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Section: *CS 483 AL1 (76324)*

ECE 408/CS483 Milestone 3 Report

0. List Op Times, whole program execution time, and accuracy for batch size of 100, 1k, and 5k images from your basic forward convolution kernel in milestone 2. This will act as your baseline this milestone. Note: **Do not** use batch size of 10k when you profile in --queue rai_amd64_exclusive. We have limited resources, so any tasks longer than 3 minutes will be killed. Your baseline M2 implementation should comfortably finish in 3 minutes with a batch size of 5k (About 1m35 seconds, with nv-nsight).

			Total	
Batch Size	Op Time 1	Op Time 2	Execution	Accuracy
			Time	
100	0.262705	0.957244	0m1.542s	0.86
	ms	ms		
1000	2.39847	9.13148 ms	0m10.144s	0.886
	ms			
5000	11.8805	45.393 ms	0m48.627s	0.871
	ms			

1. Optimization 1: Using Streams to overlap computation with data transfer (4 points)

a. Which optimization did you choose to implement and why did you choose that optimization technique.

I chose to implement the streams to overlap computation with data transfer. In my mind, this done as I felt this could have real efficacy when scaled up to larger batch sizes, as it utilizes GPU resources even harder than a single kernel launch, while also freeing up the host code to get its end of the bargain (memory control) out of the way. The way we implemented it in class examples was very similar to how I did it, as I noticed the input and output data is able to be segmented over the image batches with minimal changes to the actual kernel, allowing for progressive data loads and reads over multiple kernel launches in a sort of batches of batches approach.

b. How does the optimization work? Did you think the optimization would increase performance of the forward convolution? Why? Does the optimization synergize with any of your previous optimizations?

Code for the involved is below and in the stream-new-forward.cu file in the submission in the m3opts folder.

The optimization starts with modifying the kernel to be launched multiple times by adding a number of streams to the host code using the cudaStreamCreate fns. Thru this, we are able to use the cudaMemcpyAsync api to achieve asynchronism with memcpys in HtoD direction right before a kernel launch is needed, and a DtoH direction right afterwords. This is while other sections of the overall Project Milestone 2 (PM2) kernel are also running at the same time. By splitting up the B batch processed images over NUM_STREAMS streams, I can run multiple kernel independently. By computing the exact amount of data needed for each one of those kernel launches, I was able to organize a set of memcpys in the loop to insert/extract data from the kernels while they are running asynchronously. This involved a rather involved piece of host code logic to account for leftover (not nicely sized) workloads. The kernel required a small adjustment to take in a streamNum in order to figure out which B image batch a thread block is running on, replacing the standard blockldx.z identifier. Code for the involved is below and in the stream-new-forward.cu file in the submission in the m3opts folder.

```
const int H out = (H - K)/S + 1;
    const int W_{out} = (W - K)/S + 1;
them, or create your own.
    // float a = in 4d(0,0,0,0)
    #define out_4d(i3, i2, i1, i0) output[(i3) * (M * H_out * W_out) + (i2) * (H_out
* W_out) + (i1) * (W_out) + i0]
    #define in 4d(i3, i2, i1, i0) input[(i3) * (C * H * W) + (i2) * (H * W) + (i1) *
(W) + i0]
    #define mask_4d(i3, i2, i1, i0) mask[(i3) * (C * K * K) + (i2) * (K * K) + (i1)
* (K) + i0]
   // same as grid setup
    int W_size = ceil(1.0f*W_out/TILE_WIDTH); // number of horizontal tiles per
    int H_size = ceil(1.0f*H_out/TILE_WIDTH); // number of vertical tiles per output
map
    int b = blockIdx.z + (ceil(1.0f * B / NUM_STREAMS) * streamNum); // batch num is
based on streamNum iteration
    int m = blockIdx.x;
    int h = (blockIdx.y / W_size) * TILE_WIDTH + threadIdx.y; // target h of output
    int w = (blockIdx.y % W_size) * TILE_WIDTH + threadIdx.x; // target w of output
    // each thread ran should be within output bounds, otherwise return
    if (w < 0 || w >= W_out || h < 0 || h >= H_out || b >= B)
        return:
    float acc = 0.0f;
    for (int c = 0; c < C; c++) { // sum over all input channels
        for (int p = 0; p < K; p++) { // loop over KxK filter
            for (int q = 0; q < K; q++) {
                int h_idx = (h * S + p);
                int w idx = (w * S + q);
                if (!(w_idx < 0 || w_idx >= W || h_idx < 0 || h_idx >= H)) {
                    acc += in_4d(b, c, h_idx, w_idx) * mask_4d(m, c, p, q);
```

```
out_4d(b, m, h, w) = acc;
   #undef out 4d
   #undef in 4d
   #undef mask 4d
 _host__ void GPUInterface::conv_forward_gpu_prolog(const float *host_output, const
float *host_input, const float *host_mask, float **device_output_ptr, float
**device_input_ptr, float **device_mask_ptr, const int B, const int M, const int C,
   // cudaError t error = cudaGetLastError();
   #define wbCheck(stmt)
       cudaError t err = stmt;
        if (err != cudaSuccess) {
            std::cout<<"Failed to run stmt: "<<#stmt<<std::endl;</pre>
            std::cout<<"CUDA error: "<<cudaGetErrorString(err)<<std::endl;</pre>
            exit(-1);
    } while (0)
    const int H out = (H - K)/S + 1;
    const int W_{out} = (W - K)/S + 1;
   size_t dop_sz = B * M * H_out * W_out * sizeof(float);
   size_t dip_sz = B * C * H * W * sizeof(float);
   size_t dmp_sz = M * C * K * K * sizeof(float);
   // device mask stream/event
   cudaStream t dms;
```

```
cudaStreamCreate(&dms);
    cudaEvent_t dme;
    cudaEventCreate(&dme);
    // would be async but we need CUDA 11.3+ for that. Minimal impact anyways
    wbCheck(cudaMalloc((void **)device output ptr, dop sz));
    wbCheck(cudaMalloc((void **)device input ptr, dip sz));
    wbCheck(cudaMalloc((void **)device_mask_ptr, dmp_sz));
    cudaHostRegister((void *) host_output, dop_sz, 0);
    cudaHostRegister((void *) host_input, dip_sz, 0);
    cudaHostRegister((void *) host_mask, dmp_sz, 0);
    // async memcpy here, record event after done to signal before kernel
    wbCheck(cudaMemcpyAsync(*device_mask_ptr, host_mask, dmp_sz,
cudaMemcpyHostToDevice, dms));
    cudaEventRecord(dme, dms);
    int streamSize = ceil(1.0f * B / NUM STREAMS); // proportion of B to batch
process in each stream
    int W size = ceil(1.0f*W out/TILE WIDTH); // number of horizontal tiles per
output map
    int H size = ceil(1.0f*H out/TILE WIDTH); // number of vertical tiles per output
    int tileNums = H_size * W_size; // total number of tiles per map
    dim3 DimBlock(TILE_WIDTH, TILE_WIDTH, 1); // output tile for untiled code
    dim3 DimGrid(M, tileNums, streamSize);
    std::cout<<"DimBlock:</pre>
"<<DimBlock.x<<"x"<<DimBlock.z<<std::endl;</pre>
    std::cout<<"DimGrid: "<<DimGrid.x<<"x"<<DimGrid.y<<"x"<<DimGrid.z<<std::endl;</pre>
    cudaStream_t* streams = (cudaStream_t*) malloc(NUM_STREAMS *
sizeof(cudaStream_t));
    for (int streamNum = 0; streamNum < NUM_STREAMS; streamNum++) {</pre>
        cudaStreamCreate(&(streams[streamNum]));
    bool ended = false;
    for (int streamNum = 0; streamNum < NUM STREAMS; streamNum++) {</pre>
        int offset = streamNum * streamSize; // number of Bs to skip for start of
stream
        int outStreamBytes = streamSize * M * H_out * W_out * sizeof(float); // how
many Bs to transfer back to host
        int inStreamBytes = streamSize * C * H * W * sizeof(float); // how many Bs
```

```
bounds sizes for input and output
        int outDiff = dop_sz - ((offset * M * H_out * W_out * sizeof(float)) +
outStreamBytes);
        int inDiff = dip_sz - ((offset * C * H * W * sizeof(float)) +
inStreamBytes);
        // if we already ended, just skip this stream
        if (ended)
            continue;
        if((outDiff <= 0 || inDiff <= 0) && !ended)</pre>
            ended = true;
        // std::cout<<"Starting stream: "<<streamNum<<", offset: "<< offset << ",
outB: "<< outStreamBytes << ", inB: "<< inStreamBytes <<", Ended:</pre>
"<<ended<<std::endl;
        if(outDiff < 0)
            outStreamBytes += outDiff;
        if(inDiff < 0)</pre>
            inStreamBytes += inDiff;
        // general "stacking" idea from Mark Harris of Nvidia
        // changed up drastically to account for not-nice parameters and leftover
work conditions, along with wait event
        cudaMemcpyAsync(&(*device_input_ptr)[offset * C * H * W], &host_input[offset
* C * H * W], inStreamBytes, cudaMemcpyHostToDevice, streamS[streamNum]);
        cudaStreamWaitEvent(streams[streamNum], dme, 0);
        conv_forward_kernel<<<DimGrid, DimBlock, 0, streams[streamNum]>>>(streamNum,
*device_output_ptr, *device_input_ptr, *device_mask_ptr, B, M, C, H, W, K, S);
        cudaMemcpyAsync((void *) &host output[offset * M * H out * W out],
&(*device_output_ptr)[offset * M * H_out * W_out], outStreamBytes,
cudaMemcpyDeviceToHost, streams[streamNum]);
    for (int streamNum = 0; streamNum < NUM STREAMS; streamNum++) {</pre>
        cudaStreamDestroy(streams[streamNum]);
    cudaEventDestroy(dme);
    cudaStreamDestrov(dms):
```

```
cudaHostUnregister((void *) host_output);
    cudaHostUnregister((void *) host input);
    cudaHostUnregister((void *) host_mask);
   // not needed because we sync after anyways
   // cudaDeviceSynchronize();
   free(streams);
   #undef wbCheck
_host__ void GPUInterface::conv_forward_gpu(float *device_output, const float
*device\_input, const float *device\_mask, const int B, const int M, const int C,
_host__ void GPUInterface::conv_forward_gpu_epilog(float *host_output, float
*device_output, float *device_input, float *device_mask, const int B, const int M,
   #define wbCheck(stmt)
        cudaError t err = stmt;
        if (err != cudaSuccess) {
            std::cout<<"Failed to run stmt: "<<#stmt<<std::endl;</pre>
            std::cout<<"CUDA error: "<<cudaGetErrorString(err)<<std::endl;</pre>
            exit(-1);
   } while (0)
   // All we need to do here is cleanup
   wbCheck(cudaFree(device input));
   wbCheck(cudaFree(device_output));
   wbCheck(cudaFree(device_mask));
   #undef wbCheck
```

```
_host__ void GPUInterface::get_device_properties()
    int deviceCount;
    cudaGetDeviceCount(&deviceCount);
    for(int dev = 0; dev < deviceCount; dev++)</pre>
        cudaDeviceProp deviceProp;
        cudaGetDeviceProperties(&deviceProp, dev);
        std::cout<<"Device "<<dev<<" name: "<<deviceProp.name<<std::endl;</pre>
        std::cout<<"Computational capabilities:</pre>
"<<deviceProp.major<<"."<<deviceProp.minor<<std::endl;</pre>
        std::cout<<"Max Global memory size: "<<deviceProp.totalGlobalMem<<std::endl;</pre>
        std::cout<<"Max Constant memory size:</pre>
"<<deviceProp.totalConstMem<<std::endl;</pre>
        std::cout<<"Max Shared memory size per block:</pre>
"<<deviceProp.sharedMemPerBlock<<std::endl;
        std::cout<<"Max threads per block:</pre>
"<<deviceProp.maxThreadsPerBlock<<std::endl;</pre>
        std::cout<<"Max block dimensions: "<<deviceProp.maxThreadsDim[0]<<" x,</pre>
"<<deviceProp.maxThreadsDim[1]<<" y, "<<deviceProp.maxThreadsDim[2]<<"
z"<<std::endl;</pre>
        std::cout<<"Max grid dimensions: "<<deviceProp.maxGridSize[0]<<" x,</pre>
"<<deviceProp.maxGridSize[1]<<" y, "<<deviceProp.maxGridSize[2]<<" z"<<std::endl;</pre>
        std::cout<<"Warp Size: "<<deviceProp.warpSize<<std::endl;</pre>
```

I thought it would increase performance simply on the fact that we are utilizing more of the GPUs resources in order to produce the same outputs. The overhead did not seem too large in theory, but I supposed the smaller batch sizes left the optimization in a tough spot of not being able to show its efficacy.

It is able to synergize with any other optimization that happens on the level of the base batch image units, so fp16 optimizations or loop unrolling would work with this approach as well.

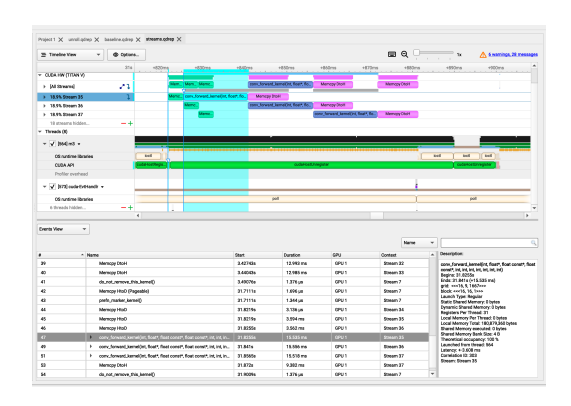
c. List the Op Times, whole program execution time, and accuracy for batch size of 100, 1k, and 5k images using this optimization (including any previous optimizations also used).

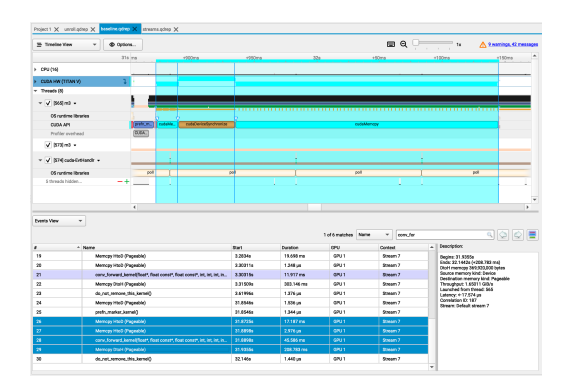
			Total	
Batch Size	Op Time 1	Op Time 2	Execution	Accuracy
			Time	
100	7.37061	6.46393 ms	0m1.715s	0.86
	ms			
1000	63.838 ms	53.3827 ms	0m10.680s	0.886
5000	176.503	148.22 ms	0m50.561s	0.871
	ms			
	NOTE:			
	layer time			
	used,			
	optimes			
	are super			
	small			

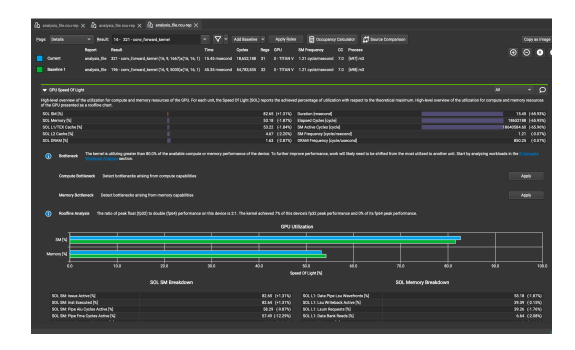
d. Was implementing this optimization successful in improving performance? Why or why not? Include profiling results from *nsys* and *Nsight-Compute* to justify your answer, directly comparing to your baseline (or the previous optimization this one is built off of).

I'll focus on 5000 batch size optime. For the baseline, we have 11.8805 + 45.393 = 57.2735. However, for the streams, the optimes were exceedingly low as due to how the stream opt needed to be done, it had to be outside of the testing fn. As such the layer times were 176.503 + 148.22 = 324.723 for the streams opt. There isn't really a direct way to compare these values of layer time and optime though, and so I feel like to show the opt, we can look directly at reports and extrapolate from there.

Below is the screenshot for the timing graph of the nsys command for the stream opt as shown via the UI for NVIDIA Nsight Systems . As you can see, the relevant memcpys and kernel launches that occur are stacked, sharing execution time just as we needed. If we were to stretch out these transfers, they would significantly total up the time, as shown via the baseline pic below it. Additionally, the host code is ahead of the device by a decent margin, which ends up freeing the host thread much sooner to do other work: the result we were looking for. The kernel is unchanged and thus Nsight-Compute is not used here, but here is an example of the reduced kernel's details screenshot, where you see the SM % util is up while the memory % went a bit down, although this is for a reduced batch size per kernel so comparisons aren't 1-1.







e. What references did you use when implementing this technique?

The refs here were NVIDIA based. The general "stacking" idea of overlaying the memcpys and kernels were from Mark Harris of Nvidia https://developer.nvidia.com/blog/how-overlap-data-transfers-cuda-cc/. Runtime api docs were also used here https://docs.nvidia.com/cuda/cuda-runtime-api/group CUDART STREAM.html#group CUDART STREAM 1g7840e39847999 https://docs.nvidia.com/cuda/cuda-runtime-api/group CUDART STREAM 1g7840e39847999 <a href="https://docs.nvidia.com/cuda/cuda-runtime-api/group cuda-runtime-api/group cuda-runtime-api/group

2. Optimization 2: FP16 arithmetic. (note this can modify model accuracy slightly) (4 points)

- a. Which optimization did you choose to implement and why did you choose that optimization technique.
 - I chose the FP16 arithmetic optimization because since the GPU is such a compute oriented device, it make sense to avoid working with abstractions such as FP32 in order to get more lower level control of the ops and utilize some more primitive instructions.
- b. How does the optimization work? Did you think the optimization would increase performance of the forward convolution? Why? Does the optimization synergize with any of your previous optimizations?

This optimization is within the fp16-new-forward.cu file in the submission. m3opts/folder.

The optimization starts with modifying the host code to convert all the float input and mask data from float to __half data type via the __float2half() fn. After that, I took memcpyed the data to the device ptrs and fed it to the kernel, which now accepts __restrict__ half ptrs, the __restrict__ to hopefully squeeze out more optimization over our assurance that the ptrs do not interleave. From there, the main operation to optimize the the multiply from the input to the mask, then the add to the output. This was able to be done in one fused op instruction via the __hfma() fn, which deals with the FP16 floats. After that, some more host code to copy back the __half ptr output and then a set of __half2float() in order to convert back into the host_output arr used for the actual submission. NOTE: conversion from floats and halfs in host code was oked by this instructor comment, otherwise a simple kernel like from MP7 could be used to do the conversions https://campuswire.com/c/G184FB646/feed/636.

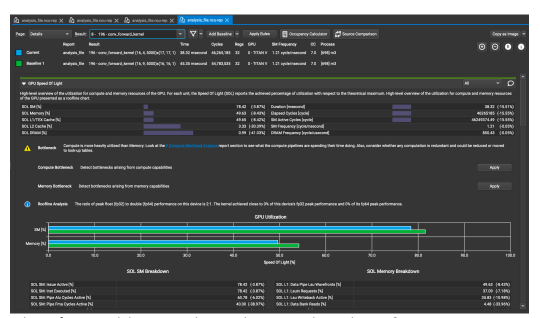
The performance should be increased due to the lessening of data volume, along with playing by the FP16 primitives the GPU actually uses under the hood (especially that super fast fused multipy add instruction). It synergizes with streams as it doesn't care about the underlying data types, as long as you swap over to __half in all relevant places. Same with the loop unrolling, as long as you swap over the ops used from a singlular to multiple hfma() calls.

 List the Op Times, whole program execution time, and accuracy for batch size of 100, 1k, and 5k images using this optimization (including any previous optimizations also used).

			Total	
Batch Size	Op Time 1	Op Time 2	Execution	Accuracy
			Time	
100	0.351344	0.837059	0m2.825s	0.86
	ms	ms		
1000	3.1836 ms	7.79319 ms	0m11.800s	0.887
5000	15.7403	38.359 ms	0m52.363s	0.8712
	ms			

d. Was implementing this optimization successful in improving performance? Why or why not? Include profiling results from *nsys* and *Nsight-Compute* to justify your answer, directly comparing to your baseline (or the previous optimization this one is built off of).

So from the pov of the optimes, I'll focus on 5000 batch size. For the baseline, we have 11.8805 + 45.393 = 57.2735. Comparing to the fp16, we have 15.7403 + 38.359 =54.0993, a nearly 3ms improvement overall. As we can also see, the smaller layer pass went thru quicker on the baseline, meaning this approach isn't as robust for smaller sizes, but the larger layer pass cut down by about 7ms, indicating these performance gains tend to scale. I will now show the comparison between the fp16 and the baseline kernel performance. As you can see, the SM% and the Memory% actually went down from the baseline; however, the timing of the actual kernel execution is actually way down from the baseline, meaning the kernel still outputs faster results despite the bandwidth hits. This tracks with this article I found on half arithmetic (https://ionthruster.medium.com/an-introduction-to-writing-fp16-code-for-nvidias-gpusda8ac000c17f) in which it states in the Performance section that "With half and 32 threads/warp — we only hit ~64B/Load, whereas to achieve good bandwidth we need to strive to hit at least 128B/Load", meaning we want to try and pack in two __half ops at the same time to save both on memory and SM performance. This could involve a refactoring to the half2 datatype if future optimizations wanted to go down that route, and I believe it would optimize this code even harder than what we have with just standard fp16 floats



e. What references did you use when implementing this technique?

I utilized the half info section of the CUDA docs https://docs.nvidia.com/cuda/cuda-math-api/group CUDA MATH HALF MISC.html, and then combined this with the docs for the __hfma() fn https://docs.nvidia.com/cuda/cuda-math-api/group CUDA MATH HALF ARITHMETIC.html#group CUDA MATH HALF ARITHMETIC 1gaec96bd410157b5813c940ee320175f2. I also utilized Ion Thrusters medium article https://ion-thruster.medium.com/an-introduction-to-

<u>writing-fp16-code-for-nvidias-gpus-da8ac000c17f</u> as a sort of post-hoc source for why the mem and SM bandwidth might be down even though we are using the __half datatype.

3. Optimization 3: Tuning with restrict and loop unrolling (considered as one optimization only if you do both) (3 points)

(Delete this section blank if you did not implement this many optimizations.)

a. Which optimization did you choose to implement and why did you choose that optimization technique.

For this optimization, I used the __restrict__ ptrs along with loop unrolling within the kernel convolution fn for various kernel params. I have heard that loop unrolling is a classic C optimization to squeeze more performance out of the processor by stacking operations in larger batches over a total. As such, it would seem natural to try it out in order to try it over more than a couple threads on a CPU. The restrict keyword is also something I have seen as a contract or hint that helps out a lot with compiler optimization, and for similar reasons would make sense to try.

b. How does the optimization work? Did you think the optimization would increase performance of the forward convolution? Why? Does the optimization synergize with any of your previous optimizations?

This optimization is within the loopunrollrestrict-new-forward.cu file in the submission. m3opts/folder.

It involves tagging all 3 data pointers in the kernel fn with __restrict__ to hint to the compiler that this is non-aliased, read-only memory that doesn't overlap. Further into the execution, the looping over the KxK mask is unrolled such that each loop iteration does multiple operations at once based on K. For sufficiently small K, it is unchanged as performance gain is minimal. For middling values, it does not perform as many unrolled ops due to overhead, but larger masks have more unrolled ops. There are a lot of local vars involved to split apart the loop unroll in order to organize the logic and remove redundant calculations. It should increase performance for the reasons listed in section A, in that __restrict__ hints to the compiler that you can go crazier on these optimizing pointers because we are being "nice" and not overlapping stuff, and the loop unrolling is meant to get away from the costly and sort of unneeded loop iteration on every single multiply-add operation. It synergizes with the streams and FP16 optimizations, as it changes the loop behavior, not the underlying operation. As long as the loops are intact in functionality, this optimization can help.

c. List the Op Times, whole program execution time, and accuracy for batch size of 100, 1k, and 5k images using this optimization (including any previous optimizations also used).

			Total	
Batch Size	Op Time 1	Op Time 2	Execution	Accuracy
			Time	
100	0.252222	0.84406 ms	0m4.207s	0.86
	ms			
1000	2.23091	7.87841 ms	0m11.137s	0.886
	ms			
5000	11.0115	39.0854 ms	0m51.149s	0.871
	ms			

d. Was implementing this optimization successful in improving performance? Why or why not? Include profiling results from *nsys* and *Nsight-Compute* to justify your answer, directly comparing to your baseline (or the previous optimization this one is built off of).

I wanted to start the optimization by tweaking the # of unrolled ops on the K cases between 3 and 8 (targeting the 5000 batch request we run after the test cases pass). I ran through various unroll factors for the 5000 batch size kernels, and these were the profiling results from the terminal output.

// unroll of 4 http://s3.amazonaws.com/files.rai-project.com/userdata/build-656bb81edbc25168da9ef6db.tar.gz

Op Time: 12.29 ms Op Time: 40.8108 ms Test Accuracy: 0.871 real 0m52.019s

// unroll of 3 http://s3.amazonaws.com/files.rai-project.com/userdata/build-656bb9b2dbc2516a4709382b.tar.qz

Op Time: 11.0115 ms Op Time: 39.0854 ms Test Accuracy: 0.871 real 0m51.149s

// unroll of 2 http://s3.amazonaws.com/files.rai-project.com/userdata/build-656bbb56dbc2516bb4aa423b.tar.gz

Op Time: 11.1923 ms Op Time: 40.2075 ms Test Accuracy: 0.871 real 0m51.068s

// unroll of 1 is just baseline so not tested

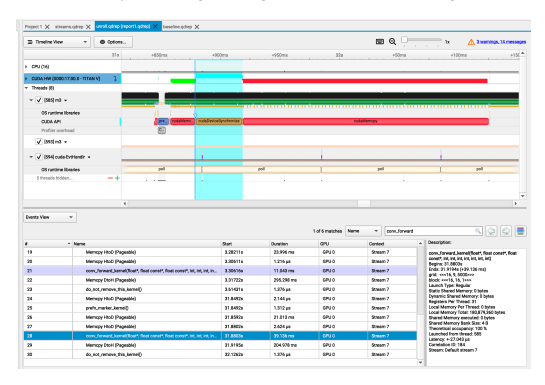
From this, it was clear that in terms of total optime, the unroll of 3 was the most performant. As such, I decided on that as my unroll factor to continue.

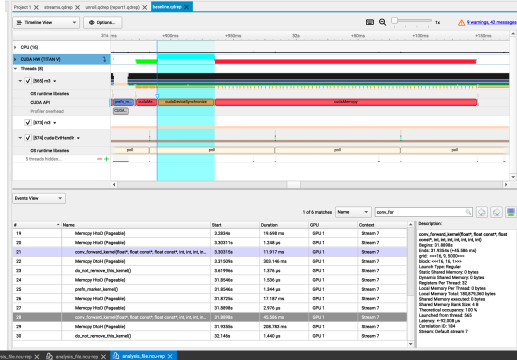
I'll focus on 5000 batch size optime. For the baseline, we have 11.8805 + 45.393 = 57.2735. For this loop unrolling logic with the unroll factor of 3, we see 11.0115 + 11.0115

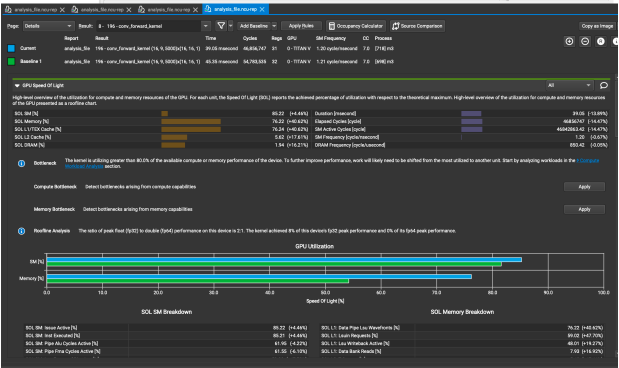
39.0854 = 50.0969, about 7ms less. From this, we can see this optimization does have the potential to enable the same amount of computation in less time. In all, it does seem like this optimization will shine more with larger masks as opposed to the rather small masks we have right now (3 or 7?) as that is the main variable to optimize against in this opt.

Below is the unroll report, followed by the baseline report. The last kernel launch (the one we care about) in the unroll report was 39.136ms, while the baseline was 45.586ms. This tracks as we see a huge ms drop off on the unroll due to that unroll factor being tuned in for the best gains in the timing department.

If we take a look at the kernels, we can see that the unroll kernel utilized 4.46% more of the memory in the SOL tests, compared to the baseline. And even more profound, the memory utilization jumped by a giant margin of 40.62%, showing this is nothing but a proper optimization that takes more advantage of the memory bandwidth. Results are overlayed with the green being the baseline and the blue being the unroll.







e. What references did you use when implementing this technique?

I was kickstarted by this forum post on the benefits of loop unrolling https://forums.developer.nvidia.com/t/automatic-loop-unrolling/10502 which refers to a section in the best practices programming guide where this is esposed as a benefit. While the general idea was rather simple (split individual for loop iterations into batches), and even mirrored on large scale via the streams op, the real time spent was in tuning the # of ops per unrolled loop iteration in order to try and get optimal results by running the profiling queue again and again. Tooling the specifics of the manual loop unroll was done by myself after settling on a design.