CPDS Laboratory Assignment Laboratory Assignment 0: Parallel Execution Environment

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Deliverable

Session 1

Experimental setup

The objective of this first assignment is to familiarise yourself with the hardware and software environment that you will be using during the semester to do all laboratory assignments in CPDS. From your PC/terminal booted with Linux¹ you will access boada, a multiprocessor server located at the Computer Architecture Department at UPC. From a machine within the UPC network you will be able to access boada directly. However, to be able to access the boada server from a computer outside the UPC you will need to use a UPC VPN connection. To connect to it you will have to establish a connection using the secure shell command: "ssh -X cpds1XYZ@boada.ac.upc.edu", being cpds1XYZ the username assigned to you. Option -X is necessary in order to forward the X11 data and be able to open remote windows in your local desktop². You should change the password for your account using "ssh -t cpds1XYZ@boada.ac.upc.edu passwd"³.

Once you are logged in you will find yourself in any of the interactive nodes:boada-6 to boada-8, where you can execute interactive jobs and from where you can submit execution jobs to the rest of the nodes in the machine. In fact, boada is composed of several nodes (named boada-1 to boada-15), equipped with five different processor generations, as shown in the following table:

Node name	Processor generation	Interactive	Partition
boada-1 to 4	Intel Xeon E5645	No	execution2
boada-6 to 8	Intel Xeon E5-2609 v4	Yes	interactive
boada-9	Intel Xeon E5-1620 v4 + Nvidia K40c	No	cuda9
boada-10	Intel Xeon Silver 4314 + 4 x Nvidia GeForce RTX 3080	No	cuda
boada-11 to 14	Intel Xeon Silver 4210R	No	execution
boada-15	Intel Xeon Silver 4210R + ASUS AI CRL-G116U-P3DF	No	iacard

However, in this course you are going to use only nodes boada-6 to boada-8 interactively, nodes boada-11 to boada-14 through the execution queue and, later on, node boada-10 through the cuda queue. Submission of jobs for *batched* execution in a queue is explained in the next section. The rest of the nodes have restricted access and CPDS users are not allowed to send jobs to their corresponding queues (Partitions). Note that the all the nodes within a particular Partition are identical. Therefore, you do not need to worry if the execution was done in node boada-11, boada-12, or any of the other 2 nodes within that partition, when submitting to the execution queue.

All nodes have access to a shared NAS (Network-Attached Storage) disk; you can access it through /scratch/nas/1/cpdsXXYY (in fact this is your home directory, check by typing pwd in the command line). In addition, each node in boada has its own local disk which can be used to store temporary files non visible to other nodes; you can access it through /scratch/1/cpdsXXYY.

¹You can also access from your laptop, booted with Linux, Windows or MacOS X, if a secure shell client is installed.

 $^{^2 \}rm Option$ –Y if you are connecting from a MacOS X laptop with XQuartz. With Windows, if you use putty (<code>https://www.chiark.greenend.org.uk/~sgtatham/putty/latest.html)</code> then you will also need xming: (<code>https://wiki.centos.org/HowTos/Xming</code>). Alternatively, you can use MobaXterm (<code>https://mobaxterm.mobatek.net/download.html</code>) which has everything integrated.

³The passwd command will be executed in boada. After entering the old password correctly twice you will be asked for your new password also twice.

We will post all necessary files to do each laboratory assignment in /scratch/nas/1/cpds0/sessions. For the session today, you need to extract the files from file a0.tar.gz from that location by uncompress it into your home directory in boada.

From the the root of your home directory unpack the files with the following command line: "tar-zxvf /scratch/nas/1/cpds0/sessions/a0.tar.gz".

In order to set up all environment variables you have to process the environment.bash file now available in your home directory with "source ~/environment.bash". Note: since you have to do this every time you login in the account or open a new console window, we recommend that you add this command line in the .bashrc file in your home directory, a file that is executed every time a new session is initiated. If you have unpacked the a0.tar.gz files in your home directory then this file has already been added automatically to your home directory and it will be used from your next connection to boada. From the next time you log in to boada it will be automatically read, and environment variables will be properly set effortless.

In case you need to transfer files from boada to your local machine (laptop or desktop in laboratory room), or viceversa, you have to use the secure copy scp command. For example if you type the following command "scp cpds1XYZ@boada.ac.upc.edu:a0/pi/pi_seq.c ." in your local machine you will be copying the source file pi_seq.c located in directory a0/pi of your home directory in boada to the current directory, represented with the ".", in the local machine, with the same name.

1.1 Node architecture and memory

The first thing you will do is to investigate the architecture of the available nodes in boada. Run sbatch submit-arch.sh command in directory a0/arch. This command will enqueue submit-arch.sh script. More detail about queue execution below. This script will execute the 1scpu and 1stopo commands in order to obtain information about the hardware in one of the nodes of execution queue (boada-11 to 14). The execution of these two commands using submit-arch.sh script generates three files, where number may be 11, 12, 13 or 14: 1) 1scpu-boada-number, 2) 1stopo-boada-number, and map-boada-number.fig. You can use the xfig command to visualise the output file generated (map-boada-number.fig) and export to a different format (PDF or JPG, for example) using File \rightarrow Export in order to include it in your deliverable for this laboratory assignment⁴.

Those files will help you to figure out:

- the number of sockets, cores per socket and threads per core in a specific node;
- the amount of main memory in a specific node, and each NUMAnode;
- the cache memory hierarchy (L1, L2 and L3), private or shared to each core/socket.

Fill in the table in the "Deliverable" section with the main characteristics of the node. Draw the architecture of the node based on the information generated by the tools above. The "--of fig map.fig" option for lstopo can be very useful for that purpose.

1.2 Execution modes: interactive vs queued

There are two ways to execute your programs in boada:

- 1. via a queueing system (in one of the nodes boada-11 to boada-14);
- 2. interactively (in any of the login nodes boada-6 to boada-8). In this case, the system limits the number of cores to be used in parallel executions to two.

⁴In the boada Linux distribution you can use xpdf to open pdf files and display to visualise graphics files. You can also use the "fig2dev -L pdf map.fig map.pdf" command to convert from .fig to .pdf; look for alternative output graphic languages by typing "man fig2dev".

It is mandatory to use option 1 when you want to execute scripts that require several processors in a node, ensuring that your job is executed in isolation (and therefore reporting reliable performance results) and to avoid adding additional load to the interactive node accessed by all users; the execution starts as soon as a node is available. When using option 2 your execution starts immediately but will share resources with other programs and interactive jobs, not ensuring representative timing results. Usually, scripts for both options (submit-xxxx.sh and run-xxxx.sh, respectively) will be provided:

- Queueing a job for execution: "sbatch [-p partition] ./submit-xxxx.sh" . Additional parameters may be specified, if needed by the script, after the script name. If you do not specify the name of the partition with "-p partition" your script will run on the execution partition by default. Use "squeue" to ask the system about the status of your job submission. You can use "scancel" followed by the job identifier to remove a job from the queueing system. Note that partition names associated to each node name are shown in the last column of the table above. After the execution in an available node associated to the specified partition, in addition to the files being generated by the script, two additional files will be created. Their name will have the script name followed by an ".e" and an ".o" and the job identifier. They will contain the messages sent to the standard error and standard output respectively during the execution of the job. You should check them to be sure results make sense.
- Interactive execution: ./run-xxxx.sh. Additional parameters may be specified after the script name. Jobs interactively executed have a short time limit to be executed.

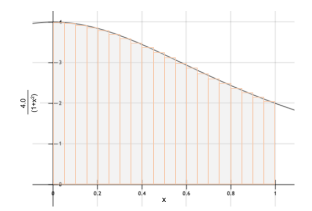
1.3 Serial compilation and execution

Next you will get familiar with the compilation and execution steps for both sequential and parallel applications. For this, we will use some codes which compute digits of number Pi. As a reference, Figure 1.1 shows a python code which computes the area under one quadrant of the unit circle, i.e. a circle of radius equal to 1, defined by the curve $y = \sqrt{1-x^2}$.

```
def midpoint_riemann_sum():
   N = 1000000000
   delta_x = (1 - 0) / N
   x = 0
   pi = 0
   start = time.time()
   while x < (1-delta_x):
       x_next = x + delta_x
       x_mid = (x_next + x) / 2
       f_x = math.sqrt(1 - math.pow(x_mid, 2))
       pi += f_x * delta_x
       x += delta_x
   pi = 4 * pi
   end = time.time()
   print(end - start)
   print("Estimate: " + str(pi))
   print("Actual: " + str(math.pi))
```

Figure 1.1: Python code for computing Pi

The execution time for this code on boada-1 took 522.1 seconds to compute the estimate 3.14159266367805 (actual value is 3.141592653589793). As you can see, it took a long time while it only commputed 7 correct digits of Pi. We aim at estimating digits of Pi much faster. For that we are going to use some C codes which we will parallelize. You are going to use a very simple code, pi_seq.c, which you can find inside the a0/OpenMP/pi directory. pi_seq.c performs the computation of the Pi number by computing the integral of the equation in Figure 1.2. This time the algorithm is using the derivative of



Mathematically, we know that:

$$\int_{0}^{1} \frac{4.0}{(1+x^2)} dx = \pi$$

We can approximate the integral as a sum of rectangles:

$$\sum_{i=0}^{N} F(x_i) \Delta x \approx \pi$$

Where each rectangle has width Δx and height $F(x_i)$ at the middle of interval i.

Figure 1.2: One way to compute Pi

the arctangent function of x, which is equal to 1 divided by $(1+x^2)$ which is another way to estimate Pi. Again, the equation can be solved by computing the area defined by the function, which at its turn can be approximated by dividing the area into small rectangles and adding up its area. This is one of the many ways to estimate digits of number Pi. Both algorithms above multiply by 4 because the functions compute $\pi/4$. Figure 1.3 shows a simplified version of the code you have in pi_seq.c. Variable num_steps defines the number of rectangles, whose area is computed in each iteration of the i loop.

Figure 1.3: Serial code for Pi

Figure 1.4 shows the compilation and execution flow for a sequential program. We will always compile programs to generate binary executable files through a Makefile, with multiple targets that specify the rules to compile each program version. In this course we are going to use gcc (the C front—end from the GNU Compiler Collection).

In the following steps you will compile pi_seq.c using the Makefile and execute it interactively and through the queueing system, with the appropriate timing commands to measure its execution time:

- Open the Makefile file, identify the target you have to use to compile the sequential code. Observe
 how the compiler is invoked. Execute the command line "make target_identified" in order to
 generate the binary executable file. Alternatively, you can just type "make" which will compile all
 programs.
- 2. Interactively execute the binary generated to compute number pi sequentially by doing 1.000.000.000.000 iterations using the run-seq.sh script ("run-seq.sh pi_seq 100000000") which returns the user and system CPU time, the elapsed time, also called wall time, and the % of CPU used (using the GNU time program in /usr/bin/time). In addition, the program itself also reports the elapsed execution time using gettimeofday. Look at the source code and identify the function invocations and data structures required to measure execution time. Note that CPU time is the quantity of

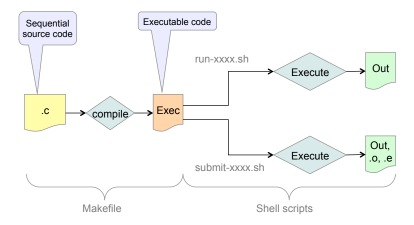


Figure 1.4: Compilation and execution flow for sequential program.

processor time taken by the process. This does not indicate duration. Elapsed time represents the total duration of the execution. If we exploit parallelism, elapsed time can be less than the total CPU time because several cores are used to do the work and it can get done faster.

3. Submit the execution to the queueing system using the "sbatch submit-seq.sh" command and use "squeue" to see that your script is running. Look at submit-seq.sh script and the results generated (the standard output and error of the script and the time-pi_seq-boada{11-14} file).

1.4 Compilation and execution of OpenMP programs

In this section we are going to use OpenMP, the standard for parallel programming using shared-memory, to express parallelism in the C programming language. Note that OpenMP will be explained in detail in the second part of the course, and the goal now is to have a first contact and observe that we can speed-up the execution of our programs. Therefore, in this section we will see how to compile and execute parallel programs in OpenMP and we will observe the performance improvements. Figure 1.5 shows the compilation and execution flow for an OpenMP program. The main difference with the flow shown in Figure 1.4 is that now the Makefile will include the appropriate compilation flag to enable OpenMP.

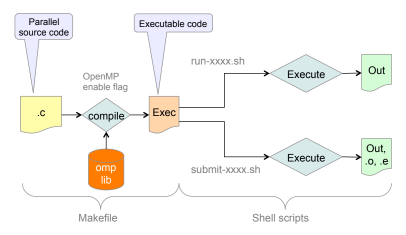


Figure 1.5: Compilation and execution flow for OpenMP.

1.4.1 Compiling OpenMP programs

1. In the same aO/OpenMP/pi directory you will find an OpenMP version of the code for doing the computation of pi in parallel (pi_omp.c). Compile the OpenMP code using the appropriate target

in the Makefile). What is the compiler telling you? Is the compiler issuing a warning or an error message? Is the compiler generating an executable file? The compilation run smoothly because we used flag -fopenmp, but if you remove the -fopenmp flag and force the recompilation you will see the difference.

1.4.2 Executing OpenMP programs

- 1. Interactively execute the OpenMP code with 8 threads (processors) and same number of iterations (1.000.000.000) using the run-omp.sh script ("run-omp.sh pi_omp 1000000000 8"). What is the time command telling you about the user and system CPU time, the elapsed time, and the % of CPU used? Take a look at the script to discover how do we specify the number of threads to use in OpenMP. Note that, since any login node boada-[678] is shared by many users and is reserved for interactive usage, it has been configured so that our processes only have 2 cores available for their execution. Thus, we will see no improvement due to the parallelization.
- 2. Use submit-omp.sh script to queue the execution of the OpenMP code ("sbatch submit-omp.sh") and measure the CPU time, elapsed time and % of CPU when executing the OpenMP program using 8 threads in isolation. Do you see a major difference between interactive and queued execution?

1.4.3 Strong vs. weak scalability

Finally in this section you are going to explore the scalability of the pi_omp.c code when varying the number of threads used to execute the parallel code. To evaluate the scalability the ratio between the sequential and the parallel execution times will be computed. Two different scenarios will be considered: strong and weak scalability.

- In *strong* scalability the number of threads is changed with a fixed problem size. In this case parallelism is used to reduce the execution time of your program.
- In weak scalability the problem size is proportional to the number of threads. In this case parallelism is used to increase the problem size for which your program is executed.

We provide you with two scripts, submit-strong-omp.sh and submit-weak-omp.sh, which should be submitted to the queueing system with the sbatch command. The scripts execute the parallel code using from 1 (np_NMIN) to 20 (np_NMAX) threads. The problem size for strong scalability is 1.000.000.000 iterations; for weak scalability, the initial problem size is 100.000.000 which grows proportionally with the number of threads. The execution will take some time because several executions are done for each test (in order to get a minimum time), please be patient! And, very important, do NOT execute interactively but, instead, submit them for execution in the execution queue. As a result each script generates several files, including a plot (a file with a name ending in .ps in Postscript format) showing the resulting parallel execution time and speed-up or parallel efficiency. Visualise⁵ the plots generated and reason about how the speed-up changes with the number of threads in the two scenarios.

Change the value for np_NMAX in the submit-strong-omp.sh from 20 to 40 and execute again. Can you explain the behaviour observed in the scalability plot? Even when Hyper-Threading is enabled and the system can take instructions quickly from 40 threads, there are only 20 cores available!

1.5 Compiling and Executing MPI programs

1.5.1 Strong scalability with processes and threads

Next, we will test ourselves how to compile and execute parallel programs in MPI, or a combination of both MPI and OpenMP. To do so, follow similar same steps as for OpenMP in the previous section, but working with the files in directory aO/MPI/pi. The code calculates number Pi with a different algorithm based on the so called Monte Carlo method. Monte Carlo methods are a broad class of computational algorithms that rely on repeated random sampling to obtain numerical results. The idea is to simulate

⁵In the boada Linux distribution you can use the ghostscript gs command to visualise Postscript files or convert the files to PDF using the ps2pdf command and use xpdf to visualise PDF files.

random (x, y) points in a 2-D plane with domain as a square of side 1 unit. Imagine a circle inside the same domain with same diameter and inscribed into the square. We then calculate the ratio of number of points that lied inside the circle and total number of generated points. To higher the number of samples, the better the precision obtained.

We can get a plot of the execution time and speed—up that is obtained when varying the number of MPI processes (strong scalability) by submitting a job to the execution queue with sbatch submit-strong-mpi.sh). In addition, we want to observe the combination of MPI and OpenMP. For that execute sbatch submit-mpi2-omp.sh) and study the results presented in the output file created after the execution finishes. Note that since the test uses 2 MPI processes the execution implies using an even number of threads between 2 and 40 even when the output presents the results for 1 to 20 threads: there are between 1 to 20 threads per MPI process; with 2 processes this results in having twice as many threads.

We hope that, with this experiments, you realize that we can speed up the code by parallelizing a code with OpenMP, with MPI, or a combination of both to use several nodes and several cores per node!

Deliverable

After completing this laboratory assignment, and before starting the next one, you will have to deliver a report in PDF format (other formats will not be accepted) containing the answers to the questions stated at the end of this document;

Your professor will open the assignment at the moodle Atenea website and set the appropriate delivery dates for the delivery. Note that, unlike the subsequent assignments which can be done in groups of 2 students, this initial assignment must be done independently by each student, requiring Individual submission through the moodle Atenea website.

Important: In the front cover of the document, please clearly state your name and surname, the identifier of the group (username cpds1XYZ), title of the assignment, date, academic course/semester, ... and any other information you consider necessary.

As part of the document, you can include any code fragment, figure or plot you need to support your explanations. In case you need to transfer files from boada to your local machine (laptop or desktop in laboratory room), or viceversa, you can use the secure copy scp command. For example "scp cpds1XYZ@boada.ac.upc.edu:a0/foobar.txt local/directory/." executed in your local machine to copy file foo.txt inside directory a0 in your home directory of boada to directory local/directory/ with the same name.

Node architecture and memory

1. Complete the following table with the relevant architectural characteristics of the different node types available in boada:

	boada-11 to boada-14
Number of sockets per node	
Number of cores per socket	
Number of threads per core	
Maximum core frequency	
L1-I cache size (per-core)	
L1-D cache size (per-core)	
L2 cache size (per-core)	
Last-level cache size (per-socket)	
Main memory size (per socket)	
Main memory size (per node)	

2. Include in the document the architectural diagram for one of the nodes boada-11 to boada-14.

Timing sequential and parallel executions

For each of the following sections show the plots or tables required. But, in addition, please **provide** some reasoning about the results obtained. For instance, comment whether the scalability is good or not, and why.

- 3. Plot the execution time and speed—up that is obtained when varying the number of threads (strong scalability) by submitting the jobs to the execution queue (section 1.4.3). Show the parallel efficiency obtained when running the weak scaling test. Explain what strong and weak scalability refer to, exemplifying your explanation with the plots that you present. Explain the reasons which cause the behaviour observed when more than 20 threads are used.
- 4. Plot the execution time and speed-up that is obtained when varying the number of MPI processes from 1 to 20 (strong scalability) by submitting the jobs to the execution queue (section 1.5.1). In addition, show in a table the elapsed execution time when executed with 2 MPI processes when varying the number of threads from 1 to 20⁶. Which was the original sequential time?; the time with 20 MPI processes?; and the time with 2 MPI processes each using 20 OpenMP threads? You can retrieve such information for 1 and 20 MPI processes from file elapsed.txt; and for 2 MPI processes each with 20 threads within the output file created after the execution of sbatch submit-mpi2-omp.sh.

⁶Note that for 2 MPI processes the scalability test implies using an even number of threads between 2 and 40.