- Your script from the OpenMM script builder should look like the window on the right
- Create a directory called "OpenMM" on the Desktop.
- Save the script as "MD_ubq.py" and move it into the "OpenMM" directory.
- Copy "1ubq.pdb" from the VMD tutorial into the same "OpenMM" directory.

```
# this script was generated by openmm-builder. to customize it further,
 # you can save the file to disk and edit it with your favorite editor.
 from __future__ import print_function
 from simtk.openmm import app
 import simtk.openmm as mm
 from simtk import unit
 from sys import stdout
 pdb = app.PDBFile('1ubq.pdb')
 forcefield = app.ForceField('amber99sbildn.xml', 'amber99_obc.xml')
 system = forcefield.createSystem(pdb.topology,
    nonbondedMethod=app.CutoffNonPeriodic, nonbondedCutoff=1.0*unit.nanometers,
    constraints=app.HBonds, rigidWater=True)
 integrator = mm.LangevinIntegrator(300*unit.kelvin, 1.0/unit.picoseconds,
    2.0*unit.femtoseconds)
 integrator.setConstraintTolerance(0.00001)
 platform = mm.Platform.getPlatformByName('CPU')
 simulation = app.Simulation(pdb.topology, system, integrator, platform)
 simulation.context.setPositions(pdb.positions)
 print('Minimizing...')
 simulation.minimizeEnergy()
 simulation.context.setVelocitiesToTemperature(300*unit.kelvin)
 simulation.reporters.append(app.DCDReporter('trajectory.dcd', 100))
 simulation.reporters.append(app.StateDataReporter(stdout, 100, step=True,
    time=True, potentialEnergy=True, temperature=True, progress=True,
    remainingTime=True, speed=True, totalSteps=1000, separator='\t'))
 print('Running Production...')
 simulation.step(1000)
 print('Done!')
                                    MD_ubq.py
OpenMM Script Builder
                                                   Save Script
                            Simulation
                                            Integrator
                                            # this script was generated by openmm-builder.
```

- We are not completely ready for the simulation because 1ubq.pdb
 - does not have hydrogen molecules
 - includes water, which does not help when we have an implicit solvent model
- OpenMM provides a package that does some simple modeling, including these required tasks.
- To make our script use this capability, open a text editor and add the five lines shown in the box

```
pdb = app.PDBFile('1ubq.pdb') =
forcefield = app.ForceField('amber99sbildn.xml', 'amber99_obc.xml') =

modeller = app.Modeller(pdb.topology, pdb.positions) =
modeller.deleteWater() =
modeller.addHydrogens(forcefield) =
pdb = modeller =
app.PDBFile.writeFile(pdb.topology, pdb.positions, open('ubq_mod.pdb', 'w')) =

system = forcefield.createSystem(pdb.topology, =
constraints=app.HBonds, rigidWater=True) =
integrator = mm.LangevinIntegrator(300*unit.kelvin, 1.0/unit.picoseconds, =
app.pdb = app.PDBFile.xml.pdb = app.PDBFile.xml.pdb = app.pdb = app
```