Parallel Computing/Programming Assignment #3: Point-2-Point Reduction vs. Collective Reduction

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February 22, 2022

DUE DATE: Thursday, March 3rd, 2022, 11:59 a.m.

1 Assignment Description

For this first MPI assignment, your are to develop a point-2-point message version of the MPI_reduce operation and compare the performance of that version you develop to the collective version, MPI_reduce across a variable number of MPI rank configurations.

1.1 MPI_P2P_reduce

Here, you will create a function called MPI_P2P_Reduce that takes the same arguments as MPI_reduce except that your implementation will only perform the MPI_SUM operation and the final reduction result goes to MPI rank 0. The core algorithm forms a binary tree of the MPI ranks. The algorithm becomes:

```
MPI_P2P_reduce(...)
{
    1. Each rank computes sum over local data array.
    2a. Compute pairwise sums using MPI_Isend/Irecv
    between MPI ranks at a stride of 1:
        0 and 1, 2 and 3, 4 and 5, ....n-2 and n-1
        with result going to lower rank id
    2b. Compute pairwise sums using MPI_Isend/Irecv
        between MPI ranks at a stride of 2:
        0 and 2, 4 and 6, ....n-4 and n-2.
        with result going to lower rank id
    2c. Compute pairwise sums using MPI_Isend/Irecv
```

```
between MPI ranks at a stride of 4:
        0 and 4, 8 and 12, ...n-8 and n-4.
    with result going to lower rank id
2d, e, ... keep going until Rank 0 has the final sum result.
}
```

As a specific example, consider a 16 MPI rank configuration. There will be $log_2(16)$ steps after the calculation of the local sum at each MPI rank. Note, for below the pair (x,y) means that MPI rank x performs a MPI_Irecv for sum data that was sent via MPI_Isend from MPI rank y. Upon receipt, MPI rank x will add rank y's value to it running sum.

```
    Rank pair lists: (0,1), (2,3), (4,5), (6,7), (8,9), (10,11), (12,13), (14,15)
    Rank pair lists: (0,2), (4,6), (8,10), (12,14)
    Rank pair lists: (0, 4), (8, 12)
    Rank pair lists: (0, 8)
    Final sum: 0
```

Use the AiMOS's clock_now function from Assignment 2 to measure the number of cycles this point-2-point reduce operation took as well as the MPI collection MPI_reduce. Exclude data allocation and initialization in your timing measurements. To use this function make sure you include the following header file on AiMOS and use per the example below.

Note, that the clock_frequency is 512,000,000 cycles per second. The outputs for the collective and point to point versions should be the sum answer from Rank 0 and the time in seconds (one space between numbers). The first output is for the point2point version (on a separate line and the second is the collective reduce version.

1.2 Data for Reducing

. The data will consist of a single long array of 1 billion (e.g., $2^{30} = 1,073,741,824$) entries of type MPI_LONG_LONG. This array will be split equally across ranks and it's data value will be initialized to it's global position index. For example, bigarray[0] = 0 while $bigarray[9999999998elements_per_rank] = 0$

999999999. This initialization is to be deterministic across all MPI rank configurations. In terms of the answer it is easy to compute via the equation N * N - 1/2 which is 576,460,751,766,552,576.

The output of your program (from MPI Rank 0 only!), must print the above correct answer in order to receive full credit for this assignment for BOTH the MPI_Reduce and MPI_P2P_Reduce implementations.

Note, that each AiMOS compute node has 512 GB of RAM. A 32 rank, 1 node run will than have 32 arrays of 32K entries each. So, you should have plenty of RAM for all the experiments.

1.3 Experiments

You will conduct a strong scaling experiment using "AiMOS", the CCI IBM DCS/AC922 supercomputer system. Here, you will have 32 MPI rank per AiMOS compute-node unless otherwise specified.

- Run 1 billion entry 2 MPI ranks (1 AiMOS nodes)
- Run 1 billion entry 4 MPI ranks (1 AiMOS nodes)
- Run 1 billion entry 8 MPI ranks (1 AiMOS nodes)
- Run 1 billion entry 16 MPI ranks (1 AiMOS nodes)
- Run 1 billion entry 32 MPI ranks (1 AiMOS nodes)
- Run 1 billion entry 64 MPI ranks (2 AiMOS nodes)
- Run 1 billion entry 128 MPI ranks (4 AiMOS nodes)
- Run 1 billion entry 256 MPI ranks (8 AiMOS nodes)

In your report, you will plot the execution time of each reduction approach as a function of the number of MPI ranks and suggest a reason why the shape of the performance curve for both implementations and why one is faster than the other.

To launch the above experiments, you will need to use the sbatch SLURM job scheduling command. Class lecture will review how to use this command as well it is documented on the CCI Wiki. For using Spectrum-MPI, make sure use the slurmSpectrum.sh example script to launch your MPI jobs from the CCI Wiki (see: https://secure.cci.rpi.edu/wiki/Slurm/.

2 HAND-IN INSTRUCTIONS

Both the report in PDF format and MPI/C-code and headerfiles are to be submitted using the Class Grading System, submitty.cs.rpi.edu.