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Performance Upper Bound Analysis and Optimization of SGEMM on Fermi and Kepler GPUs

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Abstract

In this paper, we present an approach to estimate GPU applications' performance upper bound based on algorithm analysis and assembly code level benchmarking. As an example, we analyze the potential peak performance of SGEMM (Single-precision General Matrix Multiply) on Fermi (GF110) and Kepler (GK104) GPUs. We try to answer the question of how much optimization space is left for SGEMM and why. According to our analysis, the nature of Fermi (Kepler) instruction set and the limited issue throughput of the schedulers are the main limitation factors for SGEMM to approach the theoretical peak performance. The estimated upper-bound peak performance of SGEMM is around 82.5% of the theoretical peak performance on GTX580 Fermi GPU and 57.6% on GTX680 Kepler GPU. Guided by this analysis and using the native assembly language, on average, our SGEMM implementations achieve about 5% better performance than CUBLAS in CUDA 4.1 SDK for large matrices on GTX580. The achieved performance is around 90% of the estimated upper-bound performance of SGEMM on GTX580. On GTX680, the best performance we achieve is around 77.3% of the estimated performance upper bound. We also describe how to use native assembly language directly in the CUDA runtime source code.

Categories and Subject Descriptors C.1.4 [Processor Architectures]: Parallel Architectures; C.4 [Performance of Systems]: Modeling techniques; F.2.1 [Numerical Algorithms and Problems]: Computations on matrices

General Terms Measurement, Performance

Keywords Kepler GPU, Fermi GPU, SGEMM, CUDA, Performance Upper Bound Analysis

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1. Introduction

Nowadays, vendors are putting more and more computing cores per die and the GPU architecture is evolving rapidly. Fermi and Kepler GPUs are the present generations of NVIDIA GPUs [13, 14]. There are many studies about optimizing specific kernels on Fermi GPU. However, since the architecture is changing with each generation, we may need to repeat the optimization work again very soon. Unfortunately, no practical performance upper bound evaluation is available to the developers. In practice, developers apply several optimization techniques based on the analysis to the algorithm or serial code, and their expert experience. Then developers may modify the optimizations with feedback provided by tools like NVIDIA Visual Profiler [3]. However, they can not be sure how far the obtained performance is from the best achievable performance. In this paper, we present an approach to project performance upper bound using algorithm analysis and assembly code level benchmarking.

To understand GPU performance result, there exist many works about how to project/predict CUDA applications' performance using analytical or simulation methods. Meng et al. [10] proposed a GPU performance projection framework based on annotated code skeletons. Hong and Kim [6] introduced the MWP-CWP model to predict CUDA application performance using PTX code. Bakhoda et al. [5] developed a detailed GPU simulator and the simulator also uses the PTX code as input. Recently, Sim et al. [17] extended the MWP-CWP model and utilize the assembly code of CUDA kernel to predict performance. The quantitative GPU performance model proposed by Zhang and Owens [20] is also based on the native assembly code. Since very little information about the underlying GPU architecture is disclosed, it becomes very unlikely to build accurate simulators for each new GPU generation. Luckily, the results [6, 10, 17, 20] show that we can have very good approximation of GPU performance using analytical approaches. However existing GPU performance models all rely on certain level of an application's implementation (C++ code, PTX code, assembly code...) and do not answer the question of how good the current optimized version is and whether further optimiza-

tion effort is worthwhile or not. Different from existing GPU performance models, our approach does not project the possible performance from certain implementations, but the performance upper bound that an application cannot exceed.

Researchers are also interested in the outcome of different optimization combinations on GPUs. The roofline model [19] is well known for estimating the optimization effects. The recent work by Sim et al. [17] studied the effects of different optimization techniques on GPUs using the similar approach as the roofline model. However, the chosen optimizations normally rely on the initial code version and different optimizations are likely to have complex impacts on each other. Our approach tackles the problem from the opposite angle as the roofline method. We first assume an optimistic situation on GPUs (no shared memory bank conflict, global memory accesses are all coalesced, all the auxiliary operations like address calculations are neglected, etc.). Then we try to predict a performance upper bound when mapping an application on the GPU based on the constraints introduced by the architecture, the instruction set and the application itself, or the constraints that we are not able to eliminate using optimization techniques. With a tight performance upper bound of an application, we have an evaluation on how much optimization space is left and can decide the optimization effort. Also, with the analysis, we can understand which parameters are critical to the performance and have more insights into the performance result. Hence, with these knowledge, it would be easier for the community to move to the new architecture.

As an example, we analyze the potential peak performance of SGEMM (Single-precision General Matrix Multiply) on Fermi (GF110) and Kepler (GK104) GPUs. GEMM¹ operation is essential for Level 3 BLAS (Basic Linear Algebra Subprograms) [2] routines and generally represents the practical best performance of a computer system. If we compare the performance of SGEMM from CUBLAS with the theoretical peak performance, on Fermi, it achieves around 70% and on Kepler, only around 42% of the theoretical peak performance. There are already some articles about optimizing GEMM kernels on Fermi GPU [11] [18], and an auto-tuning framework has also been presented [7]. In this research, the focus is to answer the question of how much optimization space is left for SGEMM and why. We also show that the analysis can help optimization efforts since it uncovers critical parameters. Only single precision SGEMM is evaluated, since we could only access the GTX580 Fermi and the GTX680 Kepler Geforce cards, which have much poorer double precision performance than Tesla products. It is not really worth the effort to study the DGEMM performance on Geforce GPU. In November 2012, NVIDIA has announced the new Tesla K20X Kepler GPU (GK110)

¹GEMM performs the matrix-matrix operation $C := \alpha * op(A) * op(B) + \beta * C$. α and β are scalars, and A, B and C are matrices. $op(X)$ is $op(X) = X$ or $op(X) = X^T$.

and the documented SGEMM efficiency is around 73% of the theoretical peak performance [15]. The K20X Kepler GPU (GK110) architecture is different from the GTX680 (GK104) and uses a different instruction set (each thread can utilize maximum 255 registers on the new architecture while the limit is 63 on GTX680 GPU). With a Tesla GPU card, it should not be difficult to extend the analysis to SGEMM and DGEMM on the Tesla GPU using our approach.

This paper is organized as follows: Section 2 introduces our target GPU architectures. Section 3 introduces our assembly level benchmarking approach. Section 4 presents our analysis for performance upper bound of SGEMM on Fermi and Kepler GPUs. In Section 5 assembly code level optimization methods and performance result of SGEMM are presented. Section 6 concludes this study.

2. GPU Architecture Characteristics

NVIDIA GPUs are composed of a cluster of independent SMs (Streaming Multiprocessor). Each SM includes several SPs (Streaming Processor), which is the basic computing component, scheduler, SFUs (Special Functional Unit), LD/ST (Load/Store) Units, register file and a unified shared memory/L1 cache. The SPs inside one SM are similar to a lane of SIMD engines and they share the memory resource of the SM like the registers and shared memory.

A comparison of the three generations of NVIDIA GPUs is illustrated in Table 1. From GT200 to Kepler GPU, the number of SPs increases dramatically, from 240 (GTX280, 65nm) to 1536 (GTX680, 28nm) [13, 14]. However, if we consider the memory resource (registers and shared memory) per SP, the on-die storage per SP actually decreases. Previous generations have two clock domains in the SM, the core clock for the scheduler and the shader clock for the SPs. The shader clock is roughly twice the speed of the core clock. On Kepler (GK104) GPU, shader clock no longer exists, the functional units with SMs run at the same core clock. However, to compare the different generations more easily, we still use the term shader clock on Kepler GPU. In the rest of this paper, all throughput data is calculated with the shader clock.

A typical CUDA [12] program normally creates thousands of threads to hide memory access latency or math pipeline latency. The threads are grouped into 1D to 3D *blocks*, and further into 1D or 2D *grids*. Each block is assigned to one SM at execution time. The *warp* is the basic execution and scheduling unit of a SM, and is composed of 32 threads. We define a **warp instruction** as the same instruction shared by all threads in the same warp, and a **thread instruction** as the instruction executed by one thread. So a *warp instruction* launches 32 operations or consists of 32 *thread instructions*. On the SM, only a limited set of threads can run concurrently (active threads). On one hand, the increased SPs require more active threads to hide latency. On the other hand, the register and the shared memory resource

	GT200 (GTX280)	Fermi (GTX580)	Kepler (GTX680)
Core Clock (MHz)	602	772	1006
Shader Clock (MHz)	1296	1544	1006
Global Memory Bandwidth(GB/s)	141.7	192.4	192.26
Warp Scheduler per SM	1	2	4
Dispatch Unit per SM	1	2	8
Thread Instruction issue throughput per shader cycle per SM	16	32	128?
SP per SM	8	32	192
SP Thread Instruction processing throughput per shader cycle per SM (FMAD/FFMA)	8	32	192?
LD/ST (Load/Store) Unit per SM	unknown	16	32
Shared Memory per SM	16KB	48KB	48KB
32bit Registers per SM	16K	32K	64K
Theoretical Peak Performance (GFLOPS)	933	1581	3090

Table 1. Architecture Evolution

limits the number of active threads. For the same application, the active threads that one SP supports actually decreases because of the reduced memory resource per SP from Fermi GPU to Kepler GPU. More instruction level parallelism within one thread needs to be explored (Section 4.3).

For Fermi (and Kepler GK104) instruction set, there is a hard limit of maximum 63 registers per thread (for GT200 generation the limit is 127 registers per thread) since in the instruction encoding, only 6 bits are left for one register.

3. CUDA Programming with Native Assembly Code

For performing this study, we have to develop some software components and reverse engineer many characteristics of the hardware. We use the GPU assembly code directly with an assembly tool Asfermi[1]. Asfermi was first developed to work on Fermi GPU. We patch Asfermi to support Kepler GPU (GK104) and manage to use native assembly language directly in the CUDA runtime source code.

3.1 Using Native Assembly Code in CUDA Runtime API Source Code

Programming in assembly code on NVIDIA GPUs is not publicly supported by the company. However, our analysis is requiring such programming. With an assembly tool for Fermi GPU called Asfermi [1] and a little hacking into the CUDA programming compiling stages, we manage to use hand-tuned GPU assembly code in CUDA projects using CUDA runtime APIs .

There are several advantages of using assembly code or native machine code directly instead of using high level languages like C++. First, we can carefully control the register allocation since the register resource per thread is very limited and sometimes the compiler may spill many registers for programs utilizing much register resource per thread like SGEMM. Second, the instruction order can be carefully designed to better prefetch data from global memory and mix different instruction types to get better throughput. Third,

SIMD-like instructions (LDS.64 or LDS.128) could be used intentionally to reduce the instruction number. Also, we can control the exact behavior of the machine code. For example, the compiler might choose to use wider load instructions (LDS.64 or LDS.128) based on the data alignment in shared memory. However, using wide load instructions does not always benefit the performance (Section 4.1).

3.2 Kepler GPU Binary File Format

Asfermi was first developed to work on Fermi GPU. We patched Asfermi to support CUDA sm_30 (GK104 Kepler GPU). However, although the CUDA program using assembly code kernel can still run correctly on Kepler GPU, the performance is very poor. The reason is that new control information is embedded into the CUDA binary file to help processor scheduling. According to the GTX680 white paper [14], the compiler places the scheduling information along with the actual instructions in the CUDA binary file.

The scheduling information (we call it control notation) is placed before each group of 7 instructions and has the format of 0xXXXXXXXX7 0x2XXXXXXXX. 0x7 and 0x2 are identifiers and the rest of the notation is separated into 7 fields and associated with each following instruction. It is similar to the explicit-dependence lookahead used in Tera computer system [4]. Unfortunately, NVIDIA does not disclose the encoding of the control notation. We do not know how to generate the control notation exactly as the nvcc compiler. In our implementation of SGEMM on Kepler GPU, as a compromise, we use the same control notation for same kind of instructions and try to find the best combination of those notations for major instruction types. However, our decryption of the notations is still not enough.

3.3 Math Instruction Throughput on Kepler GPU

Understanding and modeling the behavior of math instructions on Kepler GPU is a major difficulty. We use two approaches to test the throughput of math instructions. First, a kernel is written in C++ code and compiled into binary with control notations embedded by nvcc. Second, a kernel is written in assembly code directly and the controlling

FADD R0, R1, R0	128.7	FADD R0, R1, R2	132.0
		FADD R0, R1, R3	66.2
FMUL R0, R1, R0	129.0	FMUL R0, R1, R2	132.0
		FMUL R0, R1, R3	66.2
FFMA R0, R1, R4, R0	129.0	FFMA R0, R1, R4, R5	132.0
		FFMA R0, R1, R3, R5	66.2
		FFMA R0, R1, R3, R9	44.2
IADD R0, R1, R0	128.7	IADD R0, R1, R2	132.4
		IADD R0, R1, R3	66.2
IMUL R0, R1, R0	33.2	IMUL R0, R1, R2	33.2
		IMUL R0, R1, R3	33.2
IMAD R0, R1, R4, R0	33.2	IMAD R0, R1, R4, R5	33.1
		IMAD R0, R1, R3, R5	33.2
		IMAD R0, R1, R3, R9	26.5

Table 2. Examples of Math Instruction Throughput on Kepler GPU with Various Operand Register Indices

notations are embedded with our parsing tool. Each thread executes the same 8192 math instructions. Each block has 1024 threads without synchronization and 40960 blocks are spawned to keep the GPU busy.

Instruction FFMA performs single precision fused multiply-add operation ($FFMA\ RA, RB, RC, RD$ performs the operation $RA := RB * RC + RD$). With the first approach, the instruction throughput of $FFMA\ R9, R8, R9, R5$ is measured as 129.2 operations per shader cycle². With the second approach and the control notation of 0x25, the throughput is 132.0 operations per shader cycle (The actual shader clock cannot be obtained during execution. All throughput data is calculated by boost clock of 1058MHz[14]).

Some math instructions’ throughput is illustrated in Table 2 measured with the second approach. In these cases, the scheduling function units on one SM can only issue about maximum 132 thread instructions per shader cycle, which is much lower than the SP’s processing throughput (192 thread instructions per shader cycle). If some of the three source registers are the same (like $FFMA\ RA, RB, RB, RA$), with some carefully designed code structures, the FFMA throughput can approach around 178 thread instructions per shader cycle. However, considering ‘useful’ FFMA’s throughput, that is ($FFMA\ RA, RB, RC, RA$), the maximum single precision performance for many applications like SGEMM on GTX680 GPU (GK104) cannot exceed around 68.75% (132/192) of the claimed performance (3090GFlops) by NVIDIA.

Our benchmark result also shows that the instruction throughput is related to register indices. According to some other experiments, we speculate that the registers reside on four banks. Take the instruction $FFMA\ RA, RB, RC, RD$ for instance, if there are two different source registers on the same bank, the throughput drops by 50%, and if all three source registers RB, RC, RD are different registers on the same bank, the throughput is around 33.3% of the best

²The actual implementation is not 8192 $FFMA\ R9, R8, R9, R5$ instructions per thread but 4 independent FFMA instructions like $FFMA\ R9, R8, R9, R5$ unrolled by 2048 times.

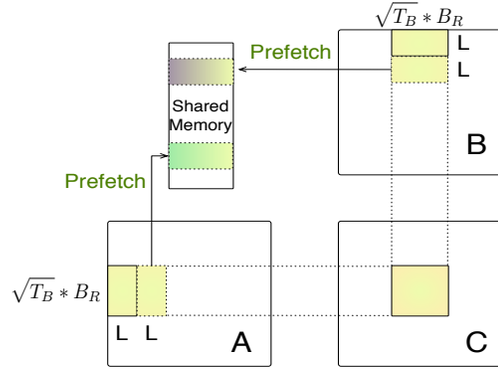


Figure 1. SGEMM Implementation

case. We name the four banks as even 0 ($R_{index} \% 8 < 4$ && $R_{index} \% 2 == 0$), even 1 ($R_{index} \% 8 \geq 4$ && $R_{index} \% 2 == 0$), odd 0 ($R_{index} \% 8 < 4$ && $R_{index} \% 2 == 1$), and odd 1 ($R_{index} \% 8 \geq 4$ && $R_{index} \% 2 == 1$). Since we implement SGEMM with assembly code directly, the register indices have to be carefully chosen to make sure there is no bank conflict. The detailed optimization is illustrated in section 5.4.

4. Analysis of Potential Peak Performance of SGEMM

The general analysis approach can be similar for all applications while the detailed analysis process may differ from application to application. Our method is applicable for applications which use a few major instruction types and a simple execution path. Many high-performance computing kernels have this characteristic, especially linear algebra routines. Our analysis requires characteristics of the architecture such as register file size, maximum number of registers per thread, shared memory size, instruction throughput for different instruction mix, etc. Those characteristics need to be collected on the real hardware and are independent of the effective application.

First, we should analyze the instruction types and percentage of a routine. Second, we should find the critical parameters which affect the different instructions’ mixing percentage. Third, we analyze how the instruction throughput changes when we vary these critical parameters. Fourth, we can use the instruction throughput with critical parameters’ optimal combination to estimate the performance upper bound. With this approach, not only we can have the performance upper bound estimation, know how much performance gap is left and decide the optimization effort, but we can also understand what parameters are essential to the performance and how to distribute our optimization effort.

For SGEMM, all well-implemented SGEMM kernels actually utilize shared memory on the GPU to reduce the global memory pressure as illustrated in Figure 1. First, data is loaded from global memory to shared memory and then threads within one block can share the loaded

data in the shared memory. For Fermi (GF110) and Kepler (GK104) GPUs, arithmetic instructions like FFMA cannot take operands from the shared memory. Since LDS instructions are needed to load data first from shared memory into registers, most of the instructions executed in SGEMM are FFMA and LDS instructions. For instance, in our SGEMM implementation with 1024x1024 matrix size, 80.5% of instructions executed are FFMA instructions and 13.4% are LDS.64 instructions. So essentially, in our analysis, we define a few key parameters and study the instruction throughput mixing FFMA and LDS.X instructions while varying these parameters.

The rest of this section is our analysis of SGEMM’s performance upper bound. We show that the analysis can give good insights about how to optimize a specific kernel (SGEMM) and help us to understand the performance result.

4.1 Using Wider Load Instructions

To achieve better performance, it is essential to minimize auxiliary instructions’ percentage. By auxiliary instructions, we mean non-math instructions, especially LDS instruction. The assembly code for CUDA sm_20 (GF110 Fermi GPU) and sm_30 (GK104 Kepler GPU) provides SIMD-like LDS.64 and LDS.128 instructions to load 64bit and 128bit data from the shared memory. Using wider load instructions can reduce the total number of LDS instructions.

According to our benchmarks, on Fermi GPU, the peak throughput for LDS instruction is 16 32bit-operations per shader clock per SM. Using LDS.64 instructions does not increase the data throughput and the LDS.128 instruction normally leads to 2-way shared memory bank conflict on Fermi GPU. LDS.128 has the throughput of only 2 thread instructions per shader cycle on one SM. On Kepler GPU, the throughput for LDS operation is measured as 33.1 64bit operations per shader clock per SM. Using the 32bit LDS operation actually decreases the data throughput in half comparing with using LDS.64 instructions and properly used LDS.128 instruction does not introduce penalty.

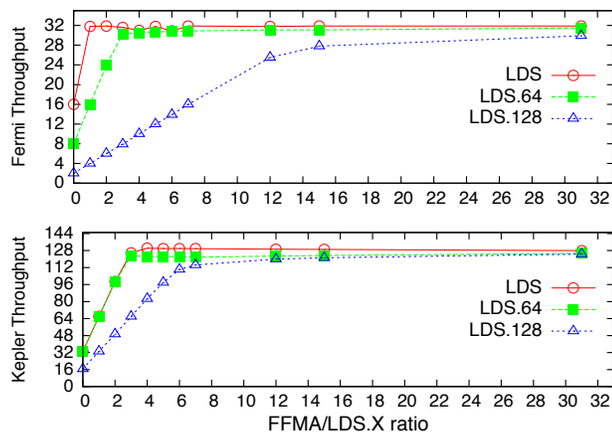


Figure 2. Thread Instruction Throughput Mixing FFMA and LDS.X

Figure 2 illustrates the instruction throughput of mixing FFMA and LDS.X instructions. While gradually increasing the ratio of FFMA instructions to LDS instructions, the overall instruction throughput approaches the FFMA’s peak processing throughput. The instruction ratio of FFMA to LDS.X depends on the algorithm parameters such as register blocking size. Apparently, the overall performance does not always benefit from using wider load instructions. However, the compiler might choose to use the wider load instructions based on the data alignment in the shared memory. With the native assembly language, it is possible for us to carefully design the data layout and use the best instruction type.

4.2 Register Blocking

As in Table 1, the scheduler of GT200 GPU can issue one warp instruction like FFMA per core cycle and since there are 8 SPs per SM, SPs need 4 shader cycles to process one warp instruction. Apparently, as the issue throughput is higher than the SP’s processing throughput, math instructions executed in SPs cannot saturate the scheduler’s issue throughput. So the scheduler has some ‘free cycles’ to issue instructions to other functional units.

On Fermi GPUs, SM are redesigned with 2 warp schedulers and 32 SPs. Each warp scheduler, equipped with one dispatch unit, issues instructions to 16 SPs. With an issue rate of one warp instruction per shader cycle, the schedulers’ ability could be fully utilized by 32 SPs. The theoretical peak performance for math instructions comes from the SPs’ performance. The percentage of other instructions becomes an issue when there are many auxiliary instructions: there are fewer cycles left for schedulers to issue *useful* instructions like FFMA.

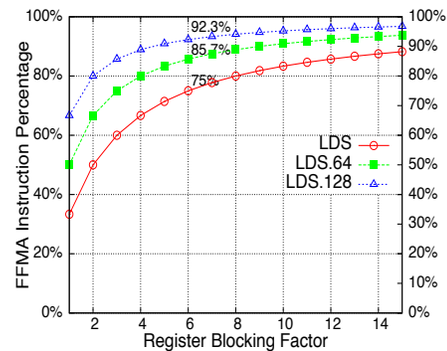


Figure 3. FFMA Instruction Percentage in SGEMM Main-loop with Different Register Blocking Factors

In the worst case, without any register reuse, 2 LDS instructions are needed to fetch data for 1 FFMA instruction in the SGEMM main loop. In that case, only 1/3 of the instructions are floating point operations. Blocking is a well-known technique to better utilize memory hierarchy for scientific programs [8, 9]. To increase the percentage of math instructions, register blocking is needed. We illustrate the percent-

age of FFMA instructions varying register blocking factors in Figure 3.

If 6-register blocking is used (which is the case of our SGEMM implementation on Fermi GPU), the FFMA/LDS.X ratios are 3:1, 6:1, and 12:1 if shared memory accesses are implemented with LDS, LDS.64 and LDS.128 respectively. The percentage of FFMA instructions is 75%, 85.7% and 92.3%. On Fermi GPU, the overall instruction throughputs for one SM in these cases are 31.3, 30.4 and 24.5 thread instructions per shader clock. Because using LDS.128 instruction may lead to extra penalties, even if all the accesses to shared memory are implemented with LDS.128, in the best case we can only achieve around 71% ($\frac{24.5}{32} * 92.3\%$) of SMs' single precision floating point performance. Also, in many cases, a lot of padding in shared memory has to be used to get proper data alignment. Apparently, it is not worth the programming effort to mix FFMA with LDS.128 for SGEMM on the Fermi GPU.

4.3 Active Threads on SM

Normally, the more active threads one SM executes, the higher performance the GPU can achieve. Since register and shared memory resource is limited per SM, only a limited set of warps can be executed concurrently (T_{SM}).

$$T_{SM} * R_T \leq R_{SM} \quad (1)$$

The registers that each thread can utilize (R_T) is less than or equal to 63 on Fermi and Kepler GPUs (R_{Max}). Furthermore, the register budget of the active warps cannot exceed the SM's register amount (R_{SM}) (Equation 1).

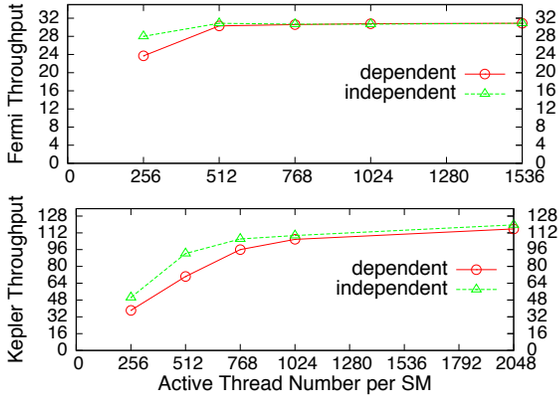


Figure 4. Instruction Throughput Mixing FFMA and LDS.64 with Ratio of 6:1

Figure 4 illustrates the instruction throughput mixing FFMA and LDS.64 instructions with ratio 6:1 under different number of active threads on one SM. We tested two cases. In the first case ('independent' in Figure 4), 6 FFMA and 1 LDS.64 instructions are all independent. In the second case ('dependent' in Figure 4), 6 FFMA instructions are dependent on one LDS.64 instruction. The second case is closer to the actual implementation of SGEMM. On Fermi

GPU, with 512 active threads, the instruction throughput of the second case is already close to the best situation. On Kepler GPU, however, with fewer than 1024 active threads, the Kepler GPU is very sensitive to the dependences between instructions.

4.4 Register and Shared Memory Blocking Factors

Larger register blocking size can introduce more register reuse within one thread and higher percentage of FFMA instructions. However, the register blocking size is limited by the register resource on the SM and the instruction set constraint. With a register blocking factor B_R , if we only consider the registers needed for blocking, we can describe the resource constraint as Equation 2.

$$B_R^2 + B_R + 1 < R_T \leq R_{Max} \quad (2)$$

This loose condition for register blocking factor B_R can be used to roughly estimate B_R . B_R^2 is the register set needed to hold C sub-matrix per thread, B_R is one column/row of A or B sub-matrix. For instance, with maximum 63 registers per thread, $B_R \leq 7$.

As depicted in Figure 1, $T_B * B_R^2$ is the size of the C sub-matrix per block (each block has T_B threads) and $\sqrt{T_B * B_R^2} * L$ is the size of a sub-matrix for A or B (L is the stride). To overlap the data transfer and the computation, extra registers are needed to fetch data from global memory to shared memory since no direct data transfer is provided between the two memory space. The stride L needs to be chosen such that each thread loads the same amount of data (Equation 3).

$$(\sqrt{T_B} * B_R * L) \% T_B = 0 \quad (3)$$

Considering data prefetching from global memory and a few registers to store the addresses of matrices in global memory and shared memory (R_{addr}), the overall strict constraint for register blocking factor can be described as Equation 4.

$$B_R^2 + \frac{2 * \sqrt{T_B} * B_R * L}{T_B} + B_R + 1 + R_{addr} \leq R_T \leq R_{Max} \quad (4)$$

Since shared memory is allocated in block granularity, for Blk active blocks, $Blk * 2 * \sqrt{T_B} * B_R * L$ is needed to store prefetched global memory data (Equation 5). The shared memory blocking factor can be defined as $B_{Sh} = \sqrt{T_B * B_R^2}$. With the shared memory blocking factor B_{Sh} , the performance bounded by global memory bandwidth can be roughly estimated using Equation 6.

$$Blk * 2 * \sqrt{T_B} * B_R * L \leq Sh_{SM} \quad (5)$$

$$\frac{P_{MemBound}}{\#GlobalMem.bandwidth} = \frac{2 * B_{Sh}^2}{2 * B_{Sh} * 4} \quad (6)$$

4.5 Potential Peak Performance of SGEMM

The instruction factor F_I is the ratio of FFMA instructions in the SGEMM main loop (We only consider FFMA and

LDS.X instructions here). It depends on the choice of LDS.X instruction and register blocking factor B_R (Figure 3). For instance, if LDS.64 is used with register blocking factor 6, $F_I = 0.5$.

The throughput factor F_T is a function of register blocking factor (B_R), number of active threads (T_{SM}), throughput of SPs ($\#SP_TP$), LD/ST units ($\#LDS_TP$) and dispatch units ($\#Issue_TP$) (Equation 7). The function f for Fermi and Kepler GPUs is illustrated in Figure 2 and in Figure 4 (only shows LDS.64) and obtained through benchmarks varying these parameters.

$$F_T = f(B_R, \#Issue_TP, \#SP_TP, \#LDS_TP, T_{SM}) \quad (7)$$

With the register blocking factor B_R , the instruction factor F_I and the throughput factor F_T , the performance bounded by SMs' processing throughput is estimated as Equation 8 and the overall performance is as Equation 9.

$$P_{SMBound} = \frac{B_R^2}{B_R^2 + B_R * 2 * F_I} * F_T * P_{theoretical} \quad (8)$$

$$P_{potential} = \min(P_{MemBound}, P_{SMBound}) \quad (9)$$

With the previous analysis, we can estimate the performance upper bound of SGEMM on Fermi and Kepler GPUs. On the Fermi GPU for instance, because of the hard limit of 63 registers (R_{Max}) per thread, considering prefetching and using the strict condition of Equation 4, the maximum blocking factor is only 6. The detailed register allocation is illustrated in Section 5.2. With the register blocking factor of 6, the register resource per SM can support up to 512 threads. Using Equation 3, we choose 256 threads per block.

To easily program the data prefetching, according to Equation 3, L could be 8, 16, 24, ... Considering the condition in Equation 4, we choose L as 16. With a 6-register blocking factor, mixing LDS or LDS.64 with FFMA instructions, the throughput can achieve close to 32 thread instructions per shader clock per SM. Using a LDS.64 instruction can increase the FFMA instruction percentage to 85.7% from 75% (using LDS). Though LDS.128 instruction can provide higher percentage of FFMA instructions, the instruction processing throughput is too low.

According to Equations 6, 8 and 9, the performance is bounded by SMs' processing throughput, and the potential peak is about 82.5% ($\frac{6^2}{6^2+6*2*0.5} * \frac{30.8}{32}$) of the theoretical peak performance for SGEMM. The main limitation comes from the nature of the Fermi instruction set and the limited issue throughput of schedulers.

It is similar to estimate the performance upper bound of SGEMM on Kepler GPU as Fermi GPU. The Kepler GPU (GK104) instruction set is very close to that of Fermi GPU. It means that the limit of 63 registers per thread still exists. Thus, 6-register blocking is also applicable. And the register resource can support 1024 active threads per SM (64K 32bit registers per SM). We can choose either 256 or 1024 threads per block. Similarly, if we use LDS.64 instructions, the FFMA instruction percentage is 85.7%. If

we use LDS.128 instructions (need padding or data layout transform), the FFMA instruction percentage is 92.3%.

Similarly, according to Equations 6, 8 and 9, the performance is bounded by SM's processing throughput, and the potential peak is about 54.6% ($\frac{6^2}{6^2+6*2*0.5} * \frac{122.4}{192}$) of the theoretical peak performance for SGEMM using LDS.64 instructions. Using LDS.128 instructions, the potential peak is about 57.6% ($\frac{6^2}{6^2+6*2*0.25} * \frac{119.9}{192}$) of the theoretical peak. The main limitation factors are still the nature of instruction set and the limited issue throughput of schedulers.

5. Assembly Code Level Optimization

The estimated performance upper bound is a limit that an actual implementation cannot exceed. It can be a little optimistic since we only consider the major performance degradation factors. Besides the considered parameters, there might be other aspects which can limit the performance. The 'real' upper bound or the best possible performance is between the estimated upper bound and the achieved performance.

Depending on whether to apply transpose operation on input matrix A or B, there are 4 variations for GEMM kernel. Figure 5 illustrates the performance of four SGEMM variations from CUBLAS and our implementation (ASM) with 2400x2400 and 4800x4800 matrices. On GTX580 GPU, we achieve around 74.2% of the theoretical peak performance, i.e., about 90% of the estimated performance upper bound, which we think is good enough. In our analysis, we only study the two main instruction types. There are other auxiliary instructions which do not devote to the GFLOPS. And also, we do not consider the effect of barriers which will harm the performance too. We show that the 'real' upper bound is within this 10% and future optimization is unlikely to achieve a lot of speedup. On Kepler GPU, although we cannot provide the optimal controlling information as discussed in section 3.2, we achieve around 77.3% of the estimated upper bound. Similar to Fermi GPU, there are some factors we do not consider in our analysis. The larger gap between our achieved performance might be due to our very limited knowledge of the undisclosed scheduling information of Kepler GPU, which is critical to performance or to some hidden characteristics that we are not able to discover due to limited documentation. Figure 6 illustrates the performance comparison on Fermi GPU between our implementation (assembly), CUBLAS from CUDA 4.1 and MAGMA library [11]. Figure 7 is the performance comparison on Kepler GPU between our implementation (assembly), CUBLAS from CUDA 4.2 and MAGMA library.

The rest of the section briefly describes our optimizations on assembly code level of SGEMM.

5.1 Optimization of Memory Accesses

Assembly code level optimization of memory accesses is similar to high level language optimization. Global memory

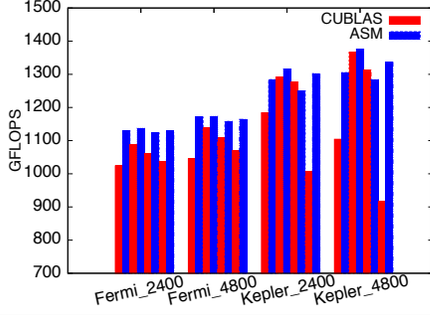


Figure 5. SGEMM Performance of CUBLAS and Our Implementation on Fermi and Kepler GPUs

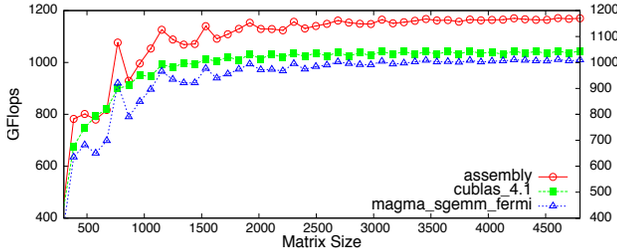


Figure 6. SGEMM_NN Performance on GTX580

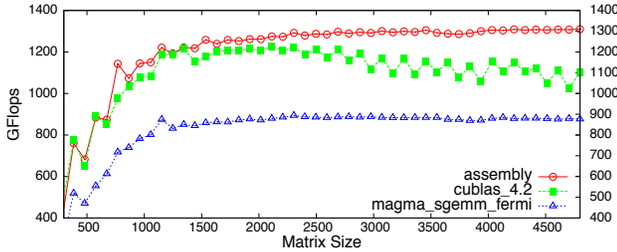


Figure 7. SGEMM_NN Performance on GTX680

requests from the threads within a warp could be grouped (coalesced) into one or more memory transactions depending on the compute capability of the device and the memory accessing pattern. To access global memory efficiently, generally it is better to let threads in a warp access continuous data elements in global memory to get coalescing. Considering the majority of instructions in the SGEMM main loop are FFMA and LDS, and it is critical to reduce the number of LDS instructions (using LDS.64 or LDS.128), sub-matrices in shared memory should be grouped such that each thread accesses continuous B_R data elements. Also, proper padding needs to be applied to reduce shared memory access conflicts and satisfy the alignment restriction of the LDS instruction.

5.2 Register Spilling Elimination

The register resource is 32K 32-bit registers per SM for the Fermi GPU and each thread can use a maximum of 63 registers. The register R1 is normally occupied as stack pointer. According to our analysis, the number of per-thread registers with prefetching is at least $B_R^2 + \frac{2*\sqrt{T_B}*B_R*L}{T_B} + B_R + 1 + R_{index}$. With the register blocking factor of 6 for Fermi GPU, the register allocation of our implementation is

as the following. Note that we use 32bit addressing to save address registers.

1. B_R^2 , 36 registers to save intermediate result for C matrix.
2. $\frac{2*\sqrt{T_B}*B_R*L}{T_B}$, 12 registers to prefetch A and B from global memory.
3. $B_R + 2$, 6 registers to load A from shared memory and 2 registers to load B from shared memory during the main loop. Using 2 registers for B is because LDS.64 instruction is used.
4. 2 registers. Track of A, B in global memory during the prefetching.
5. 1 register to store the loop end condition.
6. 2 registers. Track of A, B in shared memory during the prefetching.
7. 2 registers. Track of A, B in shared memory in the main loop.

In all, 63 registers are used. Since we do not need thread stack, R1 is used to store the loop end condition in our code. Therefore, we are able to fully eliminate the register spilling.

5.3 Instruction Reordering

Generally, we try to interleave different instruction types to get better balance between functional units within one SM and better instruction throughput. We apply the following simple reordering optimizations:

1. In the main loop, between the 2 barriers are all shared memory accesses. By moving address calculation from start of the loop to mix with the shared memory accesses, we can achieve better performance.
2. Interleaving prefetching from global memory with FFMA and LDS instructions can benefit performance.

5.4 Register Allocation for Kepler GPU

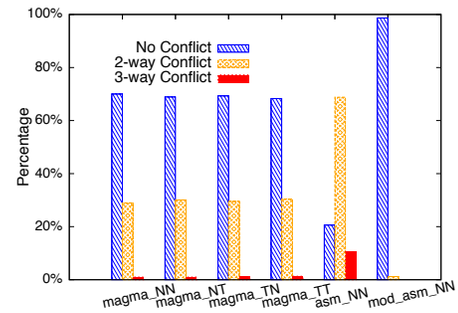


Figure 8. Register Conflict of FFMA Instruction

As we describe in Section 3.3, to get the best throughput, the 3 source registers of FFMA instructions should reside on 3 different banks if they are different. In our current implementation, 6-register blocking is used. 6 registers are used to load A from the shared memory and 2 registers to load B from the shared memory in the main loop. 36 different registers (R26~R61) hold the C sub-matrix. In this implementation, register spilling is eliminated.

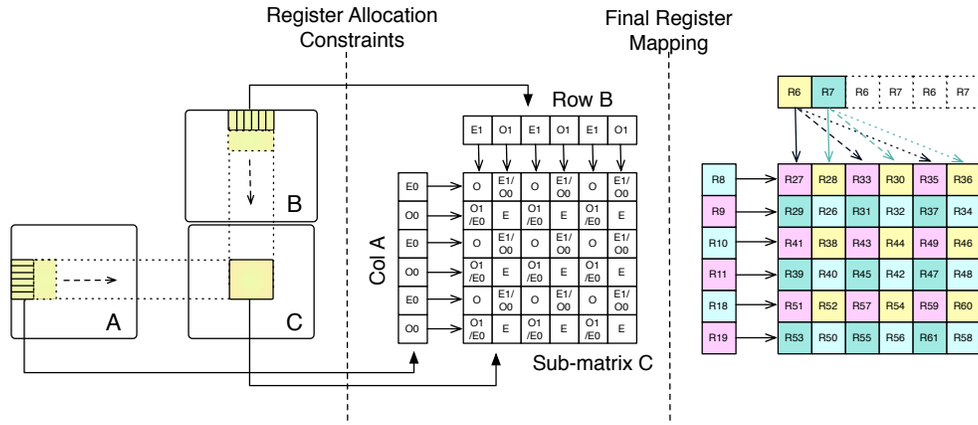


Figure 9. Register Allocation

As in Figure 8, around 30% of the FFMA instructions in the MAGMA [11] SGEMM binary for the Kepler GPU (nvcc generated) have the 2-way register bank conflict and 1% of the FFMA instructions have the 3-way register bank conflict. In our first version of SGEMM_NN on GTX680, which achieves around 1100GFLOPS, 68.8% of the FFMA instructions have the 2-way register bank conflict, and 10.6% of the 3-way conflict. After applying the optimization, the modified version, which achieve around 1300GFLOPS, has only 1.2% of the 2-way FFMA register bank conflict and the 3-way conflict is fully removed.

Our optimization is depicted in Figure 9. In the SGEMM main loop, at each stage, one column from matrix A and one row from matrix B are processed. To use the register blocking and the LDS.64 instructions, at least 6 and 2 different registers are needed for column A and row B. Of course, there are many possible implementations, here we describe one possibility. We select registers from E0 and O0 for column A. Row B uses registers from E1 and O1. Then we use the first table in Figure 9 as the constraints of register allocation. In the final mapping stage, we make sure that 36 registers of C sub-matrix have 9 registers on each bank and had our register allocation as the second table, which does not have any register bank conflict to compute the 36 elements from the C sub-matrix.

5.5 Opportunity for Automatic Tools

Our study emphasizes that for Fermi and Kepler GPUs, it is essential to study the impact of algorithm parameters on instruction throughput to get insight into the performance result. The main optimization opportunity comes from the allocation of registers. For example, the four SGEMM variations of MAGMA library compiled with nvcc spill at least 10 registers (40 Bytes) on the Kepler GPU. When the active thread number is 512, at least 20KB L1 cache is needed to make sure that the spilled data stays in the L1 cache. However, since normally the unified 64KB shared memory/L1 cache is configured as 48KB shared memory and 16KB L1 cache, some data will be spilled out of L1 cache. As the active threads increase, more data is spilled out of L1 cache and the performance will be harmed. We already show that with

careful design, register spilling could be eliminated. We also show that around 30% of FFMA instructions in the nvcc-generated SGEMM binary from MAGMA library have register bank conflict. We propose a simple solution in Section 5.4. It is possible for optimizers to detect the loop structure and remove the conflicts with proper register allocation.

An automatic tuning tool normally needs to explore a large design space and evaluate the performance of many configurations [7, 10, 16]. It may take a significant amount of time. Normally, the automatic tuning tool is application-dependent and each includes several efficient optimizations for the specific application. To build the tool relies on the developers' understanding of the application and optimization experience. With the proposed analysis approach, we can better understand which parameters are critical to the performance. The estimated upper bound actually corresponds to a set of parameters and optimization options. This knowledge can help an automatic tool to explore the design space in a relatively small region. And of course by comparing the performance of an automatic tool's output code and the estimated performance upper bound, we can judge whether the optimized version is good enough.

In our analysis, to study the instruction throughput mixing FFMA and LDS.X instructions, we manually write some benchmarks varying several key parameters such as instruction type choice (LDS.X), the mixing ratio, the blocking factor, the instructions' dependence, active threads and study these parameters' impact on the instruction throughput. For many applications with few major instruction types, a similar approach can be used to estimate the performance upper bound. The difference would be the chosen instruction types and their mixing pattern (mixing ratio, dependence, etc.). Systematic and automatic development of a set of microbenchmarks to help to estimate the performance upper bound of other applications is possible. A family of assembly level microbenchmarks could be defined and evaluated in order to provide a small database of performance references that could be used by the auto-tuning tool, and also the developer to transform the code for performance. Generally, the assembly level microbenchmarks can also help to un-

derstand the difference between different GPU architectures. For example, the benchmarks illustrated in Figure 4 show the increasing need for active threads on Kepler GPU. Assembly level benchmarking requires an assembly tool chain which is missing from the official support. We manage to make it work on Fermi GPU. But on Kepler, there are some issues like the hidden scheduling information, which we cannot fully decrypt.

6. Conclusion

In this work, we have proposed an approach to analyze GPU applications' performance upper bound. Different from existing works on GPU performance models, our approach relies on application analysis and assembly level benchmarking. As an example, we analyze the potential peak performance of SGEMM on Fermi and Kepler GPUs. We show that the nature of the Fermi (Kepler) instruction set and the limited issue throughput of schedulers are the main limitation factors for SGEMM to approach the theoretical peak performance. The general guideline is to reduce the auxiliary instructions and increase the FFMA instruction's percentage. Proper register allocation, shared memory data layout and memory access pattern need to be carefully designed to minimize the impact of memory accesses on performance. We also show that our analysis can help to decide some critical algorithm parameters and show how much optimization space exists. Guided by the analysis, we further optimize the four SGEMM kernel variations and achieve better performance on Fermi GPU (around 5% on average for large matrices) than highly optimized routine provided by NVIDIA.

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