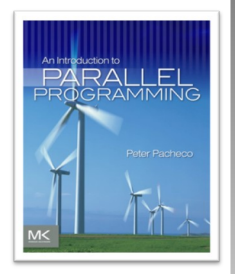


# An Introduction to Parallel Programming Peter Pacheco



#### Chapter 3

Distributed Memory Programming with MPI

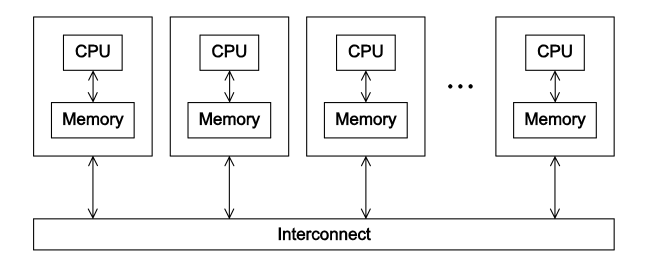


#### Roadmap

- Writing your first MPI program.
- Using the common MPI functions.
- The Trapezoidal Rule in MPI.
- Collective communication.
- MPI derived datatypes.
- Performance evaluation of MPI programs.
- Parallel sorting.
- Safety in MPI programs.

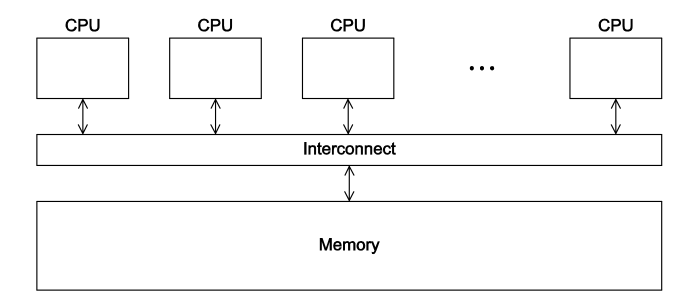


# A distributed memory system





# A shared memory system





#### **Hello World!**

```
#include <stdio.h>
int main(void) {
   printf("hello, world\n");
   return 0;
}
```



(a classic)



### **Identifying MPI processes**

Common practice to identify processes by nonnegative integer ranks.

p processes are numbered 0, 1, 2, .. p-1



### **Our first MPI program**

```
1 #include < stdio.h>
2 #include <string.h> /* For strlen
3 #include <mpi.h> /* For MPI functions, etc */
   const int MAX_STRING = 100;
   int main(void) {
      char
                 greeting[MAX STRING]:
             comm_sz; /* Number of processes */
      int
               my_rank; /* My process rank
10
      int
11
12
      MPI Init (NULL, NULL);
13
      MPI Comm size (MPI COMM WORLD, &comm sz);
      MPI Comm rank (MPI COMM WORLD, &mv rank);
14
15
      if (my_rank != 0) {
16
         sprintf(greeting, "Greetings from process %d of %d!".
17
18
               my_rank, comm_sz);
19
         MPI_Send(greeting, strlen(greeting)+1, MPI_CHAR, 0, 0,
20
               MPI COMM WORLD);
21
      } else {
22
         printf("Greetings from process %d of %d!\n", my_rank, comm_sz);
23
         for (int q = 1; q < comm_sz; q++) {
24
            MPI_Recv(greeting, MAX_STRING, MPI_CHAR, q,
25
               O, MPI COMM WORLD, MPI STATUS IGNORE);
26
            printf("%s\n", greeting);
27
28
29
30
      MPI_Finalize();
31
      return 0;
32
      /* main */
```





### **Compilation**

wrapper script to compile source file mpicc -g -Wall -o mpi\_hello mpi\_hello.c produce create this executable file name debugging (as opposed to default a.out) information turns on all warnings



#### **Execution**

mpiexec -n <number of processes> <executable>

mpiexec -n 1 ./mpi\_hello

run with 1 process

mpiexec -n 4 ./mpi\_hello

run with 4 processes



#### **Execution**

```
mpiexec -n 1 ./mpi_hello
```

Greetings from process 0 of 1!

```
mpiexec -n 4 ./mpi_hello
```

Greetings from process 0 of 4!

Greetings from process 1 of 4!

Greetings from process 2 of 4!

Greetings from process 3 of 4!



#### **MPI Programs**

- Written in C.
  - Has main.
  - Uses stdio.h, string.h, etc.
- Need to add mpi.h header file.
- Identifiers defined by MPI start with "MPI\_".
- First letter following underscore is uppercase.
  - For function names and MPI-defined types.
  - Helps to avoid confusion.



#### **MPI Components**

- MPI\_Init
  - Tells MPI to do all the necessary setup.

- MPI\_Finalize
  - Tells MPI we're done, so clean up anything allocated for this program.

```
int MPI_Finalize(void);
```



#### **Basic Outline**

```
#include <mpi.h>
int main(int argc, char* argv[]) {
   /* No MPI calls before this */
   MPI_Init(&argc, &argv);
   MPI_Finalize();
   /* No MPI calls after this */
   return 0;
```



#### **Communicators**

- A collection of processes that can send messages to each other.
- MPI\_Init defines a communicator that consists of all the processes created when the program is started.
- Called MPI\_COMM\_WORLD.



#### **Communicators**



```
int MPI_Comm_size(
    MPI_Comm comm /* in */,
    int* comm_sz_p /* out */);
```

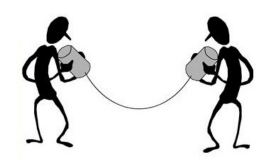
number of processes in the communicator

#### **SPMD**

- Single-Program Multiple-Data
- We compile <u>one</u> program.
- Process 0 does something different.
  - Receives messages and prints them while the other processes do the work.
- The if-else construct makes our program SPMD.



#### Communication

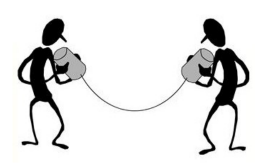


### **Data types**

MPI datatype	C datatype
MPI_CHAR	signed char
MPI_SHORT	signed short int
MPI_INT	signed int
MPI_LONG	signed long int
MPI_LONG_LONG	signed long long int
MPI_UNSIGNED_CHAR	unsigned char
MPI_UNSIGNED_SHORT	unsigned short int
MPI_UNSIGNED	unsigned int
MPI_UNSIGNED_LONG	unsigned long int
MPI_FLOAT	float
MPI_DOUBLE	double
MPI_LONG_DOUBLE	long double
MPI_BYTE	
MPI_PACKED	



#### Communication



### **Message matching**

```
MPI_Send(send_buf_p, send_buf_sz, send_type, dest, send_tag,
        send_comm);
                MPI_Send
                src = q
                                      MPI_Recv
MPI_Recv(recv_buf_p, recv_buf_sz, recv_type, src, recv_tag
         recv_comm, &status);
```



### Receiving messages

- A receiver can get a message without knowing:
  - the amount of data in the message,
  - the sender of the message,
  - or the tag of the message.







### status\_p argument

**MPI\_Status\*** status;

status.MPI\_SOURCE status.MPI TAG

MPI\_SOURCE MPI\_TAG MPI\_ERROR



# How much data am I receiving?

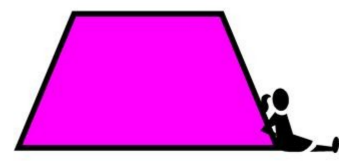




#### Issues with send and receive

- Exact behavior is determined by the MPI implementation.
- MPI\_Send may behave differently with regard to buffer size, cutoffs and blocking.
- MPI\_Recv always blocks until a matching message is received.
- Know your implementation; don't make assumptions!

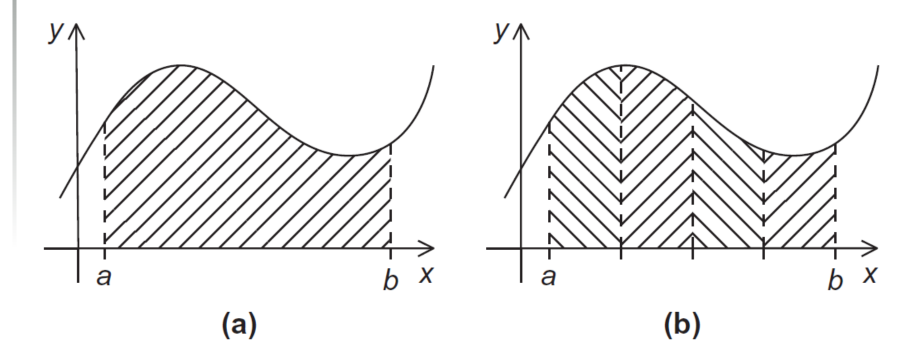




#### TRAPEZOIDAL RULE IN MPI



# **The Trapezoidal Rule**





### The Trapezoidal Rule

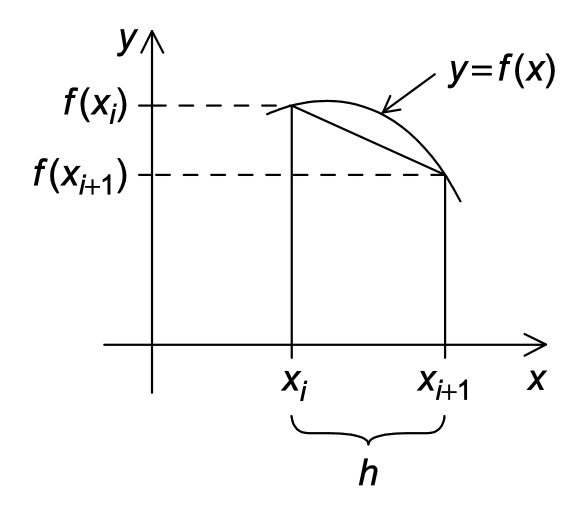
Area of one trapezoid 
$$=\frac{h}{2}[f(x_i)+f(x_{i+1})]$$

$$h = \frac{b-a}{n}$$

$$x_0 = a$$
,  $x_1 = a + h$ ,  $x_2 = a + 2h$ , ...,  $x_{n-1} = a + (n-1)h$ ,  $x_n = b$ 

Sum of trapezoid areas =  $h[f(x_0)/2 + f(x_1) + f(x_2) + \dots + f(x_{n-1}) + f(x_n)/2]$ 

### One trapezoid





### Pseudo-code for a serial program

```
/* Input: a, b, n */
h = (b-a)/n:
approx = (f(a) + f(b))/2.0;
for (i = 1; i \le n-1; i++) {
   x_i = a + i*h:
   approx += f(x_i);
approx = h*approx;
```



### Parallelizing the Trapezoidal Rule

- 1. Partition problem solution into tasks.
- 2. Identify communication channels between tasks.
- 3. Aggregate tasks into composite tasks.
- 4. Map composite tasks to cores.

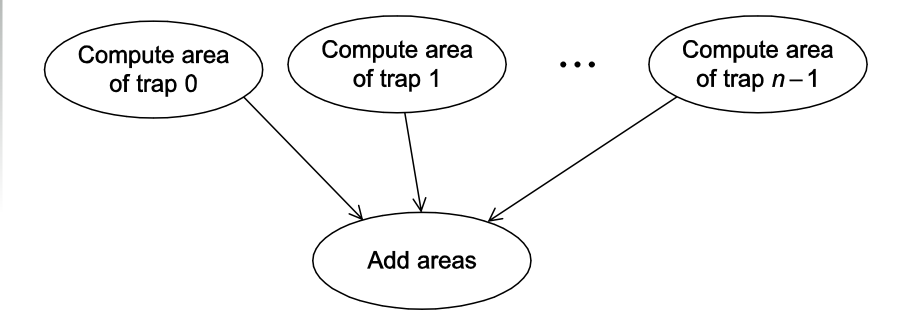


#### Parallel pseudo-code

```
Get a, b, n;
      h = (b-a)/n;
      local n = n/comm sz;
4
      local_a = a + my_rank*local_n*h;
      local_b = local_a + local_n*h;
      local_integral = Trap(local_a, local_b, local_n, h);
6
7
      if (my_rank != 0)
8
         Send local integral to process 0;
9
      else /* my_rank == 0 */
10
         total integral = local integral;
11
         for (proc = 1; proc < comm sz; proc++) {
12
            Receive local integral from proc;
13
            total_integral += local_integral;
14
15
16
      if (my_rank == 0)
17
         print result;
```



# Tasks and communications for Trapezoidal Rule





### First version (1)

```
int main(void) {
      int my rank, comm sz, n = 1024, local n;
3
      double a = 0.0, b = 3.0, h, local a, local b;
      double local int, total int;
5
      int source:
      MPI Init(NULL, NULL);
8
      MPI Comm rank (MPI COMM WORLD, &my rank);
9
      MPI Comm size (MPI COMM WORLD, &comm sz);
10
11
      h = (b-a)/n; /* h is the same for all processes */
12
      local n = n/comm sz; /* So is the number of trapezoids */
13
14
      local a = a + mv rank*local n*h;
15
      local b = local a + local n*h;
16
      local int = Trap(local a, local b, local n, h);
17
18
      if (my rank != 0) {
19
         MPI\_Send(\&local\_int, 1, MPI\_DOUBLE, 0, 0,
20
               MPI COMM WORLD);
```



### First version (2)

```
21
      } else {
22
         total_int = local_int;
23
         for (source = 1; source < comm_sz; source++) {</pre>
24
             MPI Recv(&local int, 1, MPI DOUBLE, source, 0,
25
                   MPI_COMM_WORLD , MPI_STATUS_IGNORE );
26
             total int += local int;
27
28
29
30
      if (mv rank == 0) {
31
          printf("With n = %d trapezoids, our estimate\n", n);
32
         printf("of the integral from %f to %f = %.15e\n",
33
              a, b, total int);
34
35
      MPI Finalize();
36
      return 0:
37
     /* main */
```



### First version (3)

```
double Trap(
         double left endpt /* in */,
         double right_endpt /* in */,
         int trap_count /* in */,
         double base_len /* in */) {
6
      double estimate, x;
      int i:
8
9
      estimate = (f(left\_endpt) + f(right\_endpt))/2.0;
      for (i = 1; i \le trap_count - 1; i++)
10
         x = left_endpt + i*base_len;
11
12
         estimate += f(x);
13
14
      estimate = estimate * base len;
15
16
      return estimate:
17
    /* Trap */
```



### **Dealing with I/O**

```
#include < stdio.h>
#include <mpi.h>
                                  Each process just
                                  prints a message.
int main(void) {
   int my_rank, comm_sz;
   MPI Init(NULL, NULL);
   MPI Comm size (MPI COMM WORLD, &comm sz);
   MPI Comm rank (MPI COMM WORLD, &my rank);
   printf("Proc %d of %d > Does anyone have a toothpick?\n",
         mv rank, comm sz);
   MPI_Finalize();
   return 0;
   /* main */
```



### Running with 6 processes

```
Proc 0 of 6 > Does anyone have a toothpick?

Proc 1 of 6 > Does anyone have a toothpick?

Proc 2 of 6 > Does anyone have a toothpick?

Proc 4 of 6 > Does anyone have a toothpick?

Proc 3 of 6 > Does anyone have a toothpick?

Proc 5 of 6 > Does anyone have a toothpick?
```

unpredictable output





### <u>Input</u>

- Most MPI implementations only allow process 0 in MPI\_COMM\_WORLD access to stdin.
- Process 0 must read the data (scanf) and send to the other processes.

```
MPI_Comm_rank(MPI_COMM_WORLD, &my_rank);
MPI_Comm_size(MPI_COMM_WORLD, &comm_sz);
Get_data(my_rank, comm_sz, &a, &b, &n);
h = (b-a)/n;
. . .
```



## Function for reading user input

```
void Get_input(
     int my rank /*in */.
     int comm sz /*in */,
     double* a_p /* out */,
     double* b_p /* out */,
     int * n p /* out */) {
  int dest:
  if (my rank == 0) {
     printf("Enter a, b, and n\n");
     scanf("%lf %lf %d", a p, b p, n p);
     for (dest = 1; dest < comm sz; dest++) {
        MPI_Send(a_p, 1, MPI_DOUBLE, dest, 0, MPI_COMM_WORLD);
        MPI_Send(b_p, 1, MPI_DOUBLE, dest, 0, MPI_COMM_WORLD);
        MPI Send(n p, 1, MPI INT, dest, 0, MPI COMM WORLD);
  else \{ /* my\_rank != 0 */
     MPI_Recv(a_p, 1, MPI_DOUBLE, 0, 0, MPI COMM WORLD,
           MPI STATUS IGNORE);
     MPI_Recv(b_p, 1, MPI_DOUBLE, 0, 0, MPI_COMM_WORLD,
           MPI STATUS IGNORE);
     MPI_Recv(n_p, 1, MPI_INT, 0, 0, MPI_COMM_WORLD,
           MPI STATUS IGNORE);
  /* Get_input */
```



# COLLECTIVE COMMUNICATION



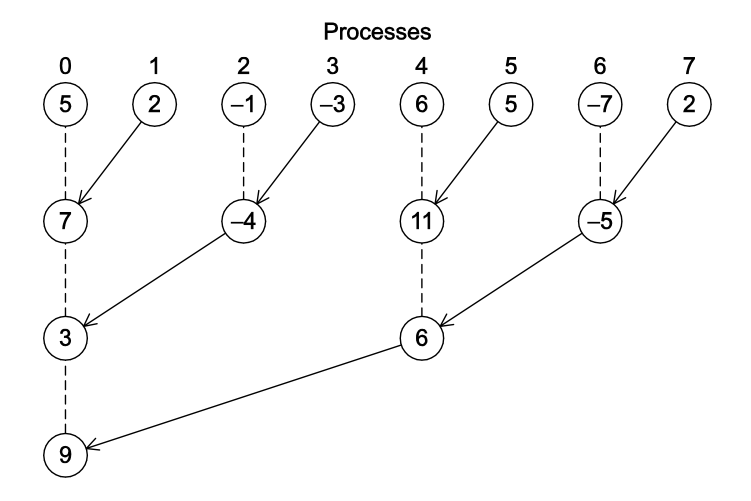


#### **Tree-structured communication**

- 1. In the first phase:
  - (a) Process 1 sends to 0, 3 sends to 2, 5 sends to 4, and 7 sends to 6.
  - (b) Processes 0, 2, 4, and 6 add in the received values.
  - (c) Processes 2 and 6 send their new values to processes 0 and 4, respectively.
  - (d) Processes 0 and 4 add the received values into their new values.
- 2. (a) Process 4 sends its newest value to process 0.
  - (b) Process 0 adds the received value to its newest value.

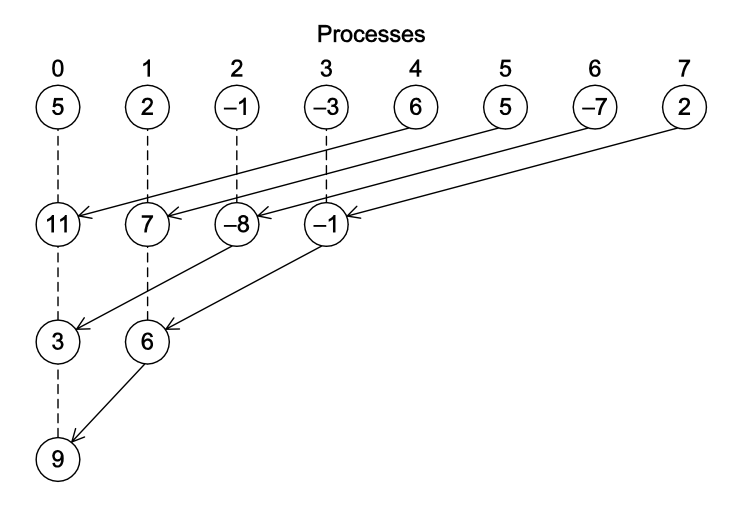


# A tree-structured global sum





# An alternative tree-structured global sum





### MPI\_Reduce

```
\label{eq:mpi_reduce} \begin{split} \texttt{MPI\_Reduce}(\&\texttt{local\_int}\,,\,\,\&\texttt{total\_int}\,,\,\,1\,,\,\,\texttt{MPI\_DOUBLE}\,,\,\,\texttt{MPI\_SUM}\,,\,\,0\,,\\ \texttt{MPI\_COMM\_WORLD}\,); \end{split}
```



# Predefined reduction operators in MPI

Operation Value	Meaning
MPI_MAX	Maximum
MPI_MIN	Minimum
MPI_SUM	Sum
MPI_PROD	Product
MPI_LAND	Logical and
MPI_BAND	Bitwise and
MPI_LOR	Logical or
MPI_BOR	Bitwise or
MPI_LXOR	Logical exclusive or
MPI_BXOR	Bitwise exclusive or
MPI_MAXLOC	Maximum and location of maximum
MPI_MINLOC	Minimum and location of minimum



All the processes in the communicator must call the same collective function.

For example, a program that attempts to match a call to MPI\_Reduce on one process with a call to MPI\_Recv on another process is erroneous, and, in all likelihood, the program will hang or crash.



- The arguments passed by each process to an MPI collective communication must be "compatible."
- For example, if one process passes in 0 as the dest\_process and another passes in 1, then the outcome of a call to MPI\_Reduce is erroneous, and, once again, the program is likely to hang or crash.



- The output\_data\_p argument is only used on dest\_process.
- However, all of the processes still need to pass in an actual argument corresponding to output\_data\_p, even if it's just NULL.



 Point-to-point communications are matched on the basis of tags and communicators.

- Collective communications don't use tags.
- They're matched solely on the basis of the communicator and the order in which they're called.



## Example (1)

Time	Process 0	Process 1	Process 2
0	a = 1; c = 2	a = 1; c = 2	a = 1; c = 2
1	MPI_Reduce(&a, &b,)	MPI_Reduce(&c, &d,)	MPI_Reduce(&a, &b,)
2	MPI_Reduce(&c, &d,)	MPI_Reduce(&a, &b,)	MPI_Reduce(&c, &d,)

### Multiple calls to MPI\_Reduce



# Example (2)

- Suppose that each process calls MPI\_Reduce with operator MPI\_SUM, and destination process 0.
- At first glance, it might seem that after the two calls to MPI\_Reduce, the value of b will be 3, and the value of d will be 6.



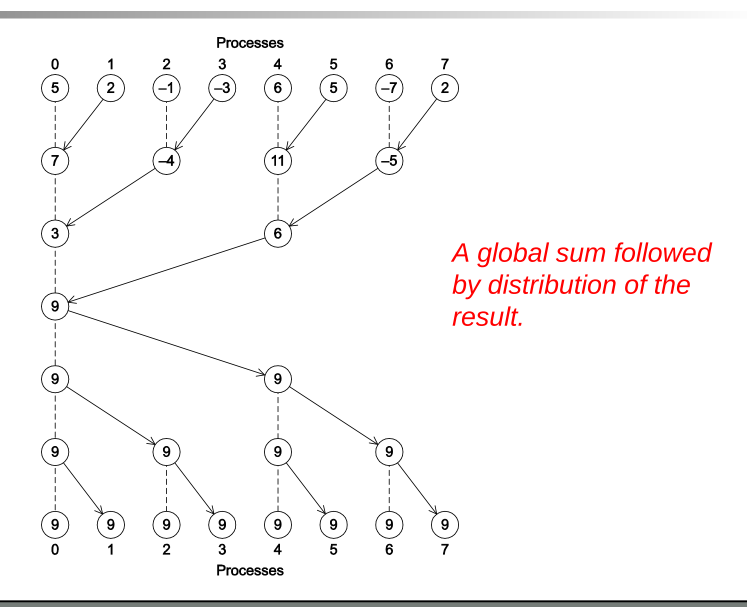
## Example (3)

- However, the names of the memory locations are irrelevant to the matching of the calls to MPI\_Reduce.
- The order of the calls will determine the matching so the value stored in b will be 1+2+1 = 4, and the value stored in d will be 2+1+2 = 5.

### MPI\_Allreduce

Useful in a situation in which all of the processes need the result of a global sum in order to complete some larger computation.

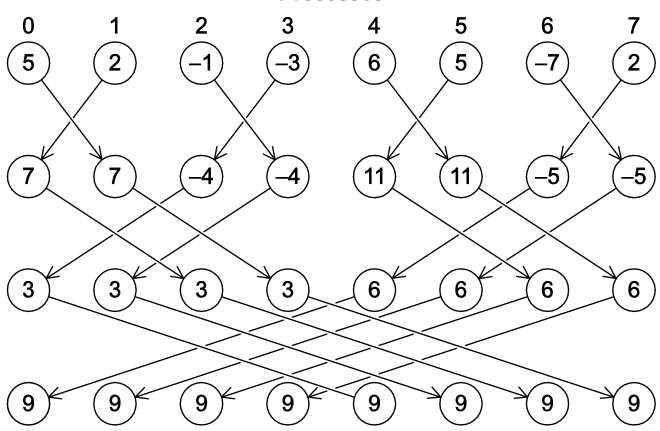








#### **Processes**



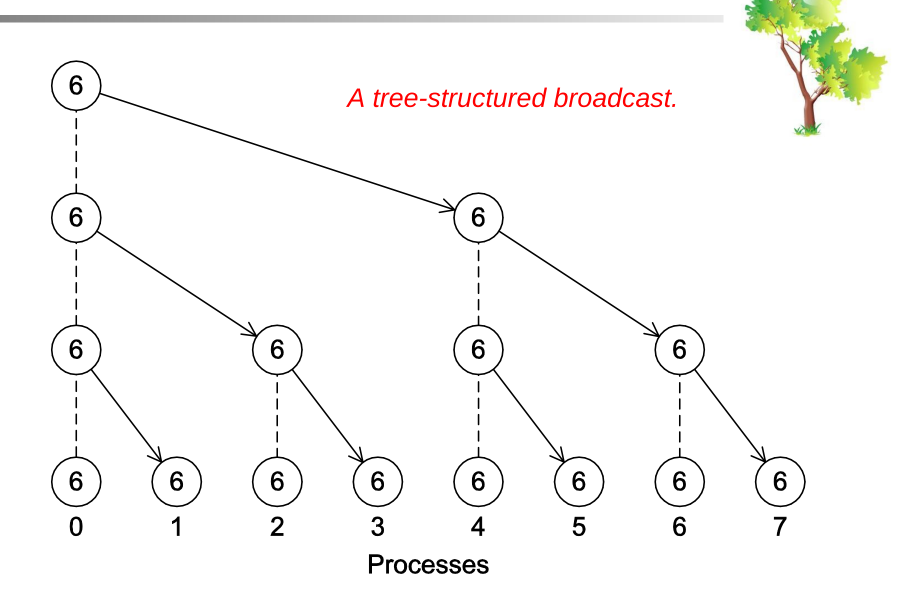
A butterfly-structured global sum.



#### **Broadcast**

 Data belonging to a single process is sent to all of the processes in the communicator.







# A version of Get\_input that uses MPI\_Bcast

```
void Get_input(
     int my_rank /* in */,
     int comm_sz /* in */,
     double * a_p /* out */,
     double* b_p /* out */,
     int * n_p /* out */) {
  if (my_rank == 0) {
     printf("Enter a, b, and n\n");
     scanf("%lf %lf %d", a_p, b_p, n_p);
  MPI_Bcast(a_p, 1, MPI_DOUBLE, 0, MPI_COMM_WORLD);
  MPI_Bcast(b_p, 1, MPI_DOUBLE, 0, MPI_COMM_WORLD);
  MPI_Bcast(n_p, 1, MPI_INT, 0, MPI_COMM_WORLD);
 /* Get_input */
```



#### **Data distributions**

$$\mathbf{x} + \mathbf{y} = (x_0, x_1, \dots, x_{n-1}) + (y_0, y_1, \dots, y_{n-1})$$

$$= (x_0 + y_0, x_1 + y_1, \dots, x_{n-1} + y_{n-1})$$

$$= (z_0, z_1, \dots, z_{n-1})$$

$$= \mathbf{z}$$

Compute a vector sum.



# Serial implementation of vector addition

```
void Vector_sum(double x[], double y[], double z[], int n) {
  int i;

for (i = 0; i < n; i++)
    z[i] = x[i] + y[i];
} /* Vector_sum */</pre>
```



# Different partitions of a 12component vector among 3 processes

		Components										
									В	loc	k-cyc	lic
Process		В	lock			Су	clic				size:	
0	0	1	2	3	0	3	6	9	0	1	6	7
1	4	5	6	7	1	4	7	10	2	3	8	9
2	8	9	10	11	2	5	8	11	4	5	10	11



### Partitioning options

- Block partitioning
  - Assign blocks of consecutive components to each process.
- Cyclic partitioning
  - Assign components in a round robin fashion.
- Block-cyclic partitioning
  - Use a cyclic distribution of blocks of components.



# Parallel implementation of vector addition

```
void Parallel_vector_sum(
    double local_x[] /* in */,
    double local_y[] /* in */,
    double local_z[] /* out */,
    int local_n /* in */) {
    int local_i;

    for (local_i = 0; local_i < local_n; local_i++)
        local_z[local_i] = local_x[local_i] + local_y[local_i];
} /* Parallel_vector_sum */</pre>
```



#### **Scatter**

• MPI\_Scatter can be used in a function that reads in an entire vector on process 0 but only sends the needed components to each of the other processes.

```
int MPI Scatter(
    void*
               send_buf_p /*in */.
    int
               send_count /* in */,
               send_type /*in */,
    MPI_Datatype
    void*
               recv_buf_p /* out */,
    int
               recv_count /* in */,
    MPI_Datatype recv_type /*in */,
    int
               src_proc /* in */,
               comm /* in */):
    MPI Comm
```



### Reading and distributing a vector

```
void Read vector(
     double local_a[] /* out */,
     int local n /* in */,
                     /* in */,
     int
              n
     char vec name [] /* in */,
     int     my_rank     /* in */,
     MPI_Comm comm /*in */) {
  double * a = NULL;
  int i:
  if (mv rank == 0) {
     a = malloc(n*sizeof(double));
     printf("Enter the vector %s\n", vec name);
     for (i = 0; i < n; i++)
        scanf("%lf", &a[i]);
     MPI Scatter(a, local n, MPI DOUBLE, local a, local n, MPI DOUBLE,
           0, comm);
     free(a);
  } else {
     MPI Scatter(a, local n, MPI DOUBLE, local a, local n, MPI DOUBLE,
           0. \text{comm}):
  /* Read_vector */
```



#### **Gather**

Collect all of the components of the vector onto process 0, and then process 0 can process all of the components.



# Print a distributed vector (1)

```
void Print_vector(
    double local_b[] /* in */,
    int local_n /* in */,
               /* in */,
    int
    char title[] /*in */,
       my_rank /* in */,
    int
    MPI Comm comm /*in */) {
  double * b = NULL;
  int i;
```



# Print a distributed vector (2)

```
if (mv rank == 0) {
  b = malloc(n*sizeof(double));
   MPI Gather(local_b, local_n, MPI_DOUBLE, b, local_n, MPI_DOUBLE,
         0. \text{comm}):
   printf("%s\n", title);
   for (i = 0; i < n; i++)
      printf("%f ", b[i]);
  printf("\n");
  free(b);
} else {
  MPI Gather (local b, local n, MPI DOUBLE, b, local n, MPI DOUBLE,
         0. \text{comm});
/* Print_vector */
```



### <u>Allgather</u>

- Concatenates the contents of each process' send\_buf\_p and stores this in each process' recv\_buf\_p.
- As usual, recv\_count is the amount of data being received from each process.



# **Matrix-vector multiplication**

$$A = (a_{ij})$$
 is an  $m \times n$  matrix

 $\mathbf{x}$  is a vector with n components

y = Ax is a vector with m components

$$y_i = a_{i0}x_0 + a_{i1}x_1 + a_{i2}x_2 + \cdots + a_{i,n-1}x_{n-1}$$

i-th component of y

Dot product of the ith row of A with x.



# **Matrix-vector multiplication**

<i>a</i> <sub>00</sub>	<i>a</i> <sub>01</sub>	• • •	$a_{0,n-1}$
$a_{10}$	$a_{11}$	:	$a_{1,n-1}$
:	:		:
$a_{i0}$	$a_{i1}$	• • • •	$a_{i,n-1}$
<i>a</i> <sub>i0</sub> :	<i>a</i> <sub>i1</sub> :	•••	$a_{i,n-1}$ :

$x_0$
$x_1$
:
$x_{n-1}$

	У0
	<i>y</i> <sub>1</sub>
	:
-	$y_i = a_{i0}x_0 + a_{i1}x_1 + \cdots + a_{i,n-1}x_{n-1}$
	:
	$y_{m-1}$

### Multiply a matrix by a vector

```
/* For each row of A */
for (i = 0; i < m; i++) {
    /* Form dot product of ith row with x */
    v[i] = 0.0;

for (j = 0; j < n; j++)
    y[i] += A[i][j]*x[j];
}</pre>
```

Serial pseudo-code



#### **C** style arrays



# Serial matrix-vector multiplication

```
void Mat_vect_mult(
     double A[] /* in */,
     double x[] /* in */,
     double y[] /* out */,
     int m /*in */,
     int n /* in */) {
  int i, j;
  for (i = 0; i < m; i++) {
     y[i] = 0.0;
     for (j = 0; j < n; j++)
        v[i] += A[i*n+j]*x[j];
 /* Mat_vect_mult */
```



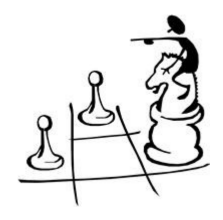
### An MPI matrix-vector multiplication function (1)

```
void Mat_vect_mult(
    double local_A[] /* in */,
    double local_x[] /* in */,
    double local_y[] /* out */,
    int local_m /* in */,
    int n /*in */,
    int local_n /* in */,
    MPI_Comm comm /* in */) {
  double * x;
  int local_i, j;
  int local_ok = 1;
```



### An MPI matrix-vector multiplication function (2)





#### **MPI DERIVED DATATYPES**



#### **Derived datatypes**

- Used to represent any collection of data items in memory by storing both the types of the items and their relative locations in memory.
- The idea is that if a function that sends data knows this information about a collection of data items, it can collect the items from memory before they are sent.
- Similarly, a function that receives data can distribute the items into their correct destinations in memory when they're received.



#### **Derived datatypes**

- Formally, consists of a sequence of basic MPI data types together with a displacement for each of the data types.
- Trapezoidal Rule example:

Variable	Address
a	24
b	40
n	48

 $\{(MPI\_DOUBLE, 0), (MPI\_DOUBLE, 16), (MPI\_INT, 24)\}$ 

#### **MPI\_Type create\_struct**

• Builds a derived datatype that consists of individual elements that have different basic types.



#### MPI\_Get\_address

- Returns the address of the memory location referenced by location\_p.
- The special type MPI\_Aint is an integer type that is big enough to store an address on the system.

```
int MPI_Get_address(
    void* location_p /* in */,
    MPI_Aint* address_p /* out */);
```



#### MPI\_Type\_commit

• Allows the MPI implementation to optimize its internal representation of the datatype for use in communication functions.

```
int MPI_Type_commit(MPI_Datatype* new_mpi_t_p /* in/out */);
```



#### MPI\_Type\_free

When we're finished with our new type, this frees any additional storage used.

```
int MPI_Type_free(MPI_Datatype* old_mpi_t_p /* in/out */);
```



### Get input function with a derived datatype (1)



### Get input function with a derived datatype (2)



### Get input function with a derived datatype (3)

```
void Get_input(int my_rank, int comm_sz, double* a_p, double* b_p,
     int* n_p) {
  MPI Datatype input mpi t;
  Build_mpi_type(a_p, b_p, n_p, &input_mpi_t);
   if (my rank == 0) 
     printf("Enter a, b, and n\n");
     scanf("%lf %lf %d", a_p, b_p, n_p);
  MPI_Bcast(a_p, 1, input_mpi_t, 0, MPI_COMM_WORLD);
  MPI Type free(&input mpi t);
  /* Get_input */
```





#### **PERFORMANCE EVALUATION**



#### **Elapsed parallel time**

 Returns the number of seconds that have elapsed since some time in the past.

```
double MPI_Wtime(void);

double start, finish;
...
start = MPI_Wtime();
/* Code to be timed */
...
finish = MPI_Wtime();
printf("Proc %d > Elapsed time = %e seconds\n"
my_rank, finish-start);
```



#### **Elapsed serial time**

- In this case, you don't need to link in the MPI libraries.
- Returns time in microseconds elapsed from some point in the past.

```
#include "timer.h"
. . .
double now;
. . .
GET_TIME(now);
```





#### **Elapsed serial time**

```
#include "timer.h"
. . .
double start, finish;
. . .
GET_TIME(start);
/* Code to be timed */
. . .
GET_TIME(finish);
printf("Elapsed time = %e seconds\n", finish-start);
```



#### **MPI\_Barrier**

 Ensures that no process will return from calling it until every process in the communicator has started calling it.

```
int MPI Barrier(MPI Comm comm /* in */);
```





#### **MPI\_Barrier**

```
double local_start, local_finish, local_elapsed, elapsed;
MPI Barrier(comm);
local start = MPI Wtime();
/* Code to be timed */
local finish = MPI Wtime();
local_elapsed = local_finish - local_start;
MPI Reduce(&local elapsed, &elapsed, 1, MPI DOUBLE,
  MPI\_MAX, 0, comm);
if (my rank == 0)
  printf("Elapsed time = %e seconds\n", elapsed);
```



### Run-times of serial and parallel matrix-vector multiplication

	Order of Matrix				
comm_sz	1024	2048	4096	8192	16,384
1	4.1	16.0	64.0	270	1100
2	2.3	8.5	33.0	140	560
4	2.0	5.1	18.0	70	280
8	1.7	3.3	9.8	36	140
16	1.7	2.6	5.9	19	71

(Seconds)



#### **Speedup**

$$S(n, p) = \frac{T_{\text{serial}}(n)}{T_{\text{parallel}}(n, p)}$$



#### **Efficiency**

$$E(n,p) = \frac{S(n,p)}{p} = \frac{T_{\text{serial}}(n)}{p \times T_{\text{parallel}}(n,p)}$$



# Speedups of Parallel Matrix-Vector Multiplication

	Order of Matrix				
comm_sz	1024	2048	4096	8192	16,384
1	1.0	1.0	1.0	1.0	1.0
2	1.8	1.9	1.9	1.9	2.0
4	2.1	3.1	3.6	3.9	3.9
8	2.4	4.8	6.5	7.5	7.9
16	2.4	6.2	10.8	14.2	15.5



### Efficiencies of Parallel Matrix-Vector Multiplication

	Order of Matrix				
comm_sz	1024	2048	4096	8192	16,384
1	1.00	1.00	1.00	1.00	1.00
2	0.89	0.94	0.97	0.96	0.98
4	0.51	0.78	0.89	0.96	0.98
8	0.30	0.61	0.82	0.94	0.98
16	0.15	0.39	0.68	0.89	0.97



#### **Scalability**

A program is scalable if the problem size can be increased at a rate so that the efficiency doesn't decrease as the number of processes increase.





#### **Scalability**

Programs that can maintain a constant efficiency without increasing the problem size are sometimes said to be strongly scalable.

Programs that can maintain a constant efficiency if the problem size increases at the same rate as the number of processes are sometimes said to be weakly scalable.



### A PARALLEL SORTING ALGORITHM



#### **Sorting**

- n keys and p = comm sz processes.
- n/p keys assigned to each process.
- No restrictions on which keys are assigned to which processes.
- When the algorithm terminates:
  - The keys assigned to each process should be sorted in (say) increasing order.
  - If 0 ≤ q < r < p, then each key assigned to process q should be less than or equal to every key assigned to process r.



#### Serial bubble sort

```
void Bubble sort(
     int a[] /* in/out */,
     int n /* in */) {
  int list length, i, temp;
  for (list_length = n; list_length \geq 2; list_length--)
     for (i = 0; i < list length -1; i++)
        if (a[i] > a[i+1]) {
           temp = a[i];
           a[i] = a[i+1];
           a[i+1] = temp;
 /* Bubble_sort */
```



#### Odd-even transposition sort

- A sequence of phases.
- Even phases, compare swaps:

$$(a[0], a[1]), (a[2], a[3]), (a[4], a[5]), \dots$$

Odd phases, compare swaps:

$$(a[1], a[2]), (a[3], a[4]), (a[5], a[6]), \dots$$

#### **Example**

Start: 5, 9, 4, 3

Even phase: compare-swap (5,9) and (4,3) getting the list 5, 9, 3, 4

Odd phase: compare-swap (9,3) getting the list 5, 3, 9, 4

Even phase: compare-swap (5,3) and (9,4) getting the list 3, 5, 4, 9

Odd phase: compare-swap (5,4) getting the list 3, 4, 5, 9



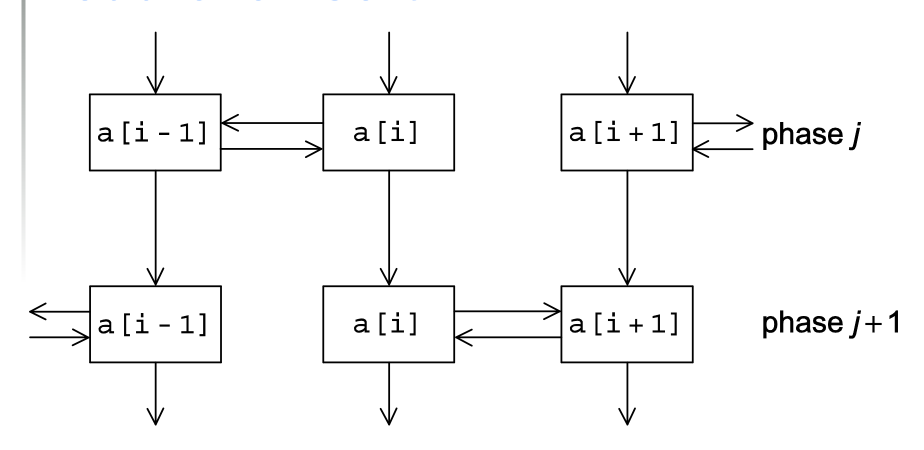
#### Serial odd-even transposition

sort

```
void Odd_even_sort(
      int a [] /* in/out */,
      int n /* in */) {
   int phase, i, temp;
   for (phase = 0; phase < n; phase++)
      if (phase % 2 == 0) { /* Even phase */
         for (i = 1; i < n; i += 2)
            if (a[i-1] > a[i]) {
               temp = a[i];
               a[i] = a[i-1];
              a[i-1] = temp;
      } else { /* Odd phase */
         for (i = 1; i < n-1; i += 2)
            if (a[i] > a[i+1]) {
               temp = a[i];
               a[i] = a[i+1];
               a[i+1] = temp;
  /* Odd_even_sort */
```



### Communications among tasks in odd-even sort



Tasks determining a[i] are labeled with a[i].



### Parallel odd-even transposition sort

	Process				
Time	0	1	2	3	
Start	15, 11, 9, 16	3, 14, 8, 7	4, 6, 12, 10	5, 2, 13, 1	
After Local Sort	9, 11, 15, 16	3, 7, 8, 14	4, 6, 10, 12	1, 2, 5, 13	
After Phase 0	3, 7, 8, 9	11, 14, 15, 16	1, 2, 4, 5	6, 10, 12, 13	
After Phase 1	3, 7, 8, 9	1, 2, 4, 5	11, 14, 15, 16	6, 10, 12, 13	
After Phase 2	1, 2, 3, 4	5, 7, 8, 9	6, 10, 11, 12	13, 14, 15, 16	
After Phase 3	1, 2, 3, 4	5, 6, 7, 8	9, 10, 11, 12	13, 14, 15, 16	



#### Pseudo-code

```
Sort local keys;
for (phase = 0; phase < comm_sz; phase++) {
   partner = Compute_partner(phase, my_rank);
   if (I'm not idle) {
      Send my keys to partner;
      Receive keys from partner;
      if (my_rank < partner)</pre>
         Keep smaller keys;
      else
         Keep larger keys;
```



#### Compute\_partner

```
if (phase % 2 == 0) /* Even phase */
  if (my_rank % 2 != 0) /* Odd rank */
     partner = my_rank - 1;
  else
                            /* Even rank */
     partner = my_rank + 1;
else
                       /* Odd phase */
   if (my_rank % 2 != 0)  /* Odd rank */
     partner = my_rank + 1;
  else
                            /* Even rank */
     partner = my_rank - 1;
if (partner == -1 || partner == comm_sz)
  partner = MPI_PROC_NULL;
```



- The MPI standard allows MPI\_Send to behave in two different ways:
  - it can simply copy the message into an MPI managed buffer and return,
  - or it can block until the matching call to MPI\_Recv starts.



- Many implementations of MPI set a threshold at which the system switches from buffering to blocking.
- Relatively small messages will be buffered by MPI\_Send.
- Larger messages, will cause it to block.



- If the MPI\_Send executed by each process blocks, no process will be able to start executing a call to MPI\_Recv, and the program will hang or deadlock.
- Each process is blocked waiting for an event that will never happen.

(see pseudo-code)



A program that relies on MPI provided buffering is said to be unsafe.

Such a program may run without problems for various sets of input, but it may hang or crash with other sets.



#### MPI\_Ssend

- An alternative to MPI\_Send defined by the MPI standard.
- The extra "s" stands for synchronous and MPI\_Ssend is guaranteed to block until the matching receive starts.



#### Restructuring communication

```
MPI_Send(msg, size, MPI_INT, (my_rank+1) % comm_sz, 0, comm);
MPI Recv(new msg, size, MPI INT, (my rank+comm sz-1) % comm sz,
       0, comm, MPI_STATUS_IGNORE.
if (my rank % 2 == 0) \{
  MPI Send(msg, size, MPI INT, (my rank+1) % comm sz, 0, comm);
  MPI Recv(new msg, size, MPI INT, (my rank+comm sz-1) % comm sz,
         O, comm, MPI STATUS IGNORE.
} else {
  MPI_Recv(new_msg, size, MPI_INT, (my_rank+comm_sz-1) % comm_sz,
         0, comm, MPI STATUS_IGNORE.
  MPI_Send(msg, size, MPI_INT, (my_rank+1) % comm_sz, 0, comm);
```



### **MPI\_Sendrecv**

- An alternative to scheduling the communications ourselves.
- Carries out a blocking send and a receive in a single call.
- The dest and the source can be the same or different.
- Especially useful because MPI schedules the communications so that the program won't hang or crash.

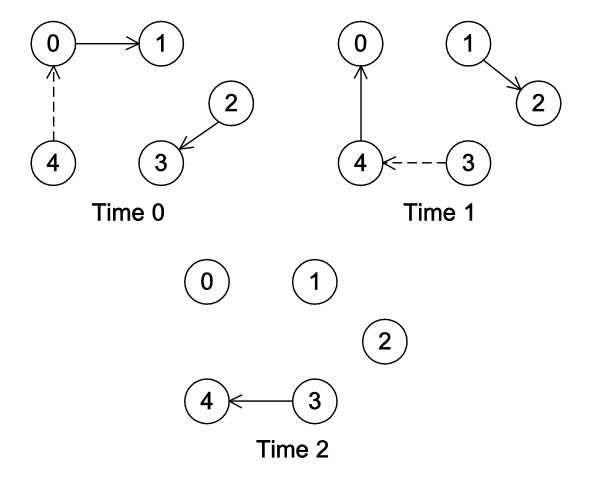


#### MPI\_Sendrecv

```
int MPI_Sendrecv(
    void*
                send_buf_p /*in */,
                send_buf_size /*in */,
    int
    MPI_Datatype send_buf_type /*in */,
                             /* in */,
     int
                dest
                send_tag /*in */,
     int
                recv_buf_p /* out */,
     void*
                recv_buf_size /* in */,
     int
     MPI_Datatype recv_buf_type /*in */,
     int
                             /* in */,
                source
                int
     MPI Comm communicator /*in */,
     MPI_Status* status_p /*in */);
```



# Safe communication with five processes





#### Parallel odd-even transposition sort

```
void Merge_low(
     int my_keys[], /* in/out */
     int recv_keys[], /* in */
     int temp_keys[], /* scratch */
     int local_n /* = n/p, in */) {
  int mi, ri, ti;
  m i = r i = t i = 0;
  while (t_i < local_n) {
     if (my_keys[m_i] <= recv_keys[r_i]) {</pre>
        temp keys[t i] = my keys[m i];
       t i++; m i++;
     } else {
        temp keys[t i] = recv keys[r i];
       t_i++; r_i++;
  for (m_i = 0; m_i < local_n; m_i++)
     my keys[m_i] = temp_keys[m_i];
} /* Merge_low */
```



## Run-times of parallel odd-even sort

	Number of Keys (in thousands)				
Processes	200	400	800	1600	3200
1	88	190	390	830	1800
2	43	91	190	410	860
4	22	46	96	200	430
8	12	24	51	110	220
16	7.5	14	29	60	130

(times are in milliseconds)



## **Concluding Remarks (1)**

- MPI or the Message-Passing Interface is a library of functions that can be called from C, C++, or Fortran programs.
- A communicator is a collection of processes that can send messages to each other.
- Many parallel programs use the singleprogram multiple data or SPMD approach.



## **Concluding Remarks (2)**

- Most serial programs are deterministic: if we run the same program with the same input we'll get the same output.
- Parallel programs often don't possess this property.
- Collective communications involve all the processes in a communicator.



## **Concluding Remarks (3)**

- When we time parallel programs, we're usually interested in elapsed time or "wall clock time".
- Speedup is the ratio of the serial run-time to the parallel run-time.
- Efficiency is the speedup divided by the number of parallel processes.



## **Concluding Remarks (4)**

- If it's possible to increase the problem size (n) so that the efficiency doesn't decrease as p is increased, a parallel program is said to be scalable.
- An MPI program is unsafe if its correct behavior depends on the fact that MPI\_Send is buffering its input.

