

# Mixed-Precision Orthogonalization Scheme and Adaptive Step Size for CA-GMRES on GPUs

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**Abstract.** We propose a mixed-precision orthogonalization scheme that takes the input matrix in a standard 32 or 64-bit floating-point precision, but accumulates its intermediate results in the doubled-precision. For a 64-bit input matrix, we use software emulation for the higher-precision arithmetics. Compared with the standard orthogonalization scheme, we require about  $8.5\times$  more computation but a much smaller increase in communication. Since the computation is becoming less expensive compared to the communication on new and emerging architectures, the relative cost of our mixed-precision scheme is decreasing. Our case studies with CA-GMRES on a GPU demonstrate that using mixed-precision for this small but critical segment of CA-GMRES can improve not only its overall numerical stability but also, in some cases, its performance. We also study an adaptive scheme to dynamically adjust the step size of the matrix powers kernel. Our experiments on multiple GPUs show that a near optimal step size can be chosen based on the performance measurements from the first restart loop of CA-GMRES.

## 1 Introduction

On modern computers, communication (e.g., data movement through memory hierarchies) is becoming expensive compared to computation (i.e., floating point operations). It is critical to take this hardware trend into consideration when designing high-performance software for new and emerging computers. To this end, we previously studied a communication-avoiding variant of the Generalized Minimum Residual method [3] (CA-GMRES) on multicore CPUs with multiple GPUs [5], focusing on the orthogonalization (*Orth*) and matrix powers kernels (*MPK*). We found that the Cholesky QR (CholQR) orthogonalization [4] obtains a superior performance based on the optimized BLAS-3 GPU kernels. Unfortunately, CholQR can be numerically unstable, and CA-GMRES may not converge even with reorthogonalization. We also found that depending on the matrix sparsity pattern, *MPK* can be faster than the standard algorithm, but it can be also slower due to the computational and/or communication overheads traded for reducing the communication latency. This is especially true with a relatively large step size preferred by *Orth*.

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 $\hat{\mathbf{x}} := \mathbf{0}$  and  $\mathbf{v}_1 := \mathbf{b}/\|\mathbf{b}\|_2$ .
repeat (restart-loop)
  Projection Subspace Generation on GPUs (inner-loop):
  for  $j = 1, s + 1, 2s + 1, \dots, m$  do
    MPK: Generate new vectors  $\mathbf{v}_{k+1} := A\mathbf{v}_k$ 
      for  $k = j, j + 1, \dots, \min(j + s, m)$ .
    BOrth: Orthogonalize  $V_{j+1:j+s+1}$  against  $V_{1:j}$ .
    TSQR: Orthogonalize the vectors within  $V_{j+1:j+s+1}$ .
  end for

  Projected Subsystem Solution on CPUs (restart):
  Compute the solution  $\hat{\mathbf{x}}$  in the generated subspace,
  which minimizes its residual norm.
  Set  $\mathbf{v}_1 := \mathbf{r}/\|\mathbf{r}\|_2$ , where  $\mathbf{r} := \mathbf{b} - A\hat{\mathbf{x}}$ .
until solution convergence do

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Fig. 1. CA-GMRES( $s, m$ ).

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Step 1: Gram-matrix formation
for  $d = 1, 2, \dots, n_g$  do
   $B^{(d)} := V_{1:s+1}^{(d)T} V_{1:s+1}^{(d)}$  on GPU
end for
 $B := \sum B^{(d)}$  (comm)

Step 2: Cholesky factorization
 $R := \text{chol}(B)$  on CPU

Step 3: Orthogonalization
for  $d = 1, 2, \dots, n_g$  do
  copy  $R$  to  $d$ -th GPU
   $V_{1:s+1}^{(d)} := V_{1:s+1}^{(d)} R^{-1}$  on GPU
end for

```

Fig. 2. CholQR.

To address the aforementioned deficiencies, in this paper, we first design and study a mixed-precision CholQR that takes the input matrix in a standard precision but accumulates its intermediate results in a higher-precision. Compared with the standard scheme, our mixed-precision scheme increases the computational cost by  $8.5\times$  but the increase in its communication cost is less significant. Since the computation is becoming less expensive compared to the communication on new and emerging architectures, we hope to improve the overall numerical stability of CA-GMRES using the mixed-precision without a significant increase in the orthogonalization time. Case studies on different GPUs demonstrate that this mixed-precision scheme can improve not only the stability of CA-GMRES but also, in some cases, its performance by allowing a larger step size, avoiding the reorthogonalization, and improving the convergence rate. We then study an adaptive scheme to dynamically adjust the step size of *MPK*. We demonstrate that a near optimal step size can be found based on the performance measurements from the first restart loop of CA-GMRES.

## 2 Communication-Avoiding GMRES

The  $j$ -th GMRES iteration generates the  $(j + 1)$ -th Krylov basis vector  $\mathbf{v}_{j+1}$  through a sparse matrix-vector multiply (*SpMV*) followed by its orthonormalization (*Orth*) against the previously-generated basis vectors. In our implementation [5], the coefficient matrix  $A$  and the basis vectors are distributed in a block row format among the GPUs on a compute node. We then generate these basis vectors on the GPUs, while the projected subsystem is solved on the CPUs.

Even on a single GPU, both *SpMV* and *Orth* require communication to move the data through the memory hierarchy of the GPU. CA-GMRES [2] aims to reduce this communication by replacing *SpMV* and *Orth* with three new kernels – matrix powers kernel (*MPK*), block orthogonalization (*BOrth*), and tall-skinny QR (*TSQR*) – that generate and orthogonalize a set of  $s$  basis vectors at once. Even on one GPU, CA-GMRES can obtain a speedup of up to two [5]. Figure 1 shows the pseudocode of CA-GMRES( $s, m$ ), where  $\mathbf{v}_j$  and  $V_{j:k}$  are the  $j$ -th

dd-multiply	# of d-instructions				dd-addition	# of d-instructions			
	Add/Sub	Mul	FMA	Total		Add/Sub	Mul	FMA	Total
dd-input	5	3	1	9	IEEE-style	20	0	0	20
d-input	3	1	1	5	Cray-style	11	0	0	11

**Fig. 3.** Number of double-precision instructions in double-double operations.

column and the submatrix consisting of the  $j$ -th through the  $k$ -th columns of  $V$ , respectively, and the iteration is restarted after  $m+1$  basis vectors are generated.

### 3 Cholesky QR factorization

In this section, we use  $V_{1:s+1}^{(d)}$  to denote the local matrix of  $V_{1:s+1}$  on the  $d$ -th GPU, and  $n_g$  is the number of available GPUs. To orthogonalize the  $s+1$  vectors  $V_{1:s+1}$  (as for  $TSQR$ ), CholQR first forms the Gram matrix  $B := V_{1:s+1}^T V_{1:s+1}$  through the matrix-matrix product  $B^{(d)} := V_{1:s+1}^{(d)T} V_{1:s+1}^{(d)}$  on the GPU, followed by the reduction  $B := \sum_{d=1}^{n_g} B^{(d)}$  on the CPU. Next, the Cholesky factor  $R$  of  $B$  is computed on the CPU. Finally, the GPU orthogonalizes  $V_{1:s+1}$  by a triangular solve  $V_{1:s+1}^{(d)} := V_{1:s+1}^{(d)} R^{-1}$ . Hence, all the required GPU-GPU communication is aggregated into a pair of messages, while the GPU computation is based on BLAS-3. Hence, both intra and inter GPU communication can be optimized. Figure 2 shows these three steps of CholQR. Unfortunately, the condition number of  $B$ ,  $\kappa(B)$ , is the square of  $\kappa(V_{1:s+1})$ . This often causes numerical instability, especially in CA-GMRES, where even using the Newton basis, the vector  $\mathbf{v}_j$  can converge to the principal eigenvector of  $A$ , and  $\kappa(V_{1:s+1})$  can be large.

## 4 Mixed-Precision Orthogonalization Scheme

### 4.1 Implementation

To improve the numerical stability of CholQR, we use the doubled-precision at the first two steps of CholQR, while the last step is in the standard precision. When the target hardware does not support a desired higher precision, software emulation is needed. For instance, double-double (dd) precision emulates the quadruple precision by representing each numerical value by an unevaluated sum of two double precision numbers, and is capable of representing the 106 bits precision, while the double-precision number is of 53 bits precision. There are two standard implementations [1] of adding two numerical values in dd-precision,  $a + b = \hat{c} + e$ , where  $e$  is the round-off error; one satisfies the IEEE-style error bound ( $e = \delta(a + b)$  with  $|\delta| \leq 2\epsilon_{dd}$  and  $\epsilon_{dd} = 2^{-105}$ ), and the other satisfies the weaker Cray-style error bound ( $e = \delta_1 a + \delta_2 b$  with  $|\delta_1|, |\delta_2| \leq \epsilon_{dd}$ ). Table 3 lists the computational costs of the dd-arithmetics required by our mixed-precision dd-CholQR. The standard CholQR in double precision (d-CholQR) performs about a half of its total flops at Step 1 and the other half at Step 3. Hence, using dd-precision for Steps 1 and 2, our dd-CholQR performs about  $8.5\times$  more

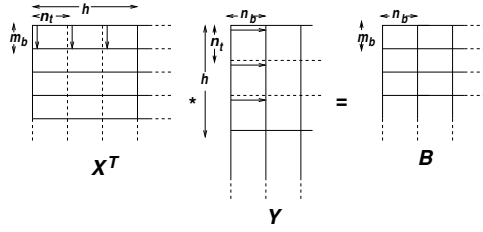


Fig. 4. *InnerProds* implementation (arrow shows data access by a GPU thread).

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double regC[m_b][n_b], regA[m_b], regB
for  $\ell = 1, \dots, \frac{h}{n_t}$  do
  for  $j = 1 \dots n_b$ 
    regA[i] =  $x_{\ell * n_t, j}$ 
  for  $j = 1, \dots, n_b$  do
    regB =  $y_{\ell * n_t, j}$ 
    for  $i = 1 \dots m_b$ 
      regC[i][j] += regA[i] * regB
    end for
  end for
end for

```

Fig. 5. *InnerProds* pseudocode.

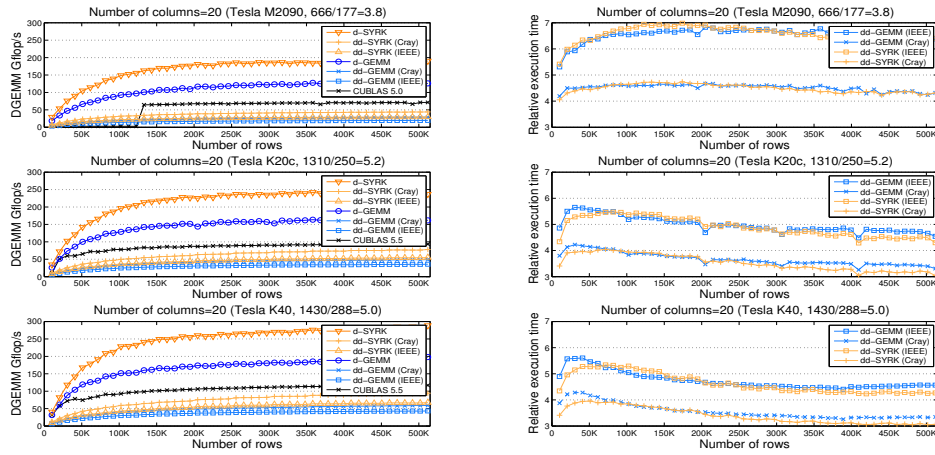


Fig. 6. Performance of *InnerProds* in double and double-double precision.

computation than d-CholQR. On the other hand, the increase in communication is less significant; our intra-GPU communication is about the same, only writing the  $s$ -by- $s$  output matrix in the dd-precision while reading the  $n$ -by- $s$  input matrix in the d-precision ( $s \ll n$ ). We communicate twice more data between the GPUs ( $16s^2$ Bytes with  $s \approx 10$ ), but with the same latency.

Since the Gram matrix is much smaller in its dimension than the coefficient matrix ( $s \ll n$ ), CholQR typically spends only a small portion of its orthogonalization time computing its Cholesky factor at Step 2. In addition, solving the triangular system with many right-hand-sides at Step 3 exhibits a high parallelism and can be implemented efficiently on a GPU. On the other hand, at Step 1, computing each element of the Gram matrix requires a reduction operation on  $n$ -length vectors. These inner-products (*InnerProds*) are communication-intensive and exhibit only limited parallelism. Hence, Step 1 often becomes the bottleneck, where standard implementations fail to obtain high-performance on the GPU. In our implementation of a matrix-matrix multiply (GEMM) to com-

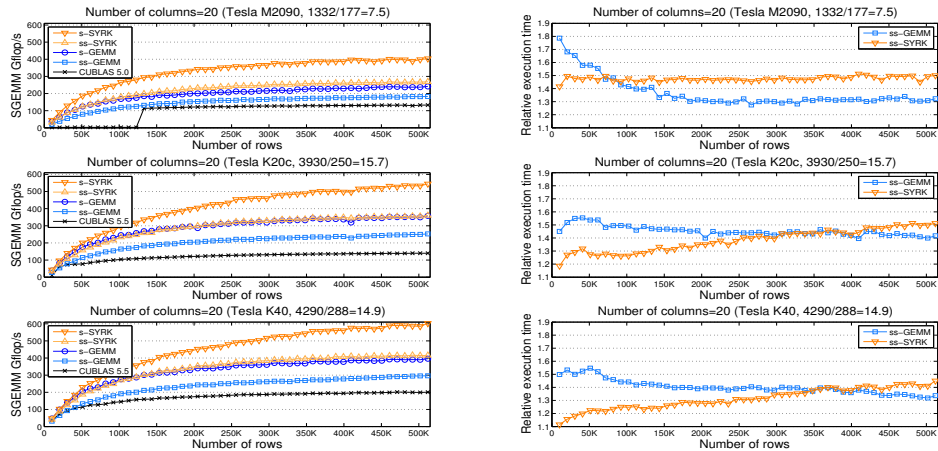


Fig. 7. Performance of *InnerProds* in single and single-single precision.

pute *InnerProds*,  $B := X^T Y$ , each thread block computes a partial product,  $B^{(i,j,k)} := X^{(k,i)T} Y^{(k,j)}$ , where  $X^{(k,i)}$  and  $Y^{(k,j)}$  are  $h$ -by- $m_b$  and  $h$ -by- $n_b$  blocks of  $X$  and  $Y$ , respectively. Within the thread block, each of its  $n_t$  threads computes its partial result in the local registries (see Figure 4 for an illustration, and Figure 5 for the pseudocode, where  $x_{\ell,j}$  is the  $(\ell, j)$ -th element of  $X^{(k,i)}$ ). Then, each thread block performs the binary reduction of the partial results among its threads, summing  $n_r$  columns at a time using the shared memory to store  $n_t \times (m_b \times n_r)$  numerical values. The final result is computed by another binary reduction among the thread blocks. Our implementation is designed to reduce the number of synchronizations among the threads while relying on the CUDA runtime and the parameter tuning to exploit the data locality.<sup>1</sup> For the symmetric (SYRK) multiply,  $B := V^T V$ , the thread blocks compute only a triangular part of  $B$  and reads  $V^{(k,j)}$  once for computing a diagonal block.

## 4.2 Performance

Figure 6 compares the *InnerProds* performance in d- and dd-precisions on different GPUs. Each GPU has a different relative cost of communication to computation, and on top of each plot, we show the ratio of the double-precision peak performance (Gflop/s) over the shared memory bandwidth (GB/s) (i.e., flop/B to obtain the peak). This ratio tends to increase on a newer architecture, indicating a greater relative communication cost. We tuned our kernel for each matrix dimension on each GPU in each precision (see the five tunable parameters

<sup>1</sup> In the current implementation, the numbers of rows and columns in  $X$  and  $Y$  are a multiple of  $h$ , and multiples of  $m_b$  and  $n_b$ , respectively, where  $n_b$  is a multiple of  $n_r$ .

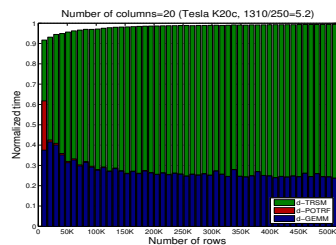


Fig. 8. d-CholQR time breakdown.

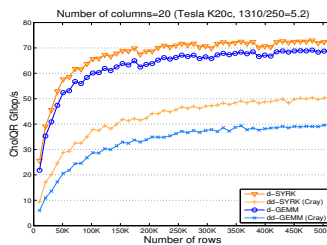


Fig. 9. d/dd-CholQR performance.

$h$ ,  $m_b$ ,  $n_b$ ,  $n_r$ , and  $n_t$  in Section 4.1), and the figure shows the optimal performance. Based on the shared memory bandwidth in the figure, the respective peak performances of d-GEMM are 442, 625, and 720Gflop/s on M2090, K20c, and K40. Our d-GEMM obtained 29, 26, 28% of these peak performances and speedups of about 1.8, 1.7, and 1.7 over CUBLAS 5.5 on these GPUs. In addition, though it performs  $16\times$  more instructions, the gap between *dd-InnerProds* and *d-InnerProds* tends to decrease on a newer architecture with a lower computational cost, and *dd-InnerProds* is only less than four times slower on K20c.

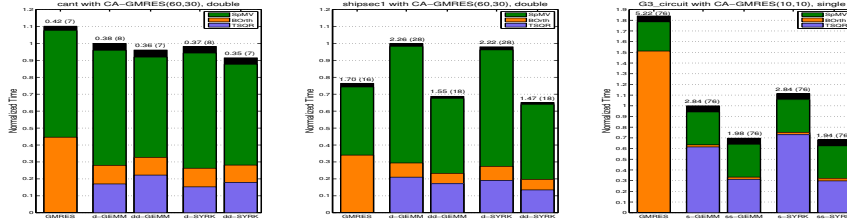
Figure 6 shows the performance of the mixed-precision *ss-InnerProds* in single precision, where the input matrix is in single precision, but the intermediate results are computed in double precision. Since the higher-precision is now supported by the hardware, even on an older GPU, the relative performance of the mixed-precision *ss-InnerProds* and the standard *s-InnerProds* is much closer than that of *dd-InnerProds* and *d-InnerProds*, where the software emulation is needed. In addition, *ss-InnerProds* is significantly more efficient obtaining over 300Gflop/s, in comparison to *d-InnerProds* that obtains just over 150Gflop/s.

Figure 8 shows the breakdown of d-CholQR orthogonalization time. Because of our efficient implementation of *InnerProds*, only about 30% of the orthogonalization time is now spent in *d-InnerProds*. As a result, while *dd-InnerProds* was about three times more expensive than *d-InnerProds*, Figure 9 shows that dd-CholQR is only about 1.7 or 1.4 times more expensive than d-CholQR when GEMM or SYRK is used for *InnerProds*, respectively. For dd-CholQR, the Cholesky factorization in dd-precision is computed using MPACK<sup>2</sup> on the CPU.

### 4.3 Case Studies with CA-GMRES

Figure 10 shows the normalized solution time of CA-GMRES with a single K20c for three sparse matrices from UF Sparse Matrix collection. Using dd-CholQR in the first two cases, even with the computationally expensive software emulation, the solution time was reduced not only because the reorthogonalization was avoided but also because CA-GMRES converged in fewer iterations. In the last case, due to the hardware support for the higher-precision, the solution time was reduced even though CA-GMRES converged with the same number of iterations.

<sup>2</sup> <http://mplapack.sourceforge.net>



**Fig. 10.** CA-GMRES Performance: On top of each bar shows total time in sec. and restart count. CA-GMRES with non-optimal  $s$  got speedups over GMRES using CGS.

## 5 Adaptive Step Size for Matrix Powers Kernel

Most of CA-GMRES implementations including ours [5] use the same step size  $s$  for  $MPK$ ,  $BOrth$ , and  $TSQR$ , while the optimal  $s$  for  $MPK$  is typically smaller than that of  $BOrth$  or  $TSQR$ . Instead of having a different  $s$  for  $MPK$  as an input, we design an adaptive scheme to dynamically adjust the step size  $\hat{s}$  of  $MPK$  based on the performance measurements from the first restart-loop of CA-GMRES.<sup>3</sup> For this, we use the following performance model:

$$MPK \text{ time} = \text{Inter-communication time} + \text{Kernel time},$$

where we let

$$\text{Inter-communication time} = \text{Latency} + \frac{\text{Communication volume}}{\text{Bandwidth}}, \text{ and}$$

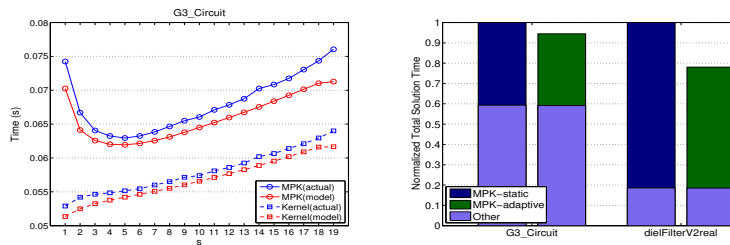
$$\text{Kernel time} = \frac{\text{Flop count}}{\text{flop/s}} + \# \text{ of random data accesses} \times \text{Data access time}.$$

In our experiments, we computed “Latency,” “Bandwidth,” “flop/s,” and “Data access time” based on the measured time of the reduction for the dot-products, point-to-point communication for  $SpMV$ ,  $SpMV$ , and data copy on the GPU, respectively. In addition, “# of random data accesses” is approximated by the aggregated number of non-local vector element accesses by  $MPK$ . To generate the  $s$  basis vectors, we invoke  $MPK$   $s/\hat{s}$  times using the step size  $\hat{s}$  before calling  $BOrth$  and  $TSQR$ . Figure 11 shows the results on three Tesla M2090 GPUs.

## 6 Conclusion

We proposed a mixed-precision orthogonalization scheme to improve the numerical stability of CA-GMRES. Our case studies demonstrated that though it requires about  $8.5\times$  more computation, using a higher-precision for this small but critical segment of CA-GMRES can improve not only its overall stability

<sup>3</sup> Our first loop is GMRES since the shifts for the Newton basis are not available.



**Fig. 11.** Result of *MPK* Performance Model (left) and its effect on CA-GMRES (right).

but also, in some cases, its performance. We also showed that the overhead of a higher-precision is decreasing on a newer architecture with an increasingly lower cost of computation. Though we only studied the effects of a higher-precision on a single GPU, on multiple GPUs of a compute node, the performance of CA-GMRES depends more on the performance of the GPU kernels (i.e., intra-GPU communication) than the inter-GPU communication [5]. Hence, similar benefits of using a higher-precision are expected on the multiple GPUs. We will study its effects on a system with a greater communication latency (e.g., distributed GPUs or CPUs) where the improvement may be greater. We are also studying the use of mixed-precision in eigensolvers where the orthogonality can be more crucial, and are writing an extended paper focusing on the numerical properties of our mixed-precision scheme. Finally, it is of our interest to apply or extend recent mixed precision efforts (e.g., reproducible BLAS<sup>4</sup> and precision tuning<sup>5</sup>) for our studies. In this paper, we also studied an adaptive scheme to adjust the step size of *MPK* on multiple GPUs. Our *MPK* is currently optimized only for the inter-GPU communication which is relatively inexpensive on a node. We expect a greater benefit of the adaptive scheme (in term of time or memory) on a larger system with greater communication cost (e.g., a distributed system).

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<sup>4</sup> <http://www.eecs.berkeley.edu/~hdnguyen/rblas>

<sup>5</sup> <http://crd.lbl.gov/groups-depts/ftg/projects/current-projects/corvette/precimonious>