Complete Genomes of Two Dipteran-Associated Spiroplasmas Provided Insights into the Origin, Dynamics, and Impacts of Viral Invasion in Spiroplasma

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Accepted: May 21, 2013

Data deposition: The genome sequences reported in this study have been deposited at DDBJ/EMBL/GenBank under the accessions CP005077 and CP005078.

Abstract

Spiroplasma is a genus of wall-less, low-GC, Gram-positive bacteria with helical morphology. As commensals or pathogens of plants, insects, ticks, or crustaceans, they are closely related with mycoplasmas and form a monophyletic group (Spiroplasma–Entomoplasmataceae–Mycoides) with Mycoplasma mycoides and its relatives. In this study, we report the complete genome sequences of Spiroplasma chrysopicola and S. syrphidicola from the Chrysopicola clade. These species form the sister group to the Citri clade, which includes several well-known pathogenic spiroplasmas. Surprisingly, these two newly available genomes from the Chrysopicola clade contain no plectroviral genes, which were found to be highly repetitive in the previously sequenced genomes from the Citri clade. Based on the genome alignment and patterns of GC-skew, these two Chrysopicola genomes appear to be relatively stable, rather than being highly rearranged as those from the Citri clade. Phylogenetic analyses suggest that the susceptibility to plectroviral invasion probably originated in the common ancestor of the Citri clade or one of its subclades. This susceptibility may be attributed to the absence of antiviral systems found in the Chrysopicola clade. Using the virus-free genomes of the Chrysopicola clade as references, we inferred the putative viral integration sites in the Citri genomes. Comparisons of syntenic regions suggest that the extensive viral invasion in the Citri clade promoted genome rearrangements and expansions. More importantly, the viral invasion may have facilitated horizontal gene transfers that contributed to adaptation in the Citri clade.

Key words: Citri–Chrysopicola–Mirum clade, Mollicutes, plectrovirus, Spiroplasma chrysopicola, Spiroplasma syrphidicola, viral insertion.

Introduction

Spiroplasma (Spiroplasmataceae, Entomoplasmatales) is a genus of low-GC, Gram-positive, wall-less bacteria that are distinguished from other genera in the class Mollicutes by their helical morphology. The majority of spiroplasmas are found to be commensals of insects or plants, whereas a small number of species are pathogens of plants, insects, and crustaceans (Gasparch et al. 2004; Gasparch 2010). Phylogenetic analyses based on 16S rDNA suggested that Entomoplasmatales (Spiroplasma, Entomoplasmata, and Mesoplasma) and Mycoplasmatales (Mycoplasma and Ureaplasma) form a major clade of Mollicutes (Mycoplasmatales–Entomoplasmatales), whereas the other major clade includes Acholeplasma and “Candidatus Phytoplasma” (Gasparch et al. 2004). The traditionally delimited Spiroplasma and Mycoplasma were found to be nonmonophyletic (Gasparch et al. 2004; Volokhov et al. 2012). Mycoplasma mycoides and several other mycoplasmas form a monophyletic group within a clade that contains entomoplasmases and mesoplasmases (Mycoides–Entomoplasmatales), which, in

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turn, is nested within a clade that otherwise comprised spiroplasmas. This latter group (Spiroplasma–Entomoplasmataceae–Mycoides) has three additional clades, including the basal kodiets, the Apis sister to the Mycoides–Entomoplasmataceae clade, and the Citri–Chrysoplicola–Mirum clade (Gasparich et al. 2004; Lo et al. 2013).

The Citri–Chrysoplicola–Mirum clade includes the only three phytopathogenic spiroplasmas discovered to date: Spiroplasma citri (Saglio et al. 1973), S. phoeniceum (Saillard et al. 1987), and S. kunkeli (Whitcomb et al. 1986), the causative agents of citrus stubborn disease, periwinkle yellowing, and corn stunt disease, respectively. Similar to the phytopathogenic phytoplasmas, all these phytopathogenic spiroplasmas are transmitted by leafhopper vectors (Saillard et al. 1987). This clade also contains species pathogenic to arthropods, including honeybees (S. melliferum; Clark et al. 1985), Drosophila (S. poulsonii; Williamson et al. 1999), crabs (S. einocheiris; Wang et al. 2004), and shrimps (S. penaei; Nunnan et al. 2005). Because of the great economical importance and early discovery of S. citri and S. melliferum, they have been extensively used for studies on Spiroplasma biology and genomics. Previous genome sequencing efforts revealed that the genomes of these two species contain highly repetitive plectrovirus-related fragments (Carle et al. 2010; Alexeev et al. 2012; Lo et al. 2013). Because these viral fragments can be relatively large in size (~7 kb) and comprise more than 20% of these genomes, the three available Spiroplasma genomes to date are in draft assemblies rather than complete sequences. In addition, similar plectroviral genes were found in the survey sequence tags (Bai and Hogenhout 2002) and in an 85-kb segment (Zhao et al. 2003) of the S. kunkeli genome, suggesting that the presence of plectroviral sequences may be a common characteristic of Spiroplasma. This is in sharp contrast to the sequenced genomes of Mycoplasma and Mesoplasma, where such repetitive regions of viral origin are not found.

Compared with the plant-pathogenic phytoplasmas and vertebrate-pathogenic mycoplasmas, spiroplasmas occupying diverse ecological niches are relatively understudied. Additionally, previous genome sequencing projects for spiroplasmas have only focused on the three species mentioned earlier, and a complete Spiroplasma genome sequence is still lacking. To have a better understanding of the evolution of the plectroviral sequences and their impacts on the Spiroplasma genomes, we chose two other species from the Citri–Chrysoplicola–Mirum clade for whole-genome sequencing.

**Materials and Methods**

**DNA Extraction and Whole-Genome Shotgun Sequencing**

The bacterial strains, S. chrysoplicola DF–1T (ATCC 43209) and S. syphidica EA–1T (ATCC 33826), were obtained from the American Type Culture Collection. For each species, 20 ml R2 medium (Moulder et al. 2002) containing 200 µl culture was incubated at 30°C for 72 h. DNA extractions were carried out using the Wizard Genomic DNA Purification Kit (Promega). One paired-end library (insert size = ~155 bp) and one mate-pair library (insert size = ~4.5 kb) were prepared for each species. Approximately 0.7-Gb 101-bp reads were sequenced for each library on the HiSeq 2000 platform (Illumina, USA) by a commercial sequencing service provider (Yourgene Bioscience, Taiwan).

**Genome Assembly and Annotation**

For each Spiroplasma species, the paired-end and mate-pair reads were used for de novo assembly by the program ALLPATHS-LG release 42781 (Gnerre et al. 2011). The assembled scaffolds were used as the starting point for our iterative assembly improvement process (Lo et al. 2013). For each iteration, we mapped all raw reads from the two libraries to the existing scaffolds using BWA v0.6.2 (Li and Durbin 2009) and visualized the results with IGV v2.1.24 (Robinson et al. 2011). Neighboring scaffolds with mate-pair support for continuity were combined, and reads overhanging at margins of contigs or scaffolds were used to extend the assembly and to fill gaps. The MPILEUP program in the SAMTOOLS v0.1.18 package (Li et al. 2009) was used to identify polymorphic sites. Polymerase chain reaction and primer walking were used to confirm these polymorphic sites and the sequences of repetitive regions. Mapping of the raw reads onto the final assembly using BWA resulted in 300- and 171-fold coverage of mate-pair reads (mapping quality of at least 37) and 586- and 645-fold coverage of pair-end reads (mapping quality of 60) for S. chrysoplicola and S. syphidica, respectively.

The complete genome sequences were processed using RNAmer (Lagesen et al. 2007), tRNAscan-SE (Lowe and Eddy 1997), and PRODIGAL (Hyatt et al. 2010) for gene prediction. The gene name and description for the protein-coding genes were assigned based on the orthologous genes identified by OrthoMCL (Li et al. [2003]; e value ≤ 1 × 10^{-15}) in S. melliferum IPMB4A, S. melliferum KC3, and S. citri Gil3-3X and BlastP (Altschul et al. 1997; Camacho et al. 2009) searches against the National Center for Biotechnology Information (NCBI) nonredundant (nr) protein database (Benson et al. 2012). For functional categorization, all protein sequences were annotated by utilizing the KAAS for comparisons with the previously sequenced Spiroplasma genomes.
tool (Moriya et al. 2007) provided by the KEGG database (Kanehisa and Goto 2000; Kanehisa et al. 2010) using the bidirectional best hit method and a set of selected reference genomes as described in Lo et al. (2013). The KEGG orthology assignment was further mapped to the COG functional category assignment (Tatusov et al. 1997, 2003). Genes that did not have any COG functional category assignment were ascribed to a custom category (category X). CRISPFinder (Grissa et al. 2007) was used to search for CRISPRs (clustered regularly interspaced short palindromic repeats) in the genomes. Open reading frames overlapping with CRISPR repeats were removed from the gene list, as the typical repeat region does not encode proteins (Haft et al. 2005). Positions of genes and viral insertions (see later), GC-skew, and GC content were drawn using Circos (Krzywinski et al. 2009).

**Molecular Phylogenetic Inference**

Previous phylogenetic analyses of the Citri–Chrysopicola–Mirum clade were either less comprehensive in terms of taxon sampling or based solely on 16S rDNA, which did not provide sufficient variable characters to resolve every node. Therefore, we constructed a phylogeny based on a combined data set of 16S rDNA and DNA-directed RNA polymerase subunit beta (rpoB), a single-copy gene with phylogenetic utility for Mycoplasmataceae (Volokhov et al. 2012) and spiroplasmas (Bi et al. 2008). Sequences of the two genes were retrieved for selected species of the Mycoplasmatales–Entomoplasmatales clade with an emphasis on the Citri–Chrysopicola–Mirum clade (supplementary table S1, Supplementary Material online), aligned using MUSCLE v3.8 (Edgar 2004) with the default settings, and concatenated into a single data set. A maximum likelihood phylogeny was inferred using PhyML v3.0 (Guindon and Gascuel 2003) with the GTR + I + G model and six substitution rate categories. Bootstrap supports were estimated from 1,000 samples of alignment generated by the SEQBOOT program of PHYLIP v3.69 (Felsenstein 1989).

**Genomic Analyses**

We aligned the genomes of *S. chrysopicola* and *S. syphidica* using Mauve 2.3.1 (Darling et al. 2010) and calculated their nucleotide sequence identity from a concatenated MUSCLE alignment of single-copy genes using the DNADIST program of PHYLIP. To have a better understanding of the distribution and impacts of the plectroviral insertions in Spiroplasma genomes, we examined the syntenic regions between the Citri and Chrysopicola clades based on the OrthoMCL homologous gene clustering results and identified syntenic regions in the Citri clade with plectroviral genes in the middle or at the margin. We define synten as the occurrence of at least four orthologous genes in the same order. Because most of the plectroviral genes in the *S. melliferum* IPMB4A assembly were removed from the contigs due to unresolved polymorphisms (Lo et al. 2013), this analysis was focused on *S. melliferum* KC3 and *S. citri*. The occurrence of plectroviral genes between two syntenic regions was regarded as a single viral insertion and mapped on the genome maps of *S. chrysopicola* and *S. syphidica* in the middle between the two regions. If a syntenic region could only be found on one side of the plectroviral genes, the viral insertion was mapped next to the ortholog nearest to the plectroviral genes.

We compared the gene content among the five sequenced Spiroplasma genomes by using *M. mycoides* (NC_005364) and *M. penetrans* (NC_004432) as the outgroups. Protein-coding genes were clustered using OrthoMCL as described earlier. Amino acid sequences of single-copy genes were extracted, aligned with MUSCLE, concatenated, and used for PhyML phylogenetic analyses with the LG model (Le and Gascuel 2008). Because the genomes of the two *S. melliferum* strains are approximately 99.9% identical at the nucleotide level based on shared single-copy genes (Lo et al. 2013), we combined these two partial genomes into a pan-genome to better represent the complete genome of *S. melliferum*. This approach is not expected to bias the results of gene content comparisons, because we focus on homologous gene clusters rather than individual genes. For the whole-genome phylogenetic analyses, genes from the strain IPMB4A were used.

**Results and Discussion**

**Phylogenetic Relationships within the Citri–Chrysopicola–Mirum Clade**

A maximum likelihood phylogeny was inferred from a combined data set of 16S rDNA and *rpoB* (fig. 1). All nodes received a bootstrap support of at least 75%. The phylogeny indicates that the Mycoplasmatales–Entomoplasmatales clade is divided into two groups, the Spiroplasma–Entomoplasmataceae–Mycoides clade, and a clade formed by the rest of the mycoplasmas. The Spiroplasma–Entomoplasmataceae–Mycoides clade is further divided into the Ixodetis, Mycoides–Entomoplasmataceae, Apis, and Citri–Chrysopicola–Mirum clades. In general, this phylogeny of spiroplasmas and mycoplasmas is consistent with previous analyses based only on 16S rDNA (Gasparich et al. 2004; Lo et al. 2013) but provides stronger support for each of the major nodes. Additionally, with a more comprehensive sampling, our phylogeny provides new insights into the relationships within the Citri–Chrysopicola–Mirum clade. Previous analyses based on 16S rDNA either failed to resolve the relationships among *S. kunkeli*, *S. phoeniceum* and the clade formed by *S. citri* and *S. melliferum* (Wang et al. 2004) or suggested that *S. kunkeli* was more closely related to *S. citri* and *S. melliferum* than *S. phoeniceum* (Bi et al. 2008; Lo et al. 2013). However, our result strongly indicates that *S. phoeniceum* and *S. kunkeli* are more closely related to each other than either is to the clade of *S. citri* and *S. melliferum*. For the
purposes of this study, we further divided the Citri–Chrysopicola–Mirum clade into three subclades named after the specific epithets of the three component species chosen by Gasparich et al. (2004): the Mirum clade (*S. mirum* and *S. eriocheiris*), the Chrysopicola clade (*S. chrysopicola* and *S. syrphidicola*), and the Citri clade, of which all characterized species except for *S. insolitum* are known to be pathogenic.

Genome Features of *S. chrysopicola* and *S. syrphidicola*

The genomes of *S. chrysopicola* and *S. syrphidicola* were sequenced to completion (fig. 2; table 1). They are approximately 1.1 Mb in size, which are much smaller than the estimated genome sizes of the Citri clade spiroplasmas (1.82 Mb in *S. citri* [Carle et al. 2010], 1.38 Mb in *S. melliforum* IPMB4A [Lo et al. 2013], and 1.55 Mb in *S. kunkei* [Dally et al. 2006]; fig. 1). The GC contents of these two species (28.8–29.2%) are higher than those of the Citri clade (25.9–27.5%) and the Mycoides–Entomoplasmataceae clade (24.0% in *M. mycoides* and 27.0% in *Mesoplasmata flororum*). This agrees with the finding based on biochemically determined GC contents that *S. chrysopicola* and *S. syrphidicola* appeared to be at the high end of the *Spiroplasma* range (Whitcomb et al. 1996, 1997; Gasparich et al. 2004). The genomes of both species demonstrate GC-skew inversions near *dnaA*, the first gene downstream of *oriC* in most bacteria (Mott and Berger 2007), and at about the opposite position of *dnaA* in the genome (fig. 2A). In between these two inversions, the GC-skew remains positive for the first half of the genome and negative for the second except for an 11-kb segment near the replication origin. These two halves roughly correspond to asymmetry in transcription polarity (rings 2 and 3 in fig. 2A). Together, these indicate that the two genomes have been relatively stable, probably with few recent rearrangements. On the contrary, several complete genomes of mycoplasmas or phytoplasmas exhibit anomalous GC-skew patterns (Minion et al. 2004; Westberg et al. 2004; Bai et al. 2006), which are related to rearrangements caused by a high proportion of repetitive sequences (Westberg et al. 2004;
**Fig. 2.**—Genome maps and alignment of *Spiroplasma chrysopicola* and *S. syrphidicola*. (A) Genome maps of *S. chrysopicola* and *S. syrphidicola*. Rings from the outermost to the center: 1) scale marks, 2) protein-coding genes on the forward strand, 3) protein-coding genes on the reverse strand, 4) tRNA (purple) and rRNA (red) genes, 5) putative sites of viral insertions in *S. melliferum* KC3 (red) relative to the respective genomes, 6) putative sites of viral insertions in *S. citri* (green) relative to the respective genomes, 7) protein-coding genes unique to the respective genomes, 8) GC skew, and 9) GC content. Protein-coding genes are color coded according to their COG categories. The positions of the genomic regions shown in figure 3A and B are indicated by arrows. (B) Whole-genome alignment between *S. chrysopicola* and *S. syrphidicola*. 

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**Color Key**

**Information storage & processing**
- [A] Translation, ribosomal structure & modification
- [B] RNA processing & modification
- [C] Transcription
- [D] Replication, recombination & repair
- [E] Chromatin structure & dynamics

**Cell processing & signaling**
- [F] Cell cycle control, cell division, chromosome partitioning
- [G] Nuclear structure
- [H] Defense mechanisms
- [I] Signal transduction mechanism
- [J] Cell wall/membrane/envelope biogenesis
- [K] Cell motility
- [L] Cytoskeleton
- [M] Extracellular structures
- [N] Intracellular trafficking, secretion & vesicular transport
- [O] Posttranslational modification, protein turnover, chaperones

**Metabolism**
- [P] Energy production & conversion
- [Q] Carbohydrate transport & metabolism
- [R] Amino acid transport & metabolism
- [S] Nucleotide transport & metabolism
- [T] Coenzyme transport & metabolism
- [U] Lipid transport & metabolism
- [V] Inorganic ion transport & metabolism
- [W] Secondary metabolites biosynthesis, transport & catabolism

**Poorly characterized**
- [X] General function prediction only
- [Y] Function unknown
- [Z] No COG assignment
Bai et al. 2006). Another characteristic of the *S. chrysopica* and *S. syphicida* genomes is related to the gene organization around the oriC. The gene order *rpnA-rpmH-oriC-dnaA-dnaN* is conserved in *Bacillus*, *Acholeplasma*, *M. penetrans*, *M. mycoides*, and *Mes. flororum*. In the Chrysopica and Citri clades, however, *guaB* is located between *rpmH* and *oriC*, which seems to be a synapomorphy of these spiroplasmas in the Citri–Chrysopica–Mirm clade.

The two genomes of the Chrysopica clade are largely syntenic, except for a 41-kb block of mainly hypothetical genes around position 1,000 kb in the *S. chrysopica* genome (fig. 2B). This block corresponds to an inverted block centered at position 500 kb of the *S. syphicida* genome. Interestingly, the positive-to-negative GC-skew inversion is not right opposite the negative-to-positive inversion in the *S. chrysopica* genome, resulting in a smaller half with positive GC-skew (fig. 2A). In addition, the positive-to-negative inversion is 43 kb upstream of the theoretical replication terminus (right opposite the origin). Taken together, these suggest that the 41-kb block was recently inverted and translocated to its present location in the lineage leading to *S. chrysopica*, resulting in asymmetrical halves with positive and negative GC-skew.

Both *S. chrysopica* and *S. syphicida* have a single rRNA operon (16S–23S–5S), which is consistent with the finding that only one rRNA operon was found in *S. citri*, *S. melliferum*, and *S. kunkeli* (Dally et al. 2006; Carle et al. 2010; Alexeev et al. 2012; Lo et al. 2013). This is different from the Mycoides–Entomoplasmataceae clade, where both *M. mycoides* (Westberg et al. 2004) and *Mes. flororum* contain two rRNA operons with the same 16S–23S–5S organization. Genes related to the helical morphology of *Spiroplasma* (Kürner et al. 2005), *fib* (fibril protein) and *mreB* (cell shape determining protein), are found in both species. It should be noted that, as in the partial genomes of *S. citri* and *S. melliferum*, there are five copies of *mreB* (*mreB1*–*mreB5*) in both species, whereas there are only three copies in *Bacillus subtilis* (Jones et al. 2001). In addition, the five *mreB* paralogs are arranged in a conserved order in both *S. chrysopica* and *S. syphicida*, with *mreB1* located 18 kb and 13 kb upstream of the rRNA operon, respectively, and *mreB2–mreB3* 2 kb upstream of *mreB4–mreB5*, which is 10 kb from *guaB* (the last gene before the replication origin). Interestingly, the cluster of genes from *mreB2* to *mreB5* comprises the first 7 kb of the 11 kb segment with positive GC-skew near the replication origin (fig. 2A). The *mreB* paralogs, along with other genes in this segment, are transcribed from a strand different from that of the flanking genes. In *S. citri*, *mreB1* is located in a small contig bounded by plectroviral sequences, whereas in both *S. melliferum* strains, *mreB1* is located 20 kb upstream of the rRNA operon in the same contig. The other four *mreB* paralogs in both *S. melliferum* and *S. citri* are arranged in the same order as in the Chrysopica clade and the distance between *mreB5* and *guaB* is 10 kb in *S. melliferum* and 13 kb in *S. citri* due to the presence of a plectroviral fragment in between. The conservation in the copy number, gene order, and genomic positions of the *mreB* orthologs between the Citri and Chrysopica clades suggest that their role as
cytoskeletal filaments (Küner et al. 2005) is essential for these spiroplasmas. In contrast to the Citri–Chrysopicola–Mirum clade, no mreB homolog has been reported from the Mycoides–Entomoplasmataceae clade, which is closely related with spiroplasmas, but characterized by pleomorphic or coccoid morphology (Tully et al. 1994; Manso-Silvá et al. 2009).

Although extensive gene decay was reported for the S. citri genome (21% of the annotated coding sequences are truncated) (Carle et al. 2010), such phenomenon is not found in S. chrysopicola, S. syrphidica, or S. melliferum. Similar gene decay is also found in an 85-kb segment of the S. kunkelii genome (Zhao et al. 2003), which, similar to the S. citri genome, has a truncated copy of pyrP (encoding uracil permease). However, this gene is intact in S. melliferum, the sister species of S. citri, and in the Chrysopicola clade. The observation of gene decay in leafhopper-transmitted phytopathogenic S. citri and S. kunkelii, but not in other closely related spiroplasmas, suggests that these vector-borne pathogenic species may have smaller effective population size and suffer elevated levels of genetic drift (Kuo et al. 2009).

Origin and Evolution of Viral Invasion in Spiroplasma

The most striking genome feature of S. chrysopicola and S. syrphidica is that they do not contain any trace of plectroviral genes (table 1), which were found to be abundant in the genomes of S. citri (Carle et al. 2010), S. kunkelii (Bai and Hogenhout 2002; Zhao et al. 2003), and S. melliferum (Alexeev et al. 2012; Lo et al. 2013). In our OrthoMCL results (supplementary table S2, Supplementary Material online), no protein-coding gene from S. chrysopicola or S. syrphidica was grouped with any viral gene of S. citri or S. kunkelii. We used the amino acid and nucleotide sequences of large plectroviral fragments from the Citri clade as queries and performed TBLASTn and blastn searches against the genomes of S. chrysopicola and S. syrphidica. However, no significant hit was found.

The plectroviral fragments in the S. citri genome have been linked to genome rearrangements (Ye et al. 1996; Lo et al. 2013). Even though the recA gene, which is involved in homologous recombination, is known to have been pseudo- genized in S. citri and S. melliferum (Marais et al. 1996; Carle et al. 2010; Lo et al. 2013), genome alignment between these two closely related species (99.0% genome-wide nucleotide sequence identity based on the alignment described earlier) reveals extensive genome rearrangements (Lo et al. 2013). These seemingly contrary observations may be explained by a hypothesis that the extensive rearrangements have occurred soon after the divergence of S. citri and S. melliferum, whereas the losses of recA in these species are relatively recent events (Marais et al. 1996; Lo et al. 2013). Moreover, a comparison between S. citri strains that have been propagated in laboratory for 10 years has found several large-scale changes in their chromosomal organization (Ye et al. 1996), suggesting high levels of genome instability even in the absence of a functional copy of recA.

In contrast to the S. citri–S. melliferum comparison, only one small translocation was observed between S. chrysopicola and S. syrphidica (fig. 2B). This result is surprising because the latter pair is much more divergent (92.2% genome-wide nucleotide sequence identity based on the alignment described earlier). The GC-skew pattern (fig. 2A) further indicates that the genomes of the Chrysopicola clade have been relatively stable. Together, these suggest that the absence of plectroviral fragments in these two spiroplasmas is unlikely to be the result of recent loss of viral fragments but is due to the lack of viral invasion throughout the lineage history of the Chrysopicola clade.

In addition to the three Spiroplasma species that have partial genome sequences available (i.e., S. melliferum, S. citri, and S. kunkelii), the presence of viral fragments was identified in S. phoeniceum based on Southern hybridization (Renaudin et al. 1988). Thus, the susceptibility to viral invasion appeared to be a derived state that originated in the common ancestor of this subgroup in the Citri clade (fig. 1). However, additional evidences suggest that this susceptibility is likely to originate in the common ancestor of the entire Citri clade. First, viruses with similar genome characteristics (single-stranded circular DNA) and morphology to those of S. citri viruses have been isolated from Drosophila-associated spiroplasmas such as S. poulsonii (Oishi et al. 1984; Cohen et al. 1987). Second, based on the genome sizes estimated by pulsed-field gel electrophoresis (fig. 1 and table 1; Carle et al. 1995; Williamson et al. 1997, 1999; Daly et al. 2006; Bi et al. 2008; Lo et al. 2013), all characterized lineages in the Citri clade appear to have experienced genome expansions. Compared with the Chrysopicola clade, the genomes of the Citri clade are larger by 0.26 Mb (S. melliferum) to 0.92 Mb (S. poulsonii). Because plectroviral genes comprise approximately 20% of protein- coding genes in the assembled parts of the S. citri genome (~84% of the whole genome; table 1) and most of the unassembled parts probably correspond to plectroviral fragments (Carle et al. 2010), plectroviral fragments could account for up to 35% of the S. citri genome. This clearly indicates that the viral fragments are a major contributor to the larger genome sizes observed in the Citri clade. This observation is intriguing among Mollicutes because genome reduction linked with host association (Ochman and Davalos 2006; Kuo et al. 2009; McCutcheon and Moran 2012) was found in mycoplasmas and phytoplasmas (Chen et al. 2012).

To the best of our knowledge, plectroviral invasion has not been reported for Mollicutes outside the Citri clade. Plectroviruses with homologous genes of S. citri plectroviruses (Sha et al. 2000) and a similar replication mechanism (Dickinson and Townsend 1984) were isolated from Acholeplasma (Gourlay and Wyld 1973; Steinick et al.

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**Genome Biol. Evol.** 5(6):1151–1164. doi:10.1093/gbe/evt084  Advance Access publication May 27, 2013  1157
another insect/plant-associated Mollicutes genus distantly related to spiroplasmas (fig. 1). Although Southern blot hybridization suggested that plectrovirus genomes seemed to be present in the cellular DNA of several A. laidlawii strains (Just et al. 1989), no viral invasion was reported for the complete genome of A. laidlawii PG-BA (Lazarev et al. 2011). However, other types of repetitive sequences have been found to promote chromosomal rearrangements in various Mollicutes lineages. For example, insertion sequences comprise 13% of the M. mycoides genome and are linked to high genomic plasticity (Westberg et al. 2004). Additionally, the potential mobile units in phytoplasmas are associated with the genome instability observed in these lineages (Bai et al. 2006; Wei et al. 2008; Toroño et al. 2010).

It should be noted that the absence of viral invasion in the Chrysoplicola and Mycocides–Entomoplasmataceae clades is unlikely to be attributed to differences in ecological niches. Plectroviruses have been isolated and shown to infect spiroplasmas from the dipteran genus Drosophila, implying that these viruses might also be found in the hosts of S. chrysoplicola and S. syphicola. Additionally, Mes. florum (originally described as Acholeplasma florum) was isolated from floral surfaces, where acholeplasmas could be found (Tully et al. 1994), and is likely to be exposed to plectroviruses, but it does not contain any plectroviral fragments. Therefore, the susceptibility to viral invasion seems to have a physiological or genetic basis that requires further investigation.

Dynamics of Insertions and Losses of Viral Fragments

By comparing the syntenic regions among the available Spiroplasma genomes, we are able to visualize the distribution of plectroviral fragments in S. citri or S. melliferum relative to the complete genomes of the Chrysoplicola clade (fig. 2A). Except for an approximately 100-kb region extending from the replication origin and an approximately 150-kb region encompassing most of the ribosomal protein genes, the plectroviral fragments are extensively distributed throughout these genomes. Additionally, the numbers of insertions drawn in figure 2A are probably underestimates, because we only mapped viral fragments that are present in the assembled parts of the S. citri and S. melliferum genomes and that are bordered by syntenic regions found in the Chrysoplicola clade. Because several plectroviral insertion sites are shared among strains within S. citri (Bébéar et al. 1996) and within S. melliferum (Alexeev et al. 2012; Lo et al. 2013), these plectroviral fragments are assumed to be inheritable. However, between-species comparisons indicate that few insertion sites are shared between S. citri and S. melliferum (fig. 2A). Moreover, these two species do not have viral fragments in the positions corresponding to those in the 85-kb segment of the S. kunkelii genome (Zhao et al. 2003). The observations of these species-specific viral insertions are in agreement with the findings that plectroviruses are commonly isolated from either S. citri (Cole et al. 1973) or S. melliferum (Liss and Cole 1981) and that plectro viral insertion into the host chromosome has been observed in laboratory-cultured strains (Dickinson and Townsend 1985; Sha et al. 1995). These suggest that integration of plectroviral genomes into the bacterial chromosome is an ongoing process in natural populations of these spiroplasmas. New insertions, rather than differential losses, may thus account for the high proportion of species-specific viral insertion sites in S. citri (24 out of the 30 mapped) and S. melliferum (10 out of the 15 mapped). Considering the close evolutionary relationship (99.0% genome-wide nucleotide sequence identity) between these two species, viral insertions appear to have occurred at a high rate in both lineages.

However, the rapid accumulation of viral insertions seems to be counterbalanced by losses of viral fragments. If viral invasion follows an insertion-only model and the last common ancestor of the Citri clade was susceptible to viral invasion and had a genome size like that of the present-day Chrysoplicola clade (1.1 Mb), 10 unique insertions in S. melliferum since its divergence from S. citri would translate into 374 insertions in the lineage leading to S. melliferum since the last common ancestor of the Citri clade (based on the branch lengths in fig. 1 and assuming a constant rate of new insertions). If the size of every insertion is 6.8 kb (Sha et al. 2000), the present-day S. melliferum would have a genome size of 3.6 Mb, of which 70% comprised plectroviral fragments. As the genomes of S. citri and S. melliferum are not overwhelmed by viral fragments and have not attained such large sizes, it implies that losses of viral fragments are also taking place at a comparable rate to counteract viral invasion. The removal of these repetitive sequences may be partially driven by the intrinsic deletional bias in bacteria that maintains their genome compactness (Kuo and Ochman 2009), as indicated by the presence of truncated viral genes (Carle et al. 2010) and partial viral fragments (Bébéar et al. 1996).

The dynamics of insertions and losses seem to have resulted in greater variability of genome sizes in the susceptible species. Three closely related S. citri strains maintained in different hosts could have a difference in genome size up to 270 kb (Ye et al. 1996). Furthermore, this difference was the result of laboratory propagation within a period of merely 10 years. In agreement with the difference in distribution of viral fragments between S. citri and S. melliferum, this also suggests that the gains and losses of viral fragments are highly dynamic.

Impacts of Viral Fragments: A Closer Look

Several previous studies suggested that the plectroviral fragments in S. citri and S. melliferum may be associated with chromosomal rearrangements and horizontal gene transfers (Ye et al. 1992, 1996; Alexeev et al. 2012; Lo et al. 2013). To test this hypothesis, we compared the syntenic regions between genomes of the Citri and Chrysoplicola clades. These
comparisons allowed us to establish the directionality of rearrangement events. Moreover, the gene-level resolution provided by the genome sequences allowed us to identify boundaries of viral integration and candidates of horizontally transferred genes. Many genes unique to either or both *S. citri* and *S. melliferum* were indeed found in the vicinity of plectroviral fragments. For example, examination of one insertion site in *S. melliferum* KC3 revealed nine such genes (fig. 3A).

These genes include a set of type I restriction enzyme and methylase (*hsdR* and *hsdM*), as well as a phosphotransferase system (PTS) component, a lysophospholipase (*yqkD*), and a glycerophosphoryl diester phosphodiesterase family protein (*glpQ*). In another example, 11 genes unique to *S. citri* are found in the vicinity of plectroviral fragments (fig. 3B). It is notable that genes unique to *S. citri* are also found at the boundaries of contigs, which are supposed to be bordered by repetitive plectroviral sequences (Carle et al. 2010).

Association of viral fragments with genes unique to the Citri clade does not unambiguously indicate that the genes were brought in along with the viral genome. However, the range of genome sizes (6.8 kb [Sha et al. 2000] to 8.5 kb [Dickinson and Townsend 1984]) and nonconservation in gene contents of plectroviruses (Sha et al. 2000) do suggest they may carry bacterial DNA through imprecise excision of prophage genome from the host chromosome. Furthermore, plectroviruses may be able to invade different strains or even different species of the Citri clade, as indicated by the experimental infection of various *S. citri* strains by viruses isolated from *S. melliferum* (Sha et al. 1995) and the infection of Drosophila-associated spiroplasmas by viruses isolated from other Drosophila spiroplasma strains (Oishi et al. 1984; Cohen et al. 1987). Because virulence-related genes could be transferred among bacteria by viruses (Beres et al. 2002), the ability to infect different spiroplasmas indicates that plectroviruses may help to spread virulence factors among different Citri species that share similar niches, such as the three leafhopper-transmitted phytopathogenic spiroplasmas (Liu et al. 1983; Whitcomb et al. 1986; Saillard et al. 1987).

The example in figure 3B also demonstrates possible rearrangement events unique to *S. citri*. In this scenario, where the gene order in *S. melliferum* and the Chrysopicola clade represents the ancestral state, a viral fragment was inserted into between celC and upp (fig. 2A). Another viral fragment was inserted into between fur and proS with an inverted orientation. Subsequent recombination(s) between these two inserted fragments replaced the genes downstream of celC for fur and its downstream genes. Other rearrangement events may have occurred involving genes downstream of tyrS, such that only the segment from fur to tyrS was

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**Figure 3.** Examples of syntenic regions with plectroviral genes in the Citri clade. Annotated plectroviral genes are highlighted in red with labels indicating the ORF number assignments. Hypothetical genes with unknown functions are in gray. Genes within plectroviral sequences, where not indicated, are found in both *Spiroplasma citri* and *S. melliferum*. Putative viral insertion positions are indicated by red arrowheads. The corresponding regions in *S. syphidicola* contain the same genes in the same order as in *S. chrysopelia* and are not shown for simplicity. In (A), the corresponding region in *S. citri* has other plectroviral fragments in the vicinity (fig. 2) and is scattered into different contigs.
rearranged from its original position (downstream of proS) to the position downstream of celC. According to the physical mapping (Ye et al. 1992; Carle et al. 2010), the effect of these rearrangements would correspond to a translocation across a genomic distance of approximately 240 kb from one side of oriC to the other.

**Comparative Analyses of Gene Content**

To better understand the potential impacts of viral invasion on the gene content of susceptible genomes, we compared the content of homologous gene clusters in the four *Spiroplasma* species with sequenced genomes (fig. 4 and supplementary table S2, Supplementary Material online). With *M. mycoides* and *M. penetrans* as the outgroups, the presence/absence patterns of homologous gene clusters were mapped onto the whole-genome phylogeny (fig. 5 and supplementary table S3, Supplementary Material online). The genes unique to the four species from the Citri and Chrysopicola clades include *mreB* and *fib*, which are related to the spiroplasma morphology (Küner et al. 2005), and *spi* (spiralin gene), which encodes an abundant lipoprotein in the plasma membrane (Killiny et al. 2005) that appears to be unique to the Citri–Chrysopicola–Mirim clade (Meng et al. 2010). We found that *treB*, encoding a PTS component for transporting trehalose, the main hemolymph sugar in insects (Becker et al. 1996), is uniquely absent in the Chrysopicola clade. Because spiroplasmas in insect hemolymph must rely on trehalose (André et al. 2003), the lack of *treB* indicates that both *S. chrysopicola* and *S. syphidicola* may be unable to proliferate within hemolymph. It has been shown that multiplication within hemolymph and subsequent invasion to other tissues are highly related to the pathogenicity of *S. citri* to its insect hosts (Liu et al. 1983). Together, these may explain why both *S. citri* and *S. melliferum* are pathogenic to their insect hosts (Liu et al. 1983; Clark et al. 1985), whereas there is no report of entomopathogenicity for *S. chrysopicola* and *S. syphidicola*.

Additionally, the genomes of the Citri and Chrysopicola clades differ in their content of antiviral systems. A Type I restriction/modification (R/M) system is intact in *S. chrysopicola* (SCHRY_v1c02520-2540) and apparently truncated in *S. citri* (SPICI01B_067-69), whereas another Type I R/M system is present in *S. melliferum* KC3 (SPM_0320-325; fig. 3A). However, only *S. chrysopicola* and *S. syphidicola* contain genes for a Type II R/M system (SCHRY_v1c02610, SCHRY_v1c06730-6740; SSYRP_v1c02700, SSYRP_v1c06950-6960). A CRISPR system is first found at approximately 1,010 kb of the *S. syphidicola* genome (CRISPR-associated [cas] genes: SSYRP_v1c09370-9400; CRISPR repeats: positions 1,005, 965–1,007,784). This is the first CRISPR system reported from the *Spiroplasma*–*Entomoplasmataceae–Mycoides* clade of Mollicutes. Although other CRISPR systems are found in *Mycoplasma* and *Ureaplasma* (Makarova et al. 2011), this *S. syphidicola* CRISPR system is more similar to that of *S. melliferum*.

![Fig. 4.—Numbers of shared and unique gene clusters among *Spiroplasma chrysopicola*, *S. syphidicola*, *S. citri*, and *S. melliferum*. Genomes of the two *S. melliferum* strains (IPMB4A and KC3) were analyzed as a single pan-genome of *S. melliferum*. For detailed lists of these orthologous gene clusters, see supplementary table S2, Supplementary Material online.](image-url)
S. chrysopicola and S. syrphidicola (figs. 1 and 5). This may be accounted for by the potential role of viral invasion in promoting gene gains through horizontal transfer. The high insertion rates of viral fragments, coupled with differentiation in ecological niches between S. citri and S. melliferum, may explain why they accumulated high numbers of lineage-specific genes shortly after their divergence.

Building a Model for Viral Invasion

Based on the phylogenetic and comparative genome analyses in the previous sections, we proposed a model for the genome evolution of spiroplasma species susceptible to viral invasion as a summary of this study (fig. 6). We hypothesize that the common ancestor of the Chrysopicola and Citri clades was resistant to virus attack and had few or no viral fragments in its genome.
genome, probably similar to the genomes of present-day *S. chrysopica* and *S. syrphidicola*. After the divergence of the Chrysopica clade, the common ancestor of the susceptible species lost its antiviral system (RVM, CRISPR, or other systems) and started to accumulate viral fragments in the genome. The insertion of these viral fragments has several consequences, including increased genome sizes, homologous recombination between similar copies of viral fragments, and acquisitions of nonviral foreign DNA via phage-mediated horizontal gene transfers. As new viral fragments are inserted into the genome, old fragments are lost due to a mutational bias toward deletions, which counterbalances the high insertion rate. The loss of viral fragments implies that the older, viral fragment-mediated rearrangements may no longer be recognizable from the current distribution of viral fragments. As a result of the accumulated effects of viral invasion, these spiroplasmas are characterized by larger and more variable genome sizes, nonconserved genome organization, and higher numbers of lineage-specific genes. In contrast, the genomes from the Chrysopica clade remain relatively small and have few incidences of rearrangements. This model may serve as a working hypothesis for genome evolution of susceptible spiroplasmas. More studies, including genome sequencing for other species from the Citri (*S. spiroplasmas*. More studies, including genome sequencing for other species from the Citri (*S. spiroplasmas*). **Supplementary Material**

**Supplementary tables S1–S3** are available at Genome Biology and Evolution online (http://www.gbe.oxfordjournals.org).

**Acknowledgments**

The authors thank the DNA Analysis Core Laboratory (Institute of Plant and Microbial Biology, Academia Sinica) for providing Sanger sequencing service. This work was supported by the research grants from the Institute of Plant and Microbial Biology at Academia Sinica and the National Science Council of Taiwan (NSC 101-2621-B-001-004-MY3) to C.H.K.

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**Associate editor:** John McCutcheon