ItHPC Lab Report

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1. Lab 1: Intro

This lab consists of introductory exercises to MPI. Therefore the answers to each question consist of only of source code, submitted electronically.

2. Lab 2: Poisson's equation

2.1. Part 1

2.1.1. Step 1

It is simple to understand that the program was indeed executed twice since two pairs of statements are written to the terminal in comparison to one pair before the modifications. Since we are now running the same program in two different nodes this behavior is expected.

The result is the following

Number of iterations : 2355

Elapsed processortime: 1.350000 s

Number of iterations : 2355

Elapsed processortime: 1.360000 s

2.1.2. Step 2

After adding the global variable and the necessary call to MPI_Comm_rank using the predefined communicator MPI_COMM_WORLD.

```
MPI_Comm_rank(MPI_COMM_WORLD, &proc_rank);
```

The following is printed to the standard output:

- (0) Elapsed processortime: 1.360000 s
- (0) Number of iterations : 2355
- (1) Elapsed processortime: 1.360000 s
- (1) Number of iterations : 2355

2.1.3. Step 3

2.1.4. Step 4

Adjusting the code so each process writes to a separate file does not affect the text displayed, so there is no need to repeat it here. In addition by executing the command

```
diff output0.dat output1.dat
```

I was able to confirm the files are indeed identical.

2.1.5. Step 5

On this step, responsible to ensure correct distribution of information originated from an input file, several statements had to be rewritten. Below is

a summary of those changes, in particular the parts that were not completely specified in the exercise manual.

To ensure only process 0 opens the file a simple comparison suffices

```
/* only process 0 may execute this if */ if (proc_rank == 0) \{ \dots \}
```

To broadcast the data read from the file it is first necessary to explain which fields the MPI_Bcast(void *buffer, int count, MPI_Datatype datatype, int root, MPI_Comm comm)) function requires.

For our situation the buffer pointer should refer to the address of the variable we want to broadcast. The count relates to the number of entries in the buffer. The datatype should describe the type of data the buffer points to, e.g. for integers this should be MPI_INT. The root is the message broadcaster, in our case node 0. Finally we will use the usual predefined communicator for the last argument comm.

Thus the broadcast calls are as follows

```
/* broadcast the array gridsize in one call */
                       , 2, MPI_INT , 0, MPI COMM WORLD);
   MPI Bcast(&gridsize
   /* broadcast precision goal */
   MPI_Bcast(&precision_goal, 1, MPI_DOUBLE, 0, MPI_COMM_WORLD);
   /* broadcast max iter */
                            , 1, MPI INT , 0, MPI COMM WORLD);
   MPI Bcast(&max iter
(\ldots)
   /* The return value of this scan is broadcast even though it is no
   MPI Bcast(&s, 1, MPI INT, 0, MPI COMM WORLD);
(...)
   /* broadcast source x */
   MPI_Bcast(&source_x , 1, MPI_DOUBLE, 0, MPI_COMM_WORLD);
   /* broadcast source_y */
   MPI_Bcast(&source_y , 1, MPI_DOUBLE, 0, MPI_COMM_WORLD);
   /* broadcast source val */
   MPI Bcast(&source val, 1, MPI DOUBLE, 0, MPI COMM WORLD);
```

2.1.6. Step 6

Following the same approach as in the previous section, only the finished version of incomplete code from the manual will be shown in the excerpt.

```
MPI_Cart_create(MPI_COMM_WORLD, 2, P_grid, wrap_around, reorder, &
    (...)
/* Rank of process in new communicator */
MPI_Comm_rank(grid_comm, &proc_rank);
/* Coordinates of process in new communicator */
MPI_Cart_coords(grid_comm, proc_rank, 2, proc_coord);
    (...)
/* rank of processes proc_top and proc_bottom */
MPI_Cart_shift(grid_comm, Y_DIR, 1, &proc_top, &proc_bottom);
/* rank of processes proc_left and proc_right */
MPI_Cart_shift(grid_comm, X_DIR, 1, &proc_left, &proc_right);
```

There a couple new function calls on this code whose arguments I will explain next. As explained in the exercise description MPI uses the MPI_Cart_* function calls to arrange tasks in a virtual process grid.

To create one the API calls needs the previous communicator, in our case we were using MPI_COMM_WORLD. ndims and dims define the number of dimensions of the grid and the number of processors in each, respectively. For our example these are 2 and P_grid. Thereafter the periods specifies if the grid is periodic or not per dimension, and finally if the ranking is reordered or not. These are replaced by the self-explanatory variables wrap_around and reorder. The new communicator is stored in the address of comm_cart. From now on all pointers to communicators refer to this one.

```
int MPI_Cart_create(MPI_Comm comm_old, int ndims, int *dims, int *perion int MPI_Cart_coords(MPI_Comm comm, int rank, int maxdims, int *coords) int MPI_Cart_shift(MPI_Comm comm, int direction, int displ, int *source
```

To define the coordinates of the process in the new communicator we use MPI_Cart_coords. Here the rank is necessarily the processor rank (proc_rank), the maximum dimensions is 2. The coordinates of each specified process are stored in the proc_coord array.

Finally the shift operation returns the shifted source and destination ranks. The direction and displacement quantity arguments are self-evident and are replaced in the program by X_DIR or Y_DIR and 1 respectively for horizontal and vertical displacements. In accordance to the outputs source and destination are stored in proc_top and proc_bottom or proc_left and proc_right depending on the direction.

The text written to the standard output now features the coordinate of each processor.

```
(0) Number of iterations: 2355
```

$$(0) (x,y) = (0,0)$$

```
(0) Elapsed Wtime: 1.414062 s (96.9% CPU)
```

- (1) Number of iterations: 2355
- (1) (x,y)=(1,0)
- (1) Elapsed Wtime: 1.464844 s (95.6% CPU)

As one can observe since there are two processor, number 0 is allocated the left half of the grid, and processor with rank 1 deals with the right half.

2.1.7. Step 7

When there are three processors the work can not be evenly split between them. This can be confirmed by inspecting the x and y variables in the Setup_grid function.

For example for processor 2 (in a 3 processor configuration) it is visible that x is always negative.

$$(2) x = -30, \dim[X_DIR] = 36$$

$$(2) y = 71, \dim[Y_DIR] = 102$$

$$(2) x = -3, \dim[X DIR] = 36$$

$$(2) y = 76, \dim[Y DIR] = 102$$

$$(2) x = -28, \dim[X DIR] = 36$$

$$(2) y = 26, \dim[Y DIR] = 102$$

2.1.8. Step 8

2.1.9. Step 9

After implementing the collective reduction operation the total number of iterations is indeed the same, as confirmed by the program's output.

```
(0) Number of iterations: 2355
```

(1) Number of iterations: 2355

(2) Number of iterations: 2355

(2) Elapsed Wtime: 1.507812 s (96.8% CPU)

2.1.10. Step 10

2.2. Part 2

$2.2.1.\ 2.1$

2.2.2. 2.2

After changing the code to accommodate for the algorithmic improvement, 5 tests were performed to compare different values for the relaxation parameter ω . The results are summarized in table 2.1. From this data we concluded that 1.93 is the optimal value for the relaxation parameter, accomplishing almost 18 times less iterations than the original with ω equal to one.

| ω | $Maximum \ \mathtt{Wtime}^1 \ (s)$ | Iterations | Reduction |
|------|------------------------------------|------------|-----------|
| 1.00 | 1.500000 | 2355 | 1.00 |
| 1.90 | 0.222656 | 220 | 10.7 |
| 1.92 | 0.199219 | 165 | 14.3 |
| 1.93 | 0.175781 | 131 | 18.0 |
| 1.94 | 0.289062 | 142 | 16.6 |
| 1.98 | 0.351562 | 419 | 5.62 |

Table 2.1: Time, number of iterations obtained and respective iteration reduction for different ω values. The topology used was pt: 4–4 1 with a grid size of g: 100 x 100.

2.2.3. 2.3

The goal of this exercise is to investigate the scaling behavior of the code with a fixed relaxation parameter. To accomplish that analysis several runs were measured with various grid sizes. In addition different *slices* were also tested.

| Grid Size | Wtime for 4 4 1 (s) | Wtime for 4 2 2 (s) | Δ (%) |
|--------------------|---------------------|---------------------|--------------|
| $200 \ge 200$ | 0.464844 | 0.699219 | 50.42 |
| 400 ± 400 | 1.054688 | 1.054688 | 0.00 |
| 800×800 | 3.742188 | 3.726562 | -0.42 |
| 2000×2000 | 21.351562 | 21.328125 | -0.11 |

Table 2.2: The maximum time for different grid sizes and different slicing arrangements. The Δ column compares the relative difference in performance between the two previous columns. The number of iterations for each run was fixed at 300.

 $^{^1{\}rm The}$ maximum time was computed over the four individual processor times for each ω value.

The results of the aforementioned experience are shown in table 2.2. As one can see as the grid size increase the difference in the different topologies become negligible. Nonetheless for small sizes the 4 4 1 topology is significantly faster.

To to determine the average time per iteration the time will be parametrized as follows

$$t(n) = \alpha + \beta \cdot n$$

3. Lab 3: Finite

4. Lab 4: Nbody

5. Lab 5: Matmul