Conserved Negative Charges in the Transmembrane Segments of Subunit K of the NADH:Ubiquinone Oxidoreductase Determine Its Dependence on YidC for Membrane Insertion

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All members of the Oxa1/Alb3/YidC family have been implicated in the biogenesis of respiratory and energy transducing proteins. In *Escherichia coli*, YidC functions together with and independently of the Sec system. Although the range of proteins shown to be dependent on YidC continues to increase, the exact role of YidC in insertion remains enigmatic. Here we show that YidC is essential for the insertion of subunit K of the NADH:ubiquinone oxidoreductase and that the dependence is due to the presence of two conserved glutamate residues in the transmembrane segments of subunit K. The results suggest a model in which YidC serves as a membrane chaperone for the insertion of the less hydrophobic, negatively charged transmembrane segments of NuoK.

In *Escherichia coli* the inner membrane contains essential energy transducing complexes such as components of the electron transport chain. The majority of inner membrane proteins are inserted cotranslationally via the general secretory pathway otherwise known as the Sec system. In this pathway, the bacterial signal recognition particle targets ribosome-bound nascent chains to the SecYEG translocase via the signal recognition particle receptor FtsY. Membrane insertion of these proteins proceeds by a cotranslational “threading mechanism” in which the accessory protein YidC is postulated to play an important role in the clearance of transmembrane segments (TMSs) from the SecYEG channel (1, 2). A small subset of integral membrane proteins are targeted directly to YidC where they are integrated into the membrane in a Sec-independent manner. YidC belongs to the evolutionarily conserved Oxa1/Alb3/YidC family. Oxa-related proteins have been identified in all genomes sequenced to date and are postulated to have a varying role in the biogenesis of respiratory complexes I, III, IV, and V as illustrated by studies in *Neurospora crassa* (15), *Podospora anseria* (16), HEK293 cells (17), and *Saccharomyces cerevisiae* (6, 7, 18–20). This highlights that although the general function of Oxa1/Alb3/YidC family proteins is known, each member plays a specific and in some cases yet to be identified role in the biogenesis of respiratory proteins.

In a recent study, we showed that YidC depletion in *E. coli* resulted in a cessation of growth under anaerobic conditions and that this growth defect may be in part due to reduced levels of the complex I homolog in bacteria, the NADH:ubiquinone reductase, or NADH dehydrogenase I, in the membrane (21). In particular, levels of the smallest membrane subunit K (NuoK) decreased. The aim of this study was to elucidate the role of YidC in NuoK membrane biogenesis and to determine the structural features of NuoK underlying this role. We found that in vitro synthesized NuoK requires both SecYEG and YidC for insertion and that two conserved negative charges in TMSs 2 and 3 determine the dependence of NuoK on YidC for insertion.

**Experimental Procedures**

**Bacterial Strains and Plasmids**—The YidC depletion strain *E. coli* FTL10 (22) was a generous gift of Frank Sargent (University of East Anglia, Norwich, UK). *E. coli* FTL10 YidC*<sup>+</sup>* and YidC*<sup>−</sup>* strains were used as described (21). Strain *E. coli* SF100 was used for SecYEG overexpression and wild type and SecYEG*<sup>+</sup>* inner membrane vesicles (IMVs) were prepared as described (23). *E. coli* BL21 (DE3) Rosetta™ (Novagen) was
YidC Function

FIGURE 1. Membrane insertion of NuoK is affected by YidC and SecYEG levels in the membrane. A, membrane topology of NuoK indicating the glutamate residues in TMSs 2 and 3. B and C, in vitro synthesis of NuoK were carried out at 37 °C for 30 min in the presence of 10 µg of IMVs. After the synthesis reaction, the membranes were collected through a sucrose cushion and resuspended in 50 mM HEPES-KOH, pH 8. Proteinase K was added to 1.6 mg/ml in the absence (lanes 2 and 3) or presence of SDS (lanes 4 and 6). The standards of 10% of the synthesis reactions are shown (lanes 1 and 4). The synthesis reactions were performed in the presence of YidC+ or YidC− IMVs (B) or wild type (WT) or SecYEG+. The background signal of proteinase K-resistant NuoK observed when no liposomes were present was deducted from the data. All of the data points shown are the averages of three independent experiments. The bars indicate the standard errors of the mean.

Materials—Sodium fumarate was purchased from Sigma-Aldrich. Antiserum against YidC was raised in chickens against purified His-tagged YidC (Agrisera AB, Vännä, Sweden). Anti-sera against subunit K of the NADH dehydrogenase I (NuoK) was a generous gift from Takao Yagi (The Scripps Research Institute). Alkaline phosphatase-conjugated anti-chicken and anti-rabbit IgG were purchased from Sigma-Aldrich.

In Vitro Synthesis and Insertion Reactions—Synthesis reactions were carried out essentially as described by Saller et al. (25). The reactions were carried out for 30 min at 37 °C using T7 polymerase (Fermentas) and EasyTag express protein labeling mix (PerkinElmer Life Sciences) in the presence of 10 µg of IMVs or proteoliposomes. A small sample of the reaction was removed as a synthesis control, and in the case of NuoK constructs, the remainder was spun through a sucrose cushion consisting of 50 mM HEPES-KOH, pH 8, 0.5 mM phenylmethylsulfonyl fluoride, and 20% (w/w) sucrose to collect the membranes. The isolated membranes were resuspended in 50 mM HEPES-KOH, pH 8, and treated with 1.6 mg/ml proteinase K for 30 min on ice in the absence or presence of 0.1% SDS. For FtsQ, the reaction was treated with proteinase K following synthesis. The samples were trichloroacetic acid-precipitated and analyzed by SDS-PAGE and phosphorimaging.

RESULTS

Membrane Insertion of NuoK Is Adversely Affected by Decreased YidC and SecYEG Levels in the Membrane—NuoK is a 11-kDa protein containing three TMSs, connected by two short (0.3 and 1.2 kDa) loops, with a C terminus of 1.4 kDa (Fig. 1A). To monitor the requirements of NuoK insertion, NuoK was synthesized in vitro from plasmid pETNuoK. This resulted in the production of a 10−11-kDa protein visualized on SDS-PAGE (Fig. 1B, lane 1). The thick diffuse banding pattern is typically observed for hydrophobic proteins. When in vitro synthesis was performed in the presence of YidC+ IMVs, proteinase K treatment of NuoK resulted in a protein band ~1.4 kDa smaller than full-length NuoK (Fig. 1B, lane 2). The fragment most likely represents digestion of the C-terminal residues (ΔC-NuoK). Solubilization of the IMVs with SDS prior to proteinase K digestion resulted in degradation of all full-length NuoK, whereas a weak signal of ΔC-NuoK remains likely because of protease protection by the SDS micelles around the three TMSs. We therefore conclude that the signal observed after proteinase K treatment represents membrane-inserted NuoK in its correct topology.

IMVs were isolated from YidC+ and YidC− strains, and YidC levels were monitored by Western blot to confirm YidC depletion (supplemental Fig. S1). When in vitro synthesis was performed in the presence of YidC+ and YidC− IMVs, NuoK insertion into IMVs was dramatically reduced in the YidC− strain.
YidC Function

![Figure 2. Insertion of NuoK is stimulated in the absence of a PMF. Upper panel, insertion assays with YEG+ IMVs were performed as described in the legend to Fig. 1 in the absence (lanes 1 and 3) or presence (lanes 2 and 4) of 3 μM nigericin/valinomycin (nig/val) to dissipate the PMF. As a control, insertion assays with in vitro synthesized FtsQ were performed (lower panel) in the absence (lanes 1 and 3) or presence (lanes 2 and 4) of 3 μM nigericin/valinomycin. The 10% standards of the synthesis reactions are shown (lanes 1 and 2). MW, molecular mass.](image)

Renders NuoK independent of YidC for insertion

YidC overexpressed from plasmid pEH1YidC was purified on a Ni²⁺-NTA column, NuoK was found to copurify with YidC (Fig. 3, lane 2). The recovery of NuoK on the Ni²⁺-NTA column was specific for the presence of His-tagged YidC because it was not recovered when the nontagged YidC was used as a control (Fig. 3, lane 1).

Both YidC and SecYEG Are Essential for NuoK Insertion into Proteoliposomes—To elucidate the minimal requirements for insertion of in vitro synthesized NuoK, insertion assays were performed with proteoliposomes containing purified YidC and SecYEG alone and both YidC and SecYEG (Fig. 4) (24). In the absence of (proteo)liposomes, almost all of the in vitro synthesized NuoK was degraded upon incubation with proteinase K (lane 10), whereas when NuoK was synthesized in the presence of empty liposomes, a small amount of proteinase K-resistant ΔC-NuoK remained (lane 9). A comparable amount of protease-resistant ΔC-NuoK was observed when NuoK was synthesized in the presence of YidC-containing proteoliposomes (lane 6) and could have arisen from interaction of the hydrophobic TMSs of NuoK with the (proteo)liposomes. A reproducible but slight increase in protease-protected NuoK was observed when SecYEG-containing proteoliposomes were present (lane 8), but it was only when both YidC and SecYEG were present that a high level of insertion was observed (lane 7). In agreement with the observation with IMVs, NuoK minimally requires both SecYEG and YidC for insertion in vitro.

Mutation of Glutamates at Positions 36 and 72 to Lysines Renders NuoK Independent of YidC for Insertion—YidC has been postulated to form a platform at which TMSs can be released into the membrane (33), and this may particularly be important for less hydrophobic or negatively charged TMSs (34). NuoK has two conserved glutamates at positions 36 and 72 important for less hydrophobic or negatively charged TMSs of NuoK with the (proteo)liposomes. A reproducible but slight increase in protease-protected NuoK was observed when SecYEG-containing proteoliposomes were present (lane 8), but it was only when both YidC and SecYEG were present that a high level of insertion was observed (lane 7). In agreement with the observation with IMVs, NuoK minimally requires both SecYEG and YidC for insertion in vitro.

NuoK Interacts with YidC—Oxa1 has been isolated as a complex with in vitro synthesized Atp9 (homologous to Fₐₒ ATP synthase, suggesting a role for Oxa1 in the assembly of the protein complex (20). In E. coli, in vitro synthesized Fₐₒ has been shown to copurify with YidC, suggesting that it too contacts its substrates and remains briefly associated with them (32). E. coli FTL10 was grown under anaerobic conditions in growth medium containing 10 mM fumarate and 0.5% (v/v) glycerol. Overexpression of YidC from plasmid pEH1YidC or pTrcYidC was induced with 0.5 mM isopropyl β-D-thiogalactopyranoside. When IMVs containing His-tagged YidC overexpressed from plasmid pEH1YidC were transformed with plasmids pEH1YidC and pTrcYidC, allowing the overexpression of His-tagged YidC (lane 2) and nontagged YidC (lane 1), respectively. Overexpression was performed under anaerobic growth conditions in media containing fumarate and glycerol. IMVs were isolated and solubilized in 2% N-dodecyl-β-D-maltoside whereupon YidC was purified by Ni²⁺-NTA chromatography. Elution fractions were run on SDS-PAGE, blotted onto polyvinylidene difluoride membrane, and probed with antibodies against NuoK. MW, molecular mass.
The Double Mutant NuoK E36K,E72K Requires Only SecYEG for Insertion into Proteoliposomes—Insertion of the mutants described above was investigated using proteoliposomes to determine the minimal requirements for insertion. The same set of (proteo)liposomes was used as for the wild type NuoK. The insertion patterns observed for the single mutants NuoK E36K and NuoK E72K, was comparable with those observed for wild type NuoK (Figs. 4 and 5). In the absence (of proteoliposomes, almost all NuoK E36K synthesized was digested by externally added protease K (Fig. 5B, top panel, lane 10). Proteoliposomes containing YidC only and empty liposomes afforded a small amount of protection in the presence of protease K (Fig. 5B, top panel, lanes 6 and 9), whereas synthesis in the presence of proteoliposomes containing only SecYEG resulted in a very slight but reproducible increase in insertion (Fig. 5B, top panel, lane 8) compared with that observed for YidC only and empty (proteo)liposomes. Only when synthesis was performed in the presence of proteoliposomes containing both YidC and SecYEG was an appreciable amount of inserted NuoK E36K observed (Fig. 5B, top panel, lane 7). Also for the mutant protein NuoK E72K, insertion was only observed in proteoliposomes containing both YidC and SecYEG (Fig. 5B, middle panel, lane 7). The observations with proteoliposomes support those made with IMVs.

In the presence of (proteo)liposomes, protease K treatment of in vitro synthesized NuoK E36K,E72K resulted in the formation of three protein bands (Fig. 5B, bottom panel, lanes 6–9). The upper band corresponds to undigested full-length protein, whereas the two lower bands, 2 and 2.5 kDa smaller than the full-length NuoK E36K,E72K, correspond to C-terminal truncations. If the mutant protein were inserted in the inverse topology, protease K digestion would most likely result in two bands corresponding to the N-terminal two TMSs (5.3–6.6 kDa) and TMS 3 plus the C terminus (4.2–5.5 kDa). This is not observed. Alternatively, if protease K did not cleave the protein in the periplasmic loop between TMSs 2 and 3, only full-length protein would be observed. It is therefore not possible to deduce the topology of the full-length protein observed after protease K digestion. In the absence of (proteo)liposomes, most of the NuoK was digested (Fig. 5B, bottom panel, lane 10). Incubation with protease K in the presence of proteoliposomes containing only YidC or empty liposomes resulted in a
FIGURE 5. Mutation of glutamates at positions 36 and 72 to lysines renders NuoK independent of YidC for insertion. A, insertion reactions were performed as described in the legend of Fig. 1. Synthesis reactions were performed in the presence of YidC\textsuperscript{+} or YidC\textsuperscript{−} IMVs, and proteinase K was added in the absence (lanes 2 and 5) or presence (lanes 3 and 6) of SDS. Synthesis reactions were performed with plasmids containing single mutants NuoK E36K (top panel) and NuoK E72K (middle panel) and double mutant NuoK E36K,E72K (bottom panel). The 10% standards of the synthesis reactions are shown (lanes 1 and 4). B, NuoK insertion was quantified from the amount of protease-protected material in the presence of YidC\textsuperscript{+} (indicated in white) and YidC\textsuperscript{−} (indicated in dark gray) IMVs. All of the data points shown are the averages of three independent experiments. The bars indicate the standard errors of the mean. C, NuoK insertion reactions were performed in the presence (lanes 1–4 and 6–9) or absence (lanes 5 and 10) of (proteo)liposomes reconstituted with or without purified SecYEG and YidC as indicated. The insertion reactions (lanes 6–10) were performed essentially as described in the legend to Fig. 1 using plasmids containing single substitutions NuoK E36K (top panel) and NuoK E72K (middle panel) and double substitutions NuoK E36K,E72K (bottom panel). 5% standards of the synthesis reactions are shown (lanes 1–5). D, NuoK insertion was quantified from the amount of protease-protected material in the presence and absence of (proteo)liposomes. The background signal of proteinase K-resistant NuoK observed when no liposomes were present was deducted from the data. All of the data points shown are averages of three independent experiments. The bars indicate the standard errors of the mean.
slight increase in protease-protected protein (Fig. 5B, bottom panel, lanes 6 and 9). However, when NuoK E36K,E72K was synthesized in the presence of SecYEG-containing proteoliposomes (either alone or together with YidC), an appreciable amount of insertion was observed (Fig. 5B, bottom panel, lanes 7 and 8). This indicates that, in contrast to wild type NuoK and the single glutamate mutants, NuoK E36K,E72K requires only SecYEG for insertion in vitro.

**DISCUSSION**

All of the subunits of the bacterial complex I, NADH dehydrogenase I, are encoded by the *nuo* operon. In *E. coli*, NADH dehydrogenase I contains 13 subunits, NuoA through to NuoN with NuoC and NuoD fused to form one protein. If any of these subunits are absent, a functional enzyme complex cannot be formed (37). In a previous study, we showed that levels of the small membrane subunit NuoK are greatly reduced upon YidC depletion (21). A decrease in the levels of NuoK in the membrane could have been due to the absence of one of the other subunits under YidC-depleting conditions, but as we now demonstrate, YidC is directly involved in the insertion of NuoK. Using IMVs we showed that the levels of YidC and SecYEG affected the efficiency of NuoK insertion. The decrease in insertion efficiency in YidC− IMVs was not due to the impaired ability of the IMVs to generate a PMF because in the absence of a PMF, the efficiency of insertion of NuoK was even enhanced. It has been shown with M13 procot Lep and CyOA derivatives that the presence of positive charges in the translocated loops of these derivatives created proteins that only inserted in the absence of a PMF (38, 39). NuoK contains one negatively charged residue in the periplasmic loop between TMSs 2 and 3, whereas the cytoplasmic loop between TMSs 1 and 2 and the C-terminal tail contain numerous positively charged amino acids. The protein topology therefore follows that predicted by the “positive inside rule” (40), and it is not immediately obvious why the PMF would inhibit protein insertion. NADH dehydrogenase I is the preferred NADH dehydrogenase under anaerobic growth conditions (41). Unlike NADH dehydrogenase II, the proton-pumping NADH dehydrogenase I is energy-conserving, which is needed under the “energy limited” conditions of growth without oxygen. It is possible that NuoK insertion is enhanced under such conditions when the PMF is reduced and NADH dehydrogenase I is needed.

Although the number of known YidC-dependent membrane proteins has increased during recent years, the exact mechanism of YidC in membrane protein insertion remains enigmatic. YidC has been shown to contact the TMSs of numerous integral membrane proteins in various cross-linking studies (24, 42, 43), and yet YidC is dispensable in the insertion of most of these proteins (26, 44, 45). Using *in vitro* insertion into proteoliposomes, we show that NuoK minimally requires both YidC and SecYEG for insertion. Subunit a of the cytochrome oxidase, CyOA, has also been shown to require both YidC and SecYEG for insertion *in vitro* (12). Based on the individual insertion requirements of constructs containing either the N- or C-terminal parts of CyOA, it has been suggested that SecYEG and YidC work sequentially in the insertion of CyOA with YidC inserting the N-terminal hairpin and SecYEG inserting the C-terminal TMS and long periplasmic C-terminal tail (13, 46). This strict sequential insertion is only alleviated when a long linker between TMSs 1 and 2 is introduced, after which the C terminus can insert independently of the N-terminal hairpin (39). Interestingly the introduction of numerous positive charges in the signal peptide and TMS 1 block the insertion of this protein via the YidC pathway but do not result in default insertion via the Sec translocon (39).

We observed that the presence of the negative charges in TM 2 and 3 determines the YidC-dependent insertion. NuoK contains single glutamates in TM2 and 3, which are required for high ubiquinone activity (35, 36). Substitution of the glutamates at positions 36 and 72 for lysines produced a protein that, like most studied integral membrane proteins, required only SecYEG for integration into the membrane. This supports a model in which the TMSs of YidC have a role in forming a membrane chaperone, assisting the integration of less hydrophobic, negatively charged TMSs, similar to that proposed for Oxa1 (34, 34). The presence of just one glutamate-containing TMS renders the entire protein still YidC-dependent, and it is only when both glutamates have been substituted for lysines that the protein can be inserted by SecYEG, unassisted by YidC. NuoK is one of many integral membrane respiratory proteins that contain membrane-negative charges. NuoA has a similar structure to NuoK with glutamates at positions 81 and 102 (TM2 and 3) and an aspartate at position 79 (TM2). It would be of interest if these similar structural features necessitate YidC in the insertion process.

In the well studied substrate of YidC, *F*, mutation of an aspartate at position 61 (TM2) to a glycine produces an ATP synthase that contains an enzymatically active F₁ part but no functional F₀ (47). There were, however, detectable levels of the mutant *F*, present in the membrane. Attempts to elucidate the insertion requirements for this mutant *in vitro* were unsuccessful because of the proteinase K-resistant nature of the protein.³ Substitution of a glycine for aspartate in TM1 of *F* results in a mutant form of the protein that is still dependent on YidC for insertion but that does not form an oligomeric ring structure as the wild type protein does (32, 48). Thus the introduction of a negative charge into a TMS is tolerated by the YidC-only mode of insertion. Of the other proteins shown to insert with the assistance of YidC, the mechanosensitive channel of large conductance, MscL, and the phase coat proteins M13 and PF3 have membrane-located negative charges. CyOA, however, does not contain any membrane-negative charges. YidC is not essential for the insertion of FtsQ (26) and LepB (49), even though interaction between these proteins and YidC during membrane insertion has been observed (1, 24, 42, 43, 50). FtsQ does not contain any membrane-embedded charges, whereas LepB contains a glutamate in TM2. There must therefore be structural features other than membrane-located negative charges that confer YidC dependence to proteins.

YidC is remarkably resilient to single amino acid substitutions, and even when TMSs 4 and 5 were swapped for unrelated TMSs, YidC activity was retained (33). It has therefore been

³ S. Kol and J. de Keyzer, unpublished data.
suggested that it is the presence of the C-terminal 5 TMSs and not the specific sequences of the hydrophobic stretches that is essential for YidC activity. Our data support the hypothesis that YidC serves as a platform from which TMSs are integrated into the membrane. Furthermore, YidC is essential for Nuok TMS integration when membrane-embedded negative charges are present. The involvement of YidC in the integration of such TMSs may provide an explanation as to its conserved function in the biogenesis of respiratory proteins that often contain TMSs. This conserved function of YidC may also be explained by its ability to facilitate the integration when membrane-embedded negative charges are present. Our data support the hypothesis that it is the C-terminal 5 TMSs and not the specific sequences of the hydrophobic stretches that is essential for YidC activity.

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REFERENCES

1. Scotti, P. A., Urbanus, M. L., Brunner, J., de Gier, J. W., von Heijne, G., van der Does, C., Driessen, A. J., Oudega, B., and Luirink, J. (2000) J. Mol. Biol. 295, 1131–1147
2. de Gier, J. W., and Luirink, J. (2001) Mol. Microbiol. 41, 119–131
3. Pohlschroeder, M., Hartmann, E., Hand, N. J., Dilks, K., and Haddad, A. (1994) EMBO J. 13, 5053–5060
4. Yen, M. R., Harley, K. T., Tseng, Y. H., and Saier, M. H., Jr. (2001) J. Biol. Chem. 276, 17771–17779
5. de Gier, J. W., and Luirink, J. (2001) FEBS Lett. 501, 1–5
6. Bauer, M., Behrens, M., Esser, K., Michaelis, G., and Pratje, E. (1994) J. Biol. Chem. 269, 20412–20419
7. Bonnefoy, N., Chalvet, F., Hamel, P., Slonimski, P. P., and Dujardin, G. (1994) J. Mol. Biol. 230, 239–250
8. M. C., Nakamaru-Ogiso, E., Matsuno-Yagi, A., and Yagi, T. (2005) Biochemical Society Transactions 33, 372–378