Assignment MPI: Process Symmetry and Collectives

The purpose of this assignment is for you to learn more about

- getting started with MPI by implementing a hello world
- implementing a static workload partitioning scheme in MPI on numerical integration using a reduction collective

As usual all time measurements are to be performed on the cluster.

To be able to compile and run an MPI program on mamba, you need to add the line module load mpich/3.2.1 at the end of the file .bashrc located in the home directory of your account on mamba. (log off and back in afterward.)

To compile an MPI application, use the mpic compiler in C and the mpicxx compiler in C++. They also serve as linker. To run an MPI application using 19 processes, you can run mpirun -n 19 ./myprogram. But you will need to have a proper node allocation first. And if you have a proper node allocation then specifying -n is not necessary because the cluster scheduler does that for you.

To queue an MPI job on mamba, you will need to specify how many nodes and how many cores per node you plan on using. You could use qsub -l nodes=2:ppn=16 to request two nodes with 16 cores each, or qsub -l procs=32 if you only care about having 32 cores independently of where they are. You can also use qsub -l nodes=4:ppn=4, to have 4 cores from 4 different nodes. There is currently a cap on mamba that prevents you from requiring more than 32 processes in total. This is abstracted in the scaffolding.

1 Hello World

This problem is fairly simple. It consists in initializing the MPI application and getting each process to print a message of the form "I am process i out of P. I am running on machine.".

Question: Go into the hello_world/ directory and write the code in hello.cpp

Question: Run the code on mamba using make run_1x16, make run_2x8, make run_4x8 and confirm that the run happen on different machines by looking at the output generated in the run.sh.oxyz files.

2 Numerical Integration: static

Adapt the numerical integration application to make it use MPI in a simple way. The first MPI process should take the first N/P iterations of the loop, the second should take the next N/P iterations of the loop, etc.. The partial integration should be accumulated on rank 0 so that it can print the correct answer to stdout and the time it took to stderr.

Question: Go into the num_int/ directory. Write the code in mpi_num_int.cpp.

Question: Run and time that program on mamba using many configurations. Use the make bench to queue all the jobs. (Note that if you run make bench twice, all your previously queued jobs will be deleted.)

Question: Generate figures of the results using make plot. Do you observe speedup higher than can be achieved on a single machine?