AGENDA

Multi-GPU Programming Models

Multi-GPU Programming with OpenACC and CUDA
SUMMIT NODE
(2) IBM POWER9 + (6) NVIDIA VOLTA V100

CPU 0

0 (0-3)
1 (4-7)
2 (8-11)
3 (12-15)
4 (16-19)
5 (20-23)
6 (24-27)

14 (56-59)
15 (60-63)
16 (64-67)
17 (68-71)
18 (72-75)
19 (76-79)
20 (80-83)

GPU 0

16 GB (HBM2)

GPU 1

16 GB (HBM2)

GPU 2

16 GB (HBM2)

CPU 1

22 (68-91)
23 (92-95)
24 (96-99)
25 (100-103)
26 (104-107)
27 (108-111)
28 (112-115)

30 (120-123)
31 (124-127)
32 (128-131)
33 (132-135)
34 (136-139)
35 (140-143)

36 (144-147)
37 (148-151)
38 (152-155)
39 (156-159)
40 (160-163)
41 (164-167)
42 (168-171)

GPU 3

16 GB (HBM2)

GPU 4

16 GB (HBM2)

GPU 5

16 GB (HBM2)

256 GB (DDR4)

135 GB/s

256 GB (DDR4)

135 GB/s

64 GB/s

NVLink2

(50 GB/s)

(900 GB/s)
MULTI-GPU PROGRAMMING MODELS
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Single Thread, Multiple GPUs
• A single thread will change devices as-needed to send data and kernels to different GPUs

Multiple Threads, Multiple GPUs
• Using OpenMP, Pthreads, or similar, each thread can manage its own GPU

Multiple Ranks, Single GPU
• Each rank acts as-if there’s just 1 GPU, but multiple ranks per node use all GPUs

Multiple Ranks, Multiple GPUs
• Each rank manages multiple GPUs, multiple ranks/node. Gets complicated quickly!
MULTI-GPU PROGRAMMING MODELS

Trade-offs Between Approaches

- Single Thread, Multiple GPUs
  - Conceptually Simple
  - Requires additional loops
  - CPU can become a bottleneck
  - Remaining CPU cores often underutilized

- Multiple Threads, Multiple GPUs
  - Conceptually Very Simple
  - Set and forget the device numbers
  - Relies on external Threading API
  - Can see improved utilization
  - Watch affinity

- Multiple Ranks, Single GPU
  - Little to no code changes required
  - Re-uses existing domain decomposition
  - Can see improved utilization
  - Watch affinity

- Multiple Ranks, Multiple GPUs
  - Easily share data between peer devices
  - Coordinating between GPUs extremely tricky
MULTI-DEVICE CUDA

CUDA by default exposes all devices, numbered 0 - (N-1), if devices are not all the same, it will reorder the “best” to device 0.

Each device has its own pool of streams.

If you do nothing, all work will go to Device #0.

Developer must change the current device explicitly
MULTI-DEVICE OPENACC

OpenACC presents devices numbered 0 - (N-1) for each device type available.

The order of the devices comes from the runtime, almost certainly the same as CUDA.

By default all data and work go to the current device.

Developers must change the current device and maybe the current device type using an API.
MULTI-DEVICE OPENMP

OpenMP devices numbered 0 - (N-1) for ALL devices on the machine, including the host.

The order is determined by the runtime, but devices of the same type are contiguous.

To change the device for data and compute a clause is added to directives.

Device API routines include a devicenum
MULTI-GPU PROGRAMMING WITH OPENACC AND CUDA
MULTI-GPU W/ CUDA AND OPENACC

The CUDA and OpenACC approaches are sufficiently similar, that I will demonstrate using OpenACC.

Decoder Ring:

<table>
<thead>
<tr>
<th>OpenACC</th>
<th>CUDA</th>
</tr>
</thead>
<tbody>
<tr>
<td>acc_get_device_type()</td>
<td>N/A</td>
</tr>
<tr>
<td>acc_set_device_type()</td>
<td>N/A</td>
</tr>
<tr>
<td>acc_set_device_num()</td>
<td>cudaSetDevice()</td>
</tr>
<tr>
<td>acc_get_device_num()</td>
<td>cudaGetDevice()</td>
</tr>
<tr>
<td>acc_get_num_devices()</td>
<td>cudaGetDeviceCount()</td>
</tr>
</tbody>
</table>
Multi-Device Pipeline

A Case Study

We’ll use a simple image filter to demonstrate these techniques.

No inter-GPU communication required

Pipelining: Breaking a large operation into smaller parts so that independent operations can overlap.

Since each part is independent, they can easily be run on different devices. We will extend the filter to run on more than one device.
Pipelining in a Nutshell

Two Independent Operations Serialized

Overlapping Copying and Computation

NOTE: In real applications, your boxes will not be so evenly sized.
Multi-device Pipelining in a Nutshell

Device 0

H2D | kernel | D2H

Device 1

H2D | kernel | D2H

H2D | kernel | D2H
Pipelined Code

#pragma acc data create(imgData[w*h*ch], out[w*h*ch])
            copyin(filter)
{
    for (long blocky = 0; blocky < nbblocks; blocky++)
    {
        long starty = MAX(0, blocky * blocksize - filtersize/2);
        long endy = MIN(h, starty + blocksize + filtersize/2);
        #pragma acc update device(imgData[starty*step:blocksize*step]) async(block%3)
            starty = blocky * blocksize;
            endy = starty + blocksize;
        #pragma acc parallel loop collapse(2) gang vector async(block%3)
            for (y=starty; y<endy; y++) for (x=0; x<w; x++)
            {
                <filter code omitted>
                out[y * step + x * ch] = 255 - (scale * blue);
                out[y * step + x * ch + 1] = 255 - (scale * green);
                out[y * step + x * ch + 2] = 255 - (scale * red);
            }
        #pragma acc update self(out[starty*step:blocksize*step]) async(block%3)
    }
#pragma acc wait
}
#pragma acc data create(imgData[w*h*ch],out[w*h*ch])
        copyin(filter)
{
    for ( long blocky = 0; blocky < nblocks; blocky++)
    {
        long starty = MAX(0,blocky * blocksize - filtersize/2);
        long endy   = MIN(h,starty + blocksize + filtersize/2);
        #pragma acc update device(imgData[starty*step:blocksize*step]) async(block%3)
        starty = blocky * blocksize;
        endy = starty + blocksize;
        #pragma acc parallel loop collapse(2) gang vector async(block%3)
        for (y=starty; y<endy; y++) for ( x=0; x<w; x++ ) {
            <filter code ommitted>
            out[y * step + x * ch] = 255 - (scale * blue);
            out[y * step + x * ch + 1 ] = 255 - (scale * green);
            out[y * step + x * ch + 2 ] = 255 - (scale * red);
        }
        #pragma acc update self(out[starty*step:blocksize*step]) async(block%3)
    }
    #pragma acc wait
}
NVPROF Timeline of Pipeline
Extending to multiple devices

Create 1 OpenMP thread on the CPU per-device. This is not strictly necessary, but simplifies the code.

Within each thread, set the device number.

Divide the blocks as evenly as possible among the CPU threads.
Multi-GPU Pipelined Code (OpenMP)

```c
#pragma omp parallel num_threads(acc_get_num_devices(acc_device_default))
{
    acc_set_device_num(omp_get_thread_num(),acc_device_default);
    int queue = 1;
    #pragma acc data create(imgData[w*h*ch],out[w*h*ch])
    #pragma omp for schedule(static)
    for ( long blocky = 0; blocky < nblocks; blocky++) {
        // For data copies we need to include the ghost zones for the filter
        long starty = MAX(0,blocky * blocksize - filtersize/2);
        long endy   = MIN(h,starty + blocksize + filtersize/2);
        #pragma acc update device(imgData[starty*step:(endy-starty)*step]) async(queue)
        starty = blocky * blocksize;
        endy   = starty + blocksize;
        #pragma acc parallel loop collapse(2) gang vector async(queue)
        for ( long y = starty; y < endy; y++ ) { for ( long x = 0; x < w; x++ ) {
            <filter code removed for space>
        }}
        #pragma acc update self(out[starty*step:blocksize*step]) async(queue)
        queue = (queue%3)+1;
    }
    #pragma acc wait
}
```
Multi-GPU Pipelined Performance

Source: PGI 17.3, NVIDIA Tesla P100 (DGX-1)
OpenACC with MPI

Domain decomposition is performed using MPI ranks

Each rank should set its own device

• Maybe acc_set_device_num

• Maybe handled by environment variable (CUDA_VISIBLE_DEVICES)

GPU affinity can be handled by standard MPI task placement

Multiple MPI Ranks/GPU (using MPS) can work in place of OpenACC work queues/CUDA Streams
Setting a device by local rank

// This is not portable to other MPI libraries
char *comm_local_rank = getenv("OMPI_COMM_WORLD_LOCAL_RANK");
int local_rank = atoi(comm_local_rank);
char *comm_local_size = getenv("OMPI_COMM_WORLD_LOCAL_SIZE");
int local_size = atoi(comm_local_size);
int num_devices = acc_get_num_devices(acc_device_nvidia);
#pragma acc set device_num(local_rank%num_devices) \
    device_type(acc_device_nvidia)

You may also try using MPI_Comm_split_type() using
MPI_COMM_TYPE_SHARED or OMPI_COMM_TYPE_SOCKET.

In the end, you need to understand how jsrun/mpirun is placing your ranks.
MPI Image Filter (pseudocode)

if (rank == 0 ) read_image();
// Distribute the image to all ranks
MPI_Scatterv(image);

MPI_Barrier(); // Ensures all ranks line up for timing
omp_get_wtime();
blur_filter(); // Contains OpenACC filter
MPI_Barrier(); // Ensures all ranks complete before timing
omp_get_wtime();

MPI_Gatherv(out);
if (rank == 0 ) write_image();

$ jsrun -n 6 -a 1 -c 1 -g 1 ...

There’s a variety of ways to do MPI decomposition, this is what I used for this particular example.
Multi-GPU Pipelined Performance (MPI)

Speed-up from one device

1 Device: 1.00X
2 Devices: 1.53X
4 Devices: 2.83X
8 Devices: 4.89X

Crosses quad boundary

Source: PGI 17.3, NVIDIA Tesla P100 (DGX-1), Communication Excluded
Multi-GPU Pipelined Performance (MPI)

Source: PGI 17.3, NVIDIA Tesla P100 (DGX-1), Communication Excluded
#pragma omp parallel
{
    cudaSetDevice(idx);
    #pragma omp for
    for (int b=0; b < nblocks; b++)
    {
        cudaMemcpyAsync(..., streams[b%3]);
        blur_kernel <<<griddim, blockdim, 0, streams[b%3]>>>();
        cudaMemcpyAsync(..., streams[b%3]);
    }
    cudaDeviceSynchronize();
}

MPI_Comm_rank(local_comm, &local_rank);
cudaSetDevice(local_rank);

for (int b=0; b < nblocks; b++)
{
    cudaMemcpyAsync(..., streams[b%3]);
    blur_kernel <<<griddim, blockdim, 0, streams[b%3]>>>();
    cudaMemcpyAsync(..., streams[b%3]);
}
cudaDeviceSynchronize();
MULTI-DEVICE OPENMP 4.5

Same Pattern, Different API

```c
#pragma omp parallel num_threads(num_dev)
{
    #pragma omp for
    for ( int b=0; b < nblocks; b++)
    {
        #pragma omp target update map(to:...)
        device(dev) depend(inout:A)
        nowait
        #pragma omp target teams distribute
        parallel for simd device(dev)
        depend(inout:A)
        for(...) { ... }
        #pragma omp target update map(from:...)
        device(dev) depend(inout:A)
        nowait
    }
    #pragma omp taskwait
}

MPI_Comm_rank(local_comm, &local_rank);
int dev = local_rank;
for ( int b=0; b < nblocks; b++)
{
    #pragma omp target update map(to:...)
    device(dev) depend(inout:A)
    nowait
    #pragma omp target teams distribute
    parallel for simd device(dev)
    depend(inout:A)
    for(...) { ... }
    #pragma omp target update map(from:...)
    device(dev) depend(inout:A)
    nowait
}
#pragma omp taskwait
```
Multi-GPU Approaches

Choosing an approach

**Single-Threaded, Multiple-GPUs** - Requires additional loops to manage devices, likely undesirable.

**Multi-Threaded, Multiple-GPUs** - Very convenient set-and-forget the device. Could possibly conflict with existing threading.

**Multiple-Ranks, Single-GPU each** - Probably the simplest if you already have MPI, decomposition is done. Must get your MPI placement correct.

**Multiple-Ranks, Multiple-GPUs** - Can allow all GPUs to share common data structures. Only do this is you absolutely need to, difficult to get right.
CLOSING SUMMARY
MULTI-GPU APPROACHES
Choosing an approach

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GPU TO GPU COMMUNICATION

- CUDA aware MPI functionally portable
  - OpenACC/MP interoperable
  - Performance may vary between on/off node, socket, HW support for GPU Direct
  - WARNING: Unified memory support varies wildly between implementations!
- Single-process, multi-GPU
  - Enable peer access for straightforward on-node transfers
- Multi-process, single-gpu
  - Pass CUDA IPC handles for on-node copies
- Combine for more flexibility/complexity!
ESSENTIAL TOOLS AND TRICK

- Pick on-node layout with OLCF jsrun visualizer
- Select MPI/GPU interaction with jsrun --smiargs
  - “-gpu” for CUDA aware, “off” for pure GPU without MPI
- Profile MPI and NVLinks with nvprof
- Good performance will require experimentation!