Package ‘rcdk’

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LazyLoad yes
Description Allows the user to access functionality in the ‘CDK’, a Java framework for chemoinformatics. This allows the user to load molecules, evaluate fingerprints, calculate molecular descriptors and so on. In addition, the ‘CDK’ API allows the user to view structures in 2D.
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**Atoms**

**Operations on atoms**

**Description**

- `getSymbol(atom)` returns the chemical symbol for an atom.
- `getPoint3D(atom)` returns the 3D coordinates of the atom.
- `getPoint2D(atom)` returns the 2D coordinates of the atom.
- `getAtomicNumber(atom)` returns the atomic number of the atom.
- `getHydrogenCount(atom)` returns the number of implicit H’s on the atom. Depending on where the molecule was read from this may be NULL or an integer greater than or equal to 0.
- `getCharge(atom)` returns the partial charge on the atom. If charges have not been set the return value is NULL, otherwise the appropriate charge.
- `getFormalCharge(atom)` returns the formal charge on the atom. By default the formal charge will be 0 (i.e., NULL is never returned).
- `isAromatic(atom)` returns TRUE if the atom is aromatic, FALSE otherwise.
- `isAliphatic(atom)` returns TRUE if the atom is part of an aliphatic chain, FALSE otherwise.
- `isInRing(atom)` returns TRUE if the atom is in a ring, FALSE otherwise.
- `getAtomIndex(atom, mol)` returns the index of the atom in the molecule (starting from 0).
- `getConnectedAtoms(atom, mol)` returns a list of atoms that are connected to the specified atom.

**Usage**

```java
getSymbol(atom)
gpoint3D(atom)
gpoint2D(atom)
gatomicNumber(atom)
ghydrogenCount(atom)
gcharge(atom)
gformalCharge(atom)
gconnectedAtoms(atom, mol)
gatomIndex(atom, mol)
isAromatic(atom)
isAliphatic(atom)
isInRing(atom)
```

**Arguments**

- `atom` A `jobjRef` representing an `IAtom` object.
- `mol` A `jobjRef` representing an `IAtomContainer` object.
Value

In the case of get.point3d the return value is a 3-element vector containing the X, Y and Z coordinates of the atom. If the atom does not have 3D coordinates, it returns a vector of the form c(NA, NA, NA). Similarly for get.point2d, in which case the return vector is of length 2.

Author(s)

Rajarshi Guha (<rajarshi.guha@gmail.com>)

See Also

get.atoms

bpdata

Boiling Point Data

Description

Structures and associated boiling points for 277 molecules, primarily alkanes and substituted alkanes.

Usage

bpdata

Format

A data.frame with two columns:

<table>
<thead>
<tr>
<th></th>
<th>SMILES</th>
<th>BP</th>
</tr>
</thead>
<tbody>
<tr>
<td>.1</td>
<td>character</td>
<td>Structure in SMILES format</td>
</tr>
<tr>
<td>.2</td>
<td>numeric</td>
<td>Boiling point in Kelvin</td>
</tr>
</tbody>
</table>

The names of the molecules are used as the row names

References


cdk.version

Get Current CDK Version

Description

Returns a string containing the version of the CDK used in this package
cdkFormula-class

Usage

  cdk.version()

Value

  A string representing the CDK version

Author(s)

  Rajarshi Guha (<rajarshi.guha@gmail.com>)

---

Class cdkFormula, a class for handling molecular formula

Description

  This class handles molecular formulae. It provides extra information such as the IMolecularFormula Java object, elements contained and number of them.

Objects from the Class

  Objects can be created using new constructor and filled with a specific mass and window accuracy.

Note

  No notes yet.

Author(s)

  Miguel Rojas-Cherto (<miguelrojasch@yahoo.es>)

References

  A parallel effort to expand the Chemistry Development Kit: http://cdk.sourceforge.net

See Also

  get.formula, set.charge.formula, get.isotopes.pattern, isvalid.formula,
compare.isotope.pattern

Compare isotope patterns.

Description

Computes a similarity score between two different isotope abundance patterns.

Usage

compare.isotope.pattern(iso1, iso2, ips = NULL)

Arguments

iso1 The first isotope pattern, which should be a jobjRef corresponding to the IsotopePattern class
iso2 The second isotope pattern, which should be a jobjRef corresponding to the IsotopePattern class
ips An instance of the IsotopePatternSimilarity class. If NULL one will be constructed automatically

Value

A numeric value between 0 and 1 indicating the similarity between the two patterns

Author(s)

Miguel Rojas Cherto

References

http://cdk.github.io/cdk/2.0/docs/api/org/openscience/cdk/formula/IsotopePatternSimilarity.html

See Also

generate.isotope.pattern.similarity
do.aromaticity

Perform Aromaticity Detection, atom typing or isotopic configuration

Description

These methods can be used to perform aromaticity detection, atom typing or isotopic configuration on a molecule object. In general, when molecules are loaded via `load.molecules` these are performed by default. If molecules are obtained via `parse.smiles` these operations are not performed and so the user should call one or both of these methods to correctly configure a molecule.

Usage

do.aromaticity(molecule)
do.typing(molecule)
do.isotopes(molecule)

Arguments

molecule The molecule on which the operation is to be performed. Should of class `jobjref` with a `jclass` attribute of `IAtomContainer`

Value

No return value. If the operations fail an exception is thrown and an error message is printed

Author(s)

Rajarshi Guha (<rajarshi.guha@gmail.com>)

See Also

`load.molecules, parse.smiles`

-------

eval.atomic.desc Evaluate an Atomic Descriptor

Description

The CDK implements a number of descriptors divided into three main groups - atomic, molecular and bond. This method evaluates the specified atomic descriptor(s) for a molecule

Usage

eval.atomic.desc(molecule, which.desc, verbose=FALSE)
eval.desc

Arguments

molecule A reference to a CDK IAtomContainer object
which.desc The fully qualified class name of the descriptor to evaluate or a vector such names
verbose If TRUE, progress will be written to the screen, otherwise the function performs silently

Value

A data.frame is returned.

Author(s)

Rajarshi Guha (<rajarshi.guha@gmail.com>)

See Also

g.get.atomic.desc.names get.desc.names eval.desc

---

**eval.desc**

*Evaluate a Molecular Descriptor*

Description

The CDK implements a number of descriptors divided into three main groups - atomic, molecular and bond. This method evaluates the specified molecular descriptor(s) for a molecule.

Usage

eval.desc(molecules, which.desc, verbose=FALSE)

Arguments

molecules A single IAtomContainer object or a list of references to CDK IAtomContainer objects
which.desc The fully qualified class name of the descriptor to evaluate or a vector such names
verbose If TRUE, progress will be written to the screen, otherwise the function performs silently

Value

A data.frame is returned. For a single molecule it will have one row, for multiple molecules it will have the number of rows equal to the number of molecules.
**generate.2d.coordinates**

**Author(s)**
Rajarshi Guha (<rajarshi.guha@gmail.com>)

**See Also**
get.desc.names get.desc.categories

**Examples**

```r
smiles <- c('CCC', 'c1cccccl', 'CC(=O)C')
mols <- sapply(smiles, parse.smiles)
dnames <- get.desc.names('constitutional')
descs <- eval.desc(mols, dnames, verbose=TRUE)
```

---

**Description**

This function will generate reasonable 2D coordinates based purely on connectivity information.

**Usage**

```r
generate.2d.coordinates(molecule)
```

**Arguments**

- **molecule** An IAtomContainer object that can be obtained by loading them from disk or drawing them in the editor.

**Value**

Returns the input molecule with 2D coordinates added.

**Author(s)**
Rajarshi Guha (<rajarshi.guha@gmail.com>)
generate.formula  

Generate a cdkFormula object.

Description

This function generate a list of cdkFormula objects given a mass.

Usage

```r
generate.formula(mass, window=0.01, elements=list(c("C",0,50),c("H",0,50),
          c("N",0,50),c("O",0,50),
          c("S",0,50)),
          validation=FALSE, charge=0.0)
```

Arguments

- **mass**: The mass value from which to be generate the formulas.
- **window**: The window accuracy in the same units as mass.
- **elements**: Elements to take into account.
- **validation**: TRUE, if the method should only generate valid formulas. If FALSE, nonsensical formulae may be generated which must be filtered out by the user.
- **charge**: The charge value of the formula.

Value

Objects of class MassToFormulaTool, from the IMolecularFormula package

Author(s)

Miguel Rojas-Cherto (<miguelrojasch@yahoo.es>)

See Also

generate.formula, set.charge.formula, get.isotopes.pattern, isValid.formula

Examples

```r
mfSet <- generate.formula(18.03383,charge=1,
          elements=list(c("C",0,50),c("H",0,50),c("N",0,50)))
for (i in mfSet) {
    print(i)
}
```
**get.atomic.desc.names**  
*Get the names of the available atomic descriptors*

**Description**

The CDK implements a number of descriptors divided into three main groups - atomic, molecular and bond. This method returns the names of the available atomic descriptors.

**Usage**

```r
get.atomic.desc.names(type = "all")
```

**Arguments**

- **type**  
  A string which can be one of "all", "topological", "geometrical" "hybrid", "constitutional", "electronic", allowing you to choose atomic descriptors of specific categories. The keyword "all" will return all available descriptors.

**Value**

A vector of fully qualified descriptor names.

**Author(s)**

Rajarshi Guha (<rajarshi.guha@gmail.com>)

**See Also**

- eval.atomic.desc  
- get.desc.names  
- eval.desc

---

**get.atoms**  
*Get the atoms from a molecule or bond*

**Description**

This function returns a list containing IAtom objects from a molecule or a bond.

**Usage**

```r
get.atoms(object)  
get.atom.count(molecule)
```

**Arguments**

- **object**  
  A jObjRef representing an IAtomContainer, IMolecule or IBond object

- **molecule**  
  A jObjRef representing an IAtomContainer
get.bonds

Value
A list containing jobjRef's to a CDK IAtom object or else the number of atoms in the molecule

Author(s)
Rajarshi Guha (<rajarshi.guha@gmail.com>)

See Also
getNbonds, getNpoints, getNsymbols

get.bonds
Get the bonds from a molecule

description
This function returns a list containing IBond objects from a molecule

Usage
get.bonds(molecule)

Arguments
molecule A jobjRef representing an IAtomContainer, IMolecule

Value
A list containing jobjRef's to a CDK IBond object

Author(s)
Rajarshi Guha (<rajarshi.guha@gmail.com>)

See Also
get.atoms, get.connected.atom,
get.connected.atom     

Derivation

This function returns the atom that is connected to a specified in a specified bond. Note that this function assumes 2-atom bonds, mainly because the CDK does not currently support other types of bonds.

Usage

get.connected.atom(bond, atom)

Arguments

- bond: A jObject representing an IBond object
- atom: A jObject representing an IAtom object

Value

A jObject representing an IAtom object

Author(s)

Rajarshi Guha (<rajarshi.guha@gmail.com>)

See Also

get.atoms

get.desc.categories     

Description

This function returns the broad descriptor categories that are available. Examples include topological, geometrical and so on. You can use a specific category to avoid calculating all descriptors for a set of molecules and saves you having to select individual descriptors by hand.

Usage

get.desc.categories()

Value

A character vector of descriptor category names
get.desc.names

Author(s)

Rajarshi Guha (<rajarshi.guha@gmail.com>)

See Also

eval.desc, get.desc.names

get.desc.names  Get Descriptor Class Names

Description

The CDK implements a number of descriptors divided into three main groups - atomic, molecular and bond. Currently the package will only evaluate molecular descriptors. This function returns the class names of the available descriptors, which can then be used to calculate descriptors for a specific molecule.

By default all available descriptor class names are returned. However it is possible to specify that a subset of the descriptors should be considered. The subset is specified by keyword and can be one of: topological, geometrical, hybrid, constitutional, protein, electronic.

Usage

get.desc.names(type = "all")

Arguments

type  Indicates which subset of molecular descriptors should be considered

Value

A character vector of descriptor class names

Author(s)

Rajarshi Guha (<rajarshi.guha@gmail.com>)

See Also

eval.desc, get.desc.categories
get.fingerprint

Evaluate Fingerprints

Description

This function evaluates fingerprints of a specified type for a set of molecules or a single molecule. Depending on the nature of the fingerprint, parameters can be specified. Currently five different fingerprints can be specified:

- **standard** - Considers paths of a given length. The default is but can be changed. These are hashed fingerprints, with a default length of 1024
- **extended** - Similar to the standard type, but takes rings and atomic properties into account
- **graph** - Similar to the standard type by simply considers connectivity
- **hybridization** - Similar to the standard type, but only consider hybridization state
- **macs** - The popular 166 bit MACCS keys described by MDL
- **estate** - 79 bit fingerprints corresponding to the E-State atom types described by Hall and Kier
- **pubchem** - 881 bit fingerprints defined by PubChem
- **kr** - 4860 bit fingerprint defined by Klekota and Roth
- **shortestpath** - A fingerprint based on the shortest paths between pairs of atoms and takes into account ring systems, charges etc.
- **signature** - A feature,count type of fingerprint, similar in nature to circular fingerprints, but based on the signature descriptor
- **circular** - An implementation of the ECFP6 fingerprint

Depending on whether the input is a single IAtomContainer object, a list or single vector is returned. Each element of the list is an S4 object of class fingerprint-class or featvec-class, which can be manipulated with the fingerprint package.

Usage

```
get.fingerprint(molecule, type = 'standard',
                fp.mode = 'bit', depth=6, size=1024, verbose=FALSE)
```

Arguments

- **molecule**
  An IAtomContainer object that can be obtained by loading them from disk or drawing them in the editor.
- **type**
  The type of fingerprint. See description for possible values. The default is the standard binary fingerprint.
- **fp.mode**
  The type of fingerprint to return. Possible values are 'bit', 'raw', and 'count'. The 'raw' mode will return a featvec-class type of fingerprint, representing fragments and their count of occurrence in the molecule. The 'count' mode is
similar, except that it returns hash values of fragments and their count of occurrence. While any of these values can be specified, a given fingerprint implementation may not implement all of them, and in those cases the return value is NULL.

**depth**
The search depth. This argument is ignored for the 'pubchem', 'maccs', 'kr' and 'estate' fingerprints

**size**
The length of the fingerprint bit string. This argument is ignored for the 'pubchem', 'maccs', 'kr', 'signature', 'circular' and 'estate' fingerprints

**verbose**
If TRUE, exceptions, if they occur, will be printed

**Value**
Objects of class `fingerprint-class` or `featvec-class`, from the `fingerprint` package. If there is a problem during fingerprint calculation, NULL is returned.

**Author(s)**
Rajarshi Guha (<rajarshi.guha@gmail.com>)

**References**

**See Also**
`load.molecules`

**Examples**

```r
## get some molecules
sp <- get.smiles.parser()
smiles <- c('CCC', 'CCN', 'CCN(C)(C)', 'c1cccccccccccccccccc', 'C1CCCC1CC(CN(C)(C))CC(=O)CC')
mols <- parse.smiles(smiles)

## get a single fingerprint using the standard
## (hashed, path based) fingerprinter
fp <- get.fingerprint(mols[[1]])

## get MACCS keys for all the molecules
fps <- lapply(mols, get.fingerprint, type='maccs')

## get Signature fingerprint
## feature, count fingerprinter
fps <- lapply(mols, get.fingerprint, type='signature', fp.mode='raw')
```
get.formula

Get the formula object from a formula character.

Description

This function returns a formula object containing mass, string character and isotopes when is given a character/string formula.

Usage

get.formula(mf, charge=0)

Arguments

mf A string containing the formula of the molecular formula of chemical object.

charge The charge of the molecular formula.

Value

Objects of class cdkFormula, from the IMolecularFormula package

Author(s)

Miguel Rojas-Cherto (<miguelrojasch@yahoo.es>)

References

A parallel effort to expand the Chemistry Development Kit: http://cdk.sourceforge.net

See Also

set.charge.formula, get.isotopes.pattern, isvalid.formula, generate.formula

Examples

formula <- get.formula('NH4', charge = 1)

formula
get.isotope.pattern.generator

*Construct an isotope pattern generator.*

**Description**

Constructs an instance of the CDK IsotopePatternGenerator, with an optional minimum abundance specified. This object can be used to generate all combinatorial chemical isotopes given a structure.

**Usage**

```java
get.isotope.pattern.generator(minAbundance = NULL)
```

**Arguments**

- `minAbundance`: The minimum abundance

**Value**

A jobRef corresponding to an instance of IsotopePatternGenerator

**Author(s)**

Miguel Rojas Cherto

**References**

http://cdk.github.io/cdk/1.5/docs/api/org/openscience/cdk/formula/IsotopePatternGenerator.html

get.isotope.pattern.similarity

*Construct an isotope pattern similarity calculator.*

**Description**

A method that returns an instance of the CDK IsotopePatternSimilarity class which can be used to compute similarity scores between pairs of isotope abundance patterns.

**Usage**

```java
get.isotope.pattern.similarity(tol = NULL)
```

**Arguments**

- `tol`: The tolerance
get.isotopes.pattern

Value
A jobbRef corresponding to an instance of IsotopePatternSimilarity

Author(s)
Miguel Rojas Cherto

References
http://cdk.github.io/cdk/1.5/docs/api/org/openscience/cdk/formula/IsotopePatternSimilarity.html

See Also
compare.isotope.pattern

generate.isotopes

Generate the isotope pattern.

Description
This function get the isotope pattern given a cdkFormula object. It modifies as the IMolecularFormula Java object as the its mass.

Usage
get.isotopes.pattern(formula,minAbund=0.1)

Arguments
formula A cdkFormula object.
minAbund Minimal abundance of the isotopes to be added in the combinatorial search.

Value
Objects of class IsotopePatternGenerator, from the IMolecularFormula package

Author(s)
Miguel Rojas-Cherto (<miguelrojasch@yahoo.es>)

References
A parallel effort to expand the Chemistry Development Kit: http://cdk.sourceforge.net

See Also
get.formula, set.charge.formula, isValid.formula, generate.formula
get.mol2formula  Parser a molecule to formula object.

Description
This function converts a molecule object to a formula object.

Usage
get.mol2formula(molecule, charge=0)

Arguments
molecule  The molecule to be parsed.
charge    The charge characterizing the molecule.

Value
Objects of class MolecularFormulaManipulator, from the IMolecularFormulaManipulator package

Author(s)
Miguel Rojas-Cherto (<miguelrojasch@yahoo.es>)

See Also
set.charge.formula, get.isotopes.pattern, isValid.formula

Examples
molecule <- parse.smiles("N")[1]
convert.implicit.to.explicit(molecule)
formula <- get.mol2formula(molecule, charge=0)

get.murcko.fragments  Molecule Fragmentation Methods

Description
A variety of methods for fragmenting molecules are available ranging from exhaustive, rings to
more specific methods such as Murcko frameworks. Fragmenting a collection of molecules can be
a useful for a variety of analyses. In addition fragment based analysis can be a useful and faster
alternative to traditional clustering of the whole collection, especially when it is large.
Note that exhaustive fragmentation of large molecules (with many single bonds) can become time
consuming.
Usage

get.murcko.fragments(mols, min.frag.size = 6, as.smiles = TRUE, single.framework = FALSE)
get.exhaustive.fragments(mols, min.frag.size = 6, as.smiles = TRUE)

Arguments

mols A molecule object or list of molecule objects. Each object should have a jclass of IAtomContainer
min.frag.size The size of the smallest fragments to be considered
as.smiles If TRUE, the fragments are returned as SMILES strings, otherwise as IAtomContainer objects
single.framework If TRUE, then a single framework (i.e., the framework consisting of the union of all ring systems and linkers) is returned for each molecule. Otherwise, all combinations of ring systems and linkers are returned

Value

get.murcko.fragments returns a list with each element being a list with two elements: rings and frameworks. Each of these elements is either a character vector of SMILES strings or a list of IAtomContainer objects. get.exhaustive.fragments returns a list of length equal to the number of input molecules. Each element is a character vector of SMILES strings or a list of IAtomContainer objects.

Author(s)

Rajarshi Guha (<rajarshi.guha@gmail.com>)

See Also

load.molecules, parse.smiles,

Examples

mol <- parse.smiles('c1ccc(c1)CN(c2cc(ccc2[N+](=O)[O-])c3c(nc(nc3CCN)N)C')[[1]]
mfl <- get.murcko.fragments(mol, as.smiles=TRUE, single.framework=TRUE)
mfl <- get.murcko.fragments(mol, as.smiles=TRUE, single.framework=FALSE)

get.properties Get All Property Values of a Molecule

Description

Returns a list of all the properties of a molecule. The names of the list are set to the property names
get.property

Usage
get.properties(molecule)

Arguments
molecule A Java object of class IAtomContainer or IMolecule

Value
A list of the property values, with names equal to the property names. NULL property values are returned as NA

Author(s)
Rajarshi Guha (<rajarshi.guha@gmail.com>)

See Also
get.property, set.property, remove.property

Examples
smiles <- 'c1ccccc1'
mol <- parse.smiles(smiles)[[1]]
set.property(mol, 'prop1', 23.45)
set.property(mol, 'prop2', 'inactive')
get.properties(mol)

get.property Get the Value of a Molecule Property

Description
This function retrieves the value of a keyed property that has previously been set on the molecule. The get.title function is simply a wrapper around get.property that directly provides access to the molecule title.

Usage
get.property(molecule, key)
get.title(molecule)

Arguments
molecule A Java object of class IAtomContainer
key A string naming the property
get.smiles

Value

The value of the property is the key is found else NA. For get.title, the title of the molecule if available otherwise NA

Author(s)

Rajarshi Guha (<rajarshi.guha@gmail.com>)

See Also

get.properties, set.property, remove.property

Examples

```r
smiles <- 'c1ccccc1'
mol <- parse.smiles(smiles)[[1]]
set.property(mol, 'prop1', 23.45)
set.property(mol, 'prop2', 'inactive')
get.property(mol, 'prop1')
```

get.smiles  

Get the SMILES for a Molecule

Description

The function will generate a SMILES representation of an IAtomContainer object. The default parameters of the CDK SMILES generator are used. This can mean that for large ring systems the method may fail. See CDK Javadocs for more information

Usage

```r
get.smiles(molecule, flavor = smiles.flavors(c('Generic')), smigen = NULL)
```

Arguments

- molecule: A Java object of class IAtomContainer
- flavor: Customizations for SMILES generation. See smiles.flavors
- smigen: An instance of SmilesGenerator, which can be useful if you are generating SMILES for a large number of molecules

Value

An R character object containing the SMILES

Author(s)

Rajarshi Guha (<rajarshi.guha@gmail.com>)
See Also

`smiles.flavors, parse.smiles`

Examples

```r
m <- parse.smiles('C1C=CCC1N(C)c1ccccc1')[[1]]
gem.smiles(m)
gem.smiles(m, smiles.flavors(c('Generic','UseAromaticSymbols')))```

get.smiles.parser  
*Get a SMILES Parser*

Description

This function returns a reference to a SMILES parser object. If you are parsing multiple SMILES strings, it is preferable to create your own parser and supply it to `parse.smiles` rather than forcing that function to instantiate a new parser for each call.

Usage

```r
gem.smiles.parser()```

Value

A `jobjRef` to a CDK SmilesParser object

Author(s)

Rajarshi Guha (<rajarshi.guha@gmail.com>)

See Also

`get.smiles, get.smiles.parser, view.molecule.2d`

get.total.charge  
*Get the Total Charges for the Molecule*

Description

gem.total.charge returns the summed partial charges for a molecule and `get.total.formal.charge` returns the summed formal charges. Currently, if one or more partial charges are unset, the function simply returns the sum of formal charges (via `get.total.formal.charge`). This is slightly different from how the CDK evaluates the total charge of a molecule (via `AtomContainerManipulator.getTotalCharge()`), but is in line with how OEChem determines net charge on a molecule.

In general, you will want to use the `get.total.charge` function.
**get.total.hydrogen.count**

**Usage**

```java
get.total.charge(molecule)
get.total.formal.charge(molecule)
```

**Arguments**

- `molecule` A Java object of class `IAAtomContainer`

**Value**

A double value indicating the total partial charge or total formal charge

**Author(s)**

Rajarshi Guha (<rajarshi.guha@gmail.com>)

---

**get.total.hydrogen.count**

*Get the Total Hydrogen Count for a Molecule*

**Description**

The function will return the summed implicit hydrogens of all atoms in the specified AtomContainer

**Usage**

```java
get.total.hydrogen.count(molecule)
```

**Arguments**

- `molecule` A Java object of class `IAAtomContainer`

**Value**

An integer value indicating the number of implicit hydrogens

**Author(s)**

Rajarshi Guha (<rajarshi.guha@gmail.com>)
Description

These methods will return the value for the corresponding descriptors. While they can always be evaluated using `eval.desc`, they are common enough that separate functions are provided.

Usage

```r
get.tpsa(molecule)
get.alogp(molecule)
get.xlogp(molecule)
get.volume(molecule)
```

Arguments

- `molecule`: A `jobjRef` representing an `IAtomContainer` object

Details

It's important to note that ALogP and XLogP assumes that the molecule has explicit hydrogens. If the molecule is read from an SD file, explicit H's are usually present. On the other hand, if the molecule is obtained from a SMILES, explicit hydrogens must be added.

The molecular volume is calculated using a group contribution method rather than the an analytical method. This allows to avoid the use of 3D structures.

Value

Single numeric value representing TPSA, ALogP, XLogP or molecular volume.

Author(s)

Rajarshi Guha (<rajarshi.guha@gmail.com>)

See Also

`eval.desc`
**hasNext**

*Does This Iterator Have A Next Element*

**Description**

`hasNext` is a generic function that indicates if the iterator has another element.

**Usage**

```r
class 'iload.molecules'
hasNext(obj, ...) # S3 method for class 'iload.molecules'
```

**Arguments**

- `obj`: an iterator object.
- `...`: additional arguments that are ignored.

**Value**

Logical value indicating whether the iterator has a next element. In the context of reading a structure file, this indicates whether there are more molecules to read.

**See Also**

- `iload.molecules`

---

**is.connected**

*Get the Largest Component in a Disconnected Molecule*

**Description**

These methods allow one to check whether a molecule is fully connected or else retrieve the largest disconnected component.

**Usage**

```r
class 'iload.molecules'
gc.
class 'iload.molecules'
```

**Arguments**

- `mol`: A jObjectRef representing an IAtomContainer object.
Value

For `get.largest.component`, if the input molecule has more than one disconnected component, the largest is returned. Otherwise, the molecule itself is returned.

For `is.connected`, `TRUE` if the molecule is fully connected, `FALSE` otherwise.

Author(s)

Rajarshi Guha (<rajarshi.guha@gmail.com>)

Examples

```r
m <- parse.smiles("CCCCCCCC")[[1]]
largest <- get.largest.component(m)
length(get.atoms(largest)) == 6
```

isvalid.formula Validate a cdkFormula object.

Description

This function validates a `cdkFormula` object. At the moment is using the nitrogen Rule and RDBE Rule.

Usage

```r
isvalid.formula(formula, rule = c("nitrogen","RDBE"))
```

Arguments

- `formula`: A `cdkFormula` object.
- `rule`: The rules to be applied: nitrogen and RDBE.

Value

Objects of class `MolecularFormulaChecker`, from the `IMolecularFormula` package

Author(s)

Miguel Rojas-Cherto (<miguelrojasch@yahoo.es>)

References

A parallel effort to expand the Chemistry Development Kit: [http://cdk.sourceforge.net](http://cdk.sourceforge.net)

See Also

generate.formula, get.isotopes.pattern, get.formula
Examples

```r
formula <- get.formula('NH4', charge = 0)
isvalid.formula(formula, rule = c("nitrogen","RDBE"))
```

Description

The CDK can read a variety of molecular structure formats. This function encapsulates the calls to the CDK API to load a structure given its filename.

Usage

```r
load.molecules(molfiles=NA, aromaticity = TRUE, typing = TRUE, isotopes = TRUE, verbose=FALSE)
iload.molecules(molfile, type="smi", aromaticity = TRUE, typing = TRUE, isotopes = TRUE, skip=TRUE)
```

Arguments

- `molfiles` A character vector of filenames. Note that the full path to the files should be provided. URL’s can also be used as paths. In such a case, the URL should start with "http://"
- `molfile` A string containing the filename to load. Must be a local file
- `type` Indicates whether the input file is SMILES or SDF. Valid values are "smi" or "sdf"
- `aromaticity` If TRUE then aromaticity detection is performed on all loaded molecules. If this fails for a given molecule, then the molecule is set to NA in the return list
- `typing` If TRUE then atom typing is performed on all loaded molecules. The assigned types will be CDK internal types. If this fails for a given molecule, then the molecule is set to NA in the return list
- `isotopes` If TRUE then atoms are configured with isotopic masses
- `verbose` If TRUE, output (such as file download progress) will be bountiful
- `skip` If TRUE, then the reader will continue reading even when faced with an invalid molecule. If FALSE, the reader will stop at the first invalid molecule

Details

Note that if molecules are read in from formats that do not have rules for handling implicit hydrogens (such as MDL MOL), the molecule will not have implicit or explicit hydrogens. To add explicit hydrogens, make sure that the molecule has been typed (this is TRUE by default for this function) and then call `convert.implicit.to.explicit`. On the other hand for a format such as SMILES, implicit or explicit hydrogens will be present.
Value
load.molecules returns a list of CDK Molecule objects, which can be used in other rcdk functions.

iload.molecules is an iterating version of the loader and is applicable for large SMILES or SDF files. In contrast to load.molecules this does not load all the molecules into memory at one go, and as a result lets you process arbitrarily large structure files.

Author(s)
Rajarshi Guha (<rajarshi.guha@gmail.com>)

See Also
view.molecule.2d, convert.implicit.to.explicit

Examples
## Not run:

```r
## load a single file
amol <- load.molecules('foo.sdf')

## load multiple files
mols <- load.molecules(c('mol1.sdf', 'mol2.smi',

## iterate over a large file
moliter <- iload.molecules("big.sdf", type="sdf")
while(hasNext(moliter)) {
  mol <- nextElem(moliter)
  print(get.property(mol, "cdk:Title"))
}

## End(Not run)
```

Description
These functions perform substructure searches of a query, specified in SMILES or SMARTS forms, over one or more target molecules and maximum common substructure searches for pairs of molecules.

Usage

```r
matches(query, target, return.matches=FALSE)
is.subgraph(query, target)
get.mcs(mol1, mol2, as.molecule = TRUE)
```
matches

Arguments

query          A SMILES or SMARTS string
target         A single IAtomContainer object or a list of IAtomContainer objects
mol1           An IAtomContainer
mol2           An IAtomContainer
return.matches If TRUE the lists of atom indices that correspond to the matching substructure are returned
as.molecule    If TRUE the MCS is returned as a new IAtomContainer object. Otherwise a atom index mapping between the two molecules is returned as a 2D array of integers

Details

For the case of isNsubgraph, the query molecule must be a single IAtomContainer or a valid SMILES string. Note that this method can be significantly faster than matches, but is limited by the fact that SMARTS patterns cannot be specified. This uses the "TurboSubStructure" SMSD method and so only searches for the first substructure match.

For MCS detection, the default SMSD algorithm is employed and the best scoring MCS is returned by default. Furthermore, one can obtain the resultant MCS either as an IAtomContainer in which the atoms and bonds are clones of the corresponding matching atoms and bonds in one of the molecule. Or else as a 2D array of dimensions N×2 of atom index mappings. Here N is the size of the MCS and the first column represents the atom index from the first molecule and the second column the atom index from the second molecule.

Note that since the CDK SMARTS matcher internally will perform aromaticity perception and atom typing, the target molecules need not have these operations done on them beforehand for matches method. However, if isNsubgraph or get.mcs is being used, the molecules should have aromaticity detected and atom typing performed explicitly.

If the atom indices of the matching substructures (in the target molecule) are desired, use the matches function directly.

Value

For matches with return.matches = FALSE, a boolean vector where each element is TRUE or FALSE depending on whether the corresponding element in targets contains the query or not. If return.matches = TRUE, the return value is a list of lists. The number of elements of the top level list equals the number of matches. Each element is a list of two elements, named "match" and "mapping". The first element is TRUE if the query matched the target. If so, the second element is a list of numeric vectors, giving the atom indices (0-indexed) of the target atoms that matched the query. If there was no match for this target molecule, this element will be NULL.

For isNsubgraph, a boolean vector, where each element is TRUE or FALSE depending on whether the corresponding element in targets contains the query or not.

For get.mcs an IAtomContainer object or a 2D array of atom index mappings between the two molecules.

Author(s)

Rajarshi Guha (<rajarshi.guha@gmail.com>)
See Also

load.molecules, get.smiles, do.aromaticity, do.typing, do.isotopes

Examples

smiles <- c('CCC', 'c1cccccl', 'C(C)(C=O)C(CC=C)c1cc1c(=O)')
mols <- sapply(smiles, parse.smiles)
query <- '[#6]=O'
doesMatch <- matches(query, mols)

## get mappings
mappings <- matches("CCC", mols, TRUE)

Molecule

Operations on molecules

Description

Various functions to perform operations on molecules.

get.exact.mass returns the exact mass of a molecule

get.natural.mass returns the natural exact mass of a molecule

convert.implicit.to.explicit converts implicit hydrogens to explicit hydrogens. This function
does not return any value but rather modifies the molecule object passed to it

is.neutral returns TRUE if all atoms in the molecule have a formal charge of 0, otherwise FALSE

Usage

get.exact.mass(molecule)
get.natural.mass(molecule)
convert.implicit.to.explicit(molecule)
is.neutral(molecule)

Arguments

molecule A jObjRef representing an IAtomContainer or IMolecule object

Details

In some cases, a molecule may not have any hydrogens (such as when read in from an MDL MOL
file that did not have hydrogens). In such cases, convert.implicit.to.explicit will add implicit
hydrogens and then convert them to explicit ones. In addition, for such cases, make sure that the
molecule has been typed beforehand.
parse.smiles

Value

- `exact_mass` returns a numeric
- `get_natural_mass` returns a numeric
- `convert_implicit_to_explicit` has no return value
- `is_neutral` returns a boolean.

Author(s)

Rajarshi Guha (<rajarshi.guha@gmail.com>)

See Also

- `get_atoms`, `do_typing`

Examples

```r
m <- parse.smiles('c1ccccc1')[[1]]

## Need to configure the molecule
do.aromaticity(m)
do.typing(m)
do.isotopes(m)

get.exact.mass(m)
get.natural.mass(m)

convert.implicit.to.explicit(m)
get.natural.mass(m)
do.isotopes(m) # Configure isotopes of newly added hydrogens
get.exact.mass(m)

is.neutral(m)
```

---

**parse.smiles**  
*Parse a Vector of SMILES Strings*

**Description**

This function parses a vector of SMILES strings to generate a list of `IAAtomContainer` objects. Note that the resultant molecule will not have any 2D or 3D coordinates.

Note that the molecules obtained from this method will not have any aromaticity perception, atom typing or isotopic configuration done on them. This is in contrast to the `load.molecules` method. Thus, you should perform these steps manually on the molecules.

**Usage**

```r
parse.smiles(smiles, kekulise=TRUE)
```
Removing Hydrogens from a Molecule

**Arguments**

- **smiles**: A SMILES string.
- **kekulise**: If set to `false`, disables electron checking and allows for parsing of incorrect SMILES. If a SMILES does not parse by default, try setting this to `false` - though the resultant molecule may not have consistent bonding. As an example, `c4ccc2c(cc1=Nc3ncccc3(Cn12))c4` will not be parsed by default because it is missing a nitrogen. With this argument set to `false`, it will parse successfully, but this is a hack to handle an incorrect SMILES.

**Value**

A list of `jobjrefs` to their corresponding CDK `IAtomContainer` objects. If a SMILES string could not be parsed, `NA` is returned instead.

**Author(s)**

Rajarshi Guha (<rajarshi.guha@gmail.com>)

**See Also**

- `load.molecules`, `get.smiles`, `get.smiles.parser`, `view.molecule.2d`, `do.aromaticity`, `do.typing`, `do.isotopes`

**Examples**

```r
smiles <- c('CCC', 'c1ccccc1', 'C(C)(C=O)C(CNC)C1CC1C(=O)')
mol <- parse.smiles(smiles[1])
mols <- parse.smiles(smiles)
```

---

**Description**

This function generates a new `IAtomContainer` object in which the hydrogens have been removed. This can be useful for descriptor calculations.

**Usage**

```r
remove.hydrogens(molecule)
```

**Arguments**

- **molecule**: A Java object of class `IAtomContainer`.

**Value**

A `jobjref` that refers to a `IAtomContainer` object.
Author(s)
Rajarshi Guha (<rajarshi.guha@gmail.com>)

remove.property Remove A Property From a Molecule

Description
This function removes a keyed property from a molecule object. This deletes the key and its value from the molecule.

Usage
remove.property(molecule, key)

Arguments
molecule A Java object of class IAtomContainer
key A string naming the property

Value
None

Author(s)
Rajarshi Guha (<rajarshi.guha@gmail.com>)

See Also
get.property, set.property

set.charge.formula Set the charge to a cdkFormula object.

Description
This function set the charge to a cdkFormula object. It modifies as the IMolecularFormula Java object as the its mass.

Usage
set.charge.formula(formula, charge=-1)
**Arguments**

- **formula**: A cdkFormula object.
- **charge**: The value of the charge to set.

**Value**

Returns the formula object with the specified charge

**Author(s)**

Miguel Rojas-Cherto (<miguelrojasch@yahoo.es>)

**References**

A parallel effort to expand the Chemistry Development Kit: [http://cdk.sourceforge.net](http://cdk.sourceforge.net)

**See Also**

- `get.formula`, `get.isotopes.pattern`, `isValid.formula`, `generate.formula`

---

**Description**

This function allows one to add a keyed property to a molecule. The key must be a string, but the value can be string, numeric or even an arbitrary Java object (of class `objRef`)

**Usage**

```
set.property(molecule, key, value)
```

**Arguments**

- **molecule**: A Java object of class `IAtomContainer`
- **key**: A string naming the property
- **value**: The value of the property. This can be character, integer, double or of class `objRef`

**Value**

None

**Author(s)**

Rajarshi Guha (<rajarshi.guha@gmail.com>)
smiles.flavors

See Also

get.property, get.properties, remove.property

Examples

```r
smiles <- 'c1ccccc1'
mol <- parse.smiles(smiles)[[1]]
set.property(mol, 'prop1', 23.45)
set.property(mol, 'prop2', 'inactive')
get.properties(mol)
```

Description

The CDK supports a variety of customizations for SMILES generation including the use of lower case symbols for aromatic compounds to the use of the ChemAxon CxSmiles format. Each 'flavor' is represented by an integer and multiple customizations are bitwise OR'ed. This method accepts the names of one or more customizations and returns the bitwise OR of them. See CDK documentation for the list of flavors and what they mean.

Usage

```r
smiles.flavors(flavors = c("Generic"))
```

Arguments

- `flavors`  
  A character vector of flavors. The default is Generic (Output non-canonical SMILES without stereochemistry, atomic masses). Possible values are
  - Absolute
  - AtomAtomMap
  - AtomicMass
  - AtomicMassStrict
  - Canonical
  - Cx2dCoordinates
  - Cx3dCoordinates
  - CxAtomLabel
  - CxAtomValue
  - CxCoordinates
  - CxFragmentGroup
  - CxMulticenter
  - CxPolymer
  - CxRadical
  - CxSmiles
• CxSmilesWithCoords
• Default
• Generic
• InChILabelling
• Isomeric
• Stereo
• StereoCisTrans
• StereoExTetrahedral
• StereoTetrahedral
• Unique
• UniversalSmiles
• UseAromaticSymbols

**Value**

A numeric representing the bitwise OR of the specified flavors

**Author(s)**

Rajarshi Guha <rajarshi.guha@gmail.com>

**References**

CDK documentation

**See Also**

get.smiles

**Examples**

```r
m <- parse.smiles('C1C=CCC1N(C)c1cccc1')[[1]]
gt.smiles(m)
gt.smiles(m, smiles.flavors(c('Generic','UseAromaticSymbols')))  
```

```r
m <- parse.smiles("OC(=O)=O)c1ccc1c1ccc1 [Sg:n:13:m:ht,Sg:n:11:n:ht"][[1]]
gt.smiles(m,flavor = smiles.flavors(c("CxSmiles")))
gt.smiles(m,flavor = smiles.flavors(c("CxSmiles","UseAromaticSymbols")))
```
Description

The CDK is capable of generating 2D structure diagrams. These methods allow one to view 2D structure diagrams. Depending on the method called a Swing JFrame is displayed which allows resizing of the image or a raster image (derived from a PNG byte stream) is is returned, which can be viewed using `rasterImage`. It is also possible to copy a 2D depiction to the system clipboard, which can then be pasted into various external applications.

Usage

```java
generateDepictor(width = 200, height = 200, zoom = 1.3, style = "cow",
annotate = "off", abbr = "on", suppressH = TRUE,
showTitle = FALSE, smalimit = 100, sma = NULL)
generateNdepictorNCwithNCnwidthNCandNCheightNCandNCzoomNCandNCstyleNCandNCannotateNCandNCabbrNCandNsuppressHNCandNshowTitleNCandNsmalimitNCandNsmaNCnull
```

Arguments

- `molecule`: If a single molecule is to be viewed this should be a reference to an `IAAtomContainer` object. If multiple molecules are to be viewed this should be a list of such objects. If a character is specified then it is taken as the name of a file and the molecules are loaded from the file.
- `depictor`: A depiction object. If NULL then one with default settings is created.
- `ncol`: The number of columns if a grid is desired.
- `width`: The width of the image.
- `height`: The height of the image.
- `zoom`: Zoom factor.
- `style`: Depiction style. Possible values are 'cow', 'wob', 'cob', 'nob'.
- `annotate`: Annotation style. By default no annotations are added. Possible values include 'number', 'mapidx', 'atomvalue', 'colmap'.
- `abbr`: Abbreviation style for functional groups. Possible values are 'groups', 'reagents', 'on'.
- `suppressH`: When TRUE show H's otherwise hide them.
- `showTitle`: When TRUE display title.
- `smalimit`: How many SMARTS patterns should be highlighted?
- `sma`: A string containing the SMARTS pattern to match.
Details

For the case of view.molecule.2d, if a jobjRef is passed it should be a reference to an IAtomContainer object. In case the first argument is of class character it is assumed to be a file and is loaded by the function.

This function can be used to view a single molecule or multiple molecules. If a list of molecule objects is supplied the molecules are displayed as a grid of 2D viewers. In case a file is specified, it will display a single molecule or multiple molecules depending on how many molecules are loaded.

For view.image.2d, the image can be viewed via rasterImage.

copy.image.to.clipboard copies the 2D depiction to the system clipboard in PNG format. You can then paste into other applications.

Due to event handling issues, the depiction will show on OS X, but the window will be unresponsive. Also copying images to the clipboard will not work. As a result, on OS X we make use of a standalone helper that is run via the system command. Currently, this is supported for the view.molecule.2d method (for a single molecule) and the copy.image.to.clipboard method.

In the future, other view methods will also be accessible via this mechanism. While this allows OS X users to view molecules, it is slow due to invoking a new process.

The depictions will work fine (i.e., no need to shell out) on Linux and Windows.

Value

get.depictor returns a depiction object that can be supplied to other methods. view.molecule.2d and copy.image.to.clipboard do not return anything. view.image.2d returns an array of the dimensions height x width x channels, from the original PNG version of the 2D depiction.

Author(s)

Rajarshi Guha (<rajarshi.guha@gmail.com>)

See Also

view.table, rasterImage, readPNG

Examples

m <- parse.smiles('c1ccccc1C(=O)NC')[[1]]

## Not run:
dep <- get.depictor(width=200, height=200)
img <- view.image.2d(m, dep)
plot(1:10, 1:10, pch=19)
rasterImage(img, 0,8, 2,10)

dep$setHeight(as.integer(400))
dep$setWidth(as.integer(400))
copy.image.to.clipboard(m, d) ## Paste into Word

## End(Not run)
view.table

View 2D Structures With Data

Description

The CDK is capable of generating 2D structure diagrams. This function can be used to view a set of molecules along with some associated data. The format of the output is a table, where the first column are the 2D images of the molecules, followed by the data columns.

Usage

view.table(molecules, dat, cellx = 200, celly = 200)

Arguments

molecules  A list of jobRef objects that represent IAtomContainer
dat  A data.frame containing numeric or character columns. If columns are named they will be used in the data table. If not, names are autogenerated. The number of rows of the data.frame should be equal to the number of molecules
cellx  Initial width of the table cells
celly  Initial height of the table cells

Details

Due to event handling issues, the depiction will show on OS X, but the window will be unresponsive. The depictions will work fine on Linux and Windows.

Value

Nothing

Author(s)

Rajarshi Guha (<rajarshi.guha@gmail.com>)

See Also

view.molecule.2d

Examples

smiles <- c('CCC', 'CCN', 'CCN(C)(C)',
'c1ccccc1C1ccccc1',
'1CCC1CC(CN(C)(C))CC(=O)CC'
mols <- parse.smiles(smiles)
dframe <- data.frame(x = runif(4),
  toxicity = factor(c('Toxic', 'Toxic', 'Nontoxic', 'Nontoxic'))),
  solubility = c('yes', 'yes', 'no', 'yes'))
## Not run: view.table(mols[1:4], dframe)
write.molecules  Write Molecules To Disk

Description

This function writes one or more molecules to an SD file on disk, which can be of the single- or multi-molecule variety. In addition, if the molecule has keyed properties, they can also be written out as SD tags.

Usage

write.molecules(mols, filename, together=TRUE, write.props=FALSE)

Arguments

- **mols**: A list of Java objects of class IAtomContainer
- **filename**: The name of the SD file to write. Note that if together is FALSE then this argument is taken as a prefix for the name of the individual files
- **together**: If TRUE then all the molecules are written to a single SD file. If FALSE each molecule is written to an individual file
- **write.props**: Should keyed properties be included in the SD file output

Details

This function can be used to write a single SD file containing multiple molecules. In case individual SD files are desired the together argument can be set to FALSE. In this case, the value of filename is used as a prefix, to which a numeric identifier and the suffix of ".sdf" is appended. In case, a single molecule is to be written to disk, simply specify the filename and use the default value of together

Value

The value of the property

Author(s)

Rajarshi Guha (<rajarshi.guha@gmail.com>)

See Also

load.molecules, set.property, get.property, remove.property
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