Periplasm homeostatic regulation maintains spatial constraints essential for cell envelope processes and cell viability.

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The cell envelope of Gram-negative bacteria consists of two membranes surrounding a periplasm and peptidoglycan layer. Molecular machines spanning the cell envelope depend on spatial constraints and load-bearing forces across the cell envelope and surface. The mechanisms dictating spatial constraints across the cell envelope remain incompletely defined. In *Escherichia coli*, the coiled-coil lipoprotein Lpp contributes the only covalent linkage between the outer membrane and the underlying peptidoglycan layer. Using proteomics, molecular dynamics and a synthetic lethal screen we show that lengthening Lpp to the upper limit does not change the spatial constraint, but rather impacts the load-bearing capacity across the outer membrane. Our findings demonstrate *E. coli* expressing elongated Lpp homeostatically counteracts periplasmic enlargement with a combination of tilting Lpp and reducing Lpp abundance. By genetic screening we identified all of the genes in *E. coli* that become essential in order to enact this homeostasis, and by quantitative proteomics discovered that very few proteins need to be up- or down-regulated in steady-state levels in order to enact this homeostasis. We observed increased levels of factors determining cell stiffness, decrease membrane integrity, increase membrane vesiculation and a dependance on otherwise non-essential tethers to maintain lipid transport and peptidoglycan biosynthesis. Further this has implications for understanding how spatial constraint across the envelope controls processes such as flagellum-driven motility, cellular signaling and protein translocation.
INTRODUCTION

Gram-negative bacteria have a cell envelope composed of two membranes sandwiching between them an aqueous space called the periplasm, in which an essential structural layer of peptidoglycan (PG) resides. The outer membrane is critical to cell growth and these bacteria face challenges to their cell biology in terms of membrane protein assembly and lipid-transport pathways that must traverse the distance from the inner membrane (IM) to the outer membrane (OM) 1. Recent work investigating the spatial demands for assembly of proteins into the outer membrane has shown precincts of active protein integration into the membrane can deliver new material to the growing outer membrane 2 and that random planar movement from these precincts drives the observed non-uniform distributions of the major proteins of the outer membrane 3, 4. By contrast to these protein components that diffuse to the outer membrane, elements of the LPS-transport machinery (eg. the Lpt complex) 5, 6, 7, 8, 9 span the OM and IM in order to fulfil their function in delivering lipid components to the outer membrane.

The PG layer is a fundamental aspect of the cell envelope, and it must be dynamically remodeled to allow growth as well as the assembly and transit of trans-envelope structures. PG synthesis and remodeling is a complex process with high levels of redundancy at various steps, involving at least 50 enzymes in E. coli 10. The Penicillin-binding proteins (PBPs) are the core components responsible for the periplasmic biosynthesis of peptidoglycan. There are multiple PBP complexes including the two core, semi-redundant PBP complexes PBP1a and PBP1b embedded in the IM which are activated by interactions with lipoproteins LpoA and LpoB embedded in the OM. Thus the activation of PG synthesis by these enzymes is spatially regulated, serving as a self-limiting molecular ruler to modulate PG thickness 11. Cells must possess either a functional PBP1a or PBP1b system for growth 12, 13.

Trans-envelope complexes such as lipopolysaccharide (LPS) transit pathways 5, 6, 7, 8, 9 and the protein translocation and assembly module (the TAM) 14, 15, 16 are also spatially constrained by the need to reach across from the IM to OM in order to function. The PG layer is covalently attached to the outer membrane by Braun’s lipoprotein (Lpp), with two recent papers addressing whether extending the length of Lpp would impact on (i) flagellar function given that the flagellum spans both membranes 17, and (ii) signal-transduction systems that span the OM to IM 18. In both cases, extending the coiled-coil structure of Lpp by 21 residues (Lpp+21) was found to be the longest form that supported close to normal growth 17, 18. Imaging of these Lpp+21 strains in both studies showed that the total periplasmic width had been stretched ~3-4 nm 17, 18. This being the case, the Lpp+21 model would provide a powerful experimental system to study how processes like OM biogenesis and PG biosynthesis can be maintained under a spatial stress on the cell wall.
To understand how trans-envelope processes in *E. coli* adapt to the presence of an enlarged periplasm, a combination of phenotypic analysis, proteomics, molecular dynamics and a synthetic lethal screen was employed to identify and characterize factors needed to maintain viability in the Lpp\(^{+21}\) strain of *E. coli*. The genetic screen demanded synthetic growth phenotypes from an array of mutants each lacking a gene that, while non-essential in wild-type *E. coli*\(^{19}\), is essential in the Lpp\(^{+21}\) strain. These genes fall into three functional categories: PG biosynthesis and remodelling, LPS biosynthesis and PG-outer membrane linkage. We show that previously non-essential proteins involved in bridging the gap between the OM and PG become essential in the context of the Lpp\(^{+21}\) strain background. These include previously known PG binding OM proteins (OmpA and Pal) as well as proteins previously not known to play an active role in linking the OM and PG (TolC and YiaD). We observed a thicker more diffuse or heterogeneous PG layer in the Lpp\(^{+21}\) strain and whole cell proteomics revealed that in response to an increased length of Lpp, *E. coli* increases the levels of a range of cell envelope proteins involved in PG turnover. We discuss the outcomes in terms of how the PG-outer membrane linkage functionalizes the periplasm, the evolutionary constrains in place to maintain this functionality, and the specific activity of Lpp in contributing to the load-bearing function of the OM.
RESULTS

Resilience and growth of the Lpp^{+21} strain

A phylogenetic assessment of Lpp lengths across diverse bacterial lineages showed a very narrow window of protein size (Fig. 1A), with Lpp being 78 residues in most species of bacteria including E. coli. Lpp lengths of 99 residues or more are at the upper end of the natural range for this protein and, in nature, these longer Lpp proteins are found in the genus Geobacter. The previously described Lpp^{+21} isoform expressed in E. coli therefore sits near the upper physiological range observed among bacteria. We introduced a gene encoding the Lpp^{+21} isoform into an E. coli background suitable for genetic screens (Fig. S1) and confirmed the size of the protein by SDS-PAGE of bacterial cell extracts (Fig. 1B). Enteric bacteria like E. coli can thrive in hyperosmotic environments compared to most laboratory growth media, such as the human gut by maintaining the periplasm and the cytoplasm in an iso-osmotic state \(^{20, 21}\). This is achieved by adjusting the solute concentration in the cell compartments by influx or efflux of water. By doing so, osmolality contributes immensely to the architectural aspects of cellular compartments \(^{21, 22}\). In this study sorbitol was used to mimic these physiological osmotic conditions. Previous studies have demonstrated that the periplasmic volume increased rapidly in response to increased osmolyte levels in the external medium. This phenomenon can reduce the cytoplasmic volume by about 30%, thereby constricting the IM inwards, and substantially increasing the periplasmic volume by around 300% \(^{21, 22, 23}\). The Lpp^{+21} isoform had little impact on growth of E. coli. On minimal growth medium, growth rates of the Lpp^{+21} strain of E. coli were equivalent to the isogenic wild-type E. coli (Fig. 1C). This was likewise true on growth media osmotically balanced with concentrations of sorbitol up to 1.0M (Fig. 1D), and on rich (LB) medium with or without sorbitol (Fig. 1D).

To establish the extent to which the periplasm had been remodeled in the Lpp^{+21} strain of E. coli, we compared the periplasms of WT and Lpp^{+21} strains using electron cryotomography in which cells are preserved in a frozen-hydrated, near-native state. To discern peptidoglycan, which was indistinct in previous studies, we increased the signal to noise ratio in images by calculating subtomogram averages instead of inspecting individual tomograms. As expected, the average distance from the middle of the IM density to the middle of the OM density was increased in both strains under hyperosmotic conditions when compared to previously reported data where cells were grown in standard laboratory media \(^{18}\). As shown in Fig. 1E, the averaged distance from the IM to the OM were somewhat greater in the Lpp^{+21} strain (32-36nm) compared to the isogenic wild-type strain (30-32nm) in line with previous Lpp+21 periplasmic width measurements. Previous studies could not discern peptidoglycan; whereas here, we were able discern peptidoglycan in subtomogram averages. The distance to the center of the PG density from the OM was slightly increased in the Lpp^{+21} strain (Fig. 1F), although by less than the anticipated ~3 nm. The PG morphology was also qualitatively different: in the wild-type strain a uniform dark PG layer could be observed in the images, whereas in the Lpp^{+21} strain...
a broader and more diffuse PG layer was evident, suggesting heterogeneity in both the density and thickness of the PG.
Figure 1. Phenotypes of *E. coli* cells encoding the Lpp+21 isoform. (A) Non-redundant Lpp sequences were identified (Methods) and the protein length charted on the x-axis. The number of non-redundant sequences showing that length is shown on the y-axis. The location of Lpp and the lengthened Lpp+21 are indicated. (B) Whole cell lysates were prepared from the indicated strains and subject to SDS-PAGE and immunoblot analysis with anti-Lpp antibodies and anti-OmpA antibodies. OmpA serves as a loading control. (C) The JW5028 – Keio BW25113 strain with *kan* gene replacing a pseudogene background and isogenic Lpp+21 strain (Fig. S1) were grown over 24 hours. The growth medium is M9, containing the indicated concentration of sorbitol as an osmolyte. (D) Growth rates for the same strains were measured in rich (LB) growth media with and without sorbitol over 20 hours. (E) The periplasmic width distribution of the indicated strains in hyperosmotic conditions. While PG layer in the wild-type strain is a uniform thin electron dense layer, the PG layer in the Lpp+21 strain is more diffuse and thicker. (F) Subtomogram averages of cell envelopes in hyperosmotic conditions. Measurements from EM views evaluate the distance between OM and PG in the Lpp+21 strain micrographs. The histogram depicts the frequency with which a given distance is observed between the OM and PG.
Cell envelope response to elongated Lpp

To determine the adaptive response to changing the distance constraint between OM and PG, quantitative whole cell proteomics was applied to evaluate the Lpp\textsuperscript{+21} strain. Triplicate samples of the wild-type and Lpp\textsuperscript{+21} strains were processed for analysis by mass spectrometry and we sought to identify those proteins where the steady-state level increases or decreases three-fold or more (Log\textsubscript{2} fold change of ± 1.6) in the Lpp\textsuperscript{+21} strain (Fig. 2A; Table S1). The level of the oligopeptide transporter subunits (OppB, OppC, OppD and OppF) are substantially increased in the Lpp\textsuperscript{+21} strain compared to the wild-type (Table S2). This suggests an increase in PG turnover and an overall increased capacity to recycle PG components, and is consistent with the concomitant increase in AmiC, one of the two major amidases involved in PG remodeling. In addition, proteins implicated in diverse stress-responses (cold shock proteins CspG, CspA, CspI and YdfK, as well as envelope stress protein ZraP and redox stress protein YfcG) were observed at increased steady-state levels in Lpp\textsuperscript{+21} strain (Fig. 2A, Table S2). The greatest decreases were seen in the steady state levels of the GatZABCD proteins involved in galactitol phosphotransferase system and DHAP synthesis (Fig. 2A). The gat\textsubscript{ABCD} genes have been shown to be responsive to factors that change \textit{E. coli} cell surface tension \cite{24} and Lpp\textsuperscript{+21} has been reported to significantly decrease cell stiffness \cite{25}.

A decrease was seen in the steady-state level of the Lpp\textsuperscript{+21} isoform in the mutant, to approximately one-eighth the level of Lpp in the wild-type strain (Fig. S2). This is consistent with the relative abundance of Lpp and Lpp\textsuperscript{+21} observed in SDS-PAGE analysis of cell extracts from the two strains (Fig. 1F). However, despite the relative decrease, Lpp\textsuperscript{+21} remains as a highly abundant component of the OM-PG linkage factors given that Lpp is present at up to 10\textsuperscript{6} protein molecules per wild-type cell \cite{1, 26}.

The mass spectrometry data was processed to allow for an analysis of sub-cellular proteomes \cite{27} (Fig S3). An initially puzzling observation was that the Lpp\textsuperscript{+21} strain has a 12\% overall reduction of total periplasmic protein compared to wild type (Fig. 2B). This was calculated as the proportion of the summed intensity from identified proteins predicted to reside in the periplasm in the STEPdb: G, E, F2, F3, I annotations \cite{27} (Fig S3). Both Lpp\textsuperscript{+21} strains and null Lpp strains of \textit{E. coli} are softer as previously adjudged by atomic force microscopy \cite{25}, and factors that increase the softness of \textit{E. coli} also increase outer membrane vesicle (OMV) production \cite{17, 28, 29, 30}. To address whether the measured depletion of periplasmic content reflects an increased production of OMVs, extracts measuring the amount of total protein in the OMV fraction were normalized to OD\textsubscript{600} (Fig 2C, Fig. S3). This confirmed that the presence of Lpp\textsuperscript{+21} promotes approximately 10-fold more total protein associated with the OMV fraction, reflecting increased OMV production. The overall level of OM proteins associated with the cells was maintained constant (Fig. S3) but a small OM integrity defect was evident from an increased sensitivity to SDS (Fig. 2D).
Figure 2. The Lpp+21 cells have softened outer membranes and increased blebbing. (A) Heat map of the significant proteomic differences observed between the wild-type and Lpp+21 mutant. Blue boxes indicate a relative reduction and red indicates a relative increase in protein level, centred on the average of the replicate samples. The grouping of the proteins is based on the similarity of the change in expression observed. (B) The Lpp+21 mutant has an overall reduction in the level of periplasmic proteins. (C) The Lpp+21 mutant has an increase in protein secreted via OMV blebbing. (D) SDS sensitivity profiles of the Lpp+21 mutant compared to the wild-type in increasing concentration of SDS in LB (solid) media. Representative data are shown from experiments performed in triplicate.
**Lpp\(^{+21}\) can be accommodated in the periplasm, but other factors become essential**

To directly address the altered phenotype induced by Lpp\(^{+21}\), we established a synthetic genetic array for factors in *E. coli* that become essential in order to maintain viability of the Lpp\(^{+21}\) strain (Fig S4). The screen demands synthetic growth phenotypes from an array of mutants each lacking a single gene that, while non-essential in wild-type *E. coli*, become essential in the Lpp\(^{+21}\) strain of *E. coli*. The endogenous lpp gene was replaced by a gene encoding Lpp\(^{+21}\) in a isogenic library of 3818 *E. coli* mutants, each of which lacks a non-essential gene. Growth on rich medium allowed the rescue of the new library for array into a format suitable for high-throughput screening with a Singer RoToR robotics platform (Fig S4). Phenotypic analysis was thereafter scored for growth by comparing the growth of the isogenic mutants in the Lpp background (Fig. 3A) with the equivalent mutants in the Lpp\(^{+21}\) background (Fig. 3B). Each of the genes that displayed a noticeable phenotype in these analyses are presented in Table 1.

The only cytoplasmic factor identified in our screen, YraN is predicted to be a Holliday-junction resolvase related protein, and we therefore speculate that this mutant failed to resolve the merodiploid condition transient in the introduction of the lpp\(^{+21}\) condition to the background strain, making the yraN mutant a technique-relevant artefact of the screen. This being the case, only functions performed in the periplasm were recovered as essential to viability for the Lpp\(^{+21}\) strain.

Most of the components of the LPS biosynthetic machinery are essential genes in *E. coli* and are thus not represented in the library of non-essential genes. Several non-essential genes in the LPS biosynthetic pathway that are in the library, become essential to the Lpp\(^{+21}\) strain (Table 1): the core LPS biosynthesis factors GalU, GmhB and RfaD were shown to be essential in the Lpp\(^{+21}\) strain.

**An essential role for keeping the OM-PG distance**

Several genes encoding proteins that could play roles in anchoring the PG within the cell envelope were identified as essential in the Lpp\(^{+21}\) background. Independently, none of the major proteins bridging the OM and PG are essential for growth in *E. coli* \(^{19}\) and all are therefore represented in the library. In a Lpp\(^{+21}\) background the genes encoding the β-barrel protein OmpA and the lipoprotein Pal become essential (Table 1). PG-binding domain PF00691 is common to these proteins: appended to a beta-barrel in OmpA, but to a lipooyl anchor in Pal, and is also conserved in other proteins across diverse Gram-negative bacteria (Fig 3C). In *E. coli* there are 4 additional proteins containing this PG-binding domain and these were mapped in a sequence similarity network analysis (Fig 3D). A protein of unknown function, YiaD, is present (Fig 3D) and it too is essential in a Lpp\(^{+21}\) background (Table 1). We suggest, therefore, that this protein plays a substantive role in OM-PG linkage. The remaining three proteins: MotB, LafU and YfiB, are more divergent to the OmpA/Pal/YiaD cluster. Neither motB, lafU nor yfiB displayed a synthetic phenotype with Lpp\(^{+21}\), and it has
been suggested previously that motB, lafU and yfiB are not expressed at detectable levels under laboratory conditions.

Detecting genes encoding drug-efflux pumps as important for growth of the Lpp^{+21} strain was initially surprising. Either the absence of the inner membrane proteins AcrB or the OM component TolC caused a reduction of growth in the Lpp^{+21} genetic background (Table 1). When antibiotic selection was removed by plating the mutants on medium without chloramphenicol, the synthetic growth defects were observed in the absence of drug selection (Fig. 3D), indicating that this synthetic phenotype is not the result of a decreased drug efflux activity. The trans-envelope AcrAB-TolC multidrug efflux pump has been shown to traverse through the PG and interact directly with PG at several defined sites, and as loss of the core AcrB and TolC components became essential, we suggest that this system could be acting as an additional OM-PG linkage that becomes essential in a Lpp^{+21} background. Together with the observation that OmpA, Pal, YiaD and TolC are also essential in the Lpp^{+21} genetic background, these data suggest that functions that maintain local areas of closer contact between the OM and PG are essential for viability.
Figure 3. Factors that become essential to mediate OM-PG linkage in Lpp+21 E. coli. (A) The growth phenotype in M9 minimal media (0.5 M sorbitol) of single gene knock outs that exhibit essentiality on Lpp+21 background (Table 1). (B) The growth phenotype in M9 minimal media (0.5 M sorbitol) of double gene mutants. The mutants are results of Hfr Cavalli lpp:lpp+21 cat crossed with 22 KanR recipients shown in panel A (methods). The double mutants are indicated and are arranged in 4 biological replicates (each having 4 technical replicates). (C) Sequence similarity network of domain (Pfam PF00691) containing proteins from across the Proteobacteria. Each circle represents a protein from a representative proteome (rp35) containing the PF00691 domain, connected by lines with a length imparted by their similarity score as defined by EFI - Enzyme Similarity Tool 68, with a cutoff of 30. Proteins are colored by their taxonomic class and the approximate location of the E. coli K12 six PF00691 proteins is indicated. (D) Synthetic lethal phenotype of the drug efflux mutants in the absence of selective antibiotics in M9 minimal media condition. Representative data are shown from experiments performed in biological triplicate.
| Cellular process                                      | \textbf{Lpp}^{+21}-essential genes | Function                                                                 |
|------------------------------------------------------|------------------------------------|--------------------------------------------------------------------------|
| LPS biosynthesis                                     | \textit{galU}                      | UDP-glucose metabolic process                                            |
|                                                      | \textit{gmhB/yaeD}                 | ADP-L-glycero-\(\beta\)-D-manno-heptose biosynthetic process            |
|                                                      | \textit{rfaD/hldD/waaD}            | ADP-L-glycero-\(\beta\)-D-manno-heptose biosynthetic process            |
| Peptidoglycan biosynthesis, turnover and remodelling | \textit{lpoA, lpoB, mrcA, mrcB}    | Regulators of PG synthases                                              |
|                                                      | \textit{ldcA}                      | L,D-carboxypeptidase involved in PG recycling/turnover                  |
|                                                      | \textit{nlpD}                      | Regulator of AmiC PG hydrolase                                          |
|                                                      | \textit{slt}                       | Lytic PG transglycosylase, degradation of uncrosslinked glycan strands  |
|                                                      | \textit{ybhC}                      | Carbohydrate esterase, PG turnover\(^*\)                               |
| PG-OM linkage                                        | \textit{acrB, mdtL\(^*\), mdtK\(^*\), ydhl\(^*\)} | IM components of drug efflux pumps.                                     |
|                                                      | \textit{tolC}                      | OM component of drug efflux pump                                       |
|                                                      | \textit{ompA}                      | \(\beta\)-barrel protein with PG binding domain                        |
|                                                      | \textit{pal}                       | Lipoprotein with PG binding domain                                     |
|                                                      | \textit{yiaD}                      | Lipoprotein with PG binding domain                                     |

\(^*\)The function of YbhC is not known, but cellular localization, structural and transcriptomic evidence indicates it may be involved in PG turnover\(^{37}\)

\(^*\)Putative IM drug efflux machinery, TolC independent with as yet unknown OM component\(^{38}\)
To compare the behaviors of Lpp and Lpp\(^{+21}\), both with and without an OmpA monomer present, molecular
dynamics (MD) simulations were run for 200 ns each (see Methods for details). The Lpp lipoprotein from \textit{E. coli} is a triple coiled-coil that is anchored to the inside face of the outer membrane by its N-terminal acyl
group with a length equating to approximately 7.5 nm\(^{39}\). The experiments were established to test the scenario
for Lpp trimers or Lpp\(^{+21}\) trimers in the absence or presence of an OmpA tether between the patch of OM and
patch of PG (Fig. 4). These MD simulations only represent a portion of the membrane, and as such we could
not account for potential heterogeneity in the protein and LPS concentrations between a WT and Lpp\(^{+21}\) cell.

For three of the systems: Lpp only, Lpp with OmpA, and Lpp\(^{+21}\) only, tilting was marginal, with tilt angles of
76.9±4.7°, 75.5±4.7°, and 82.8±2.9°, respectively (all numbers from the last 100 ns of the 200-ns trajectory).
These angles are in agreement with previous simulations of Lpp alone (~80°) and Lpp with an OmpA
monomer (~75°) PG\(^\text{36,40,41}\) In initial simulations of Lpp\(^{+21}\) with OmpA, the non-covalent connection between
OmpA and PG was quickly disrupted as Lpp\(^{+21}\) extended from its kinked state. Therefore, the simulation was
repeated with an enforced OmpA-PG connection. Lpp\(^{+21}\) was observed to both straighten and tilt within the
first 100 ns; the tilt angle measured for the last 100 ns was 49.4±2.3°.

The distance between the OM inner leaflet phosphorus atoms and the PG sugars was measured in each
scenario. In the presence of Lpp, the distances with and without OmpA were similar at 8.3±2.1 nm and 8.1±1.2
nm, respectively. This was not true for the other scenarios where the distance for Lpp\(^{+21}\) alone was 11.6±1.3
nm, but for Lpp\(^{+21}\) with OmpA, the distance was reduced significantly to 8.7±2.7 nm. Thus, we observe that
PG-binding proteins like OmpA can counteract the increased distance imposed by Lpp\(^{+21}\), inducing it to tilt
significantly in accommodation. We also compared our simulations to the distances that were measured by
EM (centre of the OM to centre of the PG). In wild-type \textit{E. coli} (i.e. Lpp+OmpA), the centre-centre distance
in the simulations is 10.7 +/- 2.2 nm (Fig 4B), similar to the 9.7 -10.8 nm measured in intact cells (Fig. 1F).
The centre-centre distance measured in the simulation of Lpp\(^{+21}\) tilted by the presence of OmpA (11.0 +/- 2.6
nm), fits the observed distances of 10.8 – 12.8 nm much better than the distance that would be created by a
perpendicular Lpp\(^{+21}\) (14.2 +/- 1.3 nm).
Figure 4. Final states of the OM-PG linkage from MD simulations. (A-B). A patch of OM with the LPS molecules depicted in orange (lipid A moiety) and yellow (core oligosaccharides), and the phospholipids in the inner leaflet of the OM depicted in grey. The PG layer (blue for glycans and green for peptide crosslinks) is attached to the OM via two trimers of Lpp (A), or a trimer of Lpp and the PG-binding domain of OmpA (red). The β-barrel anchor of OmpA is shown embedded in the OM. (C-D) Equivalent scenarios formed with Lpp+21 trimers. The distances shown are calculated from the inner face of the OM to the centre of the PG layer and represent the average over the last 100 ns of a 200-ns simulation.
DISCUSSION

We observed that over a range of osmotic conditions, and in nutrient-rich or nutrient-poor media, growth rates of the Lpp\(^{+21}\) strain of *E. coli* were equivalent to the isogenic wild-type *E. coli*, suggesting bacteria can adapt to the presence of the extended Lpp\(^{+21}\). We observed a moderate increase to the OM-IM space similar to that previously reported\(^{17,18,25}\) and reviewed\(^{25,42}\). Three adaptive features were expressed as phenotypes in the Lpp\(^{+21}\) strain: (i) the steady-state level of the Lpp\(^{+21}\) tether was reduced eight-fold compared to the level of Lpp in the isogenic wild-type strain, and other tethers that enforce a wild-type distance: OmpA, Pal, YiaD and TolC, became essential factors in the Lpp\(^{+21}\) condition, (ii) structures that depend on a wild-type OM-IM distance, such as the LPS transport system, continued to function but key components of the system became essential for cell viability, and (iii) the PG network took on characteristics of dysregulated synthesis and all components of an otherwise redundant PG biosynthesis pathway become essential to viability. In response to Lpp elongation we demonstrated a reduction in Lpp levels, as shown by the quantitative proteomic presented here, and the apparent tilting of the elongated Lpp as suggested by our molecular dynamic simulation. The reduction in the copy number of the elongated Lpp isoform observed in the quantitative proteomics was not due to differences in transcriptional regulation, as RNAseq data demonstrated a moderate increase in the level of *lpp*\(^{+21}\) transcripts in the mutant (Table S6). Taken together, this may suggest a reinterpretation of some of the previous conclusions drawn from experiments with elongated Lpp isoforms that assumed that Lpp\(^{+21}\) was present at the same amount as Lpp, and that Lpp\(^{+21}\) sips perpendicular to the membranes and thereby the 21 extra amino acids always stretch the periplasmic width by a constant amount\(^{17,18,25}\).

Tethers that enforce the distance constraint between the OM and PG layer became essential factors in the Lpp\(^{+21}\) condition

The lipoyl-N-terminus of Lpp is integrated into the OM, and the C-terminus of a protomer of the Lpp trimer is covalently linked to the PG layer. Stereochemically, only one unit of a Lpp trimer can be covalently attached to the PG layer, and this stoichiometry has been observed experimentally\(^{39,40,43}\). This makes each Lpp trimer an important bridge between the OM and PG layers of the cell wall, but it has not been clear whether the core role of Lpp is acting in compression or tension. In other words\(^{44}\), is the role of Lpp to be a supportive brace to keep the OM away from the PG, or a binding anchor to bring the OM as closely as possible to the PG? In a wild-type *E. coli* scenario, molecular dynamics simulations show that the flexible linkers in PG-binding proteins like OmpA allow for adjustments in positioning the PG relative to the OM. In the presence of Lpp, the flexible linker of the PG binding domain of OmpA needs to extend further from the OM, as the distance between the OM and PG increases when Lpp sits perpendicular to the OM\(^{40}\). In the absence of Lpp, OmpA determines the distance between the OM and PG\(^{40,41}\), but is not essential for cell viability\(^{45,46}\). Taken together with the data presented here from characterization of the Lpp\(^{+21}\) phenotypes, we suggest that Lpp functions as a brace to keep the OM away from the PG layer.
The genetic screen showed that each one of the OM-located PG-binding factors Pal, YiaD, and OmpA are essential (as is TolC) for viability of the Lpp<sup>+</sup> strain of *E. coli*. While previous high throughput synthetic genetic screens demonstrate none of Pal, YiaD, OmpA nor TolC are essential or negatively affect the growth in an Lpp null background<sup>45,46</sup>. The overlap in the distance constraints measured for cells expressing Lpp and Lpp<sup>+</sup> suggests that much of the Lpp<sup>+</sup> is in the highly tilted (49.4±2.3°) form observed in the MD simulations. The enforced tilt and reduction in steady state levels of Lpp<sup>+</sup> would severely impact its bracing force, with the measurable consequence that membrane integrity (SDS resistance) is diminished, OmpA and other tethers become essential to viability, and the OM is highly permissive to OMV formation. Furthermore, the discovery of YiaD in the genetic screen for essential factors is significant. Our proteomic assessment and previously published data<sup>26</sup> demonstrated that the copy number of YiaD is much lower than that of either Pal or OmpA, making it not immediately clear how a minor component of the membrane could be exerting such an essential role. Previously, YiaD was suggested to be a factor regulating OMP biosynthesis by the BAM complex<sup>47</sup>. Structural analysis shows YiaD to be highly similar to the PG-binding domains of Pal and OmpA<sup>48</sup>. We suggest that the primary function of YiaD is to mediate OM-PG linkage, and that this indirectly impacts on OMP biogenesis.

**Continued function of LPS transport in localized regions of the periplasm**

The bridges needed to mediate LPS transport appear to be susceptible to disruption caused by Lpp<sup>+</sup> in *E. coli*. LptA is subject to degradation if the Lpt complex is compromised<sup>7</sup>, with LptA-LptC and LptA-LptD interactions proposed as key quality control steps in the assembly of the Lpt complex<sup>6</sup>. That the Lpp<sup>+</sup> cells have enough maintained sufficient LPS in the OM to maintain membrane integrity is supported by our observations of the only minor increase in SDS sensitivity of the strain and observations by others of only minor changes in vancomycin sensitivity<sup>25</sup>. These findings are consistent with observations through electron microscopy that regions of the periplasm in Lpp<sup>+</sup> cells are maintained with OM-PG distances reflective of the wild-type condition, which would permit LPS transport to the OM.

**All components of the PG biosynthesis pathways become essential to cell viability**

In *E. coli*, the weave of the PG-layer is maintained in a uniform, open state through quality control mechanisms that depend on the regulators in the OM (LpoA and LpoB) being able to permeate it to contact the synthetases (PBP1a and PBP1b) in the IM. It has been hypothesized in this way that the OM lipoproteins may serve as a molecular ruler to modulate PG thickness, maintaining a single layer of PG equidistant from the OM layer under normal conditions<sup>11</sup>. The PG synthases encoded by *mrcA* and *mrcB* are redundant in the sense that *mrcA* mutant strains and *mrcB* mutant strains are each viable<sup>49,50,51</sup>. However, in the Lpp<sup>+</sup> strain both *mrcA* and *mrcB* (as well as their OM lipoprotein partners) are essential for viability, indicative of a compromised capability to build the PG-layer. Mutations designed to impact on these interactions lead to transient deposition.
of “high-density PG” and “multi-layered PG” through dysregulation of the synthetases. The broader and more diffuse morphology of the PG-layer observed by electron microscopy in the Lpp+21 strain could suggest a similar phenotype associated with transient or local impacts on the OM-PG distance. It’s worth noting Δlpp ΔmrcB mutants have previously shown a moderate growth defect and that this defect could be alleviated with the introduction of the lpp+21 gene, whereas in the present study we observed a synthetic lethal phenotype in the ΔmrcB lpp+21 mutants. This discrepancy could be due differences in growth conditions used (LB vs hyperosmotic minimal media) imparting additional periplasmic stress or differences in the strain background (DH300 vs BW25113).

An essential requirement was also placed on PG-layer remodeling, whereby the PG-binding factor NlpD was found to be essential in Lpp+21 cells: its function is in modulating the activity of the amidase AmiC to remodel PG strands, and AmiC was observed at increased steady-state levels in the Lpp+21 strain. Taken together with the increase in oligopeptide transporter subunits (OppB, OppC, OppD and OppF) in the Lpp+21 strain to recycle PG precursors across the IM, our results suggest a clearance of the malformed PG caused by dysregulation of the PG synthases is a crucial adaptation in the Lpp+21 strain.

**Stiffness, load-bearing and connection of OM-PG**

The concept of bacterial cell stiffness has emerged as a means to understand the physical parameters that define how readily bacteria can respond to major environmental changes. Measurements by AFM have revealed a characteristic stiffness in Gram-negative bacterial cells that is contributed by load-bearing outer membrane and its attachment to the underlying PG layer. In *E. coli*, mutants lacking Lpp, Pal or OmpA are softer than wild-type cells, and cells expressing the Lpp+21 isoform are also softer than wild-type cells. Taken together with our observation that the Lpp+21 isoform display a broader and more diffuse PG layer morphology in the subtomogram averages, this further supports the proposition that the OM is a major contributor to cell stiffness.

Although Lpp length alone does not appear to dictate periplasm width, indeed the *Pseudomonas aeruginosa* Lpp protein is only 4 amino acids longer than that of *E. coli* but its periplasm has been measured as ~3 nm larger under identical conditions, the length of Lpp seen in *E. coli* is conserved in a range of bacterial lineages. The species from the genus *Geobacter* naturally express longer Lpp proteins of 99 or more residues, equivalent to Lpp+21 in length. *Geobacter* species have a complicated periplasm housing “electron conduits” formed of transmembrane and periplasmic redox proteins, in order to transfer electrons to the external surface of the bacterial cell. There have been no direct measurements of the width the periplasm of *Geobacter*, but it would be interesting to see if these naturally longer Lpp forms correlates with an enlarged periplasm. That species of *Geobacter* serve as an exception to what is otherwise a strict rule about the length of Lpp, and thus the structurally-enforced distance constraint between the OM and PG layer, raises interesting questions.
about whether OM softness, OMV production or increased OM permeability might assist the unusual biological functions of the OM and periplasm in bacteria other than *E. coli*.
MATERIALS AND METHODS

Bacterial strains and growth conditions

E. coli BW25113 was the parental strain of all the recipient strains used in this study. JW5028, a derivative of BW25113, containing a kanamycin resistance marker in place of a pseudogene, was used as the wild-type for this study. Strains were grown in LB broth or M9 minimal media broth supplemented with 1 mM MgSO₄, 0.1 mM CaCl₂, 1.12 mM thiamine and 0.2% (w/v) glucose for a defined minimal media. To osmotically stabilize the growth medium, sorbitol was supplemented to a final concentration of 0.5 M. For overnight cultures, strains were grown in LB broth overnight at 37 °C under continuous agitation. Subculturing was done by diluting the saturated culture 1:100 using new media. The cells were then grown to mid-exponential growth phase (OD₆₀₀ = 0.5-0.6) at 37 °C under shaking. Culture media was supplemented with antibiotics for plasmid selection and maintenance or selection of mutants at the following concentrations: 100 μg/ml ampicillin, 30 μg/ml kanamycin, 34 μg/ml chloramphenicol. 15 g/l agar was added to media before autoclaving when solid media was required (Fig. S4).

Construction of Lpp⁺²¹ mutant

The endogenous lpp gene was replaced with the extended lpp⁺²¹ gene previously described¹⁷,¹⁸ with minor modifications (Fig. S1). First the replacement was done in the donor E. coli Hfr Cavalli cells using the λ–red recombinant system. A gene block was sourced (Integrated DNA Technologies) containing extra 21 amino acid residues (three heptad repeats) inserted between codon 42 and 43 of E. coli Lpp. The gene block also contained 50 bp DNA flanking 5’ and 3’ ends of the lpp⁺²¹ gene. On the 5’ end, the extension was homologous to DNA sequence upstream of lpp, while on the 3’ end, the extension was homologous to the cat gene. The gene block was combined with the cat gene by Gibson assembly, and the resulting PCR fragment was used to replace lpp. The new Lpp⁺²¹ strain was selected by plating on medium containing chloramphenicol. Chromosomal lpp⁺²¹ was moved into E. coli BW25113 background by mating with kanamycin-resistant Keio collection strain JW5028, described above, to generate a double mutant¹⁹,⁶² (Fig. S4). The mutation was verified by PCR described (Fig. S1) and sequencing.

SDS-PAGE and immunoblotting

Cells grown in M9 minimal media (0.5 M sorbitol) and normalized by OD₆₀₀nm, lysed in Laemmli SDS-loading dye and separated in 15% acrylamide gels and transferred to 0.45 μm hydrophobic Immobilon-P PVDF membrane (Merck Millipore). Immunoblotting was as described previously¹⁹. Rabbit primary antibodies; α-Lpp antibody (kindly provided by T. Silhavy) and α-OmpA were diluted 1:400,000 and 1: 30,000, respectively in 5% skim milk, TBST. The membranes were incubated with goat, α-rabbit IgG, HRP-conjugated secondary antibody (Sigma; 1: 20,000 in 5% skim milk, TBST), and washed with TBST. Detection was by enhanced
chemiluminescence with ECL prime western blotting detection reagent (GE Healthcare Life Sciences), visualized using Super RX-N film (Fujifilm).

**Outer membrane vesicle purification and quantification**

Overnight cultured cells, grown in LB without antibiotics, were washed twice in 1 x M9 salts then subcultured in 500 ml M9 minimal media supplemented with 0.5 M sorbitol (1:1000 dilution). The strains were grown to late logarithmic phase without antibiotics, \( \text{OD}_{600} \approx 0.9 \) and spun down to collect culture supernatant. Collected culture supernatant were then processed for OMVs isolation and purification using differential ultracentrifugation technique as discussed previously. OMVs were washed twice in PBS to remove sorbitol then quantified using a bicinchoninic acid assay kit (Thermo Scientific CST#23225).

**SDS sensitivity**

Streaks were made on LB (solid) media with 0% SDS – 5% SDS. After 16 hrs of incubation at 37 °C, the minimum SDS concentration inhibiting growth was obtained by analyzing growth on each concentration. The streaks were done in duplicate and repeated three times.

**Proteomics**

Saturated overnight cultures were washed twice in 1 x M9 salts and diluted 1:100 in 10 ml M9 minimal media supplemented with 0.5 M sorbitol. Cultures were further grown to logarithmic phase, collected by centrifugation and washed using PBS buffer. The cell pellet was homogenised in 4 % SDS, 100 mM Tris, pH 8.1 and boiled at 95 °C for 10 minutes. The lysate was then sonicated with a Bioruptor® Pico (Diagenode) and protein concentration was determined with Bicinchoninic Acid assay (BCA, Thermo Fisher). SDS was removed with chloroform/methanol, the protein was digested by trypsin overnight and the digested peptides were purified with ZipTips (Agilent). Using a Dionex UltiMate 3000 RSLCnano system equipped with a Dionex UltiMate 3000 RS autosampler, an Acclaim PepMap RSLC analytical column (75 µm x 50 cm, nanoViper, C18, 2 µm, 100Å; Thermo Scientific) and an Acclaim PepMap 100 trap column (100 µm x 2 cm, nanoViper, C18, 5 µm, 100Å; Thermo Scientific), the tryptic peptides were separated by increasing concentrations of 80% ACN / 0.1% formic acid at a flow of 250nl/min for 120 min and analyzed with a QExactive Plus mass spectrometer (Thermo Scientific). The instrument was operated in the data dependent acquisition mode to automatically switch between full scan MS and MS/MS acquisition. Each survey full scan (m/z 375–1575) was acquired in the Orbitrap with 60,000 resolution (at m/z 200) after accumulation of ions to a 3 x 10^6 target value with maximum injection time of 54 ms. Dynamic exclusion was set to 30 seconds. The 20 most intense multiply charged ions \((z \geq 2)\) were sequentially isolated and fragmented in the collision cell by higher-energy collisional dissociation (HCD) with a fixed injection time of 54 ms, 15,000 resolution and automatic gain control (AGC) target of 2 x 10^5.
The raw data files were analyzed using MaxQuant software suite v1.6.5.0 against Andromeda search engine for protein identification and to obtain their respective label-free quantification (LFQ) values using in-house standard parameters. The proteomics data was analyzed using LFQ-Analyst and the analysis of the data quality analysis is presented in Fig S6. Due to the 21 amino acid insertion in the Lpp²⁺ isoform, the relative levels of Lpp in the mutant had to be assessed manually. Only the unique peptide (IDQLSSDVQTLNAK) shared between the two isoforms was used to quantify the levels of Lpp and Lpp²⁺ in the wild-type and mutant strain, respectively (Fig S2).

To estimate the total relative amount of proteins from the various subcellular compartments, the raw intensities from peptides identified from proteins from different subcellular locations were summed and divided by the total summed intensity from all peptides. Sub-cellular locations annotations were applied from the STEPdb 2.0 where proteins designated F1, A, R, and N were classified as cytoplasmic; B was designated inner membrane; H, X, and F4 were designated outer membrane / extracellular; and I, G F2, F3, and E were designated as periplasmic.

Sequence similarity network analysis
Proteobacterial proteins containing the Pfam domain (PF00691) were extracted from the Representative Proteome 35% co-membership rpg-35 group and a sequence similarity network was generated with the EFI Enzyme Similarity Tool. This network was visualized with Cytoscape with a similarity score cutoff of 30. Each protein is represented by a colored circle node and each similarity match above the similarity score cutoff is represented by an edge between nodes with the length determined by the similarity score.

Lpp length distribution across bacterial species
To determine the amino acid length distribution of Lpp in Gammaproteobacteria (Table S4), amino acid sequences were sourced from the InterPRO database (version 81.0) using the Interpro Family tag - Murein-lipoprotein (IPR016367). Filtered Lpp sequences were then concatenated into representative nodes (at least >90% sequence similarity) using the online available amino acid Initiative-Enzyme Similarity Tool (EFI-EST).

Synthetic genetic interaction array
The Lpp²⁺¹ isoinform was transferred to each of the Keio collection clones by conjugation as described (Fig. S4). First, the Hfr chloramphenicol resistant Lpp²⁺¹ strain was arrayed in 384-colony density on LB agar containing chloramphenicol using the Singer rotor HAD (Singer Instruments, United Kingdom). Similarly, the Keio collection arrayed in 384-colony density was pinned on LB agar plates containing kanamycin and incubated overnight at 37 °C. Using the Singer rotor HDA, the Hfr Lpp²⁺¹ strain and the Keio collection clones from the 384-colony density were then co-pinned onto LB agar plates and incubated at 37 °C for 16 hours.
Following conjugation, the colonies were transferred to LB agar with kanamycin (selection 1) at the same colony density and incubated at 37 °C for 16 hours. To select for double mutants (selection 2), colonies from the intermediate selection were pinned on LB agar with both kanamycin and chloramphenicol and incubated at 37 °C for 14 hours. For assessment of synthetic genetic interaction in nutrient-limited media, the double mutants generated were replica pinned in M9 minimal media at the same density and incubated at 37 °C for 25-30 hours. Images were acquired using Phenobooth (Singer Instruments, United Kingdom) for analysis. Images were manually screened to cross-reference recipient plate images to the final double antibiotic selection plates images. Candidate synthetic lethal or growth-compromised mutants were then subjected to another round of screening in the same conditions as previously identified (mini-screen) for validation. Four biological replicates were included that were further arrayed in four technical replicates. Mutants were confirmed by PCR. Where further phenotypic screening of mutants was conducted, independent isogenic knock-out mutants were generated using the PCR and λ-red based homologous recombination method.

Since the Keio collection viaD mutant has been identified as containing a potential duplication event, the candidate viaD synthetic lethal interaction was confirmed through independently constructing a viaD mutant in the BW25113 strain background (Fig. S5). The lpp^+21 variant was subsequently generated in this mutant as described above. As with the Keio viaD mutant, this strain demonstrated synthetic lethality on M9 media.

Two colony PCR reactions (Fig. S4) confirmed the identity of all candidate double mutants, using a set of primers flanking the lpp gene, and a set of primers flanking the kanamycin gene (Table S5).

**Preparation of Electron Cryo-microscopy samples, data collection and analysis**

Strains were grown aerobically in M9 minimal media (0.5 M sorbitol) until an OD_{600} of 0.6 was reached. Cells were collected by spinning at 6000xg for 5 minutes and resuspended to an OD_{600} of ≈ 12. Cryo-EM, data collection and analysis were performed similarly to previous studies, except using 3-D subtomogram averages derived from whole-cell cryotomograms instead of projection images so as to discern peptidoglycan. Tilt series of WT and Lpp^+21 strains was acquired on an FEI Krios operating at 300 keV with a Gatan K2 direct detector and energy filter with a 20eV slit with a tilt range of +/- 60° using 3° increments and reconstructed using IMOD. Subtomograms were picked manually using 3DMOD along the length of all non-polar periplasm and averaged using PEET.

**Generation of simulation systems**

Initially, two systems were generated: the OM and PG, as previously detailed, with two copies of wild-type Lpp and with two copies of Lpp^+21. For wild-type Lpp, we used the homo trimer from PDB 1EQ7. For Lpp^+21, a monomer was first built using I-TASSER. Next, the trimer of Lpp^+21 was built using the wild-type Lpp trimer as a template, further optimized using Targeted Molecular Dynamics (TMD) for 1 ns. For both
Lpp and Lpp\textsuperscript{21}, the proteins were anchored in the OM via N-terminal acylation while the C-terminus of one copy from each trimer was covalently linked to the PG. The systems generated were prepared for equilibration using the following steps for 1 ns each: 1) minimization for 10,000 steps, 2) melting of lipid tails, 3) restraining only the PG and the protein, and 4) restraining the PG and the protein backbone. Both systems were equilibrated for 200 ns.

For each of the two systems (Lpp and Lpp\textsuperscript{21}), a new system was constructed with one Lpp trimer removed and OmpA inserted into the OM. The full-length OmpA structure was taken from Ortiz-Suarez et al.\textsuperscript{73}. The periplasmic domain region of the OmpA (clamp, hereafter) was lowered to the PG by shortening the distance between the PG and the clamp for 20 ns. After the clamp was lowered, it was clenched to the nearest DAP residue by using two different distance collective variables\textsuperscript{74} between the centre of mass of residue 242 or 256 from OmpA and that of a nearby PG DAP residue to maintain the connection for 110 ns. In the case of Lpp\textsuperscript{21}, PG was first pulled towards the OM to match the distance between them in wild-type Lpp, after which the clamp of OmpA was lowered and clenched for 110 ns. The wild-type Lpp/OmpA system was equilibrated for 200 ns; OmpA stayed bound to the PG without colvars. The Lpp\textsuperscript{21}/OmpA system was also equilibrated for 200 ns, but colvars were needed to maintain the OmpA-PG interaction.

**Molecular Dynamics (MD)**

All-atom molecular dynamics simulations were performed using NAMD \textsuperscript{2.11} and the CHARMM36m \textsuperscript{76} and CHARMM36 \textsuperscript{77} force-field parameters for proteins and lipids, respectively, with the TIP3P-CHARMM water model \textsuperscript{78}. Unless otherwise stated, all MD simulations were performed under a periodic boundary condition with a cut-off at 12 Å for short-range electrostatic and Lennard-Jones interactions with a force-based switching function starting at 10 Å. For long range electrostatic calculations, the particle-mesh Ewald method \textsuperscript{79} with a grid spacing of at most 1 Å was used for long-range electrostatic calculations. Bonds between a heavy atom and a hydrogen atom were maintained to be rigid, while all other bonds remain flexible. Unless otherwise stated, each system was equilibrated under an isothermal-isobaric ensemble (NPT) at 310 K and 1 bar, with a timestep of 4 fs after hydrogen mass repartitioning \textsuperscript{80}. A Langevin thermostat with a damping coefficient of 1 ps\textsuperscript{-1} was used for temperature control and a Langevin piston was used for pressure control. VMD was used for all visualization and analysis \textsuperscript{81}. 
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FIGURE LEGENDS

Figure 1. Phenotypes of E. coli cells encoding the Lpp\textsuperscript{+21} isoform. (A) Non-redundant Lpp sequences were identified (Methods) and the protein length charted on the x-axis. The number of non-redundant sequences showing that length is shown on the y-axis. The location of Lpp and the lengthened Lpp\textsuperscript{+21} are indicated. (B) Whole cell lysates were prepared from the indicated strains and subject to SDS-PAGE and immunoblot analysis with anti-Lpp antibodies and anti-OmpA antibodies. OmpA serves as a loading control. (C) The JW5028 – Keio BW25113 strain with kan gene replacing a pseudogene background and isogenic Lpp\textsuperscript{+21} strain (Fig. S1) were grown over 24 hours. The growth medium is M9, containing the indicated concentration of sorbitol as an osmolyte. (D) Growth rates for the same strains were measured in rich (LB) growth media with and without sorbitol over 20 hours. (E) The periplasmic width distribution of the indicated strains in hyperosmotic conditions. While PG layer in the wild-type strain is a uniform thin electron dense layer, the PG layer in the Lpp\textsuperscript{+21} strain is more diffuse and thicker. (F) Subtomogram averages of cell envelopes in hyperosmotic conditions. Measurements from EM views evaluate the distance between OM and PG in the Lpp\textsuperscript{+21} strain micrographs. The histogram depicts the frequency with which a given distance is observed between the OM and PG.

Figure 2. The Lpp\textsuperscript{+21} cells have softened outer membranes and increased blebbing. (A) Heat map of the significant proteomic differences observed between the wild-type and Lpp\textsuperscript{+21} mutant. Blue boxes indicate a relative reduction and red indicates a relative increase in protein level, centred on the average of the replicate samples. The grouping of the proteins is based on the similarity of the change in expression observed. (B) The Lpp\textsuperscript{+21} mutant has an overall reduction in the level of periplasmic proteins. (C) The Lpp\textsuperscript{+21} mutant has an increase in protein secreted via OMV blebbing. (D) SDS sensitivity profiles of the Lpp\textsuperscript{+21} mutant compared to the wild-type in increasing concentration of SDS in LB (solid) media. Representative data are shown from experiments performed in triplicate.

Figure 3. Factors that become essential to mediate OM-PG linkage in Lpp\textsuperscript{+21} E. coli. (A) The growth phenotype in M9 minimal media (0.5 M sorbitol) of single gene knock outs that exhibit essentiality on Lpp\textsuperscript{+21} background (Table 1). (B) The growth phenotype in M9 minimal media (0.5 M sorbitol) of double gene mutants. The mutants are results of Hfr Cavalli lpp: lpp\textsuperscript{+21} cat crossed with 22 Kan\textsuperscript{R} recipients shown in panel A (methods). The double mutants are indicated and are arranged in 4 biological replicates (each having 4 technical replicates). (C) Sequence similarity network of domain (Pfam PF00691) containing proteins from across the Proteobacteria. Each circle represents a protein from a representative proteome (rp35) containing the PF00691 domain, connected by lines with a length imparted by their similarity score as defined by EFI - Enzyme Similarity Tool \textsuperscript{68}, with a cutoff of 30. Proteins are colored by their taxonomic class and the approximate location of the E. coli K12 six PF00691 proteins is indicated. (D) Synthetic lethal phenotype of
the drug efflux mutants in the absence of selective antibiotics in M9 minimal media condition. Representative data are shown from experiments performed in biological triplicate.

Figure 4. **Final states of the OM-PG linkage from MD simulations.** (A-B). A patch of OM with the LPS molecules depicted in orange (lipid A moiety) and yellow (core oligosaccharides), and the phospholipids in the inner leaflet of the OM depicted in grey. The PG layer (blue for glycans and green for peptide crosslinks) is attached to the OM via two trimers of Lpp (A), or a trimer of Lpp and the PG-binding domain of OmpA (red). The β-barrel anchor of OmpA is shown embedded in the OM. (C-D) Equivalent scenarios formed with Lpp^{21} trimers. The distances shown are calculated from the inner face of the OM to the centre of the PG layer and represent the average over the last 100 ns of a 200-ns simulation.
**SUPPLEMENT**

**Figure S1. Construction and assessment of Lpp^{+21} E. coli.** (A) Schematic representation of the elongated Lpp^{+21} isoform. The strain was constructed by inserting (red lettering) heptad repeats between residue D42 and Q43 of Lpp. (B) Schematic detailing the isogenic replacement of the lpp gene with the lpp^{+21} gene. (C) PCR confirmation of the lpp^{+21} mutant strain (Methods).

**Figure S2. Quantitation of Lpp and Lpp^{+21} isoforms.** The sequence of the 21 residues inserted to create the Lpp^{+21} isoform is also indicated. Mass spectrometry data for Lpp vs Lpp^{+21} was reanalyzed after extraction from the whole cell proteomic data. Given the different tryptic peptides generated from the two isoforms of Lpp, a shared peptide (red) was used to quantify the relative levels of each Lpp isoform in each of the strains. The graphs document the relative levels of the peptide and show that the presence or absence of sorbitol in the growth medium has no effect on the level of Lpp^{+21} relative to Lpp.

**Figure S3. Sub-cellular proteomics of Lpp^{+21} E. coli.** (A) Relative amount of protein in different subcellular compartments, measured by the raw relative proportion of peptide intensities identified from proteins annotated to reside in cellular compartments defined in the STEPdb. (B) Comparative proteome profile of the wild-type and Lpp^{+21} strain for cell lysate and extracted outer membrane vesicles. Loading of each technical replicate was normalized to OD 600. Representative data are shown from experiments performed in biological triplicate.

**Figure S4. A synthetic lethal screen to determine genes essential to Lpp^{+21} E. coli.** An Hfr donor strain carrying a selectable marker (cat) fused to lpp^{+21}, replacing the lpp ORF, is mated on agar plates with arrayed F^- recipients (384) per plate carrying a selectable marker (kan) replacing other ORF. Upon mating, cells are subjected the first round of selection (intermediate selection) using antibiotic kanamycin and then further subjected to a second round of selection using both antibiotics; (A) depicts images of representative plates generated in each step of the procedure with imaging and manual analysis step, cross-referencing of single gene knock outs and double recombinants, included. (B) depiction of the strains as cartoons generated in each step of the procedure. (C) A representative mini screen of manually selected genes from the main synthetic lethal screen. Sterility controls were included on each mini screen. The mini screen was performed in 384-pin density with each clone arrayed in four biological replicates, each having four technical replicates (blue boxes). Synthetic lethal mutants identified from the mini screen were further verified by PCR to confirm the presence of both gene modifications and rule out partial duplication events.

**Figure S5. Construction and characterization of the validation yiaD mutant.** A kanamycin resistance cassette was amplified from pKD4 using primers with overhangs complementary to upstream and downstream
of \textit{yiaD}. The PCR fragment was electroporated in BW25113 cells harbouring the \(\lambda\)–red recombinase plasmid (pKD46). Transformants were selected on kanamycin-resistant plates and verified by PCR (methods). Primers flanking the \textit{yiaD} gene confirm replacement of \textit{yiaD} with kanamycin cassette and primers amplifying \textit{lpp} confirm \textit{lpp}^{	ext{+21}} replacement of \textit{lpp}. The sequence information for all primers used are included in Table S5.

**Figure S6. Proteomics quality control report.** (A) Principle Component Analysis (PCA) plot of data from triplicate samples of wild-type and \textit{lpp}^{	ext{+21}} strains grown on M9 medium with or without supplementation by sorbitol. (B) Sample Coefficient of variation plots for the same four samples.

**Figure S7. Molecular dynamics plots of \textit{lpp} tilt angles over time.** Plots of the angles over time, from which the averages and standard deviations were derived. Simulation were run for 200 ns and average tilt angles from the last 100 ns were calculated. To check for convergence, the \textit{lpp} simulation ran for an additional 200 ns. The angle from the last 200 ns is 78.3 +/- 4.8°, nearly identical (within 1.4°) to that from the 100-200 ns window.

**Table S1 Proteomic results**

**Table S2. Substantive changes in steady-state protein levels in cell envelope of \textit{lpp}^{	ext{+21}}**

**Table S3: Proteins used in the generation of the sequence similarity network**

**Table S4: Representative \textit{lpp} protein information**

**Table S5: Bacterial strains and primers used in the study**

**Table S6: Genes up-regulated in \textit{lpp}^{+21} strain**
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