm will be added)                                   |
| z                        | charge z of m/z peaks for calculation of real mass. 0 is for auto-detection                  |
| maxisotopes              | maximum number of isotopes shown in the resulting molecules                                  |
| elements                 | list of allowed chemical elements, defaults to CHNOPS  |
| minElements, maxElements | Molecular formulas, which contain lower and upper boundaries of allowed formula respectively |
| filter                   | NYI, will be a selection of DU, DBE and Nitrogen rules                                       |
| molecule                 | a molecule as obtained from getMolecule() or decomposeMass / decomposeIsotopes               |

## Details

Sum formulas are calculated which explain the given mass or isotope pattern.

**Value**

A list of molecules, which contain the sub-lists

|                       |                                       |
|-----------------------|---------------------------------------|
| <code>formula</code>  | potential formulae                    |
| <code>mass</code>     | exact monoisotopic mass of hypothesis |
| <code>score</code>    | calculated score                      |
| <code>isotopes</code> | a list of isotopes                    |

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**References**

For a description of the underlying IMS see: see citation("Rdisop")

**See Also**

[decomposeMass](#)

**Examples**

```
# For Glutamate:  
decomposeIsotopes(c(147.0529, 148.0563), c(100.0, 5.561173))
```

`getMolecule`

*Calculate mass and isotope information for a molecule given as sum formula*

**Description**

Parse the sum formula and calculate the theoretical exact mass and the isotope distribution.

**Usage**

```
getMolecule(formula, elements = NULL, z = 0, maxisotopes = 10)
getMass(molecule)
getFormula(molecule)
getIsotope(molecule, index)
getScore(molecule)
getValid(molecule)
```

**Arguments**

|             |  |
|-------------|--|
| formula     | Sum formula  |
| elements    | list of allowed chemical elements, defaults to full periodic system of elements                              |
| z           | charge z of molecule for exact mass calculation  |
| maxisotopes | maximum number of isotopes shown in the resulting molecules  |
| molecule    | an initialized molecule as returned by getMolecule() or the decomposeMass() and decomposeIsotope() functions |
| index       | return the n-th isotope mass/abundance pair of the molecule  |

**Details**

getMolecule() Parse the sum formula and calculate the theoretical exact monoisotopic mass and the isotope distribution. For a given element, return the different mass values.

**Value**

getMolecule: A list with the elements

|          |  |
|----------|--|
| formula  | repeated sum formula   |
| mass     | exact monoisotopic mass of molecule                                |
| score    | probability, for given molecules a dummy value which is always 1.0 |
| valid    | result of neutrogen rule check                                     |
| isotopes | a list of isotopes   |

getMass, getFormula and getScore: return the mass of the molecule as string or real value

**Author(s)**

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**References**

For a description of the underlying IMS see: see citation("Rdisop")

**Examples**

```
# For Ethanol:  
getMolecule("C2H6O")
```

**initializeCHNOPS***Initialize (a subset of) elements of the periodic system of elements (PSE)***Description**

Initialize the information about name, mass and isotopes. To reduce the number of decomposition hypotheses, subsets of elements can be created.

**Usage**

```
initializeCHNOPS()
initializeCHNOPSMgKCaFe()
initializePSE()
initializeElements(names)
```

**Arguments**

|       |                                    |
|-------|------------------------------------|
| names | vector of element names within PSE |
|-------|------------------------------------|

**Details**

These functions return full, pre-defined or user-defined (sub-) lists of elements.

**Value**

A list with the elements

|         |                          |
|---------|--------------------------|
| name    | repeated sum formula     |
| mass    | nominal mass of molecule |
| isotope | a list of isotopes       |

The initializeCharges() is special, since it allows to parse charges such as getMolecule("H3O+", elements=c(initializeCHNOPS(), "H3O+"))

**Author(s)**

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**References**

For a description of the underlying IMS see: see citation("Rdisop")

Isotope patterns obtained through wikipedia.org

**See Also**

[getMolecule](#)

*initializeCHNOPS*

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**Examples**

```
# For Ethanol:  
elements <- initializeCHNOPS()
```

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