BLISlab: A Sandbox for Optimizing GEMM

FLAME Working Note #80

Use appropriate PDF viewer so that hyperlinks (denoted with ☰) work

Jianyu Huang
Robert A. van de Geijn
Department of Computer Science and
Institute for Computational Engineering and Sciences
The University of Texas at Austin
Austin, TX 78712
{jianyu,rvdg}@cs.utexas.edu

August 31, 2016

Abstract

Matrix-matrix multiplication is a fundamental operation of great importance to scientific computing and, increasingly, machine learning. It is a simple enough concept to be introduced in a typical high school algebra course yet in practice important enough that its implementation on computers continues to be an active research topic. This note describes a set of exercises that use this operation to illustrate how high performance can be attained on modern CPUs with hierarchical memories (multiple caches). It does so by building on the insights that underly the BLAS-like Library Instantiation Software (BLIS) framework by exposing a simplified “sandbox” that mimics the implementation in BLIS. As such, it also becomes a vehicle for the “crowd sourcing” of the optimization of BLIS. We call this set of exercises

1

1 Introduction

Matrix-matrix multiplication (Gemm) is frequently used as a simple example with which to raise awareness of how to optimize code on modern processors. The reason is that the operation is simple to describe, challenging to fully optimize, and of practical importance. In this document, we walk the reader through the techniques that underly the currently fastest implementations for CPU architectures.

1.1 Basic Linear Algebra Subprograms (BLAS)

The Basic Linear Algebra Subprograms (BLAS) [10, 5, 4, 14] form an interface for a set of linear algebra operations upon which higher level linear algebra libraries, such at LAPACK [2] and libflame [19], are built. The idea is that if someone optimizes the BLAS for a given architecture, then all applications and libraries that are written in terms of calls to the BLAS will benefit from such optimizations.

The BLAS are divided into three sets: the level-1 BLAS (vector-vector operations), the level-2 BLAS (matrix-vector operations), and the level-3 BLAS (matrix-matrix operations). The last set benefits from the fact that, if all matrix operands are $n \times n$ in size, $O(n^3)$ floating point operations are performed with $O(n^2)$ data so that the cost of moving data between memory layers (main memory, the caches, and the registers) can be amortized over many computations. As a result, high performance can in principle be achieved if these operations are carefully implemented.

1https://github.com/flame/blislab
1.2 Matrix-matrix multiplication

In particular, GEMM with double precision floating point numbers is supported by the BLAS with the (Fortran) call

\[
dgemm( \text{transa}, \text{transb}, m, n, k, \alpha, A, \text{lda}, B, \text{ldb}, \beta, C, \text{ldc} )
\]

which, by appropriately choosing \text{transa} and \text{transb}, computes

\[
C := \alpha AB + \beta C; \quad C := \alpha A^T B + \beta C; \quad C := \alpha AB^T + \beta C; \quad \text{or } C := \alpha A^T B^T + \beta C.
\]

Here \( C \) is \( m \times n \) and \( k \) is the “third dimension”. The parameters dla, dlb, and dlc are explained later in this document.

In our exercises, we consider the simplified version of GEMM,

\[
C := AB + C,
\]

where \( C \) is \( m \times n \), \( A \) is \( m \times k \), and \( B \) is \( k \times n \). If one understands how to optimize this particular case of dgemm, then one can easily extend this knowledge to all level-3 BLAS functionality.

1.3 High-performance implementation

The intricacies of high-performance implementations are such that implementation of the BLAS in general and GEMM in particular was often relegated to unsung experts who develop numerical libraries for the hardware vendors, for example as part of IBM’s ESSL, Intel’s MKL, Cray’s LibSci, and AMD’s ACML libraries. These libraries were typically written (at least partially) in assembly code and highly specialized for a specific processor.

A key paper [1] showed how an “algorithms and architectures” approach to hand-in-hand designing architectures, compilers, and algorithms allowed BLAS to be written in a high level language (Fortran) for the IBM Power architectures and explained the intricacies of achieving high performance on those processors. The Portable High Performance ANSI C (PHiPAC) [3] project subsequently provided guidelines for writing high-performance code in C and suggested how to autogenerate and tune GEMM written this way. The Automatically Tuned Linear Algebra Software (ATLAS) [17, 18] built upon these insights and made autotuning and autogeneration of BLAS libraries mainstream.

As part of this document we discuss more recent papers on the subject, including the paper that introduced the Goto approach to implementing GEMM [6] and the BLIS refactoring of that approach [16], as well as other papers that are of more direct relevance.

1.4 Other similar exercises

There are others who have put together exercises based on GEMM. Recent efforts relevant to this paper are GEMM: From Pure C to SSE Optimized Micro Kernels by Michael Lehn at Ulm University and a wiki on Optimizing Gemm that we ourselves put together.

1.5 We need you!

The purpose of this paper is to guide you towards high-performance implementations of GEMM. Our ulterior motive is that our BLIS framework for implementing BLAS requires a so-called micro-kernel to be highly optimized for various CPUs. In teaching you the basic techniques, we are hoping to identify “The One” who will contribute the best micro-kernel. Think of it as our version of “HPC’s Got Talent”. Although we focus in our description on optimization for the Intel Haswell architecture, the setup can be easily modified to instead help you (and us) optimize for other CPUs. Indeed, BLIS itself supports architectures that include AMD and Intel’s x86 processors, IBM’s Power processors, ARM processors, and Texas Instrument DSP processors [15, 12, 8].
2 Step 1: The Basics

2.1 Simple matrix-matrix multiplication

In our discussions, we will consider the computation

\[ C := AB + C \]

where \( A, B, \) and \( C \) are \( m \times k, k \times n, m \times n \) matrices, respectively. Letting

\[
A = \begin{pmatrix}
a_{0,0} & \cdots & a_{0,k-1} \\
\vdots & \ddots & \vdots \\
a_{m-1,0} & \cdots & a_{m-1,k-1}
\end{pmatrix},
B = \begin{pmatrix}
b_{0,0} & \cdots & b_{0,n-1} \\
\vdots & \ddots & \vdots \\
b_{k-1,0} & \cdots & b_{k-1,n-1}
\end{pmatrix}, \text{ and } C = \begin{pmatrix}
c_{0,0} & \cdots & c_{0,n-1} \\
\vdots & \ddots & \vdots \\
c_{m-1,0} & \cdots & c_{m-1,n-1}
\end{pmatrix}
\]

\( C := AB + C \) computes

\[ c_{i,j} := \sum_{p=0}^{k-1} a_{i,p} b_{p,j} + c_{i,j}. \]

If \( A, B, \) and \( C \) are stored in two-dimensional arrays \( A, B, \) and \( C, \) the following pseudocode computes \( C := AB + C: \)

```plaintext
for i=0:m-1
    for j=0:n-1
        for p=0:k-1
            C( i, j ) := A( i, p ) * B( p, j ) + C( i, j )
        endfor
    endfor
endfor
```

Counting a multiply and an add separately, the computation requires \( 2mnk \) floating point operations (flops).

2.2 Setup

To let you efficiently learn about how to efficiently compute, you start your project with much of the infrastructure in place. We have structured the subdirectory, step1, somewhat like a project that implements a real library might. This may be overkill for our purposes, but how to structure a software project is a useful skill to learn.

Consider Figure 4, which illustrates the directory structure for subdirectory step1:

**README** Is a file that describes the contents of the directory and how to compile and execute the code.

**sourcemesh.sh** Is a file that configures the environment variables. In that file

- **BLISLAB_USE_INTEL** sets whether you use the Intel compiler (**true**) or the GNU compiler (**false**).
- **BLISLAB_USE_BLAS** indicates whether your reference **dgemm** employs an external BLAS library implementation (**true**) if you have such a BLAS library installed on your machine, or the simple triple loops implementation (**false**).
- **COMPILER_OPT_LEVEL** sets the optimization level for your GNU or Intel compiler (**O0**, **O1**, **O2**, **O3**). (Notice that, for example, **O3** consists of the capital letter "O" and the number "3".)
- **OMP_NUM_THREADS** and **BLISLAB_IC_NT** sets the number of threads used for parallel version of your code. For Step 1, you set them both to 1.

**dgemm** Is the subdirectory where the routines that implement **dgemm** exist. In it

- **bldgmemm_ref.c** contains the routine **dgemm_ref** that is a simple implementation of **dgemm** that you will use to check the correctness of your implementations, if **BLISLAB_USE_BLAS** = **false**.
- **my_dgmemm.c** contains the routine **dgemm** that that initially is a simple implementation of **dgemm** and that you will optimize as part of the first step on your way to mastering how to optimize **gemm**.
Figure 1: Structure of directory step1.

- **bl_dgemm_util.c:** contains utility routines that will later come in handy.

- **include:** This directory contains include files with various macro definitions and other header information.

- **lib:** This directory will hold libraries generated by your implemented source files (libblislab.so and libblislab.a). You can also install a reference library (e.g. OpenBLAS) in this directory to compare your performance.

- **test:** This directory contains “test drivers” and correctness/performance checking scripts for the various implementations.
  - **test_bl_dgemm.c:** contains the “test driver” for testing routine bl_dgemm.
  - **test_bl_dgemm.x:** is the executable file for test_bl_dgemm.c.
  - **run_bl_dgemm.sh:** contains a bash script to collect performance results.
  - **tacc_run_bl_dgemm.sh:** contains a SLURM script for you to (optionally) submit the job to the Texas Advanced Computing Center (TACC) machines if you have an account there.

### 2.3 Getting started

What we want you to do is to start with the implementation in my_dgemm.c and optimize it by applying various standard optimization techniques. The initial implementation in that file is the straight-forward implementation with the three loops given in Figure 2. The first thing to notice is how two-dimensional arrays are mapped to memory in so-called column-major order. The reason for this choice is that the original BLAS assumed column-major storage of arrays because the interface was for Fortran users first. Examining

\[ C( i, j ) += A( i, p ) \times B( p, j ); \]
we notice that, each operand is a MACRO. Consider early in that file

#define C( i, j ) C[ (j)*ldc + (i) ]

The linear array at address $C$ is used to store elements $C_{i,j}$ so that the $i,j$ element is mapped to location $j * ld + i$. The way to view this is that the columns of $C$ are each stored contiguously. However, think of matrix $C$ as embedded in a larger array that has $ld$ rows so that accessing a row means going through array $C$ with stride $ld$. The term leading dimension of two-dimensional array $C$ is typically used to refer to the row dimension of this larger array, hence the variable $ld$ (leading dimension of $C$). This is illustrated for all three matrices in the following figure:

in which the arrows are meant to indicate that columns are stored contiguously.

2.3.1 Configure the default implementation

By default, the exercise compiles and links with Intel’s icc compiler, which will apply compiler optimizations (level O3) to the code. You need to set the environment variable by executing:

```bash
$ source sourceme.sh
```

in the terminal, and you will see the output:

```
BLISLAB_USE_INTEL = true
COMPILER_OPT_LEVEL = O3
```

2.3.2 Compile, execute and collect results

If you do not have access to Intel’s compiler (icc), then read Subsections 2.3.2 and 2.3.3 and continue with Subsection 2.3.5.

You can compile, execute your code and collect the performance result by executing

```bash
make clean
make
```
cd test
./run_bl_dgemm.sh

in subdirectory step1. You will see the performance result output:

```
run_step1_st="
%m %k %n %MY_GFLOPS %REF_GFLOPS
16 16 16 0.82 2.15
32 32 32 0.74 5.50
48 48 48 0.85 5.66
......
];
```

You can change the sampling block size in `run_bl_dgemm.sh`. Notice that if you have errors in your code, these will be reported as, for example,

```
C[ 0 ][ 0 ] != C_ref, 1.253000E+00, 2.253000E+00
```

### 2.3.3 Draw the performance graph

Finally, you can use MATLAB to draw your performance graph with our scripts. In `misc/results` subdirectory, after executing

```
./collect_result_step1
```

you will get a MATLAB file “step1_result.m”, with the performance results. You can then execute

```
bl_dgemm_plot.m
```

in MATLAB, which will then generate the performance graph.

### 2.3.4 Change to the GNU compiler

Since we want you to explicitly learn about what kind of tricks lead to high performance, and because some of you may not have access to the Intel compiler, you should next change to using the GNU C compiler. For this, you must edit `sourceme.sh`:

```
BLISLAB_USE_INTEL=false
```

Then, similar to the default setting, you need to set the environment variable by executing:

```
$ source sourceme.sh
```

in the terminal, and you will observe:

```
BLISLAB_USE_INTEL = false
COMPILER_OPT_LEVEL = 03
```

### 2.3.5 Turn off optimization

Next, we want you to turn off the optimization performed by the compiler. This serves three purposes: first, it means you are going to have to explicitly perform optimizations, which will allow you to learn about how architectures and algorithms interact. Second, it may very well be that the optimizing compiler will try to
“undo” what you are explicitly trying to accomplish. Third, the more tricks you build into your code, the harder it gets for the compiler to figure out how to optimize.

You need first edit `sourceme.sh`:

```bash
COMPILER_OPT_LEVEL = 00
```

Then, similar to the default setting, you need to set the environment variable by executing:

```bash
$ source sourceme.sh
```

in the terminal, and you will see the output:

```bash
BLISLAB_USE_INTEL = false
COMPILER_OPT_LEVEL = 00
```

### 2.3.6 (Optional) Use optimized BLAS library as reference implementation

By default, your reference GEMM implementation is a very slow triple-loop implementation. If you have a BLAS library installed on your test machine, you can adopt the `dgemm` from that library as your reference implementation by setting:

```bash
BLISLAB_USE_BLAS=true
```

in `sourceme.sh`. If you use Intel compiler, you don’t need to explicitly specify the path of MKL. However, if you use GNU compiler, you need to specify the path of your BLAS library. For example, you may want to install our BLIS library from [https://github.com/flame/blis](https://github.com/flame/blis) in directory `/home/lib/blis` and in `sourceme.sh` set

```bash
BLAS_DIR=/home/lib/blis
```

After executing `$ source sourceme.sh`, you will observe:

```bash
BLISLAB_USE_BLAS = true
BLAS_DIR = /home/lib/blis
```

and now performance and accuracy comparisons of your implementation will be against this optimized library routine.

## 2.4 Basic techniques

In this subsection we describe some basic tricks of the trade.

### 2.4.1 Using pointers

Now that optimization is turned off, the computation of the address where an element of a matrix exists is explicitly exposed. (An optimizing compiler would get rid of this overhead.) What you will want to do is to change the implementation in `my_gemm.c` so that it instead uses pointers. Before you do so, you may want to back up the original `my_gemm.c` in case you need to restart from scratch. Indeed, at each step you may want to back up in a separate file the previous implementations.

Here is the basic idea. Let’s say we want to set all elements of \( C \) to zero. A basic loop, patterned after what you found in `my_gemm.c` might look like

```c
for ( i = 0; i < m; i ++ ) {
    for ( j = 0; j < n; j ++ ) {
```
\( C( i, j ) = 0.0; \)
}
}

Using pointers, we might implement this as

```c
double *cp;

for ( j = 0; j < n; j ++ ) {
    cp = &C[ j * ldc ]; // point cp to top of jth column
    for ( i = 0; i < m; i ++ ) {
        *cp++ = 0.0; // set the element that cp points to to zero and
        // advance the pointer.
    }
}
```

Notice that we purposely exchanged the order of the loops so that advancing the pointer takes us down the columns of \( C \).

### 2.4.2 Loop unrolling

Updating loop index \( i \) and the pointer \( cp \) every time through the inner loop creates considerable overhead. For this reason, a compiler will perform *loop unrolling*. Using an unrolling factor of four, our simple loop for setting \( C \) to zero becomes

```c
double *cp;

for ( j = 0; j < n; j ++ ) {
    cp = &C[ j * ldc ];
    for ( i = 0; i < m; i += 4 ) {
        *(cp+0) = 0.0;
        *(cp+1) = 0.0;
        *(cp+2) = 0.0;
        *(cp+3) = 0.0;
        cp += 4;
    }
}
```

Importantly:

- \( i \) and \( cp \) are now only updates once every four iterations.
- \(*(cp+0)\) uses a machine instruction known as *indirect addressing* that is much more efficient than if one computed with \(*(cp+k)\) where \( k \) is a variable.
- When data it brought in for memory into cache, it is brought in a cache line of 64 bytes at a time. This means that accessing contiguous data in chunks of 64 bytes reduces the cost of memory movement between the memory layers.

Notice that when you unroll, you may have to deal with a “fringe” if, in this case, \( m \) is not a multiple of four. For the sake of this exercise, you need not worry about this fringe *as long as you pick your sampling block size wisely*, as reiterated in Section 2.5.

### 2.4.3 Register variables

Notice that computation can only happen if data is stored in registers. A compiler will automatically transform code so that the intermediate steps that place certain data in registers is inserted. One can give
a hint to the compiler that it would be good to keep certain data in registers as illustrated in the following somewhat contrived example:

```c
double *cp;

for ( j = 0; j < n; j ++ ) {
    cp = &C[j * ldc ];
    for ( i = 0; i < m; i += 4 ) {
        register double c0=0.0, c1=0.0, c2=0.0, c3=0.0;
        *(cp+0) = c0;
        *(cp+1) = c1;
        *(cp+2) = c2;
        *(cp+3) = c3;
        cp += 4;
    }
}
```

2.5 A modest first goal

We now ask you to employ the techniques discussed above to optimize my _dgemm_. For now, just worry about trying to attain better performance for smallish matrices. In particular, consider the following picture:

What we want you to do is to write your code so that \( m_R \times n_R \) blocks of \( C \) are kept in registers. You get to choose \( m_R \) and \( n_R \), but you will want to update file `include/bl_config.h` with those choices. This ensures that the test driver only tries problem sizes that are multiples of these block sizes, so you don’t have to worry about “fringe”.

You will notice that even for smallish matrices that can fit in one of the cache memories, your implementation performs (much) worse than the implementations that are part of MKL or other optimized BLAS library that you may have installed. The reason is that the compiler is not using the fastest instructions for floating point arithmetic. These can be accessed either by using _vector intrinsic functions_, which allows you to explicitly utilize them from C, or by coding in assembly code. For now, let’s not yet go there. We will talk more about this in Step 3.

3 Step 2: Blocking

3.1 Poorman’s BLAS

Step 1 of this exercise makes you realize that with the advent of cache-based architectures, high-performance implementation of GEMM necessitated careful attention to the amortization of the cost of data movement between memory layers and computation with that data. To keep this manageable, it helps to realize that only a “kernel” that performs a matrix-matrix multiplication with relatively small matrices needs to be highly optimized, since computation with larger matrices can be blocked to then use such a kernel without an adverse impact on overall performance. This insight was explicitly advocated in [9]

This is sometimes referred to as "poorman’s BLAS" in the sense that if one could only afford to optimize matrix-matrix multiplication (with submatrices), then one could build Gemm, and other important matrix-matrix operations known as the level-3 BLAS, in terms of this. What we will see later is that actually in general this is a good idea, for the sake of modularity as well as for performance.

In the last section you already saw an example of blocking.

3.2 Blocked matrix-matrix multiplication

Key to blocking Gemm to take advantage of the hierarchical memory of a processor is understanding how to compute $C := AB + C$ when these matrices have been blocked. Partition

$$A = \begin{pmatrix} A_{0,0} & \cdots & A_{0,K-1} \\ \vdots & \ddots & \vdots \\ A_{M-1,0} & \cdots & A_{M-1,K-1} \end{pmatrix}, B = \begin{pmatrix} B_{0,0} & \cdots & B_{0,N-1} \\ \vdots & \ddots & \vdots \\ B_{K-1,0} & \cdots & B_{K-1,N-1} \end{pmatrix}, \text{ and } C = \begin{pmatrix} C_{0,0} & \cdots & C_{0,N-1} \\ \vdots & \ddots & \vdots \\ C_{M-1,0} & \cdots & C_{M-1,N-1} \end{pmatrix},$$

where $C_{i,j}$ is $m_i \times n_j$, $A_{i,p}$ is $m_i \times k_p$, and $B_{p,j}$ is $k_p \times n_j$. Then

$$C_{i,j} := \sum_{p=0}^{K-1} A_{i,p}B_{p,j} + C_{i,j}.$$  

3.3 Your mission, if you choose to accept it

We now ask you to implement the blocked matrix-matrix multiplication in my dgemm. Specifically, for small matrices you achieve better performance than for larger matrices because the smaller matrices fit in cache. Block the matrices into submatrices of the size for which you do attain higher performance, and you will see that the resulting implementation can maintain the better performance even for larger matrices.

4 Step 3: Blocking for Multiple Levels of Cache

4.1 The Goto Approach to Implementing Gemm

Around 2000, Kazushige Goto revolutionized how Gemm is implemented on current CPUs with his techniques that were first published in the paper [6].


A further "refactoring" of this approach was more recently described in [16].


The advantage of the BLIS framework is that it reduces the kernel that must be highly optimized, possibly with vector intrinsics or in assembly code, to a micro-kernel. In this section, we briefly describe the highlights of the approach. However, we strongly suggest the reader become familiar with the above two papers themselves.

Figure 3 (left) illustrates the way the Goto approach structures the blocking for three layers of cache (L1, L2, and L3). In the BLIS framework, the implementation is structured exactly this way so that only the micro-kernel at the bottom needs to be highly optimized and customized for a given architecture. In the original GotoBLAS implementation, now maintained as OpenBLAS [11], the operation starting with the second loop around the micro-kernel is instead customized. In order to get the best performance, it helps is all data is accessed contiguously, which is why at some point prior to reaching the micro-kernel, data is packed in the order indicated by the arrows:
Now, notice that each column of the block of $A$ in the above picture is multiplied by each element in the corresponding row of the block of $B$. (We call these blocks of $A$ and $B$ micro-panels.) This means that the latency to the L2 cache (the time required to bring in an element of the micro-panel of $A$ from that cache) can be amortized over $2n_B$ flops. For this reason, we can organize the computation so that the micro-panel of $A$ typically resides in the L2 cache. Actually, we can do better: while a rank-1 update is happening with $\tilde{A}$, that has been packed to reside in the L3 cache (by virtue of how the packing into $\tilde{A}$ is organized). We can do this because the packing into $\tilde{A}$ is amortized over all computations with $\tilde{B}_p$ and the packing into $\tilde{B}_p$ is amortized over computations with many blocks $A_i$. The outermost loop partitions $B$ so that the block $\tilde{B}_p$ fits in the L3 cache or, if a processor does not have an L3 cache, limits the amount of workspace for packing $\tilde{B}_p$ that is needed. This

Figure 3: Left: The GotoBLAS algorithm for matrix-matrix multiplication as refactored in BLIS. Right: the same algorithm, but expressed as loops.

```
Loop 0 for $k_c = 0 : k_c - 1$
    $C_{c_i}(I_{r,c},J_{r}) \leftarrow A_{c}(I_{r},k_r) \cdot B_{c}(k_r,J_{r})$
endfor
```

```
Loop 1 for $i_c = 0 : m_c - 1$ steps of $m_c$
    $I_c = i_c : i_c + m_c - 1$
    $A(I_c, P_c) \rightarrow A_e$ // Pack into $A_e$
    $T_c = i_c : i_c + m_c - 1$
    $B(T_c, P_c) \rightarrow B_e$ // Pack into $B_e$
endfor
```

```
Loop 2 for $j_r = 0 : n_r - 1$ steps of $n_r$
    $J_r = j_r : j_r + n_r - 1$
endfor
```

```
Loop 3 for $i_c = 0 : m_c - 1$ steps of $m_c$
    $T_c = i_c : i_c + m_c - 1$
    $A(I_c, P_c) \rightarrow A_e$ // Pack into $A_e$
    $B(T_c, P_c) \rightarrow B_e$ // Pack into $B_e$
endfor
```

```
Loop 4 for $p_c = 0 : k_c - 1$
    $P_c = p_c : p_c + k_c - 1$
    $B(P_c, J_{p,c}) \rightarrow B_e$ // Pack into $B_e$
endfor
```

```
Loop 5 for $j_c = 0 : n - 1$ steps of $n_c$
    $J_c = j_c : j_c + n_c - 1$
endfor
```

Now, notice that each column of the block of $A$ in the above picture is multiplied by each element in the corresponding row of the block of $B$. (We call these blocks of $A$ and $B$ micro-panels.) This means that the latency to the L2 cache (the time required to bring in an element of the micro-panel of $A$ from that cache) can be amortized over $2n_B$ flops. For this reason, we can organize the computation so that the micro-panel of $A$ typically resides in the L2 cache. Actually, we can do better: while a rank-1 update is happening with a column of the micro-panels of $A$ and $B$, the next column of the micro-panel of $A$ can be brought into registers so that computation masks the cost of that data movement. The fact that we want to keep the micro-panel of $B$ in the L1 cache (because it will be reused for many micro-panels of $A$) limits the blocking parameter $k_C$.

With the insights, the rest of the picture hopefully becomes clear. The first loop around the microkernel works with a block of $A$, $\tilde{A}_i$, that has been packed and resides in the L2 cache (by virtue of how the computation is ordered). This limits the blocking parameter $m_C$. That block of $A$ multiplies a block of $B$, $\tilde{B}_p$, that has been packed to reside in the L3 cache (if the processor has an L3 cache). Notice that the packing into $\tilde{A}_i$ is amortized over all computation with $\tilde{B}_p$ and the packing into $\tilde{B}_p$ is amortized over computations with many blocks $A_i$. The outermost loop partitions $B$ so that the block $\tilde{B}_p$ fits in the L3 cache or, if a processor does not have an L3 cache, limits the amount of workspace for packing $\tilde{B}_p$ that is needed. This
limits the blocking parameter $n_C$.

One may ask if the above described scheme is optimal. In [7] a theory is given that shows that under an idealized model the above is locally optimal (in the sense that assuming data is in a certain memory layer in the hierarchy, the proposed blocking at that level optimally amortizes the cost of data movement with the next memory layer). A theory that guides the choice of the various blocking parameters is given in [13].

4.2 Setup

Figure 4 illustrates the directory structure for subdirectory step3. Comparing to step1, we have modified/added the following directories/files:

- **kernels** This directory contains the micro-kernel implementations for various architecture.
  - `bl_dgemm_ukr.c` gives a naive C implementation.
  - `bl_dgemm_int_kernel.c` gives an AVX/AVX2 intrinsics micro-kernel implementation for Haswell architecture.
  - `bl_dgemm_asm_kernel.c` gives an AVX/AVX2 assembly micro-kernel implementation for Haswell architecture.

4.3 Advanced techniques

You can find the vector instructions online:

- [Intel Intrinsics Guide](#)
- [Intel ISA Extensions](#)
4.3.1 An introduction example for “axpy”

We provide you an example for the implementation of “axpy” to demonstrate how to use Intel AVX Intrinsics and Assembly (in misc/examples subdirectory). This example will serve as a good start point for you to learn basic broadcast/fma/load/store instructions. Moreover, this example is actually a primitive for the “broadcast” implementation for 4×4 rank-1 update.

4.3.2 4×4 rank-1 update

The micro-kernel implementation can be boiled down to a 4×4 rank-1 update. There are two possible implementations: one based on broadcast (Figure 5) and one of a butterfly permutation (Figure 6). You can also try other possible implementations.

4.4 Your mission, if you choose to accept it

We provide you a reference implementation of simplified BLIS framework in my_dgemm. The code is organized in the same way presented in Figure 3. However, the step size in each loop is not well chosen, and the micro-kernel implementation is a naive C version. Therefore, you will not expect high performance with the code. What we want you to do is to

- Specify the blocking parameter \( m_C, n_C, k_C \) and the micro-kernel size parameter \( m_R, n_R \) in the file include/bl_config.h; and
Figure 6: AVX 4×4 rank-1 update with butterfly permutation. Given 4×1 vector $A$ and $B$, we compute the 4×4 outer-product $C$ by 4 FMA interleaved with vectorized shuffling operations. The 3rd operands (0x5, 0x1) indicates the shuffling (permutation) type.

- Implement the efficient micro-kernel with vector intrinsics or assembly code. Place the code in kernels/bl_dgemm_int_kernel.c (for vector intrinsics), or kernels/bl_dgemm_asm_kernel.c (for assembly). You need to specify the function name of the micro-kernel by modifying BL_MICRO_KERNEL in include/bl_config.h.

5 Step 4: Parallelizing with OpenMP

The benefit of the BLIS way of structuring the GotoBLAS approach to the implementation of GEMM is that it exposes five loops in tt C which can then be easily parallelized with OpenMP directives.

5.1 To parallelize or not to parallelize, that’s the question

The fundamental question becomes which loop to parallelize. In [12]


a detailed discussion is given of what the pros and cons are regarding the parallelization of each loop. For multi-core architectures (multi-threaded architectures with relatively few cores) results can be found in the earlier paper [15]

6 Conclusion

We use GEMM as a case study to show how to program for performance.

Useful links

Documentation

- The Science of High-Performance Computing (SHPC) group website
- The FLAME project publications webpage (The umbrella project that BLIS is part of is known as the FLAME project.)
- Intel Intrinsics Guide
- Intel ISA Extensions

Software

- BLIS on GitHub
- Intel Free Software (including C/C++ compilers).
- Intel’s Math Kernels Library (MKL) website
- Download MKL for free.

Acknowledgments

We thank the other members of the Science of High-Performance Computing (SHPC) team for their support. This research was partially sponsored by the National Science Foundation grant ACI-1148125/1340293 and Intel through its funding of the SHPC group as an Intel Parallel Computing Center.

Any opinions, findings and conclusions or recommendations expressed in this material are those of the author(s) and do not necessarily reflect the views of the National Science Foundation (NSF).

References


