

DELFT UNIVERSITY OF TECHNOLOGY

INTRODUCTION TO HIGH PERFORMANCE COMPUTING  
WI4049TU

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# Lab Report

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## General Remarks

This final Lab report includes the answers for the exercises (base grad denoted in paranthesis):

0. Introductory exercise (0.5)
1. Poisson solver (1.75)
2. Finite elements simulation (1.0)
3. Eigenvalue solution by Power Method on GPU (1.75)

The optional **shining points** (e.g., performance analysis, optimization, discussion, and clarifying figures) which yield further points are usually marked by a small colored heading in the text or an additional note is added under a figure or table. For example:

**This is a shining point.**

**N.B.** The whole repository is available on [GitHub](#).

Furthermore, if not stated otherwise the code is run on the **Delft Blue** supercomputer. Due to excessive load on the system some tests (additionally denoted) and especially the development of the code was done on the local machine.

## 0 Introductory exercise

In the introductory lab session, we are taking a look at some basic features of MPI.

We start out very simple with a hello world program on two nodes.

### Hello World

```
1 #include "mpi.h"
2 #include <stdio.h>
3
4 int np, rank;
5
6 int main(int argc, char **argv)
7 {
8     MPI_Init(&argc, &argv);
9     MPI_Comm_size(MPI_COMM_WORLD, &np);
10    MPI_Comm_rank(MPI_COMM_WORLD, &rank);
11
12    printf("Node %d of %d says: Hello world!\n", rank, np);
13
14    MPI_Finalize();
15    return 0;
16 }
```

This program can be compiled with the following command:

```
mpicc -o helloworld1.out helloworld1.c
```

And run with:

```
srunk -n 2 -c 4 --mem-per-cpu=1GB ./helloworld1.out
```

We get the following output:

```
Node 0 of 2 says: Hello world!
```

```
Node 1 of 2 says: Hello world!
```

From now on I'll skip the compilation and only mention on how many nodes the program is run and what the output is / interpretation of the output.

## 0.a) Ping Pong

I used the template to check how long `MPI_Send` and `MPI_Recv` take. The code can be found in the appendix for this section.

I've modified the printing a bit to make it easier to gather the information. Then I piped the program output into a textfile for further processing in python. I ran it first on one and then on two nodes as specified in the assignment sheet. Opposed to the averaging over 5 send / receive pairs, I've done 1000 pairs. Furthermore I reran the whole program 5 times to gather more data. All this data is shown in the following graph:

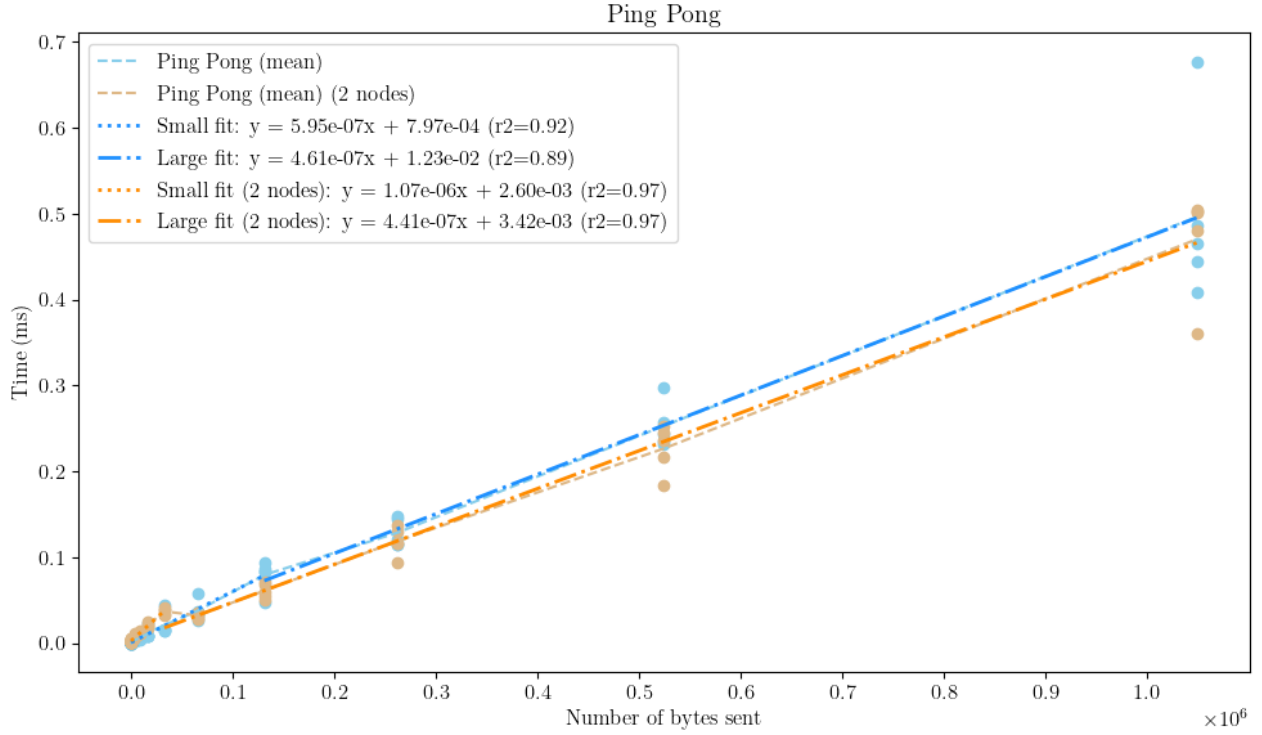


Figure 1: Ping Pong: Number of bytes sent vs. average time taken from 1000 pairs of send / receive. 5 runs shown for each size as scatter plot. Mean of these 5 runs shown as line. Blue small fit includes all data points up to 131072 bytes, blue large from there. Red small fit includes all data points up to 32768 bytes, red large from there.

As can be seen in the data and the fits, there are outliers especially for the larger data sizes. For our runs we get the following fits and  $R^2$  values:

Run Type	Data Size	Fit Equation	$R^2$ Value
Single Node	Small ( $\leq 131072$ )	$5.95 \times 10^{-7} \cdot x + 7.97 \times 10^{-4}$	0.92
Single Node	Large ( $\geq 131072$ )	$4.61 \times 10^{-7} \cdot x + 1.23 \times 10^{-2}$	0.89
Two Node	Small ( $\leq 32768$ )	$1.07 \times 10^{-6} \cdot x + 2.60 \times 10^{-3}$	0.97
Two Node	Large ( $\geq 32768$ )	$4.41 \times 10^{-7} \cdot x + 3.42 \times 10^{-3}$	0.97

Table 1: Fit Equations and  $R^2$  Values for Single Node and Two Node Runs

**Note:** Each run was performed 5 times (for 1 and 2 nodes) to get a fit on the data and calculate a  $R^2$  value.  
**TODO: Further analysis needed?**

### Extra: Ping Pong with `MPI_SendRecv`

We do the same analysis for the changed program utilizing `MPI_SendRecv`. The code can be found in the appendix for this section.

We get the following graph from the measurements which were performed in the same way as for the previous program:

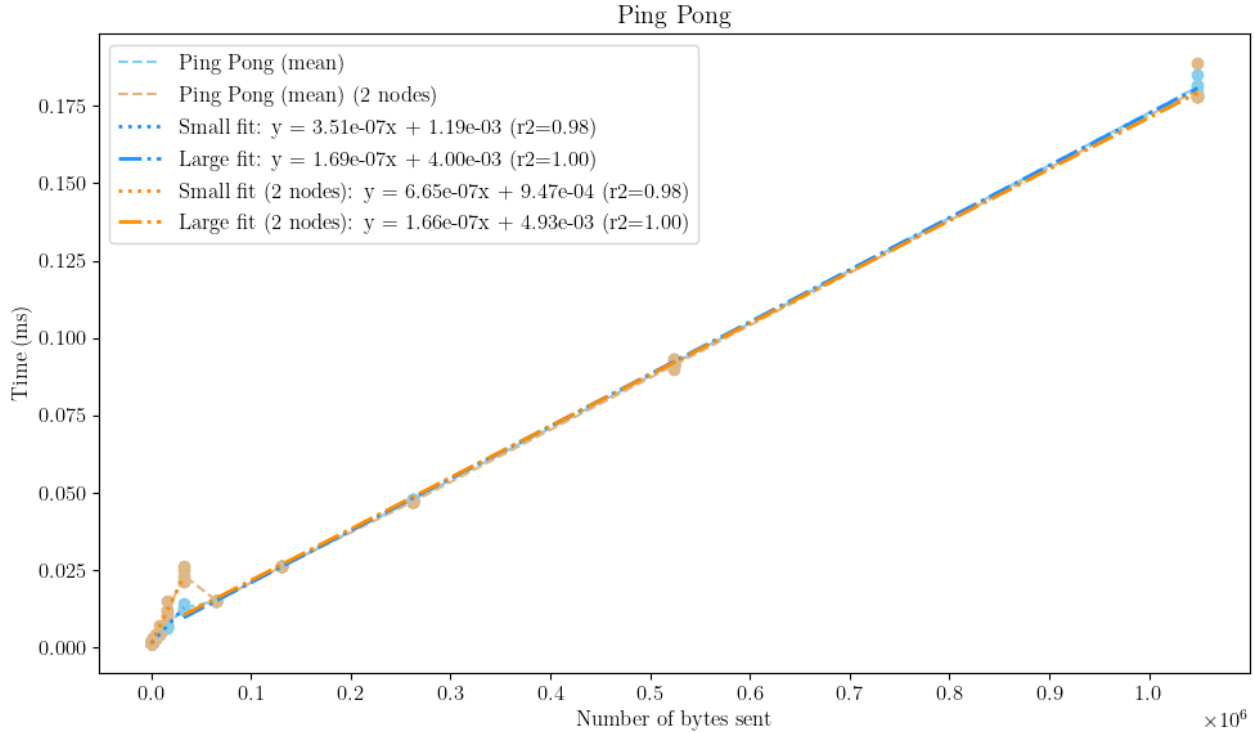


Figure 2: Ping Pong with MPI\_SendRecv: Number of bytes sent vs. average time taken from 1000 pairs of send / receive. 5 runs shown for each size as scatter plot. Mean of these 5 runs shown as line. Blue small fit includes all data points up to 32768 bytes, blue large from there. Red small fit includes all data points up to 32768 bytes, red large from there.

We get the following fits and Rš values for the runs:

Run Type	Data Size	Fit Equation	Rš Value
Single Node	Small ( $\leq 32768$ )	$3.51 \times 10^{-7} \cdot x + 1.19 \times 10^{-3}$	0.98
Single Node	Large ( $\geq 32768$ )	$1.69 \times 10^{-7} \cdot x + 4.00 \times 10^{-3}$	1.00
Two Node	Small ( $\leq 32768$ )	$6.65 \times 10^{-7} \cdot x + 9.47 \times 10^{-4}$	0.98
Two Node	Large ( $\geq 32768$ )	$1.66 \times 10^{-7} \cdot x + 4.93 \times 10^{-3}$	1.00

Table 2: Fit Equations and Rš Values for Single Node and Two Node Runs

**TODO: Further analysis needed?**

## 0.b) MM-product

After an introduction of the matrix-matrix multiplication code in the next section, the measured speedups are discussed in the subsequent section.

### Explanation of the code

For this exercise I've used the template provided in the assignment sheet as a base to develop my parallel implementation for a matrix-matrix multiplication. The code can be found in the appendix for this section.

The program can be run either in sequential (default) or parallel mode (parallel as a command line argument). For the sequential version, the code is practically unchanged and just refactored into a function for timing purposes. The parallel version is more complex and works as explained below:

First, rank 0 computes a sequential reference solution. Then rank 0 distributes the matrices in the following way in `splitwork`:

- Matrix A is split row-wise by dividing the number of rows by the number of nodes.
- The first worker (=rank 1) gets the most rows starting from row 0:  
 $\text{total\_rows} - (\text{nr\_workers} - 1) \cdot \text{floor}(\frac{\text{total\_rows}}{\text{nr\_workers}}).$
- All other workers and the master (= rank 0) get the same number of rows:  $\text{floor}(\frac{\text{total\_rows}}{\text{nr\_workers}}).$
- The master copies the corresponding rows of matrix A and the whole transposed matrix B\* into a buffer (for details on `MM_input` buffer see bellow) for each worker and sends them off using `MPI_Isend`.
- The workers receive the data using `MPI_Recv` and then compute their part of the matrix product and send only the rows of the result matrix back to the master using `MPI_Send`.
- In the meanwhile the master computes its part of the matrix product.
- Using `MPI_Waitall` the master waits for all data to be sent to the workers and only afterwards calls `MPI_Recv` to gather the results from the workers.
- Finally all results are gathered by the master in the result matrix.

Assume we have a 5x5 matrix A and 2 workers (rank 1 and rank 2) and master (rank 0). The partitioning is done row-wise as follows:

#### Partitioning Example

$$A = \begin{pmatrix} a_{11} & a_{12} & a_{13} & a_{14} & a_{15} \\ a_{21} & a_{22} & a_{23} & a_{24} & a_{25} \\ a_{31} & a_{32} & a_{33} & a_{34} & a_{35} \\ a_{41} & a_{42} & a_{43} & a_{44} & a_{45} \\ a_{51} & a_{52} & a_{53} & a_{54} & a_{55} \end{pmatrix} \rightarrow \begin{pmatrix} \text{Worker 1} \\ \text{Worker 1} \\ \text{Worker 1} \\ \text{Master} \\ \text{Master} \end{pmatrix}$$

- **Rank 0 (Master):** Rows 4 and 5 (last two rows)
- **Rank 1 (Worker 1):** Rows 1 to 3 (first three rows) - Worker 1 always gets the most rows

This partitioning can be visually represented as:

$$\text{Master (rank 0): } \begin{pmatrix} a_{41} & a_{42} & a_{43} & a_{44} & a_{45} \\ a_{51} & a_{52} & a_{53} & a_{54} & a_{55} \end{pmatrix}$$

$$\text{Worker 1 (rank 1): } \begin{pmatrix} a_{11} & a_{12} & a_{13} & a_{14} & a_{15} \\ a_{21} & a_{22} & a_{23} & a_{24} & a_{25} \\ a_{31} & a_{32} & a_{33} & a_{34} & a_{35} \end{pmatrix}$$

Each worker computes its part of the matrix product, and the master gathers the results at the end and compiles them into the final matrix.

The `MM_input` buffer is used to store the rows of matrix A and the whole matrix B for each worker. It is implemented using a simple struct:

```
1 typedef struct MM_input {
2     size_t rows;
3     double *a;
4     double *b;
5 } MM_input;
```

**\*[Optimization] Note on transposed matrix B:** It is usually beneficial from a cache perspective to index arrays sequentially or in a row-major order. However, in the matrix-matrix multiplication, we access the elements of matrix B in a column-wise order. This leads to cache misses and is not optimal. To mitigate this, we can transpose matrix B and then access it in a row-wise order. This is done in the code by the master before sending the data to the workers.

### Discussion of the speedups

The code was run on Delft's cluster with 1, 2, 4, 8, 16, 24, 32, 48, and 64 nodes. For the experiments the matrix size of  $A$  and  $B$  was set to  $2000 \times 2000$ . This means that the program has to evaluate 2000 multiplications and 1999 additions for each element of the resulting matrix  $C$ . In total this results in  $\approx 2000^3 = 8 \times 10^9$  operations. The command looked similar to the following for the different node counts:

```
srun -n 48 --mem-per-cpu=4GB --time=00:02:00 ./MM.out parallel
```

For this experiment, the execution time was measured and the speedup was calculated. The results are shown in [Table 3](#) and [Figure 3](#).

CPU Count	Execution Time / s	Approx. Speedup
1	47.11	1.0
2	10.26	4.6
4	10.30	4.6
8	5.20	9.1
16	2.97	15.9
24	2.54	18.5
32	2.29	20.6
48	2.98	15.8
64	1.72	27.4

Table 3: Execution Time vs CPU Count



Figure 3: Speedup vs CPU Count  
Black  $\times$  marks the average of the rerun for  $n = 48$ .

**Note:** The speedup is calculated as  $S = \frac{T_1}{T_p}$ , where  $T_1$  is the execution time on 1 node and  $T_p$  is the execution time on  $p$  nodes.

### Discussion:

As one can clearly discern from the data in [Table 3](#) and [Figure 3](#), the speedup increases with the number of nodes (with the exception of  $n = 48$ ). This is expected as the more nodes we have, the more work can be done in

parallel. However, the speedup is not linear. This is due to the overhead of communication between the nodes. The more nodes we have, the more communication is needed, and this overhead increases. This is especially visible in the data for  $n = 48$ . Here the speedup is lower than for  $n = 32$ . For this run the communication didn't went as smooth as for the other runs. This can potentially be attributed to the fact that one (or more) of the nodes or the network was under heavy load during this task.

**[Further investigation]** After observing this slower speed for the  $n = 48$ , I reran the tests multiple times and got a runtime of around 1.9s which was to be expected initially. Therefore, this one run is an odd one out, most likely due to the reasons mentioned above! I've also added the averaged data of the reruns as a datapoint in [Figure 3](#).

Another interesting fact can be seen when comparing the time taken for  $n = 1$  and  $n = 2$ . They don't at all scale with the expected factor of 2. This is could be due to the fact, that the resource management system prefers runs with multiple nodes instead of a single node (= sequential).

Additional notes: The flag `-mem-per-cpu=<#>GB` was set depending on the number of nodes used. For 1-24 nodes 8GB was used, for 32-48 nodes 4GB, and for 64 nodes 3GB. This had to be done to comply with QOS policy on the cluster.

**TODO: Data locality?**

# 1 Poisson solver

In this section of the lab report, we will discuss a parallel implementation of the Poisson solver. The Poisson solver is a numerical method used to solve the Poisson equation, which is a partial differential equation that is useful in many areas of physics.

**Note:** For local testing and development I'll run the code with `mpirun` instead of the `srun` command on the cluster.

## 1.1 Building a parallel Poisson solver

For the first part of the exercise we follow the steps lined out in the assignment sheet. I'll comment on the steps 1 through 10 and related questions below. The finished implementation can be found in the appendix for this section.

1. **Step:** After adding `MPI_Init` and `MPI_Finalize`, we can run the program with multiple processes. We can see that the program runs with 4 processes in [Figure 4](#) via the quadrupled output.

```
etschgi1@Deep-Thought:~/REPOS/HPC/01_lab1/src$ mpirun -np 4 ./mpi.out
Number of iterations : 2355
Number of iterations : 2355
Number of iterations : 2355
Elapsed procestime:      0.133189 s
Number of iterations : 2355
Elapsed procestime:      0.134150 s
Elapsed procestime:      0.134474 s
Elapsed procestime:      0.135356 s
```

Figure 4: MPI\_Poisson after Step 1 - Running with 4 processes

2. **Step:** To see which process is doing what, I included the rank of the process for the print statements as shown in [Figure 5](#).

```
etschgi1@Deep-Thought:~/REPOS/HPC/02_lab1/src$ mpirun -np 4 ./mpi.out
(0) Number of iterations : 2355
(2) Number of iterations : 2355
(0) Elapsed procestime:      0.135963 s
(2) Elapsed procestime:      0.137101 s
(3) Number of iterations : 2355
(3) Elapsed procestime:      0.139614 s
(1) Number of iterations : 2355
(1) Elapsed procestime:      0.142026 s
```

Figure 5: MPI\_Poisson after Step 2 - Running with 4 processes

3. **Step:** Next we define `wtime` as a global double and replace the four utility timing functions with the ones given on Brightspace. A quick verification as shown in [Figure 6](#) shows that the program still runs as expected.

```
etschgi1@Deep-Thought:~/REPOS/HPC/02_lab1/src$ mpirun -np 4 ./mpi.out
(3) Number of iterations : 2355
(1) Number of iterations : 2355
(3) Elapsed Wtime      0.134918 s ( 98.5% CPU)
(1) Elapsed Wtime      0.134459 s ( 98.5% CPU)
(0) Number of iterations : 2355
(2) Number of iterations : 2355
(0) Elapsed Wtime      0.138669 s ( 98.5% CPU)
(2) Elapsed Wtime      0.138910 s ( 98.5% CPU)
```

Figure 6: MPI\_Poisson after Step 3 - Running with 4 processes

4. **Step:** Next we check if two processes indeed give the same output. Both need 2355 iterations to converge and the `diff` command returned no output, which means that the files content is identical.
5. **Step:** Now only the process with rank 0 will read data from files and subsequently broadcast it to the others. Testing this again with 2 processes, we see an empty diff of the output files and the same number of iterations needed to converge.



6. **Step:** We create a cartesian grid of processes using `MPI_Cart_create` and use `MPI_Cart_shift` to find the neighbors of each process. We can see that the neighbors are correctly identified in [Figure 7](#).

```
(0) (x,y)=(0,0)
(0) top 1, right -2, bottom -2, left 2
(1) (x,y)=(0,1)
(1) top -2, right -2, bottom 0, left 3
(2) (x,y)=(1,0)
(2) top 3, right 0, bottom -2, left -2
(3) (x,y)=(1,1)
(3) top -2, right 1, bottom 2, left -2
```

Figure 7: MPI\_Poisson after Step 6 - Running with 4 processes on a 2x2 grid

When there is no neighbor in a certain direction, -2 (or `MPI_PROC_NULL`) is returned.

7. **Step:** We overhaul the setup to get a proper local grid for each process. Furthermore, we only save the relevant source fields in the local grid for each process.

**With for instance 3 processes you should see that 1 or 2 processes do not do any iteration. Do you understand why?**

If we have a look at the input file we see that there are only 3 source fields in the grid. This means that the process that does not have a source field in its local grid will not do any iterations (or only 1). Therefore, if we have 3 processes and the distribution of source fields as given in the input file only 1 process will do iterations if processes are ordered in x-direction and 2 if ordered in y-direction. From this we can conclude that indeed all processes have different local grids and perform different calculations.

```
etschgi1@Deep-Thought:~/REPOS/HPC/02_lab1/src$ mpirun -np 3 ./mpi.out 3 1
(0) (x,y)=(0,0)
(0) top -2, right -2, bottom -2, left 1
(1) (x,y)=(1,0)
(1) top -2, right 0, bottom -2, left 2
(2) (x,y)=(2,0)
(2) top -2, right 1, bottom -2, left -2
(0) Number of iterations : 1
(2) Number of iterations : 1
(2) Elapsed Wtime 0.000668 s ( 95.3% CPU)
(0) Elapsed Wtime 0.000917 s ( 95.9% CPU)
(1) Number of iterations : 695
(1) Elapsed Wtime 0.014772 s ( 95.2% CPU)
```

```
etschgi1@Deep-Thought:~/REPOS/HPC/02_lab1/src$ mpirun -np 3 ./mpi.out 1 3
(1) (x,y)=(0,1)
(1) top 2, right -2, bottom 0, left -2
(1) Number of iterations : 1
(2) (x,y)=(0,2)
(2) top -2, right -2, bottom 1, left -2
(0) (x,y)=(0,0)
(0) top 1, right -2, bottom -2, left -2
(1) Elapsed Wtime 0.000616 s ( 95.4% CPU)
(0) Number of iterations : 601
(2) Number of iterations : 723
(0) Elapsed Wtime 0.017636 s ( 95.3% CPU)
(2) Elapsed Wtime 0.017801 s ( 95.3% CPU)
```

Figure 8: MPI\_Poisson after Step 7 - Running with 3 processes on a 3x1 (left) vs. 1x3 (right) grid  
For the 3x1 grid, only rank 1 does iterations ( $> 1$ ), for the 1x3 grid, ranks 0 and 2 do iterations ( $> 1$ ).

8. **Step:** After defining and committing two special datatypes for vertical and horizontal communication, we setup the communication logic to exchange the boundary values between the processes. We call our `Exchange_Borders` function after each iteration (for both red / black grid points). Now we face the problem in which some processes may stop instantly (no source in their local grid). They will not supply any data to their neighbors, which will cause the program to hang. We shall fix this in the next step.
9. **Step:** Finally we need to implement the logic to check for convergence (in a global sense). We do this by using a `MPI_Allreduce` call with the `MPI_MAX` operation. This way we aggregate all deltas and choose the biggest one for the global delta which we use in the while-loop-condition to check for convergence. We can see that the program now runs as expected in [Figure 9](#).

```

(0) (x,y)=(0,0)
(0) top -1, right 2, bottom 1, left -1
(1) (x,y)=(0,1)
(1) top 0, right 3, bottom -1, left -1
(2) (x,y)=(1,0)
(2) top -1, right -1, bottom 3, left 0
(3) (x,y)=(1,1)
(3) top 2, right -1, bottom -1, left 1
(0) Number of iterations : 2355
(1) Number of iterations : 2355
(2) Number of iterations : 2355
(3) Number of iterations : 2355
(1) Elapsed Wtime      0.287549 s ( 99.9% CPU)
(2) Elapsed Wtime      0.287537 s (100.0% CPU)
(3) Elapsed Wtime      0.287537 s (100.0% CPU)
(0) Elapsed Wtime      0.295957 s ( 99.9% CPU)

```

Figure 9: MPI\_Poisson after Step 9 - Running with 4 processes on a 2x2 grid

Note that this run in Figure 9 was done with another pc and another MPI implementation. Therefore, we see  $-1$  for cells without a neighbor! However, other than that cosmetic difference it has no impact on the programm.

10. **Step:** Now we only have to fix two remaining things. First we have to make sure that each process uses the right global coordinates for the output file in the end. Therefore, we change the function a bit to include the specific x-/y-offset for each processor. The second thing is the potential problem, that different processors might start with different (red/black) parities. In order to accomplish a global parity we simply have to change the calculation in the if in Do\_Step from

```

1  if ((x + y) % 2 == parity && source[x][y] != 1)

```

to

```

1  if ((x + offset[X_DIR] + y + offset[Y_DIR]) % 2 == parity && source[x][y] != 1)

```

this guarantees that during a given iteration all processors are using the same parity.

This just leaves one question open: Are the results acutally the same?

Checking the output files of the MPI-implementation with the sequential reference indeed shows identical numerical values for the calculated points. Furthermore, the needed iterationcount is also identical which isn't a big surprise, given that the two programmes perform the exact same calculation steps.

## 1.2 Exercises, modifications, and performance aspects

For this subsection we'll define the following shorthand notation:

$pt = 414$  means 4 processors in a  $1 \times 4$  topology.

Table 4: Notation for this section

$n$ :	the number of iterations
$g$ :	gridsize
$t$ :	time needed in seconds
$pt$ :	processor topology in form $pxy$ , where:
$p$ :	number of processors used
$x$ :	number of processors in x-direction
$y$ :	number of processors in y-direction

### Note on different Versions:

For the following exercises the implementation will be slightly adapted to measure different performance aspects. To facilitate this, we will use defines to switch between different versions of the code at compile time. The final version of the poissonsolver can be found in the appendix for this section.

### Note on long scheduling times and work-arounds:

Delft Blue is especially bussy and wait times for jobs can be well over 30 minutes regularly. As we make use of these resources extensively in this lab, I've created `sbatch` scripts which run mutiple configurations at once.

For development of the tests and postprocessing I'll make use of my local machine. As discussed with the tutor of this lab, for some exercises a local run is sufficient to get the desired insights (e.g. [subsubsection 1.2.2](#) to find the optimal omega). I'll also note this in the respective subsections.

### 1.2.1 Over-relaxation (SOR)

We start of by rewriting the `Do_Step` routine to facilitate SOR updates. Furthermore, we need  $h^2$ , the grid spacing (which is 1 in our case) and the relaxation parameter  $\omega$  to calculate the updated values. A quick local test shows a speedup of roughly a factor of 10. More systematic tests will be done in the next section.

### 1.2.2 Optimal $\omega$ for 4 proc. on a 4x1 grid

With the power of a little python scripting we can easily test different values for  $\omega$  and plot the results as seen in [Figure 10](#). This test was performed locally as the results are not dependent on processors because the number of iterations needed for convergence is the same for different configurations and only depends on the algorithm and the specific problem.

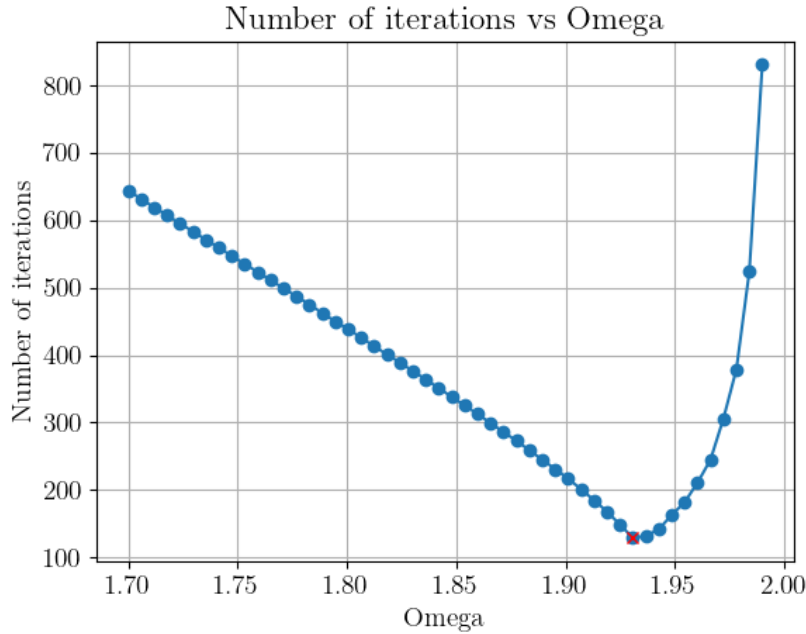


Figure 10: [Optimal  \$\omega\$  for 4 processors on a 4x1 grid](#)

We find that the optimal  $\omega$  is at about 1.93 for this setup with only 129 iterations. This constitutes a speedup of about 1825% compared to the sequential implementation.

**N.B.:** If not stated otherwise, we will use  $\omega = 1.93$  for the following exercises.

### 1.2.3 Scaling behavior with larger grids

This investigation is carried out three times: Once with a  $4 \times 1$  topology (as in the previous section), followed by a  $1 \times 4$  and a  $2 \times 2$  topology. We use grid sizes of  $200 \times 200$ ,  $400 \times 400$ ,  $800 \times 800$  and  $1600 \times 1600$  and set  $\omega = 1.95$  for all runs. All nine simulations are ran using a `sbatch` script on Delft Blue to change the appropriate input parameters and run the program subsequently (details can be found in the Appendix and online in the [repository](#)). The results are shown in [Figure 11](#).

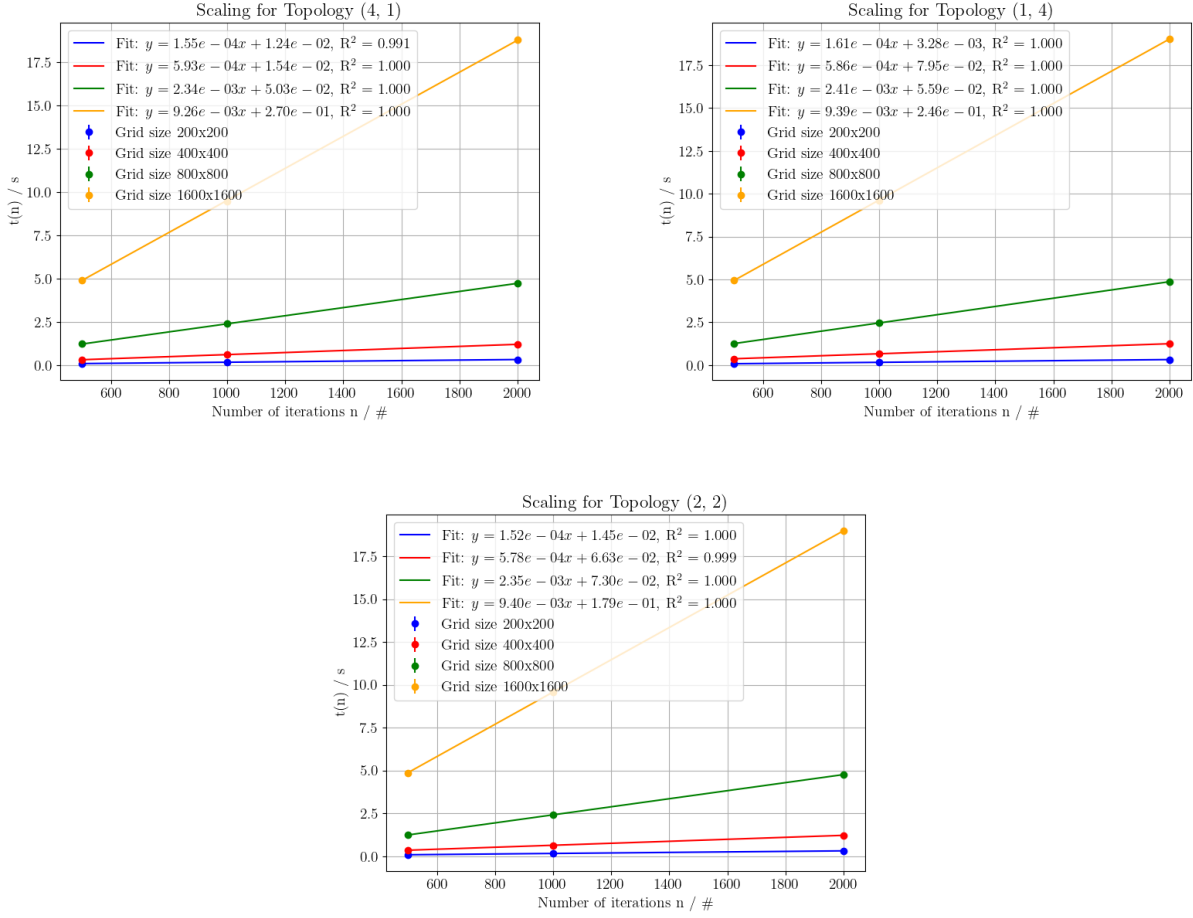


Figure 11: Scaling behavior of the Poisson solver with different grid sizes and processor topologies

As seen by the high  $R^2$  values in the plots, the scaling behavior is very close to linear for the grids. We obtain the following scaling factors for the different grid sizes and topologies from the linear fits:

Table 5: Scaling factors for different processor topologies for the Poisson solver  
Using:  $t(n) = \alpha + \beta \cdot n$  as a model

Topology	Gridsize	$\alpha$	$\beta$
$4 \times 1$	$200 \times 200$	$1.24e-02$	$1.55e-04$
$4 \times 1$	$400 \times 400$	$1.54e-02$	$5.93e-04$
$4 \times 1$	$800 \times 800$	$5.03e-02$	$2.34e-03$
$4 \times 1$	$1600 \times 1600$	$2.70e-01$	$9.26e-03$
$1 \times 4$	$200 \times 200$	$3.28e-03$	$1.61e-04$
$1 \times 4$	$400 \times 400$	$7.95e-02$	$5.86e-04$
$1 \times 4$	$800 \times 800$	$5.59e-02$	$2.41e-03$
$1 \times 4$	$1600 \times 1600$	$2.46e-01$	$9.39e-03$
$2 \times 2$	$200 \times 200$	$1.45e-02$	$1.52e-04$
$2 \times 2$	$400 \times 400$	$6.63e-02$	$5.78e-04$
$2 \times 2$	$800 \times 800$	$7.30e-02$	$2.35e-03$
$2 \times 2$	$1600 \times 1600$	$1.79e-01$	$9.40e-03$

### What can you conclude from the scaling behavior?

We see that the scaling behavior is very close to linear for all topologies. This means that the parallel implementation scales as expected with the number of grid points.

If we compare the scaling factors ( $\beta$ ) for the topologies we see that the  $2 \times 2$  topology scales slightly better than the  $4 \times 1$  and  $1 \times 4$  topologies (except for the largest grid, where all three topologies scale with a very similar

factor). This is not surprising, as the  $2 \times 2$  topology has a more balanced communication workload balance. In the  $2 \times 2$  topology every processor has two neighbors, while in the  $4 \times 1$  and  $1 \times 4$  topologies the processors at the ends only have one neighbor. This is a general trend: A topology which divides the grid into square / square-like parts will scale better than a topology which divides the grid into long and thin parts.

In essence: We want to keep the communication between processors as balanced as possible to achieve the best scaling behavior.

#### Scaling of different grid sizes:

We see that larger grids take longer for the same amount of iterations. This is also to be expected, as the number of grid points grows quadratically with the grid size. a  $800 \times 800$  grid has 4 times as many grid points as a  $400 \times 400$  grid and therefore takes roughly 4 times as long to calculate.

#### 1.2.4 Scaling behavior [Theory - no measurements]

If I could choose between a  $16 \times 1$ ,  $8 \times 2$ ,  $4 \times 4$ ,  $2 \times 8$ ,  $1 \times 16$  topology, I would choose the  $4 \times 4$  topology. This is because the  $4 \times 4$  topology has the most balanced communication workload balance, as detailed in the [Shining](#) in [subsubsection 1.2.3](#).

#### 1.2.5 Iterations needed for convergence scaling

We investigate the number of iterations needed for convergence using the  $4 \times 1$  topology square grids with sidelength: 10, 25, 50, 100, 200, 400, 800, 1600. The results for different  $\omega$  are shown in [Figure 12](#). This test was performed locally as the results are the same for different systems and only depend on the algorithm and the specific problem.

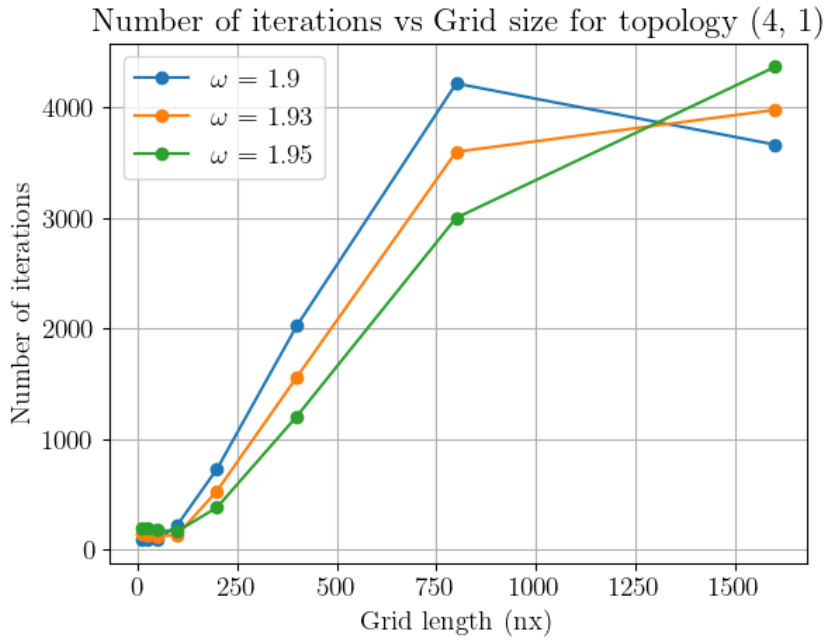


Figure 12: Iterations needed for convergence with different grid side lengths

We can clearly see that the number of iterations till convergence increases with the problem size. At first, I expected linear growth proportional to the number of gridpoints. However, it turns out that the number of iterations actually grow slower and in a square root like fashion. This can be seen by the linear behavior in the plot of grid-side length against iterations.

#### Why is the number of iterations needed for convergence $\propto \sqrt{g}$ ?

Our poisson problem is a discretized system in 2D space. The condition number of the matrix we have to solve is proportional to the number of gridpoints in our system. SOR uses the spectral properties of the matrix to solve in a way such that the dominant error mode takes time proportional to the diameter of the domain to converge. This means it is proportional to  $\sqrt{g} = \sqrt{n_x \cdot n_y}$ .

### Why does omega with the best performance change with the grid size?

As can be seen in Figure 12  $\omega = 1.9$  beats the other two values for very small and the largest gridsize. For different gridsizes we get differently sized matrices we have to solve. SOR overrelaxes high-frequency errors and underrelaxes low-frequency errors (the later for stability). The optimal  $\omega$  is indeed dependent on the gridsize and the error modes present in the system. In our current example, it might be that  $\omega = 1.9$  is a good compromise for the grid sizes we are looking at and we are so to say lucky with that specific choice.

#### 1.2.6 Error as a function of the iteration number

With the same  $4 \times 1$  topology and grid sizes of  $800 \times 800$  the error for 15000 iterations is tracked using  $\omega = 1.93$ . The results are shown in Figure 13.

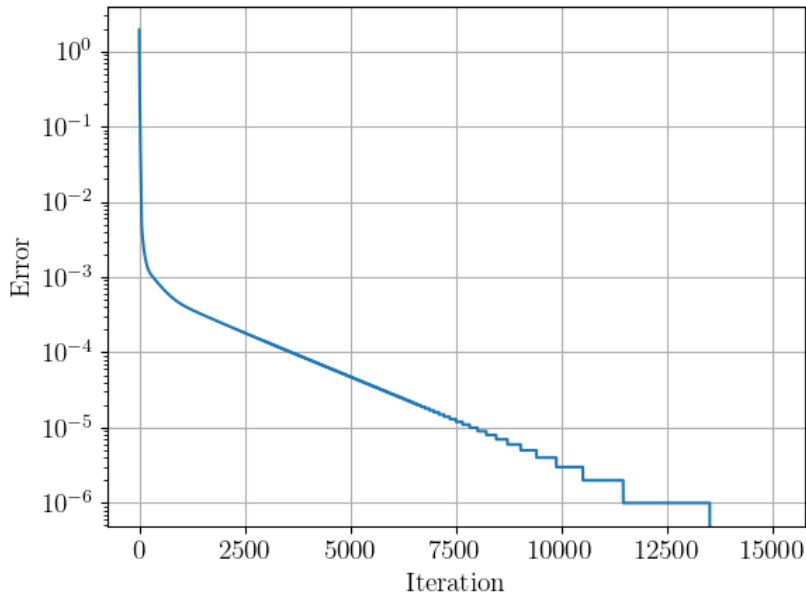


Figure 13: Error as a function of the iteration number

At first the error decreases rapidly in the first few iterations to about  $10^{-3}$  (logarithmic scale!). After that the error decreases more slowly until it is below floating point precision.

**Note:** All calculations are done using double precision floating point numbers and only the error recording was done using single precision which leaves the step-like artifacts in the plot. Obviously these steps would also be present in the double precision error calculation, but they would be much smaller at comparable iteration numbers and only become visible at much larger iteration numbers.

#### 1.2.7 Optional - Gain performance by reducing MPI\_Allreduce calls

The last subsection showed us that the error reduces monotonically. We might be able to save some time by leaving out some checks and maybe check the global error every 10th or 100th iteration only.

First, we should benchmark if it is at all wise to optimize here, by measuring how long the MPI\_Allreduce call takes. We can do this by measuring the time needed for the MPI\_Allreduce call in the Do\_Step function and summing up to get the total time spent in MPI\_Allreduce calls.

We again solve with a  $4 \times 1$  topology,  $\omega = 1.93$  and a  $800 \times 800$  grid: It takes roughly 20 seconds of which the processors spend around 1 - 2 seconds in the MPI\_Allreduce call. This is a significant amount of time ( $(7.0 \pm 0.4)\%$ ). This means we would save some time by reducing the number of MPI\_Allreduce calls and calculating 9 (0.25% of total) more iterations wouldn't hurt us too bad because it takes 3601 to converge!

We run the program three times with MPI\_Allreduce calls every 1, 10 and 100 iterations and get the speedups in MPI\_Allreduce calls as shown in Table 6.

Table 6: Speedup in MPI\_Allreduce calls for different iteration counts and calculated overall speedup (%)

Iterations	MPI_Allreduce - speedup (factor)	calculated overall speedup (%)
1	1.00	0
10	$6.0 \pm 2.0$	$5.9 \pm 0.5$
100	$62 \pm 6$	$6.9 \pm 0.4$

As can be clearly seen from the table we can gain around 6 % using MPI\_Allreduce calls every 10 iterations and around 7 % using MPI\_Allreduce calls every 100 iterations. This is a significant speedup for a very small change in the code.

**Note:** The speedup is calculated to account for fluctuations in the runtime of the program, due to other processes running on the same machine / cluster.

### 1.2.8 Reduce border communication

Another way to reduce communication overhead is to reduce the number of border exchanges. To investigate if this yields a speedup we run the program on a  $4 \times 1$  topology,  $\omega = 1.93$  and different grid sizes and track the iterations and time as seen in Figure 14.

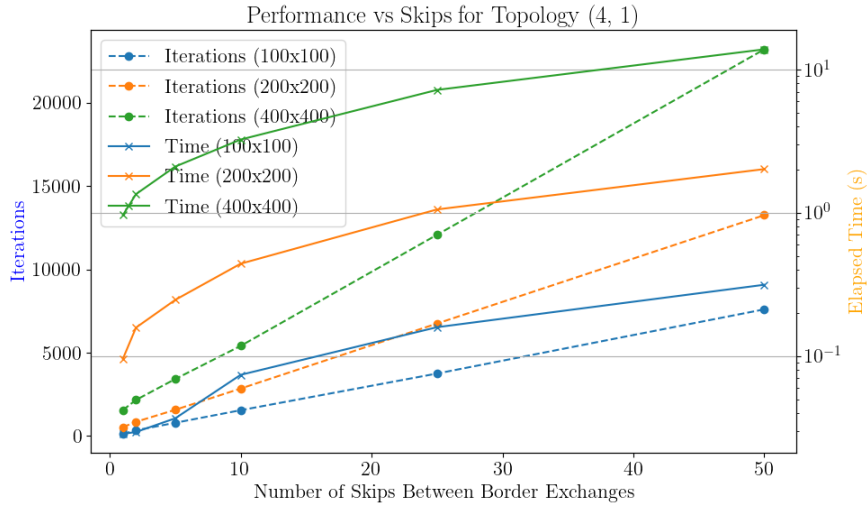


Figure 14: Speedup by reducing border exchanges (4x1 topology)

Running the with different numbers of skipped border exchanges naturally slows down convergence, meaning we need more iterations to reach the same error. For all tested grid sizes the initial SOR version without skipping border exchanges has the fewest iterations needed to convergence and also the fastest runtime.

#### What can you conclude from the results?

We can conclude that reducing the number of border exchanges does not yield a speedup. The reason for this is that we have to calculate more iterations to converge to the solution which outweighs the gains from reduced communication overhead. Interestingly, for the  $100 \times 100$  grid there exists a local minimum in time at 4 skipped border exchanges compared to 3 skipped. This is likely due to our source field distribution and thus specific to our problem.

Running the problem again with another ( $2 \times 2$  topology specifically) on our Delft Blue node we get the same qualitative result as seen in Figure 15.

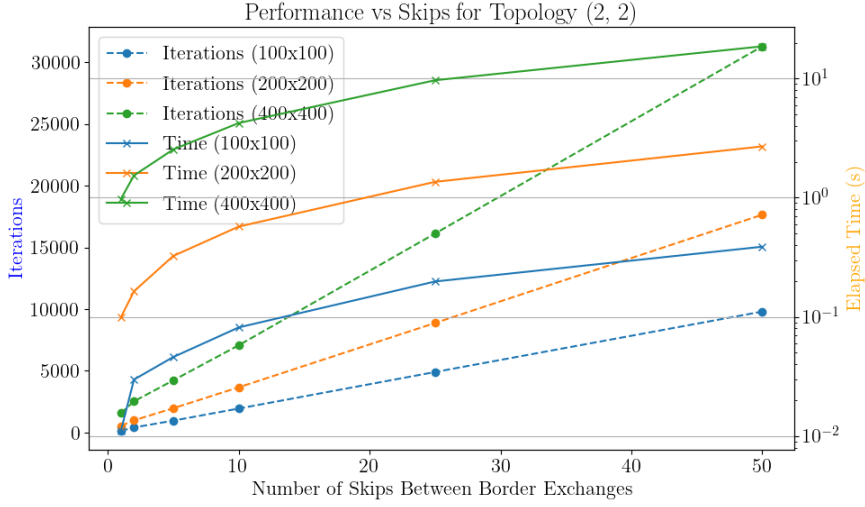


Figure 15: Speedup by reducing border exchanges (2x2 topology)

#### Taking a closer look at $2 \times 2$ vs. $4 \times 1$ topology:

While both have qualitatively the same behavior with the fastest time and lowest iterations recorded for the SOR version without skipping border exchanges, the  $2 \times 2$  topology has a worse convergence behavior in terms of iterations. This is likely due to the fact that the borders of the local grids for the processors in the  $2 \times 2$  topology are positioned in closer proximity to the source coordinates compared to the  $4 \times 1$  topology. This could explain the observed higher iteration count for convergence in all grid sizes in the  $2 \times 2$  topology.

#### 1.2.9 Optimize Do\_Step loop

In `Do_Step` we iterate over the whole grid but only update one of the two parities at a time. This means we can split the loop into two loops, one for each parity. We start out with something like this:

```

1  for (x = 1; x < dim[X_DIR] - 1; x++){
2      for (y = 1; y < dim[Y_DIR] - 1; y++){
3          if ((x + offset[X_DIR] + y + offset[Y_DIR]) % 2 == parity && source[x][y] != 1){
4              ...

```

and we change it to:

```

1  int start_y;
2  for (x = 1; x < dim[X_DIR] - 1; x++){
3      start_y = ((1 + x + offset[X_DIR] + offset[Y_DIR]) % 2 == parity) ? 1 : 2;
4      for (y = start_y; y < dim[Y_DIR] - 1; y += 2){
5          if (source[x][y] != 1){
6              ...

```

The basic idea is to avoid y-coordinates which are not in the parity we are currently updating. We measure 10 runs for a  $800 \times 800$  grid and a  $4 \times 1$  topology with  $\omega = 1.93$  and get the following times:

$$t_{\text{no\_improvements}} = (5.59 \pm 0.05) \text{ s} \quad \text{and} \quad t_{\text{loop\_improvements}} = (4.64 \pm 0.07) \text{ s}$$

So we get a minimal speedup of about 17% by optimizing the loop which is a enormous speedup for such a small change.

#### Why does this make such a difference

The reason for this is that we avoid unnecessary looping and if statements. This means that we have less overhead in the loop and can therefore calculate faster by skipping the unnecessary loop entries.

#### 1.2.10 Optional - Time spent within Exchange\_Borders

We can measure the time spent in `Exchange_Borders` by adding a timer to the function. We run the program with  $\omega = 1.93$  and different topologies<sup>1</sup> and grid sizes and get the results shown in Figure 16.

<sup>1</sup>{(2,2), (3,3), (4,4), (5,5), (6,6), (2,3), (2,4), (2,5), (3,4)}



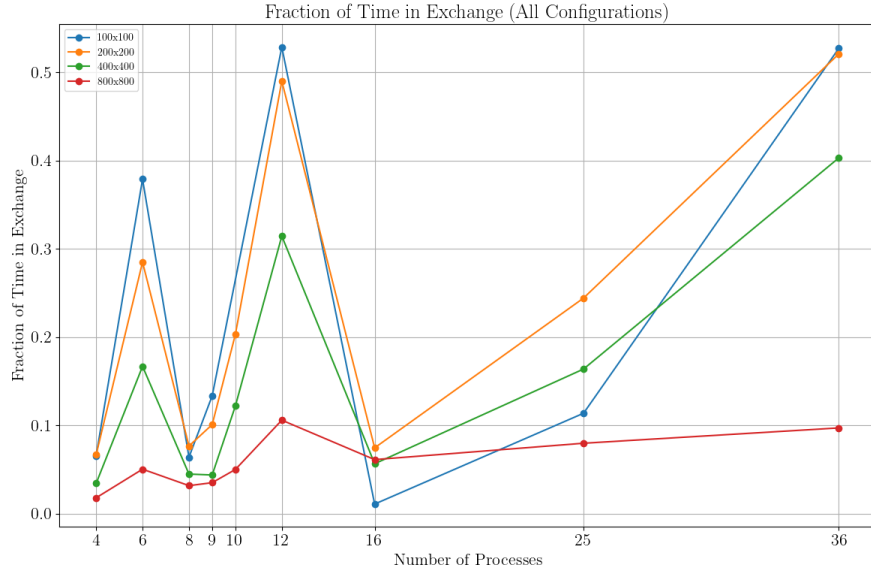


Figure 16: Fraction of total time spent in `Exchange_Borders`

As we can clearly see, the time spent in `Exchange_Borders` is initially smaller and grows with processor count with some remarkable peaks especially for small grids and certain processor topologies. The curves are generally shifted downward for larger gridsizes.

#### Interpretation:

One would expect larger grid sizes to be more computationally expensive and we have already established that iterations take longer the bigger the grid. Communication obviously also takes longer for a larger grid because we have more data to sent. However, the circumference of a square grows linearly, while the area grows quadratically. The quadratic growth of the area is the reason for the downward shift in the curves for larger grid sizes because the communication overhead grows slower than the computational overhead for larger grids. Besides the peaks at  $p = 6$  and  $p = 12$  we see a general trend of increasing time spent in `Exchange_Borders` increasing processor count. This is not surprising as we have more processes which have to communicate with each other and the data locality is worse for larger processor counts.

#### When is the time spent in `Exchange_Borders` significant / comparable to computation?

As can be seen in Figure 16 the time spent in `Exchange_Borders` is significant for all grid sizes from the start (between 2.5 and 7.5%). Thereafter it peaks for  $p = 6$  and again for  $p = 12$  to around 5% to 38% and 10% to 54% respectively. This means that the time spent in `Exchange_Borders` is significant for all grid sizes and processor counts, but especially as the processor count grows.

#### 1.2.11 Latency and bandwith in `Exchange_Borders`

We use the configurations from subsection 1.2.3:  $4 \times 1$ ,  $2 \times 2$  and  $3 \times 3$  topologies with grid sizes of  $200 \times 200$ ,  $400 \times 400$  and  $800 \times 800$  and  $\omega = 1.95$  as well as the other settings set to their defaults. We obtain the results in Table 7.

Table 7: Metrics for `Exchange_Borders` latency and bandwith

Topology	Grid Size	Latency (ms)	Latency (%)	Bandwidth ( $\text{Bs}^{-1}$ )	Total Data (B)
4x1	200x200	$2.0 \pm 0.6$	3.9	2 034 737 840	3 104 896
4x1	400x400	$9.8 \pm 0.4$	1.4	2 005 275 269	19 450 368
4x1	800x800	$51 \pm 23$	0.8	2 112 229 431	96 480 384
2x2	200x200	$3.6 \pm 1.0$	5.0	541 474 183	2 493 696
2x2	400x400	$17 \pm 7$	2.3	668 287 123	15 591 168
2x2	800x800	$92 \pm 37$	1.4	665 489 803	77 261 184
3x3	200x200	$161 \pm 80$	43.9	9 006 796	1 686 912
3x3	400x400	$105 \pm 53$	18.9	56 268 947	10 419 840
3x3	800x800	$219 \pm 67$	6.2	2 369 103 880	51 603 552

### Interpretation:

As can be seen the latency is lowest for the  $4 \times 1$  topology followed by  $2 \times 2$  and  $3 \times 3$ . This is not surprising as the  $4 \times 1$  topology has the least amount of neighbors to communicate with per processor. The worst latencies can generally be observed for the  $3 \times 3$  topology because every processor has to communicate with 2,3 or 8 neighbors. As can be seen by the huge discrepancy in the latency percentage for the  $3 \times 3$  topology, the latency is strongly dependent on the grid size (problem size). While nearly half of the time is spent in latency for the  $200 \times 200$  grid, only 6.2% of the time is spent in latency for the  $800 \times 800$  grid.

#### 1.2.12 Exchange Border potential improvements

Indeed we communicate twice as much as we need after each `Do_Step` call. We shall analyze this considering the following points:

- address of the first point to exchange: This depends on the parity of the processor. It is simply the first or second point in the grid depending on the parity (leaving out the edge case where the processor only has one data-point).
- the number of points to exchange: This normally is the number of points in the grid minus ghost cells divided by two. However, this may change if there is an odd number of points in the grid (then we have to exchange one more or one less point).
- the number of points in between grid points that have to be exchanged: The stride of the data will change. Currently we exchange every point for one direction and every `dim[Y_DIR]`-th point for the other direction. This can be optimized to exchange every second point in one direction and every second `dim[Y_DIR]`-th point in the other direction.

#### Is it worth it?

For smaller gridsizes it is not worth it to optimize the border exchange. The time spent in the border exchange might be significant in relative terms, but the absolute time is still small. For larger gridsizes it might be worth it to optimize the border exchange. As we have seen, this becomes more significant as the processor count and gridsizes grow. For our current problem and similarly sized problems the effort put into optimizing the border exchange is certainly not worth it.

## 2 Finite elements simulation

We will now shift our focus to a more general grid which is based on triangulation. In this section we will compare our parallel implementation from the previous section and dissect the differences and similarities. For that reason we shall use the same sources in our grid as given in `sources.dat`.

Note that the sections for the exercises will be labeled as (2.1, 2.2, 2.3, ...) corresponding to exercises (4.1, 4.2, 4.3, ...) in the lab manual.

### 2.1 Code understanding & Exchange\_Borders

The first step is to read through the code in `MPI_FemPois.c` and to understand it. Furthermore, we have to implement the `Exchange_Borders` function for which only a skeleton is given. The function should exchange the border values of the local grid with the neighboring processes.

The implementation of this function is quite straight forward. We only have to loop over all the neighbors of a process and send out the border values and receive the border values from the neighbors. The function is implemented as follows:

```
1 void Exchange_Borders(double *vect)
2 {
3     for (size_t i = 0; i < N_neighb; ++i)
4     {
5         MPI_Sendrecv(vect, 1, send_type[i], proc_neighb[i], 0, //send
6                     vect, 1, recv_type[i], proc_neighb[i], 0, //recv
7                     grid_comm, &status);
8     }
9 }
```

### 2.2 Time benchmarking

Next we turn our attention to timing of different sections in the code.

We have to measure:

- Time spent in computation
- Time spent exchanging information with neighbors
- Time spent doing global communication
- Idle time

We setup the following variable to measure / deduce the time spent in the different sections:

```
1 double total_time = 0.0;
2 double exchange_time_neighbors = 0.0;
3 double exchange_time_global = 0.0;
4 double compute_time = 0.0;
```

We will measure the time spent in computations by timing the solve function and subtracting the time spent in the `MPI_Allreduce` calls. The time spent in the `MPI_Allreduce` calls is the time spent in global communication. The time spent in exchanging information with neighbors is the time spent in the `Exchange_Borders` function. Finally, the idle time can be determined by summing up the differences between the cores total time and the slowest cores total time. Note that this way of calculating the idle time is an approximation since it is assumed that the slowest core doesn't have any idle time. However, I've discussed this with the TA and he said that this is a valid way of calculating the idle time for this exercise.

The commandline output for runs is as shown in [Figure 17](#) for an example run:

```

(2) - Exchange time (neighbors): 0.007183
(2) - Exchange time (global): 0.002372
(2) - Sum of times (compute + exchange (global & local)): 0.062241
(2) - Total time: 0.069645
(0) - Total time: 0.069800
(1) - Compute time: 0.051027
(1) - Exchange time (neighbors): 0.005517
(1) - Exchange time (global): 0.004273
(1) - Sum of times (compute + exchange (global & local)): 0.060816
(1) - Total time: 0.069416
(3) - Compute time: 0.051977
(3) - Exchange time (neighbors): 0.006824
(3) - Exchange time (global): 0.003341
(3) - Sum of times (compute + exchange (global & local)): 0.062142
(3) - Total time: 0.069399

```

Figure 17: Timing for the different sections.

(x) ... denotes the process rank

Compute time ... time spent in the `solve` function only on computing

Exchange time (neighbors) ... time spent in `Exchange_Borders`

Exchange time (neighbors) ... time spent in `MPI_Allreduce` calls

Sum of times ... total time spent in compute and communication (excluding setup / idle time)

Total time ... total time spent in the program

The idle time is calculated as denoted above and therefore not shown in the output in Figure 17. We get the following results from our Delft Blue runs in Table 8:

Table 8: Rank averaged time benchmark for different grid sizes and topologies.

All times are in milliseconds.

**Note:** WTime does also include setup and teardown (mallocs, frees, etc.) - therefore the sum of the times is not equal to WTime.

Top.	Grid Size	WTime (avg)	Comp. (avg)	Ex. Neighb. (avg)	Ex. Global (avg)	Idle (avg)
1x4	100x100	82.0	18.6	1.3	1.7	9.0
1x4	200x200	194.5	150.6	3.6	6.8	12.1
1x4	400x400	1498.2	1357.8	10.1	26.4	9.1
2x2	100x100	43.7	18.6	1.6	1.4	9.1
2x2	200x200	184.0	145.2	4.3	8.1	5.7
2x2	400x400	1428.7	1279.2	11.5	22.4	25.6

We see that the total time is comparable between the two topologies for all grid sizes. Furthermore, the total runtime also increases non-surprisingly with the gridsize.

### Why does computation time increase faster than linearly?

One could expect, that the grid with  $200 \times 200$  elements would take 4 times as long as the grid with  $100 \times 100$  elements. However, the runtime roughly 8 times longer. This happens because additionally to the higher number of grid points the algorithm also takes longer (more iterations) to converge and hence the computation time is longer.

### Analysis of the different times

To get a better grasp on the data a stacked bar plot, as shown in Figure 18, is created. We immediately see that the bulk of the time is spent on computing the solution. The second biggest contributor is the global exchange time and the idle time, followed by local exchange time.

That computation takes up the biggest part of the time is of non surprise especially on the bigger grids this is to be expected. Rather surprisingly the global exchange time takes up a lot of time. This begs the question why two lines of the form:

```
1 MPI_Allreduce(... , ... , 1, MPI_DOUBLE, MPI_SUM, grid_comm);
```

take up this much time. This actually comes down to masked idle time. While the operation itself only sums up the values of 4 double, the operation is blocking and therefore the other cores have to wait for the slowest core to finish. If we look back at the definition of idle time, we defined it as the difference of the total

time of the slowest core and the total time of the current core. However, as stated above, this does not take waiting time in MPI calls (like `MPI_Allreduce`) into account. Certainly, this explains now that the global exchange time is high, because every core has to wait for the others to synchronize to compute the `MPI_Allreduce`.

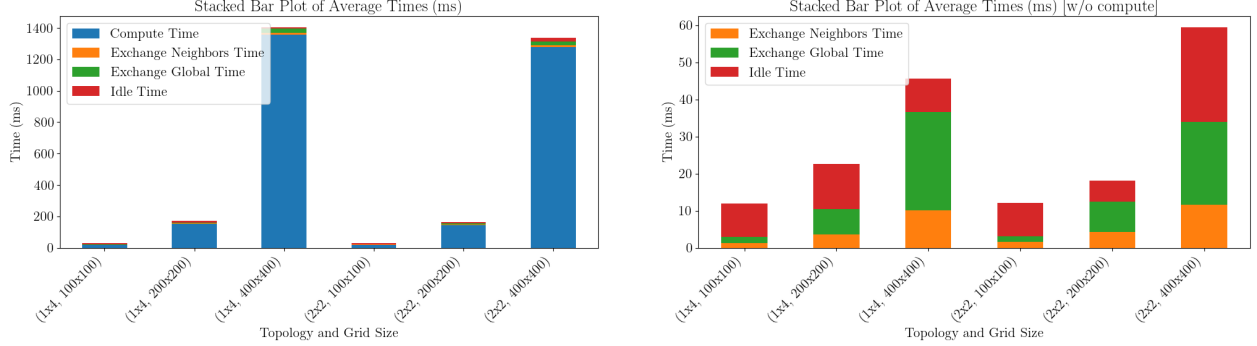


Figure 18: Scaling behavior of the Poisson solver with different grid sizes and processor topologies

We conclude that most of the time spent on communication in our solver is actually due to the waiting time in the `MPI_Allreduce` calls and the other idle time (waiting for the slowest core to finish). The actual exchange of data only represents a fraction of this time. Biggest, especially for larger grids, contributor for the time is still the computation which takes up north of 95% of the total time on larger grids.

### 2.3 Data exchange amount

This section is about the amount of data exchanged each iteration among one process with all its neighbors. We assume a uniformly triangulated grid which is partitioned stripe-wise and distribution over  $P$  processes. Furthermore, we assume that each process has to send the same amount of data. Indeed, this is not generally the case but for examples with periodic boundary conditions this is for example a valid assumption. Now we see that every process has to communicate with  $2d$  neighbors, where  $d$  is the dimension of the grid, under our assumptions there are 2 neighboring processes. With the assumption of a  $n^2$  grid, our process communicates  $n$  values with each of these neighbors. For striped partitioning we get:

$$\text{Data exchanged per Process} = 2n$$

or in total:

$$\text{Data exchanged} = 2n \cdot P$$

Let's check for the extreme case  $1000 \times 1000$  grid and  $P = 500$  processes. We get:

$$\text{Data exchanged} = 2 \cdot 1000 \cdot 500 = 1000000$$

Which is the same amount of data as there are datapoints. This makes sense because every process has to communicate the top row upwards and the bottom row downwards.

Note, that we could even do worse if we assigned every process a single row. In this case every process would have to communicate the same data upward and downward. This would result in a data exchange of  $2 \cdot 1000 \cdot 1000 = 2000000$  which is double the amount of data as there are datapoints.

For a box partitioning a process has to communicate  $n/\sqrt{P}$  (assuming divisibility with all neighbors) with each of its 4 neighbors. We get:

$$\text{Data exchanged per Process} = 4 \cdot n/\sqrt{P}$$

or in total:

$$\text{Data exchanged} = 4 \cdot n \cdot \sqrt{P}$$

This means that a striped partitioning is less efficient compared to a boxed partitioning starting from  $P = 4$ , as can be seen in Figure 19.

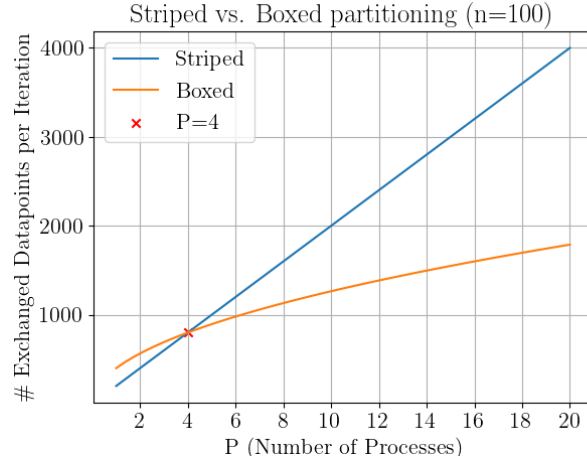


Figure 19: Striped vs. Boxed partitioning on a  $100 \times 100$  grid using  $P$  processors.

## 2.4 Unbalanced communication

In the last section we assumed that every process has to communicate with the same amount of neighbors in our given scenario. That is actually not entirely true. There is an imbalance, even if just a small one.

**But where does this imbalance come from?**

Let's take a look at a **uniformly triangulated grid** with  $P = 4$  processes. We assume that the grid is box-partitioned as given in Figure 20.

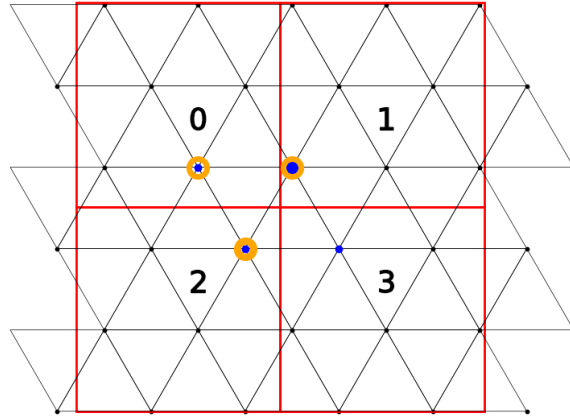


Figure 20: Box partitioning of a uniformly triangulated grid with  $P = 4$  processes.

One point (big-blue) in 1 (and similarly in 3) has to communicate its value to 3 neighbors (small-blue).

Another point (big-orange circle) in 2 (and similarly in 4) has to communicate its value to 2 neighbors (small-orange circles).

We see that the geometry of a given uniform triangulated grid leads to an imbalance in communication. This imbalance is usually not a big problem, especially for bigger grids with comparably small amounts of processors because only 2 points have to communicate with 3 neighbors. However, for smaller grids or proportionally high processor counts the imbalance is more significant.

Looking at a  $3 \times 3$  grid with  $P = 9$  processors, we see that the imbalance is more significant as shown in Figure 21.

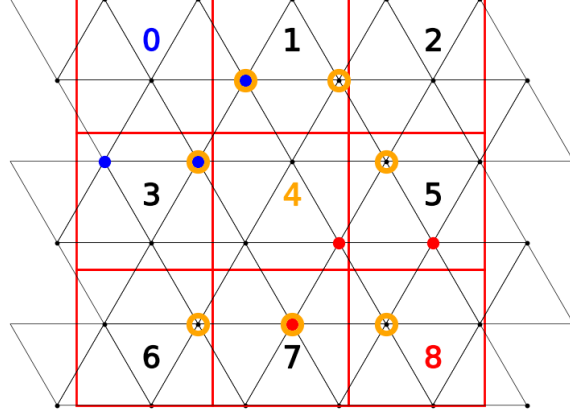


Figure 21: Box partitioning of a uniformly triangulated grid with  $P = 9$  processes. Corner (0/8 - blue/red) processes have to communicate their values to 2 or 3 neighbors respectively (1 and 3 - blue or 4, 5 and 7 - red). Central (4 - orange) processes have to communicate their value to 6 neighbors (1, 3, 5, 6, 7 and 8).

As we can see the imbalance comes from the partitioning geometry on the uniform triangulated grid. Because our processor grid is rectangular the results will always, in one way or another come out as shown in the schematic above.

For  $3 \times 3$  grid we have 4 corner processes which have to communicate with 2 or 3 neighbors and 1 central process which has to communicate with 6 neighbors. This is a significant imbalance and compared to the  $2 \times 2$  grid it is not negligible, because the central process has to communicate more than 2 extra points (whole edges of datapoints) to its neighbors.

## 2.5 Estimates for computation $\equiv$ communication time

**N.B.** This section will use two different ways to estimate the equilibrium point.

1) Let's assume that we have 4 processes and want to estimate for which grid size the computation time is equal to the communication time. One way to estimate this is by fitting a polynomial of degree 2 to the computation and a linear fit to the exchange time for our measurements in Table 8 and find the point where the times are equal. Thereafter we can use this information to determine the grid size where this happens. This process gives us a value of  $n \approx 84$  for the grid size as shown in Figure 22

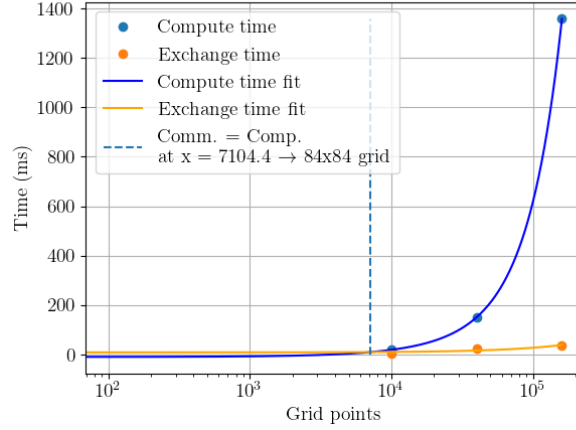


Figure 22: Fitting expected computation time  $\equiv$  communication time for 4 processes.

2) We'll also estimate this equilibrium point for a  $1000 \times 1000$  grid. However, this time around we'll actually use a different way of calculating the number of Processes. We use the definitions from the lecture for stencil type computation:

- Computation time:  $T_S = \text{number\_ops} \cdot t_{\text{op}} \sim 4t_{\text{op}} \cdot n^2$  (for stencil type - overall)
- Communication time:  $T_{\text{comm}} = \text{number\_comm} \cdot t_{\text{comm}} \sim 2n \cdot t_{\text{data}}$  (for stencil type - overall)

Note that the definitions above apply for a single iteration and **one** processor!

We can calculate the time for communication from the data in Table 8 as follows:

$$t_{\text{data}}(P) = \frac{t_{\text{global\_comm}} + t_{\text{neighbor\_comm}}}{2 \cdot n \cdot P \cdot \#_{\text{iterations}}}$$

Similarly we can calculate the time for computation as follows:

$$t_{\text{op}} = \frac{t_{\text{comp.}}}{4 \cdot n^2 \cdot \#_{\text{iterations}}}$$

Note that the later doesn't depend on the number of processes.

Using the data from Table 8 we determine that one operation takes  $t_{\text{op}} = (3.3 \pm 0.3) \times 10^{-6}$  ms. From that we can determine the computation time  $t_{\text{comp.}}$  for  $n = 1000$  as follows:

$$t_{\text{comp.}} = 4 \cdot n^2 \cdot t_{\text{op}} \cdot \#_{\text{iterations}} = 4 \cdot 1000^2 \cdot 3.3(3) \cdot 10^{-6} \cdot 1241 = (17.8 \pm 1.5) \text{ s}$$

where 1241 ( $\#_{\text{iterations}}$ ) is the number of iterations to converge. We set  $t_{\text{comp.}} = t_{\text{data}}$  and solve for  $P$ . We need values for  $t_{\text{global\_comm}}$  and  $t_{\text{neighbor\_comm}}$  and take a look at Figure 22 and decide to further investigate the trend for the exchange time ( $= t_{\text{global\_comm}} + t_{\text{neighbor\_comm}}$ ). We decide to use a linear extrapolation as shown in Figure 23. We get an estimated communication time of 100 ms.

Now we have everything to solve for  $P$  and using 1241 for  $\#_{\text{iterations}}$  again we get  $P = 29 \pm 3$  processes.



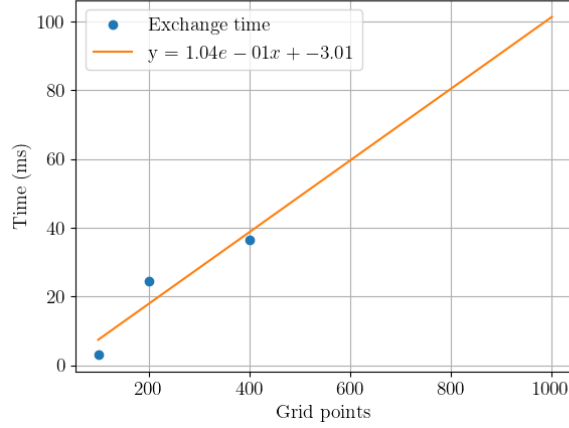


Figure 23: Extrapolating the communication time for  $n = 1000$  using a linear fit.

Our predictions are in: a  $84 \times 84$  grid for 4 processes with equal computation and communication times and a  $1000 \times 1000$  grid for 29 processes (also having equal computation and communication time).

We measure the ratio  $t_{\text{comp}}/t_{\text{comm}}$  and get the following results for a  $4 \times 1$  topology in Table 9:

Table 9: Ratio of computation to communication time for different grid sizes.

Grid Size	Ratio
10x10	$0.53 \pm 0.06$
20x20	$0.84 \pm 0.15$
30x30	$0.86 \pm 0.13$
40x40	$1.97 \pm 0.24$
50x50	$2.66 \pm 0.61$
75x75	$4.37 \pm 1.00$
84x84	$3.70 \pm 0.85$
90x90	$6.06 \pm 1.94$

We see that our initial guess of  $84 \times 84$  grid for 4 processes is off by a long-shot. However, we also notice that the ratio is highly volatile as can be seen by the high uncertainties on the ratio. A rerun of the code produced notably different ratios (e.g.  $1.35 \pm 0.25$  for  $30 \times 30$  grid). It seems that the actual position of the equilibrium point is around  $n = 35$ . As a physicist I have to say that our initial guess was of the same magnitude and therefore not too far off ☺.

Let's see how our prediction for the  $1000 \times 1000$  grid with 29 processes holds up. Using a  $29 \times 1$  topology we get  $4.04 \pm 1.83$  for the ratio. It seems we have made a lower prediction in terms of  $P$  for the equilibrium point. However, the ratio is in the same ballpark as the ratio for the  $84 \times 84$  grid which is not surprising since we used the same underlying data to make the prediction.

## 2.6 Adaptive grid

We will now make our grid denser in the source point areas. `GridDist` already implements an argument (*adapt*) for that purpose. In order to gauge the speed of convergence we let process 0 print the current precision of the solution. We perform runs with a reference and a denser grid to compare the convergence speed, convergence count and amount of computing time for a  $2 \times 2$  topology and a  $100 \times 100$ ,  $200 \times 200$  and  $400 \times 400$  grids.

We first take a look at the iterations needed for convergence using the standard precision goal of 0.0001 we get the results in Table 10.

It becomes clear that the adaptive grid needs a few more iterations to converge. This is most likely due to the fact that the dense region near the sources initially lead to a slower 'spreading' of the solution near the source.

Table 10: Iterations needed for convergence on  $2 \times 2$  topology for different grid sizes with and without adapt keyword.

Grid Size	100x100	200x200	400x400
Iterations Reference	141	274	529
Iterations Adapt	146	278	532

### Does such a distorted grid lead to faster convergence?

We can answer this question with a clear no, at least in terms of iterations. Next we'll look at the time it takes to converge.

A bar-plot of the time till convergence is shown in Figure 24. We see that the time till convergence is roughly the same for the reference and the adaptive grid. This is not surprising since the time till convergence is mainly determined by the number of iterations needed.

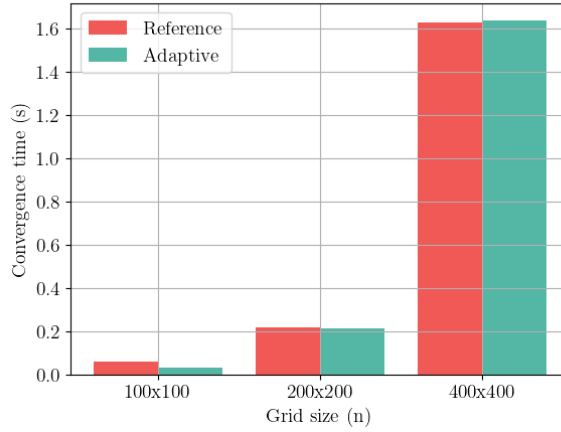


Figure 24: Time till convergence (with setup) on  $2 \times 2$  topology for different grid sizes with and without adapt keyword.

### Does it affect the speed of convergence?

Also no for this one. The time for the reference / adaptive grid is roughly the same with one exception for the  $100 \times 100$  grid.

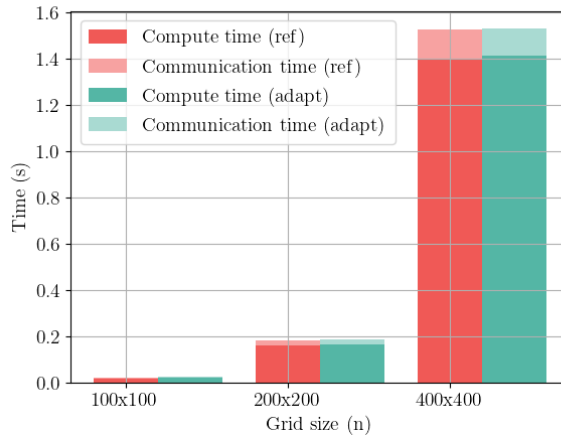


Figure 25: Time till convergence (without setup) on  $2 \times 2$  topology for different grid sizes with and without adapt keyword.

If we stack the time needed for computation and communication we see that the computation as well as the

communication time is roughly the same for the reference and the adaptive grid as shown in [Figure 25](#).

**Does it affect the amount of computing time?**

So again no, the amount of computing time is roughly the same for the reference and the adaptive grid.

### 3 Eigenvalue solution by Power Method on GPU

The last problem concerns the evaluation of eigenvalue using the power method via a parallellized CUDA code. Reference for this implementation is a sequantial CPU-code provided by the course (`power_cpu.cu`).

A scematic overview of the iteration loop for the power method is shown bellow in algorithm 1.

---

**Algorithm 1** GPU Power Method

---

```
1: Input: Matrix  $\mathbf{A}$  of size  $N \times N$ , tolerance  $\epsilon$ , maximum iterations  $max\_iter$ 
2: Output: Dominant eigenvalue  $\lambda$ 
3: Initialize  $\mathbf{v}$  with  $v_1 = 1, v_i = 0$  for  $i > 1$ 
4:  $\lambda_{Old} \leftarrow 0, \lambda \leftarrow 0$ 
5: Allocate GPU memory for  $\mathbf{A}, \mathbf{v}, \mathbf{w}$ , and  $\lambda$ 
6: Copy  $\mathbf{A}$  and  $\mathbf{v}$  to GPU memory
7:  $\mathbf{w} \leftarrow \mathbf{A} \cdot \mathbf{v}$  ▷ First iteration of  $\mathbf{w}$  computation using Av_Product kernel
8: for  $i = 0$  to  $max\_iter - 1$  do
9:   Compute norm of  $\mathbf{w}$ :  $norm \leftarrow \sqrt{\mathbf{w}^T \cdot \mathbf{w}}$  ▷ Using FindNormW kernel
10:  Normalize  $\mathbf{v}$ :  $\mathbf{v} \leftarrow \mathbf{w}/norm$  ▷ Using NormalizeW kernel
11:  Compute  $\mathbf{w} \leftarrow \mathbf{A} \cdot \mathbf{v}$  ▷ Using Av_Product kernel
12:  Compute eigenvalue:  $\lambda \leftarrow \mathbf{v}^T \cdot \mathbf{w}$  ▷ Using FindNormW kernel
13:  if  $|\lambda - \lambda_{Old}| < \epsilon$  then
14:    Break ▷ Convergence achieved
15:  end if
16:   $\lambda_{Old} \leftarrow \lambda$ 
17: end for
18: Copy  $\lambda$  back to host memory
19: Deallocate GPU memory
```

---

**Note:** A `sqrt()` was added in the `NormalizeW` kernel over `g_NormW[0]`. This way we can use the output of `FindNormW` directly in the `NormalizeW` kernel.

All of the following benchmarks are perfomed in the supplied IPython notebook on Google Collab using T4 GPUs.

#### 3.1 Step 1: Shared vs. global memory for matrix-vector multiplication

**TODO: open**

#### 3.2 Step 2: Execution time for different N and threads per block

I implemented a small loop to run the GPU code for 5 different  $N$  with  $N \in \{50, 500, 2000, 4000, 5000\}$ . The resulting time benchmarks for 32, 64 and 100 threds per block can be seen in [Figure 26](#).

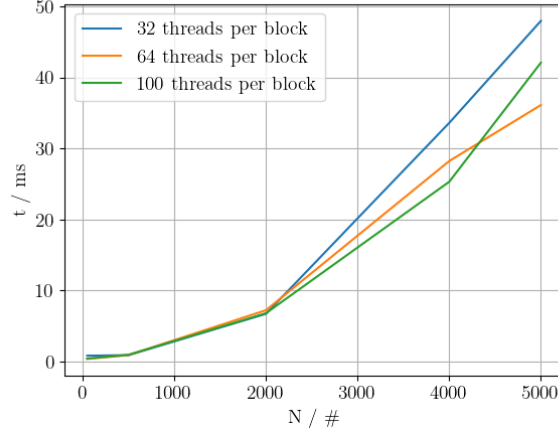


Figure 26: Runtime for  $N \in \{50, 500, 2000, 4000, 5000\}$  and for 32, 64 and 100 threads per block respectively.

**TODO: maybe add somethin, justification whatever?!**

### 3.3 Step 3: Speedups

We measure two different scenarios:

- i excluding time of memory copy from CPU  $\rightarrow$  GPU
- ii including time of memory copy from CPU  $\rightarrow$  GPU

After measuring 5 rounds without (i) and with (ii) memory access time we obtain the following [scatter plot](#) in [Figure 27](#)

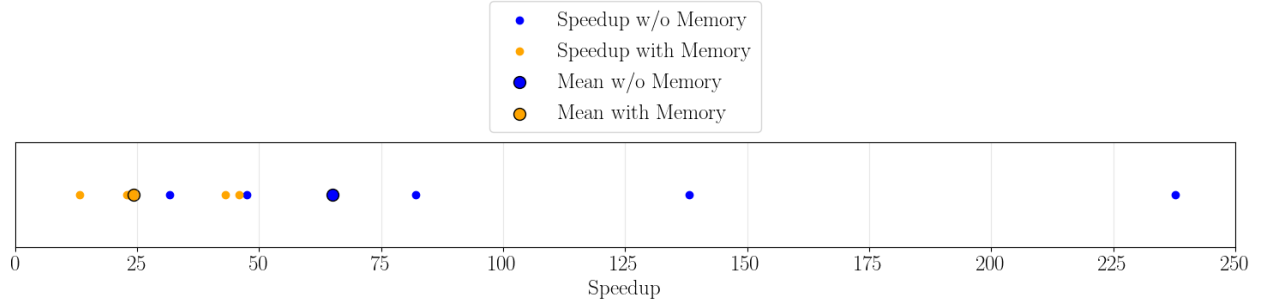


Figure 27: Speedup of GPU implementation vs. CPU with and without memory transfer times.

The mean speedup without memory access times is  $\times 65$  and with memory access timed it comes out to  $\times 24$ .

### 3.4 Step 4: Explanation of the results

## Appendix - Introductory exercise

The following code was used for the ping pong task:

```
1 #include <stdio.h>
2 #include <stdlib.h>
3 #include <mpi.h>
4
5 // Maximum array size 2^20= 1048576 elements
6 #define MAX_EXPONENT 20
7 #define MAX_ARRAY_SIZE (1<<MAX_EXPONENT)
8 #define SAMPLE_COUNT 1000
9
10 int main(int argc, char **argv)
11 {
12     // Variables for the process rank and number of processes
13     int myRank, numProcs, i;
14     MPI_Status status;
15
16     // Initialize MPI, find out MPI communicator size and process rank
17     MPI_Init(&argc, &argv);
18     MPI_Comm_size(MPI_COMM_WORLD, &numProcs);
19     MPI_Comm_rank(MPI_COMM_WORLD, &myRank);
20
21
22     int *myArray = (int *)malloc(sizeof(int)*MAX_ARRAY_SIZE);
23     if (myArray == NULL)
24     {
25         printf("Not enough memory\n");
26         exit(1);
27     }
28     // Initialize myArray
29     for (i=0; i<MAX_ARRAY_SIZE; i++)
30         myArray[i]=1;
31
32     int number_of_elements_to_send;
33     int number_of_elements_received;
34
35     // PART C
36     if (numProcs < 2)
37     {
38         printf("Error: Run the program with at least 2 MPI tasks!\n");
39         MPI_Abort(MPI_COMM_WORLD, 1);
40     }
41     double startTime, endTime;
42
43     // TODO: Use a loop to vary the message size
44     for (size_t j = 0; j <= MAX_EXPONENT; j++)
45     {
46         number_of_elements_to_send = 1<<j;
47         if (myRank == 0)
48         {
49             myArray[0]=myArray[1]+1; // activate in cache (avoids possible delay when sending
the 1st element)
50             startTime = MPI_Wtime();
51             for (i=0; i<SAMPLE_COUNT; i++)
52             {
53                 MPI_Send(myArray, number_of_elements_to_send, MPI_INT, 1, 0,
54                     MPI_COMM_WORLD);
55                 MPI_Probe(MPI_ANY_SOURCE, MPI_ANY_TAG, MPI_COMM_WORLD, &status);
56                 MPI_Get_count(&status, MPI_INT, &number_of_elements_received);
57
58                 MPI_Recv(myArray, number_of_elements_received, MPI_INT, 1, 0,
59                     MPI_COMM_WORLD, MPI_STATUS_IGNORE);
60             } // end of for-loop
61
62             endTime = MPI_Wtime();
63             printf("Rank %2.1i: Received %i elements: Ping Pong took %f seconds\n", myRank,
number_of_elements_received, (endTime - startTime)/(2*SAMPLE_COUNT));
64         }
65         else if (myRank == 1)
66         {
67             // Probe message in order to obtain the amount of data
68             MPI_Probe(MPI_ANY_SOURCE, MPI_ANY_TAG, MPI_COMM_WORLD, &status);
```

```

69     MPI_Get_count(&status, MPI_INT, &number_of_elements_received);
70
71     for (i=0; i<SAMPLE_COUNT; i++)
72     {
73         MPI_Recv(myArray, number_of_elements_received, MPI_INT, 0, 0,
74                 MPI_COMM_WORLD, MPI_STATUS_IGNORE);
75         MPI_Send(myArray, number_of_elements_to_send, MPI_INT, 0, 0,
76                 MPI_COMM_WORLD);
77     } // end of for-loop
78 }
79 }
80
81 // Finalize MPI
82 MPI_Finalize();
83
84 return 0;
85 }

```

For the bonus task, the following code was used:

```

1  #include <stdio.h>
2  #include <stdlib.h>
3  #include <mpi.h>
4
5  // Maximum array size 2^20= 1048576 elements
6  #define MAX_EXPONENT 20
7  #define MAX_ARRAY_SIZE (1<<MAX_EXPONENT)
8  #define SAMPLE_COUNT 1000
9
10 int main(int argc, char **argv)
11 {
12     // Variables for the process rank and number of processes
13     int myRank, numProcs, i;
14     MPI_Status status;
15
16     // Initialize MPI, find out MPI communicator size and process rank
17     MPI_Init(&argc, &argv);
18     MPI_Comm_size(MPI_COMM_WORLD, &numProcs);
19     MPI_Comm_rank(MPI_COMM_WORLD, &myRank);
20
21
22     int *myArray = (int *)malloc(sizeof(int)*MAX_ARRAY_SIZE);
23     if (myArray == NULL)
24     {
25         printf("Not enough memory\n");
26         exit(1);
27     }
28     // Initialize myArray
29     for (i=0; i<MAX_ARRAY_SIZE; i++)
30         myArray[i]=1;
31
32     int number_of_elements_to_send;
33     int number_of_elements_received;
34
35     // PART C
36     if (numProcs < 2)
37     {
38         printf("Error: Run the program with at least 2 MPI tasks!\n");
39         MPI_Abort(MPI_COMM_WORLD, 1);
40     }
41     double startTime, endTime;
42
43     // TODO: Use a loop to vary the message size
44     for (size_t j = 0; j <= MAX_EXPONENT; j++)
45     {
46         number_of_elements_to_send = 1<<j;
47         if (myRank == 0)
48         {
49             myArray[0]=myArray[1]+1; // activate in cache (avoids possible delay when sending
the 1st element)
50             startTime = MPI_Wtime();
51             for (i=0; i<SAMPLE_COUNT; i++)
52             {
53                 MPI_Sendrecv(myArray, number_of_elements_to_send, MPI_INT, 1,0,myArray,

```

```

    number_of_elements_to_send, MPI_INT, 1, 0, MPI_COMM_WORLD, &status);
54     }
55
56     endTime = MPI_Wtime();
57     printf("Rank %2.i: Received %i elements: Ping Pong took %f seconds\n", myRank,
    number_of_elements_to_send, (endTime - startTime)/(2*SAMPLE_COUNT));
58     }
59     else if (myRank == 1)
60     {
61         for (i=0; i<SAMPLE_COUNT; i++)
62         {
63             MPI_Sendrecv(myArray, number_of_elements_to_send, MPI_INT, 0,0,myArray,
    number_of_elements_to_send, MPI_INT, 0, 0, MPI_COMM_WORLD, &status);
64         }
65     }
66 }
67
68 // Finalize MPI
69 MPI_Finalize();
70
71 return 0;
72 }

```

The matrix multiplication used the following code:

```

1  /*****
2  * FILE: mm.c
3  * DESCRIPTION:
4  *   This program calculates the product of matrix a[nra][nca] and b[nca][ncb],
5  *   the result is stored in matrix c[nra][ncb].
6  *   The max dimension of the matrix is constraint with static array
7  *   declaration, for a larger matrix you may consider dynamic allocation of the
8  *   arrays, but it makes a parallel code much more complicated (think of
9  *   communication), so this is only optional.
10 *
11 *****/
12
13 #include <math.h>
14 #include <mpi.h>
15 #include <stdbool.h>
16 #include <stdio.h>
17 #include <stdlib.h>
18 #include <string.h>
19
20 #define NRA 2000 /* number of rows in matrix A */
21 #define NCA 2000 /* number of columns in matrix A */
22 #define NCB 2000 /* number of columns in matrix B */
23 // #define N 1000
24 #define EPS 1e-9
25 #define SIZE_OF_B NCA*NCB*sizeof(double)
26
27 bool eps_equal(double a, double b) { return fabs(a - b) < EPS; }
28
29 void print_flattened_matrix(double *matrix, size_t rows, size_t cols, int rank) {
30     printf("[%d]\n", rank);
31     for (size_t i = 0; i < rows; i++) {
32         for (size_t j = 0; j < cols; j++) {
33             printf("%10.2f ", matrix[i * cols + j]); // Accessing element in the 1D array
34         }
35         printf("\n"); // Newline after each row
36     }
37 }
38
39 int checkResult(double *truth, double *test, size_t Nr_col, size_t Nr_rows) {
40     for (size_t i = 0; i < Nr_rows; ++i) {
41         for (size_t j = 0; j < Nr_col; ++j) {
42             size_t index = i * Nr_col + j;
43             if (!eps_equal(truth[index], test[index])) {
44                 return 1;
45             }
46         }
47     }
48     return 0;
49 }

```



```

50
51 typedef struct {
52     size_t rows;
53     double *a;
54     double *b;
55 } MM_input;
56
57 char* getbuffer(MM_input *in, size_t size_of_buffer){
58     char* buffer = (char*)malloc(size_of_buffer * sizeof(char));
59     if (buffer == 0)
60     {
61         printf("Buffer couldn't be allocated.");
62         return NULL;
63     }
64     size_t offset = 0;
65     memcpy(buffer + offset, &in->rows, sizeof(size_t));
66     offset += sizeof(size_t);
67     size_t matrix_size = in->rows * NCA * sizeof(double);
68     memcpy(buffer + offset, in->a, matrix_size);
69     offset += matrix_size;
70     memcpy(buffer + offset, in->b, NCA*NCB*sizeof(double));
71     return buffer;
72 }
73
74 MM_input* readbuffer(char* buffer, size_t size_of_buffer){
75     MM_input *mm = (MM_input*)malloc(sizeof(MM_input));
76
77     mm->rows = ((size_t*)buffer)[0];
78     size_t offset = sizeof(size_t);
79     size_t matrix_size = mm->rows * NCA;
80     mm->a = (double*)malloc(sizeof(double)*matrix_size);
81     mm->b = (double*)malloc(sizeof(double)*matrix_size);
82     memcpy(mm->a, &(buffer[offset]), matrix_size);
83     offset += matrix_size;
84     memcpy(mm->b, &(buffer[offset]), NCA*NCB*sizeof(double));
85     free(buffer);
86     return mm;
87 }
88
89
90 void setupMatrices(double (*a)[NCA], double (*b)[NCB], double (*c)[NCB]){
91     for (size_t i = 0; i < NRA; i++) {
92         for (size_t j = 0; j < NCA; j++) {
93             a[i][j] = i + j;
94         }
95     }
96
97     for (size_t i = 0; i < NCA; i++) {
98         for (size_t j = 0; j < NCB; j++) {
99             b[i][j] = i * j;
100         }
101     }
102
103     for (size_t i = 0; i < NRA; i++) {
104         for (size_t j = 0; j < NCB; j++) {
105             c[i][j] = 0;
106         }
107     }
108 }
109
110 double multsum(double* a, double* b_transposed, size_t size){
111     double acc = 0;
112     for (size_t i = 0; i < size; i++)
113     {
114         acc += a[i]*b_transposed[i];
115     }
116     return acc;
117 }
118
119 double productSequential(double *res) {
120     // dynamically allocate to not run into stack overflow - usually stacks are
121     // 8192 bytes big -> 1024 doubles but we have 1 Mio. per matrix
122     double(*a)[NCA] = malloc(sizeof(double) * NRA * NCA);

```

```

123 double(*b)[NCB] = malloc(sizeof(double) * NCA * NCB);
124 double(*c)[NCB] = malloc(sizeof(double) * NRA * NCB);
125
126 /** Initialize matrices */
127 setupMatrices(a,b,c);
128
129 /* Parallelize the computation of the following matrix-matrix
130 multiplication. How to partition and distribute the initial matrices, the
131 work, and collecting final results.
132 */
133 // multiply
134 double start = MPI_Wtime();
135 for (size_t i = 0; i < NRA; i++) {
136     for (size_t j = 0; j < NCB; j++) {
137         for (size_t k = 0; k < NCA; k++) {
138             res[i * NCB + j] += a[i][k] * b[k][j];
139         }
140     }
141 }
142 /* perform time measurement. Always check the correctness of the parallel
143 results by printing a few values of c[i][j] and compare with the
144 sequential output.
145 */
146 double time = MPI_Wtime()-start;
147 free(a);
148 free(b);
149 free(c);
150 return time;
151 }
152
153 double splitwork(double* res, size_t num_workers){
154     if (num_workers == 0) // sadly noone will help me :(
155     {
156         printf("Run sequential!\n");
157         return productSequential(res);
158     }
159
160     double(*a)[NCA] = malloc(sizeof(double) * NRA * NCA);
161     double(*b)[NCB] = malloc(sizeof(double) * NCA * NCB);
162     double(*c)[NCB] = malloc(sizeof(double) * NRA * NCB);
163     // Transpose matrix b to make accessing columns easier - in row major way - better cache
164     // performance
165     setupMatrices(a,b,c);
166
167     double start_time = MPI_Wtime();
168     double (*b_transposed)[NCA] = malloc(sizeof(double) * NCA * NCB);
169     for (size_t i = 0; i < NCA; i++) {
170         for (size_t j = 0; j < NCB; j++) {
171             b_transposed[j][i] = b[i][j];
172         }
173     }
174
175     /** Initialize matrices */
176     // given number of workers I'll split
177     size_t rows_per_worker = NRA / (num_workers+1); //takes corresponding columns from other
178     // matrix
179     printf("rows per worker: %zu\n", rows_per_worker);
180     size_t row_end_first = NRA - rows_per_worker*num_workers;
181     printf("first gets most: %zu\n", row_end_first);
182
183     //setup requests
184     MPI_Request requests[num_workers];
185     MM_input *data_first = (MM_input*)malloc(sizeof(MM_input));
186     data_first->rows = row_end_first;
187     data_first->a = (double*)a; //they both start of with no offset!
188     data_first->b = (double*)b_transposed;
189     size_t total_size = sizeof(size_t) + (data_first->rows * NCA)*sizeof(double)+SIZE_OF_B;
190     char* buffer = getbuffer(data_first, total_size); //first one
191
192     // Tag is just nr-cpu -1
193     MPI_Isend(buffer, total_size, MPI_CHAR, 1, 0, MPI_COMM_WORLD, &requests[0]);
194     free(data_first);
195     total_size = sizeof(size_t) + (rows_per_worker * NCA)*sizeof(double) + SIZE_OF_B; //size

```

```

194 is the same for all other - just compute once!
195 size_t i;
196 for (i = 0; i < (num_workers-1); ++i)
197 {
198     MM_input *data = (MM_input*)malloc(sizeof(MM_input));
199     data->rows = rows_per_worker;
200     data->a = (double*)(a + (row_end_first + rows_per_worker*i));
201     data->b = (double*)(b_transposed); // send everything - all needed
202     buffer = getbuffer(data, total_size);
203     printf("nr_worker - %zu\n", i);
204     MPI_Isend(buffer, total_size, MPI_CHAR, i+2, i+1, MPI_COMM_WORLD, &requests[i+1]);
205     free(data);
206 }
207 double* my_a = (double*)(a + (row_end_first + rows_per_worker*i));
208 //I multiply the rest
209 size_t offset = 0;
210 for (size_t row = (NRA-rows_per_worker); row < NRA; row++)
211 {
212     for (size_t col = 0; col < NCB; col++)
213     {
214         res[row * NCB + col] = multsum(my_a+offset, (((double*)b_transposed)+col*NCA), NCA);
215     }
216     offset += NCA;
217 }
218 printf("My c: \n");
219 //wait for rest
220 MPI_Status stats[num_workers];
221 if(MPI_Waitall(num_workers, requests, stats) == MPI_ERR_IN_STATUS){
222     printf("Communication failed!!! - abort\n");
223 }
224 printf(">>>Everything sent and recieved\n");
225
226 // reviece rest
227 size_t buf_size = sizeof(double)*row_end_first*NCB;
228 double* revbuf;
229 offset = 0;
230 for (size_t worker = 0; worker < num_workers; worker++)
231 {
232     revbuf = (double*)malloc(buf_size); //first gets largest buffer
233     MPI_Recv(revbuf, buf_size/sizeof(double), MPI_DOUBLE, worker+1, worker, MPI_COMM_WORLD,
234     &stats[worker]);
235     memcpy(&res[offset/sizeof(double)], revbuf, buf_size);
236     free(revbuf);
237     offset += buf_size;
238     buf_size = sizeof(double)*rows_per_worker*NCB;
239 }
240 double time = MPI_Wtime()-start_time;
241 //free all pointers!
242 free(a);
243 free(b);
244 free(b_transposed);
245 free(c);
246 return time;
247 }
248
249
250 double work(int rank, size_t num_workers){
251     size_t rows_per_worker = NRA / (num_workers+1);
252     char* buffer;
253     MPI_Status status;
254     if (rank == 1) // first always get's most work
255     {
256         rows_per_worker = NRA - rows_per_worker*num_workers;
257     }
258     size_t size_of_meta = sizeof(size_t);
259     size_t size_of_a = sizeof(double)*rows_per_worker*NCA;
260     size_t buffersize = size_of_meta+size_of_a + SIZE_OF_B;
261     buffer = (char*)malloc(buffersize);
262
263     MPI_Recv(buffer, buffersize, MPI_CHAR, 0, rank-1, MPI_COMM_WORLD, &status);

```

```

264     double start = MPI_Wtime();
265     int count;
266     MPI_Get_count(&status, MPI_CHAR, &count);
267     printf("I'm rank %d and I got %d bytes (%ld doubles) of data from %d with tag %d.\n", rank
, count, (count-sizeof(size_t))/sizeof(double), status.MPI_SOURCE, status.MPI_TAG);
268
269     MM_input *mm = (MM_input*)malloc(sizeof(MM_input));
270     mm->a = (double*)&buffer[size_of_meta];
271     mm->b = (double*)&buffer[size_of_meta+size_of_a];
272
273     double *res =(double*)malloc(sizeof(double)*rows_per_worker*NCB);
274
275     size_t offset = 0;
276     for (size_t row = 0; row < rows_per_worker; row++)
277     {
278         for (size_t col = 0; col < NCB; col++)
279         {
280             res[row * NCB + col] = multsum(mm->a+offset, (((double*)mm->b)+col*NCA), NCA);
281         }
282         offset += NCA;
283     }
284     MPI_Send(res, rows_per_worker*NCB, MPI_DOUBLE, 0,rank-1, MPI_COMM_WORLD);
285     printf("[%d] sent res home\n",rank);
286     free(res);
287     return MPI_Wtime() - start;
288 }
289
290 int main(int argc, char *argv[]) {
291     int tid, nthreads;
292     /* for simplicity, set NRA=NCA=NCB=N */
293     // Initialize MPI, find out MPI communicator size and process rank
294     int myRank, numProcs;
295     MPI_Status status;
296     MPI_Init(&argc, &argv);
297     MPI_Comm_size(MPI_COMM_WORLD, &numProcs);
298     MPI_Comm_rank(MPI_COMM_WORLD, &myRank);
299     int num_Workers = numProcs-1;
300     if (argc > 1 && strcmp(argv[1], "parallel") == 0) {
301         // Variables for the process rank and number of processes
302         if (myRank == 0) {
303             printf("Run parallel!\n");
304             double *truth = malloc(sizeof(double) * NRA * NCB);
305             double time = productSequential(truth);
306             printf("Computed reference results in %.6f s\n", time);
307             printf("Hello from master! - I have %d workers!\n", num_Workers);
308             // send out work
309             double *res = malloc(sizeof(double)*NRA*NCB);
310             time = splitwork(res, num_Workers);
311             if (checkResult(res, truth, NCB, NRA)) {
312                 printf("Matrices do not match!!!\n");
313                 return 1;
314             }
315             printf("Matrices match (parallel [eps %.10f])! - took: %.6f s\n", EPS, time);
316             free(truth);
317             free(res);
318         } else {
319             double time = work(myRank, num_Workers);
320             printf("Worker bee %d took %.6f s (after recv) for my work\n", myRank, time);
321         }
322     } else // run sequential
323     {
324         printf("Run sequential!\n");
325         double *res = malloc(sizeof(double) * NRA * NCB);
326         double time = productSequential(res);
327         if (checkResult(res, res, NCB, NRA)) {
328             printf("Matrices do not match!!!\n");
329             return 1;
330         }
331         printf("Matrices match (sequential-trivial)! - took: %.6f s\n", time);
332         free(res);
333     }
334 }
335

```

```

336     MPI_Finalize();
337     return 0;
338 }

```

## Appendix - Poisson solver

The parallel Poisson solver used the following code:

Note: Sbatch scripts used for the exercises will be included after the Poisson-solver code.

```

1  /*
2  * MPI_Poisson.c
3  * 2D Poisson equation solver (parallel version)
4  */
5
6  #include <stdio.h>
7  #include <stdlib.h>
8  #include <math.h>
9  #include <time.h>
10 #include <mpi.h>
11 #include <assert.h>
12
13 #define DEBUG 0
14
15 #define max(a,b) ((a)>(b)?a:b)
16
17
18 // defines for Exercises!
19
20 #define SOR 1
21 #define MONITOR_ERROR 1
22 #define FAST_DO_STEP_LOOP
23 // #define MONITOR_ALLREDUCE 1
24 // #define ALLREDUCE_COUNT 100
25 #define MONITOR_EXCHANGE_BORDERS
26 #define SKIP_EXCHANGE
27
28 #define DEFINES_ON (SOR || MONITOR_ERROR || 0)
29 //defines end
30
31 enum
32 {
33     X_DIR, Y_DIR
34 };
35
36 // only needed for certain configs!
37 #ifdef SOR
38 double sor_omega = 1.9;
39 #endif
40 #ifdef MONITOR_ERROR
41 double *errors=NULL;
42 #endif
43 #ifdef MONITOR_ALLREDUCE
44 double all_reduce_time = 0;
45 #endif
46 #ifdef MONITOR_EXCHANGE_BORDERS
47 double total_exchange_time = 0.0; // Total time spent in exchanges
48 double total_latency = 0.0; // Total latency
49 double total_data_transferred = 0.0; // Total data transferred
50 int num_exchanges = 0; // Number of exchanges
51 #endif
52 #ifdef SKIP_EXCHANGE
53 size_t skip_exchange;
54 #endif
55
56 /* global variables */
57 int gridsize[2];
58 double precision_goal; // precision_goal of solution */
59 int max_iter; // maximum number of iterations allowed */
60 int P; //total number of processes
61 int P_grid[2]; // process grid dimensions

```

```

62 MPI_Comm grid_comm; //grid communicator
63 MPI_Status status;
64 double hx, hy;
65
66 /* process specific globals*/
67 int proc_rank;
68 double wtime;
69 int proc_coord[2]; // coords of current process in processgrid
70 int proc_top, proc_right, proc_bottom, proc_left; // ranks of neighboring procs
71 // step 7
72 int offset[2] = {0,0};
73 // step 8
74 MPI_Datatype border_type[2];
75
76 /* benchmark related variables */
77 clock_t ticks; /* number of systemticks */
78 int timer_on = 0; /* is timer running? */
79
80 /* local grid related variables */
81 double **phi; /* grid */
82 int **source; /* TRUE if subgrid element is a source */
83 int dim[2]; /* grid dimensions */
84
85 void Setup_Grid();
86 double Do_Step(int parity);
87 void Solve();
88 void Write_Grid();
89 void Clean_Up();
90 void Debug(char *mesg, int terminate);
91 void start_timer();
92 void resume_timer();
93 void stop_timer();
94 void print_timer();
95
96 void start_timer()
97 {
98     if (!timer_on){
99         MPI_Barrier(grid_comm);
100         ticks = clock();
101         wtime = MPI_Wtime();
102         timer_on = 1;
103     }
104 }
105
106 void resume_timer()
107 {
108     if (!timer_on){
109         ticks = clock() - ticks;
110         wtime = MPI_Wtime() - wtime;
111         timer_on = 1;
112     }
113 }
114
115 void stop_timer()
116 {
117     if (timer_on){
118         ticks = clock() - ticks;
119         wtime = MPI_Wtime() - wtime;
120         timer_on = 0;
121     }
122 }
123
124 void print_timer()
125 {
126     if (timer_on){
127         stop_timer();
128         printf("(i) Elapsed Wtime %14.6f s (%5.1f%% CPU)\n", proc_rank, wtime, 100.0 * ticks
129             * (1.0 / CLOCKS_PER_SEC) / wtime);
130         resume_timer();
131     }
132     else{
133         printf("(i) Elapsed Wtime %14.6f s (%5.1f%% CPU)\n", proc_rank, wtime, 100.0 * ticks
134             * (1.0 / CLOCKS_PER_SEC) / wtime);

```

```

133     }
134 }
135
136 void Debug(char *mesg, int terminate)
137 {
138     if (DEBUG || terminate){
139         printf("%s\n", mesg);
140     }
141     if (terminate){
142         exit(1);
143     }
144 }
145
146 void Setup_Proc_Grid(int argc, char **argv){
147     int wrap_around[2];
148     int reorder;
149
150     Debug("My_MPI_Init",0);
151
152     // num of processes
153     MPI_Comm_size(MPI_COMM_WORLD, &P);
154
155     //calculate the number of processes per column and per row for the grid
156     if(argc>2){
157         P_grid[X_DIR] = atoi(argv[1]);
158         P_grid[Y_DIR] = atoi(argv[2]);
159         if(P_grid[X_DIR] * P_grid[Y_DIR] != P){
160             Debug("ERROR Proces grid dimensions do not match with P ", 1);
161         }
162         #ifdef SOR
163         if (argc>3)
164         {
165             // get sor from args
166             sor_omega = atof(argv[3]);
167             printf("Set sor_omega over argv to %1.4f\n", sor_omega);
168         }
169         #endif
170         #ifndef SKIP_EXCHANGE
171         if (argc > 4)
172         {
173             skip_exchange = atoi(argv[4]);
174             printf("Set skip_exchange over argv to %zu\n", skip_exchange);
175         }
176         else{
177             skip_exchange = 1;
178             printf("Set skip_exchange to default value 1\n");
179         }
180         #endif
181     }
182     else{
183         Debug("ERROR Wrong parameter input",1);
184     }
185
186     // Create process topology (2D grid)
187     wrap_around[X_DIR] = 0;
188     wrap_around[Y_DIR] = 0;
189     reorder = 1; //reorder process ranks
190
191     // create grid_comm
192     int ret = MPI_Cart_create(MPI_COMM_WORLD, 2, P_grid, wrap_around, reorder, &grid_comm);
193     if (ret != MPI_SUCCESS){
194         Debug("ERROR: MPI_Cart_create failed",1);
195     }
196     //get new rank and cartesian coords of this proc
197     MPI_Comm_rank(grid_comm, &proc_rank);
198     MPI_Cart_coords(grid_comm, proc_rank, 2, proc_coord);
199     printf("(%) (x,y)=(%i,%i)\n", proc_rank, proc_coord[X_DIR], proc_coord[Y_DIR]);
200     //calc neighbours
201     // MPI_Cart_shift(grid_comm, Y_DIR, 1, &proc_bottom, &proc_top);
202     MPI_Cart_shift(grid_comm, Y_DIR, 1, &proc_top, &proc_bottom);
203     MPI_Cart_shift(grid_comm, X_DIR, 1, &proc_left, &proc_right);
204     printf("(%) top %i, right %i, bottom %i, left %i\n", proc_rank, proc_top,
proc_right, proc_bottom, proc_left);

```

```

205 }
206
207 void Setup_Grid()
208 {
209     int x, y, s;
210     double source_x, source_y, source_val;
211     FILE *f;
212
213     Debug("Setup_Subgrid", 0);
214
215     if(proc_rank == 0){
216         f = fopen("input.dat", "r");
217         if (f == NULL){
218             Debug("Error opening input.dat", 1);
219         }
220         fscanf(f, "nx: %i\n", &gridsize[X_DIR]);
221         fscanf(f, "ny: %i\n", &gridsize[Y_DIR]);
222         fscanf(f, "precision goal: %lf\n", &precision_goal);
223         fscanf(f, "max iterations: %i\n", &max_iter);
224     }
225     MPI_Bcast(&gridsize, 2, MPI_INT, 0, grid_comm);
226     MPI_Bcast(&precision_goal, 1, MPI_DOUBLE, 0, grid_comm);
227     MPI_Bcast(&max_iter, 1, MPI_INT, 0, grid_comm);
228     hx = 1 / (double)gridsize[X_DIR];
229     hy = 1 / (double)gridsize[Y_DIR];
230
231     /* Calculate dimensions of local subgrid */ //! We do that later now!
232     // dim[X_DIR] = gridsize[X_DIR] + 2;
233     // dim[Y_DIR] = gridsize[Y_DIR] + 2;
234
235     //! Step 7
236     int upper_offset[2] = {0,0};
237     // Calculate top left corner coordinates of local grid
238     offset[X_DIR] = gridsize[X_DIR] * proc_coord[X_DIR] / P_grid[X_DIR];
239     offset[Y_DIR] = gridsize[Y_DIR] * proc_coord[Y_DIR] / P_grid[Y_DIR];
240     upper_offset[X_DIR] = gridsize[X_DIR] * (proc_coord[X_DIR] + 1) / P_grid[X_DIR];
241     upper_offset[Y_DIR] = gridsize[Y_DIR] * (proc_coord[Y_DIR] + 1) / P_grid[Y_DIR];
242
243     // dimensions of local grid
244     dim[X_DIR] = upper_offset[X_DIR] - offset[X_DIR];
245     dim[Y_DIR] = upper_offset[Y_DIR] - offset[Y_DIR];
246     // Add space for rows/columns of neighboring grid
247     dim[X_DIR] += 2;
248     dim[Y_DIR] += 2;
249     //! Step 7 end
250
251     /* allocate memory */
252     if ((phi = malloc(dim[X_DIR] * sizeof(*phi))) == NULL){
253         Debug("Setup_Subgrid : malloc(phi) failed", 1);
254     }
255     if ((source = malloc(dim[X_DIR] * sizeof(*source))) == NULL){
256         Debug("Setup_Subgrid : malloc(source) failed", 1);
257     }
258     if ((phi[0] = malloc(dim[Y_DIR] * dim[X_DIR] * sizeof(**phi))) == NULL){
259         Debug("Setup_Subgrid : malloc(*phi) failed", 1);
260     }
261     if ((source[0] = malloc(dim[Y_DIR] * dim[X_DIR] * sizeof(**source))) == NULL){
262         Debug("Setup_Subgrid : malloc(*source) failed", 1);
263     }
264     for (x = 1; x < dim[X_DIR]; x++)
265     {
266         phi[x] = phi[0] + x * dim[Y_DIR];
267         source[x] = source[0] + x * dim[Y_DIR];
268     }
269
270     /* set all values to '0' */
271     for (x = 0; x < dim[X_DIR]; x++){
272         for (y = 0; y < dim[Y_DIR]; y++)
273         {
274             phi[x][y] = 0.0;
275             source[x][y] = 0;
276         }
277     }

```



```

278  /* put sources in field */
279  do{
280      if (proc_rank==0)
281      {
282          s = fscanf(f, "source: %lf %lf %lf\n", &source_x, &source_y, &source_val);
283      }
284      MPI_Bcast(&s, 1, MPI_INT, 0, grid_comm);
285      if (s==3){
286          MPI_Bcast(&source_x, 1, MPI_DOUBLE, 0, grid_comm);
287          MPI_Bcast(&source_y, 1, MPI_DOUBLE, 0, grid_comm);
288          MPI_Bcast(&source_val, 1, MPI_DOUBLE, 0, grid_comm);
289          x = source_x * gridsize[X_DIR];
290          y = source_y * gridsize[Y_DIR];
291          x = x + 1 - offset[X_DIR]; // Step 7 --> local grid transform
292          y = y + 1 - offset[Y_DIR]; // Step 7 --> local grid transform
293          if(x > 0 && x < dim[X_DIR] - 1 && y > 0 && y < dim[Y_DIR] - 1){ // check if in local
grid
294              phi[x][y] = source_val;
295              source[x][y] = 1;
296          }
297      }
298  }
299  while (s==3);
300
301  if(proc_rank==0){
302      fclose(f);
303  }
304 }
305
306 void Setup_MPI_Datatypes()
307 {
308     Debug("Setup_MPI_Datatypes",0);
309
310     // vertical data exchange (Y_Dir)
311     MPI_Type_vector(dim[X_DIR] - 2, 1, dim[Y_DIR], MPI_DOUBLE, &border_type[Y_DIR]);
312     // horizontal data exchange (X_Dir)
313     MPI_Type_vector(dim[Y_DIR] - 2, 1, 1, MPI_DOUBLE, &border_type[X_DIR]);
314
315     MPI_Type_commit(&border_type[Y_DIR]);
316     MPI_Type_commit(&border_type[X_DIR]);
317 }
318
319 int Exchange_Borders()
320 {
321     #ifdef MONITOR_EXCHANGE_BORDERS
322     double start_time, latency_start, latency;
323     double data_size_top, data_size_left;
324     double exchange_time;
325
326     // Measure latency with a small dummy message
327     latency_start = MPI_Wtime();
328     double dummy;
329     MPI_Sendrecv(&dummy, 1, MPI_DOUBLE, proc_top, 0, &dummy, 1, MPI_DOUBLE, proc_bottom, 0,
grid_comm, &status);
330     latency = MPI_Wtime() - latency_start;
331     total_latency += latency;
332
333     // Calculate data sizes
334     data_size_top = dim[X_DIR] * sizeof(double); // Top and bottom rows
335     data_size_left = dim[Y_DIR] * sizeof(double); // Left and right columns
336     double data_transferred = 2 * (data_size_top + data_size_left); // Total data for this
exchange
337     total_data_transferred += data_transferred;
338     #endif
339
340     Debug("Exchange_Borders",0);
341     #ifdef MONITOR_EXCHANGE_BORDERS
342     start_time = MPI_Wtime();
343     #endif
344     // top direction
345     MPI_Sendrecv(&phi[1][1], 1, border_type[Y_DIR], proc_top, 0, &phi[1][dim[Y_DIR] - 1], 1,
border_type[Y_DIR], proc_bottom, 0, grid_comm, &status);
346     // bottom direction

```

```

347 MPI_Sendrecv(&phi[1][dim[Y_DIR] - 2], 1, border_type[Y_DIR], proc_bottom, 0, &phi[1][0],
348 1, border_type[Y_DIR], proc_top, 0, grid_comm, &status);
349 // left direction
350 MPI_Sendrecv(&phi[1][1], 1, border_type[X_DIR], proc_left, 0, &phi[dim[X_DIR]-1][1], 1,
351 border_type[X_DIR], proc_right, 0, grid_comm, &status);
352 // right direction
353 MPI_Sendrecv(&phi[dim[X_DIR]-2][1], 1, border_type[X_DIR], proc_right, 0, &phi[0][1], 1,
354 border_type[X_DIR], proc_left, 0, grid_comm, &status);
355
356 #ifdef MONITOR_EXCHANGE_BORDERS
357 exchange_time = MPI_Wtime() - start_time;
358 total_exchange_time += exchange_time;
359 num_exchanges++;
360 #endif
361 return 1;
362 }
363
364 double Do_Step(int parity)
365 {
366     int x, y;
367     double old_phi, c_ij;
368     double max_err = 0.0;
369
370 #ifdef FAST_DO_STEP_LOOP
371 int start_y;
372 for (x = 1; x < dim[X_DIR] - 1; x++){
373     start_y = ((1 + x + offset[X_DIR] + offset[Y_DIR]) % 2 == parity) ? 1 : 2;
374     for (y = start_y; y < dim[Y_DIR] - 1; y += 2){
375         if (source[x][y] != 1){
376             old_phi = phi[x][y];
377             #ifndef SOR
378             phi[x][y] = (phi[x + 1][y] + phi[x - 1][y] + phi[x][y + 1] + phi[x][y - 1]) *
379 0.25;
380             #endif
381             #ifdef SOR
382             c_ij = (phi[x + 1][y] + phi[x - 1][y] + phi[x][y + 1] + phi[x][y - 1] + hx*hy*
383 source[x][y]) * 0.25 - phi[x][y];
384             phi[x][y] += sor_omega*c_ij;
385             #endif
386             if (max_err < fabs(old_phi - phi[x][y])){
387                 max_err = fabs(old_phi - phi[x][y]);
388             }
389         }
390     }
391 }
392 #endif
393 return max_err;
394 #endif
395
396 #ifndef FAST_DO_STEP_LOOP
397 /* calculate interior of grid */
398 for (x = 1; x < dim[X_DIR] - 1; x++){
399     for (y = 1; y < dim[Y_DIR] - 1; y++){
400         if ((x + offset[X_DIR] + y + offset[Y_DIR]) % 2 == parity && source[x][y] != 1){
401             old_phi = phi[x][y];
402             #ifndef SOR
403             phi[x][y] = (phi[x + 1][y] + phi[x - 1][y] + phi[x][y + 1] + phi[x][y - 1]) *
404 0.25;
405             #endif
406             #ifdef SOR
407             c_ij = (phi[x + 1][y] + phi[x - 1][y] + phi[x][y + 1] + phi[x][y - 1] + hx*hy*
408 source[x][y]) * 0.25 - phi[x][y];
409             phi[x][y] += sor_omega*c_ij;
410             #endif
411             if (max_err < fabs(old_phi - phi[x][y])){
412                 max_err = fabs(old_phi - phi[x][y]);
413             }
414         }
415     }
416 }
417 return max_err;
418 #endif
419 }
420
421 }
422

```

```

413 void Solve()
414 {
415     int count = 0;
416     double delta;
417     double global_delta;
418     double delta1, delta2;
419
420     Debug("Solve", 0);
421
422     /* give global_delta a higher value then precision_goal */
423     global_delta = 2 * precision_goal;
424
425     while (global_delta > precision_goal && count < max_iter)
426     {
427         Debug("Do_Step 0", 0);
428         delta1 = Do_Step(0);
429         #ifdef SKIP_EXCHANGE
430         if (count % skip_exchange == 0 && Exchange_Borders()) // use short circuit evaluation
431             #endif
432         #ifndef SKIP_EXCHANGE
433         Exchange_Borders();
434         #endif
435         Debug("Do_Step 1", 0);
436         delta2 = Do_Step(1);
437         #ifdef SKIP_EXCHANGE
438         if (count % skip_exchange == 0 && Exchange_Borders())
439             #endif
440         #ifndef SKIP_EXCHANGE
441         Exchange_Borders();
442         #endif
443         delta = max(delta1, delta2);
444         #ifdef MONITOR_ALLREDUCE
445         double time_ = MPI_Wtime();
446         #endif
447         #ifdef ALLREDUCE_COUNT
448         if (count % ALLREDUCE_COUNT == 0){
449             MPI_Allreduce(&delta, &global_delta, 1, MPI_DOUBLE, MPI_MAX, grid_comm);
450         }
451         #endif
452         #ifndef ALLREDUCE_COUNT
453         MPI_Allreduce(&delta, &global_delta, 1, MPI_DOUBLE, MPI_MAX, grid_comm);
454         #endif
455         #ifdef MONITOR_ALLREDUCE
456         all_reduce_time += MPI_Wtime() - time_;
457         #endif
458         #ifdef MONITOR_ERROR
459         if (proc_rank == 0)
460         {
461             errors[count] = global_delta;
462         }
463         #endif
464         count++;
465     }
466
467     printf("(%i) Number of iterations : %i\n", proc_rank, count);
468     #ifdef MONITOR_ALLREDUCE
469     printf("(%i) Allreduce time: %14.6f\n", proc_rank, all_reduce_time);
470     #endif
471     #ifdef MONITOR_EXCHANGE_BORDERS
472     printf("(%i) Exchange time: %14.6f\n", proc_rank, total_exchange_time);
473     #endif
474 }
475
476 double* get_Global_Grid()
477 {
478     Debug("get_Global_Grid", 0);
479     ///!! DEBUG only
480     for (size_t i = 0; i < dim[X_DIR]; i++)
481     {
482         for (size_t j = 0; j < dim[Y_DIR]; j++)
483         {
484             phi[i][j] = proc_rank;
485         }
486     }
487 }

```

```

486 }
487
488
489 // only process 0 needs to store all data!
490 double* global_phi = NULL;
491 if (proc_rank == 0) {
492     global_phi = malloc(gridsize[X_DIR] * gridsz[e[Y_DIR] * sizeof(double));
493     if (global_phi == NULL) {
494         Debug("get_Global_Grid : malloc(global_phi) failed", 1);
495     }
496 }
497
498 // copy own part into buffer - flatten!
499 size_t buf_size = (dim[X_DIR] - 2) * (dim[Y_DIR] - 2) * sizeof(double);
500 double* local_phi = malloc(buf_size);
501 int idx = 0;
502 for (int x = 1; x < dim[X_DIR] - 1; x++) {
503     for (int y = 1; y < dim[Y_DIR] - 1; y++) {
504         local_phi[idx++] = phi[x][y];
505     }
506 }
507 printf("I'm proc %d and i have a buffer of size %zu\n", proc_rank, buf_size);
508
509
510 // only proc 0 needs sendcounts and displacements for the gather operation
511 int* sendcounts = NULL;
512 int* displs = NULL;
513 if (proc_rank == 0) {
514     sendcounts = malloc(P * sizeof(int));
515     displs = malloc(P * sizeof(int));
516
517     // size and offset of different subgrids
518     //!! Note that this only works if every process has the same subgrid
519     if (gridsize[X_DIR] % P_grid[X_DIR] != 0 || gridsz[e[Y_DIR] % P_grid[Y_DIR] != 0)
520     {
521         Debug("!!!A grid dimension is not a multiple of the P_grid in this direction!", 1)
522     }
523
524     int subgrid_width = gridsz[e[X_DIR] / P_grid[X_DIR];
525     int subgrid_height = gridsz[e[Y_DIR] / P_grid[Y_DIR];
526     for (int px = 0; px < P_grid[X_DIR]; px++) {
527         for (int py = 0; py < P_grid[Y_DIR]; py++) {
528             int rank = px * P_grid[Y_DIR] + py;
529             sendcounts[rank] = subgrid_width * subgrid_height;
530             displs[rank] = (px * subgrid_width * gridsz[e[Y_DIR]) + (py * subgrid_height);
531         }
532     }
533 }
534 Debug("get_Global_Grid : MPI_Gatherv", 0);
535 //!! TODO this Gatherv does something wrong - all local grids are alright!!!
536 MPI_Gatherv(local_phi, (dim[X_DIR] - 2) * (dim[Y_DIR] - 2), MPI_DOUBLE, global_phi,
537             sendcounts, displs, MPI_DOUBLE, 0, MPI_COMM_WORLD);
538
539 free(local_phi);
540 if (proc_rank == 0) {
541     free(sendcounts);
542     free(displs);
543 }
544
545 return global_phi;
546 }
547
548 void Write_Grid_global(){
549     int x, y;
550     FILE *f;
551     char filename[40]; //seems dangerous to use a static buffer but let's go with the steps
552     sprintf(filename, "output_MPI_global_%i.dat", proc_rank);
553     if ((f = fopen(filename, "w")) == NULL){
554         Debug("Write_Grid : fopen failed", 1);
555     }
556     Debug("Write_Grid", 0);

```

```

557
558     for (x = 1; x < dim[X_DIR]-1; x++){
559         for (y = 1; y < dim[Y_DIR]-1; y++){
560             int x_glob = x + offset[X_DIR];
561             int y_glob = y + offset[Y_DIR];
562             fprintf(f, "%i %i %f\n", x_glob, y_glob, phi[x][y]);
563         }
564     }
565     fclose(f);
566 }
567
568 void Write_Grid()
569 {
570     double* global_phi = get_Global_Grid();
571     if(proc_rank != 0){
572         assert (global_phi == NULL);
573         return;
574     }
575     int x, y;
576     FILE *f;
577     char filename[40]; //seems dangerous to use a static buffer but let's go with the steps
578     sprintf(filename, "output_MPI%i.dat", proc_rank);
579     if ((f = fopen(filename, "w")) == NULL){
580         Debug("Write_Grid : fopen failed", 1);
581     }
582
583     Debug("Write_Grid", 0);
584
585     for (x = 0; x < gridsize[X_DIR]; x++){
586         for (y = 0; y < gridsize[Y_DIR]; y++){
587             fprintf(f, "%i %i %f\n", x+1, y+1, global_phi[x*gridsize[Y_DIR] + y]);
588         }
589     }
590     fclose(f);
591     free(global_phi);
592 }
593
594 void Clean_Up()
595 {
596     Debug("Clean_Up", 0);
597
598     free(phi[0]);
599     free(phi);
600     free(source[0]);
601     free(source);
602     #ifdef MONITOR_ERROR
603     free(errors);
604     #endif
605 }
606 void setup_error_monitor(){
607     if (proc_rank != 0)
608     {
609         return;
610     }
611
612     errors = malloc(sizeof(double)*max_iter);
613 }
614 void write_errors(){
615     if(proc_rank != 0){
616         return;
617     }
618     FILE *f;
619     char filename[40]; //seems dangerous to use a static buffer but let's go with the steps
620     sprintf(filename, "errors_MPI.dat");
621     if ((f = fopen(filename, "w")) == NULL){
622         Debug("Write_Errors : fopen failed", 1);
623     }
624
625     Debug("Write_Errors", 0);
626
627     for (size_t i = 0; i < max_iter; ++i)
628     {
629         fprintf(f, "%f\n", errors[i]);

```

```

630     }
631     fclose(f);
632 }
633
634 void Print_Aggregated_Metrics()
635 {
636     #ifdef MONITOR_EXCHANGE_BORDERS
637     if (num_exchanges > 0) {
638         double avg_exchange_time = total_exchange_time / num_exchanges;
639         double avg_latency = total_latency / num_exchanges;
640         double avg_bandwidth = total_data_transferred / total_exchange_time;
641
642         printf("\n--- Aggregated Metrics ---\n");
643         printf("Total Exchanges: %d\n", num_exchanges);
644         printf("Total Data Transferred: %.2f bytes\n", total_data_transferred);
645         printf("Total Exchange Time: %.9f s\n", total_exchange_time);
646         printf("Average Exchange Time per Call: %.9f s\n", avg_exchange_time);
647         printf("Average Latency per Call: %.9f s\n", avg_latency);
648         printf("Average Bandwidth: %.2f bytes/s\n", avg_bandwidth);
649     } else {
650         printf("No exchanges recorded.\n");
651     }
652     #endif
653 }
654
655 int main(int argc, char **argv)
656 {
657     MPI_Init(&argc, &argv);
658     Setup_Proc_Grid(argc, argv); // was earlier MPI_Comm_rank(MPI_COMM_WORLD, &proc_rank);
659     start_timer();
660
661     Setup_Grid();
662     Setup_MPI_Datatypes();
663
664     #ifdef SOR
665     if (proc_rank == 0)
666     {
667         printf("SOR using omega: %.5f\n", sor_omega);
668     }
669     #endif
670     #ifdef MONITOR_ERROR
671     setup_error_monitor();
672     #endif
673
674     Solve();
675     #ifdef MONITOR_ERROR
676     write_errors();
677     #endif
678     // Write_Grid();
679     Write_Grid_global();
680     Print_Aggregated_Metrics();
681     print_timer();
682
683     Clean_Up();
684     MPI_Finalize();
685     return 0;
686 }

```

Script used for [subsubsection 1.2.3](#):

```

1  #!/bin/bash
2  #SBATCH --job-name="scaling_123"
3  #SBATCH --time=00:03:00
4  #SBATCH --ntasks=4
5  #SBATCH --cpus-per-task=1
6  #SBATCH --partition=compute
7  #SBATCH --mem=2GB # Increased memory
8  #SBATCH --account=Education-EEMCS-Courses-WI4049TU
9
10 if [ -z "$1" ] || [ -z "$2" ]; then
11     echo "Usage: $0 <topology_x> <topology_y>"
12     exit 1
13 fi
14

```

```

15 # Move to the src directory
16 cd ../src || { echo "Error: ../src directory not found"; exit 1; }
17 basefolder="123/${1}_${2}"
18 mkdir -p ../scripts/output/$basefolder || { echo "Error creating output directory"; exit 1; }
19
20 # Define maximum iterations and grid sizes
21 maxiters=("500" "1000" "2000")
22 #grids=("50 50" "100 100" "200 200")
23 grids=("1600 1600" "3200 3200")
24
25 for grid in "${grids[@]"; do
26     nx=$(echo $grid | cut -d' ' -f1)
27     ny=$(echo $grid | cut -d' ' -f2)
28
29     mkdir -p ../scripts/output/$basefolder/${nx}x${ny} || { echo "Error creating grid
    directory"; exit 1; }
30
31     for maxiter in "${maxiters[@]"; do
32         python3 -c "
33 import util
34 util.update_input_file(nx=$nx, ny=$ny, precision_goal=0.000000000000001, max_iter=$maxiter)
35 " || { echo "Python script failed for nx=$nx, ny=$ny, max_iter=$maxiter"; exit 1; }
36
37         echo "Running with nx=$nx, ny=$ny, maxiter=$maxiter"
38         srun ./MPI_Poisson.out $1 $2 1.95 > ../scripts/output/$basefolder/${nx}x${ny}/
    maxiter_${maxiter}.txt || {
39             echo "srun failed for nx=$nx, ny=$ny, max_iter=$maxiter"; exit 1;
40         }
41         echo "Finished maxiter=$maxiter for grid ${nx}x${ny}"
42     done
43 done
44
45 python3 -c "
46 import util
47 util.reset_input_file()
48 " || { echo "Error resetting input file"; exit 1; }
49
50 echo "Job completed successfully."

```