

LUMI

A white wolf is the central focus, standing in a futuristic, blue-toned digital environment. The background is filled with vertical light beams, floating particles, and a grid-like pattern, creating a high-tech, cybernetic atmosphere. The wolf is looking slightly to the right of the camera.

Process and Thread Distribution and Binding

Kurt Lust
LUMI User Support Team (LUST)
University of Antwerp

May 2024

What are we talking about?

- Distribute processes and threads across the available resources for the job
- and bind them to the resources to ensure they stay there and only use the assigned resources
 - Across nodes: Only distribution
 - Within a node: Binding necessary
- System software level (Linux/ROCm/Slurm):
 - Control groups used at the job and job step level, sometimes at the task level
 - Affinity mask to control where a thread can get scheduled
 - ROCm runtime also has a mechanism to control access to GPUs
- Tools for verification in the lumi-CPEtools modules

When/where is it done?

- Slurm level
 - Creation of allocation: Slurm reserves resources at the node level using control groups
 - Creation of job step:
 - Distributes tasks across nodes and cores/hardware threads on nodes
 - Default in most cases: Binds tasks to CPUs (affinity mask) and GPUs (control groups unfortunately)
- Application runtime library level
 - Cray MPICH can renumber the ranks
 - OpenMP runtime: select number of CPU threads and thread binning within the resources of a task using affinity masks
 - ROCm runtime: Select GPUs using [ROCR_VISIBLE_DEVICES](#)
- Does not always make sense on nodes that are not job-exclusive!

Why do I need this?

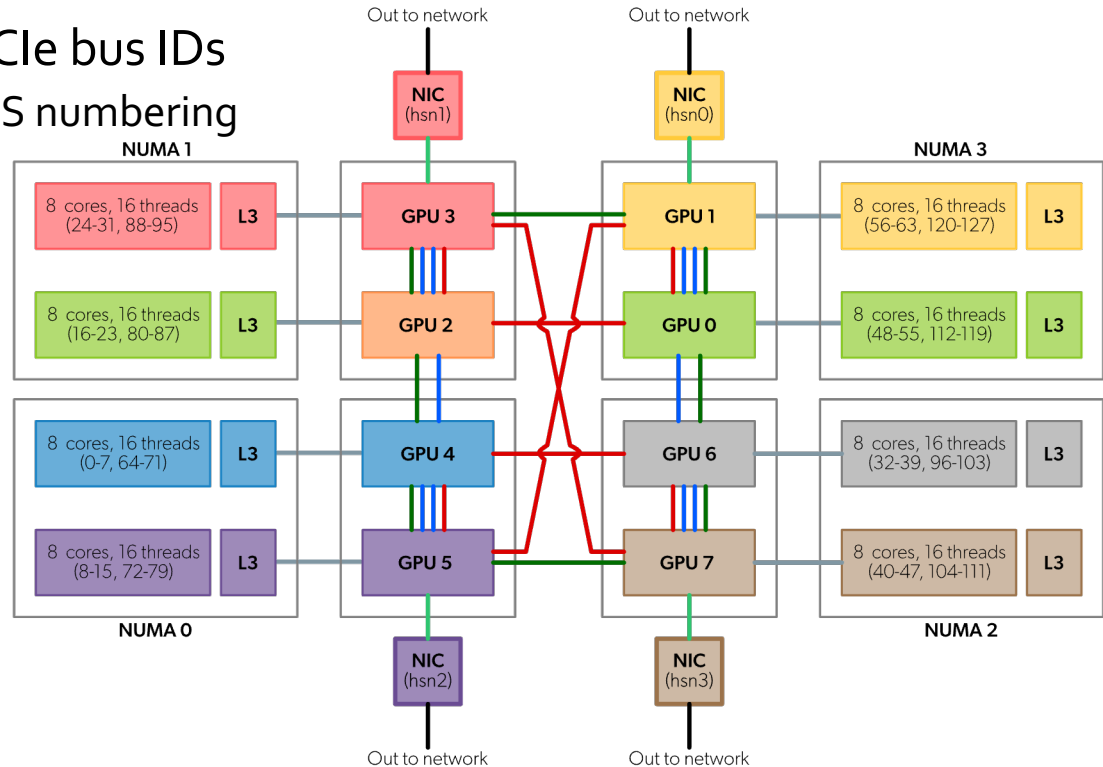
- Importance of memory locality at all levels (cache and main memory)
 - E.g.: MPI application with 14 GB/rank so 16 ranks on node: Spread out across CCDs...
 - Shared memory with lack of memory locality: Maybe need to bundle threads if the application fits in a socket
 - No solution that always works!
- Short connection between CPU and GPU sometimes essential for fast communication between both
 - Cache-coherent accesses to GPU memory by the CPU
- Mapping of MPI ranks to reduce inter-node traffic and maximise intra-node traffic which is much faster
 - Also on the GPU: Map communication pattern on the topology of a node

Core numbering

- Linux core (actually hardware thread/virtual core) numbering does not reflect the hierarchy
 - Numbers 0-127 on LUMI-C are the first hardware thread on each physical core, 128-255 then the second one, so i and $i+128$ map onto the same physical core
 - On LUMI-G: Core 0-63 first hardware thread, core 64-127 second, so i and $i+64$ map onto the same physical core
- Hardware threading on LUMI is turned on when booting a node
 - Slurm does not really turn hardware threading off when you request to, but doesn't include the second hardware thread in the affinity mask in that case
 - And slurm only does so at the regular job step level
 - The Slurm batch step will always see both hardware threads for each core!
- Technical discussion in the notes if you're interested

GPU Numbering (1)

- Very tricky
- Numbering based on the PCIe bus IDs
 - Global numbering or bare-OS numbering



GPU Numbering (2)

- Very tricky
- Numbering based on the PCIe bus IDs
 - *Global numbering or bare-OS numbering (0-7)*
- Job-level control group
 - New numbering starting from 0: *job-local numbering*
 - Same order though
- Task-level control group
 - Yet another numbering starting from 0: *task-local numbering*
 - And a headache for MPI applications
- Further restricting access via `ROCR_VISIBLE_DEVICES` will start yet another numbering in, e.g., the HIP runtime

GPU Numbering - Remarks

- Very technical demonstrations in the notes
- Slurm works differently with CPUs and GPUs on LUMI
 - CPUs: Control groups at the job level, after that affinity masks
 - GPUs: Control groups at the job and task level, even though `ROCR_VISIBLE_DEVICES` plays a bit the role of an affinity mask
- Affinity masks work differently from `ROCR_VISIBLE_DEVICES`
 - Affinity masks always refer to the global / bare OS numbering of the hardware threads
 - `ROCR_VISIBLE_DEVICES` numbering is based on the local numbering in the context where the variable is used
 - Affinity masks can only shrink as you go deeper in a hierarchy
 - `ROCR_VISIBLE_DEVICES`, being just an environment variable, can be abused to gain access to extra resources (within the confines of the control group)

Task distribution with Slurm (1)

- `srun --distribution={block|cyclic|plane=<s>}[:{block|cyclic|fcyclic}[, {Pack|NoPack}]`
- Level 1: Distribution of tasks across nodes
 - `block`: Fill first node in allocation, then fill second, etc.
 - `Pack`: Fill completely before moving to the next node
 - `NoPack`: More ballanced, trying to fill all nodes as equally as possible
 - `cyclic`: First assign one task to each node, then from the first node again assign a second task, ...
 - `plane=<s>`: As cyclic, but assigning `s` tasks at a time before moving on
 - More options that we do not discuss

Task distribution with Slurm (2)

- `srun --distribution={block|cyclic|plane=<s>}[:{block|cyclic|fcyclic}[, {Pack|NoPack}]`
- Level 2: Distribution of tasks across cores
 - **block**: Consecutive sets of cores for each task
 - **cyclic**: First assign one task to each socket on the first set of consecutive cores/virtual cores of each socket, then assign a second task on each socket on the next set of cores, ...
 - **fcyclic**: Will spread tasks out across sockets
 - Not clear where this is useful on an AMD system except for cases with one task per node and a lot of memory for that task
- Level 3 not shown in this simplified version
- Default: **block:block:nopack** but **block:*** results in **block:cyclic**
- L2 and L3 distribution conflicts with the CPU binding mechanism that we will discuss
 - But usefull with `--cpus-per-task`

Task-to-CPU binding with Slurm

- Works with affinity masks
- `srun --cpu-bind=[{quiet|verbose},]<type>`
- Some `<type>` options are for automatic binding
 - `--cpu-bind=threads` is the default behaviour on LUMI
 - Other options: See the manual
- Other `<type>` options define a list of slots to be used
 - Combination with `--distribution` L2/L3 options does not make sense
 - `--cpu-bind=map_cpu:<cpu_id_for_task_0>,<cpu_id_for_task_1>,...` : Specify a single hardware thread for each task on the node
 - For MPI programs
 - `--cpu-bind=mask_cpu:<mask_for_task_0>,<mask_for_task_1>,...` : Specify affinity mask for each task on the node.
 - For OpenMP or hybrid programs

Task-to-CPU binding with Slurm: Masks

- Slurm uses hexadecimal masks to select which CPU cores tasks should bind to
 - Bits ordered right to left
 - First bit masks core #0
 - Each task need its mask
- Single mask for 7 cores out of 8 (disabling core #0)
 - Core numbers: 76543210
 - Binary mask: 11111110
 - Hexadecimal value: 0xfe
 - Leading zeros can be omitted, but each element can still be very long

Task-to-CPU binding with Slurm: Examples

- `salloc --nodes=1 --partition=standard-g`
`module load LUMI/23.09 partition/G lumi-CPEtools/1.1-cpeGNU-23.09`
`srun --ntasks=8 --cpu-bind=map_cpu:49,57,17,25,1,9,33,41 mpi_check -r`
 - Example will be relevant for LUMI-G
- `srun --ntasks=8 --cpu-bind=mask_cpu:\`
`7e000000000000,7e00000000000000,7e0000,7e000000,7e,7e00,7e00000000,7e0000000000 \`
`hybrid_check -r`
 - Like the above but now enabling 6 cores per CCD (1-6).
 - Masks with use of both hardware threads can become extremely long, certainly on LUMI-C...
 - Playing with `--cpus-per-task` and then further restricting with OpenMP environment variables may be the easier way on LUMI-C
- Do not combine with `-c/--cpus-per-task!`

Task-to-GPU binding with Slurm

- Currently not recommended on LUMI
 - The control groups mechanism that Slurm uses breaks Peer2Peer IPC for GPU-aware MPI
- `srun --gpu-bind=[{quiet|verbose},]<type>`
- Some `<type>` options are for automatic binding
 - `--gpu-bind=none` is the most useful variant on LUMI: Turns off Slurm binding
 - `--gpu-bind=closest` is broken on LUMI
 - Other options: See the manual
- Other `<type>` options for fully manual distribution
 - `--gpu-bind=map_cpu:<gpu_id_for_task_0>,<gpu_id_for_task_1>,...` : Specify a single GPU for each task on the node
 - `--gpu-bind=mask_cpu:<mask_for_task_0>,<mask_for_task_1>,...` : Specify multiple GPUs via a mask (but only 2 hexadecimal digits as there are only 8 GPUs per node)

MPI rank redistribution with Cray MPICH

- Default behaviour: MPI rank i on task i
- Cray MPICH has its own mechanism to reorder MPI ranks on Slurm tasks that is more powerful than Slurm's
 - Best to use block distribution in Slurm for this.
 - `export MPICH_RANK_REORDER_METHOD=0` : Round-robin (like Slurm cyclic ordering)
 - `export MPICH_RANK_REORDER_METHOD=1` : Default, preserve the ordering from Slurm
 - `export MPICH_RANK_REORDER_METHOD=2` : Folded rank placement: First assign ranks on first task slot of each node from 0 till ..., then assign a rank on the second task slot but now from ... till 0, and so on.
 - `export MPICH_RANK_REORDER_METHOD=3` : Custom ordering set by the file `MPICH_RANK_ORDER` (or `$MPICH_RANK_REORDER_FILE`)
- The CPE has profiling tools that help you determine the optimal rank ordering
- See the 4-day LUMI Comprehensive course for more details

Refining core binding in OpenMP

- Slurm will assign cores up to the task/process level
 - Special case: Batch job step: All hardware threads of all cores of the first node of the job
- Thread-level control in OpenMP through library functions or environment variables
 - Debug: `export OMP_DISPLAY_AFFINITY=true`
 - `export OMP_NUM_THREADS=<num>` : Set number of threads
 - Multiple comma-separated numbers possible for multi-level parallelism
 - `OMP_PLACES` to define the places to use for binding: hardware thread level, core level or socket level, or an explicit list
 - `OMP_PROC_BIND` to set distribution and binding strategy over places
- Single level parallelism: Experiment with `omp_check` and `hybrid_check` in `lumi-CPETools`

Refining core binding in OpenMP: OMP_PLACES

- Defines the places to use for binding
 - `OMP_PLACES=threads` : OpenMP threads restricted to a single hardware thread (default)
 - `OMP_PLACES=cores` : OpenMP threads restricted to both hardware threads of a core
 - `OMP_PLACES=socket` : OpenMP threads restricted to all hardware threads of a single socket
 - Or define a set of locations (very technical)
`export OMP_PLACES="{0,1,2,3},{8,9,10,11},{16,17,18,19}"`
`export OMP_PLACES="{0:4}:3:8"`
 - Core numbers here are relative to those available to the process and not physical numbers

Refining core binding in OpenMP: OMP_PROC_BIND

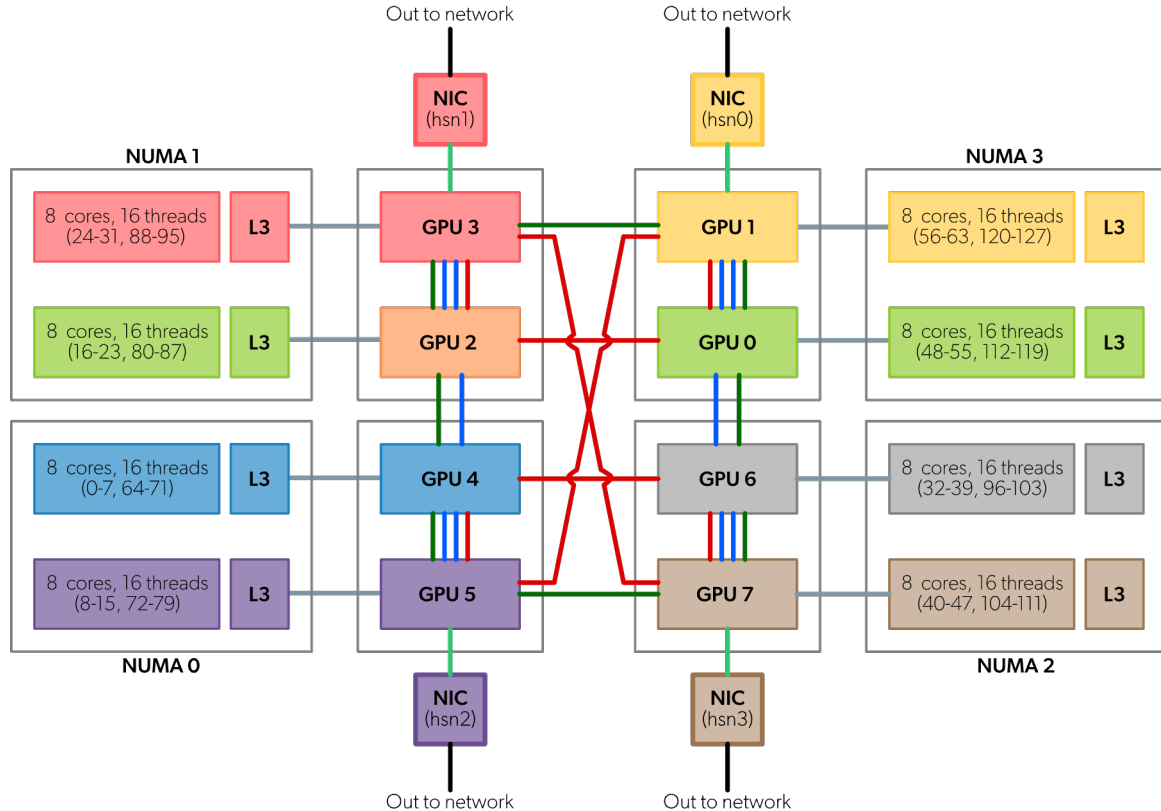
- Distribution over the places and binding selection:
 - `OMP_PROC_BIND=false` : Turn off OpenMP thread binding, use the task affinity mask
 - `OMP_PROC_BIND=close` : Try to keep the OpenMP threads as close as possible with one in each place (unless oversubscribed)
 - `OMP_PROC_BIND=spread` : Try to spread the OpenMP threads out as much as possible
 - `OMP_PROC_BIND=master` : Keep threads in the same place as the master thread.
 - Mostly useful if the place is a socket
- Multiple comma-separated options possible for nested parallelism
- Non-standard option in CCE: `auto` which is the default (other compilers: `false`)
 - CCE does a very reasonable job in many cases
- Many implementations have additional environment variables to tune the distribution

GPU binding with `ROCR_VISIBLE_DEVICES`

- Works at a very low level of the ROCm software stack
- Limits visibility to certain GPUs for all applications using the ROCm runtime
 - So also covers HIP and OpenCL
- Value: Comma-separated list of all device indices exposed to the application
 - Uses the local numbering in the control group
- Differences with affinity masks for CPUs
 - Affinity masks are OS-controlled
 - Therefore the OS can ensure you can only make masks more restrictive than the parent
 - Affinity masks always use the global numbering of hardware threads while `ROCR_VISIBLE_DEVICES` uses the local numbering in the control group

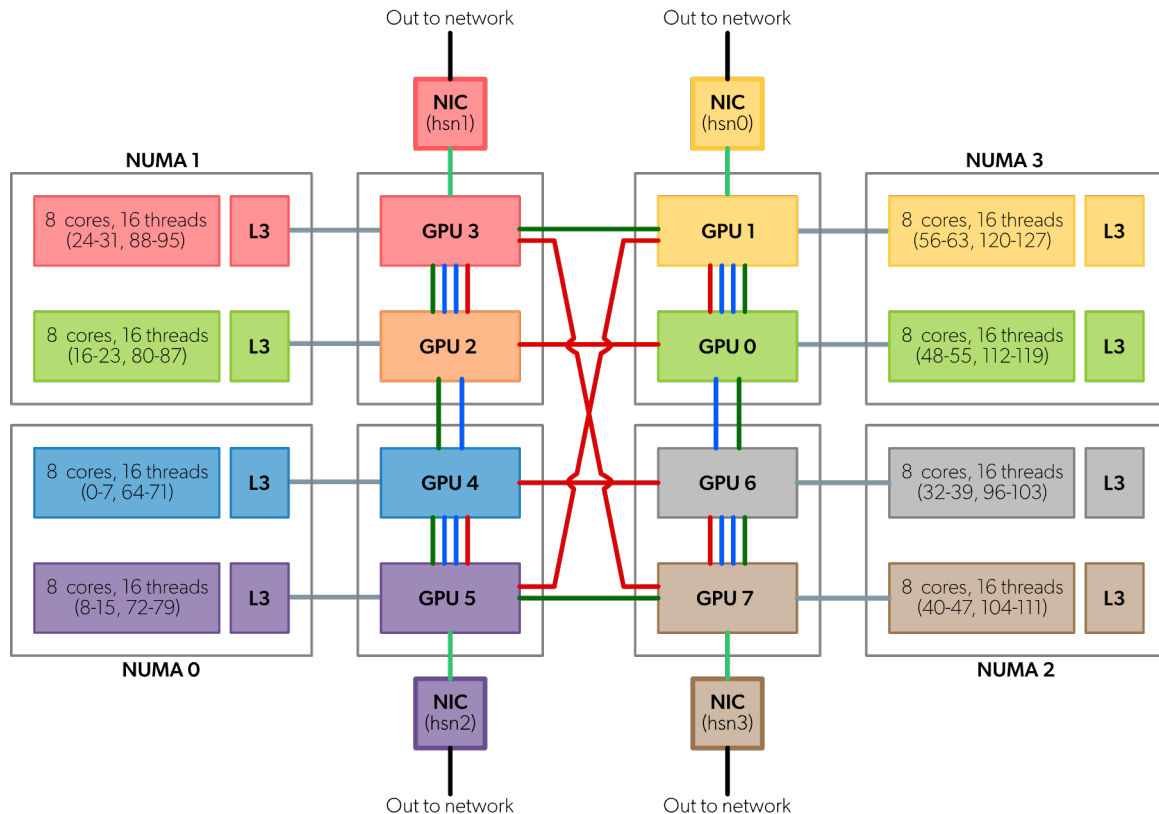
GPU binding: Optimal mapping (1)

CCD	Available HWTs	GCD
0	1-7, 65-71	4
1	9-15, 73-79	5
2	17-23, 81-87	2
3	25-32, 89-95	3
4	33-39, 97-103	6
5	41-47, 105-111	7
6	49-55, 113-119	0
7	57-63, 121-127	1



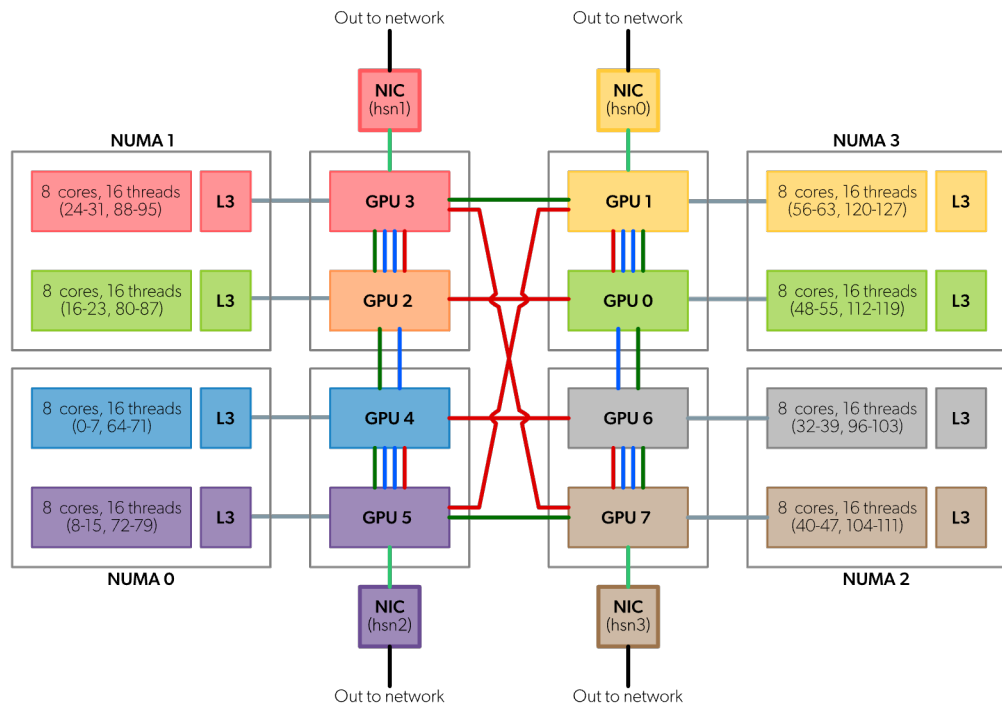
GPU binding: Optimal mapping (2)

GCD	CCD	Available HWTs
0	6	49-55, 113-119
1	7	57-63, 121-127
2	2	17-23, 81-87
3	3	25-32, 89-95
4	0	1-7, 65-71
5	1	9-15, 73-79
6	4	33-39, 97-103
7	5	41-47, 105-111



GPU binding: Embedded rings

- Green ring: 0 - 1 - 3 - 2 - 4 - 5 - 7 - 6 - 0
- Red ring: 0 - 1 - 5 - 4 - 6 - 7 - 3 - 2 - 0



GPU binding: Implementation

- Combination of two mechanisms:
 - CPU side: Use `--cpu-bind`, or in some cases simply `--cpus-per-task`
 - GPU side: Manual binding required by setting `ROCR_VISIBLE_DEVICES` because Slurm uses a mechanism with unwanted side effects.
 - Use a wrapper script that computes the proper GPU(s) from the Slurm local task id, sets `ROCR_VISIBLE_DEVICES` and then starts the application

GPU binding: Linear GCD, match cores (1)

L U M I

```
...
#SBATCH --partition=standard-g
#SBATCH --gpus-per-node=8
...
cat << EOF > select_gpu_${SLURM_JOB_ID}
#!/bin/bash
export ROCR_VISIBLE_DEVICES=\${SLURM_LOCALID}
exec \${*}
EOF
chmod +x select_gpu_${SLURM_JOB_ID}
...
CPU_BIND1="map_cpu:49,57,17,25,1,9,33,41"
srun --ntasks=$((SLURM_NNODES*8)) --cpu-bind=${CPU_BIND1} \
    ./select_gpu_${SLURM_JOB_ID} gpu_check -l
```



GPU binding: Linear GCD, match cores (2)

```
...
#SBATCH --partition=standard-g
#SBATCH --gpus-per-node=8
...
cat << EOF > select_gpu_${SLURM_JOB_ID}
#!/bin/bash
export ROCR_VISIBLE_DEVICES=\${SLURM_LOCALID}
exec \${*}
EOF
chmod +x select_gpu_${SLURM_JOB_ID}
...
CPU_BIND2="mask_cpu:0xfe000000000000,0xfe00000000000000"
CPU_BIND2="\${CPU_BIND2},0xfe0000,0xfe000000"
CPU_BIND2="\${CPU_BIND2},0xfe,0xfe00"
CPU_BIND2="\${CPU_BIND2},0xfe00000000,0xfe0000000000"
srun --ntasks=$((SLURM_NNODES*8)) --cpu-bind=${CPU_BIND2} \
    ./select_gpu_${SLURM_JOB_ID} gpu_check -l
```



GPU binding: Linear CCD, match GCD (1)

L U M I

```
...
#SBATCH --partition=standard-g
#SBATCH --gpus-per-node=8
...
cat << EOF > select_gpu_$SLURM_JOB_ID
#!/bin/bash
GPU_ORDER=(4 5 2 3 6 7 0 1)
export ROCR_VISIBLE_DEVICES=\${GPU_ORDER[\$SLURM_LOCALID]}
exec \$*
EOF
chmod +x select_gpu_$SLURM_JOB_ID
...
CPU_BIND1="map_cpu:1,9,17,25,33,41,49,57"
srun --ntasks=$((SLURM_NNODES*8)) --cpu-bind=$CPU_BIND1 \
    ./select_gpu_$SLURM_JOB_ID gpu_check -1
```



GPU binding: Linear CCD, match GCD (2)

L U M I

```
...
#SBATCH --partition=standard-g
#SBATCH --gpus-per-node=8
...
cat << EOF > select_gpu_$SLURM_JOB_ID
#!/bin/bash
GPU_ORDER=(4 5 2 3 6 7 0 1)
export ROCR_VISIBLE_DEVICES=\${GPU_ORDER[\$SLURM_LOCALID]}
exec \$*
EOF
chmod +x select_gpu_$SLURM_JOB_ID
...
CPU_BIND2="mask_cpu"
CPU_BIND2="$CPU_BIND2:0x00000000000000fe,0x000000000000fe00"
CPU_BIND2="$CPU_BIND2,0x0000000000fe0000,0x00000000fe000000"
CPU_BIND2="$CPU_BIND2,0x000000fe00000000,0x0000fe0000000000"
CPU_BIND2="$CPU_BIND2,0x00fe000000000000,0xfe00000000000000"
srun --ntasks=$((SLURM_NNODES*8)) --cpu-bind=$CPU_BIND2 \
    ./select_gpu_$SLURM_JOB_ID gpu_check -l
```



GPU binding: Linear CCD, match GCD (3)

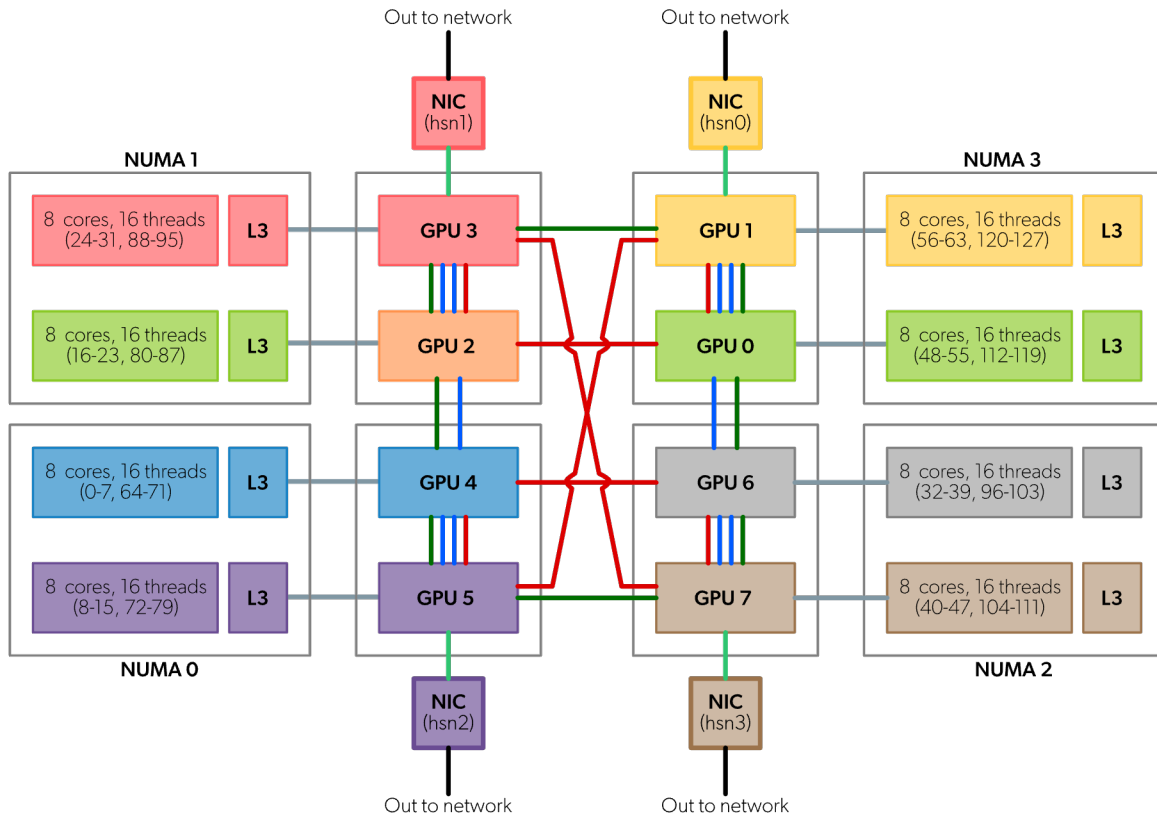
L U M I

```
...
#SBATCH --partition=standard-g
#SBATCH --gpus-per-node=8
...
cat << EOF > select_gpu_$SLURM_JOB_ID
#!/bin/bash
GPU_ORDER=(4 5 2 3 6 7 0 1)
export ROCR_VISIBLE_DEVICES=${GPU_ORDER[\\$SLURM_LOCALID]}
exec \\$*
EOF
chmod +x select_gpu_$SLURM_JOB_ID
...
srun --ntasks=$((SLURM_NNODES*8)) --cpus-per-task=7 \
    ./select_gpu_$SLURM_JOB_ID gpu_check -1
...
export OMP_NUM_THREADS=6
srun --ntasks=$((SLURM_NNODES*8)) --cpus-per-task=7 \
    ./select_gpu_$SLURM_JOB_ID gpu_check -1
```



GPU binding: Green ring (1)

Task	GCD	CCD	HWTs
0	0	6	49-55, 113-119
1	1	7	57-63, 121-127
2	3	3	25-32, 89-95
3	2	2	17-23, 81-87
4	4	0	1-7, 65-71
5	5	1	9-15, 73-79
6	7	5	41-47, 105-111
7	6	4	33-39, 97-103



GPU binding: Green ring (2)

L U M I

```
...
#SBATCH --partition=standard-g
#SBATCH --gpus-per-node=8
...
cat << EOF > select_gpu_$SLURM_JOB_ID
#!/bin/bash
GPU_ORDER=(0 1 3 2 4 5 7 6)
export ROCR_VISIBLE_DEVICES=\${GPU_ORDER[\$SLURM_LOCALID]}
exec \$*
EOF
chmod +x select_gpu_$SLURM_JOB_ID
...
CPU_BIND1="map_cpu:49,57,25,17,1,9,41,33"
srun --ntasks=$((SLURM_NNODES*8)) --cpu-bind=$CPU_BIND1 \
    ./select_gpu_$SLURM_JOB_ID gpu_check -1
```



GPU binding: Green ring (3)

L U M I

```
cat << EOF > select_gpu_${SLURM_JOB_ID}
#!/bin/bash
GPU_ORDER=(0 1 3 2 4 5 7 6)
export ROCR_VISIBLE_DEVICES=\${GPU_ORDER[\${SLURM_LOCALID}]}
exec \${*}
EOF
chmod +x select_gpu_${SLURM_JOB_ID}
...
CCD_MASK=( 0x00000000000000fe \
           0x0000000000000fe00 \
           0x0000000000fe0000 \
           0x00000000fe000000 \
           0x000000fe00000000 \
           0x0000fe0000000000 \
           0x00fe000000000000 \
           0xfe00000000000000 )
CPU_BIND2="mask_cpu"
CPU_BIND2="\${CPU_BIND2:\${CCD_MASK[6]},\${CCD_MASK[7]}}"
CPU_BIND2="\${CPU_BIND2,\${CCD_MASK[3]},\${CCD_MASK[2]}}"
CPU_BIND2="\${CPU_BIND2,\${CCD_MASK[0]},\${CCD_MASK[1]}}"
CPU_BIND2="\${CPU_BIND2,\${CCD_MASK[5]},\${CCD_MASK[4]}}"
srun --ntasks=$((SLURM_NNODES*8)) --cpu-bind=\${CPU_BIND2} \
    ./select_gpu_${SLURM_JOB_ID} gpu_check -1
```



“Allocate by resources” partitions

- Proper binding not possible
- Slurm will use a control group per task for the GPUs
 - You almost have to use `--gpus-per-task` to ensure that GPUs and tasks are on the same nodes (unless you use just a single node)
 - Problems with Peer2Peer IPC
 - Solution:
 - Turn off with `--gpu-bind=none`
 - This will number visible GPUs for the job on each node from 0,
 - and we can then again use the local task ID to assign a GPU to each task via `ROCR_VISIBLE_DEVICES` via the `select_gpu` script trick.
- Optimal mapping is not possible

Questions?

