

Rdisop

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RcppVersion

Rcpp Version and License Information

Description

RcppVersion displays the version of Rcpp/RcppTemplate that was used to build this package.

Usage

```
RcppVersion()
```

Author(s)

Dominick Samperi

Examples

```
RcppVersion()
```

addMolecules

Add/subtract sum formulae

Description

Simple arithmetic modifications of sum formulae.

Usage

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

Arguments

formula1 Sum formula

formula2 Sum formula

elements list of allowed chemical elements, defaults to full periodic system of elements

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)

Steffen Neumann <sneumann@IPB-Halle.DE>

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, maxi  
decomposeIsotopes(masses, intensities, ppm=2.0, mzabs=0.0001,  
elements=NULL, filter=NULL, z=0, maxisotopes = 10)  
isotopeScore(molecule, masses, intensities, elements = NULL, filter = NULL, z =
```

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 |
| 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

| | |
|----------|---------------------------------------|
| formula | potential formulae |
| mass | exact monoisotopic mass of hypothesis |
| score | calculated score |
| isotopes | a list of isotopes |

Author(s)

Steffen Neumann <sneumann@IPB-Halle.DE>

References

For a description of the underlying IMS see: WABI Paper

See Also

[decomposeMass](#)

Examples

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

| | |
|-------------|---|
| getMolecule | <i>Calculate mass and isotope information for a molecule given as sum</i> |
|-------------|---|

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

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

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

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

Author(s)

Steffen Neumann <sneumann@IPB-Halle.DE>

References

For a description of the underlying IMS see: WABI Paper

Examples

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

```
initializeCHNOPS      Initialize (a subset of) elements of the periodic system of elements
```

Description

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

Usage

```
initializeCHNOPS()  
initializeCHNOPS.MgKCaFe()  
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(), initializeCharges()))

Author(s)

Steffen Neumann <sneumann@IPB-Halle.DE>

References

For a description of the underlying IMS see: WABI Paper

Isotope patterns obtained through wikipedia.org

See Also

[getMolecule](#)

Examples

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

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