

## Università di Pisa

## Parallel and distributed systems: Paradigms and Models

# A parallelized version of the Jacobi Method

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#### Abstract

The Jacobi Method is an algorithm for finding the solution of a system of linear equations Ax = b. This C++ project aims to implement a sequential version of this algorithm, a parallelized version using native threads, and finally a parallel version with Fast Flow.

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#### 1 Introduction

The **Jacobi Method** is an algorithm for finding the solution of a system of linear equations Ax = b, where  $A \in \mathbb{R}^n \times \mathbb{R}^n$  is the matrix of coefficients of the system of linear equations,  $b \in \mathbb{R}^n$  is a vector of known values and  $x \in \mathbb{R}^n$  is the vector we want to approximate. At each iteration k, we calculate the new values contained in the vector x using the following formula:

$$x_i^{k+1} = \frac{1}{a_{(i,i)}} b_i - \sum_{i \neq j} a_{(i,j)} x_j^k \qquad \forall i \in 1 \dots n$$
 (1)

The pseudo-code of the algorithm, which is based on this equation, is defined in **Alg.1**.

It has been demonstrated that if the matrix A is strictly diagonally dominant, Jacobi method is guaranteed to converge. For this reason, I assumed that all the matrices examined respect that property.

The purpose of this project is the C++ implementation of this algorithm in parallel version, in order to achieve the same results with less time.

This report is organized as follows: In section 2 I described the theoretical analysis of the parallelization of the Jacobi method. In section 3 I described the methodologies used to solve the task. In section 4 I showed the results. In section 5 I concluded the work.

#### 2 Analysis

The algorithm uses two vectors x and  $tmp_{-}x$ , referring to **Eq.1**, the first vector represents  $x_i^k$  while the second vector represents  $x_i^{k+1}$ .

The values of the vector  $tmp_x$  are computed at each iteration and only at the end of an iteration do we perform the swap to x. Thus, in order to compute each value of  $tmp_x[i]$ , we need to know the corresponding previous value x[i]. This means that it is not possible to parallelize the cycle while (referring to Alg.1). However, the remaining for loops can be parallelized.

This algorithm is a perfect instance of **data parallel pattern** since for each iteration of the first for loop (lines 10-18) we have already everything

we need to compute the cycle (that is, the data contained in the vector  $\mathbf{x}$ ). In particular we can recognize map pattern, since we can apply the Jacobi method to each item of the vector independently.

Moreover, the second for loop (lines 12 - 16) can be seen as a *reduce* parallel pattern, since we have to perform an operation, the sum, that is commutative and associative. We could then perform partial sums in parallel and accumulate them with each other, as reduce pattern dictates. In practice, for this kind of problem, it is not very useful because adding this parallelization also means adding a lot of overhead.

For the same reason, I also did not parallelize the norm calculation, which is necessary for early stopping, which in theory, it is possible to parallelize. In parallel implementations, in light of the fact that each worker works (more or less) on a portion of similar size compared to all other workers, there was no need to implement load balancing techniques.

As described in more detail in the 3.2 section, the experiments on the different implementations were performed using different metrics. All of these metrics, however, are based on two values:  $T_{seq}(n)$ ,  $T_{par}(n, nw)$  which are the times required to perform the Jacobi method sequentially and in a parallel way, respectively, on a matrix of dimensions n (and nw refers to the number of workers). We now examine these two values in more detail.

Let M be the number of iterations.

Due to the way the algorithm was designed, the sequential time  $T_{seq}(n)$  can be seen in this way:

$$T_{seq}(n) = T_{init} + M \cdot (n \cdot T_{single\_iter} \cdot T_{ES})$$
 (2)

where  $T_{init}$  is the time for initializing the variables,  $T_{single\_iter}$  is the time for a single iteration and  $T_{ES}$  is the time to compute the norm for early stopping.

Instead, the time to run the algorithm in parallel,  $T_{par}(n, nw)$ , can be seen as:

$$T_{par}(n, nw) = T_{init} + T_{split} + T_{merge} + M \cdot \left(\frac{n}{nw} \cdot T_{single\_iter} + T_{ES}\right)$$
(3)

where  $T_{split}$  and  $T_{merge}$  are the overheads for splitting and merging the whole iteration among the workers.

#### 2.1 Speedup predictions

The ideal speedup is a value that grows linearly with the number of workers used. In practice, as we will see in the section 4 we will not achieve this because there are overheads and serial fractions that prevent us from achieving our ideal result.

Knowing the parallel version of the algorithm, we can try to estimate the speedup we can achieve in this way:

Given the sequential time  $T_{Seq}$  we know that the ideal time we would like to achieve using nw workers is:

$$T_{ideal} = T_{Seq}/nw$$

Since we know nw, and we can calculate  $T_{Seq}$ , so we can get the value of  $T_{ideal}$ . In addition, we can calculate the value of overheads due to the creation and join of threads and the average waiting time of a thread due to the barrier. Hence, we can then predict our speedup by imagining the parallel time  $T_{Par}$  as the sum of the ideal time and the overheads just described.

$$predicted\ speedup = \frac{T_{Seq}}{(T_{ideal} + barrier\ overhead + thread\ creation\ overhead)}$$

Assuming then a number of workers equal to 18, and n = 4096, I measured the following values (all values are measured in  $\mu$ sec):

$$T_{Seq} = 113739$$

$$T_{ideal} = 113739/18 = 6318.833$$

$$thread\ creation\ overhead = 664$$

$$barrier\ overhead = 1228.06$$

$$predicted\ speedup = \frac{113739}{6318.833 + 1228.06 + 664} \approx 13$$

Clearly, this calculation is an approximation and is only an upper bound on the actual speedup achieved, as it does not take into account other factors such as any overhead due to caching or the behavior of the scheduler.

#### 3 Methods

#### 3.1 Setup

The software starts by parsing the arguments [utils/argument\_parser.cpp], if they are specified. If the program is compiled without making the arguments explicit, the default values for each argument are obtained using the constants defined in setup/constants.h. For more information on compilation options, see appendix B.

At this point, the software runs unit tests; these were very useful for me to make sure, while writing the project, that the functions tested were correct in their required behavior. Unit tests are managed by the UnitTestManager class.

Once the unit tests have been run, the software generates a strictly diagonally dominant matrix A and a vector b, both with random values distributed in a range [RAND\_LOW\_VALUE, RAND\_HIGH\_VALUE], where these values are constants in setup/constants.h.

Once the matrix A and vector b have been generated, the software executes the Jacobi method on these values, in particular, it first executes the sequential version solutions/sequential.cpp, then the parallel version with native threads solutions/parallel\_threads.cpp, and finally the parallel version with Fast Flow solutions/fast\_flow.cpp

In the end, if required from the user, the software performs the experiments that are described in section 4.

#### 3.2 Metrics

In order to evaluate the quality of the different parallel versions of the sequential algorithm of Jacobi Method, I selected three performance indicators: **Speedup**, **Scalability** and **Efficiency**.

Each of these metrics was evaluated with respect to completion time; actually, the completion time is defined as the time required by an algorithm from when it starts until it finishes. In the case of this project, however, I thought it appropriate to exclude from this time all the calculations to perform the unit tests, the initialization of the A matrix and the b vector, and in general, anything that was not strictly related to the resolution of the algorithm itself.

#### 3.3 Sequential

The first version of Jacobi Method that I implemented was the sequential version. It is, in essence, an exact translation of the pseudo code found in appendix A.

The implementation of this version can be found in solutions/sequential.cpp.

#### 3.4 Native C++ Threads

The first way I thought of in order to parallelize the algorithm was to use a barrier in order to synchronize several threads, which work simultaneously to compute the values of  $tmp_{-}x$  (each thread works on different portions of the vector, independently). When all the threads finish, using the barrier, I updated the vector x. The barrier was implemented using the class std:barrier. The use of barriers introduced an overhead that was described in 4. The implementation of this version can be found in  $solutions/parallel\_threads.cpp$ .

#### 3.5 Fast-Flow

After developing the parallel version of the algorithm with threads, I implemented a parallel version of Jacobi's method using the library Fast-Flow[1]. In particular, I used the method ff::ParallelFor, designed to parallelize a map pattern, specifically it is able to parallelize a lambda function passed as input on a worker number, also specified as input. In our case, the function passed to the method is responsible for computing the individual values of  $tmp_{-}x$ . The implementation of this version can be found in solutions/fast\_flow.cpp.

#### 4 Experiments and Results

**Experiments** The experiments I performed were comparisons of the different implementations against the three selected metrics (Speedup, Scalability, Efficiency), and they were performed on servers provided by the University of Pisa that use an Intel Xeon Gold 5120 CPU @  $2.20 \mathrm{GHz}$  32 cores CPU. Tests were performed using matrices of size:  $1024 \times 1024$ ,  $4096 \times 4096$ ,  $16384 \times 16384$ .

Each experiment was performed 10 times and then the results were averaged: this was done to obtain more reliable results. Moreover, for a proper comparison, the seed remained constant while running the tests: thus the three matrices (of the three different dimensions) are exactly equal to each other.

To conclude, all experiments were performed with the early stopping flag active. Experiments run in batch mode at the end of main if the argument results=1 is set when the program is executed.

**Results** Figures 1,2,3 show us, respectively, the values of Speedup, Scalability and Efficiency, obtained on the three dimensions of the matrices.

We can immediately see that the size of the matrices plays a key role in choosing the best implementation, indeed in Fig. 1(a) we can see that the speedup calculated on the matrix  $1024 \times 1024$  is very far from an ideal speedup (which is linear with respect to the number of workers). Instead, the matrix  $16384 \times 16384$  (Fig. 1(b)) is closer to an ideal speedup, especially for the first few iterations.

Table 1 shows a comparison of times between native threads and Fast Flow, while Table 2 shows a comparison of metrics between native threads and Fast Flow.

The introduction of the barrier introduces a clear overhead within the implementation with native C++ threads. In particular, we can see in figure 4 that the more we increase the number of workers, the more the overhead due to the barrier increases.

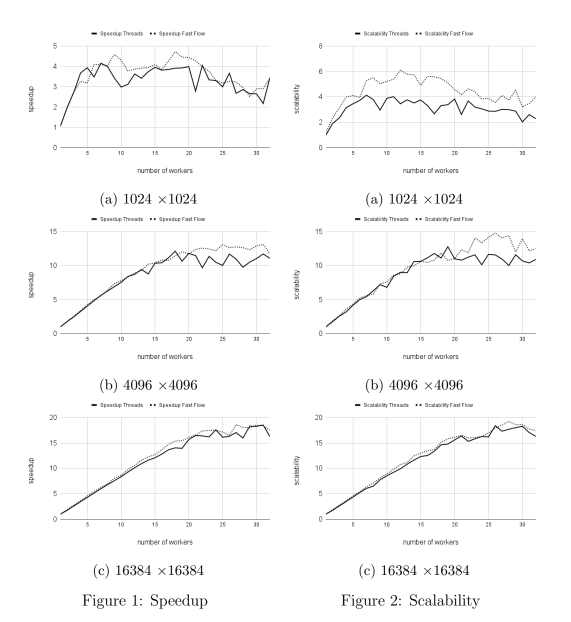
In addition, I also calculated the overhead required to perform thread creation and join, in the case of the implementation with native C++ threads. Figure 5 shows that the overhead is linear to the number of workers used, regardless of the size of the A matrix.

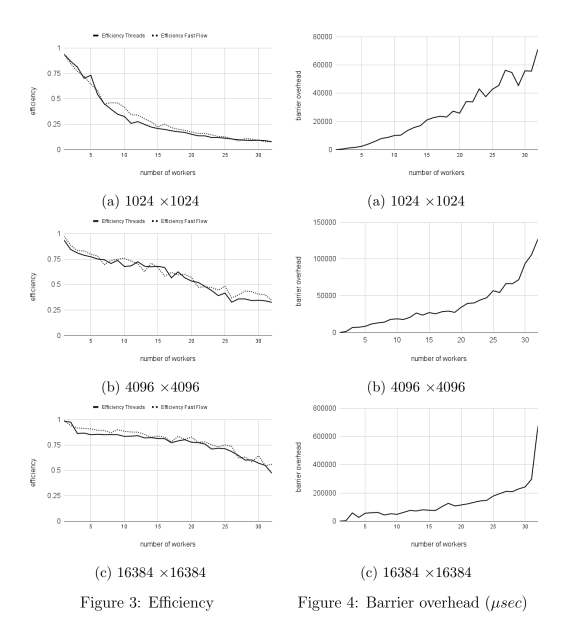
Matrix size	Implementation	Time $(\mu sec)$	nw
1024	Native Thread	2306.2	22
	Fast Flow	1977.6	18
	Sequential	9321	
4096	Native Thread	9188.8	18
	Fast Flow	$\boldsymbol{8481.2}$	31
	Sequential	111209	
16384	Native Thread	69808	31
	Fast Flow	69671.6	27
	Sequential	1291918	

Table 1: Best time

Matrix size	Implementation	Speedup	Scalability	Efficiency
1024	Native Thread	4.1574	4.13315	0.939837
	Fast Flow	4.72489	6.0993	0.939749
4096	Native Thread	12.1199	12.7777	0.937034
	Fast Flow	13.1255	14.8002	0.975051
16384	Native Thread	18.5283	18.3889	0.98332
	Fast Flow	18.6151	19.2443	0.990929

Table 2: Best results





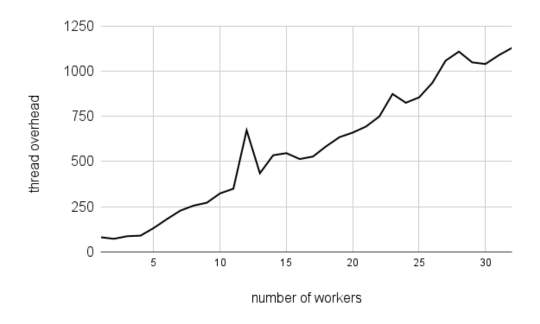


Figure 5: Thread overhead  $(\mu sec)$ 

#### 5 Conclusions

In this project, an analysis was presented for parallelization of Jacobi's method algorithm on strictly diagonally dominant matrices, through a comparison of two different types of parallel implementations (Fast Flow and native threads). From this comparison we can see that Fast Flow behaves similarly to the implementation with native threads.

If the matrix is very small, it is not worth the effort to parallelize, since the overheads involved in setting up parallelization can make parallel execution slower than sequential execution.

Finally, personally, working on this project made me realize how overheads cannot be taken for granted if we really want to use all available computing power to solve our tasks.

#### References

[1] Marco Aldinucci et al. "Fastflow: High-Level and Efficient Streaming on Multicore". In: Programming multi-core and many-core computing systems. John Wiley Sons, Ltd, 2017. Chap. 13, pp. 261–280. ISBN: 9781119332015. DOI: https://doi.org/10.1002/9781119332015. ch13. eprint: https://onlinelibrary.wiley.com/doi/pdf/10.1002/9781119332015.ch13. URL: https://onlinelibrary.wiley.com/doi/abs/10.1002/9781119332015.ch13.

## A Jacobi pseudocode

#### Algorithm 1 Jacobi Method

```
Require: \epsilon
                                          ▷ Convergence threshold (default 0.0001)
 1: function JACOBI (max_iterations, A \in \mathbb{R}^n \times \mathbb{R}^n, b \in \mathbb{R}^n)
        n = A.size()
        Initialize x as a vector of n zeros
 3:
         while true do
 4:
             if iteration = 0 then
 5:
                 break
 6:
             end if
 7:
 8:
 9:
             iteration \leftarrow iteration - 1
             for i \leftarrow 1 to n do
10:
                 sum = 0
11:
                 for j \leftarrow 1 to n do
12:
                     if i \neq j then
13:
                         sum \leftarrow sum + A_{(i,j)} * x_j
14:
                     end if
15:
                 end for
16:
                 tmp_{-}x_{i} = \frac{1}{A_{(i,i)}} \cdot (b_{i} - sum)
17:
             end for
18:
19:
             if distance(x, tmp_x) < \epsilon then
20:
                                                                         ▶ Eucl. distance
                 break
21:
             end if
22:
23:
             x \leftarrow tmp_-x
        end while
24:
25:
        return a
26: end function
```

### B How to compile and execute

To make compilation and execution easier and faster, you can run the bash script:

./run.sh

If necessary, you can still change any parameter by first compiling the program by running the command:

make

and then:

./main.out <params>

where parameters are the following:

parameter	domain	default value
n	integer $\geq 0$	5000
num_of_worker	integer $\geq 0$	100
$num\_of\_iterations$	integer $\geq 0$	10
calculate_results	boolean $(\in \{0,1\})$	0
$early\_stopping$	boolean $(\in \{0,1\})$	1
execute_unit_test	boolean $(\in \{0,1\})$	1

Table 3: Parameters