Annotation-free delineation of prokaryotic homology groups

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Abstract

Phylogenomic studies of prokaryotic taxa often assume conserved marker genes are homologous across their length. However, processes such as horizontal gene transfer or gene duplication and loss may disrupt this homology by recombining only parts of genes, causing gene fission or fusion. We show using simulation that it is necessary to delineate homology groups in a set of bacterial genomes without relying on gene annotations to define the boundaries of homologous regions. To solve this problem, we have developed a graph-based algorithm to partition a set of bacterial genomes into Maximal Homologous Groups of sequences (MHGs) where each MHG is a maximal set of maximum-length sequences which are homologous across the entire sequence alignment. We applied our algorithm to a dataset of 19 Enterobacteriaceae species and found that MHGs cover much greater proportions of genomes than markers and, relatedly, are less biased in terms of the functions of the genes they cover. We zoomed in on the correlation between each individual marker and their overlapping MHGs, and show that few phylogenetic splits supported by the markers are supported by the MHGs while many marker-supported splits are contradicted by the MHGs. A comparison of the species tree inferred from marker genes with the species tree inferred from MHGs suggests that the increased bias and lack of genome coverage by markers causes incorrect inferences as to the overall relationship between bacterial taxa.

Author summary

Assuming genes to be the basic evolutionary unit has been commonplace in bacterial genomics. For example, when quantifying the extent of horizontal gene transfer it is common to infer gene trees and reconcile them against a species tree to account for recombination-based processes. We have developed a new method which challenges this assumption by identifying contiguous regions of true homology without regards to gene boundaries and applied it to Enterobacteriaceae, a family of bacteria containing several important human pathogens. Our results show that genes are composed of distinct homologous regions with conflicting phylogenetic histories. We further demonstrate that failing to take account of this conflict, together with the functional biases we show exist...
among single-copy marker genes, significantly changes the consensus evolutionary tree of Enterobacteriaceae.

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Introduction

Studying prokaryote evolution through the lens of comparative genomics has been conducted using one set of approaches at deep timescales and another set at shallow timescales. At deep timescales where genes and genomes are highly diverged, copies of marker genes which tend to be conserved across many species are identified in the genomes being studied. A marker gene is defined as a protein-coding gene which ideally occurs as a single-copy gene in every studied genome. Multiple sequence alignments (MSAs) of each marker gene may be analyzed individually [1], or concatenated into a supermatrix and be analyzed together [2].

While early phylogenetic studies relied on one or two marker genes such as SSU rRNA [3], the advance of sequencing technologies including next generation sequencing and more recently single molecule real time sequencing has enabled the transition to phylogenomics and the construction of datasets with hundreds of genes from entire genome assemblies [1, 2, 4]. Despite the availability of whole genomes, it remains the default assumption to work with functional genes as the basic evolutionary unit for studying evolution [5]; in a typical phylogenomic analysis where a species evolutionary history is inferred from the genealogies of individual loci, a central mathematical assumption is that the evolutionary history of each locus is recombination free.

However, when functional genes are used as loci, this ignores the possibility that recombination may break up the phylogenetic history of different segments within a gene. This problem is well known with regards to eukaryotes, where meiotic recombination breaks up these histories with each generation. In that case, functional genes are known as “m-genes” and maximal segments which share a common phylogenetic history are known as coalescence genes or “c-genes” [6, 7]. These c-genes are defined as “a segment of the genome for which there has been no recombination over the phylogenetic history of a clade” [6], and it has been previously shown that the species phylogeny can be incorrectly inferred when concatenating multiple c-genes [8]. Identifying c-genes can be very challenging [9], which has led to the common phylogenomic practice of using loci that are individually short and are far apart, even if this practice does not guarantee that the loci constitute c-genes. When the genomes or genomic regions under analysis are collinear and alignable, methods such as coalescent hidden Markov models can be used to delineate c-genes [10], though these methods suffer from scalability issues [11].

While prokaryotes do not undergo meiotic recombination, they experience many other forms of recombination such as horizontal gene transfer and gene duplication/loss. These events are often non-homologous, disrupting even looser assumption of homology along the entire length of functionally equivalent genes. It has been shown that genomic regions with different lengths can be impacted, from a single nucleotide [12], multiple nucleotides [13, 14], a gene [15, 16], an operon [17–20], to even an entire chromosome [21]. Therefore, it is important both not to treat a gene as an evolutionary unit, and to identify homology groups without...
relying on marker gene annotations (homology groups were referred to as “module families” in [22]).

To understand the potential for erroneous phylogenomic inferences when “m-genes” are used instead of “c-genes” for inferring prokaryotic phylogenies, we conducted a simulation study where markers were simulated from multiple c-genes differing slightly along their length. Our results will show that this leads to extremely high bootstrap support for incorrect clades. Work on this problem has been done, but is based on proteomes derived from genome annotations rather than whole genomes. Both the methods of Wu et al. [22] and Leonard and Richards [23] are based on processing BLASTp queries to identify MHG boundaries within protein sequences, and will truncate MHGs that extend past coding regions, and may be inaccurate where errors exist in genome annotations, or where mutations are present that disrupt the coding sequences such as frameshifts or changes to stop codons.

Whole genome aligners may be used at shallow timescales to align as many homologous pairs of nucleotides as possible across input genomes. This approach reveals the relationship between different strains within a species or species complex, and the forces of molecular evolution shaping their genomes [24, 25]. While much more of the input genomes are used in this approach compared with marker genes, due to the high degree of divergence between bacteria species, state-of-the-art whole genome aligners all have difficulties at deeper timescales. For example, Parsnp [26] aligns only core genomes defined as orthologous sequences conserved by all input genomes. ProgressiveMauve [27] is extended from mauve [28] utilizing k-mer based locally collinear blocks, which is significantly slower and consumes more space when faced with highly divergent genomes. Progressive Cactus [29] requires a guide tree and performs well when aligning different samples of the same species, however its performance is questionable when aligning bacterial genomes of different species, which will be much more divergent than genomes of different primates or other mammals. SibeliaZ [30] is a de Bruijn based whole genome aligner that is faster than Progressive Cactus but has an even smaller tolerance of divergent input genomes.

Here we introduce a method that identifies groups of sequences that share common ancestry along their entire length. It is based on constructing an graph of intervals connecting segments of different genomes which have evidence for local homology at the nucleotide level, initializing each segment as a “MHG,” and walking the graph to merge and partition MHGs. The end result is a set of MHGs where each MHG contains a set of homologous sequence segments. Sequences assigned to each MHG using our method may be orthologous or paralogous, in the latter case due to hidden paralogy and because we incorporate every sequence identified as homologous from each genome, which necessarily includes gene duplications. Gene losses may result in zero sequences being included from a given genome.

Compared with the marker gene approach for deep timescales, our method has the advantage of using more of the input genomes when the divergence is low enough for sequence homology to be identifiable at a nucleotide level. The selection of marker genes is biased towards slow evolving genes, since genes with frequent point mutations will be difficult to align, and genes with structural variations should not be aligned as a single unit. These slow evolving markers may not be representative of the whole genome in terms of their structure and function, and we report evidence for this bias in our study and show how our MHGs compiled by our method by contrast are more broadly representative.

Another advantage over marker gene studies is that our algorithm does not rely on any existing genome annotation to define the boundaries of the sequences (e.g., by referring to previously inferred boundaries of domains, coding sequences or operons). This compares with a previously published method for compiling MHGs which takes protein sequences as input instead of a genome assembly [22]. That method was also targeted at Drosophila, a genus of
eukaryotes, where evolution is much slower and hence the difficulty of the problem is easier than for prokaryotes.

To analyze the performance of our method, we applied it to a select subset of Enterobacteriaceae genomes. We found that our algorithm is able to cover most of the genes with MHGs for free-living bacteria other than the presumptive outgroup taxon, especially compared with the limited number of marker genes. A further analysis of gene ontology (GO) terms associated with marker and MHG-covered genes showed that MHGs are less biased in their associated functions than marker genes are. Finally, a phylogenetic analysis revealed concerning levels of conflict between the marker genes and associated MHGs, suggesting that marker genes are an inappropriate unit for phylogenomic studies and may infer false evolutionary relationships. Our new method for the preparation of more reliable MHG-based collation of phylogenomic data is available at https://github.com/NakhlehLab/Maximal-Homologous-Groups.

Materials and methods

Algorithm for delineating maximal homologous groups of sequences

Given a set of genomes, we define a maximal homologous group of sequences (MHG) as a maximal set of maximum-length sequences (substrings of the genomes) whose evolutionary history is a single tree and involves no rearrangements. The set is maximal in that adding more sequences to it violates the single-tree assumption and the sequences are of maximum lengths in that including more nucleotides in them also violates the single-tree assumption (e.g., having fused genomic regions). MHGs are defined for any set of two or more genomes, with no regard to functional annotations.

A high-level description of the algorithm is described below and illustrated in Fig 1. A full implementation of the algorithm is available at https://github.com/NakhlehLab/Maximal-Homologous-Groups.

Our algorithm uses graphs in two key ways. Firstly, the alignment graph is used to store the connections between genome segments identified using BLASTn in an efficient way, and since segments from different connected components of the graph will never be merged into the same MHGs, the partition and merge step may be parallelized by connected component, or flushed to storage after each connected component is traversed to reduce memory usage. Secondly, MHGs are stored as graphs in order to propagate boundaries between genomes while taking into account indels, as described below. Starting from a set of genomes:

1. **All-versus-all BLASTn to find pairwise local alignments** (Fig 1B) All versus all BLASTn queries are filtered based on the bit score values [1]. For each genome, the maximum possible bit score is calculated from the set of results obtained by querying the genome against itself using BLASTn. All versus all BLASTn results are subsequently excluded if their bit score falls below a certain threshold of the maximum bit score obtained for the genome the query sequence derived from.

2. **Sequence pile-up** (Fig 1C) Given every pairwise alignment call by BLASTn, the algorithm piles up alignments that cover overlapping regions on the same genome.

3. **Initial MHGs and alignment graph construction** (Fig 1D and 1E) For piled-up alignments in every genome, the algorithm takes the union of overlapping alignments resulting in a set of non-overlapping ranges in all genomes. An alignment graph is constructed from adding the range set as nodes, and an edge is added for each local alignment to connect the two nodes that the subject and query ranges of the local alignment correspond to
There could be multiple edges between two nodes. The alignment graph serves to efficiently split up a big problem to a few subproblems where each connected component in the alignment graph can be traversed independently. In other words, A MHG can only include subsequences of nodes from the same connected component. Encountering a
new pairwise alignment mapping, the algorithm only needs to check and partition MHGs generated from the same connected component instead of searching around full set of MHGs.

4. **Alignment graph traversal** (Fig 1F) Starting from the the alignment graph, every edge is visited using a depth-first search (DFS) of every connected component. Whenever a new breakpoint is inserted, MHGs containing segments which include that breakpoint will be split into two at the breakpoint, and pairs of post-split MHGs connected by the local alignment edge will be merged. For example, when visiting edge A:B:1 in Fig 1F, encountering a first breakpoint carried by B1, MHG A1 is partitioned into A11, A12, and A11 is merged with B1 because edge A:B:1 connects A11 and B1. The split and merged MHGs will contain directions for all sequences based on the relative directions of the subject and query sequences in the local alignment (i.e. in the same direction on both genome assemblies, or opposite directions, relative to the strand represented by those assemblies). And after visiting all edges in the alignment graph, the output MHGs will contain the homology groups for the input whole genomes.

When a new breakpoint is added to an existing MHG after processing a local alignment, we use the pairwise local alignment estimated by BLASTn to ensure the new breakpoint is inserted at accurate locations. Our algorithm converts the pairwise local alignments to two bit arrays where 1 means a match, and 0 means a gap. We first locate the correct location of the breakpoint in a pairwise alignment by counting 1s from the beginning of the bit array for the source genome (for which already know the breakpoint location), to match the difference between the breakpoint coordinate in the source genome and the starting coordinate of that local alignment. Second, for the destination genome (where the breakpoint location is unknown), we count 1s in its bit array, going backwards from the breakpoint location (which is the same as identified in the source bit array) until we reach the beginning of the bit array. This then gives us the difference in position between the starting coordinate of the local alignment in the destination genome and the breakpoint. The first stage, all-vs-all BLAST search has a time complexity $O(n^2)$ where $n$ is the number of genomes, since every genome is queried by every genome. In a hypothetical scenario where all genes are present as single-copy orthologs in every genome, the alignment graph to be walked in the second stage will have $O(n^2)$ edges. Since every edge is traversed once, this stage also has a time complexity of $O(n^2)$. When an edge is visited that leads to an ortholog from a new species being added to an existing MHG, this genome will be segmented at that locus and the ortholog incorporated into the MHG through backtracking. A new R-tree [31], used to store the segmentation of each genome, will need to be constructed which in the worst case takes $O(s \cdot \log(s))$ time where $s$ is the number of segments, which will be proportional to the number of single-copy orthologs. The backtracking process is of time complexity $O(m)$ where $m$ is the number of sequences in the MHG, which will grow with $n$. However, as $n$ increases, an increasing proportion of edges will not lead to the incorporation of new sequences, hence R-tree construction and backtracking can be discounted.

For any edge visited, the R-trees of the source and destination genomes are queried. The time complexity of these queries is bounded by $O(\log(s))$ and $O(s)$, but since this will not depend on the number of genomes in this hypothetical scenario, the overall time complexity relative to the number of genomes is $O(n^2)$. Under more realistic conditions, the time complexity becomes much more difficult to analyze as the number of segments in each genome will depend on both the number of genomes and multifarious evolutionary processes.
A prokaryotic dataset

We applied our method to a set of 19 bacterial species, each from a different genus of $\gamma$-proteobacteria; 18 from within Enterobacteriaceae and one outgroup. We chose the specific genome assemblies from a previously compiled dataset of 10,575 bacterial and archaeal genomes [2]. The assembly we used for the outgroup taxon has the RefSeq identifier GCF_001887595.1, which in the analysis of Zhu et al. [2] was mislabelled as Klebsiella michiganensis when it actually is Luteibacter rhizovicinus [32]. The genome assembly GCF_001022135.1 that we used for Phytobacter ursingii was originally identified as Kluyvera intermedia when it was first published but has since been reclassified [33, 34]. The species names and NCBI reference IDs used in our analysis are: Blochmannia endosymbiont (GCF_000973545.1), Candidatus Doolittlea endobia (GCF_900039485.1), Candidatus Gullanella endobia (GCF_900048035.1), Candidatus Hoaglandella endobia (GCF_900044015.1), Candidatus Ishikawaella capsulata (GCF_000828515.1), Candidatus Risia pediculicola (GCF_000299455.1), Candidatus Tachikawaea gelatinosa (GCF_000828815.1), Cedecea neteri (GCF_000757815.1), Citrobacter freundii (GCF_000648515.1), Cronobacter sakazakii (GCF_000982825.1, Enterobacter lignolyticus SCF1 (GCF_000164865.1), Escherichia coli (GCF_000299455.1), Luteibacter rhizovicinus (GCF_001887595.1), Phytobacter ursingii (GCF_001022135.1), Kosakonia sacchari (GCF_000300455.3), Pluralibacter gergoviae (GCF_000757785.1), Salmonella enterica (GCF_000195995.1), Shigella flexneri (GCF_000006925.2), and Shimwellia blattae (GCF_000262305.1).

A marker gene is defined in an ideal sense as a single copy gene which occurs in every studied genome. In practice this requirement is too strict to construct a useful data-set with, and is relaxed from “every” to “most” genomes. In this paper, marker genes were previously identified as protein-coding genes [2], and we used the best tBLASTn [35] hit from the proteome translated from the corresponding genome to identify the marker gene for that species, but only where it was identified as present in that species in the original study. Reference IDs for each marker gene from the original publication were used to retrieve the query protein sequences from UniProt [36]. And in order to calculate whether a marker gene is covered by any MHG, we used the best hit from tBLASTn to locate the nucleotide sequence in each genome corresponding to every known marker gene protein sequence. Marker coverage of each genome and its genes was calculated based on the full lengths of the specific genes identified in that genome, as above.

We constructed a nucleotide BLAST database from all 19 genomes to conduct the all verses all BLASTn search. For the search, we used a word size of 9, a gap open penalty of -1, and a gap extend penalty of -1. These parameter values were chosen so that our method will work even for very divergent genomes. A shorter word size than the default enables BLASTn to align regions sharing shorter exactly matched k-mers, and choosing the smallest possible mismatch and gap penalties should maximize the lengths of locally homologous alignments identified by BLASTn.

Next, we used the algorithm described above to infer the set of MHGs for those 19 genomes. Coverage of genomes and genes by MHGs was computed using BEDtools [31]. Since MHGs are non-overlapping, base pair coverage refers to the proportion of a genomic sequences covered by any MHG.

Simulation of marker genes

To simulate marker genes with internal recombination we began with the species phylogeny previously inferred by concatenating marker genes [2] which was pruned to contain the above 19 bacteria. For each of 10 replicate marker genes, the first component tree was obtained by
performing a single nearest-neighbor interchange (NNI) operation on the initial tree, the second component tree was obtained by performing a single NNI operation on the first component tree, and so on. We chose NNI as a reasonable and conservative facsimile of horizontal recombination, since it will only exchange segments between closely related taxa.

We simulated 15 such component trees, generated 100bp sequence alignments with equal transition rates and base frequencies using Seq-Gen [37] on each component tree, and concatenated them to produce the marker MSA for a total length of 1,500bp, chosen because it is close to the average length of marker genes included in our study, which we calculated as being 1,503bp. The choice of 100bp was made to ensure phylogenetic histories of each segment were to an extent still resolvable while simulating a substantial number of recombination events within each gene, and equal rates and frequencies used since this study does not focus on the effects of varying substitution models or parameter values.

IQ-TREE [38] was used to infer maximum likelihood trees with bootstrap support for the 100bp non-recombining segments and the simulated markers with internal recombination. We pre-specified the HKY substitution model for IQ-TREE, of which the true model is a special case. Robinson–Foulds (RF) distances between known and inferred trees were calculated using phangorn [39].

**Phylogenetic analysis**

For each marker and for each MHG containing at least 4 segments, we ran the FFT-NS-i [40] algorithm implemented in MAFFT with 500 maximum iterations. For each resulting MSA, IQ-TREE [38] was run with default settings to build a gene tree. Finally we used ASTRAL-Pro [41] to infer a species tree from the marker gene trees, and another species tree from the MHG gene trees, because this method accounts for gene duplication and our MHGs may contain more than one sequence per genome, and it has been shown to be robust to duplication and loss rates [42].

Because methods for quantifying phylogeny conflict are based on comparing trees with uniquely labeled tips, we excluded MHGs with more than one sequence derived from any single genome. For the nucleotide sequences of marker genes and overlapping MHGs with at least four taxa, an original tree and 100 bootstrap trees are inferred using the IQ-TREE maximum-likelihood method [38]. MHGs with an internal branch length less than 0.0005 in their original trees were regarded as unresolved and filtered out. This threshold value is smaller than the reciprocal of the average marker gene length of 1,503bp, so branch lengths below this threshold imply no mutations occurred along this branch (at least under a Jukes–Cantor model) and the relationship between the four subtrees adjacent to this branch is impossible to resolve. After removing ambiguities, every bootstrap tree in every marker and MHG will yield a set of bipartition splits for each internal node. We used DendroPy [43] to calculate bipartitions, which DendroPy encodes as bitmasks. Since a marker and its overlapping MHGs may consist a different set of taxa, a bitwise normalization operation is performed to ensure each bipartition is represented by the same bitmask, and by exactly one bitmask, for the marker and all overlapping MHGs.

**Functional annotation**

To classify genes—either marker genes, or a gene associated with a MHG because they are overlapping—by function, we used a GO slim. This aggregates gene classifications into a small enough set of categories to be interpretable and reveal any bias in the distribution of gene function. We used a custom script to traverse the GO hierarchy, and categorized genes based on whether any of their GO terms had an “is_a” or “part_of” relationship to any one of the
Alliance for Genome Resources (ARG) GO slim terms. We allowed multiple categories per gene when their GO terms mapped to multiple ARG slim terms.

**Results**

**Simulation study**

To understand whether it is better to use marker genes or MHGs as the basic unit of evolutionary inference, we inferred maximum likelihood trees from simulated marker genes, their non-recombining segments, along with bootstrap trees for both. The typical bootstrap support for branches inferred from simulated markers was much higher than component trees (Fig 2A and 2D) due to the longer length and higher information content. However this is deeply misleading, as the normalized Robinson-Foulds (RF) distance between the inferred marker gene tree and the true component trees is centered around 0.5, meaning about half the branches are incorrect (Fig 2C). This is worse than the accuracy of the trees inferred individually for each non-recombining component, which was typically below 0.5 (Fig 2F). For the non-recombining segments, bootstrap support for true branches was higher than incorrect branches, suggesting that bootstrap support values for inferred MHG trees can be used to identify true splits in the local phylogeny, as opposed to the clonal frame (Fig 2B and 2E).

**MHG occurrence across genomes**

The dataset we analyzed consists of both free-living and endosymbiotic bacteria. Severe gene loss is common among endosymbionts, and as a result relatively few MHGs contain sequences from those species (Fig 3). In addition to the explicitly labeled Blochmannia endosymbiont, these species include all candidati.

In terms of the coverage of genomes by MHGs at a nucleotide level, a substantial fraction of genes were covered by MHGs, even when we limited the MHGs to those with sequences from at least eight species (Fig 4). The difference between gene length and genome size was roughly

![Fig 2. Support for branches in simulated marker genes and component trees.](https://doi.org/10.1371/journal.pcbi.1010216.g002)
proportional across genomes, and the gap between the two corresponds to the amount of intergenic sequence in each genome. Unlike eukaryotes, prokaryotes have diminutive genomes with consistently high gene density [46, 47].

Intergenic coverage was also good for free living bacteria other than *Luteibacter rhizovicinus*, but not for endosymbionts, indicating that endosymbionts have undergone substantial disruption to their genomes between as well as within genes. Because of the gene loss among endosymbionts, many genes in free living bacteria bear no homology to genes in endosymbionts, and therefore relatively few genes in free living bacteria are covered by MHGs with sequences from 16 or more species (Fig 4).

Fewer MHGs contained sequences from *L. rhizovicinus*, although the proportion of genic sequences covered by MHGs converged with other taxa for MHGs with more genomes included (Figs 3 and 4). By contrast, there was almost no intergenic coverage, even for MHGs...
with 2 or more genomes represented. This is due to the fact that *L. rhizovicinus* is highly divergent from the other genomes in our analysis, being from an entirely different family (Rhodanobacteraceae) within \(\gamma\)-proteobacteria. Nonetheless, the substantial coverage of genic sequences demonstrates the suitability of using MHGs for interfamilial analysis.

**Escherichia coli** gene functionality analysis

In order to explore whether marker genes or genes which overlap MHGs are biased towards specific functions, we analyzed the functional categorization of both those kinds of genes in *Escherichia coli*. We observed substantial over and under-representation among many gene categories by the set of marker genes (Fig 5). The three categories of gene function most over-represented among marker genes were catalytic activity, small molecule binding, and carbohydrate derivative binding. The three categories most under-represented were plasma membrane, establishment of localization, and transporter activity. While genes overlapping highly conserved MHGs (those with sequences from 16 or more species) saw similar biases, the representation of all categories was mostly unbiased among fuller subsets of MHGs (Fig 5).

**Conflict between markers and MHGs**

MHGs inferred using our algorithm should be homologous across the entire length of every sequence in the MHG, indels excepted. In contrast, marker genes may have undergone rearrangements. Our method inferred multiple MHGs for each marker gene (S1 Appendix) and since our simulation study showed that conflicting phylogenetic histories within a gene can...
produce misleadingly high bootstrap support values, we compared the support values of splits from each MHG overlapping a marker gene with the support value of that split from the corresponding marker gene.

MHGs with fewer than four sequences were excluded because they contain no splits other than the tip branches, which are uninformative. Additionally, MHGs with more than one sequence for a single genome were excluded as this would make the analysis of support for phylogenetic splits intractable. Only 20.6% of MHGs with four or more sequences were excluded on that basis, and the remaining 79.4% of MHGs we term “single-locus informative MHGs.”

For each marker-overlapping MHG, we considered the full set of splits observed in the entire bootstrap distribution for that MHG and the corresponding marker gene. Since the number of possible splits grows exponentially with the number of taxa, but only a few will be well-supported by the actual sequence alignments, a very long tail of splits with low MHG and/or marker gene support results (Fig 6). More interesting is that strong split support by marker gene alignments is not a good predictor of strong support by MHGs. When considering the marker genes with greater than 90% support, there appears to be almost no correlation with MHG support across the entire range of MHG support values (Fig 6). This is consistent with the aforementioned simulation study findings, further supporting the notion that high bootstrap support values derived from marker gene alignments may be misleading.

We also quantified phylogenetic (in)congruence using the extended quadripartition inter-node certainty (EQP-IC) metric [48]. Comparing to other statistics, EQP-IC aims at measuring the correctness of a given branch. It is robust when there are errors in the input reference tree.
And in our case, EQP-IC values were calculated for each marker gene tree split, based on support by overlapping MHG gene trees. The modal value was around zero, indicating either a lack of information or a balanced mix of phylogenetic congruence and incongruence (Fig 7). However, given our observations of the number of highly supported splits, a lack of information is more likely (Fig 6).

Fig 6. Conflicting support for splits between markers and MHGs. Bootstrapping was used to compute support. Each unique split from the MHG and marker gene tree bootstrap distributions, from each pair of MHG and corresponding marker gene, was counted once towards the frequency.

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Fig 7. Extended quadripartition internode certainty (EQP-IC) of MHGs overlapping marker genes. Values closer to 1 indicates concordant phylogenetic histories between markers and MHGs, values closer to -1 indicate discordant histories, and values around 0 indicate either a lack of information, or a mix of concordance and discordance.

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While very few marker splits were well supported by their overlapping MHGs according to this metric, a substantial number of splits were highly contradicted by those MHGs, with a spike in EQP-IC values observed around -1 (Fig 7). Values close to -1 indicate strong phylogenetic incongruence.

**Species tree comparison**

There are many approaches to problems in comparative genomics, but a central concept that many other tools rely on, and which is used to explain and communicate the relationship among a set of species, is the species phylogeny. To understand whether the lack of coverage of genomes by marker genes, observable bias in functionality, and phylogenetic conflict with associated MHGs would impact the species phylogeny estimated from marker genes, we compared a species tree inferred from marker gene trees with a species tree inferred from MHG gene trees. For this analysis, we used all markers to build the marker species tree, but only used MHGs ≥ 400bp to build the MHG species tree. This was to exclude the effects of random error in shorter MHGs, as we found the distribution of bootstrap support for the branches of MHG gene trees ≥ 400bp was similar to markers, but the bootstrap support for the branches of more inclusive sets of MHG gene trees was generally lower (Fig 8). Again only MHGs with at least four taxa were included.

The method we used to infer species trees (ASTRAL-Pro) is able to use gene trees with multiple gene copies as input. Unlike markers genes, MHGs are not restricted to single copy sequences, so to understand if this changes the species tree result we ran ASTRAL-Pro with all MHGs ≥ 400bp and also with only single-locus informative MHGs ≥ 400bp, but the same topology was returned for both sets of MHGs (Fig 9).

![Fig 8. Bootstrap support for splits. The cumulative density distribution of bootstrap support is shown across all non-trivial splits from the maximum likelihood trees inferred from the original marker gene sequence alignments (blue). The same distribution is shown for the splits inferred from the original MHG sequence alignments (orange). In addition, the distribution of split bootstrap support from two restricted sets of MHG sequence alignments are shown; those with a length of at least 100bp (green) and those with a length of at least 400bp (red).](https://doi.org/10.1371/journal.pcbi.1010216.g008)
However, the MHG-based species tree topology differed in two ways from the marker-based topology. The more substantial difference is the placement of the endosymbiont *Candidatus Ishikawaella*, which the marker-based analysis groups with *Candidatus Riesia* and *Tachikawaea*. The EQP-IC support from marker genes for this grouping is strongly positive (0.66) indicating most marker genes containing relevant taxa provide support for this clade, but the EQP-IC support from MHGs is the most negative possible value (-1), indicating that virtually all MHGs containing relevant taxa reject it.

In addition, *Pluralibacter gergoviae* and *Enterobacter lignolyticus* are a clade in the marker-based species tree but not in the MHG-based species tree. The difference in EQP-IC support is however minor, with 0.06 support among markers and -0.03 support around MHGs, indicating a lack of signal or consensus for this grouping among both methods (Fig 9).

**Intergenic MHG analysis**

Besides the potential bias introduced by an unrepresentative distribution of gene function, it is known that gene and intergenic sequences are subject to different evolutionary processes. For example, the rate of point mutations leading to non-synonymous amino acid changes may be substantially higher or lower than the rate for synonymous changes, depending on whether loci are under predominantly positive or negative selection [49]. This does not apply to intergenic sequences which definitionally do not encode for proteins even if they are functionally important. Marker genes do not include intergenic sequences, whereas our method does not exclude them, so we studied the possible effects of their inclusion.

We inferred species trees with EQP-IC support values from two subsets of MHGs. The first subset was restricted to MHGs which only contained sequences from inside genes, and the second subset was restricted to MHGs which only contained intergenic sequences. Because
homology is more difficult to identify between genomes at intergenic loci we reduced the MHG length threshold for this analysis from 400bp to 100bp. Still, the outgroup taxon *L. rhizovicinus* and all but one of the endosymbionts had no representation within intergenic-only MHGs. Furthermore, the remaining endosymbiont *Candidatus Doolittlea* was only included in two intergenic-only MHGs. For this reason, we excluded the outgroup and endosymbionts when estimating the intergenic-only MHG species tree and EQP-IC values.

After reducing the length threshold and excluding the two intergenic-only MHGs with *Candidatus Doolittlea* sequences, there were 3485 MHGs included in the first subset and 172 in the second, a similar ratio to the overall ratio of gene to intergenic genome content. The difference in topology between the gene and intergenic trees was an exchange of the *Phytobacter* and *Kosakonia* clade with the *Enterobacter* and *Pluralibacter* clade (Fig 10). This a “nearest-neighbor” interchange and as such is a minor difference. The support values for either arrangement is close to zero, indicating this change is likely sampling noise than a genuine difference between gene and intergenic evolutionary histories (Fig 10).

Reducing the length threshold appears to have reduced EQP-IC support, presumably because shorter MHGs have a lower information content and there is greater conflict between their splits. For either ≥100bp tree, the only non-endosymbiont clade supported by EQP-IC values was the *Escherichia* and *Shigella* sister relationship (Fig 10). Furthermore, empirically sound rooting of the intergenic-only tree was impossible since the outgroup was excluded, so we choose the root that best matches the gene-only MHG species tree, and this rooting should not be considered reliable.

### Discussion

The rapid advance in throughput and availability of sequencing technology has led to a deluge of genomic data. For bacteria, more than 430,000 assembled genomes were available as of April 2020 [50]. Studying the history of bacterial genome evolution offers an understanding of both the history of individual genes and operons, as well as broader evolutionary processes [51–54]. A medically significant implication is the acquisition of antibiotic resistance, both *de novo* and from other species [55, 56].

The complexity of genome evolution in bacteria is due to rampant horizontal gene transfer [57–60], gene duplication and loss [61], and *de novo* sequence evolution. Existing methods of
whole genome alignment work well at shallow timescales, but methods for studying evolution at deeper timescales in bacteria are hindered by this complexity, and rely on the assumption of non-recombination within marker genes. We have shown that this assumption, when broken, is not detectable or mitigated by low support values for branches affected by recombination. Not only that, marker gene methods rely on the accuracy of existing genome annotations. For newly sequenced genomes, annotations may not even be available [62].

By applying our method to 19 bacteria from a diversified sampling of Enterobacteriaceae, we have demonstrated its practical utility and more advantages over marker genes. Our method uses more sites from each genome, and is less biased in the functions of the corresponding genes that are included. Given a genome that is highly divergent from the rest, a family-level difference can be accurately captured by MHG. Instead of being limited by the constraint of high conservation as a marker gene, an MHG-based analysis is able to properly classify homologous relationships from any subset of the input genomes. We have shown that many highly supported splits from inferred marker trees are not supported by corresponding MHGs, concordant with our expectations from our simulation study where components have different phylogenetic histories. The EQP-IC measure provides evidence for where intragenic conflict affects the support for various taxonomic hypotheses, and for which clades are robustly inferred regardless of this conflict.

Of particular interest is our findings on the evolution of Enterobacteriaceae endosymbionts. Both marker and MHG phylogenies strongly support the monophyly of this group, in contrast to a previous concatenation analysis which claimed it has multiple origins [63]. These vertically transmitted endosymbionts provide essential amino acids to their insect hosts, and transmission may occur externally after oviposition [64, 65]. Neither marker nor MHG phylogeny is compatible with pure vertical transmission however, with the mealybug endosymbionts Candidatus Hoaglandella, Doolittlea and Gullanella being paraphyletic or polyphyletic. Based on the placement of those taxa, we hypothesize that this symbiotic relationship originated in mealybugs before diversifying to other insects. The major difference between phylogenies is that the MHG phylogeny is incompatible with this diversification occurring within a single lineage, as the EQP-IC measure strongly rejects a clade containing the four non-mealybug endosymbionts.

While the alignment graph enables some parallelization and space efficiency, improved scalability will be an important future direction for this algorithm and our implementation. The bottleneck of the current MHG tool is the polynomial (under ideal conditions) growth in runtime with the number of genomes. One avenue for alleviating this is replacing the all-vs-all BLAST with a subset of pairwise (genome by genome) BLAST queries, since MHGs are transitive in nature, complete connectivity between genomes is not required. This could use a phylogenetic guide tree, although rigorous study is needed to determine the best approach of choosing the subset of pairwise queries to minimize any loss of sensitivity. As with many algorithms, a divide-and-conquer strategy may also improve walltime performance, although not overall CPU time or power consumption. Beyond scalability, due to the stochasticity and heuristic nature of BLASTn, using the boundaries of BLASTn queries to define MHG boundaries may cause inferred MHGs to be shorter than necessary. This may be addressed in the future through post-processing, or by reconciling BLASTn query boundaries before or simultaneously with the partition and merge step.

Our work here demonstrates the fundamental problems of existing methods for prokaryotic phylogenetics at deeper timescales. Not only is it immediately useful on smaller, but still whole-genome, datasets, it provides the starting point for improved and scalable phylogenomic algorithms and implementations that are coherent with the actual complexity of the evolutionary histories across bacterial genomes.
Supporting information

S1 Appendix. Maximal homology group overlap of marker genes. Each marker gene is labeled with the UniProt accession number of its reference protein. MHG sequences are drawn as colored arrows overlapping the sequences from each species for each marker gene. MHG sequences drawn with the same color belong to the same MHG, although colors are repeated when more than 20 MHGs overlap a marker gene. Arrowheads indicate the relative orientation of MHG sequences to show inversions.

(Accessed via PDF link)

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References

1. Lerat E, Daubin V, Moran NA, Penny D. From gene trees to organismal phylogeny in prokaryotes: the case of the γ-Proteobacteria. PLoS biology. 2003; 1(1):e19. https://doi.org/10.1371/journal.pbio.0000019 PMID: 12975687
2. Zhu Q, Mai U, Pfeiffer W, Janssen S, Asnicar F, Sanders JG, et al. Phylogenomics of 10,575 genomes reveals evolutionary proximity between domains Bacteria and Archaea. Nature communications. 2019; 10(1):1–14. https://doi.org/10.1038/s41467-019-13443-4 PMID: 31792218
3. Eisen JA. The RecA protein as a model molecule for molecular systematic studies of bacteria: Comparison of trees of RecAs and 16S rRNAs from the same species. Journal of Molecular Evolution. 1995; 41(6):1105–1123. https://doi.org/10.1007/BF00173192 PMID: 8587109
4. Williams KP, Gillespie JJ, Sobral BWS, Nordberg EK, Snyder EE, Shallom JM, et al. Phylogeny of Gammaproteobacteria. Journal of Bacteriology. 2010; 192(9):2305–2314. https://doi.org/10.1128/JB.01480-09 PMID: 20207755
5. Eisen JA, Fraser CM. Phylogenomics: intersection of evolution and genomics. Science (New York, NY). 2003; 300(5626):1706–7. https://doi.org/10.1126/science.1086292 PMID: 12805538
6. Springer MS, Gatesy J. The gene tree delusion. Molecular phylogenetics and evolution. 2016; 94:1–33. https://doi.org/10.1016/j.ympev.2015.07.018 PMID: 26238460

7. Doyle JJ. Defining coalescent genes: theory meets practice in organelle phylogenomics. Systematic Biology. 2021.

8. Kubatko LS, Degnan JH. Inconsistency of phylogenetic estimates from concatenated data under coalescence. Systematic biology. 2007; 56(1):17–24. https://doi.org/10.1080/10635150601146041 PMID: 17366134

9. Springer MS, Gatesy J. Delimiting coalescence genes (c-genes) in phylogenomic data sets. Genes. 2018; 9(3):123. https://doi.org/10.3390/gene9030123

10. Hoboth A, Christensen OF, Mailund T, Schierup MH. Genomic relationships and speciation times of human, chimpanzee, and gorilla inferred from a coalescent hidden Markov model. PLoS genetics. 2007; 3(2):e7. https://doi.org/10.1371/journal.pgen.0030007 PMID: 17319744

11. Liu X, Ogilvie HA, Nakleh L. Variational inference using approximate likelihood under the coalescent with recombination. Genome Research. 2021; 31(11):2107–2119. https://doi.org/10.1101/gr.273631.120 PMID: 34426513

12. Hommais F, Pereira S, Acquaviva C, Escobar-Parâmo P, Denamur E. Single-nucleotide polymorphism phylotyping of Escherichia coli. Applied and environmental microbiology. 2005; 71(8):4784–92. https://doi.org/10.1128/AEM.71.8.4784-4792.2005 PMID: 16085876

13. Wang Y, Zhang Z. Comparative sequence analyses reveal frequent occurrence of short segments containing an abnormally high number of non-random base variations in bacterial rRNA genes. Microbiology (Reading, England). 2000; 146 (Pt 11):2845–54. https://doi.org/10.1099/00221287-146-11-2845 PMID: 11065363

14. Chan CX, Darling AE, Beiko RG, Ragan MA. Are protein domains modules of lateral genetic transfer? PloS one. 2009; 4(2):e4524. https://doi.org/10.1371/journal.pone.0004524 PMID: 19229333

15. Matic I, Rayssiguier C, Radman M. Interspecies gene exchange in bacteria: the role of SOS and mismatch repair systems in evolution of species. Cell. 1995; 80(3):507–15. https://doi.org/10.1016/0092-8674(95)90501-4 PMID: 7859291

16. Abby SS, Tannier E, Gouy M, Daubin V. Lateral gene transfer as a support for the tree of life. Proceedings of the National Academy of Sciences of the United States of America. 2012; 109(13):4962–4967. https://doi.org/10.1073/pnas.1116871109 PMID: 22416123

17. Yap WH, Zhang Z, Wang Y. Distinct types of rRNA operons exist in the genome of the actinomycete Thermomonospora chromogena and evidence for horizontal transfer of an entire rRNA operon. Journal of bacteriology. 1999; 181(17):5201–9. https://doi.org/10.1128/jb.181.17.5201-5209.1999 PMID: 10464188

18. Igarashi N, Harada J, Nagashima S, Matsuura K, Shimada K, Nagashima KV. Horizontal transfer of the photosynthesis gene cluster and operon rearrangement in purple bacteria. Journal of molecular evolution. 2003; 52(4):333–41. https://doi.org/10.1007/s00239010163 PMID: 11343129

19. Omelchenko MV, Makarova KS, Wolf YI, Rogozin IB, Koonin EV. Evolution of mosaic operons by horizontal gene transfer and gene displacement in situ. Genome biology. 2003; 4(9):R55. https://doi.org/10.1186/gb-2003-4-9-r55 PMID: 12952534

20. Castillo-Ramírez S, Vázquez-Castellanos JF, González V, Cevallos MA. Horizontal gene transfer and diverse functional constraints within a common replication-partitioning system in Alphaproteobacteria: the repABC operon. BMC genomics. 2009; 10:536. https://doi.org/10.1186/1471-2164-10-536 PMID: 19919719

21. Akagi Y, Akamatsu H, Otani H, Kodama M. Horizontal chromosome transfer, a mechanism for the evolution and differentiation of a plant-pathogenic fungus. Eukaryotic cell. 2009; 8(11):1732–8. https://doi.org/10.1128/EC.00135-09 PMID: 19749175

22. Wu YC, Rasmussen MD, Kellis M. Evolution at the subgene level: domain rearrangements in the Drosophila phylogeny. Molecular biology and evolution. 2012; 29(2):689–705. https://doi.org/10.1093/molbev/msr222 PMID: 21900599

23. Leonard G, Richards TA. Genome-scale comparative analysis of gene fusions, gene fissions, and the fungal tree of life. Proceedings of the National Academy of Sciences of the United States of America. 2012; 109(52):21402–7. https://doi.org/10.1073/pnas.1210090110 PMID: 23236161

24. Didelot X, Lawson D, Darling A, Falush D. Inference of Homologous Recombination in Bacteria Using Whole-Genome Sequences. Genetics. 2010; 186(4):1435–1449. https://doi.org/10.1534/genetics.110.120121 PMID: 20923983

25. Sakoparnig T, Field C, van Nimwegen E. Whole genome phylogenies reflect the distributions of recombination rates for many bacterial species. Elife. 2021; 10:e65366. https://doi.org/10.7554/eLife.65366 PMID: 33416498
26. Treangen TJ, Ondov BD, Koren S, Phillippy AM. The Harvest suite for rapid core-genome alignment and visualization of thousands of intraspecific microbial genomes. Genome biology. 2014; 15(11):1–15. https://doi.org/10.1186/s13059-014-0152-4 PMID: 25410596

27. Darling AE, Mau B, Perna NT. progressiveMauve: multiple genome alignment with gene gain, loss and rearrangement. PloS one. 2010; 5(6):e11147. https://doi.org/10.1371/journal.pone.0011147 PMID: 20593022

28. Darling AC, Mau B, Blattner FR, Perna NT. Mauve: multiple alignment of conserved genomic sequence with rearrangements. Genome research. 2004; 14(7):1394–1403. https://doi.org/10.1101/gr.2289704 PMID: 15231754

29. Armstrong J, Hickey G, Diekhans M, Novak AM, Deran A, et al. Progressive Cactus is a multiple-genome aligner for the thousand-genome era. Nature. 2020; 587(7833):246–251. https://doi.org/10.1038/s41586-020-2871-y PMID: 33177663

30. Minkin I, Medvedev P. Scalable multiple whole-genome alignment and locally collinear block construction with SibeliaZ. Nature communications. 2020; 11(1):1–11. https://doi.org/10.1038/s41467-020-19777-8 PMID: 33303762

31. Quinlan AR, Hall IM. BEDTools: a flexible suite of utilities for comparing genomic features. Bioinformatics. 2010; 26(6):841–842. https://doi.org/10.1093/bioinformatics/btp033 PMID: 2101278

32. Aamot HU, Hofgaard IS, Lysøe E. Complete genome sequence of Lutheibacter rhizovicinus strain LJ96T, isolated from the rhizosphere of barley (Hordeum vulgare L.) in Denmark. Genomics Data. 2017; 11:104–105. https://doi.org/10.1016/j.gdata.2016.12.012 PMID: 28123952

33. Thele R, Gumpert H, Christensen LB, Worning P, Schønning K, Weth H, et al. Draft genome sequence of a Kluyvera intermedia isolate from a patient with a pancreatic abscess. Journal of Global Antimicrobial Resistance. 2017; 10:1–2. https://doi.org/10.1016/j.jgar.2017.05.007 PMID: 28576740

34. Ma Y, Yao R, Li Y, Wu X, Li S, An Q. Proposal for Unification of the Genus Metakosakonia and the Genus Phytobacter to a Single Genus Phytobacter and Reclassification of Metakosakonia massiliensis as Phytobacter massiliensis comb. nov. Current Microbiology. 2020; 77(8):1945–1954. https://doi.org/10.1007/s00286-020-02004-4 PMID: 32506064

35. Gertz EM, Yu YK, Agarwala R, Schäffer AA, Altschul SF. Composition-based statistics and translated nucleotide searches: improving the TBLASTN module of BLAST. BMC biology. 2006; 4(1):1–14. https://doi.org/10.1186/1741-7007-4-41 PMID: 17156431

36. Consortium TU. UniProt: The universal protein knowledgebase in 2021. Nucleic Acids Research. 2021; 49(D1):D480–D489. https://doi.org/10.1093/nar/gkaa1100

37. Rambaut A, Grass NC. Seq-Gen: an application for the Monte Carlo simulation of DNA sequence evolution along phylogenetic trees. Bioinformatics. 1997; 13(3):235–238. https://doi.org/10.1093/bioinformatics/13.3.235 PMID: 9183526

38. Minh BQ, Schmidt HA, Chernomor O, Schrempf D, Woodhams MD, Von Haeseler A, et al. IQ-TREE 2: new models and efficient methods for phylogenetic inference in the genomic era. Molecular biology and evolution. 2020; 37(5):1530–1534. https://doi.org/10.1093/molbev/msaa015 PMID: 32011700

39. Schliep KP. phangorn: phylogenetic analysis in R. Bioinformatics. 2011; 27(4):592–593. https://doi.org/10.1093/bioinformatics/btq354 PMID: 21169378

40. Catoh K, Misawa K, Kuma K, Miyata T. MAFFT: a novel method for rapid multiple sequence alignment based on fast Fourier transform. Nucleic Acids Research. 2002; 30(14):3059–3066. https://doi.org/10.1093/nar/gkd436 PMID: 12166088

41. Zhang C, Scornavacca C, Molloy EK, Mirarab S. ASTRAL-Pro: Quartet-Based Species-Tree Inference despite Paralogy. Molecular biology and evolution. 2020; 37(11):3292–3307. https://doi.org/10.1093/molbev/msaa139 PMID: 32886770

42. Morel B, Schade P, Lutteropp S, Williams TA, Szöllösi GJ, Stamatakis A. SpeciesRax: A tool for maximum likelihood species tree inference from gene family trees under duplication, transfer, and loss. bioRxiv. 2021.

43. Sukumaran J, Holder MT. DendroPy: a Python library for phylogenetic computing. Bioinformatics. 2010; 26(12):1569–1571. https://doi.org/10.1093/bioinformatics/btp228 PMID: 20421198

44. Kishore R, Arnaboldi V, Van Slyke CE, Chan J, Nash RS, Urbano JM, et al. Automated generation of gene summaries at the Alliance of Genome Resources. Database. 2020; 2020. https://doi.org/10.1093/database/babaa037 PMID: 32559296

45. Wernegreen JJ. Endosymbiosis. Current Biology. 2012; 22(14):R555–R561. https://doi.org/10.1016/j.cub.2012.06.010 PMID: 22835786

46. Elliott TA, Gregory TR. What’s in a genome? The C-value enigma and the evolution of eukaryotic genome content. Philosophical Transactions of the Royal Society B: Biological Sciences. 2015; 370(1678):20140331. https://doi.org/10.1098/rstb.2014.0331
47. Kuo CH, Moran NA, Ochman H. The consequences of genetic drift for bacterial genome complexity. Genome Research. 2009; 19(8):1450–1454. https://doi.org/10.1101/gr.091785.109 PMID: 19502381

48. Zhou X, Lutteropp S, Czech L, Stamatakis A, Looz MV, Rokas A. Quartet-based computations of inter-node certainty provide robust measures of phylogenetic incongruence. Systematic biology. 2020; 69(2):308–324. https://doi.org/10.1093/sysbio/syz058 PMID: 31504977

49. Hughes AL, Nei M. Nucleotide substitution at major histocompatibility complex class II loci: evidence for overdominant selection. Proceedings of the National Academy of Sciences. 1989; 86(3):958–962. https://doi.org/10.1073/pnas.86.3.958 PMID: 2492668

50. Segerman B. The most frequently used sequencing technologies and assembly methods in different time segments of the bacterial surveillance and RefSeq genome databases. Frontiers in Cellular and Infection Microbiology. 2020; 10. https://doi.org/10.3389/fcimb.2020.527102 PMID: 33194784

51. Ochman H, Moran NA. Genes lost and genes found: evolution of bacterial pathogenesis and symbiosis. Science. 2001; 292(5519):1096–1099. https://doi.org/10.1126/science.1058543 PMID: 11352062

52. Ahmed N, Dobrindt U, Hacker J, Hasnain SE. Genomic fluidity and pathogenic bacteria: applications in diagnostics, epidemiology and intervention. Nature reviews microbiology. 2008; 6(5):387–394. https://doi.org/10.1038/nrmicro1889 PMID: 18392032

53. MacLean RC, San Millan A. The evolution of antibiotic resistance. Science. 2019; 365(6458):1082–1083. https://doi.org/10.1126/science.aax3879 PMID: 31515374

54. Maiden MC. Horizontal genetic exchange, evolution, and spread of antibiotic resistance in bacteria. Clinical Infectious Diseases. 1998; 27(Supplement_1):S12–S20. https://doi.org/10.1086/514917 PMCID: 9710667

55. Levin BR, Perrot V, Walker N. Compensatory mutations, antibiotic resistance and the population genetics of adaptive evolution in bacteria. Genetics. 2000; 154(3):985–997. https://doi.org/10.1093/genetics/154.3.985 PMID: 10757748

56. Taylor DE, Courvalin P. Mechanisms of antibiotic resistance in Campylobacter species. Antimicrobial Agents and Chemotherapy. 1988; 32(8):1107–1112. https://doi.org/10.1128/aac.32.8.1107 PMID: 3056250

57. Lan R, Reeves PR. Gene transfer is a major factor in bacterial evolution. Molecular biology and evolution. 1996; 13(1):47–55. https://doi.org/10.1093/oxfordjournals.molbev.a025569 PMID: 8583905

58. Thomas CM, Nielsen KM. Mechanisms of, and barriers to, horizontal gene transfer between bacteria. Nature reviews microbiology. 2005; 3(9):711–721. https://doi.org/10.1038/nrmicro1234 PMID: 16138099

59. Nakhleh L, Ruths D, Wang LS. RIATA-HGT: a fast and accurate heuristic for reconstructing horizontal gene transfer. In: Proceedings of the Eleventh International Computing and Combinatorics Conference (COCOON 05). LNCS #3595. Kunming: Springer; 2005. p. 84–93.

60. Linz S, Radlke A, von Haeseler A. A likelihood framework to measure horizontal gene transfer. Molecular biology and evolution. 2007; 24(6):1312–9. https://doi.org/10.1093/molbev/msm052 PMID: 17374878

61. Koskineni S, Sun S, Berg OG, Andersson DI. Selection-driven gene loss in bacteria. PLoS genetics. 2012; 8(6):100277. https://doi.org/10.1371/journal.pgen.1002787 PMID: 22761588

62. Wang Z, Wu M. A phylum-level bacterial phylogenetic marker database. Molecular biology and evolution. 2013; 30(6):1258–1262. https://doi.org/10.1093/molbev/mst059 PMID: 23519313

63. Husnık F, Chrudimský T, Hypša V. Multiple origins of endosymbiosis within the Enterobacteriaceae (γ-Proteobacteria): convergence of complex phylogenetic approaches. BMC biology. 2011; 9(1):1–18. https://doi.org/10.1186/1741-7007-9-87 PMID: 22201529

64. Kaiwa N, Hosokawa T, Nikoh N, Tanahashi M, Moriyama M, Meng XY, et al. Symbiont-supplemented maternal investment underpinning host’s ecological adaptation. Current Biology. 2014; 24(20):2465–2470. https://doi.org/10.1016/j.cub.2014.08.065 PMID: 25264255

65. Fukatsu T, Hosokawa T. Capsule-transmitted gut symbiotic bacterium of the Japanese common plataspid stinkbug, Megacopta punctatissima. Applied and Environmental Microbiology. 2002; 68(1):389–396. https://doi.org/10.1128/AEM.68.1.389-396.2002 PMID: 11772649