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Phylogeny of Gammaproteobacteria[∇]§

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The phylogeny of the large bacterial class *Gammaproteobacteria* has been difficult to resolve. Here we apply a telescoping multiprotein approach to the problem for 104 diverse gammaproteobacterial genomes, based on a set of 356 protein families for the whole class and even larger sets for each of four cohesive subregions of the tree. Although the deepest divergences were resistant to full resolution, some surprising patterns were strongly supported. A representative of the *Acidithiobacillales* routinely appeared among the outgroup members, suggesting that in conflict with rRNA-based phylogenies this order does not belong to *Gammaproteobacteria*; instead, it (and, independently, “*Mariprofundus*”) diverged after the establishment of the *Alphaproteobacteria* yet before the betaproteobacteria/gammaproteobacteria split. None of the orders *Alteromonadales*, *Pseudomonadales*, or *Oceanospirillales* were monophyletic; we obtained strong support for clades that contain some but exclude other members of all three orders. Extreme amino acid bias in the highly A+T-rich genome of *Candidatus Carsonella* prevented its reliable placement within *Gammaproteobacteria*, and high bias caused artifacts that limited the resolution of the relationships of other insect endosymbionts, which appear to have had multiple origins, although the unbiased genome of the endosymbiont *Sodalis* acted as an attractor for them. Instability was observed for the root of the *Enterobacteriales*, with nearly equal subsets of the protein families favoring one or the other of two alternative root positions; the nematode symbiont *Photorhabdus* was identified as a disruptor whose omission helped stabilize the *Enterobacteriales* root.

Although *Gammaproteobacteria* has only the taxonomic rank of class within the phylum *Proteobacteria*, it is richer in genera (~250) than all bacterial phyla except *Firmicutes* (11). It includes the paradigmatic bacterium *Escherichia coli*, well-known pathogens *Salmonella*, *Yersinia*, *Vibrio*, and *Pseudomonas*, additional pathogens occurring at the more basal and less well-resolved positions in the phylogeny like *Coxiella* and *Francisella*, and endosymbionts that can be required for survival of their insect hosts. Members exhibit broad ranges of aerobicity, of trophism, including chemoautotrophism and photoautotrophism, and of temperature adaptation (31). Morphologies include rods, curved rods, cocci, spirilla, and filaments, and the class contains the largest known bacterial cells (30). Interesting combinations of phenotypes occur, such as terrestrial bioluminescence with pathogenicity (toward insects and humans) and symbiosis (with nematodes) in the genus *Photorhabdus* (37). One feature alone, 16S rRNA sequence relationship, has been used to define the class (11).

Previous phylogenetic studies of this group indicate deep branches that make it difficult to obtain a large well-resolved phylogeny (10, 40). Taking advantage of the large number of genomes available for the class, we have followed a multiprotein approach to gammaproteobacterial phylogeny (39). A potential challenge to the validity of phylogenetic reconstruction

in bacteria is the view that horizontal transfer is so pervasive that the tree interpretation is artifactual (2). However, a central trend in the phylogenetic signals from multiple bacterial protein families can be detected that probably represents their shared vertical inheritance (25). Our methodology sought to include single-copy families most likely to reflect this central trend, applying a filter to remove the families most discordant with the majority. The resulting tree breaks with current taxonomy.

In contrast to previous studies with a small number of genomes, indicating high phylogenetic concordance among gammaproteobacterial protein families (20, 24), we found that nearly equal numbers of protein families favored one or the other of two roots for *Enterobacteriales*. However, much of this effect could be ascribed to a single taxon that was absent from the earlier studies.

MATERIALS AND METHODS

Genome selection by collapse of a preliminary tree. On 27 July 2007, there were 215 complete or incomplete genomes available for nominal *Gammaproteobacteria* at NCBI. Because the phylogenetic distribution was biased, we selected a smaller number with balanced distribution. Protein sets were taken for the 215 *Gammaproteobacteria* and for two outgroup species each from the *Alphaproteobacteria* and *Betaproteobacteria*.

An initial set of protein families that approximated one occurrence per genome among 97 *Gammaproteobacteria* were selected from the GeneTrees database (36), and the alignments were used for HMMsearch using HMMer (3) to find instances in the full set of 215 genomes. The 20 gene families with the lowest standard deviation of members per genome, the closest to single copy, were processed further. Phylogenetic trees were built for each family to resolve 54 cases of duplicate or split genes. A multiprotein species tree was prepared, aligning sequences with MUSCLE version 3.6 (7) in default mode and removing ambiguous portions of alignments with Gblocks version 0.91b (35) in default mode except for using the intermediate setting for gap tolerance; the concatenation of masked alignments containing 5,538 characters was used to prepare the tree applying MrBayes version 3.2 (28) in model-jumping mode as described

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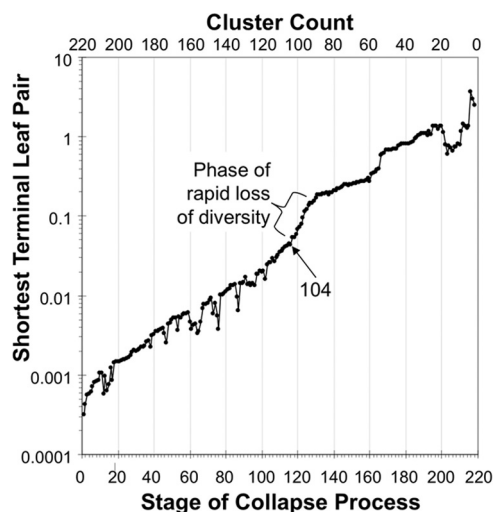


FIG. 1. Taxon selection by collapse of a preliminary tree. Collapse was a sequential process of identifying the shortest leaf pair and replacing it with a single representative leaf. This maximized diversity between clusters of the available genomes. The number of genomes selected this way was 104, since this preceded a phase in the collapse process with an especially high loss of diversity.

previously (39). A process of sequential tip pair collapse was applied to this tree, in which at each stage the terminal node with the shortest total length of its two leaves was collapsed into a single leaf whose length was the average of the two. The distance between the pair of collapsed tips, a measure of diversity reduction, was plotted against the stage of collapse (Fig. 1). The stage with 104 clusters was chosen because it preceded a phase of steep loss of genome diversity and yet yields a reasonable number for phylogenetic analysis. One genome was selected from within each of the 104 clusters, favoring complete genomes and better-known taxa. The total was raised to 110 with additional genomes: three from unrepresented genera that had become available at NCBI more recently, two that allowed comparison to a classical set of 13 taxa (20), and a nonpublic genome that became available to us. Later, two incomplete genomes were rejected because as described below their rRNA sequences suggested that they were not bacteriologically pure. The remaining 108 taxa contained four outgroup taxa, at least one member from each of the 14 gammaproteobacterial orders, and 11 taxa classified at NCBI as *Gammaproteobacteria* but not assigned to an order (see Table S1 in the supplemental material).

rRNA sequence assay of genome purity. All complete or incomplete 16S and 23S rRNA sequences were collected from 110 genomes using a curated set of seed sequences and a BLASTN-based script, resulting in 1,135 sequences (529 of 16S rRNA and 606 of 23S rRNA), of which 791 were unique. These sets were aligned using MUSCLE. For 54 rRNA fragments, sequences from the ends of a contig that aligned poorly to whole rRNAs were trimmed, leaving 337 16S rRNA and 452 23S rRNA unique sequences that were realigned using MUSCLE, manually correcting its error of splitting off small terminal portions of incomplete sequences. For all intragenomic pairs whose alignment overlapped >40 nt, the subalignment for the overlap region was taken for all available rRNA sequences (78 subalignments for 16S rRNA, 124 for 23S rRNA). Intragenomic sequence pairs for 16S and 23S rRNA were 409 and 658 for the full alignments and 228 and 558 for the subalignments, respectively. For each full and subalignment, a distance matrix was taken using distmat (26). All cases where an intragenomic pair had a distance score above the tenth percentile (39) for a partner were investigated further.

(i) In the 23S rRNA subalignment of positions 808 to 1075, a fragment from *Beggiatoa* sp. strain SS had a higher distance score to its intragenomic partner than to 99% of all its distance scores and also ranked highly in other 23S subalignments. BLAST found one of the pair to have 83% identity to *Coxiella*, but the other had 96% identity to *Haemophilus influenzae* PittGG. Because this incomplete genome, derived from a single bacterial filament isolated from sea sediment, contained highly divergent 23S fragments, we considered that its protein sequences might be contaminated with those from additional microbes, and it was rejected from our analysis.

(ii) A 23S rRNA fragment from the incomplete *Azotobacter vinelandii* genome

had high-ranking intragenomic distance scores and in one subalignment ranked as high as 98% among all distance scores. This fragment showed 96% identity to *Desulfovibrio* 23S rRNA, while the other eight sequences from this genome matched *Azotobacter* 23S rRNA. We rejected this genome from our analysis due to the potential for contamination with protein sequences from other organisms. The complete *A. vinelandii* genome now available no longer has this rRNA contamination.

(iii) A short high-scoring 23S rRNA fragment from the complete *Chromohalobacter* genome proved to be a likely regulatory region of an operon encoding proteins known to bind that region of 23S rRNA (38).

(iv) One high-scoring intragenomic rRNA distance was ascribed to misalignment by MUSCLE.

(v) The remaining high-scoring intragenomic rRNA distances were from incomplete genomes, and the top BLAST hit of each contender was to the nominal genus though low scoring. These cases could be ascribed to low-quality sequence data.

rRNA trees. Analysis began with the rRNA sequences from 108 genomes remaining after the filters of the genomic purity assay. In cases of multiple rRNA sequences, a single full-length sequence was chosen as having the shortest branch length in a preliminary tree or, if only fragments, were available, they were merged. With single representative 16S and 23S rRNA sequences for each genome, the MUSCLE alignments were further evaluated using the secondary structure models available at the comparative RNA web site (5). Sequences were manually adjusted to adhere to the predicted helical and unpaired regions of the rRNAs, using compensatory base change evidence to support homology assignment (17). Regions of the alignment that included length heterogeneity were minimized by flanking structural support (i.e., compensatory base change evidence) with remaining regions of ambiguous alignment masked (12). In several cases of mismatch to the structural model, the flanking ends of partial sequences were also masked. Resulting alignments were processed using tools from the jRNA web site (13) to create alignments, base pair analyses, and input files for MrBayes annotated for use of the doublet model, which assigns an RNA substitution matrix to base pairs.

Bayesian phylogenetic trees were generated for the 16S and 23S rRNA masked alignments and for a concatenation of the two. For each, MrBayes was run three times as a single MCMC chain for 2×10^6 generations, with a GTR + Γ + I model (Γ distribution approximated by 4 rate categories) for among-site rate variation, differing at each site, and the doublet substitution model for base pairs. In all cases, burn-in was attained well before 1.5×10^6 generations, and the trees from the final 0.5×10^6 generations were pooled from triplicate runs (7,500 trees total) to build a consensus tree.

Multiprotein data sets. An automated workflow for gene family selection, discordance filtering, and tree building was implemented through a series of Perl scripts. All proteins for a group of genomes were sorted into families using all-versus-all BLASTP followed by OrthoMCL version 1.4 (21). Approximately single-copy protein families were selected by choosing (i) a core set of complete and well-behaved ingroup genomes from among the taxa and tolerances for (ii) the number of core genomes missing from the family and (iii) the number of genomes with multiple members in the family (see Table 2, footnote a).

For the full taxon set of 108 study taxa, this process yielded 404 families, with two taxa especially poorly represented: *Candidatus* Carsonella in 69 families and *Candidatus* Endoriftia in 58. Since only 182 proteins are annotated for the extremely small *Ca. Carsonella* genome, its low family representation was not pursued further. The incomplete *Ca. Endoriftia* genome had many genes split among different contigs, frameshifted by apparent sequencing errors or simply uncalled. This genome was queried by all members of its unrepresented families using TBLASTN, and 317 of the unrepresented families received good hits, of which the 148 most reliable models for missing proteins were added to the corresponding families and to the *Ca. Endoriftia* protein file.

The resulting families were subjected to discordance filtering, that is, removal of the 10% with the most anomalous phylogenetic signal (e.g., potential cases of lateral gene transfer). Family members were aligned (MUSCLE) and the alignments were masked to remove unreliably aligned regions (Gblocks, with intermediate gap tolerance); families obliterated by Gblocks were rejected. For each family, the preferred amino acid substitution model among the 20 available in RAxML 7.0.4 was chosen, most frequently the WAG model, based on a script provided with RAxML (34). Fifty quick bootstraps (RAxML option -x) were taken for the family to determine high-support bipartitions (HSBs) of the taxa according to a chosen cutoff (see Table 2). Families with no HSBs were immediately rejected; otherwise, pairwise normalized HSB conflict counts were taken for all pairwise comparisons of the families, after modifying HSBs for shared taxa. Unshared taxa were removed, causing some HSBs to merge and others to become trivial. Conflicting HSBs were counted and normalized, dividing by the

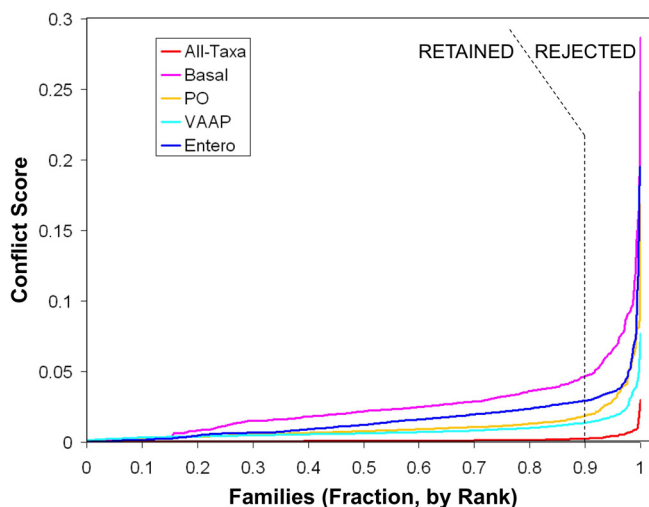


FIG. 2. Family conflict. The conflict scores for each set of protein families identified those with egregious phylogenetic signals. Families scoring above the 90th percentile were rejected.

product of the two families' HSB counts. A conflict score was assigned to each family (its average pairwise normalized HSB conflict count), which had little correlation with numbers of characters, taxa, or HSBs (data not shown). A given fraction (10%) of families with the highest conflict scores were removed. Figure 2 shows that 10% is an effective and safe rejection rate. Note that discordance filtering does not refer to (and thereby circularly reinforce) any particular phylogenetic tree topology.

Phylogenetic analysis. The masked alignments for the remaining families were concatenated. We found that each individual MrBayes run would canalize into a single tree topology, but duplicate runs would find slightly different topologies. We also found that the product of the faster (MIX) mode of RAxML depended on the starting tree provided. Instead we used the slowest RAxML mode, with gamma-distributed rate heterogeneity, which yielded a single final tree whether we started with random trees or with the products from the MIX-mode runs. Protein families were assigned their preferred substitution models found previously. To provide support values, tree building was repeated for 50 whole-protein jackknife (random removal of half the protein families) resamplings and for all single-taxon jackknife resamplings.

The above approach was applied to the full taxon set and to the four main taxon subgroups as detailed in Table 1. In addition to the use of multiple amino acid substitution matrices with the gene-partitioned concatenations, we tested uniform usage of the new LG matrix (18). Maximum likelihood trees were prepared with uniform (unpartitioned) use of LG for the concatenations of the full taxon set and all four subgroups using RAxML version 7.2.3. All trees had topologies identical to those of their counterparts that had been produced with multimodel partitioning, except that the Basal subset and the full taxon set (in the Basal region) each had a single case of a branch crossing a single node. Both cases involved the node with the poorest jackknife support in the whole tree (represented by multifurcation in Fig. 3). Comparing the two different trees in these two cases, using each of the two substitution matrix sets by the Approximately Unbiased test, the differences were not significant at the 0.05 level, and the trees from multimodel partitioning scored higher than those from uniform application of the LG matrix.

Amino acid bias. To examine amino acid bias, the 61 proteins from the 356 all-taxon set that included *Ca. Carsonella* were used to measure amino acid bias for each taxon relative to all the gammaproteobacterial taxa according to the method of Karlin (15) (Fig. 4). To examine the significance of amino acid bias, the chi-squared test implemented in TreePuzzle version 5.2 (29) was applied; 79 of 108 genomes were biased at the 5% level. As a second test of the significance of amino acid bias, we generated an artificial distribution of amino acid bias scores by making 1 million simulated 61-protein concatenations where each character state was chosen randomly from each column of the study supermatrix; bias against the gammaproteobacterial taxa was calculated for each, and significance levels were taken from the sorted list of scores. According to this test, 105 of the 108 genomes were biased at the $P < 0.05$ level, and 81 (which included all 79 of those identified by TreePuzzle) were biased at the $P < 0.000001$ level.

Statistical tree tests. The Approximately Unbiased test as implemented in CONSEL v0.1j (32) was applied, and site likelihood scores were prepared with RAxML.

RESULTS

Taxon selection. Over 200 complete and incomplete gammaproteobacterial genomes were available when our project began, but they had a biased taxonomic distribution; the six genera *Escherichia*, *Haemophilus*, *Pseudomonas*, *Shewanella*, *Vibrio*, and *Yersinia* accounted for over half the genomes. These near-duplicate genomes contain little additional phylogenetic information; we selected a subset of the genomes with maximized diversity by (i) building a preliminary phylogenetic tree, (ii) sequentially clustering close relatives, and (iii) choosing a stage in the clustering process that preceded a phase of rapid loss of diversity (Fig. 1). Two incomplete genomes were rejected because analysis of rRNA sequences suggested that these two projects might be contaminated with foreign DNA. The final set of 108 genomes included four outgroup taxa (two each from the sister class *Betaproteobacteria* and the subtending class *Alphaproteobacteria*), at least one representative of each of the 14 orders listed in *Bergey's Manual* (11), and 11 taxa that had been assigned at NCBI to the class *Gammaproteobacteria* but not to a particular order (see Table S1 in the supplemental material). This genome selection process is summarized in Table 1.

rRNA trees. As important a tool as 16S rRNA phylogeny has been in bacteriology, we and others have found that this single gene contains insufficient phylogenetic information to robustly recover deep relationships, such as those of interest in this work. For our 108 selected genomes, we made trees for 16S and 23S rRNAs (see Fig. S1 and S2 in the supplemental material). Both trees were questionable because numerous ingroup genomes (21 and 17, respectively) were placed between the two *Alphaproteobacteria* and *Betaproteobacteria* outgroups; moreover, the two trees differed substantially at

TABLE 1. Selection of genomes

Step	No. of genomes remaining		
	Ingroup	Core ^a	Outgroup
Collect gammaproteobacterial genomes from NCBI	215		0
Select alpha-/betaproteobacterial outgroups	215		4
Collapse preliminary 20-protein tree	100		4
Complete the "classical" genome set (20)	102		4
Add recently available genomes from unrepresented genera	106		4
Apply rRNA purity filter	104		4
Select core genomes ^a for family selection	104	31	4
Generate main tree, split into isolated subgroups			
Basal subgroup	19	14	3
PO subgroup	27	17	2
VAAP subgroup	37	24	2
Entero subgroup	19	8	2

^a The core genomes are a subset of the ingroup that have complete sequences and were further selected based on their diversity and their low compositional bias; these were used in the near-single-copy filter for protein families.

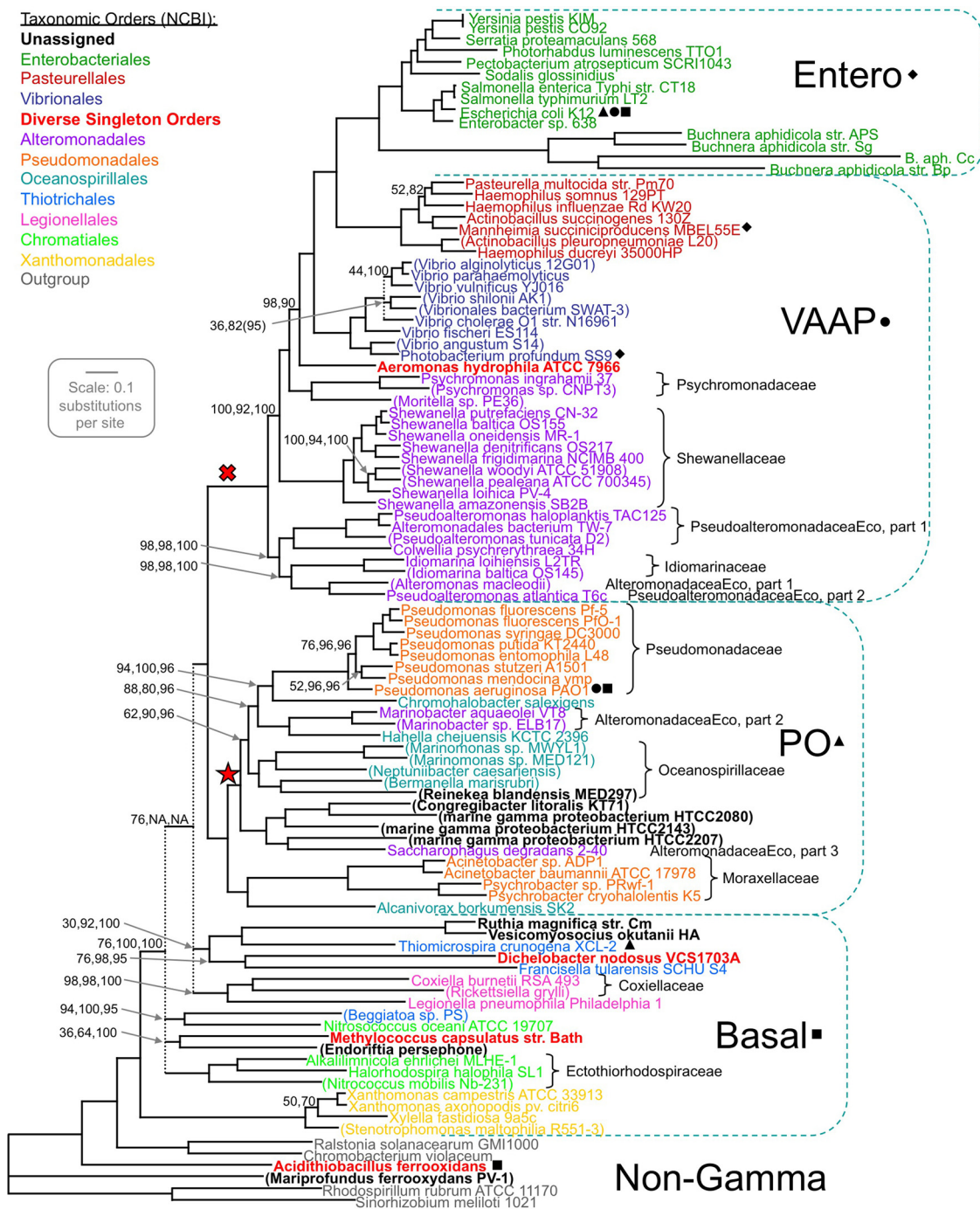


FIG. 3. Gammaproteobacterial phylogeny. Four subgroups of the taxa (excepting *Ca. Carsonella*, which is omitted here) were found not to mingle, by protein-jackknife subsampling of an all-taxon concatenation of 356 protein alignments. Longer concatenations were prepared for each taxon subgroup (Table 1), and tree topologies were built for each; small black geometric symbols link each subgroup to the outgroups used in building its subtree. The four regional trees were merged back into the topology for the all-taxon data set. Branch lengths for the merged topology were computed based on the all-taxon data set, and the four of its bifurcations that had less than 25% protein-jackknife support were collapsed, leaving three multifurcating nodes (dotted lines). Support values are shown (in the following order: all-taxon protein 50% jackknifing, taxon-subgroup protein 50% jackknifing, and taxon-subgroup single-taxon jackknifing) when any of these are <100%; cases marked NA (not applicable) were for nodes not included in taxon subgroup studies. The Entero region subtree was prepared by a two-step procedure described in the text that makes the usual support values inapplicable; however, for the first-step subtree prepared for the core *Enterobacteriales* taxa whose topology persists in this figure, protein and taxon jackknifing gave 100% support at each node. Taxon names are given in parentheses for incomplete genomes and are color coded according to the taxonomic order designation at NCBI. Taxonomic families represented by more than one taxon, except in cases in which the lone family represents the order, are bracketed, extending brackets to include unassigned taxa as necessary; only two of these nine families are split in our tree. Two nodes of interest are marked: the X indicates a node supported by a rare indel (10), and the star indicates a fully supported node that groups members of *Pseudomonadales*, *Alteromonadales*, and *Oceanospirillales* while excluding other members of each order. All known members of the latter clade, and no other known bacteria, autoregulate a ribosomal protein operon through strong mimicry of a 23S rRNA domain (38).

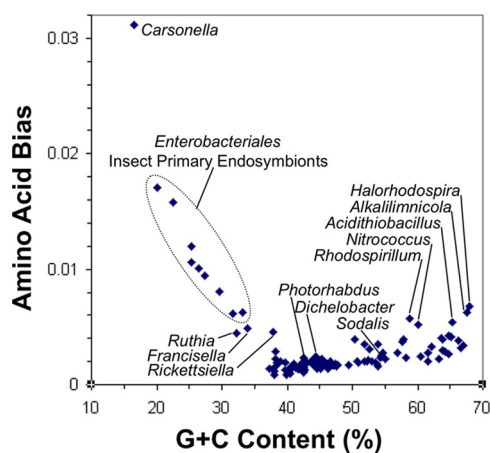


FIG. 4. Compositional bias. Amino acid bias was measured according to the method of Karlin (15) for the set of 61 all-taxon protein families that contain *Ca. Carsonella* and plotted against nucleotide composition, producing a horn of A+T-rich endosymbionts with *Ca. Carsonella* at its tip.

the deeper nodes. A tree for the concatenated alignment of the two rRNAs had the same problems (see Fig. S3 in the supplemental material). In a multiprotein study of alphaproteobacterial phylogeny, it was found that the combined rRNAs had the phylogenetic inaccuracy (measured as the level of disagreement between two different phylogenetic reconstruction methods) of ~ 2.5 single-copy proteins combined; moreover, accuracy increased as more proteins were employed, at least up to 100 proteins (39). Therefore, we took a multiprotein approach to gammaproteobacterial phylogeny.

Telescoping multiprotein approach. The first step in our multiprotein approach (Table 2) was to select a large number of protein families with (nearly) single-copy distribution patterns, because these are likely to have similar histories that reflect the organismal phylogeny. A strict single-copy criterion would only have allowed one gene family, so the criterion was slightly relaxed. A set of 31 core taxa was selected that had complete genomes of ≥ 1 Mbp and were phylogenetically representative (see Table S1 in the supplemental material), and the relaxed single-copy criterion allowed families with up to two core taxa unrepresented and up to six core taxa with duplicate membership. Cases of multiple membership for a taxon were expediently resolved by removing both (or all) members from that taxon in that family. This yielded 405 gene families.

The alignments for these families were masked by removing ambiguously aligned regions. Families with no sequence remaining after masking were rejected, as were families whose bootstrap trees produced no highly supported taxon bipartitions. These two filters eliminated families that were grossly deficient in phylogenetic signal.

To further refine the gene families and minimize the potential influence of horizontal transfer, a phylogenetic discordance filter was imposed to eliminate the 10% of protein families least concordant with the others. The set of highly supported taxon bipartitions for each family were compared to generate a normalized conflict matrix (Fig. 2). Families with high average discordance were eliminated without referring to (and

thereby circularly reinforcing) an a priori tree topology, leaving 356 families (see Table S2 in the supplemental material). Fifteen of 19 COG (clusters of orthologous groups) categories were significantly overrepresented (translation proteins especially) or underrepresented (uncategorized proteins especially) among these families (see Fig. S4 in the supplemental material).

The masked alignments for these families were concatenated into a supermatrix, and a maximum likelihood tree was produced. Its robustness was evaluated by whole-protein 50% jackknifing, that is, generating subsets of the data from which a random half of the protein families had been removed and computing a tree for each. Support for each node in the main tree was then evaluated as the fraction of protein jackknife trees that produced the node. A rationale for the use of this support measure is presented in the Discussion. Several nodes received very low support. However, the protein jackknife data revealed five regions of the tree whose boundaries were never crossed (except by the anomalous taxon *Ca. Carsonella*, which we discarded, as further described below), indicating isolated groups of taxa that could be studied separately, reuniting the resulting telescoped trees into a better-resolved composite tree. One group was the *Enterobacteriales* clade (“Entero” in Fig. 3). The second group, termed “VAAP,” consists mainly of *Vibrionales*, *Alteromonadales*, *Aeromonadales*, and *Pasteurellales*; the Entero clade emerges from the VAAP group. A sister clade to the VAAP and Entero groups is the “PO” group, consisting mainly of *Pseudomonadales* and *Oceanospirillales*. The “Basal” region is a paraphyletic group of the most anciently diverging lineages in the class. Finally, the “Non-Gamma” region contains the alpha- and betaproteobacterial outgroups and additionally contains “*Mariprofundus*,” which had been assigned to the *Gammaproteobacteria* by the NCBI taxonomy but to a novel lineage outside the *Gammaproteobacteria* by rRNA analysis (8), and, surprisingly, *Acidithiobacillus*, which has been considered a member of the *Gammaproteobacteria* (16). The *Acidithiobacillus* result is not due to an anomaly of the particular genome project studied here; a second *Acidithiobacillus* project has since become available, and it shows near-identity to the earlier project in all the protein families employed here.

TABLE 2. Selection of protein families

Step	No. of families remaining (by genome group)				
	All-taxon	Basal	PO	VAAP	Entero
OrthoMCL	25,292	8,227	12,727	12,482	7,365
Near-single-copy filter ^a	405	684	679	712	1,403
Masking filter ^b	404	684	679	712	1,403
High-support bipartition filter ^c	396	680	678	712	1,402
Discordance filter	356	616	611	640	1,262

^a This filter removed families exceeding the tolerance for either absent or multiple representation of core taxa in the family membership. The following tolerances were used (absent/multiple): all-taxon, 2/6; Basal, 5/1; PO, 1/1; VAAP, 2/1; Entero, 0/0.

^b Families with no sequence remaining after masking by Gblocks was removed.

^c Taxon bipartitions for a protein family were counted as highly supported when they appeared among $\geq 75\%$ of the bootstrap trees for the family (except for the all-taxon families, where a 95% cutoff was applied); the filter removed families with no high-support bipartitions.

Aiming to improve resolution, each of the four gammaproteobacterial regions (Basal, PO, VAAP, and Entero) of the tree were separately reanalyzed with larger numbers (611 to 1,262) of protein families, by the same selection/discordance-filtering/tree-building approach described above, and as detailed in Table 2. Even the very large Entero collection (1,262 protein families) shows functional bias compared to all 4,149 encoded proteins of *E. coli*; overrepresented COG categories were translation, coenzyme metabolism, cell division, DNA metabolism, and protein processing (chi-squared *P* values of $<10^{-10}$, $<10^{-10}$, $<10^{-7}$, $<10^{-7}$, $<10^{-4}$, and $<10^{-3}$, respectively), while the uncategorized, carbohydrate metabolism, and transcription genes were underrepresented ($<10^{-10}$, $<10^{-7}$, and <0.03 , respectively) (see Fig. S4 in the supplemental material). Also, none of the 185 *E. coli* genes annotated in genomic islands were among the Entero families. Support values for the nodes in each regional tree were then evaluated by whole-family 50% jackknifing as described above and also by single-taxon jackknifing, in which, for each in-group taxon, a tree was built after that taxon was removed. Before reuniting these telescoped regional trees, the *Enterobacteriales* region was examined further, initially because it contained the anomalous *Ca. Carsonella* genome.

Compositional attraction. The *Ca. Carsonella* genome (22) is unusual for many reasons: it has an extremely small and A+T-rich genome and is represented in only 61 of our 356 families. It was located on an extremely long branch in our original tree within a group of other insect endosymbionts, themselves (except for *Sodalis*) on long branches and with small, A+T-rich genomes. It was the only taxon that crossed a tree regional boundary, found among the insect endosymbiont *Enterobacteriales* in 48 of the whole-protein jackknife replicates but with the A+T-rich *Francisella* or *Rickettsiella* genomes (Basal region) in the other two. Either or both of these tree positions may have been artifacts resulting from amino acid bias, in turn arising from nucleotide bias in these genomes. Using the protein families that contain *Ca. Carsonella*, we measured the amino acid bias of each taxon relative to all *Gammaproteobacteria* combined (15). In a plot of amino acid bias against nucleotide bias (Fig. 4), some taxa fell into a prominent A+T-rich horn, with *Ca. Carsonella* at its far tip, eight insect endosymbionts that are its usual neighbors in the trees, and three members of the Basal region: *Candidatus Ruthia*, *Francisella*, and *Rickettsiella*. We tested more directly the possibility that the placement of *Ca. Carsonella* was artificial by building a tree after removing its eight neighboring A+T-rich taxa from the analysis; in this tree, *Ca. Carsonella* jumped to the group with the next-most-similar composition, appearing with *Francisella*. In a similar tree built after further removing *Ca. Ruthia*, *Francisella*, and *Rickettsiella*, *Ca. Carsonella* appeared in yet a new region of the tree, among *Pseudomonas* species. We conclude that either (i) the placement of *Ca. Carsonella* among the insect endosymbionts is artifactual, (ii) its secondary placement with *Francisella* is artifactual, or (iii) both these placements are artifactual such that *Ca. Carsonella* may not even belong to the *Gammaproteobacteria*. *Ca. Carsonella* was excluded from further analysis.

The group of insect endosymbionts with small, A+T-rich genomes, consisting of *Buchnera*, *Candidatus Baumannia*, *Candidatus Blochmannia*, and *Wigglesworthia*, appeared as a single

clade in the Entero region of initial trees. These genomes all have high amino acid bias relative to the remaining *Gammaproteobacteria* (Fig. 4), raising the possibility that the clade may have been an artifact of compositional attraction. We tested these genomes as we had *Ca. Carsonella*, building a tree by adding each one singly after omitting all their A+T-rich neighbors. Each member of the group was confirmed to fall among the *Enterobacteriales* in single-member tests using the all-taxon data set. Although each of the *Ca. Baumannia*-*Ca. Blochmannia*-*Wigglesworthia* strains were placed as before next to *Sodalis*, each *Buchnera* strain was placed at the base of the *Enterobacteriales*. These placements were reproduced when the single-member tests were repeated with the *Enterobacteriales*-focused data set (Fig. 5A and B, part i). This split placement supports the idea that the original grouping of all these endosymbionts was due to compositional attraction.

The presence of *Sodalis* was suspected to have influenced these results, since it is itself an insect endosymbiont (of the same insect host as *Wigglesworthia*) with a somewhat reduced, though compositionally unbiased (54% G+C), genome. Omitting *Sodalis* affected the placement of those endosymbionts that had clustered with it in the single-member tests; *Ca. Blochmannia* and *Wigglesworthia* moved to the base of the *Enterobacteriales*, while *Ca. Baumannia* moved to an intermediate position (Fig. 5A and B, part i). Thus, the basal position of the *Buchnera* strains was independent of *Sodalis*, while the positions of the other endosymbionts were *Sodalis* dependent and therefore uncertain.

Instability of the *Enterobacteriales* root. The above experiments revealed instability in the root position of the *Enterobacteriales*. Our all-taxon and *Enterobacteriales*-focused trees, most of their taxon- or protein-jackknife support trees, and most of the endosymbiont study trees show *Photorhabdus* at the root ("Plu-root" in Fig. 5A), but a substantial number of members of the last group show the *Escherichia*-*Salmonella*-*Enterobacter* clade at the root ("Eco-root" in Fig. 5A). These two frequent rooting positions are separated by two internodes that themselves are strongly disfavored as rooting positions by the Approximately Unbiased (AU) test (32) (Fig. 5C).

The *Enterobacteriales* present an interesting situation where each of two different trees are supported by nearly equal subsets of the protein families. When the A+T-rich endosymbionts were excluded and the *Enterobacteriales* protein families for the remaining core taxa (and containing at least one out-group) were sorted into those favoring either the Plu- or the Eco-root, the family counts were similar (464 and 428, respectively) (see Table S3 in the supplemental material). The *Sodalis* effect on endosymbiont placement was reexamined with the *Enterobacteriales* Plu- or Eco-root-favoring family subsets and found to occur with each subset (Fig. 5B, parts ii and iii). These two family subsets were not distinct in terms of functional classification (COG types) or in terms of fast- or slow-evolving genes as judged by total tree branch length for each family, nor were they substantially segregated when mapped to the *E. coli* genome. For each of the two family subsets, the supermatrix produced a tree with the respective root, with 100% bootstrap support (Fig. 5B, parts ii and iii). One explanation for such a situation could be a massive horizontal transfer event followed by gene sorting; however, there is no single within-*Enterobacteriales* genetic exchange path that would convert one tree to

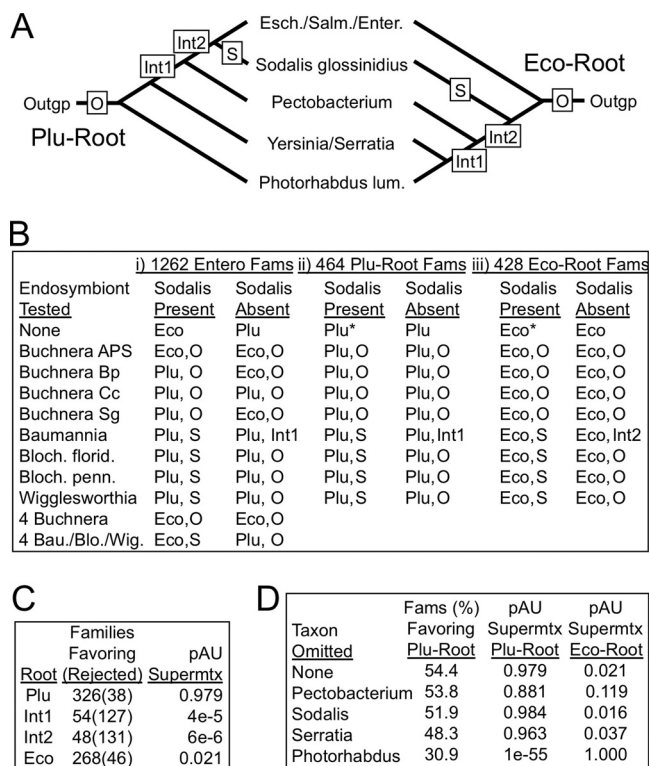


FIG. 5. Positions of the root and endosymbionts among the *Enterobacteriales*. (A) Two root positions (Plu and Eco) were obtained in our study; positions taken by A+T-rich endosymbionts (O and S) and two tested intermediate root positions (Int1 and Int2) are marked. Esch., *Escherichia*; Salm., *Salmonella*; Enter., *Enterobacter*; lum., *luminescens*; outgp, outgroup. (B) Positions taken by endosymbionts and their effect on rooting. The full *Enterobacteriales* taxon set was reduced to a core by removing all eight A+T-rich genomes (retaining the two outgroups), then each alone, or in groups of four, was added back to the data set. The compositionally unbiased endosymbiont *Sodalis* was excluded or not, and either the full set of Entero protein families (Fams) (part i) or subsets with at least one outgroup and favoring either the Plu-root (part ii) or the Eco-root (part iii) were used. The resulting root position and the position taken by the tested endosymbiont is reported. *, bootstrap analysis was performed for these trees, with support measuring 100% for every node. Bau., *Ca. Baumannia*; Blo. or Bloch., *Ca. Blochmannia*; Wig., *Wigglesworthia*; florid., *floridanus*; penn., *pennsylvanicus*. (C) Intermediate root positions disfavored. Each of the 696 protein families containing all core taxa and both outgroups were tested for preference of four hypothetical root positions using the *P* value of the Approximately Unbiased test (pAU), and pAU is reported for the supermatrix (Supermtx) that combined these 696 protein families. A pAU of <0.05 is typically taken as a rejection. (D) Identifying *Photorhabdus* as a disruptive taxon. For each of the 696 Entero families used in panel C, the indicated core taxa were singly omitted and pAU was taken for either the Plu-Root or Eco-Root topology, to determine the topology favored by the family; pAU was also taken for the two topologies based on the corresponding supermatrix.

the other, because the alternate topologies are separated by two disfavored internodes. We identified the families among our 356-family all-gammaproteobacterial data set corresponding to the *Enterobacteriales* Plu- or Eco-root-favoring families, and we built all-gammaproteobacterial trees with the supermatrix for each subset. This might have identified a gammaproteobacterial source for a hypothetical massive horizontal gene

transfer event; instead, the two all-taxon family subsets produced trees like that for the whole set, except again with the respective *Enterobacteriales* rooting. This test also ruled out the concern that the alternate-root phenomenon was an artifact of the particular two outgroup genomes chosen for the *Enterobacteriales*-focused data set, since the test effectively increased the outgroup from 2 to 89 members.

The observation of two large gene family subsets that favor different tree topologies appears to stand in contrast to earlier studies that found high concordance among gammaproteobacterial protein families (20, 24). These studies included only three genera (*Escherichia*, *Salmonella*, and *Yersinia*) from our core *Enterobacteriales* set; thus, the extra genomes in our study (*Sodalis*, *Serratia*, *Photorhabdus*, and *Pectobacterium*) may be responsible for the difference. Omitting each of these genomes one by one from either the Plu- or Eco-root-favoring supermatrix had no effect on remaining tree topology, except that when *Photorhabdus* was omitted, both supermatrices produced the Eco-root; this identified *Photorhabdus* as a disruptive taxon. These four taxa were also removed one by one from each of the 696 *Enterobacteriales* protein families that contain all core taxa and both outgroups, assessing preference for the Plu- or Eco-root (Fig. 5D). Omission of *Photorhabdus* made the families more concordant, shifting the fraction of Plu-favoring families from 0.54 to 0.31, confirming *Photorhabdus* as a disruptor.

Final tree. Based on the difficulties with the *Enterobacteriales*, its regional tree was reconstructed as follows. The A+T-rich genomes and *Photorhabdus* were omitted, and a tree was built only for the remaining taxa; support for each node by whole-family and single-taxon jackknifing was 100%. Since the four *Buchnera* genomes all separately appeared in the same position of this subtree, independently of *Sodalis*, they were included in a second round of tree building, together with *Photorhabdus*; these taxa were added back to the data set to build a larger tree, constraining to the topology of the initial subtree (since without constraint the *Buchnera* and *Photorhabdus* genomes altered that topology [Fig. 5B, part i]). Since its membership in the *Enterobacteriales* is unconfirmed, *Ca. Carsonella* was excluded from the regional tree, as were the endosymbionts *Ca. Baumannia*, *Ca. Blochmannia*, and *Wigglesworthia*, due to uncertainty arising from their sensitivity to the presence of *Sodalis*.

The four regional trees were then merged back into their positions in the original all-taxon tree, yielding the final tree depicted in Fig. 3. Support values reported are the 50% whole-protein jackknife values for the all-taxon data set, the corresponding values from each regional subtree, and single-taxon jackknife values from each regional subtree. A few support values were so low as to warrant the introduction of two multifurcations in the Basal region and one in the VAAP region.

DISCUSSION

Multiprotein phylogeny produces more robust trees than single-gene approaches, which better serve bacteriologists in interpreting biological data. A concrete example is the fully supported clade marked by a star in Fig. 3 whose members (*Pseudomonas*, *Marinomonas*, *Saccharophagus*, etc.) share a molecular biological trait that is unique among bacteria: their

S19 ribosomal protein operon contains a strong mimic of a 23S rRNA fragment that likely confers autoregulation on the operon (38). This trait would have required a more complex explanation than simple vertical inheritance according to 16S rRNA trees, since to our knowledge no such trees, including our own, recover this clade. (It was recovered with weak support in our 23S rRNA tree but was again lost in the combined 16S/23S rRNA tree.)

Although performance is improved, multiprotein phylogeny is still subject to some of the same artifacts and difficulties as single-gene approaches, such as long-branch attraction and poor resolution of branches that are deep and short. Telescoping allowed data from more protein families to be applied to isolated subgroups of taxa, but this was still insufficient to resolve all nodes or solve difficulties with highly biased genomes. We identified one genome as a phylogenetic attractor (*Sodalis*) and one as a disruptor (*Photorhabdus*); full explanation of the mechanisms of these effects will require further study and targeted genomic sequencing of more members of these clades.

This phylogenetic analysis based on hundreds of proteins for over a hundred taxa strongly supports the splitting of two families (*Alteromonadaceae* and *Pseudoalteromonadaceae*) and three orders (*Alteromonadales*, *Pseudomonadales*, and *Oceanospirillales*). A fourth order, *Thiotrichales*, also appears split but with lower support. Finally, the unity of the order *Chromatiales* could not be established. Thus, the classification based on 16S rRNA breaks down occasionally at the family level but more frequently at the order level.

The tree assigns taxonomy to previously unclassified genomes: *Reinekea* to *Oceanospirillaceae* and *Ca. Ruthia* and *Candidatus Vesicomysocius* to *Piscirickettsiaceae*; the affiliation of *Ca. Endoriftia* with *Methylococcales* requires confirmation as more genomes become available. For four genomes (*Congregibacter* and the marine strains HTCC2080, HTCC2143, and HTCC2207) the closest assigned genome was *Saccharophagus* of the split *Alteromonadaceae*. Two genomes fell among the outgroup and are therefore not part of the *Gammaproteobacteria*. For one of these, *Mariprofundus*, this placement is consistent with previous analysis of its 16S rRNA sequence that placed it as a deep branch within the phylum *Proteobacteria* (8). The other newly identified non-gammaproteobacterium is *Acidithiobacillus*, surprisingly, since this genus has long been considered a member of *Gammaproteobacteria* (16). A recent tree (K. P. Williams, unpublished data) prepared using 173 protein families from 124 bacterium-wide genomes, chosen to represent each available bacterial order, showed the same relationships of *Acidithiobacillus* and *Mariprofundus* to other proteobacteria as depicted in Fig. 3, confirming that these two genera represent independent sister groups to the beta-/gammaproteobacteria clade that arose after divergence from the *Alphaproteobacteria*. All these observations suggest revisions of the taxonomy at multiple ranks in the class and phylum.

An early multiprotein study of the *Gammaproteobacteria* used over 200 proteins, but only 13 genomes were available at that time (20). Later studies have used far fewer protein families (10 to 35 proteins) and fewer representative genomes (28 to 55 genomes) than the present analysis (4, 6, 10, 40). All these studies and ours agree on the basal branching pattern connecting the five orders *Enterobacteriales*, *Pasteurellales*,

Vibrionales, *Pseudomonas*, and *Xanthomonadales*. One of these nodes in particular (marked by an "X" in Fig. 3) has additional support from a unique 4-amino-acid deletion in RpoB; however, our results do not agree with two other suggestions based on rare indels: (i) exclusion of *Francisella* from the *Gammaproteobacteria* and (ii) a particular split of *Alteromonadales* (10). Some of the intermingling of bacterial orders observed here had been noted in one of these studies (40), but in the other studies the taxa employed or the extent of multifurcation precluded its detection.

Nearly half of the bacterial genomes currently available are incompletely sequenced, a fraction that may increase in the future, given short-read technologies and sequencing projects that do not include a goal of closing the genome. These incomplete genomes add greatly to the taxonomic diversity available for study and are nearly as rich in protein information as the complete ones, so they are worth including in such analyses despite the minor problems they raise. However, some incomplete genomes are contaminated with DNA from additional taxonomic sources and should be rejected; an rRNA impurity test was employed here to identify some such mixed genomes. Although highly divergent rRNA alleles can occur within a single genome (41), the caution exercised here was warranted; the highly divergent rRNA allele that we found in the incomplete *Azotobacter vinelandii* genome, thereby rejecting the genome, did not appear in the recently completed version of this genome project.

When our five multiprotein supermatrices (i) were partitioned according to the substitution matrix favored by each family or (ii) had the new LG substitution matrix applied uniformly, nearly identical trees resulted, with only two cases of crossing nodes, and these nodes were the least well supported in the whole tree. This agrees with a recent survey of many alignments and supermatrices that concluded that although model choice can affect tree topology, "it rarely affects evolutionary inferences drawn from the data because differences are mainly confined to poorly supported nodes" (27).

While both the jackknifing and bootstrapping approaches to determining support values remove columns from the alignment supermatrix, jackknifing does not then duplicate the remaining columns and therefore produces smaller subsamplings that can speed processing for large data sets. We prefer the random removal of whole proteins rather than single columns across all protein alignments, to better assess variation at a coarser granularity and as a double-check against possible remaining incongruity of protein families. This 50% protein jackknifing is a more stringent measure of support than the usual per-column bootstrapping, counteracting the misleadingly high support values that the latter brings to long supermatrices (23).

Compositional attraction through amino acid bias made it difficult to infer the phylogeny of the insect endosymbiont genomes, whose composition was the most biased in our taxon set. As in most previous studies, the endosymbionts joined together in a clade, with *Sodalis* when present, when all were included in a simple maximum likelihood analysis. When instead the endosymbionts were tested one at a time, *Buchnera* consistently joined at the base of the *Enterobacteriales*; unity of the *Buchnera* genomes is strongly supported by gene order studies (4). *Ca. Blochmannia* and *Wigglesworthia* may also derive from this point yet consistently branch as sister to *Sodalis*

when that taxon is included, jumping past two tree nodes. *Ca. Baumannia* appeared with *Sodalis* when present and, unlike other endosymbionts, at nearby positions when absent, weakly supporting the idea that *Ca. Baumannia* has a unique origin (Fig. 5A and B, part i). Our finding of multiple origins for the endosymbionts agrees with those of other studies that have sought to avoid compositional attraction artifacts, through the use of either nonhomogeneous substitution matrices or analysis of genomic rearrangements (4, 14). It would appear that the pattern of an *Enterobacteriales* member adopting the endosymbiotic lifestyle and becoming highly A+T-rich has occurred in multiple independent cases.

Surprisingly, nearly equal subsets of *Enterobacteriales* proteins favored one or the other of two root positions. Massive horizontal transfer is a plausible explanation in principle, although we could not identify a particular single transfer path; the pattern could be a result of numerous transfers along multiple paths. It should be noted that our family selection method removed the genes best known for horizontal transfer, those on genomic islands. Earlier studies of a classical set of 13 gammaproteobacterial genomes that includes *Escherichia*, *Salmonella*, and *Yersinia* but not *Sodalis*, *Pectobacterium*, or *Photorhabdus* found very few of the single-copy genes with evidence of horizontal transfer. We found that omitting *Photorhabdus* shifted the support by protein families from a near balance for two root positions to a preponderance for one of these roots, identifying *Photorhabdus* as a disruptor. The ambiguities left by this study show that the *Enterobacteriales* present rich problems in phylogenetic reconstruction that are not resolved simply by accumulating larger genome-scale protein supermatrices. These problems are probably due both to horizontal transfer that is especially favored by close contact with diverse cohabitants in enteric environments and to the extensive genomic alterations in multiple isolated symbiotic lineages. Future analysis of these problems within the *Enterobacteriales* is favored by the detailed mechanistic information on mutation and gene transfer known from *E. coli* and relatives.

Much of the failure to fully resolve the deeper regions of the tree can probably be ascribed to stochastic accumulation of noise that obscures reconstruction of short and ancient internodes. It may be further compounded by residual cases of horizontal gene transfer that passed through our incongruence filter. Compositional bias may be the greatest challenge facing studies on such broad scales of bacterial phylogeny as this one. The *Gammaproteobacteria* make it clear that as clades develop nucleotide bias, they can concomitantly develop amino acid bias (33), which would be expected to produce local asymmetries in amino acid substitution rates. Simple statistical tests would have rejected 73% of the taxa from our study on the basis of bias. Thus, our data set (like other data sets of broad phylogenetic scope) violated the typical assumptions of phylogenetic reconstruction algorithms regarding homogeneity and reversibility of amino acid substitution. We were able to manage the problem in some cases by the approach of placing individual biased taxa in the absence of similarly biased attractors. Some possible future solutions may be to counter bias on a per-column basis in the supermatrix, to use mixture models (19), and to use maximum-likelihood programs that do not demand model homogeneity (9). Another promising area for

improving multiprotein phylogeny is the use of rare indels (10), especially if the detection of these markers was automated and if tree building could properly weight indel data in combination with alignment data; most of these are removed in our current protocol.

The multiprotein approach to gammaproteobacterial phylogeny, applied here with some methodological advances, has improved resolution for this challenging group, and we anticipate that additional advances are within reach that will further improve performance of the approach. As more genomes accrue, the multiprotein approach should become a new standard, supplanting 16S rRNA as the basis for phylogenetic reconstructions (1).

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