Roles of Amphipathic Helices and the Bin/Amphiphysin/Rvs (BAR) Domain of Endophilin in Membrane Curvature Generation^{*S}

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Control of membrane curvature is required in many important cellular processes, including endocytosis and vesicular trafficking. Endophilin is a bin/amphiphysin/rvs (BAR) domain protein that induces vesicle formation by promotion of membrane curvature through membrane binding as a dimer. Using site-directed spin labeling and EPR spectroscopy, we show that the overall BAR domain structure of the rat endophilin A1 dimer determined crystallographically is maintained under predominantly vesiculating conditions. Spin-labeled side chains on the concave surface of the BAR domain do not penetrate into the acyl chain interior, indicating that the BAR domain interacts only peripherally with the surface of a curved bilayer. Using a combination of EPR data and computational refinement, we determined the structure of residues 63–86, a region that is disordered in the crystal structure of rat endophilin A1. Upon membrane binding, residues 63–75 in each subunit of the endophilin dimer form a slightly tilted, amphipathic α-helix that directly interacts with the membrane. In their predominant conformation, these helices are located orthogonal to the long axis of the BAR domain. In this conformation, the amphipathic helices are positioned to act as molecular wedges that induce membrane curvature along the concave surface of the BAR domain.

Biological membranes are subject to constant remodeling, and the control of membrane shape and curvature is essential for many vital cellular functions, such as cell division and motility, endocytosis, and vesicular trafficking (1). Recent work has demonstrated that these processes appear to be governed by proteins that can sense and induce membrane curvature (1, 2), and the molecular mechanisms through which these proteins act have become of interest.

Epsin and bin/amphiphysin/rvs (BAR) domain proteins are among the proteins thought to regulate membrane curvature in endocytosis (3–7). In vitro experiments show that these proteins can bind to larger vesicles and induce the formation of either small and highly curved vesicles (vesiculation) or narrow and highly curved tubules (3–6, 8). High resolution structural information has been obtained for the aqueous forms of all of these proteins (4–6, 8, 9) and provides a starting point for studying the molecular mechanisms of membrane curvature induction. The crystal structures of the BAR domains of endophilin and amphiphysin have the striking feature that they adopt curved, banana-shaped dimers, as illustrated for the structure of rat endophilin A1 (5) in Fig. 1A. The shapes of these dimers are complementary to that of the curved membranes with which the proteins interact. The concave surfaces of the dimers also have a high density of positively charged residues that are likely to interact favorably with negatively charged membranes, and multiple simultaneous mutations of these residues reduce the membrane interaction (5, 6). These data suggest that the concave surface of the BAR domain plays an important role in membrane curvature generation and that it might act as a rigid, positively charged scaffold (5, 6, 8, 10–16).

It has become clear that additional regions outside the BAR domain also play important roles in membrane curvature induction. These regions are unstructured in solution and are not resolved in crystal structures, but their removal or mutation can abolish or inhibit liposome binding activity and tubulation (5, 6, 8). Using site-directed spin labeling and EPR spectroscopy, we have shown that the endophilin N terminus undergoes a structural reorganization from an unfolded state in solution to an amphipathic helix (helix 0) that inserts into the membrane at the level of the head group (5). The inserted helix is likely to act as a molecular wedge that creates a strain in the outer leaflet and thereby promotes membrane curvature. Endophilin also has a central region (residues ~62 to 86) that is disordered in the crystal structure of the rat protein (5) but partly helical (residues 63 to 70) in the crystal structure of the human protein (8). This region has also been proposed to form a helix that inserts into the membrane (5, 17), but detailed experimental evidence is still lacking.

The abbreviations used are: BAR, bin/amphiphysin/rvs; NiEDDA, nickel ethylenediamine-N,N′-diacetic acid; SAMD, simulated annealing molecular dynamics; DEER, double electron-electron resonance.
Unreacted spin label was separated using PD10 columns. iosulfonate, which generates the spin-labeled side chain R1. gas and desiccated overnight. They were resuspended in 20 mM chased from Sigma-Aldrich. Lipids were dried with nitrogen

Single and double cysteine mutants of rat endophilin A1 were expressed and purified as described previously (5). Proteins recently been supported by genetic rescue experiments in Dro sophila (18).

Here, we investigated the roles of helix insertion and scaffolding by examining the structure of rat endophilin A1 upon vesiculation. Previous work on annexins showed that curvature-dependent membrane interactions can lead to major conformational reorganization of a protein (19). Thus, our first goal was to test whether the overall structure of the endophilin BAR domain is retained upon membrane interaction and, if so, how its concave surface interacts with the membrane. We then investigated the structure of the central insert (residues 64 – 86) in the membrane-bound protein using site-directed spin labeling and EPR spectroscopy. The information obtained was then used in a computational refinement to generate three-dimensional atomistic models of membrane-bound endophilin.

EXPERIMENTAL PROCEDURES

Preparation of Spin-labeled Rat Endophilin A1 Derivatives—Single and double cysteine mutants of rat endophilin A1 were expressed and purified as described previously (5). Proteins were reacted with 5× molar excess of the spin label MTSL (1-oxy-2,2,5,5-tetramethyl-d-pyrolone-3-methyl)-methanethiosulfonate, which generates the spin-labeled side chain R1. Unreacted spin label was separated using PD10 columns.

Liposome Preparation—Folch fraction I, type I, was purchased from Sigma-Aldrich. Lipids were dried with nitrogen gas and desiccated overnight. They were resuspended in 20 mM Hepes, pH 7.4, 150 mM NaCl, treated to brief bath sonication, and extruded through 400-nm pore filters using the Avanti mini-extruder.

Continuous Wave EPR Experiments—2 μM spin-labeled protein was combined with 1.4 mg/ml Folch liposomes and incubated at room temperature for 20 min. After incubation, unbound proteins were separated from membrane-bound proteins by high speed centrifugation at 152,800 × g for 20 min at 22 °C. EPR spectra were then obtained for the membrane-bound portion and also for samples in the absence of liposomes. Spectra were collected using a Bruker EMX spectrometer fitted with a dielectric resonator at 1.59-mW incident microwave power and a scan width of 100 gauss. Power saturation experiments to determine O2 and NiEDDA accessibility (O2 and NiEDDA, respectively) have previously been described (20). Accessibility data were obtained using oxygen from air in equilibrium with buffer, and 10 mM NiEDDA. The Φ parameter was calculated by the relationship Φ = ln[O2]/[O2/NiEDDA]). The immersion depth was calibrated using 1-palmitoyl-2-stearoyl-(n-DOXYL)-sn-glycero-3-phosphocholine (Avanti Polar Lipids) as described previously (20). We obtained the following relation between immersion depth (d) and Φ: d[A] = 6.3*Φ – 3.9. This immersion depth represents the depth of the nitroxide moiety.

Pulsed EPR and Distance Analysis—Samples were prepared by combining 4 μM protein and 1.4 mg/ml Folch liposomes. After incubation for 20 min at room temperature, samples were centrifuged at 152,800 × g for 20 min, and the pellet was recovered for measurement. The pellet was resuspended in buffer, and 20–25% sucrose was used as cryoprotectant before loading into quartz capillaries. Samples were flash frozen, and data were acquired at 78 K. Four-pulse double electron-electron resonance (DEER) experiments were performed using a Bruker Elexys E580 X-band pulse EPR spectrometer fitted with a 3-mm split ring (MS-3) resonator, a continuous flow helium cryostat (CF935; Oxford Instruments), and a temperature controller (ITC503S; Oxford Instruments). Data were fit using gaussian distributions, as implemented in the DEERAnalysis2008 package (21).

Structural Refinement—The crystal structure of the rat endophilin dimer (Protein Data Bank code 2C08) (5) was used as a starting point for computational refinement based on distance and depth constraints derived from EPR data. A complete description of the details of this procedure is given in the supplemental material. An in-house algorithm (PRONOX) was used to add residues 64 – 86 to each subunit of the endophilin dimer: add spin labels at positions 63, 66, 70, 73, 74, 75, 77, and 83; locate the starting structure at the surface of an imaginary lipid vesicle; and produce a set of constraints for simulated annealing molecular dynamics (SAMD) calculations. Based on the secondary structure determined by EPR, residues 64 – 75 were built as helices and constrained using hydrogen-bonding constraints to maintain a helical structure. These constraints, interdimer distance constraints measured from continuous wave EPR and four-pulse DEER EPR, lipid depths for residues 63, 66, and 70, and other system constraints (supplemental Tables S1 and S2) were used in SAMD simulations in AMBER8 (22). During these calculations, the helix and loop were not covalently linked to each other or to the BAR domain to allow
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TABLE 1

| Mutant     | EPR distance | Crystal structure |
|------------|--------------|-------------------|
|            | A            | A                 |
| 227R1      | 9 (CW)       | 8                 |
| 63R1       | 9 (CW)       | 11                |
| 96R1       | 38 (DEER)    | 36                |
| 216R1      | 37 (DEER)    | 34                |
| 166R1-210R1| 43 (DEER)    | 35                |
| 178R1-200R1| 31 (DEER)    | 33                |

Distances obtained from continuous wave (CW) or four-pulsed DEER experiments (DEER) were fit using gaussian models (Fig. 1 and supplemental Fig. S1). All distances were obtained for the indicated endophilin derivatives bound to membrane. The experimental distances are given as the peak of the gaussian distributions and are in good agreement with distances expected from the crystal structure which was taken as the α-carbon distance between the indicated residues.

Results

Rat Endophilin A1-BAR Is a Dimer upon Membrane Interaction—To investigate whether the BAR domain of the rat endophilin A1 dimer observed crystallographically is retained upon membrane interaction, we generated a series of spin-labeled endophilin derivatives (Fig. 1A) and incubated them with liposomes comprised of Folch lipids. The experimental conditions were chosen such that the Folch liposomes were predominantly vesiculated into highly curved vesicles with an average diameter of ~230 Å, as determined by electron microscopy (supplemental Fig. S1A). We then measured intra- and intersubunit distances and compared them with the respective α-carbon distances of the crystal dimer. The distances measured by EPR are those between the nitroxide moieties of R1, and therefore, they would be expected to be similar but not necessarily identical to the α-carbon distances. Distances >20 Å were investigated using four-pulse DEER (24–27), whereas shorter distances were investigated using continuous wave EPR (28, 29) (see “Experimental Procedures”).

The DEER measurements are illustrated with the example of membrane-bound endophilin spin-labeled in helix 3 at position 216 (Fig. 1A). A pronounced periodic oscillation is observed for this 216R1 derivative (Fig. 1B), and frequency analysis of this oscillation yields a well defined distance distribution with a maximum at 37 Å (Fig. 1C). This distance is in good agreement with the respective intersubunit α-carbon distance of 34 Å in the crystal structure of endophilin (Table 1). Similarly, the intradimer distances of membrane-bound 96R1 (in helix 1b), 63R1 (in the helix 1 insert), and the previously reported (5) distance for 227R1 (in helix 3) are in good agreement with the crystal structure (Table 1 and supplemental Fig. S1, B and C). As a control, we also measured the intersubunit distance for one of the derivatives (96R1) in solution and obtained the same distance as that for the membrane-bound form (supplemental Fig. S1D).

The tip-to-tip distance of the endophilin dimer in the crystal is nearly 130 Å. Because this range is beyond the detection limit for intersubunit contacts, we employed a different approach and measured distances between two sites located toward the tip region in the same subunit. As shown in Table 1 and supplemental Fig. S1B, the distance of 31 Å between 178R1 and 200R1 (located directly at the tip and in helix 3, respectively) is also in good agreement with the α-carbon distances of the crystal structure (33 Å). It has been found previously that the structures of different families of BAR domain proteins overlay well in the central region but are much more divergent toward the tips or wings (15). To test for potential movement between the central region and the wing region, we measured the distance between 166R1 and 210R1 (located in helix 2 and 3, respectively). The distance between these residues for membrane-bound endophilin is in good agreement with the expected distance from the crystal structure (Table 1 and supplemental Fig. S1B). Moreover, the same distance was obtained when distances were measured between these residues in solution (supplemental Fig. S1D). Collectively, this distance analysis supports the notion that the overall structure of the BAR domain dimer is largely retained upon membrane interaction.

Concave Surface of the BAR Domain Does Not Penetrate Deeply into the Acyl Chain Interior of the Membrane—Next, we used mobility and accessibility analysis of selected spin-labeled derivatives to investigate the mechanisms by which the BAR domain interacts with the membrane. Spin labels were introduced at positions that were either located on the concave surface of the BAR domain (positions 96, 159, and 166) or at positions expected to face away from the membrane (positions 108, 172, and 247) (Fig. 1A).

The EPR spectra for these derivatives in the soluble (black trace, Fig. 2) and membrane-bound forms (red trace, Fig. 2) are qualitatively similar and consistent with the respective locations in the crystal structure. For example, the EPR spectra for 247R1 (Fig. 2B) are characterized by three sharp and narrowly spaced lines in both conditions. The high motion indicated by these spectra is consistent with the location of this site at the C

![FIGURE 2. EPR analysis of sites in the BAR domain. Continuous wave X-band EPR spectra of soluble (black) and membrane-bound (red) endophilin derivatives spin-labeled at the indicated positions (A) or facing away from the concave surface (B). All spectra were normalized to the same number of spins. Spectra for 247R1 were of much higher intensity and, therefore, were shown at 2.5-fold reduced intensity. The scan width is 100 gauss. The depth parameter (ϕ) values for all derivatives in membrane-bound form are strongly negative (less than −1), indicating that the respective spin-labeled sites do not significantly penetrate into the hydrocarbon layer of the membrane.](image)
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Although these data clearly indicate the lack of significant acyl chain penetration of the nitroxide moiety of the spin-labeled derivative (Fig. 3, black) and membrane-bound (red) endophilin derivatives are shown. The line broadening and loss of spectral amplitude of all sites upon membrane interaction reflect, to a significant extent, a reduction in mobility and ordering. Additional broadening from spin-spin interaction between the same sites only has minor spectral contributions because the labeled sites are relatively far apart (also see discussion of Table 2). Scan width is 100 gauss.

To investigate the proximity of the labeled sites to the membrane bilayer, we used accessibility of the labels to the paramagnetic colliders, O2 and NiEDDA. Although the concentration of the nonpolar O2 increases in the membrane, the more polar NiEDDA preferentially partitions into the solvent. Thus, the deeper a spin label penetrates into the acyl chain region, the more it becomes accessible to O2 and inaccessible to NiEDDA. The accessibilities to O2 and NiEDDA (II02 and IIINiEDDA) are conveniently summarized by the depth parameter $\Phi$ ($\Phi = \ln(\text{II02}/\text{IIINiEDDA})$). Interestingly, the $\Phi$ values for positions 96, 159, and 166 (on the concave face) or 108, 172, and 247 (toward the convex face) were all in the range from $-1.2$ to $-2.1$ (Fig. 2). Thus, the depth parameter $\Phi$ indicates little membrane penetration by residues on the concave surface of the BAR because the values of $\Phi$ do not differ between labels at three such positions and those at solvent-exposed positions. Although these data clearly indicate the lack of significant acyl chain penetration of the nitroxide moiety of the spin-labeled sites, they do not exclude the possibility that the concave surface of endophilin interacts with the interfacial region of the bilayer (see below).

Central Insert Region Becomes an Amphipathic Helix upon Membrane Interaction—To investigate potential membrane-induced conformational changes in the helix insert region, we conducted a nitroxide scanning experiment. The EPR spectra of the soluble derivatives (Fig. 3, black) are generally of higher amplitude with sharper spectral lines than those from the respective membrane-bound forms (Fig. 3, red). Control experiments, in which sucrose was used to slow down the tumbling of the dimer in solution, showed little spectral changes, indicating that the membrane binding-induced change in tumbling rate did not play a significant spectral role (data not shown). The line shapes of 70R1 to 77R1 indicate a largely disordered and unfolded structure in solution, whereas a more ordered structure is induced upon membrane binding. Only residues 64–69 are ordered in solution, and pronounced immobilization can be seen for 64R1, 65R1, 68R1, and 69R1. Such immobilization could be caused by tertiary contacts. This region was largely disordered in the crystal structure of rat endophilin (5) but was resolved in the structure of human endophilin (8), in which residues 63–76 are found to form a short helical segment. Moreover, residues 64, 68, and 69 are pointing directly toward the BAR domain in the latter structure. Thus, the EPR spectra in the absence of membranes are consistent with a structure akin to that in the crystal of human endophilin (also see below). Residues 64–69 are also the most ordered region of the membrane-bound form, but in the presence of membranes the ordered region becomes much more extended.

To obtain more detailed secondary structure information in the membrane-bound form, we performed O2 and NiEDDA accessibility measurements for 63R1 to 77R1. As summarized with the depth parameter $\Phi$ (Fig. 4A), the accessibility data give rise to a periodic oscillation. The local $\Phi$-maxima (residues 63, 66, 70, 73, and 74; magenta) represent preferentially O2 accessible, lipid-exposed sites, and these positions cluster on the hydrophobic face of a helical wheel (Fig. 4B). Conversely, the local minima (green) correspond to solvent-exposed sites that fall onto the opposite face of the helical wheel. The formation of an amphipathic $\alpha$-helical structure is further supported by a comparison of the $\Phi$-data with a cosine curve that has the standard periodicity of an $\alpha$-helix (3.6 amino acids/turn). This curve agrees well with the experimentally observed periodicity from residues 63 to 75, but itstarts to diverge for residues 76
and 77. For these residues, an increase in the \( \Phi \)-values would be predicted, but the opposite is observed. Thus, these residues may no longer be part of the helix.

Previous studies demonstrated that \( \Phi \)-values are proportional to the membrane immersion depth of R1 side chains that are penetrating into the acyl chain region. This depth can be calibrated using spin-labeled derivatives of phospholipids (20, 30). Based on such a calibration (see “Experimental Procedures”), we obtain average immersion depths of the nitroxide moieties of 63R1, 66R1, and 70R1 that are on the order of 5–6 Å. Previous work has shown that an R1 side chain is typically at a distance of 7–10 Å from the center of an \( \alpha \)-helix to which it is attached (31). Thus, we can estimate that the center of the helix is located about 1–4 Å above the phosphate layer. The \( \Phi \)-values for residues 73 and 74 are somewhat lower, and the reduced immersion depth of these sites is consistent with a slight helix tilt. However, residues 73 and 74 are also located on the periphery of the hydrophobic face (Fig. 4B) where the side chains might project laterally rather than directly toward the bilayer interior.

**Dimeric Amphipathic Helices in the Central Insert Region Are Antiparallel**—To analyze the orientation of the amphipathic helices with respect to each other, we measured intersubunit distances between the same sites in each subunit using DEER (supplemental Fig. S2). As shown in Table 2, the distance between these sites generally increases with increasing residue number, indicating that the helices must be facing away from each other. This feature is best illustrated with the lipid-exposed sites 63R1, 66R1, and 70R1. For these sites, we can assume that the nitroxide side chains are facing in the same direction (i.e., into the membrane) and that the interlabel distances are likely to be close to those between the respective \( \alpha \)-carbon atoms. An interesting feature of the DEER distances is that they increase by about 10 Å for each helical turn, which is in good agreement with antiparallel helices because an individual helix is extended by \( \sim 5 \) Å/turn. The other distances in Table 2 are also in agreement with a helical structure (also see below).

Since Masuda and colleagues (8) observed a short antiparallel helical structure for residues 63–71 in the crystal structure of human endophilin A1, we sought to determine whether the distances in solution are similar to those in the membrane-bound form. As shown in supplemental Table S3, the intersubunit distances for 63R1, 67R1, and 70R1 in the absence of membranes differ from those observed in the membrane-bound state. Compared with the \( \alpha \)-carbon distances from the crystal structure, the measured distances in solution are in good agreement. These data suggest that the structure of the insert taken up in the crystal of human endophilin A1 is similar to that of rat endophilin A1 in solution and that the structure taken up in the membrane-bound state is different.

**Structural Refinement of the Central Insert Suggests Staggered Helices with High Mobility**—To generate atomistic models of the membrane-bound form of endophilin, we used the crystal structure of the soluble rat endophilin A1 dimer as a starting structure and built and refined the structure of the insert region (residues 63–86) using a modified version of our previously reported (23) SAMD refinement constrained by the EPR results. These calculations included intersubunit constraints based on DEER distances, depth constraints based on EPR accessibility data, constraints to maintain the helicity of the insert region as detected by EPR mobility and accessibility data, and symmetry constraints on the insert regions of the two subunits. The intersubunit constraints were implemented as the mean distance \( \pm 5 \) Å to reflect the relatively large width of the EPR distance distributions.

Endophilin structures were collected at the end of each of 60 SAMD cycles. The insert helices clearly adopted one of three positions with an angle between the helical axis and the long axis of the BAR domain of about 140°, 90°, and 40°, respectively (supplemental Fig. S3), with 18, 33, and 9 occurrences, respectively, over the 60 SAMD cycles. The three positions of the insert helices are interconvertible, and all may occur in a population of membrane-bound endophilin proteins. This may account for the width of the distance distribution for the intersubunit DEER data. However, in the SAMD cycles shown in supplemental Fig. S3A and in other calculations performed with minor changes in parameters (see the supplemental material), we consistently find that the most common conformation is that with the insert helices orthogonal to the BAR domain axis (green helices in supplemental Fig. S3B).

To examine the interaction of the endophilin dimer with a curved lipid surface, we used a representative structure with orthogonal insert helices and constructed part of a lipid vesicle around the protein (Fig. 5). We were able to do this because the predicted structure is positioned in space based on depth constraints relating the center of the lipid vesicle to the protein. Two main findings for the insert helices emerge from the SAMD calculations and subsequent building of the lipid surface. First, the helices (residues 63–75) are tilted such that the N terminus of each helix is located below the phosphate of the lipid head group, whereas the C terminus and the loop (residues 76–86) are positioned above the lipid surface (Fig. 5B). Second, the axes of the helices adopt a staggered position with respect to each other (Fig. 5A).

The detailed positions of amino acids 63–75 are shown in Fig. 5C. The amino acid positions obtained by computational refinement using EPR distance, depth and mobility data, and geometrical and symmetry constraints show remarkably good agreement with the predicted orientations based on ac-
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In this study, we investigated the structure of endophilin bound to small and highly curved vesicles. Under these conditions, the BAR domain of endophilin retains a dimeric structure similar to that found in the crystal. This demonstrates that endophilin does not undergo major conformational refolding upon membrane binding, unlike the case of annexin B12 (19). Moreover, we find that a central insert region (residues 63–75) becomes helical and inserts into the membrane at an immersion depth that is comparable (albeit slightly shallower) to that of the N-terminal helix (5). The insert helix in each subunit is tilted such that the side chains of Ala-63 and Ala-66 at the N terminus are below the level of the phosphate head groups of the lipid. The position of this helix with respect to the BAR domain is flexible, but seems to adopt a preferential position with the helical axes orthogonal to the BAR domain long axis.

The width of the DEER distance distributions for intersubunit distances between residues in the insert helices suggested some structural heterogeneity. This manifested in the computational refinement as three distinct positions with angles of about 140°, 90°, and 40° between the helical axis and the BAR domain. Interestingly, the 40° position is very close to the position adopted by the insert helices in the x-ray structure of human endophilin (8). In this crystal structure, Arg-65 in the insert helix forms a salt bridge with Glu-56, and a similar salt bridge forms in the 40° structures obtained in our SAMD refinement (supplemental Fig. S5A). In an analogous manner, Lys-67 in the insert helix forms a salt bridge with Glu-56 (in the opposite subunit) in the 140° structures generated in the SAMD calculation (supplemental Fig. S5B), whereas the salt bridge between Arg-65 and Glu-56 (same subunit) is broken. In the orthogonal structures, neither Arg-65 nor Lys-67 is located close enough to Glu-56 in the same or the opposite subunit, and the positive charges are therefore free to interact with the lipid head groups (supplemental Fig. S5C). Thus, a picture emerges of a dynamic helix that might occupy a range of conformations based on these electrostatically favorable interactions. It appears that the orthogonal orientation should be favorable in the membrane-bound state, but the other orientations of the helix might be accessible to adjust to different membrane and curvature conditions. This behavior and the detailed orientations of the side chains of the helices may be amenable to analysis by theoretical calculations along the lines of those performed previously for membrane-bound endophilin (17).

Two mechanisms have been suggested by which insertion of amphipathic helices can induce curvature. By inserting helices into only the outer leaflet of the bilayer, an imbalance is created between the two leaflets, and the tighter packing in the outer leaflet causes an expansion that, in turn, leads to bending of the membrane. In addition, insertion of helices at the level of the head groups could induce additional curvature strains by preferentially increasing the packing density in the head group rather than the acyl chain region (32). In the case of membrane-inserted amphipathic helices, these mechanisms are expected to promote membrane curvature in a direction perpendicular to the membrane plane. This behavior and the detailed orientations of the side chains of the helices may be amenable to analysis by theoretical calculations along the lines of those performed previously for membrane-bound endophilin (17).
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to that of the helix axis. Thus, the antiparallel insert helices in their most predominant orientation are ideally positioned to induce curvature along the concave surface of the BAR domain.

In contrast, the Φ-values for sites labeled on the concave surface of the BAR domain did not indicate penetration into the acyl chain region, and the endophilin BAR domain must, therefore, be positioned farther away from the membrane than its amphipathic helices. Thus, if the concave surface of endophilin interacts with the bilayer, it likely does so by interaction with the more distal portions of the head group. This general behavior is consistent with the model in Fig. 5, in which the BAR domain is docked onto a lipid vesicle according to the immersion depth information obtained for the insert helix. In this model, the BAR domain follows the general shape of the vesicle and is positioned to interact with the outer region of the membrane head groups. Future studies will have to show whether the structural features found under the present lipid conditions will also apply to vesicles of different sizes or to tubules.

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