RDKit Documentation

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AN OVERVIEW OF THE RDKIT

1.1 What is it?

- Open source toolkit for cheminformatics
 - BSD licensed
 - Core data structures and algorithms in C++
 - Python (2.x) wrapper generated using Boost.Python
 - Java and C# wrappers generated with SWIG
 - 2D and 3D molecular operations
 - Descriptor generation for machine learning
 - Molecular database cartridge for PostgreSQL
 - Cheminformatics nodes for KNIME (distributed from the KNIME community site: http://tech.knime.org/community/rdkit)
- Operational:
 - http://www.rdkit.org
 - Supports Mac/Windows/Linux
 - Releases every 6 months
 - Web presence:
 - * Homepage: http://www.rdkit.org

Documentation, links

* Github (https://github.com/rdkit)

Bug tracker, git repository

* Sourceforge (http://sourceforge.net/projects/rdkit)

Mailing lists, Downloads

* Google code (http://code.google.com/p/rdkit/)

wiki

- Mailing lists at https://sourceforge.net/p/rdkit/mailman/, searchable archives available for rdkitdiscuss and rdkit-devel
- History:

- 2000-2006: Developed and used at Rational Discovery for building predictive models for ADME, Tox, biological activity
- June 2006: Open-source (BSD license) release of software, Rational Discovery shuts down
- to present: Open-source development continues, use within Novartis, contributions from Novartis back to open-source version

1.2 Functionality overview

- Input/Output: SMILES/SMARTS, SDF, TDT, SLN 1, Corina mol2 1, PDB
- "Cheminformatics":
 - Substructure searching
 - Canonical SMILES
 - Chirality support (i.e. R/S or E/Z labeling)
 - Chemical transformations (e.g. remove matching substructures)
 - Chemical reactions
 - Molecular serialization (e.g. mol <-> text)
- 2D depiction, including constrained depiction
- 2D->3D conversion/conformational analysis via distance geometry
- UFF and MMFF94/MMFF94S implementations for cleaning up structures
- Fingerprinting: Daylight-like, atom pairs, topological torsions, Morgan algorithm, "MACCS keys", etc.
- · Similarity/diversity picking
- 2D pharmacophores ¹
- Gasteiger-Marsili charges
- Hierarchical subgraph/fragment analysis
- · Bemis and Murcko scaffold determination
- RECAP and BRICS implementations
- Multi-molecule maximum common substructure ²
- Feature maps
- · Shape-based similarity
- · RMSD-based molecule-molecule alignment
- Shape-based alignment (subshape alignment ³) ¹
- Unsupervised molecule-molecule alignment using Open3DAlign algorithm
- Integration with PyMOL for 3D visualization

¹ These implementations are functional but are not necessarily the best, fastest, or most complete.

² Contribution from Andrew Dalke

³ Putta, S., Eksterowicz, J., Lemmen, C. & Stanton, R. "A Novel Subshape Molecular Descriptor" *Journal of Chemical Information and Computer Sciences* 43:1623–35 (2003).

⁴ Tosco, P., Balle, T. & Shiri, F. Open3DALIGN: an open-source software aimed at unsupervised ligand alignment. *J Comput Aided Mol Des* **25:777–83** (2011).

- · Functional group filtering
- · Salt stripping
- Molecular descriptor library:
 - Topological (κ 3, Balaban J, etc.)
 - Compositional (Number of Rings, Number of Aromatic Heterocycles, etc.)
 - Electrotopological state (Estate)
 - clogP, MR (Wildman and Crippen approach)
 - "MOE like" VSA descriptors
 - Feature-map vectors ⁵
 - MQN⁶
- Similarity Maps ⁷
- Machine Learning:
 - Clustering (hierarchical, Butina)
 - Information theory (Shannon entropy, information gain, etc.)
- Tight integration with the IPython notebook and qtconsole.

1.3 The Contrib Directory

The Contrib directory, part of the standard RDKit distribution, includes code that has been contributed by members of the community.

• LEF: Local Environment Fingerprints

Contains python source code from the publications:

- Vulpetti, U. Hommel, G. Landrum, R. Lewis and C. Dalvit, "Design and NMR-based screening of LEF, a library of chemical fragments with different Local Environment of Fluorine" *J. Am. Chem.* Soc. 131 (2009) 12949-12959. http://dx.doi.org/10.1021/ja905207t
- Vulpetti, G. Landrum, S. Ruedisser, P. Erbel and C. Dalvit, "19F NMR Chemical Shift Prediction with Fluorine Fingerprint Descriptor" *J. of Fluorine Chemistry* 131 (2010) 570-577. http://dx.doi.org/10.1016/j.jfluchem.2009.12.024

Contribution from Anna Vulpetti

• M Kossner:

Contains a set of pharmacophoric feature definitions as well as code for finding molecular frameworks.

Contribution from Markus Kossner

• PBF: Plane of best fit

Contains C++ source code and sample data from the publication:

⁵ Landrum, G., Penzotti, J. & Putta, S. "Feature-map vectors: a new class of informative descriptors for computational drug discovery" *Journal of Computer-Aided Molecular Design* **20:751–62** (2006).

⁶ Nguyen, K. T., Blum, L. C., van Deursen, R. & Reymond, J.-L. Classification of Organic Molecules by Molecular Quantum Numbers. *ChemMedChem* **4:1803–5** (2009).

⁷ Riniker, S. & Landrum, G. A. Similarity maps - a visualization strategy for molecular fingerprints and machine-learning methods. *Journal of Cheminformatics* **5:43** (2013).

 (a) Firth, N. Brown, and J. Blagg, "Plane of Best Fit: A Novel Method to Characterize the Three-Dimensionality of Molecules" *Journal of Chemical Information and Modeling* 52 2516-2525 (2012). http://pubs.acs.org/doi/abs/10.1021/ci300293f

Contribution from Nicholas Firth

• mmpa: Matched molecular pairs

Python source and sample data for an implementation of the matched-molecular pair algorithm described in the publication:

Hussain, J., & Rea, C. "Computationally efficient algorithm to identify matched molecular pairs (MMPs) in large data sets." *Journal of chemical information and modeling* **50** 339-348 (2010). http://dx.doi.org/10.1021/ci900450m

Includes a fragment indexing algorithm from the publication:

Wagener, M., & Lommerse, J. P. "The quest for bioisosteric replacements." *Journal of chemical information and modeling* **46** 677-685 (2006).

Contribution from Jameed Hussain.

• SA_Score: Synthetic assessibility score

Python source for an implementation of the SA score algorithm described in the publication:

Ertl, P. and Schuffenhauer A. "Estimation of Synthetic Accessibility Score of Drug-like Molecules based on Molecular Complexity and Fragment Contributions" *Journal of Cheminformatics* **1:8** (2009)

Contribution from Peter Ertl

• fraggle: A fragment-based molecular similarity algorithm

Python source for an implementation of the fraggle similarity algorithm developed at GSK and described in this RDKit UGM presentation: https://github.com/rdkit/UGM_2013/blob/master/Presentations/Hussain.Fraggle.pdf

Contribution from Jameed Hussain

1.4 License

This document is copyright (C) 2013-2014 by Greg Landrum

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The intent of this license is similar to that of the RDKit itself. In simple words: "Do whatever you want with it, but please give us some credit."

CHAPTER

TWO

INSTALLATION

Below a number of installation recipies is presented, with varying degree of complexity.

2.1 Linux and the Mac

2.1.1 Installation from repositories

Ubuntu 12.04 and later

Thanks to the efforts of the Debichem team, RDKit is available via the Ubuntu repositories. To install:

sudo apt-get install python-rdkit librdkit1 rdkit-data

Fedora, CentOS, and RHEL

Gianluca Sforna creates binary RPMs that can be found here: ">http://giallu.fedorapeople.org/rdkit-20XX.XX/>

MacOS

Eddie Cao has produced a homebrew formula that can be used to easily build the RDKit https://github.com/rdkit/homebrew-rdkit_

2.1.2 Building from Source

Prerequisites

Installing prerequisites as packages

Ubuntu and other debian-derived systems Install the following packages using apt-get:

flex bison build-essential python-numpy cmake python-dev sqlite3 libsqlite3-dev libboost-dev libboost-python-dev libboost-regex-dev

Fedora, CentOS (5.7+), and RHEL Install the following packages using yum:

cmake tk-devel readline-devel zlib-devel bzip2-devel sqlite-devel @development-tools

Packages to install from source (not required on RHEL/CentOS 6.x):

- python 2.7 : use ./configure CFLAGS=-fPIC -enable-unicode=ucs4 -enable-shared
- numpy : do export LD_LIBRARY_PATH="/usr/local/lib" before python setup.py install
- boost 1.48.0 or later: do ./bootstrap.sh –with-libraries=python,regex; ./b2; ./b2 install

Older versions of CentOS Here things are more difficult. Check this wiki page for information: https://code.google.com/p/rdkit/wiki/BuildingOnCentOS

Installing prerequisites from source

- · Required packages:
 - cmake. You need version 2.6 (or more recent). http://www.cmake.org if your linux distribution doesn't have an appropriate package.

Note: It seems that v2.8 is a better bet than v2.6. It might be worth compiling your own copy of v2.8 even if v2.6 is already installed.

- The following are required if you are planning on using the Python wrappers
 - * The python headers. This probably means that you need to install the python-dev package (or whatever it's called) for your linux distribution.
 - * sqlite3. You also need the shared libraries. This may require that you install a sqlite3-dev package.
 - * You need to have numpy (http://www.scipy.org/NumPy) installed.

Note: for building with XCode4 on the MacOS – there seems to be a problem with the version of numpy that comes with XCode4. Please see below in the (see *Frequently Encountered Problems*) section for a workaround.

- · Optional packages
 - If you would like to install the RDKit InChI support (first available in the Q2 2011 release), follow the instructions in \$RDBASE/External/INCHI-API to get a copy of the InChI source and put it in the appropriate place.

Installing Boost If your linux distribution has a boost-devel package including the python and regex libraries, you can use that and save yourself the steps below.

Note: if you *do* have a version of the boost libraries pre-installed and you want to use your own version, be careful when you build the code. We've seen at least one example on a Fedora system where cmake compiled using a user-installed version of boost and then linked against the system version. This led to segmentation faults. There is a workaround for this below in the (see *Frequently Encountered Problems*) section.

- download the boost source distribution from the boost web site
- extract the source somewhere on your machine (e.g. /usr/local/src/boost_1_45_0)
- build the required boost libraries:

- cd \$BOOST
- If you want to use the python wrappers: ./bootstrap.sh --with-libraries=python, regex
- If not using the python wrappers: ./bootstrap.sh --with-libraries=regex
- Building on 32 bit systems: ./b2 install
- Building on 64 bit systems: ./b2 address-model=64 cflags=-fPIC cxxflags=-fPIC install

If you have any problems with this step, check the boost installation instructions.

Building the RDKit

Fetch the source, here as tar.gz but you could use git as well:

```
wget http://downloads.sourceforge.net/project/rdkit/rdkit/QX_20XX/RDKit_20XX_XX_X.tgz

* Ensure that the prerequisites are installed
* environment variables:

  * RDBASE: the root directory of the RDKit distribution (e.g. ~/RDKit)
  * *Linux:* LD_LIBRARY_PATH: make sure it includes $RDBASE/lib and wherever the boost shared librar.
  * *Mac:* DYLD_LIBRARY_PATH: make sure it includes $RDBASE/lib and wherever the boost shared librar.
  * The following are required if you are planning on using the Python wrappers:
  * PYTHONPATH: make sure it includes $RDBASE

* Building:

  * cd to $RDBASE
  * ''mkdir build''
  * ''cd build''
  * ''cmake ..'' : See the section below on configuring the build if you need to specify a non-default.
```

* ''make'' : this builds all libraries, regression tests, and wrappers (by default).

See below for a list of Frequently Encountered Problems and solutions.

Testing the build (optional, but recommended)

- cd to \$RDBASE/build and do ctest
- you're done!

* ''make install''

Advanced

Specifying an alternate Boost installation

You need to tell cmake where to find the boost libraries and header files:

If you have put boost in /opt/local, the cmake invocation would look like:

```
cmake -DBOOST_ROOT=/opt/local ..
```

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Specifying an alternate Python installation

You need to tell cmake where to find the python library it should link against and the python header files.

Here's a sample command line:

```
cmake -D PYTHON_LIBRARY=/usr/lib/python2.5/config/libpython2.5.a -D PYTHON_INCLUDE_DIR=/usr/include/j
```

The PYTHON_EXECUTABLE part is optional if the correct python is the first version in your PATH.

Disabling the Python wrappers

You can completely disable building of the python wrappers by setting the configuration variable RDK_BUILD_PYTHON_WRAPPERS to nil:

```
cmake -D RDK_BUILD_PYTHON_WRAPPERS= ..
```

Building the Java wrappers

Additional Requirements

- SWIG v2.0.x: http://www.swig.org
- Junit: get a copy of the junit .jar file from https://github.com/KentBeck/junit/downloads and put it in the directory \$RDBASE/External/java_lib (you will need to create the directory) and rename it to junit.jar.

Building

- When you invoke cmake add -D RDK_BUILD_SWIG_WRAPPERS=ON to the arguments. For example: cmake -D RDK_BUILD_SWIG_WRAPPERS=ON ..
- Build and install normally using *make*. The directory \$RDBASE/Code/JavaWrappers/gmwrapper will contain the three required files: libGraphMolWrap.so (libGraphMolWrap.jnilib on the Mac), org.RDKit.jar, and org.RDKitDoc.jar.

Using the wrappers

>>> m.getNumAtoms()

To use the wrappers, the three files need to be in the same directory, and that should be on your CLASSPATH and in the java.library.path. An example using jython:

```
% CLASSPATH=$CLASSPATH:$RDBASE/Code/JavaWrappers/gmwrapper/org.RDKit.jar; jython -Djava.library.path=
Jython 2.2.1 on java1.6.0_20
Type "copyright", "credits" or "license" for more information.
>>> from org.RDKit import *
>>> from java import lang
>>> lang.System.loadLibrary('GraphMolWrap')
>>> m = RWMol.MolFromSmiles('clcccc1')
```

Frequently Encountered Problems

In each case I've replaced specific pieces of the path with

Problem:

6L

```
Linking CXX shared library libSLNParse.so
/usr/bin/ld: .../libboost_regex.a(cpp_regex_traits.o): relocation R_X86_64_32S against 'std::basic_st
.../libboost_regex.a: could not read symbols: Bad value
collect2: ld returned 1 exit status
make[2]: *** [Code/GraphMol/SLNParse/libSLNParse.so] Error 1
make[1]: *** [Code/GraphMol/SLNParse/CMakeFiles/SLNParse.dir/all] Error 2
make: *** [all] Error 2
```

Solution:

Add this to the arguments when you call cmake: -DBoost_USE_STATIC_LIBS=OFF

More information here: http://www.mail-archive.com/rdkit-discuss@lists.sourceforge.net/msg01119.html

Problem:

```
.../Code/GraphMol/Wrap/EditableMol.cpp:114: instantiated from here
.../boost/type_traits/detail/cv_traits_impl.hpp:37: internal compiler error: in make_rtl_for_nonlocal
Please submit a full bug report,
with preprocessed source if appropriate.
See <URL:http://bugzilla.redhat.com/bugzilla> for instructions.
Preprocessed source stored into /tmp/ccgSaXge.out file, please attach this to your bugreport.
make[2]: *** [Code/GraphMol/Wrap/CMakeFiles/rdchem.dir/EditableMol.cpp.o] Error 1
make[1]: *** [Code/GraphMol/Wrap/CMakeFiles/rdchem.dir/all] Error 2
make: *** [all] Error 2
```

Solution:

Add #define BOOST_PYTHON_NO_PY_SIGNATURES at the top of Code/GraphMol/Wrap/EditableMol.cpp

More information here: http://www.mail-archive.com/rdkit-discuss@lists.sourceforge.net/msg01178.html

Problem:

Your system has a version of boost installed in /usr/lib, but you would like to force the RDKit to use a more recent one.

Solution:

This can be solved by using cmake version 2.8.3 (or more recent) and providing the -D Boost_NO_SYSTEM_PATHS=ON argument:

```
cmake -D BOOST_ROOT=/usr/local -D Boost_NO_SYSTEM_PATHS=ON ..
```

Problem:

Building on the Mac with XCode 4

The problem seems to be caused by the version of numpy that is distributed with XCode 4, so you need to build a fresh copy.

Solution: Get a copy of numpy and build it like this as root: as root:

```
export MACOSX_DEPLOYMENT_TARGET=10.6
export LDFLAGS="-Wall -undefined dynamic_lookup -bundle -arch x86_64"
export CFLAGS="-arch x86_64"
ln -s /usr/bin/gcc /usr/bin/gcc-4.2
ln -s /usr/bin/g++ /usr/bin/g++-4.2
```

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```
python setup.py build
python setup.py install
```

Be sure that the new numpy is used in the build:

```
PYTHON_NUMPY_INCLUDE_PATH /Library/Python/2.6/site-packages/numpy/core/include
```

and is at the beginning of the PYTHONPATH:

```
export PYTHONPATH="/Library/Python/2.6/site-packages:$PYTHONPATH"
```

Now it's safe to build boost and the RDKit.

2.2 Windows

2.2.1 Prerequisites

- Python 2.7 (from http://www.python.org/)
- numpy (from http://numpy.scipy.org/ or use pip install numpy). Binaries for win64 are available here: http://www.lfd.uci.edu/~gohlke/pythonlibs/#numpy
- PIL: (from http://www.pythonware.com/products/pil/index.htm or use pip install PIL)

Recommended extras

- aggdraw: a library for high-quality drawing in Python. Instructions for downloading are here: http://effbot.org/zone/aggdraw-index.htm The new (as of May 2008) drawing code has been tested with v1.2a3 of aggdraw. Despite the alpha label, the code is stable and functional.
- matplotlib: a library for scientific plotting from Python. http://matplotlib.sourceforge.net/
- ipython: a very useful interactive shell (and much more) for Python. http://ipython.scipy.org/dist/
- win32all: Windows extensions for Python. http://sourceforge.net/projects/pywin32/

2.2.2 Installation of RDKit binaries

- Get the appropriate windows binary build from: https://sourceforge.net/projects/rdkit/files/rdkit/>_
- Extract the zip file somewhere without a space in the name, i.e. c:/
- The rest of this will assume that the installation is in c:/RDKit_2012_12_1
- Set the following environment variables: * RDBASE: c:/RDKit_2012_12_1 * PYTHONPATH: %RDBASE% if there is already a PYTHONPATH, put; %RDBASE% at the end. * PATH: add; %RDBASE%/lib to the end

In Win7 systems, you may run into trouble due to missing DLLs, see one thread from the mailing list: http://www.mailarchive.com/rdkit-discuss@lists.sourceforge.net/msg01632.html You can download the missing DLLs from here: http://www.microsoft.com/en-us/download/details.aspx?id=5555

2.2.3 Installation from source

Extra software to install

- Microsoft Visual C++: The Express version has everything necessary and can be downloaded for free (http://www.microsoft.com/express/download/). This is a big installation and will take a while. The RDKit has been successfully built with all version of visual c++ since 6.0, so the current version of VC++ (2010 as of this writing) should be fine.
- cmake: (http://www.cmake.org/cmake/resources/software.html) should be installed.
- boost: It is strongly recommended to download and use a precompiled version of the boost libraries from http://sourceforge.net/projects/boost/files/boost-binaries/. When you run the installer, the only binary libraries you need are python, regex, and system. If you want to install boost from source, download a copy from http://www.boost.org and follow the instructions in the "Getting Started" section of the documentation. Make sure the libraries and headers are installed to c:boost
- a git client: This is only necessary if you are planning on building development versions of the RDKit. This can be downloaded from http://git-scm.com/downloads.
- · Optional packages
 - If you would like to install the RDKit InChI support, follow the instructions in \$RDBASE/External/INCHI-API/README to get a copy of the InChI source and put it in the appropriate place.
 - If you would like to install the RDKit Avalon toolkit support, follow the instructions in \$RD-BASE/External/AvalonTool/README to get a copy of the Avalon toolkit source and put it in the appropriate place.

Setup and Preparation

This section assumes that python is installed in c:\Python27, that the boost libraries have been installed to c:\boost, and that you will build the RDKit from a directory named c:\RDKit. If any of these conditions is not true, just change the corresponding paths.

- If you install things in paths that have spaces in their names, be sure to use quotes properly in your environment variable definitions.
- If you are planning on using a development version of the RDKit: get a copy of the current RDKit source using git. If you're using the command-line client the command is: git clone https://github.com/rdkit/rdkit.git c:\RDKit
- If you are planning on using a released version of the RDKit : get a copy of the most recent release and extract it into the directory c:\RDKit
- Set the required environment variables (you can set this in cygwin or in windows. If you set them in windows, be sure to restart your cygwin window)

```
- RDBASE = c:\RDKit
```

- Make sure c:\Python27 is in your PATH
- Make sure c:\RDKit\lib is in your PATH
- Make sure c:\boost\lib is in your PATH.
- Make sure c:\RDKit is in your PYTHONPATH

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Building from the command line (recommended)

- Create a directory c:\RDKit\build and cd into it
- Run cmake. Here's an example basic command line for 64bit windows that assumes the InChI and
 Avalon toolkit sources are available (see above): cmake -DRDK_BUILD_PYTHON_WRAPPERS=0N
 -DAVALONTOOLS_DIR=c:/avalontoolkit_beta/sourcedistribution
 -DBOOST_ROOT=c:/boost -DRDK_BUILD_INCHI_SUPPORT=ON -DRDK_BUILD_AVALON_SUPPORT=ON
 -G"Visual Studio 10 Win64" ..
- Build the code. Here's an example command line: c:/Windows/Microsoft.NET/Framework64/v4.0.30319/MSBui/m:4 /p:Configuration=Release INSTALL.vcxproj

Building the Code Using GUIs (not recommended)

- Environment variables: if cmake complains about not being able to find it, define the environment variable BOOST_ROOT to point to the directory containing the boost source.
- Configure the build:
 - Start the cmake gui
 - tell it where the source code is (e.g. c:/RDKit) and where to build the binaries (recommended: c:/RDKit/build)
 - click "Configure", select your compiler, and wait until the basic configuration is complete, you'll see a bunch of red entries in the main windows.
 - click "Configure" again
 - click "Generate"
- Build:
 - open the solution file that cmake created (c:/RDKit/build/RDKit.sln) with Visual Studio.
 - check to be sure that you're building a Release build (for some reason CMake produces solution files that default to doing a Debug build)
 - build the "ALL_BUILD" target; this will take a while and generate warnings, but there should be no errors.
 Note: if you are building the SWIG wrappers you may get an error the first time you try to build them. If you see this error, try building ALL_BUILD again; it should work the second time.
 - build the "INSTALL" target

Testing the Build (optional, but recommended)

- cd to c:\RDKit\build and run ctest.
- · you're done!

CHAPTER

THREE

GETTING STARTED WITH THE RDKIT IN PYTHON

3.1 What is this?

This document is intended to provide an overview of how one can use the RDKit functionality from Python. It's not comprehensive and it's not a manual.

If you find mistakes, or have suggestions for improvements, please either fix them yourselves in the source document (the .rst file) or send them to the mailing list: rdkit-devel@lists.sourceforge.net

3.2 Reading and Writing Molecules

3.2.1 Reading single molecules

The majority of the basic molecular functionality is found in module rdkit.Chem:

```
>>> from rdkit import Chem
```

Individual molecules can be constructed using a variety of approaches:

```
>>> m = Chem.MolFromSmiles('Cclcccc1')
>>> m = Chem.MolFromMolFile('data/input.mol')
>>> stringWithMolData=file('data/input.mol','r').read()
>>> m = Chem.MolFromMolBlock(stringWithMolData)
```

All of these functions return a rdkit.Chem.rdchem.Mol object on success:

```
>>> m <rdkit.Chem.rdchem.Mol object at 0x...>
```

or None on failure:

```
>>> m = Chem.MolFromMolFile('data/invalid.mol')
>>> m is None
True
```

An attempt is made to provide sensible error messages:

```
>>> m1 = Chem.MolFromSmiles('CO(C)C')
```

displays a message like: [12:18:01] Explicit valence for atom # 1 O greater than permitted and

```
>>> m2 = Chem.MolFromSmiles('clccl')
displays something like: [12:20:41] Can't kekulize mol. In each case the value None is returned:
>>> m1 is None
True
>>> m2 is None
True
```

3.2.2 Reading sets of molecules

Groups of molecules are read using a Supplier (for example, an rdkit.Chem.rdmolfiles.SDMolSupplier or a rdkit.Chem.rdmolfiles.SmilesMolSupplier):

You can easily produce lists of molecules from a Supplier:

```
>>> mols = [x for x in suppl]
>>> len(mols)
4
```

or just treat the Supplier itself as a random-access object:

```
>>> suppl[0].GetNumAtoms()
20
```

A good practice is to test each molecule to see if it was correctly read before working with it:

```
>>> suppl = Chem.SDMolSupplier('data/5ht3ligs.sdf')
>>> for mol in suppl:
... if mol is None: continue
... print mol.GetNumAtoms()
...
20
24
24
```

An alternate type of Supplier, the rdkit.Chem.rdmolfiles.ForwardSDMolSupplier can be used to read from file-like objects:

```
>>> inf = file('data/5ht3ligs.sdf')
>>> fsuppl = Chem.ForwardSDMolSupplier(inf)
>>> for mol in fsuppl:
... if mol is None: continue
... print mol.GetNumAtoms()
...
20
24
24
26
```

This means that they can be used to read from compressed files:

```
>>> import gzip
>>> inf = gzip.open('data/actives_5ht3.sdf.gz')
>>> gzsuppl = Chem.ForwardSDMolSupplier(inf)
>>> ms = [x for x in gzsuppl if x is not None]
>>> len(ms)
180
```

Note that ForwardSDMolSuppliers cannot be used as random-access objects:

```
>>> fsuppl[0]
Traceback (most recent call last):
    ...
TypeError: 'ForwardSDMolSupplier' object does not support indexing
```

3.2.3 Writing molecules

Single molecules can be converted to text using several functions present in the rdkit. Chem module.

For example, for SMILES:

```
>>> m = Chem.MolFromMolFile('data/chiral.mol')
>>> Chem.MolToSmiles(m)
'CC(0)clcccc1'
>>> Chem.MolToSmiles(m,isomericSmiles=True)
'C[C@H](0)clcccc1'
```

Note that the SMILES provided is canonical, so the output should be the same no matter how a particular molecule is input:

```
>>> Chem.MolToSmiles(Chem.MolFromSmiles('C1=CC=CN=C1'))
'c1ccncc1'
>>> Chem.MolToSmiles(Chem.MolFromSmiles('c1ccnc1'))
'c1ccncc1'
>>> Chem.MolToSmiles(Chem.MolFromSmiles('n1cccc1'))
'c1ccncc1'
```

If you'd like to have the Kekule form of the SMILES, first Kekulize the molecule, then use the "kekuleSmiles" option:

```
>>> Chem.Kekulize(m)
>>> Chem.MolToSmiles(m,kekuleSmiles=True)
'CC(0)C1=CC=CC=C1'
```

Note: as of this writing (Aug 2008), the smiles provided when one requests kekuleSmiles are not canonical. The limitation is not in the SMILES generation, but in the kekulization itself.

MDL Mol blocks are also available:

```
1 2 1 0
2 3 1 0
3 4 1 0
4 1 1 0
M END
```

To include names in the mol blocks, set the molecule's "_Name" property:

```
>>> m2.SetProp("_Name", "cyclobutane")
>>> print Chem.MolToMolBlock(m2)
cyclobutane
    RDKit
 4 4 0 0 0 0 0 0 0 0 0999 V2000
   0.0000
            0.0000
                    0.0000 C
                               0 0
                                     0 0
                                          0
                                             0
                                                0
                                                  0
                                                     0
   0.0000
            0.0000
                     0.0000 C
                               0 0 0
                                       0
                                          0
                                             0
                                                0
                                                   0
                                                     0
                                                        0
   0.0000
            0.0000
                     0.0000 C
                               0
                                  0
                                    0
                                       0
                                          0
                                             0
                                                0
                                                   0
                                                     0
            0.0000
                   0.0000 C
                               0 0 0 0
                                          0
                                             0
                                               0
                                                  0
   0.0000
 1 2 1 0
 2 3 1 0
 3 4 1 0
 4 1 1 0
M END
```

It's usually preferable to have a depiction in the Mol block, this can be generated using functionality in the rdkit.Chem.AllChem module (see the Chem vs AllChem section for more information).

You can either include 2D coordinates (i.e. a depiction):

```
>>> from rdkit.Chem import AllChem
>>> AllChem.Compute2DCoords(m2)
>>> print Chem.MolToMolBlock(m2)
cyclobutane
   RDKit
               2D
 4 4 0 0 0 0 0 0 0 0 0999 V2000
   1.0607 -0.0000 0.0000 C 0 0 0 0 0 0 0 0 0 0 0
  -0.0000 -1.0607 0.0000 C 0 0 0 0 0 0 0 0 0 0 0
  -1.0607 0.0000 0.0000 C 0 0 0 0 0 0 0 0 0 0 0
   0.0000 1.0607 0.0000 C 0 0 0 0 0 0 0 0 0 0
 1 2 1 0
 2 3 1 0
 3 4 1 0
 4 1 1 0
M END
```

Or you can add 3D coordinates by embedding the molecule:

The optimization step isn't necessary, but it substantially improves the quality of the conformation.

To get good conformations, it's almost always a good idea to add hydrogens to the molecule first:

```
>>> m3 = Chem.AddHs(m2)
>>> AllChem.EmbedMolecule(m3)
0
>>> AllChem.UFFOptimizeMolecule(m3)
0
```

These can then be removed:

```
>>> m3 = Chem.RemoveHs(m3)
>>> print Chem.MolToMolBlock (m3)
cyclobutane
   RDKit
             3D
 4 4 0 0 0 0 0 0 0 0 0999 V2000
  1.0352 -0.2833 0.0743 C
                      0 0 0 0 0 0 0 0 0 0 0
 -0.2851 -1.0372 0.0171 C 0 0 0 0 0 0 0 0 0 0 0
 -1.0352 0.2833 -0.0743 C 0 0 0 0 0 0 0 0 0 0 0
 1 2 1 0
 2 3 1 0
 3 4 1 0
 4 1 1 0
M END
```

If you'd like to write the molecules to a file, use Python file objects:

```
>>> print >>file('data/foo.mol','w+'),Chem.MolToMolBlock(m2)
>>>
```

3.2.4 Writing sets of molecules

Multiple molecules can be written to a file using an rdkit.Chem.rdmolfiles.SDWriter object:

```
>>> w = Chem.SDWriter('data/foo.sdf')
>>> for m in mols: w.write(m)
...
>>>
```

An SDWriter can also be initialized using a file-like object:

```
>>> from StringIO import StringIO
>>> sio = StringIO()
>>> w = Chem.SDWriter(sio)
>>> for m in mols: w.write(m)
...
>>> w.flush()
>>> print sio.getvalue()
```

```
mol-295
   RDKit
               3D
20 22 0 0 1 0 0 0 0 0999 V2000
   2.3200 0.0800
                -0.1000 C 0 0 0 0 0 0 0 0 0 0 0
   1.8400
                0.1200 C 0 0 0 0 0 0
        -1.2200
                                        0
                                          0
                                             0
                                               0
 1 3 1 0
 1 4 1 0
 2 5 1 0
M END
$$$$
```

Other available Writers include the rdkit.Chem.rdmolfiles.SmilesWriter and the rdkit.Chem.rdmolfiles.TDTWriter.

3.3 Working with Molecules

3.3.1 Looping over Atoms and Bonds

Once you have a molecule, it's easy to loop over its atoms and bonds:

```
>>> m = Chem.MolFromSmiles('C10C1')
>>> for atom in m.GetAtoms():
... print atom.GetAtomicNum()
...
6
8
6
>>> print m.GetBonds()[0].GetBondType()
SINGLE
```

You can also request individual bonds or atoms:

```
>>> m.GetAtomWithIdx(0).GetSymbol()
'C'
>>> m.GetAtomWithIdx(0).GetExplicitValence()
2
>>> m.GetBondWithIdx(0).GetBeginAtomIdx()
0
>>> m.GetBondWithIdx(0).GetEndAtomIdx()
1
>>> m.GetBondBetweenAtoms(0,1).GetBondType()
rdkit.Chem.rdchem.BondType.SINGLE
```

Atoms keep track of their neighbors:

```
>>> atom = m.GetAtomWithIdx(0)
>>> [x.GetAtomicNum() for x in atom.GetNeighbors()]
[8, 6]
>>> len(x.GetBonds())
```

3.3.2 Ring Information

Atoms and bonds both carry information about the molecule's rings:

```
>>> m = Chem.MolFromSmiles('OC1C2C1CC2')
>>> m.GetAtomWithIdx(0).IsInRing()
False
>>> m.GetAtomWithIdx(1).IsInRing()
True
>>> m.GetAtomWithIdx(2).IsInRingSize(3)
True
>>> m.GetAtomWithIdx(2).IsInRingSize(4)
True
>>> m.GetAtomWithIdx(2).IsInRingSize(5)
False
>>> m.GetBondWithIdx(1).IsInRingSize(3)
True
>>> m.GetBondWithIdx(1).IsInRingSize(3)
```

But note that the information is only about the smallest rings:

```
>>> m.GetAtomWithIdx(1).IsInRingSize(5)
False
```

More detail about the smallest set of smallest rings (SSSR) is available:

```
>>> ssr = Chem.GetSymmSSSR(m)
>>> len(ssr)
2
>>> list(ssr[0])
[1, 2, 3]
>>> list(ssr[1])
[4, 5, 2, 3]
```

As the name indicates, this is a symmetrized SSSR; if you are interested in the number of "true" SSSR, use the GetSSSR function.

```
>>> Chem.GetSSSR(m)
2
```

The distinction between symmetrized and non-symmetrized SSSR is discussed in more detail below in the section The SSSR Problem.

For more efficient queries about a molecule's ring systems (avoiding repeated calls to Mol.GetAtomWithIdx), use the rdkit.Chem.rdchem.RingInfo class:

```
>>> m = Chem.MolFromSmiles('OC1C2C1CC2')
>>> ri = m.GetRingInfo()
>>> ri.NumAtomRings(0)
0
>>> ri.NumAtomRings(1)
1
>>> ri.NumAtomRings(2)
2
>>> ri.IsAtomInRingOfSize(1,3)
True
>>> ri.IsBondInRingOfSize(1,3)
True
```

3.3.3 Modifying molecules

Normally molecules are stored in the RDKit with the hydrogen atoms implicit (e.g. not explicitly present in the molecular graph. When it is useful to have the hydrogens explicitly present, for example when generating or optimizing the 3D geometry, the rdkit.Chem.rdmolops.AddHs function can be used:

```
>>> m=Chem.MolFromSmiles('CCO')
>>> m.GetNumAtoms()
3
>>> m2 = Chem.AddHs(m)
>>> m2.GetNumAtoms()
```

The Hs can be removed again using the rdkit. Chem. rdmolops. RemoveHs function:

```
>>> m3 = Chem.RemoveHs(m2)
>>> m3.GetNumAtoms()
3
```

RDKit molecules are usually stored with the bonds in aromatic rings having aromatic bond types. This can be changed with the rdkit.Chem.rdmolops.Kekulize function:

```
>>> m = Chem.MolFromSmiles('clcccc1')
>>> m.GetBondWithIdx(0).GetBondType()
rdkit.Chem.rdchem.BondType.AROMATIC
>>> Chem.Kekulize(m)
>>> m.GetBondWithIdx(0).GetBondType()
rdkit.Chem.rdchem.BondType.DOUBLE
>>> m.GetBondWithIdx(1).GetBondType()
rdkit.Chem.rdchem.BondType.SINGLE
```

The bonds are still marked as being aromatic:

```
>>> m.GetBondWithIdx(1).GetIsAromatic()
True
```

and can be restored to the aromatic bond type using the rdkit.Chem.rdmolops.SanitizeMol function:

```
>>> Chem.SanitizeMol(m)
rdkit.Chem.rdmolops.SanitizeFlags.SANITIZE_NONE
>>> m.GetBondWithIdx(0).GetBondType()
rdkit.Chem.rdchem.BondType.AROMATIC
```

The value returned by SanitizeMol() indicates that no problems were encountered.

3.3.4 Working with 2D molecules: Generating Depictions

The RDKit has a library for generating depictions (sets of 2D) coordinates for molecules. This library, which is part of the AllChem module, is accessed using the rdkit.Chem.rdDepictor.Compute2DCoords function:

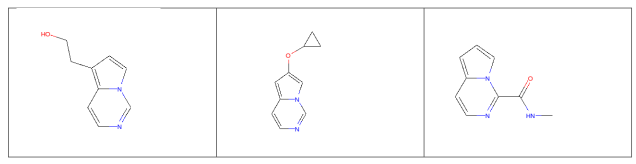
```
>>> m = Chem.MolFromSmiles('clnccc2nlccc2')
>>> AllChem.Compute2DCoords(m)
0
```

The 2D conformation is constructed in a canonical orientation and is built to minimize intramolecular clashes, i.e. to maximize the clarity of the drawing.

If you have a set of molecules that share a common template and you'd like to align them to that template, you can do so as follows:

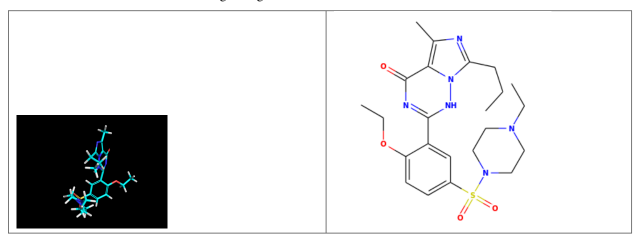
```
>>> template = Chem.MolFromSmiles('clnccc2nlccc2')
>>> AllChem.Compute2DCoords(template)
0
>>> AllChem.GenerateDepictionMatching2DStructure(m,template)
```

Running this process for a couple of other molecules gives the following depictions:



Compute2DCoords Another option for allows VOII to generate 2D depictions molecules that closely mimic 3D conformations. This available using the function rdkit.Chem.AllChem.GenerateDepictionMatching3DStructure.

Here is an illustration of the results using the ligand from PDB structure 1XP0:



More fine-grained control can be obtained using the core function rdkit.Chem.rdDepictor.Compute2DCoordsMimicDistm but that is beyond the scope of this document. See the implementation of GenerateDepictionMatching3DStructure in AllChem.py for an example of how it is used.

3.3.5 Working with 3D Molecules

The RDKit can generate conformations for molecules using distance geometry. ¹ The algorithm followed is:

- 1. The molecule's distance bounds matrix is calculated based on the connection table and a set of rules.
- 2. The bounds matrix is smoothed using a triangle-bounds smoothing algorithm.
- 3. A random distance matrix that satisfies the bounds matrix is generated.
- 4. This distance matrix is embedded in 3D dimensions (producing coordinates for each atom).
- 5. The resulting coordinates are cleaned up somewhat using a crude force field and the bounds matrix.

¹ Blaney, J. M.; Dixon, J. S. "Distance Geometry in Molecular Modeling". Reviews in Computational Chemistry; VCH: New York, 1994.

Multiple conformations can be generated by repeating steps 4 and 5 several times, using a different random distance matrix each time.

Note that the conformations that result from this procedure tend to be fairly ugly. They should be cleaned up using a force field. This can be done within the RDKit using its implementation of the Universal Force Field (UFF). ²

The full process of embedding and optimizing a molecule is easier than all the above verbiage makes it sound:

```
>>> m = Chem.MolFromSmiles('C1CCC1OC')
>>> m2=Chem.AddHs(m)
>>> AllChem.EmbedMolecule(m2)
0
>>> AllChem.UFFOptimizeMolecule(m2)
0
```

The RDKit also has an implementation of the MMFF94 force field available. ³, ⁴, ⁵, ⁶, ⁷ Please note that the MMFF atom typing code uses its own aromaticity model, so the aromaticity flags of the molecule will be modified after calling MMFF-related methods.

```
>>> m = Chem.MolFromSmiles('C1CCC1OC')
>>> m2=Chem.AddHs(m)
>>> AllChem.EmbedMolecule(m2)
0
>>> AllChem.MMFFOptimizeMolecule(m2)
0
```

Note the calls to *Chem.AddHs()* in the examples above. By default RDKit molecules do not have H atoms explicity present in the graph, but they are important for getting realistic geometries, so they generally should be added.

With the RDKit, also multiple conformers can be generated. The option numConfs allows the user to set the number of conformers that should be generated. These conformers can be aligned to each other and the RMS values calculated.

rmslist contains the RMS values between the first conformer and all others. The RMS between two specific conformers (e.g. 1 and 9) can also be calculated. The flag prealigned lets the user specify if the conformers are already aligned (by default, the function aligns them).

```
>>> rms = AllChem.GetConformerRMS(m2, 1, 9, prealigned=True)
```

² Rappé, A. K.; Casewit, C. J.; Colwell, K. S.; Goddard III, W. A.; Skiff, W. M. "UFF, a full periodic table force field for molecular mechanics and molecular dynamics simulations". *J. Am. Chem. Soc.* **114**:10024-35 (1992).

³ Halgren, T. A. "Merck molecular force field. I. Basis, form, scope, parameterization, and performance of MMFF94." *J. Comp. Chem.* **17**:490–19 (1996).

⁴ Halgren, T. A. "Merck molecular force field. II. MMFF94 van der Waals and electrostatic parameters for intermolecular interactions." *J. Comp. Chem.* **17**:520–52 (1996).

⁵ Halgren, T. A. "Merck molecular force field. III. Molecular geometries and vibrational frequencies for MMFF94." *J. Comp. Chem.* 17:553–86 (1996)

⁶ Halgren, T. A. & Nachbar, R. B. "Merck molecular force field. IV. conformational energies and geometries for MMFF94." *J. Comp. Chem.* 17:587 615 (1996)

⁷ Halgren, T. A. "MMFF VI. MMFF94s option for energy minimization studies." *J. Comp. Chem.* **20**:720–9 (1999).

More 3D functionality of the RDKit is described in the Cookbook.

Disclaimer/Warning: Conformation generation is a difficult and subtle task. The 2D->3D conversion provided within the RDKit is not intended to be a replacement for a "real" conformational analysis tool; it merely provides quick 3D structures for cases when they are required.

3.3.6 Preserving Molecules

Molecules can be converted to and from text using Python's pickling machinery:

```
>>> m = Chem.MolFromSmiles('clccnccl')
>>> import cPickle
>>> pkl = cPickle.dumps(m)
>>> type(pkl)
<type 'str'>
>>> m2=cPickle.loads(pkl)
>>> Chem.MolToSmiles(m2)
'clccnccl'
```

The RDKit pickle format is fairly compact and it is much, much faster to build a molecule from a pickle than from a Mol file or SMILES string, so storing molecules you will be working with repeatedly as pickles can be a good idea.

The raw binary data that is encapsulated in a pickle can also be directly obtained from a molecule:

```
>>> binStr = m.ToBinary()
```

This can be used to reconstruct molecules using the Chem.Mol constructor:

```
>>> m2 = Chem.Mol(binStr)
>>> Chem.MolToSmiles(m2)
'clccncc1'
>>> len(binStr)
123
>>> len(pkl)
475
```

Note that this huge difference in text length is because we didn't tell python to use its most efficient representation of the pickle:

```
>>> pkl = cPickle.dumps(m,2)
>>> len(pkl)
157
```

The small overhead associated with python's pickling machinery normally doesn't end up making much of a difference for collections of larger molecules (the extra data associated with the pickle is independent of the size of the molecule, while the binary string increases in length as the molecule gets larger).

Tip: The performance difference associated with storing molecules in a pickled form on disk instead of constantly reparsing an SD file or SMILES table is difficult to overstate. In a test I just ran on my laptop, loading a set of 699 drug-like molecules from an SD file took 10.8 seconds; loading the same molecules from a pickle file took 0.7 seconds. The pickle file is also smaller – 1/3 the size of the SD file – but this difference is not always so dramatic (it's a particularly fat SD file).

3.3.7 Drawing Molecules

The RDKit has some built-in functionality for creating images from molecules found in the rdkit.Chem.Draw package:

```
>>> suppl = Chem.SDMolSupplier('data/cdk2.sdf')
>>> ms = [x for x in suppl if x is not None]
>>> for m in ms: tmp=AllChem.Compute2DCoords(m)
>>> from rdkit.Chem import Draw
>>> Draw.MolToFile(ms[0],'images/cdk2_mol1.png')
>>> Draw.MolToFile(ms[1],'images/cdk2_mol2.png')
```

Producing these images:

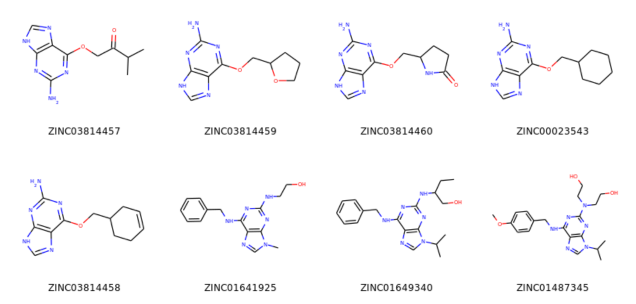
It's also possible to produce an image grid out of a set of molecules:

```
>>> img=Draw.MolsToGridImage(ms[:8],molsPerRow=4,subImgSize=(200,200),legends=[x.GetProp("_Name") for
```

This returns a PIL image, which can then be saved to a file:

```
>>> img.save('images/cdk2_molgrid.png')
```

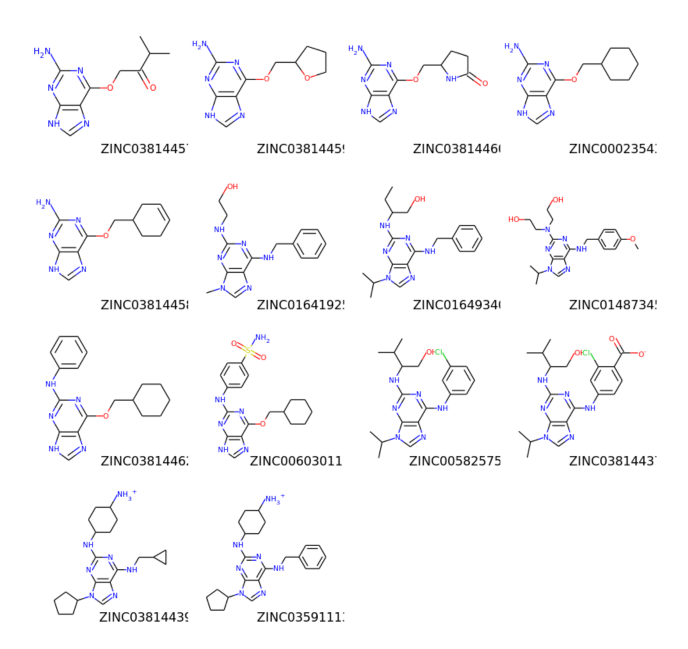
The result looks like this:



These would of course look better if the common core were aligned. This is easy enough to do:

```
>>> p = Chem.MolFromSmiles('[nH]1cnc2cncnc21')
>>> subms = [x for x in ms if x.HasSubstructMatch(p)]
>>> len(subms)
14
>>> AllChem.Compute2DCoords(p)
0
>>> for m in subms: AllChem.GenerateDepictionMatching2DStructure(m,p)
>>> img=Draw.MolsToGridImage(subms,molsPerRow=4,subImgSize=(200,200),legends=[x.GetProp("_Name") for
>>> img.save('images/cdk2_molgrid.aligned.png')
```

The result looks like this:



3.4 Substructure Searching

Substructure matching can be done using query molecules built from SMARTS:

```
>>> m = Chem.MolFromSmiles('clcccc10')
>>> patt = Chem.MolFromSmarts('cc0')
>>> m.HasSubstructMatch(patt)
True
>>> m.GetSubstructMatch(patt)
(0, 5, 6)
```

Those are the atom indices in m, ordered as patt's atoms. To get all of the matches:

```
>>> m.GetSubstructMatches(patt) ((0, 5, 6), (4, 5, 6))
```

This can be used to easily filter lists of molecules:

```
>>> suppl = Chem.SDMolSupplier('data/actives_5ht3.sdf')
>>> patt = Chem.MolFromSmarts('c[NH1]')
>>> matches = []
>>> for mol in suppl:
... if mol.HasSubstructMatch(patt):
... matches.append(mol)
...
>>> len(matches)
```

We can write the same thing more compactly using Python's list comprehension syntax:

```
>>> matches = [x for x in suppl if x.HasSubstructMatch(patt)]
>>> len(matches)
22
```

Substructure matching can also be done using molecules built from SMILES instead of SMARTS:

```
>>> m = Chem.MolFromSmiles('C1=CC=CC=C1OC')
>>> m.HasSubstructMatch(Chem.MolFromSmarts('CO'))
True
>>> m.HasSubstructMatch(Chem.MolFromSmiles('CO'))
True
```

But don't forget that the semantics of the two languages are not exactly equivalent:

```
>>> m.HasSubstructMatch(Chem.MolFromSmiles('COC'))
True
>>> m.HasSubstructMatch(Chem.MolFromSmarts('COC'))
False
>>> m.HasSubstructMatch(Chem.MolFromSmarts('COC')) #<- need an aromatic C
True</pre>
```

3.4.1 Stereochemistry in substructure matches

By default information about stereochemistry is not used in substructure searches:

```
>>> m = Chem.MolFromSmiles('CC[C@H](F)Cl')
>>> m.HasSubstructMatch(Chem.MolFromSmiles('C[C@H](F)Cl'))
True
>>> m.HasSubstructMatch(Chem.MolFromSmiles('C[C@@H](F)Cl'))
True
>>> m.HasSubstructMatch(Chem.MolFromSmiles('CC(F)Cl'))
True
```

But this can be changed via the *useChirality* argument:

```
>>> m.HasSubstructMatch(Chem.MolFromSmiles('C[C@H](F)Cl'), useChirality=True)
True
>>> m.HasSubstructMatch(Chem.MolFromSmiles('C[C@@H](F)Cl'), useChirality=True)
False
>>> m.HasSubstructMatch(Chem.MolFromSmiles('CC(F)Cl'), useChirality=True)
True
```

Notice that when *useChirality* is set a non-chiral query **does** match a chiral molecule. The same is not true for a chiral query and a non-chiral molecule:

```
>>> m.HasSubstructMatch(Chem.MolFromSmiles('CC(F)Cl'))
True
>>> m2 = Chem.MolFromSmiles('CCC(F)Cl')
>>> m2.HasSubstructMatch(Chem.MolFromSmiles('C[C@H](F)Cl'),useChirality=True)
False
```

3.5 Chemical Transformations

The RDKit contains a number of functions for modifying molecules. Note that these transformation functions are intended to provide an easy way to make simple modifications to molecules. For more complex transformations, use the Chemical Reactions functionality.

3.5.1 Substructure-based transformations

There's a variety of functionality for using the RDKit's substructure-matching machinery for doing quick molecular transformations. These transformations include deleting substructures:

```
>>> m = Chem.MolFromSmiles('CC(=0)0')
>>> patt = Chem.MolFromSmarts('C(=0)[OH]')
>>> rm = AllChem.DeleteSubstructs(m,patt)
>>> Chem.MolToSmiles(rm)
'C'
```

replacing substructures:

```
>>> repl = Chem.MolFromSmiles('OC')
>>> patt = Chem.MolFromSmarts('[$(NC(=0))]')
>>> m = Chem.MolFromSmiles('CC(=0)N')
>>> rms = AllChem.ReplaceSubstructs(m,patt,repl)
>>> rms
(<rdkit.Chem.rdchem.Mol object at 0x...>,)
>>> Chem.MolToSmiles(rms[0])
'COC(C)=O'
```

as well as simple SAR-table transformations like removing side chains:

```
>>> m1 = Chem.MolFromSmiles('BrCCclcncnc1C(=0)0')
>>> core = Chem.MolFromSmiles('clcncnc1')
>>> tmp = Chem.ReplaceSidechains(m1,core)
>>> Chem.MolToSmiles(tmp)
'[*]clcncnc1[*]'
```

and removing cores:

```
>>> tmp = Chem.ReplaceCore(m1,core)
>>> Chem.MolToSmiles(tmp)
'[*]C(=0)0.[*]CCBr'
```

To get more detail about the sidechains (e.g. sidechain labels), use isomeric smiles:

```
>>> Chem.MolToSmiles(tmp,True)
'[1*]CCBr.[2*]C(=0)0'
```

By default the sidechains are labeled based on the order they are found. They can also be labeled according by the number of that core-atom they're attached to:

```
>>> m1 = Chem.MolFromSmiles('clc(CCO)ncnc1C(=0)0')
>>> tmp=Chem.ReplaceCore(m1,core,labelByIndex=True)
>>> Chem.MolToSmiles(tmp,True)
'[1*]CCO.[5*]C(=0)0'
```

rdkit.Chem.rdmolops.ReplaceCore returns the sidechains in a single molecule. This can be split into separate molecules using rdkit.Chem.rdmolops.GetMolFrags:

```
>>> rs = Chem.GetMolFrags(tmp,asMols=True)
>>> len(rs)
2
>>> Chem.MolToSmiles(rs[0],True)
'[1*]CCO'
>>> Chem.MolToSmiles(rs[1],True)
'[5*]C(=0)O'
```

3.5.2 Murcko Decomposition

The RDKit provides standard Murcko-type decomposition ⁸ of molecules into scaffolds:

```
>>> from rdkit.Chem.Scaffolds import MurckoScaffold
>>> cdk2mols = Chem.SDMolSupplier('data/cdk2.sdf')
>>> m1 = cdk2mols[0]
>>> core = MurckoScaffold.GetScaffoldForMol(m1)
>>> Chem.MolToSmiles(core)
'clncc2nc[nH]c2n1'
or into a generic framework:
>>> fw = MurckoScaffold.MakeScaffoldGeneric(core)
>>> Chem.MolToSmiles(fw)
'clccc2cccc2c1'
```

3.6 Maximum Common Substructure

The FindMCS function find a maximum common substructure (MCS) of two or more molecules:

```
>>> from rdkit.Chem import rdFMCS
>>> mol1 = Chem.MolFromSmiles("O=C(NCc1cc(OC)c(O)cc1)CCCC/C=C/C(C)C")
>>> mol2 = Chem.MolFromSmiles("CC(C)CCCCCC(=O)NCC1=CC(=C(C=C1)O)OC")
>>> mol3 = Chem.MolFromSmiles("c1(C=O)cc(OC)c(O)cc1")
>>> mols = [mol1,mol2,mol3]
>>> res=rdFMCS.FindMCS(mols)
>>> res

<pre
```

⁸ Bemis, G. W.; Murcko, M. A. "The Properties of Known Drugs. 1. Molecular Frameworks." J. Med. Chem. 39:2887-93 (1996).

```
>>> res.canceled False
```

It returns an MCSResult instance with information about the number of atoms and bonds in the MCS, the SMARTS string which matches the identified MCS, and a flag saying if the algorithm timed out. If no MCS is found then the number of atoms and bonds is set to 0 and the SMARTS to ''.

By default, two atoms match if they are the same element and two bonds match if they have the same bond type. Specify atomCompare and bondCompare to use different comparison functions, as in:

```
>>> mols = (Chem.MolFromSmiles('NCC'), Chem.MolFromSmiles('OC=C'))
>>> rdFMCS.FindMCS(mols).smartsString
''
>>> rdFMCS.FindMCS(mols, atomCompare=rdFMCS.AtomCompare.CompareAny).smartsString
'[#7,#8]-[#6]'
>>> rdFMCS.FindMCS(mols, bondCompare=rdFMCS.BondCompare.CompareAny).smartsString
'[#6]-,=[#6]'
```

The options for the atomCompare argument are: CompareAny says that any atom matches any other atom, CompareElements compares by element type, and CompareIsotopes matches based on the isotope label. Isotope labels can be used to implement user-defined atom types. A bondCompare of CompareAny says that any bond matches any other bond, CompareOrderExact says bonds are equivalent if and only if they have the same bond type, and CompareOrder allows single and aromatic bonds to match each other, but requires an exact order match otherwise:

```
>>> mols = (Chem.MolFromSmiles('clcccc1'), Chem.MolFromSmiles('ClCCCC=C1'))
>>> rdFMCS.FindMCS(mols, bondCompare=rdFMCS.BondCompare.CompareAny).smartsString
'[#6]1:,-[#6]:,-[#6]:,-[#6]:,-[#6]:,-[#6]:,-1'
>>> rdFMCS.FindMCS(mols, bondCompare=rdFMCS.BondCompare.CompareOrderExact).smartsString
','
>>> rdFMCS.FindMCS(mols, bondCompare=rdFMCS.BondCompare.CompareOrder).smartsString
'[#6](:,-[#6]:,-[#6]:,-[#6]:,-[#6]:,-[#6]):,-[#6]'
```

A substructure has both atoms and bonds. By default, the algorithm attempts to maximize the number of bonds found. You can change this by setting the maximizeBonds argument to False. Maximizing the number of bonds tends to maximize the number of rings, although two small rings may have fewer bonds than one large ring.

You might not want a 3-valent nitrogen to match one which is 5-valent. The default matchValences value of False ignores valence information. When True, the atomCompare setting is modified to also require that the two atoms have the same valency.

```
>>> mols = (Chem.MolFromSmiles('NC10C1'), Chem.MolFromSmiles('C10C1[N+](=0)[0-]'))
>>> rdFMCS.FindMCS(mols).numAtoms
4
>>> rdFMCS.FindMCS(mols, matchValences=True).numBonds
2
```

It can be strange to see a linear carbon chain match a carbon ring, which is what the ringMatchesRingOnly default of False does. If you set it to True then ring bonds will only match ring bonds.

```
>>> mols = [Chem.MolFromSmiles("C1CCC1CCC"), Chem.MolFromSmiles("C1CCCCC1")]
>>> rdFMCS.FindMCS(mols).smartsString
'[#6](-[#6]-[#6])-[#6]-[#6]-[#6]'
>>> rdFMCS.FindMCS(mols, ringMatchesRingOnly=True).smartsString
'[#6](-[#6]-[#6])-[#6]'
```

You can further restrict things and require that partial rings (as in this case) are not allowed. That is, if an atom is part of the MCS and the atom is in a ring of the entire molecule then that atom is also in a ring of the MCS. Set completeRingsOnly to True to toggle this requirement and also sets ringMatchesRingOnly to True.

```
>>> mols = [Chem.MolFromSmiles("CCC1CC2C1CN2"), Chem.MolFromSmiles("C1CC2C1CC2")]
>>> rdFMCS.FindMCS(mols).smartsString
'[#6]1-[#6]-[#6]-[#6]-[#6]-[#6])-[#6]'
>>> rdFMCS.FindMCS(mols, ringMatchesRingOnly=True).smartsString
'[#6](-[#6]-[#6]-[#6]-[#6])-[#6]'
>>> rdFMCS.FindMCS(mols, completeRingsOnly=True).smartsString
'[#6]1-[#6]-[#6]-[#6]-[#6]-1'
```

The MCS algorithm will exhaustively search for a maximum common substructure. Typically this takes a fraction of a second, but for some comparisons this can take minutes or longer. Use the timeout parameter to stop the search after the given number of seconds (wall-clock seconds, not CPU seconds) and return the best match found in that time. If timeout is reached then the canceled property of the MCSResult will be True instead of False.

```
>>> mols = [Chem.MolFromSmiles("Nclccccc1"*100), Chem.MolFromSmiles("Nclcccccccc1"*100)]
>>> rdFMCS.FindMCS(mols, timeout=1).canceled
True
```

(The MCS after 50 seconds contained 511 atoms.)

3.7 Fingerprinting and Molecular Similarity

The RDKit has a variety of built-in functionality for generating molecular fingerprints and using them to calculate molecular similarity.

3.7.1 Topological Fingerprints

```
>>> from rdkit import DataStructs
>>> from rdkit.Chem.Fingerprints import FingerprintMols
>>> ms = [Chem.MolFromSmiles('CCOC'), Chem.MolFromSmiles('CCO'),
... Chem.MolFromSmiles('COC')]
>>> fps = [FingerprintMols.FingerprintMol(x) for x in ms]
>>> DataStructs.FingerprintSimilarity(fps[0],fps[1])
0.6...
>>> DataStructs.FingerprintSimilarity(fps[0],fps[2])
0.4...
>>> DataStructs.FingerprintSimilarity(fps[1],fps[2])
0.25
```

The fingerprinting algorithm used is similar to that used in the Daylight fingerprinter: it identifies and hashes topological paths (e.g. along bonds) in the molecule and then uses them to set bits in a fingerprint of user-specified lengths. After all paths have been identified, the fingerprint is typically folded down until a particular density of set bits is obtained.

The default set of parameters used by the fingerprinter is: - minimum path size: 1 bond - maximum path size: 7 bonds - fingerprint size: 2048 bits - number of bits set per hash: 2 - minimum fingerprint size: 64 bits - target on-bit density 0.3

You can control these by calling rdkit.Chem.rdmolops.RDKFingerprint directly; this will return an unfolded fingerprint that you can then fold to the desired density. The function rdkit.Chem.Fingerprints.FingerprintMols.FingerprintMol (written in python) shows how this is done.

The default similarity metric used by rdkit.DataStructs.FingerprintSimilarity is the Tanimoto similarity. One can use different similarity metrics:

```
>>> DataStructs.FingerprintSimilarity(fps[0],fps[1], metric=DataStructs.DiceSimilarity)
0.75
```

Available similarity metrics include Tanimoto, Dice, Cosine, Sokal, Russel, Kulczynski, McConnaughey, and Tversky.

3.7.2 MACCS Keys

There is a SMARTS-based implementation of the 166 public MACCS keys.

```
>>> from rdkit.Chem import MACCSkeys
>>> fps = [MACCSkeys.GenMACCSKeys(x) for x in ms]
>>> DataStructs.FingerprintSimilarity(fps[0],fps[1])
0.5
>>> DataStructs.FingerprintSimilarity(fps[0],fps[2])
0.538...
>>> DataStructs.FingerprintSimilarity(fps[1],fps[2])
0.214...
```

The MACCS keys were critically evaluated and compared to other MACCS implementations in Q3 2008. In cases where the public keys are fully defined, things looked pretty good.

3.7.3 Atom Pairs and Topological Torsions

Atom-pair descriptors ⁹ are available in several different forms. The standard form is as fingerprint including counts for each bit instead of just zeros and ones:

```
>>> from rdkit.Chem.AtomPairs import Pairs
>>> ms = [Chem.MolFromSmiles('C1CCC1OCC'), Chem.MolFromSmiles('CC(C)OCC'), Chem.MolFromSmiles('CCCCC')
>>> pairFps = [Pairs.GetAtomPairFingerprint(x) for x in ms]
```

Because the space of bits that can be included in atom-pair fingerprints is huge, they are stored in a sparse manner. We can get the list of bits and their counts for each fingerprint as a dictionary:

```
>>> d = pairFps[-1].GetNonzeroElements()
>>> d[541732]
1
>>> d[1606690]
2
```

Descriptions of the bits are also available:

```
>>> Pairs.ExplainPairScore(558115)
(('C', 1, 0), 3, ('C', 2, 0))
```

The above means: C with 1 neighbor and 0 pi electrons which is 3 bonds from a C with 2 neighbors and 0 pi electrons

The usual metric for similarity between atom-pair fingerprints is Dice similarity:

```
>>> from rdkit import DataStructs
>>> DataStructs.DiceSimilarity(pairFps[0],pairFps[1])
0.333...
>>> DataStructs.DiceSimilarity(pairFps[0],pairFps[2])
0.258...
>>> DataStructs.DiceSimilarity(pairFps[1],pairFps[2])
0.56
```

⁹ Carhart, R.E.; Smith, D.H.; Venkataraghavan R. "Atom Pairs as Molecular Features in Structure-Activity Studies: Definition and Applications" J. Chem. Inf. Comp. Sci. 25:64-73 (1985).

It's also possible to get atom-pair descriptors encoded as a standard bit vector fingerprint (ignoring the count information):

```
>>> pairFps = [Pairs.GetAtomPairFingerprintAsBitVect(x) for x in ms]
```

Since these are standard bit vectors, the rdkit.DataStructs module can be used for similarity:

```
>>> from rdkit import DataStructs
>>> DataStructs.DiceSimilarity(pairFps[0],pairFps[1])
0.48
>>> DataStructs.DiceSimilarity(pairFps[0],pairFps[2])
0.380...
>>> DataStructs.DiceSimilarity(pairFps[1],pairFps[2])
0.625
```

Topological torsion descriptors ¹⁰ are calculated in essentially the same way:

```
>>> from rdkit.Chem.AtomPairs import Torsions
>>> tts = [Torsions.GetTopologicalTorsionFingerprintAsIntVect(x) for x in ms]
>>> DataStructs.DiceSimilarity(tts[0],tts[1])
0.166...
```

At the time of this writing, topological torsion fingerprints have too many bits to be encodeable using the BitVector machinery, so there is no GetTopologicalTorsionFingerprintAsBitVect function.

3.7.4 Morgan Fingerprints (Circular Fingerprints)

This family of fingerprints, better known as circular fingerprints ¹¹, is built by applying the Morgan algorithm to a set of user-supplied atom invariants. When generating Morgan fingerprints, the radius of the fingerprint must also be provided:

```
>>> from rdkit.Chem import AllChem
>>> m1 = Chem.MolFromSmiles('Cclcccc1')
>>> fp1 = AllChem.GetMorganFingerprint(m1,2)
>>> fp1
<rdkit.DataStructs.cDataStructs.UIntSparseIntVect object at 0x...>
>>> m2 = Chem.MolFromSmiles('Cclnccc1')
>>> fp2 = AllChem.GetMorganFingerprint(m2,2)
>>> DataStructs.DiceSimilarity(fp1,fp2)
0.55...
```

Morgan fingerprints, like atom pairs and topological torsions, use counts by default, but it's also possible to calculate them as bit vectors:

```
>>> fp1 = AllChem.GetMorganFingerprintAsBitVect (m1,2,nBits=1024)
>>> fp1
<rdkit.DataStructs.cDataStructs.ExplicitBitVect object at 0x...>
>>> fp2 = AllChem.GetMorganFingerprintAsBitVect (m2,2,nBits=1024)
>>> DataStructs.DiceSimilarity(fp1,fp2)
0.51...
```

The default atom invariants use connectivity information similar to those used for the well known ECFP family of fingerprints. Feature-based invariants, similar to those used for the FCFP fingerprints, can also be used. The feature definitions used are defined in the section Feature Definitions Used in the Morgan Fingerprints. At times this can lead to quite different similarity scores:

¹⁰ Nilakantan, R.; Bauman N.; Dixon J.S.; Venkataraghavan R. "Topological Torsion: A New Molecular Descriptor for SAR Applications. Comparison with Other Descriptors." *J. Chem.Inf. Comp. Sci.* **27**:82-5 (1987).

¹¹ Rogers, D.; Hahn, M. "Extended-Connectivity Fingerprints." J. Chem. Inf. and Model. 50:742-54 (2010).

```
>>> m1 = Chem.MolFromSmiles('clcccn1')
>>> m2 = Chem.MolFromSmiles('clccco1')
>>> fp1 = AllChem.GetMorganFingerprint(m1,2)
>>> fp2 = AllChem.GetMorganFingerprint(m2,2)
>>> ffp1 = AllChem.GetMorganFingerprint(m1,2,useFeatures=True)
>>> ffp2 = AllChem.GetMorganFingerprint(m2,2,useFeatures=True)
>>> DataStructs.DiceSimilarity(fp1,fp2)
0.36...
>>> DataStructs.DiceSimilarity(ffp1,ffp2)
0.90...
```

When comparing the ECFP/FCFP fingerprints and the Morgan fingerprints generated by the RDKit, remember that the 4 in ECFP4 corresponds to the diameter of the atom environments considered, while the Morgan fingerprints take a radius parameter. So the examples above, with radius=2, are roughly equivalent to ECFP4 and FCFP4.

The user can also provide their own atom invariants using the optional invariants argument to rdkit.Chem.rdMolDescriptors.GetMorganFingerprint. Here's a simple example that uses a constant for the invariant; the resulting fingerprints compare the topology of molecules:

```
>>> m1 = Chem.MolFromSmiles('Cclcccc1')
>>> m2 = Chem.MolFromSmiles('Cclncncn1')
>>> fp1 = AllChem.GetMorganFingerprint(m1,2,invariants=[1]*m1.GetNumAtoms())
>>> fp2 = AllChem.GetMorganFingerprint(m2,2,invariants=[1]*m2.GetNumAtoms())
>>> fp1==fp2
True
```

Note that bond order is by default still considered:

```
>>> m3 = Chem.MolFromSmiles('CC1CCCCC1')
>>> fp3 = AllChem.GetMorganFingerprint(m3,2,invariants=[1]*m3.GetNumAtoms())
>>> fp1==fp3
False
```

But this can also be turned off:

```
>>> fp1 = AllChem.GetMorganFingerprint(m1,2,invariants=[1]*m1.GetNumAtoms(),
... useBondTypes=False)
>>> fp3 = AllChem.GetMorganFingerprint(m3,2,invariants=[1]*m3.GetNumAtoms(),
... useBondTypes=False)
>>> fp1==fp3
True
```

Explaining bits from Morgan Fingerprints

Information is available about the atoms that contribute to particular bits in the Morgan fingerprint via the bitInfo argument. The dictionary provided is populated with one entry per bit set in the fingerprint, the keys are the bit ids, the values are lists of (atom index, radius) tuples.

```
>>> m = Chem.MolFromSmiles('clcccnclC')
>>> info={}
>>> fp = AllChem.GetMorganFingerprint(m,2,bitInfo=info)
>>> len(fp.GetNonzeroElements())
16
>>> len(info)
16
>>> info[98513984]
((1, 1), (2, 1))
```

```
>>> info[4048591891] ((5, 2),)
```

Interpreting the above: bit 98513984 is set twice: once by atom 1 and once by atom 2, each at radius 1. Bit 4048591891 is set once by atom 5 at radius 2.

Focusing on bit 4048591891, we can extract the submolecule consisting of all atoms within a radius of 2 of atom 5:

```
>>> env = Chem.FindAtomEnvironmentOfRadiusN(m,2,5)
>>> amap={}
>>> submol=Chem.PathToSubmol(m,env,atomMap=amap)
>>> submol.GetNumAtoms()
6
>>> amap
{0: 3, 1: 5, 3: 4, 4: 0, 5: 1, 6: 2}
```

And then "explain" the bit by generating SMILES for that submolecule:

```
>>> Chem.MolToSmiles(submol)
'ccc(C)nc'
```

This is more useful when the SMILES is rooted at the central atom:

```
>>> Chem.MolToSmiles(submol,rootedAtAtom=amap[5],canonical=False)
'c(nc)(C)cc'
```

An alternate (and faster, particularly for large numbers of molecules) approach to do the same thing, using the function rdkit.Chem.MolFragmentToSmiles:

3.7.5 Picking Diverse Molecules Using Fingerprints

A common task is to pick a small subset of diverse molecules from a larger set. The RDKit provides a number of approaches for doing this in the rdkit.SimDivFilters module. The most efficient of these uses the MaxMin algorithm. 12 Here's an example:

Start by reading in a set of molecules and generating Morgan fingerprints:

```
>>> from rdkit import Chem
>>> from rdkit.Chem.rdMolDescriptors import GetMorganFingerprint
>>> from rdkit import DataStructs
>>> from rdkit.SimDivFilters.rdSimDivPickers import MaxMinPicker
>>> ms = [x for x in Chem.SDMolSupplier('data/actives_5ht3.sdf')]
>>> while ms.count(None): ms.remove(None)
>>> fps = [GetMorganFingerprint(x,3) for x in ms]
>>> nfps = len(fps)
```

The algorithm requires a function to calculate distances between objects, we'll do that using DiceSimilarity:

¹² Ashton, M. et al. "Identification of Diverse Database Subsets using Property-Based and Fragment-Based Molecular Descriptions." *Quantitative Structure-Activity Relationships* **21**:598-604 (2002).

```
>>> def distij(i,j,fps=fps):
... return 1-DataStructs.DiceSimilarity(fps[i],fps[j])
```

Now create a picker and grab a set of 10 diverse molecules:

```
>>> picker = MaxMinPicker()
>>> pickIndices = picker.LazyPick(distij,nfps,10,seed=23)
>>> list(pickIndices)
[93, 109, 154, 6, 95, 135, 151, 61, 137, 139]
```

Note that the picker just returns indices of the fingerprints; we can get the molecules themselves as follows:

```
>>> picks = [ms[x] for x in pickIndices]
```

3.7.6 Generating Similarity Maps Using Fingerprints

Similarity maps are a way to visualize the atomic contributions to the similarity between a molecule and a reference molecule. The methodology is described in Ref. 13 . They are in the rdkit.Chem.Draw.SimilarityMaps module:

Start by creating two molecules:

```
>>> from rdkit import Chem
>>> mol = Chem.MolFromSmiles('COclccc2cc(C(=0)NCCCCN3CCN(c4ccc5nccnc54)CC3)oc21')
>>> refmol = Chem.MolFromSmiles('CCCN(CCCCN1CCN(c2cccc2CC)CC1)Cc1ccc2cccc2c1')
```

The SimilarityMaps module supports three kind of fingerprints: atom pairs, topological torsions and Morgan fingerprints.

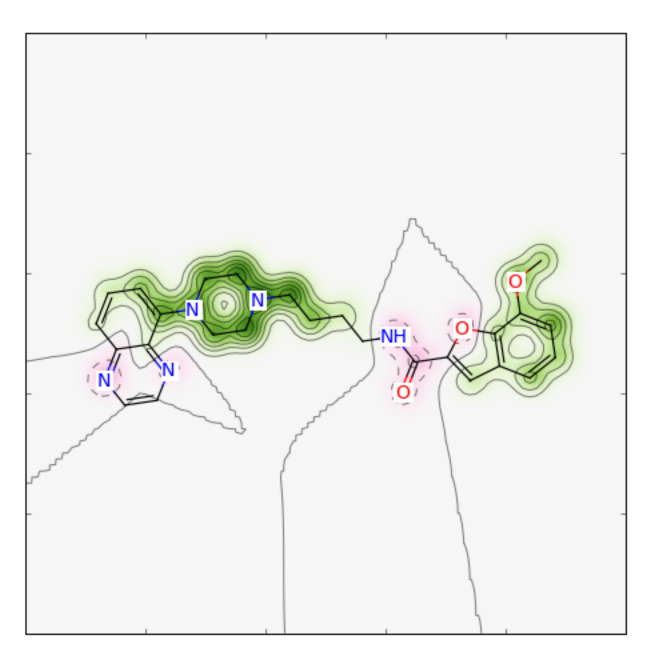
```
>>> from rdkit.Chem import Draw
>>> from rdkit.Chem.Draw import SimilarityMaps
>>> fp = SimilarityMaps.GetAPFingerprint(mol, fpType='normal')
>>> fp = SimilarityMaps.GetTTFingerprint(mol, fpType='normal')
>>> fp = SimilarityMaps.GetMorganFingerprint(mol, fpType='bv')
```

The types of atom pairs and torsions are normal (default), hashed and bit vector (bv). The types of the Morgan fingerprint are bit vector (bv, default) and count vector (count).

The function generating a similarity map for two fingerprints requires the specification of the fingerprint function and optionally the similarity metric. The default for the latter is the Dice similarity. Using all the default arguments of the Morgan fingerprint function, the similarity map can be generated like this:

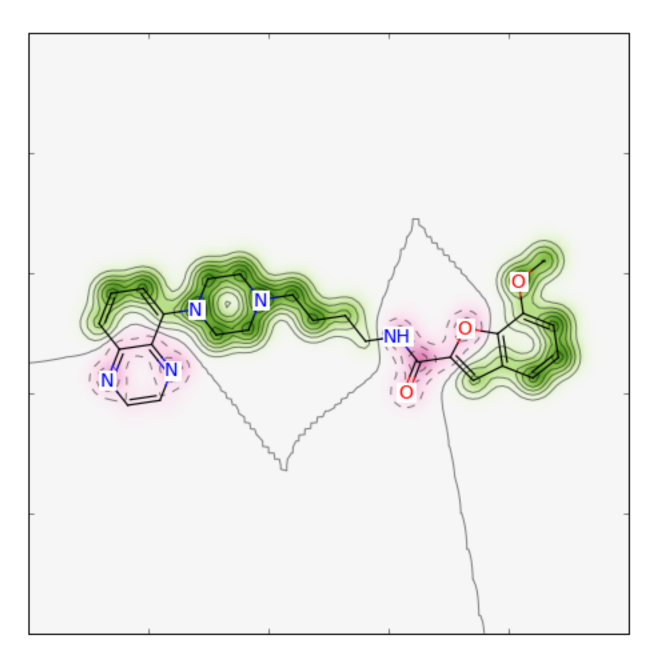
```
>>> fig, maxweight = SimilarityMaps.GetSimilarityMapForFingerprint(refmol, mol, SimilarityMaps.GetMo:
```

¹³ Riniker, S.; Landrum, G. A. "Similarity Maps - A Visualization Strategy for Molecular Fingerprints and Machine-Learning Methods" *J. Cheminf.* **5**:43 (2013).



For a different type of Morgan (e.g. count) and radius = 1 instead of 2, as well as a different similarity metric (e.g. Tanimoto), the call becomes:

```
>>> from rdkit import DataStructs
>>> fig, maxweight = SimilarityMaps.GetSimilarityMapForFingerprint(refmol, mol, lambda m,idx: SimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimilarityMaps.GetSimil
```

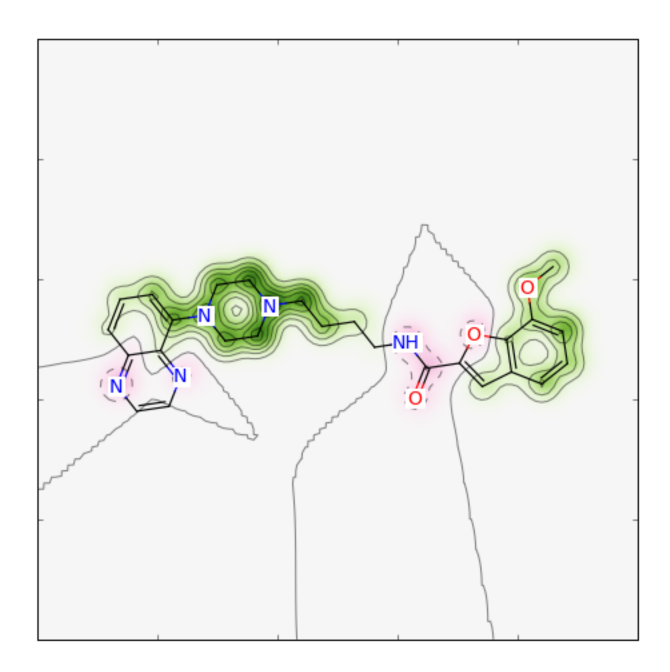


The convenience function GetSimilarityMapForFingerprint involves the normalisation of the atomic weights such that the maximum absolute weight is 1. Therefore, the function outputs the maximum weight that was found when creating the map.

```
>>> print maxweight 0.0574712643678
```

If one does not want the normalisation step, the map can be created like:

```
>>> weights = SimilarityMaps.GetAtomicWeightsForFingerprint(refmol, mol, SimilarityMaps.GetMorganFing
>>> print ["%.2f" % w for w in weights]
['0.05', ...
>>> fig = SimilarityMaps.GetSimilarityMapFromWeights(mol, weights)
```



3.8 Descriptor Calculation

A variety of descriptors are available within the RDKit. The complete list is provided in List of Available Descriptors. Most of the descriptors are straightforward to use from Python via the centralized rdkit.Chem.Descriptors module:

```
>>> from rdkit.Chem import Descriptors
>>> m = Chem.MolFromSmiles('clcccc1C(=0)0')
>>> Descriptors.TPSA(m)
37.3
>>> Descriptors.MolLogP(m)
1.3848
```

Partial charges are handled a bit differently:

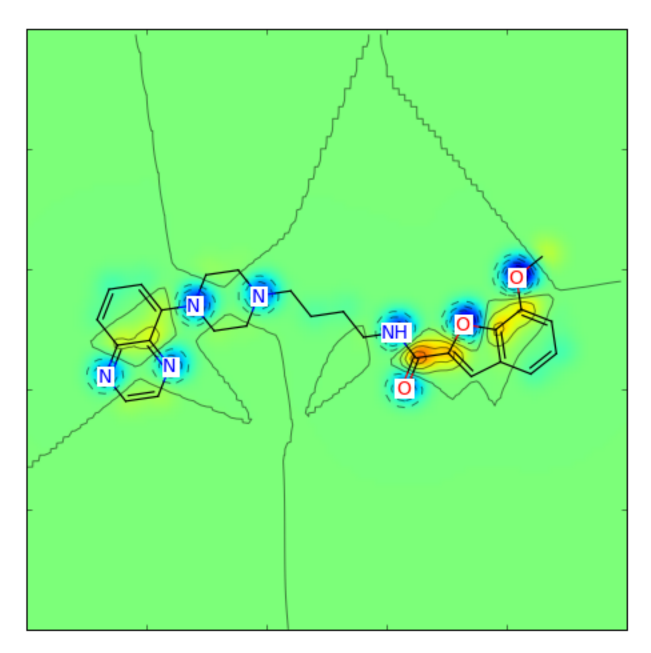
```
>>> m = Chem.MolFromSmiles('clcccclC(=0)0')
>>> AllChem.ComputeGasteigerCharges(m)
>>> float(m.GetAtomWithIdx(0).GetProp('_GasteigerCharge'))
-0.047...
```

3.8.1 Visualization of Descriptors

Similarity maps can be used to visualize descriptors that can be divided into atomic contributions.

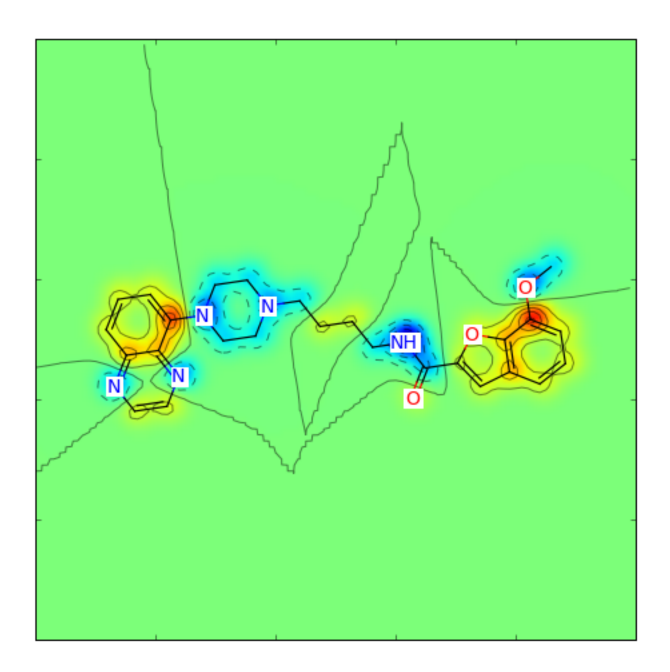
The Gasteiger partial charges can be visualized as (using a different color scheme):

```
>>> from rdkit.Chem.Draw import SimilarityMaps
>>> mol = Chem.MolFromSmiles('COclccc2cc(C(=0)NCCCCN3CCN(c4ccc5nccnc54)CC3)oc21')
>>> AllChem.ComputeGasteigerCharges(mol)
>>> contribs = [float(mol.GetAtomWithIdx(i).GetProp('_GasteigerCharge')) for i in range(mol.GetNumAtc)
>>> fig = SimilarityMaps.GetSimilarityMapFromWeights(mol, contribs, colorMap='jet', contourLines=10)
```



Or for the Crippen contributions to logP:

```
>>> from rdkit.Chem import rdMolDescriptors
>>> contribs = rdMolDescriptors._CalcCrippenContribs(mol)
>>> fig = SimilarityMaps.GetSimilarityMapFromWeights(mol,[x for x,y in contribs], colorMap='jet', contribs]
```



3.9 Chemical Reactions

The RDKit also supports applying chemical reactions to sets of molecules. One way of constructing chemical reactions is to use a SMARTS-based language similar to Daylight's Reaction SMILES ¹⁴:

```
>>> rxn = AllChem.ReactionFromSmarts('[C:1](=[0:2])-[OD1].[N!H0:3]>>[C:1](=[0:2])[N:3]')
>>> rxn
<rdkit.Chem.rdChemReactions.ChemicalReaction object at 0x...>
>>> rxn.GetNumProductTemplates()
1
>>> ps = rxn.RunReactants((Chem.MolFromSmiles('CC(=0)0'),Chem.MolFromSmiles('NC')))
>>> len(ps) # one entry for each possible set of products
```

¹⁴ A more detailed description of reaction smarts, as defined by the rdkit, is in the *The RDKit Book*.

```
1
>>> len(ps[0]) # each entry contains one molecule for each product
1
>>> Chem.MolToSmiles(ps[0][0])
'CNC(C)=0'
>>> ps = rxn.RunReactants((Chem.MolFromSmiles('C(COC(=0)0)C(=0)0'), Chem.MolFromSmiles('NC')))
>>> len(ps)
2
>>> Chem.MolToSmiles(ps[0][0])
'CNC(=0)OCCC(=0)0'
>>> Chem.MolToSmiles(ps[1][0])
'CNC(=0)CCCC(=0)0'
Reactions can also be built from MDL rxn files:
>>> rxn = AllChem.ReactionFromRxnFile('data/AmideBond.rxn')
>>> rxn.GetNumReactantTemplates()
```

```
>>> rxn = AllChem.ReactionFromRxnFile('data/AmideBond.rxn')
>>> rxn.GetNumReactantTemplates()
2
>>> rxn.GetNumProductTemplates()
1
>>> ps = rxn.RunReactants((Chem.MolFromSmiles('CC(=0)O'), Chem.MolFromSmiles('NC')))
>>> len(ps)
1
>>> Chem.MolToSmiles(ps[0][0])
'CNC(C) = O'
```

It is, of course, possible to do reactions more complex than amide bond formation:

```
>>> rxn = AllChem.ReactionFromSmarts('[C:1]=[C:2].[C:3]=[*:4][*:5]=[C:6]>>[C:1]1[C:2][C:3][*:4]=[*:5
>>> ps = rxn.RunReactants((Chem.MolFromSmiles('OC=C'), Chem.MolFromSmiles('C=CC(N)=C')))
>>> Chem.MolToSmiles(ps[0][0])
'NC1=CCCC(0)C1'
```

Note in this case that there are multiple mappings of the reactants onto the templates, so we have multiple product sets:

```
>>> len(ps)
4
```

You can use canonical smiles and a python dictionary to get the unique products:

Note that the molecules that are produced by the chemical reaction processing code are not sanitized, as this artificial reaction demonstrates:

```
>>> rxn = AllChem.ReactionFromSmarts('[C:1]=[C:2][C:3]=[C:4].[C:5]=[C:6]>>[C:1]1=[C:2][C:3]=[C:4][C:5]
>>> ps = rxn.RunReactants((Chem.MolFromSmiles('C=CC=C'), Chem.MolFromSmiles('C=C')))
>>> Chem.MolToSmiles(ps[0][0])
'C1=CC=CC=C1'
>>> p0 = ps[0][0]
>>> Chem.SanitizeMol(p0)
rdkit.Chem.rdmolops.SanitizeFlags.SANITIZE_NONE
```

```
>>> Chem.MolToSmiles(p0)
'clccccl'
```

3.9.1 Advanced Reaction Functionality

Protecting Atoms

Sometimes, particularly when working with rxn files, it is difficult to express a reaction exactly enough to not end up with extraneous products. The RDKit provides a method of "protecting" atoms to disallow them from taking part in reactions.

This can be demonstrated re-using the amide-bond formation reaction used above. The query for amines isn't specific enough, so it matches any nitrogen that has at least one H attached. So if we apply the reaction to a molecule that already has an amide bond, the amide N is also treated as a reaction site:

```
>>> rxn = AllChem.ReactionFromRxnFile('data/AmideBond.rxn')
>>> acid = Chem.MolFromSmiles('CC(=0)O')
>>> base = Chem.MolFromSmiles('CC(=0)NCCN')
>>> ps = rxn.RunReactants((acid,base))
>>> len(ps)
2
>>> Chem.MolToSmiles(ps[0][0])
'CC(=0)N(CCN)C(C)=O'
>>> Chem.MolToSmiles(ps[1][0])
'CC(=0)NCCNC(C)=O'
```

The first product corresponds to the reaction at the amide N.

We can prevent this from happening by protecting all amide Ns. Here we do it with a substructure query that matches amides and thioamides and then set the "protected" property on matching atoms:

```
>>> amidep = Chem.MolFromSmarts('[N;$(NC=[0,S])]')
>>> for match in base.GetSubstructMatches(amidep):
... base.GetAtomWithIdx(match[0]).SetProp('_protected','1')
```

Now the reaction only generates a single product:

```
>>> ps = rxn.RunReactants((acid,base))
>>> len(ps)
1
>>> Chem.MolToSmiles(ps[0][0])
'CC(=0)NCCNC(C)=0'
```

3.9.2 Recap Implementation

Associated with the chemical reaction functionality is an implementation of the Recap algorithm. ¹⁵ Recap uses a set of chemical transformations mimicking common reactions carried out in the lab in order to decompose a molecule into a series of reasonable fragments.

The RDKit rdkit.Chem.Recap implementation keeps track of the hierarchy of transformations that were applied:

```
>>> from rdkit import Chem
>>> from rdkit.Chem import Recap
>>> m = Chem.MolFromSmiles('clcccc1oCCoC(=0)CC')
```

¹⁵ Lewell, X.Q.; Judd, D.B.; Watson, S.P.; Hann, M.M. "RECAP-Retrosynthetic Combinatorial Analysis Procedure: A Powerful New Technique for Identifying Privileged Molecular Fragments with Useful Applications in Combinatorial Chemistry" *J. Chem. Inf. Comp. Sci.* 38:511-22 (1998).

```
>>> hierarch = Recap.RecapDecompose(m)
>>> type(hierarch)
<class 'rdkit.Chem.Recap.RecapHierarchyNode'>
```

The hierarchy is rooted at the original molecule:

```
>>> hierarch.smiles
'CCC(=0)OCCOclcccc1'
```

and each node tracks its children using a dictionary keyed by SMILES:

```
>>> ks=hierarch.children.keys()
>>> ks.sort()
>>> ks
['[*]C(=0)CC', '[*]CCOC(=0)CC', '[*]CCOclcccccl', '[*]OCCOclcccccl', '[*]clcccccl']
```

The nodes at the bottom of the hierarchy (the leaf nodes) are easily accessible, also as a dictionary keyed by SMILES:

```
>>> ks=hierarch.GetLeaves().keys()
>>> ks.sort()
>>> ks
['[*]C(=0)CC', '[*]CCO[*]', '[*]CCOclcccccl', '[*]clcccccl']
```

Notice that dummy atoms are used to mark points where the molecule was fragmented.

The nodes themselves have associated molecules:

```
>>> leaf = hierarch.GetLeaves() [ks[0]]
>>> Chem.MolToSmiles(leaf.mol)
'[*]C(=0)CC'
```

3.9.3 BRICS Implementation

The RDKit also provides an implementation of the BRICS algorithm. ¹⁶ BRICS provides another method for fragmenting molecules along synthetically accessible bonds:

```
>>> from rdkit.Chem import BRICS
>>> cdk2mols = Chem.SDMolSupplier('data/cdk2.sdf')
>>> m1 = cdk2mols[0]
>>> list(BRICS.BRICSDecompose(m1))
['[4*]CC(=0)C(C)C', '[14*]clnc(N)nc2[nH]cnc12', '[3*]0[3*]']
>>> m2 = cdk2mols[20]
>>> list(BRICS.BRICSDecompose(m2))
['[3*]OC', '[1*]C(=0)NN(C)C', '[14*]c1[nH]nc2c1C(=0)c1c([16*])cccc1-2', '[5*]N[5*]', '[16*]c1ccc([16*])cccc1-2', '[5*]N[5*]', '[5*
```

Notice that RDKit BRICS implementation returns the unique fragments generated from a molecule and that the dummy atoms are tagged to indicate which type of reaction applies.

It's quite easy to generate the list of all fragments for a group of molecules:

```
>>> allfrags=set()
>>> for m in cdk2mols:
...    pieces = BRICS.BRICSDecompose(m)
...    allfrags.update(pieces)
>>> len(allfrags)
90
```

¹⁶ Degen, J.; Wegscheid-Gerlach, C.; Zaliani, A; Rarey, M. "On the Art of Compiling and Using 'Drug-Like' Chemical Fragment Spaces." *ChemMedChem* 3:1503–7 (2008).

```
>>> list(allfrags)[:5]
['[4*]CC[NH3+]', '[14*]clcnc[nH]1', '[16*]clccc([16*])c(Cl)c1', '[15*]ClCCCC1', '[7*]ClC(=0)Nc2ccc(S
```

The BRICS module also provides an option to apply the BRICS rules to a set of fragments to create new molecules:

```
>>> import random
>>> random.seed(127)
>>> fragms = [Chem.MolFromSmiles(x) for x in allfrags]
>>> ms = BRICS.BRICSBuild(fragms)
```

The result is a generator object:

```
>>> ms
<generator object BRICSBuild at 0x...>
```

That returns molecules on request:

```
>>> prods = [ms.next() for x in range(10)]
>>> prods[0]
<rdkit.Chem.rdchem.Mol object at 0x...>
```

The molecules have not been sanitized, so it's a good idea to at least update the valences before continuing:

```
>>> for prod in prods:
... prod.UpdatePropertyCache(strict=False)
...
>>> Chem.MolToSmiles(prods[0],True)
'O=[N+]([0-])clccc(C2CCC02)cc1'
>>> Chem.MolToSmiles(prods[1],True)
'clccc(C2CCC02)cc1'
>>> Chem.MolToSmiles(prods[2],True)
'NS(=0)(=0)clccc(C2CCC02)cc1'
```

3.9.4 Other fragmentation approaches

In addition to the methods described above, the RDKit provide a very flexible generic function for fragmenting molecules along user-specified bonds.

Here's a quick demonstration of using that to break all bonds between atoms in rings and atoms not in rings. We start by finding all the atom pairs:

```
>>> m = Chem.MolFromSmiles('CC1CC(0)C1CCC1CC1')
>>> bis = m.GetSubstructMatches(Chem.MolFromSmarts('[!R][R]'))
>>> bis
((0, 1), (4, 3), (6, 5), (7, 8))
```

then we get the corresponding bond indices:

```
>>> bs = [m.GetBondBetweenAtoms(x,y).GetIdx() for x,y in bis]
>>> bs
[0, 3, 5, 7]
```

then we use those bond indices as input to the fragmentation function:

```
>>> nm = Chem.FragmentOnBonds(m,bs)
```

the output is a molecule that has dummy atoms marking the places where bonds were broken:

```
>>> Chem.MolToSmiles(nm,True)
'[*]C1CC([4*])C1[6*].[1*]C.[3*]O.[5*]CC[8*].[7*]C1CC1'
```

By default the attachment points are labelled (using isotopes) with the index of the atom that was removed. We can also provide our own set of atom labels in the form of pairs of unsigned integers. The first value in each pair is used as the label for the dummy that replaces the bond's begin atom, the second value in each pair is for the dummy that replaces the bond's end atom. Here's an example, repeating the analysis above and marking the positions where the non-ring atoms were with the label 10 and marking the positions where the ring atoms were with label 1:

```
>>> bis = m.GetSubstructMatches(Chem.MolFromSmarts('[!R][R]'))
>>> bs = []
>>> labels=[]
>>> for bi in bis:
       b = m.GetBondBetweenAtoms(bi[0],bi[1])
       if b.GetBeginAtomIdx() == bi[0]:
           labels.append((10,1))
. . .
       else:
. . .
           labels.append((1,10))
. . .
       bs.append(b.GetIdx())
>>> nm = Chem.FragmentOnBonds(m,bs,dummyLabels=labels)
>>> Chem.MolToSmiles(nm,True)
'[1*]C.[1*]CC[1*].[1*]O.[10*]C1CC([10*])C1[10*].[10*]C1CC1'
```

3.10 Chemical Features and Pharmacophores

3.10.1 Chemical Features

Chemical features in the RDKit are defined using a SMARTS-based feature definition language (described in detail in the RDKit book). To identify chemical features in molecules, you first must build a feature factory:

```
>>> from rdkit import Chem
>>> from rdkit.Chem import ChemicalFeatures
>>> from rdkit import RDConfig
>>> import os
>>> fdefName = os.path.join(RDConfig.RDDataDir,'BaseFeatures.fdef')
>>> factory = ChemicalFeatures.BuildFeatureFactory(fdefName)
and then use the factory to search for features:
```

>>> m = Chem.MolFromSmiles('OCc1cccc1CN')
>>> feats = factory.GetFeaturesForMol(m)
>>> len(feats)
g

The individual features carry information about their family (e.g. donor, acceptor, etc.), type (a more detailed description), and the atom(s) that is/are associated with the feature:

```
>>> feats[0].GetFamily()
'Donor'
>>> feats[0].GetType()
'SingleAtomDonor'
>>> feats[0].GetAtomIds()
(0,)
>>> feats[4].GetFamily()
'Aromatic'
```

```
>>> feats[4].GetAtomIds()
(2, 3, 4, 5, 6, 7)
```

If the molecule has coordinates, then the features will also have reasonable locations:

```
>>> from rdkit.Chem import AllChem
>>> AllChem.Compute2DCoords(m)
0
>>> feats[0].GetPos()
<rdkit.Geometry.rdGeometry.Point3D object at 0x...>
>>> list(feats[0].GetPos())
[2.07..., -2.335..., 0.0]
```

3.10.2 2D Pharmacophore Fingerprints

Combining a set of chemical features with the 2D (topological) distances between them gives a 2D pharmacophore. When the distances are binned, unique integer ids can be assigned to each of these pharmacophores and they can be stored in a fingerprint. Details of the encoding are in the *The RDKit Book*.

Generating pharmacophore fingerprints requires chemical features generated via the usual RDKit feature-typing mechanism:

```
>>> from rdkit import Chem
>>> from rdkit.Chem import ChemicalFeatures
>>> fdefName = 'data/MinimalFeatures.fdef'
>>> featFactory = ChemicalFeatures.BuildFeatureFactory(fdefName)
```

The fingerprints themselves are calculated using a signature (fingerprint) factory, which keeps track of all the parameters required to generate the pharmacophore:

```
>>> from rdkit.Chem.Pharm2D.SigFactory import SigFactory
>>> sigFactory = SigFactory(featFactory,minPointCount=2,maxPointCount=3)
>>> sigFactory.SetBins([(0,2),(2,5),(5,8)])
>>> sigFactory.Init()
>>> sigFactory.GetSigSize()
885
```

The signature factory is now ready to be used to generate fingerprints, a task which is done using the rdkit.Chem.Pharm2D.Generate module:

```
>>> from rdkit.Chem.Pharm2D import Generate
>>> mol = Chem.MolFromSmiles('OCC(=O)CCCN')
>>> fp = Generate.Gen2DFingerprint(mol, sigFactory)
>>> fp
<rdkit.DataStructs.cDataStructs.SparseBitVect object at 0x...>
>>> len(fp)
885
>>> fp.GetNumOnBits()
57
```

Details about the bits themselves, including the features that are involved and the binned distance matrix between the features, can be obtained from the signature factory:

```
>>> list(fp.GetOnBits())[:5]
[1, 2, 6, 7, 8]
>>> sigFactory.GetBitDescription(1)
'Acceptor Acceptor |0 1|1 0|'
>>> sigFactory.GetBitDescription(2)
```

```
'Acceptor Acceptor |0 2|2 0|'
>>> sigFactory.GetBitDescription(8)
'Acceptor Donor |0 2|2 0|'
>>> list(fp.GetOnBits())[-5:]
[704, 706, 707, 708, 714]
>>> sigFactory.GetBitDescription(707)
'Donor Donor PosIonizable |0 1 2|1 0 1|2 1 0|'
>>> sigFactory.GetBitDescription(714)
'Donor Donor PosIonizable |0 2 2|2 0 0|2 0 0|'
```

For the sake of convenience (to save you from having to edit the fdef file every time) it is possible to disable particular feature types within the SigFactory:

```
>>> sigFactory.skipFeats=['PosIonizable']
>>> sigFactory.Init()
>>> sigFactory.GetSigSize()
510
>>> fp2 = Generate.Gen2DFingerprint(mol,sigFactory)
>>> fp2.GetNumOnBits()
36
```

Another possible set of feature definitions for 2D pharmacophore fingerprints in the RDKit are those published by Gobbi and Poppinger. ¹⁷ The module rdkit.Chem.Pharm2D.Gobbi_Pharm2D has a pre-configured signature factory for these fingerprint types. Here's an example of using it:

```
>>> from rdkit import Chem
>>> from rdkit.Chem.Pharm2D import Gobbi_Pharm2D,Generate
>>> m = Chem.MolFromSmiles('OCC=CC(=O)O')
>>> fp = Generate.Gen2DFingerprint(m,Gobbi_Pharm2D.factory)
>>> fp
<rdkit.DataStructs.cDataStructs.SparseBitVect object at 0x...>
>>> fp.GetNumOnBits()
8
>>> list(fp.GetOnBits())
[23, 30, 150, 154, 157, 185, 28878, 30184]
>>> Gobbi_Pharm2D.factory.GetBitDescription(157)
'HA HD |0 3|3 0|'
>>> Gobbi_Pharm2D.factory.GetBitDescription(30184)
'HA HD HD |0 3 0|3 0 3|0 3 0|'
```

3.11 Molecular Fragments

The RDKit contains a collection of tools for fragmenting molecules and working with those fragments. Fragments are defined to be made up of a set of connected atoms that may have associated functional groups. This is more easily demonstrated than explained:

```
>>> fName=os.path.join(RDConfig.RDDataDir,'FunctionalGroups.txt')
>>> from rdkit.Chem import FragmentCatalog
>>> fparams = FragmentCatalog.FragCatParams(1,6,fName)
>>> fparams.GetNumFuncGroups()
39
>>> fcat=FragmentCatalog.FragCatalog(fparams)
>>> fcgen=FragmentCatalog.FragCatGenerator()
>>> m = Chem.MolFromSmiles('OCC=CC(=0)O')
```

¹⁷ Gobbi, A. & Poppinger, D. "Genetic optimization of combinatorial libraries." *Biotechnology and Bioengineering* **61**:47-54 (1998).

```
>>> fcgen.AddFragsFromMol(m, fcat)
3
>>> fcat.GetEntryDescription(0)
'C<-O>C'
>>> fcat.GetEntryDescription(1)
'C=C<-C(=0)O>'
>>> fcat.GetEntryDescription(2)
'C<-C(=0)O>=CC<-O>'
```

The fragments are stored as entries in a rdkit.Chem.rdfragcatalog.FragCatalog. Notice that the entry descriptions include pieces in angular brackets (e.g. between '<' and '>'). These describe the functional groups attached to the fragment. For example, in the above example, the catalog entry 0 corresponds to an ethyl fragment with an alcohol attached to one of the carbons and entry 1 is an ethylene with a carboxylic acid on one carbon. Detailed information about the functional groups can be obtained by asking the fragment for the ids of the functional groups it contains and then looking those ids up in the rdkit.Chem.rdfragcatalog.FragCatParams object:

```
>>> list(fcat.GetEntryFuncGroupIds(2))
[34, 1]
>>> fparams.GetFuncGroup(1)
<rdkit.Chem.rdchem.Mol object at 0x...>
>>> Chem.MolToSmarts(fparams.GetFuncGroup(1))
'*-C(=0)-,:[0&D1]'
>>> Chem.MolToSmarts(fparams.GetFuncGroup(34))
'*-[0&D1]'
>>> fparams.GetFuncGroup(1).GetProp('_Name')
'-C(=0)0'
>>> fparams.GetFuncGroup(34).GetProp('_Name')
'-O'
```

The catalog is hierarchical: smaller fragments are combined to form larger ones. From a small fragment, one can find the larger fragments to which it contributes using the rdkit.Chem.rdfragcatalog.FragCatalog.GetEntryDownIds method:

```
>>> fcat=FragmentCatalog.FragCatalog(fparams)
>>> m = Chem.MolFromSmiles('OCC(NC1CC1)CCC')
>>> fcgen.AddFragsFromMol(m, fcat)
15
>>> fcat.GetEntryDescription(0)
'C<-0>C'
>>> fcat.GetEntryDescription(1)
'CN<-cPropyl>'
>>> list(fcat.GetEntryDownIds(0))
[3, 4]
>>> fcat.GetEntryDescription(3)
'C<-0>CC'
>>> fcat.GetEntryDescription(4)
'C<-0>CN<-cPropyl>'
```

The fragments from multiple molecules can be added to a catalog:

```
>>> suppl = Chem.SmilesMolSupplier('data/bzr.smi')
>>> ms = [x for x in suppl]
>>> fcat=FragmentCatalog.FragCatalog(fparams)
>>> for m in ms: nAdded=fcgen.AddFragsFromMol(m, fcat)
>>> fcat.GetNumEntries()
1169
>>> fcat.GetEntryDescription(0)
'Cc'
>>> fcat.GetEntryDescription(100)
```

```
'cc-nc(C)n'
```

The fragments in a catalog are unique, so adding a molecule a second time doesn't add any new entries:

```
>>> fcgen.AddFragsFromMol(ms[0],fcat)
0
>>> fcat.GetNumEntries()
1169
```

Once a rdkit.Chem.rdfragcatalog.FragCatalog has been generated, it can be used to fingerprint molecules:

```
>>> fpgen = FragmentCatalog.FragFPGenerator()
>>> fp = fpgen.GetFPForMol(ms[8],fcat)
>>> fp
<rdkit.DataStructs.cDataStructs.ExplicitBitVect object at 0x...>
>>> fp.GetNumOnBits()
189
```

The rest of the machinery associated with fingerprints can now be applied to these fragment fingerprints. For example, it's easy to find the fragments that two molecules have in common by taking the intersection of their fingerprints:

```
>>> fp2 = fpgen.GetFPForMol(ms[7],fcat)
>>> andfp = fp&fp2
>>> obl = list(andfp.GetOnBits())
>>> fcat.GetEntryDescription(obl[-1])
'ccc(cc)NC<=0>'
>>> fcat.GetEntryDescription(obl[-5])
'c<-X>ccc(N)cc'
```

or we can find the fragments that distinguish one molecule from another:

```
>>> combinedFp=fp&(fp^fp2) # can be more efficent than fp&(!fp2)
>>> obl = list(combinedFp.GetOnBits())
>>> fcat.GetEntryDescription(obl[-1])
'cccc(N)cc'
```

Or we can use the bit ranking functionality from the rdkit.ML.InfoTheory.rdInfoTheory.InfoBitRanker class to identify fragments that distinguish actives from inactives:

```
>>> suppl = Chem.SDMolSupplier('data/bzr.sdf')
>>> sdms = [x for x in suppl]
>>> fps = [fpgen.GetFPForMol(x,fcat) for x in sdms]
>>> from rdkit.ML.InfoTheory import InfoBitRanker
>>> ranker = InfoBitRanker(len(fps[0]),2)
>>> acts = [float(x.GetProp('ACTIVITY')) for x in sdms]
>>> for i, fp in enumerate(fps):
... act = int(acts[i] > 7)
     ranker.AccumulateVotes(fp,act)
>>> top5 = ranker.GetTopN(5)
>>> for id, gain, n0, n1 in top5:
. . .
     print int(id),'%.3f'%gain,int(n0),int(n1)
702 0.081 20 17
328 0.073 23 25
341 0.073 30 43
173 0.073 30 43
1034 0.069 5 53
```

The columns above are: bitId, infoGain, nInactive, nActive. Note that this approach isn't particularly effective for this artificial example.

3.12 Non-Chemical Functionality

3.12.1 Bit vectors

Bit vectors are containers for efficiently storing a set number of binary values, e.g. for fingerprints. The RDKit includes two types of fingerprints differing in how they store the values internally; the two types are easily interconverted but are best used for different purpose:

- SparseBitVects store only the list of bits set in the vector; they are well suited for storing very large, very sparsely occupied vectors like pharmacophore fingerprints. Some operations, such as retrieving the list of on bits, are quite fast. Others, such as negating the vector, are very, very slow.
- ExplicitBitVects keep track of both on and off bits. They are generally faster than SparseBitVects, but require more memory to store.

3.12.2 Discrete value vectors

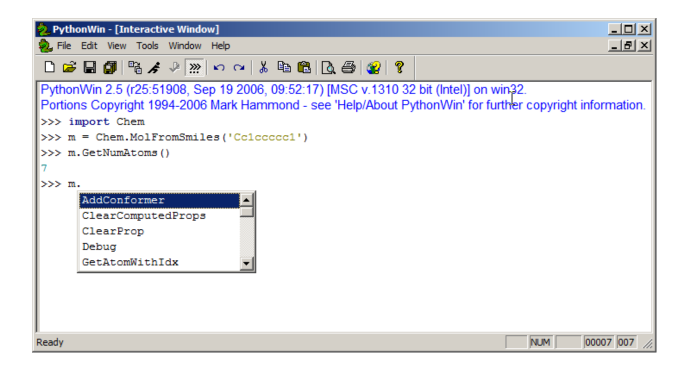
3.12.3 3D grids

3.12.4 Points

3.13 Getting Help

There is a reasonable amount of documentation available within from the RDKit's docstrings. These are accessible using Python's help command:

When working in an environment that does command completion or tooltips, one can see the available methods quite easily. Here's a sample screenshot from within Mark Hammond's PythonWin environment:



3.14 Advanced Topics/Warnings

3.14.1 Editing Molecules

Some of the functionality provided allows molecules to be edited "in place":

```
>>> m = Chem.MolFromSmiles('clcccc1')
>>> m.GetAtomWithIdx(0).SetAtomicNum(7)
>>> Chem.SanitizeMol(m)
rdkit.Chem.rdmolops.SanitizeFlags.SANITIZE_NONE
>>> Chem.MolToSmiles(m)
'clccncc1'
```

Do not forget the sanitization step, without it one can end up with results that look ok (so long as you don't think):

```
>>> m = Chem.MolFromSmiles('clcccc1')
>>> m.GetAtomWithIdx(0).SetAtomicNum(8)
>>> Chem.MolToSmiles(m)
'clcccc1'
```

but that are, of course, complete nonsense, as sanitization will indicate:

```
>>> Chem.SanitizeMol(m)
Traceback (most recent call last):
   File "/usr/lib/python2.6/doctest.py", line 1253, in __run
      compileflags, 1) in test.globs
   File "<doctest default[0]>", line 1, in <module>
            Chem.SanitizeMol(m)
ValueError: Sanitization error: Can't kekulize mol
```

More complex transformations can be carried out using the rdkit.Chem.rdchem.RWMol class:

```
>>> m = Chem.MolFromSmiles('CC(=0)C=CC=C')
>>> mw = Chem.RWMol(m)
>>> mw.ReplaceAtom(4,Chem.Atom(7))
>>> mw.AddAtom(Chem.Atom(6))
7
>>> mw.AddAtom(Chem.Atom(6))
8
>>> mw.AddBond(6,7,Chem.BondType.SINGLE)
7
>>> mw.AddBond(7,8,Chem.BondType.DOUBLE)
8
>>> mw.AddBond(8,3,Chem.BondType.SINGLE)
9
>>> mw.RemoveAtom(0)
>>> mw.RemoveAtom(0)
>>> mw.GetNumAtoms()
```

The RWMol can be used just like an ROMol:

```
>>> Chem.MolToSmiles(mw)
'O=CC1C=CC=CN=1'
>>> Chem.SanitizeMol(mw)
rdkit.Chem.rdmolops.SanitizeFlags.SANITIZE_NONE
>>> Chem.MolToSmiles(mw)
'O=Cc1ccccn1'
```

It is even easier to generate nonsense using the RWMol than it is with standard molecules. If you need chemically reasonable results, be certain to sanitize the results.

3.15 Miscellaneous Tips and Hints

3.15.1 Chem vs AllChem

The majority of "basic" chemical functionality (e.g. reading/writing molecules, substructure searching, molecular cleanup, etc.) is in the rdkit.Chem module. More advanced, or less frequently used, functionality is in rdkit.Chem.AllChem. The distinction has been made to speed startup and lower import times; there's no sense in loading the 2D->3D library and force field implementation if one is only interested in reading and writing a couple of molecules. If you find the Chem/AllChem thing annoying or confusing, you can use python's "import ... as ..." syntax to remove the irritation:

```
>>> from rdkit.Chem import AllChem as Chem
>>> m = Chem.MolFromSmiles('CCC')
```

3.15.2 The SSSR Problem

As others have ranted about with more energy and eloquence than I intend to, the definition of a molecule's smallest set of smallest rings is not unique. In some high symmetry molecules, a "true" SSSR will give results that are unappealing. For example, the SSSR for cubane only contains 5 rings, even though there are "obviously" 6. This problem can be fixed by implementing a *small* (instead of *smallest*) set of smallest rings algorithm that returns symmetric results. This is the approach that we took with the RDKit.

Because it is sometimes useful to be able to count how many SSSR rings are present in the molecule, there is a rdkit.Chem.rdmolops.GetSSSR function, but this only returns the SSSR count, not the potentially non-unique set of rings.

3.16 List of Available Descriptors

| Descriptor/Descriptor Family | Notes |
|---|--|
| Gasteiger/Marsili Partial Charges | Tetrahedron 36 :3219-28 (1980) |
| BalabanJ | Chem. Phys. Lett. 89 :399-404 (1982) |
| BertzCT | J. Am. Chem. Soc. 103:3599-601 (1981) |
| Ipc | J. Chem. Phys. 67 :4517-33 (1977) |
| HallKierAlpha | Rev. Comput. Chem. 2:367-422 (1991) |
| Kappa1 - Kappa3 | Rev. Comput. Chem. 2:367-422 (1991) |
| Chi0, Chi1 | Rev. Comput. Chem. 2:367-422 (1991) |
| Chi0n - Chi4n | Rev. Comput. Chem. 2:367-422 (1991) |
| Chi0v - Chi4v | Rev. Comput. Chem. 2:367-422 (1991) |
| MolLogP | Wildman and Crippen <i>JCICS</i> 39 :868-73 (1999) |
| MolMR | Wildman and Crippen <i>JCICS</i> 39 :868-73 (1999) |
| MolWt | |
| ExactMolWt | |
| HeavyAtomCount | |
| HeavyAtomMolWt | |
| NHOHCount | |
| NOCount | |
| NumHAcceptors | |
| NumHDonors | |
| NumHeteroatoms | |
| NumRotatableBonds | |
| NumValenceElectrons | |
| NumAmideBonds | |
| Num{Aromatic,Saturated,Aliphatic}Rings | |
| Num{Aromatic,Saturated,Aliphatic}{Hetero,Carbo}cycles | |
| RingCount | |
| FractionCSP3 | |
| TPSA | J. Med. Chem. 43:3714-7, (2000) |
| LabuteASA | J. Mol. Graph. Mod. 18:464-77 (2000) |
| PEOE_VSA1 - PEOE_VSA14 | MOE-type descriptors using partial charges and surface area contribution |
| SMR_VSA1 - SMR_VSA10 | MOE-type descriptors using MR contributions and surface area contribu |
| SlogP_VSA1 - SlogP_VSA12 | MOE-type descriptors using LogP contributions and surface area contributions |
| EState_VSA1 - EState_VSA11 | MOE-type descriptors using EState indices and surface area contribution |
| VSA_EState1 - VSA_EState10 | MOE-type descriptors using EState indices and surface area contribution |
| MQNs | Nguyen et al. <i>ChemMedChem</i> 4 :1803-5 (2009) |
| Topliss fragments | implemented using a set of SMARTS definitions in \$(RDBASE)/Data/F |

3.17 List of Available Fingerprints

| Fingerprint Type | Notes |
|------------------|--|
| RDKit | a Daylight-like fingerprint based on hashing molecular subgraphs |
| Atom Pairs | JCICS 25:64-73 (1985) |
| Topological | JCICS 27:82-5 (1987) |
| Torsions | |
| MACCS keys | Using the 166 public keys implemented as SMARTS |
| Morgan/Circular | Fingerprints based on the Morgan algorithm, similar to the ECFP/FCFP fingerprints JCIM |
| | 50 :742-54 (2010). |
| 2D | Uses topological distances between pharmacophoric points. |
| Pharmacophore | |
| Pattern | a topological fingerprint optimized for substructure screening |

3.18 Feature Definitions Used in the Morgan Fingerprints

These are adapted from the definitions in Gobbi, A. & Poppinger, D. "Genetic optimization of combinatorial libraries." *Biotechnology and Bioengineering* **61**, 47-54 (1998).

| Fea- | SMARTS | |
|--------|---|---------------|
| ture | | |
| Donor | [\$([N;!H0;v3,v4&+1]),\$([O,S;H1;+0]),n&H1&+0] | |
| Ac- | [\$([O,S;H1;v2;!\$(*-*=[O,N,P,S])]),\$([O,S;H0;v2]),\$([O,S;-]),\$([N;v3;!\$(| N-*=[O,N,P,S] |
| cep- | | |
| tor | | |
| Aro- | [a] | |
| matic | | |
| Halo- | [F,Cl,Br,I] | |
| gen | | |
| Basic | [#7;+,\$([N;H2&+0][\$([C,a]);!\$([C,a](=0))]),\$([N;H1&+0]([\$([C,a]);!\$([C, | a](=0))])[\$(|
| Acidic | [\$([C,S](=[O,S,P])-[O;H1,-1])] | |

3.19 License



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The intent of this license is similar to that of the RDKit itself. In simple words: "Do whatever you want with it, but please give us some credit."

CHAPTER

FOUR

THE RDKIT BOOK

4.1 Misc Cheminformatics Topics

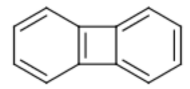
4.1.1 Aromaticity

Aromaticity is one of those unpleasant topics that is simultaneously simple and impossibly complicated. Since neither experimental nor theoretical chemists can agree with each other about a definition, it's necessary to pick something arbitrary and stick to it. This is the approach taken in the RDKit.

Instead of using patterns to match known aromatic systems, the aromaticity perception code in the RDKit uses a set of rules. The rules are relatively straightforward.

Aromaticity is a property of atoms and bonds in rings. An aromatic bond must be between aromatic atoms, but a bond between aromatic atoms does not need to be aromatic.

For example the fusing bonds here are not considered to be aromatic by the RDKit:



```
>>> from rdkit import Chem
>>> m = Chem.MolFromSmiles('C1=CC2=C(C=C1)C1=CC=CC=C21')
>>> m.GetAtomWithIdx(3).GetIsAromatic()
True
>>> m.GetAtomWithIdx(6).GetIsAromatic()
True
>>> m.GetBondBetweenAtoms(3,6).GetIsAromatic()
False
```

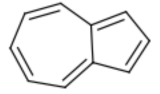
A ring, or fused ring system, is considered to be aromatic if it obeys the 4N+2 rule. Contributions to the electron count are determined by atom type and environment. Some examples:

| Fragment | Number of pi electrons |
|----------|------------------------|
| c(a)a | 1 |
| n(a)a | 1 |
| An(a)a | 2 |
| o(a)a | 2 |
| s(a)a | 2 |
| se(a)a | 2 |
| te(a)a | 2 |
| O=c(a)a | 0 |
| N=c(a)a | 0 |
| *(a)a | 0, 1, or 2 |

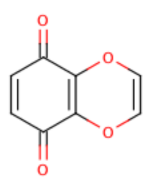
Notation a: any aromatic atom; A: any atom, include H; *: a dummy atom

Notice that exocyclic bonds to electronegative atoms "steal" the valence electron from the ring atom and that dummy atoms contribute whatever count is necessary to make the ring aromatic.

The use of fused rings for aromaticity can lead to situations where individual rings are not aromatic, but the fused system is. An example of this is azulene:



An extreme example, demonstrating both fused rings and the influence of exocyclic double bonds:



```
>>> m=Chem.MolFromSmiles('O=C1C=CC(=O)C2=C1OC=CO2')
>>> m.GetAtomWithIdx(6).GetIsAromatic()
True
>>> m.GetAtomWithIdx(7).GetIsAromatic()
True
>>> m.GetBondBetweenAtoms(6,7).GetIsAromatic()
False
```

A special case, heteroatoms with radicals are not considered candidates for aromaticity:



```
>>> m = Chem.MolFromSmiles('C1=C[N]C=C1')
>>> m.GetAtomWithIdx(0).GetIsAromatic()
False
>>> m.GetAtomWithIdx(2).GetIsAromatic()
False
>>> m.GetAtomWithIdx(2).GetNumRadicalElectrons()
1
```

Carbons with radicals, however, are still considered:



```
>>> m = Chem.MolFromSmiles('C1=[C]NC=C1')
>>> m.GetAtomWithIdx(0).GetIsAromatic()
True
>>> m.GetAtomWithIdx(1).GetIsAromatic()
True
>>> m.GetAtomWithIdx(1).GetNumRadicalElectrons()
1
```

Note: For reasons of computation expediency, aromaticity perception is only done for fused-ring systems where all members are at most 24 atoms in size.

4.1.2 Ring Finding and SSSR

[Section taken from "Getting Started" document]

As others have ranted about with more energy and eloquence than I intend to, the definition of a molecule's smallest set of smallest rings is not unique. In some high symmetry molecules, a "true" SSSR will give results that are unappealing. For example, the SSSR for cubane only contains 5 rings, even though there are "obviously" 6. This problem can be fixed by implementing a *small* (instead of *smallest*) set of smallest rings algorithm that returns symmetric results. This is the approach that we took with the RDKit.

Because it is sometimes useful to be able to count how many SSSR rings are present in the molecule, there is a GetSSSR function, but this only returns the SSSR count, not the potentially non-unique set of rings.

4.2 Chemical Reaction Handling

4.2.1 Reaction SMARTS

Not SMIRKS ¹, not reaction SMILES ², derived from SMARTS ³.

The general grammar for a reaction SMARTS is:

Some features

Mapped dummy atoms in the product template are replaced by the corresponding atom in the reactant:

```
>>> from rdkit.Chem import AllChem
>>> rxn = AllChem.ReactionFromSmarts('[C:1]=[0,N:2]>>[C:1][*:2]')
>>> [Chem.MolToSmiles(x,1) for x in rxn.RunReactants((Chem.MolFromSmiles('CC=O'),))[0]]
['CCO']
>>> [Chem.MolToSmiles(x,1) for x in rxn.RunReactants((Chem.MolFromSmiles('CC=N'),))[0]]
['CCN']
```

but unmapped dummy atoms are left as dummies:

```
>>> rxn = AllChem.ReactionFromSmarts('[C:1]=[0,N:2]>>[*][C:1][*:2]')
>>> [Chem.MolToSmiles(x,1) for x in rxn.RunReactants((Chem.MolFromSmiles('CC=0'),))[0]]
['[*]C(C)0']
```

"Any" bonds in the products are replaced by the corresponding bond in the reactant:

```
>>> rxn = AllChem.ReactionFromSmarts('[C:1]~[0,N:2]>>[*][C:1]~[*:2]')  
>>> [Chem.MolToSmiles(x,1) for x in rxn.RunReactants((Chem.MolFromSmiles('C=0'),))[0]]
```

¹ http://www.daylight.com/dayhtml/doc/theory/theory.smirks.html

² http://www.daylight.com/dayhtml/doc/theory/theory.smiles.html

³ http://www.daylight.com/dayhtml/doc/theory/theory.smarts.html

```
['[*]C=O']
>>> [Chem.MolToSmiles(x,1) for x in rxn.RunReactants((Chem.MolFromSmiles('CO'),))[0]]
['[*]CO']
>>> [Chem.MolToSmiles(x,1) for x in rxn.RunReactants((Chem.MolFromSmiles('C#N'),))[0]]
['[*]C#N']
```

Intramolecular reactions can be expressed flexibly by including reactants in parentheses. This is demonstrated in this ring-closing metathesis example ⁴:

```
>>> rxn = AllChem.ReactionFromSmarts("([C:1]=[C;H2].[C:2]=[C;H2])>>[*:1]=[*:2]")
>>> m1 = Chem.MolFromSmiles('C=CCOCC=C')
>>> ps = rxn.RunReactants((m1,))
>>> Chem.MolToSmiles(ps[0][0])
'C1=CCOC1'
```

Chirality

This section describes how chirality information in the reaction defition is handled. A consistent example, esterification of secondary alcohols, is used throughout ⁵.

If no chiral information is present in the reaction definition, the stereochemistry of the reactants is preserved:

```
>>> alcohol1 = Chem.MolFromSmiles('CC(CCN)O')
>>> alcohol2 = Chem.MolFromSmiles('C[C@H](CCN)O')
>>> alcohol3 = Chem.MolFromSmiles('CC(@H](CCN)O')
>>> acid = Chem.MolFromSmiles('CC(=O)O')
>>> rxn = AllChem.ReactionFromSmarts('[CH1:1][OH:2].[OH][C:3]=[O:4]>>[C:1][O:2][C:3]=[O:4]')
>>> ps=rxn.RunReactants((alcohol1,acid))
>>> Chem.MolToSmiles(ps[0][0],True)
'CC(=O)OC(C)CCN'
>>> ps=rxn.RunReactants((alcohol2,acid))
>>> Chem.MolToSmiles(ps[0][0],True)
'CC(=O)O[C@H](C)CCN'
>>> ps=rxn.RunReactants((alcohol3,acid))
>>> Chem.MolToSmiles(ps[0][0],True)
'CC(=O)O[C@H](C)CCN'
```

You get the same result (retention of stereochemistry) if a mapped atom has the same chirality in both reactants and products:

```
>>> rxn = AllChem.ReactionFromSmarts('[C@H1:1][OH:2].[OH][C:3]=[O:4]>>[C@:1][O:2][C:3]=[O:4]')
>>> ps=rxn.RunReactants((alcohol1,acid))
>>> Chem.MolToSmiles(ps[0][0],True)
'CC(=0)OC(C)CCN'
>>> ps=rxn.RunReactants((alcohol2,acid))
>>> Chem.MolToSmiles(ps[0][0],True)
'CC(=0)O[C@H](C)CCN'
>>> ps=rxn.RunReactants((alcohol3,acid))
>>> Chem.MolToSmiles(ps[0][0],True)
'CC(=0)O[C@H](C)CCN'
```

A mapped atom with different chirality in reactants and products leads to inversion of stereochemistry:

```
>>> rxn = AllChem.ReactionFromSmarts('[C@H1:1][OH:2].[OH][C:3]=[0:4]>>[C@@:1][0:2][C:3]=[0:4]')
>>> ps=rxn.RunReactants((alcohol1,acid))
>>> Chem.MolToSmiles(ps[0][0],True)
```

⁴ Thanks to James Davidson for this example.

⁵ Thanks to JP Ebejer and Paul Finn for this example.

```
'CC(=0)OC(C)CCN'
>>> ps=rxn.RunReactants((alcohol2,acid))
>>> Chem.MolToSmiles(ps[0][0],True)
'CC(=0)O[C@@H](C)CCN'
>>> ps=rxn.RunReactants((alcohol3,acid))
>>> Chem.MolToSmiles(ps[0][0],True)
'CC(=0)O[C@H](C)CCN'
```

If a mapped atom has chirality specified in the reactants, but not in the products, the reaction destroys chirality at that center:

```
>>> rxn = AllChem.ReactionFromSmarts('[C@H1:1][OH:2].[OH][C:3]=[0:4]>>[C:1][0:2][C:3]=[0:4]')
>>> ps=rxn.RunReactants((alcohol1,acid))
>>> Chem.MolToSmiles(ps[0][0],True)
'CC(=0)OC(C)CCN'
>>> ps=rxn.RunReactants((alcohol2,acid))
>>> Chem.MolToSmiles(ps[0][0],True)
'CC(=0)OC(C)CCN'
>>> ps=rxn.RunReactants((alcohol3,acid))
>>> Chem.MolToSmiles(ps[0][0],True)
'CC(=0)OC(C)CCN'
```

And, finally, if chirality is specified in the products, but not the reactants, the reaction creates a stereocenter with the specified chirality:

```
>>> rxn = AllChem.ReactionFromSmarts('[CH1:1][OH:2].[OH][C:3]=[O:4]>>[C@:1][O:2][C:3]=[O:4]')
>>> ps=rxn.RunReactants((alcohol1,acid))
>>> Chem.MolToSmiles(ps[0][0],True)
'CC(=0)0[C@H](C)CCN'
>>> ps=rxn.RunReactants((alcohol2,acid))
>>> Chem.MolToSmiles(ps[0][0],True)
'CC(=0)0[C@H](C)CCN'
>>> ps=rxn.RunReactants((alcohol3,acid))
>>> Chem.MolToSmiles(ps[0][0],True)
'CC(=0)0[C@H](C)CCN'
```

Note that this doesn't make sense without including a bit more context around the stereocenter in the reaction definition:

```
>>> rxn = AllChem.ReactionFromSmarts('[CH3:5][CH1:1]([C:6])[OH:2].[OH][C:3]=[O:4]>>[C:5][C@:1]([C:6])
>>> ps=rxn.RunReactants((alcohol1,acid))
>>> Chem.MolToSmiles(ps[0][0],True)
'CC(=0)O[C@H](C)CCN'
>>> ps=rxn.RunReactants((alcohol2,acid))
>>> Chem.MolToSmiles(ps[0][0],True)
'CC(=0)O[C@H](C)CCN'
>>> ps=rxn.RunReactants((alcohol3,acid))
>>> Chem.MolToSmiles(ps[0][0],True)
```

Note that the chirality specification is not being used as part of the query: a molecule with no chirality specified can match a reactant with specified chirality.

In general, the reaction machinery tries to preserve as much stereochemistry information as possible. This works when a single new bond is formed to a chiral center:

```
>>> rxn = AllChem.ReactionFromSmarts('[C:1][C:2]-0>>[C:1][C:2]-S')
>>> alcohol2 = Chem.MolFromSmiles('C[C@@H](O)CCN')
>>> ps=rxn.RunReactants((alcohol2,))
```

'CC(=0)0[C@H](C)CCN'

```
>>> Chem.MolToSmiles(ps[0][0],True)
'C[C@@H](S)CCN'
```

But it fails if two or more bonds are formed:

```
>>> rxn = AllChem.ReactionFromSmarts('[C:1][C:2](-0)-F>>[C:1][C:2](-S)-Cl')
>>> alcohol = Chem.MolFromSmiles('C[C@@H](O)F')
>>> ps=rxn.RunReactants((alcohol,))
>>> Chem.MolToSmiles(ps[0][0],True)
'CC(S)Cl'
```

In this case, there's just not sufficient information present to allow the information to be preserved. You can help by providing mapping information:

Rules and caveats

- 1. Include atom map information at the end of an atom query. So do [C,N,O:1] or [C;R:1].
- 2. Don't forget that unspecified bonds in SMARTS are either single or aromatic. Bond orders in product templates are assigned when the product template itself is constructed and it's not always possible to tell if the bond should be single or aromatic:

```
>>> rxn = AllChem.ReactionFromSmarts('[#6:1][#7,#8:2]>>[#6:1][#6:2]')
>>> [Chem.MolToSmiles(x,1) for x in rxn.RunReactants((Chem.MolFromSmiles('C1NCCCC1'),))[0]]
['C1CCCCC1']
>>> [Chem.MolToSmiles(x,1) for x in rxn.RunReactants((Chem.MolFromSmiles('c1ncccc1'),))[0]]
['c1ccccc-1']
```

So if you want to copy the bond order from the reactant, use an "Any" bond:

```
>>> rxn = AllChem.ReactionFromSmarts('[#6:1][#7,#8:2]>>[#6:1]~[#6:2]')
>>> [Chem.MolToSmiles(x,1) for x in rxn.RunReactants((Chem.MolFromSmiles('clnccc1'),))[0]]
['clcccc1']
```

4.3 The Feature Definition File Format

An FDef file contains all the information needed to define a set of chemical features. It contains definitions of feature types that are defined from queries built up using Daylight's SMARTS language. ³ The FDef file can optionally also include definitions of atom types that are used to make feature definitions more readable.

4.3.1 Chemical Features

Chemical features are defined by a Feature Type and a Feature Family. The Feature Family is a general classification of the feature (such as "Hydrogen-bond Donor" or "Aromatic") while the Feature Type provides additional, higher-resolution, information about features. Pharmacophore matching is done using Feature Family's. Each feature type contains the following pieces of information:

- A SMARTS pattern that describes atoms (one or more) matching the feature type.
- Weights used to determine the feature's position based on the positions of its defining atoms.

4.3.2 Syntax of the FDef file

AtomType definitions

An AtomType definition allows you to assign a shorthand name to be used in place of a SMARTS string defining an atom query. This allows FDef files to be made much more readable. For example, defining a non-polar carbon atom like this:

```
AtomType Carbon_NonPolar [C&!$(C=[O,N,P,S])&!$(C#N)]
```

creates a new name that can be used anywhere else in the FDef file that it would be useful to use this SMARTS. To reference an AtomType, just include its name in curly brackets. For example, this excerpt from an FDef file defines another atom type - Hphobe - which references the Carbon NonPolar definition:

```
AtomType Carbon_NonPolar [C&!$(C=[O,N,P,S])&!$(C#N)]
AtomType Hphobe [{Carbon_NonPolar},c,s,S&H0&v2,F,Cl,Br,I]
```

Note that {Carbon_NonPolar} is used in the new AtomType definition without any additional decoration (no square brackes or recursive SMARTS markers are required).

Repeating an AtomType results in the two definitions being combined using the SMARTS "," (or) operator. Here's an example:

```
AtomType d1 [N&!H0]
AtomType d1 [O&!H0]
```

This is equivalent to:

```
AtomType d1 [N&!H0,O&!H0]
```

Which is equivalent to the more efficient:

```
AtomType d1 [N,O;!H0]
```

Note that these examples tend to use SMARTS's high-precendence and operator "&" and not the low-precedence and ";". This can be important when AtomTypes are combined or when they are repeated. The SMARTS "," operator is higher precedence than ";", so definitions that use ";" can lead to unexpected results.

It is also possible to define negative AtomType queries:

```
AtomType d1 [N,O,S]
AtomType !d1 [H0]
```

The negative query gets combined with the first to produce a definition identical to this:

```
AtomType d1 [!H0;N,O,S]
```

Note that the negative AtomType is added to the beginning of the query.

Feature definitions

A feature definition is more complex than an AtomType definition and stretches across multiple lines:

```
DefineFeature HDonor1 [N,O;!H0]
Family HBondDonor
Weights 1.0
EndFeature
```

The first line of the feature definition includes the feature type and the SMARTS string defining the feature. The next two lines (order not important) define the feature's family and its atom weights (a comma-delimited list that is the same length as the number of atoms defining the feature). The atom weights are used to calculate the feature's locations based on a weighted average of the positions of the atom defining the feature. More detail on this is provided below. The final line of a feature definition must be EndFeature. It is perfectly legal to mix AtomType definitions with feature definitions in the FDef file. The one rule is that AtomTypes must be defined before they are referenced.

Additional syntax notes:

- Any line that begins with a # symbol is considered a comment and will be ignored.
- A backslash character, , at the end of a line is a continuation character, it indicates that the data from that line is continued on the next line of the file. Blank space at the beginning of these additional lines is ignored. For example, this AtomType definition:

```
AtomType tButylAtom [$([C;!R](-[CH3])(-[CH3])(-[CH3])),\
$([CH3](-[C;!R](-[CH3])(-[CH3]))]

is exactly equivalent to this one:

AtomType tButylAtom [$([C;!R](-[CH3])(-[CH3])(-[CH3])),$([CH3](-[C;!R](-[CH3])))]

(though the first form is much easier to read!)
```

Atom weights and feature locations

4.3.3 Frequently Asked Question(s)

• What happens if a Feature Type is repeated in the file? Here's an example:

```
DefineFeature HDonor1 [O&!H0]
Family HBondDonor
Weights 1.0
EndFeature

DefineFeature HDonor1 [N&!H0]
Family HBondDonor
Weights 1.0
EndFeature
```

In this case both definitions of the HDonor1 feature type will be active. This is functionally identical to:

```
DefineFeature HDonor1 [O,N;!H0]
Family HBondDonor
Weights 1.0
EndFeature
```

However the formulation of this feature definition with a duplicated feature type is considerably less efficient and more confusing than the simpler combined definition.

4.4 Representation of Pharmacophore Fingerprints

In the RDKit scheme the bit ids in pharmacophore fingerprints are not hashed: each bit corresponds to a particular combination of features and distances. A given bit id can be converted back to the corresponding feature types and

distances to allow interpretation. An illustration for 2D pharmacophores is shown in Figure 1: Bit numbering in pharmacophore fingerprints.

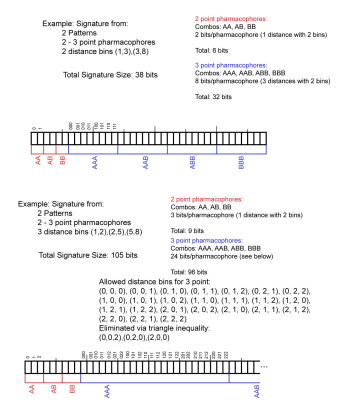


Figure 4.1: Figure 1: Bit numbering in pharmacophore fingerprints

4.5 Atom-Atom Matching in Substructure Queries

When doing substructure matches for queries derived from SMARTS the rules for which atoms in the molecule should match which atoms in the query are well defined.[#smarts]_ The same is not necessarily the case when the query molecule is derived from a mol block or SMILES.

The general rule used in the RDKit is that if you don't specify a property in the query, then it's not used as part of the matching criteria and that Hs are ignored. This leads to the following behavior:

| Molecule | Query | Match |
|----------|---------|-------|
| CCO | CCO | Yes |
| | CCO | Yes |
| CC[O-] | | |
| CCO | CC[O-] | No |
| CC[O-] | CC[O-] | Yes |
| CC[O-] | CC[OH] | Yes |
| CCOC | CC[OH] | Yes |
| CCOC | CCO | Yes |
| CCC | CCC | Yes |
| CC[14C] | CCC | Yes |
| CCC | CC[14C] | No |
| CC[14C] | CC[14C] | Yes |
| OCO | С | Yes |
| OCO | [CH] | No |
| OCO | [CH2] | No |
| OCO | [CH3] | No |
| OCO | O[CH3] | Yes |
| O[CH2]O | С | Yes |
| O[CH2]O | [CH2] | No |

4.6 Implementation Details

4.6.1 "Magic" Property Values

The following property values are regularly used in the RDKit codebase and may be useful to client code.

ROMol (Mol in Python)

| Property Name | Use |
|------------------------|---|
| MolFileComments | Read from/written to the comment line of CTABs. |
| MolFileInfo | Read from/written to the info line of CTABs. |
| _MolFileChiralFlag | Read from/written to the chiral flag of CTABs. |
| _Name | Read from/written to the name line of CTABs. |
| _smilesAtomOutputOrder | The order in which atoms were written to SMILES |

Atom

| Property Name | Use |
|------------------|---|
| _CIPCode | the CIP code (R or S) of the atom |
| _CIPRank | the integer CIP rank of the atom |
| _ChiralityPossi- | set if an atom is a possible chiral center |
| ble | |
| _MolFileRLabel | integer R group label for an atom, read from/written to CTABs. |
| _ReactionDe- | set on an atom in a product template of a reaction if its degree changes in the reaction |
| greeChanged | |
| _protected | atoms with this property set will not be considered as matching reactant queries in reactions |
| dummyLabel | (on dummy atoms) read from/written to CTABs as the atom symbol |
| molAtomMap- | the atom map number for an atom, read from/written to SMILES and CTABs |
| Number | |
| molfileAlias | the mol file alias for an atom (follows A tags), read from/written to CTABs |
| molFileValue | the mol file value for an atom (follows V tags), read from/written to CTABs |
| molFileInver- | used to flag whether stereochemistry at an atom changes in a reaction, read from/written to |
| sionFlag | CTABs, determined automatically from SMILES |
| molRxnCompo- | which component of a reaction an atom belongs to, read from/written to CTABs |
| nent | |
| molRxnRole | which role an atom plays in a reaction (1=Reactant, 2=Product, 3=Agent), read from/written |
| | to CTABs |
| smilesSymbol | determines the symbol that will be written to a SMILES for the atom |

4.6.2 Thread safety and the RDKit

While writing the RDKit, we did attempt to ensure that the code would work in a multi-threaded environment by avoiding use of global variables, etc. However, making code thread safe is not a completely trivial thing, so there are no doubt some gaps. This section describes which pieces of the code base have explicitly been tested for thread safety.

Note: With the exception of the small number of methods/functions that take a numThreads argument, this section does not apply to using the RDKit from Python threads. Boost.Python ensures that only one thread is calling into the C++ code at any point. To get concurrent execution in Python, use the multiprocessing module or one of the other standard python approaches for this.

What has been tested

- Reading molecules from SMILES/SMARTS/Mol blocks
- Writing molecules to SMILES/SMARTS/Mol blocks
- Generating 2D coordinates
- Generating 3D conformations with the distance geometry code
- Optimizing molecules with UFF or MMFF
- · Generating fingerprints
- The descriptor calculators in \$RDBASE/Code/GraphMol/Descriptors
- Substructure searching (Note: if a query molecule contains recursive queries, it may not be safe to use it concurrently on multiple threads, see below)
- The Subgraph code
- The ChemTransforms code

- The chemical reactions code
- The Open3DAlign code
- The MolDraw2D drawing code

Known Problems

- InChI generation and (probably) parsing. This seems to be a limitation of the IUPAC InChI code. In order to allow the code to be used in a multi-threaded environment, a mutex is used to ensure that only one thread is using the IUPAC code at a time. This is only enabled if the RDKit is built with the RDK_TEST_MULTITHREADED option enabled.
- The MolSuppliers (e.g. SDMolSupplier, SmilesMolSupplier?) change their internal state when a molecule is read. It is not safe to use one supplier on more than one thread.
- Substructure searching using query molecules that include recursive queries. The recursive queries modify their internal state when a search is run, so it's not safe to use the same query concurrently on multiple threads. If the code is built using the RDK_BUILD_THREADSAFE_SSS argument (the default for the binaries we provide), a mutex is used to ensure that only one thread is using a given recursive query at a time.

4.7 License



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The intent of this license is similar to that of the RDKit itself. In simple words: "Do whatever you want with it, but please give us some credit."

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CHAPTER

FIVE

RDKIT COOKBOOK

5.1 What is this?

This document provides examples of how to carry out particular tasks using the RDKit functionality from Python. The contents have been contributed by the RDKit community.

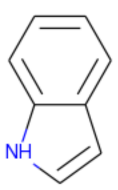
If you find mistakes, or have suggestions for improvements, please either fix them yourselves in the source document (the .rst file) or send them to the mailing list: rdkit-discuss@lists.sourceforge.net (you will need to subscribe first)

5.2 Miscellaneous Topics

5.2.1 Using a different aromaticity model

By default, the RDKit applies its own model of aromaticity (explained in the RDKit Theory Book) when it reads in molecules. It is, however, fairly easy to override this and use your own aromaticity model.

The easiest way to do this is it provide the molecules as SMILES with the aromaticity set as you would prefer to have it. For example, consider indole:



By default the RDKit considers both rings to be aromatic:

```
>>> from rdkit import Chem
>>> m = Chem.MolFromSmiles('N1C=Cc2cccc12')
>>> m.GetSubstructMatches(Chem.MolFromSmarts('c'))
((1,), (2,), (3,), (4,), (5,), (6,), (7,), (8,))
```

If you'd prefer to treat the five-membered ring as aliphatic, which is how the input SMILES is written, you just need to do a partial sanitization that skips the kekulization and aromaticity perception steps:

```
>>> m2 = Chem.MolFromSmiles('N1C=Cc2cccc12', sanitize=False)
>>> Chem.SanitizeMol(m2, sanitizeOps=Chem.SanitizeFlags.SANITIZE_ALL^Chem.SanitizeFlags.SANITIZE_KEKU:
rdkit.Chem.rdmolops.SanitizeFlags.SANITIZE_NONE
>>> m2.GetSubstructMatches(Chem.MolFromSmarts('c'))
((3,), (4,), (5,), (6,), (7,), (8,))
```

It is, of course, also possible to write your own aromaticity perception function, but that is beyond the scope of this document.

5.3 Manipulating Molecules

5.3.1 Cleaning up heterocycles

Mailing list discussions:

- http://www.mail-archive.com/rdkit-discuss@lists.sourceforge.net/msg01185.html
- http://www.mail-archive.com/rdkit-discuss@lists.sourceforge.net/msg01162.html
- http://www.mail-archive.com/rdkit-discuss@lists.sourceforge.net/msg01900.html

• http://www.mail-archive.com/rdkit-discuss@lists.sourceforge.net/msg01901.html

The code:

```
""" sanifix4.py
 Contribution from James Davidson
from rdkit import Chem
from rdkit.Chem import AllChem
def _FragIndicesToMol(oMol,indices):
    em = Chem.EditableMol(Chem.Mol())
    newIndices={}
    for i,idx in enumerate(indices):
        em.AddAtom(oMol.GetAtomWithIdx(idx))
        newIndices[idx]=i
    for i,idx in enumerate(indices):
        at = oMol.GetAtomWithIdx(idx)
        for bond in at.GetBonds():
            if bond.GetBeginAtomIdx() == idx:
                oidx = bond.GetEndAtomIdx()
            else:
                oidx = bond.GetBeginAtomIdx()
            # make sure every bond only gets added once:
            if oidx<idx:</pre>
                continue
            em.AddBond(newIndices[idx],newIndices[oidx],bond.GetBondType())
    res = em.GetMol()
    res.ClearComputedProps()
    Chem.GetSymmSSSR(res)
    res.UpdatePropertyCache (False)
    res._idxMap=newIndices
    return res
def _recursivelyModifyNs(mol,matches,indices=None):
    if indices is None:
        indices=[]
    res=None
    while len (matches) and res is None:
        tIndices=indices[:]
        nextIdx = matches.pop(0)
        tIndices.append(nextIdx)
        nm = Chem.Mol(mol)
        nm.GetAtomWithIdx(nextIdx).SetNoImplicit(True)
        nm.GetAtomWithIdx(nextIdx).SetNumExplicitHs(1)
        cp = Chem.Mol(nm)
        try:
            Chem.SanitizeMol(cp)
        except ValueError:
            res, indices = _recursivelyModifyNs(nm, matches, indices=tIndices)
        else:
            indices=tIndices
            res=cp
    return res,indices
def AdjustAromaticNs(m, nitrogenPattern='[n&D2&H0;r5,r6]'):
```

```
n n n
       default nitrogen pattern matches Ns in 5 rings and 6 rings in order to be able
       to fix: O=c1ccncc1
    Chem.GetSymmSSSR(m)
    m.UpdatePropertyCache(False)
    # break non-ring bonds linking rings:
    em = Chem.EditableMol(m)
    linkers = m.GetSubstructMatches(Chem.MolFromSmarts('[r]!@[r]'))
    plsFix=set()
    for a,b in linkers:
        em.RemoveBond(a,b)
        plsFix.add(a)
        plsFix.add(b)
    nm = em.GetMol()
    for at in plsFix:
        at=nm.GetAtomWithIdx(at)
        if at.GetIsAromatic() and at.GetAtomicNum() == 7:
            at.SetNumExplicitHs(1)
            at.SetNoImplicit(True)
    # build molecules from the fragments:
    fragLists = Chem.GetMolFrags(nm)
    frags = [_FragIndicesToMol(nm,x) for x in fragLists]
    # loop through the fragments in turn and try to aromatize them:
    for i, frag in enumerate(frags):
        cp = Chem.Mol(frag)
        try:
            Chem.SanitizeMol(cp)
        except ValueError:
            matches = [x[0]] for x in frag.GetSubstructMatches(Chem.MolFromSmarts(nitrogenPattern))]
            lres, indices=_recursivelyModifyNs (frag, matches)
            if not lres:
                #print 'frag %d failed (%s)'%(i,str(fragLists[i]))
                ok=False
                break
            else:
                revMap={}
                for k, v in frag._idxMap.iteritems():
                    revMap[v]=k
                for idx in indices:
                    oatom = m.GetAtomWithIdx(revMap[idx])
                    oatom.SetNoImplicit(True)
                    oatom.SetNumExplicitHs(1)
    if not ok:
        return None
    return m
Examples of using it:
smis= ('0=c1ccc2cccc2n1',
       'Cc1nnnn1C',
       'CCclccc2nc(=0)c(cc2cl)Cclnnnn1ClCCCCCl',
       'clcnc2cc3ccnc3cc12',
       'clcc2cc3ccnc3cc2n1',
       'O=c1ccnc(c1)-c1cnc2cc3ccnc3cc12',
```

```
'O=c1ccnc(c1)-c1cc1',
for smi in smis:
      m = Chem.MolFromSmiles(smi,False)
      try:
          m.UpdatePropertyCache(False)
          cp = Chem.Mol(m)
          Chem.SanitizeMol(cp)
          m = cp
          print 'fine:',Chem.MolToSmiles(m)
      except ValueError:
          nm=AdjustAromaticNs(m)
          if nm is not None:
              Chem.SanitizeMol(nm)
              print 'fixed:',Chem.MolToSmiles(nm)
          else:
              print 'still broken:',smi
This produces:
fixed: O=c1ccc2cccc2[nH]1
fine: Cclnnnn1C
fixed: CCclccc2[nH]c(=0)c(Cc3nnnn3C3CCCCC3)cc2c1
fine: C1=Cc2cc3c(cc2=N1)C=CN=3
fine: C1=Cc2cc3c(cc2=N1)N=CC=3
fixed: O=c1cc[nH]c(C2=CN=c3cc4c(cc32)=NC=C4)c1
still broken: O=c1ccnc(c1)-c1cc1
```

5.3.2 Parallel conformation generation

Mailing list discussion: http://www.mail-archive.com/rdkit-discuss@lists.sourceforge.net/msg02648.html

The code:

```
""" contribution from Andrew Dalke """
import sys
from rdkit import Chem
from rdkit. Chem import AllChem
# Download this from http://pypi.python.org/pypi/futures
from concurrent import futures
# Download this from http://pypi.python.org/pypi/progressbar
import progressbar
## On my machine, it takes 39 seconds with 1 worker and 10 seconds with 4 \cdot
## 29.055u 0.102s 0:28.68 101.6% 0+0k 0+3io 0pf+0w
#max_workers=1
## With 4 threads it takes 11 seconds.
## 34.933u 0.188s 0:10.89 322.4% 0+0k 125+1io 0pf+0w
max_workers=4
# (The "u"ser time includes time spend in the children processes.
# The wall-clock time is 28.68 and 10.89 seconds, respectively.)
# This function is called in the subprocess.
# The parameters (molecule and number of conformers) are passed via a Python
```

```
def generateconformations(m, n):
   m = Chem.AddHs(m)
    \verb|ids=AllChem.EmbedMultipleConfs(m, numConfs=n)|\\
    for id in ids:
        AllChem.UFFOptimizeMolecule(m, confId=id)
    # EmbedMultipleConfs returns a Boost-wrapped type which
    # cannot be pickled. Convert it to a Python list, which can.
    return m, list(ids)
smi_input_file, sdf_output_file = sys.argv[1:3]
n = int(sys.argv[3])
writer = Chem.SDWriter(sdf_output_file)
suppl = Chem.SmilesMolSupplier(smi_input_file, titleLine=False)
with futures.ProcessPoolExecutor(max_workers=max_workers) as executor:
    # Submit a set of asynchronous jobs
    jobs = []
    for mol in suppl:
        if mol:
            job = executor.submit(generateconformations, mol, n)
            jobs.append(job)
    widgets = ["Generating conformations; ", progressbar.Percentage(), " ",
              progressbar.ETA(), " ", progressbar.Bar()]
    pbar = progressbar.ProgressBar(widgets=widgets, maxval=len(jobs))
    for job in pbar(futures.as_completed(jobs)):
        mol,ids=job.result()
        for id in ids:
            writer.write(mol, confId=id)
writer.close()
```

5.3.3 Neutralizing Charged Molecules

Mailing list discussion: http://www.mail-archive.com/rdkit-discuss@lists.sourceforge.net/msg02648.html

The code:

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```
""" contribution from Hans de Winter """
from rdkit import Chem
from rdkit.Chem import AllChem
def _InitialiseNeutralisationReactions():
    patts= (
        # Imidazoles
        ('[n+;H]','n'),
        # Amines
        ('[N+;!H0]','N'),
        # Carboxylic acids and alcohols
        ('[$([O-]);!$([O-][#7])]','O'),
        # Thiols
        ('[S-;X1]','S'),
        # Sulfonamides
        ('[$([N-;X2]S(=0)=0)]','N'),
        # Enamines
```

```
('[\$([N-;X2][C,N]=C)]','N'),
        # Tetrazoles
        ('[n-]','[nH]'),
        # Sulfoxides
        ('[\$([S-]=0)]','S'),
        # Amides
        ('[\$([N-]C=O)]','N'),
    return [(Chem.MolFromSmarts(x), Chem.MolFromSmiles(y, False)) for x,y in patts]
_reactions=None
def NeutraliseCharges(smiles, reactions=None):
    global _reactions
    if reactions is None:
        if _reactions is None:
            _reactions=_InitialiseNeutralisationReactions()
        reactions=_reactions
    mol = Chem.MolFromSmiles(smiles)
    replaced = False
    for i,(reactant, product) in enumerate(reactions):
        while mol.HasSubstructMatch(reactant):
            replaced = True
            rms = AllChem.ReplaceSubstructs(mol, reactant, product)
            mol = rms[0]
    if replaced:
        return (Chem.MolToSmiles(mol,True), True)
    else:
        return (smiles, False)
Examples of using it:
smis=("c1cccc[nH+]1",
      "C[N+](C)(C)C", "c1ccccc1[NH3+]",
      "CC(=0)[0-]", "c1ccccc1[0-]",
      "CCS",
      "C[N-]S(=0)(=0)C",
      "C[N-]C=C", "C[N-]N=C",
      "c1ccc[n-]1",
      "CC[N-]C(=0)CC")
for smi in smis:
    (molSmiles, neutralised) = NeutraliseCharges(smi)
    print smi,"->", molSmiles
This produces:
clcccc[nH+]1 -> clccncc1
C[N+](C)(C)C -> C[N+](C)(C)C
clccccc1[NH3+] -> Nclccccc1
CC (=0) [O-] -> CC (=0) O
clccccc1[0-] -> Oclccccc1
CCS -> CCS
C[N-]S(=0)(=0)C -> CNS(C)(=0)=0
C[N-]C=C -> C=CNC
C[N-]N=C -> C=NNC
clccc[n-]1 \rightarrow clcc[nH]c1
CC[N-]C(=0)CC \rightarrow CCNC(=0)CC
```

5.3.4 3D functionality in the RDKit

The RDKit contains a range of 3D functionalities such as:

| Shape alignment |
|---------------------------------|
| RMS calculation |
| Shape Tanimoto Distance |
| Shape Protrude Distance |
| 3D pharmacophore fingerprint |
| Torsion fingerprint (deviation) |

There are two alignment methods currently available in the RDKit. As an example we use two crystal structures from the PDB of the same molecule.

The code:

```
from rdkit import Chem, RDConfig
from rdkit. Chem import AllChem, rdMolAlign
# The reference molecule
ref = Chem. MolFromSmiles ('NC(=[NH2+])c1ccc(C[C@@H](NC(=0)CNS(=0)(=0)c2ccc3cccc3c2)C(=0)N2CCCC2)cc1
# The PDB conformations
mol1 = Chem.MolFromPDBFile(RDConfig.RDBaseDir+'/rdkit/Chem/test_data/1DWD_ligand.pdb')
mol1 = AllChem.AssignBondOrdersFromTemplate(ref, mol1)
mol2 = Chem.MolFromPDBFile(RDConfig.RDBaseDir+'/rdkit/Chem/test_data/1PPC_ligand.pdb')
mol2 = AllChem.AssignBondOrdersFromTemplate(ref, mol2)
# Align them
rms = rdMolAlign.AlignMol(mol1, mol2)
print rms
# Align them with OPEN3DAlign
pyO3A = rdMolAlign.GetO3A(mol1, mol2)
score = pyO3A.Align()
print score
This produces:
1.55001955728
0.376459885045
```

If a molecule contains more than one conformer, they can be aligned with respect to the first conformer. If a list is provided to the option RMSlist, the RMS value from the alignment are stored. The RMS value of two conformers of a molecule can also be calculated separately, either with or without alignment (using the flag prealigned).

Examples of using it:

```
from rdkit import Chem
from rdkit.Chem import AllChem
mol = Chem.MolFromSmiles('NC(=[NH2+])clccc(C[C@@H](NC(=0)CNS(=0)(=0)c2ccc3cccc3c2)C(=0)N2CCCC2)ccl
cids = AllChem.EmbedMultipleConfs(mol, numConfs=50, maxAttempts=1000, pruneRmsThresh=0.1)
print len(cids)
# align the conformers
rmslist = []
AllChem.AlignMolConformers(mol, RMSlist=rmslist)
print len(rmslist)
# calculate RMS of confomers 1 and 9 separately
rms = AllChem.GetConformerRMS(mol, 1, 9, prealigned=True)
This produces:
```

50

49

For shape comparison, the RDKit provides two Shape-based distances that can be calculated for two prealigned molecules or conformers. Shape protrude distance focusses on the volume mismatch, while Shape Tanimoto distance takes the volume overlay overall into account.

Examples of using it:

```
from rdkit import Chem, RDConfig
from rdkit.Chem import AllChem, rdMolAlign, rdShapeHelpers
ref = Chem.MolFromSmiles('NC(=[NH2+])clcc(C[C@@H](NC(=0)CNS(=0)(=0)c2ccc3cccc3c2)C(=0)N2CCCC2)ccl
mol1 = Chem.MolFromPDBFile(RDConfig.RDBaseDir+'/rdkit/Chem/test_data/1DWD_ligand.pdb')
mol1 = AllChem.AssignBondOrdersFromTemplate(ref, mol1)
mol2 = Chem.MolFromPDBFile(RDConfig.RDBaseDir+'/rdkit/Chem/test_data/1PPC_ligand.pdb')
mol2 = AllChem.AssignBondOrdersFromTemplate(ref, mol2)
rms = rdMolAlign.AlignMol(mol1, mol2)
tani = rdShapeHelpers.ShapeTanimotoDist(mol1, mol2)
prtr = rdShapeHelpers.ShapeProtrudeDist(mol1, mol2)
print rms, tani, prtr
```

This produces:

```
1.55001955728 0.18069102331 0.0962800875274
```

A 3D pharmacophore fingerprint can be calculated using the RDKit by feeding a 3D distance matrix to the 2D-pharmacophore machinery.

Examples of using it:

```
from rdkit import Chem, DataStructs, RDConfig
from rdkit.Chem import AllChem
from rdkit.Chem.Pharm2D import Gobbi_Pharm2D, Generate

ref = Chem.MolFromSmiles('NC(=[NH2+])clccc(C[C@@H](NC(=0)CNS(=0)(=0)c2ccc3cccc3c2)C(=0)N2CCCCC2)ccl
mol1 = Chem.MolFromPDBFile(RDConfig.RDBaseDir+'/rdkit/Chem/test_data/1DWD_ligand.pdb')
mol1 = AllChem.AssignBondOrdersFromTemplate(ref, mol1)
mol2 = Chem.MolFromPDBFile(RDConfig.RDBaseDir+'/rdkit/Chem/test_data/1PPC_ligand.pdb')
mol2 = AllChem.AssignBondOrdersFromTemplate(ref, mol2)
# pharmacophore fingerprint
factory = Gobbi_Pharm2D.factory
fp1 = Generate.Gen2DFingerprint(mol1, factory, dMat=Chem.Get3DDistanceMatrix(mol1))
fp2 = Generate.Gen2DFingerprint(mol2, factory, dMat=Chem.Get3DDistanceMatrix(mol2))
# Tanimoto similarity
tani = DataStructs.TanimotoSimilarity(fp1, fp2)
print tani
```

This produces:

```
0.451665312754
```

The RDKit provides an implementation of the torsion fingerprint deviation (TFD) approach developed by Schulz-Gasch et al. (J. Chem. Inf. Model, 52, 1499, 2012). The RDKit implementation allows the user to customize the torsion fingerprints:

- In the original approach, the torsions are weighted based on their distance to the center of the molecule. By default, this weighting is performed, but can be turned off using the flag useWeights=False
- The similarity between atoms is determined by comparing the hash codes from the Morgan algorithm at a given radius (default: radius = 2).
- In the original approach, the maximal deviation used for normalization is 180.0 degrees for all torsions (default). If maxDev='spec', a torsion-type dependent maximal deviation is used for the normalization.

Examples of using it:

```
from rdkit import Chem, RDConfig
from rdkit.Chem import AllChem, TorsionFingerprints
ref = Chem.MolFromSmiles('NC(=[NH2+])clcc(C[C@@H](NC(=0)CNS(=0)(=0)c2ccc3cccc3c2)C(=0)N2CCCCC2)ccl
mol1 = Chem.MolFromPDBFile(RDConfig.RDBaseDir+'/rdkit/Chem/test_data/1DWD_ligand.pdb')
mol1 = AllChem.AssignBondOrdersFromTemplate(ref, mol1)
mol2 = Chem.MolFromPDBFile(RDConfig.RDBaseDir+'/rdkit/Chem/test_data/1PPC_ligand.pdb')
mol2 = AllChem.AssignBondOrdersFromTemplate(ref, mol2)
tfd1 = TorsionFingerprints.GetTFDBetweenMolecules(mol1, mol2)
tfd2 = TorsionFingerprints.GetTFDBetweenMolecules(mol1, mol2, useWeights=False)
tfd3 = TorsionFingerprints.GetTFDBetweenMolecules(mol1, mol2, maxDev='spec')
print tfd1, tfd2, tfd3

This produces:
[0.0645...] [0.168...] [0.0675...]
```

If the TFD between conformers of the same molecule is to be calculated, the function GetTFDBetweenConformers() should be used for performance reasons.

Examples of using it:

```
from rdkit import Chem, RDConfig
from rdkit.Chem import AllChem, TorsionFingerprints
ref = Chem.MolFromSmiles('NC(=[NH2+])clccc(C[C@@H](NC(=O)CNS(=O)(=O)c2ccc3cccc3c2)C(=O)N2CCCC2)ccl
mol1 = Chem.MolFromPDBFile(RDConfig.RDBaseDir+'/rdkit/Chem/test_data/1DWD_ligand.pdb')
mol1 = AllChem.AssignBondOrdersFromTemplate(ref, mol1)
mol2 = Chem.MolFromPDBFile(RDConfig.RDBaseDir+'/rdkit/Chem/test_data/1PPC_ligand.pdb')
mol1.AddConformer(mol2.GetConformer(), assignId=True)
tfd = TorsionFingerprints.GetTFDBetweenConformers(mol1, confIds1=[0], confIds2=[1])
print tfd
This produces:
```

[0.0645...]

For the conformer RMS and TFD values, the RDKit provides convenience functions that calculated directly the symmetric matrix which can be fed into a clustering algorithm such as Butina clustering. The flag reordering ensures that the number of neighbors of the unclustered molecules is updated every time a cluster is created.

Examples of using it:

```
from rdkit import Chem
from rdkit.Chem import AllChem, TorsionFingerprints
from rdkit.ML.Cluster import Butina
mol = Chem.MolFromSmiles('NC(=[NH2+])clccc(C[C@@H](NC(=0)CNS(=0)(=0)c2ccc3ccccc3c2)C(=0)N2CCCCC2)ccl
cids = AllChem.EmbedMultipleConfs(mol, numConfs=50, maxAttempts=1000, pruneRmsThresh=0.1)
# RMS matrix
rmsmat = AllChem.GetConformerRMSMatrix(mol, prealigned=False)
# TFD matrix
tfdmat = TorsionFingerprints.GetTFDMatrix(mol)
# clustering
num = mol.GetNumConformers()
rms_clusters = Butina.ClusterData(rmsmat, num, 2.0, isDistData=True, reordering=True)
tfd_clusters = Butina.ClusterData(tfdmat, num, 0.3, isDistData=True, reordering=True)
```

5.3.5 Using scikit-learn with RDKit

scikit-learn is a machine-learning library for Python containing a variety of supervised and unsupervised methods. The documention can be found here: http://scikit-learn.org/stable/user_guide.html

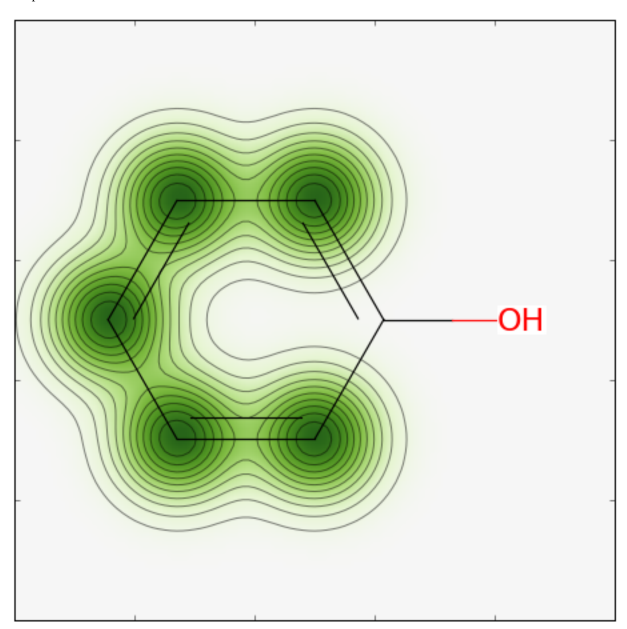
RDKit fingerprints can be used to train machine-learning models from scikit-learn. Here is an example for random forest:

The code:

```
from rdkit import Chem, DataStructs
from rdkit.Chem import AllChem
from sklearn.ensemble import RandomForestClassifier
import numpy
# generate four molecules
m1 = Chem.MolFromSmiles('clcccc1')
m2 = Chem.MolFromSmiles('c1cccc1CC')
m3 = Chem.MolFromSmiles('clccncc1')
m4 = Chem.MolFromSmiles('clccncc1CC')
mols = [m1, m2, m3, m4]
# generate fingeprints: Morgan fingerprint with radius 2
fps = [AllChem.GetMorganFingerprintAsBitVect(m, 2) for m in mols]
# convert the RDKit explicit vectors into numpy arrays
np\_fps = []
for fp in fps:
  arr = numpy.zeros((1,))
  DataStructs.ConvertToNumpyArray(fp, arr)
  np_fps.append(arr)
# get a random forest classifiert with 100 trees
rf = RandomForestClassifier(n_estimators=100, random_state=1123)
# train the random forest
# with the first two molecules being actives (class 1) and
# the last two being inactives (class 0)
ys_fit = [1, 1, 0, 0]
rf.fit(np_fps, ys_fit)
# use the random forest to predict a new molecule
m5 = Chem.MolFromSmiles('clcccc10')
fp = numpy.zeros((1,))
DataStructs.ConvertToNumpyArray(AllChem.GetMorganFingerprintAsBitVect(m5, 2), fp)
print rf.predict(fp)
print rf.predict_proba(fp)
The output with scikit-learn version 0.13 is:
    [1]
    [[ 0.14 0.86]]
Generating a similarity map for this model.
The code:
from rdkit.Chem.Draw import SimilarityMaps
# helper function
def getProba(fp, predictionFunction):
  return predictionFunction(fp)[0][1]
```

```
m5 = Chem.MolFromSmiles('clcccc10')
fig, maxweight = SimilarityMaps.GetSimilarityMapForModel(m5, SimilarityMaps.GetMorganFingerprint, land
```

This produces:



5.3.6 Using custom MCS atom types

Mailing list discussion: http://www.mail-archive.com/rdkit-discuss@lists.sourceforge.net/msg03676.html

IPython notebook: http://nbviewer.ipython.org/gist/greglandrum/8351725 https://gist.github.com/greglandrum/8351725

The goal is to be able to use custom atom types in the MCS code, yet still be able to get a readable SMILES for the MCS. We will use the MCS code's option to use isotope information in the matching and then set bogus isotope values that contain our isotope information.

The code:

```
from rdkit import Chem
from rdkit.Chem import rdFMCS
# our test molecules:
\verb|smis=["COcloce(C(Nc2nc3c(ncn3COCC=0)c(=0)[nH]2)(c2cccc2)c2cccc2)cc1",
      "COclccc(C(Nc2nc3c(ncn3COC(CO)(CO)CO)c(=0)[nH]2)(c2cccc2)c2cccc2)cc1"]
ms = [Chem.MolFromSmiles(x) for x in smis]
def label(a):
  " a simple hash combining atom number and hybridization "
 return 100*int(a.GetHybridization())+a.GetAtomicNum()
# copy the molecules, since we will be changing them
nms = [Chem.Mol(x) for x in ms]
for nm in nms:
 for at in nm.GetAtoms():
      at.SetIsotope(label(at))
mcs=rdFMCS.FindMCS(nms,atomCompare=rdFMCS.AtomCompare.CompareIsotopes)
print mcs.smartsString
```

This generates the following output:

```
 [406*] - [308*] - [306*] 1 : [306*] : [306*] : [306*] : [306*] : [306*] : [306*] : [306*] : [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*] - [307*]
```

That's what we asked for, but it's not exactly readable. We can get to a more readable form in a two step process:

- 1. Do a substructure match of the MCS onto a copied molecule
- 2. Generate SMILES for the original molecule, using only the atoms that matched in the copy.

This works because we know that the atom indices in the copies and the original molecules are the same.

That's what we were looking for.

5.3.7 Clustering molecules

For large sets of molecules (more than 1000-2000), it's most efficient to use the Butina clustering algorithm.

Here's some code for doing that for a set of fingerprints:

```
def ClusterFps(fps,cutoff=0.2):
    from rdkit import DataStructs
    from rdkit.ML.Cluster import Butina

# first generate the distance matrix:
    dists = []
```

```
nfps = len(fps)
for i in range(1,nfps):
    sims = DataStructs.BulkTanimotoSimilarity(fps[i],fps[:i])
    dists.extend([1-x for x in sims])

# now cluster the data:
cs = Butina.ClusterData(dists,nfps,cutoff,isDistData=True)
return cs
```

The return value is a tuple of clusters, where each cluster is a tuple of ids.

Example usage:

```
from rdkit import Chem
from rdkit.Chem import AllChem
import gzip
ms = [x for x in Chem.ForwardSDMolSupplier(gzip.open('zdd.sdf.gz')) if x is not None]
fps = [AllChem.GetMorganFingerprintAsBitVect(x,2,1024) for x in ms]
clusters=ClusterFps(fps,cutoff=0.4)
```

The variable *clusters* contains the results:

```
>>> print clusters[200] (6164, 1400, 1403, 1537, 1543, 6575, 6759)
```

That cluster contains 7 points, the centroid is point 6164.

5.3.8 RMSD Calculation between N molecules

Introduction

We sometimes need to calculate RMSD distances between two (or more) molecules. This can be used to calculate how close two conformers are. Most RMSD calculations make sense only on similar compounds or, at least, for common parts in different compounds.

Details

The following program (written in python 2.7) takes an SDF file as an input and generates all the RMSD distances between the molecules in that file. These distances are written to an output file (user defined).

So for an SDF with 5 conformers we will get 10 RMSD scores - typical n choose k problem, without repetition i.e. 5! / 2!(5-2)!

```
#!/usr/bin/python
,,,
calculates RMSD differences between all structures in a file
@author: JP <jp@javaclass.co.uk>
,,,
import os
import getopt
import sys

# rdkit imports
from rdkit import Chem
from rdkit.Chem import AllChem
```

```
Write contents of a string to file
def write_contents(filename, contents):
 # do some basic checking, could use assert strictly speaking
 assert filename is not None, "filename cannot be None"
 assert contents is not None, "contents cannot be None"
 f = open(filename, "w")
 f.write(contents)
 f.close() # close the file
Write a list to a file
def write_list_to_file(filename, list, line_sep = os.linesep):
 # do some basic checking, could use assert strictly speaking
 assert list is not None and len(list) > 0, "list cannot be None or empty"
 write_contents(filename, line_sep.join(list))
Calculate RMSD spread
def calculate_spread(molecules_file):
 assert os.path.isfile(molecules_file), "File %s does not exist!" % molecules
  # get an iterator
 mols = Chem.SDMolSupplier(molecules_file)
 spread_values = []
  # how many molecules do we have in our file
 mol_count = len(mols)
  # we are going to compare each molecule with every other molecule
  # typical n choose k scenario (n choose 2)
  # where number of combinations is given by (n!) / k! (n-k)!; if my maths isn't too rusty
 for i in range(mol_count - 1):
      for j in range(i+1, mol_count):
          # show something is being done ... because for large mol_count this will take some time
          print "Aligning molecule #%d with molecule #%d (%d molecules in all)" % (i, j, mol_count)
          # calculate RMSD and store in an array
          # unlike AlignMol this takes care of symettry
          spread_values.append(str(AllChem.GetBestRMS(mols[i], mols[j])))
  # return that array
 return spread_values
def main():
 try:
      # the options are as follows:
      # f - the actual structure file
      opts, args = getopt.getopt(sys.argv[1:], "vf:o:")
 except getopt.GetoptError, err:
      # print help information and exit:
     print str(err) # will print something like "option -a not recognized"
     sys.exit(401)
  # DEFAULTS
 molecules_file = None
```

```
output_file = None
for opt, arg in opts:
    if opt == "-v":
        print "RMSD Spread 1.1"
        sys.exit()
    elif opt == "-f":
       molecules_file = arg
    elif opt == "-o":
       output_file = arg
    else:
        assert False, "Unhandled option: " + opt
# assert the following - not the cleanest way to do this but this will work
assert molecules_file is not None, "file containing molecules must be specified, add -f to command
assert output_file is not None, "output file must be specified, add -o to command line arguments"
# get the RMSD spread values
spread_values = calculate_spread(molecules_file)
# write them to file
write_list_to_file(output_file, spread_values)
```

This program may be executed at the command line in the following manner (provided you have your python interpreter at /usr/bin/python, otherwise edit the first line; the funnily named shebang):

```
calculate_spread.py -f my_conformers.sdf -o my_conformers.rmsd_spread.txt
```

TL;DR: The line AllChem.GetBestRMS(mol1, mol2) returns the RMSD as a float and is the gist of this program. GetBestRMS() takes care of symettry unlike AlignMol()

5.4 License

main()

if __name__ == "__main__":

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The intent of this license is similar to that of the RDKit itself. In simple words: "Do whatever you want with it, but please give us some credit."

CHAPTER

SIX

THE RDKIT DATABASE CARTRIDGE

6.1 What is this?

This document is a tutorial and reference guide for the RDKit PostgreSQL cartridge.

If you find mistakes, or have suggestions for improvements, please either fix them yourselves in the source document (the .rst file) or send them to the mailing list: rdkit-discuss@lists.sourceforge.net (you will need to subscribe first)

6.2 Tutorial

6.2.1 Introduction

6.2.2 Creating databases

Configuration

The timing information below was collected on a commodity desktop PC (Dell Studio XPS with a 2.9GHz i7 CPU and 8GB of RAM) running Ubuntu 12.04 and using PostgreSQL v9.1.4. The database was installed with default parameters.

To improve performance while loading the database and building the index, I changed a couple of postgres configuration settings in *postgresql.conf*

And to improve search performance, I allowed postgresql to use more memory than the extremely conservative default settings:

Creating a database from a file

In this example I show how to load a database from the SMILES file of commercially available compounds that is downloadable from emolecules.com at URL http://www.emolecules.com/doc/plus/download-database.php

If you choose to repeat this exact example yourself, please note that it takes several hours to load the 6 million row database and generate all fingerprints.

First create the database and install the cartridge:

```
~/RDKit_trunk/Data/emolecules > createdb emolecules  
~/RDKit_trunk/Data/emolecules > psql -c 'create extension rdkit' emolecules
```

Now create and populate a table holding the raw data:

```
~/RDKit_trunk/Data/emolecules > psql -c 'create table raw_data (id SERIAL, smiles text, emol_id international content of the sequence "raw_data_id_seq" for serial column "raw_data_id CREATE TABLE  
~/RDKit_trunk/Data/emolecules > zcat emolecules-2013-02-01.smi.gz | sed 'ld; s/\\\\/g' | psql -c
```

Create the molecule table, but only for SMILES that the RDKit accepts:

```
~/RDKit_trunk/Data/emolecules > psql emolecules
psql (9.1.4)
Type "help" for help.
emolecules=# select * into mols from (select id,mol_from_smiles(smiles::cstring) m from raw_data) tmp
WARNING: could not create molecule from SMILES 'CN(C)C(=[N+](C)C)Cl.F[P-](F)(F)(F)(F)F'
... a lot of warnings deleted ...
SELECT 6008732
emolecules=# create index molidx on mols using gist(m);
CREATE INDEX
```

The last step is only required if you plan to do substructure searches.

Loading ChEMBL

Start by downloading and installing the postgresql dump from the ChEMBL website ftp://ftp.ebi.ac.uk/pub/databases/chembl/ChEMBLdb/latest

Connect to the database, install the cartridge, and create the schema that we'll use:

```
chembl_14=# create extension if not exists rdkit;
chembl_14=# create schema rdk;
```

Create the molecules and build the substructure search index:

```
chembl_14=# select * into rdk.mols from (select molregno,mol_from_ctab(molfile::cstring) m from composed select 1210823
chembl_14=# create index molidx on rdk.mols using gist(m);
CREATE INDEX
chembl_14=# alter table rdk.mols add primary key (molregno);
NOTICE: ALTER TABLE / ADD PRIMARY KEY will create implicit index "mols_pkey" for table "mols"
ALTER TABLE
```

Create some fingerprints and build the similarity search index:

```
chembl_14=# select molregno,torsionbv_fp(m) as torsionbv,morganbv_fp(m) as mfp2,featmorganbv_fp(m) as
SELECT 1210823
chembl_14=# create index fps_ttbv_idx on rdk.fps using gist(torsionbv);
CREATE INDEX
chembl_14=# create index fps_mfp2_idx on rdk.fps using gist(mfp2);
CREATE INDEX
chembl_14=# create index fps_ffp2_idx on rdk.fps using gist(ffp2);
CREATE INDEX
chembl_14=# create index fps_ffp2_idx on rdk.fps using gist(ffp2);
CREATE INDEX
chembl_14=# alter table rdk.fps add primary key (molregno);
NOTICE: ALTER TABLE / ADD PRIMARY KEY will create implicit index "fps_pkey" for table "fps"
ALTER TABLE
```

6.2.3 Substructure searches

Example query molecules taken from the eMolecules home page:

```
chembl_14=# select count(*) from rdk.mols where m@>'clccc2c1nncc2';
count
   2.81
(1 row)
Time: 184.043 ms
chembl_14=# select count(*) from rdk.mols where m@>'clccnc2clnccn2';
count
   671
(1 row)
Time: 449.998 ms
chembl_14=# select count(*) from rdk.mols where m@>'clcncc2n1ccn2';
count
   930
(1 row)
Time: 568.378 ms
chembl_14=# select count(*) from rdk.mols where m@>'Nc1ncnc(N)n1';
count
  4478
(1 row)
Time: 721.758 ms
chembl_14=# select count(*) from rdk.mols where m@>'clscnn1';
count
10908
(1 row)
Time: 701.036 ms
chembl_14=# select count(*) from rdk.mols where m@>'clccc2clncs2';
count
12823
(1 row)
Time: 1585.473 ms
chembl_14=# select count(*) from rdk.mols where m@>'c1ccc2c1CNCCN2';
count
 1155
(1 row)
Time: 4567.222 ms
```

Notice that the last two queries are starting to take a while to execute and count all the results.

Given we're searching through 1.2 million compounds these search times aren't incredibly slow, but it would be nice to have them quicker.

One easy way to speed things up, particularly for queries that return a large number of results, is to only retrieve a

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limited number of results:

SMARTS-based queries

Oxadiazole or thiadiazole:

This is slower than the pure SMILES query, this is generally true of SMARTS-based queries.

Using Stereochemistry

Note that by default stereochemistry is not taken into account when doing substructure queries:

```
Time: 9.447 ms
```

This can be changed using the *rdkit.do_chiral_sss* configuration variable:

```
chembl_14=# set rdkit.do_chiral_sss=true;
SET
Time: 0.241 ms
chembl_14=# select * from rdk.mols where m@>'NC(=O)[C@@H]1CCCN1C=O' limit 10;
     1295889 | COclccc(C[C@0H](C(=O)NCC(N)=O)N(C)C(=O)[C@0H]2CCCN2C(=O)[C@H](CC(C)C)NC(=O)C(C)NC(=O)OCC
     1293021 | CCC(C)C1NC(=0)C(NC(=0)C(CC(C)C)N(C)C(=0)[C@H]2CC(0)CN2C(=0)[C@H](C)O)C(C)OC(=0)[C@H](Cc:
     1293647 | CCC(C) [C@@H]1NC(=O) [C@@H]2CCCN2C(=O)C(CC(O)CC1)OC(=O)CCNC(=O)[C@H](C)N(C)C(=O)[C@H](C)C(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O)CCNC(=O
     1290320 | C=CCOC(=0) [C@@H]1C[C@@H](OC(C)(C)C)CN1C(=0) [C@@H]1[C@H]2OC(C)(C)O[C@H]2CN1C(=0)OCC1c2ccc
     1281392 | COC1=CC2C(=O)N(C)[C@@H](C)C(=O)N3NCCC[C@@H]3C(=O)N3[C@@H](C[C@@]4(O)c5ccc(Cl)cc5N[C@@H]3-
     1007418 | C/C=C\C=C\C(=0)N1CC2(CC(c3cccc(NC(=0)/C=C\C=C/C)c3)=N02)C[C@H]1C(N)=0
       785530 \mid C/C=C/C (=0) N1CC2 (CC (c3cccc (NC (=0) CC) c3) = NO2) C [C@H] 1C (N) = 0
     1292152 | CCCCCCCC(=O)N[C@H](C(=O)N[C@H](C(=O)N(C)[C@H](C(=O)N1CCC[C@H]1C(=O)N(C)[C@H](C)C(=O)NCc1
     1281390 | CC(C) [C@@H]1NC(=0) [C@@H]2C[C@@]3(O)c4ccc(Cl)cc4N[C@H]3N2C(=0) [C@H]2CCCNN2C(=0) [C@@H](C)N
     1057962 | CC[C@H](C)[C@@H]1NC(=0)[C@H](CCCNC(=N)N)NC(=0)[C@H](CC(=0)0)NC(=0)[C@H](CCSC)NC(=0)[C@H]
(10 rows)
Time: 35.383 ms
```

6.2.4 Similarity searches

Basic similarity searching:

Time: 826.886 ms

Usually we'd like to find a sorted listed of neighbors along with the accompanying SMILES. This SQL function makes that pattern easy:

```
chembl_14=# create or replace function get_mfp2_neighbors(smiles text)
   returns table (molregno integer, m mol, similarity double precision) as
 select molregno, m, tanimoto_sml (morganbv_fp (mol_from_smiles ($1::cstring)), mfp2) as similarity
 from rdk.fps join rdk.mols using (molregno)
 where morganbv_fp(mol_from_smiles($1::cstring))%mfp2
 order by morganbv_fp(mol_from_smiles($1::cstring)) <%>mfp2;
 $$ language sql stable ;
CREATE FUNCTION
Time: 0.856 ms
chembl_14=#
molregno |
                                                         | similarity
  472512 | Cc1ccc2nc(-c3ccc(NC(=0)C4CCN(C(=0)c5cccs5)CC4)cc3)sc2c1
                                                          | 0.772727272727273
  471317 | Cc1ccc2nc(-c3ccc(NC(=0)C4CCCN(S(=0)(=0)c5cccs5)C4)cc3)sc2c1 | 0.657534246575342
  471461 | Cc1ccc2nc(-c3ccc(NC(=0)C4CCN(S(=0)(=0)c5cccs5)CC4)cc3)sc2c1 | 0.647887323943662
```

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```
1032469 | O=C(Nc1nc2ccc(C1)cc2s1)[C@@H]1CCCN1C(=O)c1cccs1
                                                       | 0.623188405797101
  751668 | COclccc2nc(NC(=0)[C@@H]3CCCN3C(=0)c3cccs3)sc2c1
                                                       | 0.619718309859155
  740754 | Cc1ccc (NC (=0) C2CCCN2C (=0) c2cccs2) cc1C
                                                       | 0.606060606060606
                                                       | 0.602941176470588
  732905 | O=C (Nc1ccc(S(=0)(=0)N2CCCC2)cc1)C1CCCN1C(=0)c1cccs1
 1087495 | Cc1ccc(NC(=0)C2CCCN2C(=0)c2cccs2)c(C)c1
                                                       | 0.597014925373134
(10 rows)
Time: 5453.200 ms
chembl_14=# select * from get_mfp2_neighbors('Cc1ccc2nc(N(C)CC(=0)0)sc2c1') limit 10;
molregno |
                      m | similarity
  412312 | Cc1ccc2nc(N(C)CCN(C)c3nc4ccc(C)cc4s3)sc2c1
                                                  | 0.692307692307692
  470082 | CN(CC(=0)0)clnc2cc([N+](=0)[0-])ccc2s1
                                                   | 0.583333333333333
 1040255 | CC (=0) N (CCCN (C) C) clnc2ccc (C) cc2s1
                                                   | 0.571428571428571
  773946 | Cl.CC(=0)N(CCCN(C)C)clnc2ccc(C)cc2s1
                                                   | 0.549019607843137
                                                  | 0.518518518518518
 1044892 | Cclccc2nc(N(CCN(C)C)C(=0)c3cc(Cl)sc3Cl)sc2c1
 1040496 | Cc1ccc2nc(N(CCCN(C)C)C(=0)CCc3ccccc3)sc2c1
                                                  | 0.517857142857143
 1049393 | Cc1ccc2nc(N(CCCN(C)C)C(=0)CS(=0)(=0)c3ccccc3)sc2c1 | 0.517857142857143
                                                  | 0.510204081632653
 441378 | Cc1ccc2nc(NC(=0)CCC(=0)0)sc2c1
 (10 rows)
Time: 1797.656 ms
```

Adjusting the similarity cutoff

By default, the minimum similarity returned with a similarity search is 0.5. This can be adjusted with the *rd-kit.tanimoto_threshold* (and *rdkit.dice_threshold*) configuration variables:

```
chembl_14=# select count(*) from get_mfp2_neighbors('Cc1ccc2nc(N(C)CC(=0)O)sc2c1');
count.
   18
(1 row)
Time: 1199.751 ms
chembl_14=# set rdkit.tanimoto_threshold=0.7;
SET
Time: 0.191 ms
chembl_14=# select count(*) from get_mfp2_neighbors('Cc1ccc2nc(N(C)CC(=0)O)sc2c1');
   Ω
(1 row)
Time: 826.058 ms
chembl_14=# set rdkit.tanimoto_threshold=0.6;
Time: 0.220 ms
chembl_14=# select count(*) from get_mfp2_neighbors('Cc1ccc2nc(N(C)CC(=0)O)sc2c1');
count
    1
(1 row)
```

6.2.5 Using the MCS code

The most straightforward use of the MCS code is to find the maximum common substructure of a group of molecules:

The same thing can be done with a SMILES column:

It's also possible to adjust some of the parameters to the FMCS algorithm, though this is somewhat more painful as of this writing (the 2015_03_1 release). Here are a couple of examples:

Note The combination of "AtomCompare": "Any" and a value of "Threshold" that is less than 1.0 does a quite generic search and can results in very long search times. Using "Timeout" with this combination is recommended:

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[#8]=[#6](-[#7]-[#6]1:[#6]:[#6]:[#6](:[#6]:[#6]:1)-[#6](=[#8])-[#7]1-[#6]-[#6]-[#6]-[#6]-[#6]-[#6]2:[#6]1:[#6]0:[#

Available parameters and their default values are:

- MaximizeBonds (true)
- Threshold (1.0)
- Timeout (-1, no timeout)
- MatchValences (false)
- MatchChiralTag (false) Applies to atoms
- RingMatchesRingOnly (false)
- CompleteRingsOnly (false)
- MatchStereo (false) Applies to bonds
- AtomCompare ("Elements") can be "Elements", "Isotopes", or "Any"
- BondCompare ("Order") can be "Order", "OrderExact", or "Any"

6.3 Reference Guide

6.3.1 New Types

- *mol*: an rdkit molecule. Can be created from a SMILES via direct type conversion, for example: 'c1ccccc1'::mol creates a molecule from the SMILES 'c1cccc1'
- *qmol*: an rdkit molecule containing query features (i.e. constructed from SMARTS). Can be created from a SMARTS via direct type conversion, for example: 'clcccc[c,n]l'::qmol creates a query molecule from the SMARTS 'clcccc[c,n]l'
- sfp: a sparse count vector fingerprint (SparseIntVect in C++ and Python)
- *bfp* : a bit vector fingerprint (*ExplicitBitVect* in C++ and Python)

6.3.2 Parameters

- rdkit.tanimoto_threshold: threshold value for the Tanimoto similarity operator. Searches done using Tanimoto similarity will only return results with a similarity of at least this value.
- rdkit.dice_threshold: threshold value for the Dice similarity operator. Searches done using Dice similarity will only return results with a similarity of at least this value.
- rdkit.do_chiral_sss: toggles whether or not stereochemistry is used in substructure matching. (available from 2013_03 release).
- rdkit.sss_fp_size: the size (in bits) of the fingerprint used for substructure screening.
- rdkit.morgan_fp_size : the size (in bits) of morgan fingerprints

- rdkit.featmorgan_fp_size : the size (in bits) of featmorgan fingerprints
- rdkit.layered_fp_size : the size (in bits) of layered fingerprints
- rdkit.rdkit_fp_size: the size (in bits) of RDKit fingerprints
- rdkit.torsion_fp_size : the size (in bits) of topological torsion bit vector fingerprints
- rdkit.atompair_fp_size : the size (in bits) of atom pair bit vector fingerprints
- rdkit.avalon fp size: the size (in bits) of avalon fingerprints

6.3.3 Operators

Similarity search

- %: operator used for similarity searches using Tanimoto similarity. Returns whether or not the Tanimoto similarity between two fingerprints (either two sfp or two bfp values) exceeds rdkit.tanimoto_threshold.
- #: operator used for similarity searches using Dice similarity. Returns whether or not the Dice similarity between two fingerprints (either two sfp or two bfp values) exceeds rdkit.dice_threshold.
- <%>: used for Tanimoto KNN searches (to return ordered lists of neighbors).
- <#>: used for Dice KNN searches (to return ordered lists of neighbors).

Substructure and exact structure search

- @>: substructure search operator. Returns whether or not the *mol* or *qmol* on the right is a substructure of the *mol* on the left.
- <@: substructure search operator. Returns whether or not the *mol* or *qmol* on the left is a substructure of the *mol* on the right.
- @=: returns whether or not two molecules are the same.

Molecule comparison

- < : returns whether or not the left *mol* is less than the right *mol*
- > : returns whether or not the left *mol* is greater than the right *mol*
- = : returns whether or not the left mol is equal to the right mol
- <= : returns whether or not the left *mol* is less than or equal to the right *mol*
- >= : returns whether or not the left *mol* is greater than or equal to the right *mol*

Note Two molecules are compared by making the following comparisons in order. Later comparisons are only made if the preceding values are equal:

Number of atoms # Number of bonds # Molecular weight # Number of rings

If all of the above are the same and the second molecule is a substructure of the first, the molecules are declared equal, Otherwise (should not happen) the first molecule is arbitrarily defined to be less than the second.

There are additional operators defined in the cartridge, but these are used for internal purposes.

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6.3.4 Functions

Fingerprint Related

Generating fingerprints

- morgan_fp(mol,int default 2): returns an sfp which is the count-based Morgan fingerprint for a molecule using connectivity invariants. The second argument provides the radius. This is an ECFP-like fingerprint.
- morganbv_fp(mol,int default 2): returns a bfp which is the bit vector Morgan fingerprint for a molecule using connectivity invariants. The second argument provides the radius. This is an ECFP-like fingerprint.
- featmorgan_fp(mol,int default 2): returns an sfp which is the count-based Morgan fingerprint for a molecule using chemical-feature invariants. The second argument provides the radius. This is an FCFP-like fingerprint.
- featmorganbv_fp(mol,int default 2): returns a bfp which is the bit vector Morgan fingerprint for a molecule using chemical-feature invariants. The second argument provides the radius. This is an FCFP-like fingerprint.
- rdkit_fp(mol): returns a bfp which is the RDKit fingerprint for a molecule. This is a daylight-fingerprint using hashed molecular subgraphs.
- atompair_fp(mol): returns an sfp which is the count-based atom-pair fingerprint for a molecule.
- atompairbv_fp(mol): returns a bfp which is the bit vector atom-pair fingerprint for a molecule.
- torsion_fp(mol): returns an sfp which is the count-based topological-torsion fingerprint for a molecule.
- torsionby_fp(mol): returns a bfp which is the bit vector topological-torsion fingerprint for a molecule.
- *layered_fp(mol)*: returns a *bfp* which is the layered fingerprint for a molecule. This is an experimental substructure fingerprint using hashed molecular subgraphs.
- maccs_fp(mol): returns a bfp which is the MACCS fingerprint for a molecule (available from 2013_01 release).

Working with fingerprints

- *tanimoto_sml(fp,fp)*: returns the Tanimoto similarity between two fingerprints of the same type (either two *sfp* or two *bfp* values).
- dice_sml(fp,fp): returns the Dice similarity between two fingerprints of the same type (either two sfp or two bfp values).
- *size(bfp)*: returns the length of (number of bits in) a *bfp*.
- add(sfp,sfp): returns an sfp formed by the element-wise addition of the two sfp arguments.
- subtract(sfp,sfp): returns an sfp formed by the element-wise subtraction of the two sfp arguments.
- *all_values_lt(sfp,int)*: returns a boolean indicating whether or not all elements of the *sfp* argument are less than the *int* argument.
- *all_values_gt(sfp,int)*: returns a boolean indicating whether or not all elements of the *sfp* argument are greater than the *int* argument.

Fingerprint I/O

• bfp_to_binary_text(bfp): returns a bytea with the binary string representation of the fingerprint that can be converted back into an RDKit fingerprint in other software. (available from Q3 2012 (2012_09) release)

• bfp_from_binary_text(bytea): constructs a bfp from a binary string representation of the fingerprint. (available from Q3 2012 (2012_09) release)

Molecule Related

Molecule I/O and Validation

- is_valid_smiles(smiles): returns whether or not a SMILES string produces a valid RDKit molecule.
- is_valid_ctab(ctab): returns whether or not a CTAB (mol block) string produces a valid RDKit molecule.
- is valid smarts(smarts): returns whether or not a SMARTS string produces a valid RDKit molecule.
- is_valid_mol_pkl(bytea): returns whether or not a binary string (bytea) can be converted into an RDKit molecule. (available from Q3 2012 (2012_09) release)
- mol_from_smiles(smiles): returns a molecule for a SMILES string, NULL if the molecule construction fails.
- mol_from_smarts(smarts): returns a molecule for a SMARTS string, NULL if the molecule construction fails.
- mol_from_ctab(ctab, bool default false): returns a molecule for a CTAB (mol block) string, NULL if the molecule construction fails. The optional second argument controls whether or not the molecule's coordinates are saved.
- mol_from_pkl(bytea): returns a molecule for a binary string (bytea), NULL if the molecule construction fails. (available from Q3 2012 (2012 09) release)
- *qmol_from_smiles(smiles)*: returns a query molecule for a SMILES string, NULL if the molecule construction fails. Explicit Hs in the SMILES are converted into query features on the attached atom.
- *qmol_from_ctab(ctab, bool default false)*: returns a query molecule for a CTAB (mol block) string, NULL if the molecule construction fails. Explicit Hs in the SMILES are converted into query features on the attached atom. The optional second argument controls whether or not the molecule's coordinates are saved.
- *mol_to_smiles(mol)* : returns the canonical SMILES for a molecule.
- mol_to_smarts(mol): returns SMARTS string for a molecule.
- mol_to_pkl(mol): returns binary string (bytea) for a molecule. (available from Q3 2012 (2012_09) release)
- mol_to_ctab(mol,bool default true): returns a CTAB (mol block) string for a molecule. The optional second argument controls whether or not 2D coordinates will be generated for molecules that don't have coordinates. (available from the 2014_03 release)

Substructure operations

- substruct(mol,mol): returns whether or not the second mol is a substructure of the first.
- *substruct_count(mol,mol,bool default true)*: returns the number of substructure matches between the second molecule and the first. The third argument toggles whether or not the matches are uniquified. (*available from 2013_03 release*)

Descriptors

- mol_amw(mol): returns the AMW for a molecule.
- $mol_logp(mol)$: returns the MolLogP for a molecule.

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- mol_tpsa(mol): returns the topological polar surface area for a molecule (available from Q1 2011 (2011_03) release).
- mol_fractioncsp3(mol): returns the fraction of carbons that are sp3 hybridized (available from 2013_03 release).
- mol_hba(mol): returns the number of Lipinski H-bond acceptors (i.e. number of Os and Ns) for a molecule.
- mol_hbd(mol): returns the number of Lipinski H-bond donors (i.e. number of Os and Ns that have at least one H) for a molecule.
- mol numatoms(mol): returns the total number of atoms in a molecule.
- mol_numheavyatoms(mol): returns the number of heavy atoms in a molecule.
- mol_numrotatablebonds(mol): returns the number of rotatable bonds in a molecule (available from Q1 2011 (2011_03) release).
- mol_numheteroatoms(mol): returns the number of heteroatoms in a molecule (available from Q1 2011 (2011_03) release).
- mol_numrings(mol): returns the number of rings in a molecule (available from Q1 2011 (2011_03) release).
- mol_numaromaticrings(mol): returns the number of aromatic rings in a molecule (available from 2013_03 release).
- mol_numaliphaticrings(mol): returns the number of aliphatic (at least one non-aromatic bond) rings in a molecule (available from 2013_03 release).
- mol_numsaturatedrings(mol): returns the number of saturated rings in a molecule (available from 2013_03 release).
- mol_numaromaticheterocycles(mol): returns the number of aromatic heterocycles in a molecule (available from 2013_03 release).
- mol_numaliphaticheterocycles(mol): returns the number of aliphatic (at least one non-aromatic bond) heterocycles in a molecule (available from 2013_03 release).
- mol_numsaturatedheterocycles(mol): returns the number of saturated heterocycles in a molecule (available from 2013_03 release).
- mol_numaromaticcarbocycles(mol): returns the number of aromatic carbocycles in a molecule (available from 2013_03 release).
- mol_numaliphaticcarbocycles(mol): returns the number of aliphatic (at least one non-aromatic bond) carbocycles in a molecule (available from 2013_03 release).
- mol_numsaturatedcarbocycles(mol): returns the number of saturated carbocycles in a molecule (available from 2013 03 release).
- mol_inchi(mol): returns an InChI for the molecule. (available from the 2011_06 release, requires that the RDKit be built with InChI support).
- mol_inchikey(mol): returns an InChI key for the molecule. (available from the 2011_06 release, requires that the RDKit be built with InChI support).
- mol_formula(mol,bool default false, bool default true): returns a string with the molecular formula. The second argument controls whether isotope information is included in the formula; the third argument controls whether "D" and "T" are used instead of [2H] and [3H].

(available from the 2014_03 release)

Connectivity Descriptors

- mol_chi0v(mol) mol_chi4v(mol): returns the ChiXv value for a molecule for X=0-4 (available from 2012_01 release).
- mol_chi0n(mol) mol_chi4n(mol): returns the ChiXn value for a molecule for X=0-4 (available from 2012_01 release).
- mol_kappa1(mol) mol_kappa3(mol): returns the kappaX value for a molecule for X=1-3 (available from 2012_01 release).

MCS

- fmcs(mols): an aggregation function that calculates the MCS for a set of molecules
- fmcs_smiles(text, json default ''): calculates the MCS for a space-separated set of SMILES. The optional json argument is used to provide parameters to the MCS code.

Other

• rdkit_version(): returns a string with the cartridge version number.

There are additional functions defined in the cartridge, but these are used for internal purposes.

6.4 Using the Cartridge from Python

The recommended adapter for connecting to postgresql is pyscopg2 (https://pypi.python.org/pypi/psycopg2).

Here's an example of connecting to our local copy of ChEMBL and doing a basic substructure search:

```
>>> import psycopg2
>>> conn = psycopg2.connect(database='chemb1_16')
>>> curs = conn.cursor()
>>> curs.execute('select * from rdk.mols where m@>%s',('clcccc2clnncc2',))
>>> curs.fetchone()
(9830, 'CC(C)Sclccc(CC2CCN(C3CCN(C(=0)c4cnnc5ccccc54)CC3)CC2)cc1')
```

That returns a SMILES for each molecule. If you plan to do more work with the molecules after retrieving them, it is much more efficient to ask postgresql to give you the molecules in pickled form:

```
>>> curs.execute('select molregno,mol_send(m) from rdk.mols where m@>%s',('clccc2clnncc2',))
>>> row = curs.fetchone()
>>> row
(9830, <read-only buffer for 0x...>)
```

These pickles can then be converted into molecules:

```
>>> from rdkit import Chem
>>> m = Chem.Mol(str(row[1]))
>>> Chem.MolToSmiles(m,True)
'CC(C)Sclccc(CC2CCN(C3CCN(C(=0)c4cnnc5cccc54)CC3)CC2)cc1'
```

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The intent of this license is similar to that of the RDKit itself. In simple words: "Do whatever you want with it, but please give us some credit."

CHAPTER

SEVEN

ADDITIONAL INFORMATION

- Python API Documentation
- C++ API Documentation