Activators of the Glutamate-Dependent Acid Resistance System Alleviate Deleterious Effects of YidC Depletion in Escherichia coli

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The function of the essential inner membrane protein (IMP) YidC in Escherichia coli has been studied for a limited number of model IMPs and primarily using targeted approaches. These studies suggested that YidC acts at the level of insertion, folding, and quality control of IMPs, both in the context of the Sec translocon and as a separate entity. To further our understanding of YidC’s role in IMP biogenesis, we screened a random overexpression library for factors that rescued the growth of cells upon YidC depletion. We found that the overexpression of the GadX and GadY regulators of the glutamate-dependent acid resistance system complemented the growth defect of YidC-depleted cells. Evidence is presented that GadXY overexpression counteracts the deleterious effects of YidC depletion on at least two fronts. First, GadXY prepares the cells for the decrease in respiratory capacity upon the depletion of YidC. Most likely, GadXY-regulated processes reduce the drop in proton-motive force that impairs the fitness of YidC-depleted cells. Second, in GadXY-overproducing cells increased levels of the general chaperone GroEL cofractionate with the inner membranes, which may help to keep newly synthesized inner membrane proteins in an insertion-competent state when YidC levels are limiting.

Approximately 20% of the proteins encoded by the Escherichia coli genome function in the inner membrane (IM). These inner membrane proteins (IMPs) first are targeted to and then inserted into the IM, after which they fold and, if required, assemble into oligomeric complexes (51). The Sec translocon is the main protein-conducting channel used to insert IMPs (18, 52). The translocon consists of the heterotrimeric channel complex SecYEG and the accessory components YidC and SecDF YajC. It functions not only in the lateral transfer of transmembrane segments (TMs) of IMPs into the lipid bilayer but also in the vectorial translocation of protein (domains) into the periplasm.

YidC has been identified as an essential membrane factor that plays an important but poorly understood role in the biogenesis of IMPs in bacteria. YidC can act together with the SecYEG channel and also independently as a membrane insertase. Cross-linking studies have shown that YidC contacts the TMs of substrate IMPs upon their lateral exit from the Sec translocon, possibly to assist lipid partitioning (41). While YidC can act in concert with the Sec translocon, as demonstrated for the integral inner membrane lipoprotein CyoA (subunit II of the cytochrome bo oxidase), here, YidC is required for the insertion of the N-terminal region of CyoA into the membrane, followed by the translocation of the more complex C-terminal domain by the Sec translocon (4, 9, 43). While YidC can act in concert with the Sec translocon, other studies have shown that YidC also can function as an independent insertase for small proteins, such as the M13 and Pf3 phage coat proteins, the subunits a and c of the F1,F0 ATPase (F1,a and F1,c), and subunit K of the NADH dehydrogenase complex (27, 30, 31, 46). It has been shown that both aerobic and anaerobic respiratory chain complexes are affected in YidC-depleted cells (27, 28, 46), resulting in a reduction of the proton-motive force (PMF). PMF reduction leads to a strong upregulation of the stress response protein PspA, which is part of the Psp shock response thought to help maintain the PMF upon membrane stress (5, 17, 50).

Besides YidC’s role as an insertase, it is required for cotranslational folding into a stable conformation and/or assembly of multimeric IMP complexes, like MalFGK2 and MscL (23, 26, 47). The observation that YidC depletion induces the cell envelope stress response further supports a role for YidC in the folding of IMPs (35, 48). At an even later stage, a role...
for YidC in the quality control of IMPs has been implied from its association with the membrane protease and chaperone complex HikC-FtsH (42).

Although these studies have provided insight into the various roles of YidC, it remains difficult to imagine how one protein can combine the diverse functions ascribed to YidC. Pressing questions include the following: why is YidC essential, and how does YidC operate in various structural contexts, alone or connected to the Sec translocon? To gain more insight in the key processes that determine why YidC is essential, we have used an unbiased genetic screen to select factors that are able to complement the growth of cells that express YidC at sublethal levels. We found that the overexpression of gadX and gadY, regulators of the glutamic acid decarboxylase (Gad)-based acid resistance system in *E. coli*, rescues growth at low YidC levels and suppresses the concomitant PspA response.

### MATERIALS AND METHODS

**Enzymes and materials.** Restriction endonucleases and other DNA-modifying enzymes were obtained from Roche and Invitrogen. All other chemicals were supplied by Sigma. Antiserum against the His tag and the hemagglutinin (HA) tag were purchased from Roche and Sigma, respectively. Antiserum against YidC, PsPA, Lep, and GadAB have been described previously (33, 39). Antiserum against GroEL was kindly donated by Peter Lund (University of Birmingham, Birmingham, United Kingdom).

**Strains, plasmids, growth conditions, and primers.** *E. coli* strain Top10F (Invitrogen) was used for the cloning and maintenance of plasmid constructs. The YidC depletion strain FTL10 and its isogenic parental arabinose-resistant strain MC400-A were generous gifts from Frank Sargent (University of East Anglia, Norwich, United Kingdom) (13). All strains were grown routinely in Luria Bertani (LB) medium or in M9 medium where indicated, with appropriate antibiotics. To monitor cell growth, cultures were pregrown in medium with 0.2% L-arabinose, washed in fresh medium to remove L-arabinose, and diluted as appropriate for the following growth experiments. Strains, plasmids, and primers used in this study are listed in Table 1. For the construction of plasmids pEH3.GadBC-HA, the sequence encoding GadB and GadC was PCR amplified using genomic DNA of MC4100-A and the primers GadB Forw and GadC-HA Rev. The PCR product was digested with XbaI and SacI and cloned into pEH3. The nucleotide sequence was verified by DNA sequencing.

**Complementation of YidC depletion screen.** FTL10 cells were electroporated with a pBR322-based MC4100 ΔmarK mutant genomic DNA library (kindly donated by Pierre Genevaux, CNRS Université Paul Sabatier, Toulouse, France) or empty vector pBR322 as a control (Fermentas). The resulting transformants were selected for growth on agar plates containing no t-arabinose or low concentrations of t-arabinose that are unable to support growth with pBR322 as indicated. From colonies that grew on the selective agar plates, plasmids were isolated and retransformed into FTL10 to confirm the rescue phenotype. The boundaries of the inserted genomic fragments in the plasmids subsequently were determined by DNA sequencing.

**Quantitative real-time PCR.** Total RNA was isolated using an RNeasy minikit (Qiagen) from bacterial strains grown to mid-log phase in M9 and treated with RNase-free DNase (Qiagen) according to the manufacturer's protocol. The RNA quantity was determined by Nanodrop (ND-100 spectrophotometer), and the quality was assessed by Nanochip (Agilent Technologies). Only RNA with RNA integrity numbers (RINs) higher than 9 was used for subsequent analysis. cDNA samples were synthesized from total RNA using the SuperScript II reverse transcriptase kit (Invitrogen) with random primer oligonucleotides (Invitrogen). Quantitative real-time PCR (qPCR) was performed with each specific primer pair (Table 1) using Platinum SYBR green qPCR SuperMix-UDG (Invitrogen). The reactions were run on an ABI7500 detection system (Applied Biosystems); the fluorescence signal due to SYBR green intercalation was monitored to quantify the double-stranded DNA product formed in each PCR cycle. Melting curve analysis was performed after each qPCR to verify the specificity of each primer pair. The *marC* gene of 16S rRNA was used to normalize the cDNA input. For each time point and condition, four biological replicates were taken.

**Bacterial strains were grown in M9,** and cells were separated from the medium by centrifugation at 4°C for 5 min at 8,000 × g. The spent medium was assayed for lactate by high-performance liquid chromatography (HPLC) (LKB) with a REZEX organic acid analysis column (Phenomenex) at 45°C with 7.2 mM H2SO4 as the eluent, using an RI 1530

### TABLE 1. Strains, plasmids, and primers

<table>
<thead>
<tr>
<th>Strain, plasmid, or primer</th>
<th>Description</th>
<th>Reference or source</th>
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<tr>
<td><strong>Strains</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Top10F`</td>
<td>F' <em>mcrAΔ(mrr-hsdRMS-mcrBC)</em> Δ680lacZΔM15 ΔlacYQ4اسينع G carA1 araD139 Δ(xylS-leuY)7697 galE15 galK16</td>
<td>Invitrogen</td>
</tr>
<tr>
<td>MC400-A</td>
<td>F' ΔlucU69 araD139 rpsL150 relA1 psf1 flbB5301; arabinose resistant</td>
<td>13</td>
</tr>
<tr>
<td>FTL10</td>
<td>MC4100-A ΔyidC attB:*araC' pBAD yidC' * Kan'</td>
<td>13</td>
</tr>
<tr>
<td><strong>Plasmids</strong></td>
<td></td>
<td></td>
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<tr>
<td>pBR322</td>
<td>Cloning vector</td>
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<tr>
<td>pGadXY</td>
<td>pBR322 containing gadXY fragment (Fig. 1)</td>
<td>This study</td>
</tr>
<tr>
<td>pCL.CyoA-HA</td>
<td>cyoA ORF with C-terminal HA tag</td>
<td>43</td>
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<tr>
<td>pEH3.GadBC-HA</td>
<td>gadB ORF and gadC ORF with C-terminal HA tag</td>
<td>This study</td>
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<tr>
<td><strong>Primers</strong></td>
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<tr>
<td>YidC Forw</td>
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<td>YidC Rev</td>
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<td>PsPA Forw</td>
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<td>GadC Rev</td>
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<tr>
<td>RssC Forw</td>
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<td>RssC Rev</td>
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<tr>
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<tr>
<td>GadC-HA Rev</td>
<td>TCCGAGCTTATTAGCAGATGTCGCGACGTTAGGAATGATAAGATCCTGTTGTCATT</td>
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<td>CATCAG</td>
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193, 2011 GadXY RECOVERS GROWTH AT LOW YidC LEVELS 1309
TABLE 2. Expression levels of genes in FTL10 as analyzed by qPCR

<table>
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<tr>
<th>Cell type and gene</th>
<th>30 min Fold change</th>
<th>P value</th>
<th>5 h Fold change</th>
<th>P value</th>
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<td>yidC</td>
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<td>0.009</td>
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<td>0.005</td>
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<td>pspA</td>
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<td>0.632</td>
<td>22.2</td>
<td>0.000</td>
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<tr>
<td>gadX</td>
<td>1.76</td>
<td>0.269</td>
<td>1.6</td>
<td>0.362</td>
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<tr>
<td>gadC</td>
<td>2.66</td>
<td>0.055</td>
<td>1.0</td>
<td>0.929</td>
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<tr>
<td>Overexpressing pGadXY versus control in absence of 1- arabinose</td>
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<td></td>
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<tr>
<td>yidC</td>
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<td>0.238</td>
<td>1.8</td>
<td>0.221</td>
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<tr>
<td>pspA</td>
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<td>0.778</td>
<td>8.4</td>
<td>0.001</td>
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<tr>
<td>gadX</td>
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<td>0.000</td>
<td>42.2</td>
<td>0.000</td>
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<tr>
<td>gadC</td>
<td>9.3</td>
<td>0.002</td>
<td>6.7</td>
<td>0.004</td>
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RESULTS

Genetic screening for restored growth upon YidC depletion.
To gain insight in the function(s) of YidC, we performed a genetic screen to identify factors that suppress the lethality observed upon the depletion of YidC in E. coli strain FTL10 in which yidC is under the control of a 1- arabinose promoter (13). Chromosomal DNA of E. coli was partially digested with the restriction enzyme Sau3AI, ligated in the vector pBR322, and introduced into FTL10. FTL10 transformed with empty pBR322 was unable to grow on plates that contained no or low concentrations of 1- arabinose (0.001 to 0.01% range). FTL10 cells transformed with the genetic library also were plated on plates with no or low concentrations of 1- arabinose. Plasmids were extracted from clones that grew on these plates. After confirming the phenotype upon the retransformation of the plasmids into fresh FTL10 competent cells, the 5’ and 3’ ends of the inserts were sequenced.

Only one clone was obtained that was able to grow in the absence of 1- arabinose. Not surprisingly, this clone contained the yidC gene, validating our screening conditions (data not shown). In contrast, screening in the presence of a low concentration of 1- arabinose (0.001%) resulted in numerous colonies. The sequence analysis of 26 clones revealed that 10 of them contained overlapping inserts containing at least the gadX and gadY genes. The plasmid of a clone that contained only the gadXY genes (pGadXY) was further analyzed (Fig. 1).

The presence of pGadXY partially complements growth and represses the PspA response. We first studied in more detail the effect of pGadXY on growth when YidC was depleted. FTL10 cells harboring pGadXY or pBR322 were grown to mid-log phase in the presence of 1- arabinose (to sustain YidC expression) and then spotted in serial dilutions on solid LB medium containing different concentrations of 1- arabinose (Fig. 2A). The pGadXY plasmid was not able to support growth on plates without 1- arabinose, indicating that the complete absence of YidC was not complemented by the plasmid. However, cells harboring pGadXY efficiently supported growth on LB containing as low as 0.001% 1- arabinose (the concentration used in the screening), whereas cells harboring pBR322 already were strongly affected in growth at 0.01% 1- arabinose. In addition, we analyzed the effect of YidC depletion in liquid culture by diluting cells grown to mid-log phase in M9 medium containing no or 0.2% 1- arabinose. The FTL10 cells with pGadXY were less affected in growth upon YidC depletion than control cells (Fig. 2B). The difference in growth was less pronounced when cells were cultured in LB medium (data not shown). Apparently the presence of

![FIG. 1. Schematic representation of the pGadXY construct containing gadY, gadX, and partial gadA ORFs.](http://jb.asm.org/)
ulatory proteins and RNAs, including those encoded by phase stress conditions via a complex interplay of various regulators. The expression of these effector proteins is upregulated in response to low pH and stationary-extracellular glutamate (6, 53). The expression of these effector genes (36, 39). The factor that plays a critical role in the activation of Gad effector genes (Fig. 1), is encoded by the gadX gene, adjacent to gadY. The gadX gene encodes an AraC-like transcription factor that plays a critical role in the activation of Gad effector genes (36, 39). The gadY gene, adjacent to gadX (Fig. 1), is transcribed in the opposite direction and encodes a small non-coding RNA (ncRNA) that interacts with the 3′-untranslated region (UTR) of gadX mRNA to stabilize it (25). To examine the effect of gadX or gadY individually, most of the coding region of gadX or gadY was deleted in pGadXY, and the resulting constructs were tested for their ability to restore the growth of FT10 upon the depletion of YidC. The presence of the plasmids containing the individual genes showed only partial effects, but intact gadX and gadY both were required for the optimal restoration of growth (results not shown). A plausible explanation for the partial rescuing effect of gadY on its own is that gadY exerts a stabilizing effect on the endogenous gadX mRNA (25). The overexpression of gadX, cloned under the control of an inducible promoter, led to a severe inhibition of cell growth in both wild-type (WT) and YidC-depleted cells. This toxic effect probably resulted from the excessive expression of the GadX activator protein, which has been observed previously (39). For these reasons, we decided to continue with pGadXY as isolated in the screen.

A possible explanation for the rescue phenotype is that GadXY overproduction stabilizes YidC, and therefore less L-arabinose is needed to maintain sufficient YidC levels in the cells, which would lead to residual YidC levels in the absence of L-arabinose. To study the effects of GadXY overproduction on YidC expression, we analyzed the levels of YidC in FT10 cells grown in the absence of L-arabinose by Western blotting. As shown in Fig. 3A, YidC was depleted to comparable low levels in both control and pGadXY-complemented FT10 cells grown with low L-arabinose concentrations. Furthermore, qPCR analysis showed that after 30 min of the growth of FT10 in the absence of L-arabinose, the mRNA level of YidC was 5-fold lower, and after 5 h 7-fold lower, compared to that of cells grown in the presence of L-arabinose (Table 2). Furthermore, there was no significant difference in yidC mRNA levels in YidC-depleted cells with or without pGadXY (Table 2). We therefore conclude that GadXY overexpression does not lead to altered levels of YidC in FT10.

The depletion of YidC has been shown to induce a massive expression of PspA, a stress protein that is believed to respond to the dissipation of the PMF (5). This in turn is caused by the impaired membrane insertion of CyoA and Fc, subunits that are essential for the functional assembly of the cytochrome bo₃ oxidase and F₁,Fₒ ATPase complex, respectively (46). To investigate if GadXY overexpression is involved in suppressing
this strong PspA response, lysates from FTL10 cells (with or without pGadXY) were analyzed for PspA content by Western blotting (Fig. 3A). Strikingly, PspA showed little upregulation in FTL10 cells harboring pGadXY (lanes 5 to 8) despite the low levels of YidC, in contrast to what is observed in the control cells (lanes 1 to 4). Moreover, qPCR analysis revealed a strong upregulation of pspA mRNA (22-fold up after 5 h) in FTL10 cells (Table 2), while this upregulation was reduced in cells complemented with pGadXY (8-fold down after 5 h) (Table 2). Taken together, these data demonstrate that GadXY overexpression strongly represses the PspA response upon YidC depletion. Apparently, the PspA response is not a compulsory consequence of YidC depletion.

The presence of GadXY does not restore insertion of CyoA and F_{o,c}. As shown above, GadXY overexpression does not prevent the depletion of YidC in FTL10 cells grown without l-arabinose, but it does significantly repress the PspA response. One possible explanation is that GadXY overexpression leads to the improved insertion or assembly of YidC-dependent subunits of the respiratory chain complexes, thereby relieving the loss of PMF and subsequent PspA response. In concept, GadXY could do so by improving the insertase function of the remaining YidC under depletion conditions or by stimulating YidC-independent alternative insertase mechanisms. To examine the influence of pGadXY presence on the insertase function of YidC, we monitored the processing of CyoA, which depends solely on YidC for its initial insertion in the inner membrane (4, 9, 43). The membrane insertion of the lipoprotein CyoA can conveniently be monitored by analyzing the cleavage of its signal peptide by the lipoprotein-specific signal peptidase II. A plasmid expressing HA-tagged CyoA (44) was transformed into FTL10 containing either pGadXY or pBR322. Cells were grown in the absence or presence of l-arabinose, and CyoA-HA processing was followed by Western blotting using anti-HA antibodies. Consistently with previous data, the processing of CyoA-HA was impaired upon YidC depletion (Fig. 3B). The processing of CyoA-HA remained hampered in the presence of pGadXY under these steady-state conditions, suggesting that the insertase function of YidC is not improved by GadXY or that alternative mechanisms cannot complement this function. In addition, the absorption spectra of reduced versus oxidized eIM fractions that were prepared from WT, YidC-depleted, and pGadXY-containing cells demonstrated that the loss of cytochrome bo_{3} oxidase (46) was not rescued by the presence of pGadXY (results not shown).

Only a few natural substrates are known that use YidC as an independent insertase. If CyoA and F_{o,c} were the only substrates of YidC that are critical for maintaining the PMF and the viability of the cell under the experimental conditions, the overexpression of both proteins might restore the PMF and bypass the requirement of YidC for growth. However, the overexpression of either CyoA or F_{o,c} in FTL10 cells depleted of YidC did not result in the recovery of growth (data not shown). Apparently, YidC's lethality was not bypassed by CyoA and F_{o,c} overexpression and spontaneous insertion in the absence of YidC.

Overexpression of GadXY but not GadBC complements the growth defect and PspA response in YidC-depleted cells. As discussed above, neither improved YidC stability nor improved YidC insertase activity are likely explanations for the positive effect of GadXY on YidC-depleted cells. We therefore considered the explanation that the overproduction of GadXY mitigated indirect effects of YidC depletion that might be related to acidic stress. To examine this, we first studied the effects of YidC depletion on gadY, gadX, and gadBC transcription by the analysis of reporter gene activity using lacZ transcriptional fusions to the promoters of gadY, gadX, and gadBC. The β-galactosidase activities of all fusions were decreased by ~50% in YidC-depleted cells compared to those of control cells (Table 3). We did not observe significant differences in gadX and gadC mRNA levels in the qPCR analysis (Table 2). However, Western blot analysis revealed that GadAB levels were increased in FTL10, in the presence of l-arabinose, and in its parental strain MC4100-A (Fig. 4A, lanes 3, 4, and 7) (6). In contrast, FTL10 cells grown for 4 h in the absence of l-arabinose to deplete YidC showed decreased GadAB levels, in line with data from β-galactosidase assays (Fig. 4A, lane 8, and Table 3).

We further analyzed the effect of GadXY overproduction on GadAB protein levels (Fig. 4B). As expected, the levels of GadAB were higher in cells with pGadXY after 2 h of growth than the basal levels present in FTL10 cells harboring the control plasmid pBR322 (compare lanes 5 and 6 to 1 and 2). After 4 h of growth, the levels of GadAB still were higher in cells harboring pGadXY than in cells with the control plasmid (compare lanes 7 and 8 to 3 and 4), even in the absence of YidC (lane 8). Additional analysis of gadC and gadX mRNA levels by qPCR showed their increase in FTL10 with pGadXY (Table 2). Taken together, these data indicate that in YidC-depleted cells the Gad response is not switched on, but it can be reestablished by the overexpression of GadXY. The reason for the reduced GadB levels remains unexplained, but it is of interest that GadC levels were observed to be 20-fold lower in YidC-depleted inverted membrane vesicles (IMVs), hinting at a direct involvement of YidC in GadC biogenesis (J.-W. de Gier, personal communication).

Assuming that the depletion of YidC might lead to the acidification of the cells, the inability to mount a Gad response and the low levels of GadC could be an explanation for the growth defect upon YidC depletion. Following this scenario, the overexpression of GadBC would alleviate this effect. Therefore, we introduced a pEH3 plasmid containing the gadBC operon under the control of the lac promoter in FTL10. However, the overexpression of GadBC did not influence the growth of the cells or the induction of the PspA response when YidC was depleted (data not shown). Hence, other as-yet unidentified factors of the poorly characterized GadXY regulon likely are important causes of the observed effects. This also

### Table 3. β-Galactosidase activity (in Miller units) as directed by gad gene promoters

<table>
<thead>
<tr>
<th>Promoter</th>
<th>FTL10 + Ara</th>
<th>FTL10 − Ara</th>
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<tbody>
<tr>
<td>PgdY (−78/+65)</td>
<td>1,379</td>
<td>767 (56%)</td>
</tr>
<tr>
<td>PgdX (−222/+121)</td>
<td>1,899</td>
<td>1,150 (60%)</td>
</tr>
<tr>
<td>PgdBC (−244/+77)</td>
<td>1,767</td>
<td>910 (52%)</td>
</tr>
</tbody>
</table>

*Numbering refers to the transcriptional start site of each gene and indicates the length of the DNA regions used to generate the lacZ fusions.*
would explain why the gadBC operon was not among the plasmids selected in our genetic screen.

To rule out the hypothesis that YidC-depleted cells were experiencing acid stress, we measured the internal pH of WT, YidC-depleted, and pGadXY cells using methods previously described (34). The internal pH of wild-type cells grown in the presence of 0.2% L-arabinose was slightly acidic (pH 6.7 \pm 0.11), while cells depleted for YidC for 4 h had an internal pH of 8.1 \pm 0.57. In the presence of pGadXY and YidC depletion, the internal pH was 7.4 \pm 0.14. The internal pH values measured here would not have a suppressive effect on the growth of *E. coli* and are within the range of values reported for cells growing at neutral to mildly acidic pH. During YidC depletion, we noted no significant change in the external pH of the growth medium (i.e., pH ranged from 6.4 to 7.2), supporting the proposal that YidC-depleted cells were not experiencing either external or internal pH stress.

**YidC-depleted cells shift to fermentation.** When a low YidC level in cells is accompanied by a decrease in cytochrome bo₃ oxidase and the F₁Fₒ ATPase, the capacity of the cell to reduce O₂ would be rapidly impaired. This could affect the ratio of ubiquinone and ubiquinol (2), resulting in a switch to fermentation as a last resort to generate ATP and to oxidize NADH. In aerobic batch conditions with glucose as the sole carbon source, *E. coli* cells that lack respiratory capacity secrete lactate into the medium (1). To study this phenomenon, we determined the levels of lactate using HPLC in spent M9 medium, supplemented with 0.2% glucose, from YidC-depleted and control FTL10 cells. We found that lactate was present in medium from cells that were grown for prolonged times under YidC depletion conditions (Fig. 5). Medium from control cells did not contain lactate. Strikingly, FTL10 cells harboring pGadXY did not secrete lactate independently of the presence of YidC, indicating that the presence of pGadXY prevented the shift to fermentation, possibly by increasing the nonrespiratory NADH-oxidizing capacity.

The protein profiles of eIMs isolated from FTL10 cells grown with L-arabinose were compared to those of cells grown without L-arabinose using BN-PAGE. The gel showed clearly reduced amounts of a high-molecular-weight protein complex (Fig. 6A, indicated by an open circle) in YidC-depleted cells. To identify the proteins in this complex, it was excised from the gel and subjected to peptide mass fingerprinting. AceE, AceF, and LpdA were unambiguously identified in the complex. They form the large (4.5-MDa) multienzyme pyruvate dehydrogenase complex (PDH) that catalyzes the oxidative decarboxylation of pyruvate with the concomitant formation of acetyl-coenzyme A (CoA). Importantly, the complex is known to be downregulated upon the accumulation of NADH, again indicating that the expression of GadXY results in higher nonrespiratory NADH-oxidizing capacity (7). Another high-molecular-weight complex (Fig. 6A, indicated by an asterisk) also was identified as PDH by peptide mass fingerprinting and appeared to be reduced in eIM isolated from YidC-depleted cells, but to a lesser degree than the smaller PDH (sub)complex. In contrast, upon the overexpression of GadXY, the PDH complexes remained detectable upon YidC depletion. Taken together, these data suggest that as a result of YidC depletion, cells were unable to reoxidize the NADH pool and, therefore, showed a switch to fermentation, as evidenced by the secretion of lactate. However, the cells that overexpressed GadXY continued to form PDH complexes, indicative of NADH oxidation, and, therefore, did not show the concomitant lactate fermentation.

**GadXY overexpression leads to a decreased level of PspA complexes and increased levels of GroEL complexes.** Two
First, in a high-molecular-mass band of other observations were made from the BN-PAGE analysis. L analysis of eIMs isolated from FTL10 cells grown with or without vate dehydrogenase complex and GroEL in the IM. (A) BN-PAGE oligomerize in the main constituent of that complex by peptide mass finger-detected in the YidC-depleted eIMs, PspA was identified as 1314 YU ET AL. J. B
sates were analyzed by SDS-PAGE followed by Western blotting identified by MALDI-TOF-TOF mass spectrometry. (B) Cell ly-protein amounts were loaded on gels. The indicated bands were sho.png
molecular-weight PspA complex was absent from the eIM frac-tion obtained from YidC-depleted cells overexpressing GadXY, in accordance with data from the Western blotting of whole-cell lysates (Fig. 3A). Second, a protein complex around 0.7 MDa was strongly increased in intensity in eIMs from the YidC-depleted cells upon GadXY overexpression. It was identi-fied as the general heat shock chaperone GroEL by peptide mass fingerprinting. GroEL is a homotetradecamer of 57-kDa subunits that are arranged as two heptameric rings stacked back to back (20). Western blot analysis showed that YidC-depleted cells had slightly upregulated GroEL content, while GadXY-overexpressing cells had strongly increased levels of GroEL compared to the unaltered level of the YidC-inde-pendent IMP leader peptidase (Lep) (Fig. 6B). Although GroEL is not known to be part of the GadX regulon, it has been re-portd that GadY overexpression results in the upregulation of GroEL (25). However, the sole overexpression of GroEL and GroES from an inducible expression plasmid appeared insufficient to restore growth upon YidC depletion (results not shown).

FIG. 6. Effect of GadXY overproduction on the levels of pyru-vate dehydrogenase complex and GroEL in the IM. (A) BN-PAGE analysis of eIMs isolated from FTL10 cells grown with or without l-arabinose in the presence or absence of pGadXY. Comparable protein amounts were loaded on gels. The indicated bands were identified by MALDI-TOF-TOF mass spectrometry. (B) Cell ly-sates were analyzed by SDS-PAGE followed by Western blotting using antisera against GroEL and Lep.

other observations were made from the BN-PAGE analysis. First, in a high-molecular-mass band of >1.2 MDa, which was detected in the YidC-depleted eIMs, PspA was identified as the main constituent of that complex by peptide mass fingerprinting. Consistently, purified PspA has been shown to oligomerize in ~1-MDa complexes (12). The diffuse, high-molecular-weight PspA complex was absent from the eIM fraction obtained from YidC-depleted cells overexpressing GadXY, in accordance with data from the Western blotting of whole-cell lysates (Fig. 3A). Second, a protein complex around 0.7 MDa was strongly increased in intensity in eIMs from the YidC-depleted cells upon GadXY overexpression. It was identi-fied as the general heat shock chaperone GroEL by peptide mass fingerprinting. GroEL is a homotetradecamer of 57-kDa subunits that are arranged as two heptameric rings stacked back to back (20). Western blot analysis showed that YidC-depleted cells had slightly upregulated GroEL content, while GadXY-overexpressing cells had strongly increased levels of GroEL compared to the unaltered level of the YidC-inde-pendent IMP leader peptidase (Lep) (Fig. 6B). Although GroEL is not known to be part of the GadX regulon, it has been re-portd that GadY overexpression results in the upregulation of GroEL (25). However, the sole overexpression of GroEL and GroES from an inducible expression plasmid appeared insufficient to restore growth upon YidC depletion (results not shown).

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DISCUSSION

YidC is an essential E. coli IMP that plays an important but poorly defined role in the biogenesis of IMPs (18, 52). So far, targeted approaches have revealed a plethora of YidC func-tions that seem almost too varied to be combined in one protein. Here, we have screened a random genetic library to identify factors that improve the survival of YidC-depleted cells and may reveal why YidC is an essential protein. The gadX and gadY genes were selected multiple times in the screen and were found to partly compensate for the growth defect and the PspA response that accompanies YidC deple-tion (46). The genes encode regulatory proteins of the major acid resistance system in E. coli (25, 36, 38), hinting at an unexpected link between YidC and the acid stress response.

The gadX gene encodes a transcriptional regulator belonging to the AraC family, while gadY, encoding a small ncRNA, stabilizes the gadX transcript (25, 36). GadX regulates tran-scription of the glutamate decarboxylase isozymes GadA and GadB, which convert glutamate to GABA while consuming a proton from the cytoplasm (53). The antipporter GadC expels the decarboxylated products in exchange for glutamate. The resulting net export of protons is thought to increase the internal pH to sustain membrane integrity, normal physiological processes, and protein stability. The control of this acid resis-tance system is reported to involve at least 15 regulatory fac-tors whose functions respond to the environmental pH, growth phase, medium composition, and oxygen levels. The regulators include the AraC-like transcription factors GadX, GadW, and YdeO; the LuxR-like transcriptional regulator GadE; the two-component regulatory system EvgAS; the alternative sigma factor RpoS; TorRS; Crp; H-NS; Dps; TrmE; and three small ncRNAs, GadY, DsrA, and GcvB (11, 15, 16, 19, 21, 29, 40). Why so many regulators and control circuits are needed is not clear, but it probably reflects a need to connect the induction of this system to many facets of cell physiology. The complexity is underscored by the fact that the regulon of GadX contains many genes that have no direct link with acid resistance (15). In addition, GadX enhances multidrug resistance by activating the mdtEF efflux genes (24). Notably, the Gad response is not exclusively a consequence of acidic stress but also occurs upon salt stress and in the stationary growth phase (6).

Considering this complexity, it is intriguing that gadXY was repeatedly selected in our screen. Other regulators of the acid stress response, such as gadE or gadW, were not selected, although our screen may not have been exhaustive. Furthermore, members of the GadX regulon, like GadBC, also were not selected, nor were other genes encoding amino acid decarboxylases and acid stress regulators (data not shown). Apparently, only the overexpression of GadXY leads to a re-sponse of effector proteins that together have a beneficial effect on YidC-depleted cells. Consistently, while we demon-strated the upregulation of the main resistance effectors in response to acid stress, GadBC alone could not compensate for the growth defect upon YidC depletion.

We ruled out that the rescuing effect of the pGadXY plas-mid was due to a direct influence on YidC expression or sta-bility. Furthermore, GadXY overexpression did not improve the processing of the YidC-dependent IMP CyoA, nor were the levels of the cytochrome bo oxidase restored, suggesting
that the insertase function of YidC was not restored as a result of GadXY overexpression. Although we cannot exclude that the Sec-associated YidC functions are improved by GadXY, these functions are less important to sustain cell growth (41, 45). Therefore, it is more likely that GadXY overexpression alleviates negative secondary effects of the absence of the essential YidC protein.

Previous studies have shown that YidC-depleted cells suffer from a reduced PMF, which triggers a strong PspA response. This is probably a consequence of the fact that YidC is strictly required for the membrane insertion of CyoA and F_{1}F_{o} ATPase, respectively. The absence of these proteins may result in the accumulation of NADH. Indicative for the accumulation of NADH is the strongly regulated multi-enzyme PDH complex that connects glycolysis with the tricarboxylic acid cycle, which was much less abundant in YidC-depleted cells. This limits the capacity of the cell to convert pyruvate to acetyl-CoA, and therefore the only remaining alternative to generate ATP, although less efficient, is a shift to lactic acid fermentation. Indeed, we detected the secretion of lactate in YidC-depleted cells, indicating that such a shift had occurred. In contrast, cells that overproduced GadXY from the pGadXY plasmid retained higher levels of the PDH complex and did not secrete lactate. However, it remains unclear how the switch to fermentation is prevented in the GadXY-overexpressing cells.

An explanation of the beneficial effect of GadXY on YidC-depleted cells might lie in a compensation of intracellular acidification as a consequence of fermentation, which produces weak acids such as acetate and lactate. Under normal circumstances, the F_{1}F_{o} ATPase would counter intracellular acidification by pumping out protons at the expense of ATP. However, in YidC-depleted cells the F_{1}F_{o} ATPase is compromised too. Intracellular pH measurements indicated that YidC-depleted cells were not suffering from internal acid stress. In contrast, the intracellular pH was more alkaline than that of the WT and returned to neutral values in the presence pGadXY. Consistently, the sole overproduction of GadBC, the effectors that are upregulated by GadXY to bring about a net export of protons, could not compensate for the growth reduction upon the depletion of YidC, supporting the hypothesis that the inhibition of growth was not related to cytoplasmic acidification.

It also has been proposed that the exchange of GABA for glutamate contributes to a functional PMF (14, 34), explaining the reduced PspA response in the YidC-depleted but GadXY-overexpressing cells. However, as mentioned, the upregulation of the glutamate-dependent acid response system is unlikely to be the only reason for the improved fitness of YidC-depleted cells. Other genes in the GadX regulon probably are involved as well. Interestingly, genes encoding chaperones and proteases (ycgG, yehA, yhcA, and lon) also are induced by GadX (15). In addition, the heat shock chaperone GroEL copurified with membranes and was upregulated upon GadXY overexpression (Fig. 6). GroEL is known to act on a wide range of unfolded or partially folded proteins, and with the assistance of GroES, it helps them to reach their fully folded and active state (20). GroEL levels are increased upon YidC depletion, and it has been shown that part of GroEL is redistributed to the membrane under these conditions (48). When YidC levels are low, GroEL might be recruited to the IM through interaction with IMPs that are targeted but not yet inserted or by binding to unfolded domains of improperly assembled IMPs that accumulate near the membrane. GroEL may keep such unfolded proteins in an insertion-competent state while they anticipate their insertion into the membrane by the limited amount of YidC available. In addition, GroEL has been implicated in the targeting of proteins to the Sec translocon based on its affinity for SecA (3). The further upregulation of GroEL expression by GadY ncRNA (either directly or indirectly via GadX) (25) might increase the fitness of YidC-depleted cells by preventing IMPs from aggregating near the membrane. This eventually could allow them to insert into the membrane, albeit at a lower efficiency. Overall, we have revealed here, for the first time, that the lethality of YidC depletion may involve the acid stress response, the response to a drop in PMF, and protein quality control. The intricate relationship between YidC and these processes will be the subject of further study.

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