Lipid binding promotes oligomerization and focal adhesion activity of vinculin

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A dherens junctions (AJs) and focal adhesion (FA) complexes are necessary for cell migration and morphogenesis, and for the development, growth, and survival of all metazoans. Vinculin is an essential regulator of both AJs and FAs, where it provides links to the actin cytoskeleton. Phosphatidylinositol 4,5-bisphosphate (PIP2) affects the functions of many targets, including vinculin. Here we report the crystal structure of vinculin in complex with PIP2, which revealed that PIP2 binding alters vinculin structure to direct higher-order oligomerization and suggests that PIP2 and F-actin binding to vinculin are mutually permissive. Forced expression of PIP2-binding-deficient mutants of vinculin in vinculin-null mouse embryonic fibroblasts revealed that PIP2 binding is necessary for maintaining optimal FAs, for organization of actin stress fibers, and for cell migration and spreading. Finally, photobleaching experiments indicated that PIP2 binding is required for the control of vinculin dynamics and turnover in FAs. Thus, through oligomerization, PIP2 directs a transient vinculin sequestration at FAs that is necessary for proper FA function.

Introduction

Inositol phospholipids are important regulators of cell physiology, and their head group interactions play fundamental roles in controlling membrane–cytosol interfaces. In addition to affecting signal transduction at the cell surface, these lipids regulate membrane traffic, nuclear events, the permeability and transport functions of membranes, and the cytoskeleton (Di Paolo and De Camilli, 2006). Phosphatidylinositol 4,5-bisphosphate (PIP2) plays roles in the recruitment and activation of a wide variety of actin regulatory proteins at the plasma membrane in order to regulate the release of cytokines, and to affect cell shape and motility, as well as several other processes (Ling et al., 2002, 2006; McLaughlin et al., 2002; Yin and Janmey, 2003). Binding of PIP2 is also critical for the function of actin-binding proteins that serve to link cell–cell or cell–matrix adhesion complexes at the plasma membrane with the actin cytoskeleton. There currently is a major conceptual gap in our understanding of how PIP2 binds to and alters the function of these adaptor/scaffold proteins, such as vinculin, at the molecular level.

Vinculin functions as a central regulator of adherens junctions (AJs) and focal adhesions (FAs) and is comprised of five loosely packed, antiparallel α-helix bundle domains. Four of these helix bundles are organized in a head (VH) domain that is connected to a five-helix bundle vinculin tail (Vt) domain by a disordered proline-rich region (Borgon et al., 2004), which functions as a hinge for the molecule. The inactive closed state of vinculin is characterized by extensive intramolecular hydrophobic head–tail interactions of its N-terminal helix bundle domain (Vh1) with Vt, and physiological triggers such as talin that activate vinculin disrupt these interactions, and allow the tail domain of the open conformer to bind to other ligands, including PIP2 (but not other lipids; Weekes et al., 1996; Palmer et al., 2009) and F-actin (Johnson and Craig, 1995; Peng et al., 2011). PIP2 levels are locally elevated at cell adhesion sites and PIP2 has been proposed to prevent F-actin binding (Steimle et al., 1999), to regulate FA turnover (Chandrasekar et al., 2005; Saunders et al., 2006), and to activate vinculin (Bakolitsa et al., 1999). This latter model was challenged by mutational analyses, which revealed that vinculin mutants that cannot bind to PIP2 still localize to FAs (Chandrasekar et al., 2005) and by the structure of full-length vinculin, which established that the basic patch thought to

Abbreviations used in the paper: CC, correlation coefficient; FA, focal adhesion; MEF, mouse embryonic fibroblast; PC, phosphatidylcholine; PIP2, phosphatidylinositol 4,5-bisphosphate; VBS, vinculin binding site; VH, vinculin head; Vt, vinculin tail.

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the formation of dimers, trimers, and higher-order oligomers (Hüttelmaier et al., 1998; Bakolitsa et al., 1999; Witt et al., 2004; Palmer et al., 2009). Although cross-linking and limited proteolysis analyses have suggested that self-association, PIP_2-induced dimerization, and F-actin–induced dimerization of vinculin are distinct (Bakolitsa et al., 1999; Johnson and Craig, 2000), extensive mutagenesis studies (Weekes et al., 1996; Johnson et al., 1998; Bakolitsa et al., 1999; Johnson and Craig, 2000; Saunders et al., 2006; Diez et al., 2008, 2009; Palmer et al., 2009; Wirth et al., 2010) have failed to reveal the PIP_2 binding sites.

Here we report the oligomeric structure of human vinculin in complex with PIP_2 at a 3.2-Å resolution. This structure and analyses of bona fide PIP_2-binding defective mutants establish essential functions for this signaling lipid in driving protein oligomerization, and in modulating interactions with the actin cytoskeleton and RNA-binding protein raver1. Importantly, this structure lays the foundation for understanding the roles of signaling lipids in other cytoskeletal proteins.

| Statistics | Overall | Last shell |
|------------|---------|------------|
| Resolution | 45.17–3.20 Å | 3.37–3.20 Å |
| No. reflections (working set) | 18,212 | 2,594 |
| No. of reflections (test set) | 967 | 144 |
| R_{work}/R_{free} | 0.206/0.253 | 0.235/0.277 |
| No. atoms | | |
| Protein | 8,090 | |
| PIP_2 | 141 | |
| Solvent | 75 | |
| B-factors | | |
| Protein | 147 Å^2 | |
| PIP_2 | 235 Å^2 | |
| Solvent | 92 Å^2 | |
| Root mean square deviations | | |
| Bond lengths | 0.010 Å | |
| Bond angles | 1.11° | |
| Ramachandran plot | | |
| Favored | 96.95% | |
| Allowed | 3.05% | |
| Generously allowed | 0% | |
| Outliers | 0% | |

CC, correlation coefficient.

aPrecision-indicating merging R factor (Weiss and Hilgenfeld, 1997; Weiss, 2001).

bPearson CC calculated between the average intensities of each random half of measurements of unique reflections (Karplus and Diederichs, 2012).

cCC* estimates the value of the true level of signal CC_{true} (the correlation of the averaged and thus less noisy data set with the noise-free true signal), based on a finite-size sample (Karplus and Diederichs, 2012).
Results

Architecture of PIP$_2$-bound vinculin oligomers

Full-length vinculin is monomeric in solution, and, once activated, vinculin oligomerizes at sites of FAs (Hüttelmaier et al., 1998). Binding of the inositol head group and the hydrophobic acyl chain poses difficulties in generating PIP$_2$-bound protein crystals. Indeed, the Vt–PIP$_2$ complex crystal structure reported here represents only the third nonmembrane protein–PIP$_2$ structure (out of five in total; Saad et al., 2006; Hansen et al., 2011; Kono et al., 2013; Poon et al., 2014). Further, the propensity of Vt to self-associate posed difficulties. We initially obtained ~5 Å diffraction, which could only be improved to 3.2 Å (space group $P3_121$; $a = b = 102.58$ Å, $c = 190.77$ Å) by using an R1060A mutation in the presence of a shorter chain (C8) derivative of PIP$_2$ (Tables 1 and 2). The Arg-1060 side chain is solvent exposed and residues $>$13 Å from the PIP$_2$ binding site (Fig. 1 A), and this alanine substitution has no effect on the conformation of vinculin nor upon the data and conclusions presented herein. Consistent with proteolysis studies (Bakolitsa et al., 1999), in the Vt–PIP$_2$ structure, one PIP$_2$ ligand is sandwiched between two Vt molecules by interacting with the loop connecting the first $\alpha$-helices, H1 and H2 (Fig. 1 B). Dimer formation is mediated by the interactions of the 4' and 5' phosphates with Vt (molecule “I”) Lys-915 and Lys-1061, and with Ser-913 through a water-mediated interaction. Lys-915 is also within hydrogen bonding distance to the carbonyl of the inositol. PIP$_2$ bridges one Vt (I) molecule to another Vt (II), where the carbonyl of the glycerol backbone and of the inositol reside within hydrogen bonding distance of Lys-915 and where there is a water-mediated interaction of the carbonyl of the glycerol backbone with Ser-913 (Fig. 1, C–E). Further, PIP$_2$ directs trimer formation, where the sandwiched PIP$_2$ interacts with $\alpha$-helix H3 of a third

Figure 1. PIP$_2$ directs vinculin oligomerization by unfurling the N terminus of Vt. (A) The R1060A mutation used for crystallization does not affect PIP$_2$ (oxygen, phosphor, and carbon atoms are shown as red, orange, and white sticks, respectively) binding. Vt $\alpha$-helices are colored sequentially across the color spectrum from red (H1) to blue (H5), monomers are labeled as I and II by binding to the $\alpha$-1-$\alpha$-2 loops and the C terminus of Vt. (C–E) Close-up view of the PIP$_2$-directed oligomerization of Vt ($\alpha$-helices and bonds are colored sequentially across the color spectrum from red to blue). Residues Lys-914 and Arg-945 are contributed from a third (III) monomer within the trimer. (D and E) View of the second (D; shown in blue) and third (E; shown in green) PIP$_2$ in the asymmetric unit in the same orientation as in C, which indicates the specificity. (F) The PIP$_2$-directed Vt trimer. Vt $\alpha$-helices are colored sequentially across the color spectrum from red to blue, PIP$_2$ is shown as spheres, and the three Vt molecules are labeled I–III. (G) With the exception of the loop connecting the last $\alpha$-helix H5 to its C-terminal extended coil, unbound Vt (shown in cyan) superposes onto PIP$_2$-bound Vt with a root-mean-square deviation (RMSD) of <0.5 Å for ~970 atoms (depending on the monomer in the asymmetric unit). However, the PIP$_2$ binding site coincides with the N-terminal extended coil (in particular residues 881–886, indicated by a red “N”), which is unfurled in the PIP$_2$-bound structure.
Figure 2. Pull-down assay to analyze the lipid-binding capacity of vinculin. (A) The interaction of wild-type and mutant Vt with VH was analyzed by electrophoretic mobility shift assays on a native gel, incubating the respective Vt proteins with VH. All proteins formed a Vt–VH complex. Data shown are representative of three independent experiments. (B) Unilamellar vesicles of the phospholipids PC and 15% PIP\(_2\) were incubated with wild-type and mutant Vt (top: K915Q, K944Q/R945Q, K970Q, R1057Q, and K1061Q; bottom: K915Q/K1061Q, K944Q/R945Q/K1061Q, and K915Q/K944Q/R945Q) and sedimented. Supernatant (S) and pellet (P) fractions were detected on Coomassie-stained SDS-PAGE gel. Inactive and talin-VBS3 (residues 1945–1970) activated full-length (FL) vinculin were run as the nonbinding and binding controls, respectively. Densitometric values averaged over three experiments together with their standard deviations are shown above or below the respective experiments. (C) Previously identified residues on \(\alpha\)-helix H3 reside near the PIP\(_2\)-induced Vt–Vt interface. Shown is a close-up view of the PIP\(_2\)-induced Vt–Vt interface. Red and blue spheres represent positive and negative charges, respectively. Residues Lys-952, Lys-956, and Arg-963, which were previously thought to bind to PIP\(_2\), reside near or at the Vt–Vt interface, which might explain the decreased binding of the potentially oligomer-defective mutant (K952Q, K956Q, and R963Q) Vt compared with wild-type Vt.

**Figure 2**

![Image of Figure 2](image-url)
In the context of the full-length vinculin structure, the PIP$_2$ binding site is on the surface of the protein and is distinct from the head–tail interface. However, PIP$_2$ binding to full-length vinculin is impaired (Fig. 2 B and Fig. S1 D) because the N-terminal extended coil occludes the binding site (Fig. 3 A). Indeed, truncation of residues 879–897 increases Vt binding to PIP$_2$ approximately twofold (Palmer et al., 2009). In contrast, talin VBS-activated full-length vinculin is fully permissive for lipid binding (Fig. 2 B and Fig. S1 D). Our structural and biochemical data fully support a PIP$_2$-directed full-length vinculin dimer as a building block for higher-order vinculin oligomers (Fig. 3 B).

The PIP$_2$ and F-actin binding sites of vinculin are distinct
Activated vinculin binds directly to F-actin (Jockusch and Isenberg, 1981; Wilkins and Lin, 1982) and caps actin filaments to promote their polymerization (Ramarao et al., 2007; Le Clainche et al., 2010). This is thought to play critical roles at the leading edge of migrating cells (Dominguez and Holmes, 2011). Both PIP$_2$ and F-actin bind to the Vt domain, and PIP$_2$ binding has been reported to inhibit F-actin binding (Steimle et al., 1999). We therefore tested the structure-based PIP$_2$-binding–deficient mutants of vinculin for their ability to interact with F-actin. Notably, all PIP$_2$-binding deficient mutants cosedimented with F-actin (Fig. 4 A and Fig. S2), and also bundled F-actin, as seen with wild-type vinculin (Fig. 4 B and Fig. S3). Although the Vt F-actin binding sites have been hotly debated (Shen et al., 2011; Janssen et al., 2012; Thompson et al., 2013, 2014), the recently characterized I997A and V1001A mutants do show decreased binding to F-actin (Thievessen et al., 2013). These residues reside >12 Å from the PIP$_2$-binding site (Fig. 4 C). Thus, the PIP$_2$ and F-actin binding sites of vinculin are distinct.

The raver1–vinculin complex does not bind to PIP$_2$

The RNA-binding protein raver1 simultaneously binds to vinculin mRNA and the Vt domain of activated vinculin, and this interaction has thus been postulated to control local production of vinculin to facilitate FA assembly (Lee et al., 2009, 2012; Madl and Sattler, 2009; Rangarajan et al., 2011). Interestingly, Arg-945 that binds PIP$_2$ is also involved in electrostatic interactions with the raver1 residue Glu-120 (Fig. 5, A and B). To test if the lipid-binding–deficient K944Q/R945Q mutant binds raver1 and if the raver1–Vt complex binds PIP$_2$, we ran the wild-type and mutant Vt–raver1 complexes on a size exclusion chromatography column (Fig. 5, C and D) and performed lipid cosedimentation assays (Fig. 5 E). Interestingly, although the K944Q/R945Q vinculin mutant still bound to raver1, the complex eluted earlier than the wild-type Vt–raver1 complex. Further, the crystal structures suggested that the raver1 and PIP$_2$-binding sites are overlapping (Fig. 5, A and B), and indeed the Vt–raver1 complex did not bind to PIP$_2$ (Fig. 5 E). Thus, although Vt–raver1 can form a ternary complex with F-actin (Lee et al., 2009), the Vt–raver1 complex does not bind to the membrane.

The vinculin–PIP$_2$ interaction affects the actin cytoskeleton, cell migration, and FA turnover

PIP$_2$ binding to vinculin has been proposed to play roles in the turnover of FAs (Chandrasekar et al., 2005; Saunders et al., 2006). Vinculin-null mouse embryonic fibroblasts (MEFs) have few and very unstable FAs, and those present fail to form links to the actin cytoskeleton, which is disorganized in
Figure 4. The lipid-binding–deficient vinculin mutants bind and bundle F-actin. (A) Wild-type and mutant Vt (top: K915Q, K944Q/R945Q “944,” K1061Q, K915Q/K944Q/R945Q “915 944,” K915Q/K1061Q; K944Q/R945Q/K1061Q “944 1061”; bottom: K924Q, K970Q, and R1057Q) were cosedimented with F-actin and analyzed on Coomassie-stained SDS-PAGE. Full-length (FL) vinculin was run as the nonbinding control. S, supernatant; P, pellet. The white lines indicate the removal of unrelated lanes for presentation purposes. Data shown are representative of three independent experiments. The corresponding full-length gels are shown in Fig. S2 (A–C). (B) Confocal images of TRITC-phalloidin–labeled F-actin in the absence or presence of wild-type or mutant (K1061Q or K915Q/K944Q/R945Q/K1061Q) Vt domain. Data shown are representative of three independent experiments. Larger micrographs corresponding to these panels are shown in Fig. S3. Bars, 20 µm. (C) Model based on the superposition (not depicted) of the cryo-EM Vt–F-actin structure (Thompson et al., 2014; F-actin subunits are shown in gray or black) onto the PIP2-bound Vt dimer (α-helices colored sequentially across the color spectrum from red [H1] to blue [H5], monomers are labeled I–III, and raver1 is shown in gray). Superposition of lipid-bound Vt (I) onto the raver1-bound Vt shows overlapping binding of Vt (III) and raver1. (B) Close-up view of some interface residues. In its raver1-bound state, vinculin residue Arg-945 engages in interactions with raver1 residues Glu-120 and Arg-117, whereas in its lipid-bound state, Arg-945 points toward PIP2. (C) Size exclusion chromatogram (SEC) of Vt–raver1 (blue line; peak #4 eluting at 14.56 ml; apparent mol wt = 54.6 kD; polypeptide chain mol wt = 51.2 kD), Vt-K944Q/R945Q/raver1 (black line; peak #5, 15 ml; apparent mol wt = 45 kD; polypeptide chain mol wt = 51.2 kD), raver1 (dotted line; peak #3, 15.77 ml; apparent mol wt = 32.19 kD; polypeptide chain mol wt = 31.5 kD), and Vt (gray line; peak #1, 17.2 ml; apparent mol wt = 17.22 kD; polypeptide chain mol wt = 19.7 kD). (D) SDS-PAGE of peak fractions from the SEC shown in the previous panel. The lane numbers correspond to the peak numbering in the previous panel. (F) Lipid cosedimentation assay of the Vt–raver1 complex. The Vt–raver1 complex does not cosediment with PIP2 and remains in the soluble (S) fraction. P, pellet.
with the notion that the PIP<sub>2</sub>–vinculin interaction might play roles in FA turnover.

Direct measurement of wild-type GFP-vinculin turnover in FAs by FRAP revealed that the entire population within the FA is turning over with a half-time of ~20 s (Fig. 8), a value in the range of what was previously reported (Chandrasekar et al., 2005). However, FRAP analysis of the lipid-binding–deficient vinculin mutants revealed that a large fraction (~45% for the K1061Q mutant and >90% for the K944Q/R945Q mutant) of the GFP-vinculin within these FAs is immobile or much more tightly bound in that it did not recover within the time frame that results in complete recovery for the wild-type protein (Fig. 8). Notably, the mobile fraction of the K1061Q mutant vinculin in FAs displayed recovery kinetics that were significantly faster than wild-type vinculin, which is indicative of a much lower binding affinity. The immobile or tightly bound fraction may represent static nonfunctional interactions within the FA, which is consistent with the inability of these cells to organize the actin network and suggest that the dynamic turnover of vinculin is compromised in the PIP<sub>2</sub>-binding–deficient mutants. We conclude that the PIP<sub>2</sub>–vinculin interaction plays important roles in the stabilization and dynamic turnover of FAs.
and activate vinculin to stabilize nascent cell–matrix adhesions via its associations with F-actin (Galbraith et al., 2002; Giannone et al., 2003; Zaidel-Bar et al., 2003). The role of PIP$_2$ binding in the regulation and function of vinculin has been hotly debated (Gilmore and Burridge, 1996; Bakolitsa et al., 2004;)

**Discussion**

The binding of integrins to extracellular matrix components at FAs triggers mechanical force on associated talin molecules, which opens up helix bundles and exposes VBSs that then bind to and activate vinculin to stabilize nascent cell–matrix adhesions via its associations with F-actin (Galbraith et al., 2002; Giannone et al., 2003; Zaidel-Bar et al., 2003). The role of PIP$_2$ binding in the regulation and function of vinculin has been hotly debated (Gilmore and Burridge, 1996; Bakolitsa et al., 2004;)

*Figure 7. MEFs expressing the PIP$_2$-binding-deficient mutant of vinculin are impaired in wound closure. Vinculin-null cells (left) that were engineered to express comparable levels of wild-type vinculin (center left) or K944Q/R945Q or K1061Q mutant vinculin (center right and right) were plated onto fibronectin-coated 12-well plates. Monolayers were then wounded, and the extent of cell migration into the wound was recorded at 0, 4, 8, and 12 h. Vinculin-null cells migrate in a rapid yet chaotic fashion, whereas wound closure in these cells expressing wild-type vinculin is an ordered process. Vinculin-null cells expressing K944Q/R945Q or K1061Q mutant vinculin had reduced migration and wound closure versus cells expressing wild-type vinculin. Data shown are representative of three independent experiments. Bars, 20 µm.*
Izard et al., 2004; Bois et al., 2006; Chen et al., 2006). Our studies establish that the PIP2 binding site is occluded in full-length vinculin and they demonstrate that PIP2 binding is necessary to mediate interactions of vinculin with F-actin at FAs. Our studies also suggest that F-actin and PIP2 binding to vinculin is mutually permissive, as they clearly have distinct binding sites on vinculin. These findings are consistent with a model whereby PIP2 binding triggers the formation of higher-order oligomers of activated vinculin that are competent to make multiple contacts with F-actin that would stabilize FAs, particularly because the cryo-EM structure suggests that one molecule of vinculin can be sandwiched between two molecules of F-actin (Janssen et al., 2006).

The crystal structure reveals that PIP2 principally affects vinculin function by directing vinculin oligomerization. Specifically, PIP2 directs the formation of vinculin dimers by releasing a flexible N-terminal extended coil present in the Vt five-helix bundle, and by bridging the flexible C terminus of one Vt molecule with α-helix H1 of a second Vt molecule, thereby restraining the conformation of the C terminus and stabilizing the dimer. Further, this signaling lipid directs vinculin trimerization via interactions with two lysine residues on α-helix H3 that are at the head–tail interface in inactive vinculin. Other basic residues on H3 have been proposed to direct the vinculin–lipid interaction (Chandrasekar et al., 2005), and examination of our crystal structure reveals that these residues are rather near the lipid-induced Vt–Vt interface (Fig. 2 C). Thus, mutating these residues might have prevented this Vt–Vt association by hindering vinculin oligomerization, and this might explain the decreased lipid binding. Finally, the crystal structures explain how higher-order vinculin oligomers can occur after PIP2 binding, a response that allows amplification of the interactions of vinculin with talin, and with F-actin, at FAs.

The RNA-binding protein raver1 has been proposed to regulate the local assembly of nascent FAs by binding to mRNAs encoding components of FAs in order to allow their localized translation at these sites. Specifically, raver1 binds to both Vt and vinculin mRNA, and the ternary Vt–raver1–vinculin mRNA complex is permissive for binding to F-actin. In contrast, the raver1 and PIP2 binding sites on vinculin overlap, and our studies have shown that the vinculin–raver1 complex does not bind to PIP2. This is consistent with a model where raver1 binding to vinculin is spatiotemporally regulated and where the F-actin–vinculin–raver1 complex bound to vinculin mRNA is not attached to the plasma membrane.

While the F-actin binding sites on vinculin are also controversial (Shen et al., 2011; Janssen et al., 2012; Thompson et al., 2013; Thompson et al., 2014), the recently identified measurements for each cell line are displayed. The t50 values for wild-type (19 s) and the K1061Q mutant (7 s) were determined by fitting the curves to a single exponential. A t50 value for the K944Q/R945Q mutant could not be determined due to the absence of significant recovery. The immobile fraction is represented by the difference between the normalized plateau value and the prebleach fluorescence, and was estimated to be <5% for the wild-type cells, ~45% for the K1061Q mutant, and >90% for the K944Q/R945Q mutant. Error bars indicate SD.
I997A and V1001A mutants showed decreased F-actin binding. Notably, the crystal structure of the vinculin–PIP₂ complex establishes that Ile-997 and Val-1001 are not involved in the PIP₂ binding sites of vinculin. Further, PIP₂–binding–deficient mutants of vinculin still bind F-actin in vitro; thus, binding of PIP₂ and F-actin to vinculin seems mutually permissive. Finally, these findings suggest similar mechanisms of action for this signaling lipid in controlling the structure and functions of other lipid-and-actin-binding cytoskeletal proteins, where we propose that PIP₂ may play roles in oligomerization of these targets that are also permissive for binding to F-actin.

We present three observations suggesting that binding of PIP₂ by vinculin is an integral component of proper FA function. The exogenous expression of wild-type GFP vinculin, but not lipid-binding–deficient mutants, in vinculin-null MEFs resulted in the reestablishment of a well-organized actin network. Vinculin-null MEFs expressing lipid-binding–deficient GFP–vinculin displayed a severe impairment of cell migration in wound closure assays. Finally, direct measurement of vinculin recruitment to FAs by photobleaching revealed that wild-type vinculin displays dynamic turnover, with the entire population being replaced within the time frame of minutes. However, we find that a large portion of the PIP₂–binding–deficient mutant vinculin is immobile or tightly bound, possibly due to static nonfunctional interactions within the FAs. Thus, although lipid-binding–deficient vinculin does localize to FAs, the observations noted here would suggest that the interactions within these FAs are distinct from those of wild-type vinculin and that these differences compromise FA function, which is consistent with the inability of these cells to organize the actin network or to close a wound. Previously, a modified FRAP procedure was used to show that a PIP₂–binding–defective vinculin mutant was recruited to FAs, but disassembly was depressed (Chandrasekar et al., 2005). In a subsequent study, interference reflection microscopy was used to demonstrate that FAs in cells expressing a lipid-binding–deficient mutant displayed reduced turnover and translocation (Saunders et al., 2006). The results from these studies are in agreement with our conclusion that the PIP₂–vinculin interaction is critical for the dynamic turnover of FAs and that this interaction is compromised in the PIP₂–binding–deficient mutants. Our vinculin–PIP₂ structure and functional data will aid future studies to understand how lipid-binding–deficient vinculin mutants affect the actomyosin contractile forces as these mutants significantly disturb the F-actin network.

Materials and methods

Protein preparation

Full-length human vinculin cDNA (residues 1–1,066) was cloned into a pET3 expression vector, and a noncleavable His₆ tag was added downstream to the vinculin coding sequence. Modified pET-3 vector was used for cloning the head domain of vinculin (VH; residues 1–843) with a PreScission protease cleavable C-terminal hexahistidine tag. This expression vector contains a T7 promoter and contains an ampicillin resistance gene. For vinculin and VH purification, cell pellets were resuspended in 20 mM Tris-HCl, pH 8, and 150 mM NaCl, lysed by sonication, and clarified by ultracentrifugation (100,000 g for 45 min). The supernatant was applied onto a GSTrap FF column, and, after extensive washing with the buffer (20 mM Tris-HCl, pH 8, and 400 mM NaCl), the bound GST-VH was eluted with 20 mM of reduced glutathione, pH 8. GST affinity tags were removed by overnight cleavage with PreScission protease at 4°C and dialyzed into 20 mM Tris, pH 8, 400 mM NaCl, 1 mM DTT, and 0.1 mM EDTA. All proteins were further purified by gel filtration using a 26/60 Superdex 75 column (GE Healthcare) that was preequilibrated with 20 mM Tris-HCl, pH 8, 400 mM NaCl, 1 mM DTT, and 0.1 mM EDTA. Peak fractions were collected and concentrated to 10 mg/ml.

VH and full-length vinculin were purified as described previously (Rangarajan et al., 2010) and dialyzed against 20 mM Tris, pH 8, 10 mM DTT, 1 mM EDTA, and 1 mM EGTA, and concentrated to ~29 mg/ml. Talin-VBS3 (resides 1,945–1,970) was synthesized.

The RRM1-3 (residues 39–321) domains of human raver1 were cloned into the pET28a vector (EMD Millipore) to produce N-terminal hexahistidine–tagged proteins. This expression vector has a 17 promoter and six-histidine residue coding sequence upstream of the RRM1-3 (residues 39–321) domain coding sequence. This bacterial expression vector contains a kanamycin resistance gene. RRM1-3 was expressed in Escherichia coli strain BL21-DE3 (Invitrogen). Bacteria were grown at 37°C in Luria-Bertani medium containing 20 µg/ml kanamycin. Bacterial cultures were induced at OD₆₀₀ = 0.8 by adding isopropyl-β-thiogalactopyranoside to 0.5 mM and incubating at 20°C, and cells were harvested after 24 h. The cell pellet was resuspended in 20 mM Tris-HCl, pH 8, and 150 mM NaCl, lysed by sonication, and clarified by ultracentrifugation (100,000 g for 45 min). The cell lysate was loaded onto a HiTrap chelating nickel affinity column (GE Healthcare), which was equilibrated with 20 mM Tris·HCl, pH 8, and 150 mM NaCl, and we eluted the proteins using an imidazole gradient. Further, it was purified by gel filtration using a Superdex 75 column (GE Healthcare) equilibrated in 20 mM Tris·HCl, pH 8.0, and 150 mM NaCl. Protein fractions were collected and concentrated to 15 mg/ml.

VH–PIP₂ crystallization

Human VI (residues 891–1,066) was incubated with 20 µM PIP₂ dioleoyl (Avanti Polar Lipids) on ice for 30 min. Excess precipitate was removed by centrifugation (15,800 g for 25 min at 4°C). We screened ~1,000 commercially available crystallization conditions and obtained a crystallization hit from the P41 plate kit of Hampton Research. Crystals were grown within 2 d by hanging drop vapor diffusion from 50 mM Hepes, pH 7, 12% (wt/vol) PEG 3350, and 2% (wt/vol) Tryptone, at 20°C. They diffracted x-rays at various beam lines (11-1 at Stanford Synchrotron Radiation Laboratory [SSRL], or 22-ID and 22BM at the Advanced Photon Source of the Argonne National Laboratory [APS/ANL]) to ~5 Å Bragg spacings. All crystallization strategies and truncations failed to improve the diffraction, but ultimately, crystals of the PIP₁–bound binding–deficient mutant proteins were obtained with 5% increments of glyc erol to 25% and stored in liquid nitrogen at 100 K.

X-ray data collection and processing

X-ray diffraction data were collected at the Advanced Photon Source 22-ID beam line (wavelength of 1 Å) and integrated and scaled using XDS and SCALA as implemented in the CCP4 suite (Winn et al., 2011). Data reduction statistics are provided in Table 1.
Structure determination and crystallographic refinement
Phases were obtained by molecular replacement using the program MOLREP (Vagin and Teplyakov, 1997), where the search model consisted of Vt from the Vh1–Vt complex (Protein Data Bank [PDB] accession no. 1RKE). A clear solution with six molecules in the asymmetric unit was obtained in space group P3_{1}2_{1}2_{1}, while for the enantiomer in P3_212, the R factor remained >0.5 during refinement. An acceptable packing density, Vm (Matthews, 1968), was calculated at 2.45 A\(^{3}\)/Da, corresponding to a solvent content of ~0.495.

Initial crystallographic refinement used a 3 Å cutoff, but best results were obtained using data up to 3.2 Å Bragg spacings. Maximum likelihood crystallographic refinement was performed using autoBUSTER (Bricogne et al., 2011) by imposing target restraints using the high-resolution structure. The model was improved by local noncrystallographic symmetry restraints through local structure similarity restraints (LSLR; Smart, O.S., Brandl, P. Flensburg, W. Keller, C. Paciorek, C. Vonrhein, T. Womack, and G. Bricogne. 2008. Proceedings of the annual meeting of the American Crystallographic Association. Abstr. TP139). Iterative cycles of model building were performed using Coat (Emesly and Cowtan, 2004), and model bias was minimized by building into composite omit maps. Optimized ligand coordinates and ligand restraints were obtained from the Grade web server (grade.globalphasing.org). The electron density map was sharpened using Coat (Emesly and Cowtan, 2004) to ensure the directionality and identity of the α-helices. The quality of the final model was assessed using MolProbity (Davis et al., 2004), which revealed no outliers with >0.03 of the amino acid residues in the favored region of the Ramachandran plot. Crystallographic refinement statistics are provided in Table 2. The final model contains six Vt polypeptide chains and three Pip2 molecules. Residues 1,051–1,053 and 1,066 are missing due to weak electron density in this region.

Lipid cosedimentation assay
Pip2 binding to Vt mutants was assayed as described previously (Palmer et al., 2009). In brief, lipid vesicles of phosphatidylcholine (PC) and Pip2 were prepared by mixing together in chloroform to have a final composition of 85% PC and 15% Pip2. The lipid mixture was vacuum dried and re-suspended in 20 mM Tris, pH 8, 400 mM NaCl, and 3 mM DTT. Unilamellar vesicles were then produced by sonication after incubating the reaction mixture was pelleted at 125,000 g for 1 h at 4°C for 1 h followed by centrifugation at 75,815 g for 1 h. The supernatant and pellet were separated, and the pellet was washed two times with buffer A and resuspended in 12.5 µl. The supernatant and pellet samples were analyzed by SDS-PAGE.

Native PAGE assay of Vt proteins binding to the VH domain
About 5 µM Vh (residues 1–840) was mixed with 2–3-fold excess of wild-type or mutant (K915Q, K944Q/R945Q, K1061Q, K915Q/K944Q/R945Q, and K944Q/R945Q/K1061Q) Vt (residues 891–1,066) in 20 mM Tris, pH 8, 400 mM NaCl, and 1 mM DTT. Samples containing 50 µg of total lipid in 10 µl suspension and 40 µM Vt (wild type or mutants) were incubated at 4°C for 1 h followed by centrifugation at 75,815 g for 30 min. The supernatant and pellet were separated, and the pellet was washed two times with buffer A and resuspended in 12.5 µl. The supernatant and pellet samples were analyzed by SDS-PAGE.

F-actin cosedimentation assay
F-actin bundling assay
Protein bands with Coomassie blue.

F-actin cosedimentation assay was performed as described previously (Palmer et al., 2009). In brief, F-actin was incubated for 1 h at 4°C with wild-type or K944Q/R945Q mutant vinculin (200 µM Tris, pH 8, 400 mM NaCl, and 1 mM DTT) in a 1:2 molar ratio and run on an analytical Super 2000 size exclusion chromatography column (equilibrated in 20 mM Tris, pH 8, 400 mM NaCl, and 3 mM DTT) with a flow rate of 0.5 ml/min. Vt and raver1 proteins, both at 100 µM, were also run separately. Fractions were analyzed on a 10–15% gradient PHAST gel with SDS-PAGE buffer strips.

Cell culture and immunofluorescence
Vinculin-deficient (Vinc−/−) MEFs were provided by K. Burridge (University of North Carolina, Chapel Hill, NC). They were originally prepared by E. Adamson (Sanford-Burnham Medical Research Institute, La Jolla, CA): vinculin-null cells were isolated by targeting both alleles of the vinculin gene by homologous recombination in which exon 3 has been replaced by a neo cassette resistance gene (Kuo et al., 1998). Vinculin-null cells were cultured in DMEM containing 4.5 g/liter glucose and 2 mM l-glutamine (Invitrogen) supplemented with 10% FBS, 100 U/ml penicillin, and 100 µg/ml streptomycin at 37°C and 5% CO\(_2\) in a humidified incubator, and passed every 3–4 d. Vinc+/− MEFs were transfected with the pcDNA1 plasmid DNA encoding full-length human wild-type or mutant vinculin as an N-terminal His6-EGFP fusion, or with the vector control, using Lipofectamine LTX plus reagent (Invitrogen) according to the manufacturer's protocol. All human vinculin constructs and Pip2-binding-deficient mutants used for cell biology studies were cloned into pcDNA4.1/TO/MycHisB vector (Life Technologies). The basic feature of these plasmids is a CMV promoter and two tandem tetracycline (Zn2+ Tetr) operator sequences upstream of the coding sequence of EGFP. EGFP coding sequence and a PreScission cleavage site (LEVLFQGP) downstream to it was introduced between the TET operator and the coding sequence of HV or the mutants. Further, a three-amino linker was introduced in-between the PreScission cleavage site and the coding sequence of vinculin. The vinculin coding sequence is terminated by a stop codon (TGA) before the myc and hexahistidine tag. This mammalian expression vector contains Zeocin and ampicillin resistance genes. Stables were prepared using Zeocin. Cells were grown on six-well tissue culture treated plates for 48 h and were initially selected with medium containing 800 µg/ml Zeocin for 1–2 wk, then passed every 3–4 d with fresh media containing 200 µg/ml Zeocin.

Preparation of cells for immunofluorescence detection for localization of vinculin in FAs was performed by plating the cells stably expressing vinculin or mutants of vinculin (VH, K944Q/R945Q) or on a gelatin-coated coverslip. 24 h after plating, the cells were fixed for 10 min with 4% paraformaldehyde in PBS, pH 7.4, and permeabilized with 0.2% Triton X-100 in PBS. Blocking was performed with 1% BSA in PBS for 20 min and cells were stained with TRITC-phalloidin (5 µg/ml) for actin and with DAPI (0.1 µg/ml) for detecting nuclei. Additionally, cells (7 × 10^4) were also seeded on 12-mm coverslips coated with 20 µg/ml fibronectin (Sigma-Aldrich) and incubated for 6 h. To visualize colocalization of vinculin and paxillin (another FA marker), cells were fixed and stained sequentially with mouse anti-paxillin antibody (BD) and Alexa Fluor 568 conjugated anti–mouse IgG (Life Technologies) followed by staining with DAPI for visualizing the nucleus. For imaging, the coverslips were mounted on a glass slide using Vectashield mounting medium (Vector Laboratories).

Fluorescence images were collected on confocal laser scanning microscope (LSM 780; Zeiss), controlled with Zen software, using a GaAsP detector. Cells were randomly selected and imaged at room temperature (23°C) using either a 63×/1.3 NA Apochromat multi-immersion objective lens using water as the imaging medium, and further adjusted for brightness/contrast with Fiji (ImageJ).

Size exclusion chromatography
Purified raver1 (100 µM in 20 mM Tris, pH 8, and 150 mM NaCl) was incubated for 1 h at 4°C with wild-type or K944Q/R945Q mutant vinculin (200 µM Tris, pH 8, 400 mM NaCl, and 1 mM DTT) in a 1:2 molar ratio and run on an analytical Super 2000 size exclusion chromatography column (equilibrated in 20 mM Tris, pH 8, 400 mM NaCl, and 3 mM DTT) with a flow rate of 0.5 ml/min. Vt and raver1 proteins, both at 100 µM, were also run separately. Fractions were analyzed on a 10–15% gradient PHAST gel with SDS-PAGE buffer strips.

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Immunoblotting

Cells showing stable and comparable levels of expression of vinculin or vinculin mutants were cultured to 70–80% confluency on gelatin-coated 10-cm Petri dishes in media supplemented with 200 µg/ml Zeocin, and were collected through trypsinization. The cells were washed twice with ice-cold PBS and resuspended in RIPA buffer (10 mM Tris-HCl, pH 8, 140 mM NaCl, 1 mM EDTA, 0.5 mM EGTA, 1% Triton X-100, 0.1% sodium deoxycholate, 0.1% SDS, 1 mM PMSF, and protease inhibitor cocktail [Roche]) and briefly sonicated. The lysed samples were then separated by SDS-polyacrylamide gel, electroblotted on to Immobilon membrane (EMD Millipore) using a semidry transfer apparatus (Bio-Rad Laboratories), and incubated sequentially with anti-vinculin primary antibody and HRP-conjugated secondary antibody. The blots were then developed after treatment with chemiluminescence substrate [Thermo Fisher Scientific] according to the manufacturer’s instructions.

Wound assays

Vinculin-null MEFs and the cells stably expressing full-length wild-type or mutant vinculin were seeded at a density of 2 × 10⁶ cells/well on a 12-well plate, precoated with 20 µg/ml fibronectin. The cells were then incubated overnight at 37°C and 5% CO₂ to achieve confluency the following day. The wound healing assay was initiated by scraping a line on the culture dish using a 200-µl pipette tip on the confluent cell monolayer. The culture dish was then gently washed with PBS to remove any cell debris and replenished with 1 ml of fresh media (DMEM with 10% bovine calf serum, 100 U/ml penicillin, and 100 µg/ml streptomycin). Cells were imaged on an EVOS FL cell imaging system [Life Technologies] equipped with a color charge-coupled device image acquisition system (ICX285AQ; Sony) through software built in to the onboard microprocessor. A 10x/0.25 dry objective lens was used for bright field imaging immediately afterwards in order to represent the zero time point, and the wound healing was monitored every 4 h, returning plates for incubation after every imaging session. All assays were repeated in triplicate. Each time, images were taken at six different points and the positions that were similar in wound distance were used for direct comparison.

FRAP assays

MEF cells were plated and observed in fibronectin-coated 35-mm glass-bottom culture dishes [MatTek Corporation]. FRAP was performed on a laser scanning confocal microscope [TCS SP8; Leica] using the 488-nm line of an argon laser essentially as described previously (George et al., 2010). All experiments were performed at 37°C, and imaging was performed with a Plan-Apochromat 63x/1.3 NA objective lens using the FRAP module of the LAS AF software (Leica). Bleaching of a single FA per cell was achieved using a single bleach scan of 273-ms duration at maximal laser power. Recovery was monitored for 500–575 nm and recorded for 90 s. Although movement of cells and/or FAs precluded longer measurements, a recovery plateau was reached within this time period for most of the imaged cells, except where noted. Identical imaging settings (5% laser intensity) were used for all pre- and post-bleach images. Scan rates in the bidirectional at the highest possible rate using a 4x zoom with a pinhole of 1 Airy unit. Image analysis was performed within the LAS AF software. Datasets were imported into Origin 6.0 and normalized recovery rates were fitted to a single exponential recovery model in OriginLab software using the equation \( F(t) = 1 - e^{-kt} \), where \( F(t) \) is the fluorescence at time \( t \), \( k \) is the rate constant, and the 1/2 recovery was calculated using \( k \ln(2)/t_{1/2} \). All datasets consisted of at least eight cells per experiment.

Online supplemental material

Fig. S1 shows the full gels shown in Fig. 2c. Fig. S2 shows the full gels shown in Fig. 4. Fig. S3 shows larger panels compared with those shown in Fig. 4 B as an additional confocal images of TRITC-phalloidin labeled F-actin in the presence of Tris-HCl), pH 8. Fig. S4 shows confocal immunofluorescence studies of exogenously expressed wild-type and lipid-binding-deficient mutant GFP-tagged vinculin colocalized with paxillin in FAs in vinculin-null MEFs, as well as immunoblots obtained with vinculin antibody after SDS-PAGE to analyze the expression of wild-type and mutant vinculin proteins in vinculin-null cells using the primary antibody an antihuman antibody and a secondary HRP-conjugated anti-mouse IgG. Online supplemental material is available at http://www.jcb.org/cgi/content/full/jcb.201404128/DC1.

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