MASA: a Multi-Platform Architecture for Sequence Aligners with Block Pruning

Edans F. de O. Sandes, Department of Computer Science, University of Brasilia, Brazil
Guillermo Miranda, Barcelona Supercomputing Center, Spain
Xavier Martorell, Barcelona Supercomputing Center and Universitat Politecnica de Catalunya, Spain
Eduard Ayguade, Barcelona Supercomputing Center and Universitat Politecnica de Catalunya, Spain
George Teodoro, Department of Computer Science, University of Brasilia, Brazil
Alba C. M. A. de Melo, Department of Computer Science, University of Brasilia, Brazil

Biological sequence alignment is a very popular application in Bioinformatics used routinely worldwide. Many implementations of biological sequence alignment algorithms have been proposed for multicores, GPUs, FPGAs and CellBEs. These implementations are platform-specific and porting them to other systems requires considerable programming effort. This paper proposes and evaluates MASA, a flexible and customizable software architecture that enables the execution of biological sequence alignment applications with three variants (local, global and semi-global) in multiple hardware/software platforms with block pruning, which is able to reduce significantly the amount of data processed. To attain our flexibility goals, we also propose a generic version of block pruning and developed multiple parallelization strategies as building blocks, including a new asynchronous dataflow based parallelization, which may be combined to implement efficient aligners in different platforms. We provide four MASA aligner implementations for multicores (OmpSs and OpenMP), GPU (CUDA) and Intel Phi (OpenMP), showing that MASA is very flexible. The evaluation of our generic block pruning strategy shows that it significantly outperforms the previously proposed block pruning, being able to prune up to 66.5% of the cells when using the new dataflow based parallelization strategy.

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Additional Key Words and Phrases: Biological Sequence Alignment, Parallel Algorithms, GPU, multicores, Intel Phi

1. INTRODUCTION

The astonishing evolution of DNA sequencing techniques is producing an overwhelming number of new biological sequence data to be analyzed. Also, the number of laboratories that are analyzing the sequences is quickly increasing. To cope with this, Life Sciences laboratories are facing the challenge of producing accurate results in a very short time.

Once a new biological sequence is produced, its functional/structural characteristics must be established. In order to do that, the newly discovered sequences are compared against other sequences, looking for similarities. This is done in a daily basis, all over the world.

Biological sequence comparison is, therefore, a very important operation in Bioinformatics [Mount 2004]. It produces (a) a score, indicating the similarity between the sequences and, optionally, (b) an alignment, which highlights the regions of similarities/differences between the sequences.

Biological sequences can be DNA, RNA or protein sequences. Protein and RNA sequences are rather small and their sizes range from hundreds to tens of thousands of residues (amino acids and nucleotide bases, respectively). On the other hand, DNA sequences can be very long, often composed of Millions of base pairs (Mbp).

There are three types of comparisons: (a) global, where all the characters of the sequences belong to the alignment; (b) local, where a subset of the characters belongs to the alignment and (c) semi-global, where the head/tail of the sequences is discarded. Depending on the analysis, the biologists may choose among the types of the sequences
(DNA, RNA or protein), the comparison type (local, global, semi-global) and the output produced (score, score and alignment).

The Needleman-Wunsch (NW) algorithm [Needleman and Wunsch 1970] is an exact algorithm that retrieves the optimal score/alignment of a global pairwise sequence comparison. It is based on Dynamic Programming (DP) and calculates a DP matrix of size $n \times m$, where $n$ and $m$ are the sizes of the sequences. NW has quadratic time and space complexity $O(nm)$.

Later, in 1981, the Smith-Waterman (SW) algorithm [Smith and Waterman 1981] was proposed. It calculates the optimal score/alignment of a local sequence comparison by slightly modifying the recurrence relation used in the computation of the NW DP matrix. As in NW, SW has time and space complexity $O(nm)$.

In order to reduce the execution time of the NW and SW exact methods, heuristic methods such as BLAST [Altschul et al. 1990] were proposed. These methods combine exact pattern matching with dynamic programming in order to produce good solutions faster. BLAST can align sequences in a very short time, still producing good results. Nevertheless, its accuracy is expected to be worse than the accuracy of the exact methods. In this paper, we focus on the exact algorithms.

Genome-wide and chromosome-scale alignments are typically carried out during assembly projects. The second-generation sequence technologies increased the sequencing throughput but reduced the read lengths to hundreds of base pairs per read, leading to assembly issues related with repetitive and redundant regions of a genome [Wang 2013]. In the emerging third-generation sequence technologies, as in PacBio single-molecule real-time (SMRT) sequencing, the read lengths are increasing to thousands of base pairs per read [Ainala et al. 2015][Zhang et al. 2014] and long sequence alignments are likely to gain much more importance in the near future.

The alignment of biological sequences longer than 10 Mbp with exact DP algorithms such as SW and NW is still considered unfeasible by most of the researchers and the use of these algorithms in chromosome-wide scale has not been fully explored yet. In order to allow a fair comparison among heuristic and exact methods for chromosome-wide alignments, optimized tools able to produce exact long DNA alignments are necessary. We claim that local/global/semi-global optimal chromosome-wide alignments can be very useful since they can reveal complementary information that is not obtained with other tools/techniques.

In the last years, many parallel variants of SW have been proposed for CPUs with Single Instruction Multiple Data (SIMD) vector instructions [Farrar 2007], clusters [Rajko and Aluru 2004] [Chen and Schmidt 2003], Field Programmable Gate Arrays (FPGAs) [Yamaguchi et al. 2011] [T. F. Oliver 2005], Cell Broadband Engines (CellBEs) [Sanchez et al. 2010] [Sarje and Aluru 2009] [Wirawan et al. 2009], Graphics Processing Units (GPUs) [Sandes and Melo 2013a] [Liu et al. 2013] [Manavski and Valle 2008] [Xiao et al. 2009] [Korpar and Sikic 2013] and, more recently, for the Intel Phi [Liu and Schmidt 2014b]. Even though these SW proposals commonly share some similar features or code blocks that could be reused, they usually have a design which is highly platform-dependent. Therefore, porting them to other High Performance Computing platforms tends to involve a significant programming effort.

Still, in the last years, there have been proposals to use these exact methods in Megabase sequences. Besides CUDAAlign [Sandes and Melo 2013a], in the recent literature we can find (a) SW# [Korpar and Sikic 2013], which executes SW comparisons of $33 \text{ Mbp} \times 46 \text{ Mbp}$ on GPUs and (b) SWAPIH-LS [Liu et al. 2014], which presents comparisons of $42 \text{ Mbp} \times 50 \text{ Mbp}$ on the Intel Phi. As the need for long sequence comparison is supposed to receive more attention in the next years, it is expected that these tools will gain a lot of visibility and many more optimizations must be proposed.
in order to benefit from the new computing platforms, aiming to exploit their potential performance.

CUDAlign [Sandes and Melo 2013a] is an optimized parallel variant of SW that is able to compare Megabase DNA sequences (>1 Mbp) using GPU. Differently from other variants of SW, which calculate the whole DP matrix, CUDAlign proposed the Block Pruning (BP) optimization, which is able to reduce the number of cells calculated while still guaranteeing that the optimal local alignment will be produced. BP is suitable for regular processing patterns (e.g. diagonal by diagonal) and it was able to reduce the computation in more than 50%, when the sequences have high similarity. Since the code of CUDAlign is highly dependent on CUDA, it only runs on NVidia GPUs. Nevertheless, we identified that more than 90% of the CUDAlign code was platform-independent, thus the full reimplementation to a new platform would require only a small amount of coding effort in order to rewrite the 10% of the code related to the platform-specific section.

This paper proposes and evaluates MASA (Multi-Platform Architecture for Sequence Aligners), a software architecture for sequence aligners, based on CUDAlign. The main contributions of MASA are: (a) a flexible and customizable architecture for deployment of pairwise biological sequence aligners in different hardware/software platforms; (b) a generic version of the Block Pruning (BP) optimization; (c) a novel asynchronous parallelization strategy based on the dataflow model and (d) four parallel MASA Aligner implementations for multicores and accelerators.

In order to design MASA, we faced the challenge of modifying CUDAlign in order to decouple the small platform-specific code from the platform-independent one, creating a simple integration API without loosing performance. For the sake of this, a generic version of BP needed to be proposed to support the processing of the DP matrix in an asynchronous way, where the regular diagonal processing pattern is not guaranteed.

Besides proposing the MASA architecture, in this paper we design and implement four MASA aligners, which consists of rewriting the platform-specific code to four distinct hardware/software platforms. MASA-CUDAlign runs on NVidia GPUs. ASA-OpenMP/Phi is a sequence aligner that runs on the Intel Phi accelerator, programmed with OpenMP. MASA-OpenMP/CPU is the OpenMP version that runs on multicores. Finally, MASA-OmpSs/CPU is an OmpSs [Duran et al. 2011] version that also runs on multicores. With these 4 implementations, we show that (a) multiple hardware platforms can be integrated to MASA and that (b) high-level parallel programming environments such as OmpSs and OpenMP can also be integrated.

In addition, we included in MASA not only the local sequence alignment exact algorithm (SW), but also the global (NW) and semi-global variants. Finally, a new block pruning strategy was created in the MASA portable code in such way to allow the computation of the DP matrix in a generic execution order, not restricted to antidiagonals, as it is implemented in CUDAlign. The source code of MASA is freely available and can currently be found in https://github.com/edanssandes/MA SA-Core.

Experimental results obtained with real DNA sequences show that our multicore implementations (MASA-OmpSs/CPU and MASA-OpenMP/CPU) outperform MASA-CUDAlign and MASA-OpenMP/Phi for small sequences. For longer sequences, MASA-CUDAlign presents the best execution times. We also show that generic BP combined with the OmpSs dataflow model is able to prune more cells than the original BP and, as a consequence, MASA-OmpSs/CPU outperforms MASA-OpenMP/CPU in most cases. Finally, we present the alignment results for (a) two strains of the Amycolaptosis mediterranei bacteria (10 Mbp x 10 Mbp) and (b) the human x chimpanzee homologous chromosomes 21 (47 Mbp x 32 Mbp).

The rest of this paper is organized as follows. Section 2 presents the biological sequence alignment problem. Section 3 presents the related work in the area of biologi-
2. BIOLOGICAL SEQUENCE ALIGNMENT

DNA biological sequences are treated as strings composed by elements of the alphabet \( \Sigma = \{A, T, G, C\} \). To compare two sequences, it is necessary to find an alignment between them, placing one sequence above the other and making clear the correspondence between the characters [Mount 2004]. In the alignment, spaces (gaps) can be introduced in one of the sequences, in order to improve the alignment quality. Each alignment has a score, which measures the similarity between the sequences. Our goal is to obtain the alignment with the highest score.

2.1. NW and SW Algorithms

The NW algorithm [Needleman and Wunsch 1970] is based on Dynamic Programming (DP) and obtains the optimal global alignment in quadratic time and space. It takes as input sequences  \( S_0 \) and  \( S_1 \), with sizes  \( m \) and  \( n \), respectively. The DP matrix  \( H \) is calculated as follows. The first row and column of  \( H \) are filled with  \(-g \cdot i\), where  \( i \) is the size of the non-empty sequence and  \( g \) is the gap penalty. The remaining cells are calculated with Equation 1, where  \( ma \) and  \( mi \) are respectively the punctuation for match (\( S_0(i) = S_1(j) \)) and mismatch (\( S_0(i) \neq S_1(j) \)). Each cell keeps an indication of the cell that was used to produce the value (arrows in Figure 1). The optimal global score is the value contained in  \( H_{m,n} \). To retrieve the alignment, a traceback procedure is executed from the bottom right cell, following the arrows until the top left cell is attained (Figure 1 (a)).

\[
H_{i,j} = \max \begin{cases} H_{i-1,j-1} + (\text{if } S_0[i] = S_1[j] \text{ then } ma \text{ else } mi) \\ H_{i-1,j} - g \\ H_{i,j-1} - g \end{cases}
\]

The SW algorithm [Smith and Waterman 1981] is used to obtain the optimal local alignment. It is similar to NW, with three differences. First, the initial row and column are filled with zeroes. Second, Equation 2 is used to compute the cells. Third, the traceback starts in the cell that has the optimal local score (highest value in  \( H \)) and stops when a zero-valued cell is reached (Fig. 1 (b)).

Fig. 1. DP matrices and alignments for  \( S_0 \) and  \( S_1 \) (\( mi=-1, ma=+1, g=-2 \)) using NW and SW.
**Hi,j** = \(\max\ \begin{cases} 
H_{i-1,j-1} + \text{if } S_0[i] = S_1[j] \text{ then ma else mi} \\
H_{i-1,j} - g \\
H_{i,j-1} - g \\
0 
\end{cases}\) \(2\)

NW and SW assign a constant cost \(g\) to gaps. However, gaps tend to occur together. So, a higher penalty is associated to the gap opening and a lower penalty to the gap extensions. This is called the **affine-gap** model, which calculates 3 DP matrices: \(H\), \(E\) and \(F\) (Equations 3, 4 and 5), where \(E\) and \(F\) keep track of gaps in each sequence [Gotoh 1982].

\[
H_{i,j} = \max \begin{cases} 
H_{i-1,j-1} + \text{if } S_0[i] = S_1[j] \text{ then ma else mi} \\
E_{i,j} \\
F_{i,j} \end{cases} \]

\(3\)

\[
E_{i,j} = \max \begin{cases} 
E_{i,j-1} - G_{ext} \\
H_{i,j-1} - G_{open} \end{cases} \]

\(4\)

\[
F_{i,j} = \max \begin{cases} 
F_{i-1,j} - G_{ext} \\
H_{i-1,j} - G_{open} \end{cases} \]

\(5\)

### 2.2. Semi-global alignments

The algorithm for semi-global alignment mixes the matrix initialization and NW/SW recurrence relations in order to ignore gap penalties in the head and/or tail of the sequences. [Durbin et al. 2002] define the overlap alignment as a special semi-global case where the alignment starts in the first row or first column and ends in the last row of last column of the matrix. [Liu and Schmidt 2014a] additionally define semi-global alignment as an alignment that starts in the first row and ends in the last row, or, symmetrically, starts in the first column and ends in the last column.

In general terms, an alignment type may combine other possibilities, using different initialization formulae and optimal score retrieval positions. So, we can classify the alignment types considering where the alignment may start and where it may end. The following paragraphs define symbols for each edge possibility and state the changes that must be done in the alignment algorithm.

**Regarding the beginning of the alignment:**

 (+) **in the first cell:** initializes the first row and column with gap penalties and uses the NW recurrence relation;

 (1) **in the first row:** initializes the first row with zeroes and the first column with gap penalties and uses the NW recurrence relation;

 (2) **in the first column:** initializes the first row with gap penalties and the first column with zeroes and uses the NW recurrence relation;

 (3) **in the first row or column:** initializes the first row and column with zeroes and uses the NW recurrence relation;

 (*) **anywhere in the matrix:** initializes the first row and column with zeroes and uses the SW recurrence relation;

**Regarding the end of the alignment:**

 (+) **in the last cell:** optimal score resides exactly at the last cell;

 (1) **in the last row:** seeks for the optimal score in the last row;

 (2) **in the last column:** seeks for the optimal score in the last column;
(3) **in the last row or column:** seeks for the optimal score in the last row or column;
(*) **anywhere in the matrix:** seeks for the optimal score in any cell of the matrix;

In this way, we may have $5 \times 5 = 25$ possible alignment types. We will represent the alignment type with the symbol $(b/e)$ when beginning at position $b$ and ending at position $e$. For instance, type $(2/\ast)$ starts in the first column and ends anywhere in the matrix. Some combinations are already defined as a global alignment $(+/+)$ and local alignment $(\ast/\ast)$. All other combinations may be classified as specific semi-global alignment types, for instance, the overlap alignment $(3/3)$.

### 2.3. Linear Space Algorithms

Many SW/NW applications only retrieve the score. In this case, the traceback procedure is not executed and space complexity is linear. To retrieve the alignment with SW/NW for huge sequences, quadratic space is required and, for this reason, several Petabytes of memory may be needed. Therefore, another approach must be used.

The algorithm proposed by [Hirschberg 1975] retrieves alignments in linear space. It was further adapted by Myers-Miller (MM) [Myers and Miller 1988] to the affine-gap model. MM is an NW variant that retrieves alignments in linear space with affine-gaps. It uses a divide and conquer technique with a matching procedure to find one point where the optimal alignment occurs and recursively splits the DP matrix to obtain the alignment. This approach can double the execution time, in the worst case [Myers and Miller 1988], when compared with NW.

In order to apply MM to local alignments, the beginning and the end positions of the local alignment must be previously found with the SW algorithm. With these positions, the problem can be treated as a global alignment and, thus, the MM algorithm can be applied. [Liu et al. 2009] is one of the first proposals that apply the MM algorithm for local alignments in GPUs.

### 2.4. Parallel SW/NW

In SW/NW and its variants, most of the time is spent calculating the DP matrices and this is the part which is usually parallelized. Each cell $(i, j)$ of the DP matrix depends on three other cells: $(i-1, j)$, $(i-1, j-1)$ and $(i, j-1)$ (Equations 1 to 5). Respecting these data dependencies, the DP matrix can be processed in different ways (Figure 2): by row (a), by column (b), by antidiagonal (c), by wave-square (d), by wave-antisquare (e) or in a generic order (f).

Considering the execution orders presented in Figure 2, the antidiagonal strategy, also called wavefront [Pfister 1995], is a very straightforward and commonly used method that permits parallel computation of the cells. Since the cells in the same antidiagonal do not have direct dependencies among them, all cells in each diagonal can be computed in parallel. The linear strategies presented in Figures 2(a) (rows) and 2(b) (columns) are also suited for parallel execution using prefix computations [Aluru et al. 2003; Rajko and Aluru 2004]. Square wave strategies (Figures 2(d) and 2(e)) alternate the computation of rows and columns in the same execution. Finally, the DP matrix can be processed in a generic order as presented in Figure 2(f), allowing cells in different rows, columns or antidiagonals to be processed simultaneously. This generic execution order is well suited for dynamic schedulers that resolve task dependencies at runtime. This strategy may generate irregular computation patterns with reduced synchronization points, while still respecting the data dependencies.

### 3. RELATED WORK

In the last decades, several strategies have been proposed to efficiently execute biological sequence comparison algorithms in HPC platforms. Nowadays, HPC platforms are
usually composed of CPUs and accelerators such as FPGAs (Field Programmable Gate Arrays), CellBEs (Cell Broadband Engines), GPUs (Graphics Processing Units) and, more recently, Intel Phis. In the literature, we can find more than 50 papers in this research field and it is not our intention to discuss extensively all these works. Therefore, in the following paragraphs we will discuss solutions that: (a) calculate more than one type of alignment; (b) use multiple hardware/software platforms; and (c) compare Megabase sequences.

A parallel CellBE approach to retrieve optimal global, syntenic and spliced alignments is proposed in [Sarje and Aluru 2009]. The MM algorithm (Section 2.3), combined with Parallel Prefix (PP) computations and antidiagonal parallelizations (Figure 2(c)), is used to retrieve these 3 types of alignments. [Sarkar et al. 2010] designed an Application-Specific Integrated Circuit (ASIC) composed of tiny Processing Elements (PEs) integrated through a NoC (Network-on-Chip) to retrieve global, local and semi-global alignments. A unidirectional systolic array was designed that can calculate alignments with either the original SW/NW equations (Section 2.1) or with Parallel Prefix (PP). Each PE was implemented in RTL (Register-Transfer Level) and the switches were designed with Candence Spectra Tools, generating a custom hardware. [Maleki et al. 2014] used linear algebra to transform DP problems, including local, global and sequence-profile comparisons, into independent sets of matrix-vector multiplications (MVM) and predecessor products, using the concept of rank convergence to obtain optimal alignments in CPU. The algorithm executes iteratively, stage by stage, where each stage is executed in parallel. To compute local alignments, Farrar’s algorithm [Farrar 2007], which uses SSE (Streaming SIMD Extension) vector instructions that process according to the striped pattern with query profile, was used inside each stage. [Liu and Schmidt 2014a] proposed GSWABE, a GPU strategy that computes global, semi-global and local alignments with the Gotoh algorithm (Section 2.1) that compares short DNA reads with whole genomes. In this strategy, the DP matrix is divided into tiles of size 4x4 and each thread executes a different comparison, using antidiagonal parallelism (Figure 2(c)).

In [Aldinucci et al. 2010], the Farrar SSE implementation [Farrar 2007] was re-implemented according to the dataflow model (Figure 2(f)) and ported to multiple high-level programming environments for shared memory multicores (OpenMP, Cilk, TBB and FastFlow). [Benkrid et al. 2012] create different SW codes to locally compare pro-

![Fig. 2. Different orders to calculate the DP matrix.](image-url)
tein sequences which are highly optimized for different hardware platforms (FPGA, GPU and CellBE) programmed respectively in Handel-C, CUDA and Cell SDK. The FPGA and GPU implementations are fine-grained and explore the antidiagonal parallelism (Figure 2(c)) whereas the CellBE implementation is coarse-grained, i.e., each thread compares the same query sequence to a different sequence. [Liu et al. 2013] proposed CUDA SW++ 3.0, which uses simultaneously CPUs and GPUs to compare protein sequences. Static load distribution based on clock frequencies, number of cores (CPUs) and number of SMs (GPUs) is used to assign work to the CPUs or GPUs. The CPU code is based on SWIPE [Rognes 2011], and the GPU code uses CUDA PTX SIMD instructions combined with the ideas of [Farrar 2007] to accelerate the computation. In [Hamidouche et al. 2013], the authors propose the use of a high-level parallel programming library called BSP++ [Hamidouche et al. 2010] to execute local comparisons in several HPC platforms. A fine-grained parallel implementation for a cluster of CellBEs and multicores is proposed that uses a block-cyclic approach to distribute square partitions of the DP matrix among the processing units.

A parallel exact algorithm that generates global alignments for Megabase sequences was proposed in [Rajko and Aluru 2004] for a cluster of CPUs. The key idea of this algorithm is to use Hirshberg’s algorithm (Section 2.3) combined with Parallel Prefix (PP) computations to find a partial balanced partition between subsequences of $S_0$ and $S_1$. This partition allows the subdivision of the original problem in independent subproblems that can be solved in parallel. CUDAAlign 2.1 was proposed by [Sandes and Melo 2013b] and it is a combination of the Gotoh (Section 2.1) and MM algorithms (Section 2.3), retrieving optimal local alignments in linear space for Megabase sequences in GPU. It computes optimal local alignments in 5 stages, where stage 1 obtains the optimal score with antidiagonal parallelism (Figure 2(c)) and the block pruning optimization, saving some rows to disk. Stages 2 to 5 implement the traceback, executing a modified version of MM, retrieving the coordinates of the points that belong to the optimal local alignment in a divide-and-conquer way. [Korpar and Sikic 2013] proposed SW#, which is an approach that implements the MM algorithm (Section 2.3) with the parallelization strategy and the block pruning optimization proposed in CUDAAlign 2.1 [Sandes and Melo 2013b] for retrieving the local alignment between Megabase DNA sequences in GPU. [Liu et al. 2014] proposed SWAPHI-LS, a strategy to execute local comparisons of Megabase sequences with one or more Intel Phis. As output, the optimal score is provided. The DP matrix is divided into blocks which are processed by antidiagonals (Figure 2(c)). Each block is further divided into small tiles of fixed size, which are distributed to a team of threads. Each thread calculates its tile in a vectorized way, using SIMD instructions.

Table I presents a comparative view of the papers discussed in this section. In this table, we present the paper, the maximum query sequence size used in the experimental results, the supported alignment types, the output produced and, finally, the processing units/programming environments used in each paper. In Table I, the cited papers grouped in three different classes. In the “Multiple Type of Alignments” group, the papers were able to obtain the alignments for local, global, semi-global, synthenic and sequence profile comparison types. Although they used different types of hardware (CellBE, ASIC, CPU and GPU), these papers restricted the length of query sequences to some thousands of characters. In the “Multiple HW/SW Platforms” group, the papers used many platforms simultaneously (CPU, FPGA, CellBE, GPU) and different frameworks (OpenMP, Cilk, TBB, FastFlow, Handle-C, Cell SDK, CUDA, BSP++) to obtain the score of the optimal local alignment, without retrieving the full alignment. The maximum sequence sizes were greater than the previous group, increasing to sequences of more than a million base pairs. The group “Comparison of Megabase Sequences” present papers that aligned sequences larger than 1 Mbp. They all retrieve
Table I. Comparative View of HPC Biological Sequence Comparison Papers

<table>
<thead>
<tr>
<th>Paper</th>
<th>Max Size</th>
<th>Alignment Type</th>
<th>Output</th>
<th>Processing Units (Programming)</th>
</tr>
</thead>
<tbody>
<tr>
<td>[Sarje and Aluru 2009]</td>
<td>$10^4$</td>
<td>local, spliced, synt</td>
<td>score, align</td>
<td>CellBE (Cell SDK)</td>
</tr>
<tr>
<td>[Sarkar et al. 2010]</td>
<td>$10^4$</td>
<td>local, global, semi-g</td>
<td>score, align</td>
<td>ASIC (RTL and Cadence)</td>
</tr>
<tr>
<td>[Maleki et al. 2014]</td>
<td>$10^4$</td>
<td>local, global, seq-prof</td>
<td>score, align</td>
<td>GPU (SSE and MPI)</td>
</tr>
<tr>
<td>[Liu and Schmidt 2014a]</td>
<td>$10^4$</td>
<td>local, global, semi-g</td>
<td>score, align</td>
<td>GPU (CUDA)</td>
</tr>
<tr>
<td>[Aldinucci et al. 2010]</td>
<td>$10^4$</td>
<td>local</td>
<td>score</td>
<td>CPU (OpenMP or Cilk or TBB or FastFlow)</td>
</tr>
<tr>
<td>[Benkrid et al. 2012]</td>
<td>$10^3$</td>
<td>local</td>
<td>score</td>
<td>FPGA (Handel-C)</td>
</tr>
<tr>
<td>[Liu et al. 2013]</td>
<td>$10^3$</td>
<td>local</td>
<td>score</td>
<td>GPU (CUDA)</td>
</tr>
<tr>
<td>[Hamidouche et al. 2013]</td>
<td>$10^3$</td>
<td>local</td>
<td>score</td>
<td>CellBE (BSP++)</td>
</tr>
<tr>
<td>[Hamidouche et al. 2013]</td>
<td>$10^7$</td>
<td>local</td>
<td>score</td>
<td>GPU (CUDA)</td>
</tr>
<tr>
<td>[Liu et al. 2013]</td>
<td>$10^7$</td>
<td>local</td>
<td>score</td>
<td>GPU (CUDA)</td>
</tr>
<tr>
<td>[Liu et al. 2014]</td>
<td>$10^7$</td>
<td>local</td>
<td>score</td>
<td>Phi (IMCI)</td>
</tr>
<tr>
<td>This work (MASA)</td>
<td>$10^7$</td>
<td>local, global, semi-g</td>
<td>score, align</td>
<td>GPU (MASA and CUDA)</td>
</tr>
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<td></td>
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<td></td>
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<td>Phi (MASA and OpenMP)</td>
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<td>CPU (MASA and OpenMP)</td>
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<td></td>
<td></td>
<td>CPU (MASA and OmpSs)</td>
</tr>
</tbody>
</table>

the optimal local score and many of them were able to retrieve the full alignment. The target platforms were CPU (using MPI), CUDA GPU’s and the Intel Phi.

Our work (last row in Table 1) is able to compare sequences with more than 10 Mbp, as do the papers in the third group, generating global, local or semi-global alignments as do the papers in the first group. Differently from other works, in MASA we are able to reuse the features/optimizations that belong to the platform-independent code in different implementations, enhancing the portability to other HPC platforms and potentially reducing the time to produce the codes.

4. THE CUDALIGN ALGORITHM

Fig. 3. The CUDAlign Algorithm. (a) Stage 1: finds the optimal score and its position. Special rows are saved and some blocks are pruned (gray); (b) Stage 2: computes crosspoints between optimal alignment and special rows. Special columns are saved; (c) Stage 3: finds more crosspoints inside partitions; (d) Stage 4: executes Myers-Miller (MM) algorithm between successive crosspoints; (e) Stage 5: aligns partitions and obtains full alignment. Stage 6 is not represented in the figure since it is optional and it only creates graphical and textual versions of the alignment.

In this section, we give an overview of CUDAlign 2.1, that was used to generate the MASA code. CUDAlign 2.1 [Sandes and Melo 2013a] is a GPU algorithm that retrieves
the optimal local alignment between two Megabase DNA sequences with affine gaps in linear space. In order to find the alignment, CUDAlign combines Gotoh and MM (Sections 2.1 and 2.3) and iteratively obtains coordinates of the optimal alignment, which are incrementally refined until the full alignment is obtained.

CUDAlign 2.1 is computed in 6 stages (Figure 3). The first three stages execute in GPU and Stages 4, 5 and 6 execute in CPU. In general terms, the algorithm calculates an increasing number of crosspoints in Stages 1 to 4, and Stage 5 is employed to align and concatenate results from all partitions. Further, Stage 6 is optionally used for visualization of the alignment.

Stage 1 is the most compute-intensive phase of CUDAlign since it calculates the Gotoh's matrix in linear space. Its goal is to retrieve the optimal score and the position in the DP matrix where it occurs. This position marks the end of the optimal alignment. An optimization called block pruning (BP) [Sandes and Melo 2013a] is executed in this stage to reduce the number of cells calculated. A pruned block has such a small departing score that it is mathematically impossible to exist an optimal alignment that crosses it. Thus, a pruned block does not contribute to the optimal score and therefore does not need to be calculated. Also in this stage, some rows (called special rows) are saved to disk in order to accelerate the computation in the next stages. This strategy is derived from FastLSA [Driga et al. 2006]. The number of special rows is a parameter of CUDAlign, chosen by the user. As the number of special rows increases, the area processed by Stages 2 and 3 is reduced. Figure 3(a) illustrates Stage 1.

Stage 2 computes the semi-global recurrence relation using the MM matching procedure (Section 2.3) in the reverse direction of Stage 1, starting at the end point of the alignment obtained in Stage 1. The main goal of Stage 2 is to find all the points that belong to the optimal alignment and cross the special rows (crosspoints), including the start point. The orthogonal execution optimization [Sandes and Melo 2011] is used to reduce the area calculated by the MM matching procedure. Special columns are saved to disk. Figure 3(b) presents Stage 2.

Stage 3 is very similar to Stage 2 (Figure 3(c)). The main difference is that Stage 3 uses the special columns saved to disk in the previous stage to compute more crosspoints. These new crosspoints are found inside the partitions (area delimited by consecutive crosspoints) obtained in Stage 2.

Stage 4 is a multi-threaded version of MM (Section 2.3) that computes the points that belong to the alignment inside partitions found in Stage 3. It adds more crosspoints to the solution until very small partitions are obtained (Figure 3(d)).

Stage 5 executes the NW algorithm (Section 2.1) to align the partitions formed by crosspoints found in Stage 4 and concatenates the results to obtain the full optimal alignment (Figure 3(e)).

Stage 6 is an optional stage used only for visualization purposes.

5. DESIGN OF MASA

The main goal of MASA is to provide a flexible and customizable infrastructure to develop sequence aligners in multiple hardware/software platforms. It proposes and implements a set of platform-independent modules which may be reused by multiple platform-specific implementations. It also allows platform-independent optimizations to be deployed at once to many platform-specific implementations.

In order to design MASA, we analyzed the code of CUDAlign 2.1 (Section 4) to determine which parts of the code are platform-independent or platform-specific. Most of the execution time of CUDAlign is spent in CUDA kernels, calculating the recurrence relation (SW/NW), and this part is platform-specific. There are other parts that are platform-independent, such as input/output operations, profiling and stage coordination.
Fig. 4. Overview of MASA: (a) presents the MASA modules, including platform-independent optimizations that are selected by the developer, such as Block Pruning and Parallelization strategy, as well as the platform-specific code executed on the target processor to compute the NW/SW recurrences; (b) shows multiple Aligners we developed on top of MASA with their choice of block pruning and parallelization strategies.

We also analyzed Block Pruning and the Parallelization Strategy. Since Block Pruning (BP) (Section 4) is the most relevant CUDAlign optimization, we decided to implement it in a platform-independent way, with sufficient flexibility to be used even when a generic block execution order is used (Figure 2(f)). In the current version of MASA, Block Pruning works for local alignments, as in the previous versions of CUDAlign. The parallelization strategy employed by CUDA depends on the GPU hardware, and thus is platform-specific. Nevertheless, other implementations may benefit from parallelization strategies. Thus, as with BP, we decided that the parallelization strategy should be implemented in a platform-independent way. We call “MASA implementation” the union of the platform-specific and platform-independent codes, with selected customizations. Each implementation creates a single binary file that can be executed in a given platform.

Although the MASA code was originally based from the CUDAlign 2.1 code, new features were also included in a platform-independent way. For instance, CUDAlign 2.1 only stores special rows in disk, but MASA can store rows in memory and disk, with simultaneous usage if necessary. Multinode support was also included in Stage 1, with the ideas presented in [Sandes et al. 2014b], with heterogeneous platform support [Sandes et al. 2014a]. Stage 3 was also redesigned in order to iterate many times if the partitions are still too large for Stage 4. Furthermore, MASA can also produce the optimal alignment based on any of the 25 types defined in Section 2.2. With all these features, we can note the benefit of using the MASA platform, since a new feature can be easily delivered to all different implementations.

5.1. MASA Architecture

The architecture of MASA is divided in 5 modules, as shown in Figure 4(a). Data Management, Statistics and Stage Management are modules used by all implementations. Block Pruning and Parallelization Strategy are customizable modules and the developer can choose the strategies provided by MASA or provide his/her own strategy. The platform-specific code calculates the SW/NW equations.

Data Management: This module is responsible for managing data such as input sequences, user parameters, special rows/columns, optimal alignment and score. Once a MASA implementation is dispatched for execution, it queries this module to retrieve the data used as input as well as to store results.
Statistics: This module provides information about the execution such as execution time of each stage, amount of memory/disk used and percentage of blocks pruned, among others.

Stage Management: This module is responsible for coordinating the execution of Stages 1 to 3 and for executing Stages 4 to 6 in CPU. During the execution of Stages 1 to 3, it divides the DP matrix into partitions (Figure 5) and dispatches each partition to be computed by the platform-dependent code. This code receives the first row and first column of a partition and provides as output the last row and last column computed, as well as the highest score found and the special rows/columns. In Stage 1, there is a single partition whose size is the whole matrix whereas in the other stages there are several smaller partitions, creating boundaries over the special rows/columns. When this module receives the border cells of a partition, it starts to search for crosspoints with the MM matching procedure with the orthogonal execution optimization (Section 4) and, as soon as it finds them, the computation of the partition stops and the next partition is dispatched.

Block Pruning (BP): We redesigned the BP optimization proposed in CUDAlign 2.1 in a platform-independent way and called it Diagonal BP in MASA. To perform the pruning, Diagonal BP must keep track of a non-prunable window \([k_s..k_e]\) containing the blocks that must be computed in each diagonal. So, two pointers \(k_s\) and \(k_e\) are stored in memory, being updated after each diagonal computation (Figure 6(a)). Since MASA aims to support multiple implementations of aligners, including generic dataflow parallelization, we have extended Diagonal BP and created a new Generic BP strategy (Figure 6(b)). In Generic BP, we cannot maintain only two pointers as in Diagonal BP because many blocks from different diagonals can be calculated in parallel, in a considerable number of concurrency scenarios. Thus, Generic BP maintains a matrix containing the prunable state of each block. If the neighbors of a block are all prunable, we infer that this block is also prunable, thus expanding the prunable area (Figure 6(b)). This implementation uses \(O(B_h \times B_v)\) of memory, where \(B_h\) is the number of horizontal blocks in the DP matrix and \(B_v\) is the number of vertical blocks. A detailed description of Generic BP is presented in Section 6.

Parallelization strategy: SW, NW and Gotoh’s recurrence relations have the same dependency pattern, i.e., the computation of a cell \((i,j)\) depends on cells \((i-1,j)\), \((i-1,j-1)\), and \((i,j-1)\) (Section 2.4). To achieve better results, cells are grouped in blocks, maintaining the same dependency pattern between the blocks. Considering this, MASA provides two strategies to exploit parallelism: Diagonal and Dataflow.

In the Diagonal method, computations start at the top-left corner block of cells and propagate diagonally (Figure 2(c)). Blocks composing the same diagonal can be computed in parallel. The main limitation here is the frequent synchronization points at the end of each diagonal computation.
Dataflow parallelization is proposed to reduce the synchronization steps of the Diagonal method. In this case, the generic method (Figure 2(f)) is implemented as a dataflow where each node of the dataflow is a block of cells. When the dependencies of a block are resolved, this block is ready for execution. Data dependencies are resolved during the execution, reducing synchronization overheads of the Diagonal method.

5.1.1. MASA-API. The MASA architecture was developed based on the object-oriented paradigm, using the C++ programming language. The MASA-API is presented as a class hierarchy in Figure 7. The IALIGNER class is the interface point between MASA stages and Aligners implementations. Each MASA implementation must create its own Aligner class that implements the IALIGNER virtual methods. The Aligner implementation communicates with the stages using an IMANAGER implementation, that contains 4 type of methods: GET methods, which return alignment parameters; MUST methods, that dictate runtime behavior for the Aligner; RECEIVE methods, which transfer to the Aligner initial rows and columns of the matrix; and DISPATCH methods, that send matrix cells to the MASA stages modules. Some of the IMANAGER methods are listed in Table II.

Each IALIGNER implementation must implement some methods in order to be instantiated. The INITIALIZE and FINALIZE methods are called only once during the alignment lifecycle and they must be used to initialize and finalize any required resource (e.g. memory and accelerator hardware) for the alignment execution. Then, each stage calls the IALIGNER::SETSEQUENCE and IALIGNER::UNSETSEQUENCE to define the sequence interval and direction that will be used during the stage. During the stage execution, one or more partitions are aligned, and the aligner executes this job in the IALIGNER::ALIGNPARTITION method.

In order to simplify the creation of an IALIGNER subclass, there is a class hierarchy with different types of aligners, which are represented in Figure 7 as yellow classes. The ABSTRACTALIGNER class encapsulates the IMANAGER methods and initializes the grid and the block pruning operations. Block pruning is handled by the ABSTRACTBLOCKPRUNING class and its subclasses, shown in Figure 7 as blue classes. Then, there are two other types of aligners: ABSTRACTBLOCKALIGNER and ABSTRACTDIAGONALALIGNER.

ABSTRACTBLOCKALIGNER calculates the matrix in a block basis, using the GENERICBP block pruning strategy. Each ABSTRACTBLOCKALIGNER subclass (OpenMP and OmpSs Aligners) must implement its own scheduler mechanism and each block is pro-
cessed by a subclass of an AbstractBlockProcessor, represented in Figure 7 as green classes. The CPUBlockProcessor is a subclass that processes blocks using the conventional SW/NW code in CPU. Other Block Processors could, for instance, use FPGA or SSE instructions to process the blocks.

The AbstractDiagonalAligner class calculates the DP matrix with the wavefront strategy, using the DiagonalBP block pruning strategy. The CUDAligner subclass extends this subclass and, whenever the processDiagonal method is called, it executes a new diagonal of blocks in the GPU. Although the AbstractBlockAligner can also implement a scheduler mechanism for diagonal processing, the AbstractDiagonalAligner allows the entire diagonal to be executed at once in the GPU. Thus, the CUDA architecture is able to schedule the blocks using its own scheduler.

![Diagram](image-url)  
Fig. 7. MASA-API: Class Diagram.
### 5.2. Creating a MASA implementation

To create a new MASA implementation, the programmer develops an aligner class (which must implement the IALIGNER interface or extend one of the abstract aligner classes) and supplies one instance of this class to the MASA entry point. Then, the platform-independent code processes the arguments, reads the sequences and coordinates the stages execution. When the SW/NW equation needs to be computed, the aligner object is invoked (alignPartition procedure). It receives the boundary coordinates of the partition, the first column/row and computes the SW/NW equation. Then, the aligner uses MASA functions to inform (dispatch) the best score, the special rows and the last column of the partition.

Algorithm 1 illustrates a simplified MASA implementation based on the ABSTRACT-BLOCKALIGNER. The concrete aligner class will simply be called ALIGNER. The basic memory initialization, the grid partitioning and the block pruning (BP) initialization are transparently done by the ABSTRACT-BLOCKALIGNER initialization. Then, the specific scheduleBlocks method (lines 2-6) iterates through the grid per diagonal and calls alignBlock for each block (line 4). The alignBlock procedure receives from MASA the first row (line 10) or column (line 11) of the neighbor blocks. The BP test is made by MASA (line 12) and, if this block is not pruned, the aligner calls processBlock (line 13) in order to compute the SW/NW recurrence relation. The best score found is dispatched to MASA (line 14). Special rows (line 16) and the last column (line 17) are also dispatched to MASA. The program entry point (lines 20-23) creates the aligner object and passes it to the MASA entry point (line 22), using the default CPUBLOCKPROCESSOR (line 21).

The processBlock method is implemented in the ABSTRACT-BLOCKALIGNER class and it essentially reads the first row/column from the block.row and block.col arrays, computes the SW/NW recurrence relation and stores the last row/column into the same block.row and block.col arrays. Rows and columns of the blocks are chained so that the last row/column of a block is the first row/column of the next block. The processBlock procedure is compute-intensive, and this code is very suitable for platform-specific optimization. Furthermore, other third party tools that execute optimized DNA sequence comparison may be adapted to MASA. This can be done with the reimplementation of the processBlock method using the recurrence relation source code of the third party tool, with some modifications to fit the processBlock input/output parameters.

### 5.3. MASA Implementations

In this section, we present our four MASA implementations, which use different programming models/tools (OpenMP, OmpSs and CUDA) and target different hardware platforms (multicore, GPU, Intel Phi). Each one of our MASA implementations (Figure 4(b)) used the affine gap model (Section 2.1) and present some modification in Algo-

---

**Table II. Some IManager Methods**

<table>
<thead>
<tr>
<th>Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>GETRECURRENCETYPE</td>
<td>Recurrence type: SW or NW</td>
</tr>
<tr>
<td>GETSPECIALROWINTERVAL</td>
<td>Minimum Distance between special rows</td>
</tr>
<tr>
<td>RECEIVING</td>
<td>Receives the first row of a partition</td>
</tr>
<tr>
<td>RECEIVINGFIRSTROW</td>
<td>Receives the first column of a partition</td>
</tr>
<tr>
<td>RECDATAXCORE</td>
<td>Outputs special rows of the partition</td>
</tr>
<tr>
<td>RECEIVEFIRSTCOLUMN</td>
<td>Outputs special columns of the partition</td>
</tr>
<tr>
<td>DISSECONDROW</td>
<td>Outputs the best score found in each block</td>
</tr>
<tr>
<td>MUSTCONTINUE</td>
<td>If false, the Aligner must stop execution</td>
</tr>
<tr>
<td>MUSTPRUNEBLOCKS</td>
<td>Defines if the Aligner can prune blocks</td>
</tr>
</tbody>
</table>
Algorithm 1 Aligner Pseudocode - Block Based

1: procedure ALIGNER::SCHEDULE BLOCKS
2:  for each diagonal do
3:      for each block in diagonal do
4:         ALIGN BLOCK(block)
5:  end for
6:  end for
7: end procedure
8: 
9: procedure ALIGNER::ALIGN BLOCK(block)
10:  if block is in first row then block.row ← RECEIVE FIRST ROW
11:  if block is in first column then block.col ← RECEIVE FIRST COLUMN
12:  if Not IS BLOCK PRUNED(block) then
13:      block.score := PROCESS BLOCK(block)
14:      DISPATCH SCORE(block.score)
15:  end if
16:  if isSpecialRow(block.row) then DISPATCH ROW(block.row)
17:  if block is in last column then DISPATCH COLUMN(block.col)
18: end procedure
19: 
20: procedure MAIN(args)
21:  processor = new CPUBlockProcessor()
22:  MASA::ENTRY POINT(args, new Aligner(processor))
23: end procedure

To give an idea in terms of number of code lines (excluding blank lines and comments), the platform specific code contains 116 (OpenMP), 187 (OmpSs) and 1493 (CUDAAlign) lines, whereas the platform independent source code contains more than 15,000 lines.

5.3.1. MASA-OpenMP/CPU. MASA-OpenMP/CPU uses OpenMP to compute blocks that belong to the same diagonal in parallel. In Algorithm 1, "#pragma omp parallel for schedule(dynamic,1)" is inserted before line 3. Dynamic scheduling is used because the size of the parallel loop is usually larger than the number of threads. The processBlock function is a CPU implementation of SW/NW, without vectorized instructions (SIMD).

5.3.2. MASA-OpenMP/Phi. MASA-OpenMP/Phi employs the same parallelization strategy as MASA-OpenMP/CPU, where independent threads process each diagonal in parallel. It also uses the dynamic OpenMP scheduler, since this option leads to better performance. Our implementation uses the Intel Phi native execution mode, i.e., the entire application (including MASA) runs within the coprocessor. This was possible because Intel Phi runs a specialized Linux kernel that provides the necessary OS level services.

Because Intel Phi is equipped with 512-bit wide SIMD instructions, we have tried to modify the align partition code, as compared to the MASA-OpenMP/CPU, so that computation of cells in a diagonal are an internal loop in the computation. With this modification, we were able to leverage the Intel compiler tools and vectorize this operation. However, the vectorized version of our Phi based code did not attain significant gains on top of the non-vectorized code that was generated from a cross-compilation of the MASA-OpenMP/CPU code. This occurred because the modified code used for vectorization has a larger number of instructions and branch statements in the inner loop as compared to the original code. In special, the branch statements may strongly limit improvements with vectorization [Tian et al. 2013]. This finding is similar to that of [Farrar 2007], which proposed a striped version of SW with no branching in the inner computation loop in order to maximize improvements with vectorization. Still, our Intel Phi based aligner attains good performance and compares well to other CPU-based implementations. Therefore, we use this implementation to demonstrate...

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the MASA flexibility. As a future work, we intend to port the Farrar striped NW algorithm [Farrar 2007] to Intel Phi, such as proposed in [Liu and Schmidt 2014b], and include it in MASA.

5.3.3. MASA-OmpSs/CPU. MASA-OmpSs/CPU uses the OmpSs parallel programming environment [Duran et al. 2011] and therefore its dataflow-based parallelization.

OmpSs proposes a unified programming model for heterogeneous systems, incorporating ideas of OpenMP and adding support for irregular and asynchronous parallelism. OmpSs applications are described as dataflows and use several optimizations, including efficient and automatic data movement, performed at compiler and runtime system levels. Parallelization of an application in OmpSs is carried out with a set of "#pragma" directives. The programmer defines portions of code that should be executed as tasks in the dataflow. The dependencies among tasks are derived from pragma options defining the task variables, which can be used as input and/or output. OmpSs applications also benefit from several task scheduling policies.

In MASA-OmpSs/CPU, the code sections that invoke processBlock, dispatchRow and dispatchColumn methods are annotated as OmpSs tasks, so that the OmpSs compiler can create the appropriate structures to dispatch the tasks. The same annotation also includes hints about the type of data used (inout or output). With this, data dependencies among tasks are identified and used by the runtime system to guarantee correctness in the execution.

MASA-OmpSs/CPU uses the dataflow parallelization strategy (Figure 4(b)). Each call to alignBlock (Algorithm 1 lines 9-18) creates up to three tasks: (1) processBlock task: lines 12-15; (2) dispatchRow task: line 16; and (3) dispatchColumn task: line 17. Task dependencies are created considering the block.row and block.col vectors. Since OmpSs is able to execute blocks in generic order (Figure 2(f)), we set the block priorities to create a preferable execution order in square waves (Figure 8(a)). Considering block \((bx, by)\), its priority is \(\min(-bx, -by)\). The OmpSs hysteresis throttle mechanism was enabled to limit the number of tasks in the task graph, reducing the amount of memory used by OmpSs. When the number of tasks per thread reaches a limit, task creation stops until it drops below a given value. Our goal is to keep a sufficient level of parallelism while reducing the amount of memory used by OmpSs. To prevent loss of parallelism during the hysteresis drop down phase, the task creation order was changed to follow the priority order. So, tasks are created in lanes (Figure 8(b)). The width of the lanes is the number of parallel threads, which is equal to the number of cores in the running environment. Inside each lane, the block creation order follows a diagonal creation (third lane in Figure 8(b)).

5.3.4. MASA-CUDAlign. Since CUDAlign was created before MASA, the original CUDAlign 2.1 code was basically kept the same as in [Sandes and Melo 2013a], with
some modifications to fit in the aligner classes hierarchy. Instead of using the `AbstractBlockAligner`, MASA-CUDAlign uses the `AbstractDiagonalAligner` as a base class. Thus, a `processDiagonal` method is called for each diagonal and the CUDAAligner class launches a kernel in GPU. All the memory allocation and transfers are made by the aligner class, so much more control is needed by this class. Calls to MASA functions (i.e. dispatch and receive methods) are always made by the CPU, usually preceded/followed by `cudaMemcpY` calls to transfer data from/to the GPU.

6. GENERIC BLOCK PRUNING

The block pruning optimization was designed to avoid the calculation of blocks of cells that do not contribute to the optimal local alignment. [Sandes and Melo 2013a] have proposed a block pruning technique for antidiagonal execution order, whereas in this paper we propose and implement block pruning for generic execution order.

The antidiagonal block pruning must keep track of a non-prunable window \([k_s..k_e]\) containing the blocks that must be computed in the current diagonal. As such, the pointers \(k_s\) and \(k_e\) are stored in memory, and they are updated after each diagonal computation (Figure 6(a)). In the generic execution order, blocks from multiple diagonals may be computed concurrently and, as a consequence, keeping those two pointers is not sufficient (Figure 6(b)). Therefore, we have created a more elaborate block pruning strategy to work with generic execution order, which is required by the novel parallelization strategies proposed and implemented in MASA.

6.1. Definitions

Let \(S_0\) and \(S_1\) be the sequences being aligned, with sizes \(|S_0| = m\) and \(|S_1| = n\). The punctuation for matches and mismatches are, respectively, \(ma\) and \(mi\). Suppose that cell \((i,j)\) of the DP matrix has score \(H(i,j)\). The maximum score of any alignment that passes through cell \((i,j)\) is defined in Equation 6,

\[
H_{\text{max}}(i,j) = H(i,j) + H_{\text{inc}}(i,j)
\]

where \(H_{\text{inc}}(i,j)\) is the incremental score considering a matching of all remaining characters from subsequences \(S_0[i..m]\) and \(S_1[j..n]\). Additionally, the \(H_{\text{inc}}\) function may be calculated using Equation 7 for local alignments, and it is also presented in Figure 9.

\[
H_{\text{inc}}(i,j) = \min(m-i, n-j)ma
\]

Fig. 9. Representation of the cell based pruning definitions.
We also define two special types of cells that are used in the pruning algorithms: 
*prunable* and *pruned* cells. A **prunable cell** is a cell that cannot be part of an alignment with score higher than the maximum score currently known \( (H_{\text{best}}) \). In other words, a cell \((i,j)\) is prunable if \( H_{\text{max}}(i,j) \leq H_{\text{best}} \), since its computation will have no impact on the final optimal score. If cells \((i-1,j)\), \((i-1,j-1)\) and \((i,j-1)\) are already known to be prunable, then it may be inferred that cell \((i,j)\) is also prunable. In this case, cell \((i,j)\) is called a **pruned cell**. As it may be noticed, the **pruned cell** is considered prunable even before its computation.

Pruning conditions for the block-based execution can be similarly derived. Assume that the input DP matrix is divided into \( B_h \times B_w \) blocks. Each block \( b_{i,j} \) depends on blocks \( b_{i-1,j}, b_{i,j-1} \) and \( b_{i-1,j-1} \) (when they exist). For each block, we define \((i',j')\) as the top-left cell coordinates and \((i'',j'')\) as the bottom-right cell coordinates. The height and width of a block \( b_{i,j} \) are given, respectively, by \( \Delta_i \) and \( \Delta_j \). Therefore, the maximum possible score of any alignment that passes through block \((i,j)\) can be defined as shown in Equation 8:

\[
H'_{\text{max}}(b_{i,j}) = H'(b_{i,j}) + H'_{\text{inc}}(b_{i,j})
\]

where \( H'(b_{i,j}) \) is the best score of that block and \( H'_{\text{inc}}(b_{i,j}) \) is the highest \( H_{\text{inc}}(i,j) \) from all cells \((i,j)\) inside the block. An upper bound for \( H'_{\text{inc}}(b_{i,j}) \) may be calculated as follows: considering that the best score of the block is in the top-left cell \((i',j')\), we can define the \( H'_{\text{inc}} \) function as presented in Equation 9:

\[
H'_{\text{inc}}(b_{i,j}) = H_{\text{inc}}(i',j'), \text{ma}
\]

In a similar way to the cells case, a **prunable block** is a block whose cells cannot generate an alignment with score higher than the currently best score found, i.e. \( H'_{\text{max}} \leq H'_{\text{best}} \). Therefore, all the cells of a prunable block will not contribute to the optimal score. A block \((b_{i,j})\) is a **pruned block** if blocks \( (b_{i-1,j}), (b_{i,j-1}) \) and \( (b_{i,j-1}) \) are already known to be prunable. The difference between a prunable and a pruned block is that we need to calculate the score of the first in order to know that the block status is prunable. The pruned block, on the other hand, is known prior to its execution by inference from the status of neighboring blocks.

### 6.2. Generic Block Pruning Procedure

The generic block pruning procedure maintains a matrix \( k \) with an entry per block containing the prunable state of each block. Cell \((i,j)\) of block \( k_{(i,j)} \) is true if block \((i,j)\) is prunable. We assume blocks coordinates starting from 1, i.e., the first block is \((1,1)\). For sake of simplicity, we created an additional row/column in \( k \) (row/column 0) that is initialized with \text{true}, to represent pruned blocks, except for \( k_{0,0} \) that is set to \false in order to force the computation of the first block \((1,1)\). The remaining of the matrix cells are set to \false. The matrix \( k \) is initialized with Equation 10.

\[
\begin{array}{cccccc}
\text{false} & \text{true} & \text{true} & \cdots & \text{true} \\
\text{true} & \text{false} & \text{false} & \cdots & \text{false} \\
\text{true} & \text{false} & \text{false} & \cdots & \text{false} \\
\cdots & \cdots & \cdots & \cdots & \cdots \\
\text{true} & \text{false} & \text{false} & \cdots & \text{false} \\
\end{array}
\]

The matrix initialization is the first step of the generic pruning procedure, as presented in Algorithm 2.

The \text{isPruned} function (lines 2 to 8) is a pre-test invoked before the execution of each block to identify if that block is a pruned one, which would avoid the block computation.
It simply infers the status of the current block \((i,j)\) from its neighbors, and returns true if \(k_{i-1,j}, k_{i-1,j-1}\) and \(k_{i,j-1}\) are true.

If the current block is not pruned, its computation is carried out normally and the \texttt{pruningUpdate} method (lines 14 to 19) is executed after the block calculation to update the status of that block in matrix \(k\). The \texttt{pruningUpdate} method tries to update the current maximum score \((H'_{\text{best}})\) with the score just obtained as the result of the computation of block \(b_{i,j}\).

Further, \texttt{IsPrunable} (lines 9 to 13) is executed to evaluate if block \(b_{i,j}\) could still contribute to the optimal score. In other words, it computes whether block \(b_{i,j}\) is prunable or not, considering its own maximum score \(H'\) and the current overall \(H'_{\text{best}}\) score. The \(H'_{\text{inc}}\) function must be defined according to Equation 9.

**Algorithm 2 Pruning Strategy - Generic**

1: \[ k[0..B_h][0..B_w] \leftarrow \text{Equation 10} \]

2: function \texttt{ISPRUNED}(\(b_{i,j}\))
3: if \((k[i-1][j] \text{ is true}) \text{ and } (k[i][j-1] \text{ is true}) \text{ and } (k[i-1][j-1] \text{ is true})\) then
4: return true;
5: else
6: return false;
7: end if
8: end function

9: function \texttt{ISPRUNABLE}(\(b_{i,j}, H', H'_{\text{best}}\))
10: \((i',j') \leftarrow \text{GETMINCOORD}(b_i, b_j)\)
11: \(H'_{\text{max}} \leftarrow H' + H'_{\text{inc}}(b_{i,j})\)
12: return \(H'_{\text{max}} < H'_{\text{best}}\)
13: end function

14: procedure \texttt{PRUNINGUPDATE}(\(b_{i,j}, H'\))
15: \(H'_{\text{best}} \leftarrow \max(H'_{\text{best}}, H')\)
16: if \texttt{IsPrunable}(\(b_{i,j}, H', H'_{\text{best}}\)) then
17: \(k[i][j] \leftarrow \text{true}\)
18: end if
19: end procedure

The generic block pruning algorithm presented here uses \(O(B_h \times B_w)\) of memory to keep the pruning status of the blocks (size of \(k\)). Considering that the best performance of the algorithm is attained with a number of blocks proportional to the logarithmic of the sequence sizes (i.e. \(B_h = O(\log(m))\) and \(B_w = O(\log(n))\)), in practice, the memory complexity would be \(O(\log(m) \times \log(n))\). Therefore, it results into a moderate memory complexity increase, as compared to the Diagonal BP, with the advantage of allowing a general block execution order.

**7. EXPERIMENTAL RESULTS**

MASA was implemented in C/C++ and MASA implementations (Section 5.3) were programmed in CUDA 4.1, OpenMP 3.0 and OmpSs 1.0. The Minotauro GPU cluster, hosted in the Barcelona Supercomputing Center, was used in our tests. Minotauro is composed of 128 nodes where each node has two 6-core Intel Xeon E5649 and two NVidia Tesla M2090 boards. In our tests, we used only one Minotauro node. For the MASA-OpenMP/CPU and the MASA-OmpSs/CPU executions, we used 12 cores. MASA-CUDAAlign used one GPU NVidia Tesla M2090. For the MASA-Phi executions, we used the Intel Xeon Phi SE10P coprocessor.
Table III. Sequences used in the tests.

<table>
<thead>
<tr>
<th>Cmp.</th>
<th>Sequence 1</th>
<th>Sequence 2</th>
<th>Optimal Score</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Accession</td>
<td>Accession</td>
<td>Local</td>
</tr>
<tr>
<td>10K</td>
<td>AF133821.1</td>
<td>10K</td>
<td>5091</td>
</tr>
<tr>
<td>50K</td>
<td>NC_001715.1</td>
<td>57K</td>
<td>52</td>
</tr>
<tr>
<td>150K</td>
<td>NC_000898.1</td>
<td>172K</td>
<td>18</td>
</tr>
<tr>
<td>50K</td>
<td>NC_003064.2</td>
<td>548K</td>
<td>48</td>
</tr>
<tr>
<td>1M</td>
<td>CP000051.1</td>
<td>1M</td>
<td>88353</td>
</tr>
<tr>
<td>3M</td>
<td>BA000035.2</td>
<td>3M</td>
<td>4226</td>
</tr>
<tr>
<td>5M</td>
<td>AE016879.1</td>
<td>5M</td>
<td>522060</td>
</tr>
<tr>
<td>10M</td>
<td>NC_017186.1</td>
<td>10M</td>
<td>10235188</td>
</tr>
<tr>
<td>23M</td>
<td>NT_033779.4</td>
<td>23M</td>
<td>9063</td>
</tr>
<tr>
<td>47M</td>
<td>NC_000021.7</td>
<td>47M</td>
<td>27206434</td>
</tr>
</tbody>
</table>

7.1. Sequences used in the tests
We compared real DNA sequences (Table III) retrieved from the National Center for Biotechnology Information (NCBI) at www.ncbi.nlm.nih.gov. The sequence sizes vary from 10 Kbp (Thousand base pairs) to 47 Mbp (Million base pairs). The SW/NW parameters used were: \( m_a \) (match): +1; \( m_i \) (mismatch) −3; \( G_{open} \): −5; \( G_{ext} \): −2. For validation purposes, the optimal scores obtained during our tests are also shown for local, overlap, semi-global and global comparisons. In our experiments, the semi-global alignments are defined as the ones that contain all characters of sequence 2 (type (2/2) in Section 2.2), and overlap alignments are defined as in [Durbin et al. 2002] (type (3/3)).

7.2. Execution times for local alignments
In our tests, the default block size for MASA-OpenMP/CPU and MASA-OmpSs/CPU was set to 1024 × 1024, but this size was automatically reduced if the number of blocks in a diagonal is less than \( 2 \times p \), where \( p \) is the number of cores. For MASA-CUDAlign, we used \( T = 128 \) threads, \( B = 512 \) blocks and \( \alpha = 4 \), resulting in a block height of \( \alpha \cdot T \times \frac{n}{B} = 512 \times \frac{n}{512} \), where \( n \) is the width of the partition. For MASA-OmpSs/Phi, we empirically defined the optimal block size for each comparison by increasing the size of block until it does not have any effect in the performance.

Table IV presents the Special Rows Area (SRA) used, the time spent in each stage, the overall runtime, GCUPS, the percentage of pruned blocks and the block size in Stage 1. The overall runtime is the wallclock time, including all initializations and I/O times outside the stages. GCUPS are calculated with \( \frac{m \cdot n \cdot t}{10^9} \), where \( m \) and \( n \) are the sizes of the sequences and \( t \) is the overall runtime. Some implementations did not execute all comparisons due to time constraints. The SRA was set to be stored in the RAM memory up to 4GB, whereas the remaining bytes were saved into the disk.

MASA-CUDAlign presented the best performance for sequences \( \geq 50K \), with GCUPS ranging from 2.31 up to 54.74. These results are comparable with those presented in [Sandes and Melo 2013a], observing that the M2090 GPU is slightly faster than the GTX 560 Ti GPU. This first set of results show that the MASA architecture added negligible overheads to the GPU execution. For the smaller sequence (10K), MASA-CUDAlign is outperformed by other implementations, mainly because of the lack of parallelism to fully utilize the GPU.

The GCUPS varied from 0.57 to 4.81 (MASA-OpenMP/CPU), 0.41 to 5.95 (MASA-OmpSs/CPU) and 0.06 to 4.03 (for MASA-OpenMP/Phi). It is important to note that the MASA-OmpSs/CPU, MASA-OpenMP/CPU and MASA-OpenMP/Phi implementations may be optimized with the use of a striped version of the code that could efficiently leverage optimizations for the vector instructions [Farrar 2007]. Thus, as a future work, we intend to improve these versions by taking advantage of this kind of parallelism.
As expected, comparisons with higher similarity score (Table III) present higher block pruning efficiency. Sequences 10K, 5M, 10M and 47M are very similar, with block pruning rates above 40%. Sequence 1M presents medium similarity, with pruning above 10%, whereas other sequences have very few pruned blocks (below 1%).

The comparison of MASA-OmpSs/CPU (generic BP + dataflow parallel strategy) and MASA-OpenMP/CPU (generic BP + diagonal parallel strategy) execution times shows that the first is more efficient in most of the cases. The better performance of the MASA-OmpSs/CPU aligner is due mainly to the higher pruning rates combined with the flexible parallel execution achieved by the dataflow strategy. For instance, for the 5M and 10M comparisons, MASA-OmpSs/CPU processed 21% less blocks than MASA-OpenMP/CPU, reducing the stage 1 execution time in the same proportion. For the 1M comparison, the GCUPS rate of the MASA-OmpSs/CPU is also better than MASA-OpenMP/CPU, but in only 1% since the sequences are not so similar and the pruning rates are almost the same.

MASA-OmpSs/CPU presented the best pruning rate because the block priorities conducted the execution order to the square wave shape which, in the case of very similar
sequences, obtains the highest scores near the main diagonal much earlier than the traditional diagonal wavefront method. The execution order of the matrix calculation is shown in Figure 10, where we see every 1000 executed blocks in a different shade of gray. The first 100x100 blocks (marked in a square in the figure) were executed similar to square wave, respecting the priorities set to each block. After the 100x100 blocks calculation, we can see that the lanes used during the task creation are much more noticeable. This happens because the hysteresis throttle pauses the task creation whenever the number of queued tasks reaches a maximum value, forcing threads to process nearby blocks, respecting the wavefront inside each lane. As soon as the threads are processing blocks in a diagonal wave inside each lane, this shape tends to persist until the end of the computation.

![Wavefront](image)

**Fig. 10.** MASA-OmpSs/CPU Processing Pattern.

Although the block pruning and parallelization strategies in the MASA-OpenMP/CPU and MASA-OpenMP/Phi are the same, the percentage of pruned blocks varies slightly in some cases. This is due the different block sizes used in both strategies. Specially in the 10K comparison, MASA-OpenMP/Phi achieved the best pruning efficiency compared with the other implementations since the blocks were much smaller.

### 7.3. Pruning results for perfect match

This section presents the pruning results for each of the parallelization and pruning strategies used for the perfect match case (identical sequences) using local alignments. Figs. 11(a), 11(b) and 11(c) shows, respectively, the pruned areas for MASA-CUDAlign (55.2%), MASA-OpenMP/CPU (57.3%) and MASA-OmpSs/CPU (66.2%) when comparing sequence CP000051.1 (1 Mbp) with itself (perfect match). The line in the diagonal represents the optimal alignment and it divides the pruned area in left and the right sides. MASA-OmpSs/CPU presents the biggest pruned area in both sides. Both MASA-CUDAlign and MASA-OpenMP/CPU process the matrix by diagonals, but MASA-CUDAlign has wider blocks, leading to a less inclined wavefront and a different pruned area. Comparing MASA-CUDAlign and MASA-OpenMP/CPU, MASA-CUDAlign has a bigger pruned area in the left side and MASA-OpenMP/CPU in the
right side. Nevertheless, the difference in the right side area is more significant than in the left side, leading to a better pruning performance in the MASA-OpenMP/CPU.

Figure 12 presents the MASA-CUDAlign pruned area for perfect matches with larger sequences (23 Mbp, 32 Mbp and 47 Mbp), where the pruning shape is visibly identical, with 53% of pruned blocks. The shapes shown in Figure 12 (23 Mbp, 32 Mbp and 47 Mbp comparisons) are slightly different from the shape shown in Figure 11(a) (1 Mbp comparison), especially in the right upper border and in the middle row. This happens because the block heights are proportionally larger as we reduce the sequence size, increasing the diagonal wavefront angle.

![Fig. 11. Pruned areas (gray) when comparing sequence CP000051.1 (1 Mbp) with itself (perfect match).](image)

![Fig. 12. Other Pruned areas (gray) with perfect match, using MASA-CUDAlign.](image)

7.4. Execution times for global, semi-global and overlap alignments

In this section we compare the global, semi-global and overlap alignments against the local alignment, considering the MASA-CUDAlign implementation. For a fair comparison, the block pruning was disabled and a new execution was made for the four types of comparisons. In some cases, the optimal overlap alignment can be very short and resides very near the corners of the matrix, limiting the results to positive values.

Table V presents the execution times and GCUPS for the MASA-CUDAlign without block pruning, for the four types of alignment (local, overlap, semi-global and global). Since the global, semi-global and overlap recurrence relation (NW) does not have an if clause to avoid negative values, its Stage 1 performance is slightly better than the local recurrence relation (SW). This difference can be seen in Table V, where the local alignment comparisons are around 3% slower than the other alignment types, in most cases.
Stages 2 to 6 execution times depend on the size of the alignment. The global alignment usually resides in the main diagonal of the matrix, and its length is at least the size of the largest sequence. So, the global alignment is often larger than the other types of alignment, usually leading to a longer traceback time. The 47M local alignment (Figure 14(b)) and the overlap alignment reside in a shifted diagonal, shorter than the global and semi-global alignments which inserted many gaps before the beginning of sequence 1. Nevertheless, in comparisons 5M and 10M, all alignment types produce almost the same results, producing similar traceback time.

Figure 13 presents the shapes of the local, overlap, semi-global and global optimal alignments of the 1M comparison. In this figure, the different types of alignment resides in distinct edges of the matrix. The local alignment (Figure 13(a)) starts in the middle of the matrix and ends in the left edge. The overlap alignment (Figure 13(b)) starts in the left edge and ends in the bottom edge. The semi-global alignment (Figure 13(c)) starts in the left edge and ends in the right edge, representing all the characters of the second sequence. The global alignment (Figure 13(d)) begins and ends in the upper-left and bottom-right corners, larger than the other alignment types although with lower optimal score. These differences in the edges of the alignment produce different scores (Table III), i.e, 88353 (local), 62525 (overlap), -588728 (semi-global) and -1189459 (global).

7.5. Alignment results

7.5.1. Amycolaptosis mediterranei. In this section, we show the results obtained with the comparison of the strains S699 (NC_017186.1) and U32 (NC_014318.1) of *Amycotpisis mediterranei*. *Amycotpisis mediterranei* is a Gram-positive actinomycete which produces an important antibiotic (ricamycin) and it is extensively studied in the literature [Verma et al. 2011] [Zhao et al. 2010].

Figure 14(a) shows the optimal local alignment between these two sequences. In this figure, the green and orange parts represent the cells of the matrix that were calculated whereas the black part represents the pruned cells. As can be seen, it is almost a perfect match. The percentages of matches, mismatches and gaps are, respectively, 99.996%, 0.003% and 0.001%. This is consistent with the result reported in [Verma et al. 2011]. The optimal score is 10,235,188.

7.5.2. Human x Chimpanzee Chromosomes 21. In this section, we discuss results obtained with MASA-CUDAlign for the human x chimpanzee chromosome 21 comparison. The analysis of these two chromosomes is a very active research area and new genes and transcripts are being discovered at a high rate [Scarpato et al. 2014]. Very recently, a new genetic analysis of human chromosome 21 [Letourneau et al. 2014] advanced further the knowledge needed to fully understand Down’s Syndrome.
Table V. Execution times and GCUPS for the MASA-CUDAAlign without block pruning (local, overlap, semi-global and global alignments)

<table>
<thead>
<tr>
<th>Cmp.</th>
<th>SRA</th>
<th>Type</th>
<th>Stages (sec.)</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Local</td>
<td>~ ~ 0.2 ~</td>
<td>~1.3</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Overl.</td>
<td>~ ~ 0.2 ~</td>
<td>0.6</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Semi</td>
<td>~ ~ 0.2 ~</td>
<td>1.6</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Global</td>
<td>~ ~ 0.2 ~</td>
<td>1.5</td>
</tr>
<tr>
<td>10K</td>
<td>1M</td>
<td>Local</td>
<td>0.2 ~ ~ ~</td>
<td>0.4</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Overl.</td>
<td>0.2 ~ ~ ~</td>
<td>1.2</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Semi</td>
<td>0.2 ~ ~ 0.5</td>
<td>2.0</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Global</td>
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<td>2.3</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Local</td>
<td>1.2 ~ ~ ~</td>
<td>2.0</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Overl.</td>
<td>1.2 ~ ~ ~</td>
<td>2.6</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Semi</td>
<td>1.2 0.4 0.5</td>
<td>5.8</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Global</td>
<td>1.2 0.4 0.6</td>
<td>5.4</td>
</tr>
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<td>3M</td>
<td>Local</td>
<td>10.3 ~ ~ ~</td>
<td>10.7</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Overl.</td>
<td>10.0 ~ ~ ~</td>
<td>11.3</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Semi</td>
<td>10.0 1.3 0.9</td>
<td>18.0</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Global</td>
<td>9.9 1.4 1.0</td>
<td>19.1</td>
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<td>5M</td>
<td>Local</td>
<td>37.5 1.2 0.9</td>
<td>45.2</td>
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<tr>
<td></td>
<td></td>
<td>Overl.</td>
<td>36.5 0.8 0.6</td>
<td>43.0</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Semi</td>
<td>36.4 2.2 1.8</td>
<td>51.7</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Global</td>
<td>36.2 2.5 2.0</td>
<td>52.9</td>
</tr>
<tr>
<td>500K</td>
<td>50M</td>
<td>Local</td>
<td>332 ~ ~ 0.3</td>
<td>333</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Overl.</td>
<td>321 ~ ~ ~</td>
<td>322</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Semi</td>
<td>321 11.2 6.0</td>
<td>378</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Global</td>
<td>320 11.1 6.0</td>
<td>377</td>
</tr>
<tr>
<td>1M</td>
<td>250M</td>
<td>Local</td>
<td>871 17.5 9.6</td>
<td>952</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Overl.</td>
<td>846 17.1 9.6</td>
<td>928</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Semi</td>
<td>846 17.2 9.7</td>
<td>928</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Global</td>
<td>844 16.9 9.6</td>
<td>925</td>
</tr>
<tr>
<td>3M</td>
<td>1G</td>
<td>Local</td>
<td>1189 0.2 ~</td>
<td>1191</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Overl.</td>
<td>1152 0.2 ~</td>
<td>1154</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Semi</td>
<td>1152 19.6 9.2</td>
<td>1232</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Global</td>
<td>1150 22.0 10.9</td>
<td>1259</td>
</tr>
<tr>
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<td>3G</td>
<td>Local</td>
<td>3317 113.5 18.7</td>
<td>3557</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Overl.</td>
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<td>3463</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Semi</td>
<td>3211 114.9 18.9</td>
<td>3454</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Global</td>
<td>3208 114.8 18.9</td>
<td>3449</td>
</tr>
<tr>
<td>7M</td>
<td>3G</td>
<td>Local</td>
<td>17793 0.2 ~</td>
<td>17796</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Overl.</td>
<td>17299 0.2 ~</td>
<td>17293</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Semi</td>
<td>17260 392.7 42.8</td>
<td>17908</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Global</td>
<td>17253 407.7 44.7</td>
<td>17962</td>
</tr>
<tr>
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<td>4G</td>
<td>Local</td>
<td>48498 445.1 61.2</td>
<td>49357</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Overl.</td>
<td>47043 440.3 61.2</td>
<td>47898</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Semi</td>
<td>47042 477.7 62.9</td>
<td>47970</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Global</td>
<td>47026 475.5 62.6</td>
<td>47950</td>
</tr>
</tbody>
</table>

Figure 14(b) presents the plot of optimal local alignment between these two whole chromosomes and Figure 15 shows a small part of the text file that contains the alignment. In Figure 14(b), the human chromosome 21 (NC_000021.7) is in the x axis and the chimpanzee chromosome 21 (BA000046.3) is in the y axis. The optimal alignment is shown as a blue line and it starts in position 13,841,681 (NC_000021.7) and 1 (BA000046.3). The percentages of matches, mismatches and gaps are, respectively, 94.380%, 1.537% and 4.082%.

In order to further analyze the optimal alignment, we calculated the percentages of A-T and C-G for both sequences. The results are shown in Figure 16. We can notice...
that the human chromosome 21 has a high concentration of NNs in the beginning of the sequence. This is probably the reason why the optimal local alignment is shifted.

We also determined regions in the alignment that present a high rate of mismatches and gaps. These regions were detected by sliding a window (size = 50000) throughout the alignment and marking the areas with less than 70% matches. In each of these areas, the regions with the lowest score are selected as an unmatched region and the regions that overlap are joined together (Figure 17). The sizes of the regions and the location inside the optimal alignment are shown in Table VI. As it can be seen, we were able to identify 10 regions of interest and detailed biological analysis of these regions seems to be highly recommended.

8. CONCLUSION AND FUTURE WORK

This paper presented Multi-platform Architecture for Sequence Aligners (MASA), a software architecture for the implementation of aligners in different hardware/software platforms. In MASA, the platform-independent code was decoupled from the CUDAlign 2.1 code [Sandes and Melo 2013a] and it was successfully reused in 4 implementations. The MASA-CUDAlign implementation presented results comparable with CUDAlign 2.1, indicating that the MASA architecture did not insert considerable over-
Fig. 16. ATGC and N frequencies of chromosomes 21 (human and chimpanzee).

Fig. 17. Largest unmatched regions in the human x chimpanzee chromosome 21 optimal alignment.

head. The MASA-OmpSs/CPU implementation used a dataflow parallelization strategy able to compute blocks of the DP matrix in a generic order, respecting the data dependencies of the SW/NW algorithm.

In order to allow the execution of Block Pruning (BP) in the dataflow parallelization strategy, a Generic BP algorithm was proposed and implemented. Generic BP was
used in MASA-OmpSs/CPU, MASA-OpenMP/CPU and MASA-OpenMP/Phi, with the difference that the MASA-OmpSs/CPU processed blocks in the dataflow strategy and the others in the diagonal strategy. The experimental results showed that Generic BP was more efficient using the dataflow strategy, since the MASA-OmpSs/CPU increased the GCUPS up to 23.7% compared with MASA-OpenMP/CPU.

As future work, we intend to use vector instructions for the Intel Phi and CPU MASA implementations. We also aim to create a MASA network interconnecting heterogeneous implementations to process the same DP matrix. Considering that the processing nodes may be shared with other processes, we intend to propose a dynamic load balancing mechanism for this MASA network. Finally, we plan to analyze in detail the unmatched regions found in the human x chimpanzee chromosome 21 optimal local alignment and to implement the BP strategy for other recurrence relations.

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