

# Sri Lanka Institute of Information Technology



## SE3082 - Parallel Computing Assignment 03

Student ID	Name
IT23190016	M.S. SAMARASINGHE

BSc (Hons) in Computer Science

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## Serial Implementation/Baseline

```
M /c/Users/ASUS/Desktop/PC_Assignment_03
ASUS@LAPTOP-500RALBN MINGW64 /c/Users/ASUS/Desktop/PC_Assignment_03
$ ls
PC_Assingment3  cuda_code.cu      mpi_code.c      openmp_code.c    serial_code.c
a.exe           cuda_program.exe  mpi_code.exe    openmp_code.exe

ASUS@LAPTOP-500RALBN MINGW64 /c/Users/ASUS/Desktop/PC_Assignment_03
$ gcc serial_code.c -o serial_code -lm

ASUS@LAPTOP-500RALBN MINGW64 /c/Users/ASUS/Desktop/PC_Assignment_03
$ ./serial_code
Serial execution:
  Threads used      : 1
  Execution time    : 5.512000 seconds

ASUS@LAPTOP-500RALBN MINGW64 /c/Users/ASUS/Desktop/PC_Assignment_03
$
```

- This execution time of 5.512 seconds, measured in a single-threaded environment, will serve as the baseline for evaluating speedup performance across different parallel computing approaches, including OpenMP, MPI, and CUDA.

## Part A: Parallel Implementations

- Repository: [it23190016/PC\\_Assingment3](https://github.com/it23190016/PC_Assingment3)
- This repository contains the implementations of parallel computing approaches for the assignment, including OpenMP, MPI, and CUDA versions. Each implementation is designed to evaluate performance improvements over a single-threaded baseline through efficient utilization of multiple cores or GPUs.

## Part C: Documentation and Analysis

### 1. Parallelization Strategies

#### A. OpenMP Strategy

OpenMP uses shared memory parallelism through compiler directives. The ***#pragma omp parallel for private(j, k)*** directive automatically distributes loop iterations among CPU threads. All threads share the same memory space, eliminating communication overhead. The runtime system handles thread creation and synchronization barriers automatically.

##### Key Features:

- Automatic work distribution via compiler directives
- Shared memory model with implicit synchronization
- Minimal code changes required
- Limited to CPU cores

#### B. MPI Strategy

MPI employs distributed memory parallelism by decomposing the problem across processes. Each process computes assigned rows using ***rows\_per\_proc = N / size***. Matrix B is distributed using ***MPI\_Bcast*** since processes have separate memory spaces. This approach scales across multiple compute nodes.

##### Key Features:

- Row-wise domain decomposition
- Explicit message passing for data distribution
- Independent memory spaces per process
- Scalable across multiple nodes

#### C. CUDA Strategy

CUDA leverages GPU's massive parallel architecture using a 2D thread grid. Each thread computes one matrix element using unique thread indices. Explicit memory management transfers data between host and device. Different thread block configurations (1×1 to 32×32) optimize performance.

##### Key Features:

- Massive parallelism with thousands of threads
- 2D thread grid mapping to matrix elements
- Explicit host-device memory management
- Configurable thread block sizes

## 2. Runtime Configurations

### A. Hardware Specifications

- **System:** ASUS TUF Gaming F15 FX507ZU4 (x64-based PC)
- **Processor:** 12th Gen Intel Core i7-12700H @ 2300 MHz
  - 14 physical cores (6 P-cores + 8 E-cores), 20 logical threads
- **Memory:** 16 GB RAM, 32 GB Page File
- **GPU:** NVIDIA GeForce RTX 4050 Laptop GPU (6 GB GDDR6)
  - CUDA Version: 13.0, Driver: 581.29
  - Max threads per block: 1024, Compute Capability: 8.9

### B. Software Environment

- **Operating System:** Microsoft Windows 11 Home (Build 26100)
- **Compilers and Libraries:**
  - **Serial:** GCC compiler, C standard library
  - **OpenMP:** GCC with -fopenmp flag, omp.h library
  - **MPI:** MPICC wrapper, MPICH implementation
  - **CUDA:** NVCC compiler (v13.0), CUDA runtime API
- **Development Environment:** MSYS2 MinGW64 (OpenMP/MPI), Windows CMD (CUDA)

### C. Configuration Parameters by Implementation

#### 1. Serial Implementation

- Matrix Size: 1000×1000 (N = 1000)
- Compilation: gcc -o serial\_code.exe serial\_code.c
- Timing: clock() function from time.h

#### 2. OpenMP Implementation

- Matrix Size: 1000×1000 (N = 1000)
- Thread Configurations: 1, 2, 4, 8, 16, 20 threads
- Compilation: gcc -fopenmp -o openmp\_code.exe openmp\_code.c
- Environment Variable: export OMP\_NUM\_THREADS=<value>
- Timing: omp\_get\_wtime() function

### 3. MPI Implementation

- Matrix Size: 1000×1000 (N = 1000)
- Process Configurations: 1, 2, 4, 8, 16 processes
- Compilation: `mpicc -o mpi_code.exe mpi_code.c`
- Execution: `mpiexec -np <processes> ./mpi_code.exe`
- Timing: `MPI_Wtime()` function

### 4. CUDA Implementation

- Matrix Size: 1000×1000 (N = 1000)
- Thread Block Configurations: 1×1, 2×2, 4×4, 8×8, 16×16, 32×32
- Compilation: `nvcc cuda_code.cu -o cuda_code.exe`
- Timing: `clock()` with `cudaDeviceSynchronize()`

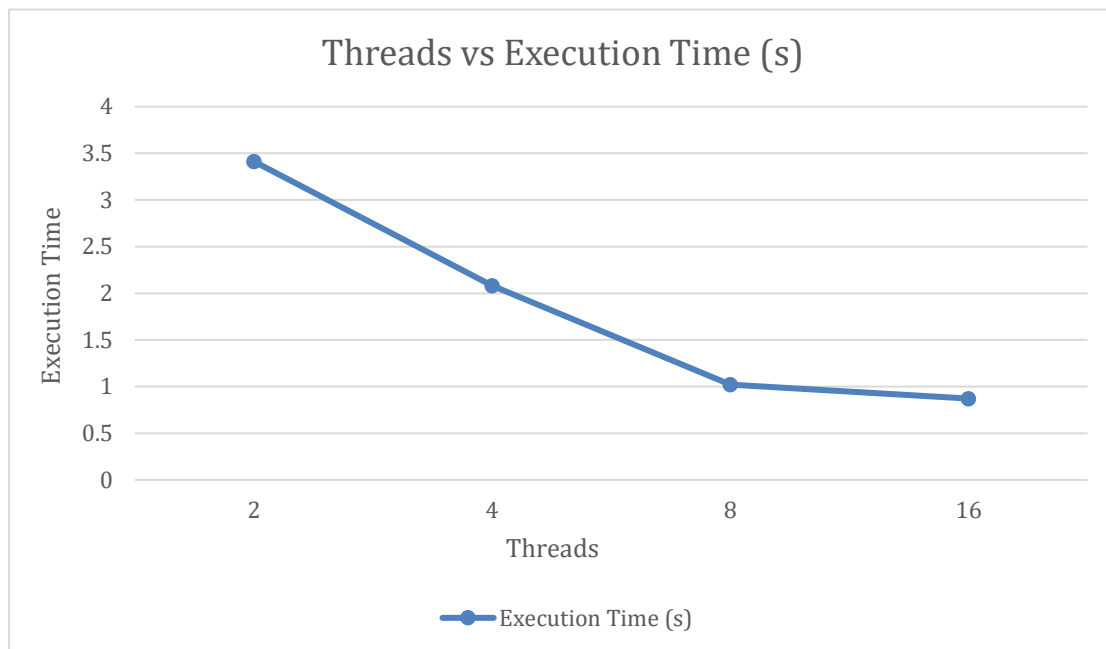
### 3. Performance Analysis

#### A. OpenMP Evaluation

#### Graphs

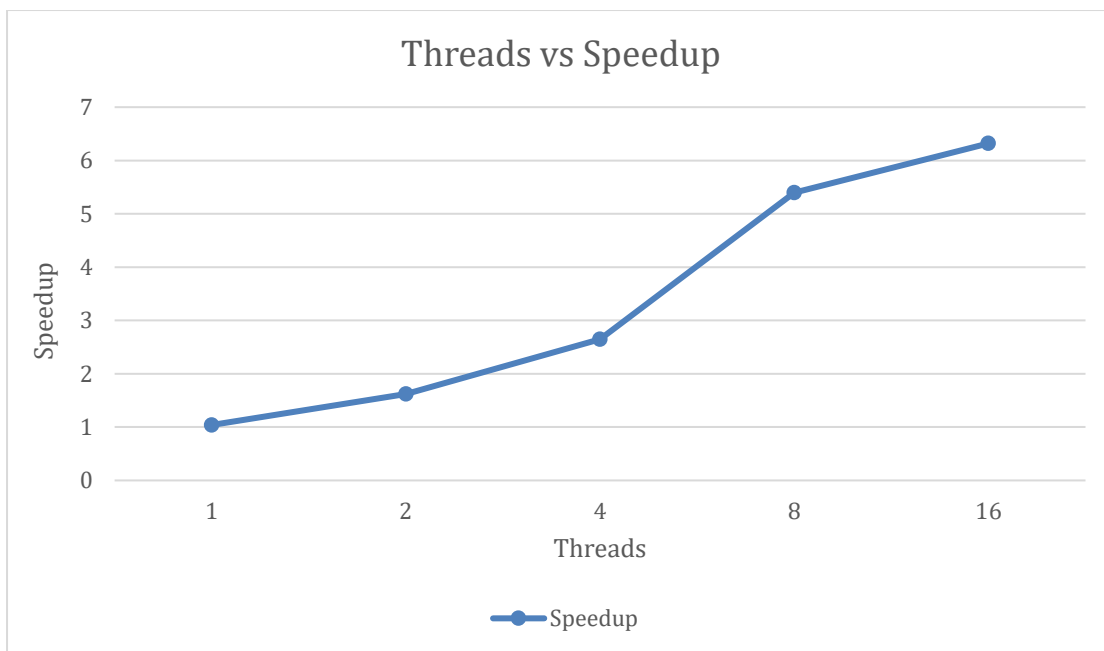
##### I. Number of threads vs Execution time

Number of Threads	Execution Time (s)
1	5.305
2	3.413
4	2.081
8	1.021
16	0.872



## II. Number of threads vs Speedup

Number of Threads	Speedup
1	1.04
2	1.62
4	2.65
8	5.40
16	6.32





## Screenshots

### I. Threads=01

```
ASUS@LAPTOP-500RALBN MINGW64 /c/Users/ASUS/Desktop/PC_Assignment_03
$ export OMP_NUM_THREADS=1
./openmp_code
OpenMP execution:
Threads used: 1
Time: 5.305000 sec
Speedup: 1.04x
```

### II. Threads=02

```
ASUS@LAPTOP-500RALBN MINGW64 /c/Users/ASUS/Desktop/PC_Assignment_03
$ export OMP_NUM_THREADS=2
./openmp_code
OpenMP execution:
Threads used: 2
Time: 3.413000 sec
Speedup: 1.62x
```

### III. Threads=04

```
ASUS@LAPTOP-500RALBN MINGW64 /c/Users/ASUS/Desktop/PC_Assignment_03
$ export OMP_NUM_THREADS=4
./openmp_code
OpenMP execution:
Threads used: 4
Time: 2.081000 sec
Speedup: 2.65x
```

### IV. Threads=08

```
ASUS@LAPTOP-500RALBN MINGW64 /c/Users/ASUS/Desktop/PC_Assignment_03
$ export OMP_NUM_THREADS=8
./openmp_code
OpenMP execution:
Threads used: 8
Time: 1.021000 sec
Speedup: 5.40x
```

### V. Threads=16

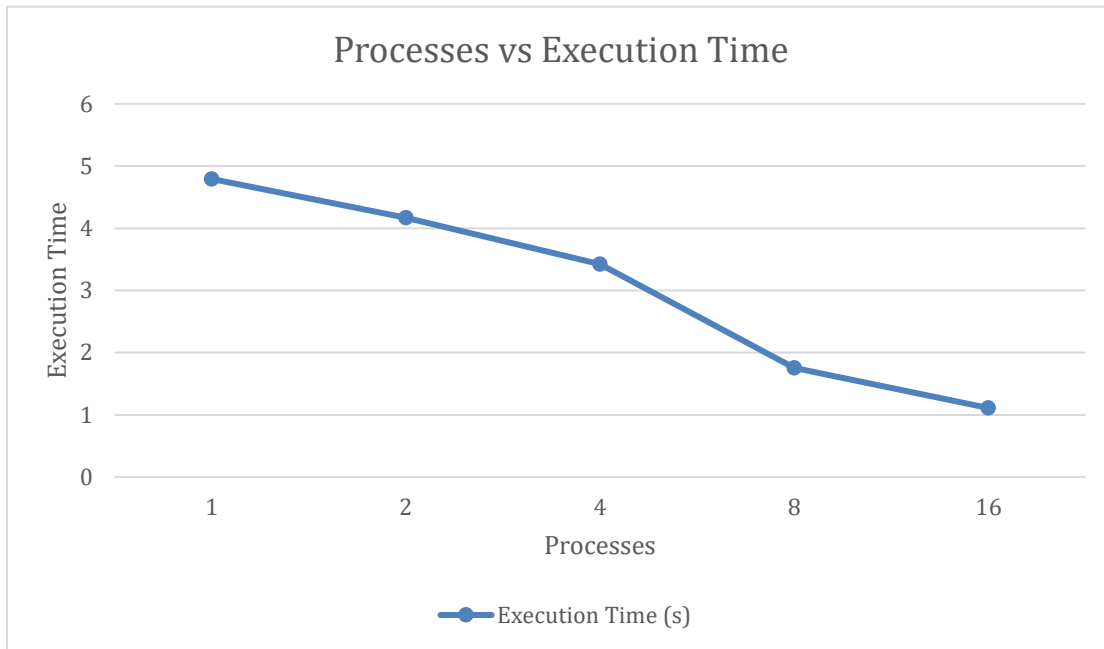
```
ASUS@LAPTOP-500RALBN MINGW64 /c/Users/ASUS/Desktop/PC_Assignment_03
$ export OMP_NUM_THREADS=16
./openmp_code
OpenMP execution:
Threads used: 16
Time: 0.872000 sec
Speedup: 6.32x
```

## B. MPI Evaluation

### A. Graphs

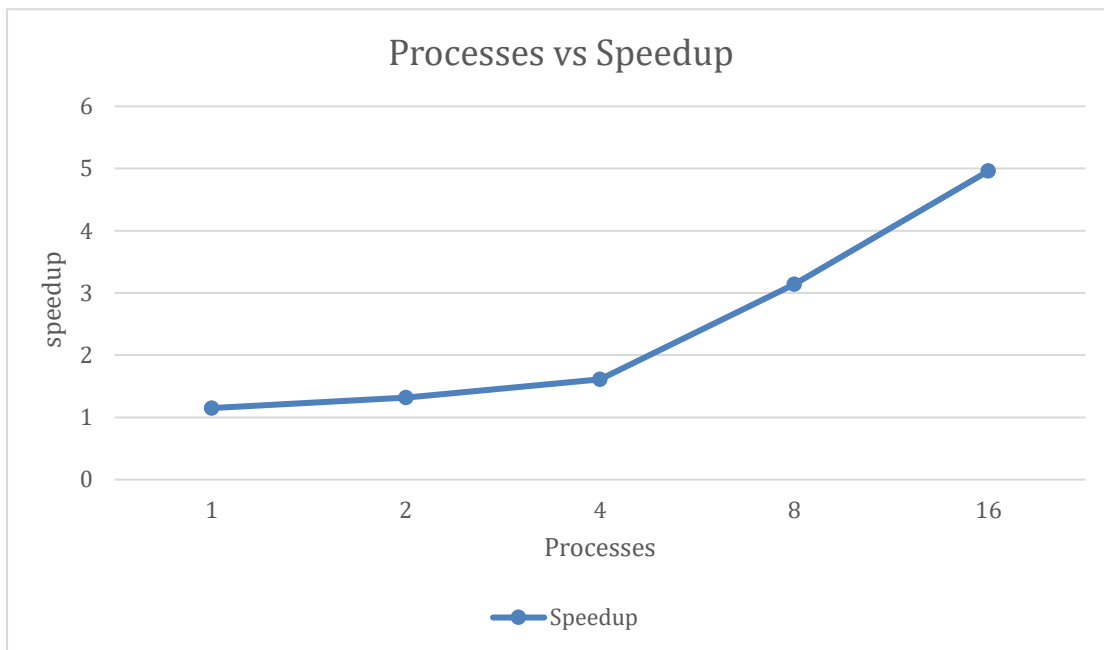
#### I. Number of Processes vs Execution Time

Number of Processes	Execution Time (s)
1	4.792355
2	4.165871
4	3.420793
8	1.755429
16	1.110914



## II. Number of Processes vs Speedup

Number of Processes	Speedup
1	1.15
2	1.32
4	1.61
8	3.14
16	4.96



## B. Screenshots

### I. Processors=1

```
ASUS@LAPTOP-500RALBN MINGW64 /c/Users/ASUS/Desktop/PC_Assignment_03
$ mpicc mpi_code.c -o mpi_code

ASUS@LAPTOP-500RALBN MINGW64 /c/Users/ASUS/Desktop/PC_Assignment_03
$ mpiexec -np 1 ./mpi_code
MPI execution:
Processes used: 1
Time: 4.792355 sec
Speedup: 1.15x
```

### II. Processors=2

```
ASUS@LAPTOP-500RALBN MINGW64 /c/Users/ASUS/Desktop/PC_Assignment_03
$ mpiexec -np 2 ./mpi_code
MPI execution:
Processes used: 2
Time: 4.165871 sec
Speedup: 1.32x
```

### III. Processors=4

```
ASUS@LAPTOP-500RALBN MINGW64 /c/Users/ASUS/Desktop/PC_Assignment_03
$ mpiexec -np 4 ./mpi_code
MPI execution:
Processes used: 4
Time: 3.420793 sec
Speedup: 1.61x
```

### IV. Processors=8

```
ASUS@LAPTOP-500RALBN MINGW64 /c/Users/ASUS/Desktop/PC_Assignment_03
$ mpiexec -np 8 ./mpi_code
MPI execution:
Processes used: 8
Time: 1.755429 sec
Speedup: 3.14x
```

### V. Processors=16

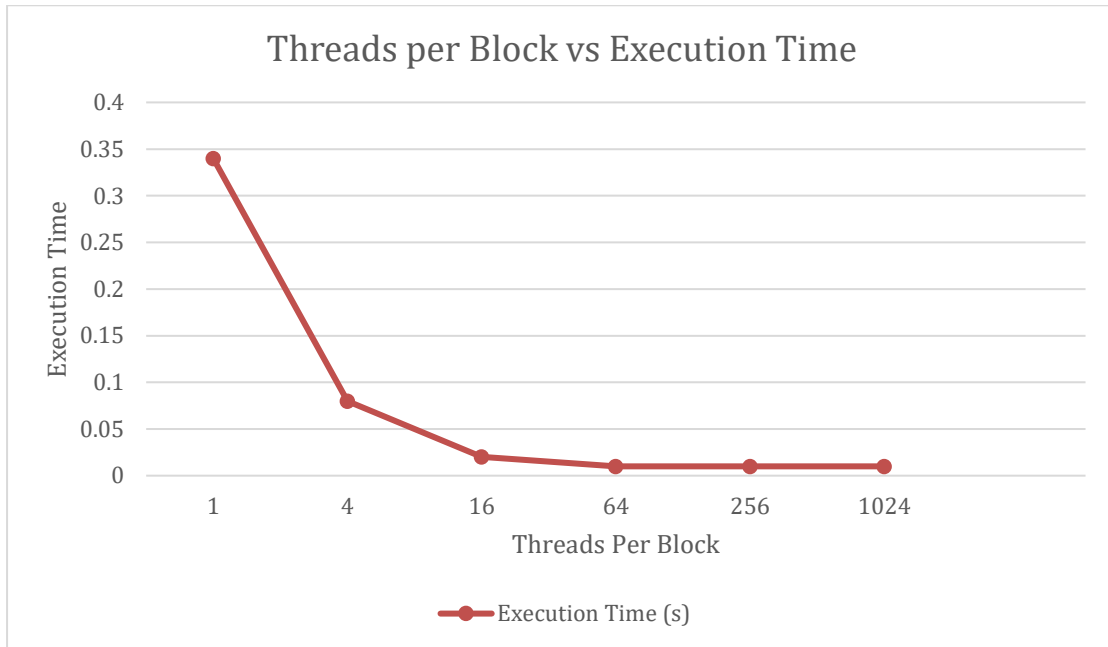
```
ASUS@LAPTOP-500RALBN MINGW64 /c/Users/ASUS/Desktop/PC_Assignment_03
$ mpiexec -np 16 ./mpi_code
MPI execution:
Processes used: 16
Time: 1.110914 sec
Speedup: 4.96x
```

## C. CUDA Evaluation

### A. Graphs

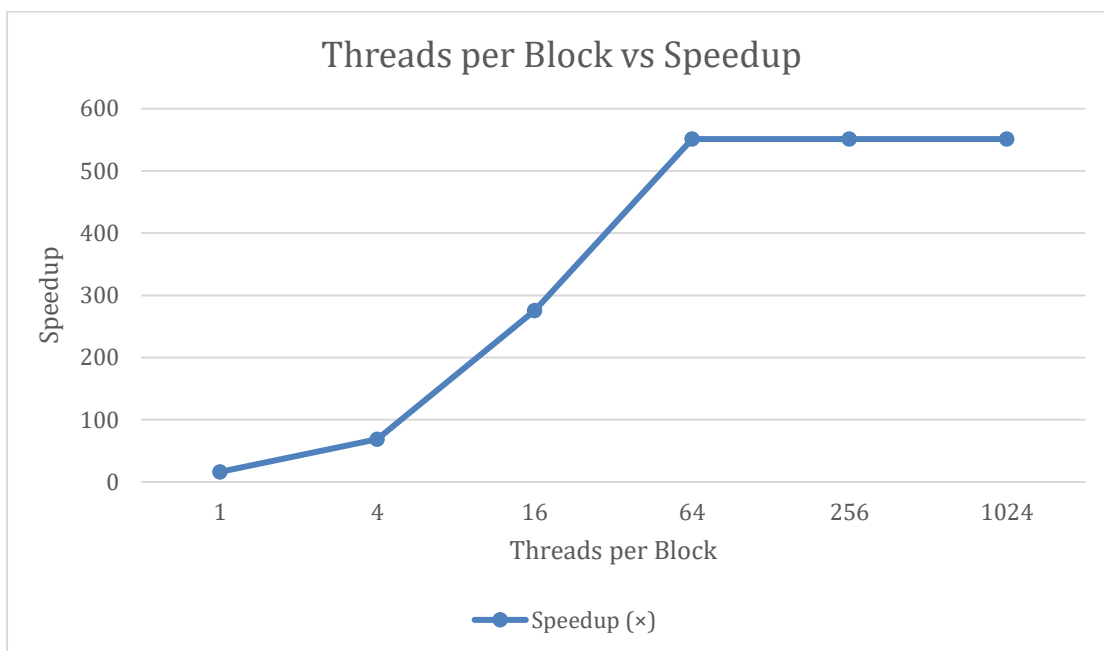
#### I. Configuration parameters vs Execution time

Threads per Block	Blocks (Grid Size)	Execution Time (s)
1	1000 × 1000	0.340
4	500 × 500	0.080
16	250 × 250	0.020
64	125 × 125	0.010
256	63 × 63	0.010
1024	32 × 32	0.010



## II. Configuration parameters vs Speedup

Threads per Block	Blocks (Grid Size)	Speedup (×)
1	1000 × 1000	16.21
4	500 × 500	68.90
16	250 × 250	275.60
64	125 × 125	551.20
256	63 × 63	551.20
1024	32 × 32	551.20



## B. Screenshots

```
C:\Users\ASUS\Desktop\PC_Assignment_03>nvcc cuda_code.cu -o cuda_code.exe
cuda_code.cu
tmpxft_00002590_00000000-10_cuda_code.cudafe1.cpp

C:\Users\ASUS\Desktop\PC_Assignment_03>cuda_code.exe
CUDA execution:
Threads per block: (1*1) = 1
Number of blocks: (1000,1000)
Time: 0.340000 sec, Speedup: 16.21x

CUDA execution:
Threads per block: (2*2) = 4
Number of blocks: (500,500)
Time: 0.080000 sec, Speedup: 68.90x

CUDA execution:
Threads per block: (4*4) = 16
Number of blocks: (250,250)
Time: 0.020000 sec, Speedup: 275.60x

CUDA execution:
Threads per block: (8*8) = 64
Number of blocks: (125,125)
Time: 0.010000 sec, Speedup: 551.20x

CUDA execution:
Threads per block: (16*16) = 256
Number of blocks: (63,63)
Time: 0.010000 sec, Speedup: 551.20x

CUDA execution:
Threads per block: (32*32) = 1024
Number of blocks: (32,32)
Time: 0.010000 sec, Speedup: 551.20x
```

## 4. Comparative Analysis

### A. Comparison on the Same Dataset / Problem Size

- All three implementations were executed on the same matrix multiplication task (N=1000). The performance results show clear differences in scalability and computational efficiency.

#### Execution Time Summary

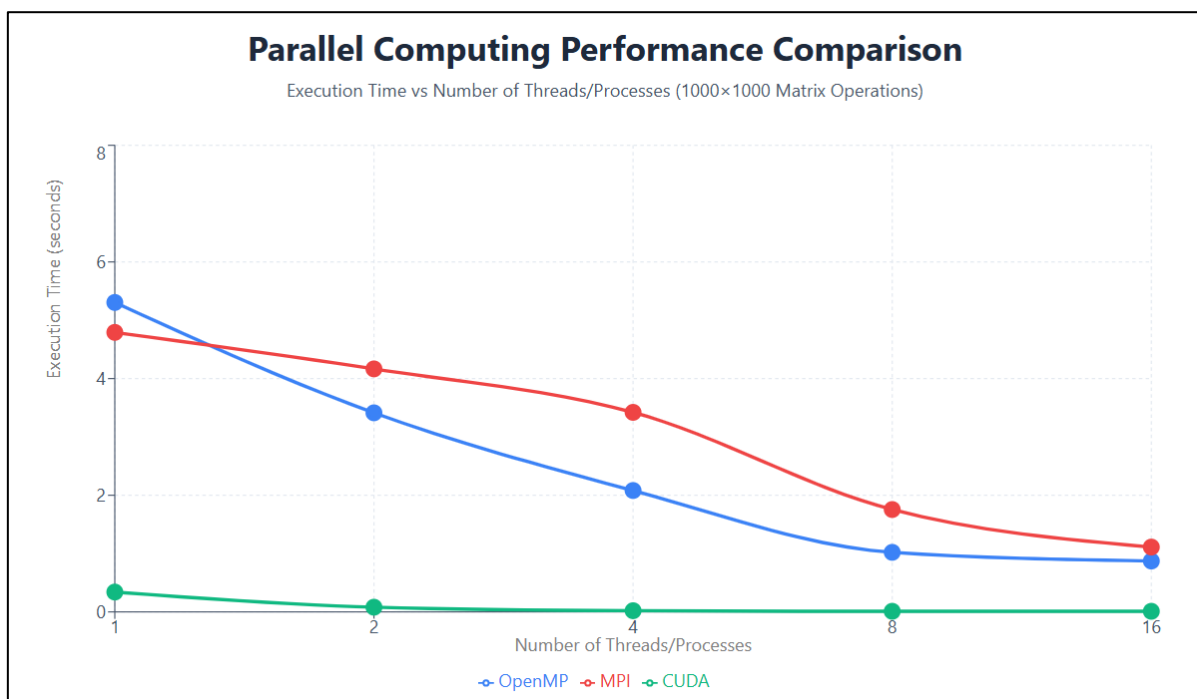
Method	Max Parallel Unit	Best Time (sec)	Worst Time (sec)
OpenMP	16 threads	0.872	5.305
MPI	16 processes	1.110	4.792
CUDA	1024 threads/block	0.010	0.340

#### Speedup Summary

Method	Max Speedup
OpenMP	6.32
MPI	4.96
CUDA	551.20

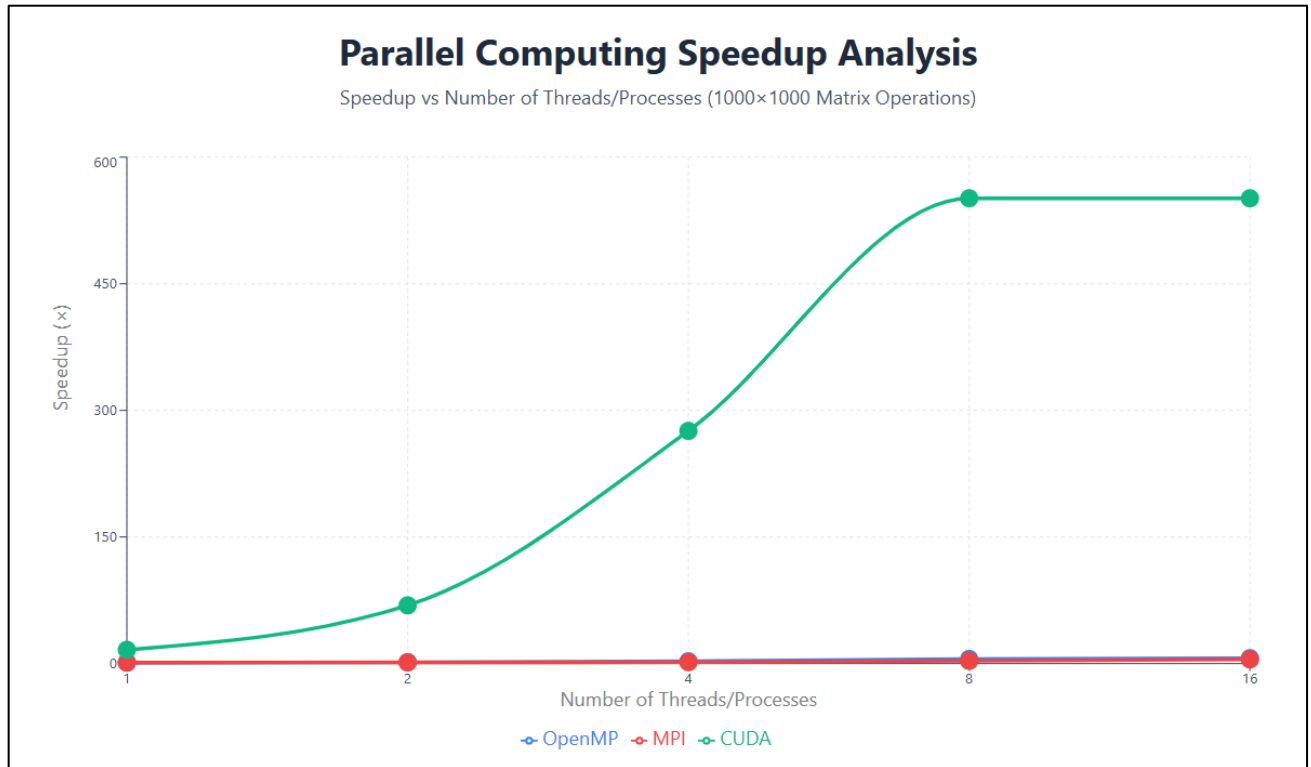
#### Visual Comparison

##### A. Execution time VS Threads/Processes across OpenMP, MPI, CUDA





## B. Execution time vs Threads/Processes across OpenMP, MPI, CUDA



## B. Comparative Discussion

### OpenMP

#### Strengths

- Very easy to integrate into existing C/C++ code using pragmas.
- Great when the workload fits a shared-memory architecture (single machine).
- Good scaling up to a moderate number of threads.

#### Weaknesses

- Limited by shared-memory hardware.
- Thread contention and cache issues become noticeable after 8–16 threads.
- Not suitable for distributed systems.

#### Observation:

OpenMP scales nicely on your CPU, but clearly hits a saturation point around 16 threads.

## MPI

### Strengths

- Designed for distributed-memory systems (clusters).
- Good control over data distribution and communication.
- Scales beyond a single machine if hardware is available.

### Weaknesses

- Communication overhead increases significantly with process count.
- For small-to-medium matrix sizes, overhead outweighs raw compute advantages.
- More complex programming model than OpenMP.

### Observation:

MPI underperforms on a single PC because communication cost dominates. On a proper multi-node cluster, MPI would catch up or surpass OpenMP.

## CUDA

### Strengths

- Massive parallelism—thousands of GPU cores.
- Very high arithmetic throughput.
- Best suited for data-parallel, highly regular operations like matrix multiplication.

### Weaknesses

- Requires GPU hardware and CUDA toolkit.
- Extra complexity in kernel design, memory transfers, and block/thread configuration.
- Performance depends heavily on correct block size tuning.

### Observation:

- CUDA absolutely crushes CPU-based methods:
- **551× speedup** is undeniable proof that GPUs dominate this workload type.

### C. Which Implementation Is Most Appropriate?

- If you have access to a capable NVIDIA GPU, CUDA is unquestionably the best for matrix multiplication.
- The speedups (up to 551×) demonstrate that GPUs are built precisely for this kind of dense numerical computation.
- If GPU resources were *not* available:
  - OpenMP is the best on a single machine.
  - MPI becomes the best only when multiple physical nodes are available.

## 5. Critical Reflection

### A. Challenges Faced

#### CUDA

- Large matrices (N=1000) caused GPU memory issues.
- Dynamic memory allocation and block/thread configuration were complex.
- Modifying configurations required significant code changes.

#### OpenMP

- Static arrays failed; dynamic allocation with malloc/free was needed.
- Hardcoded serial time required for accurate speedup calculation.

#### MPI

- Large matrices caused stack overflow; resolved with dynamic allocation.
- Complex logic required for distributing matrices across processes.

### B. Limitations

- **Memory:** GPU and stack limits restricted large matrices; dynamic allocation added minor overhead.
- **Implementation Complexity:** CUDA required major restructuring; all implementations relied on a fixed serial baseline.
- **Debugging:** Memory-related crashes were challenging to diagnose.

### C. Solutions Implemented

- **Dynamic Allocation:** Used malloc/free and cudaMalloc/cudaFree.
- **Consistent Timing:** SERIAL\_TIME used for speedup; proper timing around computation sections.
- **Modular Code:** Separated allocation, computation, cleanup; added error checks; adjustable thread/block configurations.

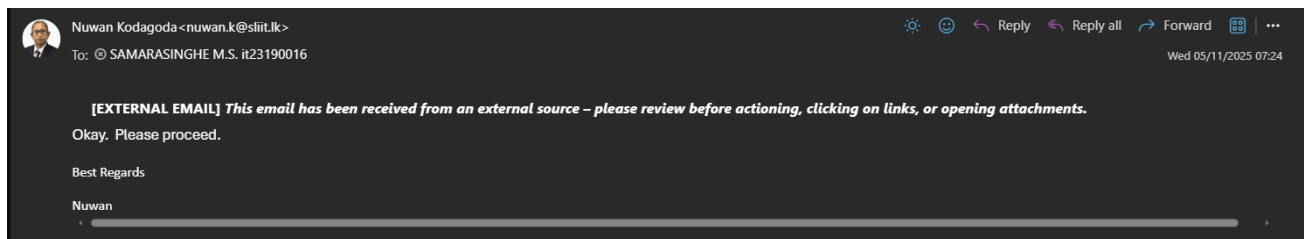
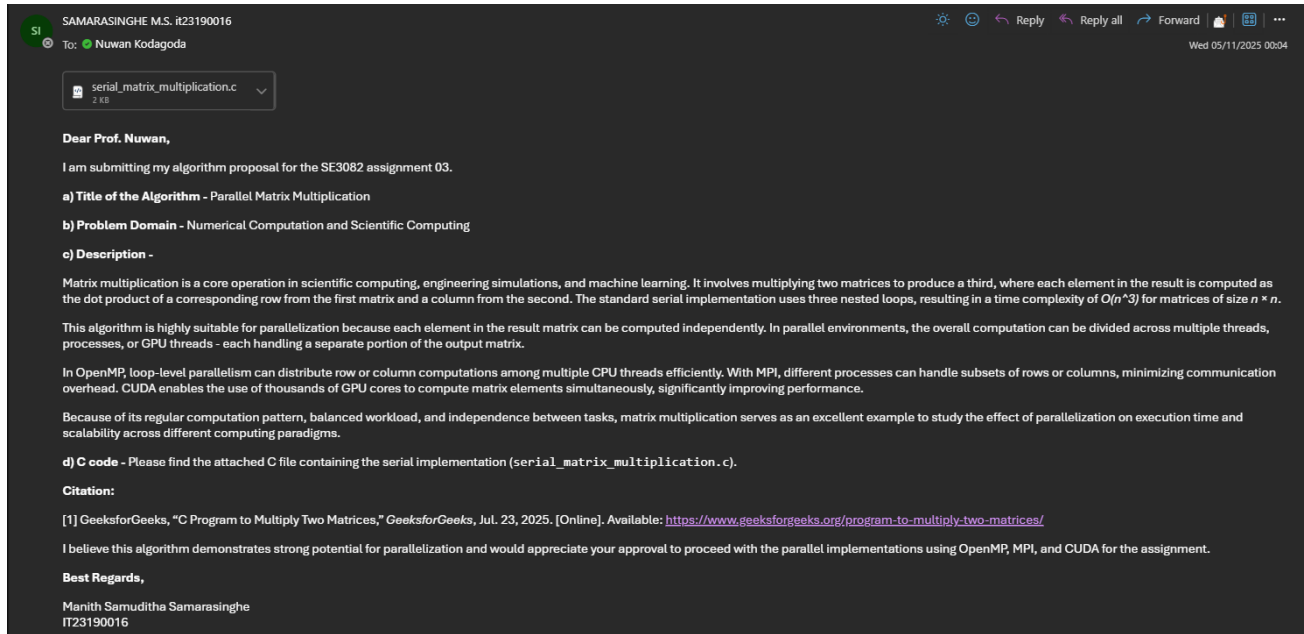
### D. Lessons Learned

- Dynamic memory is essential for large-scale parallel computing.
- Baseline timing consistency is crucial for performance comparison.
- CUDA has the highest complexity; OpenMP is simplest; MPI adds communication challenges.
- Configuration parameters strongly affect performance.

### E. Future Improvements

- Memory pooling and GPU unified memory for efficiency.
- Configurable thread/block selection and automatic memory sizing.
- Adaptive configuration and hybrid parallel approaches.
- Integrated profiling and error-handling mechanisms.

## Proposal Email and Approval



## Code Execution Demo

[Assignment03 Demo.mp4](#)