

Structural, Functional and Computational Studies of Membrane Recognition by Plasmodium Perforin-Like Proteins 1 and 2

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Abstract

Perforin-like proteins (PLPs) play key roles in mechanisms associated with parasitic disease caused by the apicomplexan parasites *Plasmodium* and *Toxoplasma*. The *T. gondii* PLP1 (TgPLP1) mediates tachyzoite egress from cells, while the five *Plasmodium* PLPs carry out various roles in the life cycle of the parasite and with respect to the molecular basis of disease. Here we focus on *Plasmodium vivax* PLP1 and PLP2 (PvPLP1 and PvPLP2) compared to TgPLP1. Determination of the crystal structure of the membrane-binding APC β domain of PvPLP1 reveals notable differences with TgPLP1, reflected in its inability to bind lipid bilayers as TgPLP1 and PvPLP2 do. Molecular dynamics simulations combined with site-directed mutagenesis and functional assays allow dissection of the binding interactions of TgPLP1 and PvPLP2 on lipid bilayers, and reveal similar tropisms for lipids enriched in the inner leaflet of the mammalian plasma membrane. In addition PvPLP2 displays a secondary synergistic interaction side-on from its principal bilayer interface. This study underlines the substantial differences between the biophysical properties of the APC β domains of apicomplexan PLPs, which reflect their significant sequence diversity. Such differences will be important factors in determining the cell targeting and membrane-binding activity of the different proteins in parasitic life cycles and disease.

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Introduction

Plasmodium, the parasitic causative agent of malaria, has a complex lifecycle involving two hosts, dominated by processes of migration through host tissue followed by cycles of intracellular replication, and *Plasmodium* Perforin-like proteins (PPLPs) have key roles in facilitating these processes.^{1–3} PPLPs are found across apicomplexan species, and contain a conserved central membrane attack complex/perforin (MACPF)

domain, as well as a unique C-terminal domain, named the Apicomplexan PLP C-terminal β -pleated sheet (APC β) domain,⁴ and a variable N-terminal domain.⁵ The MACPF superfamily represents an ancient homologous group of pore forming proteins, which function principally in cellular attack or defense. These proteins are secreted by a range of cells including bacterial and immune cells, bind to and oligomerize on cell membranes, and in this way form pores that typically result in target cell death.^{5–7} Key members of this family are perforin-

1 in cytotoxic lymphocytes,^{8,9} bactericidal perforin-2,^{10,11} the complement system itself,¹² and a range of toxins produced by organisms as disparate as stonefish¹³ and the oyster mushroom.¹⁴ Perforin family proteins may have a direct toxic effect (as complement does and also bacterial homologues) or be responsible for delivery of a secondary lethal agent (such as the delivery of granzymes by perforin-1).^{9,15}

The Apicomplexan PLPs (ApiPLPs), on the other hand, play important roles in cell traversal, invasion and egress by the parasites within their mammalian and insect hosts.^{1,16–19} The presence of MACPF domains in ApiPLPs suggests that membrane disruption by ApiPLPs is key to facilitation of these forms of cell behavior enabling infection and disease. *Plasmodium* species express 5 PLPs (PPLPs): two of them (PPLP1 and PPLP2) are involved in the mammalian host stage and the others (PPLP3–5) in the mosquito stage of the parasite life cycle. PPLP1 is implicated in cell traversal by sporozoites across the mammalian dermis and into the liver to invade hepatocytes and replicate within a parasitophorous vacuole (PV), as well as in the evasion of host phagocytic immune cells.^{1–3} Knockout of PPLP1 significantly disrupts parasite transmission in *P. berghei* and *P. falciparum*.^{2,20} PPLP1 also plays a role in erythrocyte egress by the merozoite stage (gametocyte) of the *Plasmodium* life cycle, via membrane disruption.²¹ PPLP2, too, has a function in gametocyte egress from erythrocytes for *P. berghei*²² and *P. falciparum*²³: exflagellation was disrupted in a PPLP2 knockout mutant, and was responsible for a subsequent inability to escape from erythrocytes. In contrast, PPLPs 3, 4 and 5 are all expressed during the parasite's ookinete stage and facilitate traversal across the mosquito midgut epithelium.^{16,17,24–26} Disruption of any of these three genes results in an inability by the ookinete to invade the mosquito midgut, suggesting distinct non-redundant roles for each of the three proteins.^{17,27,28} They may thus act as part of a single complex in a similar sense to the assembly of the membrane attack complex.¹²

Despite the significance of PPLPs in the malaria parasite life cycle, structural characterization has been lacking. A homologue from another member of the Apicomplexan family, *Toxoplasma gondii*, TgPLP1 facilitates parasite egress following intracellular replication.²⁹ X-ray crystallographic studies have previously revealed the structures of both the MACPF and APC β domains of TgPLP1.^{30,31} The TgPLP1 APC β crystal structures revealed the basic characteristics of the domain, which consists of three repeats of a β -sandwich fold arranged in a prism architecture.^{30,31} It binds to target membranes via an extended loop at its base^{30,31} as similar features found in proteins such as perforin-1, perforin-2 and the CDCs do too.^{11,32}

Here we present the first crystal structure of a *Plasmodium* PLP APC β domain from *Plasmodium*

vivax PLP1 at 3.15 Å resolution, and confirm a shared domain architecture between TgPLP1 and the PvPLP APC β domains. On the basis of these two structures we generate a homology model for the PvPLP2 APC β domain and use molecular dynamics (MD) simulations and lipid binding assays to characterize their membrane tropism. Our results reveal that PvPLP1 does not bind to any membranes tested while PvPLP2 shows a binding preference for lipids of the inner plasma membrane leaflet similar to that of TgPLP1. Free energy calculations and *in silico* mutational studies then identify a common mechanism of membrane anchoring between TgPLP1 and PvPLP2 APC β domains, clearly absent from PvPLP1. The variation in membrane interaction properties suggests possible functional and regulatory differences across the *Plasmodium* PLP family.

Results

Crystal structure of the PvPLP1 C-terminal APC β domain

The structure of PvPLP1 APC β domain was determined by X-ray crystallography at 3.15 Å resolution, to reveal as expected three β -stranded repeat motifs arranged in a prism (Figure 1(A); see Table 1 for data collection and refinement statistics). Each repeat consists of an inner four-stranded β -sheet and an outer two-stranded β -sheet, locked together by two disulfide bonds. Comparison with TgPLP1 APC β domain (root mean standard deviation, RMSD = 0.978 Å between 170 atom pairs) reflects a conserved C-terminal domain architecture across the *Toxoplasma* and *Plasmodium* APC β domains (Figure S1). Flexible loops between the inner and outer beta-sheets in the third repeat, which are absent in the crystal structure, were subsequently modelled in for molecular dynamics (MD) studies (see Materials and Methods below). Three 100 ns atomistic MD simulations of the PvPLP1 APC β domain structure indicated that the overall structural core was stable, at an average RMSD of 2.5 Å, while the modelled loops were more flexible (Figure S2).

The loops at the base of TgPLP1 APC β , especially the extended tryptophan-tipped hydrophobic loop (“dagger”) are important for anchoring the protein to the membrane.^{30,31} The overall sequence identity among *Plasmodium vivax* PLPs is very low, with an average of 20.2 % (Figure S3 and Figure S4). Residues in the membrane binding loop of TgPLP1 APC β ,^{30,31} although poorly conserved across all ApiPLPs, are significantly more conserved within each ApiPLP individually, indicating protein-specific functional importance (Figure S4(C)). Comparison of loops of APC β in PvPLP1 and TgPLP1 reveals that PvPLP1 lacks an extended loop and any aromatic residues at that point in the structure, containing instead several

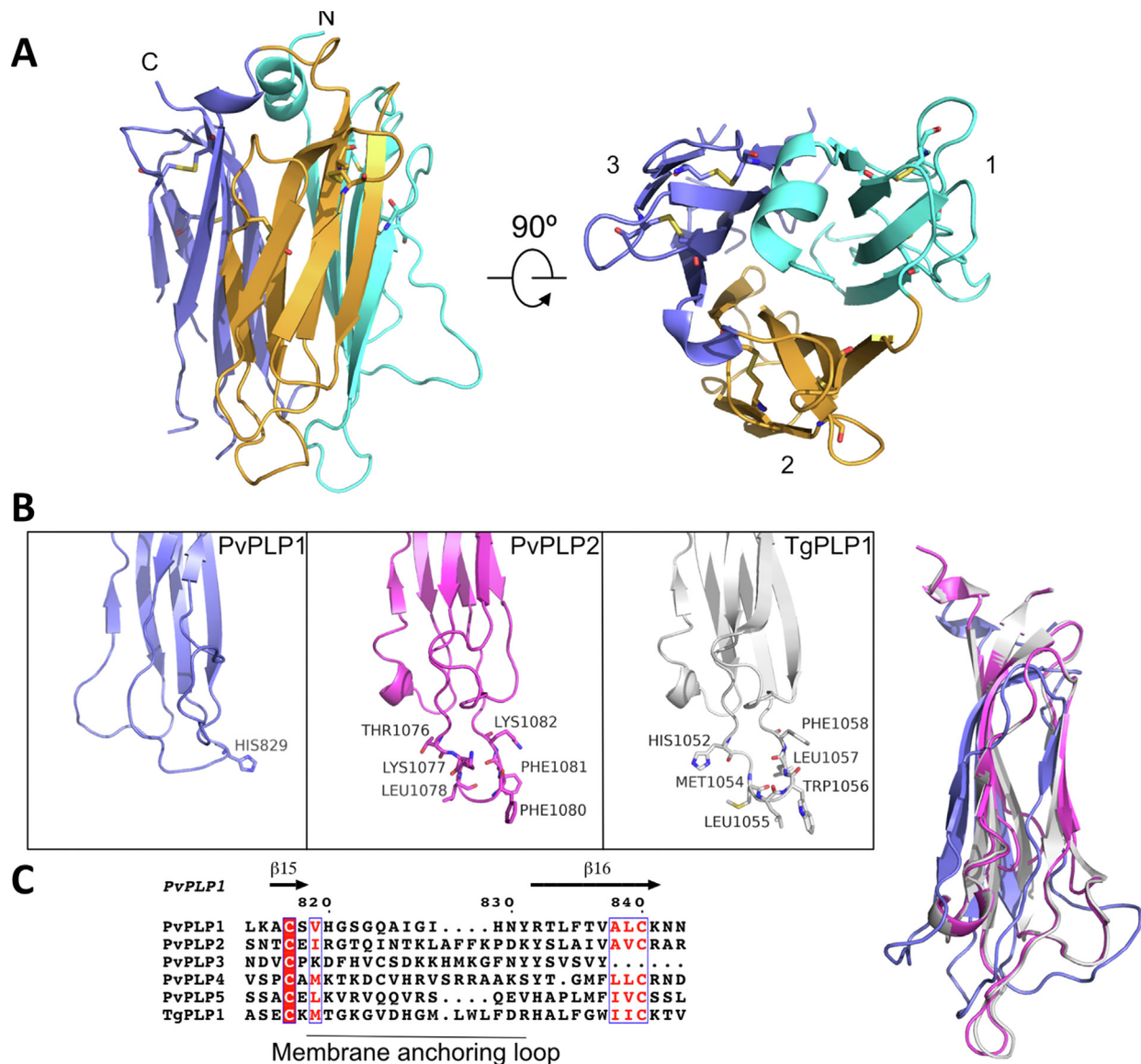


Figure 1. The crystal structure of PvPLP1 APC β domain and structural comparison with other ApiPLPs. (A) Front and top view of the 3.15 Å crystal structure of PvPLP1 APC β . The structure is composed of three repeats of a β -sandwich fold, each colored in cyan, orange and purple. Density for the turns at the base of repeat 3 was missing. (B) Boxes, comparison of the structures of the PvPLP1 and TgPLP1 APC β domain with the homology model of the PvPLP2 APC β , focusing on the third beta-sandwich module and highlighting hydrophobic and basic residues in the proposed membrane anchoring amphipathic loop. On the right is shown a superposition of the three structures, highlighting the agreement between the TgPLP1 and PvPLP2 models. (C) Sequence alignments of *Plasmodium vivax* PPLPs and TgPLP1 amphipathic loop sequences shows low sequence conservation at this region, and truncation of the third repeat in PvPLP3.

polar and aliphatic hydrophobic residues, as well as a histidine residue at its end rather than a tryptophan. In order to enable a side-by-side comparison of PvPLP1 and PvPLP2 we produced a homology model of PvPLP2 APC β using the TgPLP1 and PvPLP1 structures, facilitated by conservation of cysteine residues and hydrophobic core residues (Figure S1). The homologous membrane binding loop in PvPLP2 (¹⁰⁷⁷KLAFFK¹⁰⁸²) is longer than TgPLP1, but similarly tipped by two aromatic resi-

dues (Figure 1(B)). The hydrophobic and basic residues in the loop may contribute to membrane anchoring as it is amphipathic (later discussed as “loop 1”).

Modelling of PvPLP4 and PvPLP5 was possible on the same basis as PvPLP2, however calculation of sequence identity of each β -sandwich repeat separately found the third repeat to be the most variable, and its truncation in the PvPLP3 APC β (Figure 1(C) and Figure S3) meant

Table 1 Data collection and refinement statistics.

Wavelength	0.9686
Resolution range	63.01–3.15 (3.263–3.15)
Space group	P 1 21 1
Unit cell	73.74 42.02 87.22 90 104.95 90
Total reflections	122,538 (2207)
Unique reflections	9099 (790)
Multiplicity	13.5 (2.8)
Completeness (%)	98.40 (84.67)
Mean I/sigma(I)	11.32 (1.68)
Wilson B-factor	85.91
R-merge	0.1942 (0.5668)
R-meas	0.201 (0.6824)
R-pim	0.04997 (0.371)
CC1/2	0.995 (0.695)
CC*	0.999 (0.906)
Reflections used in refinement	9098 (790)
Reflections used for R-free	431 (37)
R-work	0.2134 (0.3271)
R-free	0.2715 (0.4303)
CC(work)	0.936 (0.742)
CC(free)	0.909 (0.015)
Number of non-hydrogen atoms	3684
Protein residues	483
RMS(bonds)	0.012
RMS(angles)	1.44
Ramachandran favored (%)	86.12
Ramachandran allowed (%)	13.23
Ramachandran outliers (%)	0.65
Rotamer outliers (%)	0.47
Clashscore	16.73
Average B-factor	84.11

Statistics for the highest-resolution shell are shown in parentheses.

we could not confidently generate a model of its structure. The PvPLP4 and PvPLP5 APC β models indicated an absence of the key hydrophobic loop compared to those of PvPLP1 and PvPLP2 (Figure S5(B)). We nevertheless hypothesise that the function of the APC β domain is conserved across all PLPs as a membrane binding module, allowing for adaptation to the diverse range of cellular environments within which *Plasmodium* PLPs function, and that their sequence and structural variation in this region are likely to relate to different binding preferences (Figure 1(B)). Also, the possible formation of a membrane-targeting complex by PvPLPs 3–5 means that they all do not need to have membrane recognition or binding properties, as is also the case with the complement membrane attack complex component proteins.¹²

PvPLP1 does not bind to membranes, while PvPLP2 binds in two distinct orientations

The membrane binding capacity of PvPLP1 and PvPLP2 APC β domains was probed firstly by lipid dot-blot assays. PvPLP1 did not bind to any mammalian membrane lipid species tested while PvPLP2 recognized anionic lipid species, with a strong preference for phosphatidylinositol 4-

phosphate (PI(4)P), followed by palmitoyloleoylphosphatidylserine (PS) and Cardiolipin, and a weak association with the other phosphatidylinositol phosphates (PIPs) (Figure 2). The weak association with other PIPs is somewhat surprising, but suggests that the bulkier substitutions presented by di- and triphosphates to the inositol ring may have a sterically blocking effect.

These results were confirmed by liposome sedimentation assays. Affinity of PvPLP1 and 2 for membranes containing Cholesterol, Sphingomyelin (SM), Cardiolipin, PS, L- α -phosphatidic acid (Egg-PA), L- α -lysophosphatidyl serine (Lyso-PS) and brain and liver total lipid extract were tested. PvPLP1 showed no binding to any membranes, while PvPLP2 bound to membranes containing negatively-charged lipid species like Cardiolipin, PS and Egg-PA (Figure 2 (B and C)).

Membrane association simulations were next performed to further probe APC β membrane binding characteristics. Although such studies were restricted to relatively simple model membrane systems with simulations run for microsecond timescales, they provide valuable insights into the principles governing biological function, allowing for other limitations including

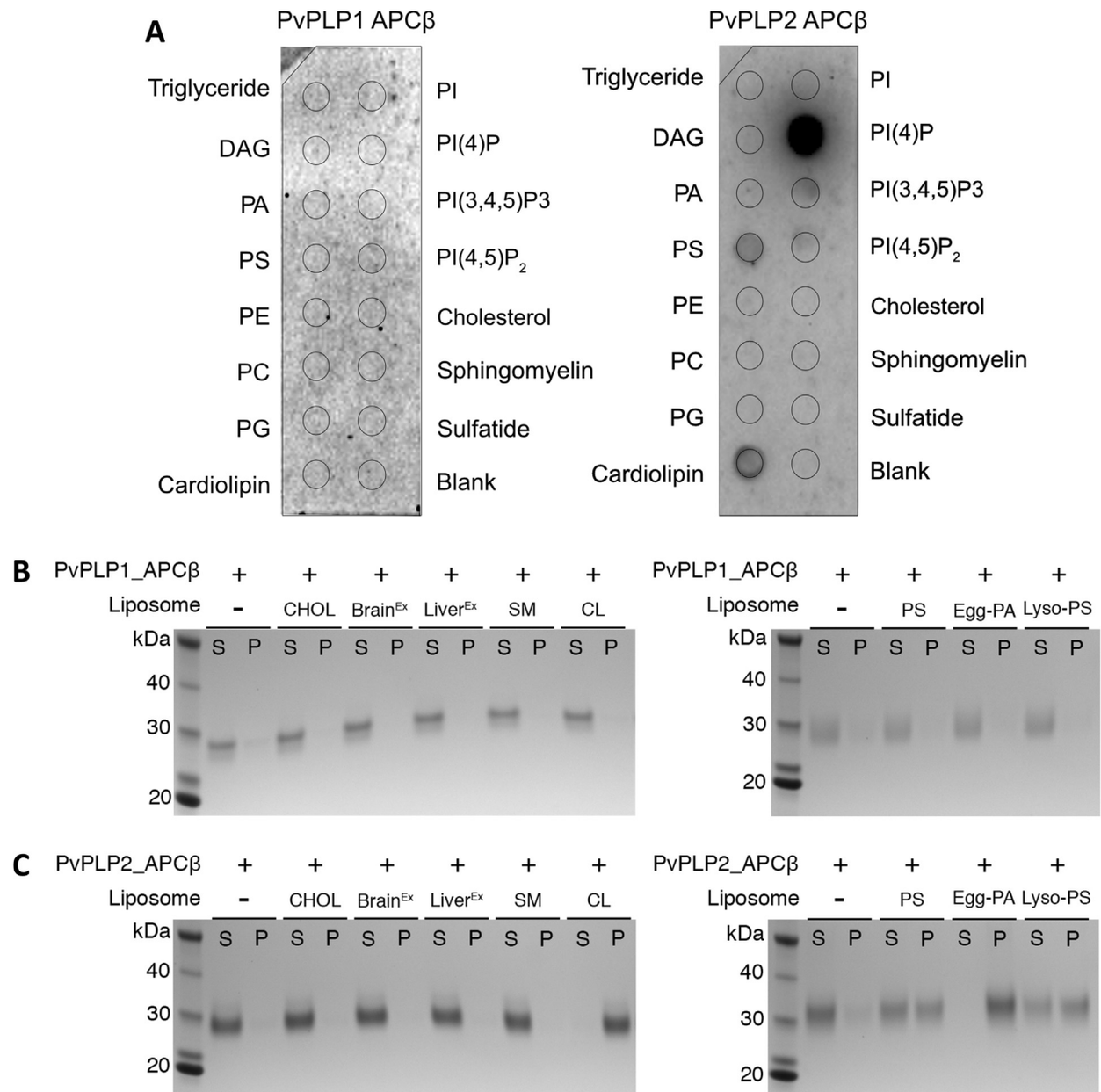


Figure 2. PvPLP1 and PvPLP2 APC β show different membrane binding capacities. (A) Representative lipid dot-blots. APC β s were incubated with commercially available nitrocellulose membrane dotted with common mammalian membrane lipid species. (B–C) Liposome sedimentation assays of APC β domains of PvPLP1 (B) and PvPLP2 (C). All liposomes are composed of 40% phosphatidylcholine (PC), 30% phosphatidylethanolamine (PE) and 30% of varied lipids. CHOL, Cholesterol; Brain^{Ex}, Brain Total Lipid Extract; Liver^{Ex}, Liver Total Lipid Extract; SM, Sphingomyelin; CL, cardiolipin; PS, phosphatidylserine; Egg-PA, L- α -phosphatidic acid; Lyso-PS, L- α -lyso-phosphatidylserine.

choice of force field and parametrizations of some specific molecular types and interactions, and finite system scale.³³ In addition, the use of a combination of atomistic and coarse-grained simulations can make longer timescales accessible without necessarily compromising on resolution.^{33–35} Table 2 provides a summary of all the simulations performed for this work. As can be seen,

the model membranes used here do represent very simple models and did not include cholesterol.

PvPLP1 and PvPLP2 APC β s were positioned above a bilayer containing lipids palmitoylcholine (PC), palmitoylcholine (PE) and PS (45:45:10) representative in simple terms of the eukaryotic membrane inner leaflet, and allowed freely to associate with the membrane over 2 μ s in

Table 2 Summary of the simulations performed in this work.

Protein	Lipids	Composition (%)	No. Simulations/length (ns)	Simulation Type (CG/AT)	
PvPLP1	–	–	3/100	AT	
	PC:PE:PS	45:45:10	20/2000	Unbiased (CG)	
	PC:PE:PS	45:45:10	25/1000	Umbrella Sampling (CG)	
PvPLP2	PC:PE:PS	45:45:10	20/2000	Unbiased (CG)	
	PC	100	25/1000	Umbrella Sampling (CG)	
	PC:PE	80:20	25/1000	Umbrella Sampling (CG)	
	PC:PS	80:20	25/1000	Umbrella Sampling (CG)	
	Upright orientation	PC:PE:PS	45:45:10	25/1000	Umbrella Sampling (CG)
	Side orientation	PC:PE:PS	45:45:10	25/1000	Umbrella Sampling (CG)
		PC:PE:LPA	45:45:10	25/1000	Umbrella Sampling (CG)
		PC:PE:PS:PI(4)P	45:42:10:3	25/1000	Umbrella Sampling (CG)
		PC:PE:PS	45:45:10	3/100	Unbiased, upright orientation (AT)
		PC:PE:PS	45:45:10	3/100	Unbiased, side orientation (AT)
		PC:PE:PS	45:45:10	25/1000	Umbrella Sampling (CG)
TgPLP1	PC	100	25/1000	Umbrella Sampling (CG)	
	PC:PE	80:20	25/1000	Umbrella Sampling (CG)	
	PC:PS	80:20	25/1000	Umbrella Sampling (CG)	
	PC:PE:PS	45:45:10	25/1000	Umbrella Sampling (CG)	
		PC:PE:PS:PI(3,4,5)P	45:42:10:3	25/1000	Umbrella Sampling (CG)
	W ¹⁰⁵⁶ A	PC:PE:PS:PIP3	45:42:10:3	25/1000	Umbrella Sampling (CG)

coarse grained (CG) Martini simulations. Sets of 20 simulations for each protein were performed. Only two binding events were observed in PvPLP1 simulations (Figure 3 and Figure S6). In one case,

transient binding lasted for 500 ns, where PvPLP1 bound in a tilted orientation facilitated by association with the long loop between $\beta 11$ and $\beta 12$ in the third repeat. In the other case, PvPLP1

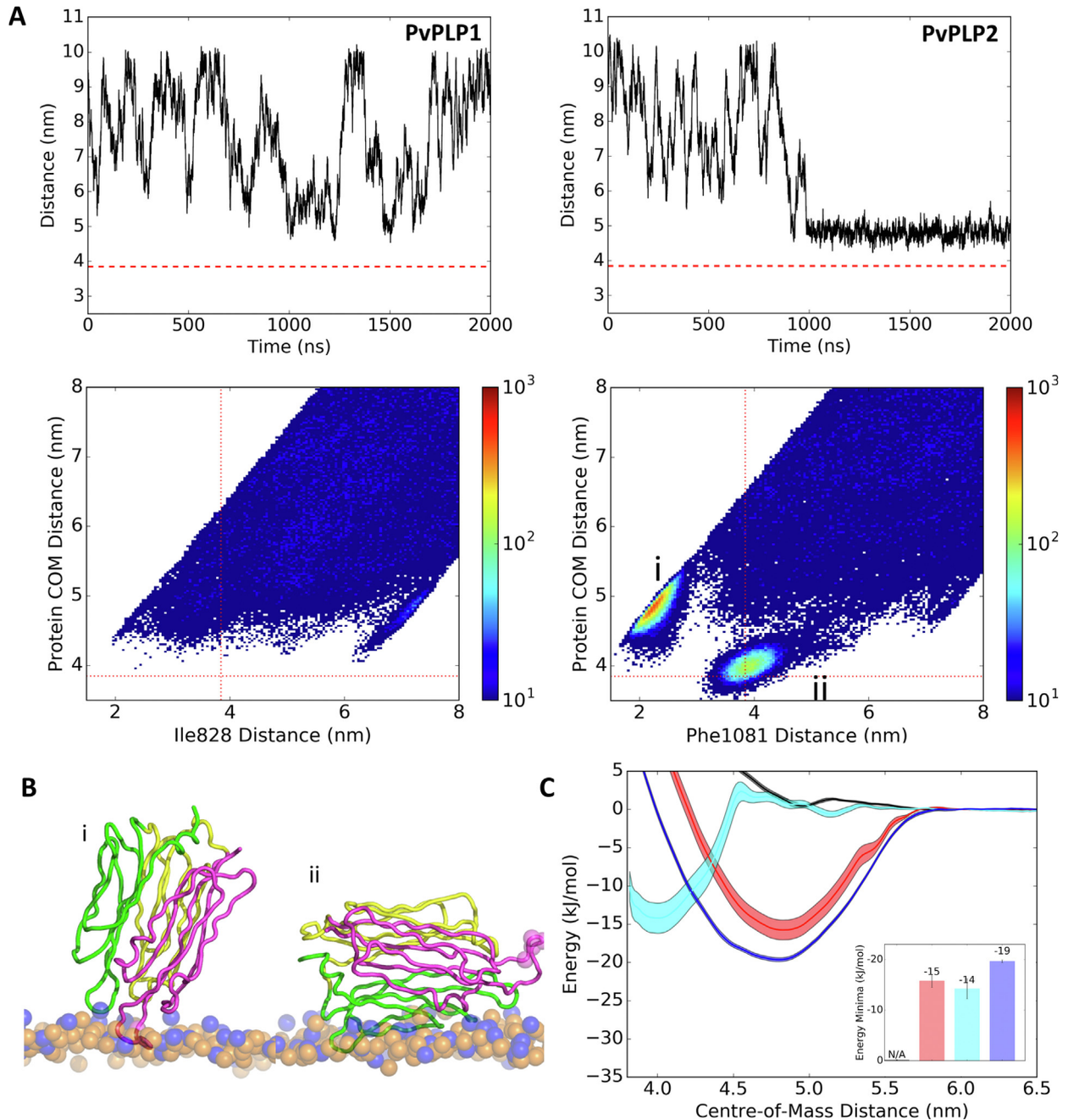


Figure 3. PvPLP1 does not bind to PC:PE:PS (45:45:10) bilayers in simulations while PvPLP2 reveals two distinct binding poses. (A) Top panels: Representative time-distance plots of PLP membrane association simulations. Distance was calculated between the center of mass (COM) of the protein and bilayer. The dashed red line indicates lipid head phosphate distance from COM of membrane. Bottom panels: Density plots indicating COM distance and selected tip residue of the amphipathic loop to bilayer COM distance. Plots show cumulative data of all 20 repeats of each simulation type. (B) Representations of two binding poses identified for PvPLP2 APC β from simulations. (C) PMF profile for PvPLP1 (black), PvPLP2 upright orientation (red), PvPLP2 side orientation (cyan) and TgPLP1 (blue) APC β domains bound to PC:PE:PS membranes. Distance is measured between COM of the protein and bilayer. Error estimates were obtained using bootstrap analysis. Computed free energy values at the energy minima for each profile are inset.

bound in an upside-down orientation, with the top corner of the domain associated with the bilayer. Given that the MACPF domain would sit above the domain at this location due to the short linker between the two domains, this orientation was deemed non-physiological, as in a previous study with the isolated TgPLP1 APC β domain.³¹ In contrast, PvPLP2 bound to the bilayer in 11 out of 20 (55%) of simulations, in two different orientations, both seemingly of physiological relevance (Figure 3 (B)). In the primary binding orientation (73% of binding events), PvPLP2 sits upright on the bilayer with the phenylalanine-tipped loop at its base anchoring it within the bilayer. This orientation is analogous to that seen in previous studies of TgPLP1.³¹ In the second orientation PvPLP2 lies along the membrane surface with its longitudinal axis parallel to the bilayer (27% of binding events). Visual inspection and lipid contact analysis indicates that for this pose the first tandem repeat of the APC β domain fold consistently lies along the face of the bilayer, suggesting a specific interaction of this repeat (Figure S6(B)). Although the starting states for these simulations make some form of interaction with the membrane very likely, the marked difference in behavior between PvPLP1 and PvPLP2 suggests a significant difference in binding affinity.

Potential of mean force calculation for PvPLPs APC β binding to membrane

As unbiased MD simulations suggested preference for PvPLP2 binding to the membrane when compared with PvPLP1, we calculated the free energy of membrane binding by PvPLP1 and PvPLP2. For comparison, the same procedure was also performed with TgPLP1. The free energy of binding to PC:PE:PS (45:45:10) bilayers was calculated using orientations taken from the membrane association simulations. Energies were calculated using umbrella sampling simulation methods to measure the potential of mean force (PMF) for binding events. In this approach, each membrane binding domain is placed at a series of positions along a 1-dimensional pathway defining an approach to the membrane long which the PMF will be calculated, and the reaction coordinate path is split into windows each of which is sampled in a separate simulation³⁶ (Figure 4 (A)). As PvPLP1 did not bind to the membrane in the orientations defined by PvPLP2 and TgPLP1, the protein was aligned with TgPLP1 on the membrane to generate a comparable binding orientation. Similar binding energies of -15 and -19 kJ mol⁻¹ were obtained for PvPLP2 and TgPLP1 respectively, while the absence of an energy well for PvPLP1 indicates it has no affinity for the PC:PE:PS (45:45:10) bilayer (Figure 3(C)). The fact that PMF calculations showed similar free energy of binding of PvPLP2 in a sideways orientation to that of the upright primary binding pose again indicates that they are both of biological relevance, though

the greater occupancy of the upright state suggests it is the ultimate and preferred orientation, especially since it is based on a significantly smaller interaction surface area than the sideways orientation.

Lipid preferences of PvPLP2 by PMF calculations

To explore the lipid preference of PLPs further, we measured free energy of binding on bilayers of different lipid compositions. PMF calculations were first performed with 100% PC, and PC:PE and PC:PS in 80:20 ratio. This was then complemented with two negatively charged lipids, in bilayers of PC:PE:PS (as above) and PC:PE:Egg-PA (molar ratio 45:45:10). PvPLP2 had a low affinity for PC bilayers alone, with addition of PE and PS increasing free energy of binding by -4 and -3 kJ mol⁻¹ respectively (Figure 4(B)). Addition of both PS and PE complement each other to increase binding energy by a further -5 kJ mol⁻¹. PvPLP2 had a greater affinity for Egg-PA (LPA) over PS, increasing binding energy by -12 kJ mol⁻¹ over PC:PE alone. Accordingly, PC, PE and PS and Egg-PA all contribute to binding of PvPLP2, and complementation with both PE and a negatively charged lipid improved binding affinity. Umbrella sampling simulations with a PC:PE:PS:PI(4)P bilayer (molar ratio 45:42:10:3) were also performed. Free energy of binding of PvPLP2 to a PI(4)P containing bilayer was -41 kJ mol⁻¹, a substantial increase in energy of -26 kJ mol⁻¹ over the PC:PE:PS bilayer consistent with the lipid dot blot result (Figure 2(A)).

In comparison, TgPLP1 had a similar increase in binding energy with the addition of either PE and PS of -8 and -7 kJ mol⁻¹, respectively (Figure 4(C)). However, presence of both PS and PE only increased binding energy by ~ -3 kJ mol⁻¹ for TgPLP1. Our previous work found that TgPLP1 showed a preference for phosphatidylinositol 3,4,5-triphosphate (PI(3,4,5)P₃) and phosphatidylinositol 4,5-bisphosphate (PI(4,5)P₂) containing bilayers.³¹ Consistently, PMF calculations with a PC:PE:PS:PI(3,4,5)P₃ bilayer (molar ratio 45:42:10:3) showed an increase in affinity of -10 kJ mol⁻¹ over a PC:PE:PS bilayer.

Taken together, these results indicate that PvPLP2 and TgPLP1 have an affinity for negatively charged lipids that generally reside in cytoplasmic leaflets. In contrast, PvPLP1 does not bind to membranes under conditions tested here. The significantly increased affinity of -26 kJ mol⁻¹ of PvPLP2 to PI(4)P, compared to -10 kJ mol⁻¹ for TgPLP1 on PI(3,4,5)P₃ containing bilayers suggests that electrostatic interactions may play a more important role in PvPLP2 membrane binding than in TgPLP1.³¹

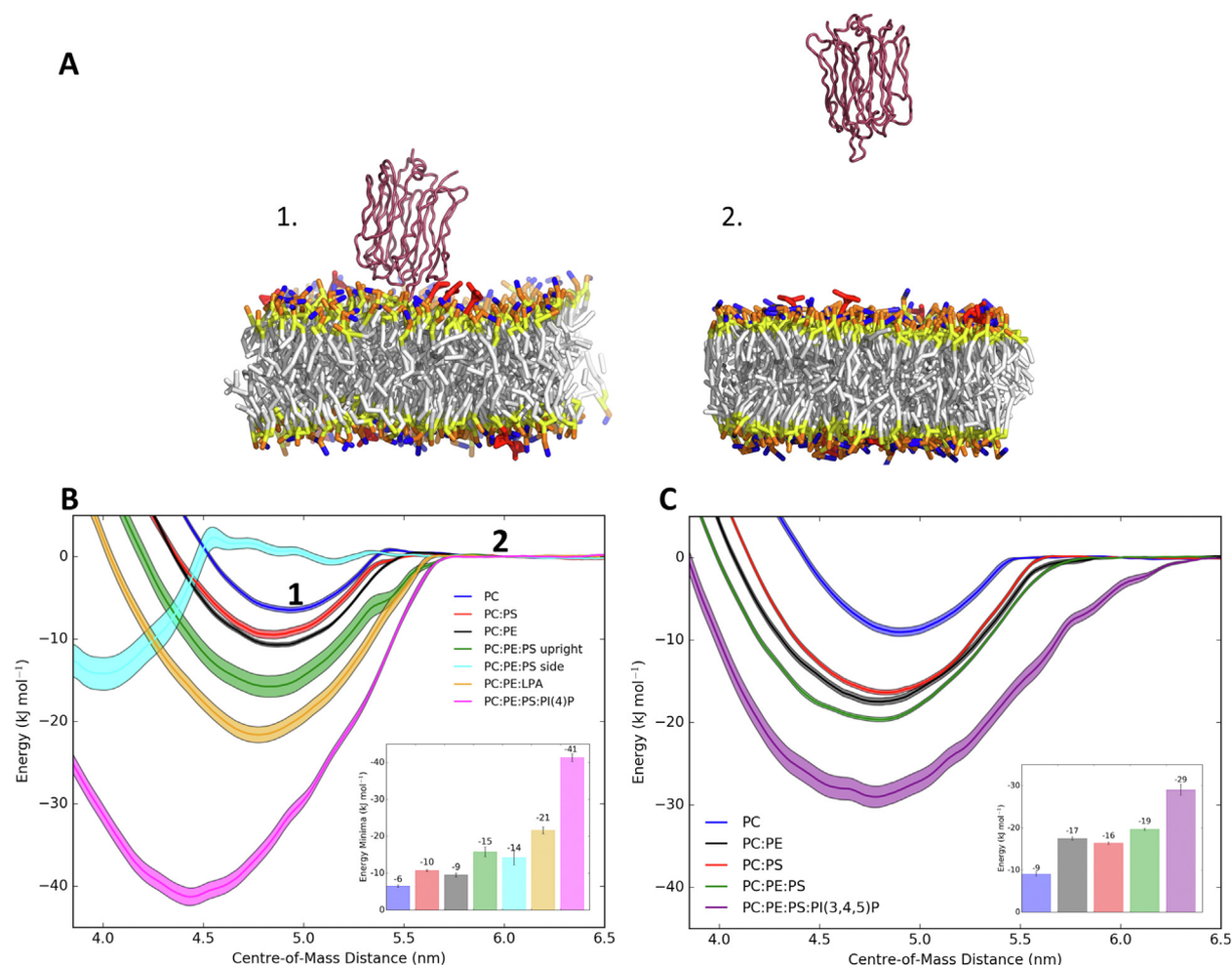


Figure 4. PvPLP2 lipid preferences and mutations within the membrane binding loop. (A) Cartoon representations of PvPLP2 APC β in the upright orientation bound to the PC:PE:PS (45:45:10) bilayer. Steered MD was used to pull PvPLP2 away from the initial position (1) on the bilayer into solvent (2) to generate umbrella sampling windows. (B) PMF profiles of PvPLP2 APC β bound to several membranes of different lipid compositions. Positions (1) and (2) are marked. PC:PE and PC:PS were in 80:20 ratios, PC:PE:PS and PC:PE:LPA (Egg-PA) in 45:45:10 ratios, and PC:PE:PS:PI(4)P in the ratio 45:42:10:3. (C) PMF profiles of TgPLP1 APC β on several bilayers of different lipid composition: PC:PE and PC:PS were 80:20, PC:PE:PS 45:45:10 and PC:PE:PS:PI(3,4,5)P 45:42:10:3.

Aromatic, hydrophobic and basic residues contribute to membrane binding of PvPLP2

To investigate the protein-lipid interactions of PvPLP2 in greater detail, the two orientations were simulated further in full atomistic simulations. Representative systems were taken from CG simulations and converted to atomistic resolution (see Materials and Methods), following which three repeats of 100-ns duration were produced for each orientation. For each orientation the protein had similar overall stability (Figure S7) and remained bound to the membrane. Over the course of one repeat of the upright orientation, PLP2 began to tilt sideways and transit to the sideways orientation. The other two repeats maintained their initial upright position. Simulations in the sideways orientation remained in a

consistent orientation throughout the 100 ns. Again, this perhaps relates to the greater buried surface in this binding orientation. The short 100 ns timescale means that our insights with atomistic simulations provide however restricted insights into binding dynamics and principally support the stability of orientations identified from CG analysis.

Lipid contact analysis following atomistic simulations reaffirmed that the phenylalanine tipped loop (loop 1) anchored the protein into the bilayer. In this orientation, a secondary lysine-rich loop (⁸⁷⁸KNK⁸⁸⁰, “loop 2”) further stabilized the membrane interactions (Figure 5(A)). A series of hydrophobic residues protruding from the surface of repeat 1 of the APC β domain also had high contact with the bilayer in the sideways orientation, with several lysines and an asparagine

residue also contributing to the membrane binding (Figure 5(B)). We termed a concentrated subset of these residues ($^{903}\text{FGKK}^{906}$) “loop 3”.

To investigate the energetic contribution of these residues to membrane binding, a series of *in silico* mutations were then produced. The equivalent *in silico* mutation of $^{1077}\text{KLAFFK}^{1082} \rightarrow \text{AAAAAA}$ (loop 1) entirely abolished membrane affinity in the upright orientation (Figure 6(A)). This result also occurred when a double mutant of F1080A: F1081A was tested, while mutation of both lysines, K1077 and K1082, in the same loop only minimally reduced binding by 2 kJ mol^{-1} over the wild-type protein. F1080 was identified in lipid contact analysis as the primary phenylalanine responsible for membrane anchoring. Point mutation of this residue to alanine substantially reduced computed membrane affinity to -4 kJ mol^{-1} . Atomistic simulations also identified that K880 on loop 2 was frequently in contact with the membrane. K880A showed only a minimal reduction in binding energy. Although the double loop mutant shows a deeper energy well of -17 kJ mol^{-1} , this is unrelated to binding to the membrane surface – it does not show the relaxation to 0 kJ mol^{-1} that the wild type protein does and its energetic profile is, like that of K880A and F1080A, truncated. A possible explanation of the energy minimum is the hydrophobic effect as

this mutant form of PvPLP2 APC β replaces 6/7 polar and charged residues with seven alanines (one alanine substitutes a phenylalanine) with consequent effects via water on the free energy of the system. As the tip tryptophan in the extended membrane binding loop of TgPLP1 was identified as a key residue in membrane binding, analogous to F1081 in PvPLP2, a W1056A mutant was compared with PvPLP2 mutants. W1056A reduced the free energy of binding to a PC:PE:PS:PIP3 bilayer by 15 kJ mol^{-1} (Figure 6(B)), consistent with liposome sedimentation assays which show mutations in this extended loop reduced membrane binding.^{30,31}

Mutation of aromatic residues in membrane anchoring loops reduces membrane binding

Following identification, via MD, of the membrane binding role of hydrophobic and positively charged residues at the base of the protein, we produced two full-length PvPLP2 mutants targeting these regions. In the first mutant, the primary membrane binding loop (“loop 1”) was mutated to alanine ($^{1077}\text{KLAFFK}^{1082} \rightarrow \text{AAAAAA}$). The second mutant targets the positively charged residues in loop 2 and 3 at the base of PvPLP2, with $^{878}\text{KNK}^{880}$ (loop 2) and $^{903}\text{FGKK}^{906}$ (loop 3) mutated to alanines. Liposome sedimentation assays with these mutants showed a similar

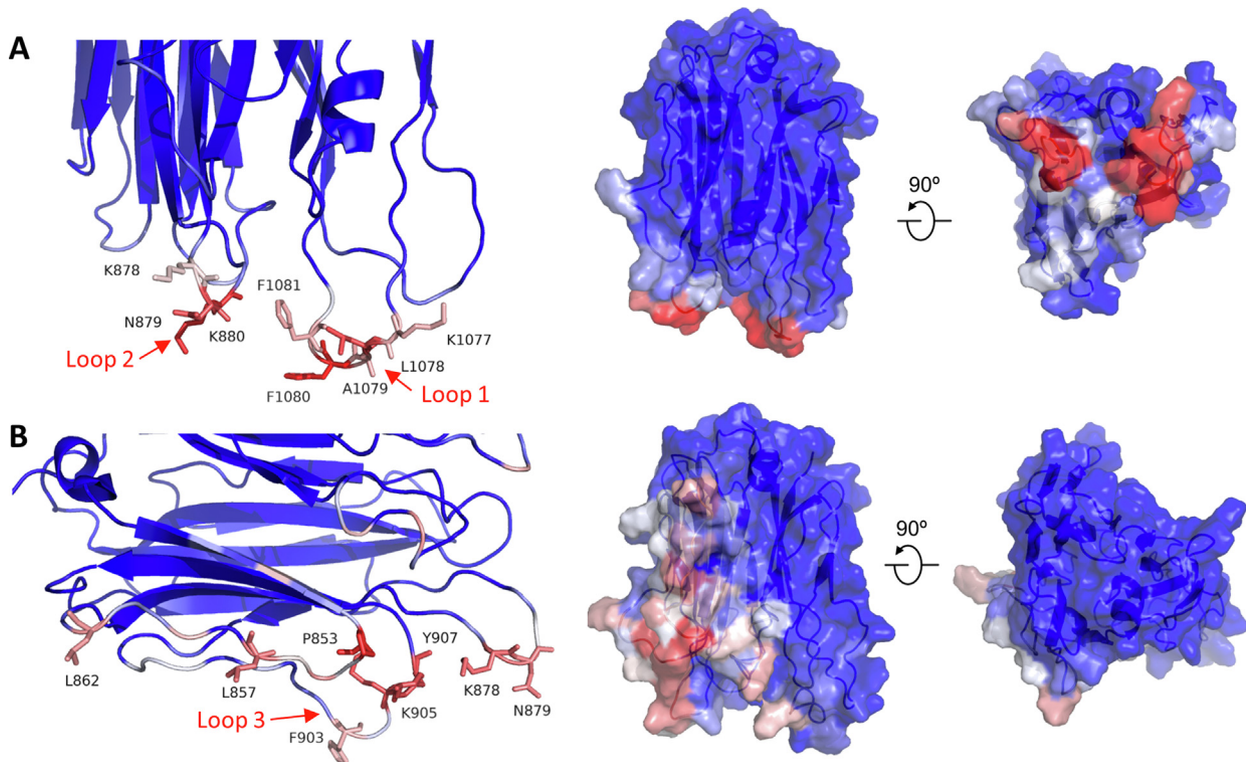


Figure 5. Protein-lipid contacts for 2 orientations of PvPLP2. Cartoon (left) and surface representations (right) of PvPLP2 APC β , colored by normalised number of lipid contacts averaged over three 100 ns atomistic simulations. A: upright orientation, B: side orientation.

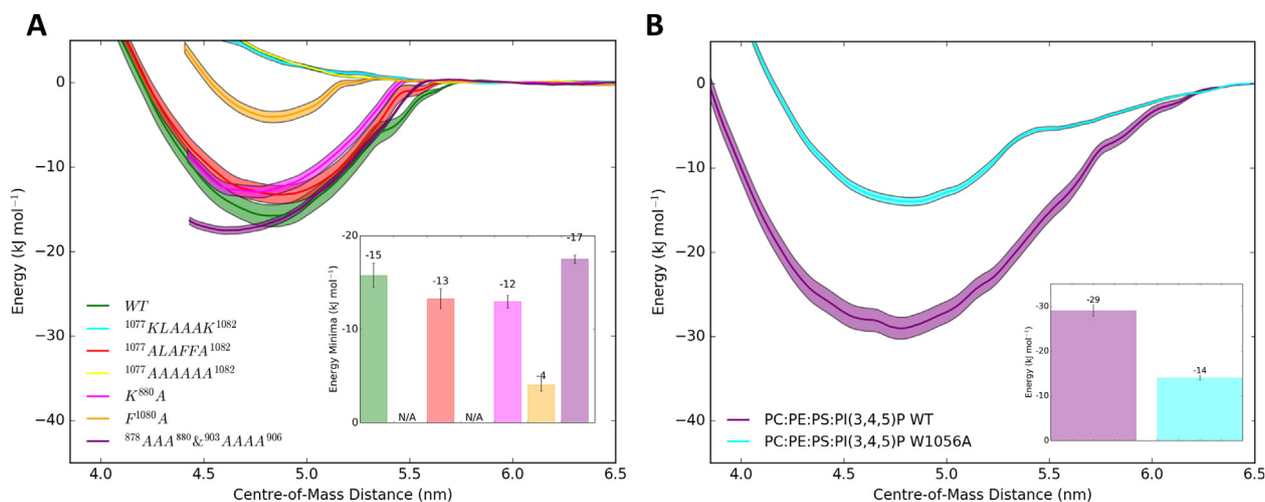


Figure 6. Lipid binding preferences and tryptophan mutation of TgPLP1. (A) PMF profiles for PvPLP2 APC β wildtype and membrane binding loop mutants FF \rightarrow AA and KK \rightarrow AA on bilayers composed of PC, PE and PS (45:45:10). Bootstrap analysis was used to generate error estimates. (B) PMF profiles for TgPLP1 APC β wildtype and W1056A mutation on a PC:PE:PS:PI(3,4,5)P₃ (45:42:10:3) bilayer. Bootstrap analysis was used to generate error estimates.

effect: the binding of the PvPLP2 to three different forms of liposomes was substantially reduced compared to that of the wild-type proteins (Figure 7).

Discussion

PPLPs disrupt membranes during several stages of the *Plasmodium* lifecycle, acting in both cell traversal and egress. Given the diversity of membrane compositions at different stages, PPLPs might have evolved with specific membrane binding characteristics at each stage. Here we show the first crystal structure of a *Plasmodium* PLP C-terminal APC β domain, that of PvPLP2 which we find to be similar to the TgPLP1 APC β domain. The similarity in structure, conservation of disulfide locking cysteines and hydrophobic core residues across the PPLP family further confirm a shared domain structure across the ApiPLPs. Combined with MD simulation studies as well as lipid dot blots and liposome sedimentation assays, we show that PvPLP2 APC β binds to membranes, while PvPLP1 APC β had little affinity for the membrane lipids. PvPLP1 may rely on a different method for membrane anchoring, possibly via an unidentified glycolipid or a protein receptor, or requiring an unidentified mechanism of activation. This is supported by the distinctly different amino acid composition of the PvPLP1 basal long loop, responsible for primary interaction of PvPLP2 and TgPLP1 with membranes. Similar to PvPLP1, PvPLP4 and 5 lack a phenylalanine/tryptophan containing loop (Figure S5), which indicates that they too may

bind to membranes with a different mechanism to PvPLP2 and TgPLP1.

Despite low sequence identity, we showed that PvPLP2 APC β bound to membranes in a similar mechanism to TgPLP1: an extended loop at the base of the domain tipped with an aromatic residue facilitates primary membrane anchoring for both proteins³¹ (Figures 4–7). PvPLP2 had affinity for PE, PS and PI(4)P, which are enriched in the inner leaflet of the plasma membrane. We observed similar affinity towards inner leaflet lipids for TgPLP1, which confirms previous experimental observations.^{30,31} However, PvPLP2 contains additional positively-charged loops (loop 2 and loop 3), which stabilize the primary binding through charge-charge interactions (Figure 7). Consistently, PMF calculations revealed a marked increase in binding energy of PvPLP2 in PI(4)P containing membranes compared to the more moderate increase in TgPLP1 binding upon addition of PI(3,4,5)P₃ (Figure 4). Mutation of these residues substantially decreased the binding efficiency to negatively-charged membranes (Figures 6 and 7). These findings suggest that anionic lipids may play a more important role in PPLP2 membrane binding, whereas TgPLP1 may have a broader binding profile with less specificity. This could reflect the more confined role of gametocyte egress in which PvPLP2 functions, with PvPLP2 targeting the inner plasma membrane of the cell, compared to the broader replicative niche in which *Toxoplasma* operates.

Interestingly, besides the primary upright membrane anchoring orientation via an extended phenylalanine tipped loop, PvPLP2 also bound to the membrane on its side, an orientation that has

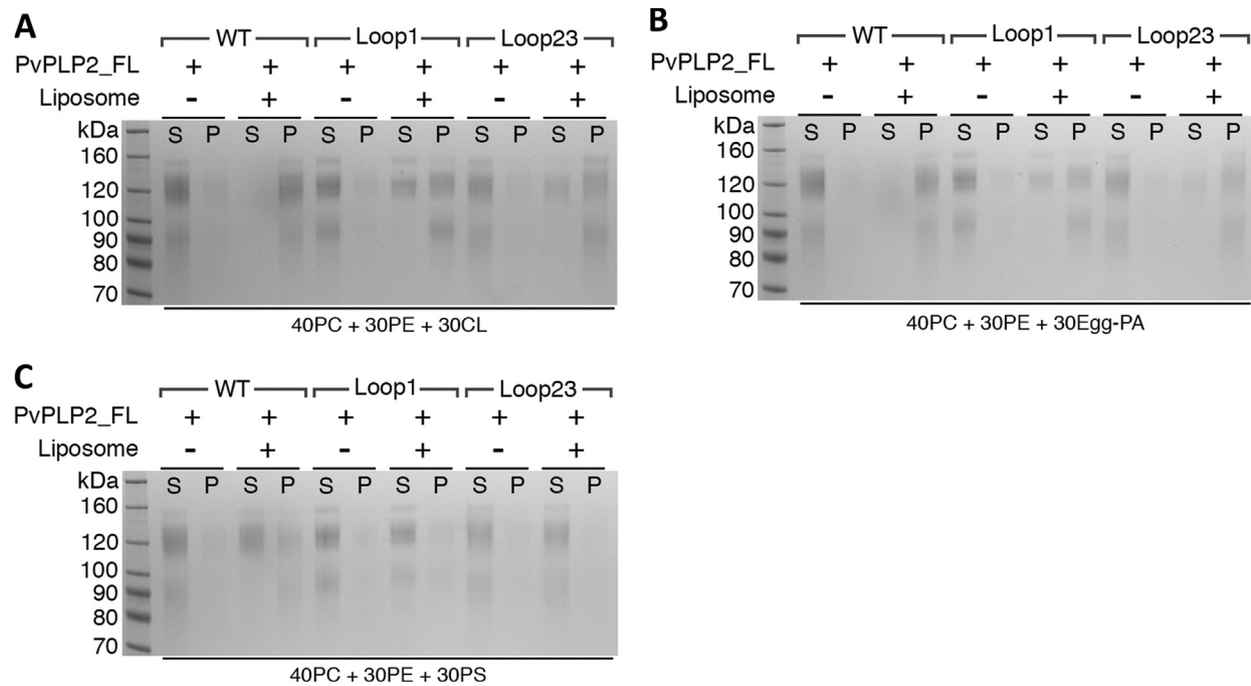


Figure 7. PvPLP2 mutant liposome binding assays. (A–C) Ultracentrifugation-based liposome-binding assays showing that mutations in the hydrophobic tip and charged residues within the APC β domain (Loop1 and Loop2/3 mutants) reduce the binding of PvPLP2 full-length protein to Cardiolipin (A); L- α -phosphatidic acid (B) and phosphatidylserine (C), indicating the role of APC β domain for membrane binding of PvPLP2. “Loop1” represents the mutant that affects the Phenylalanine loop tip (1077 KLAFFK 1082 – AAAAAA); “Loop2/3” indicates the mutant gives charge neutralization (878 KNK 880 – AAA & 903 FGKK 906 – AAAA).

not been observed for other PPLPs. PMF calculations showed that PvPLP2 had a similar affinity for the membrane in both orientations, though the more upright state appeared to be the preferred pose. Additionally, loop 2 (878 KNK 880) formed a contact point for both orientations and might be able to act as a pivot point between them. Given the preference for the upright pose, despite its smaller buried surface area, we interpret the side-on binding event as a synergy interaction increasing the chance of membrane binding.

This paper provides the first study of the molecular basis of *Plasmodium* PLP APC β membrane binding. We have revealed structural and functional similarities between members of the ApiPLP family, while also identifying key differences in membrane binding between *Plasmodium* PLP1 and PLP2. PPLPs function in multiple different environments, cell types and in both processes of cell traversal and egress, throughout the *Plasmodium* lifecycle. The differences identified here indicate that membrane binding may be an important regulatory step for PPLPs and varies across the protein family. Regulation of ApiPLP membrane binding is likely to be important for apicomplexans to prevent off-target membrane disruption and damage,

especially during motile stages which require microneme secretion without membrane damage, such as cell invasion. Specificity for negatively charged inner leaflet lipids, in addition to the role of pH, may prevent binding to the outer surface of the plasma membrane during cell invasion. Extensive further study will be needed to elucidate the different membrane binding functions and mechanisms of action of PPLPs during both cell egress and traversal. As ookinete midgut traversal is a bottleneck in transmission of malaria it will be important to further understand the molecular determinants of function of PvPLPs 3–5.

Materials and Methods

Construct cloning and site-directed mutagenesis

In this study, both PvPLP1 (PlasmoDB: PVX_000810) and PvPLP2 sequence (PlasmoDB: PVX_123515) were codon-optimized (by ThermoFisher, GeneArt) for a mammalian expression system. The APC β domains of *PvPLP1* (residues 584 to 843) and *PvPLP2* (residues 836 to 1104) and the full length of *PvPLP2* (residues 25–1104) were cloned into the pHLsec vector³⁷ using restriction sites (AgeI/KpnI), appending a signal peptide in the N-terminus and a

KTHHHHHH tag in the C-terminus of the protein. During protein secretion into the media, the signal peptide was cleaved, leaving an additional three residues Glu-Thr-Gly at the protein N-terminus. Site-directed mutagenesis of the full-length PvPLP2 (loop1 and loop2&3 mutants) was conducted by overlapping polymerase chain reaction (PCR).

Protein expression and purification

All the PvPLP1 and PvPLP2 proteins were produced recombinantly from mammalian HEK293T cells using a transient transfection and expression protocol as described.^{30,31,38} To express the APC β domains of PvPLP1 and 2, the plasmid DNA (12 mg) was transfected into 6 liters of HEK293T cells in the presence of the glycosylation inhibitor kifunensine (final concentration of 5 mM) with 12 mg of polyethylenimine. Four to five days post transfection, the cell media containing secreted recombinant protein was harvested by centrifugation at 5000 rpm for 45 minutes to remove the cell debris. The media was filtered through 0.22 μ m Amicon filters and dialyzed against a 10 \times volume of phosphate-buffered saline [10 mM sodium phosphate dibasic, 1.5 mM sodium phosphate monobasic, and 300 mM NaCl, pH 7.5] at 4 $^{\circ}$ C overnight. The dialyzed media was filtered again and 5 mM imidazole was added to the media before loading onto a Histrap column (GE healthcare). The protein was eluted using a linear imidazole gradient (20–500 mM) in 20 mM Tris (pH7.5) and 500 mM NaCl over 60 ml. Pooled protein fractions were deglycosylated with endoglycosidase F1 overnight at 4 $^{\circ}$ C and then concentrated for further purification with a size exclusion column (HiLoad 16/60 or 26/60 superdex 75 prep grade columns, GE Healthcare) in a buffer containing 20 mM Hepes (pH 7.5) and 150 mM NaCl. The eluted fraction corresponding to the monomeric species was concentrated to \sim 20 mg/ml for PvPLP1 and \sim 7 mg/ml for PvPLP2 and used for further studies.

The full-length PvPLP2 proteins (wild type and loop mutants) were expressed and purified in a similar way. In brief, kifunensine was not added during cell transfection and the harvested cell media was concentrated and buffer exchanged with PBS to a final volume of 500 ml as mentioned above. The dialysed medium was loaded to HisTrap HP (GE Healthcare) column and buffered changed with 50 mM MES (pH6.0) and 50 mM NaCl to load onto a cation exchange column (HiTrapTM SP HP, GE Healthcare). A linear salt gradient (50 mM–1 M NaCl) in 50 mM MES was used for the elution over 60 ml. The protein fractions were further purified using SEC (Superdex 200) in 20 mM Hepes (pH7.5) and 150 mM NaCl. Several smaller protein bands appeared in purified PvPLP2 full-length samples, which may be caused by partial proteolysis. The protein was concentrated to \sim 1.5 mg/ml, flash-

frozen in liquid nitrogen and stored in -80° C until further use.

Crystallization, diffraction data collection, and structure determination

Crystallization of PvPLP1 APC β domain (\sim 20 mg/ml) was set up using a sitting-drop vapor diffusion method in CrystalQuick 96-well plates by mixing 100 nl of protein with 100 nl reservoir solution with equilibration against 95 μ l of reservoir. The initial crystals appeared after \sim 20 days in 20% PEG 3350, 0.2 M Tri-potassium citrate. As the initial crystals were small and highly branched, further optimisation screening using a dilution series of this condition yielded more pin-like crystals. The resulting crystals were harvested with 20% (v/v) glycerol as a cryoprotectant and flash-frozen in liquid nitrogen. The crystal diffraction data were collected on beamline I24 at Diamond Light Source (Didcot, UK). The crystal diffracted to 3.15 \AA resolution in space group $P2_1$. The structure was determined by molecular replacement using TgPLP1 APC β (PDB 5OUO) as a model, using phenix.phaser.^{39,40} The structure was refined in phenix.refine⁴¹ with manual corrections in Coot.⁴²

Lipid dot-blot assay

Commercial membrane lipid strips spotted with 100 pmol of 15 membrane lipids (Echelon Biosciences P-6002) were used for this assay. Membranes were first soaked in PBST (PBS plus 0.1% Tween20), then blocked using 3% (w/v) non-fat dry milk dissolved in PBST for 1 hour at room temperature. The membrane strips were then washed and incubated with 0.5 μ g/ml protein in PBST plus 3% milk at 4 $^{\circ}$ C overnight. Membranes were washed 5 times with PBST for 10 minutes each time and incubated with rabbit anti-penta-Histidine polyclonal antibody for 1 hour. Membranes were washed again as described above, and incubated with secondary anti-rabbit-HRP conjugate for another hour. Following a final washing step of the membrane, the lipid-bound protein was detected by enhanced chemiluminescence (ECL, BIO-RED).

Liposome sedimentation assay

POPC (1-palmitoyl-2-oleoyl-*sn*-glycero-3-phosphocholine), POPE (1-palmitoyl-2-oleoyl-*sn*-glycero-3-phosphoethanolamine), Cholesterol, Brain Total lipid Extract, Liver Total Lipid Extract, Sphingomyelin (brain, Porcine), Cardiolipin (bovine heart), POPS (1-palmitoyl-2-oleoyl-*sn*-glycero-3-phosphoserine), Egg-PA (L- α -phosphatidic acid, LPA) and Lyso-PS (L- α -lysophosphatidylserine) were purchased from Avanti Polar Lipids Inc. Liposomes with indicated compositions were prepared as described.¹¹ In brief, different

lipids dissolved in chloroform were mixed (1 mg in total) and dried in a clean pyrex tube overnight in a desiccator attached to a VARIO-SP diaphragm pump (Vacuubrand). The lipid film was then rehydrated in 0.5 ml solubilisation buffer (20 mM HEPES pH 7.5, 150 mM NaCl) by vigorous vortexing and 5–10 freeze–thaw cycles. The hydrated mixture was extruded through 100 nm polycarbonate membrane (Whatman) for 11 times and the resulting liposomes were stored at 4 °C and used within 2 days. To perform the assays, 2.5 µg of protein was incubated with 50 µl liposomes (2 mg/ml) for 1 hour at 37 °C. Protein mixed with 50 µl solubilisation buffer was used as a negative control. The protein-buffer/liposome mixtures were then centrifuged at 67,000 rpm in an ultracentrifuge (Optima™ TL with TLA100.4 rotor) for 20 min at 10 °C. The supernatant (S) was collected to examine the liposome-unbound proteins. The pellets (P) were resuspended in an equal volume of the supernatant using solubilisation buffer. Then equal volumes of supernatant and pellet were loaded for SDS-PAGE analysis. Each liposome sedimentation assay was performed at least 3 times.

Bioinformatics

Homologous proteins to PvPLP1 were identified and a multiple sequence alignment produced using HMMER (<https://www.hmmer.org/>, <https://www.ebi.ac.uk/Tools/hmmer/>). Conservation scores were mapped onto the structure using the ConSurf server.⁴³ Multiple sequence alignments were generated and plotted using using ESPript 3.0.⁴⁴ Homology models of APCβ domains and missing loop regions from the PvPLP1 APCβ crystal structure were generated using MODELLER software.⁴⁵ All homology models were evaluated via the MODELLER objective function that combines spatial restraints and force field terms enforcing proper stereochemistry.⁴⁶ The best templates had 29% sequence identity for PvPLP2, 19% for PvPLP4 and 21% for PvPLP5 homology models.

Coarse grained MD simulations

Coarse grained (CG) simulations were performed using the Martini forcefield, v2.2^{47,48} with protein secondary structure predefined used a force constant of 1000 kJ mol⁻¹ nm⁻². The lower elastic bond cut-off used was 0.5 nm, and the upper cut-off was 1.0 nm. Lennard-Jones interactions were shifted to zero between 0.9 and 1.1 nm. The electrostatic potential energy was shifted to zero between 0 and 1.1 nm. To set up CG simulations, the protein atomic structure was first converted into a Martini CG representation^{34,35} using the Martinize script publicly available from Martini.⁴⁹ Water and ions were added to solvate the system, at an ion concentration of 150 mM NaCl. The pressure was coupled semi-isotropically using the Berendsen algorithm⁵⁰ with a coupling constant of 4 ps, and maintained

at 1 bar with a compressibility of 5×10^{-6} bar⁻¹. (Semi-isotropic coupling maintains the pressure separately for xy and z dimensions of the simulations box which is useful for membrane simulations and reduces distortions in the bilayer.) As above, the system was kept at a constant temperature of 310 K coupled using velocity rescaling,⁵¹ with protein, water and ions, and lipids coupled separately and a coupling constant of 1 ps. The time-step for integration was 20 fs, and particle coordinates were written to the trajectory file every 200 ps.

CG simulations of APCβ domain association with membranes were performed following Yamamoto et al.⁵² Proteins placed 10 nm away from a preformed bilayer were run through 20 simulations with the APCβ domain rotated sequentially around its x, y and z axis to reduce directional bias. Each simulation was run for 1 µs to allow the protein to associate with the membrane. Analysis for each ensemble of association simulations involved production of a 2-dimensional histogram plotting distance between the protein residues and membrane to determine binding orientations. Lipid contacts were also calculated, in which the frequency of contacts between lipids and protein residues were calculated for each simulation and normalised to produce a fingerprint plot of lipid contacts. Contacts were defined as any lipid and residue within 0.6 nm proximity. This distance was reduced to 0.4 nm in the case of lipid contact calculations in atomistic simulations (see below).

Representative CG simulation outputs were converted for atomistic simulations as previously described.^{34,35} In brief, the lipid and protein regions of the CG outputs were separately converted, via alignment of the starting protein structure with the CG protein position and orientation, and via the use of atomistic fragments to build a full lipids model based on their CG particles. After an energy minimization step the lipid and protein regions were combined, followed by addition of water and counter-ions, to generate a full atomistic system for simulation. See 34 for further details.

Atomistic MD simulations

Atomistic simulations were performed using Gromacs v5.1^{53,54} with the Amber 99SB-ILDN force field⁵⁵ and the SPC water model⁵⁶ with bond lengths and angles constrained using the LINCS algorithm.⁵⁷ To reduce computational burden a cut off of 1 nm was used for electrostatic and Van der Waal's interactions; coulombic interactions were also treated using Particle-Mesh Ewald (PME) electrostatics.⁵⁸ Constant temperature at 310 K was achieved using velocity rescaling, with protein, lipid and solvent (water and ions) separately coupled to an external heat bath.⁵¹ Pressure was maintained at 1 bar with the Parrinello-Rahman barostat with a coupling constant of 1 ps.⁵⁹ After initial energy minimization, equilibration was performed with the positions of heavy atoms restrained using a force

constant of $1000 \text{ kJ mol}^{-1} \text{ nm}^{-3}$, allowing the solvent to equilibrate around the protein and lipid molecules. Restraints were then removed for final production runs which were produced in triplicate, with each simulation started with random velocities. Time steps of 0.002 ps were used with the writing out of atomic coordinates every 10 ps.

PMF calculations

Potential of mean force (PMF) calculations driven by umbrella sampling were used to determine the free energy of binding of APC β domains to membranes.⁶⁰ To set up the system, a steered MD simulation was used to pull the protein from a membrane bound conformational state along the membrane normal. A series of overlapping windows along this reaction coordinate were then defined, typically every 0.1 nm, and each window was simulated for 1000 ns. Lipids were fixed along the Z-component of the simulation using a force constant of $10 \text{ kJ mol}^{-1} \text{ nm}^{-2}$. The centre of mass of the protein relative to the membrane was restrained in each simulation with a force constant of $1000 \text{ kJ mol}^{-1} \text{ nm}^{-2}$. The PMF was calculated using GROMACS g_wham,^{61,62} with error estimates obtained using bootstrap analysis. All values were shifted so that the PMF value was 0 in bulk water. Convergence was judged based on the change in PMF value calculated across consecutive 100 ns time intervals. All umbrella sampling was performed using Martini CG simulations and produced using GROMACS⁶³ with protein structure visualisation using the Pymol molecular graphics suite.⁶⁴ Data were presented using the Matplotlib Python plotting software⁶⁵ and the Seaborn Python data visualization library (<https://zenodo.org/record/1313201#.YGCml9zTU2x>).

Table 2 provides a summary of all the simulations performed for this work.

Accession numbers

The PvPLP1 APC β domain crystal structure has been deposited in the RCSB PDB, accession code 7PLN.

CRedit authorship contribution statement

Sophie I. Williams: Conceptualization, Formal analysis, Funding acquisition, Investigation, Methodology, Validation, Visualization, Writing – original draft, Writing – review & editing. **Xiulian Yu:** Conceptualization, Formal analysis, Investigation, Methodology, Validation, Visualization, Writing – original draft, Writing – review & editing. **Tao Ni:** Conceptualization, Formal analysis, Investigation, Methodology, Supervision, Validation, Visualization, Writing – original draft, Writing – review & editing. **Robert J. C. Gilbert:** Conceptualization, Formal analysis, Funding acquisition, Investigation, Methodology,

Project administration, Resources, Supervision, Validation, Visualization, Writing – original draft, Writing – review & editing. **Phillip J. Stansfeld:** Conceptualization, Formal analysis, Funding acquisition, Investigation, Methodology, Project administration, Resources, Supervision, Validation, Visualization, Writing – original draft, Writing – review & editing.

DATA AVAILABILITY

Data will be made available on request.

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Declaration of Interest

The authors all declare that they have no conflicts of interest.

Appendix A. Supplementary material

Supplementary data to this article can be found online at <https://doi.org/10.1016/j.jmb.2022.167642>.

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Abbreviations:

PLP, perforin-like protein; Tg, *Toxoplasma gondii*; Pv, *Plasmodium Vivax*; APC β , Apicomplexan PLP C-terminal β -pleated sheet

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