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Structural basis of dynamic membrane recognition

by trans-Golgi network specific FAPP proteins

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Running title: Dynamic membrane recognition by FAPP proteins

Abstract

Glycosphingolipid metabolism relies on selective recruitment of the pleckstrin homology (PH) domains of FAPP proteins to the trans-Golgi network (TGN). The mechanism involved is unclear but requires recognition of phosphatidylinositol 4-phosphate (PI4P) within the Golgi membrane. We investigated the molecular basis of FAPP1-PH domain interactions with PI4P bilayers in liposome sedimentation and membrane partitioning assays. Our data reveals a mechanism in which FAPP-PH proteins preferentially target PI4P-containing liquid disordered membranes, while liquid ordered membranes were disfavored. Additionally, NMR spectroscopy was used to identify the binding determinants responsible for recognizing TGN-like bicelles including phosphoinositide and neighboring lipid molecules. Membrane penetration by the FAPP1-PH domain was mediated by an exposed, conserved hydrophobic wedge next to the PI4P recognition site and ringed by a network of complementary polar residues and basic charges. , Our data illuminates how insertion of a structured loop provides selectivity for sensing membrane fluidity and targeting to defined membrane zones and organelles. The determinants of this membrane sensing process are conserved across the CERT, OSBP and FAPP (COF) family. Hence lipid gradients result not only in differential membrane ordering along the secretory pathway, but also specifically localize diverse proteins through recognition of ensembles of lipid ligands in dynamic and deformable bilayers in order to promote anterograde trafficking.

Keywords

pleckstrin homology domain; lipid microdomains; membrane trafficking; phosphoinositide recognition; nuclear magnetic resonance spectroscopy

Abbreviations

Brain sphingomyelin (BSM), CERT, OSBP and FAPP (COF), chemical shift perturbation (CSP), cholera toxin subunit B (CTx), diheptanoyl phosphatidylcholine (DH₇PC), dihexanoyl phosphatidylcholine (DH₆PC), dimyristoyl phosphatidylcholine (DMPC), dioctanoyl (C₈), dioleoyl phosphatidylcholine (DOPC), n-dodecyl phosphocholine (DPC), dipalmitoyl phosphatidylcholine (DPPC), heteronuclear single quantum coherence spectroscopy (HSQC), giant unilamellar vesicle (GUV), phosphatidylinositol-four-phosphate adapter protein 1 (FAPP1), liposome sedimentation assay (LSA), membrane insertion loop (MIL), palmitoyl-oleyl phosphatidylcholine (POPC), pleckstrin homology (PH), paramagnetic relaxation enhancement (PRE), phosphatidylinositol-4-phosphate (PI4P), surface plasmon resonance (SPR), trans-Golgi network (TGN), Texas-Red dihexadecanoyl phosphoethanolamine (TR-DHPE)

Introduction

Diverse biological membranes within cells selectively recruit thousands of proteins using a phosphoinositide (PIP) recognition code [1]. Such proteins transiently associate with membrane surfaces by recognizing phospholipids that exhibit organelle-specific distributions [2]. Cognate membrane complexes are further stabilized by complementary electrostatics [3-5] and insertion of nearby motifs of aliphatic, aromatic [6-8] or lipidated residues [9-11] into the bilayer. However, whether bilayer dynamics play a determining role in membrane recognition remains poorly defined [12, 13], necessitating a closer examination of the structures of functional complexes.

The largest superfamily of membrane interactive proteins is that defined by the presence of PH domains. Its members include COF proteins, which traffic ceramide, sterols and glycosphingolipids at the TGN [14-17]. Amongst them, the FAPP1 and FAPP2 proteins have

become the best understood paradigms for recruitment to membranes enriched in PI4P, which is the most abundant monophosphorylated inositol lipid in the Golgi membrane. Hence their PH domains are often used as Golgi markers in cellular studies [18, 19]. The distribution of PI4P however is not restricted to the Golgi apparatus, as this lipid signal also has critical roles beyond the Golgi [20]. Moreover, although COF family PH domains recognize PI4P physiologically [21-23], their *in vitro* PI specificities are not absolute [24-28]. This conundrum suggests that other conserved determinants also help to selectively attract these proteins to different organelle surfaces.

The two FAPP proteins differ mainly by the presence of a glycolipid transfer protein domain which is only found in the C-terminus of FAPP2 that has a critical role in the intra-Golgi vectorial transfer of glucosylceramide [21, 29] and in the synthesis of the globo-series of glycosphingolipids [30]. Both their FAPP-PH domains possess a similar hydrophobic wedge that inserts into bilayers, although its contribution to ligand specificity remains unclear [27, 28, 31]. An analogous membrane insertion loop is found in PI-binding FYVE and PX domains [32-34], with FAPP-PH domains being distinguished by Golgi-specific functions including membrane tubule budding and vesiculation [21, 31].

Here, the structural basis of lipid bilayer recognition by FAPP1 is resolved by analysis of protein partitioning into liposomal systems, mutagenesis and NMR. This yields experimental restraints for calculating the structure of ternary complexes of the protein with bound micelle and lipid ligand, thus explaining the requirement for loosely packed bilayers and pinpointing determinants for nonspecific and specific membrane engagement. We propose that modulation of PIs between micro-environments of varying membrane fluidity, as determined by the ratio of lipids including cholesterol and sphingomyelin (Fig. 1), provides a general switch for whether embedded lipid signals are accessible for protein recruitment at specific sub-organelle compartments.

Results

Specificity of FAPP-PH for disordered bilayers

In order to determine whether membrane order is important for FAPP-PH PI4P binding, we systematically explored sectors of the composition-dependent phase diagram of ternary lipid mixtures consisting of dioleoyl phosphatidylcholine (DOPC), cholesterol and brain sphingomyelin BSM [35-37] and supplemented with 2% dipalmitoyl (DP) PI4P unless stated otherwise. Membrane order was validated with GUVs, employing fluorescent lifetime images with environmentally sensitive di-4-ANEPPDHQ (di-4) membrane reporter (Fig. S1). It was evident that association with PI4P-containing membranes decreased as ratios of both cholesterol and brain sphingomyelin (BSM) increased. In pure liquid ordered (L_o) liposomes formed of an equimolar ratio of BSM and cholesterol, the FAPP1-PH domain interacted only marginally with the membrane (Fig. 2A). The most attractive PI4P containing membranes were those forming liquid disordered (L_d) membrane phases (Fig. 2A), suggesting that lipid packing and dynamics could play a key role in efficient PI4P recognition and binding.

Next, we investigated the bilayer packing density as a function of the acyl chain saturation. The headgroups of DPPC, palmitoyl-oleyl (PO) PC and DOPC were compared as they occupy areas of 64, 68 and 72 Å/nm², respectively, with concomitant effects on bilayer thickness [38]. Binding of FAPP1 to the set of PI4P/PC-based liposomes revealed the highest level of binding to loosely packed DOPC vesicles, intermediate density POPC vesicles exhibited medium binding, while the most densely packed DPPC vesicles excluded the FAPP1-PH (Fig. 2A). Thus the lipid packing density was inversely related to protein binding. Hence the membrane specificity of FAPP1-PH combines both essential chemical (i.e. PI4P) and physical (i.e. membrane fluidity)

factors. Interestingly, this preference was also found with the full-length construct of FAPP2 (Fig. 2A), indicating broader relevance across the COF family.

The specific role played by membrane order in FAPP interactions with PI4P was further validated by fluorescent microscopy using GUVs. An equimolar ratio of cholesterol, DOPC and BSM was used to follow FAPP1-PH binding to PI4P in membranes exhibiting coexisting L_d and L_o domains. The fluorescent cholera toxin subunit (CTx) bound to the ganglioside GM1 [39] and Texas-Red dihexadecanoyl-phosphoethanolamine (TR-DHPE) [40] were used to mark the respective membrane domains, with the latter label preferentially partitioning to L_d domains (Fig. 2B). The presence of protein was detected by visualizing FAPP1-PH labelled with Oregon Green to an exposed unique cysteine (Cys37) (FAPP1-PH^{green}) (Fig. 2C). The FAPP1-PH^{green} protein localized strictly to L_d domains in phase-segregated GUVs, consistent with the liposome sedimentation experiments (Fig 2A). Furthermore, this localization to disordered regions could be similarly mediated by either naturally (brain extract) or synthetically derived PI4P (dipalmitoyl or dioctanoyl) (Fig. 2C). This indicates that the phosphoinositide headgroup (rather than the respective acyl chains which differed) was the primary determinant of PI4P recognition in liquid disordered membrane regions. Binding to vesicles could be initiated by injecting soluble short chain dioctanoyl (C_8) PI4P molecules into the chamber of reaction. In particular, an approximate four-fold enhancement in protein co-localization with the disordered phase was observed after addition of 2 μ M C_8 -PI4P. This increase can be attributed in part to the inherent preference of short acyl chain lipids for disordered phases [41]. Thus FAPP1-PH associates with L_d membranes phases potentially due to a favorable access to lipid headgroup, non-stereospecific contacts with PC molecules or suitable dynamics therein. Once bound to the disordered bilayer, the protein can conceivably diffuse laterally until it recognizes the headgroup of its PI4P ligand, thus forming a stable complex.

Bicelles mimic the bilayer for FAPP1-PH binding

Before characterizing the structural basis of dynamic bilayer recognition we needed to select an appropriate bilayer mimic for quantitative analysis of the interactions. The binding of the FAPP1-PH domains with micelles and bicelles were compared as these both mimic biological bilayers but have different chain lengths and curvature properties. FAPP1-PH exhibited similar patterns of amide signal changes when bound to PI4P in the presence of micelles composed of n-dodecyl phosphocholine (DPC) or bicelles composed of dimyristoyl phosphocholine (DMPC) and dihexanoyl phosphatidylcholine (DH₆PC), indicating consistent insertion modes. Moreover their ligand affinities were similar, with the dispersed methyl resonance migrations yielding C₈-PI4P dissociation constants of $5.3 \pm 2.4 \mu\text{M}$ and $8.8 \pm 3.3 \mu\text{M}$, (Figs. S2A,B), comparable to the affinity of small unilamellar vesicles (Fig. 6C). Furthermore, the chemical shifts of the protein bound to isotropic bicelles matched those of the state saturated with DPC:CHAPS micelles. Minor spectral differences were observed between PI4P-micelle and PI4P-bicelle titrations. These arose largely from the absence of assigned crosspeaks from the bound states due to overlapped or broadened resonances. Thus no significant difference was evident between micelle and bicelle complexes, and either state suitably represents how FAPP1-PH orients on bilayers *via* dynamic membrane insertion.

Resolving multi-step bicelle binding by NMR

Previous studies have investigated how FAPP1-PH interacts with soluble PI4P molecules and associates with micelles [27], providing a basis for elucidating how the protein assembles on fluid bilayers which contain PI4P. In order to distinguish the determinants of transient nonspecific bilayer association and stable recognition of bilayer-embedded PI4P, we compared the binary and ternary complexes formed by FAPP1-PH, PI4P and micelles or bicelles in solution. This approach allowed identification of the multiple structural states formed by the FAPP1-PH domain as it interacts with membrane mimics of increasing size and complexity. First the micelle and bicelle-saturated forms of FAPP1-PH were contrasted to identify the respective interactions and

to model the PI4P-specific complexes formed. To gain maximal resolution each state was characterized by monitoring backbone and sidechain ^1H , ^{13}C and ^{15}N signals upon addition of soluble PI4P and stable bicelle formulations. Entry of the free state from solution into the bilayer was examined by tracking methyl and amide peaks as bicelles were added stepwise. This allowed the progressive changes induced by each binding event to be mapped at atomic resolution (Fig. 3A,B), showing that the interaction was principally mediated by the $\beta 1$ - $\beta 2$ hairpin which spans FAPP1-PH residues Trp8 to Gln16, as was confirmed by results from paramagnetic relaxation enhancement (PRE) studies (Fig. 3C) [27]. Moreover, the data indicated a second, distal interaction site that includes the $\beta 6$ - $\beta 7$ sheet residues Met61, Glu62, Leu63, Ile64, Glu68, His70 and Tyr72. Together this provides experimental evidence for two-pronged stable insertion *via* the $\beta 1$ - $\beta 2$ and $\beta 6$ - $\beta 7$ elements.

Next the structural orientation of the FAPP1-PH protein on bilayers was characterized by adding C_8 -PI4P to the protein-micelle (Fig. 4A,D) and protein-bicelle assemblies (Fig. 4C,E). Intermolecular distance restraints were derived from doxylated lipid molecules (Fig. 3C, 4B) and soluble gadolinium agents (Fig. 5A,B). This illuminated the multiple states of free, nonspecifically PC-associated and PI4P-specific bilayer bound protein, respectively. The experimental setup used to elucidate the structure of the micelle complex was adapted for bicelles, with the association being followed using PREs induced by paramagnetic gadolinium to identify solvent-exposed groups in the free, bicelle and bicelle-PI4P complexes. This general protocol provides a broadly applicable basis for experimentally-based elucidation and validation of multiple bilayer-complexed structural states.

Structural basis of PI4P-bilayer recognition

Having defined the conditions for elucidating the consensus solution structure in bicelles and micelles, the wild type FAPP1-PH domain responsible for dynamic bilayer insertion was

determined by triple resonance NMR methods. Its structure is largely identical to the mutant version solved previously (PDB: 2KCJ), with chemical shift differences being localized to the native Cys which had been previously replaced with a Ser to minimize opportunities for cross-linking. The structure of the monomeric native FAPP1-PH domain was defined by 1448 distance derived NOE, 128 dihedral angle and 34 hydrogen bond restraints, with the structural statistics being summarized in Table 1.

The micelle complexed structure of the FAPP1-PH domain was solved using the HADDOCK program. The restraints included 16 intermolecular distances measured from ^{13}C methyl and ^{15}N amide-resolved PREs to the micelle, as obtained with the 5-doxyl PC spin label (Tables 1&2). A flexible zone of neighbouring residues was defined as those which exhibited substantial chemical shift changes. Solvent accessibility data was derived from titrating in the gadolinium agent into ^{15}N and ^{13}C isotopically labeled protein samples. Bicelles composed of dihexanoyl (DH₇-) and dimyristoyl (DM-) PC (q=0.3, 0.25%) were added stepwise. The advantage of the new bicelle formulation over conventional bicelles composed of DH₆PC and DMPC formulations was apparent as small increments could be added from initial concentrations as low as 0.1% (w/v). This enabled monitoring of the bicelle-bound state during titration experiments. This formulation was optimized for low concentrations [42], and adapted herein to track protein resonances while conserving a signal that was sufficiently strong and resolved to be useful for PRE analysis. The solvent-exposed and micelle-embedded protein surfaces were mapped by analysis of PRE data collected using water-soluble gadolinium or doxyl PC spin labels, respectively, thus defining their complementary areas (Figure 5A-C). Interfacial residues were defined as being those with resonances that were broadened by both gadolinium and doxylated phospholipid. Together this provided cross-validated definition of the FAPP1-PH protein groups that were exposed, interfacial or deeply inserted.

The structural model of the complex based on PRE data reveals that membrane recruitment of FAPP1 involved deep insertion of Tyr11 and Leu12. Their sidechains penetrated furthest into the hydrophobic interior of the bilayer, while in solution they formed a solvent-exposed extremity at the tip of the structured β 1- β 2 loop. Their sidechains orient beneath the surface of phospholipid headgroups in either the micelle or bicelle complex in the presence or absence of PI4P, indicating a constitutive role in insertion. Moreover, the specific PI4P recognition event uniquely involved insertion of the sidechain of β 7 residue Lys74. This basic residue is conserved across the COF protein family members, and interfaces peripherally with the bilayer (Fig. 4F), thus contributing to the higher affinity of the PI4P-containing membrane complex.

The deep burial of the Leu12 and Thr13 methyl groups was confirmed by the substantial changes in solvent accessibility observed after gadolinium addition (Fig. 5B). These residues were encircled by an extensive network of interfacial contacts mediated by polar backbone and sidechain groups of residues in the β 1- β 2 loop including Asn10 and Gln16, which form hydrogen bonds and ionic interactions with PC headgroups (Table 1). The Trp8 and Trp15 residues act as aromatic buttresses against the membrane surface (Fig. 6A-B), delimiting the total buried surface area of either 1208 Å² or 1232 Å² between the protein and micelle in the absence or presence of PI4P, respectively. This insertion element is ordered with the exception of the most exposed Leu12 residue based on model-free relaxation analysis (Fig. S3) [43]. Together this reveals that overlapping structured protein surfaces mediate transient nonspecific and PI4P-specific binding of fluid bilayer phases, respectively. Thus we infer that initial, weak bilayer entry of the β 1- β 2 loop with primarily loosely packed PC molecules leads to a PI4P-dependent adjustment of the orientation of the inserted complex.

The structural organization with the ternary FAPP1-PH:PI4P: bilayer complex can be ascertained by NMR following C₈-PI4P addition to the micelle complex. Phosphoinositide addition results in the unprecedented stable micelle burial of the Met73 side chain in the β 7 strand. Considering the

absence of 5-doxyl PRE effects in the Met73 resonances, this change in exposure could be due to its enclosure within the stable PI4P-micelle complex (Fig. 5C). This indicates a PI4P ligand-dependent bilayer interaction by this structured element, with the proximal Leu63 and Ile64 residues also engaged based on their chemical shift changes. This ternary complex exhibits a slower off-rate and extensive zone of perturbations midway between the most deeply inserted β 1- β 2 extremity and the exposed termini (Fig. 4D-E). The large perturbed zone observed in the PI4P-specific complex involves all the β -strands including β 1:Val4-Tyr6, β 2:Thr13, β 3:Leu27,Tyr29, β 4:Gly42, β 5:Glu50, β 6:Met6, Leu63, Ile64, β 7:Phe71-Lys74 residues (Fig. 4D-E). The micelle-based burial of these residues was confirmed by changes in solvent accessibility upon addition of the paramagnetic gadodiamide agent, with a broader interface being buried when PI4P was present, including His54, Thr59, Glu62, Phe71 and Lys74 (Fig. 5A). Together with the entire lower half of the protein exhibiting chemical shift perturbations (CSP) and the interfacial position of the β 5- β 7 sheet, this indicates that specific recognition of PI4P-containing bilayers involves stable rather than transient dipping of the hydrophobic β 1- β 2 tip into the bilayer interior [27], and adjustment to a more substantially buried state.

In addition to the hydrophobic contribution, it is well known that electrostatics also influence the orientation of proteins on membrane surfaces. Indeed, this is reflected by the dipolar nature of the FAPP1-PH surface (Fig. 5D). Two distinct positively charged isocontours facing the micellar interface correspond to the PI4P binding pocket and elements of the β 7 strand including the Lys74 side chain [23]. The resulting electrostatic complementarity could account for the two-fold increase in binding affinity observed when negatively charged phosphatidylserine is added to the membrane to stabilize the specific PI4P-containing complex [27]. Indeed, analogous preferences for phosphatidylserine co-association is apparent in PI3P-specific FYVE and PX domains [44, 45], indicating a wider role for this accessory phospholipid as a co-determinant of intracellular membrane binding.

The ring of charge encircling FAPP1's hydrophobic membrane insertion loop (MIL) appears to delimit the depth of membrane insertion. In order to estimate the depth and angle of protein penetration into the bilayer in nonspecific and specific membrane engagement, we compared how FAPP1-PH inserted into PI4P-free versus PI4P-containing micelles, taking advantage of the sensitivity of the NMR method to even transient interactions. The methyl and amide PRE data yielded average distances of 38.47 Å and 37.14 Å between protein and micelle centers, indicating similar depths in the specific and nonspecific membrane complexes, respectively (Fig. 3, 4 and 6A). That is, the specific complex penetrated only an angstrom deeper and its insertion angle was only slightly more acute. This can be attributed to PI4P-containing micelle contacts formed by Lys74, the neighboring β 5- β 6 and β 6- β 7 loops and the extremity of the β 1- β 2 hairpin. PI4P-binding by the micelle-saturated complex also induced alterations in hydrophobic core packing of the protein based on perturbations of methyl resonances of either partially buried Val4 and Ile64 residues or completely buried Leu5, Ile44, Leu63, Ile65, Met73 residues. We propose that this network reinforces the slowly exchanging PI4P bilayer complex, and increases the residency time and local concentration of protein molecules on membranes. Upon bilayer association, the hydrophobic wedge of the FAPP1-PH domain is not symmetrically inserted. Rather, it is oriented such that the backbone residues prior to the β 2 strand (Thr13, Gly14, Trp15) deeply insert into micelles while the backbone of conserved residues following the β 1 strand binds the interfacial inositide headgroup *via* Trp8 and Thr9 contacts. The local ring of basic charge capped by an acidic patch (Fig. 5D) provides long-range guidance for the protein's entry into the membrane. Together this allows progressive binding and tilting of the bilayer complex, and could have implications for interaction of downstream protein partners such as Arf1 with TGN vesiculation machinery [46].

Mutational analysis of membrane binding determinants

The residues identified in the structural model as membrane interacting were mutated to delineate their roles. All mutants were tested for being structurally intact and functionally altered (Table S1). Binding of FAPP-PH proteins to membranes was measured using liposome sedimentation experiments. The most deeply inserted Tyr11 and Leu12 sidechains are known to be essential [31], but the contributions of other residues of the loop, including Trp8, Asn10, Thr13, Trp15 and Gln16 residues have not been determined. Each Trp residue extends its sidechain to pack against the micelle surface, and hence was mutated to an Ala, Glu or Tyr residue. The W8E and W8Y mutants of FAPP1-PH did not bind detectably (Table 3, Fig. 6C), indicating that Trp8 makes critical contributions in both nonspecific and specific membrane complexes. This is consistent with its interfacial positions in the binary and ternary structures with PI4P and micelle.

Mutations of the non-conserved Thr13 residue were designed to conserve the hydrogen-bond acceptor (T13N) and to alter the exposed hydrophobicity (T13F). Both mutations were tolerated with only minor reductions in binding by a factor of two, consistent with the peripheral role of Thr13 in nonspecific insertion. Moreover Thr13 is not conserved across the COF family, which maintains basic or small hydrophilic residues at this position [27]. Conversely, the N10T mutation compromised bilayer affinity by a small but significant degree, as can be explained by the insertion of its sidechain between PC-based headgroups and its role in hydrogen bonding (Table 2). The substitution of Trp15 with a Tyr (but not Glu, which results in unstable protein) was tolerated, consistent with the critical role of its aromatic ring in insertion as well as stabilising the β 1- β 2 hairpin. The replacement of Gln16 with an Arg residue was originally intended to mimic the polybasic motif found in PH domains which recognizes 3-phosphoinositides [47]. Unexpectedly, this mutation reduced liposome association, independent of the PIP species used. The major role of Gln16, which is strictly conserved across the mammalian COF family,

can be explained by the hydrogen bonds it consistently forms with PC molecules when fully docked.

Together this reveals that liquid disordered membrane interaction is mediated by essentially the entire breadth of the ordered β 1- β 2 hairpin loop (Fig. S3) which is ordered except for the hydrophobic tip. In particular, the hydrophobic extremity offered by Tyr11 and Leu12 is bordered by non-essential Asn10 and Thr13 contacts, and the highly conserved aromatic groups contributed by Trp8 and Trp15 form critical struts against disordered bilayer surfaces. We propose that sliding of this structured loop through the leaflet allows the protein to diffuse in the two dimensional plane of a membrane, with electrostatic forces supporting the protein's positioning on the bilayer for PI headgroup entry. Once a PI4P molecule is bound, the mobility of the resulting complex would be reduced by the additional bound bulk, and the conformational dynamics in the protein-bilayer complex including in angle and depth of insertion and core packing would shift to that of the fully occupied state. The accompanying displacement of lipid molecules and resulting perturbation of local pressure and surface area created could contribute to the deformation of membranes during budding or tubulation events.

Discussion

The ability of FAPP proteins to specifically recognize PI4P-enriched Golgi membranes is determined by a set of unique features which are revealed here by NMR. The mechanism involves proximal penetration of the structured MIL residues and PI4P acyl chains into a liquid disordered bilayer, introducing substantial protein volume into the leaflet. This insertion would naturally yield positive local curvature, and hence would be opposed by the tendency of cholesterol to induce negative membrane curvature. The FAPP1-PH domain inserts deeply *via* not only the longest structured MIL studied to date but also by the β 7 strand, as supported in the

case of FAPP1 by ^{13}C - and ^{15}N -resolved backbone and sidechain groups for micelle- or bicelle- embedded and soluble spin labels. The use of both heteronuclei for gathering PRE restraints and the use of optimized bicelles yielded the highest density of experimental protein-bilayer structural restraints to date to our knowledge.

TGN bilayer binding mediated by FAPP1-PH is initiated by nonspecific phospholipid interaction followed by specific binding to a PI4P molecule. These events differ surprisingly little in the depth or angle of bilayer penetration. The most significant differences are the structuring of the PI4P site and conformational adjustment within the core as the ligand is bound. This yields a slowly exchanging complex with a binding affinity of $5\ \mu\text{M}$ for the monomeric protein. The fact that the MIL is largely structured is unanticipated given its long and irregular nature, as is the $\beta 7$ binding element, yet could explain their specificity for dynamic bilayers. The interfacial region involves functionally critical tryptophanes of the loop and cationic residues that engage the surface of the bilayer and support the orientation of the embedded protein. In particular, Trp8 and Trp15 form struts at either extremity of the MIL, while Lys74 is opposite the PI4P binding site and forms hydrogen bonds with the membrane surface. These aromatic and basic residues are highly conserved in the FAPP family, and occupy similar positions around the basic patch of the CERT structure [23], inferring a common mechanism.

This general FAPP-bilayer binding mechanism is depicted in Figure 7. The process involves the electrostatic approach and insertion of the FAPP1 protein into disordered bilayer within a restricted membrane territory [48]. The resulting reorientation of lipids within the leaflet includes displacement of dynamic lipid allowing PI4P molecules to be more readily encountered before being stably bound within the appropriately positioned PI4P site. The resulting asymmetric insertion of protein bulk within the bilayer would induce local positive bilayer curvature. Interestingly, the insertion depth and angle are only slightly perturbed by the transition from nonspecific to specific complex, instead the original contacts become reinforced, resulting in the

tighter FAPP:PI4P complex becoming specifically localized and crowded within dynamic TGN zones. This process leaves a complementary site on the FAPP1-PH domain largely available for Arf1 docking [46] that is necessary but not sufficient to localize FAPPs at the TGN [21]. We note that while preparing this manuscript, a study addressed the mechanism of yeast Arf1 binding to human FAPP1-PH [49], and is largely consistent with our results.

The crucial role of the various COF proteins in lipid trafficking [16, 29, 30, 50] suggests that they share recognition determinants. The commonalities are most obvious for CERT [17, 51] and OSBP1 [52], localization of which has also been linked to Golgi membrane composition. The key residues are shared across the COF family, inferring similar assembly and membrane deformation processes. In phase segregated GUVs, the preference seen for *Ld* domains is such that these proteins could all be essentially completely directed only to disordered phases (Fig. 7). *In vitro*, such proteins would become crowded until reaching a critical concentration where buds form and tubules can then emanate. Biological membrane tubule formation by FAPPs has yet to be confirmed under physiological conditions. Nonetheless, tubules have been observed *in vitro* [31] and in cell-based assays [18, 21], and specific roles of contributing residues can now be tested in cellular TGN systems. Broader applicability of the general mechanism to other systems can be envisaged. Lipid enzymes may be similarly regulated by bilayer order. For example, phosphatidylinositol-4-phosphate kinase type II α , which produces half of the PI4P at the Golgi, is active once bound there within rigid microdomains [53].

In summary, we propose that membrane malleability represents a fundamental means of controlling protein recruitment to specific regions of organelles such as Golgi subcompartments. In endomembranes, lipid concentration gradients across the secretory pathway are found in opposing directions (Fig. 1), with the ratio of glycerophospholipids decreasing and sphingolipids and cholesterol increasing [54, 55]. Moreover, the saturation of the acyl chains for different lipid

species increase along the anterograde pathway [56]. As a result, the membrane rigidity and thickness are both enhanced as one travels from the endoplasmic reticulum towards the plasma membrane, with the TGN having an intermediate lipid composition prone to phase separations [57]. Hence, a membrane order dependent PI4P binding model would account for the absence of FAPPs at the plasma membrane despite pools of PI4P having been identified there [18]. Furthermore, changes of membrane rigidity could efficiently control the recruitment of CERT, OSBP and FAPP proteins to the TGN to maintain local lipid homeostasis. Thus, as sphingolipids and cholesterol are recruited at the TGN by leaving vesicles, reduced membrane packing densities would lead to an enhanced lipid transfer rate of FAPP2 (glucosylceramide), CERT (ceramide), OSBP (sterol) which would progressively re-establish the rigidity, acting as a negative feedback loop on the lipid transfer proteins. In other words, PI4P molecules in membranes with opposing membrane fluidities molecules could simplistically represent 'on' and 'off' signaling states for recruiting FAPP proteins to regions in the TGN membrane gradients. Although PI4P is present in other subcellular membranes where it critically contributes to other biological processes [20], it may not be visible there to these proteins due to its ordered microenvironment. An analogous phenomenon has been invoked for recognition of the sphingolipid GM1 by cholera toxin B subunit [58, 59]. The presence of cholesterol in the membranes forced GM1 headgroups to bend, in which conformation the toxin no longer recognizes them. Thus we propose that this principle of binary lipid order recognition states also applies to other phosphoinositides and lipids, which may be similarly visible or invisible to binding partners depending on the local conformational dynamics of the membrane[60].

Materials and Methods

Protein expression

FAPP1-PH was expressed in a pGEX-6P-1 vector (G.E. Healthcare, Little Chalfont, UK) as a glutathione S-transferase (GST) fusion protein and purified as previously described [27]. Uniform isotopic labelling with ^{15}N or $^{13}\text{C}/^{15}\text{N}$ was carried out in M9 media. The cell lysate was passed through a GST Trap column and the protein was cleaved overnight using Prescission protease (G.E. Healthcare). The FAPP1-PH protein was separated by anion exchange using a linear gradient of NaCl from 0 to 0.5 M (Tris pH 8, 20 mM), and exchanged into 20 mM Tris pH 7, 100 mM NaCl, 1 mM DTT and 1mM NaN_3 (TB). The mutants were generated with a Quik Change Lightening kit (Stratagene, Santa Clara, USA) and verified by DNA sequencing. The proteins were expressed and purified as previously described [27] using optimized salt gradients to separate FAPP and GST proteins during the final purification step.

Protein fluorescent tagging

Conjugation of FAPP1-PH protein to the Oregon Green maleimide fluorescent probe (Life Technologies) was performed according to the manufacturer's protocols. Briefly, complete reduction of disulfide bonds was achieved in TB with 10 mM DTT for 1 hour. Subsequently, the sample of 100 μM protein was buffer exchanged with TB and incubated for 4h at room temperature with maleimide fluorescent probe in a 10 times excess and purified on PD-10 columns.

Lipid binding assays

Lipids, detergents and natural extract of PI4P (brain extract) were purchased from Avanti Polar Lipids (Alabaster, USA) and synthetic PIPs including C_8 -PI4P from Echelon Biosciences (Salt Lake City, USA). Micelle and bicelle titration experiments were carried out by stepwise additions of buffer-matched stock solutions into NMR samples. Bicelles were generated using DMPC and DH_6PC mixed in chloroform, dried under a flux of nitrogen and left under high vacuum overnight. A stock solution at 25% was prepared and diluted with the protein immediately before each

experiment. Bicelles with a ratio $q=0.25$ were used at a 5% (w/v) to prevent excessive line broadening and alignment of the bicelles with the magnetic field. Bicelles containing DH₇PC and DMPC ($q=0.3$, 0.25%) were prepared for acquisition of ¹⁵N-edited HSQC experiments.

Lipids were mixed in chloroform, the organic solvent was successively dried under a nitrogen stream, and samples were placed under high vacuum overnight. The lipids were resuspended in TB to a lipid concentration of 2 mM, and suspension was submitted to ten cycles of freezing in liquid N₂ and thawing at 52°C. For the assays, 75 μL of the lipid suspension was mixed with 25 μL of protein at 8 μM and incubated at room temperature for 10 min. The pellet collected after centrifugation (55 000 rpm, 4°C, 10 min) with a TLA-55 rotor (Beckman Coulter, High Wycombe, UK) was washed three times with TB and resuspended in 100 μL. The supernatant and the pellet of each assay were loaded on precast 26 wells Criterion gel (Biorad, Hemel Hempstead, UK). After electrophoresis, the proteins were stained by blue Coomassie and quantified by gel imaging (Syngene, Cambridge, UK). Values represent the mean and standard deviations from triplicate experiments.

Surface Plasmon Resonance

The surface plasmon resonance (SPR) experiments were carried out in TB on a Biacore 3000 instrument using sensor Chip L1 (G.E. Healthcare). A suspension of 1 g/L lipids DOPC:PI4P (98:2) or DOPC were submitted to ten FAT cycles and extruded through a 100nm membrane (Avestin). A total of 150 μL of this lipid suspensions were coated on the lanes at 5 μL/min, washed with 10 mM NaOH and coated with 25 μL BSA and cleaned again with 10mM NaOH. Equilibrium measurements were carried out at 2-3μL/min, and the sensograms were obtained for an analyte range 0.1 to 20 μM and corrected by subtracting the reference signals from the DOPC lanes. The apparent dissociation constant was deduced from the fitting $B = \frac{B_{\max} \cdot P_{\text{free}}}{(K_d + P_{\text{free}})}$ where B is the binding, B_{\max} is the maximum signal under saturation and P_{free}

is the concentration of protein present in the solution injected. In all experiments, the response of FAPP1-PH^{WT} at 20 μ M was chosen as standard to normalize the response of the mutants.

GUV formation

A volume of 25 μ L of the lipid mixtures at 1 g/L in chloroform were spread on ITO-coated slides (Sigma) and dried under vacuum for at least two hours. GUVs were grown at 52 °C in 150 mM sucrose. A sinusoidal current (1.1 Vpp, 12Hz) was applied for 2 to 3 hours followed by a squared current (1.5 Vpp, 5Hz). The GUVs were collected and re-suspended in a Tris buffer (10mM, pH 7, NaCl 50mM, DTT 1mM). Chambers were built using double sided tapes and the passivation of the surface was achieved with a solution containing 1 mg/mL of casein. After washing with resuspending buffer, the GUVs were injected into the observation cell. Alternatively, glass bottom dishes (MatTek, Ashland, US) were used for microscopic observations to allow uniform injections across the sample. To visualize different phases, GM1 at 1% (mol/mol) was included in the lipid mixtures. The fluorescent probes Cholera Toxin subunit B tagged with AlexaFluor 488 (CTx) (Life Technologies, Carlsbad, CA) and Texas Red 1,2-dihexadecanoyl-*sn*-glycero-3-phosphoethanolamine (TR-DHPE) (Life Technologies) were used to mark ordered and disordered phases, respectively. The image processing and were performed within ImageJ software [61]. Green emitting dyes were excited at 488 nm and red emitting dyes at 543 nm.

Fluorescence lifetime imaging microscopy

GUVs were prepared according to established methods [62] and stained with 2 μ M di-4-ANEPPDHQ (di-4) dye (Invitrogen). Fluorescence lifetime imaging microscopy (FLIM) was performed at 23 °C with a LSM 510 microscope (Carl Zeiss) equipped with a dedicated PicoQuant FLIM system. The probe was excited with a 473nm pulsed laser diode (50 MHz) and observed with a 63x oil immersion objective and fluorescence was collected through a 495nm long wave pass filter. Laser power was adjusted to give an average photon rate of 10^4 – 10^5

photons to avoid pile-up effects. The acquisition time was of the order of 200 s to achieve at least 10^3 photons per pixel.

Thermal shift assay

In order to determine the stabilities of mutant proteins, fluorescent signals were measured using excitation at 492 and emission at 568 from solutions containing 4 μ M protein and SYPRO Orange (Life Technologies) in 50mM HEPES buffer pH 7, 100mM NaCl. Signals were followed along a linear temperature gradient between 25°C and 95°C. The experiments were carried out on an MxPro3005P qPCR detection system and processed with MxPro software (Stratagene). The unfolding transition temperature, T_m , of each protein corresponded to the point of inflexion of the curves [63].

Nuclear magnetic resonance spectroscopy

NMR experiments were acquired at 298K on an 800 MHz Varian INOVA spectrometer or a Bruker AVANCE III spectrometer equipped with 5mm cryogenic probes, using samples containing 100 to 700 μ M FAPP1-PH protein. Slowly exchanging amide protons were identified from ^{15}N HSQC spectra acquired following dissolution in 99.96% D_2O . Backbone and side-chain resonances were assigned in part by referring to those of the C94S mutant protein. NMR experiments were run using Varian BioPack pulse sequences, including BEST HNC0, HNCA, HN(CO)CA and ^{13}C -edited NOESY-HSQC (100ms mixing time), acquired from $^{13}\text{C}/^{15}\text{N}$ -labelled protein samples containing 10% D_2O . The ^{15}N relaxation experiments were acquired using pulse sequences available in the Bruker standard library and with ^{15}N -labelled protein.

Backbone generalised order parameters squared, S^2 , were determined with the model-free formalism [64, 65] from ^{15}N relaxation data using the diffusion tensor obtained for an axially symmetric motional model. The ^{15}N R_1 and R_2 relaxation rates and $\{^1\text{H}\}$ - ^{15}N heteronuclear NOE values for FAPP1-PH were measured at a ^1H frequency of 600 MHz at 298K using established

methods [66, 67]. To estimate the R_1 and R_2 values, monoexponential two-parameter decay functions were fit to peak intensity versus measured relaxation delay profiles using the Analysis program from the CcpNmr software suite. NOE values were determined from the ratio of peak intensity in the proton saturated spectrum versus peak intensity in the unsaturated spectrum for a given resonance. Errors were calculated from repeat measurements (R_1 , R_2), or from an analysis of background noise in the spectrum when repeat spectra were not available. For model-free analysis, an initial estimate of the rotational diffusion tensor was obtained from the R_2/R_1 ratios of individual so-called “rigid” residues and the PDB coordinates of the FAPP1-PH solution structure using the programs `pdbinertia`, `r2r1_diffusion` and `quadric_diffusion` (A.G. Palmer, Columbia). Residues were considered rigid if they satisfied the following criteria: 1) NOE > 0.65, and 2) of the remaining residues, $\langle R_2 \rangle - R_{2,n} < 1$ S.D. (where S.D. is standard deviation from the mean, $\langle R_2 \rangle$). Fitting of the R_2/R_1 ratios from 51 rigid residues in total was performed using different rotational diffusion tensors: isotropic, axial and fully anisotropic with established model selection criteria [68]. Amide proton-nitrogen bond lengths of 1.02 Å and ^{15}N chemical shift anisotropy of -160 ppm were assumed for all residues. Order parameters were subsequently determined following the flowchart of Mandel et al, using their software ModelFree v. 4.2 [68] and FAST ModelFree [69]. Uncertainties in the model-free order parameters were estimated from 500 Monte Carlo simulations.

PRE experiments were performed as described elsewhere [27]. DPPC was substituted for 5 doxyl-PC in reference experiments to ensure that only 5-doxyl-dependent NMR intensity differences were measured. The gadodiamide agent (Selleckchem, Newmarket, UK) was added to protein:lipid assemblies in NMR samples from a concentrated stock solution in NMR buffer. The chemical shifts perturbations were calculated as $\Delta = (\Delta_{\text{HN}}^2 + \alpha \Delta_{\text{HX}}^2)^{1/2}$ where α is the ratio of the magnetic ratios of nuclei (0.1 for $X = ^{15}\text{N}$ and 0.25 for $X = ^{13}\text{C}$). The dissociation constant K_d was calculated from $\Delta = \Delta_{\text{max}} (L_T + P_T + K_d - [(L_T + P_T + K_d)^2 - 4L_T P_T]^{1/2}) / 2P_T$ where L_T and P_T are the ligand and protein

concentrations and Δ represents the chemical shift change. The chemical shift perturbations were calculated with 2mM DPC:CHAPS as a reference to account for possible nonspecific interactions of unimolecular PC molecules in solution. The perturbations were calculated relative to micellar concentrations while bicelle-dependent changes were measured in reference to a protein NMR sample free of any lipid. The NMR spectra were processed using NMRPipe [70], the resonance assignments and the structure calculations were carried out in CcpNmr analysis suite [71] and Aria2 [72], respectively. The structures were generated by restrained torsion angle dynamics in eight iterations using Aria2. After a final water refinement, the 20 lowest energy structures out of 100 calculated were selected to represent the ensemble of FAPP1-PH structures (see details in Supplementary Information). The coordinates and resonance assignment were deposited at the PDB and BMRB [73] databases under the identifiers 2MDX and 19508, respectively.

Molecular docking

Flexible docking of FAPP1-PH and micelle structures was carried out using an established protocol [74]. Developments incorporated here include the detection of buried ^1H , ^{13}C -group PREs from doxyl-PC and solvent-exposed groups from the gadolinium agent. Briefly, the active residues were defined based on micelle and bicelle-induced chemical shift changes and the solvent accessibility was estimated with NACCESS. Ambiguous distance restraints between the micelle centre and protein groups were set for atoms significantly broadened by 5-doxyl micelles. The docking protocol began with 400 randomly oriented and spatially separated protein and micelle structures taken from the representative ensembles.

Database linking

Structure of wild-type human FAPP1-PH domain sequence PDB ID **2MDX** and the corresponding BMRB entry **19508**.

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Figures

Figure 1: The gradients of lipids are shown for the anterograde pathway from the endoplasmic reticulum (ER) where many are synthesized, through the Golgi where they and proteins are modified, to the plasma membrane (PM), and would influence the degree of lipid disorder in each compartment. The relative amount of PI4P within the compartments is illustrated by red dots.

Figure 2: FAPP-PH binds specifically to liquid disordered (L_d) phases. (A) Liposomes composed of BSM, cholesterol and DOPC ratios were selected from different regions of the ternary lipid phase diagram, as shown on the right, and the amount of bound protein was quantified by gel imaging. Unless stated otherwise, all liposomes contained 2% (mol/mol) PI4P. Data from liposomes composed of DPPC, POPC, DOPC or a mixture of DPPC, DOPC and cholesterol (5:3:2) are plotted on the left. The percentage of liposome-bound FAPP1-PH is indicated for each lipid composition. The phase diagram of DOPC, BSM and cholesterol mixtures is shown on the right. The regions of liquid ordered and disordered phases are indicated as L_o and L_d , respectively. These two phases coexist in vesicles formed by the lipid

mixtures situated within the grey zone. The zone of transition between ordered and disordered vesicles is indicated in blue. **(B)** Disordered phases were monitored by the fluorescence signals of TR-DHPE and overlapped with FAPP1-PH^{green}. The bars in the confocal images represent 10 μm . **(C)** The PI4P ligand was included as either natural (brain extract) or synthetic (C_8 or C_{16}) forms in the vesicles formed by a 1:1:1 mixture of BSM:Chol:DOPC, and resulted in similar FAPP1-PH protein localization to the segregated disordered phase as revealed by the specific dyes TR-DHPE (L_d) and Oregon Green CTx (L_o) as seen by confocal microscopy.

Figure 3: NMR signal changes induced in FAPP1-PH by PI4P independent bilayer binding. A transient interaction is evident from significant CSPs consistently induced by interactions with bicellar **(A)** and micellar **(B)** systems as measured from ^{15}N -HSQC experiments in the absence of PI4P. This defined the active residues in HADDOCK calculations of the protein nonspecifically bound to the micelle. Backbone and side chain resonance changes are indicated by black and red bars, respectively. **(C)** Distance restraints used to calculate the structural model were taken from PREs using doxyl-broadened FAPP1-PH resonances. Those restraints from ^{15}N and ^{13}C HSQC spectra are shown above and below the x-axes. Residues with significantly perturbed resonances (mean + standard deviation) are indicated. The inset cartoons show the relevant complex of protein, 5-doxyl-PC, and micelle in black, purple and grey, respectively, for each panel.

Figure 4: Interaction of FAPP1-PH with PI4P assemblies. **(A)** The structural characterization of the specific complex was based on perturbations of FAPP1-PH resonances after addition of C_8 -PI4P to samples containing $^{15}\text{N}/^{13}\text{C}$ -labelled protein and either micelles or **(B)** bicelles. Chemical shift changes of ^{15}NH (top) and $^{13}\text{CH}_3$ (bottom) groups after addition of C_8 -PI4P to

samples containing 4mM DPC:CHAPS (3:1) micelles or 5% DH₆PC/DMPC bicelles are shown (B) as are PREs following addition of 5-doxyl PC-bound micelle versus PC-micelle controls, with the spin label shown as a red molecule in the cartoon. Black and red bars indicate backbone and sidechain perturbations, respectively. The dissociation constants of FAPP1-PH for C₈-PI4P containing micelles and bicelles are $5.3 \pm 2.4 \mu\text{M}$ and $8.8 \pm 3.3 \mu\text{M}$ respectively. Groups involved in micelle (D) and (E) bicelle binding based on CSPs and PREs upon PI4P addition are indicated as blue, green and red spheres in the structure for those groups with perturbed ¹⁵N backbone, ¹⁵N sidechain and ¹³C methyl signals, respectively. Significant PREs are represented by large spheres. The ¹³C-HSQC spectra of FAPP1-PH from ligand titrations are superimposed with peaks of key residues labeled (F). The trace for the micelle bound (red) and micelle/PI4P bound state (blue) are shown in the left panel for Lys74_ε sidechain resonance. Peak intensities for the diamagnetic and paramagnetic samples (red and yellow for micelle bound; blue and cyan for micelle/PI4P) are shown in the left panel.

Figure 5: The solvent-exposed surface of bilayer-bound FAPP1-PH mapped by PREs. (A) The gadodiamide (Gd³⁺)-induced PREs are expressed as a percentage of reduction of the signal intensity observed in paramagnetic vs diamagnetic samples. The intensity reduction of backbone amide signals of each residue of the free state of FAPP1-PH (top) and the bicelle-bound (middle) and bicelle:PI4P-complexed (bottom) forms are shown. The residues indicated in the lower panel and colored in blue exhibit a significant change in solvent accessibility upon formation of the bicelle:PI4P:FAPP1-PH complex. (B) The protection factors for ¹³C methyl groups of FAPP1-PH bound to PI4P-containing micelle are plotted. The color gradient indicates the degree of solvent protection. The y axis corresponds to the solvent accessibility calculated with Naccess for the free state of FAPP1-PH. Protected methyl groups are indicated with vertical lines. (C) The 1D traces were extracted from 2D ¹³C HSQC spectra (red: diamagnetic, blue:

paramagnetic) show the Met73C ϵ resonance of the free state, FAPP1-PH:micelle and FAPP1-PH:micelle:PI4P complexes. **(D)** Electrostatic maps of the FAPP1-PH structure calculated with APBS [75] showing the isocontours and superimposed with the surface accessible area (between -1 and 1 kT/e). FAPP1-PH is oriented as in Figure 4D.

Figure 6: FAPP1-PH membrane binding: **(A)** Comparison of pure and PI4P-containing micelle docked structural orientations. Back calculated distances (micelle center to protein backbone HN) for the 20 best models of micelle bound (grey) and PI4P bound (black) models are represented. Micelle and protein centers are indicated by purple dotted lines, the radial distribution ($g(r)$) of phosphorus (yellow) and nitrogen (blue) atoms of the choline headgroups are indicated by small spheres. The radial distributions were calculated from a 200 ps free molecular dynamic calculations in explicit water within Xplor. The structure closest to the mean is shown with the distance between the protein and micelle centers labeled, as well as the change in insertion angle upon PI4P binding by the micelle complex. The $\beta 1$ and $\beta 2$ strands are indicated. **(B)** A detailed model of FAPP1-PH showing the side chains (balls and sticks) is shown. The micelle surface (orange) and β strands are indicated as are the key interfacial residues. The hydrophobic wedge is delimited by Trp8 and Gln16, and is oriented such that $\beta 2$ is fully inserted into micelles while $\beta 1$ is interfacial. Residues with significant PREs are shown. **(C)** Mutations of membrane inserting residues reduce affinities for PI4P-containing membranes based on liposome sedimentation assays and surface plasmon resonance. The co-sedimented fraction and relative response is plotted for different FAPP1-PH concentrations.

Figure 7: Membrane binding model. The proposed states of FAPP proteins at the TGN involve nonspecific insertion into the loosely packed bilayer regions (green), diffusing laterally until a PI4P molecule (red) is encountered and bound, thus increasing the residency time such that

protein accumulates, and boosting lateral pressure in the bilayer. The proposed membrane recognition by the PH domain and Arf1 interaction could be extended to other COF family proteins to drive specific organelle targeting. Lipids transferred by COF proteins to the membrane increase the membrane rigidity and result in a progressive release of the PH domain of the COF proteins, thus halting the activity of COF proteins.

Table 1 NMR and refinement statistics for solution structures and micelle docked structures

Solution structure of FAPP1-PH	
NMR distance and dihedral constraints	
Distance constraints	
Total NOE	1448
Unambiguous	1223
Long range ($ i - j > 5$)	235
Medium ($4 \leq i - j \leq 5$)	41
Short ($2 \leq i - j \leq 3$)	83
Sequential / intra	864
Ambiguous	225
Hydrogen bonds	34
Total dihedral angle restraints	128
ϕ, ψ	64, 64
Structure statistics	
Violations	
Distance constraints (Å) (>0.5 Å)	0
Dihedral angle constraints (°)	0
Deviations from idealized geometry	
Bond lengths (Å)	0.006573 ±0.000167
Bond angles (°)	0.791 ±0.016
Improper (°)	1.777 ±0.089
Average pairwise r.m.s. deviation*(Å)	
Heavy , backbone	0.86, 0.32
Energies (Kcal.mol⁻¹)	
E_{noe}	442.77 ±10.85
E_{cdih}	6.25 ±1.63
E_{bond}	74.39 ±3.73
E_{improper}	109.62 ±8.94
E_{angle}	295.85 ±12.06
E_{vdw}	-185.74 ±26.89
E_{dihe}	582.8 ±7.53
Ramachandran statistics**	

Residues in core regions	73.86%	
Residues in allowed regions	22.33%	
Residues in generous regions	2.5%	
Residues in disallowed regions	1.4%	
Docked structures	Micelle	Micelle:PI4P
Intermolecular energies		
Buried surface (Å ²)	1208.22 ±125.87	1232.09 ±117.94
E _{vdw} (Kcal.mol ⁻¹)	-58.50 ±5.26	-59.29 ±6.05
E _{elec} (Kcal.mol ⁻¹)	-80.48 ±39.22	-122.72 ±33.94
Insertion		
θ (deg.)	16.80 ±4.50	14.23 ±4.32
ψ (deg.)	166.52 ±16.29	153.52 ±30.39
r (mass centers) (Å)	38.47 ±1.00	37.14 ±1.13
Interactions ^{***}	8, 10 , 11,12,13,15 , 16	8,10,11,12,13,15,16,74

* RMSD is calculated for secondary structural elements including residues 1-7,16-22,26-30,43-44,50-53,61-64,69-73 and 78-94 calculated for ten representative structures.

** non-glycine residues

*** more than 7 of the 20 models. Bold residues are hydrogen bonded between the headgroup of DPC and W8_ε, N10_δ, T13, W15_ε, Q16_ε and K74_ε.

Table 2: Experimental data used for deriving structural restraints to model the FAPP1-PH interaction with PI4P membrane mimicking micelles in the HADDOCK calculations based on an established approach [74].

Lipid model	NMR data	Nuclei	Resonance
Micelle (PI4P+DPC)	5-doxyl PC PRE	¹⁵ N	Trp8 _ε , Gly14, Q16 _ε Asn10, Asn10 _δ , Leu12, Thr13, Trp15, Trp15 _ε
		¹³ C	Leu12 _δ , Thr13 _γ , Lys74 _ε
	CSP	¹⁵ N	Leu5, Lys7-Gln16, Ile 44, Ile 63-Ile65, Glu68, His70, Phe72, Met73, Trp8 _ε
		¹³ C	Val4, Leu5, Leu12, Ile 44, Ile63-Ile65, Met73
Bicelles	Gd ³⁺ spin label	¹⁵ N	Thr9, Asn10, Thr13, Gly14, Ser53, Thr59, Glu62, Ile71, Lys74
		¹³ C	Leu12 _δ , Thr13 _γ
	CSP	¹⁵ N	Tyr6, Trp8, Trp8 _ε , N10, N10 _δ , Thr13, Lys27, Tyr29, Gly42, Glu50, Met61, Leu63, Ile64, Ile64, Ile71, Tyr72, Lys74
		¹³ C	L12 _δ , T13 _γ , Ile63 _β , Ile64 _γ
Loss of function by mutation			
Lys7A, Trp8E, Trp8Y, Asn10T, Tyr11G, Leu12G, Gln16R, Arg18L			
HADDOCK restraints			
Deeply inserted	Trp8 _ε , Asn10(N, _δ), Leu12(N, _δ), Thr13(N, _γ), Gly14N, Trp15(N, _ε), Gln16 _ε		
Interfacial	Lys74 _ε		

Table 3: Dissociation constants measured for FAPP1-PH membrane interactions

Protein	Dissociation constant estimated from	
	liposome sedimentation assays (μM)	SPR (μM)
WT	2.05 \pm 0.90	2.68 \pm 0.71
W8E	>1000	>1000
W8Y	>1000	>1000
N10T	5.12 \pm 1.07	17.32 \pm 5.74
T13F	5.1	n.d.*
T13N	5.86 \pm 0.83	5.15 \pm 0.54
W15Y	2.67 \pm 0.09	2.48 \pm 0.61
W15E	n.d.*	n.d.*
Q16R	>1000	>1000
R18L	>1000	>1000

*not determined due to the instability of the mutant

Figure 1

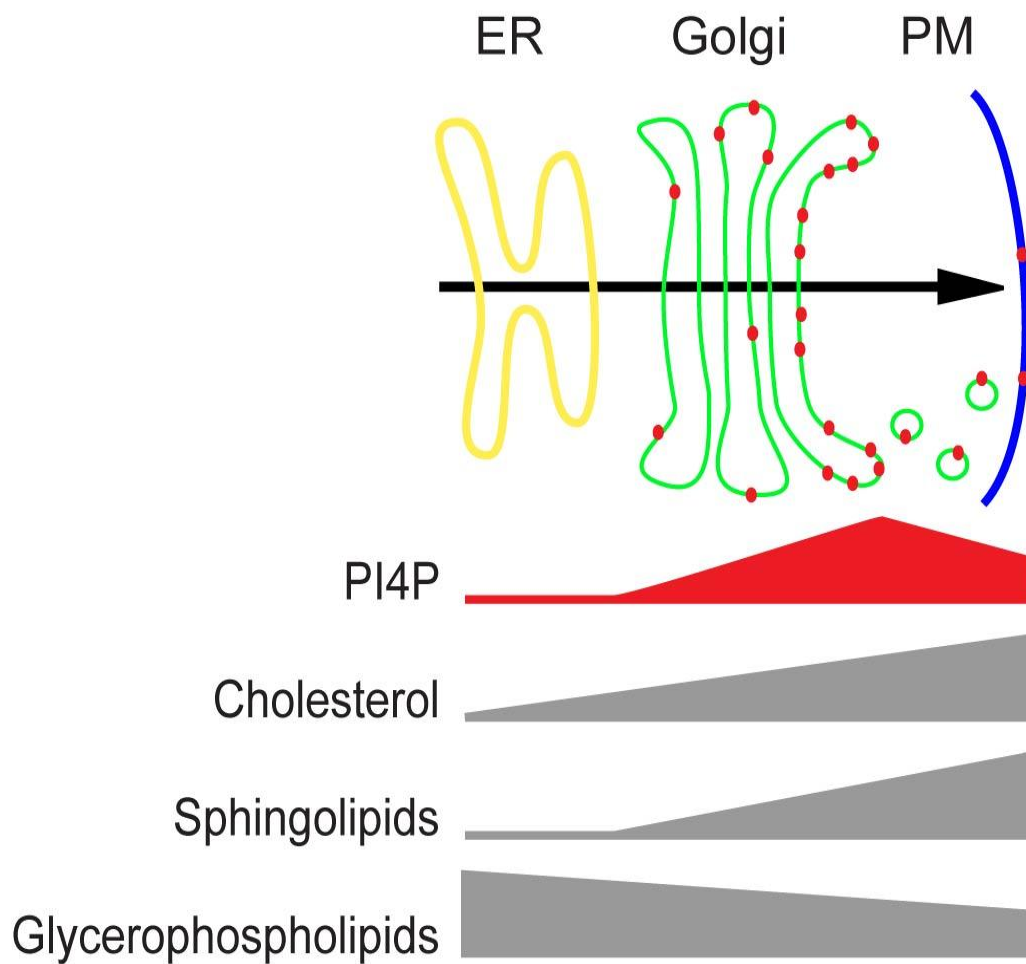


Figure 2

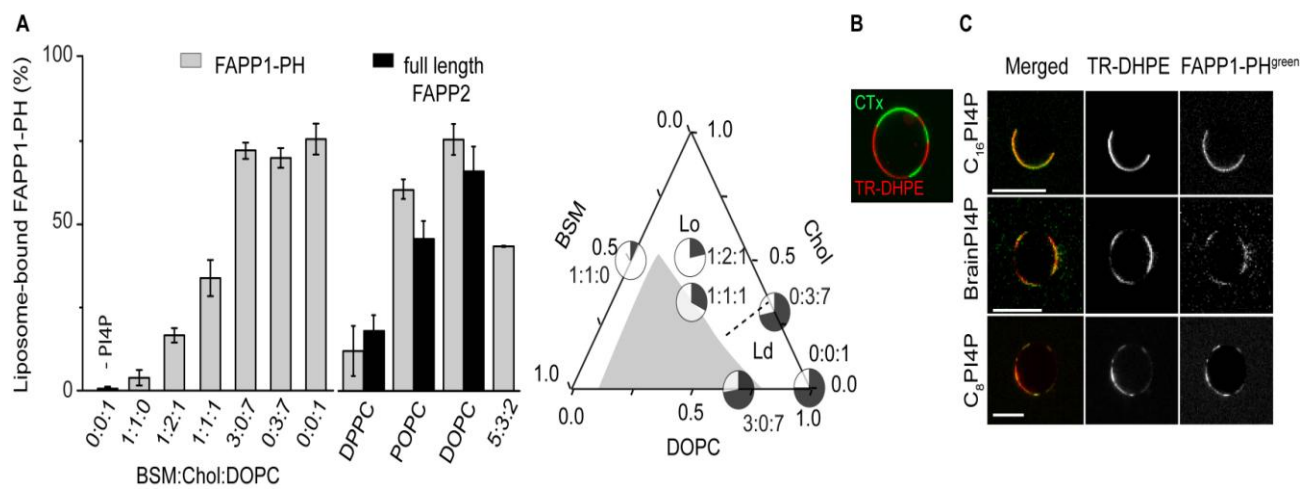


Figure 3

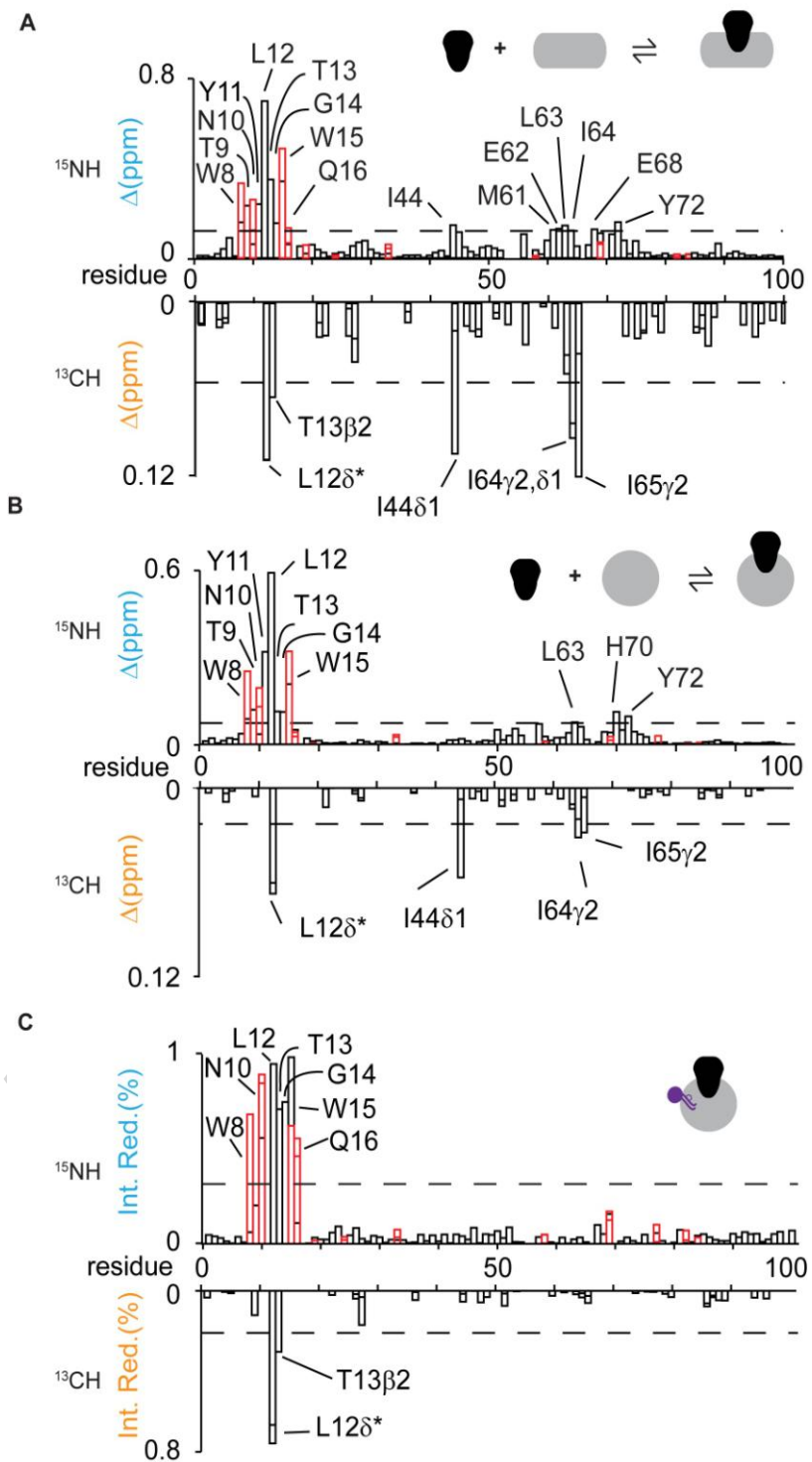


Figure 4

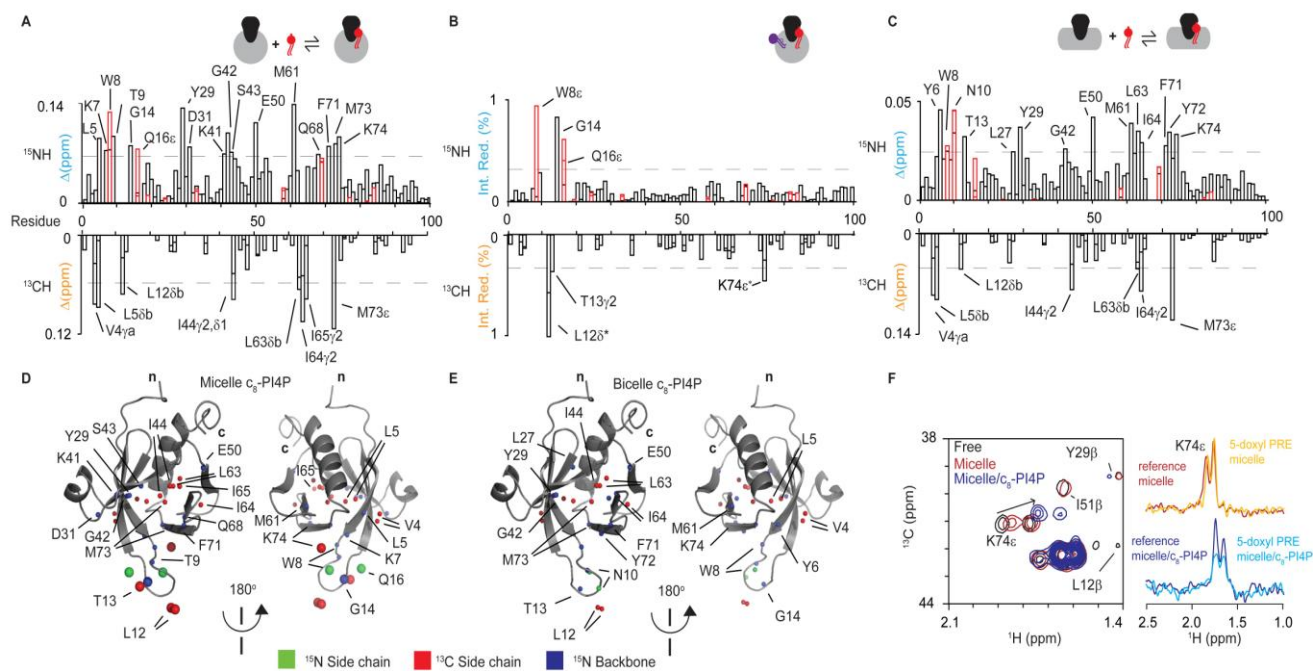


Figure 5

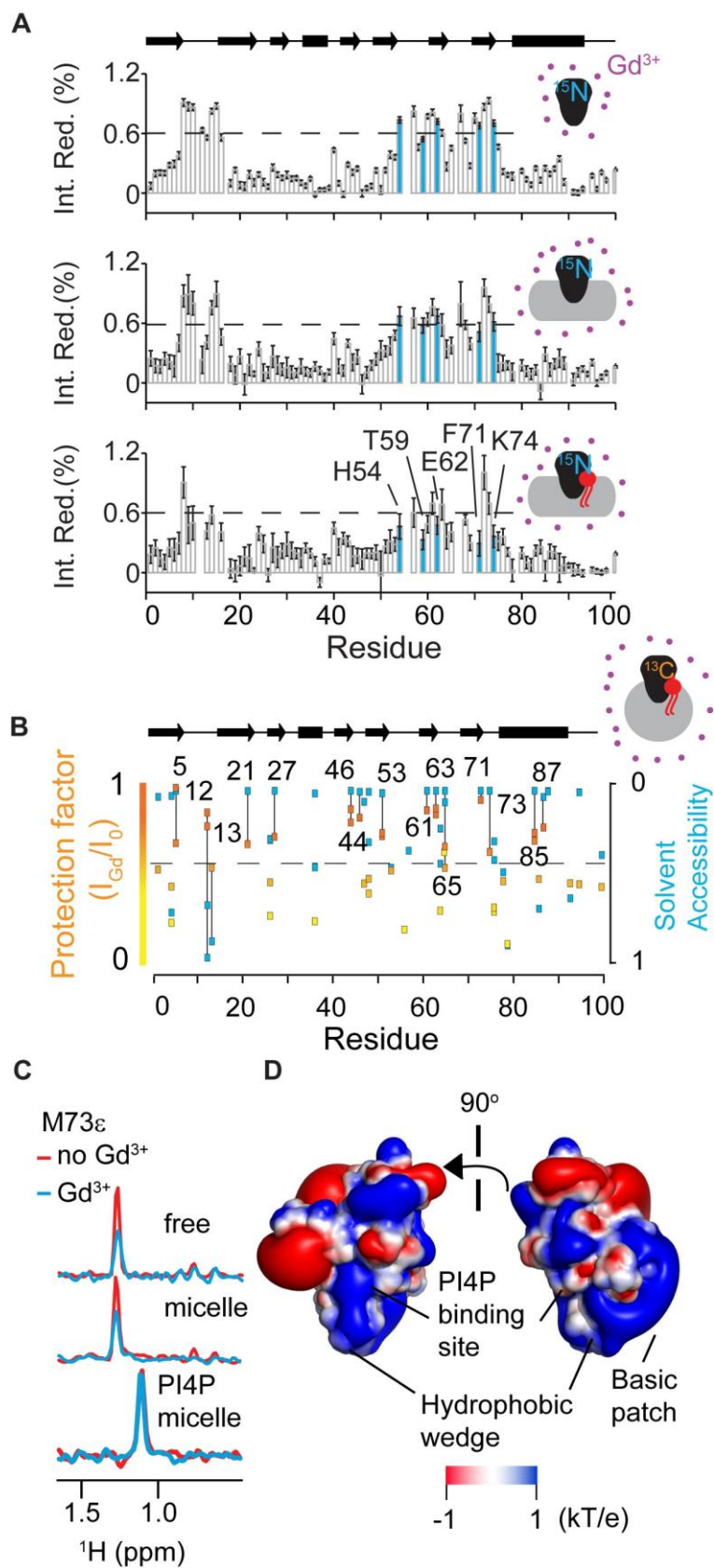


Figure 6

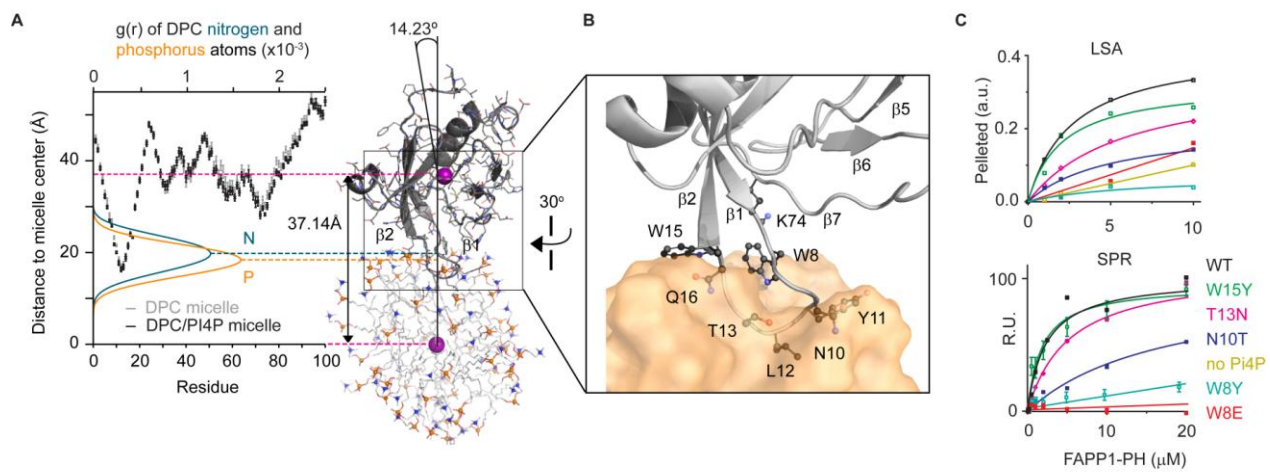
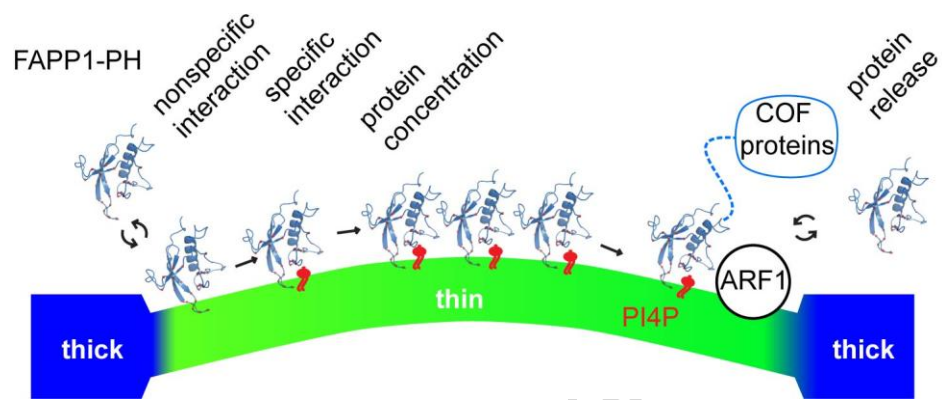
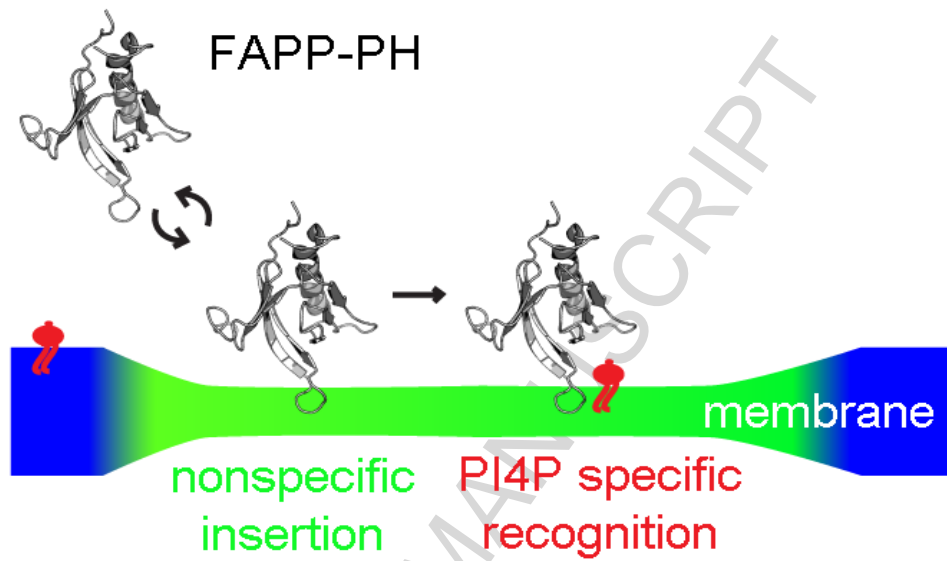


Figure 7





Graphical abstract

Highlights

- FAPP-PH domains selectively recognize PI4P within disordered membrane domains
- Two-pronged multistep binding mode mediates protein insertion into fluid bilayer
- Protein targeting to Golgi driven by disorder gradient created by lipid composition