

# Batched BLAS (Basic Linear Algebra Subprograms) 2018 Specification

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## Abstract

This document describes an API for Batch Basic Linear Algebra Subprograms (Batched BLAS or BBLAS). We focus on many independent BLAS operations on small matrices that are grouped together and processed by a single routine, called a Batched BLAS routine. The extensions beyond the original BLAS standard are considered that specify a programming interface not only for routines with uniformly-sized matrices and/or vectors but also for the situation where the sizes vary. The aim is to provide more efficient, but portable, implementations of algorithms on high-performance manycore platforms. These include multicore and many-core CPU processors; GPUs and coprocessors; as well as other hardware accelerators with floating-point compute facility.

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## 1 Introduction

### 1.1 The Batched BLAS

The specifications for the level 1, 2 and 3 BLAS have been very successful in providing a standard for vector, matrix-vector and matrix-matrix operations respectively, citeLawson1979l1blas, [1], citedongarra1990a. Vendors and other developers have provided highly efficient versions of the BLAS, and by using the standard interface have allowed software, calling the BLAS to be portable.

With the need to be able to solve larger and larger problems on today’s high-performance computers, the methods used in a number of applications such as tensor contractions, finite element methods and direct linear equation solvers, require a large number of small vector, or matrix operations to be performed in parallel. So a typical example might be to perform

$$C_i \leftarrow \alpha_i A_i B_i + \beta_i C_i, \quad i = 1, 2, \dots, \ell,$$

where  $k$  is large, but  $A_i, B_i$  and  $C_i$  are small matrices. A routine to perform such a sequence of operations is called a batched basic linear algebra subprogram, or Batched BLAS, or BBLAS.

### 1.2 History and Motivation

The origins of the Basic Linear Algebra Subprograms (BLAS) standard can be traced back to 1973, when Hanson, Krogh, and Lawson wrote an article in the SIGNUM Newsletter (Vol. 8, no. 4, p. 16) describing the advantages of adopting a set of basic routines for problems in linear algebra. This led to the development of the original BLAS [2], which indeed turned out to be advantageous and very successful. It was adopted as a standard and used in a wide range of numerical software, including LINPACK [3]. An extended, Level 2 BLAS, was proposed for matrix-vector operations [1, 4]. Unfortunately, while successful for the vector-processing machines at the time, Level 2 BLAS was not a good fit for the cache-based machines that emerged in the 1980’s. With these cache based machines, it was preferable to express computations as matrix-matrix operations. Matrices were split into small blocks so that basic operations were performed on blocks that could fit into cache memory. This approach avoids excessive

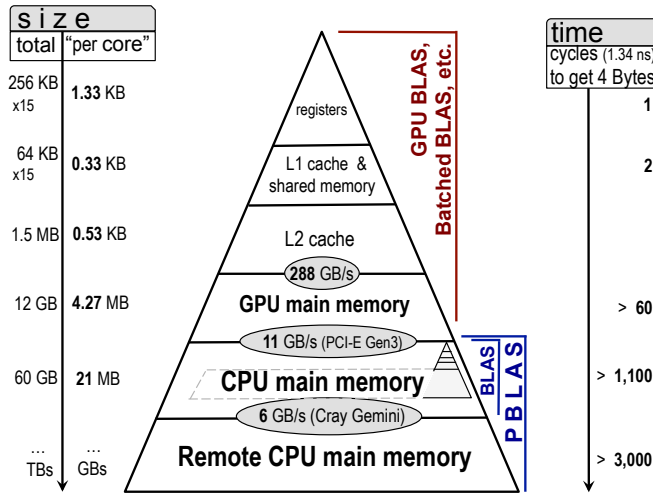


Figure 1: Memory hierarchy of a heterogeneous system from the point of view of a CUDA core of an NVIDIA K40c GPU with 2, 880 CUDA cores.

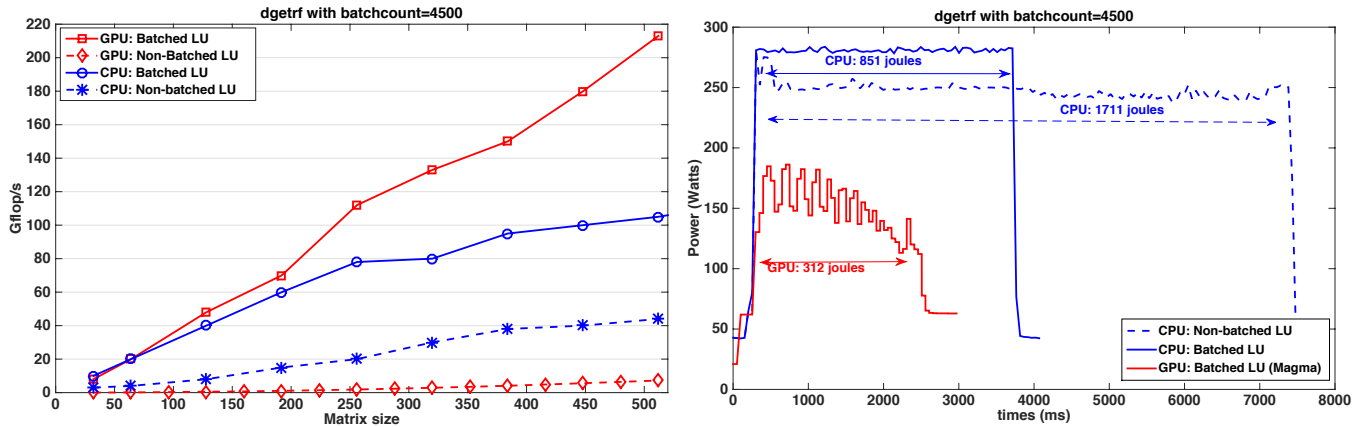


Figure 2: Speedup (Left) and power consumption (Right) achieved by the MAGMA batch LU factorization on NVIDIA K40c GPU vs. 16 cores of Intel Xeon ES-2670 (Sandy Bridge) 2.60GHz CPUs.

movement of data to and from memory and gives a surface-to-volume effect for the ratio of operations to data movement. Subsequently, Level 3 BLAS was proposed [5, 6], covering the main types of matrix-matrix operations, and LINPACK was redesigned into LAPACK [7] to use the new Level 3 BLAS where possible. For the emerging multicore architectures of the 2000's, the PLASMA library [8] introduced tiled algorithms and tiled data layouts. To handle parallelism, algorithms were split into tasks and data dependencies among the tasks were generated, and used by runtime systems to properly schedule the tasks' execution over the available cores, without violating any of the data dependencies. Overhead of scheduling becomes a challenge in this approach, since a single Level 3 BLAS routine on large matrices would be split into many Level 3 BLAS computations on small matrices, all of which must be analyzed, scheduled, and launched, without using information that these are actually independent data-parallel operations that share similar data dependencies.

In the 2010's, the apparently relentless trend in high performance computing (HPC) toward large-scale, heterogeneous systems with GPU accelerators and coprocessors made the near total absence of linear algebra software optimized for small matrix operations especially noticeable. The typical method of utilizing such hybrid systems is to increase the scale and resolution of the model used by an application, which in turn increases both matrix size and computational intensity; this tends to be a good match for the steady growth in performance and memory capacity of this type of hardware (see Figure 1 for an example of the memory hierarchy of this type of hardware). Unfortunately,

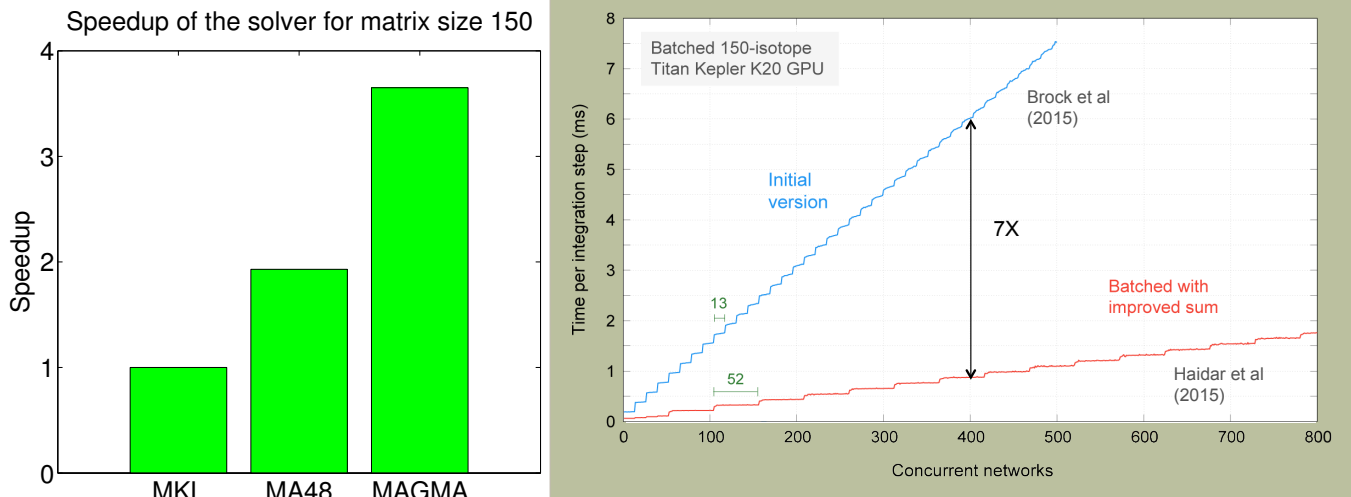


Figure 3: Acceleration of various applications by using batch approach.

numerous modern applications are cast in terms of a solution of many small matrix operations; that is, at some point in their execution, such programs must perform a computation that is cumulatively very large, but whose individual parts are very small; when such operations are implemented naively using the typical approach, they perform poorly. Applications that suffer from this problem include those that require tensor contractions (as in quantum Hall effect), astrophysics [9], metabolic networks [10], CFD and resulting PDEs through direct and multifrontal solvers [11], high-order FEM schemes for hydrodynamics [12], direct-iterative preconditioned solvers [13], quantum chemistry [14], image [15], and signal processing [16]. Batch LU factorization was used in subsurface transport simulation [17], whereby many chemical and microbiological reactions in a flow path are simulated in parallel [18]. Finally, small independent problems also occur as a very important aspect of computations on hierarchical matrices (H-matrices) [19].

One might expect that such applications would be well suited to accelerators or coprocessors, like GPUs. Due to the high levels of parallelism that these devices support, they can efficiently achieve very high performance for large data parallel computations when they are used in combination with a CPU that handles the part of the computation that is difficult to parallelize [20, 21, 22]. But for several reasons, this turns out not to be the case for applications that involve large amounts of data that come in small units. For the case of LU, QR, and Cholesky factorizations of many small matrices, we have demonstrated that, under such circumstances, by creating software that groups these small inputs together and runs them in large “batches,” we can dramatically improve performance by exploiting the increased parallelism that the grouping provides as well as the opportunities for algorithmic improvements and code optimizations [23, 24]. By using batch operations to overcome the bottleneck, small problems can be solved two to three times faster on GPUs, and with four to five times better energy efficiency than on multicore CPUs alone (subject to the same power draw). For example, Figure 2, Left illustrates this for the case of many small LU factorizations – even in a multicore setting the batch processing approach outperforms its non-batch counterpart by a factor of approximately two, while the batch approach in MAGMA<sup>1</sup> on a K40c GPU outperforms by about 2× the highly optimized CPU batch version running on 16 Intel Sandy Bridge cores [23]. Moreover, similarly to the way LAPACK routines benefit from BLAS, we have shown that these batch factorizations can be organized as a sequence of Batched BLAS calls, and their performance be portable across architectures, provided that the Batched BLAS needed are available and well optimized. Note that NVIDIA is already providing some optimized Batched BLAS implementations in CUBLAS [25], and Intel has also included a batch matrix-matrix product (GEMM BATCH) in MKL [26]. Subsequently, batch factorizations, and the underlying Batched BLAS, can be used in applications. For example, the batch LU results were used to speed up a nuclear network simulation – the XNet benchmark, as shown in Figure 3(a) – up to 3.6×, vs. using the MKL Library, and up to 2× speedup over the MA48 factorization from

<sup>1</sup>icl.utk.edu/magma

the Harwell Subroutine Library [27], by solving hundreds of matrices of size  $150 \times 150$  on the Titan supercomputer at ORNL [28]. Another example shown in Figure 3(b) is the astrophysical thermonuclear networks coupled to hydrodynamical simulations in explosive burning scenarios [29] that was accelerated  $7\times$  by using the batch approach.

Given the fundamental importance of numerical libraries to science and engineering applications of all types [30], the need for libraries that can perform batch operations on small matrices has clearly become acute. Therefore, to fill this critical gap, we propose standard interfaces for Batched BLAS operations.

The interfaces are intentionally designed to be close to the BLAS standard and to be *hardware independent*. They are given in C for use in C/C++ programs, but can readily be called from other languages and packages. The goal is to provide the developers of applications, compilers, and runtime systems with the option of expressing many small BLAS operations as a single call to a routine from the new batch operation standard, and thus to allow the entire linear algebra (LA) community to collectively attack a wide range of small matrix problems.

### 1.3 Community Involvement

A large number of people have contributed ideas to the Batched BLAS project. A number of the contributions in the form of papers and talks can be found at <http://icl.utk.edu/bblas/>. Two workshops were held in May 2016 and February 2017 [31, 32], Birds of a Feather sessions were held at SC17 in Denver and at ISC18 in Frankfurt, and a BBLAS minisymposium was held at SIAM PP18 in Tokyo, as well as a talk in an NLAFFET minisymposium.

## 2 Naming Conventions

### 2.1 Data Type and Functionality Conventions

The name of a Batched BLAS routine follows, and extends as needed, the conventions of the corresponding BLAS routine. In particular, the name is composed of 5 or 6 characters, specifying the BLAS routine and described below, followed by the suffix `_batch`:

- The first character in the name denotes the data type of the matrix (denoted as a type template `fp_t`), as follows:
  - S indicates **float**
  - D indicates **double**
  - C indicates **complex**
  - Z indicates **double complex**
  - H indicates **short float** (if available)<sup>2</sup>
  - Q indicates **long double** (if available)
- Characters two and three in the name refer to the kind of matrix involved, as follows:
  - GE All matrices are general rectangular
  - HE One of the matrices is Hermitian
  - SY One of the matrices is symmetric
  - TR One of the matrices is triangular
- The fourth and fifth, and in one case sixth, characters in the name denote the operation. For example, for the Level 3 Batched BLAS, the operations are given as follows:
  - MM represents: Matrix-matrix product
  - RK represents: Rank-k update of a symmetric or Hermitian matrix
  - R2K represents: Rank-2k update of a symmetric or Hermitian matrix

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<sup>2</sup>Half precision is available in Fortran and an extension to C/C++ language is being considered. IEEE 754 2018 floating-point standard has it available as a compute (rather than just storage) precision.

- SM represents: Solve a system of linear equations for a matrix of right-hand sides

The Level 1 and Level 2 Batched BLAS operations follow the corresponding Level 1 and Level 2 BLAS operations.

## 2.2 Argument Conventions

We follow a convention for the list of arguments that is similar to that for BLAS, with the necessary adaptations concerning the batch operations. The order of arguments is as follows:

1. Integer that specifies the number of matrices in the batch
2. Integer array that specifies batch sizes
3. Argument specifying row-, or column-major layout
4. Array of arguments specifying options
5. Array of arguments defining the sizes of the matrices
6. Array of descriptions of the input-output matrices
7. Array of input scalars (associated with input-output matrices)
8. Array of input scalars
9. Array of info parameters

Note that not every category is present in each of the routines.

### 2.2.1 Arguments Specifying Options

The arguments that specify options are of enumeration type with names such as `side`, `transa`, `transb`, `trans`, `uplo`, and `diag`. These arguments, along with the values that they can take, are described below:

- `layout` has two possible values which are used by the routines as follows:
  - `BlasColMajor`: specifies column-major layout of matrix elements;
  - `BlasRowMajor`: specifies row-major layout of matrix elements.
- `side` has two possible values which are used by the routines as follows:
  - `BlasLeft`: Specifies to multiply a general matrix by symmetric, Hermitian, or triangular matrix on the left;
  - `BlasRight`: Specifies to multiply general matrix by symmetric, Hermitian, or triangular matrix on the right.
- `trans_A`, `trans_B`, and `trans` can have three possible values each, which is used to specify the following:
  - `BlasNoTrans`: Operate with the matrix as it is;
  - `BlasTrans`: Operate with the transpose of the matrix;
  - `BlasConjTrans`: Operate with the conjugate transpose of the matrix.

Note that in the real case, the values `BlasTrans` and `BlasConjTrans` have the same meaning.

- `uplo` is used by the Hermitian, symmetric, and triangular matrix routines to specify whether the upper or lower triangle is being referenced, as follows:
  - `BlasLower`: Lower triangle

- `BlasUpper`: Upper triangle.
- `diag` is used by the triangular matrix routines to specify whether the matrix is unit triangular, as follows:
  - `BlasUnit`: Unit triangular;
  - `BlasNonUnit`: Nonunit triangular.

When `diag` is supplied as `BlasUnit`, the diagonal elements are not referenced.

### 2.2.2 Arguments defining the sizes

The sizes of matrices  $A_i$ ,  $B_i$ , and  $C_i$  for the  $i^{\text{th}}$  BLAS operation are determined by the corresponding values of the arrays  $m$ ,  $n$ , and  $k$  at position  $i$  (see the routine interfaces in Section 4). It is permissible to call the routines with  $m = 0$  or  $n = 0$ , in which case the routines do not reference their corresponding matrix arguments and do not perform any computation on the corresponding matrices  $A_i$ ,  $B_i$ , and  $C_i$ . If  $m > 0$  and  $n > 0$ , but  $k = 0$ , the Level 3 BLAS operation reduces to  $C = \beta C$  (this applies to the `gemm`, `syrk`, `herk`, `syr2k`, and `her2k` routines). The input-output matrix ( $B$  for the `tr` routines,  $C$  otherwise) is always  $m$  by  $n$  if working with rectangular  $A$ , and  $n$  by  $n$  if  $A$  is a square matrix. If there is only a single group of matrices of the same sizes (see Section 2.3.2), the  $m$ ,  $n$ , and  $k$  values for all matrices are specified by the `m[0]`, `n[0]`, and `k[0]` values, respectively.

### 2.2.3 Arguments describing the input-output matrices

The description of the matrix consists of the array name (`A`, `B`, or `C`) followed by an array of the leading dimension as declared in the calling function (`ld_A`, `ld_B`, or `ld_C`). The  $i^{\text{th}}$  values of the `A`, `B`, and `C` are pointers to the arrays of data  $A_i$ ,  $B_i$ , and  $C_i$ , respectively. Similarly, the values of `ld_A[i]`, `ld_B[i]`, and `ld_C[i]` correspond to the leading dimensions of the matrices  $A_i$ ,  $B_i$ , and  $C_i$ , respectively. For batch style with the same leading dimensions (see Section 2.3.2), the leading dimensions are specified by `ld_A[0]`, `ld_B[0]`, and `ld_C[0]` for all corresponding  $\{A_i\}$ ,  $\{B_i\}$ , and  $\{C_i\}$  matrices.

### 2.2.4 Arguments defining the input scalar

Arrays of scalars are named `alpha` and `beta`, where values at position  $i$  correspond to the  $\alpha$  and  $\beta$  scalars for the BLAS operation involving matrices  $A_i$ ,  $B_i$ , and  $C_i$ . For batch style with the same scalars (see Section 2.3.2), the scalars are given by `alpha[0]` and `beta[0]`.

## 2.3 Groups of Same-Size Batched BLAS Routines

During the past standardization meetings [31, 32] a consensus emerged to amend the previous draft of the Batched BLAS standard [33] to include in the proposed interface the situation where the sizes of matrices in the batch vary by group. The following formula calculates the argument formerly called `batch_count` (the total number of matrices in a single call) from the number and size of individual groups of matrices:

$$\text{batch\_count} = \sum_{i=0}^{\text{group\_count}-1} \text{group\_sizes}[i] \quad (1)$$

### 2.3.1 Specification of the number of matrices

The total number of matrices involved in a single call may be derived from two arguments: `group_count` and `group_sizes`. Formerly, this was known as the `batch_count` argument [33] – an integer that indicated the number of matrices to be processed. If there is more than one group of matrices, Eq. (1) may be used for calculating `batch_count`.

### 2.3.2 Batch Style Specification

The `batch_opts` argument from the previous proposal [33] was an enumerated value that specified the style for the batch computation. Permitted values were either `BLAS_BATCH_FIXED` or `BLAS_BATCH_VARIABLE`, which stood for computation of matrices with the same or group-varying sizes (including operation options, sizes, matrix leading dimensions, and scalars), respectively. This was superseded by the group interface.

**Note** that through the group interface one can specify constant size or variable size Batched BLAS operations. If a constant size batch is requested, the arguments point to the corresponding constant value. The goal of this interface is to remove the need for users to prepare and pass arrays whenever they have the same elements. Through an internal dispatch and based on the group sizes, an expert routine specific to the value/style can be called while keeping the top interface the same.

### 2.4 Error handling defined by the INFO array

For the Batched BLAS the argument `info` is an input/output argument.

On input, the value of `info` should have one of the following values:

- `BBLAS_ERRORS_REPORT_ALL`, which indicates that all errors will be specified on output. The length of the `info` array should be greater than or equal to the batch count.
- `BBLAS_ERRORS_REPORT_GROUP`, which indicates that only a single error will be reported for each group, independently. The length of the `info` array should be greater than or equal to the group count.
- `BBLAS_ERRORS_REPORT_ANY`, which indicates that the occurrence of errors will be specified on output as a single integer value. The length of the `info` array should be at least one.
- `BBLAS_ERRORS_REPORT_NONE`, which indicates that no errors will be reported on output. The length of the `info` array should be at least one.

The following values of arguments are invalid:

- Any value of the character arguments `side`, `trans_A`, `trans_B`, `trans`, `uplo`, or `diag` whose meaning is not specified;
- If any of `m`, `n`, `k`, `ld_A`, `ld_B`, or `ld_C` is less than zero.

The behaviour of the error handling is, of course, determined by the input value of `info`, see Section 3.2 for further details, but with full reporting, if a routine is called with an invalid value for arguments: `group_count` and `group_sizes` then the routine will return an error in `info[0]`. Errors related to other arguments are signaled with the number of the group in which the invalid argument was encountered (counting from one because the value of 0 is reserved for the return without an error). In other words, if a routine is called with an invalid value for any of its other arguments for a Batched BLAS operation in group  $g$  for matrix  $i$ , the routine will return an error in position `info[1+p+i]` that refers to the number of the first invalid argument (counting from one with number 0 reserved for successful completion) where  $p$  is the number of matrices in groups 1 through  $g - 1$ .

## 3 Error Handling

### 3.1 Legacy Error Reporting Methods

Historically—with the exception of Level 1 BLAS, which had no error reporting—BLAS used a method outside of the call stack to notify the caller about potential problems occurring during the execution of any routine. This design decision was made in the 1980s, and hardware architectures and software practices have changed significantly in recent decades. Nevertheless, we give a more detailed account of how this design causes problems in modern software development.

There are a few advantages of the BLAS error-handling method. First, the default implementation of `XERBLA()` guarantees that errors are reported regardless of whether the caller code checks for errors. If there is a reason for ignoring errors, then the user can simply override the default implementation, and the errors no longer need to be



reported. A similar technique, in the form of the `ILAENV()` routine, could be used to take over the tuning mechanism of LAPACK.

Another advantage is user familiarity because this mechanism corresponds to the ways UNIX® reports problems with a combination of `kill()` and `signal()` calls.

Additionally, unlike in C programs that often return an integer error code, this is not an accepted practice in Fortran. For FORTRAN 77 libraries, using a subroutine instead of a function minimizes the spelling of a routine declaration to just one line, as shown below.

```
1! just one line required for declaring a subroutine
2  EXTERNAL DGEMM
3! two lines required for declaring a function
4  INTEGER DGEMMF
5  EXTERNAL DGEMMF
```

On the other hand, adding an additional output parameter (as was done for LAPACK) clobbers the API and adds boiler plate code at the calling site and at the implementation site when the user has no intention of using the error code parameter.

The development workflow that uses `XERBLA()` for simple codes would be comprised of the following generic steps:

1. A code is written and a bug causes it to pass incorrect parameters to BLAS.
2. While executing the offending code, the reported errors are recorded.
3. Corrections of the reported errors are made by noting the routine name and the source of the problem.

### 3.1.1 Specific Issues with `XERBLA()` in BLAS

Unfortunately, the default `XERBLA()` implementation is not sufficiently precise for complex runtime error scenarios. If a BLAS routine is called in a loop, then the input/output (I/O) buffer or the console screen will be flooded with error messages. This would require one to, for example, suppress error messages in a custom `XERBLA()` implementation.<sup>3</sup>

Another problem is the fact that the same routine may be called from two distinct locations in a user's code. The default implementation of `XERBLA()` cannot account for this nor differentiate between the two. A custom `XERBLA()` implementation could communicate with the calling code through global variables to indicate the exact location of the call (e.g., the source code file name and the line number), but this requires modification of the calling routine, which is what the `XERBLA()` method is trying to avoid.

In summary, the main issue with using `XERBLA()` as an error reporting method is that it is a non-local approach that decouples the call site from the error site and requires out-of-band messaging (e.g., global variables) to sufficiently contextualize the source and reason for the invalid behavior.

Below is a specific list of issues associated with the `XERBLA()` mechanism.

**Use of global state.** `XERBLA()` requires global variables for non-trivial customization and information passing between the user and the BLAS library.

**Dependence on platform-specific features.** Often, dynamic libraries require special features in the binary format of the operating system (OS) to overload a function. This is not hardware specific but also involves the accompanying software stack, including the OS and the compiler-linker tool chain.

**Limited customization.** There can only be one `XERBLA()` per executable, and there is no mechanism for chaining or queuing its invocations in case two different call sites would like to install different error handlers. Furthermore, there is no way to establish a protocol between call sites for cooperative error handling because the only feature available is the linker name replacement system, which is available in Linux and Mac OS X and used when creating ELF or Mach-O object files.

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<sup>3</sup>Only some development workflows support this mode of operation.

**Language-specific behavior dependent on name mangling.** Modern BLAS standards and their implementations expose the Fortran API and the C API. The older CBLAS standard implements functions like `cblas_dgemm()` and the newer standard uses `BLAS_dgemm()`. The `XERBLA()` mechanism requires resolving the coexistence of both language bindings (Fortran and C), sometimes in the same binary. Neither of these languages necessarily share the same I/O streams, and—in a mixed programming language environment—it is not obvious which `XERBLA()` binding needs to be reimplemented to take over the BLAS error handling.

**Mixing computational capabilities with I/O facilities.** According to the standard’s definition of `XERBLA()`, the use of I/O streams is required by the default implementation inside the BLAS library. This obviously causes issues for headless mode operation when the access to the I/O facilities is restricted to accommodate custom environments. For example, on an embedded system or on a cloud platform, the only available I/O stream might occur during limited system logging or extra overheads generated by system-wide synchronization.

**Lack of support for the asynchronous interface.** The `XERBLA()` error handling mechanism is not meant for the asynchronous and event-based processing that has become prevalent on modern HPC architectures. Modern computing hardware features multiple execution streams that add flexibility to scheduling at the hardware level but do not guarantee a specific order of completion for independent subroutine calls. This means that the `XERBLA()`-based library cannot be wrapped inside such an interface because error delivery is independent of the error-causing invocation. Connecting the two would also add unnecessary complexity and synchronization and thus diminish the potential benefits of asynchronous execution.

**Lack of support for multithreading.** The BLAS interface with `XERBLA()` is inherently single threaded. Multiple threads that asynchronously call BLAS and cause invocation of `XERBLA()` must be synchronized to provide coherent error reporting. The behavior under such circumstances is unspecified, and extra care has to be devoted to recognize the calling thread, e.g., with calls to `pthread_self()` or `omp_get_num_threads()` and contextualize the error reporting accordingly.

### 3.1.2 Specific Issues with `XERBLA()` in BATCHED BLAS

Compared to the classic single-operation BLAS, BATCHED BLAS presents additional concerns for error handling through `XERBLA()`. For example, the batched operations may develop errors for all matrices, some matrices, or just one matrix. Also, for the group-based interface, the error can be per matrix, per group, or per the entire batch. All of these scenarios make error tracking through `XERBLA()` even more complicated, which leads to much higher overhead when an error does occur. For these reasons, `XERBLA()` is not the appropriate error-handling mechanism for BATCHED BLAS.

## 3.2 Design Goals for an Error Reporting Mechanism

It is worth mentioning that the `XERBLA()` mechanism, for all its shortcomings, does address a range of important error handling scenarios, described below.

**All errors reported.** This mode corresponds to a development stage where the user is uncertain about the correctness of the BLAS invocations and would like an immediate notification of errors before they propagate through the code base. Also, in this mode the user may discover any mismatch between the behavior that is expected and the behavior that is observed. This may occur if the user misunderstands the BATCHED BLAS standard, if the implementation is non compliant, or if the error checking is incorrect.

**Some errors reported.** In this mode, the code is composed of sections with correct invocations and of sections with potentially erroneous calls. The former corresponds to a production-hardened code that can be trusted because of its prior verification and compliance with a robust testing profile. The latter, on the other hand, is development code that requires error notifications to isolate its effect on the rest of the code.

**No errors reported.** This is the production run scenario when error reporting is unnecessary and performance is essential.

Any potential BATCHED BLAS error handling mechanism must address all three of these scenarios as they cover the large majority of software engineering and performance-optimization practices in HPC and scientific computing. This can be accomplished by adding an I/O parameter, `info`, for an integer array type to the BATCHED BLAS calls. On input, the value of `info` will have one of the following values:

- `BBLAS_ERRORS_REPORT_ALL`, which indicates that all errors will be specified on output. The length of the `info` array should be greater than or equal to the batch count.
- `BBLAS_ERRORS_REPORT_GROUP`, which indicates that only a single error will be reported for each group, independently. The length of the `info` array should be greater than or equal to the group count.
- `BBLAS_ERRORS_REPORT_ANY`, which indicates that the occurrence of errors will be specified on output as a single integer value. The length of the `info` array should be at least one.
- `BBLAS_ERRORS_REPORT_NONE`, which indicates that no errors will be reported on output. The length of the `info` array should be at least one.

On output, when the input for `info` is set to `BBLAS_ERRORS_REPORT_ALL`, the value of `info` is modified to indicate an error for each individual problem in the batch. Only a single error will be reported when `info` is set to `BBLAS_ERRORS_REPORT_ANY` on input.

Unlike the uncaught signals, BATCHED BLAS routines will not exit the program (e.g., like when executing the `exit` system call) and will always return an error code in the `info` parameter.

In Level 2 BLAS, the use of `XERBLA()` was always limited to interface errors, where it mostly handled invalid or inconsistent parameter values. However, in some BLAS routines, it is possible to create numerical conditions that could be considered errors. As an example, consider detecting a zero value on the diagonal in `TRSM()`: it is not considered an error in BLAS, and `XERBLA()` is not called in that situation.<sup>4</sup> Depending on the floating-point hardware and system settings, this may generate positive or negative infinities—and NaNs, subsequently—throughout the resulting matrix. Alternatively, a floating point exception could be raised, and an appropriate handler would be invoked to deal with the problem. As a result, the implementation of the routine does not require the extra code for handling such numerical issues—an omission that often leads to faster and more compact code, which is an important consideration for a performance-portable library like BLAS.

The downside to offloading this task from the routine is that the details of handling such situations become hardware and OS-specific. For this reason, LAPACK includes routines called `STRTRS`, `DTRTRS`, `CTRTRS`, and `ZTRTRS` that perform the triangular solves and handle zeros on the diagonal, explicitly, as needed by the calling routine. The intention of BATCHED BLAS is to follow the same policy and not report numerical issues, including checks for special values (i.e., `Inf` and `NaN`) in scalars, vectors, or matrices, regardless of whether those values originate in user data or are produced during BATCHED BLAS calculations.

A sample code that ignores errors in C might look like this:

```
1 int info[1] = {BBLAS_ERRORS_IGNORE};
2
3 BBLAS_dtrsm(..., info);
4 BBLAS_dgemm(..., info);
```

## 4 Specification of Batched BLAS Routines

### 4.1 Scope And Specifications of the Level 3 Batched BLAS

The Level 3 Batched BLAS routines described here have been derived in a fairly obvious manner from the interfaces of their corresponding Level 3 BLAS routines. The advantage in keeping the design of the software as consistent as possible with that of the BLAS is that it will be easier for users to replace their BLAS calls by calling the Batched BLAS when needed, and to remember the calling sequences and the parameter conventions. In real arithmetic, the operations proposed for the Level 3 Batched BLAS have an interface described as follows.

---

<sup>4</sup>Note that BLAS and LAPACK were written before the IEEE 754 standard and before a consistent meaning of all numerical exceptions and results was established.

### 4.1.1 General matrix-matrix products GEMM

This routine performs a batch of one of the matrix-matrix operations described below:

- $C \leftarrow \alpha \cdot A \times B + \beta C, A \in \mathbb{R}^{m \times k}, B \in \mathbb{R}^{k \times n}, C \in \mathbb{R}^{m \times n}$
- $C \leftarrow \alpha \cdot A^T \times B + \beta C, A \in \mathbb{R}^{k \times m}, B \in \mathbb{R}^{k \times n}, C \in \mathbb{R}^{m \times n}$
- $C \leftarrow \alpha \cdot A^H \times B + \beta C, A \in \mathbb{C}^{k \times m}, B \in \mathbb{C}^{k \times n}, C \in \mathbb{C}^{m \times n}$
- $C \leftarrow \alpha \cdot A \times B^T + \beta C, A \in \mathbb{R}^{m \times k}, B \in \mathbb{R}^{n \times k}, C \in \mathbb{R}^{m \times n}$
- $C \leftarrow \alpha \cdot A \times B^H + \beta C, A \in \mathbb{C}^{m \times k}, B \in \mathbb{C}^{n \times k}, C \in \mathbb{C}^{m \times n}$
- $C \leftarrow \alpha \cdot A^T \times B^T + \beta C, A \in \mathbb{R}^{k \times m}, B \in \mathbb{R}^{n \times k}, C \in \mathbb{R}^{m \times n}$
- $C \leftarrow \alpha \cdot A^H \times B^H + \beta C, A \in \mathbb{C}^{k \times m}, B \in \mathbb{C}^{n \times k}, C \in \mathbb{C}^{m \times n}$

The calling routine is described as follows:

```
BLAS_gemm_batch<fp_t> (
  group_count : int           : In ,
  group_sizes : int [group_count] : In ,
  layout      : enum Layout   : In ,
  trans_A     : enum Transpose [count] : In ,
  trans_B     : enum Transpose [count] : In ,
  m           : int [count]   : In ,
  n           : int [count]   : In ,
  k           : int [count]   : In ,
  alpha       : fp_t [count]  : In ,
  A           : fp_t [count]  : In ,
  ld_A        : int [count]   : In ,
  B           : fp_t [count]  : In ,
  ld_B        : int [count]   : In ,
  beta        : fp_t [count]  : In ,
  C           : fp_t [count]  : InOut ,
  ld_C        : int [count]   : In ,
  info        : int [count]   : InOut
)
```

where `fp_t` denotes one of the four standard floating-point arithmetic precisions (`float`, `double`, `complex`, or `double complex`). The `trans_A` and `trans_B` arrays can be of size one for the same size batch and of size at least `batch_count` for the variable sizes case. For the latter, each value defines the operation on the corresponding matrix. In the real precision case, the values `BlasTrans` and `BlasConjTrans` have the same meaning. The `m`, `n`, and `k` arrays of integers are of size at least `batch_count`, where each value defines the dimension of the operation on each corresponding matrix. The `alpha` and `beta` arrays provide the scalars  $\alpha$  and  $\beta$ , described in the equation above. They are of the same precision as the arrays `A`, `B`, and `C`. The arrays of pointers `A`, `B`, and `C` are of size at least `batch_count` and point to the matrices  $\{A_i\}$ ,  $\{B_i\}$ , and  $\{C_i\}$ . The size of matrix  $C_i$  is `m[i] * n[i]`. The sizes of the matrices  $A_i$  and  $B_i$  depend on `trans_A[i]` and `trans_B[i]`; their corresponding sizes are mentioned in the equation above. The arrays `ld_A`, `ld_B`, and `ld_C` define the leading dimension of each of the matrices  $\{A_i[\text{ld}_A[i]][*]\}$ ,  $\{B_i[\text{ld}_B[i]][*]\}$ ,  $\{C_i[\text{ld}_C[i]][*]\}$ , respectively.

If there is only one group of matrices (`group_count == 1`) only `transa[0]`, `transb[0]`, `m[0]`, `n[0]`, `k[0]`, `alpha[0]`, `lda[0]`, `ldb[0]`, `beta[0]`, and `ldc[0]` are used to specify the `gemm` parameters for the batch.

The array `info` defines the error array. It is an output array of integers of size `batch_count` where a value at position  $i$  reflects the argument error for the `gemm` with matrices  $A_i$ ,  $B_i$ , and  $C_i$ .

#### 4.1.2 Hermitian and symmetric matrix-matrix products: HEMM and SYMM

This routine performs a batch of matrix-matrix products, each expressed in one of the following forms:

- $C \leftarrow \alpha \cdot A \times B + \beta C$  if `side==BlasLeft`;  
 $A \in \mathbb{R}^{m \times m}; B, C \in \mathbb{R}^{m \times n}$
- $C \leftarrow \alpha \cdot B \times A + \beta C$  if `side==BlasRight`;  
 $A \in \mathbb{R}^{m \times m}; B, C \in \mathbb{R}^{m \times n}$

where the matrices  $A$ ,  $B$ , and  $C$  are real symmetric (`symm_batch`), complex symmetric (`symm_batch`), or complex Hermitian (`hemm_batch`), and  $\alpha$  and  $\beta$  are real or complex scalars.

The calling routine is described as follows:

```
BLAS_symm_batch-<fp_t> (
  group_count : int           : In ,
  group_sizes : int [group_count] : In ,
  layout      : enum Layout   : In ,
  side       : enum Side [count] : In ,
  uplo      : enum UpLo [count] : In ,
  m         : int [count]     : In ,
  n         : int [count]     : In ,
  alpha     : fp_t [count]    : In ,
  A         : fp_t [count]    : In ,
  ld_A     : int [count]     : In ,
  B         : fp_t [count]    : In ,
  ld_B     : int [count]     : In ,
  beta     : fp_t [count]    : In ,
  C         : fp_t [count]    : InOut ,
  ld_C     : int [count]     : In ,
  info     : int [count]     : InOut
)
```

```
BLAS_hemm_batch-<fp_t> (
  group_count : int           : In ,
  group_sizes : int [group_count] : In ,
  layout      : enum Layout   : In ,
  side       : enum Side [count] : In ,
  uplo      : enum UpLo [count] : In ,
  m         : int [count]     : In ,
  n         : int [count]     : In ,
  alpha     : fp_t [count]    : In ,
  A         : fp_t [count]    : In ,
  ld_A     : int [count]     : In ,
  B         : fp_t [count]    : In ,
  ld_B     : int [count]     : In ,
  beta     : fp_t [count]    : In ,
  C         : fp_t [count]    : InOut ,
  ld_C     : int [count]     : In ,
  info     : int [count]     : InOut
)
```

The `side` array is of size at least `batch_count` and each value defines the operation on each matrix as described in the equations above. The `uplo` array is of size at least `batch_count` and defines whether the upper or the lower triangular part of the matrix is to be referenced. The `m` and `n` arrays of integers are of size at least

`batch_count` and define the dimension of the operation on each matrix. The `alpha` and `beta` arrays provide the scalars  $\alpha_i$  and  $\beta_i$  described in the equation above. They are of the same precision as the arrays `A`, `B`, and `C`. The arrays `A`, `B`, and `C` are the arrays of pointers of size `batch_count` that point to the matrices  $\{A_i\}$ ,  $\{B_i\}$ , and  $\{C_i\}$ . The size of matrix  $C_i$  is `m[i]` by `n[i]`. The sizes of the matrices  $A_i$  and  $B_i$  depend on `side[i]`; their corresponding sizes are mentioned in the equations above. The arrays `lda`, `ldb`, and `ldc` define the leading dimension of each of the matrices  $\{A_i[\text{ld}_A[i]][*]\}$ ,  $\{B_i[\text{ld}_B[i]][*]\}$ ,  $\{C_i[\text{ld}_C[i]][*]\}$ , respectively.

The array `info` defines the error array. It is an output array of integers of size `batch_count` where a value at position  $i$  reflects the argument error for the hemm/symm with matrices  $A_i$ ,  $B_i$ , and  $C_i$ .

#### 4.1.3 Rank-k updates of a symmetric/Hermitian matrix HERK and SYRK

This routine performs a batch of rank-k updates of real symmetric (`syrc_batch`), complex symmetric (`syrc_batch`), or complex Hermitian (`herk_batch`) matrices in the form:

- $C \leftarrow \alpha \cdot A \times A^T + \beta \cdot C$  for `syrc` if `trans==BlasNoTrans`;  
 $A \in \mathbb{R}^{n \times k}; C \in \mathbb{R}^{n \times n}$
- $C \leftarrow \alpha \cdot A^T \times A + \beta \cdot C$  for `syrc` if `side==BlasTrans`;  
 $A \in \mathbb{R}^{k \times n}; C \in \mathbb{R}^{n \times n}$
- $C \leftarrow \alpha \cdot A \times A^H + \beta \cdot C$  for `herk` if `trans==BlasNoTrans`;  
 $A \in \mathbb{C}^{n \times k}; C \in \mathbb{C}^{n \times n}$
- $C \leftarrow \alpha \cdot A^H \times A + \beta \cdot C$  for `herk` if `side==BlasTrans`;  
 $A \in \mathbb{C}^{k \times n}; C \in \mathbb{C}^{n \times n}$

The calling routine is described as follows:

```
BLAS_syrc_batch_<fp_t> (
  group_count : int           : In ,
  group_sizes : int [group_count] : In ,
  layout      : enum Layout   : In ,
  uplo       : enum UpLo [count] : In ,
  trans      : enum Transpose [count] : In ,
  n          : int [count]     : In ,
  k          : int [count]     : In ,
  alpha     : fp_t [count]    : In ,
  A         : fp_t [count]    : In ,
  ld_A     : int [count]      : In ,
  beta     : fp_t [count]    : In ,
  C        : fp_t [count]    : InOut ,
  ld_C     : int [count]      : In ,
  info     : int [count]      : InOut
)
```

The `uplo` array is of size at least `batch_count` and defines whether the upper or the lower triangular part of the matrix is to be referenced. The `trans` array is of size at least `batch_count` where each value defines the operation on each matrix. In the real precision case, the values `BlasTrans` and `BlasConjTrans` have the same meaning. In the complex case, `trans == BlasConjTrans` is not allowed in `syrc` case. The `n` and `k` arrays of integers are of size at least `batch_count` and define the dimensions of the operation on each matrix. The `alpha` and `beta` arrays provide the scalars  $\alpha$  and  $\beta$  described in the equation above. They are of the same precision as the arrays  $A_i$  and  $C_i$ . The arrays of pointers `A` and `C` are of size `batch_count` and point to the matrices  $\{A_i\}$  and  $\{C_i\}$ . The size of matrix  $C_i$  is `n[i]` by `n[i]`. All matrices  $\{C_i\}$  are either real or complex symmetric. The size of the matrix  $A_i$  depends on `trans[i]`; its corresponding size is mentioned in the equation above. The arrays `ld_A` and `ld_C` define the leading dimension of each of the matrices  $\{A_i[\text{ld}_A[i]][*]\}$ ,  $\{C_i[\text{ld}_C[i]][*]\}$ , respectively.

The array `info` defines the error array. It is an output array of integers of size `batch_count` where a value at position  $i$  reflects the argument error for the `syrk` with matrices  $A_i$  and  $C_i$ .

```
BLAS_herk_batch_<fp_t> (
  group_count : int           : In ,
  group_sizes : int [group_count] : In ,
  layout      : enum Layout    : In ,
  uplo        : enum UpLo [count] : In ,
  trans       : enum Transpose [count] : In ,
  n           : int [count]     : In ,
  k           : int [count]     : In ,
  alpha       : fp_t [count]    : In ,
  A           : fp_t [count]    : In ,
  ld_A        : int [count]     : In ,
  beta        : fp_t [count]    : In ,
  C           : fp_t [count]    : InOut ,
  ld_C        : int [count]     : In ,
  info        : int [count]     : InOut
)
```

This routine is only available for the complex precisions. It has the same parameters as the `syrk` batch except that the `trans == BlasTrans` is not allowed in `herk` batch and that `alpha` and `beta` are real. The matrices  $\{C_i\}$  are complex Hermitian.

The array `info` defines the error array. It is an output array of integers of size `batch_count` where a value at position  $i$  reflects the argument error for the `herk` with matrices  $A_i$  and  $C_i$ .

#### 4.1.4 Rank-2k updates of a symmetric/Hermitian matrix HER2K and SYR2K

This routine performs batch rank-2k updates on real symmetric (SYR2K), complex symmetric (SYR2K), or complex Hermitian (HER2K) matrices of the form:

- $C \leftarrow \alpha \cdot A \times B^T + \alpha \cdot B \times A^T + \beta \cdot C$  for `syr2k` if `trans == BlasNoTrans`;  $A, B \in \mathbb{R}^{n \times k}$ ;  $C \in \mathbb{R}^{n \times n}$
- $C \leftarrow \alpha \cdot A^T \times B + \alpha \cdot B^T \times A + \beta \cdot C$  for `syr2k` if `trans == BlasTrans`;  $A, B \in \mathbb{R}^{k \times n}$ ;  $C \in \mathbb{R}^{n \times n}$
- $C \leftarrow \alpha \cdot A \times B^H + \alpha \cdot B \times A^H + \beta \cdot C$  for `her2k` if `trans == BlasNoTrans`;  $A, B \in \mathbb{C}^{n \times k}$ ;  $C \in \mathbb{C}^{n \times n}$
- $C \leftarrow \alpha \cdot A^H \times B + \alpha \cdot B^H \times A + \beta \cdot C$  for `her2k` if `trans == BlasConjTrans`;  $A, B \in \mathbb{C}^{k \times n}$ ;  $C \in \mathbb{C}^{n \times n}$

The calling routine is described as follows:

```
BLAS_syr2k_batch_<fp_t> (
  group_count : int           : In ,
  group_sizes : int [group_count] : In ,
  layout      : enum Layout    : In ,
  uplo        : enum UpLo [count] : In ,
  trans       : enum Transpose [count] : In ,
  n           : int [count]     : In ,
  k           : int [count]     : In ,
  alpha       : fp_t [count]    : In ,
  A           : fp_t [count]    : In ,
  ld_A        : int [count]     : In ,
  beta        : fp_t [count]    : In ,
  C           : fp_t [count]    : InOut ,
  ld_C        : int [count]     : In ,
  info        : int [count]     : InOut
)
```

The `uplo` array is of size `batch_count` and defines whether the upper or the lower triangular part of the matrix is to be referenced. The `trans` array is of size `batch_count` where each value defines the operation on each matrix. In the real precision case, the values `BlasTrans` and `BlasConjTrans` have the same meaning. In the complex case, `trans == BlasConjTrans` is not allowed in `syr2k` batch. The `n` and `k` arrays of integers are of size `batch_count` and define the dimensions of the operation on each matrix. The `alpha` and `beta` arrays provide the scalars  $\alpha$  and  $\beta$  described in the equations above. They are of the same precision as the arrays `A`, `B`, and `C`. The arrays `A`, `B`, and `C` are the arrays of pointers of size `batch_count` that point to the matrices  $\{A_i\}$ ,  $\{B_i\}$ , and  $\{C_i\}$ . The size of matrix  $C_i$  is `n[i]` by `n[i]`. All matrices  $\{C_i\}$  are either real or complex symmetric. The size of the matrices  $A_i$  and  $B_i$  depends on `trans[i]`; its corresponding size is mentioned in the equation above. The arrays `ld_A`, `ld_B`, and `ld_C` define the leading dimension of the matrices  $\{A_i[\text{ld\_A}[i]][*]\}$ ,  $\{B_i[\text{ld\_B}[i]][*]\}$ ,  $\{C_i[\text{ld\_C}[i]][*]\}$ , respectively.

The array `info` defines the error array. It is an output array of integers of size `batch_count` where a value at position  $i$  reflects the argument error for the `syr2k` with matrices  $A_i$ ,  $B_i$ , and  $C_i$ .

```
BLAS_her2k_batch<fp_t> (
  group_count : int           : In ,
  group_sizes : int [group_count] : In ,
  layout      : enum Layout    : In ,
  uplo       : enum UpLo [count] : In ,
  trans      : enum Transpose [count] : In ,
  n          : int [count]      : In ,
  k          : int [count]      : In ,
  alpha     : fp_t [count]      : In ,
  A         : fp_t [count]      : In ,
  ld_A      : int [count]      : In ,
  B         : fp_t [count]      : In ,
  ld_B      : int [count]      : In ,
  beta     : fp_t [count]      : In ,
  C         : fp_t [count]      : InOut ,
  ld_C      : int [count]      : In ,
  info      : int [count]      : InOut
)
```

This routine is only available for the complex precision. It has the same parameters as the `syr2k` batch routine except that the `trans == BlasTrans` is not allowed in `her2k` batch and that `beta` is real. The matrices  $\{C_i\}$  are complex Hermitian.

The array `info` defines the error array. It is an output array of integers of size `batch_count` where a value at position  $i$  reflects the argument error for the `her2k` with matrices  $A_i$ ,  $B_i$ , and  $C_i$ .

#### 4.1.5 Multiplying a matrix by a triangular matrix TRMM

This routine performs a batch of one of the following matrix-matrix products, where the matrix  $A$  is an upper or lower triangular matrix, and  $\alpha$  is scalar:

- $B \leftarrow \alpha \cdot A \times B; A \in \mathbb{R}^{m \times m}, B \in \mathbb{R}^{m \times n}$  if `side == BlasLeft` and `trans == BlasNoTrans`
- $B \leftarrow \alpha \cdot A^T \times B; A \in \mathbb{R}^{m \times m}, B \in \mathbb{R}^{m \times n}$  if `side == BlasLeft` and `trans == BlasTrans`
- $B \leftarrow \alpha \cdot A^H \times B; A \in \mathbb{C}^{m \times m}, B \in \mathbb{C}^{m \times n}$  if `side == BlasLeft` and `trans == BlasConjTrans`
- $B \leftarrow \alpha \cdot B \times A; A \in \mathbb{R}^{m \times m}, B \in \mathbb{R}^{m \times n}$  if `side == BlasRight` and `trans == BlasNoTrans`
- $B \leftarrow \alpha \cdot B \times A^T; A \in \mathbb{R}^{m \times m}, B \in \mathbb{R}^{m \times n}$  if `side == BlasRight` and `trans == BlasTrans`
- $B \leftarrow \alpha \cdot B \times A^H; A \in \mathbb{C}^{m \times m}, B \in \mathbb{C}^{m \times n}$  if `side == BlasRight` and `trans == BlasConjTrans`



```

BBLAS_trmm_batch_<fp_t>(
  group_count : int           : In,
  group_sizes : int [group_count] : In,
  layout      : enum Layout   : In,
  side        : enum Side [count] : In,
  uplo        : enum UpLo [count] : In,
  trans       : enum Transpose : In,
  diag        : enum Diagonal  : In,
  m           : int [count]    : In,
  n           : int [count]    : In,
  alpha       : fp_t [count]   : In,
  A           : fp_t [count]   : In,
  ld_A        : int [count]    : In,
  B           : fp_t [count]   : In,
  ld_B        : int [count]    : In,
  info        : int [count]    : InOut
)

```

The `side` array is of size `batch_count` and each value defines the operation on each matrix as described in the equations above. The `uplo` array is of size `batch_count` and defines whether the upper or the lower triangular part of the matrices  $\{A_i\}$  are to be referenced. The `trans` is an array of size `batch_count` where each value defines the operation on each matrix. In the real precision case, the values `BlasTrans` and `BlasConjTrans` have the same meaning. The `diag` array is of size `batch_count` where each value defines whether the corresponding matrix  $A$  is assumed to be unit or non-unit triangular. The `m` and `n` arrays of integers are of size `batch_count` and define the dimension of the operation on each matrix. The `alpha` array provides the scalars  $\alpha$  described in the equation above. It is of the same precision as the arrays `A` and `B`. The arrays of pointers `A` and `B` are of size `batch_count` and point to the matrices  $\{A_i\}$  and  $\{B_i\}$ . The size of matrix  $B_i$  is  $m[i]$  by  $n[i]$ . The size of matrix  $A_i$  depends on `side[i]`; its corresponding size is mentioned in the equation above. The arrays `ld_A` and `ld_B` define the leading dimension of the  $\{A_i[\text{ld}_A[i]][*]\}$ , and  $\{B_i[\text{ld}_B[i]][*]\}$  matrices, respective.

The array `info` defines the error array. It is an output array of integers of size `batch_count` where a value at position  $i$  reflects the argument error for the `trmm` with matrices  $A_i$  and  $B_i$ .

#### 4.1.6 Solving triangular systems of equations with multiple right-hand sides TRSM

This routine solves a batch of matrix equations. Each equation is described below, where the matrix  $A$  is an upper or lower triangular matrix, and  $\alpha$  is scalar:

- $B \leftarrow \alpha \cdot A^{-1} \times B; A \in \mathbb{R}^{m \times m}, B \in \mathbb{R}^{m \times n}$  if `side == BlasLeft` and `trans == BlasNoTrans`
- $B \leftarrow \alpha \cdot A^{-T} \times B; A \in \mathbb{R}^{m \times m}, B \in \mathbb{R}^{m \times n}$  if `side == BlasLeft` and `trans == BlasTrans`
- $B \leftarrow \alpha \cdot A^{-H} \times B; A \in \mathbb{C}^{m \times m}, B \in \mathbb{C}^{m \times n}$  if `side == BlasLeft` and `trans == BlasConjTrans`
- $B \leftarrow \alpha \cdot B \times A^{-1}; A \in \mathbb{R}^{m \times m}, B \in \mathbb{R}^{m \times n}$  if `side == BlasRight` and `trans == BlasNoTrans`
- $B \leftarrow \alpha \cdot B \times A^{-T}; A \in \mathbb{R}^{m \times m}, B \in \mathbb{R}^{m \times n}$  if `side == BlasRight` and `trans == BlasTrans`
- $B \leftarrow \alpha \cdot B \times A^{-H}; A \in \mathbb{C}^{m \times m}, B \in \mathbb{C}^{m \times n}$  if `side == BlasRight` and `trans == BlasConjTrans`

```

BLAS_trsm_batch_<fp_t>(

```

```

group_count : int           : In ,
group_sizes : int [group_count] : In ,
layout      : enum Layout   : In ,
side        : enum Side [count] : In ,
uplo        : enum UpLo [count] : In ,
trans       : enum Transpose : In ,
diag        : enum Diagonal  : In ,
m           : int [count]    : In ,
n           : int [count]    : In ,
alpha       : fp_t [count]   : In ,
A           : fp_t [count]   : In ,
ld_A        : int [count]    : In ,
B           : fp_t [count]   : In ,
ld_B        : int [count]    : In ,
info        : int [count]    : InOut
)

```

The `side` array is of size `batch_count` where each value defines the operation on each matrix as described in the equation above. The `uplo` array is of size `batch_count` and defines whether the upper or the lower triangular part of the matrices  $\{A_i\}$  are to be referenced. The `trans` array is of size `batch_count` where each value defines the operation on each matrix. In the real precision case, the values `BlasTrans` and `BlasConjTrans` have the same meaning. The `diag` array is of size `batch_count` where each value defines whether the corresponding matrix  $A$  is assumed to be unit or non-unit triangular. The `m` and `n` arrays of integers are of size `batch_count` and define the dimension of the operation on each matrix. The `alpha` array provides the scalars  $\alpha$  described in the equation above. It is of the same precision as the arrays  $A$  and  $B$ . The arrays of pointers  $A$  and  $B$  are of size `batch_count` and point to the matrices  $\{A_i\}$  and  $\{B_i\}$ . The size of matrix  $B_i$  is `m[i]` by `n[i]`. The size of the matrix  $A_i$  depends on `side[i]`; its corresponding size is mentioned in the equation above. The arrays `ld_A` and `ld_B` define the leading dimension of the matrices  $\{A_i[\text{ld\_A}[i]][*]\}$ , and  $\{B_i[\text{ld\_B}[i]][*]\}$ , respectively.

The array `info` defines the error array. It is an output array of integers of size `batch_count` where a value at position `i` reflects the argument error for the TRSM with matrices  $A_i$  and  $B_i$ .

## 4.2 Scope and Specifications of the Level 1 and Level 2 Batched BLAS

Similarly to the derivation of a Level 3 Batched BLAS from the Level 3 BLAS, we derive Level 1 and Level 2 Batched BLAS from the corresponding Level 1 and Level 2 BLAS routines. Examples are given below for the Level 1 AXPY:  $y \leftarrow \alpha \cdot x + y$  and the Level 2 GEMV:  $y \leftarrow \alpha \cdot A \times x + \beta \cdot y$  BLAS routines.

### 4.2.1 Scaling a vector and adding another vector AXPY

```

BLAS_axpy_batch_<fp_t> (
  group_count : int           : In ,
  group_sizes : int [group_count] : In ,
  n           : int [count]    : In ,
  alpha       : fp_t [count]   : In ,
  X           : fp_t [count]   : In ,
  inc_X       : int [count]    : In ,
  Y           : fp_t [count]   : In ,
  inc_Y       : int [count]    : In ,
  info        : int [count]    : InOut
)

```

Here `inc_X[i]` and `inc_Y[i]` from the  $i^{\text{th}}$  BLAS operation must not be zero and specify the increments for the elements of `X[i]` and `Y[i]`, respectively.

## 4.2.2 General matrix-vector products GEMV

```
BLAS_gemv_batch<fp_t> (  
    group_count : int           : In ,  
    group_sizes : int [group_count] : In ,  
    layout      : enum Layout    : In ,  
    trans_A     : enum Transpose [count] : In ,  
    m           : int [count]    : In ,  
    n           : int [count]    : In ,  
    alpha       : fp_t [count]   : In ,  
    A           : fp_t [count]   : In ,  
    ld_A        : int [count]    : In ,  
    beta        : fp_t [count]   : In ,  
    Y           : fp_t [count]   : InOut ,  
    inc_Y       : int [count]    : In ,  
    info        : int [count]    : InOut  
)
```

Array `inc_Y[i]` at the  $i^{\text{th}}$  position must not be zero and specifies the increment for the elements of `Y[i]`.

## 5 Numerical Stability

Although it is intended that the Batched BLAS be implemented as efficiently as possible, as with the original BLAS, this should not be achieved at the cost of sacrificing numerical stability. See Section 7 of [5] and Section 4.13 of [7].

## 6 Specification of Batch LAPACK Routines

The batch approach to BLAS can be applied to higher-level libraries, and in particular to LAPACK. In this extension, the Batch LAPACK routines are derived from the interfaces of their corresponding non-batch LAPACK routines, similarly to the derivation of Batched BLAS from the classic non-batch BLAS. For example, the specification for the batch LU factorization with partial pivoting based on row interchanges of general M-by-N matrices specified through `A`, is derived from the LAPACK GETRF routine to arrive at the following batch version:

```
LAPACK_getrf_batch<fp_t> (  
    group_count : int           : In ,  
    group_sizes : int [group_count] : In ,  
    layout      : enum Layout    : In ,  
    m           : int [count]    : In ,  
    n           : int [count]    : In ,  
    A           : fp_t [count]   : In ,  
    ld_A        : int [count]    : In ,  
    piv         : int [count]    : In ,  
    info        : int [count]    : InOut  
)
```

## 7 Implementation of the Batched BLAS

The key to efficient BLAS implementation is to hierarchically block the BLAS computation into tasks that operate on data that fits into the corresponding hierarchical memory levels of the computer architecture at hand (see for example the K40 GPU memory hierarchy on Figure 1). The goal is to reduce expensive data movements by loading the data required for a task into fast memory and reusing it in computations from there as many times as possible. An example for achieving this on Level 3 BLAS for GPUs is the MAGMA GEMM [34]. This GEMM harnesses hierarchical blocking on the memory levels available on the Kepler GPUs, including a new register blocking, and is still in use on current GPUs. Hierarchical blocking and communications are needed for optimal performance even for memory-bound computations like Level 2 BLAS, e.g., see the matrix-vector kernels developed and optimized for Xeon Phi architectures [35].

Thus, splitting an algorithm into hierarchical tasks that block the computation over the available memory

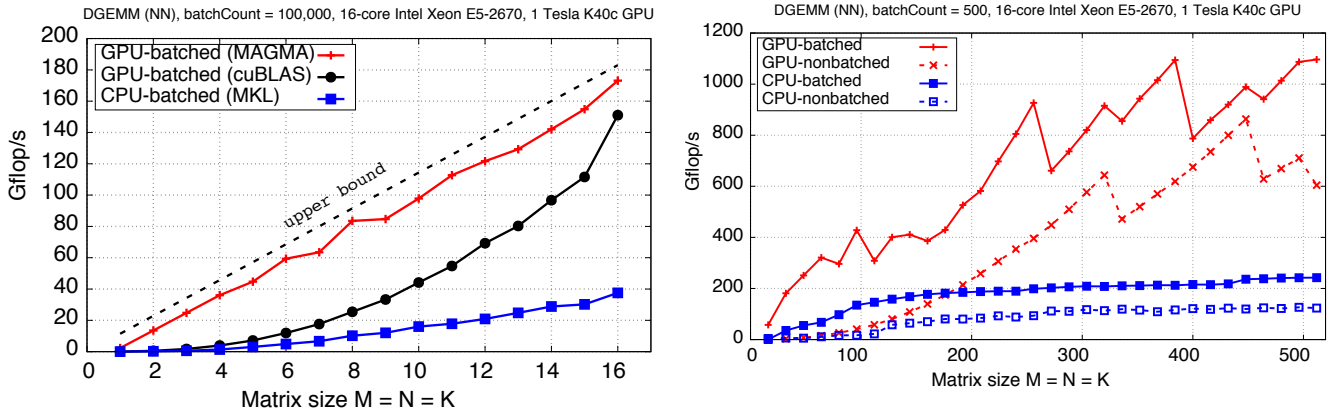


Figure 4: Performance of batch DGEMM versions on matrices of size less than 32 (Left) and larger (Right) on a K40c GPU and 16 cores of Intel Xeon ES-2670 (Sandy Bridge) 2.60 GHz CPUs.

hierarchies (in order to reduce data movement) is essential for implementing high-performance BLAS. Details on how these techniques can be extended to develop high-performance Batched BLAS, and in particular, the extensively used batch GEMM, can be found elsewhere [36]. The routines developed thereby [36] are released through the MAGMA library, providing a model Batched BLAS implementation for GPUs. The goal of this model implementation and the API proposed here is that similarly to BLAS, hardware vendors adopt the Batched BLAS API and maintain highly tuned implementations for their corresponding platforms.

The MAGMA performance is shown in Figure 4. Besides hierarchical blocking, specialized kernels are designed for various sizes, and a comprehensive autotuning process is applied to all kernels. For very small matrix sizes, e.g., sub-vector/warp in size, the performance is memory bound. Techniques like grouping several GEMMs to be executed on the same multiprocessor, vectorization across GEMMs, along with data prefetching optimizations, are used in order to reach 90+% of the theoretical peak on either multicore CPUs or GPUs [37, 38] (see Figure 4, Left). This performance is obtained on CPUs using compiler intrinsics, while on GPUs peak still can be reached by coding in CUDA. For larger sizes on GPUs, e.g., up to about 200 on K40 GPUs, best results are obtained by mapping a single GEMM (from the batch) to a multiprocessor, where the usual hierarchical blocking is applied. For larger matrix sizes, streaming is applied to GEMMs tuned for larger sizes. This results in using more than one multiprocessor for a single GEMM (see Figure 4, Right). For these sizes, similar to CPUs, coding multilevel blocking types of algorithms on GPUs must be in native machine language in order to overcome some limitations of the CUDA compiler or warp scheduler (or both) [39]. Assembly implementations [40, 41] are used today in cuBLAS for Kepler and Maxwell GPUs to obtain higher performance than corresponding CUDA codes. Running these types of implementations through different streams gives the currently best performing batch implementations for large size matrices.

## 8 Future Directions and Final Remarks

Defining a Batched BLAS interface is a response to the demand for acceleration of new (batch) linear algebra routines on heterogeneous and manycore architectures used in current applications. While expressing the computations in applications through matrix algebra (e.g., Level 3 BLAS) works well for large matrices, handling small matrices brings new challenges. The goal of the Batched BLAS is to address these challenges on a library level. The proposed API provides a set of routines featuring BLAS-inspired data storage and interfaces. Similarly to the use of BLAS, there are optimization opportunities for batch computing problems that cannot be folded into the Batched BLAS, and therefore must be addressed separately. For example, these are cases where operands  $\{A_i\}$ ,  $\{B_i\}$ , and  $\{C_i\}$  share data, operands are not directly available in the BLAS matrix format, or where expressing a computation through BLAS may just lose application-specific knowledge about data affinity. For instances where the operands originate from multi-dimensional data, which is a common case, in future work we will look at new interfaces and data abstractions, e.g., tensor-based, where

1. explicit preparation of operands can be replaced by some index operation;

2. operands do not need to be in matrix form, but instead, can be directly loaded in matrix form in fast memory and proceed with the computation from there;
3. expressing computations through BLAS will not lead to loss of information, e.g., that can be used to enforce certain memory affinity or other optimization techniques, because the entire data abstraction (tensor/s) will be available to the routine (and to all cores/multiprocessors/etc.) [37, 42].

Finally, we reiterate that the goal is to provide the developers of applications, compilers, and runtime systems with the option of expressing many small BLAS operations as a single call to a routine from the new batch operation standard. Thus, we hope that this standard will help and encourage community efforts to build higher-level algorithms, e.g., not only for dense problems as in LAPACK, but also for sparse problems as in preconditioners for Krylov subspace solvers, sparse direct multifrontal solvers, etc., using Batched BLAS routines. Some optimized Batched BLAS implementations are already available in the MAGMA library, and moreover, industry leaders like NVIDIA, Intel, and AMD, have also noticed the demand and have started providing some optimized Batched BLAS implementations in their own vendor-optimized libraries.

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