

Quartet-based phylogeny reconstruction from gene orders^{*}

Tao Liu¹, Jijun Tang², and Bernard M.E. Moret¹

¹ Department of Computer Science, U. of New Mexico, Albuquerque, NM 87131

² Department of Computer Science & Engineering, U. of South Carolina, Columbia, SC 29208

Abstract. Phylogenetic reconstruction based on gene rearrangements is attracting increasing attention from biologists and computer scientists. Methods used in reconstruction include distance-based methods, parsimony methods using sequence encodings, and direct optimization. The latter, pioneered by Sankoff and extended by us with the software suite GRAPPA, is the most accurate approach; however, its exhaustive nature means that it can be applied only to small datasets (of fewer than 15 taxa). While we have successfully scaled it up to 1,000 taxa by integrating it with a disk-covering method, yielding DCM-GRAPPA, the recursive decomposition in the DCM may require many levels of recursion to handle datasets with 1,000 or more taxa. In order to handle larger datasets and reduce the need for recursive decomposition, we investigate quartet-based approaches, which directly decompose the datasets into subsets of four taxa each. Such approaches have been well studied for sequence data, but not for gene-order data. We give an optimization algorithm for the NP-hard problem of computing optimal trees for each quartet, present a variation of the dyadic method (using heuristics to choose suitable short quartets), and use both in simulation studies. We find that our quartet-based method can handle more taxa than the base version of GRAPPA, thus enabling us to reduce the number of levels of recursion in DCM-GRAPPA, but is more sensitive to the rate of evolution, with error rates rapidly increasing when saturation is approached.

1 Introduction

Modern techniques can yield the ordering and strandedness of genes for genomes; each chromosome can then be represented by an ordering of signed genes, where the sign indicates the strand. Rearrangement of genes under inversion, transposition, and other operations such as duplications, deletions and insertions, is an important evolutionary mechanism [7]. Reconstructing phylogenies from gene-order data has been studied intensely since the pioneering papers of Sankoff [2, 23]. Because they capture the complete genome, gene-order data do not suffer from the gene tree vs. species tree problem; and because rearrangements of genes are *rare genomic events* [19], gene-order data enable the reconstruction of evolutionary events far back in time. Many biologists have embraced this new source of data in their phylogenetic work [7, 17, 18], while computer scientists are slowly solving the difficult problems posed by the manipulations of these gene orders [16]. Studies conducted by our group [15, 26, 29–31] confirm that gene-order data support very accurate reconstructions.

^{*} Contact author: Bernard Moret, moret@cs.unm.edu

The main software package for analyzing gene-order data is GRAPPA, based on the BPAanalysis software of Sankoff and Blanchette [23]. GRAPPA achieved a billion-fold speed-up over BPAanalysis [15]; it can analyze datasets of 13 organellar genomes in 20 minutes on a laptop. Extensive testing has shown that the trees returned by GRAPPA are better than those returned by other methods based on gene orders, such as distance-based methods and parsimony based on encodings [6, 31]. However, since GRAPPA examines every possible tree, it only handle only small datasets—a 17-taxon analysis would take a month on today’s most powerful computers. We integrated GRAPPA with DCM, a divide-and-conquer approach pioneered by Warnow [11], to produce DCM-GRAPPA [29], which can analyze datasets of up to 1,000 taxa without loss of accuracy.

DCM-GRAPPA works in three steps: it first decomposes the dataset into overlapping subproblems (disks), then runs GRAPPA on the subproblems, and finally uses a specialized supertree method [20] to build a tree for the original dataset from the trees returned by GRAPPA for the subproblems. Because the decomposition technique of DCM can still produce subproblems too large for GRAPPA to handle, we call DCM recursively until each subproblem size falls below a given threshold. Because the threshold is small (14 taxa), large problems require many levels of recursive decomposition, which is time-consuming and also risks propagating and amplifying errors in the assembly of the subtrees. On 1,000 taxa, DCM-GRAPPA needs 6–7 levels of recursion if limited to disks of at most 14 taxa, but only 2–3 levels if allowed disks of up to 20 taxa.

One can also decompose the set of taxa into the smallest possible subsets for which meaningful answers exist, namely *quartets*, sets of four taxa. (Sets of two or three taxa can produce only one tree, but a quartet can give rise to three distinct unrooted trees.) While there are many such quartets, their tiny size should make them easy to compute. If every quartet tree is computed correctly from noiseless data, then there is a single tree compatible with all $\binom{n}{4}$ quartet trees and that tree is the true tree [4]; in practice, of course, many of the quartet trees are in error and no single tree is compatible with all quartet trees. With sequence data, biologists have long used the heuristic method known as quartet-puzzling [25], while computer scientists have developed several theoretical methods, such as quartet cleaning [1, 3, 12]—see [24] for a review and experimental comparison of these methods. In the case of gene orders, however, finding the best quartet tree is NP-hard—it includes finding the median of three genomes, a known NP-hard problem [5], as a special case.

We present algorithmic and experimental results that lead to a reconstruction method from gene-order data which overcomes some of the problems associated with quartet methods. After some background review, we describe in Section 3 our exact method to compute optimal quartet trees under breakpoint distances

and present experimental results on our approach to resolving the quartets as well as on our use of the dyadic inference rule [10] to obtain a sufficient set of quartets from a selected subset of short quartets (inspired from the short-quartet method [8, 32]). In Section 5, we summarize our experiments on simulated and biological datasets, the results of which suggest that our method can produce accurate topologies (much more accurate than neighbor-joining) for datasets of up to 25 taxa within reasonable time, provided that the genomes are large enough to avoid saturation.

2 Definitions and Notation

A quartet is a quadruple of taxa; a quartet tree is an unrooted binary tree with four leaves labelled by these taxa. Given a quartet $\{a, b, c, d\}$, we say that a quartet tree on this set is *unresolved* if it is a star (four edges, each touching a leaf) and denote it by $(abcd)$. If the quartet tree has an internal edge separating two pairs of leaves we say that it is *resolved* and, if the pairs are a, b and c, d , denote it by $ab | cd$. The four possible quartet trees induced by a quartet are depicted in Figure 1. We will use quartet in lieu of quartet tree when the sense is clear.

Quartet $ab | cd$ agrees with tree T if all four of its taxa are leaves of T and the path from a to b in T does not intersect with the path from c to d in T . Equivalently, $ab | cd$ agrees with a tree if the subtree induced in T by the four-taxon subset $\{a, b, c, d\}$ is the quartet tree itself. Quartet $ab | cd$ is *in error* with respect to the tree T if it does not agree with T . If Q_T denotes the set of all quartets that agree with T , then T is uniquely characterized by Q_T ; moreover, T can be reconstructed in polynomial time from Q_T [8]. (Of course, the set Q of $\binom{n}{4}$ quartets that we can construct is only an approximation of Q_T .) In Figure 2, quartet $ac | bd$ would be in error, since it does not appear in Q_T .

Quartet-based methods operate in two phases. First, they construct a set Q of resolved quartets—usually by determining the preferred tree for each of the $\binom{n}{4}$ quartets. Because each dataset has size 4, any phylogenetic method can be used to estimate the quartet tree, including maximum likelihood and maximum parsimony (see [27]), neighbor-joining [21], the relaxed four-point method [9], and the ordinal quartet method [13]. In the second phase, the resolved quartets

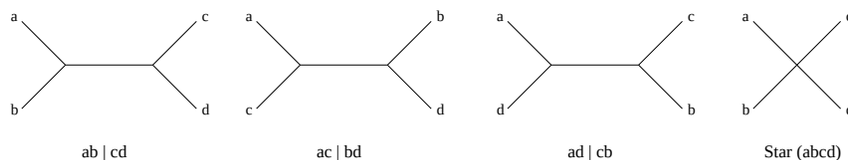


Fig. 1. The four possible quartet trees for quartet $\{a, b, c, d\}$.

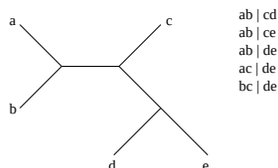


Fig. 2. An evolutionary tree T and its set Q_T of induced quartet trees.

are used to build a single tree on the full set of taxa. This second phase is simple when all quartets are compatible, but a challenge when some of the quartets conflict with others—a common occurrence when insufficient data are present [24].

3 Inferring Quartet Topologies

We can identify maximally parsimonious quartet trees by examining each of the three possible trees, assigning gene orders to the two internal nodes so as to minimize the score (the sum of the lengths of the five edges) of each tree, and returning the tree with the lowest score. However, identifying such gene orders is NP-hard even for just one internal node and the simplest distance measures [5].

An easy approach is to use GRAPPA to construct a tree for each quartet; although the result need not be optimal, our experiments, as well as earlier ones [14], show that optimality is reached in most realistic cases. If we limit ourselves to breakpoint distances, we can solve the NP-hard optimization problem directly, using variants of the reduction to TSP devised by Sankoff and Blanchette [22]; we devised two such variations, which we call $Qtsp$ and $Qedge$.

Sankoff’s reduction to TSP can be summarized as follows. Given three genomes with n genes, we build the complete graph K_{2n} on the $2n$ vertices $g_1, -g_1, g_2, -g_2, \dots, g_n, -g_n$. Edge (g, h) in this graph is assigned a weight as follows: if we have $g = -h$, then we set the weight to a large negative value to ensure that (g, h) is part of any solution, otherwise we set it to $3 - \text{adj}(g, h)$, where $\text{adj}(g, h)$ is the number of times that $-g$ and h are adjacent in the given genomes. If $s = s_1, -s_1, s_2, -s_2, \dots, s_n, -s_n$ is the solution to the TSP, then the median of the given genomes is $g = s_1, s_2, \dots, s_n$.

This result applies to one unknown genome, but we need to identify two such for quartets. Our first algorithm views the two as forming a pair and remaps the problem into a universe where pairs form the unit of computation and where a single tour (of pairs) defines both genomes; our second algorithm retains the original formulation, but looks for a pair of tours.

3.1 The $Qtsp$ Method

We look for one permutation of size n , each entry of which consists of a pair of genes (g_i, g_j) . Let the two desired internal genomes be e and f , where genome

e is connected to genomes a , b , and f , and genome f is connected to genomes c , d , and e . Build a complete graph, here on $(2n)^2$ vertices, each a pair of signed genomes. For each edge $\{(g_{i1}, g_{i2}), (g_{j1}, g_{j2})\}$ in this graph, we set the weight of the edge as follows: if we have $g_{i1} = g_{j1}$ and $g_{i2} = g_{j2}$, then we set the weight to a large negative value to ensure that this edge is part of any solution, otherwise, we set it to $4 - u_1(g_{i1}, g_{j1}) - u_2(g_{i2}, g_{j2})$, where $u_1(g_{i1}, g_{j1})$ is the number of times $-g_{i1}$ and g_{j1} are adjacent in the genomes a and b and $u_2(g_{i2}, g_{j2})$ is the number of times $-g_{i2}$ and g_{j2} are adjacent in the genomes c and d .

Proposition 1. *If $s = (s_{11}, s_{12}), (-s_{11}, -s_{12}), \dots, (s_{n1}, s_{n2}), (-s_{n1}, -s_{n2})$ is the solution to the TSP on G , then an optimal assignment for the internal genomes is $e = s_{11}, s_{21}, \dots, s_{n1}$ and $f = s_{12}, s_{22}, \dots, s_{n2}$.*

After transforming the problem, we can use the efficient TSP routine of GRAPPA to search for the optimal solution (after modifying it to introduce one more bucket of costs, since now the edge costs of interest are 1, 2, and 3, not just 1 and 2). The problem is that the instance thus created is of size quadratic in the number of genes; combined with the extra bucket of costs, the large instance size makes it difficult to obtain solutions to sizeable instances.

3.2 The Qedge Method

This method uses the original TSP formulation, of size linear in the number of genes, but seeks simultaneously to optimize tours in two separate graphs. Let a , b , c , d , e , and f be as before. We set up two complete graphs on $2n$ vertices each—one for e and one for f . For each edge $\{g, h\}$, we set its weight as follows: if we have $g = -h$, we set the weight to a large negative value to force its inclusion; otherwise, we set it to $2 - u(gh)$, where $u(gh)$ is the number of times $-g$ and h are adjacent in the genomes a and b (for an edge in the first graph) or in the genomes c and d (for an edge in the second graph). Now, when adding an edge to the two tours under construction, we can either pick different edges from the two graphs, each with the minimum weight in its own graph, and add one breakpoint between e and f to the total cost; or pick the same edge in both graphs, even if not locally optimal, thereby saving a breakpoint between e and f ; our algorithm computes the cost of each choice and picks the choice of lower cost.

3.3 Experimental Results for Qtsp and Qedge

We ran tests for these two methods and GRAPPA on quartets of 10, 20, 30, 40, and 50 genes under evolutionary rates (the expected numbers of events per edge of the model tree) of $r = 1, 2, 3, 4$, and 5. The running times of Qtsp and Qedge are shown in Figure 3. As expected, Qedge runs much faster than Qtsp; however, its running time depends strongly on the quartet score, so that it may prove unusable in the reconstruction of large trees, where many of the quartets will have large scores. Both Qtsp and Qedge reconstructed the same optimal quartet

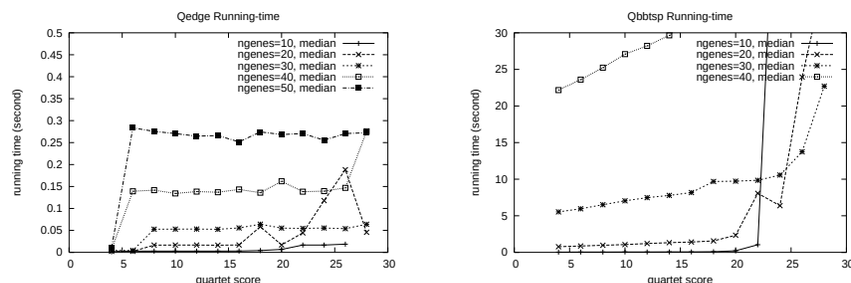


Fig. 3. Running times of Q_{tsp} (left) and Q_{edge} (right) as a function of quartet score

trees (except for ties), which were the same as the model tree in 97% of cases (although their scores were usually lower than the model tree scores); most of the 3% came from the 10-gene case, which gets quickly saturated for evolutionary rates above $r = 3$. GRAPPA did almost as well: 96% of the quartets it returned matched the model tree—and it returned answers in a few microseconds. Since reconstructing phylogenies from quartets requires the computation of a large set of quartets, the running time is critical. Therefore, based on our results, we chose GRAPPA as the method to resolve the quartets.

4 Phylogenetic Reconstruction from Quartets

The computational challenge of quartet recombination (building a single tree from the collection of quartet trees) is how to deal with quartet errors. Most optimization problems related to tree reconstruction from quartets are NP-hard, such as the Maximum Quartet Compatibility problem [12], which seeks a tree T for a given set of quartet Q such that $|Q_T \cap Q|$ is maximized. Various methods have been designed to handle quartet errors. The dyadic-closure method simply issues an error message and quits [10]. The Q^* method seeks the maximally resolved tree T' that obeys $Q(T') \subseteq Q$, a very conservative method that generally produces many polytomies [4]. Quartet-cleaning methods establish a bound on the number of quartet errors around each reconstructed tree edge [1, 3, 12]. None of these methods produces satisfactory results on sequence data [24]. It is theoretically possible to produce the true tree by selecting a subset of short quartets and adding further quartets according to an inference rule [8], but this result assumes perfect data and gives no simple method by which to select the subset of quartets. We thus set out to design a selection rule and investigate its performance, using the dyadic inference rules [10]:

1. If $ab | cd$ is a valid quartet, so are $ba | cd$ and $cd | ab$.
2. If $ab | cd$ and $ac | de$ are valid quartets, so are $ab | ce$, $ab | de$, and $bc | de$.
3. If $ab | cd$ and $ab | ce$ are valid quartets, so is $ab | de$.

4.1 Selecting a Subset of Quartets

Figure 4 shows two possible resolutions for quartet $\{a, b, c, d\}$. In the first topology, the two pairs of genomes $\{a, b\}$ and $\{c, d\}$ are far apart from each other, but in the second topology the two pairs $\{a, c\}$ and $\{b, d\}$ are quite close: the first topology is more likely to be correct, an observation supported by the relaxed four-point method [9]:

Compute pairwise distances among a , b , c , and d ; return $ab \mid cd$ if we have $d_{ab} + d_{cd} < \min(d_{ac} + d_{bd}, d_{ad} + d_{bc})$, but return a star if all sums are equal.



Fig. 4. Two quartet trees; the left has a higher probability of correctness

Since computing all $\binom{n}{4}$ quartets takes too long, we can use this relaxed four-point method to choose quartets and reduce the overall running time. After resolving the quartets, we can assign a weight to each resolved quartet to measure our confidence in that quartet: for example, we can use the inversion distance between the two internal nodes.

4.2 Fixing Quartet Errors

Although GRAPPA is reliable and although we can pick only quartets of larger weight, quartet errors still arise, especially when we are forced to select some quartets of low weight in order to resolve every internal tree edge. We propose a simple new method, quite distinct from quartet cleaning, to handle errors. Since the quartets are weighted and since we place more trust in quartets of higher weight, we examine the source of quartet errors and, whenever two quartets are incompatible, we remove the one with lower weight.

Starting from a large initial set of resolved quartets only returns us to a version of the NP-hard problem Maximum Quartet Compatibility. Instead, we proceed incrementally. We select a high weight threshold and only retain quartets (computed on the fly with GRAPPA) with weights above that threshold; if we find quartet errors, we remove the incompatible quartets of lower weight. We then apply the dyadic inference rules to augment our collection of compatible quartets. Finally, if the resulting set of quartets fully resolves the tree, we are done (a method like Q^* will recover the tree), otherwise we lower the threshold and add to our set the newly eligible quartets. By controlling the decrease in the weight threshold, we can control the tradeoff between running time and quality.

Since we do not know the weight of a quartet until we resolve it, but want to avoid resolving useless quartets, we need a fast method to select quartets to resolve. Given quartet $q = \{a, b, c, d\}$, define the *width* of q as

$$q_w = \max(d_{ab} + d_{cd}, d_{ac} + d_{bd}, d_{ad} + d_{bc}) - \min(d_{ab} + d_{cd}, d_{ac} + d_{bd}, d_{ad} + d_{bc})$$

As q_w increases, the two pairs of genomes move further apart and the weight increases: hence we can decide which quartets to resolve by comparing their width with the weight threshold. Even if the threshold is lowered to zero, the set of compatible quartets may remain inadequate to resolve the tree—in which case we have no choice but to leave these unresolved polytomies in the output.

5 Experimental Results

If the true tree has an edge defining a bipartition with no equivalent in the reconstructed tree, that edge is a *false negative (FN)*; conversely, if the reconstructed tree has an edge with no equivalent in the true tree, that edge is a *false positive (FP)*. FP edges are more problematic than FN edges.

We generated model tree topologies from the uniform distribution on binary trees, each with 12, 16 and 20 leaves respectively. On each tree, we evolved signed permutations of 40, 60 and 80 genes, using evolutionary rates (the expected numbers of events along a tree edge) of 2, 4, 6. For each combination of parameters, we generated 20 trees; the final results are averaged on the 20 datasets. We computed quartets using GRAPPA and built the resulting tree using our algorithms for selecting quartets of high weight, eliminating conflicting quartets, and expanding the set with the dyadic rules. Figure 5 shows FP and FN rates for datasets with 80 genes. Our method did well, but saturation (high evolutionary rates leading to ill-defined estimates of distances) causes a small increase in the error rate. This observation is confirmed by our results on datasets with 40 genes, where saturation occurs much sooner and the results are unacceptable, as seen in Figure 6. It can also be seen in Figure 5 that, with very low evolutionary rates, many quartets cannot be satisfactorily resolved (we get equalities and thus star topologies), leading to poor resolution and many false negatives.

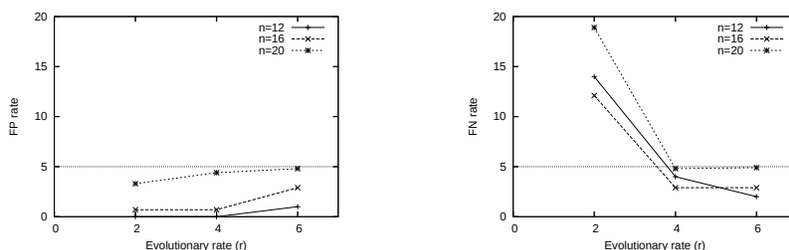


Fig. 5. Performance of our method on genomes of 80 genes.

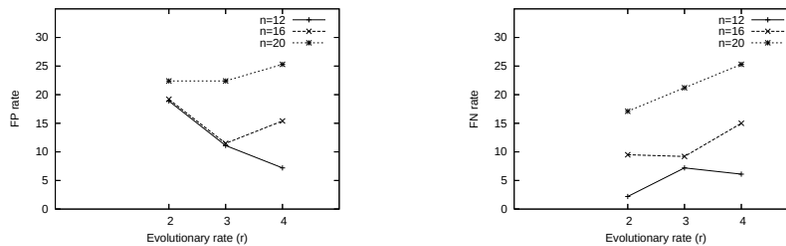


Fig. 6. Performance of our method on genomes of 40 genes.

Our tests verified that a small subset of quartets suffices to infer the complete set of quartets. For datasets with 12 genomes, only 75 quartets (15% of the total) are needed; with 20 genomes, only 270 quartets (6%) are needed. Our selection rule worked well: of the quartets selected, fewer than 2% overall were found to be incompatible. Interestingly, the set of resolved quartets produced by our method produced very accurate reconstructions, while the set produced directly by the relaxed four-point method gave very poor results.

We compared the results obtained by our method with those obtained by simply running the (very fast) neighbor-joining (NJ) method on breakpoint and inversion distance matrices (computed by GRAPPA) for each dataset. For 80-gene genomes, the Robinson-Foulds rate (the average of FP and FN rates) for NJ varied from 20% (for $r = 2$) down to 2–5% (for $r = 6$, with lower rates for 12 genomes and larger rates for 20 genomes), compared to a maximum of 10% (for $r = 2$) down to 1.5–4.5% (for $r = 6$) for our method. For 40-gene genomes, as we observed, our method suffers from saturation effects with 16 or 20 genomes, where its error rate roughly matches that of NJ (10–20%); for 12 genomes, where saturation is less of a problem, our method again easily surpasses NJ, with a median error rate of 8.5% compared to NJ’s rate of 14%.

6 Conclusions

We have presented a quartet-based phylogeny reconstruction method for gene-order data and reported its performance on simulated datasets. Our method produces accurate topologies for trees with up to 25 leaves in reasonable time when the datasets do not exhibit significant saturation. The results we have obtained promise well, especially because we have many possible avenues of improvement. For instance, we have recently developed a linear-programming method that can accurately estimate the edge lengths of fairly small trees [28]; by using this method to estimate the length of quartet edges, we can further improve our quartet selection, in terms of both speed and accuracy. Such improvement should also enable us to handle datasets with larger pairwise distances. We de-

signed this method to extend the range of base methods that can be used in conjunction with a disk-covering method: thus the limitation to sets of 20–30 taxa is not an issue, but in fact a potentially significant gain over the direct use of GRAPPA as a base method, since this last is limited to 12–15 taxa.

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