

Programming GPUs with CUDA

Day 2

Slides: <http://www.sharcnet.ca/~syam/CUDA2.pdf>

Sergey Mashchenko
SHARCNET
syam@sharcnet.ca

CO Summer School East
University of Ottawa, July 8 - 12, 2019

Outline

- Preliminary remarks
- C language extensions
- Graham GPUs, P100
- CUDA code optimization

Quiz 1

Link: <http://www.socrative.com>

Room: CUDADAY2

C language extensions

C language extensions

- Kernels (one block)

```
// Kernel definition
__global__ void VecAdd (float* d_A, float* d_B, float* d_C)
{
    int i = threadIdx.x;
    d_C[i] = d_A[i] + d_B[i];
}

int main()
{
    ...

// Kernel invocation with N threads
    VecAdd <<<1, N>>> (d_A, d_B, d_C);

    ...
}
```

Pointers to
device
addresses!



C language extensions

- Kernels (multi-block)

```
// Kernel definition
__global__ void VecAdd (float* d_A, float* d_B, float* d_C)
{
    int i = threadIdx.x + blockDim.x * blockIdx.x;
    d_C[i] = d_A[i] + d_B[i];
}

int main()
{
    ...

// M blocks with N threads each:
    VecAdd <<<M, N>>> (d_A, d_B, d_C);
    ...
}
```

C language extensions

- Static global device arrays

// Device code (outside of any function):

__device__ float d_A[10][50];

// Host code (outside of any function):

float h_A[10][50];

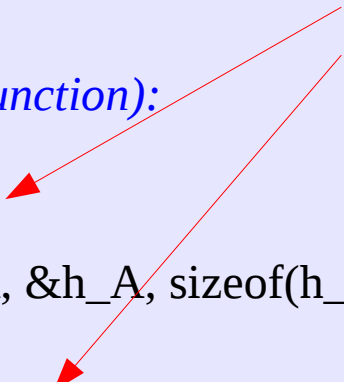
// Host to device copying:

cudaMemcpyToSymbol (d_A, &h_A, sizeof(h_A), 0, cudaMemcpyHostToDevice);

// Device to host copying:

cudaMemcpyFromSymbol (&h_A, d_A, sizeof(h_A), 0, cudaMemcpyDeviceToHost);

*Replaces two function calls:
cudaGetSymbolAddress();
cudaMemcpy();*



C language extensions

- Dynamic global device arrays

// Host code:

// Host array allocation (often cudaHostAlloc() is a better way):

```
size_t size = N * sizeof(float);  
float* h_A = (float*) malloc (size);
```

// Device array allocation:

```
float* d_A;  
cudaMalloc(&d_A, size);
```

// Host to device copying:

```
cudaMemcpy (d_A, h_A, size, cudaMemcpyHostToDevice);
```

// Device to host copying:

```
cudaMemcpy (h_A, d_A, size, cudaMemcpyDeviceToHost);
```


C language extensions

- Device functions

```
// Device code

// Function:
__device__ double my_function (double x)
{
    double f;
    ...
    return f;
}

// Kernel:
__global__ void my_kernel ();
{
    double f1, x1;
    f1 = my_function (x1);
}
```

C language extensions

- Shared memory
 - Can be much faster than global (device) memory
 - Shared across the threads in a single block
 - The amount is very limited, so it is often a limiting factor for CUDA optimization
 - Typically statically defined

// Device code

```
__shared__ double A_loc[BLOCK_SIZE];
```

C language extensions

- Execution synchronization on device
 - Can only be done within a single block.
 - As a result, only up to 1024 (usually between 64 and 256) threads can be synchronized.
 - Used when a data dependence exists between different parts of a kernel

```
// Device code
```

```
__syncthreads();
```

C language extensions

- Execution synchronization on device
 - Handling a data dependence between different parts of a kernel

```
// Kernel code  
__shared__ float A[BLOCK_SIZE];  
  
// Initializing all threads in a block:  
A[threadIdx.x] = 1;  
  
// This is needed as the previous section is executed by sequential groups of  
// 32 (warp size) threads, in an undetermined order  
__syncthreads();  
  
// A primitive example of a data dependence:  
if (threadIdx.x == 0)  
    for (i=0; i<BLOCK_SIZE; i++)  
        sum = sum + A[i];
```

C language extensions

- Synchronization between host and device
 - Kernel calls and some CUDA memory operations are executed asynchronously on host
 - But in some cases you need host-device synchronization, e.g. before using a host timer, for profiling:

```
// Host code
```

```
CudaDeviceSynchronize ();
```

C language extensions

- Synchronization between host and device
 - Example: when profiling the code

// Host code

```
struct timeval tdr0, tdr1;
```

```
gettimeofday (&tdr0, NULL);
```

```
my_kernel <<<M, N>>> ();
```

*// Without synchronization, tdr1-tdr0 will not measure time spent inside the kernel
// (it will be much smaller):*

```
CudaDeviceSynchronize ();  
gettimeofday (&tdr1, NULL);
```

Quiz 2

Link: <http://www.socrative.com>

Room: CUDADAY2

C language extensions

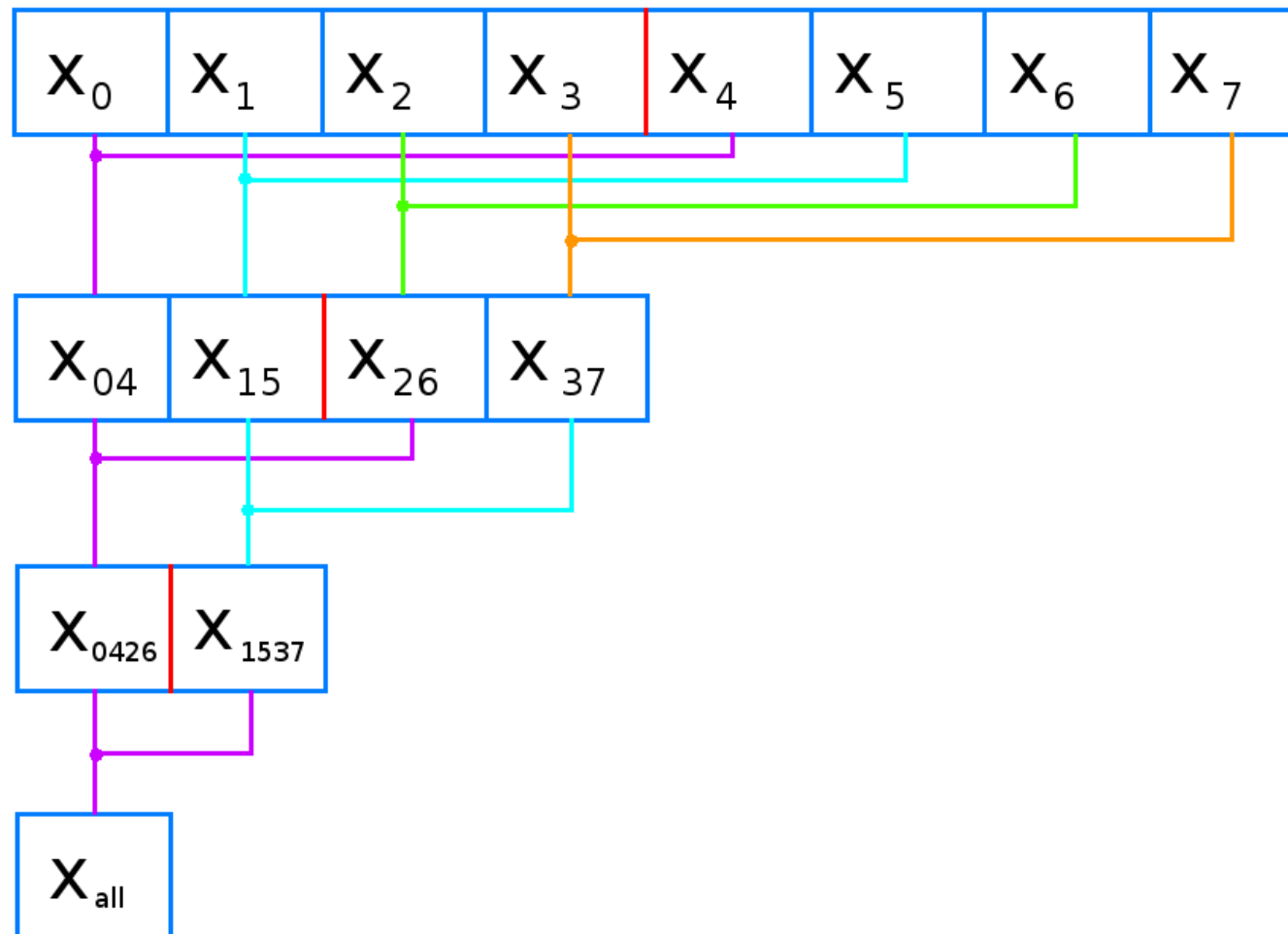
- Reductions in CUDA
 - Reductions: min/max, average, sum, ...
 - Can be a significant bottleneck for the performance, because it breaks pure data parallelism.
 - There is no perfect way to do reductions in CUDA. The two commonly used approaches (each with its own set of constraints) are
 - Binary reductions
 - Atomic reductions

C language extensions

- Binary reductions
 - The most universal type of reductions (e.g., the only way to do floating point min or max)
 - Even when using single precision (which is faster than double precision), binary summation will be more accurate than atomic summation, because it employs more accurate *pairwise summation*.
 - Usually the more efficient way to do reductions: time scales as $\log_2(N)$ **as long as there are enough of free GPU cores.**

C language extensions

Binary reduction



C language extensions

- Binary reductions
 - But: typically relies on (very limited) shared memory – placing constraints on how many reductions per kernel one can do
 - Relies on thread synchronization, which can only be done within a single block – places constraints on how many threads can participate in a binary reduction (usually 64 ... 256; maximum 1024)
 - For a large number of data elements (>1024), this leads to the need to do multi-level binary reductions, with storing the intermediate data in device memory; this can reduce the performance
 - Can be less efficient for small number of data elements (<64)
 - Significantly complicates the code

C language extensions

- Examples: binary summation with the number of elements being a power of two. The result is in sum[0].

```
__shared__ double sum[BLOCK_SIZE];  
...  
__syncthreads(); // To make sure all sum[] elements were initialized  
int nTotalThreads = blockDim.x; // Total number of active threads;  
// only the first half of the threads will be active.  
  
while(nTotalThreads > 1)  
{  
    int halfPoint = nTotalThreads / 2; // Number of active threads  
  
    if (threadIdx.x < halfPoint)  
    {  
        int thread2 = threadIdx.x + halfPoint; // the second element index  
        sum[threadIdx.x] += sum[thread2]; // Pairwise summation  
    }  
    __syncthreads();  
    nTotalThreads = halfPoint; // Reducing the binary tree size by two  
}  
  
// The result is in sum[0]
```

C language extensions

- Examples: binary min/max with the number of elements being a power of two.

```
__shared__ double min[BLOCK_SIZE];  
...  
__syncthreads(); // To make sure all min[] elements were initialized  
int nTotalThreads = blockDim.x;  
  
while(nTotalThreads > 1)  
{  
    int halfPoint = nTotalThreads / 2; // Number of active threads  
    if (threadIdx.x < halfPoint) {  
        int thread2 = threadIdx.x + halfPoint; // the second element index  
        double temp = min[thread2];  
        if (temp < min[threadIdx.x])  
            min[threadIdx.x] = temp;  
    }  
    __syncthreads();  
    nTotalThreads = halfPoint; // Reducing the binary tree size by two  
  
    // The result is in min[0]}
```

C language extensions

- Examples: multiple binary reductions.

```
__shared__ double min[BLOCK_SIZE], sum[BLOCK_SIZE];
...
__syncthreads(); // To make sure all array elements were initialized
int nTotalThreads = blockDim.x;

while(nTotalThreads > 1)
{
    int halfPoint = nTotalThreads / 2; // Number of active threads
    if (threadIdx.x < halfPoint) {
        int thread2 = threadIdx.x + halfPoint;
        sum[threadIdx.x] += sum[thread2]; // First reduction

        double temp = min[thread2];
        if (temp < min[threadIdx.x])
            min[threadIdx.x] = temp; // Second reduction
    }
    __syncthreads();
    nTotalThreads = halfPoint; // Reducing the binary tree size by two
}

// The result is in sum[0], min[0]
```

C language extensions

- Examples: two-level binary reduction

```
// Host code  
#define BSIZE 1024 // Always use a power of two; can be 32...1024  
// Total number of elements to process: 1024 < Ntotal < 1024^2  
  
int Nblocks = (Ntotal+BSIZE-1) / BSIZE;  
  
// Low level (the results should be stored in global device memory):  
x_prerreduce <<<Nblocks, BSIZE >>> ();  
  
// High level (will read the input from global device memory):  
x_reduce <<<1, Nblocks >>> ();
```

C language extensions

- Examples: binary reduction with an arbitrary number of elements (BLOCK_SIZE).

```
__shared__ double sum[BLOCK_SIZE];  
...  
__syncthreads(); // To make sure all sum[] elements were initialized  
int nTotalThreads = blockDim_2; // Total number of threads, rounded up to the next power of two  
  
while(nTotalThreads > 1)  
{  
    int halfPoint = nTotalThreads / 2; // Number of active threads  
  
    if (threadIdx.x < halfPoint)  
    {  
        int thread2 = threadIdx.x + halfPoint;  
        if (thread2 < blockDim.x) // Skipping the fictitious threads blockDim.x ... blockDim_2-1  
            sum[threadIdx.x] += sum[thread2]; // Pairwise summation  
    }  
    __syncthreads();  
    nTotalThreads = halfPoint; // Reducing the binary tree size by two  
}  
  
// The result is in sum[0]
```


C language extensions

- Continued: binary reduction with an arbitrary number of elements.
 - You will have to compute `blockDim_2` (`blockDim.x` rounded up to the next power of two), either on device or on host (and then copy it to device). One could use the following function to compute `blockDim_2`, valid for 32-bit integers:

```
int NearestPowerOf2 (int n)
{
    if (!n) return n; // (0 == 2^0)

    int x = 1;
    while(x < n)
    {
        x <<= 1;
    }
    return x;
}
```

Quiz 3

Link: <http://www.socrative.com>

Room: CUDADAY2

C language extensions

- Atomic reductions
 - Very simple and elegant code
 - Almost no change compared to the serial code
 - A single line code: much better for code development and maintenance
 - No need for multiple intermediate kernels (saves on overheads related to multiple kernel launches)
 - Requires no code changes when dealing with any number of data elements – from 2 to millions
 - Usually more efficient when the number of data elements is small (<64)

C language extensions

- Atomic reductions
 - But: atomic operations are serialized, which usually means **much** worse performance
 - No atomic functionality for some basic reductions (like floating point min / max)
 - A commonly employed good compromise is to use binary reduction at the lower level, and then use atomic reduction at the higher level.
 - All the above means that to find the right way to carry out a reduction in CUDA, with the right balance between code readability, efficiency, and accuracy, one often has to try different approaches, and choose the most efficient.

C language extensions

- Examples: atomic reductions.

```
// In global device memory:
```

```
__device__ float xsum;  
__device__ int isum, imax;
```

```
// In a kernel:
```

```
float x;  
int i;  
__shared__ imin;  
...  
atomicAdd (&xsum, x);  
atomicAdd (&isum, i);  
atomicMax (&imax, i);  
atomicMin (&imin, i);
```

- Some other atomic operations:
 - atomicExch, atomicAnd, atomicOr

C language extensions

- Binary at the lower level, atomic at the higher level

```
__shared__ float sum[BLOCK_SIZE];  
// Initialize sum[] array here  
__syncthreads(); // To make sure all sum[] elements were initialized  
int nTotalThreads = blockDim.x; // Total number of active threads;  
// only the first half of the threads will be active.  
  
while(nTotalThreads > 1){  
    int halfPoint = nTotalThreads / 2; // Number of active threads  
  
    if (threadIdx.x < halfPoint)  
    {  
        int thread2 = threadIdx.x + halfPoint; // the second element index  
        sum[threadIdx.x] += sum[thread2]; // Pairwise summation  
    }  
    __syncthreads();  
    nTotalThreads = halfPoint; // Reducing the binary tree size by two  
}  
if (threadIdx.x == 0)  
    atomicAdd (&xsum, sum[0]); // Atomic reduction
```

Hands on instructions (salloc)

```
$ ssh guest####@graham.computecanada.ca
```

```
$ salloc --time=0-01:00 --ntasks=1 -A cosse19-wa_gpu  
--mem=4G --reservation=cosse19-wr_gpu --gres=gpu:1
```

```
$ cp -r /home/syam/CUDA_day2 ~ (ignore read errors)
```

```
$ cd CUDA_day2
```

```
$ module load cuda
```

- Text editors: vim, emacs, nano . For proper nano syntax highlighting:

```
$ cp ~syam/.nanorc ~
```

Hands on instructions (sbatch)

```
$ ssh guest####@graham.computecanada.ca
$ cp -r /home/syam/CUDA_day2 ~ (ignore read errors)
$ cd CUDA_day2
$ module load cuda
```

- Text editors: vim, emacs, nano . For proper nano syntax highlighting:
\$ cp ~syam/.nanorc ~

```
$ nano job.sh
#SBATCH --reservation=cosse19-wr_gpu
#SBATCH -A cosse19-wa_gpu
#SBATCH --mem=4G
#SBATCH --time=0-00:10
#SBATCH --gres=gpu:1
#SBATCH --ntasks=1
./your_code
```

```
$ sbatch job.sh
```


Hands on exercise #1

- [CUDA_day2 / Reduction](#): implementing hybrid reduction scheme

C language extensions

- Concurrent execution and streams
 - Concurrency (parallel execution) between GPU and CPU is either a default, or easily enabled behaviour
 - Kernel launches are always asynchronous with regards to the host code; one has to use explicit device-host synchronization any time a kernel needs to be synchronized with the host:
`CudaDeviceSynchronize ()`
 - The default behaviour of GPU<->CPU memory copy operations is asynchronous for small transfers (<64kB; only host->device), and synchronous otherwise. But one can enforce any memory copying to be asynchronous by adding `Async` suffix, e.g.:
 - `cudaMemcpyAsync ()`
 - `cudaMemcpyToSymbolAsync ()`
 - For debugging purposes, one can enforce everything to be synchronous by setting the `CUDA_LAUNCH_BLOCKING` environment variable to 1.

C language extensions

- Concurrent execution and streams
 - Concurrency between different device operations (kernels and/or memory copying) is a completely different story
 - On a hardware level, modern GPUs are capable of running multiple kernels and memory transfers both to and from the device concurrently
 - By default, everything on device is done serially (no concurrency)
 - To make use of the device concurrency features, one has to start using multiple **streams** in the CUDA code
 - But even with multiple streams, there are some limitations to concurrency on GPU

C language extensions

- Concurrent execution and streams
 - A stream is a sequence of commands (possibly issued by different host threads) that execute in order
 - If stream ID is omitted, it is assumed to be “NULL” (default) stream. For non-default streams, the IDs have to be used explicitly.
 - For concurrent memory copying on GPU, one has to both add the Async suffix and specify the stream ID.

```
mykernel <<<Nblocks, Nthreads, 0, ID>>> ();  
cudaMemcpyAsync(d_A, h_A, size, cudaMemcpyHostToDevice, ID);
```

C language extensions

- Concurrent execution and streams
 - Before using, streams have to be created. At the end, they have to be destroyed

```
// Host code
cudaStream_t ID[2];

// Creating streams:
for (int i = 0; i < 2; ++i)
    cudaStreamCreate (&ID[i]);

// These two commands will run concurrently on GPU:
mykernel <<<Nblocks, Nthreads, 0, ID[0]>>> ();
cudaMemcpyAsync (d_A, h_A, size, cudaMemcpyHostToDevice, ID[1]);

// Destroying streams:
for (int i = 0; i < 2; ++i)
    cudaStreamDestroy (ID[i]);
```

C language extensions

- Concurrent execution and streams
 - Limitations:
 - For memory copying operations to run concurrently with any other device operation (kernel or another memory copying operation), the host memory has to be *page-locked* (or *pinned*; allocated with [cudaMallocHost](#) instead of `malloc`; static variables can be made pinned using [cudaHostRegister](#))
 - Up to 128 kernels can run concurrently (P100)
 - Concurrency on GPU is not guaranteed (e.g., if kernels use too much local resources, they will not run concurrently)

C language extensions

- Concurrent execution and streams
 - Other stream-related commands
 - `cudaDeviceSynchronize()` : global synchronization (across all the streams and the host);
 - `cudaStreamSynchronize (ID)` : synchronize stream ID with the host;
 - `cudaStreamQuery (ID)` : tests if the stream ID has finished running.

Quiz 4

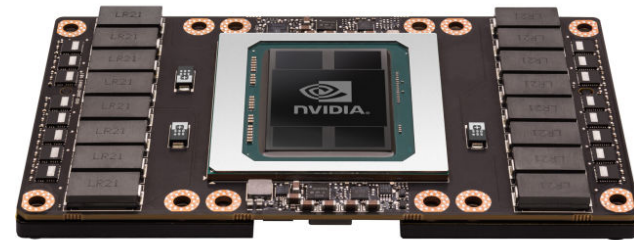
Link: <http://www.socrative.com>

Room: CUDADAY2

Graham GPUs, P100

From Fermi to Pascal

- Important GPU generations (https://en.wikipedia.org/wiki/Nvidia_Tesla):
 - Fermi: 2010
 - Kepler: 2012
 - [Maxwell: 2014]
 - **Pascal: 2016**
 - Volta: 2017
 - Turing: 2018
- (Maxwell didn't have any HPC GPUs.)
- Our cluster Graham has 320 of HPC Pascal GPUs, P100.
(Cedar has 584 P100's; Beluga has 688 Volta's)



Evolutionary changes

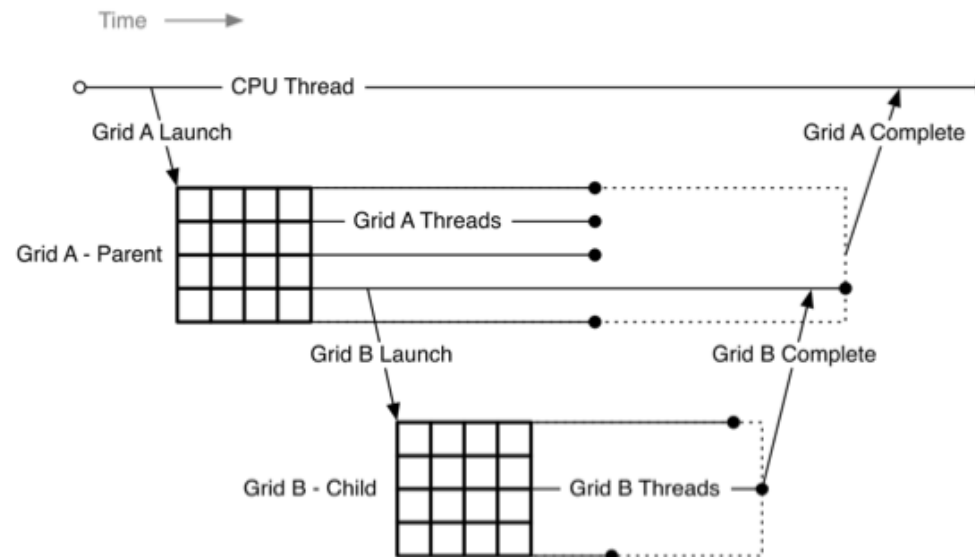
<i>Specification</i>	<i>Monk</i>	<i>Graham</i>
CUDA cores:	448	3584
SP flops:	1.03 TFlops	9.3 TFlops
Device memory:	5.2 GB	12 GB
Memory bandwidth:	148 GB/s	549 GB/s

Revolutionary changes

- **CUDA Dynamic Parallelism** (CDP): new hard/software feature allowing for dynamic workload generation on GPU (kernels launched from kernels). Makes GPU much more general purpose computing device. First appeared in Kepler GPUs.
- **Hyper-Q**: in previous generations, multiple CPU threads could only access the GPU sequentially (one queue); Kepler / Pascal expand that to 32 parallel queues. This should significantly accelerate mixed MPI/CUDA and OpenMP/CUDA codes, without any code modifications. Also great for GPU farming.

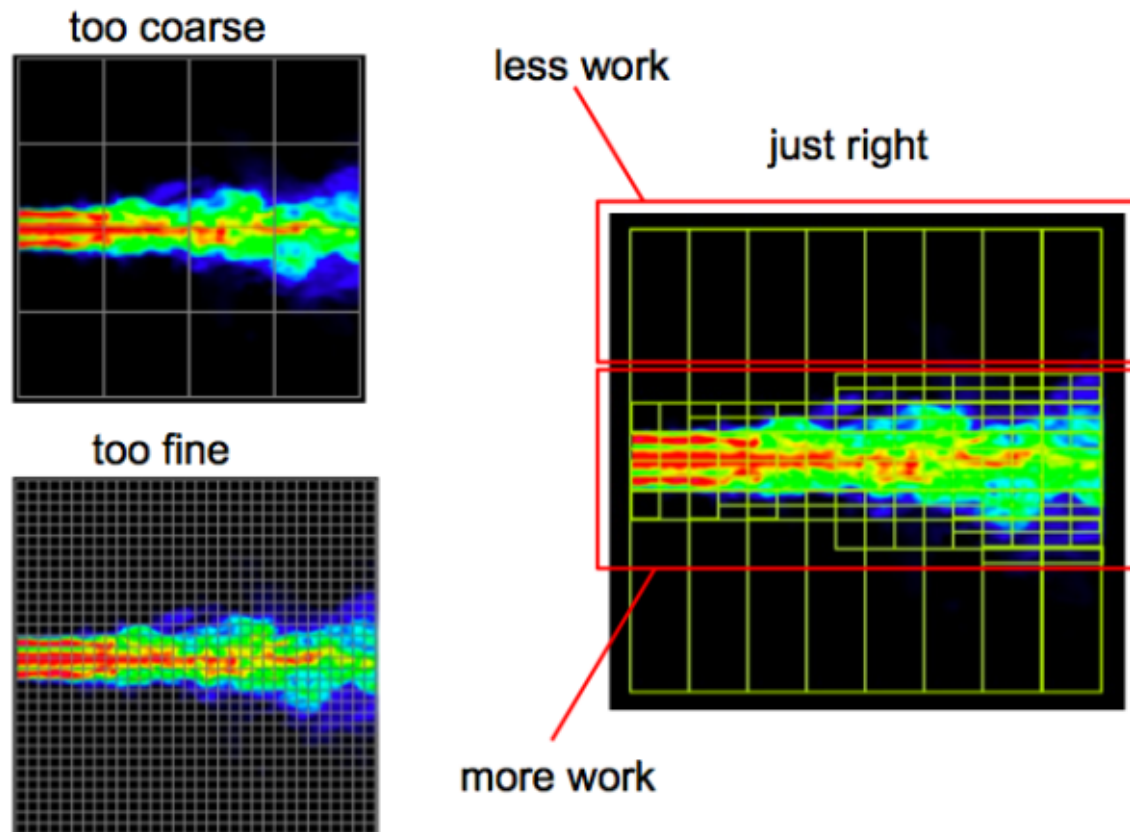
Dynamic Parallelism

- Dynamic parallelism (DP) is available in CUDA 5.0 and later on devices of Compute Capability 3.5 or higher (sm_35 for Kepler; sm_60 for Pascal).
- Under DP, an application can launch a coarse-grained kernel which in turn launches finer-grained kernels to do work where needed.



Dynamic Parallelism

- DP is perfect for adaptive grid codes and codes with recursion.



DP: simple example

- DP allows one to move almost everything to GPU.

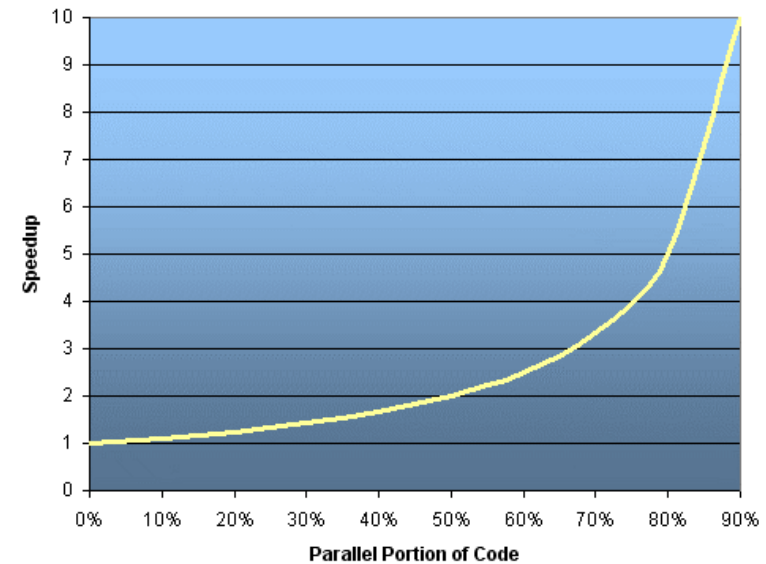
```
// On device:  
// Second level kernels (multi-threaded):  
__global__ void kernel1 (){}  
__global__ void kernel2 (){}  
  
// Top level kernel (single-threaded):  
__global__ void main_kernel (){  
    if (threadIdx.x == 0) {  
// These second level kernels will run sequentially (would need streams for concurrency)  
        kernel1<<<Nblocks, Nthreads>>>();  
        kernel2<<<Nblocks, Nthreads>>>();  
  
        ..  
    }  
}  
// On host:  
int main() {  
    main_kernel<<<1,1>>>();  
}
```

Amdahl's Law

- **Amdahl's Law** states that potential program speedup is defined by the fraction of code (P) that can be parallelized:

$$\text{speedup} = \frac{1}{1 - P}$$

- If none of the code can be parallelized, = 1 (no speedup). If all of the code is parallelized, the speedup is infinite (in theory).
- If 50% of the code can be parallelized, maximum speedup = 2, meaning the code will run twice as fast.



Amdahl's Law (2)

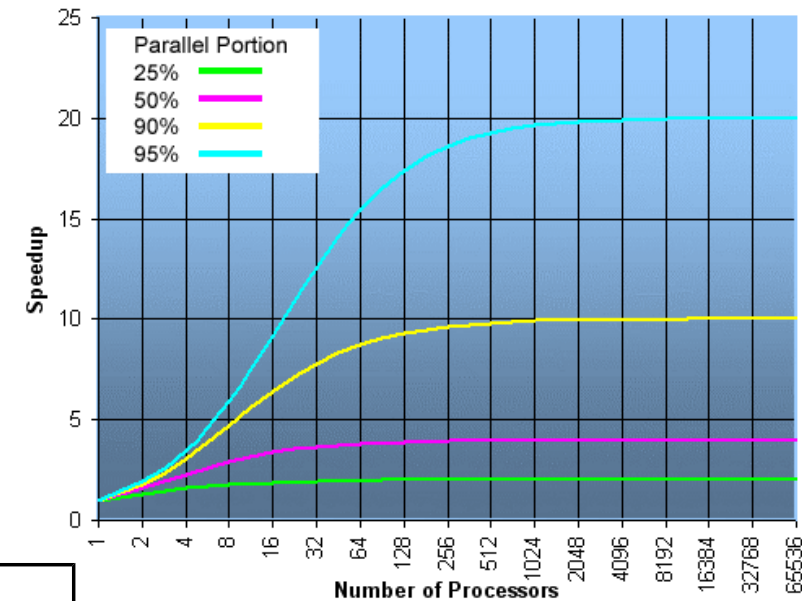
- Introducing the number of processors performing the parallel fraction of work, the relationship can be modeled by:

$$\text{speedup} = \frac{1}{\frac{P}{N} + S}$$

where P = parallel fraction, N = number of processors and S = serial fraction.

Amdahl's Law (3)

- It soon becomes obvious that there are limits to the scalability of parallelism. For example, at $P = .50$, $.90$ and $.99$ (50%, 90% and 99% of the code is parallelizable):



N	speedup		
	P = .50	P = .90	P = .99
10	1.82	5.26	9.17
100	1.98	9.17	50.25
1000	1.99	9.91	90.99
10000	1.99	9.91	99.02

Hyper-Q: why is it important?

- GPUs work well when you saturate them with data-parallel threads.
- Graham GPU has 8 times more cores (so need 8x more threads to get saturated) than the Monk GPU.
- From the Amdahl's law, a code which runs well on Monk will likely perform poorly on Graham.
- Hyper-Q helps to mitigate this, by allowing to share one GPU between different CPU threads.

Live demo of Hyper-Q

- A simple code, `primes_HQ`, only runs one block of threads per kernel.
- This mimics a realistic code which doesn't have enough of parallelism to saturate a modern GPU.
- Important: Hyper-Q is usually not enabled by default.

Other new features

- Atomic operations improvements:
 - atomicAdd now supports FP64 (integer and float)
 - atomicMin and atomicMax now support INT64
- Half precision (FP16) at twice speed of FP32
- Quantitative improvements:
 - Grid length (1D): 65,535 -> 2e9
 - 32-bit registers per thread: 63 -> 255
 - Concurrent kernels per device: 16 -> 128

Quiz 5

Link: <http://www.socrative.com>

Room: CUDADAY2

CUDA code optimization

CUDA code optimization

- Converting a code to CUDA can be considered an advanced exercise in code optimization
 - You should start profiling CUDA code from the very beginning, from the first kernel you write
 - You should start the conversion from the most cpu-intensive parts of the code
 - You often have to play with different approaches until you get the best performance in a given part of the code
- We will consider a few common optimization strategies

CUDA code optimization

- Kernels: how many?
 - You have to start a new kernel every time there is a global (across multiple blocks) data dependence
 - Example: two-level binary reduction shown previously
 - Another example: you need a separate kernel to initialize variables used to store an atomic reduction result:

```
// In global device memory:  
__device__ double d_sum;
```

```
// On host:  
// Initializing d_sum to zero:  
init_sum <<<1, 1>>> ();
```

```
// Here d_sum is used to store atomic summation result from multiple blocks  
compute_sum <<<Nblocks, BSIZE>>> ();
```

CUDA code optimization

- Kernels: how many?
 - You can try to split a kernel if it uses too many registers (*register pressure*)
 - It happens e.g. if the kernel has many complex algebraic expressions using lots of parameters
 - Register pressure can be identified when profiling the code with NVIDIA CUDA profilers (e.g., it will manifest itself via low *occupancy number*)
 - It is very non-intuitive: sometimes the register pressure can be decreased by making the kernel *longer* (presumably, because sometimes adding more lines of code gives CUDA compiler more flexibility to re-arrange register usage across the kernel)

CUDA code optimization

- Kernels: how many?
 - You should end a kernel when there is a device-host dependence
 - Example:

// On host:

```
kernel1 <<<N, M>>> (d_A);  
cudaMemcpy (h_A, d_A, size, cudaMemcpyDeviceToHost);
```

// Host code dependent on kernel1 results:

```
library_function1 ();
```

CUDA code optimization

- Kernels: how many?
 - Otherwise, you should try to make kernels as large as possible
 - Because each kernel launch has an overhead, in part because one has to store and then read the intermediate results from a slow (device or host) memory
 - You shouldn't worry that the kernel code won't fit on GPU: modern GPUs have large a limit of 512 million instructions per kernel
 - To improve readability, parts of the kernel can be modularized into device functions

CUDA code optimization

- What should be computed on GPU?
 - You start with the obvious targets: cpu-intensive data-parallel parts of the code
 - What should you do with the leftover code (not data-parallel and/or not very cpu intensive)?
 - If not a lot of data needs to be copied from device to host and vice versa for the leftover code, it may be beneficial to leave these parts of the code on host
 - If on the other hand the leftover code needs an access to a lot of intermediate results from CUDA kernels, then it may be more efficient to move everything to the GPU – even purely serial (single-thread) computations. This way, no intermediate (scratch) data will ever need to leave GPU.

CUDA code optimization

- Moving leftover code to GPU

// On host:

// First chunk of data-parallel code goes here:

```
kernel1 <<<N, M>>> ();
```

// Copying kernel1 results to host:

```
cudaMemcpy (h_A, d_A, size,  
            cudaMemcpyDeviceToHost);
```

// Non-parallelizable part of the code:

```
serial_computation (h_A, h_B);
```

// Copying serial_computation results to device:

```
cudaMemcpy (d_B, h_B, size,  
            cudaMemcpyHostToDevice);
```

// CudaDeviceSynchronize (); - Why?

// Second chunk of data parallel code which depends on d_B:

```
kernel2 <<<N, M>>>
```

// On host:

// First chunk of data-parallel code goes here:

```
kernel1 <<<N, M>>> ();
```

// Now it is executed on GPU, serially:

```
serial_computation_kernel <<<1, 1>>> ();
```

// Second chunk of data parallel code which depends on d_B:

```
kernel2 <<<N, M>>>
```

Quiz 6

Link: <http://www.socrative.com>

Room: CUDADAY2

CUDA code optimization

- Optimizing memory copying between GPU and CPU
 - GPU - device memory bandwidth is much (~20x) larger than GPU - host memory bandwidth
 - As a result, minimizing amount of data copied between GPU and CPU should be a high priority
 - One possible solution is described on the previous slide: move to GPU the “leftover” code (even if it poorly performs on GPU) if it helps to cut significantly on GPU-CPU memory copying

CUDA code optimization

- Optimizing memory copying between GPU and CPU
 - Sometimes one can reduce or eliminate the time spent on GPU-CPU data copying if it is done in parallel (asynchronously) with **host** computations:

// On host:

// This memory copying will be asynchronous only in regards to the host code:
`cudaMemcpyAsync(d_a, h_a, size, cudaMemcpyHostToDevice, 0);`

// This host code will be executed in parallel with memory copying
`host_computation ();`

CUDA code optimization

- Optimizing memory copying between GPU and CPU
 - One can also run memory transfer operation concurrently with another (opposite direction) memory transfer operation, or a kernel. For that, one has to create and use streams.
 - Only works with pinned host memory

```
// This memory copying will be asynchronous in regards to the host and stream ID[1]:  
cudaMemcpyAsync(d_a, h_a, size, cudaMemcpyHostToDevice, ID[0]);
```

```
// The kernel doesn't need d_a, and will run concurrently with the previous line:  
kernel1 <<<N, M, 0, ID[1]>>> ();
```

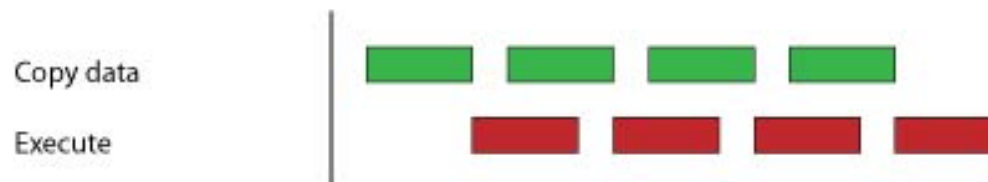
CUDA code optimization

- Optimizing memory copying between GPU and CPU
 - Staged concurrent copy and execute

One stream scenario:



You need two streams for this:



Hands on exercise #2

- [CUDA_day2 / Staged](#): using streams to stage copying and computing

CUDA code optimization

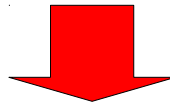
- Optimizing memory copying between GPU and CPU
 - To save on memory copying overheads, one should try to bundle up multiple small transfers into one large one
 - This can be conveniently achieved by creating a single structure, with the individual memory copying arguments becoming elements of the structure

CUDA code optimization

- Optimizing memory copying between GPU and CPU

// Host code:

```
cudaMemcpyToSymbol (d_A, &h_A, sizeof(h_A), 0, cudaMemcpyHostToDevice);  
cudaMemcpyToSymbol (d_B, &h_B, sizeof(h_B), 0, cudaMemcpyHostToDevice);  
cudaMemcpyToSymbol (d_C, &h_C, sizeof(h_C), 0, cudaMemcpyHostToDevice);
```



// Header file:

```
struct my_struct {  
    double A[1000];  
    double B[2000];  
    int C[1000];  
};  
__device__ struct my_struct d_struct;  
struct my_struct h_struct;
```

// Host code:

```
cudaMemcpyToSymbol (d_struct, &h_struct, sizeof(h_struct), 0, cudaMemcpyHostToDevice);
```

CUDA code optimization

- Optimizing memory copying between GPU and CPU
 - If you use dynamic memory allocation on host, you can usually accelerate copying to/from the device by using `cudaMallocHost` instead of `malloc`.
 - This will force the compiler to use page-locked memory for host allocations, which has much higher bandwidth to the device
 - Use this sparingly, as the performance can actually degrade when not enough of system memory is available for paging

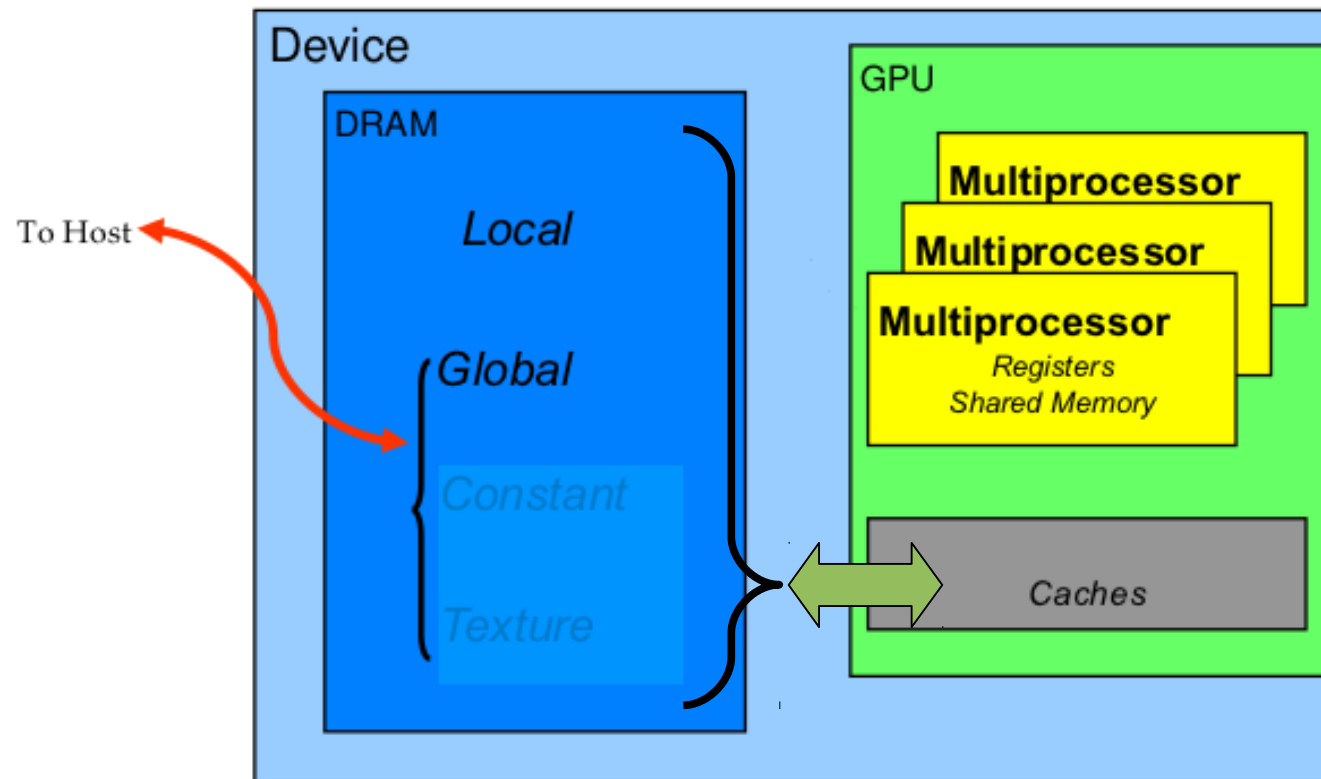
// Host code:

```
float *h_A;
```

```
cudaMallocHost (&h_A, N*sizeof(float));
```

CUDA code optimization

- Optimizing memory access on GPU
 - Memory spaces on GPU

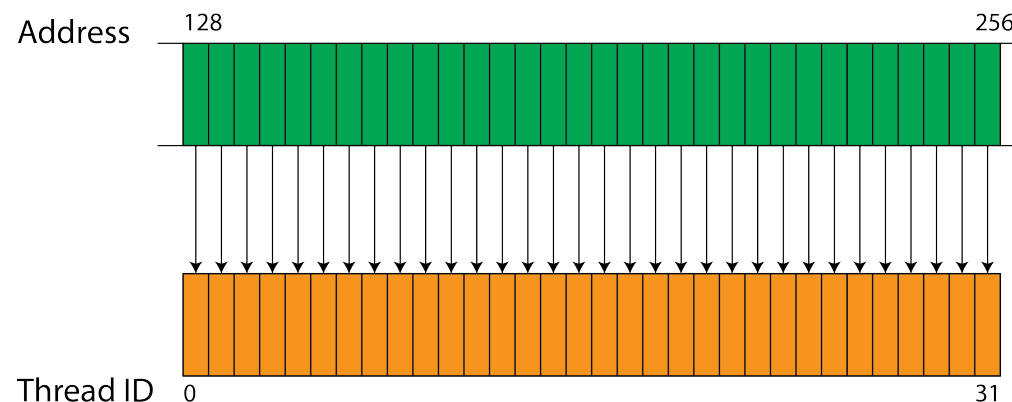


CUDA code optimization

- Optimizing memory access on GPU
 - Registers \leftrightarrow “Local” memory are not under your direct control, making it harder to optimize
 - Global and shared memory, on the other hand, are under direct programmer's control, so they are easier to optimize.
 - Main strategies for optimization:
 - Global memory: coalescence of memory accesses
 - Shared memory: minimizing bank conflicts

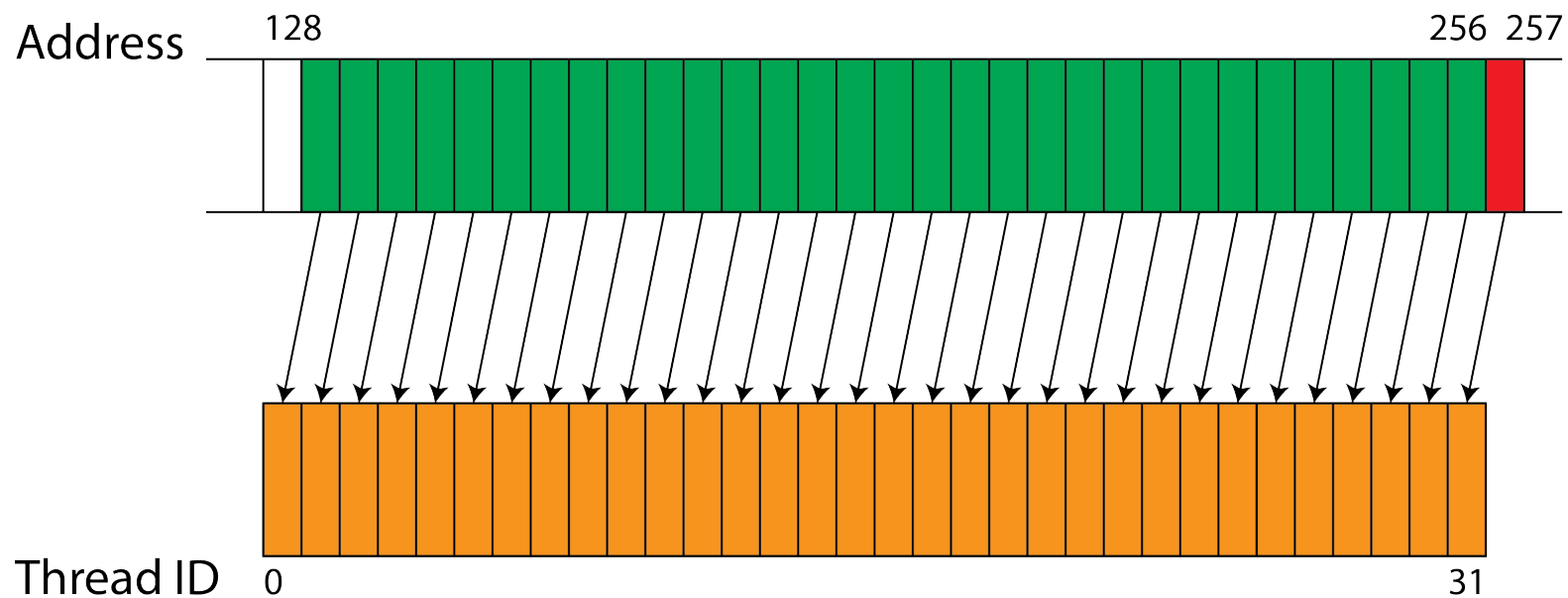
CUDA code optimization

- Optimizing memory access on GPU
 - Global memory: coalescence of memory accesses
 - Global memory loads and stores by threads of a warp are coalesced by the device into as few as one transaction when certain access requirements are met
 - By default, all accesses are cached through L1 as 128-byte lines
 - Coalescence is the best when accessing flat arrays (unit stride) consecutively.



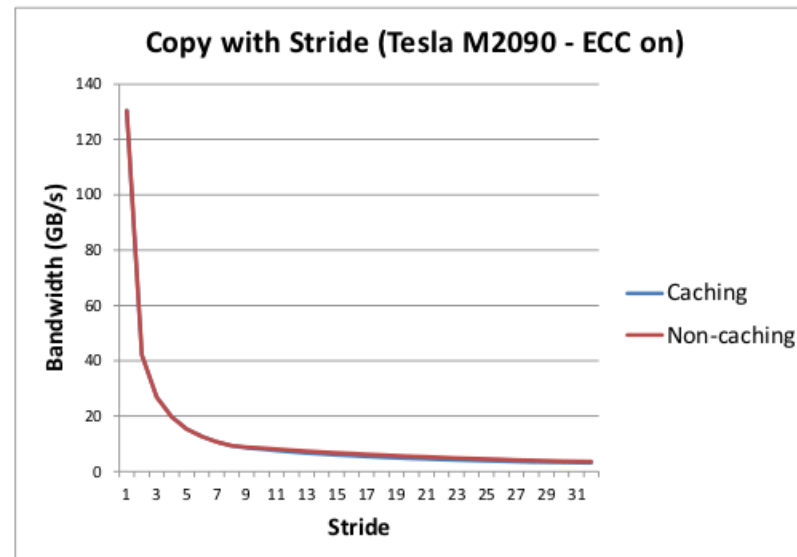
CUDA code optimization

- Optimizing memory access on GPU
 - Global memory: coalescence of memory accesses
 - Misaligned access degrades the performance, but not dramatically



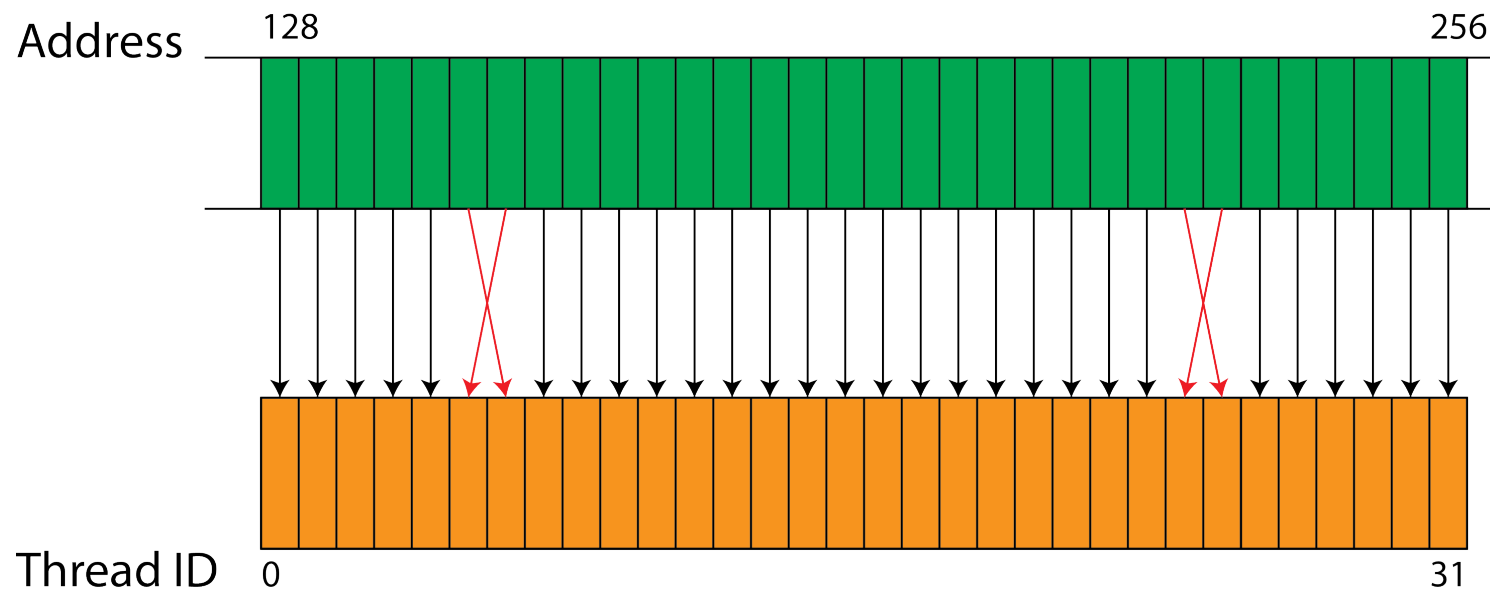
CUDA code optimization

- Optimizing memory access on GPU
 - Global memory: coalescence of memory accesses
 - Non-unit stride access (e.g. multi-D arrays), on the other hand, degrades the performance very rapidly, as the stride increases from 2 to 32:



CUDA code optimization

- Optimizing memory access on GPU
 - Global memory: coalescence of memory accesses
 - Any kind of non-sequential memory access (not just $\text{stride} > 1$) is bad*



CUDA code optimization

- Optimizing memory access on GPU
 - Global memory: coalescence of memory accesses
 - The strategy with multi-D arrays is to either
 - flatten them yourself (the only way if >3 dimensions), or
 - use special CUDA functions `cudaMallocPitch()` and `cudaMalloc3D()` to allocate properly aligned 2D and 3D arrays, respectively, or
 - at the very least, convert row-major arrays to **column-major** ones

CUDA code optimization

- Optimizing memory access on GPU

```
// Using cudaMallocPitch, 2D case
```

```
// Host code
```

```
int width = 64, height = 64;  
float* devPtr;  
size_t pitch;
```

```
cudaMallocPitch (&devPtr, &pitch, width * sizeof(float), height);  
MyKernel <<<64, 64>>> (devPtr, pitch);
```

```
// Device code
```

```
__global__ void MyKernel (float* devPtr, size_t pitch)  
{  
    int ix = blockIdx.x;  
    int iy = threadIdx.x;  
    float* row = (float*)((char*)devPtr + ix * pitch);  
    float element = row[iy]; // Coalesced access  
}
```

CUDA code optimization

- Optimizing memory access on GPU

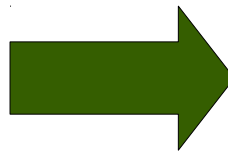
```
// Flattened, good for any D; individual dimensions can be arbitrary
// On device:
#define N_TOTAL N1*N2*N3*N4
__device__ float d_A[N_TOTAL];
__global__ void mykernel () {
    int i = threadIdx.x + blockDim.x * blockIdx.x;
    if (i < N_TOTAL) {
        d_A[i] = ...
        // You compute individual indexes only if they are needed for the computations:
        int i1 = i % N1; int m = i / N1;
        int i2 = m % N2; m = m / N2;
        int i3 = m % N3;
        int i4 = m / N3;
    }
}
// On host:
int Nblocks = (N_TOTAL + BLOCK_SIZE - 1) / BLOCK_SIZE;
mykernel <<<Nblocks, BLOCK_SIZE>>> ();
```


CUDA code optimization

- Optimizing memory access on GPU
 - Global memory: coalescence of memory accesses
 - If you have to use non-flattened static multi-D arrays, transpose them to “column-major” if they are “row-major”:

// Row-major (non coalesced)

```
float A[N][30];  
...  
A[threadIdx.x][0]=...;  
A[threadIdx.x][1]=...;
```



// Column-major (coalesced)

```
float A[30][N];  
...  
A[0][threadIdx.x]=...;  
A[1][threadIdx.x]=...;
```

CUDA code optimization

- Optimizing memory access on GPU
 - Global memory: coalescence of memory accesses
 - For the same reason, use **structures of arrays** instead of **arrays of structures** (the latter results in a memory access with a large stride)

// Array of structures behaves like row major accesses (non coalesced)

```
struct Point { double x; double y; double z; double w; } A[N];
```

```
...  
A[threadIdx.x].x = ...
```



// Structure of arrays behaves like column major accesses (coalesced)

```
struct PointList { double *x; double *y; double *z; double *w; } A;
```

```
...  
A.x[threadIdx.x] = ...
```

Quiz 7

Link: <http://www.socrative.com>

Room: CUDADAY2

CUDA code optimization

- Optimizing memory access on GPU
 - Using shared memory to optimize access to global memory
 - Shared memory is much faster than global memory; also, access to shared memory doesn't need to be coalesced
 - Shared memory can be viewed as a “user-managed cache for global memory”
 - One can store in shared memory frequently used global data
 - One can use shared memory to make reading data from global memory coalesced

CUDA code optimization

- Optimizing memory access on GPU

// Straightforward and inefficient way

```
__global__ void simpleMultiply(float *a, float* b, float *c, int
N)
{
    int row = blockIdx.y * blockDim.y + threadIdx.y;
    int col = blockIdx.x * blockDim.x + threadIdx.x;
    float sum = 0.0f;
    for (int i = 0; i < TILE_DIM; i++) {
        sum += a[row*TILE_DIM+i] * b[i*N+col];
    }
    c[row*N+col] = sum;
}
```

*// Using shared memory to both store frequently used global
// data and to make the access coalesced – 2.3x faster on K20*

```
__global__ void sharedABMultiply(float *a, float* b, float *c, int
N)
{
    __shared__ float aTile[TILE_DIM][TILE_DIM],
                    bTile[TILE_DIM][TILE_DIM];

    int row = blockIdx.y * blockDim.y + threadIdx.y;
    int col = blockIdx.x * blockDim.x + threadIdx.x;
    float sum = 0.0f;
    aTile[threadIdx.y][threadIdx.x] =
        a[row*TILE_DIM+threadIdx.x];
    bTile[threadIdx.y][threadIdx.x] = b[threadIdx.y*N+col];
    __syncthreads();
    for (int i = 0; i < TILE_DIM; i++) {
        sum += aTile[threadIdx.y][i] * bTile[i][threadIdx.x];
    }
    c[row*N+col] = sum;
}
```

CUDA code optimization

- Optimizing memory access on GPU
 - Shared memory: minimizing bank conflicts
 - Shared memory has 32 banks that are organized such that successive 32-bit words are assigned to successive banks
 - A bank conflict only occurs if two or more threads access the same bank

// No bank conflicts for 32-bit data is when the stride is odd ($s = 1, 3, \dots$)

```
__shared__ float shared[BLOCK_SIZE];  
float data = shared[BaseIndex + s * threadIdx.x];
```

// No bank conflicts for 64-bit data:

```
__shared__ double shared[BLOCK_SIZE];  
double data = shared[BaseIndex + threadIdx.x];
```

CUDA code optimization

- Minimizing warp divergence
 - The smallest independent execution unit in CUDA is a warp (a group of 32 consecutive threads in a block)
 - Within a warp, execution is synchronous (that is, warp acts as a 32-way vector processor)
 - Any flow control instruction (if, switch, do, for, while) acting on individual threads within a warp will result in **warp divergence** (with the different execution paths serialized), resulting in poor performance
 - Warp divergence minimization is hence an important CUDA optimization step

CUDA code optimization

- Minimizing warp divergence
 - Ideally, controlling conditions should be identical within a warp:

```
// On device:
__global__ void MyKernel ()
{
    int i = threadIdx.x + blockDim.x * blockIdx.x;
    int warp_index = i / warpSize; // Remains constant within a warp

    if (d_A[warp_index] == 0) // Identical execution path within a warp (no divergence)
        do_one_thing (i);
    else
        do_another_thing (i);
}
```


CUDA code optimization

- Minimizing warp divergence
 - As warps can't span thread blocks, conditions which are only a function of block indexes result in non-divergent warps

// On device:

```
__global__ void MyKernel ()  
{  
    int i = threadIdx.x + blockDim.x * blockIdx.x;  
  
    if (d_A[blockIdx.x] == 0) // No divergence, since warps can't span thread blocks  
        do_one_thing (i);  
    else  
        do_another_thing (i);  
}
```

CUDA code optimization

- Minimizing warp divergence
 - More generally, making a condition to span at least a few consecutive warps results in acceptably low level of warp divergences (even when the condition is not always aligned with warp boundaries)

```
// On device:
__global__ void MyKernel ()
{
    int i = threadIdx.x + blockDim.x * blockIdx.x;
    int cond_index = i / N_CONDITION; // Is okay if N_CONDITION >~ 5*warpSize

    if (d_A[cond_index] == 0) // Only a fraction of warps will have divergences
        do_one_thing (i);
    else
        do_another_thing (i);
}
```

CUDA code optimization

- Accuracy versus speed
 - Situation with double precision speed in CUDA improved dramatically in the recent years, but it is still slower than single precision
 - The ratio was 1:8 for capability 1.3 (old cluster angel)
 - The newer cluster monk (capability 2.0) has the ratio 1:2
 - But newest NVIDIA GPUs seem to be moving in the “wrong” direction (1:3 for K20; 1:32 for Maxwell; **back to 1:2 for Pascal**)
 - Use double precision only where it is absolutely necessary

CUDA code optimization

- Optimal CUDA parameters
 - **Number of threads per block** (BLOCK_SIZE): total range 1...1024; much better if multiples of 32; better still if multiples of 64.
 - **Number of threads per multiprocessor**: at least 768 for capability 2.x to completely hide read-after-write register latency. That means at least 10,752 threads per kernel for the whole monk GPU. (And probably >43,000 for P100.)
 - **Number of blocks in a kernel**: at least equal to the number of multiprocessors (≥ 56 for P100), to keep all multiprocessors busy.

Hands on exercise #3

- [CUDA_day2 / Primes](#): converting a serial code for the largest prime number search to CUDA