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A New Algorithm for the Reconstruction of Near-Perfect Binary Phylogenetic Trees

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Abstract

In this paper, we consider the problem of reconstructing near-perfect phylogenetic trees using binary characters. A perfect phylogeny assumes that every character mutates at most once in the evolutionary tree. The algorithm for reconstructing a perfect phylogeny for binary characters is computationally efficient but impractical in most real settings. A near-perfect phylogeny relaxes this assumption by allowing characters to mutate a constant number of times. We show that if the number of additional mutations required by the near-perfect phylogeny is bounded by q, then we can reconstruct the optimal near-perfect phylogenetic tree in time $2^{\circ}({}^{q}{}^{2}nm^{2})$ where n is the number of taxa and m is the number of characters. This is a significant improvement over the previous best result of $nm^{\circ}N2^{\circ Aq^{2r^{2}A}}$ where r is the number of states per character (2 for binary). This improvement could lead to the first practical phylogenetic tree reconstruction algorithm that is both computationally feasible and biologically meaningful. We finally outline a method to improve the bound to $q = {}^{\circ n}nm^{2}$.

1 Introduction

One of the classical problems of computational biology is to reconstruct an evolutionary tree for a set of taxa based on character data. Parsimony is one of the widely used metrics to solve the problem. It has been known to be particularly useful in the case when the evolutionary tree reconstructed is over a short period of time.

We borrow some of the following definitions and notations from [5]. The input to the problem is generally represented by a matrix / where rows R are strings of states corresponding to taxa. The columns $C = \{1, \bullet \bullet, m\}$ are referred to as characters. The set of states corresponding to any character c is denoted by A_c , therefore every taxon $s \ e \ A \setminus x \bullet \bullet x \ Am$. In a phylogenetic tree, each vertex v corresponds to a taxon and has an associated label $l(v) \ e \ A \setminus x \bullet \bullet \bullet x \ Am$. We use the terms phylogeny, phylogenetic tree or just tree interchangeably.

Definition 1: A *phylogeny* for a set of n taxa R is a tree T(V, E) with the following properties:

- 1. if a taxon seR then s E l(V(T))
- 2. for all (rz, v) E E(T), H(l(u), l(v)) = 1 where H is the hamming distance

Definition 2: The *length of a phylogeny* T, *length*(T) = |E(T)|. **Definition** 3: The penalty of a phylogeny *T* is defined as

$$penalty{T) = length(T) - \sum_{cec} (|A_c| - 1)$$

Minimizing the length of a phylogeny is the problem of finding the most parsimonious tree, a well known NP-complete problem [6]. A phylogenetic tree *T* constructed on input /, is called a perfect phylogeny if *penalty*(*T*) = 0. As summarized by [5], reconstructing a perfect phylogeny was proved to be NP-haid independently by Bodlaender et al. [2] and Steel [10]. This led researchers to work on either sophisticated heuristics (for e.g [7], [3]) or solve optimally for fixed parameter versions of the problem (for e.g, [1], [9]). Gusfield considered an important special case of the perfect phylogeny moblem, when the number of states is bounded by 2. We call such a tree as a binary perfect phylogeny. Gusfield showed that binary perfect phylogenies can be reconstructed in linear time [8]. Lagergren and Fernandez-Baca, considered the problem of reconstructing near-perfect phylogenies [5]. The assumption of a 'near'-perfect phylogeny is that *penalty*(*T*) is small for the most parsimonious tree. Their algorithm runs in time $nm^{\circ}2^{\circ Aq^{2r^2}}$ where r is the number of states per character, q is the penalty, n is the number of taxa and m is the number of characters.

Our Results: In our work, we consider an important special case of the problem when $r = |A_c| = 2$. The case when r = 2 is primarily important because Single Nucleotide Polymorphisms (SNPs) are bi-allelic. We can therefore use the algorithm for reconstructing trees where the taxa are DNA sequences and the characters are SNP markers. We show that if the penalty of the most parsimonious phylogeny is bounded by q, then we can reconstruct the phylogenetic tree in time $2^{\circ n}nm^2$. In section 4 we briefly describe how this time-bound can be improved to $q^{\circ n}nm^2$. More details on the improved algorithm will be available in [4]. Our algorithm is almost entirely self-contained and its understanding requires only some fundamental theorems on phylogenetic trees. Although some existential proofs are hard, the algorithm itself is not very complicated to implement. We also expect the algorithm to perform significantly better in practice than the worst case bounds.

 create a conflict graph G s.t there exists a bijection c : V(G) *-+ C and (tx, v) E E(G) iff \(\begin{aligned} \langle G_{c(uMv)} \rangle =4 \end{aligned} 4 \end{aligned} 2 \end{aligned} g s.t Q is a set of columns corresponding to the vertex cover of G, return nil if none exists
(a) for all trees T s.t there exists an onto function /i: E(T) >>> Q
i. for each vertex v eV(T)
A. find the subset of rows b(v) 'compatible' to v
B. build a perfect phylogeny T_v for the set b(v)
C. replace v with tree T_v
D. complete the tree by linking all the T_v's
ii. if cost(T) < min thenT_f = T,min = cost(T_f)

Figure 1: Overview of the algorithm

2 Overview of the Algorithm

We first sketch the outline of the algorithm and then explain the details of each step in subsequent sections. Specifically, given an input / and a penalty q the algorithm finds most parsimonious phylogeny with penalty at most q or *nil* if none exists. The following definitions are useful:

Definition 1 The set of gametes Gij restricted to two characters i,j is defined as: Gij = {(fc,i)|3r E R, r[i] = k, r[j] = l}.

Definition 2 Two columns i, $j \in C$ are said to contain (all) four gametes when |Gij| = 4

A high level pseudo-code of the algorithm is given in 1. The following sections elaborate on each of the steps of this pseudo-code which we will refer to as the main pseudo-code.

Pre-processing It is well known that the most parsimonious phylogeny reconstructed is imperfect if and only if the input / contains the four-gamete property. Before running our algorithm, we pre-process the input as follows. We remove all the columns that that have only single state (i.e. they do not mutate at all). We then look for the pairs of input columns i, j for which $\langle Gij \rangle = 2$. In such a case, both the columns contain the same information. We remove one of the two columns from the input. We show in the Appendix (Theorem 12) that this preprocessing steps does not alter the overall running time or the correctness.

At the end of pre-processing, we have the input / that satisfies $|Gij| \ge 3$ for all columns i, j. **Conflict graph G and the vertex cover-columns** Q: A conflict graph G corresponding to input / is constructed as follows. Each vertex $v \in V(G)$ of the graph represents a character $c(v) \in /$. An edge (u, v)is added if and only if all four gametes are present in c(u) and c(v). The function /i : $E(T) *-\pm C$ maps mutation events to the character that mutates. Note that in any phylogeny if ji(u, v) = c then $l(u)[c] \land Kv)[c]$. . *

To find the vertex cover, we use the classical 2-approximation algorithm. We set Q to be the set of columns corresponding to the vertex cover returned by the algorithm. We assert that there exists no g-near perfect phylogeny if $\langle Q \rangle > 2q$.

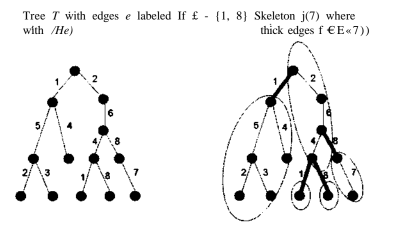


Figure 2: A phylogenetic tree and its associated skeleton. The bit vectors of the species are left out for simplicity.

Lemma 1 There exists no q-near-perfect phytogeny if |Q| > 2q.

Proof: First notice that for any pair of characters (i, j) s.t $\langle Gij \rangle = 4$ the tree *T* should contain either two edges e, e' such that /x(e) = /i(e') = i or /i(e) = ji(e') = j. If $\langle Q \rangle > 2q$ then there exists no vertex cover of size *q*. This implies that there exists at least one edge $(u, v) \in E\{G\}$ s.t neither c(u) nor c(v) mutates multiple times in T, a contradiction.

For the step 2a of the pseudo-code, we enumerate by brute force all possible trees T that mutate the columns in Q at least once.

Definition 3 A vertex v is bad w.r.t a pair of characters (i, j) and a set of columns Q, if $i, j \notin Q$ and $(l(v)[i], l(v)[j]) \notin G_{i,j}$.

From now on we adhere to the following naming convention. The tree T (used in step 2a that mutates the columns in the vertex cover Q is referred to as the **skeleton tree**. The vertices of the a skeleton tree T are referred to as **super nodes**, since an entire tree replaces each of these nodes in a final phylogenetic tree. Figure 2 shows an example of a phylogenetic tree along with its skeleton.

3 Details

3.1 Finding a compatible set for all vertices

This is the most complicated step of the algorithm. We want to partition the rows of the input matrix such that each super node v is assigned a subset of 'compatible' rows b(v). By arbitrarily rooting the tree T, the structure of the tree defines the bits for the characters in Q. We can think of every super node v in the tree being *tagged* with the states &i, &2>' " ? $b \setminus Q \setminus$ and use the notation t(v) to denote the tag. Note that t(v) is the same as l(v) restricted to the columns of Q. We can therefore partition the set of rows if the tags t(v) are all unique. The problem arises when two super nodes say $v \setminus V2$ are both assigned the same tags - and therefore

the same subset of rows S. We need to further partition S into two sets S and 52 one for each v and V2-We need several lemmas to determine how to make this partition.

Definition 4 We use the notation $V_{VliV2}(T)$ to be the path between (and including) vertices (or super nodes) $v \setminus and V2$ of any tree T.

Note that if $v \downarrow V2$ are super nodes and *T* is the phylogeny (not skeleton), then the notation $V_{Vb V2}(T)$ is used to denote the path in the tree T that connects super nodes $v \lor$ and V2 of the corresponding skeleton s(T).

The proof for the following theorem is given in Appendix 4.

Theorem 2 If there exists a q-near-perfect phylogeny that has bad vertices (w.r.t Q) then there exists a q-near-perfect phylogeny that does not contain any bad vertices (w.r.t Q).

For the rest of the paper, we fix an optimal phylogeny T^t that does not contain any bad vertices. We will impose additional restrictions on T[^] later.

Corollary 3 In Top_{tt} if there exists two edges $e = (v \setminus 2)$ and $d = (3, v \pm)$ such that $fi(e) = n(e^{f})$ then for all columns $j \notin Q$, $\langle e \in 7 \setminus 3, v_3(\text{Topt}) | ji(e) = j \rangle$ is even and therefore $l\{vi\}[i] = l(v_A)[i]$ for all $i \notin Q$.

For any phylogeny T, the skeleton of T, s(T), with respect to a set of columns Q (vertex-cover) is the tree of super-nodes where each super node is attached to edges $e \in T$ s.t /i(e) G Q. A super node can be viewed as a set of vertices of the final phylogenetic tree and therefore gives us the following obvious properties of containment. For a super node $v \mid E s(T)$, we say that edge $e = (u, v) G v \mid \text{iff } u \in v \mid_{g} v G v \mid$ and (tx, v) G E(s(T)). Similarly, we say that column $j G v \mid \text{iff } 3e G v \mid \text{st. } n(e) = j$.

Definition 5 Two super nodes $v \setminus and V2$ of s(T) are tag-adjacent if

- $L t(v \mid) = t(v2)$ and
- 2. there exists no super node V3 G V_{VliV2} {s(T)) such that $t(v^{\wedge}) = t(v)$

Definition 6 Two super nodes $v \mid and V2$ of s(T) are paired-tag-adjacent if there exists no super node v\$ G $V_{V_{u}V2}(s(T))$ such that $t(v_3) = t(v_i)$ ort $\{v_3\} = t\{v_2\}$

Definition 7 An edge e G E(T) or column /i(e) is distinguishing w.r.t a phylogeny T if for any two tagadjacent super nodes $v \mid and V2$, $\{e^f \in V_{VliV2}(T) \mid fi(e') = /i(e)\}|$ is odd.

It is easy to see that for the optimal near-perfect phylogeny T[^] there exists a distinguishing edge for every pair of tag-adjacent super nodes.

3.1.1 Partition

We can define equivalence classes $V_{t(v)}$ containing super nodes based on the equality of tags t(v) of super nodes. It is straightforward to partition the rows *R* for each equivalence class of super nodes. This is performed by just examining the tags of each equivalence class. The pseudo-code in Figure 3 provides the details of the steps necessary to compute the partitions. At any point in time, the partition algorithm maintains a partition $P(V^{\wedge})$ for each equivalence class. Each recursive call refines one of the partitions of one of the equivalence classes. Just to avoid excessive notations we will use $P\{V_{ti}\} = \{Si, \dots, S^{k}\}$ for all *i*. It should be obvious from the context as to which equivalence class 5, belongs to. We say that a column *i* partitions a set *Sj* if there exists two rows n, *T2* G *Sj* such that $r \setminus [i] = 0$ and r2[t] = 1. The partition can be written as $S_{jo} = \{n \in Sj \setminus ri[i] = 0\}$ and $S_h = \{r_2 \in Sj \setminus r_2[i] = 1\}$.

An overview of the approach is to first discover the distinguishing columns that partition two different equivalence classes but only mutates once in T^{+} . In the second step we guess the distinguishing columns

that partition two different equivalence classes and mutates multiple times. The columns guessed are added to the skeleton so that the skeleton expands in size.

The value mark(v) denotes our guess of whether or not the super node v contains any real vertices in T_{opt} . The value is set to 'real' if we guess that there exists at least one real vertex in v and 'steiner' otherwise. A partition $P(V_{t_i})$ is complete when the number of sets in $P(V_{t_i})$ is equal to the number of super nodes v in V_{t_i} with mark(v) = real, that is: $|P(V_{t_i})| = |\{v \in V_{t_i} | mark(v) = real\}|$.

The partition function initially finds the set D_2 consisting of columns that refines the partition in two different equivalence classes. If $|D_2| > q + 1$, then the function picks an arbitrary set of q + 1 columns and performs 2(q+1) recursive calls on partition induced by each column in the set on two different equivalence classes that the column refines. If $|D_2| \le q$, then the function guesses all the mutations of the columns in D_2 and adds them to s(T). Note that each added edge splits a single super node into two new super nodes.

At the end of the function partition, we add mutations and expand the skeleton. Before the expansion the set D_2 was empty. The following Lemma proves that it remains empty even after the skeleton expands. Moreover, it remains empty throughout the execution of function use-interface which will be described shortly.

Lemma 4 If D_2 is empty at any time, then it remains empty when expanding the skeleton by the addition of non-vertex cover columns (specifically in steps 3a, 3c of function partition described in Figure 3 and step 5 of function use-interface described in Figure 4).

Proof: For the sake of contradiction, assume that D_2 was empty and later became non-empty. Let c be a column in D_2 . Let t_1 and t_2 be the two tags that are split by c. Note that if the t_1 and t_2 are different on the columns in Q, then the column c would have been found in function partition (Figure 3) itself. Therefore it must be the case that t_1 and t_2 are identical on columns in Q. Since t_1 and t_2 are identical on columns in Q, the algorithm found a column d that distinguishes them. Now note that c and d together must form 4 gametes. This is a contradiction to the fact that neither c nor d belongs to Q.

Correctness:

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A partition $P(V_{t_i}) = \{S_1, \dots, S_k\}$ is defined to be *good*, if it can be refined by further partitions (no merges/unions) into that of T_{opt} . The set of partitions $\{P(V_{t_1}), \dots, P(V_{t_p})\}$ is good if every partition is good.

The correctness can be proved inductively by making the claim that if the argument set of partitions is good for a function call of *partition* then the arguments for at least one of the recursive calls is good. Consider any one function call to partition and assume inductively that its argument is good. We recursively perform partition in step 4(b)ii. If $|D_2| \ge q + 1$ then there exists at least one column $i \in D_2$ that mutates only once in T_{opt} . A column *i* that mutates only once induces a partition on $P(V_{t_j})$ only when two super nodes (of the same equivalence class V_{t_j}) contain *e* with $\mu(e) = i$ in the path connecting them or if *i* mutates inside a super node $v \in V_{t_j}$. We know however that *i* partitions at least two different equivalence classes $V_{t_j}, V_{t_{j'}}$. Since *i* mutates once, it can mutate inside the super node of at most one of V_{t_j} or $V_{t_{j'}}$ and lie in the path connecting two super nodes of the other equivalence class. We recurse on both the partition induced on $P(V_{t_j})$ and $P(V_{t_{j'}})$. Moreover, since the arguments of partition is good, the partition induced by using *i* on at least one of $S_j \in P(V_{t_j})$ or $S_j \in P(V_{t_{j'}})$ is good. This completes the recursive case. The other case function partition ($P(V_{t_1}), \cdots, P(V_{t_p})$, int list selectedCols)

- 1. if $\forall i, P(V_{t_i})$ is complete then
 - (a) guess the assignments of the partitions S_i to the super nodes of the skeleton
 - (b) expland the skeleton into S' st columns in selectedCols mutate once and the expanded tags of the super nodes are compatible with the rows assigned
 - (c) return S'
- 2. let D_2 be the set of columns st, $\forall i \in D_2, \exists_{>1}j, \exists k \text{ st } i \text{ partitions } S_k \in P(V_{t_i})$
- 3. if $|D_2| \le q+1$
 - (a) guess multiple mutating columns among set D_2 and add to skeleton
 - (b) guess the assignments of the partitions S_i to the super nodes of the skeleton
 - (c) expland the skeleton into S' such that columns in selectedCols mutate once and the expanded tags of the super nodes are compatible with the rows assigned
 - (d) guess mark(v) for all super nodes $v \in S'$ corresponding to the expanded skeleton
 - (e) use-interface (S')

4. else

- (a) consider any $D'_2 \subseteq D_2$ s.t $|D'_2| = q + 1$
- (b) $\forall i \in D'_2$ and for any two j s.t $\exists S_k \in P(V_{t_i})$, i partitions S_k
 - i. let $P(V_{t_i}) = (P(V_{t_i}) \setminus \{S_k\}) \cup \{S_{k_0}\} \cup \{S_{k_1}\}$
 - ii. partition ($P(V_{t_1}), \cdots, P(V_{t_p})$, selectedCols $\cup \{i\}$)

Figure 3: Algorithm to partition and assign rows of the input to super nodes

function use-interface (skeleton S)

- 1. let $D \mid$ be the set of columns s.t, V'i; $G D \mid$, 3ij, s.t *i* partitions R(Vj).
- 2. select some *Vj* that is incomplete
- 3. select a lowest pair $v \mid V_i$ G Vj of tag-adjacent super nodes, such that $mark(v \mid) = mark(v2) = real$
- 4. if 3/i G Du vs G V? and t7₄ G V^{\wedge}/, for some $t^1 \wedge t^{77}$ and $* \wedge t^7$, t" such that
 - (a) morfc(v3) = marfc(174) = real
 - (b) /i partitions J?(VV) U R(V?>)
 - (c) t7i , $v_2(5)$ and V_{V3} , $v_4(S)$ share a super-node in S
 - (d) (vi, vs) and $(v_2, V4)$ are paired-tag-adjacent then
- 5. guess extension S^{f} of skeleton 5 by mutating column h (once or multiple) times
- 6. guess mark(v) for super nodes adjacent to mutation(s) of h
- 7. use-interface (S^*)

Figure 4: Algorithm to find interface edges and extend the skeleton

is when we guess all multiple mutations among columns *i* that partition at least two different equivalence classes in step 3a. Since the size of the set of columns from which we guess is bounded by q_9 this step can clearly be performed in $O(2^q)$ time to find the columns mutating multiple times and $2^{\circ} q^{\circ}$ to place the edges in the skeleton tree. Note that we also guess the number of times a column mutates.

The step lb expands the skeleton by adding columns of selectedCols.

As before, the edges of the expanded skeleton define the tags associated with each super node. We now assign for each super node a partition S_i . While making this assignment it could be the case that the number of partitions is less than the the total number of super nodes. Specifically, two super nodes of the same tag are assigned the same partition. However, the number of distinct tags in the skeleton should be equal to the number of partitions. This step can be naively implemented by once again enumerating all skeletons using the columns of the current skeleton and selectedCols. We can then discard any skeleton that is not compatible to the set of partitions. The running time for extending the skeleton and assigning the rows can be bounded as follows. The recursion tree has height q, and any function call to *partition* makes at most 2(q + 1) recursive calls. Therefore the number of leaves of the recursion tree is bounded by $2^q [q + 1)^q$. Each leaf nock contains at most q columns in its list *selectedCols*. Since the number of trees with 2q vertices is bounded by $2^{\circ}Aq^{\circ (<q)}$ step 1b can be implemented in $2^{\circ}Aq^{\circ \wedge}$ time. This assures that the total time spent by partition function is bounded by $2^{\circ}Aq^{\circ \wedge q}$.

3.1.2 Interface

For discovering distinguishing edges, we are just left with the last case - distinguishing columns that mutate once or more but partition only one equivalence class. We define R(Vt) Q R for any equivalence class Vt to be the set of rows that match tag t. A vertex v of phylogeny T is called real if l(v) G R - informally if the label of the vertex is present in the input (equivalently, the vertex is non-steiner).

Consider the partition defined by T^{t} on any one equivalence class Vt of super nodes. We say that a

column *i* is an internal column if the partition induced by *i* on R[Vt) is not good (similar to the definition used in function partition). Informally, this implies that edge *e* with n(e) = i is present inside some super node *v e Vt* and there exists real vertices $r, r' \in v$ s.t r[i] = 0 and r'[i] = 1. We define a column that is not internal as interface. Note that an interface column induces a good partition. Informally a distinguishing interface edge is present in the path between two connected components of real vertices of the super nodes u, v G Vt s.t there exists real vertex r G ti, r[i] = 0 and real vertex r' G v, $r^{i} = 1$.

Definition 8 A pair of tag adjacent super nodes $v \mid and V2$ is defined to be a **lowest pair** in s(Topt) if there exists a vertex x such that for all super nodes $v \notin G$ siTopt) with $t(v_3) = t(vi)$, x G $V_{VliV3}(T_{apt})$ and x G $V_{V2iV3}(Topt)$. The vertex x is called the branch point of $v \mid and V2$.

T

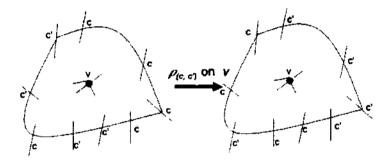


Figure 5: Application of transform p. Note that the bits of steiner vertex v changes at positions c and d after the transform.

Transform $p_c \leq J$

We now describe a very simple transform p_c^{\wedge} that operates on any tree T and is defined over two columns c, d and a steiner vertex v. To apply transform $p_{c?c'}$ on steiner vertex v, all vertices v^l such that v^l is the only real vertex in $V_{vy}(T)$ should satisfy the following property. There exists edge e G $V_{vjt'}(T)$ s.t /x(e) = c or /x(e) = c*. An example of when such a transform can be applied is shown in Figure 5. For all such paths from v to t/, the transform replaces the first mutation of c with c' and vice-versa. The result after applying the transform is shown in Figure 5. Clearly, the transform does not change the cost of the tree.

Lemma 5 In T^t, for every lowest pair super nodes vi, V2 G Vt with branch point x such that every distinguishing column of v and V2 is an interface column, there exists at least one distinguishing interface column h with the following properties:

- L column h partitions $R{Vt}$
- 2. 3vs G Vff, V4 G Vfti, $t \wedge t^7$, $if^d s.t v \setminus v$ \$ and V2, t>4 are paired-tag-adjacent and h partitions $R(Vf) \cup R(Vt'')$ and

3. the paths $V_{Vl,V2}(sC^{7})$ and $V_V \wedge_{VA}(\$(2^{\wedge}))$ share a super-node

Proof: First note that the column *h* that partitions R(Vtr)UR(V?') can only partition it into $\{R(V?), R(Vf')\}$. This is because *h* induces a partition only in one equivalence class V_t and $t \land if$, *". For the same reason $t' \land t''$. Throughout the proof of the lemma, the distance between two vertices (or edges) on a tree is simply the number of edges in the path connecting the vertices (or edges).

We know that for an optimal phylogeny, there has to exist at least one distinguishing edge h for $v \mid$ and V^2 . Let $u \mid$ and U2 be the vertices in $v \mid$ and V2 respectively, that lie on either ends of $V_{VI} \wedge_{V2}(T_{opt})$. Therefore h has to mutate odd number of times in exactly one of $V_{UI} \wedge_{I} Topt$) and $V_{Xi}u_2(Topt)$. Without loss of generality assume that it mutates an odd number of times in V_Ui , x- Consider the edge e closest to $u \mid$ such that fi(e) = h. In *Tapt* if the path from e to $u \mid$ consists entirely of degree 2 steinei* vertices, then the mutation of e can be incrementally swapped with the mutation of the neighboring edge, until the mutation of h is moved inside super node $v \mid$. Note that swapping adjacent edges of a degree 2 steiner vertex is just a trivial application of p on the vertex. After several applications of p_9 column h is no longer distinguishing between $v \mid$ and V^2 in the new optimal solution.

If the path from e to ui contains a vertex w of degree greater than 2, then every branching path from w leads to a real vertex (since steiner vertices can not be a leaf). Consider any such real vertex r, the super node in which r lies cannot have tag t since it contradicts the assumption that v_{\setminus} and V2 are a lowest pair. Now consider the case when for all such real vertices r, the path $V_{Wi}r(Topt)$ contains a mutation of h. Using transform p, the mutation of h can be swapped with neighboring edges until it is adjacent to w. Once again let e be the edge adjacent to w_9 st /x(e) = h. Let e' G $V_{Uu}x$ -, e ^ e' be another edge adjacent to w. We can now apply transform p^{Λ} on vertex w. The result of the transform is that the mutations of h and /i(e') swap positions. We can therefore keep applying transform p to move the mutation of h closer to u_{\setminus} . There are only two cases when we can not apply the transform any further. In both the cases described below, we find super node V3 that contains a real vertex.

Case 1: If the path from e to $u \mid \text{contains a real vertex r, that belongs to a super node vs say, then <math>t(vs) \wedge t(vi)$, since we assumed that $v \mid \text{and } V2$ are tag-adjacent.

Case 2: If the path from *e* to *u*\ contains a vertex *w* of degree greater than 2, that contains a branch leading to a real vertex r G V3, where $\pounds(^3) / t\{v\}$ such that the path from *w* to r does not contain a mutation of *h*. Note that $\pounds(^3) 7^{f} \pounds(^*)$ since *v*\ and *V*2 are a lowest pair.

Now consider the edge $e \ G \ _{Ul}_r(T_{op}t)$ such that p(e) = h and the distance of e from $v \mid$ is the largest. Using a series of transforms p we can move the mutation of h towards x. Once again with a similar argument we can either claim that there exists a real vertex r' in a super node $u \pm$ that either lies in the path from e to a; or is in the induced tree of w which lies in the path from e to x. If this is the case, then r and r' are two real vertices separated by an odd number of mutations of h. Also, $t(v \mid) \ \land \land(\land 3), \pounds(\#4)$. This satisfies all the properties of the lemma.

Now consider the case when the mutation of *h* can be moved so that it is adjacent to *x*. Now consider any vertex cover column *c* that mutates an odd number of times between *x* and V2. There has to exist at least one such edge since *x* can not belong to a super node of tag *t* as v and *V*2 are tag-adjacent. Consider the edge e' such that $p_i(e') = c$ and the distance from e' to *V*2 is maximum. First assume that all the vertices in $T^>u_{2,x}\{T_opt\}$ arc of degree 2 and steiner. Clearly now, using the trivial *p* transform, the mutation of *c* can be moved so that it is adjacent to *x*. We can now perform transform p_c^{\wedge} on *x* to obtain a new optimal tree where the mutations of *h* and *c* adjacent to *x* have switched places. Note that the condition of $p_{Ci}h$ requires that every path from a: to a real vertex contains either a mutation of *h* or a mutation of *c*. This can be proved as follows for the optimal tree T^. Consider any path from *x* to the first real vertex r' that is in a super node V4. If $t(v\pm) = t(v2)$ then the path from *x* to V4 should contain an odd number of mutations of *c*, since the path from *x* to V2 contains an odd number of mutations of *c*. If $\pounds(^4) \wedge tfa)$, then the path from *x* to $v\pm$ should contain an odd number of mutations of h_9 otherwise r' and r that lie in v\$ and v± respectively satisfy the properties of the Lemma. After applying p_{Cjh} , we can move the mutation of h inside v_2 since all the vertices are degree 2 the result of which is that h is no longer distinguishing between v\ and v < i.

The last case that is left to analyze is when the path from x to e' contains either real vertices or steiner vertices of degree greater than 2. If it contains a real vertex r', then r^* along with r satisfy the properties of the Lemma. If there exists a vertex w that has degree greater than 2, then the induced subtree rooted at w contains real vertices, and all paths to real vertices should contain a mutation of h (otherwise we can use the real vertex along with r to satisfy the properties of the lemma). Therefore /xistse''' $E V_{XiU2}$ wit*¹/i(e^{://}) = h_9 since otherwise we can move the mutation of h using a series of p transforms so that it is adjacent to x. This is a contradiction to optimality since we already have e adjacent to x with fi(e) - h. Therefore, we can directly apply transform p^{\uparrow}_c on vertex x. We can then move the mutation of h into u < i with a series of p transforms.

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Putting everything together, we have die fact that if for any distinguishing column h there exists no super nodes that satisfy the properties of the lemma, then the edge corresponding to the mutation of h can be moved such that it is no longer distinguishing. Furthermore, no new column becomes distinguishing in this process. We can continue this argument for every distinguishing column. If none satisfy the properties of the lemma; then eventually the first and last vertices of the path V_{VI} , v_2 (*Topt*) are identical, a contradiction to optimality.

Lemma 5 proves that for every pair of tag-adjacent super nodes, there exists at least one distinguishing interface edge that has the properties that the pseudo-code uses to identify them in step 4 of function *use-interface*. However the converse that only distinguishing interface edges have the property is not true. We will however prove that rally internal edges that mutate multiple times could satisfy the properties. Intuitively, this proves that not too many internal columns are falsely identified as interface columns. Note that at step 4 of the pseudo-code column *h* partitions *V*% and also partitions *R(Vf)* U *R(Vt'')*. If *h* is an internal column, then there exists internal edge e'' in some super node $v \in Vt$ s.t /i(e'') = *h*. Super node $v \notin V_{VZ,VA}$ because we assumed that $(v \setminus v \$)$ and $(^2,^4)$ are paired-tag adjacent. Assume that *h* lies in $V_{U3,U4}$ in *Topt* where u₃ G V? and n₄ G Vf. Note that for *h* to partition *R(Vf)* U *R(Vt*)* it has to be the case that both 1*3 and U4 contain at least one real vertex. Note that one of 1x3 or U4 could be v \$ or $v \pm$ respectively.

Consider the two paths $V_{v} V_{UZ}$ and $7_{4jl}*_4$ (shown in Figure 6). If *h* mutates just once, then it should be the case that the mutation ofh_9 edge e", should lie in one of the two paths. This is however a contradiction since *h* now partitions two equivalence classes *Vt* and one of *Vt** or *Vf* (since V3, U3, V4 and U4 each contain at least one real vertex in *Topt*). Therefore, it is not possible to add an internal column to the set *I* unless it mutates more than once in T[^].

At each recursive call, we either find a column that mutates only once thereby partitioning rows into two super nodes or discover multiple mutant edges. Therefore the depth of the recursion is bounded by O(q). Since the degree of any node of the recursion tree is bounded by $2^q q^q_y$ a naive analysis gives a bound of $2^{\circ}(q^2)q^{\circ\wedge 2}$ number of calls to the function use-interface.

At the end of the recursive calls, one of the leaves of the recursion tree corresponds to the expanded skeleton of *Topt*- **Note:** We showed that the depth in the recursion tree of the leaf that corresponds to the expanded skeleton of *Topt* is bounded by cq for some constant c. Therefore we can safely terminate the recursion at depth cq.

3.2 Linking perfect phytogenies

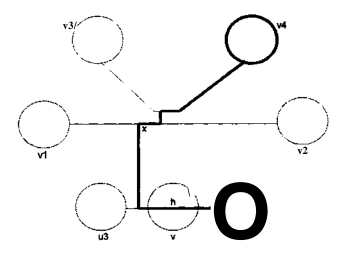


Figure 6: Proof that column h has to mutate multiple times to be an internal column that also satisfi es the properties of a distinguishing interface column.

function link-trees (skeleton S)

- 1. For each leaf super-node Si of 5 do
 - (a) build perfect phylogeny on the vertices in 5;
 - (b) let Pi < r- the set of vertices of Si
 - (c) for each mutation in Si do
 - i. if $S \setminus Si$ contains only a single state (say 0) on column c, then remove all in *Pi* that contain 1 on column *c*
 - (d) "guess" a vertex w from Pj, and let w^f be obtained from w by mutating the vertex cover between *Si* and p(Si).
 - (e) add w^{l} to p(Si and remove the leaf Si from S
- 2. repeat step 1 until 5 is empty

Figure 7: Algorithm to build the tree using the skeleton and partition

We now show how to complete the the near-perfect phylogeny once the correct partition on the super nodes have been computed (step 2(a)iD of the main pseudo-code). Note: Let the initial skeleton built using the vertex cover columns in step 2a of the main pseudo-code be S. Assume that S is isomorphic to s(Topt). Since s(Topt) does not contain any bad vertices and is optimal, all the supemodes contain perfect phytogenies.

Lemma 6 Every super node u of the skeleton $s^{n}pt$) contains a perfect phylogeny. **Proof:** For the sake of contradiction, assume not. Consider edges e, e' G u with i(e) = n(e') = c. Consider the path V connecting edges e and e'. There exists no column $d \pm Q$ that mutates an odd number of times in V according to Corollary 3.1. By the definition of the super-nodes and skeleton we know that there exists no edge e E u with fi(e) = c if c G Q. For optimality, we can not have two vertices v and v' with $l(v) = l\{v^f\}$. Therefore there can not exists edges e and e' as stated.

Using the partitions of the expanded skeleton T, we can find the partitions for the skeleton S. For the rest of the paper, we will work with skeleton S rather than T. We lose some information in this step, but it keeps the analysis simple. We now show how to construct a perfect phylogeny for the rows assigned to a super node of S. For any pair of columns that mutate within a super node, there can be at most three gametes. We can now reconstruct a unique perfect phylogeny within each super node as follows. If two columns induce two gametes, then we can arbitrarily remove one of them. If two columns induce just one gamete then we can remove both the columns. After this step, every pair of columns contain exactly three gametes. Therefore a unique perfect phylogeny can be built. We can now add back columns *i* that were removed since (\dot{z}, j) induced only two gametes. This is done by adding the mutation of *i* adjacent to the mutation of *j*. The relative ordering of *i* and *j* are determined based on which of the third gamete between *i* and *j* is present in the input (and absent from the rows of this super node). Hence we can build a unique perfect phylogeny that does not contain any bad vertices.

The notation p(v) is used to denote the parent of a vertex v in a directed tree. In the following section the term subtree T (say rooted at v) of the rooted tree T^, is used to refer to the induced tree T rooted at vertex v obtained by removing the edge (v,p(r;)) from T^.

Lemma 7 In the rooted tree T^{\wedge} , consider any induced subtree T^{f} . If $T^{\wedge \lambda^{l}}$ contains vertices v and v^{f} such that l(v)[c] = 0 and $l(v^{r})[c] = 1$ then there exists at least one edge e s.t /x(e) = c in $T_{opt} \setminus T^{f}$ **Proof:** Assume not, then the mutation of column c occurs in T'. But T' and $Tapt \setminus T^{f}$ are connected by a single edge. Therefore, $TaptXr^{l}$ contains only one of either 0 or 1 in column c, contradiction.

Corollary 8 If both T" and Tapt $\backslash T^{f}$ each contain a pair of vertices v, v' s.t l(v)[c] = 0 and l(v')[c] = I for some column c then there are at least 2 edges e, e' G T^t \$-* A*(e) = /i(e') = c.

Note that in the rooted *Topu* the perfect phylogeny inside each super node u can be decomposed into a minimal connected component of perfect phylogeny on real vertices T£, called core tree. The edges in T_U\TJ are called peripheral edges. There could be several connected components of peripheral edges inside each super node. Note that all interface edges are peripheral edges. At this point, it is easy to compute the unique core tree, which is just the perfect phylogeny on the rows that have been partitioned to any super node. For this last part of the proof, we will be considering the optimal tree T^{\wedge} with the following additional property: no super node u (except the root) contains a degree two vertex that is adjacent to both a peripheral edge and the vertex cover edge connecting u to super node p(u).

Lemma 9 There exists an optimal tree T^t pt that contains no vertex v in super node u with the property that degree(v) = 2 and v is adjacent to both an peripheral edge and the vertex cover edge connecting u to super node p(u).

Proof: Assume not, then consider the optimal tree T^{\wedge} that contains the minimum number of vertices v with the above property. One edge adjacent to v is the vertex cover edge say e_v , with say $/i(e_v) = k$. The second edge connects to a vertex v^l inside super node u and let /i(v, v') = /. Now we can flip the ordering of mutations of e^{\wedge} and (v, t/) such that $^{(e^{\wedge})} = I$ and /i(v, v') = k to obtain a new tree T $^{\wedge}$. Note that this is just a trivial operation of transform $p^{\wedge i}$ on steiner vertex v. It is easy to see that the only pair of columns in v of T^{\wedge}_t that could contain a new gamete not present in the original tree $T^{\wedge}t$ contains one column of the vertex cover. Therefore in T $^{\wedge}$, all the vertices are good (assuming they were in the original tree). The number of vertices in the super node p(u) increased by one and the number of super nodes in u decreased

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by one since v was previously part of u and now is part of $p\{u\}$. Therefore the number of vertices with the property given in the claim decreased by one, a contradiction to the assumption.

In s(Topt), consider a super node ti, that has children super nodes ci, •••, c*. The goal is to discover all the edges of T^{\wedge} bottom-up. Let 5, be the tree present inside the super node c* in T^. Note that we are using the super node *Si* to refer to the super node of both the skeleton *S* and that of (7^{\wedge}) . Let 5J be the subtree of $T^{\wedge}pt$ rooted on a vertex of *a*, obtained by removing the vertex cover edge connecting *Ci* to *u*. Note that the tree S[restricted to the nodes of c, is I }.

Assume inductively that we have computed the trees inside super nodes Si descendants and all the of Si. In otherwords the subtrees 5_{t-}^{r} of T^{\wedge} have been found. Now we proceed to construct the tree inside super node u. Note that this amounts to building the core tree of u and connecting vertices of u to vertices in 5_i as in *Tapt* (which is equivalent to computing the peripheral edges). Consider one by one each of the trees S%. We start by adding all vertices of Si into a selected set P. At any time if |P| < q, then we have completed processing Si. If not, then for each mutation of column c in Si, we check if $S \setminus S_{1}^{r}$ contains both states 0 and 1 on c. If so, we ignore c. By claim 7, the fact that there are at most q multiple mutations for any character c and that Si is a perfect phytogeny, we ignore at most q mutations of \pounds ;. If not, then wlog say 0 is the only state in column c that occurs in $S \setminus T_{1}^{r}$. If any vertex in T* contains a 1 in column c and is present in P, then we remove it from P.

Claim 10 Only the selected vertices in P can be vertices of T^{h} that connects Ci ton.

Proof: There exists exactly one edge $e = (vo, ^i)$, s.t fi(e) = c in *Si*. The edge e partitions *Si* into *Sf* and *SJ* based on the value on the column c and assume $VQ \in Sf$ and $v \in 5/$. For the sake of contradiction assume that in T^t , a vertex v from $S \in v$ connects to u via the vertex cover edge. Clearly every path to a real vertex in *Tapt*S from v contains an edge $e' = (v[, v'_o)$ with /i(e') = c.

Consider the edges e' that is the first mutation of column c in every path from e to a real vertex in $Topt Si^{2}$. Let M be the set of columns that mutate in the path connecting vo and v. Since there are no bad vertices, all mutations of columns except vertex cover, between e and e^{r} occur even number of times. Therefore, specifically vo and v'_{o} are identical in all the columns in M. Therefore, connecting VQ instead of v to u via vertex cover and deleting all mutations of M that occurs in the path between e and e' results in a tree of smaller cost, a contradiction to optimality.

We continue by finding sets *P* for each of 5^{*}. Now, using exhaustive search, we pick one by one every possible way of selecting a vertex from each of the selected sets *P* for each of *S%*. One would be the correct set of vertices as used in *T*[^]. For each *v* selected from *Si*, we mutate the column corresponding to the vertex cover edge between super nodes *Ci* and *u* and add to set *A*. We now add *A* to the super node *u* and re-construct the perfect phytogeny of *u*. It follows from the claim that we have now completely identified the tree in *u*. It is easy to implement the above procedure in $q^{\circ}M$ time.

Theorem 11 The new algorithm solves the q-near-perfect phytogeny problem in $O(2^{\circ \wedge q^2} \setminus nm + n^2))$ time. **Proof:** The proof follows from the above lemmas and Theorem 2.

4 Discussion

We now give a quick summary of how the running time can be improved to $q^{\circ n}m^2$. The only step that needs to be improved is use-interface. Intuitively, instead of building the skeleton top-down, we can

construct it bottom-up as follows. Assume that we have a skeleton T and we complete the use-interface step as described above to obtain a tree where the super nodes have unique tags. Let this final skeleton be T2. We are now going to show that we can arrive at tree T using a modified routine of use-interface. It is easy to guess the final topology of the skeleton T2 in time q^{\bullet} even before executing the function us e - in t e r f a c e since T2 has O(q) edges. Let this unlabeled tree be T'. We can guess the labels /i(e) for all edges e G T where /i(e) is either a vertex cover column or a column found in function partition. This gives us a partially labelled tree T'', where the only unlabeled edges are those that were found using the old u s e - i n t e r f a c e function. Note that in the current tree T', the connected components of unlabeled edges are exactly the super nodes of the skeleton T. We can now execute the u s e - i n t e r f a c e function as described above, except that we do not add mutations of columns h into an existing skeleton (as in step 5). Instead we guess all unlabeled edges e G T' such that n(e) = h. If we guess that column h mutates i number of times, then this step can be performed in time $\binom{2}{i}$, where p is the number of unlabeled edges currently in the tree. As before we perform u s e - i n t e r f a c e on each guess recursively. The following recursion bounds the number of calls to u s e - i n t e r f a c e when the current tree has p unlabeled edges:

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$$T(p) \leq \sum_{i=1}^{p} p^{i} T(p-i)$$

This recursion can be solved for $T(p) = q^{2q}$. Note that the new function $u \le int \le r \le n$, exactly explores the same trees as the old function. This is just a tighter analysis of the running time obtained by making sure that we don't repeatedly explore the same tree several times. Such a bottom-up construction can be extended to include the partition function as well. The algorithm now becomes simple to implement, since we can guess the final skeleton's topology up front and just guess the labels of the skeleton as the algorithm progresses.

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Appendix A: Proof of Theorem 12

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Theorem 12 It is possible to construct a q-near-perfect phylogeny on the original input (before preprocessing) using the above algorithm.

Proof: Let I be the original input and I' be the pre-processed input. If there is a tree with penalty q on I, then there exists a tree with penalty at most q on I'. Since the above algorithm goes through all trees with less than q penalty, we can add back the deleted columns and check if the penalty is still less than q. In order to show that such an algorithm will find an optimal tree, we need to prove that there always exists an optimal tree with this structure.

Consider any optimal tree T on the input I. Consider two identical columns a and b in I. We claim that we can modify the optimal tree in such a way that only the mutations of a or b are changed and the resulting tree will have all the mutations of a and b adjacent to each other. Suppose a mutation of a doesn't have a mutation of b adjacent to each other, then the vertices between them are steiner. Therefore, we can apply the transform τ_a to the neighborhood of steiner vertices. As a result of the transform, we get the mutations of a and b adjacent to each other. This also shows that the number of mutations of a and b in tree T is exactly same.

Appendix B: Proof of Theorem 2

In this section, we prove Theorem 2. Recall that a vertex v is *bad* w.r.t a pair of columns (b, c) if it has a gamete for that pair which doesn't appear in the input. Theorem 2 asserts that there is an optimal tree which doesn't have any bad vertices, i.e. all its vertices have gametes that appear in the input.

The main idea is to take any optimum solution and to transform it into one which has the above property. This is achieved through a series of transformations that reduce the number of bad vertices. We first prove some simple facts about the OPT solution.

Definition 9 We call a column c clean in a tree T, if all the vertices of T are good w.r.t (c, x) for all columns x.

Fact 13 If the column x is clean in the tree T and y is any other column, then the following properties hold:

- 1. Between any two mutations of column x, there is an even number of mutations of column y.
- 2. Between any two mutations of column y there is an even number of mutations of x.

Definition 10 Let u be a bad vertex w.r.t (b, c). We define the bad neighborhood of the vertex u w.r.t (b, c) to be

 $N_{b,c}(u) = \{v \mid v \text{ is bad w.r.t } (b,c) \& all vertices on the path <math>u \to v \text{ are bad w.r.t } (b,c)\}.$

Definition 11 The boundary of $N_{b,c}(u)$ is the set of edges that connect vertices in $N_{b,c}(u)$ to the rest of the tree T.

Fact 14 $N_{b,c}(u)$ is the connected component of bad vertices of tree T containing u.

Fact 15 The boundary of $N_{b,c}(u)$ consists of only the mutations of columns b and c.

Claim 16 The optimal tree contains equal number of mutations of columns b and c.

Proof: If not then the following transformation reduces the number of mutations in the tree. This contradicts the optimality of the tree. \Box

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Transform τ We now describe our transform that reduces the number of bad vertices in the optimum tree. Consider all the vertices in the optimum tree that are bad w.r.t (b, c). Let V_1, V_2, \ldots, V_k denote the bad neighborhoods w.r.t (b, c). The transform τ_b (resp. τ_c) changes V_1, \ldots, V_k simultaneously as follows: delete the mutations of column b (resp. c) from the boundaries of the bad components and replace every mutation of c on the boundaries by a mutation of c followed by a mutation of b (resp. replace b by bc). Let T' denote the tree obtained by applying the transform τ_b on tree T. Note that both T and T' are optimal trees.

We prove the following two properties about the transform τ .

Lemma 17 After applying the transform τ_b (or τ_c), all vertices of the tree T' are good w.r.t (b, c).

Proof: The vertices that were good w.r.t (b, c) stay good even after the transform τ_b . All the bad vertices w.r.t (b, c) were replaced by good vertices during transform τ_b . So no bad vertices are left.

Fact 18 If the tree T does not have any bad vertices w.r.t (c, x), then the resulting tree T' after applying transform τ_b will not have any bad vertices w.r.t (c, x).

Lemma 19 If the column x is clean, then the transform τ_b will not create any bad vertices w.r.t (b, x).

Proof of Theorem 2: Consider a lexicographic order on the columns c_1, c_2, \ldots . We start with an optimum tree T. We first make column c_1 clean by considering bad vertices w.r.t (c_1, c_i) for each i and applying transform τ_{c_i} . Note that the transform τ_{c_i} will only affect pairs $(c_i, c_{i'})$. It does not create bad vertices w.r.t $(c_1, c_{i'})$. From Lemma 17, it follows that at the end of this step column c_1 will be clean.

Now inductively assume that columns c_1, \ldots, c_j are clean. Next we look at all the bad vertices w.r.t (c_{j+1}, c_i) (for i > j + 1) for each *i* and apply transform τ_{c_i} . Note that the transform τ_{c_i} doesn't create any bad vertices w.r.t $(c_j, c_{i'})$. Moreover, by Lemma 19, we can say that the resulting tree has columns c_1, \ldots, c_j clean. Moreover, it follows from Lemma 17 that at the end of this step the column c_{j+1} is also clean.

Therefore, in the end all the columns are clean. In other words, there are no bad vertices in the tree. \Box

We prove the Lemma 19 through the following series of claims.

Claim 20 If the column x is clean in the tree T, then mutations of x in T are all outside $\cup_i V_i$ or all inside.

Proof: For the sake of contradiction, assume that there is a mutation of *x* inside $V \setminus$ and another one outside *UiVi*. Consider a path between these two mutations. It is easy to see that either 6 or c mutates odd number of times between the two mutations of *x*. This contradicts the fact that *x* is clean.

Claim 21 Transform r& does not create odd number of mutations of between two mutations of a clean column x.

Proof: Using the result of Claim 20, we consider following two cases.

case 1 (All x 's are outside) Consider a pair of mutations of x. If it doesn't have any mutation of b or c between them, then their parity will not change. If there were an even number of mutations of 6 in between, then in transform r&, an even number of mutations (twice the number of bad neighborhoods on the path) will get deleted and hence the parity won't change. If there were an even number of c's, then two mutations of b will be added for every bad neighborhood on the path. Overall, parity does not change.

case 2 (All x 's are inside) In this case, every c gets replaced by cb. Since there were even number of 6's and c's an even number of 6's get created and an even number of 6's get deleted between any pair of x's. Thus the parities do not change.

Remark 1 *Claim 21 is required only if the column x mutates multiple times in the optimal tree.*

Note that in transform r& only 6's are deleted or added. So a similar claim for *c* holds automatically.

Claim 22 The tree T^{l} resulting from transform T > does not leave odd mutations of a clean column x's between two mutations of b's.

Proof: We show that between every adjacent pair of mutations of *b* in tree T', there are even number of mutations of *x*. For the sake of contradiction assume that a pair of 6's has odd number of re's in tree T^d resulting after the transform r& was applied. Let $b \mid$ and 62 denote the two mutations of 6 that sandwich an odd number of mutations of *x*. Note that both 61 and 62 cannot be preset in the tree T. Without loss of generality, assume that 61 was created in tree T". Therefore, there is a mutation of column c next to 61. Call itci.

In the tree T, the column x was clean. Since the transform 77, does not change in mutations of c, no mutation of x is sandwiched between two mutations of c in the new tree T'.

If 62 is a new mutation added to tree T', then let C2 be a mutation of the column *c* which is next to 62 in the tree T'. In this case, the odd number of mutations of 2; are sandwiched between c and C2 which is not possible. Therefore 62 is present in the tree *T*.

We use Claim 20 to separate following two cases.

case 1 (All x's are outside) Let N_c^{\wedge} denote the bad neighborhood where 61 was created. Let 61 be a mutation of 6 that was deleted from the boundary of iV&₃. Then the path from 61 to $c \setminus$ in tree *T* contains no mutations of *x*, while the path from $c \setminus$ to 62 contains an odd number of mutations of *x*. Therefore, in tree *T* there are an odd number of mutations of *x* between 61 and 62, which contradicts the fact that *x* was clean in tree T.

case 2 (All x 's are inside) Note that 62 could not have been on the boundary of a bad neighborhood in the tree T, otherwise it would have been removed in the tree T'. Since all the mutations of x were inside bad neighborhoods, at least one of the neighborhoods on the path from 61 to 62 had an odd number of mutations. If this bad neighborhood is the one where 61 was created in the transform, then the path 61 to 62 includes the mutation c. In T'', all the vertices are good w.r.t (b, c). Hence there must be another mutation of c (call it C2) between 61 and 62- Note that the path c and C2 contains an odd number of mutations of x. This is a contradiction. Therefore, the bad neighborhood containing odd number of mutations of x is not the one where 61 was created. But in that case, consider the path from 61 to 62 in the old tree T. It must have entered and exited the bad neighborhood via mutations of b. Thus the odd number of mutations of x were sandwiched between two mutations of b in the old tree T. This is a contradiction to the fact that x was clean.

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