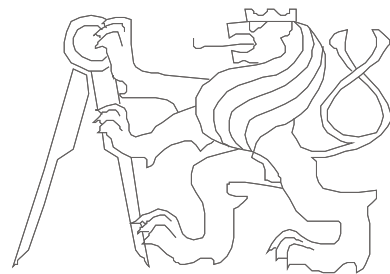


Advanced Computer Architectures

Parallel systems programming – Part II. OpenMP a MPI



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Overview: OpenMP vs. MPI

OpenMP

(Open Multi-Processing)

- Only for SMS
- Simple parallelization (using directives)
- Implicit communication

MPI

(Message Passing Interface)

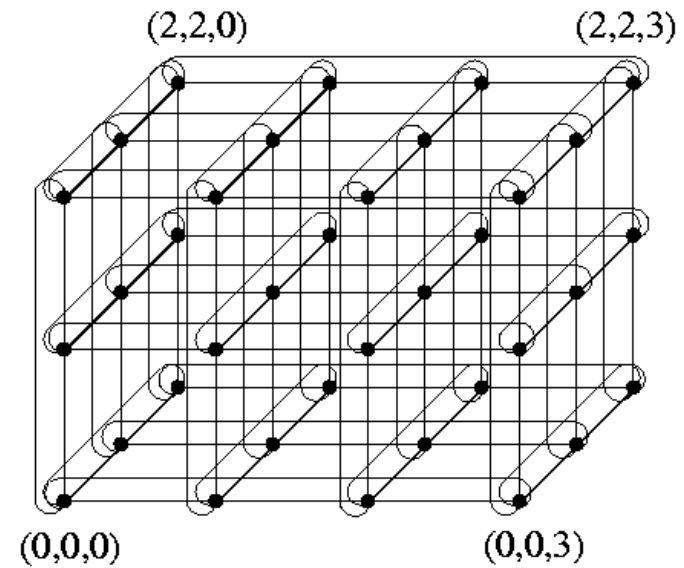
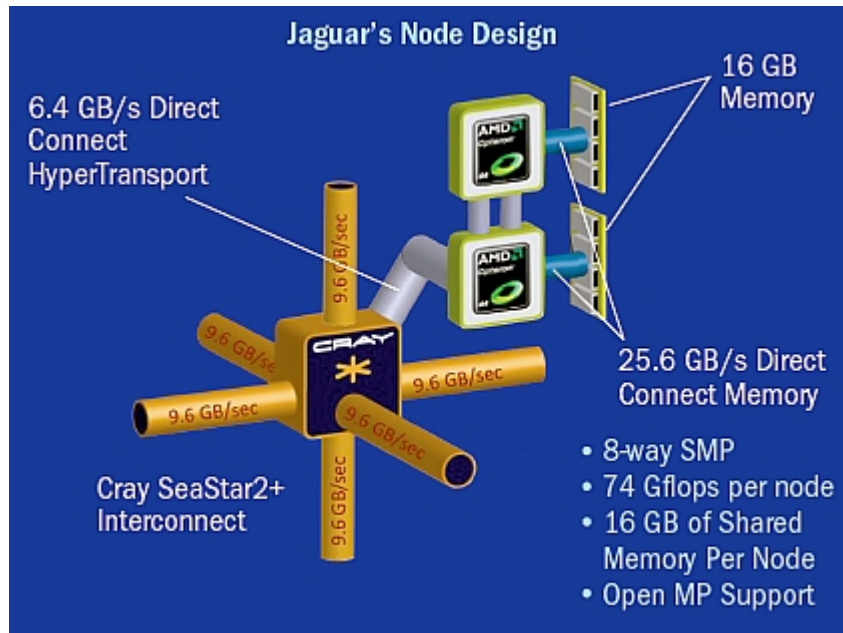
- For both SMS and DMS
- Demanding for programmer (parallelization, debugging)
- Explicit communication

Hybrid approach: OpenMP + MPI

- the current software trend in parallel computing
- MPI across cluster nodes, OpenMP inside node (resource efficient, suppressing intra-node communication overload, dynamic load balancing)
- it is possible to achieve higher speedup than when using only OpenMP or MPI

MPP – Cray XT5-HE

- $\approx 37\,000$ computation nodes (224 162 PE), service nodes for I/O
- One node – two 6-core processors – both have access to shared memory
- Each node – 25.6 GB/s into local memory (XT6: 85.3 GB/sec), 6.4GB/s into network,
- Nodes are interconnected into 3D toroid
- 12 OpenMP or MPI tasks on node simultaneously, MPI between nodes

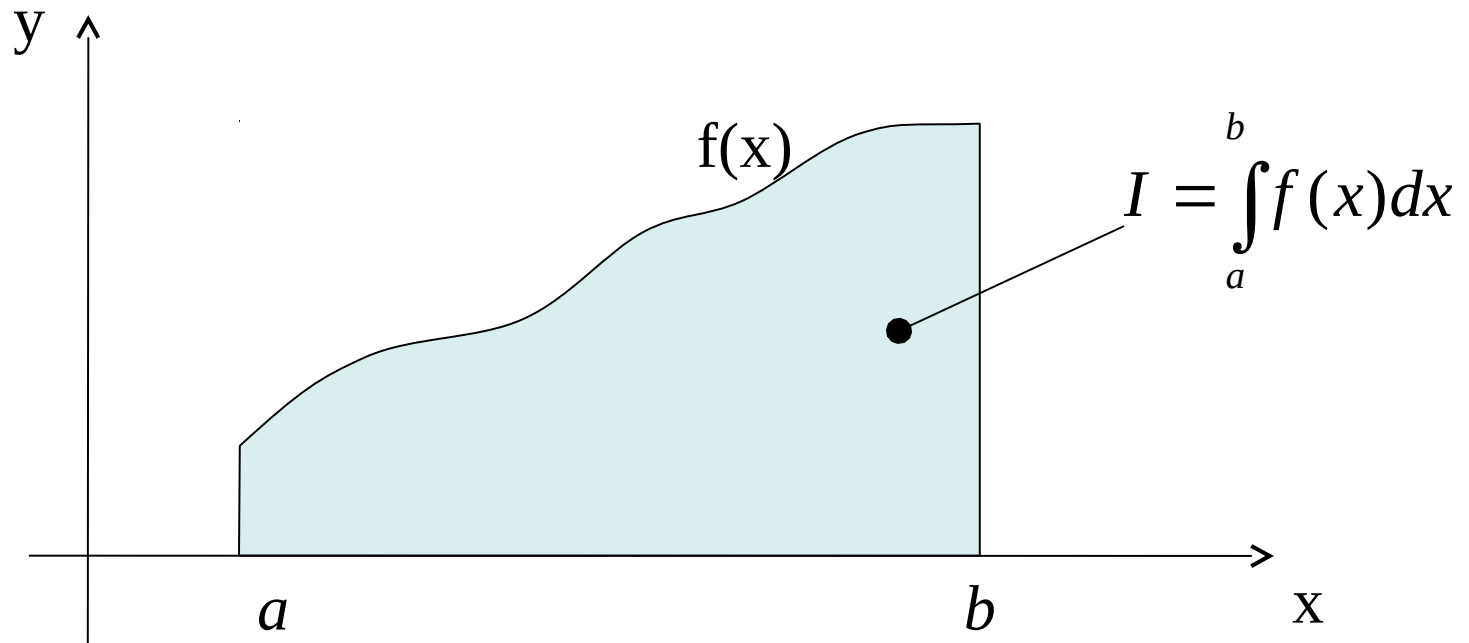


Cluster

- provides computing power by connecting multiple computers (usually a medium price category)
- used by 85% of today's 500 most powerful systems
- the advantage is lower price, good scalability



Example – calculation of a definite integral



Example – calculation of a definite integral

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

```
double integral(double a, double b, double (*f)(double), unsigned long int N)
{
    double sum=0, dx=(b-a)/N, x=a+dx;
    long int i;

    for(i=1; i<N; i++)
        sum += f(x+i*dx);
    sum += (f(a)+f(b))/2;
    return sum*dx;
}
```

```
double polynomial(double x) { return 2*x+1; }
```

```
void main() {
    long int N=5e8;
    double res, a=4, b=5;
    res = integral(a, b, polynomial, N);
    printf("I = %lf ",res);
}
```

Example – calculation of a definite integral – OpenMP simple

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

double integral(double a, double b, double (*f)(double), unsigned long int N)
{
    double sum=0, dx=(b-a)/N, x=a+dx;
    long int i;
#pragma omp parallel for shared(x,N,dx,f) private(i) reduction(+:sum)
    for(i=1; i<N; i++)
        sum += f(x+i*dx);
    sum += (f(a)+f(b))/2;
    return sum*dx;
}
```

```
double polynomial(double x) { return 2*x+1; }
```

```
void main() {
    long int N=5e8;
    double res, a=4, b=5;
    res = integral(a, b, polynomial, N);
    printf("I = %lf ",res);
}
```

Speedup: 1,85 !!!

OpenMP – overview

- All starts with: **#pragma omp directive [clause list]**
- program is executed sequentially until parallel directive is specified
- clause list specifies conditions for parallelization, number of threads, data access, sharing, partitioning and manipulation with them...
 - conditional parallelization: **if(scalar expression)**
 - degree of parallelization: **num_threads(integer expression)**
 - data access methods and rules: **shared(variable list), private(variable list), firstprivate(variable list)**
 - collection of private data at final stage of parallel execution: **reduction(operator: variable list)**, operators: +, *, -, &, |, ^, &&, ||
 - partitioning and scheduling: **schedule(scheduling_class[, parameter])**, where you can choose from: static, dynamic, guided, and runtime
- relaxed-consistency (if the shared variable must be seen identically by all threads, the programmer must take care of it – FLUSH)

OpenMP – *for* loops parallelization

```
#pragma omp parallel default (private) shared (n)
{
    #pragma omp for
    for (int i = 0; i < n; i++) {
        /* loop body */
    }
}
```

or combined in short:

```
#pragma omp parallel for default (private) shared (n)
for (int i = 0; i < n; i++) {
    /* loop body */
}
```

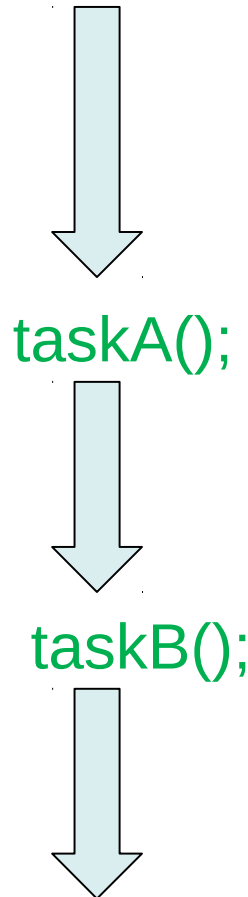
OpenMP – *for* loops parallelization

```
#pragma omp parallel default (private) shared (n)
{
    #pragma omp for
    for (i = 1; i < n; i++) {
        a[i] = i + a[i-1];
    }
}
```

?

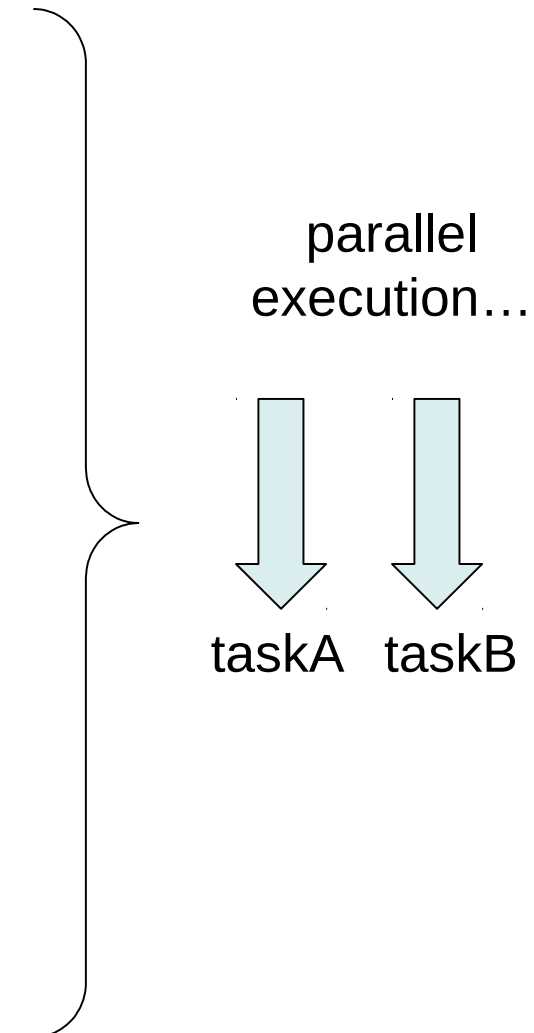
OpenMP – each thread works on unique problem partition

Sequential execution...



OpenMP – each thread works on unique problem partition

```
1 #pragma omp parallel
2 {
3     #pragma omp sections
4     {
5         #pragma omp section
6         {
7             taskA();
8         }
9         #pragma omp section
10        {
11             taskB();
12         }
13     }
14 }
```



OpenMP – each thread works on unique problem partition

or in short:

```
1 #pragma omp parallel sections
2 {
3     #pragma omp section
4     {
5         taskA();
6     }
7     #pragma omp section
8     {
9         taskB();
10    }
11 }
```

OpenMP – what it provides to programmer?

- nested parallelism (OMP_NESTED set to TRUE)
- explicit barriers specification (point of synchronization)
- mark code which should be executed by one thread only (arbitrarily selected or master only)
- mark critical section (only one thread can enter and execute at given time instant; others are required to wait or already have the given section done)
- ordered execution (as in serial execution)
- enforcing memory update and read (variable update ...) – flush; when is flush used implicitly? A flush is implied at a barrier, at the entry and exit of critical, ordered, parallel, parallel for, and parallel sections blocks and at the exit of for, sections, and single blocks. A flush is not implied if a nowait clause is present. It is also not implied at the entry of for, sections, and single blocks and at entry or exit of a master block.

OpenMP – what are other provided functionalities?

- void omp_set_num_threads (int num_threads);
 - int omp_get_num_threads ();
 - int omp_get_max_threads ();
 - int omp_get_thread_num ();
 - int omp_get_num_procs ();
 - int omp_in_parallel();
-
- void omp_init_lock (omp_lock_t *lock);
 - void omp_destroy_lock (omp_lock_t *lock);
 - void omp_set_lock (omp_lock_t *lock);
 - void omp_unset_lock (omp_lock_t *lock);
 - int omp_test_lock (omp_lock_t *lock);

And many other functions...

Example – calculation of a definite integral – serial

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

```
double integral(double a, double b, double (*f)(double), long N);
```

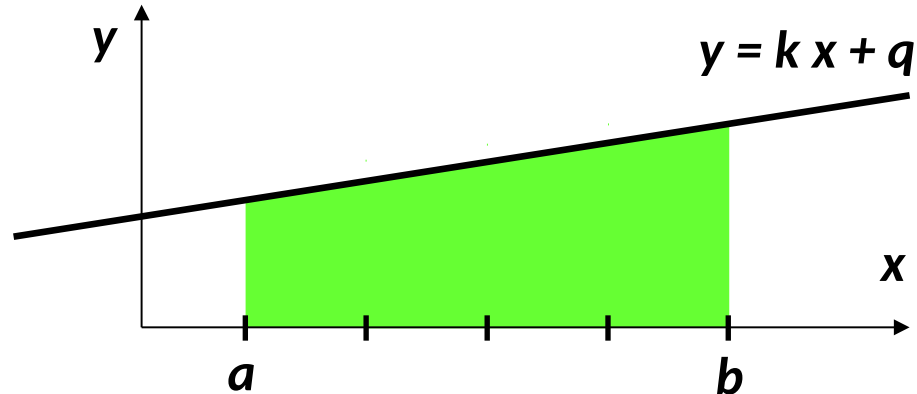
```
double polynomial(double x) {  
    return 2*x+1;  
}
```

```
void main() {  
    long int N=5e8;  
    double res, a=4, b=5;
```

```
    res = integral(a, b, polynomial, N);
```

```
    printf("I = %lf ",res);
```

```
}
```



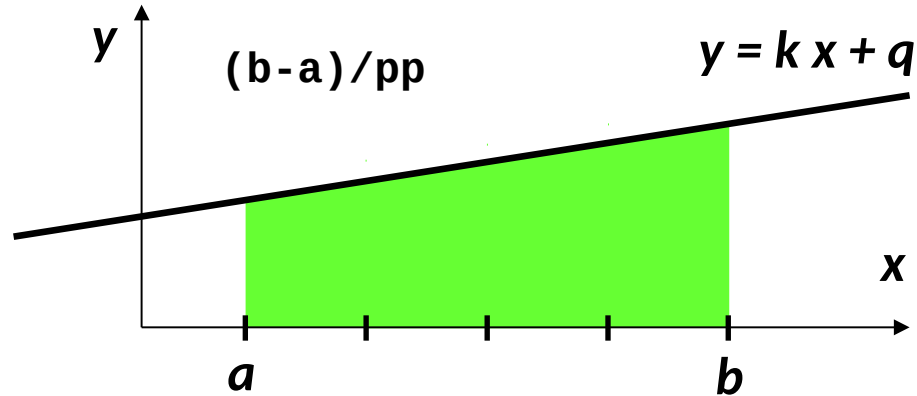
Example – calculation of a definite integral – preparation

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

```
double integral(double a, double b, double (*f)(double), long N);
```

```
double polynomial(double x) {  
    return 2*x+1;  
}
```

```
void main() {  
    long int N=5e8;  
    double res, a=4, b=5;  
    int i=0, pp=1;
```



$\text{res} = \text{integral}(a + ((b-a)/pp)*i, a + ((b-a)/pp)*(i+1), \text{polynomial}, N/pp);$

```
printf("I = %lf ",res);  
}
```

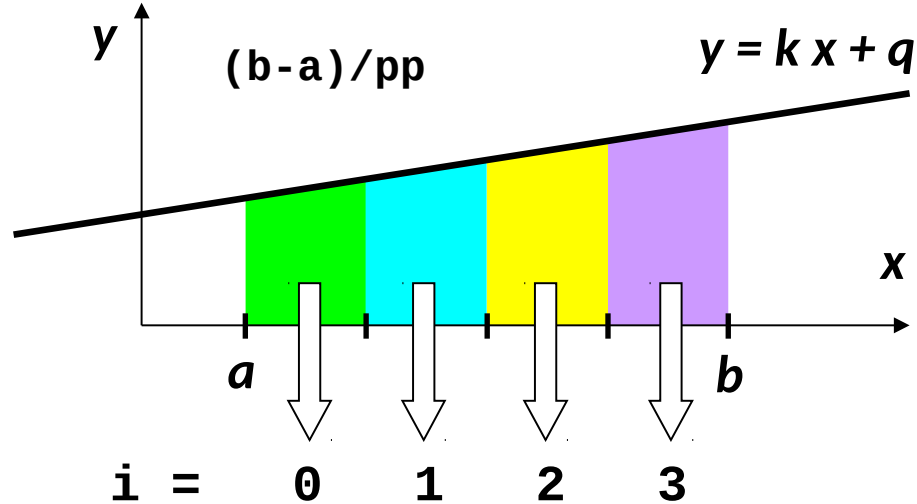
Example – calculation of a definite integral – parallel

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

```
double integral(double a, double b, double (*f)(double), long N);
```

```
double polynomial(double x) {  
    return 2*x+1;  
}
```

```
void main() {  
    long int N=5e8;  
    double res, a=4, b=5;  
    int i=???, pp=4;
```



```
res = integral(a+((b-a)/pp)*i, a+((b-a)/pp)*(i+1), polynomial, N/pp);
```

```
printf("I = %lf ",res);
```

```
}
```

Example – calculation of a definite integral – parallel

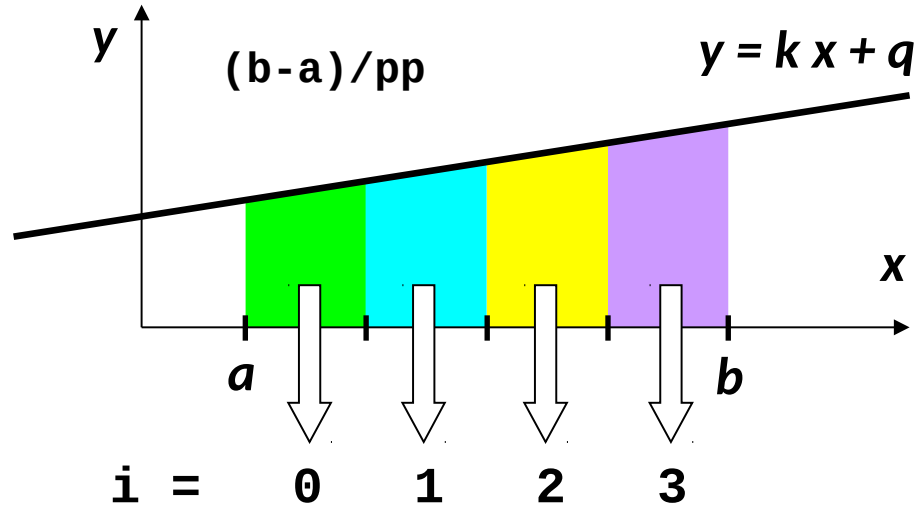
```
#include <stdio.h>
#include <omp.h>
```

```
double integral(double a, double b, double (*f)(double), long N);
```

```
double polynomial(double x) {
    return 2*x+1;
}
```

```
void main() {
    long int N=5e8;
    double res, a=4, b=5;
    int i, pp=4;
```

```
#pragma omp parallel private(i) reduction(+:res) num_threads(pp)
{
    i = omp_get_thread_num();
    res = integral(a+((b-a)/pp)*i, a+((b-a)/pp)*(i+1), polynomial, N/pp);
}
printf("I = %lf ",res);
}
```



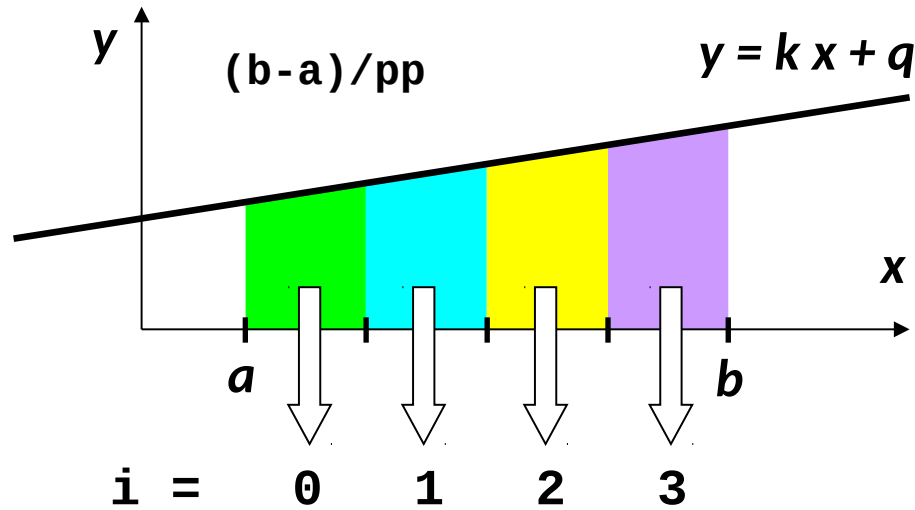
Example – calculation of a definite integral – parallel

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

```
double integral(double a, double b, double (*f)(double), long N);
```

```
double polynomial(double x) {
    return 2*x+1;
}
```

```
void main() {
    long int N=5e8;
    double res, a=4, b=5;
    int i, pp;
    pp = omp_get_num_procs();
    #pragma omp parallel private(i) reduction(+:res) num_threads(pp)
    {
        i = omp_get_thread_num();
        res = integral(a+((b-a)/pp)*i, a+((b-a)/pp)*(i+1), polynomial, N/pp);
    }
    printf("I = %lf ",res);
}
```



OpenMP – another example

```
#include <omp.h>
int a, b, i, tid;
#pragma omp threadprivate(a)

main () {
    omp_set_dynamic(0); /* Explicitly turn off dynamic threads */

    #pragma omp parallel private(b,tid)
    {
        tid = omp_get_thread_num();
        a = b = tid;
        printf("Thread %d: a,b,%d %d \n",tid,a,b);
    } /* end of parallel section */

    #pragma omp parallel private(tid)
    {
        tid = omp_get_thread_num();
        printf("Thread %d: a,b,%d %d \n",tid,a,b);
    } /* end of parallel section */
}
```

MPI – Message Passing Interface

Explicit communication, that is why:

- Most of MPI functions require communicator as argument
- **communicator** can be thought of as a handle (object) to a group of processes which can communicate with each other
- communicator `MPI_COMM_WORLD` includes all processes of given program
- processes belonging to the given communicator have assigned their unique identification number – `RANK`, which is integer in $(0, N-1)$ – N is number of processes
- communication is either collective (all the communicator processes are involved) or point-to-point

MPI – Message Passing Interface

Collective communication:

- is always blocking
- uses predefined MPI types
(MPI_CHAR, MPI_INT, MPI_LONG,...)
- division according to purpose:
 - synchronization (processes are waiting at a given location)
 - collective computation (reduction) – when one process collects data from all the other processes and final result is computed by mutual operation between partial results
 - data movement (Broadcast, Scatter, Gather, All-to-All)

MPI – almost minimal program

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

void main(int argc, char *argv[]) {
    int numprocs, rank, namelen;
    char proc_name[MPI_MAX_PROCESSOR_NAME];

    MPI_Init(&argc, &argv);
    MPI_Comm_size(MPI_COMM_WORLD, &numprocs);
    MPI_Comm_rank(MPI_COMM_WORLD, &rank);
    MPI_Get_processor_name(proc_name, &namelen);

    printf("Process %d on %s out of %d\n", rank, proc_name,
        numprocs);

    MPI_Finalize();
}
```

Example – calculation of a definite integral – MPI

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

inline double f(double x) { return 2*x+1; }

int main(int argc, char *argv[])
{
    int myid, numprocs, volba=0;
    unsigned long int i, n;
    double startwtime, endwtime, a = 4.0, b = 5.0;
    double total, integral = 0.0;

    MPI_Init(&argc, &argv);
    MPI_Comm_size(MPI_COMM_WORLD, &numprocs); // obtain number
                                                // of processes
    MPI_Comm_rank(MPI_COMM_WORLD, &myid); // obtain process number
```

Example – calculation of a definite integral – MPI

```
do {
    if (myid == 0) {
        printf("\n Enter number of intervals (0 to exit): "); fflush(stdout);
        scanf("%ld", &n);
        if(n==0) break;
        startwtime = MPI_Wtime();
    }

// send variable n, with vector length 1, type int, from process #0 to others
MPI_Bcast(&n, 1, MPI_INT, 0, MPI_COMM_WORLD);
```

Example – calculation of a definite integral – MPI

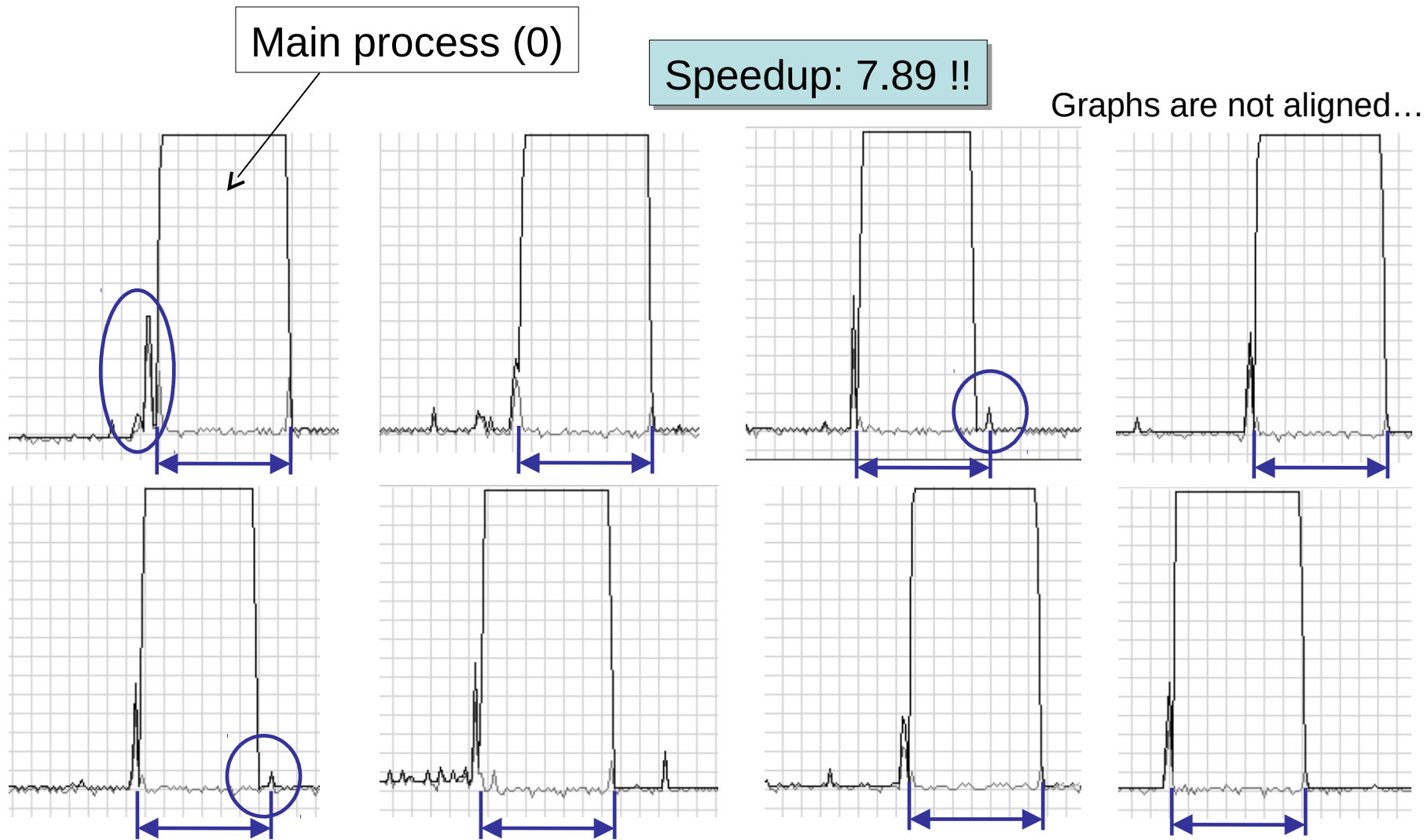
```
if (n != 0) {  
    double h = (b - a) / n;  
    double i1 = myid*(n/ numprocs);  
    double i2 = (myid+1)*(n/numprocs);  
  
    integral= ( f(a+i1*h) + f(a+i2*h) ) / 2;  
  
    for( i=i1+1 ; i<i2 ; i++ )  
        integral += f(a+i*h);  
}  
// summarize integral variable from all processes to total in process 0,  
// length 1, type double, operation sum/addition  
MPI_Reduce(&integral, &total, 1, MPI_DOUBLE, MPI_SUM, 0,  
MPI_COMM_WORLD);
```

Example – calculation of a definite integral – MPI

```
if (myid == 0) {  
    endwtime = MPI_Wtime();  
    printf("I= %f\n", total);  
    printf("spent time: %f s\n", endwtime - startwtime); fflush(stdout);  
}  
} while (n != 0);  
  
MPI_Finalize();  
return 0;  
}
```

Example – calculation of a definite integral – MPI

Computation on 8 single-core processors:



Combination of OpenMP a MPI – almost minimal program

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

int main(int argc, char *argv[]) {
    int numprocs, rank, namelen, iam = 0, np;
    char processor_name[MPI_MAX_PROCESSOR_NAME];

    MPI_Init(&argc, &argv);
    MPI_Comm_size(MPI_COMM_WORLD, &numprocs);
    MPI_Comm_rank(MPI_COMM_WORLD, &rank);
    MPI_Get_processor_name(processor_name, &namelen);

#pragma omp parallel default(shared) private(iam, np)
{
    np = omp_get_num_threads();
    iam = omp_get_thread_num();
    printf("Hello from thread %d out of %d from process %d out of %d on
%s\n", iam, np, rank, numprocs, processor_name);
}
    MPI_Finalize();
    return 0;
}
```