

Introduction to MPI-Distributed Computing with GPUs





Multi-GPU Computing: What you will learn

- CUDA-aware MPI
- Example: Jacobi-Solver
- Under the hood (why you should use CUDA-aware MPI)
 - GPUs in Clusters
 - CUDA Unified Virtual Addressing
 - GPUDirect P2P and GPUDirect RDMA



Message Passing Interface - MPI

- Standard to exchange data between processes via messages
 - Defines API to exchange messages
 - Point to Point: e.g. MPI Send, MPI Recv
 - Collectives: e.g. MPI_Reduce, MPI_Allreduce, MPI_Bcast
- Multiple implementations (open source and commercial)
 - Bindings for C/C++, Fortran, Python, ...
 - e.g. MPICH, OpenMPI, MVAPICH, IBM Spectrum MPI, Cray MPT, ParaStation MPI, ...



Example MPI Program

```
#include <mpi.h>
#include <stdio.h>

int main(int argc, char** argv) {
    // Initialize the MPI environment
    MPI_Init(NULL, NULL);
    int size;
    MPI_Comm_size(MPI_COMM_WORLD, &size);
    int world_rank;
    MPI_Comm_rank(MPI_COMM_WORLD, &world_rank);
    MPI_Finalize();
}
```

```
mpicc -o hello_mpi.out hello_mpi.c
mpirun -n 4 ./hello_mpi.out
```

CUDA-aware MPI

CUDA-aware MPI allows you to use Pointers to GPU-Memory as source and destination

```
//MPI rank 0
MPI_Send (s_buf_d, n, MPI_BYTE, size-1 , tag, MPI_COMM_WORLD);

//MPI size-1
MPI_Recv (r_buf_d, n, MPI_BYTE 0, tag, MPI_COMM_WORLD, MPI_STATUS_IGNORE);
```

Pointer to GPU memory!



LAUNCH MPI+CUDA

- Launch one process per GPU
- How to use CUDA-aware MPI
 - MVAPICH: \$MV2_USE_CUDA=1 mpirun -np \${np} ./myapp <args>
 - Open MPI: CUDA-aware features are enabled per default (using UCX)
 - Cray: MPICH RDMA ENABLED CUDA
 - IBM Spectrum MPI: \$mpirun -gpu -np \${np} ./myapp <args>
 - ParaStation MPI (using UCX): \$PSP_CUDA=1 mpirun -np \${np} ./myapp <args>
- On JUWFI S Booster:
 - Load CUDA-aware OpenMPI or ParaStation MPI modules
 - GPU-tuning is done via module loading
 - srun --gres=gpu:4 -n {np} ./myapp <args>



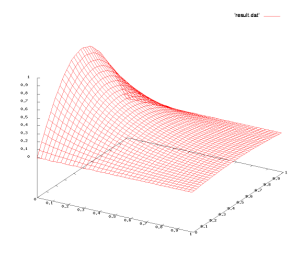
How to compile

```
nvcc -o my_kernel.o $(NVCC_FLAGS) my_kernels.cu -c
mpicc -o my_multiGPUapp -lcudart my_kernel.o my_multiGPUapp.c
```



Example: Jacobi Solver

- Solves the 2D-Laplace Equation on a rectangle
- $\Delta u(x,y) = \mathbf{0} \ \forall \ (x,y) \in \Omega \backslash \delta\Omega$
- Dirichlet boundary conditions on left/right boundaries (constant values on boundaries)
- Reflecting boundaries on top and bottom
- Iterative solver: u(t+1) = f(u(t))

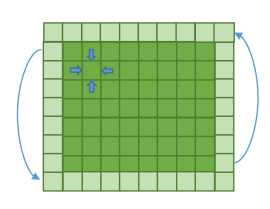




Example: Jacobi solver

While not converged

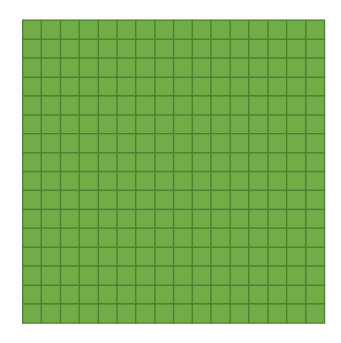
do Jacobi step:



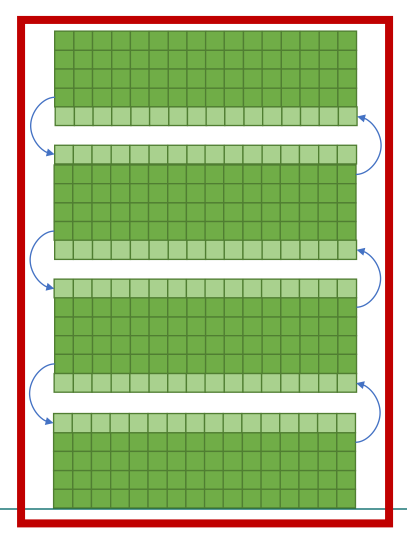
apply boundary Condition swap a_new and a next iteration

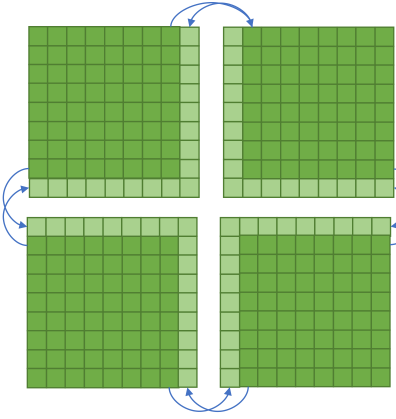


Domain Decomposition



Minimize number of neighbors: Communicate to less neighbors Optimal for latency bound communication, Continuous Transfers



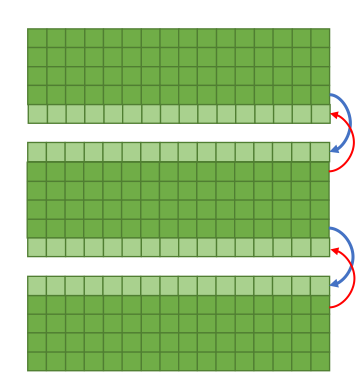


Minimize surface area/volume ratio: Communicate less data Optimal for bandwidth bound communication Non-Continuous Transfers



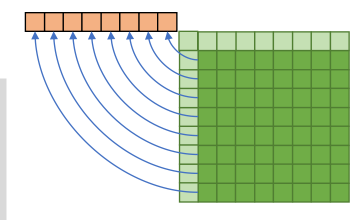
Jacobi example: Top and Bottom Boundaries

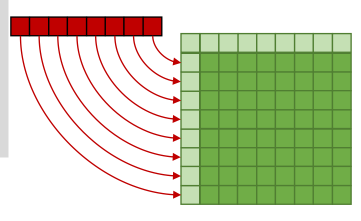
bottom





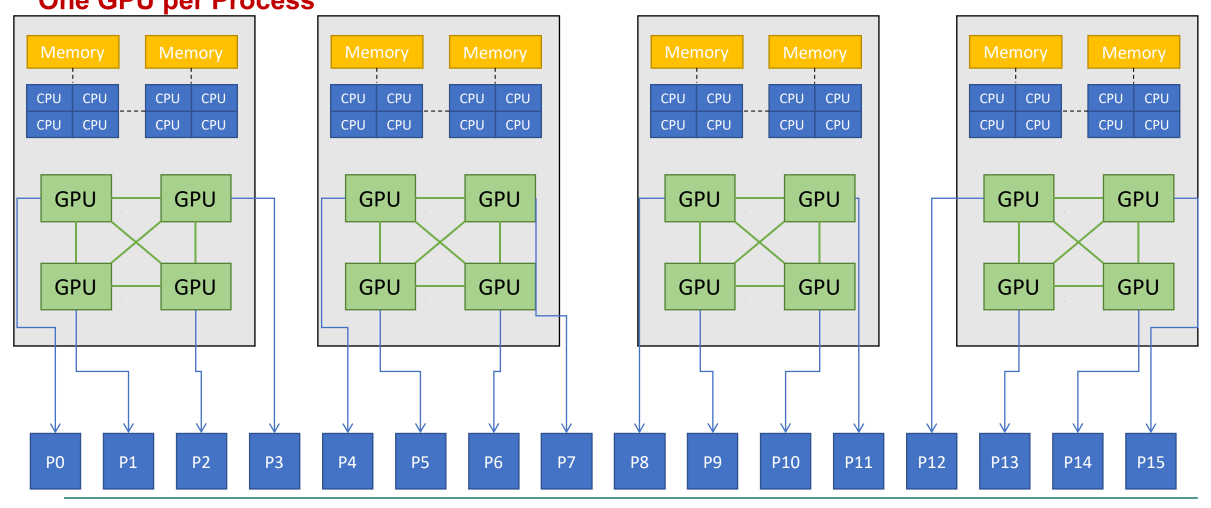
Jacobi: Left (and right) neighbours







Process Mapping on Multi GPU Systems One GPU per Process

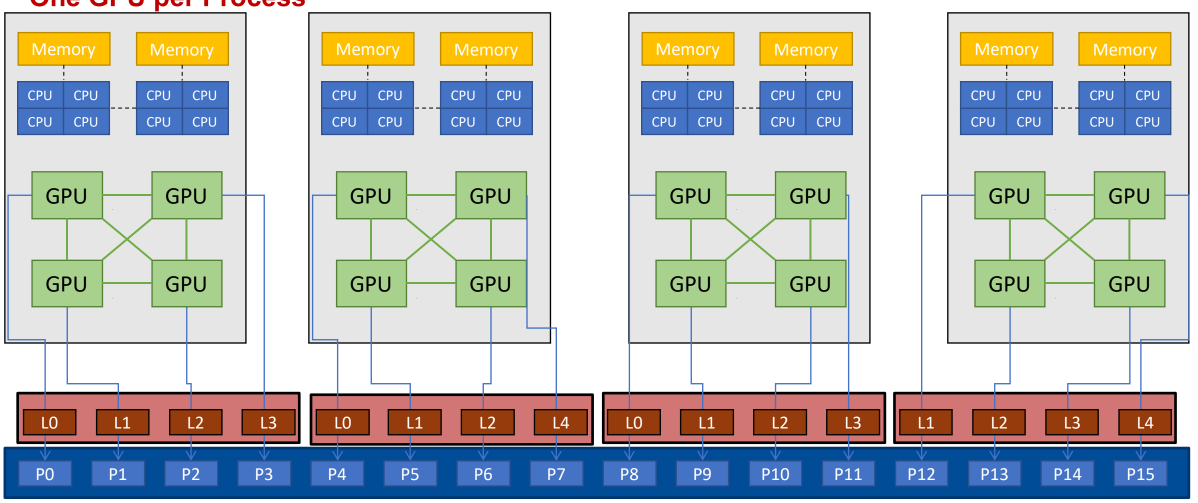




Distribute GPUs to local Nodes



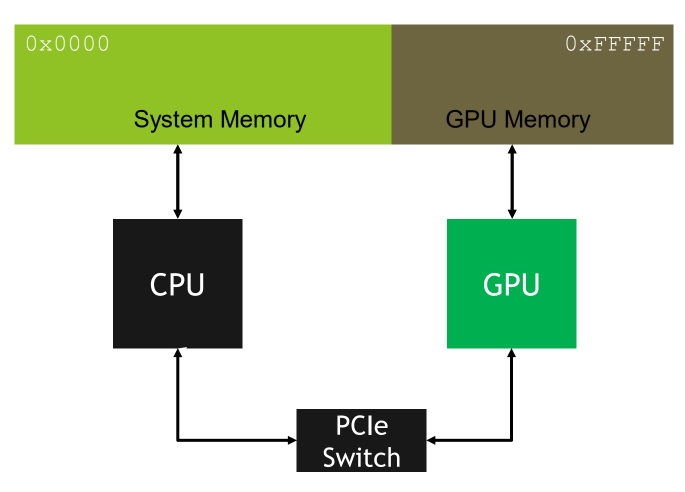
Process Mapping on Multi GPU Systems One GPU per Process





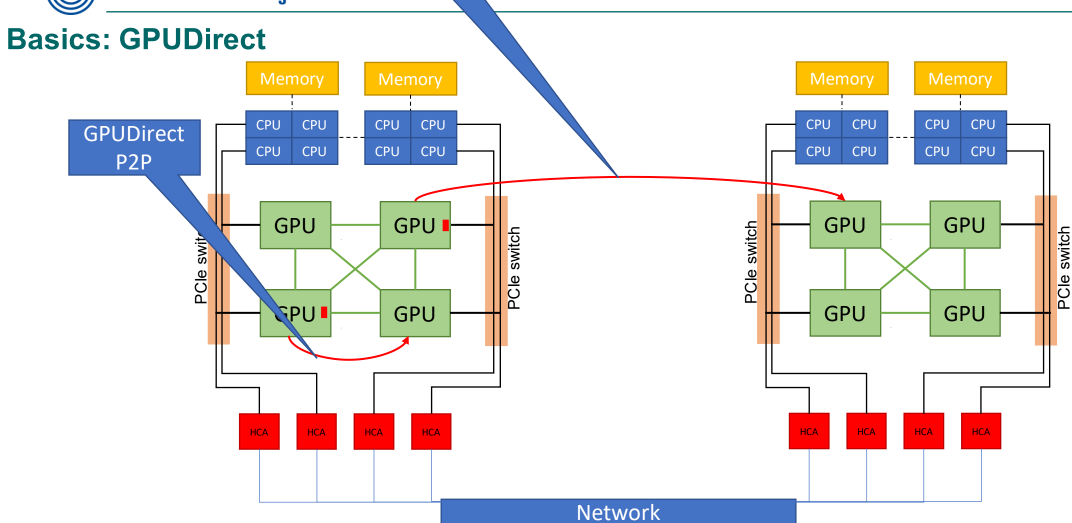
CUDA Unified Virtual Addressing

- One address space for all CPU and GPU memory
 - Determine physical memory location from a pointer value
 - Enable libraries to simplify their interfaces (e.g. MPI)
- Supported on devices with compute capability 2.0+ for
 - 64-bit applications on Linux and Windows (+TCC)

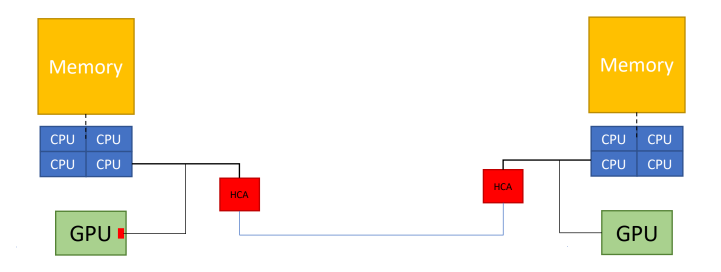




GPUDirect RDMA



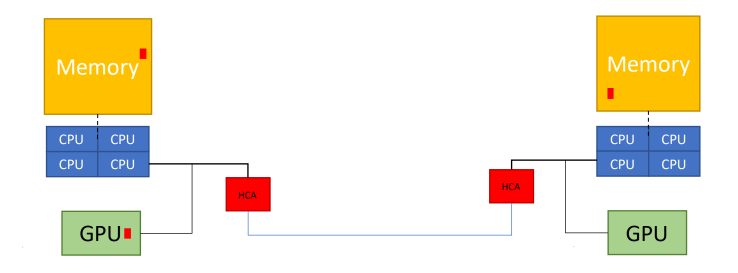
CUDA-aware MPI with GPUDirect RDMA



```
MPI_Send(s_buf_d,size,MPI_BYTE,1,tag,MPI_COMM_WORLD);
MPI_Recv(r_buf_d,size,MPI_BYTE,0,tag,MPI_COMM_WORLD,&stat);
```

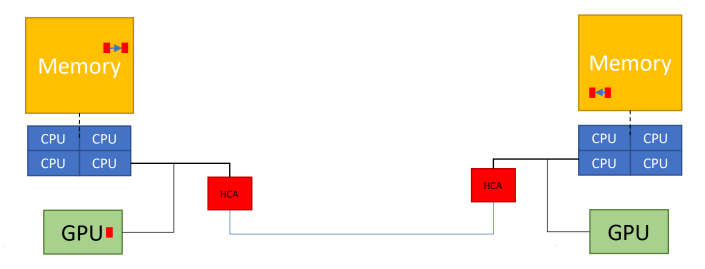


CUDA-aware MPI without GPUDirect RDMA



```
MPI_Send(s_buf_d,size,MPI_BYTE,1,tag,MPI_COMM_WORLD);
MPI_Recv(r_buf_d,size,MPI_BYTE,0,tag,MPI_COMM_WORLD,&stat);
```

Regular MPI



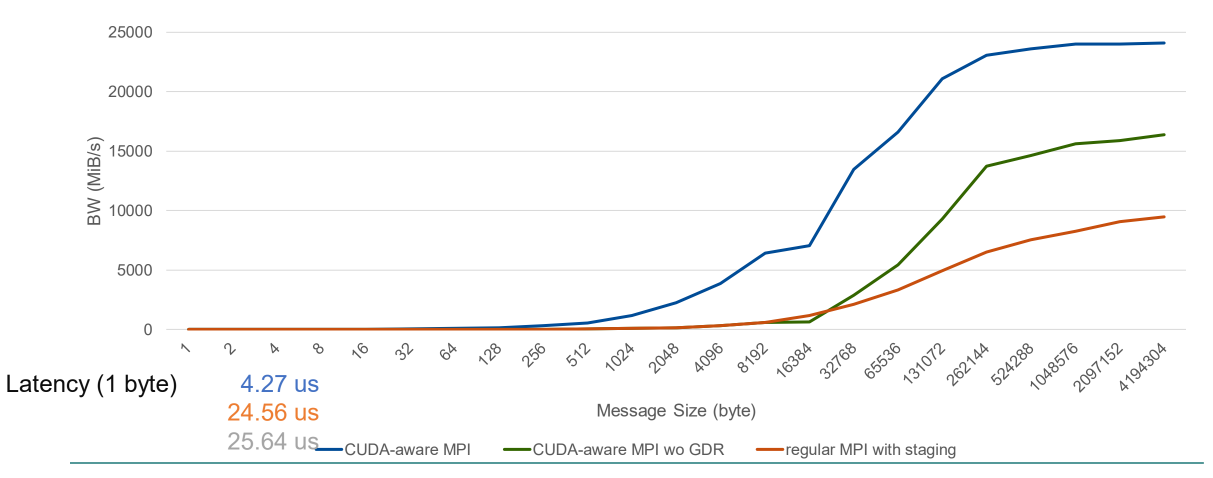
```
cudaMemcpy(s_buf_h,s_buf_d,size,cudaMemcpyDeviceToHost);
MPI_Send(s_buf_h,size,MPI_BYTE,1,tag,MPI_COMM_WORLD);

MPI_Recv(r_buf_h,size,MPI_BYTE,0,tag,MPI_COMM_WORLD,&stat);
cudaMemcpy(r_buf_d,r_buf_h,size,cudaMemcpyHostToDevice);
```



OpenMPI 4.1.0RC1 + UCX 1.9.0 on JUWELS Booster

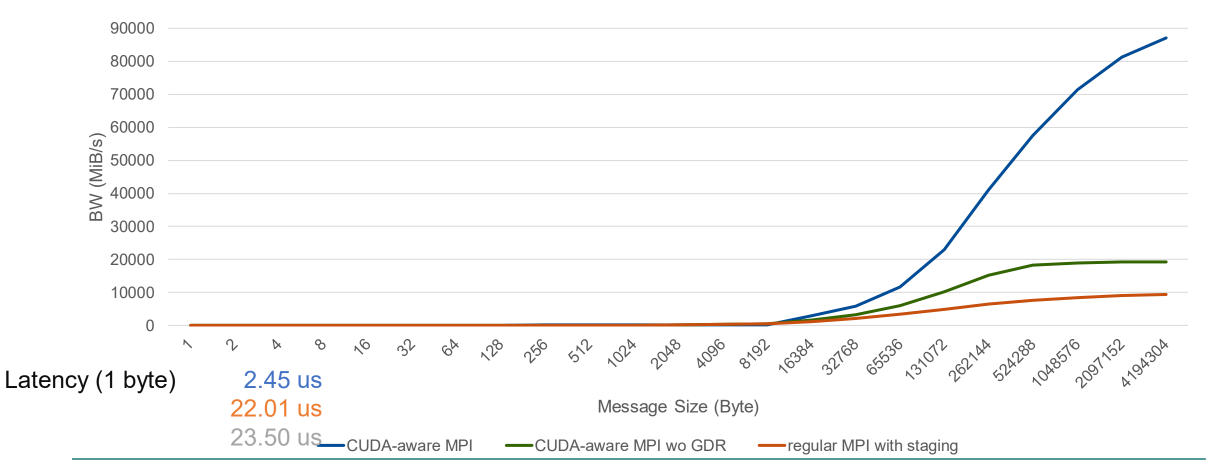
Performance Results GPUDirect RDMA





OpenMPI 4.1.0RC1 + UCX 1.9.0 on JUWELS-Booster

Performance Results GPUDirect P2P





Summary

- CUDA-aware MPI allows efficient communication for multi-GPU applications
 - Allows MPI-communication operations from GPU memory buffers
 - Simplified programming
 - Use GPUDirect technologies for performance
 - Minimizes data copies
- Most MPI versions have support for CUDA-aware MPI
- Best practice: One process per rank
 - Use local communicator in MPI