



**UNIVERSITÀ DI PISA**

Computer Science Department

Thesis for Master Degree in Computer Science

**GP-GPU:  
from Data Parallelism  
to Stream Parallelism**

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# CHAPTER 1

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## Introduction

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In a scenario where image processing needed to get more and more sophisticated, we saw *graphic processors* follow the change getting increasingly powerful, not only in computation speed but also in flexibility.

The new elasticity provided by **GPUs** made possible to exploit their benefits for a wide range of non-image-processing problems. This is the beginning of **GP-GPUs** era.

Despite this, enthusiasm slowed down when scientific community had to deal with problems that seemed to be unsuitable for GP-GPUs. However, several studies and researches showed some good results and possibilities, giving oxygen to keep trying mapping to GPU apparently unsuitable problems. This is the core of this work too.

## 1.1 Goals

The main goal of this thesis is to study the behavior of GPUs when used for different purposes, with respect to the usual ones. In particular, we want to use a GPU to run

code that modeled after a *stream parallel pattern*, to understand if we can exploit **Streaming Multiprocessors** (SM) in parallel and what is the relative efficiency.

This means we'll investigate the possibility to perform stream parallelism among "small" data parallel tasks, exploiting the different Streaming Multiprocessors, available on the GP-GPU, as workers.

Then we observed ongoing, in terms of *completion time* and *speedup*.

The latter will consider as sequential version the case that doesn't use CUDA Streams, while the parallel version using them. From these speedups we expected to see an improvement slightly below the number of used CUDA Streams, giving us an assessment on the number of used Streaming Multiprocessors.

The different experiments<sup>1</sup>, gave us results that we expected. In particular we observed that, using CUDA streams, is possible to achieve a gain proportional to the number of SMs available on a GPU. We could see this behavior especially in arithmetic-intensive applications<sup>2</sup>.

In next sections, we're going to show some preliminary details about the concepts we've just introduced.

### 1.1.1 GPU Architecture and Data Parallelism

**GPU** (*Graphics Processing Unit*) is a co-processor, generally known as a highly parallel multiprocessor optimized for parallel graphics computing. Compared with multicore CPUs, manycore GPUs have different architectural design points, one focused on executing many parallel threads efficiently on many cores. This is achieved using simpler cores and optimizing for data parallel behavior among groups of threads[1].

In most of situations, visual processing can be associated to a *data parallel pattern*. In general, we can roughly think to an image as a given and known amount of

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<sup>1</sup>Experiments will be presented in detail in Chapter 5.

<sup>2</sup>We'll show in Chapter 4 the different applications considered and implemented in this work. Chapter 5 will show all results and speedups relative to the different study cases.

*independent* data upon which we want to do the same computations on each of the different data. In most of cases, once the proper granularity of the problem has been chosen, this work should be done for each portion of the image. Considering the above scenario and given that generally a GPU should have to process huge amount of data, we wish to have a lot of threads (lot of cores consequently) doing "the same things" on all data portions.

And that's why GPUs performs their best on data parallel problems.

### 1.1.2 Other Applications

However in recent years we're moving to **GP-GPUs** (***General-purpose computing on graphics processing units***). In other words, lately GPUs have been used for applications other than graphics processing.

One of the first attempts of executing non-graphical computations on a GPU was a matrix-matrix multiply. In 2001, low-end graphics cards had no floating-point support; floating-point color buffers arrived in 2003. For the scientific community the addition of floating point meant no more problems on fixed-point arithmetic overflow.

Other computational advances were possible thanks to programmable shaders, that broke the rigidity of the fixed graphics pipeline (for example LU factorization with partial pivoting on a GPU was one of the first common kernels, that ran faster than an optimized CPU implementation).

The introduction of **NVIDIA's CUDA** (***Compute Unified Device Architecture***) in 2007, ushered a new era of improved performance for many applications as programming GPUs became simpler: terms such as texels, fragments, and pixels were superseded with *threads*, *vector processing*, *data caches* and *shared memory* [2].

In our work we took advantage of CUDA features <sup>3</sup>.

One thing we should point out from GP-GPUs birth: initially scientific applications

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<sup>3</sup>We'll show some further informations about CUDA in Section 1.4.

on GP-GPUs started from matrix (or vector) computations, that mainly could be referred to as ***data parallel problems***. But over time scientific community felt the need to cover other applications, that not necessarily fit data parallel model.

In particular some of latest researches are moving towards ***Task parallel*** applications (sometimes also known as *Irregular-Workloads parallel patterns*).

An example of non-data parallel problem is the *backtracking paradigm*. It's often at the core of compute-and-memory-intensive problems and we can find its application in different cases, such as: constraint satisfaction in AI, maximal clique enumeration in graph mining and k-d tree traversal for ray tracing in graphics.

Some computational parallel patterns perform effectively on a GPU, while the effectiveness of others is still an open issue. In several studies it was highlighted that memory-bound algorithms on the GPU perform at the same level or worse than the corresponding CPU implementation.

Task-parallel systems must deal with different situations with respect to those present in data parallelism, e.g.:

- Handle divergent workflows;
- Handle irregular parallelism;
- Respect dependencies between tasks;
- Implementing efficient load balancing.

Those requirements can lead to inefficient use of the GPU memory hierarchy and SIMD-optimized GPU multi-processors.

However, there have been backtracking-based or other task-parallel algorithms successfully mapped onto the GPU: the most visible example is in *ray tracing* rendering technique; other examples are *H.264 Intra Prediction* video compression encoding, *Reyes Rendering* and Deferred Lighting.



However, in general we cannot expect an order of magnitude increase in performance. Rather, a more realistic goal is to perform at one-two times the CPU performance, which opens up the possibility of building future non-data-parallel algorithms on heterogeneous hardware (such as CPU-GPU clusters) and performing workload-based optimizations [3].

### 1.1.3 GP-GPUs and Stream Parallel

In this work we were interested to a particular type of task parallelism: ***Stream parallelism***.

This means that our tasks are elements of an input stream, of which we don't know a priori the length or the interval times between tasks.

Once the stream elements are available, parallel workers will make independent computations over them and, finally, the manipulated elements should be delivered to some output stream.

We recall as main stream parallel patterns the ***Farm*** and the *Pipeline*, the former being the main subject in this work.

The **Farm parallel pattern**<sup>4</sup> is used to model embarrassingly parallel computations.

This pattern computes in parallel the same function  $f : \alpha \rightarrow \beta$  over all the items appearing in an input stream of type  $\alpha$  **stream** delivering the results on the output stream of type  $\beta$  **stream**.

The model of computation of the task-farm pattern consists of three logical entities: the *Emitter*, that is in charge of accepting input data streams and to assign the data to the Workers; a pool of *Workers* which compute the function  $f$  in parallel over different stream elements; the *Collector* that non-deterministically gathers Workers' partial

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<sup>4</sup>The Farm parallel pattern will be seen in detail in Chapter 3.

results and eventually produces the final result.

The Emitter, the set of Workers and the Collector interact in a pipeline way using a data-flow model which can be implemented in several different ways depending on the target platform. For example, the Emitter and Collector, could be implemented in a centralized way using a single thread, or in a partially or fully distributed way.

Farm's workers can be any other patterns. In the general case, input/output data ordering may be altered due to the different relative speeds of the workers executing the distinct stream items. If ordering is important, it can be enforced by the Collector or by the scheduling/gathering policies of the farm pattern.

*Master-Worker* is a specialization of the task-farm pattern where the Emitter and Collector are collapsed in a single entity (called *master*).

The Workers deliver computed results back to the master. The master schedules received input tasks toward the pool of workers trying to balance their workload [14].

In the case of this thesis, we exploited SMs as Farm Workers, each computing one or more kernel executions, so the function  $f$ , in our case, is given by the kernel code. Furthermore we assumed that Emitter and Collector were both managed by CPU-side. This means that we considered *GPU as Worker* and *CPU as Master*, in particular our implementation is roughly organized as follows:

- ***host-side code*** (code executed by CPU) manages input stream and forwards items to GPU according to *Round-Robin* scheduling policy, furthermore host manages results arriving from GPU;
- ***device-side code*** (code executed by GPU) mainly executes the worker function  $f$ , here called *kernel*<sup>5</sup>.

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<sup>5</sup>In Chapter 2 kernels will be explained, together with other main features from CUDA C++ language.

Since each kernel call is executed by a certain SM, we assumed Streaming Multiprocessors as workers.

For completeness, we also briefly introduce *data parallelism*. It refers to those problems where more workers perform the same task on different portions of data. Generally this is achieved when we have different parallel entities (e.g. threads), such that they execute the same code on different parts of the input data structure.

One of the most important data parallel patterns is *Map*. It computes a given function  $f : \alpha \rightarrow \beta$  over all the data items of an input collection whose elements have type  $\alpha$ . The produced output is a collection of items of type  $\beta$ . Given the input collection  $x_1, x_2, \dots, x_N$ , the output collection is  $y_1, y_2, \dots, y_N$  where  $y_i = f(x_i)$  for  $i = 1, 2, \dots, N$ . Here each data item in the input collection is independent from the other items, so all the elements can be computed in parallel [14].

The model of computation of the map pattern is very similar to the one described for the farm pattern.

The key difference is in the input/output data type:

- *data structures* for Map;
- *streams of items* for Farm.

So, the farm pattern works on streams of independent data (a stream may be unbounded), while the map pattern receives a data collection, of a fixed number of items, that is partitioned among the available computing resources.

The above difference points out one of the main problems of this work: the *Data Transfer times* between **host memory** (CPU side) and **device memory** (GPU side), and vice versa.

In particular, host/device memory copies overhead is a problem because:

- it represents an amount of time spent in memory operations, instead of necessary computations;
- it introduces host/device synchronizations, for example GPU waits for input data copy to end and CPU waits for output data to be fully copied back<sup>6</sup>.

So data transfers, may represent a bottleneck, especially with respect to the arrival rate of input stream items.

We'll show in detail all aspects of this and other minor problems, together with respective solutions, in Chapter 3.

## 1.2 Expectations

The main expectation was to show that a not suitable problem, such as Farm parallel pattern, could fit in a GPU.

It's important to point out that, in particular, we're modeling a Streaming parallel pattern having small data parallel portions as tasks.

In other words, running on GPU our streaming parallel code, it calls a stream of light kernels, each computing a small data parallel task.

Then we wanted to see that in this way we could take an advantage near the order of the number of **SMs** (*Streaming Multiprocessors*).

Looking closer at that this expected results, it means that:

- Data transfer time has to be hidden, in some way, by computation time;
- Kernel executions should take enough time in computations, in order to have chances to achieve overlapping between different kernel executions;

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<sup>6</sup>We'll see in Chapters 2-3 that we'll try to hide this overload by using asynchronous calls.

- The GPU had to achieve a good *Occupancy* <sup>7</sup>.

Once we could achieve these factors, no matter what kind of feature GPU has, we expected to get a *Speedup*  $\approx$  *number of SMs*. The reason why we wanted to see such a speedup is all about gaining some advantages with respect to CPU processing:

- We can delegate streaming problems to the GPU while the CPU can compute other things, this allows to not saturate the CPU (especially when stream has high throughput or each element requires high computation intensity);
- We can split the amount of work between CPU and GPU, the best would be to give respective quantities based on completion time <sup>8</sup>;
- We hopefully want to see a GPU speedup with respect to the CPU, or see the same performances at worst.

## 1.3 Results

At this point, it was useful to experiment different applications, i.e. different kernels codes. More precisely we implemented three types:

- Compute-bound, where the amount of computations, performed by the kernel, is greater than the amount of memory operations;
- Memory-bound, dominated by memory operations;
- Divergent flows (and memory bound too), this means we have a kernel with a lot of branching code<sup>9</sup>.

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<sup>7</sup>We'll insist on occupancy topic in Chapter 3.

<sup>8</sup>See Section 3.5

<sup>9</sup>Chapter 5 explains details about compute-bound, memory-bound and divergent flows in kernels.

In the case of our kernels, memory-bound operations are mainly given by load/store from/to the Global Memory<sup>10</sup>.

We observed that compute-bound kernel gave us the expected results for GPU Farm, that is gaining a good overlap between kernel executions and, so, a speedup proportional to SMs number.

While memory-bound gave a really low amount of overlap and speedup, even worst for the divergent flow case, where we had no speedup at all.

Anyway, the above results represent what we expected.

## 1.4 Tools

As mentioned in Subsection 1.1.2 we mainly exploited NVIDIA's CUDA Toolkit.<sup>11</sup> In particular:

- The code was implemented in `CUDA C++` language, so the compiler was `nvcc`;
- The profiling of GPU code performances was supported by `nvprof` and by its advanced visual version `NVIDIA Nsight`;
- The debugging was made by using `cuda-gdb`;
- Studies on GPU Occupancy have been done with *CUDA Occupancy Calculator spreadsheet* and *Occupancy APIs*.

Tests on the code were implemented as bash scripts and they've been run on two machines:

- The first with four NVIDIA GPUs **Tesla P100-PCIE-16GB**;
- The second with four NVIDIA GPUs **Tesla M40**.

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<sup>10</sup>In Chapter 2 we'll see an overview on the memory organization in GPUs.

<sup>11</sup>In Chapter 2 will be shown all features and details about the aforementioned tools.

The code was developed with the following environments:

- *Visual Studio Code* for CUDA C++, Makefile, bash scripts;
- *Gedit* for Python scripts.

In next chapters all the concepts briefly outlined in this introduction will be discussed in depth.

Chapter 2 introduces an accurate description of all employed tools and how they were used.

Then Chapter 3 explains the logic of the project, with both text and graphical illustrations. In other words here we point out the main ideas and concerns behind our approach and solutions.

Chapter 4 presents and explains main implementation choices and there will be listed some fundamental part of the code.

Then Chapter 5 shows either how experiments and test are set, obtained results and some respective plots.

Finally, conclusions give an overall view of the thesis and some final remarks.

## CHAPTER 2

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### Tools

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Everything in this project has been developed using a GNU/Linux environment. We had two available remote computers, to which we connect via `ssh` command.

In particular we worked on the following machines:

#### 1. Local host

- Ubuntu 14.04 LTS, (4.4.0-148-generic x86\_64)
- 1 CPU Intel® Core™ i5 CPU M 450 @ 2.40GHz x 4

#### 2. P100 remote server

- Ubuntu 18.04.2 LTS (4.15.0-43-generic x86\_64)
- 40 CPUs Intel(R) Xeon(R) CPU E5-2698 v4 @ 2.20GHz, 2 way hyperthreading
- 4 GPUs Tesla P100-PCIE-16GB, 56 SMs with 64 CUDA cores each, 16281 MBytes total Global Memory, 1.33GHz



### 3. M40 remote server

- Ubuntu 16.04.6 LTS (4.4.0-154-generic x86\_64)
- 24 CPUs Intel(R) Xeon(R) CPU E5-2670 v3 @ 2.30GHz, 2 way hyperthreading
- 4 GPUs NVIDIA Tesla M40, 24 SMs with 128 CUDA cores each, 11449 MBytes total Global Memory, 1.11GHz

Given that this work is focused on the use of the remote GPUs, their main specifics are listed in Table 2.1. All the following informations have been get by executing `cudaDeviceQuery` application (located inside samples of CUDA Toolkit).

In this work we mainly made use of the different tools available in the CUDA Toolkit. In the following section will be presented all of employed stuff, with some specifications and how they've been exploited during this project.

## 2.1 NVIDIA Architecture and CUDA

The NVIDIA GPU architecture is built around a scalable array of multithreaded **Streaming Multiprocessors** (SMs).

When a CUDA program on the host CPU invokes a kernel grid, the blocks of the grid are enumerated and distributed to multiprocessors with available execution capacity. The threads of a thread block execute concurrently on one SM, and multiple thread blocks can execute concurrently on one multiprocessor.

As thread blocks terminate, new blocks are launched on the vacated multiprocessors.

	<b>Tesla P100</b>	<b>Tesla M40</b>
<b>Driver/Runtime Version</b>	10.1	10.1
<b>CUDA Capability</b>	6.0	5.2
<b>Tot. global memory amount</b>	16281 MBytes	11449 MBytes
<b>Multiprocessors</b>	56	24
<b>CUDA Cores/MP (Tot. CUDA cores)</b>	64 (3584)	128 (3072)
<b>GPU Max Clock rate</b>	1329 MHz (1.33 GHz)	1112 MHz (1.11 GHz)
<b>Tot. amount constant memory</b>	65536 bytes	65536 bytes
<b>Tot. amount shared memory/block</b>	49152 bytes	49152 bytes
<b>Tot. #registers available/block</b>	65536	65536
<b>Warp size</b>	32	32
<b>Maximum #threads/multiprocessor</b>	2048	2048
<b>Max #threads/block</b>	1024	1024
<b>Max thread block dimensions (x,y,z)</b>	(1024, 1024, 64)	(1024, 1024, 64)
<b>Max grid size dimensions (x,y,z)</b>	(2147483647, 65535, 65535)	(2147483647, 65535, 65535)
<b>Concurrent copy &amp; kernel exec</b>	Yes with 2 copy engine(s)	Yes with 2 copy engine(s)

Table 2.1: GPUs specifics for the two remote machines employed in this project.

A multiprocessor is designed to execute hundreds of threads concurrently. To manage them, it employs a unique architecture called **SIMT** (***Single-Instruction, Multiple-Thread***) that is described in SIMT Architecture. SIMT Architecture and Hardware Multithreading describe the architecture features of the streaming multiprocessor that are common to all devices.

The multiprocessor creates, manages, schedules, and executes *threads in groups of 32 parallel threads* called *warps*. Individual threads, composing a warp, start together at the same program address, but they have their own instruction address counter and register state and are, therefore, free to branch and execute independently. The term warp originates from weaving, the first parallel thread technology. A half-warp is either the first or second half of a warp.

When a multiprocessor is given one or more thread blocks to execute, it partitions them into warps and each warp gets scheduled by a **warp scheduler** for execution. The way a block is partitioned into warps is always the same; each warp contains threads of consecutive and increasing thread IDs, with the first warp containing thread 0. **Thread Hierarchy** describes how thread IDs relate to thread indexes in the block, it will be discussed in the Subsection 2.2.2.

A warp executes one common instruction at a time, so *full efficiency is realized when all 32 threads of a warp agree on their execution path*. If threads of a warp diverge via a *data-dependent conditional branch*, the warp executes each branch path taken, disabling threads that are not on that path. Branch divergence occurs only within a warp; different warps execute independently regardless of whether they are executing common or disjoint code paths. So, this is the situation that occurs in *kernels with divergent flows*, as we mentioned in Chapter 1.

The SIMT architecture is akin to **SIMD** (***Single Instruction, Multiple Data***) vector organizations in that a single instruction controls multiple processing elements. A key difference is that SIMD vector organizations expose the SIMD width to the

software, whereas SIMT instructions specify the execution and branching behavior of a single thread. In contrast with SIMD vector machines, SIMT enables programmers to write thread-level parallel code for independent, scalar threads, as well as data-parallel code for coordinated threads. For the purposes of correctness, the programmer can essentially ignore the SIMT behavior; however, substantial performance improvements can be realized by taking care that the code seldom requires threads in a warp to diverge[19, 9].

The threads of a warp that are participating in the current instruction are called the **active threads**, whereas threads not on the current instruction are *inactive* (disabled). Threads can be inactive for a variety of reasons including having exited earlier than other threads of their warp, having taken a different branch path than the branch path currently executed by the warp, or being the last threads of a block whose number of threads is not a multiple of the warp size[5].

## Hardware Multithreading

The execution context (program counters, registers, etc.) for each warp, processed by a SM, is maintained on-chip during the entire lifetime of the warp. Therefore, switching from one execution context to another has no cost, and at every instruction issue time, a warp scheduler selects a warp that has threads ready to execute its next instruction (the active threads of the warp) and issues the instruction to those threads.

In particular, each multiprocessor has a set of 32-bit **registers** that are partitioned among the warps, and a parallel data cache or **shared memory**<sup>1</sup> that is partitioned among the thread blocks.

The number of blocks and warps that can reside and be processed together on the multiprocessor, for a given kernel, depends on:

- the amount of registers and shared memory used by the kernel;
- the amount of registers and shared memory available on the SM;

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<sup>1</sup>In next sections the GPU memory hierarchy will be explained.

- maximum number of resident blocks per SM;
- maximum number of resident warps per SM.

These limits are a function of the **compute capability**<sup>2</sup> of the device.

If there are not enough registers or shared memory available per SM to process at least one block, the kernel will fail to launch.

The total number of warps in a block is as follows:

$$\text{ceil}(\frac{Th}{W_{size}}, 1)$$

Where  $Th$  is the number of threads per block,  $W_{size}$  is the warp size (which is equal to 32),  $\text{ceil}(x, y)$  is equal to  $x$  rounded up to the nearest multiple of  $y$ [5].

In November 2006, NVIDIA introduced CUDA, a general purpose parallel computing platform and programming model provided for compute engine in NVIDIA GPUs, to solve many complex computational problems (sometimes in a more efficient way than on a CPU).

Three key abstractions are supported, they are exposed to the programmer using a minimal set of language extensions:

- A hierarchy of thread groups;
- Shared memories;
- Barrier synchronization.

These abstractions provide fine-grained data parallelism and thread parallelism, nested within coarse-grained data parallelism and task parallelism.

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<sup>2</sup>Compute Capability is a classifications of features and technical specifications associated to each compute device.

This makes possible to partition the problem into coarse sub-problems –solved independently in parallel by *blocks* of threads –, and each sub-problem into finer pieces –solved cooperatively in parallel by all *threads* within the block –.

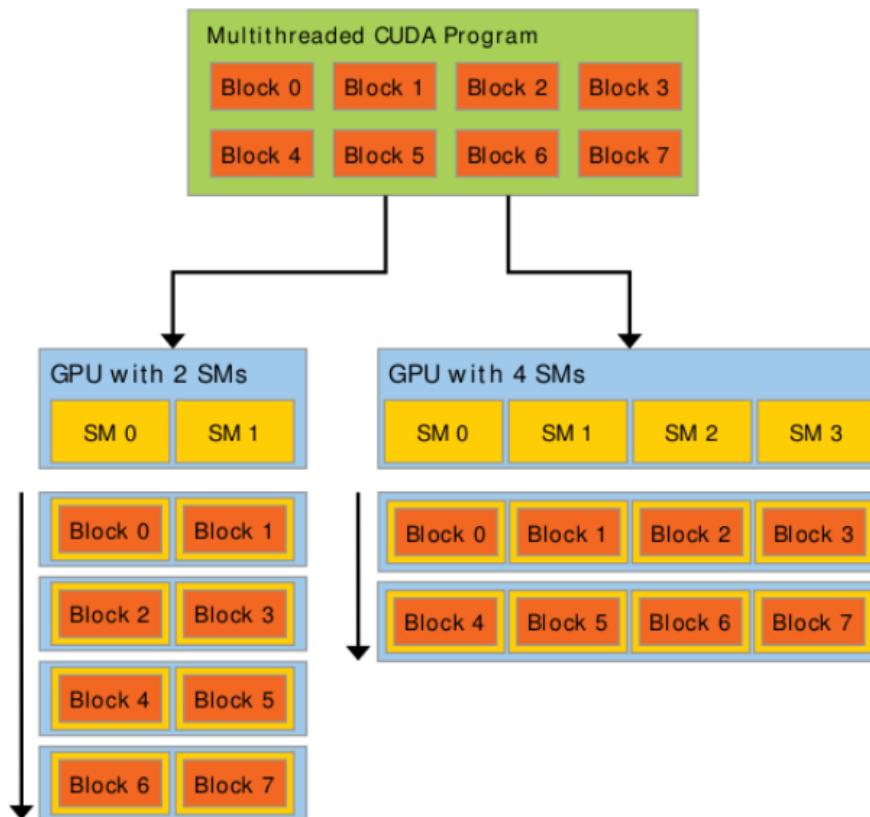


Figure 2.1: GPU scalability.

Indeed, *each block of threads can be scheduled on any of the available Streaming Multiprocessors within a GPU, in any order, concurrently or sequentially*, such that a compiled CUDA program can execute on any number of SM, as illustrated by Figure 2.1, and only the runtime system needs to know the physical multiprocessor count. This programming model scales on the number of multiprocessors and memory partitions [5].

We’ll see in next section further details on CUDA programming model, especially the extensions for C/C++ since, in this thesis, we used CUDA C++.

### 2.1.1 Copy engines and Memory organization

On all operating systems that run CUDA, host memory is virtualized. The operating system component that manages virtual memory is the *Virtuam Memory Manager* (VMM), it provides services to hardware drivers, in order to facilitate direct access of host memory by hardware.

In modern computers, many peripheral, as GPUs, can read or write host memory using a facility known as ***Direct Memory Access*** (DMA). It avoids a data copy and enables the hardware to operate concurrently with the CPU, furthermore hardware may achieve better bus performances over DMA.

To facilitate DMA, operating system VMMs provide a service named *page-locking*. Page-locked memory is ineligible for eviction and its physical addresses cannot change. Once memory is page-locked, drivers can program their DMA hardware to reference the physical addresses of the memory.

Memory that isn't page-locked is known as *pageable*, otherwise it's defined as ***pinned*** memory<sup>3</sup>.

We'll see that ***host pinned memory***, with respect to the GPUs, is generally coupled with CUDA streams and it represents a page-locked portion of host memory, set up for DMA by the current CUDA context<sup>4</sup>.

We'll see in subsection 2.2.3 CUDA Streams and we'll see that they are linked to DMA for another reason. In particular, an important DMA mechanism is the so called ***copy engine***.

DMA allows the transfer of data between host and device, while eventually a kernel is executing on the GPU.

In older GPU architectures there was a single copy engine, while in newer there are

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<sup>3</sup>It's called *pinned* since its physical addresses cannot be changed by the OS.

<sup>4</sup>To be precise page-locked memory and CUDA's pinned aren't really the same thing: pinned memory, registered by CUDA, is mapped for direct access by the GPU; ordinary page-locked memory is not.

usually 2.

The benefit of dual copy engines, coupled with the fact that PCIe<sup>5</sup> is a full duplex interconnection, allows to perform more operations simultaneously, for example:

- do a memory copy from device to host;
- Run kernel that operates on available data in memory;
- do another memory copy from host to device.

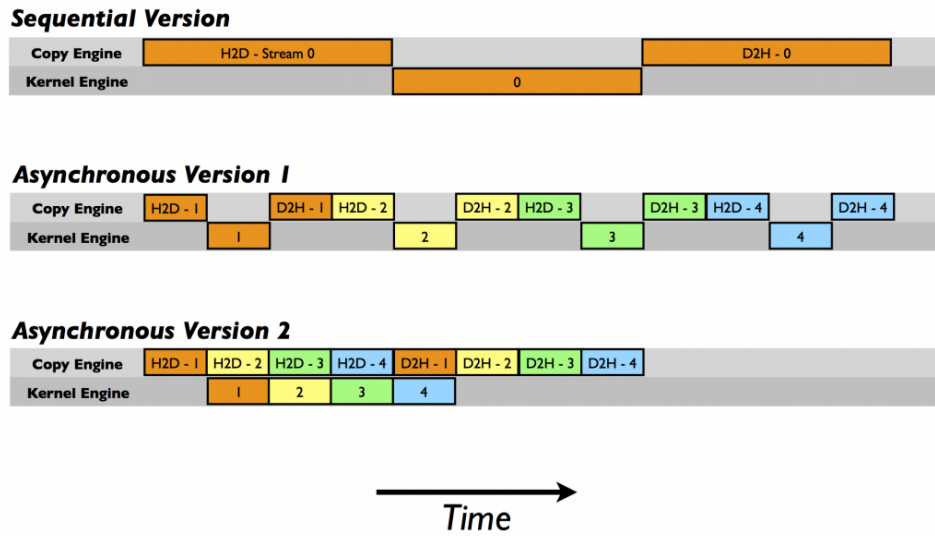


Figure 2.2: Comparison between no CUDA streams, one copy engine and streams, two copy engines and streams.

In the ideal case, all data copies are completely overlapped by kernel execution, i.e. the transfers overheads are hidden by the kernel time<sup>6</sup>.

With a single copy engine, instead, first and last steps above cannot run concurrently<sup>7</sup>[16].

<sup>5</sup>text

<sup>6</sup>We'll see what overlapping really means and how it's implemented in subsection 2.2.3

<sup>7</sup>Note that the simultaneous upload/download of large amounts of data across PCIe can use up a significant portion of available system memory bandwidth.



In Figure 2.2, starting from the top, we have a comparison between the cases of: memory copies and kernel executions completely serial; memory copy and kernel execution overlapping (one copy engine); memory copies and kernel execution overlapping (two copy engines).

To maximize performance, CUDA uses different types of memory, depending on the expected usage.

As mentioned before, host memory refers to the memory attached to the CPU(s) in the system. Device memory is attached to the GPU and accessed by a dedicated memory controller and data should be copied explicitly<sup>8</sup> between host and device memory in order to be processed by the GPU.

Device memory can be allocated and accessed in different of ways:

- **Global memory** may be allocated statically or dynamically and accessed via pointers in CUDA kernels, which translate to global load/store instructions;
- **Constant memory** is read-only memory accessed via different instructions that cause the read requests to be serviced by a cache hierarchy, optimized for broadcast to multiple threads;
- **Local memory** contains the stack, that is local variables that cannot be held in registers, parameters, and return addresses for subroutines;
- **Texture memory** (in the form of CUDA arrays) is accessed via texture and surface load/store instructions; like constant memory, read requests from texture memory are serviced by a separate cache that is optimized for read-only access.

---

<sup>8</sup>A new CUDA functionality, called *Unified Memory* allows to abstract both device and host memories as if they are only one, this avoids the use of explicit copies between host and device memory. However this feature wasn't used in this work, as it wasn't suitable to allow tests and measures we needed to perform in experiments.

Shared memory is an important type of memory in CUDA that is not supported by device memory. Instead, it is an abstraction for an on-chip memory that can be used for fast data interchange between threads within a block. Physically, shared memory comes in the form of built-in memory on the SM.

In this work we mainly exploited Global memory, it is the main abstraction by which CUDA kernels read or write device memory.

The device pointer base resides in the device address space, separate from the CPU address space used by the host code in the CUDA program. As a result, host code in the CUDA program can perform pointer arithmetic on device pointers, but they may not dereference them.

CUDA kernels can read or write global memory using standard C semantics such as pointer indirection (operator\*, operator-&) or array subscripting (operator[])[16].

## 2.2 CUDA C/C++

Here we'll briefly introduce main concepts behind the CUDA programming model, by outlining how they are exposed in C. Especially we'll show important notions about features involved in this project and how/why these were included.

### 2.2.1 Kernels

CUDA C allows to define particular C functions, called *kernels*.

when called, these are executed N times in parallel by N different CUDA threads, instead of executing only once, like it happens in regular C functions.

A kernel code is defined using the `__global__` declaration specifier. The number of CUDA threads that will execute the kernel for a given call is specified using this special execution configuration syntax: `<<<...>>>`.

Each thread executing the kernel is given a unique thread ID, accessible within the kernel through the built-in `threadIdx` variable[5].

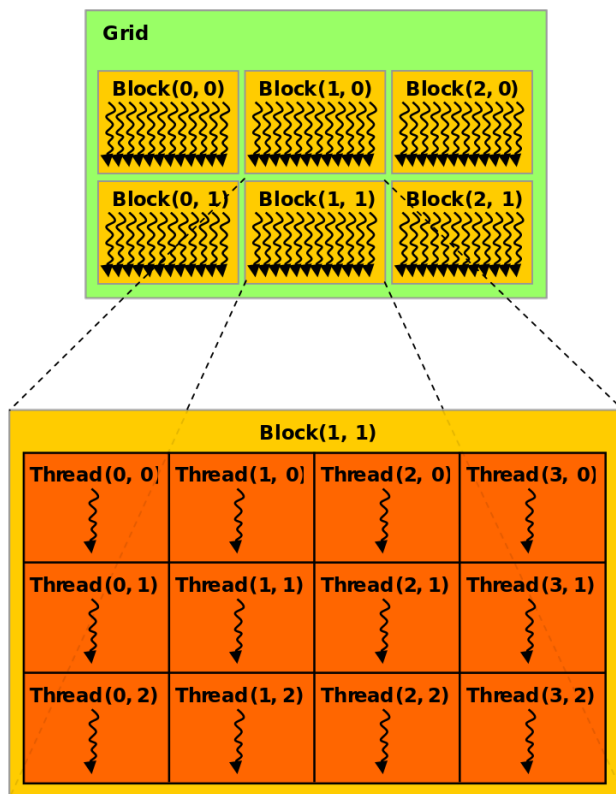


Figure 2.3: Above: a Grid formed by Blocks.

Below: a Block formed by Threads.

## 2.2.2 Thread Hierarchy

In practice `threadIdx` is a 3-component vector, so that threads can be identified using either one, or two, or three dimensional thread index. In turn these threads will form either one, or two, or three dimensional block of threads, called a *thread block*. This provides a way to invoke computation across the elements in domains such as a vectors, matrices, or volumes.

There is a limit to the number of threads per block, since *all threads of a block are expected to reside on the same processor core and must share the limited memory resources of that core*. On current

GPUs, and on the two we worked on, a thread block may contain up to 1024 threads <sup>9</sup>. However, a kernel can be executed by multiple equally-shaped thread blocks, so that

$$Totalnumberofthreads = \#threadsPerBlock \cdot \#blocks$$

Blocks in turn are organized into either one, or two, or three dimensional *grid of thread blocks* as illustrated by Figure 2.3. So, the number of blocks in a grid is usually dictated by the size of the data being processed or the number of processors in the

<sup>9</sup>see Table 2.1 for limits in the machines we used).

system. The number of *threads per block* and the number of *blocks per grid* specified in the <<<...>>> syntax can be of type `int` or `dim3`.

The dimension of the thread block, block index and thread index are accessible within the kernel through the respective built-in variables: `blockDim`, `blockIdx`, `threadIdx`. [5].

### 2.2.3 CUDA Streams

CUDA is best known for enabling fine-grained concurrency, with hardware facilities that enable threads to closely collaborate within blocks, using a combination of shared memory and thread synchronization.

But it also has hardware and software facilities that enable more *coarse-grained concurrency*:

- *CPU/GPU concurrency*. Since they are separate devices, the CPU and GPU can operate independently of each other;
- *Memcpy/kernel processing concurrency*. For GPUs that have one or more copy engines, *host*  $\leftrightarrow$  *device* memcpy can be performed while the SMs are processing kernels;
- *Kernel concurrency*. SM, in 2.x compute capability and later hardware, can run up to 4 kernels in parallel;
- *Multi-GPU concurrency*. For problems with enough computational density, multiple GPUs can operate in parallel.

CUDA streams enable these types of concurrency, but we're interested mainly in the first three types listed above.

Within a given stream, operations are performed in sequential order, but operations in different streams may be performed in parallel.

CUDA events complement CUDA streams by providing the synchronization mechanisms needed to coordinate the parallel execution enabled by streams. CUDA events may be used for CPU/GPU synchronization, for synchronization between the engines on the GPU, and for synchronization between GPUs.

They also provide a GPU-based timing mechanism that cannot be perturbed by system events such as page faults or interrupts from disk or network controllers [16].

### Overlap of Data Transfer and Kernel Execution

Some devices can perform an asynchronous memory copy to or from the GPU concurrently with kernel execution. It's possible to query this capability by checking the `asyncEngineCount` device property<sup>10</sup>, which is greater than zero for devices that support it.

When host memory is involved in the asynchronous copy, it must be *page-locked* to possibly achieve a kernel-copy overlap[18].

Applications manage the concurrent operations described above through **CUDA streams**. A stream is a sequence of commands (possibly issued by different host threads) that execute in order. On the other hand, a stream may execute their commands out of order or concurrently with respect to another; this behavior is not guaranteed and should not be relied upon for correctness (e.g. inter-kernel communication is undefined)[5, 15].

A brief code example:

Listing 2.1: CUDA Strams creation

```
cudaStream_t stream[2];
for (int i = 0; i < 2; ++i)
    cudaStreamCreate(&stream[i]);
float* hostPtr;
cudaMallocHost(&hostPtr, 2 * size);
```

---

<sup>10</sup> See *Device Enumeration* in Table 2.1, for both of our machines, from the *deviceQuery*, we get 2 copy engines.

Each of these streams is defined, by the following code sample, as a sequence of one memory copy *host*  $\rightarrow$  *device*, one kernel launch, and one memory copy *host*  $\leftarrow$  *device*:

Listing 2.2: CUDA Strams and Async example

```
for (int i = 0; i < 2; ++i) {
    cudaMemcpyAsync(inputDevPtr + i * size, hostPtr + i * size, size,
        cudaMemcpyHostToDevice, stream[i]);
    MyKernel <<<100, 512, 0, stream[i]>>> (outputDevPtr + i * size, inputDevPtr + i * size
        , size);
    cudaMemcpyAsync(hostPtr + i * size, outputDevPtr + i * size, size,
        cudaMemcpyDeviceToHost, stream[i]);
}
```

Each stream copies its portion of input array `hostPtr` to array `inputDevPtr` in device memory, processes `inputDevPtr` on the device by calling `MyKernel()`, and copies the result `outputDevPtr` back to the same portion of `hostPtr`. Note that `hostPtr` must point to page-locked host memory for any overlap to occur. Streams are released by calling `cudaStreamDestroy()`.

Listing 2.3: CUDA Strams destroy

```
for (int i = 0; i < 2; ++i)
    cudaStreamDestroy(stream[i]);
```

## 2.2.4 nvcc compiler

To compile the CUDA C++ code it is necessary to use a special compiler, included in CUDA Toolkit, that is `nvcc`.

The CUDA compiler driver hides the details of CUDA compilation from developers. It accepts a range of conventional compiler options, for example for the project we could define macros or include library paths <sup>11</sup>.

---

<sup>11</sup>NVCC documentation: <https://docs.nvidia.com/cuda/cuda-compiler-driver-nvcc/index.html#introduction>

All non-CUDA compilation steps are forwarded to a C++ host compiler that is supported by `nvcc`. Source files compiled with `nvcc` can include a mix of host code and device code. `nvcc`'s basic workflow consists in separating device from host code and then:

- compiling the device code into an assembly form (PTX code) and/or binary form (cubin object),
- and modifying the host code by replacing the `<<<...>>>` syntax introduced in Kernels <sup>12</sup> by the necessary CUDA C runtime function calls, to load and launch each compiled kernel from the PTX code and/or cubin object.

The modified host code is output either as C code that is left to be compiled using another tool, or as object code directly by letting `nvcc` invoke the host compiler during the last compilation stage. Applications can then either link to the compiled host code (most common case), or ignore the modified host code (if any) and use the CUDA driver API to load and execute the PTX code or cubin object.

This compiler can be used almost the same way as a classic `gcc`, for example:

```
nvcc -std=c++14 -g -G -o exec source.cu
```

This is an example that compiles the file `source.cu` in the executable file `exec`, keeping debug information with `-g -G` flags. Here we present compiler versions installed in our two machines:

- **Tesla P100**

`nvcc`: NVIDIA (R) Cuda compiler driver, release 10.1, V10.1.168

- **Tesla M40**

`nvcc`: NVIDIA (R) Cuda compiler driver, release 10.1, V10.1.105

---

<sup>12</sup>See subsection 2.3.1

### 2.2.5 `cuda-gdb` debugger

CUDA-GDB is the NVIDIA tool for debugging CUDA applications (available on Linux). It is an extension to the x86-64 port of GDB, the GNU Project debugger.

CUDA-GDB main features are:

- it provides an environment that allows simultaneous debugging of both GPU and CPU code within the same application;
- as programming in CUDA C is an extension to C programming, debugging with CUDA-GDB is an extension to debugging with GDB, so the existing GDB debugging features are present for debugging the host code (additional features have been provided to support CUDA device code);
- it allows to set breakpoints, to single-step CUDA applications, and also to inspect and modify the memory and variables of any given thread running on the hardware;
- it supports debugging all CUDA applications, whether they use the CUDA driver API, the CUDA runtime API, or both.
- it supports debugging kernels that have been compiled for specific CUDA architectures, but also supports debugging kernels compiled at runtime, referred to as just-in-time (JIT) compilation.

CUDA-GDB was used to debug all compiled source files, both `.cu` and `.cpp`. Mainly it was really helpful in this project to step device code, inspect runtime errors thrown by CUDA API calls and check exactly what code was doing inside our kernels.

## 2.3 Profilers

NVIDIA profiling tools were useful to optimize performances of our CUDA applications. We used two different versions, that will be showed in the following sections.



### 2.3.1 nvprof

Then `nvprof` provides a tool to collect and view profiling data from the command-line. `nvprof` was added to CUDA Toolkit with CUDA 5. It is a command-line profiler, so it's a GUI-less version of the graphical profiling features available in the NVIDIA Visual Profiler.

The `nvprof` profiler enables the collection of a timeline of CUDA-related activities on both CPU and GPU, including kernel execution, memory transfers, memory set and CUDA API calls and events or metrics for CUDA kernels.

After all data is collected, profiling results are displayed in the console <sup>13</sup> or can be saved in a log file for later viewing <sup>14</sup> [6, 7].

`nvprof` operates in different modes: *Summary Mode*, *GPU-Trace and API-Trace Modes*, *Event/metric Summary Mode* and *Event/metric Trace Mode*.

For our purposes we used only *Summary Mode*, this is the default operating mode, where we have a single result line for each kernel function and each type of CUDA memory copy performed by the application (for each operation type are shown number of calls and total, max, min and average time) [6].

We used it, in some situations, as a quick check, for example we exploited it to see if the application wasn't running kernels on the GPU at all, or it was performing an unexpected number of memory copies, etc. To this aim it's enough to run the application with

```
nvprof ./myApp arg0 arg1 ...
```

Given that we wanted to consult profiling results whenever it was necessary, we used `--log-file` option to redirect the output to files for deferred examination.

`nvprof` revealed peculiarly suitable for remote profiling. That's because of the fact command line is faster to check and save an application profiling.

---

<sup>13</sup>The textual output of the profiler is redirected to `stderr` by default.

<sup>14</sup>Or for later import into either `nvprof` or the NVIDIA Visual Profiler).

### 2.3.2 NVIDIA Visual Profiler

The NVIDIA Visual Profiler, introduced in 2008, is a performance profiling tool providing visual feedback for optimizing CUDA C/C++ applications.

The Visual Profiler displays a timeline of an application's activity on both the CPU and GPU to make performance improvement, it analyzes the application to detect potential bottlenecks, using graphical views, that allow to check memory transfers, kernel launches, and other API functions on the same timeline [6].

Visual Profiler was also used in developing phase, to check if code was properly written to hide as much as possible data transfers <sup>15</sup>, even though sometimes it may happen that profilers introduce some sampling synchronization, giving some little imprecision in visual results.

## 2.4 Visual Studio Code

Visual Studio Code is an open-source code editor by Microsoft that also runs on Linux Operating Systems. It includes support for debugging, embedded Git control and GitHub, syntax highlighting, intelligent code completion, snippets, and code refactoring <sup>16</sup>.

Since it's customizable we could add C++ and CUDA editor extensions, furthermore an SFTP extension allowed us to quickly upload/download files from remote machines.

---

<sup>15</sup>We'll see in detail the Overlapping topic and how it was managed in Chapter 3.

<sup>16</sup>Other infos: [https://en.wikipedia.org/wiki/Visual\\_Studio\\_Code](https://en.wikipedia.org/wiki/Visual_Studio_Code)

Documentation: <https://docs.microsoft.com/it-it/dotnet/core/tutorials/with-visual-studio-code>

Website: <https://code.visualstudio.com/>

## 2.5 Tests, Result gathering, Plots

Some other additional tools have been used in this project. In particular we needed:

- to serially run executions of our applications, possibly varying input dataset on interest values;
- to group all results on text files;
- to implement a script to compute averages, speedups and other interest metrics, from results on text files;
- a generator of graphs on important values from the obtained calculations.

### 2.5.1 Bash scripts

Bash (Bourne-Again SHell) is the shell, or command language interpreter, for the GNU operating system; the latter provides other shells, but Bash is the default shell<sup>17</sup>.

**Bash scripts** were needed to implement tests, that run more executions of a certain CUDA application, also varying input dataset. We programmed bash scripts (**.sh**) tests to contain several command as compile a certain CUDA application, run many times the related executable, profile it with **nvprof** and so on.

Below we give a brief example on how we used bash scripts to implement tests:

Listing 2.4: Tests on bash scripts example

```
#!/bin/bash

...
let GPU=3
let B=1024
let nTests=4

Mitters=(10000 400000 800000)
Ns=(57344 114688 229376 458752 917504 1835008)
```

---

<sup>17</sup><https://www.gnu.org/software/bash/manual/>

```

#parameters: deviceID, BLOCK, N_elements, M_iterations, cuStr[bool], strNum
make cleandplow
echo -e "${BLUE}compiling stream...${NC}"

make streamlow

echo ***STREAM_LOW_NOSTREAM_test***

for N in "${Ns[@]}"
do
    for m in "${Mitters[@]}"
    do
        echo sizeN = $N elements, kerM = $m , CUstreams = 0
        echo -n execIndex =

        for((j=0; j<nTests; j+=1));
        do
            echo -n "$j , "
            ./bin/streamlow.out $GPU $B $N $m 0 0 >> ./results/
                dev_cos_dp_stl.txt
        done
        echo -ne "$nTests \n"
        nvprof --log-file ./profiling/dev_str_low$N-$m-$B-0.txt ./bin/
            streamlow.out $GPU $B $N $m 0 0 >> ./results/dev_cos_dp_stl.txt
    done
done
done

```

Here, for example, the code is setting up all input data set, it compiles the code with the **streamlow** rule into an executable, then it loops over all data sets to run the executable in each configuration.

Here we can see that each setting is repeated more times (to allow outliers elimination) and one of those executions is performed via the **nvprof** profiler, in any case all outputs are redirected to consultable **.txt** files.

## 2.5.2 Python scripts

As Python <sup>18</sup> has dynamic typing, together with its interpreted nature, it's an ideal language for scripting and rapid application development.

In the case of this work was most useful to quickly implement a result filter: given text files with time measures, we computed averages and some speedups.

Furthermore we implemented a script to generate some plots on averages and speedups, exploiting the library `matplotlib`<sup>19</sup>.

Below we show a part of the code to compute speedups and generate some plots:

Listing 2.5: Portion of speedup and plots Python script

```
def main():
    res_path = "./output/"+sys.argv[1]+"/"
    for file in os.listdir(res_path):
        if file.endswith("avgs.csv"):
            f=os.path.join(res_path, file)

            resType=file[0:3]
            if resType=="cos":
                dataPar,zeroStream,threeStream,coresStream =
                    getDividedData(dpCosTest,strCosTest,f)
            elif resType=="mat":
                dataPar,zeroStream,threeStream,coresStream =
                    getDividedData(dpMatTest,strMatTest,f)
            ...

            zeroThree,zeroCore,coreDp = getSpeedup(zeroStream,threeStream,
                coresStream,dataPar)

            csvPath=res_path+resType+"_sp.csv"
            ##write speedup to CSV##
            with open(csvPath, "wb") as fcsv:
                writer = csv.writer(fcsv)
                printSpToCSV("Tseq/T3",zeroThree,resType,writer) #num,
                    size, time
```

---

<sup>18</sup>The Python version used to compile, on local host, our scripts is: Python 2.7.6.

See documentation: <https://docs.python.org/2/>

<sup>19</sup><https://matplotlib.org/>

```

        printSpToCSV("Tseq/Tsm",zeroCore,resType,writer)
        printSpToCSV("Tsm/Tdata",coreDp,resType,writer)

#####
# Plots #
#####
# Completion Time
launchPlots(resType,zeroStream,"Zero Streams")
launchPlots(resType,threeStream,"Three Streams")
launchPlots(resType,coresStream,"#SM Streams")
# Speedup
streamNums=[]
if sys.argv[1]=="M40":
    streamNums=[1,3,24]
elif sys.argv[1]=="P100":
    streamNums=[1,3,56]

if resType=="cos":
    plotSpeedup(streamNums, zeroStream, zeroThree,
                zeroCore, resType, cosN, cosM)
elif resType=="mat":
    plotSpeedup(streamNums, zeroStream, zeroThree,
                zeroCore, resType, matNum, matSize)
...

#####
# Speedup plot function #
#####
def plotSpeedup(streamNums,zeroStream, zeroThree , zeroCore, resType,param1,param2):
    title="Speed Up"
    title=title.upper()

    plt.figure(figId)
    plt.plot([0]+streamNums, [0]+streamNums, marker='.',label='Ideal speed up',
             linestyle='--')

    tmp=[]
    j=0
    lineW=0.5
    alphaVal=1.0
    if resType=="cos":
        indexes=[0, param2-1, len(zeroCore)-param2, len(zeroCore)-1]
        for i in indexes:

```

```

        tmp=[1.0, zeroThree[i][2],zeroCore[i][2]]
        lblLine=str(zeroThree[i][0]) + " - " +str(zeroCore[i][1])
        plt.plot(streamNums, tmp, '- ', label = lblLine, marker=markers
                 [j], linestyle='-', linewidth=lineW, alpha=alphaVal)

        lineW+=0.5
        alphaVal -=0.2
        j+=1

    ...

    plt.xticks(streamNums)
    plt.yticks(streamNums)
    plt.legend(loc="upper left")
    plt.grid(axis='both')
    plt.ylabel("Speedup")
    plt.xlabel("#CUDA Streams")
    plt.show()
    plotId+=1
    figId+=1

```

The code above starts in function `main()`, it initially reads from a `.csv` the average time measures for all types of execution by one of the three applications we tested. Then in the main we split times of all execution types (e.g. zero, three or number of SM amount of CUDA streams used). At this point code has what it needs to get all speedups<sup>20</sup> in `getSpeedup`, this function computes speedup (from input parameters) then it saves results in a `.csv` file and output them to the main function too. Finally the main calls the functions that provide completion time and speedup plots, respectively called `launchPlots` and `plotSpeedup`. The latter plots all different speedups, given by the different input data sets of executions, compared with the line of ideal speedup.

---

<sup>20</sup>We'll see in Chapter 5 all types of speedup we compute, how and why.

## CHAPTER 3

---

### Project Logic

---

In this work we started considering the features of problems that can be recognized as a Farm parallel pattern. Then the study moved to consider how a GPU works, its main architectural characteristics and facing its data parallel nature. Next we had to think how to "merge" two such different behaviors, in order to reach reasonable performances<sup>1</sup>, i.e. almost competitive with a classic data parallel problem. Finally, once the main idea behind the development was clear, we had to make some tunings. All of these steps will be shown in detail in next sections.

### 3.1 Stream Parallelism: Farm pattern

Stream parallel patterns describe problems exploiting parallelism among computations relative to different, independent data items appearing on the program input stream at different times. Each independent computation ends with the delivery of one single

---

<sup>1</sup>About expected performances see Section 3.3 and Section 3.4 for more clarifications



item on the program output stream.

We focused on farm pattern, modeling embarrassingly parallel stream parallelism.

The only functional parameter of a farm is the function  $f$  needed to compute the single task. Given a stream of input tasks [4]

$$x_m, \dots, x_1$$

the farm with function  $f$  computes the output stream as

$$f(x_m), \dots, f(x_1)$$

Its parallel semantics ensures it will process the single task in a time close to the time needed to compute  $f$  sequentially. The time between the delivery of two different task results, instead, can be made close to the time spent to compute  $f$  sequentially divided by the number of parallel agents used to execute the farm, i.e. its parallelism degree.

For example, consider the correspondent higher order function description:

```
let rec farm f =
function
EmptyStream -> EmptyStream
| Stream(x,y) -> Stream((f x),(farm f y));;
```

whose type is

$$farm :: (\alpha \rightarrow \beta) \rightarrow \alpha stream \rightarrow \beta stream$$

*The parallel semantics, associated to the higher order functions, states that the computation of any items appearing on the input stream may be performed in parallel, according to the number of available workers.*

In the farm case, according to the parallel semantics, a number of parallel agents, computing function  $f$  onto input data items, equal to the number of items appearing

onto the input stream could be used. This is not realistic, however, for two different reasons:

1. items in the stream do not exist all at the same time, since a stream is not a vector. Items of the stream do appear at different times. Actually, when we talk of consecutive items  $x_i$  and  $x_{i+1}$  of the stream we refer to items appearing onto the stream at times  $t_i$  and  $t_{i+1}$  with  $t_i < t_{i+1}$ . As a consequence, it makes no sense to have a distinct parallel agent for all the items of the input stream, as at any given time only a fraction of the input stream will be available.
2. if we use an agent to compute item  $x_k$ , presumably the computation will end at some time  $t_k$ . If item  $x_j$  appears onto the input stream at a time  $t_j > t_k$  this same agent can be used to compute item  $x_j$  rather than picking up a new agent.

This is why the parallelism degree of a task farm is a critical parameter: a small parallelism degree doesn't exploit all the parallelism available (thus limiting the speedup), while a large parallelism degree may lead to inefficiencies as part of the parallel agents will be probably idle most of time [4].

### 3.1.1 Farm performance model

It's important to consider which kind of performance indicators are useful. When dealing with performance of parallel applications, we are in general interested in two distinct kind of measures:

- those measuring the absolute (wall clock) time spent in the execution of a given (part of) parallel application. Here we can include measures such as
  1. **Latency** ( $L$ ). the time spent between the moment a given activity receives

input data and the moment the same activity delivers the output data corresponding to the input.

2. **Completion time** ( $(T_c)$ ). the overall latency of an application computed on a given input data set, that is the time spent from application start to application completion.
- those measuring the throughput of the application, that is the rate achieved in the delivering of the results. Here In this case we can include measures such as
    1. **Service time** ( $T_s$ ) the time intercurring between the delivery of two successive output items (or alternatively, the time between the acceptance of two consecutive input items), and
    2. **Bandwidth** ( $B$ ) the inverse of the service time.

These are the basic performance measures of interest in parallel/distributed computing. Applications with a very small service time (a high bandwidth) have a good throughput but not necessarily small latencies.

As an example, if we are interested in completing a computation within a deadline, we will be interested in the application completion time rather than in its service time.

Each performance measure has an associated “semantics”:

- $L, T_c$  the latency and the completion time represent “wall clock time” spent in computing a given parallel activity. We are interested in minimizing latency when we want to complete a computation as soon as possible;
- $T_s, B$  service time represents (average) intervals of time incurring between the delivery of two consecutive results (acceptance of two consecutive tasks). We are interested in minimizing service time when we want to have results output as frequently as possible but we don’t care of latency in the computation of a single result. As bandwidth is defined as the inverse of the service time, we are interested in bandwidth maximization in the very same cases.

Given those measures of interest, we can describe an *approximate performance model* for Farm parallel pattern.

First we start considering the service time strictly for workers activity

$$T_s = \frac{T_w}{n_w}$$

So in task farm the service time is given by the average service time of the workers ( $T_w$ ) in the farm, divided by the number of workers ( $n_w$ ), as hopefully  $n_w$  results will be output by the workers every  $T_w$  units of time.

We can add to this model the times spent by emitter and collector, but this will depend strictly on how the Farm is implemented. For example, suppose we have an emitter/collector single process, it gathers input tasks from the input stream and schedules these tasks for execution on one of the available workers. Workers, in turn, receive tasks and once workers compute tasks, they send back to the emitter/collector process the results<sup>2</sup>. This template is often referred to as master-worker pattern.

The three activities –task scheduling, task computation and results gathering and dispatching –happen to be executed in pipeline<sup>3</sup>. Therefore the service time of the master worker is approximated by the maximum of the service times of the three stages. However, first and third stages are executed on the same processing element (the one hosting the emitter/collector concurrent activity), so the model approximation will be

$$T_s(n) = \max\{\frac{T_w}{n_w}, (T_e + T_c)\}$$

Another example could be to have two distinct processing elements, one for the emitter and one for collector, so we could consider the performance model exactly as a three

---

<sup>2</sup>The emitter/collector process –concurrently to its task scheduling activity –collects results, possibly re-order them in such a way the input task order is respected in the output result sequence, and eventually delivers such re-ordered results onto the farm output stream.

<sup>3</sup>A performance model of the pipeline service time states that its service time is the maximum of the service times of the pipeline stages, as it is known that the slowest pipeline stage, sooner or later, will behave as the computation bottleneck.

stages pipeline. This will change the service time approximation in

$$T_s(n) = \max\{\frac{T_w}{n_w}, T_e, T_c\}$$

However, in this work, we wanted to observe behaviors and performances in terms of completion time. We focused on decreasing as much as possible the time needed to compute Streaming parallel problems on GPUs.

Furthermore, when modeling performance of parallel applications, we may also be interested in some derived performance measures. In this case we wanted to derive, from completion times, a speedup measure.

**Speedup** is the ratio between the best known sequential execution time (on the used target architecture) and the parallel execution time. Speedup is a function of  $n$ , the parallelism degree of the parallel execution<sup>4</sup>.

For the moment, considering Farm components (emitter, worker, collector) as pipeline stages, we can give an approximation to completion time performance model for the Farm pattern as follows

$$T_{comp} = T_{emit} + \frac{N}{n_w}(T_{work}) + T_{coll}$$

Where  $N$  is the (theoretically unknown) number of tasks from the input stream and  $n_w$  is the number of workers. Above  $T_{work}$  is the average completion time it takes for a worker to compute a task.

We recall that we don't know a priori the input/output stream length, it can be indefinitely long. But, the above formula means that no matter how many tasks arrive as input stream, the Farm should ideally divide the total completion time among all the workers.

Furthermore, from the above formula, we can see that for each task we'll pay a cost equal to the sum of emitter time, computation time and collector time.

But a more detailed analysis will be given in experiments, Chapter 5.

---

<sup>4</sup>We'll see details on speedup, how it's calculated and how we introduced it in this thesis, in Chapter 5

## 3.2 CPU-GPGPU: heterogeneous architecture

The target of this project was to exploit GP-GPUs high parallelism to lighten the CPU from computation intensive problems, in particular associated to the above explained Farm parallel pattern.

So we had to think what could happen if we wanted to manage such computations on input/output streams, in a way such that:

1. Input stream arrives from host side, being directly generated by CPU or acquired from a remote source;
2. Items are sent from host (main memory) to the GPU (global memory);
3. GPU cores apply specific computations on all items of the stream (as soon as they available in global memory);
4. Finally, computed elements will be copied back to host side and will become the output stream.

In this list our main concern is about data transfer, i.e. step 2 and 4 (as we mentioned in Section 1.1.3). Indeed these phases introduce an overhead per se, but especially in Farm parallel pattern they can represent a not negligible bottleneck.

We should not forget that the input we're handling, is a stream of items: even if elements are available with a high throughput, they come "one by one". As we mentioned in the previous section, we only have a part of input available at one point, so it wouldn't be realistic to have one worker per stream element.

Furthermore, in our case, the items in the input/output streams are data parallel tasks, i.e. each Farm worker will have to process an embarassingly parallel function on the input task.

This means that the tasks are small collections, made up by simple elements such as

float numbers, for example. Here "small collection" means that a single data parallel task has much smaller dimensions, than classical data parallel computations for GPUs<sup>5</sup>.

In our case we can suppose Streaming Multiprocessors to be the Farm workers, so if we don't give enough work (tasks) to each core we would have a resources under-utilization.

Furthermore, we surely don't want to transfer from/to GPU one element at time but, at the same time, in some way we have to keep as much as possible the nature of a Stream parallel patterns.

As a consequence, we had to figure out some mechanisms to:

- Hide as much as we can data transfer times, both in *Host*  $\rightarrow$  *Device* and *Device*  $\rightarrow$  *Host* direction;
- Exploit almost completely our workers resources, in other words try to make Streaming Multiprocessors as busy as possible.

### 3.2.1 Overlapping: Data Transfer hiding

In Section 2.3.3, we introduced CUDA Streams and we recall that they can perform asynchronous memory copies to or from the GPU concurrently with kernel execution. Since the machines we worked on both have concurrency on data copy and kernel execution and both include 2 copy engines<sup>6</sup>, we exploited this capability combined with CUDA streams.

In this way we aimed at achieving a situation in which we could overlap, as much as it was possible, the time it took for the GPU to execute a kernel and the time it took to transfer data back and forth.

As an example see Figure 3.1 to understand a simple case with 3 CUDA Streams. In that diagram we can see the expected behavior of three streams, but we can extend

---

<sup>5</sup>In Chapter 5 we'll see applications and management of Farm streams and tasks.

<sup>6</sup>Chapter 2 explains copy engines utility and functioning.

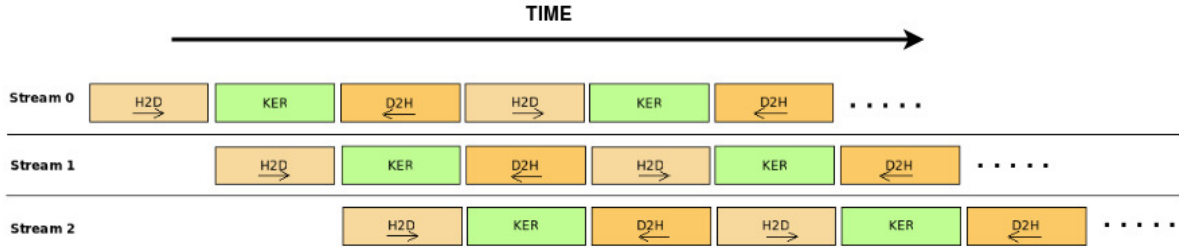


Figure 3.1: Ideal behavior for 3 CUDA Streams.

our expectations to more than three streams, without forgetting that we can have at most 2 data transfer at the same time (given that we've two copy engines).

It's important to point out that not always we can have an ideal behavior in Streams, finding a performance improvement lower than the amount we expected.

Overlap amount depends on several factors: on the order in which the commands are issued to each stream, whether or not the device supports overlap of data transfer and kernel execution, concurrent kernel execution, and/or concurrent data transfers and finally the relative weight of data transfers time and kernel executions time.

Given that our GPUs supported all kind of above mentioned mechanisms, among the above factors the only ones that can have some impact in our case are the order in which commands are issued to each stream, and the kernel/data transfer time ratio.

To improve the potential for concurrent kernel execution, synchronization of any kind should be delayed as long as possible. So, we were careful to avoid *Implicit synchronization*<sup>7</sup> and all unnecessary *Explicit synchronizations*<sup>8</sup>.

Another important face of overlapping, is that it requires to balance Kernels work in such a way it's sufficient to hide the time spent in data transfers, as we quoted just

<sup>7</sup>Implicit synchronization automatically happens when certain host operations are issued in-between commands given from different streams. E.g. this happens in case of: pinned memory allocations, device memory allocations, non-asynchronous memory operations etc.[18].

<sup>8</sup>In CUDA there are several command to force synchronization either between host and device, or between streams etc.



above. This said we can have two possibly unfair scenarios:

- Data transfers take a small amount of time, while kernels are doing lot of computations;
- Data transfers take a big amount of time, with respect to time spent in kernel execution.

The former case may arise when we have heavy computations or an *irregular kernel*. By "irregular" we mean that we're facing an efficient kernel, due to any flow control instruction (`if`, `switch`, `do`, `for`, `while`), that can significantly affect the instruction throughput by causing threads of the same warp to diverge; that is, to follow different execution paths.

If this happens, the different execution paths must be serialized, increasing the total number of instructions executed for this warp. When all the different execution paths have completed, the threads converge back to the same execution path. So we should avoid different execution paths within the same warp[5].

However, sometimes having short memory copy and long kernels can be profitable. As an example we'll see, in Chapter 5, that having more long kernels makes them to overlap more likely both in between them and with data transfers.

The latter case, in the above list, can happen when we move an amount of data such that it takes much more time at each transfer, than it takes in calculations. So in this case the dominant overhead factor would be the data transfer.

### 3.2.2 Occupancy of GPU cores

Once we carried out the stream logic, we had to understand how to try to exploit almost every Streaming Multiprocessor at any given time. This means that we wanted to launch as many kernels as needed to exploit SMs at their best, sometimes this means to arrive near the full *Occupancy* of an SM for each kernel execution, in other

situations it's better to decrease this resource exploitation<sup>9</sup>.

Clearly, when we start, we'll have a portion of time, a sort of "*warm up*" phase, where we'll have first data transfers and kernels launches. So we'll have a narrowed number of running kernels. But as soon as we could have enough data transfers, and therefore enough kernels to execute, we would reach a workload peak on GPU.

In practice, when we just said *lot of kernels*, we meant a lot of small groups of (data parallel) items on which computations have to be applied, in other words each of these elements will be assigned to a thread block.

Let's spend a bit to explain better what Occupancy means.

To ***maximize utilization*** the application should be structured in a way that it exposes as much parallelism as possible and efficiently maps this parallelism to the various components of the system to keep them busy most of the time.

The main ways to maximize utilization can be classified as follows:

1. **Application Level** At a high level, the application should maximize parallel execution between the host, the devices, and the bus connecting the host to the devices, by using *asynchronous functions* calls and streams;
2. **Device Level** At a lower level, the application should maximize parallel execution between the multiprocessors of a device. Multiple kernels can execute concurrently on a device, so maximum utilization can also be achieved by using streams to enable enough kernels to execute concurrently;
3. **Multiprocessor Level** At an even lower level, the application should maximize parallel execution between the various functional units within a multiprocessor. In particular, *a GPU multiprocessor relies on thread-level parallelism to maximize utilization of its functional units.*

---

<sup>9</sup>Chapter 5 will also discuss occupancy and the role it played in our applications and experiments.

From the above, it's clear that occupancy is directly linked to the number of resident warps<sup>10</sup>.

At every instruction issue time, a warp scheduler selects a warp that is ready to execute its next instruction, if any, and issues the relative instructions to the active threads of the warp.

The number of clock cycles it takes for a warp to be ready to execute its next instruction is called the **latency**, and *full utilization is achieved when all warp schedulers always have some instruction to issue for some warp at every clock cycle during that latency period, or in other words, when latency is completely "hidden"*[19, 9].

The most common reason a warp is not ready, to execute its next instruction, is that the instruction's input operands are not available yet.

If all input operands are registers, latency is caused by register dependencies, i.e. some of the input operands are written by some previous instruction(s) whose execution has not completed yet.

So, in this case, the latency is equal to the execution time of the previous instruction and the warp schedulers must schedule instructions for different warps during that time.

Another reason a warp is not ready, to execute its next instruction, is that it is waiting at some *memory fence* (*Memory Fence Functions*) or synchronization point.

A synchronization point can force the multiprocessor to stay idle as more and more warps wait for other warps in the same block to complete execution of instructions.

So, having multiple resident blocks per multiprocessor can help reduce idling in this case, as warps from different blocks do not need to wait for each other at synchronization points.

The *number of blocks and warps residing on each multiprocessor for a given kernel*

---

<sup>10</sup>We recall that warps are groups of 32 threads running in parallel on a set of instructions. Resident warps are those warps that, at a given time, are active on a certain thread block. In Chapter 2 we gave a detailed explanation of these concepts.

*call depends on the execution configuration of the call* (grid and block dimensions), the memory resources of the multiprocessor, and the resource requirements of the kernel [5].

Register and shared memory are others important Occupancy variables, but we didn't focused much on them as on execution configuration. This is because, in our applications, those factors had a negligible impact on eventual occupancy.

At this point, we had to reason about how to maximize Occupancy in our Farm parallel pattern. We shouldn't forget that after looking for full occupancy, experiments can give a prove that a lower one could be better.

First, we have to make some assumptions:

- no shared memory was used;
- we took a really poor amount of registers, given the really simple nature of our example Kernels <sup>11</sup>.

Anyway, chosen kernels represent some *important application categories*.

Then we mainly put our attention on kernel *execution configuration* and number of kernels launched (by different CUDA streams), in order to try to maximize the number of active warps inside each Streaming Multiprocessor.

### 3.2.3 Occupancy drawbacks

Occupancy is a very important factor to take into account, but it's more important to be aware that **occupancy isn't the only factor to take care of**.

In other words, not always trying to achieve maximum occupancy is the best idea, in some cases lower occupancy gives even better performances.

It is common to recommend running more threads per Streaming Multiprocessor and/or

---

<sup>11</sup>We'll see what kind of kernels we used to test the farm parallel pattern, with some code slices in Chapter 4.

running more threads per thread block; the motivation is that this is the main way to hide *latencies*.

Indeed, common beliefs are: multithreading is the only way to hide latency on GPU; shared memory is as fast as registers. Those facts aren't always true.

Some studies demonstrated how was possible to hide arithmetic latency or to hide memory latency using fewer threads, leading to code that runs faster. The *Latency* is the time required to perform an operation, for arithmetic operations it takes  $\approx 20$  cycles; for memory we have  $\approx 400+$  cycles instead.

This, in particular, means that we can't start a dependent operation during these times, but they can be hidden by overlapping with other operations.

```
x = a + b; // takes about 20 cycles to execute
y = a + c; // independent, can start anytime(stall)
z = x + d; // dependent, must wait for completion
```

So *latency hiding* means to do other operations when waiting for latency, this will make code run faster (not faster than the peak). For example another way to hide latency is *Instruction Level Parallelism*<sup>12</sup>.

Furthermore another common belief is that occupancy is a metric of utilization, but, as we anticipate, it's only one of the contributing factors.

Another type of latency is memory-bounded, let's take an example:

```
--global-- void memcpy( float *dst, float *src){
    int block = blockIdx.x+ blockIdx.y* gridDim.x;
    int index = threadIdx.x+ block * blockDim.x;
    float a0 = src[index];
    dst[index] = a0;
}
```

To hide memory latency, using even fewer threads, we can do more parallel work per thread:

---

<sup>12</sup>In general ILP in a kernel is intended as assigning more instructions (possibly equal between them) to each single thread, instead of having a lot of threads executing a lower amount of instructions each. Furthermore ILP can be used to execute independent instructions between two dependent ones[19].

```

__global__ void memcpy( float*dst, float*src){
    int iblock= blockIdx.x+ blockIdx.y* gridDim.x;
    int index = threadIdx.x+ 2 * iblock* blockDim.x;
    float a0 = src[index];
    //no latency stall
    float a1 = src[index+blockDim.x];
    //stall
    dst[index] = a0;
    dst[index+blockDim.x] = a1;
}

```

Note: threads don't stall on memory access, they stall on data dependency instead.

Performances improve copying more floats per thread, instead of copying one and run more blocks and allocate shared memory to control occupancy.

For example some common concepts<sup>13</sup> on CUDA state that:

- "In general, more warps are required if the ratio of the number of instructions with no off-chip memory operands (...) to the number of instructions with off-chip memory operands is low";
- "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.."

For the former there are studies that show how a reduced quantity of warps gives good performances on memory intensive kernels.

For the latter, in reality shared memory bandwidth is lower than register bandwidth, in fact we should use registers to run close to the peak. But requiring more registers can result in having a low occupancy.

So, in many cases, this can be accomplished by computing multiple outputs per thread (see above example on multiple floats copy)[8, 9].

---

<sup>13</sup>From CUDA Programming Guide.

### 3.3 Overall Logic

We have an input stream of items, in particular they are small data parallel tasks, we don't know how much they are and their arrival frequency. The logic of this work can be summarized in the following steps:

1. As items start to arrive, we start to spread them among different CUDA streams, according to a Round-Robin policy;
2. On a certain stream, say `streams[k]`, we send out a task (e.g. the  $k$ -th item that has arrived from the input stream) to the device (GPU Global memory);
3. Immediately after the data transfer call, we launch the kernel execution, with a certain *execution configuration*<sup>14</sup>. The kernel call will be placed to `streams[k]` as well;
4. Once the kernel ends its computations, we copy back to host, on `streams[k]`, the chunk of manipulated data as output item;
5. We'll send each result element onto the output stream.

This behavior is illustrated graphically in Figure 3.2. Here we can see our input stream, from that at a given time we get a certain item, i.e. a task. In the diagram we simplified the concept of (data parallel) tasks, from the input stream, as it was a buffer of size  $k$ ; clearly this is only a graphical simplification. Then, we transfer the several items to Global memory of GPU.

For reasons we showed in the previous sections, it would be unfeasible to work on single simple elements (e.g. float numbers) but, at the same time, we should maintain a pattern as close as possible to Stream parallel. That's why we're working on small data parallel tasks<sup>15</sup> of  $k$  items, where  $k$  is empirically determined to be a relatively small

---

<sup>14</sup>We recall that this is given by grid and block sizes that we set for a certain kernel call.

<sup>15</sup>We represented tasks as arrays, but they can be either small matrices or tiny images, as we'll see in Chapter 4.

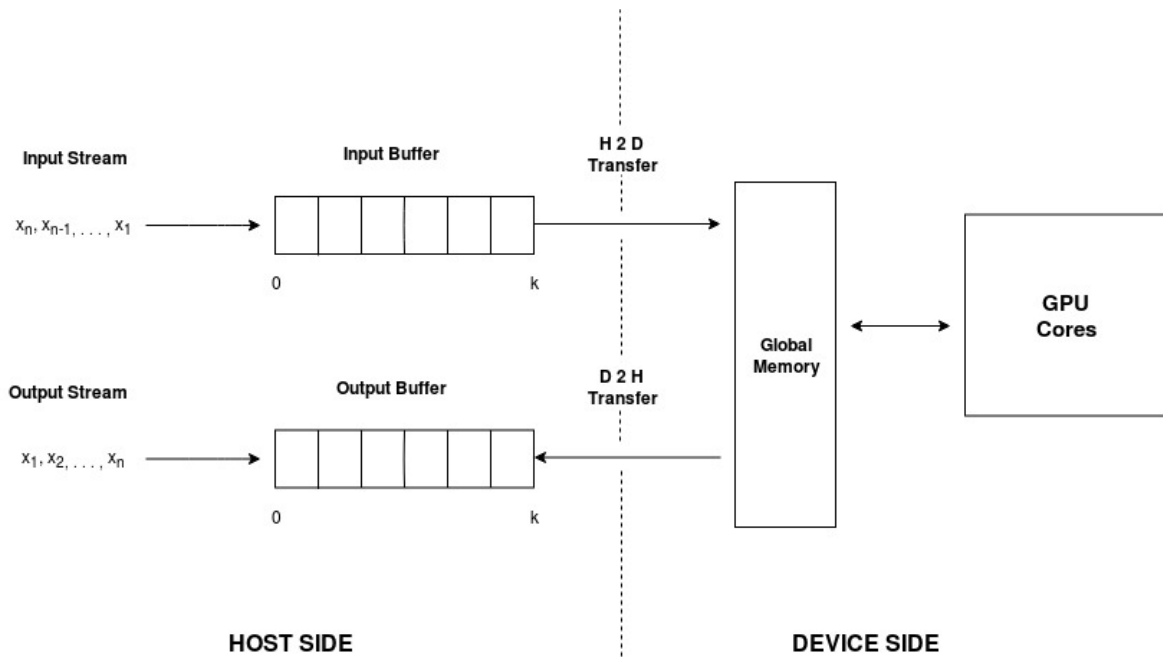


Figure 3.2: Here we have a general and broad graphical representation of our idea on how to fit a Farm parallel pattern on GPU architecture.

number of items and strictly related to execution configuration on kernel, in particular to block size.

Once we start to have several items available on GPU memory, they will be spread all over SMs that have enough available resources. Each task will be splitted in warps (32-thread groups), so each active thread in a warp will compute one instruction per time, in parallel with other threads in the warp. The instruction that will be executed are essentially those specified inside kernel code.

This means that we're executing in parallel the same operations over all the content in a task; in fact, as we mentioned before, each task is a small data parallel collection of items, upon which we're performing data parallel computations.

Since Figure 3.2 is a simplification, it may seems we're sending only one item at time to/from GPU, this would correspond almost to a farm with one worker, processing



one item per time. And this isn't completely what we wanted.

So, here's where CUDA Streams <sup>16</sup> come into play and we used them relying on the following ideas:

1. We have as many streams as Streaming Multiprocessors <sup>17</sup> and, at any given time, each of them hopefully issues a data transfer or a kernel executions;
2. We should come to the point where each stream have issued at least one kernel launch, ideally we expect that each kernel execution is taken over by a certain multiprocessor. So, at a given time, we want to reach a work peak, where almost all SMs are busy;
3. Obviously each kernel execution configuration should be well tuned, in order to take advantage of the maximum of resources in a multiprocessor (according to a certain kernel nature).

All of those parameters have been initially established taking into account of the NVIDIA GPUs' nature, then experimental proves have been performed <sup>18</sup>. Measures and other estimation lead us to consider specific values for those variable parameters. So from the above facts is clear that we're trying to exploit each SM as a Farm parallel worker, furthermore, in such a way that all of those workers are as busy as possible. Note that our **SMs-workers** apply a **kernel-function** to all **tasks**.

Bringing all pieces together we can summarize all project logic in Figure 3.3.

---

<sup>16</sup>Don't confuse input/output stream in Farm parallel pattern with CUDA Streams.

These are two completely different notions: the first refers to the parallel pattern input/output data type, the last refers to special CUDA commands (shown in Section 2.3.3).

<sup>17</sup>Again CUDA Streams are a different concepts with respect to Streaming Multiprocessors. The first are a set of commands, the last are physical processing units inside the GPU.

<sup>18</sup>We'll see in next section more informations about Tunings.

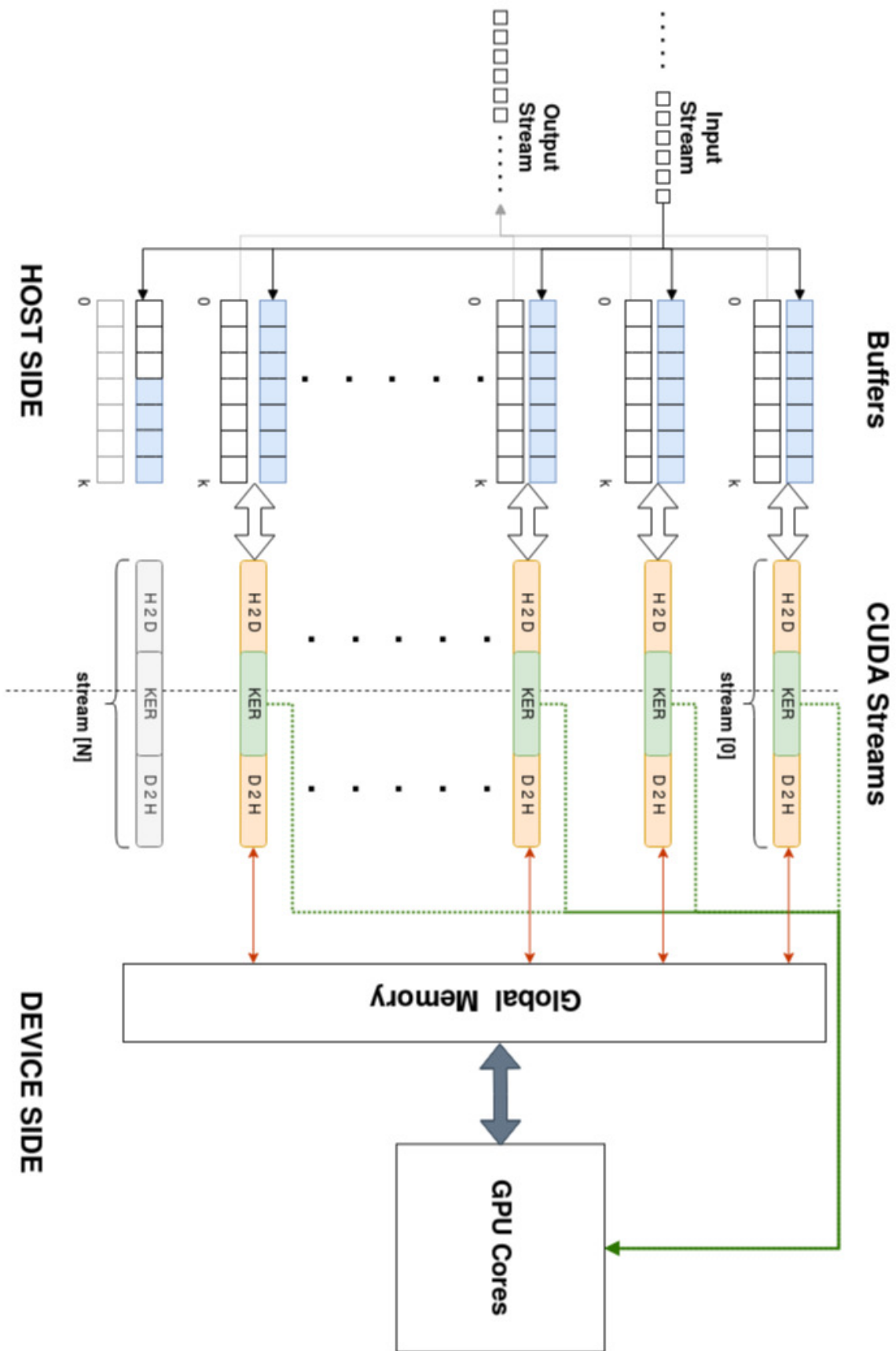


Figure 3.3: Here we have a general and broad graphical representation of our idea on how to fit a Farm parallel pattern on GPU architecture.

The schema may be further detailed as follows:

- We have  $N$  CUDA streams, where  $N$  is the number of Streaming Multiprocessors on the target machine;
- As input stream items arrive, in a Round-Robin way, we spread them all over the CUDA streams as follows:
  1. As soon as we get the  $i^{th}$  task, it's asynchronously sent on `stream [i]` to the GPU, with the command  

```
cudaMemcpyAsync( devBuffer, hostBuffer, bytes, cudaMemcpyHostToDevice,  
                stream [i]);
```
  2. Immediately after we put kernel call, again on the `stream [i]`, to schedule the desired computations on that input item;
  3. Then, asynchronously again, we bring back results to host side, using the instruction  

```
cudaMemcpyAsync( devBuffer, hostBuffer, bytes, cudaMemcpyHostToDevice,  
                stream[i]).
```

Hopefully this approach should make each Streaming Multiprocessor (or at least a part) busy. Initially only few cores will be really busy, but as soon as CUDA streams get full, the pressure on the GPU<sup>19</sup> should increase, so we expected that workload should be enough to almost fill all of Streaming Multiprocessors. In particular, for us this translates in trying to have, at any given time, the maximum number possible of active threads, having to execute instructions, inside each SM.

Figure 3.5 gives a closer look to what we just said.

---

<sup>19</sup>In other words the amount of tasks assigned to each CUDA stream and so to each SM.

From that scheme, looking at violet numbered labels, we can see the order in which we issue commands in a stream, and this will be the order in which they will be issued to device side too, for that stream. The behavior of overlapping between different streams, isn't predictable. Anyway we should take advantage of the fact that, considering two different CUDA streams, we can overlap data transfer and/or kernel execution in a `stream [i]` with the ones in a `stream [k]` (for some  $k \in [0, N - 1]$ , for  $N = \#SMs$ ). Obviously, when the number of CUDA streams is greater than 3, we can have only 2 data transfer operations issued at the same time (by two distinct streams) <sup>20</sup>.

Note that in the Figure 3.5 we represented a single kernel execution as fully occupying an entire SM; in reality not always we'll have this behavior, sometimes it's even convenient to not fully occupy an entire SM with a single kernel launch<sup>21</sup>.

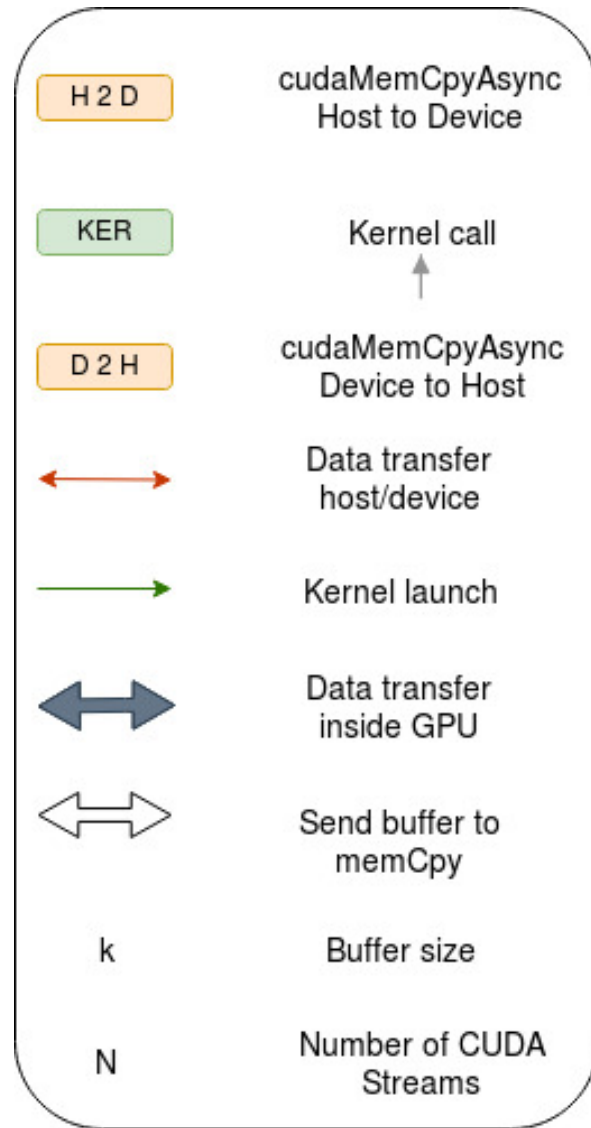


Figure 3.4: Legend about Figure 3.3.

Essentially if all of our reasoning and theories are right, we would expect that we can have an improvement, on completion time, roughly in the order of SMs number

<sup>20</sup>As mentioned in Chapter 2, for concurrent memory copy between host and device, we have 2 copy engines.

<sup>21</sup>We'll see how we practically tried out Streaming Multiprocessors *occupancy* in Chapter 5

with respect to the *serial approach*.

This similarly means that if, for example, we have 3 CUDA Stream we would expect to take an advantage on only at most 3 SMs (at peak work flow), so this should give us an improvement, in completion time, of at most 3 times compared to classical approach. The case of serial approach, instead, processes input stream items without any type of overlapping, it is equivalent to:

- send input to device and wait on host for data transfer completion;
- call the kernel;
- host calls the copy back, from device to host, and keeps waiting until the end of data transfer;
- in the meanwhile kernel is running on GPU;
- finally, only when all computations ended up, results are transferred back to the host, that at this point finishes to wait for output and, so, it can continue with next task.

## 3.4 Tunings

We showed a lot of peculiar behavior and architecture characteristics, as they were taken into account for different implementations, tests datasets and results analysis.

It's clear that, once we decided how to organize our Farm parallel pattern for the GPU, we had to perform experiments and empirical evaluations. This is because:

- It was important to think about a general logic, that wasn't architecture-dependent<sup>22</sup>;
- To validate our idea we had to make a lot of experiments, time measures, examples and counterexamples too;

---

<sup>22</sup>At least we can say that the overall view, showed in Fig. 3.3, can be plausible with almost all NVIDIA architecture having more than two copy engines and allowing concurrent kernel execution.

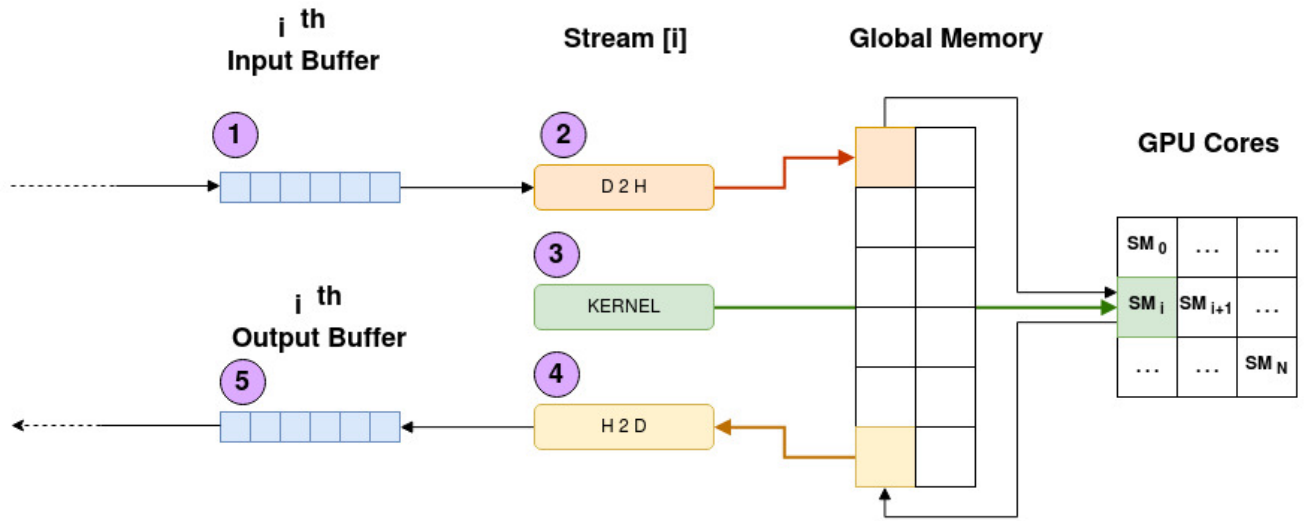


Figure 3.5: In this picture we can see what exactly happens in a certain CUDA Stream. Light violet numbered labels shows the order in which commands are issued by host to a certain stream.

- Clearly experiments required to get a little deeper on NVIDIA GPUs architecture, considering the good practices, apart from the considered application;
- Finally, we had to consider some feature totally application-bounded to launch tests and obtain results of interest.

So after the logical phase, we went through a *tuning phase*, that first had to face general NVIDIA GPUs behavior and structure<sup>23</sup>. Some of the important best practices we tried out in tuning phase were:

- The effect of execution configuration on performance for a given kernel call generally depends on the kernel code, so experimentation is recommended and in fact we followed that approach;
- The number of threads per block should be chosen as a multiple of the warp size (generally equal to 32 threads) to avoid wasting computing resources with under-populated warps<sup>24</sup>;

<sup>23</sup>In Chapter 5 we'll mainly see tunings based on the GPUs used to run tests –**P100** and **M40**.

<sup>24</sup>That's because kernels issue instructions in warps (groups of 32 threads). For example, if we have a

- We exploited *Occupancy Calculator*<sup>25</sup> both in spreadsheet and API functions<sup>26</sup> formats[5].

Given those initial guidelines, it's important to highlight what are variable parameters in Figure 3.3, on which the tuning was made:

- The number of CUDA streams;
- The number of threads per block (block size);
- The number of blocks (grid size).

The first parameter changes mainly as the target machine changes<sup>27</sup>.

The second parameter, in our case, generally depends both on input tasks dimensions and CUDA limits on thread blocks dimensions. But the threads per block choice may be also influenced by the kernel nature and a tuning according to performance measures.

The third is generally roughly determined dividing the size of the input by the block size. However Farm is a particular case and sometimes it's useful to determine the grid size empirically.

The Chapter 5 shows all main parameters tested and their respective performances.

---

block size of 50 threads, the GPU will still issue commands to 64 threads, so we would waste 14 of them idling.

<sup>25</sup>Those tools are included in CUDA Toolkit, they assist programmers in choosing thread block size based on kernel behavior, register and shared memory requirements.

<sup>26</sup>These are special function to call inside the code, we'll see in Chapter 4 a code example on how and where they are used.

<sup>27</sup>We'll see in Chapter 5 that the experiments are performed on zero and three CUDA streams. Furthermore we test the case in which we have as many CUDA streams as SM in the target GPU, so this value will change between P100 and M40 too.

### 3.4.1 Tuning on block and grid dimensions

It's important to understand some main concepts, that are the basis for the logic of this project.

As we mentioned above, the variation on *thread block size* and *grid size*, can affect heavily performances, especially in an extreme scenario as the case study of this thesis<sup>28</sup>. There's a tight correlation in between *Occupancy*, *kernel execution configuration* and kernel code nature. First note that a certain block, whatever its dimension is, will be run on a single Streaming Multiprocessor and once it's assigned it will never be moved. When resources are allocated for a thread block in an SM, it will become an *active block*.

In an SM we can have multiple blocks running independently, each of which grabbing its portion of resources, ie we can have multiple active blocks on a SM until they don't hit the maximum allowed[19, 5]. So, inside a certain SM, some particular cases may happen:

1. Maximum block dimension (aka lot of threads per block), can lead to have a smaller number of blocks<sup>29</sup>;
2. Small block dimension, can give a higher number of blocks running on the same SM<sup>30</sup>.

In the first scenario, we can have cases of good performances in some situations. For example, if few blocks have lot of work to do, it's more likely that they will monopolize

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<sup>28</sup>Here we mention some useful discussions on occupancy, block and grid dimensions from Stack Overflow:  
<https://stackoverflow.com/questions/9985912/>  
<https://stackoverflow.com/questions/5643178/>  
<https://stackoverflow.com/questions/54715373/>

<sup>29</sup>Even only one, as it happens for one of the applications we tested. Anyway, from best practices guide and from profiling this should be a inefficient configuration, so we should be very careful on performances from these situations.

<sup>30</sup>We recall that GPUs have also a limit on the number of active blocks per SM.



resources of the SM in which they're active, making all other waiting blocks scheduled in the same SM, idle for too long (for example in our GPUs it may happens for *blocksize* = 1024).

In the second scenario we can have cases of really poor performances due to low resources exploitation, but for some kinds of kernel we may have a gain, especially in cases as the above mentioned monopolization. So smaller blocks size, means more active blocks on a SM, at any given time.

In general, there is a performance *sweet spot* for middle values on thread block size(for example usually identified in *blocksize* = 512).

So our tests had to face with the above explained behavior too, without forgetting the nature of the various implemented kernels and the relative latencies.

## CHAPTER 4

---

### Implementation

---

Given the logical schema of the previous chapter, we had to build different types of code to test its correctness and efficiency.

In particular, we wanted to distinguish some kernels of interest and adapt them to the Farm parallel pattern on GPU. So, in this project we've implemented the following applications:

- Simple-computational kernel;
- Matrix multiplication;
- Blur Box filter.

### 4.1 Kernels

As we anticipated in the previous section, our kernels mainly doesn't use shared memory. Each kernel clearly is designed such that each thread executes a different element from the input data parallel task, that is one of the input stream's items.

### 4.1.1 Simple-computational kernel

This is a very synthetic kernel, where, given as inputs a data parallel task and a number  $M$ , it computes the cosine applied  $M$  times, for each element in the task. In this case small data parallel tasks are implemented as an array, of size  $N$ , formed by floating point numbers. So, the output will be again a floats array (given by cosines of the input).

This particular kernel has been chosen because it allows:

- to vary the load of computation  $T_{comp}$ , changing the amount of cosine to compute inside the kernel ( $M$ );
- to vary the communication weight  $T_{comm}$ , changing the size of the task ( $N$ ).

The first parameter models the arithmetic intensity, while the second the amount of memory operations. The second parameter, in this case, may include all those mechanisms that perform host/device transfers, but also time spent in transfer between global memory and registers inside the GPU. However, in Farm performance model, the main cost of time we focus on is host/device memory copy, neglecting memory operations internal to GPU.

Note that, since here we're working on one-dimensional data structures, we'll use one-dimensional thread blocks for convenience.

Listing 4.1: Implementation for Simple-Computation Kernel

```
__global__ void cosKernel(int M, int N, float *x_d){
    int idx = offset+blockIdx.x*blockDim.x + threadIdx.x;
    if(idx<N){
        for(int j=0; j<M; ++j)
            x_d[idx]=cosf(x_d[idx]);
    }
    return ;
}
```

This is almost a regular kernel, with no branching and an equal workload for each thread that executes that code.

It is a very useful kernel, because changing a single parameter we could then test situations either of low or high iterations amount, i.e. different workloads.

Moreover this is a computation-bounded application<sup>1</sup>. In particular it models all those kernels that perform a consistent amount of work, on a restrained quantity of data (low amount of memory operations) and that don't diverge during their execution.

### 4.1.2 Matrix multiplication

Here we have the most classical and simple version of matrix multiplication in CUDA. We're always working on input/output streams of small data parallel tasks. However, there's an important difference from the application above, i.e. now we've matrices as tasks, both in input and output. So, for convenience, we'll use two-dimensional thread blocks.

Each couple of threads perform the computation of a single element in result matrix. For example, assume we have `thread[ROW]` and `thread[COL]`, then they will perform `sum += A[ROW, i] * B[i, COL];`, where `i = 0, ..., N` and `sum` will be `C[ROW, COL]`, ie one of the items result matrix.

Listing 4.2: Implementation for Matrix Multiplication Kernel, both non-square and square

```

/**** MATMUL ****/
__global__ void matMulKernel(float* A, float* B, float* C, int m, int k, int n) {
    int ROW = blockIdx.x*blockDim.x+threadIdx.x;
    int COL = blockIdx.y*blockDim.y+threadIdx.y;

    if (ROW<m && COL<n) {
        float tmpSum = 0.0f;

        for (int i = 0; i < k; ++i) {
            tmpSum += A[(ROW*k)+i] * B[(i*n)+COL];
        }
        C[(ROW*n)+COL] = tmpSum;
    }
    return ;
}
```

---

<sup>1</sup>Chapter 5 explains the differences between computation-bound and memory-bound.

```

}

/**** SQUARE MATMUL ****/
__global__ void squareMatMulKernel(float* A, float* B, float* C, int N) {

    int COL = blockIdx.x*blockDim.x+threadIdx.x;
    int ROW = blockIdx.y*blockDim.y+threadIdx.y;

    if (ROW<N && COL<N) {
        float tmpSum=0.0f;

        for (int i = 0; i < N; ++i) {
            tmpSum += A[(ROW*N)+i] * B[(i*N)+COL];
        }
        C[(ROW*N)+COL] = tmpSum;
    }
    return ;
}

```

Matrix multiplication is one of the most widespread applications in GPU computing, this is the basis of other applications too.

It is well known that this kind of very trivial matrix multiplication is quite inefficient. In fact, each for loop iteration will have to perform a multiplication and a sum but, at the same time, the GPU will have to access global memory three times <sup>2</sup>.

This means we have a low arithmetic intensity w.r.t. memory accesses, so threads won't hide memory access latency. That's why in general other optimized algorithms are used, the most known of them is the one decomposing matrices in tiles that will fit in *shared memory*.

Even if this implementation is poor in performances, to our testing purposes is very useful, because it represent a counterexample w.r.t. the implementation presented in the previous section. In fact it's almost the opposite from Simple-computational kernel, as it's a kernel having a low arithmetic intensity with respect to the number of elements involved in computations.

---

<sup>2</sup>We'll have two load from memory for A[i, k] and B[k, j] and a store for C[i, j].

As we mentioned before, just looking at each single iteration we've more memory accesses than arithmetic operations. That's why this type of kernel is considered memory-bound.

### 4.1.3 Blur Box filter

The last type of application implemented in this project is an image processing kernel computing a blur filter.

Again we're working on input/output stream of data parallel tasks that, in this case, are given by small images.

Here input and output pictures are represented as char buffer, items are **RGB** values in  $[0, 255]$  that represent pixels as color tuples.

For each pixel, in the input image, we take the average of each of the pixels in neighborhood (inside the limits of filter size) and we write the average value to the pixel of the output image. This filter is known as a Box blur<sup>3</sup>.

Listing 4.3: Implementation for Image processing Kernel (Blur Box Algorithm)

```
/**** BLURBOX ****/
__global__ void blurBoxFilterKer(unsigned char* input_image, unsigned char*
    output_image, int width, int height) {

    const unsigned int offset = blockIdx.x*blockDim.x+threadIdx.x;
    int dim = width*height*3;
    if(offset<dim){
        int x = offset % width;
        int y = (offset-x)/width;
        int fsize = 5; // Filter size
        if(offset < width*height) {
            float output_red = 0;
            float output_green = 0;
            float output_blue = 0;
            int hits = 0;
```

---

<sup>3</sup>Another blur filter is the *Gaussian blur*, generally preferred as to be more accurate. In fact Box blurs are frequently used to approximate a Gaussian blur. By the central limit theorem, repeated application of a box blur will approximate a Gaussian blur.

```

        for(int ox = -fsize; ox < fsize+1; ++ox) {
            for(int oy = -fsize; oy < fsize+1; ++oy) {
                if((x+ox) > -1 && (x+ox) < width && (y+oy) >
                    -1 && (y+oy) < height) {
                    const int currentoffset = (offset+ox+
                        oy*width)*3;
                    output_red += input_image[
                        currentoffset];
                    output_green += input_image[
                        currentoffset+1];
                    output_blue += input_image[
                        currentoffset+2];
                    hits++;
                }
            }
        }
        output_image[offset*3] = output_red/hits;
        output_image[offset*3+1] = output_green/hits;
        output_image[offset*3+2] = output_blue/hits;
    }
}
return;
}

```

This type of kernel is another counterexample with respect to the Simple-computational kernel.

In fact this application model all those kernels that have low arithmetic intensity, opposed to a consistent amount of memory operations and, furthermore, having a lot of branching instructions. So in this case we're considering not only a memory-bound kernel, but also an example of kernel with divergent flows.

## 4.2 Parallel Patterns implementation on GPU

What really makes the difference in the implementation is how we send data to the GPU and kernel executions configurations. This is what really determines a behavior associated to either a Stream Parallel pattern or a Data Parallel pattern.

### 4.2.1 Stream Parallel on GPU

The code translates the diagram in Fig. 3.3. Let's start by explaining the setting phase:

- The block dimension is provided as command line parameter;
- Then grid dimensions are derived;
- Here we can have two different settings
  - The block has the same dimension of the task, given the capabilities in our machines means having a size  $< 1024$ . The grid will have size one;
  - The task has equal size to the maximum allowed number of active threads in a SM, for both our machines is 2048. Here the grid will be adjusted to cover the task dimension w.r.t. the maximum block dimension (1024 for both machines), i.e.  $GRID = \text{maxThreads}/BLOCK$ .

Listing 4.4: Kernel Launch configuration, ie Grid and Block dimensions setting

```
#ifdef LOWPAR
    GRID = 1;
    chunkSize = BLOCK*GRID;
#else
    GRID = maxThreads/BLOCK;
    chunkSize = BLOCK*GRID;
#endif
```

Now we outline the core of the implementation. The number of CUDA streams to spawn, as we previously told, is equal to the number of Streaming multiprocessor in the target machine. So, we start by distinguish two different cases:

- Number of streams equal to zero, this means we won't use CUDA stream
  1. We allocate space on device for the task, with a `cudaMalloc`;



2. As input stream items arrive, we send them to the device with a simple `cudaMemcpy`<sup>4</sup> (because here we're not using CUDA streams);
3. We launch the Kernel, that will execute as soon as input data is fully copied;
4. We call another `cudaMemcpy` to bring back results to host.

Listing 4.5: Data transfer host/device and kernel call, NO-CUDA Streams version

```
void cosKer(int m, int chunk, float *x, float *cosx, float *x_d)
{
    int xBytes = chunk*sizeof(float);

    gpuErrchk( cudaMemcpy(x_d, x, xBytes, cudaMemcpyHostToDevice) );

    cosKernel<<<GRID, BLOCK>>>(m, chunk, x_d);
    #ifndef MEASURES
        gpuErrchk( cudaPeekAtLastError() );
        gpuErrchk( cudaDeviceSynchronize() );
    #endif

    gpuErrchk( cudaMemcpy( cosx, x_d, xBytes, cudaMemcpyDeviceToHost) );
}
```

- Number of streams greater than zero, this means we will use CUDA streams, so steps are analogous to the previous ones, except for some specific mechanism used, such as:
  1. We allocate on host the space for the tasks, with a `cudaMallocHost` (in order to use CUDA streams and gain best overlapping possible, host should allocate memory as *Pinned*);
  2. We allocate on device the space for the tasks, with a `cudaMalloc`;
  3. In a Round-Robin fashion, we send tasks in an asynchronous way, using `cudaMemcpyAsync`<sup>5</sup>;

---

<sup>4</sup>Note that `cudaMemcpy` is a blocking operation w.r.t. the host, this means that other CUDA calls from the host will be issued after data is fully copied to device.

<sup>5</sup>Note that `cudaMemcpyAsync` is non-blocking for the host, in parameters we'll have to specify which

4. We launch the Kernel in the same CUDA stream of the copy for input data;
5. We call another `cudaMemcpyAsync` to bring back results to host, on the same stream as before.

Note that in first two steps we allocate space not for a single task, but for as many tasks as the number of CUDA streams. This allows us to avoid some explicit synchronizations and to have space reserved for a certain task for each CUDA stream.

Listing 4.6: Data transfer host/device and kernel call, CUDA Streams version

```
void cosKerStream(int m, int chunk, float *x, float *cosx, float *x_d,
                 cudaStream_t strm, int strBytes)
{
    gpuErrchk( cudaMemcpyAsync(x_d, x, strBytes, cudaMemcpyHostToDevice, strm) );

    cosKernel<<<GRID, BLOCK, 0, strm>>>(m, chunk, x_d);

    #ifndef MEASURES
        gpuErrchk( cudaPeekAtLastError() );
        gpuErrchk( cudaDeviceSynchronize() );
    #endif
    gpuErrchk( cudaMemcpyAsync( cosx, x_d, strBytes, cudaMemcpyDeviceToHost,
                               strm ) );
}
```

The reason why we implemented these two versions, of Farm Parallel Pattern on GPU, is that we want to show the gain obtained with CUDA Stream.

In particular we want to compare:

- the implementation without CUDA streams, that represents the serial version for each of our applications;
- the implementation using CUDA streams, that correspond to the parallel version for each of our applications.

---

stream will issue the copy.

Given that we're working on a Streaming parallel problem, suppose we have an input stream of items, that are "small" data parallel tasks.

So, in the first case from the above list, we'll have a serial computation of the input stream's tasks.

While, in the second case, we consider SMs as *Farm workers*, that executes different tasks in parallel.

Therefore, we want to show that with CUDA Streams and some other adjustments<sup>6</sup>, a Stream Parallel Pattern would have performances near to Data Parallel ones (comparing them for the same application and the same total amount of computed data).

It may be interesting to see how the Round Robin scheduler sends buffers to the function `cosKerStream` via CUDA streams. We present a pseudo-code version that shows only its main features:

Listing 4.7: Host side pseudo-code: input stream + kernel launcher function

```
const int streamBytes = chunkSize*sizeof(float) ;
int strSize = nStreams*chunkSize;
//host pinned mem
gpuErrchk( cudaMallocHost((void **)&x, strSize*sizeof(float)) );
gpuErrchk( cudaMallocHost((void **)&cosx, strSize*sizeof(float)) ); //pinned cosx
//device memory
gpuErrchk( cudaMalloc((void **)&x_d, strSize*sizeof(float)) );
//stream array and events creation
cudaStream_t streams[nStreams];
streamCreate(streams, nStreams);
createAndStartEvent(&startEvent, &stopEvent);

int k=0;
while (InputStream) {
    if (buffer x[i: i+chunkSize] is full)
    {
        int i = k%nStreams;
        int strOffs = i*chunkSize;
```

---

<sup>6</sup>Chapter 5 shows how we set kernel execution configurations and other relevant assumptions.

```

        cosKerStream( M_iterations, chunkSize, x[i: i+chunkSize], cosx[i: i+
            chunkSize], x_d[i: i+chunkSize], streams[i], streamBytes);

        send output buffer cosx[i: i+chunkSize] to output stream

        ++k;
    }
    else
    {
        add item to buffer x[i: i+chunkSize]
    }
}
msTot = endEvent(&startEvent, &stopEvent);
streamDestroy(streams, nStreams);

```

It's interesting to highlight the use of *CUDA Events*, they were useful to measure the completion time of memory copies and kernel executions. So they allowed us to make device side measures, that are the main concern in this project <sup>7</sup>.

As we can see we presented most important code listings relative to the Simple-computational kernel. The structure and the implementation to execute Matrix multiplication and Blur Box are similar to Simple-comp kernel, since they are really different applications but each of them is, however, modeling a Farm parallel pattern with small data parallel tasks.

## 4.2.2 Data Parallel un GPU

To prove that our Farm Pattern had acceptable performances, it was useful to compare it with its respective Data Parallel version.

Clearly, to have control on time probes and to compare such two different models, we set a maximum length on input stream for Stream parallel version. In the reality we know that we cannot have such informations on input/output streams. So we make the assumption to know input stream length only for a time measuring purpose.

So, supposing to have an input stream of `N_size` items length, this allows us to

---

<sup>7</sup>We'll explain other details on measures on Chapter 5.

compare our Farm model with a Data Parallel one. The latter will send to the GPU a single data structure in once, having a suitable size such that in total it will allow to perform the same amount of work that is done in Stream parallel version.

So pure data parallel computes a single and big data structure, in a canonical configuration kernel<sup>8</sup> and sends back the output again <sup>9</sup>

Listing 4.8: Optimal Kernel launcher for Simple-Computation kernel, uses APIs to get best Block configuration

```
x = (float *) malloc(N_size*sizeof(float));
cosx = (float *) malloc(N_size*sizeof(float));
gpuErrchk( cudaMalloc((void**)&x_d, N_size*sizeof(float)) );

generate N_size items and put in the "x" data structure

createAndStartEvent(&startEvent, &stopEvent);

float msKer = optimalCosKer(M_iter, N_size, x, cosx, x_d, clocks, clocks_d);

/** Kernel launcher */
float optimalCosKer( int m, int n, float *x, float *cosx, float *x_d){
    int gridSize;    // The actual grid size needed, based on input size
    int minGridSize; // The min grid size needed to achieve the maximum occupancy
                     // for a full device launch
    cudaEvent_t startEvent, stopEvent;

    cudaOccupancyMaxPotentialBlockSize( &minGridSize, &BLOCK, cosKernel, 0, 0);
    GRID = (n + BLOCK - 1) / BLOCK; // Round up according to array size

    gpuErrchk( cudaMalloc((void**)&clocks_d, GRID*sizeof(int)) );
    createAndStartEvent(&startEvent, &stopEvent);

    gpuErrchk( cudaMemcpy(x_d, x, n*sizeof(float), cudaMemcpyHostToDevice) );

    cosKernel<<<gridSize, blockSize>>>(m, n, x_d);
```

---

<sup>8</sup>A classic configuration, in general, is to experiment for block size and get grid size dividing the data dimension by the block size.

<sup>9</sup>And this is how generally the GPU is meant to be used and this is the kind of problem a GPU is designed for.

```

    gpuErrchk( cudaMemcpy( cosx, x_d, n*sizeof(float), cudaMemcpyDeviceToHost) );

    gpuErrchk( cudaPeekAtLastError() );
    cudaDeviceSynchronize();
    float ms = endEvent(&startEvent, &stopEvent);

    return ms;
}

```

The peculiarity of this Kernel is that we exploited CUDA Occupancy APIs. The occupancy-based launch configurator APIs, `cudaOccupancyMaxPotentialBlockSize`, heuristically calculate an execution configuration (thread block and grid sizes) that achieves the maximum multiprocessor-level occupancy[5].

This was one example of use for CUDA Occupancy calculator tools, in this case we used it to achieve the best block and grid configuration possible for our Kernel.

Again the code presented above is relative to Cosine kernel, but the implementation structure is analogous to the one for Matrix multiplication and Blur Box.

## CHAPTER 5

---

### Experiments

---

In this chapter will be shown all the experiments performed and their results. The first section starts from what we expect to get from different code tests and, to this aim, what kind of comparisons will be made.

The second section, for each kernel implementation, will explain how tests are set up, i.e. the chosen datasets for each type of code to test, then some scripts main features and, finally, what results we get. In particular, here we'll show time measures and plots with some remarks. The last section gives a brief summary and some final consideration. Furthermore it gives comparisons between stream parallel and data parallel version, for each kernel type.

### 5.1 Expectations

As previously mentioned, what we want to see is that our model and implementation for Farm parallel pattern can fit in a GPU. To this aim is necessary to gain a speedup in the order of the number of Streaming multiprocessors of the GPU we're running code.

Let's clarify some concepts in the sentence above:

- The speedup will be estimated in terms of **GPU completion time**, i.e. the total time needed to perform all host/device data transfers and kernel executions for a certain application;
- We expect to have the best speedup only when we have certain conditions;
- The best speedup would be in the order of multiprocessors number.

The last point means we can't expect to reach greater gain than the available amount of hardware resources.

Further and specific definitions about these concepts will be given in Speedup subsection. The second point above means we can expect best performances in the following cases:

- When we have a regular kernel, that is a kernel with the lowest possible amount of branching and, thus, very low (or absent) threads divergence;
- When the kernel is more computational-bound than memory-bound, the less access to Global memory the less data transfer latency will slow down execution and this may generally lead to bad occupancy and/or kernels that could not overlap (or they do it rarely);
- When the kernel execution takes an amount of time near the one for data transfer and/or other kernel calls.

When one, or more, of the above conditions isn't met, we're aware to have a considerably lower speedup than what we expected.

### 5.1.1 Measures: What and How

Before the test setup and writing, it's important to understand what we should measure, in order to get significant comparisons.



First we recall that the measures of interest are relative to *data transfers* and *kernel execution* and all completion times are reported in **milliseconds**.

In the case where CUDA streams are used, we have an additional time cost to create and destroy streams, especially when lot of streams are spawned.

If we want to have as many CUDA Stream as many SMs in device, then creation costs in the order of hundreds of milliseconds and destruction in the order of hundred of microseconds<sup>1</sup>. However, we won't sum them up with measures on data transfer and kernel execution. This is because, even if the streams overhead can be notable, it's a one-time cost to pay.

This means that it won't weigh on performances of a Stream parallel application, given that initially we create CUDA streams, then we'll run kernels on a indefinitely long input stream (theoretically) and, only when input is totally consumed out, CUDA streams will be destroyed. So on a reasonably long input stream, the CUDA streams APIs cost should become negligible.

So focusing on data transfers and kernels, we put two time probes, one before the start of the input stream loop and one at the end. The time probes are implemented using **CUDA Events**.

Below will be reported a pseudo-code to clarify how the probes are placed inside the code:

```
/**** Code with events time probes *****/
streamCreate(streams, nStreams); // Create CUDA streams

createAndStartEvent(&startEvent, &stopEvent); // Create "start" and "stop" events,
start recording

int k = 0;
while (InputStream) {
    if (buffer x[i: i+chunkSize] is full)
    {
```

---

<sup>1</sup>Measures on CUDA Streams spawn/deletion were collected with *nvprof* log file, where all CUDA APIs time are precisely measured.

```

        int i = k%nStreams;

        kernelCaller(input_host, output_host, input_device, output_device,
                      streams[i], streamBytes, ...);

        . . . .

        ++k;
    }
    else
    {
        add item to buffer x[i: i+chunkSize]
    }
}
msTot = endEvent(&startEvent, &stopEvent);
cudaEventDestroy();

**** Events Creation and start ****
void createAndStartEvent(cudaEvent_t *startEvent, cudaEvent_t *stopEvent)
{
    gpuErrchk( cudaEventCreate(startEvent) );
    gpuErrchk( cudaEventCreate(stopEvent) );
    gpuErrchk( cudaEventRecord(*startEvent,0) );
}

**** Events end and time measure collection ****
float endEvent(cudaEvent_t *startEvent, cudaEvent_t *stopEvent)
{
    float ms = 0.0f;
    gpuErrchk( cudaEventRecord(*stopEvent, 0) );
    gpuErrchk( cudaEventSynchronize(*stopEvent) );
    gpuErrchk( cudaEventElapsedTime(&ms, *startEvent, *stopEvent) );
    return ms;
}

**** Kernel caller example ****
void kernelCaller(input_host, output_host, input_device, output_device, streams[i],
                  streamBytes, ...)
{
    // H2D mem copy
    gpuErrchk( cudaMemcpyAsync(input_device, input_host, streamBytes,
                               cudaMemcpyHostToDevice, streams[i]) );
    // Kernel call

```

```

        ernel<<<GRID, BLOCK, 0, streams[i]>>>(input_device, output_device, ...);
#ifdef MEASURES
        gpuErrchk( cudaPeekAtLastError() );
#endif
        // D2H mem copy
        gpuErrchk( cudaMemcpyAsync( output_host, output_device, streamBytes,
                                    cudaMemcpyDeviceToHost, streams[i]) );
}

```

CUDA event APIs are a device-bound tool and they were chosen as inside-code measurement for several reasons. Another approach could be to use any CPU timer provided for C++ in a way such as:

```

t1 = myCPUTimer();
Kernel<<<GRID, BLOCK>>>(param0, param1, ...);
cudaDeviceSynchronize();
t2 = myCPUTimer();

```

A problem with using host-device synchronization points, such as `cudaDeviceSynchronize()`, is that they stall the GPU pipeline. Events, instead, provide a relatively light-weight<sup>2</sup> alternative to CPU timers via the *CUDA event API*. This API includes calls to create and destroy events, record events, and compute the elapsed time in milliseconds between two recorded events, exactly as it's shown in code Listing 5.1.1.

CUDA events are of type `cudaEvent_t` and are created and destroyed with `cudaEventCreate()` and `cudaEventDestroy()`. In the above code `cudaEventRecord()` places the start and stop events into the default stream, or `stream 0` (also called the “*Null Stream*”). This holds for all device timers we introduced in our code.

The `cudaEventRecord()` will record a time stamp in device for the event, but only when that event is reached in the specified stream. The function `cudaEventSynchronize()` blocks CPU execution until the specified event is recorded.

The `cudaEventElapsedTime()` function returns the number of milliseconds elapsed between the recording of *start* and *stop*. This value has a resolution of approximately 0.5 microseconds [10, 5]. So those timers will be enough accurate for our purpose, since

---

<sup>2</sup>CUDA events make use of the concept of CUDA streams.

we'll see that almost all elapsed times will be from tens to thousands milliseconds.

It's important to point out why we used events on the default stream. Given the asynchronous nature of CUDA calls, that we perform in non-default stream, the behavior and order in between different streams is unpredictable. This means that a call from a different non-null stream can actually be issued in between two events we're trying to recording, even if they were issued from the same non-default stream.

This is one of the reasons why we chose to put timers outside the loop over input stream. We could insert events inside the loop, instead, in that way we'd have measured singularly each iteration<sup>3</sup> and sum up all those elapsed times. There would have been three problems with that approach:

- Each "end" event, must be sure to measure everything until the ending event, that's why it's necessary to introduce `cudaEventSynchronize()`;
- Given that the input stream should be quite long, all those timers in each loop iteration would have introduced an amount of undesired sampling overhead, apart from synchronization time.

For first problem, we recall that `cudaEventSynchronize()` blocks CPU execution until the specified event is recorded, but we really want to avoid that. We should avoid as much (explicit) host-device synchronizations as we can: given that we're working on input/output streams of items from host, "stopping" this flow on host side at each iteration would invalidate the gain of our model, increasing the overall completion time (probably for a non negligible amount).

The second problem is related to the first. Even if events are a light-weight solution for device activities timing, it doesn't mean they don't introduce a bit of overhead (in addition to the synchronization one) in both host and device side.

For completeness, we'll show some performances case of interest measured by profilers, in addition to those from timers. In designing and implementation phase, this

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<sup>3</sup>And so measure each single memory copy H2D, Kernel execution and memory copy D2H.

allowed us, not only to observe the correctness of some measurements, but also to check some special cases and their relative technical details and performance analysis.

### 5.1.2 Tests setup

First we must point out that the target machines where we have run tests on, weren't available in exclusive access. So, all collected time measures, could be affected by a perturbation introduced from other processes.

Anyway, during the tests execution we set an automatic monitor of the resources occupancy of target machines. In particular, we made log files where it was printed some informations from `top` and `nvidia-smi` commands<sup>4</sup>.

Once we determined the time measure criterion, we had to decide what behaviors we wanted to observe from our code.

Note that for each type of input dataset, we run multiple times (more precisely 5 times) the executable so that, for a certain input setup, we can collect several time measures. This allows us to delete some *outliers* completion times, as they may distort the result, and then we take the mean value among the remaining measures.

Moreover, as we mentioned in Subsection 2.5.1, we implemented our tests as bash scripts. These scripts will cover the task of:

- Compiling a certain executable, exploiting the rules available in our Makefile;
- Run that executable  $N_{test} - 1$  times and then redirect the output, of the running application, to a specific `.txt` file;
- Run for the  $N_{test}^{th}$  time the executable via `nvprof`, redirecting the profiler output to a folder of `.txt` log files

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<sup>4</sup>`top` gives us informations on the machine state on host side, while `nvidia-smi` allows to monitor NVIDIA GPUs state. By state we mean the main running processes, utilization percentage, used memory etc.

In next sections we'll show, for each type of kernel, what type of tests have been made and relative results.

It's important to recall that *input stream length shouldn't be known a priori*, but in tests we'll see that we have to give an input limit. This is for time measuring purpose only, because we need to have a knowledge on what and how much data we are measuring.

### 5.1.3 Speedup

An important metric related to performance and parallelism is **speedup**, it's a derived measure from completion time measures. Speedup compares the latency for solving a certain computational problem with its *best known* sequential implementation (on the target architecture available), versus solving the same problem on P hardware units, generally called *workers*, as below

$$speedup = S_P = \frac{T_{seq}}{T_P}$$

where  $T_{seq}$  is the sequential execution time and  $T_P$  is the parallel completion time.

An algorithm that runs P times faster on P processors is said to exhibit **linear speedup**. It is rare in practice, since there is extra work, involved in distributing work to processors and coordinating them. This extra work clearly introduces extra time, also known as **overhead**.

While **sublinear speedup** is the norm[12], there may be exceptions, i.e. occasional programs exhibiting **superlinear speedup**<sup>5</sup>.

**Amdahl's Law** gives an important limit on speedup: it considers speedup as P varies and the problem size remains fixed.

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<sup>5</sup>In general, some causes of superlinear speedup may be: restructuring a program for parallel execution can cause it to use memory better (cache in CPU implementations), even with a single worker; or the parallel algorithm may be able to avoid work that its serialization would be forced to do.

Amdahl identified in the execution time  $T_{seq}$  of a program, two categories: time spent doing *non-parallelizable serial work* and time spent doing *parallelizable work*. Call these  $T_{ser}$  and  $T_{par}$ , respectively.

Given  $P$  workers available to do the parallelizable work, the times for sequential and parallel execution are:

$$\begin{aligned} T_{seq} &= T_{ser} + T_{par} \\ T_P &\geq T_{ser} + \frac{T_{par}}{P} \end{aligned}$$

The bound on  $T_P$  assumes no superlinear speedup, and is an exact equality only if the parallelizable work can be perfectly parallelized.

Plugging these relations into the definition of speedup, we gave before, yields **Amdahl's Law**:

$$S_P \leq \frac{T_{ser} + T_{par}}{T_{ser} + T_{par}/P}$$

Let  $f$  be the non-parallelizable serial fraction of the total work. Then the following equalities hold:

$$\begin{aligned} T_{ser} &= f \cdot T_{seq} \\ T_{par} &= (1 - f) \cdot T_{seq} \end{aligned}$$

Substituting these into speedup equation:

$$S_P \leq \frac{1}{f + (1-f)/P} \Rightarrow S_\infty \leq \frac{1}{f}$$

Speedup is *limited by the fraction of the work that is not parallelizable*, even using an infinite number of processors[12].

### 5.1.4 Results: gathering and evaluation

From all `.txt` files, containing time measures for all the execution that have been run from tests, we have to manipulate results and do some calculations.

In particular, implemented Python scripts <sup>6</sup> provides a tool to:

- First of all, output all necessary `.csv` containing all averages, of the times got by the multiple runs for a certain input <sup>7</sup>;
- Then from all those average Completion times, in `.csv` format, another script computes all **Speedups**;
- Finally, the same script that computes speedups, outputs plots on most significant measures and results.

It's important to point out what kind of speedups will be computed, so that in next sections we can presents numeric and graphic results.

Remembering what we introduced in Section 5.1.3, the speedup, in brief, is the ratio of the time spent in sequential version to the time spent on parallel version, having P workers:

$$speedup = S_P = \frac{T_{seq}}{T_P}$$

Here we have to define what those times correspond in our implementation:

- $T_{seq}$ , the sequential version, is the case in which CUDA Streams aren't used<sup>8</sup>. In a sense, this corresponds to serialize all data transfers and kernel execution as

$$H2D_0, Ker_0, D2H_0, H2D_1, Ker_1, D2H_1, \dots$$

So, even if we have a stream of items as input, this means sending only a small task per time to the device, thus using only a small amount of computational resources at time;

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<sup>6</sup>See Chapter 2.

<sup>7</sup>Remember that outliers values are rejected, and mean values are computed on remaining values.

<sup>8</sup>More precisely only default stream is used, instead non-default streams aren't.



- $T_P$ , the parallel version, is the case in which CUDA Streams are used, where  $P$  will be the number of non-default streams spawned.

In the particular scenario of a Farm for GPU, the number of workers has a more complex meaning. Those workers, more precisely, corresponds to how many Streaming Multiprocessors we're going to use in the GPU, i.e. the target number of SMs we want to make busy at computation peak time.

So the speedup, obtained from those two implementations, should also give us an indicator on how many SMs will be effectively used. In other words, here we can see a CUDA stream as a sort of channel in which we put tasks and we want to see if those channels will successfully overlap, hiding data transfer from/to device, executing multiple kernels at the same time and other possible latencies. So, the computed speedups are:

1.  $S_3 = \frac{T_{seq}}{T_3}$  as we mentioned before, this is a base case for CUDA streams usage;
2.  $S_{\#SM} = \frac{T_{seq}}{T_{\#SM}}$  this is the special case that aims to show the Farm parallel pattern fitting in GPUs architecture.

The way we defined speedup lead us to ask what is the best we can achieve and this is where Amdahl's law can be applied.

First we should define what completion times, upon which we base the analysis, are. We start by focusing on the total completion time:

$$T_{Tot} = n \cdot (T_{InStream} + T_{H2D} + T_{Kernel} + T_{D2H} + T_{OutStream})$$

Below we'll explain this formula's components:

1.  $n$  is the total number of tasks appearing on the input stream<sup>9</sup>;

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<sup>9</sup>Always consider this as an indefinitely big number.

2.  $T_{Tot}$  is the overall time it takes to compute  $n$  elements from an input stream, i.e. latency from the first available item on the input stream, until the last result item is sent to the output stream (this can involve both host and device elapsed times);
3.  $T_{Instream}$  this is the time it takes to get a –data parallel small –task from input stream (host measure);
4.  $T_{H2D}$  is the time spent in transferring a task from host to device (device time);
5.  $T_{Kernel}$  is the time needed by the GPU to compute a certain kernel on the input item (device time);
6.  $T_{D2H}$  is the time spent in transferring back result item from device to host (device time);
7.  $T_{Outstream}$  the time it takes to send results to the output stream (host measure).

Obviously the measures we performed and all next considerations will be based only on device completion times, i.e. points 4, 5 and 6 of the above list.

We can reduce our test to an important assumption: we don't know how much it is  $n$  (the real length of the input stream), that's why we focus our measures on a reasonably long portion of the input stream, e.g. take as limit an  $l$  such that ( $l \leq n$  elements will be the stream limit).

So we'll focus our attention exclusively to:

$$T_{Device} = l \cdot T_{Comp}$$

where

$$T_{Comp} = T_{H2D} + T_{Kernel} + T_{D2H}$$

In other words  $T_{Comp}$  is the time needed to: send an item of the input stream to the GPU; make computations on that item and then send back the result item to host.

As we told before, host elapsed times aren't in the study domain of this thesis. So,

what we want to parallelize is  $T_{Device}$ , in particular we want all  $l$  tasks to run in parallel. Clearly, we're also assuming that  $l > \#SM$ , i.e. our tests will run on a limited input stream, but still ensuring that we have several tasks to send for each CUDA streams, consequently more tasks per SM.

Following all these assumptions, it's normal to ask what would it be the maximum reachable speedup and to define this we exploit Amdahl law, showed in previous section. Recall that according to Amdahl, serial code can be identified in two categories: serial fraction and parallelizable fraction.

Since we're only focusing on  $T_{Device}$  and, since we wish to parallelize all of it, ideally all operations included in this completion time may be considered parallelizable. This leads trivially to:  $f = 0$  and  $(1 - f) = 1$ .

Merging this with the Amdahl's upper bound we obtain:

$$S_P \leq \frac{1}{f + (1-f)/P} = \frac{1}{1/P} = P$$

And since we have a limited amount of resources, given by the Streaming Multi-processors number, we can conclude that the maximum speedup we can achieve is:  $Sp_{\#SM} \leq \#SM$ . This formally proves our expectations.

In the next sections there will be reported all completion times and speedups, that are mostly representative. For each type of kernel, that was implemented, we'll show inputs, tests and results (with some graphics and plots).

### 5.1.5 Computation-bound and memory-bound

In Chapter 3 we presented a performance model for Farm parallel pattern, coupled with speedup and Amdahl's law. These types of performance model are the most used to predict (approximately) performances.

Anyway, in this study we're interested also in another performance model: ***Roofline Model***.

Often off-chip memory bandwidth is a constraining resource in system performance.

Hence, we want a model that relates processor performance to off-chip memory traffic. To this aim we define some important concepts:

- **work**  $W$  denotes the number of operations performed by a given kernel or application. This metric may refer to any type of operation according to the study necessity (e.g. the number of integer operations, the number of floating point operations (FLOPs), etc.). In the majority of the cases however,  $W$  is expressed as FLOPs.

Note that the work is a property of the given kernel or application and thus only partially depend on the platform characteristics.

- **memory traffic**  $Q$  denotes the number of bytes of memory transfers incurred during the execution of the kernel or application. In contrast to  $W$ ,  $Q$  is heavily dependent on the properties of the target machine.
- **arithmetic intensity**  $I$ , also referred to as operational intensity, is the ratio of the work  $W$  to the memory traffic  $Q$ :

$$I = \frac{W}{Q}$$

and denotes the number of operations per byte of memory traffic. When the work  $W$  is expressed as FLOPs, the resulting arithmetic intensity  $I$  will be the ratio of floating point operations to total data movement (FLOPs/byte)[23].

In particular, the term “operational intensity” generally means operations per byte of DRAM traffic. That is, we measure traffic between the caches and memory rather than between the processor and the caches. Thus, operational intensity predicts the DRAM bandwidth needed by a kernel on a particular computer[20, 21].

The *naive Roofline* is obtained by applying simple bound and bottleneck analysis. In this formulation of the Roofline model, there are only two parameters, the *peak*

*performance* and the *peak bandwidth* of the specific architecture, and one variable, the *arithmetic intensity*.

The peak performance, in general expressed as GFLOPS<sup>10</sup>, usually it's derived from architectural manuals, while the peak bandwidth, that references to peak DRAM bandwidth to be specific, is instead obtained via benchmarking.

The resulting plot<sup>11</sup> is then derived by the following formula:

$$P = \min \begin{cases} \pi \\ \beta \times I \end{cases}$$

where  $P$  is the attainable performance,  $\pi$  is the peak performance,  $\beta$  is the peak bandwidth and  $I$  is the arithmetic intensity.

The point at which the performance saturates at the peak performance level  $\pi$  (that is where the diagonal and horizontal roof meet), is defined as ridge point.

The ridge point offers insight on the machine's overall performance, by providing the minimum arithmetic intensity required to achieve peak performance.

A given kernel or application is then characterized by a point given by its arithmetic intensity  $I$  (on the x-axis). The attainable performance  $P$  is then computed by drawing a vertical line that hits the Roofline curve.

So, the kernel or application is said to be **memory-bound** if  $I \leq \pi/\beta$ .

Conversely, if  $I \geq \pi/\beta$ , the computation is said to be **compute-bound**[20, 22].

We'll give an example on our kernel applications, mainly to prove whether they're memory or compute bound.

To simplify this example, consider the amount of computations and memory operations needed to produce a single floating point number as output. Moreover, we consider memory traffic as simple count of loads/stores, instead of single bytes (as we're consid-

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<sup>10</sup>Floating point operations per second are called FLOPS, it is a measure of computer performance, useful when we've floating-point calculations. GFLOPS, are simply giga FLOPS.

<sup>11</sup>in general with both axes in logarithmic scale

ering an example on individual floats, that is always 4 bytes).

First, consider the Simple-computational kernel and assume to set it to perform  $M$  iterations (e.g. take 10 000<sup>12</sup>) to produce a single float as output. At each iteration we perform a single-precision cosine (`cosf`) that is internally computed in approximately 10 FLOP<sup>13</sup>. Furthermore, to get a single output, we perform 2 memory operations, one load and one store between registers and global memory.

So we can get as computational intensity:  $I \approx \frac{10}{2} \cdot M = 5 \cdot M$ , where  $M$  is the number of times we repeat arithmetic operations. For example for  $M = 10\,000$  we get  $I = 50\,000$ .

Now we make an example on matrix-multiplication kernel. Call  $k$  the number of sums of multiplications that the kernel performs to output each floating-point number. So, for each output, the kernel performs  $2 \cdot k$  FLOPs<sup>14</sup> (one sum, one multiplication for each kernel iteration). Furthermore, we have  $2 \cdot k$  loads (the couples of elements from input matrices that will be multiplied) and 1 store, between global memory and registers (one element of the result matrix).

So computational intensity will be given by:  $I = \frac{2 \cdot k}{(2 \cdot k) + 1}$ .

Summarizing the results of the above examples, we can observe that in first kernel, computation intensity simply will proportionally grow, as the number of iterations will increase.

While the second kernel shows a different behavior, i.e. computation intensity doesn't really depend on the number of iterations of the kernel, since it will give always a result  $< 1$  (we have more memory than compute operations).

According to the Naive Roofline model we can determine if an application is memory

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<sup>12</sup>We'll see in next sections that this is the minimum amount of computations that kernel had to perform in our tests.

<sup>13</sup>Here we can see a discussion on amount of FLOP per mathematical operation.

<sup>14</sup>Here intended as FLoating POint operationS, and not FLOPs per second.

or compute bound, by check respectively if  $I \leq \pi/\beta$  or  $I \geq \pi/\beta$ . From architecture manuals[17] and specifications<sup>15</sup>, we obtained  $\frac{\pi}{\beta} \approx \frac{10600 \text{ GFLOP/s}}{732 \text{ GB/s}} = 14,48 \text{ FLOP/B}$  for the P100 device and  $\frac{\pi}{\beta} \approx \frac{6840 \text{ GFLOP/s}}{288 \text{ GB/s}} = 23,75 \text{ FLOP/B}$  for the M40 GPU.

Merging it with the above kernel examples of arithmetic intensity, we get<sup>16</sup>:

- simple-computational kernel  $I \approx \frac{10}{4(\text{bytes})} \cdot M = 2.5 \cdot M$
- matrix multiplication  $I = \frac{8 \cdot k}{(12 \cdot k) + 1} \approx 1$

So we can conclude that for matrix multiplication computation intensity will be  $I \leq \pi/\beta$  for both target machines.

While in simple-computational kernel, we have a computation bound behavior for a small value of  $M$  (e.g. for  $M=10$  the kernel is compute-bound for both target machines, we recall that the minimum amount we used in tests is  $M = 10\,000$ ).

## 5.2 Simple-computation kernel

For this computation-bound kernel, for each type of input dataset, we identified three different values of kernel iterations number, call it  $M$ , it takes the following values: 10000, 400000, 800000.

These values identify how many times the kernel will have to repeat a certain mathematical operation (in our case the Cosine). This is what makes the computation load variable.

Another important parameter is the Block size that we set to  $BLOCK = (1024, 1, 1)$ . We recall that 1024 is the maximum we can give to  $x$  and  $y$  block dimension, this holds for both of the GPUs we used to run tests, ie **P100** and **M40**.

The choice for 1024 was made according to CUDA Occupancy APIs, that suggested this as best block size for the considered application. In general, but it's not a strict

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<sup>15</sup>We checked the CUDA Device Query too.

<sup>16</sup>Now we'll consider memory operations in terms of bytes, instead of load/stores number, to uniform them with the model.

<b>Tesla P100</b>	<b>Tesla M40</b>
57 344	24 576
114 688	49 152
229 376	98 304
458 752	196 608
917 504	393 216
1 835 008	786 432

Table 5.1: Input dataset for Simple-Computation kernel, these are the input stream length for both devices.

rule, computation-bound kernels perform at their best on higher block size, because this should allow us to use the maximum number of threads possible and, thus, to use as much computational resources as possible.

All types of tests performed on Simple-computational kernel are:

### 1. **Classic data parallel approach**

Here we launch the execution of our simple-computation kernel, as it would be classically used: as fully data parallel application.

This means that we have a single big data structure, that we'll send entirely to the GPU for data parallel computations.

Clearly this collection of data will execute the same kernel and will compute the same amount of work w.r.t. the Stream parallel version.

So, essentially, we're trying to model a problem from the perspectives of two different parallel pattern. It's important to point out that stream parallel will model an input/output stream of small data parallel tasks, while data parallel is modeling a fixed size unique big data structure<sup>17</sup>.

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<sup>17</sup>E.g. Simple-computation's data parallel version could be used to make data parallel computations over



In Table 5.1 we show length that was used. In data parallel those values are the size of the data structure upon which data parallel computations will be performed.

Instead, in stream parallel, the same amount of computations will be performed, but it will be done in different input streams items, given by small data parallel tasks.

Note that we should not wrongly think that data parallel version is the equivalent of bringing together tasks from input stream of stream parallel problem<sup>18</sup>.

2. **Streaming parallel with smaller tasks** Here we're facing the Farm parallel pattern for GPU, but with smaller tasks size. As we mentioned in Chapter 3, we're trying to get maximal occupancy, especially in a computational-bound kernel. So, given that the goal of our code is that each called kernel could fill fully or partially a SM, we had to take into account of:

- How big should be each task for each kernel execution;
- Consequently, how many thread blocks our kernel will issue.

These choices followed from our devices features, though the two GPUs are located in different Compute Capabilities (P100 is c.c. 6.0, M40 is c.c. 5.2) they have the same limits for

- Resident **threads** per SM = 2048 (equivalent to 64 resident warps per SM);
- Resident **thread blocks** per SM = 32.

The second limit means that we can have at most 32 thread blocks active, and so running, on a certain Streaming Multiprocessor.

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a big array of floats.

<sup>18</sup>Here, as in all data parallel versions we implemented, we don't make use of CUDA Streams, they'd be useless since we're launching a single kernel on a single huge data structure

The first limit, instead, is our main goal here. Having at most 2048 active threads in a SM and having configuration of `blocks = (1024, 1, 1)`, we will have at most two resident blocks in a SM.

The execution configuration for smaller buffers is such that we want to have 1024 buffers and kernel configuration such as `<<<1, 1024>>>`. So here each launched kernel will have one block containing 1024 threads and this theoretically should correspond to half the occupancy of a SM.

Clearly the code will send a lot of tasks to device, i.e. enough to hopefully fill all SMs. We recall that the number of chunks will be limited according to values in Table 5.1.

All of the above mentioned configurations will be tested for the following CUDA streams cases:

- **Zero** CUDA Streams. This is the scenario where we use any non-default CUDA stream, so we'll have serial and synchronous data transfers. Kernel are still an asynchronous call, but immediately after we want to have data back from device and this means have a `cudaMemcpy`, ie a synchronous call (w.r.t. the host);
- **Three** CUDA Streams. Here we'll use 3 non-default streams, because we want to observe the behavior of our code in a sort of base case. In general, using three stream is the classic configuration for devices with two copy engines. This means it's the minimum to expect an overlap such as a kernel and at most two simultaneous data transfers;
- **$N_{SM}$**  CUDA Streams, with  $N_{SM} = \#Streaming\ Multiprocessors$ . This is the special case because, in general, applications don't use such a high number of CUDA Streams. But in our case it's necessary to try to achieve the expected speedup with respect to the version without non-default streams (we'll sometimes refer to as "zero version").

Clearly, at a certain time say  $t_i$ , we can have at most two data transfer but there's no limit on kernel calls, clearly they will be effectively executed as long as there are available resources on the device. So this is the key point why all of kernel launches, at peak CUDA stream filling, should be spread in SMs, as soon as requested resources will be available.

3. **Streaming parallel with bigger tasks** This tests setup has similar premises to the one for smaller (data parallel) tasks, clearly the only thing is changing is the task size, tha will be set to 2048.

Thus, having always `blockSize=(1024, 1, 1)`, the code will set `gridSize=(2, 1, 1)`, this is because we'll have two blocks, each covering calculations on one half of the task. This can sound as having better performances, with respect to smaller tasks, but, as we said before, it's not a strict rule to have better performances on maximum occupancy. We'll see from results that instead this approach behaves worse than smaller tasks.

Clearly we repeated the above mentioned amount of CUDA streams, so for this configuration we executed code using: **Zero**, **Three** and **N<sub>SM</sub>** CUDA Streams.

## 5.2.1 Results

All the above tests on Simple-Computation Kernel give us the measures of device times, on which most of observations will rely on.

We can group the measures into two parts:

### Smaller buffers

All collected elapsed times for 1024-sized tasks are reported in Table 5.2, for the zero-streams version, and Table 5.3, for the SM-streams version.

From measures in Table 5.2(zero-streams) we can see that Completion Time, fixed an

	<b>Tesla P100 (zero stream)</b>		<b>Tesla M40 (zero stream)</b>	
M iterations	Event Times	N elements	Event Times	N elements
10000	4622.86	57344	693.747	24576
400000	181465.333		27453.933	
800000	361199.666		54888.933	
10000	9294.18	114688	1382.363	49152
400000	281507		54901.6333	
800000	407750.666		109783.666	
10000	10217.933	229376	2765.323	98304
400000	407779.666		109799.333	
800000	815513.666		219553	
10000	20433.633	458752	5528.96	196608
400000	815561		219589	
800000	1631013.333		439097.666	
10000	40865.4	917504	11058.433	393216
400000	1631096.666		439192.333	
800000	3261986.666		878195	
10000	81731.733	1835008	22112.6	786432
400000	3262250		878433	
800000	6617950		1756373.333	

Table 5.2: Device completion times for Simple-computation kernel, without using CUDA Streams, results are reported for both machines (P100 and M40).

	<b>Tesla P100 (56 Streams)</b>		<b>Tesla M40 (24 Streams)</b>	
M iterations	Event Times	N elements	Event Times	N elements
10000	104.772	57344	30.6074	24576
400000	3968.913		1178.72	24576
800000	7818.843		2355.413	24576
10000	205.729	114688	60.208	49152
400000	7828.193		2358.656	49152
800000	15691.833		4714.36	49152
10000	407.712	229376	119.3446	98304
400000	15687.966		4715.123	98304
800000	31396		9425.92	98304
10000	803.223	458752	238.249	196608
400000	31422.033		9429.89	196608
800000	62818.7		18853.966	196608
10000	1619.586	917504	475.590	393216
400000	62793.4		18856.8	393216
800000	125575		37705.666	393216
10000	3229.063	1835008	949.497	786432
400000	125547.666		37711.9	786432
800000	251503		75445.266	786432

Table 5.3: Device completion times for Simple-computation kernel, using as many CUDA Streams as SM number, results are reported for both machines (P100 and M40).

input stream length, increases proportionally with iterations number (e.g. completion times for 40 000 iterations kernel compared to the one for 1000, is almost  $40\times$  bigger).

This holds on both of devices.

This is a further sign that this type of kernel is computation-bound.

Furthermore, in the input data set for tests on Simple-computational kernel, we set input stream tasks to increase in number by a factor of  $2^{19}$ .

Always looking at Table 5.2, fixed a number of iterations, we can see that even completion times grows by a factor of 2.

This confirms that, no matter how many tasks the input stream sends, no matter how many iterations the kernel does, *we'll have a completion time directly proportional to the computations amount* performed by the simple-computation kernel.

Now turning on Table 5.3, we can see the same behavior just presented for zero-streams version. In SM-streams version too we can observe that measures grows as computations amounts grows. However it's immediate to see that zero-streams and SM-streams have completely different completion times.

This leads to compute speedups, following the approach explained in Section 5.1.4. All of the speedups are shown in Table 5.4. In this table the most important columns are  $Sp(3)$   $Sp(SM)$ , those columns stands for:

$$Sp(3) = \frac{T_{seq}}{T_3} \text{ and } Sp(SM) = \frac{T_{seq}}{T_{SM}}.$$

To have an overall view on speedup we also present plots for both P100, in Figure 5.1, and M40, in Figure 5.2.

The two plots show the speedups only for a part of the real input dataset; in particular, each plot shows the smaller and bigger input stream length and for both it shows the smaller and bigger kernel iterations number.

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<sup>19</sup>This means we're increasing the pressure of tasks that each SM will have to compute in total

	Tesla P100 (56 Streams)			Tesla M40 (24 Streams)		
M iterations	N elements	Sp(3)	Sp(56)	N elements	Sp(3)	Sp(24)
10000	57344	5.337	44.122	24576	2.976	22.665
400000		5.246	45.721		2.963	23.291
800000		5.221	46.196		2.963	23.303
10000	114688	5.366	45.176	49152	2.967	22.959
400000		4.069	35.960		2.962	23.276
800000		2.947	25.984		2.963	23.287
10000	229376	2.988	25.061	98304	2.970	23.170
400000		2.986	25.993		2.963	23.286
800000		2.986	25.975		2.962	23.292
10000	458752	2.988	25.439	196608	2.967	23.206
400000		2.986	25.955		2.963	23.286
800000		1.940	25.963		2.963	23.289
10000	917504	1.635	25.231	393216	2.967	23.251
400000		1.689	25.975		2.963	23.290
800000		2.861	25.976		2.963	23.290
10000	1835008	2.998	25.311	786432	2.968	23.288
400000		2.996	25.984		2.964	23.293
800000		2.654	26.313		2.963	23.280

Table 5.4: Here are showed speedups for all data sets of simple-computation kernel. Results are reported for both devices.

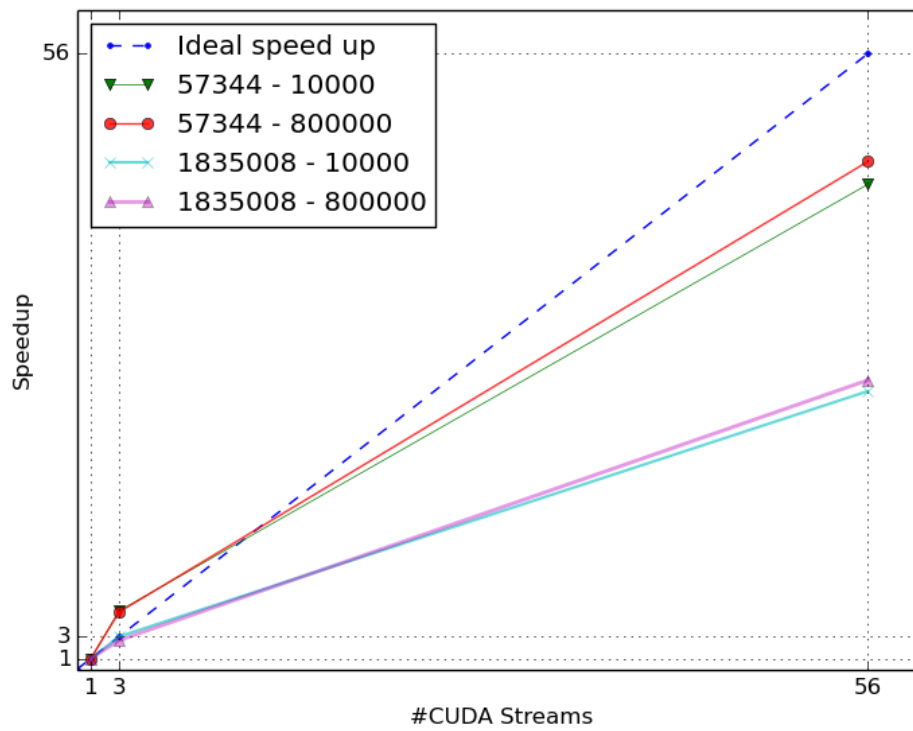


Figure 5.1: Speedup for 3 and 56 CUDA streams on P100 device.

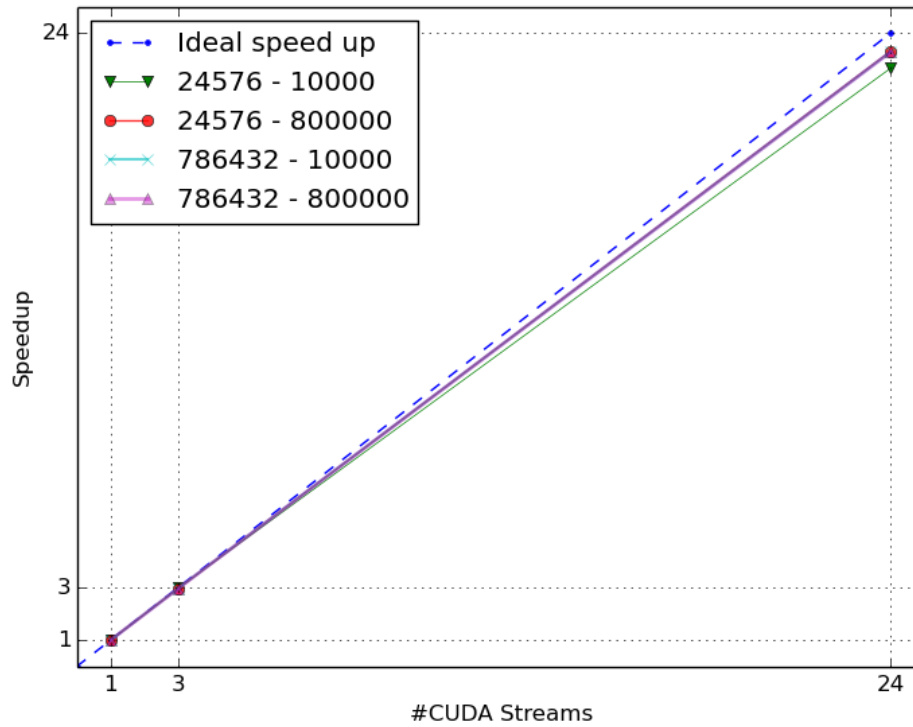


Figure 5.2: Speedup for 3 and 56 CUDA streams on M40 device.



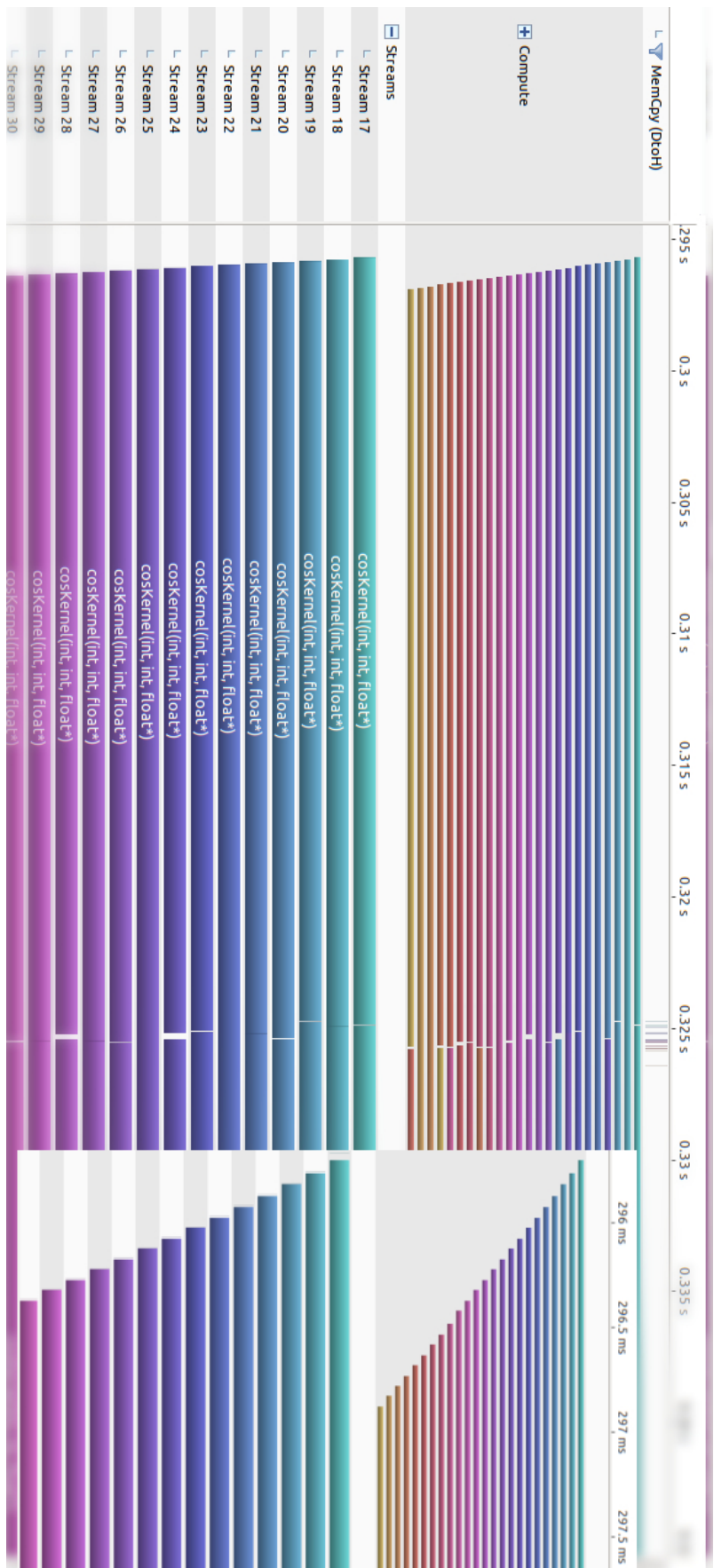


Figure 5.3: Profiling for an example execution: limit for input stream 786432, kernel iterations 10 000, 24 CUDA streams, on M40 device.

From the two plots we can clearly see how performances increase proportionally with the number of CUDA streams<sup>20</sup>.

We can have a further proof of the achieved speedup, by running the **NVIDIA Visual Profiler**, we can see a portion of timeline representation in Figure 5.3. It's evident that we have a good overlapping between CUDA Streams, on the right side of the figure we can see a zoom in to the "*stabilization phase*", i.e. the time interval it takes for the program to fill buffers and send them out to the device, until we reach a peak work rate and that "stairs" behavior vanishes.

However the profiler analysis reported some issues in our code, for example the profiling showed a too much low grid/block size and a poor memory copy overlapping (having 2 data transfers at the same time). But all these issues are really dependent on the "extreme" use case we're approaching to achieve stream parallel computations and, furthermore, they don't have a bad impact in this particular application.

A good signal from the profiler is that the average usage of SMs, for a certain kernel call, is equal to one almost fully used SM. And this is exactly what we wanted to achieve, so this allows CUDA streams to spread kernel calls on all SMs, as soon as we have the maximum work rate.

## Bigger tasks

As expected, we get about half the ideal speedup.

This is because chunks of 2048 should take up the maximum possible for active threads per SM. This means having, for each kernel call, a grid containing 2 blocks, each of size 1024.

So it seems that performances, in this kernel, are related to grid size. Recalling the smaller tasks, the profiler showed an almost full occupancy of one SM at <<<1, 1024>>>

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<sup>20</sup>More precisely this gain is bounded to the number of Streaming Multiprocessors the GPU exploits.

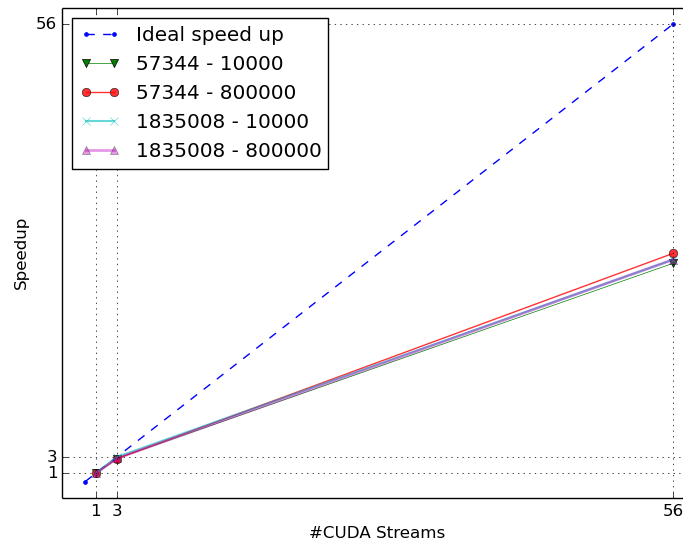


Figure 5.4: Sublinear speedup in bigger buffers execution, the performances drop by a factor of 2.

yet. So it makes sense to deduce that,  $\langle\langle\langle 2, 1024 \rangle\rangle\rangle$  is a kernel configuration such that it occupies two SMs simultaneously.

This is the same of having about the half of available resources. We show a plot about this result, tested on P100, in Figure 5.4

## 5.3 Matrix Multiplication

With Matrix multiplication we're facing a memory-bound kernel, so we had to make some slightly different tests with respect to the ones in simple-computation kernel.

Note that, even if the Farm logic is the same as the previous application, there are some details to redefine.

First of all, before we were dealing with an input stream of small data parallel tasks built by floats, here those small data parallel tasks are built by small *matrices*. In particular as soon as we have two available input matrices (say  $A$  and  $B$ ), they're sent out to device to apply the matrix multiplication kernel.

Finally we get back to host with the result matrix (say  $C$ ), that will be one of the output stream components.

For simplicity, we're measuring for square matrices case, even if code was implemented

Tesla P100		Tesla M40	
Mat. Order	In stream limit	Mat. Order	In stream limit
128	225	64	100
256	441	128	196
512	900	256	400
1024	1764	512	784
2048		1024	
Data Parallel		Data Parallel	
1920		1280	
2816		1792	
3840		2560	
5632		3584	
7680		5120	
11264		7168	

Table 5.5: Input dataset for Matrix Multiplication kernel. Above Stream parallel configuration, below Data Parallel correspondent.

for non-square case too.

Another assumption is that, for each Matrix Multiplication test, the kernel execution configuration was set up as follows:

assume  $N$  to be the matrices order, then  $BLOCK = 32$  and  $GRID = (N/BLOCK) + BLOCK - 1$ , so we have:

```
blockSize = (BLOCK, BLOCK, 1)
gridSize=(GRID, GRID, 1)
```

We performed the following executions:

1. **Classic data parallel approach** The fully data parallel version here, is totally

analogous to the one explained for Simple-computational kernel.

Here we have again big data parallel data structures (in this case a big matrices), upon which we're performing data parallel computations (i.e. matrix multiplication between two big matrices). In Table 5.5, in the lower portion, are showed the matrices orders we used for data parallel version of matrix multiplication.

Again we shouldn't confuse a totally data parallel matrix multiplication on single and big data structures, with a stream parallel version having as items small data parallel tasks.

So data parallel must not be confused as a grouping of tasks by stream parallel version. Instead, the values presented in Table 5.5 for Data Parallel and Stream parallel should be considered as simple indicators to compare the two versions in a situation in which they're computing the same amount of overall work (and clearly they're modeling the same problem but on totally different types of input).

2. **Streaming parallel** As we mentioned before, we have to put a limit on the stream length for both streams<sup>21</sup>.

In the upper portion of Table 5.5 are reported all input stream limits and, for each of them, we test different input tasks sizes (matrices order).

Note that input stream limits from Table 5.5 means how many tasks will be arriving from each of the two input streams (and, so, how many items should be sent to the output stream). For every combination given by the input parameters, we'll test for different numbers of CUDA streams: **Zero**, **Three** and **N<sub>SM</sub>** CUDA Streams (with  $N_{SM} = \#Streaming\ Multiprocessors$ ). The above test on different numbers of non-default streams, is implemented in a totally analogous way to the one for Simple-computation Kernel.

---

<sup>21</sup>As in matrix multiplication we have to multiply 2 matrices per time obtaining one as result, we suppose to have two input streams, giving us two tasks per worker. Workers in turn will output one item per time on the output stream.

### 5.3.1 Results

All the above tests, on Matrix Multiplication Kernel, give us the measures of device times.

Below we'll see that completion times and performance will notably be different, with respect to the previous computation-bound application.

All collected elapsed times are reported in Table 5.6, for the zero-streams version, in Table 5.7, for the three-streams one, and Table 5.8, for the SM-streams version.

From these tables we can highlight some behaviors:

- input streams of tasks have lengths that grow by a factor of 2 (the task number per SM doubles as execution input parameter) and it's easy to see that this makes a proportional increase in completion times, i.e. even measures grows by factors of 2;
- input task sizes again grow of  $2\times$  each, but in this case the completion times don't grow proportionally;
- for zero-streams we can see that, as tasks size grows by a factor 2, the completion time can increase from  $\approx 4\times$  to  $\approx 7\times$ ;
- for three-streams we can see that, as the task size grows by  $2\times$ , the completion time can increase from  $\approx 5\times$  to  $\approx 8\times$ ;
- finally for SM-streams we can see that, the completion time can increase from  $\approx 7\times$  to  $\approx 8\times$ .

Those evidences hold for both machines measures and they give some hints on matrix multiplication nature.

The first point tells us that: the elapsed time to send/receive to/from the device, grows linearly with the number of tasks, so this parameter would not affect particularly performances. Especially, no matter the CUDA streams amount we decide to use, the

Tesla P100 (zero Streams)			Tesla M40 (zero Streams)		
Event Times	Number of Mats	Mat. Order	Event Times	Number of Mats	Mat. Order
86.8854	225	128	17.4869	100	64
175.189	441		34.8778	196	
359.9716	900		70.9718	400	
725.5573	1764		139.3896	784	
334.0376	225	256	36.2095	100	128
672.9463	441		74.2685	196	
1435	900		147.7336	400	
2828.5366	1764		299.138	784	
1673.2133	225	512	186.3913	100	256
3325.6533	441		368.6813	196	
6611.7566	900		786.7536	400	
12919.6666	1764		1603.4933	784	
10998.7666	225	1024	1256.4	100	512
21511.1666	441		2479.4333	196	
43828.7666	900		5162.6333	400	
85853.0333	1764		9791.98	784	
80764.8666	225	2048	9075.22	100	1024
158136.3333	441		17849.5666	196	
309724.6666	900		36441.3666	400	
604324	1764		72396.8666	784	

Table 5.6: Device completion times for Mat-Mul kernel, without using CUDA Streams (zero streams), results are reported for both P100 and M40.

Tesla P100 (3 Streams)			Tesla M40 (3 Streams)		
Event Times	Number of Mats	Mat. Order	Event Times	Number of Mats	Mat. Order
25.5549	225	128	5.2046	100	64
53.2203	441		10.7422	196	
102.0575	900		23.9773	400	
193.2436	1764		47.1808	784	
148.1446	225	256	21.5294	100	128
289.6773	441		41.9395	196	
590.6776	900		74.5879	400	
1157	1764		143.3123	784	
1173.3466	225	512	129.985	100	256
2298.3866	441		254.2516	196	
4690.97	900		518.3303	400	
9205.5	1764		1016.28	784	
9371.7766	225	1024	1033.9166	100	512
18371.2666	441		2027.17	196	
37480.4	900		4136.54	400	
73258.6666	1764		8113.0933	784	
74966.4666	225	2048	8273.1066	100	1024
146156.3333	441		16194.1	196	
285788.3333	900		33041.1	400	
559469.6666	1764		64763.6666	784	

Table 5.7: Device completion times for Mat-Mul kernel, with three CUDA Streams, results are reported for both P100 and M40.



Tesla P100 (56 Streams)			Tesla M40 (24 Streams)		
Event Times	Number of Mats	Mat. Order	Event Times	Number of Mats	Mat. Order
20.8758	225	128	2.739	100	64
40.5783	441		5.0942	196	
74.6636	900		10.0277	400	
145.4766	1764		19.8252	784	
147.765	225	256	19.3538	100	128
288.5343	441		37.8560	196	
588.9643	900		65.6809	400	
1153.7333	1764		128.317	784	
1173.32	225	512	130.0533	100	256
2298.3966	441		254.281	196	
4690.9633	900		518.615	400	
9202.3333	1764		1016.55	784	
9371.0433	225	1024	1034.0666	100	512
18374.7	441		2027.2866	196	
37474.7	900		4136.7066	400	
73348.3333	1764		8110.9966	784	
74971.6666	225	2048	8262.7533	100	1024
146175.6666	441		16202.4666	196	
285955.6666	900		33059.2666	400	
559425	1764		64786.5333	784	

Table 5.8: Device completion times for Mat-Mul kernel, with as many CUDA Streams as SM number, results are reported for both P100 and M40.

increase by 2x of tasks quantity, will always give a growth of 2x in measures.

To have a visual comparison, we show plots for completion times in Figures 5.5 - 5.6, where we can have a graphical view of the completion time variation, as the tasks size grows (respectively on M40 and P100. In Figures 5.7-5.8, instead, we have similar plots, but for the time changing as the number of tasks increments.

The other points, emerging from completion times behavior, tell us that, as task size increases, we'll get worser and worser performances. Clearly this doesn't depend on CPU/GPU data transfers overhead, otherwise we'd have the same behavior when the number of tasks grows.

So, the cause must reside on what happens inside the GPU. In reality, the classic matrix multiplication is a well known problem in GPUs paradigm and the classical implementation is known as a not efficient.

This is because, the simpler implementation, at each iteration, spends more time in *global memory/registers* transfers than in effective calculations. So that's why this kind of matrix multiplication is considered memory-bound.

So, the more elements a matrices have, the more data transfers (internal to the device) the GPU will have to perform and the more active threads will stall waiting for data to be available for computations.

So we'll now focus on speedups, to see that this memory-bound behavior will be negatively reflected on GPU Farm approach. All speedups are listed in Table 5.9.

From those results, we can mainly observe that:

- $Sp(\mathcal{I})$  gives results near to the expected value, ie  $\approx 3$  for the smaller tasks sizes (128-256 for the P100, 64 for the M40);
- $Sp(SM)$  gives a really poor gain w.r.t.  $Sp(\mathcal{I})$ ;
- all speedups degrade to  $\approx 1$  as the task dimension grows.

This behavior translates in the following: when the tasks get bigger, even if CUDA

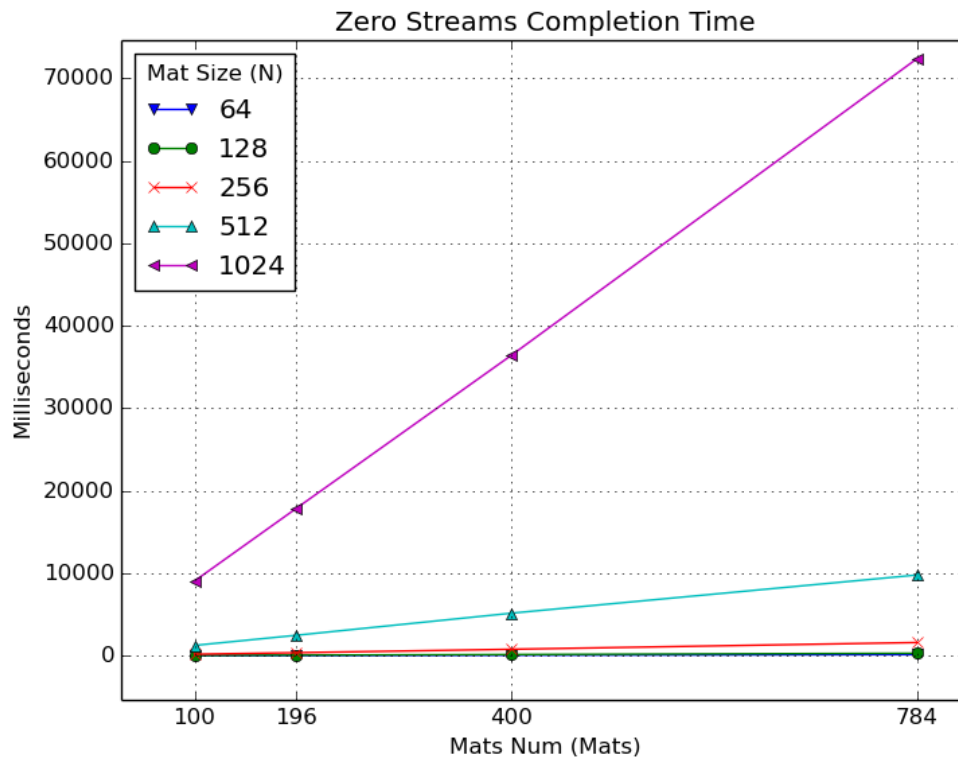


Figure 5.5: Completion Time as the matrix order changes on M40.

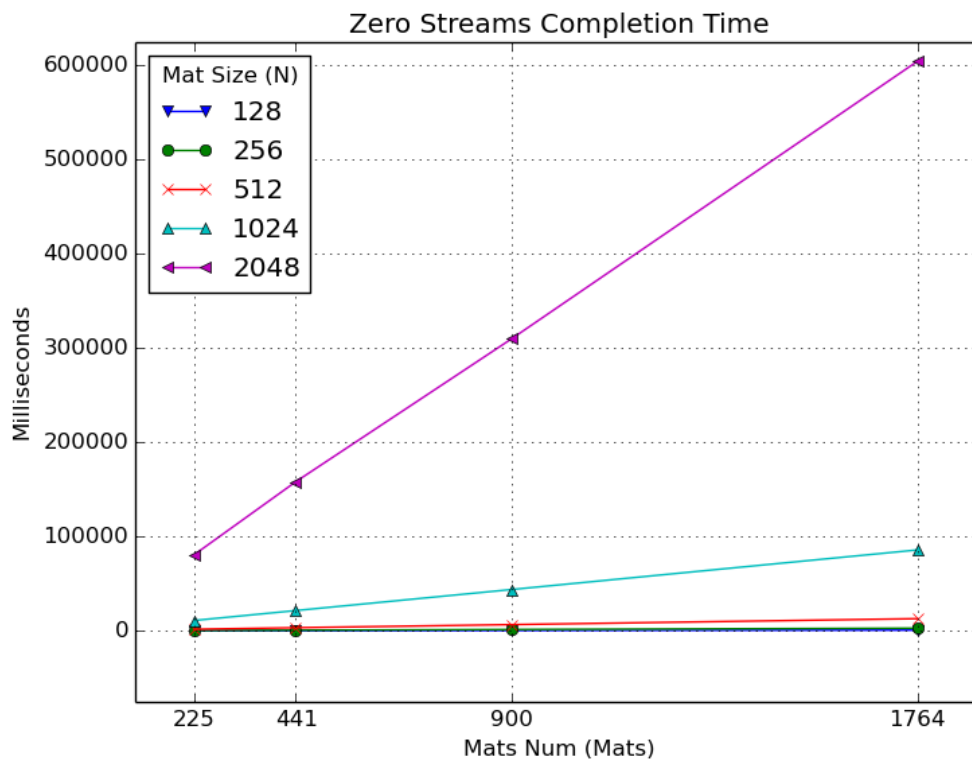


Figure 5.6: Completion Time as the matrix order changes on P100.

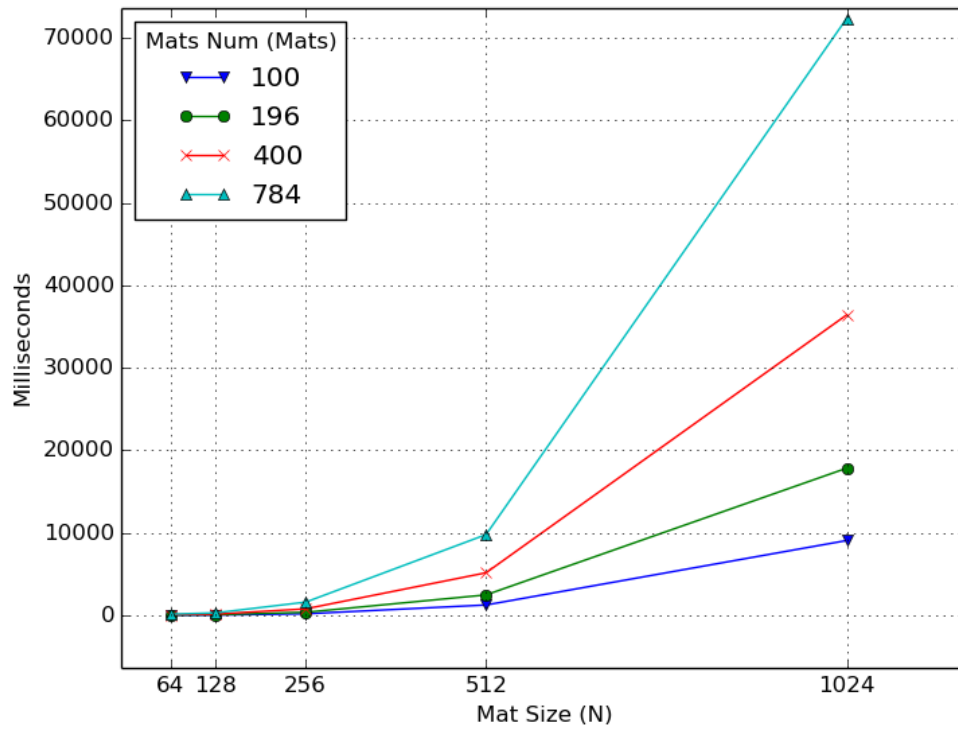


Figure 5.7: Completion Time as the number of matrices changes on M40.

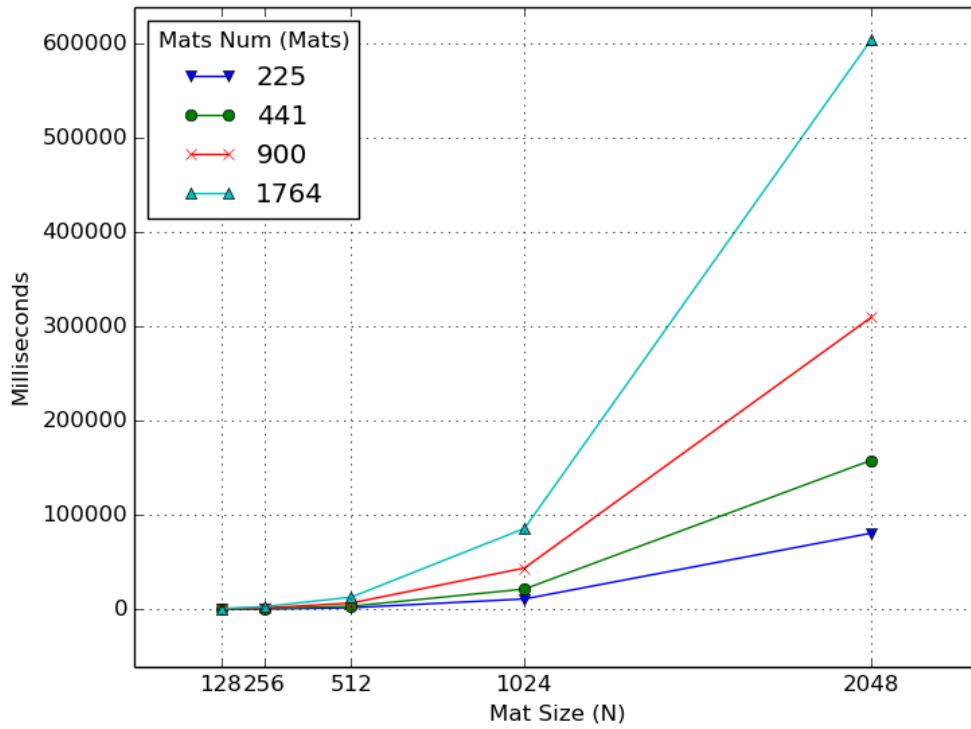
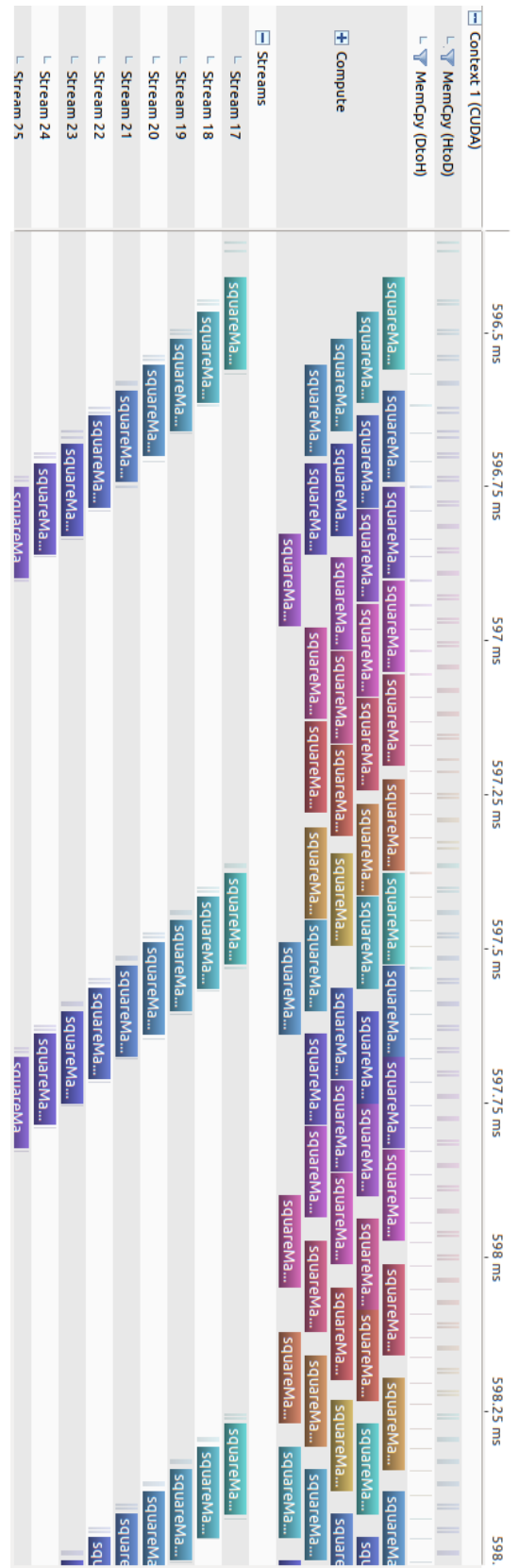


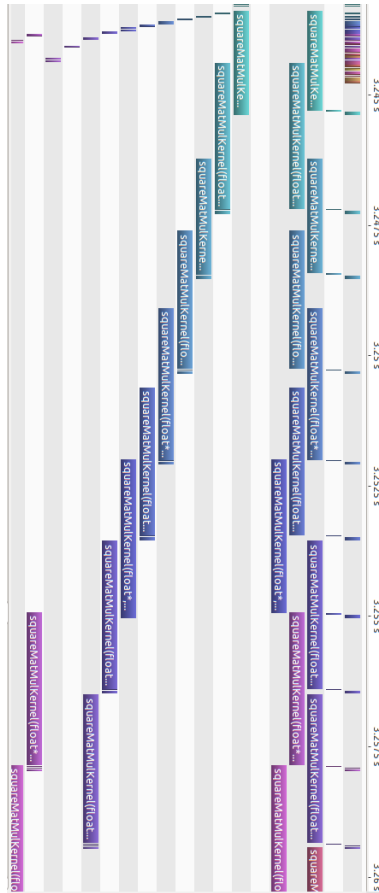
Figure 5.8: Completion Time as the number of matrices changes on P100.

Tesla P100 (56 Streams)				Tesla M40 (24 Streams)			
Mat. Number	Mat. Order	Sp(3)	Sp(56)	Mat. Number	Mat. Order	Sp(3)	Sp(24)
225	128	3.3999	4.1620	100	64	3.3598	6.3839
441		3.2917	4.3173	196		3.2467	6.8464
900		3.5271	4.8212	400		2.9599	7.0775
1764		3.7546	4.9874	784		2.9543	7.0309
225	256	2.2548	2.2606	100	128	1.6818	1.8709
441		2.3230	2.3322	196		1.7708	1.9618
900		2.4294	2.4364	400		1.9806	2.2492
1764		2.4447	2.4516	784		2.0873	2.3312
225	512	1.4260	1.4260	100	256	1.4339	1.4331
441		1.4469	1.4469	196		1.4500	1.4498
900		1.4094	1.4094	400		1.5178	1.5170
1764		1.4034	1.4039	784		1.5778	1.5773
225	1024	1.1736	1.1736	100	512	1.2151	1.2150
441		1.1709	1.1706	196		1.2231	1.2230
900		1.1693	1.1695	400		1.2480	1.2480
1764		1.1719	1.1704	784		1.2069	1.2072
225	2048	1.0773	1.0772	100	1024	1.0969	1.0983
441		1.0819	1.0818	196		1.1022	1.1016
900		1.0837	1.0831	400		1.1029	1.1023
1764		1.0801	1.0802	784		1.1178	1.1174

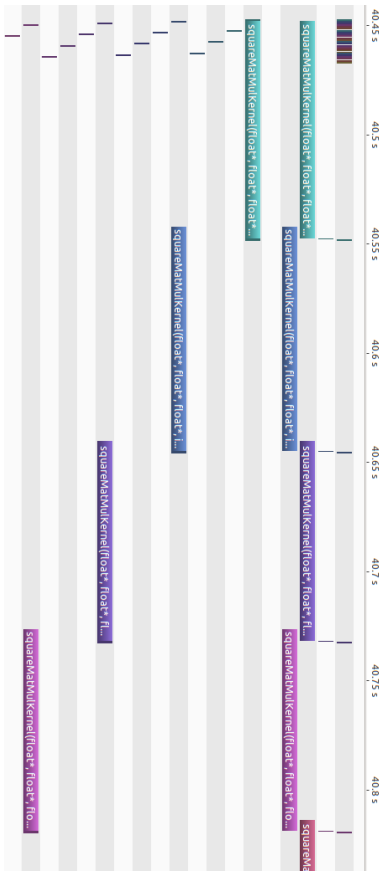
Table 5.9: Here are showed speedups for all data sets of matrix multiplication kernel. Results are reported for both devices.



(c) Matrix size 64



(b) Matrix size 256



(a) Matrix size 1024

Figure 5.9: NVIDIA Visual profiler generated *timeline* on M40, using 24 CUDA streams and running code for 784 matrices.

Streams push to have more simultaneous mat-mul, we'll have a lot of active threads (and so Multiprocessors) busy and probably waiting on gathering floats from global memory and instruction dependencies in memory operations.

This, in fact, inevitably leads to a very limited amount of gain, even when using a lot of CUDA streams. Furthermore, those results tell us that we will fit in Multiprocessor less matrix multiplication than we wish <sup>22</sup>.

Furthermore, the necessity of having multiple blocks on grid, for this specific use case, translates in a monopolization of SMs resources by a small amount of kernel calls.

Profiling the application for some key data set we can inspect the above described facts and the relative reasons.

In Figure 5.9 we can have a visual cue on what is happening during an execution of mat-mul on M40, with 24 CUDA streams, having an input stream of 784 tasks (matrices) of sizes:  $64 \times 64$  (Figure 5.9 (c)),  $256 \times 256$  (Figure 5.9 (b)),  $1024 \times 1024$  (Figure 5.9 (a)).

Analyzing the figure we can see that the amount of overlapping, between operations in different streams, is really limited in general, this just confirms what we saw from speedups.

From the above mentioned pictures, we can observe that in 64-sized case we're having a slightly better overlapping and a little more kernels running at the same time. In this case this may happens because grid and block sizes are smaller for each kernel launch, so this allow to have multiple kernels fit in SMs.

Having each thread block such that it contains  $32 \times 32$  threads, then a  $64 \times 64$  result matrix (say C) is managed by a grid of  $2 \times 2$  blocks, while a  $256 \times 256$  C is managed by a grid of  $8 \times 8$  blocks and, finally, a  $1024 \times 1024$  C is managed by a grid of  $32 \times 32$  blocks<sup>23</sup>.

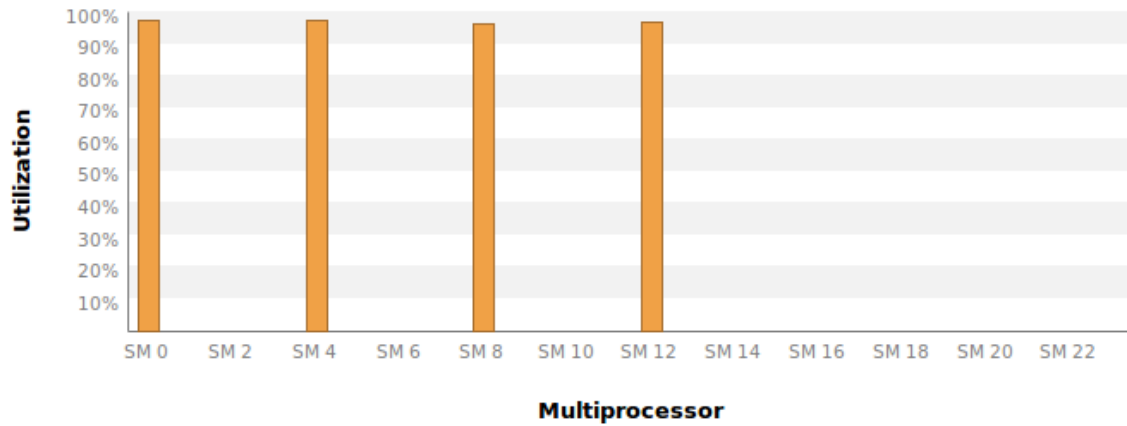
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<sup>22</sup>Clearly this strictly holds for this type of matrix multiplication we implemented.

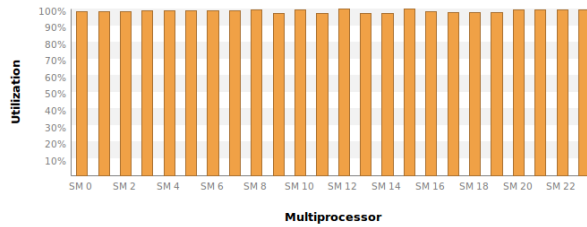
<sup>23</sup>This depends on how we implemented kernel launch, that is the classical kernel launch approach setting blocks at the maximum size possible. Remember implementations in Chapter 4 and remember the kernel

### i Multiprocessor Utilization

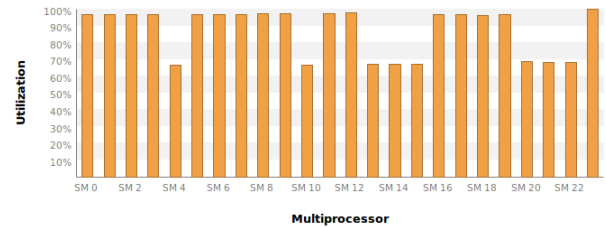
The kernel's blocks are distributed across the GPU's multiprocessors for execution. Depending on the number of blocks and the execution duration of each block some multiprocessors may be more highly utilized than others during execution of the kernel. The following chart shows the utilization of each multiprocessor during execution of the kernel.



(a) Matrix size 64



(b) Matrix size 1024



(c) Matrix size 256

Figure 5.10: NVIDIA Visual profiler generated *timeline* on M40, using 24 CUDA streams and running code for 784 matrices.

This means that, in the two latter cases, we don't even have all blocks, from a single kernel launch, fitting in all the SMs. If this may theoretically give a full occupancy of the GPU by a certain kernel, from the other side means saturate the cores without permitting other launches to fit, until the residing kernels ends or, at least, some resources are freed.

We can confirm this fact by looking at the occupancy graphs (generated from Visual Profiler), showed in Figure 5.10. In 64-sized case, we've only 4 SMs almost fully employed, while in the other cases all SMs are almost busy (this holds on average in a launch configuration explained above).



## Sample distribution

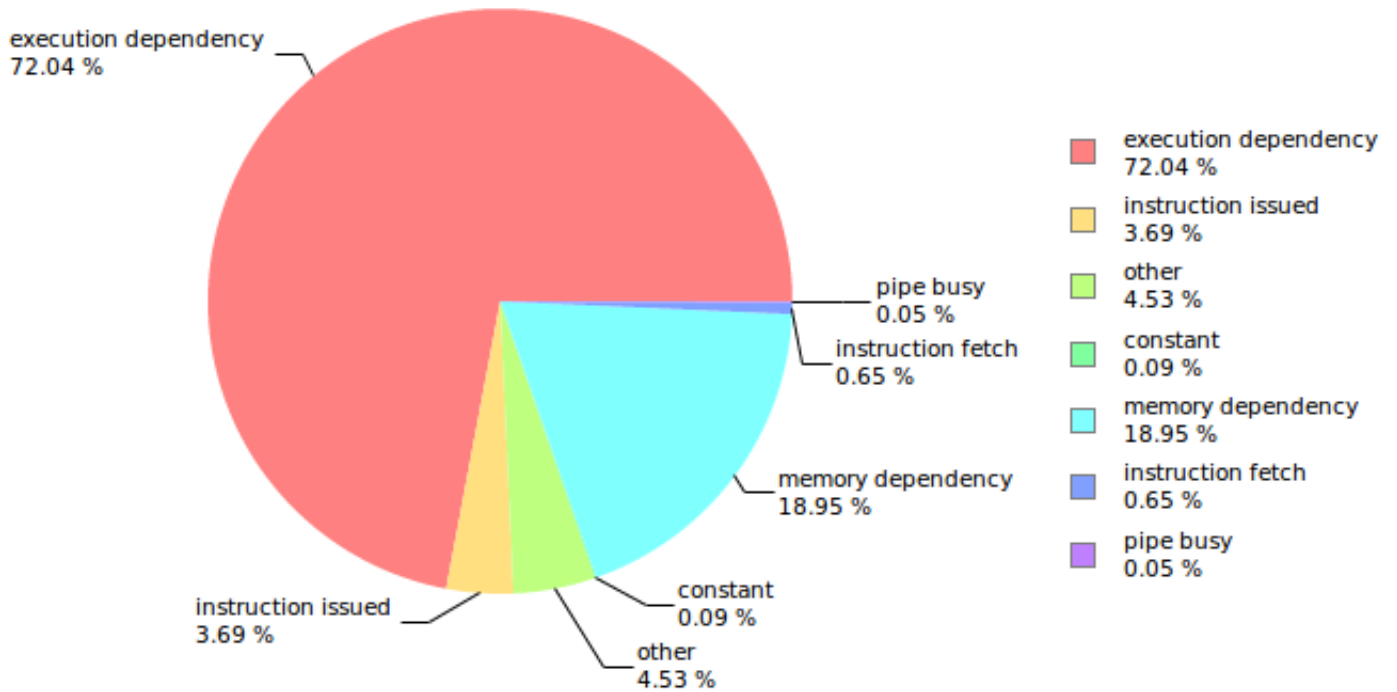


Figure 5.11: Nsight profiler execution on M40, 24 CUDA streams, 784 matrices of size 256. This graph gives the types and amounts of latencies inside mat-mul kernel execution.

single kernel launch).

The above cases exposes an example of the fact that *not always a high or full occupancy may give better performances* [8, 13], it strongly depends on the kernel nature.

On the other side as we decrease the tasks size we can't exploit enough resources. As we can see from Figure 5.9 we have a better overlapping, but it seems that kernels lasts too little, so the host can't push kernels quickly enough to fill SMs. In fact Visual Profiler suggests us that those kernels perform a really poor amount of computations, especially with respect to memory latencies, see figure Figure 5.11.

This pie chart shows that most of the time kernels is idle on:

- execution dependency<sup>24</sup>, i.e. an input required by the issued instruction isn't yet available;
- memory dependency<sup>25</sup>, a load/store cannot be made because the required resources aren't available or are fully utilized, or too many requests of a given type are outstanding.

This further demonstrates the fact that matrix multiplication is a memory-bound problem in GPU and, so, poor in computation amount.

This is why we have too much short kernels for smaller input data size and heavy kernels for bigger input sizes.

## 5.4 Image processing

With image processing, i.e. Blur Box algorithm, we're facing a memory-bound kernel and especially rich of divergent execution flows, in fact we made similar tests to the ones for Matrix multiplication.

For each Image Processing we're working on input/output streams of small data parallel tasks, given in this case by small images. So, assume that task size is represented by  $N$ , that is the image resolution, more precisely given by  $N \times N$ , then in tests, the kernel execution configuration was set up as follows:

$$\text{BLOCK} = 1024$$

and

$$\text{GRID} = (\text{N}/\text{BLOCK}) + \text{BLOCK} - 1$$

---

<sup>24</sup>Execution dependency stall can be potentially reduced by increasing instruction-level parallelism (ILP). We saw in Chapter 3, that we can improve parallelism by increasing the amount of instructions per thread. This can translate in less threads having a greater work-load each. Furthermore we can put independent operations between dependent ones, to cover their latencies.

<sup>25</sup> Memory dependency stall type can potentially be reduced by optimizing memory alignment and access patterns.

Tesla P100 & Tesla M40	
Img. Order	In stream limit
128	64
256	256
512	1024
Data Parallel Tesla P100 & Tesla M40	
	1024
	2048
	4096
	8192

Table 5.10: Input dataset for Image Processing kernel. Above Stream parallel configuration, below the relative Data Parallel.

so we have

```
blockSize = (BLOCK, 1, 1)
gridSize=(GRID, 1, 1)
```

Then we performed the following executions:

1. **Classic data parallel approach** If we think to a picture as a matrix of pixels, then it's easy to see the analogy to the previous application.

In particular the Data Parallel approach will be tested giving a single image of large dimensions.

Again we're considering a single big image, such that it's comparable with some Stream Parallel versions. Similarly to matrix multiplication case, we remember that in data parallel version we have a data parallel big structure, it will be sent to the GPU and it will perform data parallel computations all over the data structure elements.

This is totally different from stream parallel case, where we have a stream of small data parallel images. However we can find some interest sizes for data parallel structures, such that they allow to perform the same amount of work of an overall stream parallel execution. In the lower part of Table 5.3 we show the order of dimension of the single and full data parallel images used to compare to some streaming parallel cases.

## 2. Streaming parallel

As previous cases of Farm parallel pattern, we have to put a limit on the input stream's tasks number (we recall that tasks are given by small images, that are small data parallel tasks).

In the upper portion of Table 5.10 are reported input stream limits and, for each of them, we test the three types of tasks size (small images resolution). Note that tasks are square images, so, for example, a size of 128 stands for a picture of  $(128 \times 128)$  resolution (16 384 pixels).

For every combination given by the variation on input parameters, we'll test for different numbers of CUDA streams: **Zero**, **Three** and **N<sub>SM</sub>** CUDA Streams (with  $N_{SM} = \#Streaming\ Multiprocessors$ ). The above test on different numbers of non-default streams, is implemented in a totally analogous way to the ones for the two applications showed above.

### 5.4.1 Results

As expected this image processing kernel demonstrates a bad fitting for Farm parallel pattern in GPU. It gives even worse performances than matrix multiplication.

We report in Table 5.11 all completion times, for zero-streams, three-streams and SM-streams versions.

We can see that the input stream limit, so the number of tasks per SM, is tested making it growing by a factor 4, in fact completion times follow this trend by increasing  $\approx 4\times$

CUDA Streams	Img Number	Img Size	Tesla M40	Tesla P100
0	64	128	1608.7367	1425.9833
	256		6153.5800	5670.5567
	1024		25422.7333	22774.7333
	64	256	5099.54	4648.4467
	256		20633.2	18077.1667
	1024		81256	73411.1333
	64	512	15652.6333	14281.8
	256		63566.5333	56694.1333
	1024		255928.3333	224584.6667
3	64	128	1547.5200	963.1717
	256		5477.0967	3851.38
	1024		21754.4667	15387
	64	256	4557.1400	3933.31
	256		17582.4	15724.8
	1024		71291.5	60029.0667
	64	512	14315.9333	11833.2
	256		56507.8667	49691.8
	1024		224832.3333	197940.3333
24 - 56	64	128	1482.0533	1043.4733
	256		5651.3833	4178.6333
	1024		21790.8333	16213.8667
	64	256	4461.85	3766.4833
	256		17751.7667	15682.6667
	1024		70421.1	63726.1667
	64	512	13900.0333	12902.0667
	256		54006.6667	51737.7333
	1024		226638	208308.3333

Table 5.11: Device completion times for Image processing kernel, all types of tested CUDA Streams number are reported, results are given for both P100 and M40.

		Tesla M40 (24 Streams)		Tesla P100 (56 Streams)	
Img Number	Img Size	Sp(3)	Sp(24)	Sp(3)	Sp(56)
64	128	1.0396	1.0855	1.4805	1.3666
256		1.1235	1.0889	1.4723	1.3570
1024		1.1686	1.1667	1.4801	1.4046
64	256	1.1190	1.1429	1.1818	1.2342
256		1.1735	1.1623	1.1496	1.1527
1024		1.1398	1.1539	1.2229	1.1520
64	512	1.0934	1.1261	1.2069	1.1069
256		1.1249	1.1770	1.1409	1.0958
1024		1.1383	1.1292	1.1346	1.0781

Table 5.12: Here are showed speedups for all data sets of image processing kernel. Results are reported for both devices.

proportionally with number of tasks arriving from the input stream.

Instead, fixed a certain number of tasks given from the input stream, we vary the task size to experiment dimensions that double from one to another. This makes the respective completion times growing by a slightly bigger factor, ie  $\approx 3\times$ .

But the really evident and important behavior is that we don't have much difference between the version not using CUDA Streams and the ones using them.

This is, in fact, confirmed by the speedups in Table 5.12, where we can see that almost everywhere the best speedup we can achieve is about 1, that means no speedup at all. So, we almost have no overlapping, we expected a similar behavior though.

## 5.5 Results Summary

Merging all obtained results, we can state that, under specific assumptions and adjustments, *Farm parallel pattern can give a speedup really near to linear one.*

We can observe this behavior especially in computation-bound case, where we're below the ideal speedup for a small quantity. Anyway, we can't achieve perfectly the ideal for more possible reasons:

- First we've to take into account that we have a "*stabilization phase*", meaning that when input stream starts to send first items we've a little interval where CUDA streams and SMs need to fill up, until we reach the peak occupation;
- We can have the rare situation where in a certain portion of time, say  $[t_i, t_i + \Delta_t]$ , we have multiple requests (from different CUDA Streams) for memory copy, ie we've  $> \text{numberOfCopyEngines}(= 2)$  simultaneous requests.

However, we only proved theoretically that  $Sp_{\#SM} \leq \#SM$ , just for completeness we could show some tests for a greater number of CUDA Streams, for example

$$\#CUDAStreams \geq 2 \cdot \#SM.$$

We briefly show the result for Simple-computational kernel, comparing the SM-version and the double-SM-version<sup>26</sup>. So this means we're trying to use an amount of CUDA streams that is the double of SMs number, just to see that anyway we can't achieve a speedup such that  $Sp_{\#SM} > \#SM$ .

And, as we expected, the Table 5.13 confirms the theoretical upper bound determined with Amdahl's law. The table clearly shows that the speedup is still  $Sp_{\#SM} \leq \#SM$ .

---

<sup>26</sup>We only report results for M40 machine, but performances are analogous on P100.

N items	M iterations	Comp.Time (48 Streams)	Sp(48)
245	10000	35.6340	19.4686
	400000	1204.41	22.7945
	800000	2379.3233	23.0691
49152	10000	70.2144	19.6877
	400000	2380.3433	23.0645
	800000	4734.9566	23.1857
98304	10000	135.2063	20.4526
	400000	4741.3866	23.1576
	800000	9537.72	23.0194
196608	10000	282.214	19.5913
	400000	9893.6566	22.1949
	800000	19760.9333	22.2204
393216	10000	519.2003	21.2989
	400000	18951.0666	23.1750
	800000	38050.7333	23.0795
786432	10000	969.7603	22.8021
	400000	37473.3333	23.4415
	800000	74944.7333	23.4355

Table 5.13: Here are showed completion time and speedup for  $48 = 2 \cdot \#SM$  CUDA Streams.



### 5.5.1 Stream parallel compared to Data parallel

As introduced on the tests settings, we executed and collected completion time measures for data parallel version too.

The latter clearly is setup in such a way that it computes the same workload that we're computing in stream version.

But we enforce again the concept that we're comparing the same application from the two different perspective of two totally different parallel patterns.

In fact, we have to remember that pure data parallel version will take a single data structure, having known size both theoretically and practically, and over all the content of this collection we perform data parallel computations, so as result we'll have again a big data structure having the same size as the input.

The Farm version, instead, takes a stream of small tasks that are sent to the GPU to perform data parallel computations on each of those small items and, as output, we'll have back the same amount of tasks, having the same size each.

So we're not clearly comparing directly input types and dimensions, but we're comparing the overall amount of work given by the two different versions.

Below we'll show results for each kernel type:

- **Simple-computation kernel**

From the Table 5.14 we can clearly see that data parallel has really similar performances with respect to stream parallel version(in SM-CUDA-streams setting). This is what we expected, since, the almost-linear speedup we obtained, means that we successfully overlapped different executions for small tasks. Formally, suppose that for any data/stream parallel comparison, we consider the same application and the same overall work load.

Call  $T_{DataPar}$  the amount of time needed to: copy the whole single data structure from host to device, perform data parallel computations on the whole data struc-

		<b>Tesla P100 (56 Streams)</b>		<b>Tesla M40 (24 Streams)</b>	
N items	M iterations	56 Streams	Data Parallel	24 Streams	Data Parallel
57344	10000	104.7727	51.3901	30.6074	34.9234
	400000	3968.9133	1825.36	1178.72	1178.71
	800000	7818.8433	3648.6333	2355.4133	2352.63
114688	10000	205.7297	90.2111	60.2087	58.0608
	400000	7828.1933	3574.74	2358.6567	2303.43
	800000	15691.8333	7145.6633	4714.36	4601.0867
229376	10000	407.712	179.2643	119.3447	112.1287
	400000	15687.9667	7124.46	4715.1233	4456.6633
	800000	31396	24437.4333	9425.92	8910.3567
458752	10000	803.2237	669.5133	238.249	241.901
	400000	31422.0333	26757.7	9429.89	9627.5267
	800000	62818.7	52896.3667	18853.9667	19256.4333
917504	10000	1619.5867	1361.9133	475.5907	441.5377
	400000	62793.4	53258.9667	18856.8	17591.4667
	800000	125575	105657	37705.6667	35190.9
1835008	10000	3229.0633	2682.6767	949.497	884.6433
	400000	125547.6667	105891	37711.9	35013.4667
	800000	251503	209018.6667	75445.2667	70397.2

Table 5.14: Simple-computational kernel. Comparison between completion times for stream parallel (max stream -56 and 24 respectively-) and data parallel versions. Results are reported for both devices.

ture, then copy back the whole result data structure. For stream parallel version we'll wish to have instead

$$T_{StreamPar} \approx \Delta_t + T_{DataPar}$$

where  $\Delta_t$  is a not predictable overhead, given by the different behavior on GPU of stream parallel w.r.t. data parallel.

In this  $\Delta_t$  we may also think to include those overheads due to imperfect streams overlapping, i.e. host/device copy or kernel execution that couldn't hide with other activities (partially or totally). So, in those cases where  $\Delta_t$  can be negligible with respect to the overall completion time, we hope to achieve:

$$T_{DP} \approx T_{SP}$$

where in general we expect that it's more likely to have  $T_{DP} \leq T_{SP}$ , than the contrary.

- **Matrix Multiplication kernel**

In M40 Table and P100 Table we'll see a really particular behavior, because on both devices we get consistently better performances from streaming parallel version, with respect to the data parallel version.

For Completeness in M40 Table we also introduced a column to compare the data parallel version with serial version (streaming parallel with zero CUDA Streams). From that column we can see that Data parallel version also has worse performances than serial version. Furthermore we introduced a column to compute the ratio  $T_{dataParallel} / T_{\#SMstreams}$ , and as we can see that from Stream Parallel version we have a gain of  $\approx 20\times$ .

This singular behavior again depends on the memory-bound nature of this implementation.

Mat Number	Mat Order	56 Streams	Event Time	Data Par Mat Order
225	128	20.8758	289.4027	1920
441		40.5783	898.0440	2688
900		74.6636	2256.0733	3840
1764		145.4767	7063.8233	5376
225	256	147.7650	2256.0733	3840
441		288.5343	7063.8233	5376
900		588.9643	17878.3667	7680
1764		1153.7333	56190.3667	10752
225	512	1173.3200	17878.3667	7680
441		2298.3967	56190.3667	10752

Table 5.15: Matrix multiplication kernel. Comparison between completion times for stream parallel (max stream -56) and data parallel versions (Partial dataset is of stream version is considered). Results are reported for P100.

Having a huge single data structure, computing a poor amount of arithmetic on a huge amount of items residing in global memory, inevitably leads to a over-occupancy in SMs (mainly due to latencies and thread stalls), dropping dramatically performances.

- **Image Processing kernel**

In this case, looking at Table 5.17, we can observe a fluctuation in behavior. Sometimes Data Parallel performs better than Stream Parallel, and sometimes the vice versa happens. But we recall that this type of kernel is peculiarly chosen to have divergent flows. Indeed, even the profiler reported the latency due to diverging flows as an issue.

Mat Number	Mat Order	Zero Streams	24 Streams	Data Parallel	Mat Order Data Par	(DataPar)/(24Str)
100	128	36.2095	19.3538	172.6997	1280	8.9233
196		74.2686	37.8561	456.3713	1792	12.0554
400		147.7337	65.6809	1315.6467	2560	20.0309
784		299.1380	128.3170	3590.4367	3584	27.9810
100	256	186.3913	130.0533	1315.6467	2560	10.1162
196		368.6813	254.2810	3590.4367	3584	14.1200
400		786.7537	518.6150	10437.7	5120	20.1261
784		1603.4933	1016.55	28711	7168	28.2436
100	512	1256.4	1034.0667	10437.7	5120	10.0938
196		2479.4333	2027.2867	28711	7168	14.1623

Table 5.16: Matrix multiplication kernel. Comparison between completion times for stream parallel (max stream -24-) and data parallel versions (Partial dataset is of stream version is considered).

Results are reported for M40.

		<b>Tesla M40 (24 Streams)</b>		<b>Tesla P100 (56 Streams)</b>		
M40 Img Number	Img Size	Streams 56	Data Par	Streams 24	Data Par	Img Size
64	128	1043.4733	1042.1233	1482.0533	1928.8767	1024
256		4178.6333	2522.0767	5651.3833	7369.49	2048
1024		16213.8667	25430.7667	21790.8333	29222.2	4096
64	256	3766.4833	2522.0767	4461.85	7369.49	2048
256		15682.6667	25430.7667	17751.7667	29222.2	4096
1024		63726.1667	46776.4667	70421.1	50971.8333	8192
64	512	12902.0667	25430.7667	13900.0333	29222.2	4096
256		51737.7333	46776.4667	54006.6667	50971.8333	8192

Table 5.17: Here is showed the data parallel vs. stream parallel comparison for image processing kernel. Results are reported for both devices.

## CHAPTER 6

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### Conclusions

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The main goal of this thesis was to experiment if a Farm parallel pattern could fit in GPU architecture and, if this was the case, how.

Even though a Streaming parallel pattern may seem so far from the concept of normal GPU use, we founded our attempt on the increasing and pervasive concept of General-Purpose computing. Nowadays it's a common practice to use the high parallelism and huge computational power of GPUs as co-processors, even if it isn't strictly for graphical problems.

Also the research moved, in last years, the focus on problems that generally are assigned to CPUs. Clearly, in General Purpose (GP) it's easy to spot applications that are clearly embarrassingly parallel; we recall that GPUs are mostly well suited in data parallel approaches.

However, there are many others problems that are really far from data parallel behavior and some of those cases GP-GPUs demonstrate a fair behavior (generally with some adjustments).

So it makes perfectly sense to inspect for new non-data parallel applications to fit in GPU model. The main goal, in general, is to exploit the high computation potential of GPUs.

### **6.1.2 Evaluation of the problem**

The starting point of this study was to consider and understand some main features and the functioning of a graphic processor, in particular taking into account of the organization about parallelism, threads, cores, internal memory and so on. We showed main GPU's and NVIDIA CUDA characteristics, briefly introducing them in Chapter 1 and deepening on more specific concepts in Chapter 2 and Chapter 3.

In the latter we also showed how some best practices and considerations were exploited to evaluate, implement and then test our model.

Once we had an overall view on tools and NVIDIA GPUs architecture, we had a base knowledge for the next step, i.e. to design a Farm parallel pattern for a graphic processor. Obviously some key problems have arisen:

- Handle the difference on input/output w.r.t. canonical data parallel problems, i.e. dealing with streams of data parallel tasks, instead of a single and completely data parallel structure;
- Manage the way we send data to device;
- Experiment different executions, each for different sizes of small tasks (from farm input stream);
- How to hide the overhead due to data transfers between host and device;
- How to execute many "small" kernels at the same time, instead of a single "big" one;



- How to exploit the resources of the GPU at their best possible, taking into account the considered application nature.

The first two points were accomplished by thinking to a *particular way* to use CUDA streams and asynchronous host/device memory copies.

In other words, we decided to use of many non-default streams, even an high amount w.r.t. the usual way they're used. This hopefully should have allowed us to hide data transfers and be able to run, at the same time, a lot of "small" data parallel tasks on the several GPU multiprocessors.

Clearly we could expect this parallel running of different kernels, since the considered data parallel tasks, arriving from a farm stream, were reasonably small to occupy a part of resources, but without saturate too much SMs and, so, allowing other small tasks to fit in multiprocessors.

The third point was mainly linked to the concept of *occupancy* evaluations. By both *empirical approach* and a study on *best practices* for GPUs, as we showed in Chapter 3, we get interesting results and we could experiment were occupancy was a benefit. However, we also saw how occupancy may not be a relevant factor; a lot of performances bottlenecks may depend on the kernel nature. We have to face some **latencies** that happens inside the Streaming Multiprocessors, in our study we mainly pointed out two types of bottleneck in kernels: **memory-bounded** code and **diverging flows**.

Those concepts are straightly linked to the problem of last point in the above list.

The fourth and fifth points are again strongly related to a powerful programming technique in CUDA: *Asynchronous calls* and *CUDA Streams*.

We recall that here asynchronous is from a host side point of view, with respect to the device. That is, host can continue executing his code, after invoking a memory copy (or any other call that is generally blocking). Any asynchronous call will be forwarded to GPU that will "silently" work, sending back eventual results to CPU, that hasn't to wait on device calls to end.

We pointed out that often asynchronous calls require to have some synchronization point. Sometimes CUDA codes with asynchronous calls are implemented introducing *explicit synchronizations*, in order to have correct results and avoid memory overwriting. We also met that problem, having a lot of CUDA streams trying to write back results, possibly at the same time, this sometimes led to overwriting data from another non-default stream<sup>1</sup>.

This problem mainly showed up in device memory: we needed sufficient GPU memory locations for all streams.

In device side, the possible solutions are, in general, to:

- Have a reduced amount of space in global memory and use some explicit synchronization and that's the most used approach;
- Reserve enough memory to support data transfers for each single CUDA streams and, so, avoid to use explicit synchronization.

The first approach can be used in problems that are more suitable for GPUs, as probably it introduces a negligible amount of overhead, but in a stream parallel context it can cause a performance drop. That's why we decided to use the second approach. Synchronization, with such a particular use of CUDA streams, can give a bottleneck, slowing down the amount of tasks we're capable to send to SMs.

The first impression could be to risk for a saturation in global memory, but in our case it didn't happen, since we were using relatively small tasks of data parallel data (even if the reserved space for tasks has to be as much as number of CUDA streams, i.e. SMs number).

This doesn't mean that it cannot exist any stream parallel application where synchronization may not result in an disadvantage<sup>2</sup>.

---

<sup>1</sup>We recall that operations issued inside a certain non-defaults stream are executed serially in between them. In commands, belonging to different CUDA streams, instead, we can have an undefined behavior, i.e. we don't have any guarantee on the execution order.

<sup>2</sup>Maybe to hide some other computations that are happening in the same time. Again it's a matter of

Furthermore trying a *hybrid approach* could be a starting point for future works, where hybrid means having less allocated global memory locations (than CUDA Streams number) and introduce only few explicit synchronizations.

### 6.1.3 Implementation and tests

Those ideas and designs were implemented as described in Chapter 4. We decided to implement different kind of kernels to experiment the behavior of Farm parallel pattern in different conditions.

We recall that we decided to implement three types of kernels: Simple-computational, Matrix Multiplication and Image processing.

The first would have been the one from which we expected good performances, while we expected worse completion times and speedups from the second kernel type and even worse from the third one.

The next step has been to build tests, gather results and make the following considerations.

Tests have been set up in such a way to observe the performances of our model in different situations, such as varying the tasks size and varying the pressure on CUDA Streams, i.e. the number of these tasks sent on a certain non-default stream and so the number of tasks that a certain SM has to execute.

What we wanted to mainly measure was:

- the global time spent to "consume" an input stream by transferring data one task per time, do all computations of the kernel and send back results. This is what we considered the *serial version* for our applications, in other words the approach without CUDA Streams;
- the global time spent to "consume" an input stream by overlapping more data

---

experimenting according to the type of problem we're facing.

transfers and more kernel executions. This is what we considered the *parallel version*, in other words the approach using CUDA Streams (three as base case or equal to the number of SMs as special case);

- the completion time of the relative data parallel version, i.e. assuming that the input is given by a single data structure, it should anyway compute an amount of work equal to the overall work computed by all small farm tasks.

The latter point means only that we’re doing a fair comparison, but the nature of the two problems and input/output data is completely different<sup>3</sup>.

### 6.1.4 Results and considerations

The results, in Farm parallel pattern for GPU, using CUDA streams, gave just what we expected:

1. Simple-computational kernel showed a good overlapping, especially among kernels, clearly with some appropriate adjustments. We get the expected speedups and the version with the maximum number of CUDA Streams<sup>4</sup> performs almost as the data parallel version;
2. Matrix multiplication kernel showed a low overlapping behavior, especially as matrices size increased. We got poor speedups, however the version with CUDA Streams often performed better with respect to the data parallel version;
3. Image processing kernel showed an almost inexistent overlapping behavior. We get no speedups and the version with the maximum number of CUDA Streams performed sometimes better sometimes worse than the data parallel version.

---

<sup>3</sup>Farm has discussed and compared with Map mainly in Chapter 3

<sup>4</sup>That is the version with an amount of non-default streams equal to the number of Streaming Multi-processors.

From those results we understand that we have the best gain when we have long computations on each single task.

In fact, as we showed in chapter 5, the first type of kernel has a high computational intensity, while the others two are memory bound and, so, provide a counterexample for Farm on GPU.

Even if host/device data transfers introduce a not-negligible overhead, in Farm pattern, we're carrying small data parallel groups of items.

Furthermore, these small tasks aren't available all at the same time, they arrive one at time, as they're generated from an input stream. This leads potentially to a low data transfers overlap, just for a timing matter.

That's why we should mostly rely on overlapping kernels, as they should lasts longer than memory copy<sup>5</sup> and so we've more chances to achieve an overlap.

About this consideration we recall 5.3.

That's why most of the problems in performances appeared in memory-bound kernels, because we have a lot of memory operations, merged with a really small amount of work per thread to do. This leads to inefficient kernels, that, in any case, last too short to afford a good overlap with other kernels or transfers.

Furthermore, we saw that this behavior in memory-bound can even degrade as the tasks, sent to the GPU, grow in size. It's like in our Matrix Multiplication case, because we had a high number of thread blocks occupying all hardware resources, for a single matrix multiplication. Anyway, in this application, having longer kernels lead to completely monopolizing Multiprocessors, without permitting to other kernels to fit in SMs.

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<sup>5</sup>This isn't a rule, it just often happens, as in our applications. There may still be cases in which this statement is false.

### 6.1.5 Final remarks and further works

The results and considerations just discussed in previous section, expose the following necessities for Farm parallel pattern:

- It better performs in high-computational intensity scenarios;
- We get the best advantages from parallelizing executions, more than host/device memory copies;
- It relies on overlapping between CUDA streams, meaning it needs quite long kernels executions to hide the host latency deriving from the acquisition of items from an input stream;
- Kernel launches should be configured such that they don't monopolize many multiprocessors (the best would be at most one SM occupied by a single kernel call).

The second and third points, again underline the necessity of having a small amount of memory traffic with respect to computations.

Maybe it would be a good compromise even to have only a slightly higher amount of computations w.r.t. memory operations. This means that maybe we can observe acceptable performances, from GPU Farm, even in an average case.

In this work, in fact, we considered application having an opposite behavior.

In terms of arithmetic intensity and Roofline model this concept would be expressed by  $I \approx \pi/\beta$ , meaning that we're working on a kernel having a behavior in the middle of computation and memory-bound.

The above requirements may translates in a quite challenging effort, especially in evaluating problems, experimenting and profiling performances, more than in implementation difficulty.

As we said before, this especially holds in all those cases where we have memory-bound

problems. However, in this case may be tried some future workarounds to improve performances, even in apparently not suitable applications, in particular:

- using efficient memory access patterns for GPU memory (especially needed for global);
- assigning more work to each thread, for example giving more instructions to execute per kernel (Instruction Level Parallelism) and alternate dependent instruction with independent ones[13, 8];
- exploiting shared memory, it has smaller dimensions but it's much more faster than global memory;
- changing the scheduling policy of input tasks among the CUDA Streams.

These stratagems may expand to a lot of further applications using Farm parallel pattern on GPUs in the future.

As an example, numerous studies have been made on matrix multiplication to optimize device global memory latencies with shared memory. Other studies showed that we can give smaller kernel configurations, in order to make each threads perform several matrix multiplications[8], instead of computing only one element of the result matrix for each threads (as it happens in our simple and inefficient approach).

So, merging those optimizations with Farm parallel pattern may give, in future, some interesting results.

Given that this thesis based all hypothesis on equally sized tasks during a certain Farm execution on an input stream, i.e. on balanced workloads for each kernel, another interesting further study could be done in those scenarios treating unbalanced tasks of works, leading again to a stream of data parallel tasks, but giving different workloads among the respective kernel launches.

Suppose, for example, a scenario where the input stream sends items at fluctuating

speeds, so the chunk size may be established according to a time interval instead of a predefined buffer size. This means send portions of items of unknown size to the GPU.

Another assumption on which we based all this study was in having a certain type of scheduling.

In particular, we scheduled small data parallel tasks, arriving from the input stream, towards CUDA Streams with a Round Robin policy.

In the problems treated in this thesis or in other Farm applications where we're trying to achieve a performance improvement (memory-bound ones for example), maybe other scheduling techniques can be adopted.

This potentially may result in a more efficient spreading of tasks between CUDA streams. So, if this gives a better exploitation in Multiprocessors resources or a better work distribution, this may give a performance improvement.



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