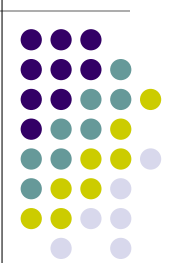
Practical Parallel Computing (実践的並列コンピューティング) 2021 No. 11

Part3: MPI (1) May 20, 2021

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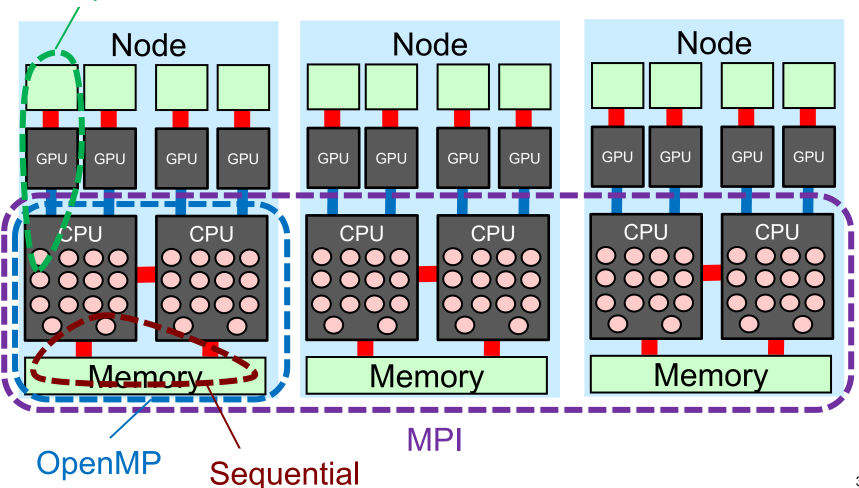
- Part 0: Introduction
 - 2 classes
- Part 1: OpenMP for shared memory programming
 - 4 classes
- Part 2: GPU programming

 - OpenACC (1.5 classes) and CUDA (2.5 classes)
- Part 3: MPI for distributed memory programming

Parallel Programming Methods on TSUBAME

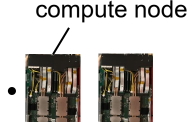


OpenACC/CUDA



How We Can Use Many Nodes

- Submit several jobs into job scheduler
 - cf) Program executions with different parameters ->
 Parameter Sweep
 - Jobs are dependent, and no cooperation















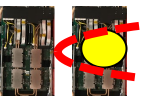






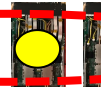
- Use distributed memory programming → A single job can use multiple nodes
 - Socket programming, Hadoop, Spark...
 - And MPI





















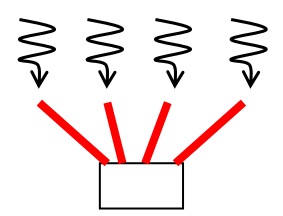
Classification of Parallel Programming Models

Sequential

Shared memory prog. model

Process/
Thread

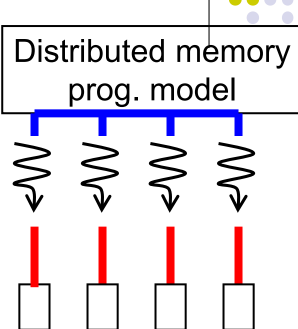
Data



Programming without parallelsim

Threads have access to shared data

- OpenMP
- pthread
- Java thread...



Need communication among processes

- MPI
- socket
- Hadoop, Spark...

MPI (message-passing interface)



- Parallel programming interface based on distributed memory model
- Used by C, C++, Fortran programs
 - Programs call MPI library functions, for message passing etc.
- There are several MPI libraries
 - OpenMPI (default) ← OpenMPI ≠ OpenMP ⊗
 - Intel MPI, SGI MPE, MVAPICH, MPICH...

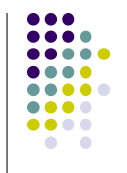
Differences from OpenMP



In MPI,

- An execution consists of multiple processes (not threads)
 - We can use multiple nodes ©
 - The number of running processes is basically constant
- No variables are shared. Instead message passing is used
 - Data distribution has to be programmed
- No smart syntaxes such as "omp for" or "omp task" ☺
 - Task distribution has to be programmed ☺





/gs/hs1/tga-ppcomp/21/hello-mpi

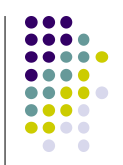
```
[make sure that you are at a interactive node (r7i7nX)]
module load cuda openmpi [Do once after login]
cd ~/t3workspace [In web-only route]
cp -r /gs/hs1/tga-ppcomp/21/hello-mpi .
cd hello-mpi
make
[An executable file "hello" is created]
mpiexec -n 7 --oversubscribe ./hello
```

Number of processes

Needed on the terminal (May 20, 2021)

Name of program (using options are ok)

Compiling and Executing MPI Programs



Case of OpenMPI library on TSUBAME3.0

- - module load cuda openmpi, and then use mpicc
 - For sample programs, "make" command works
- To execute Number of processes
 - mpiexec -n 7 --oversubscribe ./hello

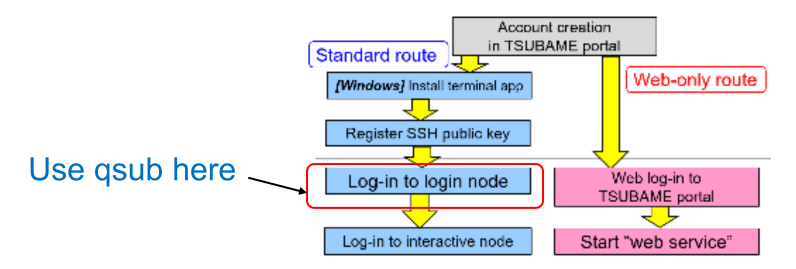
↑ These methods uses 1 (current) node.

To use multi-nodes, we need "job submission"!

Notes on Job Submission in "Standard route"



- On an interactive node via "standard route", qsub/qstat commands may not be found
- Please use qsub/qstat on a login node
 - (out of iqrsh)



Submit an MPI Job (case of OpenMPI)



 We are going to execute it with 4 processes × 2 nodes = 8 processes

(1) Make a script file: job.sh

(2) Submit the job with "qsub"

qsub job.sh

 $\uparrow \leq 0:10:00$ and ≤ 2 nodes

preparation qsub -g tga-ppcomp job.sh

 \uparrow >0:10:00 or \geq 3nodes

Be careful for TSUBAME point

Number of Noncesses pr

Number of processes per node

Program name (and option)

11

Notes in This Lecture (Also see OpenMP(4) slides on Apr 29)



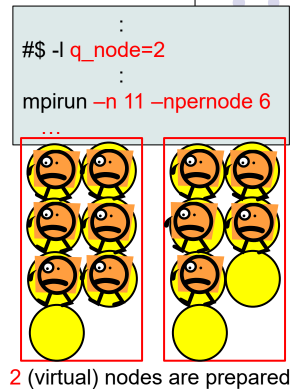
- Usually, avoid consumption of TSUBAME points
- 通常は無料利用の範囲にとどめてください
 - h_rt <= 0:10:00
- If necessary for reports, you can use up to 36,000 points in total per student
- 本講義のレポートの作成に必要な場合、一人あたり合計で 36,000ポイントまで利用を認めます
 - f_node x 1node x 10 hours
- Please check point consumption on TSUBAME portal
- The TSUBAME group name is tga-ppcomp







Nodes, Cores, MPI Processes















- 2 (virtual) nodes are preparedEach node has 4 cores (q_core)
- 4 processes are created per node. Totally 8 are created → 2 nodes are used
- 8 (virtual) nodes are preparedEach node has 1 cores (s_core)
- 1 processes are created per node. Totally 8 are created → 8 nodes are used
- 6 processes are created per node. Totally 11 are created → 2 nodes are used

Each node has 7 cores (q node)

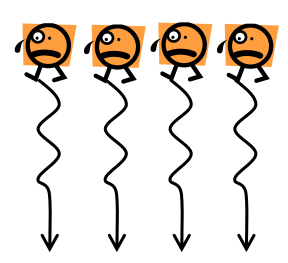
(There are idle cores)

An MPI Program Looks Like



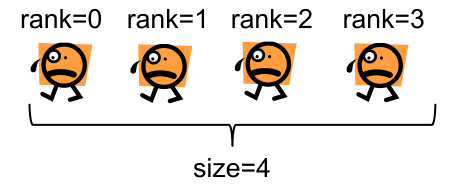
```
#include <stdio.h>
#include <mpi.h>
int main(int argc, char *argv[])
  MPI_Init(&argc, &argv); ← Initialize MPI
  (Computation/communication)
  MPI_Finalize();
                          ← Finalize MPI
```

If number of processes=4



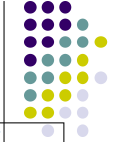
ID of Each MPI Process

- Each process has its ID (0, 1, 2...), called rank
 - MPI_Comm_rank(MPI_COMM_WORLD, &rank);
 - → Get its rank
 - MPI_Comm_size(MPI_COMM_WORLD, &size);
 - → Get the number of total processes
 - 0 ≤ rank < size
 - The rank is used as target of message passing





"mm" sample: Matrix Multiply

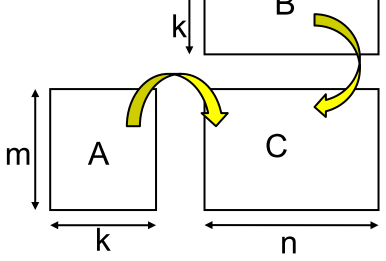


MPI version available at /gs/hs1/tga-ppcomp/21/mm-mpi/

A: a (m × k) matrix, B: a (k × n) matrix C: a (m × n) matrix

 $C \leftarrow A \times B$

- Algorithm with a triple for loop
- Supports variable matrix size.
 - Each matrix is expressed as a 1D array by column-major format



Execution:

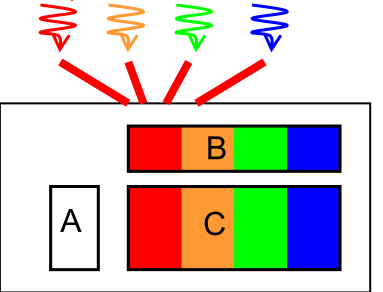
mpiexec -n [np] --oversubscribe ./mm [m] [n] [k] (interactive) mpiexec -n [np] -npernode [nn] ./mm [m] [n] [k] (in job script)

Why Distributed Programming is More Difficult (case of mm-mpi)



Shared memory with OpenMP:

Programmers consider how computations are divided

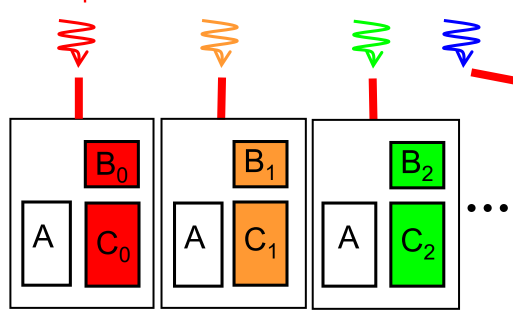


In this case, matrix A is accessed by all threads

→ Programmers do not have to know that

Distributed memory with MPI:

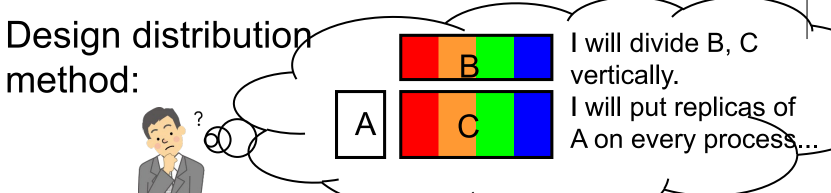
Programmers consider how data and computations are divided



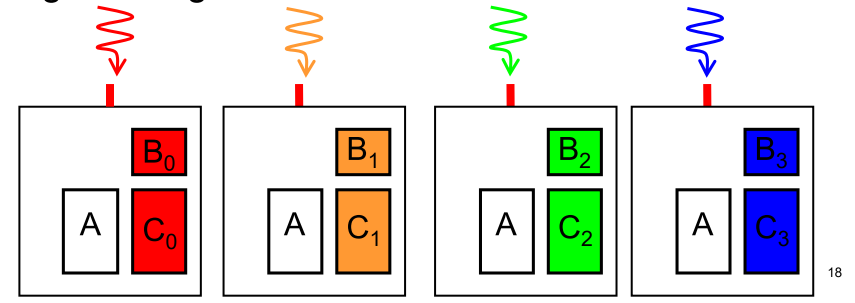
Programmers have to design which data is accessed by each process

Programming Data Distribution

(case of mm-mpi)

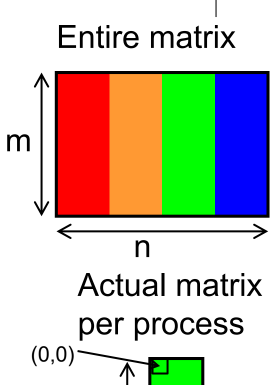


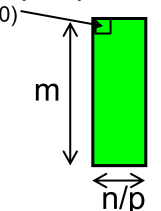
Programming actual location:



Programming Actual Data Distribution

- We want to distribute a m ×n matrix among p processes
 - We assume n is divisible by p
- Each process has a partial matrix of size m × (n/p)
 - We need to "malloc" m*(n/p)*sizeof(data-type) size
 - We need to be aware of relation between partial matrix and entire matrix
 - (i,j) element in partial matrix owned by Process r ⇔
 (i, n/p*r + j) element in entire matrix

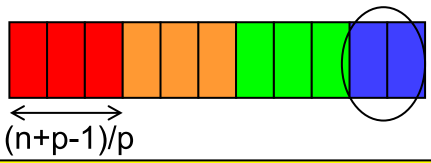




What is Done for Indivisible Cases

- What if data size n is indivisible by p?
- We let n=11, p=4
 - How many data each process take?
 - n/p = 2 is not good (C division uses round down). Instead, we should use round up division
 - \rightarrow (n+p-1)/p = 3 works well

Note that the "final" process takes less than others

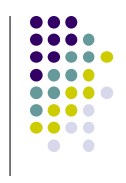


See divide_length() function in mm-mpi/mm.c It calculates the range the process should take (first index s and last index e)

Notes in Time Measurement



- In mm-mpi, gettimeofday() is used for time measurement
- For accurate measurement, we should call MPI_Barrier(MPI_COMM_WORLD) before measurement
 - This synchronizes all processes
 - All processes need to call this

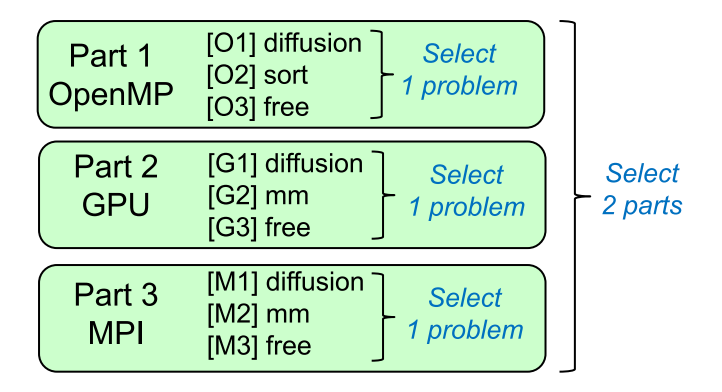


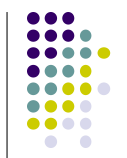
Some slides are deleted (and had bugs). They are moved to May 24 slides

Assignments in this Course



 There is homework for each part. Submissions of reports for 2 parts are required





Assignments in MPI Part (1)

Choose one of [M1]—[M3], and submit a report

Due date: June 10 (Thursday)

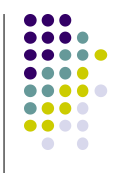
[M1] Parallelize "diffusion" sample program by MPI.

- Do not forget to change Makefile and job.sh appropriately
- Use deadlock-free communication
 - see neicomm_safe() in neicomm-mpi sample

Optional:

- To make array sizes (NX, NY) variable parameters
- To consider the case with NY is indivisible by p
 - see divide_length() in mm_mpi sample
- To improve performance further. Blocking, 2D division, etc



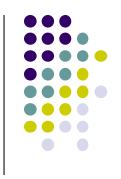


[M2] Improve "mm-mpi" sample in order to reduce memory consumption

Optional:

- To consider indivisible cases
- To try advanced algorithms, such as SUMMA
 - the paper "SUMMA: Scalable Universal Matrix Multiplication Algorithm" by Van de Geijn
 - http://www.netlib.org/lapack/lawnspdf/lawn96.pdf

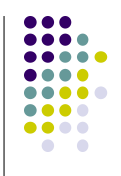




[M3] (Freestyle) Parallelize any program by MPI.

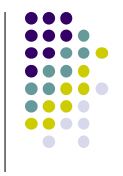
- cf) A problem related to your research
- More challenging one for parallelization is better
 - cf) Partial computations have dependency with each other

Notes in Report Submission (1)



- Submit the followings via T2SCHOLA
 - (1) A report document
 - PDF, MS-Word or text file
 - 2 pages or more
 - in English or Japanese (日本語もok)
 - (2) Source code files of your program
 - Try "zip" to submit multiple files

Notes in Report Submission (2)



The report document should include:

- Which problem you have chosen
- How you parallelized
 - It is even better if you mention efforts for high performance or new functions
- Performance evaluation on TSUBAME
 - With varying number of processes

Updated on May 24

- On a interactive nodes, 1 ≤ [number of processes] ≤ 14
- For >7 processes, use "mpiexec -n [P] --oversubscribe ..."
- To use more CPU cores, you need to do "job submission"
- With varying problem sizes
- Discussion with your findings
- Other machines than TSUBAME are ok, if available

Next Class

- MPI (2)
 - How to parallelize diffusion sample with MPI
 - Related to [M1]