The architecture of the OmpC–MlaA complex sheds light on the maintenance of outer membrane lipid asymmetry in *Escherichia coli*

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A distinctive feature of the Gram-negative bacterial cell envelope is the asymmetric outer membrane (OM), where lipopolysaccharides and phospholipids (PLs) reside in the outer and inner leaflets, respectively. This unique lipid asymmetry renders the OM impermeable to external insults, including antibiotics and bile salts. In *Escherichia coli*, the complex comprising osmoporin OmpC and the OM lipoprotein MlaA is believed to maintain lipid asymmetry by removing mislocalized PLs from the outer leaflet of the OM. How this complex performs this function is unknown. Here, we defined the molecular architecture of the OmpC–MlaA complex to gain insights into its role in PL transport. Using in vivo photo-cross-linking and molecular dynamics simulations, we established that MlaA interacts extensively with OmpC and is located entirely within the lipid bilayer. In addition, MlaA forms a hydrophilic channel, likely enabling PL translocation across the OM. We further showed that flexibility in a hairpin loop adjacent to the channel is critical in modulating MlaA activity. Finally, we demonstrated that OmpC plays a functional role in maintaining OM lipid asymmetry together with MlaA. Our work offers glimpses into how the OmpC–MlaA complex transports PLs across the OM and has important implications for future antibacterial drug development.

The outer membrane (OM)⁴ of Gram-negative bacteria is an extremely asymmetric bilayer, comprising lipopolysaccharides (LPS) in the outer leaflet and phospholipids (PLs) in the inner leaflet (1, 2). LPS molecules pack tightly together in the presence of divalent cations to form an outer layer with markedly reduced fluidity and permeability (3). Thus, the OM serves as an effective barrier against toxic compounds including detergents and antibiotics. This function is fully dependent on the establishment and maintenance of lipid asymmetry; cells generally become more sensitive to external insults when OM lipid asymmetry is disrupted, which is typically characterized by the accumulation of PLs in the outer leaflet (4, 5). The OM is also essential for viability.

The requisite lipid asymmetry of the OM is likely initially established by direct placement of LPS and PLs into the outer and inner leaflets, respectively. LPS assembly into the outer leaflet of the OM is mediated by the well-established Lpt (lipopolysaccharide transport) machinery (6), but proteins that transport and insert PLs into the inner leaflet have not been identified. For entropic reasons, there is a natural tendency for PLs to appear in the outer leaflet of the OM, although how they traverse the bilayer is unclear. This occurs more readily with perturbations in the OM, especially when assembly of other OM components is disrupted (4, 5, 7). Because loss of lipid asymmetry compromises the barrier function of the OM, several mechanisms exist to remove PLs aberrantly localized in the outer leaflet of the membrane: (i) the OM phospholipase OmpLA hydrolyzes both acyl chains from outer leaflet PLs (8), (ii) the OM acyltransferase PagP transfers an acyl chain from outer leaflet PLs to LPS (9) or phosphatidylglycerol (10), and (iii) the OmpC–MlaA system, a putative PL trafficking pathway,
removes outer leaflet PLs and shuttles them back to the inner membrane (IM) (11, 12).

The OmpC–Mla system comprises seven proteins located across the cell envelope. Removing any component results in PL accumulation in the outer leaflet of the OM and therefore sensitivity to SDS/EDTA (11, 12). The OM lipoprotein MlaA forms a complex with osmoporin OmpC that likely extracts PLs from the outer leaflet of the OM (12). The periplasmic protein MlaC serves as a lipid chaperone and is proposed to transport lipids from the OmpC–MlaA complex to the IM (11, 13, 14). At the IM, MlaF and MlaE constitute an ATP-binding cassette family transporter together with two auxiliary proteins, MlaD and MlaB (14, 15); this complex presumably receives PLs from MlaC and inserts them into the membrane. MlaD has been shown to bind PLs, whereas MlaB is important for both assembly and activity of the transporter (15). Recently, the function of the OmpC–Mla system in retrograde (OM-to-IM) PL transport has been demonstrated in *Escherichia coli* (7).

The molecular mechanism by which the OmpC–MlaA complex extracts PLs from the outer leaflet of the OM, presumably in an energy-independent manner, is an interesting problem. Aside from the LptDE machine, which assembles LPS on the surface (4, 5), the OmpC–MlaA complex is the only other system proposed to catalyze the translocation of lipids across the OM. OmpC is a classical trimeric porin that typically only allows passage of hydrophilic solutes across the OM, although MlaA is believed to be anchored to the inner leaflet of the membrane; how the two proteins are organized in a complex for the translocation of amphipathic PLs is not known. In this paper, we establish that MlaA is in fact an integral membrane protein that forms a channel adjacent to OmpC trimers in the OM, likely allowing the passage of PLs. We first demonstrated that MlaA binds the OmpC trimer within the OM bilayer by mapping the interaction surfaces using *in vivo* cross-linking. Using a recently predicted structural model of MlaA (17), we obtained molecular views of the OmpC–MlaA complex by molecular dynamics (MD) simulations, and experimentally established the existence of a hydrophilic channel within OM-embedded MlaA. Combining charge mutations in this channel modulated MlaA activity, suggesting functional importance. Furthermore, mutations altering the flexibility of a hairpin loop that could interact with the hydrophilic channel led to predictable *in vivo* effects on MlaA function. Finally, we identified a key residue on OmpC found at the OmpC–MlaA interacting surface that is important for proper function of the complex. Our findings provide important mechanistic insights into how PLs may be translocated across the OM to ensure proper lipid asymmetry.

**Results**

**The OmpC trimer contacts MlaA directly along its membrane-facing dimeric interfaces**

To develop a detailed architectural understanding of the OmpC–MlaA complex, we carried out *in vivo* photo-cross-linking to map intermolecular interactions within the complex. Guided by the crystal structure of the OmpC trimer (18), we introduced the UV-cross-linking amino acid, *para*-benzoyl-l-phenylalanine (pBpa), at 49 positions in OmpC via amber suppression (19). Initial selection focused on residues that are either solvent-accessible (*i.e.* loop and lumen) or located near the membrane–water boundaries (*i.e.* aromatic girdle). Upon UV irradiation, a ~65-kDa cross-linked band that contains both OmpC (~37 kDa) and MlaA (~28 kDa) could be detected in cells expressing OmpC variants substituted with pBpa at three positions (Leu50, Gln83, or Phe267) (Fig. 1A). These residues are found on the periplasmic turns at the dimeric interfaces of the OmpC trimer (Fig. 1C), thus localizing possible binding sites for MlaA. We have previously proposed that OmpC may allow MlaA to traverse the bilayer and gain access to PLs that have accumulated in the outer leaflet of the OM (12). Because none of the six selected residues in the OmpC lumen cross-linked to MlaA, we decided to probe for interactions between MlaA and the membrane-facing side walls of OmpC, specifically around the dimeric interfaces of the OmpC trimer. Remarkably, of the additional 49 positions tested in this region, 10 residues allowed photoactivated cross-links between OmpC and MlaA when replaced with pBpa (Fig. 1B and Fig. S1). In total, these 13 cross-linking residues clearly demarcate an extensive MlaA-interacting surface on OmpC (Fig. 1C). This explains why OmpC exhibits strong interactions and can be co-purified with MlaA on an affinity column, as we have previously reported (12). Two positions, Tyr149 and Leu340, are located right at or near the membrane–water boundary exposed to the extracellular environment, suggesting that MlaA traverses the entire width of the OM. We conclude that MlaA binds along the dimeric interfaces of the OmpC trimer in the membrane.

**MlaA resides entirely in the OM bilayer and contacts the OmpC trimer via two specific regions**

We sought to map in greater detail the OmpC–MlaA interacting surface *in vitro*. To do that, we first overexpressed and purified the OmpC–MlaA complex to homogeneity. We showed that this complex forms a single peak on size-exclusion chromatography (SEC) (Fig. 2A). OmpC within this complex exhibits the characteristic heat-modifiable gel shift commonly observed for OM β-barrel proteins (20), consistent with the presence of the folded trimer. Multitangle light scattering (MALS) analysis revealed that one copy of MlaA interacts with the OmpC trimer (Fig. S2), suggesting that only one of the three dimeric interfaces within the trimer is available for binding (Fig. 1C). We next performed protease digestion experiments to identify specific region(s) on MlaA that may interact stably with OmpC. OM β-barrel proteins such as OmpC are known to be protease-resistant (21). Given that some parts of MlaA contact OmpC within the membrane, we expect these bound regions to also be protected from proteolytic degradation. Treatment of the purified OmpC–MlaA complex with trypsin results in almost complete degradation of MlaA, with the OmpC trimer remaining intact (Fig. 2A). Following SEC, however, we found that an ~8-kDa peptide (presumably from MlaA) remains stably bound to the trimer. N-terminal sequencing and tandem MS analyses revealed that this peptide corresponds to MlaA<sub>D61–K124</sub> (Fig. 2, A and C, and Fig. S3). These results sug-
gest that MlaA interacts strongly with OmpC in the membrane in part via this specific region.

To define how OmpC contacts the MlaAD61–K124 peptide, we next attempted to overexpress and purify \( p \)Bpa-containing OmpC variants in complex with MlaA and determine which of the previously identified 13 OmpC residues interacts with MlaAD61–K124 in vitro. We sequentially performed UV cross-linking and trypsin digestion to potentially link MlaAD61–K124 to specific residues on OmpC. This approach may also allow trapping of other potential interacting regions of MlaA, which might not have been stably retained on the WT complex after trypsin digestion. We successfully detected trypsin-resistant cross-linked products for seven OmpC\(_{pBpa}\)–MlaA complexes (Fig. 2B); they appeared slightly above OmpC between 37 and 50 kDa, indicating that peptides in the range of \( \sim 6–10 \) kDa are cross-linked to OmpC. N-terminal sequencing of six of the seven adducts showed the presence of OmpC and an MlaA peptide beginning at residue Asp\(^{61}\) (Fig. 2, C and D, and Fig. S3C). Given the approximate sizes of the cross-linked adducts, we concluded that all these residues interact with MlaAD61–K124. Interestingly, an additional peptide on MlaA starting at residue Phe\(^{133}\) was also found to cross-link at two (Tyr\(^{149}\) and Leu\(^{340}\)) of these six positions on OmpC (Fig. 2, B–D, and Fig. S3C). These adducts can be detected by an \( \alpha \)-MlaA antibody that recognizes an epitope within Val\(^{182}\)–Gln\(^{195}\) on MlaA (Fig. 2B), suggesting that a peptide from Phe\(^{133}\) to at least Arg\(^{205}\) (next trypsin cleavage site) may be cross-linked (Fig. 2C). Thus, in addition to MlaAD61–K124, our cross-linking strategy revealed a second point of contact (MlaAF\(_{133}–R205\)) between OmpC and MlaA.

At the point of these findings, there was no available molecular structure for MlaA; however, a structural model has been predicted based on residue–residue contacts inferred from co-evolution analysis of metagenomic sequence data (17). Using a
rigorously validated quality score, this method of structure
determination has generated reliable models for 614 protein
families with currently unknown structures. We experimen-
tally validated the model for MlaA by replacing residue pairs
far apart on the primary sequence with cysteines and showed
that only those that are highly co-evolved (and predicted to
be residue–residue contacts) allow disulfide-bond formation
in cells (Fig. S4). We therefore proceeded to use this MlaA
model to understand the organization of the OmpC–MlaA
complex. Interestingly, the positions of the two OmpC-con-
tacting peptides on the MlaA model are spatially separated
in a way consistent with the arrangement of the residues on
OmpC that cross-link to each peptide (Fig. 2D). This not
only reveals how MlaA may potentially be oriented and orga-
nized around the dimeric interface of the OmpC trimer but
also suggests that the entire MlaA molecule may reside in
the membrane. In fact, the overall surface of MlaA, other than
the putative periplasmic-facing region, is largely hydropho-
bic (Fig. S5). Furthermore, using all-atomistic MD simula-
tions, we found that the structural fold of MlaA appears to be
more stable in a lipid bilayer than in an aqueous environment
(Fig. S6). Consistent with this, we noted that even without its
N-terminal lipid anchor, MlaA is largely not extracted from
total membranes by short washes of high pH, chaotropic, or
mild detergent solutions (Fig. S7); its membrane extraction
profile is similar to /H9252–barrel proteins but different from
known peripheral membrane proteins, indicating that MlaA
is stably inserted in the OM as an integral membrane pro-
tein. Collectively, these observations lend strong support to
the validity of the predicted MlaA structure.

**Figure 2. OmpC contacts two specific regions on MlaA.**

A, SEC profiles and SDS–PAGE analyses of purified OmpC–MlaA–His complex before (black) or
after (red) treatment with trypsin. Peak fractions from SEC were subjected to denaturing SDS–PAGE (15% Tris-HCl gel), followed by Coomassie Blue (CB) staining (right panel). Non-trypsin-treated samples were also analyzed by seminative SDS–PAGE (left panel). Edman degradation and tandem MS analyses revealed that the MlaA peptide that remains bound to OmpC following trypsin treatment begins at Asp61 (Fig. S3).

B, SDS–PAGE (15% Tris-HCl gel) and immunoblot analyses of purified OmpC<sub>p</sub>Bpa–MlaA–His complexes following sequential UV irradiation and trypsin digestion. The resulting OmpC<sub>p</sub>Bpa–MlaA<sub>p</sub>Bpa cross-linked products were N-terminally sequenced (see Fig. S3). C, amino acid sequence of MlaA with the two peptides found
cross-linked to OmpC<sub>p</sub>Bpa, highlighted (red: MlaA<sup>D61–K124</sup>, blue: MlaA<sup>F133–R205</sup>). The signal sequence and /H9251–MlaA binding epitope are
derlined and annotated. D, cartoon representations of the crystal structure of *E. coli* OmpC with positions that cross-link to specific MlaA peptides indicated (left) and
a structural model of MlaA (17) with peptides cross-linked by OmpC<sub>p</sub>Bpa highlighted (right). The OM boundary is indicated as gray dashed lines. The
figures were generated using the program Chimera (54).
MlaA forms a channel in the OM adjacent to OmpC

To obtain a physical picture of how OmpC interacts with MlaA within the complex, we used MD simulations to dock MlaA onto the OmpC trimer within a PL bilayer. Using a previously reported protocol (22), we first docked the MlaA model onto the OmpC trimer structure, both as rigid bodies. Interestingly, all initial docked structures contained MlaA binding at one dimeric interface of OmpC. Based on information derived from cross-linking, we selected two most consistent models differing slightly in how MlaA is oriented with respect to OmpC for unrestrained refinements using all-atomistic simulations. Multiple simulations were run for each MlaA orientation in a PL bilayer until overall root-mean-square deviations (RMSDs) stabilized; remarkably, the resulting equilibrium models fulfilled all observed experimental cross-linking data. We performed clustering on all available trajectories and identified the most populated conformations of the OmpC–MlaA complex in our simulations (Fig. 3A and Fig. S8A for one MlaA orientation and Fig. 3B and Fig. S8B for the other). These conformational models all show MlaA sitting in the bilayer, tucked nicely into the dimeric interface of the OmpC trimer. Evidently, the MlaA_{D61–K124} peptide interacts extensively with OmpC in these models, consistent with why this peptide remains stably bound to OmpC after protease digestion (Fig. 2A). We also mutated several MlaA residues found at the OmpC–MlaA interfaces to pBpa and performed in vivo cross-linking experiments. We identified one position, Leu^{109}, that allowed strong photoactivatable cross-linking to OmpC when replaced with pBpa (Fig. 3C). This residue lies within the MlaA_{D61–K124} peptide (Fig. 3, A and B), confirming that this region does in fact contact OmpC in cells.

One striking feature present in all the simulated OmpC–MlaA models is a negatively charged hydrophilic channel within MlaA that spans the lipid bilayer (Fig. 4A and Fig. S9). To test the existence of this hydrophilic channel in MlaA, especially in the context of the native OM, we selected 27 residues in and around the putative channel in a representative model (Fig. 4 and Fig. S10) and determined their solvent accessibility in cells using the substituted cysteine accessibility method (SCAM). We first showed that these cysteine mutants are functional (Fig. S10C). Solvent-exposed residues are expected to be reactive with the charged membrane-impermeable thiol-labeling reagent sodium {2-sulfonatoethyl}-methanethiosulfonate (MTSES). Remarkably, we found that residues predicted to be within the putative channel (Fig. 4, A and B) or at the membrane–water boundaries (Fig. S10, A and B) are indeed accessible to MTSES. We therefore conclude that MlaA forms a hydrophilic channel in the OM in cells.

Very recently, crystal structures of MlaA in complex with trimeric porins have in fact been solved (23). Interestingly, the experimentally determined structures of MlaA closely resembled the initial MlaA model predicted from co-evolution analysis (Fig. S11A). These structures revealed that MlaA interacts with trimeric porins in the membrane at one or more of their dimeric interfaces (Fig. S11B), in an orientation similar to one of our simulated OmpC–MlaA models (Fig. 3B). They also showed that MlaA contains a hydrophilic channel. Although these structural and our biochemical/modeling studies are in good agreement, there are some discrepancies between the solved structures and our biochemical data. Specifically, the distances separating porin residues equivalent to Tyr^{149}/Leu^{280} in OmpC and the MlaA_{F133–R205} peptide in the solved structures (Fig. S11B) are not consistent with the detection of strong in vivo photoactivatable cross-links between these regions in the complex (Fig. 2B). In addition, the proposed locations of some MlaA residues, specifically Met^{39}, Phe^{42}, and Asn^{43}, at the interior of the lipid bilayer (Fig. S11B) are not in agreement with high or partial solvent accessibility of these sites in cells, as highlighted in the SCAM data (Fig. S10B). These observations suggest that MlaA might adopt a different conformation in its native environment in the OM than that in a crystal lattice.

Figure 3. Molecular models of the OmpC–MlaA complex depict how MlaA may interact with OmpC in the OM bilayer. A and B, representative MlaA structures bound to OmpC in two possible orientations selected from all-atomistic MD simulation trajectories. MlaA_{D61–K124} and MlaA_{F133–R205} peptides are highlighted in red and blue, respectively, as in Fig. 2D. Leu^{109} is labeled and depicted as black spheres on MlaA. The OM boundaries are indicated as gray dashed lines. The figures were generated using the program VMD (55). C, immunoblots showing UV-dependent formation of a cross-link between MlaA and OmpC in ∆mlaA cells expressing MlaA_{D61–K124}–His. As expected, the cross-linked product also exhibits heat-modifiable gel shift, indicative of the presence of OmpC.
The MlaA channel is functionally important for OM lipid asymmetry

Based on its function in removing PLs from the outer leaflet of the OM, we hypothesize that this channel may allow passage of charged head groups as PLs translocate across the membrane. To ascertain whether this channel is functionally important, we separately mutated 19 polar and (negatively) charged residues near or within the channel to alanine or arginine and tested for MlaA function. However, all of these MlaA mutants are functional (Fig. 5B and Fig. S12A), indicating that single residue changes are not sufficient to perturb channel properties to affect PL transport. We noticed that four (Asp$^{160}$, Asp$^{161}$, Asp$^{164}$, and Asp$^{167}$) of the five negatively charged channel residues are in close proximity (Fig. 5A). To alter channel properties more drastically, we combined arginine mutations for these aspartates; interestingly, only the D161R/D167R double mutant (Fig. S12B) and the D160R/D161R/D164R triple mutant (MlaA$_{3D3R}$) disrupted function; cells expressing these variants are highly sensitive to SDS/EDTA (Fig. 5B and Fig. S12B). For the MlaA$_{3D3R}$ mutant, this is likely due to the accumulation of PLs in the outer leaflet of the OM (as judged by PagP-mediated acylation of LPS; Fig. 5C). In fact, this mutant exhibits OM defects that are more pronounced than the ΔmlaA strain (Fig. 5B). Furthermore, the 3D3R mutation gives rise to the same defects in strains also expressing the WT mlaA allele, revealing a dominant-negative phenotype (Fig. 5, B and C). We showed that MlaA$_{3D3R}$ is produced at levels comparable with WT MlaA on a plasmid (Fig. S13A) and is still able to interact strongly with OmpC (Fig. S13B). Taken together, these results suggest that the MlaA channel plays a functional role in the maintenance of lipid asymmetry.

Flexibility of a hairpin loop adjacent to the channel in MlaA is required for function

The dominant-negative phenotype of the mlaA$_{3D3R}$ mutant is similar to a previously reported mlaA* (or mlaA$_{SNP}$) mutant (Fig. 5, B and C) (24), suggesting that these mutations may have similar effects on MlaA structure and/or function. Interestingly, the positions of these mutations on the OmpC–MlaA models flank a hydrophobic hairpin loop (Gly$^{141}$–Leu$^{158}$) within MlaA (Fig. 6A). Therefore, we hypothesized that the loop could play a functional role in MlaA and that these mutations may affect interactions with this loop. To examine this possibility, we created three separate mutations at the hairpin structure and tested each variant for MlaA function. Two of these mutations, $^{147}$YVQL$^{154}$A (L1) and $^{152}$FYGSF$^{159}$A (L2),
are designed to disrupt interactions with other regions of MlaA. The other mutation, P151A, removes a proline that may be critical for the hairpin turn structure. The N terminus of the loop is connected to the rest of MlaA via an unstructured glycin-rich linker, which we reasoned may influence conformation of the entire hairpin structure. Thus, we constructed two additional mutants, \texttt{141GVGYG} \texttt{141AVAYA (3G3A)} and \texttt{141GVGYG} \texttt{141PVPYP (3G3P)}, to reduce possible flexibility in this region. Remarkably, \texttt{L1}, \texttt{L2}, and \texttt{3G3P} mutations resulted in similar extents of SDS/EDTA sensitivity (Fig. 6B), as well as OM outer leaflet PL accumulation (Fig. 6C), when compared with the \texttt{Delta mlaA} mutation. Given that these mutations also do not affect MlaA levels or interaction with OmpC (Fig. S13), we conclude that they are loss-of-function mutations. The hairpin loop, along with its surrounding structures, forms an important functional region on MlaA.

Phenotypes observed for the loop rigidifying mutation (3G3P) and dominant-negative mutations (3D3R and mlaA*)
suggest that flexibility in the hairpin loop is critical for MlaA function. We hypothesize that the hairpin loop may exist in two distinct conformations. The 3D3R or mlaA* mutations could alter interactions with the loop, resulting in it adopting one conformation and somehow giving rise to exacerbated and thus dominant-negative phenotypes; in the case of mlaA*, it was
proposed that this mutation caused MlaA to be in a “leaky” or “open” state and allowed PLs to flip out to the outer leaflet of the OM (24). In contrast, the 3G3P mutation may lock the hairpin loop in a second conformation, where MlaA is in a “closed” state, thus abolishing function in PL transport. If these were true, we predict that rigidification of the hairpin loop with the 3G3P mutation would be able to correct dominant-negative phenotypes observed in the 3D3R and/or mlaA* mutants. Indeed, the 3G3P/3D3R and 3G3P/mlaA* combination mutants no longer exhibit dominant-negative phenotypes but behave like the 3G3P or null mutants (Fig. 6D). Again, these variants are expressed at comparable levels to the single mutants and still interact with OmpC (Fig. S13). Taken together, these results indicate clear importance of dynamics in the hairpin loop in controlling the function of MlaA.

A specific residue in the dimeric interface of OmpC is important for its function in maintaining lipid asymmetry

Given that MlaA sits in the membrane and provides a channel that putatively allows movement of PLs across the OM, it is not clear why it should bind at one of the dimeric interfaces of OmpC trimers and what the exact role of OmpC may be. To understand the importance of OmpC–MlaA interaction, we attempted to engineer monomeric OmpC constructs that we predict would no longer interact with MlaA. We installed specific mutations (G19W and/or R92L) in OmpC that were found previously to disrupt the oligomerization state of its homolog OmpF (G19W and R100L correspondingly) in vitro (25) (Fig. 7A). Both the OmpC<sub>G19W</sub> and OmpC<sub>R92L</sub> single mutants can interact with MlaA and still form trimers in vitro, albeit slightly destabilized compared with WT OmpC (Fig. S14). Combination of these mutations further weakens the OmpC trimer, with noticeable monomer population at physiological temperature. Intriguingly, both the double mutant and the R92L single mutant accumulated PLs in the outer leaflet of the OM (Fig. 7B), indicating that Arg<sup>92</sup> is important for the role of OmpC in OM lipid asymmetry. Consistent with this idea, we demonstrated that cells expressing the OmpC<sub>R92A</sub> variant also exhibit perturbations in OM lipid asymmetry. We further showed that all these ompC alleles can rescue severe SDS/EDTA sensitivity known for cells lacking OmpC (Fig. 7C), suggesting normal porin function; however, cells expressing OmpC<sub>G19W/R92L</sub> and OmpC<sub>R92A</sub>, unlike WT and OmpC<sub>R92L</sub>, are still sensitive to SDS at higher concentrations of EDTA. These phenotypes mirror those observed for cells lacking MlaA, suggesting the loss of Mla function in these mutant strains. It appears that the Arg<sup>92</sup> residue is critical for this function, although it is not clear why the single R92L mutation did not result in SDS/EDTA sensitivity. The exact role of Arg<sup>92</sup> is not known, but the phenotypes observed for the single R92A mutation cannot be due to disruption of OmpC trimerization (Fig. S14). We conclude that OmpC has a functional role in maintaining OM lipid symmetry together with MlaA.

Discussion

Osmoporin OmpC interacts with MlaA to maintain lipid asymmetry in the OM; how this complex is organized to extract PLs from the outer leaflet of the OM is not known. In this study, we have employed photoactivatable cross-linking and MD simulations to gain insights into the molecular architecture of the OmpC–MlaA complex. We have established that MlaA interacts extensively with OmpC at one of the dimeric interfaces of the porin trimer and resides entirely within the OM lipid bilayer. We have also demonstrated that MlaA forms a hydrophilic channel, likely allowing PLs to translocate across the OM. This overall organization of the OmpC–MlaA complex is quite remarkable, especially how MlaA spans the OM and possibly gains access to outer leaflet PLs. Very few lipoproteins are known to span the OM; some notable examples include the LptDE and Wza translocons, which transport LPS and capsular polysaccharides, respectively. In the LptDE complex, the OM lipoprotein LptE serves as a plug and stretches across the bilayer through the lumen of the LptD β-barrel (26, 27). In the octameric Wza translocon, each protomer provides a C-terminal α-helix to form a pore that spans the membrane (28). MlaA is unique in that it is essentially an integral membrane protein, capable of forming a channel on its own. In many ways, MlaA behaves like typical OM β-barrel proteins, even though it is predominantly α-helical. Being overall a hydrophobic protein also poses a problem for MlaA to transit across the periplasmic space. How MlaA is shielded from the aqueous environment, in addition to the requirement of the Lol system (29), necessitates further investigation.

The recently solved crystal structures of MlaA in complex with trimeric porins revealed findings that converged with the main conclusions derived from our biochemical and modeling studies. It is quite remarkable that our studies effectively validate each other. We have noted, however, a couple of inconsistencies between the solved structures and our biochemical data, specifically in vivo cross-linking and SCAM (Fig. S11B). These discrepancies suggest that the crystal structure may represent only one of several possible stable conformations of MlaA, which may exist as part of the mechanism associated with lipid transport in the native OM environment. Upon close examination of these structures, we found multiple nonnative MlaA–OmpC and MlaA–porin contacts in all the crystal forms (Fig. S11C); these artificial crystal contacts, some with substantial buried surface areas, may have influenced the observed conformation of MlaA. Given that the MlaA–porin structures were
also not solved in the context of a native lipid bilayer, we suspect that they do not yet provide the complete picture.

Our functional data on the OmpC–MlaA complex provide a glimpse of how PLs may translocate across the OM during maintenance of lipid asymmetry. One key aspect of MlaA function resides in a hairpin loop structure juxtaposed against the hydrophilic channel. Dynamics of this loop appear to control whether MlaA exists in a "closed" or "open" state and thus access of PLs through the channel. A mutation that likely rigidifies the loop locks MlaA in the nonfunctional closed state (Fig. 6), whereas mutations that possibly affect interactions with the loop favors the open state, and gives rise to exacerbated phenotypes (Fig. 5). Therefore, the hairpin loop may directly gate the channel, providing access for outer leaflet PLs across the OM. This idea has also been suggested based on the solved MlaA structures, where a putative disulfide bond that apparently locks the loop in position renders MlaA nonfunctional (23). It is likely that the hydrophilic channel provides a route for the passage of PL head groups across the membrane; what is not clear, however, is how the acyl chains pass through the MlaA protein. The overall path for PLs across MlaA remains to be determined.

How OmpC participates in maintaining OM lipid asymmetry as part of the complex is not clear, especially given that MlaA alone provides a channel for PL translocation. It is possible that OmpC may play a passive role and simply be important for stabilizing the structure and orientation of MlaA in the OM. We have previously shown that MlaA also interacts with OmpF, which is expressed at comparable levels to OmpC under low osmolarity conditions (12). Removing OmpC alone, whether under high or low osmolarity conditions, causes perturbations in OM lipid asymmetry; however, removing OmpF alone has minimal effects. These results argue against a mere passive role for OmpC in PL translocation. Furthermore, we have now identified a specific residue, Arg^92, on OmpC that is required for the maintenance of OM lipid asymmetry (Fig. 7). Therefore, we believe that OmpC plays a key functional role in the process. Arg^92 lies in the dimeric interface of the OmpC trimer, which incidentally is where MlaA binds. Even though
this residue has been shown to be important for gating the porin (30), it is not obvious how this gating function may influence the translocation of PLs by MlaA. Instead, we speculate that being close to Tyr149 and Leu340, Arg92 may somehow affect interactions between OmpC and the MlaA<sub>133–205</sub> peptide, possibly influencing MlaA conformation in the OM and, in turn, properties of the hydrophilic channel. The role that Arg<sup>92</sup> plays in OmpC–MlaA function should be further characterized.

The OmpC–MlaA complex is proposed to extract PLs from the outer leaflet of the OM and hand them over to MlaC, which resides in the periplasm. Consistent with this idea, *E. coli* MlaC has been crystallized with a bound PL and shown to interact with a complex of OmpF–MlaA <em>in vitro</em> (14). However, it is not clear how transfer of PLs from OmpC–MlaA to MlaC takes place at the OM. Because PL movement from the outer to inner leaflets of the OM is entropically disfavored, it is likely that translocation of PLs by the OmpC–MlaA complex would be coupled to transfer to MlaC; i.e., extracted PLs do not go into the inner leaflet. This model circumvents the problem of having to work against a PL concentration gradient between the two leaflets and allows PL transfer to MlaC to occur in an energy-independent, possibly affinity-driven, fashion. If this were true, it may be possible that MlaC also influences the function of the OmpC–MlaA complex. Specifically, binding of MlaC to the complex may be required for PL extraction from the outer leaflet of the OM. MlaC could alter the structure and/or dynamics of MlaA in the OmpC–MlaA complex, ultimately leading to efficient PL translocation across the OM.

Lipid asymmetry is critical for the OM to function as an effective permeability barrier. Thus, understanding mechanistic aspects of how bacterial cells maintain OM lipid asymmetry would guide us in designing strategies to overcome the barrier. Our work on elucidating the architecture and function of the OmpC–MlaA complex has revealed critical insights into the role of a hairpin loop on MlaA in modulating activity, a feature that can be exploited in drug discovery efforts. In particular, aspects of how bacterial cells maintain OM lipid asymmetry.
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before pelleting and resuspended in 1 ml of ice-cold TBS (20 mm Tris, pH 8.0, 150 mm NaCl). The samples were either directly or irradiated with UV light at 365 nm for 20 min at 4 °C or room temperature. All samples were pelleted again and finally resuspended in 200 μl of 2× Laemmli buffer, boiled for 10 min, and centrifuged at 21,000 x g in a microcentrifuge for 1 min at room temperature; 15 μl of each sample subjected to SDS–PAGE and immunoblot analyses.

Overexpression and purification of OmpC–MlaA–His complexes

All proteins were overexpressed in and purified from BL21(DE3) derivatives. We found that BL21(DE3) strains from multiple labs do not actually produce OmpC (the gene is somehow missing in these strains); therefore, to obtain OmpC–MlaA complexes, we deleted ompF from the chromosome and introduced ompC on a plasmid. OmpC–MlaA–His protein complexes were overexpressed and purified from BL21(DE3) cells with chromosomal ΔompF::kan background co-transformed with either pDSW206ompC_pBpa pSup–BpaRS–6TRN and pCDFdmlaA–His (for in vitro cross-linking experiments), or pACYC184ompC and pET22b(+)_dmlaA–His (for characterization of the WT complex). An overnight 10-ml culture was grown from a single colony in LB broth supplemented with appropriate antibiotics at 37 °C. The cell culture was then used to inoculate a 1-liter culture and grown at the same temperature before pelleting and resuspended in 1 ml of ice-cold TBS (20 mm Tris, pH 8.0, 150 mm NaCl). The samples were either directly or irradiated with UV light at 365 nm for 20 min at 4 °C or room temperature. All samples were pelleted again and finally resuspended in 200 μl of 2× Laemmli buffer, boiled for 10 min, and centrifuged at 21,000 x g in a microcentrifuge for 1 min at room temperature; 15 μl of each sample subjected to SDS–PAGE and immunoblot analyses.

Affinity-purification experiments

Affinity-purification experiments were conducted using ΔmlaA strains expressing MlaA–His at low levels from the pET23/42 vector. For each strain, a 1.5-liter culture (inoculated from an overnight culture at 1:100 dilution) was grown in LB broth at 37 °C until A600 of ~0.6. The cells were pelleted by centrifugation at 4,700 x g for 20 min and then resuspended in 10-ml TBS containing 1 mM PMSF (Calbiochem) and 30 mM imidazole (Sigma–Aldrich). The cells were lysed with three rounds of sonication on ice (38% power, 1-s pulse on, 1-s pulse off for 3 min). Cell lysates were incubated overnight with 1% n-dodecyl β-d-maltoside (DDM; Calbiochem) at 4 °C. Cell debris was removed by centrifugation at 24,000 x g for 30 min at 4 °C. Subsequently, supernatant was incubated with 1 ml of nickel–nitriotriacetic acid nickel resin (Qiagen), pre-equilibrated with 20 ml of wash buffer (TBS containing 0.025% DDM and 80 mM imidazole) in a column for 1 h at 4 °C with rocking. The mixture was allowed to drain by gravity before washing vigorously with 10 x 10 ml of wash buffer and eluted with 5 ml of elution buffer (TBS containing 0.025% DDM and 500 mM imidazole). The eluate was concentrated in an Amicon Ultra 100-kDa cutoff ultrafiltration device (Merck Millipore) by centrifugation at 4,000 × g to ~500 μl. The proteins were further purified by SEC system (AKTA Pure; GE Healthcare) at 4 °C on a prepacked Superdex 200 increase 10/300 GL column, using TBS containing 0.025% DDM as the eluent. Protein samples were used either directly or irradiated with UV at 365 nm for in vivo photoactivatable cross-linking experiments.

SEC–MALS analysis to determine absolute molar masses of OmpC–MlaA–His complex

Prior to each SEC–MALS analysis, a preparative SEC was performed for BSA (Sigma–Aldrich) to separate monodisperse monomeric peak and to use as a quality control for the MALS detectors. In each experiment, monomeric BSA was injected before the protein of interest, and the settings (calibration constant for TREF detector; Wyatt Technology) that gave the well-characterized molar mass of BSA (66.4 kDa) were used for the molar mass calculation of the protein of interest. SEC purified OmpC–MlaA–His was concentrated to 5 mg/ml and injected into Superdex 200 Increase 10/300 GL column pre-equilibrated with TBS and 0.025% DDM. Light scattering and refractive index (n) data were collected online using miniDAWN TREF (Wyatt Technology) and Optilab T-rEX (Wyatt Technology), respectively, and analyzed by ASTRA 6.1.5.22 software (Wyatt Technology). Protein-conjugate analysis available in ASTRA software was applied to calculate non-proteinaceous part of the complex. In this analysis, the refractive index increment dn/dc values (where c is sample concentration) of 0.143 ml/g and 0.185 ml/g were used for DDM and protein complex, respectively (34). For BSA, UV extinction coefficient of 0.66 ml/(mg·cm) was used. For the OmpC–MlaA–His complex, the UV extinction coefficient was calculated to be 1.66 ml/(mg·cm), based on its predicted stoichiometric ratio OmpC3MlaA.

Trypsin digestion for protein N-terminal sequencing and MS analyses

A 1 mg/ml solution of purified OmpC–MlaA–His (OmpC was either WT or substituted with pBpa at selected positions) complex was incubated with or without 50 μg/ml trypsin (Sigma–Aldrich) for 1 h at room temperature. pBpa-substituted samples were irradiated with UV at 365 nm before trypsin digestion. The samples were then analyzed by SDS–PAGE, followed by SEC. Peak fractions from SEC for each sample were pooled, concentrated using an Amicon Ultra 100-kDa cutoff
ultrafiltration device (Merck Millipore), and resuspended in 2× Laemmli sample buffer before analyses by SDS–PAGE and immunoblot using α-MlaA antibody. For N-terminal sequencing, the samples were transferred onto polyvinylidene fluoride membrane, followed by Coomassie Blue staining (1–2 s). The desired protein bands were carefully excised with a surgical scalpel. For tandem MS, protein bands were excised from a Coomassie Blue–stained Tricine gel. Samples prepared for N-terminal sequencing and tandem MS were kept in sterile 1.5-ml centrifuge tubes before submission for analyses at Tufts University Core Facility (Boston, MA) and Taplin Biological Mass Spectrometry Facility, Harvard Medical School (Boston, MA), respectively.

**Substituted cysteine accessibility method**

1 ml of cells was grown to exponential phase (A\textsubscript{600} = ~0.6), washed twice with TBS (pH 8.0), and resuspended in 480 μl of TBS. For the blocking step, four tubes containing 120 μl of cell suspension were either untreated (positive and negative control tubes added with deionized H\textsubscript{2}O) or treated with 5 mM thiol-reactive reagent N-ethylmaleimide (NEM; Thermo Scientific) or MTSES (Biotium). Because MTSES is membrane-impermeable, it is expected to react with the free cysteine in MlaA variants only when the residue is near or at the membrane–water boundaries or in a hydrophilic channel. In contrast, NEM is expected to label all MlaA cysteine variants because it is membrane-permeable. Reaction with MTSES or NEM blocks the particular cysteine site from subsequently labeling by maleimide-PEG (Mal-PEG; 5 kDa, Sigma–Aldrich). After agitation at room temperature for 1 h, the cells were washed twice with TBS, pelleted at 16,000 × g, and resuspended in 100 μl of lysis buffer (10 mM urea, 1% SDS, 2 mM EDTA in 1 M Tris, pH 6.8). Both NEM- and MTSES-blocked samples and the positive control sample were exposed to 1.2 mM Mal-PEG-5k. After agitation for another hour with protection from light, all samples were added with 120 μl of 2× Laemmli buffer, boiled for 10 min, and centrifuged at 21,000 × g in a microcentrifuge for 1 min at room temperature; 20 μl from each sample tube were subjected to SDS–PAGE and immunoblot analyses.

**In vivo disulfide bond analysis**

Strain NR1216 (ΔdsbA) harboring pET23/42mla–His expressing MlaA–His with site-specific cysteine substitutions was grown overnight in LB broth at 37 °C. 0.5 ml of overnight culture was normalized by optical density, added with trichloroacetic acid at a final concentration of ~14%, and mixed thoroughly at 4 °C. This step was performed to prevent scrambling of disulfide bond formed in the cysteines substituted MlaA–His. Proteins precipitated for at least 30 min on ice were centrifuged at 16,000 × g for 10 min at 4 °C. The pellet was washed with 1 ml of ice-cold acetone and centrifuged again at 16,000 × g for 10 min at 4 °C. Supernatants were then aspirated, and the pellet was air dried at room temperature for at least 20 min. The samples were resuspended thoroughly with 100 μl of either 100 mM Tris-HCl, pH 8.0, 1% SDS (for nonreduced samples), or the same buffer supplemented with 100 mM of DTT (for reduced samples) and incubated for 20 min at room temperature. The samples were finally mixed with 4× Laemmli buffer, boiled for 10 min, and subjected to SDS–PAGE and immunoblotting analyses using α-MlaA antibody.

**Membrane extraction profile analysis**

Adapted from a membrane extraction protocol as described (35), 500 ml of cells were grown to exponential phase (A\textsubscript{600} = ~0.6). The cells were pelleted by centrifugation at 4700 × g for 20 min and then resuspended in 7.5 ml of TBS containing 1 mM PMSF (Calbiochem). The cells were lysed with three rounds of sonication on ice (38% power, 1-s pulse on, 1-s pulse off for 3 min). Unbroken cells were removed by centrifugation at 4,000 × g for 10 min. The cell lysate was collected and centrifuged at 24,000 × g for 1 h. The total membrane fraction (pellet) was resuspended in 4-ml TBS; 1-ml each of resuspended total membranes were pelleted at 21,100 × g for 1 h at 4 °C and thereafter resuspended in 1 ml of TBS, 0.1 M alkaline Na\textsubscript{2}CO\textsubscript{3} (pH 11), 4 M urea, or 1% (v/v) Triton X-100, respectively. After incubation for 1 h at room temperature with occasional gentle agitation, the samples were centrifuged at 21,100 × g for 1 h at room temperature to collect supernatant (extracted) and pellet (membrane) fractions. The pellet fractions were each resuspended in 1 ml of TBS. All pellet and supernatant fractions were then resuspended thoroughly with 1 ml of 2× Laemmli buffer, boiled for 10 min, and centrifuged at 21,100 × g in a microcentrifuge for 1 min at room temperature; 20 μl from each sample tube was subjected to SDS–PAGE and immunoblot analyses.

**Docking of MlaA to OmpC**

The ClusPro server (22) was used to dock MlaA (ligand, Uniprot ID P76506, GREMLIN model) (17) to OmpC (receptor, Protein Data Bank code 2JIN) (18). The default server settings were used in the docking procedure. The minimum distance between six residues on OmpC and the corresponding cross-linked peptide regions of MlaA was calculated for all the predicted structures obtained from the server. Two OmpC–MlaA model with the smallest average minimum distance of all residue and peptide pairs were selected as the initial structures for use in the all-atom simulations.

**Simulation procedures and setup**

All simulations were performed using version 5.1.4 of the GROMACS simulation package (36, 37).

**All-atom simulations**

In total, seven all-atom simulations were performed (Table S4). The simulations were performed using the CHARMM36 force filed parameter set (38). The equations of motion were integrated using the Verlet leapfrog algorithm with a step size of 2 fs. Lengths of hydrogen bonds were constrained with the LINCS algorithm (39). Electrostatic interactions were treated using the smooth particle mesh Ewald method (40), with cutoff for short-range interactions of 1.2 nm. The van der Waals interactions were switched smoothly to zero between 1.0 and 1.2 nm. The neighbor list was updated every 20 steps. The Nose-Hoover thermostat (41, 42) with a coupling constant of 1.0 ps was used to maintain a constant system temperature of 313 K. The protein, membrane, and solvent (water and ions) were coupled to separate thermostats. The Parrinello–Rahman barostat
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(43) with a coupling constant of 5.0 ps was used to maintain a pressure of 1 bar. Semi-isotropic pressure coupling was used for all the membrane systems, whereas isotropic coupling was used for the solvent-only system. Initial velocities were set according to the Maxwell distribution.

Proteins were inserted into a pre-equilibrated, symmetrical 1,2-dimyristoyl-phosphatidylethanolamine membrane over 5 ns using the membed tool (44) in the GROMACS simulation package. Subsequent equilibration, with position restraints of 1000 kJ mol$^{-1}$ placed on all nonhydrogen protein atoms, was performed for 20 ns to allow the solvent and lipids to equilibrate around the proteins. The position restraints were removed before performing the production runs.

All of the simulations performed are summarized in Table S4. For the OmpC trimer docked with MlaA in orientation 1, three separate production simulations with different initial velocities were performed for the OmpC–MlaA complex, resulting in three trajectories of 500, 320, and 130 ns in length, respectively. For the OmpC trimer docked with MlaA in orientation 2, two separate production simulations with different initial velocities were performed for the complex, resulting in two trajectories of 500 ns in length. Clustering was performed on the MlaA structures obtained from a combined trajectory of all three (MlaA orientation 1) or two (MlaA orientation 2) atomistic simulations: a total of 4750 frames spaced every 0.2 ns. The structures were assigned to clusters using RMSD with a 0.1-nm cutoff. Four and two clusters, respectively, were observed for MlaA in the two orientations to contain greater than 100 frames. The central structure of these four clusters was used to generate the representative OmpC–MlaA models (Fig. S8). tri_cavity was used to identify the location the pore cavity (45), and Hole was used to create the pore profile (46).

Temperature titration for chromosomal ompC mutants

Purified WT and mutant OmpC–MlaA–His complexes were aliquoted into 1.5-mL centrifuge tubes and incubated in water bath set at different temperatures for 10 min. 20 μL of each sample were transferred into separate tubes and mixed immediately with equal volume of 2× Laemmli buffer and subjected to SDS–PAGE in 12% Tris HCl gels, followed by Coomassie Blue staining (Sigma–Aldrich).

LPS labeling and lipid A isolation

Mild acid hydrolysis of $^{32}$P-labeled cultures was used to isolate lipid A according to a procedure described previously (4, 12, 47) with some modifications. 5-mL cultures were grown (inoculated with overnight cultures at 1:100 dilution) in LB broth at 37 °C until $A_{600}$ reached between 0.4–0.6. The cells were normalized by optical density, first diluted to $A_{600} = 0.1$ ($\sim 10^8$ cells) and then serially diluted (10-fold) in LB broth using 96-well microtiter plates. 2.5 μL of the diluted cultures were manually spotted onto the plates, dried, and incubated overnight at 37 °C. Plate images were visualized by G:Box Chemi-XT4 (Genesys version 1.4.3.0, Syngene).

OM permeability studies

OM sensitivity against SDS/EDTA was judged by CFU analyses on LB agar plates containing indicated concentrations of SDS/EDTA. Briefly, 5–mL cultures were grown (inoculated with overnight cultures at 1:100 dilution) in LB broth at 37 °C until $A_{600}$ reached between 0.4–0.6. The cells were normalized by optical density, first diluted to $A_{600} = 0.1$ ($\sim 10^8$ cells) and then serially diluted (10-fold) in LB broth using 96-well microtiter plates. 2.5 μL of the diluted cultures were manually spotted onto the plates, dried, and incubated overnight at 37 °C. Plate images were visualized by G:Box Chemi-XT4 (Genesys version 1.4.3.0, Syngene).

SDS–PAGE, immunoblotting, and staining

All samples subjected to SDS–PAGE were mixed 1:1 with 2× Laemmli buffer. Except for temperature titration experiments, the samples were subsequently either kept at room temperature or subjected to boiling at 100 °C for 10 min. Equal volumes of the samples were loaded onto the gels. As indicated in the figure legends, SDS–PAGE was performed using either 12% or 15% Tris–HCl gels (48) or 15% Tricine gel (49) at 200 V for 50 min. After SDS–PAGE, the gels were either visualized by Coomassie Blue staining or subjected to immunoblot analysis. Immunoblot analysis was performed by transferring protein bands from the gels onto polyvinylidene fluoride membranes (Immun-Blot 0.2 μm; Bio-Rad) using semidyed electroblotting system (Trans-
ECL™ anti-mouse IgG–HRP and anti-rabbit IgG–HRP were diluted for 1–3 h at room temperature. Secondary antibody conjugated to the horseradish peroxidase (HRP) (Qiagen) at 1:5000 dilution for 1–3 h at room temperature. Secondary antibody ECL™ anti-mouse IgG–HRP and anti-rabbit IgG–HRP were used at 1:5000 dilutions. Luminata Forte Western HRP Substrate (Merck Millipore) was used to develop the membranes, and chemiluminescence signals were visualized by G:Box Chemi-XT4 (Genesys version 1.4.3.0, Syngene).

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