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Hardness of Covering Alignment: Phase Transition in Post-Sequence Genomics

Romeo Rizzi, Massimo Cairo, Veli Mäkinen, Alexandru I. Tomescu and Daniel Valenzuela

Abstract—Covering alignment problems arise from recent developments in genomics; so called pan-genome graphs are replacing reference genomes, and advances in haplotyping enable full content of diploid genomes to be used as basis of sequence analysis. In this paper, we show that the computational complexity will change for natural extensions of alignments to pan-genome representations and to diploid genomes. More broadly, our approach can also be seen as a minimal extension of sequence alignment to labelled directed acyclic graphs (labelled DAGs). Namely, we show that finding a covering alignment of two labelled DAGs is NP-hard even on binary alphabets. A covering alignment asks for two paths \( R_1 \) (red) and \( G_1 \) (green) in DAG \( D_1 \) and two paths \( R_2 \) (red) and \( G_2 \) (green) in DAG \( D_2 \) that cover the nodes of the graphs and maximize the sum of the global alignment scores:

\[
\text{as}(\text{sp}(R_1), \text{sp}(R_2)) + \text{as}(\text{sp}(G_1), \text{sp}(G_2)),
\]

where \( \text{sp}(P) \) is the concatenation of labels on the path \( P \). Pair-wise alignment of haplotype sequences forming a diploid chromosome can be converted to a two-path coverable labelled DAG, and then the covering alignment models the similarity of two diploids over arbitrary recombinations. We also give a reduction to the other direction, to show that such a recombination-oblivious diploid alignment is NP-hard on alphabets of size 3.

Index Terms—alignment, edit distance, directed acyclic graph, diploid genome, pan-genome, NP-hard problem

1 INTRODUCTION

For decades, sequence alignments have played a central role in computational molecular biology and especially in computational genomics. Interestingly, being a fundamental computer science problem, there has been a constant interplay with developments in theoretical computer science forums around the problem, and the development of practical bioinformatics tools. Most notably, this connection is visible in the so-called read aligners [12], [13] that use Burrows-Wheeler indexing techniques [7], [9]. A recent breakthrough [2] connects the difficulty of finding sub-quadratic time solution to pair-wise alignment to a complexity theory question. There are still open questions around the basic sequence alignment setting (e.g. indexed approximate pattern matching), but at the same time the computational genomics community is moving towards abstractions beyond sequences, where even the most fundamental questions are open. One of the latest trends is to replace a reference genome with a pan-genome variant graph [18], with a backbone consisting of a reference sequence and alternative paths encoding common variants observed in a population. A basic theoretical question and decisive technological issue is how the role of the sequence alignment toolbox and conceptual framework can scale up in elaborating this more structured data, a world intrinsically populated by labelled directed graphs, that in many cases we can assume to be acyclic at least to a large extent (labelled DAGs) (see e.g. [18]). One possible formulation is to ask for the minimum number of edits to convert one graph to another; this is MAX SNP-hard even when the input DAGs are unordered trees [1].

In this paper, we refine the tractability border of alignments by focusing on graphs that are as close to sequences as possible. Namely, we focus on labelled DAGs that are coverable by two paths. A covering alignment asks for two paths \( R_1 \) (red) and \( G_1 \) (green) in DAG \( D_1 \) and two paths \( R_2 \) (red) and \( G_2 \) (green) in DAG \( D_2 \) that cover the nodes of the graphs and maximize the sum of the global alignment scores:

\[
\text{as}(\text{sp}(R_1), \text{sp}(R_2)) + \text{as}(\text{sp}(G_1), \text{sp}(G_2)),
\]

where \( \text{sp}(P) \) is the concatenation of labels on the path \( P \). We show that this problem is NP-hard even on binary alphabets. A more principled way to derive this formulation comes from modeling diploid genomes [16], where the labelled DAG is a grid graph denoting a pair-wise alignment of haplotypes. We defer the detailed derivation and applications of this natural similarity measure in the context of diploid alignments to Section 4. We show that this restricted variant of covering alignment, called recombination-oblivious diploid alignment problem, is NP-hard on alphabets of size 3. This problem becomes polynomial time solvable once one of the input alignments needs not be covered by the optimal solution, or when the problem is otherwise similarly relaxed [11], [16], [17] (see Section 4).

We hope these results are starting points for a more systematic study of sequence analysis in the era of post-sequence genomics: Our reduction from multiple alignment to covering alignment of two labelled DAGs would seem to indicate that problems that are NP-hard on many sequences become NP-hard already on two inputs with higher level abstractions. As the reduction works on a binary alphabet,
on DAGs minimally harder than sequences, and natural relaxations of the problem become solvable in polynomial time, we have thus identified a phase transition between polynomially-solvable and NP-hard alignment problems.

Our reductions follow a general approach introduced in [19] to show the NP-completeness of the problem of deciding whether a string is a square.

2 PRELIMINARIES

2.1 Problem definition

Let $\Sigma$ be a finite alphabet. We use $\Sigma^*$ to denote the set of all strings over $\Sigma$ and $\Sigma^+$ to denote the set of all non-empty strings over $\Sigma$. In this paper we will also use the term sequence to denote a string. The empty string is denoted by $\varepsilon$, and $\Sigma_\varepsilon$ denotes $\Sigma \cup \{\varepsilon\}$. For a string $A = a_1 a_2 \cdots a_n \in \Sigma_\varepsilon$ denotes its length, namely $\ell$, and $A[j]$ denotes its $j$th symbol, namely $a_j$. We will use the indexed product notation $\prod$ to denote repeated concatenation of strings. The edit distance of strings $A$ and $B$, denoted $ed(A, B)$, is defined as the minimum number of deletions, insertions and substitutions of symbols to convert $A$ into $B$.

For $\Sigma' \in \{\Sigma, \Sigma_\varepsilon, \Sigma^*, \Sigma^+\}$, a $\Sigma'$-DAG is a tuple $D = (D, \ell)$, where $D = (V, E)$ is a DAG with $|V|$ nodes and $|E|$ edges, and $\ell : V \mapsto \Sigma'$ is a total function on $V$. For a path $P = v_1, \ldots, v_l$ in $D$, the spelling of $P$, denoted $sp(P)$, equals $\ell(v_1) \cdots \ell(v_l)$. We say that a set $P$ of paths in $D$ is a path cover if every node in $V$ appears in some $P \in P$. The width of $D$ equals the minimum cardinality of a path cover of $D$.

For $\Sigma' \in \{\Sigma, \Sigma_\varepsilon, \Sigma^*, \Sigma^+\}$ we consider the following problem:

**Path covers of minimum edit distance in two $\Sigma'$-DAGs (PC-MIN-ED-$\Sigma'$)**

**INPUT:** Two $\Sigma'$-DAGs $D_1 = (D_1, \ell_1)$ and $D_2 = (D_2, \ell_2)$ of width 2.

**OUTPUT:** A path cover $\{R_1, G_1\}$ of $D_1$ and a path cover $\{R_2, G_2\}$ of $D_2$ minimizing $ed(sp(R_1), sp(R_2)) + ed(sp(G_1), sp(G_2))$.

Here $R$ works as an analogy to red path and $G$ works as an analogy to green path.

2.2 Edit distance and optimal alignments

The notion of edit distance is tightly connected with that of a pair-wise alignment (see e.g. [15] for an introduction to these notions). A pair-wise alignment of two sequences $A, B \in \Sigma^*$ is a pair of strings $(A', B')$ where:

- $A', B' \in (\Sigma \cup \{'-\'}\)^*$, where $'-'$ is a special gap symbol;
- $A', B'$ have the same length $L$;
- each $A'$ and $B'$ contains exactly $L - |A|$ and $L - |B|$ gap symbols, respectively.

Thus, $A$ and $B$ are subsequences of $A'$ and $B'$, respectively, in the sense that they can be obtained from them by deleting zero or more symbols. A pair $(A'[i], B'[i])$ is called

- **identity**, if $A'[i], B'[i] \in \Sigma$ and $A'[i] = B'[i]$;
- **substitution**, if $A'[i], B'[i] \in \Sigma$ and $A'[i] \neq B'[i]$;
- **deletion**, if $A'[i] \in \Sigma$ and $B'[i] = '-'$;
- **insertion**, if $B'[i] \in \Sigma$ and $A'[i] = '-'$.

An insertion or deletion is also called a gap. The set of all pair-wise alignments of $A$ and $B$ is denoted by $A(A, B)$. The edit distance of $A$ and $B$ can also be expressed in terms of alignments, as

$$ed(A, B) = \min_{(A', B') \in A(A, B)} \sum_{i \in \{1, \ldots, |A'|\}} s(A'[i], B'[i]).$$

An optimal alignment of $A$ and $B$ is an alignment of maximum global alignment score. With the scoring scheme $s(\varepsilon, \varepsilon) = s(c, c) = -1, s(a, b) = -1$ for all $a \neq b \in \Sigma$, and $s(a, a) = 0$, for all $a \in \Sigma$, finding the optimal global alignment score of $A$ and $B$ is equivalent to computing their edit distance. Unless otherwise stated, in the rest of this paper we assume that an “optimal alignment” refers to this scoring scheme for edit distance.

In Section 3, we prove that the MIN-ED-2PC-$\Sigma$ problem (and hence the MIN-ED-2PC-$\Sigma^*$ problem) is NP-hard in all of the above variants. Remarkably, these negative results hold also in the case of a binary alphabet $\Sigma := \{0, 1\}$. The instances resulting from the reduction can also be cast as inputs to the Recombination-Oblivious Diplod Alignment Problem (see Section 4); the two problems are polynomially equivalent on these instances and this proves that Recombination-Oblivious Diplod Alignment Problem is also NP-hard.

2.3 Further notations for strings and graphs

A string $S$ over $\Sigma$ of length $n$ can be expressed as a $\Sigma$-DAG $\overrightarrow{S}$ with $n$ nodes and of width 1 consisting of a single path $P = v_1, v_2, \ldots, v_n$, with $\ell(v_i) = S[i]$ (equivalently, $sp(P) = S$).

Let $D_1 = (D_1, \ell_1)$ be a $\Sigma^*$-DAG with a single sink $t_1$ and $D_2 = (D_2, \ell_2)$ be a $\Sigma^*$-DAG with a single source $s_2$. The $\Sigma^*$-DAG obtained by adding the arc $(t_1, s_2)$ to the disjoint union of $D_1$ and $D_2$ is denoted by $D_1 \cup D_2$, juxtaposing the aliases, just as with strings, to suggest the concatenation in series of the actual objects.

Given a $\Sigma^*$-DAG $D$, we denote by $D_\square$ the $\Sigma_\varepsilon$-DAG obtained from $D$ by expanding nodes labeled by strings of length more than 1 into paths. Namely, each node $v$ labelled by $S$, $|S| \geq 2$ is replaced by the path $\overrightarrow{S}$; the arcs incident to $v$ get updated as follows: the arcs entering (exit, resp.) $v$ now enter (exit, resp.) the first (the last, resp.) node of $\overrightarrow{S}$.

Analogously, given a $\Sigma^*$-DAG $D$, we denote by $D_\square$ the $\Sigma^*$-DAG obtained from $D$ in the same manner as above.

3 NP-HARDNESS OF PC-MIN-ED-$\Sigma^*$

In this section, the NP-hardness of PC-MIN-ED-$\Sigma^*$ is shown for the case in which the empty string can occur as a label for some of the nodes, i.e., the labeling function is not total on $V$. 
Let \( \mathbb{N}_n := \{0, 1, \ldots, n - 1\} \). For brevity, we also denote the binary \( \text{mod} \) operation by \( \% \). The reduction, which we will describe in Section 3.1, is from the following problem:

**Longest Common Subsequence (LCS)**

**INPUT:** \( n \) strings \( S_0, \ldots, S_{n-1} \).

**OUTPUT:** A longest possible string \( S \) that is a subsequence of every \( S_i, i \in \mathbb{N}_n \).

LCS is known to be NP-hard even when the input strings are all binary and of the same length \( \ell \) [14]. Moreover, we can assume that each \( S_i \) contains both a 0 and a 1. Given \( n \) input strings \( S_0, \ldots, S_{n-1} \) of the same length \( \ell \) to the LCS problem, we show how to construct two \( \Sigma^* \)-DAGs \( A \) and \( B \) of width 2 such that the following two lemmas hold.

**Lemma 1.** Let \( S' \) be a common subsequence for \( S_0, \ldots, S_{n-1} \), and let \( \delta = \ell - |S'| \). Then there exists a path cover \( \{A_r, A_g\} \) of \( A_c \), and a path cover \( \{B_r, B_g\} \) of \( B_c \), such that \( \text{ed}(\text{sp}(A_r), \text{sp}(B_r)) = 0 \) and \( \text{ed}(\text{sp}(A_g), \text{sp}(B_g)) = 2\delta \). Hence, \( \text{ed}(\text{sp}(A_r), \text{sp}(B_r)) + \text{ed}(\text{sp}(A_g), \text{sp}(B_g)) = 2\delta \).

**Lemma 2.** Let \( \{A_r, A_g\} \) be a path cover for \( A_c \), and let \( \{B_r, B_g\} \) be a path cover for \( B_c \). Let \( d := \text{ed}(\text{sp}(A_r), \text{sp}(B_r)) + \text{ed}(\text{sp}(A_g), \text{sp}(B_g)) \). Then there exists a common subsequence \( S'' \) for \( S_0, \ldots, S_{n-1} \) with \( d \geq 2(\ell - |S'|) \).

As the reader will check, the construction can be easily performed in polynomial time. As a consequence, the above two lemmas (whose formal proofs will be given later, after describing the construction) will prove the NP-hardness of PC-MIN-ED-\( \Sigma_c \).

### 3.1 The reduction, and the general idea behind it

Let \( S_0, \ldots, S_{n-1} \) be \( n \) binary strings of the same length \( \ell \), each having both a 0 and a 1.

In the reduction, we will use an integer \( M \) that will play the role of a sufficiently big constant. A string \( T \) whose length depends on \( M \) will play the role of a firm tab gadget, capable of forcing an optimal alignment to align the \( i \)-th occurrence of \( T \) in one string to the \( i \)-th occurrence of \( T \) in the other string. We now explain how to choose \( T \).

A *linear de Bruijn sequence* of order \( k \) over a binary alphabet is a string in which every binary string of length \( k \) appears as substring exactly once [5], [20]. Let \( DB(k) \) denote one such string. The string \( DB(k) \) has length \( 2^k + k - 1 \) and can be constructed in linear time by taking the spelling of an Eulerian cycle in a de Bruijn graph of order \( k - 1 \) [3], [8].

**Lemma 3.** Let \( \alpha_1, \ldots, \alpha_g \) and \( \beta_1, \ldots, \beta_h \) be strings of length at most \( M \). Let \( k \) be such that \( |DB(k)| = \Theta(qM \log qM + qM^2) \) and let \( T = DB(k) \). Then the two strings

\[
A := \alpha_1 T \alpha_2 T \ldots \alpha_{g-1} T \alpha_g
\]

\[
B := \beta_1 T \beta_2 T \ldots \beta_{h-1} T \beta_h
\]

have an optimal alignment that aligns perfectly the \( q - 1 \) occurrences of \( T \) in each string.

**Proof:** Take an optimal alignment and suppose that the \( k \)-th character of the \( i \)-th occurrence of \( T \) in \( A \) is aligned with the same \( k \)-th character of the \( j \)-th occurrence of \( T \) in \( B \). Then, it can be assumed that these occurrences of \( T \) are wholly aligned, without losing optimality (see Figure 1).

Hence, it is sufficient to rule out any optimal alignment where some occurrence of \( T \) in \( A \) has \( n \) character aligned with any other occurrence of \( T \) in \( B \). We show that such an alignment has cost \( \omega(qM) \), so it is worse than aligning only the \( q - 1 \) occurrences of \( T \), thus it is not optimal.

Suppose by contradiction that the \( i \)-th occurrence of \( T \) in \( A \) (denoted with \( T_i \)) is such that:

- for no \( 1 \leq k < |T| \) and \( 1 \leq j \leq q \), the \( k \)-th character of \( T_i \) is aligned with the \( k \)-th character the \( j \)-th occurrence of \( T \) in \( B \),
- the cost of aligning \( T_i \) with the smallest substring of \( B \) containing the aligned characters (denoted with \( B' \)) is \( o(qM) \).

Observe that \( T_i \) is aligned by identities with at least one substring \( B'' \) of \( B' \) of size

\[
|T|/o(qM) = \omega(|T|/qM) = \omega(qM \log qM/qM + qM^2/qM) = \omega(qM + M) = \omega(M + \log |T|).
\]

(See Figure 2 for the reasoning.)

This substring may include up to \( M \) characters from some \( \beta_h \), but then it includes at least \( \omega(\log |T|) = \omega(k) \) consecutive characters from an occurrence of \( T \) in \( B \), contradicting \( T \) being a de Bruijn sequence (after fixing suitable constants in the asymptotic notation).

The high-level structure of the two \( \Sigma^* \)-DAGs \( A \) and \( B \) is depicted in Figure 3. The value \( N \), which we choose to be \( 2n\ell \), plays again the role of a sufficiently big number. The strings \( T_1, T_2, \ldots, T_{N+1} \) are just identical copies of the tab gadget \( T \); their subscripts are there only to indicate their position in \( \Sigma^* \)-DAG; we will refer to this subscript as depth. Figure 4 defines the content of the \( D(i) \) gadget, for \( i \in \mathbb{N}_n \).
In our reduction, $M \leq \ell$ will suffice, and we will always apply Lemma 3 for $q = N$ or $q = N + 1$ strings. Thus the tab gadget $T$ will be of length
\[
\Theta(N\ell \log N\ell + N\ell^2) = \Theta(n\ell^2 \log n\ell^2 + n\ell^3).
\]

These two $\Sigma^*$-DAGs, and their expanded versions $A_e$ and $B_e$, can be clearly constructed in polynomial time.

We next give the proofs of the lemmas and then extend the reduction to diploid alignments.

### 3.2 Proofs of the lemmas

The proofs depict green and red paths through the gadgets, as well as their alignments. Figures 5 and 6 help to follow the constructions.

**Proof of Lemma 1:** For $i \in \mathbb{N}_n$, let $S'_i$ denote the subsequence of $S_i$ obtained by deleting the symbols selected by its subsequence $S'$ (i.e., $S'_i$ is the complement of $S'$ in $S_i$).

Assume that we underline in green the $i$-th symbol of $S'_i$ which originate from $G_i$. Notice that there exist two (disjoint) paths $R_i$ and $G_i$ forming a path cover of the $\Sigma^*$-DAG $D(i)$ and such that
\[
sp(R_i) = \psi_i[1]\psi_i[2] \cdots \psi_i[\ell] \quad \text{and} \quad sp(G_i) = S'.
\]

The reader should now check that $A_e$ is jointly covered by two disjoint paths $A_r$ and $A_g$ such that:
\[
sp(A_r) = \left( \prod_{i=1}^{N} (T sp(R_{i\%n})) \right) T,
\]
\[
sp(A_g) = S_0 \prod_{i=1}^{N} (T sp(G_{i\%n})) = S_0 \prod_{i=1}^{N} (T S')
\]
\[
= S_0 \left( \prod_{i=1}^{N-1} (T S') \right) T S'.
\]

The reader is also invited to check that $B_e$ is jointly covered by two disjoint paths $B_r$ and $B_g$ such that:
\[
sp(B_r) = \left( \prod_{i=1}^{N} (T sp(R_{i\%n})) \right) T = T \left( \prod_{i=1}^{N} (sp(R_{i\%n}) T) \right)
\]
\[
= sp(A_r),
\]
\[
sp(B_g) = \left( \prod_{i=1}^{N} (sp(G_{i\%n}) T) \right) S_1 = \left( \prod_{i=1}^{N} (S' T) \right) S_1
\]
\[
= S' \left( T \prod_{i=1}^{N-1} (S' T) \right) S_1 = S' \left( \prod_{i=1}^{N-1} (T S') \right) T S_1.
\]

Clearly, $ed(sp(A_r), sp(B_r)) = 0$. By applying Lemma 3 to the strings
\[
\alpha_1 = S_0, \alpha_2 = \cdots = \alpha_{N+1} = S', \quad \beta_1 = \cdots = \beta_N = S', \beta_{N+1} = S_1
\]
we have that any optimal alignment of $sp(A_g)$ and $sp(B_g)$ perfectly aligns the $N$ occurrences of $T$. Thus:
\[
ed(sp(A_g)_g) = ed(S_0, S') + ed(S', S_1) = \delta + \delta = 2\delta.
\]

To prove Lemma 2, we introduce one more notation related to subgraphs. For a string $S$, an $S$-subpath of a $\Sigma^*$-DAG $D$ is a $\Sigma^*$-DAG $P$ such that $P$ is a subgraph of $D$, it is a path and $sp(P) = S$.

**Proof of Lemma 2:** We assume $d < 2\ell$ since otherwise the thesis holds vacuously.

Since $A$ has two sources, namely $T_1$ and $S_0$, we have that each of $A_r$ and $A_g$ starts in precisely one of them. To simplify notation in what follows, let now $A_r$ denote that path starting in $T_1$ (and thus let $A_g$ be that path starting in $S_0$).

Notice that $A_e$ $(B_e)$ contains precisely $2N+1$ $T$-subpaths (which we also call tab subpaths), and these are displaced as follows. For $i = 1, \ldots, N$, $A_r$ $(B_r, \text{resp.})$ contains two parallel tab subpaths at depth $i$ (at depth $i + 1$, resp.) and precisely one tab subpath at depth $N + 1$ (at depth 1, resp.).

The idea here is that within $A_e$ $(B_e)$ we can reach the nodes in a tab subpath at depth $i$ from the nodes in a tab subpath at depth $i - 1$. Clearly, once a solution path of $A_e$ $(B_e)$ passes through the first and the last node of a tab subpath, it traverses it entirely, holding it as a subpath of itself.

Notice that each one of the paths $A_g$ and $A_r$ $(B_g$ and $B_r$, resp.) must necessarily traverse precisely one tab subpath from any pair of parallel tab subpaths, i.e., precisely one tab subpath of depth $i$, for $i = 1, 2, \ldots, N$ (for $i = 2, 3, \ldots, N + 1$, resp.).

Also, at least one among $A_g$ and $A_r$ $(B_g$ and $B_r$, resp.) also traverses the single tab subpath of depth $N + 1$ (of depth 1, resp.). We claim that in fact, precisely one among $A_g$ and $A_r$ $(B_g$ and $B_r$, resp.) also traverses the single tab subpath of depth $N + 1$ (of depth 1, resp.).

Indeed, assume for a contradiction that $A_r$ ends immediately before the tab subpath at depth $N + 1$. This implies that $A_g$ ends with $T_{N+1}$. We claim that in this case we have $d \geq 2\ell$, contradicting the assumption made at the beginning of this proof. First, note that $A_g$ has $N + 1$ of tab subpaths. If $B_g$ had a different number of tab subpaths (i.e., $N$), then $ed(A_g, B_g) \geq |T|$. From the choice of $T$, we have that $T \geq 2\ell$ and thus $d \geq ed(A_g, B_g) \geq 2\ell$, which is the desired contradiction.

We now have that $sp(A_g)$ has $S_0$ as prefix and contains $N + 1$ tab subpaths, and $B_g$ has $S_1$ as suffix and contains $N + 1$ tab subpaths. By Lemma 3 we have that the $N + 1$ occurrences of the tab subpath are perfectly aligned, and thus $d \geq ed(A_g, B_g) \geq |S_0| + |S_1| = 2\ell$, again the desired contradiction.

By a symmetric argument we obtain that also precisely one among $B_g$ and $B_r$ starts with the tab subpath of depth 1.

At this point, we summarize the situation as follows:
1) the tab subpaths of $A_g$ are precisely $N$, namely those at depth $1, 2, \ldots, N$;
2) the tab subpaths of $A_r$ are precisely $N + 1$, namely those at depth $1, 2, \ldots, N, N + 1$;

The existence of this identity zone allows us to define a common subsequence $S'$ to $S_0, \ldots, S_{n-1}$. Namely, the identities restricted to the content of $S_0$ and $S_1$ picked by an optimal covering alignment of $D(0)$ and $D(1)$ inside $A_g$ and $B_g$ fixes a common subsequence $S' = S_1[i_1]S_1[i_2] \cdots S_1[i_{p_0}]$ of $S_0$ and $S_1$, and we need to show that this subsequence is common to all $S_0, \ldots, S_{n-1}$.

Since $A_r$ must be picking in $D(0)$ a complementary subsequence $S'_0 = S_0[j_1]S_0[j_2] \cdots S_0[j_{p_0}]$ of $S_0$, where $i_{k_1} \neq j_{k_2}$ for all $k_1$ and $k_2$, to guarantee $S_0$ is covered by $S' \cup S'_0$, then for an identity alignment, $B_r$ must be picking in $D(0)$ a subsequence $S''$ of $S_0$ matching perfectly with $S'_0$. If one removes two identical subsequences $S'$ and $S'_0$ from the same string ($S_0$), the resulting string is the same. Hence one can modify $B_r$ to pick $S'_0$ instead of $S'$ without changing the alignment score. The analogous modification of $B_g$ inside $D(0)$ to pick $S'$ within $D(0)$ also does not change the score. One can continue propagating these modifications to the left and, analogously, to the right until one has proven $S'$ to be a subsequence of all $S_0, \ldots, S_{n-1}$.

Since the identity region contains all different types of subDAG pairs, one can obtain an alignment with cost $d = 2(\ell - |S'|)$ as follows. Copy the zero cost identity alignments to all places; with the same propagation argument as above, one observes that $S_0$ is aligned against $S'$ (being the prefix of $B_g$ before the first tab) and $S_1$ is aligned against $S'$ (being the suffix of $A_g$ after the last tab); all other parts of the alignments have cost zero. Since $S'$ is a subsequence of $S_0$ and $S_1$, the optimal edits to make them match cost exactly $2(\ell - |S'|)$. On the other hand, there cannot be any better alignments: Each edit located between the identity zone and before $S_0$ is propagated as an extra symbol or missing symbol from $S'$ to the prefix of $B_g$ matched against $S_0$. In the former case, the extra symbols may improve the alignment of $S_0$ to the prefix of $B_g$, but these improvements cancel out with the cost of introducing these edits in the first place. In the latter case, the missing symbols just increase the cost. The case of edits between identity zone and $S_1$ is analogous.

As a consequence of the above two lemmas, we obtain the claimed result.

**Theorem 4.** Problem PC-MIN-ED-$\Sigma_e$ is NP-hard on a binary alphabet.
of a pair-wise alignment (which positions a recombination is possible. A of their haploid sequences, that is, telling in synchronization itself be expressed as a pair-wise alignment that stores the forming a chromosome pair. Such a diploid chromosome haplotype sequences of each of the homologous sequences can be less common [11], [16], [17]. The motivation to model (proteins) [6], [10], but extensions to diploid organisms have been less common [11], [16], [17]. The motivation to model

for some $i$. We also overload the notation $sp(\cdot)$, and denote by $sp(A')$ the initial string $A$, that is, $sp(A')$ is an operation removing the gap symbols ‘$-$’ from $A’$. We obtain the following problem.

**Recombination-Oblivious Diploid Alignment Problem**

**INPUT**: Alignments $(A', B')$ and $(C', D')$ of strings $A$ and $B$, and $C$ and $D$, respectively.  
**OUTPUT**: Alignments $(A''', B''')$ and $(C''', D''')$ resulting from a series of recombinations to $(A', B')$ and $(C', D')$, respectively, maximizing

$$as(sp(A'''), sp(C''')) + as(sp(B'''), sp(D'''))$$

Notice that even if $(A', B')$ and $(C', D')$ represent diploid chromosomes of two siblings, their recombination patterns are independent, so the formulation gives a way to measure just the sequence similarity not penalizing on the natural recombination phenomenon. Other applications are in comparing haplotyping results between two tools even on the same data; haplotyping algorithms usually return blocks of correctly phased variants, but not on full chromosomes.

The Recombination-Oblivious Diploid Alignment Problem was defined in [16], but its complexity was left open. Related notions on comparing two sequences to a third with edit distance and crossover were studied in [11]; polynomial dynamic programming algorithms were derived, and extensions to multiple sequences were shown NP-hard. These notions and dynamic programming algorithms were further generalized in [16], [17]. The case where the third sequence is an alignment, and one needs find a recombination of it to minimize the sum of edit distances of the resulting haplotypes to the two other input sequences, is still polynomial time solvable [16], [17]. Moreover, these algorithms extend for the case where all three inputs are alignments, but only one of them needs to be covered by the paths through the two other alignments [16], [17]. Complexity changes when one needs to cover more than one alignment: We have already seen the analogous result on labelled DAGs, but in the following we state this same result in the refined model of diploid alignments, which is slightly more specific.

4 **Recombination-Oblivious Diploid Alignment**

Pair-wise sequence alignments have been extended to capture many biological sequence features, such as mutation biases, repeats (DNA), splicing (RNA), and alternative codons (proteins) [6], [10], but extensions to diploid organisms have been less common [11], [16], [17]. The motivation to model diploid alignment comes from the recent developments in sequencing and in haplotyping algorithms; it can be foreseen that one day we will have reasonably accurate haplotype sequences of each of the homologous sequences forming a chromosome pair. Such a diploid chromosome can itself be expressed as a pair-wise alignment that stores the synchronization of their haploid sequences, that is, telling in which positions a recombination is possible. A recombination of a pair-wise alignment $(A'[1..L], B'[1..L])$ of strings $A$ and $B$ is

$$(A'[1..i]B'[i + 1..L], B'[1..i]A'[i + 1..L]),$$

for some $i$. We also overload the notation $sp(\cdot)$, and denote by $sp(A')$ the initial string $A$, that is, $sp(A')$ is an operation removing the gap symbols ‘$-$’ from $A’$. We obtain the following problem.

**Recombination-Oblivious Diploid Alignment Problem**

**INPUT**: Alignments $(A', B')$ and $(C', D')$ of strings $A$ and $B$, and $C$ and $D$, respectively.  
**OUTPUT**: Alignments $(A''', B''')$ and $(C''', D''')$ resulting from a series of recombinations to $(A', B')$ and $(C', D')$, respectively, maximizing

$$as(sp(A'''), sp(C''')) + as(sp(B'''), sp(D'''))$$

Notice that even if $(A', B')$ and $(C', D')$ represent diploid chromosomes of two siblings, their recombination patterns are independent, so the formulation gives a way to measure just the sequence similarity not penalizing on the natural recombination phenomenon. Other applications are in comparing haplotyping results between two tools even on the same data; haplotyping algorithms usually return blocks of correctly phased variants, but not on full chromosomes.

The Recombination-Oblivious Diploid Alignment Problem was defined in [16], but its complexity was left open. Related notions on comparing two sequences to a third with edit distance and crossover were studied in [11]; polynomial dynamic programming algorithms were derived, and extensions to multiple sequences were shown NP-hard. These notions and dynamic programming algorithms were further generalized in [16], [17]. The case where the third sequence is an alignment, and one needs find a recombination of it to minimize the sum of edit distances of the resulting haplotypes to the two other input sequences, is still polynomial time solvable [16], [17]. Moreover, these algorithms extend for the case where all three inputs are alignments, but only one of them needs to be covered by the paths through the two other alignments [16], [17]. Complexity changes when one needs to cover more than one alignment: We have already seen the analogous result on labelled DAGs, but in the following we state this same result in the refined model of diploid alignments, which is slightly more specific.
Theorem 5. The Recombination-Oblivious Diploid Alignment Problem is NP-hard when alphabet size is 3 or larger.

Proof. We use alphabet $\Sigma = \{0, 1, t\}$ and fix the scoring scheme $s(r, c)$ as follows:

<table>
<thead>
<tr>
<th>$s$</th>
<th>0</th>
<th>1</th>
<th>$t$</th>
<th>$\ell$</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0</td>
<td>-1</td>
<td>$\infty$</td>
<td>-1</td>
</tr>
<tr>
<td>1</td>
<td>-1</td>
<td>0</td>
<td>$\infty$</td>
<td>-1</td>
</tr>
<tr>
<td>$t$</td>
<td>$\infty$</td>
<td>$\infty$</td>
<td>0</td>
<td>$\infty$</td>
</tr>
<tr>
<td>$\ell$</td>
<td>-1</td>
<td>-1</td>
<td>$\infty$</td>
<td>0</td>
</tr>
</tbody>
</table>

Here $s(r, c)$ is given by the value at row $r$ and column $c$.

The DAGs $A$ and $B$ from Section 3 can be cast as pairwise alignments by taking each column of the gadgets (as in the visualization) and considering the following cases:

(i) if a column contains two nodes $v$ and $w$ with the same label $T = \ell(v) = \ell(w)$, construct a block $(t, t)$ in the alignment;

(ii) if a column contains two nodes $v$ and $w$ with one of them, say $w$, with label $\ell(w) = \varepsilon$ construct a block $(\ell(v), \ell(v))$ in the alignment;

(iii) if a column contains only one node $v$ labeled $\ell(v) = S_0$ or $\ell(v) = S_1$, construct a block $(\ell(v), \ell(v))$ in the alignment;

(iv) if a column contains only one node $v$ labeled $\ell(v) = T$, construct a block $(\ell(v), \ell(v))$ in the alignment.

Concatenating these blocks from left to right creates pairwise alignments $(A', B')$ and $(C', D')$ corresponding to DAGs $A$ and $B$, respectively. The resulting pairwise alignment $(A', B')$ is shown in Figure 7.

![Fig. 7. High-level structure of pair-wise alignment (A', B'). The contents of blocks D', are shown in Figure 8. All the t_i corresponds to the symbol t; the subscripts are to show the relationship with the graph A.](image)

Consider a series of recombinations of $(A', B')$ into $(A'', B'')$ and a series of recombinations of $(C', D')$ into $(C'', D'')$, that maximize

$$\text{as}(sp(A''), sp(C'')) + \text{as}(sp(B''), sp(D''))$$

under the scoring function defined above. We claim that

$$-(\text{as}(sp(A''), sp(C'')) + \text{as}(sp(B''), sp(D''))) - 2\ell$$

equals the optimal solution of covering alignment of DAGs $A$ and $B$ with the unit cost edit distance. For the reverse implication, one can map the alignments of red and green paths in the proof of Lemma 1 to form alignments of $(sp(A''), sp(C''))$ and $(sp(B''), sp(D''))$, where $S_0$ and $S_1$ are deleted from the head and tail, respectively, of the alignment corresponding to red paths. Alignment corresponding to that of green paths is identical, with respect to the mapping of nodes to symbols derived above. The claimed equality then follows considering the definition of the scores. For the forward implication, since all tab symbols $t$ need to align in their occurrence order as in the proof of Lemma 2, and since recombinations inside the head $(S_0, S_1)$ and tail $(S_1, S_1)$ of $(A', B')$ and $(C', D')$, respectively, are non-effective, an optimal series of recombinations is in one-to-one correspondence with the covering red and green paths as in the reverse implication.

Hence, solving Recombination-Oblivious Diploid Alignment Problem on these instances solves the PC-MIN-ED-$\Sigma$' on $\Sigma$-DAGs and due to Lemmas 1 and 2 would solve the LCS problem. □

5 NP-HARDNESS WITHOUT EMPTY LABELS

Recall that problem PC-MIN-ED-$\Sigma$ differs from PC-MIN-ED-$\Sigma_4$ in that each node of the graph needs to have a non-empty label. Our plan is to modify as little as possible the construction offered in Section 3 and for this purpose we consider indel edit distance, rather than unit cost edit distance: In this scoring scheme, substitutions have cost $\infty$, identities have cost zero, and insertions and deletions have cost 1. We also increase the alphabet from binary to size 4 by replacing all empty labels in $A$ with a new symbol $a$, and all empty labels in $B$ with a new symbol $b$. Obviously, any pair of covering alignments needs to have gap symbols aligned with each $a$ and each $b$. This cost is invariant and does not alter the relative order of alignments when sorted by their overall cost. One can thus modify systematically proofs of Lemmas 1 and 2 taking this invariant into account to conclude that PC-MIN-ED-$\Sigma$ is NP-hard with this scoring scheme and alphabet size 4. An analogous modification to the proof of Theorem 5 gives that Recombination-Oblivious Diploid Alignment Problem is NP-hard when its input contains no gap symbols and the alphabet size is 5. The Appendix demonstrates a subtle phase transition for this latter problem variant, as a slight relaxation of it is in P. For the interested reader, the last paragraph of the Appendix revisits the reduction to show that the derived partially covering relaxation indeed achieves better edit distance for the reduction instances than the NP-hard solution; such pair of alignments has quite a specific structure and gives also more insight to the reduction itself.

6 DISCUSSION

It is evident that the reductions given here generalize to scoring functions beyond those considered here. We leave such development for future work. Notice that similar fine-grained complexity analysis has been conducted for the LCS problem [4].

The reduction technique developed here is likely to find other applications in the area of computational pan-genomics [18]. A direct consequence is that comparing two pan-genome representations is NP-hard, if accepting the notion of covering alignment developed here as the basis. Namely, the general optimization problem is to find
minimum number \( k \) of paths to cover each of the two input DAGs, and then among such covers one that maximizes the sum of \( k \) alignment scores among the \( k \) pairings. Since case \( k = 2 \) is NP-hard, case \( k = 1 \) is the classical quadratic time solvable sequence alignment problem, and our reduction works on binary alphabet, we have identified a phase transition for this notion of similarity (see also the Appendix for an even closer phase transition). As the labeled DAG representation loses the connectivity information on variations in the pan-genomic setting, one could resort back to a multiple alignment of haplotypes, and adjust the notion of recombinations to allow only limited number of those. This notion allows parameterized complexity analysis. Indeed, let us consider the Recombination-Oblivious Diploid Alignment Problem from this angle. Given a limit \( r \) for the number of recombinations in one alignment, a naive algorithm is to consider all \( \binom{n}{r} \) \( r \) recombinations on both input alignments and then compute the global alignment of the resulting haplotype pairs. This results into an \( O(n^{2r+2}) \) time algorithm. One can speed this up to \( O(n^{r+3}) \) by considering all recombinations only in one input alignment and then resorting to the algorithm in [16]. We believe there is room for further work around the parameterized tractability border of this problem. For the general covering alignment problem on DAGs a plausible direction is to look for approximation algorithms or approximation hardness.

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