Overview Load balancing Work pools Summary and next lecture

XJCO3221 Parallel Computation

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Lecture 13: Load balancing

Previous lectures

Several times in this module we have mentioned the concept of **load balancing**:

- Poor load balancing results in processing units spending time idle.
- Usually realised when **synchronising** threads or processes.
- First encountered for the Mandelbrot set generator [Lecture 3].
- Important for parallel performance for all architectures shared and distributed memory CPU, and GPU.

This lecture

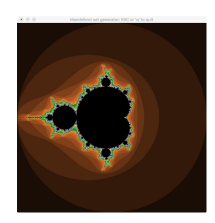
Here we will look at load balancing more closely, and how to reduce its performance penalty:

- Return to the example of the Mandelbrot set generator, this time in MPI.
- Understand how heterogeneity in the problem results in poor load balancing.
- See how a task scheduler can improve load balancing at runtime.
- Go through a concrete example of a work pool.

The Mandelbrot set (c.f. Lecture 3)

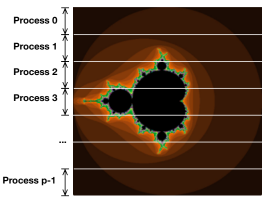
Code on Minerva: Mandelbrot_MPI.c plus makefile

- Domain is $-2 \le x \le 2$ and $-2 \le y \le 2$.
- Calculation performed iteratively for each (x, y).
- Pixel coloured according to the number of iterations.
- Here, the black region corresponds to a high number of iterations.
- No upper bound some points will iterate indefinitely if allowed.



Strip partitioning

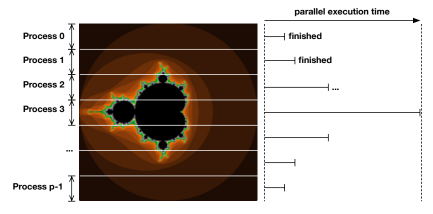
Partition the domain [cf. last lecture] into horizontal strips¹:



¹Equivalent results for partitioning into vertical strips, or blocks.

Load imbalance

Because some pixels take longer to calculate the colour than others, the load is **unevenly distributed** across the processes:



Load balancing

- Parallel execution time determined by the last processing unit to finish.
- Poor load balancing results in significant idle time for at least one process/thread.
- Inefficient use of available resources.

Definition

The goal of **load balancing** is for each **processing unit** (thread or process) to perform a **similar volume of computations**, and therefore finish at roughly the same time.

Up until now most problems we have encountered have been **naturally load balanced**.

For example, for vector addition between two *n*-vectors, assigning each processing unit to equal numbers of vector elements results in good load balancing.

• Each unit performs n/p additions.

Note that the Mandelbrot set is a map, *i.e.* an **embarrassingly parallel problem** (since there are no data dependencies).

• Still a challenge to attain good performance.

Static load balancing

Definition

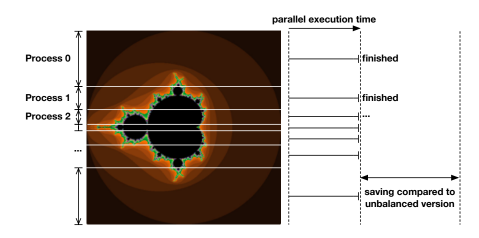
Sometimes it is possible to determine (approximately) equal loads at compile time. This is known as static load balancing.

For the Mandelbrot set example, we could assign **larger** domains to regions where the calculations should be **fast**.

Should improve load balancing.

However, an **exact** expression is not available. Therefore any such **heuristic** can only achieve **approximate** load balancing.

Static load balancing (ideal case)



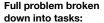
Dynamic load balancing

Definition

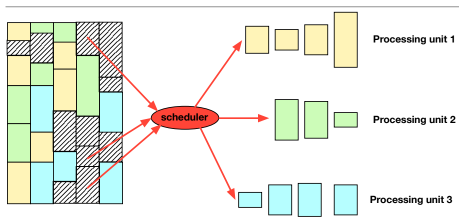
Dynamic load balancing is performed **at runtime**. No *a priori* knowledge of computation times is required.

Basic idea:

- Break the problem down into small independent tasks.
- 2 Each processing unit performs one task at a time.
- When it is complete, it starts/is assigned another task.
- Repeat 3 until all tasks are complete.



Tasks performed sequentially on different threads/processes:



Functional or task parallelism

Up to now we have largely considered parallelising the same operation to a (large) data set.

- Known as data parallelism.
- When performed in a loop, can also be termed loop parallelism.

Now we are parallelising a number of tasks.

- Called task parallelism or functional parallelism.
- Be warned that these terms are sometimes used to refer to slightly different concepts.
- More on task parallelism in Lecture 19.

Work pools

The **scheduler** assigns tasks to processing units **at runtime**.

- Often part of the parallel/concurrency runtime system.
- Introduces a (small) overhead.
- Various algorithms exist for efficient task scheduling.

To understand the role of a scheduler, we will look at a simple scheduler implemented in MPI - a **centralised work pool**.

- One process (usually rank 0) performs the scheduling this is the **main** process¹.
- Remaining processes action the tasks the workers¹.

¹You may see 'master' (for main) and 'slaves' (for workers) in the literature.

Worker pseudocode

Function workerProcess() in Mandelbrot_MPI.c

```
initialise(); // Including MPI_Init().
3 while( true )
4 {
    // Wait for message from the main (rank 0).
5
    MPI_Recv( message, ... );
6
7
    // Is this a termination request?
    if( message == TERMINATE ) break;
9
10
    // Else perform calculation and send back to rank 0.
    result = actionTask( message );
12
    MPI_Send( result, ... );
13
14 }
15
16 finalise(); // Including MPI_Finalize().
```

Main process pseudocode (1)

Function mainProcess() in Mandelbrot_MPI.c

```
initialiseAndOpenWindow();

// Initialise variable that tracks progress.
int numActive=0;

// Send initial request to each worker.
for( p=1; p<numProcs; p++ )
{
    MPI_Send(task,...,p,...);
    numActive++;
}</pre>
```

For this Mandelbrot example, each task is a **row of pixel colours** to be calculated.

Keep track with an incrementing variable row.

Main process pseudocode (2)

Function idle() in Mandelbrot_MPI.c

```
while( numActive>0 )
2 {
    // Get result from ANY worker process.
3
    MPI_Recv(result,...,MPI_ANY_SOURCE,...,&status);
4
5
    numActive --:
6
7
    // Send request IMMEDIATELY to the SAME worker.
    if (!finished)
8
    {
9
      MPI_Send(task,...,status.MPI_SOURCE,...);
      numActive++;
    }
12
13
    // Action the message.
14
    actionResult( result );
15
16 }
```

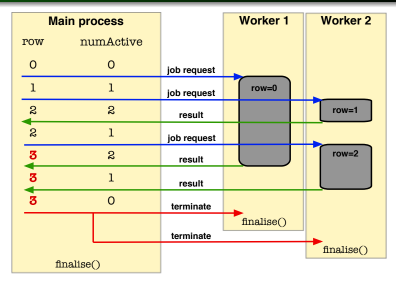
Main process pseudocode (3)

Function idle() in Mandelbrot_MPI.c

```
1 // Tell all workers to terminate.
2 for( p=1; p<numProcs; p++ )
3 MPI_Send(TERMINATE,...);
4
5 finaliseAndCloseWindow();</pre>
```

- MPI_ANY_SOURCE in place of source in MPI_Recv() receives a message from any process.
- Used status.MPI_SOURCE to recover the rank of the sending process.
- Send next request before the (potentially slow) call to actionResult().

Example: 3 rows and 2 workers



Modern schedulers

There are many more types of **work pool**, such as those with no 'main' process — a **decentralised** work pool¹.

A common approach is to use **deques** or <u>double ended queues</u>:

- Each processing unit maintains its own deque of tasks.
- Performs tasks sequentially, starting from the front.
- Once the deque is empty, 'steals' a task from the back of a randomly selected 'victim' (work stealing).

¹Wilkinson and Allen, Parallel Programming (Pearson, 2005).

OpenMP scheduler

OpenMP can also schedule loops using its schedule clause:

```
#pragma omp parallel for schedule(dynamic,chunk)
for( i=0; i<n; i++ ) { ... }</pre>
```

This breaks down the loop into 'chunks' of size chunk at runtime.

Can also be used for **static** scheduling:

```
#pragma omp parallel for schedule(static,chunk)
```

There is also a **guided** option that decreases the chunk size exponentially **at runtime** to the final value chunk:

```
#pragma omp parallel for schedule(guided,chunk)
```

In all cases, chunk is optional and defaults to 1.

MIMD at last!

Up until today we have mostly performed the **same** calculations on each processing unit.

- SIMD (Single Instruction Multiple Data) software . . .
- ...on MIMD (<u>M</u>ultiple <u>Instruction M</u>ultiple <u>D</u>ata) **hardware**.

Today is the first clear¹ example where we have implemented the MIMD pattern **in software**.

 The main process perform entirely different calculations to workers — there is a division of labour.

¹Ignoring trivial cases like e.g. rank 0 distributing global arrays.

Summary of distributed memory systems

Lec.	Content	Key points
8	Architectures	Clusters and supercomputers; intercon-
	and MPI	nect network; starting with MPI.
9	Point-to-point	Blocking send and receives; buffering;
	communication	deadlock for cyclic communication.
10	Data reorgani-	Scatter and gather; $t_{ m comm}$; collective
	sation	communication in MPI.
11	Reduction	Binary trees; OpenMP and MPI.
12	Asynchronous	Non-blocking send and receives; domain
	communication	partitioning and ghost cells.
13	Load balancing	Task parallelism; schedulers; work pools.

Next lecture we start looking at programming **general purpose graphics processing units** or GPGPUs.