SPECIES TREE INFECTION BY THE STAR METHOD, 
AND GENERALIZATIONS

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ABSTRACT. The multispecies coalescent model describes the generation of gene trees from a rooted metric species tree, and thus provides a framework for the inference of species trees from sampled gene trees. We prove that the STAR method of Liu et al., and generalizations of it, are statistically consistent methods of topological species tree inference under this model. We discuss the impact of gene tree sampling schemes for species tree inference using generalized STAR methods, and reinterpret the original STAR as a consensus method based on clades.

1. Introduction

While not always emphasized in practical analyses of biological sequences, the gene trees which standard phylogenetic methods infer may differ from the species tree relating the taxa from which the samples are taken. A frequent reason for this incongruence between gene trees and species trees is the population genetic effect of incomplete lineage sorting. Although this source of incongruence has been recognized for many years, recently methods have been proposed to bring a statistical treatment of it into data analysis, through the framework of the multispecies coalescent model (Rannala and Yang, 2003; Liu and Pearl, 2007; Kubatko et al., 2009; Liu et al., 2009; Heled and Drummond, 2010; Larget et al., 2010). However, the subject is still young, with both theoretical issues concerning some of the methods not fully established, and practical understanding of their appropriate application not yet developed.

Here we focus on the STAR method, of Liu et al. (2009), for inferring a rooted topological species tree from rooted topological gene trees. It is a fast approach to species tree construction that shows promise of good performance both in simulations (Liu et al., 2009) and in empirical studies (Lee et al., 2012).

Since STAR bases its inference on topological gene trees, it discards all metric information one might have on them. While at first this may seem wasteful, there are good reasons one might prefer such an approach. In practice, the gene trees are likely to be themselves inferred by standard phylogenetic methods, so their edge lengths are often considered to be less reliably known than their topological structure. Moreover, modeling the evolution of sequences on gene trees that were produced under the coalescent

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model requires reconciling time units on gene trees (reflecting total amounts of substitution), with time units on the species tree (reflecting population size and true time). This is usually resolved in part by assuming a molecular clock, thus forcing gene trees to be ultrametric. Since variations in base substitution rates along gene lineages can be large, and inferred metric gene trees are often far from ultrametric, it is unclear what impact this approach has on data analysis. The authors of the Bayesian inference software BEST have, in fact, warned that violations of the molecular clock could have ‘significant impacts’ (Castillo-Ramírez et al., 2010). By using only topological gene tree topologies, STAR circumvents such problems.

STAR proceeds by first combinatorially encoding each topological gene tree by a distance matrix. Averaging these over all gene trees produces a distance matrix that is then used to build a species tree. Finally, the metric information on the inferred species tree is discarded, so only its topology is retained. Though Liu et al. (2009) suggest UPGMA or Neighbor Joining (NJ) for this step, in fact any distance method, including ones based on optimality criteria such as Minimum Evolution, could be used. That an algorithmic tree construction process such as Neighbor Joining is fast accounts only in part for the speed of STAR. Its avoidance of calculating the theoretical gene tree probabilities that would be needed in either Maximum Likelihood or Bayesian analysis is also a major factor in its speed, so use of other distance methods can still be expected to have computational advantages.

Although not investigated by Liu et al. (2009), it is a rather surprising fact that the expected distance matrix obtained by STAR applied to the theoretical gene tree distribution under the multispecies coalescent model will exactly fit a rooted tree with the same topology as the species tree. That this is true is crucial to the claim that STAR is a statistically consistent method of inference. In this work we prove this claim, and as a consequence establish the statistical consistency of the STAR method when any of a large class of distance methods are used for the final selection of the species tree.

In the course of doing this, we also show that the method can be generalized into a parameterized family of STAR methods that are all consistent. The parameters specify a node numbering scheme which controls the way in which gene trees are encoded in distance matrices. (Although the particular node numbering scheme of the original method is referred to as a ‘ranking’ by Liu et al. (2009), we avoid that terminology as it conflicts with the common use of the term ‘ranked tree’ to indicate that internal nodes are assigned distinct values, often to indicate the temporal ordering of branching events in different parts of the tree (Semple and Steel, 2003).) The original STAR method arises from one particular choice of these parameters.

Our work also shows that under the coalescent the expected distance tables obtained in all generalized STAR methods are ultrametric. This is important for understanding the behavior of UPGMA, Neighbor Joining, or
any distance method when applied to the expected distance table, a point that was not addressed by Liu et al. (2009). It also suggests that in the final selection of the species tree from the average distances, it might be preferable to use a method that enforces ultrametricity.

In closing, we show that the standard STAR method can be viewed as using only the clade structure of the gene trees. This observation allows a reinterpretation of the STAR distance as one by which taxa are viewed as close if they appear together in many clades across the gene trees. The STAR method can thus be viewed as a type of greedy consensus method using the clade structure of gene trees to infer the species tree. That it is a consensus method attuned in particular to the species tree/gene tree problem is the import of the fact that STAR is consistent under the coalescent model.

Although our emphasis in this paper is on theory, we also discuss some possible implications for data analysis, focusing on various sampling schemes and how generalizations of the STAR method might possibly be used to gain a measure of confidence in species tree estimates. Additional work is needed to further develop the suggestions here.

2. THE STANDARD STAR NODE NUMBERING SCHEME AND VARIANTS

The encoding of an $n$-taxon binary rooted gene tree topology by a distance matrix in STAR is accomplished through first associating numbers to each node of the tree. The standard node numbering scheme used by STAR is as follows: Assign $n$ to the root of the tree. Then if an internal node has been assigned a number $m$, assign to each of its children that are internal nodes the number $m - 1$. These numbers are then interpreted as distances from the leaves in an ultrametric tree, so that the distance between two leaves is twice the number assigned to their most recent common ancestor. In Figure 1, for example, the distance between genes $A$ and $C$ is 6 on the left tree, and 10 on the right tree.

An alternative viewpoint is that the standard STAR metrization of a gene tree is obtained from the rooted gene tree topology by assigning all internal edges length 1, and then choosing lengths of pendant edges so that all leaves are distance $n$ from the root.

It is immediately clear that some trivial variations on this numbering scheme will have no effect on the output of STAR. For instance, suppose instead of assigning the root the number $n$, one assigned to it any number $\ell \geq n - 2$, but otherwise followed the standard scheme. Then all internal branches would still have length 1, and pendant ones would have non-negative length, regardless of the topology of the gene tree. Indeed pendant edge lengths would only change by the addition of $\ell - n$. The net effect on the pairwise distance matrix for each gene tree is to add $2(\ell - n)$ to each off-diagonal entry. Thus the empirical average of these matrices over a sample of gene trees, and the theoretical expected distance matrix change in the same way. (Note that these distance matrices will still have non-negative off-diagonal entries.)
Figure 1. Two 6-taxon gene trees with a STAR node-numbering. Both trees are drawn ultrametrically according to the standard STAR numbering with \((a_0, a_1, a_2, a_3, a_4) = (6, 5, 4, 3, 2)\). Note that only the caterpillar tree shape (left) has distinct numberings for each node. For a generalized node numbering, the \(a_i\) can be any non-negative decreasing sequence with at least two distinct terms. Node numbers are interpreted as remetrizing the gene tree, by specifying distances of the nodes from their descendant leaves.

It is easy to see that this will have no effect on the topology of the species tree output by methods such as UPGMA and NJ. Similarly, assuming one chooses a sufficiently large number to assign to the root, one could number descendant nodes by any constant decrement from their parents, with no effect on the output of the topological species tree by UPGMA or NJ, since this would merely change the average distance matrix by a rescaling and addition of a constant.

Thus assuming the output of standard STAR from the theoretical gene tree distribution does agree topologically with the species tree, there are certainly some variations on the numbering scheme that will have the same property. Less obvious variations of node numberings that can also be used for species tree inference in a generalized STAR method are the focus of the next section.

By way of contrast, one might consider the following seemingly natural node numbering: Assign to each node the count of its leaf descendants. That this ‘descendant-count’ numbering scheme behaves differently from the STAR scheme is seen by considering the 5-taxon caterpillar species tree

\[
\sigma = (((a, b): x, c): y, d): z, c),
\]
where \( x, y, z \) are in coalescent units. Taking \( x = z = \infty \) and \( y = 0 \), the only gene trees with non-zero probability under the coalescent model are

\[
T_1 = (((a, b), c), d), \quad \mathbb{P}_\sigma(T_1) = 1/3,
\]

\[
T_2 = (((a, b), d), c), \quad \mathbb{P}_\sigma(T_2) = 1/3,
\]

\[
T_3 = (((a, b), (c, d)), e), \quad \mathbb{P}_\sigma(T_3) = 1/3.
\]

Using these probabilities as weights in averaging the distance matrices

\[
D_{T_1} = \begin{pmatrix}
0 & 4 & 6 & 8 & 10 \\
4 & 0 & 6 & 8 & 10 \\
6 & 6 & 0 & 8 & 10 \\
8 & 8 & 8 & 0 & 10 \\
10 & 10 & 10 & 10 & 0
\end{pmatrix}, \quad D_{T_2} = \begin{pmatrix}
0 & 4 & 8 & 6 & 10 \\
4 & 0 & 8 & 6 & 10 \\
8 & 8 & 0 & 8 & 10 \\
6 & 6 & 0 & 8 & 10 \\
10 & 10 & 10 & 10 & 0
\end{pmatrix},
\]

\[
D_{T_3} = \begin{pmatrix}
0 & 4 & 8 & 8 & 10 \\
4 & 0 & 8 & 8 & 10 \\
8 & 8 & 0 & 4 & 10 \\
8 & 8 & 4 & 0 & 10 \\
10 & 10 & 10 & 10 & 0
\end{pmatrix},
\]

we see

\[
\mathbb{E}_\sigma(D_T) = \begin{pmatrix}
0 & 4 & 22/3 & 22/3 & 10 \\
4 & 0 & 22/3 & 22/3 & 10 \\
22/3 & 22/3 & 20/3 & 0 & 10 \\
22/3 & 22/3 & 20/3 & 0 & 10 \\
10 & 10 & 10 & 10 & 0
\end{pmatrix}.
\]

This expected distance matrix exactly fits the rooted tree

\[
(((a:2, b:2):5/3, (c:10/3, d:10/3):1/3):4/3, e:5)
\]

which is topologically different from the species tree, even if both are treated as unrooted. Similar examples with positive and finite edge lengths can easily be constructed, by taking \( x, z \) to be large while \( y \) is small, since the expected distance matrix depends continuously on the edge lengths.

If, instead, the standard STAR numbering were used in the above example, \( D_{T_1} \) and \( D_{T_2} \) would be unchanged, but in \( D_{T_3} \) all entries of 4 would be replaced by 6. How this effects \( \mathbb{E}_\sigma(D_T) \), and thus changes the tree it fits, we leave as an easy exercise for the reader.

### 3. Consistent node numbering schemes

We next show that any node numbering scheme with several simple properties, when used in STAR with UPGMA, NJ, or any well-behaved distance method for the selection of the species tree, leads to consistent inference. By ‘well-behaved’ we simply mean one that when applied to any distance table in some neighborhood of one exactly fitting \( T \) will return the topology of \( T \).

The properties we require for a node numbering scheme are:
(1) The number assigned to any internal node on a gene tree depends only on the count of nodes between it and the root in the topological gene tree. Thus the node numbers on all gene trees come from a common node numbering sequence \( a_0, a_1, \ldots, a_{n-2} \), with \( a_0 \) assigned to all roots, \( a_1 \) to their children, and more generally \( a_i \) to all internal nodes at depth \( i \) from the root.

(2) A non-negative number is assigned to every node, with the number assigned to a child always less than or equal to that assigned to its parent, with at least one instance of strict inequality. That is, \( a_0 \geq a_1 \geq \cdots \geq a_{n-2} \geq a_{n-1} = 0 \) with some \( a_\ell > a_{\ell+1} \) for some \( \ell \).

Equivalently, one could define an edge length sequence \( b_1, b_2, \ldots, b_{n-1} \), where \( b_i \geq 0 \), and at least one \( b_i > 0 \), and then assign length \( b_i \) to all internal edges at depth \( i \) from the root. If a pendant edge is at depth \( i \), it is assigned the length \( \sum_{j \geq i} b_j \). This is related to a node numbering sequence by defining \( b_i = a_{i-1} - a_i \).

Both of these properties are natural. Property (2) ensures all internal branch lengths are non-negative, with at least one positive, while property (1) ensures a connection to the number of coalescent events in a gene’s lineage that occur above (i.e., temporally before) the specified node.

Note that the standard STAR scheme is simply the special choice of the node numbering sequence \( n, n-1, n-2, \ldots, 2 \), or, equivalently, of the edge length sequence \( 1, 1, 1, \ldots, 2 \). On the other hand, the descendant-count scheme mentioned in the previous section fails to satisfy property (1). Though it is tied to counting coalescent events, that scheme counts events below the node.

We consider first a generalized STAR method where for each gene exactly one individual is sampled per taxon. Extensions to sampling multiple individuals will be discussed later.

**Definition 3.1.** A \( n \times n \) numerical matrix, with rows and columns indexed by an \( n \)-element set of taxa \( \mathcal{X} \), weakly fits a topological tree \( \psi \) on \( \mathcal{X} \) if there is some assignment of non-negative lengths to the edges of \( \psi \) such that the resulting pairwise tree distances between leaves give the entries of the matrix. If the edge lengths are positive, then we say the matrix strongly fits \( \psi \). If the assignment of edge lengths yields an ultrametric tree, we say the matrix fits \( \psi \) ultrametrically.

For a complete presentation of the multispecies coalescent model, which describes the generation of gene trees from a metric species tree, we refer to earlier works, for instance (Allman et al., 2011b). The primary feature of the coalescent model that we use is the exchangeability of lineages: for a collection of lineages present in a population at a particular time, the probability of any pattern of coalescence (moving backwards in time) of these lineages is the same for all permutations of the lineages.
We denote a species tree on a set of taxa $\mathcal{X}$ by $\sigma = (\psi, \lambda)$, where $\psi$ is a rooted topological tree and $\lambda$ is a vector of edge lengths measured in coalescent units. We use lower case letters $i, j, k$ to denote taxa, and the corresponding upper case letters $I, J, K$ to denote gene samples from those taxa. For any $\mathcal{Y} \subseteq \mathcal{X}$, by the most recent common ancestor of $\mathcal{Y}$, MRCA($\mathcal{Y}$), we mean the vertex on $\psi$ that is ancestral to all elements of $\mathcal{Y}$, and a descendant of any other vertex ancestral to all of $\mathcal{Y}$. Thus $\mathcal{Y}$ is always a subset of the set of descendants of MRCA($\mathcal{Y}$). An MRCA on a gene tree is defined similarly.

Our main result is the following.

**Theorem 3.2.** Under the multispecies coalescent model on a metric species tree $\sigma = (\psi, \lambda)$, let $P_{\sigma}(T)$ denote the probability of a rooted topological gene tree $T$. For a node numbering sequence satisfying properties (1) and (2), let $D_T$ denote the pairwise distance matrix obtained from applying the sequence to $T$. Then for all $\lambda_i \in (0, \infty)$ the expected value of this matrix,

$$E_{\sigma}(D_T) = \sum_T P_{\sigma}(T)D_T$$

is a pairwise distance matrix that strongly fits $\psi$ ultrametrically.

Of course the pairwise distance matrix $E_{\sigma}(D_T)$ of this theorem does not generally match the pairwise distances on the species tree $\sigma$.

**Proof of Theorem 3.2.** For a binary topological species tree $\psi$ with positive edge lengths $\lambda_i$, it is enough to show that, for any three taxa $i, j, k$, if $\psi$ displays the rooted triple $(\langle i, j \rangle, k)$, then with $D = E_{\sigma}(D_T)$,

$$D(i, j) < D(i, k) = D(j, k).$$

This is simply the 3-point condition for a distance to determine an ultrametric tree (Semple and Steel, 2003).

So suppose $\psi$ displays $(\langle i, j \rangle, k)$, and let $w = \text{MRCA}([i, j, k])$ on $\sigma$. Using shorthand such as $C(I, J) < w$ to mean the coalescence of the (gene) lineages of $I$ and $J$ occurs below node $w$ in the species tree, and $P_{\sigma}(T, C(I, J) < w)$ to mean the joint probability of the gene tree $T$ and the event $C(I, J) < w$, we see

$$D(i, j) = \sum_T P_{\sigma}(T)D_T(i, j)$$

$$= \sum_T P_{\sigma}(T, C(I, J) < w)D_T(i, j) + \sum_T P_{\sigma}(T, C(I, J) > w)D_T(i, j).$$

But we claim

$$\sum_T P_{\sigma}(T, C(I, J) < w)D_T(i, j) < \sum_T P_{\sigma}(T, C(I, J) < w)D_T(i, k).$$

To see this, first note that if the lineages of $I, J$ coalesce below $w$ and a gene tree $T$ is observed, then $T$ must display the rooted triple $(\langle I, J \rangle, K)$. But
for such gene trees, $D_T(i,j) ≤ D_T(i,k)$. Thus a weak version of inequality (2) holds termwise.

Note now that under the coalescent model on a species tree with finite edge lengths, if $C(I,J) < w$, then every labeled gene tree $T$ which displays $((I,J),K)$ is realizable. In particular, since $a_ℓ > a_{ℓ+1}$, and there is a gene tree displaying $((I,J),K)$ for which MRCA($I,J$) is at depth $ℓ+1$ or greater, and the MRCA($I,K$) is at depth $ℓ$ or less, one has $D_T(i,j) < D_T(i,k)$ for some gene tree $T$ with $P_σ(T,C(I,J) < w) > 0$. Thus at least one pair of corresponding terms in inequality (2) satisfies a strict inequality, and (2) is established.

We claim also that

\[ \sum_T P_σ(T,C(I,J) > w)D_T(i,j) = \sum_T P_σ(T,C(I,J) > w)D_T(i,k). \]

Indeed, up to multiplication by a positive constant, these sums respectively give the conditional expectation of $D(i,j)$ and $D(i,k)$, given that the lineages of $I,J,K$ are distinct at $w$. But by property (1) of the numbering scheme, and the exchangeability of lineages under the coalescent, these conditional expectations are the same.

From equations (1), (2), and (3) the inequality $D(i,j) < D(i,k)$ now follows.

To see that $D(i,k) = D(j,k)$, observe

\[
D(i,k) = \sum_T P_σ(T)D_T(i,k) \\
= \sum_T P_σ(T,C(I,J) < w)D_T(i,k) + \sum_T P_σ(T,C(I,J) > w)D_T(i,k).
\]

However

\[ P_σ(T,C(I,J) < w)D_T(i,k) = P_σ(T,C(I,J) < w)D_T(j,k), \]

since if the probability appearing in this equation is non-zero then the MRCA of $I,K$ on $T$ is the same as the MRCA of $J,K$. Moreover,

\[ \sum_T P_σ(T,C(I,J) > w)D_T(i,k) = \sum_T P_σ(T,C(I,J) > w)D_T(j,k) \]

by an exchangeability argument similar to that used in establishing inequality (3). Thus $D(i,k) = D(j,k)$, and the 3-point condition is established.

For non-binary $ψ$, one must also check that if $ψ$ displays the unresolved 3-taxon subtree $(i,j,k)$, then

\[ D(i,j) = D(i,k) = D(j,k). \]

This is done using exchangeability of lineages, as in the justification for equation (3).

\[ \square \]

**Corollary 3.3.** Consider the generalized STAR method using any node numbering sequence satisfying properties (1) and (2), and any method of tree selection that from a distance table strongly fitting a binary tree returns
that tree, and whose output is continuous at such tables. Then under the multispecies coalescent model on \( \sigma = (\psi, \lambda) \), with \( \psi \) binary and \( \lambda_i > 0 \), the method is statistically consistent for inferring the species tree topology \( \psi \).

Proof. Let \( N \) be the number of sampled gene trees, and \( \epsilon > 0 \). By the law of large numbers, as \( N \to \infty \), the probability that the average distance matrix differs from \( \mathbb{E}_\sigma(D_T) \) by more than \( \epsilon \) in any entry approaches 0. Thus by continuity of the distance method of tree selection, as \( N \to \infty \) the probability that the inferred tree will have the same topology as that from \( \mathbb{E}_\sigma(D_T), \psi \), approaches 1.

\[ \square \]

Atteson (1997) showed that the Neighbor Joining tree construction algorithm satisfies the continuity hypothesis of this corollary. A proof that UPGMA is continuous is straightforward.

Example 3.4. As an extreme example of a node numbering scheme leading to consistent inference, for \( n \) species let \( a_i = 2 \) for \( 0 \leq i < n - 2 \) and \( a_{n-2} = 1 \). Interpreting the \( a_i \) values as distances from the leaves, this scheme converts every non-caterpillar gene tree into a completely unresolved tree, and every caterpillar gene tree into a gene tree with one non-trivial clade of two leaves. In effect, the resulting generalized STAR method discards all information gathered from non-caterpillar gene trees yet, by Theorem 3.2 the species tree topology can still be recovered. Thus it is possible, in principle, to reconstruct the species tree topology using only caterpillar gene trees as input, even if the species tree is not a caterpillar. Of course, we do not recommend this for practical inference.

It is natural to focus especially on node numbering sequences that are strictly decreasing. Indeed, if all branch lengths on \( \sigma = (\psi, \lambda) \) are very long, then with high probability a finite sample of topological gene trees will include only those matching \( \psi \). If a node numbering is not strictly decreasing, however, then an estimate of \( \mathbb{E}_\sigma(D_T) \) from the sample would give \( D_\psi \), which may only weakly fit \( \psi \), leading to a poorly-resolved inferred species tree under STAR.

The next theorem shows that strictly decreasing node numbering sequences lead to better behavior of generalized STAR methods, in the sense that longer internal branches on the species tree do not make it more difficult to infer a fully-resolved species tree from a fixed sample size of gene trees. Since longer edges in a species tree increase the probability of the gene trees showing the associated split, this is intuitively desirable, and would certainly be a useful characteristic for data analysis.

Recall now that the probability of any gene tree under the coalescent is a polynomial in the transformed species tree branch lengths, \( \exp(-\lambda_i) \), as is explained, for instance, by Allman et al. (2011b). Thus for any collection \( \{D_T\} \) of matrices associated to gene trees (whether or not it arises from a node numbering sequence), the expected value \( \mathbb{E}_\sigma(D_T) \) also has entries that
are polynomial in the $\exp(-\lambda_i)$. As a result, the expected value also makes sense for a branch length of $\infty$, by setting the transformed branch length to 0. (Treating a branch length of $\infty$ in this way is equivalent to taking a limit as the branch length goes to $\infty$.) While an infinite branch length of course has no real biological meaning, we will allow branch lengths in $(0, \infty] = (0, \infty) \cup \{\infty\}$, both to simplify the presentation of arguments, and to easily describe behavior as branches grow long.

Note also that the strong fitting of $\psi$ by $E(\sigma(D_T))$ for finite $\lambda$ is expressible by equalities and strict inequalities in the matrix entries, arising from the 3-point conditions. By continuity then, if some $\lambda_i \to \infty$ the same equalities and non-strict versions of the inequalities must hold. That is, if branch lengths are in $(0, \infty]$, then $E(\sigma(D_T))$ will certainly weakly fit $\psi$. By requiring that the node numbering is strictly decreasing, however, this can be improved.

**Theorem 3.5.** Suppose a node numbering sequence is strictly decreasing. Then for any $\psi$ and all $\lambda_i \in (0, \infty]$, the expected distance matrix $E(\sigma(D_T))$ strongly fits $\psi$, ultrametrically.

Moreover, if the node numbering sequence is not strictly decreasing, then there is a binary $\psi$ and choices of $\lambda_i \in (0, \infty]$ for which $E(\sigma(D_T))$ only weakly fits $\psi$.

**Proof.** The proof of the first claim follows the argument of Theorem 3.2. The primary modification is in the justification of inequality (2), when some of the internal edge lengths are $\lambda_i = \infty$. But since a strictly decreasing node numbering implies $D_T(i, j) < D_T(i, k)$ for all gene trees $T$ displaying $((I, J), K)$, the inequality is immediately clear.

For the second claim, suppose $a_\ell = a_{\ell+1}$ and pick any binary species tree $\psi$ which has an internal edge between nodes $u$ of depth $\ell$ and $v$ of depth $\ell + 1$. Assign edge lengths of $\infty$ to all edges incident to the ancestors of $u$, to the edge descending from $u$ that is not incident to $v$, and to the two edges descending from $v$. Pick taxa $i, j$ that are descendants of $v$ through the two different edges, and taxon $k$ that is a descendant of $u$ but not of $v$. Then regardless of the other edge lengths, the only gene trees realizable under the coalescent will be those with one of MRCA($I, J$), MRCA($I, K$), and MRCA($J, K$) at depth $\ell + 1$, say MRCA($I, J$), and thus MRCA($I, K$) = MRCA($J, K$) located at depth $\ell$ as the parent of MRCA($I, J$). Since $a_\ell = a_{\ell+1}$, we have $E(\sigma(D_T)(i, j) = E(\sigma(D_T)(i, k) = E(\sigma(D_T)(j, k) = 2a_\ell$ are all equal, so $E(\sigma(D_T))$ cannot strongly fit $\psi$. □

**Remark.** While we have established that properties (1) and (2) are sufficient for statistical consistency of STAR, we have not shown they are necessary. In fact, addressing this question for the full STAR process which includes the selection of the species tree by some algorithm or optimality criterion would be difficult, since even if an expected distance matrix does not exactly fit a tree, the fitting processes in the distance method may overcome the misfit.
Avoiding the impact of the final tree selection step, one can investigate the existence of alternative schemes in which distance matrices might be assigned to gene trees in such a way that for any species tree the expected distance matrix under the coalescent fits the topological species tree. Preliminary investigations indicate that at least for small trees such assignments exist that do not arise from generalized STAR numbering schemes.

The theorems above were stated and proved under a gene tree sampling scheme in which one individual is sampled per taxon. We next consider extensions to multi-individual sampling.

In the case of a sampling scheme in which \( k_i \) individuals are sampled from taxon \( i \) for every gene, a simple device for extending STAR is the extended species trees: At leaf \( i \) on the original species tree attach \( k_i \) descendant edges in a multifurcation, leading to new taxa \( i_j, 1 \leq j \leq k_i \). These new pendant edges need not be assigned lengths, but all edges arising from the original species tree retain their lengths.

The coalescent model on the original species tree with multiple individuals sampled per taxon then produces exactly the same distribution of gene trees as the coalescent on the extended species tree with one sample per taxon. Thus Theorem 3.2 applies.

Since this extended species tree is not binary if any \( k_i > 2 \), Corollary 3.3 does not apply directly. However, the procedure proposed by Liu et al. (2009) for multi-individual sampling is to first average over these individuals so that a distance matrix is obtained relating the original taxa. We sketch an argument that this leads to consistent inference. Since the average distance table for the gene trees will approximate the expected one, which by Theorem 3.2 strongly fits the extended species tree ultrametrically, so will the result of further averaging this empirical distance table over its image under all permutations of individuals within each taxon. In this averaged table, distances between individuals in different taxa depend only on the two taxa, and not the individuals. Deleting all but one individual per taxon from this table will then yield a table that in expectation strongly fits the original species tree, and whose entries are those calculated by the procedure of Liu et al. (2009). Thus as long as the original species tree is binary, one still has a consistent inference scheme. For an example of an empirical study that uses STAR with a variable number of individuals per species, see Lee et al. (2012).

An even more general sampling scheme might vary the number of individuals sampled by both taxon and gene. For instance if taxon \( i \) is sampled \( k_{ij} \) times for locus \( j \), we could define a sampling scheme for locus \( j \) as \( k_j = (k_{1j}, \ldots, k_{nj}) \), where \( n \) is the number of species. Letting \( \mathcal{K} = \{k\} \) be the finite set of all possible schemes used in a particular study design, if each locus has a sampling scheme independently chosen from some distribution on \( \mathcal{K} \), then as the number of loci approaches infinity, the average empirical distance table for each sampling scheme approaches the expected distance.
table for that sampling scheme by the Law of Large Numbers. Thus, the expected distance table over $\mathcal{K}$ is a weighted average over the expected distance tables for each $k \in \mathcal{K}$, and hence also ultrametrically fits the species tree by the 3-point condition. Thus the multiple-allele version of STAR is consistent method of inference even with different numbers of alleles sampled at different loci.

Finally, one could generalize further to allow different STAR node numberings for different sampling schemes. More precisely, if $k_j \in \mathcal{K}$ specifies both a sampling scheme and numbering scheme for locus $j$, chosen according to some distribution on $\mathcal{K}$, then the STAR method will still be statistically consistent. Of course with such an approach an increased number of loci are likely to be needed for observed average distances to be close to their expected values.

4. Finite samples of loci

The statistical consistency of STAR methods, as established in the preceding section, is a statement about asymptotic behavior as the number of sampled loci approaches infinity. In this section we collect some observations on possible pitfalls of naively applying STAR when the number of sampled loci is small.

We begin with issues concerning multiple samples of individuals within taxa, where the number of individuals varies with the locus.

First, imagine a rather extreme sampling scheme in which $N$ loci are sampled, but that at locus $j$, taxon $i$ is sampled from $i + j$ individuals. Thus each sampling scheme is used for exactly one locus. The statistical consistency of STAR in this situation is not helpful to us, since without multiple uses of every sampling scheme we do not expect to see asymptotic behavior. The Law of Large Numbers does not apply, and the observed average distance table might be quite different from its expectation.

If in a real study the number of sampling schemes used at different loci is very large in comparison to the number of loci, an appeal to the Law of Large Numbers may be similarly questionable. For example, if each of $n$ species has between 1 and 3 individuals sampled at each locus, then there are potentially $3^n$ sampling schemes. We would need to sample enough loci to ensure that each of these exponentially many sampling schemes is used many times for the asymptotic claim of consistency to give us much confidence in an analysis.

If, on the other hand, a study is designed for 3 individuals sampled per species and sequenced at all loci, a small amount of missing data might result in only a few of the $3^n$ sampling schemes being used, with each used repeatedly. Especially if the data were missing at random, it might be reasonable to use the multiple-allele versions of STAR.

As another example of potentially problematic sampling, imagine sampling a large number of loci with only one individual per taxon, and a
single locus with one individual for most taxa, but many individuals for one taxon. (Such a data set is not farfetched, especially if it were built from data originally collected for other studies.) Then using the standard STAR numbering, the tree for this last locus will have a much larger number assigned to its root. If there is substantial incomplete lineage sorting between the single-sample taxa and the multi-sample one, then on this tree internode distances between coalescent events involving only single-sample taxa will be larger than for the other trees. Thus it appears that it might be desirable to downweight this tree in the distance averaging, to avoid overemphasizing the evidence for relationships between the single-sample taxa that it offers. On the other hand, if there is little incomplete lineage sorting between the single-sample taxa and the multi-sample one, then downweighting will result in an undesirable deemphasis of the evidence of relationships between the single-sampled taxa given by this locus. It is simply not clear how the anomalously-sampled locus could be used without potentially biasing the analysis.

We note too that for finite sample sizes, the standard STAR numbering scheme and its variations may have biases for tree shape. To illustrate this, consider the case of sampling a single individual per taxon for four taxa.

Now suppose UPGMA is applied to the average distance table using the standard STAR numbering with \((a_0, a_1, a_2) = (4, 3, 2)\), with ties being resolved randomly. If there are exactly two input gene trees, then STAR-UPGMA will return a symmetric tree in exactly the following cases:

(i) both input trees are symmetric;
(ii) one input tree is symmetric and one is asymmetric, and they share a cherry (a 2-taxon clade), in which case a symmetric tree is returned with probability 1/2; and
(iii) both input trees are asymmetric, and their cherries do not have overlapping taxa, in which case a symmetric tree is returned with probability 2/3.

In case (iii), for instance, if the input trees are \(((AB)C)D)\) and \(((CD)B)A)\), then the expected distance table averaged over both loci is

\[
E(D_T) = \begin{pmatrix}
0 & 6 & 7 & 8 \\
6 & 0 & 6 & 7 \\
7 & 6 & 0 & 6 \\
8 & 7 & 6 & 0 \\
\end{pmatrix},
\]

which results in an asymmetric tree if B and C are clustered first (which occurs with probability 1/3), and otherwise results in a symmetric tree.

Assuming that gene trees evolve on an unresolved species tree, unlabeled gene trees have a 1/3 probability of being symmetric and 2/3 probability
of being asymmetric under the coalescent. The probability that STAR-UPGMA returns a symmetric tree in this situation is therefore
\[
\frac{1}{3} \cdot \frac{1}{3} + 2 \cdot \frac{1}{3} \cdot \frac{2}{3} \cdot \frac{1}{2} + \frac{2}{3} \cdot \frac{1}{3} \cdot \frac{2}{3} \cdot \frac{2}{3} = \frac{19}{81} < \frac{1}{3},
\]
indicating that there is some bias against symmetric trees in this case. (To understand the second summand in the expression on the left, for example, notice that the two trees are symmetric and asymmetric with probability \(2 \cdot \left(\frac{1}{3}\right)\left(\frac{2}{3}\right)\), that these two trees share a cherry with probability \(\frac{1}{3}\), and that UPGMA will return the symmetric tree with probability \(\frac{1}{2}\).)

While these computations show bias is present, we note that this analysis has discarded the edge lengths on the UPGMA tree, even though they carry information about when a tree is close to a different topological structure.

Now it is possible that numbering schemes can be chosen to reduce bias when only finitely many loci are sampled. For example, for the numbering scheme \((a_0, a_1, a_2) = (4, 3, 1)\), case (iii) always results in STAR returning a balanced tree, and the total probability of returning a symmetric tree is \(21/81\), suggesting less bias than the standard STAR numbering, at least for this example.

Whether topological bias of STAR for small samples can be reduced for all species trees simultaneously, or at least on average under some model of species tree generation, is an interesting topic for future investigation.

5. STAR and clades

The standard node numbering scheme, in which all internal edges on gene trees are given length 1, produces distances on remetrized gene trees that has an interpretation regarding clades on gene trees. In this section we develop this connection.

As a consequence, we recover a previously known result that the topological rooted species tree can be identified from the collection of gene clade probabilities \(\text{[Allman et al. 2011a]}\). Additionally, with this viewpoint, we obtain an efficient algorithm for the identification.

This connection between STAR and clades makes clear that whatever information STAR uses for inference is found in the clade probabilities, and not in the more detailed distribution of gene trees.

Consider a gene tree \(T\) on a collection of taxa \(\mathcal{X}\). By a clade on \(T\) we mean a subset \(\mathcal{C} \subseteq \mathcal{X}\) for which the set of descendants of MRCA(\(\mathcal{C}\)) is precisely \(\mathcal{C}\). There is thus a bijection between nodes \(v\) of \(T\) and clades \(\mathcal{C}\), by which \(v = \text{MRCA}(\mathcal{C})\). The trivial clades are \(\mathcal{X}\), with MRCA at the root of the tree, and singleton subsets of \(\mathcal{X}\), with MRCA at the leaves. The trivial clades occur on all gene trees on \(\mathcal{X}\).

Given any two leaves \(a, b\) of \(T\), let \(H_{a,b}\) denote the set of non-trivial clades of \(T\) which contain both \(a\) and \(b\), and let \(c_T(a, b) = |H_{a,b}|\). Elements of \(H_{a,b}\) correspond to the non-root vertices of \(T\) lying on the path from the root to the MRCA(\(\{a, b\}\)). Thus \(c_T(a, b)\) is simply the distance from the root to
the MRCA({\(a,b\)}) in \(T\) remetrized by the standard STAR numbering. Since this tree is ultrametric, with all leaves \(n = |X|\) from the root, one finds

\[
c_T(a,b) = n - \frac{D_T(a,b)}{2}.
\]

This shows that, at least on individual gene trees, the distances used in STAR are essentially counts of clades, with gene samples being judged closer when they occur together in more clades.

Now fixing a species tree \(\sigma\), the coalescent model determines gene tree probabilities \(P_{\sigma}(T)\). If \(C\) is a non-trivial clade, and \(P_{\sigma}(C)\) denotes the probability of clade \(C\), then

\[
P_{\sigma}(C) = \sum_{T \text{ displaying } C} P_{\sigma}(T).
\]

**Proposition 5.1.** Suppose \(\sigma = (\psi, \lambda)\) is an \(n\)-taxon rooted binary metric species tree. Then the clade probabilities \(\{P_{\sigma}(C)\}\) determine \(E_{\sigma}(D_T)\).

**Proof.** Define two indicator functions:

\[
I_C(T) = \begin{cases} 
1 & \text{if } T \text{ displays } C, \\
0 & \text{otherwise},
\end{cases}
\]

and

\[
I_{a,b}(C) = \begin{cases} 
1 & \text{if } a,b \in C, \\
0 & \text{otherwise}.
\end{cases}
\]

Then using equation (4),

\[
E_{\sigma}(D_T(a,b)) = \sum_{T} P_{\sigma}(T) 2(n - c_T(a,b))
\]

\[
= 2n - 2 \sum_{T} P_{\sigma}(T) c_T(a,b)
\]

\[
= 2n - 2 \sum_{T} P_{\sigma}(T) \left( \sum_{\text{non-trivial clades } C} I_C(T) I_{a,b}(C) \right)
\]

\[
= 2n - 2 \sum_{\text{non-trivial clades } C} \left( \sum_{T} P_{\sigma}(T) I_C(T) \right) I_{a,b}(C)
\]

\[
= 2n - 2 \sum_{\text{non-trivial clades } C} P_{\sigma}(C) I_{a,b}(C),
\]

and the expected STAR distance between \(a\) and \(b\) is computable from clade probabilities.

**Remark.** Though this statement and proof involve theoretical distributions and expected STAR distances, a similar statement and argument also apply to empirical clade frequencies and empirical STAR distances computed from a gene tree sample.

From Theorem 3.2 we immediately obtain the following:
Corollary 5.2. The rooted species tree topology $\psi$ is identifiable from clade probabilities under the multispecies coalescent.

Remark. Proposition 5.1 also suggests an efficient algorithm for computing $\psi$ from $\{P(C)\}$, or an estimate of $\psi$ from estimates of $\{P(C)\}$.

We note that the program BUCKy (Ané et al., 2007) uses estimated clade probabilities to construct a concordance tree from the clades with highest estimated probabilities one clade at a time using a greedy consensus approach. Motivated by the observation that this method can be misleading (Degnan et al., 2009), a quartet version of BUCKy was developed that builds the species tree from the most supported quartets (Larget et al., 2010). However, the observation that STAR distances can be computed from clade probabilities suggests an alternative method for computing the concordance tree using Equation (5) that would be consistent under the multispecies coalescent.

6. Discussion

The main results of this paper are that the STAR algorithm and its generalizations to other node numbering schemes, using any well-behaved distance method for the selection of the species tree, do provide statistically consistent methods of species tree inference. As demonstrated by Liu et al. (2009), the method is also fast and, at least in simulations, exhibited good performance in comparison to some other methods of species tree inference.

While our focus here has been theoretical, our work indicates several insights that may be helpful in improving practical data analysis. First, because the expected distance tables under STAR are ultrametric, it may be preferable to use UPGMA rather than Neighbor Joining as the tree building method in STAR, as UPGMA enforces an ultrametric assumption while NJ does not. Alternatively, if NJ is used, then even though the reconstructed distances on the inferred species tree are ultimately discarded, it might be worthwhile to first examine them to see if they roughly give ultrametricity. If not, then one might doubt either the fit of the multispecies coalescent model to the gene tree data, or that one had a sufficiently large sample of gene trees for adequate inference. One should also consider using a more elaborate distance method based on an optimality criterion, such as Minimum Evolution, although this would require a tree search and therefore should slow runtimes. Fast heuristic searches under such a criterion, possibly adapted to enforce ultrametricity, might improve STAR’s performance. Simulation studies are needed to further investigate these issues.

Second, the generalizations of the STAR method using alternate numbering schemes also need further investigations. Although all such schemes lead to consistent inference, this is an asymptotic statement that concerns behavior with large samples. It is still possible that some numbering schemes lead to more efficient inference, in that with a small sample of gene trees
they are more likely to return the correct species tree, or one close to it. Understanding this issue, perhaps for specific species trees, or better still for specific models of species tree generation, would be desirable.

Even without more detailed theoretical results on the behavior of the generalized STAR methods, they might still be useful in data analysis. For instance, for a fixed sample of gene trees, one could repeatedly infer a species tree using randomly chosen node numbering schemes satisfying the criteria in Section 3. If all such species trees have the same topology, one might be more confident in the result. Significant differences in the inferred topologies might lead one to doubt the validity of all the trees, again due to either poor model fit of the coalescent or inadequate sample size. More minor variations in the inferred trees might suggest which features of the species tree one might be confident of, based on their shared characteristics. Because of the computational speed of STAR, calculations with many random node numberings should be feasible.

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References

Allman, E. S., Degnan, J. H., and Rhodes, J. A. (2011a). Determining species tree topologies from clade probabilities under the coalescent. J. Theor. Biol., 289:96–106.

Allman, E. S., Degnan, J. H., and Rhodes, J. A. (2011b). Identifying the rooted species tree from the distribution of unrooted gene trees under the coalescent. J. Math. Biol., 62(6):833–862.

Ané, C., Larget, B., Baum, D., Smith, S., and Rokas, A. (2007). Bayesian estimation of concordance among gene trees. Mol. Biol. Evol., 24:412–426.

Atteson, K. (1997). The performance of the neighbor-joining method of phylogeny reconstruction. In Mathematical hierarchies and biology (Piscataway, NJ, 1996), volume 37 of DIMACS Ser. Discrete Math. Theoret. Comput. Sci., pages 133–147. Amer. Math. Soc., Providence, RI.

Castillo-Ramírez, S., Liu, L., Pearl, D., and Edwards, S. (2010). Bayesian estimation of species trees: A practical guide to optimal sampling and analysis. In Knowles, L. and Kubatko, L., editors, Estimating Species Trees: Practical and Theoretical Aspects, pages 15–33. Wiley-Blackwell, College Station, Texas.
Degnan, J. H., DeGiorgio, M., Bryant, D., and Rosenberg, N. A. (2009). Properties of consensus methods for inferring species trees from gene trees. *Syst. Biol.*, 58(1):35–54.

Heled, J. and Drummond, A. J. (2010). Bayesian inference of species trees from multilocus data. *Mol. Biol. Evol.*, 27:570–580.

Kubatko, L. S., Carstens, B. C., and Knowles, L. L. (2009). STEM: species tree estimation using maximum likelihood for gene trees under coalescence. *Bioinformatics*, 25(7):971–973.

Larget, B. R., Kotha, S. K., Dewey, C. N., and Ané, C. (2010). BUCKy: Gene tree / species tree reconciliation with Bayesian concordance analysis. *Bioinformatics*, 26:2910–2911.

Lee, J. Y., Joseph, L., and Edwards, S. V. (2012). A species tree for the Australo-Papuan fairy-wrens and allies (Aves: Maluridae). *Syst. Biol.*, 61:253–271.

Liu, L. and Pearl, D. K. (2007). Species trees from gene trees: Reconstructing Bayesian posterior distributions of a species phylogeny using estimated gene tree distributions. *Syst. Biol.*, 56:504–514.

Liu, L., Yu, L., Pearl, D. K., and Edwards, S. V. (2009). Estimating species phylogenies using coalescence times among sequences. *Syst. Biol.*, 58:468–477.

Rannala, B. and Yang, Z. (2003). Bayes estimation of species divergence times and ancestral population sizes using DNA sequences from multiple loci. *Genetics*, 164:1645–1656.

Semple, C. and Steel, M. (2003). *Phylogenetics*, volume 24 of *Oxford Lecture Series in Mathematics and its Applications*. Oxford University Press, Oxford.

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