### NAME

**RDKitUtil** 

#### **SYNOPSIS**

import RDKitUtil

# **DESCRIPTION**

RDKitUtil module provides the following functions:

GetInlineSVGForMolecule, GetInlineSVGForMolecules, GetMolName, GetSVGForMolecule, GetSVGForMolecules, IsMolEmpty, MoleculesWriter, ReadAndValidateMolecules, ReadMolecules, ReadMoleculesFromMol2File, ReadMoleculesFromMolFile, ReadMoleculesFromSDFile, ReadMoleculesFromSMILESFile, WriteMolecules

#### **FUNCTIONS**

#### GetInlineSVGForMolecule

```
GetInlineSVGForMolecule(Mol, Width, Height, Legend = None, AtomListToHighlight = None,
BondListToHighlight = None, BoldText = True)
```

Get SVG image text for a molecule suitable for inline embedding into a HTML page.

### Arguments:

```
Mol (object): RDKit molecule object.
Width (int): Width of a molecule image in pixels.
Height (int): Height of a molecule image in pixels.
Legend (str): Text to display under the image.
AtomListToHighlight (list): List of atoms to highlight.
BondListToHighlight (list): List of bonds to highlight.
BoldText (bool): Flag to make text bold in the image of molecule.
```

#### Returns:

```
str : SVG image text for inline embedding into a HTML page using "img"
tag: <img src="data:image/svg+xml;charset=UTF-8,SVGImageText>
```

## GetInlineSVGForMolecules

```
GetInlineSVGForMolecules(Mols, MolsPerRow, MolWidth, MolHeight, Legends = None,
AtomListsToHighlight = None, BondListsToHighLight = None, BoldText = True)
```

Get SVG image text for molecules suitable for inline embedding into a HTML page.

# Arguments:

```
Mols (list): List of RDKit molecule objects.

MolsPerRow (int): Number of molecules per row.

Width (int): Width of a molecule image in pixels.

Height (int): Height of a molecule image in pixels.

Legends (list): List containing strings to display under images.

AtomListsToHighlight (list): List of lists containing atoms to highlight for molecules.

BondListsToHighlight (list): List of lists containing bonds to highlight for molecules

BoldText (bool): Flag to make text bold in the image of molecules.
```

### Returns:

```
str : SVG image text for inline embedding into a HTML page using "img"
tag: <img src="data:image/svg+xml;charset=UTF-8,SVGImageText>
```

# GetMolName

```
GetMolName(Mol, MolNum = None)
```

Get molecule name.

### Arguments:

```
Mol (object): RDKit molecule object.
MolNum (int or None): Molecule number in input file.
```

# Returns:

```
str : Molname corresponding to _Name property of a molecule, generated
from specieid MolNum using the format "Mol%d" % MolNum, or an
empty string.
```

### GetSVGForMolecule

```
GetSVGForMolecule(Mol, Width, Height, Legend = None, AtomListToHighlight = None,
BondListToHighlight = None, BoldText = True)
```

Get SVG image text for a molecule suitable for viewing in a browser.

```
Arguments:
```

```
Mol (object): RDKit molecule object.

Width (int): Width of a molecule image in pixels.

Height (int): Height of a molecule image in pixels.

Legend (str): Text to display under the image.

AtomListToHighlight (list): List of atoms to highlight.

BondListToHighlight (list): List of bonds to highlight.

BoldText (bool): Flag to make text bold in the image of molecule.
```

#### Returns:

str : SVG image text for writing to a SVG file for viewing in a browser.

# GetSVGForMolecules

GetSVGForMolecules(Mols, MolsPerRow, MolWidth, MolHeight, Legends = None, AtomListsToHighlight =
None, BondListsToHighlight = None, BoldText = True)

Get SVG image text for molecules suitable for viewing in a browser.

#### Arguments:

```
Mols (list): List of RDKit molecule objects.

MolsPerRow (int): Number of molecules per row.

Width (int): Width of a molecule image in pixels.

Height (int): Height of a molecule image in pixels.

Legends (list): List containing strings to display under images.

AtomListsToHighlight (list): List of lists containing atoms to highlight for molecules.

BondListsToHighlight (list): List of lists containing bonds to highlight for molecules

BoldText (bool): Flag to make text bold in the image of molecules.
```

#### Returns:

str : SVG image text for writing to a SVG file for viewing in a browser.

#### IsMolEmpty

IsMolEmpty(Mol)

Check for the presence of atoms in a molecule.

# Arguments:

Mol (object): RDKit molecule object.

# Returns:

bool : True - No atoms in molecule; Otherwise, false.

### MoleculesWriter

MoleculesWriter(FileName, \*\*KeyWordArgs)

Set up a molecule writer.

### Arguments:

```
FileName (str): Name of a file with complete path.
**KeyWordArgs (dictionary) : Parameter name and value pairs for writing and
processing molecules.
```

### Returns:

```
RDKit object : Molecule writer.
```

The file extension is used to determine type of the file and set up an appropriate file writer.

### ReadAndValidateMolecules

```
{\tt ReadAndValidateMolecules(FileName, **KeyWordArgs)}
```

Read molecules from an input file, validate all molecule objects, and return a list of valid and non-valid molecule objects along with their counts.

### Arguments:

```
FileName (str): Name of a file with complete path.
**KeyWordArgs (dictionary) : Parameter name and value pairs for reading and
processing molecules.
```

### Returns:

list : List of valid RDKit molecule objects.

```
int : Number of total molecules in input file.
int : Number of valid molecules in input file.
```

The file extension is used to determine type of the file and set up an appropriate file reader.

#### ReadMolecules

```
ReadMolecules(FileName, **KeyWordArgs)
```

Read molecules from an input file without performing any validation and creation of molecule objects.

#### Arguments:

```
FileName (str): Name of a file with complete path.
**KeyWordArgs (dictionary) : Parameter name and value pairs for reading and
processing molecules.
```

#### Returns:

```
list : List of RDKit molecule objects.
```

The file extension is used to determine type of the file and set up an appropriate file reader.

#### ReadMoleculesFromMol2File

```
ReadMoleculesFromMol2File(FileName, Sanitize = True, RemoveHydrogens = True)
```

Read molecule from a Tripos Mol2 file.

#### Arguments:

```
FileName (str): Name of a file with complete path. Sanitize (bool): Sanitize molecules. RemoveHydrogens (bool): Remove hydrogens from molecules.
```

#### Returns:

```
list : List of RDKit molecule objects.
```

#### ReadMoleculesFromMolFile

ReadMoleculesFromMolFile(FileName, Sanitize = True, RemoveHydrogens = True, StrictParsing = True)

Read molecule from a MDL Mol file.

# Arguments:

```
FileName (str): Name of a file with complete path.
Sanitize (bool): Sanitize molecules.
RemoveHydrogens (bool): Remove hydrogens from molecules.
StrictParsing (bool): Perform strict parsing.
```

# Returns:

```
list : List of RDKit molecule objects.
```

### ReadMoleculesFromPDBFile

```
ReadMoleculesFromPDBFile(FileName, Sanitize = True, RemoveHydrogens = True)
```

Read molecule from a PDB file.

# Arguments:

```
FileName (str): Name of a file with complete path.
Sanitize (bool): Sanitize molecules.
RemoveHydrogens (bool): Remove hydrogens from molecules.
```

### Returns:

```
list : List of RDKit molecule objects.
```

### ReadMoleculesFromSDFile

```
ReadMoleculesFromSDFile(FileName, Sanitize = True, RemoveHydrogens = True, StrictParsing = True)
```

Read molecules from a SD file.

# Arguments:

```
FileName (str): Name of a file with complete path. Sanitize (bool): Sanitize molecules. RemoveHydrogens (bool): Remove hydrogens from molecules. StrictParsing (bool): Perform strict parsing.
```

### Returns:

```
list : List of RDKit molecule objects.
```

#### ReadMoleculesFromSMILESFile

```
ReadMoleculesFromSMILESFile(FileName, SMILESDelimiter = ' ', SMILESColIndex = 0,
SMILESNameColIndex = 1, SMILESTitleLine = 1, Sanitize = 1)
```

Read molecules from a SMILES file.

# Arguments:

```
SMILESDelimiter (str): Delimiter for parsing SMILES line SMILESColIndex (int): Column index containing SMILES string. SMILESNameColIndex (int): Column index containing molecule name. SMILESTitleLine (int): Flag to indicate presence of title line. Sanitize (int): Sanitize molecules.
```

#### Returns:

list : List of RDKit molecule objects.

#### WriteMolecules

```
WriteMolecules(FileName, Mols, **KeyWordArgs)
```

Write molecules to an output file.

### Arguments:

```
FileName (str): Name of a file with complete path.
Mols (list): List of RDKit molecule objects.
**KeyWordArgs (dictionary) : Parameter name and value pairs for writing and processing molecules.
```

#### Returns:

```
int : Number of total molecules.
int : Number of processed molecules written to output file.
```

The file extension is used to determine type of the file and set up an appropriate file writer.

#### **AUTHOR**

Manish Sud <msud@san.rr.com>

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The functionality available in the script is implemented using RDKit, an open source toolkit for cheminformatics developed by Greg Landrum.

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