Lawrence Berkeley National Laboratory
Lawrence Berkeley National Laboratory

Title
MPI-hybrid Parallelism for Volume Rendering on Large, Multi-core Systems

Permalink
https://escholarship.org/uc/item/4r0575d1

Author
Howison, Mark

Publication Date
2010-07-13

Peer reviewed
MPI-hybrid Parallelism for Volume Rendering on Large, Multi-core Systems

M. Howison, E. W. Bethel, and H. Childs
Visualization Group, Lawrence Berkeley National Laboratory

Abstract

This work studies the performance and scalability characteristics of “hybrid” parallel programming and execution as applied to raycasting volume rendering – a staple visualization algorithm – on a large, multi-core platform. Historically, the Message Passing Interface (MPI) has become the de-facto standard for parallel programming and execution on modern parallel systems. As the computing industry trends towards multi-core processors, with four- and six-core chips common today and 128-core chips coming soon, we wish to better understand how algorithmic and parallel programming choices impact performance and scalability on large, distributed-memory multi-core systems. Our findings indicate that the hybrid-parallel implementation, at levels of concurrency ranging from 1,728 to 216,000, performs better, uses a smaller absolute memory footprint, and consumes less communication bandwidth than the traditional, MPI-only implementation.

Categories and Subject Descriptors (according to ACM CCS):
D.1.3 [Programming Techniques]: Concurrent Programming—Parallel programming
F.1.2 [Computation by Abstract Devices]: Modes of Computation—Parallelism and concurrency
I.3.3 [Computer Graphics]: Picture/Image Generation—Display algorithms

1. Introduction

It is well accepted that the future of parallel computing involves chips that are comprised of many (smaller) cores. With this trend towards more cores on a chip, many in the HPC community have expressed concern that parallel programming languages, models, and execution frameworks that have worked well to-date on single-core massively parallel systems may “face diminishing returns” as the number of computing cores on a chip increase [ABC+06]. In the broader high-performance computing community, this general topic has engendered much interest but little published research to date.

In this context, the focus of our work is on exploring the performance and scalability of a common visualization algorithm – raycasting volume rendering – implemented with different parallel programming models and run on a large supercomputer comprised of six-core chips. In this study, we compare a traditional implementation based on message-passing against a “hybrid” parallel implementation, which uses a blend of traditional message-passing (inter-chip) and shared-memory (intra-chip) parallelism. The the-

Figure 1: This 46082 image of a combustion simulation result was rendered by our MPI+pthreads implementation running on 216,000 cores of the JaguarPF supercomputer.
sis we wish to test is whether there are opportunities in the hybrid-parallel implementation for performance and scalability gains that result from using shared-memory parallelism among cores within a chip.

Over the years, there has been a consistent and well-documented concern that the overall runtime of large-data visualization algorithms is dominated by I/O costs (e.g., SKP96, KMM 01, PYR 09). During our experiments, we observed results consistent with previous work: there is a significant cost associated with scientific data I/O. In this study, however, we focus exclusively on the performance and scalability of the ray casting volume rendering algorithm, not on parallel I/O performance. This approach is valid for many visualization use cases, such as creating multiple images from a single dataset that fits entirely within the memory footprint of a large system, or creating one or more images of data that is already resident in memory, as in the case of in-situ visualization.

Our findings (Section 4) show that there is indeed opportunity for performance gains when using hybrid-parallelism for raycasting volume rendering across a wide range of concurrency levels. The hybrid-parallel implementation runs faster, requires less memory, and for this particular algorithm and set of implementation choices (Section 3), consumes less communication bandwidth than the traditional, MPI-only implementation.

2. Background and Previous Work

2.1. Parallel Volume Rendering

Volume rendering is a common technique for displaying 2D projections of 3D sampled data [Lev88, DCH88] and is computationally, memory, and data I/O intensive. In the quest towards interactivity, as well as to address the challenges posed by growing data size and complexity, there has been a great deal of work over the years in the space of parallel volume visualization (see Kaufman and Mueller [KM05] for an overview).

Within the field of parallel volume rendering, our work focuses on examining a form of “hybrid parallelism,” which is distinct and different from the term “hybrid volume rendering.” Hybrid volume rendering refers to using combination of image- and object-order techniques to perform rendering [KM05]. In contrast, hybrid parallel refers to the use of a mixture of shared- and distributed-memory algorithms, programming, and execution environments [LG08].

However, our hybrid-parallel implementation also makes use of hybrid volume rendering (e.g., [MPHK03, TH94, Ma05, BIP09]), starting with an object-order partitioning to distribute source data blocks to processors where they are rendered using ray casting [Lev88, Sab88, DCH88, UK88]. Within a processor, we then use an image-space decomposition, similar to [NL92], to allow multiple rendering threads to cooperatively generate partial images that are later combined via compositing into a final image ( [DCH88, Lev88, UK88]).

Hybrid volume rendering techniques have proven successful in achieving scalability and tackling large data sizes. The TREX system [KMM 01] is a hybrid volume rendering algorithm on a shared-memory platform that uses object-parallel data domain decomposition and texture-based, hardware-accelerated rendering followed by a parallel, software-based composition phase with image-space partitioning. The design choices for which part of the SGI Origin to use for different portions of the algorithm reflect a desire to achieve optimal performance at each algorithmic stage and to minimize inter-stage communication costs. Childs et al. [CDM06] present a hybrid scheme for volume rendering massive datasets (with one hundred million unstructured elements and a 3000^2 rectilinear data set). Their approach parallelizes over both input data elements and output pixels, and is demonstrated to scale well on up to 400 processors. Peterka et al. were the first to run a hybrid volume rendering algorithm at massive concurrency, rendering 4480^3 data sizes with 32,000 cores on an IBM BG/P system [PYR 09]. They demonstrated generally good scalability and found that the compositing phase slowed down when more than ten thousand cores were involved, likely due to hardware or MPI limitations. To address this problem, they reduced the number of processors involved in the compositing phase.

The most substantial difference between our work and previous work in hybrid volume rendering is that we are exploiting hybrid parallelism. In addition, we contribute to the knowledge of hybrid volume rendering at massive concurrency: we performed experiments with up to 216,000 cores, which is more than six times larger than previously published results. Further, our study reproduces the degradation in compositing performance first discovered by Peterka et al. [PYR 09] via a different software implementation and supercomputer, and we present results that characterize this effect in more detail.

2.2. Traditional and Hybrid Parallelism

The Message Passing Interface (MPI) evolved as the de-facto standard for parallel programming and execution on machines consisting of single-core CPUs interconnected via a high-speed fabric [SOHL 98]. To use MPI, an application developer must explicitly add MPI library calls to an application to implement fundamental parallel execution motifs: data scatter and gather, execution synchronization, and so forth. In MPI parlance, a processing element (PE) is the fundamental unit of execution, and historically each MPI PE has mapped one-to-one to the processors of a massively parallel (MPP) system. To support more recent multi-core MPPs, vendors’ MPI implementations provide support for MPI PEIs to be mapped onto one or more cores of multi-core chips.
However, on multi-core platforms, there may be opportunities for more efficient inter-PE communication through local, high-speed, shared memory that bypasses the MPI interface.

Shared-memory parallel applications are somewhat easier to develop than distributed memory ones as there is no need to explicitly move data among parallel program elements. Instead, each execution thread has access to the same shared memory within a single address space. Common programming models for shared-memory parallelism include POSIX threads [But07], OpenMP [CDK01], and Intel Thread Building Blocks [Rei07]. These APIs allow applications to manage creation and termination of execution threads, and synchronize thread execution through semaphores and mutexes. The scalability of shared-memory codes is typically limited by physical constraints: there are typically only a few cores in a single CPU. Four to six cores per CPU are common today, although the trend seems to be towards hundreds to thousands of cores per chip.

In all of the above models, the developer must explicitly design for parallelism, as opposed to relying on a compiler to discover and implement parallelism. Other approaches, which allow the developer to express parallelism implicitly via language syntax, include data-parallel languages (e.g., CUDA [NV108]), languages with data parallel extensions (e.g., High Performance Fortran [Hig93]), and partitioned global address space (PGAS) languages (e.g., Unified Parallel C (UPC) [EGCSY05], which offers the developer a single-address space view of memory, even on distributed memory platforms).

Hybrid parallelism has evolved in response to the widespread deployment of multi-core chips in distributed-memory systems. An MPI-hybrid model allows data movement among nodes using traditional MPI motifs like scatter and gather, but within nodes using shared-memory parallelism via POSIX threads or OpenMP. Previous work comparing MPI-only with MPI-hybrid implementations (e.g., [HJR09, MTT+09]) has focused on benchmarking well-known computational kernels. In contrast, our study examines this space from the perspective of visualization algorithms.

The previous studies point to several areas where MPI-hybrid may outperform MPI-only. First, MPI-hybrid tends to require less memory for applications with domain decomposition (e.g., parallel volume rendering), since fewer domains means less “surface area” between domains and hence less exchange of “ghost zone” data. Second, the MPI runtime allocates various tables, buffers, and constants on a per-PE basis. Today, the gain from using fewer PEs to reduce this memory overhead may seem small with only four or six cores per chip, but the trend towards hundreds of cores per chip with less memory per core will magnify these gains. Third, MPI-hybrid can use only one PE per node for collective operations such as scatter, gather and all-to-all, thereby reducing the absolute number of messages traversing the inter-connect. While the size of the messages in this scenario may be larger or smaller depending upon the specific problem, a significant factor influencing overall communication performance is latency, which is reduced by using fewer messages.

The primary contributions of our work is a comparison of the performance and resource requirements of hybrid- and traditional-parallel implementations of raycasting volume rendering. Our methodology and results (Section 4) are consistent with those identified in previous studies of hybrid parallelism from the HPC community (e.g., [HJR09]).

3. Implementation

From a high level view, our parallel volume rendering implementation repeats a design pattern that forms the basis of previous work (e.g., [MPHK93, TH94, Ma95, BIJP99]).

![Diagram of our system architecture.](image-url)
to produce a set of image fragments. Next, each of the \( n \) processes participates in a compositing stage in which fragments are exchanged and combined into a final image. Each of the \( n \) processes then sends its portion of the completed image to process 0 for display or I/O to storage. Figure 2 provides a block-level view of this organization. Our distributed memory parallel implementation is written in C++ and C and makes calls to MPI [SOHL’98]. The portions of the implementation that are shared-memory parallel are written using a combination of C++ and C and either POSIX threads [But97] or OpenMP [CDK’01], so that we are actually comparing two hybrid implementations that we refer to as MPI+Pthreads and MPI+OpenMP.

The MPI-only and MPI-hybrid implementations differ in several key respects. First, the raycasting volume rendering algorithm itself is serial on each MPI-only PE or is shared-memory multicore-parallel in the MPI-hybrid case. We discuss this issue in more detail in Section 3.1. Second, the communication topology in the compositing stage differs slightly, and we discuss this issue in more detail in Section 3.2. A third difference is in how data is partitioned across the pool of parallel processes. In the MPI-only implementation, each PE loads and operates on a disjoint block of data. In the MPI-hybrid case, each MPI PE loads a disjoint block of data and each of its worker threads operate in parallel on that data using an image-parallel decomposition [NL92].

### 3.1. Parallel, Multicore Raycasting Volume Rendering

Our raycasting volume rendering code implements Levoy’s method [Lev88]: we compute the intersection of a ray with the data block, and then compute the color at a fixed step size along the ray through the volume. All such colors along the ray are integrated front-to-back using the “over” operator. Output from our algorithm consists of a set of image fragments that contain an \( x, y \) pixel location, \( R, G, B, \alpha \) color information, and a \( z \)-coordinate. The \( z \)-coordinate is the location in eye coordinates where the ray penetrates the block of data. Later, these fragments are composited in the correct order to produce a final image (see Section 3.2).

In the MPI-only case, this serial implementation is invoked on each of the MPI PEs. Each operates on its own disjoint block of data. As we are processing structured rectilinear grids, all data subdomains are spatially disjoint, so we can safely use the ray’s entry point into the data block as the \( z \)-coordinate for sorting during the later composition step.

In the MPI-hybrid case, the raycasting volume renderer on each MPI PE is a shared-memory parallel implementation consisting of \( T \) threads all executing concurrently to perform the raycasting on a single block of data. As in [NL92], we use an image-space partitioning: each thread is responsible for raycasting portion of the image. Our image-space partitioning is interleaved, where the image is divided into many small tiles that are distributed amongst the threads. Through a process of manual experimentation, we determined that an image tile size of \( 32 \times 32 \) pixels produced consistently better performance than other tile sizes for a variety of volume sizes on the six-core AMD Opteron processor. We also found that a dynamic work assignment of tiles to threads minimized load imbalance.

### 3.2. Parallel Compositing

The compositing algorithm takes the fragments generated by the raycasting algorithm and produces the final image. The compositing begins by partitioning the pixels of the final image amongst the MPI PEs. It then performs an all-to-all communication step, where MPI PEs send and receive fragments. The sent fragments are the fragments from that MPI PE’s raycasting phase. The MPI PE identifies where to send fragments using the partition information. The received fragments correspond to the pixels of the final image that that MPI PE owns. The fragments are exchanged using an MPI_Alltoallv call. This call provides the same functionality as direct sends and receives, but bundles the messages into fewer point-to-point exchanges, and thus is more efficient. After the fragments are exchanged, each MPI PE is able to perform the final compositing for its portion of the larger image, using the “over” operator on the fragments. The final step is to collect the image subpieces to MPI PE 0 and output the entire image.

Peterka et al. [PYR’09] reported scaling difficulties for compositing when using more than 8,000 MPI PEs. They solved this problem by reducing the number of MPI PEs receiving fragments to be no more than 2,000. We emulated this approach, again limiting the number of MPI PEs receiving fragments, although we experimented with values higher than 2,000.

In the hybrid implementations, only one thread per socket participates in the compositing phase. That thread gathers fragments from all other threads in the same socket, packs them into a single buffer, and transmits them to other compositing PEs. This approach results in fewer messages than if all threads in the hybrid parallel implementation were to send messages to all other threads on all other CPUs. Our aim here is to better understand the opportunities for improving performance in the hybrid-parallel implementation. The overall effect of this design choice is an improvement in communication characteristics, as indicated in Section 4.4.2.

### 4. Results

#### 4.1. Methodology

Our methodology is designed to test the hypothesis that an MPI-hybrid implementation exhibits better performance and resource utilization than the MPI-only implementation. We compare the cost of MPI runtime overhead and corresponding memory footprint in Section 4.3.1; compare the absolute
amount of memory required for data blocks and ghost zone (halo) exchange in Section 4.3.2; compare scalability of the raycasting and compositing algorithms in Sections 4.4.1 and 4.4.2; and compare the levels of communication required during the compositing phase in Section 4.4.2.

Our test system, JaguarPF, is a Cray XT5 located at Oak Ridge National Lab and was recently ranked by the Top500 list as the fastest supercomputer in the world with a theoretical peak performance of 2.3 Petaflops [Top09]. Each of the 18,688 nodes has two sockets, and each socket has a six-core 2.6GHz AMD Opteron processor, for a total of 224,256 compute cores. With 16GB per node (8GB per socket), the system has 292TB of aggregate memory and roughly 1.3GB per core.

We conducted a strong scaling study where we rendered a 4608$^3$ image from a 4608$^3$ dataset (roughly 97.8 billion cells) at concurrency levels from 1,728-way parallel to 216,000-way parallel. On JaguarPF, a minimum of 1,728 cores is required to accommodate this particular problem size. In the hybrid case, we shared a data block among six threads and used one sixth as many MPI PEs. Although we could have shared a data block among as many as twelve threads on each dual-socket six-core node, we chose not to because sharing data across sockets results in non-uniform memory access. Based on preliminary tests, we estimated this penalty to be around 5 or 10% of the raycasting time. Because the time to render is view-dependent, we executed each raycasting phase ten times over a selection of ten different camera locations (see Figure 3). The raycasting times we report are an average over all locations.

In the compositing phase, we tested five different ratios of total PEs to compositing PEs. We restricted the compositing experiment to only two views (the first and last) because it was too costly to run a complete battery of all view and ratio permutations. Since the runtime of each trial can vary due to contention for JaguarPF’s interconnection fabric with other users, we ran the compositing phase ten times for both views. We report mean and minimum times over this set of twenty trials. Minimum times most accurately represent what the system is capable of under optimal, contention-free conditions, while mean times help characterize the variability of the trials.

### 4.2. Source Data and Decomposition

Starting with a 512$^3$ dataset of combustion simulation results $\dagger$, we used trilinear interpolation to upscale it to arbitrary sizes in memory. We scaled equally in all three dimensions to maintain a cubic volume. Our goal was to choose a problem size that came close to filling all available memory (see Table 1). Although upscaling may distort the results for a data-dependent algorithm, the only data dependency during raycasting is early ray termination. However, we found that for our particular dataset and transfer function, there was always at least one data block for which no early terminations occurred. Moreover, the cost of the extra conditional statement inside the ray integration loop to test for early termination added a 5% overhead. Therefore, we ran our study with early ray termination turned off, and we believe that upscaling the dataset does not effect our results.

$\dagger$ Sample data courtesy J. Bell and M. Day, Center for Computational Sciences and Engineering, Lawrence Berkeley National Laboratory.

![Figure 3: The raycasting phase used these ten camera locations to approximate an “average” use case.](image)

<table>
<thead>
<tr>
<th>Color</th>
<th>Camera</th>
</tr>
</thead>
<tbody>
<tr>
<td>Black</td>
<td>(1,0,0)</td>
</tr>
<tr>
<td>Black</td>
<td>(0,1,0)</td>
</tr>
<tr>
<td>Black</td>
<td>(0,0,1)</td>
</tr>
<tr>
<td>Cyan</td>
<td>(0,1,1)</td>
</tr>
<tr>
<td>Magenta</td>
<td>(1,0,1)</td>
</tr>
<tr>
<td>Yellow</td>
<td>(1,1,0)</td>
</tr>
<tr>
<td>Red</td>
<td>(2,1,1)</td>
</tr>
<tr>
<td>Green</td>
<td>(1,2,1)</td>
</tr>
<tr>
<td>Blue</td>
<td>(1,1,2)</td>
</tr>
<tr>
<td>Gray</td>
<td>(1,1,1)</td>
</tr>
</tbody>
</table>

Table 1: Problem size configurations and per-node memory requirements for our experiment. The memory usage for MPI-hybrid and MPI-only at 216,000 cores differs because they have different dataset dimensions.

<table>
<thead>
<tr>
<th></th>
<th>MPI-only</th>
<th>MPI-hybrid</th>
</tr>
</thead>
<tbody>
<tr>
<td>MPI PEs</td>
<td>Block Dimensions</td>
<td>MPI PEs</td>
</tr>
<tr>
<td>12$^3$=1728</td>
<td>384 × 384 × 384</td>
<td>288</td>
</tr>
<tr>
<td>24$^3$=13824</td>
<td>192 × 192 × 192</td>
<td>2304</td>
</tr>
<tr>
<td>36$^3$=46656</td>
<td>128 × 128 × 128</td>
<td>7776</td>
</tr>
<tr>
<td>48$^3$=110592</td>
<td>96 × 96 × 96</td>
<td>18432</td>
</tr>
<tr>
<td>60$^3$=216000</td>
<td>76 × 76 × 76</td>
<td>36000</td>
</tr>
</tbody>
</table>
Table 2: Memory usage as measured directly after MPI initialization.

<table>
<thead>
<tr>
<th>Cores</th>
<th>Mode</th>
<th>MPI PE</th>
<th>Per PE (MB)</th>
<th>Per Node (MB)</th>
<th>Aggregate (GB)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1728</td>
<td>MPI-hybrid</td>
<td>288</td>
<td>67</td>
<td>133</td>
<td>19</td>
</tr>
<tr>
<td>1728</td>
<td>MPI-only</td>
<td>1728</td>
<td>67</td>
<td>807</td>
<td>113</td>
</tr>
<tr>
<td>13824</td>
<td>MPI-hybrid</td>
<td>2304</td>
<td>67</td>
<td>134</td>
<td>151</td>
</tr>
<tr>
<td>13824</td>
<td>MPI-only</td>
<td>13824</td>
<td>71</td>
<td>857</td>
<td>965</td>
</tr>
<tr>
<td>46656</td>
<td>MPI-hybrid</td>
<td>7776</td>
<td>68</td>
<td>136</td>
<td>518</td>
</tr>
<tr>
<td>46656</td>
<td>MPI-only</td>
<td>46656</td>
<td>88</td>
<td>1055</td>
<td>4007</td>
</tr>
<tr>
<td>110592</td>
<td>MPI-hybrid</td>
<td>18432</td>
<td>73</td>
<td>146</td>
<td>1318</td>
</tr>
<tr>
<td>110592</td>
<td>MPI-only</td>
<td>110592</td>
<td>121</td>
<td>1453</td>
<td>13078</td>
</tr>
<tr>
<td>216000</td>
<td>MPI-hybrid</td>
<td>36000</td>
<td>82</td>
<td>165</td>
<td>2892</td>
</tr>
<tr>
<td>216000</td>
<td>MPI-only</td>
<td>216000</td>
<td>176</td>
<td>2106</td>
<td>37023</td>
</tr>
</tbody>
</table>

Because the compute nodes on an XT5 system have no physical disk for swap, and hence no virtual memory, exceeding the physical amount of memory causes program termination. Our total memory footprint was four times the number of bytes in the entire dataset: one for the dataset itself, and the other three for the gradient data, which we computed by central difference and used in shading calculations. Although each node has 16GB of memory, we could reliably allocate only 10.4GB for the data block and gradient field at 1,728-way concurrency because of overhead from the operating system and MPI runtime.

We chose concurrencies that are cubic numbers to allow for a clean decomposition of the entire volume into cubic blocks per MPI PE. In the MPI-hybrid case, however, these blocks are rectangular because we aggregated six blocks ($1 \times 2 \times 3$) into one shared block. For 216,000 cores, we could not evenly divide the 4608$^3$ dataset, and chose to round down to a 4560$^3$ dataset for MPI-only and a 4560$ \times $4590$ \times $4600 dataset for MPI-hybrid. As a result, the MPI-only dataset is approximately 1.4% smaller than the MPI-hybrid dataset. While this difference might seem to give an advantage to the MPI-only implementation, results in later sections show that the MPI-hybrid performance and resource utilization are uniformly better than for MPI-only.

4.3. Memory Usage

4.3.1. Overhead at Initialization

Because MPI-hybrid uses fewer MPI PEs, it incurs less memory overhead from the MPI runtime environment and from program-specific data structures that are allocated per PE. Table 2 shows the memory footprint of the program as measured directly after calling MPI_Init and reading in command-line parameters. We collected the VmRSS, or “resident set size,” value from the /proc/self/status interface. Memory usage was sampled only from MPI PE 0 through 6, but those values agreed within 1–2%. Therefore, the per-PE values we report in Table 2 are from PE 0 and the per-node and aggregate values are derived from the per-task value: MPI-only uses twelve PEs per node versus MPI-hybrid’s two PEs per node, and in the aggregate MPI-hybrid has one sixth as many PEs. At 216,000 cores, the per-PE runtime overhead of MPI-only is more that 2× that of MPI-hybrid and the per-node and aggregate memory usage is another factor of six larger for MPI-only because it uses 6× as many PEs as MPI-hybrid. Thus, MPI-only uses nearly 12× as much memory per-node and in-aggregate than MPI-hybrid to initialize the MPI runtime at 216,000-way concurrency.

<table>
<thead>
<tr>
<th>Cores</th>
<th>MPI-only</th>
<th>MPI+pthreads</th>
<th>MPI+OpenMP</th>
</tr>
</thead>
<tbody>
<tr>
<td>1728</td>
<td>24.88s</td>
<td>22.31s</td>
<td>22.23s</td>
</tr>
<tr>
<td>13824</td>
<td>3.10s</td>
<td>2.84s</td>
<td>2.83s</td>
</tr>
<tr>
<td>46656</td>
<td>0.92s</td>
<td>0.85s</td>
<td>0.85s</td>
</tr>
<tr>
<td>110592</td>
<td>0.38s</td>
<td>0.37s</td>
<td>0.37s</td>
</tr>
<tr>
<td>216000</td>
<td>0.19s</td>
<td>0.21s</td>
<td>0.20s</td>
</tr>
</tbody>
</table>

Figure 4: Ghost data required by our implementations.
4.3.2. Ghost Data

Two layers of ghost data are required in our raycasting phase: the first layer for trilinear interpolation of sampled values, and the second layer for computing the gradient field using central differences (gradients are not precomputed for our dataset). Because the MPI-hybrid approach uses fewer, larger blocks in its decomposition, it requires less exchange and storage of ghost data by roughly 40% across all concurrencies (see Figure 4).

4.4. Scaling Study

4.4.1. Raycasting

Our raycasting phase scales nearly linearly because it involves no inter-processor communication (see Figure 5). Each MPI PE obtains its data block, then launches either one (MPI-only) or six (MPI-hybrid) raycasting threads. Working independently, each thread tests for ray-triangle intersections along the data block’s bounding box and, in the case of a hit, integrates the ray by sampling data values at a fixed interval along the ray, applying a transfer function to the values, and aggregating the resulting color and opacity in a fragment for that ray’s image position. For these runs and timings, we use trilinear interpolation for data sampling along the ray as well as a Phong-style shader. The final raycasting time is essentially the runtime of the thread that takes the most integration steps. This behavior is entirely dependent on the view. Our approach, which is aimed at understanding “average” behavior, uses ten different views (see Figure 3 in Section 4.1) and reports their average runtime.

Overall, we have achieved linear scaling up to 216,000 for the raycasting phase with MPI-only (see Figure 5). MPI-hybrid exhibits different scaling behavior because it has a different decomposition geometry: MPI-only has a perfectly cubic decomposition, while MPI-hybrid aggregates 1 × 2 × 3 cubic blocks into rectangular blocks that are longest in the z-direction (see Table 1). The interaction of the decomposition geometry and the camera direction determine the maximum number of ray integration steps, which is the limiting factor for the raycasting time. At lower concurrencies, this interaction benefits MPI-hybrid, which outperforms MPI-only by as much as 11% (see Table 3). At higher concurrencies the trend flips, and MPI-only outperforms MPI-hybrid by 10%. We expect that if we were able to run on an eight-core system with a 2 × 2 × 2 aggregation factor for MPI-hybrid, both implementations would scale identically. We also note that at 216,000 cores, raycasting is less than 20% of the total runtime (see Figure 6), and MPI-hybrid is over 50% faster because of gains in the compositing phase that we describe next.

4.4.2. Compositing

We observe the same effect that Peterka et al. [PYR’09] report: for higher concurrencies, it is more beneficial to use...
only a subset of PEs for compositing. While Peterka et al. found that 1,000 to 2,000 compositors were optimal for up to 32,768 total PEs, we have found that at 46,656 PEs and above the optimal number of compositors is closer to 4,000 to 8,000 (see Figure 7). We note that there are many differences between our study and theirs in the levels of concurrency, architectures, operating systems, communication networks, and MPI libraries, each potentially introducing variation in the ideal number of compositors.

Above 1,728 cores, we observe that the compositing times are systematically better for the MPI-hybrid implementation. The primary cost of compositing is the MPI_Alltoallv call that moves each fragment from the PE where it originated during raycasting to the compositing PE that owns the region of image space where the fragment lies. Because MPI-hybrid aggregates these fragments in the memory shared by six threads, it uses on average about $6 \times$ fewer messages than MPI-only (see Figure 8). In addition, MPI-

Figure 5: The speedups (referenced to 1,728 cores) for both the raycasting phase and the total render time (raycasting and compositing). The raycasting speedup is linear for MPI-only, but sublinear for MPI-hybrid: this effect is caused by the difference in decomposition geometries (cubic versus rectangular). At higher concurrencies, however, raycasting accounts for a smaller proportion of the total time and MPI-hybrid exhibits better overall scaling because of its gains in the compositing phase.

Figure 6: In terms of total render time, MPI-hybrid outperforms MPI-only at every concurrency level, with performance gains improving at higher concurrency.
hybrid exchanges less fragment data because its larger data blocks allow for more compositing to take place during ray integration.

We observe a critical point in the compositing performance beginning at 13,824 cores where the elapsed time for MPI-only begins to increase with additional compositors. A similar critical point occurs for MPI-hybrid at 46,656 cores. We believe this critical point exists due to the characteristics of the underlying interconnect fabric. Because MPI-hybrid generates fewer and smaller messages, the critical point occurs at a higher level of concurrency than for MPI-only. Also, our MPI-hybrid version outperforms the MPI-only version across all configurations in these tests. Future work will examine this issue in more detail and on different architectures to better illuminate factors contributing to performance of this stage of processing.

4.4.3. Overall Performance

At 216,000 cores, the best compositing time for MPI-hybrid (0.35s, 4500 compositors) is 67% less than for MPI-only (1.06s, 6750 compositors). Furthermore, at this scale compositing time dominates rendering time, which is roughly 0.2s for both MPI-only and MPI-hybrid. Thus, the total render time is 55% faster for MPI-hybrid (0.56s versus 1.25s). Overall, the scaling study shows that the advantage of MPI-hybrid over MPI-only becomes greater as the number of cores increases (see Figure 6).

5. Conclusion and Future Work

The multi-core era offers new opportunities and challenges for parallel applications. This study has shown that hybrid parallelism offers performance improvements and better resource utilization for raycasting volume rendering on a large, distributed-memory supercomputer comprised of multi-core CPUs. The advantages we observe for hybrid parallelism are reduced memory footprint, reduced MPI overhead, and reduced communication traffic. These advantages are likely to become more pronounced in the future as the number of cores per CPU increases while per-core memory size and bandwidth decrease.

We found that at high concurrency, fragment exchange during the compositing phase is the most expensive operation. Our compositing algorithm relies entirely on the implementation of the MPI_Alltoallv call in the Cray MPI library. Our finding that using a subset of compositing PEs reduces communication overhead agrees with what Peterka et al. [PYR'09] found for the MPI implementation on an IBM BG/P system, suggesting that both libraries use similar implementations and optimizations of MPI_Alltoallv. A separate paper by Peterka et al. [PGR'09] introduces a new compositing algorithm, “Radix-k,” that shows good scaling up to 16,000-way concurrency. This approach, which is compatible with our hybrid parallel system, may lead to even faster compositing times. Studying their combined performance, especially at concurrentities in the hundreds of thousands, is an avenue for future work.

Acknowledgments

This work was supported by the Director, Office of Science, Office and Advanced Scientific Computing Research, of the U.S. Department of Energy under Contract No. DE-AC02-05CH11231 through the Scientific Discovery through Advanced Computing (SciDAC) program’s Visualization and Analytics Center for Enabling Technologies (VACET). This research used resources of the National Center for Computational Sciences at Oak Ridge National Laboratory, which is supported by the Office of Science of the U.S. Department of Energy under Contract No. DE-AC05-00OR22725. Additionally, preliminary results for this work were collected using resources of the National Energy Research Scientific Computing Center (NERSC), which is supported by the Office of Science of the U.S. Department of Energy under Contract No. DE-AC02-05CH11231.

References


[BUP99] Baiac C., Jim I., Joo G., Park S.: Parallel ray casting of visibly human on distributed memory architectures. In


