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Permalink
https://escholarship.org/uc/item/9m9634ss

ISBN
9781424446432

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Publication Date
2010-07-01

DOI
10.1109/IPDPS.2010.5470415

Peer reviewed
Optimizing and Tuning the Fast Multipole Method for State-of-the-Art Multicore Architectures

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Abstract

This work presents the first extensive study of single-node performance optimization, tuning, and analysis of the fast multipole method (FMM) on modern multicore systems. We consider single- and double-precision with numerous performance enhancements, including low-level tuning, numerical approximation, data structure transformations, OpenMP parallelization, and algorithmic tuning.

Among our numerous findings, we show that optimization and parallelization can improve double-precision performance by 25× on Intel’s quad-core Nehalem, 9.4× on AMD’s quad-core Barcelona, and 37.6× on Sun’s Victoria Falls (dual-sockets on all systems). We also compare our single-precision version against our prior state-of-the-art GPU-based code and show, surprisingly, that the most advanced multicore architecture (Nehalem) reaches parity in both performance and power efficiency with NVIDIA’s most advanced GPU architecture.

1 Introduction

This paper presents the first extensive study of single-node performance optimization, tuning, and analysis of the fast multipole method (FMM) [5] on state-of-the-art multicore processor systems. We target the FMM because it is broadly applicable to a variety of scientific particle simulations used to study electromagnetic, fluid, and gravitational phenomena, among others. Importantly, the FMM has asymptotically optimal time complexity with guaranteed approximation accuracy. As such, it is among the most attractive solutions for scalable particle simulation on future extreme scale systems. This study focuses on single-node performance since it is a critical building-block in scalable multi-node distributed memory codes and, moreover, is less well-understood.

2 Fast Multipole Method

This section provides an overview of the fast multipole method (FMM), summarizing the key components that are relevant to this study. For more in-depth algorithmic details, see Greengard, et al. [5, 16].
Given a system of $N$ source particles, with positions given by $\{y_1, \ldots, y_N\}$, and $N$ targets with positions $\{x_1, \ldots, x_N\}$, we wish to compute the $N$ sums,

$$f(x_i) = \sum_{j=1}^{N} K(x_i, y_j) \cdot s(y_j), \quad i = 1, \ldots, N \quad (1)$$

where $f(x)$ is the desired potential at target point $x$; $s(y)$ is the density at source point $y$; and $K(x, y)$ is an interaction kernel that specifies “the physics” of the problem. For instance, the single-layer Laplace kernel, $K(x, y) = \frac{1}{4\pi ||x-y||}$, might model electrostatic or gravitational interactions.

Evaluating these sums appears to require $O(N^2)$ operations. The FMM instead computes approximations of all of these sums in optimal $O(N)$ time with a guaranteed user-specified accuracy, where the desired accuracy changes the complexity constant [5].

The FMM is based on two key ideas: (i) a tree representation for organizing the points spatially; and (ii) fast approximate evaluation, in which we compute summaries at each node using a constant number of tree traversals with constant work per node.

We implement the kernel independent variant of the FMM, or KIFMM [16]. KIFMM has the same structure as the classical FMM [5]. Its main advantage is that it avoids the mathematically challenging analytic expansion of the kernel, instead requiring only the ability to evaluate the kernel. This feature of the KIFMM allows one to leverage our optimizations and techniques and apply them to new kernels and problems.

**Tree construction:** Given the input points and a user-defined parameter $q$, we construct an oct-tree $T$ (or quad-tree in 2-D) by starting with a single box representing all the points and recursively subdividing each box if it contains more than $q$ points. Each box (octant in 3-D or quadrant in 2-D) becomes a tree node whose children are its immediate sub-boxes. During construction, we associate with each node one or more neighbor lists. Each list has bounded constant length and contains (logical) pointers to a subset of other tree nodes. These are canonically known as the $U$, $V$, $W$, and $X$ lists. For example, every leaf box $B \in \text{leaves}(T)$ has a $U$ list, $U(B)$, which is the list of all leaves adjacent to $B$. Figure 1 shows a quad-tree example, where neighborhood list nodes for $B$ are are labeled accordingly.

Tree construction has $O(N \log N)$ complexity, and so the $O(N)$ optimality refers to the evaluation phase (below). However, tree construction is typically a small fraction of the total time; moreover, many applications build the tree periodically, thereby enabling amortization of this cost over several evaluations.

**Evaluation:** Given the tree $T$, evaluating the sums consists of six distinct computational phases: there is one phase for each of the $U$, $V$, $W$, and $X$ lists, as well as upward (up) and downward (down) phases. These phases involve traversals of $T$ or subsets of $T$. Rather than describe each phase in detail, which are well-described elsewhere [5, 15, 16], we summarize the main algorithmic characteristics of each phase in Table 1.

For instance, in the $U$ list phase, we traverse all leaf nodes, where for each leaf node $B \in \text{leaves}(T)$ we perform a direct evaluation between $B$ and each of the nodes $B' \in U(B)$. For each leaf $B$, this direct evaluation operates on $O(q)$ points and has a flop cost of $O(q^2)$ for each box. By contrast, the $V$-list operates on $O(q)$ points and performs $O(q \log q)$ flops for each box, and so has lower computational intensity. Generally, we expect the cost of the $U$-list evaluation phase to dominate other phases when $q$ is sufficiently large.

There are multiple levels of concurrency during evaluation: across phases (e.g., the upward and $U$-list phases can be executed independently), within a phase (e.g., each leaf box can be evaluated independently during the $U$-list phase), and within the per-octant computation (e.g., vectorizing each direct evaluation). In this paper, we consider intra-phase and per-octant parallelization only, essentially using OpenMP and manual SIMD parallelization, respectively. For the $Upward$ and $Downward$ phases, which both involve tree traversals, there is an obvious dependency between a parent and its child boxes. However, the children themselves are independent and can be computed concurrently with the amount of work per level increasing toward the leaves.

**Related work:** For parallel FMM, most of the recent work of which we are aware focuses on distributed memory codes with GPU-based acceleration [1, 6, 9, 12]. Indeed, the present study builds on our own state-of-the-art parallel 3-D KIFMM implementation, which uses MPI+CUDA [9]. However, these works have not yet considered conventional multicore acceleration and tun-
Table 1. Characteristics of the computational phases in KIFMM. \(N\) is the number of source particles, the number of boxes is \(M \sim N/q\) and \(p\) denotes the number of expansion coefficients. The user chooses \(p\) to trade-off time and accuracy, and may tune \(q\) to minimize time. \(^{1}\)Size is determined by the chosen accuracy, generally smaller than \(q\).

<table>
<thead>
<tr>
<th>Phase</th>
<th>Computational Complexity</th>
<th>Algorithmic Characteristics</th>
</tr>
</thead>
<tbody>
<tr>
<td>Upward</td>
<td>(O(Np + Mp^2))</td>
<td>postorder tree traversal, small (^{1}) (\matvec{s})</td>
</tr>
<tr>
<td>U-list</td>
<td>(O(27Nq))</td>
<td>direct computation as in Equation 1 ((\matvec{s}) on the order of (q))</td>
</tr>
<tr>
<td>V-list</td>
<td>(O(Mp^{3/2}\log p + 189Mp^{3/2}))</td>
<td>consists of small FFTs, pointwise vector multiplication (convolution)</td>
</tr>
<tr>
<td>X-list</td>
<td>(0) (\text{uniform distribution} \quad O(Nq)) (\text{non-uniform distribution})</td>
<td>matvecs</td>
</tr>
<tr>
<td>W-list</td>
<td>(0) (\text{uniform distribution} \quad O(Nq)) (\text{non-uniform distribution})</td>
<td>matvecs</td>
</tr>
<tr>
<td>Downward</td>
<td>(O(Np + Mp^2))</td>
<td>preorder tree traversal, small (^{1}) (\matvec{s})</td>
</tr>
</tbody>
</table>

In Section 5, we compare our multicore optimizations to this prior use of GPU acceleration, with the perhaps surprising finding that a well-tuned multicore implementation can match a GPU code. Coulaud, et al., propose Pthreads- and multithreaded BLAS-based multicore parallelization within node [3]. However, we use a larger set of optimizations and provide cross-platform performance and power analysis.

There are numerous non-GPU studies of single-core distributed memory FMM implementations [8,11] (see also references in Ying, et al. [15]), most based on the classical tree-based N-body framework of Warren and Salmon [14], including our own prior KIFMM work [9,15]. Researchers have considered a variety of data structures with attractive communication properties, again in the distributed context [7]. To our knowledge, the present study is the first to consider extensive multicore-centric optimizations, data structures, and cross-platform analysis.

For direct \(O(N^2)\) methods, tuning, and special-purpose hardware (e.g. MDGRAPE), see the references in related papers [2,10].

### 3 Experimental Setup

We explore FMM performance as we vary architecture, floating-point precision, and initial particle distribution. To facilitate comparisons to prior work, we select a commonly used kernel \(K\) (Laplace kernel in Section 2). The desired accuracy is fixed to a typical minimum setting that is also sensible for single-precision (yielding 4–7 decimal digits of accuracy). Moreover, because tuning can dramatically change the requisite number of floating-point operations, we define and defend our alternate performance metrics. These aspects are discussed in detail below.

#### 3.1 Architectures

This section summarizes the key differences, as they pertain to the FMM, among the three dual-socket multicore SMPs used in this study: Intel’s quad-core Xeon (Nehalem), AMD’s quad-core Opteron (Barcelona), and Sun’s chip-multithreaded, eight-core UltraSparc T2+ (Victoria Falls). Our final analysis references our prior GPU-only accelerated results [9]. The key parameters of these systems appear in Table 2.

**Basic microarchitectural approach:** Nehalem and Barcelona are x86, superscalar, out-of-order architectures with large per-thread caches and hardware prefetchers. Victoria Falls, by contrast, employs fine-grained chip multithreading (CMT) and smaller per-thread caches. Consequently, Victoria Falls requires the programmer to express roughly an order of magnitude more parallelism than the x86 systems in order to achieve peak performance or peak bandwidth. Luckily, there is ample fine-grained thread-level parallelism in the FMM.

**Computational peak:** The two x86-based systems have similar peak floating-point performance, but 4× higher than Victoria Falls partly due to the SIMD units on x86. SIMD enables up to to 4 flops (multiply and add) per cycle per core in double-precision (DP) and 8 in single-precision (SP). Because the FMM has high computational intensity in at least one of its major phases (U-list), we may expect superior performance on the x86 systems compared to Victoria Falls.

Unfortunately, many kernels \(K(x,y)\) also require square root and divide operations, which on all three systems are not pipelined and therefore are extremely slow. For example, on Nehalem, double-precision divide and square root run at 0.266 GFlop/s (5% of peak multiply/add performance) and 0.177 GFlop/s (3%), respectively. To address this deficiency, both x86 systems (but not Victoria Falls) have a low latency and pipelined single-precision approximate reciprocal square-root op-
Table 2. Architectural Parameters. All power numbers, save the GPU, we obtained using a
digital power meter. *reciprocal square-root approximate. †shared among cores on a socket.
‡max server power (of which the 2 active CPUs consume 160W) plus max power for two GPUs.

<table>
<thead>
<tr>
<th></th>
<th>Intel X5550 (Nehalem)</th>
<th>AMD 2356 (Barcelona)</th>
<th>Sun T5140 (Victoria Falls)</th>
<th>NVIDIA T10P (S1070)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Architecture</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Frequency (GHz)</strong></td>
<td>2.66 GHz</td>
<td>2.30 GHz</td>
<td>1.166 GHz</td>
<td>1.44 GHz</td>
</tr>
<tr>
<td><strong>Sockets</strong></td>
<td>2</td>
<td>2</td>
<td>2</td>
<td>2 (+2 CPUs)</td>
</tr>
<tr>
<td><strong>Cores/Socket</strong></td>
<td>4</td>
<td>4</td>
<td>8</td>
<td>8 (GPU)</td>
</tr>
<tr>
<td><strong>Threads/Core</strong></td>
<td>2</td>
<td>4</td>
<td>8</td>
<td>8 (GPU)</td>
</tr>
<tr>
<td><strong>SIMD (DP, SP)</strong></td>
<td>2-way 4-way</td>
<td>2-way 4-way</td>
<td>1 1</td>
<td>1 8-way</td>
</tr>
<tr>
<td><strong>GFlop/s (DP, SP)</strong></td>
<td>85.33 170.6</td>
<td>73.60 146.2</td>
<td>18.66 18.66</td>
<td>N/A 2073.6</td>
</tr>
<tr>
<td><em><em>rsqrt/s</em> (DP, SP)</em>*</td>
<td>0.853 42.66</td>
<td>0.897 73.60</td>
<td>2.26 —</td>
<td>N/A 172.8</td>
</tr>
<tr>
<td><strong>L1/L2/L3 cache local store</strong></td>
<td>32/256/8192 KB</td>
<td>64/512/2048 KB</td>
<td>8/4096 KB</td>
<td>—</td>
</tr>
<tr>
<td><strong>DRAM Bandwidth</strong></td>
<td>51.2 GB/s</td>
<td>21.33 GB/s</td>
<td>64.0 GB/s</td>
<td>16 KB</td>
</tr>
<tr>
<td><strong>Power</strong></td>
<td>375W</td>
<td>350W</td>
<td>610W</td>
<td>325W+400W‡</td>
</tr>
<tr>
<td><strong>Compiler</strong></td>
<td>icc 10.1</td>
<td>icc 10.1</td>
<td>cc 5.9</td>
<td>nvecc 2.2</td>
</tr>
</tbody>
</table>

KIFMM on the NCSA Lincoln Cluster [9], where each node is a dual-socket × quad-core Xeon 5410 (Harpertown) CPU server paired with two NVIDIA T10P GPUs. We use the GPUs only for control, and run all phases (except tree construction) on the GPUs. That is, there is one MPI process on each socket, and each process is assigned to one GPU; processes communicate via message passing and to their respective GPUs via PCIe. For our energy comparisons, we bound power using two configurations: aggregate peak GPU power plus zero CPU power, and aggregate peak GPU power plus the peak CPU power. With 12× the compute capacity and over 5× the bandwidth, one would naively expect the GPU implementation to considerably outperform all other platforms.

3.2 Kernel, Precision, and Accuracy

In this section, we describe the interaction kernel, precision, and desired accuracy, and their implications for implementation and optimization.

**Kernel:** Following prior work, we use the single-layer Laplacian kernel (Section 2) owing to its widely-recognized importance [16].

**Precision:** We consider both single and double-precision in our study. Single-precision is an interesting case for a variety of reasons. First, an application may have sufficiently low accuracy requirements, due to uncertainty in the input data or slow time-varying behavior. In this case, using single-precision can yield significant storage and performance benefits. Next, on current x86 architectures, SIMD instructions are 2-wide in double precision and 4-wide in single. However, forthcoming x86 Advanced Vector Extensions (AVX) will double these widths. As such, using single precision SIMD on today’s Nehalem is a proxy for double precision performance on tomorrow’s Sandy Bridge. Finally, current architectures provide fast reciprocal square-root methods in single-precision, but not double. By exploring the benefits in single, we may draw conclusions as to potential benefit future architectures may realize by implementing equivalent support in double.

**Accuracy:** One of the inputs to FMM is numerical accuracy desired in the final outcome, expressed as the desired “size” of the multipole expansion. In our experiments, we choose the desired accuracy to deliver the equivalent of 6 decimal digits in double-precision and 4 digits in single. We verify the delivered accuracy of our all of our naïve, optimized, parallel, and tuned implementations.

3.3 Particle Distributions

We examine two different particle distributions namely, a spatially uniform and a spatially non-uniform (elliptical or ellipsoidal) distribution. The uniform case is analyzed extensively in prior work; the non-uniform case is where we expect tree-based methods to deliver performance and accuracy advantages over other nu-
merical paradigms (e.g., particle-mesh methods). In both cases, our test problems use 4 million source particles plus an additional 4 million target particles.

**Uniform**: In this case, we distribute points uniformly at random within the unit cube. In 3D, for boxes not on the boundary, the U-list (neighbor list) contains 27 boxes and the V-list (interaction list) contains 189 boxes. The X- and W-lists are empty since the neighbors of a box are adjacent boxes in the same level. Thus, the time spent in the various list computations will differ from the non-uniform case and tuning will favor a different value for $q$, the maximum points per box.

**Elliptical**: In this case, particles are angularly-uniformly (in spherical coordinates) distributed on the surface of an ellipsoid with an aspect ratio 1:1:4. For an uniform distribution, a regular octree is constructed. However, the elliptical case requires an adaptively refined octree. As such, the depth of the computation tree could be quite large, resulting in high tree construction times as shown in Figure 5.

### 3.4 Performance Metrics

Since our optimizations and tuning of $q$ can dramatically change the total number of floating-point operations, we use time-to-solution (in seconds) as our primary performance metric rather than GFlop/s. In our final comparison, we present relative performance (evaluations per second), where higher numbers are better.

This choice has ramifications when assessing scalability. In particular, rather than examining GFlop/s/core or GFlop/s/thread to assess per-core (or per-thread) performance, we report thread-seconds: that is, the product of execution time by the number of threads. When it comes to energy efficiency, we present the ratio relative to the optimized and parallelized Nehalem energy efficiency (evaluations per Joule).

### 4 Optimizations

We applied numerous optimizations to the various computational phases (Section 2). Beyond optimizations traditionally subsumed by compilers, we apply numerical approximations, data structure changes, and tuning of algorithmic parameters. Table 3 summarizes our optimizations and their applicability to the FMM phases and our architectures. Note that not all optimizations apply to all phases.

Figure 2 presents the cumulative benefit as each optimization is successively applied to the serial reference KIFMM implementation [16]. We will refer to this figure repeatedly as we describe each optimization. If an optimization has associated tuning parameters (e.g. unrolling depth), we tune it empirically.

#### 4.1 SIMDization

We found it necessary to apply SIMD vectorization manually, as the compiler was unable to do so. All Laplacian kernel evaluations and point-wise matrix multiplication (in the V-list) are implemented using SSE intrinsics; specifically, in double-precision, we use SSE instructions like addpd, mulpd, subpd, divpd, and sqrtpd. Note that the Laplacian kernel performs 10 flops (counting each operation as 1 flop) per pairwise interaction, and includes both a square-root and divide.

In single-precision on x86, there is a fast (pipelined) approximate reciprocal square-root instruction: rsqrtps. As such, with sufficient instruction and data-level parallelism, we may replace the traditional scalar fsqrt/fdiv combination with not simply a rsqrtps/divps combination, but entirely with one rsqrtps. Doing so enables four reciprocal square-root operations per cycle without compromising our particular accuracy setting.

Figure 2 shows the speedup from SIMDization. The top three figures in Figure 2 use an uniform particle distribution and the bottom three use an elliptical distribution. SIMD nearly doubles Nehalem performance for all kernels except V-list, where FFTW (see Section 4.5) is already SIMDized. The benefit on Barcelona was much smaller (typically less than 50%), which we will investigate in future work. As there are no double precision SIMD instructions in SPARC/Victoria Falls, SIMD related optimizations are not applicable.

#### 4.2 Fast Reciprocal Square Root

A conventional double-precision SIMDized code would perform the reciprocal square-root operation using the intrinsics sqrtpd and divpd as above. Unfortunately, these instructions have long latencies (greater than 20 cycles) and cannot be pipelined, thus limiting performance. As we have abundant
instruction-level parallelism, we can exploit x86’s fast single-precision reciprocal square-root instruction to accelerate the double-precision computations [10]. That is, we replace the \texttt{sqrtpd/divpd} combination with the triplet, \texttt{cvtpd2ps (convert double to single)/rsqrtps/cvtps2pd} (single to double). To attain the desired accuracy, we apply an additional Newton-Raphson refinement iteration. This approach requires more floating-point instructions, but they are low latency and can be pipelined.

Figure 2 shows that the Newton-Raphson approach improves Nehalem performance by roughly 100% over SIMD. Since there are relatively few kernel evaluations in the V-list, we don’t see an appreciable benefit. Surprisingly, the benefit on Barcelona is relatively modest; the cause is still under investigation.

4.3 Structure-of-Arrays Layout

Our reference implementation uses an array-of-structures (AOS) data structure where all components of a point are stored contiguously in memory (e.g. \(x_1, y_1, z_1, x_2, y_2, z_2, ..., x_n, y_n, z_n\)). This layout is not SIMD-friendly as it requires a reduction across every point and unrolling the inner loop twice (or 4 times in single-precision).

Instead, we explore using the structure-of-arrays (SOA) or structure splitting layout [17] in which the components are stored in separate arrays. This transformation simplifies SIMDization since we can load two (or four) components into separate SIMD registers using a single instruction.

Changing the data layout further improved the overall U-list performance on Nehalem up to 300% speedup over the reference. The transformation does not affect the V-list phase due to its relatively low computational intensity. Moreover, the data layout change substantially improved Barcelona performance on most phases.

Unfortunately, the data layout change increased tree construction time due to lack of spatial locality. This tradeoff (dramatically reduced computational phase time for slightly increased tree construction time) is worthwhile if tree construction time is small compared to the total evaluation execution time.
4.4 Matrix-free calculations

To use tuned vendor BLAS routines, our reference code explicitly constructs matrices to perform matrix-vector multiplies (matvecs), as done by others [3]. However, we can apply what is essentially interprocedural loop fusion to eliminate this matrix, instead constructing its entries on-the-fly and thereby reducing the cache working set and memory traffic.

For example, recall that the U-list performs a direct evaluation like Equation 1, which is a matvec, between two leaf boxes. Rather than explicitly constructing the kernel matrix $K$ and performing the matvec, we can fuse the two steps and never store this matrix, reducing the memory traffic from $O(q^2)$ to $O(q)$, if the maximum points per box is $q$. The idea applies to the $Up$(leaf), $W$-list, $X$-list, and $Down$(leaf) phases as well; the matrices arise in a different way, but the principle is the same.

As seen in Figure 2, this technique improves Nehalem performance by an additional 25%, and often improved Barcelona and Victoria Falls performance by better than 40%. Cumulatively including this optimization (aside from V-list) improved Nehalem performance by more than 400%, Barcelona by more than 150%, and Victoria Falls by 40%.

4.5 FFTW

In our KIFMM implementation, the V-list phase consists of (i) small forward and inverse FFTs, once per source/target box combination; and (ii) pointwise multiplication $I$ times for each target box, where $I$ is the number of source boxes in said target box’s V-list.

Local FFTs are performed using FFTW [4]. Typically, one executes an FFTW plan for the array with which the plan was created using the function `fftw_execute`. As the sizes and strides of the FFTs in V-list are not only quite small but are also identical, we may create a single plan and reuse it for multiple FFTs, using the `fftw_execute_dft_r2c` function. We ensure alignment by creating the plan with the `FFTW_UNALIGNED` flag coupled with an aligned `malloc()`.

FFTW only benefits V-list computations. Nevertheless, on both x86 machines, FFTW substantially improved V-list performance for elliptical distributions. The benefit on uniform distributions was less dramatic since the relative time spent in the V-list tends to be smaller. Unfortunately, Victoria Falls saw little benefit from the plan reuse within FFTW; this will be addressed in future work.

4.6 Tree Construction

There are numerous studies of parallel tree construction [1,7,13]. In this paper, we focus on accelerating the evaluation phases of FMM for two main reasons. First, tree construction initially constituted a small fraction of execution time. Second, in many real simulation contexts, particle dynamics may be sufficiently slow that tree reconstruction can be amortized. Future work will involve parallelization of tree construction keeping in mind both uniform and non-uniform distributions.

4.7 Parallelization and Tuning

After applying serial optimization, we parallelize all phases via OpenMP. We apply inter-box parallelization for all phases except $Up$ward and $Down$ward. That is, we assign a chunk of leaf boxes to each thread and exploit parallelism within each phase. For $Up$ward and $Down$ward which has dependencies across the levels of the tree, we exploit the concurrency at each level. By convention, we exploit multiple sockets, then multiple cores, and finally threads within a core.

Algorithmically, the FMM is parameterized by the maximum number of particles per box, $q$. As $q$ grows, the U-list phase quickly increases in cost even as other phases become cheaper as the tree height shrinks; this dependence is non-trivial to predict, particularly for highly non-uniform distributions. We exhaustively tune $q$ for each implementation on each architecture, as discussed in Section 5. Auto-tuning $q$ will be the subject of our future work.

5 Performance Analysis

The benefits of optimization, threading, and tuning are substantial. When combined, these methods delivered speedups of $25\times$, $9.4\times$, and $37.6\times$ for Nehalem, Barcelona, and Victoria Falls, respectively, in double-precision for the uniform distribution; and $16\times$, $8\times$, and $24\times$, respectively, for the elliptical case. In this section we first tune our parallel implementation for the FMM’s key algorithmic parameter, $q$, the maximum particles per box. We then analyze the scalability for each architecture. Finally, we compare the performance and energy efficiency among architectures.

5.1 Tuning particles per box

Figure 3(a) presents the FMM execution time as a function of optimization, parallelization, and particles per box $q$ on Nehalem with an elliptical particle distribution. Although we performed this tuning for all architectures, particle distributions, and precisions, we only present Nehalem data due to space limitations.

The optimal setting of $q$ varies with the level of optimization, with higher levels of optimization enabling larger values of $q$. Since we did not parallelize tree construction, we consider just the evaluation time in Figure 3(b). Thus, one should only tune $q$ (or other parameters affected by parallelization) after all other optimizations have been applied and tuned.

Figure 3(c) decomposes evaluation time by phase. The $Up$ward traversal, $Down$ward traversal, and V-list
execution times decrease quickly with increasing $q$. However, execution time for the other lists, especially U-list, grow with increasing $q$. Observe a crossover point of $q = 250$ where time saved in Up, Down, and V-list can no longer keep pace with the quickly increasing time spent in U-list.

### 5.2 Scalability

Exploiting multicore can be challenging as multiple threads share many resources on a chip like caches, bandwidth, and even floating-point units. Thus, the benefit of thread-level parallelism may be limited.

Figure 4 presents the performance scalability by architecture as a function of thread-level parallelism for the double-precision, uniform distribution case (thus, has no W or X list phases). Threads are assigned first to multiple sockets before multiple cores, and multiple cores before simultaneous multithreading (SMT). For clarity, we highlight the SMT region explicitly. On all machines, the 2 thread case represents one thread per socket.

The top figures show overall times for two cases: (i) assuming tree construction before evaluation, and (ii) the asymptotic limit in which the tree is constructed once and can be infinitely reused. (Recall that tree construction was the only kernel not parallelized.) Observe that Nehalem delivers very good scalability to 8 threads at which point HyperThreading provides no further benefit. Unfortunately, the time required for tree construction becomes a substantial fraction of the overall time. Thus, further increases in core count will deliver sublinear scaling due to Amdahl’s Law.

In the bottom figures, we report thread-seconds to better visualize the scalability of the code for each phase. A flat line denotes perfect scalability and a positive slope denotes sublinear scaling. The U-list initially dominates the overall evaluation time and delivers very good scalability to 8 cores. This observation is not surprising since this phase has high arithmetic intensity. However, the V-list computations, which initially constitute a small fraction of overall time, show relatively poorer scalability, eventually becoming the bottleneck on Nehalem. The V-list has the lowest arithmetic intensity and is likely suffering from bandwidth contention. The other two kernels show good scalability to 4 cores, but remain a small fraction of the overall time.

Barcelona shows good scalability to four cores (2 per socket) but little thereafter. The bottom figure shows the problem: U-list scalability varies some but is reasonably good, while the V-list scales poorly and eventually constitutes 50% more time than U-list. Given Barcelona’s smaller L3 cache and diminished bandwidth, the effects seen on Nehalem are only magnified.

Victoria Falls, with its ample memory bandwidth and paltry floating-point capability, scales perfectly to 16 cores (lines are almost perfectly flat). Once again, performance is dominated by the U-list. As multithreading within a core is scaled, the time required for U-list rockets. Although, we manage better than a $2 \times$ speedup using four threads per core, we see none thereafter.

Figure 5 extends these results to the non-uniform elliptical distribution, which will now include W- and X-list phases. On x86, for the same number of particles, the elliptical time-to-solution is less than the uniform case. This occurs because more interactions can be pruned in the non-uniform case. However, as a consequence, tree construction also becomes a more severe impediment to scalability.

As with the uniform distribution, evaluation time scales well on Nehalem up to 8 threads (one thread per core). Thereafter, it reaches parity with tree construction time. Additional cores or optimization yield only
Figure 4. Double-precision, multicore scalability using OpenMP for a uniform particle distribution. Top: time as a function of thread concurrency showing relative time between list evaluations and tree construction. Bottom: break down of evaluation time by list. Note: “Thread-seconds” is essentially the inverse of GFlop/s/core, so flat lines denote perfect scaling:

an additional $2 \times$ speedup. Also observe that the relative time of each phase changes. U-list time dominates at all concurrencies, and V-list time scales well since the V-lists happen to have fewer boxes for this distribution.

The behavior on Barcelona is similar except, the U-list time scales somewhat more poorly with the elliptical distribution than in the uniform case. Interestingly, most phases scale poorly in the multicore region on Victoria Falls, though U-list time still dominates.

5.3 Architectural Comparison

Beyond the substantial differences among architectures in serial speedups, tuning, and multicore scalability, we observe that the raw performance among processors differs considerably as well. Moreover, performance varies dramatically with floating-point precision.

Figure 6 compares the double-precision performance for our three machines for both particle distributions. For each distribution, performance is normalized to the reference Nehalem implementation. Initially, Nehalem is only about 25% faster than Barcelona; however, after optimization, parallelization, and tuning, Nehalem is more than 3× faster. Although the initial performance differences are expected given the differences in frequency, the final difference is surprising as Nehalem has comparable peak performance and only about twice the bandwidth. Moreover, and unexpectedly, with only 4.5× the peak flops Nehalem is as much as 10× faster than Victoria Falls. Victoria Falls’ small per-thread cache capacity may result in a large number of cache misses. Across the board, we observe the benefit of tree construction amortization is nearly a factor of two.

Unlike the scalar Victoria Falls, x86 processors can more efficiently execute single-precision operations using 4-wide SIMD instructions. GPUs take this approach to the extreme with an 8-wide SIMD-like implementation. To that end, we repeated all optimizations, parallelization, and tuning for all architectures and distributions in single-precision. We then re-ran the GPU-accelerated code of prior work by Lashuk, et al. [9].

For the GPU comparison, we consider 1 node (2 CPU sockets) with either 1 GPU or 2 GPUs. The GPUs perform all phases except tree construction, with
New tree constructed for every force evaluation as asymptotic limit (force evaluation time only)

Figure 5. Double-precision, multicore scalability using OpenMP for an elliptical particle distribution. Top: Time as a function of thread concurrency, comparing evaluations and tree construction components. Bottom: Breakdown of evaluation time by list. Note: "Thread-seconds" is essentially the inverse of GFlop/s/core, so flat lines indicate perfect scaling.

The CPU used only for control and thus largely idle. This experiment allows us to compare not only different flavors of homogenous multicore processors, but also compare them to heterogeneous computers specialized for single-precision arithmetic. Note that the GPU times include host-to-device data structure transfer times.

These single-precision results appear in Figure 6. Barcelona saw the most dramatic performance gains of up to $7.2 \times$ compared with double-precision. This gain greatly exceeds the $2 \times$ increase in either the peak flop rate or operand bandwidth. Nehalem’s gains, around $4 \times$, were also surprisingly high. Victoria Falls typically saw much less than a factor of two performance increase, perhaps due to nothing more than the reduction in memory traffic. We attribute this difference to x86’s single-precision SIMD advantage, as well as the ability to avoid the Newton-Raphson approximation in favor of one rsqrtps (reciprocal square root approximation) instruction without loss of precision.

Although Nehalem’s performance advantage over Victoria Falls increased to as much as $21 \times$, its advantage over Barcelona dropped to as little as $1.6 \times$ — which is still high given the architectural similarities.

Perhaps the most surprising result is that with optimization, parallelization, and tuning, Nehalem is up to $1.7 \times$ faster than one GPU and achieves as much as $\frac{3}{2} \times$ the 2-GPU performance. Where Nehalem’s optimal particles per box was less than 250, GPUs typically required 1K to 4K particles per box. In order to attain a comparable time-to-solution, the GPU implementation had to be configured to prioritize the computations it performs exceptionally well — the computationally intense regular parallelism found in U-list.

5.4 Energy Comparison

We measured power usage using a digital power meter for our three multicore systems. As it was not possible to take measurements on the remote GPU-based system, we include two estimates: peak power (assuming full GPU and full CPU power) and maximum GPU-only power (assuming the CPUs consume zero power). We report the resultant power efficiency relative to parallelized and tuned Nehalem (higher is better). Figure 7
Reference + Optimized + OpenMP

Amortized Energy assuming CPU consumes no power

Double Precision

Nehalem Barcelona VF

Uniform Distribution Elliptical Distribution

Nehalem Barcelona VF

+1 GPU +2 GPUs

Uniform Distribution Elliptical Distribution

Performance Relative to Out-of-the-box Nehalem

Single Precision

Double Precision

Single Precision

Figure 6. Performance relative to out-of-the-box Nehalem for each distribution. Note, performance is on a log scale. Labels show the final execution time (secs) after all optimizations.

shows this relative energy-efficiency as a function of architecture, distribution, and precision.

Nehalem still manages a sizable energy efficiency over all other CPU architectures, although its energy advantage over Barcelona is less than its performance advantage. Conversely, the FBDIMM-based Victoria Falls consumes at least 66% more power than any other CPU-based machine. As such, for FMM, Nehalem is as much as 35 × more energy-efficient than Victoria Falls.

The GPU-based systems, by our estimates, consumes as much as 725W. Thus, Nehalem is as much as 2.37 × and 1.76 × more energy-efficient than systems accelerated using 1 or 2 GPUs, respectively. Even under the optimistic assumption that the largely idle CPUs consumed no power, then Nehalem’s energy-efficiency is still between 0.97 × and 1.65 ×.

6 Conclusions

Given that single-node multicore performance and power efficiency will be critical to scalability on next-generation extreme scale systems, we believe our extensive study of parallelization, low-level and algorithmic tuning at run-time (i.e., the input-dependent maximum points per box, q), numerical approximation, and data structure transformations contributes a solid foundation for the execution of FMM on such machines.

Surprisingly, given a roughly comparable implementation effort, a careful multicore implementation can deliver performance and energy efficiency on par with that of a GPU-based approach, at least for the FMM [9]. We believe this finding is a significant data point in our collective understanding of the strengths and limits of using heterogeneous computers.

Looking forward, we see numerous opportunities. First, we relied largely on bulk-synchronous parallelism, which resulted in working sets that did not fit in cache. Dataflow and work-queue approaches may mitigate this issue. Next, optimal algorithmic parameters like particles per box will vary not only with architecture, but also optimization and scale of parallelism. Finally, our manual SIMD transformations and their interaction with data layout was a significant performance win, and should be a priority for new compiler and/or programming model efforts.

Acknowledgments

We would like to thank the University of California Parallel Computing Laboratory for access to machines. All authors from LBNL were supported by the ASCR Office in the DOE Office of Science under contract number DE-AC02-05CH11231. Authors from UCB were also supported by Microsoft (Award #024263), Intel (Award #024894), and by matching funding by U.C. Discovery (Award #DIG07-10227).

Authors at Georgia Tech were supported in part by the National Science Foundation (NSF) under award number 0833136, and grants from the Defense Advanced Research Projects Agency (DARPA) and Intel Corporation. Any opinions, findings and conclusions or recommendations expressed in this material are those of the authors and do not necessarily reflect those of NSF, DARPA, or Intel.
Figure 7. Energy efficiency relative to optimized, parallelized, and tuned Nehalem for each distribution. Note, efficiency is on a log scale. (higher is better).

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