

Final Report: Neural Networks on CUDA

Goal of the final project is to implement a neural network with parallel matrix-matrix operations across four GPUs. Our neural network will be used to identify images of hand-written digits from the MNIST dataset.

Section 1: Accelerated Matrix Multiplication Idea: parallelize our generalized in-place matrix multiplication (GEMM) $C = \alpha * A * B + \beta * C$ across cores of one GPU. We will be able to use this to execute multiple steps in the neural network's feed forward and back propagation operations.

- **Algorithm #1:** Our first (naive) implementation is correct but very inefficient. We ask each thread to compute a single value of the output matrix C. This requires each thread to fetch from global memory an entire row of matrix A and column from matrix B, resulting in a lot of repeated global memory requests for the same information. Note, Algorithm 1 was used for part 1 of the project.
- **Algorithm #2:** Our second implementation makes use of shared memory, which affords us increased bandwidth. In particular, algorithm 2 loads a 32x32 block from matrix A and matrix B into shared memory before performing a partial update on the corresponding 32x32 block of output matrix C. Each thread still only performs an update for a single cell in the output matrix.
- **Algorithm #3:** Our third implementation uses a specific thread block of size 4x16 to compute a partial update of a 64x16 block of output matrix C. Each thread in this case updates one 1x16 row of this 64x16 block, which reduces latency and increases overall arithmetic intensity compared to algorithms 1 and 2. This implementation makes use of shared memory as well as local (register) memory.

In Figure 1, we can see that our algorithms progressively improve towards (but do not match) the reference cublas solution for FP32 numbers. However, for double precision FP64 numbers, algorithms 2, 3 and the reference all perform similarly, with algorithm 3 actually surpassing the reference. This is likely because the cublas solution is optimized for FP32 given most use cases (e.g. neural network training) do not require double precision.

Figure 2 shows a heat map with speed tests for different GEMM matrix dimension combinations (assuming FP32). Since algorithm 1 < algorithm 2 < algorithm 3 < reference for all settings, I have consistently implemented algorithm 3 throughout my neural network kernels. It is interesting to note that this relative differences shift quite a bit, but never enough to justify a change of algorithm for specific settings. Further analysis could take into account other input parameter options such as size of thread block, number of processes, etc.

GEMM correctness. See grading output below from GEMM Algorithm 3. Our results match `cublas_gemm` values to within precision error for FP32 numbers (rel difference of 10^{-7}). This is consistent across test cases.

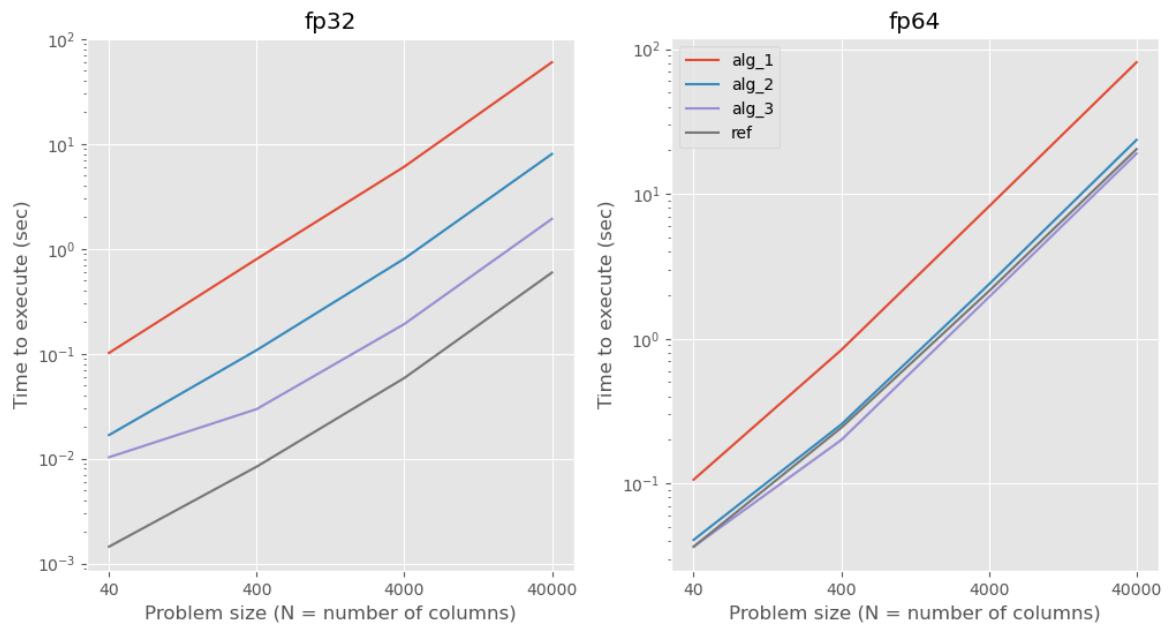


Figure 1: Speed test of GEMM algorithms for float and double precision

M	N	K	Alg 1	Alg 2	Alg 3	Reference
32	4000	40	0.0005	0.0002	0.0001	6E-05
32	4000	400	0.0068	0.0012	0.0012	0.0003
32	4000	4000	0.0932	0.0119	0.0114	0.0019
32	4000	40000	1.2541	0.1185	0.1168	0.0222
320	4000	40	0.0045	0.0015	0.0006	0.0002
320	4000	400	0.0598	0.0104	0.0029	0.0011
320	4000	4000	0.8132	0.1063	0.0294	0.0092
320	4000	40000	14.33	1.0533	0.3143	0.1844
3200	4000	40	0.0432	0.0149	0.0034	0.0026
3200	4000	400	0.5857	0.1009	0.0206	0.0077
3200	4000	4000	8.1372	1.0237	0.2423	0.0762
3200	4000	40000	145.26	10.316	2.6411	0.8254
32000	4000	40	0.4299	0.1517	0.0271	0.0252
32000	4000	400	5.8446	1.0237	0.2287	0.0999
32000	4000	4000	80.56	10.418	2.4419	0.8511
32000	4000	40000	timed out	106.68	28.67	9.7136

Figure 2: Speed test of GEMM algorithms for different M x N dimensions

*** Grading mode 4 ***

main -g 4

Number of MPI processes = 1

Number of CUDA devices = 4

Entering GEMM Benchmarking mode! Stand by.

Starting GEMM 1: M = 3200; N = 4000; K = 3136

GEMM matched with reference successfully! Rel diff = 1.00042e-07

Time for reference GEMM implementation: 0.0690576 seconds

Time for my GEMM implementation: 0.21447 seconds

Completed GEMM 1

Starting GEMM 2: M = 3200; N = 400; K = 4000

GEMM matched with reference successfully! Rel diff = 4.57243e-07

Time for reference GEMM implementation: 0.00814952 seconds

Time for my GEMM implementation: 0.0293013 seconds

Completed GEMM 2

Starting GEMM 3: M = 3200; N = 40; K = 4000

GEMM matched with reference successfully! Rel diff = 7.29707e-07

Time for reference GEMM implementation: 0.00140982 seconds

Time for my GEMM implementation: 0.0101932 seconds

Completed GEMM 3

*** Tests are complete ***

Section 2: Parallelize Neural Network (Single GPU) Idea: training a neural network involves repeatedly updating weights and biases at each layer through forward and backwards propagation. Internally, these operations are mostly various adaptations of generalized matrix multiplication! We want to take existing sequential code and parallelize each using our optimized GEMM algorithm from Section 1.

Based on profiling analysis from my part 1 of the project ('naive' version in table below), I implemented a series of updates to improve the performance of my code.

version	procs	mode_1	mode_2	mode_3	
naive	4	8.352	2.405	0.888	project part 1 implementation
update_1	4	6.420	2.026	0.694	updated kernels to use algorithm 3
update_2	4	6.218	1.989	0.664	incorporated tranpose into kernels
update_3	4	6.194	1.947	0.618	combined gradient descent step
update_4	4	6.129	1.947	0.622	switched to in-place operations
reference	4	2.580	0.980	0.590	

- **Update 1: GEMM Algorithms.** My generalized matrix multiplication from part 1 was very inefficient: each thread computes just one output value and does not make use of shared memory. Algorithm 3 from Section 1 is an order of magnitude faster due to its use of shared memory and sophisticated memory access pattern (more coalesced). My first update was therefore to adapt all variations of GEMM used in our neural network training to use the Algorithm 3 implementation.
- **Update 2: Tranpose Kernels.** My initial version of used additional memory and a dedicated kernel (`kernel_transpose`) to perform the three matrix tranpose steps required in back propagation. While this was nicely modular and therefore useful for unit testing, it was inefficient. This second update eliminates the need to store `d_XT`, `d_aOT`, and `d_W1T` in my cache (removing associated cuda malloc statements) and no longer requires three separate instantiations of the transpose kernel. I did this by simply writing additional GEMM kernel variants that handle transposing input matrices.
- **Update 3: Graident Descent.** My normalization, regularization and gradient descent steps all previously were implemented in separate kernels. It turns out that these steps can be readily combined and re-expressed together as the following matrix addition expression (e.g. for `W[0]`):

$$W[0] -= l * (\frac{1}{n} dW[0] + rW[0])$$

- **Update 4: In-place Operations.** I allocate memory on the device for a bunch of temporary variables to assist with intermediate calculations during feed forward and back propagation steps. Looking at the profiler output, these malloc steps are quite expensive. I implemented updates to eliminate `z0`, `z1`, `yc` and `1ma0` from my cache

variables by overwriting existing cache variables where possible (e.g. sigmoid function can update `a0` in-place).

Each update above resulted in a net improvement for the `mode 1` speed test at $N=4$ processes. The increases were diminishing in magnitude, with `Update 1` having the largest impact (25%) and `Update 4` being almost within margin of error given natural variability of icme gpu runtimes. The best result was 6.129 seconds for `mode 1` after `Update 4`.

Neural Network correctness. For all of the parameter variations tested, the weights and biases produced by the parallel neural network training algorithm match within machine accuracy the CPU sequential code (i.e. max and l2 norms differ by 10^{-7}). Furthermore, when tested using double precision floating point numbers, our norm errors reduce to $O(10^{-16})$ as we would expect if the differences were purely machine accuracy rather than logic inconsistencies.

`* Mode 1 *`

```
mpirun -np 3 /home/jelc/cme213-para/project/main -g 1
```

```
Number of MPI processes = 4
```

```
Number of CUDA devices = 4
```

```
num_neuron=100, reg=0.0001, learning_rate=0.0005, num_epochs=40, batch_size=800
```

```
Loading training data
```

```
Training data information:
```

```
Size of x_train, N = 60000
```

```
Size of label_train = 60000
```

```
Start Parallel Training
```

```
Time for Parallel Training: 6.12856 seconds
```

```
Precision on validation set for parallel training = 0.829167
```

```
Grading mode on. Now checking for correctness...
```

```
Max norm of diff b/w seq and par: W[0]: 1.70544e-07, b[0]: 1.02758e-06
```

```
l2 norm of diff b/w seq and par: W[0]: 1.94657e-07, b[0]: 6.68161e-07
```

```
Max norm of diff b/w seq and par: W[1]: 1.16507e-07, b[1]: 1.61697e-07
```

```
l2 norm of diff b/w seq and par: W[1]: 1.43914e-07, b[1]: 2.84616e-07
```

`* Mode 2 *`

```
mpirun -np 3 /home/jelc/cme213-para/project/main -g 2
```

```
Number of MPI processes = 4
```

```
Number of CUDA devices = 4
```

```
num_neuron=100, reg=0.0001, learning_rate=0.001, num_epochs=10, batch_size=800
```

```
Loading training data
```

```
Training data information:
```

```
Size of x_train, N = 60000
```

```
Size of label_train = 60000
```

Start Parallel Training

Time for Parallel Training: 1.94655 seconds

Precision on validation set for parallel training = 0.756

Grading mode on. Now checking for correctness...

Max norm of diff b/w seq and par: W[0]: 8.80379e-08, b[0]: 5.97242e-07

l2 norm of diff b/w seq and par: W[0]: 1.20557e-07, b[0]: 5.32801e-07

Max norm of diff b/w seq and par: W[1]: 1.03612e-07, b[1]: 2.44647e-07

l2 norm of diff b/w seq and par: W[1]: 1.2233e-07, b[1]: 2.04047e-07

* Mode 3 *

mpirun -np 3 /home/jelc/cme213-para/project/main -g 3

Number of MPI processes = 4

Number of CUDA devices = 4

num_neuron=100, reg=0.0001, learning_rate=0.002, num_epochs=1, batch_size=800

Loading training data

Training data information:

Size of x_train, N = 60000

Size of label_train = 60000

Start Parallel Training

Time for Parallel Training: 0.621657 seconds

Precision on validation set for parallel training = 0.463667

Grading mode on. Now checking for correctness...

Max norm of diff b/w seq and par: W[0]: 3.38911e-08, b[0]: 6.03991e-07

l2 norm of diff b/w seq and par: W[0]: 5.41403e-08, b[0]: 3.4692e-07

Max norm of diff b/w seq and par: W[1]: 5.75366e-08, b[1]: 4.68431e-07

l2 norm of diff b/w seq and par: W[1]: 7.41761e-08, b[1]: 3.59052e-07

Remarks on debugging. This step required significant debugging effort given the inherent complexity of indexing many different variations of parallel matrix operations. My approach here was to add `#if` and `#endif` statements after each major step of the sequential algorithm and try to reproduce that step using my parallel code. To test whether the two implementations matched, I re-used existing `checkErrors` and `checkNNErrors` functions from the provided starter code test suite.

Section 3: Parallelize Training Batches (Multiple GPUs) Idea: while each epoch needs to be executed sequentially, we can perform forwards and backwards propagation on batches within each epoch independently (and therefore in parallel). Our code from Section 2 should already make good use of hardware resources on a single GPU, however, in this project we have access to multiple GPUs! We use MPI to help coordinate sending different training batches ('mini batches') to different GPUs as well as receiving back and aggregating outputs from each batch.

Steps implemented:

- **MPI_Scatter.** We use `MPI_Scatter` to distribute mini batches of input images X and one-hot encoded target variables y across our 4 processes. Each process then conducts its own feed forward and back propagation on its respective minibatch of images and produces a set of partial gradients for weights and biases.

Bonus: While the above approach works for $N=1,2$ and 4 processes (i.e. when number of processes divides batch size), we needed a more advanced code to handle $N=3$. To do this, we implemented a more customizable variation of `MPI_Scatter` called `MPI_Scatterv`. This allows for the user to adjust the number of bytes sent to each process rather than requiring all to receive the same.

- **MPI_AllReduce.** We use `MPI_AllReduce` to sum our partial gradients that have been computed across our 4 processes together and broadcast combined output back to all processes. To do this, we first needed to copy the partial gradients from device to host of each process. Gradient descent can then be computed on all processes (a bit wasteful) to update our neural network parameters (weights and biases).

Note: we needed to be careful to avoid accidentally scaling our gradients and regularization term by the number of processes. To address this, we remove these steps from the back propagation internals and instead apply after `MPI_AllReduce`.

Run times before and after MPI for our naive implementation.

Mode 1: (before) 7.85949 vs (after) 6.56678 seconds

Mode 2: (before) 2.24912 vs (after) 2.03137 seconds

Mode 3: (before) 0.56508 vs (after) 0.69288 seconds

The above runtime statistics show an improvement for mode 1 (epochs = 40) but actually a decrease in performance for mode 3 (epochs = 1). This is because we incur a lot of MPI setup overhead regardless but do not compute enough iterations to benefit from distributing across 4 GPUs (processes).

Section 4: Profiling Optimized Code Idea: use NVIDIA's Nsight Systems and Nsight Compute software tool to interrogate performance of code and assess how to improve. This section from the preliminary report informed the updates described in Section 2 of this document.

Nsight Systems: Used to look across the whole program timeline and identify the largest relative sources of performance loss.

First takeaway (Figure 3) from looking at the updated Nsight Systems timeline was that the `cudaMalloc` is not impacted as much as I would have hoped from removing temporary cache variables. This mirrors the speed test analysis which did not show meaningful improvements from **Update 4** and is likely because we do not touch the larger memory allocation statements (e.g. input image data).

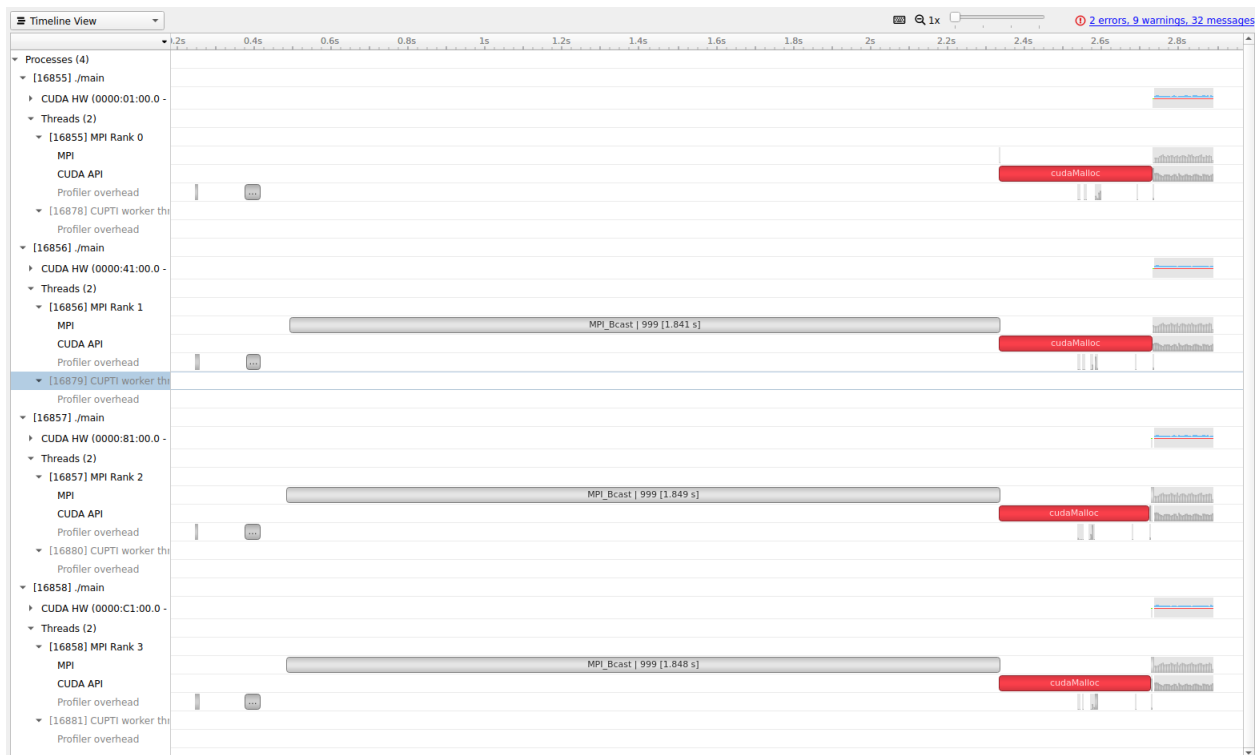


Figure 3: Nsight Systems timeline of entire program for single epoch

Second takeaway (Figure 4) was relative expense of `MPI_Scatter` and `cudaMemcpy` operations compared with our feed forward, back propagation and gradient descent kernels. To me this highlights the importance to reducing the number of movements to and from the device, especially within the batch loops as is currently the case for X and y as well as gradients pre and post `MPI_AllReduce`. A future implementation might consider splitting work differently across my 4 GPUs (processes) to reduce `cudaMemcpy` and `MPI` statements. Currently I divide each batch into 'mini batches', however, instead I could think about dividing my model parameters (weights and biases) across the different processes.

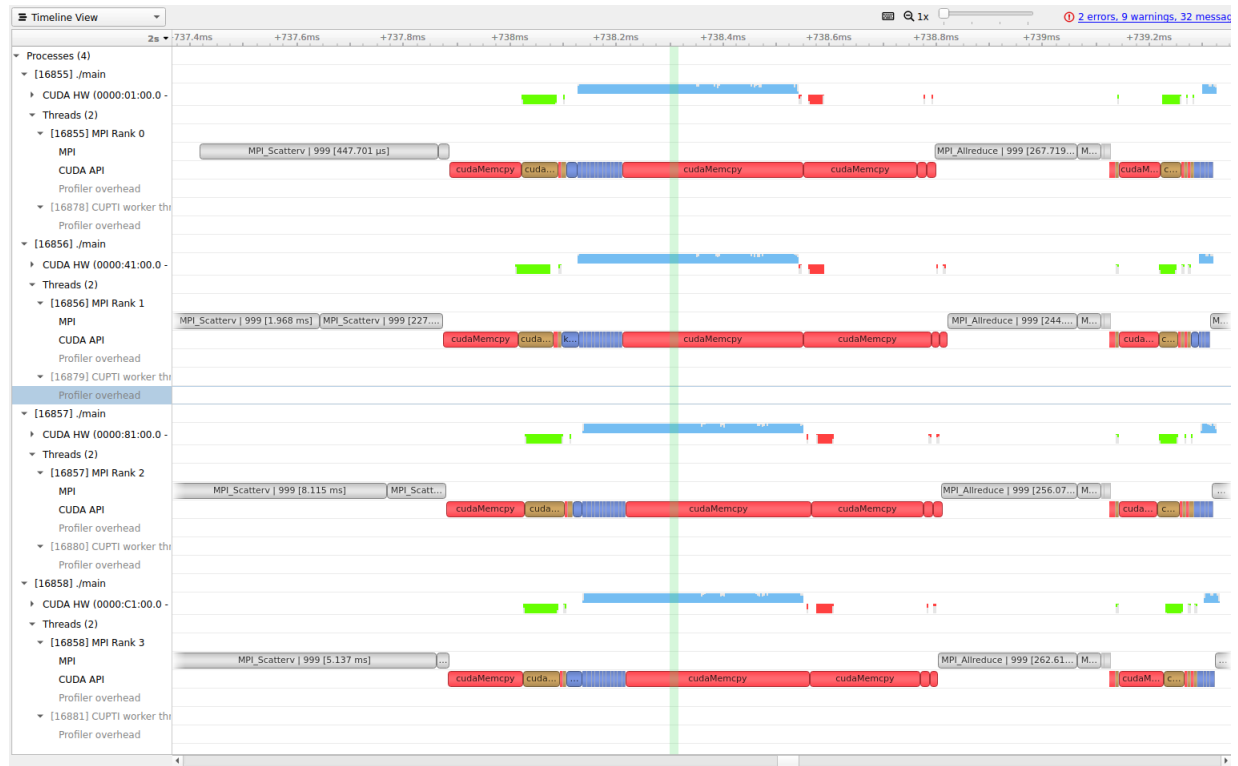


Figure 4: Nsight Systems timeline for training of one image batch

Third takeaway (Figure 5) was that our training pipeline takes noticeably (approx. 50%) less time for each epoch! This looks to be because (a) we execute far fewer kernels (16 vs 25 for each training cycle), and (b) our GEMM related kernels are all more performant. This is great news and also agrees with our runtime statistics where updates 1 and 2 made the largest difference.

Nsight Compute: Used to take a deeper look into the performance of individual kernels. In particular, our generalized matrix multiplication kernel.

Main takeaway is ... we are approaching memory bound... better arithmetic intensity for double precision ...

Finally, Nsight Compute's launch statistics suggests that I launch too small a grid for the available hardware specs. Given fixed thread block dimensions and `numXPerThread`, our grid size is fixed to match the input matrices and so in some sense we are constrained. That said, it would be interesting to experiment with smaller thread block or `numXPerThread` settings that allow for a larger grid and potentially use more of the available hardware resource at the expense of memory access being less coalesced.

