CUDAlign 4.0: Incremental Speculative Traceback for Exact Chromosome-Wide Alignment in GPU Clusters

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Abstract—This paper proposes and evaluates CUDAlign 4.0, a parallel strategy to obtain the optimal alignment of huge DNA sequences in multi-GPU platforms, using the exact Smith-Waterman (SW) algorithm. In the first phase of CUDAlign 4.0, a huge Dynamic Programming (DP) matrix is computed by multiple GPUs, which asynchronously communicate border elements to the right neighbor in order to find the optimal score. After that, the traceback phase of SW is executed. The efficient parallelization of the traceback phase is very challenging because of the high amount of data dependency, which particularly impacts the performance and limits the application scalability. In order to obtain a multi-GPU highly parallel traceback phase, we propose and evaluate a new parallel traceback algorithm called Incremental Speculative Traceback (IST), which pipelines the traceback phase, speculating incrementally over the values calculated so far, producing results in advance. With CUDAlign 4.0, we were able to calculate SW matrices with up to 60 Peta cells, obtaining the optimal local alignments of all Human and Chimpanzee homologous chromosomes, whose sizes range from 26 Millions of Base Pairs (MBP) up to 249 MBP. As far as we know, this is the first time such comparison was made with the SW exact method. We also show that the IST algorithm is able to reduce the traceback time from 2.15 × up to 21.03 × , when compared with the baseline traceback algorithm. The human × chimpanzee chromosome 5 comparison (180 MBP × 183 MBP) attained 10,370.00 GCUPS (Billions of Cells Updated per Second) using 384 GPUs, with a speculation hit ratio of 98.2%.

Index Terms—Bioinformatics, sequence alignment, parallel algorithms, GPU

1 INTRODUCTION

In comparative genomics, biologists compare the sequences that represent organisms in order to infer functional/structural properties. Sequence comparison is, therefore, one of the most basic operations in Bioinformatics [1], usually solved using heuristic methods due to the excessive computation times of the exact methods.

Smith-Waterman (SW) [2] is an exact algorithm to compute pairwise local comparisons. It is based on Dynamic Programming (DP) and has quadratic time and space complexities. The SW algorithm is divided in two phases, where the first phase is responsible to calculate a DP matrix in order to obtain the optimal score and the second phase (traceback) obtains the optimal alignment. SW is usually executed to compare (a) two DNA sequences or (b) a protein sequence (query sequence) to a genomic database. In the first case, a single SW matrix is calculated and all the Processing Elements (PEs) cooperate in this calculation, communicating to exchange border elements (fine-grained computation). For Megabase DNA sequences, a huge DP matrix with several Petabytes is computed. In the second case, multiple small SW matrices are calculated usually without communication between the PEs (coarse-grained computation). With the current genomic databases, often hundreds of thousands SW matrices are calculated in a single query × database comparison.

In the last decades, SW approaches for both cases have been parallelized in the literature, using multiprocessor/multicores [3, 4], CellBEs (Cell Broadband Engines) [5], FPGAs (Field Programmable Gate Arrays) [6], ASICs (Application Specific Integrated Circuits) [7], Intel Xeon Phis [8] and GPUs (Graphics Processing Units) [9, 10, 11, 12]. The SW algorithm is widely used by biologists to compare sequences in many practical applications, such as identification of orthologs [13], and virus integration detection [14]. In this last application, an FPGA-based platform [6] was used to compute millions of SW alignments with small query sequences in short time.

Nowadays, executing SW comparisons with Megabase sequences is still considered unfeasible by most researchers, which currently limits its practical use. We claim that important bioinformatics applications such as whole genome alignment (WGA) [15] could benefit from exact pairwise comparisons of long DNA sequences. WGA applications often construct global genome alignments by using local alignments as building blocks [16] [17]. In [18], the authors state that SW local alignments would be the best choice in this case. However, in order
to compare 1 MBP x 1 MBP sequences, the SW tool took more than 5 days, preventing its use.

In this paper, we focus on GPU solutions since they provide a very good compromise between programmability and performance for SW applications. GPUs are highly parallel architectures that may execute data parallel problems much faster than a general-purpose processor. A number of works have already examined the use of GPUs to accelerate SW computation. Some of them use only one GPU [19, 20, 11, 21, 22], whereas several approaches have been recently proposed to execute SW in multiple GPUs [12, 9, 10].

Some strategies [3, 20, 11, 10] are able to obtain the optimal local alignment of Megabase sequences longer than 1 Million Base Pairs (MBP). These strategies use linear space techniques to obtain the alignment with a reasonable amount of memory. As far as we know, there are no implementations of SW that obtained the optimal alignment between sequences longer than 60 MBP.

To measure the performance of SW strategies, the metric GCUPS (Billions of Cells Updated per Second) is often used. GCUPS are calculated using the formula

\[ \text{GCUPS} = \frac{mn}{t} \]

where \( m \) and \( n \) are the sequence sizes, and \( t \) is the execution time. As far as we know, the best GCUPS (6,020.00) was obtained by the Rivyera FPGA-based platform [6], which calculates SW scores using 128 FPGAs. To our knowledge, the best GCUPS for GPU platforms is 1,782.00, achieved by CUDAlign 3.0 [12]. Therefore, we decided to incorporate our traceback procedure in CUDAlign, since we target GPU platforms.

CUDAlign is a parallel application able to execute the SW algorithm with affine-gap in GPU for huge DNA sequences and it has evolved to many versions with incremental optimizations [19, 20, 11, 10]. CUDAlign 2.1 [11] is able to obtain the optimal alignment of sequences up to 33MBP in a single GPU. The first phase of SW is executed with a new optimization called block pruning and the second phase of SW executes a modified Myers-Miller algorithm [24] in 4 stages. CUDAlign 3.0 [23] was able to obtain the optimal local score comparing sequences up to 228MBP in multiple GPUs [12].

In this paper, we propose and evaluate CUDAlign 4.0, a new version of CUDAlign that is able to retrieve the optimal SW alignment of huge DNA sequences executing both phases of the SW algorithm using multiple GPUs. In CUDAlign 4.0, we faced the challenge of parallelizing the traceback phase of SW, which is essentially serial and consumes more than 50% of the overall execution time in some cases. A straightforward parallelization of this phase would certainly compromise the scalability of the multi-GPU solution. This challenge motivated the development of a new traceback technique called Incremental Speculative Traceback (IST), which takes advantage of otherwise idle times of GPUs to speculate the location of the optimal alignment in order to speedup the traceback phase. As long as the points that actually belong to the optimal alignment are obtained by one GPU, our speculation technique uses them to trigger new speculations in the neighbor GPUs, correcting the former values in an incremental way.

CUDAlign 4.0 was implemented in CUDA, C++ and Pthreads. In order to execute the first phase of the SW algorithm with multiple GPUs, CUDAlign 4.0 extends CUDAlign 3.0 [12] including modifications to allow the execution of the traceback phase of SW. The alignment of the homologous human x chimpanzee chromosome 5 sequences (180 MBP x 183 MBP) was obtained in 53 minutes using 384 GPUs, with a processing rate of 10,370.00 GCUPS, surpassing the best GCUPS in the literature. We also show that our IST technique is very effective, executing the traceback phase up to 21.03x faster than the baseline pipelined traceback. Since IST speculates during the time the GPUs would otherwise be idle, its overhead is negligible.

The rest of this paper is organized as follows. Section 2 presents the biological sequence comparison problem and the SW algorithm. In Section 3 we discuss implementations of SW using multiple Processing Elements. Section 4 briefly reviews the CUDAlign algorithm for a single GPU. Section 5 presents our parallelization of the SW algorithm for multiple GPUs. In Section 6 we present experimental results. Finally, Section 7 concludes the paper and outlines future work.

### 2 DNA Sequence Comparison

A DNA sequence is represented by an ordered list of nucleotide bases. DNA sequences are treated as strings composed of characters of the alphabet \( \Sigma = \{ A, T, G, C \} \). To compare two sequences, we place one sequence above the other, possibly introducing spaces, making clear the correspondence between similar characters [1]. The result of this placement is an alignment.

Given an alignment between sequences \( S_0 \) and \( S_1 \), a score is associated to it as follows. For each pair of characters, we associate (a) a punctuation \( ma \), if both characters are identical (match); or (b) a penalty \( mi \), if the characters are different (mismatch); or (c) a penalty \( g \), if one of the characters is a space (gap). The score is the addition of all these values. Figure 1 presents one possible alignment between two DNA sequences. In this figure, \( ma = +1, mi = -1 \) and \( g = -2 \).

#### 2.1 Smith-Waterman (SW) Algorithm

The algorithm SW [2] is based on Dynamic Programming (DP), obtaining the optimal pairwise local alignment in quadratic time and space. It is divided in 2 phases: calculate the DP matrix and obtain the alignment (traceback).
Phase 1 - This phase receives as input sequences $S_0$ and $S_1$, with sizes $|S_0| = m$ and $|S_1| = n$. The DP matrix is denoted $H_{m+1,n+1}$, where $H_{i,j}$ contains the score between prefixes $S_0[1..i]$ and $S_1[1..j]$. At the beginning, the first row and column are filled with zeroes. The remaining elements of $H$ are obtained from Equation 1.

$$H_{i,j} = \max \begin{cases} H_{i-1,j-1} + \text{(if } S_0[i] = S_1[j] \text{ then } ma \text{ else } mi) \\ H_{i-1,j} + \text{g} \\ H_{i,j-1} + \text{g} \\ 0 \end{cases} \quad (1)$$

In addition, each cell $H_{i,j}$ contains information about the cell that was used to produce the value. The highest value in $H_{i,j}$ is the optimal score.

Phase 2 (traceback) - The second phase of SW obtains the optimal local alignment, using the outputs of the first phase. The computation starts from the cell that has the highest value in $H$, following the path that produced the optimal score until the value zero is reached.

Figure 2 presents a DP matrix with $score = 5$. The arrows indicate the alignment path. In this figure, two DNA sequences with sizes $m = 12$ and $n = 8$ are compared, resulting in a $9 \times 13$ DP matrix. In order to compare Megabase sequences of, for instance, 60 MBP, a matrix of size 60,000,001 $\times$ 60,000,001 (3.6 Peta cells) is calculated.

The original SW algorithm assigns a constant cost $g$ to each gap. However, gaps tend to occur together rather than individually. For this reason, a higher penalty is usually associated to the first gap and a lower penalty is given to the remaining ones (affine-gap model). Gotoh [25] proposed an algorithm based on SW that implements the affine-gap model by calculating 3 values for each cell in the DP matrix: $H$, $E$ and $F$, where values $E$ and $F$ keep track of gaps in each sequence. As in the original SW algorithm, time and space complexities of the Gotoh algorithm are quadratic.

2.2 Myers-Miller Algorithm (MM)

For long sequences, space is a limiting factor for the optimal alignment computation. Myers and Miller (MM) proposed an algorithm [24] that computes optimal global alignments in linear space. It is based on Hirschberg [26], but applied over Gotoh (Section 2.1).

Hirschberg’s algorithm uses a recursive divide and conquer procedure to obtain the longest common subsequence (LCS). The idea of this algorithm is to find the midpoint of the LCS using the information obtained from the forward and the reverse directions, maintaining in memory only one row for each direction (thus in linear space). Given this midpoint, the problem is divided in two smaller subproblems, that are recursively divided in more midpoints.

The MM algorithm works as follows [24]. Let $S_0$ and $S_1$ be the sequences, with sizes $m$ and $n$ respectively, and $i^* = \frac{m+n}{2}$ the middle row of the DP matrices. In the forward direction, $CC(j)$ is the minimum cost of a conversion of $S_0[1..i^*]$ to $S_1[1..j]$ that ends without a gap and $DD(j)$ is the minimum cost of a conversion of $S_0[1..i^*]$ to $S_1[1..j]$ that ends with a gap. In the reverse direction, $RR(n-j)$ is the minimum cost of a conversion of $S_0[i^*..m]$ to $S_1[j..n]$ that begins without a gap and $SS(n-j)$ is the minimum cost of a conversion of $S_0[i^*..m]$ to $S_1[j..n]$ that begins with a gap.

To find the midpoint of the alignment, the algorithm executes a matching procedure between: a) vectors $CC$ and $RR$; b) vectors $DD$ and $SS$. The midpoint is the coordinate $(i^*, j^*)$, where $j^*$ is the position that satisfies the maximum value in Formula 2 [24].

$$max_{j \in [0,n]} \left\{ \max \left\{ CC(j) + RR(n-j) \right\} \right\}$$

After the midpoint is found, the problem is recursively split into smaller subproblems, until trivial problems are found. Figure 3 illustrates the execution of 2 steps of the MM algorithm. In the first step, the whole DP matrix is processed and the midpoint is found. In the second step, the top left and the bottom right rectangles are processed, producing 2 new alignment points. Both points are used to split the rectangles, generating four light gray areas which will be processed in the third step.

2.3 Parallel Smith-Waterman

In SW, most of the time is spent calculating the DP matrices and this is the part which is usually parallelized. In Equation 1, we can notice that cell $H_{i,j}$ depends on three other cells: $H_{i-1,j}$, $H_{i-1,j-1}$ and $H_{i,j-1}$. This kind of dependency is well suited to be parallelized using the wavefront method [27]. Using this method, the DP
matrix is calculated by diagonals, and all cells in each diagonal can be computed in parallel.

Figure 4 illustrates the wavefront method. In step 1, only one cell is calculated in diagonal $d_1$. In step 2, diagonal $d_2$ has 2 cells, that can be calculated in parallel. In the further steps, the number of cells that can be calculated in parallel increases until it reaches the maximum parallelism in diagonals $d_5$ to $d_9$, where 5 cells are calculated in parallel. In diagonals $d_{10}$ to $d_{12}$, the parallelism decreases until only one cell is calculated in diagonal $d_{13}$. The wavefront strategy suffers from reduced parallelism during the beginning of the calculation (filling the wavefront) and the end of the computation (emptying the wavefront).

3 Related Work

There are two main types of SW computations: fine-grained or coarse-grained. Fine-grained proposals are the ones that use more than one PE (Processing Element) to compute the same DP matrix; otherwise, the proposal is classified as coarse-grained. Even though there are several multi-PE coarse-grained approaches for SW in the literature ([28, 9, 22, 6]), we will provide in this section a discussion of fine-grained approaches, which are more closely related to our work.

<table>
<thead>
<tr>
<th>Ref.</th>
<th>Year</th>
<th>Output</th>
<th>Number of PEs</th>
<th>GCUPS</th>
<th>Max. Size</th>
</tr>
</thead>
<tbody>
<tr>
<td>[3]</td>
<td>2004</td>
<td>align.</td>
<td>60 x CPUs</td>
<td>0.25</td>
<td>1,100,000</td>
</tr>
<tr>
<td>[4]</td>
<td>2013</td>
<td>score</td>
<td>6,144 x CPUs</td>
<td>15.50</td>
<td>24,894,699</td>
</tr>
<tr>
<td>[10]</td>
<td>2013</td>
<td>align.</td>
<td>2 x GPUs</td>
<td>65.20</td>
<td>2,799,110</td>
</tr>
<tr>
<td>[8]</td>
<td>2014</td>
<td>score</td>
<td>4 x Xeon Phi</td>
<td>114.40</td>
<td>50,000,000</td>
</tr>
<tr>
<td>[12]</td>
<td>2014</td>
<td>score</td>
<td>64 x GPUs</td>
<td>1,762.00</td>
<td>228,333,387</td>
</tr>
</tbody>
</table>

Table 1 lists fine-grained SW implementations for platforms composed of multiple PEs. The maximum number of PEs used by each paper is presented in column 4. As can be seen, the maximum number of PEs employed is 6,144 (cluster of multicore), 64 (GPU) and 4 (Xeon Phi). GCUPS (column 5) range from 0.25 (2004) to 1,762.00 (2014), showing a spectacular increase in performance in 10 years. These GCUPS values, however, cannot be compared directly because they were obtained in different platforms, with different sequences. Nevertheless, they provide an indication of the potential of each platform. For fine-grained comparisons, the best GCUPS was obtained by CUDAlign 3.0 (1,762.00 GCUPS) [12] with 64 GPUs. If we also consider the coarse-grained approaches, the best GCUPS achieved so far is 6,020.00, obtained by the Rivyera platform [6] with 128 FPGAs. In this case, query sequences of exactly 100 characters are compared with the SW algorithm (linear gap function). The maximum size of the sequences compared in the papers is also presented in Table 1.

In Table 1 (column 3), two approaches were able to retrieve the optimal alignment of long sequences with multiple PEs. The first approach [3] uses Parallel Prefix (PP) to compute SW alignments. PP breaks the dependency among the DP cells, making it possible to compute a whole row/column in parallel, with a communication step at the end of each row/column computation. PP was also implemented for SW executions in the CellBE [5]. In [29], the wavefront method is compared to PP for SW computations. Using their implementations, the authors conclude that PP is faster than wavefront parallelization for sequences up to 4K; for sequences of length 8K or higher, wavefront is faster. Also, in [7], a hardware platform composed of a network-on-chip and ASICs is proposed that implements PP and wavefront parallelization (AD) to execute SW for small sequences. Even though the authors conclude that PP is faster in their platform for 1K x 1K comparisons, the execution times for PP increase when the number of PEs is increased from 64 to 128, limiting the scalability. This behavior does not happen with AD, showing a better potential for scalability. Based on these results, we opted not to use PP to retrieve SW alignments in our design but we think PP can be explored in future work.

The second approach that retrieves the optimal alignment for long sequences with multiple PEs is SW# ([10]), which implements the MM algorithm (Section 2.2), with some optimizations introduced by CUDAlign 2.1 [11]. This solution executes with up to 2 GPUs and, for this reason, we think that SW# may present limited scalability for the sequences longer than 200 MBP. Therefore, we opted to extend CUDAlign 3.0, which executed successfully with up to 64 GPUs, to retrieve SW alignments of huge sequences with the wavefront method.
stored in texture memory. The DP matrix is divided in a grid with \( \frac{n}{\alpha T} \times B \) blocks, where \( B \) is the number of CUDA blocks executed in parallel and \( T \) is the number of threads in each block. A thread is responsible to process \( \alpha \) rows, so a block processes \( \alpha T \) rows. A diagonal of blocks is called external diagonal and a diagonal of threads, inside a block, is called internal diagonal.

To reduce the execution time of Stage 1, three optimizations were proposed [20, 11]: Cells Delegation, Phase Division and Block Pruning. Cells delegation avoids the wavefront (Section 2.3) to be emptied and filled at each external diagonal calculation, providing full parallelism during almost all the time. To do this, the GPU blocks have a parallelogram shape, instead of the typical rectangular one. The right block processes the pending cells of its left block, with cells delegation.

To avoid concurrency hazards, each external diagonal is processed in two phases. The short phase calculates the first \( T \) internal diagonals of the block and the long phase calculates the remaining ones. Each phase corresponds to a kernel, and an optimized kernel is used in the long phase, achieving faster execution time [20].

Block pruning was proposed in CUDAlign 2.1 [11] and its goal is to eliminate the calculation of blocks of cells that surely do not belong to the optimal alignment. This optimization may reduce the execution time in more than 50% if the sequences are very similar.

In order to accelerate the execution of the traceback phase (Section 2.1), CUDAlign stores some rows in a disk space called special rows area (SRA). Figure 5(a) shows the outputs of Stage 1. In this figure, special rows are flushed to disk after each group of 4 blocks is calculated. Special rows are illustrated as a darker horizontal line in the figure, the position of the optimal score is illustrated as a cross and pruned blocks are marked in gray.

### 4.2 Obtaining the Optimal Alignment (Stages 2–6)

Stage 2 executes in GPU a semi-global alignment in the reverse direction starting from the position where the optimal score was found in Stage 1. The goal of Stage 2 is to find all the crosspoints that belong to the optimal alignment and cross the special rows, including the start point of the alignment. The interceptions are found using the Myers-Miller matching procedure (Section 2.2). During the computation of Stage 2, some special columns are saved to disk. Figure 5(b) shows the outputs of Stage 2, including the special columns in vertical lines. The computation starts at the optimal score position (bottom-most cross). Then, in the reverse direction, it finds the crosspoint in the next special row. At the end of Stage 2, the start and end points of the alignment in each partition are known and this information is used to iteratively compute more points that are part of the alignment in the further stages.

Stage 3 (Figure 5(c)) has partitions with defined start and end points, unlike Stage 2, that only has the end point of the alignment and the start point is unknown. The goal of Stage 3 is then to obtain more crosspoints inside each partition.

Stage 4 executes in CPU a modified Myers and Miller algorithm between each successive pair of crosspoints found on Stage 3, using multiple threads. The goal of Stage 4 is to increase the number of crosspoints until the distance between any successive pair of crosspoints is smaller than a given limit. Figure 5(d) shows the additional crosspoints generated during Stage 4.

Stage 5 aligns in CPU each partition formed by the crosspoints found in Stage 4. Then it concatenates all the results, giving as output the full optimal alignment, as can be seen in Figure 5(e). Stage 6 is an optional stage used only for visualization of the alignment.

Algorithm 1 presents the chain of inputs and outputs of the CUDAlign stages. The variable best (line 1) contains the optimal score and its position found in Stage 1. Arrays list2–4 (lines 2 to 4) hold crosspoints found in Stages 2 to 4. The bin (line 5) and txt (line 6) variables refer to the actual alignment files in binary representation and textual representation, respectively.

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**Algorithm 1 Overview of CUDAlign (single GPU)**

```plaintext
1: best ← STAGE1()
2: list2 ← STAGE2(best)
3: list3 ← STAGE3(list2)
4: list4 ← STAGE4(list3)
5: bin ← STAGE5(list4)
6: txt ← STAGE6(bin)
```
5 CUDALIGN DESIGN FOR MULTIPLE GPUs

CUDAlign 3.0 [12] was proposed as a Multi-GPU strategy that executes the first phase of the SW computation (Stage 1) in homogeneous and heterogeneous environments [23]. CUDAlign 4.0, proposed in this paper, extends the ideas of CUDAlign 3.0 in order to use multiple GPUs in both SW phases, obtaining the optimal alignment in a Cluster of GPUs. In this section we will present the design of CUDAlign 4.0. Subsection 5.1 focuses on the Stage 1 (based on CUDAlign 3.0 with some modifications) and Subsection 5.2 explains the new parallel traceback strategies proposed for Stages 2-4.

5.1 Multi-GPU Strategy for Stage 1

CUDAlign 3.0 [12] faced the problem of parallelizing Stage 1 using multi-GPUs. The parallelization was done using a multi-GPU wavefront method, where the GPUs are logically arranged in a linear way, i.e., the first GPU is connected to the second, the second to the third and so on (Figure 6). Each GPU computes a range of columns of the DP matrix and the GPUs transfer the cells of their last column to the next GPU. For environments with homogeneous GPUs, an even distribution of columns is used [12]. For heterogeneous environments, the column distribution works as well using a weighted proportion that considers the computing power of each GPU [23].

5.1.1 Multi-Threaded Design

In CUDAlign 3.0, each GPU is bound to one process and each process has 3 CPU threads: one manager thread ($T_M$) that manages GPU computation, and two communication threads ($T_C$) that handle communication (Figure 7). Iteratively, the $T_M$ invokes GPU kernels for the current external diagonal and transfers cells of the first/last column between GPU and CPU. The short phase kernel (Section 4) was modified to read/write the first/last columns from global memory, allowing the $T_M$ to transfer cells between GPU/CPU. The $T_M$ interacts with the $T_C$ threads to receive its first column from the previous GPU and to send its last column to the next GPU. The transfers are made over chunks of cells with the same height of the block.

Inter-process communication is performed using sockets. If one host contains more than one GPU, CUDAlign 3.0 forks one process per local GPU. Figure 7 illustrates the communication of 4 GPUs, where each host has two GPUs and each GPU has an associated process, with one $T_M$ and two $T_C$.

5.1.2 Input/Output Buffers

Each $T_C$ is associated to a circular buffer, that can be used as an input buffer (if it receives the column from the previous process) or as an output buffer (if it stores the column to be sent to the next process). If the input buffer is full or the output buffer is empty, its respective $T_C$ blocks. On the other hand, if the input buffer is empty or the output buffer is full, the $T_M$ blocks. It is important that the $T_M$ does not block often, allowing us to overlap communication with computation.

Figure 8 illustrates the buffers between 4 GPUs. Buffers $I_2$, $I_3$ and $I_4$ are the input buffers and buffers $O_1$, $O_2$ and $O_3$ are the output buffers. Each output-input pair of buffers ($O_j \rightarrow I_j$) from successive GPUs is continuously transferring data. These buffers are responsible to hide the inter-process communication in such a way that the overhead and small variations in the network may not be perceptible to the wavefront performance. Also, the amount of data waiting in each buffer may be an indication of the balancing quality. For instance, the pair of buffers ($O_1 \rightarrow I_3$) in Figure 8 has more data waiting in the input buffer $I_2$ than in the output buffer $O_1$, giving an indication that GPU1 is producing data faster than GPU2 can process. On the other hand, buffers ($O_3 \rightarrow I_4$) are almost empty, indicating that GPU3 and GPU4 are processing in almost the same speed.

5.1.3 Modifications in Stage 1 for CUDAlign 4.0

In order to execute the traceback in Multi-GPUs, we introduced two modifications in CUDAlign 3.0. First, the CUDA GPU texture memory has a limit of $2^{27}$ linear elements [30] and this forbids its usage for sequences larger than 134MBP. In CUDAlign 3.0, the texture was disabled for such large sequences, slowing down the computation in more than 7% [12]. In order to use the texture memory without the 134MBP size limitation,
we implemented in CUDAlign 4.0 a subpartitioning strategy. In this approach, the partitions that do not fit into the texture memory are divided in a grid of sub-partitions. The subpartitions are computed one after the other, where the first column/row of one subpartition is obtained from the last column/row of its neighbor subpartitions. The second modification was the writing of special rows in the file system. These special rows are used in the further stages to find points where the optimal alignment crosses them (Section 4.2). The special rows written by one GPU are only read by the same GPU, unless for the border rows, which are shared using the TCP sockets and stored inside a filesystem.

5.2 Multi-GPU Strategy for Traceback

In this section, we detail the multi-GPU Incremental Speculative Traceback (IST), which is proposed to obtain the optimal local alignment. IST uses otherwise idle GPU cycles to speculate the locations in which the optimal alignment is likely to be, before the optimal alignment locations are actually known. The IST strategy is built upon a baseline Pipelined Traceback (PT) strategy (Section 5.2.1). The IST strategy is then presented in Section 5.2.2.

5.2.1 Pipeline Traceback (PT) - Baseline Strategy

Our baseline multi-GPU parallelization strategy (Pipelined Traceback - PT) employs the multi-GPU structure of CUDAlign 3.0 [12] combined with a pipelined strategy to compute Stages 2 to 4 (Section 4.2) from different partitions in parallel. For the sake of simplicity, whenever we state that GPUi is processing Stage 4, 5 or 6, we actually mean that the process associated with GPUi is executing it in CPU.

After Stage 1 is completed, the last GPU starts the execution of Stage 2 for the last partition. Whenever the crosspoint from the border column is calculated, its coordinates are sent to the previous GPU, which is responsible for the computation of the neighboring partition. The previous GPU may then start Stage 2. A dependency chain does exist among multiple GPUs. This dependency creates a critical execution path that results in idle time between the end of Stage 1 and the beginning of Stage 2. However, after the dependencies for computation of Stage 2 have been resolved, the execution of Stages 2, 3, and 4 may be computed in pipeline. This strategy is used for all GPUs. The Stages 5 and 6 are less compute intensive and, as a consequence, they are executed by the first GPU.

Figure 9(a) illustrates the Pipelined Traceback execution timeline. The y-axis refers to time and x-axis represents each of the GPUs used, whereas each point shows the stage that a GPU is computing. The white area on the top of the plot (T_a) illustrates the wavefront startup time. The gray areas represent the computational time used for Stages 1 to 6 (T_{1,2,3,4,5,6}). The white area T_b between Stages 1 and 2 is the idle time after a GPU is done with Stage 1 for its partition, waiting to start the computation of Stage 2.

5.2.1.1 Complexity Analysis: In this section, we assume that the manager threads do not block. Considering that p GPUs are used in Stage 2 and that there are more GPUs than special rows, we can roughly estimate that each GPU will compute an area of $\frac{mn}{p}$, resulting in a time complexity of $O(\frac{mn}{p})$ per GPU. Because the execution of the $p$ GPUs has a dependency chain, their execution is serialized and the overall execution time is $O(\frac{mn}{p})$ — the same execution time complexity of Stage 1. Although the pipelined traceback improves performance by executing Stages 2, 3, and 4 in parallel to the computation of other partitions, still there is a critical execution path resulting from dependencies in Stage 2 (Figure 9(a)).

5.2.1.2 Pseudocode: The pseudocode for the Pipelined Traceback strategy is presented in Algorithm 2, which changes the original CUDAlign 2.1 traceback (Algorithm 1) to execute in multi-GPU environments. Variable $n$ represents the $n$th GPU and $recv_i$ and $send_i$ are the message calls used to receive data from the $i$th GPU and send data to the $j$th GPU. To simplify the pseudocode, we ignored some border cases (first and last GPU), where the $recv_i$ and $send_i$ calls are not executed. Using the column distribution discussed in Section 5.1, Stage 1 (line 1) is executed for the columns assigned to the $n$th GPU and the variable $best_n$ receives the local maximum score and its position in this range of columns. Then, the cumulative best score $best_n$ from GPUs 1 to $n$ is passed through the GPUs (lines 2 to 4). In the last GPU, the variable $crosspoint$ will contain the overall best score and its position (line 6), and Stage 2 will start the execution (line 10) from the position with best score (if it resides in the last GPU column range). Further, the top-left crosspoint $list_{0}[0]$ (line 11) found in the border column by Stage 2 is sent to the previous GPU (line 12), which receives it in the $crosspoint$ variable (line 8). While the GPU that just received the crosspoint computes Stage 2, the previous GPU pipelines the execution of Stages 3 and 4 (lines 13 and 14). After each GPU is done with Stage 4, it will receive the crosspoints computed by the right-neighbor GPUs (line 15), concatenate them with its own crosspoints, and send them to the left-neighbor GPU (line 17). This process continues up to the first GPU (or the one containing the beginning of the alignment). This GPU computes the final alignment by executing Stages 5 and 6 (lines 19 and 20).

5.2.2 Incremental Speculative Traceback (IST)

We have noticed in analyses of several tracbacks that the crosspoints found in border columns tend to coincide with the best score in these columns. The Incremental Speculative Traceback (IST) strategy uses this insight to estimate where the optimal alignment will cross intermediate border columns, allowing the execution of Stage 2 for multiple partitions in parallel even before the optimal alignment crosspoints are known. In parallel to this speculative computation, Stage 2 also starts its
results speculated by neighbor GPUs. With this approach, we have modified our solution for using partial looking at local results to find the point for speculation, more difficult to be predicted. Therefore, instead of only to the next GPU. Otherwise, the partition is recalculated.

if that point was guessed correctly and has already been calculated, the GPU will not recompute its partition and will rather use the point that belongs the optimal alignment. Figure 9 illustrates the Incremental Speculative Traceback (IST) and the Pipelined Traceback (PT) for the case of two GPUs.

Algorithm 2 Pipelined Traceback (nth GPU)

```
1: best’n ← STAGE1()
2: RECEVN−1(bestn−1)
3: bestn ← max(bestn−1, best’n)
4: SENDn+1(bestn)
5: if n is last GPU then
6:   crosspoint ← bestn
7: else
8:   RECEVN+1(crosspoint)
9: end if
10: list2 ← STAGE2(crosspoint)
11: left_crosspoint ← list2[0]
12: SENDn−1(left_crosspoint)
13: list4 ← STAGE4(list4)
14: list4 ← STAGE4(list4)
15: RECEVN+1(list4)
16: if crosspoint.score ≠ 0 then
17:   SENDn−1(list4 + list4)
18: else
19:   bin ← STAGE5(list4 + list4)
20:   txt ← STAGE6(bin)
21: end if
```

We have also observed that, in regions with many gaps and mismatches, the actual crosspoint is usually more difficult to be predicted. Therefore, instead of only looking at local results to find the point for speculation, we have modified our solution for using partial results speculated by neighbor GPUs. With this approach, each GPU will forward the crosspoint found during its speculation to the neighbor GPU, which uses it as a new guess. This strategy performs an incremental speculation with increasing probability of success. It is important to highlight that the results of each speculative computation, e.g., special rows, must be saved. As such, we have limited the number of speculation attempts (max_spec_tries) in order to limit memory utilization.

Figure 9(b) illustrates the Incremental Speculative Traceback. Stage 1 execution is the same as in the Pipelined Traceback (Section 5.2.1), but speculation in Stage 2 is carried out in previously idle times between the Stages 1 and 2. The incremental speculation is illustrated as many recomputation lanes going diagonally down-left. The recomputation lane starting from the last GPU refers to the critical execution path in Stage 2, which goes down in cases of recalculations, due to wrongly speculated crosspoints. Note that Stages 3 to 4 follow the execution of Stage 2 for the actual crosspoints in the optimal alignment.

5.2.2.1 Complexity Analysis: In this section, we also assume that the manager threads do not block. The time complexity of Stage 2 with IST using p GPUs may be formalized based on the time spent in the recomputation lanes. First we must note that Stage 2 speculations start as soon as Stage 1 finishes in each GPU, and this happens in different times because of the wavefront propagation. Let T1i be the Stage 1 execution time in GPUi, defined by the equation T1i = T1i−1 + (i−1) · t0 where t0 is the time shift between two successive GPUs caused by the wavefront initialization. Let l be the length of the recomputation lane that starts at GPUi and ends at GPUi−1. Each Stage 2 recalculation spends t2 seconds. So, the recomputation lane that begins in GPUi executes
in the time range starting at $T_1$ and ending at $T_1 + l_1 \cdot t_2$.

Let $T_2$ be the additional time executed by Stage 2 in all $p$ GPUs after the last GPU finished executing Stage 1. $T_2$ depends on both the length $l_i$ and the start position of each recomputation lane, considering that a long lane that starts early may end before a short lane that starts late. Thus, $T_2$ is defined by Equation 3.

$$T_2 = \max_{1 \leq i \leq p}(l_1 \cdot t_2) - T_1$$

If $t_2$ is $O\left(\frac{mn}{p^2}\right)$, the time complexity of Stage 2 is $O(l \cdot \frac{mn}{p^2})$ where $l \ (1 \leq l \leq p)$ depends on the length and position of the recomputation lanes. In the best case, all the speculations are correct. So $l = 1$ and the time complexity is $O\left(\frac{mn}{p}\right)$. In the worst case, all speculations are wrong. Thus $l = p$ and the time complexity is $O\left(\frac{mn}{p}\right)$, which is the time complexity of PT. In order to estimate $l_i$ we observed that the recomputation lanes are usually small, but they can be large in two circumstances. The first case happens in regions with high occurrences of mismatches and gaps. Small variations in the alignment of this region may lead to higher number of matches and greedily provide locally better scores. These variations will force the alignment to follow a further path that will introduce much more penalties, leading to a lower score in the far end of the alignment. The second case occurs when there is a repetition of short DNA patterns, either caused by tandem-repeats or repetitions of N’s characters due to unsequenced regions in the genome assembly (which tends to be reduced in each assembly release). The repeated pattern may increase the score in many different positions, reducing probability of hit.

5.2.2.2 Pseudocode: The pseudocode for the IST is presented in Algorithm 3. Without loss of generality, we assume that the optimal alignment position found by Stage 1 is in the last GPU. For the full IST algorithm, lines 5 to 25 of Algorithm 3 replace lines 5 to 12 of Algorithm 2. In Algorithm 3, the best score of the last column calculated by each GPU, with the exception of the last GPU, is used as the first speculated crosspoint (line 9). Further, the speculation loop (lines 12 to 25) computes Stage 2 with different crosspoint coordinates until a not speculated crosspoint is received: meaning that the actual crosspoint from the optimal alignment has already been found and it is being used. During the speculation phase, if the speculated crosspoint was not previously computed, Stage 2 is executed (line 14) and results are cached (line 15), otherwise results are obtained from the cache (line 17). The top-left crosspoint list2[0] (line 19) computed in Stage 2 is sent to the left-neighbor GPU (line 21) and a new speculated crosspoint is received from the GPU to the right (line 23).

The optimal alignment starts in the last GPU, which uses the optimal score as the first crosspoint (line 6) and defines it as a not speculated crosspoint (line 7). Whenever a not speculated crosspoint is found, the next top-left crosspoint is marked with the same tag (line 20), which is cascaded through all the GPUs and finalizes the execution loop (line 25). The loop ends for each GPU when Stage 2 has been computed for the optimal alignment, and Stages 3 and 4 are pipelined for computation after Stage 2 as discussed in the PT algorithm.

Algorithm 3 was simplified for presentation purpose. In the actual implementation, there is a limitation in the number of times (max_spec_tries) that Stage 2 is executed speculatively. Therefore, when the Cache size reaches max_spec_tries, IST will wait for the actual crosspoint in order to either retrieve the results from cache or re-execute Stage 2 in that GPU.

6 Experimental Results
The experiments were carried out using the XSEDE Keeneland Full Scale (KFS) system, which is a cluster with 264 HP SL25G08 compute nodes. Each node is equipped with two 8-core Intel Sandy Bridge processors, three NVIDIA M2090 GPUs, and 32 GB of RAM. The nodes communicate through a Mellanox FDR InfiniBand interconnect and are attached to a Lustre distributed filesystem. The CUDAlign implementation used standard C++, CUDA 5.5, Pthreads, and Sockets. CUDAlign 4.0 kernels were launched using $B = 64$ CUDA blocks, $T = 128$ threads in each block, and each thread processes $\alpha = 4$ rows of the matrix. We used 8MB local communication buffers, which is sufficient to hold up to 1 million cells. The SW score parameters used were: match: +1; mismatch −3; first gap: −5; extension gap: −2.

We intended to compare the results of CUDAlign 4.0 with a state-of-the-art tool. In order to do that, we downloaded SW# [10], which compares long sequences in GPUs (Section 3) and is freely available. In our tests, we confirmed that SW# executes nucleotide comparisons with up to two GPUs, which would lead to excessively long execution times for the long chromosome comparisons.
6.1 Sequences used in the tests

Our experiments used real DNA sequences retrieved from the National Center for Biotechnology Information (NCBI), available at www.ncbi.nlm.nih.gov. We compared all human (GRCh37) and chimpanzee (panTro4) homologous chromosomes, which produces a test set with 25 pairs of sequences.

The accession numbers and sizes of the chromosomes are presented in Table 2. They vary from 26MBP (Million Base Pairs) to 249MBP, and produce DP matrices from 1.56 to 56.91 Peta Cells. For the sake of validation, optimal local scores, alignment length, coverage, percentage of matches and mismatches, and gaps obtained are also presented in the same table. In this first moment, we compared the chromosomes with the purpose of obtaining the optimal alignments, using a variable number of GPUs. The optimal local alignments are presented in Figure 10, where we can see that many comparisons produced huge local alignments.

6.2 Performance of Pipelined Traceback

The performance and scalability of the baseline PT (Section 5.2.1) was evaluated using sequences chr22 and chr16. The number of nodes was varied from 1 to 128 (3 to 384 GPUs). Table 3 presents the experimental results for the entire execution (Total), which are also broken into the time spent in Stage 1 and the remaining stages (Traceback). As shown, the speedups attained with 128 nodes for chr22 and chr16 were, respectively, 26.9× and 29.7× (21.0% and 23.2% of parallel efficiency).

The breakdown of the total execution shows that the Stage 1 of CUDAlign has a much better scalability. Stage 1 attained speedups of 84.0× and 97.3× with 128 nodes (65.6% and 76.0% of parallel efficiency), resulting in a peak performance of 8.3 and 9.7 TCUPS for chr22 and chr16, respectively. Stage 1 results of chr22 and chr16 are consistent with the ones obtained in CUDAlign 3.0 [12]. The PT traceback phase, on the other hand, was not able to efficiently use the parallel environment and, as a consequence, limited the scalability of the whole application. For instance, the percentage of the execution time spent in the traceback phase increased from about 4% to 71% as the number of nodes used was scaled from 1 to 128. This negative impact of the traceback to the whole application performance is highly reduced when IST is used, as shown in Section 6.3.
6.3 Impact of Incremental Speculative Traceback

The experimental evaluation of the impact of IST to the performance was carried out using 5 pairs of homologous chromosomes: chr22, chr16, chr13, chr8, and chr5. These sequences were selected intending to provide a wide range of variation in the DP matrix size calculated (2.55, 8.13, 13.26, 21.07, 33.04 Peta cells, respectively).

The execution times and TCUPS for all stages, Stage 1, and Stage 2-6 (traceback) are presented in Table 4 for IST and PT. As shown, we have attained up to 10.37 TCUPS for all stages and 11.08 TCUPS if only Stage 1 is considered. Additionally, higher TCPUS are achieved when longer sequences are used. PT and IST execute the same code in Stage 1 and, as such, the execution times are very similar and most of the cases have a difference of up to about 1%, except for some cases highlighted in Section 6.5.

The speculation hits of the IST strategy for comparisons chr22, chr16, chr13, chr8 and chr5 (with 128 nodes) were 69.2%, 88.3%, 79.4%, 99.7% and 98.2% respectively. Considering the high speculation effectiveness, the results show that IST was able to significantly improve the performance of PT for all sequences and number of nodes used. The performance improvement of IST over PT for the traceback phase is also presented in Table 4. IST improved this phase in 2.15–2.81, 4.35–5.03, 3.85–7.66 for comparisons using, respectively, chr22, chr16, and chr13. Additionally, the traceback gains with IST for chr8 and chr5 with 128 nodes were of 18.30 and 21.03, respectively. Regarding the overall execution time, IST was up to 2.93 faster than PT for chr8 with 128 nodes. It must be noted that the speculations do not introduce noticeable overhead since IST executes during the time the GPU would otherwise be idle.

The performance improvement variations are result of the IST ability to correctly speculate, which depends on the characteristics of the alignments. The next section presents a detailed evaluation of the impact of these characteristics to the performance.

6.4 Effect of Alignment Characteristics to IST

To evaluate the impact of the alignment characteristics to the IST performance we have created timelines of the executions for chr8 and chr16 with 128 nodes (Figures 11 and 12). The light gray area on the top of the plots represents Stage 1 execution, while diagonal recomputation lanes pointing to the bottom-left corner refer to the traceback phase (Stages 2 to 5). In the PT timelines, the white area between the Stage 1 and the traceback are idle times as a consequence of dependencies in Stage 2. In the IST timelines, the speculated phase of Stage 2 presents black diagonal lines right after Stage 1. In both cases, a cross (×) marks the total execution time.

The tracebacks with PT (Figures 11 (a) and 12 (a)) are presented as diagonal lines to the bottom-left corner. Here we can clearly notice the large amount of idle time between Stages 1 and 2. Stages 3-6 are pipelined and they are usually executed in a very short time.

In tracebacks with IST (Figures 11 (b) and 12 (b)) we can notice a different behavior, where the speculated
Stage 2 executions are presented as a diagonal line right after the end of Stage 1. From this line, there are other lines that go in the bottom-left direction, representing the recomputation lanes due to IST. Some of these lanes span over many GPUs, as can be seen in the central GPUs of chr16 (Figure 12(b)). In chr8, the speculative prediction was very efficient (Figure 11(b)), producing a traceback path almost straight to the left and reducing the traceback time from 4495s to 246s (18.30x).

In chr16 with PT (Figure 12(a)) we noticed a higher elapsed time in Stages 3-6 between GPUs 110 and 140. This occurred because there is a large gap region around GPU 140, and this produced disproportional partitions that required more time in Stage 4. This additional time was propagated to other GPUs (110-139) due to the stage 4 communication path (line 15 in Algorithm 2). The same behavior happened in chr16 with IST (Figure 12(b)), but the communication spanned more GPUs (1-139) because all these GPUs precomputed the partitions successfully during the Stage 2 speculation phase, and, consequently, all these GPUs needed to wait for the Stage 4 results calculated by GPU 140.

As stated in Section 5.2.2.1, one reason to occur a recomputation lane is the existence of regions with very high incidence of N’s, caused by unsequenced regions in the genomic assembly. Figure 13 illustrates the sizes and positions of the N-regions (in black) in each chromosome. The length and coverage percentage of the largest N-regions in the human chr22, chr16, chr13, chr8 and chr5 are, respectively, 16MBP (31%), 11MBP (12%), 19MBP (16%), 3MBP (2.1%) and 3MBP (1.7%).

We observed that the percentages of N-regions and their locations give a very good prediction of the traceback speedup for the IST method (Table 4). For instance, the chr22 alignment has the largest N-region (in percentage) and it presented the smallest traceback speedups (below 2.81x). On the other hand, chr8 and chr5 alignments have the smallest N-regions, presenting higher speedups (18.30x and 21.03x, respectively).

### 6.5 Buffer Analysis of Stage 1

In Figures 11 and 12, we have also represented the output buffer usage in Stage 1 (Section 5.1.2) using different levels of dark gray columns, which indicate an unbalanced usage of the buffer. When a GPU is slower than its left-neighbor GPU, its buffer starts to be filled until it eventually becomes full, cascading this slow down to previous GPUs. For instance, in chr16 with IST (Figure 12(b)) there are dark gray lines close to the middle GPUs indicating an unbalanced performance in that GPU. This behavior produced a Stage 1 time difference of 6% (55s) when comparing PT and IST executions (Table 4).

IST and PT execute the same code in Stage 1 and these performance differences are not related to the traceback method. This unbalancing may be the result of the following factors: 1) very similar sequence regions, leading to frequent updates in the current best scores, causing a
small overhead due to additional instructions executed in the GPU kernel; 2) concurrent resource usage with other jobs in the cluster, which may slow down the file system performance or increase the network latency. The first factor is meant to be reproducible between runs with the same sequences, but the second factor is usually irreproducible. Additionally, the second factor may produce long-term impact in the wavefront balancing, considering that job concurrency is systemic. With these considerations and analyzing the timelines, we may infer that most of the unbalancing scenarios were caused by irreproducible factors due to intra-cluster concurrency.

7 Conclusion and Future Works

This paper presented CUDAlign 4.0, a multi-GPU algorithm able to compute the optimal alignment of huge DNA sequences. CUDAlign 4.0 was tested in the KFS system, a cluster with 264 homogeneous hosts with 3 GPUs each. We were able to pairwise align the human × chimpanzee homologous chromosomes using the SW algorithm. As far as we know, there is no other implementation in the literature that is able to execute such chromosome comparison using an exact algorithm.

The sizes of the sequences used in the tests varied from 26 MBP up to 249 MBP and the tests computed DP matrices from 1.56 up to 60.20 petacells. In KFS, CUDAlign 4.0 presented results up to 10.37 TCUPS when using 384 GPUs. To our knowledge, this is the highest CUPS rate obtained for SW executions. By incrementally speculating the points that belong to the optimal alignment, the proposed IST strategy was able to improve 2.15x up to 21.03x the traceback execution time, when compared to the baseline strategy (PT). For instance, for the chr5 comparison with 384 GPUs, the traceback time was reduced from 3508s down to 167s.

This paper did not intend to give detailed biological analysis of the chromosomes, but it showed that CUDAlign 4.0 is able to produce the optimal alignment even for the largest human chromosomes. With CUDAlign 4.0, the use of exact methods is now feasible for whole chromosome alignment with huge sequences.

As future work, we intend to integrate our Block Pruning method [11] to the Stage 1 of CUDAlign 4.0, which may significantly reduce the execution time in this phase. We also want to produce sub-optimal alignments to increase the alignment coverage of homologous chromosomes, as well as to produce alignments in the reverse complement strands. Additionally, the static column distribution presented in this paper is suited for dedicated environments, but it may be improved with dynamic load balancing [31] for non-dedicated environments.

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