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Phylogenomics and signature proteins for the alpha Proteobacteria and its main groups

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Abstract

Background: Alpha proteobacteria are one of the largest and most extensively studied groups within bacteria. However, for these bacteria as a whole and for all of its major subgroups (viz. *Rhizobiales*, *Rhodobacterales*, *Rhodospirillales*, *Rickettsiales*, *Sphingomonadales* and *Caulobacterales*), very few or no distinctive molecular or biochemical characteristics are known.

Results: We have carried out comprehensive phylogenomic analyses by means of Blastp and PSI-Blast searches on the open reading frames in the genomes of several α-proteobacteria (viz. *Bradyrhizobium japonicum*, *Brucella suis*, *Caulobacter crescentus*, *Gluconobacter oxydans*, *Mesorhizobium loti*, *Nitrobacter winogradskyi*, *Novosphingobium aromaticivorans*, *Rhodobacter sphaeroides* 2.4.1, *Silicibacter sp.* TM1040, *Rhodospirillum rubrum* and *Wolbachia* (*Drosophila*) endosymbiont). These studies have identified several proteins that are distinctive characteristics of all α-proteobacteria, as well as numerous proteins that are unique repertoires of all of its main orders (viz. *Rhizobiales*, *Rhodobacterales*, *Rhodospirillales*, *Rickettsiales*, *Sphingomonadales* and *Caulobacterales*) and many families (viz. *Rickettsiaceae*, *Anaplasmataceae*, *Rhodospirillaceae*, *Acetobacteraceae*, *Bradyrhizobiaceae*, *Brucellaceae* and *Bartonellaceae*). Many other proteins that are present at different phylogenetic depths in α-proteobacteria provide important information regarding their evolution. The evolutionary relationships among α-proteobacteria as deduced from these studies are in excellent agreement with their branching pattern in the phylogenetic trees and character compatibility cliques based on concatenated sequences for many conserved proteins. These studies provide evidence that the major groups within α-proteobacteria have diverged in the following order: (*Rickettsiales*(*Rhodospirillales* (*Sphingomonadales* (*Rhodobacterales* (*Caulobacterales*-*Parvularculales* (*RhizobialesSphingomonadales* and *Rhodospirillales* species, respectively. The results presented here also provide support for the grouping of *Hyphomonadaceae* and *Parvularcula* species with the *Caulobacterales* and the placement of *Stappia aggregata* with the *Rhizobiaceae* group.

Conclusion: The α-proteobacteria-specific proteins and indels described here provide novel and powerful means for the taxonomic, biochemical and molecular biological studies on these bacteria. Their functional studies should prove helpful in identifying novel biochemical and physiological characteristics that are unique to these bacteria.

Background

The α -proteobacteria form one of the largest groups within bacteria that includes numerous phototrophs, chemolithotrophs, chemoorganotrophs and aerobic photoheterotrophs [1]. They are abundant constituents of various terrestrial and marine environments [2]. *Pelagibacter oblique*, which is the smallest known free-living bacteria, is believed to be the most numerous bacteria on this planet (about 10^{28} cells) comprising about 25% of all microbial cells in the oceans [2]. The intimate association that many α -proteobacteria exhibit with the eukaryotic organisms is of central importance from agricultural and medical perspectives [3,4]. Symbiotic association of the *Rhizobiaceae* family members with the plant root nodules is responsible for most of the atmospheric nitrogen fixation by plants [4-6]. Many other α -proteobacteria such as *Rickettsiales*, *Brucella* and *Bartonella* have adopted intracellular life styles and they constitute important human and animal pathogens [3,7-9]. Additionally, the α -proteobacteria have also played a seminal role in the origin of the eukaryotic cell [10,11].

In the current taxonomic scheme based on 16S rRNA, α -proteobacteria are recognized as a Class within the phylum Proteobacteria [1,12,13]. They are subdivided into 7 main subgroups or orders (viz. *Caulobacterales*, *Rhizobiales*, *Rhodobacterales*, *Rhodospirillales*, *Rickettsiales*, *Sphingomonadales* and *Parvularculales*) [12]. The α -proteobacteria and their main subgroups are presently distinguished from each other and from other bacteria primarily on the basis of their branching in phylogenetic trees [14,15]. However, we have previously described several conserved inserts and deletions (indels), as well as whole proteins, that are specific for these bacteria [16,17].

In the past 3–4 years, the numbers of α -proteobacterial genomes that have been sequenced has increased markedly. In addition to > 60 complete genomes (Table 1), sequence information is available for a large number of other species. These genomes cover all of the main groups within α -proteobacteria (Table 1) and provide an enormously valuable resource for identifying molecular characteristics that are unique to them. This information can be used to identify unique sets of genes or proteins that are distinctive characteristics of various higher taxonomic groups (e.g. families, orders, etc.) within α -proteobacteria. Such genes/proteins provide valuable tools for taxonomic, biochemical and molecular biological studies [17-26]. With this goal in mind, in the present work, we have performed comprehensive phylogenomic analyses of α -proteobacterial genomes to identify proteins/ORFs that are distinctive characteristics of the various higher taxonomic groups within α -proteobacteria. The phylogenomic distribution of these proteins is also compared with the branching patterns of these species in phyloge-

netic trees and in the character compatibility cliques to develop a reliable picture of α -proteobacterial evolution.

Results and discussion

Phylogeny of alpha proteobacteria

For comparing and interpreting the results of phylogenomic analysis, it was necessary at first to examine the evolutionary relationships among α -proteobacteria in phylogenetic trees. Phylogenetic analyses for α -proteobacteria was carried out based on concatenated sequences for 12 highly conserved proteins (see Methods). The relationships among these species were examined by both traditional phylogenetic methods (viz. neighbour-joining (NJ), maximum parsimony (MP) and maximum-likelihood (ML)) and by the character compatibility approach [27]. Figure 1 presents a neighbour-joining distance tree for α -proteobacteria, showing the bootstrap scores for various nodes using the NJ, MP and ML methods. In this tree, all of the main groups or orders within α -proteobacteria (viz. *Caulobacterales*, *Rhizobiales*, *Rhodobacterales*, *Rhodospirillales* and *Sphingomonadales*), except the *Rickettsiales*, formed well-resolved clades by different methods. For the *Parvularculales*, sequence information was available from a single species and it branched with the *Caulobacterales*. The two main families of the *Rickettsiales* i.e. *Rickettsiaceae* and *Anaplasmataceae* did not form a monophyletic clade, although they constituted the deepest branching lineages within α -proteobacteria. Except for the NJ method, the relative branching of different main groups within α -proteobacteria was not resolved by other methods. The branching orders of different groups as seen here is similar to that observed previously in the rRNA trees [1,15,16,28]. Recently, after this work was completed, Williams et al. [29] have reported similar results based on phylogenetic analysis of a different large set of protein sequences.

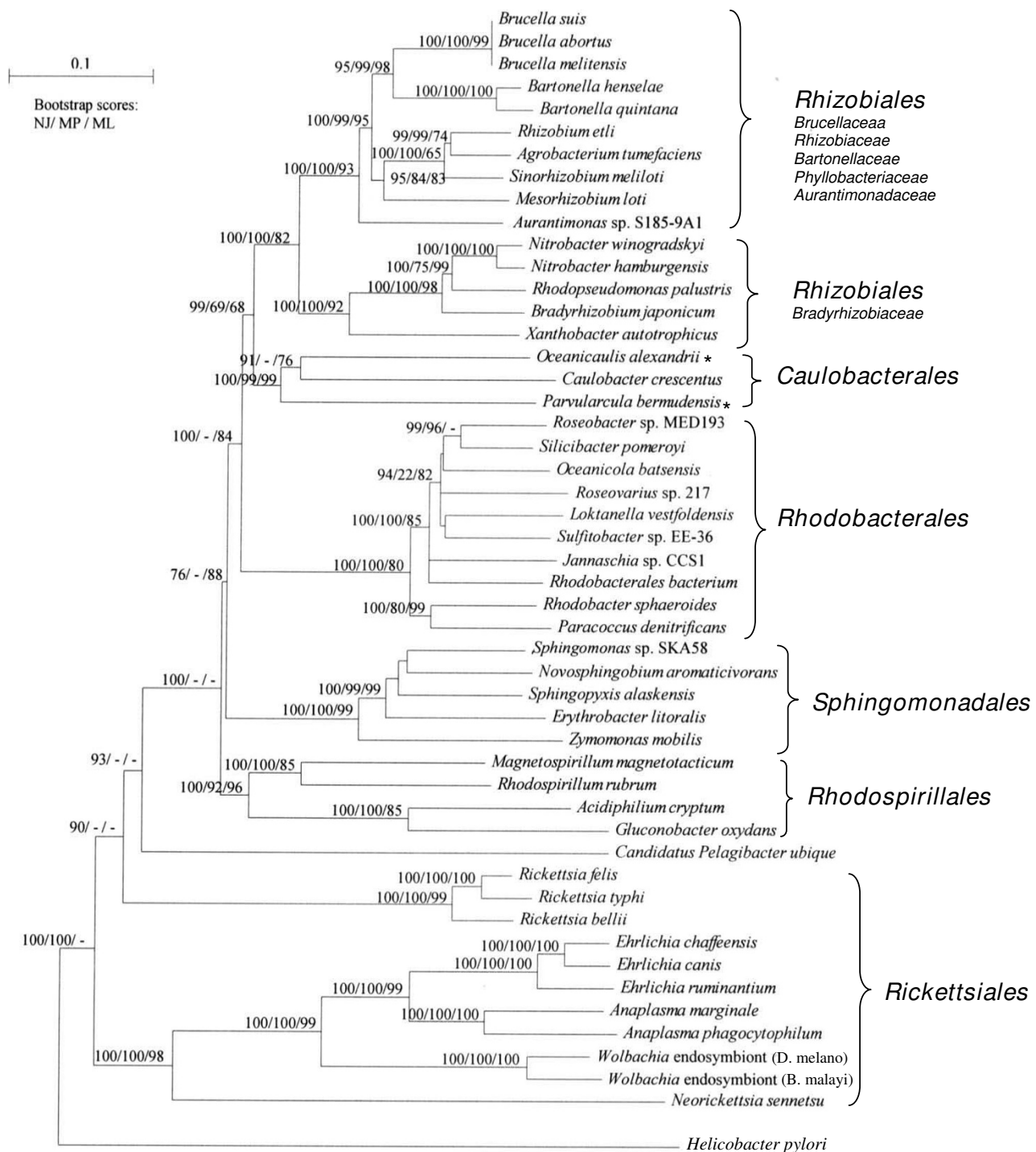
The evolutionary relationships among α -proteobacteria were also examined using the character compatibility approach [27]. This method removes all homoplastic and fast-evolving characters from the dataset [27,30,31] and it has proven useful in obtaining correct topology in cases which have proven difficult to resolve by other means [31-33]. These analyses were carried out on a smaller set of 27 species containing all main groups of α -proteobacteria and two ϵ -proteobacteria. The concatenated sequence alignment for the 12 proteins contained 896 positions that were useful for these studies (i.e. those sites where only two amino acids were found with each present in at least two species). The compatibility analysis of these sites resulted in 12 largest cliques each containing 350 mutually compatible characters. The two main relationships observed in these cliques are shown in Fig. 2. The other cliques differed from those shown only in the relative branching positions of various *Rhizobiaceae* species, which varied from each other by a single character and are

Table I: Genome sizes, protein numbers and GC contents of sequenced alpha-proteobacteria

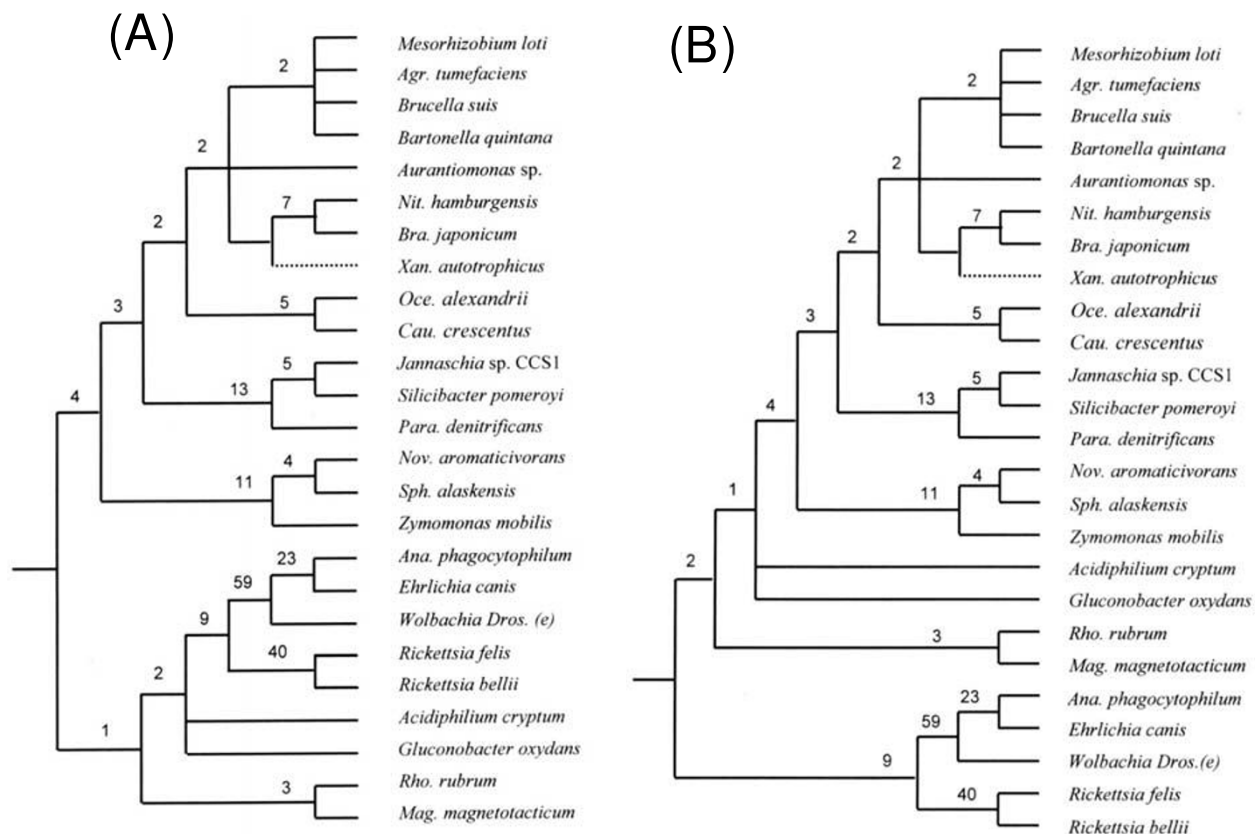
Species Name	Order	Family	Genome Size (Mb)	GC content (%)	Protein Number	Reference
<i>Bartonella bacilliformis</i> KC583	Rhizobiales	Bartonellaceae	1.4	38.2	1283	TIGR
<i>Bartonella henselae</i> str. Houston-I	Rhizobiales	Bartonellaceae	1.93	38.2	1488	[7]
<i>Bartonella quintana</i> str. Toulouse *	Rhizobiales	Bartonellaceae	1.58	38.8	1142	[7]
<i>Bradyrhizobium japonicum</i> USDA 110*	Rhizobiales	Bradyrhizobiaceae	9.1	64.1	8317	[5]
<i>Bradyrhizobium japonicum</i> BT Ail	Rhizobiales	Bradyrhizobiaceae	8.53	64.8	7622	DOE-JGI
<i>Bradyrhizobium</i> sp. ORS278	Rhizobiales	Bradyrhizobiaceae	7.5	65.5	6717	Genoscope
<i>Nitrobacter hamburgensis</i> X14	Rhizobiales	Bradyrhizobiaceae	5.01	61.6	4326	DOE-JGI
<i>Nitrobacter winogradskyi</i> Nb-255*	Rhizobiales	Bradyrhizobiaceae	3.4	2.5	3122	[42]
<i>Rhodopseudomonas palustris</i> BisA53	Rhizobiales	Bradyrhizobiaceae	5.51	64.4	4878	DOE-JGI
<i>Rhodopseudomonas palustris</i> BisB18	Rhizobiales	Bradyrhizobiaceae	5.51	65.0	4886	DOE-JGI
<i>Rhodopseudomonas palustris</i> BisB5	Rhizobiales	Bradyrhizobiaceae	4.89	64.8	4397	DOE-JGI
<i>Rhodopseudomonas palustris</i> CGA009	Rhizobiales	Bradyrhizobiaceae	5.47	65.0	4820	[41]
<i>Rhodopseudomonas palustris</i> HaA2	Rhizobiales	Bradyrhizobiaceae	5.33	66.0	4683	DOE-JGI
<i>Brucella abortus</i> biovar 1 str. 9-941	Rhizobiales	Brucellaceae	3.29	57.2	3085	[69]
<i>Brucella melitensis</i> 16M	Rhizobiales	Brucellaceae	3.29	57.2	3198	[38]
<i>Brucella melitensis</i> biovar Abortus 2308	Rhizobiales	Brucellaceae	3.28	57.2	3034	[39]
<i>Brucella suis</i> 1330*	Rhizobiales	Brucellaceae	3.32	57.3	2123	[40]
<i>Brucella ovis</i> ATCC 25840	Rhizobiales	Brucellaceae	3.3	57.2	2892	TIGR
<i>Mesorhizobium loti</i> MAFF303099*	Rhizobiales	Phyllobacteriaceae	7.6	62.5	7372	[6]
<i>Mesorhizobium</i> sp. BNC1	Rhizobiales	Phyllobacteriaceae	4.94	61.1	4543	DOE-JGI
<i>Agrobacterium tumefaciens</i> str. C58	Rhizobiales	Rhizobiaceae	5.67	59.0	4661	[37]
<i>Rhizobium etli</i> CFN 42	Rhizobiales	Rhizobiaceae	6.53	61.0	5963	[35]
<i>Rhizobium leguminosarum</i> bv. <i>viciae</i> 3841	Rhizobiales	Rhizobiaceae	7.75	61.0	7263	[81]
<i>Sinorhizobium meliloti</i> 1021	Rhizobiales	Rhizobiaceae	6.69	62.2	6205	[36]
<i>Caulobacter crescentus</i> CB15*	Caulobacterales	Caulobacteraceae	4.02	67.2	3737	[51]
<i>Hyphomonas neptunium</i> ATCC 15444	Caulobacterales+	Hyphomonadaceae	3.71	61.9	3505	[46]
<i>Maricaulis maris</i> MCS10	Caulobacterales+	Hyphomonadaceae	3.37	62.7	3063	DOE-JGI
<i>Jannaschia</i> sp. CCS1	Rhodobacterales	Rhodobacteraceae	4.4	62.2	4212	DOE-JGI
<i>Paracoccus denitrificans</i> PD1222	Rhodobacterales	Rhodobacteraceae	5.25	66.8	5077	DOE-JGI
<i>Rhodobacter sphaeroides</i> 2.4.I*	Rhodobacterales	Rhodobacteraceae	4.45	68.8	4242	DOE-JGI
<i>Rhodobacter sphaeroides</i> ATCC 17025	Rhodobacterales	Rhodobacteraceae	4.54	68.2	4333	DOE-JGI
<i>Rhodobacter sphaeroides</i> ATCC 17029	Rhodobacterales	Rhodobacteraceae	4.42	69.0	4132	DOE-JGI
<i>Roseobacter denitrificans</i> OCh 114	Rhodobacterales	Rhodobacteraceae	4.3	58.9	4129	[44]
<i>Silicibacter pomeroyi</i> DSS-3	Rhodobacterales	Rhodobacteraceae	4.6	64.1	4252	[45]
<i>Silicibacter</i> sp. TM1040*	Rhodobacterales	Rhodobacteraceae	4.15	60.1	3864	DOE-JGI
<i>Gluconobacter oxydans</i> 621H*	Rhodospirillales	Acetobacteraceae	2.92	60.8	2664	[56]
<i>Granulibacter bethesdensis</i> CGDNIH1	Rhodospirillales	Acetobacteraceae	2.7	59.1	2437	Rocky Mountain Lab
<i>Acidiphilum cryptum</i> JF-5	Rhodospirillales	Acetobacteraceae	3.97	67.1	3564	DOE-JGI
<i>Magnetospirillum magneticum</i> AMB-1	Rhodospirillales	Rhodospirillaceae	4.97	65.1	4559	[57]
<i>Rhodospirillum rubrum</i> ATCC 11170*	Rhodospirillales	Rhodospirillaceae	4.41	65.4	3841	DOE-JGI
<i>Candidatus Pelagibacter ubique</i> HTCC1062	Rickettsiales	-	1.3	29.7	1354	[2]
<i>Anaplasma marginale</i> str. St. Maries	Rickettsiales	Anaplasmataceae	1.2	49.8	949	[70]
<i>Anaplasma phagocytophylum</i> HZ	Rickettsiales	Anaplasmataceae	1.47	41.6	1264	[71]
<i>Ehrlichia canis</i> str. Jake	Rickettsiales	Anaplasmataceae	1.32	29.0	925	DOE-JGI
<i>Ehrlichia chaffeensis</i> str. Arkansas	Rickettsiales	Anaplasmataceae	1.18	30.1	1105	[71]
<i>Ehrlichia ruminantium</i> str. Gardel	Rickettsiales	Anaplasmataceae	1.5	27.5	950	[72]
<i>Ehrlichia ruminantium</i> str. Welgevonden	Rickettsiales	Anaplasmataceae	1.52	27.5	888	[73]
<i>Neorickettsia sennetsu</i> str. Miyayama	Rickettsiales	Anaplasmataceae	0.86	41.1	932	[71]
<i>Rickettsia bellii</i> RML369-C	Rickettsiales	Rickettsiaceae	1.52	31.6	1429	[74]
<i>Rickettsia conorii</i> str. Malish 7	Rickettsiales	Rickettsiaceae	1.27	32.4	1374	[75]
<i>Rickettsia felis</i> URRWXCal2	Rickettsiales	Rickettsiaceae	1.59	32.5	1512	[76]
<i>Rickettsia prowazekii</i> str. Madrid E	Rickettsiales	Rickettsiaceae	1.11	29.0	835	[77]
<i>Rickettsia typhi</i> str. Wilmington	Rickettsiales	Rickettsiaceae	1.11	28.9	838	[78]
<i>Wolbachia endosymbiont</i> (<i>Drosophila</i>)*	Rickettsiales	Rickettsiaceae	1.27	35.2	1195	[79]
<i>Wolbachia endosymbiont</i> (<i>Brugia malayi</i>)	Rickettsiales	Rickettsiaceae	1.08	34.2	805	[80]
<i>Erythrobacter litoralis</i> HTCC2594	Sphingomonadales	Erythrobacteraceae	3.05	63.1	3011	GBM Foundation
<i>Novosphingobium aromaticivorans</i> DSM 12444*	Sphingomonadales	Sphingomonadae	3.56	65.2	3937	DOE-JGI
<i>Sphingopyxis alaskensis</i> RB2256	Sphingomonadales	Sphingomonadae	3.37	65.5	3195	DOE-JGI
<i>Sphingomonas wittichii</i> RW1	Sphingomonadales	Sphingomonadae	5.93	67.9	5345	DOE-JGI
<i>Zymomonas mobilis</i> subsp. <i>mobilis</i> ZM4	Sphingomonadales	Sphingomonadae	2.06	46.3	1998	[53]

* indicates species that were used in blast searches

+ Although these species are classified as Rhodobacterales, results presented here and elsewhere [29,47] suggest their placement in the Caulobacterales.
DOE-JGI, Department of Energy, Joint Genome Institute; GBM, The Gordon and Betty Moore Foundation Marine Microbiology Initiative; TIGR, The Institute for Genome Research.

**Figure 1**

A neighbour-joining distance tree based on concatenated sequences for 12 conserved proteins. The numbers on the nodes indicate bootstrap scores (out of 100) observed in the neighbour-joining (NJ), maximum parsimony (MP) and maximum-likelihood (ML) analyses (NJ/MP/ML). The species marked with * are presently not part of the Caulobacterales order, but the results of phylogenetic and phylogenomic studies presented here suggest their placement in this group.

**Figure 2**

Character compatibility cliques showing the two largest cliques of mutually compatible characters based on the two states sites in the concatenated sequence alignment for 12 conserved proteins. The cliques consisted of 350 mutually compatible characters. The numbers of characters that distinguished different clades are indicated on the nodes. Rooting was done using the sequences for *Helicobacter pylori* and *Campylobacter jejuni*. *, as in Figure 1.

shown as unresolved in Fig. 2. In both these cliques, the species from all main orders within α -proteobacteria were clearly distinguished by multiple unique characters. Further, in contrast to the phylogenetic trees where *Rickettsiaceae* and *Anaplasmataceae* did not form a distinct clade (Fig. 1), their monophyletic grouping was strongly supported by 9 unique shared characters (Fig. 2). The *Rickettsiales* and *Rhodospirillales* formed the deepest branching lineages in these cliques and other groups within α -proteobacteria branched in the following order: (*Sphingomonadales* (*Rhodobacterales* (*Caulobacterales* (*Rhizobiales*)))). This branching order was supported by multiple unique characters at each node giving confidence in the results.

The two main cliques obtained in these analyses differed from each other in terms of the branching position of the species belonging to the order *Rhodospirillales*. In one clique (Fig. 2A), the *Rhodospirillales* branched with the

Rickettsiales, whereas in the other this group of species was found to branch after the *Rickettsiales* and it formed outgroup of the other α -proteobacteria (Fig. 2B). However, only a single character supported the former relationship indicating that it was not reliable. The two families within the *Rhodospirillales* order (*Rhodospirillaceae* and *Acetobacteraceae*), although they branched close to each other, no unique character common to them was identified, indicating that they are highly divergent. The exact branching position of the *Xanthobacter* was also not resolved in these cliques. In some cliques, it appeared as an outgroup of the *Bradyrhizobiaceae* (as shown by the dotted lines in Fig. 2), whereas in others it was placed in the middle of the *Rhizobiaceae* and *Bradyrhizobiaceae* families (as seen for the *Aurantiimonas*).

Phylogenomic analyses of alpha proteobacteria

Table 1 lists some characteristics of various α -proteobacterial genomes that have been sequenced. The genomes vary in size from less than 1 Mb for *Neorickettsia sennetsu* to more than 9.0 Mb for *Bradyrhizobium japonicum*. To identify proteins that are distinguishing features of various higher taxonomic groups within α -proteobacteria, systematic Blastp searches were performed on each ORF in the genomes of *B. japonicum* USDA 110, *Brucella suis* 1330, *Caulobacter crescentus* CB15, *Gluconobacter oxydans* 621H, *Mesorhizobium loti* MAFF303099, *Nitrobacter winogradskyi* Nb-255, *Novosphingobium aromaticivorans* DSM 12444, *Rhodobacter sphaeroides* 2.4.1, *Silicibacter* sp. TM1040, *Rhodospirillum rubrum* ATCC 11170 and *Wolbachia* (*Drosophila melanogaster*) endosymbiont (see Methods section). The genomes chosen covered all main taxonomic groups within the sequenced α -proteobacteria. These analyses have identified large numbers of proteins that are uniquely found in particular groups of α -proteobacteria. A brief description of these results is given below.

Proteins that are distinguishing features of all (or most) alpha proteobacteria

We previously described 6 proteins (viz. CC1365, CC1725, CC1887, CC2102, CC3292 and CC3319) that appeared distinctive characteristics of α -proteobacteria [17]. The α -proteobacterial specificity of these proteins in earlier work was only assessed by means of Blastp searches and it was not confirmed by PSI-Blast, as in the present work (see Methods). Further, since the earlier work, the number of sequenced α -proteobacteria and other genomes has more than doubled. Hence, it was important to confirm the α -proteobacteria specificity of these proteins. Our results reveal that four of these proteins viz. CC1365, CC2102, CC3292 and CC3319, are indeed specific for the α -proteobacteria as a whole, whereas for the remaining two proteins homologs showing significant similarities are also found in other bacteria. Of these four proteins, CC3292 is present in all sequenced α -proteobacteria including *Candidatus Pelagibacter ubique*, which is the smallest known free-living bacterium [2]. The protein CC2102 is only missing in *P. ubique*, while the other two proteins are only missing in 1–2 rickettsiae species (CC1365) and *P. ubique* (CC3319). These proteins, which are uniquely present in virtually all α -proteobacteria, provide distinguishing markers for this Class of bacteria (Fig. 3).

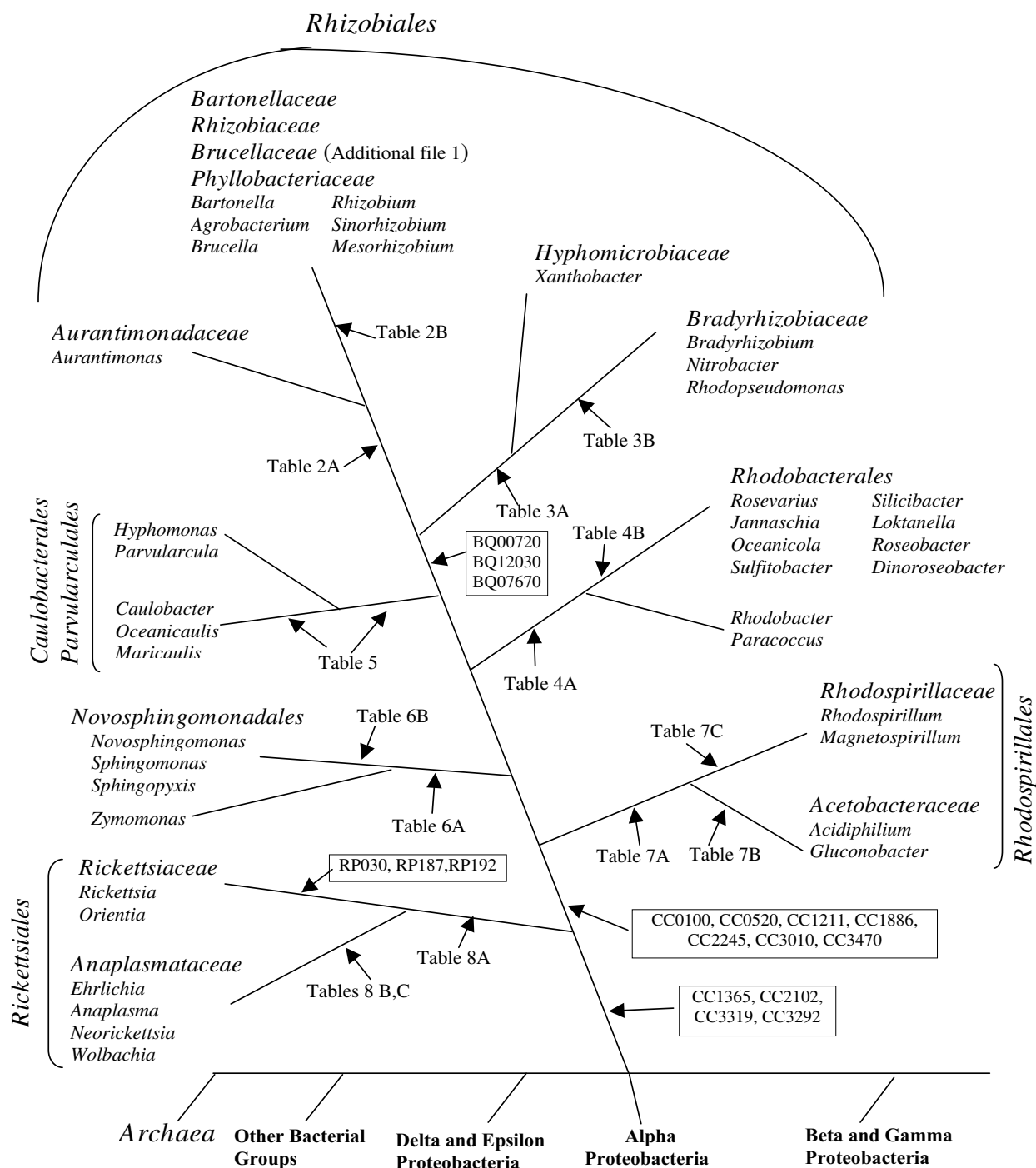
In earlier work, 9 proteins were identified that were present in nearly all α -proteobacteria, except the *Rickettsiales* [17]. These latter species are all intracellular bacteria that have lost many genes that are not required under these conditions [3,34]. The Blastp and PSI-Blast reexamination of these proteins have confirmed that 7 of these proteins (CC0100, CC0520, CC1211, CC1886, CC2245,

CC3010 and CC3470) exhibit the indicated specificity. Except for CC0520 and CC3470, the other five proteins are also found in *P. ubique*, providing evidence for its placement within α -proteobacteria [2]. These results also provide evidence that *P. ubique* is not specifically related to the *Rickettsiales*. The phylogenomic distributions of these genes/proteins can be explained by either their evolution after the divergence of the *Rickettsiales* (Fig. 3), or by gene loss from this lineage.

Proteins that are distinguishing features of the Rhizobiales species

The *Rhizobiales* species comprise more than 1/3rd of the sequenced α -proteobacterial genomes (see Table 1). This order includes a wide assortment of species many of which interact with the eukaryotic organisms to produce diverse effects. This group includes various rhizobia (*Rhizobium*, *Mesorhizobium*, *Sinorhizobium*) and *Bradyrhizobium* species that induce root nodules in plants and live symbiotically within them to enable nitrogen fixation [4-6,35,36]. Another *Rhizobiaceae* species, *A. tumefaciens*, induces Crown gall disease (tumors) in plants [37]. *Bartonella* and *Brucella* species are intracellular pathogens responsible for a number of diseases in human and animals including trench fever and brucellosis [7,38-40]. Other members of this order exhibit enormous versatility in terms of their metabolic capabilities and life styles [1,41,42]. Earlier studies identified six proteins that appeared distinctive characteristics of the *Rhizobiales* species [17]. Reexamination of these proteins by the more stringent criteria used in the present work confirmed that three of these proteins (BQ00720, BQ07670 and BQ12030) are indeed specific for the *Rhizobiales* and they are present in virtually all sequenced species from this large order. In addition to the *Rhizobiales*, these proteins are also present in *Stappia aggregata*, which is presently grouped with the *Rhodobacterales* [12], but was originally known as a strain of *Agrobacterium* [43].

New blast searches on the genomes of *B. japonicum*, *B. suis*, *M. loti* and *N. winogradskyi* have identified large number of other proteins that are specific for different subgroups within the *Rhizobiales* order. Seven proteins listed in Table 2A are uniquely present in most of the sequenced species belonging to the *Rhizobiaceae* (*Rhizobium*, *Sinorhizobium*, *Agrobacterium*), *Brucellaceae*, *Phyllobacteriaceae* (*Mesorhizobium*) and *Aurantimonadaceae* families. The absence of many of these proteins in *Bartonella* species is probably due to gene loss [7,34]. These groups form a well-defined clade with high bootstrap scores in the NJ, MP and ML trees (see Fig. 1) and the genes for them likely evolved in a common ancestor of these genera/families (Fig. 3). Another 9 proteins (Table 2B) are uniquely present in most of the above species except *Aurantimonas*, which forms outgroup of the *Rhizobiaceae*, *Brucellaceae*, *Bartonel-*

**Figure 3**

Summary of the phylogenomic analyses showing the species distribution of various α -proteobacteria-specific proteins and the suggested evolutionary stages where the genes for these proteins have likely evolved. The genes IDs for some proteins described in earlier work are indicated [17]. The information for all other proteins can be found in the indicated Tables or Additional files. A large numbers of conserved indels that are specific for different groups or clades within α -proteobacteria shown here have also been identified in our earlier work [16] (not shown here). The branching order of α -proteobacteria relative to other bacteria has been established in earlier work [32,58].

Table 2: Proteins specific for the Rhizobiaceae and related species

Gene ID	Accession Number	Function	Gene ID	Accession Number	Function
A. Proteins unique to <i>Aurantimonas</i>, <i>Mesorhizobium</i>, <i>Sinorhizobium</i>, <i>Rhizobium</i>, <i>Agrobacterium</i>, <i>Bartonella</i> and <i>Brucella</i>					
mll0062 ¹	<u>NP_101943</u>	Hypothetical	mll0789 ¹	<u>NP_102519</u>	Hypothetical
mll4068 ¹	<u>NP_105027</u>	Hypothetical	mll3016	<u>NP_104217</u>	Omp10
mll7791 ^{1,4}	<u>NP_108034</u>	Hypothetical	mlr6526 ¹	<u>NP_107016</u>	Hypothetical
mlr0777 ¹	<u>NP_102510</u>	Hypothetical			
B. Proteins unique to <i>Mesorhizobium</i>, <i>Sinorhizobium</i>, <i>Rhizobium</i>, <i>Agrobacterium</i>, <i>Bartonella</i> and <i>Brucella</i> (Missing in <i>Aurantimonas</i>)					
mll0122 ^{1,4}	<u>NP_101988</u>	Hypothetical	mll5001 ^{1,4}	<u>NP_105743</u>	Hypothetical
mll1268 ²	<u>NP_102895</u>	Hypothetical	mll8359 ^{1,4}	<u>NP_108472</u>	Hypothetical
mll2847 ^{2,3}	<u>NP_104087</u>	Hypothetical	mlr1823 ¹	<u>NP_103319</u>	Hypothetical
mll2898 ⁴	<u>NP_104130</u>	Hypothetical	mlr0094 ^{1,4}	<u>NP_101965</u>	MhpC (COG0596)
mll4298 ^{1,2}	<u>NP_105201</u>	Hypothetical			
C. Proteins unique to <i>Mesorhizobium</i> and <i>Rhizobiaceae</i> species					
mll0080	<u>NP_101954</u>	Hypothetical	mll6703 ⁴	<u>NP_107159</u>	Hypothetical
mll0867	<u>NP_102577</u>	Hypothetical	mlr1904	<u>NP_103376</u>	Hypothetical
mll9619	<u>NP_109472</u>	Hypothetical	mlr3274	<u>NP_104418</u>	Hypothetical
mlr5174 ⁴	<u>NP_105883</u>	Hypothetical	mlr4951	<u>NP_105704</u>	NodF
mll6303	<u>NP_106835</u>	Hypothetical			
D. Proteins specific to <i>Mesorhizobium</i> and either <i>Rhizobium</i> or <i>Sinorhizobium</i>					
mll0459	<u>NP_102252</u>	Hypothetical	mll2007	<u>NP_103455</u>	Hypothetical
mll1779	<u>NP_103286</u>	Hypothetical	mlr1999	<u>NP_103450</u>	Hypothetical
mll6195	<u>NP_106741</u>	Hypothetical	mlr2029	<u>NP_103476</u>	Hypothetical
mll8758	<u>NP_106740</u>	Hypothetical	mlr6601	<u>NP_107075</u>	Hypothetical
mlr3037	<u>NP_104236</u>	Transcriptional regulator			

¹ Missing in *Bartonella*² Missing in *Agrobacterium*³ Missing in *Rhizobium*.⁴ Also found in *Stappia aggregata*

laceae and *Phyllobacteriaceae* families (Figs. 1 and 2) [29]. Thus, the genes for these proteins have evolved after the branching of *Aurantimonadaceae* (Fig. 3). For six of the proteins in Tables 2A and 2B (marked with *), homologs with low E values are also found in *S. aggregata*, providing further evidence for its relatedness to the *Rhizobiaceae* family. Nine other proteins in Table 2C are found in most *Rhizobiaceae* species and *M. loti*, but they are missing in *Bartonella* and *Brucella* species. Although these proteins suggest a closer relationship between the *Rhizobiaceae* and *Phyllobacteriaceae* families, it is more likely that their genes have been lost from the *Bartonella* and *Brucella* species [34], which are intracellular bacteria. Additionally, some proteins were only found in either *Mesorhizobium* and *Rhizobium*, or *Mesorhizobium* and *Sinorhizobium* (Table 2D). These analyses have also identified 43 proteins that are uniquely present in all four sequenced *Brucella* species and many other proteins that are present in either three or

two of the sequenced *Brucella* species (see Additional file 1).

The analyses of proteins in the genomes of *B. japonicum* and *N. winogradskyi* have identified 12 proteins that are uniquely present in either all (or most) of the sequenced *Bradyrhizobiaceae* species as well as *X. autotrophicus* (Table 3A). The species from these two families form a strongly supported clade in the phylogenetic tree (Fig. 1)[29]. Sixty-two additional proteins in Table 3B are uniquely present in various species belonging to the *Bradyrhizobiaceae* family (i.e. *Bradyrhizobium*, *Nitrobacter*, *Rhodopseudomonas*). Many other proteins (see Additional file 2) are only found in two of the three *Bradyrhizobiaceae* genera and their distributions can result from gene losses, lateral gene transfers (LGTs), or other mechanisms.

Table 3: Proteins that are specific for the Bradyrhizobiaceae group

Gene ID	Accession Number	Function	Gene ID	Accession Number	Function
A. Proteins Unique to Bradyrhizobiaceae Family and Xanthobacter					
bll6014 ^{1,2}	<u>NP_772654</u>	Putative general secretion pathway protein M	Nwi_2179	<u>YP_318785</u>	Hypothetical
Nwi_1093	<u>YP_317707</u>	Hypothetical	Nwi_2432 ¹	<u>YP_319038</u>	Hypothetical
Nwi_1227	<u>YP_317841</u>	Hypothetical	Nwi_2476 ¹	<u>YP_319081</u>	Putative bacterioferritin
Nwi_1786 ¹	<u>YP_318399</u>	Hypothetical	Nwi_2572	<u>YP_319177</u>	Hypothetical
Nwi_1788	<u>YP_318401</u>	Hypothetical	Nwi_2623	<u>YP_319228</u>	Hypothetical
Nwi_2147 ^{1,3}	<u>YP_318753</u>	Hypothetical	Nwi_2707	<u>YP_319312</u>	Hypothetical
B. Proteins Unique to Bradyrhizobiaceae Family					
bll5899 ²	<u>NP_772539</u>	Hypothetical	Nwi_2021	<u>YP_318632</u>	Hypothetical
blr6106 ^{1,2}	<u>NP_772746</u>	Hypothetical	Nwi_2063	<u>YP_318673</u>	Hypothetical
Nwi_0278	<u>YP_316897</u>	Hypothetical	Nwi_2064	<u>YP_318674</u>	Hypothetical
Nwi_0503	<u>YP_317122</u>	Hypothetical	Nwi_2163	<u>YP_318769</u>	Hypothetical
Nwi_0528	<u>YP_317147</u>	Hypothetical	Nwi_2173	<u>YP_318779</u>	Hypothetical
Nwi_0605 ¹	<u>YP_317224</u>	Hypothetical	Nwi_2183 ¹	<u>YP_318789</u>	Hypothetical
Nwi_0710 ^{1,d}	<u>YP_317328</u>	Hypothetical	Nwi_2208 ³	<u>YP_318814</u>	Hypothetical
Nwi_0925	<u>YP_317539</u>	Hypothetical	Nwi_2244	<u>YP_318850</u>	Hypothetical
Nwi_0966 ^{1,d}	<u>YP_317580</u>	Hypothetical	Nwi_2247	<u>YP_318853</u>	Hypothetical
Nwi_1084	<u>YP_317698</u>	Hypothetical	Nwi_2379	<u>YP_318985</u>	Hypothetical
Nwi_1092	<u>YP_317706</u>	Hypothetical	Nwi_2381 ²	<u>YP_318987</u>	Hypothetical
Nwi_1107 ³	<u>YP_317721</u>	Hypothetical	Nwi_2414 ³	<u>YP_319020</u>	Hypothetical
Nwi_1108	<u>YP_317722</u>	Hypothetical	Nwi_2489	<u>YP_319094</u>	Hypothetical
Nwi_1336	<u>YP_317949</u>	Hypothetical	Nwi_2492 ^{1,3}	<u>YP_319097</u>	Hypothetical
Nwi_1139	<u>YP_317753</u>	Hypothetical	Nwi_2500	<u>YP_319105</u>	Hypothetical
Nwi_1247 ³	<u>YP_317861</u>	Hypothetical	Nwi_2506 ³	<u>YP_319111</u>	Hypothetical
Nwi_1270	<u>YP_317883</u>	Hypothetical	Nwi_2509 ³	<u>YP_319114</u>	Hypothetical
Nwi_1275 ³	<u>YP_317888</u>	Hypothetical	Nwi_2531	<u>YP_319136</u>	Hypothetical
Nwi_1454 ¹	<u>YP_318067</u>	Hypothetical	Nwi_2575	<u>YP_319180</u>	Hypothetical
Nwi_1498	<u>YP_318111</u>	Hypothetical	Nwi_2577	<u>YP_319182</u>	Hypothetical
Nwi_1512	<u>YP_318125</u>	Hypothetical	Nwi_2588 ^{1,3}	<u>YP_319193</u>	Hypothetical
Nwi_1581 ³	<u>YP_318194</u>	Hypothetical	Nwi_2630	<u>YP_319235</u>	Hypothetical
Nwi_1582	<u>YP_318195</u>	Hypothetical	Nwi_2676	<u>YP_319281</u>	Hypothetical
Nwi_1586	<u>YP_318199</u>	Dihydrofolate reductase	Nwi_2677 ³	<u>YP_319282</u>	Hypothetical
Nwi_1649 ^{1,3}	<u>YP_318262</u>	Hypothetical	Nwi_2769	<u>YP_319374</u>	Hypothetical
Nwi_1674	<u>YP_318287</u>	Hypothetical	Nwi_2789 ¹	<u>YP_319394</u>	Hypothetical
Nwi_1705	<u>YP_318318</u>	Hypothetical	Nwi_2984 ³	<u>YP_319586</u>	Hypothetical
Nwi_1711	<u>YP_318324</u>	Hypothetical	Nwi_2959	<u>YP_319561</u>	Hypothetical
Nwi_1785	<u>YP_318398</u>	Hypothetical	Nwi_3035	<u>YP_319637</u>	Hypothetical
Nwi_1793	<u>YP_318406</u>	Hypothetical	Nwi_3140	<u>YP_319739</u>	Hypothetical
Nwi_1800 ^{1,2}	<u>YP_318413</u>	Hypothetical	Nwi_3141 ¹	<u>YP_319740</u>	Hypothetical

¹ Missing in one or more strains of Rhodopseudomonas² Missing in one or more species of Nitrobacter³ Missing in Bradyrhizobium japonicum USDA 110

Proteins that are distinguishing features of the *Rhodobacterales* species

The order *Rhodobacterales* is a heterogeneous lineage of bacteria that exhibit much diversity in terms of their metabolism and cell division cycles [1,13]. This group includes many photosynthetic bacteria that are capable of CO₂ as well as nitrogen fixation and also many chemоорganotrophs that can metabolize various sulfur-containing

compounds [44,45]. A number of budding, stalk forming and prosthecate bacteria also belong to this group [46]. In addition to many completely sequenced genomes (see Table 1), information for several other species belonging to this order (e.g. *Sulfitibacter*, *Oceanicola*, *Loktanella*, *Jannaschia*, *Dinoroseobacter*, *Roseovarius* and *Sagittula*) is available in the NCBI database. To identify proteins that are specific for the *Rhodobacterales*, phylogenomic analyses

were carried out on various ORFs in the genomes of *R. sphaeroides* 2.4.1 and *Silicibacter sp. TM1040*.

These studies have identified 29 proteins that are present in all-available *Rhodobacterales* species (Table 4A), but these proteins as well as those listed in Table 4B and 4C are not found in *H. neptunium*, *Oceanicaulis alexandrii*, *Maricaulis maris* or *Stappia aggregata*. These latter species are presently grouped with the *Rhodobacterales* [12], how-

ever, the absence of various *Rhodobacterales*-specific proteins in them and phylogenetic studies indicate that the placement of these species within this order is incorrect and needs be revised. In phylogenetic trees based on concatenated sequences for many proteins, *O. alexandrii* and *M. maris* consistently branched with the *Caulobacter* rather than the well-defined clade of *Rhodobacterales* (Figs. 1 and 2) [29]. The studies by Badger et al. [47] also provide

Table 4: Proteins that are specific for the Rhodobacterales

Gene ID	Accession Number	Function	Gene ID	Accession Number	Function
A. Proteins specific for Rhodobacterales (<i>Oceanicola</i>, <i>Loktanella</i>, <i>Paracoccus</i>, <i>Roseovarius</i>, <i>Roseobacter</i>, <i>Jannaschia</i>, <i>Silicibacter</i>, <i>Sulfitobacter</i>, <i>Dinoroseobacter</i>, <i>Sagittula</i>)					
TM1040_0093 ¹	YP_612088	Hypothetical	TM1040_1842	YP_613837	Phasin, PhaP
TM1040_0184 ²	YP_612179	Hypothetical	TM1040_1967 ³	YP_613961	Putative CheA signal transduction
TM1040_0236 ^{1,2}	YP_612231	Hypothetical	TM1040_1988	YP_613982	Hypothetical
TM1040_0471	YP_612466	Putative rod shape-determining protein MreD	TM1040_2263 ²	YP_614257	Hypothetical
TM1040_0586 ²	YP_612581	Hypothetical	TM1040_2370	YP_614364	Hypothetical
TM1040_0587 ²	YP_612582	Hypothetical	TM1040_2425 ³	YP_614419	Hypothetical
TM1040_0697	YP_612692	Hypothetical	TM1040_2466	YP_614460	GCN5-related N-acetyltransferase COG045
TM1040_0750 ^{2,4}	YP_612745	Hypothetical	TM1040_2487 ²	YP_614481	Hypothetical
TM1040_0752 ¹	YP_612747	Lipoprotein, putative	TM1040_2582 ³	YP_614576	Hypothetical
TM1040_1063	YP_613058	Gene transfer agent	TM1040_2999	YP_614993	Hypothetical
TM1040_1064	YP_613059	Gene transfer agent	TM1040_3077 ³	YP_611313	Hypothetical
TM1040_1247 ³	YP_613242	Hypothetical	TM1040_3749	YP_611978	Hypothetical
TM1040_1350 ²	YP_613345	Hypothetical	TM1040_3759	YP_611988	Lipoprotein, putative
TM1040_1406	YP_613401	Outer membrane chaperone Skp (OmpH)	TM1040_3764 ³	YP_611993	Putative transmembrane protein
TM1040_1567	YP_613562	Hypothetical			
B. Proteins unique to various Rhodobacterales but missing in <i>Rhodobacter</i> and <i>Paracoccus</i>					
TM1040_1558 ¹	YP_613553	Hypothetical	TM1040_2157 ⁴	YP_614151	Hypothetical
TM1040_1735 ¹	YP_613730	Hypothetical	TM1040_2443 ^{1,4}	YP_614437	Lipolytic enzyme, G-D-S-L
TM1040_1844 ^{1,5,6}	YP_613839	Hypothetical	TM1040_2680 ⁷	YP_614674	Hypothetical
C. Proteins Unique to <i>Silicibacter</i> and <i>Roseobacter</i>					
TM1040_1099 ⁸	YP_613094	Hypothetical	TM1040_3189	YP_611425	
TM1040_1423 ⁸	YP_613418	Hypothetical	TM1040_3202 ⁸	YP_611438	
TM1040_1451	YP_613446	Hypothetical	TM1040_3208 ⁸	YP_611444	
TM1040_1986 ⁸	YP_613980	Hypothetical	TM1040_3226 ⁸	YP_611462	
TM1040_2106	YP_614100	Hypothetical	TM1040_3529 ⁸	YP_611763	
TM1040_2139 ⁸	YP_614133	Hypothetical	TM1040_3626 ⁸	YP_611855	
TM1040_3075 ⁸	YP_611311	Hypothetical			

¹ Missing in *Loktanella vestfoldensis* SKA53

² Missing in one or more *Rhodobacter sphaeroides* strains

³ Missing in *Paracoccus denitrificans* PD12222

⁴ Missing in *Rhodobacterales bacterium* HTCC2654

⁵ Missing in one or more species of *Roseovarius*

⁶ Missing in *Oceanicola batsensis* HTCC2597

⁷ Missing in one or more species of *Roseobacter*

⁸ Missing in *Silicibacter pomeroyi* DSS-3

strong evidence for the grouping of *H. neptunium* with the *Caulobacterales*.

Six additional proteins in Table 4B are present in most of the *Rhodobacterales* species, but they are missing in *R. sphaeroides* and *P. denitrificans*, which form a distinct clade that appears as the outgroup of other *Rhodobacterales* species (Fig. 1)[29]. Thus, the genes for these proteins have likely evolved after the branching of these two genera. Thirteen additional proteins, which are only found in *Silicibacter* and *Roseobacter* genera (Table 4C) support a close relationship among them, as seen in phylogenetic trees (Fig. 1).

Of the proteins that are specific for *Rhodobacterales*, two of them (YP_613058 and YP_613059 in Table 4A, corresponding to proteins ABK27256 and ABK27255 in *R. capsulatus* genome) were previously identified as part of a complex referred to as gene transfer agent [48], based on similarity to certain virus-like elements. Another protein in the same category (viz. ABK27253) is specific for the *Rhodobacter* genus. The significance of these results is presently unclear.

Proteins that are distinctive characteristics of the *Caulobacterales*

The order *Caulobacterales* is comprised of a single family with only four genera [12,49]. These chemoorganotrophic bacteria are commonly found in marine aerobic environments and they are distinguished by their ability to form stalked cells and unusual cell division cycle [49,50]. The complete genome of only *C. crescentus*, which is the best-studied organism from this group, is presently available [51]. However, as discussed above, a number of other species which are presently classified as *Rhodobacterales* viz. *O. alexandrii*, *M. maris*, *H. neptunium* and also *Parvularcula bermudensis*, consistently branch with *C. crescentus* in different phylogenetic trees (Figs. 1 and 2) [29,47]. Thus, phylogenomic analysis of the ORFs from *C. crescentus* genome was of much interest.

These analyses have identified 2 proteins (CC0486 and CC2480), which are uniquely present in this species as well as *O. alexandrii*, *M. maris*, *H. neptunium* and *P. bermudensis* (Table 5). One additional protein, CC2764 is present in all of these species except *H. neptunium*. The remaining eight proteins in Table 5 are only found in *C. crescentus*, *O. alexandrii* and *M. maris*, indicating that the latter two species are more closely related to *C. crescentus* in comparison to either *H. neptunium* or *P. bermudensis*. These results are strongly supported by the branching patterns of these species in phylogenetic trees (Figs. 1 and 2) [29,47]. Previously, Badger et al. [46] have reported identification of 62 proteins that were only found in *C. crescentus* and *H. neptunium*. However, the blast threshold used in this study to infer the absence of these proteins in other species was very high i.e. 1e-10. By the criteria used in the present work (see Methods), none of these 62 proteins was found to be unique to these two species.

Proteins that are distinguishing characteristics of the *Sphingomonadales*

The species belonging to the order *Sphingomonadales* are present in both terrestrial and aquatic environments [52]. A distinguishing characteristic of many species from this group is the presence of glycosphingolipids in their cell envelope rather than lipopolysaccharides [52,53]. Several species from this group (e.g. *N. aromaticivorans*) can degrade a wide variety of aromatic hydrocarbons [52], whereas others such as *Zymomonas mobilis*, can highly efficiently ferment sugar to ethanol [53], making them of much interest and importance from biotechnological standpoints. This group also includes phototrophic organisms (e.g. *Erythrobacter litoralis*), which contain bacteriochlorophyll *a* and can derive significant fraction of their metabolic energy via anaerobic photosynthesis [54]. The complete genomes of 5 species from this order are now available (see Table 1). In addition, large numbers of sequences for *Sphingomonas sp. SKA58*, are also available in the NCBI database. Blast searches on the ORFs in the genome of *N. aromaticivorans* have identified 16 proteins

Table 5: Proteins specific for the Caulobacter and related species

Proteins Unique to Caulobacter, Oceanicaulis and Maricaulis (Some also found in Hyphomonas and Parvularcula)					
Gene ID	Accession Number	Function	Gene ID	Accession Number	Function
CC0486 ¹	<u>NP_419305</u>	Hypothetical	CC1066	<u>NP_419882</u>	Hypothetical
CC2480 ¹	<u>NP_421283</u>	Hypothetical	CC1586	<u>NP_420397</u>	Hypothetical
CC2764 ²	<u>NP_421560</u>	Hypothetical	CC2207	<u>NP_421010</u>	Hypothetical
CC3101	<u>NP_421895</u>	Hypothetical	CC2628	<u>NP_421428</u>	hfaA protein
CC0512	<u>NP_419331</u>	Hypothetical	CC2639	<u>NP_421438</u>	Hypothetical
CC1064	<u>NP_419880</u>	Hypothetical			

All of the proteins in this Table are present in *C. crescentus* as well as in *O. alexandrii* and *M. maris*.

¹ These proteins are also present in *H. neptunium* and *P. bermudensis*.

² Also found in *P. bermudensis*.

(Table 6A) that are uniquely present in all 6 of the *Sphingomonadales* species for which information is available. Thirteen additional proteins (Table 6B) are present in all other *Sphingomonadales* species, except *Z. mobilis*, which is the deepest branching species within this group (Fig. 1). The genes for these proteins likely evolved after the branching of *Z. mobilis*. Many other proteins are present in only 3 or 4 of these species (viz. *N. aromaticivorans*, *E. litoralis*, *S. alaskensis*, *S. wittichii* and *Sphingomonas sp. SKA58*) (see Additional file 3) and their phylogenomic distribution can result from a variety of mechanisms including shared ancestry, gene loss and LGTs among these species.

We have also identified a 4 aa insert in a highly conserved region of Gyrase B that is mainly specific for the species from this order (Fig. 4). This indel is present in all available *Sphingomonadales* species, but it is not found in most other alpha proteobacteria or other bacteria (results not shown). Besides *Sphingomonadales*, a similar size indel is also present in three other species (viz. *C. leidya*, *R. blasticus*, and *Pseudorhodobacter incheonensis*). Because other *Rhodobacterales* or *Caulobacter* species lack this insert, the presence of this indel in these three species could be either due to LGTs or possibly due to taxonomic misclassification of these species.

Proteins and indels that are specific for the *Rhodospirillales* species

The order *Rhodospirillales* is comprised of diverse species including some photosynthetic and magnetotactic bacteria, some acidophiles as well as other bacteria commonly associated with flowers, fruits and fermented beverages that are involved in the partial oxidation of carbohydrates and alcohols [55]. The order is made up of two main families, *Rhodospirillaceae* and *Acetobacteraceae* [12,55]. The complete genomes of four species, two from each family, *Rhodospirillaceae* (*Magnetospirillum magnicum*, *R. rubrum*) and *Acetobacteraceae* (*Acidiphilum cryptum* and *G. oxydans*), are available (Table 1) [56,57]. Phylogenomic analyses of various ORFs in the genomes of *G. oxydans* and *R. rubrum* have led to identification of one protein, GOX0963, which is uniquely found in all of these species (Table 7A). Three other proteins in this Table are present in at least 3 of the 4 species from this order. This table also lists 14 proteins each that are distinctive characteristics of either the *Acetobacteraceae* (Table 7B) or the *Rhodospirillaceae* (Table 7C) families, providing molecular markers for these families. We have also identified a 25 aa insert in a conserved region of the RNA polymerase beta subunit (RpoB) that is unique to various sequenced *Rhodospirillales* species, but not found in any other bacteria (Fig. 5).

Table 6: Proteins that are specific for the *Sphingomonadales* group of species

Gene ID	Accession Number	Function	Gene ID	Accession Number	Function
A. Proteins Unique to <i>Sphingomonadales</i> Order (Novosphingobium, Erythrobacter, Sphingomonas, Sphingopyxis and Zymomonas)					
Saro_0018	YP_495301	Hypothetical	Saro_1291	YP_496569	Hypothetical
Saro_0052 ¹	YP_495335	Hypothetical	Saro_1378	YP_496656	Hypothetical
Saro_0087	YP_495370	Hypothetical	Saro_1914 ¹	YP_497188	Hypothetical
Saro_0150	YP_495433	Hypothetical	Saro_2130	YP_497403	Hypothetical
Saro_0232	YP_495514	Hypothetical	Saro_2788	YP_498058	Hypothetical
Saro_0409	YP_495691	Hypothetical	Saro_2958	YP_498227	Hypothetical
Saro_1088	YP_496367	Hypothetical	Saro_3138	YP_498407	Hypothetical
Saro_1144	YP_496423	Hypothetical	Saro_3213	YP_498482	Hypothetical
B. Proteins Unique to <i>Sphingomonadales</i> but missing in Zymomonas					
Saro_0044	YP_495327	Hypothetical	Saro_1748 ²	YP_497022	
Saro_0154	YP_495437	Hypothetical	Saro_1785	YP_497059	
Saro_0415 ³	YP_495697	Hypothetical	Saro_1972	YP_497246	
Saro_0458	YP_495740	Hypothetical	Saro_2036	YP_497309	
Saro_1078	YP_496357	Hypothetical	Saro_2037	YP_497310	
Saro_1126	YP_496405	Hypothetical	Saro_2333	YP_497604	
Saro_1160	YP_496439	Hypothetical	Saro_2548	YP_497818	
Saro_1163	YP_496442	Hypothetical			

¹ Missing in *Sphingomonas wittchii*

² Blast scores for *C. crescentus* and *H. neptunium* are also significant

³ Significant blast scores for *C. crescentus*

		95		140
	Novo. aromaticivorans	87201322	EGVSAAEVIMTQLHAGGKF	
	Sphingopyxis alaskensis	103488449	ENTS	DDNAYKVSGGLHGVGVSVNALS
	Erythrobacter litoralis	85375173		-----
	Erythrobacter sp. NAP1	85710072		-----
	Sphingomonas sp. SKA58	94496546		-----
	Erythrobacter sp.	19909663	D--N	-A-----
	Sphingomonas wittichii	118761537		-----
	Erythrobacter sp.	19909665		-----
	Zymomonas mobilis	6580765	D-NA	NS-----
<i>Sphingomonadales</i>	Caulobacter leidyi	19909581		-----
	Rhodobacter blasticus	18157440	N---DD	SG-----
	Pseudorhod. incheonensis	90903296	N---DE	GG-----
<i>Other Alpha with inserts</i>	Brucella suis	23501039		-Q-S-----
	Brucella melitensis	17988106		-Q-S-----
	Mesorhizobium sp. BNC1	110635929		-Q-S-----
	Bartonella quintana	49473721	--I-----	-Q-S-----
	Agrobacterium tumefaciens	15887371		-Q-S-----
	Rhizobium leguminosarum	116249778		-Q-S-----
	Mesorhizobium loti	13474326	--I-----	-Q-S-----
	Bradyrhizobium sp. BTAl	78694144		-Q-S-----
	Nitrobacter hamburgensis	92115637	--I-----	-Q-S-----
	Nitrobacter winogradskyi	75674203		-Q-S-----
	Xantho. autotrophicus	89360513		NQ-S-----
	Rhodopseudomonas palustris	91974486	--I-----	-Q-S-----
	Bradyrhizobium japonicum	27375934	--I-----	-Q-S-----
<i>Rhizobiales</i>	Caulobacter sp. K31	113933904		-Q-S-----
	Caulobacter fusiformis	19909539		-Q-S-----
	Caulobacter crescentus	16124415		-Q-S-----
	Oceanicaulis alexandri	83945027	--Q	-Q-S-----
	Parvularcula bermudensis	84701583	--Q-----	-Q-S-----
	Rhodospirillum rubrum	83591344		-Q-S-----
	Mag. magnetotacticum	46200851	--I-----AL-K-----	EQ-SNRI-S-----I-----
	Roseovarius nubinhibens	83950587		-S-S-----
	Oceanicola batsensis	84501255	--I-----	-Q-S-----I-----
	Rhodobacteres bacterium	84684515		-S-S-----
	Stappia aggregata	118590457		-Q-S-----
	Roseobacter sp. MED193	86139413		-S-S-----
	Sulfitobacter sp. NAS-14.1	83855146		-S-S-----
	Roseobacter denitrificans	110677620		-S-S-----
	Paracoccus denitrificans	69935301		-Q-S-----
	Jannaschia sp. CCS1	89052495		-S-S-----
	Silicibacter pomeroyi	56695080		-S-S-----
	Silicibacter sp. TM1040	99079848		-S-S-----
	Rhodobacter sphaeroides	77464924		-Q-S-----
	Wolbachia endo. (Dros)	42520018	--I-----	-S-T-----I-----
	Wolbachia pipiens	67809845	--I-----	-S-T-----I-----
	Ehrlichia chaffeensis	68171653	--M-----	-H-S-----I-----
	Ehrlichia canis	73667042	--M-----	-HSS-----I-----
	Anaplasma phagocytophilum	88607620	--I-----	-H-S-----
	Anaplasma marginale	56416812		-H-S-----
	Rickettsia conorii	15892807	--I-----	-Q-----A-----
	Rickettsia rickettsii	53732213	--I-----	-Q-----A-----
	Rickettsia prowazekii	15604433	--I-----	-QQS--I-----
<i>Caulobacterales</i>				
<i>Rhodospirillales</i>				
<i>Rhodobacterales</i>				
<i>Rickettsiales</i>				

Figure 4

Partial sequence alignments of DNA Gyrase B showing a 4 aa insert that is mainly specific for the *Sphingomonadales* species. A 4–5 aa insert present in some other α-proteobacteria could be due to either LGTs or taxonomic anomalies. The dashes (-) denote identity with the amino acid on the top line. Sequence information for other groups of bacteria (which do not contain this insert) is not shown.

Proteins that are specific for the Rickettsiales

The *Rickettsiales* species are intracellular pathogens responsible for a number of diseases in humans and other animals [3,34]. This order is comprised of three families, *Rickettsiaceae*, *Anaplasmataceae* and *Holosporaceae*. All of the sequenced genomes are from the first two families. Phylogenomic analysis of various ORFs from the genome of *Wolbachia* (*D. melanogaster*) endosymbiont has identified 3 proteins viz. WD0161, WD0715 and WD0771 (Table 8A) that are specific for the entire *Rickettsiales*

order. Five other proteins in Table 8B (viz. WD0083, WD0157, WD0821, WD0827 and WD0863) are present in all sequenced *Ehrlichia*, *Anaplasma*, *Wolbachia* and *Neorickettsia* species (belonging to the *Anaplasmataceae* family), but they are not found in any of the *Rickettsiaceae* or other bacteria. Ten additional proteins (Table 8C) are uniquely present in the *Ehrlichia*, *Anaplasma* and *Wolbachia* species, but are absent in *Neorickettsia*. In view of the deep branching of *Neorickettsia* in comparison to these other genera (Fig. 1), the genes for these proteins have

Table 7: Proteins that are specific for the Rhodospirillales group

Gene ID	Accession Number	Function	Gene ID	Accession Number	Function
A. Proteins Unique to Rhodospirillales Order (<i>Gluconobacter</i>, <i>Magnetospirillum</i>, <i>Rhodospirillum</i> and <i>Acidiphilium</i>)					
GOX0633 ¹	AAW60410	Hypothetical	GOX0963	AAW60735	Hypothetical
GOX0695 ²	AAW60472	Hypothetical	GOX1258 ³	AAW61019	Hypothetical
B. Proteins Unique to Acetobacteraceae Family (<i>Gluconobacter</i> and <i>Acidiphilium</i>)					
GOX0143	AAW59936	Hypothetical	GOX1616	AAW61357	Hypothetical
GOX0343	AAW60126	ANK, ankyrin repeats	GOX2216	AAW61951	Hypothetical
GOX1212	AAW60973	Phage portal protein	GOX2275	AAW62008	Hypothetical
GOX1215	AAW60976	Putative phage protein	GOX2316	AAW62049	Hypothetical
GOX1222	AAW60983	Putative phage protein	GOX2452	AAW62183	Hypothetical
GOX1224	AAW60985	Putative phage protein	GOX2454	AAW62185	Hypothetical
GOX1233	AAW60994	Hypothetical	GOX2456	AAW62187	Hypothetical
C. Proteins Unique to Rhodospirillaceae Family (<i>Rhodospirillum</i> and <i>Magnetospirillum</i>)					
Rru_A0125	YP_425217	Putative diguanylate phosphodiesterase	Rru_A2592	YP_427676	Hypothetical
Rru_A0152	YP_425244	Hypothetical	Rru_A2828	YP_427912	Hypothetical
Rru_A0531	YP_425622	Hypothetical	Rru_A3562	YP_428643	Hypothetical
Rru_A1689	YP_426776	Hypothetical	Rru_A3636	YP_428717	Hypothetical
Rru_A1756	YP_426843	Hypothetical	Rru_A3662	YP_428743	Hypothetical
Rru_A2112	YP_427199	Hypothetical	Rru_A3739	YP_428820	Hypothetical
Rru_A2510	YP_427597	Predicted transcriptional regulator	Rru_A3800	YP_428881	Hypothetical

¹ Missing in *Rhodospirillum*² Missing in *Acidiphilium*³ Missing in *Magnetospirillum*

likely evolved after the branching of *Neorickettsia*. In earlier work, a number of proteins that appeared specific for the *Rickettsiaceae* family were also identified [17]. Additional Blastp and PSI-Blast searches on these proteins confirm that three of these proteins viz. RP030, RP187 are RP192, are indeed specific for the *Rickettsiaceae* family. Of these, RP030 is also found in *Orientia tsutsugamushi*.

Conclusion

In this work, we have used a combined phylogenetic and phylogenomic approach to examine the evolutionary relationships among α -proteobacteria. Our analyses have identified large numbers of genes/proteins that are uniquely found in α -proteobacteria at various phylogenetic depths (Fig. 3). These include several proteins that are distinctive characteristics of all α -proteobacteria, as well as many proteins that constitute the unique repertoires of either all of the main orders of α -proteobacteria (viz. *Rhizobiales*, *Rhodobacterales*, *Rhodospirillales*, *Rickettsiales*, *Sphingomonadales* and *Caulobacterales*) or its different families (viz. *Rickettsiaceae*, *Anaplasmataceae*, *Rhodospirillaceae*, *Acetobacteraceae*, *Bradyrhizobiaceae*, *Bryellaceae* and *Bartonellaceae*). In addition, numerous other

α -proteobacteria-specific proteins are present at different phylogenetic depths and they provide important information regarding the evolution of these bacteria. This work also describes two novel conserved indels in important housekeeping genes (viz. Gyrase B and RpoB) that are distinctive characteristics of the *Sphingomonadales* and *Rhodospirillales* orders, respectively. These indels are in addition to many other α -proteobacteria-specific indels that have been described in earlier work [16,58].

Based upon these α -proteobacteria-specific proteins and conserved indels, it is now possible to define nearly all of the higher taxonomic groups (i.e. most orders and many families) within α -proteobacteria in clear and definitive molecular terms based upon multiple characteristics (Fig. 3). The species distribution profiles of these α -proteobacteria-specific proteins and indels also provide important information regarding their branching order and interrelationships, which are highly concordant with each other (c.f. Fig. 26 in ref. [16] with Fig. 3 in this work). Importantly, the relationships that emerge from these phylogenomic analyses (Fig. 3) are in excellent agreement with the branching patterns of these species in different phylo-

		203	RIDRRRKLPPVTTLLYALEGAAATLREARTAEGRSVPDPSEIAGMTSE	EILDFYYQKLSYKRDDKG WKVDFKPDLHRLGVKLMFDLVDAATGE
Rhodospirillales	Acetobacteraceae	83594027	-----D-LNTAQ--V-A---GLEQ---K-----	---SY-G-VV-T-GB---TP-DAER-K---VS-I-K---
		46202801	-----QRTEA-IASTDPQ-LP-H-A-A-	---SY-TQVVFSTP---ARP-D-EAF---IT-AEA-I-D-T
		114327212	-----NYL-Q---QKI---GD-EGLD-R-DQD	---SY-EAVFPL-LGGE-ARP-D-AF-L---LSP----D-
		58038861	-----K-----R-T	---TY-G-VPFT---GE-A-ARP-EPETF---A---VE---E---
		88938867	-----E-A-E-K-E-LEIG-R-DA-	---RT-GUVT-SKEK---TA-DGARM---DI-I-K-K
		Micavibrio sp. EPB	-----R-DSSMTADY-AKCEENDQF---LMVQ-SN-	---ST-NRVT-E-QAD -R-P-DAER-K-F-ASV-I-DS-
		77921157	-----K-----S-F-GDGE	---T-T-SL-Q---GE- -R-P-O-A-K-Q-TLA-MI-D---
		23006310	-----I-S-M-GMDGE	---ST-N-IT-V-AGDH-RIP-NVERF---I-AVG---D-
		Rhodospseudomonas palustris	-----I-S-M-GMDGE	D---ST-N-VT-E-GD- -RIPYSV-RFK-M-VS-I-DS-
		39936331	-----S-MF-GDGE	---ST-KTVT-T-GDN -RIPYSAERFK-M-IIS---D---
		Bradyrhizobium japonicum	-----I-S-MF-GDGE	---ST-N-IL---TKE- -R-P-DVNR-E-YSTMN-I-D-K
		36333357	-----I-S-MF-GDGE	A---ST-K-IL---TKE- -R-P-DANRF-YSTMN-I-D-K
		Mesorhizobium loti	-----I-S-M-GMDGE	---ST-E-IFPT-EKG- -RMP-D-KRMK-Y-AVA-I-D-
		13470542	-----S-M-GMDGE	---ST-K-IQ---IKD- -R-P-VA-RF-YSTMN-I-DS-K
		Agrobacterium tumefaciens	-----I-S-M-GMD-S	---NH-DR-VFE-AEN- ---P-MVENA-S-PA-V-K---
		15889250	-----I-S-M-GMD-S	---H---NTVITW-AAGEG-IP-E-EAV-Q-PT-A---K-
		Bartonella henselae	-----I-S-M-GMD-S	Q---HH-N-ITTFV-G-G- ---PYVAESW-R-PF---IINGD-
		Bru. melitensis	-----A-S-M-GMDGE	---AQ---N-VVFL-GEGL- ---QIPYVAEAW-M-PA---I-KS-
		Rhodopseudomonas palustris	-----S-MF-GDGE	D---HH-DTIVTA-EAG- ---P-VLEQW-S-PA---I-KS-
		39936331	-----I-S-MF-GDGE	---TT-DVVFPEKRSG- -ATPY-ERW---PE-P---D-
		Xanthobacter autotrophicus	-----I-S-M-GDNE	---ST-GRVT-EAGKD- -TMP-VKERW-I-PQR-I-KS-
		89358263	-----I-S-M-GDNE	D---CL-SVTHDAVEG- FSTT-D-EK-K-T-SER---
		Afipia clevelandensis	-----I-S-M-GDNE	A-MNAV-NTVT-RFEKNKG-VTK-F-ERV- -TRFA-
		36333343	-----I-S-M-GDNE	---ST-N-NTVITKVKQ- -R-P-NA-RV-A-PER---I-K-K
		Erythrobacter litoralis	-----I-S-M-GMTGE	A-M-AY-DTII- -LEKNKG-VAP-F-RV- -TRPTY---S-
		103486971	-----K-----GMTGE	G-M-AY-ETVNPHQKNRG-VTR-P-ERV- -TRPTY---
		Zymomonas mobilis	-----K-----GLDSE	A-MNAV-NTVN-TLK-GEG-VAP-P-RV- -TRPSY---
		56551627	-----K-----GLN-E	G-M-AY-DTVEFL-LKKNKG-VTK-P-ERV- -TRET---N-
		Sphingomonas sp. SKA58	-----K-----GLN-E	G-M-AY-PEKTV- -RLEK-RG-VTP-P-ERV- -TRPTY- -I-S-
		Novo. Aromaticivorans	-----K-----GLDSE	G-M-AY-ETVD- -LRKGQG-VTK-P-ERV- -TRPTY---DS-
		87198050	-----K-----GLDSE	G-M-AY-NTVN-LEKSRG-VTP-P-ERV- -TRPTY---
		Caulobacter crescentii	-----A-F---GMDGE	G-M-AY-ATVTF- -MVRNNG-STK-F-ERV- -TRFTT---K-
		16124757	-----A-F---GMDGE	Q---Q---TR-SL-R-K- -TS-RA-AWK---PE-E-I---K-K
		Parvularcula bermudensis	-----A-G-YQDE	---ST-DTIVT- -ASK- -T-PYVAKERW---PAR---K-D
		84703909	-----A-G-YQDE	D-M-AY-DTIVT- -THRKRGZ-VTR-P-ERI- -TRFTQ-V---
		Parvibaculum lavamentivorans	-----A-G-YQDE	D---R-E-E-IK-VKHKD- ---P-V-KFK- -R-P---M-
		121525595	-----A-G-YQDE	D---A-D-VI-T-C--- -R-P-IV-REFK- -R-SY- -M-
		Roseobacter denitrificans	-----S-GLDQE	D---T-D-IR-VKSE- -V-P-VA-RF---R-SH- -M-S
		110681145	-----S-GLDQE	D-FAQ-CEVSECRLTKD- K-T-C-V-EKF- -R-QY- -IN-E---
		Rhodobacter sphaeroides	-----S-GLDQE	--IK-Y-NSVT- -FVKNNKG-S-K-1-Q-ITAHR-TS---D-N
		Sulfurobacter sp. NAS-14.1	-----S-GLDQE	--MK-Y-NSVT- -CIKNKG-S-K-1-Q-ITAHR-TS---D-N
		Roseovarius nubinhibens	-----S-GLDQE	--IK-Y-NSVT- -LVKNNKG-A-K-1-Q-ITAHR-TS---D-N
		Silicibacter pomeroyi	-----S-GLDQE	--MK-Y-NSVT- -CIKNKG-S-K-1-Q-ITAHR-TS---D-N
		Oceanicola batisensis	-----S-GLDQE	
		Silicibacter sp. TM1040	-----S-GLDQE	
		Jannaschia sp. CCS1	-----S-GLDQE	
		89053059	-----S-GLDQE	
		Hypomonas neptunum	-----S-GLDQE	
		114799111	-----S-GLDQE	
		Maricaulicus maris MCS10	-----S-GLDQE	
		114570352	-----S-GLDQE	
		Dinoroseobacter shibae	-----L-----GLDQE	
		118735795	-----L-----GLDQE	
		Wolbachia (Drosophila endo.)	-----V---K---ISV---K---GLSN	
		42519942	-----V---K---ISV---K---GLSN	
		Anaplasma marginale	-----KK---SY---K---GMSSNN	
		56416546	-----KK---SY---K---GMSSNN	
		Ehrlichia chaffeensis	-----K---SL---R---GLSN	
		68171621	-----K---SL---R---GLSN	
		Neorickettsia sennetsu	-----KK---F---R---GLSNK	
		88608380	-----KK---F---R---GLSNK	
		Rickettsia prowazekii	-----K---YA---R---IGMSTE	
		15604016	-----K---YA---R---IGMSTE	
		Rickettsia typhi	-----K---YA---R---IGMNT	
		51473339	-----K---YA---R---IGMNT	
		Rickettsia conorii	-----K---YT---R---IGMSTE	
		6652720	-----K---YT---R---IGMSTE	
		Rickettsia typhi	-----K---YA---R---IGMSTE	
		5813831	-----K---YA---R---IGMSTE	

Figure 5

Partial sequence alignments of RNA polymerase β subunit (RpoB) showing a large insert (boxed) that is a distinctive characteristic of various *Rhodospirillales* species and not found in any other bacteria. There are two homologs of RpoB in *Magentospirillum* (*Mag.*) *magneticum* and only one of these contains the insert. The dashes (-) denote identity with the amino acid on the top line.

genetic trees (Figs. 1 and 2) [29], giving high degree of confidence in the derived inferences. It should be noted that both in our work (Fig. 1) and that by Williams et al.

[29], when analyses were performed using the traditional phylogenetic methods (viz. NJ, MP or ML analyses), the branching of *Caulobacterales* with respect to *Rhizobiales*

Table 8: Protein that are specific for the Rickettsiales group of species

Gene ID	Accession Number	Function	Gene ID	Accession Number	Function
A. Proteins Unique to Rickettsiales (Wolbachia, Ehrlichia, Anaplasma, Rickettsia and Neorickettsia)					
WD0161	NP_965979	Hypothetical	WD0771 ¹	NP_966526	Hypothetical
WD0715	NP_966474	Hypothetical			
B. Proteins Unique to Anaplasmataceae Family (Wolbachia, Ehrlichia, Anaplasma and Neorickettsia)					
WD0083	NP_965909	Hypothetical	WD0827	NP_966580	Hypothetical
WD0157	NP_965975	Hypothetical	WD0863	NP_966613	Hypothetical
WD0821	NP_966574	Hypothetical			
C. Proteins Unique to Anaplasmataceae Family but missing in Neorickettsia					
WD0148	NP_965966	Hypothetical	WD0772	NP_966527	Hypothetical
WD0412	NP_966202	Hypothetical	WD1025	NP_966750	Hypothetical
WD0467	NP_966253	Preprotein translocase, SecG	WD1056	NP_966779	Hypothetical
WD0757	NP_966513	Hypothetical	WD1220	NP_966932	Hypothetical
WD0764	NP_966520	Hypothetical	WD1230	NP_966942	Hypothetical

¹ Missing in Neorickettsia and Orientia

and *Rhodobacterales* was not resolved. In contrast, using the character compatibility approach, the *Caulobacterales* and related species were found to consistently branch in between the *Rhizobiales* and the *Rhodobacterales* (Fig. 2). Previously, this approach has also proven useful in clarifying the phylogenetic placement of *Salinibacter ruber*, which was not resolved by other methods [33,59].

The phylogenetic studies presented here reveal that a number of species belonging to the *Hyphomonadaceae* family (viz. *M. maris*, *O. alexandrii* and *H. neptunium*), for which sequence information is available and that are presently grouped with the *Rhodobacterales*, branch reliably with the *Caulobacterales* (Figs 1 and 2)[29,47]. The same is also true for *P. bermudensis*, which is the only species from *Parvularculales* order, for which sequence information is available. The grouping of these species with the *Caulobacterales* is independently strongly supported by phylogenomic studies, where a number of proteins that are unique for *C. crescentus* were present in these species and at the same time many proteins that are distinctive characteristics of the *Rhodobacterales* were absent in them. These results make a strong case for the transfer of these *Hyphomonadaceae* species and also *P. bermudensis* to an expanded *Caulobacterales* order [29,47]. Another taxonomic anomaly identified by the present study concerns the phylogenetic position of *Stappia aggregata*. This species was originally identified as an *Agrobacterium*-related species, but later transferred to the *Rhodobacterales* order [43]. However, the shared presence of many *Rhizobiales* -specific proteins by *S. aggregata* and the absence of various *Rhodobacterales*-specific proteins in it, strongly suggest that it should be regrouped with the *Rhizobiaceae-Phyllobacteriaceae* species.

The overwhelming majority of the identified α -proteobacteria-specific proteins do not have a homolog showing significant similarity in any other bacteria. The group-specificities of these proteins indicate that their genes have evolved in a common ancestor of these particular groups or clades of α -proteobacteria (Fig. 3). The clade specificities of these proteins also provide evidence that following their evolution, their genes have been transmitted primarily in a vertical manner, and that non-specific mechanisms such as LGTs have not played a significant role in their species distribution. Similar inferences have been reached in earlier studies for proteins that are specific for other higher taxa of bacteria [20,22-24,33,60]. Most of the α -proteobacteria-specific proteins identified in the present work are of unknown function. A number of these proteins are present in the genomes in clusters of two or three, suggesting that they may form functional units and could be involved in related functions [18,26,48,61]. The retention of these α -proteobacteria-specific proteins and conserved indels by the indicated clades of α -proteobac-

teria over long evolutionary periods strongly suggests that they serve essential functions in these groups of bacteria. Hence, studies on their cellular functions may lead to the discovery of novel biochemical and physiological characteristics that are distinctive characteristics of either all α -proteobacteria or their particular subgroups. Lastly, the primary sequences of many of these genes/proteins are highly conserved and they provide novel means for the identification and characterization of these bacteria by PCR-based and immunological methods.

Methods

Identification of proteins that are specific for alpha proteobacteria

The Blastp searches were carried out on each ORF in the genomes of *B. japonicum* USDA 110, *B. suis* 1330, *C. crescentus* CB15, *G. oxydans* 621H, *M. loti* MAFF303099, *N. winogradskyi* Nb-255, *N. aromaticivorans* DSM 12444, *R. sphaeroides* 2.4.1, *Silicibacter sp.* TM1040, *R. rubrum* ATCC 11170 and *Wolbachia* (*D. melanogaster*) endosymbiont to identify proteins that are uniquely present in α -proteobacteria species at different phylogenetic depths. The blast searches were performed against all organisms (i.e. non-redundant (nr) database) using the default parameters, without the low complexity filter [62]. The proteins that were of interest were those where either all significant hits were from the indicated groups (or orders) of α -proteobacteria, or which involved a large increase in E values from the last hit belonging to a particular group to the first hit from any other group and the E values for the latter hits were $> 1e^{-4}$, indicating weak similarity that could occur by chance. However, higher E values were often considered significant for smaller proteins as the magnitude of the E value depends upon the length of the query sequence [62]. All promising proteins were further analyzed using the position-specific iterated (PSI)-Blast program [62]. In this study, we have also retained a few proteins where 1 or 2 isolated species from other groups had acceptable E values, as they provide possible cases of LGTs. For all of the proteins that are specific for α -proteobacteria, their protein ID's, accession numbers and any information regarding cellular functions (such as COG number or presence of any conserved domain) were tabulated and are presented. In describing various proteins in the text, "bll", "bsl", "blr", "BQ", "BR or BRA", "CC", "GOX", "ml", "Nwi", "Saro", "RSP", "TM1040", "Rru" and "WD" indicate the identification numbers of the proteins in the genomes of *B. japonicum*, *Bartonella quintana*, *B. suis*, *C. crescentus*, *G. oxydans*, *M. loti*, *N. winogradskyi* Nb-255, *N. aromaticivorans*, *R. sphaeroides* 2.4.1, *Silicibacter sp.* TM1040, *R. rubrum* and *Wolbachia* (*Dros.*) endosymbiont, respectively.

Phylogenetic analysis

The amino acid sequences for 12 conserved proteins (viz. RNA polymerase β and β' subunits, alanyl-tRNA synthetase, phenylalanyl-tRNA synthetase, arginyl-tRNA synthetase, protein synthesis elongation factors EF-Tu and EF-G, RecA, Gyrase A, Gyrase B, Hsp60 and Hsp70) for different species were downloaded from the NCBI database and aligned using the CLUSTAL x program [63]. In addition to the sequences for 50 α -proteobacteria species, sequences for two deep-branching species viz. *Helicobacter pylori* and *Campylobacter jejuni* [58], were also included for rooting purposes. The sequence alignments for all 12 proteins were concatenated into a single large file and poorly aligned regions were removed using the Gblocks 0.91b program [64]. The final sequence alignment that was used for phylogenetic analyses contained 7652 aligned positions. A neighbour-joining tree based on this alignment was constructed based on Kimura's two parameter model distances using the TREECON program [65]. Maximum-likelihood and MP trees were computed using the WAG+F model plus a gamma distribution with four categories using the TREE-PUZZLE [66] and Mega 3.1 program [67], respectively. All trees were bootstrapped 100 times [68], unless otherwise indicated.

The character compatibility analysis was performed on a concatenated sequences for the above 12 proteins for 25 α -proteobacteria species representing all its main orders plus two outgroup species (i.e. *H. pylori* and *C. jejuni*) [32]. Using the program "DUALSITE" [32], all sites in the sequence alignments where only two amino acid states were found, with each state present in at least two species, were selected. All columns where any gap was present in any of the species were omitted. The useful two state sites were converted into a binary file of "0, 1" characters using the DUALSITE program and this file was used for compatibility analysis [32]. The compatibility analysis was carried out using the CLIQUE program from the PHYLIP (ver. 3.5c) program package [68] to identify the largest clique(s) of compatible characters. The cliques were drawn and the numbers of characters that distinguished different nodes were indicated.

Identification of conserved indels specific for α -proteobacteria subgroups

Multiple sequence alignments for various proteins constructed in this work were visually inspected to search for any indels in a conserved region that was unique to particular subgroups or orders of α -proteobacteria. The group-specificity of any indel was evaluated by carrying out blast searches on a short segment of the sequence (between 80–120 aa) containing the indel and flanking conserved regions against the non-redundant database. The sequence information for various α -proteobacteria was compiled into signature files that are presented. Sequence

information for all other groups of bacteria, which lack these inserts, is not shown.

Authors' contributions

The initial blastp searches on various genomes were carried out by RSG with the computer assistance provided by Venus Wong. AM analyzed the results of these blast searches to identify various group-specific proteins and confirmed their specificities by means of PSI-blast and genomic blasts. Phylogenetic analyses and identification of conserved indels was done by RSG. RSG was also responsible for directing this study, for final evaluation of the results, and for writing this manuscript. All authors have read and approved the submitted manuscript.

Additional material

Additional file 1

Proteins that are specific for the Brucella species. Many of these proteins are specific for all sequenced Brucella species (viz. B. abortus, B. melitensis, B. ovis and B. suis), whereas others are present in only 2 or 3 of these species. The proteins only found in a single Brucella species are not listed here.

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Additional file 2

Bradyrhizobiaceae-specific proteins that are missing in some species. For the proteins listed in this table, all significant hits in Blastp and PSI-Blast searches are from Bradyrhizobiaceae species. However, unlike the proteins listed in Table 3, which are present in all sequenced Bradyrhizobiaceae species belonging to the genera Bradyrhizobium, Nitrobacter and Rhodopseudomonas, these proteins are generally missing in species from one of these three genera.

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Additional file 3

Proteins that are specific for the Sphingomonadales but missing in one or more species. The proteins listed in part (A) of this table are present in three of the following four Sphingomonadales species (Novosphingobium, Erythrobacter, Sphingomonas and Sphingopyxis), whereas those listed in part (B) are present in Novosphingobium and either Sphingomonas or Erythrobacter.

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