Using the ProtoMol/MDL 3.0 TestSuite

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1 Obtaining the TestSuite

The testsuite for ProtoMol/MDL 3.0 is obtainable by checking out the most recent version of the software from the sourceforge.net server, using Subversion. You may checkout the software anonymously using the following command:

svn co https://protomol.svn.sourceforge.net/svnroot/protomol protomol

Upon executing this command, a protomol/ subdirectory will be produced in your working space. Change to this directory and you will find a subdirectory protomol-test which contains all regression tests for both PROTOMOL and MDL, along with Python scripts to execute the tests. Thus before using the testsuite, you must minimally have version 2.5 of Python installed (MDL requires this version anyway).

2 Running Tests

The testsuite expects two things to be contained in your executable PATH: the Python interpreter and the ProtoMol executable. Thus you must compile ProtoMol before running the testsuite, and must provide its containing directory in the PATH environment variable. Then to run the entire testsuite, from the protomol-test directory you can just run:

```
python testHarness
```

which will produce output similar to the following:

```
Test Name: [file...]
                             Result
LeapfrogTest:
                             [PASS]
NMLTest:
                             [PASS]
BornForceTest:
                             [PASS]
LeapfrogTruncatedShadowTest:
                             [PASS]
CoulombForceCutoffTest:
                             [PASS]
VelacTest:
                             [PASS]
. . . .
```

* * * * * * * * * * * * * * *	Test	Suite:	MDL	* * * * * * * * * * * * * * * * *
LeapfrogTest:				[PASS]
NMLTest:				[PASS]
BornForceTest:				[PASS]

The testsuite contains tests for both PROTOMOL (the Config testsuite) and MDL (the MDL testsuite). Each of their listings contains a set of tests which are each designed individually to test one feature of the software. For example, LeapfrogTest tests the Leapfrog propagator, NML tests Normal Mode Langevin dynamics, BornForceTest tests computation of the Born radii for implicit solvent simulations. Assuming you checked out a fresh version of the software and did not make any source code modifications, all PROTOMOL and MDL tests should initially pass.

2.1 Options

Running the testHarness script with the -h option will provide help:

```
> python testHarness -h
Usage: testHarness [options] [command]
Command:
 enable <test>
                    enable a previously disabled a test
 disable <test>
                    disable a test
  init <test>
                    initialize a test
                    reset a test by removing its .expect files
 reset <test>
 run <test>
                    run a specific test
 try <test>
                   run a test and view its results (implies -I)
                    view a test's files
 view <test>
  If no command is given all tests will be run.
Options:
 -h, --help
                      show this help message and exit
 -I, --interactive run tests in interactive mode
 -V, --valgrind
                     run tests under valgrind
                       increase verbosity
 -v
  -C, --disable-color disable color output
```

Thus the testHarness script accepts both flags or special actions to take for certain tests. As the help output states, if no command is given then all tests will be run, which we have already seen. We can enable or disable certain tests, i.e.:

> python testHarness disable ConfigTests/LeapfrogTest

It is important to note that disabling or enabling a test has effect until rerunning enable or disable, even after the test harness stops executing. Initializing a test must be performed when adding a new test to the testsuite. Alternatively, assuming correct file placement the new test will run automatically but an UNINITIALIZED message will be displayed when the test harness is executed, prompting for initialization. We outline this in Chapter 2. Resetting a test is appropriate if at some point the results produced by a test in the suite are discovered to be incorrect, this will remove all *benchmark* output which is being compared, and require the test to be reinitialized. By using the run command, we can just run one test, i.e.:

> python testHarness run ConfigTests/NMLTest

which is useful if one test is being debugged, to avoid rerunning all tests upon every modification. The try command is similar, but provides interactivity if the test fails, which we outline below. Benchmark files for a specific test can be viewed using the view command.

The flags are fairly self explanatory; we have already seen -h for the help screen, -v provides more helpful output upon a test failure, -C removes color and -v will also profile execution using ValGrind (valgrind.org) assuming it is installed. A particularly helpful flag is the -I flag, which runs tests interactively. Individual tests can be run interactively with the try command as outlined above. Interactivity provides the ability for a user to choose the action to take upon a test failure. When a test fails, output similar to the following is produced:

There are several actions that can subsequently be taken. The $\langle a \rangle$ key will abort the test harness script, and $\langle c \rangle$ and $\langle q \rangle$ take identical action by continuing the harness without taking any action. The failed test can be disabled ($\langle D \rangle$), reset ($\langle r \rangle$) to use the output currently generated as the new benchmark, or initialized ($\langle i \rangle$). Benchmark output can be viewed by pressing the $\langle v \rangle$ key. Pressing the lowercase $\langle d \rangle$ will run the diff command to generate a description of differences between currently generated and benchmark output. The $\langle 1 \rangle$ key displays any logged output from the testsuite, and $\langle t \rangle$ will rerun the test (useful if the test has been initialized or reset). Note the current test remains active until $\langle a \rangle$, $\langle c \rangle$ or $\langle q \rangle$ is pressed.

3 Creating New Tests

3.1 Test Architecture

In order to define new PROTOMOL and MDL tests, you must understand the testsuite structure and components. Within protomol-test, you will find two subdirectories ConfigTests and MDLTests. Each of these subdirectories contains a set of small Python scripts, one per test. A requirement enforced by the testsuite is that each of these scripts end in Test. To show an example, let's look at AngleForceTest:

```
#!/usr/bin/python
import os
```

os.system('python ProtoMolCheck AngleForce xyz')

Note that the last line of the script is a system call to the Python interpreter, passing as arguments a script ProtoMolCheck, the current test name (without the Test suffix), and a set of one or more file suffixes. In general, the tests for PROTOMOL will run the ProtoMolCheck Python script and the MDL tests will run the MDLCheck script. These are also contained within their respective testsuite subdirectories.

ProtoMolCheck takes as arguments the test name and file suffixes. All data files are contained within the data/ subdirectory of ConfigTests. ProtoMolCheck will execute PROTOMOL on a configuration file with the same name as the test name, concatenated with a .conf suffix. By default, ProtoMolCheck checks three types of output: standard output, standard error, and the return code (which should always be zero upon a successful execution). These outputs are compared to files in the data directory with the .expect suffix – for example, AngleForceTest.return.expect, AngleForceTest.stdout.ex etc. If file suffixes are provided as arguments to ProtoMolCheck, the DataComparator script is run to compare numerical values between a .orig file and an output file generated at runtime consisting of the test name concatenated with the appropriate suffix (for example, the above case would compare AngleForceTest.xyz and AngleForceTest.xyz.orig). Thus it is mandatory that in each test configuration file output file names which should be compared are named properly. Once compared, the generated output files are removed from the data/ directory.

The DataComparator source code is contained within a subdirectory of the protomol-test root, called src/. When the protomol-test scons script is run, the DataComparator executable is produced in the protomol-test root directory. For any test which check file output as well as defaults, the DataComparator is run. The DataComparator accepts a command line argument for tolerance when comparing values, which in the ProtoMolCheck script is set to 0.0000001. When a PROTOMOL-generated output file is compared to a .orig, each numerical value is compared to ensure the absolute value of the difference falls within this tolerance. If it does not, a Files do not match message is sent to standard output and the test fails. The DataComparator can recognize XYZ trajectory and snapshot files, as well as PDB files and DCD trajectories, and energy files.

MDL tests work very similar to those for PROTOMOL, with the differences being (1) that they are contained within a subdirectory MDLTests instead of ConfigTests, (2) that scripts with the test name concatenated with the .py extension are executed as opposed to running the PROTOMOL executable on a .conf file, and (3) the driver script is named MDLCheck instead of ProtoMolCheck. Everything else works the same.

3.2 Creating a PROTOMOL Test

Now suppose we wanted to create our own test for PROTOMOL, for a new integrator FooBar. For the sake of simplicity, we'll assume that the integrator is single timestepping (STS) and its only necessary parameter is a timestep. Suppose we wanted to test the integrator on alanine dipeptide in vacuum. An appropriate name for our new test would be FooBarTest, so our PROTOMOL configuration file must be named FooBarTest.conf and might look something like this:

```
firststep 0
numsteps 1
# Constraints
angularMomentum 0
comMotion 0
exclude scaled1-4
```

```
posfile alanine.pdb # from solvated simulation with same name
psffile alanine.psf
parfile alanine.par
temperature 310
boundaryConditions vacuum
cellManager Cubic
Integrator {
 level 0 FooBar {
        timestep 0.5
    force Bond
    force Angle
    force Dihedral
    force Improper
    force LennardJones Coulomb
       -algorithm NonbondedSimpleFull
  }
}
finpdbposfile FooBarTest.pdb
```

Note that we generated two types of output: positions in a PDB snapshot (most appropriate to test when dealing with integrators) and an energies file, and that we named them using the test name concatenated with the appropriate suffix. We place this file in the directory ConfigTests/data. Now, we must create a FooBarTest script within the ConfigTests/ directory one level up in the hierarchy. We must call ProtoMolCheck through a system call to the Python interpreter, passing the test name and appropriate suffixes which are in this case pdb and energies:

import os

os.system('python ProtoMolCheck FooBar pdb energies')

allenergiesfile FooBarTest.energies

The test has now been created, because the testHarness automatically executes all Python test scripts within the ConfigTests and MDLTests subdirectories of the root. When the testHarness script is executed, you'll see the following output:

FooBarTest:

[UNINITIALZED]

If a test is UNINITIALIZED, that means that the .expect and .orig files have not been created yet. To initialize and create these files, you'll want to run the testHarness with the -I option, which pauses on any test that does not pass and provides you with choices:

python testHarness -I

which will stop after detecting an uninitialized test:

```
FooBarTest: [UNINITIALZED]
(a)bort (c)ontinue (D)isable (d)iff (i)nitialize (l)og (r)eset
(t)est (q)uit (v)iew?
```

Now you will want to press the <I> key to initialize this test. Once initialized, the testHarness script will continue. If you run the script a second time, the test should pass:

FooBarTest:

[PASS]

3.3 Creating an MDL Test

Once the new FooBar integrator has been wrapped for MDL, we can also develop an analogous MDL test for the integrator. The Python script FooBarTest.py for running the same simulation protocol as the configuration file for the PROTOMOL test, would look like this:

```
from MDL import *
phys = Physical()
io = IO()
io.readPDBPos(phys, "data/alanine.pdb")
io.readPSF(phys, "data/alanine.psf")
io.readPAR(phys, "data/alanine.par")
phys.bc = "Vacuum"
phys.temperature = 310
forces = Forces()
ff = forces.makeForceField(phys, "charmm")
io.files = {'energies':('data/FooBarTest.energies', 1)}
prop = Propagator(phys, forces, io)
prop.propagate("FooBar", steps=1, dt=0.5, forcefield=ff)
io.writePDBPos(phys, 'data/FooBarTest.pdb')
```

This file is placed in the MDLTests/data directory, and we next must create a FooBarTest script one level higher in the directory hierarchy, which runs MDLCheck:

```
import os
os.system('python MDLCheck FooBar pdb energies')
```

That's it! Rules with respect to initialization also apply to MDL tests.

4 **Regression Testing Rules**

As a rule of thumb, the regression testsuite should be executed upon any change of PROTOMOL or MDL source code viewed as a permanent change. This of course has different meanings for developers and users; for developers the testsuite should be executed before committing updated source code to the centralized repository, for users the testsuite should be executed to ensure the software still runs properly after any local source code modifications.

Note that changing PROTOMOL source code can affect the precompiled binaries of MDL, and so both sets of regression tests should be run upon changes to PROTOMOL. Changes to pure Python libraries of MDL only require the MDL tests to be run to ensure validity. This is an implication of the multi-tiered design of the problem solving environment: higher level layers (i.e. a domain-specific language like MDL) only depend on levels lower in the hierarchy (i.e. the PROTOMOL computational back end).

5 Contact Info

If you have questions or comments, please direct them to:

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