

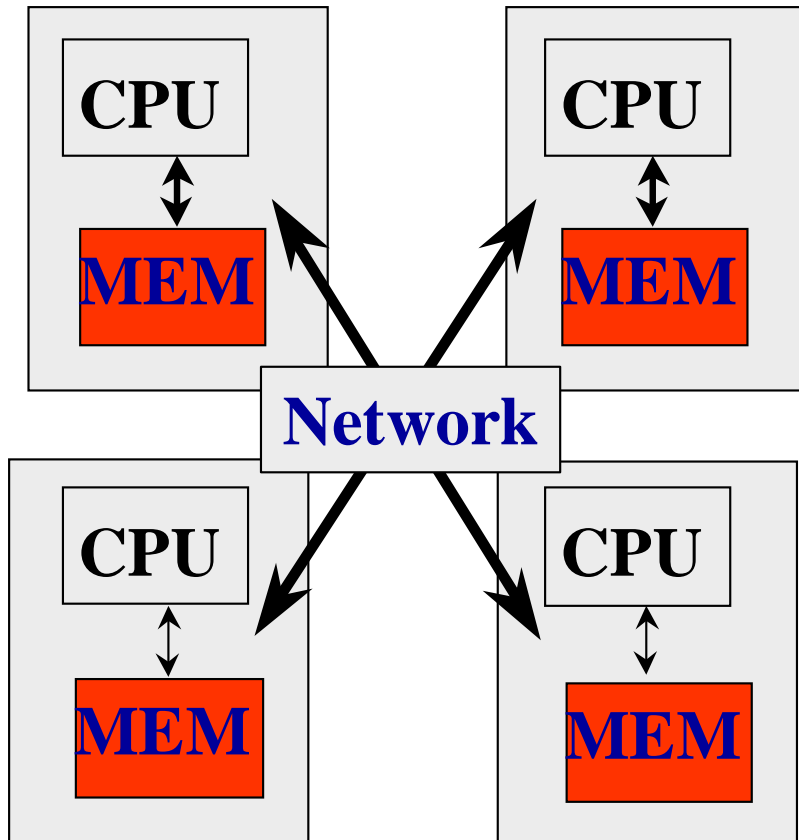
Programming for High Performance Computing

Programming Environment

Dec 5, 2013

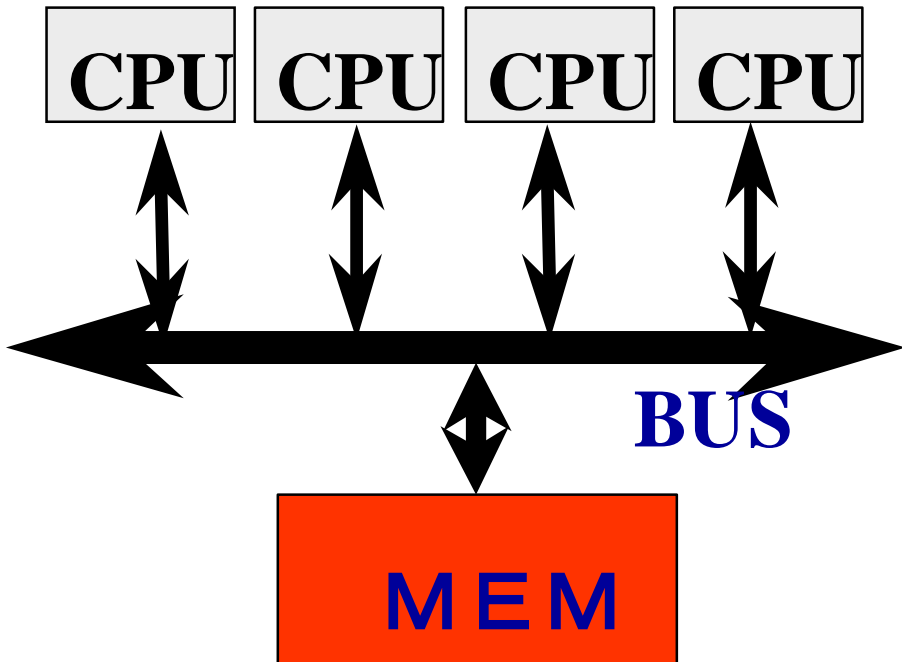
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Distributed Memory Machine



- ◆ Parallel machine that consists computers (CPU and memory) connected by a network
- ◆ Parallel program is executed on each machine, communicating data (message) by the network
- ◆ MPP (Massively Parallel Processor)
 - ◆ Using proprietary technology
- ◆ PC Cluster
 - ◆ Using commodity technology

Shared Memory Machine



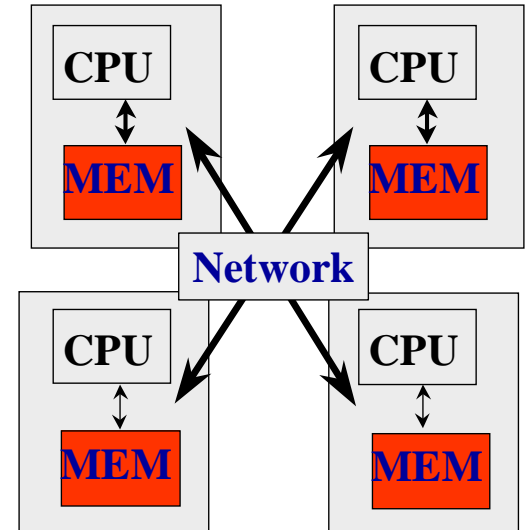
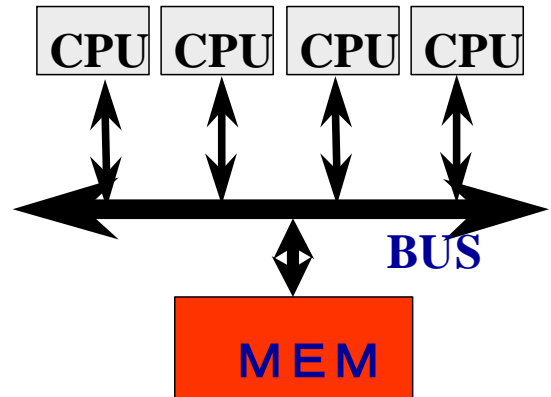
- ◆ Parallel machine such that CPUs access the same memory
- ◆ Parallel program is executed by accessing the data on the shared memory to communicate each other
- ◆ Large scale server, multicore CPU
- ◆ UMA (uniform memory access) and NUMA (non-uniform memory access)

Benefit of parallel processing

- Increase computational power
 - More than one CPU
- Increase memory BW
 - Each CPU can access each memory
- Increase I/O BW
 - Each CPU can access each storage in parallel
- Increase cache/memory size
 - Even though the data size does not fit in the cache size in a single processor, the cache memory can be efficiently used by dividing the data in the problem
- Cost effective
 - When using microprocessors



PC Cluster Technology



Parallel Programming (1)

- Message passing programming
 - For Distributed memory machines
 - Can be used for shared memory machines
 - Complicated and rather difficult
 - Data transfer should be programmed
 - Scalable in terms of # processors
- Shared memory programming
 - For shared memory machines
 - Can be used for distributed shared memory (DSM) system on distributed memory machines
 - Easy to program (incremental from serial program)
 - Data transfer is taken care by the system
 - Often not scalable in terms of # processors

Parallel Programming (2)

- Message passing programming
 - MPI, PVM
- Shared memory programming
 - Multithread programming
 - Pthread, Solaris thread, NT thread
 - OpenMP
 - Annotation for parallel execution
 - Thread control, ... for shared memory machines
 - High Performance Fortran (HPF)
 - Annotation for parallel execution
 - Parallel construct
 - Distribution of array, ... for distributed memory machines
- Automatic parallelization
 - Parallelize serial programs by compiler
 - Parallelization analysis by compiler is limited
 - Sometimes hint is required for parallelization by annotation
- Fancy parallel programming languages

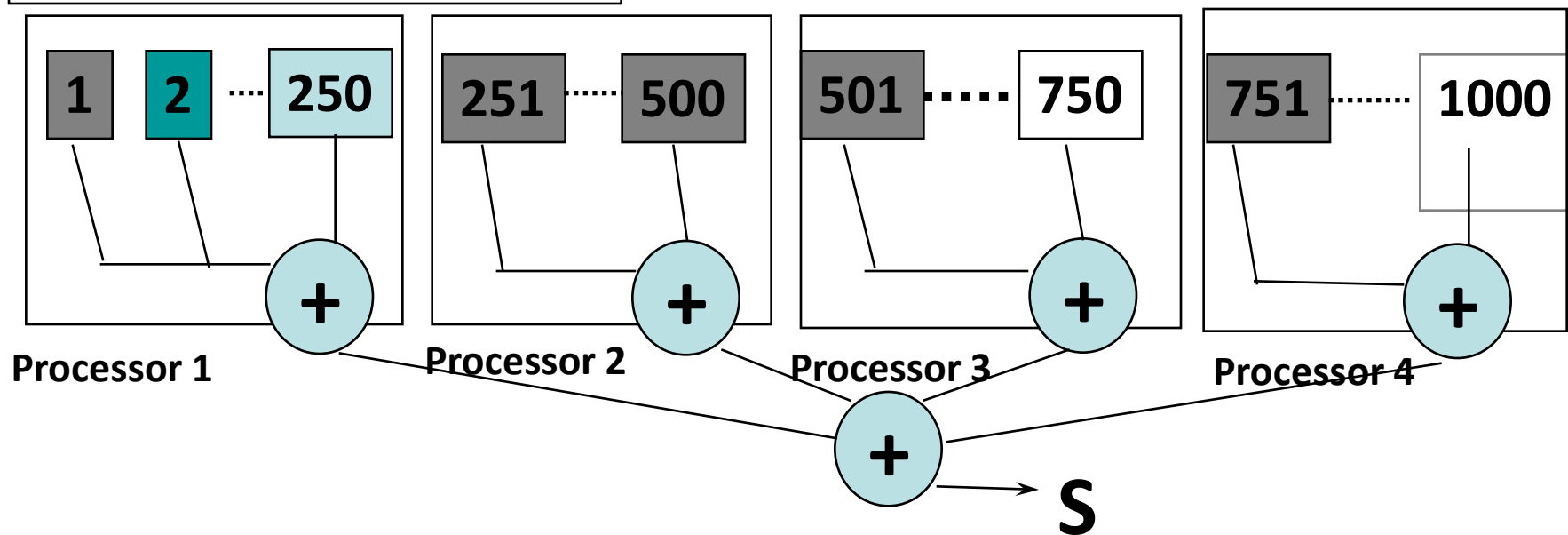
Example of parallel computing

Serial computation

```
for (i = 0; i < 1000; i++)  
    S += A[i]
```



Parallel computation



Multithread Programming

- Thread creation

Pthread, Solaris thread

```
for (t = 1; t < n_thd; t++)  
    pthread_create(&th[t], NULL, thd_main, t)  
thd_main(0);  
for (t = 1; t < n_thd; t++)  
    pthread_join(th[t]);
```

- Divide the summation loop
- Atomic addition

```
double s; /* global */  
pthread_mutex_t mutex  
    = PTHREAD_MUTEX_INITIALIZER;  
int n_thd; /* number of threads */  
void *thd_main(void *a)  
{ int c, b, e, i, id = a; double ss;  
    c = 1000 / n_thd;  
    b = c * id;  
    e = b + c;  
    ss = 0.0;  
    for (i = b; i < e; i++) ss += a[i];  
    pthread_mutex_lock(&mutex);  
    s += ss;  
    pthread_mutex_unlock(&mutex);  
}
```

Programming in OpenMP

Just it is!

```
#pragma omp parallel for reduction(+:s)
for (i = 0; i < 1000; i++) s += a[i];
```

About OpenMP

- Parallel programming model for shared memory machine
 - Extend the base languages (Fortran/C/C++) by using directive
- Specification discussed internationally (mainly US compiler vendors)
 - Oct. 1997 Fortran ver.1.0 API
 - Oct. 1998 C/C++ ver.1.0 API
 - July 2011 OpenMP 3.1
- URL
 - <http://www.openmp.org/>

Programming in MPI

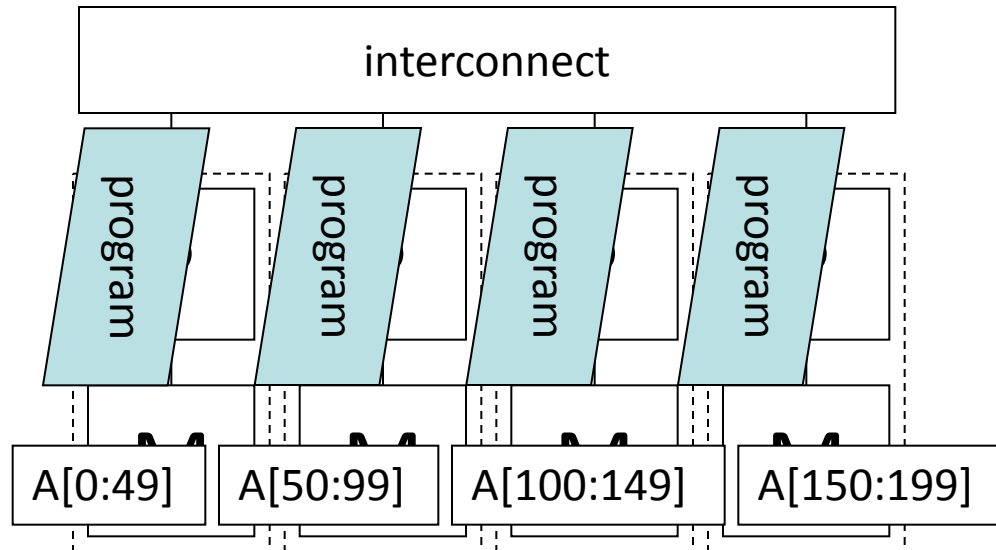
- MPI (Message Passing Interface)
- Standard programming model for distributed memory machines
 - Required for more than 100 nodes
 - Complicated but required for performance
 - Like assembly language
- Message passing (point-to-point communication)
 - Send/Receive
- Collective communication
 - Summation, ...

MPI – The Message Passing Interface

- Standard of message passing interface
- MPI-1.0 released in 1994
 - Portable parallel library, application
 - 8 communication modes, collective communication, communication domain, process topology
 - Defined more than 100 interfaces
 - C, C++, Fortran
 - Specification <http://www.mpi-forum.org/>
 - MPI-3.0 released in September, 2012
 - Japanese translation <http://phase.hpcc.jp/phase/mpi-j/ml/>

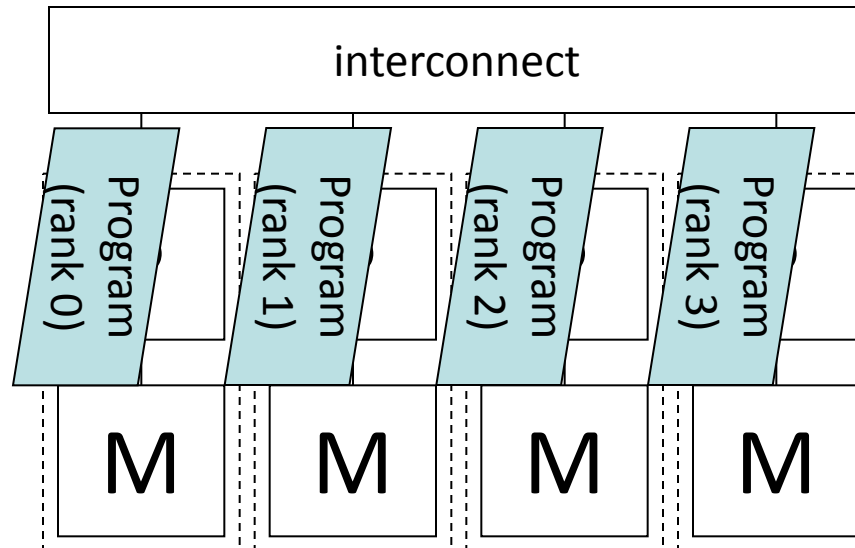
SPMD – Single Program, Multiple Data

- Parallel execution of the same single program independently (cf. SIMD)
- The same program but processes different data
- Parallel program is interacted with each other by message exchange

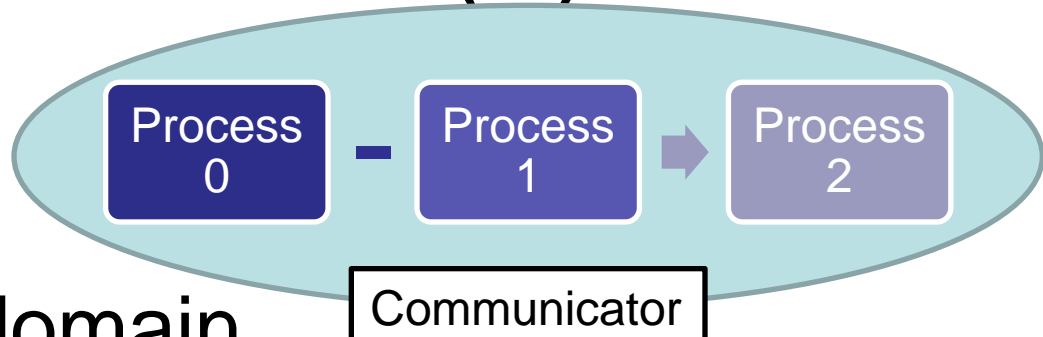


MPI execution model

- Execute the same program on each processor
 - Execution is not synchronous (if no communication happens)
- Each process has its own process rank
- Each process is communicated in MPI



Communicator (1)



- Communication domain
 - Set of processes
 - # processes, process rank
 - Process topology
 - 1D ring, 2D mesh, torus, graph
- **MPI_COMM_WORLD**
 - Initial communicator including all processes

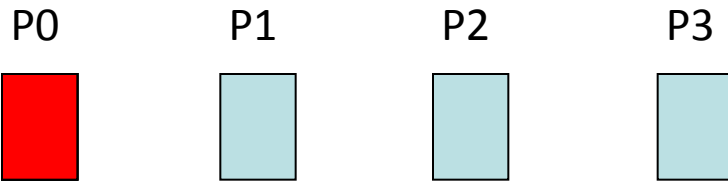
Communicator (2)

- “Scope” of collective communication (communication domain)
- Can divide set of processes
 - Two thirds of processes compute weather forecast, the rest one third compute the initial condition of the next iteration
- Intra-communicator and inter-communicator

Collective communication

- Message exchange among **all processes** specified by a communicator
- Barrier synchronization (no data transfer)
- Global communication
 - Broadcast, gather, scatter, gather to all, all-to-all scatter/gather
- Global reduction
 - Reduction (sum, maximum, logical and, ...), scan (prefix computation)

Global communication



- broadcast

- Transfer $A[*]$ of the root process to all other processes



- gather

- Gather sub arrays distributed among processes into a root process
 - Allgather gather sub arrays into all processes



- scatter

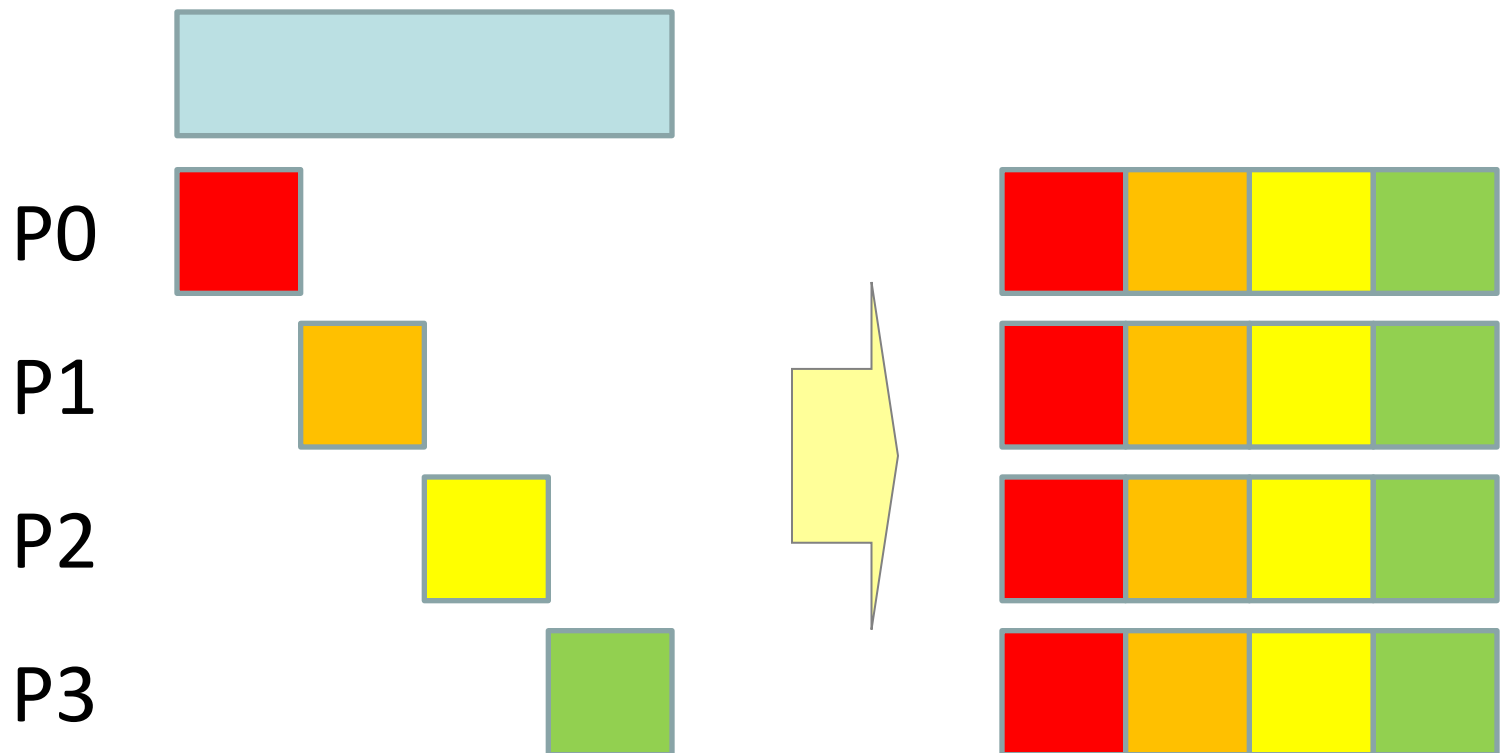
- Scatter $A[*]$ of the root process to all processes

- Alltoall

- Scatter/gather data from all processes to all processes
 - Distributed matrix transpose $A[:][*] \rightarrow A^T[:,[*]]$ (: means this dimension is distributed)

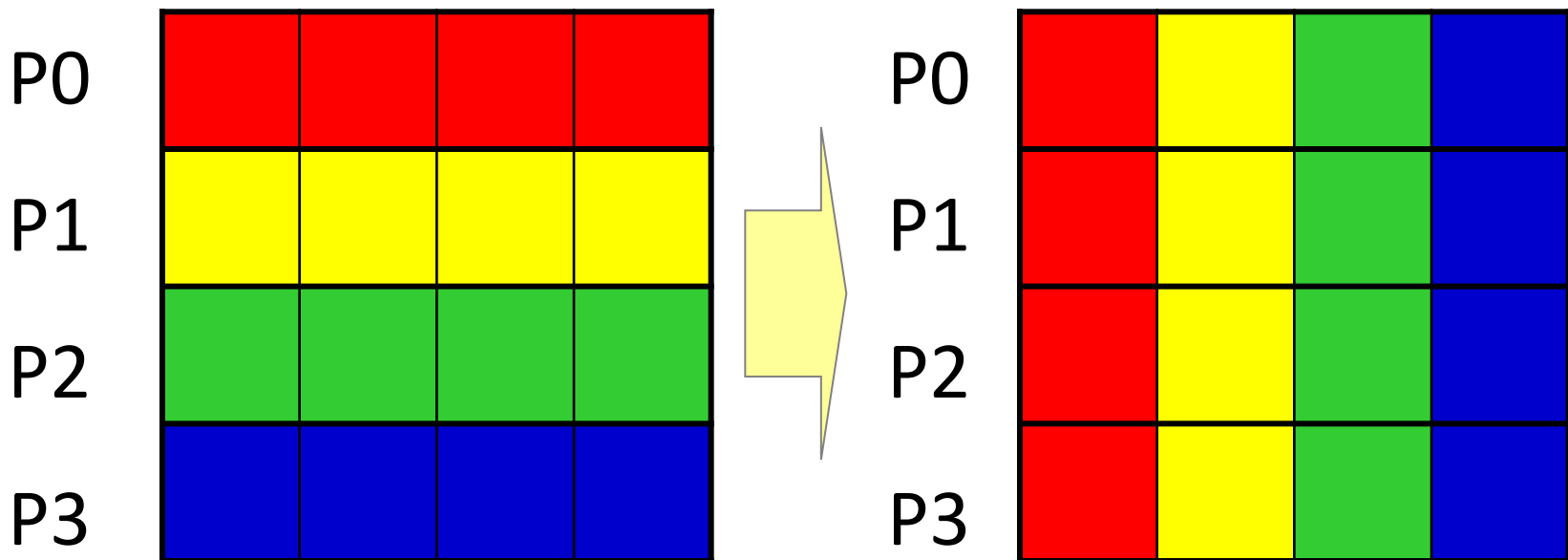
allgather

- Gather sub array of each process, and broadcast the whole array to all processes



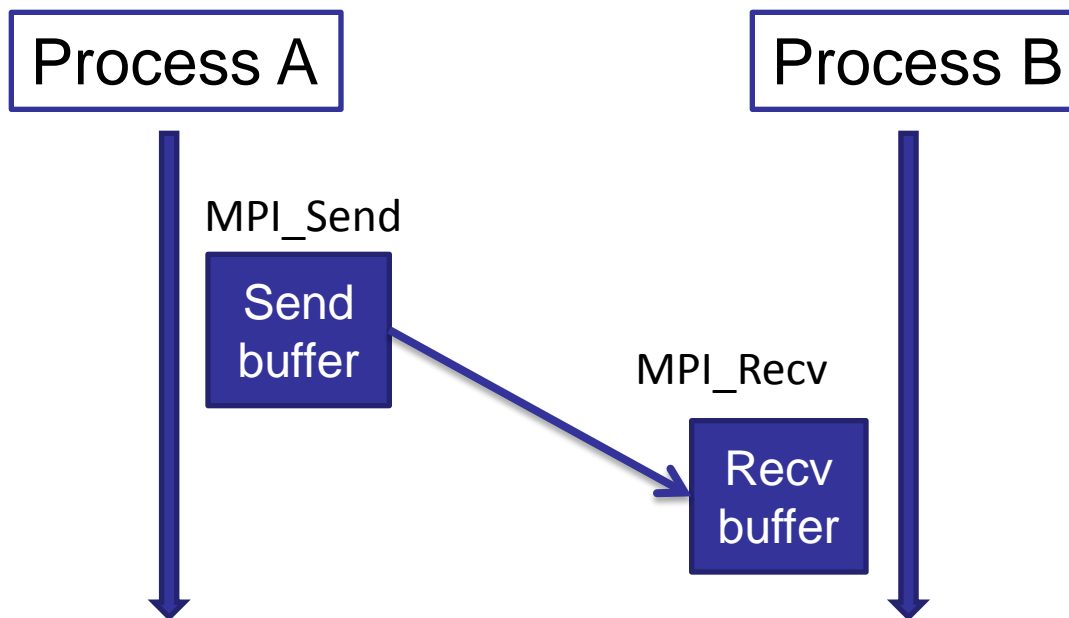
alltoall

- Matrix transformation of (row-wise) distributed 2D array



Point-to-point communication (1)

- Data transfer among two process pair
 - Process A sends a data to process B (send)
 - Process B receives the data (from the process A) (recv)



Point-to-point communication (2)

- Data is typed
 - Basic data type
 - MPI_INT, MPI_DOUBLE, MPI_BYTE, ...
 - Array, structure, vector, user-defined data type
- Send and the corresponding receive are specified by Communicator, message tag, process rank of source and destination

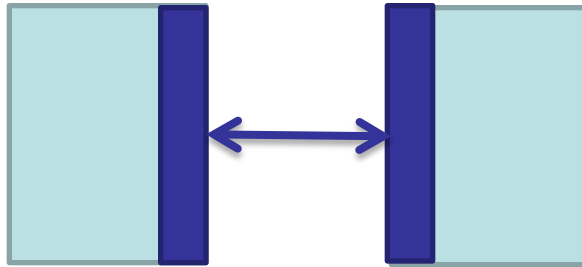
Point-to-point communication (3)

- Semantics of blocking communication
 - Send call returns when the send buffer can be reused
 - Receive call returns when the receive buffer is available
- When MPI_Send(A, . . .) returns, A can be safely modified
 - It may be that A is just copied into the communication buffer of the sender
 - It does not mean message transfer completion

Non-blocking point-to-point communication

- Nonblocking communication
 - post-send, complete-send
 - post-receive, complete-receive
- Post-{send,recv} initiates the send/receive operations
- Complete-{send,recv} waits for the completion
- It enables the overlap of computation and communication to improve performance
 - Multithread programming also enables the overlapping, but nonblocking communication often more efficient

Message exchange



- Blocking P2P

```
...  
MPI_Send(destination, data)  
MPI_Recv(sender, data)  
...
```

- It can be executed only if MPI_Send is buffered
 - Unless, the **dead lock** occurs
- Or, use MPI_Sendrecv

- Nonblocking P2P

```
...  
MPI_Isend(destination, data, &req[0])  
MPI_Recv(sender, &req[1])  
MPI_Waitall(2, req, status)  
...
```

- Safely executed
- Portable

Sample program (1): hostname

```
#include <stdio.h>
#include <mpi.h>

int
main(int argc, char *argv[])
{
    int rank, len;
    char name[MPI_MAX_PROCESSOR_NAME];

    MPI_Init(&argc, &argv);
    MPI_Comm_rank(MPI_COMM_WORLD, &rank);
    MPI_Get_processor_name(name, &len);
    printf("%03d %s\n", rank, name);
    MPI_Finalize();
    return (0);
}
```

Explanation

- Include `mpi.h` to use MPI
- Each process executes the main function
- SPMD (single program, multiple data)
 - A single program is executed on each node
 - Each program accesses different data (ie. data in their own running process)
- Initialize the MPI process
 - `MPI_Init`

Explanation (continued)

- Obtain the process rank
 - `MPI_Comm_rank(MPI_COMM_WORLD, &rank);`
 - Obtain the self rank in the communicator `MPI_COMM_WORLD`
 - Communicator is an opaque object. The information can be access by API
- Obtain the node name
 - `MPI_Get_processor_name(name, &len);`
- All processes should finalize the MPI process
`MPI_Finalize();`

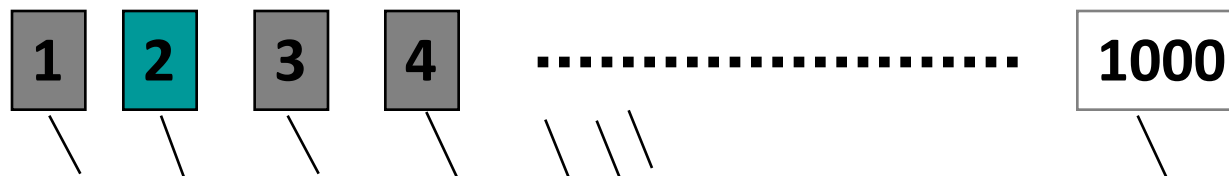
Operation for communicator

- int **MPI_Comm_size**(MPI_Comm *comm*, int **size*);
 - Returns the total number of processes *size* in the communicator *comm*
- int **MPI_Comm_rank**(MPI_Comm *comm*, int **rank*);
 - Returns the process rank *rank* in the communicator *comm*

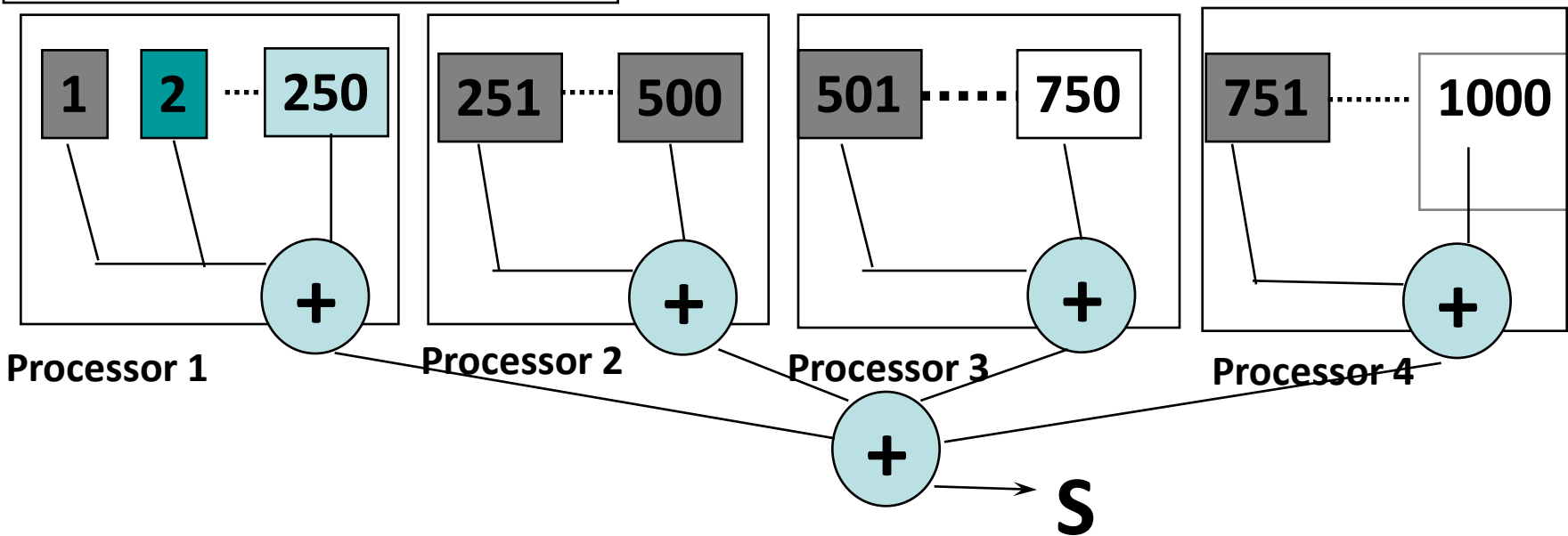
Sample program (2): summation

Serial computation

```
for (i = 0; i < 1000; i++)  
    S += A[i]
```



Parallel computation



```
#include <mpi.h>

double SubA[250];      // subarray of A

int main(int argc, char *argv[])
{
    double sum, mysum;

MPI_Init(&argc,&argv);
    mysum = 0.0;
    for (i = 0; i < 250; i++)
        mysum += SubA[i];
MPI_Reduce(&mysum, &sum, 1, MPI_DOUBLE,
            MPI_SUM, 0, MPI_COMM_WORLD);
MPI_Finalize();
    return (0);
}
```

Explanation

- Allocate a different part of **sub-array** of A is in each process
- Computation and communication
 - Each process computes a partial sum, and communicates with all processes to sum it up by collective communication
`MPI_Reduce(&mysum, &sum, 1, MPI_DOUBLE,
MPI_SUM, 0, MPI_COMM_WORLD);`
 - Combines *mysum* (an array of MPI_DOUBLE with size 1) using MPI_SUM, and returns the combined value *sum* of the root process (rank 0)

Sample program (3): Cpi

- Calculate the PI by the integral calculus
- Test program of MPICH
 - Riemann Sum
 - Broadcast n (number of divided parts)
 - Reduce the partial sum
 - The partial sum is computed in cyclic manner

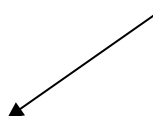
$$\pi = \int_0^1 \frac{4}{1+t^2} dt$$

...

MPI_Bcast(&n, 1, **MPI_INT**, 0, **MPI_COMM_WORLD**);

```
h = 1.0 / n;
sum = 0.0;
for (i = myid + 1; i <= n; i += numprocs){
    x = h * (i - 0.5);
    sum += f(x);
}
mypi = h * sum;
```

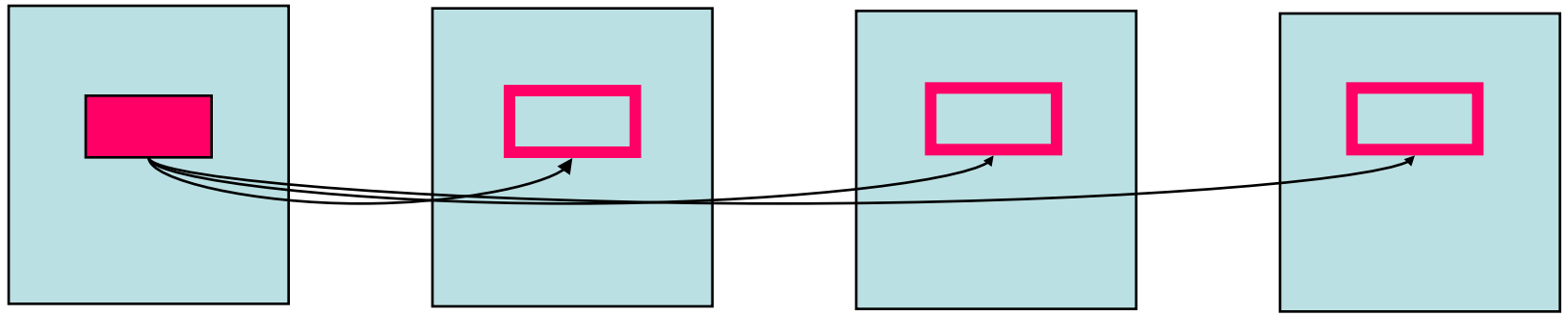
for (i = 1; i <= n; i++)



MPI_Reduce(&mypi, &pi, 1, **MPI_DOUBLE**,
MPI_SUM, 0, **MPI_COMM_WORLD**);

broadcast

```
MPI_Bcast(
    void    *data_buffer, // address of source and destination buffer of data
    int      count,      // data counts
    MPI_Datatype data_type, // data type
    int      source,      // source process rank
    MPI_Comm  communicator // communicator
);
```

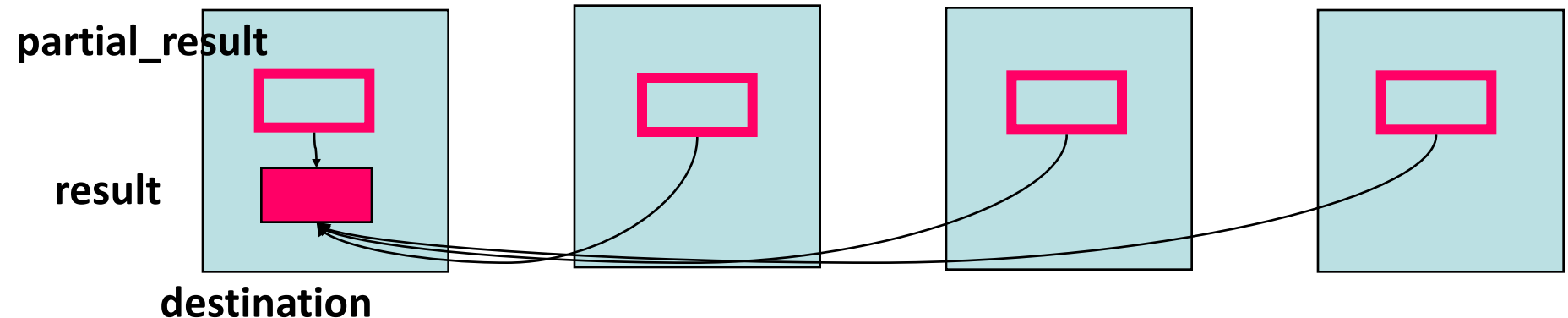


source

It should be executed on all processes in the communicator

Reduction

```
MPI_Reduce(
    void *partial_result,           // address of input data
    void *result,                  // address of output data
    int count,                     // data count
    MPI_Datatype data_type,        // data type
    MPI_Op operator,              // reduce operation
    int destination,               // destination process rank
    MPI_Comm communicator // communicator
);
```



It should be executed on all processes in the communicator

MPI_Allreduce returns the result on all processes

```
/* cpi mpi version */
#include <stdlib.h>
#include <stdio.h>
#include <math.h>
#include <mpi.h>

double
f(double a)
{
    return (4.0 / (1.0 + a * a));
}

int
main(int argc, char *argv[])
{
    int n = 0, myid, numprocs, i;
    double PI25DT = 3.141592653589793238462643;
    double mypi, pi, h, sum, x;
    double startwtime = 0.0, endwtime;
    int namelen;
    char processor_name[MPI_MAX_PROCESSOR_NAME];
```

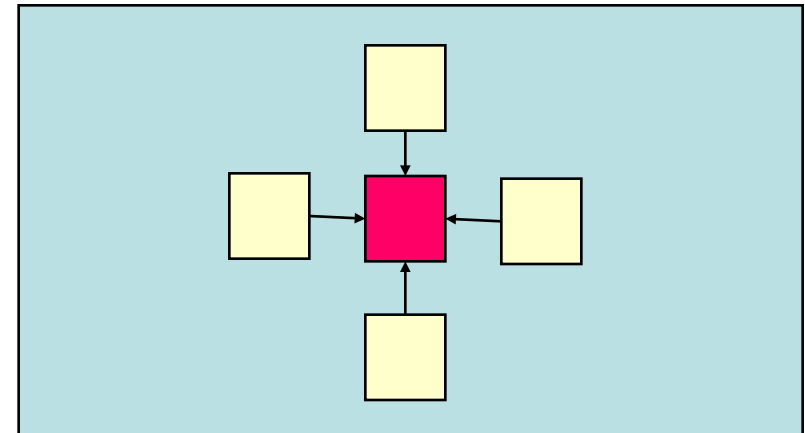
```
MPI_Init(&argc, &argv);
MPI_Comm_size(MPI_COMM_WORLD, &numprocs);
MPI_Comm_rank(MPI_COMM_WORLD, &myid);
MPI_Get_processor_name(processor_name, &namelen);
fprintf(stderr, "Process %d on %s\n", myid, processor_name);

if (argc > 1)
    n = atoi(argv[1]);
startwtime = MPI_Wtime();
/* broadcast 'n' */
MPI_Bcast(&n, 1, MPI_INT, 0, MPI_COMM_WORLD);
if (n <= 0) {
    fprintf(stderr, "usage: %s #partition\n", *argv);
    MPI_Abort(MPI_COMM_WORLD, 1);
}
```

```
/* calculate each part of pi */
h = 1.0 / n;
sum = 0.0;
for (i = myid + 1; i <= n; i += numprocs){
    x = h * (i - 0.5);
    sum += f(x);
}
mypi = h * sum;
/* sum up each part of pi */
MPI_Reduce(&mypi, &pi, 1, MPI_DOUBLE, MPI_SUM, 0, MPI_COMM_WORLD);
if (myid == 0) {
    printf("pi is approximately %.16f, Error is %.16f\n",
          pi, fabs(pi - PI25DT));
    endwtime = MPI_Wtime();
    printf("wall clock time = %f\n",
          endwtime - startwtime);
}
MPI_Finalize();
return (0);
}
```

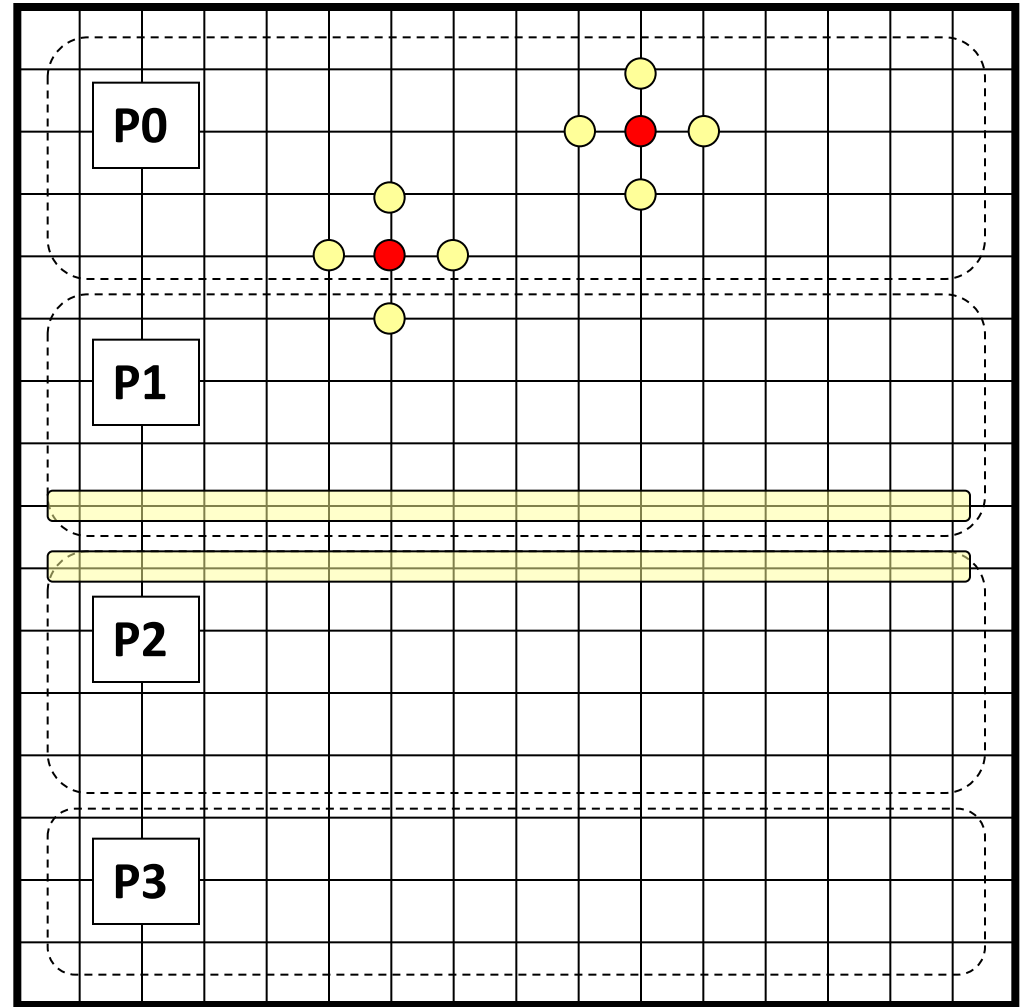
Sample program (4): laplace

- Explicit solution of Laplace equation
 - Update by averaging data of up, down, left, right four points
 - Prepare two arrays old and new to keep the old (previous) value
 - Region segmentation, region division
 - Compute the residual to check the convergence



Matrix decomposition and nearest neighbor communication

- Block distribution of 2D region
- To update boundary elements, boundary elements of neighbors are required
- Data exchange of boundary elements



Blocking point-to-point communication

- Send/Receive

```
MPI_Send(  
    void          *send_data_buffer, // address of input data  
    int           count,          // data count  
    MPI_Datatype data_type,      // data type  
    int           destination,   // destination process rank  
    int           tag,           // message tag  
    MPI_Comm     communicator,   // communicator  
) ;
```

```
MPI_Recv(  
    void          *recv_data_buffer, // address of receive data  
    int           count,          // data count  
    MPI_Datatype data_type,      // data type  
    int           source,         // source process rank  
    int           tag,           // message tag  
    MPI_Comm     communicator,   // communicator  
    MPI_Status   *status        // status information  
) ;
```

Point-to-point communication

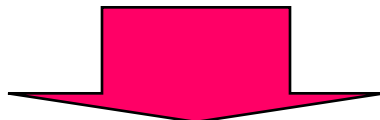
- Message is specified by address and size
 - Typed: MPI_INT, MPI_DOUBLE, ...
 - Binary data can be specified by MPI_BYTE with message size in byte
- Source/destination is specified by process rank and message tag
 - MPI_ANY_SOURCE for any source process rank
 - MPI_ANY_TAG for any message tag
- Status information includes the source rank, size, tag of the received message

Nonblocking point-to-point communication

- MPI_Isend/Irecv initiates the communication, MPI_Wait waits for the completion in semantics of blocking communication
 - Computation and communication can be overlapped if the communication can be executed in the background

```
int MPI_Isend( void *buf, int count, MPI_Datatype datatype,  
    int dest, int tag, MPI_Comm comm, MPI_Request *request )
```

```
int MPI_Irecv( void *buf, int count, MPI_Datatype datatype,  
    int source, int tag, MPI_Comm comm, MPI_Request *request )
```



```
int MPI_Wait ( MPI_Request *request, MPI_Status *status)
```

Process topology

- int **MPI_Cart_create**(MPI_Comm *comm_old*,
int *ndims*, int **dims*, int **periods*, int *reorder*,
MPI_Comm **comm_cart*);
 - Creates *comm_cart* with *ndims* dimensional hypercube topology
 - Process size of each dimension is specified by *dims*
 - *Periods* specified whether each dimension is periodical or not
 - *Reorder* specifies whether it allows renumbering of ranks between old and new communicators

Source/destination of shift communication

- int **MPI_Cart_shift**(MPI_Comm *comm*, int *direction*, int *disp*, int **rank_source*, int **rank_dest*);
 - *Direction* specified the dimension of shift communication
 - It is 0 to *ndims*-1 in *ndims* dimension case
 - Disp is a displacement of shift communication
 - It returns *rank_source* as a source rank and *rank_dest* as a destination rank
 - If the boundary is not periodical, it returns MPI_PROC_NULL if it exceeds the boundary

```

/* calculate process ranks for 'down' and 'up' */
MPI_Cart_shift(comm, 0, 1, &down, &up);

/* recv from down */
MPI_Irecv(&uu[x_start-1][1], YSIZE, MPI_DOUBLE, down, TAG_1,
            comm, &req1);
/* recv from up */
MPI_Irecv(&uu[x_end][1], YSIZE, MPI_DOUBLE, up, TAG_2,
            comm, &req2);

/* send to down */
MPI_Send(&u[x_start][1], YSIZE, MPI_DOUBLE, down, TAG_2, comm);
/* send to up */
MPI_Send(&u[x_end-1][1], YSIZE, MPI_DOUBLE, up, TAG_1, comm);

MPI_Wait(&req1, &status1);
MPI_Wait(&req2, &status2);

```

In a process of rank 0 and numprocs-1, **MPI_Cart_shift** returns **MPI_PROC_NULL**
 No need to treat specially. **MPI_Send** and **Irecv** do not do anything if
MPI_PROC_NULL is specified

```
/*
 * Laplace equation with explicit method
 */
#include <stdio.h>
#include <stdlib.h>
#include <math.h>
#include <mpi.h>

/* square region */
#define XSIZE 256
#define YSIZE 256
#define PI 3.1415927
#define NITER 10000
double u[XSIZE + 2][YSIZE + 2], uu[XSIZE + 2][YSIZE + 2];
double time1, time2;
void lap_solve(MPI_Comm);
int myid, numprocs;
int namelen;
char processor_name[MPI_MAX_PROCESSOR_NAME];
int xsize;
```

2D target domain
Uu is for new values

```
void
initialize()
{
    int x, y;

    /* initialization */
    for (x = 1; x < XSIZE + 1; x++)
        for (y = 1; y < YSIZE + 1; y++)
            u[x][y] = sin((x - 1.0) / XSIZE * PI) +
                cos((y - 1.0) / YSIZE * PI);

    /* zero clear in the boundary */
    for (x = 0; x < XSIZE + 2; x++) {
        u [x][0] = u [x][YSIZE + 1] = 0.0;
        uu[x][0] = uu[x][YSIZE + 1] = 0.0;
    }
    for (y = 0; y < YSIZE + 2; y++) {
        u [0][y] = u [XSIZE + 1][y] = 0.0;
        uu[0][y] = uu[XSIZE + 1][y] = 0.0;
    }
}
```

```
#define TAG_1 100
#define TAG_2 101

#ifndef FALSE
#define FALSE 0
#endif

void lap_solve(MPI_Comm comm)
{
    int x, y, k;
    double sum;
    double t_sum;
    int x_start, x_end;
    MPI_Request req1, req2;
    MPI_Status status1, status2;
    MPI_Comm comm1d;
    int down, up;
    int periods[1] = { FALSE };
```

```

/*
 * Create one dimensional cartesian topology with
 * nonperiodical boundary
 */
MPI_Cart_create(comm, 1, &numprocs, periods, FALSE, &comm1d);
/* calculate process ranks for 'down' and 'up' */
MPI_Cart_shift(comm1d, 0, 1, &down, &up);

x_start = 1 + xsize * myid;
x_end = 1 + xsize * (myid + 1);

```

- Create *comm1d* with one dimensional topology
 - The boundary is not periodical
- Obtain the *up* and *down* process rank
 - The boundary process may obtain MPI_PROC_NULL

```
for (k = 0; k < NITER; k++){
    /* old <- new */
    for (x = x_start; x < x_end; x++)
        for (y = 1; y < YSIZE + 1; y++)
            uu[x][y] = u[x][y];

    /* recv from down */
    MPI_Recv(&uu[x_start - 1][1], YSIZE, MPI_DOUBLE,
             down, TAG_1, comm1d, &req1);
    /* recv from up */
    MPI_Recv(&uu[x_end][1], YSIZE, MPI_DOUBLE,
             up, TAG_2, comm1d, &req2);
    /* send to down */
    MPI_Send(&u[x_start][1], YSIZE, MPI_DOUBLE,
             down, TAG_2, comm1d);
    /* send to up */
    MPI_Send(&u[x_end - 1][1], YSIZE, MPI_DOUBLE,
             up, TAG_1, comm1d);

    MPI_Wait(&req1, &status1);
    MPI_Wait(&req2, &status2);
```

```

/* update */
for (x = x_start; x < x_end; x++)
    for (y = 1; y < YSIZE + 1; y++)
        u[x][y] = .25 * (uu[x - 1][y] + uu[x + 1][y] +
                           uu[x][y - 1] + uu[x][y + 1]);
}
/* check sum */
sum = 0.0;
for (x = x_start; x < x_end; x++)
    for (y = 1; y < YSIZE + 1; y++)
        sum += uu[x][y] - u[x][y];
MPI_Reduce(&sum, &t_sum, 1, MPI_DOUBLE, MPI_SUM, 0, comm1d);
if (myid == 0)
    printf("sum = %g\n", t_sum);
MPI_Comm_free(&comm1d);
}

```

```
int
main(int argc, char *argv[])
{
    MPI_Init(&argc, &argv);
    MPI_Comm_size(MPI_COMM_WORLD, &numprocs);
    MPI_Comm_rank(MPI_COMM_WORLD, &myid);
    MPI_Get_processor_name(processor_name, &namelen);
    fprintf(stderr, "Process %d on %s\n", myid, processor_name);

    xsize = XSIZE / numprocs;
    if ((XSIZE % numprocs) != 0)
        MPI_Abort(MPI_COMM_WORLD, 1);
    initialize();
    MPI_Barrier(MPI_COMM_WORLD);
    time1 = MPI_Wtime();
    lap_solve(MPI_COMM_WORLD);
    MPI_Barrier(MPI_COMM_WORLD);
    time2 = MPI_Wtime();
    if (myid == 0)
        printf("time = %g\n", time2 - time1);
    MPI_Finalize();
    return (0);
}
```

Things to improve

- This program allocates the whole array although it is not necessary
 - When the partial array is allocated, the index of array should be computed from global index to local index
 - This is essential to solve large-scale problem using distributed memory machine
- Two dimensional distribution of 2D array is more efficient than one dimensional distribution
 - Reduce the communication size
 - Can be parallelized by more number of processors

Summary

- Parallel computing is indispensable to achieve high performance
- OpenMP is easy, but may be efficient up to 16 processors
- For more number of processors, MPI is only the choice
 - Tradeoff between the programming cost and execution time
 - We expect a new programming language that replaces MPI for longer term
- It is not **too** difficult to parallelize scientific applications
 - It has inherent data parallelism
 - Most scientific applications can be parallelized in some “parallelization patterns”

Parallel computing in coins

- viola0[1-6].coins.tsukuba.ac.jp (6 node cluster)
 - 8 cores / node
 - 2.93GHz Nehalem x 2 sockets
 - 12GB memory / node
 - 1333MHz 2GB DDR3 x 3 channels x 2
 - Network bandwidth 4GB/s
 - 4x QDR Infiniband
 - Software
 - CentOS5.4
 - OpenMPI*, MVAPICH1, MVAPICH2
 - System default is OpenMPI
 - It can be selected by mpi-selector-menu
 - gcc, gfortran, Sun JDK6
 - BLAS, LAPACK, ScaLAPACK

Initial setting

- Ssh authentication setting

```
% ssh-keygen -t rsa  
% cat .ssh/id_rsa.pub >> .ssh/authorized_keys
```

- Create known hosts file (also for hostnames of IB interface such as viola01-ib0)

```
% echo StrictHostKeyChecking no >> .ssh/config  
% ssh viola01-ib0 hostname  
viola01.coins.tsukuba.ac.jp  
% ssh viola02-ib0 hostname  
viola02.coins.tsukuba.ac.jp  
...  
% ssh viola06-ib0 hostname  
viola06.coins.tsukuba.ac.jp
```

Selection of MPI

```
$ mpi-selector-menu
```

Current system default: openmpi-1.3.2-gcc-x86_64

Current user default: <none>

System default is OpenMPI, no
user default specified

"u" and "s" modifiers can be added to numeric and "U"
commands to specify "user" or "system-wide".

- 1. mvapich-1.1.0-gcc-x86_64
- 2. mvapich2-1.2-gcc-x86_64
- 3. openmpi-1.3.2-gcc-i386
- 4. openmpi-1.3.2-gcc-x86_64
- U. Unset default
- Q. Quit

Select MVAPICH2

Selection (1-4[us], U[us], Q): **2u**

Compilation

- How to compile an MPI program

```
% mpicc -O2 a.c
```

- If you select MPI, you need to re-compile
 - Not binary compatible among OpenMPI and MVAPICH

How to execute MPI program in OpenMPI

- Create a host file

```
% cat hosts-openmpi  
viola01-ib0 slots=8  
viola02-ib0 slots=8  
...  
viola06-ib0 slots=8
```

- Execute an MPI program (OpenMPI)

```
% mpirun –hostfile hosts-openmpi –np 48 a.out
```

How to execute MPI program in MVAPICH

- Create a host file

```
% cat hosts  
viola01-ib0  
viola02-ib0  
...  
viola06-ib0  
% cat hosts hosts hosts hosts hosts hosts hosts hosts >  
hosts-mvapich
```

- Execute an MPI program

```
% mpirun_rsh –hostfile hosts-mvapich –np 48 a.out
```

- Note that the execution command in MVAPICH is mpirun_rsh

Open source implementation

OpenMP

- GNU GCC 4.2 or later
 - % cc -fopenmp . . .
- Omni OpenMP Compiler
 - <http://phase.hpcc.jp/Onni/>

MPI

- OpenMPI
 - <http://www.openmpi.org/>
- MPICH2
 - <http://www-unix.mcs.anl.gov/mpi/mpich2/>
- YAMPII
 - <http://www.il.is.s.u-tokyo.ac.jp/yampii/>