

## Outline

Introduction to parallel computation:

- Moore's law
- Parallel computing
- Parallel computers
- Single Program Multiple Data paradigm
- Measuring performances
- Amdahl's law
- Message Passing Interface

Scope: Unity makes strength!

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Computational complexity

- Not all computational problems are created equal. Some problems are so simple that even large instances may be solved rapidly. Other problems are intractable – once the problem is large enough there is essentially no hope of solving it exactly.
- Computer scientists have formalized this observation, and divided problems into computational complexity classes of varying difficulty. Knowing the complexity class of a problem won't solve it, but it will help you decide what kind of solution method is appropriate.
- If the problem is hard, finding an exact solution is apt to be costly or impractical, and you will probably have to resort to enumerative methods (which may be slow or useless for large cases) or settle for an approximate solution obtained with heuristic/stochastic methods
- How long does it take to find the maximum in a set of numbers? The answer should be a function of the size of the particular instance to be solved. The function tells how the algorithm run time grows as the input size increases. Suppose the numbers are A(i): i=1, ..., n. The obvious algorithm finds the maximum in time proportional to n. We say the algorithm runs in linear time, or is O(n).

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- If instead we wish to sort the numbers in ascending order, the fastest algorithms (e.g. quicksort) require time proportional to n log n. We say that sorting is done in O(n log n) time.
- The major distinction we make between algorithms is whether they
  take polynomial time or not. An algorithm requires polynomial time if
  for some k it has a time bound of O(nk)
- In a decision problem, given an input, we are required to give a Boolean YES/NO (1/0) answer. That is, in a decision problem we are only asked to verify whether the input satisfies a certain property.
- We denote by P the class of decision problems that are solvable in polynomial time. A standard convention is to call an algorithm "feasible" if belongs to P, i.e. if there is some polynomial p such that the algorithm runs in time at most p(n) on inputs of length n.
- Most algorithms that are not polynomial are exponential, requiring more than c<sup>dn</sup> time in the worst case, for some constants c>1 and d>0.
   Any exponential function will eventually exceed any polynomial function as n increases. In practice, exponential time algorithms are usually slower than polynomial time algorithms even for quite modest values of n.

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## NP complexity

• Let us consider the subset sum problem: Assume that we are given some integers, such as  $\{-7, -3, -2, 5, 8\}$ , and we wish to know whether some of these integers sum up to zero.

In this example, the answer is "yes", since the subset of integers 3, -2, 5} corresponds to the sum (-3) + (-2) + 5 = 0. The task of deciding whether such a subset with sum zero exists is called the subset sum problem.

• To answer if some of the integers add to zero we can create a combinatorial algorithm which obtains all the possible subsets. As the number of integers that we feed into the algorithm becomes larger, the number of subsets grows exponentially (2<sup>n</sup>) and so does the computation time, in fact:

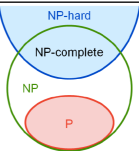
Binomial Theorem: 
$$(x+y)^n = \sum_{k=0}^n \binom{n}{k} x^{n-k} y^k \underset{x=y=1}{\Rightarrow} 2^n = \sum_{k=0}^n \binom{n}{k}$$
 # of ways of choosing k elements from the set with n elements

 However, notice that, if we are given a particular subset, we can easily verify whether the subset sum is zero, by just summing up the integers of the subset. So if the sum is indeed zero, that particular subset is the proof for the fact that the answer is "yes".

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#### NP-complete complexity

- An algorithm that verifies whether a given subset has sum zero is called a verifier
- More generally, a problem is said to be NP (nondetermistic polynomial-time) if there exists a verifier V for the problem and the answer can be obtained in polynomial time

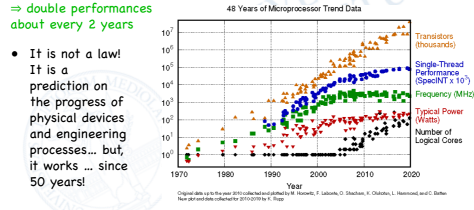


- The class of problems which are at least as hard as the hardest problems in NP is called NP-hard (nondetermistic polynomial-time hard problem)
- Finding the ground state (minimum energy state) of a generic Ising model is NP-hard (Barahona, 1982). So, maybe we can map hard problems on an Ising model ...
- Problems that are NP-hard do not have to be necessarily elements of NP; indeed, they may not even be decidable. So let's take the intersection of these two sets: NP-complete
- In computational complexity theory, NP-complete problems are contained in NP; although any given solution to an NP-complete problem can be verified quickly (in polynomial time... in fact, it is in NP), there is no known efficient way to locate a possible solution

## Moore's law

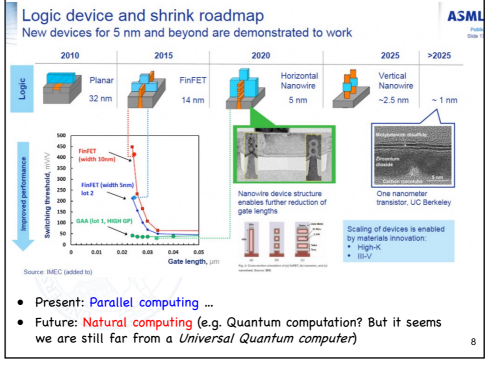
 Gordon E. Moore (co-founder of Intel) predicted (Electronics, Vol. 38, 1965) that transistor density of a semiconductor chip is going to double about every 2 years ...





 But, how long? It seems up to 2025. Note, that recently we had to move towards multi-core architectures

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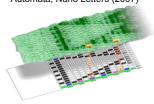


## Natural Computation

• Natural computing encompasses three classes of methods:

- 1. Methods based on the use of computers to synthesize natural phenomena (Computational Intelligence, Neural Networks, Genetic Algorithms, etc.)
- 2. Methods that employ natural materials (electrons, atoms, molecules, DNA) to compute.
- 3. Methods that take inspiration from nature for the development of novel computational architectures or problem-solving techniques
- A remarkable example of the second class, which is also a remarkable example of (this time expected) emergence/self-organization in a complex system is DNA computing.
- DNA computing is a form of parallel computing that uses DNA, biochemistry, and molecular biology hardware, instead of the traditional silicon-based computer technologies
- 4 key components coding: in principle 1 liter of fluid with 6 grams of DNA could memorize 3 Zbytes and perform up to 1 EFLOPS (Zetta=10<sup>21</sup>) (Exa=10<sup>18</sup>)

DNA thin slab self-assembly of a Sierpinski triangle as a Cellular Automata, *Nano Letters* (2007)



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## New computational architectures

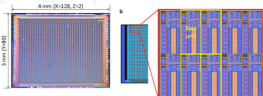
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 Along the third class of Natural computing methods, recently, several special hardwares or physical systems for solving Ising models have been proposed.

- D-Wave machines, based on quantum annealing
  - Dwave One (2011, 128 qubits)
  - Dwave Two (2013, 512 qubits)
  - Dwave 2X (2015, 1152 qubits)



 (classical) Ising chip by Hitachi (2015) 20480 spins based on CMOS (complementary metal-oxide semiconductor) annealing

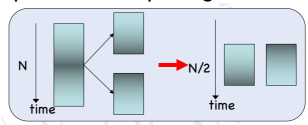


1. Coherent (quantum) Ising Machine (Optical/laser networks, 2011/2015)

## What is parallel computing?

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 Split a single application code to run across multiple processors



 Computational time may be reduced:

in principle if the number of CPU is p the time  $t_p$  involved to complete the computational task could be reduced to  $t_p=t_1/p$  (being  $t_1$  the time needed to complete the task serially)

- Memory requirements per processor may be reduced
- Rather then using one expensive processor (...maybe waiting some years to have it on the market), many less expensive processors can be used
- Programming is now more difficult, because dividing a problem to run efficiently across multiple processors is a difficult task
- The efficiency depends on the algorithms and on the characteristics of the parallel computer

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# Parallel computing examples

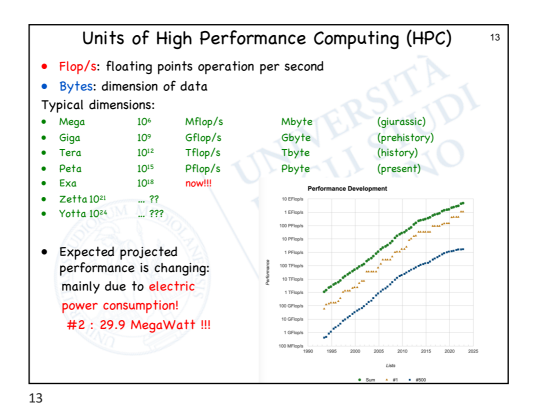
#### • Nature:

- Our brain is made of about 10<sup>11</sup> neurons
- The living organism are made from cells
- Leaves of trees
- Etc.etc.
- ... Reality?

#### Society:

- Highway tollgates
- Supermarket cash registers
- Written exams
- Etc.etc.





High Performance Computing (HPC)

• Present Number 1 (Autumn 2022):

Frontier - HPE Cray
# CPU cores: 8,730,112
Site: Oak Ridge National Laboratory, USA
Peak performance: 1,685.65 Pflop/s
Operating system: HPE Cray OS
Power: 21.1 MW

• First in Italy (N.4 in Autumn 2022)

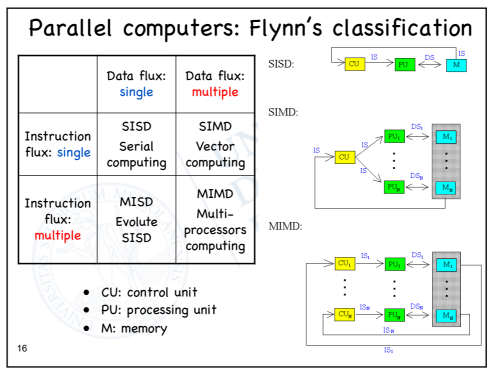
LEonardo - Atos
# CPU cores: 1,463,616
Site: CINECA (BO, Italy)
Peak performance: 255.75 Pflop/s
Operating system: Linux
Power: 5.6 MW

## Parallel computers

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- If we define a parallel computer as "a collection of computing elements able to communicate and to cooperate in order to solve quickly large computational tasks" ... then we have to answer to the following questions:
- "...a collection of computing elements..."
  - How many?
  - What is the computational power of each one of them?
  - What kind of memory they have?
- "...able to communicate..."
  - How they are connected?
  - What kind of communication protocol they use?
- "...and to cooperate..."
  - Is there any synchronization mechanism?
  - What is their operating systems?
- "...in order to solve..."
  - What algorithms are parallelizable?
  - How can I parallelize an algorithm?
  - Automatic or manual parallelization?

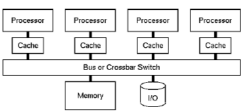
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# Parallel computers: memory classification

|                  | Shared address space            | Individual address space     |
|------------------|---------------------------------|------------------------------|
| Shared<br>memory | SMP<br>symmetric multiprocessor | N/A                          |
| Distributed      | NUMA                            | MPP                          |
| memory           | non uniform memory access       | massively parallel processor |

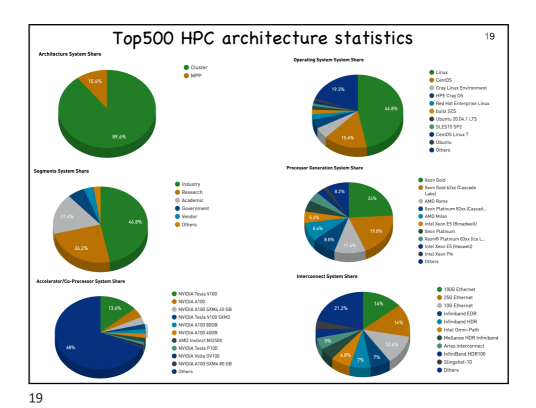
 SMP architecture uses shared system resources such as memory and I/O subsystem that can be accessed equally from all the processors.
 As shown in figure, each

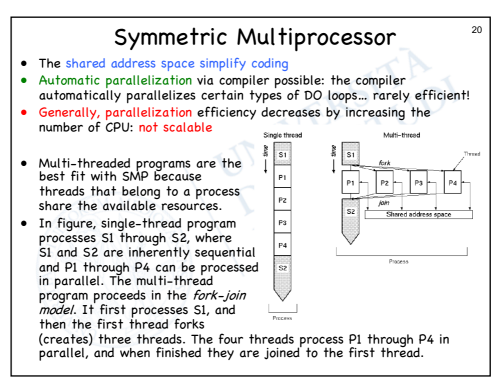


processor has its own cache which may have several levels.

- SMP machines have a mechanism to maintain coherency of data held in local caches.
- A single operating system controls the SMP machine and it schedules processes and threads on processors so that the load is balanced.

- MPP architecture consists of nodes connected by a network that is usually high-speed. Each node has its own processor, memory, and I/O subsystem (see the figure).
- The operating system is running on each node, so each node can be considered a workstation.
- Despite the term massively, the number of nodes is not necessarily large. In fact, there is no criteria. What makes the situation more complex is that each node can be an SMP node (typical situation!)
- NUMA architecture machines are built on a similar hardware model as MPP, but it typically provides a shared address space to applications using a hardware/software directory-based protocol that maintains cache coherency.
- As in an SMP machine, a single operating system controls the whole system. The memory latency varies according to whether you access local memory directly or remote memory through the interconnect. Thus the name non-uniform memory access.





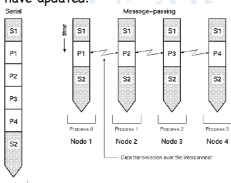
## MPP based on single-processor nodes

 If the address space is not shared among nodes, parallel processes have to transmit data over an interconnecting network in order to access data that other processes have updated.

One can have many different interconnection geometries

 Scalability is algorithmic dependent but can be efficient

- Manual parallelization
- One process runs on each node and the processes communicate with each other during the execution of the parallelizable part, P1-P4. In general, each process communicates with all the other processes. Due to

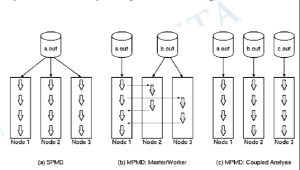


the communication overhead, work load unbalance, and synchronization, time spent for processing each of P1-P4 is generally longer in the message-passing program than in the serial program. All processes in the message-passing program are bound to S1 and S2.

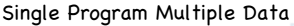
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## Message passing: parallel programming models

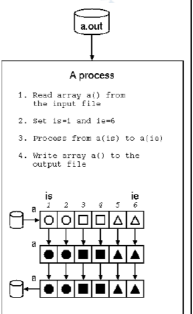
 When you run multiple processes with message-passing, there are further categorizations regarding how many different programs are cooperating in parallel execution.

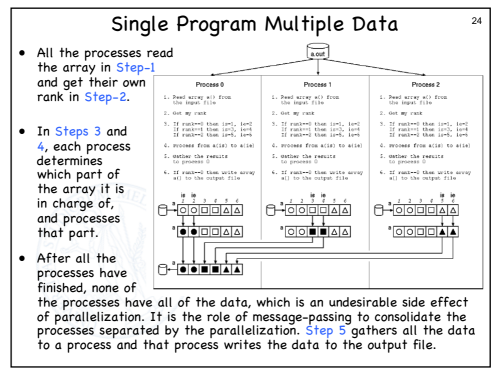


- In the SPMD (Single Program Multiple Data) model, there is only one program and each process uses the same executable working on different sets of data.
- The MPMD (Multiple Programs Multiple Data) model uses different programs for different processes, but the processes collaborate to solve the same problem. Typical usage of the MPMD model can be found in the master/worker style of execution or in the coupled analysis.



- This figure shows a sequential program that reads data from a file, does some computation on the data, and writes the data to a file.
- White circles, squares, and triangles indicate the initial values of the elements, and black objects indicate the values after they are processed.
- Remember that in the SPMD model, all the processes execute the same program. To distinguish between processes, each process has a unique integer called rank. You can let processes behave differently by using the value of rank.





#### Parallel programming: general notions

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 We will now introduce some basic notions of parallel programming via a simple example: imagine you need to compute the sum

```
S = a_1 + a_2 + ... + a_{16}
```

a total of 15 additions, with a computer with p processors.

- We will call T<sub>p</sub> (with p≥1) the time spent in computing S with a p-CPU computer via a parallel algorithm
- We will take the machine time to elaborate a single addition as standard unit; thus: T<sub>1</sub>=15
- Consider now the case p=2: the best way to divide the task among the 2 CPU is to ask to CPU-1 to compute

```
S_1 = a_1 + a_2 + ... + a_8 , and simultaneously to CPU-2 to compute S_2 = a_9 + a_{10} + ... + a_{16} .
```

Time spent to compute  $S_1$  and  $S_2$  is 7. At this point one of the two CPU can compute  $S=S_1+S_2$  while the other CPU remains idle. Total time is therefore  $T_2=8$ . We discuss now the cases p=3, 4, 8.

• (p=3) Step 1:  $S_1 = a_1 + ... + a_5$  /  $S_2 = a_6 + ... + a_{10}$  /  $S_3 = a_{11} + ... + a_{15}$  (time=4) Step 2:  $S_a = S_1 + S_2$  /  $S_b = S_3 + a_{16}$  (time=1 one CPU is idle) Step 3:  $S_1 = S_2 + S_3 + S_3 + a_{16}$  (time=1 two CPUs are idle) Total time:  $S_3 = S_3 + S_3 + S_3 + S_3 + S_3 + a_{16}$  (time=1 two CPUs are idle) Total time:  $S_3 = S_3 + S_3 + S_3 + S_3 + a_{16} + A_3 + a$ 

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#### Simple example ... continue

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• (p=4)

```
Step 1: S_1 = a_1 + ... + a_4 / S_2 = a_5 + ... + a_8 / S_3 = a_9 + ... + a_{12} / S_4 = a_{13} + ... + a_{16} (time=3)
```

Step 2:  $S_a = S_1 + S_2 / S_b = S_3 + S_4$  (time=1, two CPUs are idle)

Step 3:  $S = S_a + S_b$  (time=1, three CPUs are idle)

Total time:  $T_4=5$ 

• (p=8)

```
Step 1: S_1 = a_1 + a_2 / S_2 = a_3 + a_4 / ... / S_8 = a_{15} + a_{16} (time=1)
```

Step 2:  $S_a=S_1+S_2$  /  $S_b=S_3+S_4$  /  $S_c=S_5+S_6$  /  $S_d=S_7+S_8$  (time=1, 4 CPUs are idle)

Step 3:  $S_{\mu} = S_a + S_b / S_v = S_c + S_d$  (time=1, 6 CPUs are idle)

Step 4:  $S = S_{\mu} + S_{\nu}$  (time=1, 7 CPUs are idle)

Total time:  $T_8=4$ 

 As you should note, by increasing the number of CPUs the calculation of S is performed more and more quickly, but the efficiency decreases because more and more CPUs remain idle once the majority of the calculations have been performed

## Parallel programming: some definitions

Cost:

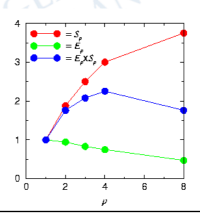
 $(S_p \le p ; if S_p \div p linear speed-up)$  $(E_p \le 1)$ Speed-up:

 $S_p = T_1/T_p$   $E_p = S_p/p = T_1/C_p$ Efficency:

In the optimization of a parallel calculation, a good compromise is obtained by trying to maximize the product  $[E_p \times S_p]$ 

Summary table:

|   |                | erranamin and  |      |      | 1 11             |
|---|----------------|----------------|------|------|------------------|
| Р | T <sub>p</sub> | C <sub>p</sub> | Sp   | Ep   | $E_p \times S_p$ |
| 1 | 15             | 15             | 1    | 1    | 1                |
| 2 | 8              | 16             | 1.88 | 0.94 | 1.76             |
| 3 | 6              | 18             | 2.5  | 0.83 | 2.08             |
| 4 | 5              | 20             | 3    | 0.75 | 2.25             |
| 8 | 4              | 32             | 3.75 | 0.47 | 1.76             |



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## Amdahl's law (1967)

- In late '60 there was considerable skepticism regarding the viability of massive parallelism; the skepticism was centered around Amdahl's law, an argument put forth by Gene Amdahl [AFIPS Conference Proceedings vol. 30 (Atlantic City, N.J., 1967) pp. 483-485] that even when the fraction of serial work in a given problem is small, say s, the maximum speed-up obtainable from even an infinite number of parallel processors is only 1/s
- Consider, in fact, a code which is composed by a fraction f that is completely parallelizable and by a fraction 1-f that is intrinsically serial and thus that cannot be parallelized (every code is in this situation). Imagine also to have a parallel computer with p CPUs. If  $T_1$  is the time spent in executing the code (serially) on a serial computer, we will have:

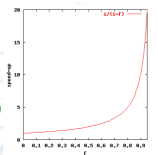


- $f \times T_1 = time$  spent in executing the fraction of the code P serially
- $(1-f)\times T_1$  = time spent in executing the fraction of the code S serially
- $(f \times T_1)/p$  = time spent in executing the fraction of code P on a parallel computer with p CPUs; thus

$$T_p = (1-f) \times T_1 + (f \times T_1)/p = T_1[f + (1-f)p]/p$$



- It now possible to obtain the speed-up:  $S_p=T_1/T_p=p/[f+(1-f)p]=1/[f/p+(1-f)]$
- By using an increasing number p of CPUs, one can reduce the ratio f/p to zero; It follows that S<sub>p</sub>≤ 1/(1-f). This is Amdahl's law. Is this the end of parallel computing?
- No! Usually the non-parallelizable fraction (1-f) tends to zero when computational complexity of the problem increase (Gustafson's law)



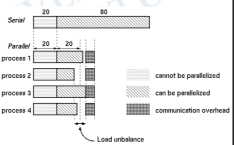
- Amdahl's Law approximately suggests: Suppose a car is travelling between two cities 120 Km apart and has already spent one hour travelling half the distance at 60 Km/h. No matter how fast you drive the last half, it is impossible to surpass 120 Km/h average (speed-up) before reaching the second city. Going infinitely fast you would only achieve 120 Km/h.
- Gustafson's Law approximately states: Suppose a car has already been travelling for some time at less than 120 Km/h. Given enough time and distance to travel, the car's average speed can always eventually reach and surpass 120 Km/h (speed-up), no matter how long or how slowly it has already travelled.

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#### Real cases

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- The situation just described is ideal, real cases are much more complicated
- Up to now we haven't considered that in real parallel computation there is a time spent in communicating (needed) partial results among
- Efficiency depends also on (communication overhead) how many data should travel across the network, on how many times data should travel on the network, on the network speed (bandwidth), on its latency



- Generally a good parallelization is obtained when:
  - The parallelizable fraction increases
  - Balance the workload of parallel processes
  - Communication is minimized as quantity of data and number of times
  - Design parallel algorithms from scratch (think parallel, write parallel!)

#### MPI: Message Passing Interface



- The Message Passing Interface (MPI) is a standard developed by the Message Passing Interface Forum (MPIF).
- MPIF, with the participation of more than 40 organizations (Convex, Cray, IBM, Intel, Meiko, nCUBE, NEC, ... but also USA Universities and National Laboratories), started working on the standard in 1992. The first draft (Version 1.0), which was published in 1994, was strongly influenced by the work at the IBM T. J. Watson Research Center.
- It specifies a portable interface for writing message-passing programs, and aims at practicality, efficiency, and flexibility at the same time.
- MPIF has further enhanced the first version to develop a second version (MPI-2) in 1997, a third version (MPI-3) in 2014/15 and a forthcoming version (MPI-4). For details about MPI and MPIF, visit <a href="http://www.mpi-forum.org">http://www.mpi-forum.org</a>
- It is a library for Fortran, C, C++, Python and other languages which implements the message passing model on parallel computers thus giving the possibility to processes running on different CPUs to synchronize information even if a common address space for memory (generally distributed) is lacking

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## MPI: Message Passing Interface



- Communication is obtained via explicit messages
- The MPI standard is small and large: it consists of functions for
  - Process identification
  - Point-to-point communication
  - Collective operations
  - Process groups
  - Synchronizations
  - Dynamical creation of processes (MPI-2; important for fault-tolerance)
  - I/O portable functions (MPI-2)
- But you can obtain everything via the following 6 functions:
  - MPI Init()
  - MPI Comm size()
  - MPI\_Comm\_rank()
  - MPI Send()
  - MPI Recv()
  - MPI Finalize()
- -> MPI initialization
- -> to know how many processes
- -> to know who am I
- -> to send a message
- -> to receive a message
- -> MPI finalization

#### A first example

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- Note that the program is executed in the SPMD (Single Program
   Multiple Data) model. All the nodes that run the program, therefore,
   need to see the same executable file with the same path name, which is
   either shared among nodes by NFS or other network file systems, or is
   copied to each node's local disk.
- This program does not include any communication, thus it is banally parallelized in the SPMD model

```
C++
                        Python
from mpi4py import MPI
                                                               #include <iostream>
comm = MPI.COMM_WORLD
                                                               using namespace std;
size = comm.Get size()
rank = comm.Get_rank()
                                                               int main(int argc, char* argv[])
print('Sono il nodo ', rank, ' dei ', size, ' che hai utilizzato!')
                                                               int size, rank:
                                                                MPI_Init(&argc,&argv);
                                                               MPI_Comm_size(MPI_COMM_WORLD, &size);
                                                               MPI_Comm_rank(MPI_COMM_WORLD, &rank);
                                                              cout<<" Sono il nodo "<<rank<<" dei "<<size<<" che hai
utilizzato!"<<endl;</pre>
                                                               MPI Finalize();
                                                               return 0;
```

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• Via the inclusion of mpi.h one defines MPI-related objects such as MPI COMM WORLD.

- MPI\_Init() is called for initializing an MPI environment.
   MPI\_Init() must be called once and only once before calling any other MPI calls.
   MPI\_Finalize() terminates MPI processing and no other MPI call can be made afterwards.
   Attention: Including code before MPI\_Init() or after MPI\_Finalize() can produce nonportable programs
- The method MPI\_Comm\_size() returns the number of processes belonging to the MPI\_COMM\_WORLD object. A communicator is an identifier associated with a group of processes. MPI\_COMM\_WORLD defined in mpi.h represents the group consisting of all the processes participating in the parallel job.
- Each process in a communicator has its unique rank, which is in the
  range [0,size-1] where size is the number of processes in that
  communicator. A process can have different ranks in each
  communicator that the process belongs to. MPI\_Comm\_rank() returns
  the rank of the process within the communicator given as the first
  argument.

- Although each process executes the same program in the SPMD model, you can make the behaviour of each process different by using the value of the rank.
- This is where the parallel speed-up comes from; each process can operate on a different part of the data or the code concurrently.
- Let's see how it works: copy&paste into example 1.cpp , then ...
  - Load MPI module (1 time): module load mpi/mpich-3.2-x86 64
  - Compile: mpicxx example 1.cpp
  - Run on 4 processors: mpiexec -np 4 a.out (on Python: mpiexec -np 4 python example 1.py )

#### Typical result:

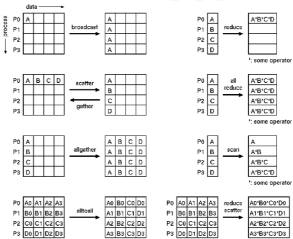
```
SHELL$ mpiexec -np 4 a.out
Sono il nodo 0 dei 4 che hai utilizzato!
Sono il nodo 1 dei 4 che hai utilizzato!
Sono il nodo 2 dei 4 che hai utilizzato!
Sono il nodo 3 dei 4 che hai utilizzato!
```

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 Collective communication allows you to exchange data among a group of processes.

The communicator argument in the collective communication subroutine calls specifies which processes are involved in the communication. In other words, all the processes belonging to that communicator must call the same collective communication subroutine with matching arguments.



All of the MPI collective communication subroutines are blocking i.e. they ends only when all the involved processes have completed their task. There are several types of collective communications, as illustrated in the picture.

#### MPI BCAST

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• The subroutine MPI\_Bcast() broadcasts the message from a specific process called *root* to all the other processes in the communicator given as an argument.

```
Python
 from mpi4py import MPI
                                                                                             #include "mpi.h"
 import numpy as np
                                                                                            #include <iostream>
                                                                                            using namespace std;
 comm = MPI.COMM_WORLD
size = comm.Get_size()
rank = comm.Get_rank()
                                                                                            int main(int argc, char* argv[])
my values = np.zeros(3)
                                                                                            MPI Init (&argc, &argv);
                                                                                            MPI_Comm_size(MPI_COMM_WORLD, &size);
MPI_Comm_rank(MPI_COMM_WORLD, &rank);
for i in range(3):
   if rank == 0:
        my_values[i] = i+1
                                                                                            int my_values[3];
for(int i=0;i<3;i++)</pre>
print('Prima: ', my_values[0], ' ', my_values[1], ' ',
my_values[2], ' per il processo: ', rank)
                                                                                             if(rank==0) my values[i]=i+1;
                                                                                            cout<< "Prima: "<< my_values[0]<< " "<< my_values[1]<<
 " "<< my_values[2]<< " per il processo "<< rank<< endl;
 comm.Bcast(my_values, root=0)
                                                                                            MPI_Boast(my_values,3,MPI_INTEGER,0, MPI_COMM_WORLD);
cout<< "Dopo: "<< my_values[0]<< " "<< my_values[1]<< "
"<< my_values[2]<< " per il processo "<< rank<< endl;</pre>
print('Dopo: ', my_values[0], ' ', my_values[1], ' ',
my_values[2], ' per il processo: ', rank)
                                                                                             return 0;
```

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## MPI data types

- Descriptions of MPI data types and communication buffers follow.
- MPI subroutines recognize data types as specified in the MPI standard. The following is a description of MPI data types in the Fortran language bindings.
- Make sure that the amount of data transmitted matches between the sending process and the receiving processes.

| MPI Data Types                       | Description (Fortran Bindings)            |
|--------------------------------------|---|
| MPI_INTEGER1                         | 1-byte integer                            |
| MPI_INTEGER2                         | 2-byte integer                            |
| MPI_INTEGER4, MPI_INTEGER            | 4-byte integer                            |
| MPI_REAL4, MPI_REAL                  | 4-byte floating point                     |
| MPI_REAL8, MPI_DOUBLE_PRECISION      | 8-byte floating point                     |
| MPI_REAL16                           | 16-byte floating point                    |
| MPI_COMPLEX8, MPI_COMPLEX            | 4-byte float real, 4-byte float imaginary |
| MPI_COMPLEX16,<br>MPI_DOUBLE_COMPLEX | 8-byte float real, 8-byte float imaginary |

| MPI Data Types            | Description (Fortran Bindings)              |
|---------------------------|---|
| MPI_COMPLEX32             | 16-byte float real, 16-byte float imaginary |
| MPI_LOGICAL1              | 1-byte logical                              |
| MPI_LOGICAL2              | 2-byte logical                              |
| MPI_LOGICAL4, MPI_LOGICAL | 4-byte logical                              |
| MPI_CHARACTER             | 1-byte character                            |
| MPI_BYTE, MPI_PACKED      | N/A   |

#### MPI GATHER

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- The subroutine MPI\_Gather() transmits data from all the processes in the communicator to a single receiving process. The length of the message sent from each process must be the same.
- The memory locations of the send buffer (isend) and the receive buffer (irecv)
  must not overlap.

```
C++
                            Python
from mpi4py import MPI
                                                                         #include "mpi.h
import numpy as np
                                                                         using namespace std;
comm = MPI.COMM_WORLD
                                                                         int main(int argc, char* argv[])
size = comm.Get size()
rank = comm.Get_rank()
                                                                         MPI_Init(&argc,&argv);
                                                                         MPI_Comm_size(MPI_COMM_WORLD, &size);
MPI_Comm_rank(MPI_COMM_WORLD, &rank);
if(size>3){cout<<"Hai scelto troppi processi"<<endl;
   exit('Hai scelto troppi processi')
irecv = np.zeros(3)
                                                                                      return 1;}
                                                                         int irecv[3];
for(int i=0;i<3;i++) irecv[i]=0;</pre>
isend = np.zeros(1)
isend[0] = rank+1
                                                                         int isend = rank + 1;
comm.Gather(isend, irecv, root=0)
                                                                         MPI_Gather(&isend,1,MPI_INTEGER,irecv,1,MPI_INTEGER,0,
print('irecv: ', irecv[0], ' ', irecv[1], ' ',
irecv[2])
                                                                         MPI COMM WORLD);
                                                                         if(rank==0) cout<< "irecv: " <<irecv[0] <<" "
<<irecv[1] <<" " <<irecv[2] <<end1;</pre>
                                                                         MPI Finalize();
                                                                         return 0;
```

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#### MPI REDUCE

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 The subroutine MPI\_Reduce() does reduction operations such as summation of data distributed over processes, and brings the result to the root process.

```
Python
                                                                                                   C++
from mpi4py import MPI
import numpy as np
                                                                     #include <iostream>
                                                                     using namespace std;
comm = MPI.COMM_WORLD
                                                                     int main(int argc, char* argv[])
size = comm.Get_size()
rank = comm.Get_rank()
                                                                    MPI_Init(&argc,&argv);
MPI_Comm_size(MPI_COMM_WORLD, &size);
MPI_Comm_rank(MPI_COMM_WORLD, &rank);
           = np.zeros(1)
Prod
         = np.zeros(1)
isend = np.zeros(2)
                                                                     int isend[2].irecv[2];
                                                                     for(int i=0;i<2;i++) isend[i]=rank+i+1;</pre>
  isend[i] = rank+i+1
                                                                     MPI Reduce (&isend[0], &irecv[0], 1, MPI INTEGER,
                                                                    MPI_SUM, 0, MPI_COMM_WORLD);
MPI_Reduce(&isend[1], &irecv[1], 1, MPI_INTEGER,
comm.Reduce(isend[0], Sum, op=MPI.SUM, root=0)
comm.Reduce(isend[1], Prod, op=MPI.PROD, root=0)
                                                                                 MPI_PROD, 0, MPI_COMM_WORLD);
  print('Sum: ', Sum, 'Product: ', Prod)
                                                                              cout<<"irecv: "<<irecv[0]<<" "<<irecv[1]<<endl;
                                                                     MPI Finalize():
                                                                     return 0;
```

#### MPI REDUCE

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- As is the case with MPI\_Gather(), the send buffer and the receive buffer cannot overlap in memory.
- The fifth argument of MPI\_Reduce(), MPI\_SUM, specifies which reduction operation to use. The MPI provides several common operators by default, where MPI\_SUM is one of them, which are defined in mpif.h

|     | Operation   | Data type   |
|-----|---|---|
| 4   | MPI_SUM (sum),<br>MPI_PROD (product)  | MPI_INTEGER, MPI_REAL,<br>MPI_DOUBLE_PRECISION, MPI_COMPLEX |
|     | MPI_MAX (maximum),<br>MPI_MIN (minimum)   | MPI_INTEGER, MPI_REAL,<br>MPI_DOUBLE_PRECISION              |
| 100 | MPI_MAXLOC (max value<br>and location),<br>MPI_MINLOC (min value<br>and location) | MPI_2INTEGER,<br>MPI_2REAL,<br>MPI_2DOUBLE_PRECISION        |
|     | MPI_LAND (logical AND),<br>MPI_LOR (logical OR),<br>MPI_LXOR (logical XOR)        | MPI_LOGICAL   |
|     | MPI_BAND (bitwise AND),<br>MPI_BOR (bitwise OR),<br>MPI_BXOR (bitwise XOR)        | MPI_INTEGER,<br>MPI_BYTE                                    |

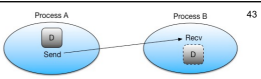
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#### Managing groups: MPI\_COMM\_SPLIT

It is possible that a problem consists of several sub-problems and each problem
can be solved concurrently. This type of application can be found in the category
of coupled analysis. MPI allows you to create a new group as a subset of an
existing group. Specifically, use MPI\_COMM\_SPLIT for this purpose.

```
Python
from mpi4py import MPI
                                                                           #include "mpi.h"
import numpy as np
                                                                           using namespace std;
int main(int argc, char* argv[])
comm = MPI.COMM WORLD
size = comm.Get_size()
rank = comm.Get_rank()
  exit('Servono 4 processi, non ',size,' !!!')
                                                                            MPI_Init(&argc,&argv);
                                                                           MPI_Comm_size(MPI_COMM_WORLD, &size);
MPI_Comm_rank(MPI_COMM_WORLD, &rank);
if rank == 0:
ikey = 2
if rank == 1:
                                                                            if(size!=4){cout<<"Servono 4 processi, non
"<<size<<"!!"<<endl; return 1;}</pre>
                                                                            int icolor, ikey;
  icolor = 1
                                                                            if (rank==0) {icolor=1;ikey=2;}
if rank == 2:
                                                                            if (rank==1) {icolor=1;ikey=1;}
                                                                            if (rank==2) {icolor=2;ikey=1;}
  icolor = 2
   ikey = 1
                                                                            if(rank==3){icolor=2;ikev=2;}
if rank == 3:
                                                                            MPI_Comm_split(MPI_COMM_WORLD,icolor,ikey,&nuovocom);
                                                                            int newsize, newrank;
   ikev = 2
                                                                            MPI_Comm_size(nuovocom, &newsize);
                                                                           MPI_Comm_rank(nuovocom, &newrank);
cout<<"Ero: "<<rank<<" di "<<size<<" ... e adesso sono:
"<<newrank<<" di "<<newsize<<endl;</pre>
newsize = newcomm.Get_size()
newsize - newcomm.Get_rank()
newrank = newcomm.Get_rank()
rank, ' di ',
 orint('Ero: ', rank, ' di ', size, ' ... e adesso sono: ', newrank, ' di ', newsize)
                                                                           MPI Finalize();
                                                                            return 0;
```

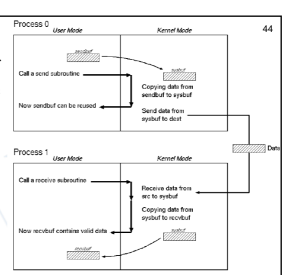
# Point to point communications



- When you use point-to-point communication subroutines, you should know about the basic notions of blocking and non-blocking communication, as well as the issue of deadlocks.
- When a message is sent from process 0 to process 1, there are several steps involved in the communication. At the sending process, the following events occur one after another.
  - 1. The data is copied to the *user buffer* by the user.
  - 2. The user calls one of the MPI send subroutines.
  - 3. The system copies the data from the user buffer to the system buffer.
  - 4. The system sends the data from the system buffer to the destination process.
- The term *user buffer* means scalar variables or arrays used in the program. The following occurs during the receiving process:
  - The user calls one of the MPI receive subroutines.
  - 2. The system receives the data from the source process and copies it to the system buffer.
  - 3. The system copies the data from the system buffer to the user buffer.
  - 4. The user uses the data in the user buffer.

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- When you send data, you cannot or should not reuse your data from user buffer to the system buffer. Also when you receive data, the data is not ready until the system completes copying data from a system buffer to a user buffer.
- In MPI, there are two modes of communication: blocking and non-blocking. When you use blocking communication subroutines such as MPI\_SEND and MPI\_RECV, the program will not return from the



subroutine call until the copy to/from the system buffer has finished. On the other hand, when you use non-blocking communication subroutines such as MPI\_ISEND and MPI\_IRECV, the program immediately returns from the subroutine call. That is, a call to a non-blocking subroutine only indicates that the copy to/from the system buffer is initiated and it is not assured hat the copy has completed.

- Therefore, you have to make sure of the completion of the copy by MPI\_WAIT. If you use your buffer before the copy completes, incorrect data may be copied to the system buffer (in case of nonblocking send), or your buffer does not contain what you want (in case of non-blocking receive).
- Why do you use non-blocking communication despite its complexity? Because non-blocking communication is generally faster than its corresponding blocking communication. Some hardware may have separate co-processors that are dedicated to communication. On such hardware, you may be able to hide the latency by computation. In other words, you can do other computations while the system is copying data back and forth between user and system buffers.
- At the simplest level one can have unidirectional communications or bidirectional communications
- When two processes need to exchange data with each other, you have to be careful about deadlocks. When a deadlock occurs, processes involved in the deadlock will not proceed any further. Deadlocks can take place either due to the incorrect order of send and receive, or due to the limited size of the system buffer.

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#### MPI SEND / MPI RECV

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 When you send a unidirectional message, there are 4 combinations of MPI subroutines to choose from depending on whether you use a blocking or non-blocking subroutine for sending or receiving data (the example show usage of blocking subroutines).

```
Python
                                                                                             C++
from mpi4py import MPI
                                                                 #include "mpi.h"
                                                                 #include <iostream>
import numpy as np
                                                                 using namespace std;
comm = MPI.COMM WORLD
                                                                 int main(int argc, char* argv[])
size = comm.Get_size()
rank = comm.Get_rank()
                                                                 int size, rank;
                                                                 MPI Init (&argc, &argv);
                                                                 MPI_Comm_size(MPI_COMM_WORLD, &size);
MPI_Comm_rank(MPI_COMM_WORLD, &rank);
imesg[0] = rank
if rank == 1:
  comm.Send(imesg, dest=0)
                                                                 int itag=1;
elif rank == 0:
                                                                 int imesg = rank;
if(rank==1)
  comm.Recv(imesg, source=1)
                                                                   MPI_Send(&imesg,1,MPI_INTEGER,0,itag,MPI_COMM_WORLD);
print('Io sono: ', rank,' Il mio messaggio: ', imesg)
                                                                 else if(rank==0)
                                                                  MPI_Recv(&imesg,1,MPI_INTEGER,1,itag,MPI_COMM_WORLD,
                                                                            &stat);
                                                                 cout<<"messaggio = "<<imesg<<endl;
                                                                 MPI_Finalize();
                                                                 return 0;
```

#### Bidirectional communications

- Case 1 Both processes call send, and then receive (dangerous if blocking send)
- Case 2 Both processes call receive, and then send (really dangerous!)
- Case 3 One process calls send and receive subroutines in this order, and the
  other calls in the opposite order (safe)

```
Python
                                                                                                    C++
                                                                     #include "mpi.h"
from mpi4py import MPI
                                                                     #include <iostream>
import numpy as np
                                                                      using namespace std;
                                                                     const int n = 100; // try to increase n
int main(int argc, char* argv[]) {
comm = MPI.COMM WORLD
size = comm.Get_size()
rank = comm.Get_rank()
                                                                     int size, rank;
                                                                     MPI Init (&argc, &argv);
n = 100 # try to increase n
                                                                      MPI_Comm_size(MPI_COMM_WORLD, &size);
imesg = np.zeros(n)
imesg2 = np.zeros(n)
                                                                     MPI Comm rank (MPI COMM WORLD, &rank);
                                                                     MPI_Status stat1, stat2;
                                                                     int* imesq = new int[n]; int* imesq2 = new int[n];
for i in range(n):
                                                                      int itag=1; int itag2=2;
   imesg[i] = rank
                                                                     for(int i=0;i<n;i++){imesg[i]=rank; imesg2[i]=rank+1;}</pre>
   imesg2[i] = rank+1
                                                                     if (rank==1) {MPI Send(&imesg[0],n,
                                                                                   MPI_INTEGER, 0, itag, MPI_COMM_WORLD);
if rank == 1:
                                                                                   MPI Recv(&imesg2[0],n,
                                                                                   MPI_INTEGER,0,itag2, MPI_COMM_WORLD,&stat2);
cout<<"messaggio = "<<imesg2[0]<<endl;}</pre>
   comm.Send(imesg, dest=0)
   comm.Recv(imesg2, source=0)
print('Io sono: ', rank,' Il mio messaggio: ',
imesg2[0])
                                                                     else if(rank==0){MPI_Send(&imesg2[0],n,
                                                                                         MPI_INTEGER,1,itag2, MPI_COMM_WORLD);
                                                                                          MPI_Recv(&imesg[0],n,MPI_INTEGER,1,
   comm.Send(imesg2, dest=1)
                                                                                         itag, MPI_COMM_WORLD,&stat1);
cout<<"messaggio = "<<imess[0]<<endl;}</pre>
   comm.Recv(imesg, source=1)
print('Io sono: ', rank,' Il mio messaggio: ',
                                                                     MPI Finalize();
imesa[01)
```

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The following code is free from deadlock because the program immediately returns from MPI ISEND and starts receiving data from the other process.

```
Python
                                                                                                 C++
                                                                   #include "mpi.h"
from mpi4py import MPI
import numpy as np
                                                                   using namespace std;
const int n = 100; // try to increase n
 comm = MPI.COMM_WORLD
size = comm.Get size()
                                                                   int main(int argc, char* argv[]) {
rank = comm.Get_rank()
                                                                    int size, rank;
                                                                   MPI_Init(&argc,&argv);
                                                                   MPI_Comm_size(MPI COMM WORLD, &size);
n = 100 # try to increase n
imesg = np.zeros(n)
imesg2 = np.zeros(n)
                                                                    MPI_Comm_rank(MPI_COMM_WORLD, &rank);
                                                                    MPI Status stat1, stat2;
 for i in range(n):
   imesg[i] = rank
   imesg2[i] = rank+1
                                                                   int* imesg = new int[n]; int* imesg2 = new int[n];
                                                                    int itag=1; int itag2=2;
                                                                    for(int i=0;i<n;i++){imesg[i]=rank; imesg2[i]=rank+1;}</pre>
   req = comm.Isend(imesg, dest=0)
                                                                   if(rank==1) {MPI_Isend(&imesg[0],n,
                                                                                MPI_INTEGER,0,itag, MPI_COMM_WORLD,&req);
MPI_Recv(&imesg2[0],n,MPI_INTEGER,0,itag2,
   comm.Recv(imesg2, source=0)
        nt('Io sono: ', rank,' Il mio messaggio: ',
                                                                                 MPI COMM WORLD, &stat2);
imesq2[0])
                                                                                 cout<<"messaggio = "<<imesg2[0]<<endl;}
elif rank == 0:
                                                                   else if (rank==0) {MPI Send(&imesg2[0],n,
   comm.Send(imesg2, dest=1)
                                                                                      MPI INTEGER, 1, itag2, MPI COMM WORLD);
comm.Recv(imesg, source=1)
print('Io sono: ', rank,' Il mio messaggio: ',
imesg[0])
                                                                                              cv(&imesg[0],n,MPI_INTEGER,
                                                                                       1,itag,MPI_COMM_WORLD, &stat1);
cout<<"messaggio = "<<imess[0]<<endl;}</pre>
                                                                   MPI Finalize();
```

#### MPI WTIME

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 Measurement of time and thus of performances can be obtained as in the following example

```
C++
                            Python
from mpi4py import MPI
                                                                         #include "mpi.h'
import numpy as np
                                                                         #include <iostream>
                                                                         using namespace std;
                                                                         int main(int argc, char* argv[]){
size = comm.Get_size()
rank = comm.Get_rank()
                                                                         int size, rank;
                                                                         MPI_Init(&argc,&argv);
                                                                        MPI_Comm_size(MPI_COMM_WORLD, &size);
MPI_Comm_rank(MPI_COMM_WORLD, &rank);
tstart = MPI.Wtime()
                                                                         MPI_Status stat;
imesg = np.zeros(n)
                                                                         double tstart = MPI_Wtime();
Sum = 0
                                                                        int n = 100;
int* imesg = new int[n];
 for i in range(n):
   imesg[i] = rank
   if rank == 1:
                                                                        int sum=0;
for(int i=0;i<n;i++){</pre>
                                                                           imesg[i]=rank;
                                                                          comm.Send(imesg, dest=0)
   elif rank == 0:
comm.Recv(imesg, source=1)
                                                                        double tend = MFI Wtime();
double dt = tend - tstart;
cout<<"io sono "<<rank<<"; somma = "<<sum<<"; tempo =
"<<dt<<end1;</pre>
tend = MPI.Wtime()
 print('Io sono:', rank, 'Ho implegato:', tend-tstart,
'secondi')
                                                                         MPI_Finalize();
                                                                         return 0;}
```

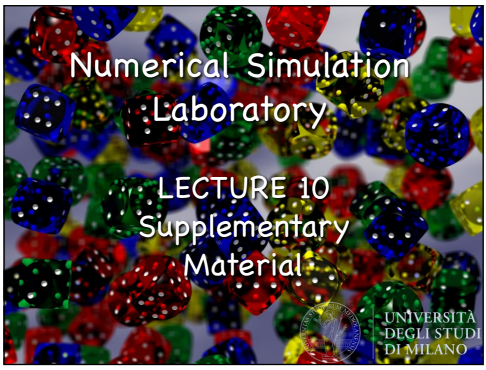
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#### Lecture 10: Suggested books

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- Gropp, Lusk, Skjellum, Using MPI, Portable Parallel Programming with the Message-Passing Interface - The MIT Press (2014)
- https://hpc-tutorials.llnl.gov/mpi/





#### Parallelizing do-loops

- In almost all of the scientific and technical programs, the hot spots are likely to be found in DO loops. Thus parallelizing DO loops is one of the most important challenges when you parallelize your program.
- The basic technique of parallelizing DO loops is to distribute iterations among processes and to let each process do its portion in parallel.
- Usually, the computations within a DO loop involves arrays whose indices are associated with the loop variable. Therefore distributing iterations can often be regarded as dividing arrays and assigning chunks (and computations associated with them) to processes.
- In block distribution, the iterations are divided into p parts, where p is the number of processes to be executed in parallel. The iterations in each part is consecutive in terms of the loop variable.
- In cyclic distribution, the iterations are assigned to processes in a round-robin fashion. Note that, in general, the cyclic distribution incurs more cache misses than the block distribution because of non-unit stride access to matrices within a loop.
- When you use some redistribution, it is often sufficient for each process to have only a part of the arrays and to do the computation associated with that part. If a process needs data that another process holds, there has to be some transmission of the data between the processes.

#### Block distribution

- ullet Suppose when you divide n by p, the quotient is q and the remainder is r.
- For example, in the case of n = 14 and p = 4, q is 3 and r is 2. One way to distribute iterations in such cases is as follows. Processes 0..r-1 are assigned q+1 iterations each. The other (p-r) processes are assigned q iterations. This distribution corresponds to expressing n as n=r(q+1)+(p-r)q

```
void blocchi(int nstart, int nend, int nprocs, int
mype, int& istart, int& iend)
#include <iostream>
using namespace std;
const int n=1000000;
                                                                                 int il= nend - nstart +1;
                                                                                int i1 = int = int i1
int i2 = i1/nprocs;
int i3 = i1 % nprocs;
int min = mype;
void blocchi(int,int,int,int,int&,int&);
int main(int argc, char* argv[])
                                                                                 if(i3<mype)
          int size, rank;
         MPI_Init(&argc,&argv);
MPI_Comm_size(MPI_COMM_WORLD, &size);
MPI_Comm_rank(MPI_COMM_WORLD, &rank);
                                                                                 min=i3;
istart = mype*i2 + nstart + min;
                                                                                 iend = istart + i2 -1;
                                                                                 if(i3>mype)
         int* a = new int[n];
          int sum=0, ssum;
                                                                                           iend++;
         int istart, iend;
         blocchi(0,n-1,size,rank,istart,iend);
          for(int i=istart;i<iend;i++){
                  a[i]=i;
                                                                                         In the example above:
MPI Reduce (&sum, &ssum, 1, MPI INTEGER, MPI SUM, 0, MPI COMM
                                                                              Rank 0 0 0 0 1 1 1 1 2 2 3 3 3 3
         if(rank==0) cout<<"somma = "<<ssum<<endl;</pre>
         MPI_Finalize();
         return 0;
```

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## Cyclic distribution

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- To avoid load unbalance when workload is not uniform among iterations one can try cyclic distribution
- The following is an example of how to implement it
- In the previous example the distribution corresponds to:

```
        teration
        1
        2
        3
        4
        5
        6
        7
        8
        9
        10
        11
        12
        13
        14

        Rank
        0
        1
        2
        3
        0
        1
        2
        3
        0
        1
        2
        3
        0
        1
```

```
#include "mpi.h"
#include <iostream>
using namespace std;

const int n = 10000;
int main(int argc, char* argv[])
{
   int size, rank;
        MPI_Init(&argc,&argv);
        MPI_Comm_size(MPI_COMM_WORLD, &size);
        MPI_Comm_rank(MPI_COMM_WORLD, &rank);

   int* a = new int[n];
   int sum=0,ssum;
   for(int i=rank;i<n;i+=size)
   {
        a[i]=i+1;
        sum += i+1;
   }

MPI_Reduce(&sum,&ssum,1,MPI_INTEGER,MPI_SUM,0,MPI_COMM_WORLD);

   if(rank==0) cout<<"somma = "<<ssum<<end1;
        MPI_Finalize();
        return 0;
}</pre>
```