

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>
 - Set of variables used:
 - rank → rank of processor
 - size → number of processors
 - left → rank of the left node
 - right → rank of the right node
 - msgleft → message to be sent to the left
 - msgright → message to be sent to the right
 - tag → 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
 - 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
 - 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

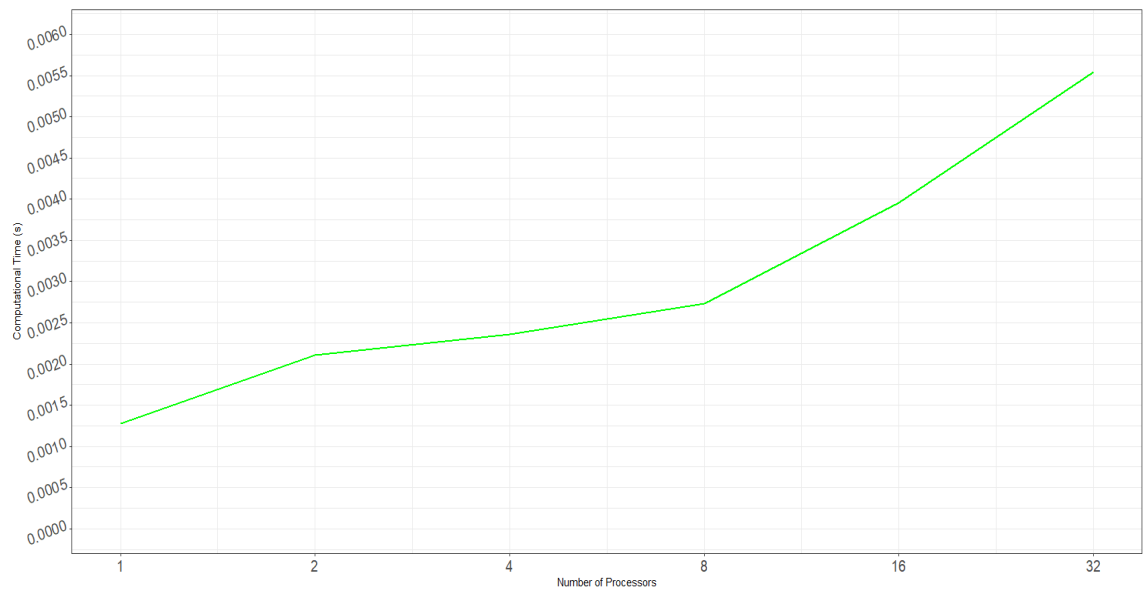


Figure 1: Computational time increases almost quadratically when number of processors doubles

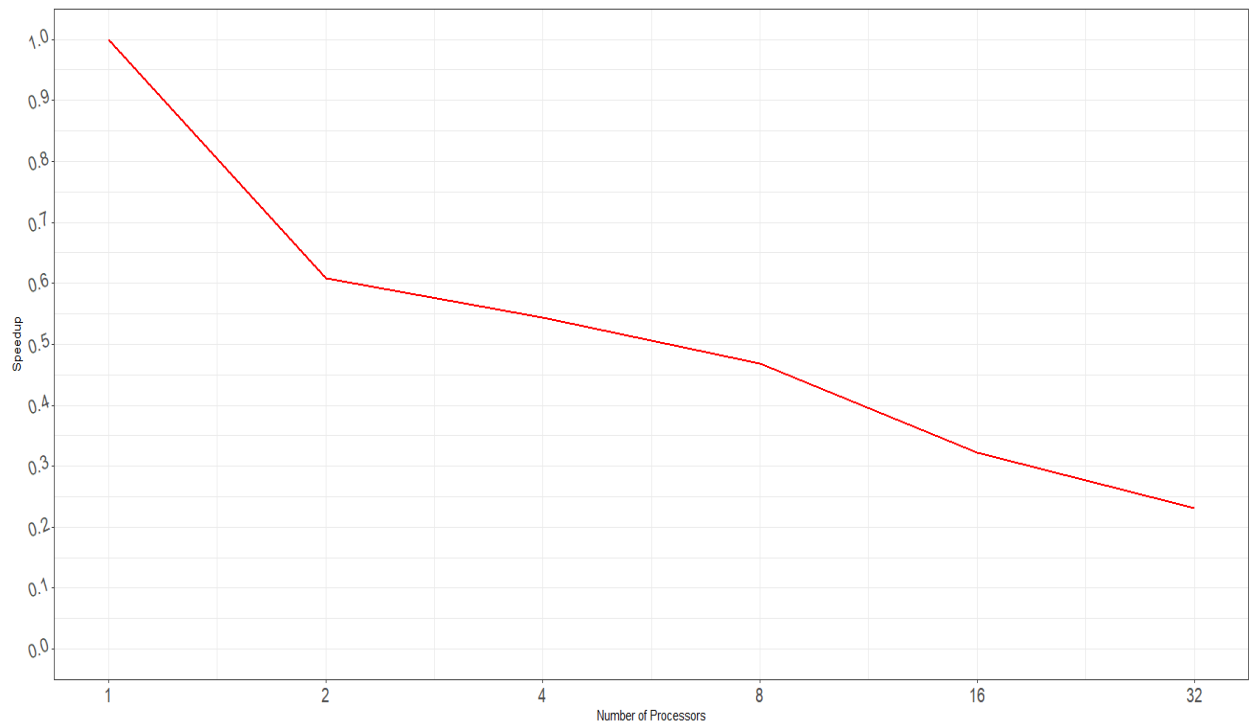


Figure 2: Speedup decrease as the number of processors increase

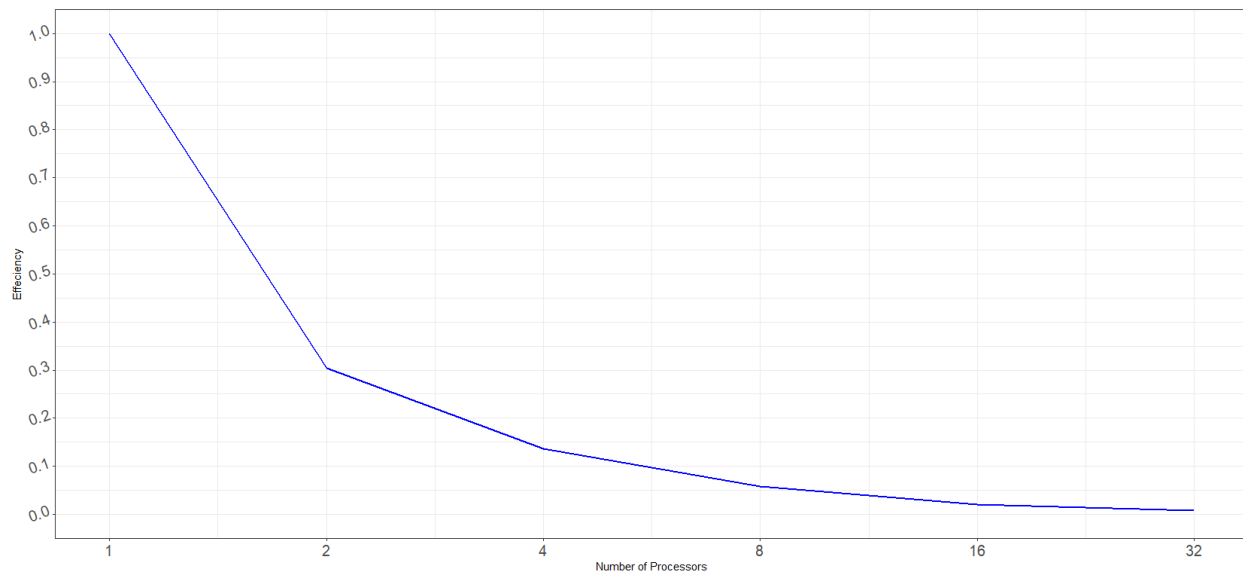


Figure 3: Efficiency decrease as the number of processors increase

- 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 → number of processors
 - x → dimension x-axis
 - y → dimension y-axis
 - z → dimension z-axis
 - x_submatrix → dimension submatrix on x-axis
 - y_submatrix → dimension submatrix on y-axis
 - z_submatrix → 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
- Results and conclusion:
 - Maximum, minimum and average time for different submatrix division for the following matrix size: 2400 x 100 x 100

	SUBMATRIX DIVISION	MAX TIME	MIN TIME	AVG 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 DIVISION	MAX TIME	MIN TIME	AVG 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 DIVISION	MAX TIME	MIN TIME	AVG 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 → map by node
 - S → map by socket
 - C → map by core
 - RB → report bindings
- 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

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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 v4.0.3)
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 contiguous processors				
INTEL PROCESSORLIST 0,1		0.42	6377	
Run on the same socket				
INTEL PROCESSORLIST 0,2		0.23	10926	

- Different mappings for MPI Benchmark comparison between GCC and INTEL

	SOCKET		NODE		CORE	
	Latency	Bandwidth	Latency	Bandwidth	Latency	Bandwidth
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	7.97	3477	5.39	7037	5.43	7000
pml = ucx, btl = self	0.41	14058	0.2	13656	0.2	13771
pml = ucx, btl = vader	0.4	14277	0.2	13646	0.2	13083
INTEL	0.43	6341	0.42	6439	0.43	6336

- 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 0 and the other process is on the other core with same socket and same node.

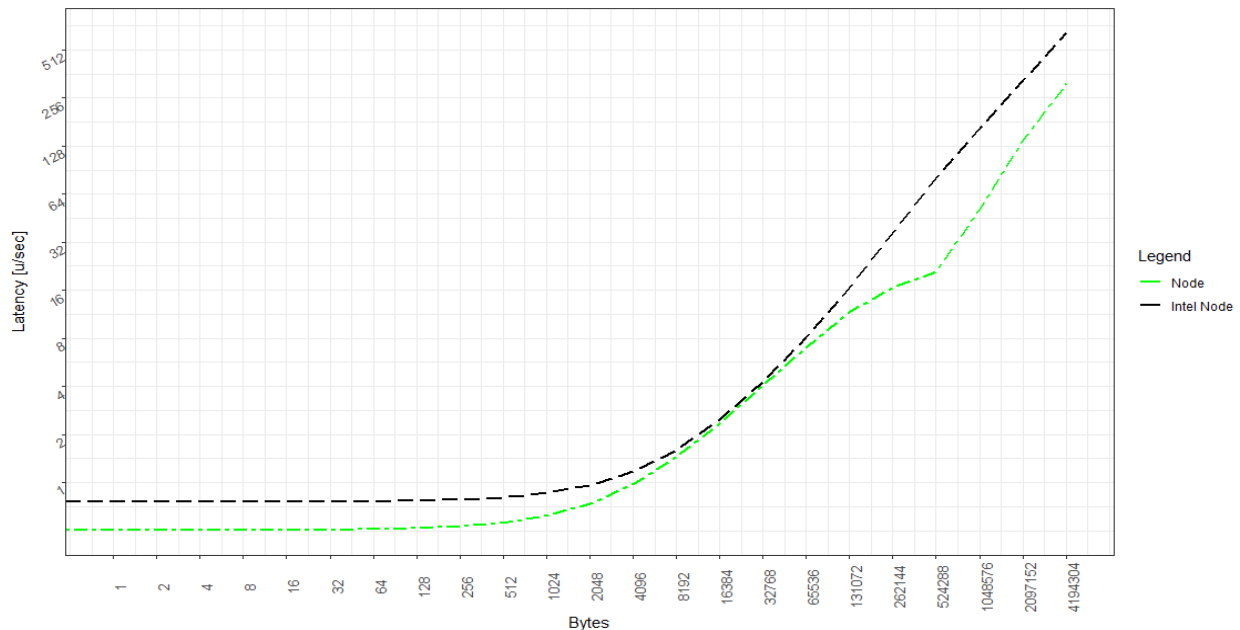


Figure 4: fitted model of latency computed for mapping by node comparing GCC and INTEL modules

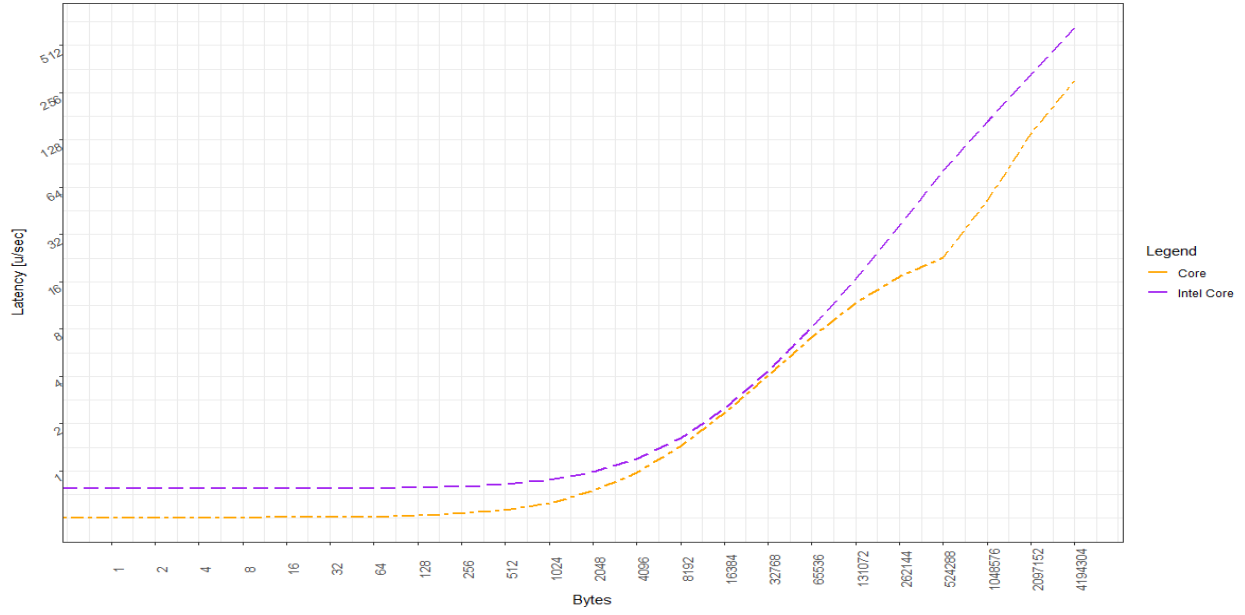


Figure 5: fitted model of latency computed for mapping by core comparing GCC and INTEL modules

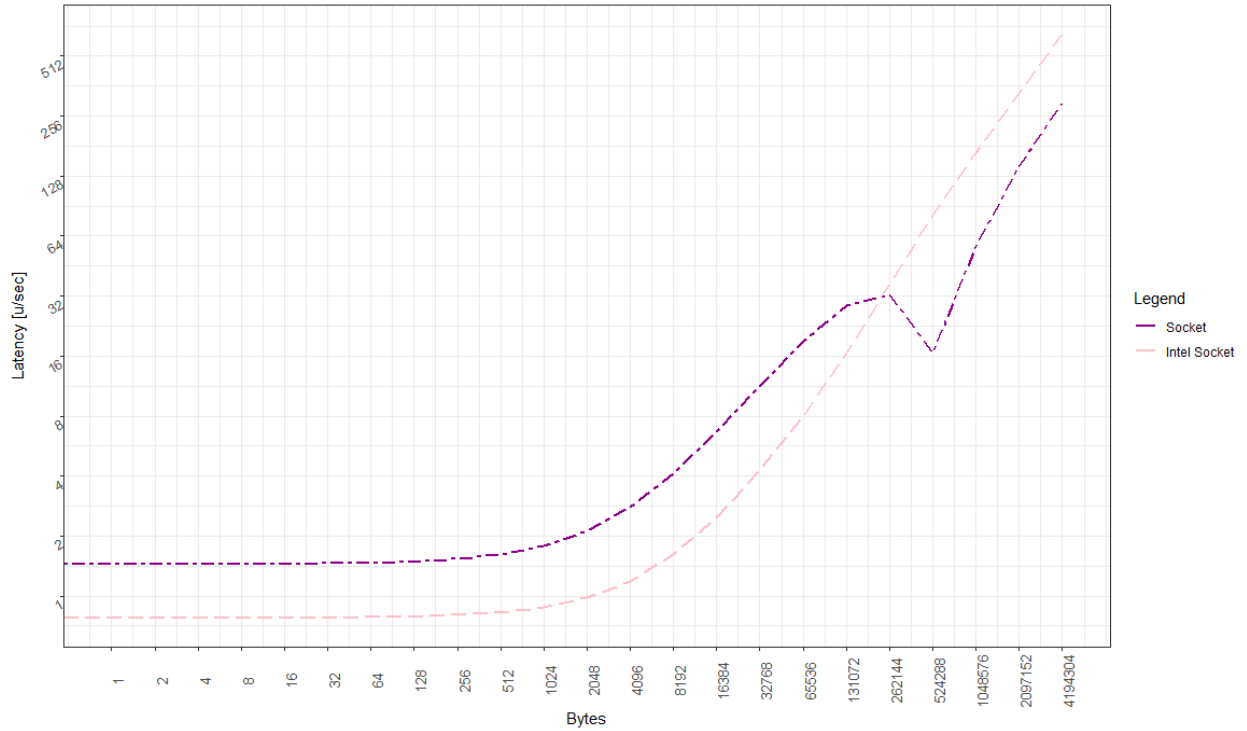


Figure 6: fitted model of latency computed for mapping by socket comparing GCC and INTEL modules

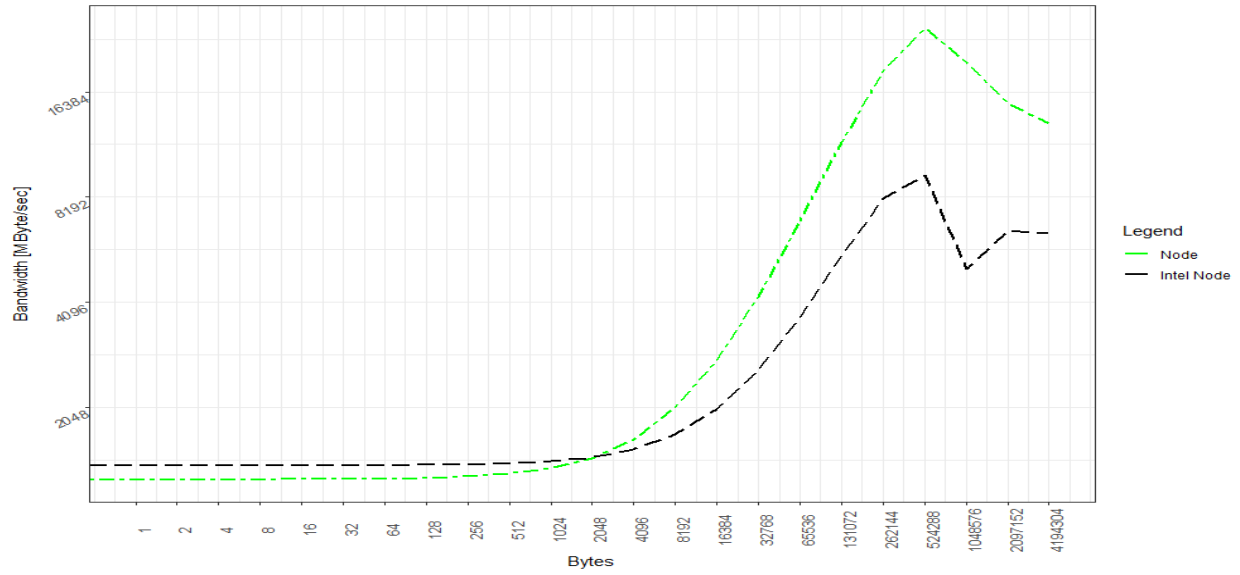


Figure 7: fitted model of Bandwidth computed for mapping by node comparing GCC and INTEL modules

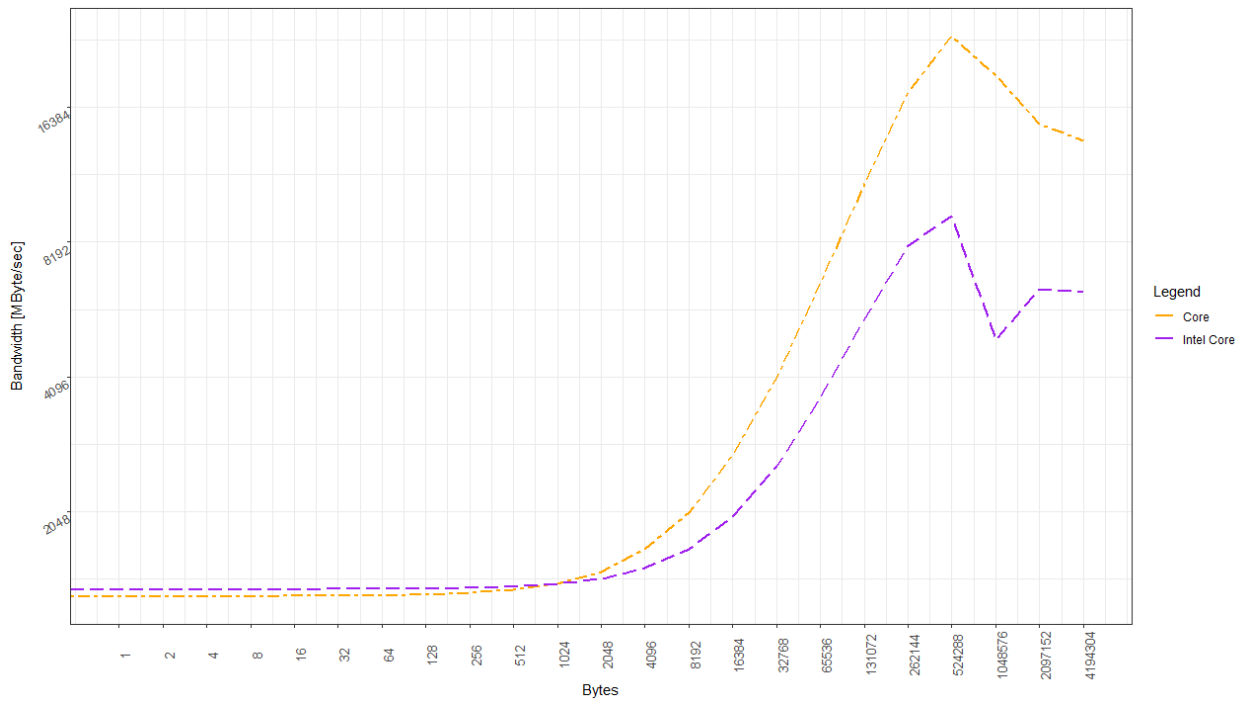


Figure 8: Fitted model of Bandwidth computed for mapping by core comparing GCC and INTEL modules

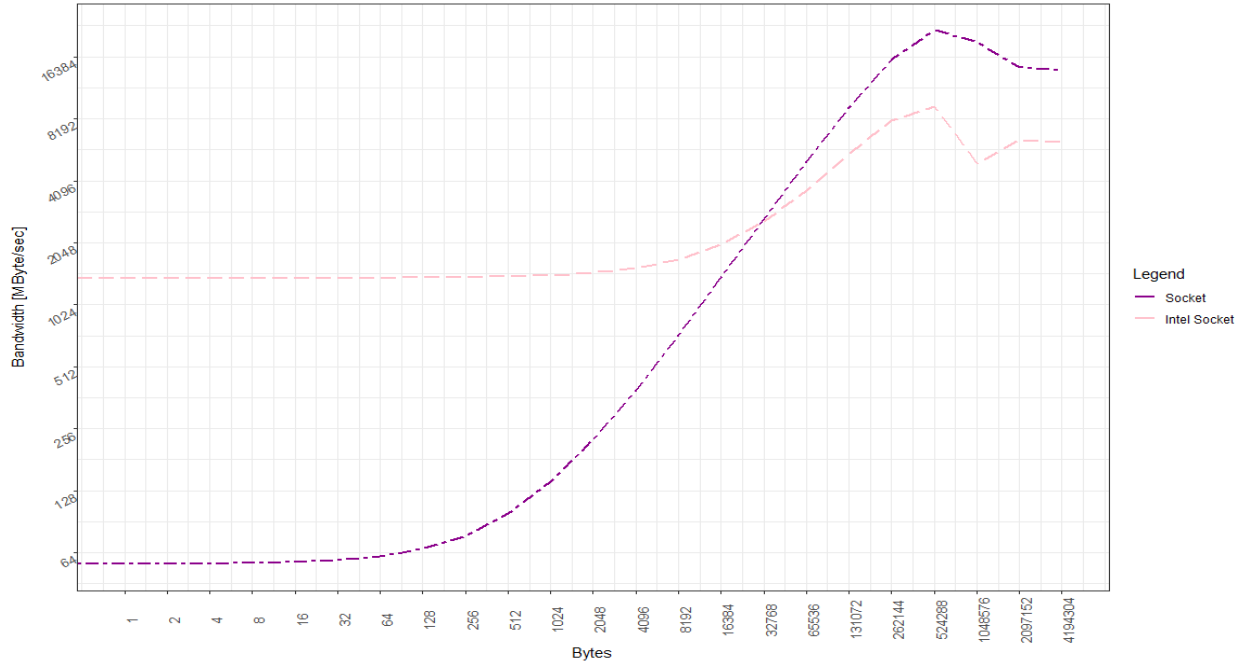


Figure 9: fitted model of Bandwidth computed for mapping by socket comparing GCC and INTEL modules

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
 - $B \rightarrow$ full-duplex bandwidth
 - $T_c(L,N) \rightarrow$ communication time
 - $T_s(L,N) \rightarrow$ raw compute time for all cell updates in a Jacobi sweep
 - $P_1(L) \rightarrow$ single-processor performance for a domain size L^3
 - $P(L,N) \rightarrow$ performance on $N = N_x N_y N_z$ processors for a particular overall problem size of $L^3 N$ grid points (using cubic subdomain of size L^3)
 - $L \rightarrow$ spatial dimension
 - $T_l \rightarrow$ latency of network
- Formulas used:

- $c(L, \vec{N}) = L^2 \cdot k \cdot 2 \cdot 8$
- $T_c(L, \vec{N}) = \frac{c(L, \vec{N})}{B} + kT_l$
- $P(L, \vec{N}) = \frac{L^3 N}{T_c(L, \vec{N}) + T_s(L)}$
- 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 is 0
 - 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	1	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	1	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	1	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
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