



# ESPResSo++ Documentation

*Release latest*

**Developer team**

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## CONTENTS

<b>1</b>	<b>Installation</b>	<b>3</b>
<b>2</b>	<b>Tutorial</b>	<b>5</b>
2.1	Basic System Setup . . . . .	5
2.2	Simple Lennard Jones System . . . . .	7
2.3	Advanced Lennard Jones System . . . . .	9
2.4	Polymer Melt . . . . .	11
2.5	AddNewPotential . . . . .	13
2.6	Appendices . . . . .	16
2.7	Adaptive Resolution Simulations . . . . .	17
2.8	Thermodynamic integration . . . . .	23
<b>3</b>	<b>User Interface</b>	<b>27</b>
3.1	analysis . . . . .	27
3.2	bc . . . . .	42
3.3	check . . . . .	44
3.4	esutil . . . . .	44
3.5	external . . . . .	44
3.6	integrator . . . . .	44
3.7	interaction . . . . .	65
3.8	io . . . . .	131
3.9	espressopp . . . . .	134
3.10	standard_system . . . . .	154
3.11	storage . . . . .	156
3.12	tools . . . . .	159
3.13	Logging mechanism . . . . .	170
<b>4</b>	<b>Credits</b>	<b>173</b>
4.1	ESPresSo++ Developers . . . . .	173
4.2	FAQ . . . . .	173
4.3	Getting Help . . . . .	173
	<b>Bibliography</b>	<b>175</b>
	<b>Python Module Index</b>	<b>177</b>
	<b>Index</b>	<b>181</b>



**Welcome to the homepage of the ESPResSo++ project**

ESPResSo++ is an extensible, flexible, fast and parallel simulation software for soft matter research. It is a highly versatile software package for the scientific simulation and analysis of coarse-grained atomistic or bead-spring models as they are used in soft matter research.

ESPResSo and ESPResSo++ have common roots and share parts of the developer/user community. However their development is independent and they are different software packages.

ESPResSo++ is free, open-source software published under the GNU General Public License (GPL).

**Please cite this, if you used ESPResSo++ in your research** J. D. Halverson, T. Brandes, O. Lenz, A. Arnold, S. Bevc, V. Starchenko, K. Kremer, T. Stuehn, D. Reith, “ESPResSo++: A Modern Multiscale Simulation Package for Soft Matter Systems”, *Computer Physics Communications*, 184 (2013), pp. 1129-1149 DOI: 10.1016/j.cpc.2012.12.004 Online access: <http://dx.doi.org/10.1016/j.cpc.2012.12.004>

[Recent publications where ESPResSo++ was used](#)



## INSTALLATION

The first step in the installation of ESPResSo++ is to download the latest release from the following location:

<https://github.com/espressopp/espressopp/releases>

On the command line type:

```
tar -xzf espressopp-latest.tgz
```

This will create a subdirectory espressopp-latest

Enter this subdirectory

```
cd espressopp-latest
```

Create the Makefiles using the `cmake` command. If you don't have it yet, you have to install it first. It is available for all major Linux distributions and also for Mac OS X. (ubuntu,debian: "apt-get install cmake" or get it from <http://www.cmake.org> )

```
cmake .
```

(the space and dot after *cmake* are necessary)

If `cmake` doesn't finish successfully (e.g. it didn't find all the libraries) you can tell `cmake` manually, where to find them by typing:

```
ccmake .
```

This will open an interactive page where all configuration information can be specified. Alternatively, if `cmake .` complains on missing BOOST or MPI4PY libraries and you had not installed them, you can try

```
cmake . -DEXTERNAL_BOOST=OFF -DEXTERNAL_MPI4PY=OFF
```

In this case, ESPResSo++ will try to use internal Boost and mpi4py libraries.

After successfully building all the Makefiles you should build ESPResSo++ with:

```
make
```

(This will take several minutes)

Before being able to use the `espressopp` module in Python you need to source the `ESPRC` file:

```
source ESPRC
```

(This sets all corresponding environment variables to point to the module, e.g. `PYTHONPATH`) You have to source this file every time you want to work with `espressopp`. It would be advisable to e.g. source the file in your `.bashrc` file ( "`source <path_to_espressopp>/ESPRC`" )

In order to use `matplotlib.pyplot` for graphical output get the open source code from:

<http://sourceforge.net/projects/matplotlib>

and follow the installation instructions of your distribution.





## TUTORIAL

### Basic System Setup

ESPResSo++ is implemented as a python module that has to be imported at the beginning of every script:

```
>>> import espressopp
```

ESPResSo++ uses an object called *System* to store some global variables and is also used to keep the connection between some other important modules. We create it with:

```
>>> system = espressopp.System()
```

Starting a new simulation with ESPResSo++ we should have an idea about what we want to simulate. E.g. how big should the simulation box be or what is the density of the system or what are the interactions and the interaction ranges between our particles.

Let us start with the size of the simulation box:

```
>>> box = (10, 10, 10)
```

In many cases you will need a random number generator (e.G. to couple to a temperature bath or to randomly position particles in the simulation box). ESPResSo++ provides its own random number generator (for the experts: see `boost/random.hpp`) so let's use it:

```
>>> rng = espressopp.esutil.RNG()
```

Our simulation box needs some boundary conditions. We want to use periodic boundary conditions:

```
>>> bc = espressopp.bc.OrthorhombicBC(rng, box)
```

We tell our system object about this:

```
>>> system.bc = bc
>>> system.rng = rng
```

Now we need to decide which parallelization scheme for the particle storage we want to use. In the current version of ESPResSo++ there is only one storage scheme implemented which is *domain decomposition*. Further parallelized storages (e.g. *atom decomposition* or *force decomposition*) will be implemented in future versions.

The *domain decomposition* storage needs to know how many CPUs (or cores, if there are multicore CPUs) are available for the simulation and how to assign the CPUs to the different domains of our simulation box. Moreover the storage needs to know the maximum interaction range of the particles. In a simple Lennard-Jones fluid this could for example be  $r_{cut} = 2\frac{1}{6}$ . This value together with the *skin* value determines the minimal size for the so called *linked cells* which are used to speed up Verlet list rebuilds (see Frenkel&Smit or Allen&Tildesley for the details).

```
>>> maxcutoff = pow(2.0, 1.0/6.0)
>>> skin = 0.4
```

Tell the system about it:

```
>>> system.skin = skin
```

In the most simple case, if you want to use only one CPU, the *nodeGrid* and the *cellGrid* could look like this:

```
>>> nodeGrid = (1,1,1)
>>> cellGrid = (2,2,2)
```

In general you don't need to take care of that yourself. Just use the corresponding ESPResSo++ routines to calculate a reasonable *nodeGrid* and *cellGrid*:

```
>>> nodeGrid = espressopp.tools.decomp.nodeGrid(espressopp.MPI.COMM_WORLD.size, box, maxcutoff)
>>> cellGrid = espressopp.tools.decomp.cellGrid(box, nodeGrid, maxcutoff, skin)
```

Now we have all the ingredients we need for the *domain decomposition* storage of our system:

```
>>> ddstorage = espressopp.storage.DomainDecomposition(system, nodeGrid, cellGrid)
```

We initialized the DomainDecomposition object with a pointer to our system. We also have to inform the system about the DomainDecomposition storage:

```
>>> system.storage = ddstorage
```

The next module we need is the *integrator*. This object will do the actual work of integrating Newtons equations of motion. ESPResSo++ implements the well known *velocity Verlet* algorithm (see for example Frenkel&Smit):

```
>>> integrator = espressopp.integrator.VelocityVerlet(system)
```

We have to tell the integrator about the basic time step:

```
>>> dt = 0.005
>>> integrator.dt = dt
```

Let's do some math in between:

---

**Note:** For 3D vectors like positions, velocities or forces ESPResSo++ provides a so called *Real3D* type, which simplifies handling and arithmetic operations with vectors. 3D coordinates would typically be defined like this:

```
>>> a = espressopp.Real3D(2.0, 5.0, 6.0)
>>> b = espressopp.Real3D(0.1, 0.0, 0.5)
```

Now you could do things like:

```
>>> c = a + b          # c is a Real3D object
>>> d = a * 1.5        # d is a Real3D object
>>> e = a - b          # e is a Real3D object
>>> f = e.sqr()        # f is a scalar
>>> g = e.abs()         # g is a scalar
```

In order to make defining vectors even more simple include the line

```
>>> from espressopp import Real3D
```

just at the beginning of your script. This allows to define vectors as:

```
>>> vec = Real3D(2.0, 1.5, 5.0)
```

---

Back to our simulation:

The most simple simulation we can do is integrating Newtons equation of motion for one particle without any external forces. So let's simply add one particle to the storage of our system. Every particle in ESPResSo++ has a unique particle id and a position (this is obligatory).

```
>>> pid = 1
>>> pos = Real3D(2.0, 4.0, 6.0)      # remember to add "from espressopp import Real3D"
>>>                                   # at the beginning of your script
>>> system.storage.addParticle(pid, pos)
```

Of course nothing will happen when we integrate this. The particle will stay where it is. Add some initial velocity to the particle by adding the follow line to the script:

```
>>> system.storage.modifyParticle(pid, 'v', Real3D(1.0, 0, 0))
```

After particles have been modified make sure that this information is distributed to all CPUs:

```
>>> system.storage.decompose()
```

Now we can propagate the particle by calling the integrator:

```
>>> integrator.run(100)
```

Check the result with:

```
>>> print "The new particle position is: ", system.storage.getParticle(pid).pos
```

Let's add some more particles at random positions with random velocities and random mass and random type 0 or 1. The boundary condition object knows about how to create random positions within the simulation box. We can add all the particles at once by creating a particle list first:

```
>>> particle_list = []
>>> num_particles = 9
>>> for k in range(num_particles):
>>>     pid = 2 + k
>>>     pos = system.bc.getRandomPos()
>>>     v = Real3D(system.rng(), system.rng(), system.rng())
>>>     mass = system.rng()
>>>     type = system.rng(2)
>>>     part = [pid, pos, type, v, mass]
>>>     particle_list.append(part)
>>> system.storage.addParticles(particle_list, 'id', 'pos', 'type', 'v', 'mass')
>>> # don't forget the decomposition
>>> system.storage.decompose()
```

To have a look at the overall system there are several possibilities. The easiest way to get a nice picture is by writing out a PDB file and looking at the configuration with some visualization program (e.g. VMD):

```
>>> filename = "myconf.pdb"
>>> espressopp.tools.pdb.pdbwrite(filename, system)
```

or (if *vmd* is in your search PATH) you could directly connect to VMD by:

```
>>> espressopp.tools.vmd.connect(system)
```

or you could print all particle information to the screen:

```
>>> for k in range(10):
>>>     p = system.storage.getParticle(k+1)
>>>     print p.id, p.type, p.mass, p.pos, p.v, p.f, p.q
```

## Simple Lennard Jones System

Lets just copy and paste the beginning from the “System Setup” tutorial:

```
>>> import espressopp
>>> from espressopp import Real3D
```

```
>>>
>>> system          = espressopp.System()
>>> box              = (10, 10, 10)
>>> rng              = espressopp.esutil.RNG()
>>> bc               = espressopp.bc.OrthorhombicBC(rng, box)
>>> system.bc        = bc
>>> system.rng        = rng
>>> maxcutoff         = pow(2.0, 1.0/6.0)
>>> skin              = 0.4
>>> system.skin       = skin
>>> nodeGrid          = (1,1,1)
>>> cellGrid          = (1,1,1)
>>> nodeGrid          = espressopp.tools.decomp.nodeGrid(espressopp.MPI.COMM_WORLD.size, box, maxcutoff, skin)
>>> cellGrid          = espressopp.tools.decomp.cellGrid(box, nodeGrid, maxcutoff, skin)
>>> ddstorage         = espressopp.storage.DomainDecomposition(system, nodeGrid, cellGrid)
>>> system.storage    = ddstorage
>>>
>>> integrator        = espressopp.integrator.VelocityVerlet(system)
>>> dt                = 0.005
>>> integrator.dt     = dt
```

And lets add some random particles:

```
>>> num_particles = 20
>>> particle_list = []
>>> for k in range(num_particles):
>>>     pid = k + 1
>>>     pos = system.bc.getRandomPos()
>>>     v = Real3D(0,0,0)
>>>     mass = system.rng()
>>>     type = 0
>>>     part = [pid, pos, type, v, mass]
>>>     particle_list.append(part)
>>> system.storage.addParticles(particle_list, 'id', 'pos', 'type', 'v', 'mass')
>>> system.storage.decompose()
```

All particles should interact via a Lennard Jones potential:

```
>>> LJPot = espressopp.interaction.LennardJones(epsilon=1.0, sigma=1.0, cutoff=maxcutoff, shift=True)
```

shift=True means that the potential will be shifted at the cutoff so that  $\text{potLJ}(\text{cutoff})=0$  Next we create a VerletList which will than be used in the interaction: (the Verlet List object needs to know from which system to get its particles and which cutoff to use)

```
>>> verletlist = espressopp.VerletList(system, cutoff=maxcutoff)
```

Now create a non bonded interaction object and add the Lennard Jones potential to that:

```
>>> NonBondedInteraction = espressopp.interaction.VerletListLennardJones(verletlist)
>>> NonBondedInteraction.setPotential(type1=0, type2=0, potential=LJPot)
```

Tell the system about the newly created NonBondedInteraction object:

```
>>> system.addInteraction(NonBondedInteraction)
```

We should set the langevin thermostat in the integrator to cool down the random particle system:

```
>>> langevin          = espressopp.integrator.LangevinThermostat(system)
>>> langevin.gamma     = 1.0
>>> langevin.temperature = 1.0
>>> integrator.addExtension(langevin)
```

and finally let the system run and see how it relaxes or explodes:

```
>>> espressopp.tools.analyse.info(system, integrator)
>>> for k in range(100):
>>>     integrator.run(10)
>>> espressopp.tools.analyse.info(system, integrator)
```

Due to the random particle positions it may happen, that two or more particles are very close to each other and the resulting repulsive force between them are so high that they ‘shoot off’ in different directions with very high speed. Usually the numbers are then larger than the computer can deal with. A typical error message you get could look like this:

---

**Note:** ERROR: particle 5 has moved to outer space (one or more coordinates are nan)

---

In order to prevent this, systems that are setup in a random way and thus have strong overlaps between particles have to be “warmed up” before they can be equilibrated.

In ESPResSo++ there are several possible ways of warming up a system. As a first approach one could simply constrain the forces in the integrator. For this purpose ESPResSo++ provides an integrator Extension named CapForces. The two parameters of this Extension are the system and the maximum force that a particle can get. The following python code shows how CapForces can be used. Add it to your Lennard-Jones example just after adding the Langevin Extension:

```
>>> print "starting warmup with force capping ..."
>>> force_capping = espressopp.integrator.CapForce(system, 1000000.0)
>>> integrator.addExtension(force_capping)
>>> # reduce the time step of the integrator to make the integration numerically more stable
>>> integrator.dt = 0.0001
>>> espressopp.tools.analyse.info(system, integrator)
>>> for k in range(10):
>>>     integrator.run(1000)
>>> espressopp.tools.analyse.info(system, integrator)
```

After the warmup the time step of the integrator can be set to a larger value. The CapForce extension can be disconnected after the warmup to get the original full Lennard-Jones potential back.

```
>>> integrator.dt = 0.005
>>> integrator.step = 0
>>> force_capping.disconnect()
>>> print "warmup finished - force capping switched off."
```

## Task 1:

write a python script that creates a random configuration of 1000 Lennard Jones particles with a number density of 0.85 in a cubic simulation box. Warm up and equilibrate this configuration. Examine the output of the command

```
>>> espressopp.tools.analyse.info(system, integrator)
```

after each integration step. How fast is the energy of the system going down ? How long do you have to warmup ? What are good parameters for dt, force\_capping and number of integration steps ?

## Advanced Lennard Jones System

This tutorial needs the matplotlib.pyplot and numpy libraries and also VMD to be installed.

```
>>> import espressopp
```

After importing espressopp we import several other Python packages that we want to use for graphical output of some system parameters (e.g. temperature and energy)

```
>>> import math
>>> import time
>>> import matplotlib
>>> matplotlib.use('TkAgg')
>>> import matplotlib.pyplot as plt
>>> plt.ion()
```

We setup a standard Lennard-Jones system with 1000 particles and a density of 0.85 in a cubic simulation box. ESPResSo++ provides a “shortcut” to setup such a system:

```
>>> N = 1000
>>> rho = 0.85
>>> L = pow(N/rho, 1.0/3)
>>> system, integrator = espressopp.standard_system.LennardJones(N, (L, L, L), dt=0.0001)
```

We also add a Langevin thermostat:

```
>>> langevin = espressopp.integrator.LangevinThermostat(system)
>>> langevin.gamma = 1.0
>>> langevin.temperature = 1.0
>>> integrator.addExtension(langevin)
```

We do a very short warmup in the beginning to get rid of “extremely” high forces

```
>>> force_capping = espressopp.integrator.CapForce(system, 1000000.0)
>>> integrator.addExtension(force_capping)
>>> espressopp.tools.analyse.info(system, integrator)
>>> for k in range(10):
>>>     integrator.run(100)
>>>     espressopp.tools.analyse.info(system, integrator)
```

Now let’s initialize a graph. So that we can have a realtime-view on what is happening in the simulation:

```
>>> plt.figure()
```

We want to observe temperature and energy of the system:

```
>>> T = espressopp.analysis.Temperature(system)
>>> E = espressopp.analysis.EnergyPot(system, per_atom=True)
```

x will be the x-axixs of the graph containing the time. yT and yE will be temperature and energy as y-axes in 2 plots:

```
>>> x = []
>>> yT = []
>>> yE = []
>>> yTmin = 0.0
>>> yEmin = 0.0
>>> x.append(integrator.dt * integrator.step)
>>> yT.append(T.compute())
>>> yE.append(E.compute())
>>> yTmax = max(yT)
>>> yEmax = max(yE)
```

Initialize the two graphs (‘ro’ means red circles, ‘go’ means green circles, see also pyplot documentation)

```
>>> plt.subplot(211)
>>> gT, = plt.plot(x, yT, 'ro')
>>> plt.subplot(212)
>>> gE, = plt.plot(x, yE, 'go')
```

We also want to observe the configuration with VMD. So we have to connect to vmd. This command will automatically start vmd (vmd has to be found in your PATH environment for this to work)

```
>>> sock = espressopp.tools.vmd.connect(system)
>>> for k in range(200):
>>>     integrator.run(1000)
>>>     espressopp.tools.vmd.imd_positions(system, sock)
```

Update the x-, and y-axes:

```
>>> x.append(integrator.dt * integrator.step)
>>> yT.append(T.compute())
>>> yE.append(E.compute())
>>> yTmax = max(yT)
>>> yEmax = max(yE)
```

Plot the temperature graph

```
>>> plt.subplot(211)
>>> plt.axis([x[0], x[-1], yTmin, yTmax*1.2 ])
>>> gT.set_ydata(yT)
>>> gT.set_xdata(x)
>>> plt.draw()
```

Plot the energy graph

```
>>> plt.subplot(212)
>>> plt.axis([x[0], x[-1], yEmin, yEmax*1.2 ])
>>> gE.set_ydata(yE)
>>> gE.set_xdata(x)
>>> plt.draw()
```

In the end save the equilibrated configurations as .eps and .pdf files

```
>>> plt.savefig('mypyplot.eps')
>>> plt.savefig('mypyplot.pdf')
```

## Polymer Melt

We first import espressopp and then define all the parameters of the simulation:

```
>>> import espressopp
>>> num_chains      = 10
>>> monomers_per_chain = 10
>>> L               = 10
>>> box             = (L, L, L)
>>> bondlen         = 0.97
>>> rc              = pow(2, 1.0/6.0)
>>> skin            = 0.3
>>> dt              = 0.005
>>> epsilon         = 1.0
>>> sigma           = 1.0
```

Like in the simple Lennard Jones tutorial we setup the system and the integrator. First the system with the excluded volume interaction (WCA, Lennard Jones type)

```
>>> system          = espressopp.System()
>>> system.rng       = espressopp.esutil.RNG()
>>> system.bc        = espressopp.bc.OrthorhombicBC(system.rng, box)
>>> system.skin      = skin
>>> nodeGrid         = espressopp.tools.decomp.nodeGrid(espressopp.MPI.COMM_WORLD.size, box, rc, skin)
>>> cellGrid         = espressopp.tools.decomp.cellGrid(box, nodeGrid, rc, skin)
>>> system.storage    = espressopp.storage.DomainDecomposition(system, nodeGrid, cellGrid)
>>> interaction       = espressopp.interaction.VerletListLennardJones(espressopp.VerletList(system,
>>> potLJ             = espressopp.interaction.LennardJones(epsilon, sigma, rc)
```

```
>>> interaction.setPotential(type1=0, type2=0, potential=potLJ)
>>> system.addInteraction(interaction)
```

Then the integrator with the Langevin extension

```
>>> integrator = espressopp.integrator.VelocityVerlet(system)
>>> integrator.dt = dt
>>> thermostat = espressopp.integrator.LangevinThermostat(system)
>>> thermostat.gamma = 1.0
>>> thermostat.temperature = temperature
>>> integrator.addExtension(thermostat)
```

Now we add the particles. Keep in mind that we want to create a polymer melt. This means that particles are “bonded” in chains. We setup each polymer chain as a random walk.

```
>>> props = ['id', 'type', 'mass', 'pos', 'v']
>>> vel_zero = espressopp.Real3D(0.0, 0.0, 0.0)
```

In providing bonding information for the particles we “setup” the bonded chains. For this we use the FixedPairList object that needs to know where and in which storage the particles can be found:

```
>>> bondlist = espressopp.FixedPairList(system.storage)
>>> pid = 1
>>> type = 0
>>> mass = 1.0
>>> chain = []
```

ESPResSo++ provides a function that will return position and bond information of a random walk. You have to provide a start ID (particle id) and a starting position which we will get from the random position generator of the boundary condition object:

```
>>> for i in range(num_chains):
>>>     startpos = system.bc.getRandomPos()
>>>     positions, bonds = espressopp.tools.topology.polymerRW(pid, startpos, monomers_per_chain, bondlen)
>>>     for k in range(monomers_per_chain):
>>>         part = [pid + k, type, mass, positions[k], vel_zero]
>>>         chain.append(part)
>>>     pid += monomers_per_chain
>>>     type += 1
>>>     system.storage.addParticles(chain, *props)
>>>     system.storage.decompose()
>>>     chain = []
>>>     bondlist.addBonds(bonds)
```

---

**Note:** try out the command

```
>>> espressopp.tools.topology.polymerRW(pid, startpos, monomers_per_chain, bondlen)
```

to see what it returns

---

Don’t forget to distribute the particles and the bondlist to the CPUs in the end:

```
>>> system.storage.decompose()
```

Finally add the information about the bonding potential. In this example we are using a FENE-potential between the bonded particles.

```
>>> potFENE = espressopp.interaction.FENE(K=30.0, r0=0.0, rMax=1.5)
>>> interFENE = espressopp.interaction.FixedPairListFENE(system, bondlist, potFENE)
>>> system.addInteraction(interFENE)
```

Start the integrator and observe how the system explodes. Like in the random Lennard Jones system, we have the same problem here: particles can strongly overlap and thus will get very high forces accelerating them to infinite



(for the computer) speed.

```
>>> espressopp.tools.analyse.info(system, integrator)
>>> for k in range(nsteps):
>>>     integrator.run(isteps)
>>>     espressopp.tools.analyse.info(system, integrator)
>>>     espressopp.tools.analyse.info(system, integrator)
```

## Task 2:

Try to warmup and equilibrate a dense polymer melt (density=0.85) by using the warmup methods that you have learned in the Lennard Jones tutorial.

## Hint:

During warmup you can slowly switch on the excluded volume interaction by starting with a small epsilon and increasing it during integration: You can do this by continuously overwriting the interaction potential after some time interval.

```
>>> potLJ = espressopp.interaction.LennardJones(new_epsilon, sigma, rc)
>>> interaction.setPotential(type1=0, type2=0, potential=potLJ)
```

## AddNewPotential

The aim of the tutorial is to implement a new interaction potential in ESPResSo++. We start with the Gromos fourth-power bond-stretching potential, because its functional form is simple and its implementation is somewhat similar to other potentials already implemented in ESPResSo++. Everything you learn in this tutorial will then be relevant for implementing any other more complicated potential.

Make sure you have a working, compiled version of ESPResSo++ before starting the tutorial.

For those who are not so familiar with C++ or interfacing python and C++, you will find some helpful notes in the appendix.

## Steps for adding a new interaction potential

1. Choose the potential and derive the force.
2. Choose the appropriate interaction template from those in `$ESPRESSOHOME/src/interaction`, e.g. `VerletListInteractionTemplate.hpp`, `FixedTripleListInteractionTemplate.hpp`
3. Create the `.cpp`, `.hpp` and `.py` files for your potential, place them in `$ESPRESSOHOME/src/interaction` and modify `$ESPRESSOHOME/src/interaction/bindings.cpp` and `$ESPRESSOHOME/src/interaction/__init__.py`
4. Compile.

These steps are described in more detail below for our tutorial example potential.

## Today's tutorial exercise

### Step 1

The potential we are implementing today is a two-body bonded potential with the form

$$V(r_{ij}) = \frac{1}{4}k_{ij}(r_{ij}^2 - r_{0,ij}^2)^2$$

noindent where  $r_{ij}$  is the distance between particles  $i$  and  $j$ . The potential has two input parameters  $r_0$  and  $k$ . Derive the force.

## Step 2

This is a 2-body interaction between a predefined (fixed) list of atom pairs. What is the appropriate interaction template to use? Choose one in `$ESPRESSOHOME/src/interaction`

Open the interaction template file. (When you close the file later, close it without saving, or else later on your compile time will be very long, because of the number of dependencies on the interaction template!) Identify the functions `addForces()` and `computeEnergy()`. Many interaction templates also contain functions such as `computeVirial()`, `computeVirialX()` (for calculating the virial in slabs along the x-direction) etc.

Find the function calls:

```
potential->_computeForce(force, dist)
```

in `addForces()` and

```
potential->_computeEnergy(r2l)
```

in `computeEnergy()`.

An interaction template can be combined with many different potentials (e.g. harmonic potential, Lennard Jones potential, etc.) Each potential will have its own C++ class containing functions to compute the energy and forces for that particular potential (see e.g. `Harmonic.cpp/hpp`, `LennardJones.cpp/hpp`) In turn, each potential can be combined with many different interaction templates.

You don't need to modify the interaction template file today. (Close it without saving!)

## Step 3

In this step we create the `.cpp`, `.hpp` and `.py` files for our potential. Let's call the potential `FourthPower`. The `FourthPower.py` file will contain the end-user python interface, and in the `FourthPower.cpp` and `FourthPower.hpp` files we will create a C++ class for our potential. We will also write a wrapper which will allow the user to call the C++ code from the python interface.

### 3(a) Interfacing potential class and interaction template

In many cases, it's not necessary to understand the contents of this section in order to implement a new potential. If you like, you can skip directly to Section 3(b) *Creating the new potential class*.

Now we need to understand how the interaction template will interface with our new class. This is done via a class template, e.g. in `Potential.hpp`, `AnglePotential.hpp`, `DihedralPotential.hpp` etc.

Still in `$ESPRESSOHOME/src/interaction`, open the file `Potential.hpp`. (When you close the file later, close it without saving, or else later on your compile time will be very long, because of the number of dependencies on the file!)

Find the functions `_computeForce(Real3D& force, const Real3D& dist)` and `_computeEnergy(real dist)` which you identified in the interaction template. Note that `_computeForce(Real3D& force, const Real3D& dist)` calls the function `_computeForceRaw(force, dist, distSqr)` and `_computeEnergy(real dist)` calls `_computeEnergySqr(dist*dist)` which calls `_computeEnergySqrRaw(distSqr)`. The functions `_computeForceRaw()` and `_computeEnergySqrRaw()` are the new functions we need to write for our new potential. They will be member methods of our new C++ class `FourthPower`.

You don't need to modify anything in `Potential.hpp` today. (Close it without saving!)

### 3(b) Creating the new potential class

An easy way to implement the new C++ class is to identify a previously implemented potential which somewhat resembles your new potential, e.g. here we could take the Harmonic potential, which is also a 2-body potential, and which has also been interfaced with the FixedPairListInteractionTemplate.

Still in `$ESPRESSOHOME/src/interaction`, copy the files `Harmonic.py`, `Harmonic.cpp` and `Harmonic.hpp` to new files `FourthPower.py`, `FourthPower.cpp` and `FourthPower.hpp`. In the new files, find and replace all occurrences of ‘Harmonic’ with ‘FourthPower’, and ‘HARMONIC’ with ‘FOURTH-POWER’.

First modify `FourthPower.hpp`.

Note the `#include` statement for `FixedPairListInteractionTemplate.hpp` and `Potential.hpp`, the files you examined in [Step 2](#) and [Step 3\(a\) Interfacing potential class and interaction template](#).

The Harmonic potential had parameters called `K` and `r0`. You can reuse these for the FourthPower potential, along with the setters and getters `setK`, `getK`, `setR0` and `getR0`. For better efficiency, you could also create a new variable which contains the square of `r0`.

Now we need functions `_computeForceRaw()` and `_computeEnergySqrRaw()`, as explained in [Step 3\(a\) Interfacing potential class and interaction template](#). Modify these functions to use the functional form of the fourth power potential as derived in [Step 1](#). Note that `Real3D dist`, which contains the vector between the two particles, has been defined as  $r_{p1} - r_{p2}$  (see `addForces()` in `FixedPairListInteractionTemplate.hpp`).

Next open `Harmonic.py` and `FourthPower.py`.

Here is an example of an end-user’s python script to add an interaction using the harmonic potential:

```
harmonicbondslist = espresso.FixedPairList(system.storage)
harmonicbondslist.addBonds(bond_list) #bond_list is a list of tuples [(particleindex,i,particleindex,j)]
harmonic_potential = espresso.interaction.Harmonic(K=10.0, r0=1.0, cutoff = 5.0, shift = 0.0)
harmonic_interaction = espresso.interaction.FixedPairListHarmonic(system, harmonicbondslist, potential)
system.addInteraction(harmonic_interaction)
```

Compare this to the contents of `Harmonic.py` to understand the python source code.

Our new potential `FourthPower` can be called by the end-user in a similar way. Since the Harmonic and FourthPower potentials have similar input parameters (`K`, `r0`) and both use the `FixedPairListInteractionTemplate`, you don’t need to make any further modifications to the file `FourthPower.py`, besides replacing ‘Harmonic’ with ‘FourthPower’.

Next open `FourthPower.cpp`.

Here you will find the C++/python interface, in the function `registerPython()`. If you want to understand this function, you will find details in [Exposing a C++ class or struct to python using boost](#). You don’t need to make any further modifications to this file, besides replacing ‘Harmonic’ with ‘FourthPower’.

### 3(c) Including the new class in espressopp

Finally, update the files `$ESPRESSOHOME/src/interaction/bindings.cpp` and `$ESPRESSOHOME/src/interaction/__init__.py` (for example by copying and modifying all the lines referring to the Harmonic potential so that they now refer to the FourthPower potential). You need to make three modifications: to include the new .hpp file, to call the new `registerPython()` wrapper, and to import everything in the new python module.

## Step 4

Move to the directory `$ESPRESSOHOME`. Update the makefiles and compile using the commands:

```
cmake .
make
```

## Advanced exercise

For an interaction potential of your choosing, follow the steps above to implement it, e.g. a non-bonded two-body interaction, probably using `VerletListInteractionTemplate` and based on the LennardJones potential, or a bonded three-body interaction, probably using `FixedTripleListInteractionTemplate.hpp` and based on the AngularHarmonic potential.

You will probably have to write setters and getters for the parameters in your potential in your `.hpp` file, and make the corresponding modifications to the function `registerPython()` in the `.cpp` file and the python user interface in the `.py` file.

## Appendices

### Exposing a C++ class or struct to python using boost

(See [http://www.boost.org/doc/libs/1\\_56\\_0/libs/python/doc/tutorial/doc/html/python/exposing.html](http://www.boost.org/doc/libs/1_56_0/libs/python/doc/tutorial/doc/html/python/exposing.html))

Say we have a C++ struct called `World`:

```
struct World
{
    World(std::string msg) : msg(msg) {}           // constructor
    void set(std::string msg) { this->msg = msg; }  // function set
    std::string greet() { return msg; }            // function greet
    std::string msg;                               // member variable
};
```

Now we write the C++ class wrapper for struct `World` to expose the constructor and the functions `greet` and `set` to python:

```
{
    class_<World>("World", init<std::string>())
        .def("greet", &World::greet)
        .def("set", &World::set)
    ;
}
```

If there are additional constructors we can also expose them using `def()`, e.g. for an additional constructor which takes two doubles:

```
class_<World>("World", init<std::string>())
    .def(init<double, double>())
    .def("greet", &World::greet)
    .def("set", &World::set)
;
```

We can also expose the data members of the C++ class or struct and the associated access (getter and setter) functions using `add_property()`, e.g. for the variable `myValue` with access functions `getMyValue` and `setMyValue`:

```
.add_property("myValue", &World::getMyValue, &World::setMyValue)
```

C++ classes and structs may be derived from other classes. Say we have the C++ struct `myDerivedStruct` which is derived from the struct `myBaseStruct`:

```
struct myBaseStruct { virtual ~myBaseStruct(); };
struct myDerivedStruct : myBaseStruct {};
```

We can wrap the base class `myBaseStruct` as explained above:

```
<Base> ("Base")
/*...*/
;
```

Now when we want to wrap the class `myDerivedStruct`, we tell boost that it is derived from the base class `myBaseStruct`:

```
class_<myDerivedStruct, bases<myBaseStruct> > ("myDerivedStruct")
/*...*/
;
```

## C++ templates

See <http://www.cplusplus.com/doc/oldtutorial/templates/>

## typedef

typedef declaration allows you to create an alias that can be used anywhere in place of a (possibly complex) type name

```
typedef DataType AliasName;
```

## Python notes

### Syntax for classes in python

(See also <https://docs.python.org/2/tutorial/classes.html>)

Here is a python class called `DerivedClassName` which is derived from two other base classes (`BaseClassName1` and `BaseClassName1`), is initialised with two variables `x` and `y` which have default values 1 and 2, and contains a function `myFunction`.

```
class DerivedClassName(BaseClassName1, BaseClassName2):
    """docstring""" #a way of providing some documentation for the class
    def __init__(self, x=1, y=2): #takes two variable which have default values 1 and 2
        self.x = x
        self.y = y
    def myFunction(self):
        return self.x * self.y
```

## PMI

PMI = parallel method invocation. For more details see the file `$ESPRESSOHOME/src/pmi.py`

## Adaptive Resolution Simulations

### Theory and Background

ESPResSo++ provides functionality to run adaptive resolution simulations using the Adaptive Resolution Simulation Scheme (AdResS). In AdResS molecules in different regions in a simulation box are described by different non-bonded force fields, typically atomistic (AT) and coarse-grained (CG). These different subregions are interfaced and coupled via a hybrid region, where the interaction smoothly changes. Molecules can diffuse between the different regions and change their interaction on the fly.

There are two different AdResS approaches: The force-based scheme, in which forces are interpolated, as well as the energy-based scheme (Hamiltonian AdResS or H-AdResS) which interpolates on the level of potential energies. In force-based AdResS (see, for example, Praprotnik et al., J. Chem. Phys. 123, 224106 (2005) as well as Annu. Rev. Phys. Chem. 59, 545 (2008)), we have for the net force between the molecules  $\alpha$  and  $\beta$

$$\mathbf{F}_{\alpha|\beta} = \lambda(\mathbf{R}_\alpha)\lambda(\mathbf{R}_\beta)\mathbf{F}_{\alpha|\beta}^{\text{AT}} + (1 - \lambda(\mathbf{R}_\alpha)\lambda(\mathbf{R}_\beta))\mathbf{F}_{\alpha|\beta}^{\text{CG}},$$

where  $\mathbf{F}_{\alpha|\beta}^{\text{AT}}$  is an AT force field based on the individual atoms belonging to the molecules  $\alpha$  and  $\beta$  and  $\lambda$  is a position dependent resolution function smoothly changing from 1 in the AT region to 0 in the CG region via the hybrid buffer region. It is evaluated based on the molecules' center of mass positions  $\mathbf{R}_\alpha$ . Note that there can of course also be bonded interactions, but these are typically not interpolated, as they are computationally usually much cheaper to evaluate than the non-bonded forces. For the sake of clarity, we omit them here.

In H-AdResS (see Potestio et al., Phys. Rev. Lett. 110, 108301 (2013)), interpolation is performed directly on potential energies in the Hamiltonian as

$$H = \sum_{\alpha} \sum_{i \in \alpha} \frac{\mathbf{p}_{\alpha i}^2}{2m_{\alpha i}} + \sum_{\alpha} \{ \lambda(\mathbf{R}_\alpha) V_{\alpha}^{\text{AT}} + (1 - \lambda(\mathbf{R}_\alpha)) V_{\alpha}^{\text{CG}} \},$$

where the first term corresponds to the kinetic energy and we again omitted intramolecular interactions. The forces obtained from this Hamiltonian are

$$\begin{aligned} \mathbf{F}_{\alpha i} = & \sum_{\beta \neq \alpha} \sum_{j \in \beta} \left\{ \frac{\lambda_{\alpha} + \lambda_{\beta}}{2} \mathbf{F}_{\alpha i|\beta j}^{\text{AT}} + \left( 1 - \frac{\lambda_{\alpha} + \lambda_{\beta}}{2} \right) \mathbf{F}_{\alpha i|\beta j}^{\text{CG}} \right\} \\ & - [V_{\alpha}^{\text{AT}} - V_{\alpha}^{\text{CG}}] \nabla_{\alpha i} \lambda_{\alpha}. \end{aligned}$$

The last term, the so-called drift force, comes from applying the position gradient on the position-dependent resolution function  $\lambda$ . It acts only in the hybrid region and unphysically pushes molecules from one region to the other. Therefore, it needs to be corrected. On the other hand, force-based AdResS, contrary to H-AdResS, does not allow a Hamiltonian formulation at all.

Usually, the force fields used in the different regions of the adaptive simulation setup have significantly different pressures given the same temperature and particle density. This pressure gradient leads, in addition to the drift force in H-AdResS, to particles being pushed across the hybrid region. Eventually, the system would evolve to an equilibrium state with a inhomogeneous density profile across the simulation box. Therefore, correction forces needs to be applied in the hybrid region to counter these effects. In H-AdResS one can use a so-called free energy correction (FEC), which on average cancels the drift force in the hybrid region (see Potestio et al., Phys. Rev. Lett. 110, 108301 (2013)). The FEC corresponds to the free energy difference between the subsystems and can, for instance, be derived from Kirkwood thermodynamic integration. An alternative approach which is typically used to cancel the pressure gradient in force-based AdResS is the so-called thermodynamic force (see Fritsch et al., Phys. Rev. Lett. 108, 170602 (2012)). It is derived by constructing the correction directly from the distorted density profile which is obtained without any correction and then refined iteratively.

## ESPResSo++ code

Several measures had to be taken to implement adaptive resolution simulations in ESPResSo++. On top of the normal particles, which serve as the CG particles in AdResS, another layer of extra AT particles is introduced such that one has access to both atomistic and CG particles throughout the whole system. A mapping between the two defines which atoms belong to which CG bead. The resolution function  $\lambda$  is implemented as a particle property of the CG particles that is updated after each integration step based on the new positions. This happens in an extension to the Velocity Verlet integrator. The actual adaptive resolution scheme is then implemented via new interaction templates that define how forces and energies are computed in force-based and energy-based AdResS. These templates use for particle pairs in the atomistic region the actual atoms, this is the AT particles, for the force and energy computation while in the CG region they use the CG particles. In the hybrid region, both are used, as defined in the equations above. The drift term of H-AdResS is implemented similarly. Furthermore, the AdResS integrator extension makes sure that the atomistic particles in the CG region travel along with the CG particles and that similarly the CG particles in the AT region are properly updated according to the new atomistic positions after each integration step. The FEC as well as a module to apply the Thermodynamic Force are implemented as integrator extensions.

In the following, we explain the new features step by step (more details about parameters etc. can be found in the documentation of the different classes).

## Address Domain Decomposition

When setting up the storage we have to use an appropriate domain decomposition that accomodates storage and proper interprocessor communication of both AT and CG particles.

```
# (H-)AdResS domain decomposition
system.storage = espressopp.storage.DomainDecompositionAddress(system, nodeGrid, cellGrid)
```

## Atomistic and Coarse-Grained particles

When adding particles to the storage, we have to define them as atomistic or coarse-grained. This has been implemented as the particle property “adrat”. If it is 0, the particle is coarse-grained. If it is 1, it is an atomistic particle.

```
# add particles to system
system.storage.addParticles(allParticles, "id", "pos", "v", "f", "type", "mass", "adrat")
```

When adding the particles as above, it is important that a set of atomistic particles belonging to one CG particle appears in the list of particles `allParticles` always after the corresponding CG particle.

Next, the `FixedTupleListAdress` defines which atomistic particles belong to which coarse-grained particles.

```
# create FixedTupleList object and add the tuples
ftpl = espressopp.FixedTupleListAdress(system.storage)
ftpl.addTuples(tuples)
system.storage.setFixedTuplesAdress(ftpl)
```

In this example, `tuples` is a list of tuples, where each tuple itself is another short list in which the first element is the CG particle and the other elements are the AT particles belonging to it. Note that in ESPResSo++ the CG particle is positioned always in the center of mass of its atoms.

Having set up the `FixedTupleList`, we can also set up an `AdResS` fixed pair list that defines bonds between AT particles within individual molecules. This is done in the following way:

```
# add bonds between AT particles
fpl = espressopp.FixedPairListAdress(system.storage, ftpl)
fpl.addBonds(bonds)
```

where `bonds` is a list of bonds between AT particles within CG molecules. Similarly, triple lists for angles, quadruple lists for dihedrals etc. are set up. Compared to conventional bonds, angles, etc. between different normal CG particles one just adds the suffix `Adress` to the appropriate list object and provides it also with the `FixedTupleList` (`ftpl` in the example). Note that you can define several different such fixed pair lists and you can, for example, also in `AdResS` simulations still use the normal `FixedPairList` to define bonds between regular CG particles.

## AdResS Verlet List

Next, we construct the `AdResS` Verlet list object for non-bonded interacting particle pairs:

```
# AdResS Verlet list
vl = espressopp.VerletListAdress(system, cutoff=0.8, adrcut=1.4,
                                dEx=1.5, dHy=1.0,
                                adrCenter=[Lx/2, Ly/2, Lz/2], sphereAdr=False)
```

We have to provide the cutoffs of the list as well as the sizes of the atomistic and hybrid regions. The parameter `cutoff` corresponds to the cutoff used for CG particle pairs with both particles being in the CG region, while `adrcut` is the cutoff for all other particle pairs (at least one particle of the pair is in the AT or hybrid region).



We want to stress that this pair list is build based on the CG particles' positions. Hence, for the AT and hybrid region one needs in some situations to provide a Verlet list cutoff (`adrcut`) slightly larger than the actual maximum interaction range of the potential, in order to not lose interactions between some atom pairs. Let us clarify this with an example: Thinking of a pair of water molecules, both coarse-grained into single beads, these CG beads could be farther apart than the interaction cutoff. Two hydrogen atoms pointing towards each other, however, could in fact still be in interaction range. Therefore, an appropriate buffer needs to be provided.

The `sphereAdr` flag decides how to geometrically set up the change in resolution. If it's true, the AT region is a spherical region positioned at `adrCenter` with radius `dEx`. If `sphereAdr` is false, the resolution changes along the x-axis of the system and `dEx` corresponds to half the width of the AT region. `dHy` always is the full width of the hybrid region. Instead of providing a 3D position for `adrCenter` as above, one can also provide a particle ID of a CG particle. In this case, the atomistic region will follow the movement of the particle. This should be only done, however, for force-based AdResS, since it would break the Hamiltonian character of H-AdResS, and also only when using a spherical adaptive geometry. Then, however, it is even possible to provide a list of particle IDs, in which case the AT region corresponds to the overlap of the spherical regions defined by the individual particles provided in the list. It will deform accordingly while these particle move.

## Interactions

When adding interactions to the system we have to use the corresponding interaction templates. Here is how to set up a non-bonded interaction in a H-AdResS system:

```
# H-AdResS non-bonded interaction: WCA potential between AT particles
# and tabulated potential between CG particles
interNB = espressopp.interaction.VerletListHadressLennardJones(vl, ftpl)
potWCA = espressopp.interaction.LennardJones(epsilon=1.0, sigma=1.0, shift='auto',
                                             cutoff=rca)
potCG = espressopp.interaction.Tabulated(itype=3, filename=tabCG, cutoff=rc) # CG
interNB.setPotentialAT(type1=1, type2=1, potential=potWCA) # AT
interNB.setPotentialCG(type1=0, type2=0, potential=potCG) # CG
system.addInteraction(interNB)
```

First, we define the appropriate interaction type, in H-AdResS this is `VerletListHadressLennardJones`. Next we define the actual potentials. Then we associate them with the H-AdResS interaction and add the interaction to the system. For force-based AdResS the only change required would be to use the `VerletListAdressLennardJones` interaction.

Note that the here used interaction, `VerletListHadressLennardJones`, couples only Lennard-Jones-type potentials with tabulated ones. However, there exist more such interaction templates for other potentials and potential combinations.

## AdResS Integrator Extension

Finally, we have to set up the AdResS integrator extension:

```
# AdResS integrator extension
adress = espressopp.integrator.Address(system, verletlist, ftpl, regionupdates = 1)
integrator.addExtension(adress)
```

It takes as arguments the Verlet list and the fixed tuple list. Additionally, for the case of a moving and/or deforming AdResS region based on one or more particles, the parameter `regionupdates` specifies how regularly we want to update the shape of the AdResS region in number of steps. This is to avoid as much as possible of the additional communication required to inform different processors of the change of the AdResS region. The parameter defaults to 1 and is not used at all for static AdResS regions.

Having set up the AdResS extension, we can distribute all particles in the box and place the CG molecules in the centers of mass of the atoms which they belong to. This can be done conveniently via

```
# distribute atoms and CG molecules according to AdResS domain decomposition,
# place CG molecules in the center of mass
espressopp.tools.AddressDecomp(system, integrator)
```



## Free Energy Compensation

When using H-AdResS, we probably want to also employ a FEC. This can be done as follows:

```
# set up FEC
fec = espressopp.integrator.FreeEnergyCompensation(system, center=[Lx/2, Ly/2, Lz/2])
fec.addForce(itype=3, filename="table_fec.tab", type=1)
integrator.addExtension(fec)
```

The FEC takes as arguments the system object as well as the center of the AT region. Then we add the actual force, which needs to be provided in a table (first column: resolution  $\lambda$ , second: energy, third: force). `itype` defines which type of interpolation should be used for values between the ones provided in the table. 1 corresponds to linear interpolation, 2 to akima splines, 3 to cubic splines. We suggest to use cubic splines. The FEC is applied on CG particles and distributed among the atoms belonging to the CG particle. `type` specifies the CG particle type for which this correction should be applied. One can, for example, use different FECs for different molecules types.

## Thermodynamic Force

When using force-based AdResS, or, alternatively, in addition to the FEC in H-AdResS, we can use the thermodynamic force. It can be set up in the following way, very similar to the FEC before:

```
# set up Thermodynamic Force
thdforce = espressopp.integrator.TDforce(system, verletlist)
thdforce.addForce(itype=3, filename="table_tf.tab", type=1)
integrator.addExtension(thdforce)
```

It works largely as for the FEC with the following differences: The table should not provide resolution values in the first column but actual distance values, this is, the distance from the (closest) AT region center. This allows to extend the application of the thermodynamic force slightly beyond the borders of the hybrid region where the resolution is constant. Furthermore, the Thermodynamic Force needs the `verletlist` as argument.

It is also possible to define a thermodynamic force, which is suited for an adaptive resolution setup with an AT region that is constructed via the overlap of several spherical regions. In this case, the extension needs more information:

```
# set up Thermodynamic Force
thdforce = espressopp.integrator.TDforce(system, verletlist, startdist = 0.9,
                                          enddist = 2.1, edgeweightmultiplier = 20)
thdforce.addForce(itype=3, filename="table_tf.tab", type=1)
integrator.addExtension(thdforce)
```

It gets three more parameters, `startdist`, `enddist` and `edgeweightmultiplier`. `startdist` explicitly says at which distance from the center of the closest AT region defining particle the thermodynamic force starts to act and `enddist` says where it ends. Hence, these value should correspond to what is actually written in the table. `edgeweightmultiplier` is a parameter that specifies how precisely the thermodynamic force should be applied in the overlap regions of different spheres. For most applications, however, 20 should provide reasonable results (for details, see Kreis et al., J. Chem. Theory Comput. 12, 4067 (2016)). The 3 additional parameters are of course also present with some default values in the basic case, but they are ignored unless we have an AT region that is constructed via the overlap of several spherical regions.

## Examples

We have provided several example scripts and setups that are available in the ESPResSo++ source code at `examples/adress`. Most of them are based on published papers.

The reader is strongly encouraged to play around with them and test what happens when the setups are modified. Possible questions to ask are provided at the end of the following subsections, which explain the individual examples in more detail.

## Force-AdResS: Tetrahedral Liquid

Subfolder: `fadress_tetraliquid`. This example consists of the system that was used in the initial work introducing the force-based adaptive resolution method (see Praprotnik et al., J. Chem. Phys. 123, 224106 (2005) and Phys. Rev. E 73, 066701 (2006)). A liquid composed of artificial tetrahedral molecules, i.e. each molecule consists of 4 bonded atoms arranged in a tetrahedral geometry, is coupled to a CG model which describes the molecules as individual beads.

Questions: The geometry is set in such a way that the resolution changes along the x-axis of the box. Try changing the setup such that the AT region is of spherical shape. You can also try removing the thermostat. Does the system conserve energy? Also vary the size of the atomistic region and see what happens. Can you also make the system all-atomistic or all-CG? You can also try to compare computational times.

## Force-AdResS: A Protein in Water

Subfolder: `fadress_protein`. This system is an aqueous solution of the regulatory protein ubiquitin. The atomistic protein and the atomistic water around it is coupled to a coarse-grained water model, which maps water molecules farther away from the protein to single beads. The CG water interaction was parametrized with iterative Boltzmann inversion (IBI). This system is similar to the setup which was used by Fogarty et al. (J. Chem. Phys. 142, 195101 (2015)) to study the structure and dynamics of a protein hydration shell.

Questions: The setup is significantly more complicated than the previous system. Try to understand the script. You can also have a look into the actual source code and try to understand, for example, how the gromacs parser works. The example is set up as a fully atomistic simulation by setting the size of the atomistic region to a value larger than the simulation box. Try to change the script such that it is an actual adaptive setup. Do not forget the thermodynamic force! Furthermore, how is the high-resolution region positioned now?

## Force-AdResS: Self-Adjusting Adaptive Resolution Simulations

Subfolder: `fadress_selfadjusting`. This setup demonstrates how force-based adaptive resolution simulations with self-adjusting high-resolution regions can be set up (Kreis et al., J. Chem. Theory Comput. 12, 4067 (2016)). The system is a polyaniline-9 molecule in aqueous solution. A spherical AT region is associated with each atom of the peptide such that the overall AT region formed by the overlap of all these spheres elegantly envelops the peptide. The peptide starts in an extended configuration and as it folds, the AT region surrounding it adjusts itself accordingly. At the outside, we use again a coarse-grained IBI single-bead model for the water molecules.

Questions: Can you change the system such that fewer atoms are associated with AT region, for example, only the heavy atoms? Can you change the update frequency of the shape of the AT region?

## H-AdResS: Tetrahedral Liquid

Subfolder: `hadress_tetraliquid`. This is the system used by Potestio et al. in the paper that proposed the H-AdResS method (Phys. Rev. Lett. 110, 108301 (2013)). It is again a simple system composed of tetrahedral molecules that change their resolution and become individual beads in the CG region. The interpolation occurs along the x-axis. This example has three subfolders.

The first folder `hadress_tetraliquid_plain` runs a simple H-AdResS simulation without any free energy correction. Hence, the drift force strongly pushes molecules from one region to the other. The script contains analysis routines which measure both a density and a pressure profile along the direction of resolution change while the simulation is running. Gathering enough statistics takes a while, but we have also provided reference profiles which are obtained after a sufficiently long simulation. Have a look at them and try to interpret them.

The second folder `hadress_tetraliquid_FEC` contains the same setup but with a free energy correction. For this, two tables are provided, `table_FEC_Helmholtz.dat` and `table_FEC_Gibbs.dat`. They were derived via Kirkwood thermodynamic integration. The first one is based on the Helmholtz free energy difference per particle between the two subsystem, and the second one corresponds to the Gibbs free energy difference per particle. Two density and pressure profiles obtained while applying these correction are also shown. Try to interpret them.

The third folder `hadress_tetraliquid_KTI` contains a simple implementation of Kirkwood thermodynamic integration (KTI) which could in principle, when run for long enough, be used to derive the FEC. This is not an adaptive resolution simulation. Instead, we tell the AdResS integrator extension that we want to run KTI. Then, the extension does not modify the resolution values associated with the different molecules and we can change them by hand during the simulation. In this way, we can set up a simulation in which we change the resolution of all molecules in the system every few steps and slowly proceed from a complete CG system to an all-atom one. Have a look and try to understand what is going on.

There are many more interesting things you can try out: Are the H-AdResS simulations energy conserving? Add the commented Langevin thermostat and compare. Also vary the timestep. Additionally, you can change the size of the hybrid region. What happens if it becomes smaller or larger? Furthermore, what happens if you change the system from H-AdResS to force-based AdResS?

## H-AdResS: Water

Subfolder: `hadress_water`. This is a slightly more advanced H-AdResS system in which an atomistic model is coupled to a coarse-grained one, mapping the three water atoms onto single beads.

Questions: Feel free to play around with the system. You could also try to figure out, how the gromacs parsers sets up the interactions and chooses the right H-AdResS interactions.

# Thermodynamic integration

## Theoretical explanation

Thermodynamic integration (TI) is a method used to calculate the free energy difference between two states A and B. For the theoretical background, see e.g. <http://www.alchemistry.org>. In this tutorial, we show how to perform TI calculations with ESPResSo++. We calculate the free energy of solvation of methanol in water. The complete python script is available in the ESPResSo+ source code under `examples/thd_integration_solvation`

To do TI, we define states A and B, with potentials  $U^A$  and  $U^B$ . We then construct a pathway of intermediate states between A and B by defining a parameter  $\lambda$  that takes values between 0 and 1 and writing the system potential  $U$  as a function of  $\lambda$ ,  $U^A$  and  $U^B$ . The free energy difference between the states A and B is then given by

$$\Delta A = \int_0^1 \left\langle \frac{dU(\lambda)}{d\lambda} \right\rangle_\lambda d\lambda$$

In practise, we discretise  $\lambda$  and perform a series of MD simulations with different  $\lambda$  values between 0 and 1, sampling  $\frac{dU(\lambda)}{d\lambda}$  in each simulation.

To calculate the solvation free energy of methanol in water, we use a box of water containing one methanol molecule. We simulate desolvation via two separate TI calculations. (Note that the procedure described here is decoupling, and solute-solute interactions will be treated differently if you're doing annihilation instead of decoupling, see Note 1.)

---

### Step 1: free energy change for switching off the Coulombic interactions

*State A:* methanol has full non-bonded (Coulomb and Lennard Jones) interactions with the solvent

*State B:* methanol has only Lennard Jones interactions with the solvent

---

### Step 2: free energy change for switching off the Lennard Jones interactions

*State A:* methanol has only Lennard Jones interactions with the solvent

*State B:* methanol has no interaction with the solvent

---

Step 1 can be done using a linear function of  $\lambda$ :

$$U(\lambda_C) = (1 - \lambda_C)U_C^A + U_{unaffected}$$

where  $U_C^A$  is the solute-solvent Coulombic interaction in state A. In ESPResSo++ the charges used for state A are the particle charges contained in the particle property `charge`. The charges in state B are zero, so  $U_C^B(q)$  does not appear in the expression. (The case where A and B both have non-zero charges is not implemented in ESPResSo++). The term  $U_{unaffected}$  is all other parts of the potential that don't change with  $\lambda_C$  including all bonded interactions, any solute-solute Coulombic interactions, solvent-solvent Coulombic interactions and all Lennard-Jones interactions. The parameter  $\lambda_C$  goes from 0 to 1 in Step 1.

Step 2 must be done using a softcore potential because of the singularity in the Lennard-Jones potential at  $r_{ij} = 0$ .

$$U(\lambda_L) = \sum_{i,j} U_L(r_{ij}, \lambda_L) + U_{unaffected}$$

$$U_L(r_{ij}, \lambda_L) = (1 - \lambda_L)U_H^A(r_A) + \lambda_L U_H^B(r_B)$$

$$r_A = (\alpha \sigma_A^6 \lambda^p + r_{ij}^6)^{1/6}$$

$$r_B = (\alpha \sigma_B^6 (1 - \lambda)^p + r_{ij}^6)^{1/6}$$

The terms  $U_H^A(r_A)$  and  $U_H^B(r_B)$  are the normal Lennard-Jones 12-6 hardcore potentials:

$$U_H^A(r_A) = 4.0\epsilon_A \left( \frac{\sigma_A^{12}}{r_A^{12}} - \frac{\sigma_A^6}{r_A^6} \right)$$

The sum  $\sum_{i,j} U_L(r_{ij}, \lambda_L)$  is over all solute-solvent interactions. The term  $U_{unaffected}$  is all other parts of the potential that don't change with  $\lambda_L$  including any solute-solute Lennard-Jones interactions and solvent-solvent Lennard-Jones interactions, which are treated using standard hardcore Lennard-Jones. (In this particular example of methanol, there are no solute-solute Lennard-Jones interactions). Finally  $\alpha$  and  $p$  are adjustable parameters of the softcore potential.

The ESPResSo++ C++ code allows for different values of  $\epsilon_A$ ,  $\epsilon_B$ ,  $\sigma_A$  and  $\sigma_B$  for every pair of atomtypes interacting via this potential. In this example, we will set  $\epsilon_B$  to 0 (we are switching off the Lennard-Jones interaction). The parameter  $\lambda_L$  goes from 0 to 1 in Step 2.

## ESPResSo++ code

We must perform many separate simulations, each with a different  $\lambda$  value. It is convenient to define a list of  $\lambda$  values in the python script and use an index to access a different element of the list in each separate simulation. The script for the first simulation contains these lines:

```
# Parameters for Thermodynamic Integration
stateBIndices = [1,2,3,4,5,6] #indices of the methanol atoms
lambdaVectorCoul = [0.00, 0.05, 0.10, 0.15, 0.20, 0.25, 0.30, 0.35, 0.40, 0.45, 0.50,
                    0.55, 0.60, 0.65, 0.70, 0.75, 0.80, 0.85, 0.90, 0.95, 1.00, 1.000,
                    1.000, 1.000, 1.000, 1.000, 1.000, 1.000, 1.000, 1.000, 1.000,
                    1.000, 1.000, 1.000, 1.000, 1.000, 1.000, 1.000, 1.000, 1.000,
                    1.000, 1.000, 1.000, 1.000, 1.000, 1.000, 1.000, 1.000, 1.000,
                    1.000, 1.000, 1.000]
lambdaVectorVdwl = [0.00, 0.00, 0.00, 0.00, 0.00, 0.00, 0.00, 0.00, 0.00, 0.00, 0.00, 0.00,
                    0.00, 0.00, 0.00, 0.00, 0.00, 0.00, 0.00, 0.00, 0.00, 0.00, 0.025,
                    0.050, 0.075, 0.100, 0.125, 0.150, 0.175, 0.200, 0.225, 0.250,
                    0.275, 0.300, 0.325, 0.350, 0.375, 0.400, 0.425, 0.450, 0.475,
                    0.500, 0.525, 0.550, 0.575, 0.600, 0.625, 0.650, 0.675, 0.700,
                    0.725, 0.750, 0.775, 0.800, 0.825, 0.850, 0.875, 0.900, 0.925,
                    0.950, 0.975, 1.000]
lambdaIndex = 0
lambdaTICoul = lambdaVectorCoul[lambdaIndex]
lambdaTIVdwl = lambdaVectorVdwl[lambdaIndex]
```

The list `lambdaVectorCoul` contains the values of  $\lambda_C$  and the list `lambdaVectorVdwl` contains the values of  $\lambda_L$ . The total number of simulations to do Step 1 and Step 2 will be `len(lambdaVectorCoul)` or `len(lambdaVectorVdwl)`. We must make a copy of the python script for each simulation, changing each time the value of `lambdaIndex`.

Next we set up the Coulombic interactions, assuming we already have created a system and a `verletlist`. The electrostatics method used is generalised reaction field.

```
#atTypes - list of all atomtypes (integers) used in the pairs interacting via this potential
#epsilon1,epsilon2,kappa - reaction field parameters
#annihilate=False means decoupling is used (see Note 1)
#ftpl - a FixedTupleListAdResS object (see AdResS tutorial)
#for non-AdResS simulations, simply set adress=False, and the parameter ftpl is not needed
qq_adres_interaction = gromacs.setCoulombInteractionsTI(system, verletlist, nbCutoff,
                                                         atTypes, epsilon1=1, epsilon2=80,
                                                         kappa=0, lambdaTI=lambdaTICoul,
                                                         pidlist=stateBIndices,
                                                         annihilate=False, adress=True, ftpl=ftpl)
```

Now we set up the softcore Lennard Jones interaction.

```
#atomtypeparameters - dictionary of format {atomtype: {'eps': epsilon, 'sig': sigma}}
#                       where atomtype is integer and epsilon and sigma are real
#defaults - dictionary containing a key 'combinationrule' with value 1 if the contents
#           of atomtypeparameters need to be converted from c6,c12 format to
#           epsilon,sigma format; can also be an empty dictionary if no conversion needed
#sigmaSC, alphaSC, powerSC - parameters of the softcore potential
alphaSC = 0.5
powerSC = 1.0
epsilonB = 0.0
sigmaSC = 0.3
lj_adres_interaction = gromacs.setLennardJonesInteractionsTI(system, defaults,
                                                             atomtypeparameters, verletlist, nbCutoff,
                                                             epsilonB=epsilonB, sigmaSC=sigmaSC, alphaSC=alphaSC,
                                                             powerSC=powerSC, lambdaTI=lambdaTIVdwl,
                                                             pidlist=stateBIndices, annihilate=False,
                                                             adress=True, ftpl=ftpl)
```

We open an output file. In the first line we write the values of  $\lambda_C$  and  $\lambda_L$  for this simulation.

```
dhdlF = open("dhdl.xvg","a")
dhdlF.write("#(coul-lambda, vdwl-lambda) = (" + str(lambdaTICoul) + ", " + str(lambdaTIVdwl) + ") \n")
```

During the MD run, every `x` number of MD steps, we return to the python level and calculate the derivatives of the energies with respect to  $\lambda$ .

```
dhdlCoul = qq_adres_interaction.computeEnergyDeriv()
dhdlVdwl = lj_adres_interaction.computeEnergyDeriv()
dhdlF.write(str(time) + " " + str(dhdlCoul) + " " + str(dhdlVdwl) + "\n")
```

After all simulations, we can now average  $\frac{dU(\lambda)}{d\lambda}$  for each value of  $\lambda_C$  or  $\lambda_L$ , integrate over  $\lambda_C$  and  $\lambda_L$ , add the values  $\Delta A_C$  and  $\Delta A_L$ , and take the negative (because the procedure described here is desolvation and we want the free energy of solvation).

## Some notes

1. This example given here uses decoupling (solute-solvent interactions are a function of  $\lambda$ , solute-solute interactions are not affected by changes in  $\lambda$ ). In ESPResSo++ it is also possible to do annihilation, where both solute-solvent and solute-solute interactions are a function of  $\lambda$ , by setting `annihilate=True` when creating the non-bonded interactions.
2. The procedure described here is desolvation. To get the free energy of solvation, we take the negative of the value obtained after integration.

3. The example Python code snippets here use the helper functions `gromacs.setLennardJonesInteractionsTI` and `gromacs.setCoulombInteractionsTI` contained in `$ESPRESSOHOME/src/tools/convert/gromacs.py`, but this is not necessary. You can do TI with ESPResSo++ without the Gromacs parser by directly calling `espresso.interaction.LennardJonesSoftcoreTI` and `espresso.interaction.ReactionFieldGeneralizedTI`. See the documentation of these two classes.

## USER INTERFACE

### analysis

#### espressopp.analysis.AllParticlePos

`espressopp.analysis.AllParticlePos.gatherAllPositions()`

**Return type**

#### espressopp.analysis.AnalysisBase

##### Overview

List of classes based on AnalysisBase:

#### espressopp.analysis.LBOutput

##### Overview

<code>espressopp.analysis.LBOutputScreen</code>
<code>espressopp.analysis.LBOutputVzInTime</code>
<code>espressopp.analysis.LBOutputVzOfX</code>

##### Details

Abstract output class for LB simulations. The implemented realisations are:

- `espressopp.analysis.LBOutputScreen` to output simulation progress and control flux conservation when using MD to LB coupling.
- `espressopp.analysis.LBOutputVzInTime` to output velocity component  $v_z$  on the lattice site with an index  $(0.25 * N_i, 0, 0)$  in time.
- `espressopp.analysis.LBOutputVzOfX` to output local density  $\rho$  and  $v_z$  component of the velocity as a function of the coordinate  $x$ .

---

**Note:** all derived output classes have to be called from class `espressopp.integrator.ExtAnalyze` with specified periodicity of invocation and after this added to the integrator. See examples.

---

### espressopp.analysis.LBOutputScreen

Computes and outputs to the screen the simulation progress (finished step) and controls mass flux conservation when using MD-to-LB coupling. Ideally, the sum of mass fluxes should be 0, i.e.  $j_{LB} + j_{MD} = 0$ .

**class** espressopp.analysis.LBOutputScreen (system, lb)

#### Parameters

- **system** (*shared\_ptr*) – system object defined earlier in the python-script
- **lb** (*lb\_object*) – lattice boltzmann object defined earlier in the python-script

Example:

```
>>> # initialise output to the screen
>>> outputScreen = espressopp.analysis.LBOutputScreen(system, lb)
>>>
>>> # initialise external analysis object with previously created output object
>>> # and periodicity of invocation (steps):
>>> extAnalysis = espressopp.integrator.ExtAnalyze(outputScreen, 100)
>>>
>>> # add the external analysis object as an extension to the integrator
>>> integrator.addExtension( extAnalysis )
```

### espressopp.analysis.LBOutputVzInTime

Computes and outputs the velocity component  $v_z$  in time on the lattice site with an index  $(0.25 * N_i, 0, 0)$ .

**class** espressopp.analysis.LBOutputVzInTime (system, lb)

#### Parameters

- **system** (*shared\_ptr*) – system object defined earlier in the python-script
- **lb** (*lb\_object*) – lattice boltzmann object defined earlier in the python-script

Example:

```
>>> # initialise output of the Vz as a function of time
>>> outputVzInTime = espressopp.analysis.LBOutputVzInTime(system, lb)
>>>
>>> # initialise external analysis object with previously created output object
>>> # and periodicity of invocation (steps):
>>> extAnalysis = espressopp.integrator.ExtAnalyze(outputVzInTime, 100)
>>>
>>> # add the external analysis object as an extension to the integrator
>>> integrator.addExtension( extAnalysis )
```

### espressopp.analysis.LBOutputVzOfX

Computes and outputs simulation progress (finished step) and controls flux conservation when using MD to LB coupling.

**class** espressopp.analysis.LBOutputVzOfX (system, lb)

#### Parameters

- **system** (*shared\_ptr*) – system object defined earlier in the python-script
- **lb** (*lb\_object*) – lattice boltzmann object defined earlier in the python-script

Example:



```
>>> # initialise output of the Vz as a function of x-coordinate
>>> outputVzOfX = espressopp.analysis.LBOutputVzOfX(system, lb)
>>>
>>> # initialise external analysis object with previously created output object
>>> # and periodicity of invocation (steps):
>>> extAnalysis = espressopp.integrator.ExtAnalyze(outputVzOfX, 100)
>>>
>>> # add the external analysis object as an extension to the integrator
>>> integrator.addExtension( extAnalysis )
```

### espressopp.analysis.OrderParameter

`espressopp.analysis.OrderParameter` (*system*, *cutoff*, *angular\_momentum*,  
*do\_cluster\_analysis*, *include\_surface\_particles*,  
*ql\_low*, *ql\_high*)

#### Parameters

- **system** –
- **cutoff** –
- **angular\_momentum** (*int*) – (default: 6)
- **do\_cluster\_analysis** – (default: False)
- **include\_surface\_particles** – (default: False)
- **ql\_low** – (default: -1.0)
- **ql\_high** (*real*) – (default: 1.0)

### espressopp.analysis.ParticleRadiusDistribution

`espressopp.analysis.ParticleRadiusDistribution` (*system*)

#### Parameters **system** –

### espressopp.analysis.PressureTensor

This class computes the pressure tensor of the system. It can be used as standalone class in python as well as in combination with the integrator extension ExtAnalyze.

Example of standalone Usage:

```
>>> pt = espressopp.analysis.PressureTensor(system)
>>> print "pressure tensor of current configuration = ", pt.compute()
```

or

```
>>> pt = espressopp.analysis.PressureTensor(system)
>>> for k in xrange(100):
>>>     integrator.run(100)
>>>     pt.performMeasurement()
>>> print "average pressure tensor = ", pt.getAverageValue()
```

Example of usage in integrator with ExtAnalyze:

```
>>> pt = espressopp.analysis.PressureTensor(system)
>>> extension_pt = espressopp.integrator.ExtAnalyze(pt, interval=100)
>>> integrator.addExtension(extension_pt)
>>> integrator.run(10000)
>>> pt_ave = pt.getAverageValue()
```

```
>>> print "average Pressure Tensor = ", pt_ave[:6]
>>> print "          std deviation = ", pt_ave[6:]
>>> print "number of measurements = ", pt.getNumberOfMeasurements()
```

The following methods are supported:

- **performMeasurement()** computes the pressure tensor and updates average and standard deviation
- **reset()** resets average and standard deviation to 0
- **compute()** computes the instant pressure tensor, return value: [xx, yy, zz, xy, xz, yz]
- **getAverageValue()** returns the average pressure tensor and the standard deviation, return value: [xx, yy, zz, xy, xz, yz, +-xx, +-yy, +-zz, +-xy, +-xz, +-yz]
- **getNumberOfMeasurements()** counts the number of measurements that have been computed (standalone or in integrator) does `_not_` include measurements that have been done using “compute()”

`espressopp.analysis.PressureTensor` (*system*)

Parameters **system** –

### `espressopp.analysis.PressureTensorLayer`

This class computes the pressure tensor of the system in layer `h0`. It can be used as standalone class in python as well as in combination with the integrator extension `ExtAnalyze`.

Example of standalone Usage:

```
>>> pt = espressopp.analysis.PressureTensorLayer(system, h0, dh)
>>> print "pressure tensor of current configuration = ", pt.compute()
```

or

```
>>> pt = espressopp.analysis.PressureTensorLayer(system)
>>> for k in xrange(100):
>>>     integrator.run(100)
>>>     pt.performMeasurement()
>>> print "average pressure tensor = ", pt.getAverageValue()
```

Example of usage in integrator with `ExtAnalyze`:

```
>>> pt = espressopp.analysis.PressureTensorLayer(system)
>>> extension_pt = espressopp.integrator.ExtAnalyze(pt, interval=100)
>>> integrator.addExtension(extension_pt)
>>> integrator.run(10000)
>>> pt_ave = pt.getAverageValue()
>>> print "average Pressure Tensor = ", pt_ave[:6]
>>> print "          std deviation = ", pt_ave[6:]
>>> print "number of measurements = ", pt.getNumberOfMeasurements()
```

The following methods are supported:

- **performMeasurement()** computes the pressure tensor and updates average and standard deviation
- **reset()** resets average and standard deviation to 0
- **compute()** computes the instant pressure tensor in layer `h0`, return value: [xx, yy, zz, xy, xz, yz]
- **getAverageValue()** returns the average pressure tensor and the standard deviation, return value: [xx, yy, zz, xy, xz, yz, +-xx, +-yy, +-zz, +-xy, +-xz, +-yz]
- **getNumberOfMeasurements()** counts the number of measurements that have been computed (standalone or in integrator) does `_not_` include measurements that have been done using “compute()”

`espressopp.analysis.PressureTensorLayer` (*system, h0, dh*)

Parameters

- **system** –
- **h0** –
- **dh** –

### espressopp.analysis.PressureTensorMultiLayer

This class computes the pressure tensor of the system in  $n$  layers. Layers are perpendicular to Z direction and are equidistant (distance is  $Lz/n$ ). It can be used as standalone class in python as well as in combination with the integrator extension ExtAnalyze.

Example of standalone Usage:

```
>>> pt = espressopp.analysis.PressureTensorMultiLayer(system, n, dh)
>>> for i in xrange(n):
>>>     print "pressure tensor in layer %d: %s" % (i, pt.compute())
```

or

```
>>> pt = espressopp.analysis.PressureTensorMultiLayer(system, n, dh)
>>> for k in xrange(100):
>>>     integrator.run(100)
>>>     pt.performMeasurement()
>>> for i in xrange(n):
>>>     print "average pressure tensor in layer %d: %s" % (i, pt.compute())
```

Example of usage in integrator with ExtAnalyze:

```
>>> pt = espressopp.analysis.PressureTensorMultiLayer(system, n, dh)
>>> extension_pt = espressopp.integrator.ExtAnalyze(pt, interval=100)
>>> integrator.addExtension(extension_pt)
>>> integrator.run(10000)
>>> pt_ave = pt.getAverageValue()
>>> for i in xrange(n):
>>>     print "average Pressure Tensor = ", pt_ave[i][:6]
>>>     print "          std deviation = ", pt_ave[i][6:]
>>> print "number of measurements = ", pt.getNumberOfMeasurements()
```

The following methods are supported:

- **performMeasurement()** computes the pressure tensor and updates average and standard deviation
- **reset()** resets average and standard deviation to 0
- **compute()** computes the instant pressure tensor in  $n$  layers, return value: [xx, yy, zz, xy, xz, yz]
- **getAverageValue()** returns the average pressure tensor and the standard deviation, return value: [xx, yy, zz, xy, xz, yz, +-xx, +-yy, +-zz, +-xy, +-xz, +-yz]
- **getNumberOfMeasurements()** counts the number of measurements that have been computed (standalone or in integrator) does `_not_` include measurements that have been done using “compute()”

`espressopp.analysis.PressureTensorMultiLayer` (*system*, *n*, *dh*)

#### Parameters

- **system** –
- **n** –
- **dh** –

### espressopp.analysis.Temperature

Calculate the temperature of the system (in  $k_B T$  units).

`espressopp.analysis.Temperature` (*system*)

**Parameters** `system` (*shared\_ptr*) – system object

**Returns** temperature

**Return type** `real`

Temperature of the system of  $N$  particles is calculated as:

$$T = \frac{1}{N_f} \sum_{i=1}^N m_i v_i^2,$$

where  $m_i$  and  $v_i$  are the mass and velocity of a particle  $i$ .

$N_f = 3N$  is the number of the system’s degrees of freedom.

**Example:**

```
>>> # declare an object, e.g., T:
>>> T = espressopp.analysis.Temperature(system)
>>>
>>> # later in your script compute temperature and print it:
>>> print T.compute()
```

## espressopp.analysis.Test

Test class for any analysis tool.

`espressopp.analysis.Test` (*system*)

**Parameters** `system` –

### Details

This abstract base class provides the interface and some basic functionality for classes that do analysis or observable measurements

It provides the following methods:

`espressopp.analysis.AnalysisBase.compute()`

Computes the instant value of the observable.

**Return type** a python list or a scalar

`espressopp.analysis.AnalysisBase.getAverageValue()`

Returns the average value for the observable and the standard deviation.

**Return type** a python list

`espressopp.analysis.AnalysisBase.getNumberOfMeasurements()`

counts the number of measurements that have been performed (standalone or in integrator) does `_not_` include measurements that have been done using “`compute()`”

**Return type**

`espressopp.analysis.AnalysisBase.performMeasurement()`

Computes the observable and updates average and standard deviation

**Return type**

`espressopp.analysis.AnalysisBase.reset()`

Resets average and standard deviation

**Return type**

## espressopp.analysis.Autocorrelation

`espressopp.analysis.Autocorrelation(system)`

**Parameters** `system` –

`espressopp.analysis.Autocorrelation.clear()`

**Return type**

`espressopp.analysis.Autocorrelation.compute()`

**Return type**

`espressopp.analysis.Autocorrelation.gather(value)`

**Parameters** `value` –

**Return type**

## espressopp.analysis.CMVelocity

Compute and reset (set to zero) the center-of-mass (CM) velocity of the system.

**class** `espressopp.analysis.CMVelocity(system)`

**Parameters** `system` (`espressopp.System`) – system object

---

**Note:** `CMVelocity` can be attached to the *integrator*. In this case the `reset()` method is called, so you will reset the CM-velocity every *n*-th steps.

---

### Methods

**compute()**

Compute the CM-velocity of the system

**Return type** `Real3D`

**reset()**

Reset (set to zero) the CM-velocity of the system. Done by computing the CM-velocity of the system and subtracting it then from every particle.

**Return type** `void`

Example of resetting velocity

```
>>> total_velocity = espressopp.analysis.CMVelocity(system)
>>> total_velocity.reset()
```

Example of attaching to integrator

```
>>> # This extension can be attached to integrator
>>> # and run `reset()` every `n-th` steps.
>>> total_velocity = espressopp.analysis.CMVelocity(system)
>>> ext_remove_com = espressopp.integrator.ExtAnalyze(total_velocity, 10)
>>> integrator.addExtension(ext_remove_com)
```

## espressopp.analysis.ConfigsParticleDecomp

`espressopp.analysis.ConfigsParticleDecomp(system)`

**Parameters** `system` –

`espressopp.analysis.ConfigsParticleDecomp.clear()`

**Return type**

`espressopp.analysis.ConfigsParticleDecomp.compute()`

**Return type**

`espressopp.analysis.ConfigsParticleDecomp.gather()`

**Return type**

`espressopp.analysis.ConfigsParticleDecomp.gatherFromFile(filename)`

**Parameters filename –**

**Return type**

## espressopp.analysis.Configurations

- `gather()` add configuration to trajectory
- `clear()` clear trajectory
- `back()` get last configuration of trajectory
- `capacity` maximum number of configurations in trajectory further adding (`gather()`) configurations results in erasing oldest configuration before adding new one `capacity=0` means: infinite capacity (until memory is full)
- `size` number of stored configurations

usage:

storing trajectory

```
>>> configurations = espressopp.Configurations(system)
>>> configurations.gather()
>>> for k in xrange(100):
>>>     integrator.run(100)
>>> configurations.gather()
```

accessing trajectory data:

iterate over all stored configurations:

```
>>> for conf in configurations:
```

iterate over all particles stored in configuration:

```
>>> for pid in conf
>>>     particle_coords = conf[pid]
>>>     print pid, particle_coords
```

access particle with id <pid> of stored configuration <n>:

```
>>> print "particle coord: ", configurations[n][pid]
```

`espressopp.analysis.Configurations(system)`

**Parameters system –**

`espressopp.analysis.Configurations.back()`

**Return type**

`espressopp.analysis.Configurations.clear()`

**Return type**

`espressopp.analysis.Configurations.gather()`

**Return type**

## espressopp.analysis.ConfigurationsExt

- *gather()* add configuration to trajectory
- *clear()* clear trajectory
- *back()* get last configuration of trajectory
- *capacity* maximum number of configurations in trajectory further adding (*gather()*) configurations results in erasing oldest configuration before adding new one capacity=0 means: infinite capacity (until memory is full)
- *size* number of stored configurations

usage:

storing trajectory

```
>>> configurations = espressopp.ConfigurationsExt(system)
>>> configurations.gather()
>>> for k in xrange(100):
>>>     integrator.run(100)
>>>     configurations.gather()
```

accessing trajectory data:

iterate over all stored configurations:

```
>>> for conf in configurations:
```

iterate over all particles stored in configuration:

```
>>> for pid in conf
>>>     particle_coords = conf[pid]
>>>     print pid, particle_coords
```

access particle with id <pid> of stored configuration <n>:

```
>>> print "particle coord: ", configurations[n][pid]
```

`espressopp.analysis.ConfigurationsExt` (*system*)

**Parameters** *system* –

`espressopp.analysis.ConfigurationsExt.back()`

**Return type**

`espressopp.analysis.ConfigurationsExt.clear()`

**Return type**

`espressopp.analysis.ConfigurationsExt.gather()`

**Return type**

## espressopp.analysis.Energy

`espressopp.analysis.EnergyPot` (*system*, *per\_atom*)

**Parameters**

- *system* –
- *per\_atom* – (default: False)

`espressopp.analysis.EnergyPot.compute()`

**Return type**

`espressopp.analysis.EnergyKin (system, per_atom)`

#### Parameters

- **system** –
- **per\_atom** – (default: False)

`espressopp.analysis.EnergyKin.compute ()`

#### Return type

`espressopp.analysis.EnergyTot (system, per_atom)`

#### Parameters

- **system** –
- **per\_atom** – (default: False)

`espressopp.analysis.EnergyTot.compute ()`

#### Return type

### `espressopp.analysis.IntraChainDistSq`

`espressopp.analysis.IntraChainDistSq (system, fpl)`

#### Parameters

- **system** –
- **fpl** –

`espressopp.analysis.IntraChainDistSq.compute ()`

#### Return type

### `espressopp.analysis.MeanSquareDispl`

`espressopp.analysis.MeanSquareDispl (system, chainlength)`

#### Parameters

- **system** –
- **chainlength** – (default: None)

`espressopp.analysis.MeanSquareDispl.computeG2 ()`

#### Return type

`espressopp.analysis.MeanSquareDispl.computeG3 ()`

#### Return type

`espressopp.analysis.MeanSquareDispl.strange ()`

#### Return type

### `espressopp.analysis.MeanSquareInternalDist`

`espressopp.analysis.MeanSquareInternalDist (system, chainlength)`

#### Parameters

- **system** –
- **chainlength** –



```
espressopp.analysis.MeanSquareInternalDist.strange()
```

**Return type**

## espressopp.analysis.Observable

### Overview

List of classes based on Observable:

### espressopp.analysis.AdressDensity

Class to compute radial density profiles in adaptive resolution simulations based on distance to closest AdResS center. Works also for multiple overlapping AdResS regions.

Examples:

```
>>> densityprofile = espressopp.analysis.AdressDensity(system, verletlist)
>>> # creates the class
```

```
>>> densityprofile.addExclusions([1,2,3])
>>> # defines particle to be excluded from the calculation based on list of particle ids
```

```
>>> densityprofile.compute(100)
>>> # computes the densityprofile using 100 bins
```

```
espressopp.analysis.AdressDensity(system, verletlist)
```

**Parameters**

- **system** (*shared\_ptr<System>*) – system object
- **verletlist** (*shared\_ptr<VerletListAdress>*) – verletlist object

```
espressopp.analysis.AdressDensity.compute(bins)
```

**Parameters** **bins** (*int*) – number of bins

**Return type** list of reals

```
espressopp.analysis.AdressDensity.addExclusions(pidlist)
```

**Parameters** **pidlist** (*list of ints*) – list of ids of particles to be excluded from the calculation

### espressopp.analysis.CenterOfMass

```
espressopp.analysis.CenterOfMass(system)
```

**Parameters** **system** –

### espressopp.analysis.MaxPID

```
espressopp.analysis.MaxPID(system)
```

**Parameters** **system** –

## espressopp.analysis.NeighborFluctuation

`espressopp.analysis.NeighborFluctuation` (*system*, *radius*)

### Parameters

- **system** –
- **radius** –

## espressopp.analysis.NPart

`espressopp.analysis.NPart` (*system*)

### Parameters **system** –

## espressopp.analysis.PotentialEnergy

The object that computes potential energy of different interactions.

`espressopp.analysis.PotentialEnergy` (*system*, *potential*, *compute\_method=None*)

### Parameters

- **system** (`espressopp.System`) – The system object
- **interaction** (`espressopp.interaction.Interaction`) – The interaction object.
- **compute\_method** (*str*) – If set to *ALL* (default) then compute total potential energies, if set to *CG* then compute only coarse-grained part (if feasible), if set to *AT* then compute only atomistic part of potential energy.

## espressopp.analysis.Pressure

`espressopp.analysis.Pressure` (*system*)

### Parameters **system** –

## espressopp.analysis.RadialDistrF

`espressopp.analysis.RadialDistrF` (*system*)

### Parameters **system** –

`espressopp.analysis.RadialDistrF.compute` (*rdfN*)

### Parameters **rdfN** –

### Return type

## espressopp.analysis.RDFatomistic

Class to compute radial distribution functions in adaptive resolution simulations in subregions of the box.

Option 1 (`spanbased = True`): the RDF can be calculated in a cuboid region in the center of the box (periodic in y,z, limited in x). In this case, particle pairs are considered for which at least one of them is in the defined cuboid region. This can be useful when the high resolution region has a slab geometry. No further normalization should be required.

Option 2 (`spanbased = False`): the routine can also calculate unnormalized RDFs using particle pairs with both particles being in the high resolution region (based on the resolution value `lambda`, the `span` parameter is not used then). This can be useful when atomistic region has complicated or spherical geometries.

In any case, only pairs of atomistic particles belonging to two different coarse-grained particles are considered. Furthermore, note that the routine uses  $L_y / 2$  ( $L_y$  is the box length in y-direction) as the maximum distance for the RDF calculation, which is then binned according to  $rdfN$  during the computation. Hence,  $L_y$  should be the shortest box side (or, equally short as  $L_x$  and/or  $L_z$ ).

Examples:

```
>>> rdf_0_1 = espressopp.analysis.RDFatomistic(system = system, type1 = 0, type2 = 1, spanbased = True)
>>> # creates the class for calculating the RDF between atomistic particles of type 1 and 0 between
>>> # At least one of these particles has to be within plus/minus 1.5 from the center of the box
>>>
>>> rdf_0_1.compute(100)
>>> # computes the rdf using 100 bins over a distance corresponding to  $L_y / 2.0$ 
```

`espressopp.analysis.RDFatomistic(system, type1, type2, spanbased, span)`

#### Parameters

- **system** (*shared\_ptr<System>*) – system object
- **type1** (*int*) – type of atom 1
- **type2** (*int*) – type of atom 2
- **spanbased** (*bool*) – (default: True) If True, calculates RDFs in a cuboid region of radius `span` from the center (limited in x, periodic in y,z). If False, calculates RDFs with both particles being in the high resolution region (using `lambda` resolution values and ignoring `span` parameter).
- **span** (*real*) – (default: 1.0) +/- distance from centre of box in x-direction of the cuboid region used for RDF calculation if `spanbased == True`. If `spanbased == False`, this parameter is not used.

`espressopp.analysis.RDFatomistic.compute(rdfN)`

**Parameters** `rdfN` (*int*) – number of bins

**Return type** list of reals

### espressopp.analysis.StaticStructF

`espressopp.analysis.StaticStructF(system)`

**Parameters** `system` –

`espressopp.analysis.StaticStructF.compute(nqx, nqy, nqz, bin_factor, ofile)`

#### Parameters

- **nqx** –
- **nqy** –
- **nqz** –
- **bin\_factor** –
- **ofile** – (default: None)

#### Return type

`espressopp.analysis.StaticStructF.computeSingleChain(nqx, nqy, nqz, bin_factor, chainlength, ofile)`

#### Parameters

- **nqx** –
- **nqy** –
- **nqz** –

- **bin\_factor** –
- **chainlength** –
- **ofile** – (default: None)

**Return type**

### **espressopp.analysis.XDensity**

`espressopp.analysis.XDensity` (*system*)

**Parameters** *system* –

`espressopp.analysis.XDensity.compute` (*rdfN*)

**Parameters** *rdfN* –

**Return type**

### **espressopp.analysis.XPressure**

`espressopp.analysis.XPressure` (*system*)

**Parameters** *system* –

`espressopp.analysis.XPressure.compute` (*N*)

**Parameters** *N* –

**Return type**

### **espressopp.analysis.XTemperature**

`espressopp.analysis.XTemperature` (*system*)

**Parameters** *system* –

`espressopp.analysis.XTemperature.compute` (*N*)

**Parameters** *N* –

**Return type**

### **Details**

`espressopp.analysis.Observable.compute` ()

**Return type**

### **espressopp.analysis.SystemMonitor**

SystemMonitor prints and logs to file values obtained from Observables like temperature, pressure or potential energy.

`espressopp.analysis.SystemMonitor` (*system, integrator, output*)

**Parameters**

- **system** (`espressopp.System`) – The system object.
- **integrator** (`espressopp.integrator.MDIntegrator`) – The MD integrator.

- **output** (`espressopp.analysis.SystemMonitorOutputCSV`) – The output object.

`espressopp.analysis.SystemMonitor.add_observable(name, observable, is_visible)`

The function adds new observable to SystemMonitor.

#### Parameters

- **name** (*str*) – The name of observable
- **observable** – The observable, eg. `espressopp.analysis.PotentialEnergy`
- **is\_visible** (*bool*) – If set to True then values will be print on console.

`espressopp.analysis.SystemMonitor.info()`

The method print out on console the values of observables.

### CSV Output

The output of SystemMonitor to CSV files.

`espressopp.analysis.SystemMonitorOutputCSV(file_name, delimiter)`

#### Parameters

- **file\_name** (*str*) – The name of CSV file.
- **delimiter** (*str*) – The field delimiter, by default it is tabulator.

#### Example

```
>>> interaction = espressopp.interaction.VerletListLennardJones(verletlist)
>>> interaction.setPotential(type1=0, type2=0,
                             potential=espressopp.interaction.LennardJones(epsilon=1.0, sigma=1.0,
                                                                               cutoff=2.0))
>>> system_monitor_csv = espressopp.analysis.SystemMonitorOutputCSV('out.csv')
>>> system_monitor = espressopp.analysis.SystemMonitor(
    system, integrator, espressopp.analysis.SystemMonitorOutputCSV('out.csv'))
>>> system_monitor.add_observable('pot', espressopp.analysis.PotentialEnergy(system, interaction))
>>> ext_analysis = espressopp.integrator.ExtAnalyze(system_monitor, 10)
>>> integrator.addExtension(ext_analysis)
```

## espressopp.analysis.Velocities

`espressopp.analysis.Velocities(system)`

**Parameters** *system* –

`espressopp.analysis.Velocities.clear()`

**Return type**

`espressopp.analysis.Velocities.gather()`

**Return type**

## espressopp.analysis.VelocityAutocorrelation

`espressopp.analysis.VelocityAutocorrelation(system)`

**Parameters** *system* –

## espressopp.analysis.Viscosity

`espressopp.analysis.Viscosity` (*system*)

Parameters **system** –

`espressopp.analysis.Viscosity.compute` (*t0, dt, T*)

Parameters

- **t0** –
- **dt** –
- **T** –

Return type

`espressopp.analysis.Viscosity.gather` ()

Return type

## bc

### espressopp.bc.BC

#### Overview

---

*espressopp.bc.OrthorhombicBC*  
*espressopp.bc.SlabBC*

---

#### Details

This is the abstract base class for all boundary condition objects. It cannot be used directly. All derived classes implement at least the following methods:

**class** `espressopp.bc.BC`

**getFoldedPosition** (*pos, imageBox*)

Parameters

- **pos** –
- **imageBox** – (default: None)

Return type

**getMinimumImageVector** (*pos1, pos2*)

Parameters

- **pos1** –
- **pos2** –

Return type

**getRandomPos** ()

Return type

**getUnfoldedPosition** (*pos, imageBox*)

Parameters

- **pos** –
- **imageBox** –

#### Return type

*pos*, *pos1* and *pos2* are particle coordinates ( type: *(float, float, float)* ). *imageBox* ( type: *(int, int, int)* ) specifies the

### espressopp.bc.OrthorhombicBC

Like all boundary condition objects, this class implements all the methods of the base class **BC** , which are described in detail in the documentation of the abstract class **BC**.

The OrthorhombicBC class is responsible for the orthorhombic boundary condition. Currently only periodic boundary conditions are supported.

Example:

```
>>> boxsize = (Lx, Ly, Lz)
>>> bc = espressopp.bc.OrthorhombicBC(rng, boxsize)
```

```
espressopp.bc.OrthorhombicBC (rng, boxL)
```

#### Parameters

- **rng** –
- **boxL** (*real*) – (default: 1.0)

```
espressopp.bc.OrthorhombicBC.setBoxL (boxL)
```

#### Parameters **boxL** –

### espressopp.bc.SlabBC

Like all boundary condition objects, this class implements all the methods of the base class **BC** , which are described in detail in the documentation of the abstract class **BC**.

The SlabBC class is responsible for a cuboid boundary condition that is periodic in all but the “dir” dimension. Currently, dir is set arbitrarily to “0” (the x-direction).

Example:

```
>>> boxsize = (Lx, Ly, Lz)
>>> bc = espressopp.bc.SlabBC(rng, boxsize)
```

```
espressopp.bc.SlabBC (rng, boxL)
```

#### Parameters

- **rng** –
- **boxL** (*real*) – (default: 1.0)

```
espressopp.bc.SlabBC.setBoxL (boxL)
```

#### Parameters **boxL** –

## check

### espressopp.check.System

## esutil

### espressopp.esutil.Collectives

locate the node with `here=True` (e.g. indicating that data of a distributed storage is on the local node). This is a collective SPMD function.

`here` is a boolean value, which should be `True` on at most one node. Returns on the controller the number of the node with `here=True`, or an `KeyError` exception if no node had the item, i.e. had `here=True`.

```
espressopp.esutil.locateItem(here)
```

Parameters `here` –

### espressopp.esutil.GammaVariate

```
espressopp.esutil.GammaVariate(alpha, beta)
```

Parameters

- `alpha` –
- `beta` –

### espressopp.esutil.Grid

### espressopp.esutil.NormalVariate

```
espressopp.esutil.NormalVariate(mean, sigma)
```

Parameters

- `mean` (*real*) – (default: 0.0)
- `sigma` (*real*) – (default: 1.0)

### espressopp.esutil.RNG

### espressopp.esutil.UniformOnSphere

## external

## integrator

### espressopp.integrator.Address

The `AdResS` object is an extension to the integrator. It makes sure that the integrator also processes the atomistic particles and not only the CG particles. Hence, this object is of course only used when performing `AdResS` or `H-AdResS` simulations.

In detail the `AdResS` extension makes sure:



- that also the forces on the atomistic particles are initialized and set to by `Adress::initForces`
- that also the atomistic particles are integrated and propagated by `Adress::integrate1` and `Adress::integrate2`

Example - how to turn on the AdResS integrator extension:

```
>>> adress = espressopp.integrator.Address(system, verletlist, fixedtuplelist)
>>> integrator.addExtension(adress)
```

If `KTI` is set to `True`, then the resolution parameters are not updated. This can be used for example for Kirkwood thermodynamic integration, during which one manually sets the whole system on different resolution parameters. `KTI = True` then prevents overwriting these manually set values. Furthermore, when having moving AdResS regions based on particles, `regionupdates` specifies the update frequency of the AdResS region in number of steps (or, to be more precise, calls of `communicateAdrPositions()`). Note that there is a tradeoff: The more frequently the AdResS region is updated, the more gradually and accurately the AdResS region changes and adapts its shape. This could allow for a smaller overall AdResS region and possibly a smoother simulation. However, when having many AdResS region defining particles, these frequent updates can become computationally significant and cost additional simulation time. The optimum is highly system and application dependent.

```
class espressopp.integrator.Address(_system, _verletlist, _fixedtuplelist, KTI = False, re-
                                   gionupdates = 1)
```

#### Parameters

- **\_system** (*shared\_ptr<System>*) – system object
- **\_verletlist** (*shared\_ptr<VerletListAdress>*) – verletlist object
- **\_fixedtuplelist** (*shared\_ptr<FixedTupleListAdress>*) – fixedtuplelist object
- **KTI** (*bool*) – update resolution parameter? (Yes: set `False`, No: set `True`)
- **regionupdates** (*int*) – after how many steps does the AdResS region needs to be updated?

## espressopp.integrator.AssociationReaction

## espressopp.integrator.BerendsenBarostat

This is the Berendsen barostat implementation according to the original paper [Berendsen84]. If Berendsen barostat is defined (as a property of integrator) then at the each run the system size and the particle coordinates will be scaled by scaling parameter  $\mu$  according to the formula:

$$\mu = [1 - \Delta t / \tau (P_0 - P)]^{1/3}$$

where  $\Delta t$  - integration timestep,  $\tau$  - time parameter (coupling parameter),  $P_0$  - external pressure and  $P$  - instantaneous pressure.

Example:

```
>>> berendsenP = espressopp.integrator.BerendsenBarostat(system)
>>> berendsenP.tau = 0.1
>>> berendsenP.pressure = 1.0
>>> integrator.addExtension(berendsenP)
```

**!IMPORTANT** In order to run *npt* simulation one should separately define thermostat as well (e.g. Berendsen-Thermostat).

Definition:

In order to define the Berendsen barostat

```
>>> berendsenP = espressopp.integrator.BerendsenBarostat(system)
```

one should have the System defined.

Properties:

- *berendsenP.tau*

The property ‘tau’ defines the time parameter  $\tau$ .

- *berendsenP.pressure*

The property ‘pressure’ defines the external pressure  $P_0$ .

Setting the integration property:

```
>>> integrator.addExtension(berendsenP)
```

It will define Berendsen barostat as a property of integrator.

One more example:

```
>>> berendsen_barostat = espressopp.integrator.BerendsenBarostat(system)
>>> berendsen_barostat.tau = 10.0
>>> berendsen_barostat.pressure = 3.5
>>> integrator.addExtension(berendsen_barostat)
```

Canceling the barostat:

If one do not need the pressure regulation in system anymore or need to switch the ensemble or whatever :)

```
>>> # define barostat with parameters
>>> berendsen = espressopp.integrator.BerendsenBarostat(system)
>>> berendsen.tau = 0.8
>>> berendsen.pressure = 15.0
>>> integrator.addExtension(berendsen)
>>> ...
>>> # some runs
>>> ...
>>> # disconnect Berendsen barostat
>>> berendsen.disconnect()
>>> # the next runs will not include the system size and particle coordinates scaling
```

Connecting the barostat back after the disconnection

```
>>> berendsen.connect()
```

References:

`espressopp.integrator.BerendsenBarostat(system)`

**Parameters** `system` –

## espressopp.integrator.BerendsenBarostatAnisotropic

#TODO fix these comments This is the Berendsen barostat implementation according to the original paper [Berendsen84]. If Berendsen barostat is defined (as a property of integrator) then at the each run the system size and the particle coordinates will be scaled by scaling parameter  $\mu$  according to the formula:

$$\mu = [1 - \Delta t / \tau (P_0 - P)]^{1/3}$$

where  $\Delta t$  - integration timestep,  $\tau$  - time parameter (coupling parameter),  $P_0$  - external pressure and  $P$  - instantaneous pressure.

Example:

```
>>> berendsenP = espressopp.integrator.BerendsenBarostatAnisotropic(system)
>>> berendsenP.tau = 0.1
>>> berendsenP.pressure = 1.0
>>> integrator.addExtension(berendsenP)
```

**!IMPORTANT** In order to run *npt* simulation one should separately define thermostat as well (e.g. Berendsen-Thermostat).

Definition:

In order to define the Berendsen barostat

```
>>> berendsenP = espressopp.integrator.BerendsenBarostatAnisotropic(system)
```

one should have the System defined.

Properties:

- *berendsenP.tau*

The property ‘tau’ defines the time parameter  $\tau$ .

- *berendsenP.pressure*

The property ‘pressure’ defines the external pressure  $P_0$ .

Setting the integration property:

```
>>> integrator.addExtension(berendsenP)
```

It will define Berendsen barostat as a property of integrator.

One more example:

```
>>> berendsen_barostat = espressopp.integrator.BerendsenBarostatAnisotropic(system)
>>> berendsen_barostat.tau = 10.0
>>> berendsen_barostat.pressure = 3.5
>>> integrator.addExtension(berendsen_barostat)
```

Canceling the barostat:

If one do not need the pressure regulation in system anymore or need to switch the ensemble or whatever :)

```
>>> # define barostat with parameters
>>> berendsen = espressopp.integrator.BerendsenBarostatAnisotropic(system)
>>> berendsen.tau = 0.8
>>> berendsen.pressure = 15.0
>>> integrator.addExtension(berendsen)
>>> ...
>>> # some runs
>>> ...
>>> # disconnect Berendsen barostat
>>> berendsen.disconnect()
>>> # the next runs will not include the system size and particle coordinates scaling
```

Connecting the barostat back after the disconnection

```
>>> berendsen.connect()
```

`espressopp.integrator.BerendsenBarostatAnisotropic(system)`

**Parameters** `system` –

## espressopp.integrator.BerendsenThermostat

This is the Berendsen thermostat implementation according to the original paper [Berendsen84]. If Berendsen thermostat is defined (as a property of integrator) then at the each run the system size and the particle coordinates will be scaled by scaling parameter  $\lambda$  according to the formula:

$$\lambda = [1 + \Delta t / \tau_T (T_0 / T - 1)]^{1/2}$$

where  $\Delta t$  - integration timestep,  $\tau_T$  - time parameter (coupling parameter),  $T_0$  - external temperature and  $T$  - instantaneous temperature.

Example:

```
>>> berendsenT = espressopp.integrator.BerendsenThermostat(system)
>>> berendsenT.tau = 1.0
>>> berendsenT.temperature = 1.0
>>> integrator.addExtension(berendsenT)
```

Definition:

In order to define the Berendsen thermostat

```
>>> berendsenT = espressopp.integrator.BerendsenThermostat(system)
```

one should have the System defined.

Properties:

- *berendsenT.tau*

The property 'tau' defines the time parameter  $\tau_T$ .

- *berendsenT.temperature*

The property 'temperature' defines the external temperature  $T_0$ .

Setting the integration property:

```
>>> integrator.addExtension(berendsenT)
```

It will define Berendsen thermostat as a property of integrator.

One more example:

```
>>> berendsen_thermostat = espressopp.integrator.BerendsenThermostat(system)
>>> berendsen_thermostat.tau = 0.1
>>> berendsen_thermostat.temperature = 3.2
>>> integrator.addExtension(berendsen_thermostat)
```

Canceling the thermostat:

```
>>> # define thermostat with parameters
>>> berendsen = espressopp.integrator.BerendsenThermostat(system)
>>> berendsen.tau = 2.0
>>> berendsen.temperature = 5.0
>>> integrator.addExtension(berendsen)
>>> ...
>>> # some runs
>>> ...
>>> # disconnect Berendsen thermostat
>>> berendsen.disconnect()
```

Connecting the thermostat back after the disconnection

```
>>> berendsen.connect()
```

`espressopp.integrator.BerendsenThermostat` (*system*)

Parameters **system** –

## espressopp.integrator.CapForce

This class can be used to forcecap all particles or a group of particles. Force capping means that the force vector of a particle is rescaled so that the length of the force vector is  $\leq$  capforce

Example Usage:

```
>>> capforce = espressopp.integrator.CapForce(system, 1000.0)
>>> integrator.addExtension(capForce)
```

CapForce can also be used to forcecap only a group of particles:

```
>>> particle_group = [45, 67, 89, 103]
>>> capforce = espressopp.integrator.CapForce(system, 1000.0, particle_group)
>>> integrator.addExtension(capForce)
```

`espressopp.integrator.CapForce(system, capForce, particleGroup)`

#### Parameters

- **system** –
- **capForce** –
- **particleGroup** – (default: None)

## espressopp.integrator.DPDThermostat

`espressopp.integrator.DPDThermostat(system, vl)`

#### Parameters

- **system** –
- **vl** –

## espressopp.integrator.EmptyExtension

`espressopp.integrator.EmptyExtension(system)`

#### Parameters

- **system** –

## espressopp.integrator.ExtAnalyze

This class can be used to execute nearly all analysis objects within the main integration loop which allows to automatically accumulate time averages (with standard deviation error bars).

Example Usage:

```
>>> pt = espressopp.analysis.PressureTensor(system)
>>> extension_pt = espressopp.integrator.ExtAnalyze(pt, interval=100)
>>> integrator.addExtension(extension_pt)
>>> integrator.run(10000)
>>>
>>> pt_ave = pt.getAverageValue()
>>> print "average Pressure Tensor = ", pt_ave[:6]
>>> print "          std deviation = ", pt_ave[6:]
>>> print "number of measurements = ", pt.getNumberOfMeasurements()
```

`espressopp.integrator.ExtAnalyze(action_obj, interval)`

#### Parameters

- **action\_obj** –
- **interval** (*int*) – (default: 1)

## espressopp.integrator.Extension

`espressopp.integrator.Extension.connect()`

**Return type**

`espressopp.integrator.Extension.disconnect()`

**Return type**

## espressopp.integrator.ExtForce

`espressopp.integrator.ExtForce(system, extForce, particleGroup)`

**Parameters**

- **system** –
- **extForce** –
- **particleGroup** – (default: None)

## espressopp.integrator.FixPositions

`espressopp.integrator.FixPositions(system, particleGroup, fixMask)`

**Parameters**

- **system** –
- **particleGroup** –
- **fixMask** –

## espressopp.integrator.FreeEnergyCompensation

Free Energy Compensation used in Hamiltonian Adaptive Resolution Simulations (H-AdResS). This works for spherical or slab adaptive resolution geometries. However, it only works for fixed, non-moving atomistic region (otherwise, H-AdResS is not properly defined).

Example:

```
>>> fec = espressopp.integrator.FreeEnergyCompensation(system, center=[Lx/2, Ly/2, Lz/2])
>>> # set up the fec module with the center in the center of the box
>>> fec.addForce(itype=3, filename="tablefec.svg", type=typeCG)
>>> # set up the actual force
>>> integrator.addExtension(fec)
>>> # add to previously defined integrator
```

`espressopp.integrator.FreeEnergyCompensation(system, center, sphereAdr)`

**Parameters**

- **system** (*shared\_ptr<System>*) – system object
- **center** (*list of reals*) – (default: [], corresponds to (0.0, 0.0, 0.0) position) center of high resolution region
- **sphereAdr** (*bool*) – (default: False) Spherical AdResS region (True) vs. slab geometry with resolution change in x-direction (False)

`espressopp.integrator.FreeEnergyCompensation.addForce(itype, filename, type)`

**Parameters**

- **itype** (*int*) – interpolation type 1: linear, 2: Akima, 3: Cubic

- **filename** (*string*) – filename for TD force file
- **type** (*int*) – particle type on which the TD force needs to be applied

`espressopp.integrator.FreeEnergyCompensation.computeCompEnergy()`

**Return type** `real`

## espressopp.integrator.GeneralizedLangevinThermostat

`espressopp.integrator.GeneralizedLangevinThermostat(system)`

**Parameters** `system` –

`espressopp.integrator.GeneralizedLangevinThermostat.addCoeffs` (*itype*, *filename*, *type*)

**Parameters**

- **itype** –
- **filename** –
- **type** –

**Return type**

## espressopp.integrator.Isokinetic

`espressopp.integrator.Isokinetic(system)`

**Parameters** `system` –

## espressopp.integrator.LangevinBarostat

This is the barostat implementation to perform Langevin dynamics in a Hoover style extended system according to the paper [Quigley04]. It includes corrections of Hoover approach which were introduced by Martyna et al [Martyna94]. If LangevinBarostat is defined (as a property of integrator) the integration equations will be modified. The volume of system  $V$  is introduced as a dynamical variable:

$$\dot{\mathbf{r}}_i = \frac{\mathbf{p}_i}{m_i} + \frac{p_\epsilon}{W} \mathbf{r}_i$$

$$\dot{\mathbf{p}}_i = -\nabla_{\mathbf{r}_i} \Phi - \left(1 + \frac{n}{N_f}\right) \frac{p_\epsilon}{W} \mathbf{p}_i - \gamma \mathbf{p}_i + \mathbf{R}_i$$

$$\dot{V} = dV p_\epsilon / W$$

$$\dot{p}_\epsilon = nV(X - P_{ext}) + \frac{n}{N_f} \sum_{i=1}^N \frac{\mathbf{p}_i^2}{m_i} - \gamma_p p_\epsilon + R_p$$

where volume has a fictitious mass  $W$  and associated momentum  $p_\epsilon$ ,  $\gamma_p$  - friction coefficient,  $P_{ext}$  - external pressure and  $X$  - instantaneous pressure without white noise contribution from thermostat,  $n$  - dimension,  $N_f$  - degrees of freedom (if there are no constraints and  $N$  is the number of particles in system  $N_f = nN$ ).  $R_p$  - values which are drawn from Gaussian distribution of zero mean and unit variance scaled by

$$\sqrt{\frac{2k_B T W \gamma_p}{\Delta t}}$$

**!IMPORTANT** Terms  $-\gamma \mathbf{p}_i + \mathbf{R}_i$  correspond to the thermostat. They are not included here and will not be calculated if the Langevin Thermostat is not defined.

Example:

```
>>> rng = espressopp.esutil.RNG()
>>> langevinP = espressopp.integrator.LangevinBarostat(system, rng, desiredTemperature)
>>> langevinP.gammaP = 0.05
>>> langevinP.pressure = 1.0
>>> langevinP.mass = pow(10.0, 4)
>>> integrator.addExtension(langevinP)
```

**!IMPORTANT** This barostat is supposed to be run in a couple with thermostat in order to simulate the *npt* ensemble, because the term  $R_p$  needs the temperature as a parameter.

Definition:

In order to define the Langevin-Hoover barostat

```
>>> langevinP = espressopp.integrator.LangevinBarostat(system, rng, desiredTemperature)
```

one should have the System and *RNG* defined and know the desired temperature.

Properties:

- *langevinP.gammaP*  
The property 'gammaP' defines the friction coefficient  $\gamma_p$ .
- *langevinP.pressure*  
The property 'pressure' defines the external pressure  $P_{ext}$ .
- *langevinP.mass*  
The property 'mass' defines the fictitious mass  $W$ .

Methods:

- *setMassByFrequency( frequency )*  
Set the proper *langevinP.mass* using expression  $W = dNk_bT/\omega_b^2$ , where frequency,  $\omega_b$ , is the frequency of required volume fluctuations. The value of  $\omega_b$  should be less then the lowest frequency which appears in the NVT temperature spectrum [Quigley04] in order to match the canonical distribution.  $d$  - dimensions,  $N$  - number of particles,  $k_b$  - Boltzmann constant,  $T$  - desired temperature.

**NOTE** The *langevinP.mass* can be set both directly and using the (*setMassByFrequency( frequency )*)

Adding to the integration:

```
>>> integrator.addExtension(langevinP)
```

It will define Langevin-Hoover barostat as a property of integrator.

One more example:

```
>>> rngBaro = espressopp.esutil.RNG()
>>> lP = espressopp.integrator.LangevinBarostat(system, rngBaro, desiredTemperature)
>>> lP.gammaP = .5
>>> lP.pressure = 1.0
>>> lP.mass = pow(10.0, 5)
>>> integrator.addExtension(lP)
```

Canceling the barostat:

If one do not need the pressure regulation in system anymore or need to switch the ensemble or whatever :)

```
>>> # define barostat with parameters
>>> rngBaro = espressopp.esutil.RNG()
>>> lP = espressopp.integrator.LangevinBarostat(system, rngBaro, desiredTemperature)
>>> lP.gammaP = .5
>>> lP.pressure = 1.0
>>> lP.mass = pow(10.0, 5)
```



```
>>> integrator.langevinBarostat = 1P
>>> ...
>>> # some runs
>>> ...
>>> # disconnect barostat
>>> langevinBarostat.disconnect()
>>> # the next runs will not include the modification of integration equations
```

Connecting the barostat back after the disconnection

```
>>> langevinBarostat.connect()
```

References:

`espressopp.integrator.LangevinBarostat` (*system*, *rng*, *temperature*)

**Parameters**

- **system** –
- **rng** –
- **temperature** –

## espressopp.integrator.LangevinThermostat

Langevin Thermostat

Example:

```
>>> langevin = espressopp.integrator.LangevinThermostat(system)
>>> # set up the thermostat
>>> langevin.gamma = gamma
>>> # set friction coefficient gamma
>>> langevin.temperature = temp
>>> # set temperature
>>> langevin.adress = True
>>> # set adress (default is False)
>>> integrator.addExtension(langevin)
>>> # add extensions to a previously defined integrator
```

`espressopp.integrator.LangevinThermostat` (*system*)

**Parameters** **system** (*shared\_ptr<System>*) – system object

`espressopp.integrator.LangevinThermostat.addExclusions` (*pidlist*)

**Parameters** **pidlist** (*list of ints*) – list of particle ids to be excluded from thermostating. In adaptive (AdResS) simulations, add ids of atomistic particles to be excluded (thermostats acts in this case on atomistic level). For normal simulations, add normal or coarse-grained particle ids.

## espressopp.integrator.LangevinThermostat1D

`espressopp.integrator.LangevinThermostat1D` (*system*)

**Parameters** **system** –

## espressopp.integrator.LangevinThermostatHybrid

As `LangevinThermostat`, but for use in AdResS systems, to allow the application of different thermostat friction constants ( $\gamma$ ) to different AdResS regions. Uses three values of  $\gamma$ , one for the atomistic region, one for the hybrid region, and one for the coarse-grained region.

```
>>> # create FixedTupleList object
>>> ftpl = espressopp.FixedTupleListAdress(system.storage)
>>> ftpl.addTuples(tuples)
>>> system.storage.setFixedTuplesAdress(ftpl)
>>>
>>> system.storage.decompose()
>>>
>>> # create Langevin thermostat
>>> thermostat = espressopp.integrator.LangevinThermostatHybrid(system, ftpl)
>>>
>>> # set Langevin friction constants
>>> thermostat.gamma = 0.0 # units = 1/timeunit
>>> print "# gamma for atomistic region for langevin thermostat = ", thermostat.gamma
>>> thermostat.gammahy = 10.0 # units = 1/timeunit
>>> print "# gamma for hybrid region for langevin thermostat = ", thermostat.gammahy
>>> thermostat.gammacg = 10.0 # units = 1/timeunit
>>> print "# gamma for coarse-grained region for langevin thermostat = ", thermostat.gammacg
>>>
>>> # set temperature of thermostat
>>> thermostat.temperature = kBT
>>> # kBT is a float with the value of temperature in reduced units, i.e. temperature * Boltzmann
```

No need to include the line

```
>>> thermostat.adress = True
```

as is necessary in the case of the basic LangevinThermostat, because LangevinThermostatHybrid is always only used in AdResS systems

## espressopp.integrator.LangevinThermostatOnGroup

Thermalize particles in the ParticleGroup only.

`espressopp.integrator.LangevinThermostatOnGroup(system, particle_group)`

### Parameters

- **system** (`espressopp.System`) – The system object.
- **particle\_group** (`espressopp.ParticleGroup`) – The particle group.

### Example

```
>>> pg = espressopp.ParticleGroup(system.storage)
>>> for pid in range(10):
>>>     pg.add(pid)
>>> thermostat = espressopp.integrator.LangevinThermostatOnGroup(system, pg)
>>> thermostat.temperature = 1.0
>>> thermostat.gamma = 1.0
>>> integrator.addExtension(thermostat)
```

## espressopp.integrator.LangevinThermostatOnRadius

Langevin Thermostat for Radii of Particles

Example:

```
>>> radius_mass = mass
>>> # set virtual mass for dynamics of radius
>>> langevin = espressopp.integrator.LangevinThermostatOnRadius(system, radius_mass)
>>> # set up the thermostat
```

```
>>> langevin.gamma = gamma
>>> # set friction coefficient gamma
>>> langevin.temperature = temp
>>> # set temperature
>>> integrator.addExtension(langevin)
>>> # add extensions to a previously defined integrator
```

`espressopp.integrator.LangevinThermostatOnRadius` (*system, dampingmass*)

#### Parameters

- **system** –
- **\_dampingmass** –

`espressopp.integrator.LangevinThermostatOnRadius.addExclusions` (*pidlist*)

Parameters **pidlist** (*list of ints*) – list of particle ids to be excluded from thermostatting.

## espressopp.integrator.LatticeBoltzmann

### Overview

---

*espressopp.integrator.LatticeBoltzmann*

---

### Details

The LatticeBoltzmann (LB) class controls fluid hydrodynamics and allows for hybrid LB/MD simulations. It is implemented as `espressopp.integrator.Extension` in ESPResSo++.

**class** `espressopp.integrator.LatticeBoltzmann` (*system, nodeGrid, a = 1., tau = 1., numDims = 3, numVels = 19*)

#### Parameters

- **system** (*shared\_ptr*) – system object
- **nodeGrid** (*Int3D*) – arrangement of CPUs in space
- **a** (*real*) – lattice spacing (in lattice units).
- **tau** (*real*) – time discretization (in lattice units)
- **numDims** (*int*) – dimensionality of the LB model
- **numVels** (*int*) – number of velocity vectors in the LB model

#### Returns

lb object

The LB-fluid in ESPResSo++ is aiming at simulations of complex soft matter systems. They consist of MD particles (colloids, composite nanoparticles or polymer chains) that are solved in the LB-fluid preserving hydrodynamic interactions.

The default lattice model is D3Q19 (`numDims = 3, numVels = 19`) and both lattice spacing `a` and timestep `tau` are set to 1. If some other lattice model is needed feel free to modify the code: adding 3D ones is straightforward, for 2D cases one has to make more thorough changes.

The parameters of the LB-fluid are expected in Lennard-Jones (LJ) units. This strategy helps users with MD-background think of LB-fluid in term of LJ liquid. One only has to specify its properties such as liquid density,  $\rho$ , temperature,  $T$ , and viscosity,  $\eta$ .

---

**Note:** Standard LJ fluid can be characterized by  $\rho \sim 1[\sigma^{-3}]$ ,  $T \sim 1[\epsilon]$ , and  $\eta \sim 5[\epsilon\tau/\sigma^3]$

---

### Example

```
>>> L = 20
>>>
>>> # create cubic box
>>> box = (L, L, L)
>>> # The rc+skin= lattice_size
>>> rc = 0.9
>>> skin= 0.1
>>>
>>> # initialize empty default system with the created cubic box.
>>> system, integrator = espressopp.standard_system.Default(box)
>>>
>>> # nodeGrid is determined based on the number of CPUs used for simulation among others
>>> nodeGrid=espressopp.tools.decomp.nodeGrid(espressopp.MPI.COMM_WORLD.size,box,rc,skin)
>>>
>>> # initialize lb object. The dimensions of the lattice are obtained from the
>>> # system's box dimensions employing lattice spacing 1.
>>> lb = espressopp.integrator.LatticeBoltzmann(system, nodeGrid)
```

### Methods

#### **getLBMom** (*node, moment*)

Get hydrodynamic moment from a specific node

##### Parameters

- **node** (*Int3D*) – node index
- **moment** (*int*) – hydrodynamic moment to get

Use 0 to get density  $\rho$  and 1-3 for mass flux components  $j_x$ ,  $j_y$  and  $j_z$ , correspondingly.

#### **setLBMom** (*node, moment, value*)

Set hydrodynamic moment for a specific node

##### Parameters

- **node** (*Int3D*) – node index
- **moment** (*int*) – hydrodynamic moment to set
- **value** (*real*) – value to set

#### **saveLBConf** ()

Dumps LB configuration with separate files for coupling forces, LB-fluid moments and populations (the last one is a bit overkill). The dump files are written for every CPU separately and are put in the *dump* folder

#### **keepLBDump** ()

Sets a flag to keep previously dumped LB configuration. Normally the previous dump is deleted after a new one is made.

### Example

```
>>> # set bulk viscosity
>>> for k in range (10):
>>>     integrator.run(50000)
>>>
>>> # output LB configuration
>>> lb.keepLBDump()           # flag to keep previously saved LB state
>>> lb.saveLBConf()          # saves current state of the LB fluid
```

### Properties

#### **Int3D nodeGrid**

Array of CPUs in space

### Example

```

>>> # it is advised to set nodeGrid by internal ESPResSo++ function
>>> # based on the number of CPUs
>>> nodeGrid=espressopp.tools.decomp.nodeGrid(espressopp.MPI.COMM_WORLD.size,box,rc,skin)

```

```

real a = 1.
    Lattice spacing (lattice units)

real tau = 1.
    Lattice time step (lattice units)

int numDims = 3
    Number of dimensions of the LB model (D3Q19)

int numVels = 19
    Number of velocity vectors of the LB model (D3Q19)

real visc_b
    Bulk viscosity (LJ units), affects gamma_b.

    Example

```

```

>>> # set bulk viscosity
>>> lb.visc_b = 5.

```

```

real visc_s
    Shear viscosity (LJ units), affects gamma_s.

    Example

```

```

>>> # set shear viscosity
>>> lb.visc_s = 5.

```

```

real gamma_b = 0.
    Bulk gamma (for experienced LB users)

real gamma_s = 0.
    Shear gamma (for experienced LB users)

real gamma_odd = 0.
    Odd gamma (for experienced LB users)

real gamma_even = 0.
    Even gamma (for experienced LB users)

real lbTemp = 0.
    Temperature of the LB fluid (LJ units)

    Example

```

```

>>> L = 20
>>> T = 1.
>>> N = 200
>>>
>>> # create cubic box
>>> box = (L, L, L)
>>> rc=0.9
>>> skin=0.1
>>>
>>> # initialize Lennard Jones system with the created cubic box and given temperature.
>>> system, integrator = espressopp.standard_system.LennardJones(N, box, temperature=T)
>>>
>>> # nodeGrid is determined based on the number of CPUs used for simulation
>>> nodeGrid=espressopp.tools.decomp.nodeGrid(espressopp.MPI.COMM_WORLD.size,box,rc,skin)
>>>
>>> # initialize lb object. The dimensions of the lattice are obtained from the
>>> # system's box dimensions employing lattice spacing 1.
>>> lb = espressopp.integrator.LatticeBoltzmann(system, nodeGrid)

```

```
>>>
>>> # set LB temperature to T
>>> lb.lbTemp = T
```

**real fricCoeff=5.**

Friction coefficient of the coupling (*LJ units*)

Example

```
>>> # set friction coefficient of the coupling
>>> lb.fricCoeff = 20.
```

**int nSteps = 1**

Timescale contrast (ratio) between LB and MD

Example

```
>>> # set time step contrast between LB and MD
>>> lb.nSteps = 10
```

**int profStep = 10000**

Frequency of time profiling

Example

```
>>> # set profiling frequency
>>> lb.profStep = 5000
```

**Int3D getMyNi**

Number of real and halo nodes for the CPU

## espressopp.integrator.LBInit

### Overview

<code>espressopp.integrator.LBInitPopUniform</code>
<code>espressopp.integrator.LBInitPopWave</code>
<code>espressopp.integrator.LBInitConstForce</code>
<code>espressopp.integrator.LBInitPeriodicForce</code>

### Details

This abstract class provides the interface to (re-)initialize populations and handle external forces.

**class** `espressopp.integrator.LBInit`

**createDenVel** (*rho0*, *u0*)

to set initial density and velocity of the LB-fluid.

#### Parameters

- **rho0** (*real*) – density
- **u0** (*Real3D*) – velocity

The following options for LB-fluid initialization are supported:

- `espressopp.integrator.LBInitPopUniform` A typical choice. It initializes uniformly distributed density and velocity: On every lattice site the density is *rho0* and velocity is *u0*
- `espressopp.integrator.LBInitPopWave` for uniform density at every lattice site, but harmonic velocity  $v_z(x)$  with the period of lattice sites in *x*-direction

**setForce** (*value*)

to set an external force onto LB-fluid.

**Parameters** *value* (`Real3D`) – value of the force

**addForce** (*force*)

to add a new external force to the existing one.

**Parameters** *force* (`Real3D`) – value of the force

Two main external force types are implemented:

- `espressopp.integrator.LBInitConstForce` to manage constant (gravity-like) forces acting on every lattice site and
- `espressopp.integrator.LBInitPeriodicForce` to manage periodic (sin-like) forces

### espressopp.integrator.LBInitPopUniform

This class creates LB-fluid with uniform density `rho0` and velocity `u0` (*lattice units*).

Example:

```
>>> # set initial density and velocity
>>> initDen = 1.
>>> initVel = Real3D( 0. )
>>>
>>> # create initPop object and initialize populations
>>> initPop = espressopp.integrator.LBInitPopUniform(system, lb)
>>> initPop.createDenVel( initDen, initVel )
```

### espressopp.integrator.LBInitPopWave

This class creates LB-fluid with uniform density and harmonic velocity (*lattice units*):

$v_x = 0, v_y = 0, v_z(i) = A \cdot \sin(2\pi \cdot i / N_x)$ , where  $A$  is the amplitude of the velocity wave,  $N_x$  is the number of lattice nodes in  $x$ -direction and  $i$  is the index of the node the velocity is calculated for.

This may be used to test the system: total moment is zero and the liquid tends to equilibrium, i.e. relaxes to a uniform zero velocity.

Example:

```
>>> # set initial density
>>> initDen = 1.
>>>
>>> # set initial velocity
>>> Vx = Vy = 0.
>>> ampVz = 0.0005
>>> initVel = Real3D( Vx, Vy, ampVz )
>>>
>>> # create initPop object and initialize populations
>>> initPop = espressopp.integrator.LBInitPopWave(system, lb)
>>> initPop.createDenVel( initDen, initVel )
```

### espressopp.integrator.LBInitConstForce

This class allows to set or add constant (gravity-like) external forces (*lattice units*) to the LB-fluid. At first, one has to create a force object and then set or add this force to the system.

Example to set external force:

```
>>> extForceToSet = Real3D(0., 0., 0.0005)
>>> lbforce = espressopp.integrator.LBInitConstForce(system, lb)
>>> lbforce.setForce( extForceToSet )
```

Example to add external force to the existing forces:

```
>>> extForceToAdd = Real3D(0.0001, 0., 0.)
>>> lbforce = espressopp.integrator.LBInitConstForce(system, lb)
>>> lbforce.addForce( extForceToAdd )
```

### espressopp.integrator.LBInitPeriodicForce

This class allows to set or add external periodic forces (*lattice units*) to the LB-fluid. At first, one has to create a force object and then set or add this force to the system.

---

**Note:** Please note, that a periodic (sin-like) force acts in  $z$ -direction as a function of  $x$ . The  $z$ -component of the force provides therefore the amplitude of the sin-modulation. The  $x$ - and  $y$ -components of the specified force interpreted as body forces in corresponding directions.

---

Example to set external sin-like force.

```
>>> ampFz = 0.0001
>>> Fx = Fy = 0.
>>> extForceToSet = Real3D( Fx, Fy, ampFz )
>>> lbforceSin = espressopp.integrator.LBInitConstForce(system, lb)
>>> lbforceSin.setForce( extForceToSet )
```

Example to add external sin-like force.

```
>>> ampFz = 0.0005
>>> Fx = Fy = 0.
>>> extForceToAdd = Real3D( Fx, Fy, ampFz )
>>> lbforceSin = espressopp.integrator.LBInitConstForce(system, lb)
>>> lbforceSin.addForce( extForceToAdd )
```

### espressopp.integrator.MDIntegrator

`espressopp.integrator.MDIntegrator.addExtension(extension)`

**Parameters** `extension` –

**Return type**

`espressopp.integrator.MDIntegrator.getExtension(k)`

**Parameters** `k` –

**Return type**

`espressopp.integrator.MDIntegrator.getNumberOfExtensions()`

**Return type**

`espressopp.integrator.MDIntegrator.run(niter)`

**Parameters** `niter` –

**Return type**



## espressopp.integrator.MinimizeEnergy

This is a very simple approach to perform energy minimization of the system. The module uses a [steepest descent method](#). The position of particles is updated following the equation:

$$p_{i+1} = p_i + \min(\gamma F_i, d_{max})$$

where  $p_{i+1}$  is a new position,  $p_i$  is a position at current step with corresponding force  $F_i$ . The parameters  $\gamma$  and  $d_{max}$  are set by user and control the relaxation of the energy and the maximum update of the coordinates per step.

Additionally, a variable  $\gamma$  step is also implemented. In this case, the position of particles is updated following the equation:

$$p_{i+1} = p_i + d_{max}/f_{max}F_i$$

where  $f_{max}$  is a maximum force in a single step of steepest descent method.  $\gamma = d_{max}/f_{max}$  is automatically adjusted to a force magnitude.

In both cases, the routine runs until the maximum force is bigger than  $f_{max}$  or for at most  $n$  steps.

**Please note** This module does not support any integrator extensions.

Example

```
>>> em = espressopp.integrator.MinimizeEnergy(system, gamma=0.001, ftol=0.01, max_displacement=0.0)
>>> em.run(10000)
```

Example

```
>>> em = espressopp.integrator.MinimizeEnergy(system, gamma=0.01, ftol=0.01, max_displacement=0.0)
>>> em.run(10000)
```

### API

`espressopp.integrator.MinimizeEnergy(system, gamma, ftol, max_displacement, variable_step_flag)`

#### Parameters

- **system** (`espressopp.System`) – The espressopp system object.
- **gamma** (`float`) – The gamma value.
- **ftol** (`float`) – The force tolerance
- **max\_displacement** (`float`) – The maximum displacement.
- **variable\_step\_flag** (`bool`) – The flag of adjusting gamma to the force strength.

`espressopp.integrator.MinimizeEnergy.run(max_steps, verbose)`

#### Parameters

- **max\_steps** (`int`) – The maximum number of steps to run.
- **verbose** (`bool`) – If set to True then display information about maximum force during the iterations.

**Returns** The true if the maximum force in the system is lower than ftol otherwise false.

**Return type** bool

`espressopp.integrator.MinimizeEnergy.f_max`  
The maximum force in the system.

`espressopp.integrator.MinimizeEnergy.displacement`  
The maximum displacement used during the run of MinimizeEnergy

`espressopp.integrator.MinimizeEnergy.step`  
The current iteration step.

## espressopp.integrator.OnTheFlyFEC

`espressopp.integrator.OnTheFlyFEC(system, center)`

### Parameters

- **system** –
- **center** – (default: [])

`espressopp.integrator.OnTheFlyFEC.getBins()`

### Return type

`espressopp.integrator.OnTheFlyFEC.getGap()`

### Return type

`espressopp.integrator.OnTheFlyFEC.getSteps()`

### Return type

`espressopp.integrator.OnTheFlyFEC.makeArrays()`

### Return type

`espressopp.integrator.OnTheFlyFEC.resetCounter()`

### Return type

`espressopp.integrator.OnTheFlyFEC.writeFEC()`

### Return type

## espressopp.integrator.Rattle

RATTLE algorithm for satisfying bond constraints and making the corresponding velocity corrections.

Refs:

Andersen, H. C. Rattle: A velocity version of the Shake algorithm for molecular dynamics calculations, J. Comp. Physics, 52, 24-34 (1983)

Allen & Tildesley, Computer Simulation of Liquids, OUP, 1987

RATTLE is implemented as an integrator extension, and takes as input a list of lists detailing, for each bond to be constrained: the indices of the two particles involved, the constraint distance, and the particle masses.

This implementation is intended for use with hydrogen-heavy atom bonds, which form isolated groups of constrained bonds, e.g NH<sub>2</sub> or CH<sub>3</sub> groups. The particle which participates in only one constrained bond (i.e. the hydrogen) should be listed first. The particle listed second (the heavy atom) may participate in more than one constrained bond. This implementation will not work if both particles participate in more than one constrained bond.

Note: At the moment, the RATTLE implementation only works if all atoms in an isolated group of rigid bonds are on the same CPU. This can be achieved by grouping all the particles using `DomainDecompositionAdress` and `FixedTupleListAdress`. The groups of rigid bonds can be identified using the dictionary `constrainedBondsDict` (see example below).

Note: The constraints are not taken into account in other parts of the code, such as temperature or pressure calculation.

Python example script for one methanol molecule where atoms are indexed in the order C H1 H2 H3 OH HO:

```
>>> # list for each constrained bond which lists: heavy atom index, light atom index, bond length
>>> constrainedBondsList = [[1, 2, 0.109, 12.011, 1.008], [1, 3, 0.109, 12.011, 1.008], [1, 4, 0.109, 12.011, 1.008], [2, 3, 0.109, 12.011, 1.008], [2, 4, 0.109, 12.011, 1.008], [3, 4, 0.109, 12.011, 1.008]]
>>> rattle = espressopp.integrator.Rattle(system, maxit = 1000, tol = 1e-6, rptol = 1e-6)
>>> rattle.addConstrainedBonds(constrainedBondsList)
>>> integrator.addExtension(rattle)
```

This list of lists of constrained bonds can be conveniently built using the espressopp tool `findConstrainedBonds`.

```
>>> # Automatically identify hydrogen-containing bonds among the particles whose indices are in t
>>> # pidlist - list of indices of particles in which to search for hydrogens (list of int)
>>> # masses - list of masses of all particles (list of real)
>>> # massCutoff - atoms with mass < massCutoff are identified as hydrogens (real)
>>> # bondtypes - dictionary (e.g. obtained using espressopp.gromacs.read()), key: bondtype (in
>>> # bondtypeparams - dictionary (e.g. obtained using espressopp.gromacs.read()), key: bondtyp
>>> hydrogenIDs, constrainedBondsDict, constrainedBondsList = espressopp.tools.findConstrainedBon
>>> # hydrogenIDs - list of indices of hydrogen atoms
>>> # constrainedBondsDict - dictionary mapping from a heavy atom to all the light atoms it is bo
>>> # constrainedBondsList - list of lists, constrained bonds for use with Rattle.addConstrainedB
>>> print "# found", len(hydrogenIDs), " hydrogens in the solute"
>>> print "# found", len(constrainedBondsDict), " heavy atoms involved in bonds to hydrogen"
>>> print "# will constrain", len(constrainedBondsList), " bonds using RATTLE"
```

`espressopp.integrator.Rattle` (*system*, *maxit* = 1000, *tol* = 1e-6, *rptol* = 1e-6)

#### Parameters

- **system** (`espressopp.System`) – espressopp system
- **maxit** (*int*) – maximum number of iterations
- **tol** (*real*) – tolerance for deciding if constraint distance and current distance are similar enough
- **rptol** (*real*) – tolerance for deciding if the angle between the bond vector at end of previous timestep and current vector has become too large

`espressopp.integrator.Rattle.addConstrainedBonds` (*bondDetailsLists*)

**Parameters** *bondDetailsLists* (*list of [int, int, real, real, real]*) – list of lists, each list contains pid of heavy atom, pid of light atom, constraint distance, mass of heavy atom, mass of light atom

## espressopp.integrator.Settle

`espressopp.integrator.Settle` (*system*, *fixedtuplelist*, *mO*, *mH*, *distHH*, *distOH*)

#### Parameters

- **system** –
- **fixedtuplelist** –
- **mO** (*real*) – (default: 16.0)
- **mH** (*real*) – (default: 1.0)
- **distHH** (*real*) – (default: 1.58)
- **distOH** (*real*) – (default: 1.0)

`espressopp.integrator.Settle.addMolecules` (*moleculelist*)

**Parameters** *moleculelist* –

**Return type**

## espressopp.integrator.StochasticVelocityRescaling

`espressopp.integrator.StochasticVelocityRescaling` (*system*)

**Parameters** *system* –

## espressopp.integrator.TDforce

Thermodynamic force.

Example - how to turn on thermodynamic force (except for multiple moving spherical regions)

```
>>> fthd="tablet.f.svg"
>>> thdforce = espressopp.integrator.TDforce(system,verletlist) #info about centre and shape of a
>>> thdforce.addForce(itype=3,filename="tablet.f.svg",type=typeCG)
>>> integrator.addExtension(thdforce)
```

Example - how to turn on thermodynamic force for multiple moving spherical regions

```
>>> fthd="tablet.f.svg"
>>> thdforce = espressopp.integrator.TDforce(system,verletlist, startdist = 0.9, enddist = 2.1, e
>>> thdforce.addForce(itype=3,filename="tablet.f.svg",type=typeCG)
>>> integrator.addExtension(thdforce)
```

`espressopp.integrator.TDforce` (*system, verletlist, startdist, enddist, edgeweightmultiplier*)

### Parameters

- **system** (*shared\_ptr<System>*) – system object
- **verletlist** (*shared\_ptr<VerletListAdress>*) – verletlist object
- **startdist** (*real*) – (default: 0.0) starting distance from center at which the TD force is actually applied. Needs to be altered when using several moving spherical regions (not used for static or single moving region)
- **enddist** (*real*) – (default: 0.0) end distance from center up to which the TD force is actually applied. Needs to be altered when using several moving spherical regions (not used for static or single moving region)
- **edgeweightmultiplier** (*int*) – (default: 20) interpolation parameter for multiple overlapping spherical regions (see Kreis et al., JCTC doi: 10.1021/acs.jctc.6b00440), the default should be fine for most applications (not used for static or single moving region)

`espressopp.integrator.TDforce.addForce` (*itype, filename, type*)

### Parameters

- **itype** (*int*) – interpolation type 1: linear, 2: Akima, 3: Cubic
- **filename** (*string*) – filename for TD force file
- **type** (*int*) – particle type on which the TD force needs to be applied

## espressopp.integrator.VelocityVerlet

`espressopp.integrator.VelocityVerlet` (*system*)

Parameters **system** –

## espressopp.integrator.VelocityVerletOnGroup

`espressopp.integrator.VelocityVerletOnGroup` (*system, group*)

### Parameters

- **system** –
- **group** –

## espressopp.integrator.VelocityVerletOnRadius

`espressopp.integrator.VelocityVerletOnRadius` (*system*, *dampingmass*)

### Parameters

- **system** –
- **dampingmass** –

## interaction

### espressopp.interaction.AngularCosineSquared

Calculates the Angular Cosine Squared interaction

$$U = K(\cos(\theta) - \cos(\theta_0))^2$$

`espressopp.interaction.AngularCosineSquared` (*K*, *theta0*)

### Parameters

- **K** (*real*) – (default: 1.0)
- **theta0** (*real*) – (default: 0.0)

`espressopp.interaction.FixedTripleListAngularCosineSquared` (*system*, *vl*, *potential*)

### Parameters

- **system** –
- **vl** –
- **potential** –

`espressopp.interaction.FixedTripleListAngularCosineSquared.getFixedTripleList` ()

**Return type** A Python list of lists.

`espressopp.interaction.FixedTripleListAngularCosineSquared.setPotential` (*type1*, *type2*, *potential*)

### Parameters

- **type1** –
- **type2** –
- **potential** –

**class** `espressopp.interaction.AngularCosineSquared.AngularCosineSquared`  
The AngularCosineSquared potential.

### espressopp.interaction.AngularHarmonic

Calculates the Angular Harmonic interaction

$$U = K(\theta - \theta_0)^2$$

`espressopp.interaction.AngularHarmonic` ( $K$ ,  $\theta_0$ )

**Parameters**

- **K** (*real*) – (default: 1.0)
- **theta0** (*real*) – (default: 0.0)

`espressopp.interaction.FixedTripleListAngularHarmonic` (*system*, *vl*, *potential*)

**Parameters**

- **system** –
- **vl** –
- **potential** –

`espressopp.interaction.FixedTripleListAngularHarmonic.setPotential` (*type1*,  
*type2*,  
*potential*)

**Parameters**

- **type1** –
- **type2** –
- **potential** –

`class espressopp.interaction.AngularHarmonic.AngularHarmonic`  
The AngularHarmonic potential.

## `espressopp.interaction.AngularPotential`

This is an abstract class, only needed to be inherited from.

`espressopp.interaction.AngularPotential.computeEnergy` (*\*args*)

**Parameters** *\*args* –

**Return type**

`espressopp.interaction.AngularPotential.computeForce` (*\*args*)

**Parameters** *\*args* –

**Return type**

## `espressopp.interaction.AngularUniqueCosineSquared`

Calculates the angular unique cosine squared interaction.

$$U = K(\cos(\theta) - \cos(\theta_0))^2$$

`espressopp.interaction.AngularUniqueCosineSquared` ( $K$ )

**Parameters** **K** (*real*) – (default: 1.0)

`espressopp.interaction.FixedTripleAngleListAngularUniqueCosineSquared` (*system*,  
*ftcl*,  
*po-*  
*ten-*  
*tial*)

**Parameters**

- **system** –
- **ftcl** –
- **potential** –

`espressopp.interaction.FixedTripleAngleListAngularUniqueCosineSquared.getFixedTripleList`

**Return type** A Python list of lists.

`espressopp.interaction.FixedTripleAngleListAngularUniqueCosineSquared.setPotential` (*potential*)

**Parameters** **potential** –

**class** `espressopp.interaction.AngularUniqueCosineSquared.AngularUniqueCosineSquared`  
The AngularUniqueCosineSquared potential.

## espressopp.interaction.AngularUniqueHarmonic

Calculates the Angular Unique Harmonic interaction

$$U = K(\theta - \theta_0)^2$$

`espressopp.interaction.AngularUniqueHarmonic` (*K*)

**Parameters** **K** (*real*) – (default: 1.0)

`espressopp.interaction.FixedTripleAngleListAngularUniqueHarmonic` (*system*,  
*ftal*, *potential*)

**Parameters**

- **system** –
- **ftal** –
- **potential** –

`espressopp.interaction.FixedTripleAngleListAngularUniqueHarmonic.setPotential` (*potential*)

**Parameters** **potential** –

## espressopp.interaction.AngularUniquePotential

This is an abstract class, only needed to be inherited from.

`espressopp.interaction.AngularUniquePotential.computeEnergy` (*\*args*)

**Parameters** **\*args** –

**Return type**

`espressopp.interaction.AngularUniquePotential.computeForce` (*\*args*)

**Parameters** **\*args** –

**Return type**

## espressopp.interaction.Cosine

Calculates the Cosine Interaction

$$U = K(1 + \cos(\theta - \theta_0))$$

`espressopp.interaction.Cosine(K, theta0)`

**Parameters**

- **K** (*real*) – (default: 1.0)
- **theta0** (*real*) – (default: 0.0)

`espressopp.interaction.FixedTripleListCosine(system, vl, potential)`

**Parameters**

- **system** –
- **vl** –
- **potential** –

`espressopp.interaction.FixedTripleListCosine.getFixedTripleList()`

**Return type** A Python list of lists.

`espressopp.interaction.FixedTripleListCosine.setPotential(potential)`

**Parameters** **potential** –

**class** `espressopp.interaction.Cosine.Cosine`

The Cosine potential.

## espressopp.interaction.CoulombKSpaceEwald

Coulomb potential and interaction Objects (*K* space part)

$$\frac{1}{2\pi V} \sum_{\substack{m \in \mathbb{Z}^3 \\ 0 < |m| < k_{max}}} \frac{\exp(-\frac{\pi^2}{\alpha^2} m'^2)}{m'^2} \left| \sum_{i=1}^N q_i \cdot \exp(2\pi i r_i \cdot m') \right|^2$$

This is the *K* space part of potential of Coulomb long range interaction according to the Ewald summation technique. Good explanation of Ewald summation could be found here [\[Allen89\]](#), [\[Deserno98\]](#).

Example:

```
>>> ewaldK_pot = espressopp.interaction.CoulombKSpaceEwald(system, coulomb_prefactor, alpha, kspacecutoff)
>>> ewaldK_int = espressopp.interaction.CellListCoulombKSpaceEwald(system.storage, ewaldK_pot)
>>> system.addInteraction(ewaldK_int)
```

**!IMPORTANT** Coulomb interaction needs *R* space part as well `CoulombRSpace`.

Definition:

It provides potential object *CoulombKSpaceEwald* and interaction object *CellListCoulombKSpaceEwald* based on all particles list.

The *potential* is based on the system information (System) and parameters: Coulomb prefactor (`coulomb_prefactor`), Ewald parameter (`alpha`), and the cutoff in *K* space (`kspacecutoff`).

```
>>> ewaldK_pot = espressopp.interaction.CoulombKSpaceEwald(system, coulomb_prefactor, alpha, kspacecutoff)
```

Potential Properties:

- *ewaldK\_pot.prefactor*

The property ‘prefactor’ defines the Coulomb prefactor.

- *ewaldK\_pot.alpha*

The property ‘alpha’ defines the Ewald parameter *alpha*.



- *ewaldK\_pot.kmax*

The property ‘kmax’ defines the cutoff in  $K$  space.

The *interaction* is based on the all particles list. It needs the information from Storage and  $K$  space part of potential.

```
>>> ewaldK_int = espressopp.interaction.CellListCoulombKSpaceEwald(system.storage, ewaldK_pot)
```

Interaction Methods:

- *getPotential()*

Access to the local potential.

Adding the interaction to the system:

```
>>> system.addInteraction(ewaldK_int)
```

References:

`espressopp.interaction.CoulombKSpaceEwald` (*system, prefactor, alpha, kmax*)

**Parameters**

- **system** –
- **prefactor** –
- **alpha** –
- **kmax** –

`espressopp.interaction.CellListCoulombKSpaceEwald` (*storage, potential*)

**Parameters**

- **storage** –
- **potential** –

`espressopp.interaction.CellListCoulombKSpaceEwald.getFixedPairList` ()

**Return type** A Python list of lists.

`espressopp.interaction.CellListCoulombKSpaceEwald.getPotential` ()

**Return type**

## espressopp.interaction.CoulombKSpaceP3M

Coulomb potential and interaction Objects ( $K$  space part)

This is the  $K$  space part of potential of Coulomb long range interaction according to the P3M summation technique. Good explanation of P3M summation could be found here [\[Allen89\]](#), [\[Deserno98\]](#).

Example:

```
>>> ewaldK_pot = espressopp.interaction.CoulombKSpaceP3M(system, coulomb_prefactor, alpha, kspace)
>>> ewaldK_int = espressopp.interaction.CellListCoulombKSpaceP3M(system.storage, ewaldK_pot)
>>> system.addInteraction(ewaldK_int)
```

**!IMPORTANT** Coulomb interaction needs  $R$  space part as well `CoulombRSpace`.

Definition:

It provides potential object *CoulombKSpaceP3M* and interaction object *CellListCoulombKSpaceP3M* based on all particles list.

The *potential* is based on the system information (System) and parameters: Coulomb prefactor (`coulomb_prefactor`), P3M parameter (`alpha`), and the cutoff in  $K$  space (`kspacecutoff`).

```
>>> ewaldK_pot = espressopp.interaction.CoulombKSpaceP3M(system, coulomb_prefactor, alpha, ks
```

Potential Properties:

- *ewaldK\_pot.prefactor*

The property ‘prefactor’ defines the Coulomb prefactor.

- *ewaldK\_pot.alpha*

The property ‘alpha’ defines the P3M parameter *alpha*.

- *ewaldK\_pot.kmax*

The property ‘kmax’ defines the cutoff in *K* space.

The *interaction* is based on the all particles list. It needs the information from Storage and *K* space part of potential.

```
>>> ewaldK_int = espressopp.interaction.CellListCoulombKSpaceP3M(system.storage, ewaldK_pot)
```

Interaction Methods:

- *getPotential()*

Access to the local potential.

Adding the interaction to the system:

```
>>> system.addInteraction(ewaldK_int)
```

```
espressopp.interaction.CoulombKSpaceP3M(system, C_pref, alpha, M, P, rcut, interpolation)
```

#### Parameters

- **system** –
- **C\_pref** –
- **alpha** –
- **M** –
- **P** –
- **rcut** –
- **interpolation** (*int*) – (default: 200192)

```
espressopp.interaction.CellListCoulombKSpaceP3M(storage, potential)
```

#### Parameters

- **storage** –
- **potential** –

```
espressopp.interaction.CellListCoulombKSpaceP3M.getPotential()
```

#### Return type

## espressopp.interaction.CoulombRSpace

Coulomb potential and interaction Objects (*R* space part)

$$\sum_{i=1}^N \sum_{\substack{j>i \\ r_{ij}<k_{max}}} \frac{q_i q_j}{r_{ij}} \operatorname{erfc}(\alpha r_{ij}) - \frac{\alpha}{\sqrt{\pi}} \sum_{i=1}^N q_i^2$$

This is the  $R$  space part of potential of Coulomb long range interaction according to the Ewald summation technique. Good explanation of Ewald summation could be found here [\[Allen89\]](#), [\[Deserno98\]](#).

Example:

```
>>> vl = espressopp.VerletList(system, rspacecutoff+skin)
>>> coulombR_pot = espressopp.interaction.CoulombRSpace(coulomb_prefactor, alpha, rspacecutoff)
>>> coulombR_int = espressopp.interaction.VerletListCoulombRSpace(vl)
>>> coulombR_int.setPotential(type1=0, type2=0, potential = coulombR_pot)
>>> system.addInteraction(coulombR_int)
```

**!IMPORTANT** Coulomb interaction needs k-space part as well EwaldKSpace.

Definition:

It provides potential object *CoulombRSpace* and interaction object *VerletListCoulombRSpace*

The *potential* is based on parameters: Coulomb prefactor (*coulomb\_prefactor*), Ewald parameter (*alpha*), and the cutoff in  $R$  space (*rspacecutoff*).

```
>>> coulombR_pot = espressopp.interaction.CoulombRSpace(coulomb_prefactor, alpha, rspacecutoff)
```

Potential Properties:

- *coulombR\_pot.prefactor*  
The property 'prefactor' defines the Coulomb prefactor.
- *coulombR\_pot.alpha*  
The property 'alpha' defines the Ewald parameter *alpha*.
- *coulombR\_pot.cutoff*  
The property 'cutoff' defines the cutoff in  $R$  space.

The *interaction* is based on the Verlet list (*VerletList*)

```
>>> vl = espressopp.VerletList(system, rspacecutoff+skin)
>>> coulombR_int = espressopp.interaction.VerletListCoulombRSpace(vl)
```

It should include at least one potential

```
>>> coulombR_int.setPotential(type1=0, type2=0, potential = coulombR_pot)
```

Interaction Methods:

- *setPotential(type1, type2, potential)*  
This method sets the *potential* for the particles of *type1* and *type2*. It could be a bunch of potentials for the different particle types.
- *getVerletListLocal()*  
Access to the local Verlet list.

Adding the interaction to the system:

```
>>> system.addInteraction(coulombR_int)
```

`espressopp.interaction.CoulombRSpace` (*prefactor*, *alpha*, *cutoff*)

**Parameters**

- **prefactor** (*real*) – (default: 1.0)
- **alpha** (*real*) – (default: 1.0)
- **cutoff** – (default: infinity)

`espressopp.interaction.VerletListCoulombRSpace` (*vl*)

#### Parameters **v1** –

`espressopp.interaction.VerletListCoulombRSpace.getPotential (type1, type2)`

#### Parameters

- **type1** –
- **type2** –

#### Return type

`espressopp.interaction.VerletListCoulombRSpace.getVerletList ()`

**Return type** A Python list of lists.

`espressopp.interaction.VerletListCoulombRSpace.setPotential (type1, type2, potential)`

#### Parameters

- **type1** –
- **type2** –
- **potential** –

## espressopp.interaction.CoulombTruncated

$$U = k \frac{q_i q_j}{d_{ij}}$$

where  $k$  is the user-supplied prefactor,  $q_i$  is the charge of particle  $i$ , and  $d_{ij}$  is interparticle distance

In this interaction potential, a different charge can be associated with each particle. For a truncated Coulomb interaction potential where only one  $q_i q_j$  value is specified for all interactions, see `CoulombTruncatedUniqueCharge`.

`espressopp.interaction.CoulombTruncated (prefactor, cutoff)`

#### Parameters

- **prefactor** (*real*) – (default: 1.0) user-supplied prefactor  $k$
- **cutoff** (*real*) – (default: infinity) user-supplied interaction cutoff

`espressopp.interaction.VerletListCoulombTruncated (vl)`

**Parameters** **v1** (`espressopp.VerletList`) – verlet list object defined earlier in python script

`espressopp.interaction.VerletListCoulombTruncated.getPotential (type1, type2)`

#### Parameters

- **type1** (*integer*) – type of first atom in pair
- **type2** (*integer*) – type of second atom in pair

`espressopp.interaction.VerletListCoulombTruncated.setPotential (type1, type2, potential)`

#### Parameters

- **type1** (*integer*) – type of first atom in pair
- **type2** (*integer*) – type of second atom in pair

- **potential** (*CoulombTruncated potential*) – potential object defined earlier in python script

```
espressopp.interaction.FixedPairListTypesCoulombTruncated(system, vl)
```

#### Parameters

- **system** (*espressopp.System*) – system object defined earlier in the python script
- **vl** (*espressopp.FixedPairList*) – fixedpairlist object defined earlier in the python script

```
espressopp.interaction.FixedPairListTypesCoulombTruncated.setPotential(potential)
```

#### Parameters

- **type1** (*integer*) – type of first atom in pair
- **type2** (*integer*) – type of second atom in pair
- **potential** (*CoulombTruncated potential*) – potential object defined earlier in python script

#Example:

```
>>> pref = 138.935485
>>> rc = 1.2
>>> fixedpairlist = espresso.FixedPairList(system.storage)
>>> fixedpairlist.addBonds([(1,2),(2,3)])
>>> pot = espressopp.interaction.CoulombTruncated(prefactor=pref, cutoff=rc)
>>> interaction=espressopp.interaction.FixedPairListTypesCoulombTruncated(system,fixedpairlist)
>>> interaction.setPotential(type1=0, type2=1, potential=pot)
>>> system.addInteraction(interaction)
```

**class** espressopp.interaction.CoulombTruncated.CoulombTruncated

The CoulombTruncated potential.

## espressopp.interaction.CoulombTruncatedUniqueCharge

$$U = \frac{Q}{d}$$

where  $Q$  is the product of the charges of the two particles and  $d$  is their distance from each other.

In this interaction potential, a unique  $Q = q_i q_j$  value is specified per potential. For a more flexible truncated Coulomb interaction potential where each individual particle has its own charge  $q_i$ , see `CoulombTruncated`.

```
espressopp.interaction.CoulombTruncatedUniqueCharge(qq, cutoff, shift)
```

#### Parameters

- **qq** (*real*) – (default: 1.0)
- **cutoff** – (default: infinity)
- **shift** – (default: “auto”)

```
espressopp.interaction.VerletListCoulombTruncatedUniqueCharge(vl)
```

#### Parameters vl –

```
espressopp.interaction.VerletListCoulombTruncatedUniqueCharge.getPotential(type1, type2)
```

#### Parameters

- **type1** –
- **type2** –

### Return type

`espressopp.interaction.VerletListCoulombTruncatedUniqueCharge.setPotential` (*type1*,  
*type2*,  
*po-*  
*ten-*  
*tial*)

### Parameters

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.CellListCoulombTruncatedUniqueCharge` (*stor*)

### Parameters stor –

`espressopp.interaction.CellListCoulombTruncatedUniqueCharge.setPotential` (*type1*,  
*type2*,  
*po-*  
*ten-*  
*tial*)

### Parameters

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.FixedPairListCoulombTruncatedUniqueCharge` (*system*,  
*vl*, *poten-*  
*tial*)

### Parameters

- **system** –
- **vl** –
- **potential** –

`espressopp.interaction.FixedPairListCoulombTruncatedUniqueCharge.setPotential` (*potential*)

### Parameters potential –

**class** `espressopp.interaction.CoulombTruncatedUniqueCharge.CoulombTruncatedUniqueCharge`  
The CoulombTruncatedUniqueCharge potential.

## espressopp.interaction.ConstrainCOM

This class is for calculating forces of constraining center of mass of subchains. Subchains are defined as a tuple list.

$$U = k_{com} \left( \vec{r}_{com} - \vec{R}_{com} \right)^2,$$

where  $\vec{r}_{com}$  stands for the center of mass of subchain and  $\vec{R}_{com}$  stands for the desired center of mass of subchain.

This class set 2 conditions on a tuple list defining subchains.

1. The length of all tuple must be same.
2. `int(key particle id / The length of a tuple)` must not be redundantly, where key particle id is the smallest particle id in a tuple.

Reference: Equilibration of high molecular weight polymer melts: A hierarchical strategy, Macro Lett., 2014, 3, 198

`espressopp.interaction.ConstrainCOM(k_com)`

**Parameters** *k\_com* (*real*) – (default: 100.)

`espressopp.interaction.FixedLocalTupleListConstrainCOM(system, tuplelist, potential)`

**Parameters**

- **system** –
- **tuplelist** –
- **potential** –

`espressopp.interaction.FixedLocalTupleListConstrainCOM.getPotential()`

**Return type**

`espressopp.interaction.FixedLocalTupleListConstrainCOM.setCom(particlelist)`

**Parameters** *particlelist* –

## espressopp.interaction.ConstrainRG

This class is for calculating forces of constraining radii of gyration of subchains. Subchains are defined as a tuple list.

$$U = k_{rg} \left( R_g^2 - R_g^{ideal^2} \right)^2$$

where  $R_g^{ideal}$  stands for the desired radius of gyration of subchain.

This class set 2 conditions on a tuple list. defining subchains.

1. The length of all tuple must be same.
2. `int(key particle id / The length of a tuple)` must not be redundantly, where key particle id is the smallest particle id in a tuple.

Reference: Equilibration of high molecular weight polymer melts: A hierarchical strategy, Macro Lett., 2014, 3, 198

`espressopp.interaction.ConstrainRG(k_rg)`

**Parameters** *k\_rg* (*real*) – (default: 100.)

`espressopp.interaction.FixedLocalTupleListConstrainRG(system, tuplelist, potential)`

**Parameters**

- **system** –
- **tuplelist** –
- **potential** –

`espressopp.interaction.FixedLocalTupleListConstrainRG.getPotential()`

**Return type**

`espressopp.interaction.FixedLocalTupleListConstrainRG.setRG(particlelist)`

**Parameters** *particlelist* (*python::list*) –

## espressopp.interaction.DihedralHarmonic

The dihedral harmonic potential

$$U(\phi_{ijkl}) = 0.5K[\phi_{ijkl} - \phi_0]^2$$

where the  $K$  is a constant, the angles should be provided in radians.

Reference: Gromacs Manual 4.6.1, section 4.2.11 (page 79-80), equation 4.60

`espressopp.interaction.DihedralHarmonic` ( $K$ ,  $\phi_0$ )

### Parameters

- **K** (*real*) – (default: 0.0)
- **phi0** (*real*) – (default: 0.0)

`espressopp.interaction.FixedQuadrupleListDihedralHarmonic` (*system*, *fql*, *potential*)

### Parameters

- **system** –
- **fql** –
- **potential** –

`espressopp.interaction.FixedQuadrupleListDihedralHarmonic.getFixedQuadrupleList` ()

**Return type** A Python list of lists.

`espressopp.interaction.FixedQuadrupleListDihedralHarmonic.setPotential` (*potential*)

**Parameters** *potential* –

### Example of usage

```
>>> # The following example shows how to add a torsional potential to particles 1,2,3,4
>>> fql = espressopp.FixedQuadrupleList(system.storage)
>>> fql.addQuadruples([(1,2,3,4)])
>>> #phi0 is in radians, IUPAC convention definition
>>> interaction = espressopp.interaction.FixedQuadrupleListDihedralHarmonic(system, fql, potential=0.5, phi0=0.0)
>>> system.addInteraction(interaction)
```

**class** `espressopp.interaction.DihedralHarmonic.DihedralHarmonic`

The DihedralHarmonic potential.

**class** `espressopp.interaction.DihedralHarmonic.FixedQuadrupleListDihedralHarmonicLocal` (*system*, *fql*, *potential*)

The (local) DihedralHarmonic interaction using FixedQuadruple lists.

## espressopp.interaction.DihedralHarmonicCos

$$U = K(\cos(\phi) - \cos(\phi_0))^2$$

`espressopp.interaction.DihedralHarmonicCos` ( $K$ ,  $\phi_0$ )

### Parameters

- **K** (*real*) – (default: 0.0)
- **phi0** (*real*) – (default: 0.0)



`espressopp.interaction.FixedQuadrupleListDihedralHarmonicCos` (*system, fql, potential*)

**Parameters**

- **system** –
- **fql** –
- **potential** –

`espressopp.interaction.FixedQuadrupleListDihedralHarmonicCos.getFixedQuadrupleList()`

**Return type** A Python list of lists.

`espressopp.interaction.FixedQuadrupleListDihedralHarmonicCos.setPotential` (*potential*)

**Parameters potential** –

**class** `espressopp.interaction.DihedralHarmonicCos.DihedralHarmonicCos`  
The DihedralHarmonicCos potential.

## espressopp.interaction.DihedralHarmonicNCos

The dihedral harmonic potential

$$U(\phi_{ijkl}) = K[1 + \cos(N \cdot \phi_{ijkl} - \phi_0)]$$

where the  $K$  is a constant, the angles should be provided in radians. The  $N$  is a multiplicity.

Reference: <http://www.uark.edu/ua/fengwang/DLPOLY2/node49.html>

`espressopp.interaction.DihedralHarmonicNCos` (*K, phi0, multiplicity*)

**Parameters**

- **K** (*real*) – (default: 0.0)
- **phi0** (*real*) – (default: 0.0)
- **multiplicity** (*int*) – (default: 1)

`espressopp.interaction.FixedQuadrupleListDihedralHarmonicNCos` (*system, fql, potential*)

**Parameters**

- **system** –
- **fql** –
- **potential** –

`espressopp.interaction.FixedQuadrupleListDihedralHarmonicNCos.getFixedQuadrupleList()`

**Return type** A Python list of lists.

`espressopp.interaction.FixedQuadrupleListDihedralHarmonicNCos.setPotential` (*potential*)

**Parameters potential** –

**class** `espressopp.interaction.DihedralHarmonicNCos.DihedralHarmonicNCos`  
The DihedralHarmonicNCos potential.

**class** `espressopp.interaction.DihedralHarmonicNCos.FixedQuadrupleListDihedralHarmonicNCosLo`

The (local) DihedralHarmonicNCos interaction using FixedQuadruple lists.

## espressopp.interaction.DihedralHarmonicUniqueCos

$$U = K(\cos(\phi) - \cos(\phi_0))^2$$

`espressopp.interaction.DihedralHarmonicUniqueCos (K)`

**Parameters** *K* (*real*) – (default: 0.0)

`espressopp.interaction.FixedQuadrupleAngleListDihedralHarmonicUniqueCos (system, fgal, po-ten-tial)`

**Parameters**

- **system** –
- **fgal** –
- **potential** –

`espressopp.interaction.FixedQuadrupleAngleListDihedralHarmonicUniqueCos.getFixedQuadrupleAngleList`

**Return type** A Python list of lists.

`espressopp.interaction.FixedQuadrupleAngleListDihedralHarmonicUniqueCos.setPotential (potential)`

**Parameters** *potential* –

**class** `espressopp.interaction.DihedralHarmonicUniqueCos.DihedralHarmonicUniqueCos`  
The DihedralHarmonicUniqueCos potential.

## espressopp.interaction.DihedralPotential

This is an abstract class, only needed to be inherited from.

`espressopp.interaction.DihedralPotential.computeEnergy (*args)`

**Parameters** *\*args* –

**Return type**

`espressopp.interaction.DihedralPotential.computeForce (*args)`

**Parameters** *\*args* –

**Return type**

## espressopp.interaction.DihedralRB

The proper dihedral with Ryckaert-Bellemans form.

$$U_{rb}(\phi_{ijkl}) = \sum_{n=0}^5 K_n(\cos(\theta))^n$$

where the  $\theta = \phi - 180^\circ$  and  $K_{0...5}$  are the coefficients.

By default the IUPAC convention is used, where  $\phi$  is the angle between planes  $ijk$  and  $jkl$ . The  $0^\circ$  corresponds to the *cis* configuration.

Reference: <http://www.gromacs.org/Documentation/Manual>

`espressopp.interaction.DihedralRB (K0, K1, K2, K3, K4, K5, iupac)`

### Parameters

- **K0** (*real*) – (default: 0.0)
- **K1** (*real*) – (default: 0.0)
- **K2** (*real*) – (default: 0.0)
- **K3** (*real*) – (default: 0.0)
- **K4** (*real*) – (default: 0.0)
- **K5** (*real*) – (default: 0.0)
- **iupac** – (default: True)

`espressopp.interaction.FixedQuadrupleListDihedralRB` (*system*, *vl*, *potential*)

### Parameters

- **system** –
- **vl** –
- **potential** –

`espressopp.interaction.FixedQuadrupleListDihedralRB.getFixedQuadrupleList` ()

**Return type** A Python list of lists.

`espressopp.interaction.FixedQuadrupleListDihedralRB.setPotential` (*type1*,  
*type2*,  
*potential*)

### Parameters

- **type1** –
- **type2** –
- **potential** –

## espressopp.interaction.DihedralUniquePotential

This is an abstract class, only needed to be inherited from.

`espressopp.interaction.DihedralUniquePotential.computeEnergy` (\*args)

**Parameters** \*args –

**Return type**

`espressopp.interaction.DihedralUniquePotential.computeForce` (\*args)

**Parameters** \*args –

**Return type**

## espressopp.interaction.FENE

Implementation of the Finitely Extensible Non-linear Elastic potential:

$$U(r) = -\frac{1}{2}r_{\max}^2 K \log \left[ 1 - \left( \frac{r - r_0}{r_{\max}} \right)^2 \right]$$

`espressopp.interaction.FENE` (*K*, *r0*, *rMax*, *cutoff*, *shift*)

**Parameters**

- **K** (*real*) – (default: 1.0)
- **r0** (*real*) – (default: 0.0)
- **rMax** (*real*) – (default: 1.0)
- **cutoff** (*real*) – (default: infinity)
- **shift** (*real*) – (default: 0.0)

`espressopp.interaction.FixedPairListFENE (system, pair_list, potential)`

#### Parameters

- **system** (*object*) – your system `espressopp.System()`
- **pair\_list** (*object*) – list of bonds `espressopp.FixedPairList()`
- **potential** (*object*) – `espressopp.interaction.FENE()`

`espressopp.interaction.FixedPairListFENE.getFixedPairList()`

**Return type** A Python list of lists.

`espressopp.interaction.FixedPairListFENE.getPotential()`

**Return type** object

`espressopp.interaction.FixedPairListFENE.setFixedPairList (pair_list)`

**Parameters** **pair\_list** (*fixedpairlist*) –

`espressopp.interaction.FixedPairListFENE.setPotential (potential)`

**Parameters** **potential** –

#### Example of usage

```
>>> # The following example shows how to bond particle 1 to particles 0 and 2 by a FENE potential
>>> # We assume the particles are already in the storage of the system
>>> # Initialize list of pairs that will be bonded by FENE
>>> pair_list = espressopp.FixedPairList(system.storage)
>>> # Set which pairs belong to the pair_list i.e. particle 0 is bonded to particles 1 and 2.
>>> pair_list.addBonds([(0,1),(1,2)])
>>> # Initialize the potential and set up the parameters.
>>> potFENE = espressopp.interaction.FENE(K=30.0, r0=0.0, rMax=1.5)
>>> # Set which system, pair list and potential is the interaction associated with.
>>> interFENE = espressopp.interaction.FixedPairListFENE(system, pair_list, potFENE)
>>> # Add the interaction to the system.
>>> system.addInteraction(interFENE)
```

## espressopp.interaction.FENECapped

$$U = -\frac{1}{2}r_{max}^2 K \cdot \log\left(1 - \frac{D - r_0^2}{r_{max}}\right)$$

where  $D = dist$  if

$cap_{rad}^2 > dist$

and  $D = cap_{rad}$  else.

`espressopp.interaction.FENECapped (K, r0, rMax, cutoff, caprad, shift)`

#### Parameters

- **K** (*real*) – (default: 1.0)
- **r0** (*real*) – (default: 0.0)

- **rMax** (*real*) – (default: 1.0)
- **cutoff** – (default: infinity)
- **caprad** (*real*) – (default: 1.0)
- **shift** (*real*) – (default: 0.0)

`espressopp.interaction.FixedPairListFENECapped(system, vl, potential)`

#### Parameters

- **system** –
- **vl** –
- **potential** –

`espressopp.interaction.FixedPairListFENECapped.getFixedPairList()`

**Return type** A Python list of lists.

`espressopp.interaction.FixedPairListFENECapped.getPotential()`

**Return type**

`espressopp.interaction.FixedPairListFENECapped.setFixedPairList(fixedpairlist)`

**Parameters** **fixedpairlist** –

`espressopp.interaction.FixedPairListFENECapped.setPotential(potential)`

**Parameters** **potential** –

**class** `espressopp.interaction.FENECapped.FENECapped`  
The FENECapped potential.

## espressopp.interaction.GravityTruncated

This is an implementation of a truncated (cutoff) Gravity Potential

$$U = P \cdot \frac{m_1 \cdot m_2}{|p_1 - p_2|}$$

where  $m_i$  is the mass of the  $i$  th particle,  $p_i$  its position and  $P$  a prefactor.

`espressopp.interaction.GravityTruncated(prefactor, cutoff)`

#### Parameters

- **prefactor** (*real*) – (default: 1.0)
- **cutoff** – (default: infinity)

`espressopp.interaction.VerletListGravityTruncated(vl)`

**Parameters** **vl** –

`espressopp.interaction.VerletListGravityTruncated.getPotential(type1, type2)`

#### Parameters

- **type1** –
- **type2** –

**Return type**

`espressopp.interaction.VerletListGravityTruncated.getVerletList()`

**Return type** A Python list of lists.

`espressopp.interaction.VerletListGravityTruncated.setPotential` (*type1*,  
*type2*,  
*potential*)

**Parameters**

- **type1** –
- **type2** –
- **potential** –

## **espressopp.interaction.Harmonic**

$$U = K(d - r_0)^2$$

`espressopp.interaction.Harmonic` (*K*, *r0*, *cutoff*, *shift*)

**Parameters**

- **K** (*real*) – (default: 1.0)
- **r0** (*real*) – (default: 0.0)
- **cutoff** – (default: infinity)
- **shift** (*real*) – (default: 0.0)

`espressopp.interaction.FixedPairListHarmonic` (*system*, *vl*, *potential*)

**Parameters**

- **system** –
- **vl** –
- **potential** –

`espressopp.interaction.FixedPairListHarmonic.getFixedPairList` ()

**Return type** A Python list of lists.

`espressopp.interaction.FixedPairListHarmonic.setFixedPairList` (*fixedpairlist*)

**Parameters** **fixedpairlist** –

`espressopp.interaction.FixedPairListHarmonic.setPotential` (*potential*)

**Parameters** **potential** –

`espressopp.interaction.FixedPairListTypesHarmonic` (*system*, *vl*)

**Parameters**

- **system** –
- **vl** –

`espressopp.interaction.FixedPairListTypesHarmonic.getFixedPairList` ()

**Return type** A Python list of lists.

`espressopp.interaction.FixedPairListTypesHarmonic.setFixedPairList` (*fixedpairlist*)

**Parameters** **fixedpairlist** –

`espressopp.interaction.FixedPairListTypesHarmonic.setPotential` (*type1*,  
*type2*,  
*potential*)

#### Parameters

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.FixedPairListTypesHarmonic.getPotential (type1, type2)`

#### Parameters

- **type1** –
- **type2** –

#### Return type

**class** `espressopp.interaction.Harmonic.Harmonic`  
The Harmonic potential.

## `espressopp.interaction.HarmonicTrap`

$$U = K \frac{1}{2} d^2$$

`espressopp.interaction.HarmonicTrap ()`

`espressopp.interaction.SingleParticleHarmonicTrap (system, potential)`

#### Parameters

- **system** –
- **potential** –

`espressopp.interaction.SingleParticleHarmonicTrap.setPotential (potential)`

#### Parameters **potential** –

**class** `espressopp.interaction.HarmonicTrap.HarmonicTrap`  
The HarmonicTrap potential.

## `espressopp.interaction.HarmonicUnique`

$$U = K(d - d_{cur})^2;$$

`espressopp.interaction.HarmonicUnique (K)`

**Parameters** **K** (*real*) – (default: 1.0)

`espressopp.interaction.FixedPairDistListHarmonicUnique (system, fpl, potential)`

#### Parameters

- **system** –
- **fpl** –
- **potential** –

`espressopp.interaction.FixedPairDistListHarmonicUnique.getFixedPairList ()`

**Return type** A Python list of lists.

`espressopp.interaction.FixedPairDistListHarmonicUnique.setFixedPairList (fixedpairlist)`

**Parameters** `fixedpairlist` –

`espressopp.interaction.FixedPairDistListHarmonicUnique.setPotential (potential)`

**Parameters** `potential` –

**class** `espressopp.interaction.HarmonicUnique.HarmonicUnique`

The HarmonicUnique potential.

## espressopp.interaction.Interaction

This is an abstract class, only needed to be inherited from.

`espressopp.interaction.Interaction.bondType ()`

**Return type** `int`

`espressopp.interaction.Interaction.computeEnergy ()`

**Return type** `real`

`espressopp.interaction.Interaction.computeEnergyAA ()`

**Return type** `real`

`espressopp.interaction.Interaction.computeEnergyDeriv ()`

**Return type** `real`

`espressopp.interaction.Interaction.computeEnergyCG ()`

**Return type** `real`

`espressopp.interaction.Interaction.computeVirial ()`

**Return type** `real`

## espressopp.interaction.LennardJones

$$V(r) = 4\varepsilon \left[ \left( \frac{\sigma}{r} \right)^{12} - \left( \frac{\sigma}{r} \right)^6 \right]$$

`espressopp.interaction.LennardJones (epsilon, sigma, cutoff, shift)`

**Parameters**

- **epsilon** (*real*) – (default: 1.0)
- **sigma** (*real*) – (default: 1.0)
- **cutoff** – (default: infinity)
- **shift** – (default: “auto”)

`espressopp.interaction.VerletListLennardJones (vl)`

**Parameters** `vl` –

`espressopp.interaction.VerletListLennardJones.getPotential (type1, type2)`

**Parameters**

- **type1** –
- **type2** –

**Return type**



`espressopp.interaction.VerletListLennardJones.getVerletList()`

**Return type** A Python list of lists.

`espressopp.interaction.VerletListLennardJones.setPotential(type1, type2, potential)`

**Parameters**

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.VerletListAdressLennardJones(vl, fixedtupleList)`

**Parameters**

- **vl** –
- **fixedtupleList** –

`espressopp.interaction.VerletListAdressLennardJones.setPotentialAT(type1, type2, potential)`

**Parameters**

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.VerletListAdressLennardJones.setPotentialCG(type1, type2, potential)`

**Parameters**

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.VerletListAdressLennardJones2(vl, fixedtupleList)`

**Parameters**

- **vl** –
- **fixedtupleList** –

`espressopp.interaction.VerletListAdressLennardJones2.setPotentialAT(type1, type2, potential)`

**Parameters**

- **type1** –
- **type2** –
- **potential** –

```

espressopp.interaction.VerletListAdressLennardJones2.setPotentialCG (type1,
                                                                    type2,
                                                                    po-
                                                                    ten-
                                                                    tial)

```

#### Parameters

- **type1** –
- **type2** –
- **potential** –

```

espressopp.interaction.VerletListHadressLennardJones (vl, fixedtupleList)

```

#### Parameters

- **vl** –
- **fixedtupleList** –

```

espressopp.interaction.VerletListHadressLennardJones.setPotentialAT (type1,
                                                                    type2,
                                                                    po-
                                                                    ten-
                                                                    tial)

```

#### Parameters

- **type1** –
- **type2** –
- **potential** –

```

espressopp.interaction.VerletListHadressLennardJones.setPotentialCG (type1,
                                                                    type2,
                                                                    po-
                                                                    ten-
                                                                    tial)

```

#### Parameters

- **type1** –
- **type2** –
- **potential** –

```

espressopp.interaction.VerletListHadressLennardJones2 (vl, fixedtupleList)

```

#### Parameters

- **vl** –
- **fixedtupleList** –

```

espressopp.interaction.VerletListHadressLennardJones2.setPotentialAT (type1,
                                                                    type2,
                                                                    po-
                                                                    ten-
                                                                    tial)

```

#### Parameters

- **type1** –
- **type2** –
- **potential** –

```

espressopp.interaction.VerletListHadressLennardJones2.setPotentialCG(type1,
                                                                    type2,
                                                                    po-
                                                                    ten-
                                                                    tial)
    
```

#### Parameters

- **type1** –
- **type2** –
- **potential** –

```

espressopp.interaction.CellListLennardJones(stor)
    
```

#### Parameters stor –

```

espressopp.interaction.CellListLennardJones.setPotential(type1, type2, poten-
                                                         tial)
    
```

#### Parameters

- **type1** –
- **type2** –
- **potential** –

```

espressopp.interaction.FixedPairListLennardJones(system, vl, potential)
    
```

#### Parameters

- **system** –
- **vl** –
- **potential** –

```

espressopp.interaction.FixedPairListLennardJones.getFixedPairList()
    
```

**Return type** A Python list of lists.

```

espressopp.interaction.FixedPairListLennardJones.getPotential()
    
```

**Return type**

```

espressopp.interaction.FixedPairListLennardJones.setFixedPairList(fixedpairlist)
    
```

**Parameters fixedpairlist** –

```

espressopp.interaction.FixedPairListLennardJones.setPotential(potential)
    
```

**Parameters potential** –

```

class espressopp.interaction.LennardJones.LennardJones
    
```

The Lennard-Jones potential.

## espressopp.interaction.LennardJones93Wall

This class defines a Lennard-Jones 9-3 SingleParticlePotential in the direction x.

$$V(r) = \epsilon \left( \left( \frac{\sigma}{r} \right)^9 - \left( \frac{\sigma}{r} \right)^3 \right)$$

where  $r$  is the distance from the lower or upper wall in the x direction.  $V(r) = 0$  after a distance *sigmaCutoff*.

The parameters have to be defined for every species present in the system with *setParams* and can be retrieved with *getParams*.

Example:

```
>>> LJ93 = espressopp.interaction.LennardJones93Wall()
>>> LJ93.setParams(0, 6., 1., wall_cutoff)
>>> SPLJ93 = espressopp.interaction.SingleParticleLennardJones93Wall(system, LJ93)
>>> system.addInteraction(SPLJ93)
```

```
espressopp.interaction.LennardJones93Wall()
```

```
espressopp.interaction.LennardJones93Wall.getParams(type_var)
```

**Parameters** *type\_var* –

**Return type**

```
espressopp.interaction.LennardJones93Wall.setParams(type_var, epsilon, sigma,
                                                    sigmaCutoff, r0)
```

**Parameters**

- **type\_var** –
- **epsilon** –
- **sigma** –
- **sigmaCutoff** –
- **r0** –

```
espressopp.interaction.SingleParticleLennardJones93Wall(system, potential)
```

**Parameters**

- **system** –
- **potential** –

```
espressopp.interaction.SingleParticleLennardJones93Wall.setPotential(potential)
```

**Parameters** *potential* –

```
class espressopp.interaction.LennardJones93Wall.LennardJones93Wall
```

The LennardJones93Wall potential.

## espressopp.interaction.LennardJonesAutoBonds

$$V(r) = 4\varepsilon \left[ \left( \frac{\sigma}{r} \right)^{12} - \left( \frac{\sigma}{r} \right)^6 \right]$$

```
espressopp.interaction.LennardJonesAutoBonds(epsilon, sigma, cutoff, bondlist, maxcrosslinks)
```

**Parameters**

- **epsilon** (*real*) – (default: 1.0)
- **sigma** (*real*) – (default: 1.0)
- **cutoff** – (default: infinity)
- **bondlist** – (default: None)
- **maxcrosslinks** (*int*) – (default: 2)

```
espressopp.interaction.VerletListLennardJonesAutoBonds(vl)
```

**Parameters** *vl* –

```
espressopp.interaction.VerletListLennardJonesAutoBonds.getPotential(type1,
                                                                    type2)
```

#### Parameters

- **type1** –
- **type2** –

#### Return type

`espressopp.interaction.VerletListLennardJonesAutoBonds.getVerletList()`

**Return type** A Python list of lists.

`espressopp.interaction.VerletListLennardJonesAutoBonds.setPotential` (*type1*,  
*type2*,  
*po-*  
*ten-*  
*tial*)

#### Parameters

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.VerletListAdressLennardJonesAutoBonds` (*vl*, *fixedtu-*  
*pleList*)

#### Parameters

- **vl** –
- **fixedtupleList** –

`espressopp.interaction.VerletListAdressLennardJonesAutoBonds.setPotential` (*type1*,  
*type2*,  
*po-*  
*ten-*  
*tial*)

#### Parameters

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.VerletListHadressLennardJonesAutoBonds` (*vl*, *fixedtu-*  
*pleList*)

#### Parameters

- **vl** –
- **fixedtupleList** –

`espressopp.interaction.VerletListHadressLennardJonesAutoBonds.setPotential` (*type1*,  
*type2*,  
*po-*  
*ten-*  
*tial*)

#### Parameters

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.CellListLennardJonesAutoBonds` (*stor*)

#### Parameters **stor** –

`espressopp.interaction.CellListLennardJonesAutoBonds.setPotential` (*type1*,  
*type2*,  
*potential*)

#### Parameters

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.FixedPairListLennardJonesAutoBonds` (*system*, *vl*, *potential*)

#### Parameters

- **system** –
- **vl** –
- **potential** –

`espressopp.interaction.FixedPairListLennardJonesAutoBonds.setPotential` (*potential*)

#### Parameters **potential** –

**class** `espressopp.interaction.LennardJonesAutoBonds.LennardJonesAutoBonds`

The Lennard-Jones auto bonds potential.

## espressopp.interaction.LennardJonesCapped

$$V(r) = 4\varepsilon \left[ \left( \frac{\sigma}{r} \right)^{12} - \left( \frac{\sigma}{r} \right)^6 \right]$$

where  $r$  is either the distance or the capped distance, depending on which is greater.

`espressopp.interaction.LennardJonesCapped` (*epsilon*, *sigma*, *cutoff*, *caprad*, *shift*)

#### Parameters

- **epsilon** (*real*) – (default: 1.0)
- **sigma** (*real*) – (default: 1.0)
- **cutoff** – (default: infinity)
- **caprad** (*real*) – (default: 0.0)
- **shift** – (default: “auto”)

`espressopp.interaction.VerletListLennardJonesCapped` (*vl*)

#### Parameters **vl** –

`espressopp.interaction.VerletListLennardJonesCapped.getPotential` (*type1*,  
*type2*)

#### Parameters

- **type1** –
- **type2** –

#### Return type

`espressopp.interaction.VerletListLennardJonesCapped.setPotential` (*type1*,  
*type2*,  
*potential*)

#### Parameters

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.VerletListAdressLennardJonesCapped` (*vl*, *fixedtupleList*)

#### Parameters

- **vl** –
- **fixedtupleList** –

`espressopp.interaction.VerletListAdressLennardJonesCapped.getPotentialAT` (*type1*,  
*type2*)

#### Parameters

- **type1** –
- **type2** –

#### Return type

`espressopp.interaction.VerletListAdressLennardJonesCapped.getPotentialCG` (*type1*,  
*type2*)

#### Parameters

- **type1** –
- **type2** –

#### Return type

`espressopp.interaction.VerletListAdressLennardJonesCapped.setPotentialAT` (*type1*,  
*type2*,  
*po-*  
*ten-*  
*tial*)

#### Parameters

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.VerletListAdressLennardJonesCapped.setPotentialCG` (*type1*,  
*type2*,  
*po-*  
*ten-*  
*tial*)

#### Parameters

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.VerletListHadressLennardJonesCapped` (*vl*, *fixedtupleList*)

#### Parameters

- **v1** –
- **fixedtupleList** –

`espressopp.interaction.VerletListHadressLennardJonesCapped.getPotentialAT` (*type1*,  
*type2*)

**Parameters**

- **type1** –
- **type2** –

**Return type**

`espressopp.interaction.VerletListHadressLennardJonesCapped.getPotentialCG` (*type1*,  
*type2*)

**Parameters**

- **type1** –
- **type2** –

**Return type**

`espressopp.interaction.VerletListHadressLennardJonesCapped.setPotentialAT` (*type1*,  
*type2*,  
*po-*  
*ten-*  
*tial*)

**Parameters**

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.VerletListHadressLennardJonesCapped.setPotentialCG` (*type1*,  
*type2*,  
*po-*  
*ten-*  
*tial*)

**Parameters**

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.CellListLennardJonesCapped` (*stor*)

**Parameters stor** –

`espressopp.interaction.CellListLennardJonesCapped.getPotential` (*type1*,  
*type2*)

**Parameters**

- **type1** –
- **type2** –

**Return type**

`espressopp.interaction.CellListLennardJonesCapped.setPotential` (*type1*,  
*type2*,  
*potential*)

**Parameters**



- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.FixedPairListLennardJonesCapped(system, vl, potential)`

#### Parameters

- **system** –
- **vl** –
- **potential** –

`espressopp.interaction.FixedPairListLennardJonesCapped.getPotential()`

#### Return type

`espressopp.interaction.FixedPairListLennardJonesCapped.setPotential(potential)`

#### Parameters **potential** –

**class** `espressopp.interaction.LennardJonesCapped.LennardJonesCapped`

The Lennard-Jones potential.

## `espressopp.interaction.LennardJonesEnergyCapped`

$$V(r) = 4\varepsilon \left[ \left( \frac{\sigma}{r} \right)^{12} - \left( \frac{\sigma}{r} \right)^6 \right]$$

where  $r$  is either the distance or the capped distance, depending on which is greater.

`espressopp.interaction.LennardJonesEnergyCapped(epsilon, sigma, cutoff, caprad, shift)`

#### Parameters

- **epsilon** (*real*) – (default: 1.0)
- **sigma** (*real*) – (default: 1.0)
- **cutoff** – (default: infinity)
- **caprad** (*real*) – (default: 0.0)
- **shift** – (default: “auto”)

`espressopp.interaction.VerletListLennardJonesEnergyCapped(vl)`

#### Parameters **vl** –

`espressopp.interaction.VerletListLennardJonesEnergyCapped.getPotential(type1, type2)`

#### Parameters

- **type1** –
- **type2** –

#### Return type

`espressopp.interaction.VerletListLennardJonesEnergyCapped.setPotential(type1, type2, potential)`

#### Parameters

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.VerletListAdressLennardJonesEnergyCapped` (*vl, fixedtupleList*)

#### Parameters

- **vl** –
- **fixedtupleList** –

`espressopp.interaction.VerletListAdressLennardJonesEnergyCapped.getPotentialAT` (*type1, type2*)

#### Parameters

- **type1** –
- **type2** –

#### Return type

`espressopp.interaction.VerletListAdressLennardJonesEnergyCapped.getPotentialCG` (*type1, type2*)

#### Parameters

- **type1** –
- **type2** –

#### Return type

`espressopp.interaction.VerletListAdressLennardJonesEnergyCapped.setPotentialAT` (*type1, type2, potential*)

#### Parameters

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.VerletListAdressLennardJonesEnergyCapped.setPotentialCG` (*type1, type2, potential*)

#### Parameters

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.VerletListHadressLennardJonesEnergyCapped` (*vl, fixedtupleList*)

#### Parameters

- **vl** –
- **fixedtupleList** –

`espressopp.interaction.VerletListHadressLennardJonesEnergyCapped.getPotentialAT` (*type1*, *type2*)

**Parameters**

- **type1** –
- **type2** –

**Return type**

`espressopp.interaction.VerletListHadressLennardJonesEnergyCapped.getPotentialCG` (*type1*, *type2*)

**Parameters**

- **type1** –
- **type2** –

**Return type**

`espressopp.interaction.VerletListHadressLennardJonesEnergyCapped.setPotentialAT` (*type1*, *type2*, *potential*)

**Parameters**

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.VerletListHadressLennardJonesEnergyCapped.setPotentialCG` (*type1*, *type2*, *potential*)

**Parameters**

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.CellListLennardJonesEnergyCapped` (*stor*)

**Parameters stor –**

`espressopp.interaction.CellListLennardJonesEnergyCapped.getPotential` (*type1*, *type2*)

**Parameters**

- **type1** –
- **type2** –

**Return type**

`espressopp.interaction.CellListLennardJonesEnergyCapped.setPotential` (*type1*, *type2*, *potential*)

**Parameters**

- **type1** –

- **type2** –
- **potential** –

`espressopp.interaction.FixedPairListLennardJonesEnergyCapped` (*system*, *vl*, *potential*)

#### Parameters

- **system** –
- **vl** –
- **potential** –

`espressopp.interaction.FixedPairListLennardJonesEnergyCapped.getPotential` ()

#### Return type

`espressopp.interaction.FixedPairListLennardJonesEnergyCapped.setPotential` (*potential*)

#### Parameters **potential** –

**class** `espressopp.interaction.LennardJonesEnergyCapped.LennardJonesEnergyCapped`  
The Lennard-Jones potential.

## `espressopp.interaction.LennardJonesExpand`

$$V(r) = 4\varepsilon \left[ \left( \frac{\sigma}{r} \right)^{12} - \left( \frac{\sigma}{r} \right)^6 \right]$$

`espressopp.interaction.LennardJonesExpand` (*epsilon*, *sigma*, *delta*, *cutoff*, *shift*)

#### Parameters

- **epsilon** (*real*) – (default: 1.0)
- **sigma** (*real*) – (default: 1.0)
- **delta** (*real*) – (default: 0.0)
- **cutoff** – (default: infinity)
- **shift** – (default: “auto”)

`espressopp.interaction.VerletListLennardJonesExpand` (*vl*)

#### Parameters **vl** –

`espressopp.interaction.VerletListLennardJonesExpand.getPotential` (*type1*, *type2*)

#### Parameters

- **type1** –
- **type2** –

#### Return type

`espressopp.interaction.VerletListLennardJonesExpand.setPotential` (*type1*, *type2*, *potential*)

#### Parameters

- **type1** –
- **type2** –

- **potential** –

`espressopp.interaction.CellListLennardJonesExpand(stor)`

**Parameters stor** –

`espressopp.interaction.CellListLennardJonesExpand.setPotential(type1, type2, potential)`

**Parameters**

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.FixedPairListLennardJonesExpand(system, vl, potential)`

**Parameters**

- **system** –
- **vl** –
- **potential** –

`espressopp.interaction.FixedPairListLennardJonesExpand.setPotential(potential)`

**Parameters potential** –

**class** `espressopp.interaction.LennardJonesExpand.LennardJonesExpand`

The LennardJonesExpand potential.

## espressopp.interaction.LennardJonesGeneric

This class provides methods to compute forces and energies of a generic Lennard Jones potential with arbitrary integers a and b.

$$V(r) = 4\epsilon \left[ \left( \frac{\sigma}{r} \right)^a - \left( \frac{\sigma}{r} \right)^b \right]$$

`espressopp.interaction.LennardJonesGeneric(epsilon, sigma, a, b, cutoff, shift)`

**Parameters**

- **epsilon** (*real*) – (default: 1.0)
- **sigma** (*real*) – (default: 1.0)
- **a** (*int*) – (default: 12)
- **b** (*int*) – (default: 6)
- **cutoff** – (default: infinity)
- **shift** – (default: “auto”)

`espressopp.interaction.VerletListLennardJonesGeneric(vl)`

**Parameters vl** –

`espressopp.interaction.VerletListLennardJonesGeneric.getPotential(type1, type2)`

**Parameters**

- **type1** –
- **type2** –

### Return type

`espressopp.interaction.VerletListLennardJonesGeneric.getVerletList()`

**Return type** A Python list of lists.

`espressopp.interaction.VerletListLennardJonesGeneric.setPotential(type1,  
type2,  
potential)`

### Parameters

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.VerletListAdressLennardJonesGeneric(vl, fixedtupleList)`

### Parameters

- **vl** –
- **fixedtupleList** –

`espressopp.interaction.VerletListAdressLennardJonesGeneric.setPotentialAT(type1,  
type2,  
po-  
ten-  
tial)`

### Parameters

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.VerletListAdressLennardJonesGeneric.setPotentialCG(type1,  
type2,  
po-  
ten-  
tial)`

### Parameters

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.VerletListAdressLennardJonesGeneric2(vl, fixedtu-  
pleList)`

### Parameters

- **vl** –
- **fixedtupleList** –

`espressopp.interaction.VerletListAdressLennardJonesGeneric2.setPotentialAT(type1,  
type2,  
po-  
ten-  
tial)`

### Parameters

- **type1** –

- **type2** –
- **potential** –

`espressopp.interaction.VerletListAdressLennardJonesGeneric2.setPotentialCG` (*type1*,  
*type2*,  
*po-*  
*ten-*  
*tial*)

#### Parameters

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.VerletListHadressLennardJonesGeneric` (*vl*, *fixedtu-*  
*pleList*)

#### Parameters

- **vl** –
- **fixedtupleList** –

`espressopp.interaction.VerletListHadressLennardJonesGeneric.setPotentialAT` (*type1*,  
*type2*,  
*po-*  
*ten-*  
*tial*)

#### Parameters

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.VerletListHadressLennardJonesGeneric.setPotentialCG` (*type1*,  
*type2*,  
*po-*  
*ten-*  
*tial*)

#### Parameters

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.VerletListHadressLennardJonesGeneric2` (*vl*, *fixedtu-*  
*pleList*, *KTI*)

#### Parameters

- **vl** –
- **fixedtupleList** –
- **KTI** – (default: False)

`espressopp.interaction.VerletListHadressLennardJonesGeneric2.setPotentialAT` (*type1*,  
*type2*,  
*po-*  
*ten-*  
*tial*)

#### Parameters

- **type1** –
- **type2** –
- **potential** –

```
espressopp.interaction.VerletListHadressLennardJonesGeneric2.setPotentialCG(type1,
                                                                    type2,
                                                                    po-
                                                                    ten-
                                                                    tial)
```

#### Parameters

- **type1** –
- **type2** –
- **potential** –

```
espressopp.interaction.CellListLennardJonesGeneric(stor)
```

#### Parameters stor –

```
espressopp.interaction.CellListLennardJonesGeneric.setPotential(type1,
                                                                    type2,
                                                                    potential)
```

#### Parameters

- **type1** –
- **type2** –
- **potential** –

```
espressopp.interaction.FixedPairListLennardJonesGeneric(system, vl, potential)
```

#### Parameters

- **system** –
- **vl** –
- **potential** –

```
espressopp.interaction.FixedPairListLennardJonesGeneric.getFixedPairList()
```

**Return type** A Python list of lists.

```
espressopp.interaction.FixedPairListLennardJonesGeneric.getPotential()
```

**Return type**

```
espressopp.interaction.FixedPairListLennardJonesGeneric.setFixedPairList(fixedpairlist)
```

**Parameters fixedpairlist** –

```
espressopp.interaction.FixedPairListLennardJonesGeneric.setPotential(potential)
```

**Parameters potential** –

```
class espressopp.interaction.LennardJonesGeneric.LennardJonesGeneric
    The generic Lennard-Jones potential.
```

## espressopp.interaction.LennardJonesGromacs

if  $d^2 > r_1^2$

$$U = 4\epsilon \left( \frac{\sigma^{12}}{d^{12}} - \frac{\sigma^6}{d^6} \right) + (d - r_1)^3 (ljsw3 + ljsw4(d - r_1) + ljsw5)$$



else

$$U = 4\epsilon\left(\frac{\sigma^{12}}{d^{12}} - \frac{\sigma^6}{d^6}\right)$$

`espressopp.interaction.LennardJonesGromacs` (*epsilon*, *sigma*, *r1*, *cutoff*, *shift*)

#### Parameters

- **epsilon** (*real*) – (default: 1.0)
- **sigma** (*real*) – (default: 1.0)
- **r1** (*real*) – (default: 0.0)
- **cutoff** – (default: infinity)
- **shift** – (default: “auto”)

`espressopp.interaction.VerletListLennardJonesGromacs` (*vl*)

#### Parameters *vl* –

`espressopp.interaction.VerletListLennardJonesGromacs.getPotential` (*type1*,  
*type2*)

#### Parameters

- **type1** –
- **type2** –

#### Return type

`espressopp.interaction.VerletListLennardJonesGromacs.setPotential` (*type1*,  
*type2*,  
*potential*)

#### Parameters

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.CellListLennardJonesGromacs` (*stor*)

#### Parameters *stor* –

`espressopp.interaction.CellListLennardJonesGromacs.setPotential` (*type1*,  
*type2*,  
*potential*)

#### Parameters

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.FixedPairListLennardJonesGromacs` (*system*, *vl*, *potential*)

#### Parameters

- **system** –
- **vl** –
- **potential** –

`espressopp.interaction.FixedPairListLennardJonesGromacs.setPotential` (*potential*)

### Parameters `potential` –

`class espressopp.interaction.LennardJonesGromacs.LennardJonesGromacs`  
The LennardJonesGromacs potential.

## `espressopp.interaction.LennardJonesSoftcoreTI`

This module is for performing simulations (e.g. as part of Thermodynamic Integration) where some Lennard-Jones interactions are a function of a parameter  $\lambda$ , used to construct a pathway between states A and B.

For those interactions which are a function of  $\lambda$ , the potential is softcore Lennard Jones with the following form:

$$U_S(r_{ij}, \lambda) = (1 - \lambda)U_H^A(r_A) + \lambda U_H^B(r_B)$$

$$r_A = (\alpha \sigma_A^6 \lambda^p + r_{ij}^6)^{1/6}$$

$$r_B = (\alpha \sigma_B^6 (1 - \lambda)^p + r_{ij}^6)^{1/6}$$

where  $\epsilon_A$ ,  $\epsilon_B$ ,  $\sigma_A$  and  $\sigma_B$  are the parameters of states A and B, and  $\alpha$  and  $p$  are adjustable parameters of the softcore potential. The potentials  $U_H^A(r_A)$  and  $U_H^B(r_B)$  are the normal Lennard-Jones 12-6 hardcore potentials:

$$U_H^A(r_A) = 4.0\epsilon_A \left( \frac{\sigma_A^{12}}{r_A^{12}} - \frac{\sigma_A^6}{r_A^6} \right)$$

The user specifies a list of particles, `pidlist`. For all pairs of particles with particletypes interacting via this potential, the LJ interaction between two particles *i* and *j* is calculated as follows:

**if (i not in pidlist) and (j not in pidlist):**  $U_H^A$  (full state A hardcore LJ interaction)

**if (i in pidlist) and (j in pidlist):**

**if annihilate==True:**  $U_S$  (softcore LJ interaction, function of lambda)

**if annihilate==False:**  $U_H^A$  (full state A hardcore LJ interaction)

**if (i in pidlist) xor (j in pidlist):**  $U_S$  (softcore LJ interaction, function of lambda)

The default is annihilation (interactions within `pidlist` are coupled to lambda, and cross-interactions between particles in `pidlist` and particles in the rest of the system are also coupled to lambda). The alternative is decoupling (only cross-interactions between particles in `pidlist` and particles in the rest of the system are coupled to lambda. Interactions within `pidlist` are not affected by the value of lambda.) If `annihilation==False`, then decoupling is performed. See: [http://www.alchemistry.org/wiki/Decoupling\\_and\\_annihilation](http://www.alchemistry.org/wiki/Decoupling_and_annihilation)

Exclusions apply as normal, i.e. interactions are only calculated for pairs of particles not already excluded.

This class does not do any automatic shifting of the potential.

So far only `VerletListAdressLennardJonesSoftcoreTI` is implemented, however `VerletListLennardJonesSoftcoreTI`, `VerletListHadressLennardJonesSoftcoreTI`, etc. can also be easily implemented.

The  $\lambda$  (`lambdaTI`) parameter used here should not be confused with the  $\lambda$  (`lambda_adr`) particle property used in AdResS simulations.

See also the Thermodynamic Integration tutorial.

Example python script:

```
>>> #value of lambda
>>> lambdaTI = 0.3
>>> #softcore parameters
>>> alphaSC = 0.5
>>> powerSC = 1.0
>>> #make list of indices of particles whose LJ parameters are different in TI states A and B
>>> pidlist = [1,2,3,4]
>>> #create interaction using VerletListAdress object and FixedTupleListAdress object
>>> lj_adres_interaction=espressopp.interaction.VerletListAdressLennardJonesSoftcoreTI(verletlist,
>>> #loop over list of all types for particles interacting with this atomistic potential
>>> for i in types:
```

```
>>> for k in types:
>>>     ljpot = espressopp.interaction.LennardJonesSoftcoreTI(epsilonA=epsA[i][k], sigmaA=sigA[i]
>>>     ljpot.addPids(pidlist)
>>>     lj_adres_interaction.setPotentialAT(type1=i, type2=k, potential=ljpot)
>>> system.addInteraction(lj_adres_interaction)
```

During the MD run, one can then calculate the derivative of the RF energy wrt lambda

```
>>> #calculate dU/dlambda
>>> dUdl = lj_adres_interaction.computeEnergyDeriv()
```

```
espressopp.interaction.LennardJonesSoftcoreTI(epsilonA, sigmaA, epsilonB,
sigmaB, alpha, power, cutoff,
lambdaTI, annihilate)
```

#### Parameters

- **epsilonA** (*real*) – (default: 1.0) LJ interaction parameter
- **sigmaA** (*real*) – (default: 1.0) LJ interaction parameter
- **epsilonB** (*real*) – (default: 0.0) LJ interaction parameter
- **sigmaB** (*real*) – (default: 1.0) LJ interaction parameter
- **alpha** (*real*) – (default: 1.0) softcore parameter
- **power** (*real*) – (default: 1.0) softcore parameter
- **cutoff** (*real*) – (default: infinity) interaction cutoff
- **lambdaTI** (*real*) – (default: 0.0) TI lambda parameter
- **annihilate** (*bool*) – (default: True) switch between annihilation and decoupling

```
espressopp.interaction.LennardJonesSoftcoreTI.addPids(pidlist)
```

**Parameters** *pidlist* (*python list*) – list of particle ids of particles whose interaction parameters differ in state A and B

```
espressopp.interaction.VerletListAdressLennardJones(vl, fixedtupleList)
```

#### Parameters

- **vl** (*VerletListAdress object*) – Verlet list
- **fixedtupleList** (*FixedTupleListAdress object*) – list of tuples describing mapping between CG and AT particles

```
espressopp.interaction.VerletListAdressLennardJones.setPotentialAT(type1,
type2,
po-
ten-
tial)
```

#### Parameters

- **type1** (*int*) – atomtype
- **type2** (*int*) – atomtype
- **potential** (*Potential*) – espressopp potential

```
espressopp.interaction.VerletListAdressLennardJones.setPotentialCG(type1,
type2,
po-
ten-
tial)
```

#### Parameters

- **type1** (*int*) – atomtype

- **type2** (*int*) – atomtype
- **potential** (*Potential*) – espressopp potential

**class** espressopp.interaction.LennardJonesSoftcoreTI.**LennardJonesSoftcoreTI**  
 The Lennard-Jones potential.

## espressopp.interaction.LJcos

if  $r^2 \leq border_{pot}$ , then:

$$U = 4\left(\frac{1}{r^{12}} - \frac{1}{r^6}\right) + 1 - \phi$$

else:

$$U = \frac{1}{2}\phi(\cos(\alpha r^2 + \beta) - 1)$$

espressopp.interaction.**LJcos** (*phi*)

**Parameters** **phi** (*real*) – (default: 1.0)

espressopp.interaction.**VerletListLJcos** (*vl*)

**Parameters** **vl** –

espressopp.interaction.VerletListLJcos.**getPotential** (*type1*, *type2*)

**Parameters**

- **type1** –
- **type2** –

**Return type**

espressopp.interaction.VerletListLJcos.**getVerletList** ()

**Return type** A Python list of lists.

espressopp.interaction.VerletListLJcos.**setPotential** (*type1*, *type2*, *potential*)

**Parameters**

- **type1** –
- **type2** –
- **potential** –

espressopp.interaction.**VerletListAdressLJcos** (*vl*, *fixedtupleList*)

**Parameters**

- **vl** –
- **fixedtupleList** –

espressopp.interaction.VerletListAdressLJcos.**setPotentialAT** (*type1*, *type2*, *potential*)

**Parameters**

- **type1** –
- **type2** –
- **potential** –

espressopp.interaction.VerletListAdressLJcos.**setPotentialCG** (*type1*, *type2*, *potential*)

#### Parameters

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.VerletListHadressLJcos (vl, fixedtupleList)`

#### Parameters

- **vl** –
- **fixedtupleList** –

`espressopp.interaction.VerletListHadressLJcos.setPotentialAT (type1, type2, potential)`

#### Parameters

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.VerletListHadressLJcos.setPotentialCG (type1, type2, potential)`

#### Parameters

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.CellListLJcos (stor)`

#### Parameters stor –

`espressopp.interaction.CellListLJcos.setPotential (type1, type2, potential)`

#### Parameters

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.FixedPairListLJcos (system, vl, potential)`

#### Parameters

- **system** –
- **vl** –
- **potential** –

`espressopp.interaction.FixedPairListLJcos.getFixedPairList ()`

**Return type** A Python list of lists.

`espressopp.interaction.FixedPairListLJcos.setFixedPairList (fixedpairlist)`

#### Parameters fixedpairlist –

`espressopp.interaction.FixedPairListLJcos.setPotential (potential)`

#### Parameters potential –

**class** `espressopp.interaction.LJcos.LJcos`

The Lennard-Jones potential.

## espressopp.interaction.MirrorLennardJones

This class provides methods to compute forces and energies of the Mirror Lennard-Jones potential.

$$V(r) = V_{LJ}(r_m - |r - r_m|)$$

where  $V_{LJ}$  is the 6-12 purely repulsive Lennard-Jones potential. This potential is introduced in R.L.C. Akkermans, S. Toxvaerd and & W. J. Briels. Molecular dynamics of polymer growth. The Journal of Chemical Physics, 1998, 109, 2929-2940.

`espressopp.interaction.MirrorLennardJones` (*epsilon*, *sigma*)

### Parameters

- **epsilon** (*real*) – (default: 1.0)
- **sigma** (*real*) – (default: 0.0)

`espressopp.interaction.FixedPairListMirrorLennardJones` (*system*, *vl*, *potential*)

### Parameters

- **system** –
- **vl** –
- **potential** –

`espressopp.interaction.FixedPairListMirrorLennardJones.getFixedPairList` ()

**Return type** A Python list of lists.

`espressopp.interaction.FixedPairListMirrorLennardJones.getPotential` ()

**Return type**

`espressopp.interaction.FixedPairListMirrorLennardJones.setFixedPairList` (*fixedpairlist*)

**Parameters** **fixedpairlist** –

`espressopp.interaction.FixedPairListMirrorLennardJones.setPotential` (*potential*)

**Parameters** **potential** –

**class** `espressopp.interaction.MirrorLennardJones.MirrorLennardJones`

The MirrorLennardJones potential.

## espressopp.interaction.Morse

This class provides methods to compute forces and energies of the Morse potential.

$$U = \varepsilon \left( e^{-2\alpha(r-r_{min})} - 2e^{-\alpha(r-r_{min})} \right)$$

`espressopp.interaction.Morse` (*epsilon*, *alpha*, *rMin*, *cutoff*, *shift*)

### Parameters

- **epsilon** (*real*) – (default: 1.0)
- **alpha** (*real*) – (default: 1.0)
- **rMin** (*real*) – (default: 0.0)
- **cutoff** – (default: infinity)
- **shift** – (default: “auto”)

`espressopp.interaction.VerletListMorse` (*vl*)

**Parameters** **vl** –

`espressopp.interaction.VerletListMorse.getPotential (type1, type2)`

**Parameters**

- **type1** –
- **type2** –

**Return type**

`espressopp.interaction.VerletListMorse.setPotential (type1, type2, potential)`

**Parameters**

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.VerletListAdressMorse (vl, fixedtupleList)`

**Parameters**

- **vl** –
- **fixedtupleList** –

`espressopp.interaction.VerletListAdressMorse.setPotentialAT (type1, type2, potential)`

**Parameters**

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.VerletListAdressMorse.setPotentialCG (type1, type2, potential)`

**Parameters**

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.VerletListHadressMorse (vl, fixedtupleList)`

**Parameters**

- **vl** –
- **fixedtupleList** –

`espressopp.interaction.VerletListHadressMorse.setPotentialAT (type1, type2, potential)`

**Parameters**

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.VerletListHadressMorse.setPotentialCG (type1, type2, potential)`

**Parameters**

- **type1** –
- **type2** –

- **potential** –

`espressopp.interaction.CellListMorse` (*stor*)

**Parameters** *stor* –

`espressopp.interaction.CellListMorse.setPotential` (*type1*, *type2*, *potential*)

**Parameters**

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.FixedPairListMorse` (*system*, *vl*, *potential*)

**Parameters**

- **system** –
- **vl** –
- **potential** –

`espressopp.interaction.FixedPairListMorse.setPotential` (*potential*)

**Parameters** *potential* –

**class** `espressopp.interaction.Morse.Morse`  
The Morse potential.

## espressopp.interaction.OPLS

This class provides methods to compute forces and energies of the OPLS dihedral potential. To create a new dihedral potential.

$$U = \sum_{j=1}^4 K_j (1 + \cos(j\phi))$$

`espressopp.interaction.OPLS` (*K1*, *K2*, *K3*, *K4*)

**Parameters**

- **K1** (*real*) – (default: 1.0)
- **K2** (*real*) – (default: 0.0)
- **K3** (*real*) – (default: 0.0)
- **K4** (*real*) – (default: 0.0)

`espressopp.interaction.FixedQuadrupleListOPLS` (*system*, *vl*, *potential*)

**Parameters**

- **system** –
- **vl** –
- **potential** –

`espressopp.interaction.FixedQuadrupleListOPLS.setPotential` (*type1*, *type2*, *potential*)

**Parameters**

- **type1** –
- **type2** –



- **potential** –

**class** `espressopp.interaction.OPLS.OPLS`  
 The OPLS potential.

## **espressopp.interaction.Potential**

This is an abstract class, only needed to be inherited from.

`espressopp.interaction.Potential.computeEnergy (*args)`

**Parameters** *\*args* –

**Return type**

`espressopp.interaction.Potential.computeForce (*args)`

**Parameters** *\*args* –

**Return type**

## **espressopp.interaction.PotentialUniqueDist**

This is an abstract class, only needed to be inherited from.

`espressopp.interaction.PotentialUniqueDist.computeEnergy (*args)`

**Parameters** *\*args* –

**Return type**

`espressopp.interaction.PotentialUniqueDist.computeForce (*args)`

**Parameters** *\*args* –

**Return type**

## **espressopp.interaction.PotentialVSpherePair**

This is an abstract class, only needed to be inherited from.

`espressopp.interaction.PotentialVSpherePair.computeEnergy (*args)`

**Parameters** *\*args* –

**Return type**

`espressopp.interaction.PotentialVSpherePair.computeForce (*args)`

**Parameters** *\*args* –

**Return type**

## **espressopp.interaction.Quartic**

This class provides methods to compute forces and energies of the Quartic potential.

$$U = \frac{K}{4} (d^2 - r_0^2)^2$$

`espressopp.interaction.Quartic (K, r0, cutoff, shift)`

**Parameters**

- **K** (*real*) – (default: 1.0)

- **r0** (*real*) – (default: 0.0)
- **cutoff** – (default: infinity)
- **shift** (*real*) – (default: 0.0)

`espressopp.interaction.FixedPairListQuartic` (*system*, *vl*, *potential*)

#### Parameters

- **system** –
- **vl** –
- **potential** –

`espressopp.interaction.FixedPairListQuartic.getFixedPairList` ()

**Return type** A Python list of lists.

`espressopp.interaction.FixedPairListQuartic.setFixedPairList` (*fixedpairlist*)

**Parameters** **fixedpairlist** –

`espressopp.interaction.FixedPairListQuartic.setPotential` (*type1*, *type2*, *potential*)

#### Parameters

- **type1** –
- **type2** –
- **potential** –

`class espressopp.interaction.Quartic.Quartic`

The Quartic potential.

## espressopp.interaction.ReactionFieldGeneralized

This class provides methods to compute forces and energies of the generalized reaction field.

$$U = PQ \left( \frac{1}{d} - \frac{\left( 1 + \frac{(\varepsilon_1 - 4\varepsilon_2)(1 + \kappa r_c) - 2\varepsilon_2 \kappa r_c^2}{(\varepsilon_1 + 2\varepsilon_2)(1 + \kappa r_c) + \varepsilon_2 \kappa r_c^2} \right)}{r_c^3} \cdot d^2 - \frac{3\varepsilon_2}{r_c(2\varepsilon_2 + 1)} \right)$$

where  $P$  is a prefactor,  $Q$  is the product of the charges of the two particles,  $d$  is their distance from each other, and  $r_c$  the cutoff-radius.

`espressopp.interaction.ReactionFieldGeneralized` (*prefactor*, *kappa*, *epsilon1*, *epsilon2*, *cutoff*, *shift*)

#### Parameters

- **prefactor** (*real*) – (default: 1.0)
- **kappa** (*real*) – (default: 0.0)
- **epsilon1** (*real*) – (default: 1.0)
- **epsilon2** (*real*) – (default: 80.0)
- **cutoff** – (default: infinity)
- **shift** – (default: “auto”)

`espressopp.interaction.VerletListReactionFieldGeneralized` (*vl*)

**Parameters** **vl** –

`espressopp.interaction.VerletListReactionFieldGeneralized.getPotential` (*type1*, *type2*)

#### Parameters

- **type1** –
- **type2** –

#### Return type

`espressopp.interaction.VerletListReactionFieldGeneralized.setPotential` (*type1*,  
*type2*,  
*po-*  
*ten-*  
*tial*)

#### Parameters

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.VerletListAdressReactionFieldGeneralized` (*vl*, *fixedtu-*  
*pleList*)

#### Parameters

- **vl** –
- **fixedtupleList** –

`espressopp.interaction.VerletListAdressReactionFieldGeneralized.setPotentialAT` (*type1*,  
*type2*,  
*po-*  
*ten-*  
*tial*)

#### Parameters

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.VerletListAdressReactionFieldGeneralized.setPotentialCG` (*type1*,  
*type2*,  
*po-*  
*ten-*  
*tial*)

#### Parameters

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.VerletListHadressReactionFieldGeneralized` (*vl*,  
*fixedtu-*  
*pleList*)

#### Parameters

- **vl** –
- **fixedtupleList** –

```

esspressopp.interaction.VerletListHadressReactionFieldGeneralized.setPotentialAT(type1,
                                                                    type2,
                                                                    po-
                                                                    ten-
                                                                    tial)

```

#### Parameters

- **type1** –
- **type2** –
- **potential** –

```

esspressopp.interaction.VerletListHadressReactionFieldGeneralized.setPotentialCG(type1,
                                                                    type2,
                                                                    po-
                                                                    ten-
                                                                    tial)

```

#### Parameters

- **type1** –
- **type2** –
- **potential** –

```

esspressopp.interaction.CellListReactionFieldGeneralized(stor)

```

#### Parameters stor –

```

esspressopp.interaction.CellListReactionFieldGeneralized.setPotential(type1,
                                                                    type2,
                                                                    po-
                                                                    ten-
                                                                    tial)

```

#### Parameters

- **type1** –
- **type2** –
- **potential** –

```

class espressopp.interaction.ReactionFieldGeneralized.ReactionFieldGeneralized
    The ReactionFieldGeneralized potential.

```

## esspressopp.interaction.ReactionFieldGeneralizedTI

This module is for performing simulations (e.g. as part of Thermodynamic Integration) where some interactions are a linear function of a parameter  $\lambda$ .

$$U(\lambda) = (1 - \lambda)U_C^A$$

where  $U_C^A$  is the standard Reaction Field interaction. This allows one to perform TI where the charges in TI state A ( $\lambda = 0$ ) are the particle charges contained in the particle property `charge` and the charges in TI state B ( $\lambda = 1$ ) are zero.

The user specifies a list of particles, `pidlist`. For all pairs of particles with `particletypes` interacting via this potential, the RF interaction between two particles `i` and `j` is calculated as follows:

**if (i not in pidlist) and (j not in pidlist):**  $U_{RF}$  (full RF interaction)

**if (i in pidlist) and (j in pidlist):**

**if annihilate==True:**  $(1 - \lambda)U_{RF}$  (RF interaction scaled by 1-lambda)

**if annihilate==False:**  $U_{RF}$  (full RF interaction)

**if (i in pidlist) xor (j in pidlist):**  $(1 - \lambda)U_{RF}$  (RF interaction scaled by 1-lambda)

The default is annihilation (completely turning off charges of particles in pidlist in state B, so that interactions within pidlist are turned off and also cross-interactions between particles in pidlist and particles in the rest of the system). The alternative is decoupling (only cross-interactions between particles in pidlist and particles in the rest of the system are turned off. Interactions within pidlist are not affected.) If annihilate==False, then decoupling is performed. See: [http://www.alchemistry.org/wiki/Decoupling\\_and\\_annihilation](http://www.alchemistry.org/wiki/Decoupling_and_annihilation)

Exclusions apply as normal, i.e. interactions are only calculated for pairs of particles not already excluded.

So far only VerletListAdressReactionFieldGeneralizedTI is implemented, however VerletListReactionFieldGeneralizedTI, VerletListHadressReactionFieldGeneralizedTI, etc. can also be easily implemented.

The  $\lambda$  (lambdaTI) parameter used here should not be confused with the  $\lambda$  (lambda\_adr) particle property used in AdResS simulations.

See also the Thermodynamic Integration tutorial.

Example python script:

```
>>> #value of lambda
>>> lambdaTI = 0.3
>>> #construct RF potential with parameters prefactor, kappa, epsilon1, epsilon2, cutoff as in standard
>>> pot = espressopp.interaction.ReactionFieldGeneralizedTI(prefactor=prefactor, kappa=kappa, eps
>>> #add list of indices of particles whose charge is 0 in TI state B
>>> pidlist = [1,2,3,4]
>>> pot.addPids(pidlist)
>>> #create interaction using VerletListAdress object and FixedTupleListAdress object
>>> qq_adres_interaction=espressopp.interaction.VerletListAdressReactionFieldGeneralizedTI(verlet
>>> #loop over list of all types for particles interacting with this atomistic potential
>>> for i in types:
>>>     for k in types:
>>>         qq_adres_interaction.setPotentialAT(type1=i, type2=k, potential=pot)
>>> system.addInteraction(qq_adres_interaction)
```

During the MD run, one can then calculate the derivative of the RF energy wrt lambda

```
>>> #calculate dU/dlambda
>>> dUdl = qq_adres_interaction.computeEnergyDeriv()
```

espressopp.interaction.**ReactionFieldGeneralizedTI** (*prefactor, kappa, epsilon1, epsilon2, cutoff, lambdaTI, annihilate*)

#### Parameters

- **prefactor** (*real*) – (default: 1.0) RF parameter
- **kappa** (*real*) – (default: 0.0) RF parameter
- **epsilon1** (*real*) – (default: 1.0) RF parameter
- **epsilon2** (*real*) – (default: 80.0) RF parameter
- **cutoff** (*real*) – (default: infinity) interaction cutoff
- **lambdaTI** (*real*) – (default: 0.0) TI lambda parameter
- **annihilate** (*bool*) – (default: True) switch between annihilation and decoupling

espressopp.interaction.ReactionFieldGeneralizedTI.**addPids** (*pidlist*)

**Parameters** *pidlist* (*python list*) – list of particle ids of particles whose charge is zero in state B

espressopp.interaction.**VerletListAdressReactionFieldGeneralized** (*vl, fixedtupleList*)

#### Parameters

- **vl** (*VerletListAdress object*) – Verlet list
- **fixedtupleList** (*FixedTupleListAdress object*) – list of tuples describing mapping between CG and AT particles

`espressopp.interaction.VerletListAdressReactionFieldGeneralized.setPotentialAT` (*type1*, *type2*, *potential*)

#### Parameters

- **type1** (*int*) – atomtype
- **type2** (*int*) – atomtype
- **potential** (*Potential*) – espressopp potential

`espressopp.interaction.VerletListAdressReactionFieldGeneralized.setPotentialCG` (*type1*, *type2*, *potential*)

#### Parameters

- **type1** (*int*) – atomtype
- **type2** (*int*) – atomtype
- **potential** (*Potential*) – espressopp potential

`class espressopp.interaction.ReactionFieldGeneralizedTI.ReactionFieldGeneralizedTI`  
The ReactionFieldGeneralizedTI potential.

## espressopp.interaction.SingleParticlePotential

This class is used to define single-particle interactions, typically used for external forces on the system.

The potential may depend on any of the particle properties (type, mass, etc.).

`espressopp.interaction.SingleParticlePotential.computeEnergy` (*position*, *bc*)

#### Parameters

- **position** –
- **bc** –

#### Return type

`espressopp.interaction.SingleParticlePotential.computeForce` (*position*, *bc*)

#### Parameters

- **position** –
- **bc** –

#### Return type

## espressopp.interaction.SoftCosine

This class provides methods to compute forces and energies of the SoftCosine potential.

$$V(r) = A \left[ 1.0 + \cos \left( \frac{\pi r}{r_c} \right) \right]$$

`espressopp.interaction.SoftCosine` (*A, cutoff, shift*)

### Parameters

- **A** (*real*) – (default: 1.0)
- **cutoff** – (default: infinity)
- **shift** – (default: “auto”)

`espressopp.interaction.VerletListSoftCosine` (*stor*)

### Parameters stor –

`espressopp.interaction.VerletListSoftCosine.setPotential` (*type1, type2, potential*)

### Parameters

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.CellListSoftCosine` (*stor*)

### Parameters stor –

`espressopp.interaction.CellListSoftCosine.setPotential` (*type1, type2, potential*)

### Parameters

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.FixedPairListSoftCosine` (*system, vl, potential*)

### Parameters

- **system** –
- **vl** –
- **potential** –

`espressopp.interaction.FixedPairListSoftCosine.setPotential` (*potential*)

### Parameters potential –

**class** `espressopp.interaction.SoftCosine.SoftCosine`

The SoftCosine potential.

## espressopp.interaction.StillingerWeberPairTerm

This class provides methods to compute forces and energies of 2 body term of Stillinger-Weber potential.

$$U = \varepsilon A \left[ \frac{d^{-p}}{\sigma} (B - 1) \right] \exp \left( \frac{1}{\frac{d}{\sigma} - r_c} \right)$$

where  $r_c$  is the cutoff-radius.

`espressopp.interaction.StillingerWeberPairTerm(A, B, p, q, epsilon, sigma, cutoff)`

#### Parameters

- **A** –
- **B** –
- **p** –
- **q** –
- **epsilon** (*real*) – (default: 1.0)
- **sigma** (*real*) – (default: 1.0)
- **cutoff** – (default: infinity)

`espressopp.interaction.VerletListStillingerWeberPairTerm(vl)`

#### Parameters **vl** –

`espressopp.interaction.VerletListStillingerWeberPairTerm.getPotential` (*type1*,  
*type2*)

#### Parameters

- **type1** –
- **type2** –

#### Return type

`espressopp.interaction.VerletListStillingerWeberPairTerm.getVerletList` ()

#### Return type

A Python list of lists.

`espressopp.interaction.VerletListStillingerWeberPairTerm.setPotential` (*type1*,  
*type2*,  
*po-*  
*ten-*  
*tial*)

#### Parameters

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.VerletListAdressStillingerWeberPairTerm(vl, fixedtu-`  
`pleList)`

#### Parameters

- **vl** –
- **fixedtupleList** –

`espressopp.interaction.VerletListAdressStillingerWeberPairTerm.setPotentialAT` (*type1*,  
*type2*,  
*po-*  
*ten-*  
*tial*)

#### Parameters

- **type1** –
- **type2** –
- **potential** –



`espressopp.interaction.VerletListAdressStillingerWeberPairTerm.setPotentialCG` (*type1*, *type2*, *potential*)

#### Parameters

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.VerletListHadressStillingerWeberPairTerm` (*vl*, *fixedtupleList*)

#### Parameters

- **vl** –
- **fixedtupleList** –

`espressopp.interaction.VerletListHadressStillingerWeberPairTerm.setPotentialAT` (*type1*, *type2*, *potential*)

#### Parameters

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.VerletListHadressStillingerWeberPairTerm.setPotentialCG` (*type1*, *type2*, *potential*)

#### Parameters

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.CellListStillingerWeberPairTerm` (*stor*)

#### Parameters stor –

`espressopp.interaction.CellListStillingerWeberPairTerm.setPotential` (*type1*, *type2*, *potential*)

#### Parameters

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.FixedPairListStillingerWeberPairTerm` (*system*, *vl*, *potential*)

#### Parameters

- **system** –
- **vl** –
- **potential** –

`espressopp.interaction.FixedPairListStillingerWeberPairTerm.setPotential (potential)`

**Parameters** **potential** –

**class** `espressopp.interaction.StillingerWeberPairTerm.StillingerWeberPairTerm`  
The Lennard-Jones potential.

## `espressopp.interaction.StillingerWeberPairTermCapped`

This class provides methods to compute forces and energies of 2 body term of Stillinger-Weber potential.

If the distance is smaller than the cap-radius:

$$U = A[d_{12}^{-p}(B - 1)]e^{\frac{1}{d_{12}-r_c}}$$

where  $r_c$  is the cutoff-radius.

`espressopp.interaction.StillingerWeberPairTermCapped (A, B, p, q, epsilon, sigma, cutoff, caprad)`

**Parameters**

- **A** –
- **B** –
- **p** –
- **q** –
- **epsilon** (*real*) – (default: 1.0)
- **sigma** (*real*) – (default: 1.0)
- **cutoff** – (default: infinity)
- **caprad** (*real*) – (default: 0.0)

`espressopp.interaction.VerletListStillingerWeberPairTermCapped (vl)`

**Parameters** **vl** –

`espressopp.interaction.VerletListStillingerWeberPairTermCapped.getCaprad ()`

**Return type**

`espressopp.interaction.VerletListStillingerWeberPairTermCapped.getPotential (type1, type2)`

**Parameters**

- **type1** –
- **type2** –

**Return type**

`espressopp.interaction.VerletListStillingerWeberPairTermCapped.getVerletList ()`

**Return type** A Python list of lists.

`espressopp.interaction.VerletListStillingerWeberPairTermCapped.setPotential (type1, type2, potential)`

#### Parameters

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.VerletListAdressStillingerWeberPairTermCapped` (*vl*,  
*fixed-*  
*tu-*  
*pleList*)

#### Parameters

- **vl** –
- **fixedtupleList** –

`espressopp.interaction.VerletListAdressStillingerWeberPairTermCapped.setPotentialAT` (*type1*  
*type2*  
*po-*  
*ten-*  
*tial*)

#### Parameters

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.VerletListAdressStillingerWeberPairTermCapped.setPotentialCG` (*type1*  
*type2*  
*po-*  
*ten-*  
*tial*)

#### Parameters

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.VerletListHadressStillingerWeberPairTermCapped` (*vl*,  
*fixed-*  
*tu-*  
*pleList*)

#### Parameters

- **vl** –
- **fixedtupleList** –

`espressopp.interaction.VerletListHadressStillingerWeberPairTermCapped.setPotentialAT` (*type1*  
*type2*  
*po-*  
*ten-*  
*tial*)

#### Parameters

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.VerletListHadressStillingerWeberPairTermCapped.setPotentialCG` (*type1, type2, potential*)

#### Parameters

- `type1` –
- `type2` –
- `potential` –

`espressopp.interaction.CellListStillingerWeberPairTermCapped` (*stor*)

#### Parameters `stor` –

`espressopp.interaction.CellListStillingerWeberPairTermCapped.setPotential` (*type1, type2, potential*)

#### Parameters

- `type1` –
- `type2` –
- `potential` –

`espressopp.interaction.FixedPairListStillingerWeberPairTermCapped` (*system, vl, potential*)

#### Parameters

- `system` –
- `vl` –
- `potential` –

`espressopp.interaction.FixedPairListStillingerWeberPairTermCapped.setPotential` (*potential*)

#### Parameters `potential` –

**class** `espressopp.interaction.StillingerWeberPairTermCapped.StillingerWeberPairTermCapped`  
The Lennard-Jones potential.

## `espressopp.interaction.StillingerWeberTripleTerm`

This class provides methods to compute forces and energies of the Stillinger Weber Triple Term potential.

if  $d_{12} \geq r_{c1}$  or  $d_{32} \geq r_{c2}$

$$U = 0.0$$

else

$$U = \epsilon \lambda e^{\frac{\sigma \gamma_1}{|r_{12}| - \sigma r_{c1}}} + \frac{\sigma \gamma_2}{|r_{32}| - \sigma r_{c2}} \left( \frac{r_{12} r_{32}}{|r_{12}| \cdot |r_{32}|} - \cos(\theta_0) \right)^2$$

`espressopp.interaction.StillingerWeberTripleTerm` (*gamma, theta0, lmbd, epsilon, sigma, cutoff*)

#### Parameters

- **gamma** (*real*) – (default: 0.0)
- **theta0** (*real*) – (default: 0.0)
- **lmbd** (*real*) – (default: 0.0)
- **epsilon** (*real*) – (default: 1.0)
- **sigma** (*real*) – (default: 1.0)
- **cutoff** – (default: infinity)

`espressopp.interaction.VerletListStillingerWeberTripleTerm` (*system*, *vl3*)

#### Parameters

- **system** –
- **vl3** –

`espressopp.interaction.VerletListStillingerWeberTripleTerm`.**getPotential** (*type1*,  
*type2*,  
*type3*)

#### Parameters

- **type1** –
- **type2** –
- **type3** –

#### Return type

`espressopp.interaction.VerletListStillingerWeberTripleTerm`.**getVerletListTriple** ()

**Return type** A Python list of lists.

`espressopp.interaction.VerletListStillingerWeberTripleTerm`.**setPotential** (*type1*,  
*type2*,  
*type3*,  
*po-*  
*ten-*  
*tial*)

#### Parameters

- **type1** –
- **type2** –
- **type3** –
- **potential** –

`espressopp.interaction.FixedTripleListStillingerWeberTripleTerm` (*system*, *ftl*,  
*potential*)

#### Parameters

- **system** –
- **ftl** –
- **potential** –

`espressopp.interaction.FixedTripleListStillingerWeberTripleTerm`.**getFixedTripleList** ()

**Return type** A Python list of lists.

`essopp.interaction.FixedTripleListStillingerWeberTripleTerm.setPotential` (*type1*,  
*type2*,  
*type3*,  
*po-*  
*ten-*  
*tial*)

#### Parameters

- **type1** –
- **type2** –
- **type3** –
- **potential** –

**class** `essopp.interaction.StillingerWeberTripleTerm.StillingerWeberTripleTerm`  
The StillingerWeberTripleTerm potential.

## essopp.interaction.Tabulated

`essopp.interaction.Tabulated` (*itype*, *filename*, *cutoff*)

#### Parameters

- **itype** –
- **filename** –
- **cutoff** – (default: infinity)

`essopp.interaction.VerletListAdressTabulated` (*vl*, *fixedtupleList*)

#### Parameters

- **vl** –
- **fixedtupleList** –

`essopp.interaction.VerletListAdressTabulated.setPotentialAT` (*type1*,  
*type2*,  
*potential*)

#### Parameters

- **type1** –
- **type2** –
- **potential** –

`essopp.interaction.VerletListAdressTabulated.setPotentialCG` (*type1*,  
*type2*,  
*potential*)

#### Parameters

- **type1** –
- **type2** –
- **potential** –

`essopp.interaction.VerletListHadressTabulated` (*vl*, *fixedtupleList*)

#### Parameters

- **vl** –
- **fixedtupleList** –

`espressopp.interaction.VerletListHadressTabulated.setPotentialAT` (*type1*,  
*type2*,  
*potential*)

**Parameters**

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.VerletListHadressTabulated.setPotentialCG` (*type1*,  
*type2*,  
*potential*)

**Parameters**

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.VerletListTabulated` (*vl*)

**Parameters vl –**

`espressopp.interaction.VerletListTabulated.getPotential` (*type1*, *type2*)

**Parameters**

- **type1** –
- **type2** –

**Return type**

`espressopp.interaction.VerletListTabulated.setPotential` (*type1*, *type2*, *potential*)

**Parameters**

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.CellListTabulated` (*stor*)

**Parameters stor –**

`espressopp.interaction.CellListTabulated.setPotential` (*type1*, *type2*, *potential*)

**Parameters**

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.FixedPairListTabulated` (*system*, *vl*, *potential*)

**Parameters**

- **system** –
- **vl** –
- **potential** –

`espressopp.interaction.FixedPairListTabulated.setPotential` (*potential*)

#### Parameters `potential` –

`espressopp.interaction.FixedPairListTypesTabulated` (*system*, *fil*)

#### Parameters

- **system** (`espressopp.System`) – The Espresso++ system object.
- **ftl** (`espressopp.FixedPairList`) – The FixedPair list.

`espressopp.interaction.FixedPairListTypesTabulated.setPotential` (*type1*,  
*type2*,  
*potential*)

Defines bond potential for interaction between particles of types *type1*-*type2*-*type3*.

#### Parameters

- **type1** (*int*) – Type of particle 1.
- **type2** (*int*) – Type of particle 2.
- **potential** (`espressopp.interaction.Potential`) – The potential to set up.

**class** `espressopp.interaction.Tabulated.Tabulated`  
The Tabulated potential.

## `espressopp.interaction.TabulatedAngular`

`espressopp.interaction.TabulatedAngular` (*itype*, *filename*)

#### Parameters

- **itype** (*int*) – The interpolation type: 1 - linear, 2 - akima spline, 3 - cubic spline
- **filename** (*str*) – The tabulated potential filename.

`espressopp.interaction.FixedTripleListTabulatedAngular` (*system*, *fil*, *potential*)

#### Parameters

- **system** (`espressopp.System`) – The Espresso++ system object.
- **ftl** (`espressopp.FixedTripleList`) – The FixedTripleList.
- **potential** (`espressopp.interaction.Potential`) – The potential.

`espressopp.interaction.FixedTripleListTabulatedAngular.setPotential` (*potential*)

**Parameters** **potential** (`espressopp.interaction.Potential`) – The potential object.

`espressopp.interaction.FixedTripleListTypesTabulatedAngular` (*system*, *fil*)

#### Parameters

- **system** (`espressopp.System`) – The Espresso++ system object.
- **ftl** (`espressopp.FixedTripleList`) – The FixedTriple list.

`espressopp.interaction.FixedTripleListTypesTabulatedAngular.setPotential` (*type1*,  
*type2*,  
*type3*,  
*po-*  
*ten-*  
*tial*)

Defines angular potential for interaction between particles of types *type1*-*type2*-*type3*.

#### Parameters

- **type1** (*int*) – Type of particle 1.



- **type2** (*int*) – Type of particle 2.
- **type3** (*int*) – Type of particle 3.
- **potential** (`espressopp.interaction.AngularPotential`) – The potential to set up.

**class** `espressopp.interaction.TabulatedAngular.TabulatedAngular`  
 The TabulatedAngular potential.

## `espressopp.interaction.TabulatedDihedral`

Calculates energies and forces for a dihedral tabulated potential. In the tabulated potential file, angles should be in radians, and the file should cover the range  $-\pi$  radians to  $+\pi$  radians ( $-180$  to  $+180$  degrees).

Note that this class has only been tested for symmetric tabulated potentials.

`espressopp.interaction.TabulatedDihedral` (*itype, filename*)

**Parameters** **itype** – The interpolation type: 1 - linear, 2 - akima spline, 3 - cubic spline :param  
 filename: The tabulated potential filename. :type itype: int :type filename: str

`espressopp.interaction.FixedQuadrupleListTabulatedDihedral` (*system, fql, potential*)

### Parameters

- **system** (`espressopp.System`) – The Espresso++ system object.
- **fql** (`espressopp.FixedQuadrupleList`) – The FixedQuadrupleList.
- **potential** (`espressopp.interaction.Potential`) – The potential.

`espressopp.interaction.FixedQuadrupleListTabulatedDihedral.setPotential` (*potential*)

**Parameters** **potential** (`espressopp.interaction.Potential`) – The potential object.

`espressopp.interaction.FixedQuadrupleListTypesTabulatedDihedral` (*system, fql*)

### Parameters

- **system** (`espressopp.System`) – The Espresso++ system object.
- **ftl** (`espressopp.FixedQuadrupleList`) – The FixedQuadrupleList list.

`espressopp.interaction.FixedQuadrupleListTypesTabulatedDihedral` (*system, fil*)

### Parameters

- **system** (`espressopp.System`) – The Espresso++ system object.
- **ftl** (`espressopp.FixedQuadrupleList`) – The FixedQuadruple list.

`espressopp.interaction.FixedQuadrupleListTypesTabulatedDihedral.setPotential` (*type1, type2, type3, type4, potential*)

Defines dihedral potential for interaction between particles of types type1-type2-type3-type4.

### Parameters

- **type1** (*int*) – Type of particle 1.
- **type2** (*int*) – Type of particle 2.

- **type3** (*int*) – Type of particle 3.
- **type4** (*int*) – Type of particle 4.
- **potential** (`espressopp.interaction.DihedralPotential`) – The potential to set up.

**class** `espressopp.interaction.TabulatedDihedral.TabulatedDihedral`  
 The TabulatedDihedral potential.

## `espressopp.interaction.TersoffPairTerm`

This class provides methods to compute forces and energies of 2 body term of Tersoff potential.

if  $d_{12} > R + D$

$$U = 0$$

if  $d_{12} < R - D$

$$U = Ae^{-\lambda_1 d_{12}}$$

else

$$U = \frac{1}{2} \left( 1 - \sin \left( \frac{\pi}{4D} (d_{12} - R) \right) \right) Ae^{-\lambda_1 d_{12}}$$

`espressopp.interaction.TersoffPairTerm` (*A, lambda1, R, D, cutoff*)

### Parameters

- **A** –
- **lambda1** –
- **R** –
- **D** –
- **cutoff** – (default: infinity)

`espressopp.interaction.VerletListTersoffPairTerm` (*vl*)

### Parameters **vl** –

`espressopp.interaction.VerletListTersoffPairTerm.getPotential` (*type1, type2*)

### Parameters

- **type1** –
- **type2** –

### Return type

`espressopp.interaction.VerletListTersoffPairTerm.getVerletList` ()

**Return type** A Python list of lists.

`espressopp.interaction.VerletListTersoffPairTerm.setPotential` (*type1, type2, potential*)

### Parameters

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.CellListTersoffPairTerm` (*stor*)

### Parameters stor –

`espressopp.interaction.CellListTersoffPairTerm.setPotential` (*type1*, *type2*, *potential*)

### Parameters

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.FixedPairListTersoffPairTerm` (*system*, *vl*, *potential*)

### Parameters

- **system** –
- **vl** –
- **potential** –

`espressopp.interaction.FixedPairListTersoffPairTerm.setPotential` (*potential*)

### Parameters potential –

`class espressopp.interaction.TersoffPairTerm.TersoffPairTerm`

The Lennard-Jones potential.

## espressopp.interaction.TersoffTripleTerm

This class provides methods to compute forces and energies of the Tersoff Triple Term potential.

$$U = f_{C_j} f_A \left( 1 + \left( \beta f_{C_k} \gamma \left( 1 + \frac{c_2}{d_2} - \frac{c_2}{d_2 + \left( \frac{r_{12} r_{32}}{|r_{12}| |r_{32}|} - \cos(\theta_0) \right)^2} \right) \left( e^{\lambda_3 (|r_{12}| - |r_{32}|)} \right)^m \right)^n \right)^{-\frac{1}{2n}}$$

`espressopp.interaction.VerletListTersoffTripleTerm` (*system*, *vl3*)

### Parameters

- **system** –
- **vl3** –

`espressopp.interaction.VerletListTersoffTripleTerm.getPotential` (*type1*, *type2*, *type3*)

### Parameters

- **type1** –
- **type2** –
- **type3** –

### Return type

`espressopp.interaction.VerletListTersoffTripleTerm.getVerletListTriple` ()

**Return type** A Python list of lists.

`espressopp.interaction.VerletListTersoffTripleTerm.setPotential` (*type1*, *type2*, *type3*, *potential*)

### Parameters

- **type1** –
- **type2** –
- **type3** –
- **potential** –

`espressopp.interaction.FixedTripleListTersoffTripleTerm(system, ftl, potential)`

#### Parameters

- **system** –
- **ftl** –
- **potential** –

`espressopp.interaction.FixedTripleListTersoffTripleTerm.getFixedTripleList()`

**Return type** A Python list of lists.

`espressopp.interaction.FixedTripleListTersoffTripleTerm.setPotential(type1, type2, type3, potential)`

#### Parameters

- **type1** –
- **type2** –
- **type3** –
- **potential** –

## espressopp.interaction.VSpherePair

This class provides methods to compute forces and energies of the VSpherePair potential.

$$V(r_{ij}, \sigma_{ij}) = \varepsilon \left( \frac{2\pi}{3} \sigma_{ij} \right)^{-\frac{3}{2}} e^{-\frac{3}{2} \frac{r_{ij}^2}{\sigma_{ij}}}, r_{ij} = |\vec{r}_i - \vec{r}_j|, \sigma_{ij} = \sigma_i^2 + \sigma_j^2$$

Reference: Fluctuating soft-sphere approach to coars-graining of polymer melts, Soft matter, 2010, 6, 2282

`espressopp.interaction.VSpherePair(epsilon, cutoff, shift)`

#### Parameters

- **epsilon** (*real*) – (default: 1.0)
- **cutoff** – (default: infinity)
- **shift** – (default: “auto”)

`espressopp.interaction.VerletListVSpherePair(vl)`

#### Parameters vl –

`espressopp.interaction.VerletListVSpherePair.getPotential(type1, type2)`

#### Parameters

- **type1** –
- **type2** –

**Return type**

```
espressopp.interaction.VerletListVSPHPair.getVerletList()
```

**Return type** A Python list of lists.

```
espressopp.interaction.VerletListVSPHPair.setPotential(type1, type2, potential)
```

**Parameters**

- **type1** –
- **type2** –
- **potential** –

**class** `espressopp.interaction.VSPHPair.VSPHPair`  
The Lennard-Jones potential.

## espressopp.interaction.VSPHSelf

This class provides methods to compute forces and energies of the VSPHSelf potential.

$$U = e_1 \left( \frac{4}{3} \pi \sigma^2 \right)^{\frac{3}{2}} + \frac{a_1 N_b^3}{\sigma^6} + \frac{a_2}{N_b} \sigma^2$$

Reference: Fluctuating soft-sphere approach to coars-graining of polymer melts, Soft matter, 2010, 6, 2282

```
espressopp.interaction.VSPHSelf(e1, a1, a2, Nb, cutoff, shift)
```

**Parameters**

- **e1** (*real*) – (default: 0.0)
- **a1** (*real*) – (default: 1.0)
- **a2** (*real*) – (default: 0.0)
- **Nb** (*int*) – (default: 1)
- **cutoff** – (default: infinity)
- **shift** (*real*) – (default: 0.0)

```
espressopp.interaction.SelfVSPH(self, system, potential)
```

**Parameters**

- **system** –
- **potential** –

```
espressopp.interaction.SelfVSPH.getPotential()
```

**Return type**

```
espressopp.interaction.SelfVSPH.setPotential(potential)
```

**Parameters** **potential** –

**class** `espressopp.interaction.VSPHSelf.VSPHSelf`  
The VSPHSelf potential.

## espressopp.interaction.Zero

This class provides methods for a zero potential no interactions between particles, mainly used for debugging and testing

```
espressopp.interaction.Zero()
```

```
espressopp.interaction.VerletListZero(vl)
```

**Parameters `v1` –**

`espressopp.interaction.VerletListZero.getPotential` (*type1, type2*)

**Parameters**

- `type1` –
- `type2` –

**Return type**

`espressopp.interaction.VerletListZero.setFixedTupleList` (*ftpl*)

**Parameters `ftpl` –**

`espressopp.interaction.VerletListZero.setPotential` (*type1, type2, potential*)

**Parameters**

- `type1` –
- `type2` –
- `potential` –

`espressopp.interaction.VerletListAdressZero` (*vl*)

**Parameters `v1` –**

`espressopp.interaction.VerletListAdressZero.setFixedTupleList` (*ftpl*)

**Parameters `ftpl` –**

`espressopp.interaction.VerletListAdressZero.setPotentialAT` (*type1, type2, potential*)

**Parameters**

- `type1` –
- `type2` –
- `potential` –

`espressopp.interaction.VerletListAdressZero.setPotentialCG` (*type1, type2, potential*)

**Parameters**

- `type1` –
- `type2` –
- `potential` –

`espressopp.interaction.VerletListHadressZero` (*vl, fixedtupleList*)

**Parameters**

- `v1` –
- `fixedtupleList` –

`espressopp.interaction.VerletListHadressZero.setFixedTupleList` (*ftpl*)

**Parameters `ftpl` –**

`espressopp.interaction.VerletListHadressZero.setPotentialAT` (*type1, type2, potential*)

**Parameters**

- `type1` –
- `type2` –

- **potential** –

`espressopp.interaction.VerletListHadressZero.setPotentialCG` (*type1*, *type2*, *potential*)

#### Parameters

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.CellListZero` (*stor*)

#### Parameters stor –

`espressopp.interaction.CellListZero.setPotential` (*type1*, *type2*, *potential*)

#### Parameters

- **type1** –
- **type2** –
- **potential** –

`espressopp.interaction.FixedPairListZero` (*system*, *vl*, *potential*)

#### Parameters

- **system** –
- **vl** –
- **potential** –

`espressopp.interaction.FixedPairListZero.setPotential` (*potential*)

#### Parameters potential –

**class** `espressopp.interaction.Zero.Zero`

The Zero potential.

## io

### espressopp.io.DumpGRO

- *dump()* write configuration to trajectory GRO file. By default filename is “out.gro”, coordinates are folded.

#### Properties

- *filename* Name of trajectory file. By default trajectory file name is “out.gro”
- *unfolded* False if coordinates are folded, True if unfolded. By default - False
- *append* True if new trajectory data is appended to existing trajectory file. By default - True
- *length\_factor* If length dimension in current system is nm, and unit is 0.23 nm, for example, then *length\_factor* should be 0.23
- *length\_unit* It is length unit. Can be LJ, nm or A. By default - LJ

usage:

writing down trajectory

```
>>> dump_conf_gro = espressopp.io.DumpGRO(system, integrator, filename='trajectory.gro')
>>> for i in range(200):
>>>     integrator.run(10)
>>>     dump_conf_gro.dump()
```

writing down trajectory using ExtAnalyze extension

```
>>> dump_conf_gro = espressopp.io.DumpGRO(system, integrator, filename='trajectory.gro')
>>> ext_analyze = espressopp.integrator.ExtAnalyze(dump_conf_gro, 10)
>>> integrator.addExtension(ext_analyze)
>>> integrator.run(2000)
```

Both examples will give the same result: 200 configurations in trajectory .gro file.

setting up length scale

For example, the Lennard-Jones model for liquid argon with  $\sigma = 0.34[nm]$

```
>>> dump_conf_gro = espressopp.io.DumpGRO(system, integrator, filename='trj.gro', unfolded=False,
```

will produce trj.gro with in nanometers

```
espressopp.io.DumpGRO(system, integrator, filename, unfolded, length_factor, length_unit, ap-
pend)
```

#### Parameters

- **system** –
- **integrator** –
- **filename** – (default: 'out.gro')
- **unfolded** – (default: False)
- **length\_factor** (*real*) – (default: 1.0)
- **length\_unit** – (default: 'LJ')
- **append** – (default: True)

espressopp.io.DumpGRO.dump()

#### Return type

## espressopp.io.DumpGROAdress

dumps coordinates of atomistic particles instead of coarse-grained particles in Adress simulation

- *dump()* write configuration to trajectory GRO file. By default filename is “out.gro”, coordinates are folded.

#### Properties

- *filename* Name of trajectory file. By default trajectory file name is “out.gro”
- *unfolded* False if coordinates are folded, True if unfolded. By default - False
- *append* True if new trajectory data is appended to existing trajectory file. By default - True
- *length\_factor* If length dimension in current system is nm, and unit is 0.23 nm, for example, then length\_factor should be 0.23
- *length\_unit* It is length unit. Can be LJ, nm or A. By default - LJ
- *ftpl* fixedtuplelist for the adres system

usage:

```
>>> ftpl = espressopp.FixedTupleListAdress(system.storage)
>>> ftpl.addTuples(tuples)
>>> system.storage.setFixedTuplesAdress(ftpl)
>>> system.storage.decompose()
```

writing down trajectory



```
>>> dump_conf_gro = espressopp.io.DumpGROAddress(system, ftpl, integrator, filename='trajectory.gro')
>>> for i in range(200):
>>>     integrator.run(10)
>>>     dump_conf_gro.dump()
```

writing down trajectory using ExtAnalyze extension

```
>>> dump_conf_gro = espressopp.io.DumpGROAddress(system, ftpl, integrator, filename='trajectory.gro')
>>> ext_analyze = espressopp.integrator.ExtAnalyze(dump_conf_gro, 10)
>>> integrator.addExtension(ext_analyze)
>>> integrator.run(2000)
```

Both examples will give the same result: 200 configurations in trajectory .gro file.

setting up length scale

For example, the Lennard-Jones model for liquid argon with  $\sigma = 0.34[nm]$

```
>>> dump_conf_gro = espressopp.io.DumpGROAddress(system, ftpl, integrator, filename='trj.gro', unfolded,
```

will produce trj.gro with in nanometers

```
espressopp.io.DumpGROAddress(system, fixedtuplelist, integrator, filename, unfolded,
                              length_factor, length_unit, append)
```

#### Parameters

- **system** –
- **fixedtuplelist** –
- **integrator** –
- **filename** – (default: 'out.gro')
- **unfolded** – (default: False)
- **length\_factor** (*real*) – (default: 1.0)
- **length\_unit** – (default: 'LJ')
- **append** – (default: True)

```
espressopp.io.DumpGROAddress.dump()
```

#### Return type

## espressopp.io.DumpXYZ

- *dump()*

write configuration to trajectory XYZ file. By default filename is out.xyz, coordinates are folded. DumpXYZ works also for Multiple communicators.

#### Properties

- *filename* Name of trajectory file. By default trajectory file name is out.xyz
- *unfolded* False if coordinates are folded, True if unfolded. By default - False
- *append* True if new trajectory data is appended to existing trajectory file. By default - True
- *length\_factor* If length dimension in current system is nm, and unit is 0.23 nm, for example, then length\_factor should be 0.23 Default: 1.0
- *length\_unit* It is length unit. Can be LJ, nm or A. By default - LJ
- *store\_pids* True if you want to store pids as fastwritexyz does. False otherwise (standard XYZ) Default: False

- **store\_velocities** True if you want to store velocities. False otherwise (XYZ doesn't require it) Default: False

usage:

writing down trajectory

```
>>> dump_conf_xyz = espressopp.io.DumpXYZ(system, integrator, filename='trajectory.xyz')
>>> for i in range(200):
>>>     integrator.run(10)
>>>     dump_conf_xyz.dump()
```

writing down trajectory using ExtAnalyze extension

```
>>> dump_conf_xyz = espressopp.io.DumpXYZ(system, integrator, filename='trajectory.xyz')
>>> ext_analyze = espressopp.integrator.ExtAnalyze(dump_conf_xyz, 10)
>>> integrator.addExtension(ext_analyze)
>>> integrator.run(2000)
```

Both examples will give the same result: 200 configurations in trajectory .xyz file.

setting up length scale

For example, the Lennard-Jones model for liquid argon with  $\sigma = 0.34[nm]$

```
>>> dump_conf_xyz = espressopp.io.DumpXYZ(system, integrator, filename='trj.xyz', \
>>>                                     unfolded=False, length_factor=0.34, \
>>>                                     length_unit='nm', store_pids=True, \
>>>                                     store_velocities = True, append=True)
```

will produce trj.xyz with in nanometers

```
espressopp.io.DumpXYZ(system, integrator, filename=out.xyz, unfolded=False, length_factor=1.0,
                      length_unit='LJ', store_pids=False, store_velocities=False, ap-
                      pend=True)
```

#### Parameters

- **system** –
- **integrator** –
- **filename** –
- **unfolded** (*bool*) –
- **length\_factor** (*real*) –
- **length\_unit** –
- **store\_pids** (*bool*) –
- **store\_velocities** (*bool*) –
- **append** (*bool*) –

`espressopp.io.DumpXYZ.dump()`

#### Return type

## espressopp

### espressopp.Exceptions

`espressopp.Error(msg)`

Parameters **msg** –

`espressopp.ParticleDoesNotExistHere (msg)`

**Parameters** `msg` –

`espressopp.UnknownParticleProperty (msg)`

**Parameters** `msg` –

`espressopp.MissingFixedPairList (msg)`

**Parameters** `msg` –

## **espressopp.FixedLocalTupleList**

This class can contain many tuple which store a arbitrary positive number, which should be more than 2, of local (real + ghost) particle id.

For using this class, there is 1 conditions:

Particles in one tuple must be in a same or neighbor cell list.

`espressopp.FixedLocalTupleList (storage)`

**Parameters** `storage` –

`espressopp.FixedLocalTupleList.addTuple (tuple)`

**Parameters** `tuple` (*python::list*) –

`espressopp.FixedLocalTupleList.getTuples ()`

**Return type**

`espressopp.FixedLocalTupleList.size ()`

**Return type**

## **espressopp.FixedPairDistList**

`espressopp.FixedPairDistList (storage)`

**Parameters** `storage` –

`espressopp.FixedPairDistList.add (pid1, pid2)`

**Parameters**

- `pid1` –
- `pid2` –

**Return type**

`espressopp.FixedPairDistList.addPairs (bondlist)`

**Parameters** `bondlist` –

**Return type**

`espressopp.FixedPairDistList.getDist (pid1, pid2)`

**Parameters**

- `pid1` –
- `pid2` –

**Return type**

`espressopp.FixedPairDistList.getPairs ()`

**Return type**

```
espressopp.FixedPairDistList.getPairsDist()
```

**Return type**

```
espressopp.FixedPairDistList.size()
```

**Return type**

## espressopp.FixedPairList

```
espressopp.FixedPairList(storage)
```

**Parameters** *storage* –

```
espressopp.FixedPairList.add(pid1, pid2)
```

**Parameters**

- *pid1* –
- *pid2* –

**Return type**

```
espressopp.FixedPairList.addBonds(bondlist)
```

**Parameters** *bondlist* –

**Return type**

```
espressopp.FixedPairList.getBonds()
```

**Return type**

```
espressopp.FixedPairList.remove()
```

‘remove the FixedPairList and disconnect’

```
espressopp.FixedPairList.getLongtimeMaxBond()
```

**Return type**

```
espressopp.FixedPairList.resetLongtimeMaxBond()
```

**Return type**

```
espressopp.FixedPairList.size()
```

**Return type**

## espressopp.FixedPairListAdress

The FixedPairListAdress is the Fixed Pair List to be used for AdResS or H-AdResS simulations. When creating the FixedPairListAdress one has to provide the storage and the tuples. Afterwards the bonds can be added. In the example “bonds” is a python list of the form ( (pid1, pid2), (pid3, pid4), ...) where each inner pair defines a bond between the particles with the given particle ids.

Example - creating the FixedPairListAdress and adding bonds:

```
>>> ftpl = espressopp.FixedTupleList(system.storage)
>>> fpl = espressopp.FixedPairListAdress(system.storage, ftpl)
>>> fpl.addBonds(bonds)
```

```
espressopp.FixedPairListAdress(storage, fixedtupleList)
```

**Parameters**

- *storage* –
- *fixedtupleList* –

`espressopp.FixedPairListAdress.add(pid1, pid2)`

**Parameters**

- `pid1` –
- `pid2` –

**Return type**

`espressopp.FixedPairListAdress.addBonds(bondlist)`

**Parameters** `bondlist` –

**Return type**

`espressopp.FixedPairListAdress.remove()`  
**remove the FixedPairListAdress and disconnect**

`espressopp.FixedPairListAdress.getBonds()`

**Return type**

## espressopp.FixedQuadrupleAngleList

`espressopp.FixedQuadrupleAngleList(storage)`

**Parameters** `storage` –

`espressopp.FixedQuadrupleAngleList.add(pid1, pid2, pid3, pid4)`

**Parameters**

- `pid1` –
- `pid2` –
- `pid3` –
- `pid4` –

**Return type**

`espressopp.FixedQuadrupleAngleList.addQuadruples(quadruplelist)`

**Parameters** `quadruplelist` –

**Return type**

`espressopp.FixedQuadrupleAngleList.getAngle(pid1, pid2, pid3, pid4)`

**Parameters**

- `pid1` –
- `pid2` –
- `pid3` –
- `pid4` –

**Return type**

`espressopp.FixedQuadrupleAngleList.getQuadruples()`

**Return type**

`espressopp.FixedQuadrupleAngleList.getQuadruplesAngles()`

**Return type**

`espressopp.FixedQuadrupleAngleList.size()`

**Return type**

## espressopp.FixedQuadrupleList

`espressopp.FixedQuadrupleList` (*storage*)

Parameters **storage** –

`espressopp.FixedQuadrupleList.add` (*pid1, pid2, pid3, pid4*)

Parameters

- **pid1** –
- **pid2** –
- **pid3** –
- **pid4** –

Return type

`espressopp.FixedQuadrupleList.addQuadruples` (*quadruplelist*)

Parameters **quadruplelist** –

Return type

`espressopp.FixedQuadrupleList.remove` ()  
**remove the FixedPairList and disconnect**

`espressopp.FixedQuadrupleList.getQuadruples` ()

Return type

`espressopp.FixedQuadrupleList.size` ()

Return type

## espressopp.FixedQuadrupleListAdress

`espressopp.FixedQuadrupleListAdress` (*storage, fixedtupleList*)

Parameters

- **storage** –
- **fixedtupleList** –

`espressopp.FixedQuadrupleListAdress.add` (*pid1, pid2, pid3, pid4*)

Parameters

- **pid1** –
- **pid2** –
- **pid3** –
- **pid4** –

Return type

`espressopp.FixedQuadrupleListAdress.addQuadruples` (*quadruplelist*)

Parameters **quadruplelist** –

Return type

`espressopp.FixedQuadrupleListAdress.getQuadruples` ()

Return type

`espressopp.FixedQuadrupleListAdress.size` ()

Return type

## espressopp.FixedSingleList

`espressopp.FixedSingleList (storage)`

**Parameters** `storage` –

`espressopp.FixedSingleList.add (pid1)`

**Parameters** `pid1` –

**Return type**

`espressopp.FixedSingleList.addSingles (singlelist)`

**Parameters** `singlelist` –

**Return type**

`espressopp.FixedSingleList.getSingles ()`

**Return type**

`espressopp.FixedSingleList.size ()`

**Return type**

## espressopp.FixedTripleAngleList

`espressopp.FixedTripleAngleList (storage)`

**Parameters** `storage` –

`espressopp.FixedTripleAngleList.add (pid1, pid2, pid3)`

**Parameters**

- `pid1` –
- `pid2` –
- `pid3` –

**Return type**

`espressopp.FixedTripleAngleList.addTriples (triplelist)`

**Parameters** `triplelist` –

**Return type**

`espressopp.FixedTripleAngleList.getAngle (pid1, pid2, pid3)`

**Parameters**

- `pid1` –
- `pid2` –
- `pid3` –

**Return type**

`espressopp.FixedTripleAngleList.getTriples ()`

**Return type**

`espressopp.FixedTripleAngleList.getTriplesAngles ()`

**Return type**

`espressopp.FixedTripleAngleList.size ()`

**Return type**

## espressopp.FixedTripleList

`espressopp.FixedTripleList (storage)`

Parameters **storage** –

`espressopp.FixedTripleList.add (pid1, pid2, pid3)`

Parameters

- **pid1** –
- **pid2** –
- **pid3** –

Return type

`espressopp.FixedTripleList.addTriples (triplelist)`

Parameters **triplelist** –

Return type

`espressopp.FixedTripleList.getTriples ()`

Return type

`espressopp.FixedTripleList.size ()`

Return type

`espressopp.FixedTripleList.remove ()`

**remove the FixedPairList and disconnect**

## espressopp.FixedTripleListAdress

`espressopp.FixedTripleListAdress (storage, fixedtupleList)`

Parameters

- **storage** –
- **fixedtupleList** –

`espressopp.FixedTripleListAdress.add (pid1, pid2)`

Parameters

- **pid1** –
- **pid2** –

Return type

`espressopp.FixedTripleListAdress.remove ()`

**remove the FixedTripleListAdress and disconnect**

`espressopp.FixedTripleListAdress.addTriples (triplelist)`

Parameters **triplelist** –

Return type

## espressopp.FixedTupleList

`espressopp.FixedTupleList (storage)`

Parameters **storage** –

`espressopp.FixedTupleList.size ()`



**Return type****espressopp.FixedTupleListAdress**

The FixedTupleListAdress is important for AdResS and H-AdResS simulations. It is the connection between the atomistic and coarse-grained particles. It defines which atomistic particles belong to which coarse-grained particle. In the following example “tuples” is a python list of the form ( (pid\_CG1, pidAT11, pidAT12, pidAT13, ...), (pid\_CG2, pidAT21, pidAT22, pidAT23, ...), ...). Each inner list (pid\_CG1, pidAT11, pidAT12, pidAT13, ...) defines a tuple. The first number is the particle id of the coarse-grained particle while the following numbers are the particle ids of the corresponding atomistic particles.

Example - creating the FixedTupleListAdress:

```
>>> ftpl = espressopp.FixedTupleListAdress(system.storage)
>>> ftpl.addTuples(tuples)
>>> system.storage.setFixedTuples(ftpl)
```

espressopp.**FixedTupleListAdress** (*storage*)

**Parameters** *storage* –

espressopp.FixedTupleListAdress.**addTuples** (*tuplelist*)

**Parameters** *tuplelist* –

**Return type**

**espressopp.Int3D**

espressopp.\_\_Int3D (*\*args*)

**Parameters** *\*args* –

espressopp.\_\_Int3D.**x** (*v*, [*0*])

**Parameters**

- *v* –
- [*0* –

**Return type**

espressopp.\_\_Int3D.**y** (*v*, [*1*])

**Parameters**

- *v* –
- [*1* –

**Return type**

espressopp.\_\_Int3D.**z** (*v*, [*2*])

**Parameters**

- *v* –
- [*2* –

**Return type**

espressopp.**toInt3DFromVector** (*\*args*)

**Parameters** *\*args* –

espressopp.**toInt3D** (*\*args*)

**Parameters** *\*args* –

`espressopp.Int3D.toInt3D (*args)`

Try to convert the arguments to a `Int3D`, returns the argument, if it is already a `Int3D`.

`espressopp.Int3D.toInt3DFromVector (*args)`

Try to convert the arguments to a `Int3D`.

This function will only convert to a `Int3D` if `x`, `y` and `z` are specified.

## espressopp.MultiSystem

`espressopp.MultiSystem()`

`espressopp.MultiSystem.beginSystemDefinition()`

**Return type**

`espressopp.MultiSystem.runAnalysisNPart()`

**Return type**

`espressopp.MultiSystem.runAnalysisPotential()`

**Return type**

`espressopp.MultiSystem.runAnalysisTemperature()`

**Return type**

`espressopp.MultiSystem.runIntegrator (niter)`

**Parameters** `niter` –

**Return type**

`espressopp.MultiSystem.setAnalysisNPart (npart)`

**Parameters** `npart` –

`espressopp.MultiSystem.setAnalysisPotential (potential)`

**Parameters** `potential` –

`espressopp.MultiSystem.setAnalysisTemperature (temperature)`

**Parameters** `temperature` –

`espressopp.MultiSystem.setIntegrator (integrator)`

**Parameters** `integrator` –

**class** `espressopp.MultiSystem.MultiSystem`

MultiSystemIntegrator to simulate and analyze several systems in parallel.

**class** `espressopp.MultiSystem.MultiSystemLocal`

Local MultiSystem to simulate and analyze several systems in parallel.

## espressopp.ParallelTempering

`espressopp.ParallelTempering (NumberOfSystems, RNG)`

**Parameters**

- **NumberOfSystems** (`int`) – (default: 4)
- **RNG** – (default: None)

`espressopp.ParallelTempering.endDefiningSystem (n)`

**Parameters** `n` –

**Return type**

`espressopp.ParallelTempering.exchange()`

**Return type**

`espressopp.ParallelTempering.getNumberOfCPUsPerSystem()`

**Return type**

`espressopp.ParallelTempering.getNumberOfSystems()`

**Return type**

`espressopp.ParallelTempering.run(nsteps)`

**Parameters** `nsteps` –

**Return type**

`espressopp.ParallelTempering.setAnalysisE(analysisE)`

**Parameters** `analysisE` –

`espressopp.ParallelTempering.setAnalysisNPart(analysisNPart)`

**Parameters** `analysisNPart` –

`espressopp.ParallelTempering.setAnalysisT(analysisT)`

**Parameters** `analysisT` –

`espressopp.ParallelTempering.setIntegrator(integrator, thermostat)`

**Parameters**

- `integrator` –
- `thermostat` –

`espressopp.ParallelTempering.startDefiningSystem(n)`

**Parameters** `n` –

**Return type**

## espressopp.Particle

`espressopp.Particle(pid, storage)`

**Parameters**

- `pid` –
- `storage` –

**class** `espressopp.Particle.ParticleLocal(pid, storage)`

The local particle.

Throws an exception: \* when the particle does not exists locally

TODO: Should throw an exception: \* when a ghost particle is to be written \* when data is to be read from a ghost that is not available

## espressopp.ParticleAccess

Abstract base class for analysis/measurement/io

`espressopp.ParticleAccess.perform_action()`

**Return type**

## espressopp.ParticleGroup

`espressopp.ParticleGroup(storage)`

**Parameters** `storage` –

`espressopp.ParticleGroup.add(pid)`

**Parameters** `pid` –

**Return type**

`espressopp.ParticleGroup.has(pid)`

**Parameters** `pid` –

**Return type**

`espressopp.ParticleGroup.show()`

**Return type**

`espressopp.ParticleGroup.size()`

**Return type**

## espressopp.pmi

Parallel Method Invocation (PMI) allows users to write serial Python scripts that use functions and classes that are executed in parallel.

PMI is intended to be used in data-parallel environments, where several threads run in parallel and can communicate via MPI.

In PMI mode, a single thread of control (a python script that runs on the *controller*, i.e. the MPI root task) can invoke arbitrary functions on all other threads (the *workers*) in parallel via *call()*, *invoke()* and *reduce()*. When the function on the workers return, the control is returned to the controller.

This model is equivalent to the “Fork-Join execution model” used e.g. in OpenMP.

PMI also allows to create parallel instances of object classes via *create()*, i.e. instances that have a corresponding object instance on all workers. *call()*, *invoke()* and *reduce()* can be used to call arbitrary methods of these instances.

to execute arbitrary code on all workers, *exec\_()* can be used, and to import python modules to all workers, use *‘import\_()’*.

### Main program

On the workers, the main program of a PMI script usually consists of a single call to the function *startWorkerLoop()*. On the workers, this will start an infinite loop on the workers that waits to receive the next PMI call, while it will immediately return on the controller. On the workers, the loop ends only, when one of the commands *finalizeWorkers()* or *stopWorkerLoop()* is issued on the controller. A typical PMI main program looks like this:

```
>>> # compute 2*factorial(42) in parallel
>>> import pmi
>>>
>>> # start the worker loop
>>> # on the controller, this function returns immediately
>>> pmi.startWorkerLoop()
>>>
>>> # Do the parallel computation
>>> pmi.import_('math')
>>> pmi.reduce('lambda a,b: a+b', 'math.factorial', 42)
>>>
>>> # exit all workers
>>> pmi.finalizeWorkers()
```

Instead of using `finalizeWorkers()` at the end of the script, you can call `registerAtExit()` anywhere else, which will cause `finalizeWorkers()` to be called when the python interpreter exits.

Alternatively, it is possible to use PMI in an SPMD-like fashion, where each call to a PMI command on the controller must be accompanied by a corresponding call on the worker. This can be either a simple call to `receive()` that accepts any PMI command, or a call to the identical PMI command. In that case, the arguments of the call to the PMI command on the workers are ignored. In this way, it is possible to write SPMD scripts that profit from the PMI communication patterns.

```
>>> # compute 2*factorial(42) in parallel
>>> import pmi
>>>
>>> pmi.exec_('import math')
>>> pmi.reduce('lambda a,b: a+b', 'math.factorial', 42)
```

To start the worker loop, the command `startWorkerLoop()` can be issued on the workers. To stop the worker loop, `stopWorkerLoop()` can be issued on the controller, which will end the worker loop without exiting the workers.

### Controller commands

These commands can be called in the controller script. When any of these commands is issued on a worker during the worker loop, a `UserError` is raised.

- `call()`, `invoke()`, `reduce()` to call functions and methods in parallel
- `create()` to create parallel object instances
- `exec_()` and `import_()` to execute arbitrary python code in parallel and to import classes and functions into the global namespace of pmi.
- `sync()` to make sure that all deleted PMI objects have been deleted.
- `finalizeWorkers()` to stop and exit all workers
- `registerAtExit()` to make sure that `finalizeWorkers()` is called when python exits on the controller
- `stopWorkerLoop()` to interrupt the worker loop on all workers and to return control to the single workers

### Worker commands

These commands can be called on a worker.

- `startWorkerLoop()` to start the worker loop
- `receive()` to receive a single PMI command
- `call()`, `invoke()`, `reduce()`, `create()` and `exec_()` to receive a single corresponding PMI command. Note that these commands will ignore any arguments when called on a worker.

### PMI Proxy metaclass

The *Proxy* metaclass can be used to easily generate front-end classes to distributed PMI classes. . . .

### Useful constants and variables

The pmi module defines the following useful constants and variables:

- `isController` is True when used on the controller, False otherwise
- `isWorker` = not `isController`
- `ID` is the rank of the MPI task
- `CONTROLLER` is the rank of the Controller (normally the MPI root)
- `workerStr` is a string describing the thread ('Worker #' or 'Controller')
- `inWorkerLoop` is True, if PMI currently executes the worker loop on the workers.

```
espressopp.pmi.exec_(*args)
```

Controller command that executes arbitrary python code on all (active) workers.

`exec_()` allows to execute arbitrary Python code on all workers. It can be used to define classes and functions on all workers. Modules should not be imported via `exec_()`, instead `import_()` should be used.

Each element of `args` should be string that is executed on all workers.

Example:

```
>>> pmi.exec_('import hello')
>>> hw = pmi.create('hello.HelloWorld')
```

`espressopp.pmi.import_(*args)`

Controller command that imports python modules on all (active) workers.

Each element of `args` should be a module name that is imported to all workers.

Example:

```
>>> pmi.import_('hello')
>>> hw = pmi.create('hello.HelloWorld')
```

`espressopp.pmi.create(cls=None, *args, **kwds)`

Controller command that creates an object on all workers.

`cls` describes the (new-style) class that should be instantiated. `args` are the arguments to the constructor of the class. Only classes that are known to PMI can be used, that is, classes that have been imported to `pmi` via `exec_()` or `import_()`.

Example:

```
>>> pmi.exec_('import hello')
>>> hw = pmi.create('hello.HelloWorld')
>>> print(hw)
MPI process #0: Hello World!
MPI process #1: Hello World!
...
```

Alternative: Note that in this case the class has to be imported to the calling module *and* via PMI.

```
>>> import hello
>>> pmi.exec_('import hello')
>>> hw = pmi.create(hello.HelloWorld)
>>> print(hw)
MPI process #0: Hello World!
MPI process #1: Hello World!
...
```

`espressopp.pmi.call(*args, **kwds)`

Call a function on all workers, returning only the return value on the controller.

`function` denotes the function that is to be called, `args` and `kwds` are the arguments to the function. If `kwds` contains keys that start with the prefix `'__pmictr_'`, they are stripped of the prefix and are passed only to the controller. If the function should return any results, it will be locally returned. Only functions that are known to PMI can be used, that is functions that have been imported to `pmi` via `exec_()` or `import_()`.

Example:

```
>>> pmi.exec_('import hello')
>>> hw = pmi.create('hello.HelloWorld')
>>> pmi.call(hw.hello)
>>> # equivalent:
>>> pmi.call('hello.HelloWorld', hw)
```

Note, that you can use only functions that are known to PMI when `call()` is called, i.e. functions in modules that have been imported via `exec_()`.

`espressopp.pmi.invoke(*args, **kwds)`

Invoke a function on all workers, gathering the return values into a list.

function denotes the function that is to be called, args and kwds are the arguments to the function. If kwds contains keys that start with the prefix `'__pmictr_'`, they are stripped of the prefix and are passed only to the controller.

On the controller, `invoke()` returns the results of the different workers as a list. On the workers, `invoke` returns `None`. Only functions that are known to PMI can be used, that is functions that have been imported to pmi via `exec_()` or `import_()`.

Example:

```
>>> pmi.exec_('import hello')
>>> hw = pmi.create('hello.HelloWorld')
>>> messages = pmi.invoke(hw.hello())
>>> # alternative:
>>> messages = pmi.invoke('hello.HelloWorld.hello', hw)
```

`espressopp.pmi.reduce(*args, **kwds)`

Invoke a function on all workers, reducing the return values to a single value.

`reduceOp` is the (associative) operator that is used to process the return values, function denotes the function that is to be called, args and kwds are the arguments to the function. If kwds contains keys that start with the prefix `'__pmictr_'`, they are stripped of the prefix and are passed only to the controller.

`reduce()` reduces the results of the different workers into a single value via the operation `reduceOp`. `reduceOp` is assumed to be associative. Both `reduceOp` and function have to be known to PMI, that is they must have been imported to pmi via `exec_()` or `import_()`.

Example:

```
>>> pmi.exec_('import hello')
>>> pmi.exec_('joinstr=lambda a,b: "\n".join(a,b)')
>>> hw = pmi.create('hello.HelloWorld')
>>> print(pmi.reduce('joinstr', hw.hello()))
>>> # equivalent:
>>> print(
...     pmi.reduce('lambda a,b: "\n".join(a,b)',
...                'hello.HelloWorld.hello', hw)
...     )
```

`espressopp.pmi.sync()`

Controller command that deletes the PMI objects on the workers that have already been deleted on the controller.

`espressopp.pmi.receive(expected=None)`

Worker command that receives and handles the next PMI command.

This function waits to receive and handle a single PMI command. If `expected` is not `None` and the received command does not equal `expected`, raise a *UserError*.

`espressopp.pmi.startWorkerLoop()`

Worker command that starts the main worker loop.

This function starts a loop that expects to receive PMI commands until `stopWorkerLoop()` or `finalizeWorkers()` is called on the controller.

`espressopp.pmi.finalizeWorkers()`

Controller command that stops and exits all workers.

`espressopp.pmi.stopWorkerLoop(doExit=False)`

Controller command that stops all workers.

If `doExit` is set, the workers exit afterwards.

`espressopp.pmi.registerAtExit()`

Controller command that registers the function `finalizeWorkers()` via `atexit`.

**class** `espressopp.pmi.Proxy` (*name, bases, dict*)  
 A metaclass to be used to create frontend serial objects.

**exception** `espressopp.pmi.UserError` (*msg*)  
 Raised when PMI has encountered a user error.

## espressopp.Quaternion

This class provides quaternions with the associate methods. Quaternions can be used as an efficient representation for the orientation and rotation of 3D vector objects in 3D euclidean space. A Quaternion as such has a real part and an imaginary part. For implementation purposes, the representation through one real scalar and one real 3D vector is used here. The vector part is defined using the `Real3D` class of `espressopp`.

The format of a quaternion is “(real\_part, unreal\_part)” with the types “real” and “Real3D”, respectively.

While there are other possible applications for quaternions (rotation) in the simulation code, they will be used at the C++-level in order to perform the integration of the Euler equations of motion regarding the particles angular motion, i.e. the rigid body dynamics.

### Usage:

The following methods from C++-level are available at the python-level:

- **getReal()** return the scalar part of the quaternion
- **setReal(real)** sets the scalar part of the quaternion
- **getImag()** returns the vector part of the quaternion
- **getImagItem(i)** returns element *i* of vector part of the quaternion
- **setImag(Real3D)** sets the vector part of the quaternion
- **setImagItem(i, real)** sets element *i* of vector part of the quaternion
- **sqr()** the inner product of the quaternion
- **abs()** the absolute value of the quaternion
- **normalize()** normalizes the quaternion to unit length
- **transpose()** transposes the quaternion (changes sign of unreal\_part)

The multiplication operator is overloaded in order to perform quaternion multiplication, see examples below. Furthermore, it is possible to multiply a quaternion with a scalar, in order to rescale it.

### Examples:

#### Initialize:

```
>>> espressopp.Quaternion()
Quaternion(0.0, Real3D(0.0, 0.0, 0.0))
```

```
>>> espressopp.Quaternion(0.0, 1.0, 2.0, 3.0)
Quaternion(1.0, Real3D(1.0, 2.0, 3.0))
```

```
>>> vec = espressopp.Real3D(1.0, 2.0, 3.0)
>>> Quaternion(vec)
Quaternion(0.0, Real3D(1.0, 2.0, 3.0))
```

```
>>> espressopp.Quaternion(1.0)
Quaternion(1.0, Real3D(0.0, 0.0, 0.0))
```

#### Get:



```
>>> q = espressopp.Quaternion(0.0, 1.0, 2.0, 3.0)
>>> q.getReal()
0.0
>>> q.getImag()
Real3D(1.0, 2.0, 3.0)
>>> q.getImagItem(0)
1.0
```

#### Set:

```
>>> q = espressopp.Quaternion(0.0, 0.0, 0.0, 0.0)
>>> q.setReal(1.0)
>>> vec = espressopp.Real3D(1.0, 2.0, 3.0)
>>> q.setImag(vec)
>>> q
Quaternion(1.0, Real3D(1.0, 2.0, 3.0))
>>> q.setImagItem(0, 0.0)
Quaternion(1.0, Real3D(0.0, 2.0, 3.0))
```

#### Transpose and normalize:

```
>>> q = Quaternion(0.0, 1.0, 2.0, 3.0)
>>> q.transpose()
Quaternion(0.0, Real3D(-1.0, -2.0, -3.0))
>>> q = Quaternion(0.0, 1.0, 2.0, 3.0)
>>> q.normalize()
Quaternion(0.0, Real3D(0.2672612419124244, 0.5345224838248488, 0.8017837257372732))
```

#### Inner product and absolute value:

```
>>> q = Quaternion(0.0, 1.0, 2.0, 3.0)
>>> q.sqr()
14.0
>>> q.abs()
3.7416573867739413
```

#### Quaternion multiplication (compare, e.g., wikipedia):

```
>>> p = Quaternion(0.0, 1.0, 2.0, 3.0)
>>> q = Quaternion(0.0, 1.0, 2.0, 3.0)
Quaternion(-14.0, Real3D(0.0, 0.0, 0.0))
```

`espressopp.Quaternion.toQuaternion(*args)`

Try to convert the arguments to a Quaternion, return the argument if it is already a Quaternion.

`espressopp.Quaternion.toQuaternionFromVector(*args)`

Try to convert the arguments to a Quaternion.

This function will only convert to a Quaternion if `real_part`, `unreal_part[0]`, `unreal_part[1]` and `unreal_part[2]` are specified.

## espressopp.Real3D

`espressopp.__Real3D(*args)`

**Parameters** `*args` –

`espressopp.__Real3D.x(v, [0])`

**Parameters**

- `v` –
- `[0` –

**Return type**

`espressopp.__Real3D.y(v, [1])`

**Parameters**

- **v** –
- **[1** –

**Return type**

`espressopp.__Real3D.z(v, [2])`

**Parameters**

- **v** –
- **[2** –

**Return type**

`espressopp.toReal3DFromVector(*args)`

**Parameters** **\*args** –

`espressopp.toReal3D(*args)`

**Parameters** **\*args** –

`espressopp.Real3D.toReal3D(*args)`

Try to convert the arguments to a Real3D, returns the argument, if it is already a Real3D.

`espressopp.Real3D.toReal3DFromVector(*args)`

Try to convert the arguments to a Real3D.

This function will only convert to a Real3D if x, y and z are specified.

## espressopp.RealND

This is the object which represents N-dimensional vector. It is an extended Real3D, basically, it has the same functionality but in N-dimensions. First of all it is useful for classes in ‘espressopp.analysis’.

### Description

...

`espressopp.__RealND(*args)`

**Parameters** **\*args** –

`espressopp.toRealNDFromVector(*args)`

**Parameters** **\*args** –

`espressopp.toRealND(*args)`

**Parameters** **\*args** –

`espressopp.RealND.toRealND(*args)`

Try to convert the arguments to a RealND, returns the argument, if it is already a RealND.

`espressopp.RealND.toRealNDFromVector(*args)`

Try to convert the arguments to a RealND.

This function will only convert to a RealND if x, y and z are specified.

## espressopp.System

The main purpose of this class is to store pointers to some important other classes and thus make them available to C++. In a way the System class can be viewed as a container for system wide global variables. If you need to run more than one system at the same time you can combine several systems with the help of the Multisystem class.

In detail the System class holds pointers to:

- the *storage* (e.g. DomainDecomposition)
- the boundary conditions *bc* for the system (e.g. OrthorhombicBC)
- a random number generator *rng* which is for example used by a thermostat
- the *skin* which is needed for the Verlet lists and the cell grid
- a list of short range interactions that apply to the system these interactions are added with the *addInteraction()* method of the System

Example (not complete):

```
>>> LJSystem      = espressopp.System()
>>> LJSystem.bc    = espressopp.bc.OrthorhombicBC(rng, boxsize)
>>> LJSystem.rng
>>> LJSystem.skin = 0.4
>>> LJSystem.addInteraction(interLJ)
```

`espressopp.System()`

`espressopp.System.addInteraction(interaction, name)`

**Parameters**

- **interaction** –
- **name** (*string*) – The optional name of the interaction.

**Return type** bool

`espressopp.System.getInteraction(number)`

**Parameters** **number** –

**Return type**

`espressopp.System.getNumberOfInteractions()`

**Return type**

`espressopp.System.removeInteraction(number)`

**Parameters** **number** –

**Return type**

`espressopp.System.removeInteractionByName(self, name)`

**Parameters** **name** (*str*) – The name of the interaction to remove.

`espressopp.System.getAllInteractions()`

**Return type** The dictionary with name as a key and Interaction object.

`espressopp.System.scaleVolume(*args)`

**Parameters** **\*args** –

**Return type**

`espressopp.System.setTrace(switch)`

**Parameters** **switch** –

## espressopp.Tensor

`espressopp.Tensor.toTensor (*args)`

Try to convert the arguments to a Tensor, returns the argument, if it is already a Tensor.

`espressopp.Tensor.toTensorFromVector (*args)`

Try to convert the arguments to a Tensor.

This function will only convert to a Tensor if x, y and z are specified.

## espressopp.VerletList

`espressopp.VerletList (system, cutoff, exclusionlist)`

### Parameters

- **system** –
- **cutoff** –
- **exclusionlist** – (default: [])

`espressopp.VerletList.exclude (exclusionlist)`

**Parameters** **exclusionlist** –

**Return type**

`espressopp.VerletList.getAllPairs ()`

**Return type**

`espressopp.VerletList.localSize ()`

**Return type**

`espressopp.VerletList.totalSize ()`

**Return type**

## espressopp.VerletListAddress

The VerletListAddress is the Verlet List to be used for AdResS or H-AdResS simulations. When creating the VerletListAddress one has to provide the system and specify both cutoff for the CG interaction and adrcutoff for the atomistic interaction. Often, it is important to set the atomistic adrcutoff much bigger than the actual interaction's cutoff would be, since also the atomistic part of the VerletListAddress (adrPairs) is built based on the coarse-grained particle positions. For a much larger coarse-grained cutoff it is for example possible to also set the atomistic cutoff on the same value as the coarse-grained one.

Furthermore, the sizes of the explicit and hybrid region have to be provided (dEx and dHy in the example below) and the center of the atomistic region has to be set (adrCenter). Additionally, it can be chosen between a spherical and a slab-like geometry (sphereAdr).

The AdResS region can also be defined based on one or more particles. For a single particle, in this case a spherical region moves along with the particle. For many such region defining particles, the high-resolution/hybrid region corresponds to the overlap of the different spherical regions based on the individual particles (for details see Kreis et al., JCTC doi: 10.1021/acs.jctc.6b00440). Note that more region defining particles mean a higher computational overhead as these particles need to be communicated among all processors (also see explanations in AdResS.py). Also note that region defining particles should be normal/CG particles, not atomistic/AdResS ones.

**Basically the VerListAddress provides 4 lists:**

- **adrZone**: A list which holds all particles in the atomistic and hybrid region
- **cgZone**: A list which holds all particles in the coarse-grained region

- **adrPairs**: A list which holds all pairs which have at least one particle in the `adrZone`, i.e. in the atomistic or hybrid region
- **vlPairs**: A list which holds all pairs which have both particles in the `cgZone`, i.e. in the coarse-grained region

Example - creating the `VerletListAdress` for a slab-type address region fixed in space (only the `x` value of `adrCenter` is used):

```
>>> vl = espressopp.VerletListAdress(system, cutoff=rc, adrcut=rc, dEx=ex_size, dHy=hy_size,
```

or

```
>>> vl = espressopp.VerletListAdress(system, cutoff=rc, adrcut=rc, dEx=ex_size, dHy=hy_size,
```

Example - creating the `VerletListAdress` for a spherical address region centered on `adrCenter` and fixed in space:

```
>>> vl = espressopp.VerletListAdress(system, cutoff=rc, adrcut=rc, dEx=ex_size, dHy=hy_size,
```

Example - creating the `VerletListAdress` for a spherical address region centered on one particle and moving with the particle

```
>>> vl = espressopp.VerletListAdress(system, cutoff=rc, adrcut=rc, dEx=ex_size, dHy=hy_size,
```

Example - creating the `VerletListAdress` for a address region based on the overlapping spherical regions by several particles

```
>>> vl = espressopp.VerletListAdress(system, cutoff=rc, adrcut=rc, dEx=ex_size, dHy=hy_size,
```

`espressopp.VerletListAdress` (`system`, `cutoff`, `adrcut`, `dEx`, `dHy`, `adrCenter`, `pids`, `exclusionlist`, `sphereAdr`)

#### Parameters

- **system** –
- **cutoff** –
- **adrcut** –
- **dEx** –
- **dHy** –
- **adrCenter** – (default: [])
- **pids** – (default: [])
- **exclusionlist** – (default: [])
- **sphereAdr** – (default: False)

`espressopp.VerletListAdress.addAdrParticles` (`pids`, `rebuild`)

#### Parameters

- **pids** –
- **rebuild** – (default: True)

#### Return type

`espressopp.VerletListAdress.exclude` (`exclusionlist`)

#### Parameters `exclusionlist` –

#### Return type

`espressopp.VerletListAdress.rebuild` ()

#### Return type

`espressopp.VerletListAdress.totalSize` ()

Return type

## espressopp.VerletListTriple

`espressopp.VerletListTriple` (*system, cutoff, exclusionlist*)

Parameters

- **system** –
- **cutoff** –
- **exclusionlist** – (default: [])

`espressopp.VerletListTriple.exclude` (*exclusionlist*)

Parameters **exclusionlist** –

Return type

`espressopp.VerletListTriple.getAllTriples` ()

Return type

`espressopp.VerletListTriple.localSize` ()

Return type

`espressopp.VerletListTriple.totalSize` ()

Return type

## espressopp.Version

Return version information of espressopp module

Example:

```
>>> version = espressopp.Version()
>>> print "Name" = ", version.name
>>> print "Major version number" = ", version.major
>>> print "Minor version number" = ", version.minor
>>> print "Git revision" = ", version.gitrevision
>>> print "boost version" = ", version.boostversion
>>> print "Patchlevel" = ", version.patchlevel
>>> print "Compilation date" = ", version.date
>>> print "Compilation time" = ", version.time
```

to print a full version info string:

```
>>> print version.info()
```

`espressopp.Version` ()

## standard\_system

### espressopp.standard\_system.Default

`espressopp.standard_system.Default` (*box, rc = 1.12246, skin = 0.3, dt = 0.005, temperature = None*)

Parameters

- **box** –

- **rc** (*real*) –
- **skin** (*real*) –
- **dt** (*real*) –
- **temperature** –

Return default system and integrator, no interactions, no particles are set if tempearture is != None then Langevin thermostat is set to temperature (gamma is 1.0)

## espressopp.standard\_system.KGMelt

`espressopp.standard_system.KGMelt` (*num\_chains*, *chain\_len*)

### Parameters

- **num\_chains** –
- **chain\_len** –

## espressopp.standard\_system.LennardJones

`espressopp.standard_system.LennardJones` (*num\_particles*, *box*, *rc*, *skin*, *dt*, *epsilon*, *sigma*, *shift*, *temperature*, *xyzfilename*, *xyzrfilename*)

### Parameters

- **num\_particles** –
- **box** – (default: (000))
- **rc** (*real*) – (default: 1.12246)
- **skin** (*real*) – (default: 0.3)
- **dt** (*real*) – (default: 0.005)
- **epsilon** (*real*) – (default: 1.0)
- **sigma** (*real*) – (default: 1.0)
- **shift** – (default: 'auto')
- **temperature** – (default: None)
- **xyzfilename** – (default: None)
- **xyzrfilename** – (default: None)

return random Lennard Jones system and integrator: if tempearture is != None then Langevin thermostat is set to temperature (gamma is 1.0)

## espressopp.standard\_system.Minimal

`espressopp.standard_system.Minimal` (*num\_particles*, *box*, *rc*, *skin*, *dt*, *temperature*)

### Parameters

- **num\_particles** –
- **box** –
- **rc** (*real*) – (default: 1.12246)
- **skin** (*real*) – (default: 0.3)
- **dt** (*real*) – (default: 0.005)

- **temperature** – (default: None)

Return minimal system and integrator without any interactions defined: particles have random positions in box if temperature is != None then Langevin thermostat is set to temperature (gamma is 1.0)

## espressopp.standard\_system.PolymerMelt

`espressopp.standard_system.PolymerMelt` (*num\_chains*, *monomers\_per\_chain*, *box*, *bondlen*, *rc*, *skin*, *dt*, *epsilon*, *sigma*, *shift*, *temperature*, *xyzfilename*, *xyrfilename*)

### Parameters

- **num\_chains** –
- **monomers\_per\_chain** –
- **box** – (default: (000))
- **bondlen** (*real*) – (default: 0.97)
- **rc** (*real*) – (default: 1.12246)
- **skin** (*real*) – (default: 0.3)
- **dt** (*real*) – (default: 0.005)
- **epsilon** (*real*) – (default: 1.0)
- **sigma** (*real*) – (default: 1.0)
- **shift** – (default: 'auto')
- **temperature** – (default: None)
- **xyzfilename** – (default: None)
- **xyrfilename** – (default: None)

returns random walk polymer melt system and integrator: if temperature is != None then Langevin thermostat is set to temperature (gamma is 1.0)

## storage

### espressopp.storage.DomainDecomposition

`espressopp.storage.DomainDecomposition` (*system*, *nodeGrid*, *cellGrid*)

### Parameters

- **system** –
- **nodeGrid** –
- **cellGrid** –

`espressopp.storage.DomainDecomposition.getCellGrid()`

### Return type

`espressopp.storage.DomainDecomposition.getNodeGrid()`

### Return type



## espressopp.storage.DomainDecompositionAddress

The DomainDecompositionAddress is the Domain Decomposition for AdResS and H- AdResS simulations. It makes sure that tuples (i.e. a coarse-grained particle and its corresponding atomistic particles) are always stored together on one CPU. When setting DomainDecompositionAddress you have to provide the system as well as the nodegrid and the cellgrid.

Example - setting DomainDecompositionAddress:

```
>>> system.storage = espressopp.storage.DomainDecompositionAddress(system, nodeGrid, cellGrid)

espressopp.storage.DomainDecompositionAddress (system, nodeGrid, cellGrid)
```

### Parameters

- **system** –
- **nodeGrid** –
- **cellGrid** –

## espressopp.storage.DomainDecompositionNonBlocking

```
espressopp.storage.DomainDecompositionNonBlocking (system, nodeGrid, cellGrid)
```

### Parameters

- **system** –
- **nodeGrid** –
- **cellGrid** –

## espressopp.storage.Storage

This is the base class for all storage objects. All derived classes implement at least the following methods:

- *decompose()*  
Send all particles to their corresponding cell/cpu
- *addParticle(pid, pos):*  
Add a particle to the storage
- *removeParticle(pid):*  
Remove a particle with id number *pid* from the storage.

```
>>> system.storage.removeParticle(4)
```

There is an example in *examples* folder

- *getParticle(pid):*  
Get a particle object. This can be used to get specific particle information:

```
>>> particle = system.storage.getParticle(15)
>>> print "Particle ID is      : ", particle.id
>>> print "Particle position is : ", particle.pos
```

you cannot use this particle object to modify particle data. You have to use the modifyParticle command for that (see below).

- *addAdrParticle(pid, pos, last\_pos):*  
Add an AdResS Particle to the storage

- *setFixedTuplesAdress(fixed\_tuple\_list):*
- *addParticles(particle\_list, \*properties):*

This routine adds particles with certain properties to the storage.

**param particleList** list of particles (and properties) to be added

**param properties** property strings

Each particle in the list must be itself a list where each entry corresponds to the property specified in properties.

Example:

```
>>> addParticles([[id, pos, type, ... ], ...], 'id', 'pos', 'type', ...)
```

- *modifyParticle(pid, property, value, decompose='yes')*

This routine allows to modify any properties of an already existing particle.

Example:

```
>>> modifyParticle(pid, 'pos', Real3D(new_x, new_y, new_z))
```

- *removeAllParticles():*

This routine removes all particles from the storage.

- 'system':

The property 'system' returns the System object of the storage.

Examples:

```
>>> s.storage.addParticles([[1, espressopp.Real3D(3,3,3)], [2, espressopp.Real3D(4,4,4)]], 'id', 'p
>>> s.storage.decompose()
>>> s.storage.modifyParticle(15, 'pos', Real3D(new_x, new_y, new_z))
```

`espressopp.storage.Storage.addAdrATParticle (pid, *args)`

**Parameters**

- **pid** –
- **\*args** –

**Return type**

`espressopp.storage.Storage.addParticle (pid, pos)`

**Parameters**

- **pid** –
- **pos** –

**Return type**

`espressopp.storage.Storage.addParticles (particleList, *properties)`

**Parameters**

- **particleList** –
- **\*properties** –

**Return type**

`espressopp.storage.Storage.clearSavedPositions ()`

**Return type**

`espressopp.storage.Storage.getParticle (pid)`

**Parameters** `pid` –

**Return type**

`espressopp.storage.Storage.getRealParticleIDs()`

**Return type**

`espressopp.storage.Storage.modifyParticle(pid, property, value)`

**Parameters**

- `pid` –
- `property` –
- `value` –

**Return type**

`espressopp.storage.Storage.particleExists(pid)`

**Parameters** `pid` –

**Return type**

`espressopp.storage.Storage.printRealParticles()`

**Return type**

`espressopp.storage.Storage.removeAllParticles()`

**Return type**

`espressopp.storage.Storage.removeParticle(pid)`

**Parameters** `pid` –

**Return type**

`espressopp.storage.Storage.restorePositions()`

**Return type**

`espressopp.storage.Storage.savePositions(idList)`

**Parameters** `idList` –

**Return type**

`espressopp.storage.Storage.setFixedTuplesAdress(fixedtuples)`

**Parameters** `fixedtuples` –

## tools

### information and analysis

#### Overview

---

`espressopp.tools.analyse`

---

`espressopp.tools.info`

---

`espressopp.tools.timers`

---

`espressopp.tools.vmd`

---

#### Details

## espressopp.tools.analyse

`espressopp.tools.analyse.final_info` (*system, integrator, vl, start\_time, end\_time*)  
final report on the simulation statistics

`espressopp.tools.analyse.info` (*system, integrator, per\_atom=False*)  
reports on the simulation progress

## espressopp.tools.info

`espressopp.tools.info.getAllBonds` (*system*)  
return all bonds of the system (currently only FixedPairLists are supported)

`espressopp.tools.info.getAllParticles` (*system, \*properties*)  
returns a list of all particle properties of all particles of the system (currently no atomistic AdResS particles are included)

## espressopp.tools.timers

`espressopp.tools.timers.show` (*alltimers, precision=1*)  
Python functions to print timings from C++.

## espressopp.tools.vmd

`espressopp.tools.vmd.connect` (*system, molsize=10, pqrfile=False, vmd\_path='vmd'*)  
Connects to the VMD.

### Parameters

- **system** (*espressopp.system*) – The system object.
- **molsize** (*int*) – The optional size of the molecule.
- **pqrfile** (*bool*) – If set to True then the pqr vmd.pqr file will be used otherwise (default) the vmd.pdb file will be used.
- **vmd\_path** (*str*) – The path to the executable of vmd, by default it is set to 'vmd'.

**Returns** Socket to the VMD.

## initializing particles

### Overview

<code>espressopp.tools.lattice</code>	
<code>espressopp.tools.replicate</code> (bonds, angles, x, ...)	Replicates configuration in each dimension.
<code>espressopp.tools.topology</code>	
<code>espressopp.tools.velocities</code>	
<code>espressopp.tools.warmup</code> (system, integrator)	Warm up for a system with a density of 0.85.

### Details

## espressopp.tools.lattice

`espressopp.tools.lattice.createCubic` (*N, rho, perfect=True, RNG=None*)  
Initializes particles on the sites of a simple cubic lattice. By setting `perfect=False` the particle positions will be given random displacements with a magnitude of one-tenth the lattice spacing.

`espressopp.tools.lattice.createDiamond` (*N, rho, perfect=True, RNG=None*)  
 Initializes particles on the sites of a diamond lattice.

### espressopp.tools.replicate

`espressopp.tools.replicate.replicate` (*bonds, angles, x, y, z, Lx, Ly, Lz, xdim=1, ydim=1, zdim=1*)

Replicates configuration in each dimension.

This may be used to increase the size of an equilibrated melt by a factor of 8 or more.

Presently this routine works only for semiflexible polymers. A general class should be written to deal with files containing coordinates and topology data.

`xdim = ydim = zdim = 1` returns the original system not replicated. `xdim = ydim = zdim = 2` returns the original system replicated to 8x. `xdim = ydim = zdim = 3` returns the original system replicated to 27x. `xdim = ydim = 1, zdim = 2` returns the original system replicated in the z-direction.

### espressopp.tools.topology

`espressopp.tools.topology.polymerRW` (*pid, startpos, numberOfMonomers, bondlength, return\_angles=False, return\_dihedrals=False, mindist=None, rng=None*)

Initializes polymers through random walk

### espressopp.tools.velocities

`espressopp.tools.velocities.gaussian` (*T, N, particle\_mass=None, zero\_momentum=True, seed=7654321, kb=1.0*)

Generates velocities with temperature T according to a Maxwell-Boltzmann distribution.

**Args:** T: The desired temperature expre. N: The number of particles. `particle_mass`: The list of particle mass if not then every particle has mass 1.0 `zero_momentum`: Remove the center-of-mass motion. `seed`: The seed for the random number generator. `kb`: The Boltzmann constant.

**Returns:** The tuple with lists of x,y,z components of the velocity.

### espressopp.tools.warmup

`espressopp.tools.warmup.warmup` (*system, integrator, number=80*)

Warm up for a system with a density of 0.85.

The method needs the following parameters:

- `system`, `integrator` ESPResSo system which schoul be warmed up and the correspondig integrator e.g.:

```
>>> system, integrator = espressopp.standard_system.LennardJones(100, (10, 10, 10))
```

- `number` number of steps of the warm up

for a system with a density of 0.85, if it explodes try a higher number

## decomp - Domain Decomposition python functions

- `nodeGrid` (*n, box\_size, rc, skin, eh\_size=0, ratioMS=0, idealGas=0, slabMSDims=[0,0,0]*):

It determines how the processors are distributed and how the cells are arranged. The algorithm is dimensional sensitive for both homogeneous and inhomogeneous setups. On top of such functionality it presents specific features for region divided heterogenous setups (e.g. for

AdResS) [see H.V. Guzman et. al, Phys. Rev. E, 96, 053311 (2017)] Link to the paper: <https://doi.org/10.1103/PhysRevE.96.053311>

*box\_size* - 3D vector containing the size of the simulation box *box\_size* [*L\_x*,*L\_y*,*L\_z*] *rc* - cutoff radius of interaction *skin* - skin size for the verlet list calculation *n* - total number of processes *eh\_size* - 1D length of the high-resolution region (e.g. for AdResS Atomistic/Explicit+Hybrid regions) *ratioMS* - spatial mapping ratio between high-resolution region and the low-resolution one

(e.g. for AdResS mapping the atomistic water molecule to the CG-model; leads to a *ratioMS*=3)

***idealGas* - this is a Flag for treating the low-resolution region as an Ideal Gas, when TRUE (no interactions included, thus none force computations load)**

*slabMSDims* - 3D vector containing flags describing the type of axis, if heterogeneous value is 1, else 0

- ***nodeGridSimple(n)***: It determines how the processors are distributed and how the cells are arranged. Note: Use it exclusively for Lattice-Boltzmann simulations, or non-parallelized tests. *n* - number of processes
- ***cherryPickTotalProcs(box\_size,rc,skin,MnN,CpN,percTol=0.2,eh\_size=0,ratioMS=0,idealGas=0,slabMSDims=[0,0,0])***:

To be used for heterogeneous simulations where the spatial heterogeneity is known on an a-priori manner, where this function returns *n* as the total number of processes to be used for the best decomposition of the system as a function of a tolerance ratio which depending on the giving range [0, *MnN*\**CpN*] of processors availability different combinations of *P\_x*,*P\_y* and *P\_z* can be found and hence several values of *n* this 'n' can become an array. Most of the parameters have been described for *nodeGrid(...)*, except:

*MnN* - M number of Nodes to be available (e.g. 128 processes/cores, 16 cores per Node gives a total of 8 Nodes) *CpN* - C number of Cores available in each Node (e.g. 16 Cores per Node or 20 Cores per Node) *percTol* - Axis base tolerance percentage to the ideal distribution of P-processors per axes *P\_x*,*P\_y*,*P\_z*

- ***neiListHom(node\_grid,box,rc,skin)***:

The new domain decomposition divides the subdomains in a neighborlist of coarse in a grid of 3 arrays [*N\_x*,*N\_y*,*N\_z*]. In this case, the neighbor list is homogeneous (non a-priori load imbalance). Most of the parameters have been described above, except:

*node\_grid* - M number of Nodes to be available (e.g. 128 processes/cores, 16 cores per Node gives a total of 8 Nodes)

- ***neiListAdress(node\_grid, cell\_grid,rc,skin,eh\_size,adrCenter,ratioMS,idealGasFlag=True,sphereAdr=False,slabMSDims=[1,1,1])***:

The new domain decomposition divides the subdomains in a neighborlist of coarse in a grid of 3 arrays [*N\_x*,*N\_y*,*N\_z*]. In this case, the neighbor list is homogeneous (non a-priori load imbalance). Most of the parameters have been described above, except:

*cell\_grid* - Based on the homogeneous allocation of cells per subdomain, a referential value *adrCenter* - Box center of the heterogeneous simulation box [*adrCenter\_x*,*adrCenter\_y*,*adrCenter\_z*], commonly the middle of

high-resolution region.

***idealGasFlag* - This is a Flag for treating the low-resolution region as an Ideal Gas (no interactions included, thus less load)**

*sphereAdr* - Geometry of the high-resolution region, if TRUE spherical, otherwise Slab-like

- ***cellGrid(box\_size, node\_grid, rc, skin)***:

It returns an appropriate grid of cells.

- ***tuneSkin(system, integrator, minSkin=0.01, maxSkin=1.2, precision=0.001)***:

It tunes the skin size for the current system

- `printTimeVsSkin(system, integrator, minSkin=0.01, maxSkin=1.5, skinStep = 0.01):`

It prints time of running versus skin size in the range [minSkin, maxSkin] with the step skinStep

## DumpConfigurations - read/write xyz files

`espressopp.tools.DumpConfigurations.fastreadxyz(filename)`

`espressopp.tools.DumpConfigurations.fastwritexyz(filename, system, velocities=True, unfolded=True, append=False, scale=1.0)`

`espressopp.tools.DumpConfigurations.fastwritexyz_standard(filename, system, unfolded=False, append=False)`

Fast write standard xyz file. Generally standard xyz file is

```
>>> number of particles
>>> comment line
>>> type x y z
>>> .....
>>> .....
>>> .....
```

Additional information can be found here: Wiki: [http://en.wikipedia.org/wiki/XYZ\\_file\\_format](http://en.wikipedia.org/wiki/XYZ_file_format) OpenBabel: [http://openbabel.org/wiki/XYZ\\_%28format%29](http://openbabel.org/wiki/XYZ_%28format%29)

In this case one can choose folded or unfolded coordinates. Currently it writes only particle type = 0 and pid is a line number. Later different types should be implemented.

`espressopp.tools.DumpConfigurations.readxyz(filename)`

`espressopp.tools.DumpConfigurations.readxyzr(filename)`

`espressopp.tools.DumpConfigurations.writexyz(filename, system, velocities=True, unfolded=False, append=False)`

`espressopp.tools.DumpConfigurations.xyzfilewrite(filename, system, append=False, atomtypes={0: 'Fe', 1: 'O', 2: 'C'}, velocities=False, charge=False)`

This method creates a xyz file with the data from a specific system: 1. row: number of the atoms 2. row: REMARK generated by ESPResSo++ following rows: atomsymbol positionX positionY positionZ (velocityX velocityY velocityZ) (charge) last row: END

The method needs the following parameters:

- filename

name of the file where the table should be saved in

- system

ESPResSo system which creates the data e.g.:

```
>>> system, integrator = espressopp.standard_system.LennardJones(100, (10, 10, 10))
```

- append

**=False** the data in the file will be overwritten

**=True** the data will be appended

- atomtypes the xyz file needs atom symbols, so it has to translate the numbers insert a dictionary with the right translation

- velocities

**=False** does not save the velocity vectors

**=True** creates columns for the velocity vectors and saves the data

•charge

**=False** does not save the charge

**=True** creates columns for the charges and saves the data

## espresso\_old - read espressomd files

This Python module allows one to use ESPResSo data files as the input to an ESPResSo++ simulation.

`espressopp.tools.espresso_old.read(file)`

Read ESPResSo data files.

Keyword argument: `file` – contains simulation variables, data of all particles, and information about bonds. (angles and dihedrals are currently not read)

## gromacs - parser for Gromacs files

This Python module allows one to use GROMACS data files as the input to an ESPResSo++ simulation, set interactions for given particle types and convert GROMACS potential tables into ESPResSo++ tables. It contains functions: `read()`, `setInteractions()`, `convertTable()`

Some tips for using the gromacs parser:

### Tip 1.

`topol.top` includes solvent via `#include` statements

If the included `.itp` file ONLY contains the solvent molecule you're using (e.g. `spc/e` water using `spce.itp`) then this is okay.

But if the `.itp` file contains info about many molecules (e.g. you want to use one ion from `ions.itp`), then `gromacs.py` will just take the first one listed. You must edit your `topol.top` file to explicitly include the solvent molecule you're using.

e.g. replace:

```
; Include topology for ions
#include "amber03.ff/ions.itp"
```

by:

```
; Include topology for ions
[ moleculetype ]
; molname          nrexcl
CL                  1

[ atoms ]
; id   at type          res nr  residu name      at name  cg nr  charge
1      CL              1       CL              CL       1     -1.00000
```

### Tip 2. impropers

impropers in the `topol.top` file (function type 4) need to be labelled '[ impropers ]', not '[ dihedrals ]' as in standard gromacs format"

Also, the dihedrals should be listed before the impropers (this is usually the case by default in gromacs-format files).

### Tip 3.

For rigid SPC/E water using Settle, `spce.itp` file should look like this:



```
[ moleculetype ]
; molname      nrexcl
SOL             2

[ atoms ]
; id  at type      res nr  res name  at name  cg nr  charge  mass
1   OW_spc        1       SOL      OW       1      -0.8476 15.99940
2   HW_spc        1       SOL      HW1      1       0.4238  1.00800
3   HW_spc        1       SOL      HW2      1       0.4238  1.00800

[ bonds ]
; i      j      funct  length  force.c.
1        2        1     0.1    345000  0.1    345000
1        3        1     0.1    345000  0.1    345000

[ angles ]
; i      j      k      funct  angle  force.c.
2        1      3        1    109.47 383    109.47 383
```

The bonds section is used to generate exclusions, but bond and angle parameters are not relevant if the Settle extension is used. The geometry is that specified in the python script when adding the Settle extension

Include modified spce file in `topol.top`, e.g. replace

```
#include "amber03.ff/spce.itp"
```

by

```
#include "amber03.ff/spce-for-espressopp.itp"
```

#### Tip 4.

Use absolute paths for any include files which are not in the standard gromacs topology directory (\$GMXLIB)

e.g. replace

```
#include "mynewresidue.itp"
```

by

```
#include "path/to/mynewres/file/mynewresidue.itp"
```

#### Tip 5.

The parser won't work if the particles ids in the include files conflict with the particle ids in the `topol.top` file itself, and the bonded interaction parameters in the `itp` file need to be looked up via particle type in the standard gromacs topology directory (\$GMXLIB)

i.e. Okay for an `itp` file like `spce.itp` above, where the bonds and angles parameters are given in the `itp` file, as in:

```
[ bonds ]
; i      j      funct  length  force.c.
1        2        1     0.1    345000  0.1    345000
```

Not okay for an `itp` file containing lines like:

```
[ bonds ]
; i      j      funct  length  force.c.
1        2        1
```

```
espressopp.tools.gromacs.convertTable (gro_in_file, esp_out_file, sigma=1.0, epsilon=1.0,
                                         c6=1.0, c12=1.0)
```

Convert GROMACS tabulated file into ESPResSo++ tabulated file (new file is created). First column of input file can be either distance or angle. For non-bonded files, `c6` and `c12` can be provided. Default value for `sigma`, `epsilon`, `c6` and `c12` is 1.0. Electrostatics are not taken into account (`f` and `fd` columns).

Keyword arguments: `gro_in_file` – the GROMACS tabulated file name (bonded, nonbonded, angle or dihedral). `esp_out_file` – filename of the ESPResSo++ tabulated file to be written. `sigma` – optional, depending on whether you want to convert units or not. `epsilon` – optional, depending on whether you want to convert units or not. `c6` – optional `c12` – optional

```
espressopp.tools.gromacs.read(gro_file, top_file='', doRegularExcl=True)
Read GROMACS data files.
```

Arguments: `:param gro_file` – contains coordinates of all particles, the number of particles, velocities and box size. `:type gro_file`: string `:param top_file` – contains topology information. Included topology files (.itp) are also read `:type gro_file`: string `:param doRegularExcl` – if True, exclusions are generated automatically based on the `nregxcl` parameter (see gromacs manual) `:type doRegularExcl`: bool

```
espressopp.tools.gromacs.setLennardJones14Interactions(system, defaults, atom-
typeparams, onefourlist,
cutoff)
Set lennard jones interactions which were read from gromacs based on the atomtypes
```

```
espressopp.tools.gromacs.setLennardJonesInteractions(system, defaults, atom-
typeparams, verletlist,
cutoff, hadress=False,
adress=False, fpl=None)
Set lennard jones interactions which were read from gromacs based on the atomtypes
```

```
espressopp.tools.gromacs.setLennardJonesInteractionsTI(system, defaults, atom-
typeparams, verletlist,
cutoff, epsilonB, sig-
maSC, alphaSC, pow-
erSC, lambdaTI, pidlist,
annihilate=True,
hadress=False,
adress=False,
fpl=None)
Set lennard jones interactions which were read from gromacs based on the atomtypes
```

```
espressopp.tools.gromacs.setTabulatedInteractions(potentials, particleTypes, sys-
tem, interaction)
Set interactions for all given particle types. Return value is a system with all interactions added.
```

Keyword arguments: `potentials` – is a dictionary where key is a string composed of two particle types and value is a potential. example: {"A\_A":potAA, "A\_B":potAB, "B\_B":potBB} `particleTypes` – is a dictionary where key is the particle type, and value is a list of particles of that type. example: {"A":["A1m", "A2m"],"B":["B1u","B2u"]} `system` – is the system to which the interaction will be added `interaction` – is the interaction to which to add the potentials

## io\_extended - read/write configurational files

This Python module allows one to read and write configurational files. One can choose folded or unfolded coordinates and write down velocities or not. It is similar to lammps read and write, but it writes down only: 1) number of particles + types 2) number of bonds (number of pairs) + types 3) number of angles (number of triples) + types 4) number of dihedrals (number of quadruples) + types 5) system size (Lx,Ly,Lz) 6) `p_id`, `p_type`, `p_positions` 7) velocities (if true) 8) bonds (if exist) 9) angles (if exist) 10) dihedrals (if exist)

`read` returns: `Lx`, `Ly`, `Lz`, `p_ids`, `p_types`, `pos`, `vels`, `bonds`, `angles`, `dihedrals` if something does not exist then it will return the empty list `bonds`, `angles`, `dihedrals` - will return list `[type, (x,x,x,x)]`, where `type` is the type of bond, angle or dihedral `(x,x,x,x)` is `(pid1,pid2)` for bonds,

`(pid1,pid2,pid3)` for angles

`(pid1,pid2,pid3,pid4)` for dihedrals

## lammmps - read lammmps files

This Python module allows one to use a LAMMPS data file as the input to an ESPResSo++ simulation.

## pathintegral - nuclear quantum effects

- method to automatically run the system including nuclear quantum effects using the Feynman path-integral

!!WARNING: THIS IS STILL AN EXPERIMENTAL FEATURE!!

This method creates, based on the supplied topology of the system, an path-integral representation with  $P$  beads. The path-integral system is a fully classical analog, which has to be run at an effective temperature  $P*T$ .

The method needs the following parameters:

- **allParticles** particles of the sytem
- **props** particle properties
- **types** types, e.g. read from the gromacs parser
- **system**
- **exclusions** non-bonded exclusions
- **integrator**
- **langevin** langevin integrator
- **rcut** the cutoff used for the rings non-bonded interactions
- **P** the Trotter Number (number of imaginary time slices)
- **polymerInitR** polymer radius for setting up ring in 2d plane
- **hbar** hbar in gromacs units [kJ/mol ps]
- **disableVVI** disable Virtual Verlet List (slow but safe). If false, the neighbour search is based on the VirtualParticles extension, which contain the rings. This speeds up neighbour search significantly.

## PDB - read and write pdb format

`espressopp.tools.pdbwrite(filename, system, molsize=4, append=False, typenames=None)`

Writes a file in PDB format

### Parameters

- **filename** (*string*) – output file name
- **system** (*espressopp System object*) – espressopp system
- **molsize** (*int*) – if molsize>0, the molecule count is increased every molsize particles (default 4)
- **append** (*bool*) – if True, append to filename, other over-write filename (default False)
- **typenames** (*dict, key=int, value=string*) – dictionary used for mapping from espressopp's integer particle types to the particle type strings written in a pdb file

`espressopp.tools.pdbread(filename, natoms, header)`

Reads one frame of a pdb format file

### Parameters

- **filename** (*string*) – input file name
- **natoms** (*int*) – number of atoms in pdf file
- **header** (*int*) – number of header lines to skip at start of file

Returns: index, atomname, resname, resid, x, y, z, alpha, beta, segid, element (lists of type int, str, str, int, float, float, float, float, float, str, str)

## povwrite - write povray files

## prepareComplexMolecules - set up proteins

various helper functions for setting up systems containing complex molecules such as proteins

`espressopp.tools.findConstrainedBonds(atomPids, bondtypes, bondtypeparams, masses, massCutoff = 1.1)`

Finds all heavyatom-hydrogen bonds in a given list of particle IDs, and outputs a list describing the bonds, in a format suitable for use with the RATTLE algorithm for constrained bonds

### Parameters

- **atomPids** (*list of int*) – list of pids of atoms between which to search for bonds
- **bondtypes** (*dict, key: (int, int), value: int*) – dictionary mapping from tuple of pids to bondtypeid, e.g. as returned by `tools.gromacs.read()`
- **bondtypeparams** (*dict, key: int, value: espressopp bond type*) – dictionary mapping from bondtypeid to class storing parameters of that bond type, e.g. as returned by `tools.gromacs.read()`
- **masses** (*list of float*) – list of masses, e.g. as returned by `tools.gromacs.read()`
- **massCutoff** (*float*) – for identifying light atoms (hydrogens), default 1.1 mass units, can also be increased e.g. for use with deuterated systems

### Returns

hydrogenIDs - list of pids (integer) of light atoms (hydrogens)

constrainedBondsDict - dict, keys: pid (integer) of heavy atom, values: list of pids of light atoms that are bonded to it

constrainedBondsList - list of lists, one entry for each constrained bond with format: [pid of heavy atom, pid of light atom, bond distance, mass of heavy atom, mass of light atom]

Can then be used with RATTLE, e.g.

```
>>> rattle = espressopp.integrator.Rattle(system, maxit = 1000, tol = 1e-6, rptol = 1e-6)
>>> rattle.addConstrainedBonds(constrainedBondsList)
>>> integrator.addExtension(rattle)
```

`espressopp.tools.getInternalNonbondedInteractions(atExclusions, pidlist)`

Gets the non-bonded pairs within a list of particle indices, excluding those which are in a supplied list of exclusions. Useful for example for getting the internal atomistic non-bonded interactions in a coarse-grained particle and adding them as a fixedpairlist

### Parameters

- **atExclusions** (*list of 2-tuples of int*) – list of excluded pairs
- **pidlist** (*list of int*) – list of pids among which to create pairs

**Returns** list of pairs which are not in atExclusions

**Return type** list of 2-tuples of int

`espressopp.tools.readSimpleSystem(filename, nparticles, header)`

Read in a column-formatted file containing information about the particles in a simple system, for example a coarsegrained protein.

This function expects the input file to have between 2 and 5 columns. The number of columns in the file is automatically detected. The function reads each column into a list and returns the lists. Column types are interpreted as follows: 2 columns: float, float 3 columns: float, float, int 4 columns: float, float, int, str 5 columns: float, float, int, str, str

For example in the case of a coarsegrained protein model, these could be: mass, charge, corresponding atomistic index, beadname, beadtype

#### Parameters

- **filename** (*string*) – name of file to open and read
- **nparticles** (*int*) – number of particles in file
- **header** (*int*) – number of lines to skip at start of file (default 0)

Returns: between 2 and 5 lists

`espressopp.tools.applyBoreschRestrains` (*system*, *restraintAtoms*, *restraintK*, *restraintR0*)

Applies restraints between ligand and protein as defined in Boresch et al, JPCB 2003, 107, 9535-9551 The restraints (one bond, two angles and three dihedrals) are applied between three ligand atoms A,B,C and three protein atoms a,b,c.

In espressopp, the potential for harmonic bond and angles is  $k(x-x_0)^2$ , but the harmonic dihedral potential is  $0.5*k(x-x_0)^2$ . This is taken care of in this function, i.e. the user should supply the force constants exactly as given in Boresch et al.

#### Parameters

- **system** (*System object*) – espressopp system
- **restraintAtoms** (*python dictionary*) – dictionary identifying the six atoms involved in the restraints, key = atom label (one of 'A','B','C','a','b','c'), value = atom index
- **restraintK** (*python dictionary*) – dictionary of force constants, key = restraint label ('aA' for the bond, 'baA' and 'aAB' for the angles, 'aABC', 'cbaA' and 'baAB' for the dihedrals), value = force constant (units as for the simulation forcefield, angle and dihedral constants should be in  $\text{rad}^{-2}$ )
- **restraintR0** (*python dictionary*) – dictionary of equilibrium values (distance or angle), key = restraint label, value = distance (in distance units) or angle (in degrees)

Examples of the three dictionaries:

```
>>> restraintAtoms = {'A':1981, 'B':1966, 'C':1993, 'a':1588, 'b':1581, 'c':1567}
>>> restraintK = {'aA':4184, 'baA':41.84, 'aAB':41.84, 'aABC':41.84, 'cbaA':41.84, 'baAB':41.84} #
>>> restraintR0 = {'aA':0.31, 'baA':120.0, 'aAB':90.0, 'aABC':100.0, 'cbaA':-170.0, 'baAB':-105.0}
```

## prepareAdress - setup AdResS simulation

Auxiliary python functions for preparation of an Adress Simulation based on a configuration from an all-atomistic simulation.

If one uses a configuration file from an all-atomistic simulation as start configuration for an AdResS simulation, the particles are probably all located inside the simulation box. However, in AdResS only the coarse-grained center-of-mass particles have to be in the box, the atomistic particles of the coarse grained might be outside around their CoM CG particle. When in the start configuration atomistic particles belonging to a molecule are folded such that some of the atoms are on the one side of the box while the others are folded to the other side the calculation of the center of mass goes wrong and the simulation will be incorrect. This script ensures a proper center of mass calculation and a proper folding and configuration for the AdResS simulation by simply putting the CG particle in one of the atoms (AdressSetCG) first. Then the molecules will be put together properly afterwards when calling AdressDecomp.

## PSF - read and write psf format

PSF file format given at <http://www.ks.uiuc.edu/Training/Tutorials/namd/namd-tutorial-win-html/node24.html>

`espressopp.tools.psfwrite` (*filename*, *system*, *maxdist=None*, *molsize=4*, *typenames=None*)  
Writes !NATOM and !NBOND sections of a psf format file

### Parameters

- **filename** (*string*) – output file name
- **system** (*espressopp System object*) – espressopp system
- **maxdist** (*float*) – if this is specified, only bonds in which the pair of particles are separated by a distance < maxdist are written to the !NBOND section
- **molsize** (*int*) – if molsize>0, the molecule count is increased every molsize particles
- **typenames** (*dict*, *key=int*, *value=string*) – dictionary used for mapping from espressopp's integer particle types to the particle type strings written in a psf file

`espressopp.tools.psfread` (*filename*)  
Reads !NATOM section of a psf format file

**Parameters** **filename** (*string*) – input file name

**Returns** pid, segname, resindex, resname, atomname, atomtype, mass, charge (lists of type int, str, int, str, str, str, float, float)

## tabulated - write tabulated file

`espressopp.tools.tabulated.writeTabFile` (*pot*, *name*, *N*, *low=0.0*, *high=2.5*, *body=2*)  
writeTabFile can be used to create a table for any potential Parameters are: \* *pot* : this is any espressopp.interaction potential \* *name* : filename \* *N* : number of line to write \* *low* : lowest r (default is 0.0) \* *high* : highest r (default is 2.5)

This function has not been tested for 3 and 4 body interactions

## topology\_helper

### units - convert to real units

Espresso++ returns temperature, energy, pressure, box length etc. in dimensionless units. Usually user should take care about real length, energy, mass and charge units. This python class is a helper in order to simplify the conversion which is based on basic units. However, user always should use it carefully for complicated systems.

Currently it is implemented for SI units. Make sure that you are using length in [nm] energy in [kJ/mol] mass in [amu] q in [e]

and it will return you pressure in [bar] temperature in [K] time in [ps] density in [kg/m^3]

Example:

## Logging mechanism

ESPResSo++ uses Loggers

Logging can be switched on in your python script with the following command:

```
>>> logging.getLogger("*name of the logger*").setLevel(logging.*Level*)
```

*Level* is one of the following:

ERROR	for errors that might still allow the application to continue
WARN	for potentially harmful situations
INFO	informational messages highlighting progress
DEBUG	designates fine-grained informational events

Example:

```
>>> import espressopp
>>> import logging
>>> logging.getLogger("Storage").setLevel(logging.ERROR)
```

To log everything (WARNING: this will produce **lots** of output):

```
>>> logging.getLogger("").setLevel(logging.DEBUG)
```

The following loggers are currently available:

- Configurations
- Observable
- Velocities
- BC
- Logger
- FixedListComm
- FixedPairList
- FixedQuadrupleList
- FixedTripleList
- FixedTupleList
- Langevin
- MDIntegrator
- AngularPotential
- DihedralPotential
- Interaction
- InterpolationAkima
- InterpolationCubic
- InterpolationLinear
- InterpolationTable
- Potential
- CellListAllPairsIterator
- DomainDecomposition.CellGrid
- DomainDecomposition
- DomainDecomposition.NodeGrid
- Storage
- DomainDecompositionAdress
- StorageAdress
- VerletList

- VerletList



**CREDITS**

## ESPResSo++ Developers

The core of the developer team comes from the [Polymer Theory Group](#) of Prof. Kurt Kremer at the [Max Planck Institute for Polymer Research](#) in Mainz.

A full list of active and former developers is available at the [main website](#) of ESPResSo++.

## FAQ

A short list of frequently asked questions is given [here](#).

## Getting Help

If you have any questions do not hesitate to [contact us](#).



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**e**

- espressopp.analysis.AdressDensity, 37
- espressopp.analysis.AllParticlePos, 27
- espressopp.analysis.AnalysisBase, 32
- espressopp.analysis.Autocorrelation, 32
- espressopp.analysis.CenterOfMass, 37
- espressopp.analysis.CMVelocity, 33
- espressopp.analysis.ConfigsParticleDecomp, 33
- espressopp.analysis.Configurations, 34
- espressopp.analysis.ConfigurationsExt, 34
- espressopp.analysis.Energy, 35
- espressopp.analysis.IntraChainDistSq, 36
- espressopp.analysis.LBOutput, 27
- espressopp.analysis.LBOutputScreen, 28
- espressopp.analysis.LBOutputVzInTime, 28
- espressopp.analysis.LBOutputVzOfX, 28
- espressopp.analysis.MaxPID, 37
- espressopp.analysis.MeanSquareDispl, 36
- espressopp.analysis.MeanSquareInternalDist, 36
- espressopp.analysis.NeighborFluctuation, 37
- espressopp.analysis.NPart, 38
- espressopp.analysis.Observable, 40
- espressopp.analysis.OrderParameter, 29
- espressopp.analysis.ParticleRadiusDistribution, 29
- espressopp.analysis.PotentialEnergy, 38
- espressopp.analysis.Pressure, 38
- espressopp.analysis.PressureTensor, 29
- espressopp.analysis.PressureTensorLayer, 30
- espressopp.analysis.PressureTensorMultiLayer, 31
- espressopp.analysis.RadialDistrF, 38
- espressopp.analysis.RDFatomistic, 38
- espressopp.analysis.StaticStructF, 39
- espressopp.analysis.SystemMonitor, 40
- espressopp.analysis.Temperature, 31
- espressopp.analysis.Test, 32
- espressopp.analysis.Velocities, 41
- espressopp.analysis.VelocityAutocorrelation, 41
- espressopp.analysis.Viscosity, 41
- espressopp.analysis.XDensity, 40
- espressopp.analysis.XPressure, 40
- espressopp.analysis.XTemperature, 40
- espressopp.bc.BC, 42
- espressopp.bc.OrthorhombicBC, 43
- espressopp.bc.SlabBC, 43
- espressopp.check.System, 44
- espressopp.esutil.Collectives, 44
- espressopp.esutil.GammaVariate, 44
- espressopp.esutil.Grid, 44
- espressopp.esutil.NormalVariate, 44
- espressopp.esutil.RNG, 44
- espressopp.esutil.UniformOnSphere, 44
- espressopp.Exceptions, 134
- espressopp.FixedLocalTupleList, 135
- espressopp.FixedPairDistList, 135
- espressopp.FixedPairList, 136
- espressopp.FixedPairListAdress, 136
- espressopp.FixedQuadrupleAngleList, 137
- espressopp.FixedQuadrupleList, 137
- espressopp.FixedQuadrupleListAdress, 138
- espressopp.FixedSingleList, 138
- espressopp.FixedTripleAngleList, 139
- espressopp.FixedTripleList, 139
- espressopp.FixedTripleListAdress, 140
- espressopp.FixedTupleList, 140
- espressopp.FixedTupleListAdress, 141
- espressopp.Int3D, 141
- espressopp.integrator.Adress, 44
- espressopp.integrator.AssociationReaction, 45
- espressopp.integrator.BerendsenBarostat, 45
- espressopp.integrator.BerendsenBarostatAnisotropic, 46
- espressopp.integrator.BerendsenThermostat, 47
- espressopp.integrator.CapForce, 48
- espressopp.integrator.DPDThermostat, 49

<code>espressopp.integrator.EmptyExtension,</code>	<a href="#">67</a>	<code>espressopp.interaction.AngularUniquePotential,</code>	<a href="#">67</a>
<a href="#">49</a>		<code>espressopp.interaction.ConstrainCOM,</code>	<a href="#">74</a>
<code>espressopp.integrator.ExtAnalyze,</code>	<a href="#">49</a>	<code>espressopp.interaction.ConstrainRG,</code>	<a href="#">75</a>
<code>espressopp.integrator.Extension,</code>	<a href="#">49</a>	<code>espressopp.interaction.Cosine,</code>	<a href="#">67</a>
<code>espressopp.integrator.ExtForce,</code>	<a href="#">50</a>	<code>espressopp.interaction.CoulombKSpaceEwald,</code>	<a href="#">68</a>
<code>espressopp.integrator.FixPositions,</code>	<a href="#">50</a>	<code>espressopp.interaction.CoulombKSpaceP3M,</code>	<a href="#">69</a>
<code>espressopp.integrator.FreeEnergyCompensation,</code>	<a href="#">50</a>	<code>espressopp.interaction.CoulombRSpace,</code>	<a href="#">70</a>
<code>espressopp.integrator.GeneralizedLangevinThermostat,</code>	<a href="#">51</a>	<code>espressopp.interaction.CoulombTruncated,</code>	<a href="#">72</a>
<a href="#">51</a>		<code>espressopp.interaction.CoulombTruncatedUniqueChar</code>	<a href="#">73</a>
<code>espressopp.integrator.Isokinetic,</code>	<a href="#">51</a>	<code>espressopp.interaction.DihedralHarmonic,</code>	<a href="#">75</a>
<code>espressopp.integrator.LangevinBarostat,</code>	<a href="#">51</a>	<code>espressopp.interaction.DihedralHarmonicCos,</code>	<a href="#">76</a>
<a href="#">51</a>		<code>espressopp.interaction.DihedralHarmonicNCos,</code>	<a href="#">77</a>
<code>espressopp.integrator.LangevinThermostat,</code>	<a href="#">53</a>	<code>espressopp.interaction.DihedralHarmonicUniqueCos,</code>	<a href="#">77</a>
<a href="#">53</a>		<code>espressopp.interaction.DihedralPotential,</code>	<a href="#">78</a>
<code>espressopp.integrator.LangevinThermostatDr,</code>	<a href="#">53</a>	<code>espressopp.interaction.DihedralRB,</code>	<a href="#">78</a>
<a href="#">53</a>		<code>espressopp.interaction.DihedralUniquePotential,</code>	<a href="#">79</a>
<code>espressopp.integrator.LangevinThermostatEwald,</code>	<a href="#">53</a>	<code>espressopp.interaction.FENE,</code>	<a href="#">79</a>
<a href="#">53</a>		<code>espressopp.interaction.FENECapped,</code>	<a href="#">80</a>
<code>espressopp.integrator.LangevinThermostatKSpace,</code>	<a href="#">54</a>	<code>espressopp.interaction.GravityTruncated,</code>	<a href="#">81</a>
<a href="#">54</a>		<code>espressopp.interaction.Harmonic,</code>	<a href="#">82</a>
<code>espressopp.integrator.LangevinThermostatP3M,</code>	<a href="#">54</a>	<code>espressopp.interaction.HarmonicTrap,</code>	<a href="#">83</a>
<a href="#">54</a>		<code>espressopp.interaction.HarmonicUnique,</code>	<a href="#">83</a>
<code>espressopp.integrator.LatticeBoltzmann,</code>	<a href="#">55</a>	<code>espressopp.interaction.Interaction,</code>	<a href="#">84</a>
<a href="#">55</a>		<code>espressopp.interaction.LennardJones,</code>	<a href="#">84</a>
<code>espressopp.integrator.LBInit,</code>	<a href="#">58</a>	<code>espressopp.interaction.LennardJones93Wall,</code>	<a href="#">87</a>
<code>espressopp.integrator.LBInitConstForce,</code>	<a href="#">59</a>	<code>espressopp.interaction.LennardJonesAutoBonds,</code>	<a href="#">88</a>
<a href="#">59</a>		<code>espressopp.interaction.LennardJonesCapped,</code>	<a href="#">90</a>
<code>espressopp.integrator.LBInitPeriodicForce,</code>	<a href="#">60</a>	<code>espressopp.interaction.LennardJonesEnergyCapped,</code>	<a href="#">93</a>
<a href="#">60</a>		<code>espressopp.interaction.LennardJonesExpand,</code>	<a href="#">96</a>
<code>espressopp.integrator.LBInitPopUniform,</code>	<a href="#">59</a>	<code>espressopp.interaction.LennardJonesGeneric,</code>	<a href="#">97</a>
<a href="#">59</a>		<code>espressopp.interaction.LennardJonesGromacs,</code>	<a href="#">100</a>
<code>espressopp.integrator.LBInitPopWave,</code>	<a href="#">59</a>	<code>espressopp.interaction.LennardJonesSoftcoreTI,</code>	<a href="#">102</a>
<a href="#">59</a>		<code>espressopp.interaction.LennardJonesSoftcoreTII,</code>	<a href="#">102</a>
<code>espressopp.integrator.MDIntegrator,</code>	<a href="#">60</a>		
<code>espressopp.integrator.MinimizeEnergy,</code>	<a href="#">60</a>		
<a href="#">60</a>			
<code>espressopp.integrator.OnTheFlyFEC,</code>	<a href="#">61</a>		
<code>espressopp.integrator.Rattle,</code>	<a href="#">62</a>		
<code>espressopp.integrator.Settle,</code>	<a href="#">63</a>		
<code>espressopp.integrator.StochasticVelocityRescaling,</code>	<a href="#">63</a>		
<a href="#">63</a>			
<code>espressopp.integrator.TDforce,</code>	<a href="#">63</a>		
<code>espressopp.integrator.VelocityVerlet,</code>	<a href="#">64</a>		
<a href="#">64</a>			
<code>espressopp.integrator.VelocityVerletOnGroup,</code>	<a href="#">64</a>		
<a href="#">64</a>			
<code>espressopp.integrator.VelocityVerletOnRadius,</code>	<a href="#">64</a>		
<a href="#">64</a>			
<code>espressopp.interaction.AngularCosineSquared,</code>	<a href="#">65</a>		
<a href="#">65</a>			
<code>espressopp.interaction.AngularHarmonic,</code>	<a href="#">65</a>		
<a href="#">65</a>			
<code>espressopp.interaction.AngularPotential,</code>	<a href="#">66</a>		
<a href="#">66</a>			
<code>espressopp.interaction.AngularUniqueCosineSquared,</code>	<a href="#">66</a>		
<a href="#">66</a>			
<code>espressopp.interaction.AngularUniqueHarmonic,</code>	<a href="#">66</a>		
<a href="#">66</a>			

[espressopp.interaction.LJcos](#), [104](#)  
[espressopp.interaction.MirrorLennardJones](#), [105](#)  
[espressopp.interaction.Morse](#), [106](#)  
[espressopp.interaction.OPLS](#), [108](#)  
[espressopp.interaction.Potential](#), [109](#)  
[espressopp.interaction.PotentialUniqueDist](#), [109](#)  
[espressopp.interaction.PotentialVSPHPair](#), [109](#)  
[espressopp.interaction.Quartic](#), [109](#)  
[espressopp.interaction.ReactionFieldGeneralized](#), [110](#)  
[espressopp.interaction.ReactionFieldGeneralizedII](#), [112](#)  
[espressopp.interaction.SingleParticlePotential](#), [114](#)  
[espressopp.interaction.SoftCosine](#), [114](#)  
[espressopp.interaction.StillingerWeberPairTerm](#), [115](#)  
[espressopp.interaction.StillingerWeberPairTermClamped](#), [118](#)  
[espressopp.interaction.StillingerWeberTripleTerm](#), [120](#)  
[espressopp.interaction.Tabulated](#), [122](#)  
[espressopp.interaction.TabulatedAngular](#), [124](#)  
[espressopp.interaction.TabulatedDihedral](#), [125](#)  
[espressopp.interaction.TersoffPairTerm](#), [126](#)  
[espressopp.interaction.TersoffTripleTerm](#), [127](#)  
[espressopp.interaction.VSpherePair](#), [128](#)  
[espressopp.interaction.VSphereSelf](#), [129](#)  
[espressopp.interaction.Zero](#), [129](#)  
[espressopp.io.DumpGRO](#), [131](#)  
[espressopp.io.DumpGROAdress](#), [132](#)  
[espressopp.io.DumpXYZ](#), [133](#)  
[espressopp.MultiSystem](#), [142](#)  
[espressopp.ParallelTempering](#), [142](#)  
[espressopp.Particle](#), [143](#)  
[espressopp.ParticleAccess](#), [143](#)  
[espressopp.ParticleGroup](#), [143](#)  
[espressopp.pmi](#), [144](#)  
[espressopp.Quaternion](#), [148](#)  
[espressopp.Real3D](#), [149](#)  
[espressopp.RealND](#), [150](#)  
[espressopp.standard\\_system.Default](#), [154](#)  
[espressopp.standard\\_system.KGMelt](#), [155](#)  
[espressopp.standard\\_system.LennardJones](#), [155](#)  
[espressopp.standard\\_system.Minimal](#), [155](#)  
[espressopp.standard\\_system.PolymerMelt](#), [156](#)  
[espressopp.storage.DomainDecomposition](#), [156](#)  
[espressopp.storage.DomainDecompositionAdress](#), [156](#)  
[espressopp.storage.DomainDecompositionNonBlocking](#), [157](#)  
[espressopp.storage.Storage](#), [157](#)  
[espressopp.System](#), [150](#)  
[espressopp.Tensor](#), [151](#)  
[espressopp.tools.analyse](#), [160](#)  
[espressopp.tools.decomp](#), [161](#)  
[espressopp.tools.DumpConfigurations](#), [161](#)  
[espressopp.tools.espresso\\_old](#), [164](#)  
[espressopp.tools.gromacs](#), [164](#)  
[espressopp.tools.info](#), [160](#)  
[espressopp.tools.io\\_extended](#), [166](#)  
[espressopp.tools.lammps](#), [166](#)  
[espressopp.tools.lattice](#), [160](#)  
[espressopp.tools.pathintegral](#), [167](#)  
[espressopp.tools.pdb](#), [167](#)  
[espressopp.tools.povwrite](#), [168](#)  
[espressopp.tools.prepareAdress](#), [169](#)  
[espressopp.tools.prepareComplexMolecules](#), [168](#)  
[espressopp.tools.psf](#), [169](#)  
[espressopp.tools.replicate](#), [161](#)  
[espressopp.tools.tabulated](#), [170](#)  
[espressopp.tools.timers](#), [160](#)  
[espressopp.tools.topology](#), [161](#)  
[espressopp.tools.topology\\_helper](#), [170](#)  
[espressopp.tools.units](#), [170](#)  
[espressopp.tools.velocities](#), [161](#)  
[espressopp.tools.vmd](#), [160](#)  
[espressopp.tools.warmup](#), [161](#)  
[espressopp.VerletList](#), [152](#)  
[espressopp.VerletListAdress](#), [152](#)  
[espressopp.VerletListTriple](#), [154](#)  
[espressopp.Version](#), [154](#)





## A

addForce() (espressopp.integrator.LBInit.espressopp.integrator.LBInit method), 59

AngularCosineSquared (class in espressopp.interaction.AngularCosineSquared), 65

AngularHarmonic (class in espressopp.interaction.AngularHarmonic), 66

AngularUniqueCosineSquared (class in espressopp.interaction.AngularUniqueCosineSquared), 67

## C

call() (in module espressopp.pmi), 146

connect() (in module espressopp.tools.vmd), 160

convertTable() (in module espressopp.tools.gromacs), 165

Cosine (class in espressopp.interaction.Cosine), 68

CoulombTruncated (class in espressopp.interaction.CoulombTruncated), 73

CoulombTruncatedUniqueCharge (class in espressopp.interaction.CoulombTruncatedUniqueCharge), 74

create() (in module espressopp.pmi), 146

createCubic() (in module espressopp.tools.lattice), 160

createDenVel() (espressopp.integrator.LBInit.espressopp.integrator.LBInit method), 58

createDiamond() (in module espressopp.tools.lattice), 160

## D

Default() (espressopp.standard\_system.Default.espressopp.standard\_system method), 154

DihedralHarmonic (class in espressopp.interaction.DihedralHarmonic), 76

DihedralHarmonicCos (class in espressopp.interaction.DihedralHarmonicCos), 77

DihedralHarmonicNCos (class in espressopp.interaction.DihedralHarmonicNCos), 77

DihedralHarmonicUniqueCos (class in espressopp.interaction.DihedralHarmonicUniqueCos), 78

displacement (in module espressopp.integrator.MinimizeEnergy), 61

## E

espressopp.\_\_LBInit\_\_Int3D() (in module espressopp.Int3D), 141

espressopp.\_\_Int3D.x() (in module espressopp.Int3D), 141

espressopp.\_\_Int3D.y() (in module espressopp.Int3D), 141

espressopp.\_\_Int3D.z() (in module espressopp.Int3D), 141

espressopp.\_\_Real3D() (in module espressopp.Real3D), 149

espressopp.\_\_Real3D.x() (in module espressopp.Real3D), 149

espressopp.\_\_Real3D.y() (in module espressopp.Real3D), 149

espressopp.\_\_Real3D.z() (in module espressopp.Real3D), 150

espressopp.\_\_RealND() (in module espressopp.RealND), 150

espressopp.analysis.AddressDensity (module), 37

espressopp.analysis.AddressDensity() (in module espressopp.analysis.AddressDensity), 37

espressopp.analysis.AddressDensity.addExclusions() (in module espressopp.analysis.AddressDensity), 37

espressopp.analysis.AddressDensity.compute() (in module espressopp.analysis.AddressDensity), 37

espressopp.analysis.AllParticlePos (module), 27

espressopp.analysis.AllParticlePos.gatherAllPositions() (in module espressopp.analysis.AllParticlePos), 27

espressopp.analysis.AnalysisBase (module), 32

espressopp.analysis.AnalysisBase.compute() (in module espressopp.analysis.AnalysisBase), 32

espressopp.analysis.AnalysisBase.getAverageValue() (in module espressopp.analysis.AnalysisBase), 32

espressopp.analysis.AnalysisBase.getNumberOfMeasurements() (in module espressopp.analysis.AnalysisBase), 32

espressopp.analysis.AnalysisBase.performMeasurement() (in module espressopp.analysis.AnalysisBase), 32

espressopp.analysis.AnalysisBase.reset() (in module espressopp.analysis.AnalysisBase), 32

espressopp.analysis.Autocorrelation (module), 32

espressopp.analysis.Autocorrelation() (in module  
 espressopp.analysis.Autocorrelation), 33  
 espressopp.analysis.Autocorrelation.clear() (in module  
 espressopp.analysis.Autocorrelation), 33  
 espressopp.analysis.Autocorrelation.compute()  
 (in module espressopp.analysis.Autocorrelation), 33  
 espressopp.analysis.Autocorrelation.gather() (in mod-  
 ule espressopp.analysis.Autocorrelation), 33  
 espressopp.analysis.CenterOfMass (module), 37  
 espressopp.analysis.CenterOfMass() (in module espres-  
 sopp.analysis.CenterOfMass), 37  
 espressopp.analysis.CMVelocity (class in espres-  
 sopp.analysis.CMVelocity), 33  
 espressopp.analysis.CMVelocity (module), 33  
 espressopp.analysis.CMVelocity.compute() (in module  
 espressopp.analysis.CMVelocity), 33  
 espressopp.analysis.CMVelocity.reset() (in module  
 espressopp.analysis.CMVelocity), 33  
 espressopp.analysis.ConfigsParticleDecomp (module),  
 33  
 espressopp.analysis.ConfigsParticleDecomp()  
 (in module espressopp.analysis.ConfigsParticleDecomp),  
 33  
 espressopp.analysis.ConfigsParticleDecomp.clear()  
 (in module espressopp.analysis.ConfigsParticleDecomp),  
 33  
 espressopp.analysis.ConfigsParticleDecomp.compute()  
 (in module espressopp.analysis.ConfigsParticleDecomp),  
 34  
 espressopp.analysis.ConfigsParticleDecomp.gather()  
 (in module espressopp.analysis.ConfigsParticleDecomp),  
 34  
 espressopp.analysis.ConfigsParticleDecomp.gatherFromFile()  
 (in module espressopp.analysis.ConfigsParticleDecomp),  
 34  
 espressopp.analysis.Configurations (module), 34  
 espressopp.analysis.Configurations() (in module  
 espressopp.analysis.Configurations), 34  
 espressopp.analysis.Configurations.back() (in module  
 espressopp.analysis.Configurations), 34  
 espressopp.analysis.Configurations.clear() (in module  
 espressopp.analysis.Configurations), 34  
 espressopp.analysis.Configurations.gather() (in module  
 espressopp.analysis.Configurations), 34  
 espressopp.analysis.ConfigurationsExt (module), 34  
 espressopp.analysis.ConfigurationsExt() (in module  
 espressopp.analysis.ConfigurationsExt), 35  
 espressopp.analysis.ConfigurationsExt.back() (in mod-  
 ule espressopp.analysis.ConfigurationsExt),  
 35  
 espressopp.analysis.ConfigurationsExt.clear() (in mod-  
 ule espressopp.analysis.ConfigurationsExt),  
 35  
 espressopp.analysis.ConfigurationsExt.gather()  
 (in module espressopp.analysis.ConfigurationsExt), 35  
 espressopp.analysis.Energy (module), 35  
 espressopp.analysis.EnergyKin() (in module espres-  
 sopp.analysis.Energy), 35  
 espressopp.analysis.EnergyKin.compute() (in module  
 espressopp.analysis.Energy), 36  
 espressopp.analysis.EnergyPot() (in module espres-  
 sopp.analysis.Energy), 35  
 espressopp.analysis.EnergyPot.compute() (in module  
 espressopp.analysis.Energy), 35  
 espressopp.analysis.EnergyTot() (in module espres-  
 sopp.analysis.Energy), 36  
 espressopp.analysis.EnergyTot.compute() (in module  
 espressopp.analysis.Energy), 36  
 espressopp.analysis.IntraChainDistSq (module), 36  
 espressopp.analysis.IntraChainDistSq() (in module  
 espressopp.analysis.IntraChainDistSq), 36  
 espressopp.analysis.IntraChainDistSq.compute()  
 (in module espressopp.analysis.IntraChainDistSq), 36  
 espressopp.analysis.LBOutput (module), 27  
 espressopp.analysis.LBOutputScreen (class in espres-  
 sopp.analysis.LBOutputScreen), 28  
 espressopp.analysis.LBOutputScreen (module), 28  
 espressopp.analysis.LBOutputVzInTime (class in  
 espressopp.analysis.LBOutputVzInTime), 28  
 espressopp.analysis.LBOutputVzInTime (module), 28  
 espressopp.analysis.LBOutputVzOfX (class in espres-  
 sopp.analysis.LBOutputVzOfX), 28  
 espressopp.analysis.LBOutputVzOfX (module), 28  
 espressopp.analysis.MaxPID (module), 37  
 espressopp.analysis.MaxPID() (in module espres-  
 sopp.analysis.MaxPID), 37  
 espressopp.analysis.MeanSquareDispl (module), 36  
 espressopp.analysis.MeanSquareDispl() (in module  
 espressopp.analysis.MeanSquareDispl), 36  
 espressopp.analysis.MeanSquareDispl.computeG2()  
 (in module espressopp.analysis.MeanSquareDispl), 36  
 espressopp.analysis.MeanSquareDispl.computeG3()  
 (in module espressopp.analysis.MeanSquareDispl), 36  
 espressopp.analysis.MeanSquareDispl.strange()  
 (in module espressopp.analysis.MeanSquareDispl), 36  
 espressopp.analysis.MeanSquareInternalDist (module),  
 36  
 espressopp.analysis.MeanSquareInternalDist()  
 (in module espressopp.analysis.MeanSquareInternalDist),  
 36  
 espressopp.analysis.MeanSquareInternalDist.strange()  
 (in module espressopp.analysis.MeanSquareInternalDist),  
 36

- espressopp.analysis.NeighborFluctuation (module), 37
- espressopp.analysis.NeighborFluctuation() (in module espressopp.analysis.NeighborFluctuation), 38
- espressopp.analysis.NPart (module), 38
- espressopp.analysis.NPart() (in module espressopp.analysis.NPart), 38
- espressopp.analysis.Observable (module), 40
- espressopp.analysis.Observable.compute() (in module espressopp.analysis.Observable), 40
- espressopp.analysis.OrderParameter (module), 29
- espressopp.analysis.OrderParameter() (in module espressopp.analysis.OrderParameter), 29
- espressopp.analysis.ParticleRadiusDistribution (module), 29
- espressopp.analysis.ParticleRadiusDistribution() (in module espressopp.analysis.ParticleRadiusDistribution), 29
- espressopp.analysis.PotentialEnergy (module), 38
- espressopp.analysis.PotentialEnergy() (in module espressopp.analysis.PotentialEnergy), 38
- espressopp.analysis.Pressure (module), 38
- espressopp.analysis.Pressure() (in module espressopp.analysis.Pressure), 38
- espressopp.analysis.PressureTensor (module), 29
- espressopp.analysis.PressureTensor() (in module espressopp.analysis.PressureTensor), 30
- espressopp.analysis.PressureTensorLayer (module), 30
- espressopp.analysis.PressureTensorLayer() (in module espressopp.analysis.PressureTensorLayer), 30
- espressopp.analysis.PressureTensorMultiLayer (module), 31
- espressopp.analysis.PressureTensorMultiLayer() (in module espressopp.analysis.PressureTensorMultiLayer), 31
- espressopp.analysis.RadialDistrF (module), 38
- espressopp.analysis.RadialDistrF() (in module espressopp.analysis.RadialDistrF), 38
- espressopp.analysis.RadialDistrF.compute() (in module espressopp.analysis.RadialDistrF), 38
- espressopp.analysis.RDFatomistic (module), 38
- espressopp.analysis.RDFatomistic() (in module espressopp.analysis.RDFatomistic), 39
- espressopp.analysis.RDFatomistic.compute() (in module espressopp.analysis.RDFatomistic), 39
- espressopp.analysis.StaticStructF (module), 39
- espressopp.analysis.StaticStructF() (in module espressopp.analysis.StaticStructF), 39
- espressopp.analysis.StaticStructF.compute() (in module espressopp.analysis.StaticStructF), 39
- espressopp.analysis.StaticStructF.computeSingleChain() (in module espressopp.analysis.StaticStructF), 39
- espressopp.analysis.SystemMonitor (module), 40
- espressopp.analysis.SystemMonitor() (in module espressopp.analysis.SystemMonitor), 40
- espressopp.analysis.SystemMonitor.add\_observable() (in module espressopp.analysis.SystemMonitor), 41
- espressopp.analysis.SystemMonitor.info() (in module espressopp.analysis.SystemMonitor), 41
- espressopp.analysis.SystemMonitorOutputCSV() (in module espressopp.analysis.SystemMonitor), 41
- espressopp.analysis.Temperature (module), 31
- espressopp.analysis.Temperature() (in module espressopp.analysis.Temperature), 31
- espressopp.analysis.Test (module), 32
- espressopp.analysis.Test() (in module espressopp.analysis.Test), 32
- espressopp.analysis.Velocities (module), 41
- espressopp.analysis.Velocities() (in module espressopp.analysis.Velocities), 41
- espressopp.analysis.Velocities.clear() (in module espressopp.analysis.Velocities), 41
- espressopp.analysis.Velocities.gather() (in module espressopp.analysis.Velocities), 41
- espressopp.analysis.VelocityAutocorrelation (module), 41
- espressopp.analysis.VelocityAutocorrelation() (in module espressopp.analysis.VelocityAutocorrelation), 41
- espressopp.analysis.Viscosity (module), 41
- espressopp.analysis.Viscosity() (in module espressopp.analysis.Viscosity), 42
- espressopp.analysis.Viscosity.compute() (in module espressopp.analysis.Viscosity), 42
- espressopp.analysis.Viscosity.gather() (in module espressopp.analysis.Viscosity), 42
- espressopp.analysis.XDensity (module), 40
- espressopp.analysis.XDensity() (in module espressopp.analysis.XDensity), 40
- espressopp.analysis.XDensity.compute() (in module espressopp.analysis.XDensity), 40
- espressopp.analysis.XPressure (module), 40
- espressopp.analysis.XPressure() (in module espressopp.analysis.XPressure), 40
- espressopp.analysis.XPressure.compute() (in module espressopp.analysis.XPressure), 40
- espressopp.analysis.XTemperature (module), 40
- espressopp.analysis.XTemperature() (in module espressopp.analysis.XTemperature), 40
- espressopp.analysis.XTemperature.compute() (in module espressopp.analysis.XTemperature), 40
- espressopp.bc.BC (class in espressopp.bc.BC), 42
- espressopp.bc.BC (module), 42
- espressopp.bc.OrthorhombicBC (module), 43
- espressopp.bc.SlabBC (module), 43
- espressopp.check.System (module), 44
- espressopp.Error() (in module espressopp.Exceptions), 134
- espressopp.esutil.Collectives (module), 44

- espressopp.esutil.GammaVariate (module), 44
- espressopp.esutil.GammaVariate() (in module espressopp.esutil.GammaVariate), 44
- espressopp.esutil.Grid (module), 44
- espressopp.esutil.NormalVariate (module), 44
- espressopp.esutil.NormalVariate() (in module espressopp.esutil.NormalVariate), 44
- espressopp.esutil.RNG (module), 44
- espressopp.esutil.UniformOnSphere (module), 44
- espressopp.Exceptions (module), 134
- espressopp.FixedLocalTupleList (module), 135
- espressopp.FixedLocalTupleList() (in module espressopp.FixedLocalTupleList), 135
- espressopp.FixedLocalTupleList.addTuple() (in module espressopp.FixedLocalTupleList), 135
- espressopp.FixedLocalTupleList.getTuples() (in module espressopp.FixedLocalTupleList), 135
- espressopp.FixedLocalTupleList.size() (in module espressopp.FixedLocalTupleList), 135
- espressopp.FixedPairDistList (module), 135
- espressopp.FixedPairDistList() (in module espressopp.FixedPairDistList), 135
- espressopp.FixedPairDistList.add() (in module espressopp.FixedPairDistList), 135
- espressopp.FixedPairDistList.addPairs() (in module espressopp.FixedPairDistList), 135
- espressopp.FixedPairDistList.getDist() (in module espressopp.FixedPairDistList), 135
- espressopp.FixedPairDistList.getPairs() (in module espressopp.FixedPairDistList), 135
- espressopp.FixedPairDistList.getPairsDist() (in module espressopp.FixedPairDistList), 135
- espressopp.FixedPairDistList.size() (in module espressopp.FixedPairDistList), 136
- espressopp.FixedPairList (module), 136
- espressopp.FixedPairList() (in module espressopp.FixedPairList), 136
- espressopp.FixedPairList.add() (in module espressopp.FixedPairList), 136
- espressopp.FixedPairList.addBonds() (in module espressopp.FixedPairList), 136
- espressopp.FixedPairList.getBonds() (in module espressopp.FixedPairList), 136
- espressopp.FixedPairList.getLongtimeMaxBond() (in module espressopp.FixedPairList), 136
- espressopp.FixedPairList.remove() (in module espressopp.FixedPairList), 136
- espressopp.FixedPairList.resetLongtimeMaxBond() (in module espressopp.FixedPairList), 136
- espressopp.FixedPairList.size() (in module espressopp.FixedPairList), 136
- espressopp.FixedPairListAddress (module), 136
- espressopp.FixedPairListAddress() (in module espressopp.FixedPairListAddress), 136
- espressopp.FixedPairListAddress.add() (in module espressopp.FixedPairListAddress), 136
- espressopp.FixedPairListAddress.addBonds() (in module espressopp.FixedPairListAddress), 137
- espressopp.FixedPairListAddress.getBonds() (in module espressopp.FixedPairListAddress), 137
- espressopp.FixedPairListAddress.remove() (in module espressopp.FixedPairListAddress), 137
- espressopp.FixedQuadrupleAngleList (module), 137
- espressopp.FixedQuadrupleAngleList() (in module espressopp.FixedQuadrupleAngleList), 137
- espressopp.FixedQuadrupleAngleList.add() (in module espressopp.FixedQuadrupleAngleList), 137
- espressopp.FixedQuadrupleAngleList.addQuadruples() (in module espressopp.FixedQuadrupleAngleList), 137
- espressopp.FixedQuadrupleAngleList.getAngle() (in module espressopp.FixedQuadrupleAngleList), 137
- espressopp.FixedQuadrupleAngleList.getQuadruples() (in module espressopp.FixedQuadrupleAngleList), 137
- espressopp.FixedQuadrupleAngleList.getQuadruplesAngles() (in module espressopp.FixedQuadrupleAngleList), 137
- espressopp.FixedQuadrupleAngleList.size() (in module espressopp.FixedQuadrupleAngleList), 137
- espressopp.FixedQuadrupleList (module), 137
- espressopp.FixedQuadrupleList() (in module espressopp.FixedQuadrupleList), 138
- espressopp.FixedQuadrupleList.add() (in module espressopp.FixedQuadrupleList), 138
- espressopp.FixedQuadrupleList.addQuadruples() (in module espressopp.FixedQuadrupleList), 138
- espressopp.FixedQuadrupleList.getQuadruples() (in module espressopp.FixedQuadrupleList), 138
- espressopp.FixedQuadrupleList.remove() (in module espressopp.FixedQuadrupleList), 138
- espressopp.FixedQuadrupleList.size() (in module espressopp.FixedQuadrupleList), 138
- espressopp.FixedQuadrupleListAddress (module), 138
- espressopp.FixedQuadrupleListAddress() (in module espressopp.FixedQuadrupleListAddress), 138
- espressopp.FixedQuadrupleListAddress.add() (in module espressopp.FixedQuadrupleListAddress), 138
- espressopp.FixedQuadrupleListAddress.addQuadruples() (in module espressopp.FixedQuadrupleListAddress), 138
- espressopp.FixedQuadrupleListAddress.getQuadruples() (in module espressopp.FixedQuadrupleListAddress), 138
- espressopp.FixedQuadrupleListAddress.size() (in module espressopp.FixedQuadrupleListAddress), 138
- espressopp.FixedSingleList (module), 138
- espressopp.FixedSingleList() (in module espressopp.FixedSingleList), 139
- espressopp.FixedSingleList.add() (in module espressopp.FixedSingleList), 139

- espressopp.FixedSingleList.addSingles() (in module espressopp.FixedSingleList), 139
- espressopp.FixedSingleList.getSingles() (in module espressopp.FixedSingleList), 139
- espressopp.FixedSingleList.size() (in module espressopp.FixedSingleList), 139
- espressopp.FixedTripleAngleList (module), 139
- espressopp.FixedTripleAngleList() (in module espressopp.FixedTripleAngleList), 139
- espressopp.FixedTripleAngleList.add() (in module espressopp.FixedTripleAngleList), 139
- espressopp.FixedTripleAngleList.addTriples() (in module espressopp.FixedTripleAngleList), 139
- espressopp.FixedTripleAngleList.getAngle() (in module espressopp.FixedTripleAngleList), 139
- espressopp.FixedTripleAngleList.getTriples() (in module espressopp.FixedTripleAngleList), 139
- espressopp.FixedTripleAngleList.getTriplesAngles() (in module espressopp.FixedTripleAngleList), 139
- espressopp.FixedTripleAngleList.size() (in module espressopp.FixedTripleAngleList), 139
- espressopp.FixedTripleList (module), 139
- espressopp.FixedTripleList() (in module espressopp.FixedTripleList), 140
- espressopp.FixedTripleList.add() (in module espressopp.FixedTripleList), 140
- espressopp.FixedTripleList.addTriples() (in module espressopp.FixedTripleList), 140
- espressopp.FixedTripleList.getTriples() (in module espressopp.FixedTripleList), 140
- espressopp.FixedTripleList.remove() (in module espressopp.FixedTripleList), 140
- espressopp.FixedTripleList.size() (in module espressopp.FixedTripleList), 140
- espressopp.FixedTripleListAdress (module), 140
- espressopp.FixedTripleListAdress() (in module espressopp.FixedTripleListAdress), 140
- espressopp.FixedTripleListAdress.add() (in module espressopp.FixedTripleListAdress), 140
- espressopp.FixedTripleListAdress.addTriples() (in module espressopp.FixedTripleListAdress), 140
- espressopp.FixedTripleListAdress.remove() (in module espressopp.FixedTripleListAdress), 140
- espressopp.FixedTupList (module), 140
- espressopp.FixedTupList() (in module espressopp.FixedTupList), 140
- espressopp.FixedTupList.size() (in module espressopp.FixedTupList), 140
- espressopp.FixedTupListAdress (module), 141
- espressopp.FixedTupListAdress() (in module espressopp.FixedTupListAdress), 141
- espressopp.FixedTupListAdress.addTuples() (in module espressopp.FixedTupListAdress), 141
- espressopp.Int3D (module), 141
- espressopp.integrator.Address (class in espressopp.integrator.Address), 45
- espressopp.integrator.Address (module), 44
- espressopp.integrator.AssociationReaction (module), 45
- espressopp.integrator.BerendsenBarostat (module), 45
- espressopp.integrator.BerendsenBarostat() (in module espressopp.integrator.BerendsenBarostat), 46
- espressopp.integrator.BerendsenBarostatAnisotropic (module), 46
- espressopp.integrator.BerendsenBarostatAnisotropic() (in module espressopp.integrator.BerendsenBarostatAnisotropic), 47
- espressopp.integrator.BerendsenThermostat (module), 47
- espressopp.integrator.BerendsenThermostat() (in module espressopp.integrator.BerendsenThermostat), 48
- espressopp.integrator.CapForce (module), 48
- espressopp.integrator.CapForce() (in module espressopp.integrator.CapForce), 49
- espressopp.integrator.DPDThermostat (module), 49
- espressopp.integrator.DPDThermostat() (in module espressopp.integrator.DPDThermostat), 49
- espressopp.integrator.EmptyExtension (module), 49
- espressopp.integrator.EmptyExtension() (in module espressopp.integrator.EmptyExtension), 49
- espressopp.integrator.ExtAnalyze (module), 49
- espressopp.integrator.ExtAnalyze() (in module espressopp.integrator.ExtAnalyze), 49
- espressopp.integrator.Extension (module), 49
- espressopp.integrator.Extension.connect() (in module espressopp.integrator.Extension), 50
- espressopp.integrator.Extension.disconnect() (in module espressopp.integrator.Extension), 50
- espressopp.integrator.ExtForce (module), 50
- espressopp.integrator.ExtForce() (in module espressopp.integrator.ExtForce), 50
- espressopp.integrator.FixPositions (module), 50
- espressopp.integrator.FixPositions() (in module espressopp.integrator.FixPositions), 50
- espressopp.integrator.FreeEnergyCompensation (module), 50
- espressopp.integrator.FreeEnergyCompensation() (in module espressopp.integrator.FreeEnergyCompensation), 50
- espressopp.integrator.FreeEnergyCompensation.addForce() (in module espressopp.integrator.FreeEnergyCompensation), 50
- espressopp.integrator.FreeEnergyCompensation.computeCompEnergy() (in module espressopp.integrator.FreeEnergyCompensation), 51
- espressopp.integrator.GeneralizedLangevinThermostat (module), 51



[esspressopp.integrator.GeneralizedLangevinThermostat\(\)](#) (in module [esspressopp.integrator.GeneralizedLangevinThermostat](#)), 51  
[esspressopp.integrator.GeneralizedLangevinThermostat.addCoeffs\(\)](#) (in module [esspressopp.integrator.GeneralizedLangevinThermostat](#)), 51  
[esspressopp.integrator.GeneralizedLangevinThermostat.addExclusions\(\)](#) (in module [esspressopp.integrator.GeneralizedLangevinThermostat](#)), 53  
[esspressopp.integrator.GeneralizedLangevinThermostat1D](#) (module), 53  
[esspressopp.integrator.GeneralizedLangevinThermostat1D\(\)](#) (in module [esspressopp.integrator.GeneralizedLangevinThermostat1D](#)), 53  
[esspressopp.integrator.GeneralizedLangevinThermostatHybrid](#) (module), 53  
[esspressopp.integrator.GeneralizedLangevinThermostatOnGroup](#) (module), 54  
[esspressopp.integrator.GeneralizedLangevinThermostatOnGroup\(\)](#) (in module [esspressopp.integrator.GeneralizedLangevinThermostatOnGroup](#)), 54  
[esspressopp.integrator.GeneralizedLangevinThermostatOnRadius](#) (module), 54  
[esspressopp.integrator.GeneralizedLangevinThermostatOnRadius\(\)](#) (in module [esspressopp.integrator.GeneralizedLangevinThermostatOnRadius](#)), 55  
[esspressopp.integrator.GeneralizedLangevinThermostatOnRadius.addExclusions\(\)](#) (in module [esspressopp.integrator.GeneralizedLangevinThermostatOnRadius](#)), 55  
[esspressopp.integrator.LatticeBoltzmann](#) (class in [esspressopp.integrator.LatticeBoltzmann](#)), 55  
[esspressopp.integrator.LatticeBoltzmann](#) (module), 55  
[esspressopp.integrator.LBInit](#) (class in [esspressopp.integrator.LBInit](#)), 58  
[esspressopp.integrator.LBInit](#) (module), 58  
[esspressopp.integrator.LBInitConstForce](#) (module), 59  
[esspressopp.integrator.LBInitPeriodicForce](#) (module), 60  
[esspressopp.integrator.LBInitPopUniform](#) (module), 59  
[esspressopp.integrator.LBInitPopWave](#) (module), 59  
[esspressopp.integrator.MDIntegrator](#) (module), 60  
[esspressopp.integrator.MDIntegrator.addExtension\(\)](#) (in module [esspressopp.integrator.MDIntegrator](#)), 60  
[esspressopp.integrator.MDIntegrator.getExtension\(\)](#) (in module [esspressopp.integrator.MDIntegrator](#)), 60  
[esspressopp.integrator.MDIntegrator.getNumberOfExtensions\(\)](#) (in module [esspressopp.integrator.MDIntegrator](#)), 60  
[esspressopp.integrator.MDIntegrator.run\(\)](#) (in module [esspressopp.integrator.MDIntegrator](#)), 60  
[esspressopp.integrator.MinimizeEnergy](#) (module), 60  
[esspressopp.integrator.MinimizeEnergy\(\)](#) (in module [esspressopp.integrator.MinimizeEnergy](#)), 61  
[esspressopp.integrator.MinimizeEnergy.run\(\)](#) (in module [esspressopp.integrator.MinimizeEnergy](#)), 61  
[esspressopp.integrator.OnTheFlyFEC](#) (module), 61  
[esspressopp.integrator.OnTheFlyFEC\(\)](#) (in module [esspressopp.integrator.OnTheFlyFEC](#)), 62  
[esspressopp.integrator.OnTheFlyFEC.getBins\(\)](#) (in module [esspressopp.integrator.OnTheFlyFEC](#)), 62  
[esspressopp.integrator.OnTheFlyFEC.getGap\(\)](#) (in module [esspressopp.integrator.OnTheFlyFEC](#)), 62  
[esspressopp.integrator.OnTheFlyFEC.getSteps\(\)](#) (in module [esspressopp.integrator.OnTheFlyFEC](#)), 62  
[esspressopp.integrator.OnTheFlyFEC.makeArrays\(\)](#) (in module [esspressopp.integrator.OnTheFlyFEC](#)), 62  
[esspressopp.integrator.OnTheFlyFEC.resetCounter\(\)](#) (in module [esspressopp.integrator.OnTheFlyFEC](#)), 62  
[esspressopp.integrator.OnTheFlyFEC.writeFEC\(\)](#) (in module [esspressopp.integrator.OnTheFlyFEC](#)), 62  
[esspressopp.integrator.Rattle](#) (module), 62  
[esspressopp.integrator.Settle](#) (module), 63  
[esspressopp.integrator.Settle\(\)](#) (in module [esspressopp.integrator.Settle](#)), 63  
[esspressopp.integrator.Settle.addMolecules\(\)](#) (in module [esspressopp.integrator.Settle](#)), 63  
[esspressopp.integrator.StochasticVelocityRescaling](#) (module), 63  
[esspressopp.integrator.StochasticVelocityRescaling\(\)](#) (in module [esspressopp.integrator.StochasticVelocityRescaling](#)), 63  
[esspressopp.integrator.TDforce](#) (module), 63  
[esspressopp.integrator.TDforce\(\)](#) (in module [esspressopp.integrator.TDforce](#)), 64  
[esspressopp.integrator.TDforce.addForce\(\)](#) (in module [esspressopp.integrator.TDforce](#)), 64  
[esspressopp.integrator.VelocityVerlet](#) (module), 64  
[esspressopp.integrator.VelocityVerlet\(\)](#) (in module [esspressopp.integrator.VelocityVerlet](#)), 64  
[esspressopp.integrator.VelocityVerletOnGroup](#) (module), 64

[espressopp.integrator.VelocityVerletOnGroup\(\)](#)  
 (in module [espressopp.integrator.VelocityVerletOnGroup](#)),  
[64](#)

[espressopp.integrator.VelocityVerletOnRadius](#) (module), [64](#)

[espressopp.integrator.VelocityVerletOnRadius\(\)](#)  
 (in module [espressopp.integrator.VelocityVerletOnRadius](#)),  
[65](#)

[espressopp.interaction.AngularCosineSquared](#) (module), [65](#)

[espressopp.interaction.AngularCosineSquared\(\)](#)  
 (in module [espressopp.interaction.AngularCosineSquared](#)),  
[65](#)

[espressopp.interaction.AngularHarmonic](#) (module), [65](#)

[espressopp.interaction.AngularHarmonic\(\)](#) (in module [espressopp.interaction.AngularHarmonic](#)),  
[65](#)

[espressopp.interaction.AngularPotential](#) (module), [66](#)

[espressopp.interaction.AngularPotential.computeEnergy\(\)](#)  
 (in module [espressopp.interaction.AngularPotential](#)), [66](#)

[espressopp.interaction.AngularPotential.computeForce\(\)](#)  
 (in module [espressopp.interaction.AngularPotential](#)), [66](#)

[espressopp.interaction.AngularUniqueCosineSquared](#)  
 (module), [66](#)

[espressopp.interaction.AngularUniqueCosineSquared\(\)](#)  
 (in module [espressopp.interaction.AngularUniqueCosineSquared](#)),  
[66](#)

[espressopp.interaction.AngularUniqueHarmonic](#) (module), [67](#)

[espressopp.interaction.AngularUniqueHarmonic\(\)](#)  
 (in module [espressopp.interaction.AngularUniqueHarmonic](#)),  
[67](#)

[espressopp.interaction.AngularUniquePotential](#) (module), [67](#)

[espressopp.interaction.AngularUniquePotential.computeEnergy\(\)](#)  
 (in module [espressopp.interaction.AngularUniquePotential](#)),  
[67](#)

[espressopp.interaction.AngularUniquePotential.computeForce\(\)](#)  
 (in module [espressopp.interaction.AngularUniquePotential](#)),  
[67](#)

[espressopp.interaction.CellListCoulombKSpaceEwald\(\)](#)  
 (in module [espressopp.interaction.CoulombKSpaceEwald](#)),  
[69](#)

[espressopp.interaction.CellListCoulombKSpaceEwald.getPotential\(\)](#)  
 (in module [espressopp.interaction.CoulombKSpaceEwald](#)),  
[69](#)

[espressopp.interaction.CellListCoulombKSpaceEwald.getPotential\(\)](#)  
 (in module [espressopp.interaction.CoulombKSpaceEwald](#)),  
[69](#)

(in module [espressopp.interaction.CoulombKSpaceEwald](#)),  
[69](#)

[espressopp.interaction.CellListCoulombKSpaceP3M\(\)](#)  
 (in module [espressopp.interaction.CoulombKSpaceP3M](#)),  
[70](#)

[espressopp.interaction.CellListCoulombKSpaceP3M.getPotential\(\)](#)  
 (in module [espressopp.interaction.CoulombKSpaceP3M](#)),  
[70](#)

[espressopp.interaction.CellListCoulombTruncatedUniqueCharge\(\)](#)  
 (in module [espressopp.interaction.CoulombTruncatedUniqueCharge](#)),  
[74](#)

[espressopp.interaction.CellListCoulombTruncatedUniqueCharge.setPotential\(\)](#)  
 (in module [espressopp.interaction.CoulombTruncatedUniqueCharge](#)),  
[74](#)

[espressopp.interaction.CellListLennardJones\(\)](#)  
 (in module [espressopp.interaction.LennardJones](#)), [87](#)

[espressopp.interaction.CellListLennardJones.setPotential\(\)](#)  
 (in module [espressopp.interaction.LennardJones](#)), [87](#)

[espressopp.interaction.CellListLennardJonesAutoBonds\(\)](#)  
 (in module [espressopp.interaction.LennardJonesAutoBonds](#)),  
[89](#)

[espressopp.interaction.CellListLennardJonesAutoBonds.setPotential\(\)](#)  
 (in module [espressopp.interaction.LennardJonesAutoBonds](#)),  
[90](#)

[espressopp.interaction.CellListLennardJonesCapped\(\)](#)  
 (in module [espressopp.interaction.LennardJonesCapped](#)),  
[92](#)

[espressopp.interaction.CellListLennardJonesCapped.getPotential\(\)](#)  
 (in module [espressopp.interaction.LennardJonesCapped](#)),  
[92](#)

[espressopp.interaction.CellListLennardJonesCapped.setPotential\(\)](#)  
 (in module [espressopp.interaction.LennardJonesCapped](#)),  
[92](#)

[espressopp.interaction.CellListLennardJonesEnergyCapped\(\)](#)  
 (in module [espressopp.interaction.LennardJonesEnergyCapped](#)),  
[95](#)

[espressopp.interaction.CellListLennardJonesEnergyCapped.getPotential\(\)](#)  
 (in module [espressopp.interaction.LennardJonesEnergyCapped](#)),  
[95](#)

[espressopp.interaction.CellListLennardJonesEnergyCapped.setPotential\(\)](#)  
 (in module [espressopp.interaction.LennardJonesEnergyCapped](#)),  
[95](#)

[espressopp.interaction.CellListLennardJonesExpand\(\)](#)  
 (in module [espressopp.interaction.LennardJonesExpand](#)),  
[95](#)

(in module espressopp.interaction.LennardJonesExpand), 97

espressopp.interaction.CellListLennardJonesExpand.setPotential() (in module espressopp.interaction.LennardJonesExpand), 97

espressopp.interaction.CellListLennardJonesGeneric() (in module espressopp.interaction.LennardJonesGeneric), 100

espressopp.interaction.CellListLennardJonesGeneric.setPotential() (in module espressopp.interaction.LennardJonesGeneric), 100

espressopp.interaction.CellListLennardJonesGromacs() (in module espressopp.interaction.LennardJonesGromacs), 101

espressopp.interaction.CellListLennardJonesGromacs.setPotential() (in module espressopp.interaction.LennardJonesGromacs), 101

espressopp.interaction.CellListLJcos() (in module espressopp.interaction.LJcos), 105

espressopp.interaction.CellListLJcos.setPotential() (in module espressopp.interaction.LJcos), 105

espressopp.interaction.CellListMorse() (in module espressopp.interaction.Morse), 108

espressopp.interaction.CellListMorse.setPotential() (in module espressopp.interaction.Morse), 108

espressopp.interaction.CellListReactionFieldGeneralized() (in module espressopp.interaction.ReactionFieldGeneralized), 112

espressopp.interaction.CellListReactionFieldGeneralized.setPotential() (in module espressopp.interaction.ReactionFieldGeneralized), 112

espressopp.interaction.CellListSoftCosine() (in module espressopp.interaction.SoftCosine), 115

espressopp.interaction.CellListSoftCosine.setPotential() (in module espressopp.interaction.SoftCosine), 115

espressopp.interaction.CellListStillingerWeberPairTerm() (in module espressopp.interaction.StillingerWeberPairTerm), 117

espressopp.interaction.CellListStillingerWeberPairTerm.setPotential() (in module espressopp.interaction.StillingerWeberPairTerm), 117

espressopp.interaction.CellListStillingerWeberPairTermCapped() (in module espressopp.interaction.StillingerWeberPairTermCapped), 120

espressopp.interaction.CellListStillingerWeberPairTermCapped.setPotential() (in module espressopp.interaction.StillingerWeberPairTermCapped), 120

sopp.interaction.StillingerWeberPairTermCapped), 120

espressopp.interaction.CellListTabulated() (in module espressopp.interaction.Tabulated), 123

espressopp.interaction.CellListTabulated.setPotential() (in module espressopp.interaction.Tabulated), 123

espressopp.interaction.CellListTersoffPairTerm() (in module espressopp.interaction.TersoffPairTerm), 126

espressopp.interaction.CellListTersoffPairTerm.setPotential() (in module espressopp.interaction.TersoffPairTerm), 127

espressopp.interaction.CellListZero() (in module espressopp.interaction.Zero), 131

espressopp.interaction.CellListZero.setPotential() (in module espressopp.interaction.Zero), 131

espressopp.interaction.ConstrainCOM (module), 74

espressopp.interaction.ConstrainCOM() (in module espressopp.interaction.ConstrainCOM), 75

espressopp.interaction.ConstrainRG (module), 75

espressopp.interaction.ConstrainRG() (in module espressopp.interaction.ConstrainRG), 75

espressopp.interaction.Cosine (module), 67

espressopp.interaction.Cosine() (in module espressopp.interaction.Cosine), 67

espressopp.interaction.CoulombKSpaceEwald (module), 68

espressopp.interaction.CoulombKSpaceEwald() (in module espressopp.interaction.CoulombKSpaceEwald), 69

espressopp.interaction.CoulombKSpaceP3M (module), 69

espressopp.interaction.CoulombKSpaceP3M() (in module espressopp.interaction.CoulombKSpaceP3M), 70

espressopp.interaction.CoulombRSpace (module), 70

espressopp.interaction.CoulombRSpace() (in module espressopp.interaction.CoulombRSpace), 71

espressopp.interaction.CoulombTruncated (module), 72

espressopp.interaction.CoulombTruncatedUniqueCharge (module), 73

espressopp.interaction.CoulombTruncatedUniqueCharge() (in module espressopp.interaction.CoulombTruncatedUniqueCharge), 73

espressopp.interaction.DihedralHarmonic (module), 75

espressopp.interaction.DihedralHarmonic() (in module espressopp.interaction.DihedralHarmonic), 75

espressopp.interaction.DihedralHarmonicCos (module), 76

espressopp.interaction.DihedralHarmonicCos() (in module espressopp.interaction.DihedralHarmonicCos), 76



76  
 espressopp.interaction.DihedralHarmonicNCos (module), 77  
 espressopp.interaction.DihedralHarmonicNCos() (in module espressopp.interaction.DihedralHarmonicNCos), 77  
 espressopp.interaction.DihedralHarmonicUniqueCos (module), 77  
 espressopp.interaction.DihedralHarmonicUniqueCos() (in module espressopp.interaction.DihedralHarmonicUniqueCos), 78  
 espressopp.interaction.DihedralPotential (module), 78  
 espressopp.interaction.DihedralPotential.computeEnergy() (in module espressopp.interaction.DihedralPotential), 78  
 espressopp.interaction.DihedralPotential.computeForce() (in module espressopp.interaction.DihedralPotential), 78  
 espressopp.interaction.DihedralRB (module), 78  
 espressopp.interaction.DihedralRB() (in module espressopp.interaction.DihedralRB), 78  
 espressopp.interaction.DihedralUniquePotential (module), 79  
 espressopp.interaction.DihedralUniquePotential.computeEnergy() (in module espressopp.interaction.DihedralUniquePotential), 79  
 espressopp.interaction.DihedralUniquePotential.computeForce() (in module espressopp.interaction.DihedralUniquePotential), 79  
 espressopp.interaction.FENE (module), 79  
 espressopp.interaction.FENE() (in module espressopp.interaction.FENE), 79  
 espressopp.interaction.FENECapped (module), 80  
 espressopp.interaction.FENECapped() (in module espressopp.interaction.FENECapped), 80  
 espressopp.interaction.FixedLocalTupleListConstrainCOM() (in module espressopp.interaction.ConstrainCOM), 75  
 espressopp.interaction.FixedLocalTupleListConstrainCOM.getPotential() (in module espressopp.interaction.ConstrainCOM), 75  
 espressopp.interaction.FixedLocalTupleListConstrainCOM.setCom() (in module espressopp.interaction.ConstrainCOM), 75  
 espressopp.interaction.FixedLocalTupleListConstrainRG() (in module espressopp.interaction.ConstrainRG), 75  
 espressopp.interaction.FixedLocalTupleListConstrainRG.getPotential() (in module espressopp.interaction.ConstrainRG), 75  
 espressopp.interaction.FixedLocalTupleListConstrainRG.setRG() (in module espressopp.interaction.ConstrainRG), 75  
 espressopp.interaction.FixedPairDistListHarmonicUnique() (in module espressopp.interaction.HarmonicUnique), 83  
 espressopp.interaction.FixedPairDistListHarmonicUnique.getFixedPairList() (in module espressopp.interaction.HarmonicUnique), 83  
 espressopp.interaction.FixedPairDistListHarmonicUnique.setFixedPairList() (in module espressopp.interaction.HarmonicUnique), 83  
 espressopp.interaction.FixedPairDistListHarmonicUnique.setPotential() (in module espressopp.interaction.HarmonicUnique), 84  
 espressopp.interaction.FixedPairListCoulombTruncatedUniqueCharge() (in module espressopp.interaction.CoulombTruncatedUniqueCharge), 74  
 espressopp.interaction.FixedPairListCoulombTruncatedUniqueCharge.setPotential() (in module espressopp.interaction.CoulombTruncatedUniqueCharge), 74  
 espressopp.interaction.FixedPairListFENE() (in module espressopp.interaction.FENE), 80  
 espressopp.interaction.FixedPairListFENE.getFixedPairList() (in module espressopp.interaction.FENE), 80  
 espressopp.interaction.FixedPairListFENE.getPotential() (in module espressopp.interaction.FENE), 80  
 espressopp.interaction.FixedPairListFENE.setFixedPairList() (in module espressopp.interaction.FENE), 80  
 espressopp.interaction.FixedPairListFENE.setPotential() (in module espressopp.interaction.FENE), 80  
 espressopp.interaction.FixedPairListFENECapped() (in module espressopp.interaction.FENECapped), 81  
 espressopp.interaction.FixedPairListFENECapped.getFixedPairList() (in module espressopp.interaction.FENECapped), 81  
 espressopp.interaction.FixedPairListFENECapped.getPotential() (in module espressopp.interaction.FENECapped), 81  
 espressopp.interaction.FixedPairListFENECapped.setFixedPairList() (in module espressopp.interaction.FENECapped), 81  
 espressopp.interaction.FixedPairListFENECapped.setPotential() (in module espressopp.interaction.FENECapped), 81  
 espressopp.interaction.FixedPairListHarmonic() (in module espressopp.interaction.Harmonic), 82  
 espressopp.interaction.FixedPairListHarmonic.getFixedPairList() (in module espressopp.interaction.Harmonic), 82  
 espressopp.interaction.FixedPairListHarmonic.setFixedPairList() (in module espressopp.interaction.Harmonic), 82  
 espressopp.interaction.FixedPairListHarmonic.setPotential() (in module espressopp.interaction.Harmonic), 82  
 espressopp.interaction.FixedPairListLennardJones() (in module espressopp.interaction.LennardJones), 82

sopp.interaction.LennardJones), 87  
 espressopp.interaction.FixedPairListLennardJones.getFixedPairList(sopp.interaction.LennardJonesGeneric),  
 (in module espressopp.interaction.FixedPairListLennardJones), 87  
 espressopp.interaction.FixedPairListLennardJonesGeneric.getPotential()  
 (in module espressopp.interaction.LennardJonesGeneric), 100  
 espressopp.interaction.FixedPairListLennardJones.setFixedPairList(sopp.interaction.FixedPairListLennardJonesGeneric.setFixedPairList),  
 (in module espressopp.interaction.LennardJonesGeneric), 100  
 espressopp.interaction.FixedPairListLennardJones.setPotential(sopp.interaction.FixedPairListLennardJonesGeneric.setPotential),  
 (in module espressopp.interaction.LennardJonesGeneric), 100  
 espressopp.interaction.FixedPairListLennardJonesAutoBonds(sopp.interaction.FixedPairListLennardJonesGromacs),  
 (in module espressopp.interaction.LennardJonesGromacs), 101  
 espressopp.interaction.FixedPairListLennardJonesAutoBonds.setPotential(sopp.interaction.FixedPairListLennardJonesGromacs.setPotential),  
 (in module espressopp.interaction.LennardJonesGromacs), 101  
 espressopp.interaction.FixedPairListLennardJonesCapped(sopp.interaction.FixedPairListLJcos), 105  
 (in module espressopp.interaction.LJcos), 105  
 espressopp.interaction.FixedPairListLennardJonesCapped.setPotential(sopp.interaction.FixedPairListLJcos.setFixedPairList),  
 (in module espressopp.interaction.LJcos), 105  
 espressopp.interaction.FixedPairListLennardJonesCapped.setPotential(sopp.interaction.FixedPairListLJcos.setPotential),  
 (in module espressopp.interaction.LJcos), 105  
 espressopp.interaction.FixedPairListLennardJonesEnergyCapped(sopp.interaction.FixedPairListMirrorLennardJones),  
 (in module espressopp.interaction.MirrorLennardJones), 106  
 espressopp.interaction.FixedPairListLennardJonesEnergyCapped.getPotential(sopp.interaction.FixedPairListMirrorLennardJones.getFixedPairList),  
 (in module espressopp.interaction.MirrorLennardJones), 106  
 espressopp.interaction.FixedPairListLennardJonesEnergyCapped.setPotential(sopp.interaction.FixedPairListMirrorLennardJones.setPotential),  
 (in module espressopp.interaction.MirrorLennardJones), 106  
 espressopp.interaction.FixedPairListLennardJonesExpand(sopp.interaction.FixedPairListMorse), 108  
 (in module espressopp.interaction.Morse), 108  
 espressopp.interaction.FixedPairListLennardJonesExpand.setPotential(sopp.interaction.FixedPairListMorse.setPotential),  
 (in module espressopp.interaction.Morse), 108  
 espressopp.interaction.FixedPairListLennardJonesGeneric(sopp.interaction.FixedPairListMorse), 108  
 (in module espressopp.interaction.Morse), 108  
 espressopp.interaction.FixedPairListLennardJonesGeneric.getFixedPairList(sopp.interaction.FixedPairListMorse), 108  
 (in module espressopp.interaction.Morse), 108

---

<b>Index</b>	<b>191</b>
--------------	------------



espressopp.interaction.HarmonicUnique), 83  
 espressopp.interaction.Interaction (module), 84  
 espressopp.interaction.Interaction.bondType() (in module espressopp.interaction.Interaction), 84  
 espressopp.interaction.Interaction.computeEnergy() (in module espressopp.interaction.Interaction), 84  
 espressopp.interaction.Interaction.computeEnergyAA() (in module espressopp.interaction.Interaction), 84  
 espressopp.interaction.Interaction.computeEnergyCG() (in module espressopp.interaction.Interaction), 84  
 espressopp.interaction.Interaction.computeEnergyDeriv() (in module espressopp.interaction.Interaction), 84  
 espressopp.interaction.Interaction.computeVirial() (in module espressopp.interaction.Interaction), 84  
 espressopp.interaction.LennardJones (module), 84  
 espressopp.interaction.LennardJones() (in module espressopp.interaction.LennardJones), 84  
 espressopp.interaction.LennardJones93Wall (module), 87  
 espressopp.interaction.LennardJones93Wall() (in module espressopp.interaction.LennardJones93Wall), 88  
 espressopp.interaction.LennardJones93Wall.getParams() (in module espressopp.interaction.LennardJones93Wall), 88  
 espressopp.interaction.LennardJones93Wall.setParams() (in module espressopp.interaction.LennardJones93Wall), 88  
 espressopp.interaction.LennardJonesAutoBonds (module), 88  
 espressopp.interaction.LennardJonesAutoBonds() (in module espressopp.interaction.LennardJonesAutoBonds), 88  
 espressopp.interaction.LennardJonesCapped (module), 90  
 espressopp.interaction.LennardJonesCapped() (in module espressopp.interaction.LennardJonesCapped), 90  
 espressopp.interaction.LennardJonesEnergyCapped (module), 93  
 espressopp.interaction.LennardJonesEnergyCapped() (in module espressopp.interaction.LennardJonesEnergyCapped), 93  
 espressopp.interaction.LennardJonesExpand (module), 96  
 espressopp.interaction.LennardJonesExpand() (in module espressopp.interaction.LennardJonesExpand), 96  
 espressopp.interaction.LennardJonesGeneric (module), 97  
 espressopp.interaction.LennardJonesGeneric() (in module espressopp.interaction.LennardJonesGeneric), 97  
 espressopp.interaction.LennardJonesGromacs (module), 100  
 espressopp.interaction.LennardJonesGromacs() (in module espressopp.interaction.LennardJonesGromacs), 101  
 espressopp.interaction.LennardJonesSoftcoreTI (module), 102  
 espressopp.interaction.LJcos (module), 104  
 espressopp.interaction.LJcos() (in module espressopp.interaction.LJcos), 104  
 espressopp.interaction.MirrorLennardJones (module), 105  
 espressopp.interaction.MirrorLennardJones() (in module espressopp.interaction.MirrorLennardJones), 106  
 espressopp.interaction.Morse (module), 106  
 espressopp.interaction.Morse() (in module espressopp.interaction.Morse), 106  
 espressopp.interaction.OPLS (module), 108  
 espressopp.interaction.OPLS() (in module espressopp.interaction.OPLS), 108  
 espressopp.interaction.Potential (module), 109  
 espressopp.interaction.Potential.computeEnergy() (in module espressopp.interaction.Potential), 109  
 espressopp.interaction.Potential.computeForce() (in module espressopp.interaction.Potential), 109  
 espressopp.interaction.PotentialUniqueDist (module), 109  
 espressopp.interaction.PotentialUniqueDist.computeEnergy() (in module espressopp.interaction.PotentialUniqueDist), 109  
 espressopp.interaction.PotentialUniqueDist.computeForce() (in module espressopp.interaction.PotentialUniqueDist), 109  
 espressopp.interaction.PotentialVSpherePair (module), 109  
 espressopp.interaction.PotentialVSpherePair.computeEnergy() (in module espressopp.interaction.PotentialVSpherePair), 109  
 espressopp.interaction.PotentialVSpherePair.computeForce() (in module espressopp.interaction.PotentialVSpherePair), 109



- espressopp.interaction.Quartic (module), 109
- espressopp.interaction.Quartic() (in module espressopp.interaction.Quartic), 109
- espressopp.interaction.ReactionFieldGeneralized (module), 110
- espressopp.interaction.ReactionFieldGeneralized() (in module espressopp.interaction.ReactionFieldGeneralized), 110
- espressopp.interaction.ReactionFieldGeneralizedTI (module), 112
- espressopp.interaction.SelfVSPHERE() (in module espressopp.interaction.VSPHERESelf), 129
- espressopp.interaction.SelfVSPHERE.getPotential() (in module espressopp.interaction.VSPHERESelf), 129
- espressopp.interaction.SelfVSPHERE.setPotential() (in module espressopp.interaction.VSPHERESelf), 129
- espressopp.interaction.SingleParticleHarmonicTrap() (in module espressopp.interaction.HarmonicTrap), 83
- espressopp.interaction.SingleParticleHarmonicTrap.setPotential() (in module espressopp.interaction.HarmonicTrap), 83
- espressopp.interaction.SingleParticleLennardJones93Wall() (in module espressopp.interaction.LennardJones93Wall), 88
- espressopp.interaction.SingleParticleLennardJones93Wall.setPotential() (in module espressopp.interaction.LennardJones93Wall), 88
- espressopp.interaction.SingleParticlePotential (module), 114
- espressopp.interaction.SingleParticlePotential.computeEnergy() (in module espressopp.interaction.SingleParticlePotential), 114
- espressopp.interaction.SingleParticlePotential.computeForce() (in module espressopp.interaction.SingleParticlePotential), 114
- espressopp.interaction.SoftCosine (module), 114
- espressopp.interaction.SoftCosine() (in module espressopp.interaction.SoftCosine), 115
- espressopp.interaction.StillingerWeberPairTerm (module), 115
- espressopp.interaction.StillingerWeberPairTerm() (in module espressopp.interaction.StillingerWeberPairTerm), 115
- espressopp.interaction.StillingerWeberPairTermCapped (module), 118
- espressopp.interaction.StillingerWeberPairTermCapped() (in module espressopp.interaction.StillingerWeberPairTermCapped), 118
- espressopp.interaction.StillingerWeberTripleTerm (module), 120
- espressopp.interaction.StillingerWeberTripleTerm() (in module espressopp.interaction.StillingerWeberTripleTerm), 120
- espressopp.interaction.Tabulated (module), 122
- espressopp.interaction.Tabulated() (in module espressopp.interaction.Tabulated), 122
- espressopp.interaction.TabulatedAngular (module), 124
- espressopp.interaction.TabulatedAngular() (in module espressopp.interaction.TabulatedAngular), 124
- espressopp.interaction.TabulatedDihedral (module), 125
- espressopp.interaction.TabulatedDihedral() (in module espressopp.interaction.TabulatedDihedral), 125
- espressopp.interaction.TersoffPairTerm (module), 126
- espressopp.interaction.TersoffPairTerm() (in module espressopp.interaction.TersoffPairTerm), 126
- espressopp.interaction.TersoffTripleTerm (module), 127
- espressopp.interaction.VerletListAdressLennardJones() (in module espressopp.interaction.LennardJones), 85
- espressopp.interaction.VerletListAdressLennardJones.setPotentialAT() (in module espressopp.interaction.LennardJones), 85
- espressopp.interaction.VerletListAdressLennardJones.setPotentialCG() (in module espressopp.interaction.LennardJones), 85
- espressopp.interaction.VerletListAdressLennardJones2() (in module espressopp.interaction.LennardJones), 85
- espressopp.interaction.VerletListAdressLennardJones2.setPotentialAT() (in module espressopp.interaction.LennardJones), 85
- espressopp.interaction.VerletListAdressLennardJones2.setPotentialCG() (in module espressopp.interaction.LennardJones), 85
- espressopp.interaction.VerletListAdressLennardJonesAutoBonds() (in module espressopp.interaction.LennardJonesAutoBonds), 89
- espressopp.interaction.VerletListAdressLennardJonesAutoBonds.setPotentialAT() (in module espressopp.interaction.LennardJonesAutoBonds), 89
- espressopp.interaction.VerletListAdressLennardJonesCapped() (in module espressopp.interaction.LennardJonesCapped), 91
- espressopp.interaction.VerletListAdressLennardJonesCapped.getPotentialAT() (in module espressopp.interaction.LennardJonesCapped), 91

<b>Index</b>	<b>195</b>
--------------	------------





Index 197

(in module `espres-`)

espressopp.interaction.VerletListStillingWeberPairTermCapped() (in module espressopp.interaction.StillingWeberPairTermCapped), 118  
 espressopp.interaction.VerletListStillingWeberPairTermCapped.getPotential() (in module espressopp.interaction.StillingWeberPairTermCapped), 118  
 espressopp.interaction.VerletListStillingWeberPairTermCapped.setPotential() (in module espressopp.interaction.StillingWeberPairTermCapped), 118  
 espressopp.interaction.VerletListStillingWeberPairTermCapped.getVerletList() (in module espressopp.interaction.StillingWeberPairTermCapped), 118  
 espressopp.interaction.VerletListStillingWeberPairTermCapped.setVerletList() (in module espressopp.interaction.StillingWeberPairTermCapped), 118  
 espressopp.interaction.VerletListStillingWeberTripleTerm() (in module espressopp.interaction.StillingWeberTripleTerm), 121  
 espressopp.interaction.VerletListStillingWeberTripleTerm.getPotential() (in module espressopp.interaction.StillingWeberTripleTerm), 121  
 espressopp.interaction.VerletListStillingWeberTripleTerm.setFixedTupleList() (in module espressopp.interaction.StillingWeberTripleTerm), 121  
 espressopp.interaction.VerletListStillingWeberTripleTerm.setPotential() (in module espressopp.interaction.StillingWeberTripleTerm), 121  
 espressopp.interaction.VerletListTabulated() (in module espressopp.interaction.Tabulated), 123  
 espressopp.interaction.VerletListTabulated.getPotential() (in module espressopp.interaction.Tabulated), 123  
 espressopp.interaction.VerletListTabulated.setPotential() (in module espressopp.interaction.Tabulated), 123  
 espressopp.interaction.VerletListTersoffPairTerm() (in module espressopp.interaction.TersoffPairTerm), 126  
 espressopp.interaction.VerletListTersoffPairTerm.getPotential() (in module espressopp.interaction.TersoffPairTerm), 126  
 espressopp.interaction.VerletListTersoffPairTerm.getVerletList() (in module espressopp.interaction.TersoffPairTerm), 126  
 espressopp.interaction.VerletListTersoffPairTerm.setPotential() (in module espressopp.interaction.TersoffPairTerm), 126  
 espressopp.interaction.VerletListTersoffTripleTerm() (in module espressopp.interaction.TersoffTripleTerm), 127  
 espressopp.interaction.VerletListTersoffTripleTerm.getPotential() (in module espressopp.interaction.TersoffTripleTerm), 127  
 espressopp.interaction.VerletListTersoffTripleTerm.getVerletList() (in module espressopp.interaction.TersoffTripleTerm), 127  
 espressopp.interaction.VerletListTersoffTripleTerm.setPotential() (in module espressopp.interaction.TersoffTripleTerm), 127  
 espressopp.interaction.VerletListVSpherePair() (in module espressopp.interaction.VSpherePair), 128  
 espressopp.interaction.VerletListVSpherePair.getPotential() (in module espressopp.interaction.VSpherePair), 128  
 espressopp.interaction.VerletListVSpherePair.getVerletList() (in module espressopp.interaction.VSpherePair), 128  
 espressopp.interaction.VerletListVSpherePair.setPotential() (in module espressopp.interaction.VSpherePair), 128  
 espressopp.interaction.VerletListZero() (in module espressopp.interaction.Zero), 129  
 espressopp.interaction.VerletListZero.getPotential() (in module espressopp.interaction.Zero), 130  
 espressopp.interaction.VerletListZero.setFixedTupleList() (in module espressopp.interaction.Zero), 130  
 espressopp.interaction.VerletListZero.setPotential() (in module espressopp.interaction.Zero), 130  
 espressopp.io.DumpGRO (module), 131  
 espressopp.io.DumpGRO() (in module espressopp.io.DumpGRO), 132  
 espressopp.io.DumpGRO.dump() (in module espressopp.io.DumpGRO), 132  
 espressopp.io.DumpGROAddress (module), 132  
 espressopp.io.DumpGROAddress() (in module espressopp.io.DumpGROAddress), 133  
 espressopp.io.DumpGROAddress.dump() (in module espressopp.io.DumpGROAddress), 133  
 espressopp.io.DumpXYZ (module), 133  
 espressopp.io.DumpXYZ() (in module espressopp.io.DumpXYZ), 134  
 espressopp.io.DumpXYZ.dump() (in module espressopp.io.DumpXYZ), 134  
 espressopp.MissingFixedPairList() (in module espressopp.Exceptions), 135  
 espressopp.MultiSystem (module), 142  
 espressopp.MultiSystem() (in module espressopp.MultiSystem), 142

- sopp.MultiSystem), 142
- espressopp.MultiSystem.beginSystemDefinition() (in module espressopp.MultiSystem), 142
- espressopp.MultiSystem.runAnalysisNPart() (in module espressopp.MultiSystem), 142
- espressopp.MultiSystem.runAnalysisPotential() (in module espressopp.MultiSystem), 142
- espressopp.MultiSystem.runAnalysisTemperature() (in module espressopp.MultiSystem), 142
- espressopp.MultiSystem.runIntegrator() (in module espressopp.MultiSystem), 142
- espressopp.MultiSystem.setAnalysisNPart() (in module espressopp.MultiSystem), 142
- espressopp.MultiSystem.setAnalysisPotential() (in module espressopp.MultiSystem), 142
- espressopp.MultiSystem.setAnalysisTemperature() (in module espressopp.MultiSystem), 142
- espressopp.MultiSystem.setIntegrator() (in module espressopp.MultiSystem), 142
- espressopp.ParallelTempering (module), 142
- espressopp.ParallelTempering() (in module espressopp.ParallelTempering), 142
- espressopp.ParallelTempering.endDefiningSystem() (in module espressopp.ParallelTempering), 142
- espressopp.ParallelTempering.exchange() (in module espressopp.ParallelTempering), 142
- espressopp.ParallelTempering.getNumberOfCPUsPerSystem() (in module espressopp.ParallelTempering), 143
- espressopp.ParallelTempering.getNumberOfSystems() (in module espressopp.ParallelTempering), 143
- espressopp.ParallelTempering.run() (in module espressopp.ParallelTempering), 143
- espressopp.ParallelTempering.setAnalysisE() (in module espressopp.ParallelTempering), 143
- espressopp.ParallelTempering.setAnalysisNPart() (in module espressopp.ParallelTempering), 143
- espressopp.ParallelTempering.setAnalysisT() (in module espressopp.ParallelTempering), 143
- espressopp.ParallelTempering.setIntegrator() (in module espressopp.ParallelTempering), 143
- espressopp.ParallelTempering.startDefiningSystem() (in module espressopp.ParallelTempering), 143
- espressopp.Particle (module), 143
- espressopp.Particle() (in module espressopp.Particle), 143
- espressopp.ParticleAccess (module), 143
- espressopp.ParticleAccess.perform\_action() (in module espressopp.ParticleAccess), 143
- espressopp.ParticleDoesNotExistHere() (in module espressopp.Exceptions), 134
- espressopp.ParticleGroup (module), 143
- espressopp.ParticleGroup() (in module espressopp.ParticleGroup), 144
- espressopp.ParticleGroup.add() (in module espressopp.ParticleGroup), 144
- espressopp.ParticleGroup.has() (in module espressopp.ParticleGroup), 144
- espressopp.ParticleGroup.show() (in module espressopp.ParticleGroup), 144
- espressopp.ParticleGroup.size() (in module espressopp.ParticleGroup), 144
- espressopp.pmi (module), 144
- espressopp.Quaternion (module), 148
- espressopp.Real3D (module), 149
- espressopp.RealND (module), 150
- espressopp.standard\_system.Default (module), 154
- espressopp.standard\_system.KGMelt (module), 155
- espressopp.standard\_system.KGMelt() (in module espressopp.standard\_system.KGMelt), 155
- espressopp.standard\_system.LennardJones (module), 155
- espressopp.standard\_system.LennardJones() (in module espressopp.standard\_system.LennardJones), 155
- espressopp.standard\_system.Minimal (module), 155
- espressopp.standard\_system.Minimal() (in module espressopp.standard\_system.Minimal), 155
- espressopp.standard\_system.PolymerMelt (module), 156
- espressopp.standard\_system.PolymerMelt() (in module espressopp.standard\_system.PolymerMelt), 156
- espressopp.storage.DomainDecomposition (module), 156
- espressopp.storage.DomainDecomposition() (in module espressopp.storage.DomainDecomposition), 156
- espressopp.storage.DomainDecomposition.getCellGrid() (in module espressopp.storage.DomainDecomposition), 156
- espressopp.storage.DomainDecomposition.getNodeGrid() (in module espressopp.storage.DomainDecomposition), 156
- espressopp.storage.DomainDecompositionAdress (module), 156
- espressopp.storage.DomainDecompositionAdress() (in module espressopp.storage.DomainDecompositionAdress), 157
- espressopp.storage.DomainDecompositionNonBlocking (module), 157
- espressopp.storage.DomainDecompositionNonBlocking() (in module espressopp.storage.DomainDecompositionNonBlocking), 157
- espressopp.storage.Storage (module), 157
- espressopp.storage.Storage.addAdrATParticle() (in module espressopp.storage.Storage), 158
- espressopp.storage.Storage.addParticle() (in module

- espressopp.storage.Storage), 158
- espressopp.storage.Storage.addParticles() (in module espressopp.storage.Storage), 158
- espressopp.storage.Storage.clearSavedPositions() (in module espressopp.storage.Storage), 158
- espressopp.storage.Storage.getParticle() (in module espressopp.storage.Storage), 158
- espressopp.storage.Storage.getRealParticleIDs() (in module espressopp.storage.Storage), 159
- espressopp.storage.Storage.modifyParticle() (in module espressopp.storage.Storage), 159
- espressopp.storage.Storage.particleExists() (in module espressopp.storage.Storage), 159
- espressopp.storage.Storage.printRealParticles() (in module espressopp.storage.Storage), 159
- espressopp.storage.Storage.removeAllParticles() (in module espressopp.storage.Storage), 159
- espressopp.storage.Storage.removeParticle() (in module espressopp.storage.Storage), 159
- espressopp.storage.Storage.restorePositions() (in module espressopp.storage.Storage), 159
- espressopp.storage.Storage.savePositions() (in module espressopp.storage.Storage), 159
- espressopp.storage.Storage.setFixedTuplesAdress() (in module espressopp.storage.Storage), 159
- espressopp.System (module), 150
- espressopp.System() (in module espressopp.System), 151
- espressopp.System.addInteraction() (in module espressopp.System), 151
- espressopp.System.getAllInteractions() (in module espressopp.System), 151
- espressopp.System.getInteraction() (in module espressopp.System), 151
- espressopp.System.getNumberOfInteractions() (in module espressopp.System), 151
- espressopp.System.removeInteraction() (in module espressopp.System), 151
- espressopp.System.removeInteractionByName() (in module espressopp.System), 151
- espressopp.System.scaleVolume() (in module espressopp.System), 151
- espressopp.System.setTrace() (in module espressopp.System), 151
- espressopp.Tensor (module), 151
- espressopp.toInt3D() (in module espressopp.Int3D), 141
- espressopp.toInt3DFromVector() (in module espressopp.Int3D), 141
- espressopp.tools.analyse (module), 160
- espressopp.tools.applyBoreschRestrains() (in module espressopp.tools.prepareComplexMolecules), 169
- espressopp.tools.decomp (module), 161
- espressopp.tools.DumpConfigurations (module), 163
- espressopp.tools.espresso\_old (module), 164
- espressopp.tools.findConstrainedBonds() (in module espressopp.tools.prepareComplexMolecules), 168
- espressopp.tools.getInternalNonbondedInteractions() (in module espressopp.tools.prepareComplexMolecules), 168
- espressopp.tools.gromacs (module), 164
- espressopp.tools.info (module), 160
- espressopp.tools.io\_extended (module), 166
- espressopp.tools.lammps (module), 166
- espressopp.tools.lattice (module), 160
- espressopp.tools.pathintegral (module), 167
- espressopp.tools.pdb (module), 167
- espressopp.tools.pdbread() (in module espressopp.tools.pdb), 167
- espressopp.tools.pdbwrite() (in module espressopp.tools.pdb), 167
- espressopp.tools.povwrite (module), 168
- espressopp.tools.prepareAdress (module), 169
- espressopp.tools.prepareComplexMolecules (module), 168
- espressopp.tools.psf (module), 169
- espressopp.tools.psfread() (in module espressopp.tools.psf), 170
- espressopp.tools.psfwrite() (in module espressopp.tools.psf), 170
- espressopp.tools.readSimpleSystem() (in module espressopp.tools.prepareComplexMolecules), 168
- espressopp.tools.replicate (module), 161
- espressopp.tools.tabulated (module), 170
- espressopp.tools.timers (module), 160
- espressopp.tools.topology (module), 161
- espressopp.tools.topology\_helper (module), 170
- espressopp.tools.units (module), 170
- espressopp.tools.velocities (module), 161
- espressopp.tools.vmd (module), 160
- espressopp.tools.warmup (module), 161
- espressopp.toReal3D() (in module espressopp.Real3D), 150
- espressopp.toReal3DFromVector() (in module espressopp.Real3D), 150
- espressopp.toRealND() (in module espressopp.RealND), 150
- espressopp.toRealNDFromVector() (in module espressopp.RealND), 150
- espressopp.UnknownParticleProperty() (in module espressopp.Exceptions), 135
- espressopp.VerletList (module), 152
- espressopp.VerletList() (in module espressopp.VerletList), 152
- espressopp.VerletList.exclude() (in module espressopp.VerletList), 152
- espressopp.VerletList.getAllPairs() (in module espressopp.VerletList), 152
- espressopp.VerletList.localSize() (in module espressopp.VerletList), 152



- sopp.VerletList), 152
  - espressopp.VerletList.totalSize() (in module espressopp.VerletList), 152
  - espressopp.VerletListAdress (module), 152
  - espressopp.VerletListAdress() (in module espressopp.VerletListAdress), 153
  - espressopp.VerletListAdress.addAdrParticles() (in module espressopp.VerletListAdress), 153
  - espressopp.VerletListAdress.exclude() (in module espressopp.VerletListAdress), 153
  - espressopp.VerletListAdress.rebuild() (in module espressopp.VerletListAdress), 153
  - espressopp.VerletListAdress.totalSize() (in module espressopp.VerletListAdress), 153
  - espressopp.VerletListTriple (module), 154
  - espressopp.VerletListTriple() (in module espressopp.VerletListTriple), 154
  - espressopp.VerletListTriple.exclude() (in module espressopp.VerletListTriple), 154
  - espressopp.VerletListTriple.getAllTriples() (in module espressopp.VerletListTriple), 154
  - espressopp.VerletListTriple.localSize() (in module espressopp.VerletListTriple), 154
  - espressopp.VerletListTriple.totalSize() (in module espressopp.VerletListTriple), 154
  - espressopp.Version (module), 154
  - espressopp.Version() (in module espressopp.Version), 154
  - espressopppp.integrator.Rattle() (in module espressopp.integrator.Rattle), 63
  - espressopppp.integrator.Rattle.addConstrainedBonds() (in module espressopp.integrator.Rattle), 63
  - espressopppp.interaction.CoulombTruncated() (in module espressopp.interaction.CoulombTruncated), 72
  - espressopppp.interaction.FixedPairListTypesCoulombTruncated() (in module espressopp.interaction.CoulombTruncated), 73
  - espressopppp.interaction.FixedPairListTypesCoulombTruncated.setPotential() (in module espressopp.interaction.CoulombTruncated), 73
  - espressopppp.interaction.LennardJonesSoftcoreTI() (in module espressopp.interaction.LennardJonesSoftcoreTI), 103
  - espressopppp.interaction.LennardJonesSoftcoreTI.addPids() (in module espressopp.interaction.LennardJonesSoftcoreTI), 103
  - espressopppp.interaction.ReactionFieldGeneralizedTI() (in module espressopp.interaction.ReactionFieldGeneralizedTI), 113
  - espressopppp.interaction.ReactionFieldGeneralizedTI.addPids() (in module espressopp.interaction.ReactionFieldGeneralizedTI), 113
  - espressopppp.interaction.VerletListAdressLennardJones() (in module espressopp.interaction.LennardJonesSoftcoreTI), 103
  - espressopppp.interaction.VerletListAdressLennardJones.setPotentialAT() (in module espressopp.interaction.LennardJonesSoftcoreTI), 103
  - espressopppp.interaction.VerletListAdressLennardJones.setPotentialCG() (in module espressopp.interaction.LennardJonesSoftcoreTI), 103
  - espressopppp.interaction.VerletListAdressReactionFieldGeneralized() (in module espressopp.interaction.ReactionFieldGeneralizedTI), 113
  - espressopppp.interaction.VerletListAdressReactionFieldGeneralized.setP() (in module espressopp.interaction.ReactionFieldGeneralizedTI), 114
  - espressopppp.interaction.VerletListAdressReactionFieldGeneralized.setP() (in module espressopp.interaction.ReactionFieldGeneralizedTI), 114
  - espressopppp.interaction.VerletListCoulombTruncated() (in module espressopp.interaction.CoulombTruncated), 72
  - espressopppp.interaction.VerletListCoulombTruncated.getPotential() (in module espressopp.interaction.CoulombTruncated), 72
  - espressopppp.interaction.VerletListCoulombTruncated.setPotential() (in module espressopp.interaction.CoulombTruncated), 72
  - exec\_() (in module espressopp.pmi), 145
- ## F
- fastreadxyz() (in module espressopp.tools.DumpConfigurations), 163
  - fastwritexyz() (in module espressopp.tools.DumpConfigurations), 163
  - fastwritexyz\_standard() (in module espressopp.tools.DumpConfigurations), 163
  - FENECapped (class in espressopp.interaction.FENECapped), 81
  - final\_info() (in module espressopp.tools.analyse), 160
  - finalizeWorkers() (in module espressopp.pmi), 147
  - FixedQuadrupleListDihedralHarmonicLocal (class in espressopp.interaction.DihedralHarmonic), 76
  - FixedQuadrupleListDihedralHarmonicNCosLocal (class in espressopp.interaction.DihedralHarmonicNCos), 77
- ## G
- gaussian() (in module espressopp.tools.velocities), 161
  - getAllBonds() (in module espressopp.tools.info), 160

- getAllParticles() (in module espressopp.tools.info), 160  
 getFoldedPosition() (espressopp.bc.BC.espressopp.bc.BC method), 42  
 getLBMom() (espressopp.integrator.LatticeBoltzmann.espressopp.integrator.LatticeBoltzmann method), 56  
 getMinimumImageVector() (espressopp.bc.BC.espressopp.bc.BC method), 42  
 getRandomPos() (espressopp.bc.BC.espressopp.bc.BC method), 42  
 getUnfoldedPosition() (espressopp.bc.BC.espressopp.bc.BC method), 42
- ## H
- Harmonic (class in espressopp.interaction.Harmonic), 83  
 HarmonicTrap (class in espressopp.interaction.HarmonicTrap), 83  
 HarmonicUnique (class in espressopp.interaction.HarmonicUnique), 84
- ## I
- import\_() (in module espressopp.pmi), 146  
 info() (in module espressopp.tools.analyse), 160  
 invoke() (in module espressopp.pmi), 146
- ## K
- keepLBDump() (espressopp.integrator.LatticeBoltzmann.espressopp.integrator.LatticeBoltzmann method), 56
- ## L
- LennardJones (class in espressopp.interaction.LennardJones), 87  
 LennardJones93Wall (class in espressopp.interaction.LennardJones93Wall), 88  
 LennardJonesAutoBonds (class in espressopp.interaction.LennardJonesAutoBonds), 90  
 LennardJonesCapped (class in espressopp.interaction.LennardJonesCapped), 93  
 LennardJonesEnergyCapped (class in espressopp.interaction.LennardJonesEnergyCapped), 96  
 LennardJonesExpand (class in espressopp.interaction.LennardJonesExpand), 97  
 LennardJonesGeneric (class in espressopp.interaction.LennardJonesGeneric), 100  
 LennardJonesGromacs (class in espressopp.interaction.LennardJonesGromacs), 102  
 LennardJonesSoftcoreTI (class in espressopp.interaction.LennardJonesSoftcoreTI), 104  
 LJcos (class in espressopp.interaction.LJcos), 105  
 locateItem() (espressopp.esutil.Collectives.espressopp.esutil.CollectivesLocateItem), 144
- ## M
- MirrorLennardJones (class in espressopp.interaction.MirrorLennardJones), 106  
 Morse (class in espressopp.interaction.Morse), 108  
 MultiSystem (class in espressopp.MultiSystem), 142  
 MultiSystemLocal (class in espressopp.MultiSystem), 142
- ## O
- OPLS (class in espressopp.interaction.OPLS), 109  
 OrthorhombicBC() (espressopp.bc.OrthorhombicBC.espressopp.bc.OrthorhombicBC method), 43
- ## P
- ParticleLocal (class in espressopp.Particle), 143  
 polymerRW() (in module espressopp.tools.topology), 161  
 Proxy (class in espressopp.pmi), 147
- ## Q
- Quartic (class in espressopp.interaction.Quartic), 110
- ## R
- ReactionFieldGeneralized (class in espressopp.interaction.ReactionFieldGeneralized), 112  
 ReactionFieldGeneralizedTI (class in espressopp.interaction.ReactionFieldGeneralizedTI), 114  
 read() (in module espressopp.tools.espresso\_old), 164  
 read() (in module espressopp.tools.gromacs), 166  
 readxyz() (in module espressopp.tools.DumpConfigurations), 163  
 readxyzr() (in module espressopp.tools.DumpConfigurations), 163  
 receive() (in module espressopp.pmi), 147  
 reduce() (in module espressopp.pmi), 147  
 registerAtExit() (in module espressopp.pmi), 147  
 replicate() (in module espressopp.tools.replicate), 161
- ## S
- saveLBConf() (espressopp.integrator.LatticeBoltzmann.espressopp.integrator.LatticeBoltzmann method), 56  
 setBoxL() (espressopp.bc.OrthorhombicBC.espressopp.bc.OrthorhombicBC method), 43  
 setBoxL() (espressopp.bc.SlabBC.espressopp.bc.SlabBC method), 43

[setForce\(\)](#) (`espressopp.integrator.LBInit.espressopp.integrator.LBInit` method), [58](#)  
[setLBMom\(\)](#) (`espressopp.integrator.LatticeBoltzmann.espressopp.integrator.LatticeBoltzmann` method), [56](#)  
[setLennardJones14Interactions\(\)](#) (in module `espressopp.tools.gromacs`), [166](#)  
[setLennardJonesInteractions\(\)](#) (in module `espressopp.tools.gromacs`), [166](#)  
[setLennardJonesInteractionsTI\(\)](#) (in module `espressopp.tools.gromacs`), [166](#)  
[setTabulatedInteractions\(\)](#) (in module `espressopp.tools.gromacs`), [166](#)  
[show\(\)](#) (in module `espressopp.tools.timers`), [160](#)  
[SlabBC\(\)](#) (`espressopp.bc.SlabBC.espressopp.bc` method), [43](#)  
[SoftCosine](#) (class in `espressopp.interaction.SoftCosine`), [115](#)  
[startWorkerLoop\(\)](#) (in module `espressopp.pmi`), [147](#)  
[step](#) (in module `espressopp.integrator.MinimizeEnergy`), [61](#)  
[StillingerWeberPairTerm](#) (class in `espressopp.interaction.StillingerWeberPairTerm`), [118](#)  
[StillingerWeberPairTermCapped](#) (class in `espressopp.interaction.StillingerWeberPairTermCapped`), [120](#)  
[StillingerWeberTripleTerm](#) (class in `espressopp.interaction.StillingerWeberTripleTerm`), [122](#)  
[stopWorkerLoop\(\)](#) (in module `espressopp.pmi`), [147](#)  
[sync\(\)](#) (in module `espressopp.pmi`), [147](#)

## T

[Tabulated](#) (class in `espressopp.interaction.Tabulated`), [124](#)  
[TabulatedAngular](#) (class in `espressopp.interaction.TabulatedAngular`), [125](#)  
[TabulatedDihedral](#) (class in `espressopp.interaction.TabulatedDihedral`), [126](#)  
[TersoffPairTerm](#) (class in `espressopp.interaction.TersoffPairTerm`), [127](#)  
[toInt3D\(\)](#) (in module `espressopp.Int3D`), [142](#)  
[toInt3DFromVector\(\)](#) (in module `espressopp.Int3D`), [142](#)  
[toQuaternion\(\)](#) (in module `espressopp.Quaternion`), [149](#)  
[toQuaternionFromVector\(\)](#) (in module `espressopp.Quaternion`), [149](#)  
[toReal3D\(\)](#) (in module `espressopp.Real3D`), [150](#)  
[toReal3DFromVector\(\)](#) (in module `espressopp.Real3D`), [150](#)  
[toRealND\(\)](#) (in module `espressopp.RealND`), [150](#)  
[toRealNDFromVector\(\)](#) (in module `espressopp.RealND`), [150](#)  
[toTensor\(\)](#) (in module `espressopp.Tensor`), [152](#)  
[toTensorFromVector\(\)](#) (in module `espressopp.Tensor`), [152](#)

## U

[UserError](#), [148](#)

## V

[VSpherePair](#) (class in `espressopp.interaction.VSpherePair`), [129](#)  
[VSphereSelf](#) (class in `espressopp.interaction.VSphereSelf`), [129](#)

## W

[warmup\(\)](#) (in module `espressopp.tools.warmup`), [161](#)  
[writeTabFile\(\)](#) (in module `espressopp.tools.tabulated`), [170](#)  
[writexyz\(\)](#) (in module `espressopp.tools.DumpConfigurations`), [163](#)

## X

[xyzfilewrite\(\)](#) (in module `espressopp.tools.DumpConfigurations`), [163](#)

## Z

[Zero](#) (class in `espressopp.interaction.Zero`), [131](#)