# 15-418/618, Spring 2018 Assignment 2 A Simple CUDA Renderer

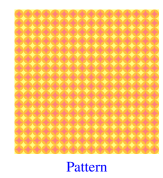
Assigned: Mon., Jan. 29

Due: Wed., Feb. 14, 11:59 pm

Last day to handin: Sat., Feb. 17









Overview

In this assignment you will write a parallel renderer in CUDA that draws colored circles. While this renderer is very simple, parallelizing the renderer will require you to design and implement data structures that can be efficiently constructed and manipulated in parallel. This is a challenging assignment so you are advised to start early. Don't be lulled into thinking that the first two parts are indicative of the time you will require to complete the third part. **Seriously, you are advised to start early.** Good luck!

Before you begin, please take the time to review the course policy on academic integrity at:

http://www.cs.cmu.edu/~418/academicintegrity.html

# **Environment Setup**

You should use the GHC machines containing NVIDIA GeForce GTX 1080 GPUs. These have host names ghcX. ghc. and rew. cmu. edu, for X between 26 and 46. The Wikipedia entry for GeForce 10 GPUs provides useful information about this model of GPU. They support CUDA compute capability 6.1.

**Note:** the performance capabilities of these GPU's varies, meaning that your results will vary across these machines. However, feel free to run your code on any of the specified machines. We have distributed executable versions of our implementations (referred to as the "targets"), which are used for evaluating the performance of your programs.

Using the GHC machines:

- 1. If you are using the ssh command to reach the machine, include the option "-Y" in your command line to enable the use of X windows.
- 2. The NVIDIA CUDA C/C++ Compiler (NVCC) is located at /usr/local/depot/cuda-8.0/bin/, which you need to add to your PATH. To do this, add the line below to the file below, according to which shell you use. (You can determine which shell you are using with the command "echo \$SHELL.")

Shell	File	Add	
bash	~/.bashrc	export	PATH=/usr/local/depot/cuda-8.0/bin:\${PATH}
csh	~/.cshrc	setenv	PATH /usr/local/depot/cuda-8.0/bin:\${PATH}

3. The CUDA shared library is located at /usr/local/depot/cuda-8.0/lib64/. It must be loaded at runtime. Add the line below to your RC file (~/.bashrc or ~/.cshrc) according to which shell you use.

```
export LD_LIBRARY_PATH=/usr/local/depot/cuda-8.0/lib64/:${LD_LIBRARY_PATH}
setenv LD_LIBRARY_PATH ${LD_LIBRARY_PATH}:/usr/local/depot/cuda-8.0/lib64/
```

- 4. Make sure you source (or reload) your RC file. If using bash, run "source ~/.bashrc." If using csh, run "source ~/.cshrc."
- 5. Download the Assignment 2 starter code from the course Github using:

```
git clone https://github.com/cmu15418/asst2-s18.git
```

#### Resources

For any C++ questions (like what does the *virtual* keyword mean), the C++ Super-FAQ is a great resource that explains things in a way that's detailed yet easy to understand (unlike a lot of C++ resources), and was co-written by Bjarne Stroustrup, the creator of C++!

The CUDA C programmer's guide is available in PDF or HTML. It is an excellent reference for learning how to program in CUDA.

You can also find a large number of examples in the CUDA SDK /usr/local/depot/cuda-8.0/samples. In addition, there are a wealth of CUDA tutorials and SDK examples on the web (just Google!) and on the NVIDIA developer site.

# 1 CUDA Warm-Up 1: SAXPY (5 pts)

To gain a bit of practice writing CUDA programs your warm-up task is to re-implement the SAXPY function from Assignment 1 in CUDA. Starter code for this part of the assignment is located in the saxpy directory.

Finish off the implementation of SAXPY in the function <code>saxpyCuda()</code> in <code>saxpy.cu</code>. You will need to allocate device global memory arrays and insert calls to move data between the host and device memories. These issues are covered in Section 3.2.2 of the Programmer's Guide.

As part of your implementation, add timers around the CUDA kernel invocation in saxpyCuda(). After your additions, your program should time two executions:

- The provided starter code contains timers that measure the **entire process** of copying data to the GPU, running the kernel, and copying data back to the CPU.
- Your timers should measure only the time taken to run the kernel. (They should not include the time of CPU to GPU data transfer or transfer of results back to the CPU.)

When adding your timing code, be careful: The CUDA kernel's execution on the GPU is asynchronous with the main application thread running on the CPU. You should place a call to <code>cudaThreadSynchronize()</code> following the kernel call to wait for completion of all CUDA work on the GPU. This call returns only when all prior CUDA work on the GPU has completed. (Without waiting for the GPU to complete, your CPU timers will report that essentially no time elapsed!) Note that in your measurements that include the time to transfer data back to the CPU, a call to <code>cudaThreadSynchronize()</code> is not necessary before the final timer (after your call to <code>cudaMemcpy()</code> that returns data to the CPU) because <code>cudaMemcpy()</code> will not return to the calling thread until after the copy is complete.

**Question:** What performance do you observe compared to the sequential CPU-based implementation of SAXPY (recall program 5 from Assignment 1)? Compare and explain the difference between the results provided by two sets of timers (the timer you added and the timer that was already in the provided starter code). Are the bandwidth values observed roughly consistent with the reported bandwidths available to the different components of the machine? *Hint*: You should use the web to track down the memory bandwidth of an NVIDIA GTX 1080 GPU, and the maximum transfer speed of the computer's PCIe-x16 bus. It's PCIe 3.0, and a 16 lane bus connecting the CPU with the GPU.

# 2 CUDA Warm-Up 2: Parallel Prefix-Sum (10 pts)

Now that you're familiar with the basic structure and layout of CUDA programs, as a second exercise you are asked to come up with parallel implementation of the function find\_repeats() which, given a list of integers A, returns a list of all indices i for which A[i] == A[i+1]. (The first element of an array has index 0.)

For example, given the array [1, 2, 2, 1, 1, 1, 3, 5, 3, 3], your program should output the array [1, 3, 4, 8]. All code for this part of the assignment is in the directory scan.

```
void exclusive scan recursive(int* start, int* end, int* output, int* scratch)
    int N = end - start;
    if (N == 0) return;
    else if (N == 1)
        output[0] = 0;
        return;
    }
    // sum pairs in parallel.
    for (int i = 0; i < N/2; i++)
        output[i] = start[2*i] + start[2*i+1];
    // prefix sum on the compacted array.
    exclusive_scan_recursive(output, output + N/2, scratch, scratch + (N/2));
    // finally, update the odd values in parallel.
    for (int i = 0; i < N; i++)
        output[i] = scratch[i/2];
        if (i % 2)
            output[i] += start[i-1];
    }
```

Figure 1: Recursive implementation of exclusive prefix sum

#### **Exclusive Prefix Sum**

We want you to implement find\_repeats() by first implementing the parallel exclusive prefix-sum operation. This operation is covered in 15-210, where it is referred to by its more general form: the "scan" operation.

Exclusive prefix sum takes an array A and produces a new array output that has, at each index i, the sum of all elements up to but not including A[i]. For example, given the array [1,4,6,8,2], the output of exclusive prefix sum [0,1,5,11,19].

A recursive implementation should be familiar to you from 15-210. As a review (or for those that did not take 15-210), the code in Figure 1 is a C implementation of a work-efficient, parallel version of scan. Details on prefix-sum (and its more general relative, scan) can be found in Chapter 6 of the 15-210 lecture notes.

While the recursive code expresses our intent well and recursion is supported on modern GPUs, its use can lead to fairly low performance due to function call and stack frame overhead. Instead we can express the algorithm in an iterative manner. The "C-like" code in Figure 2 is an iterative version of scan. We use  $parallel\_for$  to indicate potentially parallel loops. To understand this code, you may find the visualization in Figure 3 for the case of N = 16 helpful.

```
void exclusive_scan_iterative(int* start, int* end, int* output)
    int N = end - start;
    memmove(output, start, N*sizeof(int));
    // upsweep phase.
    for (int twod = 1; twod < N; twod\star=2)
        int twod1 = twod*2;
        parallel_for (int i = 0; i < N; i += twod1)</pre>
            output[i+twod1-1] += output[i+twod-1];
    output [N-1] = 0;
    // downsweep phase.
    for (int twod = N/2; twod >= 1; twod /= 2)
        int twod1 = twod*2;
        parallel_for (int i = 0; i < N; i += twod1)
            int t = output[i+twod-1];
            output[i+twod-1] = output[i+twod1-1];
            // change twod1 below to twod to reverse prefix sum.
            output[i+twod1-1] += t;
    }
}
```

Figure 2: Iterative implementation of exclusive prefix sum

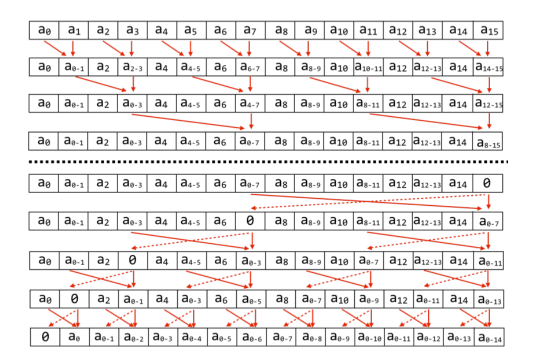


Figure 3: Visualization of exclusive prefix sum for 16 elements

You are welcome to use this general algorithm to implement a version of parallel prefix sum in CUDA. You must implement the exclusive\_scan() function in scan.cu. Your implementation will consist of both host and device code. The implementation will require multiple kernel launches.

**Note:** The scan implementation of Figure 2 assumes that the input array's length (N) is a power of 2. In the cudaScan() function in scan.cu (which calls your implementation of  $exclusive\_scan()$ ), we solve this problem by rounding the input array length to the next power of 2 when allocating the corresponding buffers on the GPU. However, we only copy back N elements from the GPU buffer back to the CPU buffer. This fact should simplify your CUDA implementation.

### **Implementing Find Repeats Using Prefix Sum**

Once you have written <code>exclusive\_scan()</code>, you should implement the function <code>find\_repeats()</code> in <code>scan.cu</code>. This will involve writing more device code, in addition to one or more calls to <code>exclusive\_scan()</code>. Your code should write the list of repeated indicies into the provided output array (in device memory), and then return the size of the output list.

When using your exclusive\_scan() implementation, remember that the contents of the start array are copied over to the output array. Also, the arrays passed to exclusive\_scan() are assumed to be in device memory.

**Grading:** We will test your code for correctness and performance on random input arrays.

For reference, a scan score table is provided below, showing the performance of a simple CUDA imple-

mentation on a Gates machine with a GTX 1080. To check the correctness and performance score of your scan() and find\_repeats() implementations, run the commands:

```
./checker.pl -m scan
./checker.pl -m find_repeats
```

Doing so will produce a reference table like the following:

Scan Score Table:

Element Count	Target Time	Your Time	Score	I
10000   100000   1000000   2000000	0.238   0.330   1.066   1.633	0.006 (F)   0.037 (F)   0.089 (F)   0.095 (F)	0   0   0   0	     
		Total score:	0/5	

Your score will be based solely on the performance of your code. In order to get full credit, it must perform within 20% of the provided target code.

**Test Harness:** For grading purposes, the test harness runs on a pseudo-randomly generated array that differs each time it is run. For debugging purposes, you may prefer a more stable environment. Passing the option "-i test1" to either checker.pl or cudaScan will use a fixed, randomly generated input. We encourage you to come up with alternate inputs to your program to help you evaluate it.

The argument "-t" to either checker.pl or your compiled program will use the Thrust Library implementation of exclusive scan. We will award up to two points of extra credit for anyone who can create an implementation that is competitive with Thrust.

# 3 A Simple Circle Renderer (85 pts)

Now for the real show!

The directory render of the assignment starter code contains an implementation of a renderer that draws colored circles. Build the code, and run the renderer with the command line "./render rgb." You will see an image of three circles appear on screen ('q' closes the window). Now run the renderer with the command line "./render snow." You should see an animation of falling snow.

The assignment starter code contains two versions of the renderer: a sequential, single-threaded C++ reference implementation, implemented in refRenderer.cpp, and an *incorrect* parallel CUDA implementation in cudaRenderer.cu.

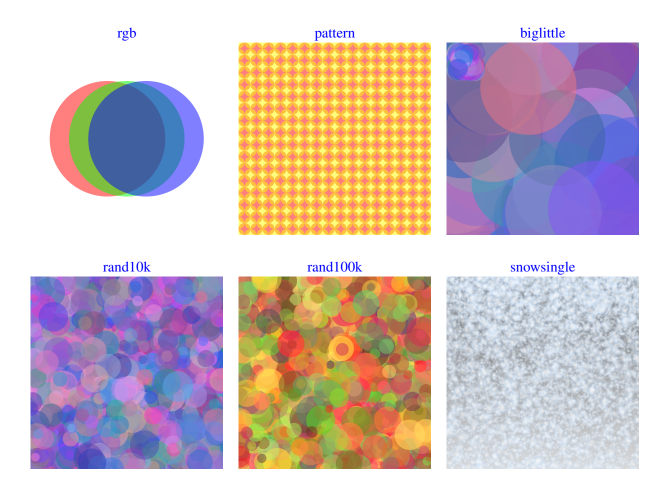


Figure 4: Sample images generated by renderer

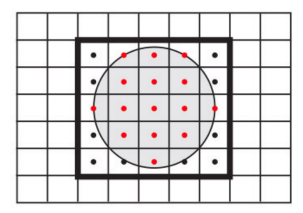


Figure 5: Computing the contribution of a circle to the output image

#### **Renderer Overview**

We encourage you to familiarize yourself with the structure of the renderer code base by inspecting the reference implementation in refRenderer.cpp. The method setup() is called prior to rendering the first frame. In your CUDA-accelerated renderer, this method will likely contain all your renderer initialization code (allocating buffers, etc). The render() method is called for each frame and is responsible for drawing all circles in the output image. The other main function of the renderer, advanceAnimation(), is also invoked once per frame. It updates circle positions and velocities. You will not need to modify advanceAnimation() in this assignment.

The renderer accepts an array of circles (3D position, velocity, radius, color) as input. The basic sequential algorithm for rendering each frame is:

Figure 5 illustrates the basic algorithm for computing circle-pixel coverage using point-in-circle tests. All pixels within the circle's bounding box are tested for coverage. For each pixel in the bounding box, the pixel is considered to be covered by the circle if its center point (shown with a dot) is contained within the circle.

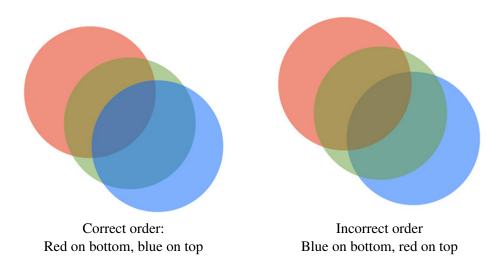


Figure 6: Ordering dependency of rendering

Pixel centers that are inside the circle are colored red, while those that are outside the circle are colored black. The circle's contribution to the image will be computed only for covered pixels.

An important detail of the renderer is that it renders **semi-transparent** circles. Therefore, the color of any one pixel is not the color of a single circle, but the result of blending the contributions of all the semi-transparent circles overlapping the pixel (note the "blend contribution" part of the pseudocode above). The renderer represents the color of a circle via a 4-tuple of red (R), green (G), blue (B), and opacity (alpha, written " $\alpha$ ") values (RGBA). Alpha value  $\alpha=1.0$  corresponds to a fully opaque circle. Alpha value  $\alpha=0.0$  corresponds to a fully transparent circle. To draw a semi-transparent circle with color  $(C_r, C_g, C_b, \alpha)$  on top of a pixel with color  $P_r, P_g, P_b$ , the renderer performs the following computation:

$$R_r = \alpha \cdot C_r + (1.0 - \alpha) \cdot P_r$$

$$R_g = \alpha \cdot C_g + (1.0 - \alpha) \cdot P_g$$

$$R_b = \alpha \cdot C_b + (1.0 - \alpha) \cdot P_b$$

Notice that composition is not commutative (object X over Y does not look the same as object Y over X), so it's important to render circles in a manner that follows the order they are provided by the application. (You can assume the application provides the circles in depth order.) For example, consider the two images shown in Figure 6, where the circles are ordered as red, green, and blue. The image on the left shows them rendered in the correct order, while the image on the right has them reversed.

#### **CUDA Renderer**

After familiarizing yourself with the circle rendering algorithm as implemented in the reference code, now study the CUDA implementation of the renderer provided in cudaRenderer.cu. You can run the CUDA implementation of the renderer using the command-line option "-r cuda."

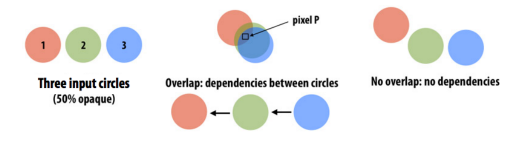


Figure 7: Illustration of ordering dependencies for rendering

The provided CUDA implementation parallelizes computation across all input circles, assigning one circle to each CUDA thread. While this CUDA implementation is a complete implementation of the mathematics of a circle renderer, it contains several major errors that you will fix in this assignment. Specifically, the current implementation does not ensure that image update is an atomic operation, and it does not preserve the required order of image updates, both of which are described below.

### **Renderer Requirements**

Your parallel CUDA renderer implementation must maintain two invariants that are preserved trivially in the sequential implementation:

- 1. **Atomicity:** All image update operations must be atomic. The critical region includes reading the four 32-bit floating-point values (the pixel's rgba color), blending the contribution of the current circle with the current image value, and then writing the pixel's color back to memory.
- 2. Order: Your renderer must perform updates to an image pixel in circle input order. That is, if circle 1 and circle 2 both contribute to pixel P, any image updates to P due to circle 1 must be applied to the image before updates to P due to circle 2. As discussed above, preserving the ordering requirement allows for a correct rendering of transparent circles. A key observation is that the definition of order only specifies the order of updates to an individual pixel. Thus, as shown in Figure 7, there is no ordering requirement between circles that do not contribute to the same pixel. These circles can be processed independently.

Since the provided CUDA implementation does not satisfy either of these requirements, the result of not correctly respecting order or atomicity can be seen by running the CUDA renderer implementation on the different scenes. You may see horizontal streaks through the resulting images, as shown in Figure 8 for the rgb scene. These streaks will change with each frame. If not streaks, you will see cases where the program renders the circles in the wrong order.

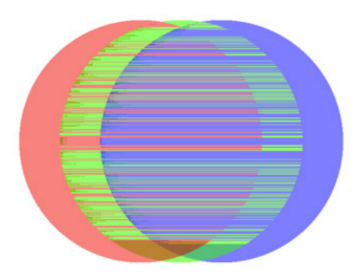


Figure 8: Incorrect rendering by provided CUDA implementation

#### What You Need To Do

Your job is to write the fastest, correct CUDA renderer implementation you can. You may take any approach you see fit, but your renderer must adhere to the atomicity and order requirements specified above. A solution that does not meet both requirements will be given no more than 10 points on part 2 of the assignment. We have already given you such a solution!

A good place to start would be to read through <code>cudaRenderer.cu</code> and convince yourself that it does not meet the correctness requirement. To visually see the effect of violation of above two requirements, compile the program with <code>make</code>. Then run:

```
./render -r cuda rand10k
```

Compare this image with the one generated by sequential code by running

```
./render rand10k
```

(This image is shown in the lower-left corner of Figure 4.)

You can get a listing of the options to the render program by running

```
./render --help
```

**Checker code:** To detect correctness of the program, render has a convenient "--check" option. This option runs the sequential version of the reference CPU renderer along with your CUDA renderer and then

<sup>&</sup>lt;sup>1</sup> There is one optimization you may *not* perform: you should render each frame independently, even if one frame contains the same circles as the previous one.

compares the resulting images to ensure correctness. The time taken by your CUDA renderer implementation is also printed.

There are a total of six circle data sets on which you will be graded for performance. However, in order to receive full credit, your code must pass the correctness test for all of our test data sets. To check the correctness and performance score of your code, run "make check" (or directly call the program checker.pl.) If you run it on the starter code, the program will print the results for the entire test set, plus a table like the following:

Score	e ta	ble:

Scene Name	Target Time	Your Time	Score	
rgb   rand10k   rand100k   pattern   snowsingle   biglittle	0.2393   2.7238   27.0537   0.3219   13.8742   21.0284	123.3879 (F)   39.4878 (F)   1045.4628 (F)   2.2350 (F)   10.5530 (F)   529.9166 (F)	0   0   0   0   0	       
		Total score:	0/72	

Note: on some runs, you *may* receive credit for some of these scenes, since the provided renderer's runtime is non-deterministic. This doesn't change the fact that the current CUDA renderer is incorrect.

"Target time" is the performance of a good solution on your current machine (in the provided render\_soln executable.) "Your time" is the performance of your CUDA renderer. Your grade will depend on the performance of your implementation compared to that of the provided implementation (see Grading Guidelines.)

Along with your code, we would like you to hand in a clear, high-level description of how your implementation works as well as a brief description of how you arrived at this solution. Specifically address approaches you tried along the way, and how you went about determining how to optimize your code (For example, what measurements did you perform to guide your optimization efforts?).

Aspects of your work that you should mention in the write-up include:

- 1. Include both partners' names and Andrew Id's at the top of your write-up.
- 2. Replicate the score table generated for your solution and specify which machine you ran your code on.
- 3. Describe how you decomposed the problem and how you assigned work to CUDA thread blocks and threads (and maybe even warps.)
- 4. Describe where synchronization occurs in your solution.

- 5. What, if any, steps did you take to reduce communication requirements (e.g., synchronization or main memory bandwidth requirements)?
- 6. Briefly describe how you arrived at your final solution. What other approaches did you try along the way. What was wrong with them?

### **Grading Guidelines**

- The write-up for the assignment is worth 13 points.
- Your implementation is worth 72 points. These are equally divided into 12 points per scene as follows:
  - 2 correctness points per scene.
  - 10 performance points per scene (only obtainable if the solution is correct). Your performance will be graded with respect to the target performance of a provided renderer,  $T_s$ .
  - No performance points will be given for solutions having time  $T > 10T_s$ .
  - Full performance points will be given for solutions within 20% of the optimized solution ( $T \leq 1.2T_s$ .)
  - For other values of T, your performance score on a scale 1 to 9 will be calculated as a linear interpolation based on the value of  $T_s/T$ .
- Up to five points extra credit (instructor discretion) for solutions that achieve significantly greater performance than required. Your write up must clearly explain your approach thoroughly.
- Up to five points extra credit (instructor discretion) for a high-quality parallel CPU-only renderer implementation that achieves good utilization of all cores and SIMD vector units of the cores. Feel free to use any tools at your disposal (e.g., SIMD intrinsics, ISPC, Pthreads.) To receive credit you should analyze the performance of your GPU and CPU-based solutions and discuss the reasons for differences in implementation choices made.

# **Assignment Tips and Hints**

Below are a set of tips and hints compiled from previous years. Note that there are various ways to implement your renderer and not all hints may apply to your approach.

- To facilitate remote development and benchmarking, we have created a "--bench" option to the render program. This mode does not open a display, and instead runs the renderer for the specified number of frames. (You'll see that the checking code runs with the command-line option "--bench 0:4.")
- When in benchmark mode, the "--file *Name*" command-line option sets the base file name for PPM images created at each frame. Created files have names of the form "*Name\_xxxx*.ppm," where *xxxx* is a 4-digit frame number. No PPM files are created if the --file option is not used.

- There are two potential axes of parallelism in this assignment. One axis is parallelism across pixels another is parallelism across circles (provided the ordering requirement is respected for overlapping circles.)
- The prefix-sum operation provided in exclusiveScan.cu\_inl may be valuable to you on this assignment (you are not required to use it.) It implements exclusive prefix-sum on a power-of-two-sized array in shared memory. The provided code does not work on non-power-of-two inputs and it also requires that the number of threads in the thread block be the size of the array.
- You are allowed to use the Thrust library in your implementation if you so choose. Thrust is not necessary to achieve the performance of the optimized CUDA reference implementations. See the documentation on prefix-sum operations in Thrust.
- Is there data reuse in the renderer? What can be done to exploit this reuse?
- The circle-intersects-box tests provided to you in circleBoxTest.cu\_inl are your friend.
- How will you ensure atomicity of image update since there is no CUDA language primitive that performs the logic of the image update operation atomically? Constructing a lock out of global memory atomic operations is one solution, but keep in mind that even if your image update is atomic, the updates must be performed in the required order. We suggest that you think about ensuring order in your parallel solution first, and only then consider the atomicity problem (if it still exists at all) in your solution.
- If you are having difficulty debugging your CUDA code, you can use printf directly from device code if you use a sufficiently new GPU and CUDA library: see this brief guide on how to print from CUDA.
- If you find yourself with free time, have fun making your own scenes!

#### **Catching CUDA Errors**

(Credit: The following was adapted from this Stack Overflow post)

By default, if you access an array out of bounds, allocate too much memory, or otherwise cause an error, CUDA won't normally inform you; instead it will just fail silently and return an error code. You can use the following macro (feel free to modify it) to wrap CUDA calls:

```
if (abort) exit(code);
}

#else
#define cudaCheckError(ans) ans
#endif
```

Note that you can undefine DEBUG to disable error checking once your code is correct for improved performance.

You can then wrap CUDA API calls to process their returned errors as such:

```
cudaCheckError( cudaMalloc(&a, size*sizeof(int)) );
```

Note that you can't wrap kernel launches directly. Instead, their errors will be caught on the next CUDA call you wrap:

```
kernel<<<1,1>>>(a); // suppose kernel causes an error!
cudaCheckError( cudaDeviceSynchronize() ); // error is printed on this line
```

All CUDA API functions, cudaDeviceSynchronize(), cudaMemcpy(), cudaMemset(), and so on can be wrapped.

**IMPORTANT:** if a CUDA function caused an error previously, but it wasn't caught, that error will show up in the next error check, even if the check wraps a different function. For example:

```
line 742: cudaMalloc(&a, -1); // executes, then continues
line 743: cudaCheckError(cudaMemcpy(a,b)); // Prints error for line 743
```

Therefore, while debugging, it's recommended that you wrap **all** CUDA API calls (at least in code that you wrote).

## **Hand-in Instructions**

You will submit your code via Autolab and your report via Gradescope. For the code, you will be submitting your entire directory tree.

#### 1. Your code

- (a) **If you are working with a partner, form a group on Autolab.** Do this before submitting your assignment. One submission per group is sufficient.
- (b) Make sure all of your code is compilable and runnable. We should be able to simply run make, then execute your programs in saxpy, scan, and render without manual intervention.

- (c) Remove all nonessential files, especially output images, from your directories.
- (d) Run the command "make handin.tar." This will run "make clean" and then create an archive of your entire directory tree.
- (e) Submit the file handin.tar to Autolab.

### 2. Your report

- (a) If this is your first time using Gradescope, go to https://gradescope.com and sign up for free with the entry code 9RW38E. Please register using your Andrew email address as id. If you already have a Gradescope account, the course should appear on your dashboard.
- (b) Please upload your report as file report.pdf to Gradescope, one submission per team, and select the appropriate pages for each part of the assignment. After submitting, you will be able to add your teammate using the *add group members* button on the top right of your submission.

Our grading scripts will rerun the checker code allowing us to verify your score matches what you submitted in the report.pdf. We might also try to run your code on other datasets to further examine its correctness.