FliO Regulation of FliP in the Formation of the Salmonella enterica Flagellum

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Abstract

The type III secretion system of the Salmonella flagellum consists of 6 integral membrane proteins: FlhA, FlhB, FliO, FliP, FliQ, and FliR. However, in some other type III secretion systems, a homologue of FliO is apparently absent, suggesting it has a specialized role. Deleting the fliO gene from the chromosome of a motile strain of Salmonella resulted in a drastic decrease of motility. Incubation of the ΔfliO mutant strain in motility agar, gave rise to pseudorevertants containing extragenic bypass mutations in FliP at positions R143H or F190L. Using membrane topology prediction programs, and alkaline phosphatase or GFPuv chimeric protein fusions into the FliO protein, we demonstrated that FliO is bitopic with its N-terminus in the periplasm and C-terminus in the cytoplasm. Truncation analysis of FliO demonstrated that overexpression of FliOA43–125 or FliO1–95 was able to rescue motility of the ΔfliO mutant. Further, residue leucine 91 in the cytoplasmic domain was identified to be important for function. Based on secondary structure prediction, the cytoplasmic domain, FliO43–125, should contain beta-structure and alpha-helices. FliOA43–125-Ala was purified and studied using circular dichroism spectroscopy; however, this domain was disordered, and its structure was a mixture of beta-sheet and random coil. Coexpression of full-length FliO with FliP increased expression levels of FliP, but coexpression with the cytoplasmic domain of FliO did not enhance FliP expression levels. Overexpression of the cytoplasmic domain of FliO further rescued motility of strains deleted for the fliO gene expressing bypass mutations in FliP. These results suggest FliO maintains FliP stability through transmembrane domain interaction. The results also demonstrate that the cytoplasmic domain of FliO has functionality, and it presumably becomes structured while interacting with its binding partners.

Citation: Barker CS, Meshcheryakova IV, Kostyukova AS, Samatey FA (2010) FliO Regulation of FliP in the Formation of the Salmonella enterica Flagellum. PLoS Genet 6(9): e1001143. doi:10.1371/journal.pgen.1001143

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Funding: CSB, IVM, and FAS were supported by Okinawa Institute of Science and Technology Promotion Corporation internal funding. ASK was supported by the UMDNJ Foundation. The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Competing Interests: The authors have declared that no competing interests exist.

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Introduction

For many bacteria, locomotion is possible using the flagellum, which functions like a helical propeller. It is a highly complex nanomachine consisting of about 30 different proteins, and it is organized into three substructures: the basal body, the hook, and the filament. Export of the components of the flagellum across the cytoplasmic membrane requires a specialized secretion apparatus at its base, which shares homology to the type III secretion system apparatus paralogue InvA from Salmonella enterica serovar Typhimurium, the secretion apparatus is postulated to consist of six integral membrane proteins: FlhA, FlhB, FliO, FliP, FliQ, and FliR; and three cytoplasmic proteins: FlhD, FlhF, and FljE [2,3]. Biochemical and genetic studies have determined the location of the membrane proteins FlhA, FlhB, FliP, and FliR to be within the flagellar basal body [4–6]. It has been demonstrated that the hook-capping protein (FliD) and hook protein (FlgE) required all of the proteins of the secretion apparatus for their export [7]. A detailed picture of the workings of the secretion apparatus is gradually being elucidated [8,9]. It has been demonstrated that the secretion apparatus harnesses the proton motive force to drive export of the external flagellar components [10,11]. FlhA and FlhB, the two largest membrane proteins of the flagellar secretion apparatus, which both have predominant C-terminal cytoplasmic domains have been most characterized. The crystal structures of the cytoplasmic domains of FlhA from Salmonella and Helicobacter pylori, and the crystal structure of the cytoplasmic domain of the type III secretion system apparatus paralogue InvA from Salmonella have recently been solved [12–14]. Also, several crystal structures of the cytoplasmic domains of paralogues of FlhB found in virulence-associated needles from enteric bacteria have recently been described [15–17].

The cytoplasmic domains of FlhA and FlhB form a docking platform for FlhD, FlhF, and FljE. The cytoplasmic domain of FlhA has been shown to bind the FlhD, FlhF, and FljE proteins, and it is also thought to be directly involved in the translocation of the export substrates into the central channel of the growing flagellar structure [18–22]. The cytoplasmic domain of FlhB has been shown to undergo autocleavage associated with interaction with FlhK, which switches specificity of export of rod/hook-like substrates to filament-type substrates [23–29]. The N-terminal transmembrane region of FlhA has been implicated to interact with the surrounding MS ring, from studies involving the isolation of extragenic suppressor mutations [5].

Less is known about the functional role of the FliO, FliP, FliQ, and FliR proteins in the secretion complex, though the protein
products of the corresponding genes have been determined [30–35]. FliP, FliQ, and FliR are very hydrophobic and were predicted to be predominantly located within the cytoplasmic membrane. However, FliO was predicted to be a bitopic membrane protein with a predominant soluble domain. Intriguingly, FliO shows the least conservation among the secretion system apparatus membrane proteins, even being absent from some systems [3,34,35]. Notably, the type III secretion system apparatus of the virulence-associated needles of *Salmonella enterica*, which consists of 6 integral membrane proteins: FliA, FliB, FliO, FliP, FliQ, and FliR. Among these proteins, FliO shows a sporadic distribution in bacteria, and its function is unknown, suggesting it might have a specialized role to play where it is present. In this study, we show that FliO has an important role in maintaining stability of FliP, which is a highly conserved member of the secretion apparatus. We have characterized the important regions of FliO through mutagenesis. We have shown that it is possible to bypass the effect of not producing the FliO protein, by encoding mutations within FliP or by overexpressing the cytoplasmic domain of FliO only.

**Author Summary**

The propeller-like flagella, which some bacteria use to swim, possess a specialized secretion apparatus, which is imbedded in the cell membrane for their formation. The components are highly conserved among flagella systems and also to the Type III secretion apparatus used by some bacteria in conjunction with virulence-associated needle complexes. The ubiquity of these secretion apparatuses and their function as intricate nanomachines has made them fascinating for biologists. The most studied flagellar system is that of *Salmonella enterica*, which consists of 6 integral membrane proteins: FliA, FliB, FliO, FliP, FliQ, and FliR. Among these proteins, FliO shows a sporadic distribution in bacteria, and its function is unknown, suggesting it might have a specialized role to play where it is present. In this study, we show that FliO has an important role in maintaining stability of FliP, which is a highly conserved member of the secretion apparatus. We have characterized the important regions of FliO through mutagenesis. We have shown that it is possible to bypass the effect of not producing the FliO protein, by encoding mutations within FliP or by overexpressing the cytoplasmic domain of FliO only.

**Results**

Incubation of a ΔfliO mutant in motility agar gave rise to pseudorevertants containing bypass mutations in fliP

We first sought to characterize the FliO protein through mutagenesis to investigate its function. To provide a genetic background for complementation studies, a ΔfliO mutant was engineered from the wild-type strain SJW1103. We found that the ΔfliO mutant, CB186, was not completely non-motile, and displayed a weakly motile phenotype after several hours in motility agar. Surprisingly, pseudorevertants arose from this strain with enhanced motility after about 42 hours extended incubation (Figure 1A). In comparison strain CB184, which encodes a non-polar ΔfliOΔfliP deletion was completely non-motile (Figure 1B). Four motile pseudorevertants from CB186 were purified, and the two most motile were characterized further. These two strains were designated CB191 and CB227.

To screen for the bypass mutations the genomic DNA of CB191 and CB227 was isolated, and the fliA, fliB, fliO, fliP, fliQ, and fliR genes were sequenced along with the fliP gene encoding the MS ring, which surrounds the secretion apparatus in the membrane. These genes were considered the most likely to contain suppressor mutations. Each pseudorevertant contained a point mutation in the fliP gene only. CB191 encoded an R143H mutation in FliP, and CB227 encoded an F190L mutation in FliP.

To investigate the physiological effects of the extragenic bypass mutations in fliP, engineered strains were constructed from SJW1103 using ΔRed genetic engineering (Figure 1C). Engi-

neered mutants encoding fliPΔR143H or fliPΔF190L mutations only showed the same motility phenotype as SJW1103. Moreover, an engineered mutant encoding ΔfliO fliPΔR143H was equally as motile as CB227, which confirmed that the fliPΔF190L mutation was responsible for the extragenic suppression of the ΔfliO deletion in strain CB227. However, an engineered strain encoding ΔfliO fliPΔR143H was not as motile as CB191. This appeared to suggest that the fliPΔR143H mutation was not the suppressor mutation responsible for the improved motility in pseudorevertant strain CB191. However, the engineered ΔfliO fliPΔR143H mutant gave rise to pseudorevertants with enhanced motility after a much shorter incubation time of 26 hours in motility agar in comparison to a ΔfliO mutant, which took at least 42 hours (Figure 1D). Therefore, in strain CB191 an additional bypass mutation must be encoded along with fliPΔR143H to overcome the ΔfliO deletion.

The FliO N-terminus is in the periplasm and its C-

The above results suggested that FliO is regulating FliP so we decided to characterize the FliO protein further to determine how this might occur. Prior to this study the membrane topology of FliO was unknown, so we started by determining the topology of FliO. We used various prediction programs for the determination of the topology of transmembrane proteins, and the majority suggested that the 125-residue FliO protein is bitopic with a short N-terminal periplasmic domain from residues 1 to 16–22, a transmembrane region between residues 17–23 to 39–43, and a large C-terminal cytoplasmic domain from residues 40–44 to 125 (Table S1). To experimentally determine the membrane topology of FliO we constructed plasmids to express chimeric gene fusions of the *phoA* gene encoding the mature form of alkaline phosphatase, at three specific points into the fliO gene. The mature form of alkaline phosphatase is only active when found in the periplasm, and its use is well established to determine membrane topology of transmembrane proteins [36–38]. The entire amino acid sequence of FliO was present in fusion product; and alkaline phosphatase was encoded between residues 6 and 7, 100 and 101, or 115 and 116 of FliO (Figure 2A). These fusion sites were permisive for motility (data not shown). Immunoblotting using anti-alkaline phosphatase antibody against whole cell lysates of cells expressing the fusions, and the lysates fractioned into insoluble membrane pellet fractions and soluble supernatant fractions demonstrated that the three FliO/PhoA chimeras were expressed stably and mainly partitioned with the insoluble membrane pellet fraction (Figure 2B). To detect expression of alkaline phosphatase, cells expressing the plasmid encoded fusions were inoculated onto L-agar plates containing 40 μg ml−1 5-Bromo-4-chloro-3-indolyl phosphate (Figure 2C). The phoA gene was deleted from the chromosome of the strains used in this experiment, to reduce background phosphatase expression. Alkaline phosphatase activity was detectable only with the FliO125::PhoA125::FliO125 fusion. These fusions were next genetically engineered onto the chromosome of *Salmonella*. Alkaline phosphatase assays were performed, and higher phosphatase
activity was measured for the FltO$_{43-125}$::PhoA$_{22-471}$::FltO$_{7-125}$ fusion, compared to the other chimeras, whether it was expressed from the chromosome only or also expressed from plasmid pTrc99A-FF4 (Table S2). This revealed that the N-terminus of FltO is in the periplasm.

GFPuv has been reported to be useful as a reporter for membrane protein topology in bacteria and it is fluorescent when it is located in the cytoplasm [39]. We constructed plasmids to encode chimeric gene fusions of the GFPuv gene between residues 6 and 7, or 115 and 116 of FltO (Figure 2D). These fusions were permissive for motility (data not shown). Immunoblotting using anti-GFP antibody against whole cell lysates of cells expressing the fusions, and the lysates fractioned into insoluble membrane pellet fractions and soluble supernatant fractions demonstrated that the two FltO/GFPuv chimeras were expressed stably and mainly partitioned with the insoluble membrane pellet fraction. However, a difference in migration between the two fusions is apparent, and the FltO$_{43-125}$::GFPuv::FltO$_{16-125}$ fusion produced multiple bands (Figure 2E). The reason for this is not clear. In previous studies, multiple bands of FltO were detected by immunoblotting, which did not appear to be physiologically important. We presume that the chimeric fusions we have constructed are somehow affecting this N-terminal modification of FltO. However, fluorescence was detectable only with the FltO$_{1-115}$::GFPuv::FltO$_{116-125}$ fusion (Figure 2F). This revealed that the C-terminus of FltO is in the cytoplasm.

Mutational analysis of FltO reveals the cytoplasmic domain of FltO is functionally important

Having defined the membrane topology of FltO, we next sought to determine the functionally important regions of the FltO protein by examining the ability of FltO proteins truncated from the N-terminus or the C-terminus to rescue the motility of a ΔfltO strain in motility agar. The truncated proteins were expressed from plasmid pTrc99A-FF4 without IPTG induction (Figure 3A). Immunoblotting using polyclonal anti-FltO$_{43-125}$-6xHis antibodies was performed first to confirm expression of the proteins. Full-length FltO$_{1-125}$, FltO$_{1-65}$, FltO$_{1-75}$, and FltO$_{1-115}$ were detected by immunoblotting (Figure 3B), FltO$_{1-125}$, FltO$_{1-65}$, FltO$_{1-105}$, and FltO$_{1-115}$ produced multiple bands of FltO, while FltO$_{1-65}$ and FltO$_{43-125}$ did not. As mentioned in the previous section, it was suggested in a previous study that FltO is subject to N-terminal cleavage or modification by an unknown mechanism and the multiple bands presumably reflect this [33]. It was not possible to detect FltO at physiological levels from whole-cell lysates of SJW1103, therefore, the proteins produced from pTrc99A-FF4 were detectable by immunoblotting, because they were being over-expressed from this vector. It was not possible to detect FltO$_{1-65}$, FltO$_{1-75}$, and FltO$_{1-85}$ by immunoblotting, which could be due to the specificity of the antibody or because these proteins were not expressed. A FLAG-tag (N-MDYKDDDDK-C) was engineered at the N-terminus of FltO$_{1-125}$, FltO$_{1-65}$, FltO$_{1-75}$, and FltO$_{1-85}$ to attempt to detect protein expression by immunoblotting using ANTI-FLAG antibody. However, only FLAG-tagged FltO$_{1-125}$ was detectable by immunoblotting, so if FLAG-tagged FltO$_{1-65}$, FltO$_{1-75}$, and FltO$_{1-85}$ were produced expression was much lower (data not shown).

Expression of FltO$_{22-125}$ or FltO$_{43-125}$, which are truncated at the N-terminus, was able to complement the ΔfltO strain CB186 (Figure 3C). FltO$_{22-125}$ encoded a methionine at position 22 in place of a natural valine and this truncation was included since it has been shown to be functional in previous studies [31,33]. Surprisingly, FltO$_{43-125}$ could complement, as this corresponds to the cytoplasmic domain of FltO, and is without the transmembrane domain. Overexpression was necessary for the cytoplasmic domain to rescue motility of CB186, since FltO$_{43-125}$ was not able to complement when expressed from the T7 promoter of plasmid pET-22b(+) , without IPTG induction, while full-length FltO$_{1-125}$ was able to rescue motility when expressed from this vector (data...
not shown). FliO

Residue leucine 91 of the cytoplasmic domain of FliO is important for function

N-terminal and C-terminal truncation analysis of FliO have defined residues 22–95 as the most important. Moreover, overexpression of the cytoplasmic domain of FliO could rescue

Figure 2. The FliO N-terminus is in the periplasm and its C-terminus is in the cytoplasm. (A) Schematic overview of chimeric fusions of alkaline phosphatase within FliO. (B) Immunoblotting using anti-alkaline phosphatase antibody against cell fractions of strains expressing FliO/alkaline phosphatase chimeras (60.3 kDa): C, whole cells; P, insoluble membrane pellet fraction; S, soluble supernatant fraction. (C) Growth of strains expressing FliO/alkaline phosphatase chimeras on L-agar plates containing 40 μg ml⁻¹ 5-Bromo-4-chloro-3-indoyl phosphate at 37°C. All strains contained a ΔphoN301 deletion. (D) Schematic overview of chimeric fusions of GFPuv within FliO. (E) Immunoblotting using anti-GFP antibody against cell fractions of strains expressing FliO/GFPuv chimeras (39.9 kDa). (F) Growth of strains expressing FliO/GFPuv chimeras on L-agar plates at 30°C. All fusions were expressed from plasmid pTrc99A-FF4 without IPTG induction.

doi:10.1371/journal.pgen.1001143.g002

not shown). FliO₁–95, FliO₁–105, and FliO₁–115, which are truncated from the C-terminus were able to restore almost full motility to CB186, so the C-terminal 30 amino acids of FliO are not essential. In comparison, FliO₁–75 and FliO₁–85 could only weakly restore motility to CB186, while FliO₁–65 could not (Figure 3C).
Figure 3. Complementation of a ΔfliO strain by FliO truncated at the N-terminus or C-terminus. (A) Schematic overview of FliO deletion mutants with the corresponding molecular weight of the protein. (B) Immunoblotting using polyclonal anti-FliO_{43-125}-6xHis antibody against 10–20% gradient SDS-PAGE separated whole cell lysates of the wild-type strain, SJW1103, containing empty pTrc99A-FF4 vector, or a ΔfliO mutant (strain CB186) containing empty vector, or strain CB186 expressing the truncated FliO proteins from plasmid pTrc99A-FF4 (without IPTG induction). (C) Complementation of strain CB186 by the truncated FliO proteins expressed from plasmid pTrc99A-FF4. SJW1103, containing empty pTrc99A-FF4 vector, was included for comparison. Soft-tryptone motility agar inoculated with the strains was incubated for 6 hours at 30°C.

doi:10.1371/journal.pgen.1001143.g003
motility of a ΔfliO mutant, suggesting that this domain is functionally important. We next undertook a site-directed mutagenesis study of the cytoplasmic domain to identify important residues to confirm it is functionally important. We aligned the FliO proteins from closely related Gammaproteobacteria, to identify conserved residues. There was considerable sequence variation, but several conserved amino acids between residues 43–95 of the cytoplasmic domain of FliO could be identified (Figure S1). To screen for these important residues, we performed site-directed mutagenesis for full-length FliO creating point substitutions for alanine or point deletions at the following positions: G64, R68, V74, G82, T84, L91, and L94. Plasmids encoding these mutations were then examined for their ability to rescue motility of the ΔfliO mutant, CB186, in soft-tryptone motility agar. FliO(L91A) was identified to cause a reduction in motility (data not shown). Immunoblotting using polyclonal anti-FliO43–125-6xHis antibodies demonstrated that FliO(L91A) and FliO(A91) could be expressed stably (Figure 4A). Strains containing the fliO(L91A) or fliO(A91) mutant alleles on the chromosome were engineered, and both strains were dramatically less motile (Figure 4B). Residue leucine 91 of the cytoplasmic domain is therefore very important for the function of FliO.

Circular dichroism (CD) spectroscopy analysis of the cytoplasmic domain of FliO

We have revealed that the cytoplasmic domain of FliO is functionally important, so we next studied this domain by circular dichroism (CD) spectroscopy to see whether it is structured. Based on secondary structure prediction the cytoplasmic domain of FliO should contain four beta-strands and one alpha-helix (Figure 5A). FliO43–125-Ala was purified using a C-terminal intein fusion tag. An additional alanine residue at the C-terminus was added to facilitate removal of the intein-tag. After cleavage of the intein fusion tag and purification FliO43–125-Ala was studied using CD spectroscopy. From the shape of the spectra, the structure of FliO43–125-Ala apparently consists mostly of random coil and some beta-structure (Figure 5B). Melting of FliO43–125-Ala was irreversible due to aggregation, this often happens with beta-structure proteins [40]. Then FliO43–125-Ala was titrated with urea and no two-state transition was observed. From these data we conclude that this peptide is disordered and its structure is a mixture of random coil and beta-sheet.

Full-length FliO increases FliP expression

We have shown that bypass mutations in fliP can rescue motility for cells deleted for the fliO gene. We have also shown through mutagenesis that the most important residues of FliO are between amino acids 22 to 95, and the cytoplasmic domain alone has functionality. To further characterize the apparent regulation of FliP by FliO, we investigated the effect of FliO co-expression on the synthesis of plasmid expressed FLAG-tagged FliP in the ΔfliO–fliP strain CB184. A FLAG-tag epitope (N–DYKDDDDK–C) was encoded between codons 22 and 23 of fliP. The FLAG-tag was added to enable detection of FliP expression by immunoblotting with ANTI-FLAG antibody. After cleavage of the N-terminal signal peptide of FliP, which occurs between residues 21 and 22, the FLAG-tag would be encoded immediately after the glutamine residue. FLAG-tagged FliP was expressed alone, or co-expressed with full-length FliO, or co-expressed with the cytoplasmic domain of FliO. FLAG-tagged FliP expression levels were not improved if the R143H or F190L bypass mutations were encoded (data not shown). Expression of full-length FliO or the cytoplasmic domain FliO43–125, was detected from the co-expression vectors by immunoblotting with polyclonal anti-FliO43–125-6xHis antibody against 10–20% gradient SDS-PAGE separated whole cell lysates of a ΔfliO mutant (strain CB186) containing empty vector, or strain CB186 expressing FliO or mutated FliO proteins (approximately 13 kDa) from plasmid pTrc99A–FF4 without IPTG induction. (B) Incubation of the wild-type strain SJW1103 or genetically engineered mutants in soft-tryptone motility agar for 6 hours at 30°C.
expression levels of FLAG-tagged FliP were not improved when the cytoplasmic domain of FliO was co-expressed (Figure 6B).

**Overexpression of the cytoplasmic domain of FliO in strains deleted for fliO and expressing bypass mutations in fliP further rescues motility**

We have revealed that full-length FliO can stabilize FliP expression, while the cytoplasmic domain of FliO cannot. We have also shown that overexpression of the cytoplasmic domain of FliO, FliO_{43–125}, can rescue motility of cells with the fliO gene deleted. We have further shown that extragenic bypass mutations within the fliP gene can also partially restore motility to the ΔfliO mutant. We next demonstrated that expression of FliO_{43–125} from pTrc99A-FF4 could further rescue motility to near wild-type levels for engineered strains encoding the ΔfliO fliP (R143H) or ΔfliO fliP (F190L) mutations (Figure 7).

**Discussion**

In this study, we have undertaken a detailed molecular analysis of the FliO secretion apparatus protein and its role in flagella assembly. FliO was previously considered to be necessary for secretion in *Salmonella* [7]. However, we have demonstrated using an engineered non-polar ΔfliO mutant that while deletion of the fliO gene leads to a dramatic reduction in motility, cells are not completely non-motile. Moreover, it was possible to readily isolate pseudorevertants containing bypass mutations in fliP, which help to rescue motility. The FliO protein is not conserved in all type III secretion systems, yet in this study we have shown that it has an important functional role in regulating FliP stability, which is a highly conserved member of the secretion system membrane proteins.

We have also shown that FliO has a predominant C-terminal cytoplasmic domain, which is in contrast to the previously predicted membrane topology of FliO, which suggested that the C-terminus is in the periplasm [32]. Coexpression of the cytoplasmic domain of FliO with FlIP did not increase FlIP expression levels, while coexpression of full-length FliO with FlIP improved FlIP expression levels. This suggests that the transmembrane region of FliO stabilizes FlIP, since the periplasmic region of FliO appears to be non-essential from our truncation analyses. However, we have shown that the cytoplasmic domain represents a functional unit beyond merely being required for FliO membrane insertion, since overexpression of the cytoplasmic domain only of FliO could improve motility of cells encoding a fliO gene deletion, with or without additional bypass mutations in fliP.

In *Buchnera* sp. APS a gene fusion of fliO and fliP exists [41]. This suggests along with the results presented here that FliO and FlIP probably also interact in *Salmonella*. This is important because FlIP has been demonstrated to be located within the flagellar basal hook and neck region, a process that is essential for the function of type III secretion systems. We have shown that FliO is necessary for FlIP localization, suggesting that FliO and FlIP may interact directly. This interaction may be critical for the proper assembly and function of the flagellar basal hook and neck region.

**Figure 6. FliO increases expression of FliP.** FLAG-tagged FlIP (25.4 kDa) was expressed alone, or co-expressed with FliO (13.1 kDa), or co-expressed with the cytoplasmic domain of FliO, FliO_{43–125} (8.8 kDa), from plasmid pTrc99A-FF4 without IPTG induction. Whole cell lysates were separated by 10–20% gradient SDS-PAGE, prior to immunoblotting. (A) Immunoblotting with polyclonal anti-FliO_{43–125}-6xHis antibody. (B) Immunoblotting with ANTI-FLAG antibody. Results are representative of the experiment performed in triplicate. doi:10.1371/journal.pgen.1001143.g006

**Figure 7. Rescue of motility by overexpression of the cytoplasmic domain of FliO.** SJW1103 is a wild-type strain. Strains genetically engineered from SJW1103 contained a fliO gene deletion with or without bypass mutations in the gene for fliP. The cytoplasmic domain of FliO, FliO_{43–125}, was expressed from plasmid pTrc99A-FF4 without IPTG induction. Soft-tryptone motility agar inoculated with the strains was incubated for 6 hours at 30°C. doi:10.1371/journal.pgen.1001143.g007
body [4]. FliP is predicted to contain 4 transmembrane loops with the 2nd and 3rd loops connected by a large (approximately 80-residue) periplasmic domain (Figure S2). The nature of the bypass mutations described here in the primary sequence of FliP, appear to be remarkably subtle. The bypass mutation F190L is located in predicted transmembrane loop 3, and was physiologically relevant by itself at improving motility in a fliO deletion background. However, the F190L mutation did not improve levels of FliP suggesting that the mutation is a gain-of-function mutation that overcomes the supporting role of FliO through a different mechanism. The FliP(R143H) mutation alone was not sufficient to greatly improve motility, rather it somehow increases the spectrum of bypass mutations permissible for rescuing motility of a fliO deletion mutant. Since arginine 143 is located in the periplasm, mutations might occur in other proteins, which interact with this domain. We are presently mapping the additional bypass mutation in strain CB191, which exists together with the fliP::R143H bypass mutation. FliP(R143H) and FliP(F190L) do not correspond to residues found at the same position for the A. aeolicus FliP orthologue or the Salmonella serovar Typhimurium FliP needle paralogs, SsaR and SpaP, which are all found in type III secretion systems without a FliO homologue (Figure S2). So these FliP homologues might consist of amino acid residues that increase their stability, or they are part of secretion systems, which do not require the supporting role(s) that FliO plays in the Salmonella system.

The most highly conserved part of the FliO protein is between residues 22 to 95. This is consistent with the results of the truncation analysis, which showed that the periplasmic domain and C-terminal 30 amino acid residues are non-essential. However, mutating residue leucine 91 of the cytoplasmic domain severely disrupted FliO function. Furthermore, overexpression of the cytoplasmic domain of FliO can partially rescue motility of a ΔfliO mutant, demonstrating functionality of this domain. Presumably, overexpression of the cytoplasmic domain of FliO overcomes the localization defect of not producing the periplasmic domain, and so overexpression enables the cytoplasmic domain to find its interaction partner(s). However, it is intriguing to know how this domain can function without being anchored to the membrane, since this suggests it might function as bridging domain to maintain protein complex stability, rather than having a catalytic role in protein translocation? Using circular dichroism spectroscopy we showed that the structure of the cytoplasmic domain FliO43–125 is a mixture of beta-sheet and random coil. We assume that the FliO cytoplasmic domain becomes structured while interacting with its binding partners similar to binding domains of many other proteins that acquire tertiary structure upon binding to their partners, such as in flagellin/flagellin or tropomodulin/tropomyosin interactions [42,43].

### Materials and Methods

#### Bacterial strains and plasmids

The bacterial strains and plasmids used in this study are listed (Table S3 and Table S4, respectively). Soft-tryptone motility agar, 0.35% (w/v), was used in motility assays [44]. Motility agar plates were maintained at 30°C, and inoculated from colonies from fresh overnight transformations. Ampicillin was used in media at 100 μg ml⁻¹ for Salmonella strains and 50 μg ml⁻¹ for E. coli strains. Kanamycin was used at 50 μg ml⁻¹, tetracycline at 15 μg ml⁻¹, chloramphenicol at 34 μg ml⁻¹, and 5-Bromo-4-chloro-3-indolyl phosphate at 40 μg ml⁻¹, where applicable. All chemicals were obtained from Sigma-Aldrich or Wako, Japan.

#### Genetic engineering procedures

The oligonucleotides used in the strain and plasmid constructions are listed (Table S5 and Table S6, respectively). To construct chromosomal gene deletions and replacements λ-Red-based recombination was used employing plasmid pKD46 [45,46]. To construct chromosomal gene deletions a kanamycin-resistance cassette was obtained from plasmid pKD13 by PCR, flanked with approximately 40-bp ends homologous to the target site. After chromosomal integration the kanamycin resistance cassette was excised using plasmid pCP20 so that the ‘scar’, which remains afterwards within the target gene(s) was encoded as an in-frame short polypeptide for the non-polar deletions, or the ‘scar’ was in the reverse orientation for the ΔfliO::391 allele. The ΔfliO::22252 deletion encodes the first 5 residues, and the final 8 residues of FliO, linked by 27-amino acid residues from an internal ‘scar’ sequence within the fliO gene. The ΔfliO::fliP::22251 deletion encodes the first 5 residues of FliO and the last 5 residues of FliP, joined by 28 residues encoded by a scar sequence.

To construct chromosomal gene replacements a tetRA tetracycline-resistance cassette was obtained by PCR from SGS3718 genomic DNA flanked by approximately 40-bp ends homologous to the target site. After chromosomal integration, it was possible to counter-select against the tetracycline-resistance cassette on medium containing fusaric acid. Then the targeted region of the chromosome was replaced using PCR-amplified DNA containing homologous-ends. To create fliO::phaA or fliO::gfpuv chimeric gene fusions in plasmid pTSO17, which carries the fliO gene, it was first linearized at the desired point by non-strand displacing PCR. Then either a phaA gene PCR product obtained from E. coli K-12 MG1655 genomic DNA or a Gfpuv gene PCR product obtained from plasmid pGFPuv (Clontech), which was flanked by 15-bp fliO homologous ends, was inserted by the homologous recombination-based In-fusion PCR cloning procedure (Clontech). Site-directed mutagenesis was performed using the QuickChange Lightning Site-Directed Mutagenesis Kit (Stratagene).

### Purification of FliO43–125-Ala and FliO43–125-6xHis

FliO43–125-Ala was purified using a C-terminal intein tag encoded by a pTXB1-based expression plasmid, according to manufacturer’s instructions (New England Biolabs). BL21 Star containing plasmid pTSO133 was cultivated. The cell pellet from 5-L culture was suspended in 200-ml buffer consisting of 40 mM Tris-HCl (pH 8.0), 1 mM NaCl, 2 mM EDTA, and 20% glycerol, and lysed by sonication. Insoluble proteins and cell debris were collected by low-speed centrifugation and suspended in 100-ml buffer containing 40 mM Tris-HCl (pH 8.0), 0.5 mM NaCl, and 8 mM urea. The suspension was incubated with stirring at room temperature for 3 hours. Insoluble material was removed by centrifugation. After centrifugation, the supernatant was diluted four times with washing buffer consisting of 40 mM Tris-HCl (pH 8.0), 0.5 mM NaCl, and 1 mM urea, and loaded onto a chitin bead column equilibrated with washing buffer (the column volume was 30-ml). The column was washed with 200-ml of washing buffer, and then with 100-ml of washing buffer containing 0.1 M DTT. After that the column was incubated at room temperature for about 20 hours. FliO43–125-Ala was eluted from the chitin column with 50-ml of washing buffer and further purified by gel-filtration on a HiLoad Superdex-75 gel filtration column (GE healthcare) equilibrated with the same buffer. Fractions containing pure FliO43–125-Ala were pooled together, and the protein was transferred by dialysis in 20 mM Na/K phosphate (pH 6.2), and 100 mM NaCl. FliO43–125-6xHis was purified according to standard procedures, using nickel affinity chromatography under denaturing conditions after expression from plasmid pES0221.
**Immunoblotting**

Immunoblotting was performed similar to a previously described method [25]. For immunoblotting of whole cell lysates, colonies of *Salmonella* serovar Typhimurium containing the desired plasmid, from an overnight transformation, were inoculated into 5-ml LB with antibiotic and grown with shaking for 6 hours at 37°C. Cells were harvested by centrifugation at 16,100-×g for 5 minutes, and the pellets were re-suspended in SDS-PAGE loading buffer (100 mM Tris-HCl, pH 6.8, 2% SDS, 1 mM β-mercaptoethanol, 7 M urea, and 0.1% bromophenol blue) and incubated at 95°C for 15 minutes. An equivalent amount of each whole cell lysate was separated by SDS PAGE and proteins were either detected by staining with Coomassie brilliant blue or transferred to PVDF membranes. Immunoblotting was performed using the anti-rabbit WesternBreeze Chromogenic Kit (Invitrogen), according to manufacturer’s instructions. The following antibodies and dilutions were used: anti-alkaline phosphatase (Rockland, PA, USA), 1:20,000 dilution; anti-GFP (Clontech), 1:5,000 dilution; ANTI-FLAG (Sigma-Aldrich), 1:5,000 dilution; and polyclonal anti-FliO 43–125-6xHis antibody, 1:10,000 dilution.

**Cell fractionation**

Cells were separated into the insoluble membrane pellet fraction and soluble supernatant fraction after lysis similar to a previously described method [47]. Briefly, from overnight transformations, colonies of *Salmonella* serovar Typhimurium containing the desired plasmid were inoculated into 50-ml LB with antibiotic and grown with shaking for 7 hours at 37°C. Cells were harvested by centrifugation at 6000-×g for 10 minutes, and the pellets were re-suspended in 11-ml 50 mM Tris-HCl (pH 8.0), 10 mM MgCl₂, and 440-µl 25× EDTA-free protease inhibitor cocktail (Roche). Cell pellets were lysed by sonication at 25% power for 5 cycles of one minute each. After incubation at room temperature for 30 minutes unlysed cell debris was removed by low-speed centrifugation at 10,000-×g for 10 minutes. The insoluble membrane pellet was obtained by high-speed centrifugation at 100,000-×g for 1 hour and resuspended in 1-ml 50 mM Tris-HCl (pH 8.0), and 10 mM MgCl₂. One-µl was added to 5-µl SDS-PAGE loading buffer for electrophoresis. The supernatant was further clarified by centrifugation at 100,000-×g for 1 hour and 10-µl was added to 5-µl SDS-PAGE loading buffer for electrophoresis.

**Observation of GFPuv fluorescence and measurement of alkaline phosphatase activity**

Expression of FliO/GFPuv chimeras was detected for strains inoculated onto L-agar plates containing the appropriate antibiotic, after 4 days incubation at 30°C, using a transilluminator set at 365 nm UV. Alkaline phosphatase assays are detailed in Text S1.

**Circular Dichroism (CD) Spectroscopy**

CD spectra of FliO 43–125-Ala was measured using an Aviv model 400 spectropolarimeter (Lakewood, NJ) in 0.1 cm cuvettes at 0°C in 20 mM Na/K phosphate, pH 6.2, 100 mM NaCl. Urea titration ellipticity at 220 nm was measured for samples in 0, 1, 2, 3, 4, 5, 6, 7 and 8 M urea at 10°C.

**Bioinformatics**

Protein secondary structure prediction was performed using Jpred 3 [48]. Programs used to align protein sequences, and programs used to predict membrane protein-membrane topology are detailed in Text S1.

**Supporting Information**

**Figure S1** Alignment and predicted membrane topologies of FliO proteins from *Gammaproteobacteria*. SALTY = *Salmonella enterica* serovar Typhimurium; ECOLI = *Escherichia coli* K-12; ERWTA = *Erwinia tasmaniensis*; YEREN = *Yersinia enterocolitica*; XANCA = *Xanthomonas campestris*; VIBCH = *Vibrio cholerae*; and PSEAE = *Pseudomonas aeruginosa*. We have shown for *Salmonella* serovar Typhimurium FliO that the underlined residues in bold type could be truncated from the N-terminus or the C-terminus, and FliO still remained partially functional. We also identified residue leucine 91 (black bold-type) is important for the function of full-length *Salmonella* serovar Typhimurium FliO. The percentage score of the FliO homologues with *Salmonella* serovar Typhimurium FliO is indicated. Found at: doi:10.1371/journal.pgen.1001143.s001 (0.03 MB PDF)

**Figure S2** Alignment and predicted membrane topologies of FliP, and FliP homologues. *Salmonella enterica* serovar Typhimurium FliP, the *Aquifex aeolicus* FliP orthologue, and the *Salmonella* serovar Typhimurium FliP paralogues, SpaP and SsaR were aligned. SALTY = *Salmonella* serovar Typhimurium, and AQUAE = *A. aeolicus*. A. aeolicus FliP, and SpaP and SsaR are found in Type III secretion systems without a FliO homologue. In this study, it was shown that bypass mutations in *Salmonella* serovar Typhimurium FliP corresponding to R143H and F190L, could partially rescue motility of a fliO deletion mutant. Residues aligned with arginine 143 and phenylalanine 190 of *Salmonella* serovar Typhimurium FliP are indicated in dark red bold type. The percentage score of the homologues with *Salmonella* serovar Typhimurium FliP is indicated. Found at: doi:10.1371/journal.pgen.1001143.s002 (0.03 MB PDF)

**Table S1** Prediction of FliO transmembrane topology. Found at: doi:10.1371/journal.pgen.1001143.s003 (0.05 MB DOC)

**Table S2** Alkaline phosphatase activity of *Salmonella enterica* serovar Typhimurium expressing chimeric fusions of alkaline phosphatase within FliO. Found at: doi:10.1371/journal.pgen.1001143.s004 (0.03 MB DOC)

**Table S3** Strains used in this study. Found at: doi:10.1371/journal.pgen.1001143.s005 (0.06 MB DOC)

**Table S4** Plasmids used in this study. Found at: doi:10.1371/journal.pgen.1001143.s006 (0.06 MB DOC)

**Table S5** Oligonucleotides used in strain constructions. Found at: doi:10.1371/journal.pgen.1001143.s007 (0.06 MB DOC)

**Table S6** Oligonucleotides used in plasmid constructions. Found at: doi:10.1371/journal.pgen.1001143.s008 (0.12 MB DOC)

**Text S1** Supplementary materials and methods. Found at: doi:10.1371/journal.pgen.1001143.s009 (0.03 MB DOC)

**Acknowledgments**

We would like to thank Vladimir Meshcheryakov (Okinawa Institute of Science and Technology) for purifying FliO 43–125-Ala. We are very grateful to FBS Osaka University, *Salmonella* Genetic Stock Center (University of
Calgary, *Escherichia coli* Genetic Stock Center (Yale University), and National Institute of Genetics (Japan) for generously providing us with plasmids and strains.

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Author Contributions

Conceived and designed the experiments: CSB ASK FAS. Performed the experiments: CSB IVM ASK FAS. Analyzed the data: CSB IVM ASK FAS. Wrote the paper: CSB FAS.