Abstract:

Graphics Processing Units (GPUs) are a class of specialized parallel architectures with tremendous computational power. The Compute Unified Device Architecture (CUDA) programming model from NVIDIA facilitates programming of general purpose applications on their GPUs. In this project, we target Parsec benchmarks to provide orders of performance speed up and reducing overall execution time on multi-core GPU platforms. Parsec benchmark suite presents and characterizes applications for CPU computation which runs on multiprocessors for providing high level of parallelism for specific applications. We extended seven of the Parsec applications to the CUDA programming language which extends parallelism to multi-GPU systems and GPU-cluster environments. Our methodology calculates execution time at different level for GPUs applications while varying the number and configuration of GPUs and the size of the input data set and allows us to look for overheads. Emphasis is placed on optimization the applications by directly targeting the architecture to best exploit the computational capabilities of NVIDIA Tesla T10 to achieve substantial speed up.

1. Introduction:

The benefits of using Graphics Processing Units (GPUs) for general purpose programming has been recognized for some time. With multiple cores and large memory bandwidth, the computational capability of GPU today is much higher than that of the CPU. Due to its high computational capability, the GPU nowadays serves not only for accelerating the graphics display but also for speeding up non-graphics applications. General purpose GPU (GPGPU) programming has become the scientific computing platform of choice mainly due to the availability of standard C libraries using NVIDIA’s CUDA programming interface running on NVIDIA GPUs and a number of efforts have explored how to reap large performance gains on CUDA-enabled GPUs.

This project is another effort to parallelize Parsec benchmark suite on GPU using CUDA to provide orders of performance speed up and reducing the overall execution time that cannot be achieved by running on CPU platform. In our CUDA implementation we compared CPU vs. GPU runtime for different size of inputs and different size of threads and we tried to identify potential overheads of each application running on GPU. Emphasis has also being made to overcome those bottlenecks and optimize the CUDA code running on GPU with some optimization techniques like Memory Coalescing, Shared Memory, Cache-efficient texture memory accesses, loop unrolling, Parallel Reduction and Page-locked Memory allocation. CUDA compute capability 1.2 and 1.3 e.g. GTX 280 used in this project offers specialized memory spaces depending on the application: Per-block shared memory, constant, and texture memories to reduce overheads. Each is suited to different data-use patterns. Although we used some optimization techniques to alleviate these issues, they remain a bottleneck for some applications.

The benchmarks have been evaluated on an NVIDIA GeForce GTX 280 GPU with a 1.33 GHz core clock, on board memory of 4GB, ultra-fast memory access with 102 GB/s peak bandwidth per GPU and varying GPU power consumption of 160W.

The rest of this paper is organized as follows. Section 2 shows glimpses of what kind of Parsec application has been transformed to CUDA, CUDA architecture and general CUDA programming model. Section 3 presents a general algorithm for CUDA programming and requirements for optimization. Section 4 explains details of the benchmark, their transformation to CUDA, results and overheads and optimization techniques used in eliminating the overheads. Section 5 presents results for execution time for different size of inputs for different size of threads.

2. Parsec Benchmark Application and Compute Unified Device Architecture Glimpse:

2.1. Parsec Benchmarks

The Princeton Application Repository for Shared-Memory Computers (PARSEC) benchmark is a benchmark suite composed of multithreaded programs. The suite focuses on emerging workloads and was designed to contain a diverse selection of applications in the field of data mining, computer vision, visualization, computer animation, financial analysis etc that represents of shared-memory programs for chip-multiprocessors. It provides a selection of next generation workloads that satisfies the following five criteria: 1. Composed of multithreaded applications. 2. Focuses on emerging workloads. 3. Diverse enough to represent the increasingly heterogeneous ways in which multiprocessors are used. 4. Workloads employ state-of-art techniques. 5. Support research

Workloads

Blackscholes This application is an Intel RMS benchmark. It calculates the prices for a portfolio of European options analytically with the Blackscholes partial differential equation (PDE).

Ferret This application is based on the Ferret toolkit which is used for content-based similarity search. It was developed by Princeton University. In the benchmark, we have configured the Ferret toolkit for image similarity search.

Fluidanimate This Intel RMS application uses an extension of the Smoothed Particle Hydrodynamics (SPH) method to simulate an incompressible fluid for interactive animation purposes.

Rastrace The Intel RMS application uses a version of the raytracing method that would typically be employed for real-time animations such as computer games.

Streamcluster This RMS kernel solves the online clustering problem. Streamcluster was included in the PARSEC benchmark suite because of the importance of data mining algorithms and the prevalence of problems with streaming characteristics.

Swaptions The application is an Intel RMS workload which uses the Heath-Jarrow-Morton (HJM) framework to price a portfolio of swaptions. Swaptions employs Monte Carlo (MC) simulation to compute the prices.
This application is an H.264 which describes the lossy compression of a video stream and is also part of ISO/IEC MPEG-4.

2.2 CUDA

The Multi-core CPUs and GPUs have been used to exploit parallelism in performing compute-intensive operations on structured data, i.e. data stored in databases. However, recent trends in data storage suggest that more and more data is being captured in the form of images, audio and video files. As a result of continued demand for programmability, modern graphics processing units (GPUs) such as the NVIDIA are designed as programmable processors employing a large number of processor cores.

The NVIDIA Tesla T10 has 30 multiprocessor units, each consisting of 8 processor cores that execute in SIMD manner. The processors (SIMD units) within a multiprocessor unit communicate through a fast on-chip shared memory, while the different multiprocessor units communicate through a slower off-chip DRAM, also called global memory. Each multiprocessor unit also has a fixed number of registers. The GPU code is launched for execution in the GPU device by the CPU (host). The host transfers data to and from GPU’s global memory. Programming GPUs for general-purpose applications is enabled through an easy-to-use C interface exposed by the NVIDIA Compute Unified Device Architecture (CUDA) model. CUDA programming is done with standard ANSI C extended with keywords that designate data-parallel functions, called kernels, and their associated data structures to the compute devices. These kernels describe the work of a single thread and typically are invoked on thousands of threads. The CUDA programming model abstracts the processor space as a grid of thread blocks (that are mapped to multiprocessors in the GPU device), where each thread block is a grid of threads (that are mapped to SIMD units within a multiprocessor). More than one thread block can be mapped to a multiprocessor unit, and more than one thread can be mapped to a SIMD unit in a multiprocessor. Threads within a thread block can efficiently share data through the fast on-chip shared memory and can synchronize their execution to coordinate memory accesses. Each thread in a thread block is uniquely identified by its thread block id and thread id. A grid of thread blocks is executed on the GPU by running one or more thread blocks on each multiprocessor. Threads in a thread block are divided into SIMD groups called warps (the size of a warp for the NVIDIA GeForce 8800 GTX is 32 threads) and periodic switching between warps is done to maximize resource utilization.

The NVIDIA Tesla T10 contains 1.4 billion transistors, 240 cores with streaming multiprocessor consists of 8 Streaming Processors (SPs), 2 Special Function Units (SFUs) 16 KB R/W shared memory and total of 16384 registers. The Tesla is capable of 1 TFLOPs/s of processing performance and comes with 4 GB of GDDR3 at 102 GB/s bandwidth. Fig 1 and 2 shows the block and grid. Block is a collection of threads. There are up to 1024 threads per SM. Each thread can communicate within the same block. Grid is a collection of blocks. It is usually one per CUDA device and used for multiple iterations. Threads cannot communicate across grids.

3. GPU Computing: CUDA Implementation

This section describes a basic algorithm of transformation of C to CUDA which in cases similar for all Parsec application. Also, this section defines basic optimization techniques used. The details of these are explained in details in section 4.

**Basic Algorithm for CUDA implementation**

The GPU is seen as a compute device to execute a portion of an application that has to be executed many times, can be isolated as a function and which works independently on different data. Such a function can be compiled to run on the device. The following steps are followed for CUDA implementation for the already written C code available in Parsec benchmarks suite 2.1.
Set the grid size
const int N = 1024;
const int blocksize = 16;

Compute Kernel
__global__ void add_matrix( float* a, int N )
{ int i = blockIdx.x * blockDim.x + threadIdx.x; }

cudaMalloc( (void**)&ad, size );
cudaMemcpy( ad, a, size, cudaMemcpyDeviceToHost );

Copy data to GPU
cudaMemcpy( ad, a, size, cudaMemcpyDeviceToHost );

Free memory
cudaFree(ad );
cudaFree(bd );
cudaFree(cd );
const int size = N*N*sizeof(float);

CPU memory Allocation
float *a = new float[N*N];
a[i] = 1.0f; };

GPU Memory Allocation
cudaMalloc( (void**)&ad, size );
cudaMemcpy( ad, a, size, cudaMemcpyDeviceToHost );

Copy data back To CPU
cudaMemcpy(c,cd,size, cudaMemcpyDeviceToHost );

3.2 Optimization
Optimization is one of the rather difficult parts of CUDA but definitely an important to achieve a good performance. This section gives a brief idea about common kind of optimization can be used or rather should be used for achieving a good performance. This part doesn’t give the details of optimization used for particular application. The details are mentioned in the upcoming section 4 for specific benchmark as we move further.

The optimization for each Parsec benchmark depends on specific requirement, kind of data involved and the size of input. But the principle for choosing optimization to the CUDA code depends on following:

- Optimize use of on-chip memory to reduce bandwidth usage and redundant execution
- Group threads to avoid SIMD penalties and memory port/bank conflicts.
- Threads within a thread block can communicate via synchronization, but there is no built-in global communication mechanism for all threads.

The optimization CUDA involves many different techniques but it is important to find out what techniques are best suited for particular application. The following optimization techniques present a brief idea of optimization:

Global Memory Throughput (Memory Coalescing): Reducing transaction sizes to half from actual size: Memory access is handled per-wraps. So, we can carry out smallest possible number of transactions. Reducing transaction size when is the first options. The details are same as the paper by [4].

Launching Configuration: This is way of optimizing by launching sufficient threads to hide latency. Use of threads depends on the access pattern (global, shared or texture) but application can be launched upto 512 threads which is sufficient. Like using global memory, it is always wise to use maximum of 512 threads but when it comes to shared or particularly texture, using 512 threads will not make a huge difference. The reason can be seen in NVIDIA programming guide.

Device Memory: This is perhaps the most important optimization for CUDA. Many applications are memory bound. So, it is important to use right memory. There are five different memories available in GPU: global, shared, local, constant and texture. Our experiment shows that shared memory is best memory to be used as it allows per block data reuse which in true in our case for many applications. It caches data to reduce global memory accesses and avoid non-coalesced access to provide serialization. If n threads access the same bank, then n accesses can be executed serially (n threads access the same word in one fetch). Rather than using only global memory, we divide the work between global memory and shared memory or texture memory. This affects the performance to much extend. Before going into the application details, let us see the memory hierarchy available in GPU.

Global memory is a large memory accessible by all threads. All device kernel code and data must fit within global memory. The global memory on the Tesla T10 is has a 128KB block size, and a latency of 600-900 cycles.

Shared memory is local to each SM, and data in shared memory may be shared between threads belonging to the same thread block. If multiple blocks are scheduled to the same SM, shared memory will be evenly partitioned between them. The size of the shared memory is 16KB per SM on the Tesla. Each memory is 16-banked and has 4 cycle latency.

Local memory provides the same latency benefits as shared memory, but is local to the SPs and can only be accessed on a per-thread basis. Local memory is typically used as a fast scratchpad.

Constant memory is a special partition of global memory meant to hold constant values. It is read-only from the device scope and can only be written to by the host. The constant memory is 2-level, with distributed, hardware-managed caches in each SM. Because the constant memory is read-only, these caches do not have any coherence protocol. On the Tesla, each constant cache is 16KB and has low access latency.

Texture memory is similar to constant memory, but also includes support for special optimized texture transformation operations. This extra support comes at the cost of higher latency than constant memory for normal read operations.

4. Applications: Details, CUDA implementation, Results and Optimization

3.1. CUDA Implementation: Transformation

3.1.1 Blackscholes: Converted from original tbb version. Used for financial analysis. It computes prices of portfolio of European Options with the Blackscholes PDE.

The value of a call option for a non-dividend paying underlying stock

\[ C(S,t) = SN(d_1) - Ke^{-rt}N(S,t) \]

\[ d_1 = \frac{ln(S/R) + (r + \sigma^2/2)T}{\sigma\sqrt{T}} \]

\[ d_2 = \frac{ln(S/R) + (r - \sigma^2/2)T}{\sigma\sqrt{T}} \]

\[ P(S,t) = Ke^{-rt} - S + SN(d_1) - Ke^{-rt}N(d_2) \]
where \( N(\cdot) \) is the cumulative distribution function of the standard normal distribution, \( T - t \) is the time to maturity, \( S \) is the spot price of the underlying asset, \( K \) is the strike price, \( r \) is the risk free rate and \( \delta \) is the volatility of the underlying asset.

For CUDA implementation, blackscholes stores the portfolio with numOptions derivatives in array OptionData. The program includes file option-Data.txt which provides the initialization and control reference values for 1,000 options which are stored in array data init. The initialization data is replicated if necessary to obtain enough derivatives for the benchmark. The CUDA program was written in global memory. The program divides the portfolio into a number of work units equal to the number of threads and processes them concurrently. The transformation involves generation random input data on host, transferring data to GPU, compute prices on GPU, transfer prices back to host, compute prices on CPU and check the results. Each options is calculated independently and each thread iterates through all derivatives in its contingent and calls function BlkSklhEqEuroNoDiv for each of them to compute its price. A simple implementation of the Blacksholess algorithm would assign each thread to a specific index of input data. But there are some hardware constraints to be taken into account. Three dimensions on are only 16-bit (i.e. the maximum size of each dimension is 65536 and maximum number of threads per block is 512. [See Appendix 3.2 for CUDA code.]

**Initial Result:**

```bash
[ray@coil release]$ blackscholes Size = 65100 options threads = 512
Throughput = 3.6903 GOptions/s Time = 1.32400 ms
Time transfer CPU to GPU = 0.4998 ms
GPU Kernel execution = 0.0425 ms
Time transfer GPU to CPU = 0.737 ms
```

**Overheads and Optimization:**
The main problem in blackscholes is the transfer time between host to device memory and vice versa. This may be due to fact that the PDE equations which calculates option pricing involves intense calculation and it needs data input for that. But, blackscholes doesn’t need all data at single time but it does access same data again and again. It is important to reduce the transfer time by some optimization method like global memory coalescing or page locked memory allocation.

For CUDA optimization, blackscholes throughput is memory bounded. So the memory access and the transfer between the device and host should be kept minimum. A technique called memory coalescing where reads and writes within each warp can be arranged sequentially, so that all memory requests can be coalesced into a single continuous block with base address aligned to 16 byte boundaries for best performance. Coalescing means that a memory read by consecutive threads in a warp is combined by the hardware into several, wide memory reads. The requirement is that the threads in the warp must be reading memory in order. For example, if you have a float array called data[ ] and we want to read many floats starting at offset n, then thread 0 in the warp must read data[n], thread 1 must read data[n+1], and so on. Those 32-bit reads, which are issued simultaneously, are merged into several 384 bit reads in order to efficiently use the memory bus. Coalescing is a warp-level activity, not a thread-level activity. Normally 16 different threads can be read from the global memory. Interleaved access to global memory by threads in a thread block is essential to exploit this architectural feature. Depending on the use of global memory and registers, the optimal number of threads after doing memory coalescing for maximum performance is in the 192-256 range. Here we can reduce the transaction size depending on all threads whose request address lies in same segment.

In GPU, execution of a program proceeds by distributing the computations across threads blocks and across threads within a thread block. Before, going into memory coalescing check it is important to check if there is any data reuse and if it does then shared memory is used to re-order non coalescing addressing. The global memory access by all threads of a half-warp is coalesced into a single memory transaction as soon as the words accessed by all threads lie in the same segment of size equal to 32 bytes if all threads access 1-byte words, 64 bytes if all threads access 2-byte words, 128 bytes if all threads access 4-byte or 8-byte words. Coalescing is achieved for any pattern of addresses requested by the half-warp, including patterns where multiple threads access the same address. This is in contrast with devices of lower compute capabilities where threads need to access words in sequence. If a half-warp addresses words in n different segments, n memory transactions are issued (one for each segment), whereas devices with lower compute capabilities would issue 16 transactions as soon as n is greater than 1. In particular, if threads access 16-byte words, at least two memory transactions are issued. Unused words in a memory transaction are still read, so they waste bandwidth. To reduce waste, hardware will automatically issue the smallest memory transaction that contains the requested words. For example, if all the requested words lie in one half of a 128-byte segment, a 64-byte transaction will be issued. More precisely, the following protocol is used to issue a memory transaction for a half-warp.

**Coalescing Algorithm:**
- Find the memory segment that contains the address requested by the lowest numbered active thread.
- Find all other active threads whose requested address lies in the same segment.
- Reduce the transaction size, if possible:
  - If the transaction size is 128 bytes and only the lower or upper half is used, reduce the transaction size to 64 bytes;
  - If the transaction size is 64 bytes and only the lower or upper half is used, reduce the transaction size to 32 bytes.
- Carry out the transaction and mark the serviced threads as inactive. Repeat until all threads in the half-warp are serviced.

![Fig 3a: Coalesced access in which all threads but one access the corresponding word in a segment](image)

![Fig 3b: Non - Coalesced access in which all threads but one access the corresponding word in a segment](image)
In the above code, all the options of price is accessed or read in order from global memory by each thread of a half-warp during the execution of a single read is coalesced. [See appendix 3.1 for coalesced code]

Finally, higher performance for data transfers between host and device is achieved by using page-locked host memory. In addition, when using mapped page-locked memory, there is no need to copy to or from device memory. Data transfers are implicitly performed each time the kernel accesses the mapped memory. For maximum performance, these memory accesses must be coalescend like if they were accesses to global memory. Assuming that the mapped memory is read or written only once, using mapped page-locked memory instead of explicit copies between device and host memory. CUDA 3.0 capabilities allow a kernel to directly access host page-locked memory – no copy to device needed. It is useful when we cannot predict what data is needed but it is less efficient if all data will be needed. Explained in [11] and [12].

3.2. Streamcluster: It solves online clustering problem [13].
This is the problem of finding a partition of a data set so that data set can be cluster under some definition of similarity. The particular definition of clustering that is the focused is the k-Median objective that of identifying k centers so that the sum of distances from each point to its nearest center is minimized. The function computes how much cost can be saved by opening a new center. For every new point, it weighs the cost of making it a new center and reassigning some of the existing points to it against the savings caused by minimizing the distance between two points x and y for all point

For CUDA implementation, a given point x, find the cost of the function: open a facility at x for points y such that the assignment distance of y exceeds dist(y, x), make y a member of x. Then for facilities y such that reassigning y and all its members to x would save cost, realize this closing and reassignment. If the cost of this operation is negative, perform this operation and return the amount of cost saved otherwise, do nothing, numcenters will be updated to reflect the new number of centers. z is the facility cost, x is the number of this point in the array points. The minimum distance for saving is given by

\[ d(x, y) = |x - y|^2 \]

The CUDA transformation code is given in Appendix 3.2 at page

Initial Result:
Input data: 8,192 input points, 10-20 centers, 64 dimensions, 8 threads

```
[raya@cell release]$ time -11.769146
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Overhead and Optimization: The streamcluster spends most time in executing the kernel i.e. executing GPU kernel. It makes sense as the algorithm itself continually need to reuse data for calculating the distance, open new centers, determines the cost for opening and updating new centers at every run. So, shared memory is a good choice to maxmize the per-block data reuse. First, shared memory caches data to reduce global memory access and second it avoids non-coalesced access. Since streamcluster performance bottlenecks is from kernel call overhead and CPU-GPU transfer, shared memory helps to access data simultaneously, thus achieving high memory bandwidth.

The global memory generates a large number of thread blocks, which is favorable for thread scheduling. However, in streamcluster the data set is densely distributed in clusters. So, if we want continuously compare the points using distance vector and creating new center and updating center table, there are large amount a data sharing between blocks, particularly as the size of the input point increases. Therefore, it incurs a large amount of read overhead because each thread must search to compare of the input points. Because of these data sharing between blocks, it is difficult to coordinate threads in different blocks; it is not favorable to use global memory. For sparse data sets it might be worthwhile to store
the dataset in cacheable constant memory, however for dense data sets it could be difficult to find enough data points in range to completely populate a warp, leading to performance degradation from unutilized processors.

While CUDA does not provide a hardware-managed cache for global memory, it is desirable to temporarily write the output distance values to shared memory as they must be read multiple times in each block and data reuse. However, CUDA does not provide any cache for global memory. While the constant & texture memory partitions are cacheable, these memories are read-only, and thus cannot be used to store the output which accounts for half of all memory references. So, shared memory is a good choice.

To overcome the lack of caching on the global memory, we use the per-SM shared memory as a software-managed cache. In streamcluster, a thread block is assigned to each point on the output grid and reads from any input samples as a cache creates many challenges for kernels in which data may be shared between different thread blocks.

The shared memory space is much faster than the local and global memory spaces because it is on-chip. In fact, for all threads of a warp, accessing the shared memory is as fast as accessing a register as long as there are no bank conflicts between the threads and it can be accessed by all threads within a thread block simultaneously. The shared memory space is divided into equal-sized memory modules called banks, which can be accessed in parallel. In the NVIDIA, the shared memory is divided into 16 banks. Successive 32-bit words are assigned to successive banks. So, any memory read or write request made of n addresses that fall in n distinct memory banks can be serviced simultaneously; yielding an effective bandwidth that is n times as high as the bandwidth of a single module.

For a given point x, the value of x can be accessed and shared by any other threads and it can be simultaneously read by each threads.

However if n threads of a half-warp access the same bank at a time, there is an n-way bank conflict, resulting in n sequential accesses to the shared memory. The hardware splits a memory request with bank conflicts into as many separate conflict-free requests as necessary, decreasing the effective bandwidth by a factor equal to the number of separate memory requests. For devices of compute capability 3.0, the warp size is 32 and the number of banks is 16, a shared memory request for a warp is split into one request for the first half of the warp and one request for the second half of the warp. As a consequence, there can be no bank conflict between a thread belonging to the first half of a warp and a thread belonging to the second half of the same warp. If an instruction executed by a warp writes to the same location in shared memory for more than one of the threads of the warp, only one thread per half-warp performs a write.

In the above case of CUDA code from the example, threads tid and tid+n access the same bank whenever s•n is a multiple of the number of banks (i.e. 16) or, equivalently, whenever n is a multiple of $16/d$ where d is the greatest common divisor of 16 and s. As a consequence, there will be no bank conflict only if half the warp size (i.e. 16) is less than or equal to $16/d$, that is only if d is equal to 1, i.e. s is odd.

![Fig 5: Bank without conflicts](image)

**The following part shows how to define shared memory:**

```c
__shared__ double shared[32];
float data = shared[BaseIndex + s * tid];
```

This implementation in streamcluster CUDA is as follows:

```c
extern __shared__ float coord_s[32];
// coordinate mapping of point[x] to shared mem //
if(threadIdx.x == 0)
for(int i=0; i<dim; i++) {
    coord_s[i] = coord_d[i*num + x]
}
```

Accesses that are larger than 32-bit per thread are split into 32-bit accesses that typically generate bank conflicts. For example, there are 2-way bank conflicts for arrays of **doubles** accessed as follows:

```c
__shared__ double shared[32];
double data = shared[BaseIndex + tid];
```

One way to avoid bank conflicts in this case is to split the double operands like in the following sample code:

```c
__shared__ int shared_lo[32];
__shared__ int shared_hi[32];
double dataIn;
shared_lo[BaseIndex + tid] = __double2loint(dataIn);
shared_hi[BaseIndex + tid] = __double2hiint(dataIn);
```

Finally, shared memory compute capability 3.0 offers a special hardware optimization in the GPU architecture that enables broadcast of the word to all threads simultaneously when servicing the dataset in cacheable constant memory, however for dense data sets it could be difficult to find enough data points in range to completely populate a warp, leading to performance degradation from unutilized processors.

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To overcome the lack of caching on the global memory, we use the per-SM shared memory as a software-managed cache. In streamcluster, a thread block is assigned to each point on the output grid and reads from any input samples as a cache creates many challenges for kernels in which data may be shared between different thread blocks.

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```c
__shared__ double shared[32];
float data = shared[BaseIndex + s * tid];
```

This implementation in streamcluster CUDA is as follows:

```c
extern __shared__ float coord_s[32];
// coordinate mapping of point[x] to shared mem //
if(threadIdx.x == 0)
for(int i=0; i<dim; i++) {
    coord_s[i] = coord_d[i*num + x]
}
```

Accesses that are larger than 32-bit per thread are split into 32-bit accesses that typically generate bank conflicts. For example, there are 2-way bank conflicts for arrays of **doubles** accessed as follows:

```c
__shared__ double shared[32];
double data = shared[BaseIndex + tid];
```

One way to avoid bank conflicts in this case is to split the double operands like in the following sample code:

```c
__shared__ int shared_lo[32];
__shared__ int shared_hi[32];
double dataIn;
shared_lo[BaseIndex + tid] = __double2loint(dataIn);
shared_hi[BaseIndex + tid] = __double2hiint(dataIn);
```

Finally, shared memory compute capability 3.0 offers a special hardware optimization in the GPU architecture that enables broadcast of the word to all threads simultaneously when servicing the dataset in cacheable constant memory, however for dense data sets it could be difficult to find enough data points in range to completely populate a warp, leading to performance degradation from unutilized processors.

While CUDA does not provide a hardware-managed cache for global memory, it is desirable to temporarily write the output distance values to shared memory as they must be read multiple times in each block and data reuse. However, CUDA does not provide any cache for global memory. While the constant & texture memory partitions are cacheable, these memories are read-only, and thus cannot be used to store the output which accounts for half of all memory references. So, shared memory is a good choice.

To overcome the lack of caching on the global memory, we use the per-SM shared memory as a software-managed cache. In streamcluster, a thread block is assigned to each point on the output grid and reads from any input samples as a cache creates many challenges for kernels in which data may be shared between different thread blocks.

The shared memory space is much faster than the local and global memory spaces because it is on-chip. In fact, for all threads of a warp, accessing the shared memory is as fast as accessing a register as long as there are no bank conflicts between the threads and it can be accessed by all threads within a thread block simultaneously. The shared memory space is divided into equal-sized memory modules called banks, which can be accessed in parallel. In the NVIDIA, the shared memory is divided into 16 banks. Successive 32-bit words are assigned to successive banks. So, any memory read or write request made of n addresses that fall in n distinct memory banks can be serviced simultaneously; yielding an effective bandwidth that is n times as high as the bandwidth of a single module.

For a given point x, the value of x can be accessed and shared by any other threads and it can be simultaneously read by each threads.

However if n threads of a half-warp access the same bank at a time, there is an n-way bank conflict, resulting in n sequential accesses to the shared memory. The hardware splits a memory request with bank conflicts into as many separate conflict-free requests as necessary, decreasing the effective bandwidth by a factor equal to the number of separate memory requests. For devices of compute capability 3.0, the warp size is 32 and the number of banks is 16, a shared memory request for a warp is split into one request for the first half of the warp and one request for the second half of the warp. As a consequence, there can be no bank conflict between a thread belonging to the first half of a warp and a thread belonging to the second half of the same warp. If an instruction executed by a warp writes to the same location in shared memory for more than one of the threads of the warp, only one thread per half-warp performs a write.

In the above case of CUDA code from the example, threads tid and tid+n access the same bank whenever s•n is a multiple of the number of banks (i.e. 16) or, equivalently, whenever n is a multiple of $16/d$ where d is the greatest common divisor of 16 and s. As a consequence, there will be no bank conflict only if half the warp size (i.e. 16) is less than or equal to $16/d$, that is only if d is equal to 1, i.e. s is odd.

![Fig 6: Bank with conflicts](image)
one memory read request. This reduces the number of bank conflicts when several threads of a half-wrap read from an address within the same 32-bit word.

The CUDA transformation and optimization code is given in Appendix 3.2 at page.

![Diagram of CUDA transformation and optimization]

**Fig 7:** Framework for optimization using shared memory

### 3.3. Ferret: This application is based on the Ferret toolkit [15] which is used for content-based similarity search of feature-rich data such as audio, images, video, 3D shapes and so on. It is a search engine which finds a set of images similar to a query image by analyzing their contents. The input is an image database and a series of query images. It involves breaking the images into segments in order to distinguish different objects in the image. The segmented images use feature extraction which mathematically describe content of segment. This is then indexed (selecting images that matches the query images) and ranked (ranking according to order of similarity). The four stages involving ferret are query image segmentation, feature extraction, indexing of candidate sets with hashing and ranking using with finding nearest neighbor search using distance vector algorithm.

Segmentation is the process of decomposing an image into separate areas which display different objects. The rationale behind this step is that in many cases only parts of an image are of interest, such as the foreground. Segmentation allows the subsequent stages to assign a higher weight to image parts which are considered relevant and seem to belong together.

After segmentation, ferret extracts a feature vector from every segment. A feature vector is a multi-dimensional mathematical description of the segment contents. It encodes fundamental properties such as color, shape and area. Once the feature vectors are known, the indexing stage can query the image database to obtain a candidate set of images. This method uses hash functions which map similar feature vectors to the same hash bucket with high probability. It then employs a step-wise approach which indexes buckets with a higher success probability first. After a candidate set of images has been obtained by the indexing stage, it is sent to the ranking stage which computes a detailed similarity estimate and orders the images according to their calculated rank. The similarity estimate is derived by analyzing and weighing the pair-wise distances between the segments of the query image and the candidate images. For a given query image X and database image Y, the ferret employs the Earth Mover's Distance (EMD) given by

$$\text{EMD} (X,Y) = \min \sum_i \sum_j f_{ij} d(X_i,Y_j)$$

To implement this algorithm in CUDA, we took the original ferret-parallel.c from the Parsec 2.1 ferret and then we assign the distance calculation of each data point to a single thread for analyzing and weighing the minimum distances between the segments of the query image and the candidate images using blocks of thread in parallel. Then the distance for each test point was sorted in parallel. Therefore each thread loop over all the initial stored data, calculating its assigned data point’s distance with query data, finding the minimum distance from its data point to a stored images using feature extraction for best match and filtering out the few matches. When all threads are finished doing this, the membership of each data point is known, copy result back from GPU to CPU.

**Result:**

[raya@coil release]$ ferret
Throughput - 31.45 GB/s, Time - 12.11s, Size - 35000 threads - 32
CPU-GPU Transfer Time - 5.71s
Calculating the distance - 3.23s
Filtering - 1.04s

**Overhead and Optimization:** The GPU-based implementation of ferret is clearly memory bound as there is more memory access. The algorithm for this pass requires intensive data sharing, and creates congestion on the memory subsystem. For example, one possible way of parallelizing this pass is to ask all threads whose data point belongs in a same group to add their values to a certain data structure in memory.

Ferret is good candidate of shared memory as ferret needs to access only the values in a correlated section of memory. In ferret, calculating distance means we need to load the current image to find the distance between the already stored images to find the best possible match. So, all the threads within the block calculate the distance. For memory throughput, the time for CPU-GPU transfer and the GPU execution holds equally for overheads. When this is the case, shared memory becomes the most effective memory to use.
Shared memory allows all threads within a block to share information. First, data is read from global memory into shared memory (accessing the images to calculate the distance in each threads and finding which one is the best match), ensuring that consecutive threads in a half-warp read from consecutive memory addresses to facilitate non-coalesced memory access. Then calculations are performed using only shared memory. Finally, the results are written to global memory. The general shared memory optimization implementation is same as we did in streamcluster avoiding all bank conflicts. Here initially, we compute the distance and filtered and stored matched images in shared memory. The process was repeated several times till we get the top 10 images and finally this top 10 images is stored in global memory. Here, we are explicitly using shared memory because just we dividing kernel into global memory for flushing the data and reading the data into shared memory and storing the value of kernel call arguments could cause for extra overhead due to limit in resources of 16KB of shared memory and limited registers. Any bank conflict in shared memory is eliminated in same way as the streamcluster.

3.4. Swaptions: A financial analysis application to price the portfolio of the swaptions. It solves a PDE equation and stores the portfolio in swaptions array. The swaptions application uses the Heath-Jarrow-Morton (HJM) framework [18] to price a portfolio of swaptions. The HJM framework describes how interest rates evolve for risk management and asset liability management for a class of models between the option seller and the option buyer whereby the option buyer is granted a right secured by the option seller, to carry out some operation at some moment in the future. Its central insight is that there is an explicit relationship between the drift and volatility parameters of the forward rate dynamics in a no arbitrage market. Swaptions therefore employs Monte Carlo (MC) simulation to compute the prices. The program stores the portfolio in the swaptions array. Each entry corresponds to one derivative. Swaptions partitions the array into a number of blocks equal to the number of threads and assigns one block to every thread.

The price of the underlying options with constant drift $\mu$ and volatility $\nu$: $dS = \mu S dt + \nu S dW_t$ (where $W_t$ is Wiener random process). Details of the equation given in [18].

For CUDA implementation, swaptions array is partitioned into number of work units threads and processed concurrently. Device memory is allocated to run compute to price. Each thread block is iterated and calls for HJM Swaption Blocking function for every entry to compute prices. It generates a random HJM path for each run. Once the path is generated, we compute the value of swaptions (expected value and confidence width). Each thread computes and sums the multiple simulation paths and stores the sum and the sum of squares into a device memory. The number of option computed is one option per thread. Then, computed sum is put back to the host memory.

```
float r = d_Random[pos];
float endStockPrice = S*exp(MuByT+BySqrtT * r);
float callProfit = fmaxf(endStockPrice - X, 0);
sumCall.Expected += callProfit;
sumCall.Confidence += callProfit*callProfit
```

Initial Result:
[raya@coil release]$  
Options : 256  
Simulation paths: 14100  
Time (ms) : 6.45204  
Kernel = 1.17525  
CPU to GPU memory = 2.142637  
GPU to CPU memory = 1.766440

Overhead and Optimization: The swaptions generates a HJM path after being implemented in CUDA and we use them to compute an expected value and pricing for the underlying option. There are multiple ways we could go about computing the mean of all of the samples. The number of options is typically in the hundreds or fewer, so computing one option per thread will likely not keep the GPU efficiently occupied. Therefore, we concentrate on using multiple threads per option. Given that, we have two choices: we can either use one thread block per option, or multiple thread blocks per option or we can use combination of both. When computing a very large number of paths per option, it will probably help us to reduce the latency of reading the random input values if we divide the work of each option across multiple blocks.

Each thread computes and sums the multiple simulation paths and stores the sum and the sum of squares into a device memory array. To compute the expected price for each option, we sum all values per option. To do so, we launch a second kernel with a process called parallel reduction [8] to compute the sums. A tree based approach used within each thread block where we synchronize across all threads once each block produces result. By doing a lot of computation per thread, we were able to reduce overhead such as the cost of kernel launching and storage to device memory. So, the process begins with storing intermediate results of the options to global memory, finish the first kernel, and then launch the parallel reduction kernel to compute the final sum. Indeed, we compute these final values using a single thread block per option by storing each thread in shared memory instead of global memory, and the parallel reduction is performed in shared memory. This saves a global store per thread and an extra kernel invocation.

![Fig 8: Parallel Reduction](image)

<table>
<thead>
<tr>
<th>Options</th>
<th>Simulation paths</th>
<th>Time (ms)</th>
<th>Kernel</th>
<th>CPU to GPU memory</th>
<th>GPU to CPU memory</th>
</tr>
</thead>
<tbody>
<tr>
<td>256</td>
<td>14100</td>
<td>6.45204</td>
<td>1.17525</td>
<td>2.142637</td>
<td>1.766440</td>
</tr>
</tbody>
</table>
The CUDA code for Parallel reduction for Address Interleaving:

```c
__device__ void sumReduceSharedMem(volatile T *sum, volatile T *sum2, int tid)
{
    // do reduction in shared mem
    if (blockSize >= 64) sum[tid] += sum[tid + blockDim.x];
    __syncthreads();
    if (blockSize >= 128) sum[tid] += sum[tid + blockDim.x * 2];
    __syncthreads();
    if (blockSize >= 256) sum[tid] += sum[tid + blockDim.x * 4];
    __syncthreads();
    if (tid == 0) g_odata[blockIdx.x] = sdata[0];
}
```

3.5. x264: The x264 application is an H.264/AVC (Advanced Video Coding) video encoder. H.264 describes the lossy compression of a video stream [13] which provides many improved features. The advancements allow H.264 encoders to achieve a higher output quality with a lower bit-rate. The flexibility of H.264 allows its use in a wide range of contexts with different requirements, from video conferencing solutions to high-definition (HD) movie distribution. Next generation HD DVD or Blu-ray video players already require H.264/AVC encoding. H.264 encoders and decoders operate on macroblocks of pixels which have the fixed size of 16x16 pixels. Various techniques are used to detect and eliminate data redundancy. The most important one is motion estimation process [9]. Motion compensation is usually the most expensive operation that has to be executed to encode a frame. Motion estimation algorithm exploits redundancy between frames, which is called temporal redundancy. A video frame is broken down into macroblocks (each macroblock typically covers 16x16 pixels), each macroblock’s movement from a previous frame (reference frame) is tracked and represented as a vector, called motion vector. Storing this vector and residual information instead of the complete pixel information greatly reduces the amount of data used to store the video.

Motion Estimation is the process of finding the lowest cost (in bits) way of representing a given macroblock’s motion. In video, pixel values often do not change much across scenes (frames), and the pixel values of a block can be predicted from the values of a similar block in a previous frame (reference frame). The motion estimation operates to find a motion vector, which indicates the similar block in the reference frame, and residuals to compensate prediction errors.
To find the motion vector, the motion estimation algorithm finds the best match in a certain window of the reference frame. The block coded with motion estimation is called inter-coded. This fact is used to aid in the compression process. Rather than transmitting the entire motion vector for a given macroblock, we only need to transmit how it differs from what we predict it would be, given how the blocks around it moved. The motion vector prediction (MVp) is the median of these three vectors. We only need to transmit how a given block’s motion vector differs from the MVp. To do that they account the cost of encoding the residual as well as the cost of encoding the motion vector as a prime area of estimating cost. In order to solve the problem of performing motion estimation in a massively parallel environment we needed to deal with the issue of needing the MVp to find the optimal motion vector. The whole explanation is beyond the scope of this project. Please refer paper [20] and [21] for further details.

For CUDA implementation of x264, we used global memory for part of input data, since data use has 2D locality. x264 required more modification when imported to CUDA, involved large scale code transformation from the Parsec parallel application code. In CUDA, the problem of needing the MV is solved by properly estimating the cost of a given motion vector by using an estimate of what the MVp will actually be like. This cost calculation uses a evaluation metrics called Sum of Absolute transferred difference [refer wikipedia]: cost=SATD+[C(MV_p−MV_p)+C(MV−MV_p)] where C(.) is the cost of encoding a motion vector differential of that length. In our implementation, a macroblock is divided into sixteen 4x4 blocks, and the SAD value of each 4x4 block is calculated in parallel for all candidate motion vectors (positions) within the search range on the reference frame. We then merge these 4x4 block SADs to form the 4x8, 8x4, 8x8, 16x8, and 16x16 block SADs, respectively. For each block size, we compare the SADs of all candidate MVs and the one with the least SAD is the integer-pixel motion vector (IMV), MV with integer pixel accuracy. Next, in order to obtain the fractional pixel motion vectors, the reference frame is interpolated using a filter as defined in the H.264/AVC standard. We calculate the SADs at 24 fractional pixel positions that are adjacent to the best integer MV, and then choose the least SAD position as the fraction-pixel motion vector (FMV).

Now one block per macroblock is assigned. We had one thread check each search position within the search window. By having one block per macroblock we were able to take advantage of thread synchronization to perform a reduction to find the lowest cost motion vector for each macroblock.

Initial Result:
[raya@coil release]$x264 Loaded 'pixel_size.pgm', '640 x 340 pixels'
Processing time: 0.118000s
81.56 pixels/sec
GPU execution = 0.0306800s
CPU-GPU transfer = 0.0631000s

Overhead and Optimization: Our implementation of x264 is bounded by both GPU execution and CPU-GPU transfer but it looks like GPU-CPU communication holds for overhead in overall computation time than GPU execution. While x264 CUDA implementation does not take advantage of shared memory to preload the needed pixel values, it does store both the current frame and the reference (previous) frame in global memory. So, x264 regularly need to access neighboring memory locations with 2D spatial locality, using texture memory instead of global memory is useful for performance increase. Global memory doesn’t provide caching; it is desirable to use texture memory. Reading device memory through texture fetching present some benefits that can make it an advantageous alternative to reading device memory from global or constant memory:

- If the memory reads do not follow the access patterns that global or constant memory reads must respect to get good performance, higher bandwidth can be achieved providing that there is locality in the texture fetches.
- Addressing calculations are performed outside the kernel by dedicated units.
- Packed data may be broadcast to separate variables in a single operation.

Now, every multiprocessor on a GPU has its own texture cache, originally designed for storing images that are reproduced many times to create the illusion of a textured object. Tesla T1 provides support for special optimized texture transformation operations with texture memory up to size of global memory which the largest size among all memories and clock cycle of > 100 cycles. This extra support comes at the cost of higher latency than constant memory for normal read operations. Whenever an element of the texture data type is read from global memory it is stored in the texture cache on the device, and all subsequent requests for this element do not need to read from global memory. As texture memory space resides in device memory and is cached in texture cache, so a texture fetch costs one memory read from device memory only on a cache miss, otherwise it just costs one read from texture cache. The texture cache is optimized for 2D spatial locality, so threads of the same warp that read texture addresses that are close together in 2D will achieve best performance. Also, it is designed for streaming fetches with a constant latency; a cache hit reduces DRAM bandwidth demand but not fetch latency. While the constant & texture memory partitions are cacheable, these memories are read-only, and thus cannot be used to store the output which accounts for half of all memory references. As, in x264, we need to predict the successive blocks depending on the previous block, it is important to store the output partially somewhere. So, we used shared memory as a good option as all the threads within a specific block are able to share information. All of the main calculations used for processing occur within shared memory, and then all of the results are transferred to texture memory.

// declare texture reference for 2D float texture:// texture<quint8_t,2,cudaReadModeElementType> currentTex;

Initially, when the pixel size is small, 16834 register was enough for SAD calculation in texture but with large pixel size and frame a lack of registers restricts the number of threads that could be scheduled, exposing the latency of texture memory. We also achieve potential throughput using loop unrolling. Loop unrolling is the most common optimization in compiler construction. Loop unrolling reduces the operations which is not part of core data computation. By unrolling the X loop by 4, we can compute MV_{px}, MV_{py}, MV_{py}, MV_{py} once and use it multiple times, much like the CPU version of
the code does not affect shared memory usage and a register is saved by removing the unrolled loop’s induction variable.

The following code shows for shared memory usage and loop unrolling in x264:

```c
__shared__ float As[16][16];
__shared__ float Bs[16][16];

// load input tile elements
As[MVx][MVy] = A[indexA];
Bs[MVpx][MVpy] = B[indexB];
indexA += 16;
indexB += 16 * widthB;

__syncthreads();
// compute results for tile
Ctemp = As[MVx][0] * Bs[0][MVy];
Ctemp += As[MVpx][15] * Bs[15][MVpy];
__syncthreads();
C[indexC] = Ctemp;

//loop unrolling
float dx1 = (abs(dx))<<2;
float dx2 = (abs(dx))<<2;
float dx3 = (abs(dx))<<2;
float dx4 = (abs(dx))<<2;

int x1Cost = round((log2f(dx1)*2+0.718f+!!dx)+ .5f);
int x2Cost = round((log2f(dx1)*2+0.718f+!!dx)+ .5f);
int x3Cost = round((log2f(dx1)*2+0.718f+!!dx)+ .5f);
int x4Cost = round((log2f(dx1)*2+0.718f+!!dx)+ .5f);
```

3.6. Fluidanimate: This Intel RMS application uses an extension of the Smoothed Particle Hydrodynamics (SPH) method to simulate an incompressible fluid for interactive animation purposes [22]. Its output can be visualized by detecting and rendering the surface of the fluid. The force density fields are derived directly from the Navier-Stokes equation which is written as follows:

\[
\rho (\frac{\partial v}{\partial t} + v \cdot \nabla v) = -\nabla p + \rho g + \mu \nabla^2 v
\]

where \(v\) is a velocity field, \(\rho\) is a density field, \(p\) a pressure field, \(g\) an external force density field and \(\mu\) the viscosity of the fluid.

The density at a location \(r\) can be calculated by equation:

\[
\rho_j(r) = \sum_j m_j W(r-r_j,h)
\]

Applying the SPH interpolation equation to the pressure term and the viscosity term of the Navier-Stokes equation yields the equations for the pressure and viscosity forces

\[
f_p = -\sum_j m_j \frac{p_i + p_j}{2\rho_j} \nabla W(r-r_j,h)
\]

\[
f_v = -\mu \sum_j m_j \frac{v_i + v_j}{\rho_j} \nabla^2 W(r-r_j,h)
\]

For CUDA implementation, it follows same principle as blackscholes each parameters like velocity field, density field, pressure field, external force density field and viscosity of the fluid are computed in device memory with threads and the final value is again copied back to host memory. All the functions use global memory. The fluidanimate represents in 3D version for node nx, ny and nz which are the number of computational nodes in the x, y and z directions for a flow domain, respectively. The 3D domain of size \(nx \times ny \times nz\) is represented by a breadth \(nx\), width \(nx\) and height \(nz\) on the host side.

// number of grid cells in each dimension
int nx, ny, nz;
// cell dimensions
struct Vec3 delta;
```

Fig 10: Dimensional array x, y, z

On the GPU side, the representation is used to store data in global memory. This 3D mapping translates to efficient data transfer between the host and the device. Several matrices are needed to represent the pressure and velocity components at different time levels. Memory allocation on the device is done only once before starting the time stepping. In our implementation, the velocity field at time \(t\) depends only on the velocity field at \(t-1\). Five different matrices are used to represent the velocity fields. The matrices are swapped at the end of each time step for reuse GPU code is of five different kernels to implement the steps of the algorithm:

Rebuild spatial index Because the smoothing kernels \(W(r-r_h)\) have finite support \(h\), particles can only interact with each other up to the distance \(h\). The program uses a spatial indexing structure in order to exploit proximity information and limit the number of particles which have to be evaluated.

```c
__device__ void RebuildGridMT(int i, int nx, int ny, int nz, struct Vec3 delta, bool* border, Cell* cells, Cell* cells2, int* cnumPars, int* cnumPars2, Grid* grids)
```

Compute densities This kernel estimates the fluid density at the position of each particle by analyzing how closely particles are packed in its neighborhood. In a region in which particles are packed together more closely, the density will be higher.

```c
__device__ void ComputeDensities2MT(int i, float hSq, float densityCoeff, int nx, int ny, bool* border, Cell* cells, Cell* cells2, int* cnumPars, int* cnumPars2, Grid* grids)
```

Compute forces Once the densities are known, they can be used to compute the forces. The kernel evaluates pressure, viscosity and also gravity as the only external influence.

```c
__device__ void ComputeForcesMT(int i, float h, float hSq, float pressureCoeff, float viscosityCoeff, int nx, int ny, int nz, bool* border, Cell* cells int* cnumPars, int* cnumPars2, Grid* grids)
```
Handle collisions with scene geometry 

The next kernel updates the forces in order to handle collisions of particles with the scene geometry.

__device__ void ProcessCollisionsMT(int i, int nx, int ny, bool* border, Cell* cells, Cell* cells2, int* cnumPars, int* cnumPars2, Grid* grids)

Update positions of particles 

Finally, the forces can be used to calculate the acceleration of each particle and update its position.

Result:

Particles size: 100K
[yray@coil release]$ fluidanimate,512 Throughput = 8.3400 GB/s, Time = 0.005972 s, Size = 100000
Time transfer CPU to GPU = 0.001998 s
GPU Kernel execution = 0.002252 s
Time transfer GPU to CPU = 0.001737 s

Overheads and Optimization: Fluidanimate uses 5 kernels, 2 of which are further divided into further parts, has peak memory throughput of approximately 10.450 GB/sec, which corresponds to an optimal balance of just over 5 math operations per double precision value loaded from memory. Because most of loops perform a large amount of math, performance of code is mainly limited by memory bandwidth. The throughput is limited because of the global memory access for all 5 kernels. For the above domain, each GPU needs neighboring data computed by other GPUs which means all GPUs need to synchronize to exchange velocity and pressure fields at each time step. But a GPU cannot directly exchange data with another GPU. Hence, cells at the GPU domain decomposition boundaries needs to be copied back to the host, which adds an extra communication overhead to the overall computation in addition to the CUDA kernel launches at every time.

For Optimization, if threads in a warp read from the same cache line in the same cycle, these reads are batched into a single operation via a process known as memory coalescing for global memory which helps conserve bandwidth while reducing effective latency, therefore speedup the whole program. The concept is similar to loading an entire cache line from memory versus loading one word at a time at CPU. Coalescing operates at half-warp granularity, so uncoalesced loads and stores waste approx. 15/16ths of available memory bandwidth. Therefore the most important optimization for memory-bound applications is to arrange work so that threads in the same warp will access sequential memory locations at the same time. However, since 5 kernel runs at the same time, not all threads participated in coalescing. It is important to divide the memory coalesced access and non-coalesced access. For the non-coalesced, the access is benefited by using shared memory. 2 or 3 kernels can be used for the shared memory which involves data reuse and updates high arithmetic intensity. This will reduce the overhead and increase performance due to data transfer to the shared memory is largely compensated by this high arithmetic intensity.

CUDA optimization algorithm

- Coalesced the global memory access. The simultaneous global memory accesses by each thread of a half-warp during the execution of a single read or write instruction will be coalesced into a single access if:
  - The size of the memory element accessed by each thread is either 4, 8, or 16 bytes
  - The elements form a contiguous block of memory
  - The N_th element is accessed by the N_th thread in the half-warp
  - The address of the first element is aligned to 16 times the element’s size

- Read from the input of the global memory and copy the sub-domain from the global memory to the shared memory.
- Computation done by the threads, using data from the shared memory and write to the shared memory.
- Read from shared memory and the final result of the computation is written back to the global memory before exiting the kernel.

This back and forth data transfer between the global memory to the shared memory creates an overhead. But due to high arithmetic intensity of the kernel the overhead of data copying in order to benefit from the shared memory implementation.

3.7. Raytrace: The raytrace application is an Intel RMS workload which renders an animated 3D scene. Ray tracing is a technique that generates a visually realistic image by tracing the path of light through a scene. Rays through each pixel in an image plane are traced back to the light source. The scatter model describes what happens when a ray hits a surface. Raytracing works by projecting a ray of vision through each pixel on the screen from the observer. When the rays through the screen intersect an object, it is projected onto the screen. For every pixel on the screen, an equation involving the ray through that pixel from the observer and each object in the scene is solved to determine if the ray intersects the object. Then, the pixel through which the ray passes is set to the color of the intersected object at the point of intersection. If more than one object is intersected, then the closer of the intersections is taken.

Reflections and shadows are achieved through what is known as recursive ray tracing. When figuring out the color of the intersection, one important factor is whether that point is in shadow or not. To find this out, the same techniques are used, but instead of the normal location of the observer, the ray starts at the point of intersection and moves toward any light sources. If it intersects something, then the point is in shadow. When a reflective Surface is intersected, a new ray is traced starting from the point of intersection. The color that this ray returns is incorporated into the color of the original intersection.

![Raytracing method](image)

Fig 11: Raytracing method
Finding intersection points follows raytracer storage of the scene graph in a Bounding Volume Hierarchy (BVH). A BVH is a tree in which each node represents a bounding volume. The bounding volume of a leaf node corresponds to a single object in the scene which is fully contained in the volume. Bounding volumes of intermediate nodes fully contain all the volumes of their children, up to the volume of the root node which contains the entire scene. If the bounding volumes are tight and partition the scene with little overlap then a ray tracer searching for an intersection point can eliminate large parts of the scene rapidly by recursively descending in the BVH while performing intersection tests until the correct surface has been found.

![Raytracing algorithm for bounding volume hierarchy](Fig 12)

For CUDA implementation, BVH uses a transversal algorithm[17] The BVHs is built in top-down fashion with surface area heuristics using the centroids of bounding boxes for scene triangles, tracing packets of rays and only the node address is saved to the stack. As a BVH does not need to store the mint and maxv values along the ray, only the node address is saved to the stack. The algorithm maps one ray to one thread block a packet to a chunk. It traverses the tree synchronously with the packet and work on one node at a time and process whole packet against it. If the node is a leaf, it intersects the rays in the packet with the contained geometry. Each thread stored the distance to the nearest found intersection. If the node is not a leaf, the algorithm loads the two children and intersects the packets with both of them to determine the traversal order. Each ray determines which of the two nodes it intersects and in which it wants to go first by comparing the signed entry distances of both children. For packet traversal, the stack can be shared by all the threads in a packet, which increases the utilization of the resources. The order of traversal among several threads is resolved by a concurrent write to the shared memory. Each thread then writes the preference to one of four entries, value one for one of the four cases: traverse left, traverse right, traverse both, and traverse none. If at least one ray wants to visit the other node then the address of this other node is pushed onto stack. In case all rays do not want to visit both nodes or after the algorithm has processed a leaf, the next node is taken from the top of the stack and its children are traversed. If the stack is empty, the algorithm terminates.

### Initial Results and Optimization:

[raya@coil release]$ raytrace "loading scene" "512 x 512 pixels"
Time = 21.5672 s

5. Experimental Results:

In this part of the section we calculated the execution time theoretically to match it with experimental value and check the degree of the correctness.

As we calculated the speed up percentage for transformation from CPU to GPU and from GPU original to GPU optimized.

We check the experimental value while running the CUDA program on GPU. And then we split the time into various parts or into various run to check the time taken by individual part to execute the code. We calculated the execution time for the CUDA code without optimization. This time division helped us to look for overhead in the code and thus we were able to reduce or eliminate that bottleneck with optimization in the description above. The initial results for CUDA run without optimization is given in each benchmark above.

Now, Let $t_{\text{gpu}}$ is the execution time for running the kernel on gpu, $t_{\text{cpu transfer}}$ is the time taken for transfer from cpu to gpu i.e. from host to device, $t_{\text{gpu transfer}}$ is the time taken to copy from gpu to cpu, $t_{\text{gpu alloc}}$ be the time for gpu memory allocation and $t_{\text{gpu free}}$ is the time taken for gpu memory free

The total time taken to execute the CUDA code:

$$t_{\text{total}} = t_{\text{cpu transfer}} + t_{\text{gpu}} + t_{\text{gpu transfer}} + t_{\text{gpu alloc}} + t_{\text{gpu free}}$$

Although Gpu memory allocation and gpu memory free accounts for very less time, we can ignore the them in our calculation. To, show an example we took the execution time from blackscholes. It shows total executed time against the individual executed time. And we calculated the theoretical value as below:

**Blackscholes:**

- Size = 13200 options, threads = 512
- Throughput = 1.4260 GOptions/s
- Time = 0.443 ms
- Time transfer CPU to GPU = 0.197 ms
- GPU Kernel execution = 0.0125 ms
- Time transfer GPU to CPU = 0.2334 ms

**Calculation:**

Total execution time ($t_{\text{total}}$) = 0.443s [Experimental result]

Theoretical results: $t_{\text{total}} = t_{\text{cpu transfer}} + t_{\text{gpu}} + t_{\text{gpu transfer}}$

$$= 0.1871 + 0.0125 + 0.2134 = 0.413ms$$
So, the theoretical value and the experimental value is approximately same which proves our correctness of the program. The little time difference is because of may be some other time like the time taken for memory allocation, memory free and other calculation. Calculating all these is beyond our scope of the project.

Table 1: The following table showing the experimental time and calculated theoretical time for each application:

<table>
<thead>
<tr>
<th>Benchmarks</th>
<th>Experimental Value</th>
<th>Theoretical Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blackscholes</td>
<td>0.443ms</td>
<td>0.413ms</td>
</tr>
<tr>
<td>Streamcluster</td>
<td>8.360s</td>
<td>8.134s</td>
</tr>
<tr>
<td>Ferret</td>
<td>11.21s</td>
<td>10.683s</td>
</tr>
<tr>
<td>Swaptions</td>
<td>8.39ms</td>
<td>8.26ms</td>
</tr>
<tr>
<td>x264</td>
<td>51.1ms</td>
<td>44.5ms</td>
</tr>
<tr>
<td>Fluidanimate</td>
<td>7.69ms</td>
<td>7.35ms</td>
</tr>
<tr>
<td>Raytrace</td>
<td>21.56s</td>
<td>18.45s</td>
</tr>
</tbody>
</table>

The Speed up equation of CUDA execution time is given by,

Between CPU and GPU:

\[
\text{Speed Up} = \frac{CPU_{time} - GPU_{time}}{CPU_{time}} \times 100\%
\]

Between GPU original and GPU optimized:

\[
\text{Speed Up} = \frac{GPU_{original} - GPU_{optimized}}{CPU_{original}} \times 100\%
\]

Note: Look for 4.29(a) at page 16 performance comparison between CPU and GPU.

Note: Look for 4.29(b) at page 17 performance comparisons between GPU original and GPU optimized.

The speed takes in account the best gain obtained while running the code in CPU and then in GPU. It is the time difference between the execution time of CPU and GPU when it is maximum.

Eg. For Blackscholes with 65K size and 256 threads

CPU time = 4.265 ms and GPU time = 1.539

\[
\text{Speed Up} = \frac{CPU_{time} - GPU_{time}}{CPU_{time}} \times 100\% = 64.3\%
\]

The following graphs shows the speed up percentage gained in GPU against the speed up percentage the application is expected to achieve when it is transformed to GPU platform.

Graph 1: Speed up comparison between original speed up, optimized speed up and expected speed up. The speed is in percentage.

Similarly, we calculated the speed up from GPU original to GPU optimized and checked how much gained performance speed up is achieved in optimizing the CUDA code.

Table 2: Speed up from GPU original to GPU optimized

<table>
<thead>
<tr>
<th>Benchmarks</th>
<th>Speed Up Percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blackscholes</td>
<td>41.66%</td>
</tr>
<tr>
<td>Streamcluster</td>
<td>48.63%</td>
</tr>
<tr>
<td>Ferret</td>
<td>23.93%</td>
</tr>
<tr>
<td>Swaptions</td>
<td>29.45%</td>
</tr>
<tr>
<td>x264</td>
<td>39.21%</td>
</tr>
<tr>
<td>Fluidanimate</td>
<td>45.4%</td>
</tr>
<tr>
<td>Raytrace</td>
<td>-</td>
</tr>
</tbody>
</table>

Graph 2: CPU-GPU Communication time between original CUDA and optimized CUDA

6. Discussion:

While reader can draw his/her conclusion from experimental results below, let us provide our interpretation of the measured data. Application running on GPU has benefited from the degree of parallelism available on GPU than CPU. Each application with different size of inputs has shown great performance increase.
Initially for small size of input the GPU execution time was same as CPU. But as we increased the size of the input, the performance became better and better. Also, Exploiting large number of threads was the major factor for speed up. Application like blackscholes and fluidanimate didn’t show much improvement but once the size of input increased they out performed CPU showing 64.3% and 79.4% of speed up which is good considering it was optimized later. And application like swaptions, ferret showed improvement from the beginning but we were unable to exploit the large number of threads for some reason. The best one remains the x264 which achieved improvement much closer as expected, achieving 17.9% against the expected value of 50%. All these application achieved a speed up without optimization.

Optimization helped the original code to rearrange the CUDA and use some techniques to improve and achieve desirable speed to be much closer to expected speed. Table 2 shows the speed up percentage for the optimized code. It was interesting to see none of application except one fluidanimate crossed speed more than 50% and streamcluster which came closer to 50%. This may be due to the fact that we did optimization purely on the GPU see adding extra features from NVIDIA compute capability 3.0, changing and rearranging the original code and making computational changes. But, we never tried to change anything in the algorithm part of application. Many papers implement their experimentation not only for CUDA code optimization but also changing the algorithm to see how it performs. This change is beyond our scope of the project. So, the performance in table 2 is reasonable considering we implemented original algorithm without any change.

Optimization like memory coalescing with input size from 50K - 100K reduces the running time dramatically for fluidanimate and blackscholes. For input size at 100K, the running time under memory coalescing for fluidanimate is 50% lower than that of original kernel execution. But for input size at 3K, 15K and 300K, the effects of memory coalescing is not as large at others. Since for this application, we only modified a portion of the data to obtain memory coalescing, the speedup could be larger if we apply the modification to every portion of the code which is impossible manually. Optimization like shared memory worked well for streamcluster for 4K and 8K size but didn’t affect much for ferret and swaptions. Also, optimization like parallel reduction in swaptions didn’t work very well as expected. Again, x264 performed well for texture memory, shared memory and loop unrolling optimization. There are several other optimization as we do in compiler construction but it is impossible to apply them manually on CUDA and check the performance.

6. Conclusion and Future Work:
In this project, we presented the parallelized CUDA code of Parsec application for blackscholes, streamcluster, ferret, swaptions, x264, fluidanimate and raytrace respectively. This paper characterized the applications in terms of inherent algorithmic characteristics, parallelization, overhead, and optimization, and showed that each application exhibits unique characteristics. We presented efficient optimization like memory coalescing, bank conflict shared memory, cached texture memory, loop unrolling, parallel reduction and demonstrated that the GPU acting as a coprocessor can effectively accelerate massive data computation. Experimental results show that with GPU, the process is 4-6 times faster than the CPU only configuration. And also our experimental result show that optimized kernel functions greatly outperform the naive kernel functions. Our future work extending Parsec benchmark to support more platforms, such as FPGAs.

7. Reference:
3) GPUPU Programming lecture by David Keith, CS 218, 2008.
5) Compile-time Data Layout Transformation for Memory Optimization on GPGPU by Yan Wang, UC Riverside, June 8, 2010.
7) Compiler Framework for Optimization of Affine Loop Nests for GPGPUs, Muthu Manikandan Baskaran Department of Computer Science and Engg, The Ohio State University.
8) Optimizing Parallel Reduction in CUDA by Mark Harris, NVIDIA Developer Technology.
10) NVISION08: Interactive Ray Tracing with CUDA.
18) http://en.wikipedia.org/wiki/Heath%E2%80%93Jarrow%E2%80%93 Morton Framework
4. 2(a) Performance Comparison between CPU and GPU

**Blackscholes 4K**

- CPU Execution Time (ms)
- GPU Execution Time (ms)

**Blackscholes 16K**

- CPU Execution Time (ms)
- GPU Execution Time (ms)

**Blackscholes 65K**

- CPU Execution Time (ms)
- GPU Execution Time (ms)

**Fluidanimate - 35K**

- CPU Execution Time (ms)
- GPU Execution Time (ms)

**Fluidanimate - 100K**

- CPU Execution Time (ms)
- GPU Execution Time (ms)

**Fluidanimate - 300K**

- CPU Execution Time (ms)
- GPU Execution Time (ms)

**Streamcluster - 16, 4K, 8K**

- CPU Execution Time (s)
- GPU Execution Time (s)
4.2(b) Performance Comparison between GPU original vs GPU Optimized

**Blackscholes**

- GPU Original (ms)
- GPU Optimized (ms)

**Streamcluster**

- GPU Original (s)
- GPU Optimized (s)

**Ferret**

- GPU Original (s)
- GPU Optimized (s)

**Swaptions**

- GPU Original (ms)
- GPU Optimized (ms)

**x264**

- GPU Original (ms)
- GPU Optimized (ms)

**Fluidanimate**

- GPU Original (ms)
- GPU Optimized (ms)
Note: This section doesn’t show the full version of the CUDA code. This shows only the portion of optimized code where the optimization has been done.

**Appendix 3.1: Blackscholes**

typedef struct OptionData_device_
  fptype sptprice;
  fptype strike;
  fptype rate;
  fptype volatility;
  fptype otime;

//Kernel Function is called and memory is coalesced//

__global__ void BlkSchlsEqEuroNoDiv( struct float* sptprice,
  float* strike, float* rate, float* volatility,
  float* time, int* otype, float* prices )
{int i = blockDim.x * blockIdx.x + threadIdx.x;
if (i < numOptions)
{
  hstrike = hsptprice + numOptions;
  hrate = hstrike + numOptions;
  hvolatility = hrate + numOptions;
  hotime = hvolatility + numOptions;
  for (i=0; i<numOptions; i++) {
    hotype[i] = data[i].OptionType;
    hsptprice[i] = data[i].s;
    hstrike[i] = data[i].strike;
    hrate[i] = data[i].r;
    hvolatility[i] = data[i].v;
    hotime[i] = data[i].t;
  }
  // Allocate memory//
  cutilSafeCall(cudaMalloc((void**) &dsptprice,
    5 * numOptions * sizeof(fptype)));
  dstrike = dsptprice + numOptions;
  drate = dstrike + numOptions;
  dvolatility = drate + numOptions;
  dotime = dvolatility + numOptions;
  cutilSafeCall(cudaMalloc((void**) &dotype,
    numOptions * sizeof(int)));
  // Transfer from host memory to device memory//
  cutilSafeCall(cudaMemcpy(dsptprice, hsptprice,
    5 * numOptions * sizeof(fptype),
    cudaMemcpyHostToDevice));
  cutilSafeCall(cudaMemcpy(dotype, hotype,
    numOptions * sizeof(int),
    cudaMemcpyHostToDevice));
  hprices = (fptype*)malloc(numOptions * sizeof(fptype));
  // Compute prices on GPU //
  BlkSchlsEqEuroNoDiv<<blocksPerGrid, threadsPerBlock>>(dsptprice, dstrike, drate, dvolatility, dotime, do
type, 0, numOptions, dprices);
  // Transfer back result from device memory to host memory//
  cutilSafeCall(cudaMemcpy(hprices, dprices,
    numOptions * sizeof(fptype),
    cudaMemcpyDeviceToHost));
  // Clean-up memory//
  free(hsptprice);
  free(hotype);
  cutilSafeCall(cudaFree(dsptprice));
  cutilSafeCall(cudaFree(dprices));
  cutilSafeCall(cudaFree(dotype));}
Appendix 3.2: Streamcluster

// streamcluster header file
typedef struct {
    float weight;
    float *coord;
    long assign; /* number of point where this one is assigned */
    float cost; /* cost of that assignment, weight*distance */
} Point;

// this is the array of points
typedef struct {
    long num; /* number of points; may not be N if this is a sample */
    int dim; /* dimensionality */
    Point *p; /* the array itself */
} Points;

// shared memory with and without bank conflict for coordinate of point[x]
extern __shared__ float coord_s[];

if(threadIdx.x == 0)
    for(int i=0; i<dim; i++)
        { coord_s[i] = coord_d[(i*num + threadIdx.x)];

    int min, stride, max, j;
    int id = threadIdx.x;
    int num_banks = 16;
    char cause_bank_conflicts = 0;
    if (cause_bank_conflicts) {
        min = id * num_banks;
        stride = 1;
        max = (id + 1) * num_banks;
    }
    else {
        min = id;
        stride = num_banks;
        max = (stride * (num_banks - 1)) + min + 1;
    }
    for (j = min; j < max; j += stride)
        mem[j] = 0;
    for (i = 0; i < iters; i++)
        for (j = min; j < max; j += stride)
            mem[j]++;
    for (j = min; j < max; j += stride)
        out[j] = mem[i];

// Allocate device and host memory and Copy host memory to device

//kernel execution

/* Determine the number of thread blocks in the x- and y-dimension */
int num_blocks_x = (int)((float)num / (float)THREADS_PER_BLOCK);
int num_blocks_y = (int)((float)num / (float)MAXBLOCKS);
int num_blocks = (int)((float)(num_blocks_x * num_blocks_y) / (float)MAXBLOCKS);

// cost between this point and point[x]: euclidean distance multiplied by weight
x_cost += (coord_d [(i*num + threadIdx.x)] - coord_s[i]) * (coord_d [(i*num + threadIdx.x)] - coord_s[i]);
x_cost = x_cost * p[threadIdx].weight;
float current_cost = p[threadIdx].cost;

// if computed cost is less then original,mark it as to reassign and if larger, save the difference
float *lower = &work_mem_d[threadIdx*(K+1)];
if (x_cost < current_cost) {
    switch_membership_d[threadIdx] = 1;
    lower += x_cost - current_cost;
}
else {
    int assign = p[threadIdx].assign;
    lower[center_table_d[assign]] += current_cost - x_cost;}

// compute the number of centers to close if we open i
low += work_mem_h[j*(K+1) + center_table[i] ];
gl_lower[center_table[i]] = low;

// if opening a center at x saves cost (i.e. cost is negative) do so otherwise, do nothing
if (is_center[i] && gl_lower[center_table[i]] > 0) is_center[i] = false;

is_center[x] = true;
*numcenters = *numcenters + 1 - numclose

Appendix 3.3: Ferret

```c
__shared__ void emd(char *data, float x2, float y2, float *z, int N, int W, int L_POS)

// Allocate memory on host and device
    z  = (float *) malloc(REC_WINDOW * sizeof(float));
    cudaMalloc((void **) &data, sizeof(char) * REC_WINDOW * REC_LENGTH);

// Transfer data from host to device
cudaMemcpy(data, sandbox, (sizeof(char)*REC_WINDOW*REC_LENGTH), cudaMemcpyHostToDevice);

// Execute kernel
    int block_size = 16;
    dim3 dimBlock(block_size);
    dim3 dimGrid((REC_WINDOW/dimBlock.x) + (!(REC_WINDOW%dimBlock.x)?0:1) );

emd<<<dimGrid,dimBlock>>>(data, x2, y2, z_d, REC_WINDOW, REC_LENGTH, LATITUDE_POS);
    cudaMemcpyHostToDevice);

// Update list of nearest neighbors
    for(i=0;i< rec_count;i++) {
        float min_dist = -1;
        int main_idx = 0;
        for( j = 0 ; j < k ; j++ ) {
            if(neighbors[j].dist > min_dist) {
                min_dist = neighbors[j].dist;
                min_idx = j;
            }
        }

        // Compare each record with min value to find the nearest neighbor
        if(z[i] < neighbors[min_idx].dist) {
            sandbox[(i+1)*REC_LENGTH-1] = '\0';
            strcpy(neighbors[min_idx].entry, sandbox +i*REC_LENGTH);
            neighbors[min_idx].dist = z[i];
        }
    }

// Copy data from device memory to host memory
    cudaMemcpy( z, z_d, sizeof(float)*REC_WINDOW, cudaMemcpyDeviceToHost );

// free memory
    free(z);
    cudaMemcpy(data);
    cudaMemcpy(z_d);
```

Appendix 3.4: Swaptions

static __global__ void swaptionsOneBlockPerOption

// Parallel Reduction and Loop Unrolling function call
#include "swaptions_reduction.cuh"

// do reduction in shared memory
for(unsigned int s=1;s<blockDim.x;s*= 2)
    if (tid % (2*s) == 0) {
        sum[tid] = max(sum[tid], sum[tid + s]);
    }

//Cycle through the entire samples array: derive end stock price for each path
for(int iSum = threadIdx.x; iSum < SUM_N; iSum += blockDim.x){
    __TOptionValue sumCall = {0, 0};
    for(int i = iSum; i < pathN; i += SUM_N){
        const int optionIndex = blockIdx.x;
        const real S = d_OptionData[optionIndex].S;
        const real X = d_OptionData[optionIndex].X;
        const real MuByT = d_OptionData[optionIndex].MuByT;
        const real VBySqrtT = d_OptionData[optionIndex].VBySqrtT;

        //Finalizing reduction for swaptionsKernel1
        static __global__ void swaptionsReduce __TOptionValue *d_Buffer, int accumN){
            const int SUM_N = THREAD_N;
            __shared__ real s_SumCall[SUM_N];
            __shared__ real s_Sum2Call[SUM_N];
            __TOptionValue *d_SumBase = &d_Buffer[blockIdx.x * accumN];

            //Allocate internal device memory
            static void initswaptionsGPU(TOptionPlan *plan){
                const int doMultiBlock = (plan->optionCount < 16) ? 64 : 16;
                const int accumN = THREAD_N * doMultiBlock;
            }

            //Deallocate internal device memory
            static void closeswaptionsGPU(TOptionPlan *plan){
                const int doMultiBlock = (plan->optionCount < 16) ? 64 : 16;
                if (doMultiBlock) cutilSafeCall( cudaFree(plan->d_Buffer) );
            }

            // Main computations: Computing the prices
            static void swaptionsGPU(TOptionPlan *plan){
                __TOptionData h_OptionData[MAX_OPTIONS];
                __TOptionValue h_CallValue[MAX_OPTIONS];

                if(plan->optionCount <= 0 || plan->optionCount > MAX_OPTIONS){
                    printf("swaptionsGPU(): bad option count.\n");
                    return;
                }

                for(int i = 0; i < plan->optionCount; i++){
                    const double T = plan->optionData[i].T;
                    h_OptionData[i].S(real)plan->optionData[i].S;

                    cutilSafeCall( cudaMemcpyToSymbol( d_OptionData, 
                        h_CallValue, sizeof(__TOptionData) 
                    )
                }
            }
        }
    }
}
}
Appendix 3.5: x264

// declare texture reference for 2D float texture:
texture<
uint8_t,
2,
cudaReadModeElementType>
currentTex;
texture<
uint8_t,
2,
cudaReadModeElementType>
lastTex

// defining all parameters for estimating cost:
void runCudaA(uint8_t *current, uint8_t *ref, int strideCur, int strideRef, int width, int height,
int2 ** mvOut, int A, int2 *mvsX);
void cuda_me(x264_t *h, int **mvX, int **mvY){
  int *mvsXD, *mvsYD;
  cudaMalloc((void**)&mvsXD, bh*bw* sizeof(int ));
  cudaMalloc((void**)&mvsYD, bh*bw* sizeof(int ));

  // Allocating Memory //
cutilSafeCall(cudaMallocArray(&currentArray, &channelDesc, h->fenc->i_width[0], h->fenc->i_lines[0]));

  // copy CPU to GPU://
cutilSafeCall(cudaMemcpy2DToArray(currentArray, 0, 0, h->fenc->plane[0], h->fenc->i_stride[0]*
sizeof(uint8_t), h->fenc->i_width[0]*sizeof(uint8_t), h->fenc->i_lines[0],cudaMemcpyHostToDevice));

  // estimating cost what will MVp will actually be://
  __device__ unsigned mvCost(int dx, int dy){
    dx = (abs(dx))<<2;
    dy = (abs(dy))<<2;
    int xCost = round((log2f(dx+1)*2 + 0.718f + !!dx) + .5f);
    int yCost = round((log2f(dy+1)*2 + 0.718f + !!dy) + .5f);
    return (LAMBDA * (xCost+yCost));
  }

  // per block for one macroblock is assigned to minimize cost of motion vector for each macroblock://
  __global__ void minSAD2(unsigned bw, int* mvXs,int* mvYs, unsigned* minSAD, unsigned* bestLoc){
    const unsigned int bx = blockIdx.x;
    const unsigned int by = blockIdx.y;
    #define FILTER(a,b,c,d) ((((a+b+1)>>1)+((c+d+1)>>1)+1)>>1)
    __device__ uint8_t val2(ImageTexture t, unsigned  x, unsigned  y){
      return FILTER(tex2D(t, x, y),tex2D(t, x+1, y),tex2D(t, x, y+1),tex2D(t, x+1, y+1));
    }
    __device__ uint8_t val4(unsigned  x, unsigned  y, ImageTexture t){
      return (val2(t, x, y)+val2(t, x+2, y)+val2(t, x, y+2)+val2(t, x+2, y+2))>>2;
    }
    __device__ uint8_t val16(unsigned  x, unsigned  y, ImageTexture t){
      return (Val4(x, y, t)+val4(x+4, y, t)+val4(x, y+4, t)+val4(x+4, y+4, t))>>2;
  }

23
__device__ unsigned SAD2(ImageTexture a, ImageTexture b, unsigned xStart, unsigned yStart, unsigned width, unsigned height, unsigned dx, unsigned dy) {
    unsigned ret = 0;
    for(unsigned x = xStart; x < xStart + 2*width; x+=2) {
        for(unsigned y = yStart; y < yStart + 2*height; y+=2) {
            ret = __usad(val2(a, x, y), val2(b, x+dx, y+dy), ret);
        }
    }
    return ret;
}

//Copy back from GPU to CPU//
//Memory Free//

Appendix 3.7: Raytrace

__device__ int traverseBvh(float3 &eye, float3 &dir, unsigned *cells, uint4* triangles, unsigned* stackNode, unsigned* stackCount, bool idle) {
    for(int i=0; i<count; i++) {
        float u, v, t;
        int a = resolve[i*4+1];
        int b = resolve[i*4+2];
        int c = resolve[i*4+3];
        intersectDirect(a, b, c, eye, dir, u, v, t);
    }
    unsigned short x = startX + blockIdx.x*8 + threadIdx.x;
    unsigned short y = startY + blockIdx.y*8 + threadIdx.y;
    dir.x = upperLeft.x + __fmul_rn(dx.x, x) + __fmul_rn(dy.x, y);
    dir.y = upperLeft.y + __fmul_rn(dx.y, x) + __fmul_rn(dy.y, y);
    dir.z = upperLeft.z + __fmul_rn(dx.z, x) + __fmul_rn(dy.z, y);
    int hitTriangle = traverseBvh<true>(origin, dir, cells, triangles, stackNode, stackCount, true);
    unsigned short x = startX + blockIdx.x*8 + threadIdx.x;
    unsigned short y = startY + blockIdx.y*8 + threadIdx.y;
    color.x = min(color.x, 1.0f);
    color.y = min(color.y, 1.0f);
    color.z = min(color.z, 1.0f);
    float3 dir;
    dir.x = upperLeft.x + __fmul_rn(dx.x, x) + __fmul_rn(dy.x, y);
    dir.y = upperLeft.y + __fmul_rn(dx.y, x) + __fmul_rn(dy.y, y);
    dir.z = upperLeft.z + __fmul_rn(dx.z, x) + __fmul_rn(dy.z, y);
    unsigned material;
    float3 reflectedDir;
    float3 normal;
    if(hitTriangle >=0) {
        float t;
        catchNormalIntersection(triangles, hitTriangle, eye, dir, normal, t, material);
        intersectionPoint = eye + dir*t;
        reflectedDir = dir - normal*dot(normal, dir)*2.0f;
    }
}