ECE 459: Programming for Performance Lab 3 — Using the GPU ¹

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In this lab, you'll learn how to improve performance of a convolutional neural network (CNN) by using the GPU to do some of the work.

Learning Objectives

- Become familiar with running code on the GPU
- · Learn how to use the CUDA interface

Lab video. https://www.youtube.com/watch?v=s3hCPNnUIcc

Background

GPUs were originally designed to accelerate computer graphics (hence the name, Graphics Processing Unit), but their architecture makes them suitable for a wide range of tasks that can be parallelized. There are two major programming interfaces between the CPU and the GPU—OpenCL and CUDA. While OpenCL is more portable and runs across a wide range of GPUs, CUDA usually offers better performance and is very widely supported. It's also easier to use. The biggest limitation of CUDA is that it only runs on Nvidia GPUs.

CUDA

The typical CUDA workflow consists of having the CPU initialize the GPU, copy data into the GPU memory, invoke the code (a "kernel") on the GPU, and copy the results from the GPU back into CPU memory. In this lab, the CPU portion of the workflow is handled by a library of Rust bindings called RustaCUDA. The GPU kernel is programmed in CUDA-C, a language similar to C++ with none of Rust's safety guarantees. As such, calling a CUDA kernel from Rust is an unsafe operation and you must carefully coordinate between your Rust code and CUDA-C code to avoid undefined behaviour.

You can find more information about the CUDA-C language here:

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

Here's a resource we pointed at last year:

http://users.wfu.edu/choss/CUDA/docs/Lecture%205.pdf

Here's another resource about parallel reductions (dividing and conqueing) in CUDA: https://developer.download.nvidia.com/assets/cuda/files/reduction.pdf

And just for fun: https://www.youtube.com/watch?v=-P28LKWTzrI

The CNN

A Convolutional Neural Network (CNN) is a machine learning algorithm that takes an image and passes it through multiple layers for processing, producing a single vector of values as output. In this lab, we'll be using a simplified version of a CNN that consists of only three layers. The inputs to the CNN are the 100×100 matrices that represent images.

¹v1, 17Feb22

Mathematically, our CNN network is transforming a 100×100 input matrix into a 10-element vector by performing a bunch of matrix operations on the input. The input typically represents an image, and the output represents probabilities of classifications. For example, if we're using the CNN classifying images of numbers, then each element of the output vector will represent the probability of the image being a given number. In the real world, CNN networks are used to make these types of classifications, but for the purposes of the lab, you can think of it as a bunch of number crunching that'd be much more suited for the GPU then the CPU.

The first layer of the CNN is a Convolution layer, consisting of 10 neurons, each containing a 5×5 matrix called a filter. Each neuron divides the input image into 5×5 sections and performs a dot product between its filter and each section of the input, producing a 20×20 output matrix of products.

The second layer is a Rectified Linear Unit (ReLU) layer, which sets any negative values in the Convolution layer output to zero.

The third layer is the Output layer, consisting of 10 neurons, each containing a 4000×1 vector called weights. Each Output neuron flattens all the matrices' output from the previous layer and concatenates them, forming a 4000×1 vector that it multiplies with its weights via a dot product. Since each Output neuron produces a single value, the entire Output layer produces an output vector of 10 values.

The Software

Your program reads in a CNN description and a number of 100×100 matrices (representing images), and runs the neural network by feeding the matrices through the CNN. The program writes its results to an output file, and reports the elapsed number of microseconds. The output files contain the output vectors for each input matrix.

All the data files (input, output and CNN descriptions) are in CSV (Comma-Separated Values) format, which is basically just a text file where every line contains values separated by commas (just as the name promised!).

We provide you with some starter code, including a main.rs file (which you will not need to modify), a cpu.rs file containing the CPU-based implementation, and a cuda.rs file containing a skeleton for the GPU-based implementation. There is also a file kernel/kernel.cu, which contains a skeleton for the code that will run on the GPU. There is also a build.rs file that gets run automatically at build time to compile the kernel.cu file into a kernel.ptx file which gets downloaded to the GPU. Once again, you can change the starter code, but don't change what the marker sees when running your code.

Running things for real. To build the code that runs on the GPU, you must use the correct version of the gcc compiler. Here's a command you can use to set that up:

```
nvcc --compiler-bindir /usr/bin/gcc-6 -ptx nbody.cu
```

There's a bash script in the kernel directory, setgcc, that does this for you.

The command line for running your program is as follows:

```
cargo run --release -- <mode> <cnn_file> <input_file> <output_file>
```

where <mode> is either "cpu" to run the CPU-based implementation or "cuda" to run the GPU implementation. All the files are pathnames. You would typically use the following commands:

```
cargo run --release -- cpu input/cnn.csv input/in.csv output/out.csv
cargo run --release -- cuda input/cnn.csv input/in.csv output/out_cuda.csv
```

The program outputs the time spent doing "actual work", i.e. converting the input matrices to the output vectors. This measurement does not include I/O or the overhead of initializing the GPU. As such, the time should be lower for the CUDA version than the CPU version. If you're curious about what's slow, you can put nvprof in front of cargo run.

Generating Inputs and Checking Results

In addition to the starter code, we provide two Python scripts. One (generate.py) is used to generate the input and CNN files and write them into the input directory as in.csv and cnn.csv. The other script (compare.py) is used to compare the two files in the output directory (out.csv, which is the output of the CPU version, and out_cuda.csv which is the output of the GPU version). If your CUDA-based implementation is correct, compare.py should not report any differences between those two output files. Note that the output files will not be perfectly identical, which is why we use a script to do the comparison.

The files generated by generate.py and the files compared by compare.py have fixed names, so it's recommended that you use the same names on the command lines that you use to run the application.

The scripts are written in Python 3, so you can run them using the python3 command (e.g. python3 compare.py).

Rubric

Implement your code on the GPU (40 marks) Your code needs to build correctly and run on the GPU.

Produce correct results (40 marks) The compare.py script should not report any discrepancies.

Report (20 marks)

- 8 marks for a clear discussion of how your kernel works
- 8 marks for a clear discussion of how cuda.rs works with the kernel
- · 4 marks for clarity

Clarifications

Can you clearly tell me which lectures are relevant to this lab? Lectures 22 and 23.

nvcc: command not found. Make sure you're on an ecetesla machine (or maybe eceubuntu4).

I'm not close enough, says compare.py. Are you using doubles to represent numbers?

```
I'm confused about the syntax for <<< ... >>>. Check out https://docs.nvidia.com/cuda/cuda-c-programming-guide/index.html#execution-configuration.
```

The arguments are gridsize, blocksize, shared mem, stream.

The third number (0) is the shared parameter. This defines the size of dynamic shared memory for the stream, but we don't need it here. The Rustacuda documentation (and/or source code!) can give you some guidance about its use if you have a use for it.

The second number (256) is the block size. The first parameter is the grid size, not the number of grids. The grid size tells us how many blocks are in there, and its number of points divided by 256, plus 1 just in case it does not divide evenly (which for something like 100k points, it does not). So for 100k points, it's 391 blocks.

We don't use the grid dimension because it's not really necessary in the calculation. gridDim.x would tell us the number of blocks in the grid in the x dimension, but that is not a useful figure in the calculation so we don't ask for it.

***. In Rust, multi-dimensional arrays are represented as a large 1D array, so your CUDA array representation has to match it. I would pass in the ConvLayer pointer as const double*. Since the dimensions of the convolution array are (CONV_LAYER_SIZE, FILTER_DIM, FILTER_DIM), a 3D array access of conv_layer[i][j][k] should be written as conv_layer[i*FILTER_DIM*FILTER_DIM + j*FILTER_DIM + k] in your kernel.

Weird compile error. "I try to compile let result_conv = launch!(self.module.convolution_layer... and it doesn't work, so I'm sad."

Use a local variable to store self.module and launch that. For more details: https://docs.rs/rustacuda/0.1.2/rustacuda/macro.launch.html.

Weird nondeterministic behaviour that's changed by adding an unused variable in the kernel. Check your kernel for any out of bounds accesses on your buffers. Also, to reiterate, make sure you're using "double" as your floating point data type in the kernel. Others have reported that moving their declaration of the input device box above the stream and module declarations makes things more deterministic:

```
let mut input_matrix = DeviceBox::new(input).unwrap(); // <-- first
let mut layer2_output = DeviceBuffer::from_slice(&[0.0f64; 4000]).unwrap();</pre>
```

Go figure.

Other weird behaviour. Make sure you're synchronizing at the appropriate times (which may be fewer than you think). Word to the wise²: "CUDA operations issued to the same stream always serialize." That is, you only need to call synchronize() once per stream and weird things may happen if you call it more than once.

I wish I could use a debugger. We had mixed reports from last year. People said print statements worked. You can use 'cuda-gdb' instead of the default 'rust-gdb' too.

I'm printing things but it's not quite right. Be sure to use %.2f to print a double (sorry).

Multiple kernels? Multiple calls to the kernel? Yeah, go ahead.

Hey, that could be slow. Does performance actually count? Again, no (wait for Lab 4); just use the GPU in a sufficiently parallel way. But we may be sad. Unless you're not actually using the GPU. In which case you may then be sad. To expand on that:

You're expected to utilize the GPU's parallelism to a reasonable degree, so don't put everything on the same thread and make sure the GPU solution is faster than the CPU solution. Consider a divide-and-conquer strategy for summing elements, since that's faster than atomicAdd. I'll use dot product as an example.

Let's say you need to do a dot product between vectors A and B, which are both 4000 elements long. The first step is to calculate the element-wise product C of A and B, which is easily parallelizable across 4000 threads (thread n calculates C[n] = A[n]*B[n]). The second step is summing C, which is the more complicated part. To do this

 $^{^2}$ https://forums.developer.nvidia.com/t/cudadevicesynchronize-needed-between-kernel-launch-and-cudamemcpy/44494

we split C into 200 equal segments with 20 elements each, and assign each segment to a different thread (200 threads in total). Each thread sums its segment (20 elements), so the 200 threads produce another vector D of 200 elements where sum(D) = sum(C). We can repeat the last step to reduce D to a 10-element vector, which can then be summed to the dot product.

An analogy would be if you wanted to collect and accumulate all the garbage across N households efficiently. First you'd send N workers to collect the garbage of all households (that's the easily parallelizable part). Then you dump the N bags of garbage into M landfills, where M < N. Finally, you assemble the garbage from all M landfills into a single garbage processing plant, completing the procedure.

Are constants constant? We won't change the constants in the CNN file, i.e. you can hardcode them if you want.

Do we commit compiled files to git? (kernel.ptx and output/out_cuda.csv) Doesn't matter, we'll regenerate.