

# Structural Insight into How Bacteria Prevent Interference between Multiple Divergent Type IV Secretion Systems

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**ABSTRACT** Prokaryotes use type IV secretion systems (T4SSs) to translocate substrates (e.g., nucleoprotein, DNA, and protein) and/or elaborate surface structures (i.e., pili or adhesins). Bacterial genomes may encode multiple T4SSs, e.g., there are three functionally divergent T4SSs in some *Bartonella* species (*vir*, *vbh*, and *trw*). In a unique case, most rickettsial species encode a T4SS (*rvh*) enriched with gene duplication. Within single genomes, the evolutionary and functional implications of cross-system interchangeability of analogous T4SS protein components remains poorly understood. To lend insight into cross-system interchangeability, we analyzed the VirB8 family of T4SS channel proteins. Crystal structures of three VirB8 and two TrwG *Bartonella* proteins revealed highly conserved C-terminal periplasmic domain folds and dimerization interfaces, despite tremendous sequence divergence. This implies remarkable structural constraints for VirB8 components in the assembly of a functional T4SS. VirB8/TrwG heterodimers, determined via bacterial two-hybrid assays and molecular modeling, indicate that differential expression of *trw* and *vir* systems is the likely barrier to VirB8-TrwG interchangeability. We also determined the crystal structure of *Rickettsia typhi* RvhB8-II and modeled its coexpressed divergent paralog RvhB8-I. Remarkably, while RvhB8-I dimerizes and is structurally similar to other VirB8 proteins, the RvhB8-II dimer interface deviates substantially from other VirB8 structures, potentially preventing RvhB8-I/RvhB8-II heterodimerization. For the *rvh* T4SS, the evolution of divergent VirB8 paralogs implies a functional diversification that is unknown in other T4SSs. Collectively, our data identify two different constraints (spatio-temporal for *Bartonella trw* and *vir* T4SSs and structural for *rvh* T4SSs) that mediate the functionality of multiple divergent T4SSs within a single bacterium.

**IMPORTANCE** Assembly of multiprotein complexes at the right time and at the right cellular location is a fundamentally important task for any organism. In this respect, bacteria that express multiple analogous type IV secretion systems (T4SSs), each composed of around 12 different components, face an overwhelming complexity. Our work here presents the first structural investigation on factors regulating the maintenance of multiple T4SSs within a single bacterium. The structural data imply that the T4SS-expressing bacteria rely on two strategies to prevent cross-system interchangeability: (i) tight temporal regulation of expression or (ii) rapid diversification of the T4SS components. T4SSs are ideal drug targets provided that no analogous counterparts are known from eukaryotes. Drugs targeting the barriers to cross-system interchangeability (i.e., regulators) could dysregulate the structural and functional independence of discrete systems, potentially creating interference that prevents their efficient coordination throughout bacterial infection.

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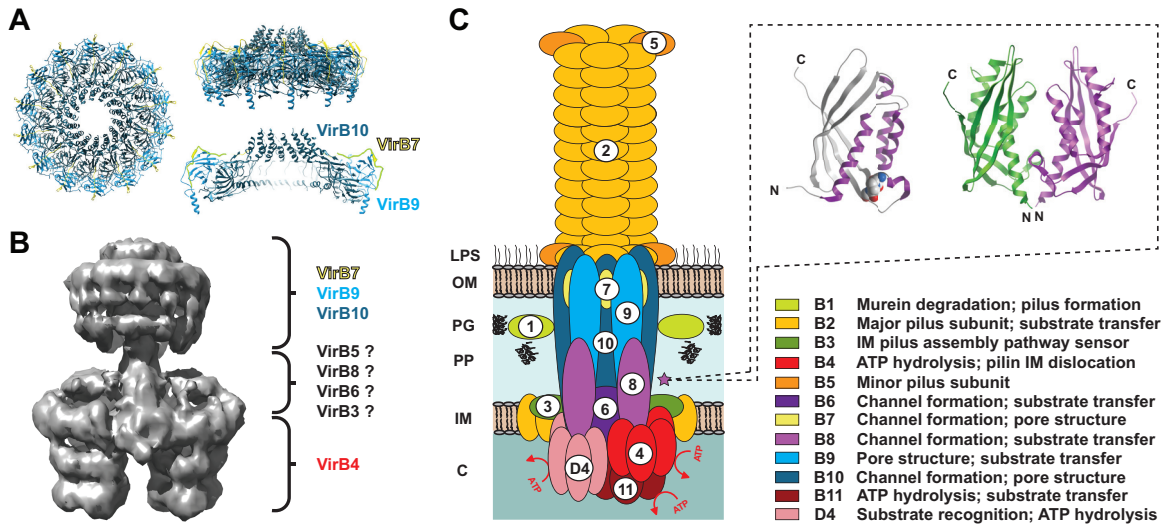
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Occurring in Gram-negative, Gram-positive, and wall-less bacteria, as well as archaea, type IV secretion systems (T4SSs) are primarily utilized for translocating substrates across the cell envelope (1, 2). T4SSs that translocate plasmid (3) and naked DNA (4, 5), as well as genomic islands (6), are major facilitators of bacterial diversification, contributing to the spread of antimicrobial resistance and virulence genes. Other T4SSs translocate nu-

cleoprotein (e.g., oncogenic T-DNA via the *vir* T4SS of *Agrobacterium tumefaciens*) or protein substrates directly into eukaryotic cells, wherein these effectors often perturb cell signaling to benefit bacterial survival (7, 8). Recently, it was reported that T4SSs may also be utilized to kill neighboring bacteria via translocation of a proteinaceous effector (9). Typically, T4SSs elaborate surface structures (i.e., pili or specialized adhesins) in a process either



**FIG 1** Architecture of P-type type IV secretion systems (P-T4SSs). (A) Bird's-eye (left) and side (right) views of the dodecameric P-T4SS core complex (CC) encoded by plasmid pKM101 of *Escherichia coli* (PDB ID 3JQO), adapted from the work of Chandran et al. (17). Colors for the CC subunits (VirB7, VirB9, and VirB10) are similar to the model in panel C. (B) Negative-station electron microscopy-generated structure of the P-T4SS encoded by the *E. coli* R388 conjugative plasmid (EMD-2567) (adapted from the work of Low et al. [19]). Colors for the CC subunits and cytosolic/IM barrels (VirB4) are similar to the model in panel C. The putative positions of other IM channel (IMC) proteins are depicted with question marks. VirB1, VirB2, VirB11, and VirD4 are not shown, as they were not included in the original structure. (C) General model of the composition of P-T4SSs, with functions for all 12 components (VirB1 to VirB11 and VirD4) listed at bottom right. The purple star depicts the bitopic VirB8 IMC proteins, with a dashed box illustrating the monomeric (left) and dimeric (right) structures for VirB8 C-terminal domains of *Agrobacterium tumefaciens* (PDB ID 2CC3) (44).

coupled to or independent of substrate translocation (10, 11). Thus, T4SSs are extraordinarily diverse in function relative to other bacterial secretion systems (12, 13).

Recently, T4SSs have been classified into eight major groups (14). One well-studied group, P-T4SSs, is typified by the *vir* T4SS of the pTi plasmid of *A. tumefaciens*, which encodes 11 scaffold components (VirB1 to VirB11) and a coupling protein (VirD4) that recruits substrates to the secretion channel (15). Tremendous architectural insight has been garnered from recent macromolecular structures generated for other P-T4SSs (16–20), allowing many of the various scaffold components to be arranged into an anatomical model (Fig. 1). Phylogenomics and bioinformatics efforts have shown that the other seven groups of T4SSs contain homologs or analogs to some (or most) of the components of this anatomical model (21), implying that a common T4SS architecture links the prokaryotic cytoplasm to the extracellular milieu. Variations on this theme, particularly regarding the distal region of the translocation channel, are thought to primarily be a consequence of coevolution with cell envelope morphology (22, 23). However, many structural innovations likely correlate with specific functions, e.g., mating pair stabilization during conjugation (24, 25) or pilus interactions with host cells during effector translocation (26).

Bacterial genomes may encode multiple divergent T4SSs, e.g., the *cag* and *com* P-T4SSs of *Helicobacter pylori* (27, 28), the *vir*, *vbh*, and *trw* P-T4SSs of certain *Bartonella* species (29), and the *dot/icm* (I-T4SS) and *lvh* (P-T4SS) T4SSs of *Legionella* species (30, 31). These divergent T4SSs typically have different functions, though cross-system interchangeability is known for *L. pneumophila dot/icm* and *lvh* T4SSs. The *lvh* P-T4SS, which is dispensable for *L. pneumophila* replication in both amoebae and macrophage hosts (32) but causes virulence phenotypes under conditions that

mimic the *L. pneumophila* aquatic phase (33), was shown to complement certain *dot/icm* mutants defective in conjugation (32). Furthermore, arrested virulence phenotypes in an *L. pneumophila dotA/lvh* double mutant were restored via complementation with the *lvh* coupling protein LvhD4 (34), implying structural and functional resilience in the face of extreme sequence divergence across *dot/icm* and *lvh* T4SSs. For other bacteria harboring multiple divergent T4SSs, studies on cross-system interchangeability are lacking, leaving a limited understanding of how distinct T4SSs achieve correct spatiotemporal assembly to execute their specific functions. Such precise regulatory mechanisms must exist, given also that many bacterial genomes are continually bombarded with integrative conjugative elements that often carry T4SS loci (35). If not quickly purged, it is likely that incoming T4SSs with compatibility to the native T4SS(s) either undergo rapid diversifying evolution or acquire mechanisms for differential regulation.

To gain further insight into T4SS cross-system interchangeability, we selected two bacterial genera with multiple divergent analogs of VirB8, a channel protein required for substrate transfer (Fig. 1). First, two or three distinct complete T4SSs are found within the genomes of some *Bartonella* species. Many of these species encode *vir* and *trw* secretion systems, which are highly divergent in sequence and function (36). Some species have an additional *vbh* T4SS homologous to the *vir* T4SS (29), wherein the biological function still remains elusive. The *Bartonella vir* T4SS is involved in protein secretion, with an arsenal of *Bartonella* effector proteins (Beps) translocated into host cells during infection (8). The *trw* T4SS is not involved in protein translocation but rather elaborates variable surface pili comprised of different combinations of duplicated VirB2 (TrwL) and VirB5 (TrwJ) homologs (37), with such structures shown to interact with host erythrocytes (38, 39). While these divergent T4SSs are differentially expressed

during *Bartonella* host cell infection (40), the potential for cross-system interchangeability is currently unknown. Second, for species of *Rickettsiales*, an entirely different strategy has evolved with a single T4SS, the *Rickettsiales* vir homolog (*rvh*), which contains duplications of several components, including a VirB8 homolog (RvhB8) (23, 41). The functional significance of these duplicate *rvh* components is unknown, particularly regarding whether a single *rvh* T4SS functions during the rickettsial life cycle or if multiple different T4SSs are assembled throughout the complex life cycle (usually involving arthropod and vertebrate hosts) (42).

Herein we report crystal structures for six proteins of the VirB8 family: three VirB8 and two TrwG proteins of several *Bartonella* species, as well as the first crystal structure of an *rvh* scaffold component, RvhB8-II of *Rickettsia typhi*. Comparative structural analysis, in conjunction with biochemical assays, protein modeling, and bioinformatics, leads us to propose two distinct mechanisms (spatiotemporal for *Bartonella trw* and *vir* T4SSs and structural for *rvh* T4SSs) that prevent cross-system interchangeability between multiple divergent T4SSs encoded within single bacterial genomes.

## RESULTS

**Origin of multiple VirB8 proteins in *Bartonella* and *Rickettsia* genomes.** To understand the origin of multiple VirB8-like proteins within *Bartonella* and *Rickettsia*, we estimated a phylogeny of selected P-T4SSs. The *Bartonella vir* and *vbh* T4SSs share common ancestry, probably arising from whole-system duplication (Fig. 2). Previously, the *Bartonella vir* and *vbh* T4SSs were hypothesized to originate via lateral gene transfer (LGT), as they are absent from the ancestral human pathogen *Bartonella bacilliformis* (29, 36). The *Bartonella trw* T4SSs are found in a different clade, which includes plasmid-encoded T4SSs of gammaproteobacterial species (Fig. 2). Thus, the *Bartonella trw* T4SSs were likely acquired via LGT from non-alphaproteobacterial sources, becoming incorporated into the chromosomes, where they evolved to mediate adhesion to host erythrocytes (29, 36).

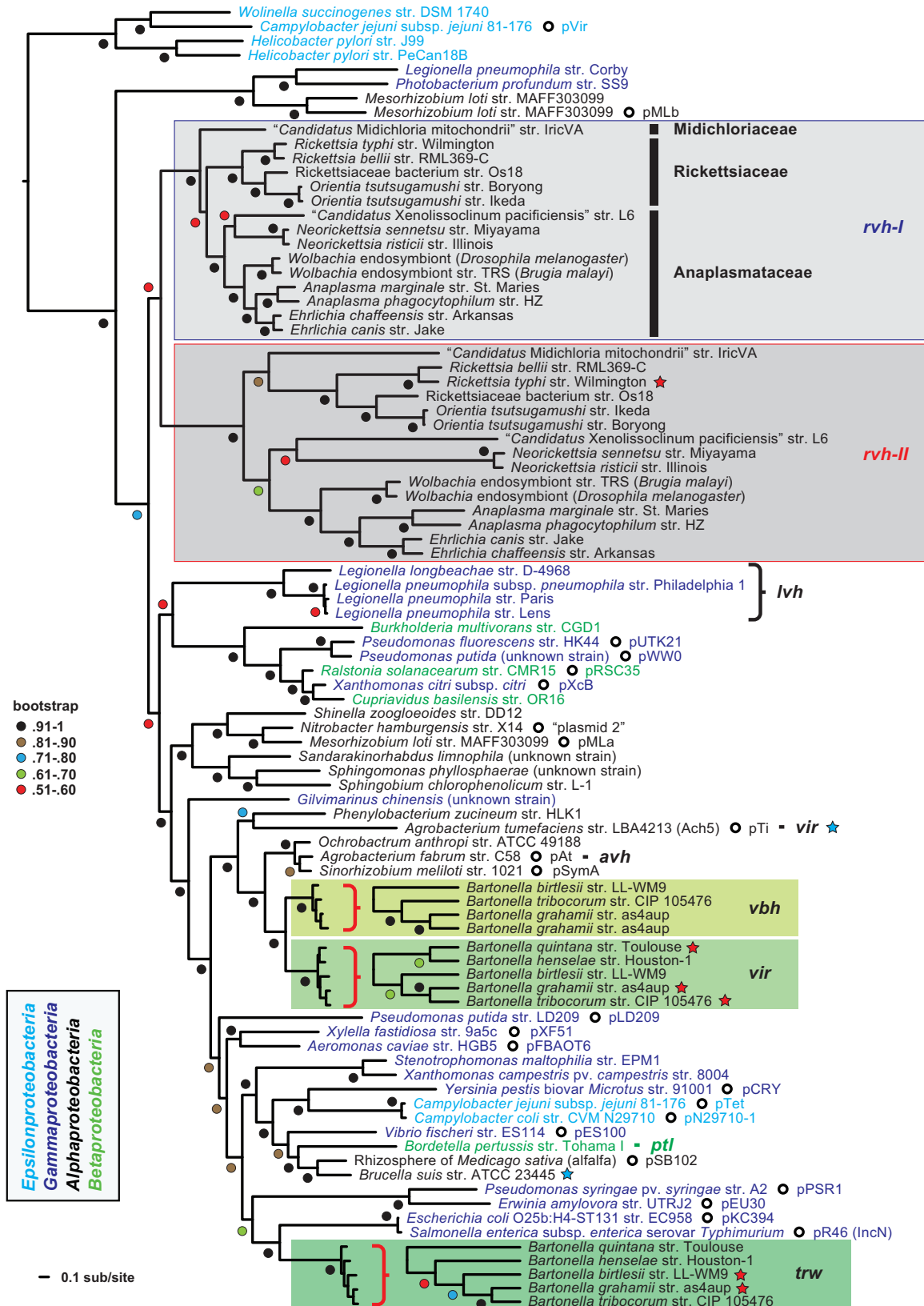
The *rvh* T4SS forms a lineage distinct from other P-T4SSs, though robust support for the relationship of this lineage to other P-T4SSs is lacking (Fig. 2; also see Text S1 in the supplemental material). We considered the paralogous genes encoding RvhB4, RvhB8, and RvhB9 to comprise different structural and functional machines, as previous phylogeny estimation supported these genes as ancient duplications that arose early in *Rickettsiales* evolution (23, 41). The genes encoding RvhB4-I, RvhB8-I, and RvhB9-I (collectively referred to as *rvh-I*) are conserved relative to analogs in other P-T4SSs and thus are anticipated to assemble into the *rvh* T4SS while it functions in protein translocation. Alternatively, the genes encoding RvhB4-II, RvhB8-II, and RvhB9-II (collectively referred to as *rvh-II*) have evolved atypical features that deviate substantially from counterparts in other P-T4SSs (see Discussion). The structural and functional significance of *rvh-II* is unknown, though a substantially higher divergence of *rvh-II* than *rvh-I* indicates different selective pressures operating on these paralogs (Fig. 2). Remarkably, this odd paralogy within the *rvh* T4SS is conserved across the three derived families of *Rickettsiales* (*Rickettsiaceae*, *Anaplasmataceae*, and “*Candidatus* Midichloriaceae”), which comprise a large, diverse assemblage of symbiotic and pathogenic species (43).

**Divergent *Bartonella* VirB8 and TrwG proteins homodimerize via interactions at an NPXG motif highly conserved across T4SSs.** We determined crystal structures of the sol-

uble C-terminal periplasmic domains of three VirB8 and two TrwG proteins of several *Bartonella* species (see Text S2 in the supplemental material). As expected by the highly conserved sequences of C-terminal periplasmic domains of *Bartonella* VirB8 (~90%) and TrwG (~90%) proteins (data not shown), the structures within each protein family are highly similar, with 5  $\alpha$ -helices and 4  $\beta$ -sheets (Fig. 3A and B). Within *Bartonella* VirB8, the pairwise root mean square deviation (RMSD) values, or measures of the average distance between the atoms of superimposed proteins, range from 0.33 to 0.60 Å for similar C $\alpha$  atoms, and within *Bartonella* TrwG, the two structures have a pairwise RMSD value of 0.88 Å. The structures of *Bartonella* VirB8 and TrwG proteins are also highly similar to each other (Fig. 3C), despite extremely low sequence conservation (~28% identity overall, ~34% identity for C-terminal periplasmic domains) (Fig. 3D; see Text S3 in the supplemental material). The pairwise RMSD values between *Bartonella* VirB8-TrwG sets range from 1.57 to 1.78 Å. The *Bartonella* VirB8 and TrwG protein structures are also similar to the *Agrobacterium tumefaciens* (44) (RMSD ranges of 1.28 to 1.33 Å and 1.52 to 1.63 Å for VirB8 and TrwG, respectively) and *Brucella suis* (45) (RMSD ranges of 1.73 to 1.78 Å and 0.98 to 1.18 Å for VirB8 and TrwG, respectively) VirB8 protein structures (Fig. 3C). This implies remarkable structural constraints for VirB8 components in the assembly of functional P-T4SSs.

With the exception of monomeric *Bartonella quintana* VirB8, all of the *Bartonella* VirB8 and TrwG proteins crystallized as dimers. Similar to the *A. tumefaciens* and *B. suis* structures, the dimerization interfaces of *Bartonella* VirB8 and TrwG structures contain major and minor dimerization sites. The major dimerization site involves helix  $\alpha$ 1 from one subunit contacting the sharp turn between helix  $\alpha$ 5 and strand  $\beta$ 4 of the other subunit (Fig. 3E). The latter motif, here referred to as the NPXG motif, is the most conserved region of proteins of the VirB8 family and was originally suggested to be critical for VirB8 dimerization (45). Bacterial 2-hybrid (B2H) experiments with selected full-length VirB8 and TrwG proteins (Fig. 3F), as well as glutaraldehyde cross-linking experiments with the corresponding soluble periplasmic domains (see Fig. S1 in the supplemental material), further indicated that a homodimer is the probable functional unit for *Bartonella* VirB8 and TrwG proteins. To assess the importance of the major dimerization site within the subunit interface, we carried out site-directed mutagenesis of *Bartonella birtlesii* TrwG to yield mutants in helix  $\alpha$ 1 (TrwG<sub>V96G/V97G</sub>) and the NPXG motif (TrwG<sub>P214A</sub>), which were utilized in a series of B2H assays (Fig. 3F). Both mutations had negligible effects on dimerization when analyzed individually with the wild-type TrwG (TrwG<sub>wt</sub>) protein (Fig. 3F). However, when both of the fusion constructs contained the mutations, we detected a significant reduction in dimerization (Fig. 3F). Thus, helix  $\alpha$ 1 and the NPXG motif are central (but not the sole) mediators of *B. birtlesii* TrwG dimerization and probably function similarly for most other VirB8-like proteins of P-T4SSs, given the high degree of conservation of these structural features.

***Bartonella* VirB8 and TrwG proteins are capable of forming heterodimers.** The strong structural conservation across *Bartonella* VirB8 and TrwG proteins suggests that individual VirB8 and TrwG monomers could form a heterodimer. A structural model of the *Bartonella grahamii* VirB8/TrwG heterodimer illustrates how such a dimer interface might form, with the overall symmetry similar to the VirB8 and TrwG homodimers (Fig. 4A). The formation of this dimerization interface is undoubtedly facilitated by the



**FIG 2** Phylogeny estimation of P-type type IV secretion systems (P-T4SSs). Phylogeny was estimated from concatenated alignments of five components (VirB4 and VirB8 to VirB11), except for *rvhB-II*, which contains homologs to VirB4, VirB8, and VirB9 only (see the text for further details on alignment and data set construction). ML-based phylogeny was estimated with RAxML on the unmasked alignment (LG + gamma + I). Branch support was assessed with 1,000 bootstrap pseudoreplications. A tree with specific bootstrap values, as well as additional phylogenies estimated from alternative alignments/models/optimality

(Continued)

high degree of sequence conservation of contacting residues within helix  $\alpha 1$  and the NPXG motif, four of which are invariant across VirB8 and TrwG sequences (Fig. 4B). Using the B2H assay, we analyzed the potential cross-system interchangeability between *B. grahamii* VirB8 and TrwG (Fig. 4C). We initially established a reference of interaction strength for *B. birtlesii* and *B. grahamii* TrwG homodimers, as well as for the *B. grahamii* VirB8 homodimer. TrwG interaction signals were much higher than VirB8 homodimerization, possibly reflecting the known biological functions of these divergent systems, i.e., rigid adhesive function of the *trw* T4SS compared to dynamic effector translocation of the *vir* T4SS (36). Subsequently, we evaluated the potential for heterodimeric interactions between *B. grahamii* VirB8 and TrwG proteins. These assays indicate the ability of VirB8 and TrwG to form heterodimers (Fig. 4C), with a strength of interactions not significantly lower than that observed for the *B. grahamii* VirB8 homodimer. Thus, it appears that in the absence of spatiotemporal barriers (e.g., differential expression), VirB8 and TrwG proteins have the potential to interfere with one another during assembly into their respective T4SSs.

**Duplicate, coexpressed VirB8 proteins encoded within the *Rickettsia typhi* genome are structurally divergent.** To lend insight into the duplicate VirB8-like proteins encoded within *Rickettsia* genomes, we set out to determine the crystal structures for these divergent proteins from the murine typhus agent, *R. typhi* strain Wilmington (RvhB8-I and RvhB8-II, which share only 18% amino acid identity). Attempts to generate a crystal structure for RvhB8-I failed; thus, we modeled both the monomer and dimer of this protein by homology, using a multitemplate approach (Fig. 5A). Because RvhB8-I contains residues conserved in other VirB8 family proteins, such as the NPXG motif (41), its sequence could be reasonably modeled to the *Bartonella* spp., *B. suis*, and *A. tumefaciens* VirB8 structures, as well as the *Bartonella* TrwG structures. Thus, the RvhB8-I structure is anticipated to fold similarly to these proteins. The protomeric structure of *R. typhi* RvhB8-I is quite similar to that of *Bartonella* VirB8 and TrwG proteins, with pairwise RMSD values of 1.46 to 1.64 Å. In contrast, the biological dimeric structure determined for *R. typhi* RvhB8-II deviates substantially from the RvhB8-I model and all seven VirB8/TrwG dimeric structures, with pairwise RMSD values of 3.20 to 3.79 Å (Fig. 5B). We previously observed that the lack of conservation in the NPXG motif of RvhB8-II, coupled with rapidly evolving sites throughout the protein, indicated a possible divergent structure and function for this protein (41). All other VirB8 and TrwG T4SS proteins with known dimeric structures exhibit the NPXG motif posterior to helix  $\alpha 5$  that interacts with helix  $\alpha 1'$  of the other protomer. The RvhB8-II sequence differs substantially in this region, and in the structure, this region does not form helix  $\alpha 5$ . Part of this region is disordered in the crystal structure; one potential explanation could be *in situ* cleavage of this region during crystallization, although mass spectrometry analysis of the protein stock solution revealed an intact protein (~21 kDa) prior to crystallization (see Text S2 in the supplemental

material). The divergent sequence and lack of helix  $\alpha 5$  perturbs the major dimerization site to generate an asymmetrical subunit interface, unlike other VirB8 and TrwG structures that had symmetric dimers. To compensate for disruption of the major dimerization site, a minor dimerization site is observed involving residues between  $\beta$ -strands  $\beta 2$  and  $\beta 3$  (Fig. 5C). Glutaraldehyde cross-linking experiments with the soluble periplasmic domains of RvhB8-I and RvhB8-II demonstrated that both proteins form homodimers in solution (see Fig. S1 in the supplemental material). Thus, despite a divergent dimeric structure adopted by RvhB8-II, both VirB8-like proteins of *R. typhi* appear to form functional dimers.

The significance of duplicate VirB8-like proteins carried by all species of *Rickettsia* is unclear. The arrangement of *rvhB8-I* and *rvhB8-II* in the *R. typhi* genome is odd (46), with the genes found in adjacent operons that encode other *rvh* genes (Fig. 5D). The conservation of these loci across all sequenced *Rickettsia* genomes implies a conserved function (41). Reverse transcription-quantitative PCR (RT-qPCR) analysis of *R. typhi* throughout one day of host cell infection indicates that *rvhB8-I* and *rvhB8-II* genes are simultaneously expressed (Fig. 5E). Whether RvhB8-I and RvhB8-II both assemble into the same *rvh* secretion machine remains unknown. A modeled RvhB8-I/RvhB8-II heterodimer suggests that such an interaction would be highly divergent from other VirB8 and TrwG homodimers, as well as the VirB8/TrwG heterodimer (see Text S4 in the supplemental material). Aside from an entirely skewed major dimerization site, there are no residues contacting across the minor dimerization site within the modeled heterodimer. Supporting these observations, B2H assays failed to detect interactions between RvhB8-I and RvhB8-II (data not shown). Furthermore, the N-terminal cytoplasmic and transmembrane-spanning (TMS) regions of RvhB8-I and RvhB8-II are extraordinarily divergent from one another (41). Collectively, if RvhB8-I and RvhB8-II both simultaneously assemble into one *rvh* T4SS, there is little support for these proteins forming heterodimers.

**Divergent RvhB8-II proteins are conserved across species of *Rickettsiales* but not found in other bacteria.** The presence of two VirB8-like genes in all species of *Rickettsiales* that carry the *rvh* T4SS prompted us to evaluate if the odd *R. typhi* RvhB8-II characteristics were present in other RvhB8-II proteins. Analysis of selected rickettsial species revealed that RvhB8-I and RvhB8-II proteins encoded within the same genome never share more than 19% amino acid identity (Fig. 6A, yellow highlighting). Across species within the six major rickettsial genera, RvhB8-I is more conserved relative to RvhB8-II (Fig. 6B). This is further witnessed by phylogeny estimation of RvhB8 proteins, which shows substantially higher divergence of RvhB8-II than RvhB8-I (Fig. 6C). Collectively, these data indicate that an ancient duplication of *rvhB8* quickly led to diversification of *rvhB8* paralogs under different selective constraints. Considering these results with the similar divergence patterns witnessed for RvhB4 and RvhB9 paralogs

#### Figure Legend Continued

criteria, is provided in Text S1 in the supplemental material. The tree was rooted with four P-T4SSs from species of *Epsilonproteobacteria*. *rvh-I* and *rvh-II* T4SSs are in light and dark gray, respectively. The *Bartonella vir*, *vbh*, and *trw* T4SSs are colored different shades of green. Red stars depict sequences used to generate VirB8 structures in this study; blue stars depict sequences previously used to determine VirB8 structures. Plasmid-encoded T4SSs are depicted with open circles, with plasmid names provided. NCBI GenBank accession numbers for all proteins are provided in Text S1 in the supplemental material.

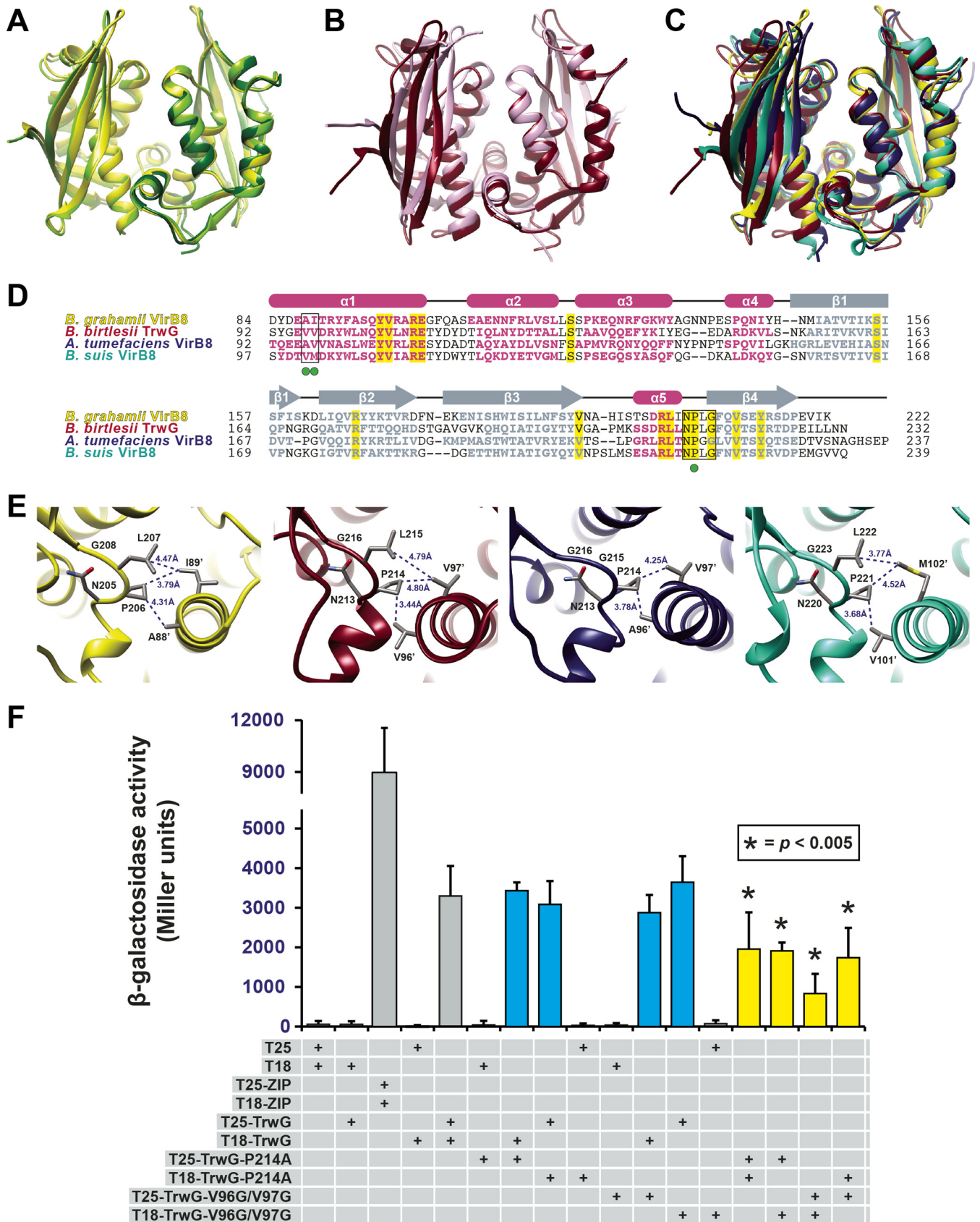


FIG 3 *Bartonella* VirB8 and TrwG proteins form structurally conserved homodimers. (A) Superimposition of the ribbon representations of the VirB8 crystal structures from *B. quintana* strain Toulouse (dark green) (PDB ID 4LSO), *B. grahamii* strain as4aup (yellow) (PDB ID 4KZ1) and *B. tribocorum* strain CIP 105476 (Continued)

(Fig. 2), it is probable that *rvh-I* and *rvh-II* contribute differently to the function and regulation of the *rvh* T4SS.

Comparison of full-length RvhB8 paralogs across selected rickettsial species provides further insight into the pattern of *rvhB8* diversification (Fig. 6D). Four residues within the periplasmic C-terminal domain are conserved across RvhB8-I and RvhB8-II proteins, as they are in most other VirB8 proteins. One of these residues, a Tyr within  $\beta$ -strand  $\beta_4$ , has been implicated in an interaction with VirB4 (47), suggesting that both RvhB8-I and RvhB8-II have the potential to interact with one or both of the RvhB4 paralogs. A Trp residue within the cytoplasmic domain is also conserved across RvhB8-I and RvhB8-II proteins. The functional significance of this Trp residue is unknown, though it demarcates the highly variable N-terminal sequences from a conserved cytoplasmic region proximal to the predicted TMS region (see Text S5 in the supplemental material). RvhB8-I proteins share 15 conserved residues; ten of these are found in the globular C-terminal domain and are also highly conserved across VirB8 proteins from other bacteria. In contrast, RvhB8-II proteins share only one conserved residue, a Tyr residue within the N-terminal cytoplasmic domain. Collectively, RvhB8-I proteins are conserved relative to other VirB8 proteins, while RvhB8-II proteins have retained only a minimal set of the conserved residues that define the VirB8 structural architecture.

Like *R. typhi* RvhB8-II, the other rickettsial RvhB8-II proteins contain a divergent NPXG motif (Fig. 6E). Specifically, the Asn and Gly residues are replaced with other residues that are not conserved across RvhB8-II homologs, with the Pro residue present only in ~60% of the proteins. This affects the interactions within the major dimerization site, suggesting that all RvhB8-II proteins form atypical dimer interfaces relative to the conserved VirB8 structure. To ascertain how conserved the NPXG motif is and, specifically, whether there is flexibility in this motif in other VirB8 proteins, we evaluated this region in 1,239 nonredundant VirB8 proteins. Only ~10% of these proteins contained divergent NPXG motifs, which were assembled into groups based on the primary sequence of their respective NPXG motifs (Fig. 6F). We then modeled the monomer and dimer structures for at least one representative from each of these groups (see Text S4 in the supplemental material). All of these predicted structures involved conserved contacts within the major dimerization site typical of VirB8 proteins but not RvhB8-II. Even replacement of the Pro residue with bulky aromatic residues does not alter the dimerization interface (Fig. 6G). Altogether, these observations suggest that the structure of RvhB8-II proteins is atypical relative to other VirB8 proteins, with the deviant NPXG motif skewing the dimerization interface, resulting in an asymmetrical dimer.

#### Figure Legend Continued

(light green) (PDB ID 4MEI). (B) Superimposition of the ribbon representations of the TrwG crystal structures from *B. birtlesii* strain LL-WM9 (burgundy) (PDB ID 4JF8) and *B. grahamii* strain as4aup (pink) (PDB ID 4NHF). (C) Superimposition of the ribbon representations for the crystal structures of *B. grahamii* VirB8, *B. birtlesii* TrwG, *Agrobacterium tumefaciens* strain C58 VirB8 (dark blue) (PDB ID 2CC3) (44), and *Brucella suis* biovar 1 (strain 1330) VirB8 (aquamarine) (PDB ID 2BHM) (45). (D) Sequence alignment of VirB8/TrwG proteins and secondary structure assignment. Sequences were extracted from a larger alignment (see Text S4 in the supplemental material). Sequences are the globular domains depicted in panel C. For each protein, residues involved in the major dimerization site are boxed. Invariant residues are highlighted in yellow. Magenta bars and gray arrows depict the  $\alpha$ -helices and  $\beta$ -strands for the VirB8/TrwG structure, with colored residues in the proteins corresponding to these structural features. Green dots mark the residues mutated within the dimerization interface of *B. birtlesii* TrwG. (E) High-resolution depiction of the major dimerization sites for the proteins illustrated in panels C and D. From left to right: *B. grahamii* VirB8, *B. birtlesii* TrwG, *A. tumefaciens* VirB8, and *B. suis* VirB8. (F) Analysis of *B. birtlesii* TrwG-TrwG interactions using the bacterial two-hybrid system. Different plasmid combinations were transformed into adenylate cyclase deficient and cAMP-specific-phosphodiesterase-deficient *E. coli* strain APE304. After overnight growth in liquid Luria-Bertani medium,  $\beta$ -galactosidase activity was measured and calculated (in Miller units). T25-ZIP and T18-ZIP are positive-interaction control plasmids encoding dimer-forming yeast transcription factor GCN4 (89). Values are from three independent experiments, all analyzed in triplicate. Blue bars depict wild-type-mutant interactions, while yellow bars depict mutant-mutant interactions.

## DISCUSSION

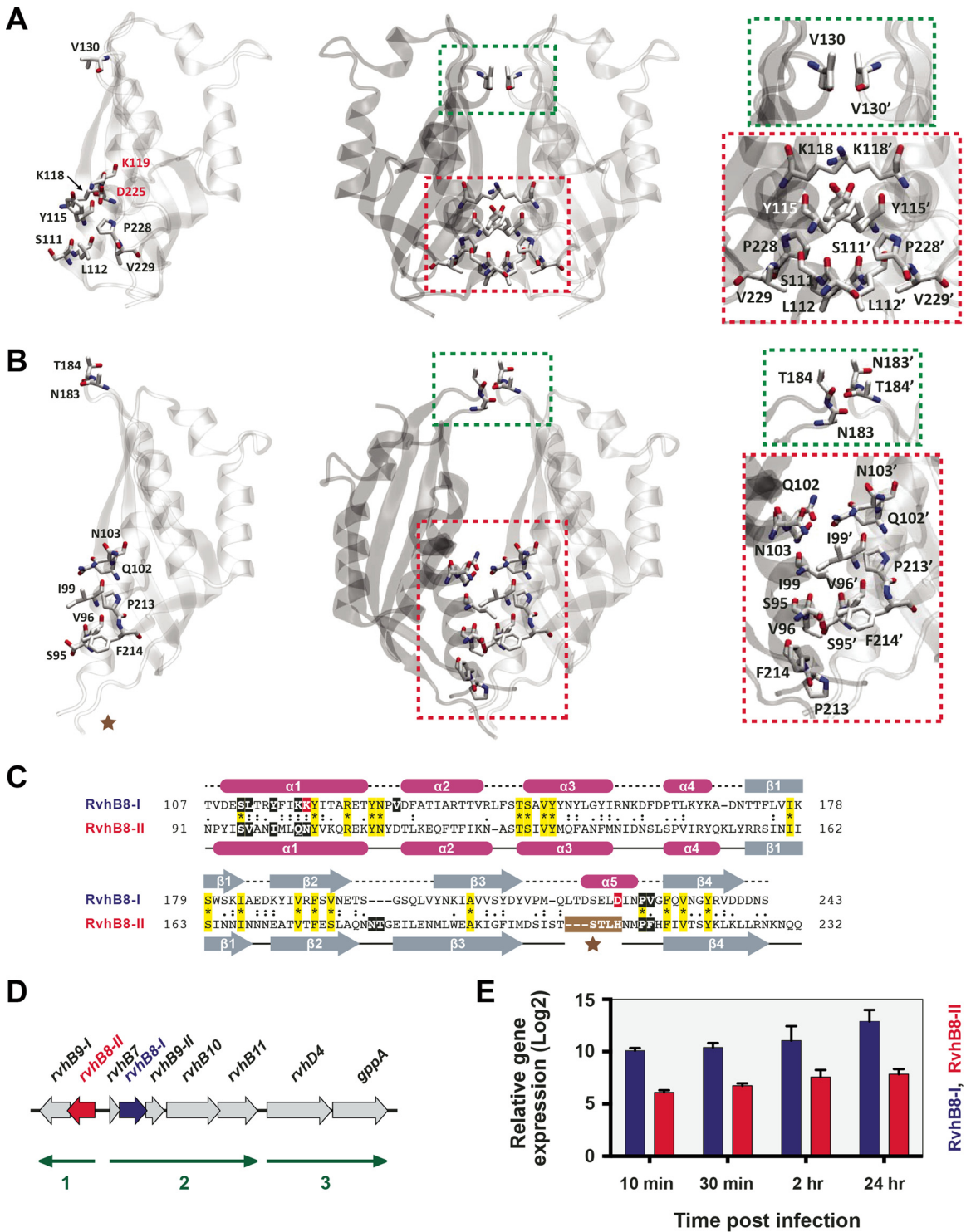
In this study, we set out to identify potential barriers to within-genome T4SS cross-system interchangeability, focusing on the *vir* and *trw* T4SSs carried by *Bartonella* species and the duplication-laden *rvh* T4SS of *Rickettsiales* species. Our data imply that these bacteria either rely on robust spatiotemporal regulation or structural constraints to maintain the cooccurrence of these analogous multiprotein complexes.

For *Bartonella* VirB8 and TrwG proteins, which share overall ~28% identity, the determined structures are highly similar to VirB8 structures previously determined for *A. tumefaciens* (44) and *B. suis* (45) (Fig. 3A to D). With the exception of monomeric *B. quintana* VirB8, all of the *Bartonella* VirB8 and TrwG proteins crystallized as dimers. Using site-directed mutagenesis of residues within the dimer interface, we demonstrate by B2H assays that contacts between helix  $\alpha_1$  of one subunit and the NPXG motif of the other subunit are important for dimerization of *B. birtlesii* TrwG (Fig. 3F). It is probable that this dimer interface area functions similarly in most other VirB8-like proteins of P-T4SSs, given the high conservation of the NPXG motifs. Strikingly, TrwG<sub>wt</sub>-TrwG<sub>V96G/V97G</sub> and TrwG<sub>wt</sub>-TrwG<sub>P214A</sub> protein pairs had a strength of interaction that was indistinguishable from those formed by the wild-type proteins. Also, the interaction signal was not completely abolished when mutations were introduced to both of the interacting proteins. These data indicate either that a certain degree of flexibility is allowed across the *B. birtlesii* TrwG dimer interface or that other areas outside the periplasmic domain of *B. birtlesii* TrwG, such as the TMS region, are important for the dimerization process. Indeed, an M102R mutation in *B. suis* VirB8 (Met102 corresponds to the Val97 of *B. birtlesii* TrwG) (Fig. 3E) abolished the homodimer formation when soluble periplasmic domains were analyzed but had only a partially attenuating effect in an intracellular macrophage survival assay that employed full-length VirB8 (47). The partial M102R mutation-mediated reduction of *B. suis* VirB8 homodimer formation has also been witnessed in B2H assays with full-length proteins (48). Taken together, our data indicate that the periplasmic domains of VirB8 proteins, specifically via the NPXG motif, are important but not the sole determinants of dimerization.

To further explore the VirB8 dimerization process, we used B2H assays to assess the possibility that the structurally conserved *B. grahamii* VirB8 and TrwG form heterodimers (Fig. 4C). These assays indicate that, while weaker than TrwG-TrwG interactions, TrwG-VirB8 interactions are not significantly different in strength from VirB8-VirB8 interactions. A generated structural model of the *B. grahamii* TrwG-VirB8 heterodimer also indicated a similar







**FIG 5** *Rickettsia typhi* expresses two structurally divergent VirB8-like proteins. (A) Structural model for *R. typhi* RvhB8-I (RT0280; YP\_067242). (Left) RvhB8-I monomer in ribbon representation with nine residues involved in the dimerization interface shown in stick representation. For clarity, residues colored red are not shown in the dimer. (Center) RvhB8-I dimer in ribbon representation with seven residues involved in the dimerization interface shown in stick representation. Green indicates a minor dimerization site involving residues within the loop between  $\alpha$ -helices  $\alpha$ 1 and  $\alpha$ 2. Red indicates a major dimerization site involving residues of  $\alpha$ -helix  $\alpha$ 1 and the NPXG motif. (Right) Higher magnification of the RvhB8-I subunit interface. (B) Ribbon representation for the monomer and dimer of the crystal structure (PDB ID 4O3V) of *R. typhi* RvhB8-II (RT0278; YP\_067240). Depiction of dimerization scheme follows the layout shown for RvhB8-I in panel A, except that the minor dimerization site (green) involved residues between  $\beta$ -strands  $\beta$ 2 and  $\beta$ 3. The brown star denotes the break in the structure. (C) Sequence alignment of RvhB8-I and RvhB8-II proteins and secondary structure assignment. Sequences are those of the globular domains depicted in panels A and B. For each protein, residues involved in the dimerization interface are within black (or red) boxes. Invariant residues are highlighted yellow. Magenta cylinders and gray arrows depict the  $\alpha$ -helices and  $\beta$ -strands, respectively, for structures shown in panels A and B. For RvhB8-II, the brown star

(Continued)

dimerization interface relative to TrwG and VirB8 homodimers (Fig. 4A). Our data suggest that *Bartonella* VirB8 and TrwG proteins have the potential to interfere with one another during assembly into their respective T4SSs. It is therefore remarkable that *vir* and *trw* T4SSs are under robust spatiotemporal regulation (40). The stringent response factors SpoT and DksA were shown to mediate regulation of the *vir* T4SS during early infection, with expression of *vir* and *bep* loci requiring the dual activation of the alternative sigma factor RpoH1 and the BatR/BatS two-component system (40). Remarkably, SpoT, DksA, and BatR/BatS were shown to negatively regulate the *trw* T4SS, possibly by moderating the KorA/KorB negative regulator, which is known to control *trw* expression (37). Taken together, spatiotemporal regulation of *vir* and *trw* T4SSs (40) likely prevents cross-system interchangeability, which was shown to be structurally possible at the level of VirB8/TrwG in our current study.

Regarding the *rvh* T4SS, we observed an entirely different relationship between its two VirB8-like proteins, which share ~16% amino acid identity across selected *Rickettsiales* species (Fig. 6A). While the sequence of RvhB8-I can be modeled to the structures of VirB8 and TrwG (Fig. 5A), the crystal structure of RvhB8-II revealed a markedly different organization of the dimerization interface. The region equivalent to the NPXG motif is largely disordered and leads to a different arrangement of contacting residues in the major dimerization site (Fig. 5B). Furthermore, relative to other VirB8 and TrwG structures, a unique minor dimerization site is formed across residues between  $\beta$ -strands  $\beta$ 2 and  $\beta$ 3 in RvhB8-II. This odd structure for RvhB8-II confirms our previous observation that a lack of conservation within the NPXG motif would likely perturb the monomer structure and dimer interface (23, 41). Despite this, the C-terminal domains of both RvhB8-I and RvhB8-II form homodimers in solution (see Fig. S1 in the supplemental material), indicating that each paralog likely forms a functional dimer. This is supported by higher conservation within RvhB8 paralogs (Fig. 6B; see Text S5 in the supplemental material) versus across RvhB8 paralogs (Fig. 6A). A modeled RvhB8-I/RvhB8-II heterodimer (see Text S4 in the supplemental material), combined with a failure to detect interactions between RvhB8-I and RvhB8-II via B2H assays (data not shown) and extraordinary sequence divergence in the N-terminal cytoplasmic/TMS regions across RvhB8 paralogs (see Text S5 in the supplemental material), indicates the unlikely ability of RvhB8-I and RvhB8-II to heterodimerize. Altogether, these data imply that, while highly divergent from one another, RvhB8-I and RvhB8-II proteins are strongly conserved in *Rickettsiales* genomes, illustrating a bizarre architecture of the *rvh* T4SS relative to the canonical *vir* archetypal T4SS.

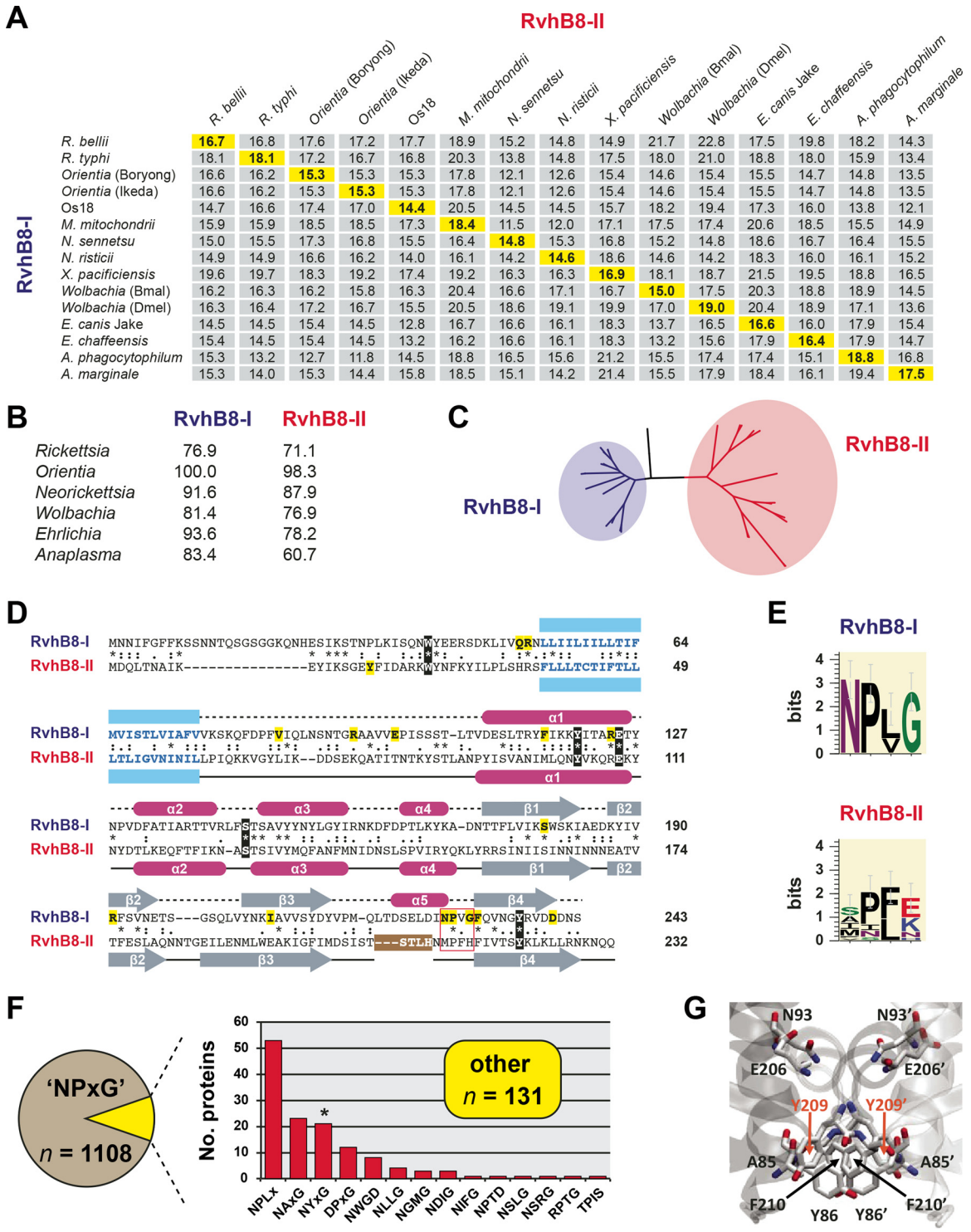
Unlike the spatiotemporal regulation of *Bartonella* *vir* and *trw* T4SSs, we detected simultaneous expression of *rvhB8-I* and *rvhB8-II* genes during *R. typhi* infection of host cells (Fig. 5E). This

suggests that, despite being located in different loci (Fig. 5D), both RvhB8 paralogs are coregulated. While regulators of the *Rickettsia* *rvh* T4SS are currently unknown, studies on other rickettsial genera have identified factors that bind the promoters of multiple *rvh* loci, which are scattered in islets similar to *Rickettsia* genomes (23). In the case of *Ehrlichia chaffeensis*, the EcrX regulator coordinately drives expression of all five *rvh* loci throughout host cell infection, indicating synchronous expression of both RvhB8 paralogs during rickettsial intracellular development (49). For the *Wolbachia* endosymbiont of *Brugia malayi*, *wBmxR1* and *wBmxR2* (homologs of EcrX) were shown to bind upstream of *rvhB9-I* and *rvhB4-II*, respectively, with both factors also binding upstream of a larger *rvh* locus (*ribA-rvhB8-I-rvhB9-II-rvhB10-rvhB11-rvhD4-wsp*) (50). Within the latter locus, it was determined that *rvhB8-I* was expressed in an operon with *ribA* (GTP cyclohydrolase-2), with RvhB8-I detected in *B. malayi* lysates. A regulator of the *rvhB8-II* locus was not determined. Additional studies on various rickettsial species have identified gene and protein expression for both RvhB8 paralogs (41), indicating conserved roles for both proteins in *rvh* T4SS assembly and function.

Despite coexpression, whether RvhB8-I and RvhB8-II both assemble into the same *rvh* secretion machine remains unknown. As it is probable that the *rvh* genes in different rickettsial species are uniquely regulated to orchestrate their specific lifestyles and adapt to their specific intracellular niches, it is tempting to speculate that *rvh* gene duplication underlies stage- and/or host-specific assembly of different *rvh* components. However, two factors do not favor this hypothesis. First, like RvhB8-II, other Rvh-II proteins (RvhB4-II and RvhB9-II) contain atypical characteristics that should prevent their contribution to functional secretion. RvhB4-II proteins lack conserved ATPase active sites that are known to be critical for VirB4 function (51–53), casting doubt on their ability to provide energy for secretion. The C-terminal domains of RvhB9-II proteins either are absent (*Rickettsiaceae*) or lack conserved residues relative to VirB9 proteins (*Anaplasmataceae* and “*Candidatus* Midichloriaceae”). As the VirB9 C-terminal domain interacts with VirB7 and the C-terminal domain of VirB10 in the dodecameric core complex (16, 17, 54) and also participates in transient interactions with other components (VirB1 and VirB8 to VirB11) (10, 55, 56), it is unlikely that RvhB9-II proteins contribute to core complex formation. Second, the *rvh-II* loci are not linked in a manner that indicates their efficient and independent regulation relative to the other *rvh* loci. *rvhB8-I* and *rvhB9-II* loci are adjacent within a large *rvh* islet that includes *rvhB10*, *rvhB11*, and *rvhD4*, while *rvhB8-II* and *rvhB9-I* are found in one distinct operon (*Rickettsia* and *Neorickettsia* species) or present as separate transcriptional units (all other derived *Rickettsiales*) (23). Furthermore, the *rvhB4-II* locus is well separated from all of these loci. Collectively, these characteristics of *rvh* gene duplication make it difficult to envision a scenario where

#### Figure Legend Continued

denotes the break in the structure, with missing residues colored white. (D) *rvhB8-I* and *rvhB8-II* are arrayed in tandem operons within the *R. typhi* genome. The schema shows nucleotide coordinates 351539 to 360917 from the *R. typhi* strain Wilmington genome (NC\_006142) (46). Nine genes are encoded within three predicted operons: 1, *rvhB9-I* and *rvhB8-II* (red); 2, *rvhB7*, *rvhB8-I* (blue), *rvhB9-II*, *rvhB10*, and *rvhB11*; 3, *rvhD4* and *gppA* (guanosine-5'-triphosphate, 3'-diphosphate pyrophosphatase). The direction of transcription for each operon is shown with green arrows. Operons were predicted with fgenesb (90). Other *rvh* genes are encoded in separate clusters within the *R. typhi* genome (41). (E) Expression of the *R. typhi* *rvhB8-I* and *rvhB8-II* genes during early host cell infection. RNA was extracted from HeLa cells infected with *R. typhi*, and gene expression of *RT0280* (encoding RvhB8-I) and *RT0278* (encoding RvhB8-II) was measured by reverse transcription-quantitative PCR (RT-qPCR). Gene expression was normalized to *R. typhi* reference genes *adr1* and *sca5* ( $2^{\Delta CT}$ ). Infections were repeated in triplicate with technical duplicate readings for RT-qPCR. Values are means and standard errors of the means.



**FIG 6** Conserved RvhB8-I and RvhB8-II paralogs are highly divergent from one another. (A) Pairwise divergence between RvhB8-I and RvhB8-II proteins from select rickettsial species. Numbers are amino acid identity (percent) as calculated across a global RvhB8 alignment (see the text for details). Highlighted values on the diagonal depict divergences between paralogs encoded within the same genome. Full species names and NCBI GenBank accession numbers for all proteins are provided in Text S1 in the supplemental material. (B) Across species and strains within the same genus, RvhB8-I is more conserved than RvhB8-II. Numbers are amino acid identity (percent) as described for panel A. Complete percent identity matrices used to estimate protein divergence are provided in Text S5 in the supplemental material. (C) Phylogeny estimation of RvhB8 proteins reveals higher divergence within the RvhB8-II clade than the RvhB8-I clade. ML-based phylogeny was estimated with RAxML on the unmasked global RvhB8 alignment (WAG + gamma + I). A complete tree, as well as phylogenies estimated from the masked alignment with other substitution models, is provided in Text S5 supplemental material. (D) Comparison of *R. typhi* RvhB8-I and RvhB8-II proteins. Sequences were aligned according to structure in SPDBV using “magic fit” followed by “improved fit” algorithms. The alignment shows predicted (top) and solved (bottom) structures for RvhB8-I and RvhB8-II, respectively. Predicted transmembrane-spanning regions (76) are colored blue. Five residues conserved

(Continued)

Rvh-II proteins assemble into a functional T4SS independent of Rvh-I proteins.

With their strict conservation across the derived rickettsial families and strikingly higher rate of evolution than Rvh-I proteins (Fig. 2), it is likely that the functions of Rvh-II proteins are linked. It is also probable that these proteins assemble in some fashion into the *rvh* T4SS. Despite defective ATPase active sites, RvhB4-II proteins are similar in size to RvhB4-I and other VirB4 proteins. Curiously, the N-terminal domains of RvhB9-II proteins are conserved, indicating that they may make contacts with RvhB4 proteins given that the N-terminal domain of VirB9 interacts with VirB4 in the periplasm near the inner membrane (18). For RvhB8-II proteins, dimers may occupy half of the inner membrane channel (IMC) in conjunction with RvhB8-I dimers, perhaps with an as-yet-unknown interaction across these complexes. In accordance with other P-T4SSs, it can be expected that 12 RvhB8 proteins occupy the IMC, as revealed in the electron-microscopic structure of the R388 P-T4SS VirB3-VirB10 complex (19). However, recent structures for VirB8-like proteins from two conjugative T4SSs of Gram-positive bacterial species, TcpC of *Clostridium perfringens* (57) and TraM of *Enterococcus faecalis* (58), revealed trimers as functional units, indicating that the manner by which VirB8 proteins oligomerize in the IMC differs across divergent T4SSs. Furthermore, for I-T4SSs, recent crystal structures of DotI (*L. pneumophila*) and TraM (IncI plasmid R64) revealed octomers and hexamers, respectively, with DotI also oligomerizing with its truncated paralog DotJ (59). Curiously, like RvhB8-II, the structures of DotI, TraM (R64), TcpC, and TraM (*E. faecalis*) all lack helix  $\alpha 5$ , which is highly conserved in RvhB8-I, VirB8, and TrwG structures. Thus, it can be anticipated that RvhB8-II might adopt a different oligomerization scheme relative to RvhB8-I within the *rvh* IMC. Elucidation of the composition of the IMC within the *rvh* T4SS will help resolve the structural and functional significance of two divergent RvhB8 proteins.

**Conclusion.** Our work here presents the first structural investigation on factors regulating the maintenance of multiple divergent T4SSs (or duplicate T4SS components) within a single bacterium. We identified the ability of VirB8 and TrwG of functionally divergent *Bartonella* T4SSs (*vir* and *trw*) to interact and therefore potentially compete in T4SS assembly. This corroborates previous studies demonstrating tight differential expression of these T4SSs (40). For the *Rickettsia rvh* T4SS, the simultaneously expressed RvhB8 proteins were shown to be structurally divergent, casting doubt on their ability to form heterodimers. Taken together, our data indicate that two distinct mechanisms (spatiotemporal for *Bartonella trw* and *vir* T4SSs, structural for *rvh* T4SSs) prevent cross-system interchangeability between divergent T4SSs encoded within a single bacterium.

T4SSs are ideal drug targets provided that no analogous counterparts are known from eukaryotes. Compounds inhibiting the transfer of broad-host-range plasmids (60) and the ATPase activ-

ity of *H. pylori* VirB11 (61) prove the efficacy of such therapeutic approaches. As an assembly factor that interacts with half of the T4SS scaffold components, VirB8 has been suggested as an appropriate target for drugs inhibiting its many interactions (62). Indeed, several inhibitors of *Brucella abortus* VirB8 have been identified, with one compound significantly limiting bacterial growth *in vivo* (63, 64). Our work in this study expands the modality for drug targeting of T4SSs, applicable to bacterial pathogens that harbor multiple T4SSs. Specifically, drugs targeting the barriers to cross-system interchangeability (i.e., regulators) could dysregulate the structural and functional independence of discrete systems, creating interference that prevents their efficient coordination throughout bacterial infection.

## MATERIALS AND METHODS

**Phylogeny estimation of P-T4SSs.** For 75 P-T4SSs encoded within diverse proteobacterial genomes, five proteins (VirB4 and VirB8 to VirB11) were selected for analysis. Additionally, second copies of RvhB4, RvhB8, and RvhB9 encoded within *Rickettsiales* genomes were treated as a minimal T4SS (*rvh-II*), resulting in 90 P-T4SSs in total (for all NCBI GenBank protein accession numbers, see Text S1 in the supplemental material). Individual proteins were separately aligned with MUSCLE (default parameters) (65) and subsequently concatenated into a single data set (unmasked data set; 3,752 amino acids [aa]). Protein alignments were also trimmed of less-conserved regions using Gblocks (66), with these masked alignments also concatenated into a single data set (masked data set; 818 characters). For both datasets, phylogenies were estimated under maximum likelihood (ML) using RAXML v.7.2.8 (67), implementing a gamma model of rate heterogeneity and estimation of the proportion of invariable sites. Two separate analyses for each data set employed the WAG or LG amino acid substitution model, resulting in four total ML-based phylogeny estimations. For all ML analyses, branch support was assessed with 1,000 bootstrap pseudoreplications.

We also analyzed the masked data set with the CAT substitution model, a nonparametric method for modeling site-specific features of sequence evolution (68, 69). The CAT model, as implemented in PhyloBayes v3.3 (70), accommodates saturation caused by convergences and reversions (71). The strong base compositional biases of *Rickettsiaceae* genomes (~30% GC) makes the CAT model highly amenable to estimating rickettsial phylogeny (72–74). Two independent Markov chains were run in parallel using PhyloBayes MPI v.1.2e (75) under the CAT-GTR model, with the bipartition frequencies analyzed at various time points using the bpcomp program. For tree building, appropriate burn-in values were determined by plotting the log likelihoods for each chain over sampled generations (time). Analyses were considered complete when the maximum difference in bipartition frequencies between the two chains was less than 0.1. Ultimately, a burn-in value of 1,000, with sampling every 2 trees, was used to build a consensus tree.

**Comparative analyses of VirB8 proteins.** For *Bartonella* and *Rickettsiales* VirB8 family proteins, predicted TMS regions were determined with TMHMM v.2.0 (76). TMS regions were used to delineate all proteins into NT and CT domains. *Bartonella* VirB8, VbhB8, and TrwG proteins were aligned with MUSCLE (default parameters), with percent identity matrices used to estimate the divergence across all sequences. The NT and CT

### Figure Legend Continued

across all RvhB8 proteins are in black, with residues conserved only in RvhB8-I ( $n = 15$ ) or RvhB8-II ( $n = 1$ ) highlighted in yellow (see Text S5 in the supplemental material). The NPXG motif is in a red box (see the description of panel E below). For RvhB8-II, the region of proteolysis (STLH) that occurred during crystallization is in a brown box. (E) Composition of the NPXG motif across 15 RvhB8-I (top) and 15 RvhB8-II (bottom) proteins. Sequence logos were generated using WebLogo v.3.3 (77). (F) Analysis of the conservation of the NPXG motif across 1,239 nonredundant proteobacterial VirB8 proteins (excluding *Rickettsiales*). Proteins lacking the conserved NPXG motif (10.6%) were placed in 14 categories based on their alternative sequences and ranked by their frequency (see Text S4 in the supplemental material for structural modeling of proteins within each category). (G) Example of a canonical interaction across the NPXG motif for *Yersinia pestis* biovar microtus strain 91001 (NP\_995427), which contains the alternative sequence NYFG.

domains were also separately aligned to determine their percent divergence relative to full-length proteins.

*Rickettsiales* RvhB8-I and RvhB8-II sequences were separately aligned, as well as being combined in one global RvhB8 alignment (see Text S5 in the supplemental material), with percent identity matrices used to estimate the divergence across all sequences. Phylogenies were estimated on the global RvhB8 alignment, which included one outgroup VirB8 protein from *Yersinia frederiksenii* (WP\_042562314). Phylogenies of the unmasked (295 aa) and masked (87 aa) alignments were estimated under maximum likelihood (ML) using RAxML, implementing a gamma model of rate heterogeneity and estimation of the proportion of invariable sites. Two separate analyses for each alignment employed the WAG or LG amino acid substitution models, resulting in four total ML-based phylogeny estimations. Branch support was assessed with 1,000 bootstrap pseudoreplications.

A structural alignment of *R. typhi* RvhB8-I and RvhB8-II was constructed, with conserved residues from the *Rickettsiales* RvhB8-I and RvhB8-II alignments superimposed to illustrate the divergent selective constraints operating on each paralog. Sequence logos depicting the conservation of the NPXG motif for RvhB8-I and RvhB8-II proteins were generated using WebLogo v.3.3 (77). To gain insight into the conservation of the NPXG motif across a broader set of VirB8 family proteins, BLASTP searches (using *A. tumefaciens* VirB8 as a query [GenBank no. AHK05288]) were conducted across the NR (all GenBank + RefSeq Nucleotides + EMBL + DDBJ + PDB) database, coupled with a search against the Conserved Domains database (78). Searches were performed with composition-based statistics across four specific databases: (i) *Alphaproteobacteria* (taxid 28211) excluding *Rickettsiales* (taxid 766); (ii) *Gammaproteobacteria* (taxid 1236); (iii) *Betaproteobacteria* (taxid 28216); and (iv) *Deltaproteobacteria* (taxid 28221) + *Epsilonproteobacteria* (taxid 29547). No filter was used. Default matrix parameters (BLOSUM62) and gap costs (existence, 11; extension, 1) were implemented, with an inclusion threshold of 0.005. A maximum of 500 unique VirB8 family proteins per database were retrieved, with all collected sequences aligned using MUSCLE (default parameters) and manually evaluated for composition of the NPXG motif within the VirB8 structure. Sequences lacking the conserved NPXG motif were selected for further analysis (see “Protein modeling” below).

**High-throughput protein expression, purification, crystallization, and structure determination.** PCR, cloning, screening, sequencing, expression screening, scale-up, and purification of proteins were performed as described previously (79, 80). DNA templates for PCR amplification were obtained from Donald Bouyer (University of Texas Medical Branch, Galveston, TX, USA) for *R. typhi* strain Wilmington and from Christoph Dehio (Biozentrum, University of Basel, Basel, Switzerland) for *Bartonella tribocorum* strain CIP 105476, *B. quintana* strain Toulouse, *B. grahamii* strain as4aup, and *B. birtlesii* strain LL-WM9. Crystal trials, diffraction, and structure solution were performed as described previously (81, 82). Further details and all associated data and summary statistics are provided in a description of the gene-to-structure pipeline (see Text S2 in the supplemental material).

**Protein-protein interactions.** For experiments measuring protein-protein interactions, all oligonucleotide primers used for generating constructs, and a description of each plasmid, see Text S6 in the supplemental material.

**(i) Cloning of bacterial two-hybrid plasmid constructs.** (a) **Wild-type genes.** The genes of interest were fused to the 3' end of the T25 fragment (pKT25) and the 3' end of the T18 fragment (pUT18c) of the *Bordetella pertussis* adenylate cyclase. Full-length *trwG* of *B. birtlesii* was PCR amplified with oligonucleotides prAPV-36 and prAPV-37 using chromosomal DNA of strain IBS325 as the template. The PCR fragment was digested with BamHI and KpnI and ligated into BamHI- and KpnI-digested pKT25, as well as pUT18c, to acquire prAPV001 and prAPV002, respectively. Full-length *trwG* of *B. grahamii* was PCR amplified with oligonucleotides prAPV-52 and prAPV-53 using chromosomal DNA of

strain as4aup as the template. The PCR fragment was digested with BamHI and KpnI and ligated into BamHI- and KpnI-digested pKT25, as well as pUT18c, to acquire prAPV003 and prAPV004, respectively. Full-length *virB8* of *B. grahamii* was PCR amplified with oligonucleotides prAPV-54 and prAPV-55 using chromosomal DNA of strain as4aup as the template. The PCR fragment was digested with BamHI and KpnI and ligated into BamHI- and KpnI-digested pKT25, as well as pUT18c, to acquire prAPV005 and prAPV006, respectively.

**(b) Mutant genes.** The pAPV001 plasmid encoding T25-TrwG of *B. birtlesii* was linearized with PCR using mutagenic 5'-phosphorylated oligonucleotide primers prAPV-38 and prAPV-39 with the mutation P214A and prAPV-40 and prAPV-41 with the double mutation V96G/V97G. The PCR products were gel isolated and religated to acquire pAPV-007 and pAPV-008, respectively. After the mutant genes were verified via sequencing, the inserts were shuttle cloned into BamHI- and KpnI-digested pUT18c to acquire pAPV-009 and pAPV-010, respectively.

**(ii) Bacterial two-hybrid experiments.** The plasmids were introduced into the adenylate cyclase-deficient and cyclic AMP (cAMP)-specific-phosphodiesterase-deficient *Escherichia coli* strain APE304 (83) by the polyethylene glycol method. Colonies (5 to 10) from freshly transformed plates were pooled and subcultured overnight in 5 ml of Luria-Bertani medium with 50  $\mu$ g/ml of kanamycin, 200  $\mu$ g/ml of ampicillin, and 100  $\mu$ M isopropyl- $\beta$ -D-thiogalactopyranoside (IPTG) at 37°C under vigorous shaking (250 rpm). The following day, the strains were diluted either 1:5 or 1:10 in phosphate-buffered saline (PBS) for the measurement of  $\beta$ -galactosidase activities. First, 10  $\mu$ l of chloroform and 10  $\mu$ l of 0.1% (wt/vol) sodium dodecyl sulfate (SDS) were added to 1 ml of the diluted bacterial suspensions. Next, 20  $\mu$ l of the vortexed bacterial suspensions were mixed in triplicate with 180  $\mu$ l of  $\beta$ -galactosidase substrate solution (1 mg/ml of 2-nitrophenyl  $\beta$ -D-galactopyranoside [catalogue no. N1127; Sigma-Aldrich] in 60 mM Na<sub>2</sub>HPO<sub>4</sub>, 40 mM NaH<sub>2</sub>PO<sub>4</sub>, 10 mM KCl, 1 mM MgSO<sub>4</sub>, 0.01% [wt/vol] SDS, and 40 mM  $\beta$ -mercaptoethanol). The reaction mixtures were incubated at room temperature for 20 to 40 min. The reactions were stopped by adding 100  $\mu$ l of 1 M Na<sub>2</sub>CO<sub>3</sub>. The end products were measured at 420 nm and 550 nm. Specific  $\beta$ -galactosidase activities (in Miller units) were calculated as follows:  $1,000 \times [A_{420} - (1.75 \times A_{550})/t \times V \times A_{600}]$ , where  $t$  is the reaction mixture incubation time,  $V$  is the volume of bacterial suspension in the reaction mixture (20  $\mu$ l), and  $A_{600}$  equals the optical density of the 1:5- or 1:10-diluted bacterial overnight cultures measured at 600 nm.

**(iii) Glutaraldehyde cross-linking experiments.** Recombinant N-terminally His-tagged proteins (10  $\mu$ g in 70  $\mu$ l of 20 mM HEPES, 300 mM NaCl, 5% glycerol, and 2 mM dithiothreitol [DTT]) were incubated for 20 min at room temperature in the presence of different concentrations of glutaraldehyde (0, 0.005, 0.01, 0.1, 0.5, 1, and 5%). The reactions were quenched by adding 20  $\mu$ l of 1 M Tris-HCl, pH 7.5. Laemmli loading dye (3 $\times$ , 45  $\mu$ l) was added, and the samples were incubated at 95°C for 10 min. Proteins from 5  $\mu$ l of the samples were separated by SDS-PAGE using 10% SDS gels and transferred onto Protran nitrocellulose transfer membranes (Whatman). The membrane was examined for the His epitope using primary mouse monoclonal anti-His antibody (1:1,000; H1029; Sigma-Aldrich) and secondary HRP-conjugated goat anti-mouse IgG (1:5,000; sc-2005; Santa Cruz Biotechnology) with the enhanced chemiluminescence (ECL) system (SuperSignal West Pico chemiluminescent substrate; Thermo Scientific).

**Protein modeling. (i) Building 3D models.** Three-dimensional (3D) structural models were built for *R. typhi* RvhB8-I and 16 VirB8 family proteins containing divergent NPXG motifs (for complete species names and sequence information, see Text S4 in the supplemental material). 3D models were generated applying the fragment assembly approach of homology modeling (84) using SPDBV/DeepView (85) and the Swiss-Model server (85, 86). Each target sequence was modeled to eight templates, including six VirB8 structures (PDB codes 4MEI from *Bartonella tribocorum*, 4LSO from *Bartonella quintana*, 4KZ1 from *Bartonella grahamii*, 2BHM from *Brucella suis* [45], 2CC3 from *Agrobacterium tumefaciens*

[44], and 4O3V from *Rickettsia typhi*) and two TrwG structures (4JF8 from *Bartonella birtlesii* and chain A of 4NHF from *Bartonella grahamii*). All structures were downloaded from the Protein DataBank (87). Template structures were structurally aligned using SPDBV/Deep View. The target sequences were computationally aligned to this set of templates and manually curated with the following criteria (position numbers refer to the X-ray structure 4MEI, VirB8 from *Bartonella tribocorum*; for a full alignment of template structures see Text S4 in the supplemental material). (i) In  $\alpha$ -helix 1, there is a semiconserved motif, e.g., EAIT in 4MEI, 4LSO, and 4KZ1. (ii) In position 5 of  $\alpha$ -helix 2, an aromatic amino acid can often be found. (iii) In position 8 of  $\alpha$ -helix 3, there is often an aromatic residue. In positions 11 and 12 of the same  $\alpha$ -helix, there is a tendency to have aromatic residues. (iv) In  $\alpha$ -helix 4, there is a tendency to have a proline in position 1. (v) In  $\beta$ -strand 1, a moderately conserved motif exists starting at position 4: hydrophobic-X-hydrophobic-X-hydrophobic-polar. In 4MEI, this motif is V151-T152-I153-K154-S155-I156-S157. (vi) In  $\beta$ -strand 2, in many cases there is the triad motif Q-hydrophobic-charged. In 4MEI, it is Q165-V166-R167. (vii) In  $\beta$ -strand 3, alignments were anchored using a moderately conserved motif from positions  $\Omega$ -4 to  $\Omega$ -2: aromatic-polar/charged-aromatic. (viii) In the loop between  $\alpha$ -helix 5 and  $\beta$ -strand 4, a conserved NPXG motif exists. (ix) Insertions and deletions were moved from inside secondary structure elements to surface loops in order to preserve the fold. This strategy followed the principle that fold is conserved over sequence.

Conserved motifs at the N and C termini were used as main alignment anchors (points 1 and 8 above). Since target sequences share low (or even very low) similarity to the sequences of the templates, only selected criteria for target to template alignment were applied. In some cases (*Colwellia psycherythrae*, *Hydrogenophaga* species, *Legionella longbeachae*, *Legionella pneumophila*, *Pseudomonas putida*, *Pseudomonas syringae*, and *Ralstonia solanacearum*), one or several of these alignment “anchors” were missing. This was especially the case for the only moderately conserved motifs in  $\beta$ -strands.

Target to template alignments in SPDBV/DeepView were saved as “projects” and submitted to the Swiss-Model server via the DeepView project mode. Although the overall quality of the models was limited due to low homology, the structures were of sufficient quality for (i) studying structural properties (i.e., neighboring residues, positions of conserved motifs, etc.) and (ii) investigating dimer interfaces.

**(ii) Building dimers. (a) Homomeric models.** In SPDBV/DeepView, the dimeric structure 2CC3 from *Agrobacterium tumefaciens* was used as a template for dimer modeling. Two versions of the same monomeric model were loaded into SPDBV/DeepView as different layers. One of these monomers was fitted to chain A of 2CC3, the other was fitted to chain B. Superposition of the monomers with the respective chains of 2CC3 was optimized using the “Improved Fit” option. The monomers were merged into a new layer to obtain a homodimer.

**(b) Heterodimeric models.** For the *Rickettsia* heterodimeric model, the structure of *R. typhi* RvhB8-II (4O3V) was used as a template. Chain A corresponded already to RvhB8-II, with chain B replaced by the 3D model generated for RvhB8-I. The TrwG/VirB8 heterodimeric structure for *Bartonella grahamii* was generated using chains A and B of the X-ray structure 4NHF, which corresponds to the TrwG domain of *B. grahamii*. Chain B was then replaced by the X-ray structure of *B. grahamii* VirB8 (4KZ1).

The dimer interface was optimized with SCWRL4 (88). Models were subsequently energy minimized by applying 20 steps of the Steepest Descent algorithm, as implemented in SPDBV/DeepView.

**RT-qPCR for RvhB8 proteins.** HeLa cells ( $\sim 1 \times 10^6$  cells) were infected with *R. typhi* at a multiplicity of infection (MOI) of  $\sim 100$  and incubated at 34°C with 5% CO<sub>2</sub> for 10 min, 30 min, 2 h, and 24 h. At each time point, cells were washed with phosphate-buffered saline (pH 7.4) and RNA was extracted using a Quick-RNA miniprep kit (Zymo Research). iScript reverse transcription supermix for RT-qPCR (Bio-Rad) was used to synthesize cDNA from 500 ng of purified RNA. qPCR was performed with VeriQuest SYBR green qPCR master mix (Affymetrix) in

a CFX384 Multicycler (Bio-Rad) (for primer sequences, see Text S6 in the supplemental material). The thermal cycling conditions included 95°C for 3 min followed by 40 cycles of amplification at 95°C for 10 s and 55°C for 60 s. A melting curve analysis was performed to confirm amplification of a single product for each primer pair. A panel of 6 reference genes (see Text S6 in the supplemental material) was tested to determine which genes were stably expressed (geNorm M score  $\leq 0.5$ ; Qbase Plus; Biogazelle) with infection. *rvhB8-I* and *rvhB8-II* gene expression was normalized to the average cycle threshold ( $C_T$ ) of *R. typhi* reference genes *adr1* and *sca5* ( $2^{\Delta C_T}$ ).

## SUPPLEMENTAL MATERIAL

Supplemental material for this article may be found at <http://mbio.asm.org/lookup/suppl/doi:10.1128/mBio.01867-15/-DCSupplemental>.

Text S1, PDF file, 2.1 MB.

Text S2, PDF file, 1.6 MB.

Text S3, PDF file, 0.5 MB.

Text S4, PDF file, 0.8 MB.

Text S5, PDF file, 1.7 MB.

Text S6, PDF file, 0.2 MB.

Figure S1, PDF file, 0.1 MB.

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