# Practical Parallel Computing (実践的並列コンピューティング)

Part 2: GPU

No 1: Overview and OpenACC May 2, 2024

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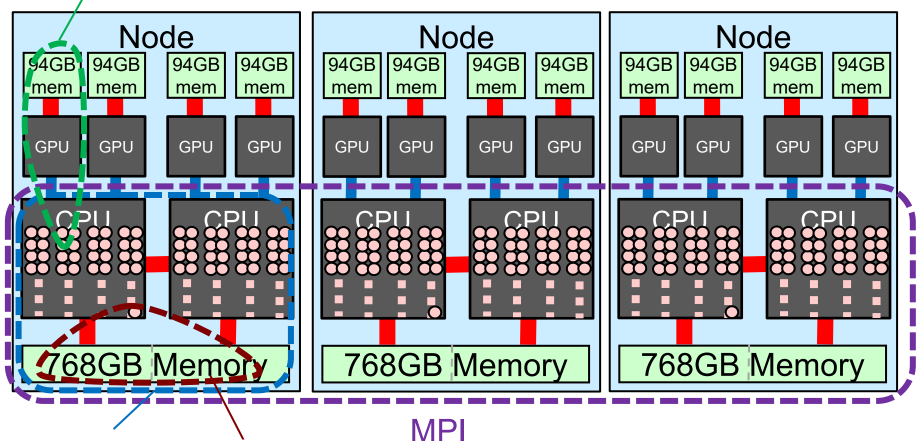
- Part 0: Introduction
  - 2 classes
- Part 1: OpenMP for shared memory programming
  - 4 classes
- Part 2: GPU programming
  - 4 classes
     We are here (1/4)
  - OpenACC (1.5 classes) and CUDA (2.5 classes)
- Part 3: MPI for distributed memory programming
  - 3 classes

# Parallel Programming Methods on TSUBAME



OpenACC/CUDA

**OpenMP** 

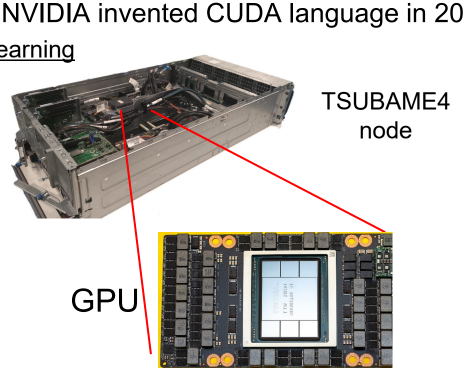


## **GPU Computing**

- Graphic processing units (GPU) have been originally used for computing graphics (including video games)
- A high performance GPU has many cores
  - CPU: 2 to 32 cores. GPU: >1000 cores
  - The concept is called GPGPU (General-Purpose computing on GPU)
- GPGPU became popular since NVIDIA invented CUDA language in 2007

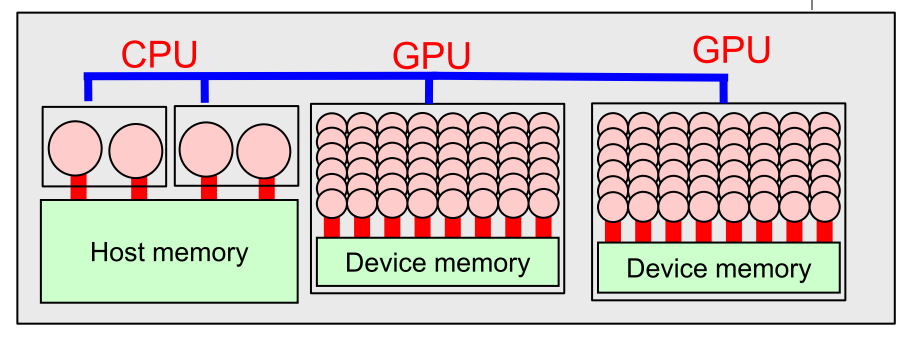
Recently it is popular for deep learning





### A Compute Node with GPU





- A GPU has its distinct memory (device memory)
  - CPU memory is called host memory
- Many cores in a GPU share its device memory
- If there are multiple GPUs, each has its device memory

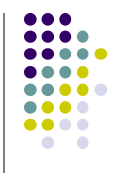


A GPU is a board or a card attached to computers

- → It cannot work alone. Driven by CPUs
- → Different programming methods

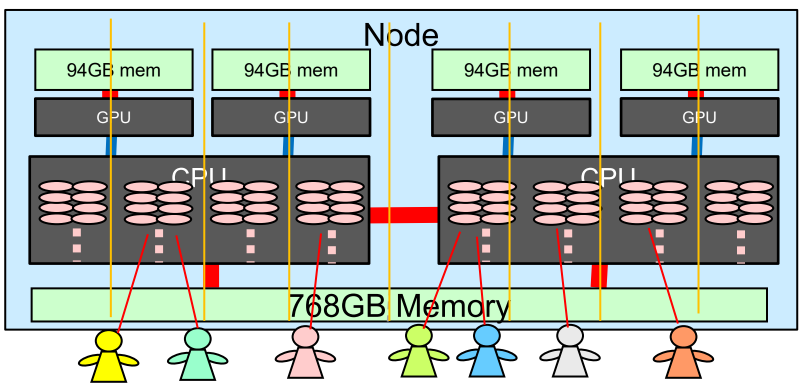
Comparing EPYC 9654 (TSUBAME4's CPU) and Tesla H100-94GB (TSUBAME4's GPU)

	1 CPU	1 GPU
Number of cores	96 cores (192 cores with 2CPUs)	16,896 CUDA cores (=128 x 132SMXs)
Clock Frequency	3.55GHz >	1.98GHz
Peak Computation Speed (double precision)	5.45TFlops	66.9TFlops
Memory Capacity	384GB > (768GB shared by 2CPUs)	94GB



## Notes on TSUBAME Interactive Node

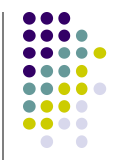




In an interactive node, each user uses 0.5 GPU

7,680 CUDA cores (128x60SMs), ~46GB memory
 If you want use "a full GPU", you may try "qsub" (described later)

## Programming Environments for NVIDIA GPUs



- CUDA ← We will use after OpenACC
  - The most famous environment, designed by NVIDIA
  - C/Fortran + new syntaxes
  - Use "nvcc" command for compile
    - module load cuda
    - nvcc ... XXX.cu
  - For more general programs than OpenACC ©
- - C/Fortran + <u>directives</u> (#pragma acc ...), Easier programming ☺
  - Supported by NVIDIA HPC SDK
    - module load nvhpc
    - nvc -acc ... XXX.c
  - For parallel programs with for-loops
- OpenMP 5, OpenCL...





C/C++/Fortran + directives

```
int a[100], b[100], c[100];
int i;
#pragma acc data copy(a,b,c)
#pragma acc kernels
#pragma acc loop independent
  for (i = 0; i < 100; i++) {
    a[i] = b[i]+c[i];
  }</pre>
```

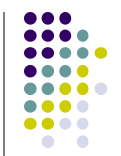
## Examples of OpenACC directives

In this case, each directive has an effect on the following block/sentence

OpenACC is not so popular as OpenMP, unfortunately

- gcc's support is not so good
- On TSUBAME4, we use NVIDIA HPC SDK

# OpenACC Version of "mm" sample



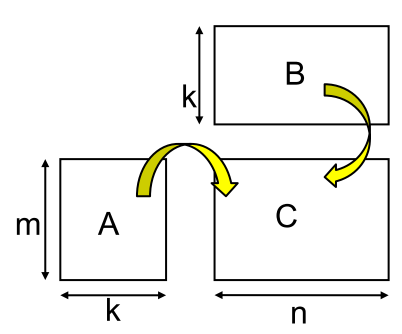
Available at /gs/bs/tga-ppcomp/24/mm-acc/

A: a  $(m \times k)$  matrix, B: a  $(k \times n)$  matrix

C: a (m × n) matrix

 $C \leftarrow A \times B$ 

- Algorithm with a triply-nested forloop
- Supports variable matrix size.
  - Each matrix is expressed as a 1D array by column-major format



Execution: ./mm [m] [n] [k]

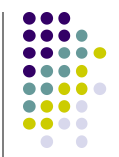


./mm 2000 2000 2000



```
[make sure that you are at a interactive node (rXn11)]
module load nvhpc [Do once after login]
cd ~/ppc24
cp -r /gs/bs/tga-ppcomp/24/mm-acc .
cd mm-acc
make
[You will see some messages, and an executable file
"mm" is created]
```

# Notes on Compiling OpenACC Programs



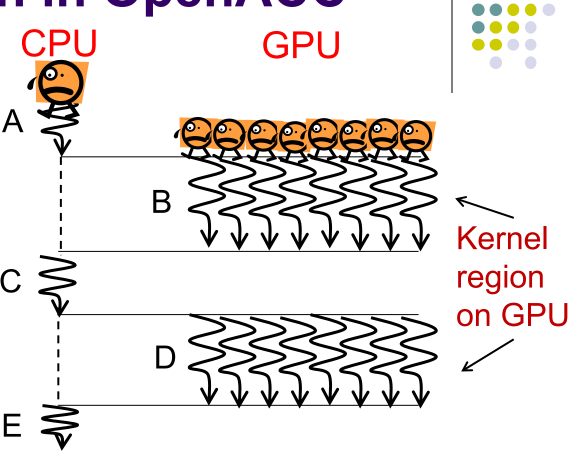
- NVIDIA HPC SDK on TSUBAME4.0
  - module load nvhpc, and then use nvc command
  - Use -acc option in compiling and linking
  - -Minfo=accel option outputs many information on parallelization

```
Example of output
:
26, Generating copyin(A[:m*k])
Generating copy(C[:m*n])
Generating copyin(B[:k*n])
Loop is parallelizable
.
```

They are not errors ©

## Kernel Region in OpenACC

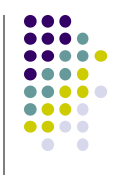
```
int main()
{
    A:
#pragma acc kernels
    {
        B;
    }
    C;
#pragma acc kernels
    D;
    E;
}
```



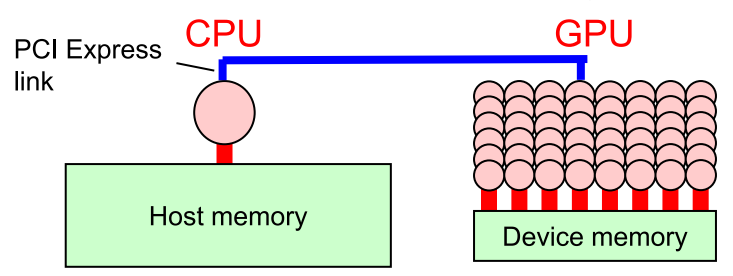
A sentence/block immediately after #pragma acc kernels is called a kernel region, executed on GPU

 We don't need to specify number of threads (it is hard to specify explicitly)

# Data Movement between CPU and GPU



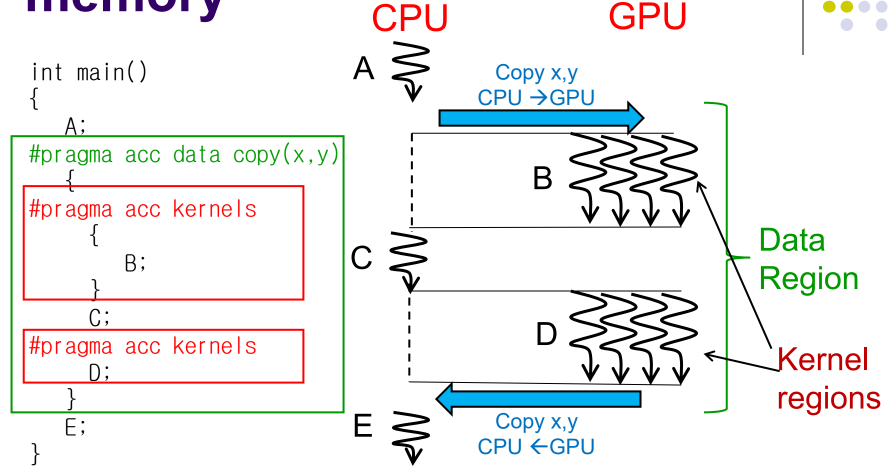
- We need to move data between CPU and GPU
  - Host (CPU) memory and Device (GPU) memory are distinct, like distributed memory
  - Threads on a GPU share the device memory



For this purpose, we use #pragma acc data directive

→ This defines a data region

Data Directives to use GPU memory



- Data region may contain 1 or more kernel regions
- Data movement occurs at beginning and end of data region

### **Data Directive (1)**

- Arrays (like a):
  - we can write array names if the sizes are statically declared → entire array is copied
- Pointers as arrays (like b):

```
cf) b [ 0 : 20 ]
start index number of elements
```

- Partial copying like b[10:5] or a[4:4] work
- Scalar variables (like x):
  - You can omit copy(x) → The compiler detects automatically ☺

```
int x;
float a[10];
double *b = (double*)
    malloc(20*sizeof(double));
:
#pragma acc data copy(x, a, b[0:20])
:
Same meaning
```

#pragma acc data copy(a[0:10], b[0:20])

## **Data Directive (2)**



- Directions of copying
  - ... data copyin(...): Copy <u>CPU→GPU</u> at the begining
  - ... data copyout(...): Copy GPU→CPU at the end
  - ... data copy(...): Do both

Optimization of data movement will help speedup

## **Loop Directive**

```
int a[100], b[100], c[100];
int i;
#pragma acc data copy(a,b,c)
#pragma acc kernels
#pragma acc loop independent
    for (i = 0; i < 100; i++) {
        a[i] = b[i]+c[i];
    }</pre>
```

- #pragma acc loop must be included in "acc kernels" or "acc parallel"
- Directly followed by "for" loop
  - The loop must have a loop counter, as in OpenMP
  - List/tree traversal is NG
- … loop independent: Iterations are done in parallel by multiple GPU threads
- … loop seq: Done sequentially. Not be parallelized
- … loop: Compiler decides

## OpenACC Version of mm (mm-acc/mm.c)

```
#pragma acc data copyin(A[0:m*k],B[0:k*n]),copy(C[0:m*n])
#pragma acc kernels
#pragma acc loop independent
  for (j = 0; j < n; j++) {
    #pragma acc loop seq
    for (l = 0; l < k; l++) {
    #pragma acc loop independent
    for (i = 0; i < m; i++) {
        Ci,j += Ai,l * BI,j;
    } }
</pre>

#pragma acc data copyin(A[0:m*k],B[0:k*n]),copy(C[0:m*n])

We can omit GPU→CPU copy of A,B
    ←For each column in C
    ←For dot product
    ←For each row in C
    ←For each row in C
```

- Each element in C can be computed in parallel (i-loop, j-loop)
- Computation of a single C element is sequential (I-loop)

## Submitting an OpenACC Job to TSUBAME4 Job Scheduler



- Sequential version
  - see mm directory

- OpenACC version
  - see mm-acc directory

#!/bin/sh

- With gpu\_1 type, you can use a full GPU
- (node\_h or node\_f types for multi-GPU)

mm/job.sh

#!/bin/sh #\$ -cwd #\$ -l cpu\_4=1 #\$ -l h\_rt=00:10:00 ./mm 2000 2000 2000

resource type
and count

maximum
run time

<u>mm-acc/job.sh</u>

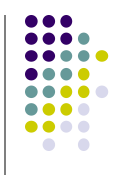
#\$ -cwd #\$ -l gpu\_1=1 #\$ -l h\_rt=00:10:00

./mm 2000 2000 2000

- Job submission
  - qsub job.sh



## Resource Types (from ppcomp-sup slide)



	Physical	Memory	
Resource type	_	(GB)	GPUs
node_f	192	768	4
node_h	96	384	2
node_q	48	192	1
node_o	24	96	0.5
gpu_1	8	96	1
gpu_h	4	48	0.5
cpu_160	160	368	0
cpu_80	80	184	0
cpu_40	40	92	0
cpu_16	16	36.8	0
cpu_8	8	18.4	0
cpu_4	4	9.2	0

← largest

same size with a partition of interactive node

← smallest

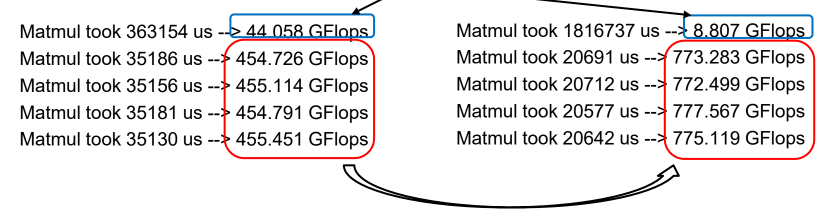
### 0.5 GPU vs 1 GPU



m=n=k=2000

Interactive node (0.5GPU) gpu\_1 (1GPU)

We can ignore speed of the first computation



About 1.7 times speedup

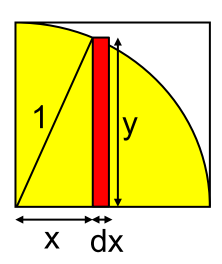
The first computation suffers slowdown by initialization of GPU etc. The similar slowdown also occurs in CPU programs

# OpenACC version of "pi" sample



Estimate approximation of  $\pi$  (circumference/diameter) by approximation of integration

- Available at /gs/bs/tga-ppcomp/24/pi-acc/
- Method
  - Let SUM be approximation of the yellow area
  - $4 \times SUM \rightarrow \pi$
- Execution:./pi [n]
  - n: Number of division
  - Cf) ./pi 100000000
- Compute complexity: O(n)



$$dx = 1/n$$
  
y = sqrt(1-x\*x)

## Algorithm of "pi"



### **OpenMP**

### **OpenACC**

```
double pi(int n) {
  int i;
  double sum = 0.0;
  double dx = 1.0 / (double)n;
#pragma omp parallel
#pragma omp for reduction(+:sum)
  for (i = 0; i < n; i++) {
     double x = (double)i * dx;
     double y = sqrt(1.0 - x*x);
     sum += dx*y;
  return 4.0*sum; }
```

```
double pi(int n) {
  int i;
  double sum = 0.0;
  double dx = 1.0 / (double)n;
#pragma acc kernels
#pragma acc loop independent reduction(+:sum)
  for (i = 0; i < n; i++) {
     double x = (double)i * dx;
     double y = sqrt(1.0 - x*x);
     sum += dx*y;
  return 4.0*sum; }
```

☆ For scalar variables, "data copy" is omitted

### **Notes on Number of Threads**

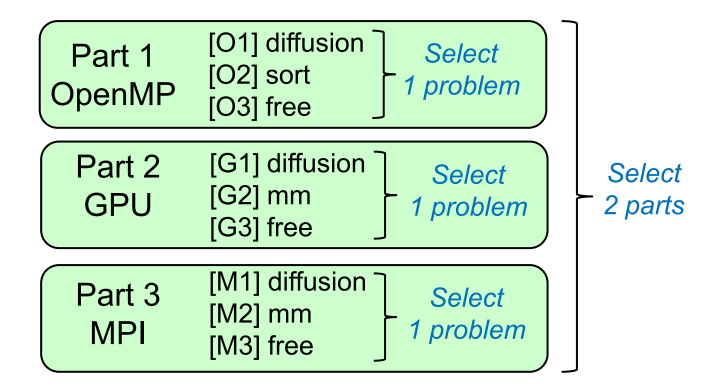


- In OpenMP, the number of threads is set by OMP NUM THREADS
- In OpenACC, the number is automatically determined per loop
- In OpenMP, thread ID is obtained by omp\_get\_thread\_num()
- In OpenACC, we cannot see thread ID

## **Assignments in this Course**



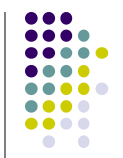
 There is homework for each part. Submissions of reports for 2 parts are required





Choose one of [G1]—[G3], and submit a report

Due date: May 30 (Thursday)



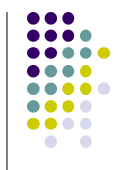
### [G1] Parallelize "diffusion" sample program by OpenACC or CUDA

 You can use Makefile in /gs/bs/tga-ppcomp/24/diffusion-acc/ or /gs/bs/tga-ppcomp/24/diffusion-cuda/

#### Optional:

- To make array sizes variable parameters
- To compare OpenACC vs CUDA
- To improve performance further
  - Different assignment of threads and elements (CUDA), etc 27





[G2] Evaluate speed of "mm-acc" or "mm-cuda" in detail

mm-acc: /gs/bs/tga-ppcomp/24/mm-acc/

mm-cuda: /gs/bs/tga-ppcomp/24/mm-cuda/

- Use various matrices sizes
- Evaluate effects of data transfer cost
- Compare with CPU (OpenMP) version

#### Optional:

- To use different loop orders
- To evaluate both mm-acc and mm-cuda
- To change/improve the program
  - Different assignment of threads and elements (CUDA) etc





[G3] (Freestyle) Parallelize any program by OpenACC or CUDA.

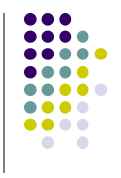
- cf) A problem related to your research
- "sort" sample on GPU?
  - The quick sort may be hard on GPU (There is no "task" syntax)
  - → Bitonic sort?
- More challenging one for parallelization is better
  - cf) Partial computations have dependency with each other

### **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
  - In Part 2, describe which you used, OpenACC or CUDA?
- 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 threads
  - With varying problem sizes
  - Discussion with your findings
  - Other machines than TSUBAME are ok, if available

### **Next Class:**

- GPU Programming (2)
  - OpenACC
    - Improving data copy
    - Improving loop parallelization
  - Introduction of CUDA

#### Schedule

- Mon, May 6: No classes (national holiday)
- Thu, May 9: GPU (2)
- Mon, May 13: GPU (3)
- Thu, May 16: No classes (cancelled/休講)

