

# Lab #9: MPI P2P Quiz + Collectives

## PCSE 2015

### Announcements

1. Would you be interested in building a compute cluster from scratch with cutting edge parts, installing the OS and a bunch of programs, measuring the performance given certain restrictions and objectives, and then competing internationally to see if you've got what it takes to take home the SC15 Student Cluster Competition Title for the fourth year in a row? More information can be found here (URL embedded). Informational meeting next week. Contact Cyrus via email if you would be interested.
2. We'll be using X11 tunneling today. Before you start PuTTY, go to the Windows Start Menu, search for Xming, and start Xming first. Next, follow the directions given here(URL embedded) for configuring PuTTY to enable X11 Forwarding.
3. We'll also be (optionally) using Allinea DDT today. Follow the directions here(URL embedded) to start DDT from a login node. Make sure to compile programs with the "-g" if you would like to profile/debug them with DDT.

### Part 1: Bug Finder!

1. I have looked through the quizzes from Tuesday and found the most common bugs. They are now coded and provided in the code/part\_1 folder. There are 7 buggy codes – ring\_1a, ring\_1b, ring\_1c, ring\_2a, ring\_2b, ring\_3a, and ring\_3b for C and Fortran.
2. Your job is to figure out what's wrong with each one. Some will run but provide incorrect answers. Some will never complete.
3. Try your hand at debugging each code with the help of DDT and/or print statements.
4. In your lab writeup, clearly identify the mistake(s) in each version. Describe the behavior of the buggy code and provide the fix for each one.

### Part 2: Collectives #1

Create a new file called `collect_1.*` where `*` is `c` or `F90` and write a code following the instructions given below:

1. Given the following variable block (C):

```
#define NPROCS 4
#define ASIZE 8
#define ASIZEPP (ASIZE / NPROCS)
#define ROOT 2
double master_vec[ASIZE], local_vec[ASIZEPP];
int i, j, rank, numprocs;
double root_sum, local_sum, global_sum;
```

Or, given the following variable block (Fortran):

```
integer, parameter :: NPROCS = 4
integer, parameter :: ASIZE = 8
integer, parameter :: ASIZEPP = ASIZE / NPROCS
integer, parameter :: ROOT = 2
double precision master_vec(ASIZE), local_vec(ASIZEPP)
integer i, j, rank, numprocs, ierr
double precision root_sum, local_sum, global_sum
```

2. If on the root process, initialize master\_vec to 100, else initialize to -1.
3. Use MPI\_Scatter to scatter the contents of master\_vec from the root process to all other processes and store in the local\_vec variable.
4. On all processes, add rank to each value of local\_vec.
5. Use the provided code block to output the value of local\_vec (C):

```
for(i=0; i<numprocs; i++){
    if(rank == i){
        printf("*** (Rank_%i):_local_vec_Output:\n",rank);
        for(j = 0; j < ASIZEPP; j++){
            printf("(Rank_%i):_%i%f\n",rank,j,local_vec[j]);
        }
    }
    MPI_Barrier(MPI_COMM_WORLD);
}
```

(Fortran):

```
do i=0, numprocs-1
    if(rank == i)then
        write(*,'(a,i0,a)') "*** (Rank_",rank,"):_local_vec_Output:"
        do j = 1, ASIZEPP
            write(*,"(2(a,i0),f10.2)") "(Rank_",rank,"):_",j,local_vec(j)
        end do
    end if
    call MPI_BARRIER(MPI_COMM_WORLD,ierr)
end do
```

to verify that each process has successfully received it's piece of master\_vec.

6. For each process, sum each element of local\_vec and store the answer in local\_sum.
7. Use MPI\_Reduce to sum each process's copy of local\_sum onto the root process in the global\_sum variable.
8. Now, use MPI\_Gather to gather the modified pieces of local\_vec back into master\_vec on the same root process that you did the scatter from.
9. For the root process, sum the contents of master\_vec and store the answer in root\_sum.

10. For each process, print the elements of master\_vec and compare global\_sum to root\_sum (they should match).
11. Use 4 MPI processes. Your output should look like (C):

```

TACC: Starting up job 5057379
TACC: Setting up parallel environment for MVAPICH2+mpispawn.
TACC: Starting parallel tasks...
***(Rank 0): local_vec Output:
(Rank 0): 0 100.000000
(Rank 0): 1 100.000000
***(Rank 1): local_vec Output:
(Rank 1): 0 101.000000
(Rank 1): 1 101.000000
***(Rank 2): local_vec Output:
(Rank 2): 0 102.000000
(Rank 2): 1 102.000000
***(Rank 3): local_vec Output:
(Rank 3): 0 103.000000
(Rank 3): 1 103.000000
***(Rank 0): master_vec Output:
(Rank 0): 0 -1.000000
(Rank 0): 1 -1.000000
(Rank 0): 2 -1.000000
(Rank 0): 3 -1.000000
(Rank 0): 4 -1.000000
(Rank 0): 5 -1.000000
(Rank 0): 6 -1.000000
(Rank 0): 7 -1.000000
(Rank 0): The ROOT process is: 2
(Rank 0): Root Sum: 0.000000
(Rank 0): MPI_Reduce Sum: 0.000000
***(Rank 1): master_vec Output:
(Rank 1): 0 -1.000000
(Rank 1): 1 -1.000000
(Rank 1): 2 -1.000000
(Rank 1): 3 -1.000000
(Rank 1): 4 -1.000000
(Rank 1): 5 -1.000000
(Rank 1): 6 -1.000000
(Rank 1): 7 -1.000000
(Rank 1): The ROOT process is: 2
(Rank 1): Root Sum: 0.000000
(Rank 1): MPI_Reduce Sum: 0.000000
***(Rank 2): master_vec Output:
(Rank 2): 0 100.000000
(Rank 2): 1 100.000000
(Rank 2): 2 101.000000
(Rank 2): 3 101.000000
(Rank 2): 4 102.000000
(Rank 2): 5 102.000000
(Rank 2): 6 103.000000
(Rank 2): 7 103.000000
(Rank 2): The ROOT process is: 2
(Rank 2): Root Sum: 812.000000
(Rank 2): MPI_Reduce Sum: 812.000000
***(Rank 3): master_vec Output:
(Rank 3): 0 -1.000000
(Rank 3): 1 -1.000000
(Rank 3): 2 -1.000000
(Rank 3): 3 -1.000000
(Rank 3): 4 -1.000000

```

```

(Rank 3): 5 -1.000000
(Rank 3): 6 -1.000000
(Rank 3): 7 -1.000000
(Rank 3): The ROOT process is: 2
(Rank 3): Root Sum: 0.000000
(Rank 3): MPI_Reduce Sum: 0.000000

```

TACC: Shutdown complete. Exiting.

### (Fortran):

```

TACC: Starting up job 5057379
TACC: Setting up parallel environment for MVAPICH2+mpispawn.
TACC: Starting parallel tasks...
*** (Rank 0): local_vec Output:
(Rank 0): 1 100.00
(Rank 0): 2 100.00
*** (Rank 1): local_vec Output:
(Rank 1): 1 101.00
(Rank 1): 2 101.00
*** (Rank 2): local_vec Output:
(Rank 2): 1 102.00
(Rank 2): 2 102.00
*** (Rank 3): local_vec Output:
(Rank 3): 1 103.00
(Rank 3): 2 103.00
*** (Rank 0): master_vec Output:
(Rank 0): 1 -1.00
(Rank 0): 2 -1.00
(Rank 0): 3 -1.00
(Rank 0): 4 -1.00
(Rank 0): 5 -1.00
(Rank 0): 6 -1.00
(Rank 0): 7 -1.00
(Rank 0): 8 -1.00
(Rank 0): The ROOT process is: 2
(Rank 0): Root Sum: 0.00
(Rank 0): MPI_Reduce Sum: 0.00
*** (Rank 1): master_vec Output:
(Rank 1): 1 -1.00
(Rank 1): 2 -1.00
(Rank 1): 3 -1.00
(Rank 1): 4 -1.00
(Rank 1): 5 -1.00
(Rank 1): 6 -1.00
(Rank 1): 7 -1.00
(Rank 1): 8 -1.00
(Rank 1): The ROOT process is: 2
(Rank 1): Root Sum: 0.00
(Rank 1): MPI_Reduce Sum: 0.00
*** (Rank 2): master_vec Output:
(Rank 2): 1 100.00
(Rank 2): 2 100.00
(Rank 2): 3 101.00
(Rank 2): 4 101.00
(Rank 2): 5 102.00
(Rank 2): 6 102.00
(Rank 2): 7 103.00
(Rank 2): 8 103.00
(Rank 2): The ROOT process is: 2
(Rank 2): Root Sum: 812.00
(Rank 2): MPI_Reduce Sum: 812.00

```

```

*** (Rank 3): master_vec Output:
(Rank 3): 1      -1.00
(Rank 3): 2      -1.00
(Rank 3): 3      -1.00
(Rank 3): 4      -1.00
(Rank 3): 5      -1.00
(Rank 3): 6      -1.00
(Rank 3): 7      -1.00
(Rank 3): 8      -1.00
(Rank 3): The ROOT process is: 2
(Rank 3): Root      Sum:      0.00
(Rank 3): MPI_Reduce Sum:      0.00

TACC: Shutdown complete. Exiting.

```

12. Try modifying NPROCS to 16, ASIZE to 64, and ROOT to 13. Do you still get the output you expect?

### Part 3: Collectives #2

Create a new file called `collect_2.*` where `*` is `c` or `F90` and write a code following the instructions given below:

1. Given the following MODIFIED variable block (C):

```

#define NPROCS 4
#define ASIZE 8
#define ASIZEPP (ASIZE / NPROCS)
#define ROOT 2
double master_vec[ASIZE], local_vec[ASIZEPP];
int i, j, rank, numprocs;
int sendcount[NPROCS], recvcnt[NPROCS], displacement[NPROCS];
double root_sum, local_sum, global_sum;

```

Or, given the following MODIFIED variable block (Fortran):

```

integer, parameter :: NPROCS = 4
integer, parameter :: ASIZE = 8
integer, parameter :: ASIZEPP = ASIZE / NPROCS
integer, parameter :: ROOT = 2
double precision master_vec(ASIZE), local_vec(ASIZEPP)
integer i, j, rank, numprocs, ierr
integer sendcount(NPROCS), recvcnt(NPROCS), displacement(NPROCS)
double precision root_sum, local_sum, global_sum

```

2. This code should be a very similar to the code you created in Collectives #1. The difference should be in the routine calls outlined below.
3. Change `MPI_Scatter` to `MPI_Scatterv`.
4. Change `MPI_Gather` to `MPI_Gatherv`.
5. Change `MPI_Reduce` to `MPI_Allreduce`.
6. You'll need to properly set `sendcount`, `recvcnt`, and `displacement` to get your `scatterv` and `gatherv` calls to act like the scatter and gather calls of Collectives #1.
7. Highlight the differences in your output as a result of your `MPI_Allreduce` call.