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TSM2X: High-performance tall-and-skinny matrix-matrix multiplication on GPUs

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ABSTRACT

Linear algebra operations have been widely used in big data analytics and scientific computations. Many works have been done on optimizing linear algebra operations on GPUs with regular-shaped input. However, few works focus on fully utilizing GPU resources when the input is not regular-shaped. Current optimizations do not consider fully utilizing the memory bandwidth and computing power; therefore, they can only achieve sub-optimal performance. In this paper, we propose two efficient algorithms - TSM2R and TSM2L - for two classes of tall-and-skinny matrix-matrix multiplications on GPUs. Both of them focus on optimizing linear algebra operation with at least one of the input matrices tall-and-skinny. Specifically, TSM2R is designed for a large regular-shaped matrix multiplying a tall-and-skinny matrix, while TSM2L is designed for a tall-and-skinny matrix multiplying a small regular-shaped matrix. We implement our proposed algorithms and test on several modern NVIDIA GPU micro-architectures. Experiments show that, compared to the current state-of-the-art works, (1) TSM2R speeds up the computation by 1.6x on average and improves the memory bandwidth utilization and computing power utilization by 18.1% and 20.5% on average, respectively, when the regular-shaped matrix size is relatively large or medium; and (2) TSM2L speeds up the computation by 1.9x on average and improves the memory bandwidth utilization by up to 9.3% on average when the regular-shaped matrix size is relatively small.

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1. Introduction

Matrix-matrix multiplication (GEMM) has been one of the most extensively used linear algebra operations in big data analytics and scientific computations. Due to many factors (such as algorithms, input data, etc.) the size or shape of input matrices for GEMM usually varies when it is used in different applications. For example, many modern highly scalable scientific simulation packages in the field of fluid dynamics, such as Finite Element Method (FEM) simulations, need to compute many GEMMs with small-sized input matrices. Artificial neural networks (ANN) involve using GEMM with small to medium input matrices. Matrix decompositions uses GEMM with large-sized input matrices

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https://doi.org/10.1016/j.jpdc.2021.02.013 0743-7315/© 2021 Elsevier Inc. All rights reserved. [7,22,27,28]. Thus, besides large-sized input, which has already been extensively optimized during the past decades, GEMM with small to medium sized input has also drawn much attention to recent researchers. For instance, Dong et al. [15] proposed MAGMA-Batched, which aims to batch small input matrices into larger ones in order to utilize the highly optimized implementations for large input size on GPUs. Heinecke et al. [18] proposed to speed up GEMM with small input using architecture and instruction level optimization on modern CPU architectures.

Although previous works have focused on optimizing GEMM with different matrix sizes, most of them only assume that the input matrices are *regular-shaped*. In other words, the size mentioned in their works usually refers to both dimensions of the input matrix. For example, a small matrix means both of its width and height are small and their magnitudes are close to each other. When the dimensions of the input matrices have significant difference, we consider them as *irregular-shaped* inputs. In particular, many irregular-shaped inputs involve *tall-and-skinny*

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matrices, in which their widths are significantly smaller then their heights. Although few works have been done to study and optimize GEMM with tall-and-skinny input, this input case has been widely used in many applications [8]. For instance, recent highly optimized K-means implementations [14,20] use GEMM as their core computation, and the input size is mostly tall-andskinny. This is because the number of centroids is usually far less than the number of input data points. Moreover, when GEMM is used for encoding checksums for many algorithm-based fault tolerance applications [3–6,19,21,29,30,36–38], the input involves a tall-and-skinny checksum weight matrix.

Previous efforts made for optimizing GEMM with regularshaped input may not work for non-regular shaped input. For instance, Chen et al. [5] illustrate that calculating GEMM with tall-and-skinny input using the vendor's highly optimized linear algebra library (e.g., cuBLAS [2]) is slower than disassembling the tall-and-skinny input matrix into several vectors and then applying matrix-vector multiplications. However, it can be easily seen that even with this workaround the computation is not efficient, since elements in input matrices are accessed by the GPU more times than necessary. Although the performance can be optimized by grouping many tall-and-skinny input matrices into large ones similar to the approach proposed, there are cases where this grouping approach is not feasible. For example, talland-skinny input matrices may be generated one at a time from a producer process in user's workflow. Grouping several of them into a large matrix requires extended waiting time, which is not applicable for time-sensitive applications. On the other hand, the memory space may limit the total number of matrices that can fit into the memory at the same time, if the input matrices are large (e.g., multiplication of regular-shaped large and tall-and-skinny matrices).

In this work, we target on optimizing the computation of GEMM with tall-and-skinny input on the GPU platform since many applications that use GEMM are deployed on GPUs. So, our optimization greatly benefits those applications. The key insight of our work is that the computation characteristic of GEMM on modern computing systems is not always unchanged as we change the shape of input matrices. For example, when the sizes of regular-shaped matrices are large (i.e., $m \simeq k \simeq n \gg 1$ for an $m \times k$ matrix multiplying an $k \times n$ matrix), the compute-to-load ratios of each element in the input matrices are $O(m) \approx O(n)$. So, the regular-shaped GEMM operations are usually computebound especially for large matrices. However, when the input is tall-and-skinny (i.e., $m \simeq k \gg n$ or $m \gg k \simeq n$), the average compute-to-load ratio is reduced to around O(1). Moreover, when k is very small (i.e., $m \gg k \simeq n$), each GPU thread would not perform enough workload to hide latency and hence low occupancy. Therefore, depending on the relationship between m, k, and n, and the performance characteristics of GPUs, the computation can be compute-bound, memory-bound, or latencybound. Specifically, when (1) $m \simeq k \gg n$, as *n* gets larger, it moves toward compute-bound; (2) $m \simeq k \gg n$, as n gets smaller, it moves toward memory-bound; and (3) $m \gg k \simeq n$, it moves toward latency-bound. To optimize GEMM with tall-andskinny input, it is critical to design a computation algorithm that considers all compute-bound, memory-bound, and latency-bound cases.

The main contributions of this paper include:

• We study the limitation of current state-of-the-art GEMM implementations with tall-and-skinny inputs (i.e., $m \simeq k \gg n$ or $m \gg k \simeq n$). With benchmarking, we find that the under-utilization of GPU resources is the main reason for performance degradation when the input is tall-and-skinny.

- To handle a broad spectrum of tall-and-skinny inputs for GEMM on GPUs, we design two classes of algorithms with optimizations focusing on different tall-and-skinny input cases: (1) TSM2R is designed to handle a large regular-shaped matrix multiplying a tall-and-skinny matrix (i.e., $m \simeq k \gg n$); (2) TSM2L is designed to handle a tall-and-skinny matrix multiplying a small regular-shaped matrix (i.e., $m \gg k \simeq n$).
- We present a performance model for TSM2R and compare it with our evaluation performance results. Moreover, we examine the inadequacies of the model for TSM2L and further improve it based on our observations.
- We carefully implement TSM2R and TSM2L using CUDA C² and evaluate them on four generations of NVIDIA GPUs including Kepler, Maxwell, Pascal, and Volta. Experiments show that our TSM2R and TSM2L can achieve 1.6x and 1.9x speedups, respectively, on average with different tall-and-skinny inputs, compared to the state-of-the-art GEMM library cuBLAS.

The rest of this paper is organized as follows. In Section 2, we give a formal definition of tall-and-skinny matrix and show some preliminary benchmark results of the GEMM with tall-and-skinny matrix using cuBLAS. In Section 3, we propose our detailed design of TSM2R and TSM2L for two different kinds of tall-and-skinny inputs. In Section 4, we present our evaluation results. In Section 5, we examine related works for tall-and-skinny inputs. In Section 6, we conclude the paper.

2. Background

2.1. Tall-and-skinny input for GEMM

In this work we restrict our scope to handle irregular-shaped inputs that involve tall-and-skinny matrices. The tall-and-skinny input size means that, for the two input matrices, at least one matrix is tall-and-skinny (i.e., one dimension is significantly smaller than the other). For example, either (i) input matrix A with size 20480×20480 and matrix B with size 20480×2 or (ii) input matrix A with size 20480 \times 2 and matrix B with size 2 \times 2 is considered as tall-and-skinny input in our work. Tall-and-skinny matrices are a typical class of matrices that can be found in irregular-shaped inputs for GEMM. In this paper, we focus on optimizing GEMM with (i) one large regular input matrix and one tall-and-skinny input matrix and (ii) one tall-and-skinny input matrix and one small regular input matrix. In this paper, for the first case, we let matrix A be the larger input matrix $(m \times k)$ and matrix B $(k \times n)$ be the tall-and-skinny input matrix, where $m \simeq k \gg n$; for the second case, we let matrix A be the talland-skinny input matrix $(m \times k)$ and matrix B $(k \times n)$ be the smaller input matrix, where $m \gg k \simeq n$. We choose these input sizes and shapes because we believe they can expose most of the challenges in processing all kinds of tall-and-skinny input, so the design idea and optimization techniques introduced in this paper can be easily applied to other cases with slight modification. Also, for simplicity's sake, we choose to let the larger matrix in (i) and smaller matrix in (ii) to be square-shaped in most of our experiments. Our optimization can work with non-square input as well with similar effects.

² The TSM2X code is available at https://github.com/codyjrivera/tsm2x-imp.

2.2. cuBLAS

One of the most commonly used standard linear algebra libraries optimized for the GPU is the cuBLAS library developed by NVIDIA. cuBLAS is the core computing library of many big data and scientific computing applications. For example, it is the GPU computing library for MAGMA heterogeneous linear algebra library [16,31,32], cuLA library [12], and cuDNN deep learning library [11]. With NVIDIA's deep optimization, the cuBLAS library is able to provide state-of-the-art performance in many use cases. For example, with large regular-shaped input matrix, their GEMM implementation is able to achieve near peak GPU performance [9].

However, we found that the GEMM subroutine is not fully optimized with certain input matrix sizes [6]. For example, with inputs that involve tall-and-skinny matrices, the GEMM operation in current best implementation (cuBLAS 9.0 running on NVIDIA Tesla K40c GPU) uses less than 10% of the theoretical peak memory bandwidth on average with n = 2 (as demonstrated in Fig. 7(a)–(b)). When n = 16, the same GEMM operation uses less than 20% of the theoretical peak memory bandwidth on average (as demonstrated in Fig. 7(g)–(h)). The resource utilization is even lower with larger input dimensions. By comparing the two input sizes, it can be seen for input with smaller *n* values, the computation utilizes higher memory bandwidth (close to memory bound). On the other hand, for input with larger n values, the computation utilizes higher computing power (close to compute bound). However, since we are unable to analyze the GEMM implementation in the closed-source cuBLAS library, it is hard to tell its exact computational characteristics.

3. Design methodologies

To handle the GEMM with two different classes of tall-andskinny inputs on GPUs described in Section 2.1, we design two efficient algorithms: TSM2R and TSM2L. TSM2R is designed to handle inputs with one large-to-medium regular-shaped matrix and one tall-and-skinny matrix, while TSM2L is designed to handle inputs with one tall-and-skinny matrix and one small regular-shaped matrix. Note that "R" or "L" means that the talland skinny matrix is multiplied on the right or left.

3.1. Design of TSM2R

In this section, we describe our proposed algorithm TSM2R for GEMM with a large regular-shaped matrix and a tall-and-skinny matrix.

3.1.1. Insight on tall-and-skinny input

For regular-shaped GEMM ($m \times k$ matrix multiplies $k \times n$ matrix), the input matrices' total size is O(mk + kn), while the computing time complexity is O(mkn), so each element in the input matrices is used O(m) or O(n) times within the entire computation process. Since loading data to the GPU from the off-chip DRAM (i.e., global memory) to GPU is expensive and to avoid extensive data load operations, one common optimization for this kind of problem is minimizing the number of times each element needs to be loaded into the GPU by using fast on-chip memory (e.g., cache, registers) to enable data reuse. As the number of loads reduces, optimized GEMM tends to be compute-bound. For example, current GEMM implementation in cuBLAS library can reach near bare-metal performance on GPUs [9].

However, unlike regular-shaped GEMM, when one matrix is tall-and-skinny (e.g., $n \ll m, k$), each element in the input matrices is used O(n) times on average:

$$\frac{(m \times k) \times n \text{ times} + (k \times n) \times m \text{ times}}{m \times k + k \times n} \approx O(n) \text{ times.}$$

Require: input matrix A $(m \times k)$ and B $(k \times n)$, output matrix C $(m \times n)$ 1: **for** i = 1 to n **do**

2: **for** j = 1 to k **do**

3: $C[global_tid, i] + = A[global_tid, j] \times B[j, i]$

4: end for

5: **end for**

Algorithm 1: Each thread's workload with inner product.

Depending on the size of n and target GPU peak computing power and memory throughput ratio, the computation can be either compute-bound or memory-bound. When n gets smaller, the computation tends to be memory-bound. Otherwise, the problem tends to be compute-bound. In either case, the problem is always near the boundary between memory bound and compute bound, so it is critical to design an algorithm that is optimized for both cases.

3.1.2. Algorithm design

Algorithm design plays a critical role in our proposed optimizations. First, we need to consider how to fit the workload of our TSM2R into the programming model of CUDA (i.e., thread hierarchy). Although the workload can be easily decomposed into many independent smaller workloads, careful consideration of the workload distribution is still necessary, since any unnecessary performance penalty can cause drastic GPU resource under-utilization. Several factors are considered in our design:

- 1. Total number of global memory accesses;
- 2. Shared and global memory access efficiency;
- 3. Utilization of overall memory bandwidth;
- 4. Parallelism of overall workload;
- 5. On-chip memory utilization;
- 6. Streaming Multiprocessor (SM) utilization;
- 7. Optimization for compute & memory-bound cases.

To achieve good performance, there must exist enough active threads in each SM of the GPU to ensure proper instruction and memory access latency hiding. So, in our algorithm we divide the workload by assigning *n* rows of matrix A to *n* different threads. Each vector-matrix multiplication is assigned to one thread (i.e., $(A[i, :] \times B))$). The benefit is three-fold: (1) this ensures high parallelism and high SM occupancy; (2) since the number of elements of matrix A is much higher than matrix B, this kind of distribution ensures that matrix A is accessed as little as possible; (3) it also enables high memory access efficiency and throughput, since all memory accesses to matrix A are naturally coalesced (assuming matrices are stored in column-major by convention).

As for the vector-matrix multiplication assigned to each thread, to further reduce the number of memory accesses to matrix A, we use outer-product style computation instead of the usual inner-product style computation. As shown in Algorithm 1, if we use inner-product, each element of matrix A is repeatedly referenced *n* times. On the other hand, if we use outer-product as shown in Algorithm 2, each element of matrix A is referenced only once. (Please note, as we will discuss in later sections, when *n* is larger than a certain threshold, elements in matrix A still need to be referenced more than once due to the limited resources available for each thread, but it is still far lower than using innerproduct.) When matrix A is large, the benefit is significant, since it greatly reduces the total number of global memory accesses during the entire GEMM computation. Also, the outer-product style does not bring any extra memory accesses to matrix B compared to inner-product style. The only cost for outer-product is extra registers holding *n* intermediate results. However, with proper tuning, the benefit of fewer memory accesses outweighs this cost.

Require: input matrix A $(m \times k)$ and B $(k \times n)$

Require: output matrix C $(m \times n)$

1: $Reg_{1:n} \leftarrow C[global_tid, 1:n]$

- 2: **for** i = 1 to k **do**
- $Reg_{1:n} + = A[global_tid, i] \times B[i, 1:n]$ 3:

```
4: end for
```

5: $C[global_tid, 1:n] \leftarrow Reg_{1:n}$

Algorithm 2: Each thread's workload with outer product.

3.1.3. Efficient off-chip memory access

One key factor of optimizing memory intensive applications is ensuring high off-chip memory access efficiency. Depending on the GPU model type or runtime configurations, global memory (off-chip) accesses of threads within the same warp can be coalesced into 128 byte- or 32 byte-transactions [10] if their access addresses fall into the same 128 byte- or 32 byte-segments in global memory, which enables efficient use of memory bandwidth. Otherwise, the GPU still loads memory in 128 byte- or 32 byte-transactions, but it may contain unrequested data that are stored in neighbor addresses, which causes inefficient memory accesses.

Since each thread reads one row of matrix A and the matrix is stored in column-major format by convention, memory accesses are naturally coalesced when threads within the same warp access elements on different rows but on the same column. So, 100% memory access efficiency is achieved on matrix A. However, for matrix B, all threads access the same element at the same time, which results in a single memory transaction containing one requested element and several unrequested neighbor elements. So, only $\frac{8 \text{ bytes}}{128 \text{ bytes}} = 6.25\%$ or $\frac{8 \text{ bytes}}{32 \text{ bytes}} = 25\%$ memory access efficiency is achieved for accessing 64-bit double floating point elements. Although the total number of elements in matrix B is small, given that each element needs to be accessed *n* times, this inefficient access pattern can still greatly impact the overall performance.

To improve the efficiency of memory accesses to matrix B, we utilize shared memory in GPU. Since it is located on-chip, shared memory gives us the speed of L1 cache and it is fully programmable. Threads within one thread block can use shared memory to share data. So, one key advantage of shared memory is that it eliminates the need for the consistency between patterns of data loading and data using pattern, which enables us to load global memory in the most efficient way and keep the way that we use data as before.

By using shared memory for accessing matrix B, we can reduce the total number of memory accesses and enable coalesced memory access. As shown in Algorithm 3, for each iteration, instead of letting threads request elements they need individually by themselves inefficiently, we now let a block of threads work together to fetch a tile of matrix B into the shared memory in a coalesce-compatible way (Line 11). Then during computation, each thread references elements in matrix B through the shared memory instead of loading each one of them individually from global memory. This reduces the total number of accesses to matrix B from global memory (from *n* to n/t_1 per element). Also, threads in a same thread block fetch elements of matrix B column by column, which enables coalesced memory access and greatly improves memory-access efficiency to 100%. Moreover, we also introduce three parameters: t_1 , t_2 , and t_3 in Algorithm 3. These parameters are used for adjusting the performance and will be discussed in later sections.

3.1.4. Optimizing use of shared memory

Although fast, elements in shared memory still need to be loaded into registers before using them [26]. Its access speed can **Require:** input matrix A $(m \times k)$ and B $(k \times n)$, output matrix C $(m \times n)$

- 1: $t_1 \leftarrow tile_size_B, t_2 \leftarrow tile_size_C, t_3 \leftarrow tile_size_A$
- 2: Register: A₁, A₂, ..., A_{t₃} 3: Register: $C_1, C_2, ..., C_{t_2}$
- 4: Shared Memory: curr**B** with size $t_1 \times t_2$
- 5: Threads per thread block $\leftarrow t_1$
- 6: Total thread blocks $\leftarrow m/t_1$
- 7: for p = 1 to *n* with step size = t_2 do
- $C_{1:t_2} \leftarrow C[global_tid, p: p + t_2 1]$ 8:
- ٩· **for** i = 0 to k with step size = t_1 **do**
- /* Load a tile of B into shared memory */
- 10: ThreadsSynchronization() *currB*[*global_tid*, 1 : t_2] \leftarrow *B*[*j* + *global_tid*, *p* : *p* + t_2 - 1] 11:
- ThreadsSynchronization() 12:
- 13: **for** l = i to $i + t_1$ with step size = t_3 **do**
- /* Load a tile of A into registers*/
- $A_{1:t_3} \leftarrow A[global_tid, l: l + t_3 1]$

```
14:
```

```
C_{1:t_2} + = A_{1:t_3} \times currB[l: l + t_3, 1: t_2]
15:
           end for
```

```
16:
```

```
17:
     end for
```

18: $C[global_tid, p: p + t_2 - 1] \leftarrow C_{1:t_2}$

```
19: end for
```

Algorithm 3: TSM2R with shared memory.

affect the overall performance. Shared memory is divided into several same-sized memory banks for fast parallel accesses. Different threads can access different memory banks simultaneously. So, having a total of *b* memory banks can speedup overall shared memory throughput by up to *b* times compared to the throughput of one single memory bank. However, if x threads in the same warp access different data from the same memory bank, an *x*-way bank conflict occurs and each request is processed sequentially, which dramatically reduces the accessing throughput by a factor of 1/x.

In our algorithm, threads in the same thread block load data from global memory into shared memory column by column to enable fast coalesced global memory access. Then threads access data from shared memory row by row during computation. How we store elements in shared memory will affect how these elements are accessed from memory banks, which affects the throughput of shared memory. We have two ways of storing a tile of matrix B in shared memory: column-major storage and row-major storage. To choose between the two ways, we need to analyze and compare which way brings the least overall bank conflict. We assume the size of one tile of matrix B is $t_1 \times t_2$ and t_1 is the multiply of total number of memory banks *b* for simplicity.

For column-major storage, elements (32-bit words or 64-bit words) in the same column of one tile of matrix B are stored in successive memory banks. So, for shared memory with b memory banks, t_1 elements of one column are stored in *b* different successive memory banks with each bank storing at most $\frac{t_1}{b}$ elements and being accessed by at most $\frac{warp \ size}{b}$ threads at the same time, which may potentially cause bank conflict if $\frac{warp \ size}{b}$ is greater than one.

For row-major storage, elements in the same row of matrix B are stored in successive memory banks. So, elements of the same column are stored in $\frac{b}{t_2}$ different banks, where each bank stores $\frac{t_1 \times t_2}{h}$ elements from one column. Since each bank has t_2 times more elements from one column, each bank has at most t_2 times more threads accessing it at the same time: $\frac{warp \ size}{k} \times t_2$, which may also potentially cause bank conflict.

On modern NVIDIA GPUs, the warp size is fixed to 32 and total number of banks is also 32 [10], so column-major storage does not cause bank conflict, since each bank can only have up to one thread accessing. Row-major storage can cause up to t_2 -way bank conflict, which decreases overall shared memory

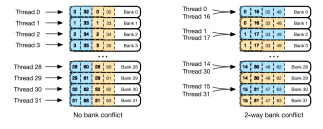


Fig. 1. Comparing column-major (left) with row-major (right) storage for storing a 64 imes 2 tile of matrix B in shared memory. Blue and yellow squares represent elements in the first and second column. When one warp of 32 threads accessing 32 elements in one column (e.g. element 0 to 31 of the first column), the column-major storage brings no bank conflict and row-major storage brings 2-way bank conflict, which reduces throughput by half. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

throughput to $\frac{1}{t_2}$ of the peak throughput. As shown in Fig. 1, we load a 64×2 matrix tile into shared memory using column-major storage (left) and row-major storage (right). When using columnmajor storage, threads in one warp all access different banks, so no bank conflict occurs. But when using row-major storage, 32 elements are stored in 16 banks causing 2-way bank conflict. When accessing elements in shared memory for computation, threads in a warp all access the same element at the same time in our algorithm. Although multiple threads are accessing one bank, they are accessing the same element, so one broadcast is initiated, which does not cause bank conflict. It is the same for both storage styles. So, we choose column-major storage as it brings no bank conflict and potentially brings the highest shared memory throughput.

3.1.5. Overlapping computation and memory access latency

During execution, for each instruction issuing moment, each warp scheduler picks an eligible warp and send it to the corresponding component for execution. A warp becomes eligible only if all operands of its next instruction are ready. However, if a warp is loading data from global memory, it will take several hundred cycles before it can be ready for execution. To hide this long latency, we can either increase the number of threads residing in each SM to ensure there always exist eligible warps [33] or put independent instructions in between data loading and data consuming operations, so that warps are also eligible for execution during memory loading time. The first approach requires us to adjust the on-chip resource usage of each thread block. We will save that discussion for the next section. In this section, we aim to add independent instructions in between data loading and data consuming operations.

A shown in Algorithm 3, Line 11 and 14 load data from global memory and Line 15 consumes data once data is loaded. However, due to data dependency, there is no independent instruction in between, so once each warp issues global memory access requests, it must wait for the requested elements to be ready before it can proceed to computation.

So, to add independent instructions, we use data prefetching to mix data loading and consumption between neighbor iterations. Specifically, instead of letting each iteration loads data that is going to be used for current iteration, we let the data needed for current iteration to be loaded by the previous iteration, so that its calculation will not be blocked by data loading (since the data are ready). When doing calculation, it also loads data that is going to be used for the next iteration. By overlapping data loading and computation, we can significantly improve memory bandwidth and SM utilization. We apply data prefetching to both matrix A and B.

Require: input matrix A $(m \times k)$ and B $(k \times n)$, output matrix C $(m \times n)$

- 1: $t_1 \leftarrow tile_size_B, t_2 \leftarrow tile_size_C, t_3 \leftarrow tile_size_A$
- 2: Register: currA₁, currA₂,...,currA_{t3}
- 3: Register: nextA₁, nextA₂,...,nextA_{t3}
- 4: Register: nextB₁, nextB₂,...,nextB_{t2}
- 5: Register: $C_1, C_2, ..., C_{t_2}$
- 6: Shared Memory: *currB* with size $t_1 \times t_2$
- 7: Threads per thread block $\leftarrow t_1$
- 8: Total thread blocks $\leftarrow m/t_1$
- 9: for p = 1 to n with step size = t_2 do
- 10: $C_{1:t_2} \leftarrow C[global_tid, p: p + t_2 - 1]$
- /*load the first tile of A and B*/
- $currB[local_tid, 1:t_2] \leftarrow B[local_tid, p:p+t_2-1]$ 11.
- $currA_{1:t_3} \leftarrow A[global_tid, 1:t_3]$ 12:
- 13. **for** i = 0 to k with step size = t_1 **do**
- ThreadsSynchronization() 14:
- /*prefetch the next tile of B into registers*/
- if $i + t_1 < n$ then 15:
- 16: $nextB_{1:t_2} \leftarrow B[j + t_1 + local_tid, p : p + t_2 - 1]$
- 17: end if
- **for** l = j to $j + t_1$ with step size = t_3 **do** 18:
- /*prefetch the next tile of A into registers*/
- 19. **if** $l + t_3 < n$ **then**
- $nextA_{1:t_3} \leftarrow A[global_tid, l + t_3 : l + t_3 + t_3 1]$ 20:
- 21: end if
- $C_{1:t_2} + = currA_{1:t_3} \times currB[l: l + t_3, 1: t_2]$ 22:
- /*load the prefetched tile of A from nextA registers into currA registers*/
- 23:
- $currA_{1:t_3} \leftarrow nextA_{1:t_3}$ 24: end for
- 25:
- ThreadsSynchronization() /*load the prefetched tile of B from nextB registers to shared memorv*/
- 26 $currB[local_tid, 1:t_2] \leftarrow nextB_{1:t_2}$
- 27: end for
- 28. $C[global_tid, p: p + t_2 - 1] \leftarrow C_{1:t_2}$
- 29: end for

Algorithm 4: TSM2R with shared memory and data prefetching.

LD C	Threads Sync.	Threads Sync.	Threads Sync.	ST C
	LD NextB	LD NextB	LD NextB	
LD	NextA LD NextA	LD NextA LD NextA	LD NextA LD NextA	
Co	mpute Compute	Compute Compute	Compute Compute	

Fig. 2. Example workload of one iteration of our optimized TSM2R with data prefetching.

As shown in Algorithm 4, we design our TSM2R with data prefetching. Note that global_tid and local_tid represent the (global) thread ID in the grid and the (local) thread ID in the block, respectively. In Line 2 and 3, we allocate two sets of t_3 registers for storing current tile of elements of matrix A and next tile of element of matrix A for prefetching. In Line 4 and 6, we allocate t_2 registers for data prefetching of elements in matrix B, and allocate $t_1 \times t_2$ for storing currently loaded tile of matrix B. Note that we cannot store current tile of matrix B in registers, because elements in matrix B need to be shared between threads during computation.

Before the core computation iteration (Line 13-27), we preload current tile of matrix A and B into registers and shared memory (Line 11 and 12), so that computation can start immediately as soon as we enter the computation loop without being blocked by any data dependency. The main computation resides in Line 22. To overlap computation with memory accesses, we initiate loading for the next tile before the computation (Line 16 for matrix B and Line 20 for matrix A). We use two loops for loading matrix A and B, because we want to have the flexibility to adjust loading pace (tile size) differently for the two matrices. We will discuss this in the next subsection. Figs. 2 and 3 show one iteration of our optimized TSM2R with data prefetching. LD C and ST C represent loading initial values from matrix C and

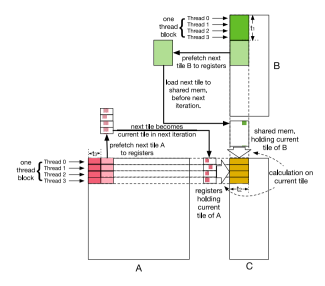


Fig. 3. Matrix view of tall-and-skinny matrix-matrix multiplication with data prefetching.

storing final results back to matrix C. Each iteration we show three sub-iterations for loading matrix B. As we can see, we compute and pre-load the next tile of matrix B concurrently to improve memory bandwidth utilization. A thread barrier is inserted in the end of each iteration. For the innermost iteration, we do the actual computation and pre-load elements from matrix A each time. Please note that the length of each rectangle does not accurately represent the exact execution time length and the ratio between number of LD nextA and LD nextB is not necessarily two in actual computation. Also, we show one thread block with four threads only for illustration proposes. As we will discuss in the next subsection that different parameter values can affect the length of each part and the ratio between number of LD nextA and LD nextB. Especially on the execution time of LD nextA and Compute, which will affect the characteristic of computation (i.e. memory-bound or compute-bound). Also, for simplicity, we ignore the data movement from next tile to current tile that occurs in each iteration.

3.1.6. Parameters definition

In Algorithms 3 and 4, we introduced three adjustable parameters: t_1 , t_2 , and t_3 . In this section, we first discuss how each parameter controls the computation of our TSM2R. Then, we introduce our performance model that estimates how certain performance metrics change with these parameters. Finally, we explain our strategies for choosing values for these parameters in order to achieve high GPU resource utilization and optimize overall performance. Please note that the following discussions are all based on Algorithm 4.

3.1.7. Behaviors of parameters

We first list the behaviors of each parameter below:

• t_1 specifies the number of rows of one tile of matrix B. To maximize use of available active threads and to avoid any inefficient thread execution caused by warp divergence, we let all threads in each thread block participate in fetching elements of matrix B. For fast coalesced global memory access, we let each thread fetch one row, so t_1 is also the total number of threads in each thread block. Also, since we let a total of *m* threads work on the computation, the total number of thread blocks can be calculated as: m/t_1 .

- *t*₂ specifies the number of elements in matrix C that each thread is working on at a time. It is used to divide the overall workload into several smaller workloads that are processed iteratively by each thread. A smaller workload makes each thread's SM resource usage smaller, which allows us to keep higher SM occupancy. However, dividing the workload means we need to load matrix A repeatedly for each small workload. So, there is a trade-off. *t*₂ also affects the ratio between total number of memory fetches and computation operations in core part of our algorithm, which allows us to adjust the computation to be compute or memory-bound (will be discussed later in detail).
- *t*₃ specifies the number of elements in matrix A that each thread fetches at a time. Since elements fetches are independent to each other, they can be done without blocking each other, so *t*₃ can be used to adjust the memory loading concurrency.

3.1.8. Performance metrics estimation

In this section, we introduce our parameter-based performance model that is used to estimate three important performance metrics: SM occupancy, memory bandwidth utilization and computing power utilization. These estimations will be used for optimizing the overall performance.

• Max SM occupancy estimation

With these parameters we can calculate the max occupancy of each SM, which is defined as max number of active threads per SM. (Some works also use max number of warps, which is similar to ours. We found that using the maximum number of threads is more consistent across our performance models. We also choose our thread block size to be the dividend of this value to ensure the expected number of threads are active.) This occupancy is mainly bound by the maximum hardware allowable number of threads (HW_MAX) and on-chip memory utilization per thread. We first calculate the total number of registers utilized per thread. Since register utilization can potentially be optimized by the nvcc compiler, we use the maximum number of registers to estimate this value. First of all, there is a relatively fix amount of registers used for CUDA initial setup, and we represent this amount as C. We get its amount through offline profiling. Then, we need two sets of t_2 registers for storing elements of matrix B for both next tile fetching and current tile calculation. Please note that although the current tile of matrix B is stored in shared memory, it still needs to be transferred to registers for calculation. Next, we need t₂ registers for keeping intermediate results of matrix C. Finally, we need two sets of t₃ registers for storing elements of matrix A for both next tile fetching and current tile calculation. So, the total number of registers is:

$$R_{thread} = (t_2 \times 3 + t_3 \times 2) \times \frac{bytes_per_element}{bytes_per_register} + C.$$

As for shared memory, although it is allocated per thread block, we calculate the average amount of shared memory that each thread uses for consistent calculation here. Since the size of allocated shared memory per thread block is $t_1 \times t_2$, and as we will discuss earlier that we set $t_1 = threads_per_threadblock$, the amount of shared memory allocated for each thread on average is:

 $S_{thread} = t_2 \times bytes_per_element.$

So, the max SM occupancy can be calculated as:

$$MaxOccup_{SM} = min(HW_MAX, \frac{R_{SM}}{R_{thread}}, \frac{S_{SM}}{S_{thread}}).$$

• Max memory bandwidth utilization estimation

Next, we estimate the max memory bandwidth utilization of our algorithm when the computation is memory-bound. In this case, loading elements of matrix A dominates the computation instead of floating point calculations in our algorithm. So, we can estimate max memory bandwidth utilization using the maximum number of concurrent global memory accesses per SM. It can be calculated as:

$Concurrent_{mem} \approx MaxOccup_{SM} \times t_3.$

Note that we only consider the memory accesses to matrix A here for simplicity. Since the majority of memory accesses are for matrix A, this only brings minor inaccuracy. Then, similar to [33,35] we calculate the least number of concurrent memory accesses per SM needed to achieve max memory bandwidth utilization using Little's Law:

$$Throughput_{max_mem} = \frac{Peak Band.}{\#_of_SM \times core_clock},$$

 $Concurrent_{max_mem} = latency_{mem} \times Throughput_{max_mem}$.

The $latency_{mem}$ is the average global memory access latency, which is considered as a constant in our model and is obtained through offline profiling. The estimated memory bandwidth utilization is:

$$Util_{mem} = \frac{Concurrent_{mem}}{Concurrent_{max_mem}}.$$

• Max computing power utilization estimation

Next, we estimate the max computing power utilization of our algorithm when the computation is compute-bound. In this case, floating point calculation dominates the computation instead of memory accesses in our algorithm. So, we can estimate max computing power utilization using the maximum number of concurrent floating point operations per SM. It can be calculated as:

$Concurrent_{comp} = MaxOccup_{SM} \times t_3 \times t_2.$

Then, also similar to [33] we calculate the least number of concurrent floating point operations per SM needed to achieve max computing power utilization using Little's Law:

$$Throughput_{max_comp} = \frac{Peak Perf.}{\#_of_SM \times core_clock},$$

 $Concurrent_{max_comp} = latency_{comp} \times Throughput_{max_comp}.$

The $latency_{comp}$ is the average latency of floating point operations in our calculations, which is considered as a constant in our model and is obtained through offline profiling. So, the estimated computing power utilization is:

$$Util_{comp} = \frac{Concurrent_{comp}}{Concurrent_{max\ comp}}$$

• Determine compute-bound or memory-bound

Given parameters and GPU specification, we can determine whether the current computation is memory or computebound. This is mainly determined by the innermost loop (Line 20–24) of Algorithm 4. The memory loading instructions (Line 21) overlap the computation (Line 23). Since Line 24 depends on memory loading results, it serves as an implicit synchronization point for memory loading and computation. The time taken for the two parts will determine whether the current computation is compute-bound or memory-bound. So, we first estimate the time taken for computation and memory access as follows:

$$time_{comp} = \frac{t_3 \times t_2}{Peak \ Perf. \times \#_of_SM \times Occupancy_{SM}}$$

 $time_{mem} = \frac{t_3 \times bytes_per_elem.}{Peak \ Band. \times \#_of_SM \times Occupancy_{SM}}$

Then, by comparing the two time costs, we can determine whether the current computation is compute-bound or memory-bound.

$$r = \frac{time_{comp}}{time_{mem}} = \frac{t_2}{bytes_per_elem.} \times \frac{Peak Band.}{Peak Perf.}$$

As we can see, when r is greater than one, the computation is compute-bound. Otherwise, the computation is memorybound. Also, since we divide the original workload into several smaller workloads using t_2 , this ratio is determined by t_2 . By adjusting t_2 , the actual computation can be shifted between compute and memory-bound. The boundary between the two cases can be calculated by setting the ratio r = 1, so we get a threshold for t_2 :

$$t_2^{\text{threshold}} = \frac{\text{Peak Perf.}}{\text{Peak Band.}} \times \text{bytes_per_elem.}$$

Similarly, we can also estimate the computation characteristics of the original problem, in which the workload is not divided into smaller workloads. In this case, t_2 is always fixed to k. So, by comparing k with $t_2^{threshold}$ we can estimate the computation characteristics. If k is greater than $t_2^{threshold}$, the original problem is compute-bound; otherwise, it is memory-bound. It can be easily seen, depending on the value of t_2 and k, the computation characteristics of the current problem and original problem can be different, which can affect the overall performance. We discuss this in later part of this section.

3.1.9. Deciding parameters

When choosing parameters, the first thing we should determine is whether we should optimize for computation or memory bandwidth. This is determined by whether the given TSM2R computation on the given GPU should be compute or memory-bound. In the last section, we proposed to estimate this characteristic by comparing n and $t_2^{threshold}$, so that we can accordingly adjust parameters to optimize the computation.

In the case where original problem is memory-bound ($n \le t_2^{threshold}$), we need to keep the actual computation memory-bound also (let $1 \le t_2 \le n$) and optimize for memory bandwidth utilization. On the other hand, if the original problem is compute-bound ($n > t_2^{threshold}$), we first try to keep the actual computation compute-bound too (let $t_2^{threshold} \le t_2 \le n$) and optimize computing power utilization. However, in the case where $t_2^{threshold}$ is too high on the given GPU, we also try to optimize it for memory-bound (let $1 \le t_2 \le t_2^{threshold}$) and output the result parameters that deliver better performance.

Algorithm 5 shows the parameter optimization procedure for t_2 and t_3 . We first determine the computation characteristic in Line 1. If it is memory-bound, we optimize for the total global memory access time (Line 4). Otherwise, we optimize for either total computation time (Line 9) or memory access time (Line 14). Please note that we only count the total amount of memory accesses to matrix A for simplicity, since total accesses to matrix B is much less than matrix A, so this simplification only brings minor inaccuracy. Also, considering the total accesses to matrix B would bring one additional parameter (t_1), which can be hard to optimize since t_1 is also related to threads organization that is hard for modeling-based estimation. The memory bandwidth

```
1: if n < t_2^{threshold} then
       Total_memory \approx m \times k \times \frac{n}{t_2} \times bytes_per_elem.
 2:
3.
       Bandwidth = PeakBand. \times \tilde{U}til_{mem}
       Use Gradient Descent to Optimize (t_2 \text{ and } t_3): Time = \frac{Total\_memory}{Bandwidth} with
 4:
     1 < t_2 < n and 1 < t_3
       Output: t_2 and t_3
5:
6: else
7:
        Total_flops = m \times k \times n \times 2
       Compute\_power = PeakPerf. \times Util_{comp}
8:
       Use Gradient Descent to Optimize (t_2 and t_3): Time_1 = \frac{Total_flops}{Compute_power}
9:
                                                                                                         with
     t_2^{threshold} \le t_2 \le k \text{ and } 1 \le t_3
10:
        t_{2(time_1)} \leftarrow t_2
11:
         t_{3(time_1)} \leftarrow t_3
        Total_memory \approx n \times n \times \frac{k}{t_2} \times bytes_per_elem.
12.
         Bandwidth = PeakBand. \times Util_{mem}
13:
        Use Gradient Descent to Optimize (t_2 \text{ and } t_3) in Time_2 = \frac{Total\_memory}{Bandwidth} with
14:
     1 \le t_2 \le t_2^{\text{threshold}} and 1 \le t_3
15:
        t_{2(time_2)} \leftarrow t_2
16.
         t_{3(time_2)} \leftarrow t_3
17:
         if Time_1 < Time_2 then
18:
            Output: t_{2(time_1)} and t_{3(time_1)}
19:
         else
20.
           Output: t_{2(time_2)} and t_{3(time_2)}
21:
        end if
22: end if
```

Algorithm 5: Parameter optimization for TSM2R.

utilization term ($Util_{mem}$) and computing power utilization term ($Util_{comp}$) is calculated using the equation mentioned before. Since we have two parameters (t_2 and t_3) in our optimization target, we use Gradient Descent (GD) to do the optimization. In GD, based on our experience, we set initial value of both t_2 and t_3 to be 1, and step size to be 0.1. The stop threshold is set to be 10^{-4} , since we do not need very accurate precision. The final t_2 and t_3 are rounded to the nearest integers.

To optimize t_1 , we find it only controls the number of threads in each thread block. Since the total number of threads is fixed to m, t_1 only determines how these threads are organized into thread blocks. There is trade-off: if t_1 is large, the total number of accesses to elements of matrix B is reduced, however, a large thread block means a large number of threads need to participate in the same synchronization, which may have an impact on performance. On the other hand, if t_1 is small, the total number of accesses to elements of matrix B is higher, but the smaller thread block makes scheduling more flexible and efficient. It is hard to determine the optimum value of t_1 theoretically, so we use offline profiling to choose the best value. Specifically, once t_2 and t_3 are determined, we benchmark different t_1 values that can divide $MaxOccup_{SM}$ as mentioned earlier, and choose the t_1 for the best performance. Although t_1 seems to have direct effect on shared memory allocation (or max SM occupancy), it actually has limited impact on it, since we fix the amount of shared memory per thread ($S_{thread} = t_2 \times bytes_per_element$).

3.2. Design of TSM2L

The algorithm proposed in the above sections – TSM2R – is optimized for the case where a large regular-shaped matrix multiples a tall-and-skinny matrix. In this section, we first propose a new algorithm TSM2L to handle the case where a tall-and-skinny matrix multiplies a small regular-shaped matrix. For example, an input matrix A of size 102400×4 multiples an input matrix B of size 4×4 , where the tall-and-skinny matrix is multiplied on the left. We then introduce two different optimization approaches to overcome the bottleneck that this kind of tall-and-skinny input poses.

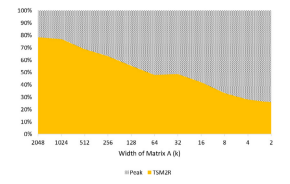


Fig. 4. Memory bandwidth usage of very small values of k with double precision (m = 15360, n = 16).

Require: input matrix A $(m \times k)$ and B $(k \times n)$ **Require:** output matrix C $(m \times n)$

1: $tcf \leftarrow thread_count_factor$

- 2: $t_1 \leftarrow tile_size_B, t_2 \leftarrow tile_size_C, t_3 \leftarrow tile_size_A$
- 3: Register: currA₁, currA₂,...,currA_{t₃}
- 4: Register: nextA₁, nextA₂,...,nextA_{t3}
- 5: Register: nextB₁, nextB₂,...,nextB_{t2}
- 6: Register: $C_1, C_2, ..., C_{t_2}$
- 7: Shared Memory: *currB* with size $t_1 \times t_2$
- 8: Threads per thread block $\leftarrow t_1$
- 9: Total thread blocks $\leftarrow m/(t_1 \times tcf)$
- 10: Total threads $\leftarrow m/tcf$
- /*loop over all the horizontal tiles of matrix A*/
- 11: **for** $r = global_tid$ to m with step size = Total threads **do**
- Perform Line 9–29 of Algorithm 4 with all occurrences of the identifier *global_tid* replaced by the identifier *r*end for
 - Algorithm 6: Proposed optimization-1 for TSM2L.

3.2.1. Performance bottlenecks

We start by adapting our previous algorithm TSM2R to handle the new case without further optimization. However, applying the algorithm to this case reveals a bottleneck. We evaluate TSM2R on this case with matrices of $15\,360 \times k$ and $k \times 16$ – where k varies – on an NVIDIA Tesla V100 GPU. As shown in Fig. 4, as the inner dimension k decreases, the memory bandwidth usage also decreases.

To explain these results, we expand upon the performance model (Algorithm 4 and 5) proposed in Section 3.1 This model assumes that the maximum theoretical occupancy is always achieved throughout the computation. However, on the one hand, since the algorithm loops $k \times \frac{n}{t^2}$ times, and *k* is very small, each thread does not perform enough workload to hide the latency, resulting in low occupancy. On the other hand, the program issues much fewer global memory reads than the case with large *k*, resulting in less efficient memory usage. Therefore, TSM2R performs in a **latency-bound mode** (neither compute-bound nor memory-bound) on this input case (i.e., a tall-and-skinny matrix multiplying a small matrix), as indicated in a prior study [33].

3.2.2. Proposed optimizations

Based on these observations, we design two further optimizations for TSM2L. Both optimizations intend to trade warp latency for memory-access latency by launching fewer threads. As a result, each thread performs more computation, and the accumulated warp latency can be replaced by the memory-access latency. Since we are launching fewer threads than the number of rows of matrix *A*, we must divide it into several horizontal tiles. Here we introduce a new parameter *tcf* to represent the tile number of matrix A in our algorithm. We launch $\frac{m}{tcf}$ threads in the new kernel.

The first optimization involves dividing the multiplication into *tcf* parts, where each part consists of multiplying a $\frac{m}{tcf}$ -row tile of matrix *A* by the entire matrix *B*. In essence, this optimization repeats the TSM2R algorithm once for each tile of matrix *A*. In this optimization, each element of matrix *A* is still accessed $\frac{n}{t_2}$ times, though each element of *B* is loaded *tcf* $\times \frac{n}{t_2}$ times, which is *tcf* times more than that in TSM2R. We describe the detail in Algorithm 6.

Require: input matrix A $(m \times k)$ and B $(k \times n)$, output matrix C $(m \times n)$

1: $tcf \leftarrow thread_count_factor$ 2: $t_1 \leftarrow tile_size_B$, $t_2 \leftarrow tile_size_C$, $t_3 \leftarrow tile_size_A$ 3: Register: currA₁, currA₂,...,currA_{t3} 4: Register: nextA₁, nextA₂,...,nextA_{t3} 5: Register: nextB₁, nextB₂,...,nextB_{t2} Register: C_1 , C_2 ..., C_2 Register: $nextC_1$, $nextC_2$..., $nextC_{t_2}$ Shared Memory: currB with size $t_1 \times t_2$ 6٠ 7. 8: 9: Threads per thread block $\leftarrow t_1$ 10: Total thread blocks $\leftarrow m/(t_1 \times tcf)$ 11: Total threads $\leftarrow m/tcf$ 12: for p = 1 to n with step size = t_2 do $currB[local_tid, 1:t_2] \leftarrow B[local_tid, p:p+t_2-1]$ 13: $currA_{1:t_3} \leftarrow A[global_tid, 1:t_3]$ 14: 15: **for** j = 0 to k with step size = t_1 **do** 16: ThreadsSynchronization() /*prefetch the next tile of B into nextB*/ $nextB_{1:t_2} \leftarrow B[j + t_1 + local_tid - 1, p : p + t_2 - 1]$ 17: $C_{1:t_2} \leftarrow C[global_tid, p: p + t_2 - 1]$ 18: /*loop over all the horizontal tiles of matrix A*/ 19: for r = global_tid to m with step size = Total threads do /*prefetch the next tile of C into registers*/ 20: **if** r + Total threads $\leq m$ **then** $nextC_{1:t_2} \leftarrow C[r + Total threads, p: p + t_2 - 1]$ 21: end if 22: 23: Perform Line 18–25 of Algorithm 4, with all occurrences of the identifier global_tid replaced by the identifier /*store the sum so far in C*/ $C[r, p: p + t_2 - 1] \leftarrow C_{1:t_2}$ /*load the prefetched tile C from nextC to C*/ 24: 25: $C_{1:t_2} \leftarrow nextC_{1:t_2}$ 26: end for ThreadsSynchronization() 27: /*load the prefetched tile B from nextB to shared memory*/ $currB[local_tid, 1:t_2] \leftarrow nextB_{1:t_2}$ 28: 29: end for 30: end for

Algorithm 7: Proposed optimization-2 for TSM2L.

The second optimization is to interleave the computation of the tiles, rapidly loading elements from different tiles of matrix A and loading and storing intermediate sums in matrix C. Once a $t1 \times t2$ tile of matrix B is loaded, the intermediate results are loaded, computed, and stored for each tile of matrix A. The C register set is loaded with values from matrix C which contains the product accumulated so far. After the computation is finished, the values are stored to matrix C again as the next tile of matrix A is prepared for computation. To quickly switch between tiles, values from matrix C are prefetched in addition to the prefetching already described in Algorithm 4. A new set of registers, $nextC_{1:t_2}$, is used to store the values of C associated with the next tile of A. The elements of *A* and *B* are accessed only $\frac{n}{t_2}$ times, though each element of *C* is accessed $\frac{k}{t_1} \times \frac{n}{t_2}$ times. However, since we do not achieve either high occupancy or high memory bandwidth in this case, we are not as concerned about issuing more memory read instructions. We describe the detail in Algorithm 7.

Fig. 5 illustrates the effects of the two optimizations on both performance and memory bandwidth usage. As fewer and fewer threads are launched, the impact of warp latency is replaced with that of different kinds of latency such as memory bandwidth latency. As a result, computation time decreases and memory access bandwidth increases in this case. Note that for this case the number of threads launched must be reduced to at least $\frac{1}{64}$ of

m before a significant decline in speedup or memory bandwidth utilization occurs. This is because the kernel must manage so many threads that perform little work when $m = 10^7$.

Therefore, we must choose an appropriate *tcf*, determining the number of threads to launch for each kernel. If the algorithm is launched with an insufficient number of threads, the parallelism becomes too low and hence the performance would suffer. If the algorithm is launched with too many threads, the performance would be impacted by warp latency just as it does in the naive adaptation of TSM2R. We thus must determine an appropriate *tcf* for each target system with offline profiling.

3.3. Design summary

We summarize the design of TSM2R and TSM2L, including our performance model, in Table 1.

4. Experimental evaluation

4.1. Experiments setup

We implement our TSM2R and TSM2L using CUDA C for single and double precision floating-point input. We disable compiler auto unrolling in favor of explicit loop unrolling for better control on register allocation. Note that since our proposed algorithms mainly target traditional scientific computing applications rather than machine learning applications, we omit an evaluation on half-precision input. We evaluate our optimized implementations on two heterogeneous testbed clusters, which are Darwin [13] at Los Alamos National Laboratory and PantaRhei [25] at the University of Alabama. We run each test on a single GPU node with single GPU. We conduct our tests on four different commonly used modern NVIDIA GPUs with four different micro-architectures: Kepler, Maxwell, Pascal, and Volta. For Kepler GPU, we use Tesla K40c, which has 1430 GFLOPS peak double floating-point performance and 288 GB/s memory bandwidth. For Maxwell GPU, we use Tesla M40, which has 213 GFLOPS peak double floating-point performance and 288 GB/s memory bandwidth. For Pascal GPU, we use Tesla P100, which has 4600 GFLOPS peak double floating-point performance and 720 GB/s memory bandwidth. For Volta GPU, we use Tesla V100, which as 7500 GFLOPS peak double floating-point performance and 900 GB/s memory bandwidth. We provide more information about our experimental clusters and GPUs in Table 2.

For comparison, we compare our TSM2R and TSM2L with GEMM in the current latest cuBLAS library [2] and latest BLASX library [34]. Also, we try to compare our work with KBLAS [1], however since its GEMM kernel is based on cuBLAS, its performance is identical to cuBLAS, so we omitted its results. Each test is repeated multiple times to reduce noise and timed using CUDA Events API. We measure performance by calculating the performance of FAMD instructions. We also measure the global memory throughput using nvprof on the command line with --metrics gld_throughput option. In addition, we use --metrics gld_efficiency option to verify if 100% global memory access efficiency is achieved in our optimization.

Our input matrices are initialized with random floating-point numbers between 0 and 1. We test the multiplication of a large squared matrix and a tall-and-skinny matrix for TSM2R and the multiplication of a tall-and-skinny matrix and a small squared matrix for TSM2L. Specifically, for TSM2R, the size of the large regular-shaped matrix is from 10240×10240 to 30720×30720 , and the size of the tall-and-skinny matrix ranges from $10240 \times n$ to $30730 \times n$ with n equals 2, 4, 8, and 16. For TSM2L, the size of the tall-and-skinny matrix ranges from $10^4 \times k$ to $10^7 \times k$ with k equals 8 or 16, and the size of the small regular-shaped matrix is 8 or 16.

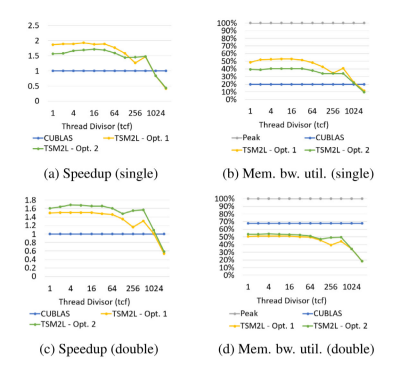


Fig. 5. Performance comparison with single and double precision using different tcf ($m = 10^7$, k = n = 16).

Table	1
Table	- 1

Summary of TSM2R and TSM2L design

TSM2R	Large guare or regtangular matrix by tall and glyinny matrix					
	Large square or rectangular matrix by tall-and-skinny matrix					
TSM2L	Tall-and-skinny matrix by small square matrix					
TSM2R optimizations	Compute and memory-bound cases					
Algorithm 1	Inner product only					
Algorithm 2	Outer product: saves global memory accesses					
Algorithm 3	Shared memory: more efficient global accesses to matrix B					
Algorithm 4	Data prefetch: overlap compute and memory operations					
TSM2L optimizations	Latency bound cases					
Algorithm 6	Divide matrix A into horizontal tiles: compute each tile sequentially					
Algorithm 7	Divide matrix A into horizontal tiles: interleave the computation of each tile					
Performance model						
Parameter t_1	Number of rows of a tile of Matrix B					
Parameter t_2	Number of elements of C each thread computes at a time					
Parameter t_3	Number of elements of A each thread fetches at a time					
Util _{comp}	Computing power utilization term					
Util _{mem}	GPU memory bandwidth utilization term					
$t_2^{threshold}$	$= \frac{Peak Perf.}{Peak Band.} \times bytes_per_elem.$					
	Determines whether a computation is compute-bound or memory-bound					

Table 2

Experimental platforms with detailed GPU information.

		PantaRhei			
CPU	Intel Xeon E5-2	Intel Xeon E5-2650v2			
Memory		251 GB		384 GB	
GPU	Tesla K40c	Tesla M40	Tesla P100	Tesla V100	
Architecture	Kepler	Maxwell	Pascal	Volta	
GPU memory	12 GB	12 GB	16 GB	16 GB	
Peak performance (single)	5046 GFLOPS	6844 GFLOPS	10 600 GFLOPS	15 000 GFLOPS	
Peak performance (double)	1430 GFLOPS	213 GFLOPS	4600 GFLOPS	7500 GFLOPS	
Memory bandwidth	288 GB/s	288 GB/s	720 GB/s	900 GB/s	

4.2. Evaluation of TSM2R

In this section, we first evaluate the performance of TSM2R with different input sizes and compare it with state-of-the-art libraries on K40c, M40, P100, and V100.

4.2.1. Tests with different optimization combinations

We use the GEMM in cuBLAS as our comparison baseline. We apply different combinations of optimization in TSM2R and compare them with GEMM in cuBLAS and BLASX. We have in total four versions of TSM2R:

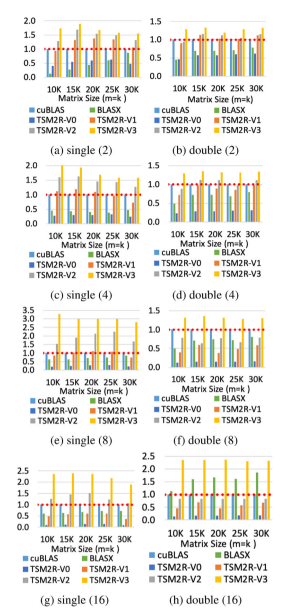


Fig. 6. Speedup comparison with single and double precision on K40c (n = 2, 4, 8, 16).

- V0: the most straightforward inner product version as described in Algorithm 1;
- V1: the outer version as in Algorithm 2. This version reduces the total number of global memory accesses from algorithm level;
- V2: based on outer production version as in Algorithm 2, we add the use of shared memory, leading to more efficient global memory access to matrix B;
- V3: based on the outer production version as in Algorithm 2 and the use of shared memory, we add data prefetch. This is the best version of our optimized implementation, which is described in Algorithm 4.

We provide detailed performance breakdowns on K40c and V100, but our optimization behaves similarly on other GPUs. To evaluate our optimization, we need to determine by which resource our program is bounded. Since, $t_{2(k40c)}^{threshold} \approx 40$, the computation is always memory bound for the given *n* values. The optimized parameters are: $t_2 = n$, $t_3 = 4$, and $t_1 = 128$. The

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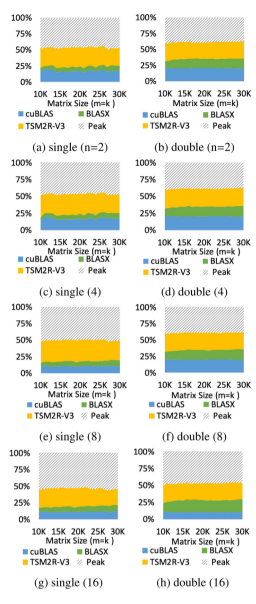


Fig. 7. Memory bandwidth utilization comparison on K40c (n = 2, 4, 8, 16).

parameters are only applied to the last two versions of TSM2R. Fig. 6 shows the speedup of different versions in single and double precision. From the results, we can see that the TSM2R-VO suffers from poor performance due to the requirement of much higher number of global memory accesses in the inner product version. TSM2R-V1, on the other hand, significantly improves the performance compared to TSM2R-V0 (2.2x~4.7x faster), since it requires much lower number of global memory accesses. TSM2R-V2 further improves the efficiency of global memory access to matrix B, which plays a vital role in the overall performance. In addition, the shared memory shares tiles of matrix B between threads within a thread block, which also reduces the total number of memory accesses to matrix B. This leads to additional 1.1x to 2.1x speedup. Finally, the data prefetch introduced in TSM2R-V3 further mitigates the memory access bottleneck, which brings additional $1.3x \sim 3.5x$ speedup (1.9x on average).

4.2.2. Memory throughput analysis

Fig. 7 shows the memory throughput of TSM2R-V3, cuBLAS and BLASX in both single and double precision on K40c GPU.

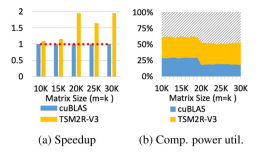


Fig. 8. Speedup and computing power utilization comparison with double precision on M40 (n = 16).

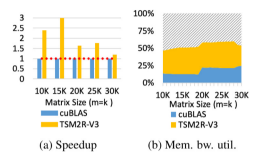


Fig. 9. Speedup and memory bandwidth utilization comparison with double precision on P100 (n = 16).

Result shows that TSM2R brings $12.5\% \sim 26.6\%$ (17.6% on average) improvement on memory bandwidth utilization compared with cuBLAS and $20.1\% \sim 38.8\%$ (24.3% on average) improvement compared with BLASX.

4.2.3. Tests on different micro-architectures

In addition to Kepler micro-architecture, we also conduct tests on newer Maxwell, Pascal, and Volta GPUs. Similar to the Kepler GPU, we get $t_{2(m40)}^{threshold} \approx 6$ and $t_{2(p100)}^{threshold} \approx 50$. Tesla M40 has lower computing power, so the computation with input with n = 16is compute bound. Our parameter optimization procedure also output parameters in favor of computing optimization: $t_2 = 8$, $t_3 = 4$, and $t_1 = 256$. As shown in Fig. 8, our optimized implementation achieves 1.1x-1.9x (1.47x on average) speedup on Tesla M40 with 7% to 37.3% (20.5% on average) computing power usage improvement compared to the GEMM function in cuBLAS 9.0. P100 has much stronger computing power, as we can see the computation with input with n = 16 is memory bound. Our parameter optimization procedure also output parameters in favor of memory optimization: $t_2 = 4$, $t_3 =$ 4, and $t_1 = 128$. As shown in Fig. 9, our optimized implementation achieves $1.1x \sim 3.0x$ (2.15x on average) speedup on Tesla P100 with 17% to 47.6% (34.7% on average) memory bandwidth utilization improvement compared to the GEMM function in cuBLAS.

We also test TSM2R on the NVIDIA Tesla V100 GPU with Volta micro-architecture. Due to larger memory space on V100, we further increase the size of regular-shaped matrix to 40 960 × 40 960. As $t_{2(v100)}^{threshold} \approx 70$, the computation is memory-bound for the given values. To ensure maximum performance and account for the Volta's architectural improvements, we optimize the parameters via brute-force. The optimized parameters are $t_1 = 128$, $t_2 = n$, and $t_3 = 32$ for single precision. For double precision, the optimized parameters are $t_1 = 128$, $t_2 = n$, and $t_3 = 12$ otherwise. As shown in Fig. 10, we exhibit gradually improving performance from TSM2R-V0 to TSM2R-V2, similar to K40c. For TSM2R-V3, our best version, speedups of

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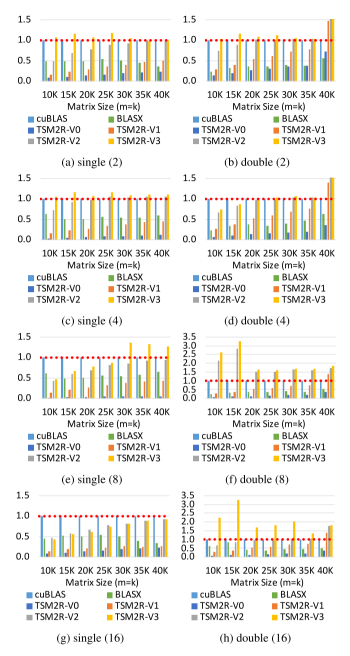


Fig. 10. Speedup comparison with single and double precision on V100 (n = 2, 4, 8, 16).

up to 1.35x (0.91x on average) are achieved on single precision, while speedups of up to 3.2x (1.5x on average) are achieved on double precision. Note that the speedup for n = 16 on single precision is slower than cuBLAS. This is due to cuBLAS's single-precision GEMM being optimized for 32×32 matrices; thus we no longer target this case. Finally, we note that the kernels achieve higher memory bandwidth utilization on V100 than on other GPUs, as shown in Fig. 11. This is partly attributed to the improvements of Volta over previous micro-architectures. More specifically, V100 with improved HBM2 memory allows more workloads to obtain up to 19% more memory bandwidth utilization than Pascal GPUs, according to its whitepaper [24]. We provide experimental metrics for our TSM2R kernels in Table 3.

Due to our performance modeling, we can predict TSM2R's performance on the Nvidia Tesla A100, with Ampere architecture. Its peak double-precision floating point performance is 9.7

Table 3

Details of TSM2R kernel. Note that the number of registers is experimental data collected from NVCC and depends on n (n = 16).

GPU	Precision	t_1	t_2	t_3	Threads/Block	Shared memory (bytes)	# Registers
K40c	Single	128	п	4	128	$512 \times (n \times 4)$	64
K40c	Double	128	п	4	128	$1024 \times (n \times 8)$	128
M40	Single	256	8	4	256	8192	40
M40	Double	256	8	4	256	16384	70
P100	Single	128	4	4	128	2048	32
P100	Double	128	4	4	128	4096	56
V100	Single	128	п	32	128	$512 \times (n \times 4)$	144
V100	Double	128	п	16	128	$1024 \times (n \times 8)$	180
V100	Double	128	п	12	128	$1024 \times (n \times 8)$	168

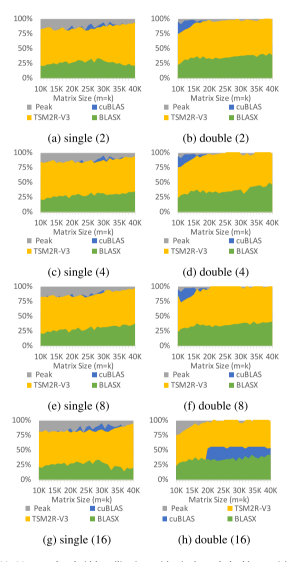


Fig. 11. Memory bandwidth utilization with single and double precision on V100 (n = 2, 4, 8, 16).

TFLOPS, $1.3 \times$ that of the V100, and its global memory bandwidth is 1555 GB/s, $1.73 \times$ that of the V100. Since all the cases considered in this paper are memory-bounded on the A100 (as $t_{2(a100)}^{threshold} \approx 50$) and our implementation has already achieved more than 90% efficiency in memory bandwidth, we expect TSM2R can achieve a speedup of about $1.7 \times$ on the A100 over the V100. Since we do not currently have access to a Nvidia Tesla A100 GPU, our estimates are based on the available whitepaper

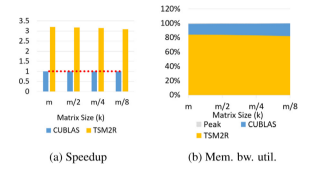


Fig. 12. Performance comparison with double-precision rectangular input on V100 (m = 15360, n = 16).

and do not take into account any other architectural improvements [23].

4.2.4. Tests on non-squared input

We also evaluate TSM2R with rectangular input matrices ($m \times k$) on V100, where k is smaller than m by certain small integer factors. Evaluating this case reveals very little performance impact, as demonstrated in Fig. 12. Although smaller than m, k is still large enough to ensure the kernel to follow our performance model. The memory bandwidth utilization of the kernel remains similar to the case where m = k, and the performance of the kernel scales linearly with the matrix size.

4.3. Evaluation of TSM2L

We next evaluate the performance of TSM2L and compare it with cuBLAS on V100. For TSM2L, we must choose the variable *tcf* for each matrix input combination. We obtain these values through experiments that vary *tcf*. As a result, for $m = 10^4$, 10^5 , 10^6 , 10^7 , we select *tcf* values as 1, 1, 2, and 8 for single precision and values 1, 1, 1, and 4 for double precision. Considering two proposed optimizations for TSM2L, we have two versions of TSM2L: TSM2L-Opt1, based on Algorithm 6, and TSM2L-Opt2, based on Algorithm 7.

As shown in Fig. 13, TSM2L can obtain speedups over cuBLAS ranging from $1.1x \sim 3.5x$ (2.5x on average) in single precision and speedups from $1.0x \sim 1.7x$ (1.3x on average) in double precision. TSM2L-Opt1 generally performs better on single precision input than TSM2L-Opt2, while TSM2L-Opt2 performs better than TSM2L-Opt1 in several double precision cases. In addition, as shown in Fig. 14, TSM2L achieves memory bandwidth utilization of up to 55% peak global memory bandwidth (40% on average). In single precision, TSM2L utilizes significantly more memory bandwidth than cuBLAS, up to 40% more (25% on average). However, in double precision, TSM2L uses only slightly more memory

Table 4

Details of TSM2L kernel on V100. Note that the number of registers is experimental data collected from NVCC and depends on n (n = 16).

· · ·							
Optimization	Precision	t_1	t_2	t_3	Threads/Block	Shared memory (bytes)	# Registers
TSM2L-Opt1	Single	128	п	32	128	$512 \times (n \times 4)$	144
TSM2L-Opt1	Double	128	п	16	128	$1024 \times (n \times 8)$	180
TSM2L-Opt1	Double	128	п	12	128	$1024 \times (n \times 8)$	168
TSM2L-Opt2	Single	128	п	32	128	$512 \times (n \times 4)$	251
TSM2L-Opt2	Double	128	п	16	128	$1024 \times (n \times 8)$	254
TSM2L-Opt2	Double	128	п	12	128	$1024 \times (n \times 8)$	252

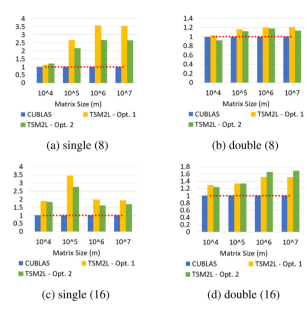


Fig. 13. Speedup comparison with single and double precision on V100 (k = n = 8, 16).

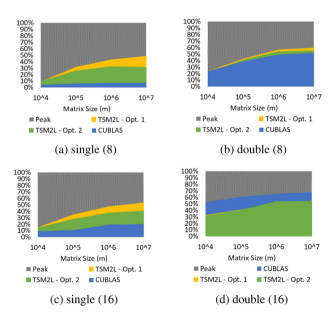


Fig. 14. Memory bandwidth utilization with single and double precision on V100 (k = n = 8, 16).

bandwidth in the case that k = n = 8, and in the case that k = n = 16, cuBLAS uses more memory bandwidth. However, since TSM2L still outperforms cuBLAS, this can be explained by inefficient memory use patterns in the GEMM kernel. We provide experimental metrics for our TSM2L kernels in Table 4.

5. Related works

A preliminary version of this work was published in [8]. It introduces our TSM2R algorithm and evaluates it on Kepler, Maxwell, and Pascal GPUs respectively. In this paper, we expand the evaluation by adding experiments on the Volta GPU V100. Moreover, we also broaden the applicability of this work through our new TSM2L algorithm to handle a new input case.

Ernst et al.'s work also focuses on optimizing tall-and-skinny GEMM [17]. It proposes two algorithms for tall-and-skinny input: TSMTTSM, where a tall-and-skinny matrix is multiplied by transposed tall-and-skinny matrix, and TSMM, where a tall-and-skinny matrix is multiplied by a small square matrix. It evaluates these algorithms on the Volta GPU V100 with double-precision real and complex floating-point numbers. Although TSMTTSM's input case is not considered by our work, TSMM's input case is the same as TSM2L's. However, TSMM and TSM2L approach this input case differently. Specifically, TSMM takes matrices in row-major format as input, so its optimizations focus on avoiding partially-written cache lines while storing columns of matrix C. TSMM launches multiple threads per row of matrix C, with each thread storing several columns of matrix C. Unlike TSMM, the design of our TSM2L takes into account the latency of launching many threads that perform little work, and its optimizations' focus on managing both warp and memory bandwidth latency. Moreover, note that our TSM2L achieves superior performance. TSMM only achieves speedups over cuBLAS where the small dimension, k = n, is less than 8, whereas our TSM2L achieves speedups with the small dimension up to 16.

6. Conclusion

In this work, we first analyze the performance bottleneck of current GEMM in the latest cuBLAS library. We identify that current implementations lack of full utilization of computing power or memory bandwidth when the input shape is tall-and-skinny. Then, we discovered the potential challenges of optimizing tall-and-skinny GEMM since its workload can vary between compute bound, memory bound, and latency bound. Next, we propose two high-performance GEMM algorithms – TSM2R and TSM2L – on GPUs for tall-and-tinny input with several optimization techniques focusing on GPU resource utilization. Finally, experiment results show that our optimized implementations can achieve speedups tall-and-skinny matrix-matrix multiplication with diverse input sizes on modern GPUs.

CRediT authorship contribution statement

Cody Rivera: Conceptualization, Methodology, Software, Writing - original draft. **Jieyang Chen:** Conceptualization, Methodology, Software, Writing - original draft. **Nan Xiong:** Software, Validation. **Jing Zhang:** Writing - review & editing. **Shuaiwen Leon Song:** Writing - review & editing. **Dingwen Tao:** Supervision, Writing - review & editing.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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