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

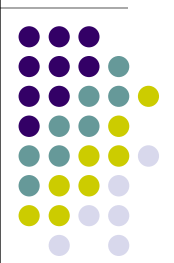
Part 2: GPU

No 3: Threads in CUDA May 11, 2023

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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 (3/4)
 - OpenACC (1.5 classes) and CUDA (2.5 classes)
- Part 3: MPI for distributed memory programming
 - 4 classes

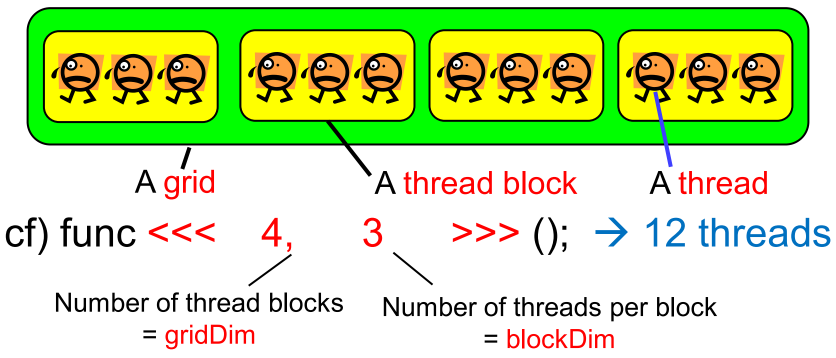
Comparing OpenMP/OpenACC/CUDA



	OpenMP	OpenACC	CUDA
Processors	CPU	CPU+GPU	
File extension	.c	, .CC	.cu
To start parallel (GPU) region	#pragma omp parallel	#pragma acc kernels	func<<<,>>>()
To specify # of threads	export OMP_NUM _THREADS=	(num_gangs, vector_length etc)	
Desirable # of threads	# of CPU cores or less	# of GPU cores or "more"	
To get thread ID	omp_thread_num()	-	blockldx, threadldx
Parallel for loop	#pragma omp for	#pragma acc loop	-
Task parallel	#pragma omp task	-	-
To allocate device memory	-	#pragma acc data	cudaMalloc()
To copy to/from device memory	-	#pragma acc data #pragma acc update	cudaMemcpy()
Functions on GPU	-	#pragma acc routine	global,device

Threads in CUDA

Specify 2 numbers (at least) for number of threads, when calling a GPU kernel function

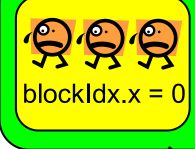


The reason is related to GPU hardware
Thread block ⇔ SMX, Thread ⇔ CUDA core

Thread Block ID, Thread ID



blockldx.x = 2threadIdx.x = 2





blockldx.x = 1



blockldx.x = 2



A grid

A thread block

A thread

For every thread, gridDim.x = 4, blockDim.x = 3

Note: In order to see the entire sequential ID, we should compute blockldx.x * blockDim.x + threadIdx.x

The Case of add-cuda Sample

- /gs/hs1/tga-ppcomp/23/add-cuda/
- We want to do for (i = 0; i < 100; i++) { DA[i] += DB[i]; }

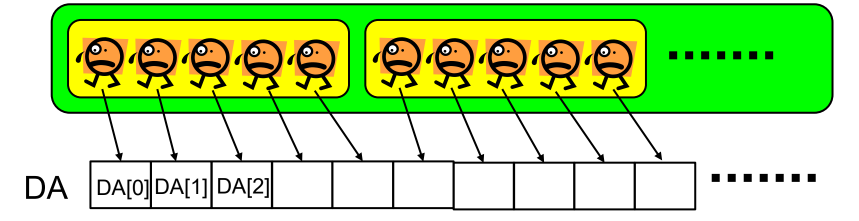
```
[CPU side]

add<<<20, 5>>>(...);

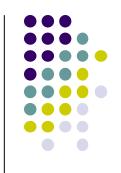
20x5=100 threads
will execute add function

[GPU side]

__global__ void add(int *DA, int *DB)
{
    int i = blockldx.x * blockDim.x + threadldx.x;
    DA[i] += DB[i];
    return;
}
```



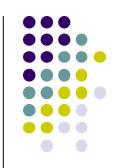
How is Number of Threads Designed? (1)



On CUDA, a different strategy is required from on OpenMP

- On OpenMP, number of threads (OMP_NUM_THREADS) should be ≤ CPU cores (or hyper threads)
 - The number is basically determined by hardware
 - \leq 14 on q_node node, \leq 56 on f_node
- On CUDA, it is better to use number of thread ≥ GPU cores
 - ■ 3584 on TSUBAME3's P100 GPU
 - You can use >1,000,000 threads!

How is Number of Threads Designed? (2)



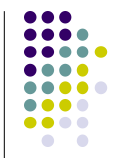
We have to deicide 2 numbers <<
block number, block size>>>

A better way would be

- (1)We decide total number of threads P
- (2)We tune each block size BS
 - Good candidates are 16, 32, 64, ... 1024
- (3) Then block number is P/BS
 - We consider indivisible cases later



"mm" sample: Matrix Multiply (related to [G2])



CUDA versions are at

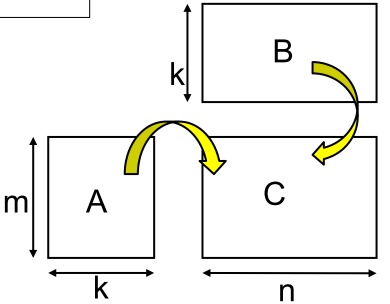
- /gs/hs1/tga-ppcomp/23/mm-v1-cuda/
- /gs/hs1/tga-ppcomp/23/mm-cuda/

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

C: a (m × n) matrix

 $C \leftarrow A \times B$

- Supports variable matrix size
- Execution:./mm [m] [n] [k]



On CUDA, We need to design

- (1) How we parallelize computation
- (2) How we put data on host memory & device memory

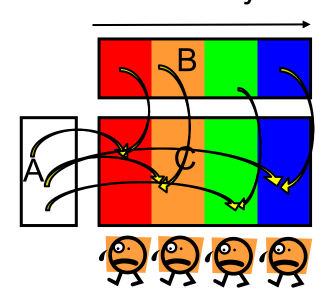


In mm, we can compute different C elements in parallel

•On the other hand, it is harder to parallelize dot-product loop

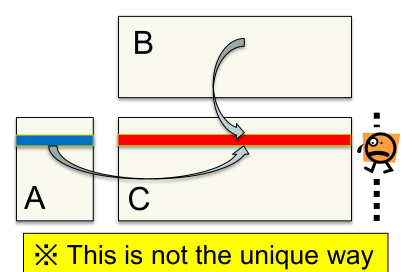
OpenMP

Parallelize column-loop (or row-loop)



CUDA (mm-v1-cuda)

- We can create many threads
- •1 thread computes 1 row
 - We use m threads



Parallelism in mm-v1-cuda

- It is ok to make >1000, >10000 threads on CUDA
- We use <u>m</u> threads for m rows computation

```
matmul_kernel<<<m/m/ss, BS>>>(....);

gridDim blockDim (BS=16 in this sample)
```

1 element for 1 row → No need of "i" loop in this sample

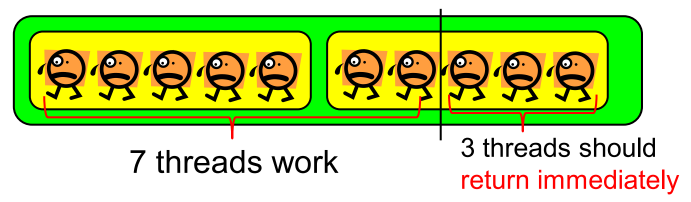
Note: <<<m, 1>>> also works, but speed is not good <<<1, m>>> causes an error if m>1024 (CUDA's rule)

If Number of Threads is Indivisible by BlockDim

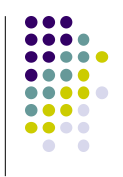


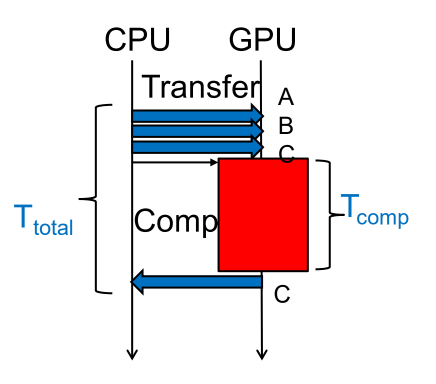
- m: the number of threads
- BS: BlockDim
- If m may be indivisible by BS, we should use <<<(m+BS-1)/BS, BS>>>
- → But # of threads may be larger m. "Extra" threads (id≧m) should not work. See mm-v1-cuda/mm.cu

Example: m=7, BS=5 \rightarrow <<<2,5>>> 10 threads start working, but 3 threads should do nothing



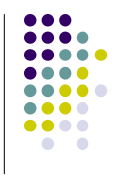
Data Transfer in mm-v1-cuda





- (1) A, B, C are copied from CPU to GPU
 - cudaMemcpy(DA, A, ...) ...
- (2) Computation is done on GPU
- (3) C is copied from GPU to CPU
 - cudaMemcpy(C, DC, ...)

Notes in Time Measurement



- clock(), gettimeofday() must be called from CPU
- For accurate measurement, we should call cudaDeviceSynchronize() before measurement
 - Actually GPU kernel function call and cudaMemcpy(HostToDevice) are non-blocking





- Measured with a P100 GPU on TSUBAME3
 - ./mm 1000 1000 1000 and so on
- The program outputs 2 speeds
 - Speed with data transfer costs → shown on the table
 - Speed without data transfer costs

m=n=k	mm-acc (Gflops)	mm-v1-cuda (Gflops)	
1000	158	15	
2000	193	29	
4000	214	50	Why slow?
6000	198	70	
8000	198	85	

Discussion on Speed (related to [G2])



mm-v1-cuda is slower than mm-acc!

- In mm-acc, i-loop and j-loop has "loop independent"
 - → m×n elements are computed in parallel
- In mm-v1-cuda, we use m threads in total
 - → We should use more threads on a GPU!
 - At least, ≥ 3,584 = number of CUDA cores
 - We see m=8000 threads are still insufficient, and slow

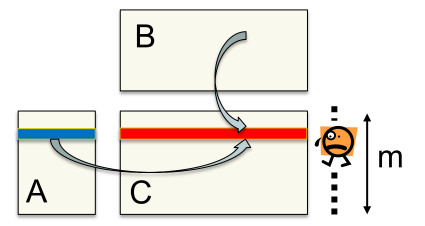
Improvement: How to Use More Threads



In mm, computation of each C element is independent with each other

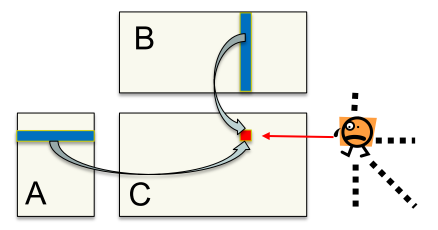
mm-v1-cuda

- •1 thread computes 1 row
 - → We use m threads



mm-cuda

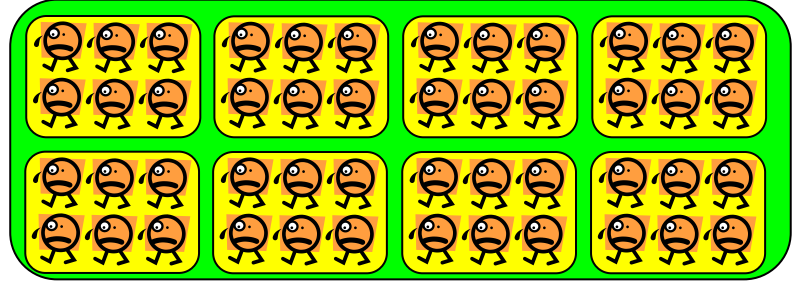
- •1 thread computes 1 element
 - → We use m × n threads !!



Creating Threads with 2D/3D IDs

- Now we want to make m*n (may be >1,000,000) threads
 - <<<(m*n)/BS, BS>>> is ok, but coding is bothersome
- On CUDA, gridDim and blockDim may have "dim3" type,
 3D vector structure with x, y, z fields

cf) func $<<< \dim 3(4,2,1), \dim 3(3,2,1) >>> (); \rightarrow 48 \text{ threads}$

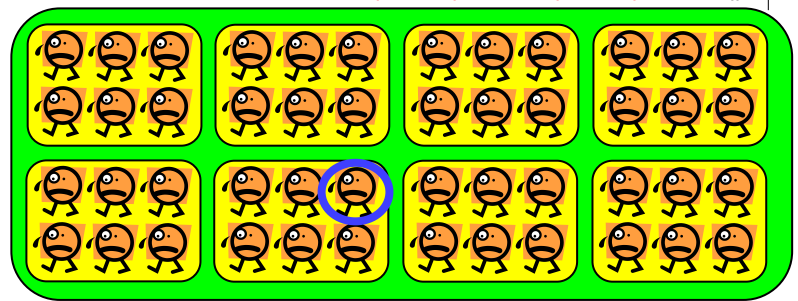


* This example is the case of 2D (Z dimensions are 1)

Thread IDs in multi-dimensional cases



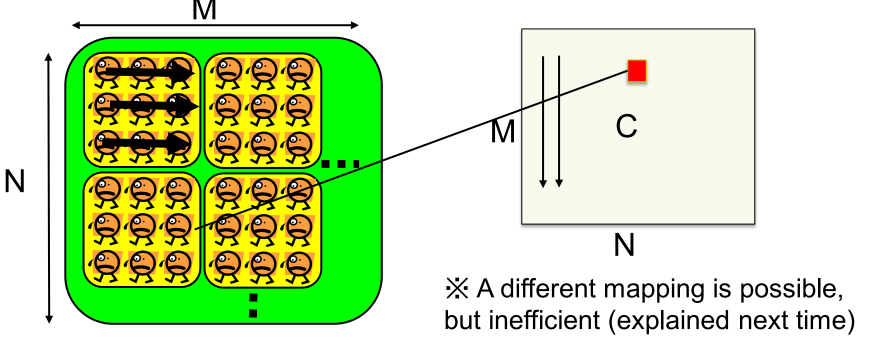
In the case of func <<< dim3(4,2,1), dim3(3,2,1) >>> ();



- For every thread, gridDim.x=4, gridDim.y=2, gridDim.z=1 blockDim.x=3, blockDim.y=2, blockDim.z=1
- For the thread with blue mark,
 blockldx.x=1, blockldx.y=1, blockldx.z=0
 threadldx.x=2, threadldx.y=0, threadldx.z=0



- The total number of threads are m*n
- How do we determine gridDim, blockDim?
 - <<m, n>>> does not work for constraints explained later ⊗
- Here, we determine blockDim as x=16, y=16 → 256 threads per block
 - Then gridDim is computed from M, N
- x is mapped to row index, y is mapped to column index (X)





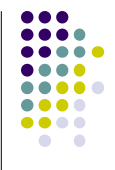


```
gridDim blockDim

matmul_kernel<<<dim3(m / BS, n / BS, 1), dim3(BS, BS, 1)>>>
(DA, DB, DC, m, n, k);
```

BS=16 in this sample Actually, we use rounding up

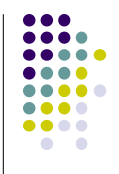
About Programming Efforts



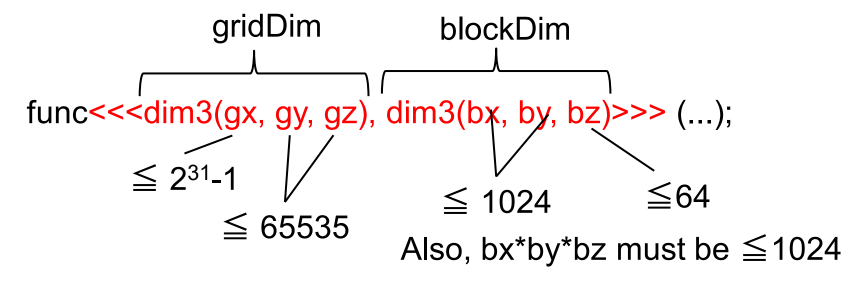
- So far, mm-cuda with (m * n) threads has been explained
 - How fast is it? Please measure it (Related to [G2])
- On the other hand, codes are more complex than mm-acc Programmer have to
 - Call cudaMalloc, cudaMemcpy
 - Determine the number of blocks, threads
 - Determine the structure of for-loop
 - We have to change code largely when parallelization method is changed
 - In OpenACC, adding "#pragma acc loop independent" works

:

CUDA Rules on Number of Threads



```
func<<<A, B>>> (...); (A, B are integers) is same as func<<<dim3(A,1,1), dim3(B,1,1)>>> (...);
```

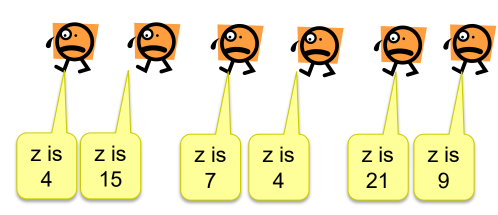


Cf) <<<m, n>>> causes an error if n>1024 ⊗

Rules for Memory/Variables

Variables declared in GPU kernel functions are "thread

private"



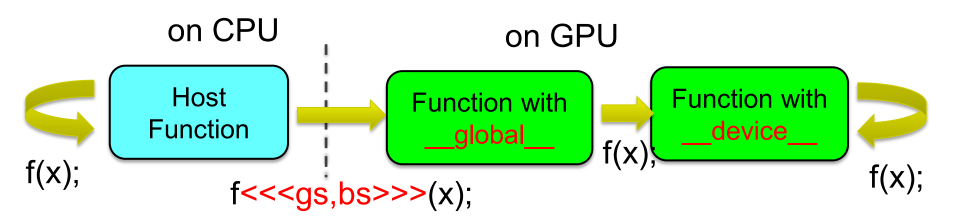
- Device memory is shared by all CUDA threads
 - Be careful to avoid race condition problem (multiple threads write same address)
 - Reading same address is ok
- Do not forget host memory and device memory are separated



Two Types of GPU Kernel Functions

- 1) Functions with **global** keyword
 - "Gateway" from CPU
 - Return value type must be "void"
- 2) Function with <u>device</u> keyword
 - Callable only from GPU
 - Can have return values
 - Recursive call is OK

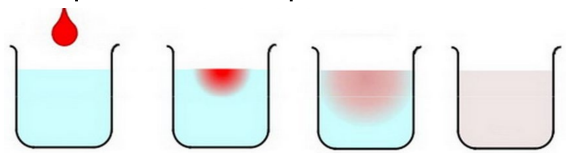
In OpenACC, #pragma acc routine



"diffusion" Sample Program related to [G1]



An example of diffusion phenomena:



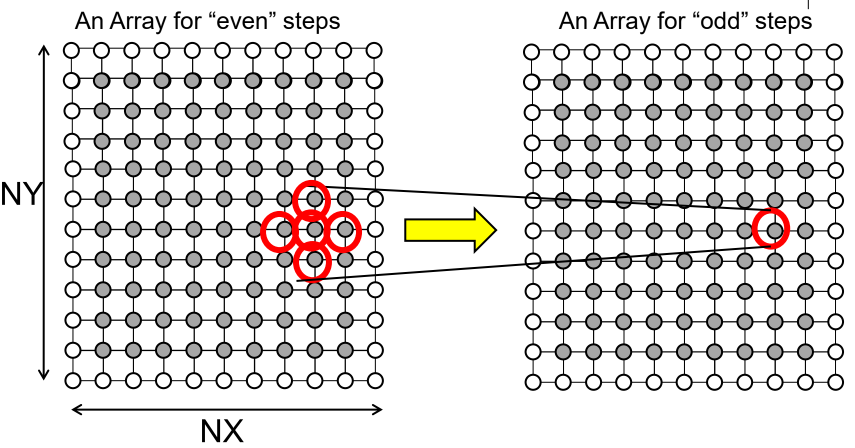
The ink spreads gradually, and finally the density becomes uniform (Figure by Prof. T. Aoki)

Available at /gs/hs1/tga-ppcomp/23/diffusion/ You can use /gs/hs1/tga-ppcomp/23/diffusion-cuda/

- Execution:./diffusion [nt]
 - nt: Number of time steps

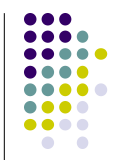






Both arrays have to be on GPU device memory when computations are done

Consideration of Parallelizing Diffusion with CUDA related to [G1]

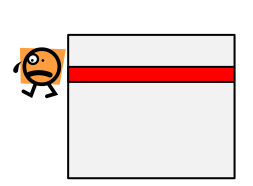


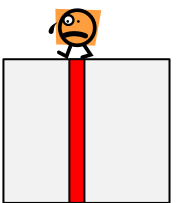
- x, y loops can be parallelized
- t loop cannot be parallelized

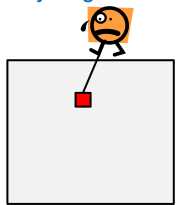
```
[Data transfer from CPU to GPU]
for (t = 0; t < nt; t++) {
                                        GPU computation must be
  for (y = 1; y < NY-1; y++) {
                                        a distinct function
    for (x = 1; x < NX-1; x++) {
                                        (GPU kernel function)
                                        It's better to transfer
                                       data out of t-loop
[Data transfer from GPU to CPU
```

Considering CUDA Threads

- How do we design threads on CUDA?
- There are several choices in [G1]
 - 1thread = 1row
 - We use NY threads in total → only x-loop in kernel function
 - 1thread = 1column
 - We use NX threads in total → only y-loop in kernel function
 - 1thread = 1element
 - We use NX x NY threads in total → No loop in kernel function!
 - This looks fast since the number of threads is very large



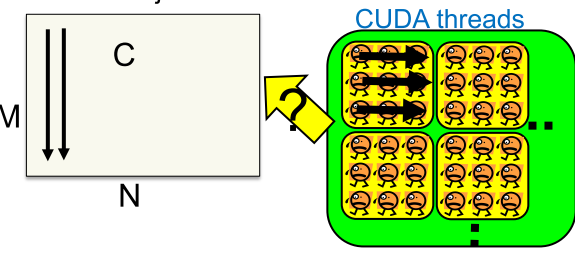




Mapping between Threads and Data

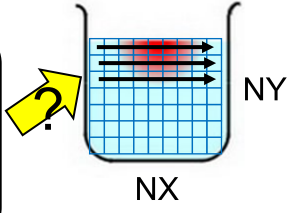
mm-cuda:

Matrices has column-major format



diffusion:

2D array has row-major format

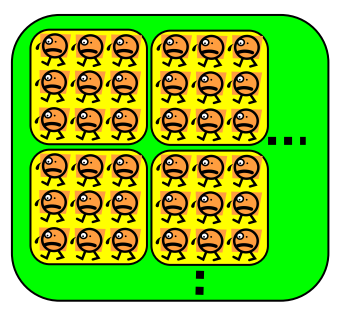


```
j = blockIdx.y * blockDim.y +
threadIdx.y;
i = blockIdx.x * blockDim.x +
threadIdx.x;
: This thread computes Cij
```

```
y = blockIdx.y * blockDim.y +
threadIdx.y;
x = blockIdx.x * blockDim.x +
threadIdx.x;
: This thread computes[y][x]
```

[Q] What if the dimensions are exchanged?

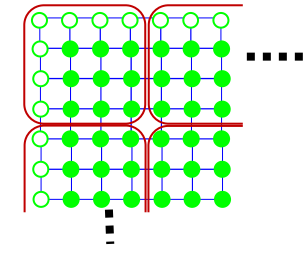




- (1) We decide total number of threads
- → (NX, NY, 1) threads
- See notes on the next page
- (2) We tune each block size (blockDim)
- → Good candidates are (4, 4, 1), (8, 8, 1), (16, 16, 1), (32, 32, 1)
- The number must be ≤ 1024
- How about non-square blocks?
- (3) Then block number (gridDim) is determined
 We should consider indivisible cases
 31

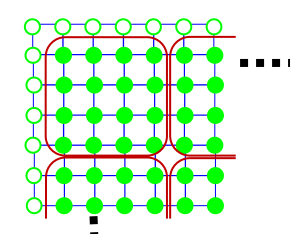
Considering gridDim/blockDim (2)

- In diffusion, Points [1, NX-1) × [1, NY-1), excluded boundary, should be computed There are choices:
 - (A) Create NX x NY threads
 - Thread (x,y) computes (x,y)
 - Threads with below IDs do nothing
 - x == 0 or y == 0 or $x \ge NX-1$ or $y \ge NY-1$
 - (B) Create (NX-2) x (NY-2) threads
 - Thread (x,y) computes (x+1,y+1)
 - Threads with below IDs do nothing
 - $x \ge NX-2$ or $y \ge NY-2$



(A)

(B)



Either is ok ©

Discussion on Data Transfer of Diffusion



Both codes will work, but how about speeds?

} [Data transfer from GPU to CPU]

Computation: O(NX NY nt)
Transfer: O(NX NY)

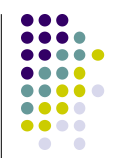
Computation: O(NX NY nt)
Transfer: O(NX NY nt)

What Can be Done in GPU Functions?



- Basic computations (+, -, *, /, %, &&, ||...) are OK
- if, for, while, return are OK
- Device memory access is OK
- Host memory access is NG
- Calling host functions is NG
- Calling most of functions in libc or other libraries for CPUs are NG
 - Several mathematical functions, sin(), sqrt()... are OK
 - printf() is OK
 - Calling malloc()/free() on GPU is OK, if the size must be small
 - Usually, use cudaMalloc() on <u>CPU</u>

Assignments in GPU Part (Abstract)



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

Due date: May 25 (Thursday)

- [G1] Parallelize "diffusion" sample program by OpenACC or CUDA
- [G2] Evaluate speed of "mm-acc" or "mm-cuda" in detail
- [G3] (Freestyle) Parallelize any program by OpenACC or CUDA.





- GPU Programming (4)
 - Discussion on speed and GPU hardware
- Also please note due date of OpenMP assignment is May 11 (Today!)