An OpenACC Example Code for a C-based heat conduction code
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Introduction

Accelerators have been considered as the leading candidate by many scientific and technical programmers to program and accelerate huge complex scientific applications. Accelerators such as GPUs have immense potential in terms of high compute capacity but programming these devices is a challenge. CUDA, OpenCL and other vendor-specific models are definitely a way to go, but these are low-level models that demand excellent programming skills; moreover, they are time consuming to write and debug. There is a need to approach GPU computing from a higher-level. This is the motivation behind OpenACC, a portable and directive-based programming model that helps programmers identify compute-intensive areas of the code and instruct compilers to offload such computations to accelerators.

In this post, we will discuss how the programmers can use OpenACC to simplify parallel programming of heterogeneous CPU/GPU systems and yet achieve ‘good’ performance. We have considered a 2D heat conduction application that is a stencil-based code as the application to run on GPUs. We chose this code primarily because the computation/communication ratio is quite high. The computation portion is intensive and it can be distributed to several threads. The performance is already very good by using GPU global memory, and it can be further improved by using GPU shared memory.

2D Heat Conduction

The formula to represent 2D heat conduction is given as follows:

$$\frac{\partial T}{\partial t} = \alpha \left( \frac{\partial^2 T}{\partial x^2} + \frac{\partial^2 T}{\partial y^2} \right)$$

where $T$ is temperature, $t$ is time, $\alpha$ is the thermal diffusivity, and $x$ and $y$ are points in a grid.

To solve this problem, one possible finite difference approximation is:
where $\Delta T$ is the temperature change over time $\Delta t$ and $i, j$ are indices in a grid. In the beginning, the boundary points of the grid are assigned with initial temperature and the inner points update their temperature iteratively. Each inner point updates its temperature by using the previous temperature of its neighboring points and itself. The temperature updating operation for all inner points in a grid needs to last long enough which means many iterations is required to get the final stable temperatures. We fix the number of iterations to 20000. The grid has 1024*1024 points.

Following is the original code snippet of 2D heat where we have inserted an OpenMP pragma. $ni$ and $nj$ are X and Y dimensions of the grid, respectively. At the beginning, the initial temperatures of all points in the grid are stored in $temp1\_h$. $temp2\_h$ duplicates the temperatures in $temp1\_h$. After each iteration step, $temp2\_h$ stores the new temperatures and then we swap the pointer so that in the next iteration the input of the kernel points to the current new temperatures. Because after all the iterations are completed, the pointers still need to be swapped. The final temperatures are stored in $temp1\_h$.

```c
void step_kernel(int ni, int nj, float fact,
    float* temp_in, float* temp_out)
{
    #pragma omp parallel for
    // loop over all points in domain (except boundary)
    for (j=1; j < nj-1; j++) {
        for (i=1; i < ni-1; i++) {
            // find indices into linear memory
            // for central point and neighbours
            i00 = I2D(ni, i, j);
            im10 = I2D(ni, i-1, j);
            ip10 = I2D(ni, i+1, j);
            i0m1 = I2D(ni, i, j-1);
            i0p1 = I2D(ni, i, j+1);
            // evaluate derivatives
            d2tdx2 = temp_in[im10]-2*temp_in[i00]+temp_in[ip10];
            d2tdy2 = temp_in[i0m1]-2*temp_in[i00]+temp_in[i0p1];
            // update temperatures
            temp_out[i00] = temp_in[i00]+tfac*(d2tdx2 + d2tdy2);
        }
    }
}

int main(int argc, char* argv[])
Parallelizing using OpenMP and OpenACC:

We will parallelize this program using both OpenMP and OpenACC and compare their performances to the serial code.

This program is running on a system consisting of two quad-core Intel Xeon x86_64 CPU (2.27GHz) with one Nvidia Tesla K20 GPU.

**STEP 1:**
In the first step we try to optimize this code for a multicore system. Here we insert OpenMP parallel for directive before the nested loop in step_kernel. This directive tells the compiler to distribute the work among all threads set in OMP_NUM_THREADS environment variable. We use the commonly used host compiler GCC to compile this program.

```
$ module load gcc/4.4.7
$ gcc -fopenmp -O3 -o heat heat.c
```

The following table shows the performance and speedup of OpenMP code compared to the serial code. It does not show significant speedup because the kernel where we have inserted the OpenMP parallel for directive is inside the main loop, therefore the overhead incurred is quite high; creating and destroying threads when entering and exiting the parallel region respectively.
<table>
<thead>
<tr>
<th>Program version</th>
<th>Execution time (s)</th>
<th>Speedup compared to serial</th>
</tr>
</thead>
<tbody>
<tr>
<td>Serial</td>
<td>134.04</td>
<td>1</td>
</tr>
<tr>
<td>OpenMP 2 threads</td>
<td>112.74</td>
<td>1.19</td>
</tr>
<tr>
<td>OpenMP 4 threads</td>
<td>66.44</td>
<td>2.02</td>
</tr>
<tr>
<td>OpenMP 8 threads</td>
<td>44.33</td>
<td>3.02</td>
</tr>
</tbody>
</table>

**STEP 2:**
In this step we will see if we can improve the performance by using OpenACC directives. The compilers that support OpenACC include HMPP, PGI and Cray. Here we will show how to parallelize the code with PGI’s compiler. First we simply insert the following one line directive:

```c
#pragma acc kernels copyin(temp_in[0:ni*nj])
copy(temp_out[0:ni*nj])
```

The directive tells the compiler to convert the following loop nest into a kernel that will be executed on the accelerator (in this case GPU is the accelerator). We also specify that data `temp_in` should be copied to the device before the kernel execution but not copied back to the host after the kernel execution. The data `temp_out` needs to be both copied to the device before the kernel execution and copied back to the host after the kernel execution. The reason that `temp_out` also needs to be copied to the device is that the kernel only updates the inner points value, while `temp_out` also includes boundary points values. If we just use `copyout(temp_out)`, then the boundary points values that transferred to the host would be garbage values. The following command is used to compile this program:
Here “-acc” flag tells the compiler to enable OpenACC and “-ta=nvidia,cuda5.0,cc35” tells the compiler that the target accelerator is NVIDIA GPU. The cuda version is 5.0 and the compiler will generate the executable for the GPU and that its computation capability is 3.5. Note that this number needs to be supported by both the PGI compiler (13.1) and the GPU hardware (K20).

The flag “-Minfo=accel” helps to view the compilation messages about the parallelization that is taking place using OpenACC directives. The flat “-Msafeptr” is used in order to remove the pointer alias issue. Because this program is very sensitive to the order of floating point operations and PGI applies some aggressive optimization in “-O3” level, the result would be slightly different from the serial version compiled by GCC. NVIDIA GPU uses Fused Multiply-Add (FMA) by default to fuse the operations of multiply and add into one operation. Using FMA for this application also delivers result different from the serial version. For easy comparison of the output produced by different versions of the program, we disable optimizations and simply use “-O0” with PGI compiler and disable FMA by passing the “nofma” option. Note that this does not mean that the GPU result is wrong. We notice that the result using FMA appeared to be more accurate than the result that does not use FMA.

For details about FMA, you can refer to the paper “Precision & Performance: Floating Point and IEEE 754 Compliance for NVIDIA GPUs”.

```bash
$module load cuda/5.0
$module load pgi/13.1
$ pgcc -acc -ta=nvidia,cuda5.0,nofma,cc35 -Minfo=accel -Msafeptr -O0 -o heat heat.c
```
After setting all of the above-mentioned flags, the compilation messages show that there are loop carried dependences in the nested loop; interestingly actually there are no dependencies in the loop. This is because we are using a computed index in a loop that the compiler is trying to automatically schedule. So we add the following directive after “acc kernels” line to give the compiler more hints:

```
#pragma acc loop collapse(2) independent
```

Now the compilation message shows the following: (we notice that the compiler no longer complains that there are loop-carried dependencies in the loop)

```
step_kernel:
  26, Generating present_or_copy(temp_out[0:nj*ni])
  Generating present_or_copyin(temp_in[0:nj*ni])
  Generating NVIDIA code
  Generating compute capability 3.5 binary
  29, Loop is parallelizable
  30, Loop is parallelizable
      Accelerator kernel generated
      29, #pragma acc loop gang /* blockIdx.y */
      30, #pragma acc loop gang, vector(128) /* blockIdx.x threadIdx.x */
```

This means that both the loops can be parallelized by and the PGI compiler automatically uses its own scheduling policy for the number of gangs and the vector length. By executing this program, we get the following:

```
$ ./heat
Time for computing: 94.92 s
```

However this is only slightly faster than the OpenMP version with 2 threads.

Let us try to find out why this is happening!

**STEP 3:**

We need performance profiling tools. We can either use NVIDIA’s command line profiler nvprof or by setting “export PGI_ACC_TIME=1” when running the program.

The output of using nvprof is as follows:

```
$ nvprof ./heat
====== NVPROF is profiling heat...
====== Command: heat
Time for computing: 109.14 s
====== Profiling result:
```
<table>
<thead>
<tr>
<th>Name</th>
<th>Time(%)</th>
<th>Time</th>
<th>Calls</th>
<th>Avg</th>
<th>Min</th>
<th>Max</th>
</tr>
</thead>
<tbody>
<tr>
<td>[CUDA memcpy HtoD]</td>
<td>58.70</td>
<td>40.98s</td>
<td>40000</td>
<td>1.02ms</td>
<td>1.01ms</td>
<td>1.19ms</td>
</tr>
<tr>
<td>[CUDA memcpy DtoH]</td>
<td>38.36</td>
<td>26.78s</td>
<td>20000</td>
<td>1.34ms</td>
<td>1.30ms</td>
<td>1.43ms</td>
</tr>
<tr>
<td>step_kernel_30_gpu</td>
<td>2.94</td>
<td>2.05s</td>
<td>20000</td>
<td>102.69us</td>
<td>101.95us</td>
<td>104.26us</td>
</tr>
</tbody>
</table>

Or by using

```bash
$ export PGI_ACC_TIME=1
```

Time for computing: 98.83 s

Accelerator Kernel Timing data
/home/rengan/GTC13/heat.c
  step_kernel NVIDIA devicenum=0
    time(us): 85,290,940
      26: data copyin reached 40000 times
          device time(us): total=51,427,629 max=9,224
          min=1,246 avg=1,285
    30: kernel launched 20000 times
        grid: [8x1022] block: [128]
          device time(us): total=2,378,772 max=1,250
          min=115 avg=118
          elapsed time(us): total=2,608,648 max=1,264
          min=124 avg=130
    47: data copyout reached 20000 times
        device time(us): total=31,484,539 max=2,519
        min=1,528 avg=1,574

Both the profiling tools show that the main performance bottleneck is due to the expensive data transfer cost.

Nvprof found that the data transfer time takes up to 97.06% of the total execution time. To reduce the data transfer time, we should try to keep the data on the GPU as long as possible, and here it is possible to do that. The solution is to add a data region outside the main loop. Note that this time `temp1_h` needs to copyin and copyout and `temp2_h` only needs to copyin. This is because after all the iterations, the pointers of `temp1_h` and `temp2_h` are swapped, and the final result is stored in `temp1_h`. Because the pointer `temp_tmp` is used just for swapping other two pointers, we make it as a device pointer so that the pointer swapping also happens on the device side.

```c
#pragma acc data copy(temp1_h[0:ni*nj]) \
    copyin(temp2_h[0:ni*nj]) \
    deviceptr(temp_tmp)
{
    // main iteration loop
```
for (istep=0; istep < nstep; istep++) {
    // CPU kernel
    step_kernel(ni, nj, tfac, temp1_h, temp2_h);
    // swap the temp pointers
    temp_tmp = temp1_h;
    temp1_h = temp2_h;
    temp2_h = temp_tmp;
}

In the kernel, the directive:

#pragma acc kernels copyin(temp_in[0:ni*nj])
copy(temp_out[0:ni*nj])

is changed to

#pragma acc kernels
present(temp_in[0:ni*nj],temp_out[0:ni*nj])

since the data in temp_in and temp_out are already present in GPU.

After compiling, the new compilation message shows:

step_kernel:
  26, Generating present(temp_out[0:nj*ni])
  Generating present(temp_in[0:nj*ni])
  Generating NVIDIA code
  Generating compute capability 3.5 binary
  29, Loop is parallelizable
  30, Loop is parallelizable
  Accelerator kernel generated
  29, #pragma acc loop gang /* blockIdx.y */
  30, #pragma acc loop gang, vector(128) /* blockIdx.x threadIdx.x */
main:
  116, Generating copyin(temp2_h[0:nj*ni])
  Generating copy(temp1_h[0:nj*ni])

By profiling again, we get:

$ nvprof ./heat
======== NVPROF is profiling heat...
====== Command: heat
Time for computing: 4.32 s
======== Profiling result:
And after setting “PGI_ACC_TIME=1”, we get the following:

Time for computing: 2.59 s

Accelerator Kernel Timing data
/home/rengan/GTC13/heat.c
step_kernel  NVIDIA  devicenum=0
time(us): 2,077,249
   30: kernel launched 20000 times
     grid: [8x1022]  block: [128]
     device time(us): total=2,077,249 max=862
min=100  avg=103
   elapsed time(us): total=2,310,052 max=874
min=111  avg=115
/home/rengan/GTC13/heat.c
main  NVIDIA  devicenum=0
time(us): 4,305
   116: data copyin reached 2 times
     device time(us): total=2,809 max=1,449
min=1,360  avg=1,404
   131: data copyout reached 1 times
     device time(us): total=1,496 max=1,496
min=1,496  avg=1,496

The profiling results show that the data transfer bottleneck does not exist any more, almost all the time is spent only in the computation part. And the data copy time is reduced from 60000 times to 3 times.

To reduce the profiling overhead, we run the program without any profiling tool and get the following:

$ ./heat
Time for computing: 2.42 s

Finally we once again compare the performances obtained by OpenACC against the OpenMP and the serial version. The following table highlights the performance numbers.

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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>serial</td>
<td></td>
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<tr>
<td>------------</td>
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<td></td>
</tr>
<tr>
<td>OpenACC</td>
<td>2.42</td>
<td></td>
</tr>
<tr>
<td></td>
<td>55.39</td>
<td></td>
</tr>
</tbody>
</table>

We see that porting a code using OpenACC is not very hard. Of course, the complexity of the code matters. With the help of profiling tools, we can check the bottleneck of the current performance, and then we could perform optimizations of the code incrementally. In this example, we get more than 55x compared to the serial version and more than 18x (55.39/3.02) compared to the OpenMP version using 8 threads. We also learnt a very important lesson that while programming GPUs, the performance bottleneck is usually due to the data transfer part. This can prove to be quite challenging when the complexity of the application grows.