

Reconciliation Revisited: Handling Multiple Optima when Reconciling with Duplication, Transfer, and Loss

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ABSTRACT

Phylogenetic tree reconciliation is a powerful approach for inferring evolutionary events like gene duplication, horizontal gene transfer, and gene loss, which are fundamental to our understanding of molecular evolution. While duplication–loss (DL) reconciliation leads to a unique maximum-parsimony solution, duplication-transfer-loss (DTL) reconciliation yields a multitude of optimal solutions, making it difficult to infer the true evolutionary history of the gene family. This problem is further exacerbated by the fact that different event cost assignments yield different sets of optimal reconciliations. Here, we present an effective, efficient, and scalable method for dealing with these fundamental problems in DTL reconciliation. Our approach works by sampling the space of optimal reconciliations uniformly at random and aggregating the results. We show that even gene trees with only a few dozen genes often have millions of optimal reconciliations and present an algorithm to efficiently sample the space of optimal reconciliations uniformly at random in $O(mn^2)$ time per sample, where m and n denote the number of genes and species, respectively. We use these samples to understand how different optimal reconciliations vary in their node mappings and event assignments and to investigate the impact of varying event costs. We apply our method to a biological dataset of approximately 4700 gene trees from 100 taxa and observe that 93% of event assignments and 73% of mappings remain consistent across different multiple optima. Our analysis represents the first systematic investigation of the space of optimal DTL reconciliations and has many important implications for the study of gene family evolution.

Key words: gene duplication, gene family evolution, gene-tree/species-tree reconciliation, horizontal gene transfer, host-parasite cophylogeny, phylogenetics.

1. INTRODUCTION

THE SYSTEMATIC COMPARISON OF A GENE TREE with its species tree under a reconciliation framework is a powerful technique for understanding gene family evolution. Specifically, gene tree/species tree reconciliation shows how the gene tree evolved inside the species tree while accounting for events like gene

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duplication, gene loss, and horizontal gene transfer, which drive gene family evolution. Thus, gene tree/species tree reconciliation is widely used and has many important applications, for example, for inferring orthologs, paralogs, and xenologs (Storm and Sonnhammer, 2002; Koonin, 2005; Wapinski et al., 2007; van der Heijden et al., 2007; Vilella et al., 2009; Sennblad and Lagergren, 2009; Mi et al., 2010); reconstructing ancestral gene content and dating gene birth (Chen et al., 2000; Ma et al., 2008; David and Alm, 2011), accurate gene tree reconstruction (Vilella et al., 2009; Rasmussen and Kellis, 2011), and whole genome species-tree reconstruction (Bansal et al., 2007; Burleigh et al., 2011).

Duplication–loss (DL) reconciliation, which accounts for only gene duplication and gene loss events, has been widely studied and extensively used (Goodman et al., 1979; Page, 1994; Mirkin et al., 1995; Eulenstein and Vingron, 1998; Bonizzoni et al., 2005; Durand et al., 2006; Górecki and Tiuryn, 2006; Chauve et al., 2008). However, since it does not account for horizontal gene transfer events, it only applies to multicellular eukaryotes, a very small part of the tree of life. An interesting and extremely useful property of DL-reconciliation is that, assuming loss events have a nonzero positive cost, the most parsimonious reconciliation is always unique (Górecki and Tiuryn, 2006). In addition, the most parsimonious reconciliation remains the same irrespective of the chosen event costs for duplication and loss. Given these properties, there is no ambiguity in interpreting the results of DL-reconciliation, which makes it extremely easy to use in practice.

The limited applicability of DL reconciliation has led to the formulation of the duplication-transfer-loss (DTL) reconciliation model, which can simultaneously account for duplication, transfer, and loss events and can be applied to species and gene families from across the entire tree of life. Indeed, the DTL-reconciliation model and its variants have been widely studied in the literature (Gorbunov and Liubetskii, 2009; Doyon et al., 2010; Tofigh, 2009; Tofigh et al., 2011; David and Alm, 2011; Chen et al., 2012; Bansal et al., 2012; Stolzer et al., 2012). In addition, DTL-reconciliation has also been indirectly studied in the context of the host–parasite cophylogeny problem (Charleston, 1998; Ronquist, 2003; Merkle and Middendorf, 2005; Libeskind-Hadas and Charleston, 2009; Merkle et al., 2010; Conow et al., 2010; Ovadia et al., 2011).

The DTL-reconciliation problem is typically solved in a parsimony framework, where costs are assigned to duplication, transfer, and loss events, and the goal is to find a reconciliation with minimum total cost. DTL-reconciliations can sometimes be *time-inconsistent*; that is the inferred transfers may induce contradictory constraints on the dates for the internal nodes of the species tree. The problem of finding an optimal *time-consistent* reconciliation is known to be NP-hard (Tofigh et al., 2011; Ovadia et al., 2011). Thus, in practice, the goal is to find an optimal (but not necessarily time-consistent) DTL-reconciliation. The problem of finding an optimal time-consistent reconciliation does become efficiently solvable (Libeskind-Hadas and Charleston, 2009; Doyon et al., 2010) if the species tree is fully dated. However, accurately dating the internal nodes of a species tree is a notoriously difficult problem (Rutschmann, 2006), which severely restricts its applicability. Thus, for wider applicability and efficient solvability, in this work, unless otherwise stated, we assume the input species tree is undated and seek an optimal (not necessarily time-consistent) DTL-reconciliation (Tofigh et al., 2011; David and Alm, 2011; Chen et al., 2012; Bansal et al., 2012). This problem can be solved very efficiently, with our own algorithm achieving the fastest known time complexity of $O(mn)$ (Bansal et al., 2012), where m and n denote the number of nodes in the gene tree and species tree respectively.

Despite its extensive literature, the DTL-reconciliation problem remains difficult to use in practice for understanding gene family evolution. The first reason for this difficulty is that there are often multiple equally optimal reconciliations for a given gene tree and species tree and for a fixed assignment of event costs. The second reason is that event costs, which can be very difficult to assign confidently, play a much more important role than in DL reconciliation, as varying the costs can result in different optimal reconciliations.

Thus, when applying DTL-reconciliation in practice, it is unclear whether the evolutionary history implied by a particular given optimal solution is meaningful, as many other optimal reconciliations exist with the same minimal reconciliation cost. Moreover, it is unclear whether the properties of an optimal reconciliation are representative of the space of optimal reconciliations, and also how large and diverse this space is. Furthermore, the number of optimal reconciliations is often prohibitively large, as it can grow exponentially in the number of events required for the reconciliation, making even the basic task of enumerating all optimal reconciliations unfeasible for all but the smallest of gene trees (Chen et al., 2012). Here, we directly address these problems and seek to make DTL-reconciliation as easy to use as the DL-reconciliation model.

It was recently shown that, when the species tree is fully dated, one can represent the set of all optimal reconciliations in a compact way by building a polynomially-sized minimum reconciliation graph

(Scornavacca et al., 2013). While this is an interesting approach to dealing with multiple optima, the fact that this is only known to work when the species tree is fully dated severely limits its applicability in practice. Moreover, since constructing the minimum reconciliation graph requires $O(mn^3)$ time and space (Scornavacca et al., 2013), where m and n denote the number of nodes in the gene tree and species tree, respectively, it can only be applied to small instances of the problem.

Our contribution. In this work, we develop an efficient and scalable approach to explore the space of optimal DTL-reconciliations and show how it can be used to infer the similarities and differences in the different optimal reconciliations for any given input instance. Our approach is based on uniformly random sampling of optimal reconciliations, and we demonstrate the utility of our approach by applying it to a biological dataset of approximately 4700 gene trees from 100 (predominantly prokaryotic) taxa (David and Alm, 2011). Specifically, our contributions are as follows:

1. We analyze the gene trees in the biological dataset and show that even gene trees with only a few dozen genes often have many millions of optimal reconciliations. This analysis provides the first detailed look into the prevalence of optimal reconciliations in biological datasets.
2. We study some basic structural properties of optimal DTL-reconciliations, which shed light on the inherent structure in optimal reconciliations and are both mathematically and biologically interesting.
3. We show how to efficiently sample the space of optimal reconciliations uniformly at random. Our algorithm produces each random sample in $O(mn^2)$ time and requires only $O(mn)$ space, where m and n denote the number of nodes in the gene tree and species tree, respectively. This algorithm is fast enough to be applied thousands of times to the same dataset and scalable enough to be applied to datasets with hundreds or thousands of taxa.
4. We use our algorithm for random sampling to explore the space of optimal reconciliations and investigate the similarities and differences between the different optimal reconciliations. We show how to distinguish between the parts of the reconciliation that have high support from those that are more variable across the different multiple optima.
5. We show that even in the presence of multiple optimal solutions, a large amount of shared information can be extracted from the different optimal reconciliations. For instance, we observed that, for fixed event costs, any internal node taken from a gene tree in the biological dataset had a 93.31% chance of having the same event assignment (speciation, duplication, or transfer) and a 73.15% chance of being mapped to the same species tree node, across all (sampled) optimal reconciliations.
6. Our method allows users to compare the space of optimal reconciliations for different event costs and extract the shared aspects of the reconciliation. This makes it possible to study the impact of using different event costs and to meaningfully apply DTL-reconciliation even if one is unsure of the exact event costs to use. We applied our method to the biological dataset using different event costs and observed that large parts of the reconciliation tend to be robust to event cost changes.

Thus, our new method allows for large-scale, systematic exploration of the space of optimal reconciliations in real biological datasets and makes it possible to deal effectively with multiple optima by being able to distinguish between the parts of the reconciliation that have high support and those that are more variable across the different optimal reconciliations.

The remainder of the article is organized as follows: The next section introduces basic definitions and preliminaries. In Section 3, we study the prevalence of multiple optimal reconciliations in biological data, and in Section 4, the basic structural properties of optimal reconciliations. We introduce our sampling-based approach and algorithms in Section 5. The results of our analysis of the multiple optimal reconciliations for the biological dataset appear in Section 6, and in Section 7, we show how our method can be applied to study the impact of using different reconciliation costs. Concluding remarks appear in Section 8.

2. DEFINITIONS AND PRELIMINARIES

We follow the basic definitions and notation from Bansal et al. (2012). Given a tree T , we denote its node, edge, and leaf sets by $V(T)$, $E(T)$, and $Le(T)$ respectively. If T is rooted, the root node of T is denoted by $rt(T)$, the parent of a node $v \in V(T)$ by $pa_T(v)$, its set of children by $Ch_T(v)$, and the (maximal) subtree of T rooted at v by $T(v)$. If two nodes in T have the same parent, they are called *siblings*. The set of *internal nodes* of T , denoted $I(T)$, is defined to be $V(T) \setminus Le(T)$. We define \leq_T to be the partial order on $V(T)$ where

$x \leq_T y$ if y is a node on the path between $rt(T)$ and x . The partial order \geq_T is defined analogously, that is, $x \geq_T y$ if x is a node on the path between $rt(T)$ and y . We say that v is an *ancestor* of u , or that u is a *descendant* of v , if $u \leq_T v$ (note that, under this definition, every node is a descendant as well as an ancestor of itself). We say that x and y are *incomparable* if neither $u \leq_T v$ nor $v \leq_T u$. Given a nonempty subset $L \subseteq Le(T)$, we denote by $lca_T(L)$ the least common ancestor (LCA) of all the leaves in L in tree T ; that is, $lca_T(L)$ is the unique smallest upper bound of L under \leq_T . Given $x, y \in V(T)$, $x \rightarrow_T y$ denotes the unique path from x to y in T . We denote by $d_T(x, y)$ the number of edges on the path $x \rightarrow_T y$. Throughout this work, unless otherwise stated, the term “tree” refers to a rooted binary tree.

We assume that each leaf of the gene tree is labeled with the species from which that gene was sampled. This labeling defines a *leaf-mapping* $\mathcal{L}_{G,S}: Le(G) \rightarrow Le(S)$ that maps a leaf node $g \in Le(G)$ to that unique leaf node $s \in Le(S)$, which has the same label as g . Note that gene trees may have more than one gene sampled from the same species. Throughout this work, we denote the gene tree and species tree under consideration by G and S respectively and will implicitly assume that $\mathcal{L}_{G,S}(g)$ is well defined.

2.1. Reconciliation and DTL-scenarios

Reconciling a gene tree with a species tree involves mapping the gene tree into the species tree. Next, we define what constitutes a valid reconciliation; specifically, we define a duplication-transfer-loss scenario (DTL-scenario) (Tofigh et al., 2011; Bansal et al., 2012) for G and S that characterizes the mappings of G into S that constitute a biologically valid reconciliation. Essentially, DTL-scenarios map each gene tree node to a unique species tree node in a consistent way that respects the immediate temporal constraints implied by the species tree and designates each gene tree node as representing either a speciation, duplication, or transfer event.

Definition 2.1 (DTL-scenario). *A DTL-scenario for G and S is a seven-tuple $\langle \mathcal{L}, \mathcal{M}, \Sigma, \Delta, \Theta, \Xi, \tau \rangle$, where $\mathcal{L}: Le(G) \rightarrow Le(S)$ represents the leaf-mapping from G to S , $\mathcal{M}: V(G) \rightarrow V(S)$ maps each node of G to a node of S , the sets Σ , Δ , and Θ partition $I(G)$ into speciation, duplication, and transfer nodes respectively; Ξ is a subset of gene tree edges that represent transfer edges, and $\tau: \Theta \rightarrow V(S)$ specifies the recipient species for each transfer event, subject to the following constraints:*

1. If $g \in Le(G)$, then $\mathcal{M}(g) = \mathcal{L}(g)$.
2. If $g \in I(G)$ and g' and g'' denote the children of g , then,
 - (a) $\mathcal{M}(g) \not\leq_S \mathcal{M}(g')$ and $\mathcal{M}(g) \not\leq_S \mathcal{M}(g'')$,
 - (b) At least one of $\mathcal{M}(g')$ and $\mathcal{M}(g'')$ is a descendant of $\mathcal{M}(g)$.
3. Given any edge $(g, g') \in E(G)$, $(g, g') \in \Xi$ if and only if $\mathcal{M}(g)$ and $\mathcal{M}(g')$ are incomparable.
4. If $g \in I(G)$ and g' and g'' denote the children of g , then,
 - (a) $g \in \Sigma$ only if $\mathcal{M}(g) = lca(\mathcal{M}(g'), \mathcal{M}(g''))$ and $\mathcal{M}(g')$ and $\mathcal{M}(g'')$ are incomparable,
 - (b) $g \in \Delta$ only if $\mathcal{M}(g) \geq_S lca(\mathcal{M}(g'), \mathcal{M}(g''))$,
 - (c) $g \in \Theta$ if and only if either $(g, g') \in \Xi$ or $(g, g'') \in \Xi$.
 - (d) If $g \in \Theta$ and $(g, g') \in \Xi$, then $\mathcal{M}(g)$ and $\tau(g)$ must be incomparable, and $\mathcal{M}(g')$ must be a descendant of $\tau(g)$, that is, $\mathcal{M}(g') \leq_S \tau(g)$.

Constraint 1 above ensures that the mapping \mathcal{M} is consistent with the leaf-mapping \mathcal{L} . Constraint 2(a) imposes on \mathcal{M} the temporal constraints implied by S . Constraint 2(b) implies that any internal node in G may represent at most one transfer event. Constraint 3 determines the edges of G that are transfer edges. Constraints 4(a), 4(b), and 4(c) state the conditions under which an internal node of G may represent a speciation, duplication, and transfer respectively. Constraint 4(d) specifies which species may be designated as the recipient species for any given transfer event.

In some cases, one may wish to restrict transfer events to only occur between coexisting species. This requires that divergence time information (either absolute or relative) be available for all the internal nodes of the species tree. In such cases, the definition of a DTL-scenario remains the same, except for the additional restriction on transfer events.

DTL-scenarios correspond naturally to reconciliations, and it is straightforward to infer the reconciliation of G and S implied by any DTL-scenario. Figure 1 shows two simple DTL-scenarios. Given a DTL-scenario, one can directly count the minimum number of gene losses (Bansal et al., 2012) in the corresponding reconciliation as follows.

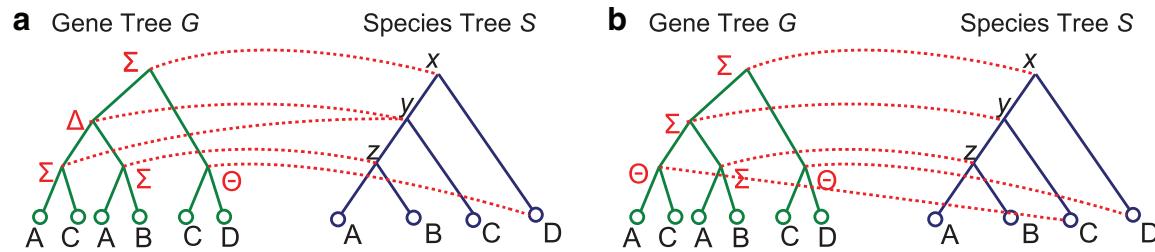


FIG. 1. Multiple optimal reconciliations. Parts (a) and (b) show two different reconciliations for the gene tree and species tree depicted in the figure. Both of the reconciliations are optimal for event costs $P_{\Delta} = 1$, $P_{\Theta} = 3$, and $P_{loss} = 1$. The reconciliation in part (a) invokes one duplication, one transfer, and two losses, while the reconciliation in part (b) invokes two transfers.

Definition 2.2 (Losses). Given a DTL-scenario $\alpha = \langle \mathcal{L}, \mathcal{M}, \Sigma, \Delta, \Theta, \Xi, \tau \rangle$ for G and S , let $g \in V(G)$ and $\{g', g''\} = Ch(g)$. The number of losses $Loss_{\alpha}(g)$ at node g is defined to be:

- $|d_S(\mathcal{M}(g), \mathcal{M}(g')) - 1| + |d_S(\mathcal{M}(g), \mathcal{M}(g'')) - 1|$, if $g \in \Sigma$
- $d_S(\mathcal{M}(g), \mathcal{M}(g'))$, if $g \in \Delta$ and $\mathcal{M}(g) = \mathcal{M}(g')$.
- $d_S(\mathcal{M}(g), \mathcal{M}(g')) + d_S(\mathcal{M}(g), \mathcal{M}(g''))$, if $g \in \Delta$, $\mathcal{M}(g) \neq \mathcal{M}(g')$, and $\mathcal{M}(g) \neq \mathcal{M}(g'')$, and
- $d_S(\mathcal{M}(g), \mathcal{M}(g')) + d_S(\tau(g), \mathcal{M}(g'))$ if $(g, g') \in \Xi$.

We define the total number of losses in the reconciliation corresponding to the DTL-scenario α to be $Loss_{\alpha} = \sum_{g \in I(G)} Loss_{\alpha}(g)$.

Let P_{Δ} , P_{Θ} , and P_{loss} denote the costs associated with duplication, transfer, and loss events respectively. The cost of reconciling G and S according to a DTL-scenario α is defined as follows.

Definition 2.3 (reconciliation cost of a DTL-scenario). Given a DTL-scenario $\alpha = \langle \mathcal{L}, \mathcal{M}, \Sigma, \Delta, \Theta, \Xi, \tau \rangle$ for G and S , the reconciliation cost associated with α is given by $\mathcal{R}_{\alpha} = P_{\Delta} \cdot |\Delta| + P_{\Theta} \cdot |\Theta| + P_{loss} \cdot Loss_{\alpha}$.

Given G and S , along with event costs P_{Δ} , P_{Θ} , and P_{loss} , the goal is to find a most parsimonious reconciliation of G and S . More formally,

Problem 1 (most parsimonious reconciliation, or MPR). Given G and S , the most parsimonious reconciliation (MPR) problem is to find a DTL-scenario for G and S with minimum reconciliation cost.

We distinguish two versions of the MPR problem: (i) the *undated MPR (U-MPR)* problem in which the species tree is undated, and (ii) the *fully-dated MPR (D-MPR)* problem in which every node of the species tree has an associated divergence time estimate (or there is a known total order on the internal nodes of the species tree), and transfer events are required to occur only between coexisting species.

Note that even if G and S are such that S contains one or more species that are not represented in G , we keep the species tree as is (i.e., we do not trim S to match the species set of G).

3. MULTIPLE OPTIMAL SOLUTIONS

In general, for any fixed values of P_{Δ} , P_{Θ} , and P_{loss} , there may be multiple equally optimal solutions to the MPR problem (both U-MPR and D-MPR). This is illustrated in Figure 1. The figure also illustrates the fundamental problem with having multiple optima: Given the different evolutionary histories implied by the different multiple optima, what is the true evolutionary history of the gene family? We address this problem in this article. But first, in this section, we investigate the prevalence of optimal reconciliations in real datasets. For our study, we use a published biological dataset of 4735 gene trees and 100 (predominantly prokaryotic) species (David and Alm, 2011). The gene trees in the dataset have median and average

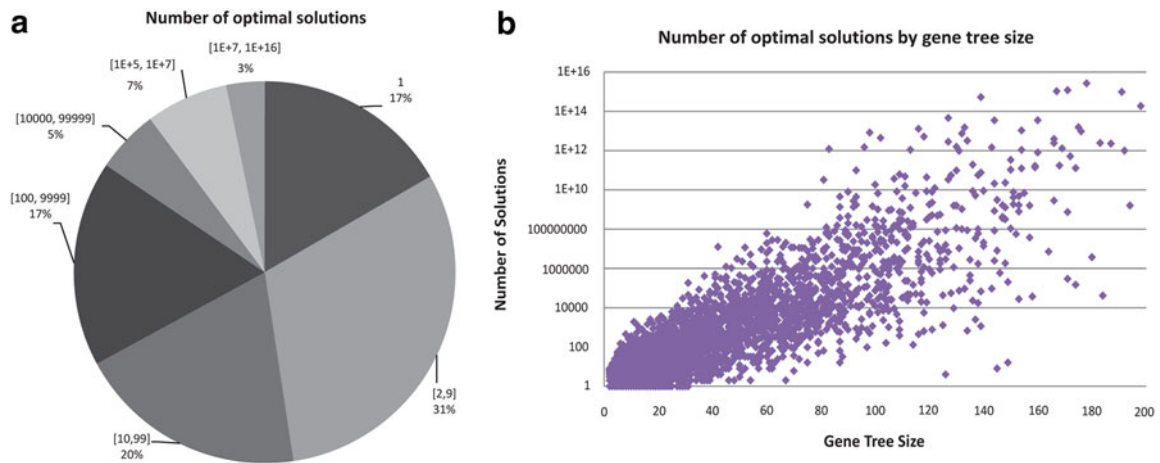


FIG. 2. Number of optimal reconciliations for gene trees in the biological dataset. The pie chart in part (a) shows the distribution of the number of optimal reconciliations for the different gene trees in the biological dataset. The dot plot in part (b) plots the size (in the number of internal nodes) and the number of optimal reconciliations for each gene tree. Due to arithmetic overflow concerns, results are only shown for the 4699 (out of 4735) gene trees that had fewer than 10^{16} optima.

leaf-set sizes of 18 and 35.1, respectively. This dataset has been previously analyzed using DTL-reconciliation but without consideration of multiple optima. In our analysis of this dataset we used the same event costs as used by David and Alm (2011) (i.e., $P_{\Delta} = 2$, $P_{\Theta} = 3$, and $P_{loss} = 1$). Since the gene trees in the dataset are unrooted, we first rooted them optimally by choosing a root that minimized the reconciliation cost. In cases where there were multiple optimal rootings, we chose one of the optimal rootings at random. We computed the number of multiple optimal reconciliations for each of the rooted gene trees by augmenting the dynamic programming algorithm used to solve the MPR problem (e.g., Bansal et al. 2012) to keep track of the number of optima for each subproblem. Further algorithmic details appear in Section 5. Unless otherwise stated, all analyses in the manuscript were performed using the undated version of DTL-reconciliation (i.e., the U-MPR problem).

Figure 2 shows the results of our analysis. As part (a) of the figure shows, only 17% of the approximately 4700 gene trees have a unique optimal reconciliation. Over half of the gene trees have over 100 optimal reconciliations and 15% have more than 10,000 optimal reconciliations. This illustrates the extent of the problem with multiple optimal reconciliations in biological datasets. As part (b) of the figure shows, the number of optimal reconciliations tends to increase exponentially with gene tree size. These results demonstrate the importance of considering multiple optima in DTL-reconciliation, and the impracticality of enumerating all optimal reconciliations for all but the smallest gene trees.

We also repeated the above analysis using the dated version of the DTL-reconciliation problem (i.e., the D-MPR problem), and observed no significant reduction in the number of multiple optima. For instance, even for the dated version, 14% of the gene trees had more than 10,000 optimal reconciliations. To make sure that the choice of event costs was not responsible for inflating the counts (since, in this case, $P_{\Theta} = P_{\Delta} + P_{loss}$, potentially making it easier to trade off transfers for duplications and losses), we also tried changing the event costs to 1.9, 2.9, and 0.9 for duplication, transfer, and loss, respectively, and observed only a small reduction in the numbers of inferred optimal reconciliations. Specifically, we observed that 11.6% of the gene trees still had over 10,000 optimal reconciliations, and 7.6% still had over 100,000 optimal reconciliations.

Recall that the gene trees in the dataset were originally unrooted. While the results above are for a fixed optimal rooting of these gene trees, we point out that about half the gene trees in the dataset have more than one optimal rooting. It may thus be necessary, in practice, to either consider all possible optimal rootings when studying multiple optimal reconciliations, or to use other information to assign a root uniquely.

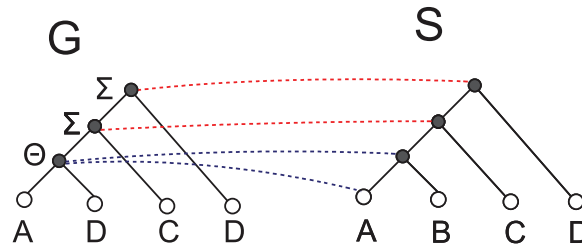


FIG. 3. Mappings from transfer events. Assuming that transfers, duplications, and losses have an equal cost, any optimal reconciliation of G and S must have a cost of 2. The figure depicts two such optimal reconciliations, one in which the transfer node maps to species A and one in which it maps to the species represented by $lca(A, B)$. These two mappings are shown by the dashed blue lines. All other nodes have identical mappings in the two reconciliations. Both reconciliations have one loss and one transfer, and the reconciliation in which the transfer node maps to $lca(A, B)$ shows that transfer node mappings are not constrained in the manner of speciation or duplication nodes as shown in Theorem 4.1.

4. BASIC STRUCTURAL PROPERTIES OF OPTIMAL RECONCILIATIONS

While there may be multiple optimal solutions for any given instance of the MPR problem, the gene tree–species tree mappings in optimal reconciliations are, as we show, strongly constrained. The following theorem applies to both the U-MPR and D-MPR problems.

Theorem 4.1. *Let g be any node in $I(G)$ and g', g'' be its two children. If loss events have a nonzero positive cost, then any optimal solution for the MPR problem on G and S must satisfy the following constraint: If $g \in \Sigma \cup \Delta$ then $\mathcal{M}(g) = lca(\mathcal{M}(g'), \mathcal{M}(g''))$.*

Proof. Let α denote any optimal reconciliation for G and S . Suppose, for the sake of contradiction, that α does not satisfy the constraint of part (1). Then, under the mapping of α , G must contain a node $h \in \Sigma_\alpha \cup \Delta_\alpha$ such that $\mathcal{M}_\alpha(h) >_S lca(\mathcal{M}_\alpha(h'), \mathcal{M}_\alpha(h''))$, where $\{h', h''\} = Ch(h)$. Let α' denote an alternative DTL-scenario obtained from α by changing the mapping of h to $lca(\mathcal{M}_\alpha(h'), \mathcal{M}_\alpha(h''))$. Clearly, α' is a valid DTL-scenario. We will show that α' has a lower reconciliation cost than α , a contradiction.

If $h \in \Sigma_\alpha$, then the constraint follows immediately from the definition of DTL-scenarios. Thus, assume that $h \in \Delta_\alpha$. Note that the number of transfers, duplications, and speciations is completely identical for α and α' . Consequently, any difference in the reconciliation costs of α and α' must be purely due to different numbers of losses. Also observe that, if $g \notin \{h \cup pa(h)\}$ then $Loss_\alpha(g) = Loss_{\alpha'}(g)$. Let $x = \mathcal{M}_\alpha(h)$ and $y = \mathcal{M}_{\alpha'}(h)$. Since $h \in \Delta_\alpha$, it follows from the definition of losses that $Loss_{\alpha'}(h) = Loss_\alpha(h) - 2 \times d(x, y)$. It also follows that $Loss_{\alpha'}(pa(h)) \leq Loss_\alpha(pa(h)) + d(x, y)$, irrespective of whether $pa(h)$ is a speciation, duplication, or transfer. Since $d(x, y) \geq 1$, it follows that $Loss_{\alpha'} < Loss_\alpha$, which implies that α' has a lower reconciliation cost than α . This contradicts the initial assumption that α is an optimal reconciliation. ■

Similar observations were made by David and Alm (2011), who used the constraints on the mappings to speed up their algorithm for the MPR problem. However, they did not provide any proofs and erroneously assumed that the constraint on the mapping of speciation and duplication events also applies to transfer events; that is, if $g \in \Theta$ and $(g, g') \in \Xi$ then $\mathcal{M}(g) = \mathcal{M}(g')$. As Figure 3 demonstrates, this does not hold true for transfer events.

5. UNIFORMLY RANDOM SAMPLING OF OPTIMAL RECONCILIATIONS

As Section 3 demonstrates, the exhaustive enumeration of all optimal reconciliation is only feasible for very small gene trees. Indeed, as Figure 2 shows, even gene trees with only a few dozen taxa often have hundreds of millions of solutions, and this number grows exponentially with gene tree size. In this section, we show how to sample the space of reconciliations uniformly at random. Random sampling makes it possible to explore the space of optimal reconciliations without exhaustive enumeration and

makes it possible to understand the variability in the different reconciliations and to distinguish between the highly supported and weakly supported parts of a given optimal reconciliation. Our algorithm for random sampling is based on the dynamic programming algorithm for the MPR problem from Bansal et al. (2012). The idea is to keep track of the number of optimal solutions for each subproblem considered in the dynamic programming algorithm. In the following, we show how to compute the number of optimal solutions at each step correctly and efficiently. First, we need a few definitions.

Given any $g \in I(G)$ and $s \in V(S)$, let $c_{\Sigma}(g, s)$ denote the cost of an optimal reconciliation of $G(g)$ with S such that g maps to s and $g \in \Sigma$. The terms $c_{\Delta}(g, s)$ and $c_{\Theta}(g, s)$ are defined similarly for $g \in \Delta$ and $g \in \Theta$ respectively. Given any $g \in V(G)$ and $s \in V(S)$, we define $c(g, s)$ to be the cost of an optimal reconciliation of $G(g)$ with S such that g maps to s . The algorithm for the MPR problem performs a nested post-order traversal of the gene tree and species tree to compute the value of $c(g, s)$ for each g and s . The dynamic programming table is initialized as follows for each $g \in Le(G)$: $c(g, s) = 0$ if $s = \mathcal{M}(g)$, and $c(g, s) = \infty$ otherwise. For $g \in I(G)$, observe that $c(g, s) = \min\{c_{\Sigma}(g, s), c_{\Delta}(g, s), c_{\Theta}(g, s)\}$.

At each step, the values of $c_{\Sigma}(g, s)$, $c_{\Delta}(g, s)$, and $c_{\Theta}(g, s)$ for any $g \in I(G)$ and $s \in V(S)$, can be computed based on the previously computed values of $c(\cdot, \cdot)$. To show how $c_{\Sigma}(g, s)$, $c_{\Delta}(g, s)$, and $c_{\Theta}(g, s)$ are computed, we need some additional notation. Let $in(g, s) = \min_{x \in V(S(s))} \{P_{loss} \cdot d_S(s, x) + c(g, x)\}$ and $out(g, s) = \min_{x \in V(S)} \text{incomparable to } s \{c(g, x)\}$. In other words: $out(g, s)$ is the cost of an optimal reconciliation of $G(g)$ with S such that g may map to any node from $V(S)$ that is incomparable to s ; and $in(g, s)$ is the cost of an optimal reconciliation of $G(g)$ with S such that g may map to any node, say x , in $V(S(s))$ but with an additional reconciliation cost of one loss event for each edge on the path from s to x . The values $c_{\Sigma}(g, s)$, $c_{\Delta}(g, s)$, and $c_{\Theta}(g, s)$ are computed as follows:

For any $g \in I(G)$ and $s \in I(S)$, let $\{g', g''\} = Ch_G(g)$ and $\{s', s''\} = Ch_S(s)$.

If $s \in Le(S)$ then,

$$c_{\Sigma}(g, s) = \infty,$$

$$c_{\Delta}(g, s) = P_{\Delta} + c(g', s) + c(g'', s), \text{ and}$$

$$\text{if } s \neq rt(S), \text{ then } c_{\Theta}(g, s) = P_{\Theta} + \min\{in(g', s) + out(g'', s), in(g'', s) + out(g', s)\}. \text{ Else, } c_{\Theta}(g, s) = \infty.$$

If $s \in I(S)$, then

$$c_{\Sigma}(g, s) = \min\{in(g', s') + in(g'', s''), in(g'', s') + in(g', s'')\}.$$

$$c_{\Delta}(g, s) = P_{\Delta} + \min \begin{cases} c(g', s) + in(g'', s'') + P_{loss}, \\ c(g', s) + in(g'', s') + P_{loss}, \\ c(g'', s) + in(g', s'') + P_{loss}, \\ c(g'', s) + in(g', s') + P_{loss}, \\ c(g', s) + c(g'', s), \\ in(g', s') + in(g'', s'') + 2P_{loss}, \\ in(g', s'') + in(g'', s') + 2P_{loss}, \\ in(g', s') + in(g'', s') + 2P_{loss}, \\ in(g', s'') + in(g'', s'') + 2P_{loss}. \end{cases}$$

If $s \neq rt(S)$, then $c_{\Theta}(g, s) = P_{\Theta} + \min\{in(g', s) + out(g'', s), in(g'', s) + out(g', s)\}$. Else, $c_{\Theta}(g, s) = \infty$.

The optimal reconciliation cost of G and S is simply: $\min_{s \in V(S)} c(rt(G), s)$, and an optimal reconciliation with that cost can be reconstructed by backtracking in the dynamic programming table. We refer the reader to Bansal et al. (2012) for further algorithmic details.

5.1. Computing the number of optimal reconciliations

To output optimal reconciliations uniformly at random we must keep track of the number of optimal reconciliations for each of the subproblems considered in the dynamic programming algorithm. We define the following: For any $g \in V(G)$ and $s \in V(S)$, let $N(g, s)$ denote the number of optimal solutions for reconciling $G(g)$ with S such that g maps to s . The idea is to compute $N(\cdot, \cdot)$ using the same nested post-order traversal used to compute the $c(\cdot, \cdot)$ values. The dynamic programming table for $N(\cdot, \cdot)$ is initialized as follows for each $g \in Le(G)$:

$$N(g, s) = \begin{cases} 1 & \text{if } g \in Le(G) \text{ and } s = \mathcal{L}(g), \\ 0 & \text{if } g \in Le(G) \text{ and } s \neq \mathcal{L}(g). \end{cases}$$

To compute $N(g, s)$, for $g \in I(G)$, we must consider all possible mappings of g' and g'' that yield a cost of $c(g, s)$. In the interest of brevity and clarity, let us assume that $s \in I(S)$ and $s \neq rt(S)$; the cases when $s \in Le(S)$ or $s = rt(S)$ are easy to handle analogously.

Let a_1 through a_{13} denote the individual expressions in the $\min\{ \}$ blocks in the equations for $c_{\Sigma}(g, s)$, $c_{\Delta}(g, s)$, and $c_{\Theta}(g, s)$ above. Specifically, let a_1 denote $in(g', s') + in(g'', s'')$, a_2 denote $in(g'', s') + in(g', s'')$, a_3 through a_{11} denote the nine expressions in the $\min\{ \}$ block for $c_{\Delta}(g, s)$, and a_{12} and a_{13} denote the two expressions in the $\min\{ \}$ block for $c_{\Theta}(g, s)$. Each of these a_i s represents a certain cost, which we denote by $c(a_i)$, and a certain number of optimal reconciliations, which we denote by $N(a_i)$. Furthermore, let b_i , for $1 \leq i \leq 13$, be binary boolean variables associated with the a_i s such that $b_i = 1$ if a_i yields the minimum cost $c(g, s)$, and $b_i = 0$ otherwise. Specifically, for $i \in \{1, 2\}$, $b_i = 1$ if and only if $c(a_i) = c(g, s)$; for $i \in \{3, \dots, 11\}$, $b_i = 1$ if and only if $c(a_i) + P_{\Delta} = c(g, s)$; and for $i \in \{12, 13\}$, $b_i = 1$ if and only if $c(a_i) + P_{\Theta} = c(g, s)$. Then, we must have:

$$N(g, s) = \sum_{i=1}^{13} b_i \times N(a_i). \quad (1)$$

Next, we show how to compute $N(a_i)$ for any i . Observe that each a_i has one term involving g' and one term involving g'' . These terms take one of the three forms: $c(\cdot, \cdot)$, $in(\cdot, \cdot)$, or $out(\cdot, \cdot)$. These terms, involving g' and g'' , can be viewed as representing the choice of optimal mappings for g' and g'' , respectively. For instance, $c(g', s)$ implies that g' must map to s , $in(g', s)$ implies that g' may map to any node $x \in V(S(s))$ for which $(P_{loss} \cdot d_S(s, x) + c(g', x))$ is minimized (recall the definition of $in(\cdot, \cdot)$), and $out(g', s)$ implies that g' may map to any node $x \in V(S)$ that is incomparable to s , for which $c(g', x)$ is minimized. Based on this observation, for any given a_i , we can compute a set of optimal mappings for g' , which we will denote by X' and a set of optimal mappings for g'' , which we will denote by X'' . It is not hard to see that the value of $N(a_i)$ must be as follows:

$$N(a_i) = \left(\sum_{x \in X'} N(g', x) \right) \times \left(\sum_{x \in X''} N(g'', x) \right). \quad (2)$$

Note that, for deriving the expression of $N(g, s)$ above, we have assumed that $s \in I(S) \setminus rt(S)$; this is because the expression for at least one of $c_{\Sigma}(g, s)$, $c_{\Delta}(g, s)$, or $c_{\Theta}(g, s)$ changes when $s \in Le(S)$ or $s = rt(S)$. The expressions for $N(g, s)$, for $s \in Le(S)$, and $s = rt(S)$ can be derived similarly by simply accounting for these minor differences in the number of terms that constitute the sum on the R.H.S. of Equation 1. Specifically, when $s \in Le(S)$, the terms $c_{\Sigma}(g, s)$, $c_{\Delta}(g, s)$, and $c_{\Theta}(g, s)$ contribute 0, 1, and 2 a_i s, respectively. Thus, in this case, there are only three a_i s: a_1 denotes $c(g', s) + c(g'', s)$, a_2 denotes $in(g', s) + out(g'', s)$, and a_3 denotes $in(g'', s) + out(g', s)$.

Similarly, when $s = rt(S)$, the terms $c_{\Sigma}(g, s)$ and $c_{\Delta}(g, s)$ contribute, as before, 2 and 9 a_i s, respectively, while $c_{\Theta}(g, s)$ contributes none; this yields eleven a_i s, a_1, \dots, a_{11} , that are identical to the first eleven a_i s used in Equation 1. Thus, the analogues of Equation 1 for the two cases when $s \in Le(S)$ and $s = rt(S)$ are, respectively:

$$N(g, s) = \sum_{i=1}^3 b_i \times N(a_i), \quad \text{and} \quad (3)$$

$$N(g, s) = \sum_{i=1}^{11} b_i \times N(a_i). \quad (4)$$

The equations for $N(g, s)$ and $N(a_i)$ derived above make it possible to compute the value $N(g, s)$ for each $g \in I(G)$ and $s \in V(S)$ by using the same nested post-order traversal that is used for computing the values $c(\cdot, \cdot)$. For completeness, a detailed description of the algorithm is given below as Procedure *ComputeNumSolutions*.

Procedure *ComputeNumSolutions*($G, S, \mathcal{L}, P_\Sigma, P_\Delta, P_\Theta$)

```

1: for each  $g \in V(G)$  and  $s \in V(S)$  do
2:   Initialize  $c(g, s)$ ,  $c_\Sigma(g, s)$ ,  $c_\Delta(g, s)$ , and  $c_\Theta(g, s)$  to  $\infty$ , and  $N(g, s)$  to 0.
3: for each  $g \in Le(G)$  do
4:   Initialize  $c(g, \mathcal{L}(g))$  to 0, and  $N(g, \mathcal{L}(g))$  to 1.
5: for each  $g \in I(G)$  in post-order do
6:   for each  $s \in V(S)$  in post-order do
7:     Let  $\{g', g''\} = Ch_G(g)$ .
8:     if  $s \in Le(S)$  then
9:        $c_\Sigma(g, s) = \infty$ .
10:       $c_\Delta(g, s) = P_\Delta + c(g', s) + c(g'', s)$ .
11:       $c_\Theta(g, s) = P_\Theta + \min\{in(g', s) + out(g'', s), in(g'', s) + out(g', s)\}$ .
12:       $c(g, s) = \min\{c_\Sigma(g, s), c_\Delta(g, s), c_\Theta(g, s)\}$ .
13:      Assign the three  $a_i$  expressions as described earlier (for the case when  $s \in Le(S)$ ) and, for each  $i \in \{1, 2, 3\}$ ,
      compute  $N(a_i)$  using Equation 2.
14:      Compute  $N(g, s)$  using Equation 3.
15:     else
16:       Let  $\{s', s''\} = Ch_S(s)$ .
17:        $c_\Sigma(g, s) = \min\{in(g', s') + in(g'', s''), in(g'', s') + in(g', s'')\}$ .
18:        $c_\Delta(g, s) = P_\Delta + \min \begin{cases} c(g', s) + in(g'', s'') + P_{loss}, \\ c(g', s) + in(g'', s') + P_{loss}, \\ c(g'', s) + in(g', s'') + P_{loss}, \\ c(g'', s) + in(g', s') + P_{loss}, \\ c(g', s) + c(g'', s), \\ in(g', s') + in(g'', s'') + 2P_{loss}, \\ in(g', s'') + in(g'', s') + 2P_{loss}, \\ in(g', s') + in(g'', s') + 2P_{loss}, \\ in(g', s'') + in(g'', s'') + 2P_{loss}. \end{cases}$ 
19:       if  $s \neq rt(S)$ , then  $c_\Theta(g, s) = P_\Theta + \min\{in(g', s) + out(g'', s), in(g'', s) + out(g', s)\}$ .
20:        $c(g, s) = \min\{c_\Sigma(g, s), c_\Delta(g, s), c_\Theta(g, s)\}$ .
21:       if  $s \neq rt(S)$  then
22:         Assign the thirteen  $a_i$  expressions as described earlier (for the case when  $s \in I(S) \setminus rt(S)$ ) and compute
          $N(a_i)$ , for each  $i \in \{1, \dots, 13\}$ , using Equation 2.
23:         Compute  $N(g, s)$  using Equation 1.
24:       if  $s = rt(S)$  then
25:         Assign the eleven  $a_i$  expressions as described earlier (for the case when  $s = rt(S)$ ) and compute  $N(a_i)$ , for
         each  $i \in \{1, \dots, 11\}$ , using Equation 2.
26:       Compute  $N(g, s)$  using Equation 4.
    
```

Let m and n denote the number of leaf nodes in G and S , respectively. We have the following:

Theorem 5.1. *Given G and S , and fixed event costs P_Δ , P_Θ , and P_{loss} , the value $N(g, s)$ for each $g \in V(G)$ and $s \in V(S)$ can be computed in $O(mn^2)$ time.*

Proof. From Bansal et al. (2012) we already know that Procedure *ComputeNumSolutions* correctly computes each $c(\cdot, \cdot)$. We will show that Procedure *ComputeNumSolutions* correctly computes each $N(\cdot, \cdot)$ in $O(mn^2)$ time. ■

Correctness: Observe that the value of $N(g, s)$ is initialized correctly for each $g \in Le(G)$ in the “for” loops of Steps 1 and 3. Let g be any node in $I(G)$ and let $\{g', g''\} = Ch_G(g)$. Let us assume that the values $N(g', x)$ and $N(g'', x)$ have been computed correctly for each $x \in V(S)$. We will show that the value of $N(g, s)$, for any $s \in V(S)$, is computed correctly as well.

For some fixed g and s , consider any a_i for which $b_i = 1$ in the equation used to compute $N(g, s)$ (i.e., in Eqs 1, 3, or 4). Let X' and X'' be as defined for Equation 2, for this chosen a_i . By definition,

any mapping from g' to a node in x' and from g'' to a node in X'' yields an optimal solution with cost $c(g, s)$ for the subproblem associated with the value $N(g, s)$. Thus, the mapping of g' represents exactly $\sum_{x \in X'} N(g', x)$ optimal reconciliations for the subproblem, while the mapping of g'' represents exactly $\sum_{x \in X''} N(g'', x)$. Since the mappings of g' and g'' can be assigned independently (for any fixed a_i), the total number of optimal reconciliations for the subproblem must be exactly as given by Equation 2.

To complete the proof we must now also show the correctness of Equation 1 (the correctness of Eqs. 3 and 4 will follow trivially). It suffices to show that the terms chosen as part of the sum in the R.H.S. of Equation 1 (i.e., for which $b_i = 1$) represent pairwise disjoint sets of reconciliations. Observe that the expressions for a_3 through a_7 and for a_{10} through a_{13} each represent a set of reconciliations that does not overlap with any other a_i (due to different constraints on the mappings for at least one of g' or g''). However, the expression for a_1 matches the expression for a_8 , both of which restrict the mapping of g' as $in(g', s')$ and the mapping of g'' as $in(g'', s'')$, and the expression for a_2 matches the expression for a_9 , both of which restrict the mapping of g' as $in(g', s'')$ and the mapping of g'' as $in(g'', s')$. However, since the terms a_1 and a_2 represent speciation while the terms a_8 and a_9 represent duplication, the lowest reconciliation costs for scenarios a_1 and a_2 will always be less than the lowest reconciliation costs for scenarios a_8 and a_9 (assuming that duplication and/or loss events have non zero positive costs). This implies that neither b_8 nor b_9 can ever be 1. Thus, the terms chosen as part of the sum in the R.H.S. of Equation 1 (i.e., for which $b_i = 1$) must all represent pairwise disjoint sets of reconciliations.

Induction completes the proof.

Complexity: We analyze the complexity of Procedure *ComputeNumSolutions* step-by-step. The “for” loops from Steps 1 through 4 require $O(mn)$ time. Steps 7 through 26 each require at most $O(n)$ time and are each executed $O(mn)$ times (through the “for” loops at lines 5 and 6), yielding a total time complexity of $O(mn^2)$ for these steps. The total time complexity of Procedure *ComputeNumSolutions* is thus $O(mn^2)$. ■

Corollary 5.1. *Given G and S , and fixed event costs P_Δ , P_Θ , and P_{loss} , the total number of optimal reconciliations of G and S can be computed in $O(mn^2)$ time.*

Proof. Let μ denote the minimum reconciliation cost of G and S , and let Y denote the set $\{s \in V(S) : c(rt(G), s) = \mu\}$. Then, the total number of optimal reconciliations of G and S is simply $\sum_{s \in Y} N(rt(G), s)$. Since all $c(\cdot, \cdot)$ and all $N(\cdot, \cdot)$ can be computed in $O(mn^2)$ time by Procedure *ComputeNumSolutions* (Theorem 5.1), the corollary follows. ■

Remarks. (1) Note that the total number of optimal reconciliations can grow exponentially as a function of gene tree and species tree size. Throughout this work, however, when analyzing the time/space complexity of our algorithm, we make the assumption that the values $N(\cdot, \cdot)$ can each be stored in some constant-sized memory block. (2) It is worth observing that the approach described above also makes it possible to compute the number of optimal reconciliations when the species tree is dated or, more generally, when constraints are imposed on the mappings or event assignments for some subset of the nodes of the gene tree.

5.2. Sampling optimal reconciliations uniformly at random

Once all the $c(\cdot, \cdot)$ and $N(\cdot, \cdot)$ have been computed, an optimal reconciliation itself can be built by backtracking through the dynamic programming table. To ensure that reconciliations are generated uniformly at random, the idea is to make the choice of mapping assignments based on the number of optimal solutions contained within each choice. For instance, if a node g has already been assigned a mapping, its two children g' and g'' must be assigned mappings jointly based on their joint probability mass.

In general, to output a reconciliation we must assign a mapping and an event (speciation, duplication, or transfer) to each node of G (see Definition 2.1). (Technically, we are also required to assign the transfer edges, but this is trivially accomplished once the mapping and event assignments are in place.) To ensure that the output reconciliation is sampled uniformly at random from the space of all optima, we generate this mapping and event assignment as shown in Procedure *RandomOptimalReconciliation* below. This algorithm assumes that Procedure *ComputeNumSolutions* has already been executed. Let μ denote the minimum reconciliation cost of G and S .

Procedure *RandomOptimalReconciliation*

-
- 1: **for** each $g \in I(G)$ in a pre-order traversal of G **do**
 - 2: **if** $g = rt(G)$ **then**
 - 3: Let P denote the set $\{s \in V(S) : c(rt(G), s) = \mu\}$.
 - 4: For any $s \in P$, assign the mapping $\mathcal{M}(g)$ to be s with probability $\frac{N(g, s)}{\sum_{s \in P} N(g, s)}$.
 - 5: Let $\{g', g''\} = Ch_G(g)$.
 - 6: Let Q denote the set $\{i : b_i = 1\}$ in the context of the Equations 1, 3, or 4, used when computing $N(g, s)$.
 - 7: Choose an a_i , where $i \in Q$, with probability $\frac{N(a_i)}{\sum_{i \in Q} N(a_i)}$.
 - 8: Assign an event type to g based on whether the chosen a_i was derived from $c_\Sigma(\cdot, \cdot)$, $c_\Delta(\cdot, \cdot)$, or $c_\Theta(\cdot, \cdot)$.
 - 9: Consider the sets X' and X'' for the chosen a_i , as defined in Equation 2.
 - 10: Assign the mapping $\mathcal{M}(g')$ to be node s from X' with probability $\frac{N(g', s)}{\sum_{s \in X'} N(g', s)}$.
 - 11: Assign the mapping $\mathcal{M}(g'')$ to be node s from X'' with probability $\frac{N(g'', s)}{\sum_{s \in X''} N(g'', s)}$.
-

Steps 6 through 11 can actually be implemented during the nested post order traversal of Procedure *ComputeNumSolutions* for all possible $g \in I(G)$ and $s \in V(S)$. This has the advantage of reducing the overall space complexity of the algorithm from $O(mn^2)$ to $O(mn)$ since we need not store all the information about the a_i s and the X' and X'' , etc., for later reuse by Procedure *RandomOptimalReconciliation*.

Based on the details of the procedure above, we have the following theorem.

Theorem 5.2. *Procedure RandomOptimalReconciliation generates each optimal reconciliation with equal probability.*

Proof. Observe that the mapping assignment for $g = rt(G)$ (Step 4) is consistent with the goal of generating each optimal reconciliation uniformly at random. We will prove the theorem by induction on the nodes of G . Consider some $g \in I(G)$ and suppose that the procedure fixes the mapping of g to some node s from $V(S)$. It suffices to show that each optimal reconciliation of $G(g)$ with S is now equally likely to be generated by the procedure, that is, each of those optimal reconciliations is generated with probability $1/N(g, s)$. By the proof of Theorem 5.1, we know that the a_i s, for $i \in Q$ (Step 6), divide the reconciliations counted in $N(g, s)$ into disjoint subsets. The procedure chooses one of these a_i s at random based on their probability mass and ensures that the generated reconciliation of $G(g)$ with S is from the chosen a_i (Steps 8 through 11). It remains to show that the procedure generates a reconciliation uniformly at random from among all the reconciliations that make up the chosen a_i . Let $\{g', g''\} = Ch_G(g)$ and recall that the term a_i consists of $\sum_{x \in X'} N(g', x) \times \sum_{x \in X''} N(g'', x)$ reconciliations (Eq. 2). To ensure a uniformly random generation of the reconciliations counted in a_i , the mapping $s' \in X'$ for g' should be chosen with probability $\frac{N(g', s') \times \sum_{x \in X''} N(g'', x)}{N(a_i)}$, which is equal to $\frac{N(g', s')}{\sum_{x \in X'} N(g', x)}$ (by Eq. 2). Similarly, the mapping $s'' \in X''$ for g'' should be chosen with probability $\frac{N(g'', s'')}{\sum_{x \in X''} N(g'', x)}$. Indeed, this is exactly how the procedure assigns the mappings of g' and g'' in Steps 10 and 11. ■

The overall time complexity of our algorithm for generating optimal reconciliations uniformly at random is dominated by that of Procedure *ComputeNumSolutions* and is consequently $O(mn^2)$. This is only a factor of n slower than the fastest known algorithm for the MPR problem (Bansal et al., 2012).

Our implementation of the random sampling algorithm will be made available as part of the next release of the RANGER-DTL software package (Bansal et al., 2012).

6. EXPLORING THE SPACE OF OPTIMAL RECONCILIATIONS

We applied our method to the biological dataset to understand the space of optimal reconciliations for the approximately 4700 gene trees in the dataset. As before, we used event costs $P_\Delta = 2$, $P_\Theta = 3$, and $P_{loss} = 1$ for this analysis. For this study, we focused on understanding how similar the different optimal reconciliations are to each other. To that end, we used our algorithm to sample 500 optimal reconciliations for

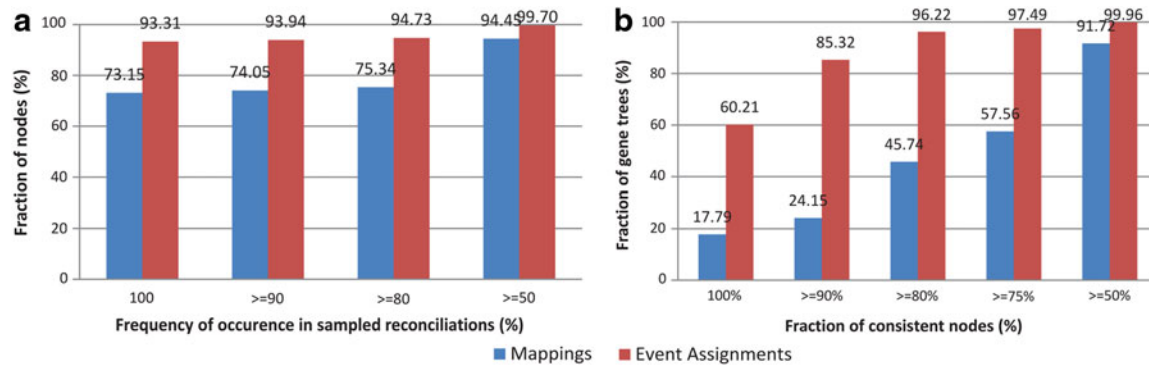


FIG. 4. Stability of mappings and event assignments. The plot in part (a) shows the fraction of internal nodes from the 4699 gene trees that have the same mapping or the same event assignment across at least a certain fraction of the 500 samples. The plot in part (b) plots the fraction of the 4699 gene trees that have at least a certain fraction of their nodes with a consistent mapping or a consistent event assignment across all 500 samples.

each gene tree and wrote a program that reads in these samples and summarizes them as follows: For each internal node in the gene tree we (i) consider the fraction of times that node is mapped to the different nodes of the species tree, and (ii) consider the fraction of times that node is labeled as a speciation, duplication, and transfer event. We used this to investigate the stability of the embedding of the gene tree into the species tree (i.e., the stability of gene node mappings), and the stability of event assignments for the internal nodes of the gene tree.

We first checked to see how stable the gene node mappings were across the internal nodes in all 4699 gene trees. Figure 4a shows the results of this analysis. Overall, we observed that mappings tended to be fairly well conserved across the different multiple optima. For instance, we observed that 73.15% of the internal gene tree nodes had the same mapping across all 500 samples. Recall that only 17% of the gene trees have a unique solution. We also repeated this analysis for event assignments and these results are also shown in Figure 4a. Amazingly, we observed that 93.31% of the nodes had a consistent event assignment across all 500 samples. This suggests that event assignments tend to be highly conserved across the different multiple optima. Thus, even in those instances where there are many different optimal reconciliations, it should be possible to confidently assign event types to most internal nodes of the gene tree (even though the mappings of the nodes themselves may not be consistent across the different multiple optima). This has important implications for understanding gene family evolution, since the inference of orthologs, paralogs, and xenologs depends only on the event assignments for gene tree nodes.

In practice, users are often interested in analyzing the evolutionary history of a specific gene family. We thus asked the following question: Given a gene tree from the biological dataset, what fraction of its nodes can be expected to have (i) a consistent mapping, and (ii) a consistent event assignment across all 500 samples. Figure 4b shows the results of this analysis. The results show that for most gene trees, event assignments are completely consistent across all samples for most of their internal nodes. For instance, we observed that 60.2% of the gene trees have a consistent event assignment for all of their internal nodes, and almost all gene trees had a consistent event assignment for at least half of their internal nodes. As we observed before, gene tree node mappings tend to be more variable, but still, over 91% of the gene trees had a consistent mapping for at least half of their internal nodes. We also tested to see if there was a correlation between the number of optimal reconciliations for a gene tree and fraction of its internal nodes with consistent mappings or consistent event assignments. To our surprise, we found no correlation (Fig. 5). Thus, even if a gene tree/species tree pair have many optimal reconciliations, a large fraction of the gene tree nodes can still be expected to have consistent mappings and event assignments.

It is worth noting that our analysis of this entire ~ 4700 gene tree, 100-taxon dataset, with 500 computed optimal reconciliations for each gene tree, required only about a week of running time on a desktop computer using a single 3 GHz processing core and 4 GB of RAM.

We also analyzed the stability of the mappings and assignments using the dated version of the DTL-reconciliation problem (i.e., the D-MPR problem) and obtained similar results. Specifically, we observed that for the dated version, 72.4% and 94.8% of the internal nodes has consistent mapping and event

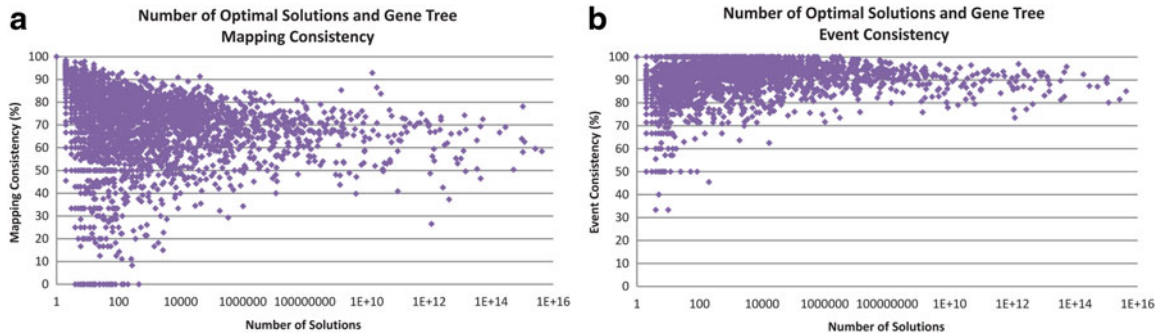


FIG. 5. Consistency of mappings and event assignments by number of optimal reconciliations. **(a)** Fraction of gene tree nodes that have a consistent mapping across all 500 samples. **(b)** Fraction of gene tree nodes that have a consistent event assignment across all 500 samples.

assignments, respectively, in each of the 100 random samples we tested. Thus, the additional restriction on transfers imposed when a fully dated species tree is used appears to have little effect on the consistency of assignments and mappings.

7. APPLICATION TO UNDERSTANDING SENSITIVITY TO EVENT COSTS

The ability to explore the space of multiple optimal reconciliations makes it possible to study the effect of using different event costs on the reconciliation. For instance, one can compare if the mapping or event assignments that are consistent across the multiple optima for a particular event cost assignment are also consistent across a different event cost assignment. Similarly, if one is unsure of which event cost assignment to use, one can try out several different event costs, compute a set of random samples for each event cost assignment, and aggregate the samples from all event cost assignments into a single analysis to understand which aspects of the reconciliation are conserved across the different event cost assignments.

We performed a preliminary study of the effect of using different event costs on the analysis of the biological dataset. Recall that our default event costs are $P_{\Delta} = 2$, $P_{\Theta} = 3$, and $P_{loss} = 1$. For this study, we kept $P_{loss} = 1$ but considered the following combinations of the duplication and transfer costs: (i) $P_{\Delta} = 2$, $P_{\Theta} = 4$, (ii) $P_{\Delta} = 2$, $P_{\Theta} = 2$, (iii) $P_{\Delta} = 3$, $P_{\Theta} = 3$, and (iv) $P_{\Delta} = 1$, $P_{\Theta} = 1$. We computed 100 random samples for each setting of event costs.

We first tested the impact of different cost assignments on the number of inferred speciation, duplication, and transfer events. Since the number of inferred events can vary across the different optimal reconciliations, even for the same fixed event cost assignment, we only considered those event assignments that are supported by all 100 samples (500 samples in case of the default cost assignment). Table 1 shows the results of our analysis. As expected, we observe that as the relative cost of a transfer event or duplication event increases, the number of inferred transfers or duplications, respectively, decreases. For instance, when the event cost assignment is changed from $P_{\Delta} = 2$ and $P_{\Theta} = 3$ to $P_{\Delta} = 2$, $P_{\Theta} = 4$, the number of transfers decreases from 62540 to 56217, with a corresponding increase in the number of speciations and duplications. Note also that the total number of speciations, duplications, and transfers with 100% support

TABLE 1. IMPACT OF EVENT COSTS ON EVENT INFERENCE

	$P_{\Delta} = 2$ $P_{\Theta} = 3$	$P_{\Delta} = 2$ $P_{\Theta} = 2$	$P_{\Delta} = 2$ $P_{\Theta} = 4$	$P_{\Delta} = 3$ $P_{\Theta} = 3$	$P_{\Delta} = 1$ $P_{\Theta} = 1$
Speciation	69174	64473	70582	70762	49973
Duplication	16501	12796	19813	12970	12385
Transfer	62540	67191	56217	65006	71977

This table shows the number of inferred speciation, duplication, and transfer events (inferred consistently across all random samples) for the different event cost assignments.

TABLE 2. IMPACT OF EVENT COSTS ON NUMBER OF OPTIMAL RECONCILIATIONS.

Number of optimal reconciliations	$P_{\Delta} = 2$	$P_{\Delta} = 2$	$P_{\Delta} = 2$	$P_{\Delta} = 3$	$P_{\Delta} = 1$
	$P_{\Theta} = 3$	$P_{\Theta} = 2$	$P_{\Theta} = 4$	$P_{\Theta} = 3$	$P_{\Theta} = 1$
1	16.7%	13.1%	17.8%	15.6%	8.3%
[2, 9]	30.9%	25.7%	30.6%	28.0%	19.8%
[10, 99]	19.5%	20.7%	19.7%	20.1%	21.1%
[100, 9999]	17.4%	20.1%	16.3%	19.4%	20.8%
[10000, 99999]	5.3%	6.4%	4.9%	5.6%	7.2%
$[10^5, 10^{16}]$	10.2%	14.0%	10.7%	11.3%	22.8%

This table shows the fraction (%) of gene trees for which the number of optimal reconciliations lies within the given ranges, for different event cost assignments.

is significantly smaller for the cost assignments $P_{\Delta} = 2$ and $P_{\Theta} = 2$, and $P_{\Delta} = 1$ and $P_{\Theta} = 1$, than for the other cost assignments, indicating more variability in the reconciliations.

Next, we tested the impact of different cost assignments on the number of optimal reconciliations. Table 2 shows the results of our analysis. Note that the number of optimal reconciliations increases significantly for the cost assignments $P_{\Delta} = 2$ and $P_{\Theta} = 2$, and $P_{\Delta} = 1$ and $P_{\Theta} = 1$, compared to the other cost assignments. This is likely related to the observation made above that the total number of speciations, duplications, and transfers with 100% support is significantly smaller for these two cost assignments than for the others.

Finally, we considered the following question: What fraction of the gene tree nodes with consistent mappings (event assignments) under the default costs also have the same consistent mappings (respective event assignments) under the alternative event costs? The results of this analysis for the four combinations of event costs listed above are as follows: For mappings, the fractions are 94%, 83.38%, 92.04%, and 63.97%, respectively. And, for event assignments, the fractions are 92.06%, 91.52%, 96.07%, and 80.37%, respectively. As the analysis indicates, consistent mappings and event assignments tend to be well conserved even when using different event costs. Even with the rather extreme event costs of $P_{\Delta} = P_{\Theta} = P_{loss} = 1$, almost 64% of the consistent mappings and over 80% of the event assignments are conserved.

8. CONCLUSION

In this work, we have presented an efficient and scalable approach to the problem of multiple optimal DTL-reconciliations. Our approach is based on random sampling, and we show how to sample the space of optimal reconciliations uniformly at random, efficiently in $O(mn^2)$ time per sample. The sampling-based approach makes it possible for users to explore the space of optimal reconciliations and to distinguish between stable and unstable parts of the reconciliation. This approach also allows users to investigate the effect of using different event costs on the reconciliation. Our analysis of the biological dataset provides the first real insight into the space of multiple optima and reveals that many, if not most, aspects of the reconciliation remain consistent across the different multiple optima, and that these can be efficiently inferred and used for understanding gene family evolution. We believe that this work represents an important step toward making DTL-reconciliation a practical method for understanding gene family evolution.

Many aspects of the space of optimal reconciliations remain to be explored. For instance, it would be interesting to investigate why so many of the input instances have millions (and more) of multiple optima. In this work, we did not consider the effect of alternative optimal gene tree rootings on the reconciliation space, and we would like to study this further. The ability to handle multiple optima also enables the systematic evaluation of the accuracy of DTL-reconciliation at inferring evolutionary history correctly, and we plan to pursue this further. It would also be instructive to study the effect of using different event costs more thoroughly. In our work, we have only considered optimal reconciliations, and it might be beneficial to also consider slightly suboptimal reconciliations; doing so in a principled way might require the use of a probabilistic model of gene family evolution by duplication, transfer, and loss (e.g., Tofigh, 2009).

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AUTHOR DISCLOSURE STATEMENT

No competing financial interests exist.

REFERENCES

- Bansal, M.S., Burleigh, J.G., Eulenstein, O., and Wehe, A. 2007. Heuristics for the gene-duplication problem: A $\Theta(n)$ speed-up for the local search. In *RECOMB*, 238–252.
- Bansal, M.S., Alm, E.J., and Kellis, M. 2012. Efficient algorithms for the reconciliation problem with gene duplication, horizontal transfer and loss. *Bioinformatics* 28, 283–291.
- Bonizzoni, P., Vedova, G.D., and Dondi, R. 2005. Reconciling a gene tree to a species tree under the duplication cost model. *Theor. Comput. Sci.* 347, 36–53.
- Burleigh, J.G., Bansal, M.S., Eulenstein, O., et al. 2011. Genome-scale phylogenetics: Inferring the plant tree of life from 18,896 gene trees. *Syst. Biol.* 60, 117–125.
- Charleston, M. 1998. Jungles: A new solution to the host-parasite phylogeny reconciliation problem. *Mathematical Biosciences* 149, 191–223.
- Chauve, C., Doyon, J.-P., and El-Mabrouk, N. 2008. Gene family evolution by duplication, speciation, and loss. *J. Comput. Biol.* 15, 1043–1062.
- Chen, K., Durand, D., and Farach-Colton, M. 2000. Notung: dating gene duplications using gene family trees. In *RECOMB*, 96–106.
- Chen, Z.-Z., Deng, F., and Wang, L. 2012. Simultaneous identification of duplications, losses, and lateral gene transfers. *IEEE/ACM Trans. Comput. Biology Bioinform.* 9, 1515–1528.
- Conow, C., Fielder, D., Ovadia, Y., and Libeskind-Hadas, R. 2010. Jane: a new tool for the cophylogeny reconstruction problem. *Algorithm. Mol. Biol.* 5, 16.
- David, L.A., and Alm, E.J. 2011. Rapid evolutionary innovation during an archaean genetic expansion. *Nature* 469, 93–96.
- Doyon, J.-P., Scornavacca, C., Gorbunov, K.Y., et al. 2010. An efficient algorithm for gene/species trees parsimonious reconciliation with losses, duplications and transfers, 93–108 In Tannier, E., ed., *RECOMB-CG*, Vol. 6398 of *Lecture Notes in Computer Science*. Springer, New York.
- Durand, D., Halldórsson, B.V., and Vernot, B. 2006. A hybrid micro-macroevolutionary approach to gene tree reconstruction. *J. Comput. Biol.* 13, 320–335.
- Eulenstein, O., and Vingron, M. 1998. On the equivalence of two tree mapping measures. *Discrete Applied Mathematics* 88, 101–126.
- Goodman, M., Czelusniak, J., Moore, G.W., et al. 1979. Fitting the gene lineage into its species lineage. a parsimony strategy illustrated by cladograms constructed from globin sequences. *Systematic Zoology* 28, 132–163.
- Gorbunov, K.Y., and Liubetskii, V.A. 2009. Reconstructing genes evolution along a species tree. *Molekuliarnaia Biologiia* 43, 946–958.
- Górecki, P., and Tiuryn, J. 2006. Dis-trees: A model of evolutionary scenarios. *Theor. Comput. Sci.* 359, 378–399.
- Koonin, E.V. 2005. Orthologs, paralogs, and evolutionary genomics. *Annual Review of Genetics* 39, 309–338.
- Libeskind-Hadas, R., and Charleston, M. 2009. On the computational complexity of the reticulate cophylogeny reconstruction problem. *J. Comput. Biol.* 16, 105–117.
- Ma, J., Ratan, A., Raney, B.J., et al. 2008. Dupcar: Reconstructing contiguous ancestral regions with duplications. *J. Comput. Biol.* 15, 1007–1027.
- Merkle, D., and Middendorf, M. 2005. Reconstruction of the cophylogenetic history of related phylogenetic trees with divergence timing information. *Theory of Biosciences* 123, 277–299.
- Merkle, D., Middendorf, M., and Wieseke, N. 2010. A parameter-adaptive dynamic programming approach for inferring cophylogenies. *BMC Bioinformatics* 11(Suppl 1), S60.
- Mi, H., Dong, Q., Muruganujan, A., et al. 2010. Panther version 7: improved phylogenetic trees, orthologs and collaboration with the gene ontology consortium. *Nucleic Acids Research* 38(suppl 1), D204–D210.

- Mirkin, B., Muchnik, I., and Smith, T.F. 1995. A biologically consistent model for comparing molecular phylogenies. *J. Comput. Biol.* 2, 493–507.
- Ovadia, Y., Fielder, D., Conow, C., and Libeskind-Hadas, R. 2011. The cophylogeny reconstruction problem is np-complete. *J. Comput. Biol.* 18, 59–65.
- Page, R.D.M. 1994. Maps between trees and cladistic analysis of historical associations among genes, organisms, and areas. *Syst. Biol.* 43, 58–77.
- Rasmussen, M.D., and Kellis, M. 2011. A bayesian approach for fast and accurate gene tree reconstruction. *Molecular Biology and Evolution* 28, 273–290.
- Ronquist, F. 2003. Parsimony analysis of coevolving species associations, 22–64. In Page, R. D. M., ed. *Tangled Trees: Phylogeny, Cospeciation and Coevolution*. The University of Chicago Press, Chicago.
- Rutschmann, F. 2006. Molecular dating of phylogenetic trees: A brief review of current methods that estimate divergence times. *Divers. Distrib.* 12, 35–48.
- Scornavacca, C., Paprotny, W., Berry, V., and Ranwez, V. 2013. Representing a set of reconciliations in a compact way. *J Bio Comp Biol.* 11, 1250025.
- Sennblad, B., and Lagergren, J. 2009. Probabilistic orthology analysis. *Syst. Biol.* 58, 411–424.
- Stolzer, M., Lai, H., Xu, M., et al. 2012. Inferring duplications, losses, transfers and incomplete lineage sorting with nonbinary species trees. *Bioinformatics* 28, 409–415.
- Storm, C.E.V., and Sonnhammer, E.L.L. 2002. Automated ortholog inference from phylogenetic trees and calculation of orthology reliability. *Bioinformatics* 18, 92–99.
- Tofigh, A. 2009. *Using Trees to Capture Reticulate Evolution : Lateral Gene Transfers and Cancer Progression* [Ph.D. thesis]. KTH Royal Institute of Technology, Stockholm.
- Tofigh, A., Hallett, M.T., and Lagergren, J. 2011. Simultaneous identification of duplications and lateral gene transfers. *IEEE/ACM Trans. Comput. Biology Bioinform.* 8, 517–535.
- van der Heijden, R., Snel, B., van Noort, V., and Huynen, M. 2007. Orthology prediction at scalable resolution by phylogenetic tree analysis. *BMC Bioinformatics* 8, 83.
- Vilella, A.J., Severin, J., Ureta-Vidal, A., et al. 2009. Ensemblcompara genetrees: Complete, duplication-aware phylogenetic trees in vertebrates. *Genome Research* 19, 327–335.
- Wapinski, I., Pferrer, A., Friedman, N., and Regev, A. 2007. Natural history and evolutionary principles of gene duplication in fungi. *Nature* 449, 54–61.

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