# Parallel Computations & Applications

National Tsing-Hua University 2019, Summer Semester



#### Outline

- Embarrassingly Computations
- Divide-And-Conquer Computations
- Pipelined Computations
- Synchronous Computations



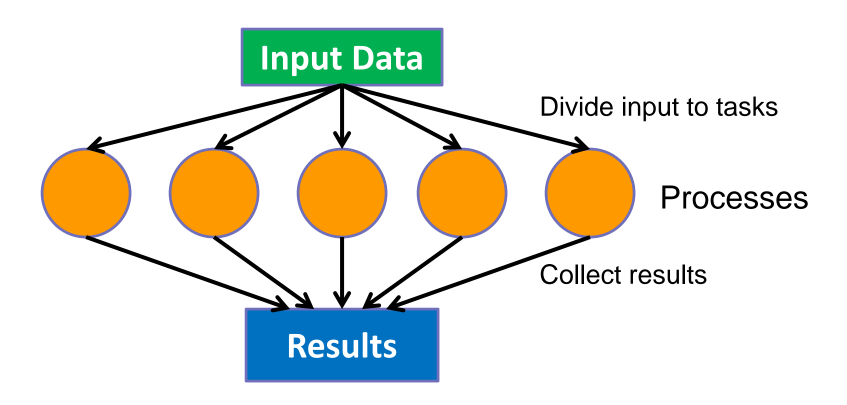
#### Outline

- Embarrassingly Computations
  - Image Transformations
  - Mandelbrot Set
  - Monte Carlo Methods
- Divide-And-Conquer Computations
- Pipelined Computations
- Synchronous Computations



# What is Embarrassingly Parallel

A computation that can be divided into a number of completely independent tasks



# Example 1: Image Transformations

- Low-level image operations:
  - Shifting: object shifted by  $\Delta x$  in the x-dimension and  $\Delta y$  in the y-dimension:

$$x' = x + \Delta x$$
,  $y' = y + \Delta y$ 



 $\triangleright$  Scaling: object scaled by a factor of Sx in the x-direction and Sy in the y-direction;

$$x' = xS_x$$
,  $y' = yS_y$ 





 $\triangleright$  Rotation: object rotated through the angle  $\theta$  about the origin of the coordinate system:

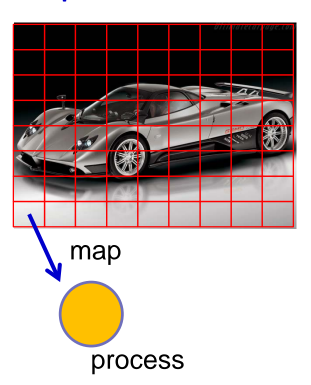
$$x' = x \cos \theta + y \sin \theta$$
$$y' = -x \sin \theta + y \cos \theta$$

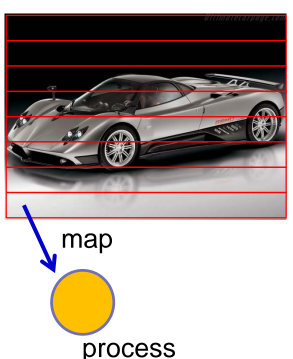


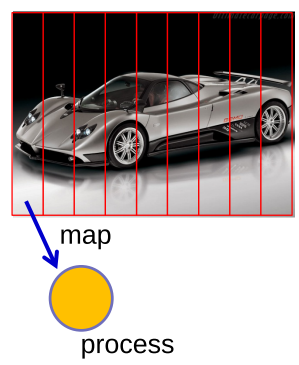




- partition
- partition
- Square region Row region Column region partition







# Pseudo-code for Image Shift

Partition region by ROW with width 10 480

```
//master process
for(i=0, row=0; i<48; i++, row+=10) // for each of 48 processes
    send(row, P<sub>i</sub>);
                                    // send row no.
for(i=0; i<480; i++) for(j=0; j<640; j++) temp_map[i][j] = 0; // initialize temp
for(i=0; i<(480*640); i++) {
                                                          // for each pixel
    recv(oldrow, oldcol, newrow, newcol, P<sub>ANY</sub>);
                                                          // accept new coordinates
    if !((newrow<0)||((newrow>=480)||(newcol<0)||((newcol>=640))
         temp map[newrow][newcol] = map[oldrow][oldcol];
for(i=0; i<480; i++) for(j=0; j<640; j++) map[i][j] = temp_map[i][j]; // update map
// slave process
recv (row, Pmaster);
for (oldrow = row; oldrow < (oldrow+10); oldrow++) // for each row in the partition
    for (oldcol = 0; oldcol < 640; oldcol++) {
                                                        // for each column in the row
                                                        // shift along x-dimension
         newrow = oldrow + delta x;
                                                        // shift along y-dimension
         newcol = oldcol + delta y;
         send(oldrow, oldcol, newrow, newcol, Pmaster); // send out new coordinates
```

640

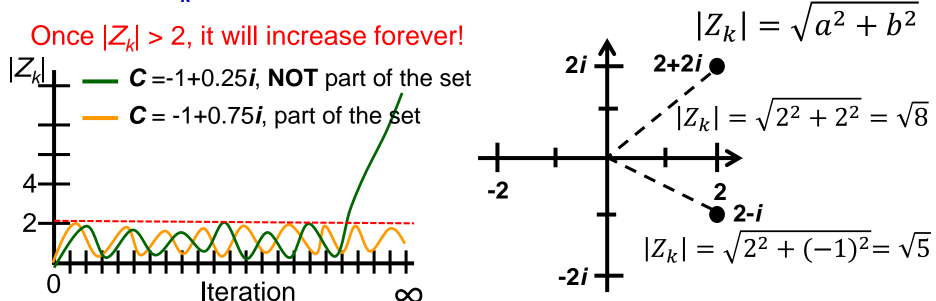
10 x 640



■ The Mandelbrot Set is a set of **complex numbers** that are quasi-stable when computed by iterating the function:

$$Z_0 = C$$
,  $Z_{k+1} = Z_k^2 + C$ 

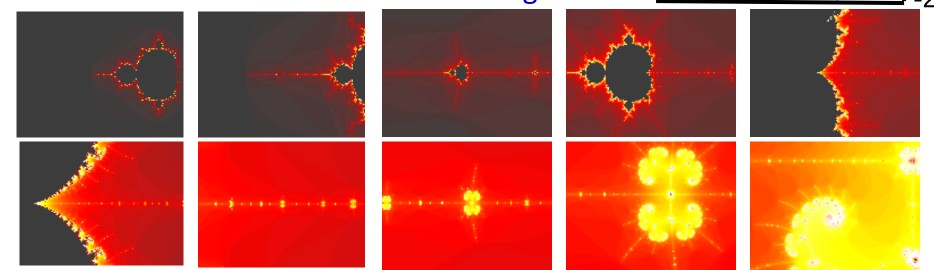
- $\triangleright$  **C** is some complex number:  $\mathbf{C} = a + b\mathbf{i}$
- $ightharpoonup Z_{k+1}$  is the (k+1)<sup>th</sup> iteration of the complex number
- ightharpoonup If  $|Z_k| \le 2$  for ANY  $k \to C$  belongs to Mandelbrot Set

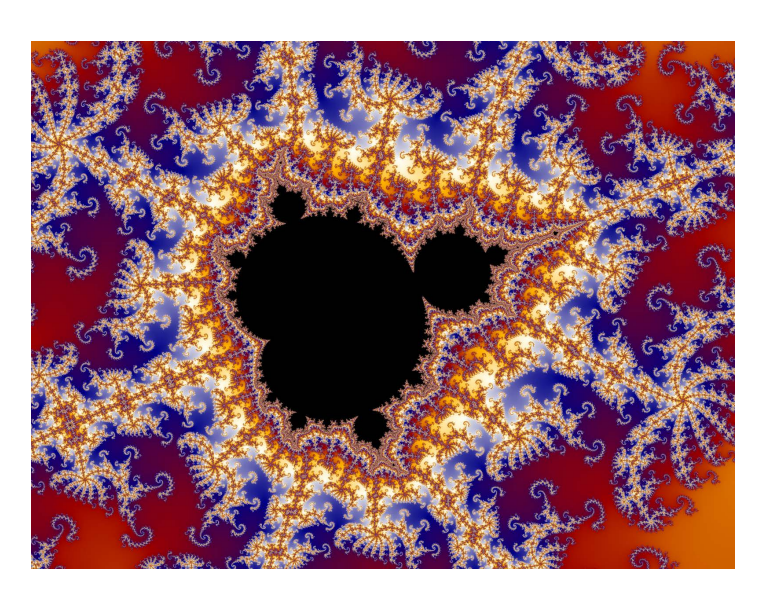


Parallel Programming - NTHU LSA Lab

#### Fractal

- What exact is Mandelbrot Set?
  - ➤ It is a **fractal**: An object that display self-similarity at various scale; Magnifying a fractal reveals small-scale details similar to the large-scale characteristics
  - ➤ After plotting the Mandelbrot Set determined by thousands of iteration:
  - Add color to the points outside the set &
     zoom in at the center of the image: -2<u>i</u>





# .

# Mandelbrot Set Program

■ Compute  $Z_{k+1} = Z_k^2 + C$ > Let  $C = C_{real} + C_{imag}i$ ,  $Z_k = Z_{real} + Z_{imag}i$ >  $Z_{k+1} = (Z_{real}^2 - Z_{imag}^2 + 2Z_{real}Z_{imag}i) + (C_{real} + Z_{real}^2)$ 

```
Struct complex {
    float real;
    float imag;
};
```

# Sequential Mandelbrot Set Program

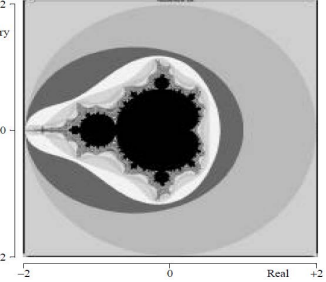
- Testing program:
  - Giving a complex number
  - $\triangleright$  Return the iteration number when  $|Z_k| > 2$
  - Let the maximum iteration is 256

```
int cal_pixel (complex c) {
  int count = 0;
                                   // number of iterations
                                   // maximum iteration is 256
  int max= 256;
  float temp, lengthsq;
                                   // initialize complex number z
  complex z;
  z.real = 0; z.imag = 0;
  do {
    temp = (z.real * z.real) - (z.imag * z.imag) + c.real; // compute next z.real
    z.imag = (2 * z.real * z.imag) + c.imag;
                                                         // compute next z.imag
    z.real = temp;
    lengthsq = (z.real * z.real) + (z.imag * z.imag);
    count++;
                                                     // update iteration counter
  } while ((lengthsq < 4.0) && (count < max));</pre>
  return count;
                                                                              12
```

# Sequential Mandelbrot Set Program

Scaling Coordinate Display Program:

- Plot the Mandelbrot Set from the coordinate system
- Color indicate the iteration number black=256, white=0
- Points are apart with a fixed distance read\_disk, imag\_dist



```
for (x=real_min; x < real_max; x += real_dist) {
    for (y=imag_min; y < imag_max; x += imag_dist) {
        c.real = x; c.img = y;
        color = cal_pixel (c);
        display(x, y, color);
    }
}</pre>
```

# Parallelizing Mandelbrot Set Program

■ Partition screen 640\*480 by row using 48 processes

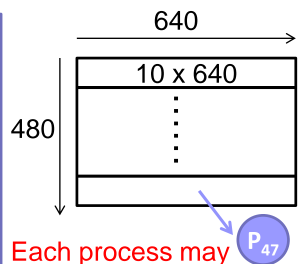
```
//master process
for(i=0, row=0; i<48; i++, row+=10)
    send(row, P<sub>i</sub>);

// for each process
// send row no.

for(i=0; i<(480*640); i++) {
    recv(&x, &y, &color, P<sub>ANY</sub>);
    display(x, y, color);
}

// for each pixel point
// receive coordinate/colors
// display pixel
}
```

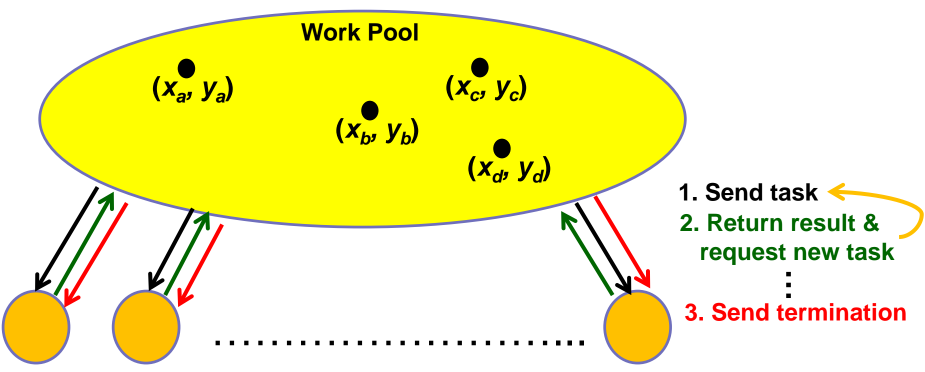
```
//slave process
recv (&row, P<sub>master</sub>);
for (x=0; x < 640; x++) {
    for (y=row; y < (row+10); y++) {
        c.real = min_real + (x * scale_real);
        c.imag = min_imag + (y * scale_image);
        color = cal_pixel (c);
        send(x, y, &color, P<sub>master</sub>);
    }
}
```



have different load!



- Work pool / Processor Farm
  - > Useful when tasks require different execution time
  - Dynamic load balancing



# Coding for Work Pool Approach

```
//master process
                                  // # of active processes
count = 0;
                             // row being sent
row = 0;
for (k=0; k<num_proc; k++) { // send initial row to each processes
    send(row, P<sub>i</sub> , data_tag);
    count++;
    row++;
do {
    recv(&slave, &r, color, P<sub>ANY</sub>, result_tag);
    count--;
    if (row < num_row) {</pre>
                              // keep sending until no new task
        send(row, P<sub>slave</sub>, data_tag); // send next row
         count++;
                                          Tag is needed to distinguish
         row++;
                                          between data and termination msg
    } else {
        send(row, P<sub>slave</sub>, terminate_tag); // terminate
    display(r, color);
                                         // display row
} while(count > 0);
```

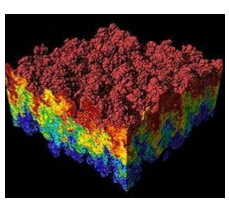


# Coding for Work Pool Approach

```
//slave process P ( i )
recv(&row, P<sub>master</sub> , source_tag);
while (source_tag == data_tag) { // keep receiving new task
    c.imag = min_imag + (row * scale_image);
    for (x=0; x<640; x++) {
        c.real = min_real + (x * scale_real);
        color[x] = cal_pixel (c); // compute color of a single row
    send(i, row, color, P<sub>master</sub>, result_tag); // send process id and results
    recv(&row, P<sub>master</sub> , source_tag);
```

# Example 3: Monte Carlo Methods

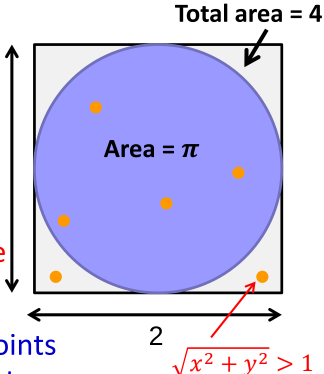
- Monte Carlo methods: a class of computational algorithms that rely on repeated random sampling to compute their results
  - Invented in 1940s by John von Neumann, Stanislaw Ulam and Nicholas Metropolis, while they were working on nuclear weapon (Manhattan Project)
  - Especially useful for simulating systems with many coupled degrees of freedom, such as fluids, disordered material



HISTORY

## Monte Carlo Methods --- $\pi$ calculation

- How to compute  $\pi$  ???
  - $\triangleright$  Definition of  $\pi$ : the area of a circle with unit radius
  - ightharpoonup We know:  $\frac{\text{Area of circle}}{\text{Area of square}} = \frac{\pi}{4}$
  - Randomly choose points from the square
  - $\triangleright$  Giving sufficient number of samples, the fraction of points **within** the circle will be  $\pi/4!!!$
  - ➤ E.g.: With 10,000 randomly sample points we expect 7854 points within the circle
    - $\rightarrow$  7854/10000 =  $\pi$ /4  $\rightarrow$   $\pi$  = 7854/10000\*4 = 3.1416



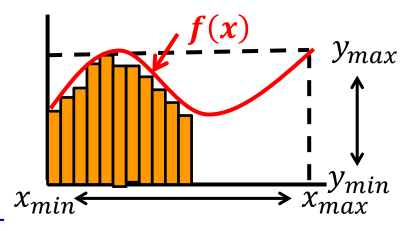
# Monte Carlo Methods --- Integral

- Monte Carlo Method can compute ANY **definite** integral!
  - max and min values of the integral must be known
  - Very inefficient....

#### ■ Method:

- $\triangleright$  Randomly choose point (x, y):
  - $\star x_{max} \le x \le x_{min}$
  - $y_{max} \le y \le y_{min}$
- Compute the area (integral) according to the ratio of points inside and outside the area
  - $\rightarrow$  just like the computation of  $\pi$

$$Area = \int_{x_{min}}^{x_{max}} f(x) dx$$



 $\triangleright$  Given any point (x, y), outside means : y > f(x)



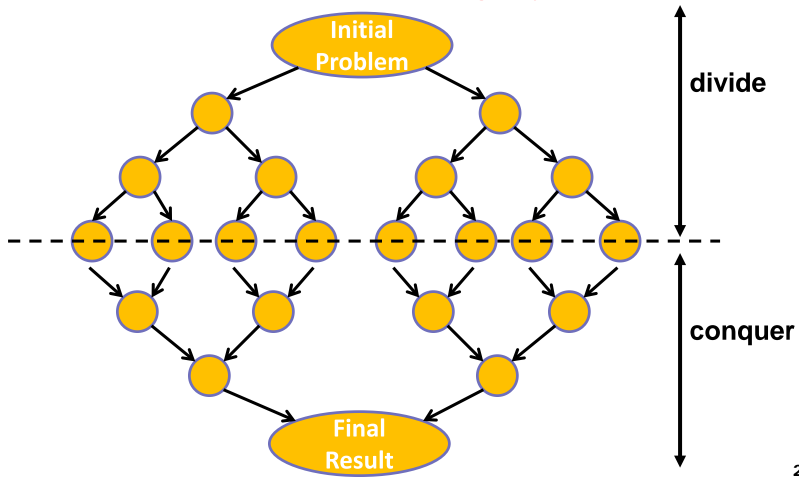
#### Outline

- Embarrassingly Computations
- Divide-And-Conquer Computations
  - Parallel Sorting Algorithm
  - N-Body Simulation
- Pipelined Computations
- Synchronous Computations



# What is Divide & Conquer

■ Recursively divide a problem into sub-problems that are of the same form as the larger problem



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#### Sorting

Re-arranging a list of numbers into increasing (strictly non-decreasing) order

> One of the most common and critical operation in

large data processing

Who gets admission to the school?

Which is the most relevant page?



Google 搜索 手气不错



Which

friend to be

LSA Lab

# Sorting in Parallel

- In sequential
  - $\triangleright$  We all know it is  $n \log n$
  - > i.e. n is the number of elements
- In parallel with *n* processors
  - Optimal parallel time complexity

$$\frac{O(n\log n)}{n} = O(\log n)$$

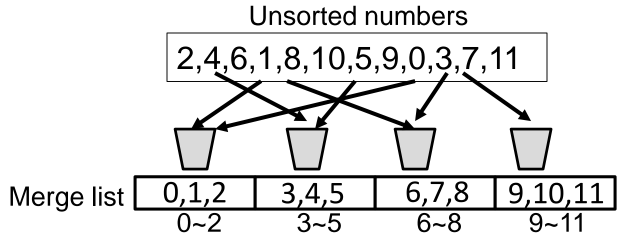
Has been obtained but the constant hidden in the order notation extremely large

LSA Lab 24



#### **Bucket Sort**

- Algorithm
  - 1. Range of numbers is divided into *m* equal regions
  - 2. One bucket is assigned for each region
  - 3. Place numbers to buckets based on the region
  - 4. Use sequential sort for each bucket



- Only effective if number of items per bucket is similar!!
  - Numbers should have a known interval ([max, min])
  - Numbers better to be uniformly distributed

# 10

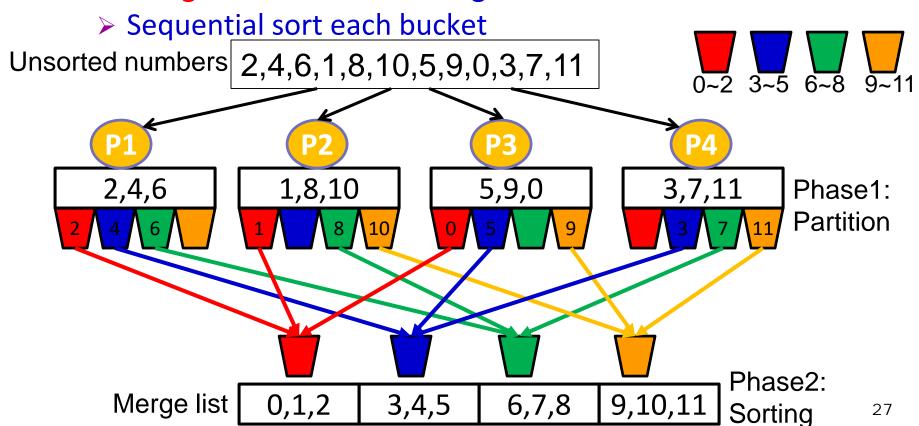
# **Complexity Analysis**

- Sequential:
  - 1. Distribute numbers to bucket: O(n)
  - 2. Sequential sort each bucket: (n/m)log(n/m) x m
  - Overall: O(n log(n/m))
- Parallelize sorting: one process per bucket
  - 1. Distribute numbers to bucket: O(n)
  - Sequential sort each bucket: (n/m)log(n/m)
  - Overall: O(n + n/m log(n/m))
  - > A single process must scan through all numbers in step1



#### Further Parallelized Bucket Sort

- Parallelize partitioning and sorting:
  - Partition numbers to m parts/processes
  - Each process divides its numbers to small buckets
  - Merge small buckets to large bucket





# Merge Sort

#### Divide & Conquer

 $\triangleright$  Sequential:  $O(n \log n)$ 

Sorted list

 $\triangleright$  Parallel: O(n)

 $T_{comm} = O\left(2\left(\frac{n}{2} + \frac{n}{4} + \dots + 1\right)\right) = O(n)$ 

#elements

 $T_{comp} = O\left(n + \frac{n}{2} + \frac{n}{4} + \cdots 2\right) = O(n)$ Unsorted list 6 5 3 5 6 Divide list Merge 3 5 5 8 6

LSA Lab



## **Quick Sort**

- Most popular sequential sorting algorithm
  - Parallel: Iteratively pick pivot and partition numbers
- Complexity:
  - $\triangleright$  Sequential:  $O(n \log n)$
  - $\triangleright$  Parallel: O(n)

Pivot Unsorted list

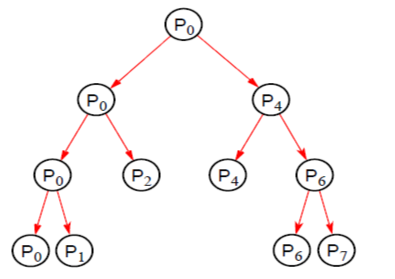
4 2 7 8 5 1 3 6

3 2 1 4 5 7 8 6

2 1 3 4 5 7 8 6

1 2 3 6 7 8

Still best choose in parallel? Not really & load might not be balanced



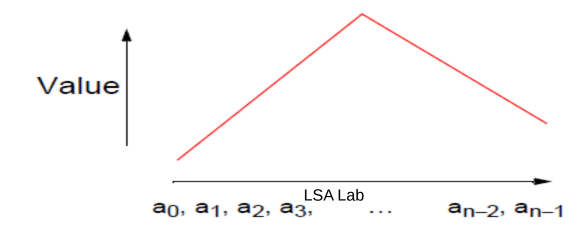
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# Bitonic Mergesort

- A parallel sorting algorithm based on bitonic sequence
  - > Remove the O(N) bottleneck for merging
- Monotonic sequence:
  - A sequence of increasing numbers

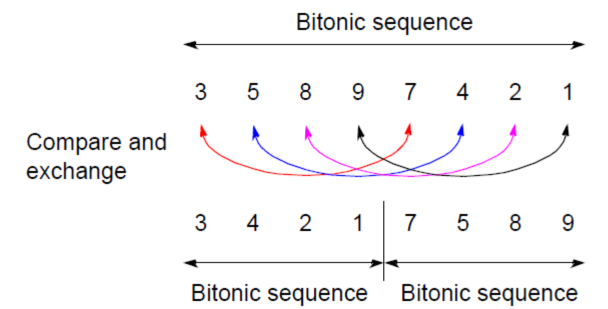
- Bitonic sequence:
  - Two sequences, one increasing and one decreasing <3, 5, 8, 9, 7, 4, 2, 1, >



# M.

# Characteristic of Bitonic Sequence

- If we perform a compare-and-exchange operation on  $a_i$  with  $a_{(i+n/2)}$  for all i
  - Get TWO bitonic sequences
  - The numbers in one sequence are all less than the numbers in the other sequence
- Example: <3,5,8,9,7,4,2,1>

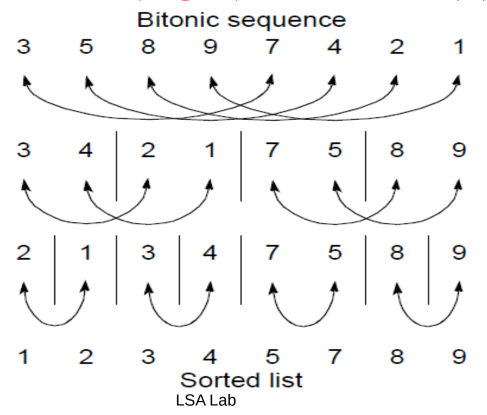




# **Bitonic Sorting Operation**

- Given a bitonic sequence, recursively performing compare-and-exchange will sort the list
- Complexity is only O(log n) instead of O(n)

Compare and exchange

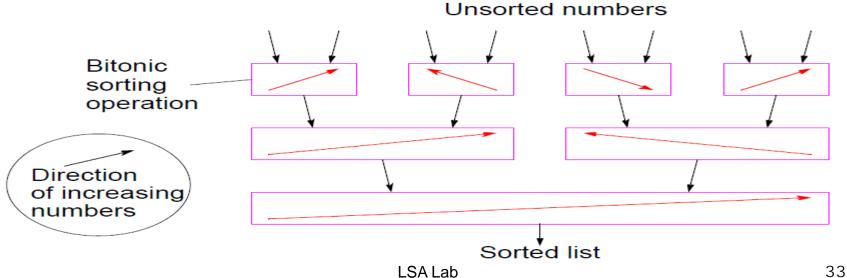




# Bitonic Mergesort

- Combine bitonic sorting into merge sorting
- Complexity:
  - $\triangleright$  Let  $n=2^k$ : there will be k phases
  - $\triangleright$  k<sup>th</sup> phase has size  $2^k \rightarrow$  need only k steps with bitonic sorting

$$T_p = O\left(\sum_{i=1}^k i\right) = O\left(\frac{k(k+1)}{2}\right) = O\left(\frac{\log n(\log n)}{2}\right) = O(\log^2 n)$$





#### Rank Sort

- Count the number of numbers that are smaller than each of the selected number
- Sequential:  $O(n^2)$

```
for(i=0; i<n; i++) {
    rank=0;
    for(j=0; j<n; j++) {
        if (a[i] < a[j]) rank++;
      }
    b[rank] = a[i];
}</pre>
```

```
rank: 1 0 2 3
a[i]: 4 1 6 8
b[i]: 1 4 6 8
```

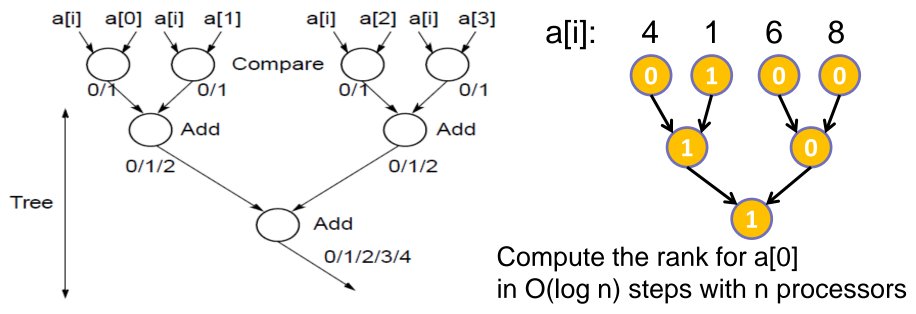
- Parallel: O(n)
  - use one processor for each number

LSA Lab 34



#### Rank Sort with n<sup>2</sup> Processors

- We can use n processor to compute the rank of a number in parallel (summation parallel algo.)
- Time complexity?  $O(\log n)$
- How about parallel efficiency? Actually decreased!



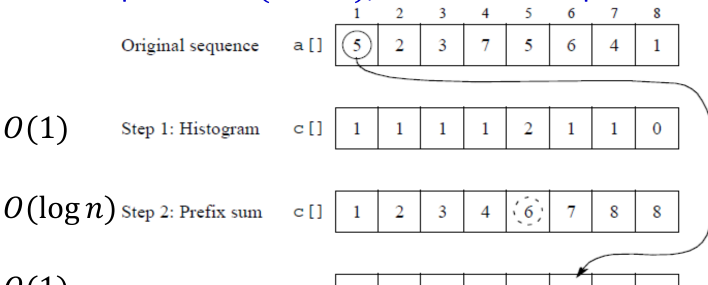
LSA Lab 35



# **Counting Sort**

- If the numbers are integers and the range is known
  - $\triangleright$  Coding the rank sort algorithm to reduce the sequential time from  $O(n^2)$  to O(n)
- Use array c[] to count the histogram of values
- Complexity





Move 5 to position 6. Then decrement c [5]

O(1)

Step 3: Sort

b[] 1 2 3 4 5 5 7

Final sorted sequence

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## Summary

#### ■ With #processors = n

	Sequential	Parallel	
Bucket Sort	$O(n \log n)$	O(n)	
Merge Sort	$O(n \log n)$	O(n)	
Quick Sort	$O(n \log n)$	O(n)	
Bitonic	$O(n \log n)$	$O(\log^2 n)$	
mergesort	2		With n <sup>2</sup>
Rank Sort	$O(n^2)$	$O(n)$ $O(\log n)$	processors
Counting Sort*	O(n)	$O(\log n)$	

<sup>\*</sup>Special case with known value range

LSA Lab 37

## Example 2: N-Body Problem

- Newtonian laws of physics
  - > The gravitational force between two bodies of masses  $m_a \& m_b$ :

$$F = \frac{Gm_am_b}{r^2}$$

Subject to the force, acceleration occurs  $F = m \times a$ 

- Let the time interval be  $\Delta t$  & current velocity  $v^t$ , position  $x^t$ 
  - $\triangleright$  New velocity  $v^{t+1}$ :

$$F = m \frac{v^{t+1} - v^t}{\Delta t} \Rightarrow v^{t+1} = v^t + \frac{F\Delta t}{m}$$

 $\triangleright$  New position  $x^{t+1}$ :

$$x^{t+1} = x^t + v^{t+1} \Delta t$$





#### Three-Dimensional Space

- Considering 2 bodies at  $(x_a, y_a, z_a) \& (x_b, y_b, z_b)$  $r = \sqrt{(x_a - x_b)^2 + (y_a - y_b)^2 + (z_a - z_b)^2}$
- The forces, velocities and positions can be resolved in the three direction independently

$$F_{x} = \frac{Gm_{a}m_{b}}{r^{2}} \left( \frac{x_{b} - x_{a}}{r} \right)$$

$$F_{y} = \frac{Gm_{a}m_{b}}{r^{2}}\left(\frac{y_{b} - y_{a}}{r}\right)$$

$$F_z = \frac{Gm_am_b}{r^2} \left(\frac{z_b - z_a}{r}\right)$$

#### N-Body Sequential Code

Assume all bodies have the same mass m

```
for (t=0; t<T; t++) {
   for (i=0; i<N; i++) {
       F = Compute_Force(i); // compute force in O(N^2)
       v_new[i] = v[i] + F *dt / m; // compute new velocity
       x_new[i] = x[i] + v_new[i] * dt; // compute new position
   for(i=0; i<N; i++){
       x[i] = x_new[i];
                                    // update position
                                    // update velocity
       v[i] = v_new[i];
```

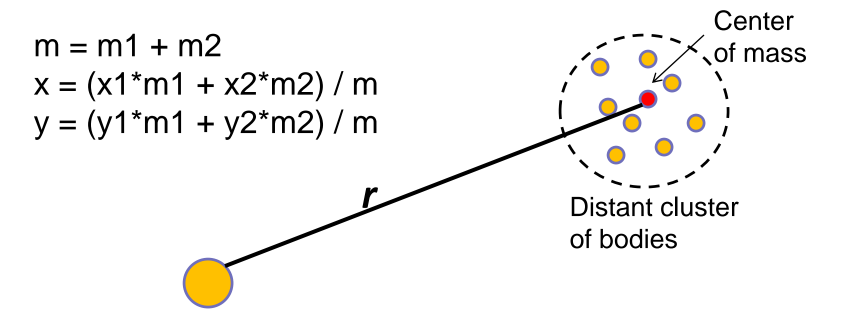
■ Non-feasible as N increases due to  $O(N^2)$  complexity



#### Approximate Algorithms

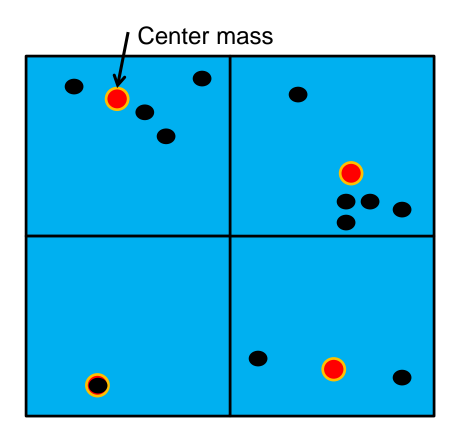
 Reduce time complexity by approximating a cluster of bodies as a single distant body

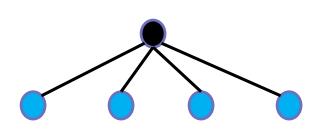
How to find those clusters of bodies?





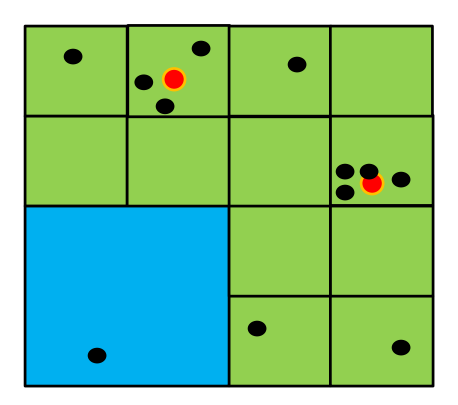
- Step1: Recursively divide space by two in each dimensions
  - > Record the center mass and position of each internal node

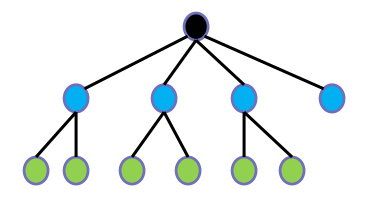






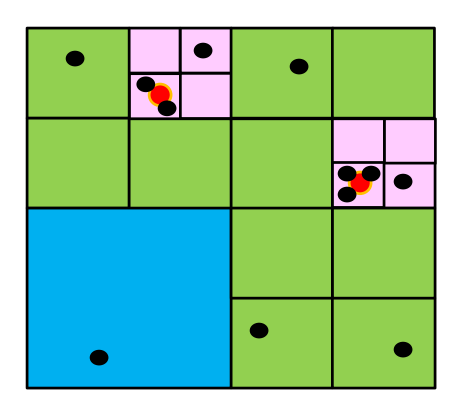
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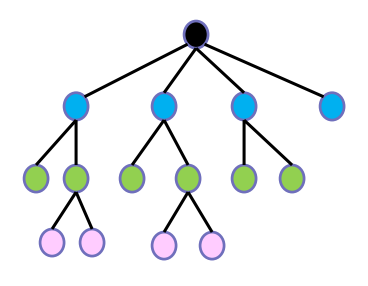






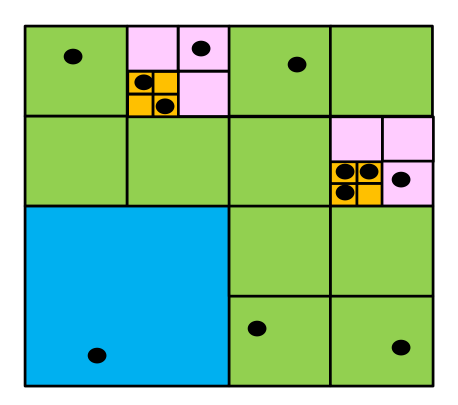
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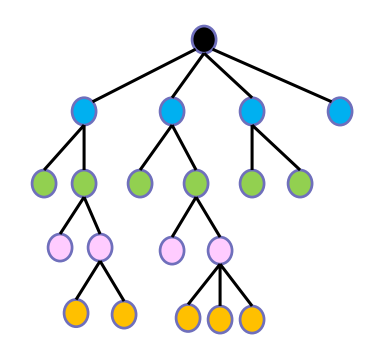






- Step1: Recursively divide space by two in each dimensions
  - > Record the center mass and position of each internal node

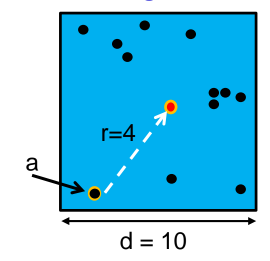


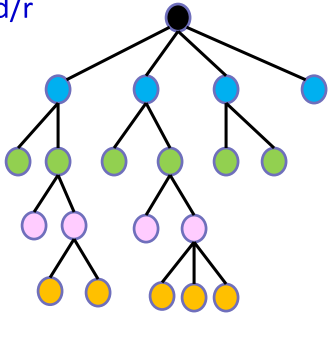


- Step2: Compute approximate forces on each object
  - 1. traverse the nodes of the tree, starting from the root.
  - 2. If the center-of-mass of an **internal node** is **sufficiently far** from the body, approximate the internal node as a single body
  - $\triangleright$  Far is determined by a parameter:  $\theta=d/r$ 
    - r: the distance between the body and the node's center-of-mass
    - ♦ d: the width of the region

Example:  $\theta$ =0.5

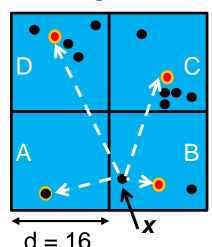
 $d/r=2.5 > \theta$ 

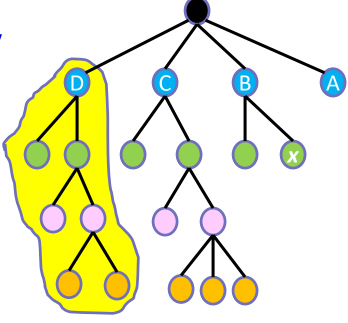




- Step2: Compute approximate forces on each object
  - 1. Traverse the nodes of the tree, starting from the root.
  - 2. If the center-of-mass of an **internal node** is **sufficiently far** from the body, approximate the internal node as a **single body**
  - $\triangleright$  Far means  $d/r < \theta$  (e.t.  $0 < \theta < 1$ )
    - r: the distance between the body and the node's center-of-mass
    - ♦ d: the width of the region

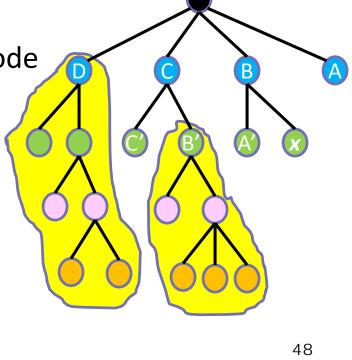
Example: 
$$\theta$$
=1  
 $d/r_A$ =16/10 >  $\theta$   
 $d/r_B$ =16/2 >  $\theta$   
 $d/r_C$ =16/15 >  $\theta$   
 $d/r_D$ =16/20 <  $\theta$ 



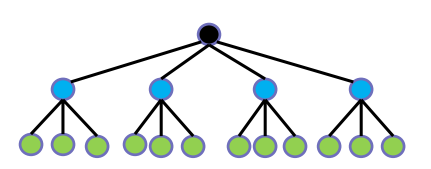


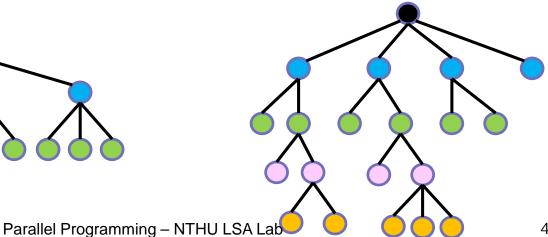
- Step2: Compute approximate forces on each object
  - 3. If it is a leaf node, calculate the force and add to the object.
  - 4. Otherwise, recursively compute the force from children of the internal node.

Example:  $\theta=1$   $d/r_{A'}=8/7 > \theta \rightarrow A'$  is a leaf node  $d/r_{B'}=8/15 < \theta \rightarrow B'$  treated like a single node  $d/r_{C'}=8/20 < \theta \rightarrow C'$  is a leaf node



- lacksquare 0 controls the accuracy and approximation error of the algorithm
  - $\Rightarrow \theta = 0 \Rightarrow d/r$  ALWAYS larger than  $\theta \Rightarrow$  same as brute force
  - $\Rightarrow$  0 = 1  $\Rightarrow$  most likely only need to consider the object within the same cluster/region
- If the tree is balanced, the complexity is  $O(n \log n)$ 
  - > But in general, the tree could be very unbalanced .......
- The tree must be re-built for each time interval



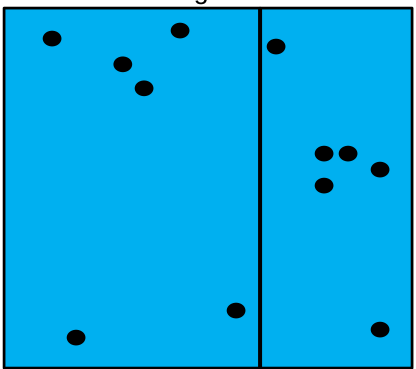


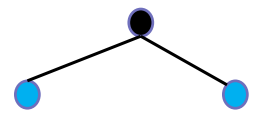
# 100

## Orthogonal Recursive Bisection Method

Recursively evenly divide space with the same number of bodies in each of the dimensions

Divide along x dimension



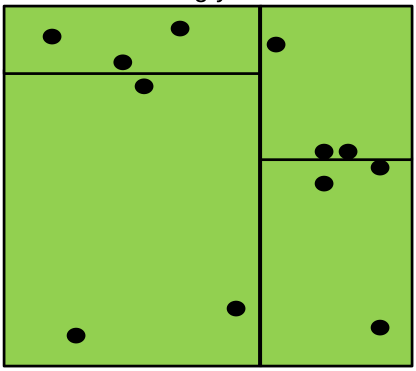


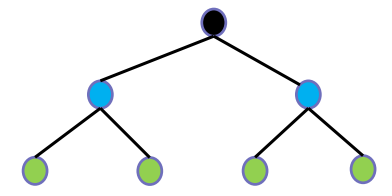
# re.

## Orthogonal Recursive Bisection Method

Recursively evenly divide space with the same number of bodies in each of the dimensions

Divide along y dimension



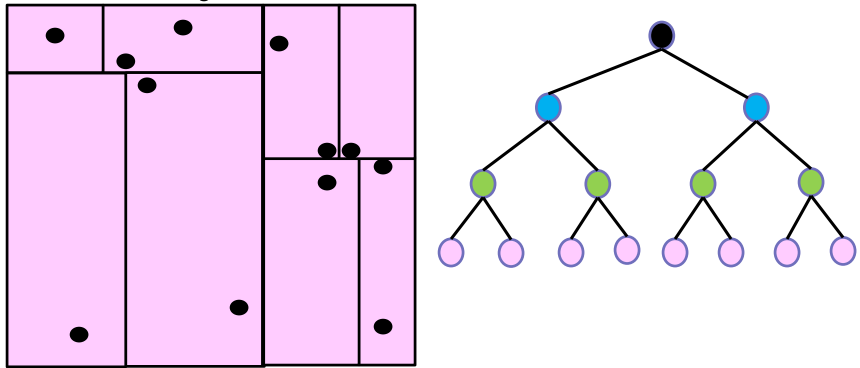




## Orthogonal Recursive Bisection Method

Recursively evenly divide space with the same number of bodies in each of the dimensions

Divide along x dimension

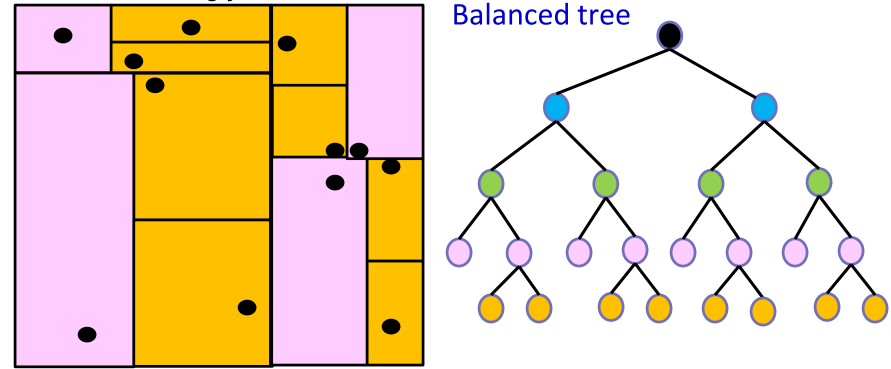


# 100

## Orthogonal Recursive Bisection Method

Recursively evenly divide space with the same number of bodies in each of the dimensions

Divide along y dimension

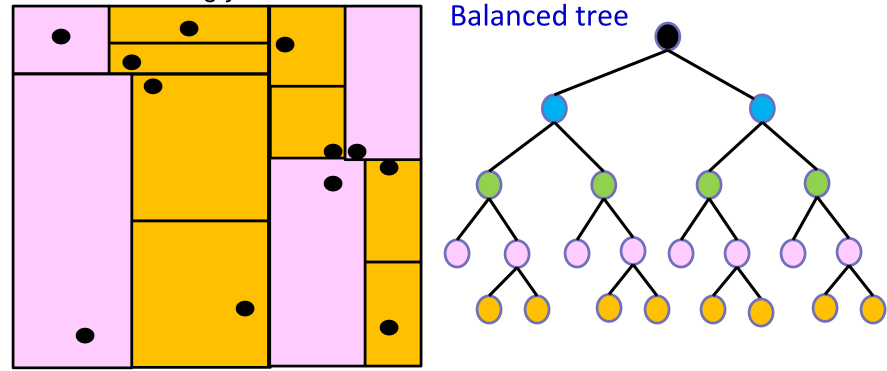




## Orthogonal Recursive Bisection Method

- It is more balanced, but less accurate
  - Objects close to each other may not in the same cluster

Divide along y dimension





#### Outline

- Embarrassingly Computations
- Divide-And-Conquer Computations
- Pipelined Computations
  - Adding Numbers
  - Sorting Numbers
  - Linear Equation Solver
- Synchronous Computations

## What is Pipelined Computations

- A problem is divided into a series of tasks
- Tasks have to be completed one after the other
- Each task will be executed by a separate process or processor



$$\rightarrow$$
 P0  $\rightarrow$  P1  $\rightarrow$  P2  $\rightarrow$  P3  $\rightarrow$ 



## Types of Pipelined Computations

- Pipelined approach can provide increased speed under three types of computations:
- If more than one instance of the complete problem is to be executed
- If a single instance has a series of data items must be processed, each requiring multiple operations
- If information to start the next process can be passed forward before the process has completed all its internal operations

## Type 1 Pipelined Computations

1. If more than one **instance** of the complete problem is to be executed

	_		p-1			_		m			
						Instance	Instance	Instance	Instance	Instance	
$P_5$						1	2	3	4	5	
D					Instance	Instance	Instance	Instance	Instance	Instance	
$P_4$					1	2	3	4	5	6	
D				Instance							
$P_3$				1	2	3	4	5	6	7	
$P_2$			Instance								
			1	2	3	4	5	6	7		
$P_1$		Instance									
		1	2	3	4	5	6	7			
$P_0$	Instance										
	1	2	3	4	5	6	7				
	- 1 -	- •		- •	1.0		_				

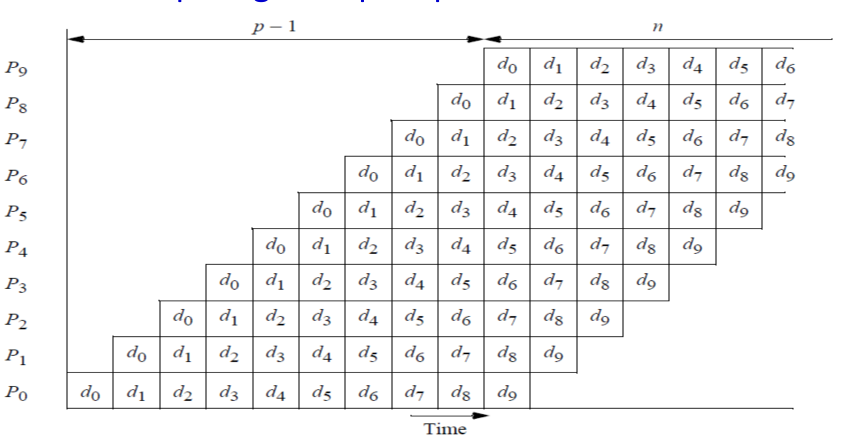
(Alternative space-time diagram) Time

- After the first (p-1) cycles, one problem instance is completed in each pipeline cycle
- The number of instance should be >> the number of



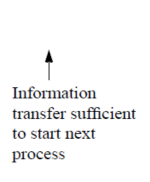
## Type 2 Pipelined Computations

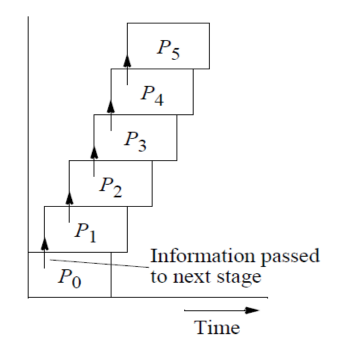
 If a series of data items must be processed, each requiring multiple operations

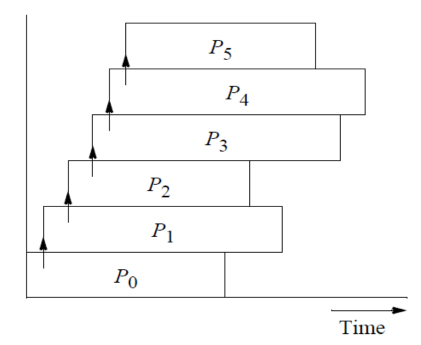


## **Types 3 Pipelined Computations**

 Only one problem instance, but each process can pass on information to the next process, before it has completed





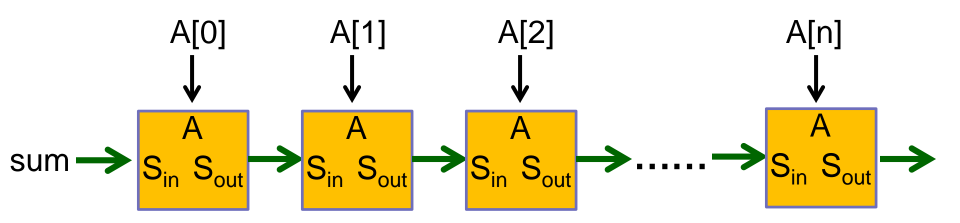


(a) Processes with the same execution time

(b) Processes not with the same execution time

# **Example1: Adding Numbers**

- Compute sum of an array:
  - for(i=0; i<n; i++) sum += A[i]</pre>
- Pipeline for an unfolded loop:
  - $\triangleright$  sum += A[0], sum += A[1], sum += A[2], .....





#### Example1: Adding Numbers

■ The basic code for Pi:

```
recv(&sum, P<sub>i-1</sub>);
sum += number;
send(&sum, P<sub>i+1</sub>);
```

■ For the first process, P0:

```
send(&sum, P<sub>i+1</sub>);
```

■ For the last process, Pn-1:

```
recv(&sum, P<sub>i-1</sub>);
sum += number;
```

■ SPMD Program:

```
// code for process Pi
if (Pi != P0) {
    recv(&sum, P<sub>i-1</sub>);
    sum += number;
}
if (Pi != Pn) {
    send(&sum, P<sub>i+1</sub>);
}
```



## **Example2: Sorting Numbers**

#### ■ Insertion Sort:



- 5 2

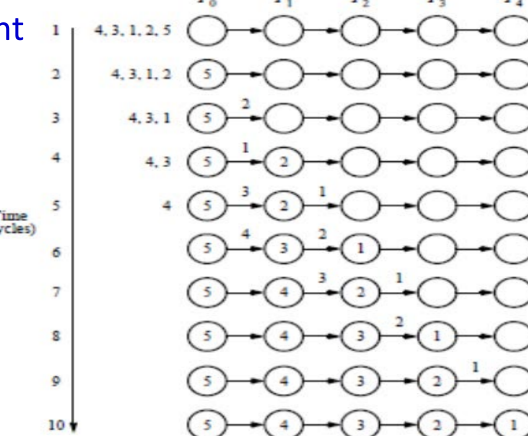


## **Example2: Sorting Numbers**

- Insertion Sort:
  - > Each process holds one number
  - Compare & move the smaller

```
number to the right
```

```
recv(&number, P<sub>i-1</sub>);
if (number > x) {
    send(&x, P<sub>i+1</sub>);
    x= number;
} else {
    send(&number, P<sub>i+1</sub>);
}
```



# v.

## Example 3: Linear Equation Solver

- Special linear equations of "upper-triangular" form
  - > a's and b's are constants, x's are unknown to be found

$$\begin{pmatrix}
a_{n-1,0} & a_{n-1,1} & a_{n-1,2} & \dots & a_{n-1,n-1} \\
a_{n-2,0} & a_{n-2,1} & \dots & a_{n-2,n-2} & 0 \\
\vdots & \vdots & \vdots & 0 & 0 \\
a_{1,0} & a_{1,1} & 0 & 0 & 0 \\
a_{0,0} & 0 & 0 & 0
\end{pmatrix}
\begin{pmatrix}
x_0 \\
x_1 \\
\vdots \\
x_{n-2} \\
x_{n-1}
\end{pmatrix} = \begin{pmatrix}
b_{n-1} \\
b_{n-2} \\
\vdots \\
b_1 \\
b_0
\end{pmatrix}$$

$$a_{n-1,0}x_0 + a_{n-1,1}x_1 + a_{n-1,2}x_2 + \dots + a_{n-1,n-1}x_{n-1} = \mathbf{b}_{n-1}$$

$$\vdots$$

$$a_{2,0}x_0 + a_{2,1}x_1 + a_{2,2}x_2 = \mathbf{b}_2$$

$$a_{1,0}x_0 + a_{1,1}x_1 = \mathbf{b}_1$$

$$a_{0,0}x_0 = \mathbf{b}_0$$



### Example 3: Linear Equation Solver

#### Back Substitution

 $\triangleright x_0$  is found from the last equation

$$x_0 = \frac{b_0}{a_{0,0}}$$

 $\triangleright$  Value for  $x_0$  is substituted into the next equation

$$x_1 = \frac{b_1 - a_{1,0} x_0}{a_{1,1}}$$

 $\triangleright$  Values for  $x_0$ ,  $x_1$  are substituted into the next equation

$$x_2 = \frac{b_2 - a_{2,0}x_0 - a_{2,1}x_1}{a_{2,2}}$$

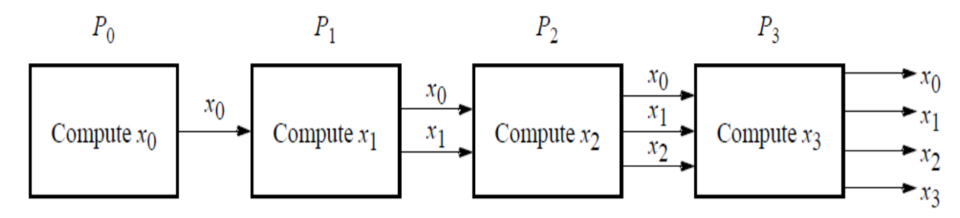
> So on until all unknowns are found ...

$$x_i = \frac{b_i - \sum_{j=0}^{i-1} a_{i,j} x_j}{a_{i,i}}$$

# 10

## Example 3: Linear Equation Solver

■ First pipeline stage computes  $x_0$  and passes  $x_0$  onto the second stage, which computes  $x_1$  from  $x_0$  and passes both  $x_0$  and  $x_1$  onto the next stage, which computes  $x_2$  from  $x_0$  and  $x_1$ , and so on



## Example 3: Linear Equation Solver

#### ■ Parallel Code

```
// code for Pi sum = 0; for (j=0; j<i; j++) { // compute partial result recv(&x[j], P_{i-1}); // once data is available send(&x[j], P_{i+1}); sum += a[i][j]*x[j]; } x[i] = (b[i] - sum) / a[i][j]; // send out final result to send(&x[j], P_{i+1}); // next process

P_5
P_4
P_3
P_2
P_1
P_1
Final computed value passed onward
```

#### ■ Time complexity:

- $> O(n^2)$  without passing forward
- $\triangleright O(n)$  with passing forward

Time



#### Outline

- Embarrassingly Computations
- Divide-And-Conquer Computations
- Pipelined Computations
- Synchronous Computations
  - Prefix Sum
  - System of Linear Equations



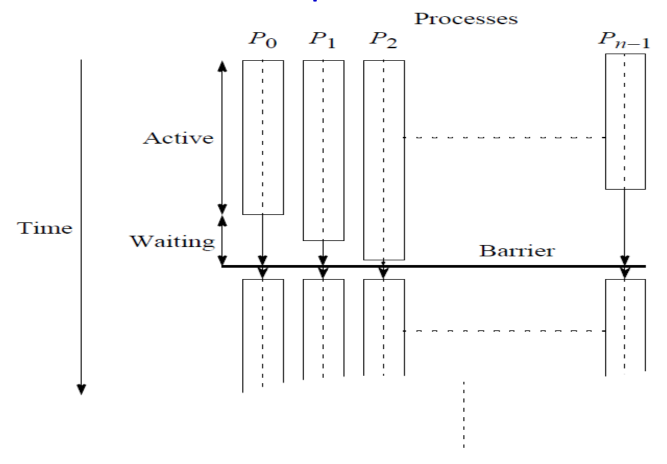
### Synchronous Computations

- **Definition:** all the processes *synchronized* at regular points
- Barrier: Basic mechanism for synchronizing processes
  - Inserted at the point in each process where it must wait
  - Message (token) is passed among processes for synchronization
- Deadlock: Common problem occurs from synchronization
  - Two or multiple processes waiting for each other



#### Barrier

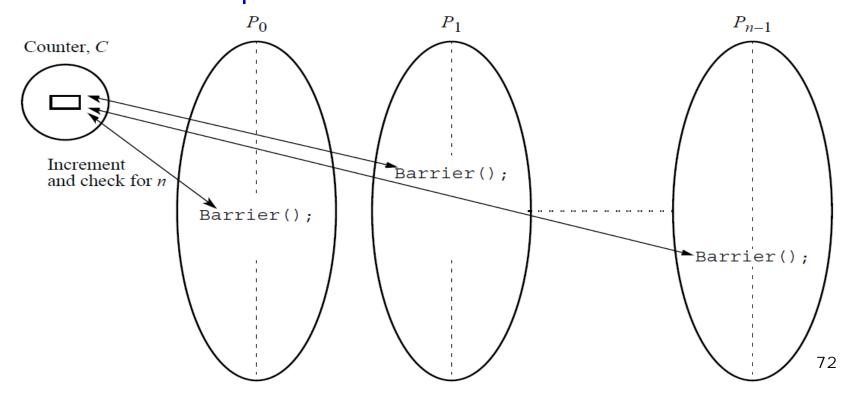
■ All processes can only continue from this POINT when all the processes have reached it



# re.

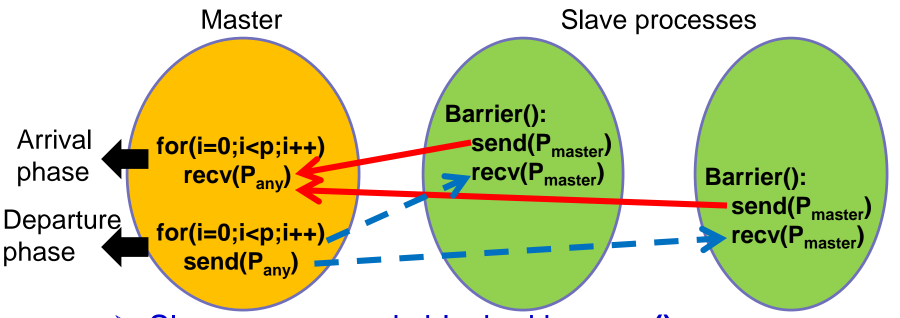
#### Counter Barrier Implementation

- A.k.a: Linear Barrier
  - Centralized counter: count # of processes reaching the barrier
  - Increase & check the counter for each barrier call
  - Processes is locked by the barrier call until counter == # processes



## Counter Barrier Implementation

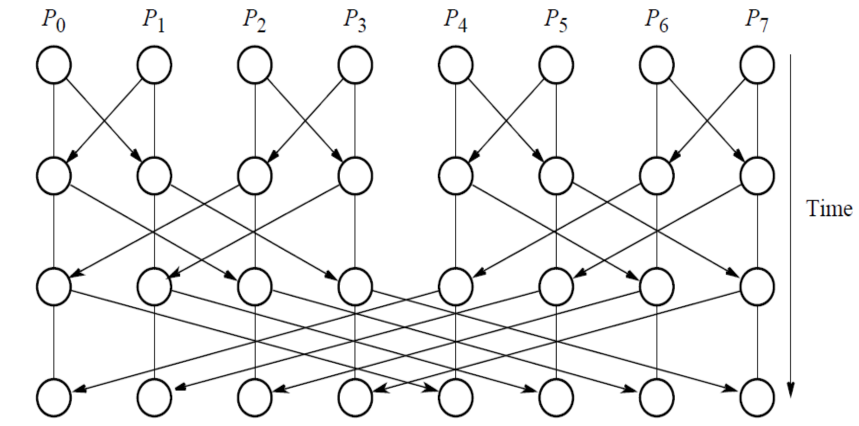
- Counter-based barrier often have two phases
  - > Arrival phase: a process enters arrival phase and does not leave this phase until all processes have arrived in this phase
  - Departure phase: Processes are released after moving to the departure phase



- Slave processes is blocked by recv()
- Master could be a bottleneck



At stage i, each process passes a token to the process with 2i distance away



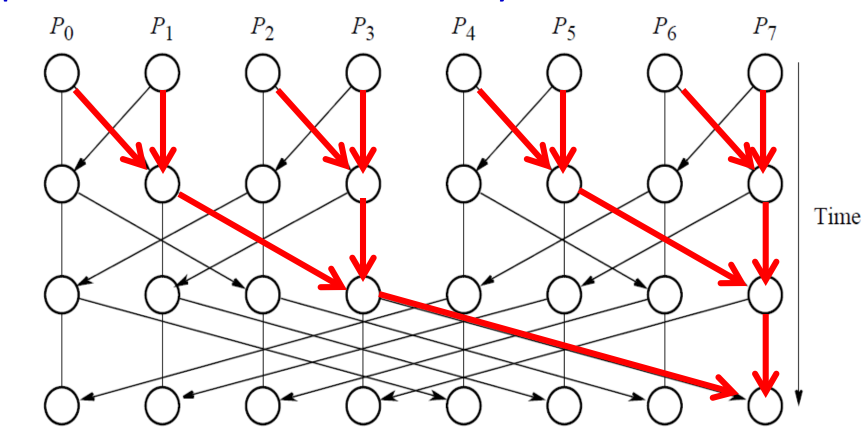
1st stage

2nd stage

3rd stage

# **Butterfly Barrier Implementation**

At stage i, each process passes a token to the process with 2i distance away



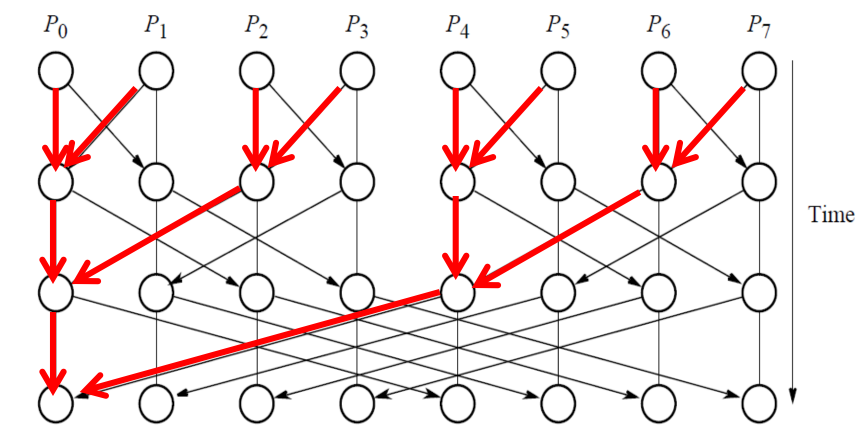
1st stage

2nd stage

3rd stage

# **Butterfly Barrier Implementation**

At stage i, each process passes a token to the process with 2i distance away



1st stage

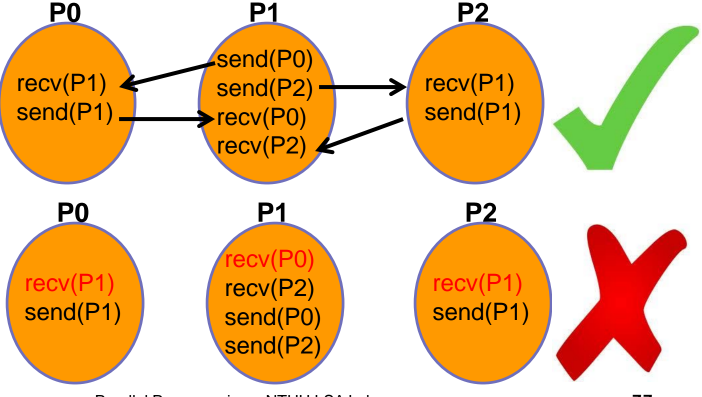
2nd stage

3rd stage

### Deadlock Problem

 A set of blocked processes each holding some resources and waiting to acquire a resource held by another process in the set

**■** Example:



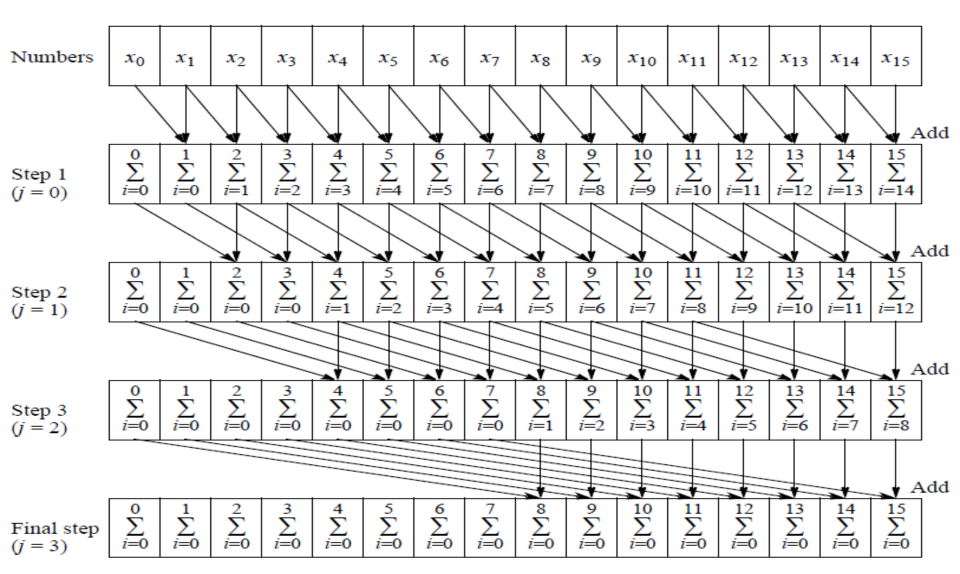


## Example 1: Prefix Sum

- Given a list of numbers  $x_0, x_1, ..., x_{n-1}$ , compute all *partial* summations
  - $> x_0; x_0 + x_1; x_0 + x_1 + x_2; \dots$
  - ➤ Could also replace operator + with AND, OR, \*, etc.
- Example:
  - x = 1,2,3,4,5
  - > Sum = 1,3,6,10,15
- Sequential code: O(n²)

```
//sequential code
for(i = 0; i < n; i++) {
    sum[i] = 0;
    for (j = 0; j <= i; j++)
        sum[i] = sum[i] + x[j];
}
```

## Data Parallelism Solution





### Data Parallelism Code

■ Sequential Code: O(n²), optimal: O(n)

```
for (j = 0; j < log(n); j++) /* at each step */
for (i = 2^{j}; i < n; i++) /* add to accumulating sum */
x[i] = x[i] + x[i - 2^{j}]
```

Parallel Code: O(log n)

```
for (j = 0; j < log(n); j++) /* at each step */
forall (i = 0; i < n; i++) /* add to accumulating sum */
if (i >= 2^{j}) x[i] = x[i] + x[i - 2^{j}];
```



Each iteration composed of several processes that start together at beginning of iteration and next iteration cannot begin until all processes have finished previous iteration

### openMP

#### MPI

```
for (j=0; j<n; j++) { // each iteration
  i = myrank;
  body(i);
  barrier(mygroup);
}</pre>
```

# Example 2: System of Linear Equations

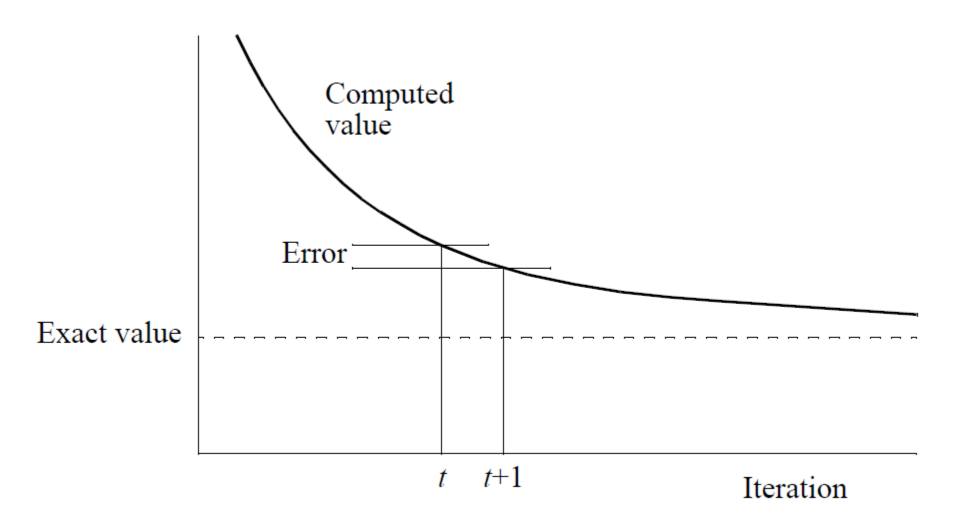
System of linear equations

$$\begin{pmatrix}
a_{0,0} & a_{0,1} & a_{0,2} & \dots & a_{0,n-1} \\
a_{1,0} & a_{1,1} & a_{1,2} & \dots & a_{1,n-1} \\
\vdots & \vdots & \vdots & \vdots & \vdots \\
a_{n-2,0} & a_{n-2,1} & a_{n-2,2} & \dots & a_{n-2,n-1} \\
a_{n-1,0} & a_{n-1,1} & a_{n-1,2} & \dots & a_{n-1,n-1}
\end{pmatrix}
\begin{pmatrix}
x_0 \\
x_1 \\
\vdots \\
x_{n-2} \\
x_{n-1}
\end{pmatrix} =
\begin{pmatrix}
b_0 \\
b_1 \\
\vdots \\
b_{n-2} \\
b_{n-1}
\end{pmatrix}$$

- Jacobi iteration algorithm:
  - > Convert *i*th iteration to  $x_i = \frac{1}{a_{i,i}} [b_i \sum_{i \neq j} a_{i,j} x_j]$
  - $\triangleright$  Initial guess with  $x_i = b_i$ , and calculate new  $x_i$  values
  - ightharpoonupRepeat until  $\begin{vmatrix} x_i & t-1 \\ x_i & -x_i \end{vmatrix}$  < error tolerance Parallel Programming NTHU LSA Lab



# Jacobi iteration algorithm



# Jacobi iteration algorithm example

$$\begin{cases} -x_0 + 2x_1 - x_2 = 2\\ 2x_0 + x_1 - 2x_2 = 2\\ 2x_0 - x_1 + 2x_2 = 2 \end{cases}$$
$$x_i = \frac{1}{a_{i,i}} [b_i - \sum_{i \neq i} a_{i,j} x_j]$$

$$\begin{cases} x_0 = 2 - \frac{(2x_1 - x_2)}{-1} \\ x_1 = 2 - \frac{(2x_0 - 2x_2)}{1} \\ x_2 = 2 - \frac{(2x_0 - x_1)}{2} \end{cases}$$

■ Iter1:  $x_0^1 = 2$ ,  $x_1^1 = 2$ ,  $x_2^1 = 2$ 

⇒ 
$$x_0^2 = 2 - \frac{2x_1^1 - x_2^1}{-1} = 4$$
,  $x_1^2 = 2$ ,  $x_2^2 = 1$   
⇒  $e_0 = |2 - 4| = 2$ ,  $e_1 = 0$ ,  $e_2 = 1$ 

$$rac{1}{2}e_0 = |2 - 4| = 2, \qquad e_1 = 0, \qquad e_2 = 1$$

■ ilter2: 
$$x_0^2 = 2 - \frac{2x_1^2 - x_2^2}{-1} = 5$$
,  $x_1^3 = -2$ ,  $x_2^3 = -1$ 

$$ightharpoonup e_0 = |4 - 5| = 1, \qquad e_1 = 4, \qquad e_2 = 2$$

## Jacobi iteration algorithm

- Sequential Code
  - > a[][] and b[] holding constants in the equations
  - x[] holding unknowns
  - fixed number of iterations

```
x_{i} = \frac{1}{a_{i,i}} [b_{i} - \sum_{i \neq j} a_{i,j} x_{j}]
```

```
for (i = 0; i < n; i++) x[i] = b[i]; /*initialize unknowns*/
for (iteration = 0; iteration < limit; iteration++) {
   for (i = 0; i < n; i++) { /* for each unknown */
       sum = -a[i][i] * x[i];
       for (j = 0; j < n; j++) /* compute summation */
          sum = sum + a[i][j] * x[j];
       new_x[i] = (b[i] - sum) / a[i][i]; /*compute unknown*/
   for (i = 0; i < n; i++) x[i] = new_x[i]; /*update to new values*/
```



## Jacobi iteration algorithm

- Parallel Code
  - Process i handles unknown x[i]

```
x[i] = b[i]; /*initialize unknown*/
for (iteration = 0; iteration < limit; iteration++) {
   sum = -a[i][i] * x[i];
   for (j = 0; j < n; j++) /* compute summation */
       sum = sum + a[i][j] * x[j];
   new_x[i] = (b[i] - sum) / a[i][i]; /* compute unknown */
   allGather(&new_x[i]); /* gather & broadcast new value */
                                    /* wait for all processes */
   barrier();
```