Revisiting an equivalence between maximum parsimony and maximum likelihood methods in phylogenetics

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Abstract

Tuftley and Steel (1997) proved that Maximum Likelihood and Maximum Parsimony methods in phylogenetics are equivalent for sequences of characters under a simple symmetric model of substitution with no common mechanism. This result has been widely cited ever since. We show that small changes to the model assumptions suffice to make the two methods inequivalent. In particular, we analyze the case of bounded substitution probabilities as well as the molecular clock assumption. We show that in these cases, even under no common mechanism, Maximum Parsimony and Maximum Likelihood might make conflicting choices. We also show that if there is an upper bound on the substitution probabilities which is ‘sufficiently small’, every Maximum Likelihood tree is also a Maximum Parsimony tree (but not vice versa).

Keywords: phylogenetics, maximum parsimony, maximum likelihood, molecular clock
1 Introduction

Stochastic models for nucleotide substitution and tree reconstruction methods for inferring phylogenetic trees are used to interpret the ever-growing amount of available genetic sequence data. Unsurprisingly, such models and methods have therefore been widely discussed in the last decades (e.g., Felsenstein, 1978; Felsenstein, 2004; Semple and Steel, 2003; Yang, 2006). Two of the most frequently used tree reconstruction methods are Maximum Parsimony (MP) and Maximum Likelihood (ML). A basic difference between these two methods is that MP, unlike ML, is not based on a specific nucleotide substitution model. If the sequences under consideration are related by a specific model of substitution, the results of MP and ML may coincide (Hendy and Penny, 1989), but there are also examples, such as the famous ‘Felsenstein Zone’, for which this is not the case (Felsenstein, 1978).

In 1997, Tuffley and Steel took an important step forward in the analysis of MP and ML (Tuffley and Steel, 1997): they showed that a particular symmetric model of substitution with ‘no common mechanism’ is sufficient for MP and ML to be equivalent when applied to a sequence of characters.

The purpose of this paper is to analyze this equivalence of MP and ML further by considering slightly modified model assumptions that are of biological relevance. For instance, MP is often assumed to be justified whenever the nucleotide substitution probabilities are small (e.g., Felsenstein, 2004, p. 101). Therefore, we restrict the model by placing an upper bound on these probabilities, and find that under no common mechanism MP and ML are no longer equivalent. Moreover, the equivalence of MP and ML under a ‘no common mechanism model’ also fails under the constraint of a molecular clock, even without a bound on the substitution probabilities. These two claims will be established by constructing counterexamples that are minimal with respect to the number of taxa. To construct our examples, we exploit a useful property of the likelihood function for a ‘no common mechanism’ model, namely that it is multilinear in the substitution probabilities.
This fact underlies Equation 18 and Lemma 2 in [Tuffley and Steel, 1997], which we use in our arguments.

We then go on to prove bounds on the probability of observing a given sequence of characters on a tree, and use them to show that it is possible to choose sufficiently small substitution probabilities (depending on the number of taxa, the number of characters and the number of states) so that every tree chosen by ML is also a most parsimonious tree.

2 Notation and Model Assumptions

Recall that a phylogenetic $X$-tree is a tree $T = (V(T), E(T))$ on a leaf set $X = \{1, \ldots, m\} \subset V(T)$ with no vertices of degree 2. Note that the tree does not have to be binary. Furthermore, recall that a character $f$ is a function $f : X \rightarrow \mathcal{C}$ for some set $\mathcal{C} := \{c_1, c_2, c_3, \ldots, c_r\}$ of $r$ character states ($r \in \mathbb{N}$). An extension of $f$ to $V(T)$ is a map $g : V(T) \rightarrow \mathcal{C}$ such that $g(i) = f(i)$ for all $i$ in $X$. For such an extension $g$ of $f$, we denote by $l_T(g)$ the number of edges $e = \{u, v\}$ in $T$ on which a substitution (mutation) occurs, i.e. where $g(u) \neq g(v)$. The parsimony score of $f$ on $T$, denoted by $l_T(f)$, is obtained by minimizing $l_T(g)$ over all possible extensions $g$. The parsimony score of a sequence of characters $S := f_1f_2\ldots f_n$ is given by $l_T(S) = \sum_{i=1}^{n} l_T(f_i)$.

Recall that a character $f$ on a leaf set $X$ is said to be informative (with respect to parsimony) if at least two distinct character states occur more than once on $X$. Otherwise $f$ is called non-informative. Note that for a non-informative character $f$, $l_{T_i}(f) = l_{T_j}(f)$ for all trees $T_i, T_j$ on the same set $X$ of leaves.

Next we describe the fully symmetric $r$-state model [Neyman, 1971], also known as the $N_r$-model, which underlies the Tuffley and Steel equivalence result.

Consider a phylogenetic $X$-tree $T$ arbitrarily rooted at one of its vertices. The $N_r$-
model assumes that a state is assigned to the root from the uniform distribution on the set of states. The state then evolves away from the root as follows. The model assumes equal rates of substitutions between any two distinct character states. For any edge \( e = \{u, v\} \in E(T) \), where \( u \) is the vertex closer to the root, let \( p_e \) denote the conditional probability \( P(v = c_i | u = c_j) \), where \( c_i \neq c_j \). The probability \( p_e \) is equal for all pairs of distinct states \( c_i \) and \( c_j \). Therefore, the probability that a substitution (\( c_j \) to a state different from \( c_j \)) occurs on the edge \( e \) is \((r - 1)p_e\). Let \( q_e \) be the conditional probability \( P(v = c_i | u = c_i) \), i.e. the probability that no substitution occurs on edge \( e \). In the \( N_r \)-model, we have \( 0 \leq p_e \leq \frac{1}{r} \) for all \( e \in E(T) \), and \((r - 1)p_e + q_e = 1\). Moreover, the \( N_r \)-model assumes that substitutions on different edges are independent. Note that for \( r = 4 \), the \( N_r \)-model coincides with the Jukes-Cantor model [Jukes and Cantor, 1969].

Let \( T \) be a phylogenetic \( X \)-tree and let \( f \) be a character on its leaf set \( X \). Let the substitution probabilities assigned to the edges of \( T \) under the \( N_r \)-model be collectively denoted by \( \bar{p} := (p_e : e \in E(T)) \). Then we denote by \( P(f|T, \bar{p}) \) the probability of observing character \( f \) given tree \( T \) and the parameter values \( \bar{p} \). Note that \( P(f|T, \bar{p}) \) does not depend on the root position, since the model is symmetric. The maximum value of this probability for fixed \( f \) and \( T \) as \( \bar{p} \) ranges over all possibilities is denoted by \( \max P(f|T) \), i.e. \( \max P(f|T) := \max_{\bar{p}} P(f|T, \bar{p}) \).

Now let \( S := f_1, \ldots, f_n \) be a sequence of characters. In this paper, we analyze sequences of characters under the \( N_r \)-model with no common mechanism. This means that the substitution probabilities on edges may be different for different characters in \( S \) without any correlation between the characters. We suppose that for each character \( f_i \) in the sequence and for each edge \( e \) of the tree, there is a parameter \( p_{e,i} \) that gives the substitution probability for \( f_i \) on edge \( e \), and that the parameters \( p_{e,i} \) are all independent. For \( i = 1, \ldots, n \), let \( \bar{p}_i := (p_{e,i} : e \in E(T)) \) be the vectors of substitution probabilities. We denote the model parameters \( (\bar{p}_i, i = 1, \ldots, n) \) collectively as \( \Theta \). Then the probability of observing the sequence of characters \( S \) on tree \( T \) for the given parameters \( \Theta \) is given by:
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\[ P(S|T, \Theta) = \prod_{i=1}^{n} P(f_i|T, \bar{p}_i), \]

which follows from the fact that the characters are independent.

We refer to \( P(S|T, \Theta) \) as the probability of observing sequence \( S \) given the phylogenetic tree \( T \) and model parameters \( \Theta \). We then define the likelihood of the tree \( T \) and the model parameters \( \Theta \) given the sequence \( S \), which we refer to as the likelihood function, as \( L(T, \Theta|S) := P(S|T, \Theta) \). The maximum likelihood method of phylogenetic tree reconstruction involves optimizing the likelihood function in two steps as described in Semple and Steel [2003]. We first maximize \( P(S|T, \Theta) \) over the space of model parameters \( \Theta \). We define:

\[ \max P(S|T) := \max_{\Theta} P(S|T, \Theta). \]

We then choose a tree \( T \) that maximizes \( \max P(S|T) \). We call such a tree a maximum likelihood tree (ML-tree) of \( S \). Thus, an ML-tree of a sequence \( S \) is \( \arg\max_{T} (\max P(S|T)) \).

Note that under the assumption of no common mechanism, we have:

\[ \max P(S|T) = \prod_{i=1}^{n} \max_{\bar{p}_i} P(f_i|T, \bar{p}_i). \]

3 Results

Using the notation introduced in the previous section, we are now in a position to state the equivalence result of Tuffley and Steel explicitly.

**Theorem 3.1. (Tuffley and Steel 1997).** Let \( T \) be a phylogenetic \( X \)-tree and let \( S := f_1, \ldots, f_n \) be a sequence of \( r \)-state characters on \( X \). Then, under the \( N_r \)-model with no common mechanism, we have:

\[ \max P(S|T) = r^{-\ell(S)} - n. \]
Thus ML and MP both choose the same tree(s).

In the following, we show that small changes to the assumptions of the $N_r$-model may be enough to make this equivalence fail. In particular, we analyze two settings of biological interest: first, we consider bounded substitution probabilities; secondly we investigate the case of a molecular clock. In both cases, we explicitly construct examples in which MP and ML choose different sets of trees under no common mechanism.

### 3.1 Bounded substitution probabilities

In this section, we consider a modification of the $N_r$-model in which the substitution probabilities on all edges are bounded above by some $u < \frac{1}{r}$. We construct character sequences for which MP and ML choose different sets of trees.

**Proposition 3.2.** Under the $N_r$-model with no common mechanism, for $r \geq 2$, there exist values of $u$ such that if the substitution probabilities are bounded above by $u$, MP and ML choose different sets of trees. In particular, we have:

1. For $r = 2$, for all values of $u \in \left(0, 1 - \frac{1}{\sqrt{2}}\right)$, there exist sequences of characters for which MP and ML choose different sets of trees.

2. For $r > 2$, for all values of $u \in (0, \frac{1}{r})$ there exist sequences of characters for which MP and ML choose unique and distinct trees.

In order to prove this proposition, it is necessary to summarize the main idea of the original proof of the Tuffley-Steel result. We state it here in a more general form so that it may be used to analyze the situation in which the substitution probabilities are bounded.

**Lemma 3.3.** Let $T$ be a phylogenetic $X$-tree and let $f$ be a character on $X$. Then under the $N_r$-model with all substitution probabilities bounded by $u$, where $0 \leq u \leq \frac{1}{r}$, the probability $P(f|T, \bar{p})$ can be maximized at a point where all substitution probabilities are either 0 or $u$. 
Lemma 3.3 is the same as Lemma 2 in [Tuffley and Steel, 1997] except that Tuffley and Steel stated their result only for \( u = \frac{1}{r} \). However, this assumption is not used in their proof and is therefore not required for the lemma to hold. Tuffley and Steel used it to explicitly maximize the probability of observing a character on a given tree under the \( N_r \)-model: for a given character \( f \) and tree \( T \) with a most parsimonious extension \( g \) of \( f \), assigning substitution probability \( \frac{1}{r} \) to edges where a substitution is induced by \( g \), and 0 elsewhere, gives \( \max_{\tilde{p}} P(f | T, \tilde{p}) \) (cf. Theorem 3 of [Tuffley and Steel, 1997]).

But it turns out that an ML solution cannot be similarly related to an MP solution when \( u < \frac{1}{r} \). That is, if \( g \) is a most parsimonious extension of a character \( f \), then we may not be able to maximize the probability by simply assigning the substitution probability \( u \) to edges on which there is a substitution in \( g \), and 0 to edges on which there is no substitution in \( g \). The probability may actually be maximized at some other corner of the feasibility region of \( \tilde{p} \). This is the idea of the following construction.

**Proof of Proposition 3.2.** We provide examples of sequences of characters for which MP and ML may choose different sets of trees. We first prove the case \( r = 2 \) with an example on five taxa, and show that in this case, there are no such examples on fewer than five taxa. Then we explicitly prove the case \( r = 3 \) with an example on four taxa and show how this example can be generalized for \( r > 3 \).

**Case \( r = 2 \):**

Let the set of character states be \( \{ \alpha, \beta \} \). Consider the two trees \( T_1 \) and \( T_2 \) shown in Figure 1 alongside the characters \( f_1 = \alpha\alpha\beta\beta\beta \) and \( f_2 = \alpha\beta\alpha\beta\beta \). We consider the character sequence \( S := f_1 f_2 \).

Note that \( l_{T_1}(f_1) = l_{T_2}(f_2) = 1 \) and \( l_{T_1}(f_2) = l_{T_2}(f_1) = 2 \). Therefore, \( l_{T_1}(S) = l_{T_2}(S) = 3 \), which means that MP will not favor either of the two trees \( T_1, T_2 \) over the other one. Moreover, as \( f_1 \) and \( f_2 \) are incompatible with one another, it can easily be seen that both trees are actually MP-trees: the minimal score of either character is 1, as two states are
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Employed, and this score is achieved when the character corresponds to a split on an edge of the underlying tree – but because of the incompatibility, the other character will have a score of at least 2. So for \( S \), a score of 3 is best possible, and thus both \( T_1 \) and \( T_2 \) are MP-trees.

For ML, the situation is different. This is because the assignments of \( f_1 \) on \( T_2 \) and \( f_2 \) on \( T_1 \) differ, as highlighted by Figure 1. In fact, character \( f_1 \) has a unique most parsimonious extension on \( T_2 \), whereas \( f_2 \) has two most parsimonious extensions on \( T_1 \). As we show in the following, for a sufficiently small upper bound \( u \), the likelihood function is maximized when these extensions both contribute to the likelihood. We use a symbolic algebra system to evaluate \( P(f_i|T, \bar{p}) \) for \( i = 1, 2 \), for all trees on five taxa and at all corners of the feasibility region of \( \bar{p} \) (see Lemma 3.3). More specifically, for the five-leaf-trees under investigation, there are seven edges to which either 0 or \( u \) can be assigned, which gives \( 2^7 = 128 \) possible parameter vectors \( \bar{p} \) at which the likelihood might be maximized. We observe that \( \max P(f_1|T_1) = \max P(f_2|T_2) = \frac{1}{2} u \), but \( \max P(f_1|T_2) = \frac{1}{2} u^2 \) and \( \max P(f_2|T_1) = \max(\frac{1}{2} u^2, u^2(1-u)^2) \). So there are choices of \( u \), namely all \( u < 1 - \frac{1}{\sqrt{2}} \), for which \( \max P(f_1|T_2) < \max P(f_2|T_1) \). In these cases, even though both \( T_1 \) and \( T_2 \) are MP-trees, ML will favor tree \( T_1 \) over \( T_2 \). Therefore, MP and ML are not equivalent in

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**Figure 1:** The characters \( f_1 \) and \( f_2 \) both correspond to a split on an interior edge of \( T_1 \) or \( T_2 \), respectively. But, as highlighted by the circled leaves, the assignment of \( f_1 \) on \( T_2 \) differs from the assignment of \( f_2 \) to \( T_1 \).
Now let sequence \( \tilde{S} \) contain \( n \) copies of character \( f_1 \) and \( n + 1 \) copies of character \( f_2 \) for some integer \( n > 0 \). Then, clearly \( \ell_{T_1}(\tilde{S}) = 3n + 2 \), but \( \ell_{T_2}(\tilde{S}) = 3n + 1 \). Therefore, MP will favor tree \( T_2 \) over \( T_1 \). Moreover, \( T_2 \) is an MP-tree (by the same incompatibility argument concerning \( f_1 \) and \( f_2 \) as above). On the other hand, we have \( \max P(\tilde{S}|T_1) = (\frac{1}{2}u)^n \cdot (u^2(1 - u)^2)^{n+1} \) and \( \max P(\tilde{S}|T_2) = \frac{u^{3n+1}}{2^{3n+1}} \) (provided \( u < 1 - \frac{1}{\sqrt{2}} \)). We choose a sufficiently large value of \( n \) so that the former value is larger than the latter. For such choices of \( n \), ML will favor tree \( T_1 \) over \( T_2 \), even though MP favors \( T_2 \). It is important to note, however, that for the sequence \( \tilde{S} \), the tree \( T_1 \) is not an ML-tree. It can be easily verified for the tree \( T_3 \) in Figure 2 that \( \max P(\tilde{S}|T_3) = (\frac{n}{2})^{n+1} (u^2(1 - u)^2)^n \), which is more than \( \max P(\tilde{S}|T_1) \). In fact, \( \max P(\tilde{S}|T_3) > \max P(\tilde{S}|T_1) \) for all \( u \leq \frac{1}{2} \). In fact, further work shows \( T_3 \) is the unique ML-tree. Moreover, \( T_3 \) is also an MP-tree. So for \( r = 2 \), it remains unclear whether MP and ML can make strictly conflicting choices.

![Figure 2](image)

Note that when \( r = 2 \), examples demonstrating the inequivalence of MP and ML cannot be constructed with fewer than five taxa. This is because given at most one interior edge, it can be easily checked that all non-informative binary characters have the same maximum probability on all trees, whereas informative binary characters on four taxa have a higher probability on the tree where they have parsimony score 1 (calculation not shown).

Case \( r = 3 \):
Let the set of character states be \( \{\alpha, \beta, \gamma\} \). We consider four taxa and the characters \( f_1 := \alpha\alpha\beta\beta \) and \( f_2 := \alpha\beta\gamma\beta \), as well as the sequence \( S \) of characters defined by \( S := f_1 f_2 \ldots f_2 \) \( n \) times. Two of the three possible trees on four taxa are shown in Figure 3: the tree \( T_4 = 12|34 \) and the tree \( T_5 = 13|24 \).

![Figure 3: Tree \( T_4 \) illustrated in (a) is the unique MP-tree for \( S \), whereas (b) depicts tree \( T_5 \), which is the unique ML-tree for \( S \) when \( n \) is chosen sufficiently large.](image)

Tree \( T_4 \) is clearly the unique MP-tree of \( S \), as the only informative character in \( S \) is \( f_1 = \alpha\alpha\beta\beta \).

The ML-trees are obtained as described at the end of Section 2. As before, we used a symbolic algebra system to evaluate \( P(f|T, \bar{p}) \) for all characters \( f \) in the sequence, for all trees on four taxa and at all corners of the feasibility region of \( \bar{p} \) (see Lemma 3.3). We observed that \( \max P(f_2|T_4) = \frac{u^2}{3} \) and \( \max P(f_2|T_5) = u^2(1-2u) \). Therefore, for all \( u < \frac{1}{r} \), we have \( \max P(f_2|T_5) > \max P(f_2|T_4) \). Now for any \( u < \frac{1}{r} \), a sufficiently large value of \( n \) may be chosen such that \( \frac{\max P(S|T_5)}{\max P(S|T_4)} > 1 \). We do not analyze the character \( f_1 \), although the actual choice of \( n \) will depend on the ratio \( \frac{\max P(f_1|T_5)}{\max P(f_1|T_4)} \) and on \( u \). Therefore, MP and ML choose different trees in this three-state setting. Moreover, it turns out that for the third topology on four taxa, namely \( T_6 = 14|23 \), we have \( \max P(S|T_6) < \max P(S|T_5) \) for all choices of \( u \leq \frac{1}{3} \) (calculations not shown). So, \( T_5 \) is the unique ML-tree, whereas \( T_4 \) is the unique MP-tree in this setting. So MP and ML make strictly conflicting choices.

**Case** \( r > 3 \):

Let the set of states be \( \mathcal{C} := \{\alpha, \beta, \gamma, \delta_1, \delta_2, \ldots, \delta_{r-3}\} \). Let \( \mathcal{D} := \{\delta_1, \delta_2, \ldots, \delta_{r-3}\} \). We analyze four taxa and the same characters \( f_1 := \alpha\alpha\beta\beta \) and \( f_2 := \alpha\beta\gamma\beta \) that were analyzed in the case \( r = 3 \), but this time under the \( N_r \)-model with \( r > 3 \). Again we
consider the sequence of characters $S := f_1 f_2 \ldots f_n$.

We only sketch the proof in this case. In particular, we indicate how the expressions for the likelihood function may be written regardless of the number of states.

The expressions for $P(f_i|T_j)$ for $i = 1, 2$ and $j = 1, 2, 3$ can be written in a simple manner since the states $\delta_i$ do not occur in $S$. For example, let the substitution probabilities on the edges of a four-taxa tree $T$ be $\bar{p} = (p_i, i = 1, 2, \ldots, 5)$, where $p_i, i = 1, 2, 3, 4$ are the substitution probabilities on the pending edges adjacent to taxa 1, 2, 3, 4, respectively, and $p_5$ is the substitution probability on the internal edge. Let $v$ and $w$ be the internal vertices of $T$. We write $P(f_i|T, \bar{p}) = \sum_g P(g|T, \bar{p})$, where the summation is over all extensions $g$ of $f_i$.

Now observe that if $g$ and $h$ are two extensions of either $f_1$ or $f_2$, then we have $P(g|T, \bar{p}) = P(h|T, \bar{p})$ if $g(v), h(v) \in \mathcal{D}$ and $g(w) = h(w) = s \notin \mathcal{D}$ (or vice versa with the roles of $v$ and $w$ interchanged).

Therefore:

$$\sum_{g: g(v) \in \mathcal{D}, g(w) = s \notin \mathcal{D}} P(g|T, \bar{p}) = (r - 3) P(h|T, \bar{p}),$$

where $h$ is an extension of $f$ for which $h(v) = \delta_1$ and $h(w) = s$.

Similarly:

$$\sum_{g: g(v) = s \notin \mathcal{D}, g(w) \in \mathcal{D}} P(g|T, \bar{p}) = (r - 3) P(h|T, \bar{p}),$$

where $h$ is an extension of $f$ for which $h(v) = s$ and $h(w) = \delta_1$.

Finally:

$$\sum_{g: g(v) \in \mathcal{D}, g(w) \in \mathcal{D}} P(g|T, \bar{p}) = (r - 3)(1 - 3p_5)p_1p_2p_3p_4.$$
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$P(f_2|T_5)$, and verified that $\max P(f_2|T_5) \geq \frac{u^2(3-2ru)}{r}$ and $\max P(f_2|T_4) = \frac{u^2}{r}$. Since $(3-2ru) > 1$ for all $u < \frac{1}{r}$, there is an $n$ for which $\max P(S|T_5) > \max P(S|T_4)$. This means that ML will favor $T_5$ over $T_4$, even though $T_4$ is the unique MP-tree in this setting. □

Remark 1. It is important to state that in the examples for $r \geq 3$ introduced in the proof of Proposition 3.2, where the number of taxa is bounded (in fact, it is only 4), as $u$ approaches $\frac{1}{r}$, we require $n$ to tend to infinity for ML and MP to make different choices. However, this is a necessary property of any such example for which the number of taxa is bounded: For any fixed character sequence $S$, the continuity of the likelihood function and the Tuffley-Steel result (Theorem 3.1) imply that there is a positive real number $\epsilon(S)$ such that if $u > \frac{1}{r} - \epsilon(S)$, then ML and MP choose the same sets of trees. Therefore, for a bounded number of taxa, since there are only finitely many sequences of length at most $k$, we set $\epsilon := \min_S(\epsilon(S))$, where the minimization takes place over all character sequences of length at most $k$, and conclude that MP and ML would be equivalent (in the sense of the Tuffley-Steel result) for all $u > \frac{1}{r} - \epsilon$, for all sequences of length at most $k$. Therefore, as $u$ approaches $\frac{1}{r}$, the sequence length $n$ of sequences for which MP and ML make conflicting choices has to tend to infinity.

We now complement the above inequivalence results by showing that for sufficiently small choices of $u$, all ML-trees are also MP-trees. To prove this result, we first establish lower and upper bounds for the maximum probability of observing a character given a tree.

Proposition 3.4. Let $T$ be a phylogenetic $X$-tree, where $|X| = m$. Let $f$ be a character on $X$. Let $0 \leq u < \frac{1}{r}$. Then under the $N_r$-model with all substitution probabilities bounded by $u$, we have

$$\left(\frac{1}{r}\right) u^{l_T(f)} \leq \max P(f|T) \leq r^{m-3} u^{l_T(f)}.$$  

Proof. For the lower bound, just as in the Tuffley-Steel approach explained above, we take a most parsimonious extension $g$ of $f$ and assign substitution probability $u$ to each
edge that has a substitution in \( g \), and 0 to all other edges. Considering the \( r \) possible root states (for an arbitrarily chosen root), this gives the lower bound for \( \max _{\bar{p}} P(f | T, \bar{p}) \).

To prove the upper bound, we observe that there are exactly \( r^{m-2} \) extensions of \( f \), where \( m - 2 \) is the number of internal vertices, each of which may be assigned any of the \( r \) states. We will now analyze these extensions. Let \( g \) be any extension of \( f \). For substitution probabilities \( p_e \in \{0, u\} \) assigned to the edges of the tree, the value of \( P(g | T, \bar{p}) \) for an assignment of probabilities that maximizes \( P(f | T, \bar{p}) \) is either 0 (if one of the edges where there is a substitution in \( g \) has been assigned a substitution probability 0) or is given by

\[
P(g | T, \bar{p}) = \frac{1}{r} u^{k_1} (1 - (r - 1)u)^{k_2} \leq \frac{1}{r} u^{k_1} \leq \frac{1}{r} u^{l_T(f)},
\]

where \( k_1 \geq l_T(f) \) is the number of edges where there is a substitution in \( g \), and \( k_2 \) is the number of edges which require no substitution in \( g \) but have been assigned substitution probability \( u \). The factor \( \frac{1}{r} \) is caused by the \( r \) different possible choices for the root state.

The upper bound now follows by summing the probabilities of all extensions.  

Now we will use the above bounds to derive the desired conclusion on ML-trees.

**Theorem 3.5.** Let \( T_a \) and \( T_b \) be two phylogenetic \( X \)-trees, where \( |X| = m \), and let \( S := f_1, f_2, \ldots, f_n \) be a sequence of characters on \( X \). Let the substitution probabilities on all edges of \( T_a \) and \( T_b \) be bounded by \( u \leq r^{(2-m)n} \). Then under the \( N_r \)-model with no common mechanism, we have:

\[
l_T(T_b) < l_T(T_a) \Rightarrow \max P(S | T_b) > \max P(S | T_a).
\]

**Proof.** By Proposition 3.4 we have:

\[
\max P(S | T_a) \leq r^{(m-3)n} u^{\sum l_T(f_i)}
\]

and

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\[
\max P(S|T_b) \geq \left( \frac{1}{r} \right)^n u^{\sum_i l_{T_b}(f_i)}. \tag{3}
\]

Note that for any positive integers \(a\) and \(b\) such that \(b < a\) and any positive constant \(c\), for sufficiently small values of \(u\), we have \(u^a < cu^b\). Now let \(b := \sum_i l_{T_b}(f_i)\) and \(a := \sum_i l_{T_a}(f_i)\) and \(c := \frac{1}{r^{n+1-m}n} = r^{(2-m)n}\). Then Equations (2) and (3) imply that \(\max P(S|T_b) > \max P(S|T_a)\). \(\square\)

The following corollary directly follows from the above theorem.

**Corollary 3.6.** Let \(S\) be sequence of \(n\) characters on a set of \(m\) taxa. Then there is an \(\epsilon = \epsilon(m, n, r)\) such that under the \(N_r\)-model with no common mechanism and with all substitution probabilities subject to an upper bound \(u \in [0, \epsilon]\), all ML-trees of \(S\) are also MP-trees.

### 3.2 Molecular clock

We now prove a statement similar to Proposition 3.2, but with substitution probabilities which conform to a molecular clock. Moreover, we consider only the three-state symmetric model. Under the \(N_3\)-model, we consider placing a bound \(p_{\text{max}}\) on the probability of each particular substitution from the root to any leaf. The value \(p_{\text{max}} = \frac{1}{3}\) means we place no bound beyond that already in the \(N_3\)-model, while \(p_{\text{max}} < \frac{1}{3}\) limits the tree depth.

**Proposition 3.7.** Under the \(N_3\)-model with no common mechanism, with the substitution probabilities constrained by a molecular clock, MP and ML are not equivalent for any bound \(p_{\text{max}} \in [0, \frac{1}{3}]\).

**Proof.** Consider the two rooted four-taxa trees \(T_1\) and \(T_2\) along with substitution probabilities \(p_i\) and \(\tilde{p}_i\), respectively, on their edges as shown in Figure 4. The trees have the same shape but different leaf labels, and possibly different probabilities of a substitution from the root to any of its leaves. Under a molecular clock, we have \(p_1 = p_2\) and \(p_3 = p_4\) in
\( T_1 \), and \( \tilde{p}_1 = \tilde{p}_3 \) and \( \tilde{p}_2 = \tilde{p}_4 \) in \( T_2 \). Let \( p, \tilde{p} \in [0, p_{max}] \) be the probabilities of a substitution from the root \( \rho \) to any leaf in \( T_1 \) and \( T_2 \), respectively.

Then under the \( N_3 \)-model, we write \( p \) and \( \tilde{p} \) in terms of the substitution probabilities on the edges of the trees as follows:

\[
p = (1 - 2p_5)p_1 + p_5(1 - 2p_1) + p_5p_1 = p_1 + p_5 - 3p_1p_5
\]

\[
= (1 - 2p_6)p_3 + p_6(1 - 2p_3) + p_6p_3 = p_3 + p_6 - 3p_3p_6.
\]

Thus \( p_5 = \frac{p - p_1}{1 - 3p_1} \) and \( p_6 = \frac{p - p_1}{1 - 3p_1} \). Similarly, on \( T_2 \), we have \( \tilde{p}_5 = \frac{\tilde{p} - \tilde{p}_1}{1 - 3\tilde{p}_1} \) and \( \tilde{p}_6 = \frac{\tilde{p} - \tilde{p}_2}{1 - 3\tilde{p}_2} \).

As in the proof of Proposition \( 3.2 \) we consider the \( N_3 \)-model with state space \( C := \{\alpha, \beta, \gamma\} \). Consider the characters \( f_1 := \alpha\alpha\beta\beta \) and \( f_2 := \alpha\beta\gamma\beta \), and a sequence of characters \( S := f_1f_2\ldots f_2 \), where \( n \) is a positive integer.

As before, \( T_1 \) is the unique MP-tree of \( S \). We claim that \( T_1 \) is not an ML-tree if \( n \) is sufficiently large. In order to show this, we show that \( \max P(S|T_2) > \max P(S|T_1) \) for a suitable choice of \( n \).

We have

\[
\frac{\max P(S|T_2)}{\max P(S|T_1)} = \frac{(\max P(f_1|T_2))(\max P(f_2|T_2))^n}{(\max P(f_1|T_1))(\max P(f_2|T_1))^n}.
\]
We now demonstrate that $\max P(f_2|T_2) > \max P(f_2|T_1)$ for all values of $p_{\text{max}}$. This allows us to choose a sufficiently large value of $n$ so that the ratio above is more than 1.

First we seek to maximize:

$$P(f_2|T_1, \bar{p}) = \sum_{c \in \mathcal{C}} P(f_2|T_1, \bar{p}, \rho = c) P(\rho = c) = \frac{1}{3} \sum_{c \in \mathcal{C}} P(f_2|T_1, \bar{p}, \rho = c).$$

Using a computer algebra system, we expand the right-hand side of this equation by summing the probabilities over all possible assignments of states to the internal nodes 5 and 6, and substitute $p_5 = \frac{p - p_1}{1 - 3p_1}$ and $p_6 = \frac{p - p_3}{1 - 3p_3}$ to obtain:

$$P(f_2|T_1, \bar{p}) = \frac{p_1p_3(3p_1p_3 - 2p_1 - 2p_3 + 1 + 2p - 3p^2)}{3}.$$  

Observe that for any fixed values of $p_1 + p_3$ and $p$, the expression above is maximized when $p_1p_3$ is maximized, i.e. when $p_1 = p_3$. Therefore, we can substitute $p_1$ for $p_3$ and maximize the resulting expression given by:

$$\frac{p_1^2(1 - p_1)(1 + 3p - 3p_1)}{3}.$$  

Under the constraint $p \in [0, p_{\text{max}}]$, straightforward arguments show that the expression shown above has a maximum at $p_1 = p = p_{\text{max}}$. Therefore:

$$\max P(f_2|T_1) = \frac{p_{\text{max}}^2(1 - 2p_{\text{max}})}{3}. \quad (4)$$

Similar calculations show that:

$$\max P(f_2|T_2) = p_{\text{max}}^2(1 - 2p_{\text{max}}). \quad (5)$$
where the maximum is obtained by setting $\tilde{p}_2 = \tilde{p}_4 = \tilde{p}_5 = 0$ and $\tilde{p}_1 = \tilde{p}_3 = \tilde{p}_6 = \tilde{p} = p_{\text{max}}$.

Equations (4) and (5) imply that $\max P(f_2|T_2) > \max P(f_2|T_1)$ for all $p_{\text{max}} \in (0, \frac{1}{3}]$. Now we can select a sufficiently large value of $n$ so that $\max P(S|T_1) < \max P(S|T_2)$, where the actual choice of $n$ will depend on the ratio $\frac{\max P(f_1|T_2)}{\max P(f_1|T_1)}$ and $p_{\text{max}}$.

This analysis does not show that $T_2$ is an ML-tree, but it shows that $T_1$, which is a unique MP-tree, is not an ML-tree. Therefore, the two methods are not equivalent under the constraint of a molecular clock, even when we assume no common mechanism.

4 Discussion and Outlook

Our main objective was to present examples of sequences of characters for which MP and ML with no common mechanism may choose different sets of trees under the $N_r$-model when the substitution probabilities are bounded above by $u < \frac{1}{r}$ or when a molecular clock is assumed. Our four-taxa examples with $r \geq 3$ character states shows that even if the upper bound $u$ is arbitrarily close to $\frac{1}{r}$, we can find sequences of characters which are sufficiently long to cause MP and ML to make conflicting choices.

The motivation for our four-taxa examples came from the idea of the so-called ‘misleading sequences’, which are sequences for which the (parsimoniously) perfect phylogeny (i.e. a tree on which the whole sequence is completely homoplasy-free) and the tree on which the derived Hamming distances are additive differ (for details, see Huson and Steel, 2004; Bandelt and Fischer, 2008). Even though this discrepancy refers to perfect phylogenies (as opposed to general MP-trees), we used a similar idea to construct our four-taxon examples. In particular, the idea underlying the construction of our sequences is based on the fact that MP ignores parsimoniously non-informative characters in any sequence, whereas ML (just as distance-based methods) does not. We exploited this fact to cause a discrepancy between MP and ML by taking sufficient non-informative characters.
It has been known that there are no binary ‘misleading sequences’: if a set of binary characters is convex on a binary phylogenetic tree, then the Hamming distances of this sequence are a tree metric on the same tree (see [Semple and Steel, 2003], Prop. 7.1.9). But for MP and ML, it is still unknown if there is a sequence of binary characters for which these methods make conflicting choices when the substitution probabilities are bounded above by $u < \frac{1}{r}$. We looked at binary characters on five-taxa trees and found character sequences for which some MP-trees are not ML-trees, but we observed that the ML-trees in our examples were also MP-trees - which means that the equivalence of MP and ML failed. But we did not find an example of strictly conflicting choices in the binary case. Also, we did not find examples of sequences for which MP and ML are not equivalent for values of $u$ which are arbitrarily close to $\frac{1}{2}$. Thus, it would be interesting to analyze two-state models further to decide if all ML-trees are MP-trees and if the equivalence between MP and ML under no common mechanism can fail for values of $u$ that are arbitrarily close to $\frac{1}{2}$.

MP is traditionally assumed to be justified (in the sense of agreement with ML) whenever substitution probabilities are small (see, for example, [Felsenstein, 2004]). Therefore, our result, which shows that an upper bound on the substitution probabilities can make the equivalence of MP and ML fail under the $N_r$-model with no common mechanism, is particularly surprising. On the other hand, we have shown that for sufficiently small choices of the upper bound, all ML-trees are at least also MP-trees (but not vice versa). So in summary, although MP has been proven to agree with ML in the $N_r$-model under the assumption of no common mechanism (and under no further constraints), our examples show that this equivalence may fail when the model is changed slightly. Therefore, we conclude that neither the presence nor the absence of a common mechanism alone can justify MP in the sense of an MP-ML equivalence. More research could be done on other models of nucleotide substitution in order to analyze conditions under which ML and MP may give conflicting results. This might highlight even more differences between MP and ML.
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