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On the Complexity of Computing MP Distance Between Binary Phylogenetic Trees

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Abstract. Within the field of phylogenetics there is great interest in distance measures to quantify the dissimilarity of two trees. Recently, a new distance measure has been proposed: the Maximum Parsimony (MP) distance. This is based on the difference of the parsimony scores of a single character on both trees under consideration, and the goal is to find the character which maximizes this difference. Here we show that computation of MP distance on two *binary* phylogenetic trees is NP-hard. This is a highly nontrivial extension of an earlier NP-hardness proof for two multifurcating phylogenetic trees, and it is particularly relevant given the prominence of binary trees in the phylogenetics literature. As a corollary to the main hardness result we show that computation of MP distance is also hard on binary trees if the number of states available is bounded. In fact, via a different reduction we show that it is hard even if only two states are available. Finally, as a first response to this hardness we give a simple Integer Linear Program (ILP) formulation which is capable of computing the MP distance exactly for small trees (and for larger trees when only a small number of character states are available) and which is used to computationally verify several auxiliary results required by the hardness proofs.

Keywords: Maximum Parsimony, phylogenetics, tree metrics, NP-hard, binary trees

1. Introduction

When present day species are considered and their evolutionary relationships are to be investigated, phylogeneticists often seek to estimate the best evolutionary tree explaining the given species data (e.g., DNA alignments). However, it is well known that different data sets on the same species can lead to different trees, or that different phylogenetic tree estimation methods, e.g., Maximum Parsimony or Maximum Likelihood or distance based methods, can lead to different trees even for the same



data set [6, 11]. Thus, in practice one is often confronted with multiple trees, and it is therefore interesting to measure how different these trees really are. A new way of determining their relative similarity has recently been proposed [5]: the Maximum Parsimony distance (or MP distance, for short).

This metric basically requires, the search for a character which has a low parsimony score on one of the trees involved and a high score on the other one. In [5] it has been shown that calculating the MP distance between two trees is NP-hard. The proof presented there required non-binary trees (sometimes also called *multifurcating* trees). This was not entirely satisfactory, for the following reason. In many branches of phylogenetics multifurcating trees are used to model uncertainty about the precise order of branching events [14], in which case the term unresolved is often used instead of multifurcating. Distance measures which interpret multifurcations this way often have the property that the distance decreases, or in the worst case stays the same, if one or both of the input trees become more unresolved [12]. However, the parsimony score of a single tree increases (or in the best case stays the same) if its edges are contracted to create multifurcations. This is why algorithms that compute Maximum Parsimony trees usually output binary trees: a non-binary solution can be refined into a binary solution without loss of quality. Given this traditional emphasis on binary trees in the parsimony literature, and the fact that evolutionary events such as mutation or speciation are understood to split a lineage into two parts, not three or more [9], it is logical to explore the complexity of MP distance on binary trees.

In this paper, we answer this question by showing that computing the MP distance between two binary trees is, unfortunately, also NP-hard. This is by no means a simple extension of the hardness proofs in [5]. To prove hardness in the present case we are required to develop a rather elaborate array of novel gadgets and arguments, with a strong graph-theoretical flavour.

Moreover, we show as a corollary to the main theorem that this hardness remains if we restrict the number of character states to four (or more). Note that this covers the most important biological applications, as the DNA and RNA alphabets consist of four character states each, and the protein alphabet consists of 20 states. However, when morphological data is analyzed, binary characters are also often relevant, which is why we consider this case, too. We show that when restricted to two character states, calculating the MP distance is not just NP-hard, but even APX-hard, which means that there exists a constant c > 1 such that a polynomial-time c-approximation is impossible unless P = NP.

As a tentative first step towards addressing the NP-hardness of the MP distance, we present a simple Integer Linear Program (ILP) which calculates this distance (both on a bounded number of states as well as in the unbounded case). The ILP is rather "explicit" in the sense that it has a static, constraint-based formulation of Fitch's algorithm embedded within it. Although faster than naive brute force algorithms, the ILP for an unbounded number of states does not scale well and is limited to trees with approximately 16 taxa. On the other hand, the ILP for binary characters is fast: it can cope with trees with up to 100 taxa in reasonable time. In both cases the ILP is fast enough to verify the MP distance of a number of gadgets used in the hardness proofs. An implementation of this ILP has been made publicly available at http://skelk.sdf-eu.org/mpdistbinary/[13].

2. Notation

Recall that an *unrooted phylogenetic X-tree* is a tree $\mathcal{T}=(V(\mathcal{T}),E(\mathcal{T}))$ on a leaf set $X=\{1,\ldots,n\}\subset V(\mathcal{T})$. Such a tree is named *binary* if it has only vertices of degree 1 (leaves) or 3 (internal vertices). A *rooted phylogenetic X-tree* additionally has one vertex specified as the *root*, and such a rooted tree is named *binary* if the root has degree 2 and all other vertices are of degree 1 (leaves) or 3 (internal vertices). Note that two leaves are said to form a *cherry*, if they are connected to the same inner node. Moreover, a rooted binary tree on three taxa is also often referred to as a *rooted triplet*, and a rooted tree with only one cherry is also called a *caterpillar tree* or *caterpillar* for short. We often denote trees in the well-known Newick format [4], which uses nested parentheses to group species together according to their degree of relatedness. For instance, the tree ((1,2),(3,4)) is a tree with two so-called cherries (1,2) and (3,4) and a root between the two.

Furthermore, recall that a *character* f is a function $f: X \to \mathcal{C}$ for some set $\mathcal{C} := \{c_1, c_2, c_3, \ldots, c_k\}$ of k *character states* $(k \in \mathbb{N})$. Often, k is assumed to equal 4 in order for \mathcal{C} to represent the DNA alphabet $\{A, C, G, T\}$, but in the present paper k is not restricted this way but can be any natural number unless stated otherwise. Note that in the special case where |f(X)| = 2, we also refer to f as a binary character. In general, when |f(X)| = r, f is called an *r-state character*. In order to shorten the notation, it is customary to write, for instance, f = AACC instead of f(1) = A, f(2) = A, f(3) = C, and f(4) = C. Note that each r-state character f on taxon set f partitions f into f non-empty and non-overlapping subsets f if and only if f(f) = f(f).

Note that in this paper, we refer to a character always with its underlying taxon set partition in mind, i.e., for instance we do not distinguish between *AACC*, *CCAA*, and *CCGG*, and so on. Moreover, when there is no ambiguity and when the stated result holds for both rooted and unrooted trees, we often just write 'tree' or 'phylogenetic tree' when referring to a phylogenetic *X*-tree.

An extension of a character f to $V(\mathcal{T})$ is a map $g\colon V(\mathcal{T})\to \mathcal{C}$ such that g(i)=f(i) for all i in X. For such an extension g of f, we denote by $l_g(\mathcal{T})$ the number of edges $e=\{u,v\}$ in \mathcal{T} on which a substitution occurs, i.e., where $g(u)\neq g(v)$. Such substitutions are also often referred to as mutations or changes. The parsimony score or parsimony length of a character f on \mathcal{T} , denoted by $l_f(\mathcal{T})$, is obtained by minimizing $l_g(\mathcal{T})$ over all possible extensions g of f. For binary trees \mathcal{T} , the parsimony score of a character f can easily be calculated with the Fitch algorithm [7], whose correctness was formally proven by Hartigan in 1973 [8]. Recall that the bottom-up phase of the Fitch algorithm starts at the labelled leaves and assigns to the unlabeled parent of two nodes the intersection of both children's label set if it is non-empty, or the union otherwise. The top-down phase then starts at the root with an arbitrary choice of the root states suggested by the bottom-up phase and keeps the current state for the descending nodes whenever this is contained in the label set of these nodes, and takes an arbitrary state out of the label set otherwise.

This paper deals with the so-called *parsimony distance* d_{MP} as introduced in [5]. This distance is defined as follows: given two phylogenetic trees \mathcal{T}_1 and \mathcal{T}_2 on the

same set X of taxa, the parsimony distance between these trees is defined as

$$d_{MP}(\mathcal{T}_1, \mathcal{T}_2) = \max_{f} \left| l_f(\mathcal{T}_1) - l_f(\mathcal{T}_2) \right|, \tag{2.1}$$

where the maximum is taken over all characters f on X. A character f which maximizes this distance is called an *optimal character*. Note that, due to the fact that the parsimony score of a tree (for a given character) is not affected by the presence or absence of a root, parsimony distance is also oblivious to whether the input trees are rooted or unrooted. Later, in Section 5, we will consider the bounded-states variant of $d_{MP}(\mathcal{T}_1, \mathcal{T}_2)$, denoted $d_{MP}^i(\mathcal{T}_1, \mathcal{T}_2)$, where $i \geq 1$. This is defined as in (2.1) except that the characters f are restricted to having at most i states. Clearly, for all i, $d_{MP}^i(\mathcal{T}_1, \mathcal{T}_2) \leq d_{MP}(\mathcal{T}_1, \mathcal{T}_2)$.

For some proofs in this paper we need the notion of a maximum agreement forest, which is closely linked to the so-called rooted subtree prune and regraft distance or rSPR distance for short. Recall that, informally, an agreement forest of two rooted phylogenetic trees is a set of subtrees which are identical in both trees and which in total contain all leaves, see, e.g., [2]. A maximum agreement forest is an agreement forest with minimum number of components. A single rSPR move involves moving to a neighboring rooted tree by detaching a branch and re-attaching it elsewhere. The rSPR distance d_{rSPR} is the minimum number of rSPR moves required to transform one rooted tree into another. Maximum agreement forests and rSPR distance are closely linked by the well-known result that, modulo a rooting technicality, an agreement forest of two rooted trees with m components yields a set of m-1 rSPR moves which turn the first tree into the second one [2].

3. Preliminaries

The following observation, which we will use extensively and implicitly throughout the article, appeared unchanged in our earlier work [5].

Observation 3.1. Let f be a character on X and T a tree on X. Let f' be any character obtained from f by changing the state of exactly one taxon. Then $l_f(T) - 1 \le l_{f'}(T) \le l_f(T) + 1$, i.e., the parsimony score can change by at most one.

Proof. Suppose $l_{f'}(\mathcal{T}) \leq l_f(\mathcal{T}) - 2$. Consider any extension of f' to the interior nodes of \mathcal{T} that achieves $l_{f'}(\mathcal{T})$ mutations. Using the same extension but on f gives at most $l_{f'}(\mathcal{T}) + 1$ mutations, because only one taxon changed state. So $l_f(\mathcal{T}) \leq l_{f'}(\mathcal{T}) + 1 \leq l_f(\mathcal{T}) - 1$, which is a contradiction. In the other direction, take any optimal extension of f and apply it to f'. At most one extra mutation will be created, so $l_{f'}(\mathcal{T}) \leq l_f(\mathcal{T}) + 1$.

A more general version of the following lemma appeared earlier in [5]. Here we have specialized the lemma and its proof to apply to rooted binary trees, which is the type of trees we will construct in the subsequent hardness reductions.

Lemma 3.2. Let f be an optimal character for two rooted, binary trees \mathcal{T}_1 and \mathcal{T}_2 , i.e., $d_{MP}(\mathcal{T}_1, \mathcal{T}_2) = \left| l_f(\mathcal{T}_2) - l_f(\mathcal{T}_1) \right|$. Without loss of generality, assume $l_f(\mathcal{T}_1) < l_f(\mathcal{T}_2)$. Then we can construct in polynomial time an optimal character f' with the following

property: $l_{f'}(T_1) < l_{f'}(T_2)$ and for each vertex u of T_1 such that both u's children are leaves (i.e., they form a cherry), f' assigns both children of u the same state.

Proof. Consider a vertex u of \mathcal{T}_1 such that both of its children are taxa, but such that f assigns the two children different states. We calculate an optimal extension of f to the interior nodes of \mathcal{T}_1 by applying Fitch's algorithm. Let s be the state allocated to u by the top-down phase of the Fitch algorithm. Choose the child of u that does not have state s and change its state to s. This yields a new character f^* . Clearly, $l_{f^*}(\mathcal{T}_1) < l_f(\mathcal{T}_1)$, simply by using the same extension that the Fitch algorithm gave. Combining this with Observation 3.1 gives $l_{f^*}(\mathcal{T}_1) = l_f(\mathcal{T}_1) - 1$ and thus $l_{f^*}(\mathcal{T}_2) = l_f(\mathcal{T}_2) - 1$ (otherwise f could not have been optimal). Hence, f^* is also an optimal character, and $l_{f^*}(\mathcal{T}_1) < l_{f^*}(\mathcal{T}_2)$. This process can be repeated for as long as necessary. Termination in polynomial time is guaranteed because each taxon has its state changed at most once.

Observation 3.3. Lemma 3.2 also holds for optimal characters under the $d_{MP}^i(\mathcal{T}_1,\mathcal{T}_2)$ model.

Proof. The transformation in the proof of Lemma 3.2 does not increase the number of states in the character.

4. MP Distance on Binary Trees Is NP-Hard

4.1. The Symmetry-Breaking Construction

In the hardness proof in Section 4.2 we will construct two trees T_E and T_V and a central fact used in the proof of correctness of the reduction is that, for all optimal characters f, $l_f(T_E) < l_f(T_V)$. In this section we show how to construct a gadget to enforce this property. Note that all the trees constructed in this section are binary. (As we demonstrated in [5] constructing such a symmetry-breaking gadget is far easier in the non-binary case.)

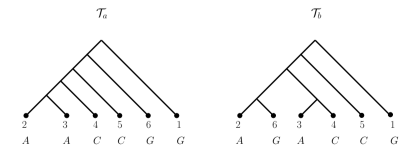


Figure 1: The two "asymmetric" trees \mathcal{T}_a and \mathcal{T}_b and an optimal character $f_{asym} = GAACCG$.

Consider the two rooted trees

$$T_a = (((((2,3),4),5),6),1),$$

$$T_b = ((((2,6),(3,4)),5),1),$$

shown in Figure 1. It can be verified computationally that $d_{MP}(\mathcal{T}_a, \mathcal{T}_b) = 2$, achieved, for example, by character $f_{asym} = GAACCG^*$ with $l_f(\mathcal{T}_a) = 2$ and $l_f(\mathcal{T}_b) = 4$.

Moreover, if f is an optimal character, then $l_f(\mathcal{T}_a) + 2 = l_f(\mathcal{T}_b)$. Expressed differently: there does not exist any optimal character f for which $l_f(\mathcal{T}_a) > l_f(\mathcal{T}_b)$, so the instance is "asymmetric". For two trees \mathcal{T}_1 and \mathcal{T}_2 , let

$$gap(\mathcal{T}_1, \mathcal{T}_2) = \left| \max_f (l_f(\mathcal{T}_2) - l_f(\mathcal{T}_1)) - \max_f (l_f(\mathcal{T}_1) - l_f(\mathcal{T}_2)) \right|,$$

where f ranges over all characters (not just optimal ones). Note that $gap(\mathcal{T}_a, \mathcal{T}_b) = 1$ because $\max_f \left(l_f(\mathcal{T}_b) - l_f(\mathcal{T}_a) \right) = 2$ and $\max_f \left(l_f(\mathcal{T}_a) - l_f(\mathcal{T}_b) \right) = 1$, where, e.g., the character f = AACCAA achieves $l_f(\mathcal{T}_a) - l_f(\mathcal{T}_b) = 2 - 1 = 1$.

We now describe an iterative construction such that, for any desired gap g, we can construct two trees \mathcal{T}_1 and \mathcal{T}_2 , both on O(g) taxa, such that $gap(\mathcal{T}_1, \mathcal{T}_2) \ge g$.

We start with \mathcal{T}_a and \mathcal{T}_b . Let \mathcal{T}_A be the rooted tree on 12 taxa obtained by taking two disjoint copies of \mathcal{T}_a and joining them together via their roots ρ_1 , ρ_2 to a newly introduced root ρ . (Here, the copying operation is assumed to introduce new taxon labels to prevent the same taxon occurring twice in the same tree.) \mathcal{T}_B is defined the same way, but with respect to \mathcal{T}_b .

Claim 4.1.
$$gap(T_A, T_B) \geq 2$$
.

Proof. We will show that $\max_f (l_f(\mathcal{T}_B) - l_f(\mathcal{T}_A)) \ge 4$ and $\max_f (l_f(\mathcal{T}_A) - l_f(\mathcal{T}_B)) \le 4$ 2, from which the claim will follow. Let f be a character such that $l_f(\mathcal{T}_a) + 2 = l_f(\mathcal{T}_b)$, i.e., f is an optimal character for \mathcal{T}_a , \mathcal{T}_b . We extend character f to become character F on T_A , T_B in the natural way, i.e., disjoint copies of the same taxon receive the same character state. If we run the bottom-up phase of Fitch's algorithm on \mathcal{T}_A and \mathcal{T}_B , we observe that each copy of \mathcal{T}_a induces 2 fewer mutations than its corresponding copy of \mathcal{T}_b . Moreover, the set of states identified (by the Fitch algorithm's bottom-up phase) to be possible at ρ_1 will be equal to the set of states identified to be possible at ρ_2 , so there will be no mutations incurred in \mathcal{T}_A on the two edges incident to its root ρ . By the same argument, there will be no mutations incurred in \mathcal{T}_B on the edges incident to its root. Hence, $l_F(\mathcal{T}_B) - l_F(\mathcal{T}_A) \ge 4$. Showing $\max_f (l_f(T_A) - l_f(T_B)) \le 2$ is possible analytically but it is technical. We therefore omit the proof, noting however that we have used an exhaustive computational search to verify that (a) $\max_f (l_f(T_B) - l_f(T_A)) = 4$, where the maximum is reached, e.g., by f = ABBCCAACCBBA and (b) $\max_{f} (l_f(T_A) - l_f(T_B)) = 2$, where the maximum is reached, e.g., by f = AABBAAAABBAA. Note that our ILP described in Section 6 can also be used to verify the claim. The computational search thus allows us to draw the slightly stronger conclusion that $gap(T_A, T_B) = 2$.

Let \mathcal{T}_A^k be the rooted tree on 12k taxa obtained by arranging k disjoint copies of \mathcal{T}_A along a caterpillar backbone. That is, $\mathcal{T}_A^1 = \mathcal{T}_A$ and for k > 1, \mathcal{T}_A^k is obtained by joining \mathcal{T}_A^{k-1} and \mathcal{T}_A via a new root. \mathcal{T}_B^k is defined analogously.

^{*} Note that for this specific character there exist optimal extensions in both trees such that the root is allocated state G.

Claim 4.2.
$$gap(T_A^2, T_B^2) \ge 3$$
.

Proof. By extending the character F to \mathcal{T}_A^2 , \mathcal{T}_B^2 in the usual fashion, and using the same Fitch algorithm-based argument as in the previous proof, we see that $\max_f \left(l_f(\mathcal{T}_B^2) - l_f(\mathcal{T}_A^2) \right) \geq 8$. On the other hand, due to the fact that $\max_f \left(l_f(\mathcal{T}_A) - l_f(\mathcal{T}_B) \right) \leq 2$, the total number of mutations incurred inside the two copies of \mathcal{T}_A can in total be at most 4 more than the total number of mutations incurred inside the two copies of \mathcal{T}_B . In the worst case, \mathcal{T}_A^2 can perhaps also suffer a single mutation on the two edges incident to the root, while \mathcal{T}_B^2 suffers none, so $\max_f \left(l_f(\mathcal{T}_A^2) - l_f(\mathcal{T}_B^2) \right) \leq 5$. The claim follows.

Lemma 4.3. For
$$k \ge 1$$
, $gap(T_A^k, T_B^k) \ge k + 1$.

Proof. We prove this statement by induction. For $k \in \{1, 2\}$ the lemma has already been proved, so assume $k \ge 3$. By continuing the arguments used in the previous claims, we see that

$$\max_{f} \left(l_{f} \left(\mathcal{T}_{B}^{k} \right) - l_{f} \left(\mathcal{T}_{A}^{k} \right) \right) \ge \max_{f} \left(l_{f} \left(\mathcal{T}_{B}^{k-1} \right) - l_{f} \left(\mathcal{T}_{A}^{k-1} \right) \right) + \max_{f} \left(l_{f} \left(\mathcal{T}_{B} \right) - l_{f} \left(\mathcal{T}_{A} \right) \right)$$

and

$$\max_{f} \left(l_f \left(\mathcal{T}_A^k \right) - l_f \left(\mathcal{T}_B^k \right) \right) \le \max_{f} \left(l_f \left(\mathcal{T}_A^{k-1} \right) - l_f \left(\mathcal{T}_B^{k-1} \right) \right)$$

$$+ \max_{f} \left(l_f (\mathcal{T}_A) - l_f (\mathcal{T}_B) \right) + 1,$$

where the 1 in the second expression accounts for the possibility that in \mathcal{T}_A^k a mutation is incurred on one of the root edges, while no such mutation is incurred in \mathcal{T}_B^k . Combining the above with the fact that $\max_f \left(l_f(\mathcal{T}_B) - l_f(\mathcal{T}_A)\right) = 4$, $\max_f \left(l_f(\mathcal{T}_A) - l_f(\mathcal{T}_B)\right) = 2$, $\max_f \left(l_f\left(\mathcal{T}_B^2\right) - l_f\left(\mathcal{T}_A^2\right)\right) \geq 8$, and $\max_f \left(l_f\left(\mathcal{T}_A^2\right) - l_f\left(\mathcal{T}_B^2\right)\right) \leq 5$, we obtain the desired result.

In addition to Lemma 4.3, we actually also need to know a (polynomial-time computable) expression for $d_{MP}(T_A^k, T_B^k)$. Conveniently, we have a closed expression for this.

Lemma 4.4. For
$$k \ge 2$$
, $d_{MP}(T_A^k, T_B^k) = 8 + 4(k-2) = 4k$.

Proof. From the proof of Lemma 4.3 we know that

$$d_{MP}\left(\mathcal{T}_{A}^{k},\mathcal{T}_{B}^{k}\right) = max_{f}\left(l_{f}\left(\mathcal{T}_{B}^{k}\right) - l_{f}\left(\mathcal{T}_{A}^{k}\right)\right).$$

Due to the recurrence shown in the proof of that lemma we see,

$$max_f\left(l_f\left(\mathcal{T}_B^k\right) - l_f\left(\mathcal{T}_A^k\right)\right) \ge 8 + 4(k-2).$$

We will complete the proof by showing $d_{MP}(T_A^k, T_B^k) \leq 8 + 4(k-2)$. To do this, we exploit the fact (proven in [5]) that $d_{MP}(T_A^k, T_B^k) \leq d_{rSPR}(T_A^k, T_B^k)$, i.e., MP distance is a lower bound on the well-known rooted subtree prune and regraft (rSPR) distance. In particular, we prove that $d_{rSPR}(T_A^k, T_B^k) \leq 8 + 4(k-2)$. We do this by showing that T_A^k , T_B^k permit an agreement forest with at most 8 + 4(k-2) + 1 = 4k + 1 components. (It is well known that an agreement forest with m components yields a set of m-1 rSPR moves that turn one tree into the other, see $[2]^{\dagger}$.) Now, observe that T_A^k , T_B^k contain 4 + 2(k-2) = 2k copies of the original T_a , T_b trees, respectively. Next, observe that an agreement forest for T_a , T_b with 3 components can be obtained by placing taxon 3 and taxon 6 each in a singleton component, and $\{1, 2, 4, 5\}$ in the remaining component. To obtain an agreement forest for T_A^k , T_B^k we put all copies of taxon 3 and all copies of taxon 6 in singleton components, yielding 4k singleton components. All remaining taxa can be placed in one large component, yielding 4k + 1 components in total.

Finally, we consider the following auxiliary observation, which will be useful later.

Observation 4.5. For each $k \ge 2$, there exists an optimal character f^k on \mathcal{T}_A^k , \mathcal{T}_B^k such that f^k has 3 states, and there exist optimal extensions of f^k to both trees, such that the roots of \mathcal{T}_A^k , \mathcal{T}_B^k both receive the same state.

Proof. As noted earlier, $f_{asym} = GAACCG$ is an optimal character for \mathcal{T}_a , \mathcal{T}_b and permits optimal extensions such that the roots of both trees can be assigned state G. We can obtain an optimal character f^k on \mathcal{T}_A^k , \mathcal{T}_B^k simply by making 2k copies of f_{asym} . The optimality of f^k follows from the fact that f_{asym} is optimal for \mathcal{T}_a , \mathcal{T}_b and that in the proof of Claim 4.1, any optimal character for \mathcal{T}_a , \mathcal{T}_b can be used. Given that each copy of \mathcal{T}_a and \mathcal{T}_b can have state G allocated to its root, it follows (by continuing the bottom-up phase of the Fitch algorithm on the remainder of \mathcal{T}_A^k and \mathcal{T}_B^k) that there exist optimal extensions of f^k such that the roots of \mathcal{T}_A^k and \mathcal{T}_B^k are both allocated state G.

4.2. The Reduction

In this section we exclusively consider simple undirected graphs. Recall that a graph G = (V, E) is *cubic* if every vertex has degree exactly 3, in which case |E| = 3|V|/2. A *proper edge colouring* of a graph G is an assignment of colours to the edges such that no two adjacent edges have the same colour, where two edges are adjacent if they have a common endpoint. Let $\chi'(G)$, the *chromatic index* of G, be the minimum number of colours required to properly colour the edges of G. The classical result of Vizing (see any standard graph-theory text, such as [3]) states that for every graph G, $\Delta(G) \leq \chi'(G) \leq \Delta(G) + 1$ where $\Delta(G)$ is the maximum degree of a vertex in G. Hence, for cubic G, $\chi'(G) \in \{3,4\}$. Even for cubic graphs it is NP-hard to distinguish between these two possibilities [10].

Theorem 4.6. Computation of $d_{MP}(\mathcal{T}_1, \mathcal{T}_2)$ on binary trees is NP-hard.

[†] To utilize this agreement forest formulation of rSPR we should first append a new taxon ρ to the root of both trees. However, in this case it is easy to check that the omission of ρ does not harm the analysis.

Proof. Let G = (V, E) be a cubic graph where n = |V|. We give a polynomial-time reduction from computation of $\chi'(G)$ to computation of d_{MP} , from which NP-hardness will follow. Specifically, we will construct two trees \mathcal{T}_E and \mathcal{T}_V such that, for a certain integer P, $d_{MP}(\mathcal{T}_E, \mathcal{T}_V) = P$ if and only if $\chi'(G) = 3$. In particular, if $\chi'(G) = 4$, then $d_{MP}(\mathcal{T}_E, \mathcal{T}_V)$ will be P-1 (or less). An important difference with [5, Theorem 4.6] is that here optimal characters f will be engineered to always have the property $l_f(\mathcal{T}_E) < l_f(\mathcal{T}_V)$ and not the other way round. Informally, at optimality \mathcal{T}_E always "wins".

The high-level idea is that in \mathcal{T}_E we will choose the colours of the edges of G. In fact, for each edge we will choose three colours, all different, representing the colour of e in three different copies of G. Due to the way we construct the two trees, there will exist optimal characters in which the edge colouring (in each of the three copies of G) is *proper*. This is because, the closer an edge colouring is to being proper, the higher the parsimony score induced in \mathcal{T}_V . Within the space of proper edge colourings, we will show that it is advantageous to use as few colours as possible, because this will give the character a low parsimony score on \mathcal{T}_E . Leveraging the fact that the colours used for the three copies of each edge are all different, we will derive the conclusion that d_{MP} can reach a certain value P if and only if there is a proper edge colouring that uses only 3 colours, i.e., $\chi'(G) = 3$.

We will prove the following:

$$\chi'(G) = 3 \Rightarrow d_{MP}(T_E, T_V) = P,$$

 $\chi'(G) = 4 \Rightarrow d_{MP}(T_E, T_V) \le P - 1.$

Let M be a large integer, at most polynomially large in n, whose value we will specify later. Letting k = M, construct \mathcal{T}_A^k , \mathcal{T}_B^k (as described in the previous section). Relabel $S_E = \mathcal{T}_A^k$ and $S_V = \mathcal{T}_B^k$. By Lemma 4.3, $gap(S_E, S_V) \ge M + 1$.

The core ingredients of \mathcal{T}_E are the subtrees B, S_E , and T^{***} . We construct B by taking an arbitrary rooted binary tree on 3|V|+|E| taxa. By appending an extra taxon α just above its root, we create a new root yielding 3|V|+|E|+1 taxa in total. Note that since α is not a taxon of B, in the following we refer to B including α or B without α to stress whether or not α is considered together with B or not.

The tree T^{***} is constructed as follows. Fix an arbitrary rooted binary tree T^* on |E| leaves, identifying the leaves with elements of E. Replace each leaf u_e of T^* , where $e \in E$, with a rooted triplet to obtain T^{**} on 3|E| leaves $u_{e,j}$ where $e \in E$ and $j \in \{1,2,3\}$. Finally, replace each leaf $u_{e,j}$ of T^{**} with a rooted triplet on three taxa $x_{e,j}^u, x_{e,j}^v$, and $x_{e,j}^{edge}$ where $u, v \in V$ are the two endpoints of e. We ensure that $x_{e,j}^u, x_{e,j}^v$ are sibling to each other (i.e., form a cherry). This is T^{***} , which is depicted in Figure 2, and it has thus 9|E| taxa.

The basic idea is that each edge $e = \{u, v\}$ occurs 3 times in total, and each such occurrence consists of a cherry representing u and v, and an extra taxon ("edge") sitting just above the cherry.

The construction of T_E is concluded by joining B including α , S_E , and T^{***} as shown in Figure 3, which also introduces auxiliary taxa β_1 , β_2 , γ_1 , γ_2 . We adopt the

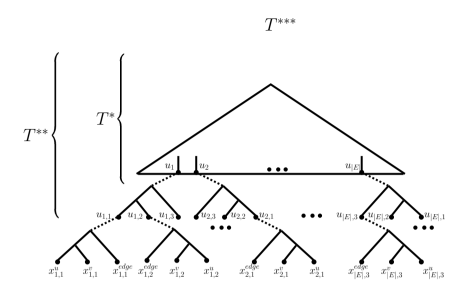


Figure 2: The tree T^{***} . Here we have identified E with the set $\{1, \ldots, |E|\}$ to simplify the figure. In the lowermost leaves we have overloaded u and v: in each case they refer to the two endpoints of the edge in question.

labels used in that figure. Summarizing, T_E contains

$$3|V| + |E| + 1 + 4 + 9|E| + 12M$$

taxa.

To construct \mathcal{T}_V we start by taking B and attaching S_V on the edge entering taxon α . Now, let

$$H = \{(v, j) | v \in V, j \in \{1, 2, 3\}\} \cup \{e | e \in E\}.$$

Clearly, |H| = 3|V| + |E|. Pick an arbitrary bijection between the taxa of B (excluding α) and the elements of H. For each edge $e \in H$, introduce a rooted triplet on the three taxa $x_{e,1}^{edge}$, $x_{e,2}^{edge}$, $x_{e,3}^{edge}$ and attach this rooted triplet on the edge entering the taxon of B corresponding to e. For each tuple $(v,j) \in H$, let $\{e,e^*,e^{**}\}$ be the 3 edges incident to v in G, introduce a rooted triplet on the three taxa $x_{e,j}^{v}$, $x_{e^*,j}^{v}$, and $x_{e^{**},j}^{v}$, and attach this rooted triplet on the edge entering the taxon of B corresponding to (v,j). Finally, we introduce a new root and join B to the new subtree on $((\beta_1, \gamma_1), (\beta_2, \gamma_2))$. This completes the construction of \mathcal{T}_V , which is depicted in Figure 4.

We are now in a position to specify the number M. We require M to be sufficiently large that, for every optimal character f, $l_f(\mathcal{T}_E) < l_f(\mathcal{T}_V)$. From Lemma 4.4 we know that there exists some character f' such that $l_{f'}(\mathcal{T}_V) - l_{f'}(\mathcal{T}_E) \ge 4M$. (In particular, we can obtain such a character by, for example, extending the character suggested by Lemma 4.4 such that all taxa outside S_E and S_V are assigned the same state.) Now, let t be the number of edges in \mathcal{T}_E that lie outside S_E . For every character f we have

$$l_f(T_E) - l_f(T_V) \le t + (4M - (M+1)).$$

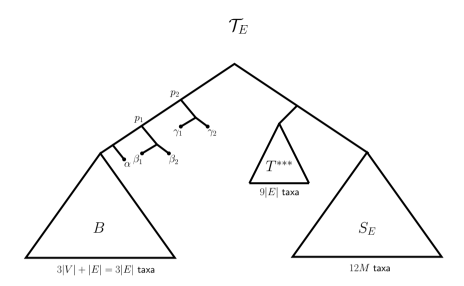


Figure 3: The tree \mathcal{T}_E . Taxon α is closely linked to subtree B as it is descending from the same root as B in \mathcal{T}_E . This root is considered in the proof. However, in \mathcal{T}_V , α shares a direct common root with S_V , not B.

The 4M term is obtained from Lemma 4.4, the (M+1) term from Lemma 4.3, and the t term arises (pessimistically) from the situation when every edge in \mathcal{T}_E (outside S_E) incurs a mutation, but no edge in \mathcal{T}_V (outside S_V) incurs a mutation. So, if we choose M such that

$$t + 4M - (M+1) < 4M,$$

it follows that for *every* optimal character f, $d_{MP}(\mathcal{T}_V, \mathcal{T}_E) = l_f(\mathcal{T}_V) - l_f(\mathcal{T}_E)$ and in particular $l_f(\mathcal{T}_V) > l_f(\mathcal{T}_E)$. Choosing M = t is therefore sufficient to achieve this. This "symmetry breaking" has far-reaching consequences which we shall heavily utilize later.

Next, let f^S be any 3-state character on the taxa in S_E and S_V such that $l_{f^S}(\mathcal{T}_V) - l_{f^S}(\mathcal{T}_E) = 4M = d_{MP}(S_E, S_V)$. This character exists and can be constructed in polynomial time thanks to Observation 4.5. Recall, in particular, that it is constructed by making many disjoint copies of the character $f_{asym} = GAACCG$.

Now, suppose $\chi'(G) = 3$. We will extend f^S to all the taxa in \mathcal{T}_E as follows, obtaining a 4-state character. Take any proper edge colouring Col of graph G using three colours red, blue, and green. We start by relabelling f^S as follows: character state G maps to blue, A maps to red and C maps to green. Next, colour all the taxa in B including α pink. Colour the cherry $\{\beta_1, \beta_2\}$ pink and the cherry $\{\gamma_1, \gamma_2\}$ blue.

Next, consider the following cyclical mapping *F*:

$$F(red, 1) \rightarrow red, F(red, 2) \rightarrow blue, F(red, 3) \rightarrow green;$$

 $F(blue, 1) \rightarrow blue, F(blue, 2) \rightarrow green, F(blue, 3) \rightarrow red;$

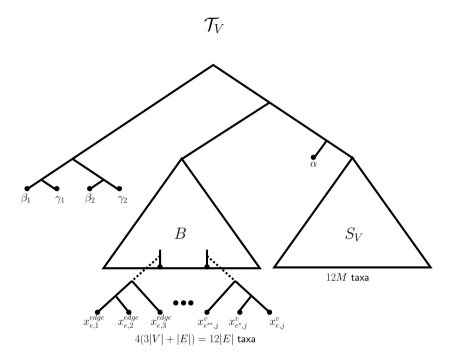


Figure 4: The tree T_V .

$$F(green, 1) \rightarrow green, F(green, 2) \rightarrow red, F(green, 3) \rightarrow blue.$$

Now, for every $e \in E$, $j \in \{1, 2, 3\}$ and letting $e = \{u, v\}$, we assign $x_{e,j}^u, x_{e,j}^v$, and $x_{e,j}^{edge}$ all the same colour: the colour F(Col(e), j) where as usual Col(e) is the colour assigned to e by the proper edge colouring Col.

Let this character be called f^{Col} . Observe that

$$l_{f^{Col}}(T_E) = 1 + 2|E| + l_{f^S}(S_E).$$

This can be confirmed by applying Fitch's algorithm. Note, in particular, that there is an optimal extension such that all the internal nodes of the tree B (including α) are coloured pink, all the nodes of the T^* part of T^{***} are blue, p_2 is blue, the root is blue, and all the unlabelled nodes are blue. The +1 is then the mutation in the transition from pink to blue on, for example, the edge between the cherries $\{\beta_1, \beta_2\}$ and $\{\gamma_1, \gamma_2\}$. There is no mutation on the edge entering the root of S_E because, by Observation 4.5 and the way we relabelled f^S , there is an optimal extension of S_E in which its root is permitted to be blue.

$$l_{f^{Col}}(\mathcal{T}_V) = 2 + 3(3|V| + |E|) + 1 + l_{f^S}(S_V).$$

The +1 term here is definitely incurred because there is an optimal extension in which the root of \mathcal{T}_V and α are both coloured pink, but pink is not used in f^S , so there will

then be a mutation on the edge entering the root of S_V . The +2 term corresponds to mutations incurred in the β_1 , β_2 , γ_1 , γ_2 region of \mathcal{T}_V .

Now, define P as follows:

$$\begin{split} P &= l_{f^{Col}}(T_V) - l_{f^{Col}}(T_E) \\ &= \left(2 + 3(3|V| + |E|) + 1 + l_{f^S}(S_V)\right) - \left(1 + 2|E| + l_{f^S}(S_E)\right) \\ &= \left(2 + 3(3|V| + |E|) + 1\right) - \left(1 + 2|E|\right) + 4M. \end{split}$$

Hence, if $\chi'(G) = 3$, $d_{MP}(T_E, T_V) \ge P$. We still need to show (1) $d_{MP}(T_E, T_V) \le P$ and (2) $d_{MP}(T_E, T_V) = P$ if and only if $\chi'(G) = 3$. Once these facts have been established NP-hardness will follow.

We approach this by starting from an arbitrary optimal character f and then transforming f step by step such that we do not lose optimality but the character attains a certain canonical form. This canonical form will be attained by accumulating one special property at a time. In all cases the argument that a new property can be obtained is based on the assumption that all earlier properties have already been accumulated. Properties are never lost, and each property can be attained in polynomial time. Thus, given an arbitrary optimal character we can transform it in polynomial time into a character that has all the described properties. Some of the proofs of the properties have been deferred to the appendix. Proofs of properties are terminated with a \blacksquare symbol.

Property 1. All cherries in \mathcal{T}_E are monochromatic. That is, if $\{x, y\}$ are two taxa that share a parent in \mathcal{T}_E , then both are assigned the same colour (i.e., state).

Proof. This is possible by combining Lemma 3.2 with the fact (established earlier) that, for every optimal character f, $l_f(\mathcal{T}_E) < l_f(\mathcal{T}_V)$.

Property 2.1. In \mathcal{T}_E , the cherry $\{\beta_1,\beta_2\}$ has a different colour to the cherry $\{\gamma_1,\gamma_2\}$.

Property 2.2. In \mathcal{T}_E , the (possibly multiple) colours used for the taxa of B (including α) are not used elsewhere in \mathcal{T}_E , except possibly $\{\beta_1, \beta_2\}$.

Property 2.3. In \mathcal{T}_E , all the taxa in B have the same colour which, with the possible exception of β_1 , β_2 , does not appear on taxa outside B and α .

Property 3. In T_E , all the taxa in B have the same colour, and cherry $\{\beta_1, \beta_2\}$ also has this colour. Moreover, this colour does not appear on any other taxa, i.e., it is unique for B (including α) and β_1, β_2 .

From now on we refer to the unique colour used by B (including α), β_1 and β_2 as pink. Property 3 is extremely important. In particular, it means that from now on we can assume the existence of optimal extensions of \mathcal{T}_V such that the root of \mathcal{T}_V is coloured pink and, moreover, that the entire image of B inside \mathcal{T}_V is coloured pink. We call these pink extensions. These greatly simplify the task of counting mutations inside \mathcal{T}_V . In particular, it means that we from now on (in \mathcal{T}_V) only need to consider mutations incurred *inside* the subtrees sibling to the taxa of B, which we call *below* pink subtrees. These subtrees never contain pink taxa.

Property 4. Let f be an optimal character with properties 1–3 and let f^* be the restriction of f to the taxa in S_V and S_E . Then $l_{f^*}(S_V) - l_{f^*}(S_E) = d_{MP}(S_E, S_V)$.

Proof. Fix a pink extension of f. From the earlier properties, f^* does not contain any pink taxa. Now, taxon α is coloured pink, because α is a taxon of B. This means that, in \mathcal{T}_V , there is unavoidably a mutation on the edge entering the root of S_V . Moreover, we know that there exist optimal characters for S_E , S_V in which the roots of S_E and S_V can be allocated the same colour blue: this is the 3-state character f^S that we constructed at the start of the proof. This means that, without loss of optimality, we can assume $f^* = f^S$, where we are free to (and should) relabel the blue inside f^S such that in \mathcal{T}_E no mutation is incurred on the edge entering the root of S_E . (This can be achieved by running the bottom-up phase of the Fitch algorithm on the subtree sibling to S_E in \mathcal{T}_E , identifying the set of colours permitted by the Fitch algorithm at the root of the subtree, and arbitrarily picking one of those colours as the relabelling colour). Optimality is assured because $(1) l_f (S_V) - l_f (S_E) = d_{MP}(S_E, S_V)$, (2) we force a mutation at the root of S_V and (3) we definitely avoid a mutation at the root of S_E .

Property 5. (a) For every edge $e = \{u, v\} \in E$ the three taxa $x_{e,1}^{edge}$, $x_{e,2}^{edge}$, $x_{e,3}^{edge}$ all have distinct colours. Moreover, (b) $x_{e,1}^{edge}$, $x_{e,1}^{u}$, $x_{e,1}^{v}$ all have the same colour, $x_{e,2}^{edge}$, $x_{e,2}^{u}$, $x_{e,2}^{v}$ all have the same colour, and finally $x_{e,3}^{edge}$, $x_{e,3}^{u}$, $x_{e,3}^{v}$ all have the same colour

Proof. (The following proof only requires Property 3, it does not use Property 4.) First, suppose for some $e \in E$ there exist $j, j' \in \{1, 2, 3\}$ such that $j \neq j'$ and $x_{e,j}^{edge}$, $x_{e,j'}^{edge}$ have the same colour. Observe that $x_{e,1}^{edge}$, $x_{e,2}^{edge}$, $x_{e,3}^{edge}$ all form a single below pink subtree in \mathcal{T}_V . Suppose we recolour $x_{e,j}^{edge}$ to some brand new colour. This raises the parsimony score of \mathcal{T}_E by at most 1. However, it also raises the parsimony score of \mathcal{T}_V by at least one, due to the introduction of a new colour into the corresponding below pink subtree. Hence, the recoloured character is optimal. We can repeat this as long as necessary to ensure that (a) eventually holds. Now, suppose for some $e \in E$ and $e \in E$

We recolour all 3 taxa with a brand new colour. This cannot raise the parsimony score of \mathcal{T}_E . On the other hand, it cannot lower the parsimony score of \mathcal{T}_V , because the three now uniquely coloured taxa all occur in different below pink subtrees of \mathcal{T}_V . (The pink taxa are critical here: without them it could happen that an optimal extension *saves* mutations by constructing monochromatic paths between some or all of the 3 taxa, causing the parsimony score of \mathcal{T}_V to actually decrease.) Hence, the recoloured character is optimal, and (a) still holds. We repeat this as long as necessary to ensure that (b) eventually also holds.

Property 6. For every $j \in \{1, 2, 3\}$, the edge colouring induced by the colours of the $x_{e, j}^{edge}$ taxa $(e \in E)$, is a proper edge colouring.

Proof. Recall that, by Property 5, each $x_{e,j}^{edge}$ taxon has the same colour as the $x_{e,j}^{u}$

and $x_{e,j}^{\nu}$ taxa below it in \mathcal{T}_E . Suppose that there is some $j \in \{1,2,3\}$ for which the induced edge colouring is not proper. Then there exists some $u \in V$ and two edges $e \neq e'$ in E incident at u such that $x_{e,j}^{u}$ and $x_{e',j}^{u}$ both have the same colour. Both these taxa are together in a below pink subtree of \mathcal{T}_V . This subtree therefore currently induces m < 2 mutations (excluding the mutation as the subtree touches the pink region). Now, suppose we introduce a brand new colour and recolour $x_{e,j}^{u}$, $x_{e,j}^{v}$, and $x_{e,j}^{edge}$ with it. This raises the parsimony score of \mathcal{T}_E by at most 1. However, it definitely also raises the parsimony score of \mathcal{T}_V , by at least 1, because the aforementioned below pink subtree now induces m+1 mutations (due to the introduction of a new colour). Hence, the new character is optimal, and all earlier properties are preserved. We can repeat this process until the induced edge colouring is proper.

Property 7. For an optimal character f,

$$l_f(\mathcal{T}_V) = 2 + 3(3|V| + |E|) + 1 + l_{fS}(S_V).$$

Proof. This is a consequence of the fact that (from Property 6) we can assume that in T_V a proper edge colouring is induced, plus the fact that a pink extension is an optimal extension. In particular, the proper edge colouring means that each of the 3|V| + |E| below pink subtrees induces 2 mutations on its internal edges and a third mutation where the subtree touches the pink region. The '2' term corresponds to the fact that the two taxa γ_1 , γ_2 are necessarily not pink. The '1' term is the mutation at the root of S_V .

Central argument.

As a consequence of Property 7, optimal characters (which we always assume to induce proper edge colourings) are only distinguished by their ability to minimize the number of mutations induced in T_E . We can already establish a strong lower bound for this number:

$$l_f(\mathcal{T}_E) \ge 1 + 2|E| + l_{f^S}(S_E).$$

Every proper edge colouring induces (at least) these mutations in \mathcal{T}_E . The '1' term is the mutation that occurs between the β_1 , β_2 and γ_1 , γ_2 taxa and the 2|E| term is a consequence of (amongst others) Property 5.

Combining this with Property 4, we see that $d_{MP}(T_E, T_V) = l_f(T_V) - l_f(T_E) \le P$, where P is the value defined earlier in the proof. We have already shown that, if G has $\chi'(G) = 3$, P is possible. We now see that this is optimal. The only thing we have left to show, is that if $\chi'(G) > 3$, that P is *not* possible. We use the contrapositive to prove this. In particular, we will show

$$l_f(\mathcal{T}_E) = 1 + 2|E| + l_{fS}(S_E) \Rightarrow \chi'(G) = 3.$$

Suppose, then, that $l_f(T_E) = 1 + 2|E| + l_{f^S}(S_E)$. This means that there are no mutations in the subtree T^{***} other than the 2|E| unavoidable mutations due to Property 5. To achieve this it must be the case that all the |E| subtrees (each containing 9 taxa) in T^{***} all have a single colour in common. Let us call this colour *blue*. Hence, for every $e \in E$, there exists exactly one $j \in \{1, 2, 3\}$ such that $x_{e,j}^{edge}, x_{e,j}^{u}, x_{e,j}^{v}$ are all

blue. We now build a proper 3-edge-colouring for G. If j=1, we assign e the colour red. If j=2, we assign e the colour blue. If j=3, we assign e the colour green. This must be a proper colouring: if it was not, then there would be some vertex $u \in V$, two incident edges e, e' incident to u, and some $j \in \{1, 2, 3\}$ such that $x_{e,j}^{edge}$ and $x_{e',j}^{edge}$ were both blue. But this would contradict Property 6. Hence, $\chi'(G)=3$.

This completes the proof. Summarising, for a given cubic graph G = (V, E),

$$\chi'(G) = 3 \Leftrightarrow d_{MP}(T_E, T_V) = P$$

from which the NP-hardness of computing $d_{MP}(T_E, T_V)$ on binary trees follows.

Corollary 4.7. For every fixed integer $i \ge 4$, computation of d_{MP}^i on binary trees is NP-hard.

Proof. This is a consequence of the fact that in the theorem only 4 states are required to construct a character achieving MP distance P. Namely, the 3 colours used in the proper edge colouring of G, plus pink.

Note that the above proof cannot (obviously) be extended to give APX-hardness. By taking multiple copies of the tree T^{***} it is possible to increase the gap between $\chi'(G)=3$ and $\chi'(G)=4$ instances to more than 1, but this is insufficient for APX-hardness.

5. Computation of d_{MP}^2 is NP-Hard on Binary Trees

As in the previous section we first require a gadget that can break symmetry between two trees.

5.1. Symmetry Breaking Gadget in the Case of 2 States

Consider the two rooted trees

$$T_a = (((5, (6, 4)), 3), ((1, (8, 2)), 7))$$

and

$$T_b = (((7, ((4, 2), 6)), 3), (8, (1, 5)))$$

shown in Figure 5.

Here, it can be verified (e.g., by exhaustive search) that $d_{MP}^2(\mathcal{T}_a, \mathcal{T}_b) = 3$, and the character $f_{asym} = AAGGGGAA$ can achieve this: $l_{f_{asym}}(\mathcal{T}_a) = 1$ and $l_{f_{asym}}(\mathcal{T}_b) = 4$.

In fact, these trees are asymmetric, in the sense that for every optimal 2-state character f, $l_f(\mathcal{T}_a) < l_f(\mathcal{T}_b)$. In particular, as can be verified by computational search (e.g., using the ILP formulation or performing an exhaustive search), $\max_f \left(l_f(\mathcal{T}_b) - l_f(\mathcal{T}_a) \right) = 3$ and $\max_f \left(l_f(\mathcal{T}_a) - l_f(\mathcal{T}_b) \right) = 2$. (The second maximum is achieved by the character AAGAAGGG, for example.) Using the same notation as in Section 4.1, but restricted to characters with at most 2 states, we therefore obtain:

$$gap(\mathcal{T}_a, \mathcal{T}_b) = 1.$$

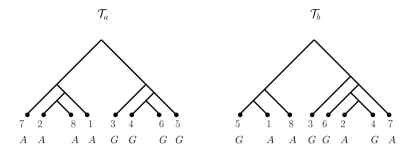


Figure 5: Two trees T_a and T_b that are "asymmetric" on characters with at most 2 states. An example of an optimal character is $f_{asym} = AAGGGGAA$.

From now on we implicitly assume that all characters have at most 2 states.

Define \mathcal{T}_A and \mathcal{T}_B in the same way as in Section 4.1. It can be verified that $gap(\mathcal{T}_A, \mathcal{T}_B) \geq 1$. This is not yet strong enough for what we require, so let \mathcal{T}_{AA} and \mathcal{T}_{BB} be obtained by joining two copies of \mathcal{T}_A , and two copies of \mathcal{T}_B , together (respectively).

Claim 5.1.
$$gap(T_{AA}, T_{BB}) \geq 2$$
.

Let \mathcal{T}_{AA}^k be the rooted tree on 32k taxa obtained by arranging k disjoint copies of \mathcal{T}_{AA} along a caterpillar backbone. That is, $\mathcal{T}_{AA}^1 = \mathcal{T}_{AA}$ and for k > 1, \mathcal{T}_{AA}^k is obtained by joining \mathcal{T}_{AA}^{k-1} and \mathcal{T}_{AA} via a new root. \mathcal{T}_{BB}^k is defined analogously.

Lemma 5.2. For
$$k \ge 1$$
, $gap(T_{AA}^k, T_{BB}^k) \ge k + 1$.

Proof. The case k = 1 is proven by Claim 5.1 and for higher k we use analogous arguments to the proof of Claim 4.2 and Lemma 4.3. We omit details.

Lemma 5.3. For
$$k \ge 1$$
, $d_{MP}^2(\mathcal{T}_{AA}^k, \mathcal{T}_{BB}^k) = 12k$.

Proof. \mathcal{T}_{AA}^k comprises 4k copies of \mathcal{T}_a . By taking 4k copies of character f_{asym} , we see that $d_{MP}^2 \left(\mathcal{T}_{AA}^k, \mathcal{T}_{BB}^k \right) \geq 4k(4-1) = 12k$. That 12k is also the upper bound, can be verified by showing $d_{rSPR} \left(\mathcal{T}_{AA}^k, \mathcal{T}_{BB}^k \right) \leq 12k$. This follows because by cutting off

all copies of taxa 2, 5, 7 into separate components, we obtain an agreement forest of \mathcal{T}_{AA}^k , \mathcal{T}_{BB}^k containing 12k+1 components.

5.2. The Reduction

We reduce from the NP-hard (and APX-hard) problem CUBIC MAX CUT [1]. Here we are given a cubic graph G = (V, E), |E| = 3|V|/2, and we are asked to partition V into two disjoint pieces $V_1 \cup V_2$ such that the number of edges that have one endpoint in V_1 and one endpoint in V_2 ("cut" edges), is maximized. Let MAXCUT(G) represent this value. We can assume, without loss of generality, that G is connected and not bipartite.

The high-level idea is similar to the 2-state hardness reduction in [5]. Namely, we will construct two trees \mathcal{T}_V and \mathcal{T}_E and apply the symmetry-breaking gadget to ensure that for all optimal characters f, $l_f(\mathcal{T}_V) < l_f(\mathcal{T}_E)$. We will model the vertices as subtrees in \mathcal{T}_V , each comprising three taxa, and argue — via a technical argument — that these subtrees are monochromatic. We will let the 2 states represent the two sides of the chosen partition $V_1 \cup V_2$. Henceforth, we will call these states red and blue. The colour of a vertex subtree thus denotes which side of the partition it is on. The tree \mathcal{T}_E will be constructed such that, the more cut edges are induced by the partition chosen by \mathcal{T}_V , the higher the parsimony score of \mathcal{T}_E . The construction will thus naturally choose a character that maximizes MAXCUT(G).

The fact that \mathcal{T}_V and \mathcal{T}_E must be binary, introduces significant complications compared to the 2-state hardness reduction in [5]. For this reason we will introduce two new special gadgets, that allow \mathcal{T}_V (respectively, \mathcal{T}_E) to be viewed as the independent union of several subtrees. In \mathcal{T}_V the gadget will be called the *cherry switch* and in \mathcal{T}_E we will have the $D(w_i)$ gadget, to be explained in due course. These independence gadgets neutralise the influence of side-effects that can occur as a consequence of the fact that \mathcal{T}_V and \mathcal{T}_E are both binary.

We begin by constructing \mathcal{T}_E . First, we construct the left-hand side subtree \mathcal{T} of \mathcal{T}_E as depicted in Figure 6. Let $\tilde{\mathcal{T}}$ be an arbitrary rooted binary tree on |E|+1 leaves $\{m_1,\ldots,m_{|E|+1}\}$. Let $I=\{w_1,\ldots,w_{|E|}\}$ be the |E| interior nodes of $\tilde{\mathcal{T}}$. Let M be a large integer whose value we will determine in due course. Let S_V be the tree \mathcal{T}_{AA}^M and S_E be the tree \mathcal{T}_{BB}^M . Let I be an arbitrary leaf of $\tilde{\mathcal{T}}$. We replace I with S_E . Next, select an arbitrary bijection between the remaining leaves of $\tilde{\mathcal{T}}$ and E. For each edge $e=\{u,v\}\in E$, replace the leaf of $\tilde{\mathcal{T}}$ corresponding to e with a cherry on two taxa $\{x_e[u],x_e[v]\}$. Now, for each internal vertex w_i , let p_i and q_i be the two children of w_i . We now introduce the independence gadget $D(w_i)$, constructed as follows. Take a rooted binary tree $(w_i^2,((w_i^0,w_i^4),(w_i^3,(w_i^5,w_i^1))))$. We replace w_i with this tree, in the following sense: delete w_i , identify w_i^0 with p_i , identify w_i^1 with q_i and if w_i had an incoming edge, identify the root of $D(w_i)$ with the head of this edge. The remaining leaves of $D(w_i)$ are $\{w_i^2, w_i^3, w_i^4, w_i^5\}$ and we regard these as taxa, so replacing each w_i with $D(w_i)$ increases the number of taxa in total by 4|E|.

 \mathcal{T} has in total 2|E|+32M+4|E| taxa, where the 32M is the number of taxa in S_E . Let \mathcal{T}' be a rooted caterpillar on |V|+3|E| leaves $\{l_1,\ldots,l_{|V|+3|E|}\}$. Replace each leaf l_i by a "double cherry" $((\alpha_i,\gamma_i),(\beta_i,\delta_i))$ where $\{\alpha_i,\beta_i,\gamma_i,\delta_i\}$ are taxa. Join \mathcal{T} and \mathcal{T}' together by a new root: this completes the construction of \mathcal{T}_E as depicted in

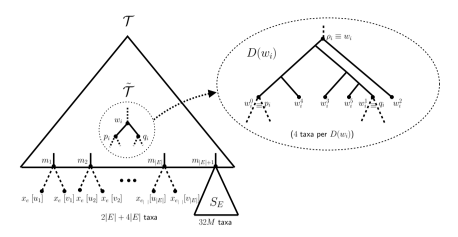


Figure 6: Tree \mathcal{T} is the left-hand side subtree of \mathcal{T}_E in the 2-state NP-hardness construction, cf. Figure 7. Every internal node w_i with children nodes p_i and q_i of the original tree \tilde{T} is replaced by tree $D(w_i)$ with root w_i and children w_i^0, \ldots, w_i^5 . Children w_i^0 and w_i^1 correspond to p_i and q_i , respectively, whereas the other children form new leaves. Therefore, each $D(w_i)$ contributes four leaves to tree \mathcal{T} . For the leaves labelled $x_{e_i}[u_i]$ and $x_{e_i}[v_i]$, u_i and v_i are the endpoints of edge e_i .

Figure 7. \mathcal{T}_E thus has in total,

$$2|E| + 32M + 4|E| + 4(|V| + 3|E|) = 32M + 18|E| + 4|V|$$

taxa.

To construct T_V we start by creating a set of taxa-disjoint trees J. The disjoint union of the taxa in the |V| + 1 + 3|E| trees in J will be exactly the set of taxa in the tree T depicted in Figure 6. J contains,

- (1) S_V ;
- (2) for each vertex $u \in V$, a rooted triplet $(x_e[u], (x_{e^*}[u], x_{e^{**}}[u]))$ where e, e^*, e^{**} are the three edges incident to u in G;
- (3) for each gadget $D(w_i)$, two single taxon trees w_i^4 and w_i^5 , and one cherry (w_i^2, w_i^3) .

Let C be a rooted caterpillar on |V|+1+3|E| leaves. Consider a directed path on |V|+3|E| edges that starts at the root of C and terminates at one of the leaves in the unique cherry of C. Let K be the edges in this path. Choose an arbitrary bijection between the leaves of C and the trees in J, and replace each leaf with its corresponding subtree. We now need to replace each edge in K with a special gadget. In particular, select an arbitrary bijection between K and $\{1,\ldots,|V|+3|E|\}$. Next, for each edge in K, subdivide it twice. From one of the vertices created by the subdivision operation, hang a cherry (α_i, β_i) , and from the other hang a cherry (γ_i, δ_i) , where i is the index given by the bijection. We call these two cherries a cherry switch — this is the

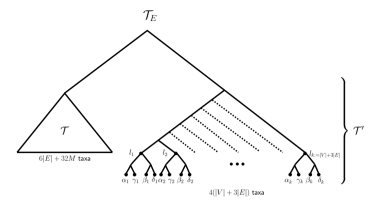


Figure 7: Tree \mathcal{T}_E for the 2-state NP-hardness construction consists of \mathcal{T} as depicted in Figure 6 on the left-hand side and \mathcal{T}' on the right-hand side. Note that \mathcal{T}_E employs in total 32M + 6|E| + 4(|V| + 3|E|) = 32M + 18|E| + 4|V| taxa.

independence gadget for \mathcal{T}_V . This completes the construction of \mathcal{T}_V , which is depicted in Figure 8.

We are now in a position to specify the number M. We require M to be sufficiently large that, for every optimal character f, $l_f(\mathcal{T}_V) < l_f(\mathcal{T}_E)$. From Lemma 5.3 we know that there exists some character f' such that $l_{f'}(\mathcal{T}_E) - l_{f'}(\mathcal{T}_V) \ge 12M$. We can obtain such a character by extending the character suggested by Lemma 5.3 such that all taxa outside S_E and S_V are assigned the same state. Now, let t be the number of edges in \mathcal{T}_V that lie outside S_V . For every character f we have

$$l_f(\mathcal{T}_V) - l_f(\mathcal{T}_E) \le t + (12M - (M+1)).$$

The 12M term is obtained from Lemma 5.3, the (M+1) term from Lemma 5.2, and the t term arises (pessimistically) from the situation when every edge in \mathcal{T}_V (outside S_V) incurs a mutation, but no edge in \mathcal{T}_E (outside S_E) incurs a mutation. So, if we choose M such that

$$t + 12M - (M+1) < 12M,$$

it follows that for *every* optimal character f, $d_{MP}(\mathcal{T}_V, \mathcal{T}_E) = l_f(\mathcal{T}_E) - l_f(\mathcal{T}_V)$ and in particular $l_f(\mathcal{T}_E) > l_f(\mathcal{T}_V)$. Choosing M = t is therefore sufficient to achieve this. Consider now the following.

Observation 5.4. Let T_1 , T_2 be two binary trees and let f be an optimal character such that $l_f(T_1) < l_f(T_2)$. Suppose T_1 contains two cherries (a,b) and (c,d) and, in T_2 , there are cherries (a,c) and (b,d) under a common parent (i.e., a "double cherry"). Then f can be modified to obtain an optimal character f'' in which (a,b) and (c,d) are both monochromatic but with different colours, and the colours of all other taxa are unchanged.

Proof. Let f be an optimal character. We first apply Lemma 3.2 to obtain an optimal character f' in which the two cherries are monochromatic in \mathcal{T}_1 . If the two cherries

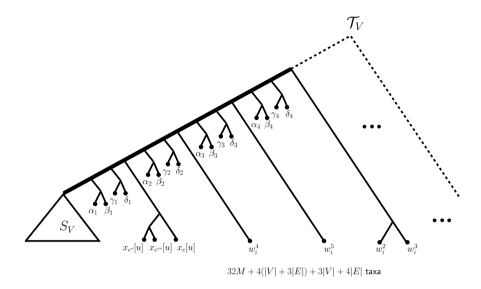


Figure 8: \mathcal{T}_V consists of a modification of a caterpillar tree with directed path K which starts at the root and leads to a leaf in a cherry. Path K is depicted in bold. Each of the |V|+3|E| edges in K carries two additional cherries (α_i, β_i) and (γ_i, δ_i) . Therefore, K contributes in total 4(|V|+3|E|) taxa. Moreover, for each of the original |E| inner nodes w_i of tree \mathcal{T} as depicted in Figure 6, \mathcal{T}_V contains four taxa w_i^2 , w_i^3 , w_i^4 , w_i^5 . This leads to 4|E| more taxa. Finally, for each vertex u in V, \mathcal{T}_V contains a triple $(x_e[u], (x_{e^*}[u], x_{e^{**}}[u]))$, which are 3|V| taxa. Using the fact that in cubic graphs we have $|V| = \frac{2}{3}|E|$, \mathcal{T}_V employs in total 32M+16|E|+7|V| = 32M+18|E|+4|V| taxa.

have different colours we are done. If not, then recolour one of the cherries to obtain f''. This raises the parsimony score of \mathcal{T}_1 by (at most) one. In \mathcal{T}_2 two new mutations are created in the cherries (a, c) and (b, d) while at most one mutation is saved on the edge entering the common parent. Hence, f'' is also optimal.

In exactly the same way as Theorem 4.6 we now give an accumulating list of properties which can be shown to be enjoyed by at least one optimal character that can be constructed in polynomial time.

Property 1. In T_V , for each $i \in |V| + 3|E|$, cherry (α_i, β_i) is monochromatic and cherry (γ_i, δ_i) is monochromatic, and the cherries have different colours.

Proof. This is an immediate consequence of Observation 5.4.

Next, observe that if a character f has Property 1, and we swap the colours used in some (or all) of the cherry switches to obtain f', then $l_f(\mathcal{T}_E) = l_{f'}(\mathcal{T}_E)$. This is because each cherry switch in \mathcal{T}_V corresponds to a double cherry in \mathcal{T}_E , and (as long as Property 1 already holds) the behaviour of the double cherries is invariant under permutation of red and blue. This is the key observation behind the next property.

Property 2. There is an optimal character f such that

$$l_f(\mathcal{T}_V) = |V| + 3|E| + \sum_{T^* \in J} l_f(T^*)$$

where $l_f(\mathcal{T}^*)$ has the expected meaning i.e., the parsimony score of \mathcal{T}^* after restricting f to the taxa in \mathcal{T}^* .

Proof. Observe that for any optimal character f, $|V| + 3|E| + \sum_{T^* \in I} l_f(T^*)$ is a lower bound on $l_f(\mathcal{T}_V)$. This can be observed by first applying Fitch's algorithm to the trees in J (which are all pendant in T_V) and then noting that, due to Property 1, each of the |V| + 3|E| cherry switches also incurs a mutation, irrespective of the states that Fitch's algorithm designates to the roots of the trees in J. To show that it is also an upper bound, first run the Fitch algorithm on the trees in J. For those trees in J that are allowed by the Fitch algorithm to have either colour at the root, pick one arbitrarily. For each cherry switch, consider the root state of the tree from J directly above it (where here "above" means: closer to the root of \mathcal{T}_V), and directly below it. There are four possibilities: red-blue (i.e., the tree from J above it requires a root state of red, the tree below it wants blue), red-red, blue-red, and blue-blue. If it is red-blue, then if necessary swap the colours on the two cherries in the cherry switch, to ensure that the red cherry is closer to the root of T_V . If it is blue-red, then ensure that the blue cherry is closer to the root of T_V . Now, irrespective of which of the four possibilities holds, there is an optimal extension which occurs exactly one mutation (and not more) per cherry switch. In the red-blue and blue-red cases the mutation will be on the edge between the two subdivision vertices (i.e., the edge between the vertices at which the two cherries are attached to the caterpillar backbone). In the cases red-red and blueblue the mutation will be on the edge feeding into the blue, respectively red cherry.

Property 3. In \mathcal{T}_V , the trees in J that are rooted triplets or cherries, are all monochromatic.

Proof. That the cherries can be made monochromatic, is simply a consequence of Lemma 3.2. That the rooted triplets are monochromatic is more subtle. Consider any triplet in J, this has the form $(x_e[u], (x_{e^*}[u], x_{e^{**}}[u]))$. We already know that $\{x_{e^*}[u], x_{e^{**}}[u]\}$ have the same colour, as they form a cherry. Now, if $x_e[u]$ also has this colour, we are done. If not, then recolour it to give it the same colour as the other two taxa. By Property 2, this *must* lower the parsimony score of \mathcal{T}_V by exactly one. Hence, the new character is also optimal. (We really need Property 2 here, since "the parsimony score of \mathcal{T}_V does not increase" — which in general is the strongest statement we can make after such a recolouring — is not strong enough for our purposes).

Property 3 basically says that, in T_V , the three taxa that represent each vertex of G all have the same colour. This will allow us to encode MAX CUT correctly. Property 3 also tells us that the $\{w_i^2, w_i^3\}$ pairs of taxa, which form part of the $D(w_i)$ gadget, will be monochromatic. This is particularly useful when combined with the fact that w_i^4 and w_i^5 are both single taxa trees in J. A tree comprising only a single taxon has

parsimony score 0, so whichever colour is allocated to the w_i^4 and w_i^5 taxa, they do not impact upon the parsimony score of \mathcal{T}_V , by Property 2. In other words, these two taxa are "free": they can be allocated any colour in an attempt to cause as many mutations as possible in \mathcal{T}_E . The $\{w_i^2, w_i^3\}$ pairs of taxa are also "free", except for the limitation that w_i^2 and w_i^3 should have the same colour. This underpins the following critical

Observation 5.5. Consider the rooted binary tree

$$D(w_i) = \left(w_i^2, \left(\left(w_i^0, w_i^4\right), \left(w_i^3, \left(w_i^5, w_i^1\right)\right)\right)\right).$$

Suppose we fix w_i^0 as red, or blue, or $\{red, blue\}$, where $\{red, blue\}$ has the same meaning as in Fitch's algorithm, i.e., "both states are possible". Suppose we do the same (independently) for w_i^1 . Then depending on our choice we can always select colours for w_i^2 , w_i^3 , w_i^4 , w_i^5 , whilst ensuring that the same colour is chosen for w_i^2 and w_i^3 , such that the parsimony score of $D(w_i)$ under the resulting character is at least 2. Moreover, it is never possible to achieve a parsimony score higher than 2 in this way.

Proof. A straightforward case-analysis is sufficient to verify the "at least 2" part of the claim. There are 3² cases, several of which are symmetrical. These are the relevant cases:

- (1) w_i⁰ and w_i¹ are both red. Then choose all other taxa to be blue.
 (2) w_i⁰ is red and w_i¹ is blue. Then choose w_i⁴ to be blue, w_i⁵ to be red, and w_i² and w_i³ to both be blue.
- (3) w_i^0 is red and w_i^1 is $\{red, blue\}$. Then choose w_i^4 to be blue, w_i^5 to be blue, and
- (3) w_i is feed and w_i to feed, some y w_i² and w_i³ to both be red.
 (4) w_i⁰ is {red, blue} and w_i¹ is red. Then choose w_i⁴ to be red, w_i⁵ to be blue, and w_i² and w_i³ to both be blue.
 (5) w_i⁰ and w_i¹ are both {red, blue}. Then choose w_i⁴ to be red, w_i⁵ to be red, and w_i²
- and w_i^3 to both be blue.

To show that 3 or more mutations are never possible, note that a character on 6 taxa can only possibly have a parsimony score of 3 if there are exactly 3 red taxa and exactly 3 blue taxa. (Otherwise, simply choose an extension that assigns the majority colour to all internal nodes of the tree, yielding at most 2 mutations.) Now, if at least one of w_i^0 and w_i^1 chooses $\{red, blue\}$, then 3 mutations are certainly not possible, because we can (again) colour all the internal nodes of the tree monochromatic in the majority colour, yielding at most 2 mutations. So, suppose without loss of generality w_i^2 and w_i^3 are both red. Then exactly one of w_i^0 and w_i^4 will be red, and the other blue. But then w_i^1 and w_i^5 will both be blue. But this character has parsimony score at most 2, contradiction.

In \mathcal{T}_E the taxa w_i^0 and w_i^1 become the roots of subtrees, and the three possible choices for each taxon in Observation 5.5 reflect the three possible decisions that Fitch's algorithm can make when, in the bottom-up phase, the root of that subtree is reached. Essentially, then, Observation 5.5 allows us to "glue" these two subtrees together with a profit of exactly 2 mutations, entirely independently of the two subtrees themselves.

Now, consider any optimal character f that has Property 3 (and thus all earlier properties too). We have

$$l_f(\mathcal{T}_V) = |V| + 3|E| + l_f(S_V)$$

since (by Property 2) the singletons, cherries and triplets in J do not internally generate any mutations and mutations along the K part of \mathcal{T}_V are already accounted for. (As usual, $l_f(S_V)$ refers to the parsimony score of the restriction of f to the taxa in S_V). Let CUT(f) be the number of cut edges induced by f, i.e., after partitioning the vertices of V according to the colours of the corresponding rooted triplets in J. We have,

$$l_f(\mathcal{T}_E) = 2(|V| + 3|E|) + l_f(S_E) + CUT(f) + 2|E|.$$

The 2(|V|+3|E|) term is the contribution of the double cherries, and the 2|E| term is the 2 mutations that we know we can definitely incur in each $D(w_i)$ gadget. Hence, an optimal character should try and make the induced cut as large as possible: there is no other freedom. Consequently,

$$\begin{split} d_{MP}^2(T_V, T_E) &= l_f(T_E) - l_f(T_V) \\ &= 2(|V| + 3|E|) + l_f(S_E) + MAXCUT(G) + 2|E| \\ &- (|V| + 3|E| + l_f(S_V)) \\ &= |V| + 5|E| + (l_f(S_E) - l_f(S_V)) + MAXCUT(G) \\ &= |V| + 5|E| + 12M + MAXCUT(G). \end{split}$$

The fact that $(l_f(S_E) - l_f(S_V))$ is equal to 12M is not entirely automatic. It is a consequence of the fact that in this context there is no point choosing a character f which, when restricted to S_V and S_E , yields an MP distance smaller than $d_{MP}^2(S_V, S_E)$ (where the latter value is equal to 12M by Lemma 5.3).

The terms can easily be rearranged to obtain MAXCUT(G) from d_{MP}^2 , which yields the overall theorem:

Theorem 5.6. Computation of d_{MP}^2 is NP-hard on binary trees.

We also obtain the following corollary.

Corollary 5.7. Computation of d_{MP}^2 is APX-hard on binary trees.

Proof. We will show that if d_{MP}^2 can be approximated in polynomial time to within a multiplicative factor of $(1 - \varepsilon)$, for some $\varepsilon > 0$, that CUBIC MAXCUT can be approximated in polynomial time to within a factor of $(1 - k\varepsilon)$ for some constant k > 0 that is independent of ε . Given that CUBIC MAX CUT is APX-hard [1] there is (by definition) some $\varepsilon' > 0$ such that a factor $(1 - \varepsilon')$ approximation or better is not possible in polynomial time unless P = NP. The APX-hardness of d_{MP}^2 will then follow[‡]: the corresponding threshold for d_{MP}^2 will be ε'/k .

[‡] Formally speaking, we should give an L-reduction here [15]. For brevity we omit the technicalities. An L-reduction can if desired easily be constructed from the information provided here.

First, suppose we obtain character f, which is a $(1-\varepsilon)$ approximation to $d_{MP}^2(\mathcal{T}_V,\mathcal{T}_E)$. We need to show that a feasible solution (i.e., a cut) can be extracted in polynomial time from f, which requires that the solution obeys all the Properties. Character f might not have these Properties, but they can be acquired in polynomial time without lowering the parsimony distance score of the character. To do this, ensure first that S_V and S_E use the duplicated character f_{asym} (which optimizes the MP distance between S_V and S_E). This ensures that $l_f(\mathcal{T}_V) < l_f(\mathcal{T}_E)$. From this point on the Properties can be accumulated one at a time: the constructive proofs describing how the Properties are obtained do not require that f is optimal, only that $l_f(\mathcal{T}_V) < l_f(\mathcal{T}_E)$.

Recall that |V| = (2/3)|E|. We need an explicit expression for M. This was set to be t, the number of edges in \mathcal{T}_V minus the edges in subtree S_V . \mathcal{T}_V has in total 32M + 18|E| + 4|V| taxa, and after subtracting the 32M this gives 18|E| + 4|V|. A rooted binary tree on |X| taxa has 2(|X|-1) edges, yielding 36|E| + 8|V| - 2, plus 2 extra edges created when the subtree S_V is re-attached, giving 36|E| + 8|V| which is (124/3)|E|. Hence,

$$\begin{split} d_{MP}^2(T_V, T_E) &= |V| + 5|E| + 12M + MAXCUT(G) \\ &= (2/3)|E| + 5|E| + 496|E| + MAXCUT(G) \\ &= (1505/3)|E| + MAXCUT(G). \end{split}$$

The size of the cut returned after processing f is at least

$$= (1 - \varepsilon)((1505/3)|E| + MAXCUT(G)) - (1505/3)|E|$$
$$= (1 - \varepsilon)MAXCUT(G) - \varepsilon(1505/3)|E|.$$

It is well known that for cubic G, $MAXCUT(G) \ge 2|E|/3$, by moving a vertex to the other side of the partition if one or fewer of its incident edges is in the cut. So,

$$\begin{split} &(1-\varepsilon)MAXCUT(G)-\varepsilon(1505/3)|E|\\ &\geq (1-\varepsilon)MAXCUT(G)-\varepsilon(1505/2)MAXCUT(G)\\ &= \left(1-\frac{1507}{2}\varepsilon\right)MAXCUT(G). \end{split}$$

This concludes the proof.

6. An Integer Linear Programming (ILP) Formulation for Binary Instances

Let \mathcal{T}_1 and \mathcal{T}_2 be two binary phylogenetic trees on $n \geq 2$ taxa. Given the hardness of MP distance it is natural to ask how well $d_{MP}(\mathcal{T}_1, \mathcal{T}_2)$ can be computed in practice. One option is to leverage the result in [5] which proves that there always exists an optimal character that is convex on one of the trees (i.e., has a parsimony score exactly one less than the number of states in the character). Hence, we can guess which of the two input trees is convex, guess the number of states s in the optimal character,

and then guess the (s-1) edges of the convex tree on which the mutations occur. Assuming the trees are unrooted, and letting $g(\mathcal{T}_1, \mathcal{T}_2)$ be any safe upper bound on s, this gives a deterministic running time of

$$O\left(\sum_{s=2}^{g(\mathcal{T}_1,\mathcal{T}_2)} {2n-3 \choose s-1}\right).$$

As the following observation shows, we can take $g(T_1, T_2) = \lfloor n/2 \rfloor$.

Observation 6.1. Let \mathcal{T}_1 and \mathcal{T}_2 be two binary phylogenetic trees on $n \ge 2$ taxa. There exists an optimal convex character with at most $\lfloor n/2 \rfloor$ states. Moreover, this bound is tight.

Proof. Let f be an optimal convex character. Suppose f has strictly more than $\lfloor n/2 \rfloor$ states. Then there exists a state t that occurs on only one taxon x. We root \mathcal{T}_1 on the edge entering x. If we run the Fitch algorithm on this rooted tree a union event will necessarily be generated at the root due to the fact that t occurs on only one taxon. Let C be the set of states in this union event, and let t' be any state in $C \setminus \{t\}$. Let f' be the character obtained from f by assigning state t' to taxon x. By re-running the Fitch algorithm we see that $l_{f'}(\mathcal{T}_1) = l_f(\mathcal{T}_1) - 1$. Moreover, f' has one fewer state than f, so f' is convex. By Observation $3.1 \ l_{f'}(\mathcal{T}_2) \ge l_f(\mathcal{T}_2) - 1$. Hence, f' is optimal, convex and has fewer states than f. By repeating this process we eventually obtain an optimal convex character with at most $\lfloor n/2 \rfloor$ states.

The trees in Figure 1 on 6 taxa are a tight example for this bound: it can easily be verified computationally that for these two trees optimal characters require at least 3 states.

Of course, even if we take $g(T_1, T_2) = \lfloor n/2 \rfloor$, such brute-force algorithms will quickly become impractical for even very small n. Hence, we turn to Integer Linear Programming (ILP), which allows us to compute d_{MP} and d_{MP}^i for larger trees. The ILP for computing d_{MP}^2 performs very well, allowing computation of d_{MP}^2 in reasonable time for trees with up to 100 taxa. Unfortunately, in the case of d_{MP} the ILP struggles to terminate in reasonable time for trees with more than 16 taxa. Future research (i.e., better ILP formulations) will hopefully improve upon this.

The ILP formulation is currently limited to binary trees but the model could be extended to non-binary trees without too much difficulty.

Let \mathcal{T}_1 and \mathcal{T}_2 be rooted, binary phylogenetic trees on the same set of taxa X, where |X| = n. Let U be the internal nodes of \mathcal{T}_1 and V the internal nodes of \mathcal{T}_2 . Let s be a constant denoting the maximum number of states that any character can have; as discussed taking $s = \lfloor n/2 \rfloor$ is a safe choice. (To compute d_{MP}^i we simply take $s \le i$.)

The following ILP maximizes $l_f(\mathcal{T}_1) - l_f(\mathcal{T}_2)$ ranging over all characters f with at most s states. To obtain the true parsimony distance the ILP should be run twice, once to compute the maximum of $l_f(\mathcal{T}_1) - l_f(\mathcal{T}_2)$ and once to compute the maximum of $l_f(\mathcal{T}_2) - l_f(\mathcal{T}_1)$.

All variables in the program are binary.

First of all we constrain that in both trees the taxa have the same state, and that each taxon chooses exactly one state. We introduce variables $x_{t,i}$ for each $t \in X$ and

 $1 \le i \le s$. For each $t \in X$ we introduce the constraint:

$$\sum_{i=1}^{s} x_{t,i} = 1.$$

We now show how the parsimony score can be computed for T_1 . The variables and constraints essentially "hard-code" Fitch's algorithm. (The encoding of T_2 is symmetrical. The two encodings are linked together via the variables that represent the states of the taxa in X, and the objective function, which we shall discuss in due course).

Given an internal node $u \in U$, let l be its left child and r its right child. Fitch's algorithm tells us to take the intersection of the states at l and r, if the intersection is non-empty, and otherwise the union (in which case we pay 1 mutation). We do this computation as follows. For $1 \le i \le s$ we introduce a variable $x_{u,i}$. The idea is that $x_{u,i}$ will be 1 if and only if state i is in the set of states at node u (in the bottom-up phase of the Fitch algorithm). We determine the set of states at u by performing the union and intersection computations directly. For that purpose, for $1 \le i \le s$ we introduce $x_{u,i}^{\cap}$ and $x_{u,i}^{\cup}$ and the following constraints:

$$x_{u,i}^{\cap} \leq x_{l,i},$$
 $x_{u,i}^{\cap} \leq x_{r,i},$
 $x_{u,i}^{\cap} \geq x_{l,i} + x_{r,i} - 1,$
 $x_{u,i}^{\cup} \geq x_{l,i},$
 $x_{u,i}^{\cup} \geq x_{r,i},$
 $x_{u,i}^{\cup} \leq x_{l,i} + x_{r,i}.$

The top group of constraints ensure that the $x_{u,i}^{\cap}$ variables reflect the intersection of the states at the children (i.e., logical AND) and $x_{u,i}^{\cup}$ the union (i.e., logical OR).

For each $u \in U$ we have variables x_u^{\cap} and x_u^{\cup} which are 1 (0, respectively) if the Fitch algorithm assigns an intersection operation to node u. We can ensure that these variables take the correct value as follows. Firstly:

$$\sum_{i=1}^{s} x_{u,i}^{\cap} \ge x_{u}^{\cap}.$$

And, secondly, we add the following constraint for each $1 \le i \le s$:

$$x_u^{\cap} \geq x_{u,i}^{\cap}$$
.

To ensure that x_u^{\cap} and x_u^{\cup} are complementary we add the constraint

$$x_{u}^{\cap} + x_{u}^{\cup} = 1.$$

Now, we have to ensure that $x_{u,i}$ takes the value $x_{u,i}^{\cap}$ whenever x_u^{\cap} is 1, and $x_{u,i}^{\cup}$ otherwise. We do this by, for each $1 \le i \le s$, adding the following four constraints:

$$x_{u,i} \geq x_{u,i}^{\cap}$$

$$x_{u,i} \le x_{u,i}^{\cup},$$

$$x_{u,i} \le x_{u,i}^{\cap} + 1 - x_{u}^{\cap},$$

$$x_{u,i} \ge x_{u,i}^{\cup} - x_{u}^{\cap}.$$

Finally, all that remains is to compute the difference between the two parsimony scores. We do this with the following objective function:

$$\text{Maximize } \sum_{u \in U} x_u^{\cup} - \sum_{v \in V} x_v^{\cup}.$$

This concludes the formulation. We have implemented it by using Java to translate the input trees into an ILP format suitable for solvers such as GLPK, SCIP, or CPLEX. We have used this to verify several of the bounds used in Section 4.1. The source code can be downloaded from [13].

We tested our ILP running CPLEX on a 3.10GHz 64-bit machine with 4 GB RAM.

We observed the following running times. For computation of d_{MP} :

- the two trees \mathcal{T}_a and \mathcal{T}_b on 6 taxa as depicted in Figure 1: total running time < 1 second.
- the two trees \mathcal{T}_A and \mathcal{T}_B on 12 taxa consisting of two copies of \mathcal{T}_a or \mathcal{T}_b , respectively: 70 seconds.

For computation of d_{MP}^2 :

- the two trees on 8 taxa as depicted in Figure 5: < 1 second.
- the two trees T_A and T_B on 16 taxa consisting of two copies of T_a or T_b , respectively: < 1 second.
- the two trees T_{AA} and T_{BB} on 32 taxa consisting of four copies of T_a or T_b : 6 seconds.

Computation of d_{MP}^i , for small i, is much faster than d_{MP} due to the greatly reduced number of binary variables. We observed that the ILP could compute d_{MP}^2 for trees with 100 taxa in approximately 140 seconds.

7. Conclusion

In this article we have proven that calculating MP distance (d_{MP}) is NP-hard on binary trees. Computation of d_{MP}^2 (the version of the problem where we are restricted to binary characters) is also NP-hard on binary trees. The latter problem is also APX-hard, and determining whether d_{MP} is APX-hard remains an open question. At the moment we do not have an NP-hardness proof for d_{MP}^3 on binary trees but given that d_{MP}^i on binary trees is NP-hard for each $i \ge 4$ we expect that this will also be hard.

We have presented and implemented a simple ILP formulation, which is publicly available at [13]. The ILP is much faster than obvious brute-force algorithms and allowed us to verify the MP-distance of the symmetry-breaking gadgets used in the hardness reductions. The ILP for d_{MP}^2 is fast but the ILP for d_{MP} does not scale well.

An important open problem is therefore to develop an ILP formulation that avoids the present approach of simply hard-coding Fitch's algorithm.

Finally, elucidating the exact relationship between MP distance and other phylogenetic metrics remains an intriguing challenge.

8. Appendix

Here, we present the proofs we omitted from Section 4.2.

Property 2.1. In \mathcal{T}_E , the cherry $\{\beta_1, \beta_2\}$ has a different colour to the cherry $\{\gamma_1, \gamma_2\}$.

Proof. Suppose this is not so. Recolour $\{\beta_1, \beta_2\}$ to some new colour not appearing elsewhere. This increases the number of mutations in \mathcal{T}_E by at most 1. However, in \mathcal{T}_V the number of mutations in the $\beta_1, \beta_2, \gamma_1, \gamma_2$ subtree increases from 0 to 2. Possibly \mathcal{T}_V then saves a single mutation at the root, but in any case the parsimony score of \mathcal{T}_V increases by at least 1. So the new character is still optimal.

Property 2.2. In \mathcal{T}_E , the (possibly multiple) colours used for the taxa of B (including α) are not used elsewhere in \mathcal{T}_E , except possibly $\{\beta_1, \beta_2\}$.

Proof. Take an optimal extension F of f by applying Fitch's algorithm. Let c be the colour allocated to the root of B by this extension. Let c^* be the colour of the parent p_1 of the root of B, and c^{**} the colour of its parent p_2 . Let c_β be the colour of the $\{\beta_1, \beta_2\}$ taxa and define c_γ similarly.

There are two cases to consider. If it is *not* true that $c = c^* = c^{**}$, then we are in the "easier" of the two cases, and proceed as follows. We recolour all the monochromatic connected components induced by the extension, and starting at some vertex of B, with brand new colours. This new character must be optimal. (The score of T_V under this new character does not decrease, so the recoloured extension must also be optimal.) Moreover, with the possible exception of β_1 , β_2 none of the colours used for taxa in B are used outside B. This is guaranteed because in this case there must be a mutation between the root of B and D_1 and/or between D_1 and D_2 . In particular, this prohibits the existence of monochromatic connected components that connect taxa in D with taxa beyond $D = \{B_1, B_2\}$. This ensures that the property holds.

The second case is much more challenging: $c = c^* = c^{**}$. We will recolour the character — and this extension — to ensure that this is no longer the case. By Property 2.1, $c_{\beta} \neq c_{\gamma}$. If $c \neq c_{\beta}$ and $c \neq c_{\gamma}$, then recolour γ_1 , γ_2 and their parent to colour c. (This lowers the parsimony score of T_E by 1, and can lower the parsimony score of T_V by at most 1, so the character — and the extension — is still optimal.) Otherwise, exactly one of c_{β} and c_{γ} is equal to c. If c_{β} has this property, then swap the colours of $\{\beta_1, \beta_2\}$ and $\{\gamma_1, \gamma_2\}$ (and their parents). So we now have $c = c_{\gamma}$ and $c \neq c_{\beta}$. In particular, there is a mutation on the edge entering the cherry $\{\beta_1, \beta_2\}$. For technical reasons we now introduce a brand new colour, *bronze* say, and recolour $\{\beta_1, \beta_2\}$ (and their parent) to be bronze. This leaves the parsimony score of T_E unchanged, and cannot decrease the parsimony score of T_V , so the character is still optimal. We do this simply to ensure that the colour of β_1, β_2 does not occur anywhere else. Run Fitch's algorithm on T_V and record the output as R.

At this point we introduce a new colour silver. Recolour the following vertices silver: β_1 , β_2 , their parent, p_1 , and the entire c-coloured connected component inside B starting at the root of B. This gives a new character and extension which saves one mutation (on the edge leading into the cherry β_1 , β_2) but creates one mutation between p_1 and p_2 . So the parsimony score of \mathcal{T}_E does not increase. It is not obvious, but the parsimony score of T_V will not drop. To see why this is, note that (under this particular recolouring) the only way the parsimony score of T_V could drop, is if the recolouring causes a mutation (i.e., union event) at the root of T_V to vanish, and at the same time does not create any additional mutations elsewhere. If R did not have a mutation event at the root of T_V anyway we are done, there is nothing to consider. If it did, then in R the union event at the root must have had the form $\{c, bronze\} \cup W$ where $W \cap \{c, bronze\} = \emptyset$ and the W is the set of states generated by the bottom-up phase of Fitch's algorithm for the root-incident right subtree of T_V , let us call this T_{right} . Now, if the recolouring causes the parsimony score of T_{right} to increase, we are also done. So suppose the parsimony score of T_{right} stays the same and T_{right} suddenly has an optimal extension (generated by any method, not necessarily the Fitch algorithm) in which its root can be coloured c or silver (which is necessary to save a mutation at the root of T_V). But then we could take this extension and remerge the colours c and silver back into c, showing that T_{right} did originally have an optimal extension in which its root could be coloured c. This would mean that R cannot possibly have been an optimal extension: it claimed a mutation was needed at the root of T_V , but we have just shown that colouring the root c would have avoided mutations on both of its outgoing edges. Contradiction to the assumed optimality of

Hence, this new character is indeed still optimal. The modified extension (on \mathcal{T}_E) is necessarily also optimal for this new character: if some other extension existed that induced fewer mutations, then this would violate the assumed optimality of the original character (i.e., because the parsimony score of \mathcal{T}_V does not decrease).

At this point we can recolour all the monochromatic connected components induced by the extension, and starting at some vertex of B, with brand new colours. This new character must be optimal. (The score of \mathcal{T}_V under this new character does not decrease, so the recoloured extension must also be optimal.) Moreover, with the possible exception of β_1 , β_2 none of the colours used for taxa in B are used outside B. This is guaranteed because the silver recolouring ensured that there are no longer monochromatic connected components that connect taxa in B with taxa beyond $B \cup \{\beta_1, \beta_2\}$.

Property 2.3 In T_E , all the taxa in B have the same colour which, with the possible exception of β_1 , β_2 , does not appear on taxa outside B and α .

Proof. Let f be an optimal character. If the taxa in B are monochromatic we are done. Otherwise, run Fitch's algorithm to generate an optimal extension on \mathcal{T}_E . (Also run the Fitch algorithm on \mathcal{T}_V and let m be the number of mutations incurred there, although we do not need to remember the corresponding extension). In \mathcal{T}_E at least one node of B must be a union event (in the bottom-up phase of the Fitch algorithm). Let u be such a node that is furthest from the root of u, and let u be the subtree of u rooted at u. Let u be the two subtrees rooted at the two children of u. The

taxa in \mathcal{T}_1 must be monochromatic with some colour c_1 , and the taxa in \mathcal{T}_2 must be monochromatic with some colour $c_2 \neq c_1$. Suppose, without loss of generality, that the optimal extension colours u with colour c_2 . This causes a mutation between uand the root of T_1 . Hence, if we recolour the entire subtree T_1 (i.e., taxa and non-taxa alike) with colour c_2 , then this generates a new character f' (and new extension) in which the parsimony score of T_E drops by (at least) 1. We argue that f' can decrease the parsimony score of \mathcal{T}_V by (at most) 1, from which the optimality of f' (and its new extension) will follow. Suppose, for the sake of contradiction, that f' generates m-2 or fewer mutations in \mathcal{T}_V . Apply the Fitch algorithm to f' on \mathcal{T}_V . Now, due to the fact that B has essentially the same topology in both T_V and T_E , the subtree T_u is topologically preserved inside T_V . In particular, the images of all vertices of T_u are unambiguously defined inside T_V . Now, in its bottom-up phase the Fitch algorithm will generate in \mathcal{T}_V no union events on the images of the nodes of \mathcal{T}_u , due to the fact that all taxa in \mathcal{T}_u have colour c_2 . (There will, however, be a union event generated at each point where a pendant rooted triplet is grafted onto the image of T_u , i.e., the point in Figure 4 where the dotted line intersects with B. Such subdivision nodes are not considered to be part of the image of \mathcal{T}_u . There will definitely be a union event on such nodes because, due to Property 2.2, c₂ is different to the colours used in the pendant rooted triplet.) At this point we recolour, in T_V , T_1 (taxa and non-taxa alike) with colour c_1 , creating in total exactly one extra mutation, on the edge between uand the root of \mathcal{T}_1 : the fact that the subdivision nodes were all union events prevents additional extra mutations from being created.

This new extension is a valid extension of f on \mathcal{T}_V but generates at most m-1 mutations, contradicting the assumption that an optimal extension of f on \mathcal{T}_V had m mutations. Hence, f' must be optimal.

If f' is not yet monochromatic for B, then we re-run the Fitch algorithm on \mathcal{T}_E to generate a fresh optimal extension, and iterate the entire process until B becomes monochromatic. This process must terminate (in polynomial time) because each iteration merges two distinctly coloured subtrees of B into one strictly larger monochromatic subtree.

Property 3. In \mathcal{T}_E , all the taxa in B (including α) have the same colour, and cherry $\{\beta_1, \beta_2\}$ also has this colour. Moreover, this colour does not appear on any other taxa, i.e., it is unique for B and β_1, β_2 .

Proof. From Property 2.3 we already know that all taxa in B have the same colour and, with the possible exception of β_1 , β_2 , this colour does not appear outside B. Let c be the colour used in B. If c is the same as the colour of $\{\beta_1, \beta_2\}$, denoted again c_β , we are done. If c is the same colour as $\{\gamma_1, \gamma_2\}$, then swapping the colours on $\{\beta_1, \beta_2\}$ and $\{\gamma_1, \gamma_2\}$ preserves optimality, and we are done. (Optimality is preserved because the parsimony score of T_E cannot increase under such a swap, and the parsimony score of T_V cannot decrease due to symmetry.) So suppose neither c_β nor c_γ is equal to c. Run the Fitch algorithm to generate an optimal extension. In the bottom-up phase the Fitch algorithm will assign states $\{c, c_\beta\}$ to p_1 and $\{c, c_\beta, c_\gamma\}$ to p_2 . Suppose, in the top-down phase, the parent of p_2 communicates a state to p_2 that is either equal to c, or not in $\{c, c_\beta, c_\gamma\}$. In this case the Fitch algorithm allows us to give p_2 colour c. We can then recolour $\{\gamma_1, \gamma_2\}$ to be c (saving at least one mutation

in \mathcal{T}_E , and saving at most one mutation in \mathcal{T}_V , thus preserving optimality) and then switch back to the earlier case. If the Fitch algorithm permits p_1 to be coloured c, we simply recolour $\{\beta_1, \beta_2\}$ to be c and we are done because this, via the same analysis, preserves optimality. The only case remaining is if every possible set of choices in the top-down phase of Fitch's algorithm leads to the conclusion that both p_1 and p_2 are coloured c_β . (This is the only remaining case because if p_2 is or can be coloured c_γ , then the Fitch algorithm will subsequently allow us to colour p_1 with colour c, due to the fact that $c_\gamma \notin \{c, c_\beta\}$, i.e., we will be in an earlier case.) So consider an extension generated by the Fitch algorithm in this case. We swap the colours on $\{\beta_1, \beta_2\}$ and $\{\gamma_1, \gamma_2\}$ (including the colours of their parents). This colour swap does not affect the number of mutations but it ensures that both edges leaving p_1 carry mutations. Hence, if we now colour $\{\beta_1, \beta_2\}$, their parent, and p_1 all c, both these mutations vanish. So we definitely save one mutation in \mathcal{T}_E , and as usual at most one mutation is saved in \mathcal{T}_V . So we are done. This concludes the proof of Property 3.

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