

Lecture 23 — GPU Programming Continued

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We've learned about how a kernel works and a bit about how to write one.

The next part is the host code.

Now, fortunately, we don't have to write the whole program in C++ or C, even though the kernel has to be written in the CUDA variant.

We're going to use the Rustacuda library!

```
#[macro_use]
extern crate rustacuda;

use rustacuda::prelude::*;
use std::error::Error;
use std::ffi::CString;

fn main() -> Result<(), Box<dyn Error>> {
    // Set up the context, load the module, and create a stream to run kernels in.
    rustacuda::init(CudaFlags::empty())?;
    let device = Device::get_device(0)?;
    let _ctx = Context::create_and_push(ContextFlags::MAP_HOST | ContextFlags::SCHED_AUTO, device)?;

    let ptx = CString::new(include_str!("../resources/add.ptx"))?;
    let module = Module::load_from_string(&ptx)?;
    let stream = Stream::new(StreamFlags::NON_BLOCKING, None)?;

    // Create buffers for data
    let mut in_x = DeviceBuffer::from_slice(&[1.0 f32; 10])?;
    let mut in_y = DeviceBuffer::from_slice(&[2.0 f32; 10])?;
    let mut out_1 = DeviceBuffer::from_slice(&[0.0 f32; 10])?;
    let mut out_2 = DeviceBuffer::from_slice(&[0.0 f32; 10])?;
```

Rust Host Code Example

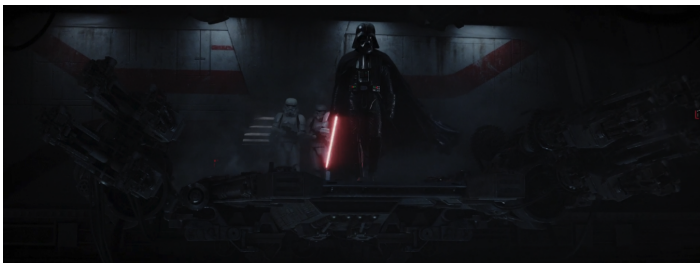
```
// This kernel adds each element in 'in_x' and 'in_y' and writes the  
    result into 'out'.  
unsafe {  
    // Launch the kernel with one block of one thread, no dynamic shared  
        memory on 'stream'.  
    let result = launch!(module.sum<<<1, 1, 0, stream>>>(  
        in_x.as_device_ptr(),  
        in_y.as_device_ptr(),  
        out_1.as_device_ptr(),  
        out_1.len()  
    ));  
    result?;  
}  
  
// Kernel launches are asynchronous, so we wait for the kernels to finish  
    executing.  
stream.synchronize()?;
```



```
// Copy the results back to host memory
let mut out_host = [0.0f32; 20];
out_1.copy_to(&mut out_host[0..10])?;
out_2.copy_to(&mut out_host[10..20])?;

for x in out_host.iter() {
    assert_eq!(3.0 as u32, *x as u32);
}

println!("Launched kernel successfully.");
Ok(())
}
```



```
extern "C" __constant__ int my_constant = 314;

extern "C" __global__ void sum(const float* x, const float* y, float* out, int
    count) {
    for (int i = blockIdx.x * blockDim.x + threadIdx.x; i < count; i +=
        blockDim.x * gridDim.x) {
        out[i] = x[i] + y[i];
    }
}
```



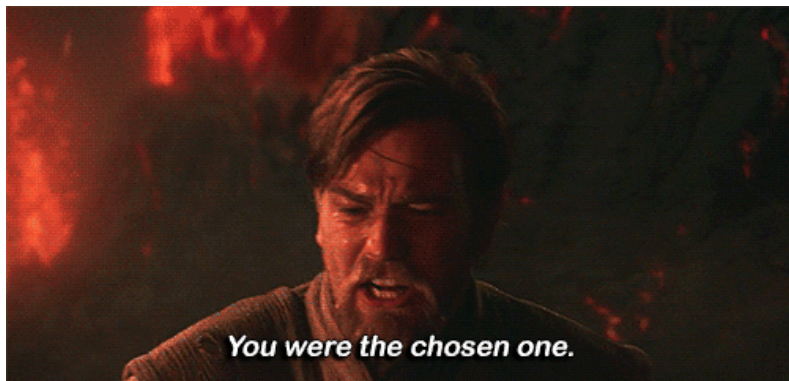
Let's take a look at the N-Body Problem code in the repo.

That's great, but how much does it speed up?

I ran this on ec2esla1.

With 100 000 points:

- nbody (sequential, no approximations): 40.3 seconds
- nbody-parallel (parallel, no approximations): 5.3 seconds
- nbody-cuda (parallel, no approximations): 9.5 seconds

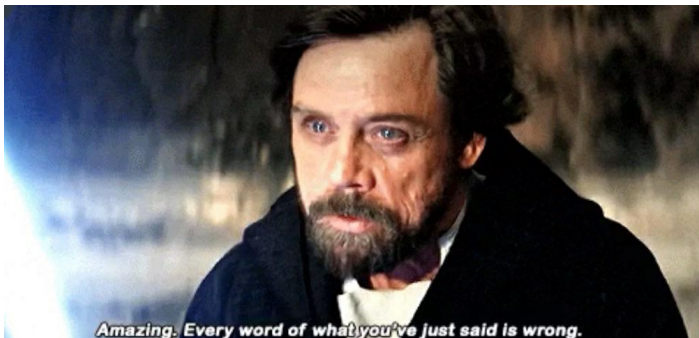


That's worse than the CPU version.

Kylo Ren is very Meme-Able

Theory: 100 000 points is not enough to overcome the overhead of setup and transferring data to and from the device.

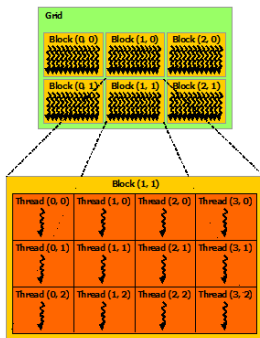




This turned out to be incorrect.

Most of the time was going to the kernel execution of the `calculate_forces` – as expected?

This Was Important



The documentation isn't super great about how grids and blocks work.

A lot of the guidance on the internet says is “experiment and try”.

The initial approach that I wrote had each work-item be its own grid.

That's inefficient, because we're not taking advantage of all the hardware that's available (the warp hardware).

The advice that I can find says the number of threads per block should always be a multiple of 32 with a maximum of 512 (or perhaps 1024).

The second guidance I can find is that numbers like 256 and 128 are good ones to start with, and you can tweak it as you need.

Then you have to adjust the grid size: divide the number of points by the threads per block to give the number of blocks.

```
let result = launch!(module.calculate_forces <<<(NUM_POINTS/256) + 1,  
    256, 0, stream>>>(points.as_device_ptr(),  
    accel.as_device_ptr(),  
    points.len()  
));  
result?;
```

I did have to add a +1 to the number of blocks, because 100 000 does not divide evenly by 256.

But just running it as-is didn't work (and led to the kernel crashing. Why?

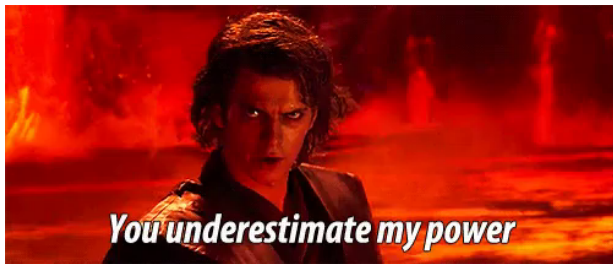
Because the indexing strategy that I used contained only the reference to the block index `blockIdx.x`.

That's fine in the scenario where every work-item gets its own block, but that's no longer the case now: 256 work-items (points) now share each block.

```
extern "C" __global__ void calculate_forces(const float4* positions, float3*
    accelerations, int num_points) {
    int idx = threadIdx.x + blockIdx.x * blockDim.x;
    if (idx >= num_points) {
        return;
    }
    float4 current = positions[idx];
    float3 acc = accelerations[idx];

    for (int i = 0; i < num_points; i++) {
        body_body_interaction(current, positions[i], &acc);
    }
    accelerations[idx] = acc;
}
```

With 100 000 points:



- nbody (sequential, no approximations): 40.3 seconds
- nbody-parallel (parallel, no approximations): 5.3 seconds
- nbody-cuda (parallel, no approximations): 9.5 seconds
- **nbody-cuda-grid (parallel, no approx., grid): 1.65 seconds**

Now that's a lot better!