

# Multiple GPU programming with MPI



Norwegian research infrastructure services

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# Motivation

**Synchronous OpenACC/OpenMP offloading: **limited to single GPU.****

**Asynchronous OpenACC/OpenMP offloading: **multiple GPUs BUT limited to a single node.****

**Accelerating existing MPI-based codes.**

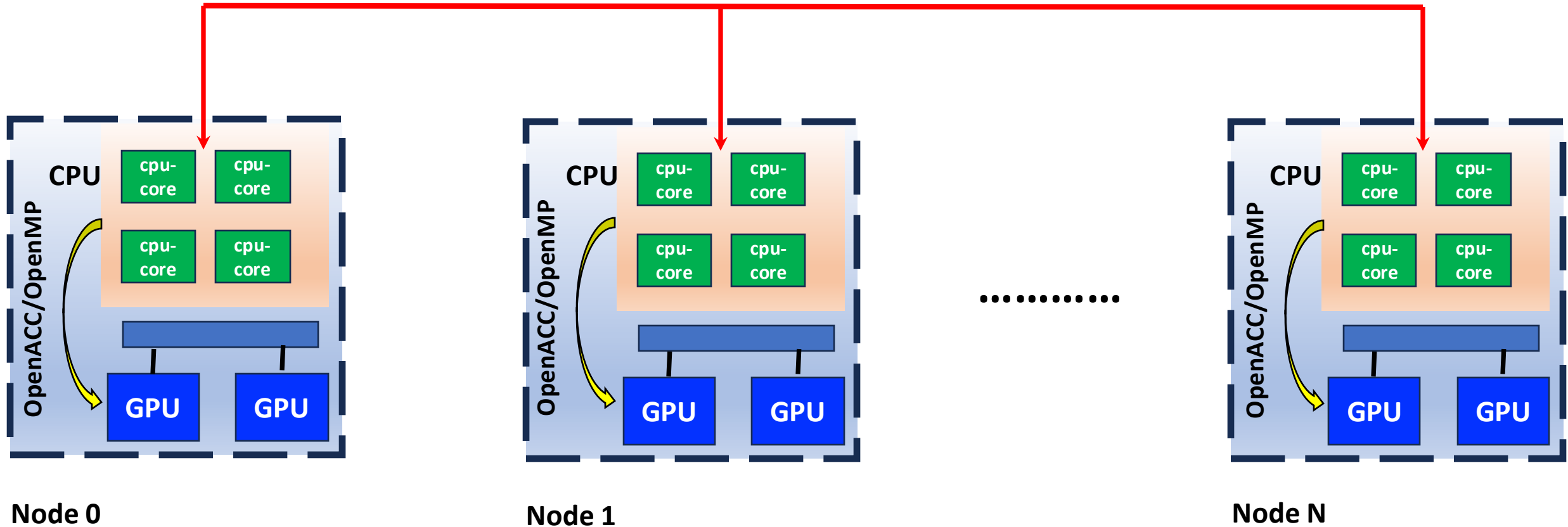
**Fully utilise the capacity of exascale supercomputer such as LUMI.**

**There is a need of multiple GPU programming.**

# Multiple GPU with MPI communications

Distributed computing

MPI communication library



# Outline

## □ Concept of:

- Traditional MPI without GPU awareness (with involvement of the CPU memory)
- MPI with GPU awareness (without involvement of the CPU memory)

# Outline

## ❑ Concept of:

- Traditional MPI without GPU awareness (with involvement of the CPU memory)
- MPI with GPU awareness (without involvement of the CPU memory)

## ❑ Application: MPI-OpenACC & MPI-OpenMP offloading

- Point-to-point communication (MPI\_Send & MPI\_Recv)

## ❑ Conclusion & Benchmark

## ❑ Hands-on examples and exercises

# Learning Outcomes

- ❑ How to assign each MPI rank to a GPU device ?
- ❑ How to combine MPI and OpenACC/OpenMP
  - MPI operations : **MPI\_Send and MPI\_Recv**
  - OpenACC directives: **update host(); update device()**
  - OpenMP directives: **update device() from(); update device() to()**
- ❑ How to perform MPI operations with GPU-awareness support
  - MPI operations : **MPI\_Send and MPI\_Recv**
  - OpenACC directive: **host\_data device\_ptr()**
  - OpenMP directive: **use\_device\_ptr()**

See our documentation for further details

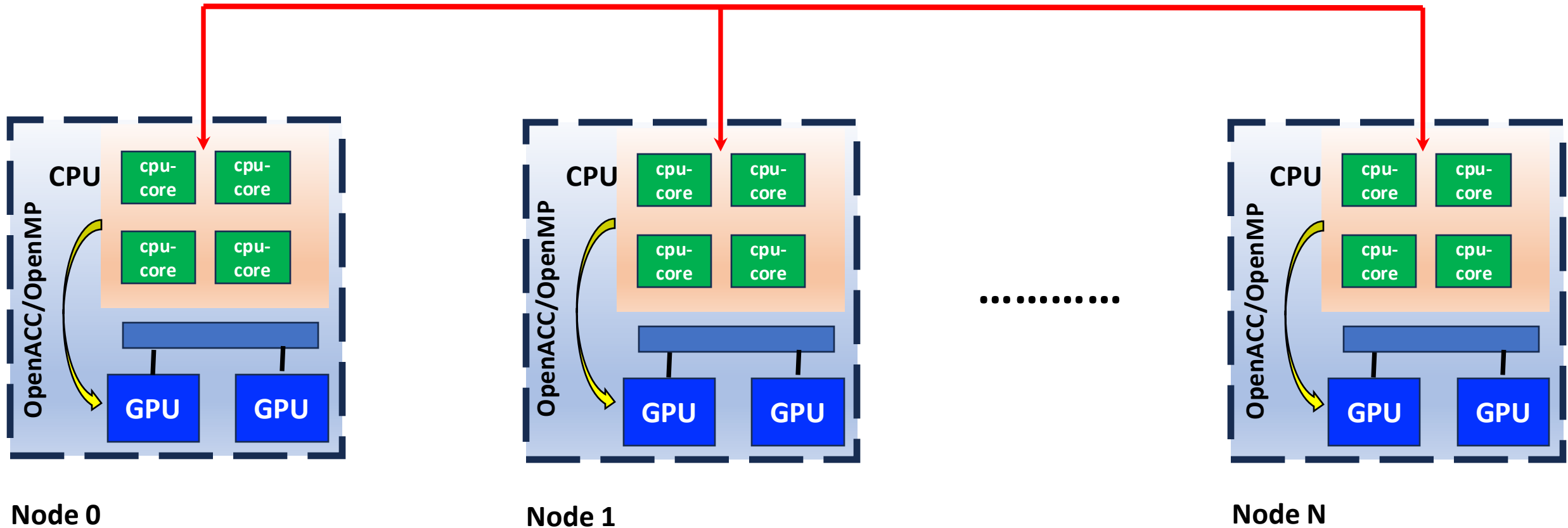
[https://enccs.github.io/gpu-programming/8-multiple\\_gpu/](https://enccs.github.io/gpu-programming/8-multiple_gpu/)

[https://documentation.sigma2.no/code\\_development/guides/gpuaware\\_mpi.html](https://documentation.sigma2.no/code_development/guides/gpuaware_mpi.html)

# Multiple GPU with MPI communications

Distributed computing

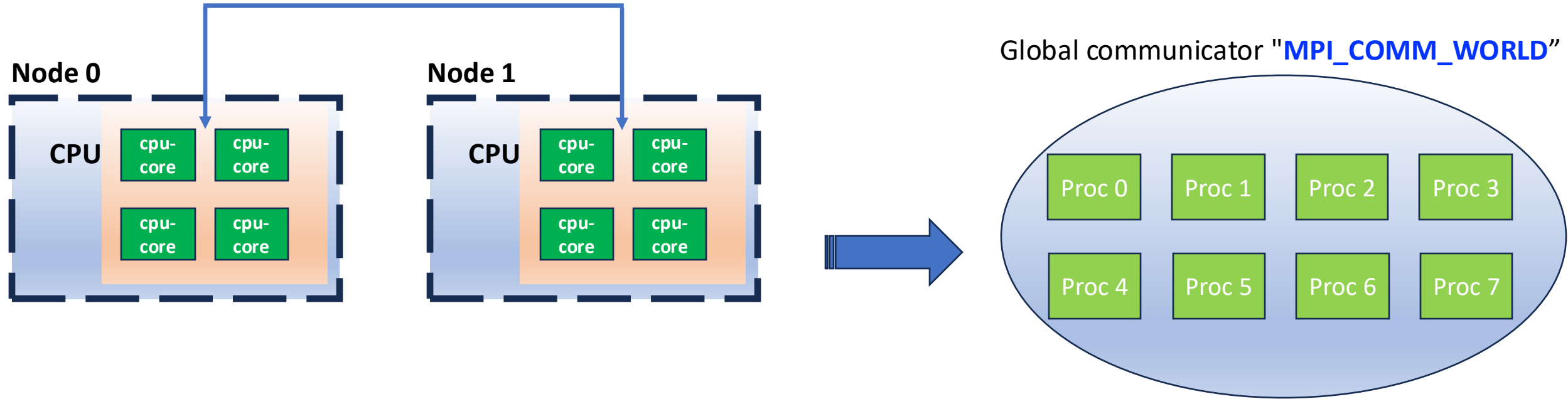
MPI communication library



How to assign each MPI rank to a GPU device ?

# How to assign each MPI rank to a GPU device ?

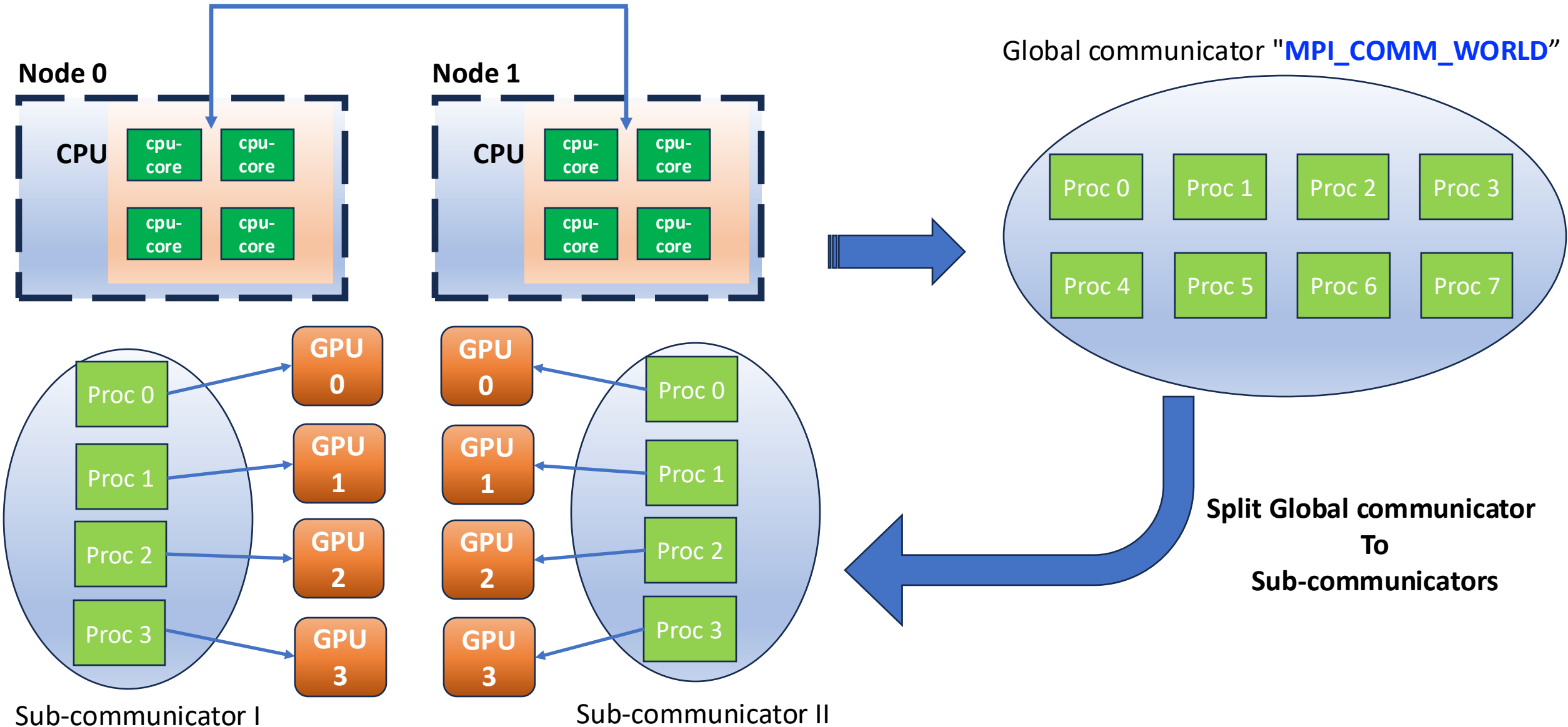
Split the global communicator "**MPI\_COMM\_WORLD**" into sub-communicators





# How to assign each MPI rank to a GPU device ?

Split the global communicator "**MPI\_COMM\_WORLD**" into sub-communicators



# How to assign each MPI rank to a GPU device ?

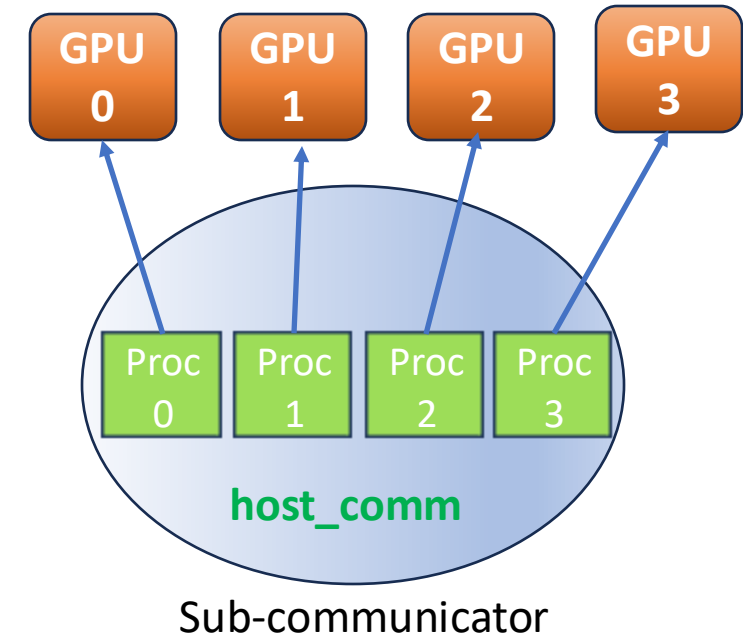
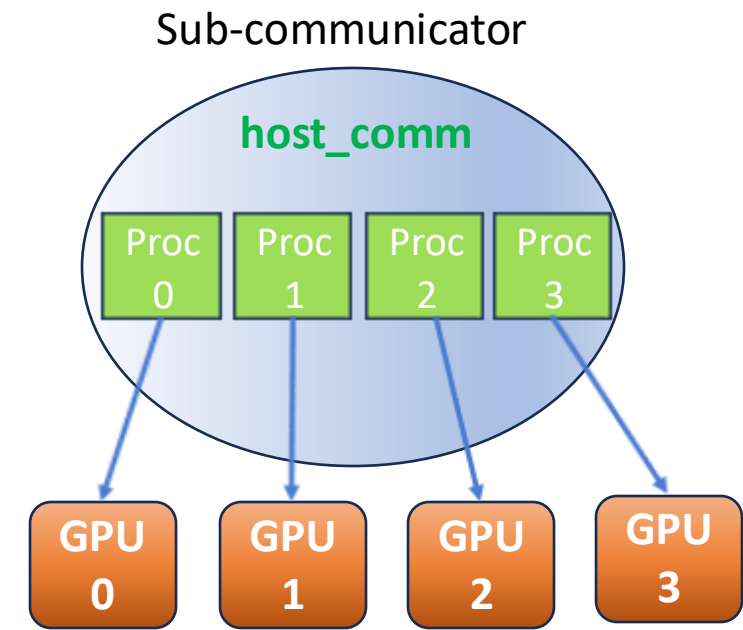
## MPI functions

```
call MPI_COMM_SPLIT_TYPE(MPI_COMM_WORLD,  
                        MPI_COMM_TYPE_SHARED, 0,  
                        MPI_INFO_NULL, host_comm, ierr)
```

```
call MPI_COMM_RANK(host_comm, host_rank, ierr)
```

Sub-communicator

New MPI rank



# How to assign each MPI rank to a GPU device ?

## MPI functions

```
call MPI_COMM_SPLIT_TYPE(MPI_COMM_WORLD,  
                        MPI_COMM_TYPE_SHARED, 0,  
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```

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call MPI_COMM_RANK(host_comm, host_rank, ierr)
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Sub-communicator

New MPI rank

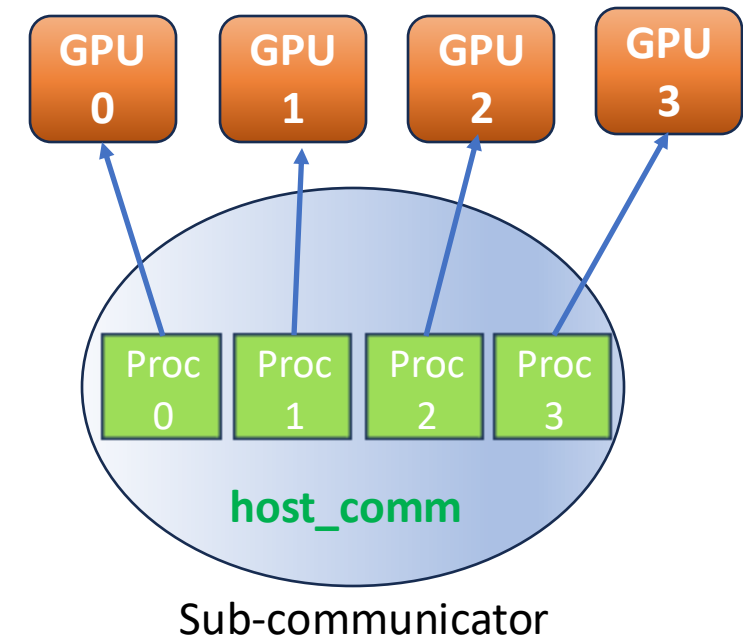
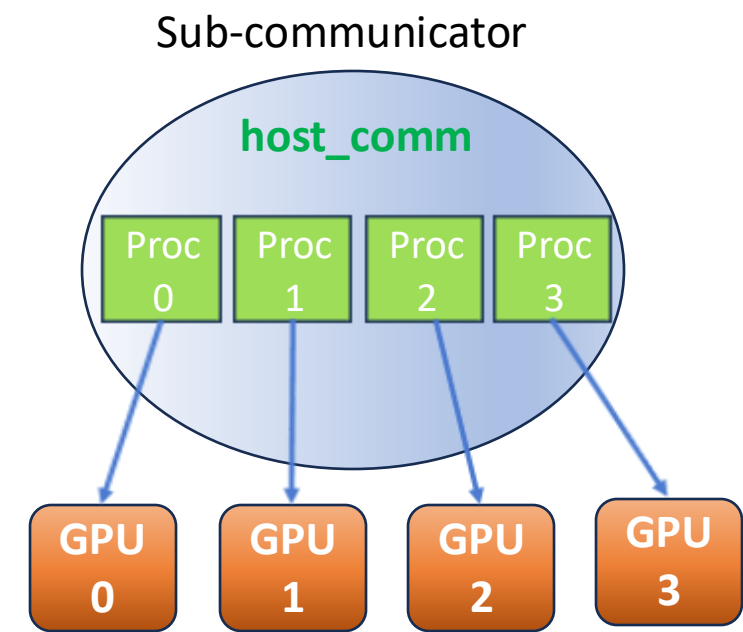
## API functions: **OpenACC**

! Sets the device INDEX and the device type to be used

```
call acc_set_device_num(host_rank, acc_get_device_type())
```

! Returns the number of devices available on the host (here on each node)

```
numDevice = acc_get_num_devices(acc_get_device_type())
```



# How to assign each MPI rank to a GPU device ?

```
call MPI_COMM_SPLIT_TYPE(MPI_COMM_WORLD,  
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                        MPI_INFO_NULL, host_comm, ierr)
```

**MPI functions**

```
call MPI_COMM_RANK(host_comm, host_rank, ierr)
```

## API functions: **OpenACC**

! Sets the device INDEX and the device type to be used

```
call acc_set_device_num(host_rank, acc_get_device_type())
```

! Returns the number of devices available on the host (here on each node)

```
numDevice = acc_get_num_devices(acc_get_device_type())
```

## API functions: **OpenMP**

! Sets the device INDEX and the device type to be used

```
call omp_set_default_device(host_rank)
```

! Returns the number of devices available on the host (here on each node)

```
numDevice = omp_get_num_devices()
```

# Hands-on Example 1:

## How to assign each MPI rank to a GPU device ?

- Download the repo

```
$ git clone https://github.com/HichamAgueny/multiGPU\_MPI\_examples
```

```
$ cd multiGPU_MPI_examples
```

- For MPI-OpenACC

```
$ cd example_1/setDevice_acc
```

- For MPI-OpenMP

```
$ cd example_1/setDevice_omp
```

- To compile and execute the code

**Load the LUMI software stack**

```
$ module load LUMI/24.03 partition/G
```

```
$ module load cpeCray
```

**Compile:** \$ ./compile.sh

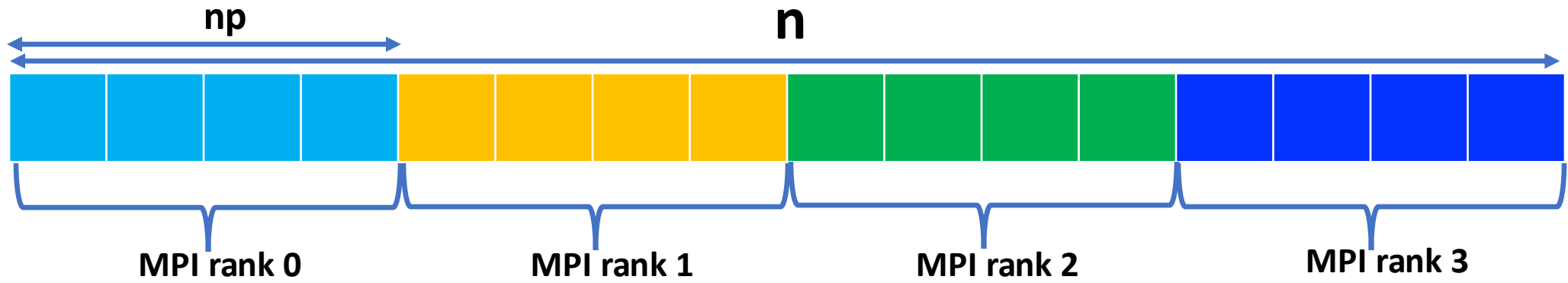
**Submit a job:** \$ sbatch script.slurm

**View the output file:** \$ vi setDevice\_accxxxxxx.out

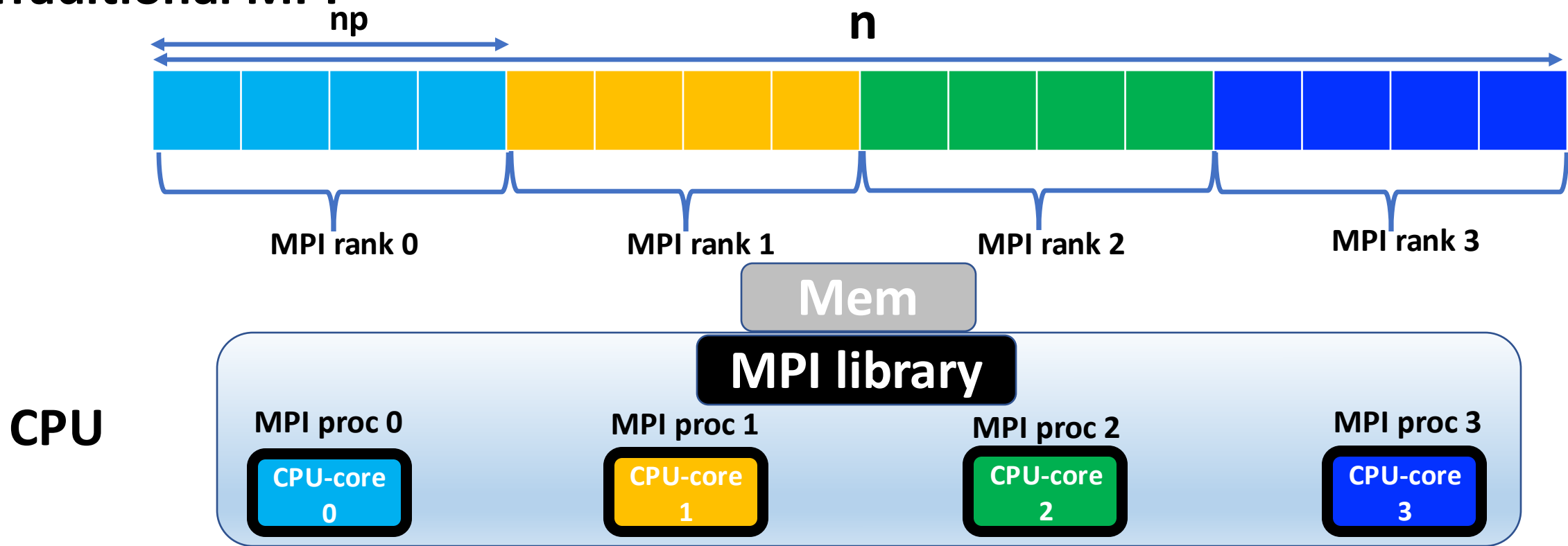
**Concept:**

**Traditional MPI without GPU awareness**

# Traditional MPI

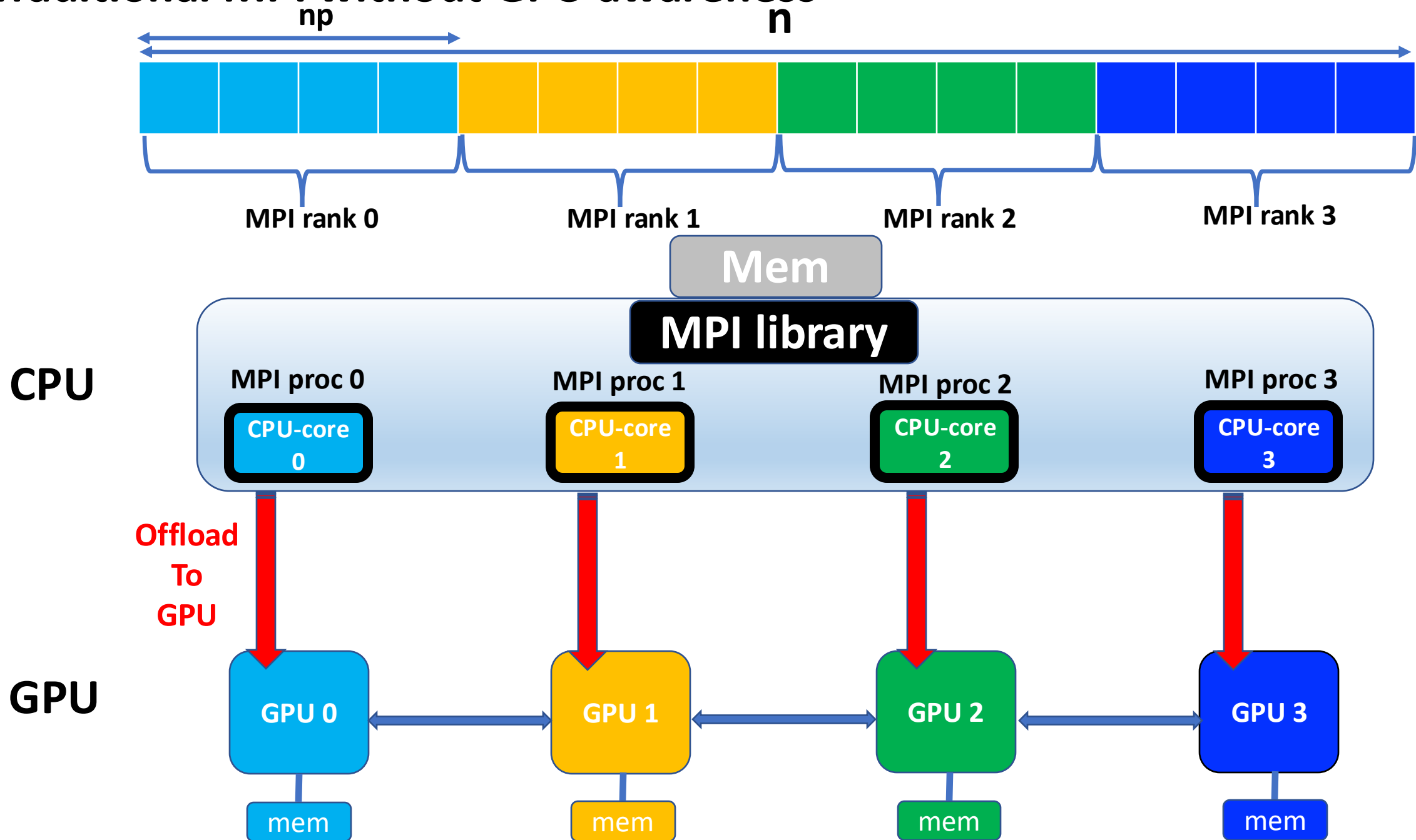


# Traditional MPI

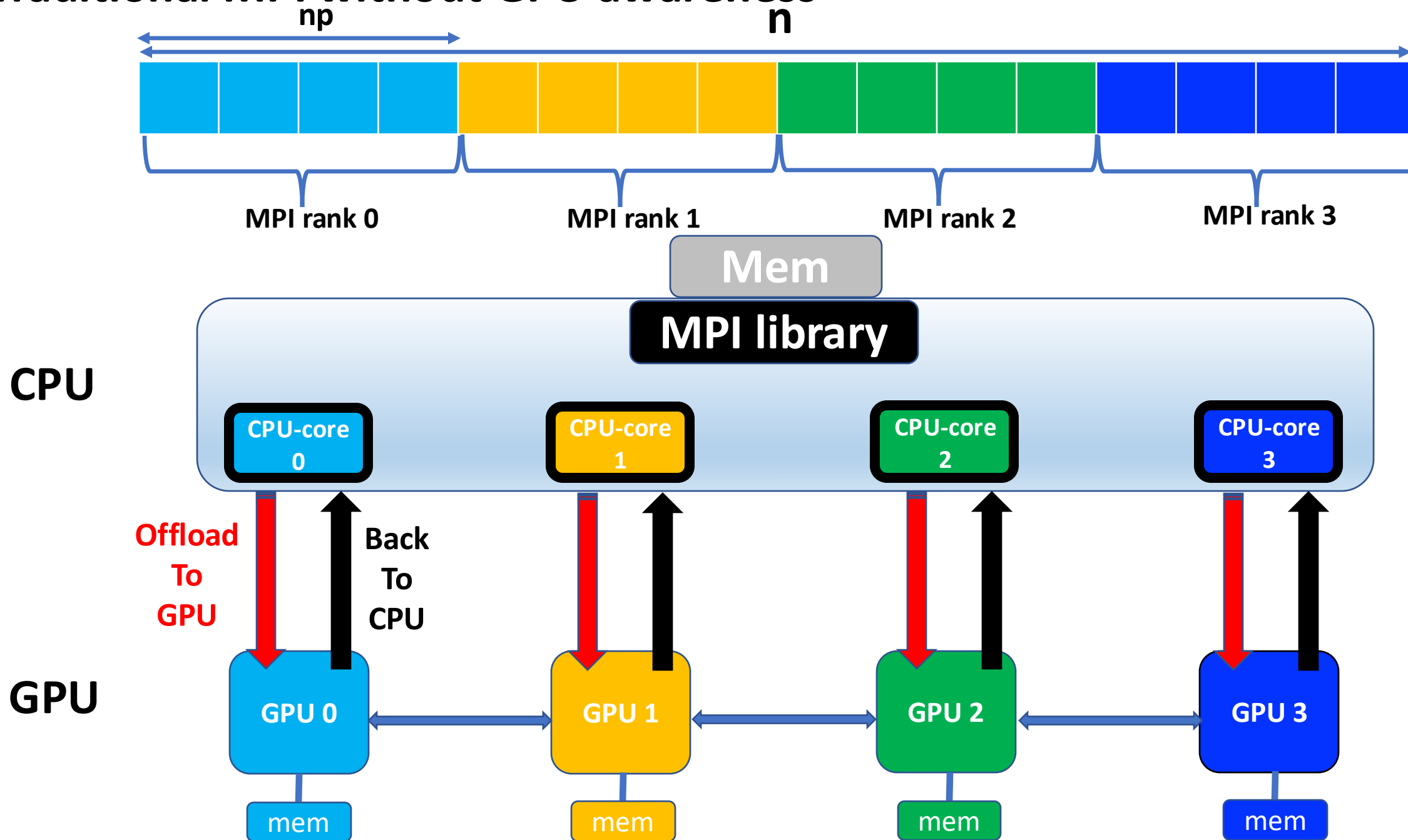




# Traditional MPI without GPU awareness



# Traditional MPI without GPU awareness



## Example: Combining MPI with OpenACC/OpenMP APIs

- MPI operations : `MPI_Send` & `MPI_Recv`
- OpenACC directives: `update host(); update device()`
- OpenMP directives: `update device() from(); update device() to()`

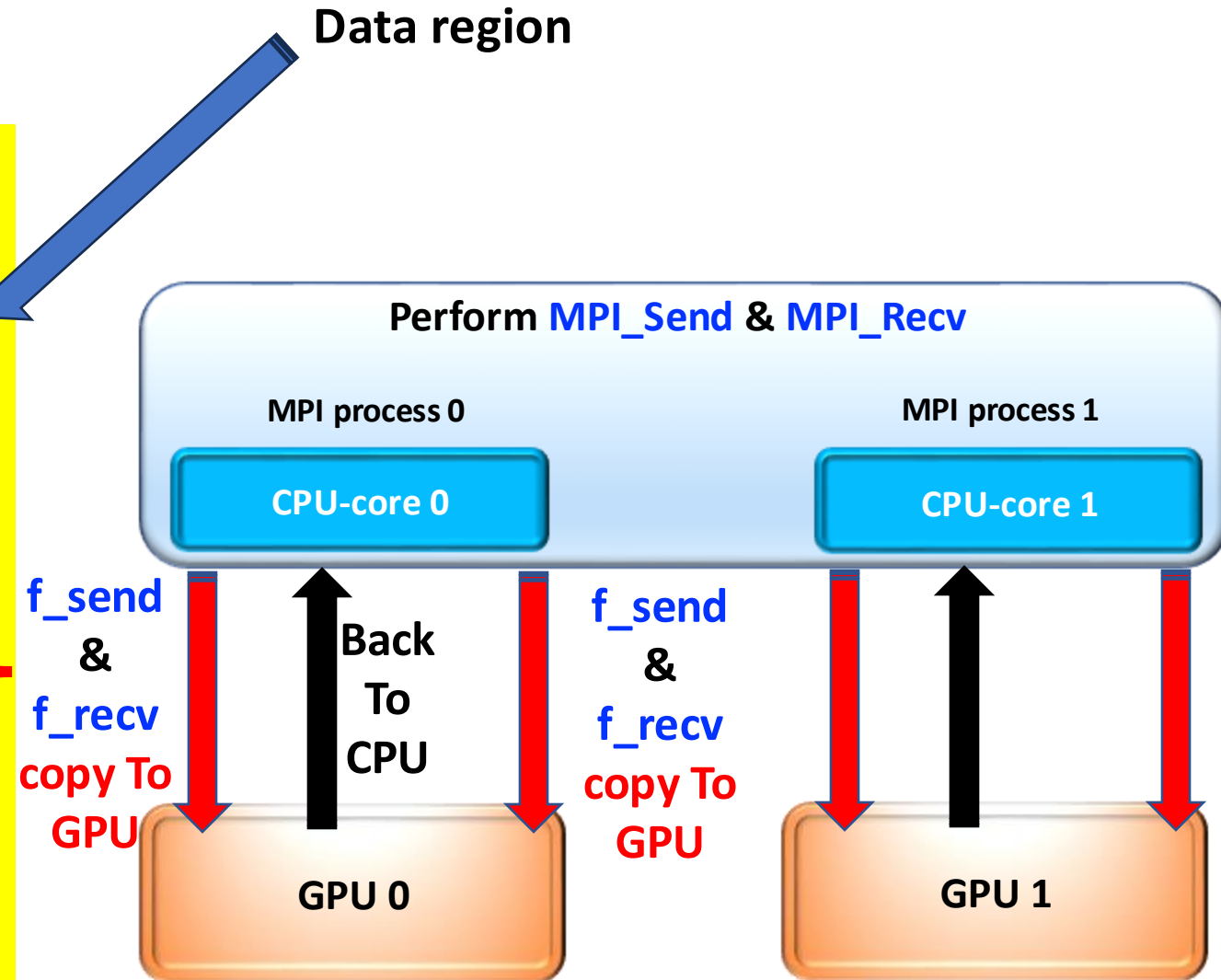
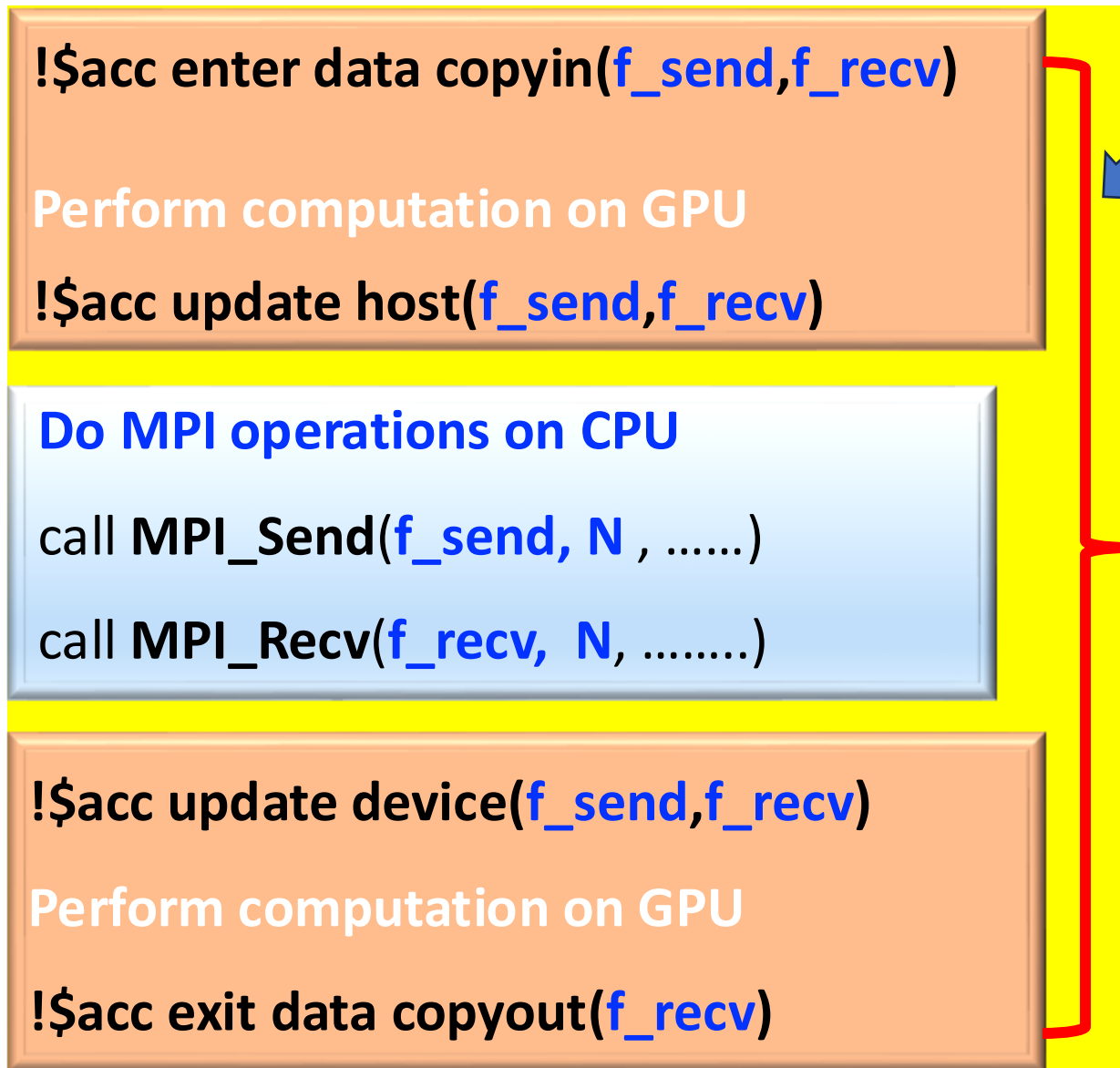
# Scenario with only 2 MPI-processes: MPI\_Send & MPI\_Recv



**Purpose:** To measure the time it takes to transfer data during **MPI communication**

```
if(MPIrank.eq.0) then
  call MPI_Send(f_send(:), N, MPI_DOUBLE_PRECISION, MPIrank=1, tag1, ..)
  call MPI_Recv(f_recv(:), N, MPI_DOUBLE_PRECISION, MPIrank=1, tag2, ..)
endif
if(MPIrank.eq.1) then
  call MPI_Recv(f_recv(:), N, MPI_DOUBLE_PRECISION, MPIrank=0, tag1, ..)
  call MPI_Send(f_send(:), N, MPI_DOUBLE_PRECISION, MPIrank=0, tag2, ..)
endif
```

# MPI-OpenACC: MPI\_Send & MPI\_Recv



## MPI-OpenACC

!\$acc enter data **copyin**(f\_send,f\_recv)

Perform computation on GPU

!\$acc **update host**(f\_send,f\_recv)

**Do MPI operations on CPU**

call MPI\_Send(f\_send, N , .....)

call MPI\_Recv(f\_recv, N , .....)

!\$acc update device(f\_send,f\_recv)

Perform computation on GPU

!\$acc exit data **copyout**(f\_recv)

## MPI-OpenMP

!\$omp target enter data **map**(to: f\_send,f\_recv)

Perform computation on GPU

!\$omp target **update from**(f\_send,f\_recv)

**Do MPI operations on CPU**

call MPI\_Send(f\_send, N , .....)

call MPI\_Recv(f\_recv, N , .....)

!\$omp target update to(f\_send,f\_recv)

Perform computation on GPU

!\$omp target exit data **map**(from: f\_recv)

# Hands-on Example 2: Traditional MPI with OpenACC/OpenMP ?

**Purpose:** To measure the time it takes to transfer data during **MPI communication**

- **Download the repo**

```
$ git clone https://github.com/HichamAgueny/multiGPU\_MPI\_examples
```

```
$ cd multiGPU_MPI_examples
```

- **For MPI-OpenACC**

```
$ cd example_2/mpiacc
```

- **For MPI-OpenMP**

```
$ cd example_2/mpiomp
```

- **To compile and execute the code**

**Load the LUMI software stack**

```
$ module load LUMI/24.03 partition/G
```

**Compile:** \$ ./compile.sh

**Submit a job:** \$ sbatch script.slurm

**View the output file:** \$ vi staging\_mpiacc-xxxxxx.out



# Output from running the code `example_2/mpiacc/mpiacc_staging.f90`

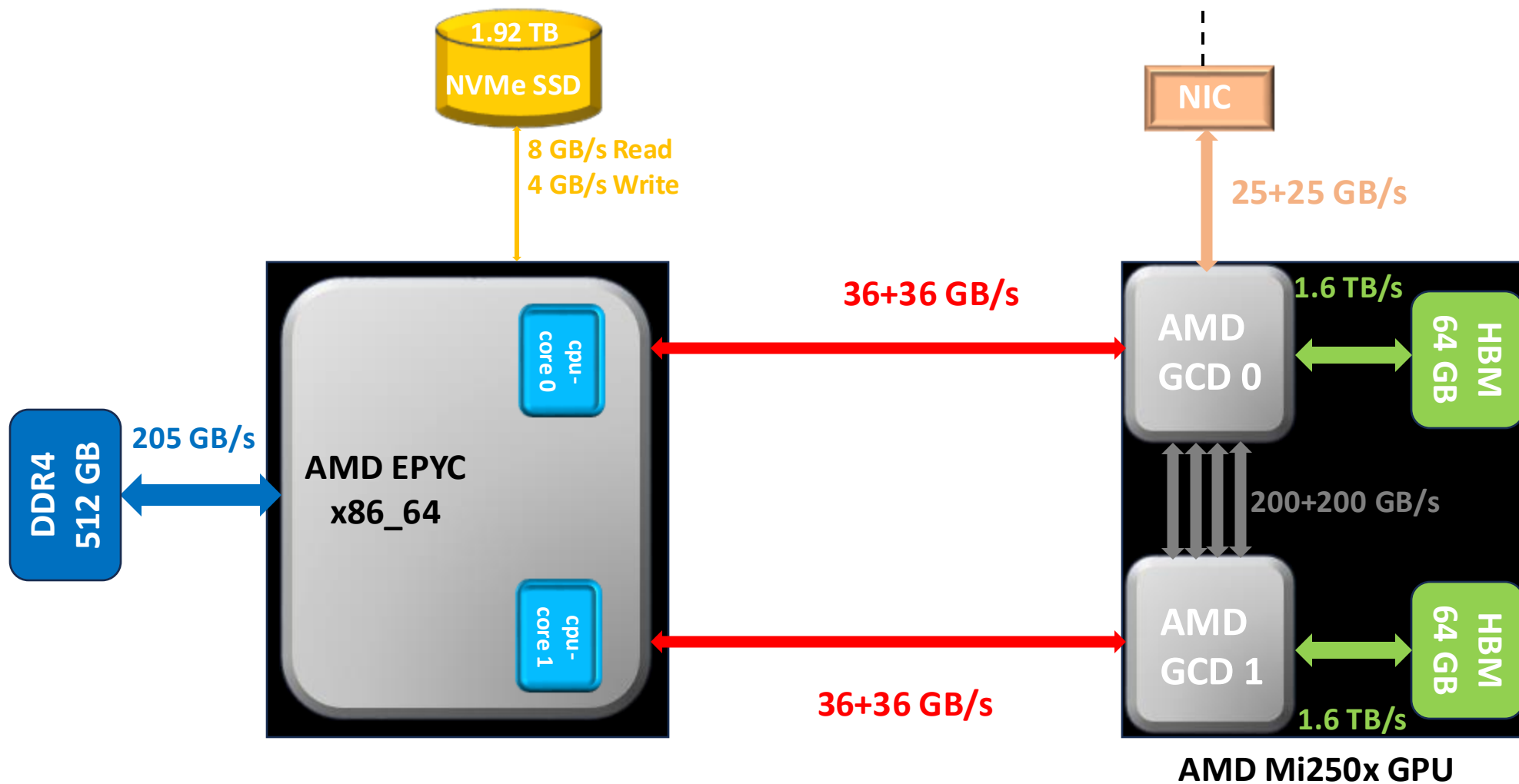
Measruing bandwidth: 2 MPI processes and 2 GPUs, single Node

--Time (s)	0.00002	Data size (B)	128	Bandwidth (GBps)	0.01187
--Time (s)	0.00002	Data size (B)	256	Bandwidth (GBps)	0.02398
--Time (s)	0.00003	Data size (B)	512	Bandwidth (GBps)	0.04033
--Time (s)	0.00003	Data size (B)	1024	Bandwidth (GBps)	0.08017
--Time (s)	0.00002	Data size (B)	2048	Bandwidth (GBps)	0.16789
--Time (s)	0.00003	Data size (B)	4096	Bandwidth (GBps)	0.27093
--Time (s)	0.00003	Data size (B)	8192	Bandwidth (GBps)	0.57978
--Time (s)	0.00003	Data size (B)	16384	Bandwidth (GBps)	1.05693
--Time (s)	0.00004	Data size (B)	32768	Bandwidth (GBps)	1.56283
--Time (s)	0.00006	Data size (B)	65536	Bandwidth (GBps)	2.12329
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--Time (s)	0.00066	Data size (B)	1048576	Bandwidth (GBps)	3.15746
--Time (s)	0.00112	Data size (B)	2097152	Bandwidth (GBps)	3.73963
--Time (s)	0.00203	Data size (B)	4194304	Bandwidth (GBps)	4.12282
--Time (s)	0.00438	Data size (B)	8388608	Bandwidth (GBps)	3.82907
--Time (s)	0.00733	Data size (B)	16777216	Bandwidth (GBps)	4.58024
--Time (s)	0.01343	Data size (B)	33554432	Bandwidth (GBps)	4.99637
--Time (s)	0.02581	Data size (B)	67108864	Bandwidth (GBps)	5.19988
--Time (s)	0.05028	Data size (B)	134217728	Bandwidth (GBps)	5.33868
--Time (s)	0.09886	Data size (B)	268435456	Bandwidth (GBps)	5.43059
--Time (s)	0.19354	Data size (B)	536870912	Bandwidth (GBps)	5.54795
--Time (s)	0.38534	Data size (B)	1073741824	Bandwidth (GBps)	5.57296

The speed of transferring **1 GB** of data between 2 MPI-processes is about **6 GB/s**

Too slow!!



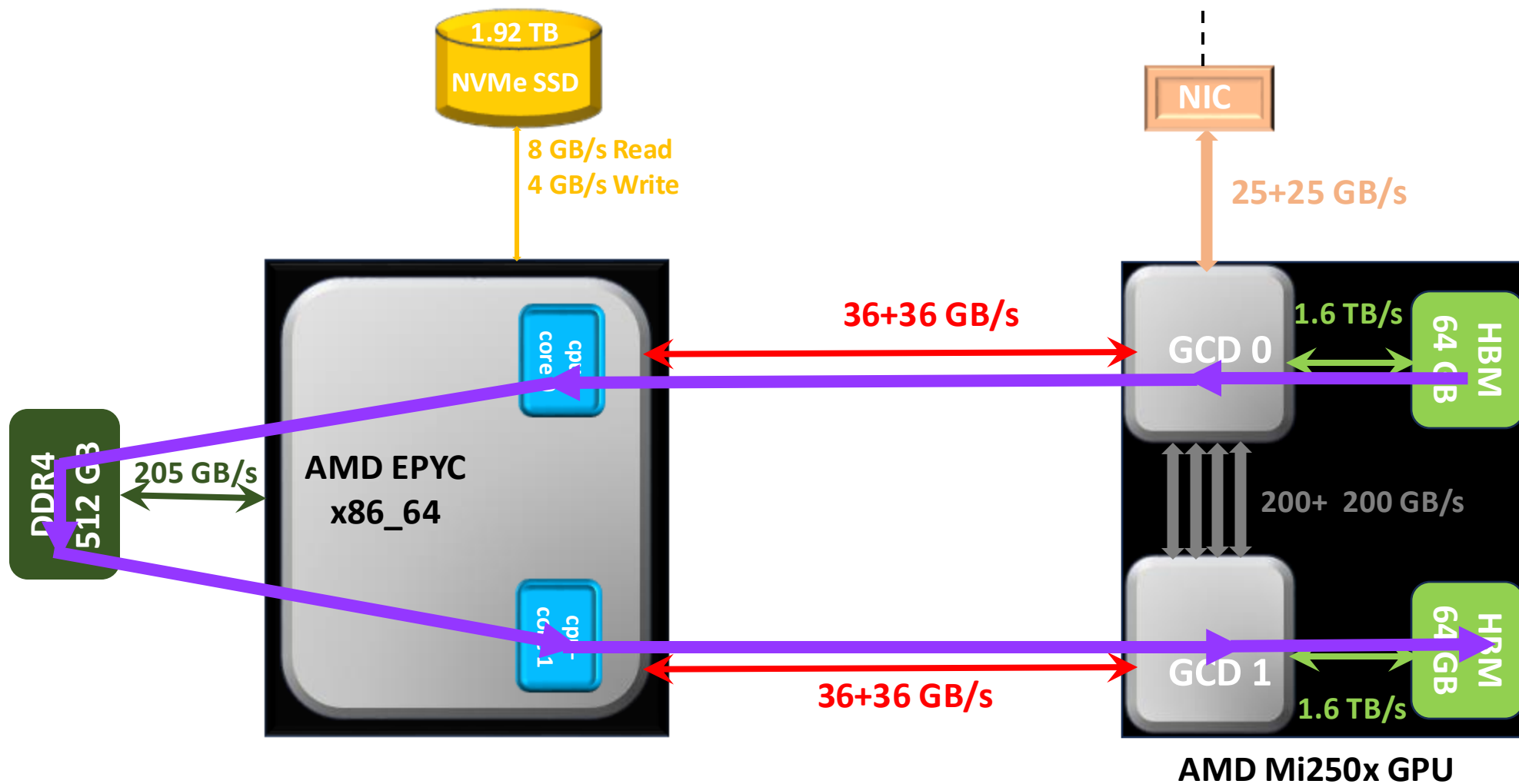


### Bidirectional bandwidths

	Infinity Fabric	36 + 36	GB/s
	Infinity Fabric	200 + 200	GB/s
	PCIe Gen4 ESM	50 + 50	GB/s
	PCIe Gen4	8 + 8	GB/s
	Ethernet	25 + 25	GB/s

GCD === GPU

*Bandwidth values are taken from LUMI doc*

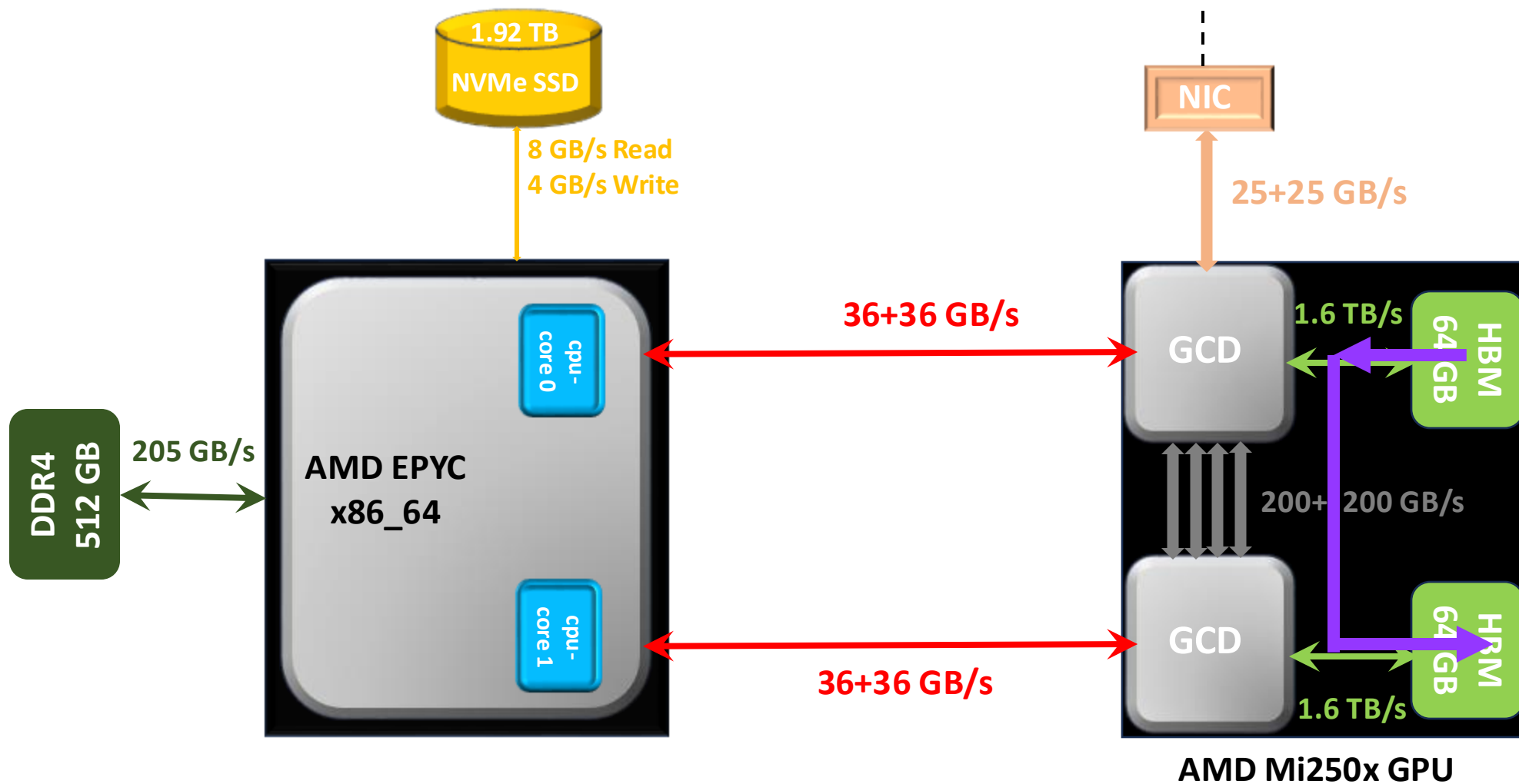


### Bidirectional bandwidths

	Infinity Fabric	36 + 36	GB/s
	Infinity Fabric	200 + 200	GB/s
	PCIe Gen4 ESM	50 + 50	GB/s
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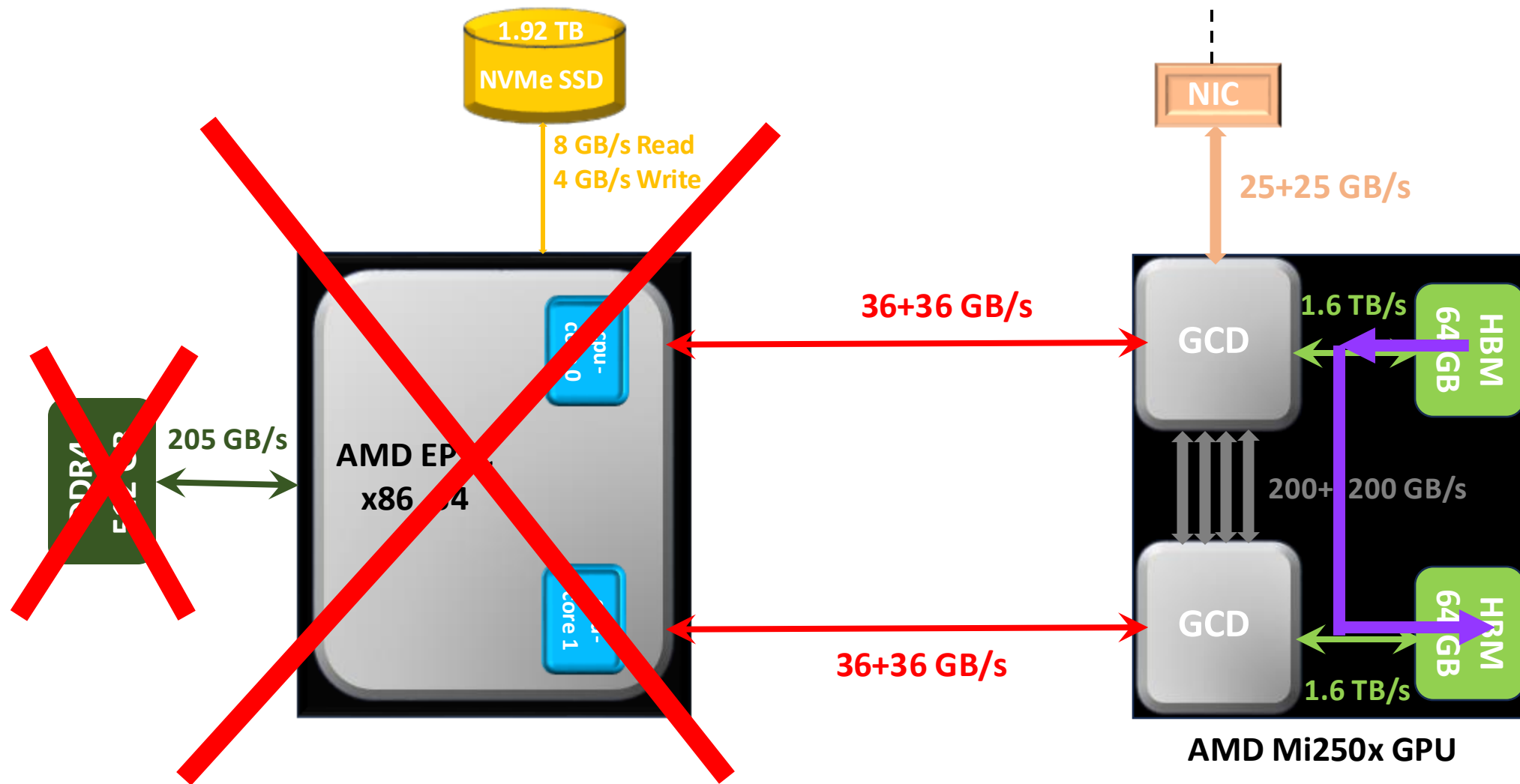
AMD Mi250x GPU

GCD === XGPU



### Bidirectional bandwidths

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### Bidirectional bandwidths

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AMD Mi250x GPU

GCD === XGPU

**Concept:**

**MPI with GPU awareness support**

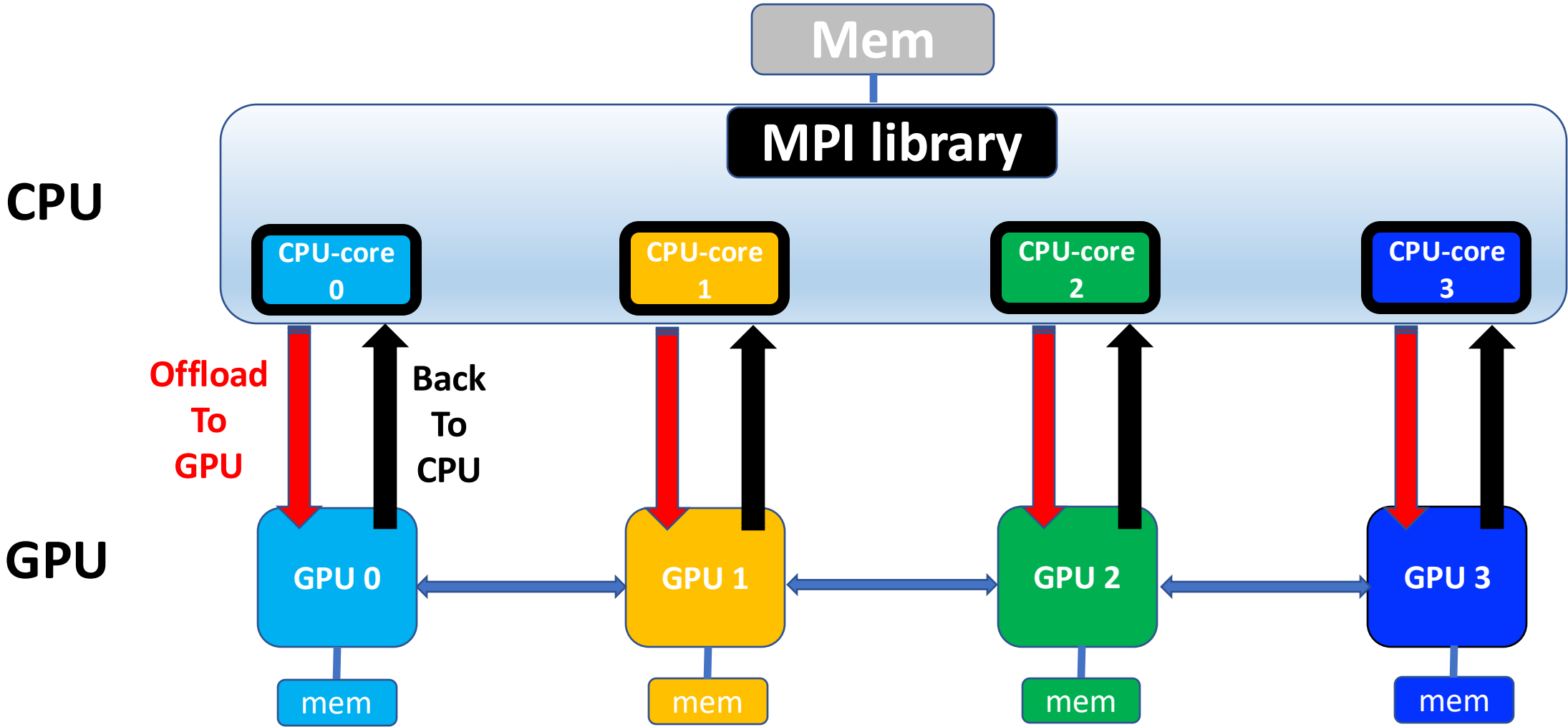
# What is GPU-aware MPI ?

In GPU-aware MPI concept:

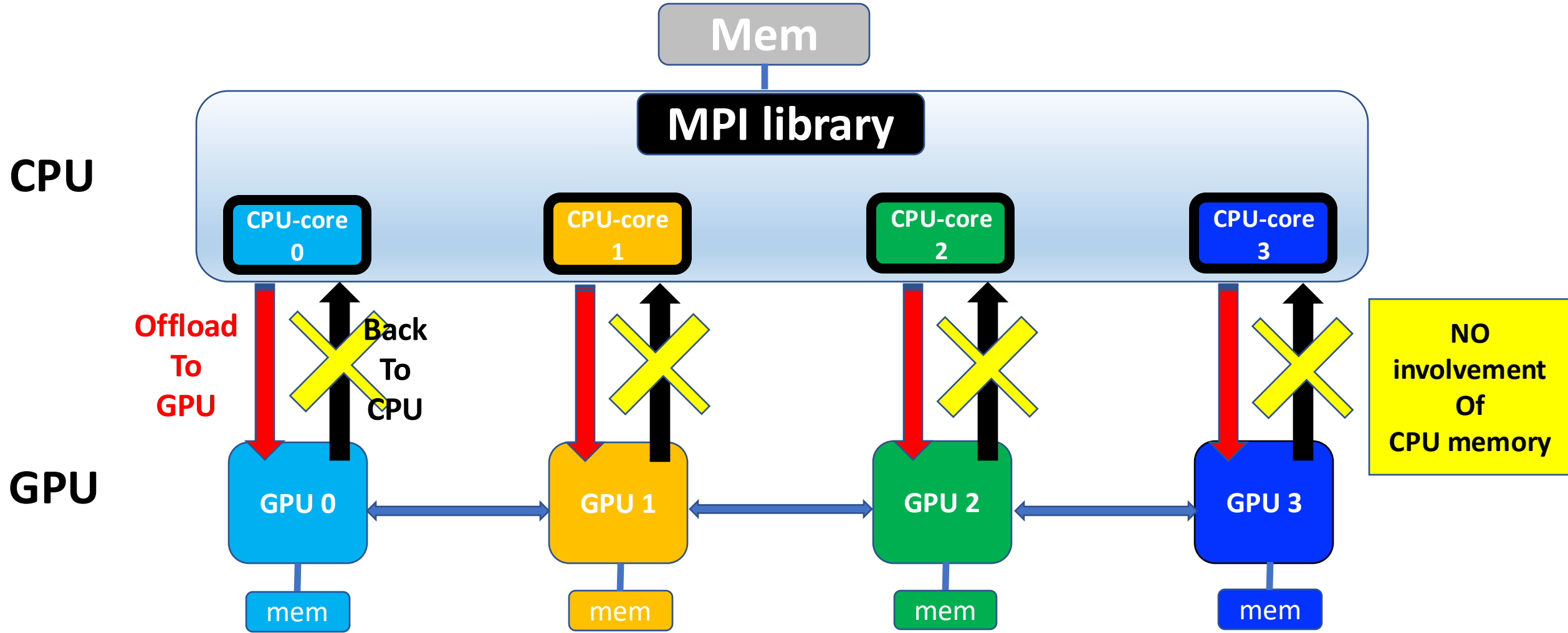
**MPI library** can **directly access the GPU memory** **without** necessarily **passing by the CPU memory**.

Device pointers are passed as arguments to an MPI routine

# Traditional MPI without GPU awareness



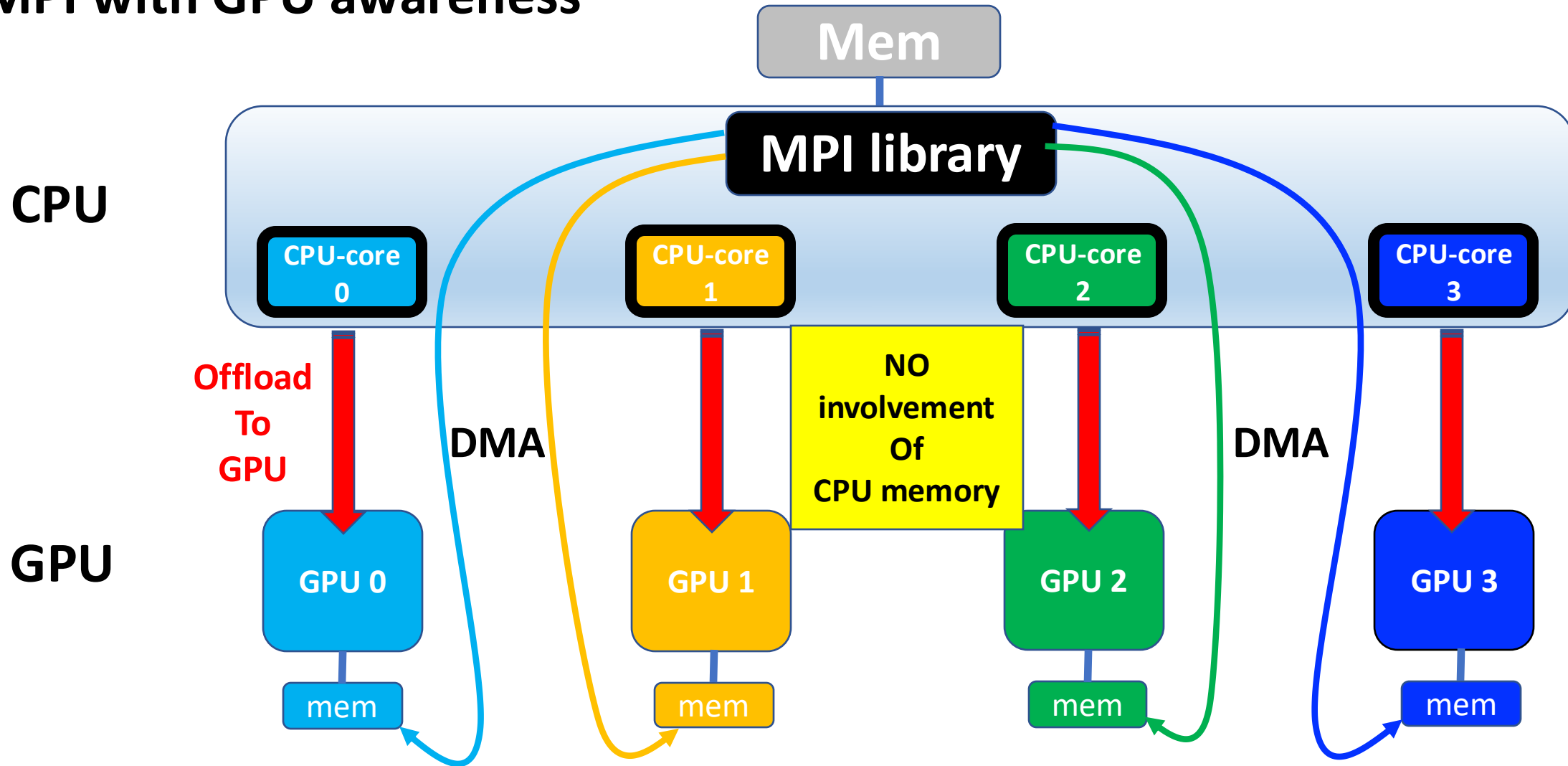
# MPI with GPU awareness



**MPI library** directly access GPU memory without passing by the CPU memory



# MPI with GPU awareness



Device pointers are passed as arguments to an MPI routine

## Example: GPU-aware MPI with OpenACC/OpenMP APIs

- MPI operations : `MPI_Send`, `MPI_Recv` and `MPI_Allreduce`
- OpenACC directive: `host_data device_ptr()`
- OpenMP directive: `use_device_ptr()`

# Scenario with only 2 MPI-processes: MPI\_Send & MPI\_Recv



**Purpose:** To measure the time it takes to transfer data during **MPI communication**

```
if(MPIrank.eq.0) then
  call MPI_Send(f_send(:), N, MPI_DOUBLE_PRECISION, MPIrank=1, tag1, ..)
  call MPI_Recv(f_recv(:), N, MPI_DOUBLE_PRECISION, MPIrank=1, tag2, ..)
endif
if(MPIrank.eq.1) then
  call MPI_Recv(f_recv(:), N, MPI_DOUBLE_PRECISION, MPIrank=0, tag1, ..)
  call MPI_Send(f_send(:), N, MPI_DOUBLE_PRECISION, MPIrank=0, tag2, ..)
endif
```

# GPU-aware MPI with OpenACC: MPI\_Send & MPI\_Recv

```
!$acc enter data copyin(f_send,f_recv)  
Do something on GPU
```

*!Device pointers f\_send & f\_recv are passed to MPI\_send & MPI\_recv*

```
!$acc host_data use_device(f_send,f_recv)
```

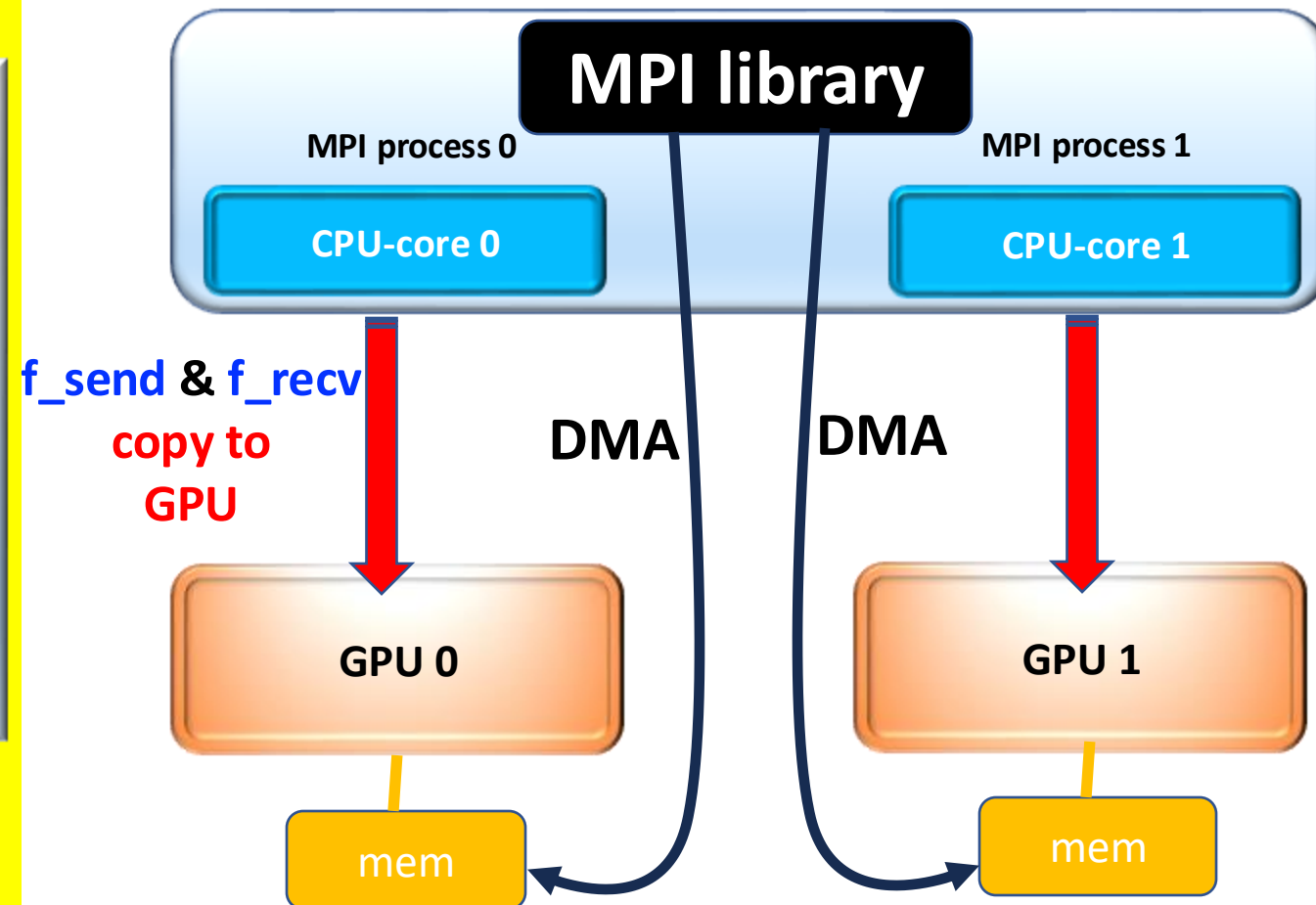
```
call MPI_Send(f_send, N , .....)
```

```
call MPI_Recv(f_recv, N, .....)
```

```
!$acc end host_data
```

```
Do something on GPU
```

```
!$acc exit data copyout(f_recv)
```



Communication can be seen as it is **between a pair of GPUs**

# GPU-aware MPI with OpenACC: MPI\_Send & MPI\_Recv

```
!$acc enter data copyin(f_send,f_recv)  
Do something on GPU
```

*!Device pointers f\_send & f\_recv are passed to MPI\_send & MPI\_recv*

```
!$acc host_data use_device(f_send,f_recv)
```

```
call MPI_Send(f_send, N , .....)
```

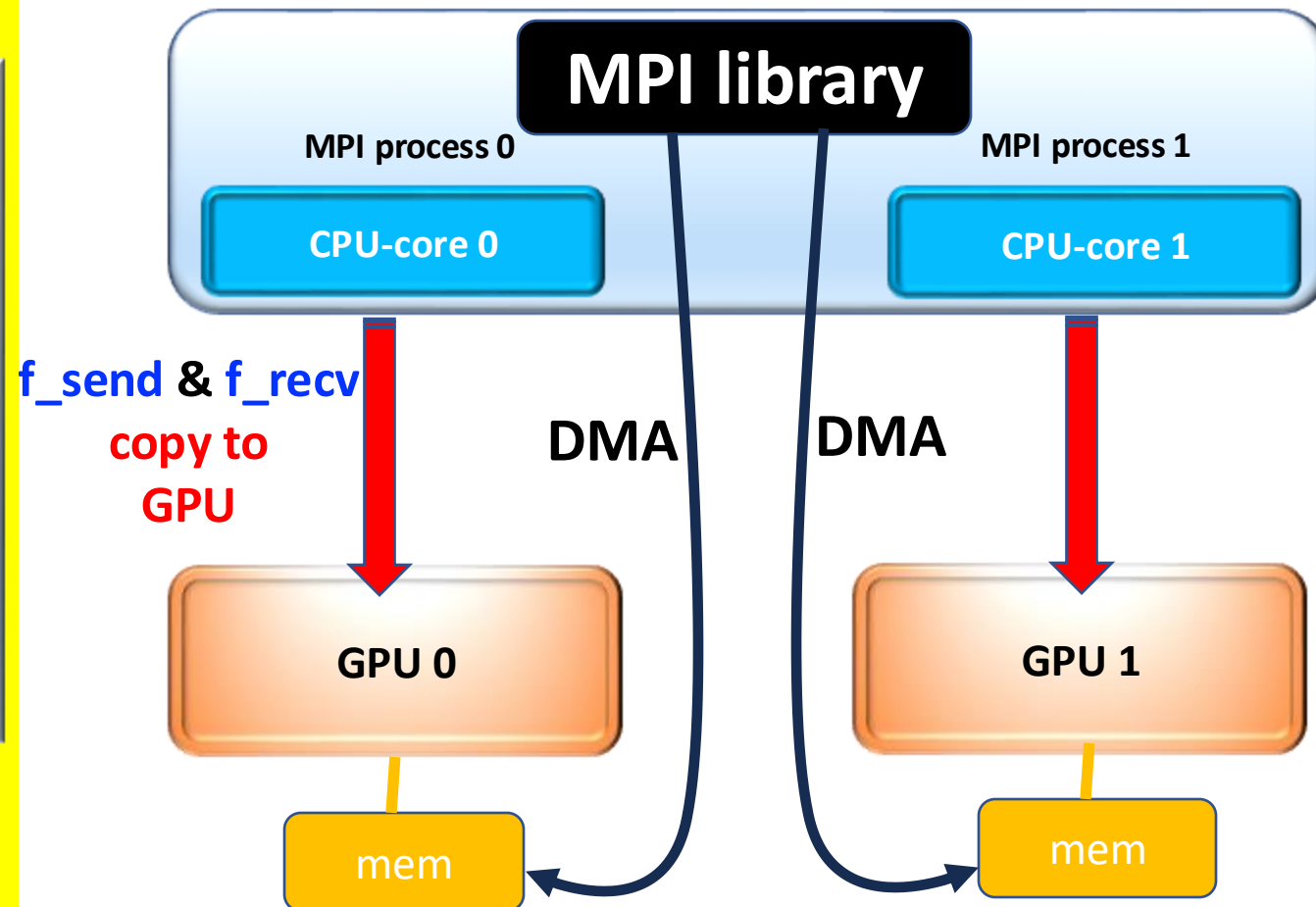
```
call MPI_Recv(f_recv, N , .....)
```

```
!$acc end host_data
```

```
Do something on GPU
```

```
!$acc exit data copyout(f_recv)
```

To enable GPU-aware support in MPICH lib  
`$ export MPICH_GPU_SUPPORT_ENABLED=1`



Communication can be seen as it is **between a pair of GPUs**

## GPU-aware MPI with OpenACC

**!\$acc enter data copyin(f\_send,f\_recv)**  
Do something on GPU

*!Device pointers f\_send & f\_recv are passed to MPI\_send & MPI\_recv*

**!\$acc host\_data use\_device(f\_send,f\_recv)**

call MPI\_Send(f\_send, N , .....)

call MPI\_Recv(f\_recv, N, .....)

**!\$acc end host\_data**

Do something on GPU

**!\$acc exit data copyout(f\_recv)**

## GPU-aware MPI with OpenMP

**!\$omp target enter data map(to: f\_send,f\_recv)**  
Do something on GPU

*!Device pointers f\_send & f\_recv are passed to MPI\_send & MPI\_recv*

**!\$omp target data use\_device\_ptr(f\_send,f\_recv)**

call MPI\_Send(f\_send, N , .....)

call MPI\_Recv(f\_recv, N, .....)

**!\$omp end target data**

Do something on GPU

**!\$omp target exit data map(from: f\_recv)**

# Hands-on Example 3: GPU-aware MPI with OpenACC/OpenMP ?

**Purpose:** To measure the time it takes to transfer data during **MPI communication**

- **Download the repo**

```
$ git clone https://github.com/HichamAgueny/multiGPU\_MPI\_examples
```

```
$ cd multiGPU_MPI_examples
```

- **For MPI-OpenACC**

```
$ cd example_3/gpuaware_mpiacc
```

- **For MPI-OpenMP**

```
$ cd example_3/gpuaware_mpiomp
```

- **To compile and execute the code**

**Load the LUMI software stack**

```
$ module load LUMI/24.03 partition/G
```

**Compile:** \$ ./compile.sh

**Submit a job:** \$ sbatch script.slurm

**View the output file:** \$ vi gpuaware\_mpiacc-xxxxxx.out



## Measruing bandwidth: 2 MPI processes and 2 GPUs, single Node

### Traditional MPI combined with OpenACC

### GPU-aware MPI with OpenACC

```
--Time (s)      0.00002 Data size (B)      128 Bandwidth (GBps)      0.01187
--Time (s)      0.00002 Data size (B)      256 Bandwidth (GBps)      0.02398
--Time (s)      0.00003 Data size (B)      512 Bandwidth (GBps)      0.04033
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--Time (s)      0.00010 Data size (B)   131072 Bandwidth (GBps)      2.51498
--Time (s)      0.00019 Data size (B)   262144 Bandwidth (GBps)      2.78053
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--Time (s)      0.00066 Data size (B)  1048576 Bandwidth (GBps)      3.15746
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--Time (s)      0.05028 Data size (B) 134217728 Bandwidth (GBps)      5.33868
--Time (s)      0.09886 Data size (B) 268435456 Bandwidth (GBps)      5.43059
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--Time (s)      0.38534 Data size (B) 1073741824 Bandwidth (GBps)      5.57296
```

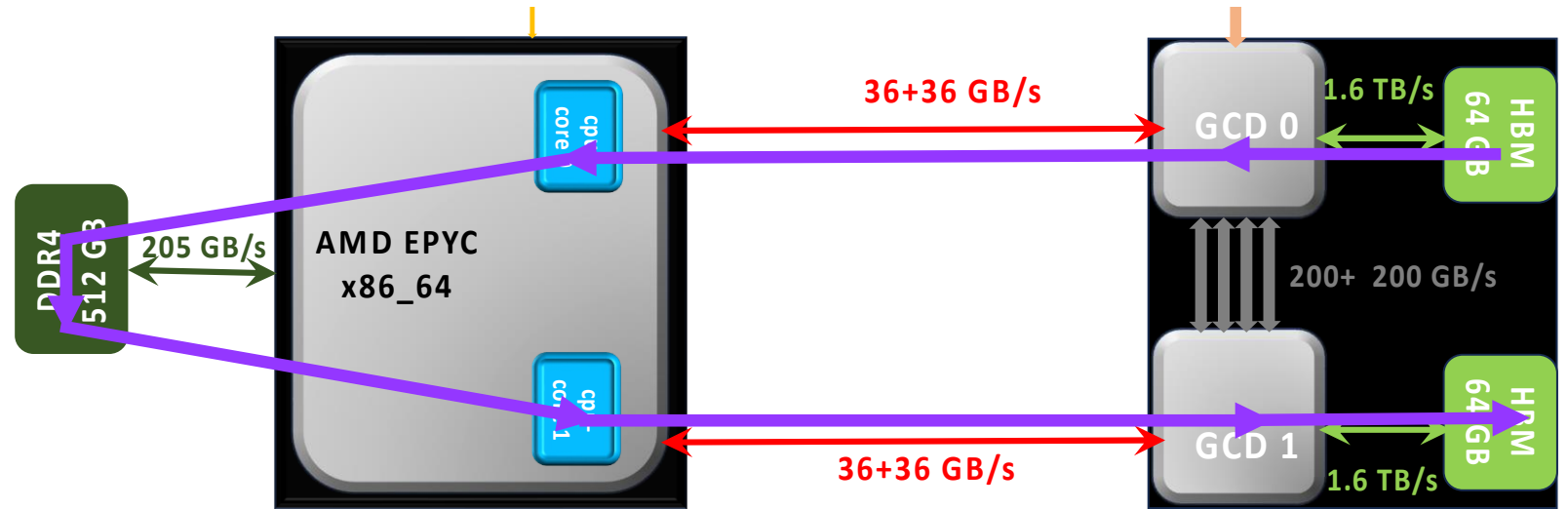
```
0.00001 Data size (B)      128 Bandwidth (GBps)      0.03195
0.00054 Data size (B)      256 Bandwidth (GBps)      0.00095
0.00001 Data size (B)      512 Bandwidth (GBps)      0.16521
0.00003 Data size (B)     1024 Bandwidth (GBps)      0.07636
0.00003 Data size (B)     2048 Bandwidth (GBps)      0.12525
0.00003 Data size (B)     4096 Bandwidth (GBps)      0.24760
0.00053 Data size (B)     8192 Bandwidth (GBps)      0.03119
0.00005 Data size (B)    16384 Bandwidth (GBps)      0.60414
0.00005 Data size (B)    32768 Bandwidth (GBps)      1.22875
0.00005 Data size (B)    65536 Bandwidth (GBps)      2.44771
0.00006 Data size (B)   131072 Bandwidth (GBps)      4.66266
0.00006 Data size (B)   262144 Bandwidth (GBps)      8.59824
0.00007 Data size (B)   524288 Bandwidth (GBps)     14.75897
0.00009 Data size (B)  1048576 Bandwidth (GBps)     23.20806
0.00013 Data size (B)  2097152 Bandwidth (GBps)     32.36915
0.00019 Data size (B)  4194304 Bandwidth (GBps)     45.31348
0.00030 Data size (B)  8388608 Bandwidth (GBps)     55.42490
0.00055 Data size (B) 16777216 Bandwidth (GBps)     61.15308
0.00098 Data size (B) 33554432 Bandwidth (GBps)     68.44087
0.00188 Data size (B) 67108864 Bandwidth (GBps)     71.25060
0.00383 Data size (B) 134217728 Bandwidth (GBps)     70.11301
0.00714 Data size (B) 268435456 Bandwidth (GBps)     75.14631
0.01384 Data size (B) 536870912 Bandwidth (GBps)     77.59176
0.02676 Data size (B) 1073741824 Bandwidth (GBps)     80.25726
```

The speed of transferring **1 GB** of data between 2 MPI-processes is about:

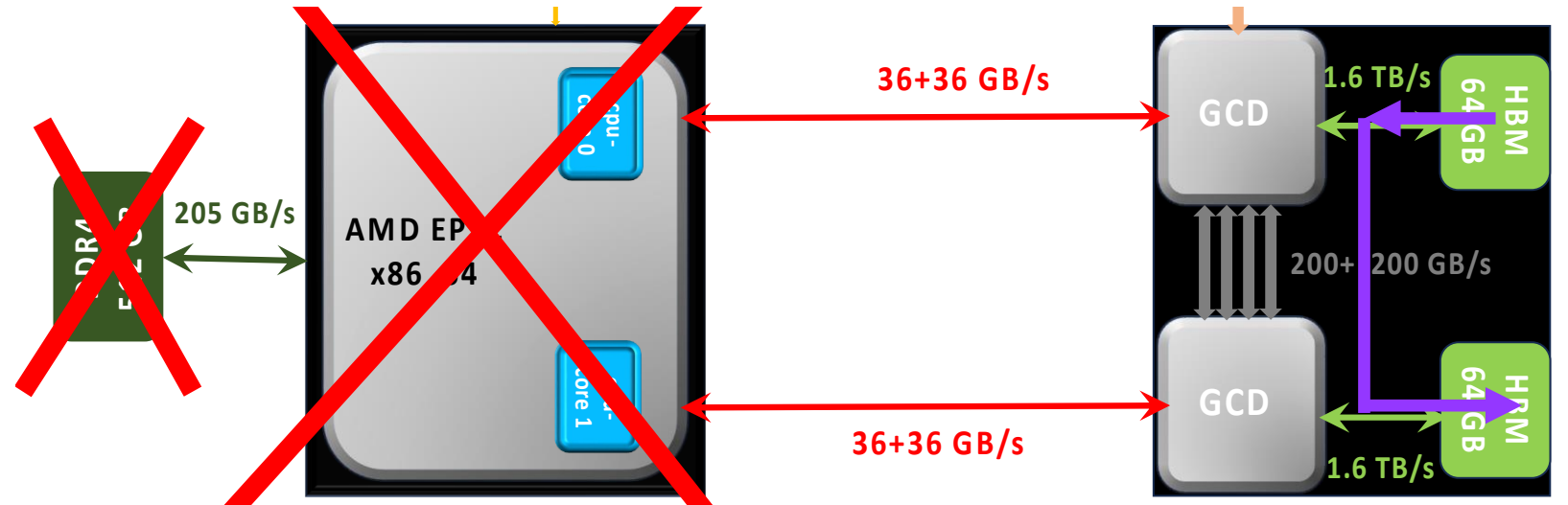
Traditional MPI **6 GB/s** vs **80 GB/s** GPU-aware MPI



## Traditional MPI



## GPU-aware MPI



# GPU-Binding (Efficient data transfer)

```
hiagueny@uan01:~> salloc -A project_465000485 -t 00:05:00 -p standard-g -N 1 --gpus 8
salloc: Pending job allocation 3636016
salloc: job 3636016 queued and waiting for resources
salloc: job 3636016 has been allocated resources
salloc: Granted job allocation 3636016
hiagueny@uan01:~> srun rocm-smi --showtoponuma
```

For NVIDIA: `srun nvidia-smi topo -m`

```
===== ROCm System Management Interface =====
===== Numa Nodes =====
GPU[0]      : (Topology) Numa Node: 3
GPU[0]      : (Topology) Numa Affinity: 3
GPU[1]      : (Topology) Numa Node: 3
GPU[1]      : (Topology) Numa Affinity: 3
GPU[2]      : (Topology) Numa Node: 1
GPU[2]      : (Topology) Numa Affinity: 1
GPU[3]      : (Topology) Numa Node: 1
GPU[3]      : (Topology) Numa Affinity: 1
GPU[4]      : (Topology) Numa Node: 0
GPU[4]      : (Topology) Numa Affinity: 0
GPU[5]      : (Topology) Numa Node: 0
GPU[5]      : (Topology) Numa Affinity: 0
GPU[6]      : (Topology) Numa Node: 2
GPU[6]      : (Topology) Numa Affinity: 2
GPU[7]      : (Topology) Numa Node: 2
GPU[7]      : (Topology) Numa Affinity: 2
===== End of ROCm SMI Log =====
hiagueny@uan01:~> srun lscpu | grep NUMA
NUMA node(s): 4
NUMA node0 CPU(s): 0-15,64-79
NUMA node1 CPU(s): 16-31,80-95
NUMA node2 CPU(s): 32-47,96-111
NUMA node3 CPU(s): 48-63,112-127
```

# Binding option: CPU-GPU affinity

```
hiagueny@uan01:~> salloc -A project_465000485 -t 00:05:00 -p standard-g -N 1 --gpus 8
salloc: Pending job allocation 3636016
salloc: job 3636016 queued and waiting for resources
salloc: job 3636016 has been allocated resources
salloc: Granted job allocation 3636016
hiagueny@uan01:~> srun rocm-smi --showtoponuma
```

=====  
===== ROCm System Management Interface =====  
===== Numa Nodes =====

GPU[0]	:	(Topology)	Numa Node: 3
GPU[0]	:	(Topology)	Numa Affinity: 3
GPU[1]	:	(Topology)	Numa Node: 3
GPU[1]	:	(Topology)	Numa Affinity: 3
GPU[2]	:	(Topology)	Numa Node: 1
GPU[2]	:	(Topology)	Numa Affinity: 1
GPU[3]	:	(Topology)	Numa Node: 1
GPU[3]	:	(Topology)	Numa Affinity: 1
GPU[4]	:	(Topology)	Numa Node: 0
GPU[4]	:	(Topology)	Numa Affinity: 0
GPU[5]	:	(Topology)	Numa Node: 0
GPU[5]	:	(Topology)	Numa Affinity: 0
GPU[6]	:	(Topology)	Numa Node: 2
GPU[6]	:	(Topology)	Numa Affinity: 2
GPU[7]	:	(Topology)	Numa Node: 2
GPU[7]	:	(Topology)	Numa Affinity: 2

NUMA node 3

NUMA node 1

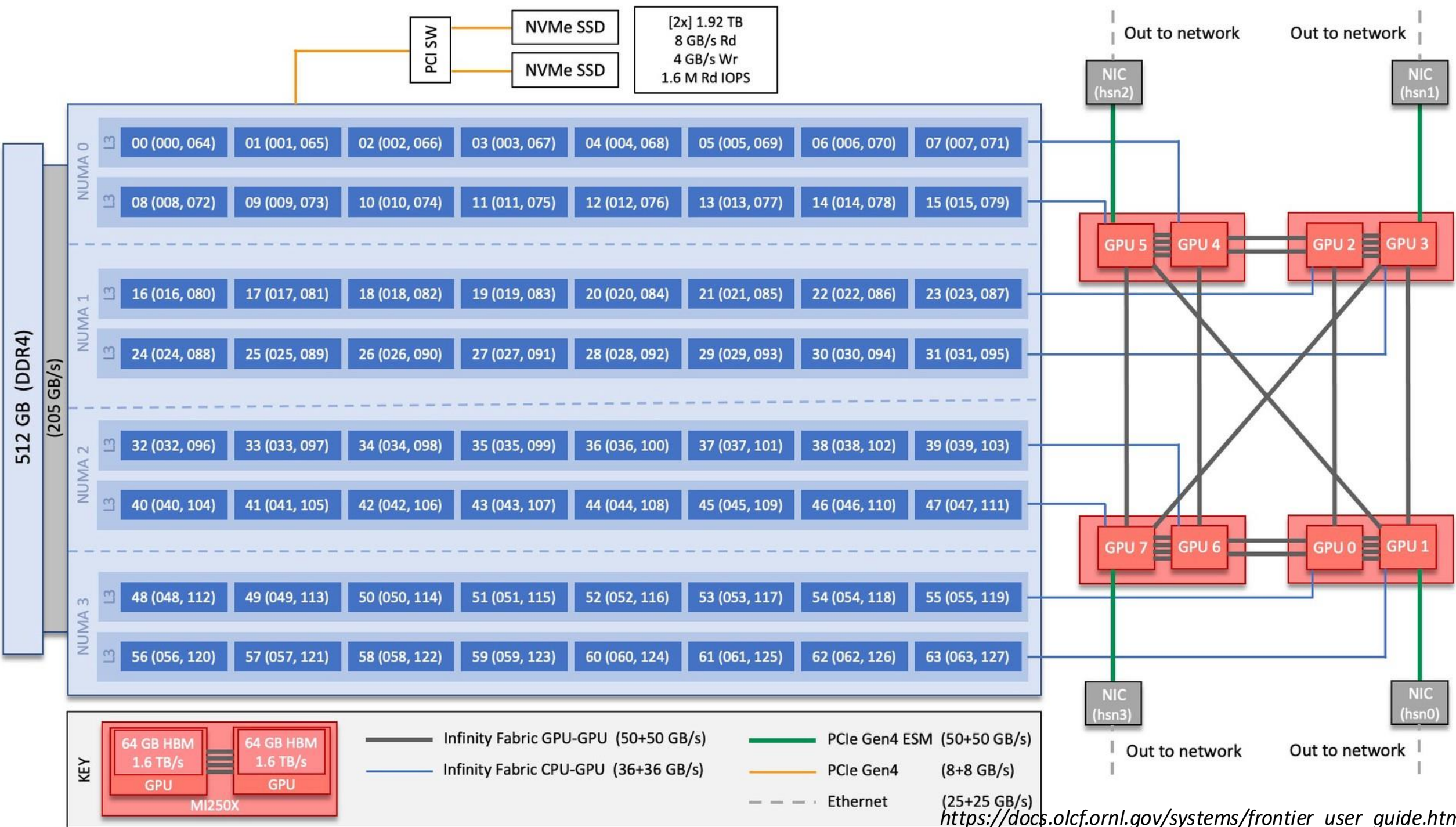
NUMA node 0

NUMA node 2

=====  
===== End of ROCm SMI Log =====

```
hiagueny@uan01:~> srun lscpu | grep NUMA
NUMA node(s): 4
NUMA node0 CPU(s): 0-15,64-79
NUMA node1 CPU(s): 16-31,80-95
NUMA node2 CPU(s): 32-47,96-111
NUMA node3 CPU(s): 48-63,112-127
```





# Binding option: CPU-GPU affinity

```
hiagueny@uan01:~> salloc -A project_465000485 -t 00:05:00 -p standard-g -N 1 --gpus 8
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===== ROCm System Management Interface =====
===== Numa Nodes =====
GPU[0]      : (Topology) Numa Node: 3
GPU[0]      : (Topology) Numa Affinity: 3
GPU[1]      : (Topology) Numa Node: 3
GPU[1]      : (Topology) Numa Affinity: 3
GPU[2]      : (Topology) Numa Node: 1
GPU[2]      : (Topology) Numa Affinity: 1
GPU[3]      : (Topology) Numa Node: 1
GPU[3]      : (Topology) Numa Affinity: 1
GPU[4]      : (Topology) Numa Node: 0
GPU[4]      : (Topology) Numa Affinity: 0
GPU[5]      : (Topology) Numa Node: 0
GPU[5]      : (Topology) Numa Affinity: 0
GPU[6]      : (Topology) Numa Node: 2
GPU[6]      : (Topology) Numa Affinity: 2
GPU[7]      : (Topology) Numa Node: 2
GPU[7]      : (Topology) Numa Affinity: 2
===== End of ROCm SMI Log =====
hiagueny@uan01:~> srun lscpu | grep NUMA
NUMA node(s): 4
NUMA node0 CPU(s): 0-15,64-79
NUMA node1 CPU(s): 16-31,80-95
NUMA node2 CPU(s): 32-47,96-111
NUMA node3 CPU(s): 48-63,112-127
```

For NVIDIA: `srun nvidia-smi topo -m`

NUMA node 3

NUMA node 1

NUMA node 0

NUMA node 2

```
#!/bin/bash
...
...
...

srun --cpu-bind=map_cpu: 49,57, 17,25, 1,9, 33,41
./application
```

# Conclusion

# Conclusion

## Traditional MPI without GPU awareness:

- Explicit data transfer between CPU and GPU (with the involvement of CPU memory).
- GPU memory is not directly accessible from/by MPI library.

 **Additional overhead caused by data movement.**

## MPI with GPU awareness:

- MPI processes can directly access GPU memory (No involvement of CPU memory).
- Elimination of CPU-GPU data transfer.

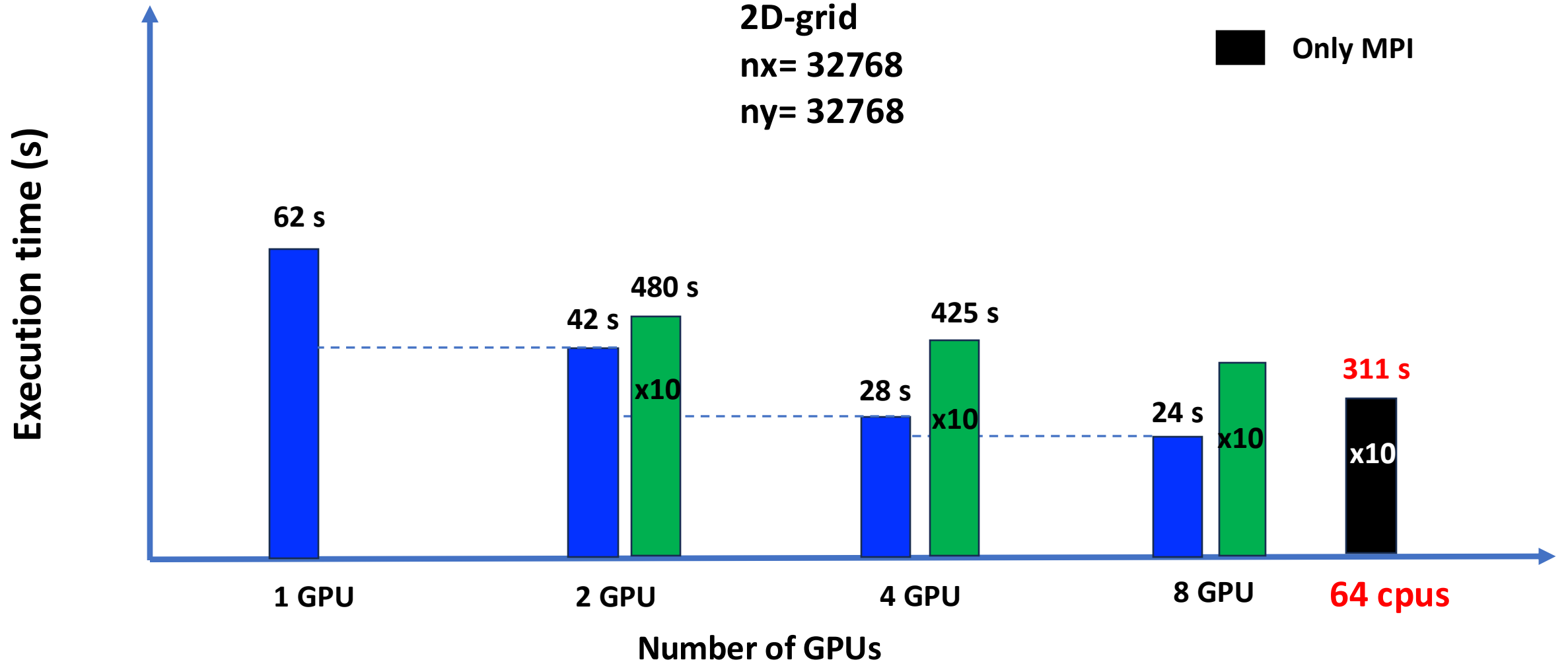
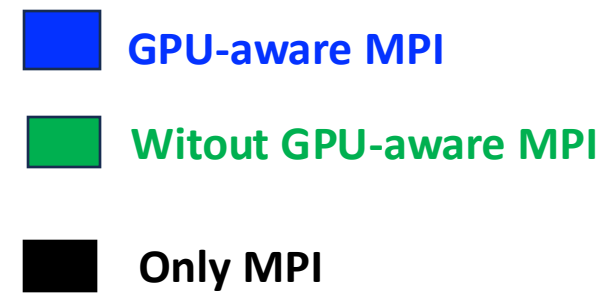
## Performance Benefits:

- Reduce data movement overhead.
- Easier to integrate directives (GPU-aware support) into existing MPI codes.

 **Significant improvement of performance**

# Benchmark (LUMI-G): 2D-Laplace Eq.

2D-grid  
nx= 32768  
ny= 32768

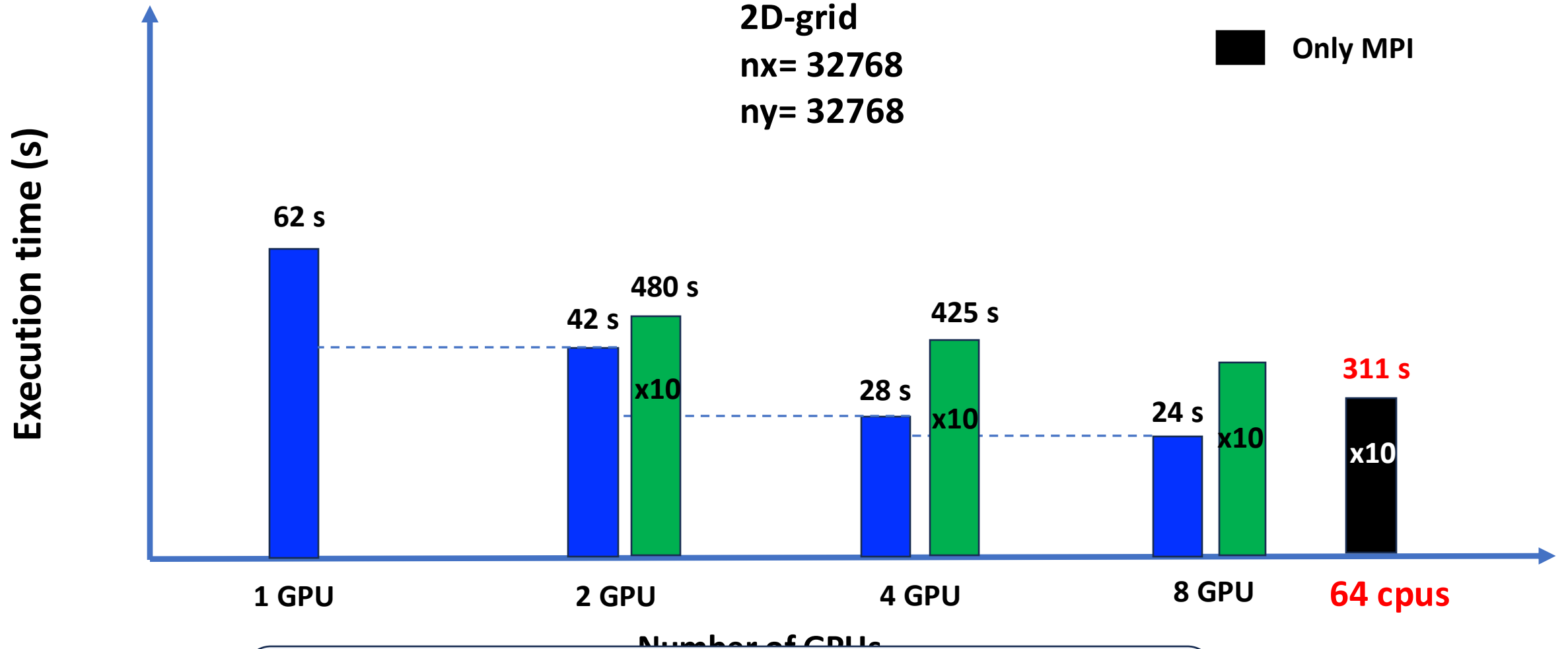




# Benchmark (LUMI-G): 2D-Laplace Eq.

2D-grid  
nx= 32768  
ny= 32768

- GPU-aware MPI
- Witout GPU-aware MPI
- Only MPI



GPU-aware MPI: speed-up by a factor  
of 10 (enhanced scalability)

# **Hands-on Exercise**

**Apply GPU-aware MPI in a 2D-Laplace equation**

# Hands-on Exercise: Apply GPU-aware MPI in a 2D-Laplace equation

- **Download the repo**

```
$ git clone https://github.com/HichamAgueny/multiGPU\_MPI\_examples
```

```
$ cd multiGPU_MPI_examples
```

- **For MPI-OpenACC**

```
$ cd exercise/mpiacc_gpuaware
```

- **For MPI-OpenMP**

```
$ cd exercise/mpiomp_gpuaware
```

- **Follow the instructions described in** `laplace_gpuaware_mpiacc.f90`  
for MPI-OpenACC or in `laplace_gpuaware_mpiomp.f90` for MPI-OpenMP

- **To compile and execute the code**

**Load the LUMI software stack**

```
$ module load LUMI/24.03 partition/G
```

**Compile:** `$ ./compile.sh`

**Submit a job:** `$ sbatch script.slurm`