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Quantifying Hybridization in Realistic Time

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Running head. Quantifying Hybridization in Realistic Time

Key words: reticulate evolution, hybridization, agreement forests, interleaving, fixedparameter tractability.

1 Abstract

Recently, numerous practical and theoretical studies in evolutionary biology aim at calculating the extent to which reticulation—for example horizontal gene transfer, hybridization, or recombination—has influenced the evolution for a set of present-day species. It has been shown that inferring the minimum number of hybridization events that is needed to simultaneously explain the evolutionary history for a set of trees is an NP-hard and also fixed-parameter tractable problem. In this paper, we give a new fixed-parameter algorithm for computing the minimum number of hybridization events for when two rooted binary phylogenetic trees are given. This newly developed algorithm is based on interleaving—a technique using repeated kernelization steps that are applied throughout the exhaustive search part of a fixed-parameter algorithm. To show that our algorithm runs efficiently to be applicable to a wide range of practical problem instances, we apply it to a grass data set and highlight the significant improvements in terms of running times in comparison to an algorithm which has previously been implemented.



2 Introduction

Molecular evolution (phylogenetics) is a lively field of research that is affected by a variety of scientific disciplines. Viewing it from the perspective of computer science, the NP-hardness of many fundamental problems in phylogenetics makes it a challenging subject to study. The theoretically well-analyzed and widely-applied tree reconstruction methods of maximum parsimony and maximum likelihood are prominent examples of such NP-hard problems (Foulds and Graham, 1982; Chor and Tuller, 2005; Roch, 2006). In this paper, we consider a particular NP-hard problem that is fundamental in the study of reticulate evolution.

Evolutionary biologists often observe inconsistencies amongst phylogenetic trees that represent the evolution of different parts of present-day species genomes. Such inconsistencies can essentially be caused either by reticulation events like horizontal gene transfer, hybridization, and recombination, or by non-biological processes like sequencing errors or signals in the data that may yield trees whose branching patterns do not always represent the correct evolutionary history. Here, we assume that hybridization (as a representative of reticulation) has led to the observed inconsistencies. In this case, it may be more appropriate to represent the evolutionary history of a set of species by a phylogenetic network rather than a phylogenetic tree since the parents of a hybrid taxa belong to two different species. It is consequently desirable to calculate a hybridization network that simultaneously explains the evolutionary histories for a given set of trees and minimizes the number of hybridization events. The reason for the latter is that it quantifies the significance of hybridization for the evolution of the species under consideration. However, computing this minimum number is NP-hard even for two trees (Bordewich and Semple, 2007a). Known as MINIMUM HYBRIDIZATION, it is this two-tree problem that is the focus of this paper.

To overcome the computational burden of NP-hard problems, one frequently resorts to approximation algorithms, heuristics, or solving polynomial-time restrictions of the

Journal of Computational Biology

problem. However, the solutions obtained from these approaches are not always acceptable; for example, this may be due to complex and expensive processes that were needed to generate certain data sets. In such cases, fixed-parameter algorithms have proven to be a valuable tool to calculate the exact solution of a computationally-hard problem. Mathematically speaking, a problem of size n, parameterized by k, is *fixed-parameter tractable* if it can be computed in O(f(k)p(n)), where f is a computable function and pis a fixed polynomial. The importance of this running time is in the separation of the variables k and n. Thus, if k is small, the problem may be tractable in reasonable time even if n is large. For a more detailed description of fixed-parameter tractability (FPT), we refer the interested reader to Downey and Fellows (1998) and Flum and Grohe (2006).

MINIMUM HYBRIDIZATION and various other problems in computational biology are known to be fixed-parameter tractable (for example, Ávila et al., 2006; Bordewich and Semple, 2007b; Gramm et al., 2008). However, practical fixed-parameter algorithms that have been applied to biological data sets rarely exist. Recently, Bordewich et al. (2007) implemented a fixed-parameter algorithm for MINIMUM HYBRIDIZATION. By applying this algorithm to a grass data set, the authors subsequently showed that many problem instances were computable within a couple of minutes. However, there were several instances to which the algorithm did not return the exact answer in reasonable time. In particular, for three tree pairs, the running time to calculate the exact solution was at least two days. Other studies in computational biology that have introduced fixed-parameter algorithms and applied them to biological or synthetic data sets are for example described in Dehne et al. (2006), and Gramm and Niedermeier (2002, 2003).

To keep up with the constant progress in molecular biology, which primarily originates from the development of efficient DNA sequencing technologies, it is of importance to develop new and to further improve existing fixed-parameter algorithms such that they can cope with an increasing data set size. A common technique for obtaining a fixedparameter algorithm for a problem is *kernelization*. The aim of this technique is to shrink the size of the initial problem instance to its difficult core by quickly resolving those parts of the problem that are easily dealt with. Kernelizaton is the technique used by Bordewich et al. (2007) in their implementation for solving MINIMUM HYBRIDIZATION. To complement kernelization, *interleaving* has been introduced as a new method in the design of fixed-parameter algorithms (Niedermeier and Rossmanith, 2000). Interleaving refers to repeated kernelization steps while one systematically processes a bounded search tree. Apart from Abu-Khzam et al. (2006) and Dehne et al. (2006), where the authors showed that interleaving has a positive impact on the overall running time of a fixedparameter algorithm, this technique has so far attracted more attention in theoretical analyses concerned with FPT than in practical studies.

Making use of interleaving, we present a fixed-parameter algorithm for solving MIN-IMUM HYBRIDIZATION that performs significantly better than that given by Bordewich et al. (2007). This improvement is highlighted by the fact that all instances of the grass data set described above can be solved in less than eleven minutes. As an example of the new algorithm's performance, an instance for which the previously implemented algorithm did not return the solution within two days, can now be calculated in less than a minute.

The new algorithm—called HYBRIDINTERLEAVE—has been implemented in Java and is available for application at

http://www.math.canterbury.ac.nz/~c.semple/software.shtml

or

http://wwwcsif.cs.ucdavis.edu/~linzs/.

To start a calculation with HYBRIDINTERLEAVE, the algorithm requires the two input trees to be given in Newick format. As output, the program provides the user with the minimum number of hybridization events that explain the two input trees.

This paper is organized as follows. The next section contains some preliminaries and

Journal of Computational Biology

formally states the decision problem MINIMUM HYBRIDIZATION for which a previously established fixed-parameter algorithm is summarized in Section 4. The new algorithm HYBRIDINTERLEAVE and its proof of correctness are given in Section 5, while Section 6 analyzes the running times of HYBRIDINTERLEAVE when applied to a grass data set and compares it with the running times of the recently implemented algorithm HYBRID-NUMBER (Bordewich et al., 2007). We end the paper with some concluding remarks in Section 7. Unless otherwise stated, the notation and terminology follows Semple and Steel (2003).

Preliminaries

This section provides preliminary definitions which are used throughout the rest of this paper and formally states the decision problem MINIMUM HYBRIDIZATION.

3.1 Phylogenetic Trees

A rooted binary phylogenetic X-tree \mathcal{T} is a rooted tree whose root has degree two while all other interior vertices have degree three. The leaf set X is the label set of \mathcal{T} and frequently denoted by $\mathcal{L}(\mathcal{T})$. Furthermore, a subset A of X is a *cluster* of \mathcal{T} if there is a vertex v whose set of descendants is precisely A. We view v as an ancestor and descendant of itself.

In the course of this paper, two types of subtrees play an important role. Let X' be a subset of X, and let \mathcal{T} be a rooted phylogenetic X-tree. The minimal rooted subtree of \mathcal{T} that connects all leaves in X' is denoted by $\mathcal{T}(X')$. Furthermore, the subtree obtained from $\mathcal{T}(X')$ by contracting all non-root degree-2 vertices is the *restriction of* \mathcal{T} to X'and is denoted by $\mathcal{T}|X'$. Lastly, a subtree is *pendant* if it can be detached from \mathcal{T} by deleting a single edge.

3.2 Hybridization Networks

A hybridization network \mathcal{H} on a set X is a rooted acyclic digraph with root ρ such that the following properties are satisfied:

- (i) X is the set of vertices of out-degree 0,
- (ii) the out-degree of ρ is at least 2, and
- (iii) for all vertices v with $d^+(v) = 1$, we have $d^-(v) = 2$,

Journal of Computational Biology

where $d^+(v)$ and $d^-(v)$ denote the out-degree and in-degree of v, respectively. The set X represents a collection of present-day taxa, and vertices of in-degree two represent an exchange of genetic information between their parents. Generically, we call these vertices *hybridization vertices*.

Figure1

To quantify the number of hybridization events, the hybridization number of \mathcal{H} is the total number of hybridization vertices. Observe that if \mathcal{H} is a rooted binary phylogenetic tree, then $h(\mathcal{H}) = 0$. Ignoring the thickness of the arcs, the left-hand-side of Figure 1 shows a hybridization network \mathcal{H} whose hybridization number is three.

Now let \mathcal{H} be a hybridization network on X, and let \mathcal{T} be a rooted binary phylogenetic X'-tree with $X' \subseteq X$. We say that \mathcal{T} is *displayed* by \mathcal{H} if \mathcal{T} can be obtained from \mathcal{H} by deleting a subset of its edges and any resulting degree-0 vertices, and then contracting edges. For example, the rooted binary phylogenetic tree \mathcal{T} shown in Figure 1 is displayed by the hybridization network \mathcal{H} of the same figure. Intuitively, if \mathcal{H} displays \mathcal{T} , then all of the ancestral relationships visualized by \mathcal{T} are visualized by \mathcal{H} . Extending the definition of the hybridization number to two rooted binary phylogenetic X-trees \mathcal{S} and \mathcal{T} , we set

 $h(\mathcal{S}, \mathcal{T}) = \min\{h(\mathcal{H}) : \mathcal{H} \text{ is a hybridization network that displays } \mathcal{S} \text{ and } \mathcal{T}\}.$

With the above definition, we now formally state MINIMUM HYBRIDIZATION.

Problem: MINIMUM HYBRIDIZATION(S, T, k)

Instance: Two rooted binary phylogenetic X-trees S and T, and an integer k. Question: Is h(S,T) < k?

Figure

3.3 Agreement Forests

Originating from an idea in Hein et al. (1996), Bordewich and Semple (2007a) showed that MINIMUM HYBRIDIZATION is NP-complete by using a characterization of the problem in terms of agreement forests. Such forests play a fundamental role in this paper. For the purpose of the upcoming definitions, we regard the root of a rooted binary phylogenetic X-tree \mathcal{T} as a vertex ρ at the end of a pendant edge adjoined to the original root. For an example of two such trees, see Figure 2. Furthermore, we view ρ as an element of the label set of \mathcal{T} ; thus $\mathcal{L}(\mathcal{T}) = X \cup \{\rho\}$. Now, let \mathcal{S} and \mathcal{T} be two rooted binary phylogenetic X-trees. An agreement forest $\mathcal{F} = \{\mathcal{L}_{\rho}, \mathcal{L}_{1}, \mathcal{L}_{2}, \dots, \mathcal{L}_{k}\}$ for \mathcal{S} and \mathcal{T} is a partition of $\mathcal{L}(\mathcal{S})$ such that $\rho \in \mathcal{L}_{\rho}$ and the following conditions are fulfilled:

- (1) For all $i \in \{\rho, 1, 2, \dots, k\}$, we have $\mathcal{S}|\mathcal{L}_i \cong \mathcal{T}|\mathcal{L}_i$.
- (2) The trees in $\{\mathcal{S}(\mathcal{L}_i) : i \in \{\rho, 1, 2, ..., k\}\}$ and $\{\mathcal{T}(\mathcal{L}_i) : i \in \{\rho, 1, 2, ..., k\}\}$ are vertex-disjoint subtrees of \mathcal{S} and \mathcal{T} , respectively.

As an example, two agreement forests for the two rooted binary phylogenetic trees S and T shown in Figure 2 are $\mathcal{F} = \{\{\rho, 7\}, \{1, 2, 3\}, \{4, 5, 6\}\}$ and $\mathcal{F}' = \{\{\rho, 1, 2, 3, 7\}, \{4\}, \{5\}, \{6\}\}.$

A characterization of the minimum number $h(\mathcal{S}, \mathcal{T})$ of hybridization events in terms of agreement forests requires an additional condition. Without going into details, this condition avoids the possibility of species inheriting genetic material from their own descendants. Let $\mathcal{F} = \{\mathcal{L}_{\rho}, \mathcal{L}_1, \mathcal{L}_2, \dots, \mathcal{L}_k\}$ be an agreement forest for \mathcal{S} and \mathcal{T} . Let $G_{\mathcal{F}}$ be the directed graph that has vertex set \mathcal{F} and an arc from \mathcal{L}_i to \mathcal{L}_j precisely if $i \neq j$, and either

- (1) the root of $\mathcal{S}(\mathcal{L}_i)$ is an ancestor of the root of $\mathcal{S}(\mathcal{L}_i)$ in \mathcal{S} or
- (2) the root of $\mathcal{T}(\mathcal{L}_i)$ is an ancestor of the root of $\mathcal{T}(\mathcal{L}_j)$ in \mathcal{T} .

Figure

We call \mathcal{F} an *acyclic-agreement forest* for \mathcal{S} and \mathcal{T} if $G_{\mathcal{F}}$ has no directed cycle. Moreover, if \mathcal{F} contains the smallest number of parts over all acyclic-agreement forests for \mathcal{S} and \mathcal{T} , we say that \mathcal{F} is a *maximum-acyclic-agreement forest* for \mathcal{S} and \mathcal{T} , in which case, we denote this number minus one by $m_a(\mathcal{S}, \mathcal{T})$. Figure 3 shows the two digraphs $G_{\mathcal{F}}$ and $G_{\mathcal{F}'}$ that are associated with the agreement forests \mathcal{F} and \mathcal{F}' , respectively, for the two rooted binary phylogenetic X-trees \mathcal{S} and \mathcal{T} depicted in Figure 2. Note that, as $G_{\mathcal{F}'}$ is acyclic, \mathcal{F}' is an acyclic-agreement forest for \mathcal{S} and \mathcal{T} while \mathcal{F} is no such forest. Indeed, \mathcal{F}' is a maximum-acyclic-agreement forest for \mathcal{S} and \mathcal{T} . Baroni et al. (2005) established the following characterization.

Theorem 1. Let S and T be two rooted binary phylogenetic X-trees. Then

 $h(\mathcal{S},\mathcal{T}) = m_a(\mathcal{S},\mathcal{T}).$

It is this characterization that was used to show that MINIMUM HYBRIDIZATION is NP-complete.

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4 Overview of a Known Fixed-Parameter Algorithm for MINIMUM HYBRIDIZATION

In this section, we summarize the basic ideas of the first fixed-parameter algorithm for MINIMUM HYBRIDIZATION. This algorithm is based on an earlier result that showed, for a pair of rooted binary phylogenetic X-trees S and T, MINIMUM HYBRIDIZATION is fixed-parameter tractable with h(S, T) being the parameter (Bordewich and Semple, 2007b). In establishing this result, the authors used two reductions—called the subtree and chain reduction—that kernelize S and T to two smaller trees whose number of leaves is linear in h(S, T).

Before detailing these reductions, we need some additional definitions. Let \mathcal{T} be a rooted binary phylogenetic X-tree. An *n*-chain of \mathcal{T} is an ordered tuple (a_1, a_2, \ldots, a_n) of elements in X such that the parent of a_1 is either the same as the parent of a_2 or a child of the parent of a_2 and, for all $i \in \{2, 3, \ldots, n-1\}$, the parent of a_i is a child of the parent of a_{i+1} . For example, referring to Figure 2, (1, 2, 3, 4, 5, 6) and (4, 5, 6) are chains of \mathcal{S} . Furthermore, let \mathcal{S} and \mathcal{T} be two rooted binary phylogenetic X-trees, and let P be a disjoint collection of 2-element subsets of X such that each pair $\{a, b\} \in P$ is a common 2-chain of \mathcal{S} and \mathcal{T} . Let the weight function $w : P \to \mathbb{Z}^+$ assign each element of P a positive integer weight. We refer to \mathcal{S} and \mathcal{T} with an associated weight function w as a pair of weighted phylogenetic trees on X. The set P is referred to as the set of weighted 2-chains of \mathcal{S} and \mathcal{T} . Unless otherwise stated, we will use w to denote the weighting of the 2-chains in P.

We now describe the reductions. Let S and T be two weighted rooted binary phylogenetic X-trees with weighted 2-chain set P.

Subtree reduction. Replace a maximal pendant subtree that is common to S and T by a single leaf with a new label. Furthermore, delete all members in P whose elements label leaves of the pendant subtree under consideration.

Journal of Computational Biology

Chain reduction. Replace a maximal *n*-chain (a_1, a_2, \ldots, a_n) with n > 2 that occurs Figure identically in \mathcal{S} and \mathcal{T} by a 2-chain with new labels a and b. Furthermore, add the 4 2-element set $\{a, b\}$ to P with an associated weight of

$$w(\{a,b\}) = n - 2 + \sum_{\substack{\{a_i,a_j\} \in P\\ \text{and } a_i, a_j \in \\ \{a_1, \dots, a_n\}}} w(\{a_i, a_j\}),$$

and delete all members in P whose elements are in $\{a_1, a_2, \ldots, a_n\}$. An explicit example of the chain reduction is shown in Figure 4, where the two rooted binary phylogenetic trees \mathcal{S}' and \mathcal{T}' have been obtained from \mathcal{S} and \mathcal{T} , which are shown in Figure 2, by replacing the 3-chain (1, 2, 3) with the 2-chain (a, b), and similarly, the 3-chain (4, 5, 6)with (c, d). Note that $w(\{a, b\}) = w(\{c, d\}) = 1$.

The correspondence between the trees resulting from repeated applications of the subtree and chain reductions, and the initial two trees is given in the next lemma. This correspondence is done via a notion of agreement forests that extends acyclic-agreement forests. An agreement forest \mathcal{F} for two rooted weighted binary phylogenetic X-trees \mathcal{S} and \mathcal{T} with weighted 2-chain set P is called *legitimate* if it is acyclic and the following property holds:

(P) For each $\{a, b\} \in P$, either $\{a\}$ and $\{b\}$ are elements in \mathcal{F} , or there exists an element of \mathcal{F} , say \mathcal{L} , such that $\{a, b\} \subseteq \mathcal{L}$.

Let \mathcal{F} be a legitimate-agreement forest for \mathcal{S} and \mathcal{T} . Set

$$w_c(\mathcal{F}, P) = \sum_{\substack{\{a,b\} \in P;\\ \{a\}, \{b\} \in \mathcal{F}}} w(\{a,b\}),$$

we define the *weight* of \mathcal{F} as

$$w(\mathcal{F}) = |\mathcal{F}| - 1 + w_c(\mathcal{F}, P)$$

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and set $f(\mathcal{S},\mathcal{T})$ to be the minimum weight of a legitimate-agreement forest for \mathcal{S} and \mathcal{T} . Note that we always have $f(\mathcal{S}, \mathcal{T}) \geq h(\mathcal{S}, \mathcal{T})$, and $f(\mathcal{S}, \mathcal{T}) = h(\mathcal{S}, \mathcal{T})$ whenever P is empty. The next two lemmas are central to showing that MINIMUM HYBRIDIZATION is fixed-parameter tractable (Bordewich and Semple, 2007b).

Lemma 2. Let S and T be two weighted rooted binary phylogenetic X-trees, and let \mathcal{S}' and \mathcal{T}' be two weighted rooted binary phylogenetic X'-trees that have been obtained from S and T, respectively, by applying the subtree or chain reduction. Then f(S,T) = $f(\mathcal{S}', \mathcal{T}').$

Lemma 3. Let S and T be two weighted rooted binary phylogenetic X-trees whose weighted 2-chain set P is empty. Furthermore, let \mathcal{S}' and \mathcal{T}' be two weighted rooted binary phylogenetic X'-trees that have been obtained from S and T, respectively, by repeatedly applying the subtree and chain reduction until no further reduction is possible. Then $|X'| \leq 14h(\mathcal{S}, \mathcal{T}).$

Cluster reduction. Besides repeatedly applying the subtree and chain reductions to Figure kernelize a problem instance of MINIMUM HYBRIDIZATION before exhaustively calculat-ing a legitimate-agreement forest of minimum weight, we can use a third reduction that breaks a problem instance of MINIMUM HYBRIDIZATION into two smaller subproblems. This reduction is depicted in Figure 5 and can be repeatedly intertwined with the other two reductions before the inevitable exhaustive search part of the algorithm. How the two smaller problem instances relate to the original instance is described in the next corollary. Due to Linz (2008), this corollary generalizes the unweighted version given by Baroni et al. (2006).

Corollary 4. Let S and T be two weighted rooted binary phylogenetic X-trees with weighted 2-chain set P, and let A be a common minimal cluster of both S and T with $|A| \geq 2$. Then,

 $f(\mathcal{S}, \mathcal{T}) = f(\mathcal{S}|A, \mathcal{T}|A) + f(\mathcal{S}_a, \mathcal{T}_a),$

where S_a and T_a are the trees obtained from S and T, respectively, by replacing the pendant subtree whose label set is precisely A with a new leaf labeled a.

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Journal of Computational Biology

In the last corollary, the weighted 2-chain sets, P_A and P_a say, of S|A and T|A, and S_a and T_a , respectively, are

$$P_A = \{\{\ell, \ell'\} : \{\ell, \ell'\} \subseteq P \text{ and } \ell, \ell' \in A\}$$

and

$$P_a = \{\{\ell, \ell'\} : \{\ell, \ell'\} \subseteq P \text{ and } \ell, \ell' \notin A\},\$$

where, for both sets, the weight of each element $\{\ell, \ell'\}$ is equal to the weighting of $\{\ell, \ell'\}$ in the weight function associated with P.

Remarks.

- (i) Note that the cluster reduction can repeatedly be applied to break S and T into as many smaller tree pairs as possible by setting A to be a minimal common cluster of S and T with |A| ≥ 2, and resetting S and T to be S_a and T_a, respectively, before applying this reduction again until S ≅ T.
- (ii) We impose maximality on a common pendant subtree and a common n-chain and minimality on a common cluster to guarantee that the corresponding label set of any such common feature intersects each member of P in either both elements or neither.

5 A New Algorithm for MINIMUM HYBRIDIZATION

In this section, we present our fixed-parameter algorithm HYBRIDINTERLEAVE for MIN-IMUM HYBRIDIZATION. It makes use of the subtree, chain, and cluster reductions, but importantly, in terms of obtaining significantly decreased running times (see Section 6), it additionally uses interleaving.

Before outlining HYBRIDINTERLEAVE and giving its pseudocode, we state two lemmas that are central to its correctness and description. Let \mathcal{T} be a rooted binary phylogenetic X-tree, and let ℓ and ℓ' be elements of X. To ease reading in this section, we use $\mathcal{T}[-\ell]$ to denote $\mathcal{T}|(\mathcal{L}(\mathcal{T}) - \{\ell\})$ and $\mathcal{T}[-\ell, \ell']$ to denote $\mathcal{T}|(\mathcal{L}(\mathcal{T}) - \{\ell, \ell'\})$. Furthermore, let S and \mathcal{T} be two weighted rooted binary phylogenetic X-trees with weighted 2-chain set P. If ℓ is contained in a member of P, we say that ℓ crosses P.

Lemma 5. Let S and T be two weighted rooted binary phylogenetic X-trees with weighted 2-chain set P. Suppose that S and T have no common pendant subtree whose leaf set size is at least 2. Then, for each $\ell \in X$, we have

$$f(\mathcal{S},\mathcal{T}) \le f(\mathcal{S}[-\ell,\ell'],\mathcal{T}[-\ell,\ell']) + 2 + w(\{\ell,\ell'\})$$

if ℓ crosses P with $\{\ell, \ell'\} \in P$, and

P, and

$$f(\mathcal{S}, \mathcal{T}) \leq f(\mathcal{S}[-\ell], \mathcal{T}[-\ell]) + 1$$

otherwise.

Proof. First assume that ℓ crosses P in an element $\{\ell, \ell'\}$. Let \mathcal{F}_{ℓ} be a legitimateagreement forest for $\mathcal{S}[-\ell, \ell']$ and $\mathcal{T}[-\ell, \ell']$ of minimum weight. Then it is easily checked that

$$\mathcal{F}=\mathcal{F}_\ell\cup\{\{\ell\},\{\ell'\}\}$$

is a legitimate-agreement forest for S and T. Moreover, we have $|\mathcal{F}| = |\mathcal{F}_{\ell}| + 2$ and

$$w_{c}(\mathcal{F}_{\ell}, P - \{\ell, \ell'\}) = w_{c}(\mathcal{F}, P) - w(\{\ell, \ell'\}). \text{ Hence,}$$

$$f(\mathcal{S}[-\ell, \ell'], \mathcal{T}[-\ell, \ell']) + 2 + w(\{\ell, \ell'\})$$

$$= |\mathcal{F}_{\ell}| - 1 + w_{c}(\mathcal{F}_{\ell}, P - \{\ell, \ell'\}) + 2 + w(\{\ell, \ell'\})$$

$$= |\mathcal{F}| - 1 + w_{c}(\mathcal{F}, P)$$

$$\geq f(\mathcal{S}, \mathcal{T}).$$
(1)

Now assume that ℓ does not cross P. Let \mathcal{F}_{ℓ} be a legitimate-agreement forest for $\mathcal{S}[-\ell]$ and $\mathcal{T}[-\ell]$ of minimum weight. Again, it is clear that

$$\mathcal{F}=\mathcal{F}_\ell\cup\{\ell\}$$

is a legitimate-agreement forest for S and T. Moreover, we have $|\mathcal{F}| = |\mathcal{F}_{\ell}| + 1$ and $w_c(\mathcal{F}_{\ell}, P) = w_c(\mathcal{F}, P)$. Thus

$$f(\mathcal{S}[-\ell], \mathcal{T}[-\ell]) + 1 = |\mathcal{F}_{\ell}| - 1 + w_c(\mathcal{F}_{\ell}, P) + 1$$

$$= |\mathcal{F}| - 1 + w_c(\mathcal{F}, P)$$

$$\geq f(\mathcal{S}, \mathcal{T}).$$

$$(2)$$

Inequalities (1) and (2) establish the lemma.

Lemma 6. Let S and T be two weighted rooted binary phylogenetic X-trees with weighted 2-chain set P. Suppose that S and T have no common pendant subtree whose leaf set size is at least 2. Then there exists an element $\ell \in X$ such that either ℓ crosses P with $\{\ell, \ell'\} \in P$ and

$$f(\mathcal{S}, \mathcal{T}) = f(\mathcal{S}[-\ell, \ell'], \mathcal{T}[-\ell, \ell']) + 2 + w(\{\ell, \ell'\}),$$

or ℓ does not cross P and

$$f(\mathcal{S}, \mathcal{T}) = f(\mathcal{S}[-\ell], \mathcal{T}[-\ell]) + 1$$

Proof. Let $\mathcal{F} = \{\mathcal{L}_{\rho}, \mathcal{L}_{1}, \mathcal{L}_{2}, \dots, \mathcal{L}_{k}\}$ be a legitimate-agreement forest for \mathcal{S} and \mathcal{T} of minimum weight. First observe that, as $G_{\mathcal{F}}$ is acyclic, it has a vertex \mathcal{L}_{i} with $i \in$

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 $\{\rho, 1, 2, \ldots, k\}$ whose out-degree is zero. Furthermore, since S and T have no common pendant subtree whose leaf set size is at least 2, \mathcal{L}_i is a singleton in \mathcal{F} . Since ρ is never a singleton in \mathcal{F} by Lemma 1 of Baroni et al. (2005), we may assume that $\mathcal{L}_i = \{\ell\}$, where $\ell \in X$.

First assume that ℓ crosses P in an element $\{\ell, \ell'\}$. Since \mathcal{F} is legitimate, $\{\ell'\} \in \mathcal{F}$ as $\{\ell\} \in \mathcal{F}$, and so

$$\mathcal{F}_\ell = \mathcal{F} - \{\{\ell\}, \{\ell'\}\}$$

is a legitimate-agreement forest for $\mathcal{S}[-\ell, \ell']$ and $\mathcal{T}[-\ell, \ell']$. Furthermore, we have $|\mathcal{F}| = |\mathcal{F}_{\ell}| + 2$ and $w_c(\mathcal{F}, P) = w_c(\mathcal{F}_{\ell}, P - \{\ell, \ell'\}) + w(\{\ell, \ell'\})$. It now follows that

$$f(\mathcal{S}, \mathcal{T}) = |\mathcal{F}| - 1 + w_c(\mathcal{F}, P)$$

$$= |\mathcal{F}_{\ell}| + 2 - 1 + w_c(\mathcal{F}_{\ell}, P - \{\ell, \ell'\}) + w(\{\ell, \ell'\})$$

$$\geq f(\mathcal{S}[-\ell, \ell'], \mathcal{T}[-\ell, \ell']) + 2 + w(\{\ell, \ell'\}).$$
(3)

Second assume that ℓ does not cross P. Since \mathcal{F} is a legitimate-agreement forest for \mathcal{S} and \mathcal{T} ,

$$\mathcal{F}_\ell = \mathcal{F} - \{\ell\}$$

is such a forest for $\mathcal{S}[-\ell]$ and $\mathcal{T}[-\ell]$. Furthermore, we have $|\mathcal{F}| = |\mathcal{F}_{\ell}| + 1$ and $w_c(\mathcal{F}, P) = w_c(\mathcal{F}_{\ell}, P)$. It now follows that

$$f(\mathcal{S}, \mathcal{T}) = |\mathcal{F}| - 1 + w_c(\mathcal{F}, P)$$

$$= |\mathcal{F}_{\ell}| + 1 - 1 + w_c(\mathcal{F}_{\ell}, P)$$

$$\geq f(\mathcal{S}[-\ell], \mathcal{T}[-\ell]) + 1.$$

$$(4)$$

Combining (3) and (4) with Lemma 5 gives the lemma.

We next give a brief outline of the algorithm HYBRIDINTERLEAVE before detailing its pseudocode. The algorithm takes as input two rooted binary phylogenetic X-trees S and T, and an integer k, and outputs h(S,T) precisely if h(S,T) < k. It starts

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Journal of Computational Biology

with initializing the variable P, the set of weighted 2-chains, that has previously been obtained by applying a chain reduction to S and T. Recall that w is the weight function associated with P. HYBRIDINTERLEAVE then directly calls the subroutine INTERLEAVE which contains the key features of this algorithm.

If k > 0, INTERLEAVE initially finds all maximal pendant subtrees that are common to \mathcal{S} and \mathcal{T} . If the resulting two trees have a label set size of at most 3, then, as $\rho \in \mathcal{L}(\mathcal{S})$, they are identical. Consequently, INTERLEAVE directly returns 0 as the minimum weight for a legitimate-agreement forest of \mathcal{S} and \mathcal{T} . Otherwise, the algorithm proceeds with replacing each maximal common *n*-chain, where $n \geq 3$, with a 2-chain. Resetting \mathcal{S} and \mathcal{T} to be the reduced weighted trees, they always have a cluster A with $2 \leq |A| < |\mathcal{L}(\mathcal{S})|$ in common which allows for an application of the cluster reduction. This reduction returns two new tree pairs. The second pair \mathcal{S}'' and \mathcal{T}'' has been obtained from \mathcal{S} and \mathcal{T} by replacing $\mathcal{S}(A)$ and $\mathcal{T}(A)$, respectively, by a new leaf while the first pair is $\mathcal{S}' = \mathcal{S}|A$ and $\mathcal{T}' = \mathcal{T}|A$ (viewing the root of \mathcal{S}' and \mathcal{T}' , respectively, as a vertex ρ' adjoined to the original root by a pendant edge). With $P' = \{\{\ell, \ell'\} \in P : \{\ell, \ell'\} \subseteq A\}$ whose associated weight function is w', the algorithm next checks whether there exists a legitimate-agreement forest for \mathcal{S}' and \mathcal{T}' with $f(\mathcal{S}', \mathcal{T}') < k$, where w' is obtained from w by restricting its domain to members that are subsets of A. To this end, the subroutine branches into |A| computational paths, where each path corresponds to an element of A and a call to INTERLEAVE. This guarantees that an element is found for which Lemma 6 holds. Furthermore, for each $\ell \in A$, the algorithm successively resets the variable h, which was originally initialized with k, to the minimum of the current value of h and the return value of the associated recursive call to INTERLEAVE increased by $2 + w'(\{\ell, \ell'\})$ if ℓ crosses P' with $\{\ell,\ell'\},$ or increased by 1 otherwise. Thus, at each step, h equals kor it contains the minimum weight over all legitimate-agreement forests for \mathcal{S}' and \mathcal{T}' that have previously been considered. After at most k iterations, INTERLEAVE($\mathcal{S}, \mathcal{T}, k$) declares h+INTERLEAVE $(\mathcal{S}'', \mathcal{T}'', w'', k - h)$, where w'' is obtained from w by restricting its domain to members that are not subsets of A. Eventually, HYBRIDINTERLEAVE either returns $h(\mathcal{S}, \mathcal{T})$ if $h(\mathcal{S}', \mathcal{T}') = h < k$ and $h(\mathcal{S}'', \mathcal{T}'') < k - h$, or it returns k.

The pseudocode for HYBRIDINTERLEAVE is given below. The pseudocodes for the subtree, chain, and cluster reductions are given in Bordewich et al. (2007). Because of this and the description given earlier, we have omitted their respective pseudocodes.

$$\begin{aligned} & \textbf{Algorithm: HybridInterleave}(\mathcal{S}, \mathcal{T}, k) \\ & \textbf{procedure Interleave}(\mathcal{S}, \mathcal{T}, w, k) \\ & \textbf{if } k \leq 0 \\ & \textbf{then return } (k) \\ & (\mathcal{S}, \mathcal{T}, w) \leftarrow \text{SUBTREEREDUCTION}(\mathcal{S}, \mathcal{T}, w) \\ & \textbf{if } |\mathcal{L}(\mathcal{S})| \leq 3 \\ & \textbf{then return } (0) \\ & (\mathcal{S}, \mathcal{T}, w) \leftarrow \text{CHAINREDUCTION}(\mathcal{S}, \mathcal{T}, w) \\ & (\mathcal{S}', \mathcal{T}', w', \mathcal{S}'', \mathcal{T}'', w'') \leftarrow \text{CLUSTERREDUCTION}(\mathcal{S}, \mathcal{T}, w) \\ & h \leftarrow k \\ & \textbf{for each } \ell \in \mathcal{L}(\mathcal{S}') - \{\rho', \ell\} \text{ such that } \{\ell, \ell'\} \in \text{domain } w' \\ & \quad \left\{ \begin{aligned} & \textbf{s}' \leftarrow \mathcal{S}'[-\ell, \ell'] \\ & \textbf{then } \\ & \left\{ \begin{aligned} & \mathcal{S}' \leftarrow \mathcal{S}'[-\ell] \\ & \mathcal{T}' \leftarrow \mathcal{T}'[-\ell] \\ & h \leftarrow \min\{h, \text{INTERLEAVE}(\mathcal{S}', \mathcal{T}', w', h-1) + 1 \\ & \textbf{return } (h + \text{INTERLEAVE}(\mathcal{S}'', \mathcal{T}'', w'', k - h)) \end{aligned} \right. \\ & \textbf{main} \\ & P \leftarrow \emptyset \\ & w : P \rightarrow \mathbb{Z}^+ \end{aligned}$$

 $k \leftarrow \text{Interleave}(\mathcal{S}, \mathcal{T}, w, k)$

return (k)

Figure

Remark. Let X' be the label set of the two rooted binary phylogenetic trees that result from repeated applications of the subtree and chain reduction in the *first* call to INTERLEAVE. By Lemma 3, we freely assume for the rest of the paper that the algorithm directly returns k if |X'| > 14k.

To illustrate the reductions that are performed throughout the algorithm HYBRID-INTERLEAVE, consider Figure 6 and the call HYBRIDINTERLEAVE(S_1, T_1, k) with k > 2. The algorithm first replaces the common pendant subtree $S_1|\{1,2\}$ of S_1 and T_1 with a single leaf labeled a. Since the resulting two trees S_2 and T_2 cannot be reduced further under any of the three reductions, INTERLEAVE is recursively called, say for $S_2[-7]$ and $T_2[-7]$. Next interleaving comes into play as S_3 and T_3 have a common 3-chain (4, 5, 6) that is replaced with the 2-chain (b, c). Again, S_4 and T_4 are fully reduced and INTERLEAVE is called, say for $S_4[-3]$ and $T_4[-3]$. The two obtained trees S_5 and T_5 are identical and the algorithm ultimately returns h = 2 for the described computational path of the search tree.

We next establish the correctness of HYBRIDINTERLEAVE.

Theorem 7. Let S and T be a pair of rooted binary phylogenetic X-trees. Then the output of HYBRIDINTERLEAVE(S, T, k) is h(S, T) if and only if h(S, T) < k; otherwise it is k.

Proof. The proof is by induction on k. If k = 0, then INTERLEAVE immediately returns 0, and so the theorem holds. Now suppose that $k \ge 1$ and that the theorem holds whenever the input parameter is at most k - 1. Because of the structure of HYBRIDINTERLEAVE and Corollary 4, to establish this part of the induction, it suffices to show that the first call to the **for each** loop correctly returns h+INTERLEAVE(S'', T'', w'', k - h) with h = f(S', T') if and only if f(S', T') < k, otherwise with h = k.

By Lemma 6, there is an $\ell \in \mathcal{L}(\mathcal{S}') - \{\rho'\}$ such that one of the following holds:

 (a) If ℓ does not cross P, then

$$f(\mathcal{S}'[-\ell], \mathcal{T}'[-\ell]) = f(\mathcal{S}', \mathcal{T}') - 1.$$

(b) If ℓ crosses P with $\{\ell, \ell'\} \in P$, then

$$f(\mathcal{S}'[-\ell,\ell'],\mathcal{T}'[-\ell,\ell']) = f(\mathcal{S}',\mathcal{T}') - 2 - w(\{\ell,\ell'\}).$$

Moreover, by Lemma 5, for all $\ell \in \mathcal{L}(\mathcal{S}') - \{\rho'\}$, we have

$$f(\mathcal{S}'[-\ell], \mathcal{T}'[-\ell]) \ge f(\mathcal{S}', \mathcal{T}') - 1$$

if ℓ does not cross P and

$$f(\mathcal{S}'[-\ell,\ell']), \mathcal{T}'[-\ell,\ell']) \ge f(\mathcal{S}',\mathcal{T}') - 2 - w(\{\ell,\ell'\})$$

if ℓ crosses P with $\{\ell, \ell'\} \in P$. It now follows by the induction assumption and Lemma 5 that if $f(\mathcal{S}', \mathcal{T}') \geq k$, then the first call to the **for each** loop correctly returns k+INTERLEAVE $(\mathcal{S}'', \mathcal{T}'', w'', 0)$. Furthermore, by the induction assumption and Lemma 6, if $f(\mathcal{S}', \mathcal{T}') < k$, then the first call to the **for each** loop correctly returns h+INTERLEAVE $(\mathcal{S}'', \mathcal{T}'', w'', k - h)$, where $h = f(\mathcal{S}', \mathcal{T}')$. This completes the proof of the theorem.

We end this section by analyzing the running time of HYBRIDINTERLEAVE and comparing it with the time complexity of a previous implemented algorithm to solve MINI-MUM HYBRIDIZATION.

Proposition 8. Let S and T be two weighted rooted binary phylogenetic X-trees whose weighted 2-chain set P is empty. Furthermore, let k be an integer. Then the running time of HybridINTERLEAVE(S, T, k) is $O((14k)^k n^3)$, where n = |X|.

Proof. By repeatedly applying the subtree and chain reductions to S and T until no further reduction is possible, it follows from Lemma 3 that the leaf set size of the obtained weighted rooted binary phylogenetic X'-trees S' and T' is at most 14h(S, T).

Furthermore, while the subtree and chain reduction can be computed in $O(n^3)$ (Bordewich and Semple, 2007b), a single application of the cluster reduction results in an O(2n) algorithm. Thus, calling all three reductions takes time $O(n^3)$.

Since HYBRIDINTERLEAVE directly returns k if |X'| > 14k, we may assume that $|X'| \le 14k$. The remaining part of this proof is by induction on k. If k = 0, then the algorithm returns k in constant time. Now suppose that the running time of HYBRID-INTERLEAVE is $O((14k')^{k'}n^3)$ for all $0 \le k' < k$. Let A be a minimal common cluster of S' and T'. As $14k \ge |A|$, the algorithm makes at most 14k calls to INTERLEAVE for the tree pair S'|A and T'|A with parameter of at most k - 1. Thus the running time is $O(n^3 + 14k(14(k-1))^{k-1}n^3)$ which is $O((14k)^k n^3)$ as claimed.

Remark. The running time of HYBRIDINTERLEAVE given in Proposition 8 is an improvable upper bound since it only considers the kernelization of the two inputted phylogenetic trees and not applications of the subtree and chain reductions to previously reduced trees for which INTERLEAVE is recursively called from within the **for each** loop. As a comparison, $O((2 \cdot 14k)^k + n^3)$ is the theoretical worst-case running time of HY-BRIDNUMBER (Bordewich and Semple, 2007b). Thinking of the exhaustive search part of both algorithms HYBRIDINTERLEAVE and HYBRIDNUMBER as successively deleting a set of edges to calculate the hybridization number, the difference in the two running times is because the bounded search tree in HYBRIDINTERLEAVE is only based on the deletion of pendant edges of the trees under consideration and not all of the edges as in HYBRIDNUMBER. The fact that one needs only consider pendant edges follows from Lemmas 5 and 6. An indication of how much better the theoretical worst-case running time of HYBRIDINTERLEAVE could possibly be is highlighted in the next section when we compare the running times of both algorithms on a biological data set.

6 Experimental Results

To evaluate the performance of HYBRIDINTERLEAVE, we applied it to a grass (*Poaceae*) data set that has previously been used for running-time analyses in the context of calculating the hybridization number (Bordewich et al., 2007) and the rooted subtree prune and regraft distance (for details, see Section 7) which is frequently used to calculate the dissimilarities between two phylogenies for when reticulation is not assumed to be its major cause. The *Poaceae* data set was originally provided by the Grass Phylogeny Working Group (2001) and contains DNA sequences for six genetic loci, each with up to 65 taxa. Details about this data set and how a gene tree was reconstructed for each locus can be found in Bordewich et al. (2007) (and references therein). Species of the *Poaceae* family are subject to numerous natural hybridization events (Ellstrand et al., 1996). Therefore, the conflicting signals in this data set are more likely to be due to hybridization than to other processes causing inconsistencies.

For each of the 15 tree pairs, we restricted the two associated phylogenies to taxa that are common to both (second column of Table 1) and calculated the hybridization number of the resulting trees. The results are summarized in Table 1, where—beside the hybridization numbers—the running times for HYBRIDNUMBER and HYBRIDINTER-LEAVE are compared for each tree pair. A detailed description of the former algorithm, is given by Bordewich et al. (2007). Note that we reran HYBRIDNUMBER to guarantee consistency among the obtained running times for both algorithms. While HYBRIDNUM-BER computes the hybridization number for eight tree pairs within a couple of minutes, HYBRIDINTERLEAVE does so for all instances of the *Poaceae* data set and performs significantly faster. The latter algorithm successfully completes each program run in less than 8 minutes and calculates hybridization numbers as high as 19 for gene tree pairs with up to 46 taxa. This seems remarkably quick since HYBRIDNUMBER cannot calculate the exact solution for three tree pairs (ndhF and ITS, rbcL and ITS, and rpoC2and ITS) within 48 hours. The running time of HYBRIDINTERLEAVE mostly depends on the exhaustive search part of this algorithm since the reductions can be computed in polynomial-time. Clearly, the running time primarily decreases with an increase in the number of taxa that can be reduced by any of the three reductions. On the other hand, if the reductions have little effect because the trees only share a limited amount of common features such as subtrees, chains, or clusters, then the running time greatly increases with the hybridization number.

effe .totrees, ch. .ation number.

7 Concluding Remarks

In this paper, we presented the new algorithm HYBRIDINTERLEAVE that exactly calculates the hybridization number for two rooted binary phylogenetic trees. The algorithm can be applied to answer questions that consider the extent to which hybridization has influenced evolution and, therefore, shaped the current diversity of species. However, from a biological point of view, the results should carefully be interpreted since the algorithm is based on the assumption that hybridization is the only cause of gene tree inconsistencies. Moreover, it is possible that the real number of hybridization events for two trees is underestimated because HYBRIDINTERLEAVE minimizes this number and the true biological scenario might be less parsimonious. Of course, calculating a hybridization network that realizes this number is a desirable extension of our work. Indeed, given a maximum-acyclic-agreement forest, there is straightforward algorithm for doing this (Semple, 2007). However, such a network is typically not unique and it is part of ongoing research to implement an algorithm that calculates all possible hybridization networks that display two rooted binary phylogenetic trees and minimizes the hybridization number. Nevertheless, we believe that HYBRIDINTERLEAVE provides an important first step towards analyzing the occurrence of hybridization within a data set and, additionally, is remarkably quick.

We have shown that interleaving is an advantageous technique to speed-up the previously implemented fixed-parameter algorithm HYBRIDNUMBER. Referring back to the running time results summarized in Table 1, it is likely that HYBRIDINTERLEAVE can also compute the exact hybridization number in a reasonable short amount of time for problem instances that either contain bigger trees or have a greater hybridization number than those of the *Poaceae* data set. In conclusion, interleaving has proven to be most effective for our purpose of providing an exact algorithm to compute the hybridization number for two phylogenies of biologically relevant size, and we look forward to seeing whether interleaving has the same positive impact when applied to other fixed-parameter

tractable problems.

We end this paper with a remark on how interleaving can also be applied to calculate the rooted subtree prune and regraft (rSPR) distance. Loosely speaking, the graph-theoretic operation of rSPR cuts (prunes) a subtree and reattaches (regrafts) it to another part of the tree. The *rSPR distance* between two arbitrary rooted binary phylogenetic X-trees S and T is the smallest number of rSPR operations that transforms Sinto T. We denote this distance by $d_{rSPR}(S,T)$ and note that it is well-defined since one can always transform S into T via a sequence of single rSPR operations. Like MINIMUM HYBRIDIZATION, calculating $d_{rSPR}(S,T)$ is NP-hard and fixed-parameter tractable (Bordewich and Semple, 2004). Furthermore, the following theorem was central to obtaining these results.

Theorem 9. Let S and T be two rooted binary phylogenetic X-trees, and let m(S, T) denote the smallest number of elements among all agreement forests for S and T minus one. Then

$$d_{\mathrm{rSPR}}(\mathcal{S},\mathcal{T}) = m(\mathcal{S},\mathcal{T}).$$

Given the strong similarities between the characterizations of $h(S, \mathcal{T})$ and $d_{rSPR}(S, \mathcal{T})$ (see Theorems 1 and 9), it is not surprising that interleaving can also be applied to calculate the latter distance. However, while it is sufficient to exclusively consider 1-element subsets in the **for each** loop of HYBRIDINTERLEAVE, for calculating the rSPR distance, we need to iterate through all proper subsets of the label set under consideration, and thus, subsequently apply analogous subtree, chain, and cluster reductions to possibly more than one tree pair. This is due to the missing acyclic property in the context of calculating $d_{rSPR}(\mathcal{T}, \mathcal{T}')$. A detailed description of how interleaving can be applied in order to speed-up the computation of $d_{rSPR}(\mathcal{T}, \mathcal{T}')$ is given by Collins (2009).

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9 Disclosure Statement

No competing financial interests exist.

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Table 1: Running time comparison of HYBRIDINTERLEAVE with HYBRIDNUMBER (Bor-

dewich et al., 20	07) for the P	<i>Poaceae</i> data set	(Grass Phylogeny	Working	Group, 2001).
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Pairwise	combination	#Taxa	Hybridization number	RT^a of Hybrid Number	RT^a of Hybrid Interleave
ndhF	phyB	40	14	5.9 h	23 s
ndhF	rbcL	36	13	5.3 h	3 s
ndhF	rpoC2	34	12	13 h	6 s
ndhF	waxy	19	9	150 s	< 1 s
ndhF	ITS	46	19	>48 h	258 s
phyB	rbcL	21	4	$< 1 \mathrm{s}$	< 1 s
phyB	rpoC2	21	7	90 s	< 1 s
phyB	waxy	14	3	$< 1 \mathrm{s}$	< 1 s
phyB	ITS	30	8	10 s	< 1 s
rbcL	rpoC2	26	13	15.2 h	8 s
rbcL	waxy	12	7	132 s	$< 1 \mathrm{s}$
rbcL	ITS	29	14	>48 h	612 s
rpoC2	waxy	10	1	< 1 s	< 1 s
rpoC2	ITS	31	15	>48 h	$57 \mathrm{s}$
waxy	ITS	15	8	330 s	< 1 s

^a Running time (RT) on a 2.66 GHz CPU, 2 GB RAM machine measured in seconds (s) and hours (h), respectively.

Figure 1: A hybridization network \mathcal{H} with $h(\mathcal{H}) = 3$, and a rooted binary phylogenetic tree \mathcal{T} that is displayed by \mathcal{H} (indicated by the thicker arcs in \mathcal{H}).

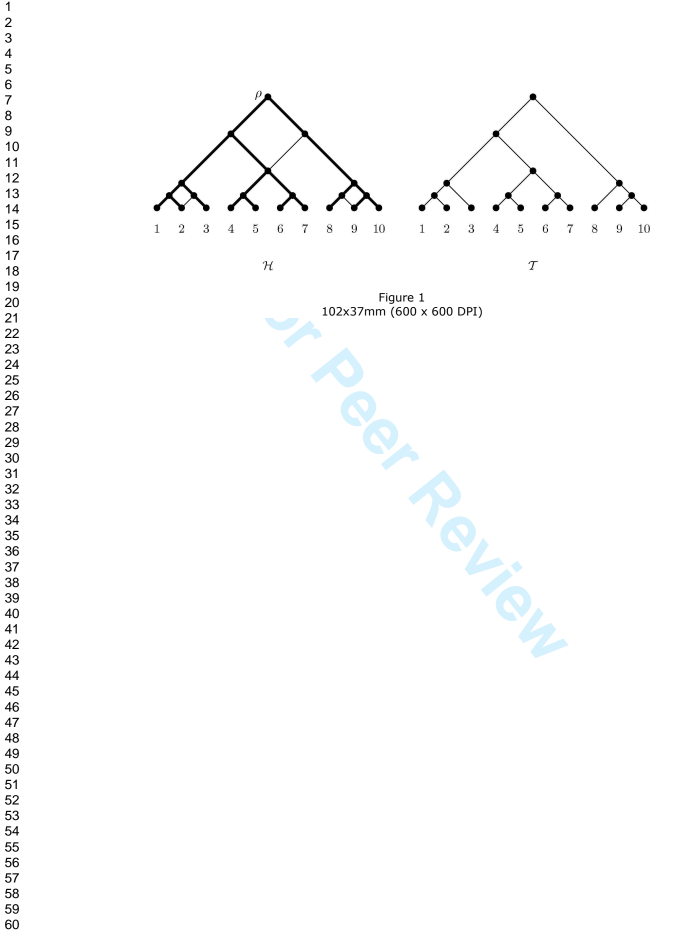


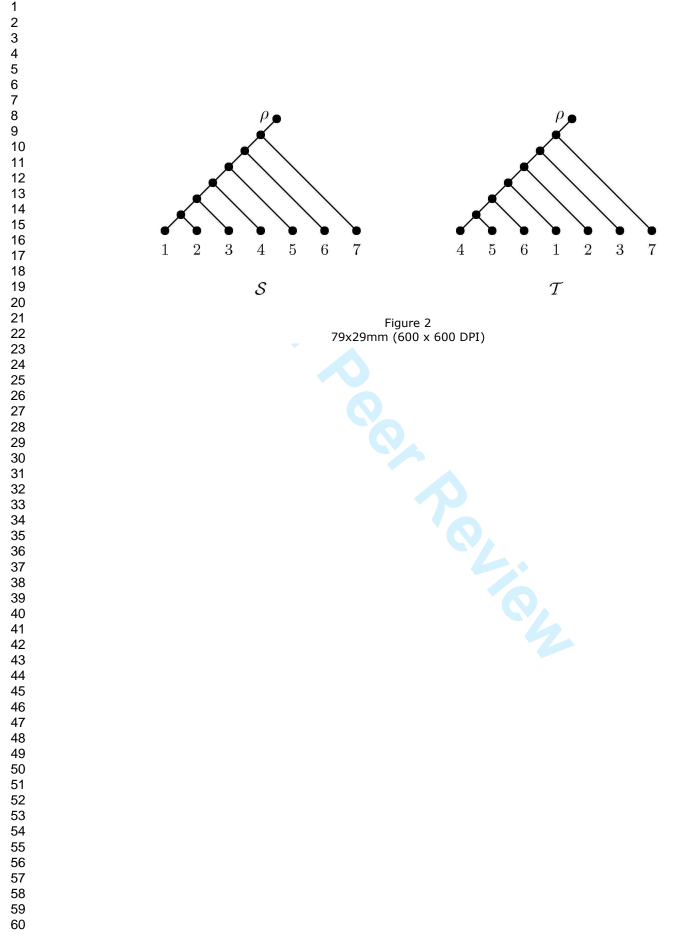
Figure 3: Two agreement forests \mathcal{F} and \mathcal{F}' and their associated digraphs $G_{\mathcal{F}}$ and $G_{\mathcal{F}'}$, respectively, for \mathcal{S} and \mathcal{T} shown in Figure 2.

Figure 4: Two rooted binary phylogenetic X-trees S' and T' that have been obtained from S and T depicted in Figure 2 by repeated applications of the chain reduction.

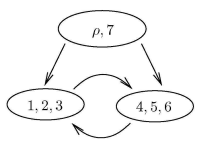
Figure 5: A cluster reduction applied to the two rooted binary phylogenetic trees S and T, where S_a and T_a have been obtained from S and T, respectively, by replacing the pendant subtree whose label set is A with a new leaf labeled a.

Figure 6: The intermediate tree pairs of a computational path in the search tree generated by a call to HYBRIDINTERLEAVE(S_1, T_1, k) with k > 2. Note that the cluster reduction is omitted here since the second tree pair which is returned from this reduction always consists of two identical trees. For details, see text.

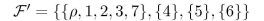




$$\mathcal{F} = \{\{
ho,7\},\{1,2,3\},\{4,5,6\}\}$$



 $G_{\mathcal{F}}$



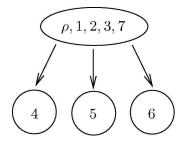




Figure 3 86x40mm (600 x 600 DPI)

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