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Hierarchical Approach for Deriving a Reproducible unblocked LU factorization

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Abstract

We propose a reproducible variant of the unblocked LU factorization for graphics processor units (GPUs). For this purpose, we build upon Level-1/2 BLAS kernels that deliver correctly-rounded and reproducible results for the dot (inner) product, vector scaling, and the matrix-vector product. In addition, we draw a strategy to enhance the accuracy of the triangular solve via iterative refinement. Following a bottom-up approach, we finally construct a reproducible unblocked implementation of the LU factorization for GPUs, which accommodates partial pivoting for stability and can be eventually integrated in a high performance and stable algorithm for the (blocked) LU factorization.

Keywords: LU factorization, BLAS, reproducibility, accuracy, long accumulator, error-free transformation, GPUs.

1. Introduction

The IEEE 754 standard, created in 1985 and then revised in 2008, has led to a considerable enhancement in the reliability of numerical computations by rigorously specifying the properties of floating-point arithmetic. This standard is now adopted by most processors, thus leading to a much better portability of numerical applications.

Exascale computing (10\textsuperscript{18} operations per second) is likely to be reached within a decade. For the type of systems yielding such performance rate, getting accurate and reproducible results in floating-point arithmetic will represent two considerable challenges \cite{9, 28}. Reproducibility is also an important and useful property when debugging and checking the correctness of codes as well as for legal issues.

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\textsuperscript{1}By accuracy, we mean the relative error between the exact result and the computed result. We define reproducibility as the ability to obtain a bit-wise identical floating-point result from multiple runs of the code on the same input data.

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The solution of a linear system of equations is often at the core of many scientific applications. Usually, this process relies upon the LU factorization, which is also its most compute-intensive part. Although there exist implementations of this factorization that deliver high performance on a variety of processor architectures—including general-purpose multicore processors, Intel Xeon Phi, and graphics processors (GPUs)—their reproducibility and, even worst, accuracy cannot be guaranteed. This problem is mainly due to the non-associativity of floating-point operations, combined with the concurrent thread-level execution of independent operations on conventional multi-core processors or the non-determinism of warp scheduling on many-core GPUs. This last type of architecture is especially appealing for the acceleration of compute-intensive kernels, as those appearing in dense linear algebra [13, 21].

In this work, we address the problem of reproducibility of the LU factorization on GPUs due to cancellations and rounding errors when dealing with floating-point arithmetic. Instead of developing a GPU implementation of the LU factorization from scratch, we rely on the hierarchical and modular structure of linear algebra libraries, and start by creating and augmenting reproducible OpenCL kernels for the BLAS (Basic Linear Algebra Subprograms [10]) that serve as building blocks in the LU factorization. In addition, we enhance the accuracy (in case of non-correctly-rounded results) of these underlying BLAS kernels for graphics accelerators.

We consider the unblocked left-looking algorithm for the LU factorization (also known as jik or jki variant [32]). The unblocked version is important for the solution of “batched” linear systems [14], where the goal is to solve a large sequence of independent small-size problems, and also as a building block to assemble high performance algorithms for the factorization. In addition, the left-looking version is especially appealing for fault tolerance, out-of-core computing, and the solution of linear systems when the coefficient matrix does not fit into the GPU memory. The unblocked left-looking algorithm can be formulated in terms of the Level-1 and Level-2 BLAS kernels for the dot product (dot), vector scaling (scal), matrix-vector product (gemv), and triangular system solve (trsv). We prevent cancellations and rounding errors in these kernels by applying the following techniques:

• We leverage a long accumulator and error-free transformations (EFTs) designed for the exact, i.e. reproducible and correctly-rounded, parallel reduction (exsum) [6] in order to derive an exact dot product (exdot). For this purpose, we combine the multi-level parallel reduction algorithm with the traditional EFT, called TwoProd [31], for the multiplication of two floating-point numbers.

• By its nature, scal is both reproducible and correctly-rounded. However, in the unblocked left-looking factorization, scal multiplies a vector by the inverse of a diagonal element, which causes two rounding errors (one to compute the inverse and one for the multiplication by the inverse). To address this issue, we provide an extension of scal (invscal) that performs the division directly, ensuring correctly-rounded and reproducible results.
We develop a reproducible and accurate implementation of GEMV by combining together a high performance GPU kernel of this operation with the exact DOT.

To improve the parallel performance of TRSV, we use a blocked variant that relies upon small TRSV involving the diagonal blocks and rectangular GEMV with the off-diagonal blocks. This approach leads to a reproducible, but not yet correctly-rounded, triangular solve (EXTRSV) [18]. We tackle this accuracy problem by applying a few iterations of iterative refinement.

Finally, we integrate partial pivoting [13] into unblocked left-looking algorithm for the LU factorization which, as part of future work, will allow us to employ this component in the solution of batched linear systems as well as a building block for high performance blocked factorization algorithms.

The paper is organized as follows. Section 2 reviews several aspects of computer arithmetic, in particular floating-point expansions and the Kulisch superaccumulator. Section 3 presents the ExBLAS library with its current set of routines. Section 4 is devoted to the presentation of the proposed reproducible LU algorithm. We evaluate our implementations in Section 5. Finally, we discuss related works and draw conclusions in Sections 6 and 7, respectively.

2. Background

In this paper, we consider the double precision format (binary64) as specified in the IEEE-754 standard. This standard requires correctly-rounded results for the basic arithmetic operations (+, −, ·, /, \sqrt{ }) which means that the operations are performed as if the result was first computed using infinite precision, and then rounded to the floating-point format. In this work, we assume the rounding-to-nearest mode.

Due to rounding errors, floating-point operations are non-associative and, therefore, non-reproducible [22]. Hence, the accuracy and reproducibility of floating-point operations strongly depend on their order [16, 29]. As a consequence, dynamic thread scheduling, which is often exploited to improve the performance of parallel algorithms, may lead to different results from one execution to another.

In the remainder of this section we present a brief overview of algorithms that constitute the foundation of our present work. Concretely, floating-point expansions with EFTs (Section 2.1) and a Kulisch superaccumulator (Section 2.2) have been proposed in the past in order to perform addition/subtraction of floating-point numbers without round-off errors; and these two algorithms have been efficiently combined to derive the hierarchical scheme for parallel summation [6] (Section 3.1) and dot product [17] (Section 3.2).

2.1. Floating-Point Expansion

Floating-point Expansions (FPE) allows us to recover and track rounding error which occurs during floating-point additions. FPE represents the result as an unevaluated sum of $p$ floating-point numbers
whose components are ordered in magnitude with minimal overlap to cover a wide range of exponents. FPEs of sizes \( p = 2 \) and \( 4 \), based on the EFT, are described in [26] and [15], respectively. The conventional EFT for the addition (TwoSum) is given in Alg. 1 [24] and, for the multiplication (TwoProd), in Alg. 2 [31]. Alg. 1 computes the addition of two floating-point numbers \( a \) and \( b \) and returns the result \( r \) and the error \( e \) such that \( r \) and \( e \) do not overlap. Similarly, TwoProd performs the multiplication of two floating-point numbers \( a \) and \( b \). For TwoProd, we use the fused-multiply-and-add (FMA) instruction to track the error that computes \( a \cdot b - r \) with only one rounding at the end.

### Algorithm 1: EFT for the summation of two floating-point numbers.

\[
\begin{align*}
\text{Function } [r, e] &= \text{TwoSum}(a, b) \\
& \quad r := a + b \\
& \quad z := r - a \\
& \quad e := (a - (r - z)) + (b - z)
\end{align*}
\]

### Algorithm 2: EFT for the product of two floating-point numbers.

\[
\begin{align*}
\text{Function } [r, e] &= \text{TwoProd}(a, b) \\
& \quad r := a \cdot b \\
& \quad e := \text{FMA}(a, b, -r)
\end{align*}
\]

Adding one floating-point number to an expansion of size \( p \) is an iterative process. The floating-point number is first added to the head of the expansion and the rounding error is next recovered as a floating-point number using the TwoSum EFT. The error is then recursively accumulated to the remainder of the expansion. As long as the dynamic range of the sum is lower than \( 2^{53 \times p} \) (for \texttt{binary64}), the FPE approach computes the accumulation of numbers without loss of accuracy.

The performance advantage of FPEs is that they can be kept in registers (after being fetched) during the computations. However, their accuracy may be insufficient for large sums or for floating-point numbers with significant variations in magnitude. Additionally, the complexity of FPEs grows linearly with their size.

### 2.2. Kulisch Superaccumulator

The Kulisch superaccumulator covers the range from the minimum representable floating-point value to the maximum value in absolute value. For the dot product of two vectors composed of \texttt{binary64} elements, Kulisch [25] proposed to use a 4,288-bit accumulator. The addition of products of \texttt{binary64} values is performed without loss of information by accumulating every floating-point number in the superaccumulator; see Fig. [1]. The superaccumulator can produce the exact sum or dot product of a very large amount of floating-point numbers with arbitrary dynamic ranges. However, the superaccumulator incurs a large memory overhead, due to the required storage hindering vectorization due to indirect memory accesses.
3. Exact BLAS Library (ExBLAS) on GPUs

This section briefly reviews the prototype implementation of the Exact BLAS (ExBLAS) library [19] that underlies our LU factorization. It starts with the parallel reduction and dot product, as they are two fundamental BLAS kernels, and then continue with the triangular solver, outlining the improvements present in this paper compared with prior work. In addition, we extend the set of routines to the scaling of a vector by the inverse of a scalar and the matrix-vector product, which represents new reproducible algorithms and implementations. While presenting these routines, we also expose how they are used within the discussed unblocked LU factorization.

3.1. Exact Parallel Reduction: exsum

In [6, 19] we introduced a multi-level approach to the summation problem. It combines the FPE and superaccumulator algorithms. The procedure splits the computation into five levels: filtering, private superaccumulation, local superaccumulation, global superaccumulation, and rounding. This 5-stage decomposition exploit the nested parallelism of modern parallel architectures such as GPUs.

The first level of this hierarchical summation approach relies on FPEs with EFTs for the accumulation of floating-point numbers (see Algs. 1 and 2). Each thread maintains its own FPE. In order to enable expansions of size $p$, we apply Alg. 3 based on Alg. 1 enhanced with superaccumulators in case the result of the accumulation cannot be represented with a FPE of size $p$.


\begin{algorithm}
\begin{algorithmic}
\Function{ExpansionAccumulate}{$x$}
\For{$i = 0 \rightarrow p - 1$}
\State $(a_i, x) := \text{TwoSum}(a_i, x)$
\EndFor
\If{$x \neq 0$}
\State \text{Superaccumulate}($x$)
\EndIf
\EndFunction
\end{algorithmic}
\end{algorithm}

In the second level, if the last rounding error $x$ is non-zero (see Alg. 3), $x$ is forwarded to the private
superaccumulator, the FPE is flushed to the superaccumulator, and the accumulation process is continued. At the end of the summation, all FPEs are forwarded to superaccumulators. Depending on the amount of memory that is available, private superaccumulators (shared among a small number of threads) are allocated in either fast local memory, e.g. cache or shared memory, or global memory.

In the third level, \( k \) private superaccumulators are merged into a single local superaccumulator, one per group of threads. In the fourth level, all local superaccumulators within a GPU are combined together into a global superaccumulator. Finally, the global superaccumulator is rounded back to the target floating-point format, in the fifth level, in order to obtain the correctly-rounded result.

We developed hand-tuned variants of a single OpenCL implementation for NVIDIA and AMD GPUs. These variants use 16 superaccumulators per workgroup of 256 threads and employ local memory to store these superaccumulators. In order to avoid bank conflicts, superaccumulators are interleaved to spread their digits among different memory banks. Concurrency between 16 threads that share one superaccumulator is settled thanks to atomic operations while scattering the input data into the corresponding digits of the superaccumulator.

3.2. Exact Dot Product: exdot

We apply the multi-level parallel reduction algorithm [6] to the dot product [17] by additionally utilizing the TwoProd EFT for the exact multiplication of two floating-point numbers. The GPU implementation of exdot is based on the same concept that underlying exsum with an auxiliary function call to TwoProd and the corresponding treatment of its outputs. The performance results of the exact dot product on GPUs present a small overhead induced by the summation of two numbers (the result and the error) after each call to TwoProd. In contrast, for large array sizes, exdot delivers both numerically reproducible and correctly-rounded results with comparable performance to its standard non-deterministic version.

3.3. Exact Vector Scaling: exscal and exinvscal

Multiplying a vector \( x \) by a scalar \( \alpha \) is a rather simple kernel that requires only one operation to be performed for each element of the vector \( (x_i := \alpha \cdot x_i) \). When the IEEE 754-2008 compliance is ensured, this operation (exscal) is both correctly-rounded and reproducible. However, the unblocked LU factorization (Alg. 7) requires a call to the scal routine in order to scale the vector by the inverse of the diagonal element. In that case, the result is not correctly-rounded due to the two rounding errors coming from the division followed by the multiplication. To ensure correctly-rounded, and reproducible results, we propose a dedicated version of scal (exinvscal). This version directly performs division of the vector elements by the diagonal element without the intermediate rounding error. Both exscal and exinvscal are easy to implement on GPUs as the update of the vector elements can be performed in parallel by a team of threads.
3.4. Exact Matrix-Vector Product: exgemv

The matrix-vector product (gemv) is one of the building blocks for the triangular solver as well as for the unblocked LU factorization. Therefore, we next present its correctly-rounded and reproducible implementation. The gemv kernel computes one of the following matrix-vector operations:

\[ y := \alpha A \cdot x + \beta y, \quad \text{or} \quad y := \alpha A^T \cdot x + \beta y, \]  

(1)

where \( \alpha \) and \( \beta \) are scalars, \( x \) and \( y \) are vectors, and \( A \) is a matrix.

We derive a reproducible and accurate algorithm for exgemv by combining its two-kernel algorithmic variants from [3] to achieve high performance; and the exact dot product, which is described in Section 3.2, to guarantee both reproducibility and accuracy of the results. We next provide a description of the implementation as a GPU kernel.

![Figure 2: Work distribution in the matrix-vector product.](image)

The proposed OpenCL implementation of the matrix-vector product splits the computations into blocks, so that each workgroup of threads compute a certain part of the output vector \( y \). Fig. 2 shows how a part of the vector \( y \) is computed using \( p \) workgroups of size \( mb \). Each thread from a workgroup is engaged in pre-loading a part of \( x \) (colored in dark blue) into local memory, making it available to the whole workgroup. Then, each thread computes its partial dot product using its own FPE, which is kept in private memory; the computation on the thread-owned FPEs is very fast. The FPEs with the partial results are flushed to local superaccumulators that are stored in the corresponding cells of the \( m \times p \) matrix (dark green). Concerning the superaccumulators, we adhere to one of the following two scenarios:

1. Hold a matrix of \( m \times p \) superaccumulators in global memory and then perform a reduction on them in order to obtain a vector of \( m \) superaccumulators that corresponds to the output vector \( y \);
2. Hold only a vector of \( m \) superaccumulators, where each superaccumulator is shared among \( p \) threads, and solve the contention for shared superaccumulators using atomic operations.
The second scenario reduces the exgemv implementation to one kernel and releases the pressure on the
global memory. That is also beneficial for the \( n - 1 \) (\( n \) is a matrix size) calls to \textsc{exgemv} within the unblocked
LU factorization. The first scenario is left for future work.

Each workgroup holds only a part of the vector \( x \) in its local memory. This allows multiple workgroups
per multiprocessor to proceed concurrently. This strategy also maximizes the reuse of cached-shared parts
of the vector \( x \) as there are 256 or 512 threads per workgroup.

In the current version of ExBLAS, we provide implementations that cover all possible cases of \textsc{exgemv}
as depicted in \([1]\).

3.5. Reproducible Triangular Solver: \textsc{extrsv}

The triangular solve involving a lower triangular coefficient matrix is one of the building blocks for the
unblocked LU factorization. We next provide the multi-level reproducible approach for triangular systems
with a single right-hand side \([18]\) and reveal our strategies for improving its accuracy.

Let us consider the system \( Lx = b \), where \( L \) is a non-singular (square) lower triangular matrix. This
system can be solved using (sequential) forward substitution, as depicted in Alg. \([4]\) where the elements of \( x \)
are computed from first to last.

\begin{algorithm}
\textbf{Algorithm 4:} Forward substitution.
\begin{align*}
x_1 & := b_1 / l_{11} \\
\text{for } i = 2 : n \text{ do} \\
& \quad s := b_i \\
& \quad \text{for } j = 1 : i - 1 \text{ do} \\
& \quad \quad s := s - l_{ij} x_j \\
& \quad \text{end} \\
& \quad x_i := s / l_{ii} \\
\text{end}
\end{align*}
\end{algorithm}

Alg. \([5]\) represents the blocked algorithm for the reproducible triangular solver with a lower triangular
matrix. Our strategy depicted in this algorithm is the following:

1. Use a blocked version of the triangular solver;
2. Apply the exact dot product \( (s := s - l_{ij} x_j) \) with FPEs and superaccumulators. We apply this
procedure through the blocked \textsc{extrsv}; within the non-blocked (called local) \textsc{extrsv} and the local
blocked \textsc{exgemv}. The array of superaccumulators is shared between these two routines. Since the
computations in the blocked \textsc{extrsv} are performed in sequential order (local \textsc{extrsv}, local \textsc{exgemv},
local \textsc{extrsv}, etc.) there is no contention for the array of superaccumulators.
Algorithm 5: Blocked extrsv
blobsz stands for a block size; acc refers to an array of superaccumulators of size n; acc(k) corresponds to one superaccumulator at the position k in this array.

for i = 0 : blsz : n do
    Local extrsv
    for k = i : i + blsz do
        for j = 1 : k - 1 do
            [r, e] := TwoProd(lkj, −xj)
            ExpansionAccumulate(r) accumulate r using Alg. 3
            ExpansionAccumulate(e) accumulate e using Alg. 3
        end
        ExpansionAccumulate(bk)
        s := RNDN(acc(k)) round to double a superaccumulator
        xk := s/lkk
    end

    Local exgemv
    for k = i + blsz : n do
        for j = i : i + blsz do
            [r, e] := TwoProd(lkj, −xj)
            ExpansionAccumulate(r)
            ExpansionAccumulate(e)
        end
    end
end

3. Correctly round the accumulated result (s := RNDN(acc(k))). One can notice that rounding to double is performed only once – at the end of the computation of each element of the solution.

4. Perform the division by the corresponding diagonal element (x_i := s/l_{ii}).

Theorem 3.1. The previous strategy yields a reproducible solution for the triangular solve.

Proof. We give a proof by induction. As x_1 := b_1/l_{11} and as the division is correctly-rounded, the latter always returns the same result, and, therefore, it is reproducible. Let us now assume that x_1, ..., x_{i-1} are reproducible. As the computation s := s − l_{ij}x_j is done with a large accumulator, there is no rounding error and the result is exact, independently of the order of computation and, consequently, reproducible. The operation s := RNDN(s) is reproducible as it is a correctly-rounded operation. Finally x_i := s/l_{ii} is reproducible as it is the result of a correctly-rounded division between two reproducible quantities. □
With the proposed approach, the matrix, both the right-hand side and the solution vectors are first split into blocks. Then computations on those blocks is organized as in Figs. 3a and 3b. With this organization, each diagonal block of size $b\text{lsz} \times b\text{lsz}$ participates in $\text{extrsv}$, and the panel underneath this block is involved in $\text{exgemv}$. The challenge lies in ensuring a balanced workload distribution among workgroups on GPUs, and a light synchronization overhead, as some parts of the algorithm (the local $\text{extrsv}$ on the diagonal blocks, see Fig. 3b) still need to be executed in the sequential order.

Each of the four aforementioned parts of the proposed triangular solver is reproducible and, therefore, the computed solution is reproducible as well. Compared with our previous work [18], in the new version of ExBLAS we covered all the four variants of the triangular solver as follows

$$T \cdot x = b, \text{ or } T^T \cdot x = b,$$

where $x$ and $b$ are $n$ elements vectors, and $T$ is an $n \times n$ unit, or non-unit, upper or lower triangular matrix.

In order to enhance the accuracy of the reproducible $\text{trs}$, we propose to apply a few iterations of iterative refinement based on the ExBLAS routines, as described in Alg. 6. The overhead of the iterative refinement can be diminished by locating the source of the accuracy problem and, consequently, applying iterative refinement directly on the diagonal blocks right after the local $\text{extrsv}$.

**Theorem 3.2.** Algorithm 6, for the triangular solve with iterative refinement, is reproducible.

**Proof.** The first step uses $\text{extrsv}$ to compute $\hat{x} := L^{-1}b$, so $\hat{x}$ is reproducible. Inside the loop, we sequentially compute $r$, $d$ and $x$, with each of them being reproducible due to the use of $\text{extrsv}$, $\text{extrsv}$ and $\text{exaxpy}$, accordingly. As a consequence, the whole algorithm is reproducible. \( \square \)
Algorithm 6: The reproducible triangular solver with iterative refinement.

\[
\hat{x} := L^{-1}b \quad \text{(EXTRSV)}
\]

for \(i = 1 : \text{nbiter}\) do
  \[
  r := b - L\hat{x} \quad \text{(EXGEMV)}
  \]
  \[
  d := L^{-1}r \quad \text{(EXTRSV)}
  \]
  \[
  \hat{x} := \hat{x} + d \quad \text{(EXAXPY)}
  \]
end

4. Reproducible LU Factorization

The LU factorization decomposes an \(m \times n\) matrix \(A\) into the product of an \(m \times r\) unit lower triangular matrix \(L\), with \(r = \min(m, n)\); an \(r \times n\) upper triangular matrix \(U\); and an \(r \times r\) permutation \(P\) such that \(PA = LU\). In order to compute this decomposition, we consider an unblocked left-looking variant, illustrated in Alg. 7 using the FLAME notation [4, 12]. This notation makes it easier to identify which regions of the matrix are updated and used; see Fig. 4. In Alg. 7, \(\text{size}(A)\) represents the number of columns of the matrix \(A\). Before the computation starts, \(A\) is virtually partitioned into four blocks \(A_{TL}, A_{TR}, A_{BL},\) and \(A_{BR}\), where \(A_{TL}\) is a \(0 \times 0\) matrix. The matrix is then traversed from the top-left to the bottom-right corner. At each iteration of the loop, \(A\) is repartitioned from \(2 \times 2\) to \(3 \times 3\) form, where \(A_{00}, A_{02}, A_{20},\) and \(A_{22}\) are matrices; \(a_{01}, a_{10}^T, a_{12}^T,\) and \(a_{21}\) are vectors; and \(\alpha_{11}\) is a scalar. The algorithm updates \(a_{01}, \alpha_{11},\) and \(a_{21}\) using \(\text{TRSV}, \text{DOT},\) and \(\text{INVSCL/GEMV},\) respectively. At the end of the computation, matrix \(A\) is overwritten by the upper triangular matrix \(U\) and the unit lower triangular matrix \(L\). Additionally, the vector \(p\) of pivots (representing the permutations contained in \(P\)) is created.

Alg. 7 shows that, thanks to the modular and hierarchical structure of linear algebra libraries, computations of more complex algorithms – such as the LU factorization – can be entirely expressed and built on top of the lower level fundamental routines – in particular, the BLAS one. We benefit from this hierarchical organization to propose a reproducible algorithmic variant of the unblocked LU factorization, Alg. 7, on top of the underlying ExBLAS routines: \(\text{EXTRSV, EXDOT, EXINVSCL, and EXGEMV}\).

To enable support for (reproducible) partial pivoting in Alg. 7, we split this process into two stages:

1. Searching for the maximum element in absolute value within the subdiagonal part of a matrix column. This operation is always reproducible.
2. Swapping two rows. This operation is also reproducible by nature.

In addition, for this particular variant of the unblocked LU factorization, we apply pivoting from the previous iteration right before the computations update the current one.

**Theorem 4.1.** Algorithm 7, for the unblocked LU factorization with partial pivoting is reproducible.
Proof. The first and last step consists in partitioning the matrix $A$ which does not bring any reproducibility issues. All computational steps of the unblocked LU factorization rely upon their reproducible counterparts, such as extrsv, exdot, exinvscal, and exgemv, plus the reproducible strategy implementing partial pivoting. As a consequence, the result computed by Alg. 7 is numerically reproducible.

Therefore, we have successfully removed all sources of indeterminism, while efficiently exploiting data-parallelism within each basic block. Eventhough the accuracy of the result computed by Algorithm 7 is improved, it is not correctly rounded. This is due to the fact that a rounding step is executed between each reproducible BLAS operations. A solution to increase the accuracy is to perform some iteration of iterative refinement. However, we believe it is not possible to prove that correct rounding can be achieved even with iterative refinement. This is due to the fact that there exists obvious cases (division by 3, or multiplications by small numbers for example) where numerical informations that could be useful for rounding could be lost due to underflow.

5. Experimental Results

In this section, we evaluate our implementations of the unblocked LU factorization with partial pivoting and its underlying ExBLAS routines on three NVIDIA architectures; see Tab. 1.

<table>
<thead>
<tr>
<th>Hardware platforms employed in the experimental evaluation.</th>
</tr>
</thead>
<tbody>
<tr>
<td>NVIDIA Quadro K420 192 CUDA cores 0.780 GHz</td>
</tr>
<tr>
<td>NVIDIA Tesla K20c 2,496 CUDA cores 0.706 GHz</td>
</tr>
<tr>
<td>NVIDIA Tesla K80 4,992 CUDA cores with a dual-GPU design 0.560-0.875 GHz</td>
</tr>
</tbody>
</table>

We develop unique OpenCL implementations for each algorithm on GPUs and tuned these implementations – e.g. by promoting loop unrolling or changing workgroup size, etc. – for each particular architecture in order to optimize performance. Moreover, our implementations strive to deliver the best performance through utilizing all the underlying resources of the target GPUs: SIMD instructions, fmas, private and local memory, as well as atomic instructions. We check the correctness of our implementations by comparing the computed results with those produced by the multiple precision sequential library MPFR for CPUs on randomly generated test sets.

As a baseline for comparison purposes, we provide our vectorized, parallelized, and optimized nondeterministic double precision implementations of the matrix-vector product, triangular solver, and the unblocked LU factorization (Alg. 7). We denote these implementations on figures as “Parallel gemv”, “Parallel trsv”, and “Parallel LU”, accordingly. We use these implementations as starting points for the integration of our reproducible solutions. Hence, these baseline implementations are very relevant in order
Algorithm 7: The left-looking algorithmic variant of the unblocked algorithm LU factorization with partial pivoting.

**Partition**

\[
A \rightarrow \begin{pmatrix} A_{TL} & A_{TR} \\ A_{BL} & A_{BR} \end{pmatrix}, \quad p \rightarrow \begin{pmatrix} p_T \\ p_B \end{pmatrix}
\]

where \( A_{TL} \) is 0 \( \times \) 0, \( p_T \) has 0 elements

**While** \( \text{size}(A_{TL}) < \text{size}(A) \) do

**Repartition**

\[
\begin{pmatrix} A_{TL} & A_{TR} \\ A_{BL} & A_{BR} \end{pmatrix} \rightarrow \begin{pmatrix} A_{00} & a_{01} & A_{02} \\ \alpha_{11} & a_{21} & A_{22} \\ a_{T0} & a_{T1} & a_{T2} \end{pmatrix}, \quad \begin{pmatrix} p_T \\ p_B \end{pmatrix} \rightarrow \begin{pmatrix} p_0 \\ \pi_1 \\ p_2 \end{pmatrix}
\]

where \( \alpha_{11} \) and \( \pi_1 \) are scalars

\[
\begin{pmatrix} a_{01} \\ \alpha_{11} \\ a_{21} \end{pmatrix} := P(\begin{pmatrix} p_0 \\ \alpha_{11} \end{pmatrix}), \quad \begin{pmatrix} a_{01} \\ a_{11} \\ a_{21} \end{pmatrix} \text{ := } A^{-1}_{00} a_{01}
\]

\[
a_{01} := \frac{\alpha_{11}}{\alpha_{21}} \quad (\text{trsv})
\]

\[
\alpha_{11} := \alpha_{11} - a_{T0}^T a_{01} \quad (\text{dot})
\]

\[
a_{21} := a_{21} - A_{20} a_{01} \quad (\text{gemv})
\]

\[
\begin{pmatrix} \alpha_{11} \\ a_{21} \end{pmatrix} := P(\begin{pmatrix} \pi_1 \\ a_{21} \end{pmatrix}), \quad \begin{pmatrix} \alpha_{11} \\ a_{21} \end{pmatrix} \text{ := } \text{PivIndex} \begin{pmatrix} \alpha_{11} \\ a_{21} \end{pmatrix}
\]

\[
a_{21} := \frac{a_{21}}{\alpha_{11}} \quad (\text{scal/invscal})
\]

Continue with

\[
\begin{pmatrix} A_{TL} & A_{TR} \\ A_{BL} & A_{BR} \end{pmatrix} \leftarrow \begin{pmatrix} A_{00} & a_{01} & A_{02} \\ \alpha_{11} & a_{21} & A_{22} \\ a_{T0} & a_{T1} & a_{T2} \end{pmatrix}, \quad \begin{pmatrix} p_T \\ p_B \end{pmatrix} \leftarrow \begin{pmatrix} p_0 \\ \pi_1 \\ p_2 \end{pmatrix}
\]

endwhile

Figure 4: Partitioning of matrix \( A \).

To assess the performance, accuracy, and reproducibility of the results. Despite some performance penalties, we would like to emphasize the importance of obtaining reproducible and, if possible, correctly-rounded results.
5.1. EXGEMV Results

While decomposing a matrix into a lower and an upper triangular matrices using the LU factorization, the matrix-vector product involves matrices of various shapes, starting from column-panels, through squarish, and then row-panels. In our experiments, we aim to evaluate these scenarios by considering the following three test cases:

1. GEMV with square matrices;
2. GEMV with row-panel matrices, where the number of rows is fixed to \( m = 256 \) and the number of columns \( n \) varies;
3. GEMV with column-panel matrices, where the number of columns is fixed to \( n = 256 \) and the number of rows varies.
Fig. 5a presents the performance results achieved by the matrix-vector algorithms for square matrices \( m \times n \) \((m = n)\) composed of double precision floating-point numbers. In the caption of the following plots, “Superacc” corresponds to the accurate and reproducible matrix-vector algorithm that is solely based on superaccumulators; “exgemv” stands for our exact implementation, which delivers the best performance, with FPEs of size \( n \) \((n = 2 : 8)\) in conjunction with the \texttt{TwoProd} EFT and superaccumulators when required. The “Superacc” implementation, which can be classified as a brute force approach to ensure reproducibility, suffers from its extensive memory usage and is an order of magnitude slower than our solution.

The performance experiments using the latter two test cases are depicted in Figs. 5b and 5c respectively. Fig. 5c shows rather constant execution time for all \texttt{gemv} implementations. This is due to the use of a column-wide array of threads that performs local dot products within \texttt{gemv}. Fig. 5b demonstrates the outcome of applying this pool of threads iteratively over columns of the matrices of size \( n \times 256 \). This turns to be beneficial from the point of view of performance, as the overhead decreases with the number of columns. Fig. 5a and Fig. 5b exhibit very close relative performance results. The best performance among the \texttt{exgemv} implementations for all the test cases is delivered by “FPE3 + Superacc”. For instance, its overhead is 4.26 times for \( m = n = 4096 \) compared to the non-deterministic \texttt{gemv} implementation. To summarize, \texttt{exgemv} delivers both reproducible and correctly-rounded results for different shapes of both transpose and non-transpose matrices.

In addition, we tested the accuracy of the double precision \texttt{gemv} and the reproducible \texttt{exgemv}. Fig. 5d demonstrates the relative forward error \( (\|y^* - \hat{y}\|/\|y^*\|) \), where \( \hat{y} \) and \( y^* \) are the computed and exact vectors, accordingly) against the condition number of the problem. This condition number is computed with the MPFR library as a ration of norms of the exact matrix-vector products, when the matrix and the vector are in their absolute and original values: \( \|\|A\|\cdot|x\|/\|A\cdot x\|\| \). In order to generate the ill-conditioned matrix-vector, we rely on the ill-conditioned dot product. So that, the \( n - 1 \) rows of the matrix \( A \) are created as random uniformly distributed numbers, e.g. between 0 and 1, while one row and the vector \( x \) correspond to the ill-conditioned dot product. The exact vector \( y^* \) is computed with the MPFR library and rounded to the double precision vector. For visual representation, we replaced all the errors that exceed 1 by 1 as there is zero digit of accuracy left. Naturally, the relative forward error of \texttt{gemv} strongly depends on the condition number and indicates the incorrectness of the computed results once the condition number reaches \( 10^{16} \). Instead, our reproducible \texttt{exgemv} ensures both correctly-rounded and reproducible results as our approach preserves every bit of the computed result until its final rounding.

5.2. extrsv Results

Fig. 6a shows a performance improvement around 35 %, compared with the results in [18], delivered by \texttt{extrsv} on K20c. This improvement is due to a new way we are handling errors produced by one of the two outputs of \texttt{TwoProd}. In [18] these errors were accumulated to the FPE starting from its head. However,
most of the time this solution were sub-optimal as the error were much smaller that the head of the FPE, leading to further error propagation along the FPE. Based on that report, we decided to add these errors closer to the tail of the expansion, namely in the last two or three slots of the expansion. The difference in terms of performance among FPEs of various sizes is less apparent as they follow the same pattern with respect to the error propagation.

Figs. 6a to 6c shows that there is still work to do on architecture-specific implementations, since for large matrices, extrsv on K420 only reports a 4× overhead, while on both K20c and K80 the overhead reaches 25×. Although OpenCL ensures portability, unfortunately, it does not come with performance portability. We have already tuned our code on higher level for various GPU architectures. However, it is clear that, for some kernels as extrsv, additional architecture specific implementations should to be provided.

Creffig:extrsv:acc:old provides a comparison for the relative forward error versus the condition number of

*Figure 6: Performance and accuracy results of trsv.*
the double precision, \texttt{trsv}, with the proposed reproducible, \texttt{extrsv}, substitution algorithms. The relative forward error corresponds to $\|x^* - \hat{x}\|/\|x^*\|$, where $\hat{x}$ and $x^*$ are respectively to the computed and exact solutions. We use the following Skeel formula \cite{34,16} to calculate the condition number of a linear system $Ax = b$ with real coefficients:

$$\text{cond}(A, x) = \frac{\|A^{-1}\| \|A\| \|x\|}{\|x\|}.$$ 

To perform those tests, we wrote a random generator of ill-conditioned triangular systems using the algorithm proposed in \cite{27}; the random generator ensures that both the matrix $L$ and the right-hand side vector $b$ are composed of double precision floating-point numbers. In order to compute the exact solution $x^*$, we rely on the MPFR library and rounded the result back to double precision. For the errors greater than 1, we cut them to 1 as the results are obviously incorrect. This experiment reveals that the relative forward error is proportional to the condition number and most importantly to the rounding error $u = 2^{-53}$. Altogether \texttt{extrsv} delivers most of the time better accuracy as the double precision triangular solver. Indeed, the accuracy of the classic triangular solver can be improved via double-double precision. However, this approach is already 9 slower than \texttt{trsv} and, moreover, it does not provide any guarantees on the reproducibility of the results. Therefore, we propose to apply iterative refinement, see Alg. \ref{alg:6}, in order to improve the accuracy of \texttt{extrsv}.

5.3. \texttt{exlu} Results

During the development of the reproducible LU factorization, we discovered a limitation of OpenCL and, therefore, the ExBLAS routines. This drawback refers to the possibility of passing a reference to a location within a matrix ($\&A[i * (lda + 1)]$) as a kernel argument. We overcame this issue by passing an offset from the beginning of the matrix to the desired location. That introduces a change in the API of the BLAS to include a few new arguments.

Figs. \ref{fig:7a} and \ref{fig:7b} report the execution times obtained by the unblocked algorithmic variant for the LU factorization with partial pivoting as a function of the matrix size ($m = n$). The execution times of the \texttt{exlu} implementations are much larger than those of each ExBLAS routine for a specific problem size. That is because \texttt{exlu} requires $n - 1$ executions of each underlying ExBLAS routine for a matrix of size $n$. For instance, \texttt{extrsv} finds the solution of a triangular system using matrices of sizes from $1 \times 1$ till $(n - 1) \times (n - 1)$. In order to ensure reproducibility, the proposed algorithm based on \texttt{exlu} requires extra computations and memory. The outcome is visible in the form of 11× and 32× performance overheads on K420 and K80, respectively, compared to the double precision unblocked LU factorization; the latter overhead is the result of the large performance overhead caused by \texttt{extrsv}.

Furthermore, we study the accuracy of the non-deterministic and reproducible implementations of the unblocked LU factorization (Alg. \ref{alg:7}) on matrices with various condition numbers, covering a range from $10^2$ to $10^{41}$; these results are depicted in Fig. \ref{fig:7c}. To produce these matrices, we modify the ill-conditioned
triangular matrix generator to cover a more general case. We compute the error $\|PA - LU\|$ as an infinity norm using the MPFR library. For those errors that exceed 1, we round them down to 1. As for extrsv, ExLU delivers more accurate results than the double precision LU factorization. However, due to the division by the diagonal element in extrsv, the correct rounding of the results is not guaranteed in ExLU. We foresee to study and enhance this by improving the accuracy of the underlying extrsv using iterative refinement. Nevertheless, the ExLU implementations always deliver reproducible results.

6. Related Work

There already exist some solutions to ensure numerical producibility, which can be classified with respect to their implementations into sequential, vectorized, multithreaded on multicores, and distributed. Those
solutions, at first, addressed sequential and parallel summation. For example, \cite{33, 35} focus on hybrid solutions that store the sum as floating-point numbers of fixed exponent without completely avoiding the previous drawbacks. Arbitrary precision libraries – like MPFR \cite{11} – are able to provide correct rounding. However, they are not designed to achieve acceptable performance for reproducible results. Moreover, MPFR is also not multi-threaded. As a result, \texttt{exsum} is three orders of magnitude faster than MPFR.

To ensure reproducibility, Intel introduced a Conditional Numerical Reproducibility (CNR) option in its MKL library \cite{23}, but this does not provide any warranty on the accuracy. In addition, a CNR-enabled execution of MKL kernels incurs a large performance overhead compared to the existing alternatives. For instance, MKL’s \texttt{dasum()} with CNR enabled is two times slower than both the conventional MKL’s \texttt{dasum()} and \texttt{exsum}.

There are two other academic efforts to guarantee reproducibility. The first \cite{7}, which is built upon \cite{33}, proposed a family of algorithms for reproducible summation in FP arithmetic. Demmel and Nguyen improved their algorithms \cite{8} by passing over the input data only once. This approach reduces the overhead to roughly 20%. Arteaga et al. \cite{2} used this approach with improved communication and obtained the same accuracy with roughly 10% overhead. Demmel and Nguyen applied their approach to the absolute value summation, dot product, and 2-norm in the ReproBLAS library. Recently, they introduced \cite{1} new concepts – such as slice (significant bits in a bin) and indexed type/sum – and reformulated the algorithms \cite{8} in terms of these concepts. They introduced the conversion algorithm that leads to improved accuracy of their reproducible summation, handling of exceptions (not implemented in software), propagation of overflows. ReproBLAS was extended to include sequential matrix-vector and matrix-matrix products. However, ReproBLAS targets only CPU and does not exploit the data-parallelism available in BLAS routines.

The approach in \cite{5} ensures reproducibility of both sequential and parallel summations. Chohra et. al. proposed to use a combination of existing solutions such as FastAccSum \cite{33} and OnlineExact \cite{35} depending on the size of the input vector. They cover absolute value summation, 2-norm, dot product and matrix-vector product, provided as the RARE-BLAS library. RARE-BLAS, which is not publicly available, runs on Intel server CPUs and Intel Xeon Phi coprocessors. The reproducibility of 2-norm is an open question as the returned result is faithfully rounded.

Alternatively, Collange et al. \cite{6} proposed a multi-level approach to compute the reproducible summation. Neal \cite{30} integrated this concept in its scalar superaccumulators of different sizes for the summation problem in the R package. This approach is based on FP expansions and Kulisch superaccumulators, discussed in Section \ref{section:2}. We showed that the numerical reproducibility and bit-perfect (correct rounding) accuracy can be achieved without performance degradation for large sums with dynamic ranges of up to 90 orders of magnitude on a large variety of computer architectures, including conventional clusters, GPUs, and coprocessors. By tracking every bit of information for a product of two FP numbers, we extended this approach to the dot product, blocked triangular solver \cite{18}, and matrix-matrix multiplication \cite{20}. The initial release
7. Conclusions and Future Work

In numerical linear algebra, algorithms and their corresponding implementation exhibit a modular and hierarchical structure. This property permits to assemble higher-level algorithms on top of fundamental kernels. In this article, we leverage this layered organization of linear algebra algorithms to derive reproducible algorithmic variants for the LU factorization. We made the demonstration that a class of reproducible higher-level linear algebra operations can be constructed by following the bottom-up approach. Through our development, we noticed that in the context of reproducibility this process is not ‘automatic’ and should be handled carefully especially if accuracy is also a major concern.

As a case study, we considered the unblocked jik variant of the LU factorization that relies upon the Level-1/2 BLAS routines `dot`, `scal`, `gemv`, and `trsv`. In particular, we derived the accurate and reproducible matrix-vector product and provided performance results on NVIDIA GPUs. Additionally, we have improved the performance of `extrsv` and drew a strategy to enhance its accuracy via iterative refinement, potentially reaching correctly-rounded results.

Although some of the performance results can be argued in some scenarios, we advocate that we should not trade off numerical stability and reproducibility to performance. As both goal are conflicting, users should start considering that some of the performance benefit brought by today’s multicore has to be devoted to the later one.

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