





## CUDA: Concurrency and Other Topics

Ben Cumming, CSCS February 18, 2016





## Going 2D and 3D

#### Launch Configuration

- so far we have used one-dimensional launch configurations
  - threads in blocks indexed using threadIdx.x
  - blocks in a grid indexed using blockIdx.x
- many kernels map naturally onto 2D and 3D indexing
  - e.g. matrix-matrix operations
  - e.g. stencils





#### Full Launch Configuration

kernels launch dimensions can be specified with dim3 structs

```
kernel<<<dim3 gridDim, dim3 blockDim>>>(...);
```

- dim3.x, dim3.y and dim3.z specify the launch dimensions
- can be constructed with 1, 2 or 3 dimensions
- unspecified dim3 dimensions are set to 1

#### launch configuration examples

```
// 1D: 128x1x1 for 128 threads dim3 a(128);
// 2D: 16x8x1 for 128 threads dim3 b(16, 8);
// 3D: 16x8x4 for 512 threads dim3 c(16, 8, 4);
```





The threadIdx, blockDim, blockIdx and gridDim can be treated like 3D vectors via the .x, .y and .z members.

```
matrix addition example
__global__
void MatAdd(float *A. float *B. float *C. int n) {
    int i = blockIdx.x * blockDim.x + threadIdx.x;
    int j = blockIdx.y * blockDim.y + threadIdx.y;
    if(i<n && j<n) {
        auto pos = i + j*n;
C[pos] = A[pos] + B[pos];
int main() {
   dim3 threadsPerBlock(16. 16):
   dim3 numBlocks(n / threadsPerBlock.x, n / threadsPerBlock.y);
   MatAdd <<< numBlocks. threadsPerBlock>>> (A. B. C. n):
```



#### Exercise: Launch Configurations

- 2D stencil in diffusion/diffusion2d.cu
  - a plotting script is provided for visualizing the results
  - use a small domain for visualization
- **extra**: can you improve performance with shared memory?

```
cd cuda/exercises/diffusion
make
aprun diffusion2d.cuda 8
module load python/2.7.6
python plotting.py
```









#### MPI with GPUs

#### MPI with data in device memory

Our GPU-accelerated applications use GPUs to parallelize onnode computation

and MPI for communication between nodes

It is likely that communication between MPI ranks will involve information on the GPU

- 1. allocate buffers in host memory
- 2. manually copy from device→host memory
- 3. perform MPI communication with host buffers
- 4. copy received data from host→device memory

This approach can be very fast:

 have a CPU thread dedicated to asynchronous host⇔device and MPI communication



#### GPU-Aware MPI

GPU-aware MPI implementations can automatically handle MPI transactions with pointers to GPU memory

- MVAPICH 2.0
- OpenMPI since version 1.7.0
- Cray MPI

#### How it works

- each pointer passed to MPI is checked to see if it is in host or device memory.
- small messages between GPUs (up to ≈8 k) are copied directly with RDMA
- larger messages are **pipelined** via host memory





#### How to use G2G communication

• set the environment variable

```
export MPICH_RDMA_ENABLED_CUDA=1
```

- if not set, MPI assumes that all pointers are to host memory, and your application will probably crash with segmentation faults
- experiment with the environment variable
   MPICH\_G2G\_PIPELINE
  - sets the maximum number of 512 kB message chunks that can be in flight (default 16)

#### MPI with G2G example

```
MPI_Request srequest, rrequest;
auto send_data = malloc_device < double > (100);
auto recv_data = malloc_device < double > (100);

// call MPI with GPU pointers
MPI_Irecv(recv_data, 100, MPI_DOUBLE, source, tag, MPI_COMM_WORLD,
    &rrequest);
MPI_Isend(send_data, 100, MPI_DOUBLE, target, tag, MPI_COMM_WORLD,
    &srequest);
```



#### Capabilities and Limitations

- support for most MPI API calls (point-to-point, collectives, etc)
- robust support for common MPI API calls
  - i.e. point-to-point operations
- no support for user-defined MPI data types





#### Exercise: MPI with G2G

- 2D stencil with MPI in diffusion/diffusion2d\_mpi.cu
- copy the kernel and kernel lanuch from previous example
  - 1. implement MPI communication with host buffering
  - 2. implement MPI communication with G2G
  - 3. can you observe any performance differences between the two?
  - 4. why are we restricted to just 1 MPI rank per node, i.e.

aprun -N 1

```
cd cuda/exercises/diffusion
make
aprun -n 2 -N 1 diffusion2d_mpi.cuda 8
module load python/2.7.6
python plotting.py
# once it gets the correct results:
sbatch job.daint
```





#### Exercises: 2D Diffusion with MPI Results

Time for 1000 t	ime step	os $128 \times 16$	,382	
	nodes	G2G off	G2G on	
	1	0.479	0.479	
	2	0.277	0.274	
	4	0.183	0.180	
	8	0.152	0.151	
	16	0.167	0.117	









## Concurrency

#### Concurrency

**Concurrency** is the ability to perform multiple CUDA operations simultaneously

- CUDA kernels
- copying from host to device
- copying from device to host
- operations on the host CPU

#### Concurrency enables

- both CPU and GPU can work at the same time
- multiple tasks can be run on GPU simultaneously
- overlapping of communication and computation





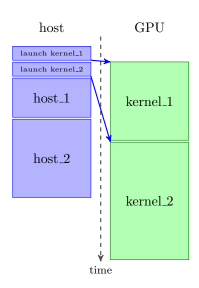
# Host code kernel\_1<<<...>>>(...); kernel\_2<<<...>>>(...); host\_1(...); host\_2(...);

#### The host:

- launches the two CUDA kernels
- then executes host calls sequentially

#### The GPU:

- executes asynchronously to host
- executes kernels sequentially





The CUDA language and runtime libraries provide mechanisms for coordinating asynchronous GPU execution

- CUDA streams can concurrently run independent kernels and memory transfers
- CUDA events can be used to synchronize streams and query the status of kernels and transfers





#### Streams

A CUDA stream is a sequence of operations that execute in issue order on the GPU

 CUDA operations are kernels and copies between host and device memory spaces

#### Streams and concurrency

- operations in different streams **may** run concurrently
  - there have to be sufficient resources on the GPU (registers, shared memory, blocks, etc)
- operations in the same stream **are** executed sequentially
- if no stream is specified, all kernels are launched in the default stream

#### Managing streams

A stream is represented using a cudaStream\_t type

- cudaStreamCreate(cudaStream\_t\* s) and
   cudaStreamDestroy(cudaStream\_t s) can be used to create and free CUDA streams respectively
- To launch a kernel on a stream specify the stream id as a fourth parameter to the launch syntax

```
kernel<<<grid_dim, block_dim, shared_size, stream>>>(...)
```

the default CUDA stream is the NULL stream, or stream 0
 (cudaStream\_t is an integer)

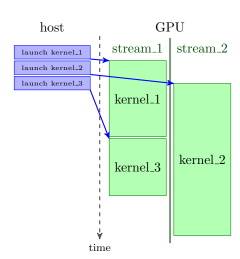
#### Basic cuda stream usage

```
// create stream
cudaStream_t stream;
cudaStreamCreate(&stream);
// launch kernel in stream
my_kernel <<<grid_dim, block_dim, shared_size, stream>>>(..)
// release stream when finished
cudaStreamDestroy(stream);
```



#### Host code kernel\_1 <<< , , , stream\_1 >>> (); kernel\_2 <<< , , , stream\_2 >>> (); kernel\_3 <<< , , , stream\_1 >>> ();

- kernel\_1 and kernel\_3 are serialized in stream\_1
- kernel\_2 can run asynchronously in stream\_2
- note that kernel\_2 will only run concurrently if there are sufficient resources available on the GPU, i.e. if kernel\_1 is not using all of the SMXs.





#### Asynchronous copy

#### cudaMemcpyAsync(\*dst, \*src, size, kind, cudaStream\_t stream = 0);

- takes an additional parameter stream, which is 0 by default
- returns immediately after initiating copy
  - host can do work while copy is performed
  - only if **pinned memory** is used
- copies in the same direction (i.e. H2D or D2H) are serialized
  - copies in opposite directions are concurrent if in different streams





#### What is pinned memory?

Pinned memory (or page-locked) memory will not be paged out to disk when memory runs low

- the GPU can safely remotely read/write the memory directly without host involvement
- only use for transfers, because it easy to run out of memory

#### Managing pinned memory

```
cudaMallocHost(**ptr, size); and cudaFreeHost(*ptr);
```

• allocate and free pinned memory (size is in bytes).

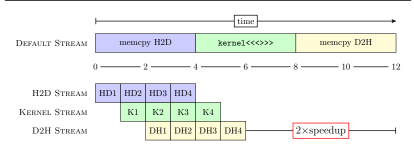




#### Asynchronous copy example: streaming workloads

Computations that can be performed independently, e.g. our axpy example:

- data in host memory has to be copied to the device, and the result copied back after the kernel is computed.
- we can overlap the copies with the kernel calls by breaking the data into chunks.





#### CUDA events

To implement the streaming workload we have to coordinate operations on the GPU. CUDA events can be used for this purpose.

- synchronize tasks in different streams, e.g.:
  - don't start kernel in kernel stream until data copy stream has finished.
  - wait until required data has finished copy from host before launching kernel
- query status of concurrent tasks
  - has kernel finished/started yet?
  - how long did a kernel take to compute?





#### Managing events

cudaEventCreate(cudaEvent\_t\*); and cudaEventDestroy(cudaEvent\_t);

create and free cudaEvent\_t

cudaEventRecord(cudaEvent\_t, cudaStream\_t);

enqueue an event in a stream

cudaEventSynchronize(cudaEvent\_t);

make host execution wait for event to occur.

cudaEventQuery(cudaEvent\_t)

• test if the work before an event in a queue has been completed

cudaEventElapsedTime(float\*, cudaEvent\_t, cudaEvent\_t);

get time between two events



#### Using events to time kernel execution

```
cudaEvent t start. end:
cudaStream_t stream;
float time_taken;
// initialize the events and streams
cudaEventCreate(&start);
cudaEventCreate(&end);
cudaStreamCreate(&stream):
cudaEventRecord(start, stream); // enqueue start in stream
my_kernel <<< grid_dim, block_dim, 0, stream>>>();
cudaEventRecord(end, stream); // enqueue end in stream
cudaEventSynchronize(end);  // wait for end to be reached
cudaEventElapsedTime(&time_taken, start, end);
std::cout << "kernel took " << 1000*time_taken << " s\n";
// free resources for events and streams
cudaEventDestroy(start);
cudaEventDestroy(end);
cudaStreamDestrov(stream):
```





#### Copy→kernel synchronization

```
cudaEvent_t event;
cudaStream_t kernel_stream, h2d_stream;
size_t size = 100*sizeof(double);
double *dptr. *hptr:
// initialize
cudaEventCreate(&event):
cudaStreamCreate(&kernel stream):
cudaStreamCreate(&h2d_stream);
cudaMalloc(&dptr, size);
cudaMallocHost(&hptr, size); // use pinned memory!
// start asynchronous copy in h2d_stream
cudaMemcpyAsync(dptr, hptr, size,
                cudaMemcpyHostToDevice, h2d_stream);
// enqueue event in stream
cudaEventRecord(event. h2d stream):
// make kernel_stream wait for copy to finish
cudaStreamWaitEvent(kernel stream. event. 0):
// enqueue my_kernel to start when event has finished
my_kernel << grid_dim, block_dim, 0, kernel_stream >>>();
// free resources for events and streams
cudaEventDestroy(event);
cudaStreamDestroy(h2d_stream);
cudaStreamDestroy(kernel_stream);
cudaFree(dptr):
cudaFreeHost(hptr);
```



#### Exercises

- 1. Open util.h in cuda/examples/async and understand copy\_to\_{host/device}\_async() and malloc\_pinned\_host()
- 2. Open CudaEvent.h and CudaStream.h
  - what is the purpose of these classes?
  - what does CudaStream::enqueue\_event() do?
- 3. Open memcopy1.cu and run
  - what does the benchmark test?
  - what is the effect of turning on USE\_PINNED? Hint: try small and large values for n (8, 16, 20, 24)
- 4. Inspect memcopy2.cu and run
  - what effect does changing the number of chunks have?
- 5. Inspect memcopy3.cu and run
  - how does it differ from memcopy2.cu?
  - what effect does changing the number of chunks have?



#### Using events to time kernel execution: with helpers

```
CudaStream stream(true);
auto start = stream.enqueue_event();
my_kernel <<< grid_dim , block_dim , 0 , stream .stream() >>> ();
auto end = stream.enqueue event():
end.wait():
auto time_taken = end.time_since(start);
std::cout << "kernel took " << 1000*time taken << " s\n":
```

#### Copy—kernel synchronization: with helpers

```
CudaStream kernel_stream(true), h2d_stream(true);
auto size = 100:
auto dptr = device_malloc < double > (size);
auto hptr = pinned_malloc < double > (size);
copy_to_device_async < double > (hptr,dptr,size,h2d_stream.stream());
auto event = h2d_stream.enqueue_event();
kernel stream.wait on event(event):
my_kernel << grid_dim, block_dim, 0, kernel_stream.stream()>>>();
cudaFree(dptr):
cudaFreeHost(hptr):
```









## Profiling CUDA

#### Profiling CUDA applications

To analyze concurrent applications we need tools that can visually represent application flow.

The CUDA toolkit provides the tools **nvprof** and **nvvp** for profiling our GPU applications

- there are visual tools for Windows and Eclipse too
- they work for OpenACC applications too





#### nvprof

#### **nvprof** is a command line tool

- can be used to generate text reports
- nvprof --help for a full list of options
- nvprof app.exe will perform basic profling of application and print text summary
- nvprof -o profile.out app.exe will save profile information to file profile.out for visualization with nvvp

#### Demonstration

Use nyprof on the memcopy test codes



#### nvvp

**nvvp** is a graphical tool for visualizing CUDA applications

- can also be used to perform interactive profiling and guided analysis
- this is not so easy on Cray systems
- we can also use the output from nvprof
  - use nvprof -o profile.out ... ./app.out to generate detailed analysis
  - this can take a long time, because each kernel has to be replayed multiple times to collect all of the information required for the report.

#### Demonstration

Use nvvp on the output of nvprof for the memcopy examples

#### Some rough guidelines for concurrency

Ideally for most workloads you don't want to rely on streams to fill the GPU with work

- a sign that the working set per GPU is not large enough
- full concurrency is difficult in practice
  - a low-level optimization strategy for the last few %
- this isn't a hard and fast rule

Streams come into their own for overlapping communication and computation

 possible to transfer data in both directions concurrently with kernels execution







## Final Thoughts

#### Is this so hard?

Each topic so far has not been too complicated:

- moving memory between host and device
- writing kernels
- parallel kernel launching
- ... ok, concurrency is hard (but concurrency is hard on the CPU too)

However, there is a lot that you have to keep in mind:

- where is my data?
- loop free thinking: individual work items to be performed in parallel
- synchronization between threads, host and device?

Combined this makes GPU programming tough to start with.

• OpenMP and OpenACC attempt to make things "easy", but they still have shortcomings

#### What next?

You have seen some features of CUDA

- not an exhaustive overview!
- there are some "essential" features that weren't covered

And more importantly, I hope that you have started to think about parallel work items

- a different perspective than thinking in loops
- an important skill for more than just GPU programming



