# Lab Guide 9

# **MPI-Based Parallelization**

## **Objective:**

- Program parallelization with distributed memory programming (message passing)

#### Introduction

This lab session aims to apply the basic MPI communication concepts studied in the previous session to parallelize simple applications.

Copy the /share/cpar/P09 Codigo folder to your home directory in the SeARCH cluster.

Compile the program in the cluster frontend using the mpic++-02-oprimes PrimeMain.cpp command. Use the sbatch primes mpi.sh command to run the application.

The primes mpi.sh file should specify the required resources and should run the MPI application.

The following example requests three PUs and spawns three MPI processes:

```
[search7edu]$ cat primes_mpi.sh
#!/bin/bash
#SBATCH --time=1:00
#SBATCH --ntasks=3
#SBATCH --partition=cpar
mpirun -np 3 ./primes
```

The number of requested resources (--ntasks) must be the same as the number of processes (-np) used in the mpirun command.

## Exercise 1 – Prime calculation using the Sieve of Eratosthenes

Consider the following sequential program, which finds all prime numbers up to a given MAXP:

```
int MAXP = 1000000;
int SMAXP = 1000;
int pack=MAXP/10;
PrimeServer *ps1 = new PrimeServer();
PrimeServer *ps2 = new PrimeServer();
PrimeServer *ps3 = new PrimeServer();
ps1->minitFilter(1,SMAXP/3,SMAXP);
ps2->minitFilter(SMAXP/3+1,2*SMAXP/3,SMAXP);
ps3->minitFilter(2*SMAXP/3+1,SMAXP,SMAXP);
int *ar = new int[pack/2];
for(int i=0; i<10; i++) {
    generate(i*pack, (i+1)*pack, ar);
    ps1->mprocess(ar,pack/2);
    ps2->mprocess(ar,pack/2);
    ps3->mprocess(ar,pack/2);
ps3->end();
```

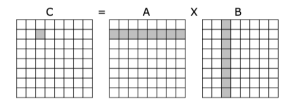
João Sobral, nov'22

- a) Parallelize the code using MPI through the implementation of a pipeline of processes that receives an array of integers, created by the generate function, and each process filters out a subset of the input. The mprocess method implements the filtering of the primes, and end prints the final amount of primes found. This pipeline should have 3 processes, one for each instance of PrimeServer performing the filtering.
- **b)** (\*) Modify the parallelization implemented in **a)** to work with an arbitrary number of processes and messages.
- c) (\*) Parallelize the sequential application through the implementation of a farm of processes behaving in a "work sharing" paradigm with dynamic scheduling.

# Exercise 2 - Distributed memory matrix multiplication with MPI collective operations

Download the matrix multiplication code from lab session 0 (base DOT version in the figure).

foreach line of A 
$$\text{foreach column of B}$$
 
$$\text{Cline,column = DOTline of A,column of B}$$
 
$$\overline{C_{ij} = DOT_{linha\_A_i,coluna\_B_j} = \sum_{k=0}^{n-1} (A_{ik} * B_{kj}) }$$



- a) Identify a strategy to implement the matrix multiplication with MPI:
  - 1. Each process will compute a subset of the elements of matrix C. Identify the main alternatives to partition the computation of this matrix among processes.
  - 2. Select the simplest alternative to partition the computations of C. What data from matrix A and matrix B is required to perform the computation of each part of C?
- b) Implement the selected approach using MPI collective operations. Relevant collection operations are:
  - 1. **Broadcast** the data from root to all other processes:

int MPI\_Bcast(void\* buffer, int count, MPI\_Datatype
datatype, int root, MPI Comm comm)

2. Scatter: scatters data from root into all other processes

int MPI\_Scatter(void\* sbuf, int scount, MPI\_Datatype
stype, void\* rbuf, int rcount, MPI\_Datatype rtype, int
root, MPI\_Comm comm )



MPI Bcast

3. **Gather**: Joints data from all processes into the root

int MPI Gather(/\* same signature of Scatter\*/)

