merAligner: A Fully Parallel Sequence Aligner

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Abstract—Aligning a set of query sequences to a set of target sequences is an important task in bioinformatics. In this work we present merAligner, a highly parallel sequence aligner that implements a seed–and–extend algorithm and employs parallelism in all of its components. MerAligner relies on a high performance distributed hash table (seed index) and uses one-sided communication capabilities of the Unified Parallel C to facilitate a fine-grained parallelism. We leverage communication optimizations at the construction of the distributed hash table and software caching schemes to reduce communication during the aligning phase. Additionally, merAligner preprocesses the target sequences to extract properties enabling exact sequence matching with minimal communication. Finally, we efficiently parallelize the I/O intensive phases and implement an effective load balancing scheme. Results show that merAligner exhibits efficient scaling up to thousands of cores on a Cray XC30 supercomputer using real human and wheat genome data while significantly outperforming existing parallel alignment tools.

I. INTRODUCTION

Recent advances in sequencing technology have made the redundant sampling of genomes extremely cost effective. Such a sampling consists mostly of short reads with low error rates that can generally be aligned to a reference genome in a straightforward way. However, the increasing depth of coverage makes the alignment of the reads to a reference sequence a computationally expensive task, requiring high degrees of parallelism for efficient execution.

The community has therefore developed several approaches for parallelizing the alignment of multiple reads (queries) to a set of reference sequences (targets). Such a class of mapping methods include the seed–and–extend algorithms (e.g. BLAST [1]). In this paradigm, the reference sequences are first indexed by constructing a seed index, and this data structure is then used to locate candidate query–to–target alignments by extracting seeds from the queries and performing seed index lookups. Finally, an extension algorithm is applied to extend a found seed, where a local alignment is returned as the result.

In some applications of this methodology, the reference genome is known a priori, thus allowing an off-line seed index construction that can then be exploited for multiple read data sets. This scenario allows for straightforward parallelization, where the seed index is replicated across a set of computational nodes, which can then independently and concurrently align their subset of the reads. Indeed, there are existing frameworks (e.g. pMap [2]) that automate the process of (1) index replication, (2) distribution of reads across nodes and (3) local alignment computation.

However, this approach can suffer from two major limitations. First, the seed index of very large genomes (e.g. wheat [3], pine [4]) may exceed the memory capacity of a singe node, thereby preventing the use of a simple seed index replication scheme. More significantly, there are important applications where the reference sequence is not known ahead of time, thus obviating the off-line approach, and requiring a high-performance implementation of the seed index construction phase to ensure efficient execution of the end-to-end parallel alignment algorithm. A well-known example of this requirement is present in most de novo genome assembly pipelines.

De novo genome assemblers reconstruct an unknown genome from a collection of short reads. Typically these assemblers first process the input reads and generate contigs, which are genome sequences significantly longer than the input reads. Next, contigs are oriented and gaps are closed during the scaffolding phase. The key first stage of the general scaffolding algorithm is aligning the reads onto the generated contigs [5]. Thus, parallel de novo genome assemblers rely on efficient aligner algorithms, where the seed index construction must be efficiently parallelized and distributed to allow high concurrency solutions for grand-challenge genomes.

The parallel alignment work presented in this paper is mo-
activated by our ongoing effort to parallelize the Meraculous [6] genome assembly pipeline [7]. Our study makes numerous contributions including:

- A highly optimized sequence alignment algorithm that parallelizes all its components from end to end, including I/O and seed index construction.
- Scalable seed index implementation that leverages software caching to extend our high-performance lock-free and communication-optimized distributed hash table.
- An efficient method to preprocess the target sequences that enables exact sequence matching with minimal communication and computation without sacrificing accuracy.
- Techniques to efficiently parallelize the I/O intensive stages and address load imbalance via randomization.
- Close-to-ideal scaling (with 0.7 - 0.78 parallel efficiency) up to 15K cores on NERSC’s Cray XC30 supercomputer, using real data sets from the human and the grand-challenge wheat genomes.
- Comparisons with existing alignment solutions, showing the significant advantage of our end-to-end parallel approach.

Overall, our work shows that efficient utilization of distributed memory architectures enables effective parallelization of sequence alignment in terms of both high scalability and reduced per-node memory requirements. An overview of our efficient end-to-end scalability with performance comparisons to BWA mem [8] and Bowtie2 [9] can be seen in Figure 1; detailed performance analysis is presented in Section VI.

II. THE MERALIGNER ALGORITHM

Algorithm 1 describes the parallel algorithm we employ to align a set of query sequences (reads) to a set of target sequences. We choose Unified Parallel C (UPC) [10] for our implementation to reduce programming complexity of global data structures and facilitate a fine-grained parallelism.

A. Extracting Seeds from Target Sequences

First, each processor \( p_i \) reads a distinct portion of the target sequences (line 4) and stores them in shared memory such that any other processor can access them. Every target sequence of length \( L \) contains \( L - k + 1 \) distinct seeds of length \( k \). The first bases \( 1 \ldots k \) of a target form the first seed, the bases \( 2 \ldots k + 1 \) form the second seed, etc. We extract seeds from the target sequences and associate with every seed the target from which it was extracted (line 5) – we also keep track of the exact offset of the seed in the target. Note that a given seed \( s \) might appear in two or more target sequences.

B. Indexing Target Sequences

Once the seeds are extracted from the target sequences, they are stored in a global hash table, henceforth referred to as the seed index (line 6), where the key is a seed and the value is a pointer to the target sequence from which this seed has been extracted. If a seed is extracted from multiple target sequences, its value in the hash table is a list of pointers to those targets. The seed index is distributed and stored in global shared memory such that any processor can access and lookup any seed. Essentially the seed index data structure provides a mapping from seed to targetSequences (see Figure 2).

C. Locating Query-to-Target Candidate Alignments

Given a seed \( s \) from a query sequence \( q \) and an index seedIndex, we perform a lookup and locate the candidate target sequences that have length \( s \) consecutive bases matching with \( q \) (line 10). Thus, each one of the query-to-target candidate alignments can be located in \( O(1) \) time (see Figure 3).

D. Identifying Alignments via Smith-Waterman

Finally, after locating a candidate target sequence \( t \) that has length \( s \) consecutive bases matching with a query sequence

Algorithm 1 Parallel sequence alignment

1: \textbf{Input:} A set of queries and a set of targets
2: \textbf{Output:} Alignments of queries with targets
3: \textbf{for} all processors \( p_i \) \textbf{in parallel do}
4: \hspace{1em} targetSeqs \leftarrow \text{READTARGETSEQUENCES} (targets)
5: \hspace{1em} seedsInTargets \leftarrow \text{EXTRACTSEEDS} (targetSeqs)
6: \hspace{1em} seedIndex \leftarrow \text{BUILDGLOBALSEEDINDEX} (seedsInTargets)
7: \hspace{1em} myQuerySequences \leftarrow \text{READQUERYSEQUENCES} (queries)
8: \hspace{1em} \textbf{for} each query sequence \( q \) \in myQuerySequences \textbf{do}
9: \hspace{2em} candidateTargets \leftarrow \text{LOOKUP} (seedIndex, \ s)
10: \hspace{2em} \textbf{for} each target \( t \) \in candidateTargets \textbf{do}
11: \hspace{3em} alignmentsSet \leftarrow \text{SMITHWATERMAN} (t, q)
increase for an $S$-fold reduction in the number of messages relative to the unoptimized approach. At the same time, our optimization solves the problem of fine-grained locking. Since each processor iterates over its local-shared stack and stores the received seeds in the appropriate local buckets of the hash table, there is no need for locks, thus allowing the resulting distributed hash table to be lock free.

We manipulate local-shared stacks atomically in order to ensure that processors $p_i$ and $p_k$ trying to store seed entries simultaneously at the local-shared stack of another processor $p_j$ do not overwrite the same locations. When processor $p_j$ stores $S$ entries to the local-shared stack of $p_k$, it needs to locate the position in $p_j$'s stack that these entries should go to. Thus, every local-shared stack is associated with its stack_ptr pointer that indicates the current position in the local-shared stack. These stack_ptr variables are shared and accessible to all processors. Therefore, if processor $p_k$ is about to store $S$ entries to processor $p_j$, it (a) reads the current value of $p_j$'s stack_ptr, called cur_pos, (b) increases the value of $p_j$'s stack_ptr by $S$ and (c) stores the $S$ entries in $p_j$'s local-shared stack into the locations cur_pos ... cur_pos+$S$-1 with an aggregate transfer. Steps (a) and (b) need to be executed atomically to avoid data hazards, for which we use the global atomic fetchadd(). Without this aggregating optimization, one would have to access a single remote bucket at a time and consequently would have to obtain one lock at a time. On the other hand, this optimization reduces the number of atomic operations by a factor of $S$ and highlights the advantage of being lock-free.

### B. Software Caching Schemes

The nature of the alignment problem enables data reuse in both the seeds and target sequences, allowing us to exploit this insight for a more efficient implementation. High throughput sequencing allows genomes to be sampled redundantly at a depth $d$, as visualized in Figure 5. Let $k$ be the length of a seed $s$ and $L$ be the read length. Any seed $s$ of the genome (yellow region) is expected to be found $f = d \cdot (1 - (k-1)/L)$ times in the read data set ($f$ is the mean of the Poisson distribution of key-frequencies [12]), thus resulting in $f$ lookups for that seed within the distributed seed index. Additionally, targets are in general sequences that are significantly longer than the reads. Thus, multiple reads are expected to be aligned with the same target and a given target $t$ is expected to be reused multiple times in the seed extension procedure.

Given this potential for data reuse, we developed a software cache architecture to reduce communication overhead as shown in Figure 6. In UPC, the address space of every node is logically divided into private memory and shared memory.
Fig. 6: Software cache architecture for the distributed seed index and the target sequences. Node $i$ has stored in its seed index cache a seed $s$ (yellow block) from Node $j$. Any lookup for the seed $s$ by processors of Node $i$ will be served by the seed index cache. Similarly, Node $i$ has stored in its target cache a sequence $t$ (red block) from Node $j$. Any processor of Node $i$ that needs to align a query with respect to $t$ will fetch $t$ from the target cache.

The private memory is thread local and can only be accessed by the UPC thread (which maps to a processor in our case) to which it has affinity. On the other hand, a location of the shared memory can be accessed by any UPC thread in the system. It is much faster to access locally stored data than to access shared memory residing on a remote node. Thus, on every node, a portion of the shared memory is dedicated for software caches that can store either remote parts of the distributed seed index (seed index cache) or target sequences owned by remote nodes (target cache). In Figure 6 consider Node $i$ which has stored in its seed index cache a seed $s$ (yellow block) that belongs to the part of the distributed seed index local to Node $j$. Any lookup for the seed $s$ by processors of Node $i$ will be served by the seed index cache resulting in much faster lookup time than accessing the original yellow block on the remote Node $j$. Similarly, consider a target sequence $t$ (red block) which has been stored to the target cache of Node $i$. Processors of Node $i$ that need to align a query in respect to $t$ will fetch $t$ from the target cache and thus avoid the expensive off-node communication.

The expected seed data reuse naturally depends on the seed distribution among processors. As discussed later in Subsection IV-B, for load balancing reasons reads are assigned to processors in a uniformly random fashion. Consider a parallel system with $p$ total processors, with $ppn$ processors per node and a seed $s$ with frequency $f$ in the read data set residing on node $i$. Following the reasoning in the previous paragraph, there are $f - 1$ additional occurrences of that seed $s$ in the read data set or equivalently there are $f - 1$ locations in the reads that include that seed.

We can then ask the question: What is the probability that at least one such read is assigned to node $i$? This problem can be reduced to the well known “bins and balls” experiment. In this case, given $f - 1$ balls (remaining occurrences of the seed $s$) and $m = p/\text{ppn}$ bins (nodes), we toss the balls (reads) uniformly at random — the probability that at least one of these balls falls in bin $i$ (node $i$) is $1 - (1 - 1/m)^{f-1}$. Therefore, with probability $1 - (1 - 1/m)^{f-1}$ our approach will perform at least one seed index lookup of $s$ resulting in a software cache hit, since there are at least two occurrences of that seed in the same node $i$. In order to assess the limits of this optimization, consider the case of a read data set with $d = 100$, $L = 100$, $k = 51$, $f = 50$ and a system with $ppn = 24$ cores per node. Figure 7 shows the probability of any seed being reused at least one time given the previous values of $f$ and $ppm$. Note, however, that this is the behavior in the ideal case of “infinite” cache. In practice, we dedicate a fraction of the nodes’ memory for software caching, and tradeoff memory for increased data reuse. For typical experimental values of $d$ and $ppm$, we expect a significant benefit from our cache optimization strategy, as demonstrated in the experimental results of Section VI.

IV. ALIGNMENT OPTIMIZATIONS

We now discuss our alignment optimizations and theoretical proofs of expected behavior.

A. Optimizing exact read matches

Here we devise a method to preprocess the target sequences to identify properties enabling exact sequence matching with minimal communication. The property we describe is based on the Lemma 1.

**Lemma 1.** Let $k$ be the length of the seeds, $q$ be a query sequence and $t$ be a target sequence where all the seeds extracted from $t$ are uniquely located in $t$ (i.e. this seed cannot be found in any other target sequence). Assume that $s$ is a subsequence in $q$ with $\text{length}(s) \geq k$ and $t$ is a candidate target to be aligned with $q$ that also includes the subsequence $s$. Then, $q$ is uniquely aligned to $t$ with respect to the subsequence $s$, in essence, there is no other target $t'$ that matches with $q$ in the subsequence $s$.

**Proof.** Since $q$ matches with $t$ in $\text{length}(s)$ bases, then all $\text{length}(s) - k + 1$ seeds in $s$ belong to both $q$ and $t$. Since all seeds in $t$ are uniquely located in $t$ we conclude that also these $\text{length}(s) - k + 1$ seeds are uniquely located in $t$ and therefore there are no other targets that include those seeds. Consequently, there are no other targets that include the subsequence $s$.  

Fig. 7: Probability of any seed being reused as a function of cores. We have picked values of $d = 100$, $L = 100$, $k = 51$, $f = 50$ and $ppm = 24$. 

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The figure shows the probability of any seed being reused as a function of the number of cores. The graph demonstrates that as the number of cores increases, the probability of reusing a seed decreases. This is because with more cores, the seed index lookup becomes less likely to be served by the seed index cache due to the increased number of seeds stored locally. The figure also includes a legend that provides the parameters used in the experiment, such as $d = 100$, $L = 100$, $k = 51$, $f = 50$, and $ppm = 24$. These parameters are used to simulate different scenarios and evaluate the performance of the seed index lookup strategy.
Consider a subsequence \( s \equiv q \), and assume that the first candidate target \( t_0 \) is to be aligned with \( q \) and it is known that all the seeds extracted from \( t_0 \) are uniquely located in \( t_0 \) (we detail in the subsequent paragraph how to identify such a property). With a fast check we can determine if \( q \) and \( t_0 \) match in exactly \( \text{length}(s) \) bases. Given this scenario, then via Lemma 1 with \( s \equiv q \) it holds that \( q \) is uniquely aligned to \( t_0 \). Thus it is not necessary to look for more candidate targets and additional seed lookups in the distributed seed index can be avoided. It is thus assured that all possible alignments of \( q \) are found (to the set of the targets) by simply performing a single seed lookup — thereby only requiring minimal communication. Further speedups can also be achieved by recognizing that a seed extension algorithm is not necessary in this case, instead a simple and fast string comparison between \( q \) and the appropriate location of \( t_0 \) can be executed.

We now explain how to identify, efficiently for all target sequences, whether the seeds extracted from \( t_0 \) are uniquely located in \( t_0 \). During the distributed seed index construction described in Subsection III-A, when a processor adds the received seeds in its local buckets of the hash table, it counts the number of occurrences of each seed — a cheap and local operation. We additionally associate a boolean single_copy_seeds flag that is initialized as true for all targets. After inserting the seeds into the seed index, a processor \( p_i \) can visit all the local seeds and if the count of an encountered seed \( s' \) is greater than 1, \( p_i \) sets the flags single_copy_seeds of the targets that \( s' \) was extracted from as false. This indicates that those targets do not have seeds uniquely located in them. At the end of this step, all the remaining targets with single_copy_seeds set to true are guaranteed to have all their seeds uniquely located in them.

To maximize the impact of this optimization, we add an additional strategy. Given the seed length \( k \), the longer a target sequence \( t \) is, the more probable it is that \( t \) contains at least one seed that is not uniquely located in \( t \), thus negating the potential of leveraging the described lookup optimization (even if some reads uniquely match to \( t \)). Now consider the case where a target sequence \( t' \) has all but one seed \( a \) uniquely located in \( t' \). If we fragment \( t' \) in two equal-length subsequences \( t'_1 \) and \( t'_2 \) (that overlap to some degree but have disjoint sets of seeds), then the non-uniquely located seed \( a \) in \( t' \) should be found (by construction) in either \( t'_1 \) or \( t'_2 \). Thus the subsequence not containing the seed \( a \) consists of uniquely located seeds, thereby enabling our described optimization.

The same reasoning can be applied recursively to address the general case where a target \( t' \) contains multiple non-uniquely located seeds. The idea is to fragment the sequence \( t' \) into \( m \) equal-length subsequences \( t'_1, t'_2, ..., t'_m \) that overlap to some degree — however the subsequences have disjoint sets of seeds and the union of their sets of seeds is exactly the set of seeds in the original sequence \( t' \). This approach increases the probability of applying the previous optimization. Note that some additional information must be stored for each of the subsequences \( t'_1, t'_2, ..., t'_m \), to allow quick locating of these subsequences later in the alignment.

### B. Load Balancing

Load balancing the queries might initially seem trivial: given \( n \) queries and \( p \) processors each processor should process \( n/p \) queries. Unfortunately, queries may differ in their processing requirements.

For instance, consider a query \( q' \) that perfectly aligns with a single target sequence. Let \( t_{\text{extractSeed}} \) be the required time to extract a seed from a query, \( t_{\text{lookupSeed}} \) the time to lookup a seed in the seed index, \( t_{\text{fetchTarget}} \) the time to fetch a target sequence, and \( t_{\text{membcmp}} \) the time to perform a membcmp() operation on \( \text{length}(q') \) bytes. Then, the time Algorithm 1 takes (after applying the previous optimization) to process \( q' \) is \( t_{q'} = t_{\text{extractSeed}} + t_{\text{lookupSeed}} + t_{\text{fetchTarget}} + t_{\text{membcmp}} \). On the other hand, consider a query \( q'' \) that can be aligned with \( C \) targets. Assume that \( t_{SW} \) is the time to execute the Smith-Waterman algorithm. Then, processing \( q'' \) takes \( t_{q''} = L \cdot (t_{\text{extractSeed}} + t_{\text{lookupSeed}}) + C \cdot (t_{\text{fetchTarget}} + t_{SW}) \) time, where \( L = \text{length}(q'') - \text{length}(seed) + 1 \). Given \( t_{\text{membcmp}} \leq t_{SW} \), it must hold that \( t_{q''} \geq \min(C, L) \cdot t_q \), thus the processing times of two queries can vary significantly.

Assume that the \( n \) queries can be divided in two categories: “fast” and “slow” (depending on their required processing time). The goal is to evenly distribute the slow queries to the available \( p \) processors. However, because it is unknown a priori if a query is fast or slow, we implement the following load balancing strategy. Before executing Algorithm 1 the order of the queries is randomly permuted in the input file and each processor is assigned a chunk of \( n/p \) consecutive queries from the corresponding file. As proven in Theorem 1, if there are \( h \) “slow” queries, \( p \) available processors and \( p \log p \ll h \leq p \log(p) \), then with high probability the load imbalance (distance of maximum “slow” load from the average “slow” load \( h/p \)) is at most: \( 2 \sqrt{\frac{2h}{p} \log p} \).

**Theorem 1.** Let \( h \) be the number of “slow” queries and \( p \) be the number of available processors and assume that \( p \log p \ll h \leq p \log(p) \). After assigning the \( h \) queries randomly to the \( p \) processors (or equivalently randomly permuting the order of the queries in the input file) then with high probability the load imbalance (distance of maximum “slow” load from the average “slow” load \( h/p \)) is at most: \( 2 \sqrt{\frac{2h}{p} \log p} \).

**Proof.** We formulate the process of randomly permuting the order of the queries in the input file as the uniformly random tossing of \( h \) balls into \( p \) bins. Let \( M \) be the random variable that counts the maximum number of balls (“slow” queries) in any bin. It therefore holds that \( \Pr[M \leq k] = 1 - o(1) \) where \( k = \frac{h}{p} + 2\sqrt{\frac{2h}{p} \log p} \) [13], i.e. with high probability the load imbalance (distance of maximum “slow” load from the average “slow” load \( h/p \)) is at most: \( 2 \sqrt{\frac{2h}{p} \log p} \). \( \square \)

### C. Restricting the Maximum Alignments per Seed

Even after applying the described load balancing scheme, there may be a few seeds that can be aligned with too many

\[ \text{polylog}(p) \text{ is some polynomial in } \log(p) \]
targets, causing a high processing time for the corresponding queries. Additionally, finding those numerous alignments may not be relevant to many genome alignment applications. Thus, a threshold can be set for the maximum number of alignments per seed, after which the candidate alignment queries are stopped. This threshold determines the sensitivity of our aligner and it can be used to trade off accuracy for speed when appropriate.

V. ADDITIONAL OPTIMIZATION

We now describe the additional set of I/O, SIMD, and compression optimizations utilized in our work.

A. Parallel I/O

A standard format to represent DNA short reads is the FASTQ format, a text file that includes one read per line with another line of the same length encoding the quality of each base pair. Unfortunately, there is no scalable way to read a FASTQ file in parallel due to its text-based nature. As previously described [7], our work has therefore been leveraging SeqDB [14], a binary format for storing DNA short reads that is implemented on HDF5 [15]. Although SeqDB was not originally designed with parallelism in mind, its HDF5 format allowed us to utilize parallel input data reading using Parallel HDF5 via the MPI-IO with modest modifications, thus enabling a portable solution. Note that the compression from FASTQ to SeqDB is a one-time lossless conversion, where the resulting file is typically 40-50% smaller than the original FASTQ file. While this compression ratio is less compact than other competing formats, SeqDB is significantly faster during the decompress phase. Overall, the SeqDB format is well suited for parallel sequence processing, and we envision it being used more widely by the community.

B. SIMD Optimized Striped Smith-Waterman

MerAligner spends a significant portion of its runtime using the Smith-Waterman (SW) algorithm for seed extension. Due to the critical role of SW, many efforts have been made to accelerate it by taking the advantages of special hardware SIMD (Single Instruction Multiple Data) instructions. In this work we incorporate such an implementation from the Striped Smith-Waterman (SSW) library [16] which has been shown to be orders of magnitude faster than reference implementations of SW in C.

C. DNA Sequence Compression

Given the \{A,C,G,T\} vocabulary of a DNA sequence, only two-bits per base are required for binary representations. We thus use a high-performance compression library that transforms the DNA sequences from text format into a binary format [7]. This approach reduces the memory footprint by 4×, while also reducing the bandwidth by 4× for communication events that involve seeds or DNA sequence transfers.

VI. EXPERIMENTAL RESULTS

We now discuss our experimental testbed and runtime results across a variety of input sets, concurrencies, and optimization schemes, as well as comparison with competing approaches.

A. Experimental Testbed

High-concurrency experiments are conducted on Edison, a Cray XC30 supercomputer at NERSC[17]. Edison has a peak performance of 2.57 petaflops/sec, with 5576 compute nodes, each equipped with 64 GB RAM and two 12-core 2.4 GHz Intel Ivy Bridge processors, and interconnected with the Cray Aries network using a Dragonfly topology.

Our experiments utilize real data sets for human and wheat genomes. The human data set contains 2.5 billion reads (252 Gbp of sequence) for a member of the CEU HapMap population (identifier NA12878) sequenced by the Broad Institute, and our goal is to align those reads onto a set of target sequences (contigs) that are generated in the Meraculous de novo genome assembly pipeline. The reads are 101 bp in length from a paired-end insert library with mean insert size 238 bp. The wheat data set, contains 2.3 billion reads (477 Gbp of sequence) for the homozygous bread wheat line 'Synthetic W7984' sequenced by the JGI and again we want to align those reads onto a set of target sequences (contigs) that are generated in the Meraculous pipeline. The reads are 100-250 bp in length from 5 paired-end libraries with insert size 240-740 bp. For all experiments the seed length is set to 51, as used in the actual scaffolding step of the Meraculous pipeline.

B. Strong Scaling of End-to-End merAligner

Figure 1 (page 1) shows the merAligner end-to-end strong scaling performance with all optimizations applied. This summarizes the main result of this study, and demonstrates the efficient utilization of distributed memory architectures for enabling scalable high performance sequence alignment. More specifically when scaling from 480 to 15,360 cores the total execution time drops from 4,147 seconds to 185 seconds (a 22× speedup), which translates to 0.7 parallel efficiency at the extreme scale for the human dataset (red curves). At the scale of 15,360 cores our approach performs alignment at 15,499,718 reads/sec. For the larger wheat data set (blue curves), scaling from 960 to 15,360 cores achieves 0.78 parallel efficiency. Note the super-linear speedup in the range of 960 — 7,680 cores, which we speculate is due to reduced congestion on the NIC since the communication is spread to even more nodes while we scale. This will be the subject of future investigation.

C. Anatomy of the Optimizations’ Benefits

We now examine the individual effect our optimization schemes, by selectively turning them off and measuring the resulting performance impact.
1) Distributed Seed Index Construction: Figure 8 illustrates the scaling of the distributed seed index construction before and after applying the “aggregating stores” optimization, given an $S = 1000$. Observe that the reduction in the communication via our optimization dramatically decreases the construction time. At 480 cores the time spent decreases from 1,229 seconds to 262 seconds (4.7× improvement), and similarly at 7,680 cores we achieve an improvement of 4.8×. For the optimized construction phase, increasing concurrency from 480 cores to 7,680 (16× core increase) results in a near-linear speedup speedup of 12.7×. These results show that our algorithm efficiently parallelizes the seed index construction in a distributed memory and enables end-to-end scaling of merAligner. These scalable seed index construction results are in contrast to serial approaches of competing alignment codes as detailed in Subsection VI-D. Also, the algorithm achieves almost perfect load balance in terms of the number of distinct seeds assigned to each processor, thanks to our use of the djb2 hash function to implement the seed to processor map.

2) Software Caching: In Figure 9 we depict the benefits of software caching on the communication time during the alignment phase (in all experiments 16 GB/node and 6 GB/node are allocated for the seed index and the target cache respectively). The red bars indicate the communication time for the seed lookups and the blue bars represent target sequence fetching overhead. Observe that the target cache is extremely efficient at all concurrencies and it essentially obviates all the communication involved with target sequences. Results also show that the seed index cache is effective at small concurrencies, where lookup time is decreased from 4,839 seconds to 3,130 seconds (~35% reduction) at 480 cores, whereas larger concurrencies see small benefits — validating our analysis in Subsection III-B. Overall, the caching scheme decreases communication overhead by 2.3×, 1.7× and 1.8× at concurrencies of 480, 1,920 and 7,680 cores respectively.

3) Exact Read Matching Optimization: Figure 10 shows the significant performance benefits of exact read matching, validating the theoretical analysis of Subsection IV-A. Here the optimization results in runtime improvement of the alignment step by factors of 2.8× and 3.1× for 480 cores and 7,680 cores respectively. Note that these gains come from both decreased communication (since in exact matching just one seed lookup is sufficient) and reduction of computation time (by avoiding Smith-Waterman execution). For example, at 480 cores our approach improves computation by 2.48× and communication by 2.82×. Finally we emphasize that ~59% of the aligned reads took advantage of this optimization, thus enabling these impressive performance gains. For the optimized aligning phase, increasing concurrency from 480 cores to 7,680 (16× core increase) results in a near-linear speedup of 15.9×.

4) Load Balancing: In order to assess the effectiveness of the load balancing scheme, we conducted experiments with and without permuting the input read files and measured the maximum, minimum and average computation time as well as alignment times (computation plus communication). Results for 480 cores for the human data set are shown in Table I. Although our load balancing scheme effectively helps reduce the maximum computation time by almost 2.5×, the total alignment time is only improved by ~5%. A closer investigation of the original data set reveals that the reads mapping to the same genome region are grouped together. Since some groups of reads did not map to any target, they...
do not require Smith Waterman execution, and thereby cause an imbalanced computing load. However, this locality made our seed index cache extremely effective and substantially decreased the communication time. Therefore, our load balancing scheme alleviates the computational load imbalance, while making the seed index cache less effective as seen in Table I. Nonetheless, our approach improves the overall execution time. We note that the read grouping in the original data set is not the common case, and thus expect our load balancer to be even more effective in the general case.

D. Comparison with Existing Parallel Aligners

To assess our optimized merAligner in the context of existing solutions, we compare human data performance with BWA-mem [8] and Bowtie2 [9] using the pMap [2] framework. Note that pMap was modified to use the latest versions of the alignment software. Our experiments are configured using 4 instances of 6 threads per Edison node, since it is not possible to run one instance per core due to memory requirements of BWA-mem and Bowtie2 (each node contains 64GB of memory, which is insufficient to hold 24 instances of the seed index). BWA-mem is run with minimum seed length equal to 51 (like merAligner) since alignments with smaller seeds are not applicable to the scaffolding computation. For Bowtie2, we set the minimum seed length to the maximum possible value (31) and we execute the experiment with the --very-fast option in order to achieve the best mapping runtime. It is important to highlight that the seed index construction for BWA-mem and Bowtie2 is performed serially. For both cases, pMap partitions the reads to the available instances, then the seed index is loaded into each instance’s memory and finally the corresponding instance is called on the set of reads assigned to it.

Table II presents comparative end-to-end performance results at 7,680 cores, and notes which computing phases are performed in serial (S) or parallel (P). As expected, the serial seed index construction is a major bottleneck for the competing codes, compared with our parallel merAligner approach. Also, pMap spends a significant amount of time in read partitioning by having a single process sending the appropriate portion of the input read files to the corresponding node (4,305 and 3,982 seconds for BWA-mem and Bowtie2 respectively). On the contrary, merAligner does not suffer from this overhead since all processors read in parallel the appropriate portions of the input read files. To make though a fair comparison, we exclude the timing of the read partitioning for the cases of BWA-mem and Bowtie2. In total, merAligner is \(20.4 \times\) and \(39.4 \times\) faster than the parallel execution of BWA-mem and Bowtie2, respectively.

A complete analysis of the accuracy of the method is outside of the scope of this paper. The algorithm is guaranteed to identify all alignments that share at least one identically matching stretch of at least length(seed) consecutive bases between query and target sequences. Whether such alignments are sufficient is largely an application-dependent question. For the purposes of the Meraculous de novo assembly toolkit, these alignments are precisely those required. Here, we simply report all alignments detected (i.e. without any percent-identity thresholding, using a commonly employed scoring matrix) and find that merAligner successfully aligned 86.3% of the reads, while BWA-mem and Bowtie2 aligned 83.8% and 82.6% of the reads respectively. The accuracy of the detailed alignments is a function of the Smith-Waterman code, and we refer the interested reader to that publication [16].

To further assess merAligner performance, we conduct experiments on the smaller 4.64 Mbp E. coli K-12 MG1655 dataset, which allows single node scalability experiments using both BWA-mem and Bowtie2 in parallel mode with threads. The execution time of all three approaches (using a seed length of 19) is shown in Figure 11. Observe that merAligner performance continues to scale using all 24 available cores, while the runtimes of BWA-mem and Bowtie2 stop improving at 18 cores. Overall, merAligner is significantly faster, exceeding BWA-mem and Bowtie2 performance on 24 cores by \(6.33 \times\) and \(7.2 \times\) respectively. Subject to the alignment correctness discussion above, we find that merAligner successfully aligned 97.4% of the reads, while BWA-mem and Bowtie2 aligned 96.3% and 95.8% of the reads respectively.

Table I: Effect of load balancing scheme on the human data set, showing the reduction of the maximum compute time

<table>
<thead>
<tr>
<th>Load Balancing</th>
<th>Computation time</th>
<th>Total Alignment time</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Min (sec)</td>
<td>Avg (sec)</td>
</tr>
<tr>
<td>Yes</td>
<td>678</td>
<td>740</td>
</tr>
<tr>
<td>No</td>
<td>515</td>
<td>690</td>
</tr>
</tbody>
</table>

Table II: End-to-end performance comparison between parallel executions of merAligner, BWA-mem and Bowtie2 using 7,680 cores on the human data set (with all times in seconds) — highlighting serial (S) or parallel (P) implementation of the phases.

<table>
<thead>
<tr>
<th>Aligner</th>
<th>Seed Index Construction</th>
<th>Mapping Time</th>
<th>Total Alignment Time</th>
<th>Speedup</th>
</tr>
</thead>
<tbody>
<tr>
<td>merAligner</td>
<td>21 (P)</td>
<td>263 (P)</td>
<td>284 sec</td>
<td>1×</td>
</tr>
<tr>
<td>BWA-mem</td>
<td>5,384 (S)</td>
<td>421 (P)</td>
<td>5,805 sec</td>
<td>20.4×</td>
</tr>
<tr>
<td>Bowtie2</td>
<td>10,916 (S)</td>
<td>283 (P)</td>
<td>11,119 sec</td>
<td>39.4×</td>
</tr>
</tbody>
</table>
A thorough survey of sequence aligners is beyond the scope of our work and we refer the reader to [18], [19], [20], [21], [22], [23]. We primarily focus on parallel sequence mapping tools and relevant methods in this section. CUSHAW2 [24], BWA [25], BWA-mem [8], Bowtie2 [9], SNAP [26], SOAP [27] and GSNAP [28] are mapping tools that employ shared memory parallelism during the aligning phase. However, these approaches are more restrictive as they are limited by the concurrency and memory capacity of the shared-memory node. CUSHAW2-GPU [29] and SOAP3-dp [30] are short read aligners that leverage GPU power on a single compute node. pMap [2] is a MPI-based tool used to parallelize existing short sequence mapping tools (like the ones mentioned above) by partitioning the reads and distributing the work among the processors. However, pMap does not leverage any parallelism during the index table construction and therefore a serialization bottleneck is introduced in the mapping pipeline. PBWA [31] employs MPI in order to execute BWA on distributed memory machines, however the index table construction and its replication are serial processes. Also, the sequence distribution is done by a single master process. Therefore, pBWA suffers from the same limitations as pMap. Menon et al. [32] parallelize the genome indexing with MapReduce, however the scaling they obtain is poor.

Bozdag et al. [33] evaluate different methods of distributed memory parallelization of a mapping pipeline. These methods fall basically in three categories: (i) partitioning the reads only, (ii) partitioning the genome (and consequently the index table) and (iii) hybrid method of (i) and (ii) that partitions both reads and the genome (and the index table). One conclusion of this study is that method (i) suffers from the serialized index table construction, method (ii) does not scale in the mapping phase, and regarding method (iii), even though it exhibits improved scalability, its scaling is not close to linear. The main reason is that in the hybrid method, the index table creation is parallelized among subgroups of processes and the reads are also partitioned among subgroups of processes. Therefore, the hybrid method does not exploit the highest possible level of concurrency. Our work does fully parallelize the index table creation and partitions the reads using all available processors.

pFANGS [34] also tries to parallelize both the index table construction and the alignment phase. It distributes the index table among the processors but the processors can not look up the distributed index in arbitrary locations. Therefore, the index lookup tasks are localized first, then an all-to-all personalized communication step is performed, the local lookups take place, and finally the lookup results are redistributed such that they are placed with the relevant queries (this redistribution is done with all-to-all personalized communication). The authors identify that the communication becomes a bottleneck because of the all-to-all communication and therefore they divide the processes in disjoint subgroups where each subgroup works independently by creating its own copy of the index table. However, in this approach the scaling of the index table construction is limited by the size of each subgroup. Orion [35] is an improvement over mpiBLAST [36] and scales the sequence matching with fine-grained parallelization. However, Orion uses mpiBLAST’s mpiformatdb tool to format and to shard the database and this process is serial.

Our previous study [7] investigated the building of distributed hash tables for the contig construction phase of Meraculous [6]. This work extends the algorithm for our distributed seed index optimizations, which now allows multiple seeds to map onto a given target. Also, in this work we enhance the distributed hash table with software caching support. Finally, Kassens et al [37] employed a PGAS language called UPC++ to parallelize Genome-Wide Association Studies and showed that this programming model is suitable for data-intensive bioinformatics applications.

VIII. CONCLUSIONS

This work presents a highly scalable sequence alignment algorithm that effectively parallelizes all computational phases, including seed index construction. Achieving our solution required numerous innovations including software-caching, accelerated exact sequence matching, I/O optimization, and load balancing via randomization. Overall we achieved near perfect scaling on up to 15K cores using real human and wheat data sets, while significantly exceeding the end-to-end performance of existing approaches by factors of 20-39×.

The approach described here was initially developed as part of a UPC adaptation of the Meraculous genome assembler [3], since aligning reads to a reference set of contigs was rate limiting after other stages of the assembler had been parallelized. Other shotgun assemblers have comparable steps, and adaptations of our method (perhaps using different detailed alignment scoring after identifying and extending seed matches) would also likely be useful as modules in other assembly settings. The Striped Smith-Waterman local alignment engine could easily be replaced with any other local alignment software tool; more broadly any seed-and-extend algorithm could be implemented with minor changes to the underlying protocols, including protein-DNA and protein-protein alignments. Importantly, for the case of alignment of read sets from multiple individuals against a reference genome, the cost of building an index can be amortized across the individuals. In contrast, for de novo assembly of a new genome a new index needs to be built for each new assembly. Therefore, the parallelization of index construction is a key element of our approach relative to existing short-read aligners.

While we have focussed on the DNA-alignment problem, and in particular its application in the context of whole-genome shotgun assembly, the work presented here is of sufficient generality that we envision merAligner providing a framework for a generic, distributed hash platform for a variety of applications. In the DNA sequence domain, large genomes (e.g., wheat or pine) may require indices that, unlike human, may not fit in the memory on a single node; our parallel approach avoids this limitation by constructing the index in parallel across distributed memories. Because merAligner
is strongly scalable, it supports the exponential growth and complexity of genome references, in addition to the rapid throughput needed to align massive numbers of genomes.

Other large sequence collections, like the GenBank [38] collection of all known genomic data, are not feasible to index using the methods of existing tools such as BWA and Bowtie; in principle, our approach is scalable to allow short reads from any source to be rapidly aligned against the complete GenBank collection. Extending our approach to other alphabets, one can also use the same methods to align protein sequences (strings of 20 characters, each corresponding to one of the 20 amino acids) against protein datasets. Finally, more general text based queries outside of the biological sequence domain may also be facilitated by our new parallel approach.

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