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Tunnel Formation Inferred from the I-Form Structures of the Proton-Driven Protein Secretion Motor SecDF

Graphical Abstract

Highlights

- Crystal structures of SecDF in I forms are described
- The structures provide insight into an energy-coupling mechanism for Sec machinery
- The structures suggest a substrate binding site and a pathway for protons

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In Brief

SecDF, a motor protein, uses proton motive force to facilitate bacterial protein translocation mediated by the SecYEG translocon and SecA ATPase. Furukawa et al. describe high-resolution (2.6–2.8 Å) structures of SecDF in I forms, providing insight into a substrate binding site and a proton transport pathway through SecDF.

Accession Numbers

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5XAP
Tunnel Formation Inferred from the I-Form Structures of the Proton-Driven Protein Secretion Motor SecDF

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SUMMARY
Protein secretion mediated by SecYEG translocon and SecA ATPase is enhanced by membrane-embedded SecDF by using proton motive force. A previous structural study of SecDF indicated that it comprises 12 transmembrane helices that can conduct protons and three periplasmic domains, which form at least two characterized transition states, termed the F and I forms. We report the structures of full-length SecDF in I form at 2.6- to 2.8-Å resolution. The structures revealed that SecDF in I form can generate a tunnel that penetrates the transmembrane region and functions as a proton pathway regulated by a conserved Asp residue of the transmembrane region. In one crystal structure, periplasmic cavity interacts with a molecule, potentially polyethylene glycol, which may mimic a substrate peptide. This study provides structural insights into the Sec protein translocation that allows future analyses to develop a more detailed working model for SecDF.

INTRODUCTION
In bacteria, there are a variety of protein secretion systems, including Sec machinery, the flagellar/type III secretion system, and the TAT (twin arginine translocation) pathway (De Geyter et al., 2016). Among them, protein translocation across the cell membrane via the Sec translocon, SecYEG membrane protein complex, is essential and conserved in all organisms as the Sec pathway (Chatzi et al., 2014; Denks et al., 2014; du Plessis et al., 2011; Rapoport, 2007; Tsirigotaki et al., 2017). In E. coli, more than 30% of the newly synthesized proteins are translocated across the cytoplasmic membrane by Sec machinery. Many proteins that have a cleavable signal sequence at the N terminus are generally targeted to the membrane, whereas chaperones such as SecB keep their unfolded states. The following protein translocation is driven by the motor protein SecA ATPase. SecDF has been identified as a secretion factor involved in this protein translocation. SecDF-deficient E. coli cells, exhibiting cold sensitivity for growth, are severely defective in protein export at any temperature (Nouwen et al., 2005; Pogliano and Beckwith, 1994). SecDF has been shown to be involved in the later steps of protein translocation (Matsuyama et al., 1993), i.e., protein release from the Sec translocon to the periplasm, using proton motive force across the membrane (Tsukazaki et al., 2011). In 2011, the first crystal structure of full-length SecDF at 3.3-Å resolution, in the F form (retermed the P1-membrane-facing form in this study) (Tsukazaki et al., 2011), revealed that SecDF consists of 12 transmembrane (TM) helices belonging to the RND superfamily (Tseng et al., 1999) as well as three periplasmic domains: P1-head, P1-base, and P4. The TM region is proposed to provide a proton pathway in which a conserved Asp is crucial for proton conduction and proton-driven protein translocation activity. The P1 region interacts with unfolded substrate proteins and probably undergoes functionally coupled conformational transitions between the F and I forms (retermed P1-intermediate forms in this study). Once SecDF captures a preprotein emerging from SecYEG on the periplasmic side, such repetitive conformational transitions are thought to occur until protein translocation is complete. However, only a predicted structural model of the full-length I form is available based on the crystal structure of P1 domain fragment. Furthermore, although SecDF proton-conducting activity was detected (Tsukazaki et al., 2011), the actual proton pathway in the TM region has not yet been structurally identified. Here, we solved the crystal structures of I form, one of which shows a plausible tunnel for proton conduction across the membrane. The structures and subsequent functional analyses led to a substantially updated working model of SecDF.

RESULTS AND DISCUSSION
Crystal Structures of SecDF in I Form
In the current study, the crystal structures of full-length I form were determined using Deinococcus radiodurans SecDF 28-760 (DrSecDF) (Figure 1A; Figures S1 and S2; Table S1).
The 4.0-Å resolution structure of DrSecDF revealed the overall structure of the I form in which the N-terminal regions except for the P1-head (TM1-6 and P1-base) and the C-terminal regions (TM7-12 and P4) are assembled pseudo-symmetrically, similar to those of the previously reported F form (Tsukazaki et al., 2011), whereas the P1-head is situated just on the periplasm side of the P1-base (Figure 1A). However, this 4.0-Å structure did not further elucidate the function of SecDF. Therefore, to improve the stability of the flexible P1 domain in the crystals, we introduced two cysteine residues at positions 143 and 268, referring to the 4.0-Å structure, to fix the P1-head with higher crystallographic B-factors onto the P1-base by disulfide bond formation (Figure S1A). The mutant SecDF crystals completely formed intramolecular disulfide bonds and diffracted X-rays to over 2.5-Å resolution, enabling us to solve the structures with space groups C2 and P2_1_2_1 at 2.6- and 2.8-Å resolution, respectively (Table S1; Figure S1). The asymmetric unit of the P2_1_2_1 crystal contains two distinct molecules designated MoIA and MoIB, whereas that of the C2 crystal contains one molecule, MoIC. In Figure 1A, the crystal structure of MoIB is shown as a representative I form of SecDF.

As the structures in I form did not differ significantly (Figure S1), we hereafter mainly discuss SecDF in I form using the higher-resolution MoIA-C structures. Each orientation of the P1-head in I form as well as in the previously determined P1 domain fragment is structurally distinct (Figure S1), presumably because of its intrinsic flexibility even in I form. Superimposition of the I (MoIB) and F forms (PDB ID 3AQP) revealed an approximately 100° rigid-body rotation of the P1-head domain (Figure 1B), with a swinging motion of the P1-head on the P1-base allowed by the two connecting hinge loops.

Interaction Site of Periplasmic Region
Unique features of the MoIB crystal structure include an ambiguous electron density at the P1-head cavity, potentially from a part of the polyethylene glycol (PEG) used as a precipitant, whereas there was no electron density map (Fo-Fc) in the same region of the MoIA and MoIC structures (Figure 1C; Figure S3A). The interaction site in the cavity provides amphipathic surfaces (Figure 1C). The P1-head structure implied that the cavity captures the emerging polypeptide from the SecYEG translocon during protein translocation. We examined this hypothesis using site-directed ultraviolet-cross-linking of pBPA, an ultraviolet-reactive amino acid derivative (Mori and Ito, 2006). pBPA molecules were introduced at each indicated position in the cavity or the exterior periplasmic surface of the Escherichia coli SecDF complex (EcSecDF) (Figures S3A and S3B). Following irradiation during protein translocation, 35S-labeled polypeptide cross-linking products were only observed upon introduction of pBPA at position 407, corresponding to Q253 in the DrSecDF cavity (Figures S3C–S3E), suggesting that the P1-head cavity...
provides interaction sites for the newly synthesized polypeptide. Whereas the I-form P1 cavities in MolA, B, C, and the P1 domain fragment possess sufficient room for the molecule binding, the F-form cavity size is reduced (Figures S4A–S4E). Thus, the I form likely holds substrates more stably than the F form, which may be closely related to the dynamic catch and release of the substrate on the periplasmic side during protein translocation.

**Tunnel Architecture in TM Region**

Additionally, MolB exhibits a partially hydrophilic, penetrated tunnel through its TM region not observed in the other SecDF I-form or F-form structures (Figures 1A and 2), in large part owing to an approximately 4-Å shift of TM5 toward the outside compared to that of MolA, MolC, and the F-form structure (Figures S4F–S4I). In fact, the root-mean-square deviations (RMSDs) of TM5 showed higher values than those of other TMs (Figure S1E). A negative electrostatic potential exists at the tunnel center arising from Asp (D365), an essential, highly conserved residue for SecDF protein translocation activity and proton transport (Tsukazaki et al., 2011). The conserved Y662 residue can associate with D365 by hydrogen bonding and possesses the only fluctuating aromatic side chain in TMs in I-form structures (Figures 2A and 2B; Figures S4F–S4I). Mutational analyses of growth complementation, protein translocation activities, and proton conductance confirmed the importance of Y662 (Figures S5F and S6B).

The crystal structures of SecDF in I form identified over ten water molecules in the TM region on the cytoplasmic side, implying their accessibility to the vicinity of D365 from the cytoplasm but not from the periplasm. Unidentified electron densities within the MolB tunnel may also be due to small molecules such as water (Figure 2B). To elucidate the tunnel formation mechanism, we performed all-atom molecular-dynamics (MD)
simulations of SecDF using the highest-resolution structure MolC, which does not have the tunnel architecture, as the initial conformation. The mutation residues I143C and L268C in MolC were reverted to the wild-type residues. For MD simulation, the SecDF model was equilibrated in phosphatidyl ethanolamine (POPE). Then, we examined the effect of the protonation state of D365, which has been predicted to have unusual pKα (7.5–8.5) by PROPKA and MCCE (Li et al., 2005; Song et al., 2009), on SecDF structural change. D365 deprotonation attracted water molecules from the cytoplasmic bulk phase, leading to an outward TM5 shift (approximately 2.4 Å) to form a tunnel as revealed in MolB; the significant shift was not observed when D365 was protonated (Figures S5A–S5C). Simulations with the deprotonated also demonstrated repeated single-file hydrogen-bonded water chain formation (Figure 3) along the vicinity of the MolB crystal structure tunnel position. Rapid growth occurred from both the cytoplasmic and periplasmic sides (Movie S1). Specifically, D365 deprotonation might reduce the large free-energy barrier (z = 5–12 Å) on the hydrophobic periplasmic side of the TM region as indicated by water molecule distribution, such that the hydrophobic side-chain fluctuations of L413 and L657 induce water chain formation. Additionally, simulations indicated Y662 side-chain conformational transition between two orientations, as observed in the MolA and MolB/MolC crystal structures (Figure 2B; Figure S5G). Y662F mutant showed minimal growth complementation ability but exhibited similar translocation activity as the control. These differences may be attributable to the varying culture conditions (Figure S5F). As observed for the various Y209 mutants, the Y662F mutant could exhibit minimal SecDF activity, suggesting that the bulky aromatic side chain is essential for providing sufficient space for tunnel formation. As D365 is critical for proton conductance (Tsukazaki et al., 2011) (Figure S6B), we therefore propose that the tunnel in the I form (MolB) reflects the proton pathway in SecDF by means of a water chain generated through D365 deprotonation, wherein the proton hops between hydrogen-bonded water molecules via the Grotthuss mechanism.

Figure 3. Hydrogen-Bonded Water Molecules along the Tunnel
(A and B) Snapshots of SecDF at 60 ns (A) and 61 ns (B) in the MD simulation when D365 was deprotonated. Some water molecules in the TM region at 61 ns formed a hydrogen-bonded single-file chain along the tunnel. (C) Time courses of z coordinates of water oxygen in the TM region in the MD simulation of deprotonated and protonated D365. Yellow shading indicates the periods wherein the water chain was created. See also Figure S5 and Movie S1.
Functional Flexibility of SecDF

In vivo proton transport analysis using a pH indicator (Morimoto et al., 2010, 2011) demonstrated that P1-head flexibility is closely related to proton permeation (Figures S6A–S6C). However, MD simulations showed that tunnel formation in the D365 deprotonation condition was not accompanied by significant periplasmic domain conformational transition, perhaps owing to the lack of systemic proton flow (Movie S1). Conversely, transduction of the conformational changes between the TM region and the P1-head arising from proton permeation should alter the configuration of the sandwiched P4 and P1-base region (Figure S6D). To confirm that this dynamic structural alternation is required for SecDF activity, we performed complementation tests and a protein translocation assay using another double-cysteine mutant of SecDF (Figures S6D–S6F). Inhibition of SecDF activity following SecDF disulfide bond-mediated P4 and P1-base fixation was reversed by the reductant DTT, indicating that P1-base and P4 domain flexibility is also crucial. Thus, the SecDF domains all appear to undergo precisely organized, dynamic conformational changes in response to proton transport.

Working Model for SecDF

We propose the following plausible working model for SecDF (Figure 4), although further structural and functional analyses are required to fully understand the SecDF mechanism. Presecretory proteins are synthesized by ribosomes and post-translationally targeted to the translocon prior to translocation across the membrane. Initially, SecA ATPase repeatedly pushes the preprotein into the SecYEG. SecDF in the F form then interacts with the emergent preprotein on the periplasmic side and the P1-head conformation changes from the F to the I form accompanied with preprotein dragging. The MolB structure may represent a snapshot in which the preprotein is captured and extended into the periplasmic space by the P1 domain. The unfolded protein on the periplasmic side (Tsukazaki et al., 2011), the MolB state would undergo further structural changes to form the transient proton transport pathway. Deprotonation of the conserved Asp at the center of the pathway may release protons in the cytoplasm and trigger opening of the proton pathway. In conjunction with proton conduction from the periplasm to cytoplasm, SecDF may release substrate proteins into the periplasmic space. After preprotein release, the tunnel may close as represented by MoIA and MoIC followed by a conformational change from the I to the F form. These dynamic transition cycles may repeat until the secretory protein is completely translocated. Notably, a SecDF paralog present in some bacteria such as Vibrio alginolyticus incorporates a sodium ion rather than a proton gradient (Ishii et al., 2015), probably through a similar penetrating tunnel of equivalent origin, which would be sufficiently large to allow influx of sodium ions via the TM region. However, the ion selectivity of SecDF remains unclear.

EXPERIMENTAL PROCEDURES

Purification and Crystallization of SecDF

The gene encoding DrSecDF(Met-28-768)-His8 registered in NCBI Reference Sequence (WP_010888457.1) was inserted into pTV118N (Takara), and the recombinant plasmid was introduced into E. coli strain BL21(DE3). The procedures from cell culture to Ni-NTA column chromatography were as described previously (Tsukazaki et al., 2006). Column fractions were concentrated using an Amicon Ultra (50-kDa cutoff; Merck Millipore) and loaded on a Superdex 200 16/600 (GE Healthcare) in 20 mM Tris-HCl, pH 8.0, 300 mM NaCl, 0.1 mM 4-[(2-aminoethyl)] benzenesulfonyl fluoride hydrochloride, and 0.1% n-dodecyl-beta-c-maltoside (DDM). SecDF was concentrated to about 20 mg/mL and crystallized using a Gryphon LCP (Art Robbins Instruments) (Kumazaki et al., 2014). SecDF crystals were grown at 20°C in a reservoir solution containing 38% PEG200, 100 mM Na-citrate, pH 5.9, and 100 mM NH₄-citrate in 7 days. The SecDF mutants were prepared and crystallized by the same procedures. The C2 and P2₁2₁2₁ crystals of DrSecDF(I143C, L268C) were grown in reservoir solutions of (41% PEG200, 100 mM Na-citrate, pH 5.9, and
100 mM Na$_2$SO$_4$ and (40%) PEG200, 100 mM Na-citrate, pH 6.6, and 100 mM Li$_2$SO$_4$, respectively.

**Data Collection and Structure Determination**

The X-ray diffraction data were collected at Spring-8 beamline BL32XU. Initial phases were calculated by molecular replacement with the SecDF structure (PDB ID 3AQP), except for the flexible P1-head, using Phaser (McCoy et al., 2007). The model was rebuilt manually using COOT (Emsley and Cowtan, 2004) and refined using PHENIX (Adams et al., 2010). Molecular graphics were generated with PyMol (Schrödinger) and CueMol2 (http://www.cuemol.org/).

**Complementation Tests and Translocation Assay of E. coli SecD and SecF Mutants**

The secD1(Cs) transformants, carrying pFLAG-CTC and a plasmid encoding hemagglutinin (HA)-tagged E. coli SecD/SecF mutant, were prepared as described previously (Tsukazaki et al., 2011). For growth tests, the secD1(Cs) cells were spotted onto Luria-Bertani (LB)-agar plates and incubated at 20°C for 48 hr. To check SecDF accumulation, the cells were grown in LB medium at 20°C for 12 hr. The accumulation of SecDF was confirmed by immunoblotting (Tsukazaki et al., 2011). In vivo translocation assay was performed as described previously (Tsukazaki et al., 2011). Dialyze the formation of SecSecDFR439C/E132C was induced by diamide (Tsukazaki et al., 2011).

**In Vitro Photo-Cross-Linking with Preprotein**

E. coli secD1(Cs) was transformed with pHM649 encoding the pBPA-incorporating system (Mori and Ito, 2006) and a secD-secF plasmid with an engineered amber codon in secD. Cells were grown in LB medium containing 0.1 mM pHBPA until mid-log phase, and production of SecD(pBPA) was then induced for 1 hr at 37°C with 0.2% arabinose. Inverted membrane vesicles (IMVs) were prepared and used to generate the 35S-proOmpA(L59) translation intermediate in a standard translation reaction with 5 mM ATP and 5 mM succinate (Tsukazaki et al., 2011; Uchida et al., 1995). Samples were mixed with an equal volume of 50 mM HEPES-KOH (pH 7.5)-5 mM MgSO$_4$ and incubated at 37°C for 60 min. For detection of translation intermediates, samples were treated with 0.6 mg/mL proteinase K at 0°C for 20 min and precipitated with 5% trichloroacetic acid (TCA). They were analyzed by SDS-PAGE and phosphor imaging. For interaction analysis, samples were either directly precipitated with 5% TCA or irradiated with UV at 40 W/m$^2$ using a UV lamp (Ultraviolet Products) at a distance of 4 cm. They were analyzed by SDS-PAGE and phosphor imaging, either directly or following immunoprecipitation with anti-SecD.

**Intracellular pH Measurement**

The E. coli BL21(DE3) was transformed with the DrsecD-encoding plasmid and pYVM063, which encodes the ratiometric pH indicator protein pHluorin (Mishima et al., 2011). The transformants were grown in LB containing 1 mM isopropl [1-3H]thiogalactopyranoside (IPTG) and 0.2% arabinose at 30°C for 8 hr. To reduce disulfide crosslinking, 5 mM DTT was added. The fluorescence spectra were measured using a fluorescence spectrophotometer (RF-5300PC; Shimadzu). Intracellular pH was determined from a pH-dependent calibration curve generated by the 410/470 nm excitation ratios of purified pHluorin (Mishima et al., 2011).

**All-Atom MD Simulation of SecDF in a POPE Lipid Bilayer**

We examined the effect of the protonation state of D365 on the conformational change of SecDF by using all-atom MD simulations with GENESIS (Jung et al., 2013). We modeled two systems (D365E and D365S), where D365 is protonated in D365E and deprotonated in D365S. The protonation states of the other acidic and basic residues were predicted with reference to pKa values calculated by PROPKA and MCCE (Li et al., 2005; Song et al., 2009). The X-ray crystal structure of the I-form SecDF (MoC) was embedded in a POPE lipid bilayer and solvated with 150 mM KCl salt solution. The mutation sites, I143C and L268C, in MoC were altered to the wild-type residues. We performed 200-ns MD simulations in the NPT ensemble at $T = 300$ K and $P = 1$ atm. The protocols and algorithms used in the MD simulations were the same as those in our previous study on SecYEG (Tanaka et al., 2015).

**ACCESS NUMBERS**

The accession numbers for the atomic coordinates of SecDF reported in this paper are PDB: 5XAM, 5XAN, and 5XAP.

**SUPPLEMENTAL INFORMATION**

Supplemental Information includes six figures, one table, and one movie and can be found with this article online at http://dx.doi.org/10.1016/j.celrep.2017.04.030.

**AUTHOR CONTRIBUTIONS**


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