# NAME

RDKitRemoveInvalidMolecules.py - Remove invalid molecules

# **SYNOPSIS**

RDKitRemoveInvalidMolecules.py -h | --help | -e | --examples

# **DESCRIPTION**

Identify and remove invalid molecules based on success or failure of RDKit molecule readers or simply count the number of invalid molecules.

The supported input file formats are: SD (.sdf, .sd), SMILES (.smi., csv, .tsv, .txt)

The supported output file formats are: SD (.sdf, .sd), SMILES (.smi)

# **OPTIONS**

-e, --examples

Print examples.

-h, --help

Print this help message.

-i, --infile <infile>

Input file name.

--infileParams <Name, Value,...> [default: auto]

A comma delimited list of parameter name and value pairs for reading molecules from files. The supported parameter names for different file formats, along with their default values, are shown below:

```
SD: removeHydrogens,yes,sanitize,yes,strictParsing,yes
SMILES: smilesColumn,1,smilesNameColumn,2,smilesDelimiter,space,
    smilesTitleLine,auto,sanitize,yes
```

Possible values for smilesDelimiter: space, comma or tab.

-m, --mode <remove or count> [default: remove]

Specify whether to remove invalid molecules and write out filtered molecules to output file or or simply count the number of invalid molecules.

-o, --outfile <outfile>

Output file name.

--outfileParams <Name, Value, ... > [default: auto]

A comma delimited list of parameter name and value pairs for writing molecules to files. The supported parameter names for different file formats, along with their default values, are shown below:

```
SD: compute2DCoords,auto,kekulize,yes
SMILES: smilesKekulize,no,smilesDelimiter,space, smilesIsomeric,yes,
    smilesTitleLine,yes,smilesMolName,yes,smilesMolProps,no
```

Default value for compute2DCoords: yes for SMILES input file; no for all other file types.

--overwrite

Overwrite existing files.

-w, --workingdir <dir>

Location of working directory which defaults to the current directory.

# **EXAMPLES**

To remove invalid molecules and generate an output file SMILES file containing valid molecules, type:

% RDKitRemoveInvalidMolecules.py -i Sample.smi -o SampleOut.smi

To count number of valid and invaid molecules without generating any output file, type:

% RDKitRemoveInvalidMolecules.py -m count -i Sample.sdf

To remove invalid molecules from a CSV SMILES file, SMILES strings in column 1, name in column 2, and generate output SD file containing valid molecules, type:

```
% RDKitRemoveInvalidMolecules.py --infileParams
"smilesDelimiter,comma,smilesTitleLine,yes,smilesColumn,1,
smilesNameColumn,2" --outfileParams "compute2DCoords,yes"
-i SampleSMILES.csv -o SampleOut.sdf
```

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# **SEE ALSO**

RDKitConvertFileFormat.py, RDKitRemoveDuplicateMolecules.py, RDKitRemoveSalts, RDKitSearchFunctionalGroups.py, RDKitSearchSMARTS.py, RDKitStandardizeMolecules.py

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

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