



Bacterial flagellar motor PL-ring disassembly subcomplexes are widespread and ancient

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The bacterial flagellum is an amazing nanomachine. Understanding how such complex structures arose is crucial to our understanding of cellular evolution. We and others recently reported that in several Gammaproteobacterial species, a relic subcomplex comprising the decorated P and L rings persists in the outer membrane after flagellum disassembly. Imaging nine additional species with cryo-electron tomography, here, we show that this subcomplex persists after flagellum disassembly in other phyla as well. Bioinformatic analyses fail to show evidence of any recent horizontal transfers of the P- and L-ring genes, suggesting that this subcomplex and its persistence is an ancient and conserved feature of the flagellar motor. We hypothesize that one function of the P and L rings is to seal the outer membrane after motor disassembly.

cryo-electron tomography | flagellar motor | evolution

The bacterial flagellum is one of the most famous macromolecular machines, made up of thousands of protein subunits that self-assemble in a highly synchronized manner into a motor, a flexible hook, and a long extracellular filament that rotates in a propeller-like fashion to move the cell (1). The process of how these different parts assemble has been studied extensively by using different biophysical and biochemical methods (2–7). These studies have resulted in the current “inside-out” model, which starts with the assembly of an inner-membrane-embedded type III secretion system (T3SS) export apparatus, a membrane/supramembrane (MS) ring, a cytoplasmic switch complex (aka C ring), and a periplasmic rod which connects the MS ring to the extracellular hook. The P (peptidoglycan) and L (lipopolysaccharide) rings surround the rod in the periplasm and are thought to act as a bushing during rotation. Finally, the hook is connected by junction proteins to the long filament. While almost all species have this conserved core, different species can have additional cytoplasmic, periplasmic, and extracellular components (8–12). For example, in some species (like *Vibrio* spp.), the P and L rings are decorated by five proteins (MotX, MotY, FlgO, FlgP, and FlgT) (13, 14). In other species, like *Legionella pneumophila* and *Pseudomonas aeruginosa*, the P ring is decorated by a ring formed by MotY (9).

Much less is known about the process of flagellar disassembly, though it is known that *Caulobacter crescentus* loses its flagellum and pili at a specific stage of its life cycle (15). We and others also recently reported that different Gammaproteobacteria species lose their flagella when starving or due to mechanical stress (7, 16–18). Interestingly, in situ imaging using cryo-electron tomography (cryo-ET) showed that this disassembly process leaves an outer-membrane-associated relic subcomplex consisting of the decorated flagellar P and L rings (referred to henceforth as PL-subcomplexes). These PL-subcomplexes plug the hole in the outer membrane that might otherwise be present after the flagellum disassembles. However, it remains unclear whether these PL-subcomplexes only persist in Gammaproteobacteria, or if the phenomenon is more widespread.

Here, using a combination of cryo-ET (19) and subtomogram averaging (20, 21), we show that the PL-subcomplex persists in

nine additional bacterial species, including *Vibrio cholerae*, *Vibrio harveyi*, and *Vibrio fischeri* (Gammaproteobacteria with sheathed flagella); *Hyphomonas neptunium*, *Agrobacterium tumefaciens*, and *C. crescentus* (Alphaproteobacteria); *Hylemonella gracilis* (Betaproteobacterium); *Campylobacter jejuni* (Epsilonproteobacterium); and *Acetonebacterium longum* (Firmicute). Bioinformatics analyses further show that the P- and L-ring genes are ancient and diverged separately in each species (were not recently transferred horizontally). Together, these results suggest that the outer-membrane-sealing role of the PL-subcomplexes is ancient and widely conserved.

Results

To examine the generality of PL-subcomplex persistence and how the presence of a membranous sheath surrounding the flagellum might affect this process, we used cryo-ET to image nine additional bacterial species from five classes (Fig. 1). All previously described PL-subcomplex subtomogram averages have been of species with unsheathed flagella: *Shewanella oneidensis*, *L. pneumophila*, *P. aeruginosa*, *Salmonella enterica*, and *Plesiomonas shigelloides* (7, 16, 17) (SI Appendix, Fig. S1). All of these feature a crater-like structure in the outer membrane (see examples in SI Appendix, Fig. S1), sealed across the bottom by either the P- or L-ring proteins or additional, as-yet-unidentified molecules. This presumably is to avoid an ~20-nm pore in the outer membrane, which might be detrimental to the cell. For this reason, we were

Significance

In order to understand the evolution of complex biological machines like the bacterial flagellar motor, it is crucial to know what each component does and when it arose. Here, we show that a subcomplex of the motor thought to act as a bushing for the spinning motor likely also serves another function—it plugs the hole in the outer membrane left when the flagellum disassembles. Moreover, this component and function is ancient, since it appears in diverse phyla without evidence of recent gene transfer.

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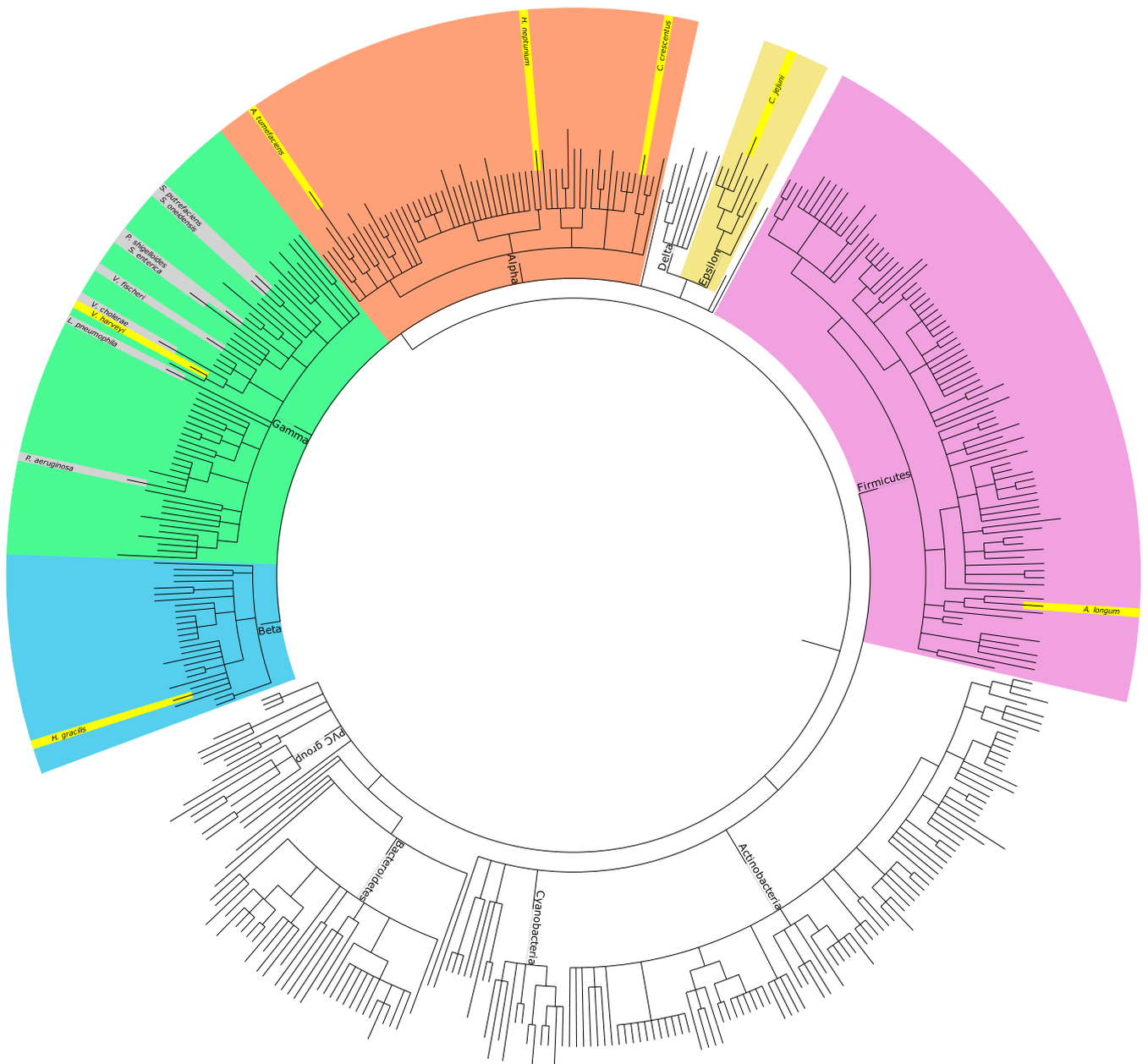


Fig. 1. A taxonomic tree of representative bacterial species. The species where PL-subcomplexes were previously reported are highlighted in gray (all in the Gammaproteobacteria class), while species with PL-subcomplexes identified in this study are highlighted in yellow. PVC, Planctomycetes–Verrucomicrobia–Chlamydiae.

first interested in whether there would be similar discontinuities in the outer membrane in species with sheathed flagella (in which the flagellum does not always penetrate the outer membrane). Images of individual PL-subcomplexes in *V. cholerae* and *V. fischeri* have been published (16), but no subtomogram averages are available. Thus, we first imaged the three Gammaproteobacterial species *V. cholerae*, *V. harveyi*, and *V. fischeri*, whose flagella are sheathed. As expected, we observed that the outer membrane of all three *Vibrio* species bent and extended to sheath the micrometers-long extracellular flagellar filaments (Fig. 2 A–C). At the base of these filaments, flagellar motors were clearly visible. Next to the fully assembled motors, we occasionally observed PL-subcomplexes (Fig. 2 D–F). Subtomogram averages of these subcomplexes confirmed that they indeed consisted of the embellished P and L rings (Fig. 2 G–I). In contrast to the structures previously

observed from unsheathed flagella, the *Vibrio* spp. structures reported here exhibit an intact, convex outer-membrane layer across the top (Fig. 2 G–I). The bottom of the PL-subcomplex is still plugged, however (Fig. 2 G–I, yellow arrows), raising the question of why.

In addition, the structure of the PL-subcomplex in *V. harveyi* has an extracellular ring located just above the outer membrane (Fig. 2I, blue arrows). Such a ring is also present in the fully assembled sheathed flagellum (SI Appendix, Fig. S2, blue arrows). However, while the diameter of this ring is 30 nm in the PL-subcomplex, it has a diameter of 36 nm in the fully assembled flagellum, suggesting that this ring collapses upon flagellar disassembly. The presence of extracellular rings has been described in the unsheathed flagellum of *S. oneidensis* (9) and the sheathed flagellum of *Vibrio alginolyticus* (22). Importantly, the structure

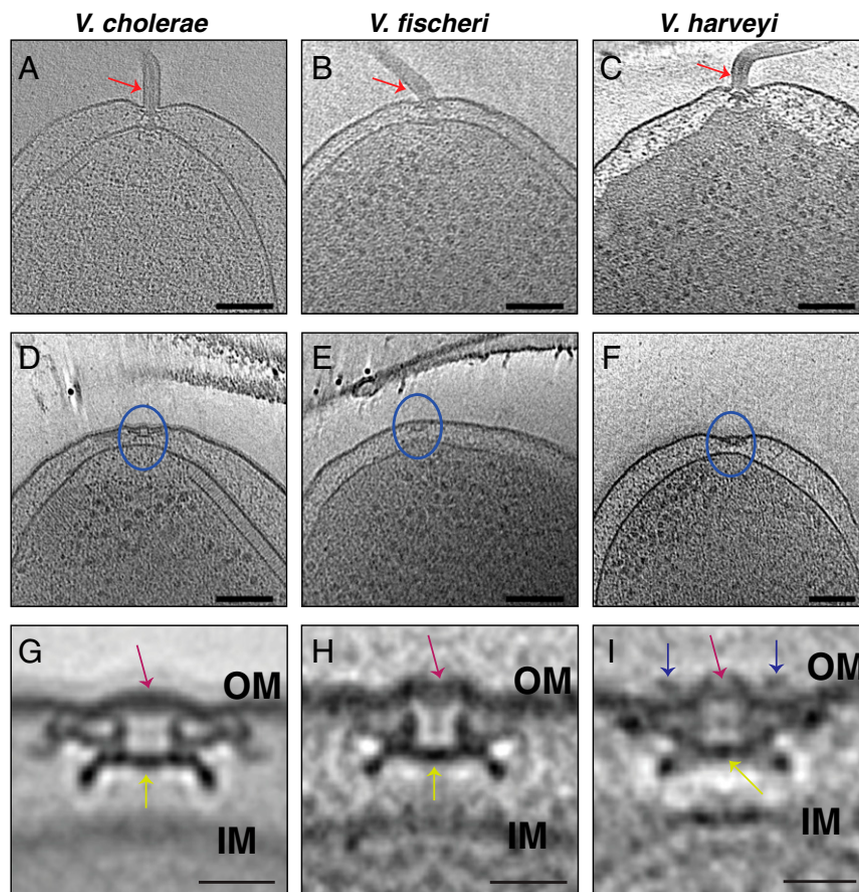


Fig. 2. Cryo-ET of the sheathed Gammaproteobacteria *Vibrio* species. (A–C) Slices through electron cryo-tomograms of *V. cholerae* (A), *V. fischeri* (B), and *V. harveyi* (C), highlighting the presence of a single polar sheathed flagellum in the three species (red arrows). (Scale bars: 100 nm.) (D–F) Slices through electron cryo-tomograms of *V. cholerae* (D), *V. fischeri* (E), and *V. harveyi* (F), highlighting the presence of flagellar disassembly PL-subcomplexes (blue circles). (Scale bars: 100 nm.) (G–I) Central slices through subtomogram averages of PL-subcomplexes in *V. cholerae* (G), *V. fischeri* (H), and *V. harveyi* (I). Purple arrows highlight the presence of intact outer membrane (OM) above the PL-subcomplexes. Yellow arrows indicate the proteinaceous plug inside the P ring. Blue arrows in I highlight the presence of an extracellular ring density in the average of *V. harveyi*. (Scale bars: 20 nm.) IM, inner membrane.

of the PL-subcomplex from *S. oneidensis* has an extra density located just at the membranous discontinuity resulting from disassembling the flagellum (SI Appendix, Fig. S1A). This density in *S. oneidensis* may also be due to the collapse of the extracellular ring present in the fully assembled flagellum.

After this comparison of the PL-subcomplexes in the sheathed and unsheathed flagella of Gammaproteobacteria, we were interested in whether PL-subcomplexes are specific to Gammaproteobacteria or present in other classes in the Proteobacteria phylum. We therefore examined five more species: *H. neptunium*, *A. tumefaciens*, and *C. crescentus* (Alphaproteobacteria [Fig. 3 A–L]); *H. gracilis* (Betaproteobacterium [Fig. 4 A–D]); and *C. jejuni* (Epsilonproteobacterium [Fig. 4 E and F]). PL-subcomplexes were observed in all of these species with the characteristic discontinuity in the outer membrane and a clear plugged base similar to their Gammaproteobacterial counterparts (not enough examples of *C. jejuni* PL-subcomplexes were available to unambiguously assign the presence of a plug). In *C. jejuni*, an inner-membrane-associated subcomplex of the flagellar motor (constituting the MS and C rings, the export apparatus, and the proximal rod) was present in the vicinity of the PL-subcomplex in a pattern reminiscent of what has recently been reported in *L. pneumophila* (7) (Movie S1 and SI Appendix, Fig. S3).

Having established that PL-subcomplexes are widespread in Proteobacteria, we next looked for them in *A. longum*, a deriderm

belonging to the class of Clostridia in the Firmicutes phylum. PL-subcomplexes were found in *A. longum* as well (Fig. 4 G and H).

The presence of PL-subcomplexes in diverse bacterial phyla could be because it is an ancient and conserved feature or because the P- and L-ring proteins were recently horizontally transferred. To explore these possibilities, we performed an implicit phylogenetic analysis on all species in which PL-subcomplexes have been found (by cryo-ET: 15 in total, including the species described here, plus those in refs. 7, 16, and 17). We compared the sequence distances among FlgI (P ring) and among FlgH (L ring) orthologs, as well as 25 single-copy well-conserved proteins (as described in ref. 23; SI Appendix, Table S1). This allowed us to investigate how P- and L-ring proteins evolved compared to the reference 25 proteins (24). If the sequence distances among FlgI (or FlgH) proteins in two species is smaller than the 25 reference proteins, this indicates a horizontal gene-transfer event (24). This analysis of pairwise comparisons of the investigated species showed that the sequence distances between FlgH proteins was at least as divergent as the 25 reference proteins, and, therefore, there is no evidence of horizontal gene transfer between these species (Fig. 5A and SI Appendix, Table S2). This same result was seen for FlgI (Fig. 5B and SI Appendix, Table S3). For the minimum and average protein

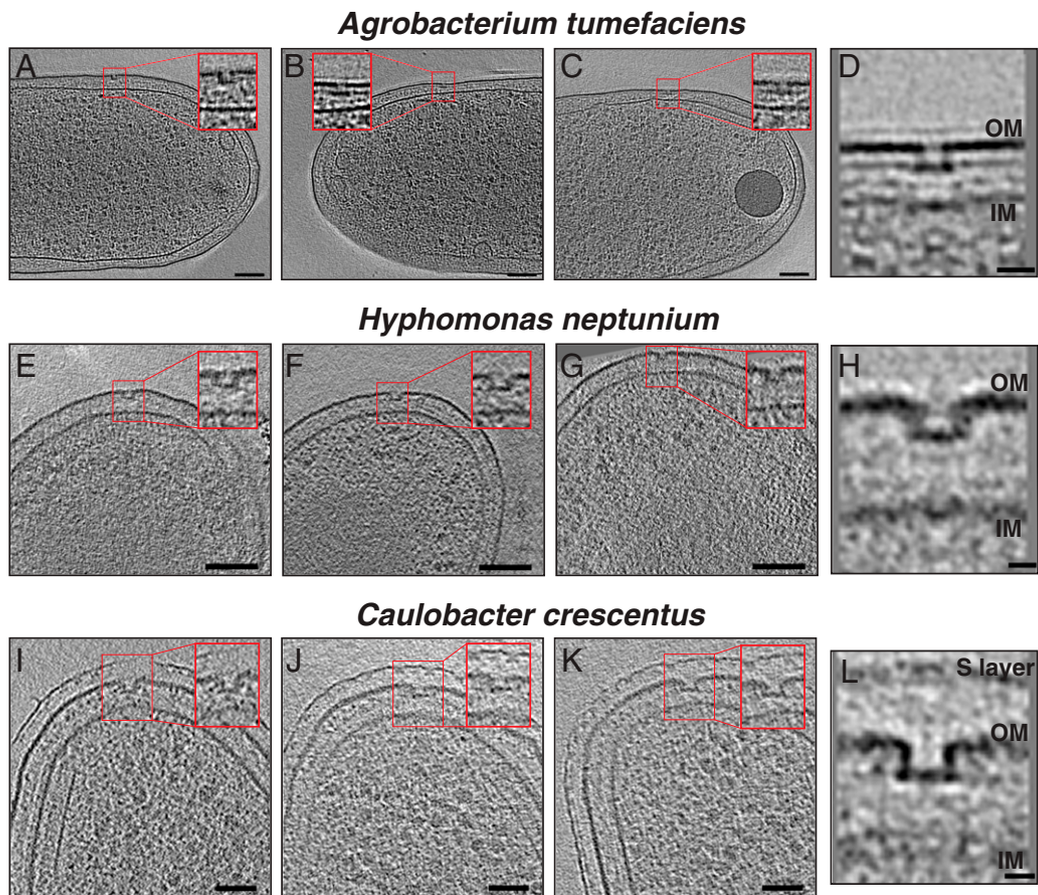


Fig. 3. Cryo-ET of the Alphaproteobacteria species. (A–C) Slices through electron cryo-tomograms of *A. tumefaciens* highlighting the presence of flagellar disassembly PL-subcomplexes with zoom-ins (*Insets*) of these subcomplexes present in the red squares. (Scale bars: 100 nm.) (D) Central slice through a subtomogram average of PL-subcomplexes in *A. tumefaciens*. (Scale bar: 20 nm.) (E–G) Same as in A–C, but for *H. neptunium*. (Scale bars: 100 nm.) (H) Central slice through a subtomogram average of PL-subcomplexes in *H. neptunium*. (Scale bar: 10 nm.) (I–K) Same as in A–C, but for *C. crescentus*. (Scale bars: 50 nm.) (L) Central slice through a subtomogram average of PL-subcomplexes in *C. crescentus*. (Scale bar: 10 nm.) IM, inner membrane; OM, outer membrane.

distances among the 15 species in this study, see [SI Appendix, Table S4](#).

In *Shewanella putrefaciens* and *P. shigelloides*, two copies for FlgI and FlgH were annotated. For both species and both genes, one copy showed more similarity to the nearest relative (*S. putrefaciens* FlgI, A4Y8M8, and FlgH, A4Y8M9; *P. shigelloides* FlgI, R8AUG5, and FlgH, R8AUH3, referred to as the primary copy). On the other hand, the other copy (referred to as the secondary copy) showed more divergence from any studied organism (*S. putrefaciens* FlgI, A4YB38, and FlgH, A4YB39; *P. shigelloides* FlgI, R8AS48, and FlgH, R8AS34; [SI Appendix, Figs. S4 and S5 and Tables S5 and S6](#)). GC-content analysis provided no evidence that any copy of FlgI and FlgH in either species is a result of horizontal gene transfer ([SI Appendix, Figs. S6 and S7 and Materials and Methods](#)).

Discussion

An important step in reconstructing the evolutionary history of biomolecular complexes is to know when certain features and functions originated. Recent studies indicate that the bacterial flagellum, one of the prime motility nanomachines in the prokaryotic world, is an ancient machine that originated from a single or few proteins through multiple gene-duplication and diversification events that preceded the common ancestor of bacteria (23). Some parts of the flagellar system are homologous to other subcomplexes present in other machines. The stator proteins MotA/B are homologous to proteins in the Tol-Pal and

TonB systems, while the motor's ATPase is homologous to the beta subunit of the adenosine triphosphate synthase (23, 25). This suggests that other, even older machines donated features and functions to the first motor. Moreover, the T3SS, also known as the injectisome, is homologous to the bacterial flagellum (though the P and L rings of the motor are not homologous to the secretin part of the injectisome) (26). Because motility preceded the evolution of eukaryotic cells, the targets of T3SS, and the T3SS is restricted mainly to proteobacteria, the injectisome likely derived from the flagellum (27, 28). On the other hand, motility in the archaeal domain of life is driven by another nanomachine—namely, the archaeellum, which is structurally related to the type-IV pilus system and not to the bacterial flagellum, suggesting different evolutionary histories for these motility nanomachines (29–31).

The proteins that form the P and L rings—namely, FlgI and FlgH, respectively—are present widely in flagellated bacteria; however, they are not as universal as other flagellar proteins known as the core proteins. For example, Spirochaetes (characterized by periplasmic flagella) and Firmicutes (many of its members are characterized by a single membrane) do not necessarily have the P and L rings. These two phyla are usually considered among the earliest evolved phyla of bacteria (32), suggesting that, although the P and L rings appeared early during the flagellar evolution, they were probably not present at first (23). However, recent studies prompted a proposal that the diderm cell plan may represent a permanent stall in one phase of

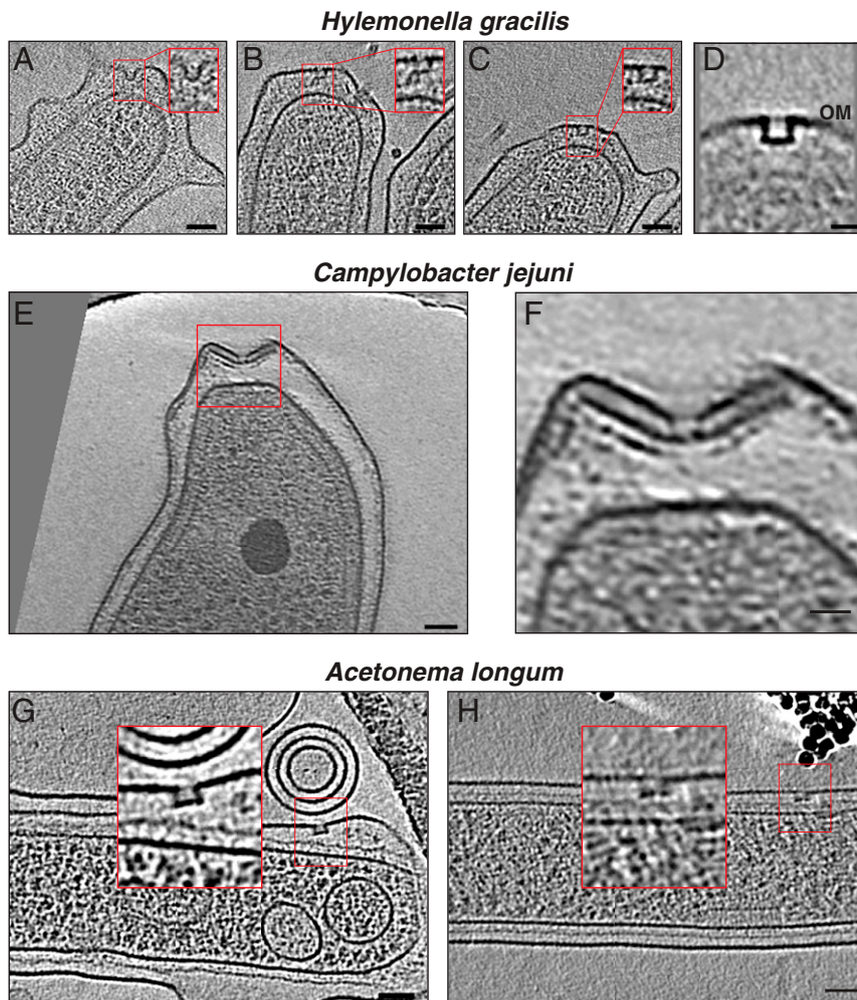


Fig. 4. Cryo-ET of Betaproteobacteria, Epsilonproteobacteria, and Firmicutes. (A–C) Slices through electron cryo-tomograms of *H. gracilis* highlighting the presence of flagellar disassembly PL-subcomplexes with zoom-ins (*Insets*) of these subcomplexes present in the red squares. (Scale bars: 50 nm.) (D) Central slice through a subtomogram average of PL-subcomplexes in *H. gracilis*. (Scale bar: 20 nm.) (E) A slice through electron cryo-tomogram of *C. jejuni* highlighting the presence of a flagellar disassembly PL-subcomplex (red square). (Scale bar: 50 nm.) (F) A zoom-in of the area enclosed in the red square in E. (Scale bar: 20 nm.) (G and H) Slices through electron cryo-tomograms of *A. longum* highlighting the presence of flagellar disassembly PL-subcomplexes with zoom-ins (*Insets*) of these subcomplexes present in the red squares. (Scale bars: 50 nm.)

sporulation and that the last common ancestor of all existing bacteria was a diderm sporulator. This model means that modern monoderms descended from a diderm that lost its outer membrane (33, 34). This hypothesis is not unreasonable in light of cataclysmic conditions of early Earth, in which only a spore might have been able to persist through some evolutionary bottleneck. All this makes the origin of PL rings very speculative. Perhaps the flagellum did evolve before diderms, implying that PL rings evolved later than other flagellar components. Alternatively, the flagellum might not have evolved until there were already diderms around, suggesting that PL rings might be as equally ancient as other flagellar parts.

While other periplasmic and extracellular components of the flagellum (the proximal and distal rods, the hook, the hook–filament junction proteins, and the filament proteins) are exported by the flagellar T3SS export apparatus, the P- and L-ring proteins are secreted through the Sec pathway (35). Also, previous studies suggested that the secreted FlgI and FlgH proteins can exist in a stable form in the periplasm before their nucleation on the rod, which could be either due to high intrinsic stability of these two proteins or due to the low protease activity in the periplasm (2, 36). This might also explain the persistence of plugged PL-

subcomplexes after flagellar disassembly. Alternatively, although the P and L rings have been thought to act as bushings supporting the rotation of the rod, the discovery that they persist in an altered, sealed form after the disassembly process could suggest an additional function—perhaps they remain to seal what would otherwise be a hole in the outer membrane. Here, we have found that PL-subcomplexes are widespread among bacteria and ancient (not the result of recent horizontal gene transfers). This indicates that the putative outer-membrane-sealing function is important enough to have been conserved since the diversification of bacterial phyla.

In addition, we showed that, in species with sheathed flagella, the outer membrane remained intact above PL-subcomplexes, but the base of the PL-subcomplexes was, nevertheless, apparently sealed. This raises questions about the nature and function of the PL-subcomplex in these species. Does it serve a function distinct from membrane-sealing in *Vibrio*, or it could be a vestige retained in their evolution from ancestors with unsheathed flagella? Finally, it will be interesting to find out whether membrane seals are needed only for flagellum disassembly, or if they might be needed in other closely related systems like the injectisome.

Data Availability. Some of the data used in this study are available in the Electron Tomography Database (ETDB)-Caltech (48). All of the data are available upon request from the corresponding author.

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