High Performance Computing Project - A.A. 2022/2023

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September 20, 2023

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Chapter 1

Benchmarking MKL, OpenBLAS, and BLIS

1.1 Introduction

In this exercise we compare the performance of three High Performance Libraries (HPC): MKL, Open-BLAS, and BLIS. In particular, we focus on the level 3 BLAS function called gemm, which multiples an $m \times k$ matrix A times a $k \times n$ matrix B and stores the result in an $m \times n$ matrix C.

This function comes in two types, one for single precision (float) and the other for double precision (double). Furthermore, it is capable of exploiting paralellism using OpenMP (OMP) to speed up the calculations, provided that we have the required computational resources.

Using squared matrices only, we perform a scalability study in two scenarios. In the first scenario, we fix the number of cores, and increase the size of the matrices from 2000 to 20000. In the second scenario, we fix the matrix size to 10000 and increase the number of cores that gemm can use by modifying the OMP_NUM_THREADS environment variable.

In both scenarios, we repeat the measurements for both single and double precision, for both THIN and EPYC nodes, using the maximum number of cores.

Furthermore, for the second scenario, we also modify the thread affinity policy of OMP in order to observe any differences.

1.2 Methodology

1.2.1 Compiling BLIS and obtaining binaries

We begin by downloading the BLIS library by using the following commands:

Where NODE can be specified as either THIN or EPYC and P are the available cores for each node, 24 and 128 respectively.

With these commands, we have compiled the BLIS library for the desired architecture.

Next, we specify the flag in the Makefile to compile for float or double using DUSE_FLOAT or -DUSE_DOUBLE. Then, we run:

```
$\salloc -n \{P\} -N1 -p \{NODE\} --time=1:0:0

$\module load mkl/latest
$\module load openBLAS/0.3.23-omp
$\export LD_LIBRARY_PATH=/path/to/myblis/lib:\$LD_LIBRARY_PATH
$\srun -n1 make cpu
```

Which will generate the binaries for the desired architecture, with floar or double precision, depending on the flag we used.

To run, we use:

At the end of this procedure, we should have the appropriate binaries for each architecture, and for each type of precision, double or float.

We now detail the steps to obtain the measurements for both scenarios.

1.2.2 Using a fixed number of cores

For this section, we use all the cores available in a THIN or an EPYC node: 24 and 128, respectively. Since we only use squared matrices, we can describe the dimensions of the matrices with a single number, which we call "size".

For both architectures, we start with a size of 2000 and end with a size of 20000, with jumps of 2000 for a total of 10 sizes. For each size, we repeat the measurement 10 times and report the average and standard deviation.

Finally, we repeat the measurements for both floating point precision and double point precision.

The scripts that were used can be found in the folder exercise2/scripts, under the name es2_1_thin.sh and es2_1_epyc.sh.

It is important to observe that in this section, since we are using the entire node, there is little possibility to play with combinations of thread affinity.

This will be done for the next section.

Furthermore, contrary to the guidelines for the exercise, we decided to use the entire node to benchmark its full capacity, and also to avoid wasting resources.

In fact, to obtain an accurate benchmark, we need to reserve the whole node, regardless of the number of cores we decide to use. This is because if other people began to use the other half of the node, this could introduce additional workloads which interfere with the benchmark.

1.2.3 Using a fixed matrix size

For this section, we fix the size of the matrices to 10000. Then, we slowly increase the number of cores to be used, until we reach the maximum.

To set the number of cores, we change the environment variable OMP_NUM_THREADS to the desired value.

For THIN nodes, which have 24 cores, we start using 1 core, then 2 and then we increase by steps of 2, for a total of 13 points.

For EPYC nodes, which have 128 cores, we start from 1, then 10 and then we increase by steps of 10 until 120. We also use 128 cores, to see what happens at full capacity. We obtain a total of 14 points.

We repeat all measurements 10 times and report the average and standard deviation.

As usual, we repeat this process for both floating and double point precision.

In this section, we have the liberty to explore different thread allocation policies since we are not always using the whole node.

We decided to use following combinations:

- 1. OMP_PLACES=cores and OMP_PROC_BIND=close
- 2. OMP_PLACES=cores and OMP_PROC_BIND=spread

The scripts that were used can be found in the folder exercise2/scripts, under the names es2_2_close_thin.sh, es2_2_close_epyc.sh, es2_2_spread_thin.sh, and es2_2_spread_epyc.sh.

1.3 Results and Discussion

Before we discuss the results of both exercises individually, we briefly introduce the equation to calculate the theoretical peak performance (T_{pp}) of a machine:

$$T_{pp} = \text{Core Count} \times \text{clock freq.} \times \text{IPC}$$
 (1.1)

Where IPC is the instructions per cycle that the architecture is capable of executing.

This equation is very intuitive. The clock frequency tells us how many cycles per second a single core is able to achieve. The ipc factor tells us how many instructions per cycle the core can execute. This number is different for single precision (SP) and double precision (DP) operations. Finally, we need to multiple this by the number of cores that our machine has.

On orfeo, THIN and EPYC nodes are composed of:

- THIN: 24 Intel(R) Xeon(R) Gold 6126 CPU's at 2.60GHz Skylake
- EPYC: 128 EPYC AMD 7H12 CPU's at 2.60GHz Zen 2 (7002 a.k.a "Rome")

Skylake architecture is reported[1] to be able to execute 64 SP FLOP per cycle and 32 DP FLOP per cycle. On the other hand, Zen 2 is reported[1] to execute 32 SP FLOP per cycle and 16 DP FLOP per cycle.

Therefore, we obtain:

Node Type	Total Cores	IPC (SP)	IPC (DP)	T_{pp} (SP)	T_{pp} (DP)
THIN	24	64	32	$\sim 4 \text{ TFLOPS}$	$\sim 2 \text{ TFLOPS}$
EPYC	128	32	16	$\sim 10.6 \text{ TFLOPS}$	$\sim 5.3 \text{ TFLOPS}$

Table 1.1: Performance table of THIN and EPYC node architectures.

Now we can proceed to discuss the results of the exercise.

1.3.1 Using a fixed number of cores

As mentioned above, in this section we keep the number of cores fixed to the maximum available in the node, namely 24 for THIN and 128 for EPYC, and we slowly increase the size of the matrices being multiplied from m = 2000 to m = 20000.

We first show the results for THIN and then for EPYC nodes.

THIN Nodes

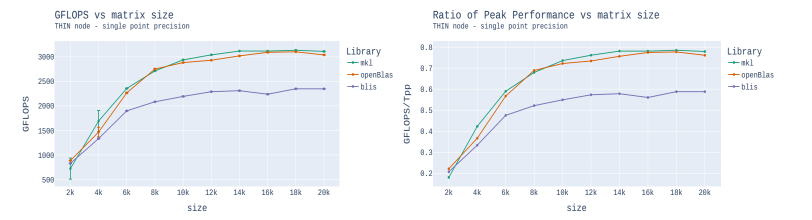


Figure 1.1: Results of SP matrix-matrix multiplication for THIN nodes. MKL and OpenBLAS perform similarly, outperforming BLIS for all matrix sizes.

We see that both MKL and OpenBLAS are able to reach ~ 3.2 TFLOPS, which is around $\sim 80\%$ of T_{pp} . On the other hand, the BLIS library is not able to exploit the full potential of the machine, arriving only to ~ 2.4 TFLOPS, which is $\sim 60\%$ of T_{pp} .

Furthermore, looking at the ratio of peak performance on the right, we observe that for small matrix sizes, none of the libraries are able to fully exploit the theoretical peak performance of the machine. This is most likely because the problem size is so small, that the majority of cores are starving for data rather than crunching numbers. In fact, we are able to reach the best performance when dealing with matrices of size 20000.

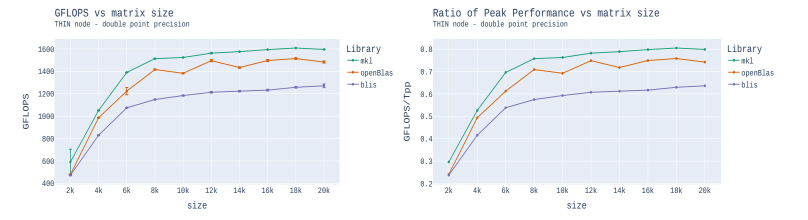


Figure 1.2: Results of DP matrix-matrix multiplication for THIN nodes. MKL performs the best, slightly above OpenBLAS. Both outperform BLIS for all matrix sizes.

We see that MKL reaches ~ 1.6 TFLOPS, while OpenBLAS is slightly lower, at ~ 1.5 TFLOPS, which is $\sim 80\%$ and $\sim 75\%$ of T_{pp} , respectively. On the other hand, BLIS arrives to ~ 1.3 TFLOPS, which is $\sim 65\%$ of T_{pp} .

Furthermore, since double precision is a heavier computation compared to single precision, we see that the libraries perform much better than the their single precision counterparts. For example, looking at the plot for single precision, for matrix size of 4000, we obtain on average around 40% of T_{pp} . On the other hand, for double precision, looking at the same size, we are already at 50% of T_{pp} .

For both SP and DP, and for all matrix size, we notice that MKL and OpenBLAS are better able to exploit the full potential of a THIN node compared to BLIS. Therefore, on THIN nodes, if we need to multiply two matrices, we should always use either MKL or OpenBLAS to get some more performance. To get the absolute best performance, it is preferable to use MKL.

These results shouldn't be surprising, considering that MKL is developed by Intel and THIN nodes are Intel-based. Therefore, it is natural to expect that this library is very fine-tuned to their own architecture and is able to exploit the performance of their machines.

Lastly, we observe the impressive results achieved by OpenBLAS which is based on the original implementation of Kazushige Goto, and is able to achieve a similar performance to MKL, which is maintained by an entire corporation.

EPYC Nodes

Now we show the results on EPYC nodes, which have a T_{pp} of 10.6 TFLOPS for SP and 5.3 TFLOPS for DP.

We first show the results of matrix-matrix multiplication for single point precision.

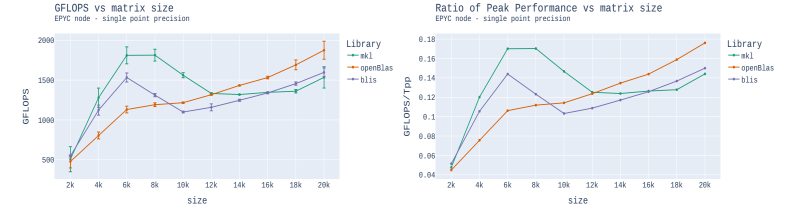


Figure 1.3: Results of SP matrix-matrix multiplication for EPYC nodes. We notice that asymptotically, OpenBLAS outperforms MKL and BLIS, while for small matrices, OpenBLAS performs the worst.

In this case, none of the libraries are able to reach more than 18% of T_{pp} . This could be an indication that to properly exploit a full EPYC node, we need to multiply much bigger matrices.

We also notice that for matrices of size ≤ 9000 , MKL and BLIS outperform OpenBLAS. Between sizes 9000 and 12000, MKL performs best and OpenBLAS begins outperform BLIS. For sizes ≥ 12000 , OpenBLAS performs the best.

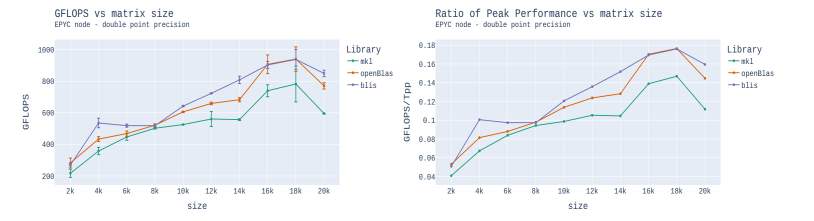


Figure 1.4: Results of DP matrix-matrix multiplication for EPYC nodes. BLIS outperforms MKL and OpenBLAS for all matrix sizes.

Again, we notice that none of the libraries are able to achieve more than 18% of T_{pp} . However, in this case, BLIS outperforms the other two libraries for all matrix sizes. The next best performer is OpenBLAS, followed by MKL, which performs the worst.

In conclusion, on EPYC nodes, it seems that for DP matrix-matrix multiplication, it is better to use BLIS, while for SP, it is very dependent on the size of the matrices. For large matrices, we should use OpenBLAS while for smaller ones, we should use MKL. Furthermore, evidence suggests that to fully exploit and EPYC node, we need to deal with much bigger matrices. So if we are multiplying matrices of size up to 20000, it is much more convenient to just use a THIN node to get more performance.

1.3.2 Using a fixed matrix size

In this section, we fix the matrix size to 10000 and we slowly increase the amount of cores that the libraries can exploit for multithreading through OMP. For THIN nodes, we arrive to 24 cores, while for EPYC nodes, we arrive to 128 cores.

Furthermore, since we are slowly increasing the number of cores that the libraries can use for multithreading, we can study the effects of using different thread allocation policies. In particular, we chose to use OMP_PROC_BIND=close and OMP_PROC_BIND=spread, while always using OMP_PLACES=cores.

In the first case, the threads will slowly occupy first one entire socket, and then, when it is full, the other one. In the second case, the threads will be placed as spread apart as possible, most likely on different sockets. In both cases, when we use the full node, we expect the results to be the same.

Furthermore, in contrast to the previous part of the exercise, we compare the GFLOPS obtained with the T_{pp} calculated with the cores that are being used. In other words, in equation 1.1, instead of using the full 24 or 128 to calculate T_{pp} for the whole node, we use the number of cores we are using for that calculation.

We first analyze THIN and then EPYC nodes.

THIN Nodes

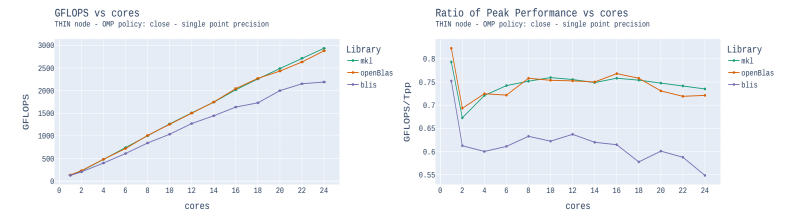


Figure 1.5: Results of SP matrix-matrix multiplication as the number of cores increase, using OMP close policy. MKL and OpenBLAS obtain the best performance.

As we can see from the graph, both MKL and OpenBLAS are able to maintain $\sim 75\%$ of T_{pp} for cores ≥ 6 . On the other hand, BLIS is able to achieve $\sim 60\%$ of T_{pp} . With 24 cores this figure decreases to 55%.

Interestingly, we notice that with 2 cores, there is a significant performance drop. This is most likely explained by the fact that both cores are mapped to the same socket due to the close policy and must share resources such as the higher level caches, causing some contention.

Furtermore, once again, we notice that the best performing library is MKL which is Intel-based, closely followed by OpenBLAS.

Now we analyze the results using a spread policy.



Figure 1.6: Results of SP matrix-matrix multiplication as the number of cores increase, using OMP spread policy. MKL and OpenBLAS are the best perfomers.

Using a spread policy, we obtain very similar results to the case where we use a close policy. The main difference is that we don't have the same performance drop at 2 cores. This is most likely due to the fact that with a spread policy, each core is mapped to its own socket and there is no contention for resources.

In fact, we notice that on average, the performance is slightly better than the close policy counterpart, and this is probably due to better resource usage from the beginning, since threads don't have to compete for resources immediately. This highlights the importance of using the correct mapping policy to obtain better performance.

Now we briefly analyze the results obtained for double precision.



Figure 1.7: Results of DP matrix-matrix multiplication as the number of cores increase, using close policy. MKL and OpenBLAS perform the best.

Similarly to the case of single precision, MKL is able to achieve around $\sim 75\%$ of T_{pp} . However, OpenBLAS suffers from a bit of performance degradation compared to the SP case.

Once again, we notice the immediate drop in performance as soon as we use 2 cores, which is probably caused by the close policy.

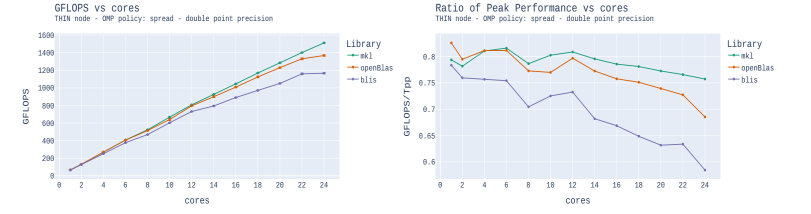


Figure 1.8: Results of DP matrix-matrix multiplication as the number of cores increase, using spread policy.

The analysis and discussion is very similar to the case of single point precision. We no longer see the drop at 2 cores, which is due to better resource usage from the beginning. Compared to the close policy, there is a slightly better performance for this scenario.

As usual, MKL, which is Intel-based, performs the best, closely followed by OpenBLAS, and finally, BLIS, which performs the worst.

Finally, we analyze EPYC nodes.

EPYC Nodes

Considering the results obtained in the first part of the exercise, we expect that MKL won't be the dominating library anymore since we are using an AMD architecture. We also expect some more fluctuation in performance among the libraries, similarly to how there was a dependency on the matrix size.

Furthermore, since we obtained low performance with matrices up to 20000, using all 128 cores, we don't expect an asymptotic improvement. What may happen however, is that when we use less cores, the performance will be much better compared to the theoretical peak performance (per number of cores this time).

We begin by analyzing the SP case with close policy.

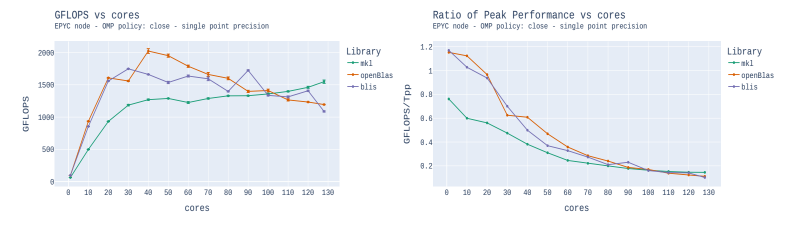


Figure 1.9: Results of SP matrix-matrix multiplication as the number of cores increase, using close policy. The best performance is obtained by OpenBLAS with 40 cores.

Looking at the graph, we see that the best absolute performance is obtained by OpenBLAS at 40 cores, however, this is only $\sim 60\%$ of T_{pp} . On the other hand, for 10-20 cores, the performance almost on part with T_{pp} . This tells us that to fully exploit the machine as intended, with a matrices of size 10000, we only need a handful of cores.

This information is consistent with our previous hypothesis that to fully exploit an EPYC node, we need to consider much larger matrices.

An immediate consequence of this observation, is that as we use more and more cores, the performance deteriorates considerably. When we use the full node, the performance is $\sim 20\%$ of T_{pp} , which is higher than what we obtained for the first part of the exercise.

Contrary to what happens in THIN, we do not notice any severe drops in performance which could be due to the close policy mapping. However, this will be more noticeable once we analyze what happens when we use the spread policy.

Now we analyze the SP case using spread policy.



Figure 1.10: Results of SP matrix-matrix multiplication as the number of cores increase, using spread policy.

There is a big change. The first thing is that now BLIS obtains the best performance, although it suffers from considerable fluctuations. Furthermore, OpenBLAS, which used to perform the best, now performs the worst out of all three libraries.

We also notice that using the spread policy causes both MKL and BLIS to improve and OpenBLAS to worsen. This is interesting because it seems to suggest that the latter library is better able to handle resource contention while the first two are better at fully exploiting resources.

Once again, we notice that as we use more cores, we get worse performance. Now we look at the DP case with close policy.

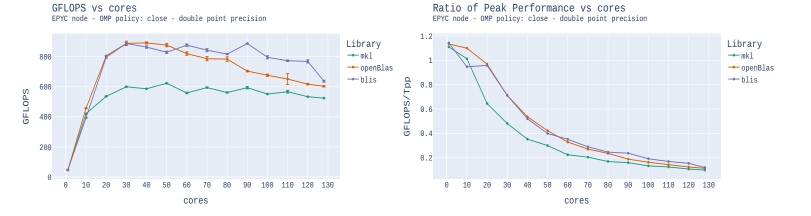


Figure 1.11: Results of DP matrix-matrix multiplication as the number of cores increase, using close policy. The best performance is obtained by BLIS, followed by OpenBLAS.

For double precision, using a close policy, we observe that BLIS performs the best for many cores, while for the first 64 cores, it performs close to OpenBLAS, which is the best. MKL performs the worst.

Finally, we analyze the DP case with spread policy.

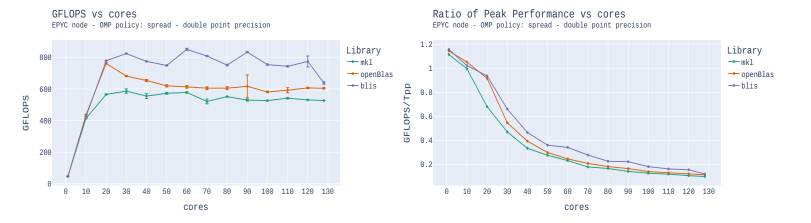


Figure 1.12: Results of DP matrix-matrix multiplication as the number of cores increase, using spread policy.

Using a spread policy makes the gap between BLIS and OpenBLAS much wider. Also, BLIS seems to be more capable at maintaining the throughput as the number of cores are increasing. Of course, this means however, that the performance compared to the theoretical peak is deteriorating.

1.4 Conclusion

After analyzing the results, we can state many conclusions. The first and most obvious one, is that it is better to use Intel-based implementations for Intel machines. This is evidenced by the fact that MKL consistently performed the best in all scenarios for THIN, although it was closely followed by OpenBLAS.

Next, if we are dealing with non-Intel architecures, such as EPYC, it is not advisable to use MKL, as it consistenly underperforms compared to OpenBLAS and BLIS. We also notice that OpenBLAS and BLIS behave completely differently, depending on the mapping policy we use.

We find that for matrices up to 20000, THIN nodes are able to achieve $\sim 75\%$ of T_{pp} , with MKL and OpenBLAS. However, such matrix dimensions are to small to be able to fully exploit an EPYC node which has 128 cores. In fact, in all of our experiments, we are unable to obtain more than 20% of the T_{pp} of the full node.

Finally, as an improvement to this exercise, it would be very interesting to analyze what happens with an EPYC node, using matrices of size 50000 or bigger perhaps, to understand whether we can obtain better performance in this case.

Chapter 2

Conway's Game of Life

2.1 Introduction

This exercise is devoted to implementing a scalable version of Conway's Game of life[2]. The game consists of a $k \times k$ grid, where each cell \mathcal{C} , at position (i, j), can be "alive" or "dead".

The grid evolves over time by looking at each cell's eight nearest neighbors and observing the following rules:

- A dead cell with exactly three live neighbors comes to live (birth).
- A live cell with two or three neighbors stays alive (survival).
- A dead or live cell with less than two or more than three neighbors dies or stays dead (death).

These seemingly simple rules give rise to many interesting behaviors and patterns[3].

There exist two methods to evolve the grid: static and ordered evolution. In static evolution, we freeze the state of the entire grid \mathcal{G}_t at time step t, and compute \mathcal{G}_{t+1} , separately, while looking at \mathcal{G}_t . This corresponds to maintaining two buffers, one for the current generation and the other for the new generation.

On the other hand, in ordered evolution, we start from a specific cell, usually in position (0,0) (top left), and update the elements inplace. This means that the state of each cell depends on the evolved state of some of its neighbors. We call this "ordered" evolution, because the choice of starting point, and the order in which we evolve the grid, lead to different results. In this scenario, the state of each cell instrinsically depends on the history of evolution of *all* the cells before it.

Our implementation must satisfy the following requirements:

- 1. Randomly initialize a square grid ("playground") of size $k \times k$ with $k \ge 100$ and save it as a binary PGM[4] file.
- 2. Load any binary PGM file and evolve for n steps.
- 3. Save a snapshot during the course of evolution with a frequency s, where s = 0 means saving only at the end.
- 4. Support both static and ordered evolution.

Lastly, it must use both MPI[5] and OpenMP[6] (OMP) to paralellize the computations and be able to process grids of considerably high dimensions.

Using our implementation, we will perform a study of the scalability of our code in three cases:

- 1. Strong OMP scalability: Using a grid of fixed dimensions, we fix the number of MPI processes to one per socket and progressively increase the number of OMP threads.
- 2. Strong MPI scalability: Using a grid of fixed dimensions, we increase the number of MPI processes, using only 1 OMP thread per process. We use the largest amount of nodes possible.
- 3. Weak MPI scalability: Changing the grid dimensions to keep the work per process constant, we use one MPI process per socket, with enough OMP threads to saturate the cores in the socket, and slowly increase the number of MPI processes among multiple nodes.

2.2 Methodology

In this section, we briefly describe some of the important choices for the implementation, as well as their implications. We will proceed by layers, tackling the more abstract and higher level problems first and then slowly going into detail about more technical choices.

The first most important step is to think about how to paralellize the problem in order to use MPI and OMP properly.

Since programs in MPI need to be written completely with paralellization in mind from the start, we must begin to conceptualize the problem in an encapsulated manner from the beginning.

At a high abstract level, we must make two important choices:

- 1. How we will decompose the problem.
- 2. How we will perform the IO.
- 3. How to mix MPI and OMP.

How we solve the first problem is of fundamental importance to ensure that our program is scalable and efficient. Also, it has important consequences on the type of paralellism we will need to use. On the other hand, the second problem will determine the organizational paradigm that we will use in our code, as we will see later. Finally, the third solution will determine the effectiveness of our hybrid code.

We briefly discuss both of these topics more in depth in the following subsections.

2.2.1 Decomposition

To exploit paralellism, we must first identify the concurrency in our application and then apply some form of decomposition. The two most important types of decomposition are domain and functional decomposition.

In domain decomposition, multiple workers are performing the same set of instructions on different portions of data (SIMD). In functional decomposition, workers are performing different instructions on possibly different data (MIMD).

In the case of static evolution, each cell can be updated independently from each other, as long as we have access to its neighbors. Therefore, one immediately obviously form of paralellism is for each MPI process to process a "part" of the whole grid. This is an example of domain decomposition, and it is shown below:

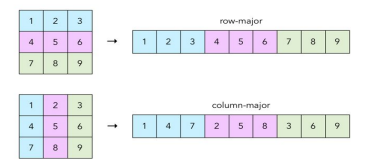


Figure 2.1: Decomposition of an array among three processes. Although the array is conceptualized as a 2D structure, internally, for efficiency, it is represented as a 1D contigous block of memory.

However, we need to decide how to split the grid. We can do a 1D decomposition by "stripes", as shown in image 2.1 or a 2D decomposition by "blocks". It is known that a 2D decomposition is more efficient as it can exploit more bandwidth as more workers will send shorter messages contemporarily. However, it is more complicated both from an implementation point of view and worse for memory access.

Let's talk briefly about this second aspect. Although we talk about the grid as a 2D array, internally, for efficiency, it will be represented as a 1D array.

If we do a 1D split, each process will work on a thick strip of data which continuous in memory. On the other hand, if we do a 2D split, each process will work on a block of memory which is not contiguous and located every certain "jumps" in the 1D array. As a consequence, as each process traverses its block of the grid, it will have more cache misses compared to the 1D case where we perform a simple a linear access. This is more easily visualized in the following image:

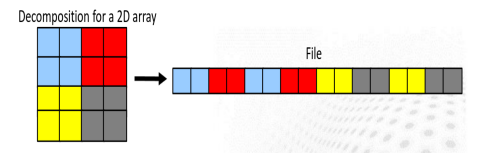


Figure 2.2: 2D decomposition of a grid. As we can see, each process needs to deal with non-contiguous memory.

Therefore, for reasons stated above, and for simplicity and readability, we decided to use the 1D splitting.

Lastly, we must consider how to handle halo regions. Since each process will process a strip composed of 1 or more rows - of the grid, the rows at the boundary of the strip must be handled with special attention. These rows - 2 per process - need information that is contained in the grid of other processes. Therefore, we need to allocate some extra space for these additional rows, and set up message passing between processes to handle the exchange of these boundary rows, also known as halo regions.

Below we show the idea:

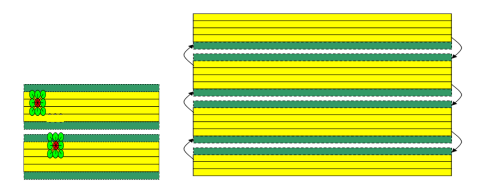


Figure 2.3: On the figure in the left, we can see that we can process cells that are internal with no problem. However, at the boundary we need information that belongs to another process. Therefore, we allocate some extra space and do a halo exchange as shown in the figure on the right.

Finally, in the case of **ordered** evolution, we cannot update each cell independently, since we modify the elements inplace in an ordered fashion. Therefore, we cannot parallelize this process. The only thing that we can do, is use MPI to be able to process much bigger grids than we could in a conventional computer. In this case, each process will compute its part of the grid *in order*, rather than in parallel. So process 0 will update its portion, and send the updated row to process 1, and so on. Of course, each process needs the appropriate halo regions.

2.2.2 MPI IO

The problem specifications require that we must save the grid with a certain frequency s. There are multiple ways to achieve this, and they all have important implications on the scalability of the code.

The simplest way is for each process to write its part of the grid to a separate file. This however, is extremely messy as we loose the flexibility of using a different number of processes between separate runs. More importantly, for a large number of processes, the filesystem will not be able to handle all these requests and will be completely overloaded and possibly crash. This means that for P >> 1 we would have very serious problems, which is undesirable, since we wish to use as many processes as we want/have.

The second solution, is to use a master-slave approach. In this paradigm, one process is designated as the master process, while all the other are its slaves. The master will coordinate the IO, and with a frequency s, it will collect information from all the slaves, and take care of writing everything to a file. This approach, although very common, is not scalable, as each process needs to send its part of the grid to the master process. So for P >> 1, the bottleneck will be in the communication.

Lastly, the third, and best solution, is to use MPI IO, which was meant to handle exactly this problem. The idea behind MPI IO is very similar to message passing. Each process will call a collective function to read/write its part of the grid and the MPI implementation will take care of exploiting the parallel filesystem and reducing the amount of read and write operations. Furthermore, this is the only solution that takes advantage of the parallel filesystem which allows for a unified logical view of the file, while allowing the file to be stored in physically different locations. This is very advantageous for scalability since many processes can contemporarily write in different physical locations, which increases the throughput of the program. We note that this is a form of functional decomposition.

Notice that this aspect is independent of the type of evolution.

2.2.3 Mixing MPI and OMP

Now that we have discussed at an abstract level how to distribute the workload among the MPI processes, we need to think about how to integrate OMP to obtain a hybrid implementation.

In static evolution, since each process can update its part of the grid independently, the most natural approach is to use OMP to parallelize this computation. For this, there are a few options.

The simplest option, is that each MPI process will spawn a parallel region at each iteration t to accelerate its computations.

Another, more advanced approach, is to use MPI_THREAD_MULTIPLE. In this mode, each OMP thread can make an MPI call contemporarily. This mode must be handled with care but it will enable us to create a single parallel region, as we will see later.

We implemented both cases to do some comparisons.

Finally, for ordered evolution, we cannot use OMP, because the update depends on the order, and it is intrinsically serial.

2.3 Implementation

In this section, we discuss the most technical details of our implementation, from the datatypes used, to compiler keywords, to optimization of branches, to memory allocation, and optimization of for loops. We will also describe how to use the code, what software stack is needed, and what tools were used.

Our implementation was done in C, so we will have to manually handle all memory aspects. This gives us great control in to what we are doing.

2.3.1 State representation

We begin by discussing how to represent data. Since each cell can be alive (1) or dead (0), theoretically, we need a single bit to represent the state. This means that every byte of data can represent the state of 8 cells. However, this approach is complex since we need to do a lot of bit manipulation to access the information of each cell. This will also make the code less readable and longer.

The next best idea, which we used, is to use the smallest datatype possible, i.e a char, which is 1 byte, to represent the state of a single cell. In this way, we are less memory efficient, but we gain in readability and ease of implementation. In particular, we use an unsigned char which is of the same

size, however, in this way, the compiler can possibly optimize the operations since it doesn't need to keep track of the sign.

All of our implementations contain the following macros:

```
#define DEAD 0
#define ALIVE 1
```

2.3.2 Distributing the grid among MPI processes

Given a $k \times k$ grid, we need to tell each MPI process how many rows it must deal with. To do this we can divide the total number of rows by the number of processes, i.e the size of the MPI Communicator, and distribute the remainder among a subset of them.

This is done through a common trick:

```
my_rows = total_rows/size + 1*(rank<(total_rows%size));</pre>
```

In this way, all processes will have an equal workload, except for some ranks which will have to process one more row.

2.3.3 Allocating data

Now that we have determined how many rows each process has to deal with, each rank must allocate the appropriate memory in heap to store its portion of the grid.

In static evolution, we need to keep two buffers, one for the current generation, and one for the next one. We look at the previous generation as we update the new generation. In ordered evolution, we only have one buffer, and we update each cell inplace as we traverse the array.

As mentioned previously, it is not efficient to allocate a 2D array, although it is more naturally aligned with the problem statement. This is because 2D arrays are not necessarily contiguous in memory and we also have an overhead of following pointers.

Therefore, we allocate 1D arrays which are contiguous in memory and we use a macro to be able to access the array as if it were a 2D structure.

Lastly, for each process, we need to allocate two additional rows to accommodate the halo rows. Also, to optimize for cache access, we explicitly align the data to the cache line size of THIN and EPYC nodes, which are 64 Bytes.

This can be seen in the following snippet:

```
// defining cache line size
#define CACHE_LINE_SIZE 64
// macro to access data in 2D fashion
#define DATA(i,j) (data[(i)*cols + (j)])
...
// aligned allocation
data = (unsigned char *) aligned_alloc(CACHE_LINE_SIZE, (my_rows + 2) * cols);
```

Of course, for static evolution, we perform two such allocations.

2.3.4 PGM Files - Header

Now that we have discusses how we perform the allocation, we briefly discuss PGM binary files.

PGM, or Portable Grey Map, are portable binary files composed of a header and of binary data. In our case, the header is the following line:

```
P5 {rows} {cols} {maxval}\n
```

The P5 is a magic number that states we are using a PGM file of binary data where each byte corresponds to a pixel in the image. The color is interpreted according to the maxval value. Without going into too much detail, by setting maxval to one, we are establishing that 1's will represent be drawn as white (alive) and 0's will be drawn as black (dead).

Whenever we save a snapshot of the evolution, we need to write this header. This will be done by one single process - rank 0.

Also, as we will see later, to use MPI IO, we need to find out the length of this header in bytes. All these operations are performed in the following snippet:

```
#define HEADER_FORMAT_STRING "P5 %lf %lf %d\n"
#define MAX_VAL 1

...

// We get the header size
header_size = snprintf(NULL, 0, HEADER_FORMAT_STRING, rows, cols, MAX_VAL);

// allocate the string
header = malloc(header_size + 1)

// write the header into the string
sprintf(header, HEADER_FORMAT_STRING, rows, cols, MAX_VAL);
```

We will use the header size in MPI IO, in the next section.

2.3.5 MPI IO

As discussed above, MPI IO is the best choice to handle the IO when we are dealing with many processes. It takes advantage of the parallel file system to write in multiple locations at once, while keeping a logically unified view of the file. In this way, we avoid the bottleneck of having to send the grid back to the master or overloading the filesystem with too many posix calls.

Below we list the code snippet that we used and then discuss the relevant details.

```
void save_grid(char * restrict fname, MPI_Comm comm, int rank, char * restrict header,
       unsigned long int header_size, MPI_Offset offset, unsigned char * restrict data,
      unsigned long int my_rows, unsigned long int cols)
3
  {
5
      MPI_File fh;
6
      // Opening the file in MPI_MODE_WRITEONLY or MPI_MODE_CREATE_ONLY
      const int err = MPI_File_open(comm, fname, MPI_MODE_WRONLY | MPI_MODE_CREATE,
      MPI_INFO_NULL, &fh);
9
      // Check that the file was opened correctly
10
      if(err != MPI_SUCCESS)
11
           fprintf(stderr, "Error opening %s\n", fname);
13
          MPI_Abort(MPI_COMM_WORLD, err);
14
16
      if(rank == 0)
18
           // we specify the file handle, the byte where we write, what we write,
19
          // the length of what we write, the data type of what we write, and the
20
           // status of the writing.
21
          MPI_File_write_at(fh, 0, header, header_size, MPI_CHAR, MPI_STATUS_IGNORE);
22
23
24
      MPI_File_write_at_all(fh, offset, data + cols, my_rows*cols, MPI_CHAR,
25
      MPI_STATUS_IGNORE);
26
      MPI_File_close(&fh);
27
28 }
```

First, all processes call the MPI_File_open() routine to open the file. The file is opened in MPI_MODE_WRONLY for write only and MPI_MODE_CREATE to create the file if it does not exist already.

Then, we choose rank 0 to write the header of the PGM file. We use the MPI function MPI_File_write_at() which allows us to specify at what position in the file, in bytes, the data must be written.

For the header, we will write at the beginning of the file, so at byte 0. The rest of the arguments are explained in the snippet.

Finally, for writing the grid, we will use the collective version of MPI_File_write_at(), namely MPI_File_write_at_all(). This function is called by all the processes. Internally, the MPI implementation will perform this operation in the most efficient way possible.

The arguments are the same as the non-collective version.

The only thing we need to explain, is how we find the byte offset for each process.

We first find the row offset, i.e. which row to start writing from, through the following function:

```
unsigned long int get_my_row_offset(unsigned long int total_rows, int rank, int size)
      // returns the offset from which the rank must get its rows
3
      // for example: for rank 0, the offset will be zero as it will read
5
      // my_rows(of rank 0) from the start. However, rank 1 will have to read
6
      // after the rows that rank O handles (accounting for the remaineder),
      // and so on.
9
      // nrows is the interger division between total_rows and size
      unsigned long int nrows = total_rows/size;
      unsigned long int remainder = total_rows%size;
12
13
      // The idea is the following, if the division was pefect, we would do
14
      // nrows st rank. However, we still need to account for the remainder.
1.5
      // For remainder processors (starting to count from rank 1 because rank 0
16
      // just gets 0 as offset), we simply need to add rank to the offset.
17
      // If we are dealing with a process id which is greater than the remainder,
18
      // then we simply need to add the whole remainder.
19
20
      unsigned long int my_offset = nrows*rank
21
          + rank*(((unsigned long int) rank) <= remainder)
22
          + remainder*(((unsigned long int) rank) > remainder);
23
24
      return my_offset;
25
26 }
```

We illustrate with a brief example. Suppose we have a 14×14 grid and four ranks. The interger division is 3 and the remainder is 2. Since the remainder of the division is 2, two ranks, 0 and 1, will need to handle an extra row.

So rank 0 will deal with 3 + 1 = 4 rows, rank 1 will also deal with 3 + 1 = 4 rows, and ranks 2 and 3 will deal with 3 rows each.

Now, for the offset, rank 0 will just have offset 0. Rank 1 will need to shift by the interger division plus 1 to account for the extra row that rank 0 has. Similarly, rank 2 has to shift by twice the interger division, plus the extra row that rank 0 has and the extra row of rank 1.

Rank 3 will have to shift by three times the interget division, plus the extra rows of the previous ranks. However, this time, rank 2 did not have an extra row. So rank 3 just needs to account for *total* extra rows, which is equal to the remainder.

This sounds complicated, but it is simple once one works through a few examples.

Since the offset for the MPI functions is in bytes, to get the final offset, we first multiply the row offset by the number of columns, and by the size of the datatype we use in bytes - for a char this is 1.

Then we need to add the size in bytes of the header.

```
const MPI_Offset header_offset = header_size * sizeof(char);

const MPI_Offset my_file_offset = my_row_offset * cols * sizeof(char);

const MPI_Offset my_total_file_offset = my_file_offset + header_offset;

...
```

Finally, reading of the file is done in the same exact way, replacing MPI_File_write_at_all with MPI_File_read_at_all.

However, we need to have an additional step to read the header, and this is done through a normal fscanf() call.

2.3.6 Updating Cells - Static Evolution

Now we discuss how we perform the cell update for static evolution.

The entire evolution process is contained inside a for loop over time steps. At each iteration, we had to use message passing to exchange the halo rows. This is done using non blocking operations to hide the latency of the communication.

```
1 // we post send and recv requests for two halo regions.
2 // matrix:
                           row 0 (HALO)
3 // -
4 // 010100011001100100100 row 1
5 // 010010000100101001111 row 2
                                              | these rows dont need
6 // ...
_7 // 001010111110011000001 row my_rows - 1 \, __| halo regions
8 // 010010010000100101000 row my_rows
9 // -----
                           row my_rows + 1 (HALO)
10
11 // send row: 1 to the previous rank
12 MPI_Isend(data_prev + cols, cols, MPI_CHAR, prev, prev_tag, MPI_COMM_WORLD, &
      prev_send_request);
// send row: my_rows to next rank
MPI_Isend(data_prev + my_rows_x_cols, cols, MPI_CHAR, next, next_tag, MPI_COMM_WORLD,
      &next_send_request);
15
16 // receive from prev and put in row 0 (halo region)
17 MPI_Irecv(data_prev, cols, MPI_CHAR, prev, next_tag, MPI_COMM_WORLD, &
     prev_recv_request);
18 // receive from next and put in row my_rows + 1 (halo region)
MPI_Irecv(data_prev + my_rows_x_cols_p_cols, cols, MPI_CHAR, next, prev_tag,
      MPI_COMM_WORLD, &next_recv_request);
```

Where data_prev and data are the buffers for the previous and next generation, respectively.

Next, while messages are exchanged, each process can process the internal portion of its grid.

Two versions were used to do this: one using a double for loop over rows and columns, and the other using a single for loop over cells. Later, when we tie everything together, we will explain the reason for these two versions.

```
2
3 // note we start from row 2 and finish at row my_rows - 1
4 for(unsigned long int row = 2; row < my_rows; ++row)</pre>
      for(unsigned long int col = 0; col < cols; ++col)</pre>
6
7
           upgrade_cell_static(data_prev, data, row, col);
8
9
10 }
11
12 ...
for(unsigned long int cell = 2*cols; cell< my_rows*cols; ++cell)</pre>
2 {
          unsigned long int i = cell/cols;
          unsigned long int j = cell - i*cols;
4
5
          upgrade_cell_static(data_prev, data, i, j);
6 }
```

Finally, at this point, there is nothing left to do, but wait until we receive the halo cells.

```
MPI_Wait(&prev_recv_request, MPI_STATUS_IGNORE);
MPI_Wait(&next_recv_request, MPI_STATUS_IGNORE);
MPI_Request_free(&prev_send_request);
MPI_Request_free(&next_send_request);
```

We then process the first and last row of the grid (internal ones).

```
for(unsigned long int col=0; col < cols; ++col)
{
    upgrade_cell_static(data_prev, data, 1, col);
}

// we only do this if we have more than one row
for(unsigned long int col=0; col < cols * (my_rows > 1); ++col)

upgrade_cell_static(data_prev, data, my_rows, col);
}
```

Then, if we need to save the grid we save using MPI IO, and then we swap the pointers and repeat. Apart from a few differences that we introduce with OMP later, these are the fundamental steps of our evolution.

The last thing is to discuss the update function itself:

```
void upgrade_cell_static(unsigned char * restrict data_prev, unsigned char * restrict
      data, unsigned long int i, unsigned long int j)
2
      DATA(i,j) = DEAD;
3
      // we use ternary operators to avoid branches
5
      // as well as the register keyword to signal to the compiler to
6
      // keep this data close
      // compute coordinates of neighbors with wrap around of dimensions
9
      register unsigned long int jm1 = j==0 ? cols-1 : j-1;
10
      register unsigned long int jp1 = j == (cols-1) ? 0 : j+1;
11
      // for rows, wrapping is ensured by the presence of halos
12
      register unsigned long int im1 = i-1;
13
      register unsigned long int ip1 = i+1;
14
      // we fetch the neighbors with 8 different instructions to saturate pipelines.
16
      // We access first the neighbors that are already most likely already
17
      // in cache due to previous updates.
18
19
      unsigned char tmp0=DATA_PREV(im1, jm1);
      unsigned char tmp3=DATA_PREV(i,jm1);
20
      unsigned char tmp5=DATA_PREV(ip1,jm1);
21
22
      unsigned char tmp1=DATA_PREV(im1, j);
23
      unsigned char tmp2=DATA_PREV(im1,jp1);
24
25
      unsigned char tmp4=DATA_PREV(i,jp1);
26
27
      unsigned char tmp6=DATA_PREV(ip1,j);
28
      unsigned char tmp7=DATA_PREV(ip1,jp1);
29
30
      unsigned char n_alive_cells = tmp0+tmp1+tmp2+tmp3+tmp4+tmp5+tmp6+tmp7;
31
32
33
      DATA(i,j) = ALIVE*(n_alive_cells == 3) + DATA_PREV(i,j)*(n_alive_cells == 2);
34 }
```

The function is very simple. First, we set the cell to be dead regardless, since if there are less than two and more than 3 live neighbors, it dies.

Then, we calculate the coordinates of the neighbors using ternary operators to ensure that we wrap around the dimensions.

Then, we use separate instructions for each neighbor to encourage pipeline saturation.

We aggregate the state of all the neighbors, and apply the rules. This version of the rules may seem off at first, but it is completely equivalent to the one we described, although in this way, it's a bit more efficient.

Finally, notice the **restrict** keyword for the two pointers so that the compiler can optimize better since it knows it will never encounter memory aliasing.

2.3.7 Updating Cells - Ordered Evolution

For ordered evolution, the idea is more complex.

We need to make sure that ranks process their parts of the grid in an ordered fashion. We first show the commented snippet and then comment:

```
// for ordered evolution, we cannot parallelize, and each process
// must work on their part of the grid in serial order.

// first, each process posts a non blocking receive operation for
// the top halo row and the bottom halo row.

// Since we have ordered evolution each process will then wait
// until the top halo row has been received to start working on
// its grid.

MPI_Irecv(data, cols, MPI_CHAR, prev, next_tag, MPI_COMM_WORLD, &prev_recv_request);
```

```
10 MPI_Irecv(data + my_rows_x_cols_p_cols, cols, MPI_CHAR, next, prev_tag, MPI_COMM_WORLD
      , &next_recv_request);
12 // however, each process will also have to send its top row
_{
m 13} // (the bottom halo row for the previous process) which is necessary
_{14} // for the previous process to calculate the grid.
_{15} // This excludes rank 0, because it needs to send its first
16 // row to rank size-1 row only AFTER it has been updated
if (rank != 0) { MPI_Isend(data + cols, cols, MPI_CHAR, prev, prev_tag, MPI_COMM_WORLD,
      &prev_send_request);}
18
19 // to get things started for rank 0, this is the first forward
20 // message of the top halo row.
if(rank == size-1) { MPI_Isend(data + my_rows_x_cols, cols, MPI_CHAR, next, next_tag,
      MPI_COMM_WORLD, &next_send_request); }
23 // All processes will wait until receive of top halo row is complete
24 // (since they need it to start ordered evolution)
25 //
_{26} // the first time, only process 0 should pass.
MPI_Wait(&prev_recv_request, MPI_STATUS_IGNORE);
29 // Since we have received the top halo row, we can process all
_{
m 30} // data EXCEPT the last row, since we are not sure that we have
^{31} // received the bottom halo yet.
32 for(unsigned long int row = 1; row < my_rows; ++row)</pre>
33 {
      for(unsigned long int col = 0; col < cols; ++col)</pre>
34
35
           upgrade_cell_ordered(data, row, col);
36
37
38 }
39 // Based on how many rows we have, we behave differently
40 if (my_rows >1)
41 {
       // in this case, the for loop above has been executed and
42
       // now that the first row has been processed, rank 0 can send it
43
44
      // to rank size-1
       if(rank == 0){ MPI_Isend(data + cols, cols, MPI_CHAR, prev, prev_tag,
45
      MPI_COMM_WORLD, &prev_send_request);}
46
      // We have finished processing all our grid except for last row.
47
      // For this, we need to make sure that bottom halo has been received.
48
49
       // wait for receive of bottom halo
50
51
      MPI_Wait(&next_recv_request, MPI_STATUS_IGNORE);
52
53
      // process last row
      for(unsigned long int col=0; col<cols; ++col)</pre>
54
55
56
           upgrade_cell_ordered(data, my_rows, col);
57
58
       // now that we have computed the last row, we need to send it
59
60
       // to the next process. This will be the top halo for the next
       // process, which in turn will finally pass the wait statement
61
      // above.
62
63
       // all except process n-1 because this will be taken care of in the
64
       // next iteration of the for loop
65
      if(rank<(size-1)){ MPI_Isend(data + my_rows_x_cols, cols, MPI_CHAR, next, next_tag</pre>
66
       , MPI_COMM_WORLD, &next_send_request);}
67
68 }
69 else
70 {
71
       // in this case we have skipped the for loop
       // we need to wait to receive the bottom halo from next
72
73
      MPI_Wait(&next_recv_request, MPI_STATUS_IGNORE);
74
// we process row 1
```

```
for(unsigned long int col=0; col<cols; ++col)</pre>
76
77
           upgrade_cell_ordered(data, my_rows, col);
78
79
       // rank zero sends this row to rank n-1
81
      if(rank == 0){ MPI_Isend(data + cols, cols, MPI_CHAR, prev, prev_tag,
82
      MPI_COMM_WORLD, &prev_send_request);}
83
       // all ranks except last, send their last row (the same one )
84
       // to the next rank
85
86
       if(rank<(size-1)){ MPI_Isend(data + my_rows_x_cols, cols, MPI_CHAR, next, next_tag</pre>
       , MPI_COMM_WORLD, &next_send_request);}
87 }
88
89 // we put a barrier since all processes need to wait before until the whole grid
90 // is processed before checking if they need to save it
91 MPI_Barrier(MPI_COMM_WORLD);
93 // Free the send operations
94 MPI_Request_free(&prev_send_request);
95 MPI_Request_free(&next_send_request);
```

The snippet is commented very thoroughly, however, we briefly explain the idea. To process its part of the grid, each rank needs to wait for the previous rank to process its last row, and send it over to be placed in top halo row.

Hence, by placing a wait request for the top halo row, we ensure that the ranks process the cell in correct order, one at a time.

Furthermore, each rank also needs the last halo row to process its last internal row. We can send this in the beginning with a non-blocking operation.

Finally, to get things started, we rank size - 1 sends its last row to rank 0, which will be the first one to receive a top halo row, and pass through the receive operation.

From there, we simply update rows 1 to the second to last, wait until we receive the bottom halo, process the last row, and then send it to the next rank so it can pass the wait request.

There is a small consideration to make in the case we have only one row, but conceptually, it is the same.

Lastly, we show the update function:

```
void upgrade_cell_ordered(unsigned char * data, unsigned long int i, unsigned long int
2 {
      register unsigned long int jm1 = j == 0 ? cols-1 : j-1;
3
      register unsigned long int jp1 = j == (cols-1) ? 0 : j+1;
      register unsigned long int im1 = i-1;
      register unsigned long int ip1 = i+1;
      unsigned char tmp0=DATA(im1, jm1);
      unsigned char tmp3=DATA(i,jm1);
9
      unsigned char tmp5=DATA(ip1, jm1);
10
      unsigned char tmp1=DATA(im1,j);
12
13
      unsigned char tmp2=DATA(im1,jp1);
14
      unsigned char tmp4=DATA(i,jp1);
15
16
      unsigned char tmp6=DATA(ip1,j);
17
      unsigned char tmp7=DATA(ip1,jp1);
18
19
20
      register unsigned char n_alive_cells = tmp0+tmp1+tmp2+tmp3+tmp4+tmp5+tmp6+tmp7;
21
      DATA(i,j) = ALIVE*(n_alive_cells==3) + DATA(i,j)*(n_alive_cells==2);
22
23 }
```

The function is virtually the same, with the only difference that we operate on a single buffer.

2.3.8 OMP Integration - Hybrid V1

In this section, we tie everything together to show the final implementation. As mentioned before, we have two implementations for static evolution.

Here we comment on the first one.

For the first implementation, we create a single parallel region outside of the for loop over the evolution time steps. Furthermore, we use MPI Multiple mode, where each thread can make an MPI call contemporarily:

The idea behind this, is to avoid the creation of a parallel region every time we iterate over the time steps. To do this, we need to initialize MPI slightly differently, as shown below:

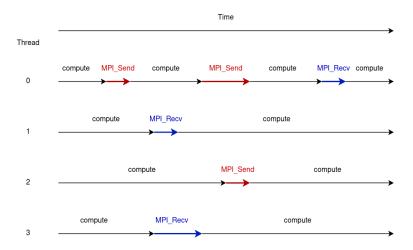


Figure 2.4: Visualization of MPI Multiple mode. Each OMP thread can make MPI calls.

```
int provided_thread_level;
MPI_Init_thread(&argc, &argv, MPI_THREAD_MULTIPLE, &provided_thread_level);
if(provided_thread_level < MPI_THREAD_MULTIPLE)

{
    printf("Can't do thread multiple mode... Aborting");
    MPI_Finalize();
}</pre>
```

Then, the for loop over the evolution time steps looks like this:

```
#pragma omp parallel
  for(unsigned long int t = 1; t < n+1; ++t)</pre>
3
      // no barrier at the end
5
      #pragma omp single nowait
6
      MPI_Irecv(data_prev, cols, MPI_CHAR, prev, next_tag, MPI_COMM_WORLD, &
      prev_recv_request);
      #pragma omp single nowait
      MPI_Irecv(data_prev + my_rows_x_cols_p_cols, cols, MPI_CHAR, next, prev_tag,
      MPI_COMM_WORLD, &next_recv_request);
      #pragma omp single nowait
      MPI_Isend(data_prev + cols, cols, MPI_CHAR, prev, prev_tag, MPI_COMM_WORLD, &
      prev_send_request);
12
      #pragma omp single nowait
      MPI_Isend(data_prev + my_rows_x_cols, cols, MPI_CHAR, next, next_tag,
13
      MPI_COMM_WORLD, &next_send_request);
14
      // remaining threads in the meantime start to work on internal part
      // of grid. We need to keep the implied barrier because waiting can
16
      // only happen if operations have been posted
17
      #pragma omp for schedule(dynamic, chunk)
18
      for(unsigned long int cell = 2*cols; cell < my_rows_x_cols; ++cell)
19
      {
20
21
               unsigned long int i = cell/cols;
              unsigned long int j = cell - i*cols;
```

```
upgrade_cell_static(data_prev, data, i, j);
23
24
      }//barrier. Needed, otherwise, if threads havent finished posting
        //\mathrm{send} and receive, the next operation will be an error.
25
26
27
      #pragma omp single
28
           MPI_Wait(&prev_recv_request, MPI_STATUS_IGNORE);
29
           MPI_Wait(&next_recv_request, MPI_STATUS_IGNORE);
30
           MPI_Request_free(&prev_send_request);
31
           MPI_Request_free(&next_send_request);
32
      } //barrier needed because we need to get halo cells to process
33
34
         //next rows
35
       // threads can split a row further
36
37
      #pragma omp for schedule(dynamic, small_chunk) nowait
      for(unsigned long int col=0; col<cols; ++col)</pre>
38
39
           upgrade_cell_static(data_prev, data, 1, col);
40
41
42
       // once done with row above, threads will start immediately on this
43
      #pragma omp for schedule(dynamic, small_chunk)
44
      for(unsigned long int col=0; col<cols*(my_rows>1); ++col)
45
46
           upgrade_cell_static(data_prev, data, my_rows, col);
47
      }// barrier since I need to wait for all of them to finish before
48
49
       // saving and swapping
50
      // implied barrier at the end to finish saving and swapping
51
      #pragma omp single
52
53
54
           ++save_counter;
           if(s>0 && save_counter == s && t<100000)</pre>
55
               sprintf(snapshot_name, "snapshot_%051d", t);
57
               save_grid(snapshot_name, MPI_COMM_WORLD, rank, header, header_size,
      my_total_file_offset, data, my_rows, cols);
               save_counter = 0;
59
          }
60
61
           tmp_data = data;
           data = data_prev;
63
           data_prev = tmp_data;
64
65
           tmp_data = NULL;
      }// barrier
66
67 } // barrier
```

- 2.3.9 OMP Integration Hybrid V2
- 2.3.10 Software Stack
- 2.3.11 Running the Code
- 2.4 Bonus Algorithm 2
- 2.5 Results and Discussion
- 2.5.1 Strong OMP Scalability
- 2.5.2 Strong MPI Scalability
- 2.5.3 Weak MPI Scalability
- 2.6 Conclusions

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