Assignment 6- Hybrid MPI + MP + CUDA Programming CSCI 596: Scientific Computing & Visualization

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November 1, 2017

The goal of this assignment is understand the basics of cuda through two examples—for calculating the pair distribution function, and to calculate π .

1 Task I- Pair-Distribution Computation with CUDA

The pair distribution function (PDF) is the probability of finding any two atoms, which are a given distance r apart, in a given volume. It is popularly used to measure the order, especially crystalline materials.

To compute the pdf, we fist calculate the histogram of atomic pair distances, i.e., calculate the distance between all pairs of atoms i and j, say r_{ij} , put them in the appropriate 'bins' Δr to generate a histogram. Note that the maximum distance between pairs under periodic boundary condition is the diagonal of half the simulation box. The algorithm is as follows:

for all histogram bins i

nhist[i] = 0

for all atomic pairs (i,j)

++nhist $[r_{ij}/r]$

Once this histogram is obtained, the pdf may then be computed using the following formula:

$$g(r_k) = \frac{\text{nhist}(k)}{2\pi r_k^2 \Delta r \rho N} \tag{1}$$

where r_k is the distance between two atoms, ρ is the number density of atoms, N is the total number of atoms. Notice that the obtaining the histogram involves computing distance between all pairs of atoms, which can get expensive with a large number of atoms. The parallel processing power of the GPUs was used to carry out these computations.

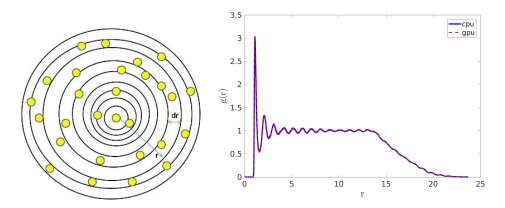


Figure 1: (a)Schematic for obtaining the histogram (b)PDF computation—CPU vs GPU

The code is attached in Appendix A. The pair distribution function computed using both the CPU and the GPUs shown for comparison in Fig 1 (the trailing off after r = 14 is explained in the notes.). The cpu implentation ran in 2.62 seconds whereas the gpu-parallel implementation ran in 0.28 seconds (almost 10 times faster!).

2 Task II– Parallel Computation of π using MPI+OMP+CUDA

In this part, a triple-decker MPI+OpenMP+CUDA program was written to compute the value of π , by modifying the double-decker MPI+CUDA program provided in class, described in the lecture note on "Hybrid MPI+OpenMP+CUDA Programming".

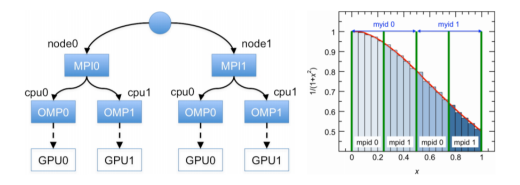


Figure 2: Output of program for computation of π

This program runs on two CPU cores, and 2 GPU devices on each compute node. This is achieved by launching one MPI rank per node, where each rank spawns two OpenMP threads, and each thread uses a different GPU device. This is illustrated in Fig 2. The above OpenMP multithreading introduces a race condition for variable pi. This can be circumvented by data privatization using reduction(+:pi), which creates a pi variable for each thread. The '+:' means that the 'addition reduction' is to be performed on this variable.

The program can be found in appendix B, and the output of the program is as shown below in Fig 2. myid refers to the process rank, mpid is the thread number within the processor, and device used refers to the gpu.

```
[kanale@hpc3026 assgn6]$ mpirun -np 2 -machinefile nodefile ./pi --bind-to none
[hpc3026:44118] mca: base: component_find: unable to open /usr/usc/openmpi/1.6.4/lib/openmpi/mca_btl_mx: libmyriexpress.
so: cannot open shared object file: No such file or directory (ignored)
[hpc3027:07438] mca: base: component_find: unable to open /usr/usc/openmpi/1.6.4/lib/openmpi/mca_btl_mx: libmyriexpress.
so: cannot open shared object file: No such file or directory (ignored)
myid = 0; mpid = 0: device used = 0; partial pi = 0.979926
myid = 0; mpid = 1: device used = 1; partial pi = 0.874671
myid = 1; mpid = 1: device used = 1; partial pi = 0.567582
myid = 1; mpid = 0: device used = 0; partial pi = 0.719409
PI = 3.141588
```

Figure 3: Output of program for computation of π

A Appendix- Calculating the PDF

Note that there were no major changes in the main() function, so it is not shown.

```
2 Program pdf0.c computes a pair distribution function for n atoms
 3 given the 3D coordinates of the atoms.
 5 #include <stdio.h>
 6 \# include < math.h >
 7 #include <time.h>
 8 #include < stdlib . h>
 9
10 #define NHBIN 2000 // Histogram size
12 float al [3];
                        // Simulation box lengths
                         // Number of atoms
13 int n;
                         // Atomic position array
14 float *r;
15 FILE *fp;
16
17 __constant__ float DALTH[3];
18 __constant__ int DN;
19 __constant__ float DDRH;
20
21 \ \_\_device\_\_ \ \mathbf{float} \ d\_SignR(\mathbf{float} \ v, \mathbf{float} \ x) \ \{\mathbf{if} \ (x > 0) \ \mathbf{return} \ v; \ \mathbf{else}
      return -v;
22
23
24 __global__ void gpu_histogram_kernel(float *r, float *nhis) {
25
    int i, j, a, ih;
26
     float rij, dr;
27
28
     int iBlockBegin = (DN/gridDim.x)*blockIdx.x;
29
     int iBlockEnd = min((DN/gridDim.x)*(blockIdx.x+1),DN);
     int iBlockBegin = (DN/gridDim.y)*blockIdx.y;
30
     int jBlockEnd = min((DN/gridDim.y)*(blockIdx.y+1),DN);
31
32
     for (i=iBlockBegin+threadIdx.x; i<iBlockEnd; i+=blockDim.x) {
       for (j=jBlockBegin+threadIdx.y; j<jBlockEnd; j+=blockDim.y) {
33
34
         if (i < j) 
         // Process (i,j) atom pair
35
36
         rij = 0.0;
37
              for (a=0; a<3; a++)
38
                dr = r[3*i+a]-r[3*j+a];
                /* Periodic boundary condition */
39
40
                   dr-d SignR(DALTH[a], dr-DALTH[a])-d SignR(DALTH[a], dr+DALTH[a]);
41
                rij += dr*dr;
42
```

```
rij = sqrt(rij); /* Pair distance */
43
44
             ih = rij/DDRH;
             // nhis[ih] += 1.0; /* Entry to the histogram */
45
46
             atomicAdd(&nhis[ih],1.0);
         \} // end if i<i
47
       \} // end for j
48
    \} // end for i
49
50 }
51
52
54 void histogram() {
55 /*---
56 Constructs a histogram NHIS for atomic-pair distribution.
57 -
     float alth [3];
58
59
     float* nhis; // Histogram array
60
     float rhmax, drh, density, gr;
61
     int a, ih;
62
63
     float * dev r; // Atomic positions
64
     float * dev nhis; // Histogram
65
    /* Half the simulation box size */
66
    for (a=0; a<3; a++) alth [a] = 0.5*al[a];
67
    /* Max. pair distance RHMAX & histogram bin size DRH */
68
    rhmax = sqrt(alth[0]*alth[0]+alth[1]*alth[1]+alth[2]*alth[2]);
69
     drh = rhmax/NHBIN; // Histogram bin size
70
71
72
73
     nhis = (float*) malloc(sizeof(float)*NHBIN);
    // for (ih=0; ih < NHBIN; ih++) nhis[ih] = 0.0; // Reset the histogram
74
75
    cudaMalloc((void**)&dev r, sizeof(float)*3*n);
76
77
     cudaMalloc((void**)&dev nhis, sizeof(float)*NHBIN);
    cudaMemcpy(dev r,r,3*n*sizeof(float),cudaMemcpyHostToDevice);
78
79
    cudaMemset(dev nhis,0.0,NHBIN*sizeof(float));
    cudaMemcpyToSymbol(DALTH, alth, sizeof(float)*3,0,cudaMemcpyHostToDevice);
80
    cudaMemcpyToSymbol(DN,&n, sizeof(int),0,cudaMemcpyHostToDevice);
81
    cudaMemcpyToSymbol(DDRH,&drh, sizeof(float), 0, cudaMemcpyHostToDevice);
82
83
84
    \dim 3 \text{ numBlocks}(8,8,1);
85
    dim3 threads per block (16,16,1);
    gpu histogram kernel <<< numBlocks, threads per block >>> (dev r, dev nhis);
86
87
88
    // Compute dev nhis on GPU: dev r[] Âő dev nhis []
89
```

```
90
     cudaMemcpy(nhis, dev nhis, NHBIN*sizeof(float), cudaMemcpyDeviceToHost);
91
92
     density = n/(al[0]*al[1]*al[2]);
93
     /* Print out the histogram */
     fp = fopen("pdf gpu.d", "w");
94
     for (ih=0; ih < NHBIN; ih++) {
95
       gr = nhis[ih]/(2*M PI*pow((ih+0.5)*drh,2)*drh*density*n);
96
        fprintf(fp, "%e %e\n", (ih+0.5)*drh, gr);
97
98
99
     fclose (fp);
100
     free (nhis);
101 }
```

pdf1writeup.cu

B Appendix Computation of π

```
1 // Hybrid MPI+CUDA computation of Pi
2 #include <stdio.h>
3 #include <mpi.h>
4 #include < cuda.h>
5 \# include < omp.h >
7 \# define NUM DEVICE 2 // \# of GPU devices = \# of OpenMP threads
8 \# define NBIN 10000000 // Number of bins
9 #define NUM BLOCK
                           // Number of thread blocks
                       13
10 #define NUM THREAD 192 // Number of threads per block
11
12 // Kernel that executes on the CUDA device
13 global void cal pi(float *sum, int nbin, float step, float offset, int
     nthreads, int nblocks) {
14
    int i;
15
     float x;
    int idx = blockIdx.x*blockDim.x+threadIdx.x; // Sequential thread
16
        index across the blocks
    for (i=idx; i<nbin; i+=nthreads*nblocks) { // Interleaved bin
17
        assignment to threads
      x = offset + (i+0.5)*step;
18
19
      sum[idx] += 4.0/(1.0+x*x);
    }
20
21 }
22
23 int main(int argc, char **argv) {
     int myid, nproc, nbin, tid, dev_used, mpid;
24
25
     float step, offset, pi=0.0, pig;
26
    dim3 dimGrid (NUM_BLOCK, 1, 1); // Grid dimensions (only use 1D)
27
    dim3 dimBlock (NUM THREAD, 1, 1); // Block dimensions (only use 1D)
```

```
float *sumHost, *sumDev; // Pointers to host & device arrays
28
29
30
     MPI Init(&argc,&argv);
31
    MPI Comm rank(MPI COMM WORLD,&myid); // My MPI rank
32
    MPI Comm size (MPI COMM WORLD, & nproc); // Number of MPI processes
     omp set num threads(NUM DEVICE);
33
     nbin = NBIN/(nproc*NUM DEVICE); // Number of bins per MPI process
34
     step = 1.0/(float)(nbin*nproc*NUM DEVICE); // Step size with
35
        redefined number of bins
36
37
    #pragma omp parallel private (mpid, offset, sumHost, tid) reduction (+:pi)
38
39
         mpid = omp get thread num();
         offset = (NUM DEVICE*myid+mpid)*step*nbin; // Quadrature-point
40
             offset
41
42
         cudaSetDevice (mpid%2);
         size t size = NUM BLOCK*NUM THREAD*sizeof(float); //Array
43
            memory size
         sumHost = (float *) malloc(size); // Allocate array on host
44
         \verb|cudaMalloc| ((\verb|void **|) \& \verb|sumDev|, \verb|size|); // Allocate array on device|
45
         {\tt cudaMemset(sumDev,0,size);} \hspace*{0.2cm} / / \hspace*{0.2cm} \textit{Reset array in device to } 0
46
47
         // Calculate on device (call CUDA kernel)
         cal pi <<<dimGrid, dimBlock>>>
48
             (sumDev, nbin, step, offset, NUM THREAD, NUM BLOCK);
         // Retrieve result from device and store it in host array
49
         cudaMemcpy(sumHost, sumDev, size, cudaMemcpyDeviceToHost);
50
         // Reduction over CUDA threads
51
         for (tid=0; tid <NUM THREAD*NUM BLOCK; tid++)</pre>
52
53
           pi += sumHost[tid];
         pi *= step;
54
         // CUDA cleanup
55
56
         free (sumHost);
         cudaFree(sumDev);
57
58
         // Reduction over MPI processes
59
         cudaGetDevice(&dev used);
60
         printf("myid = %d; mpid = %d: device used = %d; partial pi =
61
            %f \ n'', myid, mpid, dev used, pi);
62
     } // End omp parallel
     MPI Allreduce(&pi,&pig,1,MPI FLOAT,MPI SUM,MPI COMM WORLD);
63
     if (myid==0) printf("PI = \%f \setminus n", pig);
64
65
     MPI Finalize();
     return 0:
66
67 }
```

piCudaMpiOmpwriteup.cu