

ExaFMM

Fast multipole method software aiming for exascale systems

User's Manual

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Revision History

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1 Introduction

This manual describes ExaFMM, an open-source implementation of the fast multipole method (FMM) written in the C++ language. For maximum freedom of use, ExaFMM is distributed under the MIT License. Please note that you must give proper attribution in all derived works.

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ExaFMM was released for the first time in November 2011. It is freely downloadable from <https://bitbucket.org/exafmm/exafmm>, and we welcome correspondence with potential users or those who wish to extend it for specific purposes.

Please write to exafmm@barbagroup.bu.edu with any comments or questions.

2 Download & Install

2.1 Download

If you have mercurial, simply do

```
hg clone https://exafmm@bitbucket.org/exafmm/exafmm
```

if not, you may download the source from <https://bitbucket.org/exafmm/exafmm/overview>. There is a link on the right that says "get source".

2.2 Install

To use ExaFMM as a black box library do

```
cd exafmm
./configure
make lib
```

set your library path to `exafmm/wrapper`, and add ```-lexafmm``` to your library flags. An MPI C++ compiler is required. If you want to run without MPI, open `exafmm/Makefile` and change the variable `CXX` from `mpicxx` to `g++`. ExaFMM is designed so that the parallel portion of the code is an extension of the serial portion of the code using class inheritance. Therefore, all the serial parts of the code function with or without MPI, and there is no need for a separate serial version.

The minimum installation of ExaFMM does not depend on any external libraries. However, it does provide visualization routines that uses VTK (downloadable from <http://www.vtk.org/VTK/resources/software.html>) if the user chooses to use them, but the FMM itself will function without VTK.

3 Test suite

3.1 Unit tests

All unit tests can be found in the directory `unit_test`. The `Makefile` in this directory contains the commands to run all the unit tests. For a very quick demonstration of an N -body calculation for $N = 10,000$ bodies do:

```
cd unit_test
make serialrun
```

The result will be indicated by the output text. If you encounter any errors, please send the full output text to exafmm@barbagroup.bu.edu.

Here is the full list of unit tests, and a short description of what they do.

- index:** Assigns Morton index to 10^7 bodies, and sorts the bodies according to the index. If VTK is available, it will plot the group of bodies according to the index, with a toggle bar that controls the group to be shown.
- topdown:** Constructs a tree structure top-down for 10^7 bodies. If VTK is available, it will plot the group of bodies according to the leaf index, with a toggle bar that controls the group to be shown.
- bottomup:** Same as topdown, except the tree construction is bottom-up.
- kernel:** Checks the convergence of kernels by doing P2M, M2M, M2L, L2L, L2P, and M2P translations for a single pair of particles. The distance between the particles and the cells that contain them is varied and the

difference between a direct summation is monitored. The default is the Laplace kernel but changing `D.kernelName` on line 14 allows you to check different kernels, as is the case with many other unit tests.

- serialrun:** Compares serial FMM with direct summation for 10^4 bodies.
- unsort:** Same as `serialrun` except it sorts back the bodies to their initial order. (FMM reorders the bodies according to the Morton index).
- Nserial:** Same as `serialrun` except it varies the number of bodies from 10^4 to 10^7 and checks the $\mathcal{O}(N)$ scaling.
- direct_gpu:** Compares the serial direct summation on a GPU and CPU.
- mpi:** Prints the hostname, rank, and size of all MPI processes in order.
- check_gpus:** Compares the parallel direct summation on many GPUs and CPUs. Prints the MPI rank, GPU device ID, and Error between the CPU and GPU for every MPI process.
- shift:** Verification of communication routine `shiftBodies` for sending bodies to the next process in round-robin fashion. This allows the source bodies to be shifted to the next process so that an all-pairs interaction can be calculated on distributed memory systems without increasing storage requirements. If VTK is available, the same `shiftBodies` routine will be used to plot bodies on all processes without storing all of them in one process. The toggle bar can be used to show bodies on different processes.
- nth_element:** Verification of parallel `nth_element` routine. The `nth_element` finds the n^{th} element of a sorted vector without actually sorting the vector. It is an efficient method for finding the median. The parallel `nth_element` finds the global n^{th} element without performing an actual parallel sort. If VTK is available, it will plot the bodies on both sides of the `nth_element` and toggles between the two. Only the bodies on MPI rank 0 will be plotted.
- bisection:** Verification of recursive multi-section partitioning. If VTK is available, it will plot each of the partitioned domains (one per process), and toggle between all domains. It will partition the domain into equal number of bodies for any number of processes.
- let:** Verification of local essential tree (LET). Partitions the domain using multi-section, performs a parallel tree construction, and communicates the LET. If VTK is available, it will plot the bodies and cells on each process, and toggle between the bodies and cells on every process.
- parallelrun:** Parallel version of `serialrun`. Compares parallel FMM with direct summation for 10^4 bodies. If VTK is available it will plot the same thing as `let`, but this time for the actual bodies and cells that were used in the FMM calculation.

- unpartition:** Same as `parallelrun`, except it (un)partitions the bodies back to their original distribution among processes, and (un)sorts them back to their original order within each process at the end.
- ijparallelrun:** Same as `unpartition`, except it has different distribution for target bodies and source bodies.
- Nparallel:** Parallel version of `Nserial`. Varies the number of bodies from 10^4 to 10^7 and checks the $\mathcal{O}(N)$ scaling of the parallel FMM.
- skip_tree:** Verification of tree construction skipping. Same as `parallelrun` except it does two FMM calculations, and in the second one it skips the tree construction and uses the tree structure from the first FMM.
- overlap_comm:** Verification of communication overlapping. Same as `skip_tree`, except it overlaps the communication with the computation of the local tree using OpenMP.
- vtk:** Verification of VTK plotter. Plots two groups of bodies, with a bar that toggles between them.
- cuprintf:** Verification of `cuprintf` --a printf routine that prints from inside kernels that works even on old GPUs.

3.2 Regression tests

There are no regression tests at the moment.

4 Algorithm description

The fast multipole method (FMM) is an algorithm for accelerating computations of the form:

$$f(y_j) = \sum_{i=1}^N c_i \mathbb{K}(y_j, x_i) \quad (1)$$

Such a computation may represent a field value evaluated at point y_j , where the field is generated by the influence of sources located at the set of centers $\{x_i\}$. The sources are often associated with particle-type objects, such as stellar masses, or charged particles. The evaluation of the field at the centers themselves, therefore, represents the well-known N -body problem. In summary: $\{y_j\}$ is a set of evaluation points, $\{x_i\}$ is a set of source points with weights given by c_i , and $\mathbb{K}(y, x)$ is the kernel that governs the interactions between evaluation and source particles. The objective is to obtain the field f at all the evaluation points, which requires in principle $\mathcal{O}(N^2)$ operations if both sets of points have N elements. Fast algorithms aim at obtaining f approximately with a reduced operation count, ideally $\mathcal{O}(N)$.

The FMM works by approximating the influence of a cluster of particles by a single collective representation, under the assumptions that the influence of particles becomes weaker as the evaluation point is further away, *i.e.*, the kernel $\mathbb{K}(y, x)$ decays as $|x - y|$ increases, and that the approximations are used to evaluate far distance interactions. To accomplish this, the FMM hierarchically decomposes the computational domain and then it represents the influence of sets of particles by a single approximate value. The hierarchical decomposition breaks up the domain at increasing levels of refinement, and for each level it identifies a near and far sub-domain. By using the hierarchical decomposition, the far field can be reconstructed as shown in Figure 1.

Using the computational domain decomposition, the sum in Equation (1) is decomposed as

$$f(y_j) = \sum_{l=1}^{N_{near}} c_l \mathbb{K}(y_j, x_l) + \sum_{k=1}^{N_{far}} c_k \mathbb{K}(y_j, x_k) \quad (2)$$

where the right-most sum of (2), representing the far field, is evaluated approximately and efficiently.

We now need to introduce the following terminology with respect to the mathematical tools used to agglomerate the influence of clusters of particles:

Multipole Expansion (ME): a series expansion truncated after p terms which represents the influence of a cluster of particles, and is valid at distances large with respect to the cluster radius.

Local Expansion (LE): a truncated series expansion, valid only inside a sub-domain, which is used to efficiently evaluate a group of MEs.

In other words, the MEs and LEs are series (*e.g.* Taylor series) that converge in different sub-domains of space. The center of the series for an ME is the center of the cluster of source particles, and it only converges outside the cluster of particles. In the case of an LE, the series is centered near an evaluation point and converges locally.

As an example, consider a particle interaction problem with decaying kernels, where a cluster of particles far away from an evaluation point is 'seen' at the evaluation point as a 'pseudo-particle', and thus its influence can be represented by a single expression. For example, the gravitational potential of a galaxy far away can be expressed by a single quantity locally. Thus, by using the ME representing a cluster, the influence of that cluster can be rapidly evaluated at a point located far away ---as only the single influence of the ME needs to be evaluated, instead of the multiple influences of all the particles in the cluster. Moreover, for clusters of particles that are farther from the evaluation point, the pseudo-particle representing that cluster can be larger. This idea, illustrated in Figure ??, permits increased efficiency in the computation.

The introduction of an aggregated representation of a cluster of particles, via the multipole expansion, effectively permits a decoupling of the influence of the source particles from the evaluation points. This is a key idea, resulting in the factorization of the computations

of MEs that are centered at the same point, so that the kernel can be written as,

$$\mathbb{K}(x_i, y_j) = \sum_{m=0}^p a_m(x_i) f_m(y_j) \quad (3)$$

This factorization allows pre-computation of terms that can be re-used many times, thus increasing the efficiency of the overall computation. Similarly, the local expansion is used to decouple the influence of an ME from the evaluation points. A group of MEs can be factorized into a single LE so that one single evaluation can be used at multiple points locally. By representing MEs as LEs one can efficiently evaluate a group of clusters in a group of evaluation points.

4.1 Hierarchical space decomposition

In order to utilize the tools of MEs and LEs, a spatial decomposition scheme needs to be provided. In other words, for a complete set of particles, we need to find the clusters that will be used in conjunction with the MEs to approximate the far field, and the sub-domains where the LEs are going to be used to efficiently evaluate groups of MEs. This spatial decomposition is accomplished by a hierarchical subdivision of space associated to a tree structure (*quadtrees* structure in two dimensions, or an *octree* structure in three dimensions) to represent each subdivision. The nodes of the tree structure are used to define the spatial decomposition, and different scales are obtained by looking at different levels of the tree. A tree representation of the space decomposition allows us to express the decomposition independently of the number of dimensions of the space. Consider Figure ?? where a quadtree decomposition of the space is illustrated. The nodes of the tree at any given level cover the whole domain. The relations between nodes of the tree, represent spatial refinement. The domain covered by a parent box is further decomposed into smaller sub-domains by its child nodes. Thus, in the FMM the tree structure is used to hierarchically decompose the space and the hierarchical space decomposition is used to represent the near-field and far-field domains. As an example, consider Figure ?? where the near-field for the *black* colored box is represented by the light colored boxes, and the far-field is composed by the dark colored boxes.

5 Using ExaFMM in application programs

Here we discuss at a high level, how to interact with ExaFMM. This section is PENDING.

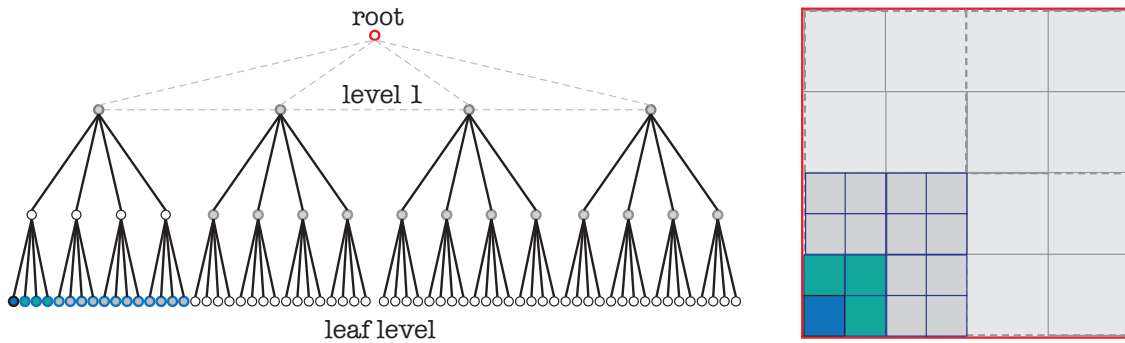


Figure 1: Tree structure associated to spatial decomposition.

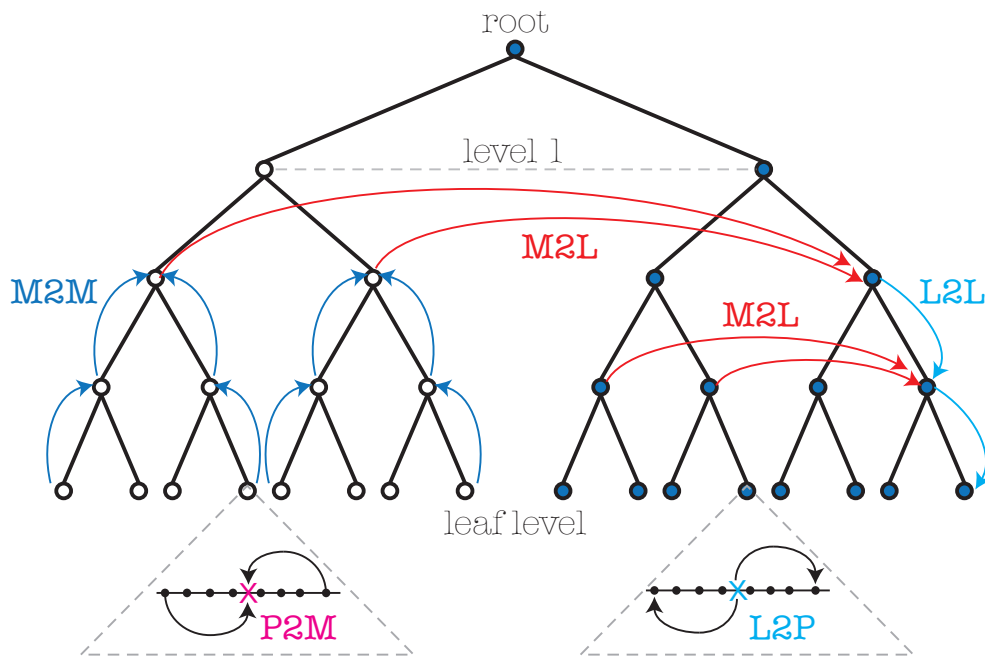


Figure 2: The sketch illustrates the *upward sweep* and the *downward sweep* stages on the tree. Each stage has been further decomposed into the following substages: *P2M*--transformation of particles into MEs (particle-to-multipole); *M2M*--translation of MEs (multipole-to-multipole); *M2L*--transformation of an ME into an LE (multipole-to-local); *L2L*--translation of an LE (local-to-local); *L2P*--evaluation of a LEs at particle locations (local-to-particle).