#### Setup

ssh to DAVINCI and grab a GPU node (if you don't already have one):

```
$ source /projects/k2i/hpc/scripts/login-gpu.sh
```

Once you have a GPU node, load the necessary packages and clone today's Github repo:

```
$ source /projects/k2i/hpc/scripts/gpu-day-one-setup.sh
```

\$ git clone git@github.com:agrippa/hpc-bootcamp.git \$HOME/bootcamp-gpu

It also may be useful to have the CUDA API references open:

http://docs.nvidia.com/cuda/cuda-runtime-api/index.html

Slides are accessible in your browser in Github and Box:

Github: <a href="https://github.com/agrippa/hpc-bootcamp">https://github.com/agrippa/hpc-bootcamp</a>

# **GPU Accelerated Computing (Day 1)**Introduction to Programming GPUs

#### **Max Grossman**

Habanero Extreme Scale Software Research Group, Rice University Principal & Co-Founder, 7pod Technologies Author, Professional CUDA C Programming

#### **Outline**

- 1. A Brief History of Computing
- 2. GPU Architecture Overview
- 3. An Overview of CUDA
- 4. Hands-On with a Scientific App
- 5. Related Topics:
  - 1. Alternative GPU Programming Models
  - 2. Alternatives to GPUs

#### **About Me**

#### Habanero Research Group

- Runtime Systems for HPC ("Big Compute")
- Data Analytics ("Big Data")
- Compilers and Programmer Tools

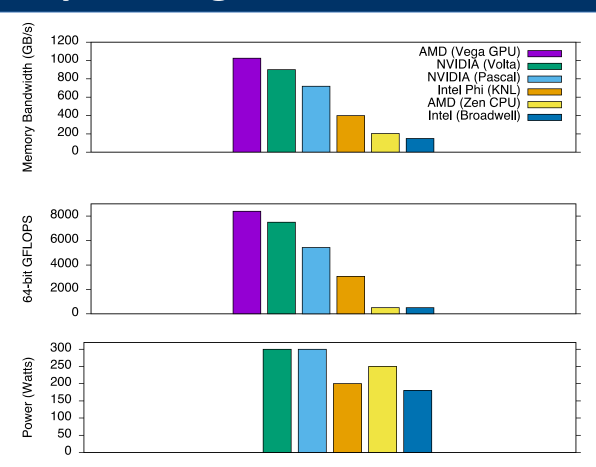
#### Co-Founder, 7pod Technologies (7pod.tech)

- Consulting on new applications of data analytics and high performance computing
- HPC and data analytics training



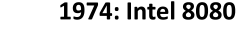
# Why GPUs?

#### **Upcoming Hardware Generations**



Is your hardware holding you back, or are you holding back your hardware?

1971: Intel 4004









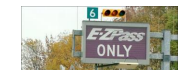
#### What do all of these applications have in common?







What do all of these applications have in common?

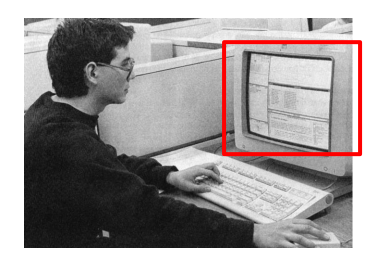


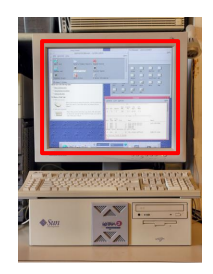
Application performance is entirely dependent on straight-line performance.





First dedicated Graphics Processing Units (GPUs) introduced in the 1990s.





#### Only became programmable in the early 2000s.

**Game**Developers

#### Standard "Grid" Computation

Initialize "view" (pixels:texels::1:1)

```
glMatrixMode(GL_MODELVIEW);
glLoadIdentity();
glMatrixMode(GL_PROJECTION);
glLoadIdentity();
glOrtho(0, 1, 0, 1, 0, 1);
glViewport(0, 0, gridResX, gridResY);
```

- For each algorithm step:
  - Activate render-to-texture
  - Setup input textures, fragment program
  - Draw a full-screen quad (1 unit x 1 unit)



NVIDIA





**GPU:** high performance growth



- Annual growth ~1.5x → decade growth ~ 60x
- Moore's law
- GPU
  - Annual growth > 2.0x → decade growth > 1000x
  - Much faster than Moore's law





NVIDIA



1

Tianhe-2 (MilkyWay-2) - TH-IVB-FEP Cluster, Intel Xeon E5-2692 12C 2.200GHz, TH Express-2, Intel Xeon Phi 31S1P

NUDT

#### 2

Titan - Cray XK7 , Opteron 6274 16C 2.200GHz, Cray Gemini interconnect, NVIDIA K20x

Cray Inc.

#### 3

**Sequoia** - BlueGene/Q, Power BQC 16C 1.60 GHz, Custom

#### 4

K computer, SPARC64 VIIIfx 2.0GHz, Tofu interconnect Fujitsu

#### 5

Mira - BlueGene/Q, Power BQC 16C 1.60GHz, Custom

6

Trinity - Cray XC40, Xeon E5-2698v3 16C 2.3GHz, Aries interconnect Cray Inc.

-, --,

Piz Daint - Cray XC30, Xeon E5-2670 8C 2.600GHz, Aries interconnect , NVIDIA K20x Cray Inc.

#### And a decade later...

Green500 Rank	MFLOPS/W	Site*	Computer*	Total Power (kW)
1	7,031.58	Institute of Physical and Chemical Research (RIKEN)	Shoubu - ExaScaler-1.4 80Brick, Xeon E5-2618Lv3 8C 2.3GHz, Infiniband FDR, PEZY-SC	50.32
2	5,331.79	GSIC Center, Tokyo Institute of Technology	TSUBAME-KFC/DL - LX 1U-4GPU/104Re-1G Cluster, Intel Xeon E5-2620v2 6C 2.1GHz, Infiniband FDR, NVIDIA Tesla K80	51.13
3	5,271.81	GSI Helmholtz Center	ASUS ESC4000 FDR/G2S, Intel Xeon E5-2690v2 10C 3GHz, Infiniband FDR, AMD FirePro S9150	57.15
4	4,778.46	Institute of Modern Physics (IMP), Chinese Academy of Sciences	Sugon Cluster W780I, Xeon E5-2640v3 8C 2.6GHz, Infiniband QDR, NVIDIA Tesla K80	65.00
5	4,112.11	Stanford Research Computing Center	XStream - Cray CS-Storm, Intel Xeon E5-2680v2 10C 2.8GHz, Infiniband FDR, Nvidia K80	190.00
6	3,856.90	IT Company	Inspur TS10000 HPC Server, Xeon E5-2620v3 6C 2.4GHz, 10G Ethernet, NVIDIA Tesla K40	58.00
7	3,775.45	Internet Service	Inspur TS10000 HPC Server, Intel Xeon E5-2620v2 6C 2.1GHz, 10G Ethernet, NVIDIA Tesla K40	110.00
8	3,775.45	Internet Service	Inspur TS10000 HPC Server, Intel Xeon E5-2620v2 6C 2.1GHz, 10G Ethernet, NVIDIA Tesla K40	110.00
9	3,775.45	Internet Service	Inspur TS10000 HPC Server, Intel Xeon E5-2620v2 6C 2.1GHz, 10G Ethernet, NVIDIA Tesla K40	110.00
10	3,775.45	Internet Service	Inspur TS10000 HPC Server, Intel Xeon E5-2620v2 6C 2.1GHz, 10G Ethernet, NVIDIA Tesla K40	110.00

TITAN	SUMMIT
18,688	~3,400
(1) 16-core AMD Opteron per node	(Multiple) IBM POWER 9s per node
(1) NVIDIA Kepler K20x per node	(Multiple) NVIDIA Volta GPUs per node
32GB (DDR3)	>512GB (HBM+DDR4)
PCI Gen2	NVLINK (5-12x PCle3)
Gemini	Dual Rail EDR-IB (23 GB/s)
9 MW	10 MW

Rank	System	Cores	(TFlop/s)	(TFlop/s)	(kW)
1	Summit - IBM Power System AC922, IBM POWER9 22C 3.07GHz, NVIDIA Volta GV100, Dual-rail Mellanox EDR Infiniband , IBM D0E/SC/Oak Ridge National Laboratory United States	2,397,824	143,500.0	200,794.9	9,783
2	Sierra - IBM Power System S922LC, IBM POWER9 22C 3.1GHz, NVIDIA Volta GV100, Dual-rail Mellanox EDR Infiniband , IBM / NVIDIA / Mellanox DOE/NNSA/LLNL United States	1,572,480	94,640.0	125,712.0	7,438
3	Sunway TainuLight - Sunway MPP, Sunway SWZ6010 Z60C 1.450HZ, Sunway , NRCPC National Supercomputing Center in Wuxi China	10,647,600	93,014.6	120,430.9	10,371
4	Tianhe-2A - TH-IVB-FEP Cluster, Intel Xeon E5-2692v2 12C 2.2GHz, TH Express-2, Matrix-2000, NUDT National Super Computer Center in Guangzhou China	4,981,760	61,444.5	100,678.7	18,482
5	Piz Daint - Cray XC50, Xeon E5-2690v3 12C 2.6GHz, Aries interconnect , NVIDIA Tesla P100 , Cray Inc. Swiss National Supercomputing Centre (CSCS) Switzerland	387,872	21,230.0	27,154.3	2,384
6	Trinity - Cray XC40, Xeon Eb-2698v3 16C 2.3GHz, Intel Xeon Phi 7250 68C 1.4GHz, Aries interconnect , Cray Inc. DOE/NNSA/LANL/SNL United States	979,072	20,158.7	41,461.2	7,578
7	Al Bridging Cloud Infrastructure (ABCI) - PRIMERGY CX2570 M4, Xeon Gold 6148 20C 2.4GHz, NVIDIA Tesla V100 SXM2, Infiniband EDR, Fujitsu National Institute of Advanced Industrial Science and Technology (AIST) Japan	391,680	19,880.0	32,576.6	1,649
8	SuperMUC-NG - ThinkSystem SD530, Xeon Platinum 8174 24C 3.1GHz, Intel Omni-Path , Lenovo Leibniz Rechenzentrum Germany	305,856	19,476.6	26,873.9	
9	Titan - Cray XK7, Opteron 6274 16C 2.200GHz, Cray Gemini interconnect, NVIDIA K20x, Cray Inc. D0E/SC/Oak Ridge National Laboratory Upitad States	560,640	17,590.0	27,112.5	8,209

#### **Today's and Tomorrow's Top 500 list**

NVIDIA GPUs to Power New Berkeley National Lab Supercomputer, Accelerating Scientific Discoveries

October 30, 2018 by GEETIKA GUPT

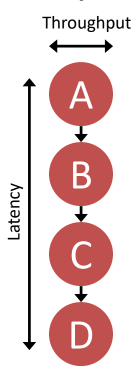
# Cray, AMD to build 1.5 exaflops supercomputer for US government

System will mix Epyc CPUs and Radeon Instinct GPUs.

PETER BRIGHT - 5/7/2019, 6:26 PM

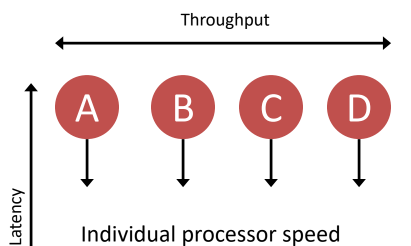
#### Latency vs. Throughput in Apps & Datasets

#### Latency-bound



Increasing the # of processors doesn't matter, only straight-line performance.

#### Throughput-bound

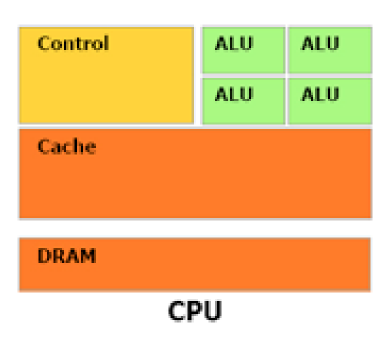


Individual processor speed increases matter less than processor count.

#### **Latency-Optimized Processors**

For most consumer (i.e. interactive) applications, latency is the #1 performance metric.

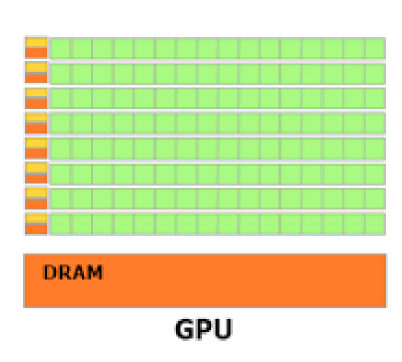
- ↑ clock frequencies
- ↑ control logic (e.g. branch prediction)
- ↑ cache per core
- parallelism/throughput



### **Throughput-Optimized Processors**

For many scientific applications, throughput is the #1 performance metric.

- ↑ cores/concurrency
- nemory bandwidth
- clock frequency
- **↓** cache
- **♦** control logic



### **Throughput-Bound Applications**

While many applications are latency-bound, the important ones (scientific, finance, medical, etc.) tend to be throughput bound.

T	ΔRI	F	6:	Har	dwar	e Con	figu	ratio

Device	Туре	Number of cores	Memory size
4 x GPU	GeForce GTX Titan 0.876 GHz	2688	6 GB Global Memory
CPU	Intel Core i7-4770 3.40 GHz	4	32 GB

#### ARLE 7: Result

Number of	Chronos 4 GPUs	Well Known	Performance	
Elements	Chronos 4 GPOS	FE Program	Improvement	
200.000	21	516 (8.6 min)	24,57 x	
500.000	43	3407 (56.78 min)	79,23 x	
1.000.000	83	Insufficient Memory	x	
2.000.000	168	Insufficient Memory	x	

"Speeding up a Finite Element Computation on GPU", GTC 15

_	Reconstruction Stage	Single CPU time	Single GPU time	Single GPU speedup	Two GPU time	Two GPU speedup
_	Preprocessing and Support Function	34.09	9.36	3.64X	7.72	4.42X
	Refraction-Corrected Ray Tracing	1899.98	63.29	30.02X	45.53	41.73X
	Compounding Views	39.33	0.84	46.71X	0.84	46.71X
	Entire Reflection Reconstruction	2108.40	79.16	26.63X	54.57	38.64X

"Multi-GPU Accelerated Refraction-Corrected Reflection Image Reconstruction for 3D Ultrasound Breast Imaging", GTC 15

#### Overview of (NVIDIA) GPU Architecture

#### **Architectural Generations**

Discussion will focus on characteristics shared across architectural generations.

When important, differences between generations will be noted.

Fermi → Kepler → Maxwell →
 Pascal → Volta

Much of the material in this section can be found in the white papers below.



http://www.nvidia.com/content/pdf/fermi\_white\_papers/nvidia\_fermi\_compute\_architecture\_whitepaper.pdf
https://www.nvidia.com/content/PDF/kepler/NVIDIA-Kepler-GK110-Architecture-Whitepaper.pdf
https://developer.nvidia.com/maxwell-compute-architecture
http://images.nvidia.com/content/pdf/tesla/whitepaper/pascal-architecture-whitepaper.pdf
https://devblogs.nvidia.com/parallelforall/inside-volta/

#### The CUDA Core

The CUDA Core is the finest granularity of execution on NVIDIA GPUs.

In a single cycle, executes either:

- 32-bit integer arithmetic instruction
- 32-bit floating point arithmetic instruction

CUDA core != x86 core.

CUDA Core

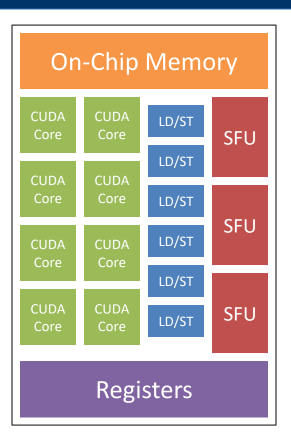
# The Streaming Multiprocessor (SM)

SM is analogous to "core" in x86 processors

#### Consists of many CUDA cores

- 32 for Fermi, 192 for Kepler, 128 for Maxwell, 64 for Pascal
- All CUDA cores in the same SM execute in lock step
- Registers + on-chip memory are shared among threads running on this SM

Think "vector core" but with more flexibility.

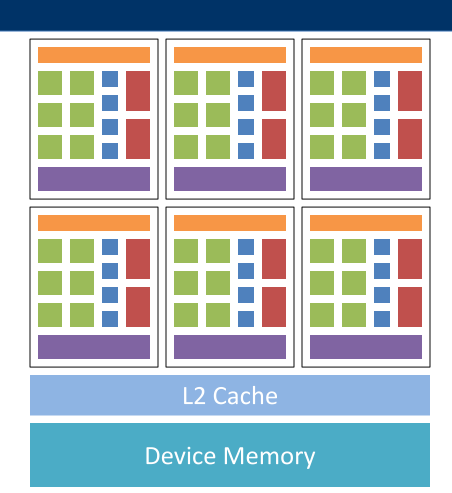


#### The GPU

Consists of O(10) SMs

L2 cache, device memory (DRAM) shared among all SMs

Off-chip latency ~100x worse than on-chip



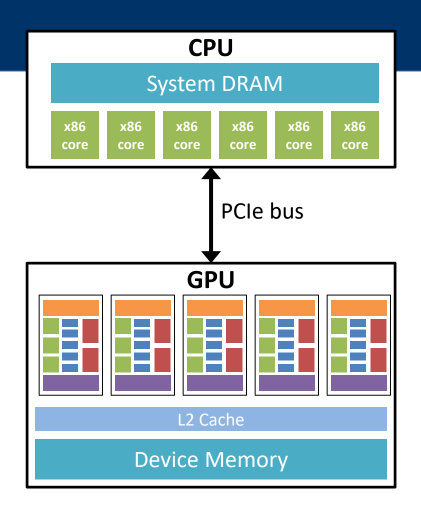
#### The Full System

GPUs are not standalone processors, require a CPU to manage them.

Commonly refer to CPU as "host" and GPU as "device".

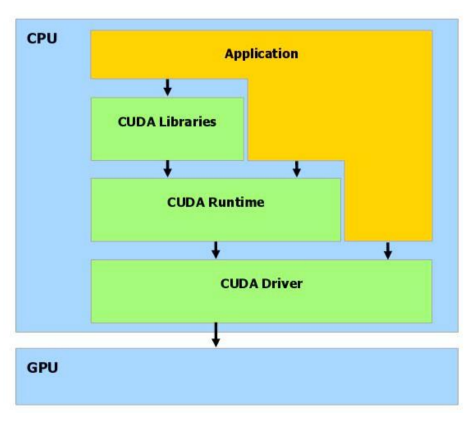
Connected by very high latency PCIe bus.

Analogous to two MPI processes, versus OpenMP/Cilk everything shared model.



### Hands-On With CUDA, a Simple Example

#### **CUDA Software Stack**



Kernel is written as single-threaded code.

A single CUDA thread per CUDA core, each CUDA thread executes same kernel (similar to SPMD).

Let's start by considering vector addition: C = B + A

```
void vector_add(int *C, int *B, int *A, int N) {
   int i;
   for (i = 0; i < N; i++) {
      C[i] = A[i] + B[i];
   }
}</pre>
```



CUDA Core

```
To enable GPU execution of this function, simply add __global__:
  global void vector add(int *C, int *B, int *A,
    int N) {
  int i;
  for (i = 0; i < N; i++) {
   C[i] = A[i] + B[i];
                                                         CUDA
```

#### CUDA function type qualifiers:

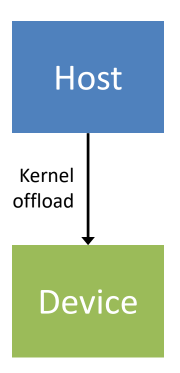
Qualifier	Executes on	Callable from
global	device	host
host	host	host
device	device	device

```
__device__ void bar(...) { ... }
__global__ void foo(...) {
  bar(...);
}
```



CUDA Core

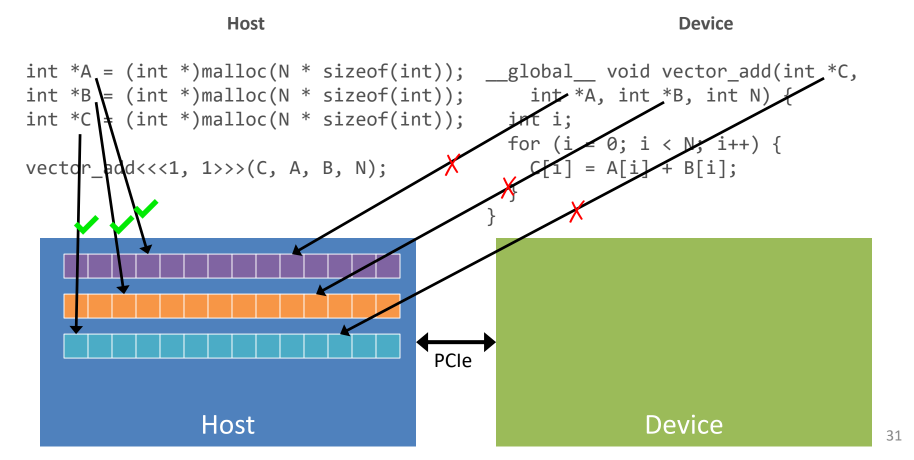
Calling this kernel from the host requires special syntax to indicate that the function call runs on the device:



One final step: the data.

Recall that host and device are physically discrete processors  $\rightarrow$  separate address spaces, a pointer on one has no meaning on the other.

```
Host Device
```



Host **Device** int \*A = (int \*)malloc(N \* sizeof(int)); \_\_global\_\_ void vector\_add(int \*C, for (i = 0; i < N; i++) { vector add<<<1, 1>>>(C, A, B, N); C[i] = A[i] + B[i];**PCle** Device Host

```
cudaError_t cudaMalloc(void **devPtr, size_t size);
vector add < <<1, 1>>> (C, A, B, N);
                   Allocate size bytes of memory on GPU.
                          Store address at *devPtr.
```

Host

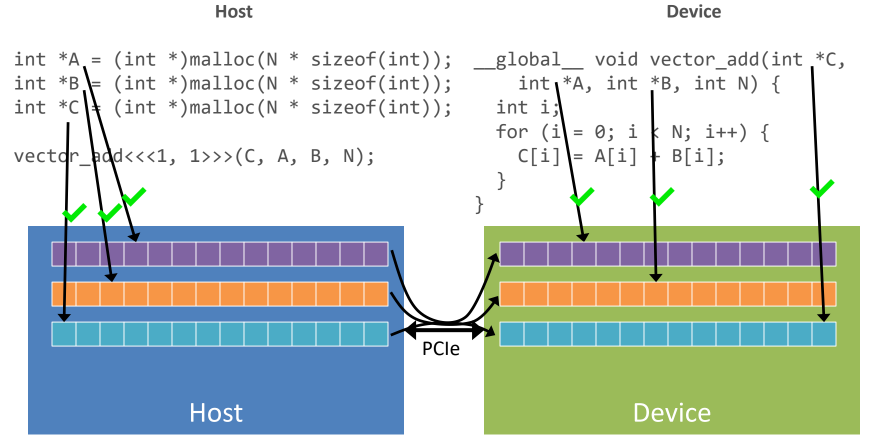
int \*A = (int \*)malloc(N \* sizeof(int)); global void vector add(int \*C, for (i = 0; i < N; i++) { vector add<<<1, 1>>>(C, A, B, N); C[i] = A[i] + B[i];**PCle** Device Host

**Device** 

Transfer size bytes from src to dst.

Address spaces must obey direction, can be:

- cudaMemcpyHostToHost
- cudaMemcpyDeviceToDevice
- cudaMemcpyHostToDevice
- cudaMemcpyDeviceToHost



# A Simple CUDA Example

```
int *A, *d A;
A = (int *)malloc(N * sizeof(int));
                                                                   Host
cudaMalloc((void **)&d A, N * sizeof(int));
cudaMemcpy(d A, A, N * sizeof(int), cudaMemcpyHostToDevice);
vector add<<<1, 1>>>(d C, d A, d B, N);
      cudaError t cudaMalloc(void **devPtr, size t size);
                                                                  Device
      cudaError t cudaMemcpy(void *dst, const void *src,
          size t size, enum cudaMemcpyKind direction);
      cudaError_t cudaFree(void *addr);
```

# A Simple CUDA Example

Building CUDA programs requires **nvcc**. Supports all of the usual flags you expect from a compiler.

<sup>\*</sup>Makefiles will be provided for your convenience

# Hands On - 00\_vecadd

Don't forget to follow the instructions on slide 1.

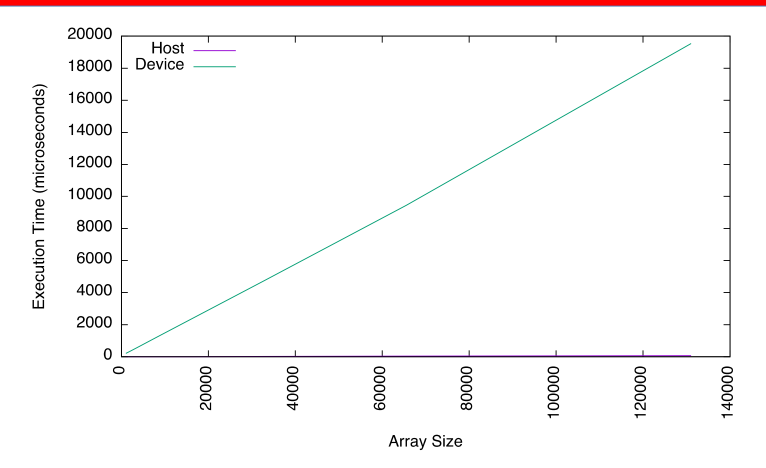
```
$ cd ~/bootcamp-gpu/src/00_vecadd
$ make
$ vim/emacs vecadd.cu # Get familiar with the code
$ ./vecadd 131072 # Vectors with 131,072 elements
```

What kind of performance do you observe?

Does it change relatively with different input sizes?

\$ git clone git@github.com:agrippa/hpc-bootcamp.git \$HOME/bootcamp-gpu

# Hands On - 00\_vecadd



40

#### **Review - Basic CUDA**

Running a function on the GPU requires 1) \_\_global\_\_ annotation, and 2) kernel launch syntax (<<<...>>>).

CUDA forces you to think about discrete address spaces and manage coherency.

Copying and pasting code is not the path to performance in CUDA.

# nvprof

nvprof is a command-line profiler for CUDA applications.

```
$ nvprof --help
Usage: nvprof [options] [CUDA-application] [application-arguments]
...
```

Can be used in a number of modes:

- Default/Summary Mode
- API Trace Mode
- Event/Metric Summary Mode
- Event/Metric Trace Mode

# **Hands On – nvprof Summary mode**

A good starting point is usually Default/Summary mode.

\$ nvprof ./vecadd 131072

# **Hands On – nvprof Summary mode**

A good starting point is usually Default/Summary mode.

\$ nvprof ./vecadd 131072

# nvprof

Diving deeper requires Event/Metric Summary Mode.

```
$ nvprof --query-events # List all enabled events
$ nvprof --query-metrics # List all enabled metrics
```

**Events** are raw data collected by the hardware, e.g. # instructions (inst\_issued)

**Metrics** are derived based on events, e.g. # instructions per cycle (issued\_ipc)

# **Hands On – nvprof Metrics**

Diving deeper requires Event/Metric Summary Mode.

- 1. ipc: # of instructions executed per cycle on the GPU
- 2. sm\_efficiency: % of time an SM on the GPU is doing useful work
- 3. alu\_fu\_utilization: indicates how well the CUDA cores are being used

```
$ nvprof --metrics ipc,sm_efficiency,alu_fu_utilization ./vecadd 131072
```

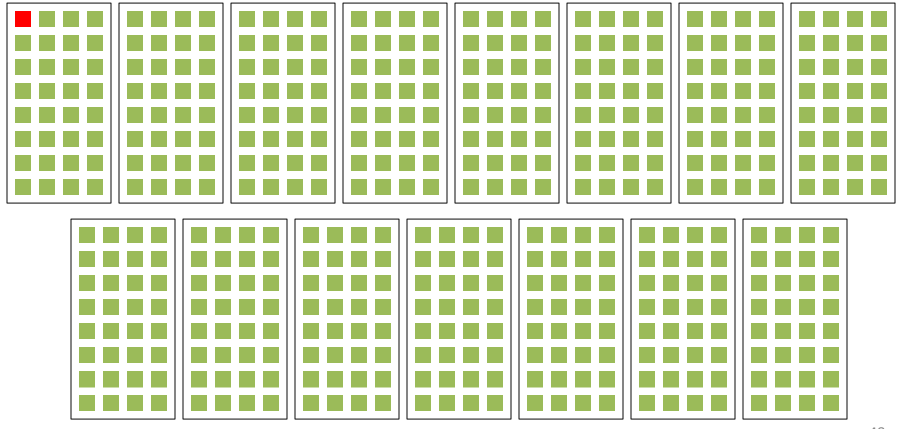
# **Hands On – nvprof Metrics**

Diving deeper requires Event/Metric Summary Mode.

```
$ nvprof --metrics ipc,sm_efficiency,alu_fu_utilization ./vecadd 131072
```

```
==30413== Warning: Some kernel(s) will be replayed on device 0 in order to collect all events/metrics.
Finished! All 131072 elements validate.
Took 621 microseconds on the host
Took 174666 microseconds on the device, 0.00356x speedup
==30413== Profiling application: ./vecadd 131072
==30413== Profiling result:
==30413== Metric result:
Invocations
                                          Metric Name
                                                                              Metric Description
                                                                                                         Min ...
Device "Tesla M2050 (0)"
        Kernel: vector add(int*, int*, int*, int)
                                        sm efficiency
                                                                         Multiprocessor Activity
                                                                                                       7.14% ...
                                                                                    Executed TPC
                                                                                                    0.103982 ...
                                                   ipc
                                                            Arithmetic Function Unit Utilization
                                                                                                     Low (1) ...
                                   alu fu utilization
```

# nvprof



Or How to Write Parallel CUDA Programs

# Flynn's Taxonomy

	Single Instruction	Multiple Instructions
Single Data	SISD	MISD
Multiple Data	SIMD	MIMD

#### Single Instruction, Single Data stream (SISD)

A sequential computer which exploits no parallelism in either the instruction or data streams. e.g., old single processor PC

#### Single Instruction, Multiple Data streams (SIMD)

A computer which exploits multiple data streams against a single instruction stream to perform operations which may be naturally parallelized. e.g. graphics processing unit

#### Multiple Instruction, Single Data stream (MISD)

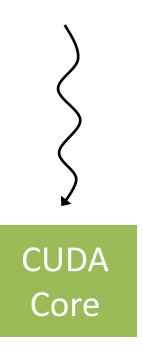
Multiple instructions operate on a single data stream. Uncommon architecture which is generally used for fault tolerance. Heterogeneous systems operate on the same data stream and must agree on the result. e.g. the Space Shuttle flight control computer.

#### Multiple Instruction, Multiple Data streams (MIMD)

Multiple autonomous processors simultaneously executing different instructions on different data. e.g. a PC cluster memory space.

#### Recall:

- Programmer writes single-threaded kernel
- 1 CUDA thread per CUDA core.

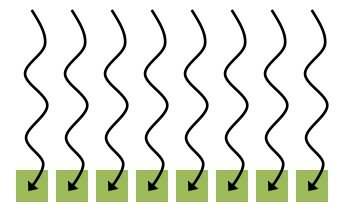


#### Recall:

- Programmer writes single-threaded kernel
- 1 CUDA thread per CUDA core.

Threads are scheduled in groups of 32, called a warp.

- Warps execute in lock step (same instruction at same cycle)
- All threads in warp on same SM



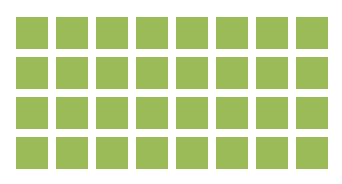
#### Recall:

- Programmer writes single-threaded kernel
- 1 CUDA thread per CUDA core.

Threads are scheduled in groups of 32, called a warp.

#### Warps are grouped into thread blocks.

- Warps in a block execute on same SM
- Warp execution in same thread block can be interleaved with each other on SM
- SM resources are partitioned among all threads in resident thread blocks



e.g. 4 warps in one thread block

#### Recall:

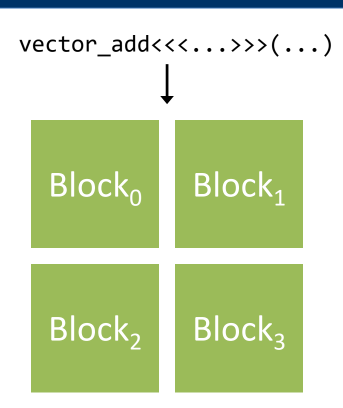
- Programmer writes single-threaded kernel
- 1 CUDA thread per CUDA core.

Threads are scheduled in groups of 32, called a warp.

Warps are grouped into **thread blocks**.

A single kernel **grid** can include many thread blocks.

 Kernel call is not complete until all thread blocks within it complete.



Blocks and grids are configurable by kernel launch arguments:

```
vector_add<<<nblocks, nthreads_per_block>>>(...)
vector_add<<<1, 1>>>(...); // 1 block with 1 thread inside it
vector_add<<<1, 32>>>(...); // 1 block with 1 full warp
vector_add<<<1, 256>>>(...); // 1 block with 256 threads (8 warps)
vector_add<<<N, 256>>>(...); // N blocks with 256 threads/block
```

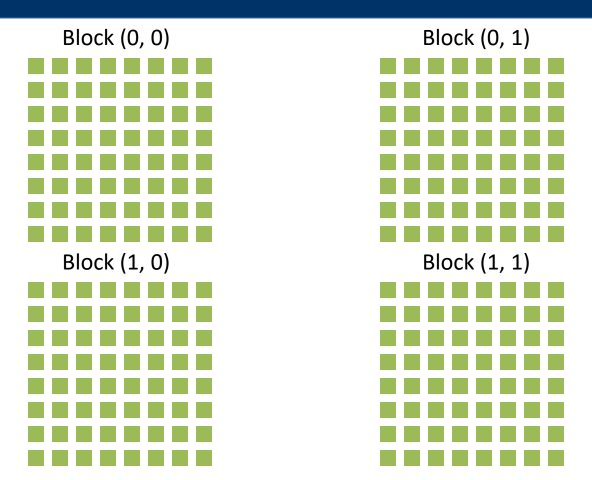
Blocks and grids can also be multi-dimensional:

```
// Equivalent to vector_add<<<1, 256>>>(...);
dim3 1d_block(256, 1, 1);
vector_add<<<1, 1d_block>>>(...);
dim3 block(16, 16); // 2D 16x16 block of threads, 256 total
dim3 grid(4, 4); // 2D 4x4 grid of thread blocks, 16 total
kernel<<<grid, block>>>(...);
```

Special CUDA kernel variables can be used to check the coordinates of a thread in an executing grid.

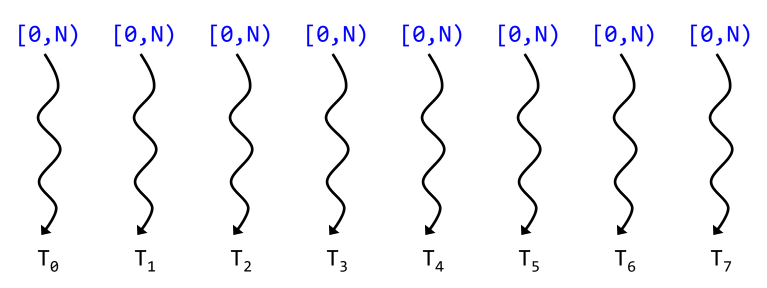
Variable Name	Description
threadIdx.x, threadIdx.y,	Offset of a thread within a thread block
blockIdx.x, blockIdx.y,	Offset of a block within a grid
blockDim.x,	# of threads per block
gridDim.x,	# of blocks in grid
<pre>blockIdx.x * blockDim.x</pre>	One way to calculate globally unique thread ID for 1D blocks and grids

```
global void matrix add(...) {
  const int row = blockIdx.x * blockDim.x + threadIdx.x;
 const int col = blockIdx.y * blockDim.y + threadIdx.y;
dim3 block(8, 8); // 2D 8x8 block of threads, 64 total
dim3 grid(2, 2); // 2D 2x2 grid of thread blocks, 4 total
vector add<<<grid, block>>>(...);
```



Launch configuration alone is insufficient for our vector\_add to run in parallel.

vector\_add<<<N / 256, 256>>>(C, A, B, N);



Must also change the work performed by the kernel.

```
global__ void vector_add(int *C, int *B, int *A, int N) {
const int i = blockIdx.x * blockDim.x + threadIdx.x;
C[i] = A[i] + B[i];
```

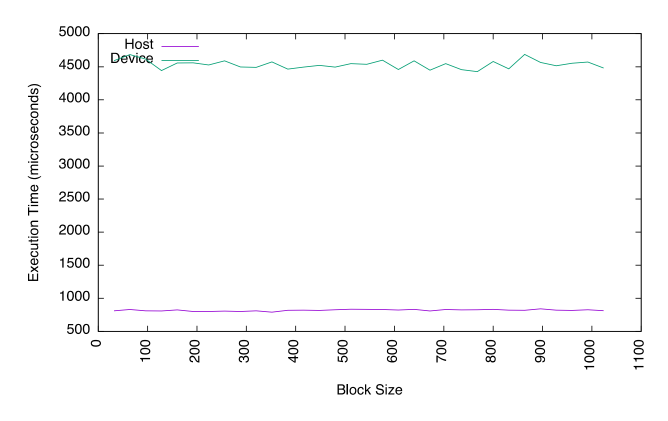
Using the previous slides as a guide, add parallelism to the vecadd example:

- 1. Change the <<<...>>> launch configuration to use B threads per block, N/B blocks
  - Be careful that the N you run is evenly divisible by B
  - CUDA supports up to 1,024 threads per block
- 2. Change the kernel to process a single data point

Does performance improve?

\$ ./vecadd 1048576

Does block size affect performance?



More similar host and device performance (was ~150,000 us).

No clear trends in performance as a function of block size (yet).

Still not great performance, time to go back to nvprof.

\$HOME/bootcamp-gpu/src/01\_vecadd contains an example solution that allows configuring threads per block from the command line:

\$ nvprof --metrics ipc,sm\_efficiency,alu\_fu\_utilization ./vecadd 1048576

```
==32216== NVPROF is profiling process 32216, command: ./vecadd 1048576
==32216== Warning: Some kernel(s) will be replayed on device 0 in order to collect all events/metrics.
Finished! All 1048576 elements validate using 256 threads per block.
Took 5032 microseconds on the host
Took 21110 microseconds on the device, 0.23837x speedup
                                                                                       0.103982
                                                                                                   7.14 %
==32216== Profiling application: ./vecadd 1048576
==32216== Profiling result:
==32216== Metric result:
Invocations
                                          Metric Name
                                                                             Metric Description
                                                                                                         Min
Device "Tesla M2050 (0)"
        Kernel: vector add(int*, int*, int*, int)
                                        sm efficiency
                                                                        Multiprocessor Activity
                                                                                                     96.20%
                                                                                    Executed IPC
                                                                                                    0.657917
                                                  ipc
                                                           Arithmetic Function Unit Utilization
                                   alu fu utilization
                                                                                                     LOW (3)
                                                                                                  Low (1)
```

#### \$ nvprof ./vecadd 1048576 256

```
[jmg3@gpu-014 01 vecadd]$ /opt/apps/software/Compiler/GCC/4.4.7/CUDA/6.5.14/bin/nvprof ./vecadd
1048576 32
==32199== NVPROF is profiling process 32199, command: ./vecadd 1048576 32
Finished! All 1048576 elements validate using 32 threads per block.
Took 5049 microseconds on the host
Took 4541 microseconds on the device, 1.11187x speedup
==32199== Profiling application: ./vecadd 1048576 32
==32199== Profiling result:
                                    Min
Time(%)
           Time
                 Calls Avg
                                                Max Name
44.85% 1.5758ms
                       1 1.5758ms 1.5758ms 1.5758ms [CUDA memcpy DtoH]
43.93% 1.5436ms 2 771.81us 767.70us 775.93us [CUDA memcpy HtoD]
 10.26% 360.38us 1 360.38us 360.38us vector add(int*, int*, int*, int)
 0.96% 33.856us
                       1 33.856us 33.856us [CUDA memset]
```

### Review – CUDA Execution Model

**nvcc** – Compile CUDA applications, shares many flags with x86 compilers, use -arch to target specific GPU generations.

**nvprof** – Tool for analyzing CUDA applications, profile-driven optimization!

CUDA thread hierarchy: thread, warp, thread block, and grid

Access thread coordinates from running kernel via **blockIdx.x**, **threadIdx.x**, **blockDim.x** 

Good to maximize parallelism of CUDA kernels over input, PCIe transfers can quickly become bottlenecks

Most APIs are synchronous by default.

• When you call malloc, you expect its return value to point to immediately addressable memory.

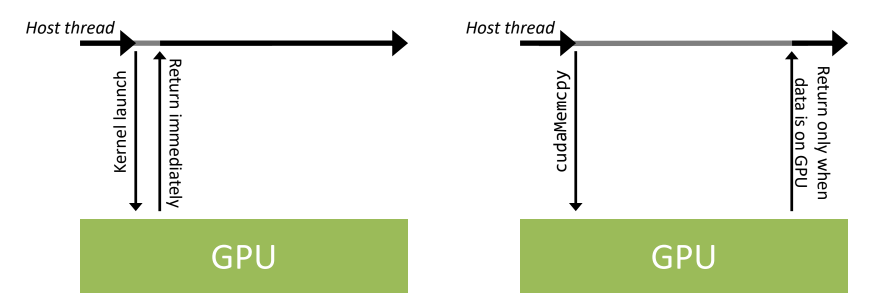
Commonly, high latency operations support some level of asynchrony:

- POSIX AIO
- NodeJS/Javascript callbacks
- E-mail

Overlapping useful work on-core with asynchronous work off-core can benefit performance.

CUDA supports both blocking and asynchronous APIs.

A kernel launch <<<...>>> is asynchronous. A cudaMemcpy is blocking.

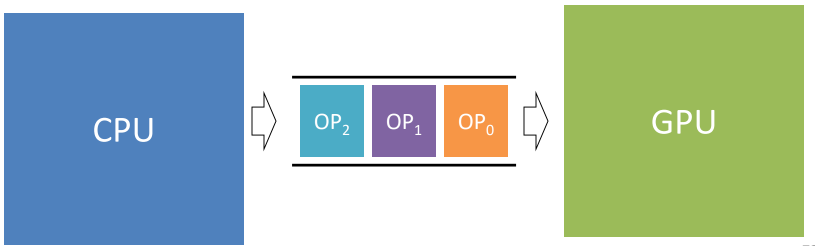


Some blocking APIs have asynchronous equivalents. Some don't.

Blocking API	Asynchronous API
cudaMalloc	
cudaMemcpy	cudaMemcpyAsync
	Kernel launch (<<<>>>)
cudaFree	

The core of CUDA asynchrony is **CUDA streams**.

A CUDA stream defines a sequence of CUDA operations (e.g. kernel launch, memory copy, etc.) to be performed in order.



Streams must be explicitly created by the programmer:

```
cudaStream_t stream;
cudaError_t cudaStreamCreate(cudaStream_t *stream);
cudaError_t cudaStreamDestroy(cudaStream_t *stream);
```

Every asynchronous operation is associated with a CUDA stream.

If the CUDA stream is not explicitly set, a default stream is used.

Implicit default stream

kernel<<<...>>(...);
cudaMemcpyAsync(...);

**Explicit stream** 

```
kernel<<<..., stream>>>(...);
cudaMemcpyAsync(..., stream);
```

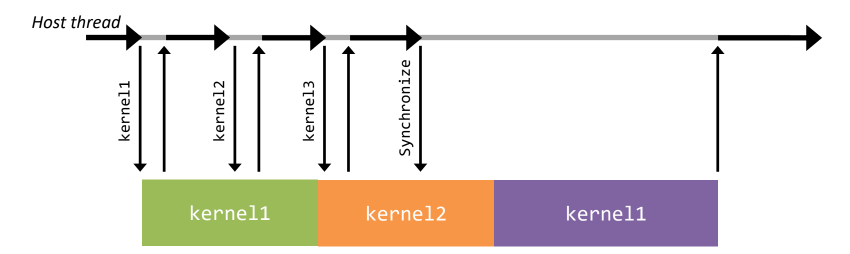
Stream status can be queried or blocked on.

Implicitly includes all past operations in that stream.

```
cudaStream_t stream;
cudaStreamCreate(&stream);
kernel<<<..., stream>>>(...);
cudaStreamSynchronize(stream);
```

```
cudaError_t cudaStreamSynchronize(cudaStream_t stream);
cudaError_t cudaStreamQuery(cudaStream_t stream);
```

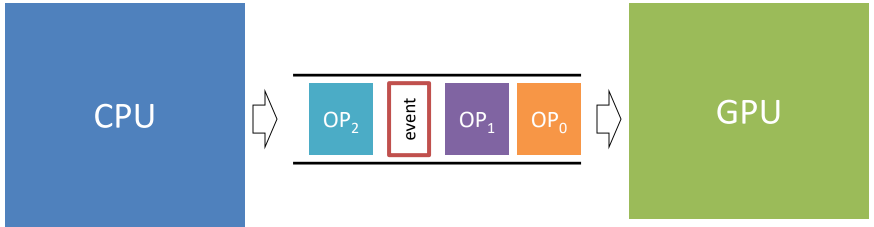
```
kernel1<<<..., stream>>>(...);
kernel2<<<..., stream>>>(...);
kernel3<<<..., stream>>>(...);
cudaStreamSynchronize(stream);
```



Also possible to query specific points in time within a stream using CUDA Events.

Events are inserted in to streams and satisfied as they exit the stream.

Enable querying of all operations inserted in to the stream prior to the event.

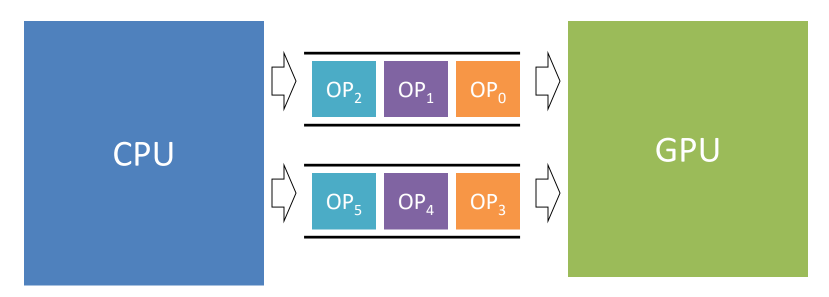


Like streams, events can be created, destroyed, queried, and blocked on.

Placing an event in a stream:

Synchronizing on all operations issued to a given device is possible with cudaDeviceSynchronize.

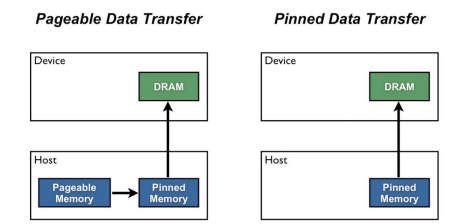
cudaError\_t cudaDeviceSynchronize();



### Special Note on cudaMemcpyAsync

Host allocations transferred through cudaMemcpyAsync must be page-locked/pinned.

CUDA provides cudaMallocHost for page-locked host allocations.



cudaError\_t cudaMallocHost(void \*\*ptr, size\_t size);

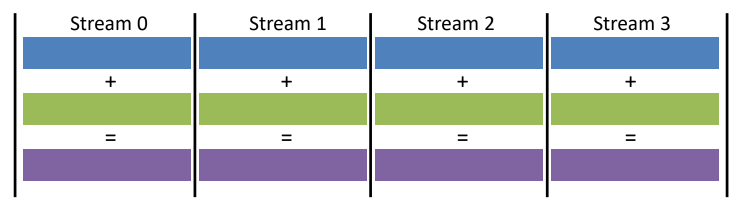
\$HOME/bootcamp-gpu/src/02\_vecadd\_streams/vecadd.cu contains an example of chunked, synchronous vector addition.



\$ ./vecadd <N> <nchunks>

### Hands On – CUDA Streams

Complete the template in \$HOME/bootcamp-gpu/src/02\_vecadd\_streams/vecadd.cu based on the TODOs contained to use a different stream on each vector chunk.



\$ ./vecadd 4194304 4 # Vector of 4,194,304 elements, 4 chunks

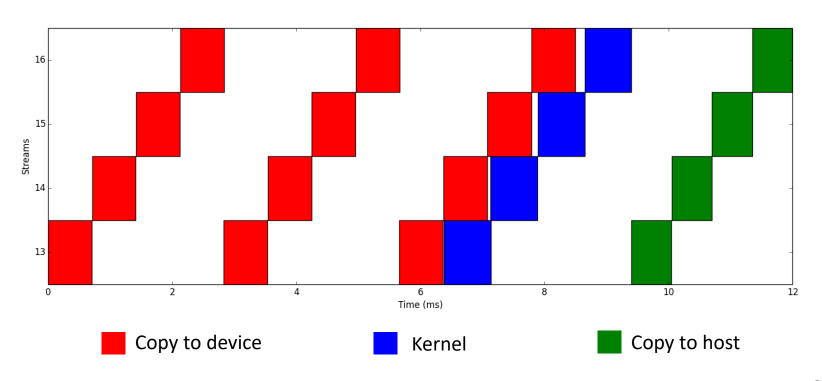
Do multiple streams significantly improve performance for this example?

### Hands On – CUDA Streams

Try using nvprof's GPU tracing mode to understand stream interleaving.

```
$ nvprof --print-gpu-trace ./vecadd_solution 4194304 1
$ nvprof --print-gpu-trace ./vecadd solution 4194304 4
```

### **Hands On – CUDA Streams**



# Review – CUDA Asynchrony

A **CUDA stream** defines a sequence of CUDA operations (e.g. kernel launch, memory copy, etc.) to be performed in order.

**CUDA events** are inserted in to streams and satisfied as they exit the stream, mark a specific point-in-time in the processing of the operations in that stream.

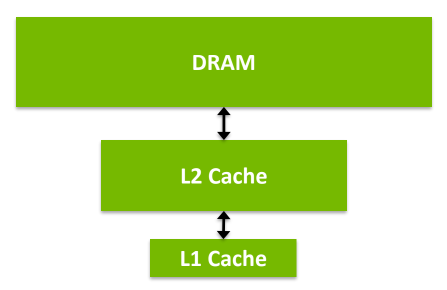
Using asynchronous APIs requires an understanding of streams (and events), can benefit performance by enabling host-device, host-copy, device-copy overlaps.

# **Memory Hierarchies**

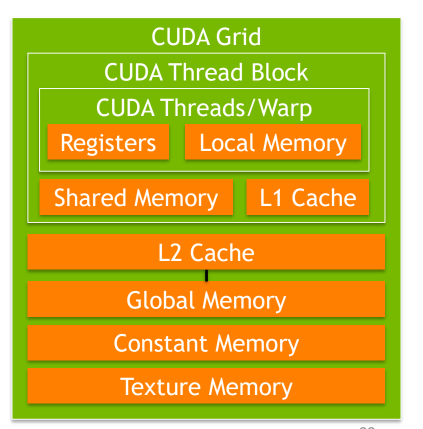
Up until now, we've glossed over efficient data access in CUDA kernels.

On CPU, this generally involves optimizing for cache line locality.

A memory hierarchy emulates a large amount of low-latency memory

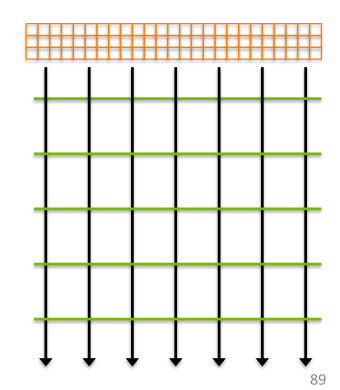


- The CUDA Memory Hierarchy is more complex than the CPU's
  - Many different types of memory, each with special-purpose characteristics
  - More explicit control over data movement



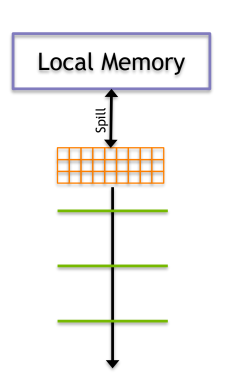
#### Registers

- Lowest latency memory space on the GPU
- Private to each CUDA thread
- Constant pool of registers per-SM divided among threads in resident thread blocks
- Architecture-dependent limit on number of registers per thread
- Registers are not explicitly used by the programmer, implicitly allocated by the compiler
- nvcc -maxrregcount allows you to limit # registers per thread



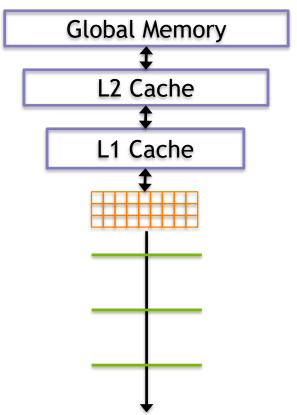
#### Local Memory

- When registers are exhausted, variables spill to local memory
- Variables likely to be placed in local memory: large local structures or arrays, local arrays whose indices cannot be determined at compile-time
- Local memory is not physical, variables stored in local memory are spilled to Global Memory, L1 cache, or L2 cache
- Not explicitly controlled by programmer



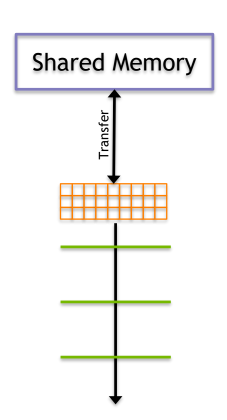
#### GPU Caches

- Behaviour of GPU caches is architecturedependent
- Per-SM L1 cache stored on-chip
- Per-GPU L2 cache stored off-chip, caches values for all SMs
- Due to parallelism of accesses, GPU caches can be difficult to reason about



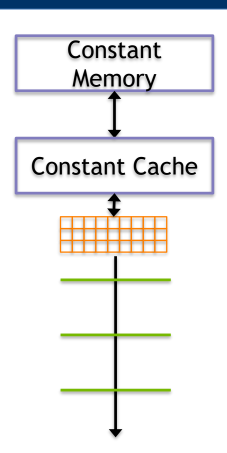
#### Shared Memory

- Declared with the \_\_shared\_\_ keyword
- Low-latency, high bandwidth
- Shared by all threads in a thread block
- Explicitly allocated and managed by the programmer, manual L1 cache
- Stored on-SM, same physical memory as the GPU L1 cache
- On-SM memory is statically partitioned between L1 cache and shared memory



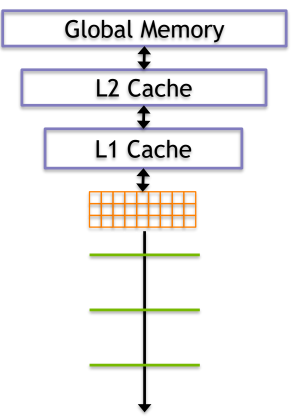
#### Constant Memory

- Declared with the constant keyword
- Read-only
- Limited in size: 64KB
- Stored in device memory (same physical location as Global Memory)
- Cached in a per-SM constant cache
- Optimized for all threads in a warp accessing the same memory cell



#### Global Memory

- Large, high-latency memory
- Stored in device memory (along with constant and texture memory)
- Can be declared statically with \_\_\_device\_\_\_
- Can be allocated dynamically with cudaMalloc
- Explicitly managed by the programmer
- Optimized for all threads in a warp accessing neighbouring memory cells



# **Review - CUDA Memory Spaces**

MEMORY	ON/OFF CHIP	CACHED	ACCESS	SCOPE	LIFETIME
Register	On	n/a	R/W	1 thread	Thread
Local	Off	†	R/W	1 thread	Thread
Shared	On	n/a	R/W	All threads in block	Block
Global	Off	†	R/W	All threads + host	Host allocation
Constant	Off	Yes	R	All threads + host	Host allocation
Texture	Off	Yes	R	All threads + host	Host allocation

# **CUDA Kernel Synchronization**

# **Kernel Synchronization**

Synchronization in CUDA is closely tied with the CUDA memory hierarchy.

Relative to CPU programming models, synchronization options are constrained in CUDA.

- CUDA is designed to be highly scalable and portable across generations
- Lots of synchronization negates both of these principles

There is no global synchronization in CUDA (i.e. no #pragma omp barrier) – global synchronization requires kernel termination.

Instead, CUDA offers local synchronization and some global memory fences.

For performance reasons, use of these should still be minimized

# **Kernel Synchronization**

Function	Action	Scope	Notes
syncthreads()	Barrier	Thread block	Most commonly useful function.
threadfence_block()	Memory fence	Thread block	Applies to shared and global mem.
threadfence()	Memory fence	Device	Same asthreadfence_block(), but also a global memory write fence relative to all threads on the device.
threadfence_system()	Memory fence	Whole system	Same asthreadfence_block(), but also a global memory write fence relative to all threads on the device and host and read fence for calling thread.

# **Review - Kernel Synchronization**

Good chance you'll only ever use \_\_syncthreads()

Avoid synchronization at all costs.

Global synchronization by kernel termination.

CPUs rely heavily on hardware-managed caches for performance.

GPU caching is a more challenging problem

- Thousands of threads cooperating on a problem
- Difficult to predict the next round of accesses for all threads

For efficient global memory access, GPUs instead rely on:

- 1. Large device memory bandwidth
- 2. Aligned and coalesced memory access patterns
- 3. Maintaining sufficient pending I/O operations to keep the memory bus saturated and hide global memory latency

Aligned and coalesced global memory accesses are key to optimizing an application's use of global memory bandwidth.

- 1. Coalesced: threads in a warp reference memory addresses that can all be serviced by a single global memory transaction (think of a memory transaction as the process of bring a cache line into the cache)
- 2. Aligned: the global memory accesses by threads within a warp start at an address boundary that is an even multiple of the size of a global memory transaction

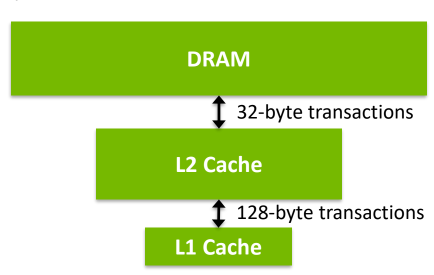
Global memory transactions are either 32 or 128 contiguous bytes.

Up to Maxwell, size of memory transaction depends on caches it passes through.

• **L1 + L2** - 128 bytes, **L2 only** - 32 bytes

With Maxwell/Pascal, all transactions are 32 bytes.

Information in the "Volta Tuning Guide" is sparse at this time, but expect similar behaviour to Pascal.



Reads Cacheable in L1?

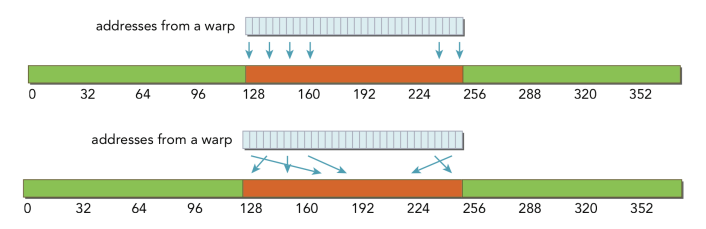
**Architecture** 

Which caches a global memory transaction passes through depends on GPU architecture and the type of access (read vs. write).

Fermi Kepler Kepler K40 + Maxwell Pascal P100 Pascal P104 Volta	Yes No Yes Yes Yes Yes Yes Yes Yes	Yes No No Yes No Yes Ves	Global memory reads cached in L2 if not in L1
Architecture Fermi	Writes Cacheable in L1?	Cached by Default?	Global memory writes cached in L2, up until Volta.
Kepler Kepler K40 through Volta	No No	No No	
Volta	Yes	Yes	103

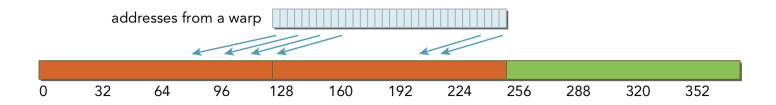
Cached by Default?

Aligned and Coalesced Memory Access (w/L1 cache)



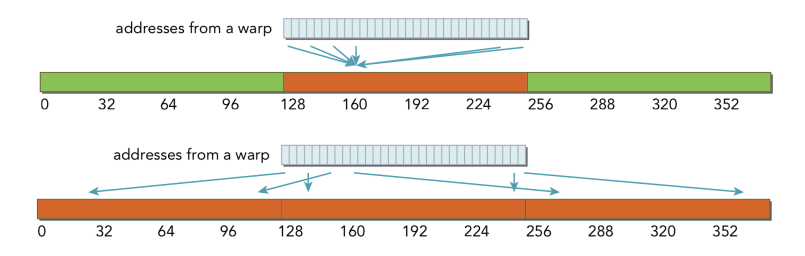
With 128-byte transactions, a single transaction is required and all of the loaded bytes are used

Misaligned and Coalesced Memory Access (w/L1 cache)



With 128-byte transactions, two memory transactions are required to load all requested bytes. Only half of the loaded bytes are used.

Misaligned and Uncoalesced Memory Access (w/L1 cache)



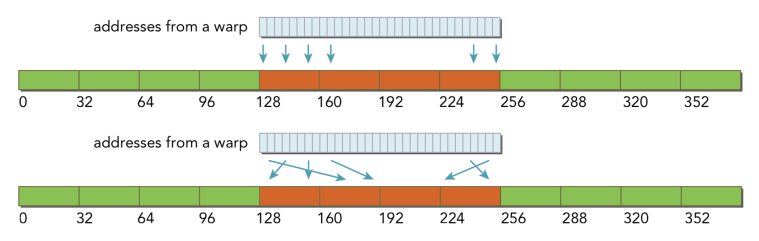
With uncoalesced loads, many more bytes loaded than requested.

Memory accesses that are not cached in L1 are serviced by 32-byte transactions

This can improve memory bandwidth utilization

However, the L2 cache is device-wide, higher latency than L1, and still relatively small → many applications may take a performance hit if L1 cache is not used for reads

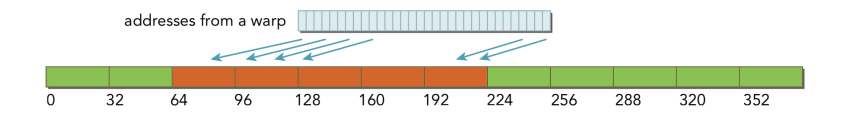
Aligned and Coalesced Memory Access (w/o L1 cache)



With 32-byte transactions, four transactions are required and all of the loaded bytes are used

### **Optimizing CUDA Memory Accesses**

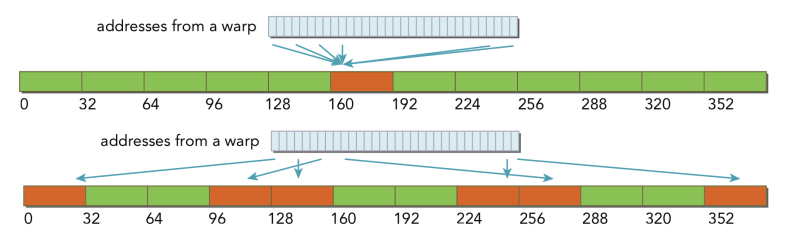
Misaligned and Coalesced Memory Access (w/o L1 cache)



With 32-byte transactions, extra memory transactions are still required to load all requested bytes but the number of wasted bytes is likely reduced compared to 128-byte transactions.

### **Optimizing CUDA Memory Accesses**

Misaligned and Uncoalesced Memory Access (w/o L1 cache)



With uncoalesced loads, more bytes loaded than requested but better efficiency than with 128-byte transactions.

# **Hands On – Memory Access Optimization**

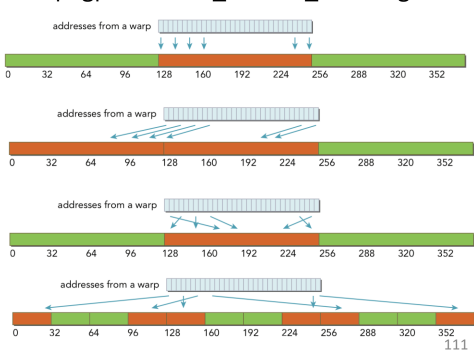
To illustrate these points, we'll experiment with a variation on the vector add micro-benchmark under \$HOME/bootcamp-gpu/src/03\_vecadd\_misaligned.

vector\_add
vector\_add\_read\_offset

vector\_add\_write\_offset

vector\_add\_weirdly\_coalesced

vector\_add\_not\_coalesced



### **Hands On – Memory Access Optimization**

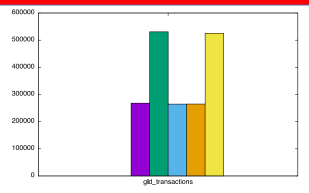
#### Experiment with several new metrics from nvprof:

- 1. gld\_transactions: # of global memory load transactions
- 2. gst\_transactions: # of global memory store transactions
- 3. gld\_transactions\_per\_request: mean # of gld txs issued to satisfy a warp's request
- 4. gst\_transactions\_per\_request: mean # of gst txs issued to satisfy a warp's request
- 5. gld efficiency: gld requests / gld transactions
- 6. gst\_efficiency: gst\_requests / gst\_transactions

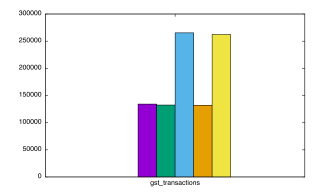
```
$ nvprof --metrics gld_transactions ./vecadd 4194304
$ nvprof --metrics gst_transactions ./vecadd 4194304
$ ...
```

What do you observe?

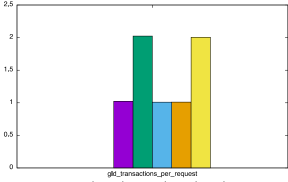
### **Hands On – Memory Access Optimization**



Misaligned, uncoalesced reads ↑ gld\_transactions

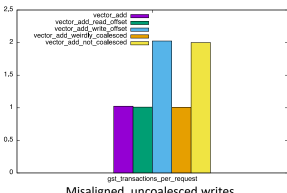


Misaligned, uncoalesced writes ↑ gst\_transactions



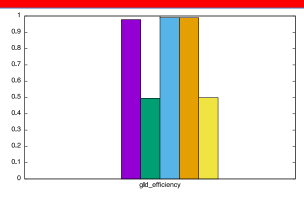
Misaligned, uncoalesced reads

↑ gld\_transactions\_per\_request

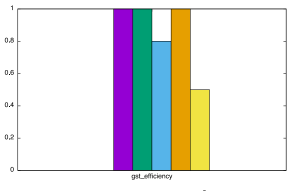


Misaligned, uncoalesced writes

↑ gst\_transactions\_per\_request



Misaligned, uncoalesced reads ♥ gld\_efficiency



Misaligned, uncoalesced writes ♥ gst\_efficiency

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## **Review - Optimizing CUDA Memory Access**

Seek **aligned** and **coalesced** global memory accesses when optimizing CUDA kernel performance.

The way in which a load/store is serviced depends on the cache(s) it passes through.

Aligned and coalesced accesses reduce the number of transactions necessary and the efficiency of each of those transactions (i.e. # bytes requested / # bytes loaded).

### **CUDA Atomic Operations (Briefly)**

### **Atomic Operations**

Atomic operations are a special class of mathematical operations in computing

- An atomic operation is performed uninterruptedly, so that there is no interference from other threads
- When a thread's atomic operation has completed, it can be certain its requested changes have been made without interference from other threads
- Atomic operations are particularly useful on the massively parallel GPU

Atomic operations may improve correctness, but can have a detrimental impact on performance

- Out of thousands of threads, only one can succeed in accessing a shared variable
- Atomicity requires reading and writing DRAM, no caching allowed

### **Atomic Operations**

```
__global__ void sumAll(int *in_array, int N, int *out_scalar) {
   int i = blockIdx.x * blockDim.x + threadIdx.x;
   if (i < N) {
        // out_scalar += out_array[i] would cause massive data race
        atomicAdd(out_scalar, in_array[i]);
   }
}</pre>
```

### **Atomic Operations Summary**

CUDA supports a variety of atomic operations. CAS can be used to build your own.

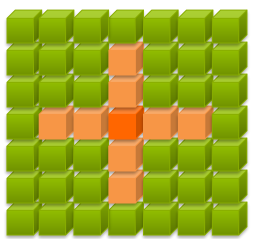
OPERATION	FUNCTION	SUPPORTED TYPES
Addition	atomicAdd	int, unsigned int, unsigned long long int, float
Subtraction	atomicSub	int, unsigned int
Unconditional Swap	atomicExch	int, unsigned int, unsigned long long int, float
Minimum	atomicMin	int, unsigned int, unsigned long long int
Maximum	atomicMax	int, unsigned int, unsigned long long int
Increment	atomicInc	unsigned int
Decrement	atomicDec	unsigned int
Compare-And-Swap	atomicCAS	int, unsigned int, unsigned long long int
And	atomicAnd	int, unsigned int, unsigned long long int
Or	atomicOr	int, unsigned int, unsigned long long int
Xor	atomicXor	int, unsigned int, unsigned long long int

### Hands on with a Real World App

## A (Slightly) More Complex Application

2D wavefront propagation (stencil)

```
for (int y = 0; y < ny; y++) {
 for (int x = 0; x < nx; x++) {
    for (int d = 1; d <= radius; d++) {
      div += c coeff[d] * (curr[y pos offset] +
         curr[y neg offset] + curr[x pos offset] +
         curr[x neg offset]);
    next[this offset] = temp + div * vsq[this offset];
```



We'll use iso2d as a more realistic example to explore optimizations.

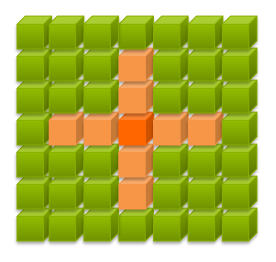
### **A More Complex Application**

#### Provided reference code:

04\_iso2d\_seq/: sequential C implementation

```
$ cd $HOME/bootcamp-gpu/src/04_iso2d_seq
$ make
```

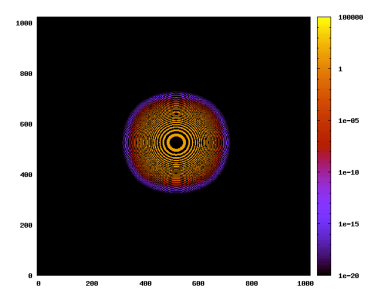
```
$ ./iso2d_seq -x 1024 -y 1024 -i 800 -t
```



### A More Complex Example

iso2d's -t flag will produce an output file to visualize for correctness.

```
$ ../iso2d_common/iso.sh -x 1024 -y 1024 -i snap.text
$ eog iso.png # Must use ssh -X to connect to DAVINCI
```



## Hands On – A Quick Review of OpenMP

Let's start by examining the parallelism possible in iso2d using OpenMP.

Where could you add a #pragma omp parallel for in iso2d.cpp for the most parallelism?

Compare the performance of your OpenMP implementation to the provided sequential code by modifying iso2d.cpp in 04\_iso2d\_seq.

Visualize the results to verify the correctness of your OpenMP implementation.

\$ ./iso2d\_omp -x 1024 -y 1024 -i 800

(If you're having trouble you can cheat and look in 05\_iso2d\_omp/)

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Next, your task is to port the OpenMP version of iso2d to CUDA, and check its performance.

06\_iso2d\_cuda/iso2d.cu contains TODOs to help in the port.

Use nvprof's Summary Mode to check where your implementation is spending time.

(If you're having trouble you can cheat and look in iso2d\_2dsolution.cu or iso2d\_1dsolution.cu for solutions using 2D and 1D thread blocks)

Use nvprof's Summary Mode to check where your implementation is spending time.

```
iso_r4_2x: 0.4557509422 s total, 0.0005638975 s/step, 1859.52 Mcells/s/step
==19313== Profiling application: ./iso2d cuda solution -x 1024 -y 1024 -i 800
==19313== Profiling result:
Time(%)
           Time
                    Calls
                              Avg
                                        Min
                                                 Max
                                                      Name
98.94% 432.51ms
                     800 540.64us 537.07us 546.56us fwd kernel
 0.85% 3.7121ms
                     804 4.6160us 1.2150us 955.81us [CUDA memcpy HtoD]
 0.21% 911.11us 1 911.11us 911.11us 911.11us
                                                      [CUDA memcpy DtoH]
==19313== APT calls:
Time(%)
           Time
                    Calls
                                        Min
                                                 Max
                              Avg
                                                      Name
82.61% 444.60ms
                     805 552.29us 9.3120us 1.2511ms
                                                      cudaMemcpy
15.56% 83.736ms
                          20.934ms 144.61us 83.285ms
                                                      cudaMalloc
                          7.9280us 7.1660us 28.247us cudaLaunch
 1.18% 6.3430ms
                   800
       1.5825ms
                             247ns
 0.29%
                     6400
                                      207ns 8.7500us
                                                      cudaSetupArgument
```

Dig deeper using nvprof metrics. We'll start by just looking at the ones we've tried in past examples.

```
$ nvprof --metrics <metric> -x 1024 -y 1024 -i 800 -t with:
```

- 1. ipc
- 2. sm\_efficiency
- 3. alu fu utilization
- 4. gld\_transactions
- 5. gst\_transactions

- 6. gld\_transactions\_per\_request
- 7. gst\_transactions\_per\_request
- 8. gld\_efficiency
- 9. gst\_efficiency

Dig deeper using nvprof metrics. We'll start by just looking at the ones we've tried in past examples.

Invocations Device "Tesla M2050 (0)	Metric Name	Metric Description	Avg
Kernel: fwd_ker			
800	<pre>sm_efficiency</pre>	Multiprocessor Activity	99.92%
800	ipc	Executed IPC	1.594381
800	gld_transactions	Global Load Transactions	19710600
800	<pre>gst_transactions</pre>	Global Store Transactions	1051120
800	<pre>gld_transactions_per_request</pre>	Global Load Transactions Per Request	1.978683
800	<pre>gst_transactions_per_request</pre>	Global Store Transactions Per Request	2.004852
800	<pre>gld_efficiency</pre>	Global Memory Load Efficiency	50.71%
800	<pre>gst_efficiency</pre>	Global Memory Store Efficiency	80.00%
800	alu_fu_utilization	Arithmetic Function Unit Utilization	High (7)

## iso2d

### What does this kernel load?

Accessed	Coalesced	Aligned?	Frequency of Access
<pre>curr[this_offset]</pre>	Yes	Maybe	A few threads
<pre>next[this_offset]</pre>	Yes	Maybe	A few threads
<pre>curr[y_pos_offset]</pre>	Yes	Maybe	A few threads
<pre>curr[y_neg_offset]</pre>	Yes	Maybe	A few threads
<pre>curr[x_pos_offset]</pre>	Yes	Maybe	A few threads
<pre>curr[x_neg_offset]</pre>	Yes	Maybe	A few threads
<pre>vsq[this_offset]</pre>	Yes	Maybe	A few threads
c_coeff[d]	No	Usually Not	All threads

### iso2d

Constant memory is optimized for broadcast access (whole warp reads same location).

Variables in CUDA constant memory must be declared statically:

```
__constant__ TYPE const_c_coeff[NUM_COEFF];
```

Can be referenced as normal arrays from CUDA kernels:

Initialized using a special-purpose cudaMemcpy API:

```
cudaMemcpyToSymbol(const c coeff, c coeff, NUM COEFF * sizeof(TYPE));
```

Try your hand at optimizing the iso2d CUDA version using constant memory.

Re-analyze the overall performance and the same profiler metrics.

A template with TODOs (along with an example solution) is available in 07\_iso2d\_cuda\_cmem.

Try your hand at optimizing the iso2d CUDA version using constant memory.

Re-analyze the overall performance and the same profiler metrics.

Version	Performance (Mcells/s/step)	Improvement
Sequential	73.35	
OpenMP	275.20	3.75
CUDA	2,708.77	9.84
+ cmem	2,812.53	1.04

-x 4096 -y 4096 -i 1000

Try your hand at optimizing the iso2d CUDA version using constant memory.

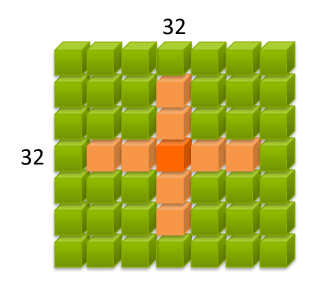
Re-analyze the overall performance and the same profiler metrics.

Invocations	Metric Name	Control Description	Avg
Device "Tesla M2050 Kernel: fwd	(0)" kernel	Multiprocessor Activity Executed IPC Global Load Transactions	
800	sm_efficiency	Multiprocessor Activity	99.90%
800	ipal	Executed IPC	1.594887
800	gld_transactions	Global Load Transactions	19714799
800	gst_transactions	Global Store Transactions	1060304
800	gld_transactions_per_request	Global Load Transactions Per Request	1.976322
800	gst_transactions_per_request	Global Store Transactions Per Request	2.016384
800	gld_efficiency	Global Memory Load Efficiency	50.60%
800	gst_efficiency	Global Memory Store Efficiency	80.00%
800	alu_fu_utilization	Arithmetic Function Unit Utilization	High (7)

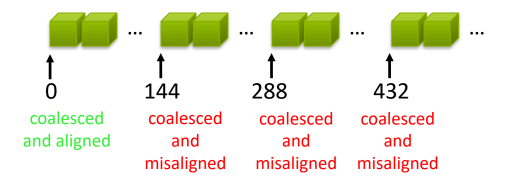
### Back to the drawing board.

Accessed	Coalesced	Aligned?	Frequency of Access
<pre>curr[this_offset]</pre>	Yes	Maybe	A few threads
<pre>next[this_offset]</pre>	Yes	Maybe	A few threads
<pre>curr[y_pos_offset]</pre>	Yes	Maybe	A few threads
<pre>curr[y_neg_offset]</pre>	Yes	Maybe	A few threads
<pre>curr[x_pos_offset]</pre>	Yes	Maybe	A few threads
<pre>curr[x_neg_offset]</pre>	Yes	Maybe	A few threads
<pre>vsq[this_offset]</pre>	Yes	Maybe	A few threads
c_coeff[d]	No	Usually Not	All threads

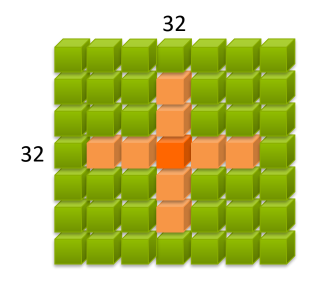
Depending on the width of the 2D grid we're applying our stencil to, there may be many accesses that are mis-aligned.



$$(2 + 32 + 2)$$
 \* sizeof(float) = 144 bytes per row

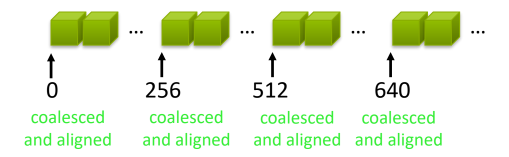


Depending on the width of the 2D grid we're applying our stencil to, there may be many accesses that are mis-aligned.



If we pad each row out to an even multiple of 128 bytes...

$$(2 + 32 + 2)$$
 \* sizeof(float) + **112** = 256 bytes per row



Depending on the width of the 2D grid we're applying our stencil to, there may be many accesses that are mis-aligned.

By padding rows in the grid to be a length evenly divisible by 128 bytes, we can improve the # of aligned accesses (thereby reducing transactions).

Try writing a version of your constant memory version of iso2d that pads all rows of curr, next, and vsq out to 128 bytes.

Re-analyze the overall performance and the same profiler metrics.

08\_iso2d\_cuda\_aligned contains a starting template and example solution.

Version	Performance (Mcells/s/step)	Improvement
Sequential	73.35	
OpenMP	275.20	3.75
CUDA	2,708.77	9.84
+ cmem	2,812.53	1.04
+ aligned	2944.26	1.05

-x 4096 -y 4096 -i 1000

Metric	Old Value	New Value
sm_efficiency	99.92%	99.90%
ipc	1.594381	1.717437
gld_transactions	19710600	14179671
gst_transactions	1051120	525216
<pre>gld_transactions_per_request</pre>	1.978683	1.423451
<pre>gst_transactions_per_request</pre>	2.004852	1.001770
gld_efficiency	50.71%	70.25%
gst_efficiency	80.00%	100.00%
alu_fu_utilization	High (7)	High (7)

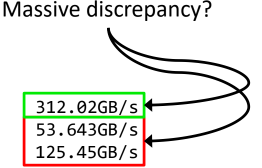
### **Hands On – nvprof Throughput Metrics**

#### Let's collect some new metrics:

- 1. gld\_throughput: Achieved throughput for global memory accesses
- 2. dram\_read\_throughput: Throughput between DRAM and L2
- 3. I2\_read\_throughput: Throughput through L2 cache

### **Hands On – nvprof Throughput Metrics**

- 1. gld\_throughput: Achieved throughput for global memory accesses
- 2. dram\_read\_throughput: Throughput between DRAM and L2
- 3. I2\_read\_throughput: Throughput through L2 cache



### iso2d

Throughput metrics suggest that we're hitting a lot in L1 cache.

Can confirm using the l1\_cache\_global\_hit\_rate metric.

```
Metric Name Metric Description Avg
Device "Tesla M2050 (0)"

Kernel: fwd_kernel

11 cache global hit rate L1 Global Hit Rate 64.03%
```

Good: we're getting lots of locality benefits.

Bad: L1 cache may be a bottleneck, losing some of our on-chip memory to shared memory.

### iso2d

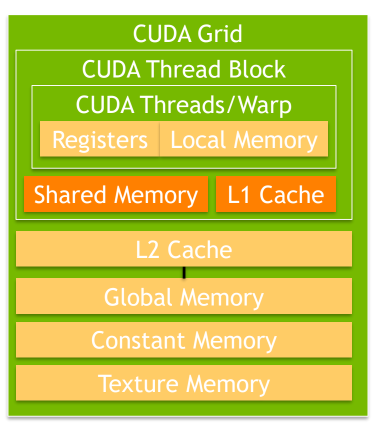
Original discussion of shared memory:

- On-chip, low latency, high bandwidth
- Same physical memory as L1 cache

Reconfigure the partitioning of on-chip memory between L1 and shared memory:

cudaError\_t cudaThreadSetCacheConfig(
 enum cudaFuncCache cacheConfig);

- cudaFuncCachePreferNone (default)
- cudaFuncCachePreferShared
- cudaFuncCachePreferL1



# Hands On – On-Chip Memory Configuration

How does cudaThreadSetCacheConfig affect the performance of iso2d?

Try changing the call to it at the start of the 08\_iso2d\_cuda\_aligned example solution (iso2d\_aligned\_solution.cu) to be either:

cudaThreadSetCacheConfig(cudaFuncCachePreferShared);

cudaThreadSetCacheConfig(cudaFuncCachePreferL1);

Then, measure any change in overall performance and L1 hit ratios.

## **Hands On – On-Chip Memory Configuration**

Version	Performance (Mcells/s/step)	Improvement
Sequential	73.35	
OpenMP	275.20	3.75
CUDA	2,708.77	9.84
+ cmem	2,812.53	1.04
+ aligned	2944.26	1.05
+ cudaFuncCachePreferShared	2943.11	1.00
+ cudaFuncCachePreferL1	3037.05	1.03

-x 4096 -y 4096 -i 1000

w/ cudaFuncCachePreferNone:64.03% L1 Global Hit Ratew/ cudaFuncCachePreferShared:64.04% L1 Global Hit Rate77.29% L1 Global Hit Rate

#### **Hands On – On-Chip Memory Configuration**

Version	Performance (Mcells/s/step)	Improvement
Sequential	73.35	
OpenMP	275.20	3.75
<b>CUDA</b> Appears to be improve		ng on-chip
+ cmem	locality <sub>2,812.53</sub>	1.04
+ aligned	2944.26	1.05
+ aligned Time to lo + cudaFuncCachePreferShared	2943.11	1.00
+ cudaFuncCachePreferL1	3037.05	1.03

-x 4096 -y 4096 -i 1000

w/ cudaFuncCachePreferNone:64.03% L1 Global Hit Ratew/ cudaFuncCachePreferShared:64.04% L1 Global Hit Rate77.29% L1 Global Hit Rate

#### **CUDA Shared Memory**

Shared memory is shared by threads in the same block.

On-SM memory

Shared memory can be allocated statically or dynamically

Statically Allocated Shared Memory

- Size is fixed at compile-time
- Can declare many statically allocated shared memory variables
- Can be declared globally or inside a device function
- Can be multi-dimensional

```
__shared__ int s_arr[256][256];
```

### **CUDA Shared Memory**

#### Dynamically Allocated Shared Memory

- Size in bytes is set at kernel launch with a third kernel launch configurable
- Can only have one dynamically allocated shared memory array per kernel
- Must be one-dimensional array

```
__global__ void kernel(...) {
    extern __shared__ int s_arr[];
    ...
}
kernel<<<nblocks, threads_per_block, shared_memory_bytes>>>(...);
```

#### **CUDA Shared Memory**

What is reused in iso2d? i.e. what can benefit from improved on-chip locality?

Take a look at using shared memory to tile curr. This is the toughest transformation we'll try today.

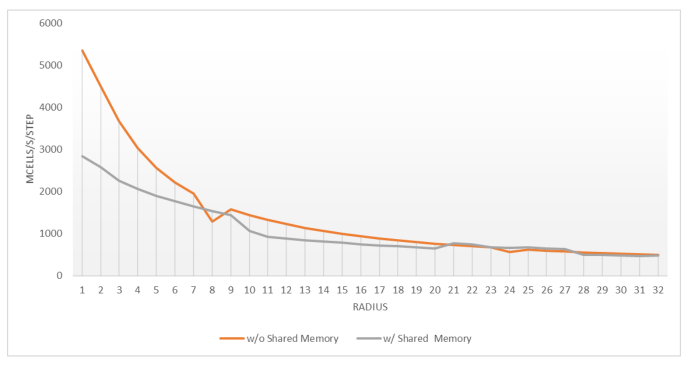
There is also a template and example solution under 09\_iso2d\_cuda\_smem/.

You can start by declaring a dynamic shared memory allocation in the kernel, and figuring out how large it needs to be from the launch:

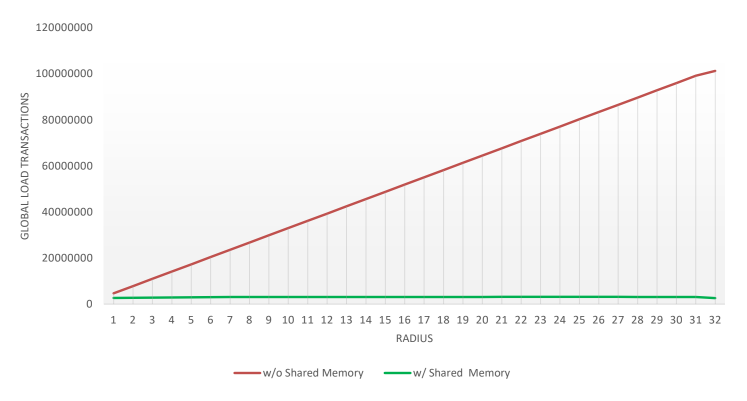
Give it a shot!

Version	Performance (Mcells/s/step)	Improvement
Sequential	73.35	
OpenMP	275.20	3.75
CUDA	2,708.77	9.84
+ cmem	2,812.53	1.04
+ aligned	2944.26	1.05
+ cudaFuncCachePreferL1	3037.05	1.03
+ smem	2063.41	0.68

Increasing radius causes more reuse, more compute to offset overheads of cache initialization => w/ shared mem is better relative to w/o shared mem.



Very effective at reducing global load transactions.



Still don't see any overall performance benefit because our manual management of shared memory is doing no better than the automatic management of L1, but incurs additional overheads (instructions executed, registers, etc).

Without shared memory, L1 Global Hit Rate = 77.29%

With shared memory, L1 Global Hit Rate = 0.13%

# iso2d Summary

Version	Performance (Mcells/s/step)	Improvement
Sequential	73.35	
OpenMP	275.20	3.75
CUDA	2,708.77	9.84
+ cmem	2,812.53	1.04
+ aligned	2944.26	1.05
+ cudaFuncCachePreferL1	3037.05	1.03
+ smem	2063.41	0.68

#### **Alternatives to CUDA**

#### **OpenCL**

A more open standard for parallel programming of a variety of processors.

• Supports GPUs, CPUs, FPGAs

Much of the OpenCL API was based on the CUDA APIs:

CUDA API	OpenCL API
cudaMalloc	clCreateBuffer
cudaFree	clReleaseMemObject
cudaMemcpyAsync	clEnqueueWriteBuffer
cudaMemcpyAsync	clEnqueReadBuffer
kernel<<<>>>();	clEnqueueNDRangeKernel

# OpenCL

Pros	Cons
Portable API, run one kernel everywhere.	More verbose, explicit API
Open source	Weaker tooling than CUDA
Not tied to a single vendor	Smaller community than CUDA, less established
Collaboration among many companies, constantly pushing the standard forward.	

#### **OpenMP Accelerators**



#### target

Create a device data environment and execute the construct on the same device.

```
#pragma omp target [clause[[,] clause],...] new-line
parallel-loop-construct | parallel-sections-construct

| C/C++ |
| Fortran |
| Somp target [clause[[,] clause],...] |
| parallel-loop-construct | parallel-sections-construct |
| Somp end target |
| Fortran |
```

#### Clauses

```
device(integer-expression)
map(list)
mapto(list)
mapfrom(list)
scratch(list)
num_threads(list)
if(scalar-expression)
```

```
sum = 0;
#pragma omp target device(acc0) map(B,C)
#pragma omp parallel for reduction(+:sum)
for (i=0; i<N; i++)
    sum += B[i] * C[i]</pre>
```

## **OpenMP Accelerators**

Pros	Cons
High-level, easy to work with	Relatively new, implementations are still coming along, community is still growing, tooling is nonexistent
Supported by a large open-source consortium	Higher abstractions mean less control over optimizations
Single parallel programming model for host and device	

#### **OpenACC**

Pragma-based (ala OpenMP) programming for GPUs, primarily supported by PGI (a.k.a. NVIDIA).

```
#pragma acc parallel loop gang deviceptr(d A, d B, d C)
    for (i = 0; i < M; i++)
#pragma acc loop worker vector
        for (j = 0; j < P; j++) {
            float sum = 0.0f;
            for (k = 0; k < N; k++)
                sum += d A[i * N + k] * d B[k * P + j];
           d_C[i * P + j] = sum;
```

# **OpenACC**

Pros	Cons
High-level, easy to work with	With OpenMP 4.0, future of OpenACC is unclear
Earliest example of pragma-based GPU programming, so likely the most stable	Higher abstractions mean less control over optimizations
Supported by NVIDIA/PGI	

### **Trilinos Project**

"The Trilinos Project is an effort to develop algorithms and enabling technologies within an object-oriented software framework for the solution of large-scale, complex multi-physics engineering and scientific problems."

Over 50 packages: <a href="https://trilinos.org/packages/">https://trilinos.org/packages/</a>

Includes packages for BLAS, Preconditioners, Linear Solvers, Nonlinear Solvers,
 Eigensolvers, Automatic Differentiation, Domain Decomposition, Partitioning, Mesh
 Manipulation, ...

Supports CUDA, OpenMP as backends

Provided free and open source by Sandia National Lab

# **Trilinos Project**

Pros	Cons
Like a super CUDA library	Long compile times due to heavy use of C++ templating
Open source and free, developed/supported by the national labs	Performance can be hit or miss

#### Kokkos, Raja

- Kokkos (Sandia), Raja (Livermore)
  - C++ APIs that expose high-level parallel operators (e.g. map, reduce, parallel-for) and use template meta-programming to map them to multiple architectures
  - Emphasize that performance bottlenecks are predominantly in memory accesses
  - Kokkos sits under Trilinos library for computational science, used in production in industry and gov't

```
Kokkos::parallel_for(numberOfAtoms,
    [=] (const size_t atomIndex) {
        atomForces[atomIndex] =
            calculateForce(data);
    }
);
```

Example of parallel-for in Kokkos. Blue shows Kokkos calls, red shows user-written kernel.

Example of parallel reduce in Kokkos.
Blue shows Kokkos calls, red shows user-written kernel

#### **GPU Acceleration of the JVM**

#### Many projects looking at JVM acceleration:

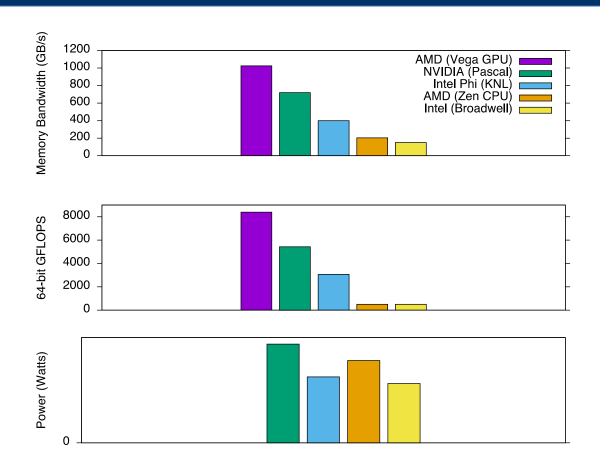
- APARAPI: <a href="https://github.com/aparapi/aparapi">https://github.com/aparapi/aparapi</a>
- Rootbeer: <a href="https://github.com/pcpratts/rootbeer1">https://github.com/pcpratts/rootbeer1</a>
- SWAT: https://github.com/agrippa/spark-swat
- IBM's J9 JVM:

http://ahayashi.blogs.rice.edu/files/2013/07/IBM Java8 GPU PACT15 cameraready-sj0tik.pdf

Pros	Cons
Broader set of GPU applications	Overheads from JIT-ing, serialization, etc may make speedup harder to achieve
High-level JVM programming languages	Most solutions are still in the research stage
Speedup relative to JVM-based apps may be > speedup relative to native apps	Little or no control over optimization

#### **Alternatives to GPUs**

#### FPGA, KNL, x86



Is your hardware holding you back, or are you holding back your hardware?

GPUs have higher hardware peaks, but also have performance characteristics that you actually stand a chance of reasoning about.

## **Summary**

## **Topics Not Covered**

Multi-GPU CUDA Programming

**Shared Bank Conflicts** 

**CUDA-Aware MPI** 

**CUDA** libraries

Advanced Stream Usage

Dynamic Parallelism

Advanced Instruction Optimization

Warp Shuffle Instructions

**Unified Memory** 

cuda-gdb

**Texture Memory** 

• • •

## **Topics Covered**

**GPU Architecture** 

nvprof

**CUDA Execution Model** 

Kernel Synchronization

**CUDA Asynchrony** 

**CUDA Memory Hierarchy** 

**Optimizing CUDA Data Access** 

Hands On with a Non-Trivial CUDA Application

**CUDA Atomic Operations** 

Alternatives to CUDA

#### **Additional Resources**

I'm not condoning theft, but...

If you happen to google "Professional CUDA C Programming PDF", the first result may or may not be a bootleg copy of my textbook, on which these slides are based.

Contact:
Max Grossman
max@7pod.tech

