

Lab1

Platform introduction & MPI

Parallel Programming
2022/09/29

Outline

- Platform introduction - Apollo
- Login to Apollo
- Linux command
- MPI hello world
- Compile and job submission
- Time measurement
- Lab1 - Pixels in circle

Platform instruction - Apollo

- 19 nodes for this course (apollo31 - 48, 50)
- Intel X5670 2x6 cores @ 2.93GHz (Hyper threading disabled)
- 96GB RAM (each node)
- 5.5TB shared RAID5 disk
- QDR Infiniband (40 Gb/s)

Software

- OS: Arch Linux
- Compilers: GCC 10.2.0, Clang 11.0.1
- MPI: Intel MPI Library, Version 2019 Update 8
- Scheduler: Slurm 20.02.5

Available resources

- 1 login node (apollo31) (200%CPU max)
- 18 compute nodes (1200% CPU max)
- Use `squeue` to view SLURM usage
- Cluster monitor: <http://apollo.cs.nthu.edu.tw/monitor>
- 48GB disk space per user
- Use `quota -s` to view disk quota

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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 pp22sXX@apollo.cs.nthu.edu.tw`
- Enter password
- You'll be ask to change your password on first login

SSH - Windows

- Tools
 - [MobaXterm](#)
 - [Putty](#)
 - Cmd or Powershell (Windows 10)
 - Windows Terminal (Windows 11)
- `ssh pp22sXX@apollo.cs.nthu.edu.tw`
- Enter password
- You'll be ask to change your password on first login

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Some useful command

- Login: `ssh pp22sXX@apollo.cs.nthu.edu.tw`
- File transfer:
`rsync -avhP filename pp22sXX@apollo.cs.nthu.edu.tw:filename`
- Editors: `vim` `emacs` `nano`
- SLURM usage: `squeue`
- Disk quota: `quota -s`
- Change password: `passwd`
- Download file: `wget` `aria2c`
- Code syntax highlighting: `pygmentize`

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MPI hello world

```
#include <mpi.h>
#include <stdio.h>

int main(int argc, char *argv[]) {
    int rank, size;
    MPI_Init(&argc, &argv);

    MPI_Comm_rank(MPI_COMM_WORLD, &rank); // the total number of process
    MPI_Comm_size(MPI_COMM_WORLD, &size); // the rank (id) of the calling process

    printf("Hello, World.  I am %d of %d\n", rank, size);

    MPI_Finalize();
    return 0;
}
```

You can download this code directly on apollo.

```
wget https://www.open-mpi.org/papers/workshop-2006/hello.c
```

MPI_Send

```
int MPI_Send(const void *buf,  
             int count,  
             MPI_Datatype datatype,  
             int dest,  
             int tag,  
             MPI_Comm comm)
```

MPI_Recv

```
int MPI_Recv(void *buf,  
             int count,  
             MPI_Datatype datatype,  
             int source,  
             int tag,  
             MPI_Comm comm,  
             MPI_Status *status)
```

MPI_Reduce

```
int MPI_Reduce(const void *sendbuf,  
               void *recvbuf,  
               int count,  
               MPI_Datatype datatype,  
               MPI_Op op,  
               int root,  
               MPI_Comm comm)
```


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Compilation

- `gcc` - C compiler
- `g++` - C++ compiler
- `mpicc` - MPI C compiler wrapper
- `mpicxx` - MPI C++ compiler wrapper
- `mpicc` and `mpicxx` actually invoke `gcc` and `g++`
- You can use `clang` instead by adding the flags `-cc=clang` for c and `-cxx=clang++` for c++
- Compile the hello world program:
`mpicc -O3 hello.c -o hello`

Run the hello world program

```
$ srun -n4 ./hello
```

Output:

Hello, World. I am 3 of 4

Hello, World. I am 1 of 4

Hello, World. I am 2 of 4

Hello, World. I am 0 of 4

Job submission

SLURM workload scheduler:

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.

SLURM prefer the following jobs:

- short jobs (you can set time limit)
- less resource demanding jobs
- jobs queued for a long time
- users that haven't run a lot of jobs recently

Job submission using srun

- `srun [options] ./executable [args]`
- Options:
 - `-N NODES`: NODES is the **number of nodes** to run the job
 - `-n PROCESSES`: PROCESSES is the number of **total process** to launch
 - `-c CPUS`: CPUS is the number of cpus available to **each process**
 - `-t TIME`: The time limit in "minutes" or "minutes:seconds"
 - `-J NAME`: The name of the job. Will be displayed on queue

Job submission using sbatch

- Using sbatch command to submit jobs in the background
- You can write a simple script to do that

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

- `$ sbatch script.sh`

Job control

- `sinfo`: view status of nodes
- `squeue`: view submitted jobs in queue
- `scancel JOBID`: cancel a job with its JOBID

Practices

Compile and run the hello world program.

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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) {
        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.

Outline

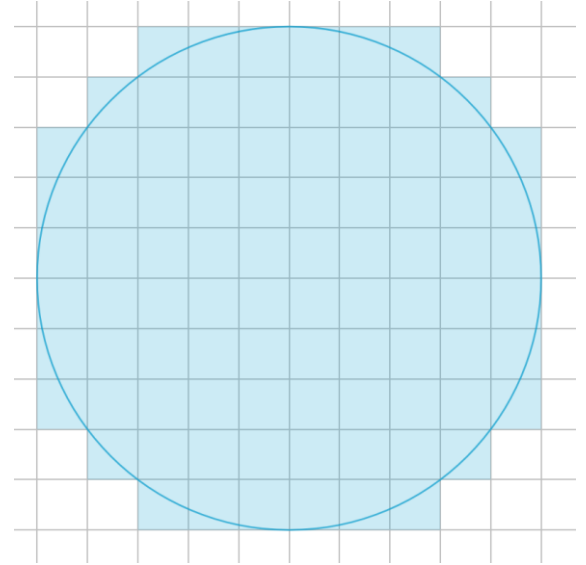
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Pixels in circle

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 $r=5$.



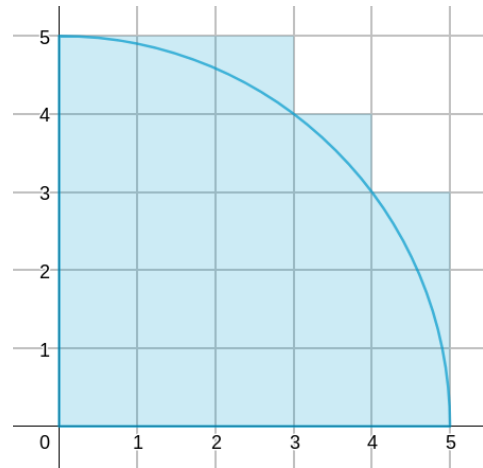
Pixels in circle

Equation:

$$\text{pixels}(r) = 4 \times \sum_{x=0}^{r-1} \left\lceil \sqrt{r^2 - x^2} \right\rceil$$

Example: $r = 5$

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



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
 - 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/pp22/share/lab1/sample`, copy these files to your home directory.
- All of the test cases can be found in `/home/pp22/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)
 - Due: 10/06 (Thu.) 23:59
- Full score for AC of all 12 test cases; otherwise, zero.

```
enmingw32@apollo31 ~/lab1-code> ls
lab1.cc  Makefile*
enmingw32@apollo31 ~/lab1-code> lab1-judge
Looking for lab1.cc: OK
Looking for Makefile: OK
Running: /usr/bin/make -C /home/pp22/enmingw32/.judge.225368503 lab1
make: Entering directory '/home/pp22/enmingw32/.judge.225368503'
mpicxx -std=c++17 -O3 lab1.cc -o lab1
make: Leaving directory '/home/pp22/enmingw32/.judge.225368503'
04.txt 1.02 accepted
03.txt 1.07 accepted
01.txt 0.82 accepted
02.txt 0.77 accepted
05.txt 1.22 accepted
06.txt 1.32 accepted
07.txt 1.27 accepted
08.txt 1.27 accepted
09.txt 1.52 accepted
10.txt 1.97 accepted
11.txt 1.92 accepted
12.txt 2.57 accepted
Removing temporary directory /home/pp22/enmingw32/.judge.225368503
Scoreboard: not updating {12 15.90} -x→ {12 16.72}
```