





# An Introduction to HPC and Scientific Computing

Lecture seven: An introduction to GPUs and how to use them.

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# **Learning outcomes**

### In this lecture you will learn about:

- Different hardware generations of GPUs.
- How GPUs execute code.
- A more detailed look at the differences between GPUs and CPUs
- How GPUs hide memory latencies.
- The GPU software environment and useful tools.
- Finally we will cover the basics of how to use GPU libraries in your codes.



# **Hardware overview**





# **CUDA** capable hardware

NVIDIA has produced eight generations of GPU architectures that are capable of executing CUDA code.

Each generation has a different compute capability. Typically as the compute capability increases the functionality of the card increases.

Over the eight generations memory capacity has increased by twenty times, bandwidth to main memory by twenty times, but compute throughput by 55 times.

Hardware generation	Card	# CUDA cores	Main memory (GB)	Bandwidth (GB/s)	Peak compute performance (Tflops)	CUDA compute capability
Tesla	C1060	240	4	103	0.93	1.3
Fermi	M2090	512	6	178	1.3	2
Kepler	K40	2880	12	288	4.2	3.5
Maxwell	M40	3072	12	288	5.8	5.2
Pascal	P100	3584	16	732	8.1	6
Volta	V100	5120	32	900	14	7
Ampere	A100	6912	40/80	1555	19.5	8
Hopper	H100	14592	80	2039	51.2	9

https://docs.nvidia.com/cuda/cuda-c-programming-guide/index.html#compute-capabilities





# **CUDA** capable hardware

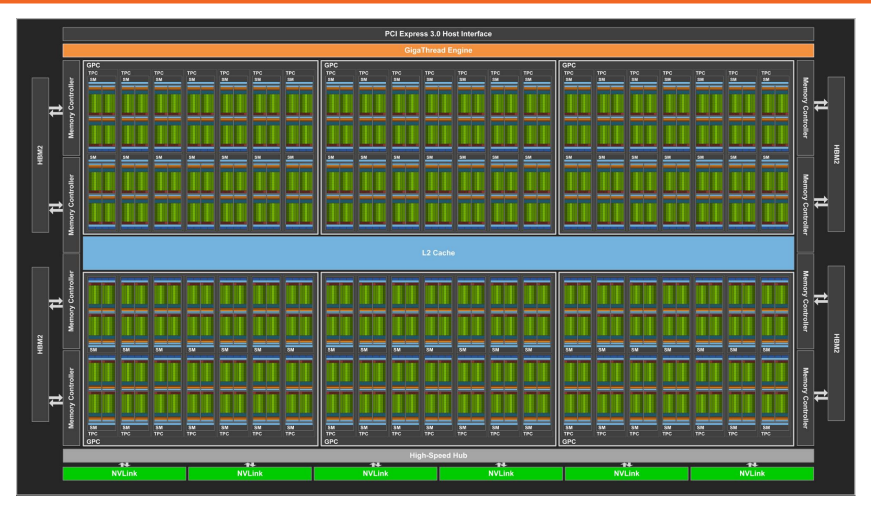
Each generation has slightly different technical specifications, for example the maximum number of thread blocks per SM (multiprocessor). These specifications are often what limits the eventual performance of your GPU code.

							Compute Capability						
Technical Specifications	3	3.2	2	3.5	3.7	5	5.2	5.3	6	6.1	6.2	7	
Maximum dimensionality of grid of thread blocks		3											
Maximum x-dimension of a grid of thread blocks		231-1											
Maximum y- or z-dimension of a grid of thread blocks		65535											
Maximum dimensionality of thread block		3											
Maximum x- or y-dimension of a block		1024											
Maximum z-dimension of a block		64											
Maximum number of threads per block		1024											
Warp size		32											
Maximum number of resident blocks per multiprocessor		16 32											
Maximum number of resident warps per multiprocessor		64											
Maximum number of resident threads per multiprocessor		2048											
Number of 32-bit registers per multiprocessor		64	K		128 K				64 K				
Maximum number of 32-bit registers per thread block	64 K	32	K		64	4 K		32 K	64 K		32 K	64 K	
Maximum number of 32-bit registers per thread		255											
Maximum amount of shared memory per multiprocessor		48 k	B		112 KB	64 KB	96 KB	64	KB	96 KB	64 KB	96 KB	
Maximum amount of shared memory per thread block		48 KB 96 KI											
Number of shared memory banks		32											





# **NVIDIA Volta V100**





# **NVIDIA Volta V100 – Streaming Multiprocessor (SM)**

A V100 has 80 SMs (see right).

The GV100 SM incorporates 64 FP32 cores and 32 FP64 cores per SM.

The SM is partitioned into four processing blocks, each with 16 FP32 Cores, 8 FP64 Cores, 16 INT32 Cores, a new L0 instruction cache, one warp scheduler, one dispatch unit, and a 64 KB Register File.







### **NVIDIA Volta V100 - SM**

Lets look at this in more detail

We've talked about a Streaming Multiprocessor, this is a collection of hardware that our thread block executes on.

A FP32 core is the execution unit that performs single precision floating point arithmetic (floats).

A FP64 core performs double precision arithmetic (doubles).







### **NVIDIA Volta V100 - SM**

Lets look at this in more detail

A INT32 Core performs integer arithmetic

The warp scheduler selects which warp (group of 32 threads) to send to which execution units.

64 KB Register File – lots of transistors used for vary fast memory.

128KB of configurable L1 (data) cache or shared memory

Shared memory is a used manged cache.

LD/ST units load and store data from cores

SFU – special function units compute things like transcendentals.







### **Code execution model**

### Software

Hardware

A thread executes on a core.

Thread

3

FP32

Core

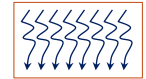
A group of threads (thread block) is executed on a SM.

Thread blocks don't migrate between SMs.

Several concurrent thread blocks can reside on a SM.

In CUDA this is a kernel launch.

Thread Block





SM

A group of thread blocks form a grid and a grid runs on the device.





**Device** 

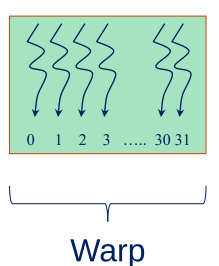




# Warps

Code executes on a GPU in groups of threads, the threads are naturally grouped into lots of 32, this is called a warp.

The reason that a warp has 32 elements is that the SIMD (recall lecture one) units on a GPU have 32 "lanes".







# Warps

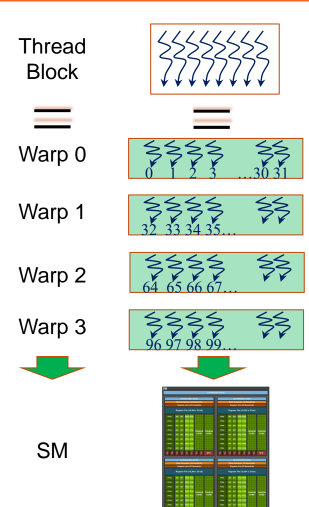
Thread blocks are formed from warps.

The warp is executed in parallel on the SM.

By this we mean that everything that happens within a warp is lock-step.

So the same operation (instruction) in thread 2, 3, 8 occurs in thread 7, 1, 4, 11... (from 0 to 31) at the same time.

So we have a Single Instruction Multiple Thread (SIMT) architecture.

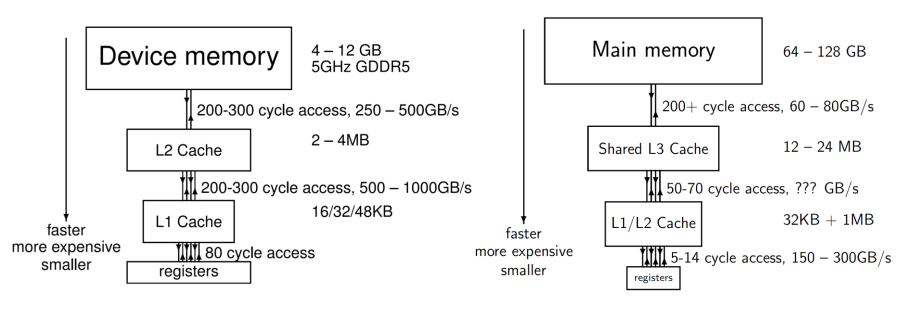






# **Memory architecture**

In lecture one we looked a the memory architecture of CPUs. Lets compare this to that of GPUs. We see that although GPUs have greater bandwidth, they have less cache and (importantly) greater latencies (the time taken to get data from memory to the processing cores).







Images courtesy of Mike Giles





CPUs are optimised for low latency access to cached data sets.

Lots of transistors spent on control logic for outof-order (OoO) and speculative execution.

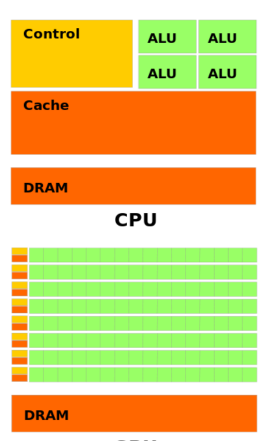
CPUs support 10's of threads.

GPUs are optimised for data parallel high throughput computation.

By having many thousands of memory requests *in flight* at any one time the larger latencies can be hidden.

Lots of transistors dedicated to computation.

GPUs support 1000's of threads.









Lets think about our GPU memory pipe as a conveyor belt.

Every two seconds a container becomes available in which we can put a single piece of data.

Bandwidth is one container per two seconds = 0.5 containers per second

Our conveyor belt has 8 containers on it at any one time.

Latency is 8 containers / 0.5 containers per second = 16 seconds







A CPU has only 10's of threads to make memory requests to hide it's memory latencies.

However a GPU has thousands of threads which can all make memory requests and so can hide larger latencies.

A GPU hides the memory latencies of one warp with the computation of another warp





A warp is executed on an SM, up to the point it stalls on a memory request.

The memory request is made and the warp removed.

The warp scheduler then looks to see what data has arrived through the memory pipe and then schedules the warp associated with that data for execution.

Eventually the data arrives for our stalled warp and then this is scheduled for execution.







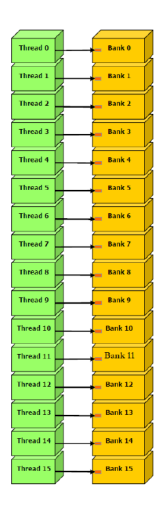
# **Memory coalescing**

Global memory access happens in transactions of 32 or 128 bytes (cache line).

So to achieve peak memory bandwidth we need to use all of the data stored in the cache line.

Coalesced memory access
A warp of threads access adjacent data
(see right) in a cache line. In the best
case this results in one memory
transaction (best bandwidth).

Uncoalesced memory access
A warp of threads access scattered data all in different cache lines.
This would result in 32 different memory transactions (giving poor bandwidth).







### Software overview

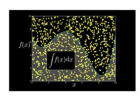
NVIDIA provides a rich ecosystem of software tools that allow you to easily utilise GPUs in your project.

During this course we will focus on the CUDA programming language, we will also look at some of the NVIDIA libraries that can be used to make programming GPUs easier.





GPU-accelerated library for Fast Fourier
Transforms



cuRAND

GPU-accelerated random number generation (RNG)



#### **CUDA Toolkit**

Provides a comprehensive environment for C/C++ developers building GPU-accelerated applications.



cuBLAS

GPU-accelerated standard BLAS library





### Libraries

### Deep Learning Libraries



GPU-accelerated library of primitives for deep neural networks



GPU-accelerated neural network inference library for building deep learning applications



Advanced GPU-accelerated video inference library

### Signal, Image and Video Libraries



cuFFT

GPU-accelerated library for Fast Fourier
Transforms



**NVIDIA Performance Primitives** 

GPU-accelerated library for image and signal processing



**NVIDIA Codec SDK** 

High-performance APIs and tools for hardware accelerated video encode and decode





### Libraries

### Linear Algebra and Math Libraries



cuBLAS

GPU-accelerated standard BLAS library



**CUDA Math Library** 

GPU-accelerated standard mathematical function library



**cuSPARSE** 

GPU-accelerated BLAS for sparse matrices



cuRAND



**cuSOLVER** 

Dense and sparse direct solvers for Computer Vision, CFD, Computational Chemistry, and Linear Optimization applications



**AmgX** 

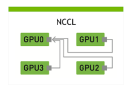
GPU accelerated linear solvers for simulations and implicit unstructured methods





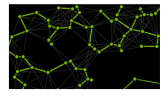
### Libraries

### Parallel Algorithm Libraries



**NCCL** 

Collective Communications Library for scaling apps across multiple GPUs and nodes



nvGRAPH

GPU-accelerated library for graph analytics



**Thrust** 

GPU-accelerated library of parallel algorithms and data structures





### Languages



#### **CUDA Toolkit**

Provides a comprehensive environment for C/C++ developers building GPU-accelerated applications.



#### **CUDA FORTRAN**

Enjoy GPU acceleration directly from your Fortran program using CUDA Fortran from The Portland Group.



#### OpenCL<sup>TM</sup>

OpenCL is a low-level API for GPU computing that can run on CUDA-powered GPUs.



#### OpenACC

Directives for parallel computing, is a new open parallel programming standard designed to enable all scientific and technical programmers.



#### Anaconda Accelerate

Enables acceleration on your GPU or multi-core processor using Python.



### PGI Accelerator Fortran and C Compilers

Accelerate applications on GPU platforms by adding compiler directives to existing code.



#### **PyCUDA**

Gives you access to CUDA fuctionality from your Python code.



### The PGI CUDA C/C++ compiler for x86

Compile and optimize their CUDA applications to run on x86-based workstations, servers and clusters.



#### Altimesh Hybridizer™

An advanced productivity tool that generates vectorized C++ (AVX) and CUDA C code from .NET assemblies (MSIL) or Java archives (bytecode)



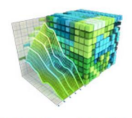
#### Alea GPU

This is a novel approach to develop GPU applications on .NET, combining the CUDA with Microsoft's F#.





### **Tools**



#### **NVIDIA Visual Profiler**

This is a cross-platform performance profiling tool that delivers developers vital feedback for optimizing CUDA C/C++ applications. First introduced in 2008, Visual Profiler supports all CUDA capable NVIDIA GPUs shipped since 2006 on Linux, Mac OS X, and Windows.



#### NVIDIA® Nsight™

The ultimate development platform for heterogeneous computing. Work with powerful debugging and profiling tools, optimize the performance of your CPU and GPU code. Find out about the Ecilipse Edition and the graphics debugging enabled Visual Studio Edition.



#### **CUDA-GDB**

Delivers a seamless debugging experience that allows you to debug both the CPU and GPU portions of your application simultaneously. Use CUDA-GDB on Linux or MacOS, from the command line, DDD or EMACS.



#### **CUDA-MEMCHECK**

Identifies memory access errors in your GPU code and allows you to locate and resolve problems quickly. CUDA-MEMCHECK also reports runtime execution errors, identifying situations that could result in an "unspecified launch failure" error while your application is running.





### **CUDA**

CUDA (Compute Unified Device Architecture) provides a set of programming extensions based on the C/C++ family of languages.

If you have a basic understanding of C and understand the concept of threads and SIMD execution, then CUDA is easy to pick up.

An introduction to the CUDA programming language will be given in lecture eight.



#### **CUDA Toolkit**

Provides a comprehensive environment for C/C++ developers building GPU-accelerated applications.

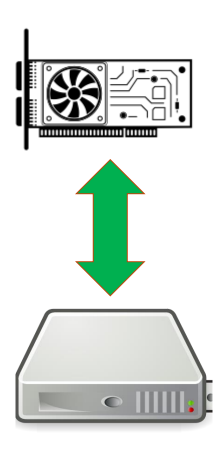


# **Working with GPUs**

Recall for our first lecture (and earlier in these slides), a GPU is attached to the computer by a PCIe bus.

It also has it's own memory (for example a V100 has 16GB of HBM2 memory). So the GPU has a separate address space.

This means that for us to work with the GPU we need to allocate memory on the card (device) and then transfer data to and from the device.







# **Working with GPUs**

To allocate memory on the device CUDA provides a function just like malloc() called cudaMalloc()

To free memory CUDA provides a function like free() called cudaFree()

Examples are given in the code snippit on the right.

```
// Allocate pointers for host and device memory
float *h input, *h output;
float *d input, *d output;
// malloc() host memory (this is in your RAM)
h input = (float*) malloc(mem size);
h output = (float*) malloc(mem size);
// allocate device memory input and output arrays
cudaMalloc((void**)&d input, mem size);
cudaMalloc((void**)&d output, mem size);
// Do something here!
// cleanup memory
free(h input);
free(h output);
cudaFree(d input);
cudaFree(d output);
```



# **Working with GPUs**

To transfer data to and from the GPU CUDA provides a function to do this called cudaMemcpy ()

As arguments cudaMemcpy() takes:

cudaMemcpy(destination pointer, origin pointer, size of memory to move, direction)

#### **Examples below**

```
// Copy host memory to device input array
cudaMemcpy(d_input, h_input, mem_size, cudaMemcpyHostToDevice);

// Do something on the GPU

// copy result from device to host
cudaMemcpy(h_output, d_output, mem_size, cudaMemcpyDeviceToHost);
```





### An example - cuRAND

The example on the right shows how to generate a distribution of random normal number using cuRAND.

This code could replace

// Do something here

In the previous slides. Combining this will cudaMemcpy() allows you to start building a GPU program.

```
// random number generation
#include <cuda.h>
#include <curand.h>

// Declare variable
curandGenerator_t gen;

// Create random number generator
curandCreateGenerator(&gen, CURAND_RNG_PSEUDO_DEFAULT) );

// Set the generator options
curandSetPseudoRandomGeneratorSeed(gen, 1234ULL) );

// Generate the randoms
curandGenerateNormal(gen, d_input, NUM_ELS, 0.0f, 1.0f) );
```

https://docs.nvidia.com/cuda/curand/introduction.html#introduction



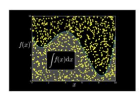
### **Practical 5**

In practical 5 you will work with different codes to familiarise yourselves with cuRAND, cuFFT and cuBLAS.





GPU-accelerated library for Fast Fourier
Transforms



cuRAND

GPU-accelerated random number generation [RNG]



#### **CUDA Toolkit**

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#### cuBLAS

GPU-accelerated standard BLAS library





### What have we learnt?

In this lecture you have learnt about different hardware generations of GPUs and some of there differences.

We've looked at how GPUs execute code.

We have looked at the differences between GPUs and CPUs in greater detail, we have also looked at how GPUs hide memory latencies.

We've looked at the GPU software environment and useful tools.

Finally we have covered the basics of how to use GPU libraries in your codes.







# **Further reading**

http://docs.nvidia.com/cuda/curand/index.html

https://developer.nvidia.com/curand

http://docs.nvidia.com/cuda/cufft/index.html

https://developer.nvidia.com/cufft

http://docs.nvidia.com/cuda/cublas/index.html

https://developer.nvidia.com/cublas

https://docs.nvidia.com/cuda/

https://developer.nvidia.com/about-cuda



### In the next lecture...

In the next lecture you will learn about the CUDA programming language and how to write a CUDA code.



