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A β-barrel for oil transport through lipid membranes: Dynamic NMR structures of AlkL

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The protein AlkL is known to increase permeability of the outer membrane of bacteria for hydrophobic molecules, yet the mechanism of transport has not been determined. Differing crystal and NMR structures of homologous proteins resulted in a controversy regarding the degree of structure and the role of long extracellular loops. Here we solve this controversy by determining the de novo NMR structure in near-native lipid bilayers, and by accessing structural dynamics relevant to hydrophobic substrate permeation through molecular-dynamics simulations and by characteristic NMR relaxation parameters. Dynamic lateral exit sites large enough to accommodate substrates such as carvone or octane occur through restructuring of a barrel extension formed by the extracellular loops.

As in any cell, the outer membrane of bacteria is composed of a lipid bilayer that separates intracellular processes from the outside environment. In addition to this lipid bilayer, Gram-negative bacteria have evolved an outer leaflet composed mainly of lipopolysaccharides (LPS) (1). While the inner membrane is permeable to hydrophobic molecules, the charged and densely packed LPS layer reduces their permeation rates 50–100-fold, as compared with phospholipid membranes (2–4) (Fig. 1). Since these membrane properties limit uptake of nutrients, bacteria make use of a wide array of passive transmembrane porins, β-barrels of sufficiently large diameter to allow nonspecific permeation of small molecules (1, 5).

The OmpW family of outer-membrane proteins (6), classified as porins, is widespread in Gram-negative bacteria. These proteins fold into eight-stranded β-barrels displaying a narrow pore diameter in the membrane, which appears too compact to permit passage of even small molecular substrates. Yet there is mounting evidence that the OmpW family protein AlkL functions as a passive importer of hydrophobic molecules. AlkL is found on the same operon together with genes necessary for oxidation of alkanes and utilization as a carbon and energy source (7) (Fig. 1). These include an alkane monooxygenase (AlkB) and associated rubredoxin (AlkG) and rubredoxin reductase (AlkT), as well as an aldehyde dehydrogenase (AlkH) and alcohol dehydrogenase (AlkK) and a positive regulatory protein (AlKS). Due to its location on the alk operon, AlkL was proposed to have a passive transport function, increasing the rate of diffusion for alkanes. Definitive evidence for such transport was recently found by incorporating the alkane hydroxylase complex of Pseudomonas putida GPO1 with and without AlkL in cells (8–10), and in even more defined nanoreactor polymersomes (11). These studies demonstrate transport of alkanes, and a cyclic methyl ketone, respectively, in line with the wide substrate utilization of P. putida. In polymersomes, the increase in transport-dependent production of product alcohol occurs with an increase in the number of incorporated AlkL molecules.

While the transport function of AlkL has been established, a controversy in the transport mechanism still exists. By analogy to other β-barrels (12–15), a lateral exit was proposed based on X-ray structures of OmpW (16) and OprG (17), formed from proline residues that break the hydrogen bonding at the end of the transmembrane β-strands. The X-ray structure reveals a β-barrel arrangement in the loops; however, the 1.3-Å radius of the putative lateral opening is incompatible with the size of known substrates of AlkL such as alkanes or monocyclic molecules. The static X-ray structures are therefore insufficient to explain the mode of action, and a view complemented by dynamics is needed (18–20). In contrast, micellar solution NMR structures lack the putative exit sites entirely, and rather display flexible extracellular loops (21, 22). This supports an alternative vision of transport characterized by hinge motions of the loops and major rearrangements of backbone segments or partial loss of secondary structure. Here we solve this controversy by determining the de novo NMR structure in near-native lipid bilayers for AlkL, and by detecting structural dynamics relevant to

Significance

Here we show how AlkL, a minimalistic outer-membrane protein from oil-consuming bacteria, exploits dynamics of extracellular loops to channel substrate across the polysaccharide barrier and into the hydrophobic interior of the outer membrane. This work represents a unique example of a side-by-side atomic-level structure determination by solution NMR in detergents and by solid-state NMR in lipid bilayers, which critically demonstrates the importance of a lipid environment to investigate function. Corroborating our experimental measurements, molecular-dynamics simulations capture substrate transit via lateral openings. The capacity to unravel membrane protein function under near-native conditions, fueled by the latest method developments, opens up a new frontier for their investigation, and provides thereby for improved fundamental insights into biological processes.
Furthermore, NMR experiments probing protein-to-lipid interactions demonstrate that AlkL is in immediate contact with lipids (23). Residues all across the β-barrel show contact with the lipid acyl chains, defining the transmembrane region of the protein (SI Appendix, Fig. S2). For sensitivity reasons, the lipid-to-protein ratio (LPR) was minimized to 0.5 by mass (~18 by mole). The resulting spectrum at LPR of 0.5 is unchanged from an LPR of 10, but spectral quality degrades with LPR of 0.25 (SI Appendix, Fig. S3). This combined evidence shows that AlkL is properly embedded in the membrane and that the protein structure is independent of concentration over the range detectable by MAS NMR.

NMR Structures. Two structures were determined from assigned resonances in 170 out of 203 residues in lipids, and 132 residues in detergent. Unassigned signals in lipids belong mainly to the flexible N-terminus, to periplasmic turns, and to short portions of the first extracellular loop, and are absent from many of the NMR spectra. In detergent, only the first loop was assigned, and displays chemical shifts characteristic of unstructured proteins. Approximately 50 additional unassigned signals in the 2D 1H-15N heteronuclear single quantum coherence fall in the same chemical shift range, indicating a lack of β-structure also for the remaining loops. This is in contrast to the DMPC preparation where assignments in all loops indicate structure. No nuclear Overhauser effect (NOE) transfer to these peaks was observed, also pointing to lack of structure. Yet additional evidence pointing toward random coiled loops in detergents was obtained by preparing a sample in D2O. In D2O only the assigned residues of the transmembrane region remained detectable, indicating protection from HD exchange. In an MSP1 nanodisc preparation (SI Appendix, Fig. S4) without detergent, mainly unassigned residues appeared, indicating that the rigid transmembrane domain is not detectable in the larger nanodisc particle. Consistent with the DMPC preparation where more loop residues are structured, some peaks that were flexible in detergent are not detected in the nanodisc preparation.

Structures were calculated from a clear pattern of proton-proton cross-strand contacts in radio-frequency-driven recoupling (RFDR) or NOE spectra. While traditional three-dimensional (3D) HHN- and HNN NOE spectroscopy (NOESY) spectra provided sufficient resolution in solution, we turned to four-dimensional (4D) HNNH-RFDR spectra in solids to remove ambiguities in the assignment of cross-strand restraints (SI Appendix, Figs. S5 and S6). The fully protonated sample together with >100-kHz MAS (24) granted access to sidechain resonance assignments and collection of sidechain-to-sidechain distance restraints which was used to refine the bilayer-bound structure. All restraints used for calculation of the two structures are listed in SI Appendix, Table S2.

The two AlkL structures determined by MAS and solution NMR are shown in Fig. 2 A and B, and share an identical transmembrane (TM) β-barrel motif. The lipid-embedded structure, however, reveals an elongated β-barrel fold, extending into the region where polysaccharide would be in LPS, similar to homologous crystal structures. Structured loop residues also occur in a sample reconstituted in the presence of LPS and DMPC lipids (SI Appendix, Fig. S7) as indicated by the detection of the same loop resonances in cross-polarization-based spectra. Only minor chemical shift perturbation (CSP) is observed. Conversely, the loops are highly flexible in OG detergent micelles, which is in turn consistent with previous determinations of homologs by solution NMR. That the bilayer-embedded structure in this case appears closer to homology models of crystal structures is in contrast to a recent solid-state NMR structure of OmpG (25) obtained in 2D crystalline lipid bilayers, which agreed better with the solution NMR conformation. In the case of AlkL, the perturbation due to

Results and Discussion
To understand the mechanism of the transport function of AlkL, we determined two separate structures, one in detergent micelles by using well-established solution NMR protocols and a second in lipid bilayers, by using the latest magic-angle spinning (MAS) NMR methodology. This is complemented by characterization of structural dynamics by both simulation and NMR relaxation measurements. The solution NMR data were acquired from octyl glucoside solution (OG), with traces of the refolding detergent LDAO using a perdeuterated AlkL sample. Both of these detergents are considered mild. Lipid-embedded AlkL data were acquired from 1,2-dimyristoyl-sn-glycero-3-phosphocholine (DMPC) bilayer samples, as well as from preparations reconstituted in the presence of LPS, using both fully protonated as well as perdeuterated AlkL. In these latter preparations, negative stain transmission electron micrographs reveal the presence of liposomes, without detectable protein aggregates (SI Appendix, Fig. S1). Furthermore, NMR experiments probing protein-to-lipid interactions

Fig. 1. The main features of the inner and outer membranes of Gram-negative bacteria, and location of key proteins of the Alk operon. Hydrophobic molecules such as alkanes and terpenoids (Inset, blue), do not readily traverse (blue dashed arrow) the LPS (purple). Instead, their permeation is assisted by porins such as AlkL (green). A permeation path through the extracellular loops with a lateral exit into the hydrophobic bilayer (blue arrow) was proposed based on crystal structures of AlkL homologs (8). Detergent micelle solution data support a more dynamic process without a specific lateral release (21, 22).
detergent micelles appears to outweigh the influence of crystal contacts.

Site-specific $^{15}$N longitudinal relaxation ($R_1$) (26) in the lipid-embedded sample explains the apparent inconsistency between the two models in lipids and in micelles, by assessing dynamic motions occurring on a fast (nanosecond) timescale (Fig. 2C and SI Appendix, Table S3). While the TM region displays uniform low relaxation rates, the extracellular portion is subject to enhanced $\eta$ backbone dynamics. These measurements define a clear junction in flexibility between the TM and extracellular regions of the structure. Considering that transport through the AlkL inner pore would require channel opening, the low $R_1$ relaxation rates of the TM region are in favor of a model where the TM region is serving solely as an anchor for the more flexible extracellular part. This complements the rigid representation of extracellular $\beta$-strands in Fig. 2A, and is consistent with the fact that a detergent environment (Fig. 2B) results in loss of structure in the loops. The same mobility that is likely needed to allow for the transport function of the protein appears to predispose the extracellular part to destabilization by even mild detergent, an effect which likely plagues other membrane proteins (27).

Since the lipid bilayer most closely mimics the native membrane, and these conditions support a folded extracellular domain, we focused on the membrane-embedded sample and solid-state NMR for further structural and dynamical analysis of transport in AlkL. The surface hydrophobicity of the bilayer NMR structure (Fig. 3A) demonstrates the expected hydrophobic exterior (tan) in the transmembrane region, with polar and charged residues lining the interior. Reverse polarity (purple) is seen on the extracellular barrel extension, which has a hydrophilic exterior.

These structural features define a long, accessible hydrophobic pore, leading up to the membrane interfacial region. Within the TM domain, the pore narrows to less than the radius of a water molecule as visualized using HOLE (28) (Fig. 3B), which along with a hydrophilic character suggests lateral release into the hydrophobic membrane rather than continued transport through the protein. A cross-sectional view (Fig. 3D) reveals the tight stacking of long internal sidechains that block diffusion of molecules through the TM region. At the same time, the structure features six small openings, two of which (I and IV) are located close to the membrane surface (Fig. 3C), in the vicinity of several proline residues that break the pattern of cross-strand hydrogen bonds. Site I is the most prominent in our NMR structure, while site IV corresponds to the previously proposed putative lateral exit (8, 16, 17). These potential exit sites would require dynamic rearrangement to accommodate the size of substrate. Detailed views of the hydrophobic residues lining the extracellular pore are displayed in Fig. 3D and E. The residue R168 is shown occluding the pore at the top of the TM barrel, and in the same view, the residues V75 and T54 are seen near the mouth of the largest potential exit site I/II.

**Dynamical Lateral Release Model.** The validity of this “dynamical lateral release” model is supported by MD simulations (Figs. 4 and 5) and NMR relaxation rates sensitive to slow motions (Fig. 6), recorded both in the presence and the absence of the substrates octane and carvone. *P. putida* utilizes a wide substrate range including linear alkanes, cyclic hydrocarbons, and even polycyclic molecules (PAHs) (7). The alkanes are metabolized by the alk proteins, whereas PAHs are utilized by additional metabolic pathways, making octane an ideal molecule to investigate the transport function of AlkL. All simulations were initiated in explicit water from the solid-state NMR structure of AlkL embedded in a membrane. Simulations in LPS membrane (details in the SI Appendix) were initiated with 1-palmitoyl-2-oleoyl-sn-glycero-3-phosphocholine on the inner leaflet (Fig. 4). These MD simulations conform to the expected behavior of the LPS, namely, that the hydrophobic octane does not traverse the barrier characterized by polar sugar moieties functionalized with charged phosphate groups. As expected, octane enters the hydrophobic pore formed by loop residues, and traverses this region of the protein up until the level of the lipid head groups. At this point, the pore is occluded by charged sidechains, in particular R168. Fig. 4A and B shows two snapshots from the MD...
trajectory, as well as a comparison of the bilayer penetration depth of octane in LPS alone and mediated by AlkL (Fig. 4A, Inset). This comparison was made with the zero depth referenced to the LPS phosphate groups. Consistent with the low rate (millisecond timescale) of transport that supports growth of yeast and E. coli on sugar (29–31), these 1 μs MD trajectories were not able to capture substrate permeation events, even when reinitialized from structures where octane was already bound in the pore. Nonetheless, the protein did readily allow deep penetration of substrate into the membrane, and we were able to observe full octane transit in MD trajectories in a simplified DMPC environment. Fig. 4C–E shows several snapshots from 1-μs trajectories in which octane escaped into the lipid membrane from three different interstrand locations, namely sites I and IV identified in Fig. 3C as well as between P147 and P171. While the LPS appears to stabilize the protein, increasing the energy barrier for such events, the events may nevertheless represent permeation pathways albeit that occur much more slowly in the native membrane.

In addition to octane, we observed permeation for the monocyclic ketone carvone, which is an intermediate-sized oil that provides a challenging test of the transport function of AlkL. Carvone has a similar structure to a cyclic methyl ketone used to demonstrate accelerated transport facilitated by AlkL (11). Fig. 5A shows the initial conformation of AlkL embedded into a DMPC membrane in explicit water. The simulations in DMPC match the conditions of the NMR structure determination, where the protein was properly folded. Over multiple microsecond trajectories, the structure is stable, and the barrel extension exhibits higher plasticity than the transmembrane region, in agreement with the experimental data. Within 100 ns, carvone molecules cluster onto the barrel extension and several molecules are seen to concurrently occupy the pore (Fig. 5B). After 1,000 ns, all carvone molecules have diffused into the membrane or remain within the barrel extension (Fig. 5C). Although this simulation time was again not sufficient to observe lateral release of carvone, release did occur in further simulations, initialized from structures where carvone was already bound in the pore (see SI Appendix, Figs. S8 and S9 for the detailed simulation protocols). As visualized by four snapshots in Fig. 5D, carvone is observed to exit toward the membrane uniquely through site I. This shows that exit of both octane, and the larger carvone molecule is possible without a pair of proline residues breaking cross-strand hydrogen bonds, and therefore represents a distinct class of lateral exit site. The simulations in DMPC do not preserve the barrier properties of LPS. They can therefore not be used to assess AlkL-mediated acceleration of permeation, but can be used to probe the permeation pathway. As expected, most carvone molecules enter the membrane directly in DMPC, and relatively few carvone molecules enter the membrane via AlkL. This nevertheless demonstrates the transport function of the protein. Since there is no active transport in AlkL, we expect nearly all octane or carvone to partition to the hydrophobic membrane, and this is indeed observed in long enough MD simulations. In a living system, consumption of the substrate occurs via AlkB in the inner membrane, necessitating exit from the inner leaflet of the outer membrane, and entry into the inner membrane lipids. To our knowledge, these phospholipid membranes do not present a substantial barrier to AlkB substrates, and no additional proteins have been identified in the passive flux of metabolites once through the LPS barrier.

![Fig. 3.](#) The bilayer structure of AlkL, and potential exit. A shows surface hydrophobicity (tan) and hydrophilicity (purple). In B, the central pore is visualized using the HOLE (18) program. Red indicates a diameter less than one water molecule (1.15 Å), while green and blue indicate a radius of one water (1.15–2.30 Å), and more than one water molecule (>2.30 Å), respectively. The positions of charged residues constraining the interior are labeled. In C, potential lateral exit sites are visualized. In D–F, slices through the β-barrel extension are characterized by a hydrophobic inner surface (D and E), with the central pore blocked by arginine in the membrane-embedded part of the protein F. In the same vicinity, a lateral opening occurs near residues T54 and V75.
Transit events through AlkL are made possible by dynamic remodeling of the pore, as evidenced by the distinct distributions of conformers in the absence and presence of carvone (Fig. 5E and F). Notably, the main conformers identified in simulations with bound carvone (Fig. 5E) mostly retained a widened potential exit site I, as compared to the NMR structure (Fig. 3C). Furthermore, there is hardly any overlap in the phase-space regions explored by the protein without and with bound carvone (Fig. 5F). Interestingly, the NMR structure is located in between these two regions. The motion along the first (largest) eigenvector in a principal component analysis (PCA) shows that the difference without and with bound carvone is related to the opening of the newly identified site I for carvone permeation (Movie S1).

We used two further NMR measurements to detect structural variations and motional processes relevant to hydrophobic substrate permeation: chemical shift perturbations (CSPs) and 15N longitudinal relaxation under RF spin lock (ρ1ρτ) (Fig. 6 and SI Appendix, Table S3). High CSPs highlight significant changes in the barrel extension for both carvone and octane. Of particular interest are large CSP values in residues flanking P76, which lies in a β-bulge separating sites I and IV. Coincident with large CSPs are enhanced values of 15N R1ρ, an indicator of transient chemical shift changes (exchange phenomena) and microsecond–millisecond motions. R1ρ rates were also sensitive to the presence of carvone in the barrel extension (SI Appendix, Fig. S10). Overall, the NMR data are direct signatures of the transient expansion of the pore captured by MD simulations, providing a connection between loop flexibility and permeation. Of the sites implicated by dynamics measurements, sites III–VI are flanked by β-strands with a well-defined hydrogen bonding pattern in lipids (SI Appendix, Fig. S11), indicating a more static structure. Only the site I/II is larger and adjacent to the highly flexible loop 1. It is no surprise that this location supports dynamic release of the larger carvone molecule.

Whether the dynamic lateral release mechanism described here is general for proteins of the OmpW family remains an open question. While for AlkL, there is no evidence for hydrophilic substrates, and indeed no CSP was observed in AlkL with leucine as substrate (SI Appendix, Fig. S12), amino acids have been proposed for homologous proteins. If an OmpW family protein does accommodate hydrophilic substrates, lateral transport may represent the first part of the mechanism, which would traverse the polysaccharide barrier of the outer membrane. Complete traversal would then require transmembrane passage through the aliphatic lipid tails, which presents a barrier to hydrophilic substrate. This might occur via oligomeric interfaces or partner proteins. For example, crystal packing interactions are found in OmpW crystals (Protein Data Bank [PDB] ID code 2F1V) in a region corresponding to the exit site in AlkL (SI Appendix, Fig. S11), and a pore large enough for glycerol is found in a homology model of OprG based on the OmpW crystal structure (32). Alternatively, OmpW family members might act in concert with other proteins present in the outer membrane in a similar way as a type 9 secretion system that exports protein through a lateral exit to a porin (33). Although no such protein has been identified, the outer membrane is rich with a plethora of membrane proteins that may play a role. Further evidence is needed to understand whether AlkL homologs exhibit the same transport function via a lateral release mechanism as proposed here for AlkL.

Conclusions. Through a combination of NMR and MD simulations, we uncovered a dynamic translocation pathway in an eight-stranded β-barrel for the import of hydrophobic molecules. The side-by-side comparison of sample conditions shows a strong impact of lipids and LPS in particular in stabilizing the protein. This demonstrates that even a β-barrel, often regarded as particularly stable, can be strongly impacted by environmental conditions in the functionally important extracellular loops. The sensitivity to environmental conditions also explains discrepancies related to environmental influences affecting crystal and solution structures. The capacity of NMR to site-specifically probe structure and dynamics of membrane proteins in lipid bilayers was the foundation for these mechanistic insights, and is key to understanding proteins in motion. Here specifically, the transmembrane transport mechanism that is implicated by CSP and MD provides the underpinning for potential future biotechnology and bioengineering applications.

Methods

Sample Preparation. Uniformly 13C,15N-labeled and 2H,13C,15N-labeled samples of AlkL (28-230) from P. putida GPo1 were expressed in Escherichia coli, purified, and reconstituted in lipids according to a protocol described previously (34), and recently adapted to accommodate isotopic labeling (35). A full description is provided in SI Appendix, Supplementary Text. Samples in the presence of octane or carvone were prepared in saturated solutions, by addition of a small drop of oil to the surface of the sample buffer. Liposomes were separated by centrifugation before packing in NMR rotors.

NMR Spectroscopy. Solution NMR spectra were recorded at ν2ρ2π = 1 GHz and 600 MHz on Bruker Avance III instruments equipped with cryogenic probes using 2H,13C,15N-labeled AlkL. Solid-state NMR spectra were recorded at ν2ρ2π = 1 GHz and 800 MHz, on Bruker Avance III instruments equipped with 0.7- and 1.3-mm HCN probes spinning at MAS rates of 111.111 kHz (13C,15N-labeled AlkL) or 60 kHz (2H,13C,15N-labeled AlkL). Spectrometer settings, as well as acquisition parameters specific for each 3D and 4D spectrum, and for site-specific 15N R1 (28) and R2 (36) rates, are discussed in SI Appendix, Supplementary Text and summarized in SI Appendix, Table S1. The spectral quality underlying resonance assignment datasets (37, 38) is demonstrated in the pair of CA-N-H correlation spectra shown in SI Appendix, Fig. S13 and the assignments are discussed in detail in a separate manuscript (39). CSPs induced by the presence of carvone or octane were
obtained from (H)CANH and (H)CONH spectra, and were calculated as the RMSD of 1H, 13CA, and 15N shifts, with relative scaling of 1, 0.3, and 0.15, respectively (40).

**Solution NMR Structure Calculation.** Several unambiguous distance restraints were manually assigned in the well-resolved 3D (H)N(HH)NH spectrum, leading, along with dihedral angle restraints, to an initial model. A set of hydrogen bond restraints was generated for resonances that showed no exchange with D2O. NOESY peak lists were added for automated assignment and structure calculation using CYANA 3.98 (41). Further details are discussed in SI Appendix, Supplementary Text, and calculation statistics are reported in SI Appendix, Table S2.

**MAS NMR Structure Calculation.** Several unambiguous distance restraints were manually assigned in the well-resolved 4D (H)NH(H)NH spectrum leading, along with dihedral angle restraints, to a first model. Hydrogen bond restraints were included where cross-peaks in the 4D spectrum were identified with β-sheet chemical shift, and forming a clear pattern of contacts expected for antiparallel β-sheets (SI Appendix, Figs. S3 and S4). The hydrogen bond restraints are indicated in SI Appendix, Fig. S11. R F D R and band-selective spectral spin diffusion–SD (35) peak lists were added for automated assignment and structure calculation using CYANA 3.98 (41). The final structure made use of both backbone and sidechain protons (SI Appendix, Fig. S14). Further details are discussed in SI Appendix, Supplementary Text, and calculation statistics are reported in SI Appendix, Table S2.

**Relaxation Data Fitting.** Signal intensity decays of both 15N R1 and R2 relaxation experiments were fit to a monoexponential function. Error estimate was performed with Monte Carlo simulations: synthetic datasets are produced by adding Gaussian random noise (with the same SD as the experimental noise) to the backcalculated decay curves. The error was determined as the SD of the ensemble of dynamical parameters obtained by fitting 1,000 synthetic datasets.

**MD Simulations.** All simulations were performed with GROMACS 2018 simulation software (42). SI Appendix, Supplementary Text discusses the details...
of the force fields and parameters used, as well as the procedures for preparing the systems prior to MD simulations. Four systems containing AlkL were simulated for 1,000 ns each (“AlkL apo” simulations). After the addition of carvone, four new systems were initially simulated for 1,000 ns each. Further simulations were spawned by monitoring the position of bound carvone and simulated for additional 100 ns in several cycles, as shown in SI Appendix, Fig. 56.

Data Availability. The atomic coordinates and restraints were deposited in the Protein Data Bank, https://www.wwpdb.org/ [PDB ID codes 6QWR (43) and 6QAM (44), and chemical shifts were deposited in the Biological Magnetic Resonance Data Bank, http://www.bmrb.wisc.edu/] [JD 34365 (45) and 34338 (46)].

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Supplementary Information for

A beta-barrel for oil transport through lipid membranes: dynamic NMR structures of AlkL

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This PDF file includes:

- Supplementary text
- Figures S1 to S14
- Tables S1 to S3
- Legend for the Supplementary Movie
- SI References

Other supplementary materials for this manuscript include the following:

- One Supplementary Movie
Supplementary Text

Methods

Sample preparation. AlkL (28-230) from Pseudomonas putida GPo1 (Uniprot Q00595) with a C-terminal His-8 tag was expressed in E. coli as inclusion bodies, purified under denaturing conditions and refolded into DDAO micelles. A detailed protocol for AlkL refolding has been described previously(1) and was recently adapted to accommodate isotopic labelling(2). Isotopic labelling was performed in minimal media containing 4 g/L 13C-glucose and 1 g/L 15NH4Cl (U-CN protein) or 4 g/L 13C,15N-glucose and 1 g/L 15NH4Cl in D2O (U-DCN protein, 100% back exchange of NH-protons during purification).

The detergent was exchanged to 2% octyl glycoside (OG) using a PD10 desalting column for solution NMR as well as for reconstitution into Lipids. The final solution NMR sample contained 330 µM U-DCN AlkL in 2% OG and NMR buffer composed of 20 mM Sodium Phosphate at pH 7, with 10 mM glutamate, 10 mM arginine, and 0.02% sodium azide. Traces of the refolding detergent DDAO may remain. To identify residues involved in hydrogen bonds, the buffer was exchanged to 20 mM Sodium Phosphate pH 7, 95% D2O using a desalting column. In order to allow solution NMR analysis of AlkL in lipids, a nanodisc sample was prepared according to published protocols using membrane scaffold protein (MSP1D1)(3).

Lipid bilayer samples for solid-state NMR were prepared by addition of DMPC (10 mg/ml, multilamellar vesicles in H2O) to AlkL in OG detergent at a lipid to protein ratio of 1:2 (w/w). Protein and lipids form soluble mixed micelles at this stage. Detergent was subsequently removed by dialysis against NMR buffer resulting in formation of multilamellar vesicles that precipitate within 24 hours. The white precipitate was collected by centrifugation using a table top centrifuge, resuspended in 0.5 ml buffer and packed into 1.3 or 0.7 mm rotors (Bruker) using an ultracentrifuge packing device (Giotto Biotech). To investigate the effect of the presence of a hydrophobic small molecule, a solid-state NMR sample was equilibrated with buffer saturated with carvone or octane and compared with a second rotor packed from the same batch of precipitated protein. The bilayer sample in LPS (Sigma, phenol extract purified from E. Coli K-235) were prepared in an identical manner, except that the reconstitution was performed from AlkL in OG detergent with a ratio of DMPC:LPS:AlkL of 1:1:2 (by weight). The pellet was nearly transparent, different from the white precipitate that forms in the presence of DMPC alone. The spectrum (Fig S5) is nearly unchanged, compared as different from the DMPC preparation.

Solution NMR Spectroscopy. Triple resonance solution NMR spectra were recorded on a 600 MHz Bruker Avance III instrument equipped with a cryogenic probe. TROSY based HSQC, HNCA, HN(CA)CO, HNCO, and HN(CA)CO spectra were used for resonance assignment(4). To obtain distance restraints, a 3D NOESY-15N-TROSY (NHH) and a 3D 15N-HSQC-NOESY-15N-HSQC (NNH) was recorded on a 1000 MHz Bruker Avance III instrument equipped with a cryogenic probe. All spectra were recorded at a sample temperature of 308 K.

Solid state NMR Spectroscopy. Spectra for the assignment of backbone and sidechain resonances were (H)NH, (H)CH, (H)CANH, (H)(CO)CA(CO)NH, (H)CONH, (HCA)CB(CA)NH, (HCA)CB(CACO)NH, (H)NCAHA, (H)N(CA)CAHA, (H)CCH, (H)COC(AHA), (H)CO(N)CAHA. These spectra were recorded on U-CN AlkL in a 0.7 mm MAS rotor at 111 kHz on a 1000 MHz Bruker Avance III spectrometer (5, 6). Additionally, we recorded spectra for assignments: (H)CANH, (HCO)CA(CO)NH, (H)CONH, (HCO)(CA)NH, (HCA)CB(CA)NH, on U-DCN AlkL in a 1.3 mm MAS rotor at 60 kHz on a 800 MHz Bruker Avance III spectrometer. To obtain distance restraints, we recorded H(NH)NH and H(H)CH using RFDR, H(NH), (H)(NH)NH and (H)(HH)CH using BASS-SD on U-CN AlkL as well as a 4D (H)NH(NH)NH using RFRD on U-DCN AlkL(2, 7). Further experimental details are tabulated in Extended Data Table 1. Relaxation rates (15N R1 and R2) were determined using a modified 3D (H)CONH experiment (pseudo-4D), with the longest relaxation delay at 54 seconds for R1 and 200 ms for R2. Chemical shift perturbations induced by the presence of either octane or carvone were measured under saturating concentrations of the small molecule, and were evaluated as a combination of 1H, 15N and 13CA shift changes, with relative scaling of 1, 0.15, and 0.3: CSP = \sqrt{\frac{1}{2}(2\Delta_\nu^2 + 0.15\Delta_\nu^2 + 0.3\Delta_\gamma^2)}). All spectra were recorded at a sample temperature of 305 ± 3 K.
| Experiment | H(H)NH | H(H)NH | (H)(HH)NH | H(H)CH | (H)(C(HH))CH | (H)NH(H)NH | (H)NH(H)NH |
|------------|--------|--------|------------|--------|-------------|------------|------------|
| Sample     |       |        |            |        |             |            |            |
| Spectrometer (MHz) | 1000  | 1000  | 1000       | 1000  | 1000        | 1000       | 800        |
| MAS, ν (kHz) | 111   | 111   | 111        | 111   | 111         | 60          | 60         |
| Heteronuclear transfer 1 | H-N CP | H-N CP | H-N CP   | H-C CP | H-C CP      | H-N CP     | H-N CP     |
| Field (kHz) | 1.4/0.4 v. | 1.4/0.4 v. | 1.4/0.4 v. | 1.4/0.4 v. | 1.6/0.6 v. | 1.7/0.7 v. |
| Time (ms)* | 1      | 0.6    | 0.6        | 0.5    | 0.35        | 1           | 1          |
| Heteronuclear transfer 2 | N-H CP | N-H CP | N-H CP | C-H CP | C-H CP     | N-H CP     | N-H CP     |
| Field (kHz) | 0.4/1.4 v. | 0.4/1.4 v. | 0.4/1.4 v. | 0.4/1.4 v. | 0.6/1.6 v. | 0.7/1.7 v. |
| Time (ms)* | 0.4    | 0.3    | 0.3        | 0.35   | 0.19        | 0.6         | 0.6        |
| H-H Transfer | RFDR  | BASS-SD | BASS-SD | RFDR  | BASS-SD  | RFDR       | RFDR       |
| Field (kHz) | 192   | 6.25   | 6.25      | 192    | 4.1        | 100         | 100        |
| Time (ms)* | 0.5    | 6      | 6         | 0.5    | 4.5        | 3           | 0.5        |
| Heteronuclear transfer 3 | –      | –      | H-N CP | –      | H-C CP     | H-N CP     | H-N CP     |
| Field (kHz) | –      | –      | 1.4/0.4 v. | –      | 1.4/0.4 v. | 1.6/0.6 v. | 1.7/0.7 v. |
| Time (ms)* | –      | –      | 0.6      | –      | 0.35       | 1           | 1          |
| Heteronuclear transfer 4 | –      | –      | N-H CP | –      | C-H CP     | N-H CP     | N-H CP     |
| Field (kHz) | –      | –      | 0.4/1.4 v. | –      | 0.4/1.4 v. | 0.6/1.6 v. | 0.7/1.7 v. |
| Time (ms)* | –      | –      | 0.3      | –      | 0.19       | 0.6         | 0.6        |
| sw (t1) (ppm) | 12.5  | 35     | 35        | 12.5   | 38         | 6.5         | 7          |
| Acq.time (t1) (ms) | 4.6   | 9      | 9         | 4.6    | 7.7        | 4.4         | 2.2        |
| sw (t2) (ppm) | 66    | 5.5    | 35        | 66     | 38         | 40          | 35         |
| Acq.time (t2) (ms) | 16.4  | 3.8    | 9         | 6.6    | 8.7        | 9.1         | 9.1        |
| sw (t3) (ppm) | 100   | 100    | 100       | 100    | 100        | 40          | 35         |
| Acq.time (t3) (ms) | 8     | 8      | 8         | 8      | 8          | 9.1         | 8.8        |
| sw (t4) (ppm) | –     | –      | –         | –      | –          | 100         | 30         |
| Acq.time (t5) (ms) | –     | –      | –         | –      | –          | 8           | 8          |
| H decoupling | swTPPM | swTPPM | swTPPM | swTPPM | swTPPM | waltz | swTPPM |
| Field (kHz) | 26    | 26     | 26        | 26     | 26         | 7.2         | 13         |
| Interscan delay (s) | 1     | 0.8    | 0.8       | 1      | 0.8        | 1.25        | 1          |
| Number of scans | 8     | 80     | 240       | 8      | 16         | 48          | 16         |
| NUS (%) | –     | –      | –         | –      | –          | 2           | –          |
| Measurement time (d) | 3     | 2.4    | 11.5      | 3      | 4.6        | 5.5         | 13         |
**Resonance Assignment.** We used FLYA (8) with peak lists generated from spectra of the fully-protonated sample automatically in Sparky (9) or CCPN (10). The threshold for peak identification was initially set such that only an estimated 5-10 false peaks were present (as determined by the number of peaks picked with incorrect sign). Subsequently, overlapping peaks and weaker peaks (that were not initially picked) were manually identified to extend the assignment. The automated assignments were verified and additional assignments completed manually.

**Structure Calculation.** Several unambiguous distance restraints were manually assigned in the well resolved 3D (H)N(HH)NH spectrum for the solution sample. Similarly, the 4D (H)NH(H)NH spectrum was used to identify unambiguous restraints in the lipid bilayer sample. We additionally used the chemical shifts to generate dihedral angle restraints using TALOS-N (11). These initial unambiguous restraints and dihedral angles clearly defined the β-barrel region, and additional structure in the loops for the lipid bilayer preparation. For the solution structure, an initial set of hydrogen bond restraints was defined for resonances that showed no exchange in D₂O. For both structures, hydrogen bond restraints were included where cross-peaks in the 4D spectrum were identified (Fig S3-4), and where chemical shifts were consistent with an extended conformation, as indicated in Fig. S9. A short stretch of helical hydrogen bonds in loop 2 was introduced for the bilayer structure, where chemical shifts indicated, and where an initial model showed a helical structure (Fig. S9). NOESY peak lists (solution) or RFDR and BASS-SD peak lists (solids) were added for automated assignment and structure calculation using CYANA 3.98 (12). Final numbers are listed in Extended Data Table 2. Figures were prepared with UCSF Chimera (13).

**Table S2:** Structure calculation statistics. Summary of identified cross-peaks and conformational restraints used in the structure calculation, and structure quality in terms of RMSD within a bundle of 20 conformers used to represent the ensemble.

| Distance restraints       | Lipids (MAS) | Detergents (solution) |
|--------------------------|--------------|-----------------------|
| Total                    | 769          | 112                   |
| Short-range (|i-j|<=1)     | 425          | 53                    |
| Medium-range (1<|i-j|<5) | 21           | 3                     |
| Long-range (|i-j|>=5)     | 323          | 56                    |
| h-bonds (manual restraints) | 94           | 64                    |
| Dihedral angle restraints | 303          | 206                   |
| Restraints per residue `*` | 3.1 (3.7`)  | 1.3 (2.8`)            |
| Backbone RMSD (Å) `*`    | 0.66         | 0.73 (#)              |
| Heavy atom RMSD (Å) `*`  | 1.40         | 1.52 (#)              |

*calculated over structured regions, residues 12-33, 44-108, 116-156, 161-205 (22+65+41+45=173 residues)

`*`excluding short-range restraints, excluding H-bond restraints to avoid overcounting.

#calculated over structured regions, residues 13-27, 52-74, 100-107, 116-126, 146-169, 193-205 (15+23+8+11+24+13=94 residues)

**Table S3.** Residue specific chemical shift perturbation (CSP) and amide $^{15}$N relaxation.

| Residue | Octane CSP (ppm) | Carvone CSP (ppm) | Apo -N R, (s⁻) | Apo -N R, (s⁻) |
|---------|------------------|-------------------|----------------|----------------|
| 12      | 0.050            | 0.024             |                |                |
| 13      | 0.094            | 0.034             |                |                |
| 14      | 0.022            | 0.033             | 0.047 ±0.004   | 12.0 ±0.06     |
| 15      | 0.026            | 0.030             |                |                |
| 16      | 0.035            | 0.036             |                |                |
| 17      | 0.049            | 0.031             |                |                |
| 18      | 0.034            | 0.082             |                |                |
| 19      | 0.063            | 0.026             | 0.019 ±0.005   | 7.7 ±0.12      |
|    |     |     |     |     |
|----|-----|-----|-----|-----|
| 20 | 0.035 | 0.039 | 9.2 ±0.11 |
| 21 | 0.167 | 0.069 | 8.6 ±0.09 |
| 22 | 0.057 | 0.056 | 0.014 ±0.003 | 10.4 ±0.10 |
| 23 | 0.050 | 0.111 | 0.024 ±0.005 |
| 24 |     |     |     |     |
| 25 | 0.159 | 0.062 |     |     |
| 26 | 0.268 |     |     |     |
| 27 | 0.080 |     |     |     |
| 28 | 0.078 |     |     |     |
| 29 | 0.097 |     |     |     |
| 30 | 0.042 | 0.044 | 0.093 ±0.045 | 17.7 ±0.26 |
| 31 | 0.332 |     | 0.040 ±0.013 | 22.8 ±0.57 |
| 32 | 0.047 |     |     | 11.7 ±0.21 |
| 33 | 0.054 |     |     |     |
| 34 |     |     |     |     |
| 35 | 0.024 |     |     |     |
| 36 | 0.142 |     |     |     |
| 37 | 0.073 |     |     |     |
| 38 |     |     |     |     |
| 39 | 0.113 | 0.032 |     |     |
| 40 | 0.030 | 0.036 |     |     |
| 41 | 0.283 | 0.107 | 0.043 ±0.010 | 16.3 ±0.21 |
| 42 |     |     |     |     |
| 43 | 0.026 | 0.067 |     |     |
| 44 | 0.072 | 0.094 |     |     |
| 45 | 0.091 | 0.113 | 0.019 ±0.003 | 9.4 ±0.11 |
| 46 | 0.147 | 0.082 | 0.024 ±0.006 | 16.3 ±0.30 |
| 47 | 0.133 | 0.096 |     |     |
| 48 | 0.098 | 0.049 | 0.015 ±0.002 | 10.3 ±0.13 |
| 49 | 0.055 | 0.035 | 0.014 ±0.004 | 5.9 ±0.06 |
| 50 | 0.055 | 0.018 |     |     |
| 51 | 0.020 | 0.033 | 0.013 ±0.003 | 7.8 ±0.06 |
| 52 | 0.046 | 0.051 | 0.025 ±0.006 | 15.0 ±0.25 |
| 53 | 0.032 | 0.053 |     |     |
| 54 | 0.034 | 0.045 |     |     |
| 55 | 0.027 | 0.040 |     |     |
| 56 | 0.024 | 0.010 |     | 108.4 ±3.8 |
| 57 | 0.030 | 0.032 | 0.023 ±0.002 | 7.7 ±0.04 |
| 58 | 0.020 | 0.025 | 0.014 ±0.002 | 5.8 ±0.03 |
| 59 | 0.035 | 0.023 |     |     |
| 60 | 0.119 | 0.090 | 0.010 ±0.003 | 8.0 ±0.06 |
| 61 | 0.045 |     |     |     |
| 62 |     |     |     |     |
| 63 | 0.028 ±0.009 |     | 12.0 ±0.14 |
| 64 |     |     |     |     |
| 65 | 0.365 | 0.188 |     | 28.0 ±0.67 |
| 66 | 0.121 |     |     |     |
| 67 | 0.168 | 0.372 | 0.040 ±0.009 | 14.6 ±0.18 |
| 68 | 0.101 | 0.230 |     |     |
| 69 | 0.169 | 0.025 |     |     |
| 70 | 0.154 | 0.166 |     |     |
|   |   |   |   |
|---|---|---|---|
| 83 | 0.084 | 0.049 | 0.023 ±0.004 |
| 84 | 0.054 | 0.018 | 0.012 ±0.009 |
| 85 | 0.029 | 0.041 | 0.037 ±0.004 |
| 86 | 0.099 | 0.062 |   |
| 87 | 0.060 | 0.039 |   |
| 88 | 0.150 | 0.049 | 0.091 ±0.029 |
| 89 | 0.087 | 0.055 | 0.040 ±0.010 |
| 90 | 0.124 | 0.068 |   |
| 91 | 0.082 | 0.071 | 0.074 ±0.011 |
| 92 | 0.028 | 0.038 | 0.052 ±0.005 |
| 93 | 0.047 | 0.041 | 0.029 ±0.005 |
| 94 | 0.026 | 0.059 | 0.017 ±0.002 |
| 95 | 0.060 | 0.043 |   |
| 96 | 0.094 | 0.099 | 0.100 ±0.007 |
| 97 | 0.028 | 0.038 | 0.052 ±0.005 |
| 98 | 0.054 | 0.018 | 0.012 ±0.009 |
| 99 | 0.339 | 0.066 | 0.047 ±0.013 |
| 100 |   |   |   |
| 101 | 0.091 | 0.099 | 0.100 ±0.007 |
| 102 | 0.087 | 0.068 | 0.088 ±0.002 |
| 103 | 0.030 | 0.020 | 0.014 ±0.002 |
| 104 | 0.026 | 0.044 | 0.013 ±0.002 |
| 105 | 0.056 | 0.057 |   |
| 106 | 0.027 | 0.042 | 0.010 ±0.002 |
| 107 | 0.004 | 0.044 |   |
| 108 | 0.123 | 0.078 | 0.006 ±0.002 |
| 109 | 0.041 |   |   |
| 110 |   |   |   |
| 111 | 0.077 | 0.011 |   |
| 112 | 0.036 | 0.029 |   |
| 113 | 0.059 | 0.027 | 0.019 ±0.003 |
| 114 | 0.015 | 0.022 | 0.013 ±0.004 |
| 115 | 0.030 | 0.055 |   |
| 116 | 0.044 | 0.045 | 0.018 ±0.004 |
| 117 | 0.101 | 0.123 | 0.018 ±0.004 |
| 118 | 0.129 | 0.087 |   |
| 119 | 0.337 | 0.170 |   |
| 120 | 0.256 | 0.176 | 0.016 ±0.007 |
| 121 | 0.022 |   |   |
| 122 | 0.021 |   |   |
| 123 | 0.031 | 0.051 | 0.034 ±0.006 |
| 124 | 0.066 | 0.135 |   |
| 125 | 0.082 | 0.072 | 0.028 ±0.004 |
| 126 | 0.014 | 0.022 |   |
| 127 | 0.029 | 0.054 | 0.030 ±0.005 |
| 128 | 0.135 | 0.046 | 0.030 ±0.006 |
| 129 | 0.046 | 0.051 |   |
| 130 | 0.067 | 0.037 |   |
| 131 | 0.052 | 0.035 | 0.016 ±0.003 |
| 132 | 0.028 | 0.017 |   |
| 133 | 0.029 | 0.015 |   |
|   |         |         |          |         |
|---|---------|---------|----------|---------|
| 141 | 0.097   | 0.063   | 6.0 ±0.06 |
| 142 | 0.223   | 0.206   | 0.041 ±0.009 | 8.7 ±0.01 |
| 143 | 0.028   | 0.022   | 0.045 ±0.005 | 7.7 ±0.04 |
| 144 | 0.133   | 0.050   | 9.4 ±0.12  |
| 145 | 0.133   | 0.104   | 9.3 ±0.21  |
| 146 | 0.064   | 0.111   | 9.8 ±0.12  |
| 148 | 0.057   | 0.017   | 8.3 ±0.06  |
| 149 | 0.044   | 0.065   |           |
| 150 | 0.043   | 0.030   | 0.009 ±0.002 | 9.1 ±0.05 |
| 151 | 0.012   | 0.031   | 0.011 ±0.002 | 7.9 ±0.04 |
| 152 | 0.035   | 0.060   |           |
| 153 | 0.022   | 0.043   | 0.022 ±0.003 | 7.7 ±0.05 |
| 154 | 0.031   | 0.051   |           |
| 155 | 0.022   | 0.024   |           |
| 156 | 0.054   | 0.040   |           |
| 157 | 0.049   |         |           |
| 158 | 0.033   |         |           |
| 161 | 0.176   | 0.102   |           |
| 162 | 0.012   | 0.033   | 0.025 ±0.008 | 10.4 ±0.11 |
| 163 | 0.031   | 0.034   |           |
| 164 | 0.026   | 0.027   |           |
| 165 | 0.035   | 0.040   | 0.029 ±0.003 | 6.6 ±0.04 |
| 166 | 0.048   | 0.028   | 0.027 ±0.006 | 9.3 ±0.09 |
| 167 | 0.069   | 0.071   | 0.014 ±0.002 | 7.1 ±0.04 |
| 168 | 0.069   | 0.115   |           |
| 169 | 0.107   | 0.070   |           |
| 171 | 0.348   | 0.283   |           |
| 173 | 0.264   | 0.204   |           |
| 174 | 0.034   |         | 0.053 ±0.007 | 10.5 ±0.01 |
| 175 | 0.144   | 0.055   | 0.090 ±0.015 | 8.9 ±0.07 |
| 176 | 0.057   | 0.017   |           |
| 177 | 0.039   | 0.033   |           |
| 178 | 0.019   | 0.032   | 0.018 ±0.002 | 6.8 ±0.04 |
| 179 | 0.065   | 0.034   | 0.026 ±0.003 | 8.0 ±0.05 |
| 180 | 0.212   | 0.042   |           |
| 181 | 0.182   | 0.055   |           |
| 183 | 0.003   | 0.017   |           |
| 185 | 0.044   | 0.020   |           |
| 186 | 0.086   | 0.027   |           |
| 187 | 0.205   | 0.105   | 0.040 ±0.009 | 11.1 ±0.15 |
| 188 | 0.112   | 0.066   | 13.0 ±0.18 |
| 189 | 0.057   | 0.053   |           |
| 190 | 0.053   | 0.022   |           |
| 191 | 0.156   |         | 14.7 ±0.17 |
| 194 |         |         |           |
| 195 | 0.352   | 0.123   |           |
| 196 | 0.033   | 0.008   | 0.014 ±0.002 | 6.1 ±0.06 |
| 197 | 0.152   | 0.129   | 0.021 ±0.005 | 11.0 ±0.23 |
| 198 | 0.031   | 0.055   |           |
MD Simulations: System preparation - AlkL NMR structure. The NMR structure of AlkL was aligned along the membrane normal using the Orientations of Proteins in Membranes (OPM) database (14) and subsequently embedded in a DMPC or LPS membrane and solvated with water molecules and 150 mM KCl using the CHARMM-GUI webserver (15). The full system contained ~160 000 atoms. The titratable groups of the protein were protonated according to their standard protonation states at pH 7, with an exception of D58, which was protonated, based on the spatial proximity of another negatively charged residue, D70 and robust binding of K+ ion to (charged) D58 in initial simulations. Further, pKa values predicted by PROPKA (16) suggested that one of these aspartates (D58 or D70) should be in its protonated form.

The initial, multi-step equilibration of the system, with a gradual release of restraints acting on protein atoms, was conducted using scripts provided by CHARMM-GUI. Subsequently, the system was equilibrated for 100 ns, without any restraints, prior to production runs.

To study interactions and possible permeation of hydrophobic molecules through AlkL, 64 carvone molecules were placed in the aqueous phase of the last snapshot from the equilibration run.

MD Simulations: System preparation - AlkL homology models. Two homology models of AlkL were built, using structures of OmpW and OprG proteins (PDB id: 2F1T and 2X27, respectively) as templates using the Swiss-model web server (swissmodel.expasy.org). Both homology models were then prepared for MD simulations in the same manner as the NMR structure.

MD Simulation details. All simulations were performed with GROMACS 2018 simulation software (17). The CHARMM36m force field was used in all simulations (18). Lipids were modeled using the CHARMM36 force field (19). The P. Aruginosa LPS was chosen with Lipid A type 1, core 2, and 2x the standard O-antigen in the CHARMM gui (www.charmm-gui.org) with calcium counterions. The parameters for carvone were obtained from the CHARMM General Force Field (CGenFF) (20) webserver (https://cgenff.paramchem.org). Water was modeled using the CHARMM version of the TIP3P model (21), with LJ interactions placed on hydrogen atoms. Standard CHARMM parameters were used for K+, Ca2+ and Cl- ions (22). All bonds were constrained using LINCS, allowing for a 2fs time step (23). Non-bonded (van der Waals) interactions were force-switched off from 0.8 to 1.2 nm. Long-range electrostatic interactions were treated with PME with a 1.2 nm real space cutoff (24). The simulated systems were kept at a temperature of 320K and a pressure of 1 bar, using a Nosé-Hoover thermostat (25, 26) and a Parrinello-Rahman barostat (27), respectively.

For LPS simulations (Fig. S6), an initial simulation was carried out for 100 ns and 10 equidistant frames from this trajectory were extracted. Into each of these 10 frames, 100 randomly placed octanes were inserted, replacing about 1000 water molecules. Each of these 10 octane-containing frames were independently equilibrated 10 times and the 100 simulations were run for a total of ~15 μs. The top 10 trajectories (defined by the octane with the most contacts with AlkL and manually verified to be close to the exit site) out of these initial 100 were chosen and another 10 replicas from each were spawned and run for a total of ~50 μs.

The simplified DMPC-octane simulations were run in the presence of 64 randomly placed octanes in 11 independent simulations for a total of ~9 μs (Fig. S6).

Four systems containing AlkL in DMPC were simulated for 1000 ns each (‘Alk apo’ simulations). After the addition of carvone, four new systems were initially simulated for 1000 ns each. Further simulations were spawned by monitoring the position of bound carvone and simulated for additional 100 ns in several cycles, as shown in Fig. S7.

| Data | Value | Value | Value | Value |
|------|-------|-------|-------|-------|
| 199  | 0.031 | 0.019 | 0.010 ±0.003 | 9.0 ±0.07 |
| 200  | 0.070 | 0.037 | 0.016 ±0.004 | 7.2 ±0.07 |
| 201  | 0.059 | 0.028 | 0.016 ±0.005 | 8.3 ±0.09 |
| 202  | 0.041 | 0.066 | 0.019 ±0.004 | 9.6 ±0.09 |
| 203  | 0.045 |       |       |       |
| 204  | 0.098 |       |       |       |

Relaxation data fitting. Signal intensity decays of both 1H R1 and R2 relaxation experiments were fitted to a monoexponential function. Error estimate was performed with Monte Carlo simulations: synthetic datasets are produced by adding Gaussian random noise (with same standard deviation as the experimental noise) to the back-calculated decay curves. The error was determined as the standard deviation of the ensemble of dynamical parameters obtained by fitting 1000 synthetic datasets.

| Data | Value | Value | Value | Value |
|------|-------|-------|-------|-------|
| 199  | 0.031 | 0.019 | 0.010 ±0.003 | 9.0 ±0.07 |
| 200  | 0.070 | 0.037 | 0.016 ±0.004 | 7.2 ±0.07 |
| 201  | 0.059 | 0.028 | 0.016 ±0.005 | 8.3 ±0.09 |
| 202  | 0.041 | 0.066 | 0.019 ±0.004 | 9.6 ±0.09 |
| 203  | 0.045 |       |       |       |
| 204  | 0.098 |       |       |       |
Systems containing homology models of AlkL were simulated with and without carvone for 1000 ns in 4 copies each. The total simulation time of all systems was ~34 μs and for the entire study, about 108 μs.

**MD Simulation analysis.** Exit pathways were identified and analysed using the MOLE 2.0 software (28) and CONAN (29). Remaining analysis was performed using GROMACS tools: cluster, covar, anaeig, distance and in-house written scripts. Molecular structures were rendered using VMD (30). Plots were made using NumPy and Matplotlib.
Supporting Figures

Figure S1. Negative stain EM images of the preparations used for NMR. Liposomes are observed. No protein aggregate was identified, and no evidence for ordered 2D crystalline arrays could be found.

Figure S2. Lipid to protein magnetization transfer shows the lipid-embedded protein surface. In (A), the lipid plane of a 3D HhNH spectrum at about 1.1 ppm is colored in yellow. In (B) selected strips from a similar 4D spectrum, HhnCANH resolves resonance overlap in the 3D spectrum. 50 ms of Nuclear Overhauser Effect (NOE) mixing was used. In C, the residues for which lipid contacts are observed are colored in yellow on the solid-state NMR structure. Spectra were acquired at a 800 MHz spectrometer with 55 kHz MAS. The protein was deuterated to suppress any protein signals arising at the proton frequency of the lipid acyl chains.
Figure S3. Spectra at varying DMPC lipid concentration. In (A), the lipid protein ratio (LPR) of 10 (blue) is compared with a LPR of 0.5 (red). The spectra are nearly identical, and notably, several well-resolved resonances from the loops are unchanged in the dilute sample. In (B), degradation of the spectral quality is observed for an over concentrated sample with LPR of 0.25 (black) compared with a sample at 0.5 LPR (red). All spectra were recorded on a 700 MHz spectrometer with a spinning frequency of 60 kHz. Spectra in (A) were recorded with deuterated protein, and those in (B) with protonated protein.
Figure S4. Solution $^{15}$N-TROSY spectra of AlkL. In (A and D), unassigned resonances are indicated with red crosses. In (B), assignments are annotated. In (C), the red spectrum was recorded after 24 hours in D$_2$O, indicating exceptionally stable hydrogen bonds, which were used in the structure calculation. In (D), the black spectrum was recorded on AlkL in MSP1D1 DMPC nanodiscs. Only signals arising from highly flexible loops are detected, prohibiting structural analysis under these conditions. Notably, many signals that were flexible and unassigned in (A) are not detected. This data is consistent with the MAS NMR spectra in which we were not able to assign 18 N- and C-terminal residues, as well as part of L1, T2 and T3, and the His$_8$ tag. The absence of peaks in assignment spectra indicates that some of these unassigned resonances correspond to residues that are highly flexible.
Figure S5. Inter-proton proximities. Selected planes from two 4D (H)NH(H)NH spectra (using RFDR for proton-proton mixing). In blue, the spectrum was recorded with high resolution, but lower sensitivity, by sampling the proton dimension to 4 ms. In red, the spectrum was recorded with shorter indirect proton evolution of 2 ms which resulted in higher sensitivity, but lower resolution. The first 2 planes (connecting residues 17-204) show an example where the peak was just above noise in the blue spectrum, and clearly found in the red spectrum. The third 2 planes (connecting residues 21-200) shows an example where the longer evolution times in the blue spectrum was necessary, and the peak is indistinguishable from the diagonal in the red spectrum. Both spectra were used in conjunction to construct a consistent set of nearly unambiguous contacts, and to build the initial model (see Fig. S6).

Figure S6. Proton-proton contact maps. In (A), automatically generated contacts from peaks picked from the (H)NH(H)NH spectra of Fig. S5. A threshold of 0.5 ppm (15N) and 0.11 ppm (1H) were used to generate all possible assignments. Contacts arising from one side of the spectrum’s diagonal are shown in red, while those from the other are shown in blue. A clear pattern of antiparallel β-strands emerges, despite some degeneracy in the 1H-15N plane used to identify starting and ending signal. In (B), the contact map of the final structure is shown, after resolution of ambiguities using CYANA, and including contacts from all spectra. The additional restraint assignment possibilities in (A) arising from degenerate chemical shifts have been removed in (B).
Figure S7. Solution and solid-state NMR spectra of AlkL. In A, the solution $^{15}$N-$^1$H HSQC spectrum (red) is compared with the dipolar $^{15}$N-$^1$H correlation acquired on the DMPC lipid bilayer sample (blue). Residues that were assigned in lipids, but could not be assigned in solution are encircled. Some of the residues outside the TM region displayed large differences in chemical shift, such as R78, Y109, L127, and W161, indicating the change from ordered to disordered. The remaining unassigned amide proton shifts in solution fall between 7.5 and 8.7 ppm, indicating a lack of β-sheet secondary structure. In B, the DMPC sample (blue) is compared with a sample reconstituted in 50% lipopolysaccharide (LPS) with a ratio of 1:0.5:0.5 AlkL:DMPC:LPS (w/w/w) (black). The spectra were contoured from 25% of the intensity of the G178 peak. The spectrum obtained in LPS, with only minor chemical shift changes, indicates that the structured extracellular loops are stable in the near native environment including lipopolysaccharides that are typically found on the outside of the asymmetric bacterial outer membrane. In the reconstituted sample, neither the AlkL, nor the LPS are oriented.
Figure S8. Simulation protocol for octane permeation through AlkL. Single numbers label individual simulation replicas. Simulations in which an octane molecule spontaneously permeated through the protein are indicated in red, along with the number of transit events observed.
Figure S9. Simulation protocol for carvone permeation through AlkL. Single numbers label individual simulation replicas. Simulations in which a carvone molecule spontaneously permeated through the protein are labelled with boxes 'carvone out'. The 40 simulations referred to as 'AlkL with carvone' in the main text are enclosed within the rounded maroon boxes.
**Figure S10.** Dynamic parameter perturbations highlight weak binding sites of carvone. In A, transverse relaxation rates are shown for apo (red) and carvone saturated (blue) samples, as determined from (H)CONH spectra. In B, the difference in rates is shown on the structure.

**Figure S11.** Topology map of AlkL as determined by NMR, both in detergent micelles (A) and in lipid bilayers (B). Assigned residues with chemical shifts consistent with β-sheet conformation are indicated in boxes, while other assigned residues are indicated with white circles. Unassigned residues are in grey circles. Blue residues point toward the interior. In lipids, a hydrophobic volume is formed in the extracellular barrel extension. The interior of the transmembrane β-barrel is packed densely with the side-chains of charged residues, while hydrophobic residues point out, as expected. Intracellular turns are numbered as T1-T3, and extracellular loops are numbered L1-L4. In both panels, in order to render the circular barrel topology, the last β-strand is repeated on the right-hand side and grayed out. The approximate position of the membrane is indicated by horizontal lines. The new and previous proposed exit sites are shown as red and yellow ovals, respectively, and packing interactions in OmpW crystals are indicated by green lines.
Figure S12. Chemical shift perturbations due to saturation with carvone (purple), which has about 7-8 mM solubility in water (31) and 1 mg/ml (8 mM) Leucine (blue). Shift changes of similar magnitude are observed with octane, which has a very low solubility in water (only about 60 nM according to the CDC NIOSH https://www.cdc.gov/niosh/npg/npgd0470.html. The shift changes due to carvone are evident in the 2D spectrum. For leucine, no shift changes are seen, consistent with the expectation that AlkL does not act as a channel for amino acids.
Figure S13. Backbone resonance assignment. Linking of sequential NH moieties in the (H)CANH (green) and (H)(CO)CA(CO)NH (purple) spectra of $^2$H,$^{13}$C,$^{15}$N AlkL in lipid bilayers. Strips corresponding to each assigned NH group are shown. The sequence specific assignment is indicated above each strip, and a slash indicates where assignments are missing.
Figure S14. Solid-state NMR $^1$H-$^{13}$C dipolar correlation spectrum recorded on lipid bilayer embedded U-$^{13}$C,$^{15}$N AlkL using 111 kHz magic-angle spinning. Assignments are shown for the C-H region (right), as well as for selected isolated side-chain peaks (left).

SI Movie. The motion along the first (largest) eigenvector in a principal component analysis of MD snapshots with and without the molecule carvone. The motion shows that the difference without and with bound carvone is related to the opening of the newly identified site I for carvone permeation.
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