# MARS: ACCELERATING MAPREDUCE WITH GRAPHICS PROCESSORS

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The Hong Kong University of Science and Technology
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This is to certify that I have examined the above M.Phil. thesis and have found that it is complete and satisfactory in all respects, and that any and all revisions required by the thesis examination committee have been made.

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### ABSTRACT

We design and implement Mars, a MapReduce runtime system accelerated with graphics processing units (GPUs). MapReduce is a simple and flexible parallel programming paradigm originally proposed by Google, for the ease of large scale data processing on thousands of CPUs. Compared with CPUs, GPUs have an order of magnitude higher computation power and memory bandwidth. However, GPUs are designed as special-purpose co-processors and their programming interfaces are less familiar than those on the CPUs to MapReduce programmers.

To harness GPUs' power for MapReduce, we developed Mars to run on NVIDIA GPUs, AMD GPUs, as well as multi-core CPUs. Furthermore, we integrated Mars into Hadoop, an open-source CPU-based distributed MapReduce system. Mars hides the programming complexity of GPUs behind the simple and familiar MapReduce interface, and automatically manages task partitioning, data distribution, and parallelization on the processors. We have implemented six representative applications on Mars and evaluated their performance on PCs equipped with GPUs as well as multi-core CPUs. The GPU acceleration with an NVIDIA GTX280 achieved a speedup of an order of magnitude over a quad-core CPU. Utilizing both the GPU and the CPU further improved GPU-only performance by 40% for some applications. Additionally, integrating Mars into Hadoop enabled GPU acceleration for a network of PCs.

### CHAPTER 1

### INTRODUCTION

MapReduce is a successful paradigm [15], originally proposed by Google, for the ease of distributed data processing on a large number of machines. In such a system, users specify two functions: (1) a *map* function to process an input key/value pair, and to generate a set of intermediate key/value pairs; (2) a *reduce* function to merge all intermediate key/value pairs associated with the same key. The system will automatically distribute and execute tasks on multiple machines [4, 15] or multiple CPUs in a single machine [28]. Thus, this paradigm reduces the programming complexity so that developers can easily exploit the parallelism in the underlying computing resources for complex tasks. Encouraged by the success of CPU-based MapReduce systems, in particular, Phoenix [28], we develop Mars, a MapReduce system accelerated with graphics processors, or GPUs.

GPUs can be regarded as massively parallel processors with an order of magnitude higher computation power (in terms of number of floating point operations per second) and memory bandwidth than CPUs [7]. Moreover, the computational performance of GPUs is improving at a rate higher than that of CPUs. However, it is a challenging task to program GPUs for general-purpose computing applications, including those that MapReduce users are familiar with. Specifically, GPUs are traditionally designed as special-purpose co-processors for dedicated graphics rendering. As such, GPU cores are SIMD (Single-Instruction-Multiple-Data), which discourages complex control flows. Furthermore, GPU cores are virtualized, and threads are managed by the hardware. Finally, GPUs manage their own on-board device memory and require programmers to explicitly transfer data between the GPU memory and the main memory. Additionally, the architectural details of GPUs vary by vendors as well as by product releases, and programmer's access to these details is limited. All these factors make desirable a GPGPU (General Purpose Computation on GPUs) framework on which users can develop correct and efficient GPU programs easily.

Recently, several GPGPU programming frameworks have been introduced, such as NVIDIA CUDA [5], and AMD Brook+ [1]. These frameworks significantly improve the programmability of GPUs; nevertheless, their interfaces are vendor-specific and their hardware abstractions may

be unsuitable for complex applications such as those running on MapReduce. Therefore, we propose Mars, a MapReduce framework to ease the programming of such applications on the GPU. Furthermore, the MapReduce framework of Mars enables the integration of GPU-accelerated code to distributed environment, like Hadoop, with the least effort. Our Mars system can run on multicore CPUs (MarsCPU), on CUDA-enabled NVIDIA GPUs (MarsCUDA) or Brook+-enabled AMD GPUs (MarsBrook), or on a combination of a multi-core CPU and a GPU on a single machine. We further integrate Mars into Hadoop [4], an open-source CPU-based MapReduce system on a network of machines, which results in MarsHadoop, where each machine can utilize its GPU with MarsCUDA or MarsBrook in addition to its CPU with the original Hadoop. No matter what GPU and/or CPU Mars runs on, the API (Application Programming Interface) to the user is the same and is similar to that of existing CPU-based MapReduce systems.

Easing up GPU programming for MapReduce applications is the main goal of our work. However, a higher-level abstraction for programming, specifically MapReduce, comes at a price of performance. In particular, we identify the following three technical challenges in implementing Mars on GPUs. First, since MapReduce divides up a task by data, load imbalance is an inherent problem in utilizing the massive thread parallelism on the GPU, especially because GPU threads are managed by the hardware. Second, GPUs lack efficient global synchronization mechanisms. Threads in Map or Reduce tasks are likely to have write conflicts on the output buffer. While atomic operations are enabled in recent GPUs, the overhead of atomic operations would harm the scalability of massive GPU threads [2]. We consider a lock free scheme to minimize the synchronization overhead among GPU threads. Third, MapReduce applications are in general data-intensive and their result sizes are data-dependent. These two characteristics pose the following requirements on programming the GPU: (a) sufficient thread parallelism to hide the high latency and to utilize the high bandwidth of the device memory; (b) pre-allocation of output buffers in the device memory for bulk DMA transfers, as GPU memory allocation is done through the CPU before the GPU program starts.

With these challenges in mind, we develop Mars for GPUs of two most common programming interfaces - CUDA and Brook+. We focus on MarsCUDA, than MarsBrook, because in our implementation and evaluation, CUDA was more flexible and had a higher performance than Brook+ for MapReduce applications.

In MarsCUDA, the massive thread parallelism on the GPU is well utilized as each thread is

automatically assigned a key/value pair to work on. In *Map*, the system evenly distributes key/value pairs to each thread. In *Reduce*, we develop a simple but effective skew handling scheme to redistribute data evenly across all reduce tasks. To avoid write conflicts between threads, we adopt a lock-free scheme that guarantees the correctness of parallel execution with little synchronization overhead.

We extend our general design of Mars from a GPU-only MapReduce framework to a MapReduce system with GPU acceleration enabled. With this extension, Mars components can work stand-alone on a single platform, e.g., MarsCUDA on a CUDA-enabled GPU or MarsCPU on a multi-core CPU, as well as to work together to utilize multiple processors, e.g., a CPU and a GPU on a single machine. We also integrate Mars into Hadoop to enable GPU-acceleration for individual machines in a distributed environment.

We evaluated the performance of Mars in comparison with its CPU-based counterparts and the native implementation without MapReduce. Our results demonstrate the effectiveness of our GPU-oriented optimization strategies. On average, our MarsCUDA is 22 times faster than the CPU-based MapReduce, Phoenix [28], and is less than 3 times slower than the hand-tuned native CUDA implementation. Additionally, the applications developed with Mars had a code size reduction up to seven times, compared with hand-tuned native CUDA code.

**Organization:** The remainder of the thesis is organized as follows. We give a brief overview of GPUs, and review prior work on GPGPU and MapReduce in Chapter 2. We present the design and implementation details of Mars in Chapter 3 and Chapter 4 respectively. We present the extension to multiple machines in Chapter 5. In Chapter 6, we present our experimental results. Finally, we conclude in Chapter 7.

### CHAPTER 2

### PRELIMINARIES AND RELATED WORK

In this chapter, we first give a brief introduction on the GPU, and then review the related work on GPGPU as well as on MapReduce.

## 2.1 Graphics Processing Units (GPUs)

The GPU is an integral component of modern computers, ranging from handheld devices to highend servers. GPUs are originally designed for gaming applications with fixed hardware pipelines for rendering. Due to the high computation power and rapidly improving programmability, they have recently become a powerful co-processor for general purpose computing [7].

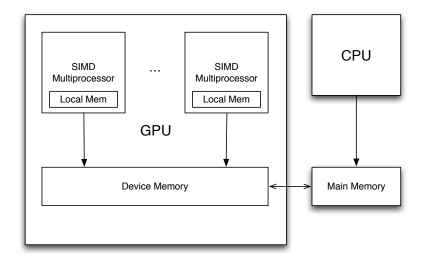


Figure 2.1. The many-core architecture model for GPUs

As shown in Figure 2.1, we model the GPU as a many-core processor, which contains a number of SIMD multiprocessors. Such a many-core model is common to both AMD and NVIDIA GPUs. On the GPU board, there is a GRAM device memory. The device memory has both a high bandwidth and a high access latency. For example, the NVIDIA GTX280 GPU has an access latency of 400 to 600 cycles, and the peak memory bandwidth between the device memory and the multiprocessors is around 140 GB/second.

Both NVIDIA CUDA and AMD Brook+ expose a parallel programming model, which does not require programmers to have knowledge of the graphics rendering pipeline. In this model, the system consists of a *host* (a CPU), and one or more *devices* (GPUs). GPUs are abstracted as massively data-parallel co-processors. CUDA and Brook+ programmers write code using C/C++ syntax with extended keywords for kernel functions, which are GPU programs to be executed on *devices*.

Programming frameworks such as CUDA and Brook+ greatly improve the programmability of the GPU. However, it is still a challenging task of developing efficient GPU programs for complex applications, such as those with MapReduce, because GPUs have a special-purpose co-processor architecture and are vendor-specific on the programming frameworks for complex applications. Although the newly introduced OpenCL [6] is an industry standard further hiding hardware details from users, Mars is at a higher level of abstraction. OpenCL is a general-purpose programming language, with which Mars or other MapReduce frameworks can be developed.

### 2.2 GPGPU

GPGPU, or General Purpose Computation on GPUs, has recently emerged in various applications, such as linear algebra [22, 31], embedded system design [16], bioinformatics [12], databases [17, 18, 21], machine learning [13], and distributed computing projects including Folding@home and Seti@home. Recently, several GPGPU languages including AMD Brook+ [1] (extended from Brook [10]) and NVIDIA CUDA [5] have been proposed by GPU vendors. They usually expose a general-purpose, massively multi-threaded computing architecture and provide a programming environment similar to C/C++. High-level programming frameworks such as Accelerator [30] and RapidMind [25] are also developed to better facilitate GPGPU programming. These programming frameworks require programmers to have knowledge of specific programming models such as the stream programming model in Brook+ [10], or even more, knowledge of the GPU hardware details. By contrast, we propose to develop a MapReduce framework accelerated with GPUs to ease the development of a more complex class of data processing tasks. It provides a uniform MapReduce interface no matter whether it runs on the GPU, on the CPU, or both.

We now briefly survey recent work that developed GPGPU primitives as building blocks for various applications, in particular, those not covered in the survey by Owens et al [27]. Sengupta et

al. [29] proposed the segmented scan primitive. He et al. [20] proposed a multi-pass scheme to optimize the scatter and the gather operations. He et al. [21] further developed a small set of primitives such as prefix sum and split for relational databases. Additionally, CUDPP [3], a CUDA library of data parallel primitives, was released for GPGPU computing. These GPU-based primitives reduce the complexity of GPU programming. However, even with the primitives, programmers need to write complex GPU code for data processing tasks. By contrast, our work further simplifies GPU programming for MapReduce programmers by providing them with a higher level and more familiar interface than the primitives.

This thesis focuses on accelerating MapReduce on the GPU, and provides a GPU-based MapReduce framework to developers. As in the original MapReduce, it is up to developers' choice whether to use MapReduce or not according to the workload's computational characteristics. Recent studies [23] have used data analysis techniques to categorize the computational characteristics of different workloads on the GPU. These techniques are helpful for developers to determine whether their workloads are suitable for Mars in specific and the GPU in general.

# 2.3 MapReduce

The MapReduce framework [15] is based on two primitives, Map and Reduce, from functional programming. The general form is as follows:

Map:  $(k_1, v_1) \to list(k_2, v_2)$ .

Reduce:  $(k_2, list(v_2)) \rightarrow list(k_3, v_3)$ .

The Map function takes an input key/value pair  $(k_1, v_1)$  and outputs a list of intermediate key/value pairs  $(k_2, v_2)$ . The Reduce function takes all values associated with the same key and produces a list of key/value pairs. Programmers implement the application logic inside the Map function and the Reduce function. The MapReduce runtime manages the parallel execution of these two functions.

The following pseudo code illustrates a program written using MapReduce. This program counts the number of occurrences of each word in a collection of documents [15]. In this program, *Map* and *Reduce* are implemented using two system-provided APIs, *EmitIntermediate* and *Emit*, respectively.

```
Map(void *doc) {
1: for each word w in doc
2: EmitIntermediate(w, 1); // count each word once
}
Reduce(void *word, Iterator values) {
1: int result = 0;
2: for each v in values
3: result += v;
4: Emit(word, result); // output word and its count
}
```

There have been several MapReduce implementations since MapReduce was proposed [15]. Hadoop [4] is an open-source MapReduce implementation on clusters. Based on Hadoop, Yang et al. [32] added the merge operation to MapReduce for the ease of relational databases operations. Phoenix [28] is an efficient MapReduce runtime system on multi-core CPUs. Kruijf et al. [14] developed MapReduce on the Cell BE. Yeung et al. [33] implemented an FPGA-based MapReduce system.

Let us briefly introduce the implementation of Phoenix [28]. A key component in Phoenix is a scheduler, for buffer management and task distribution. The scheduler starts the Map stage by evenly dividing the input buffer into small chunks, and assigns the chunks to map workers dynamically. Each map worker runs in a CPU thread. The Reduce stage does not start until all Map tasks are done. The scheduler groups the intermediate output from the Map stage by key, and a Reduce worker processes values associated with the same key. Reduce tasks are assigned to workers dynamically. Each reduce worker maintains a static array for outputting results, and sorts this static array using insertion sort. Finally, the scheduler merges all output arrays of reduce workers into a single one. Because the output data size is not known in advance, the scheduler first allocates buffers with a default small size, and then resizes the buffer as needed.

Catanzaro et al. [11], developed another MapReduce system on the GPU, but it required programmers to be aware of GPU hardware details, such as thread configuration and memory hierarchy. Finally, the Merge framework [24], focused on dynamically scheduling MapReduce tasks among multiple processors, dedicated to Intel products. By contrast, Mars hides hardware details from programmers, and works on heterogeneous GPUs, a combination of CPU and GPU on a single

machine, as well as a distributed system of multiple machines.

### CHAPTER 3

### DESIGN OF MARS

In this chapter, we present our design for Mars, with emphasis on the GPU-based component. Our design is guided by the following three goals.

- 1. *Programmability*. User code size reduction encourages programmers to use the GPU for their tasks.
- 2. Flexibility. The design should be applicable to various multi-/many-core processors, e.g., multi-core CPUs and AMD GPUs, and should be as expressive as the underlying runtime, e.g., NVIDIA CUDA, AMD Brook+ or pthreads, so that the system will work for a wide range of hardware and applications.
- 3. High Performance. The overall performance should be accelerated by the GPU effectively.

### 3.1 Overview

By examining the Phoenix design, we see that there are three potential sources of overhead. First, the tight-coupling of the Map and Reduce stages makes every application go through both stages, no matter whether they need both stages or not. Second, a dynamic thread scheduler for task assignment heavily relies on locking to implement synchronization. Third, each reduce worker may require frequent data movement for sorting the static output array, and the data movement can become a bottleneck for the overall performance. The latest paper about Phoenix also points out this problem [34].

In the Mars design, we decide to separate a MapReduce workflow into three loosely coupled stages - *Map*, *Group* and *Reduce*. The *Group* stage is designed to group *Map* output by key, which is the format for *Reduce* input. Our observation is that some applications need only the *Map* stage, some need both *Map* and *Group*, and some need all of the three stages. The Group stage is the same as running Reduce with the identity function in the original MapReduce system [15]. Our purpose

of providing an explicit Group stage is to allow a MapReduce application with high flexibility to customize its workflow, and to avoid the overhead of entering unnecessary stages. No matter what configuration of the three stages is for an application, the MapReduce interface of Mars is unchanged - users write Map and/or Reduce functions when necessary.

Moreover, we decide to use a lock-free scheme for synchronization and to perform in-advance buffer allocation. One reason is to avoid heavy overheads of locking and buffer reallocation. The other reason is that, current GPUs do not support locking or in-flight buffer reallocation. In our design, we statically distributes tasks to a massive number of GPU threads, so that we can fully utilize the parallelism of the GPU. We adopt a two-phase, lock-free scheme for result output. The basic idea is that, in the first phase, we calculate histograms on the size of output results for each thread, followed by a prefix sum operation on the histograms, so that we obtain both the exact output buffer size and the deterministic write position for each thread; in the second phase, we perform the actual computation and output. We will detail this strategy in Section 3.4.

#### 3.2 Data Structure

Data structures in Mars affect the workflow, memory access patterns, and the expressiveness of the system.

Since the GPU does not support dynamic memory allocation on the device memory during the execution of the GPU code, this limitation rules out dynamic data structures such as queues and linked lists, as used in other MapReduce implementations. Instead, we use plain arrays as the main data structure in Mars. The *Map* stage takes input records in the key/value form, and outputs intermediate result records, which are in turn the input of the *Group* stage. The output of the *Group* stage is the input of the *Reduce* stage, and *Reduce* produces final output records. Each of the three sets - the input records, the intermediate records and the output records - is stored in three arrays, i.e., the key array, the value array and the directory index array. The directory index consists of an entry of <key offset, key size, value offset, value size > for each key/value pair. Given a directory index entry, we fetch the key or the value at the corresponding offset in the key array or the value array.

Variable-sized types, such as strings, are supported with the directory index, since current GPUs have no such build-in types yet. If two key/value pairs need to be swapped, we swap their corre-

sponding entries in the directory index without modifying the key and the value arrays.

Some applications perform chained MapReduce procedures, where the output of one MapReduce procedure is the input of another one. Since the sets of input records, intermediate records and output records are all in the three-array structure uniformly, chained MapReduce is supported gracefully in Mars.

#### 3.3 Mars Workflow

Figure 3.1 illustrates the workflow of Mars, assuming the data resides in the disk at the beginning. The Mars scheduler runs on the CPU, and schedules tasks to the GPU. Mars has three stages, *Map*, *Group*, and *Reduce*.

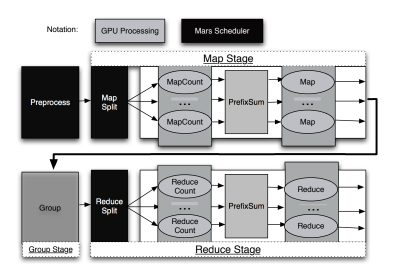


Figure 3.1. The workflow of Mars on the GPU.

Before the *Map* stage, Mars preprocesses on the CPU the input data from disk, transforming the input data to key/value pairs (input records) in main memory. After that, it transfers input records from the main memory to the GPU device memory.

In the *Map* stage, *Map Split* dispatches input records to GPU threads such that the workload for all threads is even. Each thread executes the user-defined MapCount function to compute a local histogram on the number and the total size of intermediate records that *Map* will output. Then, the runtime performs a GPU-based *Prefix Sum* on the local histograms to obtain the output size and the write position for each thread. Finally, after the CPU allocates the output buffer in the device

memory, each GPU thread executes the user-defined Map function and outputs results. Since the write position for each thread is pre-computed and has no conflict with any other threads, there will be no write conflict between concurrent threads. This lock-free scheme of MapCount, Prefix Sum, and Map is adapted from our previous work [21].

In the *Group* stage, both sort-based and hash-based approaches are available for grouping records by key. However, we adopt the sort-based, because some applications require to sort all output records, and the hash-based approach has to perform additional sort within each hash bucket.

In the *Reduce* stage, *Reduce Split* dispatches each group of records with the same key to a GPU thread. However, it may cause load imbalance between threads, since the number of records of different groups may vary widely. We adopt a skew handling scheme to alleviate the load imbalance problem (Section 3.6). The *Reduce* stage then works in a lock-free scheme, similar to that in *Map*, to obtain the result size and the write location for each thread. Finally, all Reduce workers output results to a single buffer.

Because these three stages are loosely coupled and not every application requires all stages, Mars allows users to customize the following workflows in their applications:

- MAP\_ONLY. Mars executes the *Map* stage only, and does not execute the *Group* or *Reduce* stage.
- MAP\_GROUP. Mars executes the *Map* and *Group* stages, and does not execute the *Reduce* stage.
- MAP\_GROUP\_REDUCE. Mars executes all three stages *Map*, *Group*, and *Reduce* stages.

Because usually applications need a Map to transform input records, and a Group to prepare for the intermediate records to feed to Reduce, we exclude the other workflow configurations that skip either Map or Group in the presence of Reduce.

### 3.4 Lock-free Scheme

With the array structure, we allocate the space on the device memory for the input data as well as for the result output before executing the GPU program. However, the sizes of the output from the

*Map* and the *Reduce* stages are unknown. Moreover, write conflicts occur when multiple threads write results to the shared output array. To address these two problems, we adopt a previous lock-free output scheme for relational joins [21]. Since the output scheme for the *Map* stage is similar to that for the *Reduce* stage, we present the scheme for the *Map* stage only.

First, each MapCount invocation on a thread outputs three counts, i.e., the number of intermediate results, the total size of intermediate keys (in bytes) and the total size of intermediate values (in bytes). Based on intermediate key sizes (or value sizes), Mars computes a prefix sum on these sizes and produces an array of write locations. A write location is the start location in the output array for a map task to write. Based on the number of intermediate results, Mars computes a prefix sum and produces an array of start locations in the output directory index. Through these prefix sums, we also know the sizes of the arrays for the intermediate results. Finally, Mars allocates arrays in the device memory with the exact sizes for storing the intermediate results.

Second, each Map invocation on a thread outputs the intermediate key/value pairs to the output array. Since each Map has its deterministic and non-overlapping positions to write to, the write conflicts are avoided.

The lock-free scheme is suitable for the massive thread parallelism on the GPU, even though it performs a MapCount in addition to a Map. The overhead of executing MapCount is application dependent, and is usually small. For example, this overhead is negligible in the matrix multiplication in our study, since MapCount simply emits the size without performing the actual multiplication. In addition, the code for MapCount function is also application dependent, while in most cases, programmers write one statement to emits output data sizes.

### 3.5 Rapid Group

The Group stage requires to sort intermediate records. However, we observe that some applications inherently have their intermediate records grouped after the Map phase, and each group has the same number of records. For example, [A,A,A,B,B,B,C,C,C] shows three groups with A, B, and C as the key respectively, and each group is with the same size 3. For such applications, Mars provides a configuration parameter for users to indicate whether the intermediate data is already grouped. The runtime automatically skips the time consuming sorting, and then dispatches each group of intermediate records with the same group size to Reduce workers. We name this strategy

as "Rapid Group".

# 3.6 Skew handling

We design a skew handling scheme to distribute workloads evenly across reduce workers, where the user-defined Reduce operation is commutative and associative. This scheme iteratively performs the Reduce stage in the following two steps. First, we divide the data into M equal-sized chunks. Second, we perform a reduction on each chunk. In this step, each of the M threads applies the reduce function on groups of records in a single chunk. Note, in each iteration, we perform reduction on the intermediate results with the same keys only.

### 3.7 Mars APIs

Mars provides a small set of APIs. Similar to the existing MapReduce frameworks, Mars has two kinds of APIs, the user-implemented APIs, which the users should implement by themselves, and the system-provided APIs, which the users can use as library calls. The definitions of these APIs are in Table 3.1.

Table 3.1. Mars APIs

<b>Function Name</b>	Description	Function Type
MAP_COUNT	It calculates the output buffer size of MAP.	User-implemented
MAP	The map function.	User-implemented
REDUCE_COUNT	It calculates the output buffer size of REDUCE.	User-implemented
REDUCE	The reduce function.	User-implemented
EMIT_INTERMEDIATE_COUNT	It emits the key size and the value size in	System-provided
	MAP_COUNT.	
EMIT_INTERMEDIATE	It emits the key and the value in MAP.	System-provided
EMIT_COUNT	It emits the key size and the value size in RE-	System-provided
	DUCE_COUNT.	
EMIT	It emits the key and the value in REDUCE.	System-provided

### CHAPTER 4

### SINGLE-MACHINE IMPLEMENTATIONS

In this chapter, we present the implementation details of Mars on a single machine. The current Mars system consists of four modules (Table 4.1). All these four modules share the common design of Mars, and provide the same MapReduce interface to the user. They can run on different hardware platforms: MarsCUDA on an NVIDIA GPU, MarsBrook on an AMD GPU, MarsCPU on a multi-core CPU, and the GPU/CPU co-processing module on both the CPU and the GPU through combining the aforementioned modules. Different modules in Mars allow programmers to take advantage of different processors on a single machine. Because our machines cannot host a multi-GPU configuration due to limited extension slots, we have not explored multi-GPU co-processing.

Table 4.1. Modules in the Mars system

Implementation	Software platform	Hardware platform
MarsCUDA	NVIDIA CUDA	an NVIDIA GPU
MarsBrook	AMD Brook+	an AMD GPU
MarsCPU	pthreads	a multi-core CPU
GPU/CPU co-processing	CUDA/Brook+ and pthreads	NVIDIA/AMD GPUs and multi-core CPUs

### 4.1 MarsCUDA

We implemented MarsCUDA using NVIDIA CUDA. We used the GPU Prefix Sum routine from CUDPP [3] to implement the lock-free scheme, and the GPU Bitonic Sort routine for the *Group* phase. CUDA exposes sufficient hardware details of NVIDIA GPUs, so that we can apply some optimizations in MarsCUDA runtime.

#### 1. Memory access

Coalesced access. We utilize the NVIDIA GPU feature of coalesced access to improve the memory performance. In CUDA, simultaneous device memory accesses by threads in a half-warp (warp is an NVIDIA term for a group of 32 threads for scheduling) can be coalesced into a single memory transaction, which significantly reduces the number of device memory accesses. We implement the access to the directory index arrays as coalesced.

**Local memory.** NVIDIA GPUs provide the programmable on-chip local memory (or *shared memory* [26]), for sharing data among threads running on the same multiprocessor. It is important to fully utilize the local memory to reduce the costly accesses to the GPU memory. In Mars, data sharing or communication only happens in the Group stage. MarsCUDA runtime automatically uses a GPU-based bitonic sort [19] to exploit this memory hierarchy in the Group stage. Mars does not expose the local memory to the user-defined functions in the Map and the Reduce stages. Since local memory is programmer-controlled fast memory, it introduces complexity and needs the effort from the programmer. This is a trade-off between performance and programmability. Nevertheless, users who are aware of the GPU memory hierarchy and need such data sharing can exploit the local memory in implementing the Map (or Reduce) function.

**Built-in vector types.** Data accesses in the GPU device memory should be aligned to make sure the correctness and achieve high memory bandwidth. Fortunately, GPUs support built-in vector types [26], including *float4* and *int4*. The alignment requirement is automatically fulfilled for built-in types. In addition, the GPU is able to issue a single load instruction to read data of built-in type, of size up to 16 bytes. Compared with reading an array one *float* or *int* at a time, the number of compiler-generated instructions for reading *float4* or *int4* is greatly reduced and the overall performance is improved.

Page-locked host memory. CUDA supports page-locked host memory (a.k.a pinned), which prevents the operating system from paging the locked memory buffer, yielding high transfer bandwidth between the device memory and the host memory [26]. The MarsCUDA runtime utilizes the page-locked host memory mechanism, in order to reduce the data transfer overhead. Our test demonstrated that page-locked memory can double the memory transfer rate through PCI-E bus than pageable memory.

#### 2. Parallelism

Since CUDA exposes the thread configuration, we utilize the parallelism by assigning the tasks to

a large number of threads. The thread configuration, i.e., the number of thread blocks and the number of threads per thread block, is related to both hardware and software factors: (1) the hardware configuration such as the number of multiprocessors and the on-chip computation resources such as the number of registers on each multiprocessor, and (2) the characteristics of the map and the reduce tasks, e.g., the degree of memory- or computation-intensiveness.

Since the map and the reduce functions are implemented by the developer, and their costs are unknown to the runtime system, it is difficult to find the optimal setting for the thread configuration at run time. CUDA provides an off-line calculator <sup>1</sup> for computing the multiprocessor occupancy given a CUDA program. For the program (either the map task or the reduce task), the calculator takes the number of threads per thread block and the number of registers used per thread as input, and outputs the occupancy and the number of active thread blocks per multiprocessor. The number of registers used per thread is obtained using the NVCC compiler of CUDA.

With the calculator, we iterate the number of threads per block in multiples of 32 (the schedule unit size) ranging from 32 to 512 (the maximum number of threads per thread block), until the occupancy is higher than a predefined threshold. Thus, we get the number of threads per thread block and the number of thread blocks. In practice, we set the occupancy threshold to be 2/3 so that the GPU is sufficiently busy, and each thread block receives adequate computation resources.

### 4.2 MarsBrook

We implement MarsBrook on AMD GPUs using the stream programming model Brook+ [1]. Due to the limitation of Brook+, MarsBrook is less advanced than MarsCUDA in both expressivity and performance. Nevertheless, as programming support of Brook+ improves, MarsBrook can demonstrate a higher flexibility and performance.

MarsBrook requires users to specify the data types of keys and values statically, and each record is of a fixed size. Type conversion is not allowed in Brook+. Unlike CUDA, Brook+ does not allow the developer to access data in GPU memory by arbitrary address. Instead, data in the GPU memory is accessed using a *stream*, which is essentially a sequentially-accessed array of fixed-sized elements. Random access in a stream is achieved by providing another predefined stream, consisting of indexes of target elements to access. Although the Mars APIs are the same on CUDA

<sup>&</sup>lt;sup>1</sup>http://developer.download.nvidia.com/compute/cuda/CUDA\_Occupancy\_calculator.xls

and on Brook+, as listed in Table 3.1, using Mars on CUDA is more flexible than using that on Brook+ when the Mars user develops a user-defined function.

Moreover, MarsBrook has relatively limited room for performance optimization. The reason is that Brook+ does not expose detailed hardware features, e.g., fast on-chip local memory, coalesced memory access, or GPU thread configuration.

### 4.3 MarsCPU

We implement MarsCPU using the pthreads library on linux for multi-threading. Instead of adopting lock-based task scheduling as in Phoenix, MarsCPU inherits the lock-free design of GPU-based Mars, which we expect to scale to hundreds of cores for future many-core CPUs. MarsCPU deploys CPU threads to perform Map and Reduce tasks. If there are N Map (or Reduce) tasks, and T CPU threads, where N is usually much larger than T, then a thread processes  $\lceil N/T \rceil$  tasks. We implement a CPU multi-threaded parallel mergesort for the Group stage.

## 4.4 GPU/CPU co-processing

The workflow of GPU/CPU co-processing is shown in Figure 4.1. There are also mainly three stages, Map, Group and Reduce. In the Map stage, the scheduler divides the input data into multiple chunks. The number of chunks is equal to the total number of CPUs and GPUs in the machine. The chunk sizes are determined based on the performance comparison between the CPU and the GPU. Suppose the speedup of the GPU worker over the CPU worker is S, where the S where the S defined to be the ratio of the execution time on the CPU to that on the GPU for the same amount of input data. Given the total input size of S bytes, we assign data chunks of S and S and S bytes to the GPU and the CPU workers, respectively. The speedup S can be obtained by either calibration or predictive model [23].

When a processor finishes a *Map* task, it performs a local *Group* on intermediate results. The runtime merges all intermediate results. When all the processors finish their tasks, the *Map* stage ends.

The *Reduce* stage takes the intermediate results from the *Group* stage as input. Similar to the *Map* stage, the co-processing scheduler statically assigns the data chunks to the processors. When

all the processors finish their tasks, the runtime merges all local results.

Mars dispatches workload between the GPU worker and the CPU worker only if the following conditions are satisfied. First, the Map and Reduce stages take up high proportion of the entire running time on the CPU worker. If components other than the Map and Reduce stages contribute to a large portion of running time, the GPU worker is not able to make large performance acceleration. Second, the GPU worker and the CPU worker have comparable performance. The benefit of using the CPU worker diminishes, as the speedup of the GPU worker over the CPU worker becomes higher.

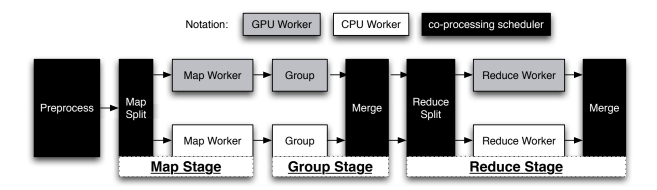


Figure 4.1. The workflow of GPU/CPU co-processing.

With the GPU/CPU co-processing module, Mars can harness the computation power of NVIDIA GPUs, AMD GPUs, and multi-core CPUs on the same machine, by integrating MarsCUDA, Mars-Brook, and MarsCPU modules as components.

### CHAPTER 5

### MULTI-MACHINE IMPLEMENTATIONS

In this chapter, we present the integration of Mars into a CPU-based distributed MapReduce system, specifically Hadoop in our implementation. This integration benefits from both worlds: Hadoop utilizes CPUs on multiple machines and provides fault-tolerance and other features of a distributed system; Mars utilizes the GPU to accelerate local computation. We denote Mars-enabled Hadoop as MarsHadoop.

We use the *Hadoop Streaming* technology <sup>1</sup> to integrate Mars into Hadoop. *Hadoop Streaming* enables the developers to use their own custom Map or Reduce implementation in Hadoop. In our implementation, we use the Mars executable to read the input from *stdin* and to emit the output to *stdout*. Thus, the Map and the Reduce tasks can be performed on the GPU, and other tasks such as task scheduling and failure handling are performed by Hadoop. Finally, since current GPUs do not support multi-tasking, we configure MarsHadoop to run GPU-based tasks sequentially on one GPU.

Figure 5.1 illustrates the workflow of MarsHadoop. A Map Worker/Reduce Worker in MarsHadoop is the same as a Map Worker/Reduce Worker shown in Figure 4.1; in other words, it can be from MarsCUDA, MarsBrook, or MarsCPU, depending on the underlying processor. In the configuration of Figure 5.1, Node 1 simultaneously runs two Map Workers, on a GPU and a CPU respectively.

<sup>&</sup>lt;sup>1</sup>http://hadoop.apache.org/common/docs/r0.15.2/streaming.html

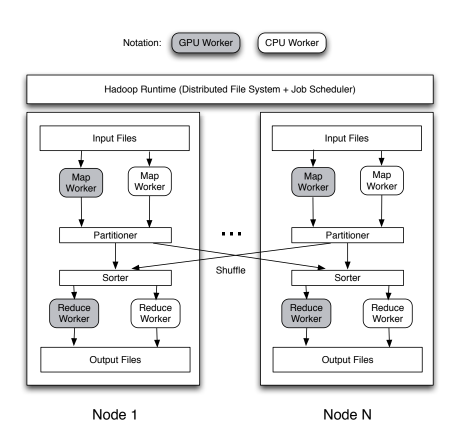


Figure 5.1. MarsHadoop. Some Map and Reduce tasks are performed on the GPU, and others are on the CPU.

### CHAPTER 6

#### EXPERIMENTAL EVALUATION

In this chapter, we evaluate Mars on a single machine using a micro-benchmark of six applications in comparison with their CPU-based counterparts and native GPU-based implementations. We also evaluate the performance of MarsHadoop on two connected machines.

## **6.1** Experimental Setup

Our experiments were performed on three PCs, A, B and C. Table 6.1 shows their hardware configuration. Both PCs A and B run 32-bit CentOS 5.1 Linux with kernel 2.6.18, NVIDIA CUDA 2.2, and the GPU driver 185.18.14. PC C runs 32-bit Windows XP Pro SP3, with Brook+ 1.01.0 beta, and the GPU driver 8.561. All hard drives on these PCs are SATA magnetic hard disks with 7200 rpm. On all PCs, the main memory and the device memory are connected by PCI-E bus with a theoretical bandwidth of 4 GB/sec.

### 6.2 Micro-benchmark

We have implemented the following six real-world applications for evaluating the MapReduce framework.

**String Match (SM):** Each Map task searches a portion of the input file to check whether the target string is in the portion. Neither the *Group* nor the *Reduce* stage is needed.

**Matrix Multiplication (MM):** Matrix multiplication is used intensively in analyzing the relationship of two documents. Given two matrices M and N, each Map task computes multiplication for a row from M and a column from N. It outputs the pair of the row ID and the column ID as the key and the corresponding result as the value. Neither the *Group* nor the *Reduce* stage is needed.

**Black-Scholes:** Black-Scholes model [8] is used for calculating the price for European options according to a partial differential equation. For each option, a Map task computes the prices for the

Table 6.1. Machine configurations

Machine	PC A	PC B	PC C
GPU	NVIDIA GTX280	NVIDIA 8800GTX	ATI Radeon HD 3870
# GPU core	240	128	320
GPU Core Clock (MHz)	602	575	775
GPU Memory Clock (MHz)	1107	900	2250
GPU Memory Bandwidth	141.7	86.4	72.0
(GB/s)			
GPU Memory Capacity (MB)	1024	768	512
CPU	Intel Core2 Quad Q6600	Intel Core2 Quad Q6600	Intel Pentium 4 540
CPU Clock (MHz)	2400	2400	3200
# CPU core	4	4	2
CPU Memory Capacity (MB)	2048	2048	1024

call and put prices of an option, and emits a structure containing the price of the option call and the price of the option put as the key, and the option id as the value. The Group stage is to rank the price of option calls. No Reduce stage is needed.

**Similarity Score (SS):** It is used in web document clustering. The characteristics of a document are represented using a feature vector of floating point numbers. Given two document features,  $\vec{a}$  and  $\vec{b}$ , the similarity score between these two documents is defined to be  $\frac{\vec{a} \cdot \vec{b}}{|\vec{a}| \cdot |\vec{b}|}$ . SS computes the pair-wise similarity score for a set of documents. Each Map task computes the similarity score for two documents. It outputs the intermediate pair with the score as the key and the pair of the two document IDs as the value. The *Group* stage is required to rank the pair-wise similarity scores and no *Reduce* stage is required.

**Principal component analysis (PCA):** This application computes the mean vector and the covariance matrix of a set of points in the first two steps in PCA. The input data is stored in a matrix. The whole process contains two MapReduce invocations in a chain. The first MapReduce procedure is to find the mean for each row in the matrix, and the second is to calculate the covariance matrix. Neither Group nor Reduce stage is needed in the first MapReduce invocation. A Map task computes the mean for a row. In the second invocation, each Map task is to calculate the covariance

of two rows. The Group stage is required to sort the row-pairs by row IDs. No Reduce phase is needed.

Monte Carlo (MC): Monte Carlo [9] is used to compute option pricing in financial engineering. The Monte Carlo numeric integration is to mathematically estimate the expectation of the price of option call. Each Map task is to compute the expected value of a random sample for an option, and to emit the option ID as the key, while the expected value of the random sample as the value. The Group stage and the Reduce stage are required to calculate the mean of all the samples for each option. In this application, all the options have the same number of samples, and the intermediate results are ordered by option ID already. Mars does not need to perform sorting in the Group stage.

The above applications are commonly used in benchmarking MapReduce implementations in the previous studies [13, 28]. SM, MM and PCA are adopted from Phoenix suite [28], SS is a common component in web applications, while BS and MC are prevalent in financial engineering, and are adopted from CUDA SDK. In particular, the workflow of these applications differ: SM and MM only have the *Map* stage, BS, SS and PCA have *Map* and *Group* stages, and MC has all the three stages. PCA has a chain of multiple MapReduce procedures, whereas other applications have only one MapReduce invocation.

Within a single machine, we used three data sets for each application (S, M and L) to evaluate the scalability of the MapReduce framework. The input for SM is textual data, and is adopted from Phoenix [28]; The input for all the other applications contains randomly generated real numbers, ranging from zero to one. All these input data are stored as files in the hard disk. We summarize the size of input data for each application in Table 6.2.

**Metrics.** The wall time is the major metric for the performance evaluation. We measure the elapsed time of each application from reading data from the disk till generating results in the main memory. We ran each experiment five times and report the average value. The variation of elapsed time between runs is negligible. The performance speedup on A over B is defined as the running time of B divided by the running time of A. The performance slowdown on A over B is defined as the running time of A divided by the running time of B.

We use the number of code lines written by the user as the metric on comparing the programmability of different MapReduce implementations as well as the native implementation with CUDA and Brook+. Note that we exclude comments and empty lines from the code size counting.

Table 6.2. The input data sizes of the micro-benchmark

Applications	Small	Medium	Large
String Match	size: 55MB	size: 105MB	size: 160MB
Matrix Multiplication	256x256	512x512	1024x1024
Black-Scholes	# option: 1,000,000	# option: 3,000,000	# option: 5,000,000
Similarity Score	# feature: 128, # doc-	# feature: 128, # doc-	# feature: 128, # doc-
	uments: 512	uments: 1024	uments: 2048
PCA	1000x256	2000x256	4000x256
Monte Carlo	# option: 500, # sam-	# option: 500, # sam-	# option: 500, # sam-
	ples per option: 500	ples per option: 2500	ples per option: 5000

## **6.3** Results on a Single Machine

On a single machine, we have compared the performance and programmability of the MapReduce frameworks between the CPU and the GPU. We have implemented the six applications on MarsCUDA, MarsCPU, and the latest release of Phoenix in version 2.0.0. We have also implemented the applications directly on CUDA and pthreads respectively, including thread configuration, data distribution, task execution, buffer management, and various memory optimizations.

We present the results on the NVIDIA GPU in detail, and briefly present the results on the AMD GPU, mainly demonstrating the feasibility.

#### 1. Results on MarsCUDA and MarsCPU

**Programmability.** Table 6.3 shows the comparison of user code size, for implementing the micro-benchmark with MarsCUDA, MarsCPU, Phoenix, and CUDA. By design, the code sizes with MarsCUDA are the same as those with MarsCPU. In general, the applications with MarsCPU have a smaller code size to those with Phoenix. Phoenix needs additional code to tune the runtime performance, for example, to setup cache sizes and data chunk size, and to specify the partition and locator functions that Mars does not require. If the *Group* stage is required, applications like SS with MarsCUDA have a much smaller code size than that is manually written using CUDA, due to an optimized but lengthy group function on CUDA. The user code size of MarsCUDA is up

to 7 times smaller than that of the native implementation with CUDA. For Matrix Multiplication, CUDA have a smaller code size, because MarsCUDA requires additional code to prepare the input key/value pairs, while the native CUDA implementation does not.

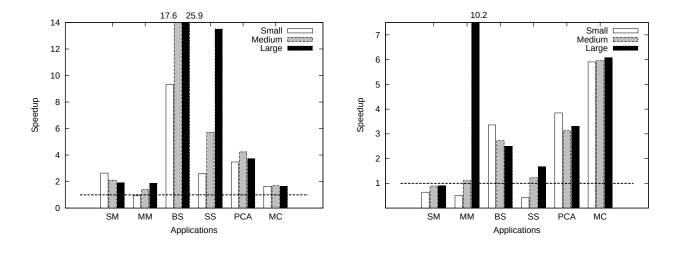
Table 6.3. Comparison of application code size on MarsCPU, MarsCUDA, Phoenix, and CUDA.

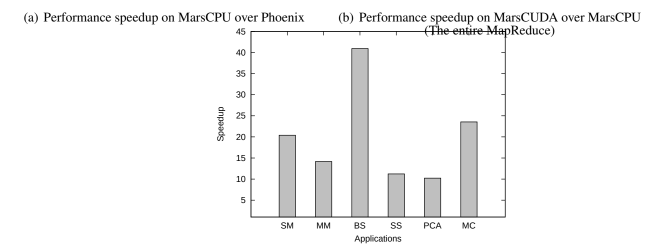
Applications	Phoenix	MarsCUDA/MarsCPU	CUDA
String Match	206	147	157
Matrix Multiplication	178	72	68
Black-Scholes	199	147	721
Similarity Score	125	82	615
Principal component analysis	297	168	583
Monte Carlo	251	203	359

**Overall performance on MapReduce.** We conducted the performance evaluation of MarsCUDA and MarsCPU on PC A by comparing with Phoenix. Figure 6.1 shows the overall performance comparison. Both MarsCUDA and MarsCPU outperform Phoenix for the six applications, due to the general lock-free design of Mars.

The overall performance of MarsCPU is generally better than that of Phoenix, achieving a speedup of up to 25.9x. Applications written using Phoenix always have a *Reduce* stage, whereas using ours they may not have. Phoenix maintains a global 2D array of pointers to keys array. Each keys array is in essence a contiguous buffer as a bucket for hashing, and is sorted by insertion sort when a new key arrives. Such design incurs two serious performance bottlenecks. First, lock-based synchronization is needed. Second, lots of memory buffer movements (calling *memmove()*) are required for insertion sort in the static array. In contrast, the design of Mars is lock-free and each Map task or Reduce task has deterministic output buffer sizes and writing positions, so neither lock nor memory management overhead would be introduced. In particular, BS and SS that require to rank distinct real numbers are over 10x slower on Phoenix than on MarsCPU. That is because Phoenix has to deploy millions of identity reduce tasks for these two applications. Our profiling results obtained from Intel VTune show that over 99% of the total execution time of BS and SS on Phoenix is contributed to the *memmove()* operations in the Reduce stage.

As shown in 6.1(c), MarsCUDA utilizes the GPU hardware to accelerate the Map and Reduce





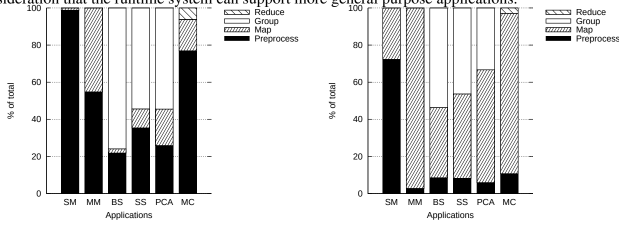
(c) Performance speedup on MarsCUDA over MarsCPU (On large dataset, Map & Reduce stages only)

Figure 6.1. Performance evaluation for MarsCPU and MarsCUDA on the micro-benchmark

stages for the 6 applications, and outperforms MarsCPU in the two stages by 21x on average, and up to 40.9x. Please note that, this speedup is obtained without specific performance tuning on the GPU code, e.g., exploiting local memory. When it turns to the overall performance, MarsCUDA has a 10x speedup over MarsCPU for MM, and 6x for MC, but not so impressive speedup for the other applications (Figure 6.1(b)).

In order to figure out the source of slowdown in overall speedup, we further investigate the time breakdown of each application on the large data set for both MarsCUDA and MarsCPU. We divide the total execution time into four components, including the time for 1) preprocessing input data ("Preprocess"), including input file I/O, generating key/value pairs, and transfering data from

main memory to device memory, 2) the *Map* stage ("Map"), 3) the *Group* stage ("Group"), and 4) the *Reduce* stage ("Reduce"). MarsCUDA generally has a larger portion of preprocess time, involving key-val pair preparation and PCI-E I/O. In addition, the GPU-based Group stage has limited speedup over the CPU-based. We use Amdahl's law to explain this speedup involving parallel and sequential executions. Take SM for example. Although the GPU accelerates the Map phase by 20 times, the Map only takes up some 25% in MarsCPU. According to Amdahl's law, the theoretical speedup of MarsCUDA over MarsCPU is at most 1.3. Our measurement is close to this theoretical speedup. The preprocess is possible to be parallelized on the multi-core CPU for MarsCUDA runtime. However, we leave the parallelization decision to programmers, for the consideration that the runtime system can support more general purpose applications.



(a) Time breakdown of MarsCUDA

(b) Time breakdown of MarsCPU

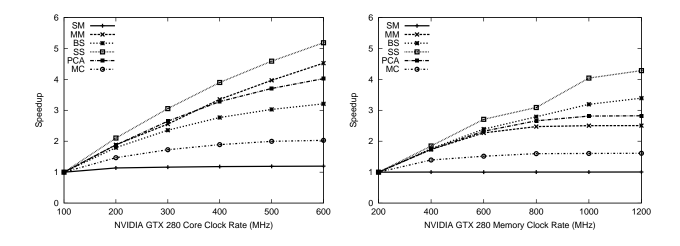
Figure 6.2. Time breakdown of MarsCUDA and MarsCPU on the micro-benchmark

**Scaling.** We used the clock rate scaling tool NVClock <sup>1</sup> to vary the NVIDIA GPU's core clock rate and memory clock rate, in order to evaluate the impact of hardware capability on MarsCUDA. Figures 6.3(a) and 6.3(b) show the performance result of the six applications running on MarsCUDA with the large data set.

In general, most applications (except for SM) on MarsCUDA are sensitive to both core clock rate and memory clock rate. This result indicates that MarsCUDA can scale well as the GPU evolves. SM is not sensitive to the hardware scaling, since its GPU computation time is relatively small (as shown in Figure 6.2(a)).

Comparison with native implementation. Figure 6.4(a) shows the performance slowdown of

<sup>&</sup>lt;sup>1</sup>http://www.linuxhardware.org/nvclock/



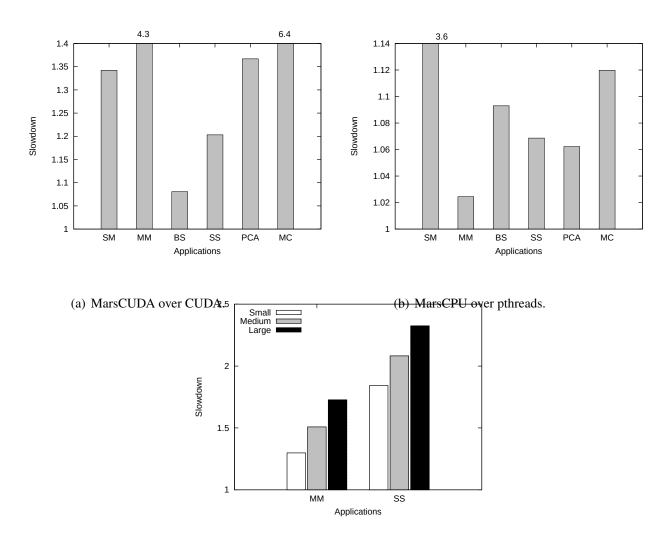
- (a) Baseline: Running at 100 MHz core clock rate. Memory clock rate: fixed to 1100 MHz.
- (b) Baseline: Running at 200 MHz memory clock rate. Core clock rate: fixed to 600 MHz.

Figure 6.3. Varying clock rates on GTX 280.

the six applications on MarsCUDA over the native implementation, with large dataset. Overall, the implementation of applications based on MarsCUDA has roughly the same performance as on CUDA. However, MM and MC perform much poorer on MarsCUDA, mainly due to two reasons. One reason is rooted at the potential deficiency of MapReduce compared with a native implementation, as a previous study has already demonstrated [28]. The other reason is that MarsCUDA does not automatically exploit the local memory to improve the temporal locality due to the lack of knowledge about specific applications. Similarly, Figure 6.4(b) illustrates that applications on MarsCPU has roughly the same performance as on pthreads.

Comparison with other GPU implementations. There are two GPU-based MapReduce implementations in parallel to our work [11, 24], while the source code is not available in public. Therefore, we are not able to conduct empirical performance study by comparing with these two implementations. The peak speedup on the NVIDIA 8800 GTX GPU over on the CPU, reported by Catanzaro [11], is better than ours (150x vs 72x). However, their MapReduce runtime implementation is highly specialized for the machine learning workloads, and they compared with the sequential CPU code, while Mars is for general purpose applications, and we compared with parallel code. The Merge framework [24] reports a peak speedup of about 23x on the Intel X3000 GPU over on the CPU, while their MapReduce design is targeted on Intel's GPUs, and Mars is a general design for different many-core processors.

#### 2. Results on MarsBrook



(c) MarsBrook over Brook+.

Figure 6.4. The performance slowdown of Mars over native implementations.

Due to the limitation of Brook+, we have developed only two numerical applications (i.e., MM and SS) on MarsBrook. Table 6.4 shows the code size of applications written in MarsBrook compared with the native implementation in Brook+. The result is consistent with the comparison between MarsCUDA and the native CUDA implementation. For example, the native implementation of SS has a much larger code size than that on MarsBrook, since SS requires a *Group* stage.

Figure 6.4(c) shows the performance slowdown of two applications by using MarsBrook over the native implementation. The implementation on top of MarsBrook is up to twice slower than the native implementation, which is the price to pay for the user code size reduction.

#### 3. Results on GPU/CPU co-processing of Mars

Table 6.4. Comparison on code sizes of MM and SS using MarsBrook and Brook+.

Applications	MarsBrook	Brook+
MM	66	93
SS	66	611

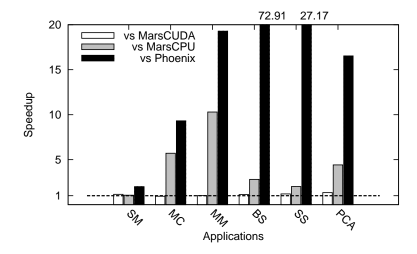
We used MarsCUDA and MarsCPU as two components in the co-processing. Figure 6.5 shows the performance speedup of the GPU/CPU co-processing module over MarsCUDA, MarsCPU, and Phoenix, on the large dataset. Overall, co-processing utilizes the computation power of both the CPU and the GPU, and yields a considerable performance improvement over using MarsCPU or Phoenix on a CPU. However, the speedup of using co-processing over using standalone MarsCUDA is limited.

The workload dispatching between MarsCUDA and MarsCPU in co-processing mainly depends on the performance comparison between the CPU processing and the GPU processing. The theoretical speedup of co-processing over MarsCUDA would be (S+1)/S, where S is the speedup of using MarsCUDA over using MarsCPU. For example, if the speedup S is 10, then using co-processing would only outperform using standalone MarsCUDA by a factor of  $\frac{10+1}{10}=1.1$ . Therefore, for compute-intensive applications MM, BS, SS, MC, and PCA, using co-processing cannot boost the performance considerably over using the standalone MarsCUDA. For SM that spends most time in preprocessing, using co-processing can hardly achieve the theoretical speedup  $\frac{1+1}{1}=2$ . Nevertheless, applications using co-processing of MarsCUDA and MarsCPU still outperforms Phoenix with a speedup of 24 times on average, and 72 times at maximum.

### 6.4 Results on MarsHadoop

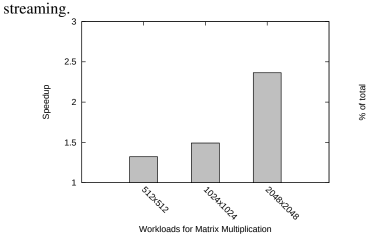
We experimented MM on MarsHadoop. We configured Hadoop on PC A and PC B: PC A as the master node, while PC A itself and PC B as slave nodes.

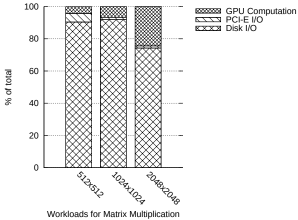
Figure 6.6(a) shows the performance speedup of MarsHadoop over the native Hadoop implementation on MM. As the matrix size varied, MarsHadoop is up to 2.8 times faster than the native Hadoop implementation. We further examine the time breakdown in the slave node, and the results are shown in Figure 6.6(b). As the matrix size increases, the ratio for the computation time grows,



 $Figure\ 6.5.\ Performance\ speedup\ of\ GPU/CPU\ co-processing\ module\ over\ MarsCUDA, MarsCPU, and\ Phoenix.$ 

indicating that Mars starts to help. The disk I/O is mainly due to the extra I/O caused by Hadoop





- (a) Performance speedup on MarsHadoop over native Hadoop.
- (b) Time breakdown on MarsHadoop.

Figure 6.6. Matrix Multiplication on MarsHadoop

### CHAPTER 7

### CONCLUSION

Graphics processors have become an efficient accelerator for high-performance computing. This thesis proposes Mars, which harnesses the GPU computation power and high memory bandwidth to accelerate MapReduce frameworks. Mars is applicable to run on NVIDIA GPUs, AMD GPUs, multi-core CPUs, and Hadoop-based distributed systems. Our empirical studies show that Mars improves the programmability of both the NVIDIA and the AMD GPUs, and the GPU-CPU co-processing of Mars on an NVIDIA GTX280 GPU and an Intel quad-core CPU outperformed Phoenix, the state-of-the-art MapReduce on the multi-core CPU with a speedup of up to 72 times and 24 times on average. Additionally, integrating Mars into Hadoop enabled GPU acceleration for a network of PCs.

The code and documentation of Mars can be found at http://www.cse.ust.hk/gpuqp/.

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