

# Recognition and targeting mechanisms by chaperones in flagellum assembly and operation

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Edited by G. Marius Clore, National Institutes of Health, Bethesda, MD, and approved July 18, 2016 (received for review May 16, 2016)

The flagellum is a complex bacterial nanomachine that requires the proper assembly of several different proteins for its function. Dedicated chaperones are central in preventing aggregation or undesired interactions of flagellar proteins, including their targeting to the export gate. FliT is a key flagellar chaperone that binds to several flagellar proteins in the cytoplasm, including its cognate filament-capping protein FliD. We have determined the solution structure of the FliT chaperone in the free state and in complex with FliD and the flagellar ATPase FliI. FliT adopts a four-helix bundle and uses a hydrophobic surface formed by the first three helices to recognize its substrate proteins. We show that the fourth helix constitutes the binding site for FlhA, a membrane protein at the export gate. In the absence of a substrate protein FliT adopts an autoinhibited structure wherein both the binding sites for substrates and FlhA are occluded. Substrate binding to FliT activates the complex for FlhA binding and thus targeting of the chaperone-substrate complex to the export gate. The activation and targeting mechanisms reported for FliT appear to be shared among the other flagellar chaperones.

flagellum | chaperones | assembly factors | NMR spectroscopy | type III secretion

he flagellum is the organelle that enables bacterial locomotion and is one of the most sophisticated protein machines (1-4). Flagella organelles may act as virulence factors because motility is crucial for the action of pathogenic bacteria (5, 6). About 25 different proteins are involved in the assembly of the flagellum, which is divided into five parts: the basal body, hook, hook-filament junction, filament, and filament cap. The basal body is a flagellum-specific type III secretion system responsible for the translocation of flagellar proteins to the distal end of the growing flagellar structure for self-assembly (7). The assembly of the hook and the filament is strictly sequential and is controlled by the export apparatus, which uses both ATP and proton motive force (PMF) to drive protein translocation (8, 9). The cytoplasmic domain of FlhA (10-12), one of the six membrane proteins of the flagellar type III export apparatus, acts as an adaptor to receive the flagella building blocks FliD and flagellin (FliC) when bound to their cognate chaperones FliT and FliS, respectively (Fig. 1) (11).

The export apparatus also comprises three cytosolic proteins: FliI, FliH, and FliJ (Fig. 1). FliI (13), the only ATPase of the system, assembles to a hexameric ring (14, 15) and is located at the base of the basal body (16). The exact role of FliI is not known, but it has been implicated in powering protein export, targeting flagellar proteins to the export gate, and functioning as a sorting platform or specificity switch (17). FliI adopts a structure that is very similar to the structure of the F<sub>1</sub> ATP synthase (13). The first 20 N-terminal residues of FliI mediate self-oligomerization (18) and are involved in binding FliH (19, 20). FliH is homologous to the  $\beta$  and  $\delta$  subunits of the F<sub>0</sub>F<sub>1</sub> ATP synthase (21) and is thought to provide the support point for the association of the FliI ring to the export gate (22). FliJ, which is essential for the export of flagellum building blocks, binds to the center of the hexametric FliI ring in a manner similar to the way the  $\gamma$  subunit binds to the  $\beta$  subunit of  $F_0F_1$  ATP synthase (23). FliJ also was proposed to recycle substrate-free flagellar chaperones at the export gate (24). FliI, FliH, and FliJ bind to FlhA and FlhB proteins at the export gate (11, 25, 26).

Chaperones dedicated to the assembly and operation of the flagellum bind and protect their cognate substrates from aggregation or premature interactions in the cytoplasm (27–29). In addition, chaperones have been implicated in targeting flagellar proteins to the export gate (11, 30, 31). FliT is a key chaperone in the assembly and operation of the flagellum. FliT interacts with several flagellar proteins, including the exported substrate FliD and the export apparatus components FliI, FliJ, and FlhA, and negatively regulates transcription of flagellar genes by inhibiting the formation of a DNA complex with the master regulator FlhDC (11, 24, 28, 32–36). FliD is exported first during the filament assembly to form a pentameric cap that promotes self-assembly of FliC (37).

The structural details of how FliT recognizes its various partners and how it targets and escorts them to the export gate are not known. We have determined the structures of FliT in the unliganded state and in complex with its substrates FliD and FliI. The structural data show that FliT adopts an autoinhibited conformation in which both the substrate- and FlhA-binding sites are occluded. Substrate binding to FliT releases the FlhAbinding site, thereby enabling targeting of the FliT–substrate complex to the export gate.

### **Significance**

The flagellum is a highly sophisticated organelle rotated by a motor that confers swarming motility to bacterial cells. Such motility is essential for the full pathogenicity of several virulence bacteria. Several proteins are required for the assembly and operation of the flagellum. Here we report the structural characterization of FliT, a key flagellar chaperone, in the unliganded state and in complex with two substrate flagellar proteins. FliT adopts an autoinhibited structure in order to avoid futile interactions with the export gate in the absence of a substrate. Substrate binding to FliT activates complex targeting to the export gate followed by either the export of the substrate or its assembly to the export apparatus.

Author contributions: N.K., P.R., and C.G.K. designed research; N.K. and P.R. performed research; A.E. contributed new reagents/analytic tools; N.K., P.R., and C.G.K. analyzed data; and N.K., P.R., and C.G.K. wrote the paper.

The authors declare no conflict of interest.

This article is a PNAS Direct Submission

Data deposition: The atomic coordinates reported in this paper have been deposited in the Protein Data Bank, www.wwpdb.org (PDB ID codes 5KS6, 5KRW, and 5KP0). The NMR chemical shifts have been deposited in the Biological Magnetic Resonance Data Bank, www.bmrb.wisc.edu (accession nos. 30136, 30134, and 30127).

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This article contains supporting information online at www.pnas.org/lookup/suppl/doi:10. 1073/pnas.1607845113/-/DCSupplemental.

www.pnas.org/cgi/doi/10.1073/pnas.1607845113



Fig. 1. A simplified schematic of the flagellum that includes the proteins studied in this work. For a detailed view, see refs. 1–4.

### Results

Solution Structure of the FliT Chaperone. Salmonella enterica serovar Typhimurium FliT consists of 122 residues (~13.7 kDa), and multiangle light scattering (MALS) analysis showed that it is a monomer in solution (Fig. S1A). At high concentrations FliT has a tendency to form a weak dimer with a dimerization dissociation constant ( $K_d$ ) of ~0.5–0.6 mM, in agreement with previously published analytical ultracentrifugation data (35). We used NMR (Fig. S1 B and C) to determine the structure of FliT in solution under conditions in which FliT is monomeric ([FliT] ~0.08 mM). The structure shows that FliT adopts a four-helix bundle (Fig. 2A and Fig. S2A). The first three helices ( $\alpha 1-\alpha 3$ , residues 2-27, 30-49, and 58-83, respectively) are longer than the fourth ( $\alpha$ 4) helix (residues 98–112). Helices  $\alpha$ 3 and  $\alpha$ 4 are connected by a long unstructured linker. Helices  $\alpha 1$ ,  $\alpha 2$ , and  $\alpha 3$ interact intimately with each other, whereas helix  $\alpha 4$  interacts primarily with helix  $\alpha 3$  (Fig. 2 *B* and *C*). The positioning of helix  $\alpha$ 4 with regard to the other helices appears to be suboptimal, and the FliT structure departs from the canonical four-helix bundle. Helices  $\alpha 1$ ,  $\alpha 2$ , and  $\alpha 3$  expose a contiguous hydrophobic surface (~1,500 Å<sup>2</sup>), and helix  $\alpha$ 4 acts as a lid to occlude it from solvent (Fig. 2D). Leu102, which docks into a hydrophobic pocket lined by nonpolar residues from the other three helices (Fig. 2 B and C), seems to be crucial in stabilizing the helix  $\alpha$ 4 position against the core of the bundle. A FliT variant in which Leu102 was replaced by Ala is prone to aggregation, and NMR spectra experience severe line broadening, especially at the interface with helix  $\alpha 4$  (Fig. S1D). Apparently, mutation of Leu102 destabilizes the docking of helix  $\alpha 4$ , and as a result the hydrophobic core is exposed to the solvent. A FliT variant lacking helix  $\alpha 4$  (FliT<sup> $\Delta \alpha 4$ </sup>) has a strong tendency to aggregate, further highlighting the role of helix  $\alpha 4$  in shielding the hydrophobic core of FliT.

The FliT structure was previously determined by X-ray crystallography to be a tetramer (35). The structures in solution and in the crystal have pronounced differences. Compared with the solution structure, helix  $\alpha$ 3 in the crystal structure extends by an additional three turns, and helix  $\alpha$ 4 is detached and interacts instead with the hydrophobic core of another FliT subunit. The differences between the solution and crystal structures appear to be the result of crystal-packing effects. Interaction Between FliT and FliD. FliT is the chaperone for the filament-capping protein FliD, but the structural basis for their interaction is not known. To understand how FliT interacts with FliD, we used NMR spectroscopy to characterize the complex. We prepared several FliD fragments and identified the last 40 residues of the C terminus as the FliT-binding site, in agreement with previous biochemical experiments (28, 32). To identify the FliD-binding site on FliT, we titrated FliD<sup>C</sup> (the FliD fragment encompassing the last 40 C-terminal residues) to isotopically labeled FliT and monitored their interaction by NMR (Fig. S3A). Chemical shift analysis shows that the FliT residues most affected by FliD binding are located in the hydrophobic core formed by helices  $\alpha 1 - \alpha 3$  (Fig. S3B). Because the hydrophobic core in free FliT is occluded by helix  $\alpha 4$  (Fig. 2D), FliD binding would entail displacement of helix  $\alpha 4$ . Indeed, NMR analysis shows that the helix  $\alpha 4$  residues experience large chemical shift perturbation, and linewidth measurements suggest that helix  $\alpha 4$ is more mobile when FliD is bound, indicating displacement of helix  $\alpha 4$  (Fig. S3B). Taken together, the NMR data show that FliD binds to the hydrophobic surface formed by helices  $\alpha 1 - \alpha 3$ in FliT, and as a result helix  $\alpha 4$  is displaced.

**Solution Structure of the FliT–FliD Complex.** To understand how FliT recognizes and binds to FliD, we sought to determine the structure of the FliT–FliD complex. The NMR spectra of FliT–FliD show significant line broadening at the interface of the complex, most likely resulting from unfavorable kinetics of complex formation and dissociation. Several different constructs were designed and prepared, and their NMR properties were tested. The best NMR spectra suitable for structure determination by NMR were provided by a construct consisting of the last 40 C-terminal residues of FliD (FliD<sup>C</sup>) fused to a FliT variant



**Fig. 2.** Solution structure of the FIIT chaperone. (*A*) FIIT is shown in a cartoon rendering and is colored using a continuous-gradient color scheme from the N terminus (blue) to the C terminus (red). The four helices ( $\alpha 1-\alpha 4$ ) are labeled. (*B* and *C*) FIIT is shown in gray cartoon with semitransparent representation of the solvent-accessible surface in two different views. The residues at the interface of the four-helix bundle are shown in ball-and-stick configuration. (*D*) The hydrophobic surface of FIIT is colored green. Helix  $\alpha 4$  acts as a lid to occlude the large hydrophobic surface formed by helices  $\alpha 1-\alpha 3$  from solvent.

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**Fig. 3.** Solution structure of the  $\text{FliT}^{\Delta\alpha4}$ - $\text{FliD}^{C}$  complex.  $\text{FliT}^{\Delta\alpha4}$  is colored green and  $\text{FliD}^{C}$  is colored orange. The solvent-exposed surface of FliT is represented in semitransparent light gray. Residues are shown in ball-and-stick configuration. The dashed lines denote hydrogen bonds. Two views are shown related by a 90° rotation about the *x* axis.

lacking the terminal helix  $\alpha 4$  (FliT<sup> $\Delta \alpha 4$ </sup>) (Fig. S4 *A*–*C*). Fusion proteins increase the effective concentration, and thus the population, of the complex and at the same time shift the kinetics of complex formation toward the slow-exchange regime. This strategy has been used extensively to determine structures of protein complexes by NMR (38). NMR analysis and comparison of the spectra of the FliT<sup> $\Delta \alpha 4$ </sup>–FliD<sup>C</sup> fusion construct with the spectra of FliT–FliD<sup>C</sup> show that the molecular interface and the structural properties are essentially identical in the two complexes (Fig. S5). The solution structure of the FliT<sup> $\Delta \alpha 4$ </sup>–FliD<sup>C</sup> complex was determined by NMR on the basis of a large number of intermolecular NOEs. The structure and NMR statistics are summarized in Table S1.

FliD<sup>C</sup> binds to FliT<sup>Δα4</sup> by forming a long amphipathic helix running from residue Ser428 through Ser467 (Fig. 3 and Fig. S2B), which juxtaposes with the three-helix bundle ( $\alpha$ 1– $\alpha$ 3) of FliT. A continuous string of hydrophobic residues, notably Tyr436, Leu443, Met446, Leu450, Thr453, Tyr456, Leu457, Phe461, and Met464, run along the 37-Å–long FliD helix. Key contacts are between FliT residues Trp11 and Tyr40 and between FliD residues Leu457 and Leu450, respectively. The only polar contacts appear to be hydrogen bonds between FliT Lys79 and FliD Ser454 and between FliT Glu45 and FliD Lys449. The face of the FliD<sup>C</sup> helix that is exposed to the solvent is decorated almost exclusively with polar amino acids. Mutation of residues at the interface between FliT and FliD<sup>C</sup> has a strong effect on the stability of the complex (Fig. S64).

FliD binding elicits drastic conformational changes in FliT. Superposition of the apo FliT structure with the structure of the FliT–FliD<sup>C</sup> complex reveals that the C-terminal FliD helix (residues Asn451–Ser467) occupies the same space as helix  $\alpha 4$  in the free FliT (Fig. 44). Thus, the structure explains why FliD binding entails displacement of FliT helix  $\alpha 4$ . In addition, the long FliD<sup>C</sup> helix pairs with FliT helix  $\alpha$ 3, resulting in the elongation of the latter by three turns of a helix. As a result, FliT–FliD<sup>C</sup> complex formation buries a total of ~2,500 Å<sup>2</sup>, with the majority of the surface (~2,060 Å<sup>2</sup>) consisting of nonpolar atoms and only ~440 Å<sup>2</sup> consisting of polar atoms.

Of note, the structural data show that FliT adopts an autoinhibited conformation in its free state. Indeed, isothermal titration calorimetry (ITC) shows that FliD binds with a 10-fold higher affinity to FliT<sup> $\Delta \alpha 4$ </sup> than to FliT (Fig. S6 *C* and *D*), confirming that helix  $\alpha 4$  acts as an autoinhibitory element in FliT.

Targeting of FliT-FliD to the Export Gate. FliT was shown by biochemical assays to deliver FliD to the export gate by interacting with the cytoplasmic domain of FlhA (FlhA<sup>C</sup>) (11, 36). To understand the targeting process, we characterized the interaction of FliT and FliD with FlhA<sup>C</sup>. NMR shows that free FliT does not bind to FlhA<sup>C</sup> (Fig. S7), in agreement with surface plasmon resonance (SPR) data (36). In contrast, the FliT-FliD<sup>C</sup> complex binds strongly to FlhA<sup>C</sup> (Fig. 4B). NMR analysis points to the FliT helix  $\alpha 4$  as the primary binding site mediating the interaction between FliT–FliD<sup>C</sup> and FlhA<sup>C</sup>. Indeed, the FliT<sup> $\Delta a4$ </sup>–FliD<sup>C</sup> variant, which lacks the FliT  $\alpha 4$  helix, does not interact with FlhA<sup>C</sup> (Fig. 4C). Taken together, the data demonstrate that free FliT does not interact with FlhA because the  $\alpha$ 4 helix, which constitutes the primary FlhA-binding site, is inaccessible (Fig. 2). FliD binding displaces the FliT  $\alpha$ 4 helix, thereby poising the FliT-FliD complex for binding to FlhA.

**Interaction Between FliT and FliI.** It has been shown previously by pull-down assays that FliT binds to the FliI ATPase (39). To



**Fig. 4.** FliD binding releases FliT helix  $\alpha$ 4 and activates the complex for FlhA binding. (A) Superposition of free FliT on the FliT–FliD<sup>C</sup> complex structure. The FliT helix  $\alpha$ 4, which our NMR data show is released upon FliD binding, is modeled in the structure. (B) <sup>1</sup>H-<sup>13</sup>C heteronuclear multiple quantum coherence (HMQC) spectra of FliT–FliD<sup>C</sup> (blue) and in the presence of FlhA<sup>C</sup> (orange). The data show that FliT–FliD<sup>C</sup> binds to FlhA<sup>C</sup> and that the interaction is mediated primarily by FliT helix  $\alpha$ 4. (C) <sup>1</sup>H-<sup>13</sup>C HMQC spectra of FliT<sup> $\Delta$ α4</sup>–FliD<sup>C</sup> (blue) and in the presence of FliA<sup>C</sup> (orange). The data show that truncation of FliT helix  $\alpha$ 4 abrogates binding to FlhA<sup>C</sup>.



**Fig. 5.** Solution structure of the  $FliT^{\Delta\alpha4}$ – $Flil^{EN}$ . Fli $T^{\Delta\alpha4}$  is colored green, and Flil<sup>EN</sup> is colored pink. The solvent-exposed surface of FliT is represented in semitransparent light gray. Residues are shown in ball-and-stick configuration. The dashed lines denote hydrogen bonds. Two views are shown related by a 90° rotation about the *x* axis.

understand how FliT interacts with FliI, we used NMR spectroscopy to characterize their complex. To identify the binding site of FliI on FliT, we added unlabeled FliI (~50 kDa) in a stepwise manner (Fig. S84) to isotopically labeled FliT and monitored their interaction by NMR (Fig. S8B). Differential broadening analysis shows that the FliT residues most affected by FliI binding are located in the hydrophobic core formed by helices  $\alpha 1-\alpha 3$  (Fig. S8C), the FliT site to which FliD binds.

To determine the FliT-binding site in FliI, we prepared several FliI constructs and tested their binding to FliT. A FliI construct lacking the extreme N-terminal region (EN, residues 1–17) (FliI<sup> $\Delta$ EN</sup>) showed no evidence of binding to FliT (Fig. S8D). Therefore the NMR data point to the FliI EN region as the

binding site for the FliT chaperone, in agreement with previously reported pull-down assays (35). The FliI EN region also mediates self-oligomerization and stimulates ATPase activity (18), thus explaining why FliT binding to FliI inhibits its ATPase activity (Fig. S8*E*). FliT binds to FliI with moderate affinity ( $K_d \sim 5 \mu$ M) as measured by ITC. As with FliD, FliI binding displaces FliT helix  $\alpha$ 4, and FliI binds with a 10-fold higher affinity to FliT<sup> $\Delta \alpha 4$ </sup> (Fig. S6 *E* and *F*).

**Solution Structure of the FliT–Flil Complex.** To determine the structure of the FliT–Flil complex, we followed an approach similar to the one described above for the FliT–FliD complex. The FliT<sup> $\Delta \alpha 4$ </sup> variant fused to the FliI EN (FliT<sup> $\Delta \alpha 4$ </sup>–FliI<sup>EN</sup>) yielded NMR spectra of high quality (Fig. S4 *D–F*). Superposition of the FliT<sup> $\Delta \alpha 4$ </sup> – FliI<sup>EN</sup> and FliT–FliI<sup>EN</sup> spectra showed that the chemical shifts are very similar (Fig. S8*F*), in agreement with the observation that helix  $\alpha 4$  does not interact with FliI<sup>EN</sup>. The solution structure of the FliT<sup> $\Delta \alpha 4$ </sup>–FliI<sup>EN</sup> complex was determined by NMR on the basis of a large number of intermolecular NOEs. The structure and NMR statistics are summarized in Table S1.

The FliT<sup> $\Delta \alpha 4$ </sup>-FliI<sup>EN</sup> complex (Fig. 5 and Fig. S2C) forms a four-helix bundle, with FliI<sup>EN</sup> adopting an amphipathic helix extending from residues Met1 to Val25 that juxtaposes with the three-helix bundle ( $\alpha 1-\alpha 3$ ) of the FliT chaperone. Complex formation buries a total of ~2,230 Å<sup>2</sup>. The majority of the surface (~1,820 Å<sup>2</sup>) consists of nonpolar atoms, and ~410 Å<sup>2</sup> consists of polar atoms. Two notable salt bridges are formed between Lys79 of FliT and Asp13 and Glu16 of FliI and between Arg4 of FliI and Glu37 of FliT. Several bulky nonpolar residues (Met1, Leu5, Trp8, Leu12, Phe15, and Met19) run along the interacting face of the FliI EN helix and insert into pockets lined by nonpolar residues in FliT. Residues Trp8 and Phe15 of FliI make key hydrophobic contacts with Tyr40 and Trp11, respectively, of FliT. Mutation of residues at the interface between FliT and FliI has a strong effect on the stability of the complex (Fig. S6B).

FliT Uses the Same Binding Site to Interact with Its Substrates. Comparison of the structures of the FliT–FliI and FliT–FliD complexes demonstrates that FliD and FliI interact with essentially the same surface of FliT (Fig. S94). To determine the site that FliT uses to interact with FliJ, we monitored the binding of unlabeled FliJ to isotopically labeled FliT (Fig. S9B). Analysis of the NMR data showed that FliJ binds to the same surface of FliT to which FliD and FliI bind. Therefore, FliT uses the same binding site to recognize and interact with its three partner proteins: FliD, FliJ, and FliI. Binding of any of the partners displaces the FliT  $\alpha$ 4 helix, and thus FliT may target these three



Fig. 6. Structures of the three flagellar chaperones: FliT (this work) (A), FlgN (PDB ID code 2FUP) (B), and FliS (40) (C). The highly conserved Tyr residues that are crucial for the interaction of the substrate-bound chaperones with FlhA are labeled. The FlgN structure was resolved only up to residue Arg106 (Salmonella numbering), and the remaining C-terminal region was modeled in the structure.



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Fig. 7. A cartoon showing how FliT may assist in the assembly of the cytoplasmic components by binding and targeting FliJ (i), Flil (ii), and FliD (iii) to the export gate (iv). See text for details.

substrates to the export gate by its direct interaction with FlhA (Fig. 4*B*).

# Discussion

Here we present the solution structures of the FliT chaperone and its complexes with the cognate substrate FliD and the FliI ATPase. FliT is a key protein in the assembly and operation of the flagellum because it binds to several flagellar proteins in the cvtoplasm (FliD, FliJ, and FliI) and ushers them to the export gate by interacting with the FlhA membrane protein. We show that FliT adopts an autoinhibited structure in which both the binding site for the partner proteins (the hydrophobic surface formed by helices  $\alpha$ 1- $\alpha$ 3) and the binding site for FlhA (helix  $\alpha$ 4) are occluded. The autoinhibited structure of FliT may serve two goals: First, it occludes from the solvent, and thus protects, the hydrophobic substate-binding surface in the absence of substrates. Indeed, our data show that FliT variants in which the docking of helix  $\alpha 4$  to the hydrophobic core is compromised are unstable and prone to aggregation. Second, the autoinhibited structure serves to mask the binding site for FlhA, which is activated for binding only when the substrate protein is to be targeted to the export gate.

Interestingly, the autoinhibition/activation and targeting mechanism presented here for FliT appears to be shared among the other flagellar proteins (Fig. 6). FlgN, the chaperone dedicated to the binding and targeting of the FlgK and FlgL hook proteins (32), likely adopts an autoinhibited structure. The overall structure of *Pseudomonas aeruginosa* FlgN [Protein Data Bank (PDB) ID code 2FUP] (Fig. 6B) resembles the structure of FliT (Fig. 6A), with the FlgN C-terminal helix packing against a hydrophobic surface formed by the first three helices that likely form the substrate-binding site. Although the structure of the complex between FlgN and its substrates is not known, biochemical data showed that truncation of the terminal C helix in FlgN increases the binding of substrates (30). The C-terminally

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truncated FlgN does not interact with FlhA (31), suggesting that, as in FliT, the C-terminal helix in FlgN serves as the FlhAbinding site. In contrast to FliT, substrate-free FlgN does interact with FlhA, albeit with lower affinity than in the complex with its substrate FlgK, probably because the FlhA-binding site in FlgN is only partially occluded (Fig. 6*B*). Similarly, FliS, the chaperone for FliC, adopts an autoinhibited structure (Fig. 6*C*), with the inhibitory structural element, located in the N terminus, released upon FliC binding (40). The N terminus of FliS serves as the FlhAbinding site, and in all three chaperones a highly conserved Tyr residue appears to be essential for efficient FlhA binding (Fig. 6) (36). Thus, the three flagellar chaperones may use similar strategies for substrate binding and activation of the complexes for binding to FlhA and thus for targeting to the export gate.

We propose, based on the current results, a simplified model of how FliT may assist with the assembly and operation of the flagellum (Fig. 7). FliT binds to FliJ and ushers it to the export gate by means of the interaction between the FliT  $\alpha$ 4 helix and FlhA (Fig. 7*i*). Then, FliT transports FliI at the membrane (Fig. 7*ii*), where FliH takes over FliI from FliT (41), and FliI, with the assistance of FliJ, forms a hexamer (Fig. 7*iii*). When it is time for FliD to be exported, FliT binds and escorts FliD to the membrane for its export and the assembly of the filament-capping structure (Fig. 7*iv*). FlgN also binds to FliJ and FliI, and thus FliT and/or FlgN may escort these proteins to the export gate.

## **Materials and Methods**

The materials and methods used in this work are described in *SI Materials* and *Methods*.

ACKNOWLEDGMENTS. This work was supported by NIH Grant Al094623 (to C.G.K.) and Fonds Wetenschappelijk Onderzoek Grant T3RecS G002516N (to A.E.).

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