







Simon Garcia de Gonzalo, Sandia National Laboratories (slides from Dr. Lena Oden, FernUniversität in Hagen)





Sandia National Laboratories is a multimission laboratory managed and operated by National Technology & Engineering Solutions of Sandia, LLC, a wholly owned subsidiary of Honeywell International Inc., for the U.S. Department of Energy's National Nuclear Security Administration under contract DE-NA0003525.

# 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_Send(s_buf_d,n,MPI_BYTE,size-1 ,tag,MPI_COMM_WORLD);

//WPI_size-1
MPI_Recv(r_buf_d,n,MPI_BYN, 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 JUWELS 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
```

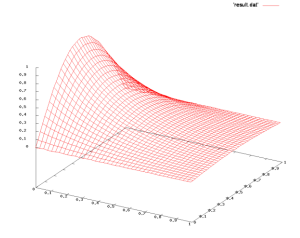
# Example: Jacobi Solver



- Solves the 2D-Laplace Equation on a rectangle
- $\Delta u(x,y) = \mathbf{0} \ \forall \ (x,y) \in \Omega \setminus \delta \Omega$



- Reflecting boundaries on top and bottom
- Iterative solver: u(t+1) = f(u(t))

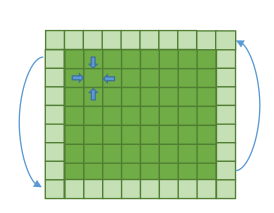


## **1**

# Example: Jacobi solver

While not converged

do Jacobi step:



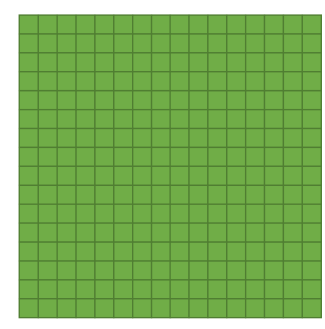
apply boundary Condition

swap a\_new and a

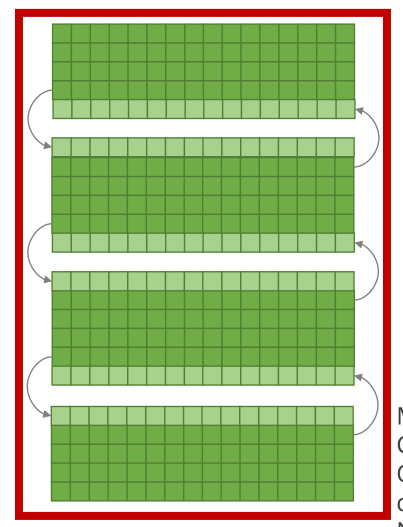
next iteration

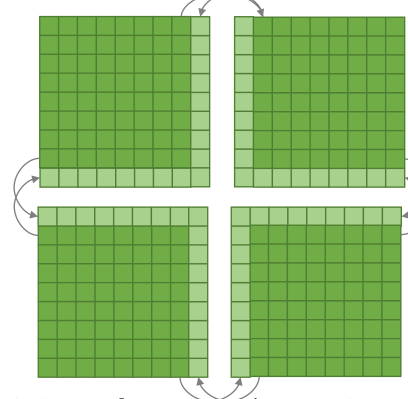
# **Domain Decomposition**





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



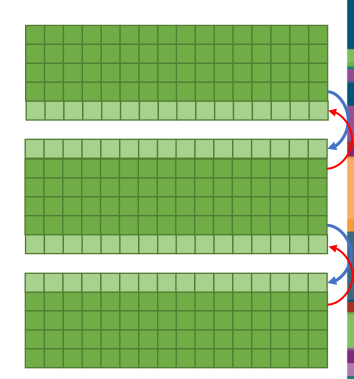
## Jacobi example: Top and Bottom Boundaries

MPI STATUS IGNORE)

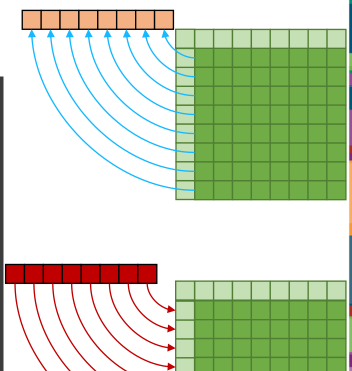
bottom

MPI DOUBLE, t nb 1, MPI COMM WORLD,

top neighbor

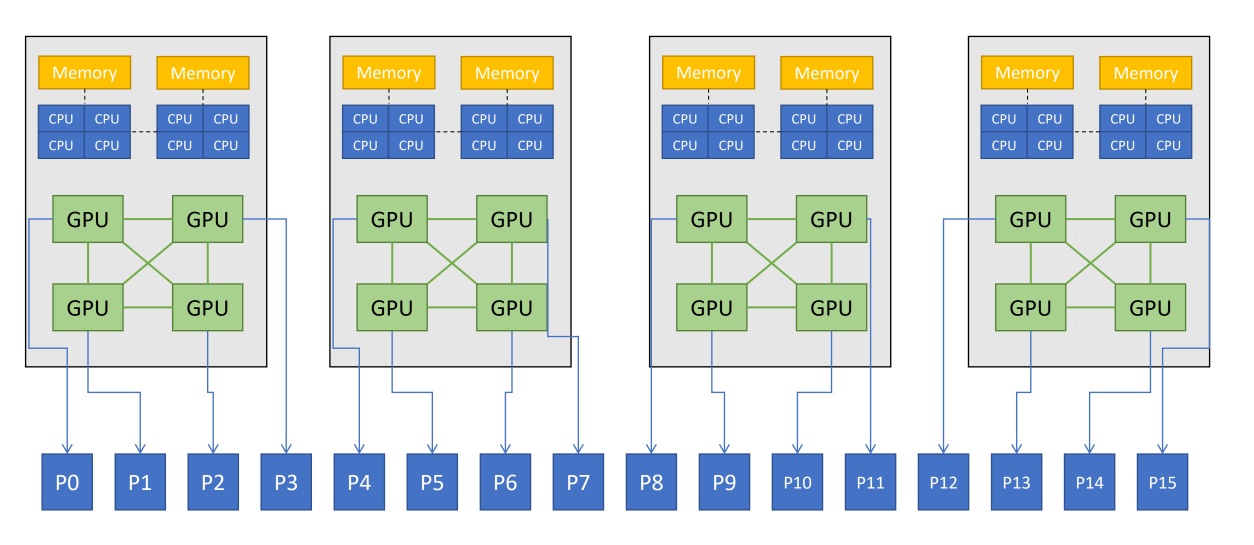


# Jacobi: Left (and right) neighbours





## Process Mapping on Multi GPU Systems One GPU per Process



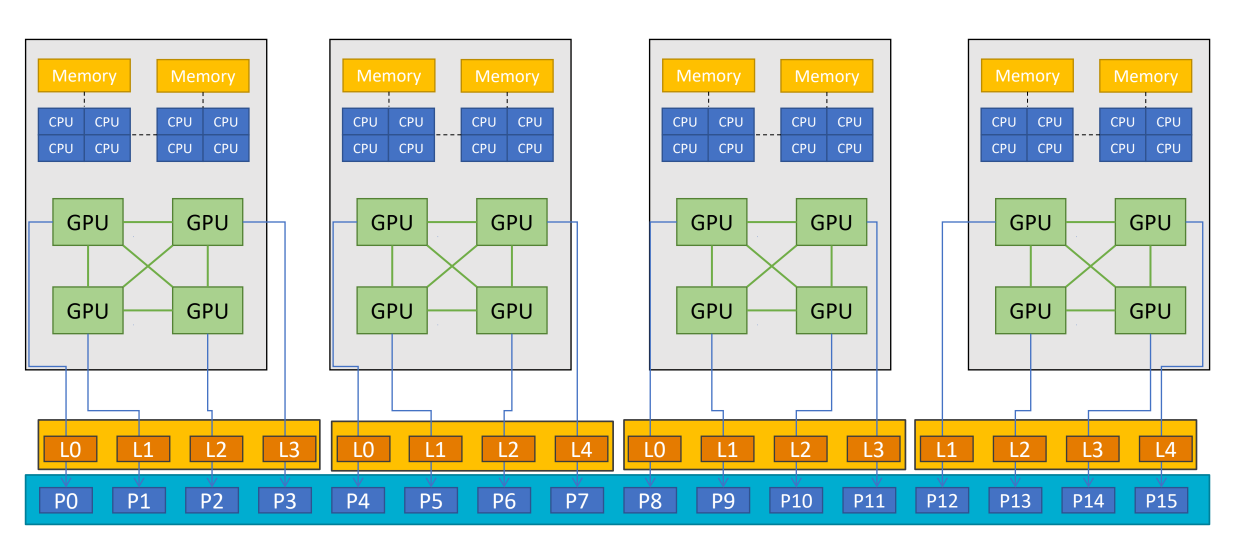
### **1**

### Distribute GPUs to local Nodes

```
MPI Comm local comm;
MPI Comm split type (MPI COMM WORLD, MPI COMM TYPE SHARED,
                                                           rank
                    MPI INFO NULL, &local_comm)
    local rank = -1;
MPI Comm rank(local comm, &local rank)
MPI Comm free(&local comm)
int num_devs = 0;
cudaGetDeviceCount(&num devs);
cudaSetDevice(local rank%num devs);
```

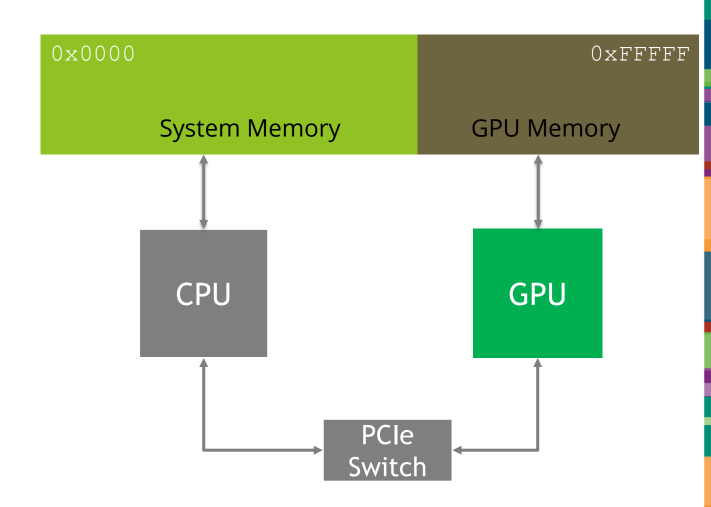


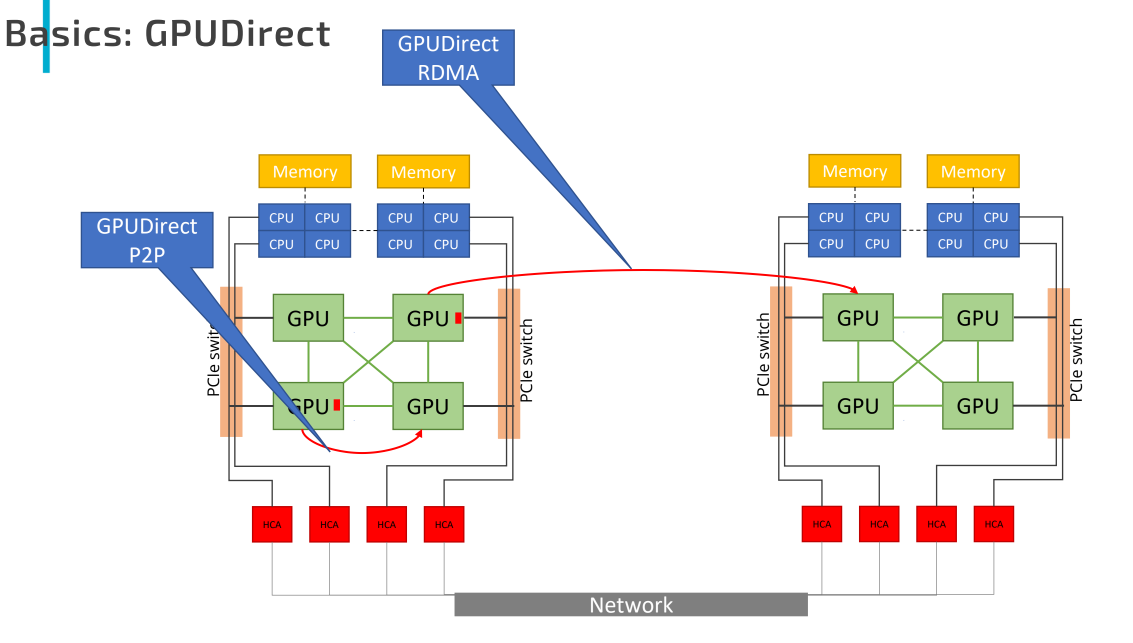
# Process Mapping on Multi GPU Systems One GPU per Process



# **CUDA Unified Virtual Addressing**

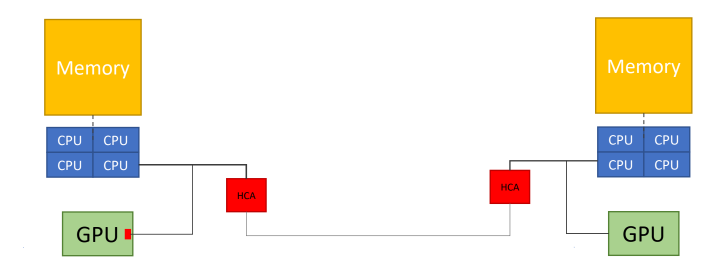
- One address space for all CPU and GPU memory
  - Determine physical memory location from a pointer value
  - Enable libraries to simplify their interfaces (e.g. MPI)
- Supported on devices with compute capability 2.0+ for
  - 64-bit applications on Linux and Windows (+TCC)







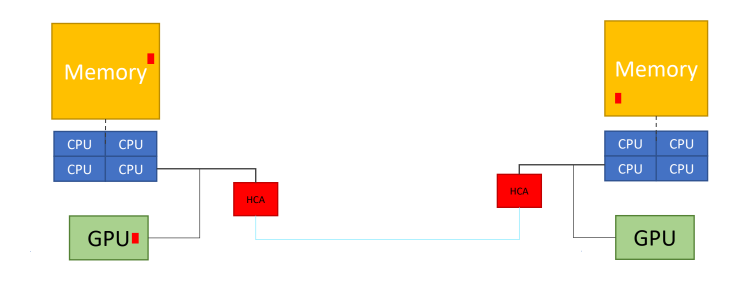
#### **CUDA-aware MPI with GPUDirect RDMA**



```
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);
```



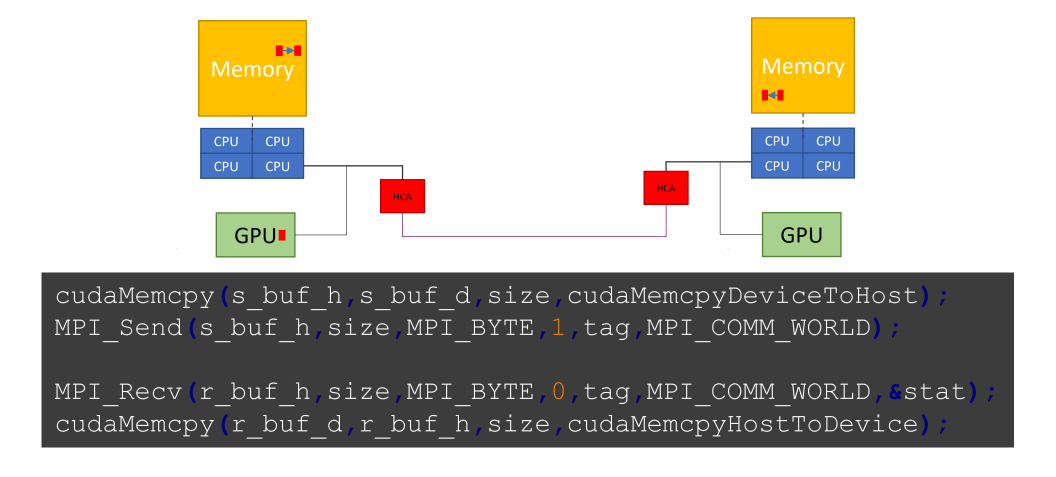
#### **CUDA-aware MPI without GPUDirect RDMA**



```
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);
```

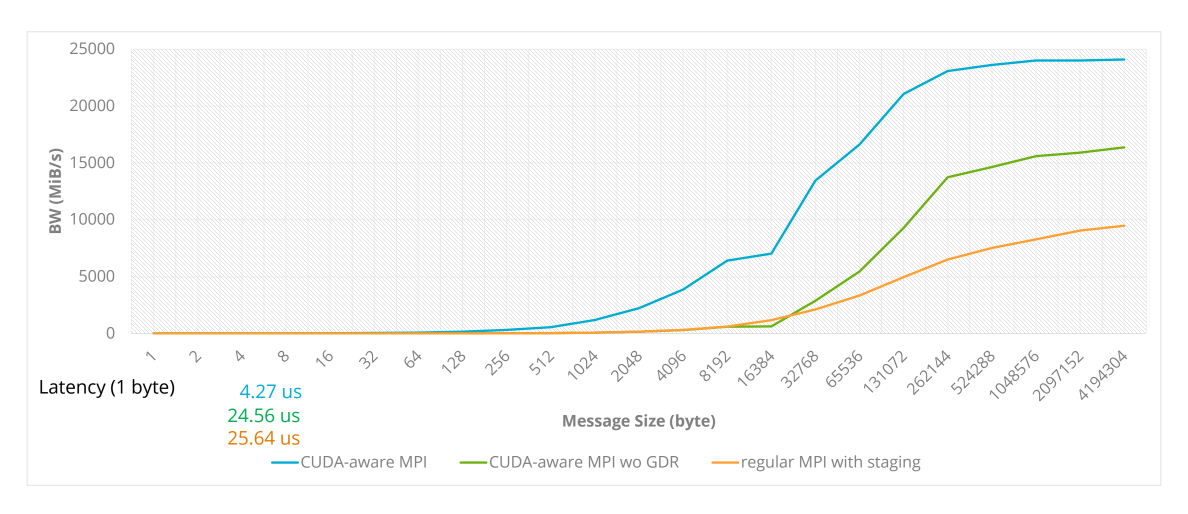


#### Regular MPI



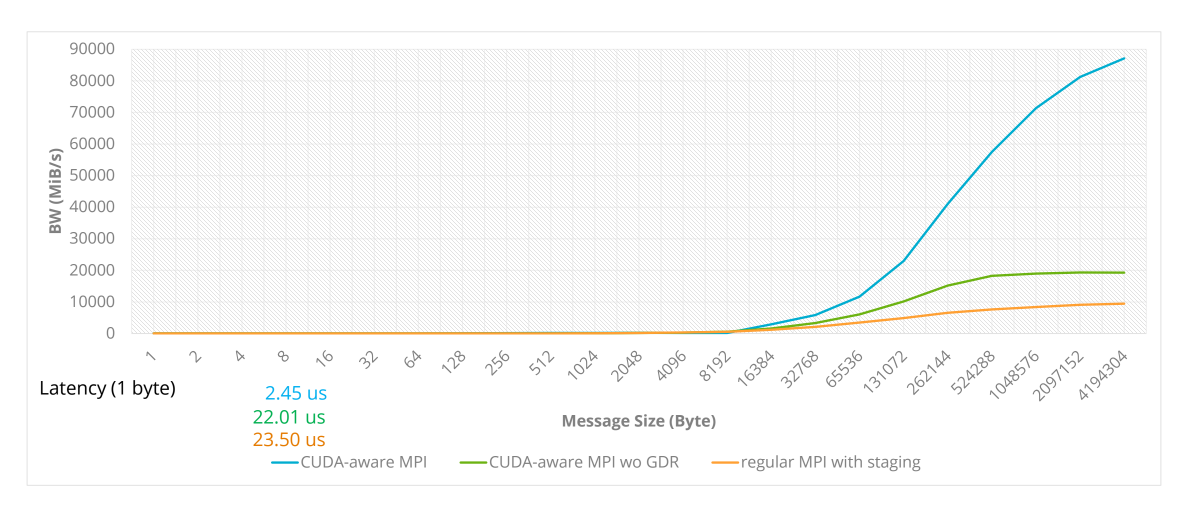
## Performance Results GPUDirect RDMA

OpenMPI 4.1.0RC1 + UCX 1.9.0 on JUWELS Booster



## Performance Results GPUDirect P2P

OpenMPI 4.1.0RC1 + UCX 1.9.0 on JUWELS-Booster



# **Summary**

**1** 

- 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