Surface proteins in pathogenic bacteria often function as antigens, and evidence indicates that their molecular evolution is driven by both positive selection and recombination (1, 28). Examples include the pilin genes from Neisseria species (17, 21), msp2 from Anaplasma marginale (12, 37), porB from Neisseria meningitidis (58), and ompL1 of the Leptospira genus (19). The examples above all involve pathogenic bacteria of vertebrates, and it is believed that recombination and rapid sequence evolution in their surface antigens are selectively advantageous by promoting avoidance of the vertebrate host immune response. Less well understood is the evolution of surface proteins of bacteria found strictly in invertebrates. Here we investigate the patterns of variation in the surface protein of Wolbachia, an intracellular bacterium found in arthropods and nematodes.

Wolbachia bacteria are among the most successful and intriguing intracellular bacteria in nature. It is estimated that 20 to 75% of insect species harbor Wolbachia bacteria (25, 62, 65), with infections also commonly found in terrestrial crustaceans, chelicerata, and filarial nematodes (3, 5, 14, 18, 49).

Transmission of Wolbachia bacteria within host populations is vertical (64); however, it is now well known that Wolbachia bacteria in arthropods can also shift host species, “jumping” to new unexplored cellular environments through mechanisms still unclear (horizontal transmission) (60, 63, 68).

As a parasite of arthropods, Wolbachia bacteria are best known to be manipulators of host reproduction. A major genetic effect of the symbiosis with Wolbachia bacteria is a distortion of the host sex ratio, through mechanisms enhancing the female proportion (the sex transmitting the bacterium), such as feminization of genetic males, parthenogenesis induction, and male killing (54, 64). Wolbachia bacteria are also able to induce cytoplasmic incompatibility between eggs from uninfected females and sperm from infected males, thus rapidly increasing the proportion of infected individuals in host populations, often to fixation (48). While in insects Wolbachia bacteria are primarily reproductive parasites, in filarial nematodes the symbiosis with Wolbachia bacteria appears to have evolved toward a mutualistic interaction (3, 4).

The genus Wolbachia (class Alphaproteobacteria, order Rickettsiales) is currently divided into six taxonomic supergroups (A to F) based primarily on 16S and ftsZ gene phylogenies. Phylogenies for these two genes are concordant at the supergroup level (31). A and B are the two main groups found in arthropods. C and D are found in filarial nematodes (3). Recently, two new supergroups, E and F, have been proposed. So far, supergroup E contains Wolbachia bacteria infecting springtails (class Collembola), a primitive insect group (15, 59), while supergroup F contains Wolbachia bacteria that infect termites and filarial species of the genus Mansonella (31, M. Casiraghi, S. R. Bordenstein, L. Baldo, N. Lo, T. Beninati, J. J. Wernegreen, J. H. Werren, and C. Bandi, unpublished data).

The vertical transmission of Wolbachia bacteria through the reproductive tissues of their hosts implies that these bacteria experience little recombination, as appears to be the case for other vertically inherited symbionts (e.g., Buchnera aphidicola) (56). However, the discordances between the phylogenies of some Wolbachia genes (27) and the discovery of recombination in bacterium-host interactions is currently unknown, but results presented here indicate that exchanges of IHR motifs are favored by natural selection. Identifying host proteins that interact with wsp variants should help reveal how these widespread bacterial parasites affect and evolve in response to the cellular environments of their invertebrate hosts.
events within the Wolbachia surface protein (wsp) (47, 66) suggested that recombination may be more common in Wolbachia bacteria than some other endosymbiotic bacteria. Furthermore, the relatively frequent occurrence of multiple infections with different Wolbachia strains in the same hosts (23, 62, 63), the presence of phages and insertion elements within the Wolbachia genome (35, 68), and lateral transfer of phage among strains (7) are consistent with a recombinogenic genome.

The Wolbachia surface protein gene wsp encodes a major surface membrane protein showing sequence similarity to the major outer membrane proteins of closely related alphaproteobacteria (9). Among the Wolbachia genes for which a large sequence data set is currently available, wsp is the most variable, showing relatively high genetic divergence among strains. Analyses of the rates of synonymous and nonsynonymous substitutions along the gene sequences show discrete regions under strong positive selection in a background of overwhelming purifying selection (2, 28). Because of its variability, wsp has been used extensively in phylogenetic analyses and for microtaxonomic subdivision of the two major clades, A and B, into subgroups (68).

Localization of the protein at the interface between the two cellular environments and the presence of regions under strong positive selection suggest a key role of the protein in the arms race expected to occur between arthropod hosts and this intracellular parasite (2, 61). Furthermore, in nematode Wolbachia bacteria, wsp has been demonstrated to play an antigenic role in stimulating the immune response of the vertebrate animals that are infected by filarial worms (6, 10, 11).

Despite the fact that Wolbachia bacteria are not found in vertebrates, their outer surface membrane protein (Wsp) shows surprising analogies with antigenic proteins of pathogens: a heterogeneous pattern of variation characterized by hypervariable regions (HVRs) flanked by conserved regions (CRs) (2, 9), strong positive selection affecting the HVRs (2, 28), and evidence of recombination affecting its sequences (26, 47, 66). All this strongly suggests a potential role of this protein in host-Wolbachia interactions.

Previous evidence of recombination in wsp has come from three studies. Werren and Bartos (66) reported the first example of recombination within supergroup B, occurring between the two Wolbachia strains of a parasitoid wasp and the fly it parasitizes. More recently, Reuter and Keller (47) showed recombination to have occurred among five strains of a parasitoid wasp and the fly it parasitizes. Among the 40 wsp sequences selected for the main analysis, 37 were already available in GenBank. Sequences from Coptotermes lacteus and Coptotermes acinaciformis were obtained during this study. The two species were gifts from Michael Lenz (CSIRO Entomology) and John Holt (James Cook University), respectively, and were collected in Melbourne and Townsville. The guts of a single worker termite from each species were removed, and DNA was extracted from the remaining tissues as described previously (32).

PCR was performed using the conditions described by Maekawa et al. (32), with primers WPestF (5′-TTAGACTGCTAAAGTGGAATT) and WPestR (5′-AACCCTGGGATAAGAAAGA). Direct sequencing of the PCR product was performed using the BigDye v2.0 terminator sequencing kit and an ABI 3700 automated sequencer.

Analysis of recombination. (i) Detection of breakpoints. To identify potential recombination breakpoints, we used the recombination detection program RDP2 (34), which implements different methods for detecting recombination. We primarily used the MAXCHI program (43, 52), which implements different methods for detecting recombination. For every possible sequence pair in the alignment, a window of set length with a partition in its center is moved along the sequences and a chi-square value is calculated, being an expression of the difference in the number of variable sites on either side of the central partition. A variable window size setting, with different proportions of variable sites (VI) per window was initially tested, providing basically similar results regardless of the VI proportions. To estimate the pattern of distribution of recombination events, the parameters were fixed as follows. Sequence triplets were scanned using a variable window size with a 0.3 fraction of VI and a highest acceptable P value of 0.001. For specific detection of breakpoints in the selected subsamples of sequences (see Fig. 3), we used the option “Manual MaxChi,” which permits the analysis to be performed selecting potential recombinant and parental sequences. The significance of chi-square peaks was more accurately determined by a permutation test (1,000 permutations). Peaks at which the observed chi-square values exceeded values in the 5% tail of the null distribution were considered significant.

(ii) Phylogenetic analyses. Phylogenetic analyses were performed on different portions of wsp. The nucleotide sequence alignment was subdivided into four
Table 1. List and features of the 40 Wolbachia strains analyzed in this study

<table>
<thead>
<tr>
<th>Host species</th>
<th>Host order</th>
<th>Wolbachia supergroup*</th>
<th>Strain identifying code</th>
<th>Accession no.</th>
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* Supergroup identification for each strain is based on published literature. A question mark indicates that no identification of the supergroup is provided.

sectors of nearly equal length (about 135 bp each) starting from the first nucleotide. Partitioning was based on the observed pattern of nucleotide sequence divergence along the alignment and on the pattern of the ratios of dN/dS (the nonsynonymous/synonymous substitution rate ratio per codon site) reported for wsp by Baldo et al. (2), identifying three regions under positive selection in arthropod Wolbachia bacteria, separated by CRs (in that study, the last HVR was only partially included in the analyses). The four sectors were divided within the middle of each CR, allowing each segment to encompass a single HVR plus part of the flanking conserved domain. Since the vast majority of nucleotide sequence variability is at the HVRs, CRs are expected to have a minor effect on the phylogenetic reconstructions.

Phylogenetic analyses of each sector were conducted using Bayesian inference (BI) and maximum parsimony. For BI, the appropriate models of sequence evolution were estimated for each of the four gene partitions using the program Modeltest 3.06 (42). In each case, this was found to be GTR+I+G (GTR, general time-reversible model; I+G, invariable or gamma distributed rates of variation at sites).

The BI analyses were performed using MrBayes 3.2 (22). One hundred thousand trees were generated, with a sample frequency of 100. The first 500 trees were considered the burn-in and discarded. From the remaining 500 trees, 50% majority rule consensus trees were generated. Maximum-parsimony 50% majority rule bootstrap trees were estimated in PAUP* (55) (1,000 replicates, 10 random-addition replicates per bootstrap replicate). All characters were weighted equally, and gaps were treated as missing. Since HVR3 and -4 were difficult to align unambiguously, we performed phylogenetic analyses on various alignments of both regions.

Test for selective pressure at the HVRs. To investigate whether the HVRs show stabilizing or diversifying selection, we used an independent-contrast approach (13) to evaluate rates of synonymous versus nonsynonymous substitution (Ks and Ka) of phylogenetically independent sets of closely related sequences. Independent contrasts were selected based on the phylogenetic analysis. The advantages of this approach are that alignment issues are avoided due to similarity in sequences and relatively short-term trends in synonymous versus nonsynonymous divergence can be evaluated. Sets of phylogenetically independent contrasts were compared using a one-tailed Wilcoxon matched-pair signed-rank test (13).

RESULTS

Pattern of variation along wsp. The pattern of nucleotide sequence divergence along wsp, based on 93 sequences, is shown in Fig. 1a. A heterogeneous distribution of variability is evident along the gene: four HVRs are seen as peaks, with similar maximum values of nucleotide sequence divergence of around 0.5, separated by regions under strong conservation (CRs). The corresponding amino acid pattern of variation mir-
alignments difficult, particularly for HVR4. This merely reflects the pattern of considerable divergence between motifs, with relatively few transitional sequences. For this reason, we do not claim that the alignment between motifs at the HVRs (shown in Fig. 2) reflects actual nucleotide sequence homologies. The alignments are very useful, however, for showing conservation within and divergence between amino acid sequence motifs.

Recombination in wsp. We have examined the wsp gene for recombination by three basic methods. First, we have evaluated the amino acid sequences for signatures of recombination between the HVRs by visual inspection. Examples of such recombination are readily apparent in Fig. 2, and a sample of these is described below. Second, we have analyzed the sequences for recombination occurrence and specific breakpoints using primarily the program MAXCHI. Third, we have used phylogenetic approaches to further support significant discordances at the nucleotide sequence level for the different HVRs of wsp and to show how relationships among sequences shift across the four HVRs.

Recombination between HVRs, which results in shuffling of motifs between sequences, is readily apparent by examination of Fig. 2. The patterns are easily visualized by the color coding of HVRs based on amino acid motifs. In HVR1, there are clear similarities among sequences 1 and 2, 3 to 6, 7 and 8, 9 and 10, 11 to 13, 14 to 23, 24 to 26, 27 to 30, 31 to 36, 37 and 38, and 39 and 40. However, these same groups of sequences can differ dramatically in their composition in other HVRs. For example, by HVR4, sequence 1 now groups with sequences 9 and 15 by amino acid motif, whereas sequence 2 now groups with 14 by amino acid motif. Basically, sequences showing almost identical amino acid motifs at some HVRs are dramatically divergent at different HVRs while grouping with different sets of sequences based upon shared motifs. Such a remarkable pattern of similarity or divergence among sequences along the gene strongly indicates a shuffling among a limited set of motifs. We note that all sequences shown in Fig. 2 are consistent with the above recombinant pattern. Below, we illustrate the recombinant pattern with three examples. These examples are also highlighted in the analysis of breakpoints at the nucleotide sequence level (Fig. 3) and phylogenetic analyses (Fig. 4).

Example 1. Sequences 1-DmelA (from the dipteran Drosophila melanogaster) and 2-CglaA (from the hymenopteran Callyrythys glandium) share 100% amino acid identity at HVR1, -2, and -3, while at HVR4 the two sequences greatly diverge. At HVR4, sequence 1-DmelA is very similar to sequences 9-BbroA (from the hymenopteran Blastophaga brownii) and 15-TdroA (from the parasitic wasp Trichopria drosophilae) with 21/25 monomorphic sites (Ms) shared between the two. In contrast, sequence 2-CglaA is almost identical to sequence 14-LwhiA (the dipteran Lutzomyia whitmani, 24/25 Ms).

Example 2. Sequence 7-PherA (from the fig wasp Pegoscapus herrei) is almost identical to sequence 8-FexsA (from the ant Formica exsecta) at HVR1 and HVR2 (respectively, 21/22 and 23/24 Ms), while at HVR3 the two sequences diverge. Sequence 7-PherA becomes almost identical to sequences 5-CuncA (from the louse Colpocephalum unciferum), 6-ScrA (from the lepidopteran Sitotroga cerealella), 9-BbroA, and 10-DsmA (from the fruit fly Drosophila simulans) (17/18 Ms), and sequence 8-FexsA converges to sequences 3-EhorA (from the
FIG. 2. Amino acid alignment of 40 wsp sequences (180 amino acid in length). The reference sequence is 1-DmelA (from the host D. melanogaster). Amino acid motifs at the HVRs are color coded based on similarity of shared polymorphisms within each HVR. HVR1 is used as the initial reference region for sequence grouping. HVR motifs with uncertain groupings were left uncolored. Based on the pattern of colors along the gene, each of the first 34 arthropod sequences shows a unique combination of four HVR motifs and thus could be regarded as a distinct protein type. A clear shuffling of HVRs between sequences can be visualized. CR1 (about 145 bp) is not shown because the region was not completely sequenced for all of the strains. Arrowheads indicate the limits of the alignment sectors (HVR+) analyzed for phylogenetic reconstructions (see Fig. 4).
FIG. 3. Examples of recombination breakpoints along \textit{wsp}. For each alignment, only the polymorphic sites are shown. Arrows indicate significant breakpoints detected by MAXCHI. Polymorphisms shared with the underlined sequence are highlighted in gray. Also shown are the positions of the four HVRs. Polymorphisms shared between all four HVRs are highlighted in gray. In all cases, recombination breakpoints fall between HVRs.
FIG. 4. Phylogenetic trees of the four portions of wsp encompassing single HVR+/- (135 bp each). The trees were generated by MrBayes (100,000 replicates, 50% majority rule) and are unrooted. Sequence identification corresponds to the description provided in Table 1. Colors highlight some of the examples discussed in the text (see Results) and show changes in phylogenetic association across HVR+/- affecting all sequences shown. Sequences of supergroups A and B do not form separate groupings at any HVR. Posterior probability values higher than 70% are shown at the nodes. For nodes supported also by parsimonious analyses, the corresponding bootstrap value is shown under the posterior probabilities.
louse Echinophthirus horridus), 4-BbovA (from the louse Bo-
vicola bovis), 26-PsiA1 (from the dipteran Protocalliphora sia-
lia), 33-EcauA (from the lepidopteran Ephestia cautella), and
34-TqueA (from the louse Trinoton quereducale) (17/18 Ms).
Then, at HVR4, sequence 7-PherA2 diverges from all the pre-
vious sequences and becomes identical to sequences 6-ScerA
and 21-BnipB (from the hymenopteran Blastophaga nipp-
onica), while sequence 8-FexA remains very close only to
sequences 4-BbovA 26-PsiA1, and 34-TqueA (24/25 Ms).

**Example 3.** Sequence 30-AvalB is from a Wolbachia bacte-
rarium present in the isopod Armadillidium vulgare and induces
feminization in this host. Phylogenetically, this Wolbachia bac-
terium is embedded within a clade of bacteria otherwise found
in insects and therefore probably represents a major host shift
from insects to isopods (8). At HVR1 and -2, this sequence
groups with sequences 29-PyrB (from the isopod Porcellion-
tes pruinosis, 22/22 and 21/24 Ms, respectively, at HVR1 and
HVR2), 28-PyR (from the isopod Porcellio spinicornis, 21/22
and 22/24 Ms), and 27-AenCB (from the lepidopteran Acraea
enedon, 17/22 and 20/24 Ms). At HVR3, it diverges dramat-
ically from the previous grouping and, interestingly, groups
strongly with some sequences from supergroup A, 31-PgenA
(from the hymenopteran Peganopus gemellus, 13/18 Ms) and
32-DeryA (from the spider Dysdera erythrina, 14/18 Ms), and
with 16-Plum? (from the louse Pediculus humanus, 13/18 Ms).
It then diverges from these in HVR4, where it does not show
strong similarities to any other HVR4 in the data set (but it
shows high similarity with other wsp sequences from isopod
hosts not in the data shown; e.g., sequences with accession no.

(i) **Recombination breakpoints.** The pattern of recombi-
nation in wsp appears highly complex. Analyses performed with
the MAXCHI program did not identify one partitioning of wsp
describing all the recombinant patterns, since breakpoint loca-
tions are quite different among sequences and recombination
can involve segments of different lengths.

For this reason, we first characterized the general pattern of
recombination within the gene. We then analyzed some of
the same examples previously reported to show the recombination
events at the nucleotide sequence level. Figure 1b presents an
analysis using MAXCHI, which calculates the number of re-
combination blocks among the 40 sequences for each position
along the gene. The number of detected hits can differ consid-
erably, depending on the parameter settings (e.g., the window
size), ranging from 480 to 1,600 for variable window sizes
ranging from 0.1 to 0.5 of variable sites (VI). However, despite
minor changes in parameter values, the pattern of distribution
of hits along the gene is quite consistent. Data indicate that
potential recombination hits have involved all sites along the
gene. The number of hits per site increases around position
215 of the alignment (in CR3, after HVR2), levels off in
HVR3, increases again around position 380 (before HVR4),
and levels off in HVR4. The fact that all sites within a single
HVR experience nearly identical numbers of events indicates
that single HVRs are generally exchanged as whole tracts. We
cannot exclude the possibility that in some cases recombi-
nation could have occurred within HVRs, involving small se-
quence tracts within these regions, but there is no significant
evidence for this from our analyses of breakpoints. Regions
with an increasing number of hits (CR3 and CR4) involve sites
that have undergone an unequal number of recombination
events, thus reflecting the occurrence of recombination break-
points. This suggests that recombination events are more likely
to occur between HVR2 and -3 and between HVR3 and -4
than between HVR1 and -2. This observation is confirmed also
by visual inspection of the protein alignment (Fig. 2) and
phylogenetic analyses (Fig. 4) showing most of the incongru-
ences in relationships occurring between the regions cited
above.

Recombinant segments can involve either single HVRs or
longer segments, encompassing two or more HVRs at a time
(in all cases, CRs can also be recombed). Some sequences
show a single recombinant breakpoint, while some show sev-
eral breakpoints leading to partitions of the gene in multiple
recombinant blocks. Consequently, recombinant sequences
can be characterized by an x segment mosaic (from two to four
segments).

Figure 3 shows four cases of recombination breakpoints at
the nucleotide sequence level. For 1-DmelaA, 2-CglAa, and
14-LwthA, a single significant breakpoint is present in the
region around position 461 of the nucleotide sequence align-
ment, corresponding to the 5' end of HVR4 (Fig. 3a). The
breakpoint divides the alignment into two portions: the first
portion encompasses HVR1, -2, and -3 (CRs included), while
the second involves only HVR4. Before the breakpoint, se-
quen 2-CglAa is identical to 1-DmelaA, sharing all 73 poly-
morphisms with it, while in the second portion it clearly be-
comes identical to sequence 14-LwthA at 48/49 polymorphic
sites. Statistically significant strings of associated polymor-
phisms strongly indicate recombination (53): the probability
of getting a string of 73 matching polymorphisms followed by
a string of 48 is remarkably low ($P < 10^{-15}$ based on the
permutation probability).

Figure 3b shows a more complex pattern, involving four
sequences and a single recombinant breakpoint around nucleo-
tide position 240, corresponding to the 3' end of HVR2.
Based on the shared polymorphisms, the breakpoint divides
sequences into two portions, clearly grouping sequences
4-BbovA with 6-ScerA and sequences 7-PherA with sequences
8-BbroA before the breakpoint and with sequences 4-BbovA
with 8-BbroA and sequences 6-ScerA with 7-PherA after the
breakpoint. This represents either independent recombinant
sequences with a common breakpoint (e.g., recombination hot-
spot) or a reciprocal exchange event.

Figure 3c shows the recombinant pattern involving sequence
30-AvalB. It shows similarity at the nucleotide sequence level
to sequence 28-PspiB before position 270, with a breakpoint
occurring within CR3, and to sequences 31-PgenA and 32-
DeryA after. A second breakpoint is localized around position
451, with 28-PspiB having high similarity to 31-PgenA and
32-DeryA, while 30-AvalB dramatically diverges from all the
sequences. This suggests recombination occurring between A
and B sequences.

Similar to the previous example, Fig. 3d shows recombinants
involving A and B sequences. Sequence 21-BnipB is iden-
tical to sequence 22-OscaB before position 400 (in CR4), shar-
ing all of the 119 polymorphisms with it. After that, it becomes
identical to sequences 7-PherA2 and 6-ScerA, sharing all of the
remaining 36 polymorphisms with them.

The origin or direction of these recombination events cannot
be reliably inferred because the DNA exchange history in wsp appears too reticulated to be resolved. A preliminary attempt to resolve relationships among some strains by examining additional genes has so far been unsuccessful. Other genes also provided ambiguous inferences about strain relationships, probably reflecting the widespread recombination in Wolbachia genomes (L. Baldo, J. H. Werren, and S. R. Bordenstein, unpublished data).

(ii) Phylogenetic analyses. As indicated by the patterns of amino acid similarity or divergence along the sequences described above, phylogenetic relationships predicted for the four HVRs are in conflict. To provide further statistical significance for this pattern and to show how relationships shift across HVRs in the whole data set, we compared the nucleotide sequence phylogenies of the four regions of wsp. The wsp gene was divided into four sectors; the breakpoints for these sectors are indicated in Fig. 2 (each sector is indicated as HVR+ to point out that it contains both the HVR and a portion of the flanking CRs).

Shown in Fig. 4 are the four consensus Bayesian trees describing the phylogeny of the four sectors of wsp. The tree topologies from bootstrap analysis were generally congruent with those estimated from Bayesian analysis, and associated bootstrap values were similar. The use of different alignments of HVR3+ and HVR4+ resulted in consistent phylogenies for the main clades: for this reason, only one phylogeny of these regions, based on the alignment showed in Fig. 2, is reported.

The goal of this analysis was not to define the precise phylogenetic relationships between motifs, as these will be influenced by alignment issues (i.e., small insertions or deletions that complicate alignment between motifs). Rather, the analysis is used to show phylogenetic relationships within motifs (where alignment is not an issue) and the shuffling of these relationships between HVRs.

As shown in Fig. 4, the evolutionary relatedness of the 40 sequences varies considerably across the four regions. The four trees greatly differ both in their topologies and in the branch lengths found between pairs of sequences, revealing striking phylogenetic incongruences.

To test the null hypothesis that the phylogenies for each portion (HVR) of the gene were not significantly different, we compared sets of the most probable trees given by MrBayes for each HVR+. MrBayes generated by MCMC search a set of about 500 trees for each HVR+, sorted by a posterior probability ($P$) of $<1$ and with a cumulative $P$ equal to 100 (expressed as a percentage). The probability that two HVRs share the same underlying tree was then determined by multiplying their joint probabilities for all trees within the set. By this analysis, no two regions shared any common trees, so the proportion of shared trees for any of the four regions with any other was zero. This indicates that the different sectors have very different phylogenetic histories. To specifically infer significant conflicts in subphylogenies inferred by the different sectors, we estimated the posterior cumulative probability at each region that two given sequences form a cluster (the $P$ value for a given clade at one sector is the sum of the posterior probabilities given by all the trees in a set inferring that clade). Comparison of the $P$ values given by each HVR provides a simple way to support incongruences of relationships. For instance, a discordance among phylogenies is highly significant when the $P$ values for a given cluster in two different HVRs are, respectively, equal to 100 and 0.

Comparison of the four tree topologies shows significant shuffling of the HVRs occurring within a single supergroup, as well as between superfamilies (see both Fig. 2 and 4 for comparison). Some of the examples previously mentioned in terms of amino acid motifs are highlighted in the phylogenetic analysis (Fig. 4). As can be seen, they show highly supported clusters at some HVRs and divergence at others. The two examples below show recombination between superfamilies A and B.

Example 1. Within supergroup A, sequence 7-PherA2 strongly clusters with sequence 8-FexA at HVR1+ and HVR2+ ($P = 100$), while at HVR3+ the two sequences diverge. Sequence 7-PherA2 now clusters with sequences 5-CuncA, 6-ScerA, 9-BhroA, and 10-DsimA ($P = 71$), and sequence 8-FexA forms a strong monophyletic group with sequences 3-EhorA, 4-BhovA, 26-PsiaA1, 33-EcauA, and 34-TqueA ($P = 100$). Then, at HVR4+, sequence 7-PherA2 again diverges from the previous cluster while strongly grouping with sequence 6-ScerA and, interestingly, with a sequence from supergroup B, 21-BnjpB ($P = 99$). Sequence 8-FexA remains phylogenetically close to sequence 26-PsiaA1.

Example 2. HVR1+ and -2+ show a consistent grouping of sequences 30-AvulB, 29-PpruB, and 28-PspiB (in both cases, $P = 100$). But at HVR3+, the cluster is no longer supported; 30-AvulB now forms a monophyletic group with supergroup A sequences 31-PgernA and 32-DeryA and with 16-Phum? and 25-PcwB ($P = 100$). Again, at HVR4+, 30-AvulB radically diverges and significantly groups with sequence 29-PpruB ($P = 100$), even if the two are separated by a relatively high level of genetic divergence (as indicated by the branch length leading to 30-AvulB).

Noticeable also is the shift in phylogenetic relationships across the HVRs of nematode sequences from superfamilies C and D with respect to each other and relative to arthropod Wolbachia sequences. Specifically, the two superfamilies are phylogenetically close at HVR1+ and -2+ but significantly diverge at HVR3+ and -4+, where each of the two superfamilies significantly clusters with different arthropod sequences. This discordance can be seen also in the alignment in Fig. 2.

Another interesting example of amino acid shuffling within wsp appears to have occurred among several superfamilies, including recently proposed supergroup F (containing Wolbachia bacteria that infect termites). The wsp sequences from termites have not been previously described. The two sequences analyzed, 39-ClacF (from Coptotermes lacteus) and 40-CaciF (from Coptotermes acinaciformis), strongly cluster across all four HVRs while considerably varying in their relatedness with respect to the other sequences. They form a phylogenetically distant clade at HVR1+ and -2+, but branch lengths decrease at HVR3+, accompanied by a relatively close association with the two sequences from nematode Wolbachia bacteria of supergroup D, 37-WbanD (from Wuchereria bancrofti) and 38-BmalD (from Brugia malayi). At HVR4+, sequence 20-PexiB (from the hemipteran Paromius exigus), which clustered with sequences 18-EpurB and 19-AscB at HVR1+ ($P = 100$) and -2 ($P = 100$), is now close to those of termites ($P = 80$), which together are still relatively close to sequences from supergroup D in nematodes ($P = 80$). The $P$ value that sequence 20-PexiB
clusters with sequences 18-EputB and 19-AencB at HVR3+ and HVR4+ is, in both cases, equal to 0. The conflict in relationships inferred by the diverse HVRs among termites and members of supergroups D and B suggests a common origin of HVR4 for the three supergroups, possibly due to intragenic recombination. Indeed, visual examination of the HVR4 protein sequence (Fig. 2) suggests a common motif shared among the two group D nematode Wolbachia sequences and insect sequences 39-ClacF, 40-CaciF, 2-CglaA, 14-LwhiA, and 20-PexiB. The above relationships are also supported by parsimony analyses of HVR4+, which groups these sequences in a single cluster with a bootstrap value of 98% (although the same cluster is not supported by a posterior probability of >0.5 in a Bayesian analysis of HVR4+).

As underlined by comparison of branch lengths among trees, several of the above sequences that shift position from one cluster to another across the HVRs (within supergroups A and B and between them) still show high nucleotide sequence identities (>95%) within both clusters. This is unlikely to be due to divergent evolution of sequences along the gene. Instead, it strongly indicates recombination among a set of wsp sequence portions. The remarkable nucleotide sequence conservation among motifs shared by distinct sequences also suggests either relatively recent horizontal DNA transfers or pressure for nucleotide sequence conservation acting on the recombined segments at synonymous sites subsequent to intragenic exchange.

**Diversifying selection at the HVRs.** We further investigated whether the HVRs are experiencing stabilizing or diversifying selection at the amino acid level by comparing rates of synonymous versus nonsynonymous substitutions ($K_s$ and $K_a$). Previous studies have used the program PAML and detected evidence of diversifying selection in wsp, particularly within HVR1 to -3, whereas HVR4 was only partially included or excluded from the analyses due to alignment issues (2, 28).

However, these analyses did not take into consideration recombination within the wsp gene. Here we augment those earlier studies by using an independent-contrast approach (13). We compare independent sets of closely related sequences at each HVR (based on the phylogenetic analysis) for rates of $K_s$ and $K_a$ (Table 2). This approach avoids problems of recombination and alignment by analyzing only closely related sequences and evaluates them for evidence of diversifying versus stabilizing selection. Although sample sizes for independent contrasts are relatively small within each HVR, a significantly higher $K_a$ was found in HVR1 (Wilcoxon $W = 21$, mean $K_a - K_s = 0.0256$ to 0.005, $P < 0.05$, $n = 6$) and HVR4 ($W = 21$, mean $K_a - K_s = 0.022$ to 0.000, $P < 0.05$, $n = 6$), and the same trend was found in the other two HVRs, though differences were not significant. Pooling across all HVRs, we analyzed a total of 31 sets of independent contrasts, finding a strong pattern of elevated rates of amino acid substitution. Indeed, among the 31 sets, 19 showed $K_a > K_s$, 11 showed $K_a = K_s$, and only one set showed $K_a < K_s$ (Wilcoxon $W = 198$, mean $K_a - K_s = 0.0181$ to 0.002, $P < 0.02$, $n = 20$). This is further evidence that the HVRs of wsp are subject to strong diversifying selection. All the analyzed sets of paired contrast sequences at single HVRs differ only by single nucleotide polymorphisms, and there were no insertions or deletions (indels). This suggests that the primary engine for early variation at HVR motifs are single nucleotide substitutions.

### TABLE 2. Average $K_a$ and $K_s$ among closely related sequences at each HVR motif

<table>
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<th>Group</th>
<th>Sequences</th>
<th>$K_a$</th>
<th>$K_s$</th>
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<tbody>
<tr>
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<td>a</td>
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<tr>
<td></td>
<td>b</td>
<td>3:4;5;6</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>c</td>
<td>7:8</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>d</td>
<td>12:13</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>e</td>
<td>18;19;20;21;22;23</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>f</td>
<td>24;25</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>g</td>
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<tr>
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<td>h</td>
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<tr>
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<tr>
<td></td>
<td>h</td>
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**GROUPS**

- Groups were extrapolated from phylogenies of HVR+s in Fig. 4. Only very closely related groups of sequences, i.e., showing $K_a$ and $K_s$ of $<0.06$, were considered. All groups but one (indicated with an asterisk), show $K_a > K_s$.
- Numbers refer to sequence codes in Table 1.
- Estimation is based on the formula of Nei and Gojobori (39a).

### DISCUSSION

Recombination in the wsp gene has implications for (i) potential functional and evolutionary interactions of this surface protein with the host cytoplasm and (ii) the widespread use of wsp for determining phylogenetic relationships among Wolbachia bacteria.

In recent years, new insights into mechanisms shaping prokaryotic genomes and their molecular evolution have highlighted a fluidity among microbial genomes associated with lateral gene transfer and recombination events (30, 44). Consistent with these new findings, we have found evidence of extensive recombination in the Wolbachia surface protein encoded by wsp, which results in shuffling of HVR motifs. We can assume that these recombinant proteins have been selectively favored, and previous analyses of synonymous and nonsynonymous substitutions across wsp support this view (2, 28).

Recombination provides an effective means for introducing variability in bacterial genomes. More specifically, homologous
intraspecific and intragenic recombinations have important implications for gene evolution. While intragenic recombination only transfers a gene type to a different background genome (giving rise to a new “genome type”), intragenic recombination may create new variants of the gene and promote the evolution of novel phenotypes through rearrangement of sequence combinations (46). If the two recombinant segments are highly divergent in portions of their sequences, intragenic recombination can be responsible for a dramatic change in the protein sequence. For instance, homologous recombination involving target amino acid motifs, which does not disrupt the correct functioning of the protein, could represent a powerful engine for protein innovation, providing otherwise “clonal” bacteria with a tool for counteracting the effects of slow accumulation of mutations, thereby escaping Muller’s ratchet (38, 39). The importance of recombinational events depends, however, on the selective advantages introduced into the novel product.

The complex pattern of recombination detected by the present study in wsp suggests a long history of recombination for the gene, which has led to a marked mosaicism of its representative sequences. Analyses of the four HVRs of wsp in the selected data set revealed a strict conflicting pattern of similarities and differences among sequences, leading to a clear partitioning of the alignment into segments with incongruent phylogenetic relationships. High posterior probabilities and bootstrap values support the shuffling of amino acid motifs within the four HVRs through horizontal DNA transfer events. The presence of a high number of polymorphisms and indels in the HVRs and the limited set of HVR motifs make it highly unlikely that the observed mosaic pattern has been simply shaped by random substitutions, via homoplastic events. Since two paralogs of wsp (wspB and wspC) have been recently annotated in the wMel genome, we also evaluated the hypothesis that these genes could represent a source of sequence fragments for recombination in wsp. However, no similarity was found between any wsp HVRs and the two paralog gene sequences. Furthermore, a BLAST search of the HVRs of the wsp sequence from wMel against the whole wMel genome but wsp resulted in no significant hits. These results appear to exclude a potential role of intragenomic recombination (with either other surface proteins or different portions of the genome) in shaping wsp.

Recombination in wsp involves both CRs and HVRs. However, recombination affecting CRs appears to have a minor effect on the amino acid structure of the gene, being masked by the high protein conservation between recombining segments in these regions. The gene rearrangement affects to a greater extent the protein sequence of HVRs.

Previous studies using the program PAML indicate that the HVRs have been subject to strong positive selection, with ratios of dN/dS of $\gg$1 (2, 28). Recombination may not necessarily invalidate these previous results since the phenomenon would basically work by recombining a preexisting variability. However, using a different approach in this study, we were able to show elevated levels of amino acid substitutions relative to synonymous substitutions in closely related HVR sequences, thus avoiding the problems of recombination and alignment. Furthermore, this approach allowed us to demonstrate positive selection in the whole of HVR4, which was only partially inferred in previous analyses due to problems of aligning more distantly related sequences. It is worth noting that similar sequences within a motif of an HVR are typically found in different insect species, often in different orders of insects. Therefore, it is unclear whether selection for amino acid changes is a result of adaptation to new host environments, of antagonistic coevolution between the host and wsp HVRs, or of some combination of these two processes. Overall, both recombination and diversifying selection appear to be responsible for the extensive divergence among wsp sequences.

As an outer membrane protein-encoding gene, wsp shows sequence similarity with genes coding for the major surface proteins of Rickettsiales bacteria (9). A BLAST search in the Conserved Domain Database (available at the National Center for Biotechnology Information) identified significant domain homology between the wsp product and proteins that mediate various pathogen-host cell interactions from several pathogenic proteobacteria (e.g., Neisseria species). The genetic similarity of wsp with these genes is restricted to motifs at CRs, while the HVRs of wsp do not show any significant homology with any sequence in GenBank (data not shown). This could suggest a basic structural role for the conserved motifs and distinct functional roles for the HVRs among the different surface proteins. The function of wsp is unknown, and its three-dimensional molecular structure, as well as its precise location in the outer membrane, has yet to be characterized. However, wsp was found to be the most abundant protein expressed by infected Drosophila eggs (9), suggesting a potentially strong influence of the protein and its surface domains in host-bacterium interactions (27, 66).

Intragenic recombination has been frequently found to affect the outer membrane protein-encoding genes of several parasitic bacteria, i.e., the intimin genes of Escherichia coli (36), ospC and ospD of Borrelia burgdorferi (24, 33, 45), and recently ompL1 of the genus Leptospira (19). Similar to the recombinant pattern found for wsp, in leptospiral ompL1 recombination involves four variable regions encoding surface-exposed loops whose variants may represent specific adaptations to host environmental constraints. Regarding wsp, it will be interesting to determine what host proteins bind to the HVR domains and whether these proteins have been subject to divergent selection in infected host species.

The wsp gene has been widely used to identify phylogenetic associations among Wolbachia strains (14, 25, 41, 49, 51, 68). In addition, a nucleotide sequence divergence of 2.5% among wsp sequences has been used to infer subgroup affiliation and to identify novel lineages (25, 29, 49, 68). Because of the high level of intragenic recombination within wsp, use of this gene for phylogenetic reconstructions could be misleading. For instance, Kikuchi and Fukatsu (29) proposed a new subgroup of Wolbachia bacteria using the wsp sequence from the heteropteran Elasmosticta putoni (18-EputB). However, as can be seen in our analysis, the wsp gene from this strain contains no unique elements but rather portions of different existing wsp sequences characterized by high nucleotide sequence similarities. An example of partitioned similarity in this sequence is its similarity with sequence 19-AencB at HVR1 and -2, 24-DinnB at HVR3, and 25-PcouB at HVR4 (Fig. 2 and 4). Therefore, although the combination of HVR motifs appears to be novel (thus leading to its appearance as a new subgroup), its HVR motifs are not. The bacterium may well be divergent, but its
phylogenetic status cannot be inferred simply based on wsp. The above example suggests that caution should be taken to avoid false subgroup affiliation of strains and confusion between novel “Wolbachia lineages” and new “wsp recombinant genotypes.” Phylogenetic relationships among Wolbachia strains can be potentially clarified through comparative analyses of different portions of the genomes (such as analysis of the flanking regions of wsp and different genes).

Our results strongly suggest that recombination has occurred not only within supergroups but also between them. Nucleotide identities of motifs shared by sequences from supergroups A and B are in some cases very high, as in Sitotroga cerealella, Pegoscapus herrei, and Blastophaga nipponica, sharing 100% nucleotide sequence identity at HVR4. Such a high homology in an HVR strongly suggests recent horizontal DNA transfer between supergroups A and B. An interesting transfer of motifs at HVR4 may also have occurred between sequences from the heteropteran species Paromius eugius and those from termites (supergroup F).

As a result, the shuffling among HVRS leads to a strong alteration of the phylogenetic signal, which not only affects the relative relatedness of the strains within a supergroup but can also weaken the major taxonomic organization of the genus into supergroups.

wsp has likely undergone both intragenic and intergenic recombination (lateral gene transfer). It is well known that Wolbachia strains experience frequent horizontal transmission, even if in some cases demonstrations are based on incongruences between wsp and host phylogenies, which leads to some circularity. However, horizontal movement of Wolbachia bacteria has also been detected using much more conserved genes such as ftsZ (63) and 16S (40, 63). It is clear that horizontal transmission of Wolbachia strains cannot be inferred solely on the basis of the wsp gene, since lateral gene transfer, rather than bacterial transfer, could be responsible. All three of these phenomena (intragenic and intergenic recombination and horizontal transmission) have the major effect of producing artificial phylogenies. For this reason, wsp phylogeny alone should no longer be considered to represent bacterial strain phylogeny, to invoke discrepancies between Wolbachia and host systematics (51, 68), or as a reference phylogeny to support potential lateral gene transfer in comparative gene phylogenies (7). For instance, the reported incompatibility between wsp and ftsZ phylogenies (27) could instead be explained in light of intragenic recombination in wsp rather than lateral transfer of the ftsZ gene.

The impact of recombination on Wolbachia genomes is still to be clarified. As recently reported, the complete genome of wMel encodes the necessary apparatus for recombination, including RecA, and shows unusual features likely to be derived from frequent intragenomic recombination and lateral DNA transfers (67). The results of our study indicate a greater impact of recombination in the Wolbachia genome than previously appreciated. However, we do not know how unique wsp is with regard to intragenic recombination. The high number of distinct recombinant sequences detected for wsp and the complex patterns of recombination affecting some of them suggest that contact of DNA among different strains has occurred quite frequently within the Wolbachia genus. Mechanisms leading to contact and exchange of DNA among distant strains are still poorly understood, although multiple infections in the same hosts provide one obvious avenue (7, 66).

The implications of widespread recombinant are clearly of great interest. Over the long term, the phenomenon disrupts bacterial clonal history and obfuscates the understanding of microbial evolution based on sequence comparisons (16). However, it also provides a potential motor for evolutionary change and the acquisition of new mechanisms among bacteria. The patterns of recombination in Wolbachia genomes could clarify important aspects of the evolution of this host-symbiont system, such as the evolution of similar phenotypes (e.g., host feminization, parthenogenesis induction, or male killing) among distantly related strains and the long persistence of these widespread parasitic bacteria in their invertebrate hosts.

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