#### **Assignment 1 – Report**

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#### Section1: MPI programming

- Code 1: ring.c
  - Code and Readme file: on github https://github.com/Fatma-Moustafa/2021Assignment01
  - o Set of variables used:
    - $rank \rightarrow rank of processor$
    - size → number of processors
    - left → rank of the left node
    - right → rank of the right node
    - $msgleft \rightarrow message to be sent to the left$
    - msgright → message to be sent to the right
    - $tag \rightarrow tag$  of the processor
    - lastLeftTag → tag of the left processor that sent the last message
    - lastRightTag → tag of the right processor that sent the last message
  - o Formulas used:
    - Speedup is the ratio between time to execute the serial workload and time to execute the workload parallelly with P processors

$$S(P) = \frac{T(1)}{T(P)}$$

 Efficiency is the average speedup by one processor in a parallel program

$$E(P) = \frac{S(P)}{P}$$

- Test cases and iterations:
  - 100 iterations were performed for 2, 4, 8, 16, 32 processors
- o Results and conclusion:
  - As the number of processors increase, the time of computation increase. This shows that the ring code has a weak scalability, since the efficiency decrease with increasing the number of processors.
- Table1: shows the computational time, speedup and efficiency as the number of processors increase

Processor	Time	S(P) = T(1) / T(P)	E(P) = S(P) / P
1	0.00128008	1	1
2	0.00210541	0.6079955923	0.3039977962
4	0.00235606	0.5433138375	0.1358284594
8	0.00272901	0.469063873	0.05863298412
16	0.00396601	0.3227626758	0.02017266724
32	0.00554364	0.230909655	0.00721592672

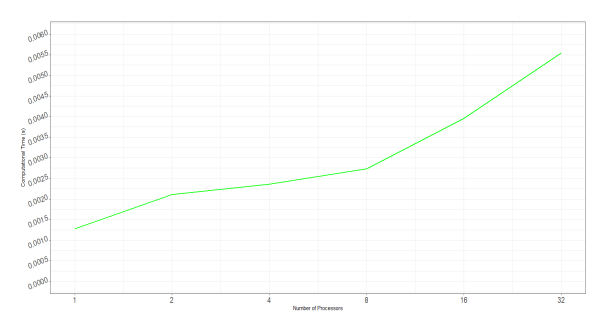


Figure 1: Computational time increases almost quadratically when number of processors doubles  $\,$ 

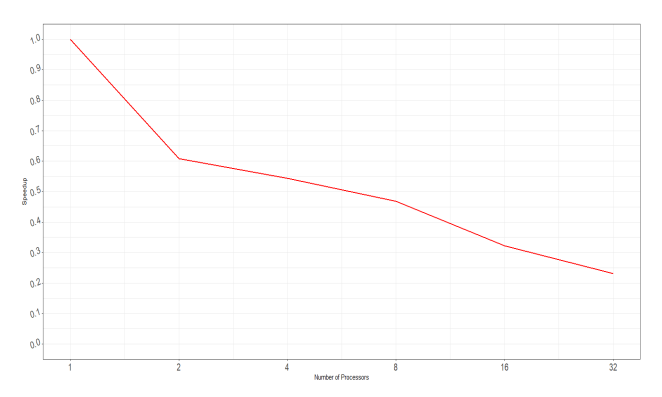


Figure 2: Speedup decrease as the number of processors increase

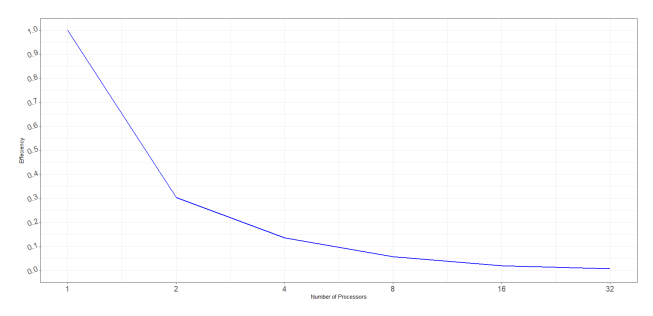


Figure 3: Efficiency decrease as the number of processors increase

- Code 2: sum3Dmatrix.cpp
  - Code and Readme file: on github https://github.com/Fatma-Moustafa/2021Assignment01
  - Set of variables used:
    - rank → rank of processor
    - $size \rightarrow number of processors$
    - $x \rightarrow dimension x-axis$
    - $y \rightarrow dimension y-axis$
    - $z \rightarrow dimension z$ -axis
    - $x \text{ submatrix} \rightarrow \text{dimension submatrix on } x$ -axis
    - y\_ submatrix → dimension submatrix on y-axis
    - z submatrix  $\rightarrow$  dimension submatrix on z-axis
    - matrix\_1, matrix\_2, matrix\_3, matrix\_1\_ submatrix, matrix\_2\_ submatrix, matrix\_3\_ submatrix allocated dynamically with corresponding dimension
    - elements\_ submatrix → Multiplication of the submatrices with smaller dimensions. This value is needed in the split of the job in multiprocessors.
  - Test cases: Matrices dimensions with number of processors kept constant at 24
    - 2400 x 100 x 100
    - 1200 x 200 x 100

- 800 x 300 x 100
- o Results and conclusion:
  - Maximum, minimum and average time for different submatrix division for the following matrix size: 2400 x 100 x 100

	SUBMATRIX	MAX	MIN TIME	AVG TIME
	DIVISION	TIME		
1D	100 * 100 * 100	0.111555	0.110219	0.111346
2D	200 * 50 * 100	0.129135	0.129024	0.129085
	400 * 25 * 100	0.106145	0.106029	0.106082
3D	400 * 50 * 50	0.107988	0.106702	0.107354
	800 * 25 * 50	0.1086	0.108506	0.108555

 Maximum, minimum and average time for different submatrix division for the following matrix size: 1200 x 200 x 100

	SUBMATRIX	MAX	MIN TIME	AVG TIME
	DIVISION	TIME		
1D	50 * 100 * 100	0.070834	0.0707541	0.0707909
2D	100 * 100 * 100	0.108833	0.107592	0.107559
	200 * 50 * 100	0.109028	0.107788	0.108683
	200 * 200 * 25	0.108971	0.108835	0.108912
	100 * 200 * 50	0.109147	0.107856	0.108938
	400 * 25 * 100	0.109756	0.10964	0.109706
3D	200 * 100 * 50	0.108592	0.107287	0.108236
	400 * 50 * 50	0.120001	0.118698	0.119639

 Maximum, minimum and average time for different submatrix division for the following matrix size: 800 x 300 x 100

	SUBMATRIX	MAX	MIN	AVG TIME
	DIVISION	TIME	TIME	
2D	100 * 100 * 100	0.108054	0.107968	0.108007
	200 * 50 * 100	0.108232	0.106991	0.10789
	400 * 25 * 100	0.105332	0.105194	0.105265
	800 * 25 * 50	0.108422	0.108318	0.108368
	800 * 50 * 25	0.108903	0.108833	0.108864
3D	200 * 100 * 50	0.106568	0.106472	0.106518

400 * 100 * 25	0.108621	0.108503	0.108557
400 * 50 * 50	0.108691	0.10857	0.108633

### Section2: Measure MPI point to point performance

- Set of variables used:
  - $N \rightarrow map by node$
  - $S \rightarrow map by socket$
  - $C \rightarrow map by core$
  - $RB \rightarrow report bindings$
- o Running Netwrok parameters:
  - MCA: modular component architecture
  - OB1: multidevice multi rail engine
  - PML: MPI point-to-point communication
  - BTL: byte transport layer
  - BML: BTL multiplexing layer
  - TCP: transmission control protocol
  - SELF: process-loopback communications
  - VADER: shared memory
  - UCX: unified communications X

```
MCA pml: v (MCA v2.1.0, API v2.0.0, Component v4.0.3)
MCA pml: cm (MCA v2.1.0, API v2.0.0, Component v4.0.3)
MCA pml: monitoring (MCA v2.1.0, API v2.0.0, Component v4.0.
MCA pml: ob1 (MCA v2.1.0, API v2.0.0, Component v4.0.3)
MCA pml: ucx (MCA v2.1.0, API v2.0.0, Component v4.0.3)
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MCA btl: self (MCA v2.1.0, API v3.1.0, Component v4.0.3)
MCA btl: openib (MCA v2.1.0, API v3.1.0, Component v4.0.3)
MCA btl: tcp (MCA v2.1.0, API v3.1.0, Component v4.0.3)
MCA btl: uct (MCA v2.1.0, API v3.1.0, Component v4.0.3)
MCA btl: vader (MCA v2.1.0, API v3.1.0, Component v4.0.3)
MCA fbtl: posix (MCA v2.1.0, API v2.0.0, Component v4.0.3)
```

- Test cases and iterations:
  - CSV files with all the runs using GCC and INTEL modules: on GitHub
  - MPI Benchmark comparison between GCC and INTEL

	Latency	Bandwidth	
GCC - No mapping	0.2	13431	

GCC - No mapping with ucx		0.21	13674
INTEL		0.42	6445
Run on 2 contigu	uou	s processors	
INTEL PROCESSORLIST 0,1		0.42	6377
Run on the s	sam	ie socket	
INTEL PROCESSORLIST 0,2		<mark>0.23</mark>	10926

 Different mappings for MPI Benchmark comparison between GCC and INTEL

	SOCKET		NODE		CORE	
	Latency	Bandwid	Latency	Bandwid	Laten	Bandwid
		th		th	cy	th
basic	0.39	14125	0.21	13270	0.2	13776
pml = ob1	0.53	10284	0.26	9685	0.25	9724
pml = ob1, btl = self	<mark>7.97</mark>	3477	<mark>5⋅39</mark>	7037	<mark>5⋅43</mark>	7000
pml = ucx, btl = self	0.41	14058	0.2	13656	0.2	13771
pml = ucx, btl =	0.4	14277	0.2	13646	0.2	13083
vader						
INTEL	0.43	6341	0.42	6439	0.43	6336

#### o Results and conclusion:

#### GENERAL OVERVIEW:

- For the OpenMPI the latency and bandwidth are 0.2 microsecond and 13431 respectively. Using a UCX protocol does not have an effect on these values.
- It is noticed that Intel has almost double the latency and half the bandwidth, even we run it on 2 contiguous processors.
- However, when run on the same socket the latency and bandwidth has values almost similar to that with the OpenMPI.
- UCX for infiniband, OB1 PML + BTL for all others

#### • OPENMPI:

- Mapping by core or by node has similar results in terms of latency and bandwidth. When mapping by socket we get almost similar bandwidth but almost double the latency.
- Using the PML as UCX maintains the values of latency and bandwidth, regardless of the BTL being self or vader.

- With the PML being OB1 the latency increases by around 25% and the bandwidth decreases by almost 25%.
- In the case of having PML as OB1 and BTL as self, the latency is the highest and the bandwidth is the lowest among all the other cases.
  - Latency shows an increase of 20 times in the map by socket case and even more when mapping is done by node or core.
  - The bandwidth decreases to quarter its value when mapping by socket and to half when mapping by core or node.
  - TCP protocol has high latency because it has high travelling distance.

#### ■ INTEL:

- Whatever the type of mapping that is being done, the values of latency and bandwidth remains almost the same for all these mappings.
- Also, they are similar to the values obtained when we run on 2 contiguous processors.
- Latency does not increase. We have 2 processes one on core o and the other process is on the other core with same socket and same node.

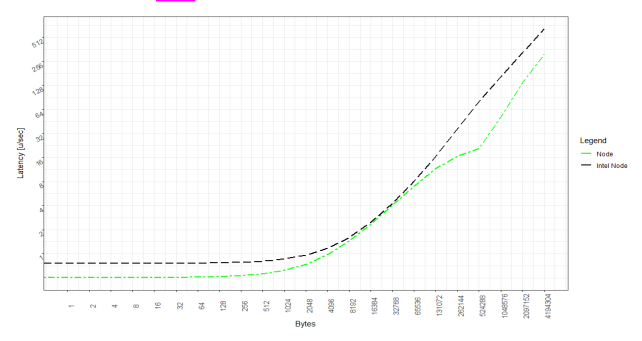


Figure 4: fitted model of latency computed for mapping by node comparing GCC and INTEL modules

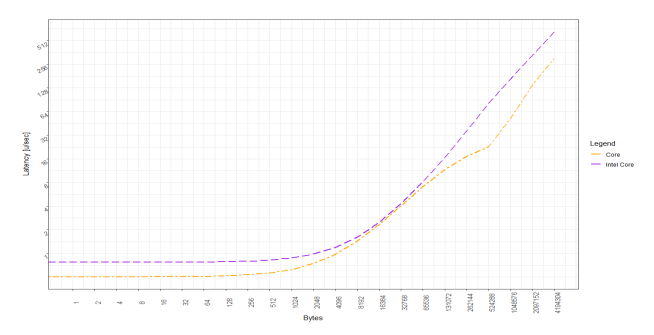


Figure 5: fitted model of latency computed for mapping by core comparing GCC and INTEL modules

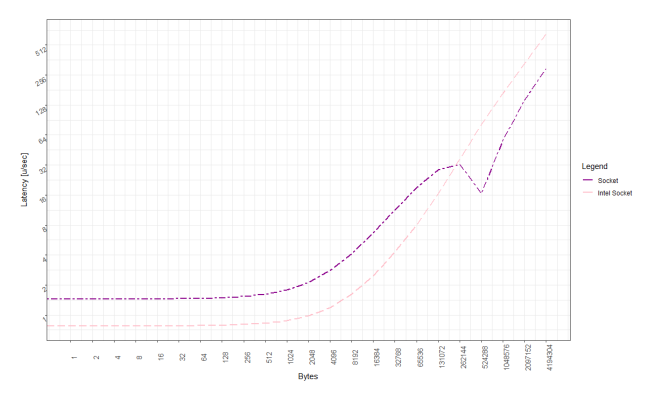


Figure 6: fitted model of latency computed for mapping by socket comparing GCC and INTEL modules  $\frac{1}{2}$ 

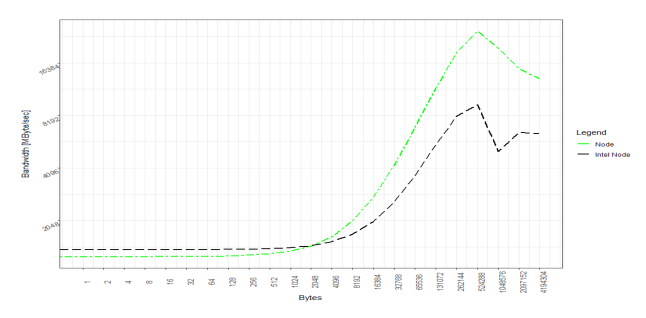


Figure 7: fitted model of Bandwidth computed for mapping by node comparing GCC and INTEL modules

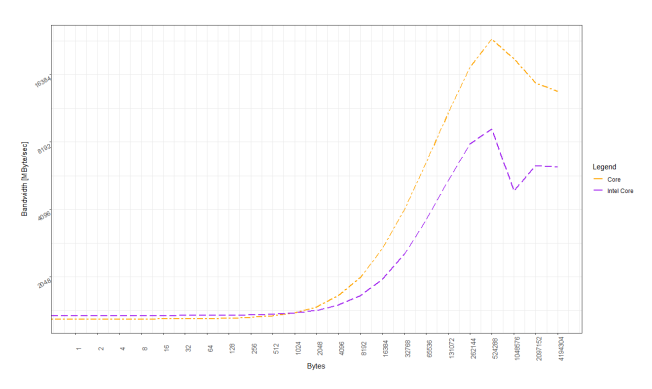


Figure 8: Fitted model of Bandwidth computed for mapping by core comparing GCC and INTEL modules  ${\sf GCC}$ 

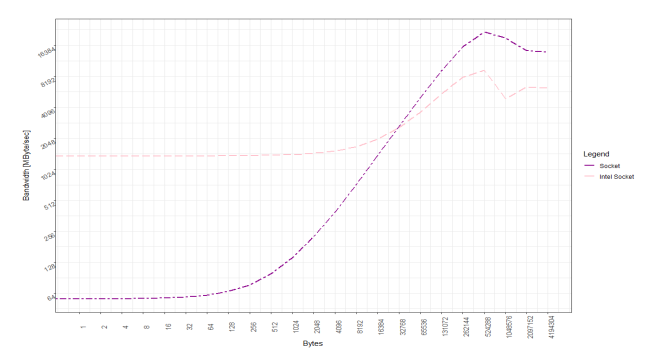


Figure 9: fitted model of Bandwidth computed for mapping by socket comparing GCC and INTEL modules

# Section3: Compare performance observed against performance model for Jacobi solver

- Set of variables used:
  - $N \rightarrow$  number of processes
  - $N_x \rightarrow$  number of processors in along x
  - $N_y \rightarrow$  number of processors in along y
  - $N_z \rightarrow$  number of processors in along z
  - $k \rightarrow$  largest number of coordinate directions in which number of processess is greater than one
  - $c(L,N) \rightarrow maximum$  bidirectional data volume transferred over a node's network link
  - $B \rightarrow \text{full-duplex bandwidth}$
  - $T_c(L,N)$  → communication time
  - $T_s(L,N)$  → raw compute time for all cell updates in a Jacobi sweep
  - $P_1(L)$  → single-processor performance for a domain size  $L^3$
  - P(L,N) → performance on N= N<sub>x</sub> N<sub>y</sub> N<sub>z</sub> processors for a particular overal problem size of L<sup>3</sup>N grid points (using cubic subdomain of size L<sup>3</sup>)
  - $L \rightarrow$  spatial dimension
  - $T_1 \rightarrow latency of network$
- Formulas used:

$$c(L, \vec{N}) = L^2 \cdot k \cdot 2 \cdot 8$$

$$T_c(L, \vec{N}) = \frac{c(L, \vec{N})}{B} + kT_b$$

$$T_c(L, \vec{N}) = \frac{c(L, \vec{N})}{B} + kT_l$$

$$P(L, \vec{N}) = \frac{L^3 N}{T_c(L, \vec{N}) + T_s(L)}$$

- Running parameters:
  - Spat\_dim: problem size = L, in this case it was 1200
  - proc\_dim: possible presets for number of processes, in this case it
  - pbc\_check = periodicity, in this case is t
- Test cases and iterations:
  - Runs on a thin node for 4,8,12 processors
  - Runs on 2 thin nodes for 12, 24, 48 processors
  - Runs on GPU node
- Results and conclusion:

In the following tables, some of the output of the runs performed are shown.

- Runs on a thin node for 4,8,12 processors
  - Map by node

N	Nx	Ny	Nz	k	c(L,N)	Tc(L,N)	P(L,N)	P(1)*N/P(L,N)
1	1	1	1	0	0	0	114.10122	1
4	2	2	1	4	92160	7.661737771	303.075668	1.505910665
8	2	2	2	6	138240	11.49260666	518.976297	1.758865998
12	3	2	2	6	138240	11.49260666	778.464446	1.758865998

#### Map by core

N	Nx	Ny	Nz	k	c(L,N)	Tc(L,N)	P(L,N)	P(1)*N/P(L,N)
1	1	1	1	0	0	0	114.10122	1
4	2	2	1	4	92160	7.48989547	305.376651	1.494563781
8	2	2	2	6	138240	11.23484321	524.047439	1.741845672
12	3	2	2	6	138240	11.23484321	786.071158	1.741845672

#### Map by socket

N	Nx	Ny	Nz	k	c(L,N)	Tc(L,N)	P(L,N)	P(1)*N/P(L,N)
1	1	1	1	0	0	0	114.10122	1
4	2	2	1	4	92160	8.08460177	297.558448	1.533832713
8	2	2	2	6	138240	12.12690265	506.905586	1.80074907
12	3	2	2	6	138240	12.12690265	760.35838	1.80074907

- Runs on 2 thin nodes for 12, 24, 48 processors
  - Map by node

N	Nx	Ny	Nz	k	c(L,N)	Tc(L,N)	P(L,N)	P(1)*N/P(L,N)
1	1	1	1	0	0	0	114.10122	1
12	3	2	2	6	138240	11.49260666	778.464446	1.758865998
24	4	3	2	6	138240	11.49260666	1556.92889	1.758865998
48	4	4	3	6	138240	11.49260666	3113.85778	1.758865998

## • Map by core

N	Nx	Ny	Nz	k	c(L,N)	Tc(L,N)	P(L,N)	P(1)*N/P(L,N)
1	1	1	1	0	0	0	114.10122	1
12	3	2	2	6	138240	11.23484321	786.071158	1.741845672
24	4	3	2	6	138240	11.23484321	1572.14232	1.741845672
48	4	4	3	6	138240	11.23484321	3144.28463	1.741845672

# • Map by socket

N	Nx	Ny	Nz	k	c(L,N)	Tc(L,N)	P(L,N)	P(1)*N/P(L,N)
1	1	1	1	0	0	0	114.10122	1
12	3	2	2	6	138240	12.12690265	760.35838	1.80074907
24	4	3	2	6	138240	12.12690265	1520.71676	1.80074907
48	4	4	3	6	138240	12.12690265	3041.43352	1.80074907

### Runs on GPU node

# • Map by node

N	Nx	Ny	Nz	k	c(L,N)	Tc(L,N)	P(L,N)	P(1)*N/P(L,N)
1	1	1	1	0	0	0	114.10122	1
4	2	2	1	4	92160	7.661737771	303.075668	1.505910665
8	2	2	2	6	138240	11.49260666	518.976297	1.758865998
12	3	2	2	6	138240	11.49260666	778.464446	1.758865998
24	4	3	2	6	138240	11.49260666	1556.92889	1.758865998
48	4	4	3	6	138240	11.49260666	3113.85778	1.758865998

# • Map by core

N	Nx	Ny	Nz	k	c(L,N)	Tc(L,N)	P(L,N)	P(1)*N/P(L,N)
1	1	1	1	0	0	0	114.10122	1
4	2	2	1	4	92160	7.48989547	305.376651	1.494563781
8	2	2	2	6	138240	11.23484321	524.047439	1.741845672
12	3	2	2	6	138240	11.23484321	786.071158	1.741845672
24	4	3	2	6	138240	11.23484321	1572.14232	1.741845672
48	4	4	3	6	138240	11.23484321	3144.28463	1.741845672

# Map by socket

N	Nx	Ny	Nz	k	c(L,N)	Tc(L,N)	P(L,N)	P(1)*N/P(L,N)
1	1	1	1	0	0	0	114.10122	1
4	2	2	1	4	92160	8.08460177	297.558448	1.533832713
8	2	2	2	6	138240	12.12690265	506.905586	1.80074907
12	3	2	2	6	138240	12.12690265	760.35838	1.80074907
24	4	3	2	6	138240	12.12690265	1520.71676	1.80074907
48	4	4	3	6	138240	12.12690265	3041.43352	1.80074907

In the following tables, the slowdown factor compared to perfect scaling for the different number of processors and different mappings are shown.

## Runs on a thin node for 4,8,12 processors

by node				by core	by socket		
N		P	N	P	N	P	
	1	1	1	1	1	1	
	4	1.505910665	4	1.494563781	4	1.533832713	
	8	1.758865998	8	1.741845672	8	1.80074907	
1	12	1.758865998	12	1.741845672	12	1.80074907	

### Runs on 2 thin nodes for 12, 24, 48 processors

by node				by core	by socket		
N		P	N	P	N	P	
	1	1	1	1	1	1	
	12	1.758865998	12	1.741845672	12	1.80074907	
	24	1.758865998	24	1.741845672	24	1.80074907	
	48	1.758865998	48	1.741845672	48	1.80074907	

#### Runs on GPU node

	by node		by core	by socket		
N	P	N	P	N	P	
1	1	1	1	1	1	
4	1.505910665	4	1.494563781	4	1.533832713	
8	1.758865998	8	1.741845672	8	1.80074907	
12	1.758865998	12	1.741845672	12	1.80074907	
24	1.758865998	24	1.741845672	24	1.80074907	
48	1.758865998	48	1.741845672	48	1.80074907	