1 Background and Motivation

Tensors, or multi-dimensional arrays, are structures that generalize vectors and matrices to higher dimensions. Specifically, $\mathbf{X} \in \mathbb{R}^{N_1 \times N_2 \times \cdots \times N_D}$ is an $D$-dimensional tensor with elements $x_{i_1,\ldots,i_D} \in \mathbb{R}$ where $1 \leq i_d \leq N_d$ for all $1 \leq d \leq D$. Tensor-based computations have witnessed a remarkable increase in popularity in the last two decades due to great advancements in tensor theory and algorithms as well as their expressive power for high-dimensional algebraic structures that appear in fundamental applications in computational biology, chemistry, physics, signal processing, data analysis, and machine learning. Tensors are particularly useful when the problem at hand either has or can be reformulated to exhibit a high-dimensional structure. In this formulation, “high-dimensional” matrices and vectors of size $N^D \times N^D$ and of size $N^D$, respectively, are expressed as $2D$- and $D$-dimensional tensors, respectively, of size $N$ in each dimension. In most cases, the inherent structure and properties of the problem that form these matrices/vectors (and corresponding tensors) imply a “low-rank” property; matrices and vectors can be expressed using polynomial number of elements in $N$ and $D$ (instead of $N^{2D}$ or $N^D$ elements), which is rendered possible via so-called tensor decompositions. Once matrices and vectors are expressed in this form, all matrix and vector operations such as matrix-vector multiplication and basic vector arithmetic (addition, subtraction, multiplication/division by a scalar, inner product) are performed under this “compressed” scheme with tremendous gains in terms of computational and memory costs.

One of the most popular tensor decompositions used in this context is called tensor-train decomposition. A matrix of size $N^D \times N^D$ is expressed via tensor-train network of $D$ 4-dimensional tensors of size $N \times N \times R \times R$, where $R$ is the rank of the matrix in the tensor-train form. Similarly, a vector of size $N^D$ comprises $D$ 3-dimensional tensors of size $N \times R \times R$. An analogous way to interpret these tensors would be to consider them as matrices (or vectors) of size $N \times N$ (or $N$), where each element in the matrix (vector) is a matrix of size $R \times R$. We will be exclusively working on a fundamental kernel involving the multiplication of such matrix/vector of matrices described in what follows.

2 Matrix-vector multiplication in Kronecker form

Matrix-vector multiplication is a fundamental computational kernel in scientific computing whose optimization plays a key role in obtaining high performance linear and non-linear solvers. For a given matrix $\mathbf{A} \in \mathbb{R}^{M \times N}$ and a vector $\mathbf{x} \in \mathbb{R}^N$, the multiplication $\mathbf{y} = \mathbf{A}\mathbf{x}$ is defined as $y_i = \sum_{j=1}^{N} A_{i,j} x_j$ where $\mathbf{y} \in \mathbb{R}^M$.

Here, we will consider a generalized version of the standard matrix-vector multiplication, involving a matrix of matrices and a vector of matrices, i.e., $\mathbf{A}_{i,j}, \mathbf{x}_j$, and $\mathbf{y}_i$ are matrices. In describing this computation, we are in need of a special operation called Kronecker product on matrices, which is defined in what follows.

The Kronecker product of two matrices $\mathbf{B} \in \mathbb{R}^{M_B \times N_B}$ and $\mathbf{C} \in \mathbb{R}^{M_C \times N_C}$ is denoted by the operation $\otimes$ and results in a matrix $\mathbf{D} = \mathbf{B} \otimes \mathbf{C} \in \mathbb{R}^{M_B M_C \times N_B N_C}$, which is defined as

$$
\mathbf{D} = \mathbf{B} \otimes \mathbf{C} = \begin{pmatrix}
B_{1,1} \mathbf{C} & \cdots & B_{1,N_B} \mathbf{C} \\
\vdots & \ddots & \vdots \\
B_{M_B,1} \mathbf{C} & \cdots & B_{M_B,N_B} \mathbf{C}
\end{pmatrix}
$$

We now define an operation we call matrix-vector multiplication in Kronecker form (MxVK) as follows. Given a matrix of $N \times N$ matrices $\mathbf{A}$ such that $\mathbf{A}_{i,j} \in \mathbb{R}^{R_A, Q_A}$ for some fixed $R_A, Q_A$ for all $1 \leq i, j \leq N$, and a vector of $N$ matrices $\mathbf{x}$ such that $\mathbf{x}_j \in \mathbb{R}^{R_x, Q_x}$ for some fixed $R_x, Q_x$ for all $1 \leq j \leq N$, the result of the MxVK yields a vector $\mathbf{y} = \mathbf{A}\mathbf{x}$ with entries

$$
y_i = \sum_{j=1}^{N} A_{i,j} \otimes x_j, \quad \mathbf{y}_i \in \mathbb{R}^{R_A R_x \times Q_A Q_x}
$$

Note that when $R_A = R_x = 1$, MxVK reduces to standard matrix-vector multiplication as $\mathbf{A}_{i,j} \mathbf{x}_j$ and $\mathbf{y}_i$ become scalars and Kronecker product reduces to scalar multiplication. We provide a pictorial representation of this operation in the following figure in which the computation of $\mathbf{y}_2 = \sum_{j=1}^{N} \mathbf{A}_{2,j} \otimes \mathbf{x}_j$ is highlighted.
MxVK is a fundamental kernel in low-rank tensor computations using tensor-train decomposition; carrying out matrix-vector multiplies in this form constitutes one of the most expensive steps in the context of an iterative solver. The goal in this task is to implement very efficient MxVK kernels and effectively parallelize them with different paradigms using OpenMP.

You are already given a skeleton code `mxvk.c` that accepts the parameters \( N, R_A, Q_A, R_x, Q_x \) then creates a matrix \( A \) and vectors \( x \) and \( y \) with these parameters, runs the MxVK function on these matrices to fill the result vector \( y \), and checks the correctness of this result. Matrices and vectors are stored in data structures provided in `matmat.c/.h` and `vecmat.c/.h`. Here, \( A \) is stored in row-major format; meaning in the memory \( A_{i,1}, A_{i,2}, \ldots, A_{i,N} \), which is followed by the second row \( A_{2,1}, \ldots, A_{2,N} \), etc. Each element \( A_{i,j}, x_j, \) and \( y_i \) are stored, however, in column-major format, i.e., \( x_j(:,1) \) is stocked first, followed by \( x(:,2) \), etc.

Do not hesitate to take your time to read the provided code and have a look at matrix/vector data structures. You will see in the code that MxVK function utilizes another subroutine to perform the Kronecker product of two matrices as in (1), then performs the MxVK using this subroutine as in (2).

\[ A \times y \]
\[ \text{RA} \]
\[ \text{RARx} \]

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\[ 3 \text{ Tasks} \]

\[ 3.1 \text{ Optimizing sequential MxVK} \]

In this first task, your goal is to optimize the given MxVK kernel as much as possible. Take particularly into consideration the fact that matrices \( A_{i,j} \) and \( X_j \) are relatively small (e.g. \( \leq 64 \) in each dimension). A good start would be determining the total number of flops performed in MxVK, the total amount of data “touched” during this computation (or total mops (memory operations)) in terms of \( N, R_A \), and \( R_x \). Finally, \( \#\text{flops}/\#\text{mops} \) gives the computational intensity of this operation, which you should aim to attain with an optimized implementation.

Explain each optimization technique you employed, and report the time difference with respect to previous baseline. Also report how well your code performs with respect to different set of parameters (i.e., small or large \( N, R_A, R_x \)), and try to fine-tune your code for edge cases if possible.

You are NOT allowed to use any sort of parallelization for this task. You are indeed allowed, however, to employ this optimized kernel in the following tasks requiring parallelization.

\[ 3.2 \text{ Parallelization using OpenMP} \]

Now that you have a working sequential implementation, it is time to optimize it to the fullest and obtain the best performance. Here, you will parallelize the optimized sequential implementation you provided in the previous case using OpenMP parallel constructs. You are free to use parallel loops or sections. You are also welcome to make significant changes to your sequential implementation, in which case you would need to report why you needed such a change from a performance point of view. Make sure your parallelization scales acceptably well for pathological input sizes as well (i.e., \( N_A \) very small, \( R_A \) small, \( R_x \) large, etc.). Explain your parallelization strategy, how you divide the work among threads, achieve load balance, handle data dependencies, perform synchronization (if necessary), etc.

\[ 3.3 \text{ Parallelization using OpenMP tasks} \]

You are NOT allowed to have any other means of parallelization for this assignment except OpenMP tasks.

You will now do another parallel implementation, but this time using OpenMP tasks. Note that your implementation should NOT be a simple re-wrapping of your previous parallelization with loops; you should try to extract
finer-grain parallel tasks that have elegant output independence hence can be executed in parallel efficiently, but are coarse enough that it would not kill the performance due to tasking overhead. If you are new to OpenMP task-based programming, you can have a look at the quick introduction and examples we provided to get going.

4 General guidelines and considerations

- You can assume that $1 \leq R_A, R_x \leq 64$, and $1 \leq N \leq 1024$.

- It might be useful to take into consideration the machine parameters (particularly the cache size at different levels, which you can query using `lscpu` command on Linux).

- Grading will be based on the speed of execution of your kernels. We will run each kernel 100 times and take the average. You should also be writing a report summarizing your approaches for optimization, and performance gains you obtained from each approach. (Grading criteria to be completed...)

- Since we focus on parallelization, you are also encouraged to parallelize the work among the group members. For instance, sequential optimization and parallelization could be done in parallel!