THE ARCTIC UNIVERSITY OF NORWAY

# Lecture 4: Embarrassingly Parallel Computations

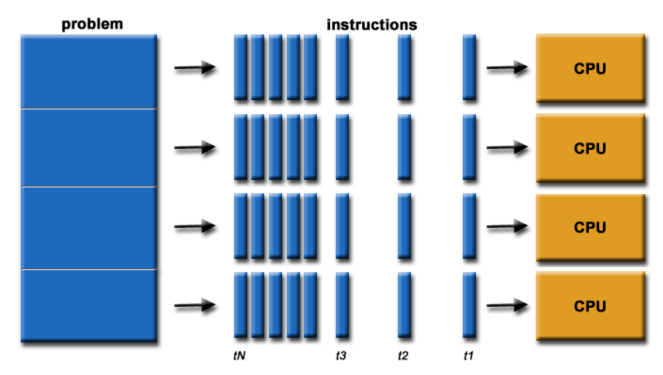
Parallel Programming (INF-3201)

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### **Parallel computations**

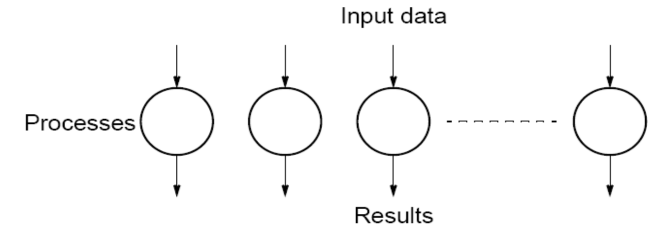
- A program is run using multiple CPUs
- A problem is broken into discrete parts that can be solved concurrently
- Each part is further broken down to a series of instructions
- Instructions from each part execute simultaneously on different CPUs



Source: Blaise Barney, "Introduction to Parallel Computing", Livermore Computing.

## **Embarrassingly Parallel Computations**

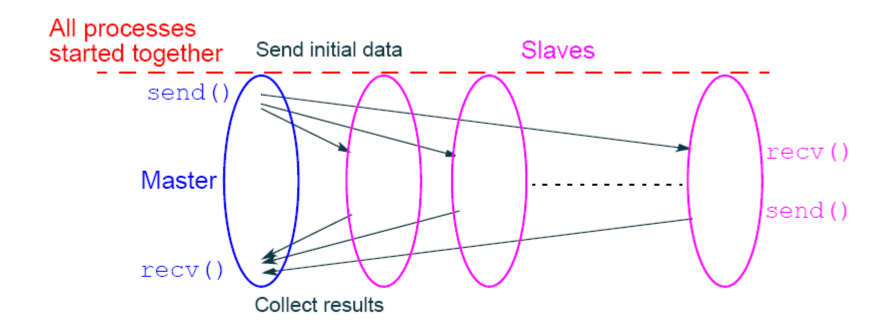
A computation that can obviously be divided into a number of completely independent parts, each of which can be executed by a separate process(or).



No communication or very little communication between processes

- Each process can do its tasks without any interaction with other processes
- Speedup, message-passing, SPMD

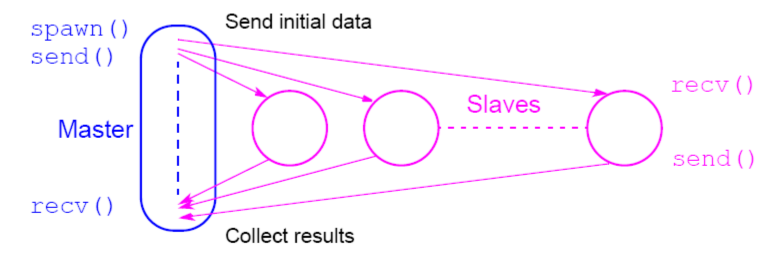
# Static process creation Master-slave approach



Usual MPI approach

# Dynamic process creation Master-slave approach

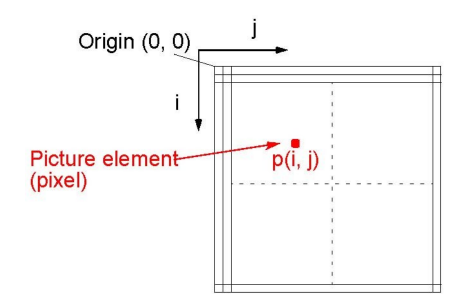
#### Start Master initially



(PVM approach)

#### **Example: Low level image processing**

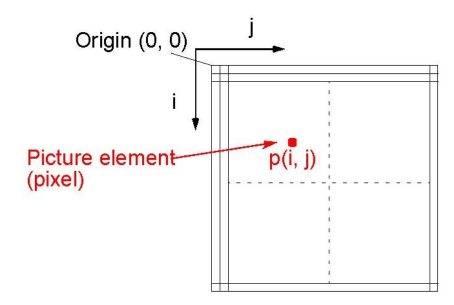
- Operates directly on stored image to improve/enhance it
- Stored image consists of two-dimensional array of pixels (picture elements/"image dots")



#### **Example: Low level image processing**

Many operations are embarassingly parallel

Example: operations that compute each pixel independently (such as shift, scale, rotate)



## Some geometrical operations

### **Shifting**

Object shifted by  $\Delta x$  in the x-dimension and  $\Delta y$  in the y-dimension:

$$\chi' = \chi + \Delta \chi$$

$$y' = y + \Delta y$$

where x and y are the original and x' and y' are the new coordinates.

### **Scaling**

Object scaled by a factor  $S_x$  in x-direction and  $S_y$  in y-direction:

$$\chi' = \chi S_{\chi}$$

$$y' = yS_v$$

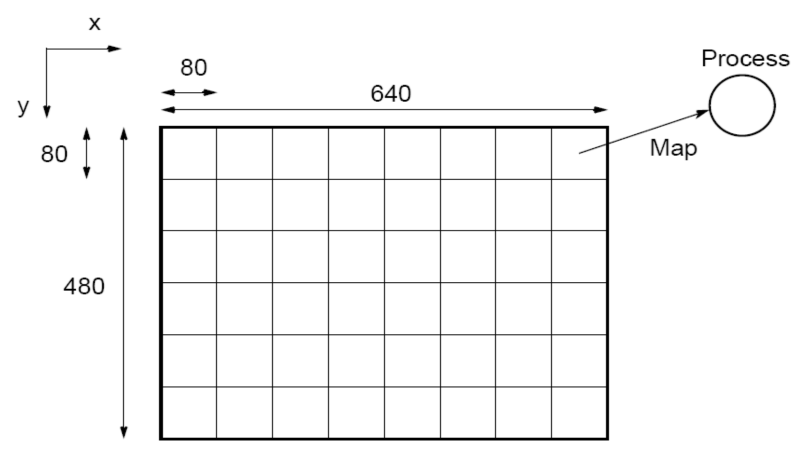
## Some geometrical operations

#### **Rotation**

Object rotated through an angle  $\theta$  about the origin of the coordinate system:

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

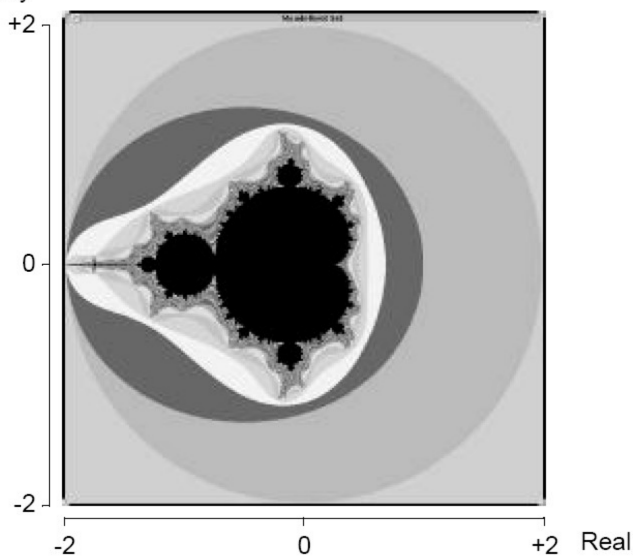
# Partitioning into regions for individual processes



Square region for each process (can also use strips)

## **Example: Mandelbrot set**

⁼lmaginary



## **Mandelbrot Set**

Set of points c in a complex plane that are quasi-stable (will increase and decrease, but not exceed some limit) when computed by iterating the function

$$z_{k+1} = z_k^2 + c$$

where  $z_{k+1}$  is the (k+1)th iteration of the complex number z = a + bi. The initial value for z is zero.

c is a complex number giving position of point in the complex plane.

Iterations continued until magnitude of z is greater than 2 or number of iterations reaches arbitrary limit.

Magnitude of z is the length of the vector given by

$$z_{\text{length}} = \sqrt{a^2 + b^2}$$

# Sequential routine computing value of one point returning number of iterations

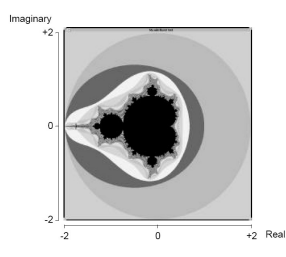
```
struct complex {
   float real;
   float imag;
};
int cal_pixel(complex c)
   int count, max;
   complex z;
   float temp, lengthsq;
   max_iter = 256;
                                  /* 256 colors */
   z.real = 0; z.imag = 0;
                                  /* number of iterations */
   count = 0;
   do {
      temp = z.real * z.real - z.imag * z.imag + c.real;
      z.imag = 2 * z.real * z.imag + c.imag;
      z.real = temp;
      lengthsq = z.real * z.real + z.imag * z.imag;
      count++;
   } while ((lengthsq < 4.0) && (count < max_iter));
   return count;
                    /* color of the pixel */
}
```

## Parallelizing Mandelbrot Set Computation

### Static Task Assignment

Simply divide the region in to fixed number of parts, each computed by a separate processor.

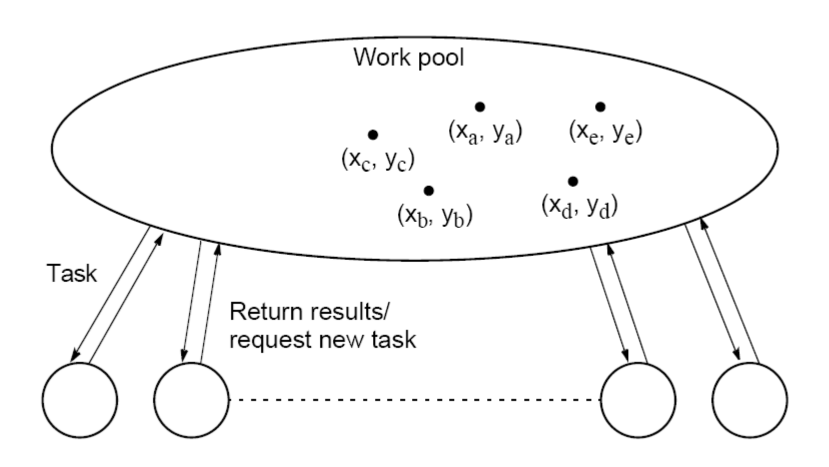
Not very successful because different regions require different numbers of iterations and time.



### **Dynamic Task Assignment**

Have processor request regions after computing previous regions

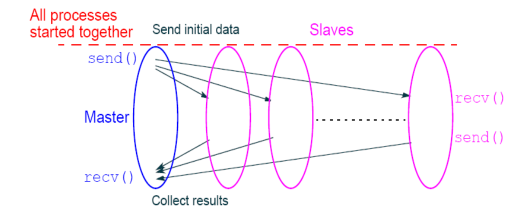
# Dynamic Task Assignment Work Pool/Processor Farms



### Parallel computation

#### Assumption

# processors/processes is given (num\_proc)

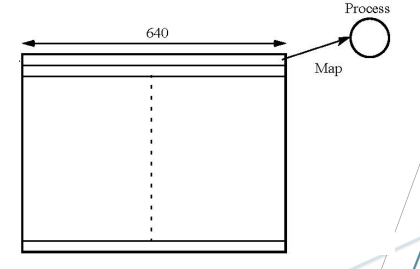


- Each processor computes 1 row at a time
  - Communication time

$$t_{\text{comm}} = t_{\text{startup}} + nt_{\text{data}}$$

The work pool holds row numbers

#### Usual MPI approach



#### Master code

```
/* counter for termination*/
count = 0;
                                                /* row being sent */
row = 0;
for (k = 0; k < num\_proc; k++) {
                                               /* assuming num_proc < disp_height */</pre>
    send(row, Pk, data_tag);
                                               /* send initial row to process */
                                                /* count rows sent */
    count++;
    row++;
} /* next row */
do {
    recv (&slave, &r, &color, P<sub>ANY</sub>, result_tag);
                                                /* reduce count as rows received */
    count - -;
    if (row < disp_height) {</pre>
         send (row, P<sub>slave</sub>, data_tag);
                                               /* send next row */
                                                /* next row */
         row++;
         count++;
    } else
        send (row, P<sub>slave</sub>, terminator_tag); /* terminate */
     display (r, color);
                                               /* display row */
} while (count > 0);
```

#### Slave code

## **Monte Carlo Methods**

Another embarrassingly parallel computation.

Monte Carlo methods use of random selections.

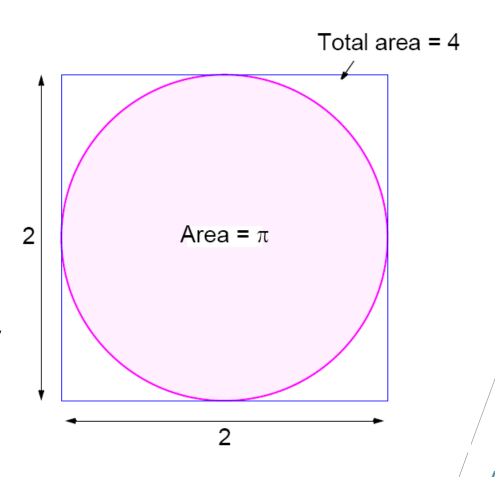
## Example - To calculate $\pi$

Circle formed within a 2 x 2 square. Ratio of area of circle to square given by:

$$\frac{\text{Area of circle}}{\text{Area of square}} = \frac{\pi(1)^2}{2 \times 2} = \frac{\pi}{4}$$

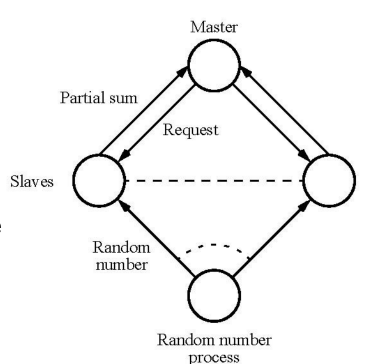
Points within square chosen randomly. Score kept of how many points happen to lie within circle.

Fraction of points within the circle will be  $\frac{\pi}{4}$ , given a sufficient number of randomly selected samples.



#### **Parallel implementation**

- Observation
  - Independent iterations ⇒ embarrassingly parallel problem
- Concern
  - Each computation must use a different random number, and
  - No correlation between the random numbers.
- Approach
  - Have a process responsible for issuing the next random number
  - (not a good one in practice why?)



#### **Parallel implementation**

#### Master

#### **Slaves**

```
sum = 0;
send(P<sub>master</sub>, req_tag);
recv(&xr, &n, P<sub>master</sub>, source_tag);
while (source_tag == compute_tag) {
   for (i = 0; i < n; i++)
        sum = sum + xr[i] * xr[i] - 3 * xr[i];
   send(P<sub>master</sub>, req_tag);
   recv(&xr, &n, P<sub>master</sub>, source_tag);
};
reduce_add(&sum, P<sub>group</sub>); //Collective routine
```

### **Parallel implementation**

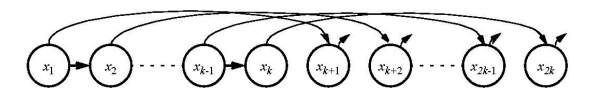
May not scale well

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- Random numbers computed one location
- Random numbers sent to each location
- Still shows principle of handing out tasks and terminating

#### Random number generation

- Goal
  - Create pseudorandom-number sequence  $x_1, x_2, ..., x_i, x_{i+1}, ..., x_n$
- Sequential version
  - Evaluate x<sub>i+1</sub> from a function f of x<sub>i</sub>
  - f must create a large sequence with correct statistical properties
  - Regular form:  $x_{i+1} = (ax_i + c) \mod m$
- Parallel version
  - Observation:  $x_{i+k} = (Ax_i + C) \mod m$ 
    - A =  $a^k \mod m$ ; C =  $c(a^{k-1} + a^{k-2} + ... + a^1 + a^0) \mod m$  (cf. next slide)
    - k: a selected jump constant (usually, #processors)



Available library: Scalable Parallel Random Number Generators (SPRNG)