

Analysis of an Engineered *Salmonella* Flagellar Fusion Protein, FliR-FlhB†

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***Salmonella* FliR and FlhB are membrane proteins necessary for flagellar export. In *Clostridium* a *fliR-flhB* fusion gene exists. We constructed a similar *Salmonella* fusion gene which is able to complement *fliR*, *flhB*, and *fliR flhB* null strains. Western blotting revealed that the FliR-FlhB fusion protein retains the FlhB protein's cleavage properties. We conclude that the FliR and FlhB proteins are physically associated in the wild-type *Salmonella* basal body, probably in a 1:1 ratio.**

The bacterial flagellum, a rotary motor, is the organelle responsible for the motility of numerous bacteria, including *Salmonella enterica* serovar Typhimurium. The flagellum consists of a basal body and motor structure, a flexible hook, and a filament. Assembly begins with the formation of the MS ring, which is imbedded in the inner membrane (14). The MS ring houses the export apparatus which is responsible for the translocation of the majority of the flagellar proteins that are external to the cell membrane. Export substrates are translocated across the cytoplasmic membrane, travel through a narrow (ca. 3-nm) channel in the nascent structure (17, 25), and then assemble at the distal end of the growing flagellum (2, 9). The flagellar export apparatus is a member of the type III secretion system family (15), which is also responsible for virulence factor secretion in pathogenic bacteria. In *Salmonella* an apparatus distinct from yet homologous to the flagellar export machinery is present for the secretion of virulence factors (8).

The flagellar export apparatus consists of six integral membrane proteins (FlhA, FlhB, FliO, FliP, FliQ, and FliR) and three associated cytoplasmic proteins (FliH, FliI, and FliJ) (14, 15). Together these proteins are responsible for the export of rod-type, hook-type, and filament-type export substrates (21, 23). FliI is a flagellum-specific ATPase (3) and is regulated by FliH (7, 20, 24). FliJ is a putative general chaperone (5, 18). This note focuses on two of the membrane components, FliR and FlhB.

The membrane components are likely located in a patch of membrane within the central pore of the MS ring (6, 29). There is direct evidence for the presence of FliP and FliR in the basal body (4) and intergenic suppression evidence for the presence of FlhA, which probably interacts with the MS ring protein FliF (12). Very little is known about the functions of FliO, FliP, FliQ, FliR, and FlhA, but their presence is required for flagellum-specific protein export (21). Although FliO, FliP,

FliQ, and FliR are extremely hydrophobic, FlhA and FlhB both have large cytoplasmic domains that interact with FliH, FliI, and FliJ (23). FlhB, along with the hook-length control protein FliK, is responsible for substrate specificity switching from rod- and hook-type export to filament-type export (22). Minamino and Macnab showed that FlhB can be divided into two domains: FlhB_{TM}, comprising the four-membrane-spanning N-terminal domain, and FlhB_C, the cytoplasmic domain (22). The ability to switch is related to the cleavage of FlhB_C into FlhB_{CN} and FlhB_{CC} (22). Fraser et al. have shown that a noncleaving mutant, FlhBN269A, does not switch from rod- and hook-type to filament-type export (6). FlhB_C binds to numerous export substrates (23) as well as FlhA_C and itself (34). Mutations in FlhB_C are able to suppress a FliK null background (32).

In many bacterial genomes, *fliR* and *flhB* are positioned as consecutive genes within the same operon (Fig. 1). In *Salmonella* the two genes are separated by about 55 kb (16). A database search for FlhB homologs found that *Clostridium acetobutylicum* (26) and *Clostridium tetani* (1) have a gene that is equivalent to a fusion of *fliR* and *flhB*. In the present study we show that an engineered *Salmonella* fusion protein has the ability to complement *fliR* and *flhB* null strains of *Salmonella* as well as a *fliR flhB* double-null strain. This strongly suggests that FliR and FlhB are in close proximity to each other within the basal body. In addition, these findings clarify the predicted membrane topology of FliR, which had previously been unclear (27), and firmly establish the presence of FlhB in the basal body.

Bacterial strains and plasmids are listed in Table 1. Luria broth and soft tryptone agar plates were prepared as previously described (23). T7 medium contains, per liter, 20 g of tryptone, 10 g of yeast extract, 5 g of NaCl, and 8.7 g of K₂HPO₄ (pH 7.2). Ampicillin was added to a final concentration of 50 µg/ml.

Construction of the *fliR-flhB* fusion gene. The *fliR-flhB* fusion gene was constructed by a method adapted from Toker et al. (30). Complementary primers coding for the 3' end of *fliR* and the 5' end of *flhB* were used with outside primers to create an NdeI site 5' to *fliR* and a BamHI site 3' to *flhB*. Our fusion protein has a short sequence of four amino acids (GAGA;

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† This paper is dedicated to Robert Macnab, who died suddenly on 7 September 2003.

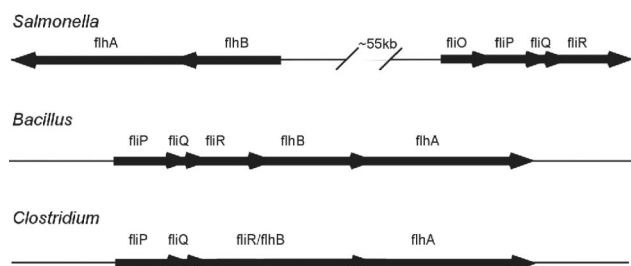


FIG. 1. The genes coding for the integral membrane proteins of the flagellar export apparatus. The genomes of *Salmonella* (16) and *Bacillus subtilis* (13) are representative of the two most common arrangements. In *Salmonella*, *flhA* and *flhB* are located in a separate operon approximately 55 kb from the operon containing *fliO*, *fliP*, *fliQ*, and *fliR*. In *Bacillus* these genes are contained within the same operon, with *flhB* immediately following *fliR*. The arrangement of genes in *Clostridium* (1, 26) is similar to that in *Bacillus*, but a gene homologous to both *fliR* and *flhB* is present in place of the separate genes.

italicized in the sequence) linking the otherwise wild-type sequence in order to maximize the possibility of a functional protein (FliR-SEMPINNP-GAGA-MAESDDDKTEAPT P-FlhB), although there is no such spacer in the *Clostridium* protein (FliR-KLVPIGLIFADSDKTEEATP-FlhB) (26).

The *fliR* and *flhB* portions of the fusion gene were amplified from plasmids pRMK3000 and pMM26, respectively, by using *Pfu* Turbo DNA polymerase (Stratagene). These PCR products were then used as templates for a second round of PCR to construct the fusion gene. Amplified DNA was purified by using the QiaQuick gel extraction kit (QIAGEN Inc.), digested with NdeI and BamHI, and cloned into pUC18. The fusion gene was then subcloned into the pTrc99A-FF4 and pET vectors for use in protein expression studies. The construction of the fusion gene *fliR-flhBN269A* used the same protocol, but pGFBN269A was used in place of plasmid pMM26. As a con-

trol, we also constructed a plasmid containing *fliR* and *flhB* as separate but consecutive genes by cloning the *fliR* gene with NdeI and BamHI and the *flhB* gene with XbaI and HindIII into pTrc99A-FF4.

Construction and complementation of null strains. The *fliR* and *flhB* null strains were constructed as previously described (11). The *fliR flhB* double-null strain was constructed by transferring the *flhB* null genotype to SJW170, a *fliR* mutant strain containing a 10-bp deletion resulting in a frameshift after amino acid 89 and having 27 foreign amino acids before reaching a stop codon. Cells were tested for their swarming ability on soft tryptone agar plates as described by Fraser et al. (5).

Complementation results are shown in Fig. 2. Strains MKM50 (*flhB* null) and MKM65 (*fliR* null) are complemented by plasmids pMM26 and pRMK3000, respectively. These null strains are also complemented by pJSV004 (fusion plasmid) and pJSV024 (coproducing plasmid). The fusion protein can complement MKM58 (*fliR flhB* double null) although providing FliR and FlhB as separate proteins (pJSV024) allows for improved swarming. This indicates that the fusion protein retains the functions of both FliR and FlhB. The complementation of MKM65 by pJSV004 is better than its complementation of the other null strains. This result could be due to the presence of wild-type FlhB (in this *fliR* null strain), which is not present in either of the other two null strains. This suggests that wild-type FlhB may be present in the basal body in addition to the FliR-FlhB fusion protein.

Although plasmid pJSV014, containing the FliR-FlhB fusion protein with the noncleavable FlhBN269A mutation, does not complement either strain MKM50 or MKM58, it does complement strain MKM65 almost as well as plasmid pRMK3000. This finding indicates that the FliR portion of FliR-FlhBN269A is able to function in the export process although it is fused to the noncleaving FlhB variant which is locked in

TABLE 1. Strains and plasmids used in this study

Strain or plasmid	Description	Source or reference
Strains		
<i>E. coli</i>		
NovaBlue	Recipient for cloning experiments	Novagen
BL21(DE3)/pLysS	Used for overproduction of proteins from pET-based vectors	Novagen
<i>Salmonella</i>		
JR501	Used for converting <i>E. coli</i> plasmids to <i>Salmonella</i> compatibility	28
SJW1103	Wild type	33
SJW170	<i>fliR</i> null	33
MKM50	<i>flhB</i> null strain	6
MKM65	<i>fliR</i> null strain	This study
MKM58	<i>fliR flhB</i> double-null strain	This study
Plasmids		
pET19b	N-terminally His-tagged T7 expression vector	Novagen
pET19bFH	N-terminally His-FLAG-tagged T7 expression vector	4
pTrc99A-FF4	Modified pTrc expression vector	27
pMM26	pTrc99A-FF4/FlhB	22
pRMK3000	pET19b/N-His-FliR	This study
pGFBN269A	pet19b/N-His-FlhBN269A	6
pJSV003	pET19bFH/N-His-FLAG-FliR-FlhB	This study
pJSV004	pTrc99A-FF4/FliR-FlhB	This study
pJSV013	pET19bFH/N-His-FLAG-FliR-FlhBN269A	This study
pJSV014	pTrc99A-FF4/FliR-FlhBN269A	This study
pJSV024	pTrc99A-FF4/FliR + FlhB	This study

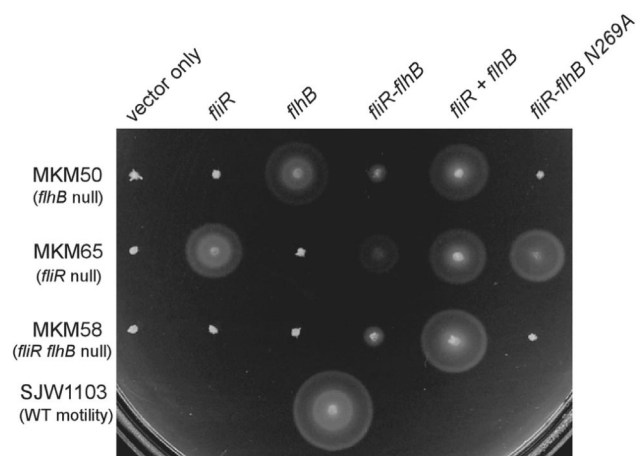


FIG. 2. Complementation analysis in null backgrounds. The plate was incubated at 30°C for 5 h. Column 1, vector only; column 2, pRMK3000 (FliR); column 3, pMM26 (FlhB); column 4, pJS004 (FliR-FlhB fusion); column 5, pJSV024 (FliR + FlhB coproducing plasmid); and column 6, pJSV014 (FliR-FlhBN269A); WT, wild type.

rod- and hook-type export. This could mean that in the *fliR* null strain the cleaved wild-type FlhB from the chromosome can interact with FlhBN269A of the fusion protein, enabling it to function at a level near that of the wild-type. Supporting this idea, several species have a separate gene encoding FlhB_{CC} in addition to full-length FlhB (15). Since FlhB_C can interact with itself (34), oligomerization of FlhB_C may be necessary for full function.

To examine the dominant-negative effects of the fusion protein in wild-type cells, we tested the swarming of strain SJW1103 transformed with plasmids pJSV004, pJSV024, and pJSV014. In the presence of 0.1 mM IPTG (isopropyl-β-D-thiogalactopyranoside), we saw a decrease in swarming with all three plasmids, but especially with the two fusion plasmids. This finding indicates that overproduction of these proteins can interfere with the assembly of the export apparatus (data not shown).

Purification of the FliR-FlhB fusion protein. In order to confirm that the FliR-FlhB fusion construct remains intact and retains the cleavage properties of FlhB, we purified the fusion protein. We found it necessary to use a system yielding high levels of protein expression in order to detect the FliR-FlhB fusion protein. In *Salmonella* we never detected the fusion protein under a variety of culture conditions. Cultures of *Escherichia coli* BL21(DE3)/pLysS transformed with plasmids pJSV003 and pJSV013, coding for N-His-FLAG-FliR-FlhB and N-His-FLAG-FliR-FlhBN269A, respectively, were grown in T7 medium (3 liters) at 30°C to an optical density at 600 nm of 0.4. IPTG was then added to a final concentration of 0.2 mM, and growth continued for an additional 4 h. Cells were harvested and resuspended in 120 ml of lysis buffer (50 mM sodium phosphate [pH 8.0], 500 mM NaCl, 20% glycerol, 10 mM β-mercaptoethanol, and 0.1 mM phenylmethylsulfonyl fluoride), sonicated, and centrifuged at 10,000 × *g* to pellet unbroken cells and inclusion bodies. The supernatant was then centrifuged at 100,000 × *g* for 45 min. Pellets were retained and resuspended by homogenization in 15 ml of solubilization buffer (lysis buffer with 6 M urea, 1% [vol/vol] Triton X-100,

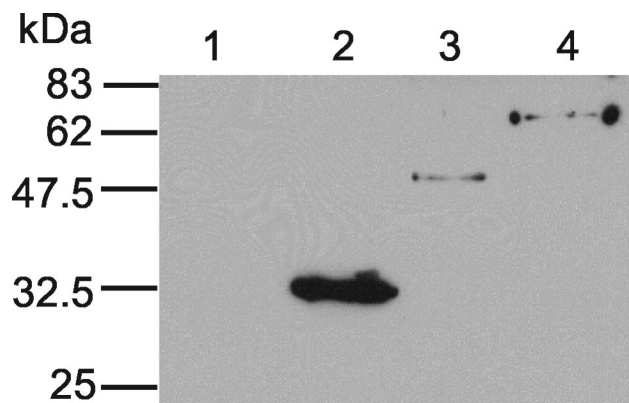


FIG. 3. Western blot of FLAG-tagged FliR-FlhB fusion constructs. Lane 1, vector only; lane 2, N-His-FLAG-FlhB; lane 3, N-His-FLAG-FliR-FlhB; and lane 4, N-His-FLAG-FliR-FlhBN269A. Mass markers (kDa) are given on the left.

and 10 mM imidazole). This suspension was stirred at 4°C for 1 h and then subjected to centrifugation at 100,000 × *g* for 45 min. The denatured, solubilized membrane fraction was retained as the supernatant.

The supernatant was loaded by gravity flow onto a 250-μl Talon (BD Biosciences) metal affinity column, washed with 20 column volumes of wash buffer (solubilization buffer with 20 mM imidazole), and eluted in the same buffer with 250 mM imidazole.

Western blotting. Samples to be probed were run on sodium dodecyl sulfate–15% polyacrylamide gels and transferred onto nitrocellulose membranes. Membranes were probed by using anti-FLAG M2 antibody (Sigma) and horseradish peroxidase-conjugated goat anti-mouse secondary antibody (Bio-Rad). Immunodetection was carried out with the SuperSignal Pico detection kit (Pierce) according to the manufacturer's instructions.

Western blotting reveals that the FliR-FlhB fusion remains intact and that normal posttranslational processing of FlhB occurs (Fig. 3, lane 3). The band migrated at ~60 kDa, consistent with an intact fusion protein in which the FlhB component has been normally cleaved at N269 (the cleavage product would not be seen with anti-FLAG antibodies). Furthermore, the uncleavable N269A fusion variant (lane 4) migrated at ~75 kDa. A vector-only control (lane 1) and wild-type N-His-FLAG-FlhB (lane 2) are also shown.

FlhB is present in the basal body. Although FlhB has been assumed to be present in the basal body due to its role in substrate specificity switching (6), attempts to locate FlhB within the basal body have been unsuccessful, probably because the detergent used in the basal body purification procedure solubilized FlhB. The present study has shown that a FliR-FlhB fusion protein can function in *Salmonella* although not as well as in the wild-type situation where the two proteins are separate. This result places FlhB within the basal body since it has previously been shown that FliR is located on the cytoplasmic face of the MS ring, probably within a membrane patch at the center of the ring (4).

Implications for export apparatus component stoichiometries. The fusion protein is functional in a *fliR flihB* double-null background, suggesting that FliR and FlhB must be present in

at least a 1:1 ratio in the wild-type basal body, in agreement with previous estimates of about 1 to 3 copies of FliR (4, 10) and about 2 copies of FlhB (34). However, the improved swarming with the fusion protein in the *fliR* null background compared to the *flhB* null background suggests that wild-type FlhB (produced from the chromosome of strain MKM65) could also be playing a role in the export process. Given this possibility, we can conclude that there is at least one FlhB for every FliR.

Topology of FliR. The present results also address the question of the membrane topology of FliR. Previously it was predicted that FliO has a single transmembrane helix, FliP could have three or four transmembrane spans, and FliQ is probably a two-span, hairpin protein (27). However, it was difficult to predict the number of helices for FliR since its C-terminal region is almost exclusively hydrophobic and could contain five or six helices. We concluded then that the C terminus of FliR could lie either in the periplasm (five spans) or in the cytoplasm (six spans). We also knew that FliR is a very insoluble protein and can form aggregates even in harsh detergent conditions (e.g., 4% sodium dodecyl sulfate) (4), consistent with a six-transmembrane-spanning protein with very little soluble sequence. It is now known that the C terminus of FliR can be fused to the cytoplasmically located N terminus of FlhB (19) and still function in the export apparatus, suggesting that FliR has six transmembrane spans and that its C terminus lies in the cytoplasm.

The presence of natural fusion proteins in export apparatus homologs underscores the importance of genomic databases in protein homology research, suggesting interactions not readily detectable due to problems encountered in studying membrane proteins with unknown functions. In *Buchnera aphidicola* (31), a gene fusion homologous to both *fliO* and *fliP* exists, suggesting another potential interaction.

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