RDKit: A software suite for cheminformatics, computational chemistry, and predictive modeling

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Availability

 SourceForge Page: http://rdkit.sourceforge.net

- Source code:
 - Browse:

http://svn.sourceforge.net/viewcvs.cgi/rdkit/trunk/

- Download using subversion:
 svn co https://svn.sourceforge.net/svnroot/rdkit/trunk rdkit
- Binaries: available for Win32
- Licensing: BSD license (except for GUI components, which are GPL)

Components

End User

- Command-line scripts
- Python interface
- Database extensions

Developer

- Python programming library
- C++ programming library

General Molecular Functionality

- Input/Output: SMILES/SMARTS, mol, SDF, TDT
- "Cheminformatics":
 - Substructure searching
 - Canonical SMILES
 - Chirality support
 - Chemical transformations
 - Chemical reactions
 - Molecular serialization (e.g. mol <-> text)
- 2D depiction, including constrained depiction and mimicking 3D coords
- 2D->3D conversion/conformational analysis via distance geometry
- UFF implementation for cleaning up structures
- Fingerprinting (Daylight-like, circular, atom pairs, topological torsions, "MACCS keys", etc.)
- Similarity/diversity picking (include fuzzy similarity)
- 2D pharmacophores
- Gasteiger-Marsili charges
- Hierarchical subgraph/fragment analysis
- Hierarchical RECAP implementation

General Molecular Functionality, cntd

- Feature maps and feature-map vectors
- Shape-based similarity
- Molecule-molecule alignment
- Shape-based alignment (subshape alignment)*
- Very fast 3D pharmacophore searching
- Integration with PyMOL for 3D visualization
- Database "cartridge" (PostgreSQL, sqlite coming) *

* functional implementations, but not really recommended for use

General "QSAR" Functionality

- Molecular descriptor library:
 - Topological (κ3, Balaban J, etc.)
 - Electrotopological state (EState)
 - clogP, MR (Wildman and Crippen approach)
 - "MOE like" VSA descriptors
 - Feature-map vectors
- Machine Learning:
 - Clustering (hierarchical)
 - Information theory (Shannon entropy)
 - Decision trees, naïve Bayes*, kNN*
 - Bagging, random forests
 - Infrastructure:
 - data splitting
 - shuffling (y scrambling)
 - out-of-bag classification
 - serializable models and descriptor calculators
 - enrichment plots, screening, etc.

* functional implementations, but not really recommended for use

Command Line Tools

- ML/BuildComposite.py: build models
- ML/ScreenComposite.py: screen models
- ML/EnrichPlot.py: generate enrichment plot data
- ML/AnalyzeComposite.py: analyze models (descriptor levels)
- Chem/Fingerprints/FingerprintMols.py: generate 2D fingerprints
- Chem/BuildFragmentCatalog.py: CASE-type analysis with a hierarchical catalog

Database Utilities

Reading/Writing Molecules

```
MolFromSmiles
                          mol = Chem.MolFromSmiles('CCCOCC')
MolFromSmarts
MolFromMolBlock
MolFromMolFile
SmilesMolSupplier
SDMolSupplier
                          mols = [x for x in Chem.SDMolSupplier('input.sdf')]
TDTMolSupplier
SupplierFromFilename
MolToSmiles
MolToSmarts
                          print >>outF, Chem.MolToMolBlock(mol)
MolToMolBlock
                          w = Chem.SDWriter('out.sdf')
SmilesWriter
                          for m in mols:
SDWriter
                             w.write(m)
TDTWriter
```

Working with molecules: Substructure matching

```
>>> def findcore(m):
>>> m = Chem.MolFromSmiles('0=CCC=0')
                                                     smi = Chem.MolToSmiles(m)
>>> p = Chem.MolFromSmarts('C=0')
                                                     patt = Chem.MolFromSmarts('[D1]')
>>> m.HasSubstructMatch(p)
                                                     while 1:
True
                                                        m2 = Chem.DeleteSubstructs(m, patt)
>>> m.GetSubstructMatch(p)
                                                        nSmi = Chem.MolToSmiles(m2)
(1, 0)
                                                        m = m2
>>> m.GetSubstructMatches(p)
                                                        if nSmi==smi:
((1, 0), (3, 4))
                                                           break
>>> m = Chem.MolFromSmiles('C1CCC1C(=0)0')
>>> p= Chem.MolFromSmarts('C=0')
                                                      else:
>>> m2 =Chem.DeleteSubstructs(m,p)
                                                            smi = nSmi
>>> Chem.MolToSmiles(m2)
                                                     return m
'0.C1CCC1'
                                               >>> m = Chem.MolFromSmiles('CCC1CCC1')
>>>
                                               >>> c = findcore(m)
                                               >>> Chem.MolToSmiles(c)
```

'C1CCC1'

Working with molecules: properties

```
>>> suppl = Chem.SDMolSupplier('divscreen.400.sdf')
>>> m = suppl.next()
>>> list(m.GetPropNames())
['Formula', 'MolWeight', 'Mol_ID', 'Smiles', 'cdk2_ic50', 'cdk2_inhib',
'cdk act bin 1', 'mol name', 'scaffold', 'sourcepool']
>>> m.GetProp('scaffold')
'Scaffold 00'
>>> m.GetProp('missing')
Traceback (most recent call last):
  File "<stdin>", line 1, in ?
KeyError: 'missing'
>>> m.HasProp('missing')
>>> m.SetProp('testing','value')
>>> m.GetProp('testing')
'value'
>>> m.SetProp('calcd','45',computed=True)
>>> list(m.GetPropNames())
['Formula', 'MolWeight', 'Mol_ID', 'Smiles', 'cdk2_ic50', 'cdk2_inhib',
'cdk_act_bin_1', 'mol_name', 'scaffold', 'sourcepool', 'testing']
```

Generating Depictions

```
>>> from rdkit import Chem
>>> from rdkit.Chem import AllChem
>>> m = Chem.MolFromSmiles('C1CCC1')
>>> m.GetNumConformers()
0
>>> AllChem.Compute2DCoords(m)
0
>>> m.GetNumConformers()
1
>>> print Chem.MolToMolBlock(m)
                           0999 V2000
                0 0
                         0
   1.0607
             0.0000
                       0.0000 C
   -0.0000
            -1.0607
                       0.0000 C
                                  0 0
                                       0 0 0 0
                                                   0 0 0 0 0
                                                                 0
   -1.0607
             0.0000
                       0.0000 C
                                                                  0
   0.0000
             1.0607
                       0.0000 C
                                                                  0
       1 0
    2
    3 1 0
    4 1 0
       1 0
  END
```

>>>

Generating 3D Coordinates

```
>>> from rdkit import Chem
>>> from rdkit.Chem import AllChem
>>> m = Chem.MolFromSmiles('C1CCC1')
>>> AllChem.EmbedMolecule(m)
0
>>> m.GetNumConformers()
1
>>> AllChem.UFFOptimizeMolecule(m)
0
>>> m.SetProp(' Name', 'testmol')
>>> print Chem.MolToMolBlock(m)
testmol
 4 4 0 0 0 0 0
                     0 0 0999 V2000
  -0.8040 0.5715 -0.2537 C
  -0.3727 -0.9165 -0.2471 C 0 0 0 0 0 0 0 0 0 0 0
   0.7942 -0.5376 0.6386 C
   0.3825
          0.8826 0.6323 C
 1 2 1 0
 2 3 1 0
  3 4 1 0
 4 1 1 0
M END
>>> list(AllChem.EmbedMultipleConfs(m, 10))
[0, 1, 2, 3, 4, 5, 6, 7, 8, 9]
>>> m.GetNumConformers()
10
```

Low-budget conformational analysis

```
>>> from rdkit import Chem
>>> from rdkit.Chem import AllChem
>>> from rdkit.Chem import PyMol
>>> v = PyMol.MolViewer()

>>> m=Chem.MolFromSmiles('OC(=0)C1CCC(CC(=0)0)CC1')
>>> AllChem.EmbedMultipleConfs(m,10)
<rdBase._vectint object at 0x00A2D2B8>

>>> for i in range(m.GetNumConformers()):
... AllChem.UFFOptimizeMolecule(m,confId=i)
... v.ShowMol(m,confId=i,name='conf-%d'%i,showOnly=False)

>>> w = Chem.SDWriter('foo.sdf')
>>> for i in range(m.GetNumConformers()):
... w.write(m,confId=i)
...
>>> w.flush()
```

Molecular Miscellany

```
>>> from rdkit import Chem
>>> from rdkit.Chem import Crippen
>>> Crippen.MolLogP(Chem.MolFromSmiles('c1ccccn1'))
1.0815999999999999
>>> Crippen.MolMR(Chem.MolFromSmiles('c1ccccn1'))
24.236999999999991
>>> AllChem.ShapeTanimotoDist(m1, m2)
>>> Chem.Kekulize(m)
>>> m = Chem.MolFromSmiles('F[C@H]([Cl])Br')
>>> Chem.AssignAtomChiralCodes(m)
>>> m.GetAtomWithIdx(1).GetProp('_CIPCode')
'R'
>>> m = Chem.MolFromSmiles(r'F\C=C/Cl')
>>> Chem.AssignBondStereoCodes(m)
>>> m.GetBondWithIdx(1).GetStereo()
Chem.rdchem.BondStereo.STEREOZ
```

Database CLI tools

```
# building a database:
% python $RDBASE/Projects/DbCLI/CreateDb.py --dbDir=bzr --molFormat=sdf bzr.sdf
# similarity searching:
% python $RDBASE/Projects/DbCLI/SearchDb.py --dbDir=bzr --molFormat=smiles \
  --similarityType=AtomPairs --topN=5 bzr.smi
[18:23:21] INFO: Reading query molecules and generating fingerprints
[18:23:21] INFO: Finding Neighbors
[18:23:21] INFO: The search took 0.1 seconds
[18:23:21] INFO: Creating output
Alprazolam, Alprazolam, 1.000, Ro13-9868, 0.918, Triazolam, 0.897, Estazolam, 0.871, U-35005, 0.870
Bromazepam, Bromazepam, 1.000, Ro05-3072, 0.801, Ro05-3061, 0.801, Nordazepam, 0.801, Ro05-2921, 0.772
Delorazepam, Delorazepam, 1.000, Ro05-4619, 0.900, Nordazepam, 0.881, Lorazepam, 0.855, Ro20-8065, 0.840
# substructure and property searching:
% python $RDBASE/Projects/DbCLI/SearchDb.py --dbDir=bzr --smarts='c1ncccc1' \
  -q 'activity>6.5' --sdfOut=search1.sdf
[18:27:25] INFO: Doing substructure query
[18:27:25] INFO: Found 1 molecules matching the guery
[18:27:25] INFO: Creating output
Bromazepam
[18:27:25] INFO: Done!
```

RDKit Developer's Overview

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Installation

- Download and install boost libraries (www.boost.org)
- Download distribution from http://rdkit.sourceforge.net (or get the latest version using subversion)
- Download and install other python dependencies (see \$RDBASE/Docs/SoftwareRequirements.txt)
- Follow build instructions in \$RDBASE/INSTALL

Documentation

Main Page | Namespace List | Class Hierarchy | Class List | <mark>Directories</mark> | File List | Class Members | File Members

C++:

Generated using doxygen: cd \$RDBASE/Code doxygen doxygen.config

Python:

Currently generated using epydoc: cd \$RDBASE/rdkit epydoc --config epydoc.config

RDCode Directories

This directory hierarchy is sorted roughly, but not completely, alphabetically:

- Catalogs
- ChemicalFeatures
- DataManip
 - MetricMatrixCalc
- DataStructs
- DistGeom
- Features
- ForceField
 - UFF
- Geometry
- GraphMol
 - Depictor
 - Descriptors
 - DistGeomHelpers
 - ◇ FeatTrees
 - FileParsers
 - Fingerprints
 - ForceFieldHelpers
 - UFF
 - ⋄ FragCatalog
 - MolAlign
 - MolChemicalFeatures

 - PartialCharges
 - ShapeHelpers
 - SmilesParse
 - Subgraphs
 - Substruct
- ML
 - Cluster
 - Murtagh
 - ∘ Data
 - InfoTheory
- Numerics
 - Alignment
 - EigenSolvers
 - Optimizer
- Query
- RDBoost
- RDGeneral
- SimDivPickers

Organization

\$RDBASE/

- External/: essential 3rd party components that are hard to get or have been modified
- Code/: C++ library, wrapper, and testing code
- rdkit/: Python library, GUI, and scripts
- Docs/: Manually and automatically generated documentation
- Data/: Testing and reference data
- Scripts/: a couple of useful utility scripts
- Web/: some simple CGI applications
- bin/: installation dir for DLLs/shared libraries

C++ "Guiding Ideas"

- If boost already has the wheel, don't re-invent it.
- Try to keep classes lightweight
- Maintain const-correctness
- Test, test, test
- Include tests with code
- Keep namespaces explicit (i.e. no using namespace std;)
- Keep most everything in namespace RDKit
- Test, test, test

Test Harness

Sample test_list.py:

```
import sys
tests=[
 ("testExecs/itertest.exe", "", {}),
  ("testExecs/MolOpsTest.exe", "", {}),
  ("testExecs/testCanon.exe", "C10CCC1 C1CC0C1", (}),
  ("python", "test list.py", { 'dir': 'Depictor'}),
  ("python", "test list.py", { 'dir': 'ShapeHelpers'})
if sys.platform != 'win32':
  tests.extend([
    ("testExecs/cptest.exe", "", {}),
    ("testExecs/querytest.exe", "", {}),
    1)
longTests=[
if name ==' main ':
  import sys
 import TestRunner
  failed,tests = TestRunner.RunScript('test list.py',0,1)
  sys.exit(len(failed))
```

Wrapping code using BPL: Intro

- boost::python is not an automatic wrapper generator: wrappers must be written by hand.
- very flexible handling of "primitive types"
- tight python type integration
- allows subclassing of C++ extension classes

Wrapping code using BPL: defining a module

DataStructs/Wrap/DataStructs.cpp

```
#include <boost/python.hpp>
#include <RDBoost/Wrap.h>
#include "DataStructs.h"
namespace python = boost::python;
void wrap SBV();
void wrap EBV();
BOOST_PYTHON_MODULE(cDataStructs)
  python::scope().attr("__doc__") =
    "Module containing an assortment of functionality for basic data structures.\n"
    "\n"
    "At the moment the data structures defined are:\n"
  python::register_exception_translator<IndexErrorException>(&translate_index_error);
  python::register_exception_translator<ValueErrorException>(&translate_value_error);
 wrap SBV();
 wrap_EBV();
```

Wrapping code using BPL: defining a class

Wrapping code using BPL: making it "pythonic"

```
struct SBV_wrapper {
  static void wrap(){
    python::class_<SparseBitVect>("SparseBitVect",
                                   "Class documentation",
                                   python::init<unsigned int>())
      .def("__len__",&SBV::GetNumBits)
      .def("__getitem__",
             (const int (*)(const SBV&, unsigned int))get_VectItem)
      .def("__setitem__",
             (const int (*)(SBV&,unsigned int,int))set_VectItem)
      .def(python::self & python::self)
      .def(python::self | python::self)
      .def(python::self ^ python::self)
      .def(~python::self)
```

Wrapping code using BPL: supporting pickling

```
// allows BitVects to be pickled
struct sbv pickle suite : python::pickle suite
  static python::tuple
  getinitargs(const SparseBitVect& self)
    return python::make_tuple(self.ToString());
 };
};
struct SBV_wrapper {
  static void wrap(){
    python::class_<SparseBitVect>("SparseBitVect",
                                   "Class documentation",
                                  python::init<unsigned int>())
      .def_pickle(sbv_pickle_suite())
```

Wrapping code using BPL: returning complex types

- IntVect is a std::vector<int>
- Convertors provided for: std::vector<int>, std::vector<unsigned>, std::vector<string>, std::vector<double>, std::vector< std::vector<int> >, std::vector< std::vector< std::vector< std::list<int>, std::list< std::vector<int> > in rdBase.dll
- · Others can be created using:

```
RegisterVectorConverter<RDKit::Atom*>();
RegisterListConverter<RDKit::Atom*>();
```

Wrapping code using BPL: accepting Python types

```
void SetBitsFromList(SparseBitVect *bv, python::object onBitList) {
  PySequenceHolder<int> bitL(onBitList);
  for (unsigned int i = 0; i < bitL.size(); i++) {</pre>
    bv->SetBit(bitL[i]);
struct SBV wrapper {
  static void wrap(){
    python::class_<SparseBitVect>("SparseBitVect",
                                   "Class documentation",
                                   python::init<unsigned int>())
. . .
          .def("SetBitsFromList", SetBitsFromList,
               "Turns on a set of bits. The argument should be a tuple or list of bit
ids.\n")
. . .
```

Wrapping code using BPL: keyword arguments

GraphMol/Wrap/rdmolfiles.cpp

```
docString="Returns the a Mol block for a molecule\n\
 ARGUMENTS:\n\
n
    - mol: the molecule\n\
    - includeStereo: (optional) toggles inclusion of stereochemical\n\
                     information in the output\n\
    - confId: (optional) selects which conformation to output (-1 = default)\n\
n
 RETURNS:\n\
n\
    a string\n\
n";
  python::def("MolToMolBlock", RDKit::MolToMolBlock,
      (python::arg("mol"), python::arg("includeStereo")=false.
       python::arg("confId")=-1),
       docString.c_str());
```

This also demonstrates defining a function and taking a complex (wrapped) type as an argument

Wrapping code using BPL: returning pointers

GraphMol/Wrap/rdmolfiles.cpp

```
docString="Construct a molecule from a SMILES string.\n\n\
   ARGUMENTS: \n\
     - SMILES: the smiles string\n\
     - sanitize: (optional) toggles sanitization of the molecule.\n\
       Defaults to 1.\n\
   RETURNS: \n\
     a Mol object, None on failure.\n\
 \n":
   python::def("MolFromSmiles", RDKit::MolFromSmiles,
       (python::arg("SMILES"),
        python::arg("sanitize")=true),
       docString.c_str(),
       python::return value policy<python::manage new object>());
GraphMol/Wrap/Mol.cpp
     .def("GetAtomWithIdx",(ROMol::GRAPH_NODE_TYPE (ROMol::*)(unsigned int))
                            &ROMol::getAtomWithIdx,
     python::return_value_policy<python::reference_existing_object>(),
     "Returns a particular Atom.\n\n"
     " ARGUMENTS:\n"
        idx: which Atom to return\n\n"
        NOTE: atom indices start at 0\n")
```