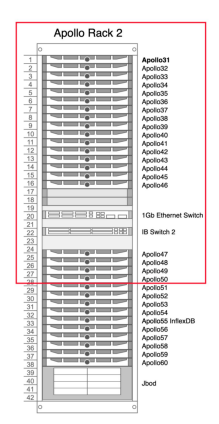
# Platform Introduction & MPI

Parallel Programming 2024

2024/09/12

Lab1

### Platform Introduction - Apollo



- 20 nodes for this course (apollo[31-50])
  - 1 head node: apollo31
  - 19 compute nodes: apollo[32-50]
- Intel X5670 2x6 cores @ 2.93GHz
  - HyperThreading is enabled only on the head node
- 96GB RAM on each node
- QDR Infiniband (40 Gb/s)
- OS: Arch Linux (kernel 5.15.90)
- Storage
  - /home & /opt : 4x RAID10 HDD
    - For storing files & codes
  - /share : BeeGFS parallel file system
    - For performance benchmarking



#### Available resources

- 1 login node (apollo31) (200%CPU max)
- 19 compute nodes (1200% CPU max)
  - Use squeue to view Slurm queues
- Cluster monitor: <a href="http://apollo.cs.nthu.edu.tw/monitor">http://apollo.cs.nthu.edu.tw/monitor</a>
  - Monitor CPU & memory usage of each node
- 48GB disk space per user
  - Use quota -s to view your disk quota

Login to the Apollo Cluster

# Login to Apollo

- Address: apollo.cs.nthu.edu.tw
- Username: check email
- Password: check email
- MINING IS PROHIBITED. Also, do not attack the server.

#### SSH - Linux and Mac

- Open terminal
- ssh pp24sXX@apollo.cs.nthu.edu.tw
- Enter password
- You'll be asked to change your password on the first login

#### SSH - Windows

- Tools
  - MobaXterm
  - o <u>Putty</u>
  - Cmd or Powershell (Windows 10)
  - Windows Terminal (Windows 11)
- ssh pp24sXX@apollo.cs.nthu.edu.tw
- Enter password
- You'll be asked to change your password on the first login

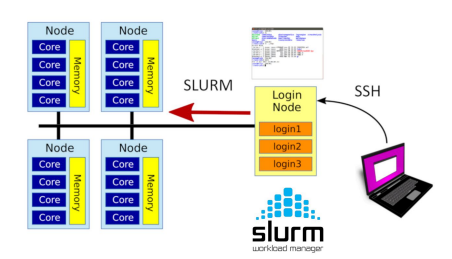
#### Some useful command

- Login: ssh pp24sXX@apollo.cs.nthu.edu.tw
- File transfer:
   rsync -avhP filename pp24sXX@apollo.cs.nthu.edu.tw:filename
- Editors: vim emacs nano
- Slurm job queue: squeue
- Disk quota: quota -s
- Change password: passwd
- Download file: wget aria2c
- Code syntax highlighting: pygmentize

Using the Apollo Cluster

On a cluster system, there are multiple users and multiple nodes. Slurm schedules jobs submitted by users across different nodes, so that the same resource is not used by two jobs at the same time (to ensure accuracy of performance-critical experiments), and also increases the utilization of the cluster.

- Note: Slurm prefers the following jobs:
  - short jobs (you can set time limit)
  - less resource demanding jobs
  - o jobs queued for a long time
  - users that haven't run a lot of jobs recently



- Show partition information
  - o sinfo
    - Displays the information about Slurm nodes and partitions, providing an overview of the cluster's status.
    - We have only one partition, which is the CPU partition
      - idle: The nodes are not allocated
      - alloc: The nodes are fully allocated
      - mix: The nodes are partially allocated
      - down / drain: The nodes are out of service (we will fix them ASAP!)

```
[pp24s000@apollo31 ~]$ sinfo
PARTITION
            AVAIL TIMELĪMIT
                                      STATE NODELIST
                              NODES
apollo-cpu*
                                        mix apollo[35-36]
                        5:00
               up
apollo-cpu*
                                     alloc apollo[32-34]
                        5:00
                                   3
               up
apollo-cpu*
                                       idle apollo[37-50]
               up
                        5:00
                                  14
```

- Interactive Job submission
  - srun [options] ./executable [args]
    - Options:
    - -N NODES: The the minimum number of nodes to run the job
    - -n PROCESSES: The number of total process to launch
    - -c CPUS: The number of cpus available to each process (i.e. threads per process)
    - -t TIME: The time limit in "minutes" or "minutes:seconds"
    - -J NAME: The name of the job. Will be displayed on squeue.
  - srun -N 2 -n 8 -c 2 ./hello\_world
    - Runs a total of 8 processes, each using 2 CPUs, on at least 2 nodes
      - 6 processes \* 2 cpus on 1st node (CPU Allocation: 12/12)
      - 2 processes \* 2 cpus on 2nd node (CPU Allocation: 4/12)

- Batch Job submission
  - sbatch script.sh

```
#!/bin/bash
#SBATCH -J BatchRun
#SBATCH -N 2
#SBATCH -n 4
#SBATCH -c 6
srun ./hello_world
```

- Tracking your jobs
  - sacct

[pp24s000@apollo31 ~]\$ sacct					
JobID	JobName Partition	Account	AllocCPUS	State	ExitCode
5117585.0	hw1	pp24	24	COMPLETED	0:0
5117586	32.txt apollo-cpu	pp24	24	COMPLETED	0:0
5117586.0	hw1	pp24	24	COMPLETED	0:0
5117587	33.txt apollo-cpu	pp24	12	RUNNING	0:0
5117587.0	hw1	pp24	12	RUNNING	0:0
5117588	34.txt apollo-cpu	pp24	12	RUNNING	0:0
5117588.0	hw1	pp24	12	RUNNING	0:0
5117589	35.txt apollo-cpu	pp24	12	PENDING	0:0

- Show the job queue
  - squeue

```
[pp24s000@apollo31 ~]$ squeue
            JOBID PARTITION
                                NAME
                                         USER ST
                                                       TIME
                                                             NODES NODELIST(REASON)
                                                                 1 (QOSMaxJobsPerUserLimit)
          5117629 apollo-cp
                              35.txt pp24s000 PD
                                                       0:00
          5117636 apollo-cp
                              07.txt
                                           vi PD
                                                       0:00
                                                                 1 (QOSMaxJobsPerUserLimit)
          5117634 apollo-cp
                              05.txt
                                          уi
                                              R
                                                       0:01
                                                                 1 apollo33
          5117635 apollo-cp
                              06.txt
                                                                 1 apollo33
                                                       0:01
          5117628 apollo-cp
                              34.txt pp24s000
                                                                 1 apollo32
                                                       0:13
          5117627 apollo-cp
                                                                 1 apollo34
                              33.txt pp24s000
                                                       0:15
```

- Cancel a job
  - scancel [Job ID]

### Environment Modules (module)

- A tool to simplify shell initialization and dynamically manage environment settings using "modulefiles".
- Dynamically update environment variables like PATH, LD\_LIBRARY\_PATH.
- Simplify the process of switching between different software versions.
- Ideal for managing complex software dependencies.
- Used on almost ALL supercomputing clusters

#### Common usages

- List Available Modules
  - module avail
- Load a Module
  - o module load <module\_name>
  - E.g., module load mpi/latest
- Unload a Module
  - o module unload <module\_name>
  - E.g., module unload mpi/latest
- List Loaded Modules
  - module list

- Show Module Info
  - o module show <module name>
  - E.g., module show openmpi
- Swap Modules
  - o module swap <module1> <module2>
  - E.g., module swap mpi/latest openmpi
- Unload All Modules
  - module purge

MPI Hello World

#### MPI Hello World (Send, Recv Test)

```
#include <string.h>
int main(int argc, char *argv[])
   int i, rank, size, namelen;
   char name[MPI MAX PROCESSOR NAME];
   MPI Status stat;
   MPI_Init(&argc, &argv);
   MPI_Comm_size(MPI_COMM_WORLD, &size);
   MPI Comm rank(MPI COMM WORLD, &rank);
   MPI Get processor name(name, &namelen);
   if (rank == 0) {
       printf("Hello world: rank %d of %d running on %s\n", rank, size, name);
           MPI_Recv(&rank, 1, MPI_INT, i, 1, MPI_COMM_WORLD, &stat);
          MPI_Recv(&size, 1, MPI_INT, i, 1, MPI_COMM_WORLD, &stat);
          MPI_Recv(&namelen, 1, MPI_INT, i, 1, MPI_COMM_WORLD, &stat);
          MPI_Recv(name, namelen + 1, MPI_CHAR, i, 1, MPI_COMM_WORLD, &stat);
          printf("Hello world: rank %d of %d running on %s\n", rank, size, name);
   } else {
       MPI_Send(&rank, 1, MPI_INT, 0, 1, MPI_COMM_WORLD);
       MPI_Send(&size, 1, MPI_INT, 0, 1, MPI_COMM_WORLD);
       MPI Send(&namelen, 1, MPI INT, 0, 1, MPI COMM WORLD);
       MPI_Send(name, namelen + 1, MPI_CHAR, 0, 1, MPI_COMM_WORLD);
  MPI Finalize();
```

Get this code on Apollo:
cp /opt/intel/mpi/test/test.c .

#### MPI\_Send

#### MPI\_Recv

### Provided Compilers & MPIs

- Compilers
  - GCC 12.2.1 (Default)
    - gcc / g++
  - Intel Compiler Classic
    - module load compiler
    - icc / icpc
  - Intel oneAPI Compiler
    - module load compiler
    - icx / icpx

- MPI Implementations
  - Intel MPI
    - module load mpi
  - OpenMPI 4.1.6 + UCX
    - module load openmpi

# Using different combinations of MPI & compilers

- Intel MPI + GCC
  - module load mpi
  - o mpicxx ...
- Intel MPI + ICC
  - module load mpi compiler
  - I\_MPI\_CXX=icpc mpicxx ...
- OpenMPI + GCC
  - module load openmpi
  - o mpicxx ...
- OpenMPI + ICC
  - module load openmpi compiler
  - OMPI\_CXX=icc mpicxx ...

#### Practice

- Compile and run the hello world program.
  - o mpicxx test.c -o test
  - ./test
- Compare different srun options

Measuring time in your code

#### Correct measurement method

- srun -n4 time ./hello
- sbatch & time srun
- MPI Wtime()
- omp\_get\_wtime()
- clock\_gettime(CLOCK\_MONOTONIC, ...)
- std::chrono::steady\_clock

```
#!/bin/bash
#SBATCH -n 4
#SBATCH -N 2
time srun ./hello
```

#### Example: MPI\_Wtime()

```
double starttime, endtime;
starttime = MPI_Wtime();
.... stuff to be timed ...
endtime = MPI_Wtime();
printf("That took %f seconds\n",endtime-starttime);
```

#### Example: clock\_gettime(CLOCK\_MONOTONIC, ...)

```
int main() {
   struct timespec start, end, temp;
  double time used;
   clock gettime(CLOCK MONOTONIC, &start);
   .... stuff to be timed ...
   clock gettime(CLOCK MONOTONIC, &end);
   if ((end.tv nsec - start.tv nsec) < 0) {</pre>
       temp.tv sec = end.tv sec-start.tv sec-1;
       temp.tv nsec = 1000000000 + end.tv nsec - start.tv nsec;
   } else {
       temp.tv sec = end.tv sec - start.tv sec;
       temp.tv nsec = end.tv nsec - start.tv nsec;
   time used = temp.tv sec + (double) temp.tv nsec / 1000000000.0;
   printf("%f second\n", time used);
```

### Wrong measurement method

- time srun -n4 ./hello: this time include queuing time
- time(NULL): the resolution is too low (1-second)
- clock(): it will count 2x time when using two threads and will not include I/O time.
- clock\_gettime(CLOCK\_REALTIME, ...): it will be affected by NTP adjustments and DST changes.
- std::high\_resolution\_clock::now(): it may be affected by NTP adjustments and DST changes.

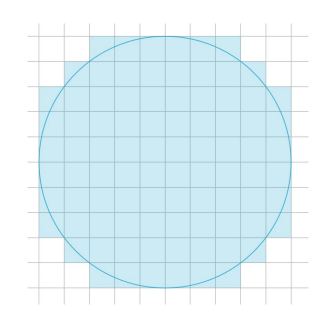
Lab 1 - Pixels in circle

#### Pixels in circle

Suppose we want to draw a filled circle of radius r on a 2D monitor, how many pixels will be filled?

We fill a pixel when any part of the circle overlaps with the pixel. We also assume that the circle center is at the boundary of 4 pixels.

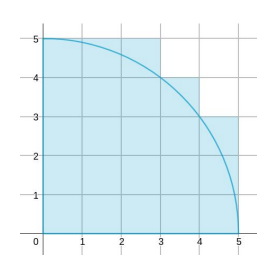
For example, 88 pixels are filled when r=5.



#### Pixels in circle

#### Equation:

$$ext{pixels}(r) = 4 imes \sum_{x=0}^{r-1} \left\lceil \sqrt{r^2 - x^2} 
ight
ceil$$



Example: r = 5

$$\begin{aligned} pixels(5) &= 4 \bigg( \bigg\lceil \sqrt{25 - 0} \bigg\rceil + \bigg\lceil \sqrt{25 - 1} \bigg\rceil + \bigg\lceil \sqrt{25 - 4} \bigg\rceil + \bigg\lceil \sqrt{25 - 9} \bigg\rceil + \bigg\lceil \sqrt{25 - 16} \bigg\rceil \bigg) \\ &= 4(5 + 5 + 5 + 4 + 3) \\ &= 88 \end{aligned}$$

### Lab Spec

- Parallelize the calculation using MPI.
- Program input format: srun -Nnode -nproc ./lab1 r k
  - node: number of nodes
  - proc: number of MPI processes
  - o r: the radius of circle, integer
  - k: integer
- Program output: pixels % k (Since the output pixels may be very large, we output the remainder instead.)
- Your program should be at least (n/2) times faster than the sequential version when running with n processes. For example, when running with 12 processes, your execution time should not exceed 1/6 of the sequential code.

#### Lab Spec

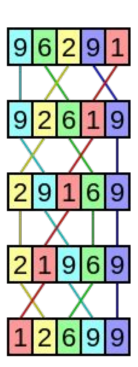
- The sequential code lab1.cc and a build file Makefile can be found at /home/pp24/share/lab1/sample, copy these files to your home directory.
- All of the test cases can be found in /home/pp24/share/lab1/testcases
- Within the same directory of lab1.cc and Makefile, run lab1-judge to check.
- Scoreboard
- Submit your code to eeclass:
  - lab1.cc
  - Makefile (optional, if you change any compile flags)
  - module.list (optional)
    - By default, will use GCC & Intel MPI to judge your code
    - If you are using other modules (such as openmpi), run module save ./module.list to generate it
- Due: 9/19 (Thu.) 23:59 (1 week)
- Full score for AC of all 12 test cases; otherwise, zero.

```
[pp24s000@apollo31 lab1]$ lab1-judge
Looking for lab1.cc: OK
Looking for Makefile: OK
Running: /usr/bin/make -C /share/judge dir
make: Entering directory '/share/judge_dir
mpicxx -std=c++17 -03 -fopenmp
make: Leaving directory '/share/judge_dir/
01.txt
          0.31
                 accepted
03.txt
          0.42
                 accepted
02.txt
          0.36
                 accepted
                 accepted
04.txt
          0.36
05.txt
          0.36
                 accepted
06.txt
          0.47
                 accepted
07.txt
          0.52
                 accepted
08.txt
          0.46
                 accepted
09.txt
          0.52
                 accepted
10.txt
          1.07
                 accepted
11.txt
          0.82
                 accepted
12.txt
          1.47
                 accepted
Removing temporary directory /share/judge_
Scoreboard: not updating {12 6.74} -x-> {1
```

#### HW1: Odd-Even Sort

• Spec

• Due: 11/3 23:59



# Next Lab on 9/26 (2 weeks later)

- Lab2 & HW2 release
  - HW2: Same deadline on 11/3, on another CPU platform (maybe)
- Profiling tools
  - Learn how to collect metrics from your application, to help identify bottlenecks and optimize the performance.
    - It's important to give ideas on how you interpret the profiling results, and how you improve your code based on the findings in your homework report.