Anchored distances for quartet-based estimation of phylogenetic trees and applications to coalescent-based analyses

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Abstract

Inferring trees based on pairwise distances has utility in phylogenetic reconstruction broadly and specifically for estimation of species trees from gene trees under the coalescent model. In this paper, we introduce a new general approach for computing distances between pairs of leaves by first computing the topology and the internal branch length of the quartet tree that includes the pair of leaves and two additional leaves, which we call "anchors". We show that using these anchored distances, a family of distance-based reconstruction methods can be designed, ranging in complexity between $\Theta(n^2)$ and $\Theta(n^4)$. We then describe how anchored distances can be used for statistically consistent estimation of species trees from discordant unrooted gene trees under the multi-species coalescent model. The resulting method, which we call DISTIQUE, is competitive with the best alternative methods. Supplementary material, code, and datasets are available at http://esayyari.github.io/DISTIQUE.

1 Introduction

Inferring trees using pairwise distances is a well-studied approach to phylogenetic reconstruction [1–4]. As maximum likelihood and Bayesian methods became faster [5–7], distance-based methods started to fall out of favor, but they still remain useful, for example, for quickly building guide trees for multiple sequence alignment [8–10]. Furthermore, many new distance-based methods have been introduced in recent years for coalescent-based analysis of gene trees [11–17].

The history of species and genes can be discordant [18], and Incomplete Lineage Sorting (ILS), modeled by the multi-species coalescent model [19], is one of the main causes of discordance. One approach for estimating the species phylogeny in the face of such discordances is to first estimate a gene tree for each gene, and to then summarize them to build a species tree. The summary method, thus, takes as input a set of gene trees and produces a species tree. One desirable property of a summary method is statistical consistency; i.e., it should be guaranteed to return the correct species tree with high probability as the number of (error-free) gene trees increases. Many statistically consistent summary methods have been developed (e.g., ASTRAL [14,20], MP-EST [15], BUCKy-population [16]), including some that are distance-based (e.g., NJst [11], and a new implementation of it called ASTRID [12], STEAC and STAR [13], and GLASS [17]). Coalescent-based species tree estimation is a vibrant field of research, with rapid advances and many recent examples of successful biological analyses [21–23] (but see [24–28] for criticism of these methods).

A powerful general approach to phylogenetic reconstruction is analyzing quartets, which are subsets of four leaves in a tree. Unrooted quartet trees can have one of only three possible topologies, making them easy to analyze. Thus, one can first infer a set of quartet trees and then combine them to build a tree on the full dataset [2,29,30]. Relatedly, one can use induced quartets to estimate

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a so-called supertree [31] from a collection of input trees [30, 32–35]. Quartet-based supertree estimation has been revived in recent years [16, 20, 36–38] because of its connections to coalescent-based analyses [39–41]. Under the coalescent model, for quartets of species, the unrooted species tree topology is always identical to the most likely unrooted gene tree (but this is not true for larger trees [40, 42]), and the length of the internal quartet species tree branch (in coalescent units) defines the probabilities of the gene tree quartet topologies. These observations are used in recent statistically consistent quartet-based species tree estimation methods. For example, ASTRAL seeks to find the species tree that shares the maximum number of quartet trees with input gene trees, an optimization problem that produces a statistically consistent estimator.

In this paper, we first introduce a method to estimate pairwise distances from quartet trees, using a general technique applicable to any measure of distance. Assume we can compute the topology and the length of the single internal branch ("quartet length" henceforth) for each quartet. We choose two arbitrary fixed "anchor" leaves, and define pairwise distance between all other leaves by computing the quartet tree topology and length for the quartet that consists of the two anchors and the two leaves of interest. We show that out measure of distance is additive, and hence, can be used for statistically consistent inference of trees (as long as the quartet topology/length estimator is consistent). These "double anchored" distances are defined for all leaves except the two anchors, but we introduce a family of methods that use double anchored distances to reconstruct trees on the complete leaf-set. These methods can be adjusted to have running times that range between $\Theta(n^2)$ and $\Theta(n^4)$ for n species (assuming other factors are constant).

A motivation of our anchored quartet-based distance estimation is to enable pairwise distance calculation when estimating quartet topology and length is straightforward but estimating leaf distances is not. Coalescent-based analyses have these properties. We introduce a family of statistically consistent coalescent-based summary methods, called DISTIQUE, which use anchoring to estimate species trees from gene trees. We evaluate the accuracy of DISTIQUE on simulated and biological data, and show that its accuracy is competitive with the best alternative methods.

2 Theoretical results

We start with general results, and then describe their applications to coalescent-based analyses.

2.1 Anchored distance matrices

Definitions: We denote the leaf-set by \mathcal{L} and let $n = |\mathcal{L}|$. We only consider unrooted trees, and for a tree T on \mathcal{L} , the set of quartet trees induced on all possible $\binom{n}{4}$ quartets of leaves is denoted by \mathcal{Q}^T . We use ab.cd to note that in the quartet tree on $\{a,b,c,d\}$, a and b are sisters (there are only three possibilities). A tree T is equivalent to a distance matrix, computed by summing lengths of the edges between pairs of leaves, and a distance matrix that corresponds to a tree is called additive. We refer to the unique tree associated with the additive matrix D as T^D and refer to the distance matrix computed on T as D^T ; we omit superscripts when clear.

To test for the additivity of a distance matrix D, we can use the four point condition [43], which states, for quartet of leaves $\{a,b,c,d\} \in \mathcal{L}$, the median and the maximum of the following three values should be the same: $\{D[a,b] + D[c,d], D[a,c] + D[b,d], D[a,d] + D[b,c]\}$. When internal branch lengths are assumed positive, as we do throughout this paper, the minimum value is strictly smaller than the median. Let w.l.o.g. D[a,b] + D[c,d] be the smallest value; the quartet topology ab.cd is induced by T^D (i.e., $ab.cd \in \mathcal{Q}^T$). If we denote the lengths of the single internal branch in the quartet tree (which we call its "quartet length") by $d_D(a,b,c,d)$, it is easy to show that $d_D(a,b,c,d) = \frac{1}{2}[D[a,c] + D[b,d] - (D[a,b] + D[c,d])]$.

Proper setting: (β, α, f) is called a proper setting if $\beta, \alpha > 0$ are two constants and f(x) is a monotonically increasing function, and $0 < f(x) < \beta$ for x > 0.

Anchored Distance Matrices: Given a tree T^D , a proper setting, and "anchor" leaves $u, v \in \mathcal{L}$,

$$D'_{uv}[a,b] = \begin{cases} \beta + \alpha . d_D(a,b,u,v) & ab.uv \notin \mathcal{Q}^T \\ \beta - f(d_D(a,b,u,v)) & ab.uv \in \mathcal{Q}^T \end{cases}$$
(1)

is a distance matrix on the leaf-set $\mathcal{L} - \{u, v\}$; we call D'_{uv} a "double anchored distance matrix", and say that D'_{uv} is induced from D anchored by u, v. Similarly, given a single anchor leaf $v \in \mathcal{L}$,

$$D'_{v}[a,b] = \sum_{u \in \mathcal{L} - \{a,b\}} D'_{uv}[a,b]$$
(2)

is a distance matrix on the leaf-set $\mathcal{L} - \{v\}$; we call D'_v a "single anchored distance matrix". And,

$$D'[a,b] = \sum_{v \in \mathcal{L} - \{a,b\}} D'_v[a,b] = \sum_{u,v \in \mathcal{L} - \{a,b\}} D'_{uv}[a,b]$$
(3)

is called an "all-pairs anchored distance matrix" and is defined on the complete leaf-set \mathcal{L} . Finally,

$$D''[a,b] = \max_{u,v \in \mathcal{L} - \{a,b\}} \max(0, \frac{D'_{uv}[a,b] - \beta}{\alpha})$$
(4)

is called an "all-pairs anchored maximum distance matrix", and is defined on the complete \mathcal{L} . Thus, D'[a,b] and D''[a,b] are the sum and max of double anchored distances for all $\binom{n-2}{2}$ ways of choosing two anchors other than (a,b).

Theorem 1 A double anchored distance matrix D'_{uv} induced from a distance matrix D^T and anchored by a pair of leaves $u, v \in \mathcal{L}$ is an additive distance matrix for the leaf-set $\mathcal{L}' = \mathcal{L} - \{u, v\}$ and corresponds to a tree that is topologically identical to $T|\mathcal{L}'$.

Theorem 2 A single anchored distance matrix D'_v induced from D^T anchored by $v \in \mathcal{L}$ is an additive distance matrix for the leaf-set $\mathcal{L}' = \mathcal{L} - \{v\}$ and corresponds to a tree that is topologically identical to $T|\mathcal{L}'$.

Theorem 3 An all-pairs anchored distance matrix D' induced from D^T is additive and correspond to a tree with identical topology to T.

Proofs of all three theorems are given in Appendix A, where we first show that the four point condition holds for an arbitrarily chosen quartet of leaves in all possible scenarios for adding u and v on a quartet tree. Proof of Theorem 2 is similar and involves adding one anchor to a quartet tree. Theorem 3 is proved by dividing Equation 3 to sums that resemble Theorems 1 and 2, and an extra case that can be easily analyzed.

Theorem 4 An All-pairs anchored maximum distance matrix D'' induced from D^T is additive and correspond to a tree with identical topology and internal branch lengths to T.

Proof. We prove that for leaves (a,b), Equation 4 returns the sum of internal branch lengths between a and b on D^T (which we call D^T_{ab}); the theorem automatically follows as D^T is additive and we are only interested in internal branch lengths. For simplicity, set $\alpha=1$, but the proof holds for other α values. If (a,b) are not sisters, there is at least an anchor pair (u,v) with quartet topology au.bv and $d_D(a,b,u,v)=D^T_{ab}$ (pick u from sister group of a after rooting T on b and vice-versa). With this choice, $D'_{uv}[a,b]-\beta=d_D(a,b,u,v)=D^T_{ab}$; moreover, $D'_{u'v'}[a,b]-\beta$ for other anchors are no bigger than D^T_{ab} (if they have ab.uv topology, $D'_{u'v'}[a,b]<\beta$, and otherwise, it will give the length for a subset of the path from a to b). Thus, max returns D^T_{ab} , as desired. When (a,b) are sisters, $D^T_{ab}=0$. For sister leaves, $D'_{uv}[a,b]<0$ for any (u,v), and thus, D''[a,b]=0, as desired.

Algorithm 1 Anchored quartet-based algorithms. $d^{\mathcal{D}}(.)$ is a quartet estimator and returns the quartet topology and length. $(\beta, \alpha, f(x))$ gives a proper setting. Stopping criteria should *at least* require that every leaf is in at least one tree in \mathcal{T} . DAD, SAD, AAD, and AMD give algorithms for double, single, all-pairs, and all-pairs maximum anchored distance-based reconstruction, respectively.

```
function SAD(\mathcal{D})
                                                                                                    function AAD(\mathcal{D})
function D'_{u,v}(a,b)
  (t,d) \leftarrow d^{\mathcal{D}}(a,b,u,v)
                                                        repeat for \{u\} \subset \mathcal{L}
                                                                                                      D' \leftarrow 0_{n \times n}
  if t = ab.uv then return \beta –
                                                          D'_u \leftarrow 0_{n-1 \times n-1}
                                                                                                      for \{a,b\}\subset\mathcal{L} do
                                                          for \{v\} \subset \mathcal{L} - \{u\} do
                                                                                                         for \{u,v\}\subset \mathcal{L}-\{a,b\} do
f(d)
                                                             for \{a,b\} \subset \mathcal{L} - \{u,v\}
                                                                                                            D'[a,b] + = D'_{u,v}(a,b)
  else return \beta + \alpha.d
                                                                                                      return NeighborJ(D')
                                                     do
function DAD(\mathcal{D})
                                                               D'[a,b] += D'_{u.v}(a,b)
                                                                                                    function AMD(D)
  repeat for \{u, v\} \subset \mathcal{L}
                                                                                                      D'' \leftarrow 0_{n \times n}
     D'_{uv} \leftarrow 0_{n-2 \times n-2}
                                                          add NeighborJ(D'_n) to \mathcal{T}
                                                                                                      for \{a,b\}\subset\mathcal{L} do
     for \{a,b\} \subset \mathcal{L} - \{u,v\} do
                                                        until stopping criteria
                                                                                                         for \{u,v\}\subset \mathcal{L}-\{a,b\} do
       D'[a,b] = D'_{u,v}(a,b)
                                                        return SuperFine(\mathcal{T})
                                                                                                            D''[a,b]
     add NeighborJ(D'_{uv}) to \mathcal{T}
                                                                                                    \max(D''[a,b], D'_{u,v}(a,b))
  until stopping criteria
                                                                                                      return NeighborJ(D'')
  return SuperFine(\mathcal{T})
```

2.2 Phylogenetic reconstruction using anchored quartet-based distances

Theorems 1-4 can be used to design a general family of tree reconstruction methods based on quartet distances. Let \mathcal{D} be the data from which we want to infer a tree; we do not make assumptions here about the nature of the data, but require a way to estimate quartet trees:

(Consistent) quartet estimator: A quartet estimator $d^{\mathcal{D}}(q)$ is a function that given a quartet of leaves $\{a, b, c, d\}$, uses \mathcal{D} to estimate the quartet tree topology and the quartet length. A quartet estimator is statistically consistent if, as size of \mathcal{D} increases, the estimated quartet tree topology and quartet length both converge in probability to correct values.

Statistically consistent quartet estimators can be designed for various types of data. For example, the four point condition gives a way to estimate quartet trees from sequence data [44], and the log-det method gives a model-based way of estimating corrected branch lengths [45]. As the next section shows, coalescent theory can also be used to design consistent quartet estimators.

Given a consistent quartet estimator $d^{\mathcal{D}}(.)$, Theorems 1-4 can be used to define a family of statistically consistent phylogenetic reconstruction methods that range in running time between $\Theta(n^2)$ and $\Theta(n^4)$ (fixing other parameters). Algorithm 1 shows general forms of these algorithms. All-pairs anchored distance-based (AAD) and all-pairs maximum distanced-based (AMD) are the simplest methods in this family, which use Equations 3 and 4 to compute the distance matrix (i.e., sum/max of double anchored distance over all possible $\binom{n-2}{2}$ anchors). They then compute the tree using a consistent distance-based method (here, neighbor joining [1]).

Theorem 5 All-pairs anchored distanced-based (AAD) and all-pairs maximum distance-based (AMD) phylogenetic reconstruction algorithms shown in Algorithm 1 are statistically consistent.

Proof (sketch). Since $d^{\mathcal{D}}(.)$ is assumed statically consistent, in limit, it will return the correct quartet topology with arbitrarily high probability, and its estimates of quartet lengths can be made arbitrarily close to true value with any desired probability. From Theorem 3 and 4, it follows that the distance matrices for AAD and AMD become arbitrarily close to additive, with high probability. The proof follows from statistical consistency of neighbor joining for distance matrices that are in limit arbitrarily close to additivity [1].

Using double or single anchored distance matrices would enable us to use only a subset of all $\binom{n-2}{2}$ anchors, but a difficulty is that these matrices do not include the complete leaf-set. To address

this difficulty, we compute a set of trees, each on n-1 or n-2 leaves, and then use a supertree method (i.e., SuperFine [46,47]) to combine them. General forms of algorithms for single anchored (SAD) or double anchored (DAD) distance-based tree inference are shown in Algorithms 1. A pair of anchors $\{u,v\}$ (or, a single anchor u for SAD) are selected with some criteria (e.g., random w/o replacement), a distance matrix is generated for $\mathcal{L} - \{u,v\}$ (or, $\mathcal{L} - \{u\}$ for SAD), and a distance method, neighbor joining [1], is used to estimate a tree on $\mathcal{L} - \{u,v\}$ (or, $\mathcal{L} - \{u\}$ for SAD). This process repeats until stopping criteria are met; the stopping criteria need to include that each leaf has been at least in one estimated tree. Finally, the collection of trees is combined using a supertree method (e.g., SuperFine) to get a single tree. If SuperFine is used, we can show:

Theorem 6 Single and double anchored distanced-based (SAD and DAD) phylogenetic reconstruction algorithms shown in Algorithm 1 are both statistically consistent.

Proof (sketch). Assume the stopping criteria are reached after k rounds. By arguments similar to those used for Theorem 5, we can argue that each individual tree T_i , $1 \le i \le k$ on n-1 (for SAD) or n-2 (for DAD) leaves is a statistically consistent estimate; thus, for any $\epsilon' < 1$ and i, there is a dataset size such that with probability $1 - \epsilon'$, T_i is recovered correctly. Take the largest of these dataset sizes; with this largest dataset size, with probability at least $1 - k\epsilon'$, every T_i is correct. Thus, setting $\epsilon' = \frac{\epsilon}{k}$ allows us to argue that for any ϵ , there is a dataset size where all T_i s are correct with probability higher than $1 - \epsilon$. Then, by Theorem 1 of [46], the strict consensus merger (and by extension SuperFine) applied to the set of T_i trees returns the correct tree on \mathcal{L} .

Branch lengths: None of our anchored methods can estimate terminal branch length (i.e., will set terminal branch lengths to zero). If a distance-based method has difficulty with handling zero length for terminal, a constant value can be added to all distances. Only AMD returns statistically consistent estimates of branch length; for other methods, the branch lengths should be ignored.

Running time analysis: AAD and AMD clearly require $\Theta(n^4)$ running time (assuming quartet estimator is constant time) to build the distance matrix; using the default neighbor joining algorithm, which requires $O(n^3)$, would result in $\Theta(n^4)$ asymptotic running time for AAD and AMD. The running times of SAD and DAD depend on the design of the stopping criteria, and also the exact distance method and SuperFine implementation used. To build the distance matrix, each iteration of DAD and SAD requires $\Theta(n^2)$ and $\Theta(n^3)$, respectively; if a fast neighbor joining algorithm is used (e.g., FNJ [48], or NINJA [4]), the running time of building each tree is $O(n^2)$. If the stopping criterion is limited by a constant, building the set of trees on n-2 or n-1 leaves using DAD and SAD become $\Theta(n^2)$ and $\Theta(n^3)$ respectively. However, the stopping criteria can limit the number of rounds to a function of n; for example, a stopping criteria that allows for up to \sqrt{n} rounds would result in $O(n^{2.5})$ and $O(n^{3.5})$ respectively. Clearly, any function between $\Theta(n^2)$ and $\Theta(n^4)$ can be achieved by adjusting the stopping criteria. Finally, DAD and SAD use SuperFine to combine the set of k trees. The running time of SuperFine depends on the supertree method used inside SuperFine; if used with a quadratic time algorithm, SuperFine becomes quadratic also, and thus, our DAD and SAD become $\Theta(n^2)$ and $\Theta(n^4)$ respectively.

2.3 DISTIQUE: anchored coalescent-based species tree estimation

Our anchored quartet-based methods enable calculation of pairwise distances, even when pairwise distances are not easy to estimate, as long as quartet distances can be computed. Species tree estimation under multi-species coalescent model [19] (which models ILS) is an example of such a scenario. Here, the input dataset \mathcal{D} is a set of gene trees, and we seek to find a species tree. We show how our anchored quartet-based distances can be used for statistically consistent species tree estimation. We assume in our theoretical analyses that the input gene trees are generated using the multi-species coalescent process, and that each gene includes an individual per species.

Coalescent-based quartet estimator: For a given quartet q, let $p_1 \leq p_2 \leq p_3$ denote frequencies of the three quartet topologies induced by gene trees; we define the coalescence-based quartet estimator $d^{\mathcal{D}}(q)$ to return the topology that that has frequency p_3 , with branch length $-\ln 3 - \ln p_1$.

Under the multi-species coalescent process, for four taxa, the species tree topology has higher probability than the two alternative frequencies [39]; moreover, for quartet length d in the species tree, the probability of the most probable quartet gene tree is $1 - \frac{2}{3}e^{-d}$, and the probability of the alternative topologies are both $\frac{1}{3}e^{-d}$ [39,49]. It immediately follows that:

Corollary 7 The coalescent-based quartet estimator is statistically consistent.

DISTIQUE: We define $\beta = \ln 3$, $\alpha = 1$, and $f(x) = \ln(3 - 2e^{-x})$. For x > 0, f(x) is clearly positive, monotonic, and bounded from above by $\beta = \ln 3$; thus, (β, α, f) is a proper setting. Moreover, it is easy to show that with these choices, the calculation of Equation 1 greatly simplifies to $D'_{uv}[a,b] = -\ln p(ab.uv)$. Thus, to define distances, we can simply find the frequency of all quartet topologies in gene trees and use methods shown in Algorithm 1 to reconstruct the species tree (replacing function $D'_{u,v}(a,b)$ with $-\ln p(ab.uv)$). We call the resulting family of methods DISTIQUE (Distance-based Inference of Species Trees from Induced QUartet Elements). Note that the all-pairs distance matrix simplifies to:

$$D'[a,b] = \sum_{u,v \in \mathcal{L} - \{a,b\}} -\ln p(ab.uv)$$

$$\tag{5}$$

Theorem 8 DISTIQUE methods are statistically consistent under multi-species coalescent model. Proof. By Corollary 7, our choice of quartet estimator is statistically consistent. Since $(\beta, \alpha, f(x))$ is a proper setting, Theorems 5 and 6 prove that DISTIQUE methods are statistical consistency.

3 Experimental evaluation

We use simulated and real datasets to evaluate the accuracy of four versions of DISTIQUE. We evaluate species tree accuracy, and measure it using False Negative (FN) rate, which is equivalent to normalized RF distance [50] here because all estimated species trees were fully resolved.

3.1 Methods

We compare four versions of DISTIQUE, described below, against each other, and against ASTRAL-II [14], which is a quartet-based method, the NJst algorithm [11], implemented in ASTRID [12], which is a distance-based method, and concatenation using RAxML [5]. We use ASTRAL and ASTRID because of their connections to DISTIQUE, and because they have been found to be more accurate than alternative summary methods [11,12,14,20,51,52].

DISTIQUE Variations In this paper, we evaluate only two variants of DISTIQUE: AAD and AMD (all-pairs sum and max). We chose the two all-pairs methods because they do not require a strategy for choosing anchors or stopping criteria, nor do they require the use of a supertree method. Thus, these are the simplest versions to test without extensive exploration of parameter choices. The two all-pairs DISTIQUE are the slowest variants, and future work needs to explore DISTIQUE family more broadly. Even these simple variants involved the following design choices.

Smoothing: The gene tree quartet probabilities are exponentially related to the species tree quartet length. Thus, for branches that are even moderately long, getting zero frequencies for some alternative quartet topologies becomes likely. For example, for a quartet species tree with length 8 (in coalescent units [49]), the probability of seeing no discordance among 1000 genes is 98%. Thus, using a simple empirical frequency estimator, we are likely to get zero probabilities, which

produce distances of infinity (Eq. 5). To avoid this problem, we use Krichevsky-Trofimov~(KT) [53] or add-half estimator, which means a pseudo-count of 0.5 is added for each quartet topology. This estimator was shown to reach the min-max cumulative loss for KL divergence asymptotically [53].

Consensus resolution: As Section 4.1 will demonstrate, the naive application of DISTIQUE, while as accurate as competing methods on model conditions with high ILS, did not give good accuracy for low ILS. This is despite the fact that lower ILS datasets are easies because of high similarity between gene treesand the species tree. When discordance is low, the true frequencies of many branches become difficult to estimate, a problem we partially tackle using smoothing. However, our solution is insufficient. When species are all separated by long branches, many quartet probabilities become exceedingly close to zero, and limited data have difficulty estimating long distances (this resembles traditional distance-based methods [2]). For example, species quartet lengths of 8, 16, or 32 are all likely to result in no gene tree discordance given 1000 genes, making it impossible to distinguish between these very different distances. We can construct examples when all gene trees are likely to be identical, ye AAD with smoothing is mislead (see Fig S8).

Long branches with low levels of discordance should be easy to recover, since they appear in most gene trees. A simple majority rule (50%) consensus of the input trees would return all these long branches. Thus, we can simply compute the majority rule consensus to recover the easy long branches, and then use DISTIQUE to resolve the polytomies in the consensus trees. To resolve a polytomy, we first assign a label to each branch pendant to it (defining a mapping from species to labels), and then build a tree using DISTIQUE with the labels as leaves; this tree defines a resolution of the polytomy. To use DISTIQUE with labels, we need to find empirical frequencies for quartets of labels. To do so, we select all quartets of species such that each species belongs to a different label (this will be a non-empty set), and set the frequency of the label quartet to the arithmetic mean of the frequencies of corresponding species quartets (we also tried geometric mean and root mean square, and observed no differences in results). We apply DISTIQUE to these label frequencies, which produce a tree on the label set and therefore a resolution of the polytomy.

Polytomies in input: DISTIQUE-AAD can seamlessly handle polytomies in input gene trees; unresolved nodes in gene trees result in unresolved quartet trees for that gene, which we exclude from our calculation of the empirical quartet frequencies.

DISTIQUE is implemented in python, using external libraries for manipulating trees (Dendropy [54]) and for recovering quartet trees from gene trees [55]. We used FastME [56] as our distance-based method, and used the same tool inside ASTRID (see Supplementary material for details); we also tried other distance-based method PhyD* [57] and observed no meaningful differences. The code is open source and publicly available at http://esayyari.github.io/DISTIQUE.

3.2 Datasets

We use three sets of simulated datasets from the literature in our studies: the mammalian dataset of [24], the avian dataset of [27,58], and the 11-taxon dataset of [51]. The first two datasets are based on biological data and have a single species tree topology, whereas the 11-taxon dataset is simulated using SimPhy [59] and has a different species tree per replicate. For the 11-taxon dataset, we have four levels of ILS, ranging from low (M1) to very high (M4), and for each case, we vary both the number of genes (100, 500, 1000) and the number of sites per gene (from 10 to 200). For the mammalian and avian datasets, we create two different collections, one where we fix the number of genes (to 200 for mammalian and to 1000 for avian) and vary the amount of ILS, and a second collection, where we fix the amount of ILS (to 0.2X for mammalian and 1X for avian) and vary the number of genes (100 to 3200 for mammalian and 200 to 2000 for avian). The amount of ILS is changed by multiplying or dividing branch lengths by 2 or 5; shorter branches (0.2X and

Table 1: Statistical significance of species tree topological error, comparing DISTIQUE-AAD-Cons versus ASTRAL and ASTRID, run on estimated gene trees, tested using ANOVA, with multiple test correction using Benjamini-Hochberg [44] (n=24; $\alpha=0.05$). The columns labelled *method* show p-values of the comparison between methods; those labelled m:ILS, m:sites, or m:genes show significance of the interaction between method and other parameters. \dagger : DISTIQUE is outperformed by ASTRAL or ASTRID; *: DISTIQUE outperforms ASTRAL or ASTRID.

	DISTIQUE versus ASTRAL				DISTIQUE versus ASTRID			
	method	m:ILS	m:sites	m:genes	method	m:ILS	m:sites	m:genes
11-taxon	0.9861	0.9861	0.9861	0.9861	$< 10^{-4*}$	$< 10^{-4}$	0.9861	$< 10^{-4}$
avian-1X	0.0856	_	_	0.7613	0.0001^{\dagger}	_	_	0.9929
avian-1000g	0.0511	$< 10^{-4}$	_	_	$< 10^{-4}$	0.8667	_	_
mammalian-0.2X	0.0856	_	_	0.6096	0.7613	_	_	0.8667
mammalian-200g	0.0856	0.0868	_	_	0.9861	0.0161	_	_

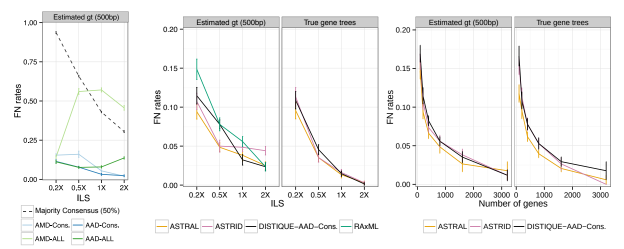


Figure 1: DISTIQUE variants on mammalian datasets with 200 genes.

Figure 2: DISTIQUE-AAD-Cons versus other methods on the mammalian dataset. (left) number of genes: 200; (right) ILS: 0.2X. Mean and standard error of species tree error is shown for true and estimated gene trees (500bp alignments).

0.5X) produce more ILS and longer branches reduce ILS See Table S1. For the biological analyses, we re-analyzed a dataset of 2022 supergene trees from an avian phylogenomic dataset [22, 27].

4 Results

4.1 Comparison between DISTIQUE variants

We compare DISTIQUE-AAD and DISTIQUE-AMD (sum and maximum of all-pairs), each applied to either the entire dataset or to polytomies of a 50% majority rule consensus (Cons). Figure 1 compares these four versions of DISTIUQE on the mammalian dataset as we vary the amount of ILS, and also shows the missing branch rate for the majority consensus tree. With very high ILS (0.2X), all methods have similar accuracy, and AAD is slightly better than AMD. As ILS decreases, a surprising pattern emerges. When DISTIQUE is applied to the entire dataset, the error unexpectedly goes up with decreased ILS, a pattern that is more pronounced for AMD. As discussed before, we attribute this pattern to difficulties of estimating long quartet lengths. When DISTIQUE is used to resolve polytomies in the consensus tree, the accuracy improves with decreased ILS, as expected. Note that even with reduced ILS, the consensus tree misses more than 25% of branches, and leaves some polytomies for DISTIQUE to resolve. Similar results were obtained on other datasets (Supplementary Figs. S1-S3). Since DISTIQUE-ADD-Cons was consistently our best method, we use this variant for comparisons to other methods.

4.2 DISTIQUE versus other methods

Mammalian: Figure 2 compares DISTIQUE against other summary methods as we change the level of ILS or the number of genes. As expected, reducing ILS and increasing the number of genes result in improved accuracy for all methods, and error is somewhat higher when estimated gene trees are used instead of true gene trees. With true gene trees, summary methods have very similar accuracy. When estimated gene trees are used, overall, DISTIQUE and ASTRID (the two distance-based methods) have similar accuracies (with no statistically significance differences according to an ANOVA test with FDR correction [44]; see Table 1). However, their relative accuracy depends on the level of ILS (the interaction effect is statistically significant - Table 1), where DISTIQUE is better with lower ILS and ASTRID with higher ILS. Comparing the two quartet-based methods, DISTIQUE and ASTRAL, it seems ASTRAL has a slight advantage but the differences are not statistically significant (p = 0.086). On the 500bp dataset with 200 genes, DISTIQUE was always at least as good as concatenation (concatenation was not performed for 0.2X ILS in [24]).

Avian: Figure 3 compares DISTIQUE against other methods on the avian datasets, when the amounts of ILS or the number of genes are varied. With true gene trees, the distance-based methods ASTRID and DISTIQUE slightly outperform ASTRAL, but differences tend to be small. On estimated gene trees, ASTRID and ASTRAL both seem to outperform DISTIQUE, but differences are statistically significant only for ASTRID (Table 1). The relative performance of DISTIQUE and ASTRAL is not significantly impacted by the number of genes, but the amount of ILS does have a statistically significant impact (Table 1); with more ILS, DISTIQUE has better accuracy and with lower ILS, ASTRAL is more accurate. The relative performance of DISTIQUE and ASTRID is impacted by neither the amount of ILS nor the number of genes. Interestingly, on this dataset, the species tree error based on estimated gene trees is much higher than the error with true gene trees (twice the error or more in most cases). Concatenation was the least accurate method in general, except on the dataset with reduced ILS (2X), where it was a bit more accurate than DISTIQUE.

11-taxon: Figure 4 shows results on the 11-taxon datasets, with varying amount of ILS, numbers of genes, and the numbers of sites per gene (which controls the amount of gene tree error). Overall, there are no statistically significant differences between DISTIQUE and ASTRAL, which have extremely close accuracies for most conditions. DISTIQUE is significantly more accurate than ASTRID on this dataset, and the magnitude of the difference depends on both the amount of ILS and number of genes, but not the number of sites (Table 1). Thus, differences become larger as the amount of ILS increases and the number of genes decreases. Finally, relative performance of concatenation (RAxML) and other methods depends on the amount of ILS and the number of sites. With low ILS, concatenation has much better accuracy than summary methods, including DISTIQUE, but as the amount of ILS increases, its error increases rapidly. Also, with few sits per gene, differences between concatenation and summary methods are small, but as the number of sites per gene increases and gene trees become more accurate, summary methods improve rapidly, while the accuracy of concatenation accuracy fails to improve as much.

4.3 Biological results

On the avian supergene trees, we ran ASTRAL, ASTRID, and DISTIQUE with multi-locus bootstrapping [60]; all three methods generated trees with relatively high bootstrap support (on average 95% for ASTRAL, 91% for NJst, and 97% for DISTIQUE), and were all different from the published tree generated using MP-EST [22] (trees shown in Figs. S6, S7). DISTIQUE and ASTRID differed on two branches, which both had low support in ASTRID. DISTIQUE and ASTRAL differed in five branches, and two of these in DISTIQUE and three of them in ASTRAL had high support. The presence of highly supported conflict between coalescent-based methods indicates a need for skeptical and careful analysis of results of any single method on real data.

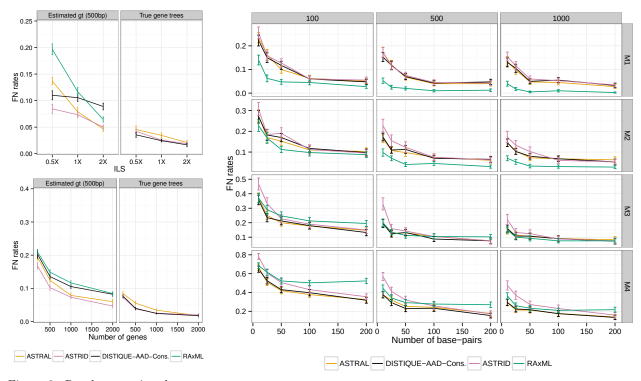


Figure 3: Results on avian datasets. (top) 1000 genes; (bottom) 1X ILS. Mean and standard error of species tree error is shown for true and estimated gene trees (20 replicates).

Figure 4: DISTIQUE versus other methods on the 11-taxon dataset. Columns show different number of genes, and rows show different levels of ILS. Mean and standard error of species tree error is shown for estimated gene trees with varying numbers of sites per gene (50 replicates).

5 Discussions and future work

We introduced a new distance-based approach for tree reconstruction by inferring topologies and internal branch lengths for quartets of leaves. Our new anchoring technique computes additive pairwise distances from quartet trees (potentially using a subset of all $\Theta(n^4)$ quartets of leaves), a technique that is useful especially when quartet trees are easy to calculate but pairwise distances are not. Coalescent-based analyses have this property. We used anchoring to design DISTIQUE, a new statistically consistent summary method for species tree estimation.

DISTIQUE was not consistently better than other summary methods in our experiments, but it was always close to the best method, and was never outperformed by more than 5% of branches. When we used true gene trees, DISTIQUE tended to match or outperform other summary methods, but it was slightly more sensitive to gene tree estimation error than ASTRAL and ASTRID were. It is possible that these differences reflect the fact that DISTIQUE, unlike ASTRAL or ASTRID (i.e., NJst), uses the expected quartet probabilities according to the coalescence model, and as a results is more sensitive to errors in gene trees, which distort these probabilities. Using methods beyond simple empirical frequency [61] might further improve the accuracy of DISTIQUE.

Despite having strong competition in ASTRAL and ASTRID (and possibly other methods we did not test), we believe there are reasons to further develop DISTIQUE in various ways. The accuracies of double or single anchored versions of DISTIQUE with $O(n^2)$ to $O(n^4)$ need to be tested to see if they can achieve similar levels of accuracy as the current version. Moreover, the accuracy of DISTIQUE may improve using other techniques such as estimating quartet lengths using parametric methods, or weighting various quartets according to coalescent expectations. Finally, we may be able to use DISTIQUE to co-estimate gene trees and the species tree using distances.

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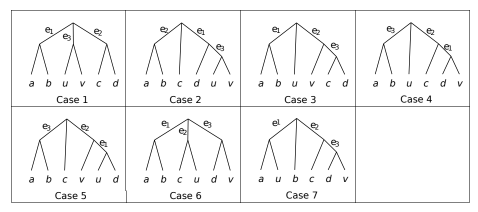
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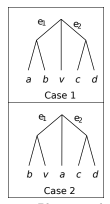


Figure 5: Placements of two anchors u and v on a given quartet topology ab.cd.

Figure 6: Placing anchor v on quartet tree ab.cd.

Appendices

A Theorem proofs

Proof of Theorem 1: We show that for arbitrary $\{a, b, c, d\} \subset \mathcal{L}$, the four point conditions holds for D'_{uv} and that the conditions are compatible with T. Thus, assuming $ab.cd \in \mathcal{Q}^T$, we show

$$D'_{uv}[a,b] + D'_{uv}[c,d] < D'_{uv}[a,c] + D'_{uv}[b,d] = D'_{uv}[a,d] + D'_{uv}[b,c]$$

$$(6)$$

Figure 5 shows all ways of placing anchors (u and v) on the quartet tree ab.cd. Anchors u and v can be sisters, in which case they can be placed on the internal branch (Case 1) or on one of the tip branches (Case 2; w.l.o.g, we use the branch leading to d). When u and v are not sisters, they can be both placed on the internal branch (Case 3), or one on the internal branch and the other on a tip branch (Case 4), or they can be both on terminal branches, which can be done in three ways: u and v can be on the same terminal branch (Case 5), on different but adjacent branches (Case 6), or on two non-adjacent branches (Case 7).

In Table 2, for each of the seven cases, we compute all parts of Equation 6; $LHS = D'_{uv}[a,b] + D'_{uv}[c,d]$, $RHS1 = D'_{uv}[a,d] + D'_{uv}[b,c]$, and $RHS2 = D'_{uv}[a,c] + D'_{uv}[b,d]$. Computation of values in Table 2 is straight-forward; each $D'_{uv}[x,y]$ should be compute using Equation 1; thus, where xy.uv is induced by the tree shown in Figure 5, we use $[\beta - f(d)]$ and otherwise we use $[\beta + \alpha d]$, where $d = d_D(x,y,u,v)$ is the quartet length (length of the internal branch) for the quartet tree induced by $\{x,y,u,v\}$. For example, for Case 1, $D'_{uv}[a,b]$ is $[\beta - f(e_1+e_3)]$ because ab.uv is induced by the tree, and the distance of the edge on the ab.uv quartet is $e_1 + e_3$; in Case 7, we set $D'_{uv}[a,b]$ to $\beta + \alpha e_1$ because ab.uv is not induced by the tree and $d_D(a,b,u,v) = e_1$.

Table 2: Four point condition for all 7 cases shown in Figure 5. For each case, LHS < RHS1 and RHS1 = RHS2.

	LHD	RHS1	RHS2
	$D'_{uv}[a,b] + D'_{uv}[c,d]$	$D'_{uv}[a,d] + D'_{uv}[b,c]$	$D'_{uv}[a,c] + D'_{uv}[b,d]$
Case 1	$[\beta - f(e_1 + e_3)] + [\beta - f(e_2 + e_3)]$	$[\beta - f(e_3)] + [\beta - f(e_3)]$	$[\beta - f(e_3)] + [\beta - f(e_3)]$
Case 2	$[\beta - f(e_1 + e_2 + e_3)] + [\beta - f(e_3)]$	$[\beta - f(e_3)] + [\beta - f(e_1 + e_3)]$	$[\beta - f(e_1 + e_3)] + [\beta - f(e_3)]$
	$[\beta - f(e_1)] + [\beta - f(e_3)]$	$[\beta + \alpha e_2] + [\beta + \alpha e_2]$	$[\beta + \alpha e_2] + [\beta + \alpha e_2]$
Case 4	$[\beta - f(e_3)] + [\beta + \alpha e_1]$	$[\beta + \alpha(e_1 + e_2)] + [\beta + \alpha e_2]$	$[\beta + \alpha e_2] + [\beta + \alpha(e_1 + e_2)]$
Case 5	$[\beta - f(e_2 + e_3)] + [\beta + \alpha e_1]$	$[\beta - f(e_2)] + [\beta + \alpha e_1]$	$[\beta - f(e_2)] + [\beta + \alpha e_1]$
Case 6	$[\beta - f(e_1)] + [\beta + \alpha(e_2 + e_3)]$	$[\beta + \alpha e_3] + [\beta + \alpha e_2]$	$[\beta + \alpha e_2] + [\beta + \alpha e_3]$
Case 7	$[\beta + \alpha e_1][\beta + \alpha e_3]$	$[\beta + \alpha(e_1 + e_2 + e_3)] + [\beta + \alpha e_2]$	$[\beta + \alpha(e_1 + e_2)] + [\beta + \alpha(e_2 + e_3)]$

We need to show that LHS < RHS1 and that RHS1 = RHS2. We remind the reader that all branches are assumed to be strictly positive, and that f is a positive and monotonically increasing function bounded from above by β . In all cases, the equality of RHS1 and RHS2 is immediately clear from Table 2. The inequality (LHS < RHS1) follows directly from the fact that f(x) is monotonically increasing in Cases 1, 2, and 5. For Case 3, because of positivity of branch lengths and f(x), we have $LHS < 2\beta < RHS$. Similarly, for Case 4, $LHS < 2\beta + \alpha e_1 < 2\beta + \alpha e_1 + 2\alpha e_2 = RHS$. Case 6 follows from the positivity of f, and Case 7 is trivially correct for positive branch lengths. \square

Proof of Theorem 2: For distances defined using Equation 2, we show that the four point condition holds for an arbitrary $\{a, b, c, d\} \subset \mathcal{L}$, assuming w.l.o.g that $ab.cd \in \mathcal{Q}^T$:

$$\sum_{u \notin \{a,b\}} D'_{uv}[a,b] + \sum_{u \notin \{c,d\}} D'_{uv}[c,d] < \sum_{u \notin \{a,d\}} D'_{uv}[a,d] + \sum_{u \notin \{a,b\}} D'_{uv}[b,c] = \sum_{u \notin \{a,c\}} D'_{uv}[a,c] + \sum_{u \notin \{b,d\}} D'_{uv}[b,d] + \sum_{u \notin \{a,b\}} D'_{uv}[a,b] + \sum_{u \notin \{a,b\}}$$

Let $\mathcal{L}' = \mathcal{L} - \{a, b, c, d\}$. Each sum can be broken into cases where $u \in \{a, b, c, d\}$ or $u \in \mathcal{L}'$:

$$\begin{split} &D'_{cv}[a,b] + D'_{dv}[a,b] + D'_{av}[c,d] + D'_{bv}[c,d] + \sum_{u \in \mathcal{L}'} D'_{uv}[a,b] + D'_{uv}[c,d] < \\ &D'_{bv}[a,d] + D'_{cv}[a,d] + D'_{av}[b,c] + D'_{dv}[b,c] + \sum_{u \in \mathcal{L}'} D'_{uv}[a,d] + D'_{uv}[b,c] = \\ &D'_{bv}[a,c] + D'_{dv}[a,c] + D'_{av}[b,d] + D'_{cv}[b,d] + \sum_{u \in \mathcal{L}'} D'_{uv}[a,c] + D'_{uv}[b,d] \end{split}$$

For terms where $u \in \mathcal{L}'$, the additivity is proved in Theorem 1. Thus, we need to prove inequalities and qualities of terms where $u \in \{a, b, c, d\}$. The single anchor v can be placed either on the internal branch (Case 1) or on a terminal branches (Case 2), as Figure 6 shows. In Case 1, for each term of the sum, we have:

 $LHS = [\beta - f(e_1)] + [\beta - f(e_1)] + [\beta - f(e_2)] + [\beta - f(e_2)] < 4\beta < [\beta + \alpha e_1] + [\beta + \alpha e_2] + [\beta + \alpha e_1] + [\beta + \alpha e_2] = RHS$ and for the second case we have:

$$LHS = [\beta + \alpha e_1] + [\beta + \alpha e_1] + [\beta - f(e_2)] + [\beta - f(e_1 + e_2)] < 4\beta + 2\alpha e_1 - f(e_1 + e_2) < [\beta + \alpha e_2] + [\beta - f(e_1)] + [\beta + \alpha e_1] + [\beta + \alpha (e_1 + e_2)] = RHS1 = RHS2$$

Since addition of additive matrices are additive, the proof follows.

Proof of Theorem 3: Similar to proof of Theorem 2, terms in Equation 3 can be of three types: $\{u, v\}$ and $\{a, b, c, d\}$ intersect in either (I) two elements, (II) one element, or (III) none. Thus, the four point condition can be written out as $(\mathcal{L}' = \mathcal{L} - \{a, b, c, d\})$:

$$\begin{split} &2D'_{ab}[c,d] + \sum_{v \in \mathcal{L'}} D'_{cv}[a,b] + D'_{dv}[a,b] + D'_{av}[c,d] + D'_{bv}[c,d] + \sum_{u,v \in \mathcal{L'}} D'_{uv}[a,b] + D'_{uv}[c,d] < \\ &2D'_{ad}[b,c] + \sum_{v \in \mathcal{L'}} D'_{cv}[a,d] + D'_{bv}[a,d] + D'_{av}[b,c] + D'_{dv}[b,c] + \sum_{u,v \in \mathcal{L'}} D'_{uv}[a,d] + D'_{uv}[b,c] = \\ &2D'_{ac}[b,d] + \sum_{v \in \mathcal{L'}} D'_{bv}[a,c] + D'_{dv}[a,c] + D'_{av}[b,d] + D'_{cv}[b,d] + \sum_{u,v \in \mathcal{L'}} D'_{uv}[a,c] + D'_{uv}[b,d] \end{split}$$

For terms of type (III) and (II), the additivity is proved in Theorems 1 and 2, respectively. Thus, we need to prove additivity only for terms of type (I), which have no anchors; we have:

$$LHS = 2D'_{ab}[c,d] = 2[\beta - f(d)] < 2\beta < 2[\beta + \alpha d] = 2D'_{ad}[b,c] = 2D'_{ac}[b,d] = RHS; \ d = d_D(a,b,c,d)$$

Thus, for all three types, inequalities and qualities needed to prove additivity hold. \Box