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Dipartimento di Ingegneria Elettrica e delle Tecnologie  
dell'Informazione



## Parallel and Distributed Computing submissions

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# Chapter 1

## Prolusion

### 1.1 Goal

A comprehensive overview of a Parallel and Distributed Computing project developed in `C` is presented. The project consists of multiple modules delivered in incremental phases.

Every algorithm is implemented within the **Single Program Multiple Data** parallel model.

This report is not intended as a user guide, but rather aims to describe the project's exploration of Parallel and Distributed Computing techniques, leveraging High Performance Computing in certain instances.

### 1.2 Environment

The project was entirely developed on macOS with the help of Xcode IDE. Naturally, this will mainly impact the build process.

### 1.3 Project directory layout

The structure of the project's root directory is outlined below.

```
parallel-distributed-computing/
├── common/
├── hpc/
│   ├── gemm/
│   ├── matmatblock/
│   ├── matmatdist/
│   └── matmatthread/
├── laplace/
├── maxsum/
├── ringsum/
└── parallel-distributed-computing.entitlements
```

`common` package serves as a library of utility functions designed to support and be reused by various modules across the project.

The remaining directories represent the individual project modules, which constitute the deliverables of the project. Within each module, the directory structure follows a standard format:

```
<module>/
├── build/
│   ├── deploy-cluster.pbs
│   └── Makefile
├── src/
│   ├── <module>/
│   │   ├── <module>.c
│   │   └── <module>.h
│   └── main.c
├── config.sh
└── run.sh
```

Most parts of the `main.c` files are provided by the project supervisor.

## 1.4 Build

The project was primarily compiled using the Clang compiler.

The build process was carried out either through the `Makefile` (some of which support compilers other than Clang) or via Xcode.

Regardless of the build process, every module of the project was compiled with:

- `-O3` optimization flag to maximize performance.

### 1.4.1 Libraries

The following are the dynamically linked libraries integrated into the project.

- `math.h`
- `mpi.h`
- `omp.h`
- `stdio.h`
- `stdbool.h`
- `stdlib.h`
- `sys/time.h`
- `unistd.h`

### 1.4.2 Xcode

When it came to build with the Xcode, the development process adhered to the workflow and conventions defined by the chosen IDE, leveraging its built-in tools and features to organize and manage the project. Particularly, this includes:

- Xcode targets
- Xcode schemes
- Xcode `.entitlements` file

## 1.5 Run

### 1.5.1 Network Requirement

Running an MPI module with no internet connection, makes the following error occur:

```
[Giulianos-MacBook-Pro.local:05355] ptl_tool: problems getting address for  
index 0 (kernel index -1)
```

```
-----  
The PMIx server's listener thread failed to start. We cannot continue.  
-----
```

```
Program ended with exit code: 213
```



# Chapter 2

## Maxsum

### 2.1 Problem at a glance

Given a matrix, the objective is to identify the row with the maximum squared root sum.

### 2.2 Algorithm

#### 2.2.1 Parallelization

The algorithm employs a multithreaded approach, each thread can compute independently its local maximum squared root sum. Once a thread has completed its task, a synchronization is needed with the other threads in order to determine the overall maximum squared root sum.

#### 2.2.2 Pseudocode

**Input:**

- global matrix
- number of thread(s) for parallel execution

**Steps:**

1. Compute the indices of the global matrix corresponding to the local matrix of the thread.
2. Compute the local maximum squared root sum.
3. Perform a thread-safe update of the overall maximum squared root sum variable.

---

**Algorithm 1:** Maxsum
 

---

**Input** :  $matrix, numThreads$   
**Output:**  $maxSumOverall$

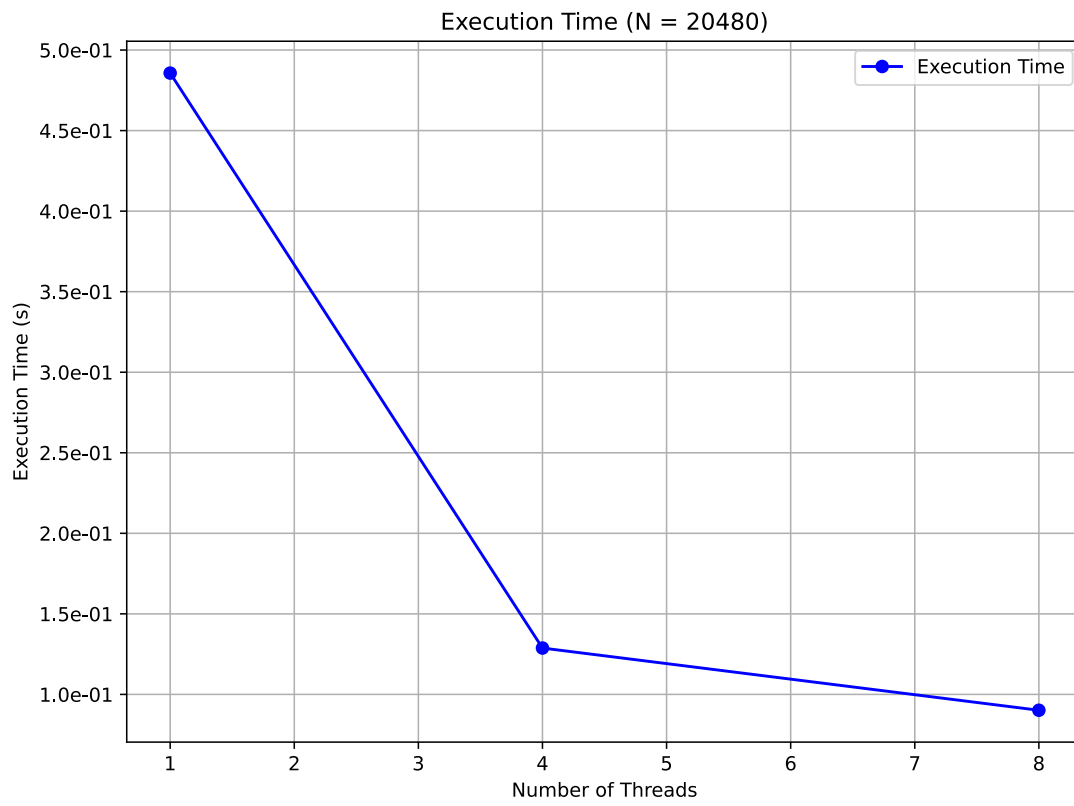
```

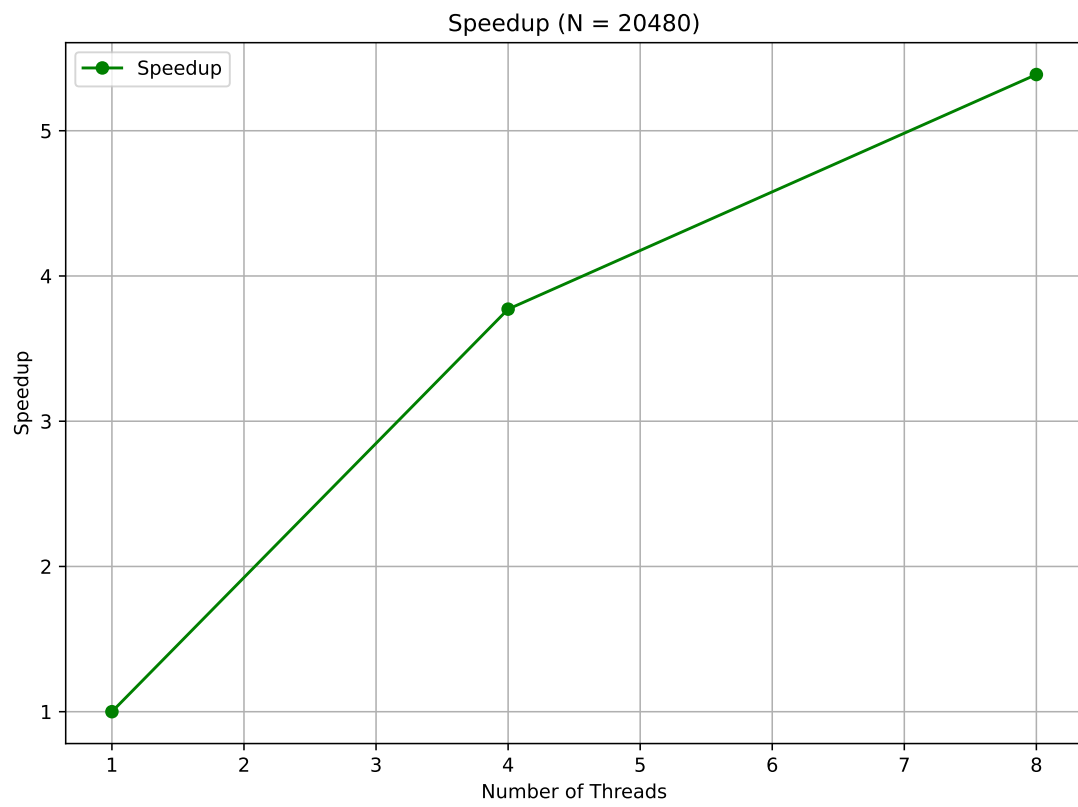
STARTTHREADS(numThreads)
 $startRow \leftarrow getStartingRowIndex(threadRank)$ 
 $endRow \leftarrow getEndingRowIndex(threadRank)$ 

for  $currRow \leftarrow startRow$  to  $endRow$  do
     $currSum \leftarrow squareRootSum(currRow)$ 
    if  $currSum > maxSum$  then
        |  $maxSum \leftarrow currSum$ 
    end
    CRITICAL
    if  $maxSumOverall < maxSum$  then
        |  $maxSumOverall \leftarrow maxSum$ 
    end
end
  
```

---

### 2.2.3 Performance Analysis









# Chapter 3

## Laplace

### 3.1 Problem at a glance

Given a matrix and a number of iterations, the objective is to compute the discrete Laplacian of the matrix over the specified iterations.

### 3.2 Algorithm

#### 3.2.1 Parallelization

The algorithm employs a multiprocess approach. A synchronization between processes is needed from the outset.

The global matrix is distributed among the processes, harnessing the advantages of parallelism, representing a spatial advantage other than a temporal one.

The algorithm comes in two versions: in the first one, communication between processes takes place in a classic way where sending a message “blocks” the execution and will resume it when a reception happens; in the second one, sending a message won’t “block” the execution, instead, execution is blocked when the expected receive message needs to be accessed.

Clearly, the second version is expected to be slightly faster than the other one.

### 3.2.2 Pseudocode

**Input:**

- local process matrix
- number of iteration(s)

**Steps:**

For each iteration:

1. If  $P_i \neq P_1$  Then
  - (a) Send to  $P_{i-1}$  the local first row
  - (b) Receive from  $P_{i-1}$  its last row
2. If  $P_i \neq P_n$  Then
  - (a) Receive from  $P_{i+1}$  its first row
  - (b) Send to  $P_{i+1}$  the local last row
3. Compute the Laplacian considering only the local inner matrix, the matrix boundaries are left out.
4. If  $P_i \neq P_1$ 
  - (a) Compute the Laplacian considering only the local top row, with the auxiliary last row from the previous process.
5. If  $P_i \neq P_n$ 
  - (a) Compute the Laplacian considering only the local bottom row, with the auxiliary first row from the next process.

An auxiliary matrix is involved to facilitate the computation.



---

**Algorithm 2:** Laplace
 

---

**Input** : *matrix, iterations***Output:** *outputMatrix*

```

for iter  $\leftarrow$  1 to iterations do
  if  $P_i \neq P_1$  then
    SEND( $P_{i-1}$ , localFirstRow)
    RECV( $P_{i-1}$ , receivedLastRow)
  end
  if  $P_i \neq P_n$  then
    RECV( $P_{i+1}$ , receivedFirstRow)
    SEND( $P_{i+1}$ , localLastRow)
  end
  laplacian(inner(matrix))
  if  $P_i \neq P_1$  then
    laplacian(matrix, receivedLastRow)
  end
  if  $P_i \neq P_n$  then
    laplacian(matrix, receivedFirstRow)
  end
  copy(outputMatrix, inner(matrix))
  if  $P_i \neq P_1$  then
    copy(outputMatrix, lastRow(matrix))
  end
  if  $P_i \neq P_n$  then
    copy(outputMatrix, firstRow(matrix))
  end
end

```

---

---

**Algorithm 3:** Laplace non blocking
 

---

**Input :** *matrix, iterations***Output:** *outputMatrix*

```

for iter  $\leftarrow$  1 to iterations do
  if  $P_i \neq P_1$  then
    IMMEDIATE_SEND( $P_{i-1}$ , localFirstRow, SEND_NEXTTOPREV)
    IMMEDIATE_RECV( $P_{i-1}$ , receivedLastRow, RECV_PREVTONEXT)
  end
  if  $P_i \neq P_n$  then
    IMMEDIATE_RECV( $P_{i+1}$ , receivedFirstRow, RECV_NEXTTOPREV)
    IMMEDIATE_SEND( $P_{i+1}$ , localLastRow, SEND_PREVTONEXT)
  end
  laplacian(inner(matrix))
  if  $P_i \neq P_1$  then
    WAIT RECV_PREVTONEXT
    laplacian(matrix, receivedLastRow)
  end
  if  $P_i \neq P_n$  then
    WAIT RECV_NEXTTOPREV
    laplacian(matrix, receivedFirstRow)
  end
  copy(outputMatrix, inner(matrix))
  if  $P_i \neq P_1$  then
    WAIT SEND_NEXTTOPREV
    copy(outputMatrix, firstRow(matrix))
  end
  if  $P_i \neq P_n$  then
    WAIT SEND_PREVTONEXT
    copy(outputMatrix, lastRow(matrix))
  end
end

```

---

### 3.2.3 Performance Analysis

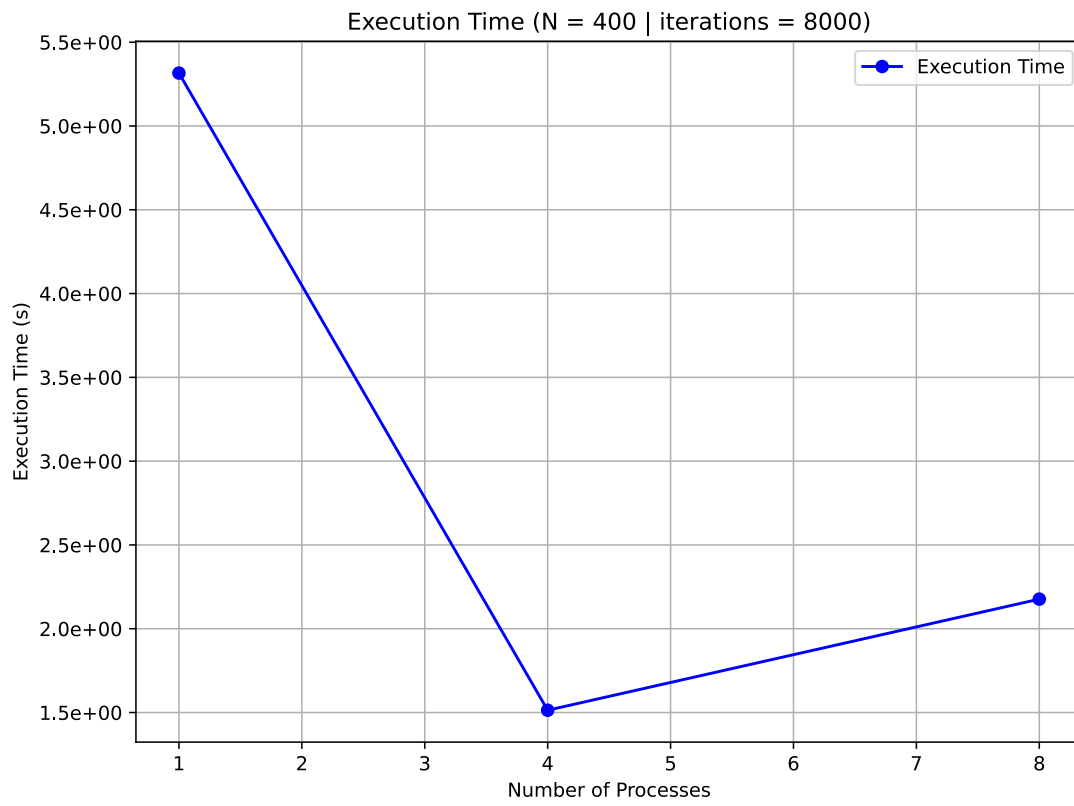


Figure 3.1: *Laplace algorithm execution time*

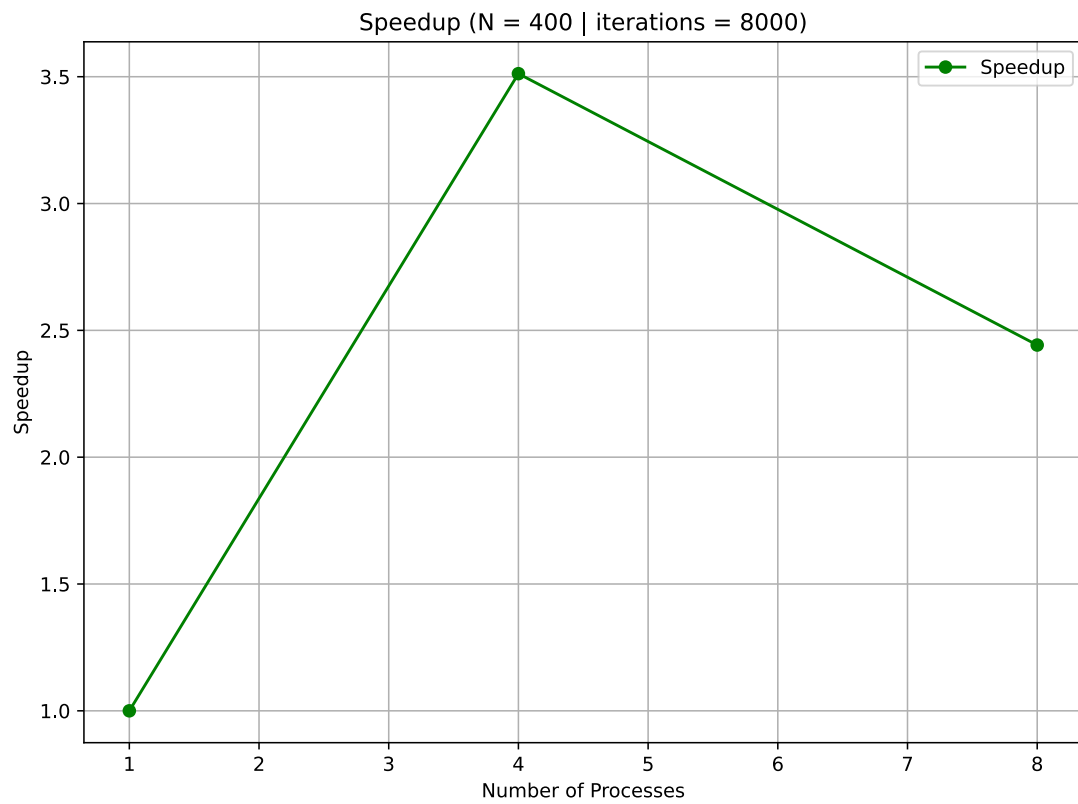


Figure 3.2: *Laplace algorithm speedup*

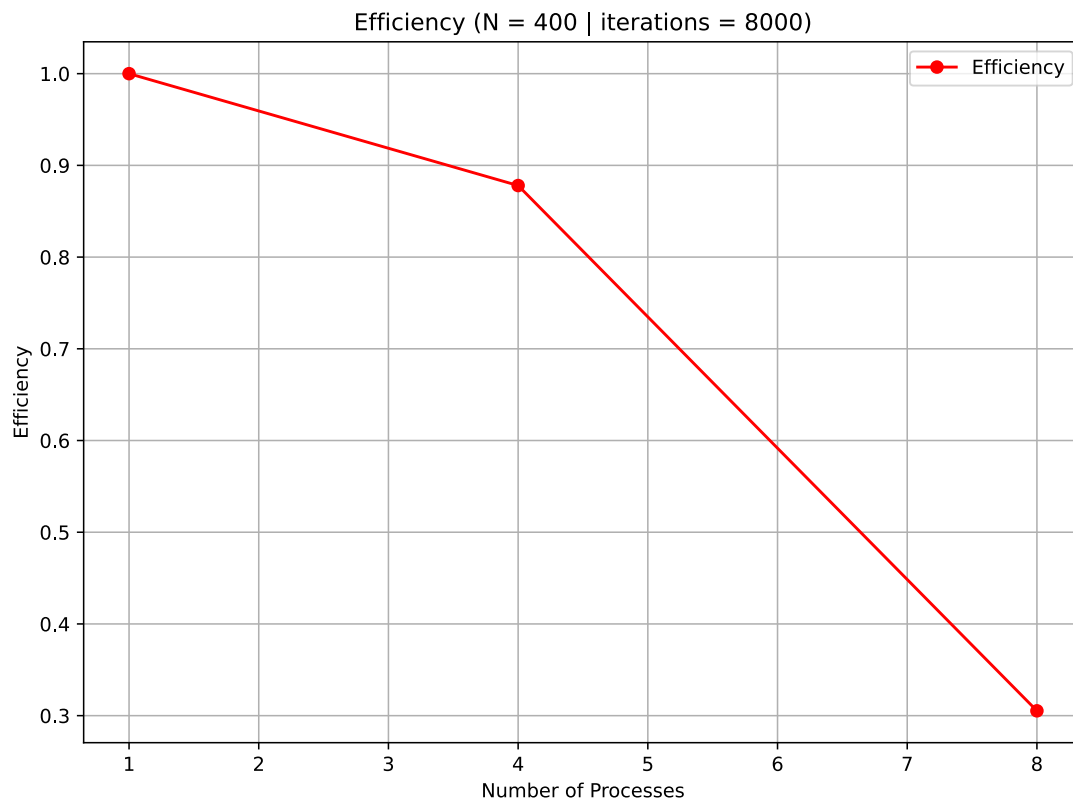


Figure 3.3: *Laplace algorithm efficiency*

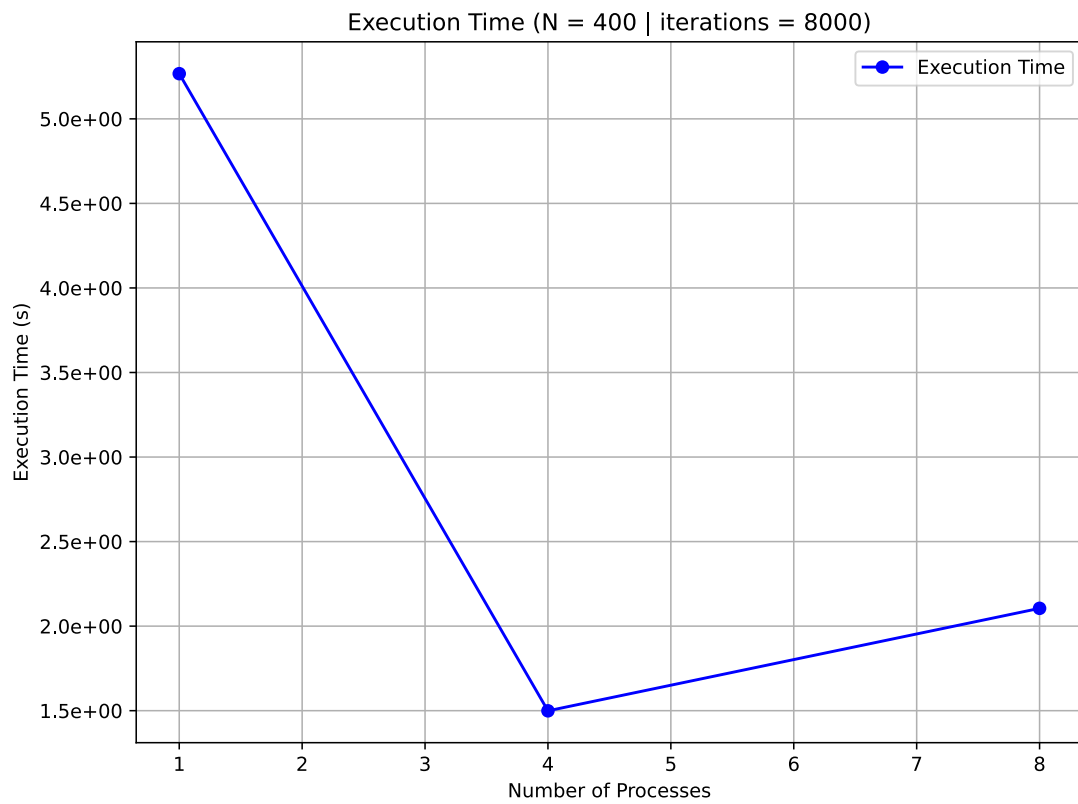


Figure 3.4: *Laplace non-blocking algorithm execution time*

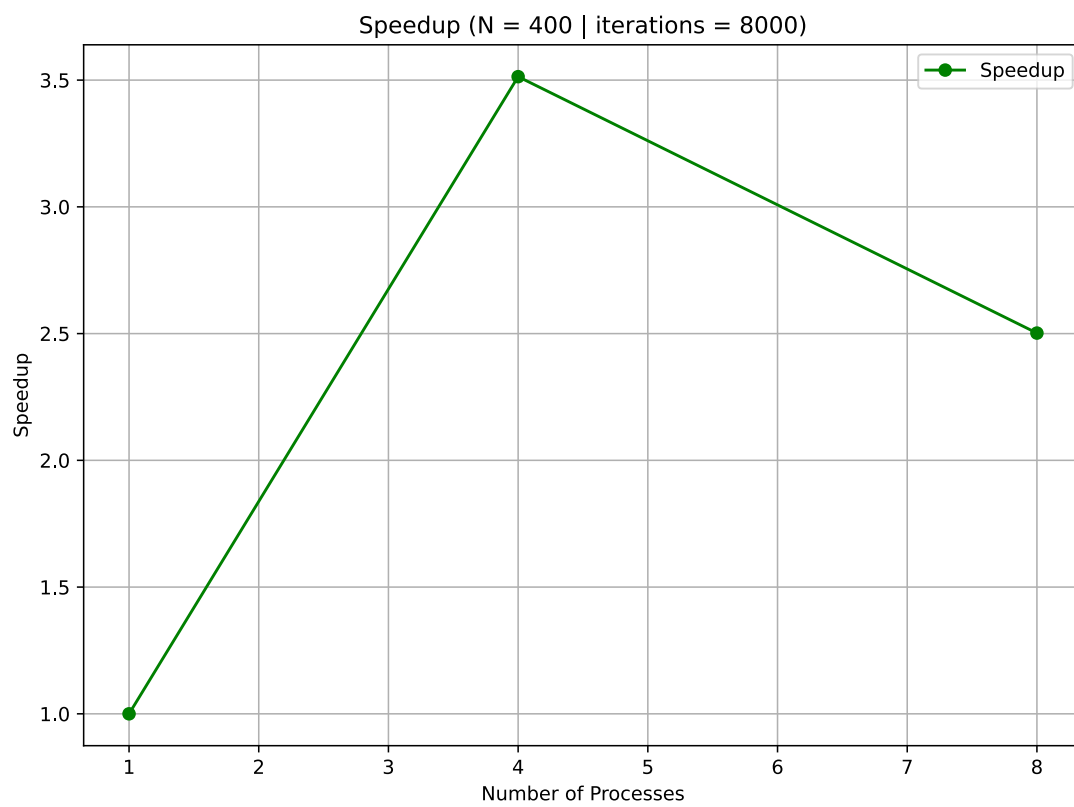


Figure 3.5: *Laplace non-blocking algorithm speedup*

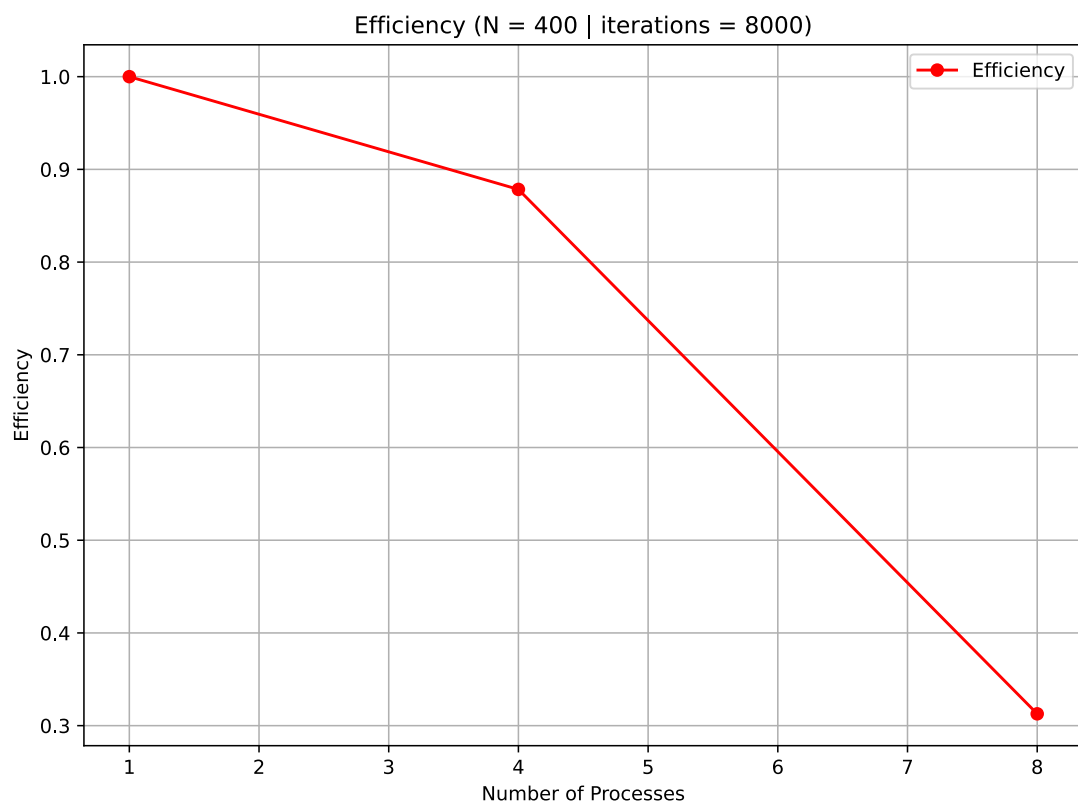


Figure 3.6: *Laplace non-blocking algorithm efficiency*



# Chapter 4

## MatMatikj

### 4.1 Problem at a glance

Given matrices  $A$ ,  $B$ ,  $C$ , the goal is to compute the GeMM operation  $C = C + A * B$  by iterating over the matrices while leveraging the ikj ordering of indices. This increase cache memory utilization.

### 4.2 Algorithm

#### 4.2.1 Parallelization

No parallelization is planned for this algorithm.

#### 4.2.2 Pseudocode

**Input:**

- $A$ ,  $B$ ,  $C$  matrices

**Steps:**

- iterate over the three input matrices in an ‘ikj’ order
- compute  $C = C + A * B$  for each step

---

**Algorithm 4:** MatMatikj

---

**Input** :  $A^{N1 \times N2}$ ,  $B^{N2 \times N3}$ ,  $C^{N1 \times N3}$

**Output:**  $C^{N1 \times N3}$

```
for  $i \leftarrow 1$  to  $N1$  do
  for  $k \leftarrow 1$  to  $N2$  do
    for  $j \leftarrow 1$  to  $N3$  do
       $C(i, j) = C(i, j) + A(i, k) + B(k, j)$ 
    end
  end
end
```

---



## Chapter 5

# MatMatBlock

### 5.1 Problem at a glance

Given matrices  $A$ ,  $B$ ,  $C$ , the goal is to compute the GeMM operation  $C = C + A * B$  in ‘blocks’, by leveraging the best performance of MatMatikj algorithm.

### 5.2 Algorithm

#### 5.2.1 Parallelization

No parallelization is planned for this algorithm.

#### 5.2.2 Pseudocode

**Input:**

- $A$ ,  $B$ ,  $C$  matrices
- $blockN1$ ,  $blockN2$ ,  $blockN3$  block dimensions of  $N1$ ,  $N2$ ,  $N3$

**Steps:**

- iterate over the three input matrices in steps of  $blockN1$ ,  $blockN2$ ,  $blockN3$
- execute MatMatikj algorithm for each step

---

**Algorithm 5:** MatMatblock
 

---

**Input** :  $A^{N1 \times N2}$ ,  $B^{N2 \times N3}$ ,  $C^{N1 \times N3}$ ,  $blockN1$ ,  $blockN2$ ,  $blockN3$   
**Output:**  $C^{N1 \times N3}$

```

for  $ii \leftarrow 1$  to  $N1$  by  $blockN1$  do
  for  $jj \leftarrow 1$  to  $N3$  by  $blockN3$  do
    for  $kk \leftarrow 1$  to  $N2$  by  $blockN2$  do
       $MatMatikj(\mathbf{A}(ii, kk), \mathbf{B}(kk, jj), \mathbf{C}(ii, jj))$ 
    end
  end
end

```

---

## Chapter 6

# MatMatThread

### 6.1 Problem at a glance

Given matrices  $A$ ,  $B$ ,  $C$ , the goal is to perform a multithreaded MatMatBlock algorithm to compute  $C = C + AB$ .

### 6.2 Algorithm

#### 6.2.1 Parallelization

#### 6.2.2 Pseudocode



## Chapter 7

# Further Algorithms

The project includes additional modules not part of the presented submissions, such as two parallel summation algorithms: **Ringsum** and **Cascadesum**.

The latter is more efficient with respect to time complexity, requiring only  $\log_2(P)$  steps for message passing, as opposed to  $P - 1$ , where  $P$  denotes the number of processes involved.









# Acronyms

**GeMM** General Matrix Multiply 21, 23

**HPC** High Performance Computing 1

**IDE** Integrated Development Environment 1, 3

**MPI** Message Passing Interface 4

**PDC** Parallel and Distributed Computing 1

**SPMD** Single Program Multiple Data 1