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Reconstitution of Sec-dependent membrane protein insertion: Nascent FtsQ interacts with YidC in a SecYEG-dependent manner

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Summary

The inner membrane protein YidC is associated with the preprotein translocase of Escherichia coli and contacts transmembrane segments of nascent inner membrane proteins during membrane insertion. YidC was purified to homogeneity and co-reconstituted with the SecYEG complex. YidC had no effect on the SecA-SecYEG mediated translocation of the secretory protein proOmpA. However, using a cross-linking approach, the transmembrane segment of nascent FtsQ was found to gain access to YidC via SecY. These data indicate the functional reconstitution of the initial stages of YidC-dependent membrane protein insertion via the SecYEG complex.

Introduction

In Escherichia coli, most secretory preproteins are translocated across the cytoplasmic membrane by a multimeric integral membrane complex termed translocase (reviewed in 157). Periplasmic and outer membrane proteins are synthesized with an amino-terminal, cleavable signal sequence (preproteins), and stabilized in a translocation-competent state by the SecB chaperone. The SecB-preprotein complex is targeted to the peripheral subunit of the translocase, the ATPase SecA. SecA drives the
translocation of the preprotein through a transmembrane pore formed by SecY, SecE and SecG (157). Insertion of most inner membrane proteins also depends on translocase, but these proteins are co-translationally targeted to the SecYEG complex by the signal recognition particle (SRP) and the SRP receptor FtsY (46, 153, 234). SRP interacts with hydrophobic transmembrane segments (TMS) of nascent membrane proteins (16, 232, 233). Upon interaction with membrane-associated FtsY, SRP delivers the ribosome nascent chain complex to the translocase (234). Insertion of TMSs can occur in the absence of SecA (182, 212), while translocation of large periplasmic loops has been shown to require SecA (8, 146, 230). Recently, the 60-kDa inner membrane protein YidC was identified as a translocase-associated component that interacts with TMSs of Sec-dependent inner membrane proteins during insertion (115, 205, 213). YidC is a homologue of the *Saccharomyces cerevisiae* Oxa1p that plays a crucial role in the biogenesis of some mitochondrial inner membrane proteins (99, 102, 103). Depletion of YidC in *E. coli* affects both Sec-dependent and Sec-independent membrane protein insertion (205). Here, we report the functional co-reconstitution of the *E. coli* YidC protein with the SecYEG complex yielding proteoliposomes that catalyze Sec-dependent membrane protein insertion.

**Results and discussion**

**YidC purifies as monomeric and dimeric forms**

YidC is a polytopic inner membrane protein with a molecular mass of 61 kDa. To facilitate its purification, a histidine tag was introduced at the C-terminus of YidC, and the gene was placed under control of the *lac* promoter yielding the expression vector pEH1hisYidC. To determine if his-tagged YidC is functional *in vivo*, pEH1hisYidC was transformed to the YidC depletion strain JS7131 (205). In this strain, the chromosomal *yidC* gene is disrupted and an intact *yidC* gene under control of the *araBAD* promotor has been introduced. JS7131 is not viable on Luria-Bertani (LB) agar plates containing 0.2 % glucose, since under these conditions expression of *yidC* from the *araBAD* promotor is tightly repressed. Transformation with pEH1hisYidC restored growth of JS7131 in the presence of glucose (Fig. 1A) indicating that plasmid-encoded his-tagged YidC is functional. For overproduction, pEH1hisYidC was transformed to strain *E. coli* SF100 (12).
YidC purification and reconstitution

Fig. 1: (A) His-tagged YidC complements YidC depletion strain *E. coli* JS7131. Cells containing pEH1hisYidC or the empty expression vector were grown on LB plates under conditions of induction (0.2 % arabinose) or repression (0.2 % glucose) of the chromosomally encoded *yidC* gene. (B) Purification of YidC. SDS-PAGE and CBB staining of solubilized IMVs (lane 1) and the unbound (lane 2), wash (40 mM imidazole) (lane 3), and eluted (400 mM imidazole) (lane 4) protein fractions of the Ni²⁺-NTA chromatography. (C) Blue Native PAGE of purified YidC. Purified YidC protein (20 µg) was pre-incubated for 15 min room temperature in the presence or absence of 1 % SDS prior to gel electrophoresis.

YidC could be overproduced up to 10 % of total membrane protein (Fig. 1B, lane 1). Inner membrane vesicles (IMVs) were solubilized with dodecylmaltoside (DDM) and the YidC was purified by Ni²⁺-NTA agarose chromatography (Fig. 1B, lane 4).
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Blue Native PAGE, two major bands were observed with molecular masses of ~80 and ~160 kDa (Fig. 1C, lane 1). These two bands were also observed when YidC was solubilized and purified in Triton X-100, n-octylglucoside or Aminoxid W35 (data not shown). After pre-treatment of the sample with 1 % SDS, only the 80 kDa band could be detected (Fig. 1C, lane 2). Apparently, the protein purifies as a mixture of monomers and dimers. We do not know if YidC is functional in a multimeric form. However, it is of interest to note that detergent-solubilized SecYEG complex also migrates as monomers and dimers on Blue Native PAGE (data not shown). Under translocation conditions, they are recruited by SecA to form a tetrameric pore (156).

Fig. 2: Protein profiles of proteoliposomes reconstituted with YidC and the SecYEG complex. Proteoliposomes were reconstituted with 0 (-) or 20 (+) µg of SecYEG complex with 0 (-), 10 (+), 30 (++) or 60 (+++) µg YidC protein. Samples were analyzed by SDS-PAGE and silver staining.
**YidC does not affect SecA-SecYEG-mediated translocation of proOmpA into proteoliposomes**

For functional studies, varying amounts of YidC were co-reconstituted with SecYEG into liposomes composed of *E. coli* phospholipids (Fig. 2). The majority of SecYEG and YidC were stably integrated into the liposomes, as 70-80% of the protein was resistant to carbonate extraction (data not shown). To analyze the effect of YidC on the SecA/SecYEG-mediated translocation of secretory preproteins, the translocation of the precursor proOmpA was tested. Increasing amounts of YidC only slightly inhibited the translocation of proOmpA (Fig. 3), and this correlated with a reduced level of the proOmpA-stimulated SecA ATPase activity (Fig. 3). As expected, YidC proteoliposomes did not stimulate the SecA ATPase (Fig. 3, lane 5). These results are consistent with *in vivo* studies (205) and demonstrate that YidC is not involved in proOmpA secretion.

![Fig. 3: Effect of reconstituted YidC on proOmpA translocation and proOmpA-stimulated SecA ATPase activity of SecYEG proteoliposomes. Translocation of $^{125}$I-proOmpA was carried out for 10 min at 37°C. SecA ATPase activity was assayed in the presence (white bars) or absence (black bars) of proOmpA. The activity measured with SecYEG proteoliposomes in the presence of proOmpA was set to 100%.](image-url)
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**Sequential interaction of the transmembrane segment of FtsQ with SecY and YidC during membrane insertion into proteoliposomes**

The *E. coli* FtsQ is a monotopic type II membrane protein that interacts with YidC in vitro (213), and that depends on YidC for efficient membrane insertion in vivo (230). Site-specific crosslinking has been used to demonstrate an interaction between YidC and the TMS of a nascent FtsQ during insertion into *E. coli* IMVs (213). To determine if YidC was functionally reconstituted into the liposomes, an insertion intermediate of FtsQ (108FtsQ) was generated by in vitro translation of truncated mRNAs and SRP-/FtsY-mediated targeting of the ribosome-bound nascent chains to the proteoliposomes. In these nascent chains, the photoreactive amino acid analogue 4-(3-trifluoromethyl-3-diazirinyl)phenylalanine [(Tmd)Phe] was introduced into the middle of the TMS (position 40) by suppression of a TAG codon using (Tmd)Phe-tRNA\textsuperscript{sup}. As shown previously, UV irradiation leads to crosslinking of SecY and YidC in IMVs (Fig. 4A, lane 1) (213, 230). With SecYEG proteoliposomes, a crosslink to SecY was obtained (Fig. 4A, lane 3). However, in SecYEG/YidC proteoliposomes, an additional crosslink to YidC was found, which increased with the amount of YidC present in the proteoliposomes (Fig. 4A, lanes 4-6). Strikingly, no YidC crosslink was observed with proteoliposomes containing YidC alone (Fig 4A, lane 7). The same results were obtained with 108FtsQ containing a unique cysteine at position 40 (Fig. 4B). In this case, crosslinking was induced by the homo-bifunctional reagent bis-maleimidoethane (BMOE). These data demonstrate that the purified YidC has been functionally reconstituted into proteoliposomes, and that YidC co-operates with SecY to mediate the membrane insertion of FtsQ.

The crosslinking data indicate that the TMS of inserted 108FtsQ is in the vicinity of SecY and YidC. However, since YidC can only be crosslinked to 108FtsQ in the presence of SecYEG, it appears that the TMS first interacts with SecY before gaining access to YidC. To test this hypothesis, a shorter nascent chain of FtsQ was used, i.e., a 77mer with the TAG codon at position 40. At this nascent chain length, the crosslinking probe is just exposed outside the ribosome. The 77FtsQ derivative was strongly crosslinked to SecY, and barely to YidC (Fig. 5). The efficiency of SecY-crosslinking to 77FtsQ increased with the amount of YidC. Apparently, YidC stabilizes the interaction between the TMS of nascent FtsQ with SecY.
**Fig. 4:** The TMS of membrane-inserted 108FtsQ interacts with reconstituted YidC in a SecYEG-dependent manner. (A) *In vitro* synthesis of an FtsQ 108mer with an UAG codon at position 40 was carried out in the presence of (Tmd)Phe-tRNA SUP. After the addition of reconstituted SRP, SRP-ribosome-nascent chain-complexes were purified by centrifugation through a high salt sucrose cushion, and used in targeting reactions containing IMVs or proteoliposomes. Samples were UV-irradiated, subsequently extracted with carbonate and analyzed on 13 % SDS-PAGE. (B) *In vitro* synthesized nascent FtsQ 108mer with a unique cysteine at position 40 was incubated with IMVs or proteoliposomes, and crosslinked with 1 mM BMOE. The higher molecular mass of the crosslinking products (X-SecY and X-YidC) in the proteoliposomes as compared to IMVs is due to the presence of histidine tags on YidC and SecY.
**Fig. 5:** The TMS of membrane-inserted 77FtsQ interacts with SecY, but not with YidC. *In vitro* translation of 77FtsQTAG40, targeting, and photocrosslinking were carried out as described in the legend of figure 4.

**Concluding remarks**

In this paper we report the functional reconstitution of purified YidC into proteoliposomes. Our crosslinking data indicate that YidC cooperates with SecYEG in mediating membrane insertion of nascent FtsQ. Remarkably, crosslinking to YidC depends on the presence of SecYEG in the proteoliposomes and on a sufficient length of the nascent chain. These results support a model, in which ribosome-bound nascent inner membrane proteins are first targeted by SRP to the SecYEG complex (234), a process that likely involves high affinity binding of the ribosome to the SecYEG complex (195). Next, the nascent chain is transferred to SecY, and concomitantly with chain elongation, inserted TMSs move to YidC. YidC presumably mediates the lateral diffusion of TMS into the lipid phase (205, 213, 230). The observation that nascent FtsQ gains access to YidC via SecYEG indicates that there is at least a transient interaction between SecYEG and YidC. However, stable complexes between SecYEG and YidC in the proteoliposomes could not be detected by immunoprecipitation (data not shown).
Our study is a first step towards the complete reconstitution of membrane protein biogenesis into proteoliposomes. Complete assembly of full length FtsQ, as monitored in IMVs by the formation of a protease-protected fragment representing the C-terminal periplasmic domain (data not shown), was not observed in SecYEG/YidC proteoliposomes with or without SecA. Other protein components might be involved in the process of FtsQ biogenesis such as SecDFyajC, a translocase-associated complex, whose exact function is not clear yet. In contrast to IMVs, proteoliposomes are also unable to generate a proton motive force, which might be required for assembly. Future studies will focus on the role of these components and factors to realize a complete reconstitution of membrane protein biogenesis into proteoliposomes.

Materials and Methods

Strains and plasmids

*E. coli* strain JS7131 (205) was used for complementation studies, and strain SF100 (12) was used for overexpression of YidC. Plasmid pEH1hisYidC encodes C-terminally His-tagged YidC under control of the *lac* promotor. The hexa-histidine-tag on *yidC* was introduced by PCR using the primers 5’-CAGTATTTCGCGACGGCGTG-3’ (forward) and 5’-CTGAAAGCTTCAGTGGTGGTGGTGGGATTTTTTCTTCTCGCGG (reverse) and plasmid pAra14-YidC (213) as template. The *Kpn*I/*Sma*I fragment of pEH1YidC (gift by J.-W. de Gier) was replaced by the *Kpn*I-digested PCR product. For the preparation of truncated mRNAs, plasmids pC4Meth108FtsQTAG40 (213), pC4Meth77FtsQTAG40 (230) and pC4Meth108FtsQCys40, constructed by replacing the TAG codon at position 40 of FtsQ in pC4Meth108FtsQTAG40 by a TGC codon, were used. Strain MC4100 was used to obtain translation lysate and IMVs for the targeting reactions.

Overexpression and purification of YidC

An overnight culture of *E. coli* SF100 carrying pEH1hisYidC was diluted 40-fold into fresh LB medium supplemented with 50 µg/ml kanamycin and grown to mid-logarithmic phase (OD₆₆₀ = 0.6). Expression of *yidC* was induced by the addition of 0.5 mM isopropyl-β-thiogalactopyranoside and growth was continued for another 2
hours. Cells were disrupted by passage through a French press (10,000 psi) and IMVs were isolated as described (124). IMVs at 1 mg of protein/ml were solubilized in 2 % (w/v) DDM, 10 mM TrisHCl (pH 8.0), 100 mM KCl, 10 mM imidazole, and 20 % (v/v) glycerol. After 1 h at 4 ºC, insoluble material was removed by centrifugation (30 min. at 100,000 g), and the supernatant was mixed with Ni²⁺-NTA agarose beads (Qiagen, Hilden, Germany) equilibrated with buffer A (10 mM TrisHCl pH 8.0, 100 mM KCl, 20 % [v/v] glycerol, 0.1 % [w/v] DDM) containing 10 mM imidazole. The suspension was gently shaken at 4 ºC, and after overnight incubation, poured into a column. Beads were washed with 5 column volumes of buffer A containing 40 mM imidazole, and bound proteins were eluted with 10 mM TrisHCl (pH 7.0), 100 mM KCl, 400 mM imidazole, 20 % (v/v) glycerol, 0.1 % (w/v) DDM. YidC-containing fractions were pooled, frozen in liquid nitrogen and stored at -80 ºC.

Reconstitution of YidC and SecYEG into proteoliposomes

Acetone/ether-washed E. coli phospholipids (100 µl; 4 mg/ml) (Avanti Polar Lipids Inc., Birmingham, AL) were solubilized with 7 mM Triton X-100, and mixed with the indicated amounts of purified SecYEG (239) and/or YidC. After 30 min on ice, the mixture was diluted with 800 µl buffer R (50 mM HEPES/KOH pH 8.0, 50 mM KCl) and supplemented with Bio-Beads SM-2 (20 mg wet weight) (Biorad Laboratories, Hercules, CA). The suspension was gently shaken at 4 ºC, and the Bio-Beads were replaced after 2, 4 and 6 h. After overnight incubation, the Bio-Beads were discarded and the proteoliposomes were collected by centrifugation (20 min at 200,000 g). The pellet was washed twice with 2 ml buffer R and finally suspended in 100 µl buffer R. Proteoliposomes were frozen in liquid nitrogen and stored at –80 ºC.

In vitro transcription, translation, targeting and crosslinking

Truncated mRNAs were prepared by linearizing the FtsQ derivative plasmids with HindIII followed by transcription using T7 polymerase according to the manufacturer conditions (Ambion Inc., Austin, TX). For photocrosslinking, (Tmd)Phe was site-specifically incorporated into nascent FtsQ by suppression of UAG stop codons using (Tmd)Phe-tRNASup in an in vitro translation system as described by Houben et al. (115). After 3 min of incubation at 37 ºC, 1 mM aurintricarboxylic acid (ICN Biomedicals Inc., Aurora, OH) was added to inhibit further initiation of translation resulting in a homogeneous nascent chain length. Translation reactions
were stopped after 20 min by adding 30 µM chloramphenicol. Reconstituted SRP (234) was added at 260 nM, and the mixture was incubated for 5 min at 37 °C. SRP-ribosome-nascent chain complexes were purified from the translation mixture by centrifugation through a sucrose cushion (112). They were resuspended in 100 mM KOAc, 5 mM Mg(OAc)₂, 50 mM Hepes-KOH (pH 7.9) and used in targeting reactions together with 0.56 µM FtsY, 50 µM GTP, 50 µM ATP and either IMVs or proteoliposomes as described (212). Photocrosslinking was induced by UV irradiation (213). Soluble and peripheral membrane proteins were extracted with 0.18 M Na₂CO₃ (pH 11.3) for 10 min on ice. Membrane fractions were collected by centrifugation (10 min, 110,000 g) and analyzed on 13 % SDS-PAGE and phosphoimaging. For chemical crosslinking, the same procedure was followed except that a standard in vitro translation system was used (234). Crosslinking was induced with 1 mM BMOE (Pierce, Rockford, IL), and after 10 min at 26 °C, reactions were quenched by the addition of 5 mM β-mercaptoethanol.

Other methods

Purification of E. coli SecA, SecB and proOmpA was carried out as described previously (239). Blue Native PAGE was performed on 6-16 % gradient gels as described (209). ProOmpA translocation (240) and SecA translocation ATPase activity (149) were assayed as described. Protein concentrations were determined with the DC Protein assay (Bio-Rad, Hercules, CA).

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