Molecular Architecture of the Bacterial Flagellar Motor in Cells

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ABSTRACT: The flagellum is one of the most sophisticated self-assembling molecular machines in bacteria. Powered by the proton-motive force, the flagellum rapidly rotates in either a clockwise or counterclockwise direction, which ultimately controls bacterial motility and behavior. Escherichia coli and Salmonella enterica have served as important model systems for extensive genetic, biochemical, and structural analysis of the flagellum, providing unparalleled insights into its structure, function, and gene regulation. Despite these advances, our understanding of flagellar assembly and rotational mechanisms remains incomplete, in part because of the limited structural information available regarding the intact rotor–stator complex and secretion apparatus. Cryo-electron tomography (cryo-ET) has become a valuable imaging technique capable of visualizing the intact flagellar motor in cells at molecular resolution. Because the resolution that can be achieved by cryo-ET with large bacteria (such as E. coli and S. enterica) is limited, analysis of small-diameter bacteria (including Borrelia burgdorferi and Campylobacter jejuni) can provide additional insights into the in situ structure of the flagellar motor and other cellular components. This review is focused on the application of cryo-ET, in combination with genetic and biophysical approaches, to the study of flagellar structures and its potential for improving the understanding of rotor–stator interactions, the rotational switching mechanism, and the secretion and assembly of flagellar components.

BACTERIAL MOTILITY AND FLAGELLA

Many bacteria require motility for their growth and survival. Motility is also essential for the infectivity of many prokaryotic pathogens. Although other types of motility exist (e.g., gliding motility), flagellum-mediated translational motion is the most common mechanism in bacteria. Flagellar rotation is driven by the proton- or sodium-motive force across the cytoplasmic membrane. In most externally flagellated bacteria, counterclockwise rotation (CCW, as viewed from the distal end of the organisms), relapsing fever (several Borrelia species), syphilis (Treponema pallidum), leptospirosis (Leptospira species), and swine dysentery (Brachyspira species). Lyme disease is the most common vector-borne infection in the United States. Syphilis is a prevalent sexually transmitted disease in many areas of the world, while leptospirosis is the most common worldwide waterborne zoonosis.

B. burgdorferi is one of the best studied spirochetes in terms of motility. In contrast to the external flagellar filaments found in most motile bacteria, spirochetes possess periplasmic flagella (PFs) that are enclosed between the outer membrane and the peptidoglycan layer within the periplasmic space (Figure 1). In B. burgdorferi, 7–11 PFs are inserted subterminally at both cell poles (Figure 1B). The PF bundles wind around a flexible protoplasmic cylinder and overlap in the middle of the cell. The PFs are essential for the distinct morphology, motility, and infectious life cycle of B. burgdorferi. The flagella at each end of the cell are coordinated to rotate in opposite directions during translational motion and in the same direction (i.e., either CW or CCW) during the spirochete equivalent of tumbling, called “flexing.” Rotation of the flagella causes a serpentine movement of the entire cell body, allowing the organism to efficiently bore its way through viscous media or tissue.

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The structure and function of bacterial flagella have been extensively studied in model systems Salmonella enterica and Escherichia coli, as summarized in several recent comprehensive reviews.4,11−14 Of approximately 50 genes involved in the expression and assembly of the flagellum, ∼20 produce integral flagellar components. The flagellum consists of the motor, the hook, and the helically shaped flagellar filament (Figure 1A,B).

The flagellar motor can be divided into several morphological domains (Figure 1C,D): the MS ring (FliF, the base for the flagellar motor), the C ring (FliG, FliM, and FliN, the switch complex regulating motor rotation), the export apparatus (a large complex exporting flagellar substrates), the rod (connecting the MS ring and the hook), the L and P rings on the rod (thought to serve as bushings at the outer membrane and at the peptidoglycan layer, respectively), and the stator, which is the motor force generator embedded in the cytoplasmic membrane.

The stator complex is composed of two transmembrane proteins, MotA and MotB, in E. coli and S. enterica. MotA has a large cytoplasmic domain, which contains conserved charged residues that are critical for the interaction with the rotor-associated protein FliG.15,16 MotB has a large periplasmic domain that is believed to bind to the peptidoglycan layer.17,18 The arrangements of MotA and MotB within the complex have been studied extensively by mutational analysis and systematic disulfide cross-linking studies.19,20 Four MotA subunits and two MotB subunits form an ion-conducting complex (MotA-MotB) that couples the proton flux to rotation of the rotor–flagellar filament assembly.21 A conserved aspartic acid residue in the transmembrane segment of MotB (Asp32 in E. coli) is the predicted proton-binding site.22 The proton binding or dissociation at this residue triggers conformational changes of the cytoplasmic domain of MotA in the stator, which are believed to drive the flagellar rotation through interactions between MotA and FliG.23

The flagellar export apparatus is responsible for the secretion of flagellar type III protein substrates, which include the polypeptide subunits of the flagellar rod, the hook, and the filament. The export apparatus uses both the ion-motive force24,25 and the energy of ATP hydrolysis26−28 to complete the export process. It is structurally and functionally homologous to the pathogenic type III secretion system (T3SS) that directly injects virulence factors into host cells.29 The export apparatus is located at the bottom of the MS ring (Figure 1C,D) and is composed of six membrane proteins (FlhA, FlhB, FliP, FliQ, FliR, and FliO) and three soluble proteins (FliI, FliH, and FliJ) in the cytoplasm.30 The membrane components are thought to form an export gate for secretion of the substrates, while the three soluble proteins form a FliH/FliI/FliJ complex that promotes the export process by binding and delivering export substrates to the export gate. It has been suggested that the FliH/FliI/FliJ complex has an architecture similar to that of F- and V-type ATPases.31,32

The structures of many flagellar proteins have been determined at atomic resolution, including components of the flagellar filament (FliC33), the C ring (FliM34, FliG35−37 FliN,38 and the FliG/FliM complex39,40), and the stator (MotB41,42). Cryo-electron microscopy (cryo-EM) studies

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**Figure 1.** Schematic models of the external flagellum of E. coli (A and C) and periplasmic flagella of B. burgdorferi (B and D). Periplasmic flagella are distinct from the external flagella, as they are enclosed within the outer membrane and their flagellar motors are considerably larger and more complex. However, the core architecture of the two flagellar types is comparable. Shared structures include the MS ring, the C ring, the rod, the hook, the filament, the stator, and the export apparatus.
have provided the most detailed structures of the purified basal body, which contains the MS ring, the C ring, the rod, and the P and L rings.43 However, these structures do not contain components of the stator and the export apparatus. In early electron microscopy studies, the stator and the export apparatus were visualized in freeze-fracture preparations of cytoplasmic membranes.44−46 The first examination of the intact flagellar motor by cryo-electron tomography (cryo-ET) was reported in 2006.47 This technique has been utilized to reveal the structural features of the stator and export apparatus in relation to the rotor elements.48−52 The combination of high-throughput cryo-ET and genetic analysis has been particularly useful for dissecting flagellar motor structure and assembly at 3.5 nm resolution.50,52,53 Therefore, this review is intended to provide an overview of this advanced imaging technique and its promising role in understanding the structure and function of intact flagellar motors.

### 3D VISUALIZATION OF INTACT FLAGELLAR MOTORS BY CRYO- ELECTRON TOMOGRAPHY AND SUBTOMOGRAM AVERAGING

Cryo-ET is a 3D imaging technique that in principle is comparable to computerized axial tomography (CAT) by which a 3D structure is reconstructed from its two-dimensional (2D) projections (Figure 2).54 The unique strength of cryo-ET lies in its potential for visualizing large macromolecular assemblies in their native environment without fixation, dehydration, or staining artifacts. The preparation of frozen hydrated specimens is a critical step in this technique. Suspensions of freshly prepared, viable bacteria are deposited onto EM holey carbon grids and then rapidly frozen in liquid ethane at approximately −180 °C. Frozen hydrated specimens (Figure 2A) are then imaged at −170 °C using a cryo-electron microscope (Figure 2B). A low-dose tilt series of images (Figure 2C), which typically cover an angular range of −64° to +64° in 1.5° increments, is collected and aligned to generate a 3D tomographic reconstruction (Figure 2D).52 The flagellar motors are readily visible in the reconstruction (Figure 2D). However, the resolution and contrast of one subtomogram from one flagellar motor are poor for fully understanding their molecular details (Figure 2E).

Subtomogram averaging and classification are the methods of choice for improving the signal-to-noise ratio and resolution of macromolecular assemblies.55,56 Multiple copies of the flagellar motors are visually identified in the cell tomograms, and the 3D density map of each flagellar motor (Figure 2E) is extracted from its cellular context. Thousands of subtomograms are aligned and averaged to obtain a higher-resolution structure with an improved signal-to-noise ratio (Figure 2F,G). To minimize potential reference bias and also identify conformational heterogeneity, multivariate statistical analysis (MSA) is a key method utilized for the classification of subtomograms.56,57 The resulting higher-resolution structures can be mapped back into their cellular context (Figure 2H), revealing the macromolecular organization at an unprecedented level.

Cryo-ET is a continuously evolving technique.58,59 Many recent developments significantly enhance the resolution and throughput, including automation,60 new methods for image processing,61 phase-plate techniques,62,63 and new generation direct electron detectors.64,65 It is expected that employment of new technologies will greatly enhance the ability to determine higher-resolution structures of intact flagellar motors in a broader range of bacterial species.
To fully understand the density maps derived from cryo-ET and subtomogram averaging, it is critical to fit ("dock") available atomic structures from individual components into 3D maps. Together with other structural and biochemical methods, cryo-ET is able to bridge the information gap from cells to molecules, which is essential for understanding the flagellar motor in its cellular context.

**DIFFERENCES BETWEEN THE PURIFIED BASAL BODY AND INTACT FLAGELLAR MOTOR IN CELLS**

Cryo-EM studies have provided a detailed structure of the purified basal body of *S. enterica*. However, many membrane-associated features are absent (Figure 3A), as demonstrated by comparing basal body reconstruction with the low-resolution map derived from cryo-ET of *S. enterica* cells (Figure 3B). Both membranes and the export apparatus are visible in the intact *S. enterica* motor, whereas little structural detail can be discerned in the stator region. In contrast, intact motor structures from relatively thin spirochetes, as exemplified by *B. burgdorferi* (Figure 3E,F), can be derived at higher resolution and reveal significantly more detail.

The boundary between the rod and the MS ring has been defined by using cryo-ET reconstructions of a rodless *B. burgdorferi ΔfliE* mutant (Figure 3G). In the absence of the rod, a socketlike domain of the MS ring is clearly revealed in the ΔfliE mutant (Figure 3H). As expected, the MS ring from *B. burgdorferi* shares a configuration similar to that in *S. enterica* (Figure 3I). Notably, the central channel of the MS ring is closed in the ΔfliE motor (Figure 3H,I). In contrast, the channel is open in the intact motor, suggesting that the rod proteins form an integrated complex with the MS ring and thereby promote the opening of the channel (Figure 3E,F).

Comparative analysis of the *S. enterica* and *B. burgdorferi* motor structures allows definition of the conserved components in flagellar motor: the MS ring, the rod, the P ring, and the export apparatus. However, the remaining *B. burgdorferi* basal body is structurally comparable to the *S. enterica* basal body (minus the C ring), suggesting that it is composed of the MS ring and the rod.

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and the rod. This is a spirochete-specific structure, and the encoding genes have not been identified at the time of this review. Other peripheral densities that span the membrane are thought to represent the stators (Figure 3F,I, outlined in blue). In spirochetes, both the collar and the putative stators exhibit an evident 16-fold symmetry. Interestingly, the stator is embedded in a curved cytoplasmic membrane and reflects the same curvature; the intrinsic flexibility of the rotor—stator interaction is likely an important factor during flagellar rotation.52

### INTACT MOTOR STRUCTURES IN DIFFERENT BACTERIAL SPECIES

Molecular architectures of intact flagellar motor from several phyla have been determined at 3–8 nm resolution by using cryo-ET and subtomogram averaging.49,51,52,68 To illustrate the degree of conservation, six representative motor structures derived from several bacteria with external or periplasmic flagella are compared in Figure 4. The *E. coli*, *Campylobacter jejuni*, *Hylemonella gracilis*, and *Treponema primitia* motor structures were obtained in a comprehensive study by Chen et al.49 Whereas the overall architectures vary between species,49 the MS ring, the C ring, the rod, and the export apparatus have the same overall configuration as those determined in *S. enterica*51 and *B. burgdorferi*,50,52,68 suggesting that the core components of the flagellar motors and their interactions are conserved (Figure 4). In general, the MS ring forms the base of each motor. The socket and central channel of the MS ring are visible in all species, suggesting that it provides an evolutionarily conserved platform for rod assembly. The C ring is consistently attached to the cytoplasmic edge of the MS ring, but the C ring diameter varies considerably in different species. The export apparatus is located at the bottom of the MS ring. A doughnut-shaped torus is well conserved among different species and is located ~6 nm below the inner membrane. Additional cytoplasmic densities likely correspond to the FlhI/FlhJ/FlhK complex (Figure 4). Notably, the motor structures from *B. burgdorferi*50,52 and *Leptospira interrogans*51 reveal significantly more detail in the export apparatus and other components, which is thought to be due in part to the higher resolution that can be achieved with thin bacterial cells in combination with the use of a large number of individual motor structures for subtomogram averaging.

The rod functions as both the secretion channel and the drive shaft that transmits torque. Five proteins (FlIE, FlgB, FlgC, FlgF, and FlgG) are involved in assembly of the rod.59 When the rod proteins are effectively expressed, the central channel of the MS ring adopts an open conformation that serves as a template for rod assembly.50 The proximal rod proteins assemble on top of the open channel of the MS ring in the following order: FlIE, FlgB, FlgC, and FlgF.50,69–71 The distal rod of external flagella is estimated to consist of four turns containing 26 FlgG subunits (~15 nm in *S. enterica*).70 In contrast, the distal rod (~4 nm) in spirochetes is too short to penetrate the outer membrane (Figure 4E), contributing to the periplasmic localization of the flagella. It is likely that FlgG polymerizes in only one turn in *B. burgdorferi*, which appears to be the case for the other rod proteins (FlgB, FlgC, and FlhO, a FlgF homologue).50 The variable rod length in species with external flagella is likely controlled by the degree of polymerization of the distal rod protein FlgG.

The P and L rings that are located around the rod of Gram-negative organisms are composed of Flgl and FlgH, respectively, and are thought to function as bushings in the cell envelope.4,11 The densities on the external surface of the rod differ considerably in external flagella and periplasmic flagella (Figure 4), consistent with the variable presence of encoding genes flgl and flgh.72 For example, *Treponema* species do not have identifiable flgl and flgh genes (Figure 4D) and lack both P and L rings.47,73 *B. burgdorferi* has flgH but does not have flgL, consistent with the presence of the P ring, but not an L ring. Inactivation of flgL resulted in the loss of the P ring structure.52 *L. interrogans* has both genes and has a large, contiguous density on the outer surface of the rod.51 The exact functions of the rod-associated rings in spirochetes are not known.

The stator complexes (MotA/MotB) assemble around the MS ring in the cytoplasmic membrane. In *S. enterica* and *E. coli*,

![Figure 4](https://example.com/figure4.png)
they are anchored to the peptidoglycan layer through the periplasmic domain of MotB. Strikingly, many bacterial species evolve the periplasmic features (Figure 4B−F, colored cyan), in comparison with the E. coli motor (Figure 4A). Some external flagella contain a disklike structure (Figure 4B,C). In spirochetes, a periplasmic collar assembles around the MS ring, and these structures appear to be quite heterogeneous in the different spirochetal genera (Figure 4D−F). The function(s) and protein composition of the collar structures are currently unknown. Interestingly, these periplasmic features are closely associated with the stator, suggesting that the stator interacts with the collar. In contrast, the motor structure derived from E. coli or S. enterica cells does not contain any apparent periplasmic features other than the rod and P and L rings (Figure 4A). The stator complexes in these organisms are known to have variable occupancy and a dynamic nature in their flagellar motors, which may contribute to the lack of a clearly defined stator structure in cryo-ET images.

Overall, cryo-ET images of the core structures (MS ring, rod, and C ring) of spirochetal flagella are similar to those of other bacteria. However, spirochetes also have a clearly discernible stator structure, a spirochete-specific collar, and variable P and L rings. The small cell diameter (0.1−0.3 μm) and orderly arrangement of flagellar motors near the cell ends in spirochetes also facilitate in situ cryo-ET analysis of these structures at 3.5 nm resolution, indicating the benefits of these cells in cryo-ET analysis. The intact flagellar motor structure of E. coli recently determined by cryo-ET has a relatively low resolution (5.9 nm). It is likely that further improvement of the cryo-ET resolution can be achieved through the use of minicells, high-efficiency electron detectors, and high-throughput techniques. Nevertheless, spirochetes (B. burgdorferi in particular) represent a valuable model system for elucidating key questions in flagellar structure and function.

### SEQUENTIAL ASSEMBLY OF BACTERIAL FLAGELLA

Flagellar assembly is a finely orchestrated biochemical process involving both highly regulated motility gene expression and ordered protein assembly. The morphogenetic pathway for flagellar synthesis has been well-established in S. enterica. Recently, the combination of cryo-ET and genetic analysis in B. burgdorferi has permitted determination of the location of specific flagellar proteins and the visualization of the process of flagellar assembly in cells. As an example, key intermediates in the flagellar assembly of B. burgdorferi can be genetically trapped by systematically targeting individual flagellar genes (encoding the rod, hook, and filament proteins) (Figure 5). Interestingly, the MS ring channel appeared to be closed in a ΔfliE mutant, and no rod-associated density was visualized; thus, assembly of the rod in B. burgdorferi is FliE-dependent, consistent with studies conducted with S. enterica. Cryo-ET analysis of each rod mutant (ΔfliE, ΔflgB, ΔflgC, ΔflhO, and ΔflgG) permitted assessment of the contribution of each rod protein to rod assembly. Similarly, analysis of hook mutant ΔflgE revealed a structure thought to represent a hook cap attached to the end of distal rod. Examination of a flaB filament deletion mutant also exhibited a filament cap structure, which is likely related to the cap protein FlhD. In this study, high-throughput cryo-ET procedures permitted the comparative analysis of seven flagellar mutants and more than 20000 gigabytes of data and thereby provide a large set of 3D flagellar structures, which may represent key intermediates during flagellar assembly (Figure 5).

### ASYMMETRIC RECONSTRUCTION REVEALS NOVEL ARCHITECTURAL ELEMENTS OF THE EXPORT APPARATUS

The flagellar motor is composed of many different components, and each component has its own symmetries. In spirochetes, the periplasmic collar has 16-fold symmetry, while the export apparatus is expected to possess different symmetries. To better understand the structure of the export apparatus, we chose the L. interrogans motor as an example for further image analysis, because of its striking structural details as shown in Figure 4F. A novel procedure in which specific substructures of interest are classified and aligned without applying rotational symmetry was utilized recently to delineate asymmetric structural assemblies in bacteriophage T7. This approach, which avoids introducing potential artificial periodicities and thus obscuring "true" symmetries, was used to determine the...
structure of the *Leptospira* motor in greater detail (Figure 6). Specifically, classification focusing on the export apparatus revealed significant details in its overall structure and interaction with the C ring (Figure 6A,B). The cytoplasmic portion of the export apparatus complex shows evident features in 6-fold symmetry or 12-fold symmetry, while the periplasmic collar and stator complexes maintain 16-fold symmetry (Figure 6A,B).

According to the structural comparison among different species of flagellar motors, the motor structure of *L. interrogans* can be segmented into several substructures (Figure 6C). The collar is a large and complex periplasmic structure that is anchored on the cytoplasmic membrane and the MS ring. Underneath the membrane, 16 ringlike particles are localized around the MS ring (Figure 6A, panel 2) and likely correspond to the stator complexes; further genetic mutation of the stator is required to define the exact structure and location of the stator. The C ring of *L. interrogans* exhibits a doughnut-shaped ring below the membrane (Figures 4 and 6C). Underneath the FlhAC ring, a spherical density with a diameter of ~10 nm is observed in most intact motors (Figure 4). This structure has been hypothesized to be the ATPase FliI, as further substantiated in studies of a Δ*fliI* mutant constructed in *C. jejuni*. FliI is a member of the ATPase family and is thought to form a spherical hexamer for protein export. The structures of FlhA and FliI not only are conserved in flagellar motors but also have structural homologues in type III injectisomes, as recently revealed by *in situ* analysis.

Conserved structures of the FlhAC ring and FliI ATPase are observed in the flagellar motor of *L. interrogans* (Figure 6C). Importantly, there are novel “linkerlike” structures with six copies in the inner part and 12 copies in the outer part that extend radially to the bottom of the C ring. The overall architecture of the export apparatus is similar to that in F- and V-type ATPase. It is also consistent with the biochemical data that show that the FlhF/FliI/FliJ complex interacts with both the FlhA ring and the C ring in a comprehensive manner. Clearly, the detailed structure of the export apparatus and its secretion mechanisms will be a fascinating topic in the future.

Figure 6. Asymmetric reconstruction of the *L. interrogans* flagellar motor. A central section of the asymmetric reconstruction is shown in panel A. Panels 1–4 are horizontal cross sections. The locations of the sections are indicated in panel A. Panels 1 and 2 are the putative collar and stator units with 16-fold symmetry. Panels 3 and 4 are the export apparatus with 12- and 6-fold symmetry, respectively. (B) 3D surface rendering of the intact motor displayed in three different views. (C) Segmentation of the motor according to the previous density outline. The complex structures inside the C ring are ascribed to be the export apparatus. The major components (FliI and FlhA) of the export apparatus in *C. jejuni* have been identified by *fliI* deletion and *flhA* cytoplasmic domain truncation mutants, as shown in the insets at the bottom right of panel C. The protein densities are colored black, and the lacking densities of FliI and FlhA are indicated by black arrows.
**ARCHITECTURE OF THE SWITCH COMPLEX**

The C ring is known as the switch complex, which plays a crucial role in flagellar switch, assembly, and rotation. FliG is directly involved in the rotor–stator interaction, while FliM and FliN interact with the signaling protein phosphorylated CheY (CheY-P) to regulate switching between CCW and CW rotation. Atomic structural information is now available for all components of the switch complex, including the full-length or fragmented FliG, FliM, FliN fragments, and the complex of FliG/FliM fragments. Together with intact C rings determined by cryo-EM and more recently by cryo-ET, these high-resolution structures become the building blocks for achieving a pseudoatomic model of the C ring.

A consensus about the relative positions and orientation of FliM and FliN has been reached, while the difference among models of the C ring is the location of the N- and C-terminal domains of FliG. FliN is organized in doughnut-shaped tetramers, which fit well into the bulge density at the bottom of the C ring. A recent crystal structure of the FliM–FliG complex from *Thermotoga maritima* (PDB-4FHR) fits well into the upper density above FliN with the charged helix on FliGC facing upward (Figure 7B). Importantly, the structures of individual domain structures within the complex are similar to those seen in other crystal structures. The FliG–FliM interface within the crystal structure is consistent with the well-defined hydrophobic interaction between residues of the EHPQR motif in FliG and the GGXG motif in FliM. The globular N-terminal domain of FliG from *Aquifex aeolicus* can also be fit to the inner lobe density of the C ring. This model is consistent with the biochemical data that show that the C-terminal domain of FliG (FliGC) interacts with the stator protein MotA while the N-terminal domain of FliG interacts with the C-terminal domain of FliF.

The C ring in many different species is in a conformation similar to that in *S. enterica* (Figure 4). However, the C ring in *Leptospira* appears to be strikingly different from others (Figure 4). Apart from the apparent extra density at the bottom of the C ring, the orientation of the upper portion is also different. Similar to those of *Bacillus subtilis* and *T. maritima*, the genome of *L. interrogans* contains two independent genes, *fliY* and *fliN*, while there is no *fliY* in *S. enterica* or *B. burgdorferi*. FliY is a flagellar protein in the CheC phosphatase family. The recent crystal structure of *L. interrogans* fits well into the extra density (Figure 7D), suggesting that FliY is closely associated with the C ring. FliY binding likely plays a critical role in the conformational changes of the switch complex that ultimately lead to the reversal of the direction of rotation. Clearly, further study is needed to understand FliY and its impact on the switch complex and flagellar switch.

**CONCLUDING REMARKS**

The bacterial flagellum is one of the most thoroughly studied prokaryotic motility organelles. Our understanding of this molecular machine has advanced dramatically over the past several decades. More atomic structures of flagellar components are emerging. However, the structure and function of the intact flagellar motor are far from fully understood at the molecular level. Many novel flagellar proteins and other motility-related proteins (YcgR, CheY-P, etc.) are often associated with the flagellar components, increasing the complexity of the flagellar structure and function. The emergence of cryo-ET and subtomogram averaging provide new avenues for studying intact flagellar motors in cells with unprecedented detail. It is expected that significant advances in cryo-ET, in combination with subtomogram averaging and molecular tools, will provide novel structural insights into many important processes of bacterial flagella: the stator–rotor interaction, protein secretion and assembly, and switching and rotation.

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**Figure 7.** Molecular architecture of the switching complex. (A) Domain organization of the switching complex in *S. enterica*: FliG, FliM, and FliN. (B) Atomic models of FliG (PDB-3hjl), the FliGMC–FliM complex (PDB-4fhr), and FliN (PDB-1yab) were fit in the EM density map of *S. enterica* as described previously. (C) Domain organization of the switching complex in *L. interrogans*: FliG, FliM, FliN, and FliY. The FliY middle domain is homologous to the middle domain of FliM. The C-terminal domain of FliY resembles FliN. (D) The atomic models of FliN, the FliM–FliGMC complex, and FliG were docked into the C ring map of *L. interrogans* as a rigid body. The atomic model of FliYM (PDB-4fhy) was fit into the extra density on the C ring.
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