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A new algorithm for “the LCS problem” with application in compressing genome resequencing data

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Abstract

Background: The longest common subsequence (LCS) problem is a classical problem in computer science, and forms the basis of the current best-performing reference-based compression schemes for genome resequencing data.

Methods: First, we present a new algorithm for the LCS problem. Using the generalized suffix tree, we identify the common substrings shared between the two input sequences. Using the maximal common substrings, we construct a directed acyclic graph (DAG), based on which we determine the LCS as the longest path in the DAG. Then, we introduce an LCS-motivated reference-based compression scheme using the components of the LCS, rather than the LCS itself.

Results: Our basic scheme compressed the Homo sapiens genome (with an original size of 3,080,436,051 bytes) to 15,460,478 bytes. An improvement on the basic method further reduced this to 8,556,708 bytes, or an overall compression ratio of 360. This can be compared to the previous state-of-the-art compression ratios of 157 (Wang and Zhang, 2011) and 171 (Pinho, Pratas, and Garcia, 2011).

Conclusion: We propose a new algorithm to address the longest common subsequence problem. Motivated by our LCS algorithm, we introduce a new reference-based compression scheme for genome resequencing data. Comparative results against state-of-the-art reference-based compression algorithms demonstrate the performance of the proposed method.

Keywords: Longest common subsequence, LCS, Longest previous factor, LPF, Compression, Biology, Genome resequencing

Background

Measuring similarity between sequences, be it DNA, RNA, or protein sequences, is at the core of various problems in molecular biology. An important approach to this problem is computing the longest common subsequence (LCS) between two strings S_1 and S_2, i.e. the longest ordered list of symbols common between S_1 and S_2. For example, when S_1 = abba and S_2 = abab, we have the following LCSs: abba and abab. The LCS has been used to study various areas (see [2, 3]), such as text analysis, pattern recognition, file comparison, efficient tree matching [4], etc. Biological applications of the LCS and similarity measurement are varied, from sequence alignment [5] in comparative genomics [6], to phylogenetic construction and analysis, to rapid search in huge biological sequences [7], to compression and efficient storage of the rapidly expanding genomic data sets [8, 9], to re-sequencing a set of strings given a target string [10], an important step in efficient genome assembly.

The basic approach to compute the LCS, between the n-length S_1 and m-length S_2, is via dynamic programming.
Using \( \text{LCS} \) to denote the dynamic programming (DP) table, the basic formulation is as follows, given \( 0 \leq i \leq n \) and \( 0 \leq j \leq m \):

\[
\text{LCS}(i, j) = \begin{cases} 
0, & \text{if } i = 0 \lor j = 0 \\
1 + \text{LCS}(i-1, j-1), & \text{if } S_1[i] = S_2[j] \\
\max\{ \text{LCS}(i, j-1), \text{LCS}(i-1, j) \}, & \text{if } S_1[i] \neq S_2[j] 
\end{cases}
\]

The above computes the length of the \( \text{LCS} \) in the last position of the table \( \text{LCS}(n, m) \). As with the edit distance computation, the actual string forming the \( \text{LCS} \) can be obtained by using a trace back on the DP table. This requires \( O(nm) \) time and \( O(nm) \) space. The \( \text{LCS} \) matrix has some interesting properties: the entries in any row or in any column are monotonically increasing, and between any two consecutive entries in any row or column, the difference is either 0 or 1. An example \( \text{LCS} \) matrix and trace are shown in Fig. 1.

Alternatively, we can formulate the problem as a two-dimensional grid, where the goal is to find the minimal cost (or maximal cost, depending on the formulation) path, from the start position on the grid (typically, \((0,0)\)), to the end position \((n,m)\). Myers et al. [11] and Ukkonen [12] used this idea to propose a minimum cost path determination problem on the grid, where the path takes a diagonal line from \((i-1, j-1)\) to \((i,j)\) if \(S_1[i] = S_2[j]\) with cost 0, and takes a horizontal or vertical line with a cost of 1, corresponding respectively to insert or delete operations. Hunt and Szymanski [13] earlier used an essentially similar approach to solve the \( \text{LCS} \) problem in \((r+n) \log n \) time, with \( n \ll m \), where \( r \) is the number of pairwise symbol matches \((S_1[i] = S_2[j])\). When two non-similar files are compared, we will have \( r \ll nm \), or \( r \) in \( O(n) \), leading to a practical \( O(n \log n) \) time algorithm. However, for very similar files, we have \( r \approx nm \), or an \( O(nm \log n) \) algorithm. This worst-case occurs, for instance, when \( S_1 = a^n \) and \( S_2 = a^m \).

Hirschberg [14] proposed space-efficient approaches to compute the \( \text{LCS} \) using DP in \( O(nm) \) time and \( O(n+m) \) space, rather than \( O(nm) \). More recently, Yang et al. [15] used the observation on monotonically increasing values in the \( \text{LCS} \) table to identify the “corner points”, where the values on the diagonals change from one row to the next. The corners define a more sparse 2D grid, based on which they determine the \( \text{LCS} \).

A generalization of the \( \text{LCS} \) problem is to find the \( \text{LCS} \) for a set of two or more sequences. This is the multiple longest common subsequence problem, which is known to be NP-hard for an arbitrary number of sequences [16]. Another interesting view of the \( \text{LCS} \) problem is in terms of the longest increasing subsequence (\( \text{LIS} \)) problem, suggested earlier in [17–19], and described in detail in [2]. The \( \text{LIS} \) approach also solves the \( \text{LCS} \) problem in \( O(n \log n) \) time (where \( m \leq n \)). In most practical scenarios, \( r < nm \).

The \( \text{LCS} \) has been used in some recent algorithms to compress genome resequencing data [20, 21]. Compression of biological sequences is an important and difficult problem, which has been studied for decades by various authors [22–24]. See [9, 25, 26] for recent surveys. Most of the earlier studies focused on lossless compression because it was believed that biological sequences should not admit any data loss, since that would impact later use of the compressed data. The earlier methods also generally exploited self-contained redundancies, without using a reference sequence. The advent of high-throughput next generation sequencing, with massive datasets that are easily generated for one experiment, have challenged both compression paradigms.

Lossy compression of high-throughput sequences admitting limited errors have been proposed in [27, 28] for significant compression. With the compilation of several reference genomes for different species, more recent methods have considered lossless compression of re-sequencing data by exploiting the significant redundancies between the genomes of related species. This observation is the basis of various recently proposed methods for reference-based lossless compression [20, 21], whereby some available standard reference genome is used as the dictionary. Compression ratios in the order of 80 to 18,000 without loss have been reported [20, 21]. The \( \text{LCS} \) is the hallmark of these reference-based approaches. In this work, we first introduce a new algorithm for the \( \text{LCS} \) problem, using suffix trees and shortest-path graph algorithms. Motivated by our \( \text{LCS} \) algorithm, we introduce an improved reference-based compression scheme for resequencing data using the longest previous factor (\( \text{LPF} \)) data structure [29–31].

**Methods**

**Preliminaries**

A string \( T \) is a sequence of symbols from some alphabet \( \Sigma \). We append a terminal symbol \( $ \notin \Sigma \) to strings for...
completeness. A string or data structure $D$ has length $|D|$, and its $i$th element is indexed by $D[i]$, where $1 \leq i \leq |D|$. A prefix of a string $T$ is $T[1 \ldots i]$ and a suffix is $T[i \ldots |T|]$, where $1 \leq i \leq |T|$. The suffix tree ($ST$) on the $n$-length $T$ is a compact trie (with $O(n)$ nodes constructed in $O(n)$ time [3]) that represents all of the suffixes of $T$. Suffixes with common prefixes share nodes in the tree until the suffixes differentiate and ultimately, each suffix $T[i \ldots n]$ will have its own leaf node to denote $i$. A generalized suffix tree ($GST$) is an $ST$ for a set of strings. A substring of $T$ is $T[i \ldots j]$, where $1 \leq i \leq j \leq n$. The longest common subsequence is defined below in terms of length-1 common substrings.

**Definition 1. Longest common subsequence (LCS):**

For the $n$-length $S_1$ and $m$-length $S_2$, the LCS between $S_1$ and $S_2$ is the length of the longest sequence of pairs $M = \{m_1, \ldots, m_M\}$, where $m_i = (u, v)$ such that $S_1[m_i.u] = S_2[m_i.v]$ for $1 \leq h \leq M$ and $m_i.u < m_{i+1}.u$ and $m_i.v < m_{i+1}.v$ for $1 \leq i < M$.

**LCS algorithm**

Below, we compute the LCS between $S_1$ and $S_2$ in the following way. (i) We use the GST to compute the common substrings (CSSs) shared between $S_1$ and $S_2$. (ii) We use the CSSs to construct a directed acyclic graph (DAG) of maximal CSSs. (iii) We compute $LCS$ by finding the longest path in the DAG. Steps (i) and (iii) are standard tasks. For step (ii), we develop new algorithms and data structures.

**Computing the CSSs**

We now briefly describe finding the common substrings (CSSs) between $S_1$ and $S_2$. In our $LCS$ algorithm, for simplicity of discussion, we will only use CSSs of length-1.

Let $A = \emptyset$. Compute the GST on $S_1$ and $S_2$, for terminals ($S_1$, $S_2$). Consider a preorder traversal of the GST. When at depth-1 for a node $N$, let $S = \emptyset$. During the preorder traversal from $N$, we collect in $S$ all of the suffix index leaves descending from $N$, which represent the suffixes that share the same first symbol. Let $S_1 = S_2 = \emptyset$. For $s \in S$, if $s \leq |S_1|$, then store $s$ in $S_1$. Otherwise, store $s$ in $S_2$. We represent all of our length-1 matches in the following structure: MATCH $(id, p_1, p_2)$. The $id$ is a unique number for the MATCH, and $p_1$ and $p_2$ are respectively the positions in $S_1$ and $S_2$ where the CSS exists. Let $id = 2$. Now, for each $s_1 \in S_1$, we create a new MATCH $m = (id++, s_1, s_2)$ for each $s_2 \in S_2$. Store each $m$ in $A$.

The running time is clearly the maximum of the GST construction and the number of length-1 CSSs.

**Lemma 2.** Say $n = |S_1|$ and $m = |S_2|$, then computing the $\eta$ CSSs of length-1 between $S_1$ and $S_2$ requires $O(\max(n + m, \eta))$ time.

**DAG construction**

Given all of the MATCHes found in $A$, our task now is to construct the DAG for $A$. For all paths of the DAG to start and end at a common node, we make MATCHES $S$ and $E$ to respectively precede and succeed the MATCHES in $A$. (Let $S$ have $id = 1$ and $E$ have $id = |A| + 2$ and then store $S$ and $E$ in $A$.) The goal of the DAG is to represent all maximal CSSs between $S_1$ and $S_2$ as paths from $S$ to $E$. We will later find the $LCS$, the longest such path.

In the DAG, the nodes will be the MATCH ids and the edges between MATCHES, say $m_1$ and $m_2$, represent that $S_1[m_1.p1] = S_2[m_2.p2]$ is chosen in the maximal common subsequence followed by $S_1[m_2.p1] = S_2[m_2.p2]$. The DAG is acyclic because, by Definition 1, the LCS is a list of ordered MATCHES. Since we cannot choose $m_1 \in M$ and then $m_h \in M$ with $h < i$, then no cycle can exist.

Our DAG construction, displayed in Algorithm 1, operates in the following way. We initialize the DAG dag by first declaring dag.gr of size $|A|$, since gr will represent all of the nodes. All outgoing edges for say the node $N \in A$ are represented by $dag.gr[N.id][1 \ldots dag.sz[N.id]]$. By setting $dag.sz = [0, \ldots, 0]$, we clear the edges in our dag. Now, setting these edges is the main task of our algorithm.

We can easily construct the edges by assuming that there exists a data structure PREV pv that can tell us the set of parents for each node $a \in A$. That is, we can call getPntrs(pv, $L$) to get the set of nodes $P$ that directly precede MATCH $L \in A$ in the final dag. By “directly precede”, we mean that in the final dag, there is connection from each $p \in P$ to $a$, i.e. each $p$ is in series with $a$, meaning that both $p$ and $a$ are chosen in a maximal CSS. Further, no $p, p_2 \in P$ can be in series with one another, and rather, they are in parallel with one another, meaning that either $p$ or $p_2$ is chosen in a maximal common subsequence.

With $P$, we can build an edge from $a_2 \in P$ to $a$ by first allocating a new space in $dag.gr[a_2.id]$ by incrementing $dag.sz[a_2.id]$ and then making a directed edge from parent to child, i.e. $dag.gr[a_2.id][1 \ldots dag.sz[a_2.id]] = a.id$. After computing the incoming edges for each node $a \in A$, the dag construction is complete.

**PREV data structure**

The simplicity of the DAG construction is due to the PREV pv, detailed here. The $pv$ is composed of four attributes.

**HashSet<int,int> p1.** Suppose that all $a.p1$ values (for $a \in A$) are placed on an integer number line. It is very unlikely that all $a.p1$ values will be consecutive and so, there will be unused numbers (gaps) between adjacent values. Since we later declare matrices on the MATCH
p1 (and p2) values, these gaps will be wasteful. With a scan of the a.p1 values (say using a Set), we can rename them consecutively without gaps; these renamed values are found by accessing HashMap<int,int> p1 with the original a.p1 value.

HashMap<int,int> p2. This is the same as the aforementioned p1, but with respect to the a.p2 values.

MATCH tbl1[][]. A fundamental data structure to support the getPrnts function is the tbl1, defined below.

Definition 3. Max Table w.r.t. p1 (tbl1): Given the set of all MATCH values A and PREV pv on A (with pv.p1 and pv.p2), the tbl1[|pv.p1|][|pv.p2|] is defined such that each tbl1[i][j] is the a ∈ A with the maximum pv.p1.get(a.p1) ≤ i, where pv.p2.get(a.p2) ≤ j. In the case that multiple such a exist, tbl1[i][j] is the a with the rightmost pv.p2.get(a.p2) ≤ j. If no such a exists, tbl1[i][j] = null.

In other words, the tbl1[i][j] stores the “closest” MATCH a with respect to the p1 values (i.e. we maximize a.p1 before a.p2). To construct tbl1, we first declare the table, tbl1[|pv.p1|][|pv.p2|] and initialize all elements tbl1[i][j] = null, signifying that no MATCHes are found. Next, we insert each a ∈ A into the list by setting tbl1[pv.p1.get(a.p1)][pv.p2.get(a.p2)] = a. Now, each tbl1[i][j] = null needs to be set as the rightmost MATCH m with the maximum m.p1 in the subtable tbl1[1...i][1...j]. This is easily computed by first moving vertically in tbl1 and setting tbl1[i][j] = tbl1[i−1][j] if tbl1[i][j] = null to propagate the maximum values vertically. Finally, we need to move horizontally in tbl1 and store in tbl1[i][j] the rightmost tbl1[i][v] (1 ≤ v ≤ j) with the maximum tbl1[i][v].p1. This is done by a left-to-right scan of each row, comparing the adjacent elements, and setting tbl1[i][v] = tbl1[i][v−1] if tbl1[i][v−1].p1 > tbl1[i][v].p1.

MATCH tbl2[][]. The tbl2 is the same as tbl1 except that we define “closest” to mean that the a.p2 value is maximized before the a.p1.

Definition 4. Max Table w.r.t. p2 (tbl2): Given the set of all MATCH values A and PREV pv on A (with pv.p1 and pv.p2), the tbl2[|pv.p1|][|pv.p2|] is defined such that each tbl2[i][j] is the a ∈ A with the maximum pv.p2.get(a.p2) ≤ j, where pv.p1.get(a.p1) ≤ i. In the case that multiple such a exist, tbl2[i][j] is the a with the rightmost pv.p1.get(a.p1) ≤ i. If no such a exists, tbl2[i][j] = null.

The construction of tbl2 is the same as tbl1, except that in the final horizontal scan, we compare tbl2[i][v].p2 and tbl2[i][v−1].p2.

Algorithm 1. Construct the DAG

```c
MATCH { int id, p1, p2 }
MATCH PRNTS { MATCH ml, m2 }
MATCH DAG { int gr[1][1], sz[] }
MATCH PREV { MATCH tbl1[1][1], tbl2[1][1];
MATCH HashMap<int,int> p1, p2 }
MATCH DAG constructDAG(Set<MATCH> A){
    int num=A.size(), sz[|num|=0,...,0]
    MATCH pv=constructPREV(A); MATCH a2
    MATCH DAG dag={ new int[|num|], sz }
    for each a in A {
        Set<MATCH> P=getPrnts(pv,L)
        for each a2 in P {
            dag.gr[a2.id]++
                        dag.sz[a2.id]=a.id
        }
    } return dag
}
```

Algorithm 2. Get the direct parent w.r.t. p1 or p2

```c
MATCH getDPrnt(PREV pv, MATCH L, bool wrtS1){
    return getDPrnt(pv, pv.p1.get(L.p1), pv.p2.get(L.p2), wrtS1)
}
```

Algorithm 3. Get all parents for c.

```c
MATCH getPrnts(PREV pv, MATCH c) {
    Set<MATCH> P
    int i=pv.p1.get(c.p1), j=pv.p2.get(c.p2)
    int q, i1, i2, j1, j2
    MATCH y, dd1, dd2
    MATCH d1=getDPrnt(pv, c, true)
    d2=getDPrnt(pv, c, false)
    if(d1=null & d2=dd) P.add(d1)
    else if(d1=null){
        P.add(dd1)
        i=dd2.p1, j=dd2.p2.get(i)
        P.add(d1)
    }
    else
    {
        MATCH d1={
            for(q=1 to 12) {
                d1=d2.p1, j=dd2.p2.get(j)
                P.add(d1)
            }
        }
    }
    return P
}
```

bool valid(MATCH m, int i1, int i2, int j1, int j2){
    return (m=null & i1.cm.p1<i1 & j1.cm.p2<j2)
}
In terms of construction time, if we assume that adding and accessing HashMap entries are constant time operations, and the Set is implemented with a HashMap, then the PREV pv on A from the n-length S1 and m-length S2 is constructed in \(O(|pv.p| \times |pv.p|)\) time. While pv.p1 and pv.p2 eliminate the gaps between the respective p1 and p2 values of \(A\), we have \(|pv.p| \in O(n)\) and \(|pv.p| \in O(m)\) in the very worst-case.

**Theorem 5.** Given the n-length \(S_1\) and m-length \(S_2\), and the set of all MATCHes \(A\), PREV pv on \(A\) is constructed in \(O(nm)\) time.

**getPrnts function**

Given the PREV pv data structure on all MATCHes \(A\), we call getPrnts(pv, L) in line 11 of constructDAG to retrieve the set of parent MATCHes \(P\) of the MATCH \(L \in A\). Recall that these parents \(P\) of the MATCH \(L\) are all MATCHes that directly precede \(L\) in the DAG, i.e. each \(p \in P\) is in series with \(L\) and no \(p, p' \in P\) are in series with one another. Using pv, we can compute, for any MATCH \(c \in A\), two direct parents that are closest to \(c\) with respect to the \(p_1\) and \(p_2\) values.

**Definition 6.** Direct Parents: Given the PREV pv on the MATCHes in \(A\) between the \(n\)-length \(S_1\) and the \(m\)-length \(S_2\), and a MATCH \(c \in A\), let \(i = pv.p1.get(c.p1)\) and \(j = pv.p2.get(c.p2)\). The direct parent of \(c\) w.r.t. \(p_1\) is:

\[
d_1 = \begin{cases} 
null, & \text{if } i \leq 1 \lor j \leq 1 \lor i > |pv.p| \lor j > |pv.p| \\
pv.tbl[i-1][j-1], & \text{otherwise}
\end{cases}
\]

The direct parent of \(c\) w.r.t. \(p_2\) is:

\[
d_2 = \begin{cases} 
null, & \text{if } i \leq 1 \lor j \leq 1 \lor i > |pv.p| \lor j > |pv.p| \\
pv.tbl[i-1][j-1], & \text{otherwise}
\end{cases}
\]

The first getDPrnt in Algorithm 2 implements Definition 6 to return the direct parents for any MATCH say \(L \in A\). In cases where we want to find the direct parent for a MATCH at a certain location in the pv.tbl1 or pv.tbl2, say pv.tbl1[i][j] or pv.tbl2[i][j], we overload getDPrnt.

The direct parents computation (getDPrnt) is the cornerstone of the getPrnts function. The following lemma, implemented in Algorithm 3, proves that the direct parents of \(c\) can be used to determine all parents of \(c\).

**Lemma 7.** Given \(A\), the MATCHes between \(S_1\) and \(S_2\), and a MATCH \(c \in A\), the two direct parents of \(c\) can be used to compute the set \(P\) with all parents of \(c\).

**Proof.** Let \(d_1\) and \(d_2\) be the direct parents of \(c\) (Definition 6). By Definition 3, \(d_1\) is a direct parent because it directly precedes \(c\) with the maximum \(p_1\) and the rightmost \(p_2\) value. Similarly by Definition 4, \(d_2\) is a direct parent of \(c\) because it directly precedes \(c\) with the maximum \(p_2\) and the rightmost \(p_1\) value. To find the remaining parents of \(c\), we now find other MATCHes that precede \(c\), which are also parallel with \(d_1\) and \(d_2\). There are three cases.

Case (a). When \(d_1 = null\), then also \(d_2 = null\) since there cannot be another MATCH preceding \(c\). Thus, \(P = \emptyset\).

**Algorithm 4.** Compression of target T (given reference R, parameter k, and LPF and POS arrays on \(Z = Ret\) with \(z = |Z|\)) into symbols file (with filename symbols_fn) and triples file (with filename triples_fn).

```
void compress(String T, String R, int k, 
              int LPF[z], int POS[z], 
              String symbols_fn, 
              String triples_fn){
    int i, posT, posZ, len, rlen = R;
    File* symbols=opn(symbols_fn, WRITE)
    File* triples=opn(triples_fn, WRITE)
    for i=r=rlen+1 to z {
        posT=i-rlen 
        len=LPF[i] 
        if (len<k){
            write_char(symbols,T[posT])
        } else {
            write_int(triples,posT)
            posZ=POS[i]
            write_int(triples,posZ)
            write_int(triples,len)
            i=i+len-1
        }
    }close(symbols)
    close(triples)
}
```

Case (b). When \(d_1 = d_2\), the nearest parents to \(c\) are the same MATCH. There are only two types of MATCHes that are parallel with \(d_1\). First, we need to consider all MATCHes, say \(m_1\), with the same endpoint \(m_1.p1 = d_1.p1\) and \(m_1.p2 \in \{1, 2, \ldots, d_1.p2 - 1\}\). Second, we need to consider the MATCHes, say \(m_2\), with the same endpoint \(m_2.p2 = d_1.p2\) and \(m_2.p1 \in \{1, 2, \ldots, d_1.p1 - 1\}\). In the LCS computation, suppose that we choose, w.l.o.g., \(m_1\) (with \(m_1.p2 = d_1.p2 - 2\)) instead of \(d_1\). Then, we cannot choose a MATCH \(m_3\) with \(m_3.p1 < d_1.p1\) and \(m_3.p2 = d_1.p2 - 1\). So, having any \(m_1\) or \(m_2\) parallel to \(d_1\) will only lead to suboptimal CSSs. Thus, only \(P = \{d_1\}\) is a parent of \(c\).

Case (c). Otherwise, \(d_1 \neq d_2\) and we have two different direct parents of \(c\). Set \(P = \{d_1, d_2\}\). Let us collect the endpoints of \(d_1\) and \(d_2\): \(i_1 = d_1.p1, i_2 = d_1.p1, j_1 = d_1.p2, j_2 = d_2.p2\). What MATCH, say \(m_3\), is parallel to \(d_1\) and \(d_2\)? By Definition 6, there cannot be any MATCH \(m_3\) directly preceding \(c\) with endpoints after \(i_2\) or \(j_2\). By (b), we do not need to consider other MATCHes.
Algorithm 5. Decompressing target $T$ using reference $R$, symbols file (with filename symbols_fn), and triples file (with filename triples_fn).

```java
String decompress(String R, String symbols_fn, String triples_fn) {
    String T = ...
    int s=1, t=1, i=1, j, beg=0, len=loc=rlen=R.length();
    int slen = num_bytes(symbols_fn)/sizeof(int);
    File* symbols = open(symbols_fn, READ);
    File* triples = open(triples_fn, READ);
    bool more = true
    while (more) {
        loc = peek_at_int(triples)
        if (loc <= slen && loc "LOCK")
            get_next_int(triples) // discard
        beg = get_next_int(triples)
        len = get_next_int(triples)
        t = 3
        for j = beg to beg + len - 1 {
            i++
            if (j < rlen) T = T[j-rlen]
            else T = R[j]
        }
        if (s <= slen) {
            i++
            s++
            T = get_next_char(symbols)
        } else more = false
    }
    close(symbols)
    close(triples)
    return T
}
```

with endpoints on either $d1$ or $d2$. So, all the possible MATCHes parallel to $d1$ and $d2$ are those with $(m3,p1 \in w \land m3.p2 \in x)$, where $w = (i1 + 1, i1 + 2, \ldots, i2 - 1)$ and $x = (j1 + 1, j1 + 2, \ldots, j2 - 1)$. To find such $m3$, we only need to find direct parents (by (b)), say $dd1$ and $dd2$, for a theoretical MATCH $m$ with $(m.p1 \in w \land m.p2 \in x)$. Then, when we have $i1 < dd1.p1 < i2$ and $j1 < dd2.p2 < j2$, this is a possible MATCH parallel with $d1$ and $d2$, which is also a possible parent of $c$, so we add $dd1$ to $P$. We do the same process for $dd2$.

Since we computed all the possible parents in $P$, additional processing on $P$ is needed to ensure that no pair of MATCHes in $P$ are in series; if any are in series, delete the MATCH furthest from $c$. With the $pv$ and $getDPrnt$, this task is simple. We simply check the direct parents (say $dd1$ and $dd2$) for each $y \in P$, and remove $dd1$ if $dd1 \in P$ and remove $dd2$ if $dd2 \in P$.

Computing the LCS

Since our dag has a single source $S$ (and all paths end at $E$), the longest path between $S$ and $E$, i.e. the LCS, is computed by giving all edges a weight of $-1$ and finding the shortest path from $S$ to $E$ via a topological sort [32].

Complexity analysis

Our LCS algorithm: (i) finds the length-1 CSSs, (ii) computes the DAG on the CSSs, and (iii) reports the longest DAG path. Here, we analyze the overall time complexity.

Step (i)

First, we find (and store in $A$) the $\eta$ length-1 CSSs in $O((n + m, \eta)$ time by Lemma 2.

Step (ii)

We then construct the DAG $dag$ on these $a \in A$ with constructDAG. In constructDAG, we initially compute the newly proposed PREV $pv$ data structure in $O(nm)$ time by Theorem 5. After constructing $pv$, the computeDAG iterates through each $a \in A$ and creates an incoming edge between the parents of $a$ and $a$. So, constructDAG executes in time $O((nm, \eta \times t_{getPrnt})$, where $t_{getPrnt}$ is the time of getPrnts. The getPrnts running time is in $O((i2 - i1) + (j2 - j1)$, with respect to the local variables $i1, i2, j1, and j2$. However, it may be the case that $i1 = j1 = 1, i2 = n, and j2 = m$, and so $O(n + m)$ time is required by getPrnts. Below we formalize the worst-case result and the case for average strings from a uniform distribution.

Lemma 8. For the $n$-length $S_1$ and the $m$-length $S_2$, the getPrnts function requires $O(n + m)$ time.

Lemma 9. For average case strings $S_1$ and $S_2$ with symbols uniformly drawn from alphabet $\Sigma$, the getPrnts function requires $O(|\Sigma|)$ time.

Proof. Since $d1$ and $d2$ are the direct parents of $c$ (see Definitions 3, 4 and 6), and since the uniformity of $S_1$ and $S_2$ means that for any symbol say $S_1[s]$ we can find every $\sigma \in \Sigma$ in $S_2[s - \Delta, \ldots, s + \Delta]$ with $\Delta \in O(|\Sigma|)$, then $(i2 - i1) \in O(|\Sigma|)$ and $(j2 - j1) \in O(|\Sigma|)$.

So, the overall constructDAG time follows.

Theorem 10. Given $A$, the length-1 MATCHes in the $n$-length $S_1$ and the $m$-length $S_2$, the constructDAG requires $O(max(nm, \eta \times max(n, m))$ time in the worst-case and $O(max(nm, \eta \times |\Sigma|)$ on average.

Step (iii)

We find the LCS with a topological sort in time linear to the dag size [32], which cannot require more time than that needed to build the dag (see Theorem 10).

Summary

Overall, (i) and (iii) do not add to the complexity of (ii). Given the above, the overall running time is as follows.

Theorem 11. The LCS between the $n$-length $S_1$ and the $m$-length $S_2$ can be computed in $O(max(nm, \eta \times max(n, m))$ time in the worst-case and $O(max(nm, \eta \times |\Sigma|)$ on average.
Compressing resequencing data

When data is released, modified, and re-released over a period of time, a large amount of commonality exists between these releases. Rather than maintaining all uncompressed versions of the data, it is possible to keep one uncompressed version, say $D$, and compress all future versions $D_t$ with respect to $D$. We refer to $D_t$ as the target and $D$ as the reference. This idea is used to compress resequencing data in [20, 21], primarily using the LCS. The LCS, however, has two core problems with respect to compression. For very similar sequences, the LCS computation time is almost quadratic, or worse, potentially leading to long compression time. Secondly, the LCS may not always lead to the best compression, especially when some CSS components are very short.

Rather than focusing on the LCS, we consider the maximal CSSs that make up the common subsequences. To intelligently choose which of the CSSs are likely to lead to improved compression, we use the longest previous factor (LPF), an important data structure in text compression [33]. Consider compressing the target $T$ with respect to the reference $R$; let $Z = R \circ T$. Suppose we choose exactly $|T|$ maximal-length CSSs, specifically, for $\beta = Z[i \ldots |Z|]$ we have $\alpha = Z[h \ldots |Z|]$ such that (1) CSSs $\alpha[1 \ldots k] = \beta[1 \ldots k]$ and (2) this is the maximal $k$ for $h < i$, where $|R| + 1 \leq i \leq |Z|$. These $k$s are computed in the LPF data structure on $Z$ at $LPF[i] = k$ and the position of this CSS is at $POS[i] = h$ [29]. (Note that LPF and POS are constructed in linear time [29–31].) The requirement that $h < i$ suits dictionary compression and compressing resequencing data because the CSS beginning at $i$ is compressed by referencing the same CSS at $h$, occurring earlier in target $T$ or anywhere in the reference $R$. Our idea is to use the LPF and POS to represent or encode CSSs that make up the target $T$ with tuples. We will then compress these tuples with standard compression schemes.

Our compression scheme

We now propose a reference-based compression scheme which scans the LPF and POS on $Z$ in a left-to-right fashion to compress $T$ with respect to $R$. This scheme is similar to the LZ factorization [29], but differs in how we will encode the CSSs. Our contribution here is (1) using two files to compress $T$, (2) only encoding CSSs with length at least $k$, and (3) further compressing the resulting files with standard compression schemes.

Initially, the two output files, triples and symbols, are empty. Let $i = |R| + 1$.

(1) If $LPF[i] < k$, we simply encode the symbol; append the (say 1-byte) char $T[i - |R|]$ to symbols and increment $i$. Otherwise $LPF[i] \geq k$, so we will encode this CSS with the triple $(pT, pZ, l)$, where $pT = i - |R|$ is the starting position of the CSS in $T$, $pZ = POS[i]$ is the starting position of the CSS in $Z[1 \ldots i - 1]$, and $l = LPF[i]$ is the length of the CSS. We write three long (say 4-byte integer) words $pT, pZ,$ and $l$ to triples. Since the triple encodes an $l$-length CSS, we set $i = i + l$ to consider compressing the suffix following the currently encoded CSS. Lastly, if $i \leq |Z|$, continue to (!).

The resulting files triples and symbols are binary sequences that can be further compressed with standard compression schemes (so, decompression will start by first reversing this process). The purpose of the $k$ and the two files (one with byte symbols and one with long triples) is to introduce flexibility into the system and encode CSSs with triples (12 bytes) only when beneficial and otherwise, encode a symbol with a byte. For convenience, our implementation encodes each symbol with a byte, but we acknowledge that it is possible to work at the bit-level for small alphabets.

The decompression is also a left-to-right scan. Let $i = 1$ and point to the beginning of triples and symbols.

(1) Consider the current long word $w_i$ in triples. According to the triple encoding, this will be the position...
of the CSS in \( T \). If \( i = w_1 \), then we pick up the next two long words \( w_2 \) and \( w_3 \) in triples. We now know \( T[i \ldots i + w_3 - 1] = Z[w_2 \ldots w_2 + w_3 - 1] \). Since we only have access to \( R \) and \( T[1 \ldots i - 1] \), then we pick up each symbol of \( Z[w_2 \ldots w_2 + w_3 - 1] \) by picking up \( R[j] \) if \( j \leq |R| \) and picking up \( T[j - |R|] \) otherwise, for \( w_2 \leq j \leq w_2 + w_3 - 1 \). We next will consider \( i = i + w_3 \). Else \( i \neq w_1 \), so we pick up the next char \( c \) in symbols since \( T[i] = c \); we next consider \( i++ \). If \( i \leq |T| \), go to (†).

The compression and decompression algorithms are detailed in Algorithms 4 and 5, respectively.

**Results and discussion**

We implemented the previously described compression scheme, selected and fixed parameter \( k \), and ran our program to compress various DNA corpora. In this section, we describe the selection of \( k \) and our final results.

### Choosing parameter \( k \)

Recall that the parameter \( k \) is a type of threshold used by our compression scheme to determine whether it is more beneficial to encode a symbol verbatim (that is, 1 byte) or encode a CSS as a triple (that is, 12 bytes). Specifically, our compression algorithm works on the LPF (which represents the CSSs of the \( n \)-length \( T \)) in a left-to-right fashion, selecting the leftmost CSS, say \( T[i \ldots i + l - 1] \) of length-\( (LPF[i] = l) \), and determining whether to encode that CSS as a triple [and then consider the next CSS \( T[i + 1 \ldots i + 1 + LPF[i + 1] - 1] \) of length-\( (LPF[i + 1]) \)], or encode the first symbol \( (T[i]) \) [and then consider the next CSS \( T[i + 1 \ldots i + LPF[i + 1] + 1] \) of length-\( (LPF[i + 1]) \)].

Obviously, it is better to encode a length-\( (l = 1) \) CSS with a 1-byte symbol, rather than a 12-byte triple. It is clearly the case that for any CSS length \( 1 \leq l < 12 \), it is better to encode the first symbol with 1-byte and take a chance that the next CSS to the right will be significantly
larger. Why can we afford to take this chance? One LPF property, which also allows for an efficient construction of the data structure (see [29]), is that $\text{LPF}[i + 1] \geq \text{LPF}[i] - 1$. That is, if we pass up on encoding the CSS at $i$ of length-$(\text{LPF}[i] = l)$ as a triple, we can encode $T[i]$ as a symbol and (1) are guaranteed that there is at least a length-$(l - 1)$ CSS with a prefix of $T[i + 1 \ldots n]$ and (2) the longest CSS common to a prefix of $T[i + 1 \ldots n]$ is of length-$\text{LPF}[i + 1]$, maybe even larger than $\text{LPF}[i]$. Clearly, we want to encode most CSSs as triples to take advantage of the concise triple representation. Now, the question becomes: how large should we set $k$, such that we can afford to take a risk passing up length-$(l < k)$ CSSs in hopes of finding even larger CSSs better suited as triples?

For this paper, we decided to select $k$ by studying the impact of the parameter on our compressed results for the Arabidopsis thaliana genome, using target TAIR9 and reference TAIR8. The compression results for various $k$ are shown in Fig. 2; since chromosome 4 does not compress as well as the others, we show it separately in Fig. 3 for improved visualization. For very small $k < 12$, we have a result that basically encodes with triples only; when $k = 1$, we are exclusively encoding CSSs as triples. We see that when $k$ is roughly between 12 and 35, we are encouraging the algorithm to pass up encoding smaller CSSs as triples, which leads to the best compression result. The results stay competitive until say $k \geq 100$, where the algorithm becomes too optimistic and passes up the opportunity to encode smaller CSSs as triples in hopes that larger CSSs will exist. Further, we see from Fig. 4 that as $k$ becomes large, it indeed becomes very expensive to pass up encoding these CSSs as triples. Also, we see from Fig. 5 that beyond say $k = 20$, there is minimal compression savings. Thus, we want to balance the expensive symbols files with the space-savings from the triples files.

In Table 1, we show the best compression results and $k$ for the Arabidopsis thaliana genome. Unless otherwise specified, our experiments below fix parameter $k$ as 31, since it is the optimal $k$ common to 4-of-5 of the Arabidopsis thaliana chromosomes and gives a competitive result for the remaining chromosome. This result follows intuition because $k$ should be at least 11 and not too large, so that we can consider CSSs that are worthy of encoding.

**Compression results**

Like [20, 21], we compress the Arabidopsis thaliana genome chromosomes in TAIR9 (target) with respect to TAIR8 (reference). In Table 2, we display the compression results. We see that all of our results are competitive with the GRS and GReEn systems, except for chromosome 4, which has the smallest average CSS length of about 326K. Nonetheless, we are able to further compress our results using compression schemes from 7-zip, with L and P respectively representing 1zma2 and ppm, to achieve even better compression.

In Table 3, we show results for compressing the genome Oryza sativa using the target TIGR6.0 and reference TIGR5.0. After compressing our algorithm’s output with 1zma2 or ppm, our results are better than both GRS

| $U$ | $k$ | $C$ |
|-----|-----|-----|
| 1   | 31–35 | 1086 |
| 2   | 16–1578 | 504  |
| 3   | 24–39  | 746  |
| 4   | 18     | 4418 |
| 5   | 19–91  | 433  |
Table 2 Arabidopsis thaliana genome: Results (in bytes) for compressing chromosome $U$ into $C$

| $U$ | $|U|$ | Our Scheme | GRS | GReEn |
|-----|------|-----------|-----|--------|
|     |      | $|C|$ | $|L(C)|$ | $|P(C)|$ | [20] | [21] |
| 1   | 30,427,671 | 1,086 | 963 | 1,037 | 715 | 1,551 |
| 2   | 19,698,289 | 504 | 584 | 605 | 385 | 937 |
| 3   | 23,459,830 | 746 | 759 | 803 | 2,989 | 1,097 |
| 4   | 18,585,056 | 4,555 | 2,507 | 3,156 | 1,951 | 2,356 |
| 5   | 26,975,502 | 433 | 502 | 520 | 604 | 618 |
| Sum | 119,146,348 | 7,324 | 5,315 | 6,121 | 6,644 | 6,559 |

Bold signifies the best result.

[20] and GReEn [21]. Note that for each of the chromosomes 6, 9, and 12, our algorithm’s output is 12 bytes, better than that of GRS [20] (14 bytes) and GReEn [21] (482 bytes, 366 bytes, and 429 bytes respectively). When we compress our result with $lzma2$ or $ppmd$, the result is bloated since more bytes are needed. So, we can further improve the overall result by not compressing chromosomes 6, 9, and 12, and further, selecting the best such compression scheme for each individual chromosome. We acknowledge that additional bits would need to be encoded to determine which compression scheme was selected.

In Table 4, we show the compression results for the Homo sapiens genome, using KOREF_20090224 as the target and KOREF_20090131 as the reference. After compressing our computed symbols and triples files with $lzma2$, we see that most all of our results are better than GRS and GReEn. Recall in previous experiments that sometimes secondary compression with 7-zip does not improve the initial compression achieved by our proposed algorithm. For this genome, we exercise the flexibility of our compression framework. In Table 4, (*) indicates that the $M$ chromosome was not further compressed with $lzma2$ due to the aforementioned reason. To indicate that $M$ was not compressed, we will simply encode a length-25 bitstring (say 4-byte) header to specify whether or not the $lzma2$ was applied. There is no need to encode $k$ in the header since it is a fixed value. Thus, the overall compressed files require 15,460,478 bytes, which is only slightly better than GRS and GReEn.

To improve this result, we exploit the difference between the Homo sapiens genome and those discussed earlier. That is, the Homo sapiens genome uses the extended alphabet {A, C, G, K, M, R, S, T, W, Y, a, c, g, k, m, n, r, s, t, w, y}. The observation is that, the alphabet size decreases roughly in half by converting to one character-case. Such a significant reduction in alphabet size will yield more significant redundancies identified by our compression algorithm. Our new decomposition method will decompose each chromosome into two parts: (1) the payload ($\rho$), representing the chromosome in one character-case, and (2) the character-case bitstring ($\alpha$), in which each bit records whether the corresponding position in the target was an upper-case character. Next, we use our previously proposed algorithm to compress $\rho$ into $C^\rho$ and $\alpha$ into $C^\alpha$.

Table 5 shows compression via decomposition for the Homo sapiens genome. Note that the $|C^\rho|$, i.e. compressed payload, column corresponds to the results reported in our initial work [1]. We observe that in various scenarios, the character-case of the alphabet

Table 3 Oryza sativa genome: Results (in bytes) for compressing chromosome $U$ into $C$

| $U$ | $|U|$ | Our Scheme | GRS | GReEn |
|-----|------|-----------|-----|--------|
|     |      | $|C|$ | $|L(C)|$ | $|P(C)|$ | [20] | [21] |
| 1   | 43,268,879 | 15,207 | 4,735 | 4,551 | 1,502,040 | 4,972 |
| 2   | 35,930,381 | 4,645 | 1,649 | 1,517 | 1,409 | 1,906 |
| 3   | 36,406,689 | 54,234 | 15,693 | 15,556 | 47,764 | 17,890 |
| 4   | 35,278,225 | 21,474 | 6,636 | 6,432 | 36,145 | 6,750 |
| 5   | 29,947,789 | 17,030 | 5,431 | 5,359 | 6,177 | 5,539 |
| 6   | 31,246,789 | 12 | 146 | 141 | 14 | 482 |
| 7   | 29,696,629 | 5,899 | 2,064 | 1,972 | 4,067 | 2,448 |
| 8   | 28,439,308 | 23,126 | 8,794 | 10,115 | 118,246 | 9,507 |
| 9   | 23,011,239 | 12 | 146 | 141 | 14 | 366 |
| 10  | 23,134,759 | 175,228 | 49,713 | 50,277 | 788,542 | 60,449 |
| 11  | 28,512,666 | 41,407 | 13,006 | 13,351 | 2,397,470 | 14,797 |
| 12  | 27,497,214 | 12 | 146 | 141 | 14 | 429 |
| Sum | 372,317,567 | 358,286 | 108,159 | 109,553 | 4,901,902 | 125,535 |

Bold signifies the best result.
### Table 4 Homo sapiens genome: Results (in bytes) for compressing chromosome U into C

| U       | |U| | Our Scheme | |L(C)| |GRS | |GReEn |
|---------|---|---|-------------|---|---|---|---|---|---|
| 1       | 247,249,719 | 2,836,652 | 1,082,859 | 1,336,626 | 1,225,767 |
| 2       | 242,951,149 | 2,871,186 | 1,050,170 | 1,354,059 | 1,272,105 |
| 3       | 199,501,827 | 2,115,410 | 790,444 | 1,011,124 | 971,527 |
| 4       | 191,273,063 | 2,398,432 | 910,898 | 1,139,225 | 1,074,357 |
| 5       | 180,857,866 | 2,064,874 | 764,458 | 988,070 | 947,378 |
| 6       | 170,899,992 | 1,902,067 | 710,355 | 906,116 | 865,448 |
| 7       | 158,821,424 | 2,326,721 | 844,194 | 1,096,646 | 998,482 |
| 8       | 146,274,826 | 1,617,884 | 617,996 | 764,313 | 729,362 |
| 9       | 140,273,252 | 1,877,509 | 704,205 | 864,222 | 773,716 |
| 10      | 135,374,737 | 1,623,010 | 617,633 | 768,364 | 717,305 |
| 11      | 134,452,384 | 1,586,558 | 604,901 | 755,708 | 716,301 |
| 12      | 132,349,534 | 1,476,523 | 566,997 | 702,040 | 668,455 |
| 13      | 114,142,980 | 1,100,576 | 399,527 | 520,598 | 490,888 |
| 14      | 106,368,585 | 1,026,227 | 377,695 | 484,791 | 451,018 |
| 15      | 100,338,915 | 1,055,663 | 398,720 | 496,215 | 453,301 |
| 16      | 88,827,254  | 1,225,378 | 443,009 | 567,989 | 510,254 |
| 17      | 78,774,742  | 1,081,739 | 396,371 | 505,979 | 464,324 |
| 18      | 76,117,153  | 865,138   | 320,361 | 408,529 | 378,420 |
| 19      | 63,811,651  | 862,129   | 320,789 | 399,807 | 369,388 |
| 20      | 62,435,964  | 605,179   | 229,418 | 282,628 | 266,562 |
| 21      | 46,944,323  | 488,340   | 180,096 | 226,549 | 203,036 |
| 22      | 49,691,432  | 568,734   | 205,244 | 262,443 | 230,049 |
| X       | 154,913,754 | 7,525,925 | 2,494,884 | 3,231,776 | 2,712,153 |
| Y       | 57,772,954  | 1,343,260 | 429,099 | 592,791 | 481,307 |
| M       | 16,571      | 151       | 151(*) | 183 | 127 |
| Sum     | 3,080,436,051 | 42,445,265 | 15,460,474 | 19,666,791 | 17,971,030 |

Bold signifies the best result.

Symbol is not significant. For example, the IUB/I-UPAC amino acid and nucleic acid codes use only upper-case letters (see http://www.bioinformatics.org/sms/iupac.html). Also, some environments and formats (such as FASTA) do not distinguish between lower-case and upper-case. According to the NCBI website for BLAST input formats (see http://blast.ncbi.nlm.nih.gov/blastcgihelp.shtml): “Sequences [in FASTA format] are expected to be represented in the standard IUB/IUPAC amino acid and nucleic acid codes, with these exceptions: lower-case letters are accepted and are mapped into upper-case; …” Further, some programs/environments use character cases for improved visualization, as is the case with the USC Genome Browser, which uses lower-case to show repeats from RepeatMasker and Tandem Repeats Finder (ftp://hgdownload.cse.ucsc.edu/goldenPath/hg38/chromosomes/README.txt).

Also, we see that further compressing the payload with lgma2 more than doubles the compression ratio. Interestingly, the payload (ρ) compresses much better than the character-case bitstring (α). Nonetheless, the compression via decomposition (in Table 5) yields a compression ratio of 360, a significant improvement over the 199 compression ratio when compressing the genome's characters in their native character-case (in Table 4). As described earlier, we do not further compress chromosome M after initial coding for the symbols and triplets, and thus encode only a 4-byte header to remember this decision, given that the payload and character-case bitstring k values are fixed. Thus, 8,556,708 bytes are needed, which is an improvement over GRS and GReEn.

Theoretically, our compression scheme requires time linear in the length of the uncompressed text, since we perform one scan of the LPF, which is constructed in linear time via the suffix array SA [29]. For the Arabidopsis...
Table 5 Homo sapiens genome: Results (in bytes) for compressing chromosome $U$ via decomposition, i.e. compressing the payload ($\rho$) into $C^\rho$ and compressing the character-case bitstring $\alpha$ into $C^\alpha$.

| $U$ | $|U|$ | Our Scheme | GReEn |
|-----|------|------------|------|
|     | $|C^\rho|$ | $|L(C^\rho)|$ | $|C^\alpha|$ | $|L(C^\alpha)|$ | $|L(C^\rho)| + |L(C^\alpha)|$ | (20) | (21) |
| 1   | 247 249 719 | 381 577 | 161 319 | 755 092 | 447 919 | 609 238 | 1 336 626 | 1 225 767 |
| 2   | 242 951 149 | 356 526 | 153 805 | 756 823 | 452 338 | 606 143 | 1 354 059 | 1 272 105 |
| 3   | 199 501 827 | 284 096 | 119 348 | 553 835 | 343 213 | 462 561 | 1 011 124 | 971 527 |
| 4   | 191 273 063 | 330 381 | 137 301 | 619 981 | 383 882 | 521 183 | 1 139 225 | 1 074 357 |
| 5   | 180 857 866 | 259 922 | 109 768 | 550 876 | 331 063 | 440 831 | 988 070 | 947 378 |
| 6   | 170 899 992 | 265 222 | 110 544 | 508 662 | 310 029 | 420 573 | 906 116 | 865 448 |
| 7   | 158 821 424 | 292 797 | 121 289 | 611 475 | 355 616 | 476 905 | 1 096 646 | 998 482 |
| 8   | 146 274 826 | 222 972 | 93 378 | 434 420 | 261 455 | 354 833 | 764 313 | 729 362 |
| 9   | 140 273 252 | 309 512 | 132 957 | 493 024 | 276 468 | 409 425 | 864 222 | 773 716 |
| 10  | 135 374 737 | 245 264 | 103 115 | 436 272 | 257 895 | 361 010 | 768 364 | 717 305 |
| 11  | 134 452 384 | 222 735 | 92 471 | 423 687 | 254 637 | 347 108 | 755 708 | 716 301 |
| 12  | 132 349 534 | 214 123 | 88 447 | 393 764 | 239 811 | 328 258 | 702 040 | 668 455 |
| 13  | 114 142 980 | 148 938 | 62 730 | 301 116 | 183 038 | 245 768 | 520 598 | 490 888 |
| 14  | 106 368 585 | 141 128 | 57 354 | 286 839 | 170 916 | 228 270 | 484 791 | 451 018 |
| 15  | 100 338 915 | 138 219 | 58 777 | 302 957 | 173 600 | 232 377 | 496 215 | 453 301 |
| 16  | 88 827 254 | 151 606 | 62 779 | 346 282 | 191 190 | 253 969 | 567 989 | 510 254 |
| 17  | 78 774 742 | 136 168 | 57 030 | 301 837 | 171 680 | 228 710 | 505 979 | 464 324 |
| 18  | 76 117 153 | 113 469 | 47 122 | 241 437 | 140 909 | 188 031 | 408 529 | 378 420 |
| 19  | 63 811 651 | 130 468 | 53 531 | 230 673 | 134 701 | 188 232 | 399 807 | 369 388 |
| 20  | 62 435 964 | 94 273 | 38 689 | 169 584 | 99 796 | 138 485 | 282 628 | 266 562 |
| 21  | 46 944 323 | 71 121 | 28 744 | 141 387 | 79 835 | 108 579 | 226 549 | 203 036 |
| 22  | 49 691 432 | 81 329 | 33 663 | 164 026 | 89 961 | 123 624 | 262 443 | 230 049 |
| X   | 154 913 754 | 523 282 | 196 868 | 1 533 249 | 875 026 | 1 071 894 | 3 231 776 | 2 712 153 |
| Y   | 57 772 954 | 152 464 | 57 002 | 300 287 | 153 582 | 210 584 | 592 791 | 481 307 |
| M   | 16 571 | 64 | 64(*) | 49 | 49(*) | 113 | 183 | 127 |
| Sum | 3 080 436 051 | 5 267 656 | 2 178 095 | 10 857 634 | 6 378 609 | 8 556 704 | 19 666 791 | 17 971 030 |

Bold signifies the best result

**thaliana** and Oryza sativa genomes, we ran our programs on a laptop; for the Homo sapiens genome, we ran our programs in an AWS EC2 m4.4xlarge environment. Consider, for example, the larger chromosomes of the Homo sapiens genome. For a payload ($\rho$), the SA construction required 2,376 sec and the LPF construction required 399 sec. Note that depending on the application, the SA and LPF may already be available. Given the LPF, our compression algorithm completed in less than one second. Decompression is also fast, and lightweight, since no data structures are required as parameters. Our future plan includes using more efficient SA and LPF constructions.

**Conclusions**

We proposed a new algorithm to compute the LCS. Motivated by our algorithm, we introduced a new reference-based compression scheme for genome resequencing data using the LPF. For the Arabidopsis thaliana genome (originally 119,146,348 bytes), our scheme compressed the genome to 5315 bytes, an improvement over the best performing state-of-the-art methods (6644 bytes [20] and 6559 bytes [21]). For the Oryza sativa genome (originally 372,317,567 bytes), our scheme compressed the genome to 108,159 bytes, an improvement over the 4,901,902 bytes in [20] and the 125,535 bytes in [21]. We also experimented with the Homo sapiens genome (originally 3,080,436,051 bytes), which was compressed to 19,666,791 bytes and 17,971,030 bytes in [20] and [21], respectively. By applying our scheme via a decomposition approach, we compress the genome to 8,556,708 bytes, and if alphabet character-case is not significant, we compress the genome to 2,178,095 bytes. Further improvement can be obtained by choosing the $k$ parameter for each specific chromosome, or each specific species.
Declarations
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Authors’ contributions
All authors contributed to the core elements of this work. DA initiated the project. RB, TA, and DA developed the LCS algorithm. DA and RB developed the compression algorithm. RB implemented the LPS and compression methods. AF and RB performed the experiments. DA coordinated the overall project. RB and DA prepared the final manuscript. All authors read and approved the final manuscript.

Competing interests
The authors declare that they have no competing interests.

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