# Coarse-grained OpenMM Documentation

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This documentation is generated automatically using Sphinx, which reads all docstring-formatted comments from Python functions in the 'cg\_openmm' repository. (See cg\_openmm/doc for Sphinx source files.)

CONTENTS 1

**CHAPTER** 

ONE

## BUILDING OPENMM OBJECTS FOR COARSE GRAINED MODELING

All OpenMM simulations require a System() and a Topology(). This chapter details procedures for building those objects for coarse grained models with user-defined properties.

## 1.1 Building an OpenMM System() for a coarse grained model.

An OpenMM System() object contains force definitions for a molecular model. The 'cg\_openmm' repository applies default definitions for all forces (see OpenMM user guide for detailed definitions).

Shown below are tools and functions needed to build and verify an OpenMM System().

```
build.cg_build.build_system(cgmodel)
```

Builds an OpenMM System() object, given a CGModel() as input.

Parameters cgmodel (class) - CGModel() class object

#### **Returns**

• system (System()) - OpenMM System() object

#### **Example**

```
>>> from foldamers.cg_model.cgmodel import CGModel
>>> cgmodel = CGModel()
>>> system = build_system(cgmodel)
>>> cgmodel.system = system
```

#### build.cg\_build.verify\_system(cgmodel)

Given a CGModel() class object, this function confirms that its OpenMM System() object is configured correctly.

Parameters cgmodel (class) - CGModel() class object

#### **Example**

```
>>> from foldamers.cg_model.cgmodel import CGModel
>>> cgmodel = CGModel()
>>> verify_system(cgmodel)
```

**Warning:** The function will force an error exit if the system is invalid, and will proceed as normal if the system is valid.

## 1.2 Building an OpenMM Topology() for a coarse grained model.

An OpenMM Topology() object contains structural definitions for a molecular model (bond assignments, residue assignments, etc.).

Shown below are tools and functions needed to build and verify an OpenMM Topology().

build.cg\_build.build\_topology (cgmodel, use\_pdbfile=False, pdbfile=None) Construct an OpenMM Topology() class object for our coarse grained model,

#### **Parameters**

- cgmodel (class) CGModel() class object
- **use\_pdbfile** (*Logical*) Determines whether or not to use a PDB file in order to generate the Topology().
- **pdbfile** (str) Name of a PDB file to use when building the topology.

#### **Returns**

• topology (Topology() ) - OpenMM Topology() object

#### **Example**

**Warning:** When 'use\_pdbfile'=True, this function will use the PDBFile() class object from OpenMM to build the Topology(). In order for this approach to function correctly, the particle names in the PDB file must match the particle names in the coarse grained model.

```
build.cq_build.verify_topology(cgmodel)
```

Given a coarse grained model that contains a Topology() (cgmodel.topology), this function verifies the validity of the topology.

Parameters cgmodel (class) - CGModel() class object.

#### **Example**

```
>>> from foldamers.cg_model.cgmodel import CGModel
>>> cgmodel = CGModel()
>>> verify_topology(cgmodel)
```

**Warning:** The function will force an error exit if the topology is invalid, and will proceed as normal if the topology is valid.

## 1.3 Configuring OpenMM Forces() for a coarse grained model.

The 'cg\_openmm' package contains multiple tools for verifying the validity of forces that are added to an OpenMM System(). These tools are shown below:

```
build.cg_build.add_force(cgmodel, force_type=None)
```

Given a 'cgmodel' and 'force\_type' as input, this function adds the OpenMM force corresponding to 'force\_type' to 'cgmodel.system'.

#### **Parameters**

- cgmodel CGModel() class object.
- type class
- **force\_type** (*str*) Designates the kind of 'force' provided. (Valid options include: "Bond", "Nonbonded", "Angle", and "Torsion")

#### **Returns**

- cgmodel (class) 'foldamers' CGModel() class object
- force (class) An OpenMM Force() object.

#### **Example**

```
>>> from foldamers.cg_model.cgmodel import CGModel
>>> cgmodel = CGModel()
>>> force_type = "Bond"
>>> cgmodel, force = add_force(cgmodel, force_type=force_type)
```

#### build.cq\_build.test\_force(cgmodel, force, force\_type=None)

Given an OpenMM Force(), this function determines if there are any problems with its configuration.

#### **Parameters**

- cgmodel (class) CGModel() class object.
- force An OpenMM Force() object.
- **force\_type** (*str*) Designates the kind of 'force' provided. (Valid options include: "Nonbonded")

#### Returns

• 'success' (Logical) - a variable indicating if the force test passed.

```
>>> from simtk.openmm.openmm import NonbondedForce
>>> from foldamers.cg_model.cgmodel import CGModel
>>> cgmodel = CGModel()
>>> force = NonbondedForce()
>>> force_type = "Nonbonded"
>>> test_result = test_force(cgmodel,force,force_type="Nonbonded")
```

#### build.cg\_build.test\_forces(cgmodel)

Given a cgmodel that contains positions and an an OpenMM System() object, this function tests the forces for cgmodel.system.

More specifically, this function confirms that the model does not have any "NaN" or unphysically large forces.

#### **Parameters**

- cgmodel CGModel() class object.
- type class

#### Returns

• success (Logical) - Indicates if this cgmodel has unphysical forces.

#### **Example**

```
>>> from foldamers.cg_model.cgmodel import CGModel
>>> cgmodel = CGModel()
>>> pass_forces_test = test_forces(cgmodel)
```

#### build.cg\_build.get\_num\_forces(cgmodel)

Given a CGModel() class object, this function determines how many forces we are including when evaluating the energy.

Parameters cgmodel (class) - CGModel() class object

#### **Returns**

• total\_forces (int) - Number of forces in the coarse grained model

```
>>> from foldamers.cg_model.cgmodel import CGModel
>>> cgmodel = CGModel()
>>> total_number_forces = get_num_forces(cgmodel)
```

## 1.4 Other tools for building and verifying the OpenMM System() and Topology()

Shown below are other utilities to build and verify a coarse grained model System()/Topology() for OpenMM:

```
build.cg_build.add_new_elements(cgmodel)
Add coarse grained particle types to OpenMM.
```

**Parameters** cgmodel (class) – CGModel object (contains all attributes for a coarse grained model).

#### Returns

• particle\_list (list) - a list of the particles that were added to OpenMM's 'Element' List.

#### **Example**

```
>>> from foldamers.cg_model.cgmodel import CGModel
>>> cgmodel = CGModel()
>>> particle_types = add_new_elements(cgmodel)
```

**Warning:** If the particle names were user defined, and any of the names conflict with existing element names in OpenMM, OpenMM will issue an error exit.

```
build.cg_build.write_xml_file (cgmodel, xml_file_name)
Write an XML-formatted forcefield file for a coarse grained model.
```

#### **Parameters**

- cgmodel (class) CGModel() class object.
- **xml\_file\_name** (*str*) Path to XML output file.

```
>>> from foldamers.cg_model.cgmodel import CGModel
>>> cgmodel = CGModel()
>>> xml_file_name = "openmm_cgmodel.xml"
>>> write_xml_file(cgmodel,xml_file_name)
```

**CHAPTER** 

**TWO** 

## OPENMM SIMULATION TOOLS FOR COARSE GRAINED MODELING

### 2.1 Building OpenMM Simulation() objects

OpenMM simulations are propagated using a Simulation() object.

Shown below are the main tools needed to build OpenMM Simulaton() objects for coarse grained modeling.

```
simulation.tools.build_mm_simulation (topology, system, positions, temper-ature=Quantity(value=300.0, unit=kelvin), simulation_time_step=None, to-tal_simulation_time=Quantity(value=1.0, unit=picosecond), out-put_pdb=None, output_data=None, print_frequency=100, test_time_step=False)
```

Build an OpenMM Simulation()

- topology (Topology()) OpenMM Topology()
- system(System()) OpenMM System()
- **positions** (Quantity() ( np.array( [cgmodel.num\_beads,3] ), simtk.unit )) Positions array for the model we would like to test
- temperature (SIMTK Unit()) Simulation temperature, default = 300.0 K
- **simulation\_time\_step** Simulation integration time step
- total\_simulation\_time Total run time for individual simulations

- output\_pdb (str) Output destination for PDB coordinates, Default = None
- **output\_data** (*str*) Output destination for non-coordinate simulation data, Default = None
- **print\_frequency** Number of simulation steps to skip when writing to output, Default = 100
- **test\_time\_step** (*Logical*) Logical variable determining if a test of the time step will be performed, Default = False

#### **Returns**

• simulation (Simulation()) - OpenMM Simulation() object

#### **Example**

```
>>> from simtk import unit
>>> from foldamers.cg_model.cgmodel import CGModel
>>> cgmodel = CGModel()
>>> topology = cgmodel.topology
>>> system = cqmodel.system
>>> positions = cgmodel.positions
>>> temperature = 300.0 * unit.kelvin
>>> simulation_time_step = 5.0 * unit.femtosecond
>>> total_simulation_time= 1.0 * unit.picosecond
>>> output_pdb = "output.pdb"
>>> output_data = "output.dat"
>>> print_frequency = 20
>>> openmm_simulation = build_mm_simulation(topology,system,
→positions, temperature=temperature, simulation_time_
→step=simulation time step, total simulation time=total simulation
→time,output_pdb=output_pdb,output_data=output_data,print_
→frequency=print_frequency,test_time_step=False)
```

simulation.tools.run\_simulation(cgmodel, output\_directory, to-tal\_simulation\_time, simulation\_time\_step, temperature, print\_frequency, output\_pdb=None, output\_data=None)

#### Run OpenMM() simulation

- cgmodel (class) CGModel() object
- output\_directory (str) Output directory for simulation data
- total\_simulation\_time Total run time for individual simulations
- **simulation\_time\_step** Simulation integration time step

- temperature Simulation temperature, default = 300.0 K
- **print\_frequency** Number of simulation steps to skip when writing to output, Default = 100

#### **Example**

```
>>> import os
>>> from simtk import unit
>>> from foldamers.cg_model.cgmodel import CGModel
>>> cgmodel = CGModel()
>>> topology = cgmodel.topology
>>> system = cgmodel.system
>>> positions = cgmodel.positions
>>> temperature = 300.0 * unit.kelvin
>>> simulation_time_step = 5.0 * unit.femtosecond
>>> total simulation time= 1.0 * unit.picosecond
>>> output_directory = os.getcwd()
>>> output_pdb = "output.pdb"
>>> output_data = "output.dat"
>>> print_frequency = 20
>>> run_simulation(cgmodel,output_directory,total_simulation_time,
→simulation_time_step, temperature, print_frequency, output_
→pdb=output pdb, output data=output data)
```

**Warning:** When run with default options this subroutine is capable of producing a large number of output files. For example, by default this subroutine will plot the simulation data that is written to an output file.

### 2.2 Running Yank replica exchange simulations

The Yank python package is used to perform replica exchange sampling with OpenMM simulations.

Shown below are the main functions and tools necessary to conduct Yank replica exchange simulations with a coarse grained model in OpenMM.

```
simulation.rep_exch.run_replica_exchange (topology, system, positions, temperature_list=None, simulation_time_step=None, total_simulation_time=Quantity(value=1.0, unit=picosecond), out-put_data='output.nc', print_frequency=100, verbose_simulation=False, exchange_attempts=None, test_time_step=False, out-put_directory=None)
```

Run a Yank replica exchange simulation using an OpenMM coarse grained model.

- topology OpenMM Topology
- system OpenMM System()
- positions Positions array for the model we would like to test
- **temperature\_list** List of temperatures for which to perform replica exchange simulations, default = None
- simulation\_time\_step Simulation integration time step
- total\_simulation\_time Total run time for individual simulations
- output\_data (string) Name of NETCDF file where we will write simulation data
- **print\_frequency** Number of simulation steps to skip when writing to output, Default = 100
- **verbose\_simulation** (*Logical*) Determines how much output is printed during a simulation run. Default = False
- **exchange\_attempts** (*int*) Number of exchange attempts to make during a replica exchange simulation run, Default = None
- **test\_time\_step** (*Logical*) Logical variable determining if a test of the time step will be performed, Default = False

• **output\_directory** (*str*) – Path to which we will write the output from simulation runs.

#### Returns

- replica\_energies ( Quantity() ( np.float( [number\_replicas,number\_simulation\_steps] ), simtk.unit ) ) The potential energies for all replicas at all (printed) time steps
- replica\_positions ( Quantity() ( np.float( [number\_replicas,number\_simulation\_steps,cgmodel.num\_beads,3] ), simtk.unit ) ) The positions for all replicas at all (printed) time steps
- replica\_state\_indices ( np.int64( [num-ber\_replicas,number\_simulation\_steps] ), simtk.unit ) The ther-modynamic state assignments for all replicas at all (printed) time steps

#### Example

Read replica exchange simulation data.

#### **Parameters**

- **system** OpenMM system object, default = None
- topology OpenMM topology object, default = None
- **temperature\_list** List of temperatures that will be used to define different replicas (thermodynamics states), default = None
- output\_data (str) Path to the output data for a Yank, NetCDF-formatted file containing replica exchange simulation data, default = None
- **print\_frequency** (*int*) Number of simulation steps to skip when writing data, default = None

#### **Returns**

- replica\_energies ( Quantity() ( np.float( [number\_replicas,number\_simulation\_steps] ), simtk.unit ) ) The potential energies for all replicas at all (printed) time steps
- replica\_positions ( Quantity() ( np.float( [number\_replicas,number\_simulation\_steps,cgmodel.num\_beads,3] ), simtk.unit ) ) The positions for all replicas at all (printed) time steps
- replica\_state\_indices ( np.int64( [number\_replicas,number\_simulation\_steps] ), simtk.unit ) The thermodynamic state assignments for all replicas at all (printed) time steps

#### Example

Make PDB files from replica exchange simulation trajectory data

#### **Parameters**

- topology OpenMM Topology
- replica\_positions Positions array for the replica exchange data for which we will write PDB files

#### **Returns**

• file\_list ( List( str ) ) - A list of names for the files that were written

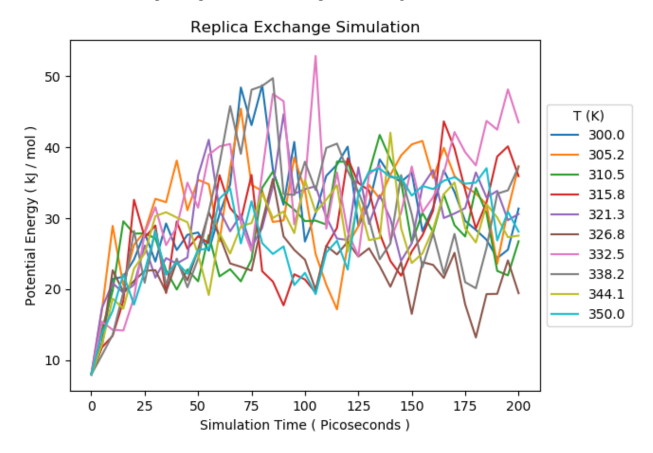
### Coarse-grained OpenMM Documentation, Release 0.0.1

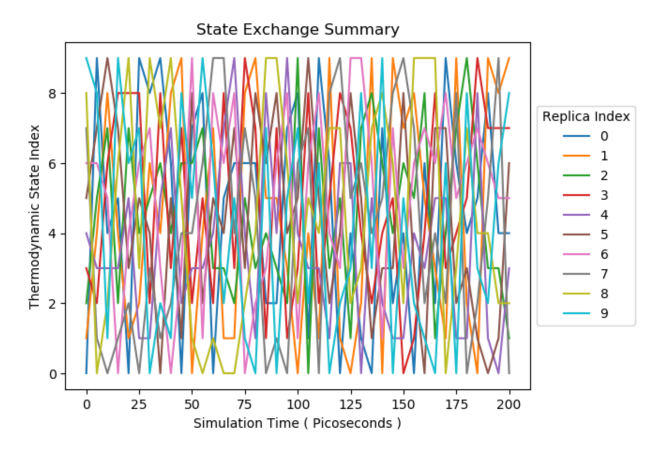
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## 2.3 Tools to plot coarse grained model simulation results

The 'cg\_openmm' package contains multiple functions to plot OpenMM and Yank simulation results using matplotlib .

Shown below is example output from a Yank replica exchange simulation run:





Shown below are functions which allow plotting of simulation results.

Plot the potential energies for a batch of replica exchange trajectories

- replica\_energies (List ( List ( float \* simtk.unit. energy for simulation\_steps ) for num\_replicas
   )) List of dimension num\_replicas X simulation\_steps, which gives the energies for all replicas at all simulation steps
- temperature\_list List of temperatures for which to perform replica exchange simulations, default = [(300.0 \* unit.kelvin).\_\_add\_\_(i \* unit.kelvin) for i in range(-20,100,10)]

- **simulation\_time\_step** Simulation integration time step
- **steps\_per\_stage** (*int*) The number of simulation steps for individual replica "stages" (period of time between state exchanges), default = 1
- **file\_name** (str) The pathname of the output file for plotting results, default = "replica\_exchange\_energies.png"
- **output\_directory** (*str*) Path to which we will write the output from simulation runs. Default = None
- **legend** (*Logical*) Controls whether a legend is added to the plot

..warning:: If more than 10 replica exchange trajectories are provided as input data, by default, this function will only plot the first 10 thermodynamic states. These thermodynamic states are chosen based upon their indices, not their instantaneous temperature (ensemble) assignment.

Plot the thermodynamic state assignments for individual temperature replicas as a function of the simulation time, in order to obtain a visual summary of the replica exchanges from a Yank simulation.

- replica\_states (List(List(float \* simtk.unit.energy for simulation\_steps)) for num\_replicas
   )) List of dimension num\_replicas X simulation\_steps, which gives the thermodynamic state indices for all replicas at all simulation steps
- **temperature\_list** List of temperatures for which to perform replica exchange simulations, default = [(300.0 \* unit.kelvin).\_\_add\_\_(i \* unit.kelvin) for i in range(-20,100,10)]
- **simulation\_time\_step** Simulation integration time step
- **steps\_per\_stage** (*int*) The number of simulation steps for individual replica "stages" (period of time between state exchanges), default = 1

- **file\_name** (str) The pathname of the output file for plotting results, default = "replica\_exchange\_state\_transitions.png"
- **legend** (*Logical*) Controls whether a legend is added to the plot
- **output\_directory** (*str*) Path to which we will write the output from simulation runs, default = None

..warning:: If more than 10 replica exchange trajectories are provided as input data, by default, this function will only plot the first 10 thermodynamic states. These thermodynamic states are chosen based upon their indices, not their instantaneous temperature (ensemble) assignment.

```
simulation.tools.plot_simulation_data(simulation_times, y_data, plot_type=None, out-put_directory=None)
```

Plot simulation data.

#### **Parameters**

- **simulation times** (*List*) List of simulation times (x data)
- y\_data (List) List of simulation data
- plot\_type (str) Form of data to plot, Default = None, Valid options include: "Potential Energy", "Kinetic Energy", "Total Energy", "Temperature"

#### **Example**

Plot all data from an OpenMM output file

#### **Parameters**

• **simulation\_data\_file** (str) – Path to file containing simulation data

- **plot\_output\_directory** (str) Path to folder where plotting results will be written.
- simulation\_time\_step Simulation integration time step

#### 2.4 Other simulation tools

Shown below are other tools which aid the building and verification of OpenMM simulation objects.

```
simulation.tools.get_mm_energy (topology, system, positions)

Get the OpenMM potential energy for a system, given a topology and set of positions.
```

#### **Parameters**

- topology OpenMM Topology()
- **system** OpenMM System()
- positions Positions array for the model we would like to test

#### **Returns**

• potential\_energy ( Quantity() ) - The potential energy for the model with the provided positions.

#### **Example**

Determine a suitable simulation time step.

- topology OpenMM Topology
- system OpenMM System()
- **positions** Positions array for the model we would like to test
- temperature Simulation temperature
- total\_simulation\_time Total run time for individual simulations
- time\_step\_list(List, default = None)-List of time steps for which to attempt a simulation in OpenMM.

#### **Returns**

- time\_step ( SIMTK Unit() ) A successfully-tested simulation time-step for the provided coarse grained model
- tolerance (SIMTK Unit()) The maximum change in forces that will be tolerated when testing the time step.

#### **Example**

```
>>> from simtk import unit
>>> from foldamers.cg_model.cgmodel import CGModel
>>> cgmodel = CGModel()
>>> topology = cgmodel.topology
>>> system = cgmodel.system
>>> positions = cgmodel.positions
>>> temperature = 300.0 * unit.kelvin
>>> total_simulation_time = 1.0 * unit.picosecond
>>> time_step_list = [1.0 * unit.femtosecond, 2.0 * unit.

-femtosecond, 5.0 * unit.femtosecond]
>>> best_time_step, max_force_tolerance = get_simulation_time_
-step(topology, system, positions, temperature, total_simulation_time,
-time_step_list=time_step_list)
```

```
simulation.tools.minimize_structure (topology, system, positions, temperature=Quantity(value=0.0, unit=kelvin), simulation_time_step=None, total_simulation_time=Quantity(value=1.0, unit=picosecond), output_pdb=None, output_data=None, print_frequency=1)
```

#### Minimize the potential energy

- topology (*Topology* ()) OpenMM topology
- **system** (System()) OpenMM system
- positions Positions array for the model we would like to test
- **temperature** Simulation temperature
- total\_simulation\_time Total run time for individual simulations
- **output\_pdb** (*str*) Output destinaton for PDB-formatted coordinates during the simulation
- output\_data (str) Output destination for simulation data

• **print\_frequency** (*int*) – Number of simulation steps to skip when writing data, default = 1

#### **Returns**

- positions ( Quantity() ( np.array( [cgmodel.num\_beads,3] ), simtk.unit )
   Minimized positions
- potential\_energy ( Quantity() Potential energy for the minimized structure.

#### **Example**

```
>>> from simtk import unit
>>> from foldamers.cg_model.cgmodel import CGModel
>>> cgmodel = CGModel()
>>> topology = cgmodel.topology
>>> system = cgmodel.system
>>> positions = cgmodel.positions
>>> temperature = 300.0 * unit.kelvin
>>> total_simulation_time = 1.0 * unit.picosecond
>>> simulation_time_step = 1.0 * unit.femtosecond
>>> output_pdb = "output.pdb"
>>> output_data = "output.dat"
>>> print_frequency = 20
>>> minimum_energy_structure, potential_energy, openmm_simulation_
→object = minimize structure(topology, system, positions,
→temperature=temperature, simulation time step=simulation time
→step, total simulation time=total simulation time, output
→pdb=output_pdb,output_data=output_data,print_frequency=print_
→frequency)
```

simulation.tools.read\_simulation\_data(simulation\_data\_file, simulation\_time\_step)

Read OpenMM simulation data

#### **Parameters**

- **simulation\_data\_file** (*str*) Path to file that will be read
- **simulation\_time\_step** Time step to apply for the simulation data

#### **Returns**

• data ( dict( "Simulation Time": list,"Potential Energy": list,"Kinetic Energy": list,"Total Energy": list,"Temperature": list ) ) - A dictionary containing the simulation times, potential energies, kinetic energies, and total energies from an OpenMM simulation trajectory.

Get an ensemble of low (potential) energy poses, and write the lowest energy structure to a PDB file if a file\_name is provided.

#### **Parameters**

- topology OpenMM Topology()
- replica\_energies (List ( List ( float \* simtk.unit. energy for simulation\_steps ) for num\_replicas
   )) List of dimension num\_replicas X simulation\_steps, which gives the energies for all replicas at all simulation steps
- replica\_positions (np.array( (float \* simtk.unit.positions for num\_beads ) for simulation\_steps )) List of positions for all output frames for all replicas
- **file\_name** Output destination for PDB coordinates of minimum energy pose, Default = None

#### Returns

• ensemble (List()) - A list of poses that are in the minimum energy ensemble.

## UTILITIES FOR COARSE GRAINED MODELING IN OPENMM

This page details the functionality of utilities in cg\_openmm/src/utilities/util.py.

```
utilities.util.distance(positions_1, positions_2)
```

Calculate the distance between two particles, given their positions.

#### **Parameters**

- **positions\_1** (Quantity() ( np.array( [3] ), simtk.unit )) Positions for the first particle
- positions\_2 Positions for the first particle

#### Returns

• distance (Quantity()) - Distance between two particles

#### Example

#### utilities.util.get\_box\_vectors(box\_size)

Given a simulation box length, construct a vector.

Parameters box\_size - Length of individual sides of a simulation box

#### **Returns**

• box\_vectors ( List( Quantity() ) ) - Vectors to use when defining an OpenMM simulation box.

```
utilities.util.lj_v (positions_1, positions_2, sigma, epsilon)
```

Calculate the Lennard-Jones interaction energy between two particles, given their positions

and definitions for their equilbrium interaction distance (sigma) and strength (epsilon).

#### **Parameters**

- positions\_1 Positions for the first particle
- positions\_2 Positions for the first particle
- **sigma** Lennard-Jones equilibrium interaction distance for two non-bonded particles
- **epsilon** Lennard-Jones equilibrium interaction energy for two non-bonded particles.

#### **Returns**

• v ( Quantity() ) - Lennard-Jones interaction energy

utilities.util.set\_box\_vectors(system, box\_size)

Impose a set of simulation box vectors on an OpenMM simulation object.

#### **Parameters**

- **system** (System()) OpenMM System()
- box\_size Length of individual sides of a simulation box

#### **Returns**

• system (System()) - OpenMM system object

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