Structural and Functional Comparison of Salmonella Flagellar Filaments Composed of FljB and FliC

Tomoko Yamaguchi1,2,§, Shoko Toma1,§, Naoya Terahara1,3, Tomoko Miyata1, Masamichi Ashihara1,5, Tohru Minamino1, Keiichi Namba1,2,4,5,*, Takayuki Kato1,§,*

1 Graduate School of Frontier Biosciences, Osaka University, 1-3 Yamadaoka, Suita, Osaka 565-0871 Japan; u523219i@fbs.osaka-u.ac.jp (T.Y.); tomax002@umn.edu (S.T.); tearhara.19r@g.chuo-u.ac.jp (N.T.); miya@fbs.osaka-u.ac.jp (T.M.); tohru@fbs.osaka-u.ac.jp (T.M.); keiichi@fbs.osaka-u.ac.jp (K.N.); galy@fbs.osaka-u.ac.jp (T.K.)
2 RIKEN Center for Biosystems Dynamics Research, 1-3 Yamadaoka, Suita, Osaka 565-0871 Japan; u523219i@fbs.osaka-u.ac.jp (T.Y.); keiichi@fbs.osaka-u.ac.jp (K.N.)
3 Faculty of Science and Engineering, Chuo University, Tokyo 112-8551, Japan; tearhara.19r@g.chuo-u.ac.jp (N.T.);
4 RIKEN SPring-8 Center, 1-3 Yamadaoka, Suita, Osaka 565-0871 Japan; keiichi@fbs.osaka-u.ac.jp (K.N.)
5 JEOL YOKOGUSHI Research Alliance Laboratories, Osaka University, 1-3 Yamadaoka, Suita, Osaka 565-0871 Japan; keiichi@fbs.osaka-u.ac.jp (K.N.)
§ These authors contributed equally.
$ Present address: Thermo Fisher Scientific K.K., 4-2-8 Shibaura, Minato-ku, Tokyo 108-0023, Japan; masamichi.ashihara@thermofisher.com (M.A.); Institute for Protein Research, Osaka University, 3-2 Yamadaoka, Suita, Osaka 565-0871 Japan; tkato@protein.osaka-u.ac.jp (T.K.)
* Correspondence: keiichi@fbs.osaka-u.ac.jp (K.N.); galy@fbs.osaka-u.ac.jp (T.K.);
Tel.: +81-6-6879-4625 (K.N. & T.K.)

Abstract: The bacterial flagellum is a motility organelle, consisting of a long helical filament as a propeller and a rotary motor that drives rapid filament rotation to produce thrust. Salmonella enterica serovar Typhimurium has two genes of flagellin, fljB and fliC, for flagellar filament formation and autonomously switches their expression at a frequency of 10^-3–10^-4 per cell per generation. We report here differences in their structures and motility functions under high viscosity conditions. A Salmonella strain expressing FljB showed a higher motility than the one expressing FliC under high viscosity. To examine the reasons for this motility difference, we carried out structural analyses of the FljB filament by electron cryomicroscopy and found that the structure is nearly identical to that of the FliC filament except for the position and orientation of the outermost domain D3 of flagellin. The density of domain D3 was much lower in FljB than FliC, suggesting that domain D3 of FljB is more flexible and mobile than that of FliC. These differences suggest that domain D3 plays an important role not only in changing antigenicity of the filament but also in optimizing motility function of the filament as a propeller under different conditions.

Keywords: bacterial flagellar motility; flagellin; Salmonella; FljB; FliC; electron cryomicroscopy; viscosity; infection

1. Introduction

Salmonella infection is one of the four major causes of disease involving diarrheas in the world. Salmonella enterica serovar Typhimurium (hereafter Salmonella) has a wide range of hosts, and it infects not only mouse, which is its original host, but also human. The infection is mainly through oral intake, and flagellar motility plays an important role in infection. The flagella enable bacteria to move through viscous environments such as mucosa, search for the host cell surface and adhere to the cell membrane for infection [1,2].
The bacterial flagellum consists of three main parts: the basal body that works as a rotary motor, the filament that functions as a screw propeller, and the hook as universal joint connecting the filament to the motor [3]. *Salmonella* has several peritrichous flagella, and the length of the filament is 10 to 15 µm long. When the cell swims straight, the motors rotate counter clockwise (CCW), and the normally left-handed supercoiled filaments form a bundle behind the cell to produce thrust. When the motors switch their rotation to the clockwise (CW) direction, the filaments switch to a right-handed supercoil in order for the bundle to fall apart so that the cell can change its orientation by tumbling to change the direction of swimming. *Salmonella* has two flagellin genes, *fljB* and *fliC*, and their expression is autonomously regulated to produce either FljB or FliC for filament formation at a frequency of $10^{-3}$–$10^{-4}$ per cell per generation. This is called phase variation [4] and is thought to be a mechanism to escape from the host immune system by changing flagellar antigenicity.

It has been reported that FliC expressing bacteria display a significant advantage for invasion of most epithelial cell lines of murine and human origin compared to FljB expressing bacteria. Differences in the swimming behaviors near surfaces have also been observed, and FliC expressing bacteria more frequently "stop" [5]. In order to understand the differences in their antigenicity and motility function, structural and functional characterization is necessary. However, structural information is available only for the FliC filament [6–8].

The three-dimensional (3D) structures of FliC and the FliC filament have been studied by X-ray crystallography and electron cryomicroscopy (cryoEM) helical image analysis, respectively [6–8]. *Salmonella* FliC from the strain SJW1103 consists of 494 amino acid residues, and the molecular mass is about 50 kDa. The molecule consists of four domains, D0, D1, D2, D3, arranged from the inner core to the outer surface of the filament. Domains D0 and D1 form the inner core of the filament and are made of α-helical coiled-coils. These domains play a critical role in forming the supercoiled structure of the filament as a helical propeller. In addition, a β-hairpin structure in domain D1 is considered to be important for switching the conformation of flagellin subunits between the two states to produce various types of supercoiled filaments in left- and right-handed forms for swimming and tumbling [6,7]. Domains D2 and D3 are in the outer part of the filament structure. These two domains increase the stability of the filament structure as well as the drag force of the filament as a propeller by increasing the diameter [6,9]. The outermost domain D3 is thought to contain epitopes for antibodies, determining the antigenicity of the flagella.

To understand the differences in the antigenicity between FljB and FliC, structural information for FljB and the FljB filament is necessary. Functional characterization of cell motility is also necessary to investigate potentially different physiological roles played by these two types of filaments, if any.

In the present study, we therefore investigated differences in the FljB and FliC filament structures and their motility functions. A *Salmonella* strain expressing FljB showed a higher motility than the one expressing FliC under high viscosity conditions. The structure of the FljB filament analyzed by cryoEM image analysis was nearly identical to that of the FliC filament except for the position and orientation of the outermost domain D3. Domain D3 of FljB also showed a higher flexibility and mobility than that of FliC. These differences suggest that domain D3 plays an important role, not only in changing antigenicity but also in optimizing motility function of the filament as a propeller under different conditions. We discuss the relationship between the structure and motility function by comparing FljB and FliC.

2. Materials and Methods

2.1. *Salmonella* strains

Bacterial strains of *Salmonella enterica* serovar Typhimurium used in this study are listed in Table 1. SJW1103 lacks the *fljAB* operon and so expresses only the FliC flagellin. To express FljB from the *fliC* promoter on the chromosome, the Δ*fliC::tetRA* allele was replaced by the *fljB* allele using the λ Red homologous recombination system [10] as described previously [11] to generate the SJW1103B strain. For cryoEM structural analyses, the strain expressing the flagellar filament of the R-type straight form (the right-handed helical symmetry) was used.
Table 1. Strains and plasmids used in this study

<table>
<thead>
<tr>
<th>Salmonella strains</th>
<th>Relevant characteristics</th>
<th>Source or reference</th>
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</thead>
<tbody>
<tr>
<td>SJW1103</td>
<td>FliC wild-type</td>
<td>Yamaguchi et al. 1984 [13]</td>
</tr>
<tr>
<td></td>
<td>Δhin-fljA-fljB</td>
<td></td>
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<tr>
<td>SJW1103ΔC</td>
<td>SJW1103 ΔfliC::tetRA</td>
<td>This study</td>
</tr>
<tr>
<td>SJW1103B</td>
<td>FljB wild-type</td>
<td>This study</td>
</tr>
<tr>
<td></td>
<td>ΔfliC::fljB</td>
<td></td>
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<tr>
<td>SJW590</td>
<td>FljB_R-type straight filament</td>
<td>This study</td>
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<tr>
<td></td>
<td>fliB(A461V), ΔfliC</td>
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2.2. Swimming motility assay

Salmonella strains, SJW1103B (only expressing FljB) and SJW1103 (only expressing FliC), were pre-cultured in 5 ml of Luria-Bertani broth (LB, 1%(w/v) tryptone, 0.5%(w/v) yeast extract, 0.5%(w/v) NaCl) with overnight shaking at 37°C. Bacterial growth was measured via an optical density at 600 nm (OD$_{600}$).

A 5 µl of the culture media was inoculated into 5 ml of fresh LB, and it was incubated for 6 hours at 37°C with shaking. The cells were diluted in the motility buffer (10 mM potassium phosphate pH 7.0, 0.1 mM EDTA, 10 mM sodium D-lactate). The viscosity of the motility buffer was adjusted by adding Ficoll PM400 (GE healthcare, Chicago, Illinois, USA) to a final concentration of 5%, 10% or 15%. Swimming motility was observed using a bright-field microscope, CX-41 (Olympus, Tokyo, Japan) with a 40× objective (PlanC N 40× NA 0.65 Olympus) and a 1.25× variable magnification lens (U-CA 1.5× Olympus) and recorded by a high-speed camera (GEV-B0620M-TC000 IMPREX, Boca Raton, FL, USA). The swimming speed was calculated by a software Move-tr2/2D (Library Co., Tokyo, Japan).

2.3. Fluorescence labeling of flagellin antibodies

IgG antibodies against FliC and FljB were purified from rabbit serum using a Melon™ Gel IgG Spin Purification Kit (Thermo Fisher Scientific, Waltham, MA, USA). Its buffer was changed to a phosphate buffered salt solution (pH 7.4) simultaneously with concentration of IgG by using a YM-50 Centriprep centrifugal filter (Merck, Darmstadt, Germany). After adjusting the concentration of IgG to 1 mg/ml, the antibodies were labelled with an Alexa Fluor™ 594 Antibody Labeling Kit (Thermo Fisher Scientific) for the FliC antibody and with an Alexa Fluor™ 488 Antibody Labeling Kit (Thermo Fisher Scientific) for the FljB antibody.

2.4. Immunofluorescent staining of the flagellar filament

SJW1103B and SJW1103 were cultured under the same conditions used for the swimming motility assay described above. The culture was diluted 50 to 100 times with the motility buffer. To label the flagellar filaments, the cells were first attached to the surface of a slide glass coated with polylysine by filling the culture medium under a cover slip for 5 minutes. Then the culture medium was replaced with a 20 µl solution of fluorescence-labeled antibodies at 25 µg/ml, and it was left for 5 to 10 minutes. The excess antibody dye was washed by gently flowing 40 µl of the motility buffer twice, and the filaments were then observed by fluorescence microscopy. These observations were performed under an IX-83 optical microscope with a UPlan-SApo 100× 1.4NA objective lens (Olympus). The number of the flagellar filaments per cell was measured using ImageJ (National Institutes of Health, Bethesda, MD, USA).

2.5. Flagellar filament purification

SJW590 was pre-cultured in 30 ml LB with shaking overnight at 37 °C, and then 15 ml of the culture was inoculated into 1 L of fresh LB. The cells were grown until the OD$_{600}$ value reached about 1.0.
culture was collected by centrifugation and resuspended in 10% sucrose buffer (10%(w/v) sucrose, 0.1 M Tris-HCl pH 8.0). EDTA (pH adjusted by HCl to 7.8) and lysozyme were added to final concentrations of 10 mM and 0.1 mg/ml, respectively. The suspension was stirred on ice for 1 hour at 4°C, and 0.1 M MgSO4 and 10% (w/v) Triton X-100 were then added to final concentrations of 10 mM and 1%(w/v), respectively. After stirring for 1 hour at 4°C, 6 ml of 0.1 M EDTA (pH adjusted by NaOH to 11.0) was added. The solution was centrifuged at 15,000×g, and the supernatant was collected. The pH was adjusted to 10.9 with 5 N NaOH, and the sample solution was re-centrifugated at 15,000×g to remove undissolved membrane fractions. The supernatant was centrifuged at 67,000×g to collect the flagellar filament with the hook basal body attached (the filament) as a pellet. The sample was resuspended in 1 ml of Buffer C (10 mM Tris-HCl pH7.8, 5 mM EDTA, 1% (w/v) Triton X-100) and was centrifuged at 7,300×g, and the supernatant was collected. The filament was purified by a sucrose density-gradient centrifugation method with a gradient of sucrose from 20% to 50%. The fraction containing the filament was collected and checked by SDS-PAGE. The fraction was two-fold diluted by Buffer A (20 mM Tris-HCl pH7.8, 150 mM NaCl, 1 mM MgCl2), and the sucrose was removed by centrifugation of the filament at 104,300×g. Finally, the filament was resuspended with 10 - 50 µl of Buffer A, and the sample solution was stored at 4 °C.

2.6. Negative staining

A 5 µl of the sample solution was mixed with 2% PTA on Pala film and was placed on a thin carbon coated, glow discharged copper grid. The extra solution was removed from the grid by blotting, and the grid was dried for 1 hour at room temperature. The sample was checked by a transmission electron microscope, JEM-1011 (JEOL, Akishima, Japan) with an accelerating voltage of 100 kV.

2.7. Electron cryomicroscopy and image processing

A 3 µl of the sample solution was applied to a Quantifoil holey carbon grid R1.2/1.3 Mo 200 mesh (Quantifoil Micro Tools GmbH, Großlöbichau, Germany) with pretreatment of both sides of the grid by glow discharge. The grids were blotted and plunged into liquid ethane at the temperature of liquid nitrogen for rapid freezing [9,14] with Vitrobot Mark IV (Thermo Fisher Scientific). Then grids were transferred into electron cryomicroscopes with a cryostage cooled by liquid nitrogen. The frozen hydrated specimens of the FljB filament on the grid were observed with a Titan Krios electron cryomicroscope (Thermo Fisher Scientific) operated at an accelerating voltage of 300 kV. Dose-fractionated movie images were recorded on a direct electron detector camera, Falcon II (Thermo Fisher Scientific) and were automatically collected using the EPU software (Thermo Fisher Scientific). Using a minimum dose system, all movies were taken by a total exposure of 2 seconds, an electron dose of 10.3 electron/Å² per frame; a defocus range of 0.2 to 1.9 μm, and a nominal magnification of 75,000×, corresponding to an image pixel size of 1.06 Å. All the seven frames of the movie image were recorded with a frame rate of 0.1 sec/frame, and the middle five frames from the 2nd to the 6th were used for image analysis. The defocus range was estimated by Gctf [15] after motion correction by RELION-3.0-β2 [16]. Using RELION-3.0-β2, the filament images were segmented and extracted in square boxes of 400 pixels with 90% overlap, and those segment images were aligned and analyzed. The overall resolution was estimated by Fourier Shell Correlation (FSC) = 0.143 (Figure S1). The summary of cryoEM data collection and statistics is shown in Table S1.

The cryoEM 3D density map has been deposited to the Electron Microscopy Data Bank under accession code EMD-9896, and the atomic model coordinates has been deposited to the Protein Data Bank under accession code 6JY0.

3. Results

3.1. Motility difference between the cells with the FljB and FliC filaments
In order to investigate whether the two different types of flagellar filaments affect swimming motility, we first measured the swimming speed and the population of motile cells with the FljB and FliC filaments in the motility buffer solutions of different viscosities adjusted by adding Ficoll. Optical microscopy was used for this observation. A wild-type *Salmonella* LT2 strain has two distinct flagellin genes, *fliC* and *fljB*, on the chromosome, and their expressions are switched autonomously. Therefore, we used two LT2 derivative strains, SJW1103 expressing only FliC and SJW1103B expressing only FljB, to examine structural and functional differences between these two flagellar filaments. These strains both showed the same growth curve (data not shown). The swimming speed of SJW1103 markedly decreased as the Ficoll concentration increased. When Ficoll was added to the final concentration of 10%, the average swimming speed decreased to 44% of that measured in the motility buffer without Ficoll. In contrast, the swimming speed of SJW1103B decreased only to 73% in the motility buffer with 10% Ficoll (Figure 1a). In addition, the decrease in the motile cell population of SJW1103B was much less than that of SJW1103. In the presence of 15% Ficoll in the motility buffer, 72% of SJW1103B cells were swimming while the motile fraction of SJW1103 cells was only 32% (Figure 1b). To examine the morphology, the length and the number per cell of the flagellar filaments, the filaments were stained by immunofluorescence staining after 6 hours of culture in LB and were observed by fluorescence microscopy. On average, most of the cells had 3 to 5 filaments, and their lengths were about 10 µm regardless of the strain (Figure 1c and d).

**Figure 1.** Motility and filament characteristics of *Salmonella* strains SJW1103 and SJW1103B. (a) Changes in the swimming speed, and (b) changes in the swimming cell population, under different viscosity conditions by adding Ficoll to the motility buffer. (c) Number per cell and (d) length of the flagellar filaments. No significant difference is observed under both conditions.

3.2. Structures of the FljB and FliC filaments

We analyzed the structure of the FljB filament by cryoEM single particle image analysis in one of the two mutant straight forms called the R-type to utilize the helical symmetry. The 3D image of the FljB filament was reconstructed at 3.6 Å resolution from about 1.1 million segment images obtained from 2,319 cryoEM movie images (Figure 2, Figure S1, Table S1). It showed a tubular structure formed by helical assembly of subunit proteins with the outer diameter of 230 Å and the inner diameter of 20 Å. The 11 protofilaments can also be recognized as in the FliC filament structure [6–8,17]. As revealed previously [6,7,17], the subunit consists of four domains, D0, D1, D2 and D3, arranged from the inner to the outer part of the tube. Domains D0 and D1 form the inner tubes while...
domains D2 and D3 form the outer part, with domain D3 exposed on the surface of the filament. The sequence regions forming these four domains are indicated in Figure S2, in which the high sequence homology between FljB and FliC are also presented. The helical parameters of the FljB filament determined in this study was also identical to that of the R-type FliC filament [8], which is reasonable considering the high sequence homology between these two flagellins.

![Figure 2. The 3D density map of the FljB filament solved by cryoEM image analysis. (a) Side view of the filament surface, (b) longitudinal section along the filament axis, and (c) cross section viewed from the distal end. The four domains of FljB, D0, D1, D2 and D3, are labeled in different colors in (b) and (c).](image)

![Table 2. The helical parameters of the FljB and FliC R-type straight flagellar filaments](table)

<table>
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<th></th>
<th>rotation (degree)</th>
<th>axial rise (Å)</th>
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<td>FljB</td>
<td>65.81</td>
<td>4.69</td>
</tr>
<tr>
<td>FliC</td>
<td>65.84</td>
<td>4.71</td>
</tr>
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</table>

To build the atomic model of the FljB filament, a homology model was first generated from the atomic model of the FliC filament (PDB ID: 1UCU) [7] by MODELLER [18], then fitted into the 3D density map and refined by UCSF Chimera [19], Coot (Crystallographic Object-Oriented Toolkit) [20] and Phenix [21] (Figure 3b). The model of FljB (PDB ID: 6JY0) in the filament structure contains residues 1–192 and 288–505, which cover domains D0, D1 and D2 (Figure S2). The missing residues 193–287, corresponding to domain D3, were not built because the resolution of the 3D map was too low to trace the chain (Figure 3b). In addition, the density level of domain D3 was much lower than other domains in FljB (Figure 3e), indicating that domain D3 is more mobile than other domains of FljB. This indicates that the two-chain hinge connection between domains D2 and D3 is relatively flexible, making domain D3 mobile. The structures of domains D0 and D1 are nearly identical between FljB and FliC, with a root mean square deviation of C\( \alpha \) atoms being 2.8 Å (Figure 3c). Domain D2 is composed of two subdomains, D2a and D2b, and while the D2b domains (residues 356–412 for FljB and 345–401 for FliC) are also nearly identical between the two, the D2a domains...
(residues 177–189, 294–355 for FljB and 177–189 and 284–344 for FliC) showed a small difference in their conformations due to 4 residue insertion in FljB (Asp 310–Gly 314) (Figure 3c, Figure S2).

**Figure 3.** Comparison of subunit structures of the FliC and FljB filaments. The 3D map and Cα ribbon model of (a) FliC (PDB ID: 1UCU) and (b) FljB (PDB ID: 6JY0). The map of FljB is the one obtained by cryoEM image analysis while that of FliC was calculated from the atomic model. The Cα ribbon model of domain D3 is not shown for FljB because the resolution of this domain was too low to trace the chain. (c) Superposition of the two maps with FljB in light green and FliC in gray. (d) The Cα ribbon model of FljB in light green is superimposed on that of FliC in light pink. The structures of FljB and FliC are nearly identical for domains D0, D1 and D2b but the folding of part of domain D2a connecting to domain D3 are distinct between the two, which is responsible for the tilt of domain D3 of FljB, making its position higher than that of FliC by about 20 Å. (e) The color coded density distribution of FljB in a longitudinal section of the molecule. The relatively low density of domain D3 indicates a high mobility of this domain.

The distinct difference in the two filament structures is the position and orientation of domain D3 relative to D2. When domains D0, D1 and D2 are superimposed between FljB and FliC, domain D3 of FljB is tilted from that of FliC by about 30°, making its position nearly 20 Å higher than that of FliC (Figure 3d). The two antiparallel chains connecting domains D2 and D3 are highly tilted in FljB to generate this difference (Figure 3c). Near this two chain domain connection, FliC has close contacts between Ala 191, Val 283 and Asp 339, with a Cα distance between Ala 191 and Asp 339 of 4.9 Å, making the conformation of this two chain domain connection stable (Figure 4a). Unlike FliC, FljB does not have these contacts, with the corresponding Cα distance between Ala 191 and Thr 350 being 18.4 Å (Figure 4b), possibly making the two chain domain connection less stable and domain D3 oriented differently than FliC. As a result, domain D3 of FljB is more mobile than that of FliC, and the FljB and FliC filaments have different lateral intermolecular interactions between domains D2 and D3 of neighboring protofilaments.
Figure 4. Interactions of residues stabilizing the position and orientation of domain D3 in FliC. (a) The Cα ribbon models of FliC (left) and FljB (right) with three residues that are important for stabilizing the position and orientation of domain D3 in space-filling representation. (b) Magnified view of the parts indicated by the boxes in (a). The ellipse in dashed line in the lower panel indicates the position of domain D3 of FljB. The corresponding residues of FliC and FljB are labeled in the same colors, orange (FliC Ala 191 and FljB Ala 191), blue (FliC Val 283 and FljB Val 292) and green (FliC Asp 339 and FljB Thr 350). In FliC, the three residues are closely packed each other by hydrophobic interactions, stabilizing the position and orientation of domain D3, while they are far apart in FljB as indicated by the distances.

In the FljB filament structure, we newly identified an interesting axial intermolecular interaction between the D0 domains that contributes to the filament assembly and stability. Along each of the 11 protofilaments, the N-terminal five residues of the subunit 1 above (the distal side of the filament) are extended perpendicular to the filament axis before forming the N-terminal α-helix and forms an antiparallel β-strands with the extended C-terminal chain connecting domains D0 and D1 of the subunit 2 below (the proximal side of the filament) (Figure 5). The residue Gln 2 of subunit 1 and Tyr 469 of subunit 2 form a hydrogen bond between the main chains (Figure 5c). This explains why the N-terminus is more important than the C-terminus for the filament stability [22].
Figure 5. The folding and intersubunit interaction of the N-terminal five residues of FljB contributing to the assembly and stabilization of the filament structure. (a) The Ca ribbon models of two neighboring FljB subunits along the protofilament are shown in rainbow color from the N-terminus in blue to the C-terminus in red. (b) The N terminal five residues are extended, lying flat on the short extended chain connecting the C-terminal α-helices of domains D0 and D1 of a subunit below along the protofilament, forming an antiparallel β-strand to stabilize the axial intersubunit interactions. (c) The residue Gln 2 of subunit 1 and Tyr 469 of subunit 2 form a hydrogen bond between the main chains (dashed line in right blue). In figure (c), only main chain back born atoms are shown (nitrogen and oxygen atoms are colored blue and red, respectively).

4. Discussion

To investigate potentially different physiological roles played by the two types of flagellar filaments formed by FljB and FliC, we carried out motility assay of two strains expressing either FljB or FliC and structural analysis of the FljB filament for comparison with the FliC filament. There were no significant differences between the strain SJW1103B producing the FljB filament and SJW1103 forming the FliC filament, either in the number of filaments per cell, their filament length or the swimming speed in the motility buffer. However, their swimming speeds were clearly different under high viscosity conditions, with a markedly smaller reduction in the swimming speed of SJW1103B than that of SJW1103 when the viscosity is increased by the addition of Ficoll in the motility buffer. These results indicated that the differences in the structure and dynamics of their flagellar filaments must be responsible for the difference in their motility.

We therefore examined the structure of the FljB filament by cryoEM single particle image analysis in one of the two mutant straight forms called the R-type to utilize the helical symmetry. The overall structure was very similar to that of the FliC filament except for the position and orientation of the outermost domain D3 exposed on the surface of the filament. Domain D3 of FljB also showed a higher flexibility and mobility than that of FliC. These differences suggest that domain D3 plays an important role not only in changing antigenicity but also in optimizing motility function of the filament as a propeller under different conditions of solvent viscosity.

These differences in the relative position and dynamics of domain D3 are well correlated with the differences in amino acid sequence between FljB and FliC, which are found in regions Val 187–Gly 189 and Ala 284–Asn 285 and residues Asp 192 and Gln 282 of FliC, which form the two antiparallel chains connecting domains D2 and D3 (Figures S2 and S3). In FliC, the position and orientation of domain D3 is stabilized by hydrophobic interactions between Ala 191, Val 283 and Asp 339 (Figure 4a). On the other hand, the distances between corresponding residues in FljB, Ala 191, Val 292 and Thr 350, are much longer than those of FliC, making the stabilizing hydrophobic interactions impossible (Figure 4b). As a result, the D3 domain of FljB is more mobile than that of FliC.
How the differences in the structure and dynamics of the D3 domains on the surface of the FljB and FliC filaments contribute to their different motility functions as a propeller is difficult to answer without actually examining the role of more mobile domain D3 by fluid dynamic simulations with these two filament structures. There must be some advantages with a higher mobility of surface domains of the helical propeller in generating higher thrust under high viscosity conditions, such as in mucosa, which pathogenic bacteria have to swim through to reach the host cells. The sequence of FljB may have been optimized for this purpose by evolution.

5. Conclusions

In this study we revealed that the flagellar filament formed by FljB has an advantage over the FliC filament by allowing cells to have higher motility under high viscosity conditions. Comparing the two structures, the subunit structures and intersubunit packing interactions are well conserved except for the position and mobility of domain D3 on the surface of the filament. The advantage of the FljB filament in the swimming motility under highly viscous conditions may have an important role in infection when they go through viscous layers of mucosa on the surface of intestinal cells or keep a biofilm in good condition by the cells with the FljB filaments acting as a nutrition deliverer [23,24]. However, in order to clarify the relationship between the filament structure and swimming motility of the cell, further computational analyses are needed.

Supplementary Materials: The following are available online at www.mdpi.com/xxx/s1, Figure S1: Fourier shell correlation of the 3D reconstruction, Figure S2: Amino acid sequence alignment of FljB and FliC, Figure S3: Differences in residue numbers and types between FliC and FljB mapped on the FliC model, Table S1: Summary of cryoEM data collection and image analysis.

Author Contributions: conceptualization, K.N. and T.K.; methodology, all authors; swimming analysis, S.T and N.T.; structural analysis, T.Y., S.T. and T.K.; resources, S.T, N.T. and T.M.; writing—original draft preparation, T.Y., T.K. and K.N.; writing—review and editing, all authors; supervision, K.N. and T.K.; project administration, K.N.; funding acquisition, K.N.

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Conflicts of Interest: The authors declare no conflicts of interest.

References


Supplementary materials

Figure S1. Fourier shell correlation of the 3D reconstruction.
Figure S2. Amino acid sequence alignment of FljB and FliC. Conserved residues are shaded in gray. The regions of domains are indicated by bars and labels in different colors underneath the sequence (D0, red; D1, orange; D2a, right blue; D2b, dark blue; D3, green). The dashed line above the sequence indicates the missing regions of the FljB model (6JY0).
**Figure S3.** Differences in residue numbers and types between FliC and FljB mapped on the FliC model. Differences are shown for residues 187 – 192 and 282 – 285 of FliC and residues 187 – 192 and 291 – 294 of FljB. The FliC chain is colored red for those residues that are different from FljB.
Table S1. Summary of cryoEM data collection and image analysis

<table>
<thead>
<tr>
<th>Data collection</th>
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<tbody>
<tr>
<td>Microscope</td>
<td>Titan Krios</td>
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<tr>
<td>Camera</td>
<td>Falcon II</td>
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<tr>
<td>Data acquisition</td>
<td>Auto / EPU</td>
</tr>
<tr>
<td>Voltage (kV)</td>
<td>300</td>
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<tr>
<td>Magnification</td>
<td>75,000</td>
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<tr>
<td>Pixel size (Å)</td>
<td>1.06</td>
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<tr>
<td>Defocus range (µm)</td>
<td>0.2 -1.9</td>
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<td>Frame rate (sec/frame)</td>
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<td>Total exposure time (sec)</td>
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<tr>
<td>Number of frames</td>
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<tr>
<td>Dose rate (e/Å²/frame)</td>
<td>10.3</td>
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<tr>
<td>Total dose (e/Å)</td>
<td>72.1</td>
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<table>
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<th>Particle statistics</th>
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<tbody>
<tr>
<td>Number of micrographs</td>
<td>2,319</td>
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<tr>
<td>Number of picked segments</td>
<td>197,442 (90% overlap)</td>
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<tr>
<td>Number of segments (final)</td>
<td>114,110</td>
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<td>Resolution (Å)</td>
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<td>FSC threshold</td>
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<th>Ramachandran plot</th>
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<td>Favored (%)</td>
<td>94</td>
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<td>Allowed (%)</td>
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<td>Outliers (%)</td>
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