Minimizing the number of episodes and Gallai’s theorem on intervals

Éva Czabarka\textsuperscript{2,3,5},
University of South Carolina, czabarka@math.sc.edu
László A. Székely\textsuperscript{1,2,3,4,5}
University of South Carolina, szekely@math.sc.edu
Todd Vision\textsuperscript{1,3}
University of North Carolina, tjv@biol.unc.edu

May 2, 2014

Abstract

In 1996, Guigo et al. [Mol. Phylogenet. Evol., 6 (1996), 189–203] posed the following problem: for a given species tree and a number of gene trees, what is the minimum number of duplication episodes, where several genes could have undergone duplication together to generate the observed situation. (Gene order is neglected, but duplication of genes could have happened only on certain segments that duplicated). We study two versions of this problem, one of which was algorithmically solved not long ago by Bansal and Eulenstein \cite{Bansal:2014aa}. We provide min-max theorems for both versions that generalize Gallai’s archetypal min-max theorem on intervals, allowing simplified proofs to the correctness of the algorithms (as it always happens with duality) and deeper understanding. An interesting feature of our approach is that its recursive nature requires a generality that bioinformaticians attempting to solve a particular problem usually avoid.

\textsuperscript{1} This work started at the Phylogeny program of the Isaac Newton Institute, Cambridge.
\textsuperscript{2} This author was supported in part by a Marie Curie Fellowship HUBI MTKD-CT-2006-042794.
\textsuperscript{3} This author was supported in part by the NIH NIGMS contract 1 R01 GM078991-01 and 3 R01 GM078991-03S1.
\textsuperscript{4} This author was supported in part by the NSF DMS contracts 0701111 and 1000475.
\textsuperscript{5} This author was supported in part by DARPA and AFOSR under the contract FA9550-12-1-0405.
1 Introduction

In 1996 Guigo et al. [7] posed the following problem: for a given species tree and a number of gene trees, what is the minimum number of episodes of gene duplication, where several genes could have duplicated in any single episode. The constraints of the problem include some vertices of the gene trees identified as duplication vertices; and duplication vertices have some associated intervals in the species tree, where a duplication of the gene represented by the gene tree must have taken place. We give mathematical definitions in Section 2 and explain the relevance of our results in Section 5.

Several variants of this problem have been investigated: [6], [14], [2], [4], [10], [5]. Bansal and Eulenstein [1] solved a version of this long-standing open problem with a greedy algorithm and proved the correctness of the algorithm by induction.

The purpose of our note is to put these problems and results into proper combinatorial context. There is no need to assume that the trees have no internal vertices of degree two or that they are binary. The intervals associated with the duplication vertices can be defined differently from the definitions in the biology literature. The greedy algorithm still works, furthermore, simple min-max theorems give a good characterization to the minimum number of episodes, even in this more general setting. As usual, the duality allows for more transparent proofs for the correctness of the optimization algorithms.

The min-max theorems are straightforward generalizations to Gallai’s Theorem on intervals (Gallai did not publish actually this theorem, and it was first printed in a paper of Hajnal and Surányi [9]):

**Theorem 1.1** [Gallai] Let us be given a finite family of closed intervals on a straight line. Denote by $\nu$ the size of the largest set of pairwise disjoint intervals, and by $\tau$ the smallest number of points that can cover all intervals i.e. every interval contains at least one of the points. Then $\nu = \tau$ holds.

As a reminder, we reproduce here the proof to Gallai’s Theorem, as our proofs were developed to generalize it. Clearly $\nu \leq \tau$, as disjoint intervals have to be covered by distinct points. We show that $\nu$ points suffice to cover all intervals. Apply the following algorithm recursively until the interval system is empty:

*Pick the leftmost right endpoint from all right endpoints of intervals from the family, and delete all intervals from the system that this point covered. Add the picked point to the list of selected points.*

We picked some right endpoints of intervals that are pairwise disjoint by the construction, therefore these endpoints are at most $\nu$ in number. These right endpoints cover all our intervals by the construction. ♠

---

2
Gallai’s Theorem have been generalized by Surányi (see [8]) essentially with the same proof:

**Theorem 1.2** [Surányi] Let us be given a finite family of subtrees on a tree. Denote by \( \nu \) the size of the largest set of pairwise disjoint subtrees, and by \( \tau \) the smallest number of points that can cover all subtrees i.e. every subtree contains at least one of the points. Then \( \nu = \tau \) holds.

2 Describing the combinatorial problems

Let us be given a finite set \( X \). The elements of \( X \) are called *taxa*. Let us be given a tree \( S \) with root \( R \), such that the leaves of \( S \) are labelled with elements from \( X \) in a one-to-one manner. Root \( R \) is joined by an edge to \( \infty \). We call \( S \) the *species tree* for the taxa in \( X \). When we talk about vertices of \( S \), we exclude \( \infty \).

Let us be given \( k \) *gene trees*, say for \( i = 1, 2, \ldots, k \) the gene tree \( G_i \). The leaves of \( G_i \) are labelled with some taxa from \( X \), but a taxon can occur in more than one leaf, and not all taxa are necessarily represented by a leaf in \( G_i \). A leaf corresponds to one taxon only. We assume that \( G_i \) also has a root \( R_i \) and one more edge going from the root to \( \infty_i \). \( \infty_i \) is not considered a vertex of the gene tree.

Vertices of \( S \) have a natural partial order, namely \( u \geq_S v \), if \( u = v \) or \( u \) separates \( v \) from \( \infty \). When we speak about the interval of \( u \) and \( v \) in \( S \), we mean the interval in this natural partial order. We call \( u \) the *upper endpoint* and \( v \) the *lower endpoint* of this interval. Vertices of \( G_i \) have a natural partial order \( \geq_i \) defined similarly. \( >_S \) and \( >_i \) will refer to strict inequalities in these partial orders.

Assume further that every \( G_i \) has a subset \( D_i \) of its vertices specified that are called *duplication vertices*. For every \( i \) and every \( d \in D_i \), we have an associated path \( P \), which is a subpath of a path connecting \( \infty \) to a leaf in the species tree \( S \). The ordered pair \((P, d)\) will be called the *duplication interval* associated to the duplication vertex \( d \in D_i \), and for more convenient notation we write it as \( P_d \). In this way we maintain names on the duplication intervals which tell which duplication vertex of which gene tree generated the duplication interval. The same intervals can have multiple names as duplication intervals: the same path \( P \) in \( S \) may be assigned as a duplication interval to vertices in different gene trees, and also to several pairs of \( \leq_i \)-comparable (or not \( \leq_i \)-comparable) duplication vertices of the same \( G_i \). Duplication intervals with different names are considered distinct objects although their underlying intervals in \( S \) are the same.

The following monotonicity assumption is made on the associated duplication intervals:

\[
\forall i \forall d, e \in D_i \quad d \geq_i e \rightarrow (\max_S P_d \geq_S \max_S P_e) \land (\min_S P_d \geq_S \min_S P_e).
\] (2.1)
We say that for two duplication intervals $P_e <_i P_d$, if $e <_i d$ for the duplication vertices $e, d$ in $G_i$. A chain of duplication intervals is a sequence of duplication intervals associated to duplication vertices $e_1 <_i e_2 <_i \ldots <_i e_m$ for some $i = 1, 2, \ldots, k$. (We freely change between speaking about chains of duplication intervals and chains of duplication vertices in the gene trees, as they are in bijective correspondence.) We may have several copies of a path in $\mathcal{S}$ present as a duplication interval, and it depends on the label of a particular copy whether it satisfies a $P_e <_i P_d$ type relation or belongs to a certain chain. Some copy of a path may do it, while another may not. Also, it may happen that a single path of $\mathcal{S}$ satisfies a strict $P_e <_i P_d$ ordering with proper duplication vertices $e, d \in D_i$.

Now we have the following models and optimization problems as:

**Discrete model.** Let $V(S)$ denote the vertex set of the species tree $S$. Let $V^*(S)$ denote the extension of $V(S)$ by allowing unlimited number of copies of the vertices. We distinguish these copies from each other, but keep the information on which vertices of $V^*(S)$ are copies of the same vertex of $V(S)$. The elements of $V^*(S)$ inherit the $\geq_S$ partial order, if they are copies of different vertices, and are incomparable when they are copies of the same vertex. We denote this extended partial order by $\geq_{S^*}$.

Consider maps $f : \bigcup_{i=1}^k D_i \rightarrow V^*(S)$, which have the property that for all $i$, the restriction $f|_{D_i} : D_i \rightarrow f(D_i)$ preserves the partial order in the following sense: $[d >_i e$ implies that $f(d) >_{S^*} f(e)$ or $f(d)$ and $f(e)$ are different copies of the same vertex]. A value of $f$ is called a (duplication) episode.

**Objective:** Minimize the quantity $|f(\bigcup_{i=1}^k D_i)|$ over all maps $f$, and/or characterize optimal solutions. (A fast algorithm for this minimization was discovered by Bansal and Eulenstein [1].)

We consider an alternative model as well:

**Continuous model.** Consider the edges of $S$ as line segments that have interior points. Let $int(S)$ denote the union of the set of interior points of all edges of $S$. The partial order $\geq_S$ on $S$ naturally extends to $\tilde{S} = V(S) \cup int(S)$. The extension will be denoted by $\geq_{\tilde{S}}$.

Consider now maps $f : \bigcup_{i=1}^k D_i \rightarrow \tilde{S}$, which have the property that for all $i$, the restriction $f|_{D_i} : D_i \rightarrow f(D_i)$ strictly preserves the partial order [i.e. $d >_i e$ implies $f(d) >_{\tilde{S}} f(e)$]. We still call the values of $f$ (duplication) episodes.

**Objective:** Minimize the quantity $|f(\bigcup_{i=1}^k D_i)|$ over all maps $f$, and/or characterize optimal solutions.

We say that a duplication interval in the species tree is degenerate, if it has only one point. For the continuous model we require, in addition, that the duplication intervals are non-degenerate, as otherwise the problem may not have a feasible solution at all. Note that the minimum number of episodes in the discrete and continuous
models can be different, even if all duplication intervals are non-degenerate.

3 The new min-max theorems

First we discuss the simpler discrete model. Let Λ be an arbitrary index set. For \( \lambda \in \Lambda \), let \( C_\lambda \) be a set of duplication intervals that make a chain with respect to one of the gene orders. We call \( \{C_\lambda : \lambda \in \Lambda\} \) a disjoint chain packing, if for every \( \lambda \neq \lambda' \), elements of \( C_\lambda \) and \( C_{\lambda'} \) do not share vertices in \( S \), i.e. \( (\cup C_\lambda) \cap (\cup C_{\lambda'}) = \emptyset \).

We call \( \sum_{\lambda \in \Lambda} |C_\lambda| \) the value of the disjoint chain packing. Fix now an arbitrary disjoint chain packing, \( \{C_\lambda : \lambda \in \Lambda\} \). Now the number of episodes needed is clearly at least as much as the value of this chain packing, as different members of a chain must belong to different episodes and vertex disjoint chains must use disjoint sets of episodes.

**Theorem 3.1** In the discrete model, the minimum number of duplication episodes equals to the maximum value of a disjoint chain packing.

We will prove the other (non-trivial) inequality in the next section.

We continue with the continuous model. For every \( \lambda \in \Lambda \), let \( C_\lambda \) be a set of duplication intervals that make a chain with respect to one of the gene orders. We call \( \{C_\lambda : \lambda \in \Lambda\} \) an almost disjoint chain packing, if the following restrictions for intersections (in \( S \)) of elements from different chains, \( C_\lambda \) and \( C_{\lambda'} \), hold:

(i) for any \( U \in C_\lambda \) and \( U' \in C_{\lambda'} \), we have \( |U \cap U'| \leq 1 \);
(ii) for any \( U \in C_\lambda \) and \( U' \in C_{\lambda'} \), \( |U \cap U'| = 1 \) imply that the single element of \( U \cap U' \) is the \( >_S \) (upper) endpoint of at least one of \( U \) and \( U' \);
(iii) if \( U \in C_\lambda \) and \( U' \in C_{\lambda'} \) intersect in a single point that is the \( >_S \) endpoint of \( U \), but not the \( >_S \) endpoint of \( U' \), then there is an \( R \in C_{\lambda'} \), such that \( U' >_S R \), \( U \cap R = U \cap U' \), and this common intersection point is the \( >_S \) (upper) endpoint of \( R \) as well.

Of course, duplication intervals from the same chain are allowed to intersect.

Note that condition (iii) means that different chains from an almost disjoint chain packing, as sets in \( \tilde{S} \), may only intersect at nodes of \( S \), and if \( v \) is a node where several chains intersect, then it is the \( >_S \) upper endpoint of all the chains that go through it with at most one exception. The chains that go through \( v \) all go down along different edges from \( v \), and the exceptional chain must contain an interval that has \( v \) as its \( >_S \) upper endpoint.

For \( v \in V(S) \), and an almost disjoint chain packing \( \{C_\lambda : \lambda \in \Lambda\} \), let \( E_\lambda(v) \) denote the number of chains \( C_\lambda \), which have elements with upper endpoint \( v \). Fix
an arbitrary almost disjoint chain packing, \( \{C_\lambda : \lambda \in \Lambda\} \). We call
\[
\sum_{\lambda \in \Lambda} |C_\lambda| - \sum_{v \in V(S), \mathcal{E}(v) \geq 1} (\mathcal{E}_\lambda(v) - 1)
\]
the \textit{value} of the almost disjoint chain packing. Now the number of duplication episodes needed is clearly at least the value of the almost disjoint chain packing, as different members of any chain must belong to different episodes, disjoint intervals also must belong to different episodes, and for any vertex \( v \) with \( \mathcal{E}_\lambda \geq 1 \), we may use \( v \) as the episode for (no more than) one of the intervals from each of the \( \mathcal{E}_\lambda(v) \) chains covering \( v \). Thus, we can save on the duplication intervals containing \( v \), by using \( v, \mathcal{E}_\lambda(v) - 1 \) episodes, compared to not using \( v \) as an episode.

\textbf{Theorem 3.2} \textit{In the continuous model, the minimum number of duplication episodes equals to the maximum value of almost disjoint chain packings.}

We will prove the other (non-trivial) inequality in the next section.

It is easy to see that Gallai’s Theorem 1.1 is a special instance of both Theorems 3.1 and 3.2 when the species tree is a path (only one taxon is present) and every gene tree has a single duplication event.

\section{Proofs}

As the algorithms and the proof of their correctness through the respective min-max theorems are very similar, we describe them in one text, and tell the differences as they arise.

The proof is mathematical induction on the total number of duplication vertices in the gene trees. There is nothing to prove if none of the gene trees contain any duplication vertex, and in this case the empty (almost) disjoint chain packing suffices. The algorithm will remove the duplication designation of certain vertices in the gene trees, but not the vertices themselves; and will solve recursively the reduced problem with the reduced number of duplication vertices. We will also provide (almost) disjoint chain packing for the reduced problem, with the right value, such that the min-max theorem holds by the inductive hypothesis for the reduced problem. Then, case by case, we show that the number of episodes from the reduced problem plus the number of episodes created by our greedy algorithm in the reduction step equals to the size of an (almost) disjoint chain packing for the original problem. This will show simultaneously the optimality of our greedy algorithm and the truth of the corresponding min-max theorem.

So we assume that we already know that the recursive algorithm solves the problem optimally in any instance when the total number of duplication vertices is less
than the current amount and that in these instances a disjoint/almost disjoint chain packing can also be built with value equal to the minimum number of episodes.

Every duplication interval has an \( \leq S \)-upper endpoint. Find a \( \leq S \)-minimal among all \( \leq S \)-upper duplication interval endpoints. Let this vertex of \( S \) be \( P \).

**Discrete model:** Let \( k \geq 1 \) be the largest integer such that \( P \) is \( \leq S \)-upper endpoint of each of the \( k \) elements of some chain \( \leq j \) for the order in a gene tree \( G_j \), say \( L_1 \leq j L_2 \leq j \cdots \leq j L_k \). Remove the duplication designation of any vertex \( d \) in any gene tree \( G_i \), if \( P \) belongs to the duplication interval of \( d \) and no \( <_i \)-chain of duplication vertices in \( G_i \) with maximum element \( d \) has length \( k + 1 \).

By induction, the same recursive algorithm solves the reduced episode problem optimally such that the min-max theorem holds for the reduced problem. Add \( P \) with multiplicity \( k \) to the system of episodes. We construct recursively a disjoint chain packing providing the same value, through the following two cases:

(i) no chain in the optimal disjoint chain packing for the reduced problem covers \( P \). Add the chain \( \{L_1 \leq j L_2 \leq j \cdots \leq j L_k\} \) to the disjoint chain packing for the reduced problem—note that we still have a disjoint chain packing. We have the following chain of inequalities: the minimum number of episodes in the original problem is at most the minimum number of episodes in the reduced problem + \( k \), which equals to the maximum value of a disjoint chain packing in the reduced problem + \( k \), which is at most the maximum value of a disjoint chain packing in the original problem. We already know the trivial inequality for the min-max theorem, hence in this case our algorithm provides the same number of episodes as the value of a disjoint chain packing.

(ii) a chain \( C \) in the optimal disjoint chain packing for the reduced problem covers \( P \). Let the lowest element in the chain \( C \) correspond to the duplication interval \( U \). By the choice of \( P \), \( P \) must be in \( U \). The duplication vertex \( d \), which is responsible for \( U \), has not been deleted from the list of duplication vertices. This means that \( d \) is the maximum element in a \((k + 1)\)-chain \( C' \) of duplication vertices in his gene tree. Merge \( C \) and \( C' \) into a single chain (it is possible as \( d \) was lowest element \( C \) but highest in \( C' \)), and add the merged chain to the optimal disjoint chain packing for the reduced problem to obtain a disjoint chain packing for the original problem. The number of episodes that we use for the original problem equals to the value of the disjoint chain packing that we constructed for the original problem.

In both cases, we constructed a disjoint chain packing, whose value is the same as the number of episodes constructed, and hence the induction proof is complete.

**Continuous model:** Note that \( P \) is not a leaf vertex in \( S \), as duplication intervals in the continuous model are non-degenerate. Assume that \( e_1, e_2, \ldots, e_\ell \) are the edges leaving \( P \) in directions different from \( \infty \) in \( S \). For \( j = 1, 2, \ldots, \ell \), let \( H_j \) denote a longest chain of duplication intervals over all gene trees with the following properties:
(α) the upper endpoint of every duplication interval from the chain is $P$.
(β) every duplication interval of the chain uses the edge $e_j$.

Set $d_j = |H_j|$. As we will not need the $d_j = 0$ terms, assume that only those edges leaving $P$ are enumerated on which $d_j \geq 1$, and for convenience those edges are still labelled as $1, 2, ..., \ell$.

We create a reduced problem by removing the duplication vertex designation of certain vertices in the gene trees. Assume that $I_f$ is a duplication interval containing $P$, with vertex $f \in D_i$ from the gene tree $G_i$. We remove the duplication designation of $f$ and the duplication interval $I_f$ if
(a) $P$ is the upper endpoint of $I_f$, or
(b) $P$ is a vertex of $I_f$ but not an endvertex (so $I_f$ uses some $e_j$ edge from $P$) and in $G_i$, there is no chain of duplication vertices of length exceeding $d_j$ in which $I_f$ is the top element, or
(c) $P$ is the lower endpoint of $I_f$, and for every $j = 1, 2, ..., \ell$, in $G_i$ there is no chain of duplication vertices of length exceeding $d_j$ in which $I_f$ is the top element and all other elements have duplication intervals passing through $e_j$.

By mathematical induction, the same recursive algorithm solves the reduced episode problem optimally such that the min-max theorem holds for the reduced problem. Add to the list of episode locations the following points in $\tilde{S}$: $P$ itself and $d_j - 1$ distinct points from the interior of $e_j$ for every $j = 1, 2, ..., \ell$. We construct recursively an almost disjoint chain packing providing the same value, through the following two cases:

(i) no chain in the optimal almost disjoint chain packing for the reduced problem covers $P$. Add to this almost disjoint system of chains, which provides the min-max result for the reduced episode problem by hypothesis, a length $d_j$ chain for every $j = 1, 2, ..., \ell$ from a gene tree, such that every duplication interval of this length $d_j$ chain has upper endpoint $P$ and uses the edge $e_j$. It is easy to see that we obtained an almost disjoint chain packing for the original problem. Simple calculation shows, analogously to the discrete case, that from the min-max result for the reduced problem, we obtain that in original problem the number of episodes equals to the value of the almost disjoint chain packing, as both sides increase by $1 + \sum_{j=1}^{\ell} (d_j - 1) = (\sum_{j=1}^{\ell} d_j) - (\ell - 1)$.

The alternative of (i) is that one or more chains in the optimal almost disjoint chain packing for the reduced problem covers $P$. Observe that $P$ may belong to two chains only if $P$ is upper endpoint of some elements of one of the chains. However, (a) has removed those elements. We are left with:

(ii) a single chain $C$ (corresponding to some gene tree $G_i$) in the optimal almost disjoint chain packing for the reduced problem covers $P$. Let the smallest element in this chain be $U$, so $P \in U \subset C$ by the choice of $P$.
The duplication vertex \( d \) that is responsible for \( U = U_d \) has not been deleted. According to the removal rules, \( P \) cannot be the upper endpoint of \( U \). Therefore \( P \) is an internal point or lower endpoint of \( U \).

If \( P \) is an internal point of \( U \) and \( U \) passes through \( e_j \), then there is a chain \( C' \) of duplication vertices in the gene tree corresponding to \( U \), in which \( U \) is the \((d_j + 1)\)th element. Create the almost disjoint system of chains for the original problem from the optimal almost disjoint system of chains for the reduced problem in the following way: replace \( C \) with \( C \cup C' \), which is still a chain; and for \( t = 1, 2, ..., \ell, t \neq j \), add the chain \( H_t \). Indeed, we obtain an almost disjoint chain packing. Simple calculation shows, analogously to the discrete case, that from the min-max result for the reduced problem, we obtain that the number of episodes equals to the value of the almost disjoint chain packing in the original problem, as both sides increase by \( 1 + \sum_{j=1}^{\ell} (d_j - 1) = \left( \sum_{j=1}^{\ell} d_j \right) - (\ell - 1) \).

If \( P \) is the lower endvertex of the duplication interval \( U \) that comes from the gene tree \( G_i \), then for some \( j \), there is a \((d_j + 1)\)-chain \( C' \) in \( G_i \) with top element \( U \), and all other elements of this chain use \( e_j \) in their duplication intervals. Create the almost disjoint system of chains for the original problem from the optimal almost disjoint system of chains for the reduced problem in the following way: replace \( C \) with \( C \cup C' \), which is still a chain; and for \( t = 1, 2, ..., \ell, t \neq j \), add the chain \( H_t \). Indeed, we obtain an almost disjoint chain packing. Simple calculation shows, analogously to the discrete case, that from the min-max result for the reduced problem, we obtain that the number of episodes equals to the value of the almost disjoint chain packing in the original problem, as both sides increase by \( 1 + \sum_{j=1}^{\ell} (d_j - 1) \).

Cases (i) and (ii) together will prove the correctness of the algorithm and the min-max theorem for the original problem.

There is one more thing to check, namely that the episodes selected for the reduced problem are distinct from the episodes selected at \( P \) and on the \( e_j \) down-edges for \( j = 1, 2, ..., \ell \). The episodes selected in internal vertices of the \( e_j \) edges are no longer in any duplication interval in the reduced problem, and therefore they cannot be selected there. In the reduced problem, the non-internal vertices selected for episodes are upper endpoints of some duplication intervals, while \( P \) is no longer the upper endpoint of any duplication interval in the reduced problem.

\section{5 Relevance for bioinformatics}

Ohno was among the first to recognize the importance of gene and genome duplications \cite{Ohno}, and the resulting opportunity for evolutionary change afforded by genetic redundancy. Gene duplication, and subsequent gene loss, are the primary drivers of changes in gene content. Rates of duplication and loss also have been shown to vary...
among lineages, and gene loss in particular is greatly elevated after whole-genome duplication events, which have occurred many times in the evolution of the eukaryotes [3,11,12]. (While gene content alone has been used for reconstruction of species trees [15], it is very sensitive to parallel or convergent gains and losses, and has not seen wide application.)

Gene trees may differ from the species tree and from each other because of repeated gene duplication and gene loss. Gene loss may eliminate the gene from a species. A species may have more than one representation in the gene tree as a result of gene duplication.

There are two possible explanations of finding duplicate genes: early genome duplications and subsequent substantial gene loss, and occasional duplication of small groups of consecutive genes, not requiring the assumption of substantial gene loss. The latter event is called duplication episode. Clearly both mechanisms are present, an early vertebrate tetraploidization seems generally accepted. (Duplication of ”medium length” segments seem unlikely.)

To reconstruct the likely history of the gene content, we should know the cost associated with genome duplication, gene loss, and duplication episodes. We do not know those costs. Minimization problems for duplication episodes look for a most parsimonious explanation.

The bioinformatics literature identifies duplication vertices in the gene trees. For every duplication vertex, the LCA (least common ancestor) mapping designates a vertex in the species tree, which is a lower bound in the $\leq_S$ partial order for the point in the species tree $S$, where the duplication of the gene could have happened. There is no absolute upper bound on this gene duplication, but the more branchings follow the gene duplication in the species tree, the more gene losses should be assumed. Therefore a common parsimony approach is to allow the shortest duplication interval for this gene: the edge between the lower bound vertex and its parent in the species tree. The assumption (2.1) follows from the way of assigning duplication intervals to duplication points in [1] and before in the literature.

In the theorems above we have assumed that all duplication intervals are closed and the upper and lower endpoints of the intervals are vertices of the species tree $S$. Let us first consider the continuous model. Without loss of generality we may assume that the duplication intervals end at vertices of $S$ by simply subdividing edges of $S$ if necessary, as we did not assume that $S$ was binary. As to the assumption of the intervals being closed: the proof really only used that the intervals are closed upwards, i.e. they include their upper endpoint in the $>_S$ ordering.

It is easy to see that the discrete model can be viewed as follows: using $k$ copies of a vertex $v$ is equivalent with placing $k$ duplication episodes on the edge leading from $v$ to its parent (not using the parent). Therefore in effect the discrete model is
equivalent with using assuming that all duplication intervals are of the form \([x, y]\), where \(y\) is an ancestor of \(x\). Thus, our theorem for the discrete model follows from the theorem on the continuous model.

Bansal and Eulenstein [1] solved the episode minimization problem for the discrete model. We leave the decision for the biology literature when the discrete or the continuous model is to be used. We provided duality results for both models. Surányi’s Theorem [12] can be understood as a min-max result for the so-called episode clustering problem: we are given duplication intervals and we want to cover the duplication intervals with the minimum number of points, while we do not require that that duplication episodes follow in strict order.

References

[1] M. S. Bansal, O. Eulenstein The multiple gene duplication problem revisited, *Bioinformatics*, 24(13), (2008), 132–138.

[2] M. S. Bansal, O. Eulenstein, An \(\Omega(n^2/\log n)\) speed-up of TBR heuristics for the gene-duplication problem *IEEE/ACM Transactions on Computational Biology and Bioinformatics (TCBB)* 5 (2008), Issue 4, 514–524.

[3] F. Brunet, H. Crollius, M. Paris, J. Aury, P. Gibert, O. Jaillon, V. Laudet, and M. Robinson-Rechavi, Gene loss and evolutionary rates following whole-genome duplication in teleost fishes, *Mol. Biol. and Evol.*, 23(9) (2006), 1808–1816.

[4] G. Burleigh, M. S. Bansal, O. Eulenstein, T. J. Vision, Inferring species trees using genome duplication episodes, *BCB ’10 Proceedings of the First ACM International Conference on Bioinformatics and Computational Biology*, 198–203.

[5] Wen-Chieh Chang, G.J. Burleigh, D.F. Fernández-Baca, and O. Eulenstein, An ILP solution for the gene duplication problem, *BMC Bioinformatics* 12(Suppl 1)(2011), S14. doi: 10.1186/1471-2105-12-S1-S14

[6] M. Fellows, M. Hallett, U. Stege, On the multiple gene duplication problem, in: *9th International Symposium on Algorithms and Computation* (ISAAC ’98), Taegen, Korea, LNCS 1533, 1998, pp. 347–356.

[7] R. Guigo et al. , Reconstruction of ancient molecular phylogeny, *Mol. Phylogenet. Evol.*, 6 (1996), 189–203.

[8] A. Gyárfás, J. Lehel, A Helly type problem in trees, in: *Combinatorial Theory and its Applications*, eds. P. Erdős, A. Rényi, V. T. Sós, North-Holland, Amsterdam, 1970, 571–584.
[9] A. Hajnal and J. Surányi, Über die Auflösung von Graphen in vollständige Teilgraphen, *Annales Univ. Sci. Bud. Eötvös* 1(1958), 115–123.

[10] C. Luo, M. Chen, Y. Chen, R. Yang, H. Liu, K. Chao, Linear-time algorithms for multiple gene duplication problems, *IEEE Transactions on Comp. Biol. and Bioinf.*, (2009).

[11] M. Lynch and J. Conery, The evolutionary fate and consequences of duplicate genes, *Science*, 290(5494), (2000), 1151–1155.

[12] J. H. Nadeau and D. Sankoff, Comparable rates of gene loss and functional divergence after genome duplications early in vertebrate evolution, *Genetics*, 147(1997), 1259–1266.

[13] S. Ohno, *Evolution by gene duplication*, Springer-Verlag, 1970.

[14] R.D.M. Page, J.A. Cotton, Vertebrate phylogenomics: reconciled trees and gene duplications, *Pacific Symposium on Biocomputing* (2002) 536–547.

[15] B. Snel, P. Bork, and M. Huynen, Genome phylogeny based on gene content, *Nature Genetics*, 21 (1999), 108–110.