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Structural characterization of FliP,

a component of the type III flagellar protein export gate

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Abstract

The bacterial flagellum is a filamentous motile organelle that protrudes from the cell body. For flagellar construction, flagellar component proteins are transported via its specific export apparatus from the cytoplasm to the growing distal end of the flagellar structure. The export gate complex is composed of six inner-membrane proteins, FliO, FliP, FliQ, FliR, FlhA and FlhB and utilizes proton motive force (PMF) as an essential energy source to drive flagellar protein export. However, it remains unknown how the export gate converts PMF to the mechanical work required for protein translocation and how the export gate complex assembles.

This thesis is focused on structure and function of FliP. FliP is an essential component for flagellar protein export and is assumed to be involved in the earliest stage of the assembly of the export gate complex. FliO is assumed to have a role in the regulation of FliP during flagellar assembly although it is not an essential component. To obtain the structural insight into FliP, I solved the structure of the periplasmic domain of FliP (FliP_P) from *Thermotoga maritia* at 2.4 Å resolution. The structural feature of Tm-FliP_P and the following *in vivo* experiments suggested that FliP forms a dimer though the FliP_P-FliP_P interaction. Structure-based mutational analyses of FliP_P indicated that the FliP_P-FliP_P interaction is required for efficient FliP assembly into the export gate.

To clarify that FliP forms oligomer, I purified full-length FliP derived from *Salmonella enterica* and observed it by electron microscopy. Full-length FliP formed the homo-hexameric ring structure. Co-expression of FliP-His with FliO and the following co-purification by Ni-NTA affinity chromatography indicated that FliP directly interacts with FliO. The FliP₆ ring dissociated from the FliO/P complex during size exclusion chromatography was more mono-dispersed than FliP alone, suggesting that FliO facilitates hexagonal FliP ring formation. Expression and purification of FliP mutant variants showed that the FliP_P-FliP_P interaction is required for efficient FliP ring formation. I propose that the FliP₆ ring is a functional unit in the export gate complex.

Abbreviations

- ATP : adenosine triphosphate
- ADP : adenosine diphosphate
- DDM : n-Dodecyl- β -D-maltoside
- DNA : deoxyribonucleic acid
- DTT : dithiothreitol
- EDTA : ethylenediaminetetraacetic acid
- EM : electron microscopy
- IPTG : isopropyl-1-thio-b-D-galactopyranoside
- LB: Luria-Bertani broth
- MALDI-TOF : matrix assisted laser desorption ionization time of flight
- MPD: 2-Methyl-2,4-pentanediol
- OD₆₀₀ : optical density (wavelength 600 nm)
- PCR : plymerase chain reaction
- PMF : proton motive force
- RMSD : root mean square deviation
- SAD : single-wavelength anomalous dispersion
- S/N : signal/noise
- SDS-PAGE : sodium dodecyl sulfate polyacrylamide gel electrophoresis
- Tris : tris (hydroxymethyl) aminomethane
- TM : transmembrane

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Chapter 1 Introduction of the bacterial flagellum

1.1 Bacterial flagellum

Many bacteria can swim in liquid environments by rotating long, herical filamentous organelle called the flagellum (Berg and Anderson, 1973) (Fig.1-1a). The flagellum has a rotary motor at the base of its filament. The flagellar motor is driven by the electrochemical potential difference of H⁺ (Larsen et al., 1974; Manson et al., 1977; Shioi et al., 1980), Na⁺ (Hirota and Imae, 1983), or both H⁺ and Na⁺ (Terahara et al., 2008) across cytoplasmic membrane. The flagellar motor of Escherichia coli and Salmonella entrica use the proton influx as the energy source to rotate counterclockwise (CCW) at the maximum speed of 300 revolution per second. When each motors rotate in CCW direction, flagellar filaments form a bundle to produce a thrust for the straight swimming. Quick reversal of motor rotation to clockwise (CW) generates a twist and induces the morphological change of the filament, thereby disrupting the flagellar bundle and so the cells tumble and change their swimming direction (Berg and Brown, 1972; Larsen et al., 1974) (Fig.1-1b). This rotational switch is caused by chemotastic signals like chemical attractants, repellents, temperature and pH changes. Therefore, bacteria can move towards more favorable environments from less favorable ones (Cluzel et al., 2000; Sourjik and Berg, 2002).

1.2 Flagellar structure

The flugellum is a large supermoleculer complex composed of 30 different proteins in copy numbers ranging from a few to a few tens of thousands. The flagellar structure can be divided into at least three parts; the basal body, the hook and the filament. The basal body is embedded in the cell membranes and acts as a rotary motor. The basal body contains a specific export apparatus, which exports protein subunits responsible for the formation of the exterior structures beyond the cellular membranes. The filament is a relatively rigid helical structure and functions as a helical propeller. The hook is flexible and acts as a universal joint (Berg, 2003; Macnab, 2003; Sarkar et al., 2009) (**Fig.1-2**).

1.2.1. Basal body

The basal body consists of the C-ring, the MS-ring, the LP-ring and the rod. spans the inner and outer membranes. About 10 stator units assemble around the basal body and so the basal body acts as a bi-directional rotary motor. The MS and C rings act as a rotor of the flagellar motor. The MS-ring is made up of a single cytoplasmic membrane protein, FliF, and works as a base for flagellar assembly. The C-ring is composed of three soluble proteins, FliG, FliM and FliN, and binds to the cytoplasmic face of the MS-ring through an interaction between FliF and FliG. The structure of the C-ring is well characterized by various structural analyses such as X-ray crystallography and cryo-electron microscopy (cryo-EM) to build its functional atomic model (Francis et al., 1992; Francis et al., 1994; Brown et al., 2002; Brown et al., 2005; Park et al., 2006; Sarkar et al., 2010; Lee et al., 2010; Mianmino et al., 2011; Paul et al., 2011). The C-ring is also called as the switch complex since FliG, FliM and FliN are involved in switching of the direction of flagellar motor rotation. A phosphorelated chemotastic

signal protein, CheY (CheY-P), which is produced by a two-component system activated from extracellular stimuli, regulates the switching frequency of the rotor ring complex. Phosphorelated CheY-P binds to both FliM and FliN to induce a conformational change in the switch complex and hence increases the probability that the motor rotates in the clockwise (CW) direction by (Bai et al., 2010; Sarkar et al., 2010a,b; Paul et al., 2011a,b). However, the detail switching mechanism is not yet clarified and several models are still present (Stock et al., 2012). The P and L rings spans the peptidoglycan layer and the outer membrane, respectively and acts as a bushing by encapsulating the rod that acts as a drive shaft (Akiba et al., 1991; Hizukuri et al., 2006, 2010).

The stator unit consists of two cytoplasmic membrane proteins, MotA and MotB. The stator functions as a proton channel that converts the energy derived from the proton influx through the channel to the rotation work (Suzuki et al., 2004; Thomas et al., 2006; Yonekura et al., 2011). Four copies of MotA and two copies of MotB form a functional stator complex (Braun and Blair, 2001; Kojima and Blair, 2004; Braun et al., 2004). MotA is a polytopic cytoplasmic membrane protein with four transmembrane helices (TM). Two conserved charged residues, Arg-90 and Glu-98, of MotA, which are located in the cytoplasmic loop between TM-2 and TM3, interact electrostatically with charged residues in the C-terminal domain of FliG to generate torque (Zhou and Blair, 1997; Zhou et al., 1998). The electrostatic interaction between MotA and FliG is also required for efficient stator assembly around the rotor. MotB is a bitopic cytoplasmic membrane protein and has a peptidoglycan-binding domain in its large C-terminal periplasmic domain (MotB_C). Dynamic conformational changes of MotB_C are required for the binding and anchoring of the stator complex to the peptidoglycan (Kojima et al., 2009), allowing the rotor to rotate relative to the cell wall.

1.2.2. The filament

The filament of *Salmonella* is a substructure of the helical tubular axial structure made of a few tens of thousands of a single protein component, flagellin (FliC or FljB). It grows around the length of $10 - 15 \mu m$. The filament propels the bacterial cell body by its rotation driven by the rotary motor (Fujime et al., 1972). The structure is composed of 11 protofilaments. Each protofilament is predicted to be in either a left-handed (L-type) or a right-handed (R-type) conformation to form the helical structure (Asakura, 1970). This is confirmed by atomic-resolution structural analyses of both L- and R-type the filaments by cryo-EM as well as by X-ray crystallographic analysis of FliC (Samatey et al., 2001; Yonekura et al., 2003; Maki-Yonekura et al., 2010).

1.2.3. The hook

The hook is a herical tubular axial substructure with its length of 55 ± 6 nm and is made of about 120 copies of a single protein, FlgE. The hook acts as a universal joint that smoothly transmits torque generated by the motor to the filament regardless of its orientation (Berg and Anderson, 1973). A whole atomic model of the hook structure is built by docking the atomic model of FlgE fragment solved by X-ray crystallography into a relative high resolution EM-density map of the hook (Samatey et al., 2004; Shaikh et al., 2005; Fujii et al., 2009). The hook subunits are tightly packed in the inner core domain whereas the outer two domains of FlgE are loosely packed in the axial inner subunit interactions in the proto-filament structure due to distinct difference of terminal alpha helices from those of the filament. So, the hook can exert the universal joint function.

1.2.4. Other structures

The filament and the hook are connected by the specific substructure, called junction, which is composed of three proteins, HAP1 (FlgK) and HAP3 (FlgL) and HAP2 (FliD). HAP2 forms the pentameric cap structure at the distal end of the filament to promote filament growth (Macnab, 2003; Berg, 2003).

The rod is a helical tubular structure and acts as a drive shaft connecting the rotor rings and the hook. The rod with an approximate length of 30 nm is a complex structure composed of five proteins, FliE, FlgB, FlgC, FlgF and FlgG. The distal rod freely rotates inside the LP-ring, of which friction is assumed to be low enough for smooth motor rotation (Khan et al., 1985; Akiba et al., 1991). Rod proteins have a strong tendency to form a fibrous structure (Saijo-Hamana et al., 2004). This makes the structural analysis of the rod components by X-ray crystallography difficult. Recently the truncation of the unfolded N- and C-terminal region of FlgG allowed its crystallization (Saijo-Hamano et al., 2013). Although the atomic model of the unusally elongated rod structure called polyrod has been built by docking atomic structure of FlgG fragment into an EM-density map of poly-rod made of FlgG (Fujii, T. 2009), the whole structure of the rod and the mechanism of the smooth rotation remain unknown.

1.3 Flagellar assembly

Flagellar assembly has been clarified by EM observation of each immature structure from various flagellar mutants (Suzuki et al., 1978; Suzuki and Komeda, 1981; Kubori et al., 1992) (Fig.1-3). The assembly of the flagellum begins with the self-assembly of the MS-ring composed of 26 copies of FliF in the cytoplasmic membrane (Fig.1-3 A). The export gate complex assembles into the central pore of the MS-ring during MS ring formation (Morimoto *et al.*, 2014). However, the assembly order of the MS-ring and the export gate complex is still unclear (Li and Sourjik, 2011; Wagner et al., 2010; Morimoto et al., 2014). After the formation of the MS-ring, the C-ring assembles at the cytoplasmic face of the MS-ring. Then the soluble components of the flagellar protein export apparatus forms the ATPase ring complex at the flagellar base (**Fig.1-3 B,C**). Recently, the ATPase ring complex has been visualized just outside the C ring by electron cryotomography (Chen *et al.*, 2011; Abrusci *et al.*, 2013; Kawamoto *et al.*, 2013). The flagellar axial structure, which consists of the rod, the hook, the junction, the filament cap and the filament, elongates in a tip-growth manner: Flagellar axial proteins are transported via the flagellar export apparatus from the cytoplasm to the distal end of the growing structure where their self-assembly occurs (Namba and Vonderviszt, 1997).

At first, five rod proteins (FliE, FlgB, FlgC, FlgF and FlgG) are exported to the periplasmic side of the MS-ring to form the rod structure with the help of the rod cap protein (FlgJ). The rod passes through the peptidoglycan layer by the peptidoglycan hydrolyzing activity of FlgJ (Nambu et al., 1999) (**Fig.1-3 D**). After the formation of the rod, FlgH and FlgI, which are exported into the periplasm by a general secretion pathway, assemble around the rod to form the LP-ring (**Fig.1-3 E, F**). After completion of the basal body, the hook component protein (FlgE) is transported to the tip of the rod and hook and self-assembles with the help of the hook cap made of five copies of FlgD until its length of 55 ± 6 nm (**Fig.1-3 G,H**) (Ohnishi et al., 1994; Hirano et al., 1994). The hook length is well controlled by an interaction between a ruler protein FliK and an export gate component FlhB. The interaction of the C-terminal domain of FliK with the C-terminal cytoplasmic domain of FlhB induces a conformational change of FlhB to switch substrate specificity of the export apparatus from rod-(FliE, FlgB, FlgC, FlgF,

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FlgG, FlgJ) and hook-type (FlgD, FlgE, FliK) to filament-type export substrates (FlgM, FlgK, FlgL, FliD, FliC). Upon hook completion, the hook cap complex is replaced by HAP1 (FlgK) (**Fig.1-3 I**), followed by the attachment of HAP3 (FlgL) on the HAP1 layer to form the hook-filament junction structure (**Fig.1-3 J**). Then the pentameric filament cap complex composed of FliD is formed at the tip of the hook-filament junction to facilitate the polymerization of flagellin (**Fig.1-3 K,L**). Flagellin subunits are transported to the distal end of the growing flagellum (**Fig.1-3 M,R**). The motor begins to rotate after the incorporation of the stator units around the rotor, and the flagellum propels cell body.

1.4 Type III flagellar protein export apparatus

For the construction of the axial flagellar structures, flagellar axial proteins must be exported to their final destination where their assembly occurs. Protein secretion beyond the cell membrane is one of the common mechanisms among all organs to maintain their biological activity. Bacteia have two major secretion pathways to secrete proteins into the periplasm (Wickner and Schekman, 2005; Natele et al., 2007). The general secretion route, termed Sec-pathway, catalyzes the transmembrane translocation of proteins in their unfolded conformation, and then the secreted proteins fold into their native structure at the trans-side of the membrane. Another pathway is the twin-arginine translocation pathway, termed Tat-pathway. It catalyzes the translocation of folded proteins. Both pathways need a specific signal sequence in the N-terminus of substrate proteins. However, among flagellar proteins, only five components FlgH and FlgI (LP-ring components), FlgA (peroplasmic chaperone), FlhE (putative plug or regulator of protone flux of flagellar protein export gate) and FliP (a component of flagellar protein export gate) have the signal peptides and are translocated via the Sec pathway (Homma et al., 1987; Jones et al., 1987, 1989; Kutsukake et al., 1994a; Nambu et al., 2000; Schoenhalas and Macnab, 1996; Minamino et al., 1994; Ohnishi et al., 1997; Paradel et al., 2004; Lee J et al., 2012; Lee J et al., 2015).

Fourteen flagellar proteins are translocated via its specific export apparatus, named the flagellar type III protein export apparatus from the cytoplasm into the central channel of the growing flagellar structure (Macnab, 2003; Minamino et al., 2008a) (**Fig.1-4**). The export apparatus is beleived to be located in the central pore of the MS-ring. The export apparatus is composed of a water-soluble ATPase complex consisting of FliH, FliI and FliJ, and a transmembrane export gate complex made of FlhA, FlhB, FliO, FliP, FliQ and FliR (Minamino and Macnab, 1999; Minamino and Macnab, 2000). In addition the export apparatus has several flagellar substrate specific chaperons (FlgN, FliA, FliS and FliT) for the ordered flagellar protein export (Vogler et al., 1991; Yokoseki et al., 1995; Mianamino and Macnab, 1999; Fraser et al., 1999; Auvray et al., 2001; Benntt et al., 2001; Aldridge et al., 2003; Minamino et al., 2012; Kinoshita et al., 2013). Recent structural analysis by cryo-electron tomography has revealed the whole complex architecture of the cytoplasmic domain of the export apparatus at low resolution (Murphy et al., 2006; Chen et al., 2011; Raddi et al, 2012; Abrusci et al., 2012; Kawamoto et al., 2013; Hu et al., 2015).

1.4.1. The ATPase complex

The ATPase complex is composed of three soluble components FliH, FliI and FliJ (**Fig.1-5 a**). Among these components, FliI shows the ATPase activity. Both amino acid sequence and structure of FliI shows high similarities with α and β subunit of F-type ATPase (Fan et al., 1996; Vogler et al., 1991; Imada et al., 2007) (**Fig.1-6 a**).

The α and β subunits form the $\alpha_3\beta_3$ ring complex in F-type ATPase whereas FliI forms a homo-hexamer (Claret et al., 2003; Kazetani et al., 2009; Minamino et al., 2006; Chen et al., 2011; Kawamoto et al., 2013; Bai et al; 2014). Although FliJ does not show a significant sequence similarity with components of both F- and V-type ATPase, its coiled-coil structure show extensive structural similarities with both γ subunit of F-type ATPase and D subunit of V-type ATPase, which locate at the central pore of $\alpha_3\beta_3$ and A₃B₃ ring, respectively (Ibuki et al, 2011) (**Fig.1-6** b). FliJ binds to the central pore of the FliI₆ ring to form FliI₆FliJ complex in a way similar to the γ and D subunits. Interestingly, FliJ can bind to the center of the A₃B₃ ring complex to act as a rotary stalk in V-type ATPase, suggesting that the type III export apparatus and F- and V-type ATPase families share the same evolutional origin (Kishikawa et al, 2013) (**Fig.1-6 c,d**). FliJ binds to not only FliI but also FliH and the axial components to prevent their aggregation in cytoplasm. The amino acid sequence of FliH also shows high similarities with b and δ subunit of F-type ATPase (Pallen et al., 2006).

Previous biochemical analysis has revealed that FliH binds to FliI to form the FliH₂FliI complex in the solution (Minamino and Macnab, 2000; Auvray et al., 2002). Since FliH interacts with a C-ring component FliN and the export gate component FlhA through its N-terminal domain, FliH is proposed to stabilize the localization of the FliI₆FliJ ring complex at the base of the flagellum (González-Pedrajo et al., 2002; Minamino et al., 2009; Hara et al., 2012). Since the FliH₂FliI complex bind to export substrates and chaperone-export substrate complexes (Thomas et al., 2004; Imada et al., 2010; Minamino et al., 2012a), the FliH₂FliI complex is proposed to deliver the export substrates to the export gate complex.

1.4.2. The export gate complex

The trasnmembrane export gate complex is composed of six integral inner membrane proteins, FlhA, FlhB, FliO, FliP, FliQ and FliR and is believed to be located in the central pore of the MS-ring (Mianmino and Macnab, 1999) (**Fig.1-5 b**). In contrast to the cytoplasmic ATPase complex, these components do not show any amino acid sequence similarities.

FlhA is the most conserved component among these proteins. Previous genetic and biochemical study indicated that FlhA interacts with all of other export gate components and the MS-ring component, FliF (Kihara et al., 2001; McMurry et al., 2004; Hara et al., 2011; Li and Sourjik, 2011; Barker and Samatey, 2012). FlhA has a large cytoplasmic domain (FlhA_C) (Minamino et al, 1994). X-ray crystallographic analysis of FlhA_C revealed FlhA_C consists of four subdomains, D1, D2, D3 and D4 and a flexible linker connecting FlhA_C with the N-terminal transmembrane domain with eight TM helices (Saijo et al., 2010; Bange et al., 2010). A highly conserved hydrophobic dimple located at an interface between D1 and D2 is involved in the interaction with flagellar secretion chaperones FlgN, FliS and FliT (Mianmono et al., 2012b; Kinoshita et al., 2013). FlgN, FliT and FliS are chaperons specific for FlgK/L, FliD and FliC respectively, and their binding affinities for FlhA_C increase when they form a complex with their substrates. The binding affinity of the FlgN/FlgK complex for FlhA_C is slightly higher than that of the FliT/FliD complex and is about 14-fold higher than that of the FliS/FliC complex. This suggests that FlhA_C interacts with the FlgN/FlgK and FlgN/FlgL complexes prior to the FliT/FliD and FliS/FliC complexes to form the hook-filament junction zone upon completion of hook assembly (Evdokimov et al., 2003; Imada et al., 2010; Mianmono et al., 2012 Kinoshita et al., 2013). The cytoplasmic domain of FlhA homologue, MxiA, formed the nonameric ring structure in

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the crystal. Structure-based mutagenesis has revealed that $MxiA_C$ ring formation is required for protein translocation. The electron densities corresponding $MxiA_C$ and FlhA_C were observed by cryo-electron tomography, indicating that FlhA forms a nonamer in the MS-ring in a way similar to MxiA (Abrusci et al., 2012; Kawamoto et al., 2013). FlhB has a large cytoplasmic domain (FlhB_C), too and is involved in export specificity swiching from rod/hook-type to filament-type export class (Mianmino et al., 2000). Self-cleavage of FlhB_C and the following interaction with the C-terminal domain of FliK after the hook completion result in the conformational change of FlhB_C to swich the export specificity (Kutsukake et al., 1994; Williams et al., 1996; Minamino and Macnab, 2000; Ferris et al., 2005; Moriya et al., 2006). Therefore, both FlhA_C and FlhB_C play important roles in coordinating protein export with flagellar assembly. In contrast to FlhA_C and FlhB_C, the function and structure of the transmembrane regions of FlhA and FlhB remain unknown.

Little of functions and structures of FliO, FliP, FliQ and FliR is known by the same reason. FliO is a 13 kDa bitopic membrane protein. FliO is not essential for flagellar construction (Barker et al., 2014). Indeed, several bacteria do not have the *fliO* gene in their genome. All bypass mutations isolated from a *fliO* null mutant have been found in the *fliP* gene, indicating that FliO has something important role to regulate FliP function (Barker et al., 2010). FliO has a large C-terminal cytoplasmic domain and is assumed to be involved in flagellar protein export. FliP is a 25 kDa polytopic cytoplasmic membrane protein and has a signal peptide at its N-terminus (Malakooti et al., 1994; Ohnishi et al., 1997). FliP is translocated and inserted in the cytoplasmic membrane by the Sec pathway. The N-terminal signal peptide of FliP is cleaved during membrane insertion, producing mature FliP with the molecular weight of 23 kDa (Ohnishi et al., 1997; Pradel et al., 2004). The mature form of FliP is predicted to have

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four TM helices and have a 10 kD periplasmic domain (FliP_P) between TM-2 and TM-3. FliP is identified in purified basal body and about four copies of FliP molecules are estimated to associate with the MS-ring (Fan et al., 1997). FliQ has two TM helices and its molecular weight is 10 kDa. The function and structure of FliQ are almost unknown. FliR is a 29 kDa cytoplasmic membrane protein and is predicted to have six TM helices. FliR is visualized to associate with the MS ring by immno-electron microscope (Fan et al., 1997). Since a fusion protein of FliR and FlhB is functional, the stoichiometry of FliR and FlhB is assumed to be 1 to 1 (Van Arnam et al., 2010).

It has been shown that the FliI ATPase activity is dispensable for flagellar protein export and that proton motive force (PMF) across the cytoplasmic membrane is a primary energy source (Minamino and Namba, 2008; Paul et al., 2008). The export gate complex can utilize only the membrane potential ($\Delta \psi$) of the PMF when FliJ interacts with the linker region of FlhA in the presence of FliH and FliI (Minamino et al., 2011). Since FliJ can work a rotary stalk in the A₃B₃ ring complex of V-type ATPase (Kishikawa et al., 2013), FliJ could rotate in the FliI₆ ring by ATP hydrolysis to induces a conformational change of FlhA thorough the interaction of FliJ with FlhA, allowing the gate to drive flagellar protein export in a $\Delta \psi$ -dependent manner (Kishikawa et al., 2013; Minamino et al., 2014) (**Fig.1-7**).

1.5 Aims and outline of my Ph.D work

Genetic and biochemical studies of the assembly mechanism of the injectisome of *Salmonella*, which is a highly homologous structure of the FBB, indicate that the export gate complex assembles in a coordinated manner prior to the assembly of the inner ring complex of the injectisome, which corresponds to the MS-ring of the FBB (Wagner et al., 2010). SpaP (FliP homologue) and SpaR (FliR homologue) form a

stable complex, which is followed by the assembly of SpaQ (FliQ homologue) and SpaS (FlhB homologue) and finally InvA (FlhA homologue). The export gate complex allows the inner ring of the injectisome to be more efficiently formed around the gate. It has been shown that FliO contributes to the protein stability of FliP within the cytoplasmic membrane (Barker et al., 2010). The assembly of FlhA-YFP into the export gate complex is dependent on FliO, FliP, FliQ and FliR but not on FlhB (Morimoto et al., 2014), raising the possibility that the assembly of the flagellar export gate complex begins with FliO and FliP, followed by FliQ and FliR, and finally FlhA and FlhB.

To understand the assembly mechanism of the export gate complex in more detail, my Ph.D work is focused on the structural characterization of FliP. Because FliP has a large periplasmic domain between TM-2 and TM-3 (**Fig.1-8**), I first characterized the structure and function of the FliP_P (Chapter 2). I solved the crystal structure of FliP_P from *Thermotoga maritima*. Structure-based mutational analyses reveal that FliP forms a homo-dimer through the FliP_P-FliP_P interaction *in vivo*. To clarify the oligomeric state of FliP, I over-expressed and purified full-length FliP of *Salmonella*. I show that FliP efficiently assembles into a hexagonal ring oligomer with the help of FliO and that the FliP_P-FliP_P interaction is required for FliP ring formation (chapter 3). Based on these results, I will discuss the assembly mechanism of FliP. Finally, I establish the co-expression system of the FliO/P/Q/R-FlhB/A-FliF/G complex to clarify the further detail molecular mechanism of the export gate assembly in the future (chapter4).



Figure1-1. Bacterial flagellum.

(a) Electron micrograph of *Salmonella enterica* serovar Typhimurium. Typical size of the cell body is about 2 μ m. In this micrograph, five flagellar filaments form a bundle. (b) A typical swimming pattern of *Salmonella* cells.



Figure1-2. Major substructures of flugellum..

Schematic diagram of the bacterial flagellum and substructures. The filament is composed of tens of thousands of a single component, FliC. The hook is composed of a single component, FlgE. At least 10-12 stator complexes surround the rotor complex. (Samatey et al., 2001;



Figure1-3. The assembly process of the flagellum proceeds from (a) to (r).



(Kawamoto et al., 2013)



Figure1-4. Architecture of flagellar protein export apparatus.

(a) *In situ* structure of the basal body revealed by the cryo-electron tomography. Red arrow indicates the cytoplasmic domain of FlhA. Green arrow indicates the FliI ATPase ring complex. (b) Schematic architecture of the flagellar protein export apparatus. The export gate utilizes proton motive force as an essential energy source. The FliI ATPase activity is required for efficient flagellar protein export.

а







(a) Components of the cytoplasmic ATPase complex. Crystal structures of FliI (PDB ID: 2DPY) and FliJ (PDB ID: 3AJW) are shown in rainbow. (b) Topology model of export gate components. Crystal structures of $FlhA_C$ (PDB ID : 3A5I) and $FlhB_C$ (PDB ID: 3B0Z) are depicted in rainbow.





(a) Superimposition of FliI (cyan) onto the β subunit of F-type ATPase (green, PDB ID: 1BMF) and the B subunit of V-type ATPase (red, PDB ID: 3VR6). (b) Superimposision of FliJ (cyan) on the γ subunit of F-type ATPase (green) and the D subunit of V-type ATPase (red). (c) 2-D cryo-EM average images of FliI₆ and A₃B₃ with or without FliJ. The structural models of FliI₆J and A₃B₃J are shown on the right. (d) Comparison between the architecture of the export apparatus and F_O-F₁ ATPase.



Figure1-7. Current model of flagellar protein export mechanism

ATP hydrolysis by FliI activates the export gate through an interaction between FliJ and FlhA, allowing the export gate complex to efficiently utilize the membrane potential $(\Delta \psi)$ of proton motive force to unfold and transport export substrates into the central channel of the flagellar axial structure.



b



Figure1-8. Multiple sequence alignment of FliP homologues and topology model of FliP

Chapter 2 Structural and functional characterization of the periplasmic domain of FliP

2.1. Introduction

It has been reported that overexpression of full-length FliP causes a remarkable growth arrest of *E. coli* host cells (Kinoshita, M., Imada, K. and Minamino, T. personal communications). This makes it difficult to carry out biochemical analysis of FliP. Therefore, to clarify the role of FliP in flagellar protein export, I first carried out structural and functional characterization of a relatively large periplasmic domain of FliP (FliP_P), which is predicted to be located between TM-2 and TM-3. Although we use *Salmonella* as a model organism for flagellar research, FliP_P from *Salmonella typhimurium* was expressed as an inclusion body and strongly tends to aggregate during its refolding and purification (Fukumura, T. 2013). In contrast, FliP_P from *Thermotoga maritima* (Tm-FliP_P) was highly soluble, and so I obtained the hexagonal crystal of Tm-FliP_P and detemined its crystal structure at 2.4 Å resolution. In this chapter, I will show the details of the Tm-FliP_P structure. I also show that the FliP_P-FliP_P interaction is critical for FliP function. Based on the obtained results, I will discuss a biological unit of FliP and will also discuss how the FliP_P-FliP_P interaction contributes to FliP assembly into the export gate complex.

2.2. Materials and Methods

1.2.1. Bacterial strains, plasmids and media

Bacterial strains and plasmids used in this chapter are listed in Table 2-1. Compositions of media used in this study are as follows:

- Luria-Bertani Broth (LB) medium : H₂O 1 l, BactoTM Tryptone (Becton, Dickinson and Company) 10 g, BactoTM Yeast Extract (Becton, Dickinson and Company) 5 g, NaCl 5 g
- 2×YT medium : H₂O 1 l, BactoTM Tryptone 16 g, BactoTM Yeast Extract 10 g, NaCl 5 g
- Terrific Broth (TB) medium : H₂O 1 l, BactoTM Tryptone 12 g, BactoTM Yeast
 Extract 24 g, glycerol 8 ml, K₂HPO₄ 9.4 g, KH₂PO₄ 2.2 g
- LA plate : LB 1 l, BactoTM Agar (Becton, Dickinson and Company) 15 g
- Soft tryptone agar plate : H₂O 1 l, BactoTM Tryptone 10 g, BactoTM Agar 7 g, NaCl
 5 g
- SeMet minimal medium : H₂O 1 l, NH₄Cl 1 g, KH₂PO₄ 3 g, Na₂HPO₄·12H₂O 8 g, glucose 20 g, MgSO₄·7H₂O 0.6 g, Fe₂(SO₄)₃ 10 mg, thiamine 10 mg, L-SeMet 50 mg (Guerrero *et al*, 2001; Sakane, 2008)

To maintain plasmids in the bacterial host cells, ampicillin were added to each media at a final concentration of 50-100 μ g/ml

1.2.2. DNA manipulations

To construct a plasmid for the crystallization, a DNA fragment encoding $FliP_P$ (Tm-FliP_P) consisting of residues 110-188 of *T. maritima* FliP was generated by PCR with a plasmid pGS1-*TmfliP* (Gene Design Inc.) as a template. The amplified PCR product was inserted into the *NdeI* and *Bam*HI sites of the pET15b vector.

To construct pUC19-based plasmids for both disulfide cross-linking experiments and genetic characterization, a DNA fragment encoding *fliP* was amplified by PCR with genomic *Salmonella* DNA as a template. The amplified DNA fragment was inserted into the *NdeI* and *Bam*HI sites of the pET3c vector. Then, the DNA fragment encoding the *fliP* gene with a ribosome-binding site of the pET3c vector was subcloned into the *XbaI-Bam*HI sites of pUC19. Each mutation for disulfide cross-linking experiments and genetic characterization was generated by quickChange site-directed mutagenesis method as described in the manufacturer's instruction (Strategene).

To construct pBAD24-based plasmids for genetic characterization, a DNA fragment encoding *fliP* with the 30 bp upstream region was amplified by PCR with genomic DNA of *Salmonella* as a template. The amplified DNA fragment was inserted into the *EcoRI* and *KpnI* sites of the pBAD24 vector. Each mutation for genetic characterization was generated by site-directed mutagenesis.

DNA sequence reactions were carried out using BigDye v3.1 (Applied Biosystems) and DNA sequencing was performed with an Applied Biosystems 3130 Genetic Analyzer (Applied Biosystems).

1.2.3. Protein expression and purification

To purify Tm-FliP_P for crystallization, a 30 ml overnight culture of *Escherichia coli* strain BL21 (DE3) harboring a plasmid encoding TmFliP_P with a hexahistidine tag attached at its N-terminus (pKY045) was inoculated into 2.51 LB medium. Cells were grown at 30 °C until the culture density had reached an $OD_{600} = 0.4$ - 0.8. Expression of _{His}Tm-FliP_P was induced by isopropyl β-D-1-thiogalactopyranoside (IPTG) at a final concentration of 0.4 mM, and then culture was incubated for another 6

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h. The cells were harvested by centrifugation (6,400 g, 10 min, 4 °C) and stored at -80 °C. The cells were thawed, suspended in Buffer A (50 mM Tris-HCl pH 8.0, 300 mM NaCl) containing 20 mM imidazole and sonicated (ASTRASON model XL2020 sonicator, Misonix Inc.). The cell lysate was centrifuged (110,000 g, 40 min, 4 °C) to remove cell debris. The supernatant was loaded onto a Ni-NTA agarose resin (QIAGEN) equilibrated with Buffer A containing 20 mM imidazole. Protein was then eluted with a 50-300 mM imidazole gradient in Buffer A and fractions containing HisTmFliP_P were collected. The His₆ tag was removed by thrombin (GE Healthcare) to the HisTm-FliP_P solution. The reaction mixture was dialyzed overnight at 4 °C against Buffer A with 200 mM imidazole, followed by Buffer A with 100 mM imidazole and finally buffer A with 20 mM imidazole. Tm-FliP_P was purified using a HisTrap HP 5 ml (GE Healthcare) to remove the N-terminally His-tagged polypeptide, non-cleaved HisTm-FliP_P and thrombin. The eluted fractions containing Tm-FliP_P was concentrated using a Vivaspin 20 (30,000 MWCO, Sartorius), dialyzed overnight against Buffer B (10 mM sodium phosphate pH 7.0, 100 mM NaCl) and loaded a HiLoad Superdex 75 (26/60) column (GE Healthcare) equilibrated with Buffer B. Fractions containing TmFliP_P were collected. The purity of the product was examined by SDS-PAGE and MALDI-TOF mass spectrometry (Voyager DE/PRO, Applied Biosystems).

To purify a SeMet derivative of Tm-FliP_P (SeMet_Tm-FliP_P) for phase determination, we used the *E. coli* B834 (DE3) strain as a host. Cells grown overnight in LB at 30 °C were harvested by centrifugation (5,000 g, 5 min, 4 °C), washed and suspended in 0.5%(w/v) NaCl. The cells were inoculated into the SeMet minimal medium and grown at 37 °C until the cell density had reached an OD₆₀₀ of 0.4. Expression of SeMet-labelled _{His}Tm-FliP_P was induced with 0.4 mM IPTG and the culture was continued for another 4 h. The cells were harvested by centrifugation (6,400

g, 10 min, 4 °C) and stored at 4 °C. SeMet_Tm-FliP_P was purified in the same way with native Tm-FliP_P. Incorporation of SeMet into TmFliP_P was confirmed by MALDI-TOF mass spectrometry.

1.2.4. Analytical ultracentrifugation of Tm-FliP_P

Analytical ultracentrifugation was carried out using a Beckman Optima XL-A analytical ultracentrifuge with an AnTi 60 rotor. The protein samples were extensively dialyzed against Buffer B prior to analytical ultracentrifugation. Sedimentation equilibrium measurements were performed at 4 °C at speeds of 22,000, 24,000 and 26,000 rpm on a sample at an initial concentration of 0.85 mg/ml, using a two channel charcoal-filled epon and quartz windows. Scans were collected at a wavelength of 280 nm at a spacing of 0.001 cm in step mode with twenty averages per step. The equilibrium of the system was judged by superimposition of the three scans. Sedimentation equilibrium data of TmFliP_P whose partial specific volume v-bar was calculated to be 0.736 ml/g from the amino acid composition of the protein were analyzed for the average molecular weight by the program OptimaTM XL-A/XLI version 6.04.

1.2.5. Crystallization of Tm-FliP_P

Purified Tm-FliP_P was concentrated and passed through a 0.20 μ m Millex-LG syringe filter (Millipore). The protein concentration was estimated based on its Abs₂₈₀ of 0.47 for a 1 mg ml⁻¹ solution. Initial crystallization screening of Tm-FliP_P was performed at 4 °C and 20 °C by the sitting-drop vapor-diffusion technique using NeXtal Evolution plates (QIAGEN) and the following screening kits: Wizard I and II, Cryo I and II (Emerald Biostructures) and Crystal Screen I and II (Hampton Research). Each

drop was prepared by mixing 1.0 μ l protein solution (20 mg ml⁻¹ Tm-FliP_P, 10 mM sodium-phosphate pH 7.0, 100 mM NaCl) with 1.0 μ l reservoir solution, and equilibrated against 60 μ l reservoir solution. The crystallization conditions were optimized by varying the precipitant concentration, pH and additives using the hanging-drop method with VDX plates (Hampton Research). Finally, crystals suitable for X-ray analysis were obtained from drops prepared by mixing 1.5 μ l protein solution (5 mg ml⁻¹) with 1.5 μ l reservoir solution containing 0.1 M phosphate-citrate pH 4.4, 36% 2-Methyl-2,4-pentanediol (MPD) at 4 °C within a week.

1.2.6. X-ray diffraction data collection and processing

All X-ray diffraction data were collected at SPring-8 beamline BL41XU (Harima, Japan). Crystals were mounted into nylon cryo-loops (Hampton Research). Since the concentration of MPD in the crystallization drops was high enough for cryoprotection, the crystals were directly transferred into liquid nitrogen for freezing. The diffraction data were recorded on a MX225HE CCD detector (Rayonix) at -173 °C using a nitrogen gas flow to reduce radiation damage. The diffraction data were indexed, integrated and scaled using the programs iMOSFLM (Battye et al., 2011) and SCALA (Evans, 2006) from the CCP4 program suite (Winn et al., 2011). The statistics of data collection are summarized in Table 1. The model was determined and refined with PHENIX (Adams et al., 2010). The missing residues in the auto-built model were manually added in COOT (Emsly et al., 2010). All structural figures were prepared by Chimera (https://www.cgl.ucsf.edu/chimera/).

1.2.7. In vivo disulfide cross-linking and immunoblotting

Salmonella cells were exponentially grown in 50 ml LB at 30 °C with shaking. The cells were harvested and resuspended in 5 ml of PPBS. For oxidization, iodine was added to the suspended *Salmonella* cells at a final concentration of 0.2 mM and incubated for 5 min at room temperature. To block disulfide bond formation for the remaining cyctein, N-ethylmalemide was added to the cell suspensions at a final concentration of 20 mM and incubated for 5 min at room temperature. The cells were washed with PBS and resuspended in 10 ml of PBS, and then disrupted by sonication (ASTRASON model XL2020 sonicator, Misonix Inc.). The cell lysate was centrifuged at 20,000g for 15 min to remove cell debris. Supernatant was ultracentrifuged at 110,000 g for 30 min to isolate the membrane fraction. The membrane fractions were resuspended in PBS to adjust the total membrane protein concentration of 10 mg ml⁻¹. The membranes were mixed with non-reducing SDS loading buffer. After sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE), immunoblot was carried out with monoclonal anti-HA antibody and detected by an ECL immunoblotting detection kit (GE Healthcare).

1.2.8. Motility assay

Fresh transformant colonies were inoculated onto soft tryptone agar plates and incubated at 30°C. For the induction from pBAD24, 0.2 % arabinose was added to soft tryptone agar plates. The motility rings were analyzed by imageJ.

1.2.9. Preparation of whole cell fractions and immunoblotting

Salmonella cells were grown at 30°C with shaking until the cell density had reached an OD_{600} of 1.2–1.8. Aliquots of culture proteins containing a constant number of cells were clarified by centrifugation. Cell pellets were resuspended in SDS-loading
buffer normalized by cell density to give a constant amount of cells. After SDS-PAGE, immunobloting was carried out with monoclonal anti-HA antibody and detected by an ECL immunoblotting detection kit (GE Healthcare).

1.2.10. Multiple sequence aliginment

Multiple sequence alignment was performed by CLUSTAL-Ω (http://www.ebi.ac.uk/Tools/msa/clustalo/). Depection of the multiple sequence alignment was carried out by similarity coloring scheme of %equivalent in ESprint3 server (http://espript.ibcp.fr/ESPript/ESPript/).

Strains or Plasmids	Relevant properties	Source or reference
E.coli		
BL21 (DE3)	Host for overexpression from the T7 promoter	Novege
Salmonella		
SJW1103	Wild type for motility and chemotaxis	(Yamaguchi et al., 1984)
TH	ΔfliP	
Plasmids		
pET3c	Cloning vector	Novagen
pET15b	Expression vector	Novagen
pBAD24	Expression vector	(Guzman et al., 1995)
pUC19	Expression and cloning vector	Invitrogen
pKY045	pET15b/TmFliP ₁₁₀₋₁₈₈	This study
pKY049	pET3c/ _{HA} FliP	This study
pKY041	pBAD24/ _{HA} FliP	This study
pKY050	pBAD24/ _{HA} FliP(F137A)	This study
pKY051	pBAD24/ _{HA} FliP(M138A)	This study
pKY052	pBAD24/ _{HA} FliP(Q141A)	This study
pKY053	pBAD24/ _{HA} FliP(Y174A)	This study
pKY054	pBAD24/ _{HA} FliP(E178A)	This study
pKY010	pUC19/ _{HA} FliP	This study
pKY014	pUC19/ _{HA} FliP(F137A)	This study
pKY055	pUC19/ _{HA} FliP(M138A)	This study
pKY056	pUC19/ _{HA} FliP(Q141A)	This study
pKY019	pUC19/ _{HA} FliP(Y174A)	This study
pKY021	pUC19/ _{HA} FliP(E178A)	This study
pKY057	pUC19/ _{HA} FliP(G130C-A182C)	This study
pKY058	pUC19/ _{HA} FliP(P133C-A182C)	This study
pKY059	pUC19/ _{HA} FliP(P133C-E178C)	This study
pKY060	pUC19/ _{HA} FliP(P133C-T181C)	This study
pKY061	pUC19/ _{HA} FliP(F137C)	This study
pKY062	pUC19/ _{HA} FliP(F137C-E178C)	This study
pKY063	pUC19/ _{HA} FliP(R140C)	This study
pKY064	pUC19/ _{HA} FliP(E144C)	This study
pKY065	pUC19/ _{HA} FliP(M123C-L153C)	This study

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pKY066	pUC19/ _{HA} FliP(M123C-T176C)	This study
pKY067	pUC19/ _{HA} FliP(Q124C-S156C)	This study
pKY068	pUC19/ _{HA} FliP(L127C-P172C)	This study

Primers	Sequence
Tm-FliPp.Fw	GGAATTCCATATGTACAACAATGCCATAACGCCG
Tm-FliPp.Re	CGCGGATCCTCATTTGAAGGCAACTTCCAGTTC
FliP_NdeI.Fw	acacaagcatATGCGCCGTTTGTTTGTTATTCCT
FliP_BamHI.Re	cgcggatccCTAACTGTAAAAGCTTTGGG
FliP+30bp_EcoRI.Fw	ccggaattcgagcttactcaagcgttccg
FliP_KpnI.Re	cggggtaccCTAACTGTAAAAGCTTTGGG
SigPep_HA.Fw	CCGCCGCCGCTGCGCAAtacccatacgatgttccagattacgctCTGCCGGGGGCTTATCAG
SigPep_HA.Re	CTGATAAGCCCCGGCAGagcgtaatctggaacatcgtatgggtaTTGCGCAGCGGCGGCGGGGGGGGGGGGGGGGGGGGGGGG
FliP_F137A.Fw	GCGCAACCGTTACGCGCGgcCATGCTGCGCCAAACCCGC
FliP_F137A.Re	GCGGGTTTGGCGCAGCATGgcCGCGCGTAACGGTTGCGC
FliP_M138A.Fw	CAACCGTTACGCGCGTTCgcGCTGCGCCAAACCCGCGAAG
FliP_M138A.Re	CTTCGCGGGTTTGGCGCAGCgcGAACGCGCGTAACGGTTG
FliP_Q141A.Fw	CGCGCGTTCATGCTGCGCgcgACCCGCGAAGCCGATCTG
FliP_Q141A.Re	CAGATCGGCTTCGCGGGGTcgcGCGCAGCATGAACGCGCG
FliP_Y174A.Fw	CGTATCCTCCTGCCCGCTgcgGTCACCAGCGAATTAAAG
FliP_Y174A.Re	CTTTAATTCGCTGGTGACcgcAGCGGGCAGGAGGATACG
FliP_E178A.Fw	CCCGCTTATGTCACCAGCGcATTAAAGACGGCGTTTCAG
FliP_E178A.Re	CTGAAACGCCGTCTTTAATgCGCTGGTGACATAAGCGGG
FliP_M123C.Fw	GCGAGCAGAAAATTTCTtgcCAGGAAGCGCTGGATAAAG
FliP_M123C.Re	CTTTATCCAGCGCTTCCTGgcaAGAAATTTTCTGCTCGC
FliP_Q124C.Fw	GAGCAGAAAATTTCTATGtgcGAAGCGCTGGATAAAGGC
FliP_Q124C.Re	GCCTTTATCCAGCGCTTCgcaCATAGAAATTTTCTGCTC
FliP_L127C.Fw	GAAAATTTCTATGCAGGAAGCGtgtGATAAAGGCGCGCAACCG
FliP_L127C.Re	CGGTTGCGCGCCTTTATCacaCGCTTCCTGCATAGAAATTTTC
FliP_G130C.Fw	CAGGAAGCGCTGGATAAAtGCGCGCAACCGTTACGCGCG
FliP_G130C.Re	CGCGCGTAACGGTTGCGCGCaTTTATCCAGCGCTTCCTG
FliP_P133C.Fw	CTGGATAAAGGCGCGCAAtgcTTACGCGCGTTCATGCTG
FliP_P133C.Re	CAGCATGAACGCGCGTAAgcaTTGCGCGCCTTTATCCAG
FliP_F137C.Fw	GCGCAACCGTTACGCGCGTgCATGCTGCGCCAAACCCGC
FliP_F137C.Re	GCGGGTTTGGCGCAGCATGcACGCGCGTAACGGTTGCGC
FliP_R140C.Fw	GTTACGCGCGTTCATGCTGtGCCAAACCCGCGAAGCCGAT
FliP_R140C.Re	ATCGGCTTCGCGGGTTTGGCaCAGCATGAACGCGCGTAAC
FliP_E144C.Fw	CATGCTGCGCCAAACCCGCtgcGCCGATCTGGCGCTGTTTG

Table 1-2. Primers used in this study of this chapter

FliP_E144C.Re	CAAACAGCGCCAGATCGGCgcaGCGGGTTTGGCGCAGCATG
FliP_L153C.Fw	CTGGCGCTGTTTGCCCGTtgcGCCAATAGCGGTCCGTTAC
FliP_L153C.Re	GTAACGGACCGCTATTGGCgcaACGGGCAAACAGCGCCAG
FliP_S156C.Fw	GTTTGCCCGTCTGGCCAATtgcGGTCCGTTACAGGGACCG
FliP_S156C.Re	CGGTCCCTGTAACGGACCgcaATTGGCCAGACGGGCAAAC
FliP_L172C.Fw	GATGCGTATCCTCCTGtgcGCTTATGTCACCAGCGAATTAAAG
FliP_L172C.Re	CTTTAATTCGCTGGTGACATAAGCgcaCAGGAGGATACGCATC
FliP_T176C.Fw	GTATCCTCCTGCCCGCTTATGTCtgcAGCGAATTAAAGACGGCG
FliP_T176C.Re	CGCCGTCTTTAATTCGCTgcaGACATAAGCGGGCAGGAGGATAC
FliP_E178C.Fw	CCCGCTTATGTCACCAGCtgcTTAAAGACGGCGTTTCAG
FliP_E178C.Re	CTGAAACGCCGTCTTTAAgcaGCTGGTGACATAAGCGGG
FliP_T181C.Fw	GTCACCAGCGAATTAAAGtgcGCGTTTCAGATCGGTTTTAC
FliP_T181C.Re	GTAAAACCGATCTGAAACGCgcaCTTTAATTCGCTGGTGAC
FliP_A182.Fw	ACCAGCGAATTAAAGACGtgcTTTCAGATCGGTTTTACG
FliP_A182.Re	CGTAAAACCGATCTGAAAgcaCGTCTTTAATTCGCTGGT

Table 1-3. Summary of the data statistics.

Values in parentheses indicate statistics for the highest resolution shell.

	Native	SeMet derivative
Space group	<i>P</i> 6 ₂ 22	<i>P</i> 6 ₂ 22
Cell dimensions		
a,b,c (Å)	114.9, 114.9, 193.8	115.3, 115.3, 193.6
α,β,γ (°)	90, 90, 120	90, 90, 120
Wavelength (Å)	1	0.9791
Resolution (Å)	43.6 - 2.4 (2.53 - 2.4)	43 - 2.8 (2.95 - 2.8)
No. of reflections	214552	257930
No. of unique reflections	30262	18573
Rmerge	0.088 (0.357)	0.097 (0.23)
Ι/σΙ	13.5 (5.5)	17.0 (5.4)
Completeness (%)	99.9 (100.0)	95.1 (67.6)
Redundancy	7.1 (7.2)	7.5 (2.4)
Figure of Merit		0.48

Data collection

[#] $R_{\text{merge}} = \sum_{\mathbf{h}} \sum_{l} |I_{\mathbf{h}l^{-}} \langle I_{\mathbf{h}} \rangle| \sum_{\mathbf{h}} \sum_{l} \langle I_{\mathbf{h}} \rangle$, where I_{l} is the *l*th observation of reflection **h** and $\langle I_{\mathbf{h}} \rangle$ is the weighted average intensity for all observations *l* of reflection **h**. ⁺ $R_{\text{ano}} = \sum_{\mathbf{h}} |\langle I(\mathbf{h}+) \rangle - \langle I(\mathbf{h}-) \rangle| \sum_{\mathbf{h}} (\langle I(\mathbf{h}+) \rangle + \langle I(\mathbf{h}-) \rangle)$, where $\langle I(\mathbf{h}+) \rangle$ and $\langle I(\mathbf{h}-) \rangle$ correspond to the average intensities of each Friedel pair for reflection **h**.

Table 1-4. X-ray refinement statistics.

Values in parentheses indicate statistics for the highest resolution shell.

Refinement		
Measurement	Value	
Resolution (Å)	42.9 - 2.4	
No. of reflections	30084	
Rwork/Rfree	0.211 / 0.258	
No. atoms		
Protein	4354	
Water	397	
B-factor		
Protein	43.6	
Water	52	
R.m.s deviations		
Bond lengths (Å)	0.004	
Bond angles (°)	0.778	
エラー! リンクが正し	くありません。	

 $R_w\!\!=\!\sum ||F_o|$ - $|F_c||$ / \sum $|F_o|,$ same as R_{free} but calculated on 5% of data set aside for refinement.

2.3. Results

2.3.1. Purification and crystallization of Tm-FliP_P

The size exclusion chromatography showed that the molecular weight of eluted Tm-FliP_P is about 40 kDa (Fig.2-1a). Since the molecular weight of Tm-FliP_P was deduced about 10 kDa, purified Tm-FliP_P would form a tetramer. To confirm whether Tm-FliP_P is actually in a tetrameric state in solution, I carried out sedimentation equilibrium analytical ultracentrifugation. A single species model with the molecular weight of 39.7 kDa produced the best fit (Fig.2-1b). Therefore, I conclude that Tm-FliP_P forms a homo-tetramer in solution. Some of the tetramers, however, were dissociated into dimers in a few days. To stabilize the tetramer, I varied pH and NaCl concentrations and found that the tetramer is quite stable at pH less than 7.0. I obtained Tm-FliP_P crystals under several conditions that contain low molecular weight compounds as a precipitant at low pH (less than 8.0). The crystals were especially well grown in a solution containing 0.1 M phosphate-citrate pH 4.4, 36% MPD at 4 °C (Fig.2-2). The TmFliP_P crystals diffracted to 2.4 Å resolution and belonged to the hexagonal space group P6222. SeMet Tm-FliPP crystals were obtained under the same condition as the native Tm-FliP_P ones. The shape and size of the SeMet Tm-FliP_P crystals were almost the same as those of the native ones and its space group was also *P*6₂22.

2.3.2. Crystal structure of Tm-FliP_P

Tm-FliP_P is composed of four α helices (α 1, α 2, α 3 and α 4) (**Fig.2-3a**). The N-terminal 13 residues were not visible in the electron density map presumably due to its conformational flexibility. Therefore, an atomic model of Tm-FliP_P contained residues from Gly-123 to Lys-188, which correspond to Gly-122 and Gln-184

respectively in FliP_P of *Salmonella*. The asymmetric unit contained eight Tm-FliP_P molecules (**Fig.2-4a**). All chains in the asymmetric unit show no structural difference (**Fig.2-4b**). The most closely contact allows four chains to form a tetramer unit (**Fig.2-5a**). The electrostatic potential map and hydrophobicity map of the tetramer surface show that the surface is polarized and hydrophilic (**Fig.2-5b,c**). Since sedimentation equilibrium analytical ultracentrifugation measurements showed that Tm-FliP_P forms a tetramer in solution, I conclude that the tetramer structure observed in the crystal is equivalent to that in solution.

2.3.3. Disulfide cross-linking of FliP_P

Two two-fold axes cross orthogonally at the center of the tetramer structure (two-fold axis1 and axis2) (**Fig.2-6a**). As a result, only two of the four C-termini of the Tm-FliP_P chains of the tetramer can be connected to the cytoplasmic membrane region in the same orientation when the tetramer is put on the cytoplasmic membrane so as to connect the C-terminus of Tm-FliP_P directly to the cytoplasmic membrane. (**Fig.2-6b**). Therefore, I assumed that full-length FliP would form a dimer through the FliP_P-FliP_P interaction. There were two self-interaction interfaces in the tetramer structure (Interface1 and Interface2) (**Fig.2-6a**, **2-7**). In order to investigate which interface is actually exist in full-length FliP, I replaced pairs of amino acid residues, which are in close proximity to each other at the interfaces of Tm-FliP_P, with cysteine to performed *in vivo* disulfide cross-linking using full-length FliP of *Salmonella*. I selected eight pairs of residues for the tight interface and four residues for the loose interface that would be possible to form a disulfide bond when they were substituted with cysteine. A *Salmonella fliP* null mutant cells that express _{HA}FliP with a pair of cysteine substitutions from pUC19, were exponentially grown in LB media at 30 °C and then, disulfide

cross-linking was induced by adding iodine, followed by SDS-PAGE and finally western blotting with anti-HA antibody. In the case of the Interface1, cross-linking products were detected under an oxidizing condition but not at a reducing condition (**Fig2-8a**). The cross-linking product was about 50 kDa that would correspond to a dimer of $_{HA}$ FliP. Almost all pairs of the Cys variants produced the cross-linking products except for $_{HA}$ FliP(P133C/E178C). The motility of the $_{HA}$ FliP(P133C/E178C) and $_{HA}$ FliP(F137C/E178C) was not restored at all, indicating E178 is critical for for FliP function. Therefore, I suggest that full-length FliP forms a dimer through the Interface1. In contrast, any FliP protein expression was not observed at the Interface2 (**Fig2-8b**), raising the possibility that these residues is critical for the protein stability of FliP. Therefore, we cannot rule out the possibility that the interactions at the Interface2 contribute to efficient export gate formation and/or export activity.

2.3.4. Mutational characterization of FliP_P

Phe-138 and Phe-178 forms T-shaped benzene dimer at the interface1 of the *Thermotoga* FliP_P structure. Glu-142 and Glu-182 form a salt bridge with Arg-134. Met-127 and Val-131 make hydrophobic interactions with Leu-183, Ala-182 and Phe-187 at the edge of the interface1 (**Fig2-9**). To clarify where these interactions are physiologically important, I selected conserved amino acid residues among the above residues and then replaced each of them by alanine in *Salmonella* full-length FliP (**Fig2-10**). To evaluate the mutational impact on FliP function, I measured the motility of the *fliP* null mutants transformed with a plasmid encoding each of the FliP point mutant proteins in soft agar. FliP mutant variants were expressed from an arabinose-inducible promoter of the pBAD24 vector by adding 0.2 % arabinose. FliP(M138A) and FliP(Q141A) restored motility to a significant degree, FliP(F137A)

and FliP(Y174A) to some degree, and FliP(E178A) did not (**Fig2-11**). Next, I overexpressed FliP variants from the pUC19 vector. Immunoblotting with anti-HA antibody revealed that the expression levels of FliP(F137A) and FliP(Q141A) are almost the same as that of wild-type FliP and that the expression levels of FliP(M138A), FliP(Y174A) and FliP(E178A) were much lower than, the wild-type level. Except for FliP(E178A), almost all variants restored the motility to the wild-type level when they were expressed from the pUC19 vector (**Fig2-11**). Furthermore, all FliP variants did not show dominant negative effect at all (**Fig2-12**).



Figure 2-1. Purification profie of Tm-FliP_P.

(a) Purification of Tm-FliP_P by size exclusion chromatography using a Superdex75 column. Eluted protein in the peak fraction was conformed by SDS-PAGE. (b) Sedimentation equilibrium analysis of Tm-FliP_P. The initial protein concentration was 0.85 mg/ml. Open circles are data points, and the solid line is a model fit. Data points are fitted to a single species model with a molecular mass of 39.7 kDa. Measurements were done at 4 °C.



Figure 2-2. Crystals of native Tm-FliP_P and its X-ray diffraction pattern.

(a) Crystals of native Tm-FliP_P. Crystals obtained using 36% MPD, 0.1 *M* phosphate-citrate pH 4.4. (b) Diffraction image of the Tm-FliP_P crystal grown from the optimized condition. A typical X-ray diffraction pattern of the TmFliP_P crystal collected at SPring-8 beamline BL41XU. The inset shows a close up view of the diffraction image. (c) Bijvoet Difference Patterson map at w = 0.33 Harker section calculated from the Tm-FliP_p Se-Met derivative data at 3.5 Å resolution. The contour lines are drawn from 2.0 s to 6.0 s with an increment of 0.5 s.



Figure 2-3. Crystal structure of Tm-FliP_P

(a) The Tm-FliP_P monomer is shown in ribbon model, viewed from two different angles rotated 180° along the vertical axis. The N-terminal 13 amino acids were disordered. (b) Multiple sequence alignment of FliP_P. Uniprot accession numbers: *Salmonella* (P54700); *Thermotoga* (Q9WZG2); *Agrobacterium* (Q44344); *Aquifex* (Q67750); *Yersinia* (Q8D0A8); *Escherichia* (P0AC05); *Bacillus* (P35528); *Pseudomonas* (Q51468); *Vibrio* (A0A085SGD5).



Figure 2-4. Unit cell and crystal contact

(a) The asymmetric unit contains eight Tm-FliP_P chains. Each chain is shown in different colors. (b) Superimposition of all Tm-FliP_P in the asymmetric unit.



Figure 2-5. Tm-FliP_P tetramer in the crystal

(a) Ribbon model of the Tm-FliP_P tetramer structure, viewed from two different angles rotated 180° along the vertical axis. (b) Electrostatic potetial surface coloring of the tetramer, viewed from the same orientations with (a). (c) Hydrophobicity surface coloring of the tetramer, viewed from the same orientations with (a).



Figure 2-6. The symmetry axes in the tetramer.

(a) The two two-fold axes cross at the center of the tetramer. Two kinds of interfaces are present in the tetramer (red and blue). (b) The orientations of each C-terminus of Tm-FliP_P chains in the tetramer structure are shown as arrows. The tetramer structure is put on the cytoplasmic membrane so as to directly connect the C-terminus of Tm-FliP_P to the cytoplasmic membrane region. The two-fold axis1 and axis2 cross the cytoplasmic membrane orthogonally in the left and the right in the figure respectively.



Figure 2-7. The interfaces in the tetramer structure.

(a) The red surface indicates interface1 and the blue indicates interface2. (b) The dimer formation that forms interface1. (c) The dimer formation that forms interface2.



Figure 2-8. Disulfide cross-linking experiments

Results of disulfide cross-linking and the mutational effects of the cysteine replacement; (a) Interface1; (b)Interface2.

Mutational sites selected for cross-linking of *Salmonella* are mapped on the Tm-FliP_P structure. C_{β} of the residues of the variants are shown as sphere. Pairs of cross-linking are shown as red lines.



Figure 2-9. Dimer model of full-length FliP and the interactions at the interface1.

(a) Dimer model of fill-length FliP through the Interface1 in the cytoplasmic membrane.
(b) Interactions in the interface1. Close-up view of green square indicates T-stacking, blue indicates salt bridge and red indicates hydrophobic interaction. The contributing residues of *Thermotoga* (and corresponding residues of *Salmonella*) are mapped on the Tm-FliP_P structure. St : *Salmonella*



Figure 2-10. Evolutionary conserved residues of FliP_P.

The figure was prepared by ConSurf (<u>http://consurf.tau.ac.il/</u>). The mutational sites of *Thermotoga* (and corresponding residues of *Salmonella*) are mapped on the Tm-FliP_P structure. St : *Salmonella*



Figure.2-11. Effect of alanine substitutions in FliP_P on motility

(a) Relative motility and representative swarming motility of a *Salmonella* $\Delta fliP$ mutant transformed with pBAD- and pUC-based plasmids encoding alanine substitution variants of _{HA}FliP. WT, wild type FliP; V, pBAD24 or pUC19 (b) Expression level of wild-type FliP and its point variants in the $\Delta fliP$ null mutant transformed with the above plasmids as judged by immunoblotting with polyclonal anti-HA antibody.



Figure.2-12. Dominant negative effect of alanine substitutions in FliP_P

(a) Relative motility and representative swarming motility of wild-type cells transformed with pUC-based plasmids encoding alanine substitution variants of $_{HA}$ FliP. WT, wild type FliP; V, pUC19; (b) Expression of wild-type and variant FliP in wild-type cells transformed with the above plasmids.

2.4. Discussion

2.4.1. Relevance between the solubility and the structure of Tm-FliP_P

Salmonella FliP_P was expressed as an inclusion body whereas $_{His}Tm-FliP_P$ was expressed in a soluble fraction. One side of the molecular surface of Tm-FliP_P monomer or dimer is high hydrophobic. However, the hydrophobic surfaces face was covered in the tetramer and hence then the hydrophobic surfaces of Tm-FliP_P are isolated from water solvent. Therefore, I suppose that *Salmonella* FliP_P might not be able to form a tetramer in a way similar to Tm-FliP_P. This might be a reason why the protein solubility is quite different between FliP_P from *Salmonella* and *Thermotoga*.

_{His}Tm-FliP_P still tends to form highly aggregates during its purification. However, the removal of an N-terminal His-tag greatly improved its solubility. The N-terminal region of Tm-FliP_P has a flexible conformation. This is in agreement with the secondary structure prediction of FliP_P indicating that N-terminus of FliP_P would form loop or turn. Since a His-tag itself tends to aggregate in general, I assume that Tm-FliP_P forms a large aggregate through its N-terminus region with His-tag. Because alanine substitution of the P133 residue, which is located in the disordered N-terminus region of Tm-FliP_P, showed a significant reduced motility of *Salmonella* cells, I suggest that the conformational flexibility of the N-terminal region of FliP_P may have important roles in FliP function.

2.4.2. The role of the FliP_P-FliP_P interation

The role of FliP in flagellar protein export remains unknown. In this study, I found that the overexpression of FliP point mutant variants restores the motility to the wild-type level. Since these mutated residues are involved in the FliP_P-FliP_P interaction at the Interface1, over-expression of FliP mutant variants could increase the probability

of FliP dimer formation, thereby fully exerting their FliP function. In agreement with this, these FliP variants did not show a dominant negative effect on wild-type motility. Therefore, I propose that the FliP_P-FliP_P interaction at the interface 1 is critical for efficient export gate formation. Since the export gate complex is located in a highly limited space within the MS-ring, the number of FliP molecules in the export gate would not increase even if the FliP variants were over-expressed. Therefore, it is possible that the FliP_P-FliP_P interaction is not directly involved in the protein export process. However, over-expression of FliP(E178A) did not restore motility to the wild-type level although its cellular level was slightly higher than that of FliP(Y174A), of which over-expression conferred wild-type motility. This raises the possibility that a highly conserved Glu-178 residue is also involved in the protein export process.

2.4.3. Effect of each mutation to the FliP_P structure

Among the point mutnant variants, FliP(F137A) reduced the motility at almost same degree with FliP(Y174A) when they were expressed from the pBAD24 vector. These residues formed T-stacking at the Interface1, suggesting that the T-stacking interaction strongly contributes to form the Interface1. The aromatic part of Phe-178 (St: Tyr-174) forms a hydrophobic core with Met-139 (St : Met-138) for the folding of α -1 and α -2. Since both expressions of FliP(M138A) and FliP(Y174A) variants were quite lower than wild type FliP, the hydrophobic core formed by Phe-178 (St: Tyr-174) and Met-139 (St : Met-138) would be important to stabilize the FliP_P structure. Glu-184 (St : Glu-178) forms a salt bridge with Arg-134 (St : Pro-133) in the crystal structure for FliP_P-FliP_P interaction (**Fig.2-9, 2-13**). Since Arg-134 (St : Pro-133) is not conserved among the FliP homologues, this salt bridge contributes to the thermal stabilization of *Thermotoga* FliP_P. Multiple sequence alignments indicated that this

Arg-134 of Tm-FliP corresponds to the Pro residue among many species and to Pro-133 of *Salmonella* FliP (**Fig.2-14**). However, FliP(E178A) variant showed the most decreased motility among the variants. The secondary structure prediction showed that the region close to Pro-133 of *Salmonella* FliP is expected to form a small random coil domain (**Fig.2-14**). These results suggest that the region of Pro-133 of *Salmonella* FliP has flexibility and Glu-178 would play an important role at the flexible region.



Figure 2-13. Electrostatic potential map of the Interface1



Figure 2-14. Secondary structure prediction of FliP used for the multiple sequence alignment. h: helix, b: beta bridge, e: extended strand, t: beta turn, s: bend region, c: random coil. The prediction was carried out by SPOMA.

Chapter 3 FliP forms the hexagonal ring structure and FliO facilitates FliP ring formation

3.1. Introduction

In the previous chapter, I have suggested that FliP forms a homo-dimer though the FliP_P-FliP_P interaction to efficiently assemble into the export gate complex. It has been reported that that 4-5 copies of FliP molecule are identified in the purified FBB of *Salmonella* (Fan et al., 1997), raising the possibility that *Salmonella* FliP forms a dimer of dimer or a trimer of dimer in the export apparatus. To clarify it, I expressed and purifed *Salmonella* FliP. Although purified FliP was not mono-dispersed, EM observation revealed that FliP formed a ring-shaped oligomer.

Previous genetic analyses of a *fliO* null mutant have suggested that FliO interacts with FliP to regulate FliP function (REF). In this study, to clarify that FliO directly interacts with FliP to regulate FliP ring formation, I constructed the FliO/P co-expression system. By the purification from this co-expression system, I found that FliO directly interact with FliP to form FliO/P complex. Furthermore, the FliP ring dissociated from the FliO/P complex shows a high mono-dispersion. 2D-EM image classification reveals that the FliP ring is hexagonal. These results let me suggest that FliP forms a homo-hexamer and that FliO act as a scaffold for efficient FliP ring formation.

3.2. Materials and Methods

3.2.1. Bacterial strains, plasmids and media

The bacterial strains and plasmids used in the chapter are listed in Table 3-1 Compositions of media used in this chapter are essentially the same as those in chapter 2.

3.2.2. DNA manipulations

To construct a pTrc99A-based plasmid, a DNA fragment encoding *fliO*, *fliP* or *fliO-fliP* was amplified by PCR with chromosomal DNA of *Salmonella* as a template. The amplified DNA fragment was inserted into the *NdeI* and *Bam*HI sites of the pTcc99A vector. Mutations for biochemical characterization were generated by site-directed mutagenesis as described in chapter 2. Peptide insertions were performed by inverse-PCR method. All substitutions and insertions were confirmed by DNA sequencing as described in chapter 2.

3.2.3. Protein expression and purification

For expression and purification of FliP, the expression vectors encoding $_{\text{His}}$ FliP (pTF) were transformed into a *S.typhimurium flhDC* null strain ($\Delta flhDC$). The resulting transformants were grown in 2×YT medium containing 100 µg ml⁻¹ at 30 °C until OD₆₀₀ = 0.4-1.0 and then incubated at 16°C for another 24 h to express $_{\text{His}}$ FliP without any induction. Cells were harvested by centrifugation at 6,400 *g* for 10 min and stored at -80 °C. The cells were thawed, resuspended in a lysis buffer containing 20 mM Tris-HCl pH 8.0, 3 mM EDTA and sonicated. The cell lysates were centrifuged at 20,000*g* for 15 min to remove cell debris. The supernatants were ultracentrifuged at 110,000*g* for 1 hour to isolate the membrane fractions. Harvested membranes were

stored at -80 °C. For purification, the membranes were solubilized in the solubilization buffer containing 50 mM Tris-HCl pH 8.0, 300 mM NaCl, 5 % glycerol, 20 mM imidazole and 1% n-dodecyl b-D-maltoside (DDM) at 4 °C for 30 min and centrifuged at 110,000g for 30 min to remove the insoluble membranes. The solubilized membranes were loaded onto a Ni-NTA agarose resin and washed extensively with the wash buffer containing 50 mM Tris-HCl pH 8.0, 300 mM NaCl, 5 % glycerol, 20 mM imidazole and 0.1% DDM. Protein was then eluted with a 50–400 mM imidazole gradient and fractions containing _{His}FliP were collected. _{His}FliP was concentrated and then purified by size exclusion chromatography (Superdex 200 10/300) in a buffer containing 20 mM Tris-HCl pH 8.0, 150 mM NaCl, 2 mM EDTA, 5 % glycerol and 0.1% DDM.

For co-expression of FliO/P, the expression vectors encoding FliO-_{His}FliP (pTF) were transformed into the *Salmonella* $\Delta flhDC$ cells. The same expression and purification procedures were applied except for using a Superose 6 10/300 column for size exclusion chromatography purification. To obtain mono-dispersed FliP from the FliO/P complex, eluted fraction of FliO/P after Ni-NTA affinity purification was incubated at room temperature for 1day, resulting in the dissociation of FliP from the complex. Further size exclusion chromatography purification was carried out in the same way as FliP alone.

3.2.4. Motility assay

Fresh transformant colonies were inoculated onto soft tryptone agar plates as described in chapter 2.

3.2.5. Preparation of whole cell fractions and immunoblotting

Salmonella cells were exponentially grown at 30°C with shaking. Aliquots of

culture proteins containing a constant number of cells were clarified by centrifugation. Cell pellets were resuspended in SDS-loading buffer normalized by cell density to give a constant amount of cells. After sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE), immunoblot detection was done with an ECL immunoblotting detection kit (GE Healthcare).

3.2.6. Negatively stained EM observation and image processing

Samples were applied to carbon-coated copper grids and negatively stained with 1.0% (w/v) uranyl acetate. Micrographs were recorded at a magnification of \times 50,000 with a JEM-1011 transmission electron microscope (JEOL, Tokyo, Japan) operated at 100 kV. 4,663 particle images were selected with BOXER (Ludtke *et al.*, 1999) and the boxed particle images were aligned and classified using the REFINE2D.PY (Ludtke *et al.*, 1999).

Strains or Plasmids	Relevant properties	Source or reference
Salmonella		
SJW1368	Δ (cheW-flhD); master operon mutant	(Ohnishi et al.,1994)
	ΔfliO	
	ΔfliP	
Plasmids		
pTrc99AFF4	Modified trc expression vector	(Ohnishi et al.,1994)
pKY069	pTrc99AFF4/ _{His} FliP	This study
pKY070	pTrc99AFF4/FliO. _{His} FliP	This study
pKY071	pTrc99AFF4/FliO. _{His} FliP(F137A)	This study
pKY072	pTrc99AFF4/FliO. _{His} FliP(E178A)	This study
pKY073	pTrc99AFF4/FliO	This study
pKY074	pTrc99AFF4/ _{HA} FliP	This study
pKY075	pUC19/ _{HA} FliP	Chapter 2

Table 3-1. Strains and plasmids used in this study of this chapter

Primers	Sequence
FliO_NdeI.Fw	gggaattccatATGATGAAGACAGAAGCCAC
FliO_BamHI.Re	cgcggatccTCAGGATCTCCCGGAACGCTTG
FliP_BamHI.Re	cgcggatccCTAACTGTAAAAGCTTTGGG
SigPep_QLHis_FliP.Fw	GCCGCCGCTGCGCAACTTcatcatcaccatcaccacCTGCCGGGGGCTTATCAGC
SigPep_QLHis_FliP.Re	GCTGATAAGCCCCGGCAGgtggtggtggtggtggtggtggtgatgatgAAGTTGCGCAGCGGCGGC
FliP_F137A.Fw	GCGCAACCGTTACGCGCGgcCATGCTGCGCCAAACCCGC
FliP_F137A.Re	GCGGGTTTGGCGCAGCATGgcCGCGCGTAACGGTTGCGC
FliP_E178A.Fw	CCCGCTTATGTCACCAGCGcATTAAAGACGGCGTTTCAG
FliP_E178A.Re	CTGAAACGCCGTCTTTAATgCGCTGGTGACATAAGCGGG

Table 3-2. Primers used in this study of this chapter

3.3. Result

3.3.1. Expression for full-length FliP of Salmonella

FliP can be expressed from a *ptac* promoter of the pTrc99A vector in Salmonella even in the absence of IPTG. The rate of the expression of FliP was quite slow and so the growth arrest did not occur. The expression level of FliP was detected by immunoblotting but not by CBB staining but by from the cell lysate (Fig. 3-1a). Therefore, I decided to attach a tag such as a His-tag and a HS-tag to the N and C-terminus of mature FliP for affinity chromatography. However, most of peptide tags dramatically reduced the FliP expression level except for an N-terminal HA tag. Therefore, I tried to modify the FliP construct by optimizing tags, signal peptides required for efficient FliP insertion into the cytoplasmic membrane, insertion positions of a tag and their combinations (Fig. 3-1b). All constructs that have a C-terminal tag destabilized FliP whereas some constructs that have a N-terminal hexa-histidine tag were stable. Next, I tested whether these constructs were functional. I checked motility of a *fliP* null mutant that expresses each FliP recombinant in soft agar. Almost all constructs that have a chimeric signal peptide showed high toxicity to *Salmonella* cells. A FliP construct, which has a hexa-histidine tag with a N-terminal single leuceine (LHHHHHH) located between Q22 and L23, was both functional and stable, and the amino acid composition was the simplest among all of the constructs (Fig. 3-1c). I used this construct as _{His}FliP for the following experiments.

3.3.2. Purification and EM observation of FliP full length

Total isolated membranes were solubilized by DDM. I carried out Ni-NTA affinity chromatography, followed by size exclusion chromatography. Purified FliP was not mono-dispersed and the highest molecular weight of eluted FliP was estimated to be
about 200 kDa as judged by size exclusion chromatography (**Fig. 3-2a**). Since purified membrane proteins are embedded in detergent micelles, they generally behave as up to two times larger than their molecular weight. Therefore, purified FliP oligomer was speculated to contain 4 to 8 copies of FliP monomer.

Next, I observed each major peak fraction of FliP (peak1 and peak2) by negatively stained electron microscope in order to visualize the FliP shape. In the peak1 fraction, I clearly observed the ring-shape structure (**Fig.3-2b**). However, the amounts of the ring structures were much lower in the peak2 fraction than in the peak 1 fraction and quite smaller fragments were seen in the peak2 fraction (**Fig.3-2b**). These results indicated that FliP forms an oligomeric ring structure and that there were several oligomeric states of FliP in the purified sample.

3.3.3. Co-expression, purification and EM observation of FliO/P

To succeed in both functional and structural analysis, the purification of a mono-dispersed protein is quite important. In general, it is necessary to carry out detergent screening to obtain mono-dispersed membrane proteins. I examined DDM, DM (n-decyl-β-D- maltopyranoside), SL1 (sucrose monolaurate), SMc (sucrose monocaprate), OG (n-octyl $-\beta$ -glucoside), CHAPS (3-[(3-Cholamidopropyl)dimethylammonio]-1-propane- sulfonate), LMNG (lauryl maltose neopentyl glycol) and DMNG (decyl maltose neopentyl glycol) for the first screenig. Only DDM and SMI made FliP dispersion but not mono-dispersion. Other detergents made FliP insoluble or soluble aggregation (void fraction in size exclusion chromatography). It is also known that some proteins are more stable when they form a complex with other components. FliP was assumed to interact with FlO, FliR and FlhA (REFS). Especially, recent genetic study indicated that FliO is not an essential

component for flagellar construction among the export gate components, and bypass mutations, which allows a *fliO* null mutant to form flagella to a considerable degree, are identified in the *fliP* gene (Barker et al., 2010; Barker et al., 2014). Furthermore, the cellular level of FliP is increased when FliP is co-expressed with FliO. Therefore, FliO is assumed to interact with FliP in direct via its transmembrane region to stabilize FliP in the cytoplasmic membrane. To test this, I co-expressed and purified FliP with FliO. I expressed FliO and FliP as an operon from the ptrac promoter of the pTrc99A vector because antibiotics except for ampicillin dramatically decreased both the FliO and FliP expression levels. I took the same purification method and condition for the FliO/P complex as FliP alone and so succeeded in the purification of the FliO/P complex. It has been reported that FliO is expressed as two forms, a long form and a short form (REF). Interestingly, both two forms of FliO were co-purified with FliP (Fig.3-3a). Although purified FliO/P complex seemed to be mono-dispersed in size exclusion chromatography, EM observation of the complex revealed that a couple of the FliP rings connect with each other presumably through an interaction between the FliP ring and FliO (Fig.3-3b). Surprisingly, I was also able to obtain the highly mono-dispersed and homogenous FliP rings from the FliO/P complexes rather compared to FliP alone (Fig.3-3a). These results suggest that FliO directly interacts with FliP to facilitate FliP ring formation.

We performed 2D classification image analysis of the FliP ring isolated from the FliO/P complexes. 4663 particles were picked up and classified into 100 classes. Many of the classes revealed the hexagonal ring with a diameter of 2nm (**Fig.3-4**).

3.3.4. Purification and EM observation of FliP variants

To test if the FliP_P-FliP_P interaction is required for FliP ring formation, I

decided to express and purify FliP(F137A), FliP(Y174A) and FliP(E178A) because these FliP mutations reduced motility (**Fig. xx**). I found that the expression levels of FliP(Y174A) and FliP(E178A) are lower than the wild-type level. Therefore, I constructed plasmids co-expressing these FliP point mutant variants with FliO because the steady cellular level of FliP is increased considerably by its co-expression with FliO. Unfortunately, I could not obtain a plasmid encording FliO and FliP(Y174A) on the pTrc99A vector presumably because high-expression of the FliO/P(Y174A) complex confers toxicity to the host cells. Therefore, I expressed and purified the FliO/P(F137A) and FliO/P(E178A) complexes in the same way with wild-type FliO/P. Both FliP(F137A) and FliP(E178A) co-purified with FliO. However size exclusion chromatography revealed that the molecular weight of each FliP variants were smaller than wild-type FliP (**Fig.3-5a**). Furthermore, FliP(E178A) dissociated from FliO much easier than wild-type FliP and FliP(F137A), indicating that the E178A mutation reduces the binding affinity of FliP for FliO. Therefore, I suggest that FliP_P contributes to a stable association of FliP with FliO.

Next, I observed purified FliP variants by negatively stained EM. Compared to wild-type FliP, the FliP(F137A) and FliP(E178A) rings were not observed efficiently compared to wild-type FliP although their much small particles were seen. These results suggest that the F137A and E178A mutations significantly reduces the probability of FliP ring formation (**Fig.3-5b**). Therefore, I suggest that the FliP_P-FliP_P interaction is required for efficient FliP ring formation.



Figure 3-1. Expression of FliP full length

а



Size exclusion chromatography (Superdex200)



(a) Size exclusion chromatography (Superdex200) of FliP. Eluted protein in each peak fraction was confirmed by CBB stain. (b) Representative negatively stained EM images of peak1 and peak2. Scale bar shows 50 nm.





Figure 3-3. Purification and EM-observation of FliO/P

(a) Size exclusion chromatography (Superose6) of FliO/P. Eluted protein in each peak fraction was confirmed by CBB stain. (b) Representative negatively stained EM images of peak1 and peak2. Scale bar shows 50 nm.



b

а



Figure 3-4. 2D classification

(a) Enlarged views of 10 representative 2D class averages show hexagonal ring structures. Scale bar shows 2 nm. (c) Auto-correlation analysis of the 2D class averaged image of the FliP ring.

Size exclusion chromatography (Superose6) Standard molecular weight marker (kDa) Void 670 158 44 Т 1 50.0 40.0 CBB stain OD₂₈₀ (mAU) (kDa 30.0 25 17 FIO 20.0 25 17 -FIIO 10.0 25 -17 -FIIO 0 20.0 8.0 12.0 16.0 Elution volume (ml)

FIIP

"FIIP (F137A)

_{sa}FliP (E178A)



Figure 3-5. Purification of FliP variants

(a) Size exclusion chromatography profiles of FliO/P (black line), FliO/P(F137A) (green line) and FliO/P(E178A) (blue line). Eluted proteins were confirmed by SDS-PAGE. (b) Representative negatively stained EM images of the FliP fraction from each variant. Scale bar shows 50 nm.

3.4. Discussion Biological unit of FliP and role of FliO

I showed that FliO directly interacts with FliP to facilitate FliP hexamer formation. I also found that a couple of the FliP rings connect with each other in the purified FliO/FliP complex. Interestingly, the FliP₆ ring dissociates from the FliO/P complex during size exclusion chromatography. These observations suggest that FliO acts as a scaffold to form FliP ring. Recent genetic study have shown that FliO dose not contribute to FliP protein stability because the bypass mutations isolated from the *fliO* null strain, FliP(R143H) and FliP(F170L), do not increase the protein stability of FliP, significantly. In contrast, I found that the co-expression of FliP with FliO increases the expression level of FliP, significantly. In agreement with this, FliO also stabilized point mutant variants of FliP. These results indicate that FliO has a role in stabilization of FliP *in vivo*. Therefore, I suggest that the bypass FliP (R143H) and FliP(F170L) mutations could increase the probability of FliP ring formation in the absence of FliO.

As discussed in chapter 2, the FliP_P-FliP_P interaction is required for FliP dimer formation. Here, I showed that both FliP(F137A) and FliP(E178A) interact with FliO. However, they did not form the FliP ring structure, indicating that the FliP_P-FliP_P interaction is also required for FliP ring formation. Furthermore, FliO easily dissociated from FliP(E178A) during size exclusion chromatography, indicating that the FliP_P-FliP_P interaction stabilizes the FliO-FliP interaction. Taken all together, I suggest that the FliP₆ ring is a timer of dimer. Because FliP(E178A) did not restore motility to a considerable degree even under high expression condition, I propose that the FliP₆ ring is a functional unit in the export apparatus. This is in agreement with the previous observation that about 4 to 5 molecules of FliP associate with the FBB. The central channel of the growing flagellar structure is a physical path for flagellar protein. Interestingly, the diameter of the central channel of the flagellar structure is only 2 nm, which is the same as that of the central pore of the $FliP_6$ ring. This raises the possibility that FliP acts as a peptide channel for export substrates. We need further experiments to confirm this hypothesis.



Fig 3-8 Assembly model of FliP

FliP self-assembles into a homo-dimer through the $FliP_P$ -FliP_P interaction in the cytoplasmic membrane. The hydrophobic interaction of F137-F137 plays an important role in this process. Although a counterpart of E178 for salt-bridge formation is not conserved, E178 is directly involved in the dimerization. The FliP dimer forms a timer of dimer with the help of FliO, which acts as scaffold for FliP₆ ring formation.

Chapter 4 Cryo-EM study of the FliP ring, and the construction of the export gate complex

4.1. Introduction

In chapter 3, I have shown that the FliP ring has a central pore with a diameter of 2 nm. This result gives rise to a new question of whether the FliP ring is a channel for export substrates. To clarify whether FliP interacts with export substrates, I would like to carry out *in vivo* photo-crosslinking experiments. I have to do two things conduct this experiment. One is that I have to obtain a high-resolution 3D structure of the FliP ring. That would be useful for me not only to make design for the photo-crosslinking experiment but also to discuss the obtained results more accurately based on the high-resolution structure. The other thing is that I have to develop the overexpression system of the entire export gate complex. So far, FliO, FliP, FliQ and FliR expressed from chromosmal DNA are unable to detected by western blotting. Overexpression of FliP alone is not sufficient since the export gate works as the complex. Furthermore, as I mentioned in chapter 1, to investigate the molecular mechanism how the export gate converts PMF to the mechanical work required for flagellar protein export in the future, the isolation of the complete export gate complex must be needed. Therefore, the overexpression system of the export gate complex must be developed.

In this chapter, I will show preliminary high-resolution structural analysis of

the FliP ring by cryo-EM and single particle image analysis. The FliP ring particles, of which detergent micelles are replaced by amphipol, are well mono-dispersed in the thin ice formed on the carbon grid. 2D-EM image classification shows a clear hexagonal ring shape, indicating structural analysis of FliP by single particle analysis could be possible. I will also show the construction of the export gate complex for the future research. All components of the export gate are sufficiently expressed. From this co-expression system, I can purify the FlhA/FliF/FliG complex.

4.2. Materials and Methods

4.2.1. Bacterial strains, plasmids and media

The bacterial strains and plasmids used in this chapter are listed in Table 4-1

4.2.2. DNA manipulation

The *Cla*I, *Eag*I and *Spe*I sites that are placed at 30 bp intervals were inserted into the pTc99A vector by invers PCR method, generating a plasmid named pTrc_CES01. To construct pTrc_SE01, the *Spe*I and *Eag*I sites were inserted into the 30bp upstream region of trc promoter and 30bp downstream region of rrnb terminator of pTrc99A, respectively by the Quick Change method. To construct pUC_CE01, a DNA fragment that contains both *Nde*I and *Bam*HI sites were removed from pUC19 by inverse PCR method. Then, the *Cla*I and *Eag*I sites were inserted by inverse PCR method, and a DNA fragment that contains a trc promoter, a multiple of cloning sites and a rrnb terminator derived from the pTrc99A vector was amplified by PCR and then inserted into the *ClaII-EagI* sites of the above vector.

4.2.3. Motility assay

Fresh transformant colonies were inoculated onto soft tryptone agar plates.

4.2.4. Preparation of whole cell fractions and immunoblotting

Salmonella cells were grown at 30°C with shaking exponentially. Aliquots of culture proteins containing a constant number of cells were clarified by centrifugation. Cell pellets were resuspended in SDS-loading buffer normalized by cell density to give a constant amount of cells. After sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE), immunoblot detection was done with an ECL immunoblotting detection kit (GE Healthcare).

4.2.5. Protein expression and purification

The FliP ring was purified in the same method as described in chapter 3. After size exclusion chromatography, total proteins were concentrated to 1 mg ml⁻¹, and supplemented with amphipol A8-35 (Anatrace) to a final concentration of 3 mg ml⁻¹. After incubation at 4 °C for 4 h, 50 mg Bio-Beads SM-2 (Bio-Rad) was added and then incubated for 6 h. Free amphipol molecules were removed by size exclusion chromatography. Peak fractions were collected for cryo-EM studies.

The export gate complex was expressed and purified in the same method as described in chapter 3.

4.2.6. Data collection and image processing

Aliquots of 2.1 µl of purified FliP at a concentration of approximately 2 mg ml⁻¹ were placed on glow-discharged holey carbon grids (Quantifoil Cu R1.2/1.3). Grids were blotted for 7 s and flash frozen in liquid ethane using an FEI Vitrobot. Grids were transferred to JEM-3200FSC electron microscope that was operating at 300 kV. Images were recorded manually using a K2 Summit detector at a super-resolution mode on the JEM-3200FSC microscope at a calibrated magnification of 40,000 (yielding a pixel size of 0.9798 Å²). The K2 detector was mounted after a Gatan Imaging Filter (GIF), but the filter was not used to remove any inelastic scattering. A dose rate of approximately 16 electrons per Å² per s and exposure time of 12 s on the K2. Defocus values in the final K2 data set ranged from 1.5–3.5 mm, and 40 video frames were recorded.

RELION1.3 was used for automated selection of 37,655 particles from 112 mmicrographs. Contrast transfer function parameters were estimated using CTFFIND3

(Mindell J.A. and Grigorieff N, 2003) and then reference-free two-dimensional class averaging was carried out.

Strains or Plasmids	Relevant properties	Source or reference		
Salmonella				
SJW1368	Δ (cheW-flhD); master operon mutant	(Ohnishi et al.,1994)		
	ΔfliP			
Plasmids				
pTrc99AFF4	Modified trc expression vector	(Ohnishi et al.,1994)		
pTrc_ES01	Modified trc expression vector	This study		
pTrc_CES01	Modified trc expression vector	This study		
pUC_CE01	Modified cloning vector	This study		
pKY076	pTrc99AFF4/FliO. _{His} FliP	Chapter 3		
pKY077	pTrc99AFF4/FliO. _{His} FliP. _{HA} FliQ.FliR _{FLAG}	This study		
pKY078	pTrc_CES01	This study		
	$/FlhB.FlhA_FliO{His}FliP{HA}FliQ.FliR_{FLAG}$			
pKY079	pTrc_CES01	This study		
	$/FlhB.FlhA_FliO.FliP_{\cdot HA}FliQ.FliR_{FLAG}_FliF.FliG_{His}$			

Table 4-1. Strains and plasmids used in this study of this chapter.

Primers	Sequence
CES3.Fw	TcggccgAGTAAATAACCCCTTGCGGATCTATGCTAGactagtcggttctggcaaatattc
CES3.Re	CTGCCGCCCAGctgctcgaagcggccACatcgatttatgatgtcg
E_pTrc99A.Fw	gttttttgcgccgacatcataacggccgcggttctggcaaatattc
E_pTrc99A.Re	gaatatttgccagaaccgcggccgttatgatgtcggcgcaaaaaac
pBAD24_S.Fw	tgtttatttttctaaatacaactagtttcaaatatgtatccgctc
pBAD24_S.Re	gagcggatacatatttgaaaactagttgtatttagaaaaataaaca
C_pTrc99A.Fw	gtttttgcgccgacatcataaatcgataggttctggcaaatattc
C_pTrc99A.Re	gaatatttgccagaacctatcgatttatgatgtcggcgcaaaaaac
pTrc99A_E.Fw	tgtttatttttctaaatacacggccgttcaaatatgtatccgctc
pTrc99A_E.Re	gagcggatacatatttgaacggccgtgtatttagaaaaataaaca
trcP_EagI.Fw	tccggccgcggttctggcaaatattctg
rrnB_SpeI.Re	ggactagttgtatttagaaaaataaacaaaaag
trcP_ClaI.Fw	tcataaatcgataggttctggcaaatattctg
rrnB_SpeI.Re	tccggccgtgtatttagaaaaataaacaaaaag
pUC_MOD.Fw	atcgatgtcggccgagactagttttcttagacgtcaggtggcac
pUC_MOD.Re	tetteegetteetegeteaetgae
pUC_MOD2.Fw	TcggccgAGTAAATAACCCCTTGCGGATCTATGCTAGactagttttcttagacgtcaggtggc
pUC_MOD2.Re	CTGCCGCCCAGetgetegaageggeeACategattetteegetteetegeteae
NdeI_FlhB.Fw	gggaattccatATGGCAGAAGAGAGCGACGAC
FlhA_BamI.Re	egeggateeTTATTTTCCTCCAATGGTC
FliO_NdeI.Fw	gggaattccatATGATGAAGACAGAAGCCAC
FliR_FLAG_BamHI.Re	cgcggatccTTActtatcgtcgtcatccttgtaatcTGGGTTATTATTTATCGGCATC
FliF_NdeI.Fw	gggaattccatATGAGTGCGACTGCATCGAC
	cgcggatccTTAGTGATGATGATGATGATGATGATGATGATGATGATGA
FliG_His10_BamHI.Re	ATCCTCGCCG

Table 4-2. Primers used in this study of this chapter

4.3. Result

4.3.1. Purification and observation of the FliPring by cryo-EM

To succeed in high-resolution structural analysis by cryo-EM and single particle analysis, it is important to embed biomolecule particles into the thin ice on the grid to obtain their high contrast images. It has been reported that detergents in the protein solution make it difficult. To remove DDM from protein solution, I replaced DDM by amphipol. The membrane proteins with amphipol are known to behave larger size than those embedded in detergent micelles in general. The size exclusion chromatography indicated that the FliP ring with amphipol also behaved larger size as well, and was well mono-dispersed as well as the FliP ring with DDM micelle (**Fig4-1**). The FliP ring particles were also mono-dispersed in a thin layer of vitreous ice formed in the carbon holes, indicating that the FliP rings are not absorbed to the carbon grid and that the rings are still in a stable state in vitreous ice (**Fig.4-2a**).

4.3.2. Construction for the expression system of the entire export gate complex

To confirm the expression level of each export gate components, it must be necessary to detect each component by immunobloting. Among the export gate components, we did not have polyclonal anti-bodies specific for FliQ and FliR. Therefore, I fused a HA-tag to the N-terminus of FliQ and a FLAG-tag to the C-terminus of FliR. These tags did not affect the export function of these two proteins at all. It has been reported that the export gate is proposed to assemble in a coordinated manner during MS ring formation (Morimoto et al., 2014). A C-ring component protein FliG is required for efficient MS ring formation in vivo (Morimoto et al., 2014). Therefore, FlhA, FlhB, FliO, FliP, FliQ, FliR, FliF and FliG must be co-expressed to obtain the complete export gate complex associated with the MS ring. To simplify the

expression system, I expressed these eight proteins from a pTrc99A-based plasmid in a *Salmonella* $\Delta flhDC$ strain as a host. *flhA* and *flhB*, *fliO*, *fliP*, *fliQ* and *fliM*, *fliF* and *fliG* are transcribed as the *flhB*, *fliL* and *fliF* operons, respectively in *Salmonella* (Kutsukake *et al.*, 1988). Three PCR products, *flhBA*, *fliOPQR*, and *fliFG*, were cloned into three distinct sites of the pTrc99A_CES01 vector. They were placed from a p*trc* promoter (**Fig.4-3**). I also constructed the FliO/P/Q/R and FlhB/A-FliO/P/Q/R, FliF/G co-expression system respectively.

As mentioned in Chapter 3, I attach a His-tag to FliP and so purification of the FliO/P/Q/R and FlhB/A-FliO/P/Q/R complexes was carried out by Ni-NTA affinity chromatography, followed by size exclusion chromatography (Fig.4-4a). The FliO/P/R complex was purified by Ni-ATA affinity chromatography from FliO/P/Q/R co-expression system and FliO was dissociated from this complex during gel filtration chromatography. The FliP/R complex showed a mono-dispersed peak. The FliO/P/R structure looked similar to the FliO/P complex (Fig.4-4b). The FliO/P/R-FlhB complex was affinity purified from the FlhB/A-FliO/P/Q/R co-expression system, indicating that FlhB directly binds to the FliO/P/R complex whereas FlhA does not. The molar ratio of FliR and FlhB seemed to be 1:1, in agreement with a previous observation that a FliR-FlhB fusion is functional (REF). However, FlhB was easily dissociated from this complex during size exclusion chromatography, indicating the interaction between FliO/P/R and FlhB could be weak. For FlhB/A-FliO/P/Q/R-FliF/G co-expression system, I fused a deca-histidine tag at the C-terminus of FliG and then the FlhA/FliF/FliG complex was able to be purified by Ni-NTA affinity chromatography, followed gel filtration chromatography (Fig.4-4a). This indicates that FlhB/FliO/P/Q/R easily dissociates from the FlhA/FliF/FliG complex during solubilization and puritication. Negatively stained EM observation of this complex revealed the MS-ring

structure, indicating that FlhA and FliG associate with the MS ring (Fig.4-4b). The stoichiometry of the FlhA/FliF/FliG complex seemed to be ~ 0.3 FlhA : 1 FliF : ~ 1.5 FliG.





(a) Size exclusion chromatography (Superdex200) after exchanging from DDM to amphipol. Eluted proteins in the peak was confirmed by SDS-PAGE. (b) Representative negatively stained EM images of purified FliP in DDM and in amphipol. Scale bar shows 50 nm.



b

4	۲	۲	۲	۲	۲	۲	۲	۲	۲
۲	۲	۲	۲	۲	۲	۲	۲	۲	۲
۲	۲	۲	۲	۲	۲	۲	-	۲	۲

Figure 4-2. Cryo-EM images and 2-D classification

(a) Representative EM image of FliPap embedded in a thin layer of vitreous ice. Fourier transform of micrograph shown in right, with Thon rings extending beyond 4 Å. (b) Representative 2D class averages.





(a) Schematic diagram of co-expression plasmids of the export gate. (b) Westernblotting of membrane fraction by antibody to each export gate component and westernblotting of whole cell fraction by antibody to FliF and FliG.



Figure 4-4. Purification and EM-observation

(a) SDS-PAGE results after Ni-NTA affinity chromatography from each co-expression system. (b) Representative negatively stained EM images FliP and complexes purified after size exclusion chromatography. Scale bar shows 50 nm.

4.4. Discussion

2D class averages of negatively stained FliP ring showed that the FliP ring is a hexagonal ring structure. Some 2D class averages of the FliP ring from cryo-EM data also showed a hexagonal ring structure (**Fig.4-2b**), indicating that the FliP ring is not distorted by negative staining. Therefore, I conclude that the FliP₆ ring is a physiological and functional relevant in the export apparatus. Since the biological unit of FliP_P is proposed to be a dimer, the FliP ring is assumed to form a trimer of dimer. Indeed, many classes of the FliP ring showed three-fold symmetric structure.

The previous genetic and biochemical study of the flagellar homologue, the needle complex, revealed that an earlier assembly of SpaP, SpaQ and SpaR (correspond to FliP, FliQ, FliR respectively) is required for the assembly of InvA and SpaS (correspond to FlhA and FlhB) into the export gate. In addition, InvA and SpaS can assemble into the export gate after completion of the assembly of the needle complex base whereas neither SpaP, SpaQ nor SpaR can. These result indicated that in flagellar structure, FliO, FliP, FliQ and FliR forms a core complex for the following assembly of FlhA and FlhB into the export gate. Since SpaP and SpaR seemed to form a complex in the export gate and their assembly is required for the assembly of SpaQ into the export gate, SpaP and SpaR forms stable complex at the earliest stage of the export gate assembly. This is in agreement with my result that FliP and FliR forms a stable complex. Although the FliP/R complex was isolated from the FliO/P/R complex, it is still unclear that the FliO/P complex structure is required for efficient interaction of FliP with FliR. FliQ was assumed to form a complex with the FliO/P/R complex, however FliQ dissociated form the FliO/P/R complex during Ni-NTA affinity chromatography, indicating that an association between FliQ and the FliO/P/R complex could be weak .

Another previous genetic study showed that fusion protein of FlhB and FliR complemented their function, indicating FlhB and FliR interact with each other at a 1 to 1 molar ratio. In this study, co-expression and purification of FlhA/FlhB -FliO/P/Q/R revealed that FlhB interacts with FliO/P/R. Furthermore the molar ratio of FlhB and FliR seemed to be 1:1. Therefore FlhB was assumed to be associated with FliO/P/R complex through the transmembrane region of FliR at a 1:1 molar ratio. However, still FliQ and FlhA were dissociated from the complex.

When FlhB, FlhA, FliO/P/Q/R and FliF/G were expressed at the same time, I succeeded in purification of the FlhA/FliF/FliG complex by Ni-Affinity chormatography, followed by size exclusion chromatography. This result was consistent with the recent genetic and fluorescent microscopic analysis that the MS-ring component, FliF, and one of the C-ring components, FliG, are required for FlhA assembly into the export gate. Co-expression and affinity purification of by C-terminal His-tag of FliG resulted in the purification of FlhA-FliF/G complex. From the result of CBB stain, the molecular ratio of FliF and FliG seemed to be 1 : 1.5. It has been reported that FliF/FliG complex was purified from FliF/G co-expression system and the molecular ratio of FliF and FliG is 1 : 1. Therefore FlhA in the MS-ring might affect the conformation of FliF or FliG to change the interaction manner between the MS-ring and FliG. Other components except for FliO and FlhB were detected by immunoblotting, however almost all components were appeared in the flow through fractions, indicating those components were dissociated form the complex when they were solubilized from the membrane or the complex flew through from the affinity resin.

To clarify the detail assemble mechanism of the export gate, it is necessary to express any combinations of the export gate components, followed by purification and observation. Furthermore to confirm these biochemical data, it is important to observe the localization and dynamics of each component by fluorescent microscope in the living cells. Moreover to isolate the complete export gate complex for understanding the molecular mechanism of the protein translocation, it is still necessary to examine the solubilization conditions from the membrane and the purification conditions.

Chapter 5 Conclusion

The PMF-driven export gate complex is located at the flagellar base to drive flagellar protein export for the construction of the flagellum beyond the cell membranes (Katayama et al. 1996; Kawamoto et al. 2013). It has been reported FliP and FliR are incorporated into the basal body along with FliF and FliG at the earliest stage (Jones and Macnab, 1990; Fan et al., 1997). Since FlhA assembles into the export gate complex along with other transmembrane export gate proteins during MS ring formation, it has been proposed that the assembly of the export gate complex may begin with FliO, FliP, FliQ and FliR, followed by FlhA and FlhB (Morimoto et al. 2014). However, it remains unknown how FliO, FliP, FliQ and FliR forms a core structure of the export gate for assembly of FlhA and FlhB. Therefore, I forced on the structure of FliP to understand the assembly mechanism of the export gate complex. Because FliP has a relatively large periplasmic domain, I first determined the structure of a periplasmic domain of FliP_P from *Thermotoga maritima* at 2.4 Å resolution. Structure-based in vivo disulfide cross-linking experiment, revealed that FliP form a homo-dimer through the FliP_P-FliP_P interaction. Highly conserved Phe-137 and Glu-178 residue located at the interface 1 contribute to the strong FliP_P-FliP_P interaction in the crystal structure. In agreement with this, the FliP(F137A) and FliP(E178A) substitutions results in a significant reduction in motility. These results suggest that FliP forms a

dimer unit through the $FliP_P$ - $FliP_P$ interaction for efficient assembly of the export gate complex.

It has been reported that about 4 or 5 copies of FliP associates with the basal body MS ring (Fan et al., 1997). To test whether if full-length FliP forms a higher oligomer in the MS ring, I overexpressed and purified full-length FliP. Although FliP did not show a mono-dispersed peak during size exclusion chromatography even though several detergents were examined for its solubilization, co-expression of FliP with FliO dramatically increased its homogeneity. FliO was co-purified with FliP, indicating FliO and FliP directly interact with each other. EM observation of FliP revealed that FliP forms a ring-shaped structure. 2D class averages of the FliP ring revealed the hexagonal ring structures, indicating FliP forms a homo-hexamer. EM observation of the FliO/P complex revealed that a couple of the FliP rings associate with each other through an interaction between FliO and the FliP ring. The FliP variants harboring a mutation that affects the FliP_P-FliP_P interaction was unable to form the stable FliP ring structure, Based on these results, I propose that the FliP form a hexamer as a trimer of dimer, and that FliO act as a scaffold to facilitate FliP ring formation. Since the diameter of the MS-ring is estimated to be around 40 nm, it is difficult to insert the FliO/P complex into the MS-ring. On the bases of the result that FliP and FliP/R easily dissociate form FliO, and the fact that FliO is indispensable among the export gate components, I assume that FliO may not localize in the MS-ring.

Co-expressions of FliO/P/Q/R and FlhB/A-FliO/P/Q/R allowed me to purify the FliO/P/R and FliO/P/R-FlhB complexes, respectively. The stoichiometry of FlhB and FliR seemed to be 1:1 that is consistent with the previous genetic analysis. Although FlhB was easily dissociated from the FliO/P/R complex during gel filtration chromatography, indicating that an association between FlhB and the FliO/P/R complex is weak. EM observation of the FliO/P/R complex revealed that its overall structure was almost the same as the FliO/P complex. The FliP/R complex was dissociated form the FliO/P/R complex during gel filtration chromatography in a way similar to FliP dissociated from the FliO/P complex, and its structure was also almost identical to the FliP ring. This suggests that FliR directly binds to the FliP ring. Co-expression of FlhB/A-FliO/P/Q/R-FliF/G allowed me to isolate the FlhA-FliF/G complex by gel filtration chromatography, suggesting that the FlhB/FliO/P/R and FliQ dissociates from the FlhA-FliF/G complex during solubilization , followed by Ni affinity chromatography. EM observation of the FlhA-FliF/G complex revealed that FliF forms the MS-ring and that FlhA and FliG associates with the MS ring . Although the stoichiometry of FliF and FliG is proposed to be 1:1, the stoichiometry of FliF and FliG in FlhA-FliF/G complex seemed to be 1:~1.5, indicating that the association of FlhA with the MS-ring might affect the interaction between FliF and FliG.

FliJ binds to the FliI₆ ring to form the FliI₆FliJ ring complex (Ibuki *et al.*, 2011). ATP hydrolysis by the FliI₆FliJ ring complex activates the export gate complex through an interaction between FliJ and FlhA, allowing the gate to efficiently utilize PMF across the cytoplasmic membrane for rapid and efficient flagellar protein export (Minamino et al., 2014). The export gate complex is a proton-protein antipoter to couples the proton flow with protein translocation (Minamino *et al.*, 2011). FlhA is an energy transducer to conduct both H^+ and Na^+ (Minamino *et al.*, unpublished data). ATP hydrolysis by FliI ATPase is coupled to proton translocation through the export gate (Morimoto *et al.*, unpublished data). Since the FliP ring seems to be a core structure for export gate assembly, the FliP ring could be located at the center of the export gate complex. The central pore of the FliP ring is 2 nm, which is the same size as the central channel of the growing flagellar structure, I propose that the FliP ring acts as

a peptide channel and that ATP hydrolysis by FliI ATPase induces proton translocation through the FlhA proton channel, allowing export substrates to go into the FliP channel and then the central channel of the growing flagellar structure. To understand the detail molecular mechanism of PMF-driven protein transport by the export gate complex, further biochemical and high-resolution structural analysis must be necessary.

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