

# Advanced HPC - Final assignment

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# Matrix-matrix multiplication

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# General algorithm

1.  $A$ ,  $B$  and  $C$  are allocated as blocks of  $n_{rows}$  rows across processes (called  $A_{loc}$ ,  $B_{loc}$  and  $C_{loc}$ ).
2. Each process initializes  $A_{loc}$  and  $B_{loc}$  randomly;
3. If using cuBLAS,  $A_{loc}$  is moved to device.
4. Considering each  $C_{loc}$  as divided into  $n_{procs}$  adjacent blocks of  $n_{rows} \times n_{cols}$  elements, the following points are iterated  $n_{procs}$  times:
  - 4a. each process communicates to all other processes only  $B_{loc}$ 's data needed to compute the  $i$ -th block of  $C_{loc}$ ;
  - 4b. each process gathers data coming from all other processes into a single matrix, called  $B_{col}$ ;
  - 4c. if using cuBLAS,  $B_{col}$  is moved to device;
  - 4d. each process computes the  $i$ -th block of  $C_{loc}$ .
5. If using cuBLAS,  $C_{loc}$  computed on device is moved back to host.

## Implementation details

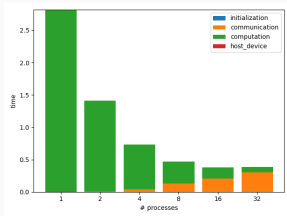
- Matrices distribution was handled by evenly distributing rows among processes. Remainder rows were assigned to first processes.
- For random initialization of  $A$  and  $B$ , a unique seed was computed randomly by process 0 and broadcasted to all other processes, which then set different seeds for each one of their thread.
- OpenMP was used to speed up the initialization of  $A$  and  $B$ , and block creation.
- MPI communications were performed using `MPI_Allgatherv()`.
- As BLAS library we chose Intel MKL 2023.2.0.
- When using BLAS and cuBLAS, we accumulated the computed entries of  $C_{loc}$  directly on the memory allocated for  $C_{loc}$ , by combining the pointer to matrix and stride arguments.
- When using cuBLAS, we avoided transpositions by computing  $B_{col}^T \times A_{loc}^T$ .

Times measured:

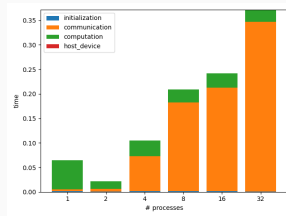
1. **Matrices initialization**, including:
  - random initialization of matrices  $A_{loc}$  and  $B_{loc}$ .
2. **Communication**, including:
  - creation of all  $B_{block}$ 's;
  - all executions of `MPI_Allgatherv()`.
3. **Computation**, including:
  - all executions of `dgemm()`.
4. **Host-device data movements** (only for cuBLAS code), including:
  - initial copy of  $A_{loc}$  from host to device;
  - copy of all  $B_{col}$ 's from host to device;
  - final copy of  $C_{loc}$  from device to host.

- All measured times were averaged over all processes.
- We placed one process per socket in non GPU-accelerated programs (i.e. two processes per node) and one per device in GPU-accelerated program (i.e. four processes per node).

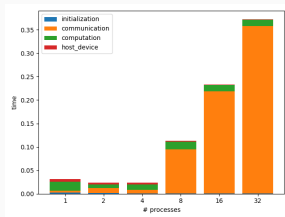
# Results - size $1000 \times 1000$



(a) "naive" - openMP  
threads: 20

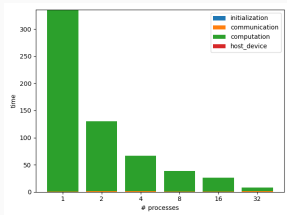


(b) BLAS; 20 openMP  
threads

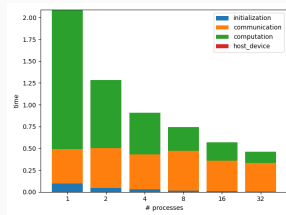


(c) cuBLAS; 8 openMP  
threads

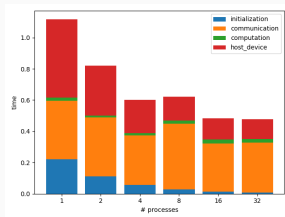
# Results - size $10000 \times 10000$



(a) "naive" - openMP  
threads: 20



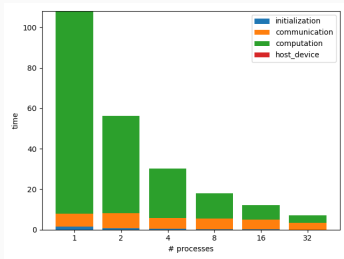
(b) BLAS; 20 openMP  
threads



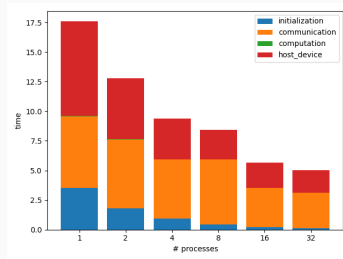
(c) cuBLAS; 8 openMP  
threads



# Results - size $40000 \times 40000$



(a) BLAS; 20 openMP threads



(b) cuBLAS; 8 openMP threads

# Observations

- Even if matrix block creation should scale, the total time for communications does not. Likely, this is because of:
  - increasing complexity in orchestrating collective communications;
  - increasing physical distance between processes.

This is particularly clear from the increasing bad communication performance as the matrix size decreases (smaller blocks but same overhead).

- Host-device data movement time does not scale perfectly, probably because of some overhead.
- When using cuBLAS the limits to performance become MPI and host-device communications.

## Jacobi method for Laplace equation

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# List of implementations

1. Simple MPI.
2. **OpenACC with CUDA-unaware MPI.**
3. **OpenACC with CUDA-aware MPI.**
4. **MPI with remote memory access.**

**All programs** were further parallelized using openMP.

# OpenACC - algorithm

1. Each process allocates its own portion of the grid. Also, an identical auxiliary matrix to store updated cells state is allocated.
2. Each process initializes its own cells.
3. Both main and auxiliary grids are moved to device.
4. For the required number of iterations, each process repeats the following:
  - 4a. communicates to the bordering processes the corresponding neighbor rows from host to host (CUDA-unaware) or from device to device (CUDA-aware);
  - 4b. (only for CUDA-unaware) received boundaries are updated on device;
  - 4c. evolves its own cells on device on the auxiliary grid, then swaps pointers on both host and device to update main grid;
  - 4d. (only for CUDA-unaware) updated border cells are copied back to host.
5. (Only for CUDA-unaware) the final grid is copied back to host.

# OpenACC - implementation details

- Grid distribution was handled by evenly distributing rows among processes.
- To speed up the grid initialization we used openMP.
- Communication of boundaries between processes was performed using `MPI_Sendrecv()`.
- Data movements were managed by placing an openACC data region around the for loop performing communications and evolution of the grid, and using clause `copy` for main and auxiliary grid.
- In the CUDA-unaware version we used the `update` clause to copy borders from host to device and vice versa.
- For CUDA-aware communications we used an openACC region with clauses `host_data` and `use_device` to "wrap" MPI calls.

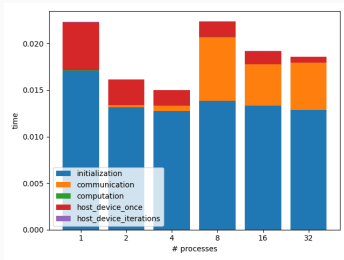
Times measured:

1. **System initialization**, including:
  - allocation and initialization of main and auxiliary matrices.
2. **One-time host-device data movements**, including:
  - initial copy of matrices from host to device;
  - final copy of main matrix from device back to host.
3. **Communication**, including:
  - all executions of `MPI_Sendrecv()`.
4. **Computation**, including:
  - all system updates on auxiliary matrix;
  - all pointer swaps for main matrix update.
5. **In-loop host-device data movements**, including:
  - all incoming borders updates after communications (host to device);
  - all outgoing borders updates after computation (device to host).

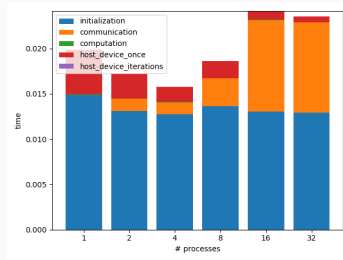
- Initialization and one-time host-device transfer times were averaged over all processes; while other times were averaged over iterations first and processes then.
- We placed one process per device (i.e. four processes per node).



# OpenACC - results - size $1200 \times 1200$

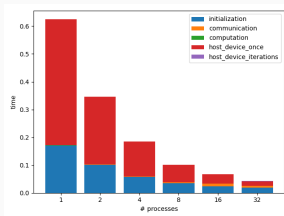


**(a)** CUDA-unaware MPI; 8 openMP threads

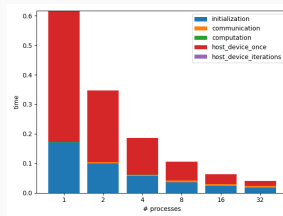


**(b)** CUDA-aware MPI; 8 openMP threads

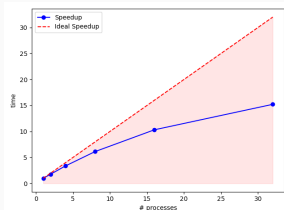
# OpenACC - results - size $12000 \times 12000$



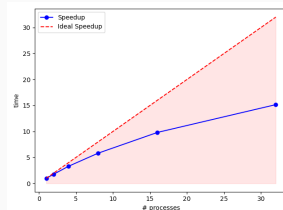
(a) CUDA-unaware MPI; 8 openMP threads



(b) CUDA-aware MPI; 8 openMP threads

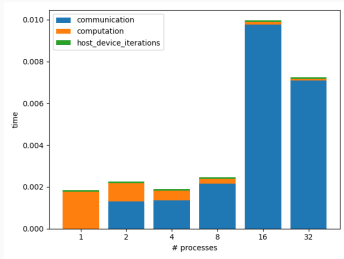


(c) CUDA-unaware MPI; 8 openMP threads

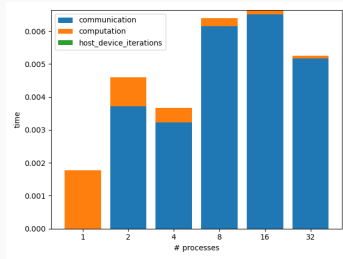


(d) CUDA-aware MPI; 8 openMP threads

# OpenACC - results - size $12000 \times 12000$ (only iterative part)

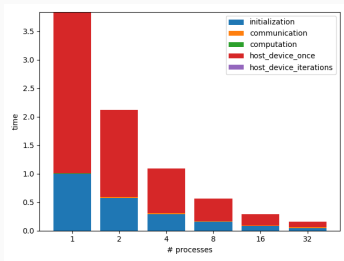


(a) CUDA-unaware MPI; 8 openMP threads

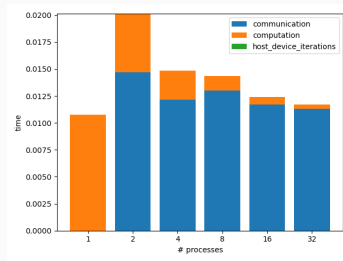


(b) CUDA-aware MPI; 8 openMP threads

# OpenACC - results - size $30000 \times 30000$



**(a)** CUDA-aware MPI; 8 openMP threads - all times



**(b)** CUDA-aware MPI; 8 openMP threads - only iterative times

- In this particular case the benefit from performing CUDA-aware MPI communications is not really significant.
- The possibility of scaling is killed by the inefficiency of MPI communications. The larger the number of iterations the stronger this effect.
- Since the difference between CUDA-aware and CUDA-unaware is not significant, MPI remote memory access could enhance scalability (see next section).

# MPI RMA - algorithm

1. Each process allocates its own portion of the grid and two additional arrays to store the upper and lower boundaries separately. Also, an auxiliary grid to store updated cells state is allocated.
2. Each process initializes its own cells.
3. For the required number of iterations, each process repeats the following:
  - 3a. opens two windows on its border arrays to allow neighbor processes to write directly;
  - 3b. writes directly to the neighbor processes' border arrays the corresponding neighbor rows (i.e. first and last), while evolves internal cells on the auxiliary grid;
  - 3c. closes the opened windows;
  - 3d. evolves cells on the first and last rows on the auxiliary grid and then swaps pointers to update main grid.

# MPI RMA - implementation details

- Grid distribution was handled by evenly distributing rows among processes. Upper and lower boundaries were separated from the actual grid.
- To speed up grid initialization and evolution we used openMP.
- For RMA operations we used `MPI_Put()` for direct writing.
- The boundary arrays were allocated contextually to window creation using `MPI_Win_allocate()`.
- We created for each process two MPI Groups containing one the previous and one the following process, and used them to expose the windows exclusively to the interested processes.
- We used `MPI_Win_post()`, `MPI_Win_start()`, `MPI_Win_complete()` and `MPI_Win_wait()` to make synchronization less "rigid"; in particular we evolved the internal cells before calling `MPI_Win_wait()`, so the evolution of these cells could be performed even if the boundaries had not been received yet.

Times measured:

1. **System initialization**, including:

- allocation of main and auxiliary matrices and window allocation for boundaries;
- main and auxiliary matrices and extreme boundaries initialization.

2. **Communication**, including:

- all MPI direct put on neighbor processes's windows execution (i.e. `MPI_Win_start()`, `MPI_Put()` and `MPI_Win_complete()`);
- all waits for MPI direct put on process's own windows to complete (i.e. `MPI_Win_wait()`).

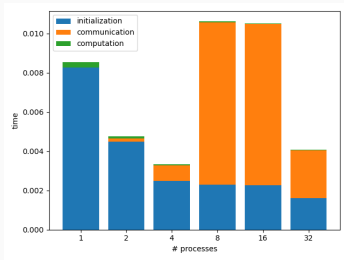
3. **Computation**, including:

- all system updates of internal rows on auxiliary matrix;
- all system updates of external rows on auxiliary matrix;
- all pointer swaps for main matrix update.

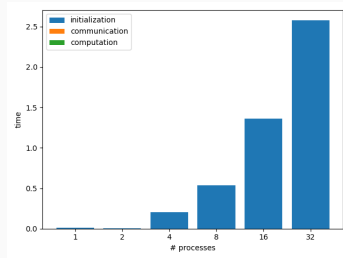


- Initialization time was averaged over all processes; while other times were averaged over iterations first and processes then.
- We placed one process per socket (i.e. two processes per node).
- **Measured communication time is not the total time for communication**: it is the additional time needed for communication w.r.t. computing time.

# MPI RMA - results - size $1200 \times 1200$

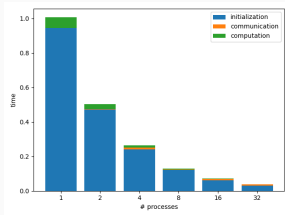


(a) standard MPI; 20 openMP threads

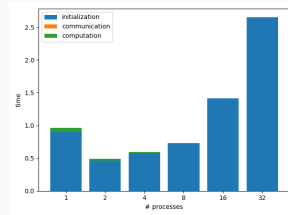


(b) MPI RMA; 20 openMP threads

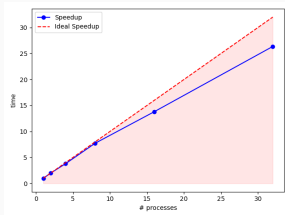
# MPI RMA - results - size $12000 \times 12000$



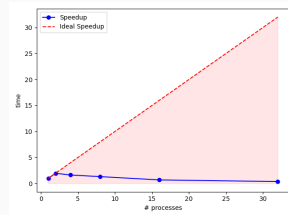
(a) standard MPI; 20 openMP threads



(b) MPI RMA; 20 openMP threads

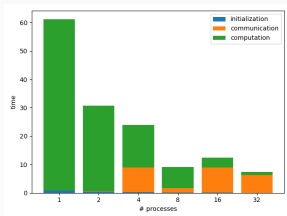


(c) standard MPI; 20 openMP threads

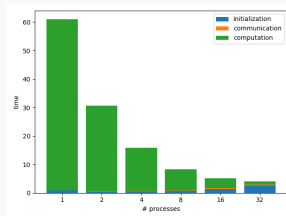


(d) MPI RMA; 20 openMP threads

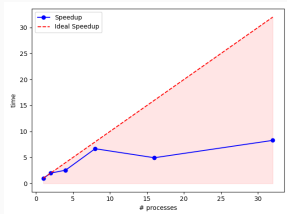
# MPI RMA - results - size $12000 \times 12000$ (1000 iterations)



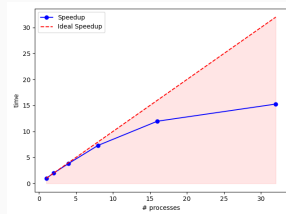
(a) standard MPI; 20 openMP threads



(b) MPI RMA; 20 openMP threads



(c) standard MPI; 20 openMP threads



(d) MPI RMA; 20 openMP threads

- Shared windows initialization is very costly, but makes communications much more efficient.
- The higher the number of iterations, the more it's beneficial to use MPI RMA (RMA "moves" the computational cost from communications to initialization).
- Cost of shared windows initialization grows with the number of processes. This effect could be mitigated by using smaller communicators, however, a overhead for communicators creation would be introduced.

Thank you!