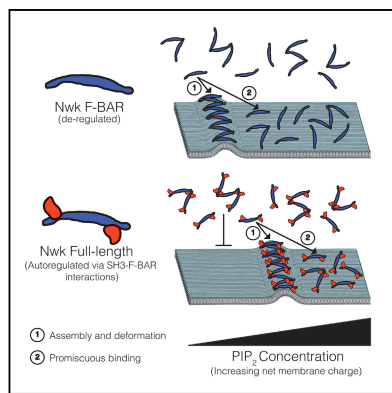
Cell Reports

Membrane Charge Directs the Outcome of F-BAR **Domain Lipid Binding and Autoregulation**

Graphical Abstract



Authors

Charlotte F. Kelley, Emily M. Messelaar, Tania L. Eskin, ..., Olga S. Sokolova, Daniela Nicastro, Avital A. Rodal

Correspondence

arodal@brandeis.edu

In Brief

Membrane dynamics are critical to many cellular processes, but the molecular mechanisms that regulate remodeling events are not well understood. Here, Kelley et al. demonstrate that for the F-BAR protein Nervous Wreck, intramolecular autoregulation and membrane charge work together to restrict remodeling to a limited range of lipid compositions.

Highlights

- Nwk F-BAR membrane binding is autoregulated by SH3 domains in vivo and in vitro
- Nwk binds membrane in an active assembled state or an inactive unassembled state
- High PIP₂ inhibits F-BAR membrane bending and favors promiscuous binding
- Autoregulation increases the charge needed for Nwk F-BAR membrane binding





Membrane Charge Directs the Outcome of F-BAR Domain Lipid Binding and Autoregulation

Charlotte F. Kelley,¹ Emily M. Messelaar,¹ Tania L. Eskin,¹ Shiyu Wang,¹ Kangkang Song,² Kalanit Vishnia,³ Agata N. Becalska,¹ Oleg Shupliakov,⁴ Michael F. Hagan,⁵ Dganit Danino,³ Olga S. Sokolova,⁶ Daniela Nicastro,² and Avital A. Rodal^{1,*}

¹Rosenstiel Basic Medical Sciences Research Center, Department of Biology, Brandeis University, Waltham, MA 02453, USA

²Departments of Cell Biology and Biophysics, UT Southwestern Medical Center, Dallas, TX 75390, USA

³Department of Biotechnology and Food Engineering, Technion - Israel Institute of Technology, Haifa 32000, Israel

⁴Department of Neuroscience, Karolinska Institutet, von Eulers väg 3, 171 77 Stockholm, Sweden

⁵Martin Fisher School of Physics, Brandeis University, Waltham, MA 02453, USA

⁶Department of Bioengineering, Faculty of Biology, M.V. Lomonosov Moscow State University, 119991 Moscow, Russia *Correspondence: arodal@brandeis.edu

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SUMMARY

F-BAR domain proteins regulate and sense membrane curvature by interacting with negatively charged phospholipids and assembling into higherorder scaffolds. However, regulatory mechanisms controlling these interactions are poorly understood. Here, we show that *Drosophila* Nervous Wreck (Nwk) is autoregulated by a C-terminal SH3 domain module that interacts directly with its F-BAR domain. Surprisingly, this autoregulation does not mediate a simple "on-off" switch for membrane remodeling. Instead, the isolated Nwk F-BAR domain efficiently assembles into higher-order structures and deforms membranes only within a limited range of negative membrane charge, and autoregulation elevates this range. Thus, autoregulation could either reduce membrane binding or promote higher-order assembly, depending on local cellular membrane composition. Our findings uncover an unexpected mechanism by which lipid composition directs membrane remodeling.

INTRODUCTION

Membrane remodeling is critical for cellular processes such as cargo trafficking, signaling, cell motility, and organelle biogenesis, and it requires the concerted action of scores of proteins that bind and actively shape cellular membranes. However, we still do not understand the underlying mechanisms that spatially and temporally target the activities of these proteins within the cell. BAR (Bin/Amphiphysin/Rvs) domain family proteins are important mediators of cellular membrane remodeling and form banana-shaped α -helical dimers that use a positively charged binding surface to interact with negatively charged membrane phospholipids (McMahon and Boucrot, 2015). Further, they assemble into stable higher-order scaffolds that

induce or stabilize membrane curvature over hundreds of nanometers (Frost et al., 2008; Mim et al., 2012; Becalska et al., 2013; Zhao et al., 2013). While it remains uncertain whether BAR domains actively generate or stabilize membrane curvature in vivo, in either case, these activities require tight cellular regulation. However, the molecular mechanisms that regulate and target these proteins have only begun to be explored (Roberts-Galbraith and Gould, 2010).

Many BAR-domain proteins possess Src-homology 3 (SH3) domains that mediate interactions with regulators of the actin cytoskeleton and other membrane-remodeling proteins such as dynamin (Owen et al., 1998; Takei et al., 1999; Itoh et al., 2005; Neumann and Schmid, 2013). Recently, SH3 domains have been implicated in directly regulating BAR domains via intramolecular interactions (Wang et al., 2009; Guerrier et al., 2009; Rao et al., 2010; Vázquez et al., 2013; Meinecke et al., 2013; Kast et al., 2014; Chen et al., 2014; Wu and Baumgart, 2014). In general, SH3 domain interactions inhibit the ability of BAR domains to induce membrane remodeling when overexpressed in cells and in vitro; however, only limited evidence exists for the role of SH3-mediated autoregulation of BAR domains in vivo (Kumar et al., 2009; Guerrier et al., 2009). Further, little is known about how SH3 domain autoregulation affects different steps of BAR domain membrane binding, higher-order assembly, and deformation.

Nervous wreck (nwk) encodes a protein containing an N-terminal Fes/Cip4 homology-BAR (F-BAR) domain and two SH3 domains. Nwk regulates the traffic and signaling output of synaptic growth receptors at the *Drosophila* neuromuscular junction (NMJ) through its interactions with the membrane, actin nucleation machinery, and other endocytic proteins including dynamin, Dap160/Intersectin, and Sorting Nexin 16 (Rodal et al., 2008; O'Connor-Giles et al., 2008; Rodal et al., 2011). These activities and interactions define a recycling route by which activated receptors can be removed from signal-permissive early endosomes, downregulating their activities and controlling synaptic growth (Rodal et al., 2011). Mammalian Nwk homologs have been implicated in membrane remodeling and receptor traffic in stereocilia and in cerebellar granule neurons (Cao et al., 2013; Sun et al., 2015). Unraveling these trafficking pathways requires a deeper understanding of the membranedeforming activities of Nwk. We previously reported that the F-BAR domain of Nwk binds to negatively charged phospholipids, similar to canonical F-BAR domains. The Nwk F-BAR self-assembles into zigzags, distinct from canonical F-BAR proteins, and thus induces membrane scallops and ridges rather than membrane tubules (Becalska et al., 2013). Here, we describe regulatory mechanisms that direct the membrane remodeling activity of the Nwk F-BAR domain in the context of the full-length protein.

RESULTS

The F-BAR Domain and C Terminus of Nwk Are Required for In Vivo Localization and Function

To investigate the importance of Nwk membrane-remodeling activity in vivo, we tested the role of the F-BAR domain in Nwk localization and synaptic growth at the Drosophila NMJ. We expressed EGFP-tagged Nwk variants (full-length Nwk [Nwk-EGFP], Nwk lacking its F-BAR domain [Nwk $^{\Delta 1-428}$ -EGFP, and the Nwk F-BAR domain alone [Nwk¹⁻⁴²⁸-EGFP]) in the nwk null mutant background (Figures 1A, 1B, S1A, and S1B). As has been previously reported for the nwk null and other endocytic traffic mutants (Coyle et al., 2004; Dickman et al., 2006), nwk mutants expressing GFP alone exhibit an increase in total synaptic bouton number at the NMJ, as well as an increase in "satellite" boutons that bud off the main axis of the axon terminal (arrows, Figure 1A). This phenotype is a result of excess growth factor signaling in the absence of normal membrane recycling pathways (Rodal et al., 2008; O'Connor-Giles et al., 2008). Nwk-EGFP expression rescued the synaptic overgrowth and satellite bouton phenotypes of *nwk* mutants, although Nwk^{$\Delta 1-428$}-EGFP exhibited no rescuing activity (Figure 1B). In contrast, Nwk1-428-EGFP exhibited a more severe synaptic overgrowth and satellite bouton phenotype than the nwk null mutant (Figure 1B). These data indicate that the C terminus of Nwk is required to properly regulate its function.

To investigate the basis of the phenotypes of Nwk¹⁻⁴²⁸-EGFP at the NMJ, we examined the localization of Nwk variants in fixed tissue in the nervous system, where endogenous Nwk is expressed (Coyle et al., 2004). At NMJs and in cell bodies in the ventral ganglion, Nwk-EGFP localized in a similar pattern to that previously reported for endogenous Nwk, in occasional puncta and in the "periactive zone" region surrounding Bruchpilot (BRP)-labeled active zones (Coyle et al., 2004). In contrast, $\mathsf{Nwk}^{\Delta 1-428}\text{-}\mathsf{EGFP}$ exhibited a diffuse cytoplasmic localization, and Nwk1-428-EGFP was tightly localized to the plasma membrane (Figure 1C). Similar results were obtained upon ectopic expression in larval salivary glands (Figure 1C), where Nwk-EGFP and Nwk^{Δ1-428}-EGFP exhibited a very similar localization to the cytoplasm and to occasional puncta, while Nwk1-428eGFP was strongly targeted to the plasma membrane. Taken together, these data indicate that selective membrane targeting of Nwk in vivo requires both the F-BAR domain and the C-terminal sequence including SH3 domains and is essential for proper NMJ development.

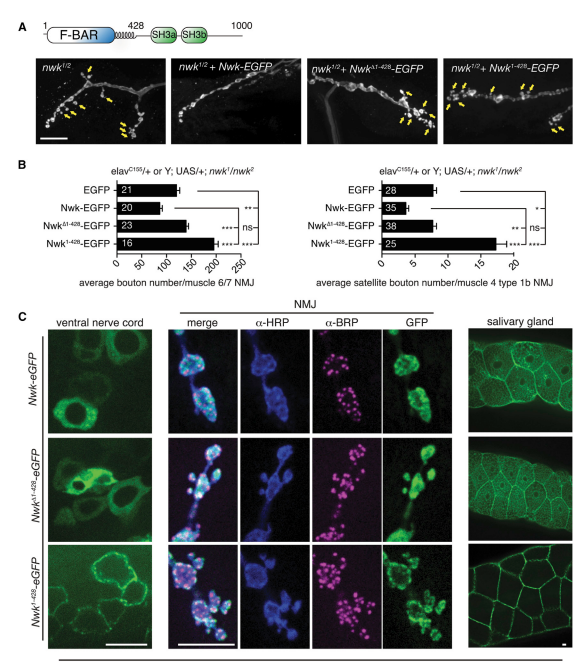
Nwk C-Terminal Sequences Regulate Its Membrane-Deforming Activity

Next, we investigated the regulation of the Nwk F-BAR domain by C-terminal sequences using a heterologous expression assay in Drosophila S2 cells, which do not express endogenous Nwk. We previously showed using this assay that the F-BAR domain of Nwk (Nwk1-428) localized to the S2 cell plasma membrane and generated membrane buds that were extended into protrusions by the actin cytoskeleton (Becalska et al., 2013; Kelley et al., 2015). We tested the activities of GFP-tagged full-length Nwk or FCHSD2, a murine Nwk ortholog, and found that unlike their respective isolated F-BAR domains, they did not generate protrusions and exhibited a predominantly cytoplasmic localization with occasional puncta (Figures 2A-2D), suggesting that the C terminus of Nwk contains a conserved inhibitory activity for its F-BAR domain. We generated a series of C-terminal truncations of Nwk and quantified protrusion formation in S2 cells, and we found that the inhibitory activity mapped to the second SH3 domain (SH3b; Figures 2A, 2B, and 2D). Given our previous finding that purified Nwk¹⁻⁷³¹ (which includes both SH3 domains) exhibits reduced membrane binding compared to the isolated F-BAR domain (Becalska et al., 2013), SH3b-domain-mediated inhibition is likely autoinhibitory, rather than due to a trans-acting factor. Further, inhibition did not require a short proline-rich stretch at the N terminus of Nwk (Nwk¹⁰⁻⁷³¹; Figures 2A, 2B, and 2D), suggesting that this sequence does not mediate autoinhibition via canonical SH3-domain/polyproline interactions.

Nwk SH3 Domains Bind to the F-BAR Domain Tips via Electrostatic Interactions

Next, we tested the prediction that autoinhibition is mediated by direct interactions between the Nwk F-BAR and SH3b domain using glutathione S-transferase (GST) pull-down assays with purified Nwk fragments, and we found that the F-BAR domain could co-precipitate with the SH3b domain (Figure 3A). We then tested the salt sensitivity of Nwk F-BAR/SH3 interactions to differentiate between hydrophobic interactions (required for canonical proline-rich ligands) or electrostatic interactions. Co-precipitation of the Nwk F-BAR domain with the GST-SH3b domain was sensitive to increasing ionic strength (Figure 3A). Further, interaction of the F-BAR domain with a tandem GST-SH3b domain was less sensitive than the GST-SH3b domain alone, suggesting that both SH3 domains contribute to the F-BAR C terminus interaction (Figure 3A).

We then used the protein homology/analogy recognition engine program Phyre (Kelley and Sternberg, 2009) to model the Nwk SH3b domain against a fold library of all existing PDB crystal structures. The highest-scoring Phyre model was based on the structure of the BIN1/AMPH2 SH3 domain (PDB structure c1mv3A). Using this model (Figures 3B and S2A), we mutated conserved charged residues in the RT-loop of the Nwk SH3b domain (E652 and E654), which are predicted to be in a similar orientation to the residues mediating electrostatic interactions of the Syndapin 1 SH3 domain with its F-BAR domain (Rao et al., 2010). In addition, we mutated conserved charged residues in the N-src loop of the Nwk SH3b domain (D673 and D674) (Figures 3C and S2A). Nwk-SH3b^{E654R} showed reduced F-BAR binding, while E652, D673, and D674 did not appear to play critical roles



elav^{C155}/+ or Y; UAS/+; nwk¹/nwk²

Figure 1. The Nwk F-BAR Domain Is Required for Its In Vivo Function

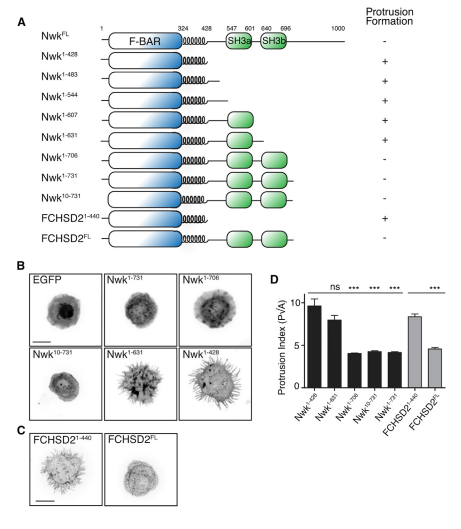
(A) Schematic of Nwk domain structure and representative confocal images of α-Cpx-stained third-instar larval NMJ morphology on muscle 4 for the indicated genotypes. Arrows indicate satellite boutons.

(B) Quantification of synaptic growth on muscle 6/7 and satellite bouton number on muscle 4 by Nwk variants in a nwk^1/nwk^2 null background. Graphs show mean \pm SEM. Numbers in bar graphs represent the number of NMJs.

(C) Localization of Nwk variants. GFP-tagged Nwk variants were expressed in the nwk^1/nwk^2 null background, under control of the GAL4 driver elav^{C155} (panneuronal and salivary glands). Third-instar larvae were fixed and stained with α -BRP and anti-horseradish peroxidase (α -HRP) antibodies. Scale bars, 10 μ m. See also Figure S1.

(Figure 3C). Interestingly, Nwk-SH3b^{E654R} did not disrupt binding to a previously identified ligand, Dap160 (Rodal et al., 2008), suggesting that the function of E654 in binding to the Nwk

F-BAR is a specific rather than general feature of the SH3 domain (Figure S2B). Finally, to test whether disrupting the interaction between the SH3b and F-BAR domains activates Nwk



membrane-remodeling activity in cells, we transfected S2 cells with Nwk^{E654R} and examined protrusion formation. Cells expressing Nwk^{E654R} exhibited membrane-remodeling activity similar to Nwk¹⁻⁴²⁸ (Figure 3D). Thus, SH3-domain-dependent inhibition of Nwk F-BAR activity is likely mediated primarily by electrostatic interactions and not by canonical SH3-proline motif interactions.

Next, we examined which site on the F-BAR domain is required for electrostatic interactions with the SH3 domain module. The tips of the Nwk F-BAR domain feature strong positive charge, which is important for high-affinity membrane association in vitro and membrane-deforming activity in S2 cells (Becalska et al., 2013). Nwk^{1-428Δtips} lacks the positively charged tips but exhibits a similar urea denaturation profile to Nwk¹⁻⁴²⁸ and is likely to be well-folded in solution (Becalska et al., 2013). Nwk SH3a and SH3b exhibited significantly reduced binding to Nwk^{1-428Δtips} compared to wild-type Nwk¹⁻⁴²⁸ (Figure 3E). These results indicate that the F-BAR tips are specifically required for Nwk SH3-F-BAR interactions in solution.

Nwk SH3 Domains Decrease F-BAR Membrane Binding

To test the role of the Nwk SH3 domains in regulation of F-BAR domain activity, we evaluated how the interaction between

Figure 2. The Activity of the Nwk F-BAR Domain in S2 Cells Is Negatively Regulated by Its SH3b Domain

(A) Schematic of Nwk constructs and summary of protrusion formation in cells.

(B and C) Activity of GFP-tagged Nwk truncations in S2 cells. Images show GFP fluorescence (in inverted contrast) of a maximum-intensity Z-projection. Scale bars, 10 μ m.

(D) Quantification of cellular morphology (perimeter [P]/square root of area [A]). Data are represented as mean \pm SEM from at least nine cells per condition; ***p < 0.001.

these domains affects membrane binding by purified Nwk. We previously showed that when compared to Nwk¹⁻⁴²⁸ in bulk liposome co-sedimentation assays, Nwk¹⁻⁷³¹ requires more phosphatidylinositol(4.5)bisphosphate

 $(Pl(4,5)P_2)$ to bind to membranes (Becalska et al., 2013). To determine if Nwk SH3 domains were required and sufficient for inhibition of membrane binding, we compared liposome co-sedimentation of Nwk¹⁻⁴²⁸, Nwk¹⁻⁷³¹, and Nwk¹⁻⁶³³ (SH3b domain truncated; Figure 4A). Compared to Nwk¹⁻⁴²⁸ and Nwk¹⁻⁷³¹, Nwk¹⁻⁶³³ required an intermediate Pl(4,5)P₂ concentration for binding (Figure 4B), suggesting that the SH3b domain is involved in, but not sufficient for, inhibiting membrane binding by purified Nwk¹⁻⁷³¹ and that intervening sequences (including the SH3a domain)

contribute to the remaining autoinhibitory effect. In contrast, in S2 cells, relief of autoinhibition depends more strictly on the electrostatic surface of the SH3b domain (Figures 2A, 2B, and 2D), suggesting that in a cellular context, a binding partner may modulate the contribution of SH3a to autoinhibition.

We then tested the effects of a purified SH3ab fragment in *trans* on Nwk¹⁻⁴²⁸ membrane binding. We found that pre-incubation of the F-BAR domain with SH3ab dramatically shifted the PI(4,5)P₂-dependent binding profile of Nwk¹⁻⁴²⁸ to more closely resemble Nwk¹⁻⁷³¹ (Figure 4C). Furthermore, the SH3ab fragment did not co-sediment with liposomes, either on its own or with low or high concentrations of Nwk¹⁻⁴²⁸ (Figures 4C and S3A). These results suggest that the interaction with the Nwk F-BAR domain directly inhibits membrane binding in solution and that the SH3 domains do not bind to the F-BAR domain when it is membrane associated.

To test whether autoregulation alters the properties of Nwk once bound to the membrane, we examined liposomes incubated with Nwk¹⁻⁴²⁸ and Nwk¹⁻⁷³¹ by cryo-electron microscopy (cryo-EM). Liposomes incubated with Nwk¹⁻⁷³¹ exhibited the same types of protein-dependent deformations

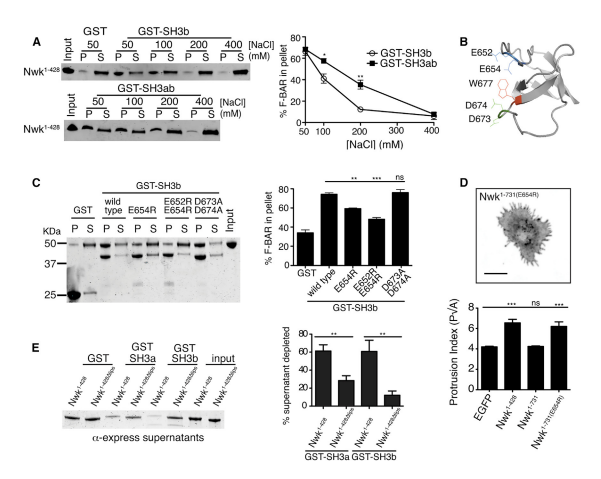


Figure 3. Direct Interactions between the Nwk SH3 and F-BAR Domains

GST fusion proteins were immobilized on glutathione agarose and incubated with the indicated purified proteins. Pellets and supernatants were fractionated by SDS-PAGE, immunoblotted or Coomassie stained, and quantified by densitometry. Graph shows the average \pm SEM of three independent reactions. *p < 0.05, **p < 0.01, and ***p < 0.001.

(A) Salt sensitivity of 6xHis-Xpress-Nwk¹⁻⁴²⁸ (1.5 μM) co-sedimentation with GST-Nwk-SH3 domains (2–3 μM). Image shows representative anti-Xpress tag immunoblot of co-sedimentation assay.

(B) Phyre model of Nwk SH3 domain (based on BIN1/AMPH2 SH3 [PDB: C1mv3A]).

(C) Co-sedimentation of Nwk¹⁻⁴²⁸ (1.5 μM) with GST or GST-Nwk-SH3b variants (3 μM). Graph shows the average ± SEM of three independent reactions. **p < 0.01, ***p < 0.001. Image shows representative Coomassie-stained gel.

(D) Activity of Nwk^{1-731(E654R)} in S2 cells. Image shows GFP fluorescence (in inverted contrast) for a maximum intensity Z-projection. Scale bar, 10 µm. Quantification of cellular morphology (perimeter [P]/square root of area [A]) is shown below. Data are presented as mean ± SEM from at least 19 cells per condition. ***p < 0.001.

(E) Co-sedimentation of Nwk¹⁻⁴²⁸ (1.5 μ M) with GST-Nwk-SH3b and GST-Nwk-SH3a (3–3.5 μ M) depends on the charged dimer tips. Image shows a representative immunoblot. Graph shows the average \pm SEM of three independent reactions. ** p < 0.01. See also Figure S2.

as we previously observed for Nwk^{1–428} (pointing, pinching, and scalloping; Figure 4D). These types of deformations did not change with longer protein-liposome incubation times (Figure S3B) and were observed more frequently with increasing protein concentration, suggesting that they are actively generated by Nwk (Figures 4D, S3E, and S3F). Finally, we did not observe any new properties such as vesiculation or tubulation for Nwk^{1–731} compared to Nwk^{1–428} under these conditions (Figures 4D and S3C–S3E). Thus, SH3 domain-mediated autoregulation via F-BAR domain tips inhibits membrane binding in solution, without changing the membrane-deforming properties of Nwk.

Nwk-Mediated Membrane Deformation Is Associated with Assembly of F-BAR Dimers into Stable Scaffolds

Next, we considered two possible explanations for reduced membrane binding for Nwk¹⁻⁷³¹ compared to Nwk¹⁻⁴²⁸ in bulk assays: either Nwk¹⁻⁷³¹ could be uniformly reduced compared to Nwk¹⁻⁴²⁸ on all liposomes (suggesting a general reduction in binding), or alternatively, the fraction of liposomes bound by Nwk¹⁻⁷³¹ could be decreased (suggesting a more complex or cooperative model for binding). To distinguish between these possibilities, we imaged Alexa Fluor 549 SNAP-tagged Nwk¹⁻⁴²⁸ or Nwk¹⁻⁷³¹ on nitro-2-1,3-benzoxadiazol-4-yl phosphatidylethanolamine (NBD-PE)-labeled giant unilamellar vesicles (GUVs) by confocal

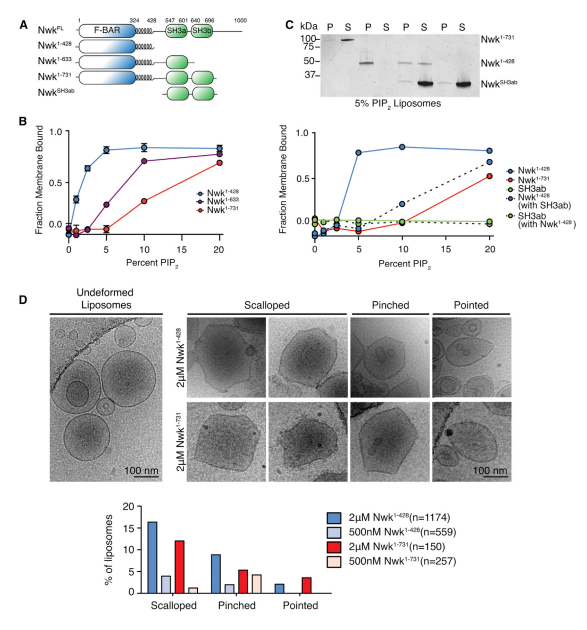


Figure 4. Nwk SH3b Inhibits Membrane Deformation by the Nwk F-BAR Domain

(A) Nwk constructs used for in vitro assays.

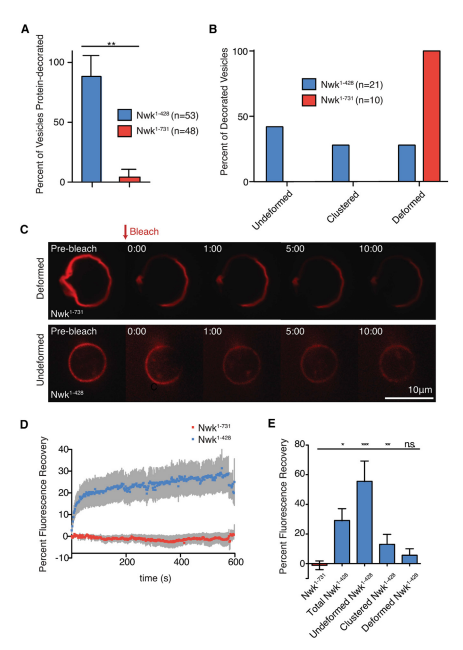
(B and C) Purified proteins (10 μM Nwk^{SH3ab} [Nwk residues 536–731]; 0.5 μM all other proteins) were incubated with liposomes of the following composition: 80-X% PC, 15% PE, 5% PS and X% PI(4,5)P₂ (where X is the concentration indicated in the graph) and subjected to liposome cosedimentation assays. Graphs show mean densitometry from one (B) or three (mean ± SEM) (C) independent experiments. (C) Image shows representative Coomassie staining of supernatant (S) and pellet (P) fractions at 10% PI(4,5)P₂.

(D) Cryo-EM of control and Nwk deformed liposomes. Both purified Nwk¹⁻⁴²⁸ and Nwk¹⁻⁷³¹ induce membrane scalloping, pointing, and pinching of 10% PI(4,5)P₂ liposomes. Scale bar, 100 nm. Bar graph summarizes vesicle morphology after 30-min incubation of Nwk¹⁻⁴²⁸ or Nwk¹⁻⁷³¹ (2 μ M and 500 nM) with 0.3 mM [DOPC:DOPE:DOPS:PI(4,5)P₂] = 70:15:5:10 liposomes. n represents the number of liposomes examined. See also Figure S3.

microscopy. Nwk^{1–731} strongly decorated a dramatically smaller fraction of GUVs compared to Nwk^{1–428} (Figure 5A), supporting the second model. We previously showed that Nwk induces deformations including membrane flattening, pinching, and clustering on GUVs (Becalska et al., 2013). Strikingly, all Nwk^{1–731}-bound GUVs were highly deformed, compared to a much smaller fraction

of Nwk^{1–428}-bound GUVs (Figure 5B). Thus, though SH3-domainmediated inhibition limits F-BAR interaction with the membrane, it unexpectedly enhances F-BAR membrane remodeling activity on the few vesicles that are decorated.

Higher-order assemblies are a conserved feature of BAR domains and likely central to their membrane-remodeling



properties. To determine if Nwk autoregulation affects its propensity to assemble on (and, by extension, deform) membranes, we next examined higher-order assembly of SNAPtagged Nwk using fluorescence recovery after photobleaching (FRAP) experiments. BAR domains can form stable scaffolds on membrane, limiting lipid fluidity and lateral diffusion of proteins (Zhao et al., 2013). Conversely, individually bound or incompletely oligomerized BAR domain dimers are expected to be mobile, either through exchange between the membrane and solution or rapid lipid diffusion. Therefore, assembled BAR domains should recover very slowly after photobleaching, while individually bound or disordered BAR domains should recover rapidly. Figure 5. Nwk SH3 Domains Limit Promiscuous Membrane Binding and Promote Stable F-BAR Protein Scaffolds Associated with Membrane Remodeling

NBD-PE-labeled GUVs incubated with SNAP-549tagged Nwk variants (500 nM) were imaged by spinning disk confocal microscopy. Lipid composition was [DOPC:POPE:DOPS:PI(4,5)P₂:NBD-PE] = 75:14.5:5:5:5. n represents the number of vesicles examined.

(A) Percent of protein-decorated GUVs after 30 min incubation with Nwk^{1–428} (blue) or Nwk^{1–731} (red). Graph represents mean \pm SEM from three independent experiments. Data are identical to 5% Pl(4,5)P₂ data point in Figures 6B and 6C.

(B) Quantification of the morphology of proteindecorated GUVs after 30-min incubation with purified Nwk¹⁻⁴²⁸ or Nwk¹⁻⁷³¹. GUVs were imaged from n independent reactions.

(C) Single spinning disc confocal slices of GUVs showing Nwk¹⁻⁷³¹ displaying limited recovery on deformed membranes, while partial recovery is observed on spherical, undeformed Nwk¹⁻⁴²⁸-coated GUVs. Scale bar, 10 μ m. See also Movies S1, S2, and S3.

(D) Quantification of recovery of protein fluorescence for Nwk¹⁻⁴²⁸ and Nwk¹⁻⁷³¹. The data are a mean of at least ten independent experiments and the error bars indicate \pm SEM.

(E) Quantification of protein fluorescence recovery according to vesicle morphology. One-way ANOVA (*p < 0.05, **p < 0.005, and ***p < 0.001).

We photobleached a region corresponding to less than ~10% of the GUV surface and examined the recovery of fluorescent Nwk into this region (Figure 5C; Movies S1, S2, and S3). For the entire population of GUVs, recovery of Nwk¹⁻⁷³¹ was significantly lower than Nwk¹⁻⁴²⁸ 10 min post-bleach (Figures 5D and 5E), and the distribution of total recovery for Nwk¹⁻⁴²⁸ was much wider, whereas Nwk¹⁻⁷³¹ recovery never exceeded 11.5%. We assessed recovery based on GUV morphology and found that spherical vesicles had significantly

higher total recovery (up to 75%) compared to deformed vesicles (less than 10%; Figure 5E). Notably, protein mobility on Nwk^{1–428} and Nwk^{1–731} GUVs that were deformed was not significantly different, indicating similar behavior of assembled F-BAR proteins at sites of deformation for both Nwk^{1–428} and Nwk^{1–731}. These results suggest that under these conditions, the isolated F-BAR domain is capable of interacting with the membrane in a range of different assembly states with distinct FRAP profiles, from individually bound and disordered with rapid turnover and protein fluorescence recovery to assembled stable protein scaffolds with slower turnover and protein fluorescence recovery. In contrast, Nwk^{1–731} protein coats do not recover after photobleaching, indicating that under these conditions all

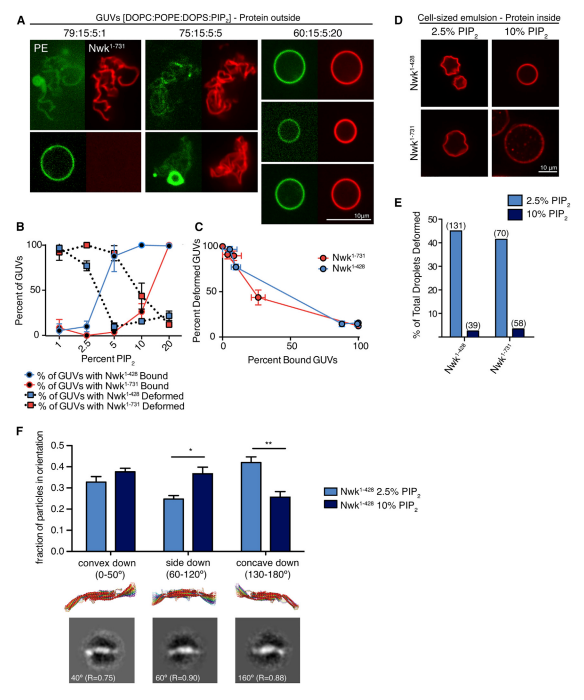


Figure 6. PI(4,5)P₂ Levels Regulate F-BAR Membrane Binding and Deformation

(A) Representative images of NBD-PE-labeled GUVs of varying PI(4,5)P₂ concentrations incubated with 500nM SNAP-549-Nwk¹⁻⁷³¹ for 30 min and imaged by spinning disk confocal microscopy.

(B) Percent of vesicles bound by Nwk¹⁻⁴²⁸ and Nwk¹⁻⁷³¹ at various Pl(4,5)P₂ concentrations (solid line) and percent of bound vesicles that are deformed (dotted line). Graph represents mean ± SEM from two to three independent experiments. 5% Pl(4,5)P₂ data are identical to Figure 5A.

(C) A plot of percent bound vesicles versus percent-deformed (from data in B) shows an inverse relationship between the number bound and frequency of deformation.

(D) Representative images of cell-sized water droplets with 1 μ M SNAP549-labeled BAR proteins. Emulsions were made with 23 mM lipid mixes of DPHPC \pm PI(4,5)P₂ in decane and incubated for 1 hr before imaging. See also Movies S4 and S5.

(E) Quantification of the percentage of total vesicles with protein-induced deformation at 2.5% and 10% Pl(4,5)P₂ for Nwk¹⁻⁴²⁸ and Nwk¹⁻⁷³¹. Number above bars represents the number of droplets examined. Associated with Figure S4.

membrane-bound Nwk¹⁻⁷³¹ is assembled into stable higher-order scaffolds.

Nwk Autoregulation Increases the Phosphoinositide Requirement for Membrane Deformation without Altering the Mechanism of Membrane Binding

Next, we tested whether SH3 domains specifically enhance assembly of Nwk F-BAR domains or if the differences in the assembly and deformation properties of Nwk¹⁻⁴²⁸ and Nwk¹⁻⁷³¹ on GUVs of a fixed lipid composition (Figure 5) reflect inherent differences in Nwk F-BAR behavior at different points on the PI(4,5)P2 binding curve (Figure 4B). We first tested the ability of Nwk¹⁻⁷³¹ and Nwk¹⁻⁴²⁸ to bind and deform GUVs with different PI(4,5)P₂ compositions. Remarkably, at 1% PI(4,5)P₂, a concentration at which Nwk¹⁻⁴²⁸ is only partially membrane associated in co-sedimentation assays (Figure 4B), we found that it decorated only a small fraction of highly deformed GUVs, much like Nwk¹⁻⁷³¹ at 5% PI(4,5)P₂ (Figures 6A and 6B). Further, at 20% PI(4,5)P₂, a concentration at which all of Nwk¹⁻⁷³¹ is bound to liposomes in co-sedimentation assays (Figure 4B), all GUVs were bound but poorly deformed, much like Nwk¹⁻⁴²⁸ at 5% PI(4,5)P₂ (Figures 6A and 6B). To rule out any quantitative differences in the behavior of Nwk¹⁻⁴²⁸ and Nwk¹⁻⁷³¹ reflecting different mechanisms of assembly or deformation, we normalized the fraction of membrane-bound Nwk to the extent of membrane deformation at the full range of $PI(4,5)P_2$ concentrations. Nwk¹⁻⁴²⁸ and Nwk1-731 exhibited quantitatively indistinguishable behavior in this analysis (Figure 6C). These results suggest an inherent bias of the F-BAR domain toward high-order scaffolds and membrane deformation when binding conditions are more stringent.

Several different parameters could together account for the observation that a small number of GUVs are highly decorated and deformed under stringent binding conditions. F-BAR proteins may exhibit nucleation behavior, resulting in recruitment of protein molecules to a small fraction of vesicles, until the concentration of protein in solution is depleted below a critical assembly threshold (nucleation mechanism). However, our observations also raise the possibility that F-BAR proteins have inherently different activities on membranes of low versus high PI(4,5)P₂ content (lipid-directed mechanism). To examine this possibility, we tested F-BAR-domain-mediated membrane deformation in cell-sized water droplets encapsulated within a phospholipid membrane. In this system, protein is emulsified in a mix of lipid and oil, resulting in the formation of droplets of protein surrounded by an interface of lipid, with head groups facing the protein in the aqueous phase (Figure S4A; Hase and Yoshikawa, 2006; Miyazaki et al., 2015). Compared to GUVs, droplets are isolated from each other and therefore not subject to nucleation effects. Both SNAP-Nwk¹⁻⁴²⁸ and SNAP-Nwk¹⁻⁷³¹ induced deformation and crumpling of PI(4,5)P2-encapsulated droplets, but not droplets without PI(4,5)P2 (Figures 6D, 6E, S4B, and S4C; Movies S4 and S5). Further, the SNAP tag alone did not deform PI(4,5)P₂-encapsulated droplets, indicating that crumpling is a specific activity of Nwk (Figure S4B). Remarkably, deformation was prevalent at 2.5% PI(4,5)P₂ for both Nwk¹⁻⁴²⁸ and Nwk¹⁻⁷³¹ but rare at 10% PI(4,5)P₂, supporting a lipid-directed mechanism that favors deformation at lower negative membrane charge (Figures 6D and 6E).

One possible mechanism to account for differential activities of the Nwk F-BAR domains with changing negative charge could be the orientation of the F-BAR on the membrane. F-BAR domains interact with membranes through positively charged concave surfaces when assembled on highly curved membranes but can also interact with membranes in a number of other orientations, including a deformation-inactive side-lying state (Frost et al., 2008; Yu and Schulten, 2013; Becalska et al., 2013). To test whether different modes of protein-membrane interaction may be favored depending on membrane charge, we used single-particle electron microscopy (EM) on lipid monolayers to compare the orientations of the Nwk F-BAR domain at 2.5% and 10% PI(4,5)P₂ (Figure 6F). The Nwk F-BAR forms an S-shaped dimer whose chirality allows single-particle averages to be assigned to concave-surface-down, side-lying, or convex-surface-down orientations (Figure 6F; Becalska et al., 2013). We found that at 2.5% PI(4,5)P₂, the concave-surface-down orientation predominates, whereas at 10% PI(4,5)P₂ Nwk does not show a strong proclivity for any one orientation. These data support a model by which limited membrane charge results in an orientation-selective binding mode, favoring assembly and membrane deformation, while high membrane charge favors promiscuous binding and diminishes the likelihood of assembly.

Nwk Deforms Cellular Membranes within a Sweet Spot of PI(4,5)P₂ Concentration

In order to test the role of PI(4,5)P2 levels in regulating Nwk F-BAR activity in vivo, we used our S2 cell assay. We acutely increased PI(4,5)P₂ using YU1422670, a small-molecule inhibitor of OCRL (Pirruccello et al., 2014), which is an important phosphoinositide-5-phosphatase in S2 cells (Ben El Kadhi et al., 2011). This compound is active in Drosophila, as it recapitulates the reported OCRL mutant phenotype of increased F-actin levels in S2 cells (Ben El Kadhi et al., 2011; Figure S5). Treatment with low concentrations of the drug (10 µM) for 15 min increased cellular deformation in Nwk1-731-EGFP-expressing S2 cells, supporting our in vitro finding that PI(4,5)P2 promotes membrane binding by Nwk¹⁻⁷³¹ (Figures 7A-7C). In contrast, high levels of drug (30 µM) decreased deformation of Nwk¹⁻⁴²⁸-EGFP-expressing cells, though Nwk was still localized to the membrane (Figures 7D–7F), supporting our finding that high levels of $PI(4,5)P_2$ decouple membrane binding and deformation. YU1422670treated cells that were not expressing Nwk had normal cellular morphology and actin localization (despite modestly increased levels of F-actin) (Figure S5), indicating that the differences

⁽F) Comparison of the orientation of Nwk¹⁻⁴²⁸ dimers on lipid monolayers with 2.5% or 10% Pl(4,5)P₂. Error bars represent fraction of particles \pm SEM from three or four independent EM grids. Shown below are superimposed 10° rotations of the predicted Nwk¹⁻³¹³ structure (Becalska et al., 2013), and representative class averages from 10% Pl(4,5)P₂, grid A, Table S1. R represents correlation coefficients to the 20-Å-filtered predicted structure. See also Table S1. Scale bars, 10 μ M.

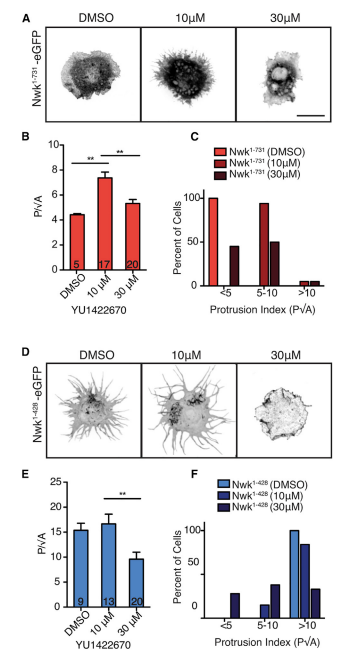


Figure 7. Cellular PI(4,5)P₂ Levels Regulate Nwk-Induced Cellular Deformation

(A and D) Increasing cellular PI(4,5)P₂ levels with the OCRL inhibitor YU1422670 alters Nwk-induced deformation. Images show GFP fluorescence (in inverted contrast) of maximum intensity Z-projections. Scale bar, 10 μ m. (B and C) Nwk¹⁻⁷³¹-EGFP expressing cells exhibit a significantly higher protrusion index (P/ \sqrt{A} ; mean \pm SEM; number in bar graph represents n cells) at low concentrations of YU1422670 compared to DMSO control-treated cells (B), with a higher percentage of cells with a protrusion index of >5 (C).

(E and F) Nwk¹⁻⁴²⁸-EGFP expressing cells have a significantly lower protrusion index (P/ \sqrt{A} ; mean \pm SEM; number in bar graph represents n cells) at high concentration (30 μ M) of YU1422670 compared to DMSO and 10 μ M YU1422670 (E) and a lower percentage of cells with a protrusion index of 10 or higher after drug treatment (F). See also Figure S5.

observed in Nwk-expressing cells after drug treatment are due to differential F-BAR activity. These results provide strong in vivo support for the model that $PI(4,5)P_2$ levels can shift the F-BAR domain between unbound, membrane-deforming, and promiscuous binding states and that autoregulation increases the $PI(4,5)P_2$ concentration at which these states occur.

DISCUSSION

Mechanisms of Nwk Regulation by Intramolecular Interactions and Membrane Composition

Here, we report that the F-BAR domain of Nwk is autoregulated by its C-terminal SH3 domains and that the Nwk C terminus is critical for regulation of its F-BAR localization and membrane remodeling function in vivo. Surprisingly, membrane composition is a key factor in the efficiency of remodeling, and autoregulation alters the optimal membrane charge requirement for maximum F-BAR activity.

Nwk is one of several F-BAR proteins with two SH3 domains (Roberts-Galbraith and Gould, 2010), and we found that the SH3b domain mediates the majority of the inhibitory effect in a cellular context. This inhibition occurs via non-canonical electrostatic interactions between the SH3b and F-BAR domains, similar to Syndapin (Rao et al., 2010). Further, we show that the autoregulatory SH3 domains are displaced from the F-BAR domain upon lipid binding. Our results indicate that regulation may directly reduce the probability of membrane binding by occluding residues important for interactions with the membrane. Our finding that binding and deformation scale similarly for Nwk¹⁻⁴²⁸ and Nwk¹⁻⁷³¹ (Figure 6C) argues that they do not bind, oligomerize, or deform membranes by distinct mechanisms and that SH3 domains do not actively promote higherorder assembly via inter-dimer interactions. Instead, the SH3 domains shift the threshold for binding so that binding and deformation occur at higher negative charge. Importantly, we show that both the isolated F-BAR domain and the full-length protein actively generate similar types of membrane deformations (Figures 4D and 6D; Movies S4 and S5), arguing against the model that these proteins merely sense curvature, even under stringent binding conditions.

To date, autoregulation of BAR domain proteins has been thought of as an "on-off switch" for membrane binding, released by binding partners (Roberts-Galbraith and Gould, 2010; Meinecke et al., 2013) or by increased membrane charge (Wu and Baumgart, 2014). However, our data support a more nuanced explanation of these inhibitory interactions that depends critically on local membrane composition. While the net effect of autoinhibition is to shift the membrane-binding and membranedeformation curves to higher negative charge, we found that the membrane-binding and membrane-deforming curves do not overlap, as would be predicted by the simple "on-off switch" model. Rather, our FRAP, GUV, and droplet deformation data indicate that higher-order assembly and deformation occur efficiently only within a range or "sweet spot" of negative charge and that high levels of PI(4,5)P2 promote promiscuous membrane binding of the F-BAR domain.

A combination of protein-directed and lipid-directed mechanisms may account for this unexpected property of the Nwk F-BAR domain. Under stringent binding conditions and when the protein to lipid ratio is limiting (e.g., on GUVs), F-BAR domains exhibit nucleation behavior, resulting in favored binding and assembly on a small population of vesicles. Importantly, this is a physiologically relevant regime, since cellular membranes and proteins are present in limiting amounts, lipid composition is constantly changing, and F-BARs are acutely released from autoinhibition by regulatory binding partners. Nucleation of F-BAR domains may occur by several potential, nonexclusive mechanisms. F-BAR domains can induce clustering of PI(4,5) P₂ (Zhao et al., 2013), which could drive further protein binding and deformation in a feed-forward mechanism. Conversely, PI(4,5)P2 clusters may form in areas of spontaneous local membrane curvature (Koldsø et al., 2014), promoting curvature-sensitive or properly oriented binding of F-BAR domains. Finally, membrane-bound F-BAR/F-BAR oligomers may increase the propensity for further protein assembly through the effects of avidity. In addition to nucleation effects, our cell-sized water droplet experiments (which eliminate protein sequestration due to nucleation) suggest that the Nwk F-BAR domain preferentially deforms membranes with lower negative charge. One mechanism for this behavior, supported by our single-particle EM data, is that the F-BAR domain is biased toward a deformation-promoting (concave-surface-down) conformation at low $PI(4,5)P_2$. Another nonexclusive possibility is that at high PI(4,5)P₂, the organization or rigidity of membrane itself may inhibit assembly, either by forming lipid clusters of a size that disfavor deformation or by restricting dynamic rearrangements of individual proteins into higher assemblies (Ruiz-Herrero and Hagan, 2015). Assessing the relative contributions of these forces will require developing a theoretical framework that describes the nucleation, oligomerization, and lipid-binding behaviors of Nwk.

Other proteins in the BAR domain family are sensitive to negative membrane charge and may be similarly regulated by nucleation behavior and membrane composition. Indeed, elevated $PI(4,5)P_2$ levels suppress the membrane deforming activity of the F-BAR protein FBP17 in vivo (Tsujita et al., 2015), suggesting that the sweet spot may be conserved feature of BAR domain activity. Further investigation will be required to determine whether BAR domains have different ranges of optimal membrane charge, which could contribute to the specificity of these proteins in the context of their cellular roles.

Contributions of Membrane Composition and Autoregulation to the Role of Nwk In Vivo

Despite advances in our knowledge of the molecular mechanisms regulating BAR domain membrane curvature-sensing and sculpting properties, as well as the effects of their binding partners, we still lack a thorough understanding of how these activities are regulated and targeted in their cellular contexts. Our data show that Nwk requires both F-BAR activity and the C-terminal regulatory SH3 domains for its in vivo role in growth signal regulation at the *Drosophila* NMJ. Removing the Nwk C terminus produces a more severe phenotype than the null mutant, suggesting that constraining and regulating Nwk F-BAR activity and targeting to the membrane via autoregulation is required for proper synapse formation. A critical next step is to consider the role of binding partners in modulating the state of the Nwk F-BAR on the membrane. Nwk interacts with dynamin and the endocytic scaffold Dap160/intersectin via the SH3a and SH3b domains, respectively (Rodal et al., 2008; O'Connor-Giles et al., 2008). These interactions, in addition to localizing BAR domains, may modulate Nwk autoinhibition to actively fine-tune membrane binding and deformation at specific levels of negative membrane charge.

Our data also raise an important issue: the physiological significance of autoregulatory mechanisms depends not only on the effects of Nwk-binding partners on its activity but also critically on the local PI(4,5)P₂ concentration at which Nwk operates in vivo. A primary defining characteristic of biological membrane compartments is their lipid composition. BAR domains selectively associate with intracellular membranes rich in charged phosphoinositides (Becalska et al., 2013; Di Paolo and De Camilli, 2006; Saarikangas et al., 2010). PI(4,5)P2 concentrations have been estimated to be \sim 0.5%–1% of total cellular membranes, but they are tightly regulated and may transiently accumulate to much higher concentrations (Di Paolo and De Camilli, 2006). Submicrometer clusters of PI(4.5)P₂ have been detected composed of as much as 80% PI(4,5)P₂ (Milosevic et al., 2005; James et al., 2008; van den Bogaart et al., 2011). This suggests that PI(4,5)P2 exists in cells at a diverse range of concentrations that are dynamically altered by phosphoinositide kinases and phosphatases. Local, regulated changes in lipid composition could shift membranebound Nwk between a disassembled state and an assembled state that is capable of deforming membranes. Using a pharmacological approach to acutely increase PI(4,5)P₂ levels in S2 cells, we found that, similar to our in vitro results, membrane association and deformation by Nwk occur at a sensitive range of cellular PI(4,5)P₂ that is elevated by autoregulation. Further tests of this lipid-directed mechanism of membrane remodeling in the nervous system will require the development of new methods to visualize endogenous Nwk-mediated membrane deformation and to acutely manipulate PI(4,5)P2 levels at the NMJ.

In summary, SH3-domain mediated autoinhibition of the Nwk F-BAR domain shifts the $PI(4,5)P_2$ dependence of higher-order assembly and promiscuous binding of individual dimers to the membrane. We predict that through this mechanism, activating or inhibitory binding partners work in concert with temporal and spatial regulation of negative membrane charge to control membrane remodeling by BAR domains.

EXPERIMENTAL PROCEDURES

Fly Stocks and NMJ Morphology Analysis

UAS-Nwk constructs were generated as described previously (Becalska et al., 2013) and injected into flies at Genetic Services using Φ c381 recombinase at the Attp40 locus. NMJs on muscle 6/7, segment A3, and muscle 4, segments A2–A3 were selected for analysis of morphology, in fixed third-instar larval fillets immunostained with α -Cpx (Huntwork and Littleton, 2007) and α -Dlg (Developmental Studies Hybridoma Bank) antibodies. Both type 1b and type 1s boutons were quantified on muscle 6/7. Only type Ib innervation, delineated by extensive postsynaptic α -Dlg staining, was quantified on muscle 4. Satellite boutons were defined as strings of five or fewer boutons extending from the main axis of the NMJ.

Cell Culture

S2 cells were cultured, transfected with Effectene, spread for 1 hr on Concanavalin-A-coated coverslips, and imaged on a spinning disk confocal microscope (see Supplemental Experimental Procedures) as described previously (Becalska et al., 2013). For S2 cell protrusion quantification, perimeter and area were calculated from maximum intensity projection images of confocal stacks of at least nine cells per sample. For the OCRL inhibitor experiment, cells were incubated with YU1422670 or 1% DMSO control for 15 min before spreading for 1 hr on Concanavalin-A-coated coverslips and fixing.

Cell-Sized Water Droplet Assay

Lipids (DPHPC [1,2-diphytanoyl-*sn*-glycero-3-phosphocholine], Avanti Polar Lipids) or DPHPC:Pl(4,5)P₂) were mixed in chloroform, dried down, and rehydrated to 23 mM (20 mg/ml) in decane. SNAP-tagged proteins were added to the lipid mix at a 1:50 volume ratio and pipetted vigorously until cloudy before imaging by spinning disk confocal microscopy.

GUV Assays

GUVs were generated by electroswelling on indium titanium oxide (ITO)coated slides. Approximately 450 μ M (350 μ g/ml) GUVs were mixed with 500 nM SNAP-tagged F-BAR proteins in 5 mM HEPES and 150 mM KCI (pH 7.5), incubated for 30 min, and imaged. For analysis of fraction of GUVs bound, fields of GUVs imaged were identified using only the NBD-PE signal. For analysis of GUV morphology, the SNAP-549 signal was used to identify bound vesicles. All non-spherical vesicles were classified as deformed. For FRAP experiments, GUVs were imaged for 10 min at 2-s intervals, with a pause for bleaching after time point 10 (20 s). Prebleach fluorescence was normalized to 1.0 in order to calculate the fraction of fluorescence recovery. Signal intensity over time from a non-bleached region of the GUV was used to correct for photobleaching.

Statistical Analyses

All error bars shown are mean \pm SEM. Statistical significance was calculated with GraphPad Prism 6 software using ANOVA followed by pairwise Tukey's tests or using Student's t tests where only two groups were compared (*p < 0.05, **p < 0.01, and ***p < 0.005). χ^2 tests for independence were used to evaluate the liposome morphology from EM images.

SUPPLEMENTAL INFORMATION

Supplemental Information includes Supplemental Experimental Procedures, five figures, one table, and five movies and can be found with this article online at http://dx.doi.org/10.1016/j.celrep.2015.11.044.

AUTHOR CONTRIBUTIONS

C.F.K., E.M.M., and A.A.R. conceived of the project and designed the experiments. C.F.K. and E.M.M performed and analyzed most of the experiments with the help of A.N.B., A.A.R., D.D., M.F.H., K.S., O.S.S., K.V., and S.W. C.F.K., E.M.M., O.S., and A.A.R. wrote the manuscript.

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REFERENCES

Becalska, A.N., Kelley, C.F., Berciu, C., Stanishneva-Konovalova, T.B., Fu, X., Wang, S., Sokolova, O.S., Nicastro, D., and Rodal, A.A. (2013). Formation of membrane ridges and scallops by the F-BAR protein Nervous Wreck. Mol. Biol. Cell *24*, 2406–2418.

Ben El Kadhi, K., Roubinet, C., Solinet, S., Emery, G., and Carréno, S. (2011). The inositol 5-phosphatase dOCRL controls PI(4,5)P2 homeostasis and is necessary for cytokinesis. Curr. Biol. *21*, 1074–1079.

Cao, H., Yin, X., Cao, Y., Jin, Y., Wang, S., Kong, Y., Chen, Y., Gao, J., Heller, S., and Xu, Z. (2013). FCHSD1 and FCHSD2 are expressed in hair cell stereocilia and cuticular plate and regulate actin polymerization in vitro. PLoS ONE *8*, e56516.

Chen, Z., Chang, K., Capraro, B.R., Zhu, C., Hsu, C.J., and Baumgart, T. (2014). Intradimer/Intermolecular interactions suggest autoinhibition mechanism in endophilin A1. J. Am. Chem. Soc. *136*, 4557–4564.

Coyle, I.P., Koh, Y.H., Lee, W.C., Slind, J., Fergestad, T., Littleton, J.T., and Ganetzky, B. (2004). Nervous wreck, an SH3 adaptor protein that interacts with Wsp, regulates synaptic growth in *Drosophila*. Neuron *41*, 521–534.

Di Paolo, G., and De Camilli, P. (2006). Phosphoinositides in cell regulation and membrane dynamics. Nature 443, 651–657.

Dickman, D.K., Lu, Z., Meinertzhagen, I.A., and Schwarz, T.L. (2006). Altered synaptic development and active zone spacing in endocytosis mutants. Curr. Biol. *16*, 591–598.

Frost, A., Perera, R., Roux, A., Spasov, K., Destaing, O., Egelman, E.H., De Camilli, P., and Unger, V.M. (2008). Structural basis of membrane invagination by F-BAR domains. Cell *132*, 807–817.

Guerrier, S., Coutinho-Budd, J., Sassa, T., Gresset, A., Jordan, N.V., Chen, K., Jin, W.L., Frost, A., and Polleux, F. (2009). The F-BAR domain of srGAP2 induces membrane protrusions required for neuronal migration and morphogenesis. Cell *138*, 990–1004.

Hase, M., and Yoshikawa, K. (2006). Structural transition of actin filament in a cell-sized water droplet with a phospholipid membrane. J. Chem. Phys. *124*, 104903.

Huntwork, S., and Littleton, J.T. (2007). A complexin fusion clamp regulates spontaneous neurotransmitter release and synaptic growth. Nat. Neurosci. *10*, 1235–1237.

Itoh, T., Erdmann, K.S., Roux, A., Habermann, B., Werner, H., and De Camilli, P. (2005). Dynamin and the actin cytoskeleton cooperatively regulate plasma membrane invagination by BAR and F-BAR proteins. Dev. Cell 9, 791–804.

James, D.J., Khodthong, C., Kowalchyk, J.A., and Martin, T.F. (2008). Phosphatidylinositol 4,5-bisphosphate regulates SNARE-dependent membrane fusion. J. Cell Biol. *182*, 355–366.

Kast, D.J., Yang, C., Disanza, A., Boczkowska, M., Madasu, Y., Scita, G., Svitkina, T., and Dominguez, R. (2014). Mechanism of IRSp53 inhibition and combinatorial activation by Cdc42 and downstream effectors. Nat. Struct. Mol. Biol. *21*, 413–422.

Kelley, L.A., and Sternberg, M.J. (2009). Protein structure prediction on the Web: a case study using the Phyre server. Nat. Protoc. *4*, 363–371.

Kelley, C.F., Becalska, A.N., Berciu, C., Nicastro, D., and Rodal, A.A. (2015). Assembly of actin filaments and microtubules in Nwk F-BAR-induced membrane deformations. Commun. Integr. Biol. 8, e1000703.

Koldsø, H., Shorthouse, D., Hélie, J., and Sansom, M.S. (2014). Lipid clustering correlates with membrane curvature as revealed by molecular simulations of complex lipid bilayers. PLoS Comput. Biol. *10*, e1003911.

Kumar, V., Fricke, R., Bhar, D., Reddy-Alla, S., Krishnan, K.S., Bogdan, S., and Ramaswami, M. (2009). Syndapin promotes formation of a postsynaptic membrane system in *Drosophila*. Mol. Biol. Cell *20*, 2254–2264.

McMahon, H.T., and Boucrot, E. (2015). Membrane curvature at a glance. J. Cell Sci. *128*, 1065–1070.

Meinecke, M., Boucrot, E., Camdere, G., Hon, W.C., Mittal, R., and McMahon, H.T. (2013). Cooperative recruitment of dynamin and BIN/amphiphysin/Rvs (BAR) domain-containing proteins leads to GTP-dependent membrane scission. J. Biol. Chem. *288*, 6651–6661.

Milosevic, I., Sørensen, J.B., Lang, T., Krauss, M., Nagy, G., Haucke, V., Jahn, R., and Neher, E. (2005). Plasmalemmal phosphatidylinositol-4,5-bisphosphate level regulates the releasable vesicle pool size in chromaffin cells. J. Neurosci. *25*, 2557–2565.

Mim, C., Cui, H., Gawronski-Salerno, J.A., Frost, A., Lyman, E., Voth, G.A., and Unger, V.M. (2012). Structural basis of membrane bending by the N-BAR protein endophilin. Cell *149*, 137–145.

Miyazaki, M., Chiba, M., Eguchi, H., Ohki, T., and Ishiwata, S. (2015). Cell-sized spherical confinement induces the spontaneous formation of contractile actomyosin rings in vitro. Nat. Cell Biol. *17*, 480–489.

Neumann, S., and Schmid, S.L. (2013). Dual role of BAR domain-containing proteins in regulating vesicle release catalyzed by the GTPase, dynamin-2. J. Biol. Chem. *288*, 25119–25128.

O'Connor-Giles, K.M., Ho, L.L., and Ganetzky, B. (2008). Nervous wreck interacts with thickveins and the endocytic machinery to attenuate retrograde BMP signaling during synaptic growth. Neuron 58, 507–518.

Owen, D.J., Wigge, P., Vallis, Y., Moore, J.D., Evans, P.R., and McMahon, H.T. (1998). Crystal structure of the amphiphysin-2 SH3 domain and its role in the prevention of dynamin ring formation. EMBO J. *17*, 5273–5285.

Pirruccello, M., Nandez, R., Idevall-Hagren, O., Alcazar-Roman, A., Abriola, L., Berwick, S.A., Lucast, L., Morel, D., and De Camilli, P. (2014). Identification of inhibitors of inositol 5-phosphatases through multiple screening strategies. ACS Chem. Biol. *9*, 1359–1368.

Rao, Y., Ma, Q., Vahedi-Faridi, A., Sundborger, A., Pechstein, A., Puchkov, D., Luo, L., Shupliakov, O., Saenger, W., and Haucke, V. (2010). Molecular basis for SH3 domain regulation of F-BAR-mediated membrane deformation. Proc. Natl. Acad. Sci. USA *107*, 8213–8218.

Roberts-Galbraith, R.H., and Gould, K.L. (2010). Setting the F-BAR: functions and regulation of the F-BAR protein family. Cell Cycle *9*, 4091–4097.

Rodal, A.A., Motola-Barnes, R.N., and Littleton, J.T. (2008). Nervous wreck and Cdc42 cooperate to regulate endocytic actin assembly during synaptic growth. J. Neurosci. 28, 8316–8325. Rodal, A.A., Blunk, A.D., Akbergenova, Y., Jorquera, R.A., Buhl, L.K., and Littleton, J.T. (2011). A presynaptic endosomal trafficking pathway controls synaptic growth signaling. J. Cell Biol. *193*, 201–217.

Ruiz-Herrero, T., and Hagan, M.F. (2015). Simulations show that virus assembly and budding are facilitated by membrane microdomains. Biophys. J. *108*, 585–595.

Saarikangas, J., Zhao, H., and Lappalainen, P. (2010). Regulation of the actin cytoskeleton-plasma membrane interplay by phosphoinositides. Physiol. Rev. *90*, 259–289.

Sun, X., Pinacho, R., Saia, G., Punko, D., Meana, J.J., Ramos, B., and Gill, G. (2015). Transcription factor Sp4 regulates expression of nervous wreck 2 to control NMDAR1 levels and dendrite patterning. Dev. Neurobiol. 75, 93–108.

Takei, K., Slepnev, V.I., Haucke, V., and De Camilli, P. (1999). Functional partnership between amphiphysin and dynamin in clathrin-mediated endocytosis. Nat. Cell Biol. *1*, 33–39.

Tsujita, K., Takenawa, T., and Itoh, T. (2015). Feedback regulation between plasma membrane tension and membrane-bending proteins organizes cell polarity during leading edge formation. Nat. Cell Biol. *17*, 749–758.

van den Bogaart, G., Meyenberg, K., Risselada, H.J., Amin, H., Willig, K.I., Hubrich, B.E., Dier, M., Hell, S.W., Grubmüller, H., Diederichsen, U., and Jahn, R. (2011). Membrane protein sequestering by ionic protein-lipid interactions. Nature 479, 552–555.

Vázquez, F.X., Unger, V.M., and Voth, G.A. (2013). Autoinhibition of endophilin in solution via interdomain interactions. Biophys. J. *104*, 396–403.

Wang, Q., Navarro, M.V., Peng, G., Molinelli, E., Goh, S.L., Judson, B.L., Rajashankar, K.R., and Sondermann, H. (2009). Molecular mechanism of membrane constriction and tubulation mediated by the F-BAR protein Pacsin/Syndapin. Proc. Natl. Acad. Sci. USA *106*, 12700–12705.

Wu, T., and Baumgart, T. (2014). BIN1 membrane curvature sensing and generation show autoinhibition regulated by downstream ligands and PI(4,5)P2. Biochemistry 53, 7297–7309.

Yu, H., and Schulten, K. (2013). Membrane sculpting by F-BAR domains studied by molecular dynamics simulations. PLoS Comput. Biol. 9, e1002892.

Zhao, H., Michelot, A., Koskela, E.V., Tkach, V., Stamou, D., Drubin, D.G., and Lappalainen, P. (2013). Membrane-sculpting BAR domains generate stable lipid microdomains. Cell Rep. *4*, 1213–1223.