Substrate Access Mechanism in a Novel Membrane-Bound Phospholipase A of *Pseudomonas aeruginosa* Concordant with Specificity and Regioselectivity

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**ABSTRACT:** PlaF is a cytoplasmic membrane-bound phospholipase A₁ from *Pseudomonas aeruginosa* that alters the membrane glycerophospholipid (GPL) composition and fosters the virulence of this human pathogen. PlaF activity is regulated by a dimer-to-monomer transition followed by tilting of the monomer in the membrane. However, how substrates reach the active site and the characteristics of the active site tunnels determine the activity, specificity, and regioselectivity of PlaF for natural GPL substrates have remained elusive. Here, we combined unbiased and biased all-atom molecular dynamics (MD) simulations and configurational free-energy computations to identify access pathways of GPL substrates to the catalytic center of PlaF. Our results map out a distinct tunnel through which substrates access the catalytic center. PlaF variants with bulky tryptophan residues in this tunnel revealed decreased catalysis rates due to tunnel blockage. The MD simulations suggest that GPLs preferably enter the active site with the acyl chain-length specific to the catalytic center. PlaF variants with bulky tryptophan residues in this tunnel revealed decreased catalysis rates due to tunnel blockage. The MD simulations suggest that GPLs preferably enter the active site with the sn-1 acyl chain first, which agrees with the experimentally demonstrated PLA₁ activity of PlaF. We propose that the acyl chain-length specificity of PlaF is determined by the structural features of the access tunnel, which results in favorable free energy of binding of medium-chain GPLs. The suggested egress route conveys fatty acid (FA) products to the dimerization interface and, thus, contributes to understanding the product feedback regulation of PlaF by FA-triggered dimerization. These findings open up opportunities for developing potential PlaF inhibitors, which may act as antibiotics against *P. aeruginosa*.

1. INTRODUCTION

*Pseudomonas aeruginosa* is an opportunistic and versatile pathogen, which causes infections in plants and humans. It is a multi-drug resistant Gram-negative bacterium and a frequent cause of nosocomial infections. The pathogenicity of *P. aeruginosa* relies on both cell-associated and extracellular virulence factors. Among those virulence factors are phospholipases, including phospholipase A₁ (PLA₁), which hydrolyze cellular glycerophospholipids (GPLs) at the sn-1 position into lysoglycerophospholipids (LGPLs) and fatty acids (FAs).

GPLs primarily form bilayers, which maintain a permeability barrier for cells and organelles, while membrane-bound LGPLs can destabilize membrane integrity in Gram-negative bacteria. GPLs and LGPLs can regulate the function and stability of membrane proteins. Interestingly, biofilm formation and growth phase transitions in *P. aeruginosa* are accompanied by the alteration of the membrane GPL composition. FAs belong to the diffusable signal factor family (DSF) and are possible signal molecules because they can diffuse through cell membranes and contribute to the regulation of diverse biological functions in various Gram-negative pathogens. In *P. aeruginosa*, DSFs promote biofilm formation and antibiotic resistance.

We recently identified PlaF, an integral, inner membrane PLA₁ that has a profound role for membrane GPL remodeling in *P. aeruginosa*. Furthermore, a *P. aeruginosa* ΔPlaF knockout strain showed strongly attenuated virulence in *Galleria mellonella* and human macrophages models compared to the wild-type, which suggests that PlaF-mediated GPL remodeling contributes to the virulence of *P. aeruginosa*. Cross-linking (in vivo and in vitro) and micro-scale thermophoresis experiments showed that PlaF exists in both monomeric and dimeric configurations, although it is active only in the monomeric state. The crystal structure of PlaF revealed that a homodimer is formed by interactions between the transmembrane (TM) and juxtamembrane (JM) regions. The homodimer contains co-crystallized endogenous ligands, myristic acid (MYR), and undecanoic acid (UND) from *P. aeruginosa* (Figure 1), which are non-covalently bound in the active site cavity. Moreover, a
OMPLA is an integral information about GPL recognition by this PLA. However, structural analysis of the outer membrane PLA (OMPLA) from the phosphate group to the polar pocket, which constrains the acyl chain in a neighboring hydrophobic pocket. Finally, the tilting orients PlaF in a con-monomer transition followed by tilting of the monomer in the membrane. The tilting orients PlaF in a configuration relative to the membrane such that substrates can directly access the cleft (Figure 1). In contrast, in the configuration observed in the crystal structure, the residues lining the opening of the active site cleft are more than 5 Å above the membrane surface (Figure 1). However, it is unknown how substrates reach the active site and how the characteristics of the active site determine the activity, specificity, and regioselectivity of PlaF for medium-chain GPLs.

The molecular mechanism underlying the access and binding of GPLs to PlaF is poorly understood in general because only a few PLA structures from microorganisms have been resolved, which either revealed closed conformations of their phospholipase domains or an accessible pocket that is predominantly hydrophobic or amphipathic. For the latter, regioselectivity was suggested to be achieved through binding of the GPL phosphate group to the polar pocket, which constrains the sn-1 acyl chain in a neighboring hydrophobic pocket. Finally, structural analysis of the outer membrane PLA (OMPLA) from *Escherichia coli* in the complex with an inhibitor provided information about GPL recognition by this PLA. However, OMPLA is an integral β-barrel protein with a hydrophobic GPL binding cleft and the active site located at the β-barrel exterior. Hence, the mechanism by which PlaF recognizes GPL substrates must be conceptually different from that of OMPLA.

Here, we combined unbiased and biased all-atom molecular dynamics (MD) simulations and configurational free-energy computations to identify access pathways to the catalytic center of PlaF. The results were validated by mutational and enzymatic studies on PlaF variants with blocked substrate access. Our results map out a distinct tunnel for substrate access within PlaF, provide explanations for the substrate specificity and PlaF activity of PlaF, and suggest egress routes for hydrolysis products. These findings enhance our understanding of the mechanism by which membrane protein function is regulated through protein–GPL interactions.

### 2. RESULTS

#### 2.1. Access Pathways to the Catalytic Site in PlaF

The crystal structure of PlaF revealed three pronounced tunnels, forming a large, T-shaped active site cleft. This cleft is compatible with binding bulky GPL substrates. However, the structural dynamics of biomolecules may lead to variations in the tunnel shape. Therefore, we reanalyzed trajectories from 10 replicas of unbiased MD simulations of 2 μs length for each of the systems di-PlaF (dimeric PlaF), PlaFα (chain A from the crystal structure), PlaFβ (chain B from the crystal structure), and t-PlaFα (chain A from the crystal structure in a tilted orientation) from our previous work using CAVER. CAVER analyzes and visualizes tunnels and channels in protein structures. We primarily focus on t-PlaFα because the tilted structure is likely in the catalytically active form. We identified the three tunnels that connect the active site of t-PlaFα to its surface like in the crystal structure (Figure 2): tunnel 1 (T1) and tunnel 2 (T2) point toward the membrane, and tunnel 3 (T3) opens to the periplasmic space.

![Figure 2. Clusters of tunnels identified in t-PlaFα ensembles. Three major tunnel clusters connect the catalytic site (black dashed circle) of PlaF to the protein surface. Tunnels T1 and T2 point toward the membrane; tunnel T3 is located >15 Å above the membrane, with its opening pointing into the periplasmic space.](https://doi.org/10.1021/acs.jcim.1c00973)

<table>
<thead>
<tr>
<th>Tunnel cluster</th>
<th>Occurrence</th>
<th>Maximum bottleneck radius</th>
<th>Average bottleneck radius</th>
<th>Average length</th>
</tr>
</thead>
<tbody>
<tr>
<td>T1</td>
<td>30.45</td>
<td>3.18</td>
<td>2.28</td>
<td>27.08</td>
</tr>
<tr>
<td>T2</td>
<td>21.80</td>
<td>2.95</td>
<td>2.21</td>
<td>23.75</td>
</tr>
<tr>
<td>T3</td>
<td>27.75</td>
<td>3.13</td>
<td>2.29</td>
<td>15.16</td>
</tr>
</tbody>
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Snapshots in which the tunnel is identified with respect to the total number of snapshots, in %. Data calculated with a probe radius of 2.0 Å in angstrom.

Between 2 and 5 Å depending on the location in the tunnel and the simulation length (Figure S1). The average bottleneck length is 2.26 ± 0.02 Å (mean ± standard error of the mean), which is close to the radius of glycerol (2.74 Å), an essential component of all GPLs, but smaller than the radius of 1,2-dilauroyl-sn-glycero-3-phosphoglycerol (DLPG) (∼4.4 Å) deduced from the lipid's...
area-per-lipid. For comparison, tunnels in monomeric PlaF, PlaFβ, and the two chains of di-PlaF show open occurrences of ~20 to ~5% (Table S1), indicating no marked differences between monomeric and di-PlaF.

To conclude, the active site of PlaF is connected to its surface with three tunnels. In the t-PlaFβ configuration, only T1 and T2 allow direct access of GPL or LGPL substrates from the membrane.

2.2. PlaF Preferentially Hydrolyses Medium-Acyt Chain LGPLs. Previously, we showed that PlaF can produce LGPLs by releasing FAs bound to the sn-1 position of GPLs. Here, we experimentally tested whether purified PlaF in vitro hydrolyses LGPLs by quantifying FAs released from a range of LGPLs varying in the head group and acyl chain length (C14–C18). The assays were performed by incubating purified PlaF (8 nmol) in n-dodecyl β-d-maltoside (DDM) micelles with the substrate, followed by quantification of the released FA by a non-esterified fatty acid (NEFA) assay. Lyso-phosphatidylethanolamine (LPE), lyso-phosphatidylglycerol (LPG), and lyso-phosphatidylcholine (LPC) contain FAs with 14–18 carbon atoms. The results revealed that PlaF can hydrolyze all tested LGPLs, with a preference for medium-acyl chain LGPLs (Figure 3). Interestingly, the lysoPLA activity of PlaF was 10- to 100-fold higher for all LGPLs than its PLA activity, indicating that hydrolysis of the first acyl chain in GPLs is much slower than that of the second one.

2.3. GPL and LGPL Substrate Extraction into Solvent and Acyl Chain Mobility. For probing the energetics of GPL and LGPL substrate extraction from the membrane into the solvent, we computed the free-energy profile for DLPG and 1-myristoyl-2-hydroxy-sn-glycero-3-phosphoglycerol (2LMG) extraction (Supporting Results), which resulted in free-energy differences between the two states of ~13 ± 0.1 and ~8 ± 0.3 kcal mol⁻¹ (Figure S2A), in very good agreement with the excess chemical potential related to these lipids’ critical micelle concentration. For access to T3, substrates would need to leave the membrane and pass through the water phase, which makes this route energetically unfavorable. Hence, T3 was not considered for further analyses.

As T1 and T2 are immersed in the hydrophilic membrane surface (Figure S3A), access of GPL and LGPL substrates to the tunnels via the head groups is plausible. However, the tunnels’ diameters are much smaller than that of a GPL-like DLPG while being in the membrane (see above). To explore the possibility that lipids access via their acyl chain instead, we probed how frequently the terminus of a GPL’s acyl chain can reach the membrane interface. The probability distribution of GPL’s acyl chains with respect to the coordinate perpendicular to the membrane (z-coordinate) was determined during the last 40 ns of 300 or 100 ns long MD simulations for membrane bilayers with or without t-PlaFβ, respectively (Figure S3A). Tails from both the upper and lower leaflet were considered. Positive z-coordinate values indicate that a tail moves toward the water–membrane interface of its leaflet; negative values indicate that it moves toward the interface of the opposite leaflet. The peak of the probability distributions is at z ≈ 2 Å, indicating the mobility of lipid termini within the leaflet (Figure S3A, see also Movies S1 and S2). The interface of the simulated membrane is at z ≈ 10 Å (Figure S3B). Notably, the cumulative probability of finding an acyl chain terminus at z > 10 Å is 1.5 and 1.0% for systems with or without PlaF, respectively. Hence, there is a finite likelihood that acyl chain termini can reach the entrances of T1 and T2. This result is also supported by the electron density profiles of the membrane components (Figure S3B).

To conclude, for t-PlaF, the access route of substrates to T3 is energetically unfavorable. In contrast, acyl chain termini of GPL lipids can reach the entrances of T1 and T2 during the time scales of our MD simulations.

2.4. Access Modes of GPL and LGPL Substrates into PlaF. As a prerequisite to computing the energetics of substrate access to the active site of PlaF, we aimed to identify favorable access modes. We applied steered MD (sMD) simulations to pull substrates inside T1 and T2 (Figure 4) via head access first or tail access first.

The closest substrate to the tunnel entrance was chosen for sMD simulations. The terminal oxygen and nitrogen atom of phosphatidylglycerol (PG) or phosphatidylethanolamine (PE) head groups, respectively, were considered for head access pulling. For tail access, the terminal carbons of respective acyl chains were considered. Substrates from the membrane were initially pulled through consecutive virtual points in T1 or T2 using four or five steps, respectively (Figure S4A, Table S2). However, pulling with terminal atoms leaves the cleavage site of the substrate distant to the catalytic S137 (Figure S4B). Therefore, the substrates were further pulled into T3, using three additional steps (Figure S4A). Depending on the access mode, the sn-1 or sn-2 sites of respective substrates were further pulled toward the nucleophilic OH group of the catalytic S137 (Table S2). Finally, this resulted in pulling pathways subdivided into eight and nine steps for T1 and T2, respectively (Table S2).

As a reaction coordinate, the distance between the pulled atom of a substrate and the consecutive virtual point was used. For each step, we repeated the pulling 50 times and computed the work done as a function of the reaction coordinate. By applying Jarzynski’s relation (eq 1), the work was related to the free-energy difference between the two states of the pulling simulation. The sMD trajectory whose work-versus-reaction coordinate profile is closest to the Jarzynski average (eq 1) was considered most favorable. Its endpoint provided the starting point for the sMD simulations in the next part of the pulling pathway. As a result, the access pathway is close to the lowest free-energy pathway of substrate access to the catalytic site.
Overall, this approach is the reversed version of sampling unbinding trajectories of ligands from proteins before applying Jarzynski’s relation \(^3\) but uses piecewise sMD simulations along the pathway to account for the curvilinear tunnels. A total of \(~27\) \(\mu\)s of sMD simulation time was used for either tunnel (Table S3).

The activity of PlaF for GPL decreases with the increasing lengths of the acyl chain between C12 and C18, irrespective of the type of head group, PG or PE.\(^\text{19}\) In addition, the number of acyl chains in a substrate also influences the PlaF activity, with LGPLs yielding a higher activity than GPLs.\(^\text{19}\) Hence, we chose DLPG with which PlaF is most active.\(^\text{19}\) 1,2-dilauroyl-sn-glycero-3-phosphoglycerol (DLPE), 1,2-distearoyl-sn-glycero-3-phosphoglycerol (DSPG), and 2LMG, a LGPL, for generating access modes (Figure 4A). Figure 4 exemplarily shows illustrations of the three access types for DLPG (see also Movies S3–S8). Work-versus-reaction coordinate profiles for all pulling simulations related to DLPG access are shown in Figure S5 for T1 and Figure S6 for T2. Based on computed potentials of mean force (PMFs) to evaluate the energetics of the access modes (see the next chapter), only tail access was considered for sMD simulations of the other GPL substrates (Figure S7). For 2LMG, head and tail access were considered for sMD simulations.

To conclude, seven access modes of GPL and two access modes of LGPL substrates into PlaF were generated for T1 and T2, resulting in 18 access modes in total.

2.5. PMFs of DLPG Access Modes. PMFs were computed from umbrella sampling (US) simulations\(^\text{3}\) and post-processed with WHAM\(^\text{3}^{5,6}\) to evaluate the energetics of substrate access for the access modes described in the previous chapter (Figure 4). As a reaction coordinate, the distance between the center of mass (COM) of the three oxygen atoms of the glycerol moiety in the substrate to the COM of C\(_{\alpha}\) atoms of the catalytic residues S137 and H286 was used. Residue D258 was not included in the reaction coordinate as its side chain is distant from the active site (Figure S4A). As the tunnels are almost straight, the reaction coordinate monotonically decreases as the substrate approaches the active site from the membrane (Figure S4B). Initially, we focused on the US simulations for the best PlaF substrate,\(^\text{19}\) DLPG. PMFs were calculated for the three access modes of DLPG across either tunnel, T1 or T2. The PMFs were evaluated for convergence, excluding the first 200 ns of 300 ns sampling time. The last 100 ns resulted in converged PMF profiles, yielding a maximal difference of \(~1\) kcal mol\(^{-1}\) in consecutive 20
ns increments of sampling time per window (Figure S8). The median overlap between the reaction coordinate distributions of neighboring windows was sufficient (≥4.8 and 3.5% for T1 and T2, respectively) (Figure S9).

The PMFs of DLPG access modes show marked differences (Figure 5A). Access with the head first is the least favorable for both T1 and T2, resulting in steep PMFs with free-energy barriers of 11 and 9 kcal mol⁻¹ (Figure 5A), in contrast to tail access. Most of the residues within a radius of 3 Å in T1 and T2 have either a neutral non-polar side chain, which likely facilitates tail access to the active site of PlaF. Furthermore, access with either one of the two tails first is more favorable in T2 than T1 (Figure 5A). Finally, access with tail 1 first in T2 is most favorable and results in no free-energy barrier until the substrate reaches the active site (Figure 5A). As the two acyl chains of DLPG are identical, these results suggest that their connection with the glycerol moiety causes differences in how the lipid interacts with the tunnel, which may explain how PlaF achieves regioselectivity to exert its PLA₁ function.

With enzyme kinetics results available only for DLPG, we used this substrate to validate our results. We computed the absolute binding free energy of DLPG to PlaF from the PMF for tail 1 access in T2, \( \Delta G_{\text{comp}}^\circ = -2.89 \pm 1.46 \text{ kcal mol}^{-1} \) (eq 4).

Assuming that product formation is slower than substrate dissociation from an enzyme, the Michaelis constant \( K_m \) is equal to the dissociation constant \( K_D \) of the enzyme–substrate complex.\(^{37,38}\) Under this assumption, from \( K_m = 7.612 \pm 1.907 \text{ mM for DLPG in PlaF} \),\(^{39}\) the experimental binding free energy \( \Delta G_{\text{exp}}^\circ = -3.07 \pm 0.30 \text{ kcal mol}^{-1} \) at \( T = 303 \text{ K} \) is calculated, which is within chemical accuracy\(^{40}\) of \( \Delta G_{\text{comp}}^\circ \). Note

**Figure 5.** PMF profiles for DLPG access. (A) PMFs of three access modes (head, tail 1, tail 2; see Figure 4B) of DLPG in T1 (blue curve) and T2 (yellow curve). For both tunnels, access with tail 1 first yields the lowest free-energy barriers to reach the active site. Furthermore, DLPG access into T2 with tail 1 first is overall the most favorable. The catalytic site is marked with a gray box. Insets within the plots illustrate the different DLPG access modes into the respective tunnels. (B) States during DLPG access via tail 1 through T2, as shown on the right, are marked in the PMF profile (left). The gray box corresponds to the integration limits used to calculate \( K_{eq} \) (eq 2) to determine \( \Delta G_{\text{comp}}^\circ \) (see the inset). State i: The starting position of DLPG (in the membrane). State ii: Tail 1 reaches the surface of the membrane close to the entrance of T2. State iii: Tail 1 enters inside T2, while tail 2 remains within the membrane. State iv: sn-1 site of tail 1 reaches the catalytic site.
that the computed binding free energy should not be confused with the free-energy barriers for substrate access from the membrane to the active site of PlaF mentioned later.

We also computed $\Delta G^\circ_{\text{comp}}$ for the other five access modes of DLPG (Eq 4). The lowest $\Delta G^\circ_{\text{comp}}$ among all six modes was obtained for tail 2 access in T1 (Table S4). However, the PMF profile (Figure 5A) reveals that the configurational free-energy minimum is not situated close to the active site but in the middle of T1. Here, one of the tails is still in the membrane, while the other is being loaded into the tunnel. If the PMF profile is integrated with two separate parts, first, a negative free energy for tail access into the tunnel results, followed by a positive free energy to reach the active site. This suggests that this access mode cannot yield a catalytically active configuration. For the other four access modes, $\Delta G^\circ_{\text{comp}} > 0.96$ kcal mol$^{-1}$ (Table S4). These findings corroborate tail 1 access of DLPG in T2 as the most likely access mode.

Along the PMF of tail 1 access of DLPG in T2, four distinct states can be identified (Figure 5B). The two tails of DLPG are immersed in the membrane at a reaction coordinate value of $\sim 38$ Å from the active site (Figure 5B, state i). The PMF remains essentially unchanged if tail 1 approaches the surface of the membrane and the entrance of T2 (state ii). This is concordant with the tail distributions along the $z$-coordinate during unbiased MD simulations (Figure S3), indicating that tail termini can reach one of the access tunnels of t-PlaFA without a considerable energetic cost. Once tail 1 enters T2, the PMF becomes negative (state iii), indicating that DLPG access that way is favorable. Finally, at $\sim 8$ Å of the reaction coordinate, the PMF has a global minimum (state iv). There, tail 1 is located in T3, and the acyl moiety at the sn-1 position of DLPG is close to the catalytic S137 of PlaF (Figure S10B) such that a nucleophilic attack can commence.

To conclude, we identified T2 as the preferred access tunnel for DLPG in PlaF. Access with tail 1 first is most favorable there. This is in line with PlaF being a PLA$_1$, which cleaves its substrates at the sn-1 position. As of T3, it is likely essential for substrate access by allowing to accommodate the substrate tail to be hydrolyzed by PlaF.

2.6. PMFs for Accesses of Other Substrates. Considering the results for DLPG, we performed US simulations for DSPG and DLPE only for tail 1 access. For the LGPL substrate, it has remained undetermined if the head or tail access is energetically favorable; hence, we performed US simulations for both access modes of 2LMG. As for DLPG, T2 is preferred over T1, regardless of the access modes (Figure 5A), we only considered T2 for computing PMFs for the other substrates. Similar to DLPG, the last 100 ns resulted in converged PMF profiles, yielding a maximal difference of $\sim 0.5$ kcal mol$^{-1}$ in consecutive 20 ns increments of the sampling time per window (Figure S11). Neighboring umbrella windows have a sufficient median overlap $\geq 3.2\%$ (Figure S12).

For DSPG and DLPE, access with tail 1 first in T2 results in pronounced free-energy barriers of 11 and 14 kcal mol$^{-1}$ (Figure 6A,B), in contrast to DLPG (0.5 kcal mol$^{-1}$). This finding indicates that a longer acyl chain or a neutral head group makes substrate access to PlaF disfavorable, which coincides with lower PlaF activities for such substrates. For 2LMG, access with the tail first is more favorable than that with the head, as for DLPG (Figure 6C,D). Furthermore, tail access by 2LMG leads to a free-energy barrier $\sim 6.5$ kcal mol$^{-1}$ lower than those for tail access by DSPG and DLPE (Figure 6A,B,D), which is concordant with the activity profile of PlaF.

To conclude, tail 1 access in T2 of GPL substrates with longer acyl chains or neutral head groups is disfavorable compared to DLPG access, in line with PlaF’s substrate specificity. For the...
LGPL substrate 2LMG, tail 1 access is also favored over head access and more favorable than DSPG and DLPE access.

2.7. Tryptophan Substitutions in T2 Hamper DLPG Access. To validate the prediction that T2 is the preferred access pathway, we identified residue positions in all identified tunnels that, when substituted with tryptophan (Trp), should constrict the tunnel and, thus, block substrate access. Earlier, this strategy has been used to block tunnels of a dehalogenase and influence its activity by limiting the rate of product release. In the case of PlaF, the products are less bulky than the substrates, such that product release should be less impacted than substrate access due to constricted tunnels.

PlaF variants were predicted subject to minimizing the structural destabilization due to the Trp substitution and preferring sites within the tunnels that influence its geometric characteristics (Table S5). We predicted four Trp substitutions for T1 and five for T2 and T3 each (Table S5). With any one of these substitutions in place, the impacted tunnel could not be identified anymore by CAVER applying the previously used probe radius of 2 Å but with a smaller probe radius of 1.2 Å (Figure 7). This indicates their constriction, also displayed by the time evolution of the tunnel profiles of the PlaF variants compared to PlaF wild-type (Figure S13).

The mutations of 14 suggested residues (Figure 7) to Trp were generated by the sequence- and ligation-independent cloning (SLIC) method, in which the whole p-PlaF expression vector was amplified. Mutations were verified by sequencing, and the wild-type PlaF (PlaFWT) and respective variants were produced in the homologous host, P. aeruginosa, following their immobilized metal affinity chromatographic purification from membranes solubilized with DDM (Figure S14). All variants showed purity comparable to that of PlaFWT (Figure S14). The specific activity of PlaF variants and PlaFWT was analyzed by measuring the hydrolysis of small (p-nitrophenyl butyrate, p-NPB) and large (DLPG) substrates (Figure 8A). The activities of all nine variants in T1 and T3 measured with p-NPB and DLPG were similar to the activity of PlaFWT. In contrast, all five T2 variants had a significantly (p < 0.001) lower activity with p-NPB and DLPG than PlaFWT.

To exclude that a substitution leads to an unstable protein, we measured the thermal stability of each variant by detecting intrinsic protein fluorescence upon unfolding. None of the variants showed a drastically reduced stability (Figure 8B). On the other hand, two T2 variants were slightly more stable (1.8–2.4 °C) than the PlaFWT, and three variants were slightly less stable (2.7–4.3 °C). Hence, Trp mutations do not affect PlaF’s stability at the temperature of enzymatic assays (30 °C).

The observation that PlaF activities with DLPG and p-NPB predominantly decreased with substitutions in T2 (Figure 8A) indicates that substitutions with the bulky Trp impact passage through T2 (Figure 8C). As expected, the activities with the larger DLPG decreased more (52–65%) than with the smaller p-NPB (15–30%).

We also determined kinetic parameters for the p-NPB hydrolysis of PlaFA221W and PlaFD74W with substitutions in T2 (Figure S15). Despite a less prominent effect of the substitutions on specific activities measured with p-NPB than DLPG, p-NPB allows for reliable determination of PlaF activities over the range of substrate concentrations and, thus, is applicable for the determination of kinetic parameters of PlaF. In contrast, enzyme kinetic experiments using hydrophobic DLPG are not feasible because of micelle formation and the slow rate of the reaction.

Although the affinities of PlaFWT, PlaFA221W, and PlaFD74W for p-NPB are similar (Figure S15, also see the table in the inset), the catalytic turnover of both variants (PlaFA221W: kcat = 314.6 ± 7.0 s−1; PlaFD74W: kcat = 403.4 ± 15.1 s−1) was significantly (p < 10−5) lower than that of PlaFWT (kcat = 487.8 ± 15.4 s−1) (Figure 8D). These results confirm that the point mutations PlaFA221W and PlaFD74W interfere with p-NPB access through T2.

To conclude, biochemical studies of 14 PlaF variants with Trp substitutions introduced in all three tunnels showed that only substitutions in T2 reduced lipolytic activity of PlaF. These results confirm that T2 is the main route for substrate access from the membrane to the catalytic site.

2.8. Potential Egress Pathways of PlaF Products. Next, we aimed at identifying potential egress pathways for products of PlaF-catalyzed hydrolysis. We performed a set of unbiased MD simulations starting from a hydrolyzed 2LMG in t-PlaF. The starting coordinates were taken from the last snapshot of the US simulations of 2LMG with tail 1 access through T2, considering the umbrella window where the sn-1 position of 2LMG was closest to the catalytic site. Then, 2LMG was cleaved into the respective products without changing their orientation in the tunnels (Figure 9A). This led to MYR being in T3 at the beginning of the simulations and the PGR (PG from LGPL, 2LMG) moiety pointing toward T2 (Figure 9B).

In 12 replicas of 3 μs length each, the products relocated within the tunnels, sometimes even diffusing into the solvent (PGR moiety in 2/12 replicas via T1 and 3/12 replicas via T2; Figure S16). MYR relocated from its original position in T3 and approached the other tunnels of PlaF during the course of the MD simulations (Figure 9C). To deduce the displacement of MYR, we measured the distance of the carboxyl carbon to the entrance of each tunnel. A cutoff of 5 Å, according to previous studies, was used to identify those replicas where MYR reaches close to the tunnel entrance. MYR moved in 7/12 replicas to the entrance of T1 and in 1/12 replicas to the entrance of T3; the entrance of T2 was not reached (Figure 9C). Interestingly, the instance of MYR reaching the entrance of T3 flips within T3 such that the carboxyl group points to the...
entrance, rather than to the active site as after hydrolysis (Figure 9B). While both MYR and PGR are able to get close to the entrance of T1, as seen for replica 11 in Figures 9 and S16, their movement is independent of each other, as evidenced by the lack of correlation between their distances (Figure S18). Altogether, MYR reaches the entrance of T1 significantly more frequently than T2 ($p = 0.0008$) and T3 ($p = 0.0047$) (Figure S17, Table S6).

To conclude, hydrolysis products of 2LMG diffuse within PlaF during time scales of 3 $\mu$s, sometimes also between tunnels.

Figure 8. Lipolytic activity of PlaF and variants with Trp substitutions in T1–T3. (A) Enzyme activities of purified PlaFWT and tryptophan variants of tunnel residues measured with DLPG and p-NPB. Only residues that were identified as part of tunnel 2 show a significant decrease in their activities, either with DLPG or p-NPB. Activities are normalized to the activity of PlaFWT, which was set as 100%. Results are means ± standard deviation of three independent measurements. Statistical analysis was performed using the t-test ($^*^p < 0.001$) of normally distributed values for DLPG ($n = 8$) and p-NPB ($n = 9$) measurements. (B) Thermal stabilities of purified PlaFWT and variants, measured by nanoDSF. Results are shown as a difference in the melting temperatures ($\Delta T_m$) of the respective PlaF variant and PlaFWT, which was $57.3 \pm 0.2$ °C. Results are means ± standard deviation of three independent measurements, each performed with three samples. (C) Results shown in panel A mapped onto the PlaF structure. The tunnels, T1–T3 (mesh view in the center), are represented as white surfaces. The investigated amino acids are shown in ball-and-stick representation. The CPK coloring scheme was used to color all atoms except the carbon atoms, which varies from pink to white (see the color scale) according to the impact on PlaF activity for DLPG of the tryptophan substitution in that position, showing a larger decrease for residues of T2. (D) $k_{cat}$ of PlaFWT and the substrate-binding-T2 variants PlaFD74W and PlaFA221W were measured using p-NPB and determined by non-linear regression analysis of the data ($n = 9$ for PlaFWT and PlaFA221W, $n = 6$ for PlaFD74W) fitted to the Michaelis–Menten equation. The lower turnover rate of the variants indicates a slower access to the catalytic site when T2 is blocked. The box plots represent the interquartile range between the first and third quartiles of the kinetic parameters determined for PlaFWT, PlaFDA74W, and PlaFA221W. The line inside the box is the median, and the whiskers represent the lowest and highest values. Statistical analysis was performed using the t-test ($^*^*^*^*^*^p < 10^{-5}$).
T1 and, to a lower extent, T3 are the most likely egress pathways of FAs from PlaF, although more sampling is required to observe actual egress.

3. DISCUSSION

Dimer−monomer transitions regulate the activity of several membrane-bound phospholipases, including PLA1 and phospholipase A2 (PLA2).45−52 Previously, we showed that PlaF becomes active due to a dimer-to-monomer transition followed by tilting of the monomer in the membrane, resulting in t-PlaFA being the active configuration of PlaF.19 Here, we addressed the questions of how membrane-bound substrates reach the active site of PlaFA and how the characteristics of the active site tunnels determine the activity, specificity, and regioselectivity of PlaF for medium-chain substrates. We performed unbiased and biased MD simulations and showed by configurational free-energy computations and mutational and enzymatic studies for t-PlaFA that (A) access of the two main PlaF substrates DLPG and 2LMG occurs most likely through tunnel T2 in a tail-first mode, (B) access of substrates with longer acyl chains or neutral head groups is less favorable, (C) tail 1 access of DLPG and 2LMG in T2 is more favorable than tail 2 access, (D) T3 accommodates the substrate tail to be hydrolyzed, and (E) T1 and T3 are potential product egress pathways.

Previous studies indicated that the characteristics of substrate access tunnels can have a decisive influence on enzyme−substrate specificity and activity.53−56 In t-PlaF, we focused on T1 and T2 because only these two allow direct access of GPL or LGPL substrates from the membrane in the t-PlaFA configuration. In contrast, to enter into T3, substrates would need to pass through the solvent, which is energetically unfavorable. In di-PlaF, T2 is closest to the membrane with a distance of 7.4 ± 1.5 Å, but T1 and T3 are at a distance >12 Å (Figure S19A). Hence, we also investigated substrate access to T2 in di-PlaF.

For assessing the energetics of substrate access, first, we generated 18 pathways, considering GPL and LGPL as substrates in T1 and T2 using sMD simulations. By relating the work along the reaction coordinate to the free-energy difference between two states of the pulling simulations via Jarzynski’s relation and considering the endpoint of the sMD trajectory closest to the Jarzynski average as the starting point for the next sMD simulation, we obtained low-free-energy pathways of substrate access to the catalytic site. sMD simulations have been widely used to explore similar biological processes such as the loading of GPL substrates into human PLA257 or recognition of arachidonic acid by cytochrome P450 2E1 across the access channel.58 The pathways served for defining reference points for subsequent US simulations, such that distributions of sampled states sufficiently overlapped, which is essential to yield accurate results in PMF computations.59 Applying US along pathways identified by sMD simulations60 or targeted simulations61,62 has been shown to be an effective method of computing PMF. Moreover, the choice of an appropriate reaction coordinate is essential for this approach.63−65 We probed for the convergence of our PMFs by comparing PMFs generated from increasing lengths of US simulations and found that US times of ∼300 ns are needed to yield PMF differences below chemical accuracy.66 Finally, we validated our PMF computations by comparing the computed absolute binding free energy of DLPG to PlaF for the most preferred access mode to an estimate of the experimental binding free energy.

Figure 9. Unbiased MD simulations of t-PlaF with bound hydrolysis products. (A) Starting configuration of the 2LMG products in t-PlaFα. MYR is represented with yellow spheres and PGR with green spheres. The catalytic S137 and H286 are shown as orange sticks. (B) Products are mapped over the respective tunnels. (C) Distance of MYR to the entrances of T1−T3 during 12 replicas of unbiased MD simulations of 3 μs. The dashed black line depicts the chosen cutoff of 5 Å, with replicas that reach this cutoff marked with an asterisk. MYR reaches a distance ≤5 Å to the entrance of T1 in seven replicas, in one replica for T3, and in none for T2. Note that in chain A of the PlaF crystal structure, MYR is found in T1.
The PMFs revealed that tail-first access through T2 is the most preferred method for DLPG and 2LMG. This finding is in line with the geometric analysis of T2, which revealed a tunnel bottleneck radius about half as large as the radius of DLPG deduced from the lipid's area-per-lipid, which can explain why a headgroup-first access is disfavorable for steric reasons. Furthermore, we showed that acyl chains of lipids embedded in a membrane can reach the interface region in unbiased MD simulations and, thus, can interact with the tunnel entrance. Such protrusions of lipid tails occur on a timescale of approximately 100 ns depending on the extent of solvent exposure. Tail-first access of GPLs into the active site has also been found for cyclopropane FA synthase. Tail-first access through T2 is favored because of the predominant hydrophobic nature of the tunnel walls. In contrast, T1 contains a higher number of charged Asp and Arg residues and fewer neutral residues than T2, which makes tail-first access there less favorable. In particular, the side chain of R80 protrudes into T1 at the tunnel entrance, which is reflected in an energy barrier of ∼3 kcal mol$^{-1}$ found there for tail-first access.

Modifications in tunnels that connect a buried active site to the bulk solvent have been shown to affect ligand binding and unbinding. Tunnel residues situated away from the active site are suitable targets for mutagenesis as their replacement should not lead to a loss of the functionality of the active site. Considering this, we introduced Trp substitutions to each of the three tunnels of PlaF and measured the activity of these PlaF variants. The Trp substitutions decreased PlaF's lipolytic activity for small and large substrates only when introduced in T2, which suggests that T2 is involved in substrate access. However, from such steady-state experiments, it cannot be excluded that the Trp substitutions influence product egress, too.

Among the investigated substrates, higher energy barriers for access to the active site were found for those with longer acyl chains and neutral head groups, concordant with PlaF's activity profile. This finding may be explained with differences in the energetics of GPL self-assembly, which is influenced by the hydrocarbon chain length and the polarity of the head group. Longer hydrocarbon chains and less polar head groups foster self-assembly, which would lead to higher energy barriers for leaving this equilibrium state and entering into PlaF. These results indicate that the energetics of access of a membrane GPL substrate to the active site through tunnel T2 contributes to the substrate specificity of PlaF.

Furthermore, of the two constitutopic acyl chains in DLPG, access via tail 1 in T2 is energetically preferred over tail 2 access. If tail 1 enters first, the carbonyl oxy group at C1 of the glycerol moiety can come closer to the nucleophilic S137 than if tail 2 enters first (Figure S10), leading to preferential hydrolysis of the carboxylic ester bond at C1. Likewise, the regioselectivity of human 5-lipoxygenase is determined by the head/tail-first type orientation of its main substrate arachidonic acid in the active site: the arachidonic acid can be positioned in the holoenzyme active site with both head-first and tail-first orientation, but only the tail-first orientation results in a configuration that yields S-lipoxygenating activity. These results indicate that the tail-first access mode of a diacyl GPL substrate determines the regioselectivity of PlaF for hydrolysis of the acyl chain bound to the sn-1 position.

As T3 is oriented to the membrane neither in the monomeric nor in the di-PlaF configuration, it likely does not contribute to substrate access. We suggest the role of T3 to accommodate the acyl chain of substrates before and products after hydrolysis. T3, with a length of ∼15 Å, provides adequate space for substrates with medium lengths of acyl chains and, thus, may affect the specificity of PlaF. Substrate tunnels that accommodate acyl chains hydrolyzed from their respective precursors have also been described for cholesterol acyltransferases. Likewise, lipid phosphate phosphatases harbor such a cavity, accommodating the substrate’s acyl chain for optimal catalysis. Site-directed mutagenesis in Candida rugosa lipase 1 revealed the role of such tunnels in determining the acyl chain length specificity.

As to di-PlaF, tail 1 access of DLPG across T2 revealed a free-energy barrier of ∼13 kcal mol$^{-1}$ (Figure S19B), in contrast to no free-energy barrier in t-PlaF (Figure S5B). This high barrier may arise because of the location of T2 in di-PlaF, ∼7 Å above the membrane. Thus, substrates would need to pass through the solvent to enter T2. These findings indicate that di-PlaF is catalytically inactive, as determined experimentally, because of energetically unfavorable substrate access.

Our results from unbiased MD simulations of products suggest that T1 and, to a lower extent, T3 are egress pathways of FAs. As to T1, this suggestion is in agreement with the crystal structure of PlaF, where FAs are found in T1. In the tilted orientation of PlaF, FAs egressing via T1 would interact with the membrane interface and could diffuse into it. FAs in a membrane can affect its fluidity and permeability and protein–lipid interactions, thereby regulating important cell processes including signal transduction, motility, and biofilm formation. Via T3, they would egress into the periplasmic space. Anchored to the cytoplasmic membrane, PlaF is not a toxin targeting the host cell membrane, but it has a direct influence on virulence adaptation of P. aeruginosa by modulating the membrane GPL composition. However, it is unknown whether FAs released from GPLs by PlaF are targeted to the external environment as, for example, diffusible FAs involved in cell-to-cell signaling. In this case, egress of FAs via T3 to the periplasm and their further passive diffusion or active transport would be possible.

In summary, we identified T2 as the preferred tunnel for substrate access to t-PlaF$_A$, while T1, and to a lesser extent T3, are likely egress routes for FAs. The energetically favorable tail 1 access of substrates is in agreement with PlaF’s PLA$_1$ function. The higher preference of PlaF for GPLs with medium-length acyl chains may be due to differences in the energetics of self-assembly and the length of T3, which accommodates them for hydrolysis (Figure 10). Finally, while t-PlaF$_A$ enables substrate access to the active site, substrate access to di-PlaF is energetically unfavorable. Our results provide an atomistic-level understanding of the unique structural feature of PlaF that its function is dependent on monomerization followed by global reorientation of the single-pass TM protein at the membrane. They may furthermore aid in understanding the feedback regulation of PlaF, which is inhibited by FAs, and open up opportunities for developing potential drugs that inhibit PlaF to combat P. aeruginosa virulence during infections.

4. MATERIALS AND METHODS

4.1. Identification of the Access Tunnels. Tunnels emerging from the active site of PlaF were identified using CAVER 3.0. The COM of the catalytic residues S137 and H286 was defined as the starting point of the search, from which the possible connections of the tunnels to the bulk solvent were identified. The catalytic residue D258 was not included in this search criteria since its side chain is distant from the catalytic site. Probe and shell radii of 2 and 6 Å were used, respectively.
investigated. For that, t-PlaFA was embedded into a membrane. Furthermore, loading of DSPG and an LGPL, 2LMG, was also prepared to compare and decipher the loading mechanism of DLPG or DLPE into t-PlaFA A. Considering the orientation and position of t-PlaFA in the membrane, one can safely assume that only substrates located in one leaflet will contact the catalytic domain of t-PlaFA and, hence, have direct access. Therefore, the composition of one leaflet was slightly modified to reflect the inclusion of the selected substrate. For this, a ratio in the upper leaflet of 6:2:1 for DLPE, DLPG, and the respective substrate was used. Using PACKMOL-Memgen, the bilayer system was prepared, solvated, and necessary counter ions were added. The minimum water distance from the membrane surface to the solvent box boundaries was increased to 35 Å to leave enough space between the substrate and the membrane surface and avoid interactions with periodic images during the extraction. Box dimensions in the x and y axes were set to 70 Å, resulting in systems comprising ~50,000 atoms.

3. Simulated Extraction of Substrates from the Membrane. MD simulations were performed using the GPU implementation of the AMBER 16 molecular simulation package,88,89 employing the ff14SB force field for the protein,90 the Lipid17 force field for the lipids,91–93 and the TIP3P water model.94 The SHAKE algorithm95 was used to constrain bond lengths of hydrogen atoms to heavy atoms, enabling a time step of 2 fs. Long-range electrostatic interactions were considered using the particle mesh Ewald algorithm.96 The system was energy-minimized by three mixed steepest descent/conjugate gradient calculations with a maximum of 20,000 steps each. First, the initial positions of the protein and membrane were restrained, followed by a calculation with restraints on the protein atoms only and finalizing with a minimization without restraints. The minimized system was then gradually thermalized in two stages. Initially, the temperature was increased from 0 to 100 K under NVT conditions, then from 100 to 300 K under NPT conditions at 1 bar, using a Langevin thermostat.97 The equilibration process continued for 5 ns, before starting with production simulations. As usual in membrane MD simulations, the NPT ensemble was used, allowing the membrane to accommodate along the trajectory.98 For US simulations, the pressure was maintained using an anisotropic Berendsen barostat,99 while for the rest of the simulations, a semi-isotropic Berendsen barostat was used, coupling the membrane (x-y) plane with the constant surface-tension dynamics. All analyses were performed using CPPTRAJ.100 Unless otherwise stated, molecular visualization was performed with PyMOL101 and visual molecular dynamics (VMD).102 The Movie Maker module within VMD was used to illustrate the acyl chain termini of lipids reaching the membrane interface and the access of substrates into PlaF.

To extract a substrate molecule from the membrane into one of the access tunnels, we selected the lipid that was closest to the entrance and pulled it from the membrane through the tunnel to the active site of PlaF, using constant velocity sMD simulations. Pulling simulations at low velocities are recommended for small polar molecules103 and large lipids104 to calculate free-energy profiles. At the lowest pulling rates, lipids have time to adopt energetically favorable conformations during the extraction process.104 In a recent study investigating GPL binding to PLaz, a constant pulling velocity of 5 Å ns⁻¹ was used.105 For the extraction of substrates, we considered all three possibilities by which a substrate may enter a tunnel: either the head group or one of the two tails. Depending on the type of head group (i.e., PG or PE), each substrate was pulled by its oxygen or nitrogen atoms at a constant velocity of 1 Å ns⁻¹ using a force constant of 5 kcal mol⁻¹ Å⁻². When pulling at the tail, the terminal carbon atom of the respective acyl chain was used.
Each tunnel was divided into several segments connected through virtual points formed by the COM of amino acids lining the respective tunnel. The number of virtual points depends on the length and shape of the respective tunnel. The virtual points guided the extraction of substrates such that the substrates followed the path of the respective tunnel. In addition, to obtain a low-energy pathway, an adaptive sMD protocol was implemented. For this, 50 replicas for each pulling simulation were carried out, and the work required was computed as a function of the reaction coordinate. The computed work was further related to free-energy difference between two states of the pulling simulation applying Jarzynski’s relation (eq 1).

$$e^{-W/kT} = e^{-\Delta F/kT}$$

Here, $\Delta F$ is the free-energy difference between two states, which is connected to work $W$ done on the system. $k$ is the Boltzmann constant, and $T$ is the temperature of the system. The replica closest to the Jarzynski’s average was considered to describe the lowest free-energy pathway and provided the starting point for the next pulling stage. Trajectories further away from that pathway were removed. This procedure results in faster convergence of PMF profiles, decreasing the overall computation needed.

For the systems without t-PlaF, the substrates were extracted with the same pulling velocity and spring constant, as mentioned above. However, to avoid edge effects, a substrate in the middle of the membrane was located. For this extraction process, the reaction coordinate was the distance between the head atom of the pulled substrate and COM of phosphorous atoms of the lipids in the opposite leaflet. Furthermore, to determine the free-energy minimum of the phospholipids in the membrane more accurately, the substrate was first pulled into the membrane (∼3 Å), before pulling it out of the membrane.

### 4.4. US and PMF Calculations

To understand the substrate access mechanism in PlaF and to identify preferential substrate access tunnels, PMFs were computed based on US, taking structures from the sMD simulations as starting points. As a reaction coordinate, the COM distance of the three oxygen atoms of the glycerol moiety in the substrate to the COM of residues S137 and H286 (only Cα atoms) of the active site was used. This reaction coordinate was also taken for all other systems for it describes the essential aspects of the structural transformation during substrate access. Consecutive positions of the substrates from the membrane to the active site as determined by pulling simulations were considered as reference points for US, with each position corresponded to one umbrella window. To achieve sufficient overlap between the umbrella windows, distances between reference points of ∼1 Å were used along the reaction coordinate. The length of individual tunnels and the size of acyl chains for respective substrates vary. Therefore, for sampling the access of different substrates, different numbers of windows were required for each tunnel. Selected positions of the lipid in the tunnel were restrained by harmonic potentials, using a force constant of 5 kcal mol$^{-1}$ Å$^{-2}$. To achieve sufficient convergence of the PMF profile, each window was sampled for 300 ns, of which the last 100 ns were used to calculate the PMF. Distance values were recorded every 2 ps and processed with WHAM. To estimate the PMF error, the data were separated into blocks according to the maximum calculated autocorrelation time of 20 ns. The correlation time was obtained for the complete trajectory, excluding the first 20 ns of sampling data for equilibration. The last 100 ns of sampling data were split into five blocks of 20 ns each, a PMF profile was calculated for each block with WHAM, and the error at each PMF point was calculated as the standard error of the mean.

Similarly, for systems without t-PlaF, trajectories obtained by pulling simulations were used to set up US simulations. Umbrella windows were extracted at distances of 1 Å from the starting point of the pulling simulation until the substrate was not interacting with the membrane anymore. The selected positions of the lipid were restrained by harmonic potentials, using a force constant of 5 kcal mol$^{-1}$ Å$^{-2}$ and as the reaction coordinate the distance of the COM of the three oxygen atoms of the glycerol moiety of the substrate to the COM of phosphorous atoms of the lower membrane leaflet. Each window was simulated for 100 ns at constant pressure (1 bar) and temperature (300 K) conditions until convergence was achieved. The first 20 ns of simulation data were discarded. WHAM was used to calculate the PMF. The PMFs were evaluated for convergence by checking the change in the free-energy profile with the increase in sampling time at every 10 ns. Furthermore, histograms of sampled configurations were visually inspected for sufficient overlap between the neighboring umbrella windows; otherwise, the iterative cycle in WHAM fails to converge, and the free-energy profiles have discontinuities.

### 4.5. Absolute Binding Free Energy from Computed PMF

The absolute binding free energy of substrates to PlaF was determined from the computed PMF using an approach modified from Chen and Kuyucak. The PMF was integrated along the reaction coordinate (eq 2) to calculate an association (equilibrium) constant ($K_\text{eq}$).

$$K_\text{eq} = \pi r^2 \int_{\text{active site}}^{\text{membrane}} e^{-W(\xi)/kT} d\xi$$

Here, $r$ is the maximum bottleneck radius of the respective tunnel, which was determined by CAVER analysis (Table 1), $\pi r^2$ is the cross-sectional area of the tunnel, $W(\xi)$ is the PMF at a specific value of the reaction coordinate, $k$ is the Boltzmann constant, and $T$ is the temperature at which the simulations were performed.

$K_\text{eq}$ was then transformed to the mole fraction scale ($K_\alpha$), taking into account the number of lipids ($N_\text{L}$) per membrane volume $V$ (eq 3).

$$K_\alpha = \frac{K_\text{eq} N_\text{L}}{V}$$

From $K_\alpha$ the difference in the free energy (eq 4) between the bound and unbound state ($\Delta G^\circ_{\text{comp}}$) of a single substrate molecule was calculated.

$$\Delta G^\circ_{\text{comp}} = -RT \ln(K_\alpha)$$

### 4.6. Blocking Access of the PlaF Substrates

To corroborate predicted access tunnels for PlaF substrates, we intended to block these by small-to-tryptophan substitutions of tunnel-lining residues. To do so, we identified possible substitution sites from our previous CAVER analyses, taking into account the tunnels’ bottleneck radii and lengths. For these analyses, the same trajectory used to search for tunnels in t-PlaF was considered. Finally, 4–5 amino acids within each tunnel were selected for substitutions.

In the first step, all the amino acids except glycines and prolines within 3 Å of individual tunnels and oriented toward a tunnel were considered. In turn, residues with an outward orientation were disregarded as a substitution there will likely...
not block the tunnel. Furthermore, as the TM and JM helices were found to be important for both the dimerization and the activity of PlaF, and residues of these helices were excluded. Finally, the catalytic residues S137, D258, and H286 and other residues of the active site were disregarded to avoid affecting the activity of PlaF.

The selected residues of each tunnel were substituted to tryptophan using FoldX, and the stability of the PlaF variants was evaluated in terms of the change in free energy (ΔΔG) with respect to the wild-type. Single amino acid substitutions were performed 10 times for each proposed residue of each tunnel, and the results were averaged. If the average ΔΔG > 3 kcal mol⁻¹, the substitution is considered destabilizing and was not further pursued. To check if the proposed substitutions will block the tunnel, the bottleneck radius of the variant tunnels was recalculated using CAVER. As done earlier, the probe radius was set to 2 Å. If no tunnel was identified with this criterium, the probe radius for tunnel search was reduced until the tunnels started to appear again.

4.7. Biological Evaluation of PlaF Activity from Mutations. 4.7.1. Site-Directed Mutagenesis, Protein Expression, and Purification. The plasmids for expression of PlaF variants with substitutions in the tunnels (Table S5) were created by PCR, using Phusion DNA polymerase (Thermo Scientific) in whole plasmid amplification, with mutagenic oligonucleotides (Table S7) designed for the SLIC method and p-plaF plasmid as a template. The presence of desired nucleotide substitutions was confirmed by DNA sequencing (MWG Biotech, Ebersberg, Germany). PlaF was purified from P. aeruginosa p-plaF membranes and solubilized with DDM, as described previously. Proteins were analyzed by polyacrylamide gel electrophoresis under denaturing conditions (sodium dodecyl sulfate-polyacrylamide gel electrophoresis) on 14% (w/v) gels, as described by Laemmli. The protein concentration was determined by measuring the A280nm using a NanoDrop 2000C spectrophotometer (Thermo Fisher Scientific Inc., Waltham, Massachusetts, USA). The extinction coefficients for PlaF and the variants were calculated with the ProtParam tool (Navia-Paldanius et al., 2012), considering the amino acid exchange and a His6-tag.

4.7.2. Enzyme Activity Assays and Kinetic Studies. The esterase activities of PlaF and variants were determined with p-NPB as substrate, as described previously, using a 96-well microplate and starting the reaction by adding 100 μL of the PlaF sample (16 nM) to the 100 μL of p-NPB solution (2 mM). Kinetic parameters (K_m and k_cat) for hydrolysis of p-NPB were determined using 8 nM enzyme, as described previously. Kinetic parameters were determined by non-linear regression analysis of data fitted to the Michaelis–Menten equation with PrismLab. Enzyme activities with DLPG and LGPLs were determined according to the established protocol in ref 113.

4.7.3. Thermal Stability. PlaF and variants (28.1 μM, 10 μL) loaded into the measuring capillaries (Prometheus NT.Plex nanoDSF Grade Standard Capillary Chips) were heated from 20 to 90 °C (1 °C min⁻¹ heating rate), and the intrinsic protein fluorescence was recorded at 330 and 350 nm using the Prometheus NT.Plex nanoDSF device (NanoTemper, Munich, Germany). The melting points were calculated from the first derivative of the ratio of F_<50,heat> and F_<50,room> using PR.ThermControl software (NanoTemper, Munich, Germany).

4.8. Egress of PlaF Products. To determine the egress pathways of PlaF products, a system with 2LMG substrate was considered. The final snapshot at 300 ns of the US simulations of the window with the substrate close to the active site was considered as the starting structure for unbiased MD simulations. 2LMG was cleaved into the products: MYR and PGR, without altering the orientation of each product within the tunnels. Atomic partial charges for the products were derived according to the restraint electrostatic potential fit procedure, as implemented in Antechamber. Geometry optimizations and subsequent single-point calculations were performed with Gaussian at the Hartree–Fock level with the 6-31G* basis set. Force field parameters for the products were taken from the general amber force field for organic molecules (GAFF, version 2). The prepared system was then minimized, thermalized, and equilibrated using the protocol described above for MD simulations. 12 replicas of production MD simulations of 3 μs length each under NPT conditions were performed. The distance of the 2LMG products to the entrance of each tunnel was computed for each replica.

DATA AND SOFTWARE AVAILABILITY
Compressed tar archived with configuration files, MD simulations data, and scripts to analyze such data are available at DOI: 10.25838/dsp-22. For molecular simulations, the AMBER16 package of molecular simulation codes was used. AMBER16 is available from http://ambermd.org/.

ASSOCIATED CONTENT
Supporting Information
The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.jcim.1c00973.

Energetics of GPL and LGPL extraction from the membrane; energetics of substrate access into T2 in dimeric PlaF; computational costs to determine PMFs related to substrate access; characteristics of tunnels identified in PlaF, PlaF_b, and di-PlaF using CAVER; pulling points across the tunnels for sMD simulations; overview of sMD simulations for the substrate access through T1 and T2 in t-PlaF; overview of computed absolute binding free energy of DLPG to t-PlaF from PMF; structural stability of proposed tunnel variants of PlaF; statistical test to determine the tendency of MYR reaching the entrance of tunnels T1−T3 in 12 replicas; list of the oligonucleotides used for generating expression plasmids; setup of US simulations for the substrate access through T1 and T2; properties of tunnels identified in t-PlaF_b ensembles; extraction of PlaF substrates from the membrane; distribution of acyl chain termini from GPLs and corresponding density profiles; substrate access pathway; work distributions obtained from 50 replicas of sMD simulations to pull DLPG across T1 and T2; convergence of PMFs for substrate access of DLPG; distribution of reaction coordinate values obtained by US for DLPG access; DLPG access to the catalytic site of t-PlaF; convergence of PMFs for other substrate access via T2; distribution of reaction coordinate values obtained by US for loading of other substrates across T2; heat map visualizing the time evolution of the tunnel profile for the proposed PlaF variants with tryptophan substitutions in
T1–T3; SDS-PAGE analysis of purified PlaF and variants; enzyme kinetics of the PlaF<sub>WT</sub> and the substrate-binding tunnel variants; unbiased MD simulations of hydrolysis products inside of t-PlaF<sub>A</sub> and within t-PlaF<sub>A</sub>; correlation of movements of lysophospholipid products; and substrate access in dimeric PlaF (PDF)

Acyl chain termini of lipids reaching the membrane interface in the presence of t-PlaF<sub>A</sub> during 40 ns long unbiased MD simulation (MPG)

Acyl chain termini of lipids reaching the membrane interface in the absence of t-PlaF<sub>A</sub> during 40 ns long unbiased MD simulation (MPG)

Extraction of a DLPG substrate molecule via the head from the membrane into T1 of t-PlaF<sub>A</sub> using sMD simulations (MPG)

Extraction of a DLPG substrate molecule via the tail 1 from the membrane into T1 of t-PlaF<sub>A</sub> using sMD simulations (MPG)

Extraction of a DLPG substrate molecule via the tail 2 from the membrane into T1 of t-PlaF<sub>A</sub> using sMD simulations (MPG)

Extraction of a DLPG substrate molecule via the head from the membrane into T2 of t-PlaF<sub>A</sub> using sMD simulations (MPG)

Extraction of a DLPG substrate molecule via the tail 1 from the membrane into T2 of t-PlaF<sub>A</sub> using sMD simulations (MPG)

Extraction of a DLPG substrate molecule via the tail 2 from the membrane into T2 of t-PlaF<sub>A</sub> using sMD simulations (MPG)

Author Contributions
H.G. conceptualization, supervision, analysis, visualization, and writing; F.K. conceptualization, supervision, analysis, visualization, and writing; K.-E.J. supervision and writing; S.A. investigation, analysis, visualization, and writing; C.H.S. investigation, analysis, visualization, and writing; and S.N.S.-V. analysis, visualization, and writing.

Notes
The authors declare no competing financial interest.

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