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DOI: 10.1016/j.jocs.2018.01.005

Document Version
Accepted author manuscript

Link to publication record in Manchester Research Explorer

Citation for published version (APA):

Published in:
Journal of Computational Science

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Batched One-sided Factorizations of Tiny Matrices using GPUs: Challenges and Countermeasures

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Abstract

The use of batched matrix computations recently gained a lot of interest for applications, where the same operation is applied to many small independent matrices. The batched computational pattern is frequently encountered in applications of data analytics, direct/iterative solvers and preconditioners, computer vision, astrophysics, and more, and often requires specific designs for vectorization and extreme parallelism to map well on today’s high-end many-core architectures. This has led to the development of optimized software for batch computations, and to an ongoing community effort to develop standard interfaces for batched linear algebra software. Furthering these developments, we present GPU design and optimization techniques for high-performance batched one-sided factorizations of millions of tiny matrices (of size 32 and less). We quantify the effects and relevance of different techniques in order to select the best-performing LU, QR, and Cholesky factorization designs. While we adapt common optimization techniques, such as optimal memory traffic, register blocking, and concurrency control, we also show that a different mindset and techniques are needed when matrices are tiny, and in particular, sub-vector/warp in size. The proposed routines are part of the MAGMA library and deliver sig-

\textsuperscript{*}This is an extended version of our conference paper \cite{1} that was invited to the JoCS special issue (https://doi.org/10.1016/j.procs.2017.05.250).

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significant speedups compared to their counterparts in currently available vendor-optimized libraries. Notably, we tune the developments for the newest V100 GPU from NVIDIA to show speedups of up to 11.8×.

Keywords: GPU computing, Matrix factorization, Batch computation

1. Introduction

Batch computations apply the same numerical algorithm to a fairly large number of relatively small problems. The batched computing workload is quite different than the common workload for just one, typically large, matrix. The latter is served well by many software packages, including LAPACK \[2\], ScaLAPACK \[3\], PLASMA \[4\], and MAGMA \[5\]. The former, however, is relatively recent, and gained a lot of attention in many scientific communities, e.g., quantum chemistry \[6\], sparse direct solvers \[7\], astrophysics \[8\], and signal processing \[9\]. Software libraries such as Intel’s MKL \[10\], NVIDIA’s cuBLAS \[11\], and MAGMA recently started to provide highly optimized batched routines for many of the BLAS and LAPACK operations.

Existing numerical linear algebra software packages rarely achieve good performance on matrices of small sizes, because most of the optimization techniques that they carry out pay off only on large matrices. For example, the hybrid lookahead technique in MAGMA \[12\] is used to overlap the panel factorization (on the CPU) with the trailing matrix update (on the GPU). This design strategy is not efficient for small sizes, since the updates are no longer compute intensive, and therefore fail to overlap the panel factorization and the CPU-GPU communication. This is why new developments with different design strategies are needed.

While there have been new developments for GPU accelerated batched computations, for example the work done by Haidar et al. \[13\] and Abdelfattah et al. \[14\], there is still room for significant improvements when the matrix sizes are tiny. For these extremely small problems, the LAPACK-style blocking cannot achieve high performance, even if it is carried out on the GPU solely. Since the
computation becomes memory bound on such small sizes, the cost of writing the factorized panel and then reading it back to perform the update becomes significant. Furthermore, the parallelization (to achieve sufficiently high occupancy) and the vectorization (for efficient warp use) become more challenging when sizes are less than 32, and must, for example, be done across matrices. Therefore, other design, parallelization, and vectorization strategies must be discovered in order to resolve the aforementioned issues regarding memory traffic and parallelization.

This paper presents highly optimized GPU kernels for batched one-sided factorizations. The paper extends the previous work by the authors for batched LU factorization and matrix inversion [1], and applies the same design principles to the QR and Cholesky factorizations. In terms of the workload size, we consider one million matrices of tiny sizes, typically from 1 up to 32. In addition to the applications already mentioned, these factorizations are of particular importance to sparse direct solvers, such as the multifrontal solvers that can be found in SuiteSparse [15]. We adopt a step-by-step methodology, where incremental improvements in the kernel design lead to incremental performance gains. Such a methodology automatically justifies all of our design choices. While all the kernels share the same optimization techniques, our design for the LU factorization adopts a unique lazy swap strategy, which eliminates the expensive intermediate row interchanges, thus leading to a much faster kernel, but that is still numerically equivalent to an LAPACK-style LU-factorization. The performance results show significant speedups against the vendor-supplied cuBLAS kernels on a Pascal P100 GPU, as well as on the new Volta V100 GPU.

2. Related Work

The design of high performance dense linear algebra (DLA) software for GPUs was originally motivated by the high performance GPUs can achieve in embarrassingly parallel, compute intensive tasks, most notably on the matrix-matrix multiplication (GEMM) [16, 17, 18]. The high performance GEMM enabled
the development of high performance DLA in libraries like MAGMA [5], where many of the LAPACK numerical algorithms are designed in a hybrid style to take advantage of both CPUs and GPUs [12].

The growing demand for high performance dense linear algebra on large batches of small matrices has led to early developments for batch matrix multiplication [19][20], which were then followed by more optimized kernels being available in cuBLAS, starting with version 8.0. The development of the batched GEMM in MAGMA enabled the development of batched one-sided factorizations routines based on the LAPACK-style blocking [13], but on a smaller scale. For example, while non-batched routines use a large blocking size (e.g., 512 to 1024) to get asymptotically optimal performance, batched routines block by much smaller sizes (e.g., 8 to 32), and rely on batched GEMM that is specifically tuned for small sizes in order to extract performance [21][22]. The developments for extremely small matrices, however, are more challenging. An approach that relies on separate panel/update stages [13] leads to redundant memory traffic. This cost can be affordable for medium sizes (e.g., 64 up to 256), but becomes significant for smaller sizes.

This is why other research efforts followed a one-kernel approach, where all computations are fused into a single GPU kernel. For example, Wang et al. [23] introduced FPGA-based parallel LU factorization of large sparse matrices, where the algorithm is reduced to factorizing many small matrices concurrently. Villa et al. [24] developed a GPU-based batched LU factorization, which has been used in subsurface transport simulation, where many chemical and microbiological reactions in a flow path are simulated in parallel [25]. Kurzak et al. [26] developed batched Cholesky factorization in single precision for sizes up to $100 \times 100$, which was used in an Alternating Least Squares (ALS) solver. Masliah et al. developed batched GEMM for very small sizes for both CPUs and GPUs [27]. Batched matrix inversion has been also introduced in the context of generating block-Jacobi preconditioners [28]. Batched QR factorization is of particular importance to H-matrix computation, as highlighted by Akbudak et al. [29], and by Boukaram et al. [30]. Kim et al. also introduced batched
\texttt{GEMM}, triangular solve, and \texttt{LU} (no pivoting) for CPUs and Intel's Xeon Phi architectures based on a compact interleaved data layout \cite{31}.

This paper follows the same one-kernel approach to improve the MAGMA performance on very small sizes. It complements the work by Haidar et al. \cite{13}, which outperforms cuBLAS for medium and large sizes, but trails it for the sizes we focus on (up to 32).

3. Contributions

Below is a list of contributions for this paper.

1. Highly optimized GPU kernels for one-sided factorization on batch workloads. The developed kernels significantly outperform the state of the art designs from the vendor provided software. We typically consider single-node workloads that involve millions of extremely small matrices.

2. A set of unified design techniques that are oblivious to the three algorithms considered (\texttt{LU}, \texttt{QR}, and Cholesky factorizations). The paper manages to find a common ground among the three algorithms to achieve high performance.

3. The paper presents a detailed study of the different choices for every aspect of the kernel design, including thread configuration, matrix storage, occupancy, and others. The paper justifies the final design choice by showing intermediate performance results for different choices. Such a detailed study can be considered as a guide for designing other algorithms on similar workloads.

4. Background

This section introduces the computational steps for the \texttt{LU}, \texttt{QR}, and Cholesky factorizations on square matrices of size $N \times N$. The description follows the \texttt{LAPACK} notations in double precision arithmetic.

The \texttt{LU} factorization computes the $L$ and $U$ factors of a general matrix $A$, such that $A = P \times L \times U$, where $P$ is a permutation matrix that reflects the row
interchanges required for pivoting. The matrix $L$ is unit lower triangular, while $U$ is upper triangular. The permutation matrix $P$ is stored in a compact format using a *pivot vector* ($\text{IPIV}$), such that for $i \in \{1,2,\cdots,N\}$, row $i$ has been swapped with row $\text{IPIV}(i)$.

There are four main steps in performing the unblocked LU factorization. Namely, these are: (1) locate the maximum absolute value in the current column ($\text{IDAMAX}$); (2) swap current row with the row with maximum absolute value in the current column ($\text{DLASWP}$); (3) scale the current column ($\text{DSCAL}$); and (4) rank-1 update ($\text{DGER}$). Algorithm 1 shows the factorization using the four steps. According to LAPACK working notes [32], the LU factorization of a square matrix performs $(\frac{2N^3}{3} - \frac{N^2}{2} + \frac{5N}{6})$ floating point operations ($\text{FLOPs}$).

**Algorithm 1: Unblocked LU factorization**

```
for i=1 to N do
    \text{IPIV}[i] = \text{max}_i = \text{IDAMAX}( \text{ABS}(\text{A}[i:N,i]) )
    \text{if} \, \text{ABS}(\text{A}[\text{max}_i,i]) = \text{ZERO} \text{ then}
        //U is singular, report error.
    \text{end}
    \text{DLASWP: Exchange rows A[i, 1:N] and A[\text{max}_i, 1:N]}
    \text{DSCAL: A[i+1:N,i] *= (1 / A[i,i])}
end
```

While the LU factorization is able to factorize symmetric positive definite (SPD) matrices, the Cholesky factorization, shown in Algorithm 2, introduces a much faster algorithm for such matrices. The algorithm factorizes an SPD matrix $A = LL^T$, where $L$ is a lower triangular matrix. In each iteration, the algorithm (1) computes the square root of the current diagonal element ($A[i,i]$), (2) scales all the elements below that element ($A[i+1:N,i]$), and (3) applies a symmetric rank-1 update to the lower triangular part of the trailing matrix ($A[i+1:N,i+1:N]$). The Cholesky factorization algorithm performs $(\frac{N^3}{3} + \frac{N^2}{2} + \frac{N}{6})$ FLOPs.
Algorithm 2: Unblocked Cholesky factorization

for \( i = 1 \) to \( N \) do
  if \( A[i, i] \leq 0 \) then
    // report error for non-positive definiteness.
  end
  \( A[i, i] = \sqrt{A[i, i]} \)
  DSCAL: \( A[i+1:N,i] *= (1 / A[i,i]) \)
  // symmetric rank-1 update (DSYR)
end

The third algorithm that we consider is the QR factorization, where a matrix \( A \) is represented as the product \( Q \times R \), where \( Q \) is an orthogonal matrix, and \( R \) is an upper triangular matrix. Following the LAPACK implementation of the algorithm, the \( Q \) matrix is not computed explicitly. It is rather represented as a product of elementary Householder reflectors, namely \( Q = H(1) \ H(2) \ \cdots \ H(N) \), where

\[
H(i) = I - \tau vv^T.
\]

Considering the reflector \( H(i) \), the corresponding vector \( v \) has \( v[1:i-1] = 0, v[i] = 1 \), which is not explicitly stored, and \( v[i+1:N] \), which is stored in \( A[i+1:N,i] \). Algorithm 3 shows the basic steps of the QR factorization: (1) generate an elementary reflector \( H(i) \) by computing \( v \) and \( \tau \) (the DLARFG routine); and (2) apply \( H(i) \) to the trailing submatrix. The QR factorization requires \( (4N^3/3 + 2N^2 + 14N/3) \) FLOPs.

Algorithm 3: Unblocked QR factorization

for \( i = 1 \) to \( N \) do
  // Generate elementary reflector \( H(i) \)
  // to annihilate \( A[i+1:N,i] \)
  \( (v, \tau) \leftarrow \text{DLARFG}(A[i:N,i]) \)
  // DLARF: Apply \( H(i) \) to \( A[i:N,i+1:N] \)
  \( A[i:N,i+1:N] -= \tau \ v \ v^T A[i:N,i+1:N] \)
end
As pointed out before, there is no need to use LAPACK-style blocking techniques, which is the strategy adopted for LU, QR, and Cholesky factorizations in the \texttt{DGETRF}, \texttt{DGEQRF}, and \texttt{DPOTRF} routines, respectively. The reason is that a very small matrix can be kept in registers or shared memory during the entire factorization. This means that it makes no difference if (additional) blocking is used or not, since all the data is blocked and accessed from fast memory until the factorization is complete.

5. System Setup

We illustrate our findings on two systems accelerated with high-end NVIDIA GPUs. The first system is equipped with two 10-core Intel Haswell processors (E5-2650 v3, running at 2.3 GHz) and a Pascal GPU (Tesla P100). The GPU has 56 streaming multiprocessors, with a base clock of 1.189 GHz. The second system is equipped with an identical CPU, but the GPU is a Volta V100, which features 80 multiprocessors, running at 1.38 GHz. Both GPUs have 16GB of CoWoS Stacked HBM2 memory, and are attached to the host CPU through a PCIE interconnect. All results are obtained using the CUDA 9.0RC toolkit. We point out that to quantify the effect of our techniques, and to justify design choices, we show incremental results on the P100 GPU only, while the final performance results are shown on both GPUs.

6. General Design Criteria

All the kernels discussed in this paper have some common design aspects. First, each matrix is factorized using exactly one CUDA thread block (TB). Upon launch, the kernel is configured with as many TBs as the number of matrices, i.e., the batch size (\texttt{batchCount}). The kernel grid in CUDA is, in general, a three dimensional array (\texttt{gx}, \texttt{gy}, \texttt{gz}). We use the \texttt{gx} dimension only to launch a 1D array of TBs. The maximum value for \texttt{gx} is $2^{31} - 1$, which means that our kernels can solve billions of matrices in a single kernel launch.
Second, we adopt an optimal memory traffic strategy. Each matrix will be read and written exactly once. Since the computation is memory bound on such small sizes, it is an important design choice to keep the entire matrix cached in a fast memory level in order to avoid any redundant memory traffic.

Third, we use C++ templates to generate fully unrolled codes. Since the sizes of interest are finite, the size of the input matrix is passed as a template parameter. For such a range of very small sizes, this is a crucial decision. Fully unrolled loops get rid of integer and branch instructions, which can be quite an overhead [27].

Finally, all kernels use unblocked computations. There is no need to factorize a panel of width $nb > 1$ so that the trailing updates use L3 BLAS operations that are rich in data reuse. As pointed out earlier, each matrix is entirely kept in registers or shared memory, which means that data reuse is preserved anyway. Every kernel factorizes one column at a time and carries out the required transformation to the trailing matrix.

### 7. Design Choice 1: Thread Configuration

For most linear algebra kernels, the configuration of TBs can be 1D or 2D, which is one of our design parameters. A 2D configuration simplifies the coding effort. To use an $N \times N$ thread configuration to factorize an $N \times N$ matrix, each thread is responsible for one element of the matrix. Thus, reading and writing the entire matrix can be done through one line of code each. In addition, the trailing matrix update is fully parallelized among threads. On the other hand, reading, writing, and updating the matrix will be written in loops if a 1D configuration of $N$ threads is used. Our analysis shows that there are multiple reasons to reject the 2D configuration in favor of the 1D alternative. In batch workloads, it is important to optimize the throughput of the processed matrices, which is achieved by maximizing the number of resident TBs per multiprocessor. Using a 2D configuration, the occupancy level of each multiprocessor are far from optimal, and thus resulting in more limited parallelism, which is likely to
produce a very poor performance. For example, the 2D configuration requires
256 threads for a $16 \times 16$ matrix. This limits the number of resident TBs per
multiprocessor to 8, which can host a maximum of 2048 threads. A modern
Kepler GPU can have up to 16 resident TBs per multiprocessor, while later
generations can host up to 32. This means that the occupancy at size 16 is
brought down by a factor of 2 on a Kepler GPU, and by a factor of 4 on later
GPUs. Using a 1D configuration, the occupancy is no longer limited by the
number of threads, but rather by the amount of memory resources required by
each TB.

Another motivation to abandon the 2D configuration is synchronization.
The use of $N^2$ threads is likely to produce too many barriers in the kernel. At
each iteration, for example, threads possessing the trailing matrix always wait
for the threads performing the factorization of the current column, and threads
performing the factorization in the next iteration have to wait for the update
to finish. On the contrary, a 1D design can be free of synchronization points.
Even with the use of the new `_syncwarp()` function introduced in CUDA 9.0,
we observe that this function is obviously more lightweight than the legacy
`syncthreads()` function. In addition, a 1D design assigns more work per
thread, which results in a better instruction-level parallelism (ILP).

We conducted a performance test for both design choices to quantify the
differences in the two techniques and to verify our analysis. For every algorithm, we designed two kernels based on both configurations. The matrix is stored in shared memory in both situations. Figures 1 through 3 show the performance for the three factorizations using both single and double precision arithmetic. All performance graphs show a clear advantage for the 1D configuration over the 2D configuration. Figure 1 shows speedups of $10.5 \times / 9.6 \times$ in single/double precision against the 2D configuration for the Cholesky factorization. A similar $9 \times$ (or more) speedup is obtained for the LU factorization in both precisions (Figure 2). Figure 3 also shows similar significant speedups for the QR factorization. It is clear that, regardless of the numerical algorithm, the 1D configuration is the winning strategy for batch workloads of small sizes. From now on, we continue to carry out further enhancements on the 1D configuration.

Figure 2: Performance comparison between the 1D configuration and the 2D configuration of the LU factorization kernel. Results are for 1M matrices on a Pascal P100 GPU.

8. Design Choice 2: Matrix Storage

The second design parameter is the matrix storage in fast memory, which can be either the register file, or the shared memory. In general, the access time of registers is faster than shared memory, but the latter provides more flexible access patterns. In either case, the 1D array of threads reads the matrix column by column from the global memory to preserve a coalesced memory access. If
threads read into registers, each thread has direct access to an entire row of the matrix. In order to access other elements, communication among threads is required, either by using shuffle operations, or through shared memory. All three factorization algorithms require accessing data from other threads, e.g., as in the row interchanges in the LU factorization, and the rank-1 updates in all algorithms. On the other hand, if the threads read the matrix into shared memory, any thread has access to the entire matrix. This comes at the cost of synchronization points (recall that CUDA now deprecates implicit warp synchronous codes, especially on the Volta GPUs). In addition, shared memory is slower than registers, and its capacity is smaller than the register file (e.g., 64KB on the P100 GPU vs. a 256KB register file). The capacity of the chosen storage is important, since it directly impacts the occupancy on the multiprocessor. Since there is a tradeoff between the two design choices, we developed two versions for each factorization algorithm to better understand the consequences of each choice per algorithm and matrix size. The first version uses shared memory for storage and communication among threads. The second uses registers for storage and warp shuffle operations for communication.

First, we consider the Cholesky factorization algorithm. The shared memory version loads the matrix into shared memory column-by-column to respect coalesced global memory access. At each iteration $i$, all threads compute the square...
Figure 4: Performance comparison between performing the Cholesky factorization in registers and in shared memory. Results are for 1M matrices on a Pascal P100 GPU.

root of $A[i,i]$, scale the column $A[i+1:N,i]$ accordingly in shared memory, and then use the result to update the lower triangular part of the trailing matrix $A[i+1:N, i+1:N]$ in shared memory, column by column. Both the scale and the update phases are fenced by synchronization points (i.e. `__synce warp()`). On the other hand, the register version must start by a shuffle operation to broadcast $A[i,i]$ to all threads. After computing the square root, each thread scales its respective element in the current column. Before the update takes place, threads must perform shuffle operations to share the elements $A[i+1:N,i]$ among all threads. Afterwards, each thread runs independently to update its respective row. It is clear that synchronization points in the shared memory version are replaced by shuffle operation in the register version. Figure 4 shows the results for the two versions on the Cholesky factorization algorithm. For both precisions, the shared memory version fails to keep up the same performance with the register version. As the sizes become larger, the shared memory capacity and bandwidth become a bottleneck, which gives the advantage to the register version.

The LU factorization requires a pivot search at each iteration $i$. On shared memory, this can be done by a linear search on the elements of the column $A[i:N,i]$. In the register version, this step is done through a binary search, which can be implemented using the butterfly warp shuffle operations shown in
Figure 5: Butterfly shuffle operation in CUDA among 8 threads.

Figure 5. While this operation requires just $\log_2(N)$ steps, it must be carried out by a number of threads that is equal to power of 2. Another limitation is that shuffle operations support only 32 bit exchanges, which means that shuffles on double precision is twice the cost of single precision. Once the pivot is found, the swap operation can be done using shared memory or using shuffle operations. The scale and update operations of the LU algorithm are similar to that of the Cholesky algorithm except that the update requires communication of the pivot row $A[i,i+1:N]$, and that all the trailing matrix is updated. Figure 6 shows the performance of the two LU versions. We observe a similar to the Cholesky factorization case advantage for the register version over the shared memory version.

Figure 6: Performance comparison between performing the LU factorization in registers and in shared memory. Results are for 1M matrices on a Pascal P100 GPU.
Finally, for the QR factorization, we begin by discussing the results first, which are shown in Figure 7. Unlike the Cholesky and LU factorizations, we observe that the shared memory version is faster than the register version, which is the opposite of our initial expectations. The explanation of this finding requires a deeper analysis for the QR factorization. As explained in Algorithm 3, the DLARFG routine in the $i^{th}$ iteration begins with a reduction operation to compute the norm of the current column $A[i:N,i]$. The reduction step can be done in exactly the same two ways as the pivot search in the LU algorithm. The main difference, however, for the QR factorization comes in the update phase, where an extra matrix-vector multiplication is required before the rank-1 update. The multiplication computes $y = v^T C$, where $C = A[i:N, i+1:N]$. Assuming that $C$ has a size of $P \times Q$, the multiplication costs $2PQ$ FLOPs.

However, there becomes a difference in the number of reduction steps required compute $y$ depending on the storage type. If registers and butterfly shuffle operations are used, then all threads collaborate to compute one element of $y$ at a time. Therefore, assuming that $N_{p^2}$ is the nearest power of 2 larger than or equal to $N$, the total number of reduction steps is $Q \times \log_2(N_{p^2})$, and not $Q \times \log_2(P)$. Recall that shuffle operations require that the number of matrices is always equal to a power of 2. On the other hand, shared memory seems to provide a better alternative. If both $v$ and $C$ are in shared memory, we can assign each element of $y$ to a single thread. Since threads are totally parallel, and using $Q$ threads, it would take only $P$ steps to find the entire vector $y$. This means that it requires much fewer steps to compute $y$ in shared memory. Since the computation of $y$ is at the innermost loop of the kernel, the shared memory version can be faster than the register version.

The results of the QR factorization in Figure 7 motivated us to try a hybrid storage design, which means to use registers as the main storage for the matrix, but use shared memory (instead of shuffle operations) for communication among threads. Another motivation for this strategy is that CUDA 9.0 deprecates all the previous shuffle operations, and replaces them by new ones that are
Figure 7: Performance comparison between performing the QR factorization in registers and in shared memory. Results are for 1M matrices on a Pascal P100 GPU.

relatively slower (significantly slower on the Volta GPU). Figure 8 shows that the register version with shared memory communication beats the pure shared memory implementation. We tried the same technique with Cholesky and LU factorization, and barely found any impact on performance.

Figure 8: Performance comparison between performing the QR factorization using hybrid storage and using shared memory only. Results are for 1M matrices on a Pascal P100 GPU.

9. Design Choice 3: Concurrency Control

The general design strategy in Section 6 states that a matrix is processed using exactly one TB. However, it does not restrict the number of matrices assigned to the same TB. In fact, we can assign multiple matrices to the same...
TB to be factorized concurrently. The motivation behind this strategy is the 1D configuration of TBs, which brings very low occupancy if the matrix size is small, and in particular, if smaller than the warp size, which is the case at hand. For example, a batch of $8 \times 8$ matrices requires a TB configuration of $(8, 1)$ threads. Such configuration does not use a full warp, which obviously wastes resources. The second issue is that such a configuration makes the CUDA runtime in full control of the number of concurrent TBs per multiprocessor. Even if the runtime does the optimal decision, it will not be able to schedule more than 32 TBs per multiprocessor, which is the hardware limit on modern GPUs. Instead, we adopt a different configuration that aggregates multiple TBs into one, thus using an $(8, n_{TB})$ TB configuration, where $n_{TB}$ controls the number of concurrent factorizations per TB. Assuming an optimal runtime behavior with $n_{TB} = 4$, the 32 TBs will be able to factorize 128 matrices per multiprocessor instead of 32, thus achieving a $4 \times$ speedup. This example assumes that there is enough memory resources to host 128 matrices of size $8 \times 8$, which is a valid assumption for double precision if the register file is used for storage. The $n_{TB}$ value is a tuning parameter, with an optimal value that depends on many factors, including the matrix size and the storage type.

Figure 9 shows the general idea of concurrency control. In general, a batch of size $BC$ can be factorized using $\left\lceil \frac{BC}{n_{TB}} \right\rceil$, so that each TB handles $n_{TB}$ matrices.
The value of the tuning parameter $nF_{TB}$ does not need to be known at compile time. We conducted an autotuning experiment that performs a full sweep over values of $nF_{TB}$ from 1 to 16. We observe that after this range, there is no gain in performance. Figures 10 through 12 show the impact of concurrency control on the three factorization algorithms in single and double precision.

![Figure 10: The impact of concurrency control on the Cholesky factorization kernel. Results are for 1M matrices on a Pascal P100 GPU.](image)

We observe that a tunable $nF_{TB}$ does not affect the performance for sizes larger than 16. For such range, the hardware rounds up the number of threads to use a full warp anyway, which means that there are no wasted resources in the TB configuration. It also means that, as long as full warps are used, the runtime is able to schedule TBs efficiently among the multiprocessors. For sizes 1 through 16, we observe performance gains in both single and double precision. The smaller the sizes, the more the speedups. For example, the Cholesky factorization has speedups that range from $1.2 \times$ to $4.3 \times$ in single precision and from $1.2 \times$ to $3.9 \times$ in double precision. The impact is more apparent in the LU and QR factorizations, with the LU factorization enjoying speedups that range from $2.0 \times$ to $3.5 \times$ in single precision, and from $1.8 \times$ to $3.7 \times$ in double precision. Similarly, the QR factorization performance is enhanced by factors that range from $2.0 \times$ to $3.8 \times$ in single precision, and from $1.8 \times$ to $3.8 \times$ in double precision.
10. Lazy Swapping for LU Factorization

The LU factorization has a unique step that is necessary for its numerical stability. At each iteration, two rows potentially exchange their positions in the matrix. This is an expensive step with pure data movement, and zero arithmetic intensity. Originally, the exchange occurs at each iteration in a greedy style. While the greedy swap is necessary for general sizes, we can take advantage of the small sizes of interest and introduce a lazy swap technique, where all the interchanges are carried once just before writing the matrix back into the global memory. Instead of explicitly changing positions in registers, each thread keeps
track of the final position of its row. A row that has been pivoted in a previous
iteration is marked with a flag so that it is excluded from the trailing matrix
update, as shown in Figure 13. Pivoted rows are also excluded from the pivot
searches in the following factorization steps. Figure 14 shows the performance
gains of the lazy swap technique, that are up to 21% and 32% in single and
double precision, respectively.

Figure 13: Trailing matrix updates in LU factorization with/without swapping.

(a) Single precision
(b) Double precision

Figure 14: The impact of the lazy swap on the LU factorization. Results are for 1M matrices
on a Pascal P100 GPU.

11. Final Performance Results

In this section, we show the final performance results of the three algorithms
against the competitive routines from the vendor library (cuBLAS). All kernels
are tuned and scheduled for the next release of the MAGMA library. The cuBLAS library provides batched routines for the LU and QR factorizations, but not for the batched Cholesky factorization. Therefore, the results for the Cholesky factorization use the cuBLAS batched LU routine instead, which is still a valid option (though suboptimal) to factorize SPD matrices. We also include a reference batched CPU implementation that we developed using sequential calls to the factorizations in the MKL library (version 11.3.0) from within an OpenMP `parallel for` loop. The number of OpenMP threads is set to 20, which is equivalent to the core count of the CPU.

![Figure 15](chart.png)

Figure 15: Final performance of the Cholesky factorization on the P100 GPU (batchCount = 1M).

![Figure 16](chart2.png)

Figure 16: Final performance of the Cholesky factorization on the V100 GPU (batchCount = 1M).
A normal behavior that we observe in most performance graphs are the spikes at sizes that are power of 2, and the drops that come right after those sizes. This is a normal behavior in GPUs, which always execute threads in groups of 32 (warps). A size that is a power of 2 allows the hardware to use full warps (especially when using the concurrency control technique discussed in Section 9). For other sizes, a round up to the nearest power of 2 is imposed either by the kernel configuration, or implicitly by the hardware. Even on the Volta V100 GPU, which has separate program counters and stacks for every thread, the same behavior is still observed.

Figure 15 shows the performance of the Cholesky factorization on the P100 GPU. The MAGMA kernels are significantly faster than cuBLAS (which uses the LU algorithm), scoring speedups that are up to $8.8 \times$ in single precision, and $8.2 \times$ in double precision. Against the reference CPU implementation, MAGMA is at least $9.3 \times$ faster in both precisions. The performance of the Cholesky factorization on the V100 GPU is also summarized in Figure 16, where we observe up to $5.7 \times / 4.1 \times$ speedups against cuBLAS in single/double precision. The MAGMA performance numbers on the V100 GPU are at least $13.2 \times$ faster than the CPU performance.

Considering the results for the LU factorization on the P100 GPU, which are summarized in Figure 17, the MAGMA kernel is faster than cuBLAS by up to $4.8 \times$ in single precision, and $5.2 \times$ in double precision. It is at least $14.3 \times$ faster than the CPU reference implementation. Looking into the results on the V100 GPU, MAGMA is still faster than cuBLAS, scoring speedups that are up to $3.8 \times$ in single precision, and up to $2.9 \times$ in double precision. The MAGMA performance on the V100 GPU is at least $22 \times$ faster than the CPU performance.

The performance gains in the QR factorization results are also significant. Considering the P100 GPU (Figure 19), MAGMA is $2.1 \times -11.04 \times$ faster than cuBLAS in single precision. It is also $2.9 \times$ to $9.4 \times$ faster in double precision. The speedup against the reference implementation is at least $16.1 \times$. We observe a kind of stagnation in the double precision performance for MAGMA. Recall
that this kernel uses both registers and shared memory to achieve the best performance. The shared memory requirements for this kernel \((N \times N)\) seem to limit the multiprocessor occupancy. The results in single precision support this explanation, since we observe no stagnation for the single precision results, as the shared memory requirements are almost half of what is needed in double precision. Considering the results on the V100 GPU (Figure 18), MAGMA is up to 11.8\times faster than cuBLAS in single precision. It is also up to 10.8\times faster in double precision. The minimum speedup obtained against the CPU reference implementation is 32.95\times across both precisions.
Figure 19: Final performance of the QR factorization on the P100 GPU (\texttt{batchCount} = 1M).

Figure 20: Final performance of the QR factorization on the V100 GPU (\texttt{batchCount} = 1M).

12. Conclusion and Future Work

This paper introduced a progressive design methodology for optimizing batched one-sided factorizations using GPUs. The paper mainly addresses extremely small matrix sizes, and introduces design techniques that are different from those used for larger matrices. Significant performance gains were achieved against the vendor library. The proposed work is of great importance for scientific applications, including astronomy, sparse multifrontal solvers, and preconditioners. Future directions include variable size batched workloads, developing an auto-tuning framework for such kernels, and integration with scientific applications where high performance batched routines have a great impact on performance.
Acknowledgements

This work is partially supported by NSF Grants SI2:SSE 1740250 and CSR 1514286, NVIDIA, and by the Exascale Computing Project (17-SC-20-SC), a collaborative effort of two U.S. Department of Energy organizations (Office of Science and the National Nuclear Security Administration) responsible for the planning and preparation of a capable exascale ecosystem, including software, applications, hardware, advanced system engineering and early testbed platforms, in support of the nations exascale computing imperative.

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