Parallel Computing/Programming Assignment #4 and #5: Massively Parallel Game of Life with MPI, Pthreads and Parallel I/O

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1 Assignment Description

This is a "double" assignment and so your grade will be counted twice (for assignment #4 and #5). You can have a team of up to 3 students.

For this assignment, your are to create an MPI implementation of *Conway's Game of Life* to leverage Pthreads within compute nodes and conduct a performance study on the IBM Blue Gene/Q. In particular, you will determining how much performance improvement is obtained by using Pthreads for intra-node shared-memory parallel vs. intra-node message passing using MPI as well as document what is the observed I/O performance.

1.1 Review: Basic Rules

The Game of Life is an example of a Cellular Automata where universe is a two-dimensional orthogonal grid of square cells (with WRAP AROUND FOR THIS ASSIGNMENT), each of which is in one of two possible states, *ALIVE* or *DEAD*. Every cell interacts with its eight neighbors, which are the cells that are horizontally, vertically, or diagonally adjacent. At each step in time, the following transitions occur at each and every cell:

- Any live cell with fewer than two live neighbors dies, as if caused by under-population.
- Any live cell with two or three live neighbors lives on to the next generation.
- Any live cell with more than three live neighbors dies, as if by over-population.

• Any dead cell with exactly three live neighbors becomes a live cell, as if by reproduction.

The initial pattern sets the entire universe to the ALIVE state. The first generation is created by applying the above rules to every cell in the seedbirths and deaths occur simultaneously, and the discrete moment at which this happens is sometimes called a "tick" The rules continue to be applied repeatedly to create further generations.

Note, a "tick" starts with Cell(0,0) and ends with Cell(N-1,N-1) in the serial case. When we get to parallel things will get much more interesting! (see below).

1.2 Adding Additional Randomness

Using the uniform distribution provided in RNG (see the template on how to use this new RNG), if value return by the RNG is greater than a set THRESHOLD, then perform the above described basic rules for each cell. Otherwise, randomly pick state of *LIVE* or *DEAD*. So if the THRESHOLD is 25%, then state updates will be made at random 25% of the time and follow the rules 75% of the time.

1.3 Parallelization Approach

To ensure that there is more true "randomness" among the cells, each row will have it's own RNG seed set. According to the template, every MPI rank will initialize 32,768 RNG streams. These streams are about 2^{70} calls apart. **SO, EACH ROW WILL HAVE IT'S OWN RNG STREAM! This is important for your Pthreads implementation.** When accessing a particular RNG stream, make sure you provide the ROW's global index and not the MPI rank's local array index. That is will be accessed using $index = local_row + (rows_per\ rank * my_mpi_rank)$

For the parallelization approach, a Blue Gene/Q compute node for example will allocate 1 MPI rank only per node and then create 63 additional threads per nodes. All threads on each node (including the MPI rank-thread) will perform an even "chunk" of rows for the Cellular Automata universe. For example, suppose you have a 1024x1024 universe using 64 threads using a single Blue Gene/Q node. If you divide the universe equally, each thread will have 16 rows or 16x1024 cells to compute. Thus, thread 0 (on node 0), will compute rows 0 to 15, thread 1 computes rows 16 to 31 and thread 2 will compute rows 32 to 47 and so on.

Now when running using more than one Blue Gene/Q node, each MPI rank only needs to allocate it's specific chunk (which is operated on by all threads within that compute node) plus space for "ghost" rows at the MPI rank boundaries (not thread boundaries). The "ghost" rows can be held outside of the main MPI rank's Cellular Automata universe.

Now, you'll notice that rows at the boundaries of MPI ranks need to be updated / exchanged prior to the start of computing each tick. For example, using the above 1024x1024 universe and 16 MPI ranks as an example, MPI rank 0 will need to send the state of row 0 (all 1024 entries) to MPI rank 15. Additionally, rank 15 will need to send row 1023 to rank 0. Also, rank 0 and rank 1 will do a similar exchange.

Note - the above example is only a 1024×1024 universe and below you will be using the larger $32,768 \times 32,768$ cell universe.

For these row exchanges, you will use the MPI_Isend and MPI_Irecv messages. You are free to design your own approach to "tags" and how you use these routines except your design should not deadlock.

1.4 Parallel I/O

When indicated for a particular experiment, you are to write out the cell universe into shared single file and time the length it took to complete that operation. All MPI ranks will participate using the MPI_file_write_at operation. Make sure your implementation performs the MPI_file_open using the MPI_COMM_WORLD communicator to ensure the file pointer is properly shared across all MPI ranks. With 128 ranks for the 32Kx32K universe, each rank will be responsible for 8,388,608 cells. You will want to make sure you compute each MPI rank's offset correctly using the cell data size, number of cells in the universe per rank and the rank's value.

1.5 Algorithm

So the algorithm becomes:

```
main(...)
   {
      Setup MPI (template has this already!)
      if rank 0 / thread 0, start time with GetTimeBase().
      Allocate My rank's chunk of the universe +
         space for "ghost" rows.
      Initalize the universe (including "ghost" rows) with every
         cell being ALIVE.
      Create Pthreads here. All threads should go into the for-loop.
      for (i = 0; i < number of ticks; <math>i++)
            Exchange row data with MPI ranks
               using MPI_Isend/Irecv from thread 0 w/i each MPI rank.
               Yes, you must correctly MPI_Test or Wait to make sure
               messages operations correctly complete.
            [Note: have only 1 MPI rank/pthread perform ALL MPI
             operations per rank/thread group. Dont' allow multiple
             threads to perform any MPI operations within MPI
             rank/thread group.]
            HERE each PTHREAD can process a row:
             - update universe making sure to use the
               correct row RNG stream
             - factor in Threshold percentage as described
             - use the right "ghost" row data at rank boundaries
             - keep track of total number of ALIVE cells per tick
               across all threads w/i a MPI rank group.
             - use pthread_mutex_trylock around shared counter
               variables **if needed**.
         }
```

```
MPI_Reduce( Sum all ALIVE Cells Counts For Each Tick);
    - Here, you will have vector of 256 ALIVE cell sum
      values which is the total number of ALIVE cells
      at each tick, t for all 256 ticks.
if rank 0 / thread 0, end time with GetTimeBase().
if needed by experiment,
   perform output of 32Kx32K cell universe using MPI_file_write_at;
   collect I/O performance stats using GetTimeBase() from
    rank 0 / thread 0;
if needed by experiment,
   construct 1Kx1K heatmap of 32Kx32K cell universe using MPI
   collective operations of your choice. Rank 0 will output
   the heatmap to a standard Unix file given it's small 1 to 4MB size.
   Make sure you an import data for graphing.
if rank 0, print ALIVE tick stats and compute (I/O if needed)
   performance stats.
MPI_Finalize();
```

A final note: your program is NOT parallel deterministic. That means that your Cellular Automata universe will differ at each step/tick in the computation depending on how many MPI ranks you use. This is because the exchange of rows is using data computed in a prior tick and does not get the benefit of the update dependencies that happen within a "tick".

Also, to link with the Pthread lib, make sure you add -lpthread to your compile command.

2 Experiments

}

You will conduct the following series of performance experiments.

2.1 Strong Scaling with Parallel I/O

a strong scaling experiment using "AMOS", the CCI Blue Gene/Q system. Here, you will have a mix of MPI ranks and threads per compute node. Below are the experiments that would be run just for 4 compute nodes on the Blue Gene/Q system.

- Run a 32768x32678 cell universe using 256 total MPI ranks, 0 Pthreads per rank on 4 BG/Q nodes for 256 ticks with a threshold of 25%.
- Run a 32768x32768 cell universe using 64 total MPI ranks, 4 Pthreads per rank on 4 BG/Q nodes for 256 ticks with a threshold of 25%.

- Run a 32768x32768 cell universe using 16 total MPI ranks, 16 Pthreads per rank on 4 BG/Q nodes for 256 ticks with a threshold of 25%.
- Run a 32768x32768 cell universe using 8 total MPI ranks, 32 Pthreads per rank on 4 BG/Q nodes for 256 ticks with a threshold of 25%.
- Run a 32768x32768 cell universe using 4 total MPI ranks, 64 Pthreads per rank on 4 BG/Q nodes for 256 ticks with a threshold of 25%.

Note, the MPI rank also counts as a Pthread within a compute node.

Now, re-perform the above experiments using 16, 64 and 128 compute nodes with the SAME RATIO of Pthreads to MPI ranks across all compute nodes used in the experiment. That is, you have runs of:

- 0 threads per MPI rank (64 MPI ranks per compute node).
- 4 threads per MPI rank (16 MPI ranks per compute node, rest are threads).
- 16 threads per MPI rank (4 MPI ranks per compute node, rest are threads).
- 32 threads per MPI rank (2 MPI ranks per compute node, rest are threads).
- 64 threads per MPI rank (1 master MPI rank per compute node but also is a thread).

In total, you will have 20 experiment with 5 experiments for each of 4, 16, 64 and 128 compute nodes. For the 128 compute node configuration, turn on the parallel I/O and output the cell universe and record the time spent performing I/O as well.

In terms of performance plots/graphs, first plot the number of ALIVE cells (Y-axis) as a function of the tick number (X-axis) for 5 experiments in the 128 node and 25% threshold case. Do you observe a significant change in ALIVE cells as a function of the tick number or as a function of the number of MPI ranks/threads?

Next, plot **compute** execution time (Y-axis) as a function of the total number of MPI ranks * Pthreads per rank (X-axis) grouping each threads per MPI rank as a graph line. Do you observe any performance differences as the threads per MPI rank changes? Also, in your report, discuss the max speedup relative to the execution time for the 256 MPI rank on 4 compute node case you obtain as well as at what point is your parallel efficiency the greatest (i.e., speedup / MPI ranks for each case and which case yields the highest value).

Now, plot the **parallel I/O** execution time (Y-axis) to write the whole 32Kx32K universe as a function of the total number of MPI ranks used in the 128 compute node series tests described in the next subsection. Discuss what write I/O performance differences you observe as the number of MPI ranks increases.

2.2 Parallel I/O and Heatmap of Final Universe State

For the parallel I/O and heatmap of final universe state conduct the following series of experiments.

• Run a 32768x32768 cell universe with a threshold of 0% for 128 ticks using the best MPI/thread performance 128 compute node configuration.

- Run a 32768x32768 cell universe with a threshold of 50% for 128 ticks using the best MPI/thread performance 128 compute node configuration.
- Run a 32768x32768 cell universe with a threshold of 75% for 128 ticks using the best MPI/thread performance 128 compute node configuration.

Last, using the previously run 25% threshold experiment configurations, construct a *heatmap* of 1024x1024 size for each configuration (4, 1Kx1K heatmaps in total) where a single element of the heatmap is the sum of a 32x32 grid of the live/dead state within the cell universe. You are allowed to use any post process tools available or modify your C-code to construct the heatmap for you using the MPI gather routines and parallel processing techniques you've learned in class. Make sure you describe your approach for heatmap construction in your report.

Additionally, include each 1Kx1K heatmap graphic where higher numbers mean more cells alive within each 32x32 grid. What do you observe about the patterns in each heatmap as the randomization threshold is increased?

For details on how to create a heatmap using GNUPlot, see: http://www.kleerekoper.co.uk/2014/05/how-to-create-heatmap-in-gnuplot.html.

3 HAND-IN INSTRUCTIONS

Submit both your PDF report and C-code to the class Submitty server, submitty.cs.rpi.edu.