

Developer Guide

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OpenMM Developer Guide

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1 Introduction

This guide describes the internal architecture of the OpenMM library. It is targeted at developers who want to add features to OpenMM, either by modifying the core library directly or by writing plugins. If you just want to write applications that use OpenMM, you do not need to read this guide; the Users Manual tells you everything you need to know. This guide is *only* for people who want to contribute to OpenMM itself.

It is organized as follows:

- Chapter 2 describes the architecture of the core OpenMM library. It discusses how the high level and low level APIs relate to each other, and the flow of execution between them.
- Chapter 3 describes in detail how to write a plugin. It focuses on the two most common types of plugins: those which define new Forces, and those which implement new Platforms.
- Chapter 4 discusses the architecture of the reference Platform, providing information relevant to writing reference implementations of new features.
- Chapter 5 discusses the architecture of the OpenCL Platform, providing information relevant to writing OpenCL implementations of new features.

This guide assumes you are already familiar with the public API and how to use OpenMM in applications. If that is not the case, you should first read the Users Manual and work through some of the example programs.

2 The Core Library

OpenMM is based on a layered architecture, as shown in the following diagram:

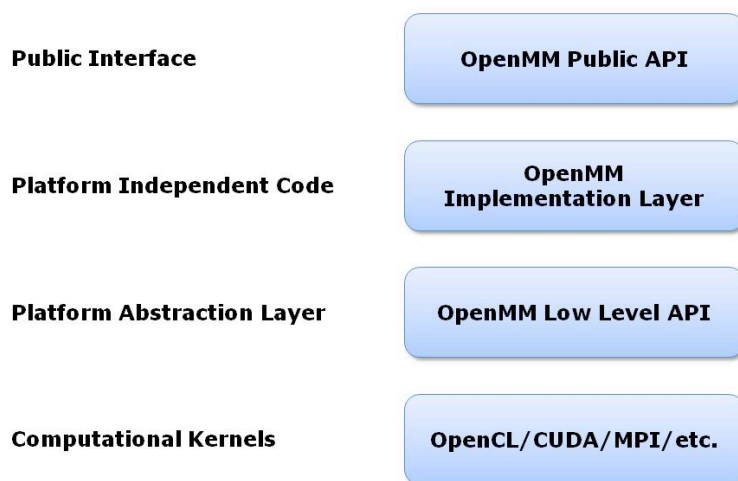


Figure 2.1: OpenMM architecture

The public API layer consists of the classes you access when using OpenMM in an application: `System`; `Force` and its subclasses; `Integrator` and its subclasses; and `Context`. These classes define a public interface but do no computation.

The next layer down consists of “implementation” classes that mirror the public API classes: `ContextImpl`, `ForceImpl`, and a subclass of `ForceImpl` for each subclass of `Force` (`HarmonicBondForceImpl`, `NonbondedForceImpl`, etc.). These objects are created automatically when you create a `Context`. They store information related to a particular simulation, and define methods for performing calculations.

Note that, whereas a `Force` is logically “part of” a `System`, a `ForceImpl` is logically “part of” a `Context`. If you create many `Contexts` for simulating the same `System`, there is still only one

System and only one copy of each Force in it. But there will be separate ForceImpls for each Context, and those ForceImpls store information related to their particular Contexts.

Also note that there is no “IntegratorImpl” class, because it is not needed. Integrator is already specific to one Context. Many Contexts can all simulate the same System, but each of them must have its own Integrator, so information specific to one simulation can be stored directly in the Integrator.

The next layer down is the OpenMM Low Level API (OLLA). The important classes in this layer are: Platform; Kernel; KernelImpl and its subclasses; and KernelFactory. A Kernel is just a reference counted pointer to a KernelImpl; the real work is done by KernelImpl objects (or more precisely, by instances of its subclasses). A KernelFactory creates KernelImpl objects, and a Platform ties together a set of KernelFactories, as well as defining information that applies generally to performing computations with that Platform.

All of these classes (except Kernel) are abstract. A particular Platform provides concrete subclasses of all of them. For example, the reference platform defines a Platform subclass called ReferencePlatform, a KernelFactory subclass called ReferenceKernelFactory, and a concrete subclass of each abstract KernelImpl type: ReferenceCalcNonbondedForceKernel extends CalcNonbondedForceKernel (which in turn extends KernelImpl), ReferenceIntegrateVerletStepKernel extends IntegrateVerletStepKernel, and so on.

We can understand this better by walking through the entire sequence of events that takes place when you create a Context. As an example, suppose you create a System; add a NonbondedForce to it; create a VerletIntegrator; and then create a Context for them using the reference Platform. Here is what happens.

1. The Context constructor creates a ContextImpl.
2. The ContextImpl calls `createImpl()` on each Force in the System, which creates an instance of the appropriate ForceImpl subclass.
3. The ContextImpl calls `contextCreated()` on the Platform(), which in turn calls `setPlatformData()` on the ContextImpl. This allows Platform-specific information to be stored in a ContextImpl. Every Platform has its own mechanism

for storing particle masses, constraint definitions, particle positions, and so on. `ContextImpl` therefore allows the Platform to create an arbitrary block of data and store it where it can be accessed by that Platform's kernels.

4. The `ContextImpl` calls `createKernel()` on the Platform several times to get instances of various kernels that it needs: `CalcKineticEnergyKernel`, `ApplyConstraintsKernel`, etc.
 - a. For each kernel, the Platform looks up which `KernelFactory` has been registered for that particular kernel. In this case, it will be a `ReferenceKernelFactory`.
 - b. It calls `createKernelImpl()` on the `KernelFactory`, which creates and returns an instance of an appropriate `KernelImpl` subclass: `ReferenceCalcKineticEnergyKernel`, `ReferenceApplyConstraintsKernel`, etc.
5. The `ContextImpl` loops over all of its `ForceImpl`s and calls `initialize()` on each one.
 - a. Each `ForceImpl` asks the Platform to create whatever kernels it needs. In this example, `NonbondedForceImpl` will request a `CalcNonbondedForceKernel`, and get back a `ReferenceCalcNonbondedForceKernel`.
6. The `ContextImpl` calls `initialize()` on the Integrator which, like the other objects, requests kernels from the Platform. In this example, `VerletIntegrator` requests an `IntegrateVerletStepKernel` and gets back a `ReferenceIntegrateVerletStepKernel`.

At this point, the Context is fully initialized and ready for doing computation. Reference implementations of various `KernelImpl`s have been created, but they are always referenced through abstract superclasses. Similarly, data structures specific to the reference Platform have been created and stored in the `ContextImpl`, but the format and content of these structures is opaque to the `ContextImpl`. Whenever it needs to access them (for example, to get or set particle positions), it does so through a kernel (`UpdateStateDataKernel` in this case).

Now suppose that you call `step()` on the `VerletIntegrator`. Here is what happens to execute each time step.

1. The `VerletIntegrator` calls `updateContextState()` on the `ContextImpl`. This gives each `Force` an opportunity to modify the state of the `Context` at the start of each time step.
 - a. The `ContextImpl` loops over its `ForceImpls` and calls `updateContextState()` on each one. In this case, our only `ForceImpl` is a `NonbondedForceImpl`, which returns without doing anything. On the other hand, if we had an `AndersenThermostat` in our `System`, its `ForceImpl` would invoke a kernel to modify particle velocities.
2. The `VerletIntegrator` calls `calcForcesAndEnergy()` on the `ContextImpl` to request that the forces be computed.
 - a. The `ContextImpl` calls `beginComputation()` on its `CalcForcesAndEnergyKernel`. This initializes all the forces to zero and does any other initialization the `Platform` requires before forces can be computed. For example, some `Platforms` construct their nonbonded neighbor lists at this point.
 - b. The `ContextImpl` loops over its `ForceImpls` and calls `calcForcesAndEnergy()` on each one. In this case, we have a `NonbondedForceImpl` which invokes its `CalcNonbondedForceKernel` to compute forces.
 - c. Finally, the `ContextImpl` calls `finishComputation()` on its `CalcForcesAndEnergyKernel`. This does any additional work needed to determine the final forces, such as summing the values from intermediate buffers.
3. Finally, the `VerletIntegrator` invokes its `IntegrateVerletStepKernel`. This takes the forces, positions, and velocities that are stored in a `Platform`-specific format in the `ContextImpl`, uses them to compute new positions and velocities, and stores them in the `ContextImpl`.

3 Writing Plugins

A plugin is a dynamic library that adds new features to OpenMM. It is typically stored in the `lib/plugins` directory inside your OpenMM installation, and gets loaded along with all other plugins when the user calls

```
Platform::loadPluginsFromDirectory(  
    Platform::getDefaultPluginsDirectory());
```

It is also possible to load plugins from a different directory, or to load them individually by calling `Platform::loadPluginLibrary()`.

Every plugin must implement two functions that are declared in the `PluginInitializer.h` header file:

```
extern "C" void registerPlatforms();  
extern "C" void registerKernelFactories();
```

When a plugin is loaded, these two functions are invoked to register any Platforms and KernelFactories defined by the plugin. When many plugins are loaded at once by calling `Platform::loadPluginsFromDirectory()`, `registerPlatforms()` is first called on all of them, then `registerKernelFactories()` is called on all of them. This allows one plugin to define a Platform, and a different plugin to add KernelFactories to it; the Platform is guaranteed to be registered by the first plugin before the second plugin tries to add its KernelFactories, regardless of what order the plugins happen to be loaded in.

3.1 Creating New Platforms

One common type of plugin defines a new Platform. There are two such plugins that come with OpenMM: one for the CUDA Platform and one for the OpenCL Platform.

To define a new Platform, you must create subclasses of the various abstract classes in the OpenMM Low Level API: a subclass of Platform, one or more subclasses of KernelFactory, and a subclass of each KernelImpl. That is easy to say, but a huge amount of work to actually do. There are many different algorithms involved in computing forces, enforcing constraints, performing integration, and so on, all of which together make up a Platform. Of course, there is no requirement that every Platform must implement every possible feature. If you do not provide an implementation of a particular kernel, it simply means your Platform cannot be used for any simulation that requires that kernel; if a user tries to do so, an exception will be thrown.

Your plugin's `registerPlatforms()` function should create an instance of your Platform subclass, then register it by calling `Platform::registerPlatform()`. You also must register the KernelFactory for each kernel your Platform supports. This can be done in the `registerKernelFactories()` function, or more simply, directly in the Platform's constructor. You can use as many different KernelFactories as you want for different kernels, but usually it is simplest to use a single KernelFactory for all of them. The support for multiple KernelFactories exists primarily to let plugins add new features to existing Platforms, as described in the next section.

3.2 Creating New Forces

Another common type of plugin defines new Forces and provides implementations of them for existing Platforms. (Defining new Integrators is not specifically discussed here, but the process is very similar.) There are two such plugins that come with OpenMM: one for the AMOEBA force field, and one that provides various forces useful in free energy calculations.

As an example, suppose you want to create a new Force subclass called `StringForce` that uses the equations of String Theory to compute the interactions between particles. You want to provide implementations of it for all three standard platforms: reference, CUDA, and OpenCL.

The first thing to realize is that this *cannot* be done with only a plugin library. Plugins are loaded dynamically at runtime, and they relate to the low level API; but you must also provide a public API. Users of your class need to create `StringForce` objects and call methods on them. That means providing a header file with the class declaration, and a (non-plugin) library with the class definition to link their code against. The implementations for particular Platforms can be in plugins, but the public API class itself cannot. Or to put it differently, the full “plugin” (from the user’s perspective) consists of three parts: the library OpenMM loads at runtime (which is what OpenMM considers to be the “plugin”), a second library for users to link their code against, and a header file for them to include in their source code.

The AMOEBA and FreeEnergy plugins go a step further: they also include their reference implementations in the non-plugin libraries, while using plugins for their CUDA implementations. This is consistent with OpenMM itself, which defines the reference Platform directly in the main library. You can do this for your own Force, or you can put the reference implementation in a plugin. Either approach is reasonable.

In any case, you will need to create the following classes:

1. `StringForce`. This is the public API for your force, and users will directly link against the library containing it.
2. `StringForceImpl`. This is the `ForceImpl` subclass corresponding to `StringForce`. It should be defined in the same library as `StringForce`, and `StringForce`’s `createImpl()` method should create an instance of it.
3. `CalcStringForceKernel`. This is an abstract class that extends `KernelImpl`, and defines the API by which `StringForceImpl` invokes its kernel. You only need to provide a header file for it, not an implementation; those will be provided by Platforms.

Now suppose you are writing the OpenCL implementation of `StringForce`. Here are the classes you need to write:

1. `OpenCLCalcStringForceKernel`. This extends `CalcStringForceKernel` and provides implementations of its virtual methods. The code for this class will probably be very complicated (and if it actually works, worth a Nobel Prize). It may execute many different GPU kernels and create its own internal data structures. But those details are entirely internal to your own code. As long as this class implements the virtual methods of `CalcStringForceKernel`, you can do anything you want inside it.
2. `OpenCLStringForceKernelFactory`. This is a `KernelFactory` subclass that knows how to create instances of `OpenCLCalcStringForceKernel`.

Both of these classes should be packaged into a dynamic library (.so on Linux, .dylib on Mac, .dll on Windows) that can be loaded as a plugin. This library must also implement the two functions from `PluginInitializer.h`. `registerPlatforms()` will do nothing, since this plugin does not implement any new Platforms. `registerKernelFactories()` should call `Platform::getPlatformByName("OpenCL")` to get the OpenCL Platform, then create a new `OpenCLStringForceKernelFactory` and call `registerKernelFactory()` on the Platform to register it. If the OpenCL Platform is not available, you should catch the exception then return without doing anything. Most likely this means there is no OpenCL runtime on the computer your code is running on.

4 The Reference Platform

The reference Platform is written with simplicity and clarity in mind, not performance. (It is still not always as simple or clear as one might hope, but that is the goal.) When implementing a new feature, it is recommended to create the reference implementation first, then use that as a model for the versions in other Platforms.

The reference Platform represents all floating point numbers with the type `RealOpenMM`, which is defined in `SimTKOpenMMRealType.h`. This allows the entire platform to be compiled in either single or double precision. By default it is double precision, but it can be changed by modifying one flag at the top of that file. The same file also defines lots of numerical constants and mathematical functions, so the correct precision version will always be used. Vector quantities (positions, velocities, etc.) are represented by `RealVec` objects. This class is identical to `Vec3`, except that its components are of type `RealOpenMM` instead of `double`.

When using the reference Platform, the “platform-specific data” stored in `ContextImpl` is of type `ReferencePlatform::PlatformData`, which is declared in `ReferencePlatform.h`. Several of the fields in this class are declared as `void*` to avoid having to include `SimTKOpenMMRealType.h` in `ReferencePlatform.h`. If you look in `ReferenceKernels.cpp`, you will find code for extracting the correct values of these fields. For example:

```
static vector<RealVec>& extractPositions(ContextImpl& context) {
    ReferencePlatform::PlatformData* data =
        reinterpret_cast<ReferencePlatform::PlatformData*>(
            context.getPlatformData());
    return *((vector<RealVec>*) data->positions);
}
```

The PlatformData's vector of forces contains one element for each particle. At the start of each force evaluation, all elements of it are set to zero. Each Force adds its own contributions to the vector, so that at the end, it contains the total force acting on each particle.

There are a few additional classes that contain useful static methods. SimTKOpenMMUtilities has various utility functions, of which the most important is a random number generator. ReferenceForce provides methods for calculating the displacement between two positions, optionally taking periodic boundary conditions into account.

5 The OpenCL Platform

The OpenCL Platform is much more complicated than the reference Platform. It also provides many more tools to simplify your work, but those tools themselves can be complicated to use correctly. This chapter will attempt to explain how to use some of the most important ones. It will *not* teach you how to program with OpenCL. There are many tutorials on that subject available elsewhere, and this guide assumes you already understand it.

5.1 Overview

When using the OpenCL Platform, the “platform-specific data” stored in `ContextImpl` is of type `OpenCLPlatform::PlatformData`, which is declared in `OpenCLPlatform.h`. The most important field of this class is `contexts`, which is a vector of `OpenCLContexts`. (There is one `OpenCLContext` for each device you are using. The most common case is that you are running everything on a single device, in which case there will be only one `OpenCLContext`. Parallelizing computations across multiple devices is not discussed here.) The `OpenCLContext` stores most of the important information about a simulation: positions, velocities, forces, an `OpenCLCommandQueue` used for executing kernels, workspace buffers of various sorts, etc. It provides many useful methods for compiling and executing kernels, clearing and reducing buffers, and so on. It also provides access to three other important objects: the `OpenCLIntegrationUtilities`, `OpenCLNonbondedUtilities`, and `OpenCLBondedUtilities`. These are discussed below.

Allocation of device memory is generally done through the `OpenCLArray` class. It takes care of much of the work of memory management, and provides a simple interface for transferring data between host and device memory.

Every kernel is specific to a particular `OpenCLContext`, which in turn is specific to a particular `OpenMM::Context`. This means that kernel source code can be customized for a

particular simulation. For example, values such as the number of particles can be turned into compile-time constants, and specific versions of kernels can be selected based on the device being used or on particular aspects of the system being simulated. `OpenCLContext::createProgram()` makes it easy to specify a list of preprocessor definitions to use when compiling a kernel.

The normal way to execute a kernel is by calling `executeKernel()` on the `OpenCLContext`. It allows you to specify the total number of work-items to execute, and optionally the size of each work-group. (If you do not specify a work-group size, it uses 64 as a default.) The number of work-groups to launch is selected automatically based on the work-group size, the total number of work-items, and the number of compute units in the device it will execute on.

5.2 Computing Forces

When forces are computed, they are stored in multiple buffers. This is done to enable multiple work-items or work-groups to compute forces on the same particle at the same time; as long as each one writes to a different buffer, there is no danger of race conditions. At the start of a force calculation, all forces in all buffers are set to zero. Each Force is then free to add its contributions to any or all of the buffers. Finally, the buffers are summed to produce the total force on each particle.

The size of each buffer is equal to the number of particles, rounded up to the next multiple of 32. Call `getPaddedNumAtoms()` on the `OpenCLContext` to get that number. The actual force buffers are obtained by calling `getForceBuffers()`. The first n entries (where n is the padded number of atoms) represent the first force buffer, the next n represent the second force buffer, and so on. More generally, the i 'th force buffer's contribution to the force on particle j is stored in element $i * \text{context.getPaddedNumAtoms()} + j$.

Depending on the device, a buffer may also be created that stores contributions to the forces in 64 bit fixed point format. On devices that support atomic operations on 64 bit integers in global memory, this can be a more efficient way of accumulating forces than using a large number of force buffers.

The potential energy is also accumulated in a set of buffers, but this one is simply a list of floats. All of them are set to zero at the start of a computation, and they are summed at the end of the computation to yield the total energy.

The OpenCL implementation of each Force object should define a subclass of `OpenCLForce`, and register an instance of it by calling `addForce()` on the `OpenCLContext`. This serves two purposes:

1. It reports how many force buffers are required when calculating this particular Force. The `OpenCLContext` sets the size of its force buffer array based on the largest number of buffers required by any Force.
2. It implements methods for determining whether particular particles or groups of particles are identical. This is important when reordering particles, and is discussed below.

5.3 Nonbonded Forces

Computing nonbonded interactions efficiently is a complicated business in the best of cases. It is even more complicated on a GPU. Furthermore, the algorithms must vary based on the type of processor being used, whether there is a distance cutoff, and whether periodic boundary conditions are being applied.

The `OpenCLNonbondedUtilities` class tries to simplify all of this. To use it you need provide only a piece of code to compute the interaction between two particles. It then takes responsibility for generating a neighbor list, looping over interacting particles, loading particle parameters from global memory, and writing the forces and energies to the appropriate buffers. All of these things are done using an algorithm appropriate to the processor you are running on and high level aspects of the interaction, such as whether it uses a cutoff and whether particular particle pairs need to be excluded.

Of course, this system relies on certain assumptions, the most important of which is that the Force can be represented as a sum of independent pairwise interactions. If that is not the

case, things become much more complicated. You may still be able to use features of `OpenCLNonbondedUtilities`, but you cannot use the simple mechanism outlined above. That is beyond the scope of this guide.

To define a nonbonded interaction, call `addInteraction()` on the `OpenCLNonbondedUtilities`, providing a block of OpenCL source code for computing the interaction. This block of source code will be inserted into the middle of an appropriate kernel. At the point where it is inserted, various variables will have been defined describing the interaction to compute:

- `atom1` and `atom2` are the indices of the two interacting particles.
- `r`, `r2`, and `invR` are the distance r between the two particles, r^2 , and $1/r$ respectively.
- `isExcluded` is a bool specifying whether this pair of particles is marked as an excluded interaction. (Excluded pairs are not skipped automatically, because in some cases they still need to be processed, just differently from other pairs.)
- `posq1` and `posq2` are float4s containing the positions (in the xyz fields) and charges (in the w fields) of the two particles.
- Other per-particle parameters may be specified, as described below.

The following preprocessor macros will also have been defined:

- `NUM_ATOMS` is the total number of particles in the system.
- `PADDED_NUM_ATOMS` is the padded number of particles in the system.
- `USE_CUTOFF` is defined if and only if a cutoff is being used
- `USE_PERIODIC` is defined if and only if periodic boundary conditions are being used.
- `CUTOFF_SQUARED` is the square of the cutoff distance (but only defined if a cutoff is being used).

Finally, two output variables will have been defined:

- You should add the energy of the interaction to `tempEnergy`.

- You should add the derivative of the energy with respect to the inter-particle distance to `dEdR`.

You can also define arbitrary per-particle parameters by calling `addParameter()` on the `OpenCLNonbondedUtilities`. You provide an array in device memory containing the set of values, and the values for the two interacting particles will be loaded and stored into variables called `<name>1` and `<name>2`, where `<name>` is the name you specify for the parameter. Note that nonbonded interactions are not computed until after `calcForcesAndEnergy()` has been called on every `ForceImpl`, so it is possible to make the parameter values change with time by modifying them inside `calcForcesAndEnergy()`. Also note that the length of the array containing the parameter values must equal the *padded* number of particles in the system.

Finally, you can specify arbitrary other memory objects that should be passed as arguments to the interaction kernel by calling `addArgument()`. The rest of the kernel ignores these arguments, but you can make use of them in your interaction code.

Consider a simple example. Suppose we want to implement a nonbonded interaction of the form $E=k_1k_2r^2$, where k is a per-particle parameter. First we create a parameter as follows

```
nb.addParameter(OpenCLNonbondedUtilities::ParameterInfo(
    "kparam", "float", 1, sizeof(cl_float),
    kparam->getDeviceBuffer()));
```

where `nb` is the `OpenCLNonbondedUtilities` for the context. Now we call `addInteraction()` to define an interaction with the following source code:

```
#ifdef USE_CUTOFF
if (!isExcluded && r2 < CUTOFF_SQUARED) {
#else
if (!isExcluded) {
#endif
    tempEnergy += kparam1*kparam2*r2;
```

```

    dEdR += 2*kparam1*kparam2*r;
}

```

An important point is that this code is executed for every pair of particles in the *padded* list of atoms. This means that some interactions involve padding atoms, and should not actually be included. You might think, then, that the above code is incorrect and we need another check to filter out the extra interactions:

```

if (atom1 < NUM_ATOMS && atom2 < NUM_ATOMS)

```

This is not necessary in our case, because the `isExcluded` flag is always set for interactions that involve a padding atom. If our force did not use excluded interactions (and so did not check `isExcluded`), then we would need to add this extra check. Self interactions are a similar case: we do not check for `(atom1 == atom2)` because the exclusion flag prevents them from being processed, but for some forces that check is necessary.

5.4 Bonded Forces

Just as `OpenCLNonbondedUtilities` simplifies the task of creating nonbonded interactions, `OpenCLBondedUtilities` simplifies the process for many types of bonded interactions. A “bonded interaction” means one that is applied to small, fixed groups of particles. This includes bonds, angles, torsions, etc. The important point is that the list of particles forming a “bond” is known in advance and does not change with time.

Using `OpenCLBondedUtilities` is very similar to the process described above. You provide a block of OpenCL code for evaluating a single interaction. This block of code will be inserted into the middle of a kernel that loops over all interactions and evaluates each one. At the point where it is inserted, the following variables will have been defined describing the interaction to compute:

- `index` is the index of the interaction being evaluated.
- `atom1, atom2, ...` are the indices of the interacting particles.

- `pos1`, `pos2`, ... are `float4`s containing the positions (in the xyz fields) of the interacting particles.

A variable called `energy` will have been defined for accumulating the total energy of all interactions. Your code should add the energy of the interaction to it. You also should define `float4` variables called `force1`, `force2`, ... and store the force on each atom into them.

As a simple example, the following source code implements a pairwise interaction of the form $E=r^2$:

```
float4 delta = pos2-pos1;
energy += delta.x*delta.x + delta.y*delta.y + delta.z*delta.z;
float4 force1 = 2.0f*delta;
float4 force2 = -2.0f*delta;
```

To use it, call `addInteraction()` on the Context's `OpenCLBondedUtilities` object. You also provide a list of the particles involved in every bonded interaction.

Exactly as with nonbonded interactions, you can call `addArgument()` to specify arbitrary memory objects that should be passed as arguments to the interaction kernel. These might contain per-bond parameters (use `index` to look up the appropriate element) or any other information you want.

5.5 Reordering of Particles

Nonbonded calculations are done a bit differently in the OpenCL Platform than in most CPU based codes. In particular, interactions are computed on blocks of 32 sequential particles at a time (which is why the number of particles needs to be padded to bring it up to a multiple of 32), and the neighbor list actually lists pairs of *blocks*, not pairs of individual particles, that are close enough to interact with each other.

This only works well if sequential particles tend to be close together so that blocks are spatially compact. This is generally true of particles in a macromolecule, but it is not true for solvent molecules. Each water molecule, for example, can move independently of other water molecules, so particles that happen to be sequential in whatever order the molecules were defined in need not be spatially close together.

The OpenCL Platform addresses this by periodically reordering particles so that sequential particles are close together. This means that what the OpenCL Platform calls particle i need not be the same as what the System calls particle i .

This reordering is done frequently, so it must be very fast. If all the data structures describing the structure of the System and the Forces acting on it needed to be updated, that would make it prohibitively slow. The OpenCL Platform therefore only reorders particles in ways that do not alter any part of the System definition. In practice, this means exchanging entire molecules; as long as two molecules are truly identical, their positions and velocities can be exchanged without affecting the System in any way.

Every Force can contribute to defining the boundaries of molecules, and to determining whether two molecules are identical. This is done through the `OpenCLForceInfo` it adds to the `OpenCLContext`. It can specify two types of information:

1. Given a pair of particles, it can say whether those two particles are identical (as far as that Force is concerned). For example, a Force object implementing a Coulomb force would check whether the two particles had equal charges.
2. It can define *particle groups*. The OpenCL Platform will ensure that all the particles in a group are part of the same molecule. It also can specify whether two groups are identical to each other. For example, in a Force implementing harmonic bonds, each group would consist of the two particles connected by a bond, and two groups would be identical if they had the same spring constants and equilibrium lengths.

5.6 Integration Utilities

The `OpenCLContext`'s `OpenCLIntegrationUtilities` provides features that are used by many integrators. The two most important are random number generation and constraint enforcement.

If you plan to use random numbers, you should call `initRandomNumberGenerator()` during initialization, specifying the random number seed to use. Be aware that there is only one random number generator, even if multiple classes make use of it. If two classes each call `initRandomNumberGenerator()` and request different seeds, an exception will be thrown. If they each request the same seed, the second call will simply be ignored.

For efficiency, random numbers are generated in bulk and stored in an array in device memory, which you can access by calling `getRandom()`. Each time you need to use a block of random numbers, call `prepareRandomNumbers()`, specifying how many values you need. It will register that many values as having been used, and return the index in the array at which you should start reading values. If not enough unused values remain in the array, it will generate a new batch of random values before returning.

To apply constraints, simply call `applyConstraints()`. For numerical accuracy, the constraint algorithms do not work on particle positions directly, but rather on the *displacements* taken by the most recent integration step. These displacements must be stored in an array which you can get by calling `getPosDelta()`. That is, the constraint algorithms assume the actual (unconstrained) position of each particle equals the position stored in the `OpenCLContext` plus the delta stored in the `OpenCLIntegrationUtilities`. It then modifies the deltas so that all distance constraints are satisfied. The integrator must then finish the time step by adding the deltas to the positions and storing them into the main position array.