α-Catenin homodimers are recruited to phosphoinositide-activated membranes to promote adhesion

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A unique feature of α-catenin localized outside the cadherin–catenin complex is its capacity to form homodimers, but the subcellular localization and functions of this form of α-catenin remain incompletely understood. We identified a cadherin-free form of α-catenin that is recruited to the leading edge of migrating cells in a phosphatidylinositol 3-kinase–dependent manner. Surface plasmon resonance analysis shows that α-catenin homodimers, but not monomers, selectively bind phosphatidylinositol-3,4,5-trisphosphate–containing lipid vesicles with high affinity, where three basic residues, K488, K493, and R496, contribute to binding. Chemical-induced dimerization of α-catenin containing a synthetic dimerization domain promotes its accumulation within lamellipodia and elaboration of protrusions with extended filopodia, which are attenuated in the α-cateninKKR<3A mutant. Cells restored with a full-length, natively homodimerizing form of α-cateninKKR<3A displayed reduced membrane recruitment, altered epithelial sheet migrations, and weaker cell–cell adhesion compared with WT α-catenin. These findings show that α-catenin homodimers are recruited to phosphoinositide-activated membranes to promote adhesion and migration, suggesting that phosphoinositide binding may be a defining feature of α-catenin function outside the cadherin–catenin complex.

Introduction

The cadherin–catenin adhesive complex is widely viewed as an important regulator of intercellular interactions. One challenge of studying this system is that adherens junction linkage components (i.e., catenins) can also localize to other parts of the cell, where they carry out distinct, “extrajunctional” functions. For example, p120 and β-catenin are not only important for adhesion but are critical cofactors for DNA-binding proteins that directly regulate cell fate decisions (McCrea and Gottardi, 2016). Thus, interpretation of catenin knockout studies are confounded by their multifunctionality, particularly given that they often engage both signaling and adhesive machinery via overlapping binding surfaces (McCrea et al., 2015).

A similar problem exists for understanding the roles of α-catenin (αCat), a filamentous F-actin–binding protein found as both junctional and extrajunctional forms (Schneider et al., 1993; Benjamin et al., 2010). αCat bound to the cadherin–catenin complex functions as a force-activated F-actin–binding protein (Buckley et al., 2014), while the epithelial isoform of αCat also exists as extrajunctional αCat in the cytosol and nucleus, where the cytoskeletal and signaling roles for αCat monomer, homodimer, and heterodimer (with β-catenin) are just emerging (Stewart and Nelson, 1997; Benjamin et al., 2010; Daugherty et al., 2014). Because αCat homodimerization is structurally incompatible with β-catenin binding (Koslov et al., 1997; Obama and Ozawa, 1997) and purified recombinant αCat homodimers show better binding to F-actin than monomeric αCat in solution (Drees et al., 2005), it was reasoned that extrajunctional αCat homodimers might serve a distinct, F-actin–regulating function. However, full understanding of this functional pool remains limited. Indeed, extrajunctional αCat can suppress in vitro actin assembly mediated by the branching protein Arp2/3 (Drees et al., 2005), possibly by...
competitive binding to actin filaments (Hansen et al., 2013), as well as antagonize lamellipodial activity in cells (Benjamin et al., 2010). However, these studies have not addressed the specific contribution of αCat homodimerization to these activities or epithelial cell behaviors. Although the relatively high $K_d$ of αCat homodimers (25–73 μM; Ishiyama et al., 2013; Pokutta et al., 2014) with respect to the estimated concentration of cytosolic αCat in cells (0.6 μM; Drees et al., 2005; Yamada et al., 2005) raises the possibility that a mechanism to increase the local concentration of αCat may be required to favor homodimerization in vivo, evidence that homodimer dissociation is kinetically blocked (Pokutta et al., 2014) could offset the need for a concentration mechanism. Nonetheless, the structural constraints of αCat homodimers clearly dictate a function outside the cadherin–catenin complex, but how and where this form of αCat is generated in the cell remain unknown.

Although αCat homodimerization is a conserved feature in Drosophila melanogaster, albeit not all αCats (Kwiatkowski et al., 2010; Dickinson et al., 2011; Desai et al., 2013; Miller et al., 2013), there are currently no minimal mutations that separate αCat heterodimerization with β-catenin from αCat homodimerization to generate a form of αCat that specifically lacks homodimer function, while sparing αCat binding to the cadherin–catenin complex. In this study, we make use of a pancadherin null cell line and a chemical-induced dimerization system to define the localization of extrajunctional αCat and function of αCat homodimers in epithelial cells. We also show that αCat contains a region of positively charged residues that contributes to phosphoinositide-dependent membrane recruitment and processes important for cell–cell adhesion.

Results

Cadherin-independent mechanism of αCat membrane recruitment

To visualize the properties of αCat outside of the cadherin–catenin complex, we took advantage of an established cell line (epidermoid A431D carcinoma cells) that completely lacks cadherins and detectable catenins (Lewis et al., 1997). Stable expression of GFP–αCat in scratch-wounded A431D cells revealed its recruitment to the leading edge of the wound front (Fig. 1 A), a phenomenon that was also observed in GFP–αCat–restored R2/7 cells, a DLD1 human colon cancer variant that expresses E-cadherin and β-catenin but lacks αCat expression and functional adhesions (van Hengel et al., 1997; Watabe-Uchida et al., 1998; Fig. 1, B and C). This pool of αCat colocalized with F-actin (Fig. 1, A and B) but not E-cadherin (Fig. 1 C), suggesting that F-actin may be a key functional target of extrajunctional αCat in cells.

Forced dimerization of αCat promotes its cortical recruitment

Because αCat homodimerization is thought to be a defining feature of extrajunctional αCat, we sought to assess the unique contributions of αCat homodimerization to its cellular localization and function. Toward this end, we used a temporal and reversible system to chemically induce homodimerization of the cytosolic pool of αCat (Fig. 1 D). We induced αCat homodimers by replacing the native, αCat N-terminal domain with the DmrB dimerization domain (iDimerize system, depicted in brown), which can be temporally controlled with a small bidentate ligand (B/B, depicted as a yellow oval) and reversed by a monodentate washout ligand (Belshaw et al., 1996; mCherry-Dmr-ΔNuCat, hereafter referred to as ΔNuCat; Fig. 1 D). This deletion was required to prevent both the native homodimerization of αCat and its recruitment to and function within the cadherin–catenin complex. By replacing this region with the DmrB dimerization domain, we ensured that αCat dimerization function was driven exclusively by the Dmr domain and B/B ligand. Because αCat lacking the N-terminal domain largely rescues cell–cell adhesion (Kadowaki et al., 1994; Imamura et al., 1999) and Drosophila embryogenesis when fused directly to the E-cadherin cytoplasmic domain (Desai et al., 2013), we reasoned that the ΔNuCat construct constituted the best way to begin assessing the contributions of extrajunctional αCat homodimers to F-actin organization and cell behaviors.

An mCherry-DmrB (hereafter referred to as mCherry) construct served as a control to verify that observed phenotypes were not due to the presence of the DmrB domain, mCherry tag, or dimerization ligands. We expressed mCherry or ΔNuCat in R2/7 DLD1 cells. Immunoblot analysis confirmed that ΔNuCat is expressed similarly to mCherry-tagged FLαCat (Fig. S1 A), and coimmunoprecipitation confirmed that ΔNuCat does not associate with the cadherin–catenin complex (Fig. S1 B). Analysis of these constructs by blue native PAGE (BN-PAGE) demonstrated B/B dose–dependent formation of dimers (Fig. 1 E). Although the ΔNuCat construct did not appear to dimerize as efficiently as mCherry, we speculate that some of the ΔNuCat is underrepresented because of dimer-dependent interactions that impede its mobility by native gel analysis. With this system, we found that forced dimerization of ΔNuCat promoted its recruitment to cell protrusions within 5 min (Video 1 and Fig. 1 F). These data suggest that homodimerization of αCat is largely sufficient to control its cortical localization in cells.

Forced dimerization of αCat promotes formation of filopodia and radiating protrusions at nascent contacts

To assess the unique contributions of αCat homodimerization to actin organization and function, we performed live-cell imaging in ΔNuCat cells coinfected with GFP-LifeAct. αCat forced dimers were rapidly recruited to the cell periphery as a function of B/B dimerization ligand, where we also observed the formation of prominent filopodia (Video 1 and Fig. 2 A). Filopodia abundance reached a maximum of 12–15 min after drug treatment. Changes in actin density were also apparent at the ultrastructural level using platinum replica electron microscopy (Fig. 2 B).

Although these studies reveal how αCat forced dimers could acutely affect F-actin organization in single cells, we also sought to determine the consequences of αCat forced dimerization during more sustained drug incubation times (3–15 h) in the context of cells proximal to neighbors. Under these conditions, we observed the formation of “radial protrusions,” structures with a broad lamellipodial base and radiating filopodia at nascent cell-cell contacts (Fig. 2, C and D). Importantly, the length of the filopodia directly correlated with the dose of B/B (Fig. 2, E and F). Because recent studies implicated filopodial protrusions as being important for cell–cell formation (Mattila and Lappalainen, 2008; Hoezle and Switkina, 2012) and compaction (Fierro-González et al., 2013), we wondered whether these protrusions enhanced cell–cell contact. Indeed, the B/B ligand promoted close cell–cell contact in ΔNuCat cells and was fully reversible with the washout ligand (Fig. 2 G).
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[arrows depict close proximity] and Video 2). Altogether, these data suggest that αCat dimerization may direct the formation of radial protrusions that appear important for initiating nascent cell–cell contact.

αCat dimers prefer phosphatidylinositol-3,4,5-trisphosphate (PIP3) phosphoinositides

Many actin-binding proteins bind phosphoinositides (PtdInsP) as a means to control their membrane proximal activation (Yin and Janmey, 2003), and the αCat homologue, vinculin, is an established phosphatidylinositol-4,5-bisphosphate (PIP2)–binding protein (Weekes et al., 1996; Palmer et al., 2009). To explore the possibility that αCat binds to PtdInsP as a way to facilitate its cortical recruitment, we first assessed the binding of purified recombinant full-length (FL) αCat dimers (separated by size exclusion chromatography; Fig. 3C) to vesicles containing seven different PtdInsP (i.e., 1-palmitoyl-2-oleoyl-sn-glycero-3-phosphocholine [POPC]/1-palmitoyl-2-oleoyl-sn-glycero-3-phosphocholine [POPC]/1-palmitoyl-2-oleoyl-sn-glycero-3-phosphoseryl [POPS]/PtdInsP [77:20:3]) by surface plasmon resonance (SPR) analysis. Sensorgrams show that the αCat
**Figure 2.** Forced dimerization of αCat enhances filopodia on radiating protrusions at nascent contacts. (A) ΔNαCat dimerization by B/B promotes filopodia formation. Filopodia were counted every 1 s during a video of force dimerization (n = 6 FOVs from two BRs; data are mean ± SD). Bars, 10 µm. See also Video 1. (B) Actin ultrastructure of dimerized ΔNαCat (±B/B) by platinum replica electron microscopy. Bars, 500 nm. Arrows show αCat enrichment at protrusions. (C) Epifluorescence microscopy of radial protrusions (RPs; white arrows) induced by homodimerization. Bars, 20 µm. (D) Blinded quantification of RPs (n > 150 cells; FOV counts ratioed to total number of cells to account for variations in cell density; Materials and methods; data are mean ± SD). (E) Structured illumination microscopy (SIM) of RPs with filopodia. Bars, 5 µm. (F) Quantification of filopodia length (n > 13 FOVs; three BRs; data are mean ± SD). Significance in D and F by ANOVA. **, P < 0.01; ***, P < 0.001; ****, P < 0.0001. (G) Time-lapse analysis of B/B-treated ΔNαCat cells coinfectected with GFP-LifeAct. Arrows indicate prolonged cell–cell contact upon homodimerization, which is reversed by washout ligand. Bars, 10 µm. See also Video 2.
dimer effectively binds these vesicles and shows selectivity for vesicles containing PIP₃ over those containing other PtdInsP (Fig. 3 A). This PIP₃ selectivity is comparable to that reported for other canonical PIP₃-binding proteins, including the PDK1 pleckstrin homology domain (Lucas and Cho, 2011; Park et al., 2016). Importantly, the PIP₃ selectivity of the αCat dimer is not due to nonspecific electrostatic effects, because it showed low affinity for POPC/POPS (65:35) vesicles that have the same net charge as POPC/POPS/PIP₃ (77:20:3) vesicles (unpublished data). Remarkably, the αCat monomer did not show any detectable binding to any of the PtdInsP-containing vesicles under these same conditions (Fig. 3 B, binding to PIP₃, shown), indicating that the dimer is the active form of αCat for PtdInsP-dependent membrane binding.

To identify domains that could serve to recruit αCat homodimers to PIP₃-activated membranes, we interrogated candidate motifs using the ScanSite database (Obenauer et al., 2003). This algorithm identified a low-stringency PIP₃-binding motif in the αCat middle M2 domain (Fig. 3 D). This motif contained three basic residues, a feature thought to promote electrostatic interactions with the plasma membrane (McLaughlin and Murray, 2005; Mulgrew-Nesbitt et al., 2006). As this motif is surface exposed in the crystal structure of the αCat homodimer (Ranagarajan and Izard, 2013; Fig. 3 D, red-filled residues), we developed an αCat mutant in which these three basic residues (positively charged at physiological pH) within this putative membrane-recruitment domain of M2 were changed to alanine residues (FL αCat K488A/K493A/R496A, hereafter referred to as KKR<3A).

To exclude the possibility that these point mutations destabilized the tertiary structure of αCat, we assessed the physical properties of purified recombinant αCatKKR<3A relative to WT αCat. The proteins showed a similar capacity to form homodimers by size exclusion chromatography coupled with multi-angle light scattering (Fig. 3 C) and BN-PAGE (Fig. 3 F). This mutant also displayed nearly identical secondary structure and melting temperatures as the FL protein (Fig. 3 E), indicating that these charge substitutions do not significantly alter the overall structure of αCat. Last, the F-actin–binding properties of αCatKKR<3A were not significantly different from the WT αCat (Fig. S1 C).

To evaluate whether these point mutations reduced the affinity of recombinant αCat homodimers for PIP₃, we first quantitatively determined the affinity of the αCat dimer to POPC/POPS/PIP₃ (77:20:3) vesicles by equilibrium SPR analysis. The apparent dissociation constant ($K_d$) calculated for this binding (i.e., 30 ± 3 nM) is comparable with that of the PDK1 pleckstrin homology domain determined under the same conditions (Lucas and Cho, 2011; Fig. 3 G). This demonstrates that the αCat dimer shows exceptionally high affinity for PIP₃-containing membranes and suggests the potential importance of PIP₃ binding to the cellular function and regulation of αCat. We then assessed the affinity of the mutant αCatKKR<3A homodimer for comparison and found that this mutant showed reduced binding to POPC/POPS/PIP₃ (77:20:3) vesicles with $K_d =$ 73 ± 15 nM (Fig. 3, G and H). Although the mutation decreased the overall membrane affinity of the αCat dimer by ~2.4-fold, it did not affect the PtdInsP selectivity of the protein, as indicated by the similar binding ratio between the WT and mutant αCat (i.e., ratio of resonance units [RU] for PIP₃ to RU for PIP₂). We reason, therefore, that the KKR basic patch in αCat contributes to membrane binding likely via PtdInsP-independent electrostatic interactions, where the αCatKKR<3A mutant can be used to assess the physiological importance of membrane binding activity of the αCat homodimer.

**αCat localization, homodimerization, and phosphatidylinositol 3-kinase (PI3K) signaling**

Consistent with the observed specificity of αCat dimers for PIP₃, it is well established that the leading edge of migrating cells is characterized by PIP₃ accumulation, rather than PIP₂, signaling lipids (Postma et al., 2004; Cai and Devreotes, 2011). Therefore, we sought to determine whether the localization of extrajunctional αCat to the leading edge generated after wounding was dependent on PI3K signaling. Indeed, a PI3K inhibitor, wortmannin, completely blocked the recruitment of αCat to the wound front (Fig. 4, A and B). Conversely, activation of the EGF receptor (highly up-regulated in A431D cells; Engelmann et al., 2005) was sufficient to increase αCat enrichments at the cell periphery (Fig. 4, A and D; and Video 3). Thus, the cadherin-free pool of αCat can respond to growth factor signals that activate PI3K.

Estimates that the $K_d$ of αCat homodimers is >40 times higher than the concentration of cytosolic αCat in cells (Drees et al., 2005) raise the possibility that PI3K-dependent localization of αCat to PIP₃-enriched membranes (Fig. 4 A) might facilitate formation of αCat homodimers by enhancing the local concentration of αCat through reduction of dimensionality. However, inhibiting PIP₃ formation with wortmannin failed to alter the monomer/dimer ratio of cytosolic αCat by native gel analysis following endogenous αCat from parental DLD1 cells (Fig. S1 D). Curiously, the detection of cytosolic αCat increased under wortmannin treatment (Fig. S1 D), perhaps consistent with the idea that αCat binding to PIP₃ promotes recruitment to activated membranes. Indeed, R2/7 cells expressing FL αCat or FLαCatKKR<3A showed that FLαCatKKR<3A was more abundant in the cytosolic fraction, whereas wortmannin treatment modestly increased the amount of FLαCat detected in the cytosolic fraction (Fig. S1 E). No differences were seen between αCat incorporation within the E-cadherin complex (Fig. S1 F). These data suggest that αCat binding to PIP₃ is not required to promote homodimer formation in cells but rather its localization to membranes, although the slow dissociation of αCat homodimers may be a confounding factor in this assessment (Pokutta et al., 2014).

**Filopodia promoted by force dimerization are reduced in an ΔNoCatKKR<3A mutant**

We next sought to evaluate whether mutation of the KKR basic patch in our forced-homodimer system (ΔNoCatKKR<3A) altered cortical actin phenotypes observed in Fig. 2. We confirmed dimerization of these constructs by BN-PAGE (Fig. 4 B). But remarkably, dimerization of ΔNoCatKKR<3A did not promote filopodia formation (Video 1 and Fig. 4 C; mean number of filopodia, 3.2 ± 0.56 for ΔNoCat vs. 11.83 ± 2.71 for ΔNoCatKKR<3A) or the substantial formation of radiating protrusions (Fig. 4, D and E). The minor PIP₃-independent contribution of αCat forced dimers to radial protrusion formation may be due to the modest overexpression of these F-actin–binding proteins (Fig. S1 A). Additionally, the filopodia observed were significantly shorter (Fig. 4, F and G), and protrusive structures were unable to promote prolonged contact between cells during the drug course study (Fig. 4 H, relative to Fig. 2 G, and Video 2). Together, these data demonstrate that proper
Figure 3. αCat homodimers prefer PIP3 phosphoinositides. (A) Phosphoinositide (PtdInsP) selectivity of WT, full-length (FL) αCat dimer. SPR sensorgrams for seven different POPC/POPS/PtdInsP (77:20:3) vesicles are shown (from top to bottom): PIP3 >> PIP2 > phosphatidylinositol-3,4-bisphosphate (PI34P2) ≈ phosphatidylinositol-5-monophosphate (PI5P) > phosphatidylinositol-4-monophosphate (PI4P) ≈ phosphatidylinositol-3,5-bisphosphate (PI35P2). 200 nM αCat was used for all measurements. Data are representative of three experiments. (B) Binding of αCat homodimer versus monomer (200 nM each) to POPC/POPS/PIP3 (77:20:3) vesicles. Data are representative of three experiments. (C) Chromatographic separation of monomer (M) and dimer (D) fractions of recombinant αCat proteins. (D) αCat schematic with KKR-basic patch overlaid on crystal structure of αCat dimer. Subunit A in gray; subunit B in green/blue; basic patch in red. (E) Circular dichroism (CD) spectra thermal denaturation analysis of purified recombinant protein; FLαCat (red trace) or FLαCatKKR≤3A (blue trace) proteins. No global differences between WT and mutant protein were detected. Wavelength scan: n = 4; temperature scan: n = 2. (F) SDS-PAGE (top) and BN-PAGE (bottom) analysis of purified proteins (10 µg). (G) Membrane binding isotherms for FLαCat dimer (open symbols) and FLαCatKKR≤3A mutant dimer (closed symbols). n = 3; data are mean ± SD. Lines represent theoretical curves constructed from apparent dissociation constant (Kd) and the maximal R_eq value (R_max) determined from nonlinear least squares analysis of the binding isotherm using the equation \( R_{eq} = R_{max}/(1 + K_d/P) \). Kd = 30 ± 3 nM for WT and 73 ± 15 nM for the mutant. (H) PIP3 versus PIP2 selectivity of WT FLαCat dimer (cyan) and FLαCatKKR≤3A mutant dimer (orange). Data are representative of three experiments. Notice that although the mutation decreased the overall membrane affinity of the αCat dimer, it did not affect the PtdInsP selectivity of the protein, as indicated by essentially the same (RU for PIP3)/(RU for PIP2) ratio for WT and the mutant. Each SPR measurement was performed in 50 mM Tris-HCl, pH 8.0, containing 0.1 M NaCl and 1 mM TCEP using L1 chip coated with POPC/POPS/PtdInsP (77:20:3) vesicles as the active surface. 200 nM αCat was used for both WT and mutant proteins. POPC vesicles were used to coat the control surface for most experiments. For A, POPC/POPS (80:20) vesicles were used for the control surface to eliminate the potential contribution of POPS to PtdInsP selectivity.
Figure 4. Filopodia promoted by force-dimerization are reduced in ΔαCatKKR<3A mutant. (A) αCat localization is sensitive to wortmannin and EGF. Schematic of PIP3 synthesis and impact of drugs to right. n = 3, >14 FOVs. Data indicate mean ± SD. Significance by ANOVA. Arrows show αCat enrichment at protrusions. Bars, 20 µm. See also Video 3. (B) BN-PAGE analysis of dimer formation (D) relative to monomer (M); B/B treatment 3 h. (C) ΔαCatKKR<3A dimerization by B/B reduces filopodia formation compared with ΔαCat. As in Fig. 2, filopodia were counted every 1 s during a video of force dimerization. Bars, 10 µm; n = 6 FOVs from two BRs; data are mean ± SD. (D) Epifluorescence microscopy of radial protrusions (RPs; white arrows) reduced in ΔαCatKKR<3A mutants. Bars, 20 µm. (E) Blinded quantification of RP (n > 150 cells; FOV counts ratioed to total number of cells to account for variations in cell density; Materials and methods). (F) Length of filopodia decreased in ΔαCatKKR<3A, as imaged in structured illumination microscopy (SIM) of RP filopodia. Bars, 5 µm. (G) Quantification of filopodia length (n > 13 FOVs from three BRs), table of results below. Significance in E and G by ANOVA; data are mean ± SD. ** P < 0.01; *** P < 0.001; **** P < 0.0001. (H) Time-lapse analysis of B/B-treated ΔαCatKKR<3A cells coinfected with GFP-LifeAct. Prolonged cell–cell contact upon homodimerization was not observed in mutant construct. Bars, 10 µm. See corresponding Videos 1 and 2.
recruitment of the homodimer to PIP$_3$-activated membranes may be critical for its activity, such that even modest reduction of membrane recruitment via KKR<3A mutations can attenuate αCat homodimer function.

**Exogenous PtdInsP are sufficient to recruit endogenous αCat**

Because liposome-based in vitro assays often fail to recapitulate presentation of PtdInsP as in vivo (Narayan and Lemmon, 2006), we sought to test whether PIP$_3$ could recruit endogenous αCat to bona fide cellular membranes by introducing ectopic BODIPY-tagged PIP$_3$ into the apical membrane using histone (H1) carriers (PIP$_3$/H1; Fig. 5 A; Gassama-Diagne et al., 2006). We validated the preferential recruitment of αCat to PIP$_3$, by comparing colocalization (Pearson’s coefficient) of BODIPY-tagged PIP$_3$, phosphatidylinositol-3,5-bisphosphate or PIP$_2$ in each slice of the Z-stack. In polarized MDCK cells grown on filters for 2 wks, we observed statistically significant colocalization of αCat with PIP$_3$, relative to other PtdInsP or the histone carrier alone (Fig. 5, B and C). These data validate the preferential in vitro binding of αCat to PIP$_3$ (Fig. 3) and demonstrate that PIP$_3$ and, to a lesser extent, PIP$_2$ are sufficient to recruit αCat to membranes.

To evaluate the contribution of the basic patch in αCatKKR<3A to PIP$_3$ recruitment in cells, we repeated the previous experiment in the αCat-null R2/7 variant of DLD1 cells expressing an mCherry-tagged WT FLαCat or FLαCatKKR<3A mutant. We observed gradual apical recruitment of αCat in R2/7 cells during a 1-h PIP$_3$/H1 integration (Fig. 6 A) and apical enrichments colocalized with PIP$_3$, after washout of excess PIP$_3$/H1 (arrow). Enrichments were not due to an increase in the overall expression of αCat (Fig. 6 B). These data are consistent with previous PtdInsP-interactome studies showing enrichment of αCat with PIP$_3$ (Catimel et al., 2009; Jungmichel et al., 2014). Because the expression of αCat in these cell lines is not as definitively junctional (relative to MDCK cells in Fig. 5), we analyzed colocalization with PIP$_3$ by rendering 3D surfaces of PIP$_3$ integration and αCat from the Z-stacks to compare 3D colocalization in Imaris. When we compared colocalization of FLαCat with PIP$_3$ compared with FLαCatKKR<3A, we found a statistically significant decrease in colocalization upon cellular loss at these residues (Fig. 6, C and D), suggesting that the basic patch promotes recruitment to PIP$_3$ membranes. Importantly, this difference is not due to differences in PIP$_3$/H1 integration (total volume) or to the size of individually integrated PIP$_3$/H1 volumes (Fig. 6 C) or to the histone carrier alone. Importantly, E-cadherin was not recruited to PIP$_3$/H1 (Fig. 6 E), suggesting that recruitment by PIP$_3$ may be a unique feature of extrajunctional αCat. Altogether, these data show that a basic patch in the M2 region of αCat contributes to its recruitment to PIP$_3$-loaded membranes.

**αCat basic patch contributes to adhesion and migration**

Although the aforementioned studies clearly establish that forced dimerization of αCat promotes its membrane recruitment and alters F-actin organization, the ∆NaCat construct is not a native αCat protein. We sought, therefore, to assess the importance of this basic patch in the context of a FL αCat that could participate in both cadherin–catenin complex and homodimeric functions by comparing the adhesive activities of FLαCat and FLαCatKKR<3A expressing R2/7 cells (immuno-blot in Fig. S1 A). As with the recruitment of extrajunctional αCat to the leading edge of wounded cells (Fig. 1), FLαCat was enriched at the wound front compared with FLαCatKKR<3A (Fig. 7, A and B). In addition, epithelial sheets expressing FLαCatKKR<3A closed twice as much wound area as those expressing FLαCat (Fig. 7, C and D; and Video 4), suggesting that αCat recruitment to PIP$_3$-activated membranes contributes to the coordination of this process. Indeed, in FLαCat cells, actin organization along the wound was more uniform, with actin cables parallel to the wound coordinated across cells (Fig. 7 E, top), reminiscent of the F-actin cables formed during epidermal closure in invertebrates (Jacinto et al., 2002). In contrast, FLαCatKKR<3A cells were often separated from their immediate neighbors along the wound front and did not form these actin cables (Fig. 7 E, bottom). To visualize F-actin during live-cell migration, we cotransfected the cells in Fig. 7 A with GFP-LifeAct. We found that monolayers expressing FLαCat displayed prominent protrusions with long filopodia (Fig. 7 F and Video 4) and colocalized with F-actin as before (Fig. 7 F), particularly in cells directly following the leading cells, compared with FLαCatKKR<3A cells. Moreover, when the migrating cells were imaged by phase contrast, the FLαCatKKR<3A cells were more rounded with refractile cell–cell contacts (Fig. 7 F), suggesting a reduction in cell–cell adhesion. To test this directly, we used a standard epithelial sheet fragmentation assay (Escobar et al., 2015). We found that mechanical disruption of monolayers expressing FLαCatKKR<3A mutant formed significantly more fragments than FLαCat cells (Fig. 7, G and H). Because this charge mutant form of αCat shows similar incorporation into the cadherin/β-catenin complex (Fig. S1 F) and binding to F-actin by cosedimentation as WT αCat (Fig. S1 C), we reason that the αCatKKR<3A mutant alters cell–cell adhesive activities by targeting the membrane recruitment of homodimeric αCat. Although it is difficult to rule out the possible contribution of αCat recruitment by PIP$_3$, within the cadherin–catenin complex, our evidence that E-cadherin is not readily recruited to apically added PIP$_3$ (Fig. 6 E), supports the idea that αCat recruitment by PIP$_3$ outside of the cadherin-catenin complex strengthens intercellular adhesion.

**Discussion**

We show that an extrajunctional form of αCat can be localized to PIP$_3$-enriched membranes, where this recruitment and αCat homodimerization appear important for cell contact formation, strength, and coordinated migration. Because recruitment by PIP$_3$ appears to be a unique feature of αCat homodimers, and currently, there are no known mutants that specifically disable the homodimerization function of αCat from the β-catenin/E-cadherin binding function, we speculate that targeting the membrane-binding function of αCat may specifically attenuate the role of extrajunctional αCat homodimer function in cells.

αCat homodimers are implicated in a number of in vitro F-actin–binding activities that may contribute to the phenotypic behaviors measured in this study. For example, αCat dimers not only bind F-actin but can protect it from severing by cofillin (Hansen et al., 2013), can promote its bundling (Koslov et al., 1997), and can induce a conformational change in the actin filament that limits barbed-end growth and branching by the Arp2/3 complex (Drees et al., 2005; Hansen et al., 2013). Consistent with the ability of αCat to limit Arp2/3 in vitro, a previous study...
suggested that the extrajunctional pool of αCat could limit lamellipodial dynamics and cell migration, using a clever construct approach that effectively diverted the extrajunctional pool of αCat toward or away from the plasma membrane (Benjamin et al., 2010). Although forced αCat dimerization promoted the formation of radial protrusions with prominent elongated filo-
Figure 6. αCat<sup>KKR</sup> mutant shows reduced recruitment to exogenous PIP<sub>3</sub>. (A) PIP<sub>3</sub>, apical membrane integration and recruitment assay; mCherry αCat (red), BODIPY–PIP<sub>3</sub> (green) with quantification of total colocalized surface. Arrows show colocalized surfaces as yellow (top) and purple (bottom) when reconstructed as a volume rendering. (B) BODIPY–PIP<sub>3</sub> did not alter expression of αCat (quantification not shown; n = 3). (C) Quantification of colocalization volumes between αCat constructs and PIP<sub>3</sub> (n = 45 from three BRs). Significance by unpaired t test: ***, P < 0.0005; ****, P < 0.0001. Error bars reflect the SD of the mean. (D) Surface renderings of colocalization volumes (purple) relative to BODIPY–PIP<sub>3</sub> integration (green). Slice shows en face view of Z-stack; arrows show colocalization (0.2-µm steps). (E) E-cadherin was not recruited to ectopic BODIPY–PIP<sub>3</sub> integrations, suggesting that this is a unique feature of extrajunctional αCat. Samples were fixed after live-cell imaging, so integration of BODIPY–PIP<sub>3</sub> was prolonged relative to Z-stacks in A and D. Bars, 20 µm.
Figure 7. αCatKR<3A mutant alters cell-cell adhesion and sheet migration. (A) Live imaging of αCat-expressing R2/7 cells imaged for 12 h postwounding. Arrows indicate αCat recruitment in FLαCat but not FLαCatKR<3A. (B) Quantification of enrichments (n > 150 FOVs inclusive of all time points, two BRs). Data are mean ± SD. (C) Representative image of wound closure area. Pink overlay depicts area quantified. (D) Quantification of wound closure (n = 24 FOVs, two BRs, data are mean ± SD). See corresponding Video 4. (E) Phalloidin stain of F-actin at wound front. Red arrows indicate actin cables parallel to the front; black arrows indicate actin cables adjoining cells via junctions. High-resolution analysis of F-actin across adjacent cells by SIM (right). (F) Live imaging of GFP-LifeAct and mCherry FLαCat during wound migration. (Left) Fluorescent image; white arrows indicate protrusions with long filopodia; yellow arrowheads show areas of colocalization. (Right) Bright-field image. Bars, 50 µm. (G) Representative images from a mechanical disruption assay with quantification in H (Materials and methods; n = 5 BRs). Data are mean ± SD; significance in all panels by unpaired t test: ****, P < 0.0001.
podia, and cells expressing FL αCat closed the wound slower than αCatKKR<3A-expressing cells, activities that suggest Arp2/3 suppression, we did not find that αCat recruitment to membranes correlated with a reduction in Arp2/3 accumulation at the periphery (Fig. S1 G). Therefore, future studies will be required to understand how these distinct biochemical activities are temporally and spatially coordinated at the membrane surface by PtdInsP, PIP3 in particular, and account for how αCat homodimers contribute to intercellular adhesive behaviors.

Further studies will also be required to understand how αCat homodimers, but not monomers, show selective binding to PIP3 and, to a lesser extent, PIP2. Although we show that disruption of an algorithm-identified charge cluster in αCat (KKR; MIT Scansite) impaired αCat homodimer binding to PIP3−liposomes by SPR analysis, as well as its recruitment to BODIPY-labeled PIP3 exogenously added to cells, the αCatKKR-basic region cannot explain the PtdInsP binding selectivity of αCat homodimers. Thus, these data suggest that the αCatKKR-basic region likely contributes to electrostatic membrane interactions (e.g., via phosphatidylserine), while other residues in αCat contribute to PIP3 head-group specificity. Efforts to identify candidate PtdInsP binding pockets by molecular modeling are limited by the low resolution (3.66 Å) of the nearly FL αCat dimer crystal structure (Protein Data Bank 4GGG; Rangarajan and Izard, 2013). Moreover, such efforts may be irrelevant if αCat homodimers bind to PIP3 through a pocket formed at an alternative dimer interface not observed in the crystal structure, with two monomers contributing residues to form one PIP3 binding site, analogous to how its homologue, vinculin, binds PIP2 (Chinthalapudi et al., 2014). Indeed, the remarkable affinity difference measured for αCat homodimer and monomer binding to PIP3 cannot be explained by avidity alone, suggesting the possibility that a conformational change in the αCat homodimer may expose a PtdInsP-binding surface that is inaccessible in the monomer. Last, it is worth noting that although the αCatKKR<3A mutant does not apparently perturb its incorporation into the cadherin complex or binding to F-actin, we cannot presently rule out a role for these residues in tension-dependent M-domain interactions that may also contribute to the cell-cell adhesive phenotypes described.

Evidence that the Kd of αCat homodimers is much higher than the concentration of cytosolic αCat in cells (Drees et al., 2005) raises the intriguing possibility that the localization of αCat to PIP3−enriched membranes might serve to facilitate αCat homodimerization by enhancing its local effective concentration. Although inhibiting PIP3 formation with wortmannin failed to alter the monomer/dimer ratio of cytosolic αCat by native gel analysis, it is likely that basic patches in αCat mediate electrostatic interactions with the plasma membrane, locally increasing the concentration of αCat to drive homodimerization. Once at the membrane, homodimer diffusion may be restricted by PIP3, as a way to localize F-actin binding activities of αCat.

In summary, this study demonstrates that αCat homodimers bind to PIP3, respond to PI3K-activated signals, and control actin organization and behaviors that promote cell–cell adhesion, which has implications for the specific targeting of extrajunctional αCat homodimer function in cells.

Materials and methods

Plasmid generation and constructs

All αCat sequences used are based on the epithelial form of αCat. The GFP-tagged IRES-Puro-rat αCat lentiviral vector (pLVX) was provided by J. de Rooij (Center for Molecular Medicine at the University Medical Center Utrecht, Utrecht, the Netherlands). The mutated basic region of the M2 domain (KKR<3A) was creating using QuikChange II site-specific mutagenesis (Stratagene) and QuikChange Primer Design Program in the context of the human αCat protein (pGEX; Escobar et al., 2015). FL and KKR<3A human αCat proteins were cloned into the pLVX vector downstream and in-frame with an N-terminal monomeric mCherry-tag, to avoid dimerization effects mediated by the GFP tag (Shaner et al., 2005). For dimerezize constructs, the first 267 aa of the human αCat sequence (encoding overlapping β-catenin hetero- and αCat homodimerization domains) were removed with the Clontech In-Fusion HD Cloning kit. This transformed truncation, ΔNotCat, was cloned into Lentivirus Inducible Homodimer System (Clontech) that had been N-terminally tagged with mCherry. The puromycin gene was removed from the pLVX backbone to accommodate the size of the αCat-expression constructs. Primers used for truncated ΔNotCat and ΔNotCatKKR<3A constructs are available upon request. A lentiviral LifeAct-GFP was purchased from Addgene (pLenti.PGK.LifeAct-GFP; Plasmid 51010).

Cell culture and stable cell line selection

MDCK, A431D, and DLD1 cells (American Type Culture Collection) and DLD1 R2/7 αCat-deficient colon carcinoma cells (a gift from F. van Roy, Ghent University, Ghent, Belgium) were maintained in DMEM (Corning), containing 10% FBS (Atlanta Biologicals or JRS Scientific), 100 U/ml penicillin, and 100 µg/ml streptomycin (Corning). iDimerize cell lines were sorted for positive selection of mCherry fluorescence by flow cytometry using FACSAria 5 (BD Biosciences). mCherry FLαCat or FLαCatKKR<3A cell lines were selected in 5 µg/ml puromycin (Sigma-Aldrich) and were subsequently sorted for mCherry expression using flow cytometry. In drug treatment assays, cells were treated with one of the following: DMSO control (Sigma-Aldrich) or 1–20 µM wortmannin (W1628; Sigma-Aldrich) for 3 h. In forced-dimerization assays, cells were treated with 25, 250, or 500 nM B/B homodimer (635059; Sigma-Aldrich) and puromycin (5 µg/ml) for 20 min to 15 h, which was sometimes followed by treatment with 1 µM washout ligand (635088; Clontech).

Antibodies

The following primary antibodies were used: polyclonal rabbit anti–αCat (C3236; Cell Signaling), hybridoma mouse anti-αCat (5B11; Daugherty et al., 2014), monoclonal mouse anti–β-catenin (610154; BD Biosciences), polyclonal rabbit anti–β-catenin (06-734; EMD Millipore), monoclonal mouse anti–E-cadherin (610182; BD Biosciences), monoclonal mouse anti–E-cadherin (HECD1, 13-1700; TaKaRa), polyclonal rabbit anti–GAPDH (FL-335, sc-25778; Santa Cruz), anti–p34-Arc/ARPC2 (07-227; EMD Millipore), monoclonal mouse anti–GAPDH (9484; Abcam), hybridoma mouse anti-tubulin (DM1A, T9026; Sigma-Aldrich), polyclonal rabbit anti-mCherry (5993; Bios, Vision), and Alexa Fluor 488 or 568 phalloidin (A12379; Invitrogen). Secondary antibodies for Western blotting included HRP-conjugated goat anti–mouse and anti–rabbit antibodies (Bio-Rad) or fluorescently labeled donkey anti–mouse and anti–rabbit antibodies (680RD or 800RD; LiCor Biosciences). Secondary antibodies for immunofluorescence included IgG Alexa Fluor 488– or 568–conjugated goat anti–mouse or anti–rabbit antibodies (Invitrogen).
Immunofluorescence and Imaging

Cells were grown on coverslips, fixed in 4% PFA (Electron Microscopy Services) for 15 s, quenched with glycine, permeabilized with 0.3% Triton X-100 (Sigma-Aldrich), and blocked with normal goat serum (Sigma-Aldrich). Primary and secondary antibody incubations were performed at RT for 1 h, interspaced by multiple washes in PBS, and followed by mounting coverslips in ProLong Gold fixative (Life Technologies). Images of asymmetrical αCat recruitment to the wound front (Fig. 1, A–C; and Fig. 4 A) were captured at RT with an Axioplan2 microscope (Zeiss) equipped with a 100× Plan-Neofluar with oil, NA 0.75 objective, and AxioCam HR Camera using AxioVision 4.8 software. Wide-field fluorescence live-cell microscopy was used to generate videos of the iDimerize drug time course and single-cell filopodia projections (Fig. 1 F; Fig. 2, A and G; Fig. 4, C and H; and Videos 1, 2, and 3) with a Nikon Ti Eclipse inverted microscope equipped with a Yokogawa CSU-X1 spinning disk head, Perfect Focus system (Nikon), and an Andor xIOn electron-multiplying charge-coupled device (CCD) camera (Andor Technology) controlled by MetaMorph v.7.7.7.0 software (Molecular Devices). Cells were maintained at 37°C plus 5% CO₂ during imaging using a Tokai-Hit stage-top incubator (Tokai-Hit) or an Okolab gas mixer (Okolab). Time-lapse sequences were acquired at 15-s intervals using the 561-nm and/or 488-nm laser. For all experiments, cells were plated on glass-bottomed dish coverslips 24 h before imaging. Imaging of radiating protrusions and filopodia (Figs. 2 E and 4 F) was performed using a structured illumination superresolution microscope (N-SIM; Nikon). Samples were illuminated with spatially high-frequency patterned excitation light (100× objective lens, NA 1.49; TiE N-SIM microscope [Nikon] and iXon X3 897 camera [Andor Technology]). The moiré patterns were produced and analyzed with analytically processed (Nikon Elements v.4.20.01, 3D SIM reconstruction tool) to reconstruct the subresolution structure of the samples. Platinum replica electron microscopy (Fig. 2 B) was performed as previously described (Korobova and Svitkina, 2008; Svitkina, 2009). Cells were grown subconfluently on coverslips and preextracted with 0.5% Triton X-100, 0.25% glutaraldehyde in PEM buffer (100 mM Pipes, pH 6.9, 1 mM EGTA, and 1 mM MgCl₂). Cells were then fixed in 2% glutaraldehyde in 0.1 M Na-cacodylate, pH 7.3, for 20 min at RT. One set of cells was then stained with 488 phalloidin for visualization of all cell peripheries in the red channel. A structure was counted as a radial protrusion if a lamellipodial fan, distinct from the main cell body, was observed. Adjacent structures were counted as one unless they were clearly supported by distinct bases, and structures were not counted if saturation inhibited a clear visualization of structure. This main lamellipodial fan was often concurrent with enrichment of αCat localization and radiating filopodia from the central structure, though the latter two features were not a requirement for a structure to be counted as a radial protrusion. To account for variability in radial protrusion readouts, >150 cells were counted, and each FOV was ratioed against the total number of cells to account for variations in cell density. Ratios were averaged for each condition and compared via one-way ANOVA with Tukey’s multiple-comparisons test. To measure filopodia length from structured illumination microscopic images, reconstructed ND2 files were imported to ImageJ, and filopodia length (actin channel) was measured with a freehand line. Statistics were calculated using one-way ANOVA with Tukey’s multiple-comparisons test (GraphPad Prism). Images presented in the paper were adjusted for brightness and contrast but were otherwise unprocessed.

Image analysis and quantification

To quantify the number of enrichments at the leading edge (Fig. 4 A), cells were fixed and imaged with an Axioplan2 microscope (in the previous section). To qualify for quantification, enrichment needed to be brighter than the cell body, and distinct from adjacent enrichments. Enrichments were only counted within the first two cell rows of the leading edge, not in leading cells farther back in the epithelial sheet. The file names were randomly generated (RandomNames Script from HowToGeek.com) to eliminate sample bias before quantification of four or five fields of view (FOVs) for three biological replicates (BRs; n = 14–17) for each condition (>124 cells). Significance was determined by multiple-comparisons ANOVA. Images presented in the paper were adjusted for brightness and contrast but were otherwise unprocessed.

To quantify the number of filopodia observed in the drug-time course videos of ΔNaCat and ΔNaCatAKR-GA cells tagged with GFP-LifeAct (Fig. 3 D), still images (every 1 min, six BRs) from the image sequence were imported into ImageJ. Filopodia were counted if there was a distinct actin projection extending from the cell body. Error bars here and throughout this study represent SDs. Representative images are shown with white arrows pointing to filopodia projections; images were adjusted for brightness and contrast but were otherwise unprocessed.

To quantify the number of radial protrusions observed (Figs. 2 C and 4 D), cells were fixed and stained with Hoechst and imaged with an Axioplan2 microscope (in the previous section). Three to eight FOVs for each of the three BRs were imaged, exported as channel-separated JPEG images, and quantified using the multipoint tool to count protrusions in FIJI/ImageJ software. The file names were randomly generated (RandomNames Script from http://www.howtogeek.com) to eliminate sample bias before quantification by a third party in a double-blind experiment. Total cell numbers were first counted using the Cell Counter plugin after thresholding the Hoechst-stained nuclei. Brightness and contrast were adjusted to visualize all cell peripheries in the red channel. A structure was counted as a radial protrusion if a lamellipodial fan, distinct from the main cell body, was observed. Adjacent structures were counted as one unless they were clearly supported by distinct bases, and structures were not counted if saturation inhibited a clear visualization of structure. This main lamellipodial fan was often concurrent with enrichment of αCat localization and radiating filopodia from the central structure, though the latter two features were not a requirement for a structure to be counted as a radial protrusion. To account for variability in radial protrusion readouts, >150 cells were counted, and each FOV was ratioed against the total number of cells to account for variations in cell density. Ratios were averaged for each condition and compared via one-way ANOVA with Tukey’s multiple-comparisons test. To measure filopodia length from structured illumination microscopic images, reconstructed ND2 files were imported to ImageJ, and filopodia length (actin channel) was measured with a freehand line. Statistics were calculated using one-way ANOVA with Tukey’s multiple-comparisons test (GraphPad Prism). Images presented in the paper were adjusted for brightness and contrast but were otherwise unprocessed.

PIP₃-histone apical recruitment assay

13 µl 100 µM PIP₃, phosphatidylinositol-3,5-bisphosphate, or PIP₂ tagged with BODIPY-FL was incubated with 13 µl 100 µM Histone H1 Carrier (Echelon) for 10 min after vigorous pipetting to facilitate complex formation. Complexes were dripped into confluent cells covered in 100 µl FluoroBrite DMEM (Life Technologies), and PtdInsP–histoneH1 complexes were allowed to integrate for 1 h. After 1 h of integration, the medium was removed and replaced with fresh FluoroBrite medium. For both experiments, addition of histone H1 alone served as a negative control.

For MDCK cells grown on filters (Fig. 5) and fixed after PdInsP integration, slides were imaged at RT with 60x oil Apo TIRF NA 1.49 objective on the Nikon A1R laser scanning confocal inverted microscope equipped with two standard photomultiplier tubes (408 and 640) and two high-sensitivity gallium arsenide photodiode detectors (488 and 561). The system is equipped with the Perfect Focus focal drift compensation mechanism and automated XY stage. Live-cell imaging of R2/7 cells Z-stack images (0.2-µm steps; Fig. 6) were captured (before and after PdInsP/H1 integration) using a Nikon Ti Eclipse inverted microscope equipped with a Yokogawa CSU-X1 spinning disk head, Perfect Focus system, a 100× 1.49 NA APO TIRF objective, and an Andor xIOn electron-multiplying CCD camera. Cells were maintained...
at 37°C plus 5% CO₂ during imaging using a Tokai-Hit stage-top incubator or an Okolab gas mixer.

For MDCK cells (Fig. 5), colocalization was quantified in Nikon Elements v.4.60.00, build 1170. Pearson’s correlation coefficients for each slice of the Z-stack were calculated using the colocalization tool. To show how colocalization changed across the stack, these data were plotted in Prism as a function of Z-stack height. Coefficients for each stack were combined (n = 3) to compare colocalization of each PtdInsP/H₁. Statistical analysis was performed using one-way ANOVA with multiple comparisons. Images presented in the paper were adjusted for brightness and contrast but were otherwise unprocessed.

For R2/7 cells (Fig. 6), 3D colocalization of PIP and αCat was quantified in Imaris (Bitplane). This method was chosen because the localization of αCat in R2/7 cells has increased cytoplasmic localization relative to the junctional localization in MDCK cells and thereby cannot quantitatively measure colocalization via Pearson’s coefficient, as in Fig. 5. Z-stacks were imported to Imaris, and new surfaces rendered for quantitatively measure colocalization via Pearson’s coefficient, as in

\[ P_{eq} = \frac{R_{max}}{1 + \frac{R_{max}}{[P]_0}} \]

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\[ P_{eq} = \frac{R_{max}}{1 + \frac{R_{max}}{[P]_0}} \]

For kinetic measurements, the flow rate was maintained at 10 µl/min. For equilibrium binding measurements of αCat were performed at a flow rate of 10 µl/min, which allowed enough time for the R values of the association phase to reach near equilibrium levels (Rₐ; Ananthanarayanan et al., 2003). Each sensorgram was background-corrected by subtracting the control surface response from the active surface response. Protein solutions with different concentrations were injected to collect a set of Rₐ values that were plotted against the protein concentrations ([P]₀).

An apparent dissociation constant (Kₐ) was then determined by nonlinear least squares analysis of the binding isotherm using the equation

\[ R_{eq} = R_{max} \frac{[P]_0}{1 + K_a [P]_0} \]

Because the concentration of lipids coated on the sensor chip cannot be accurately determined, Kₐ is defined as [P]₀ yielding half-maximal binding with a fixed lipid concentration. Each measurement was repeated at least three times to determine mean and SD values. For kinetic measurements, the flow rate was maintained at 10 µl/min.

**Actin pelleting assay**

Monomeric rabbit skeletal muscle actin was polymerized in the polymerization buffer (5 mM Tris-HCl, pH 8.0, 50 mM KCl, 2 mM MgCl₂, 1 mM ATP, 0.2 mM CaCl₂, and 0.5 mM DTT) for 1 h at RT. 5 µM actin and 5 µM αCat were mixed in the binding buffer (5 mM Tris-HCl, pH 8.0, 50 mM KCl, 40 mM NaCl, 2 mM MgCl₂, 1 mM ATP, 0.2 mM CaCl₂, 0.5 mM DTT, and 0.5 mM TCEP) for 1 h, and supernatant and pellet fractions were separated by centrifugation at 100,000 g for 20 min. Fractionated protein samples were analyzed by SDS-PAGE with Coomassie blue stain, and gel band intensity was measured by ImageJ.
using the rectangular selection tool followed by the gel analyzer tool (n = 3; data are mean ± SD).

**Cell fractionation analyses**

Hypotonic, detergent-free lysis for assessing the cytosolic fraction of αCat (Fig. S1 E): cells were washed twice with PBS**, scraped, and pelleted at 1,400 rpm for 3 min. The cell pellet was resuspended in hypotonic lysis buffer (30 mM Tris-HCl, pH 7.5, and 1 EGTA-free proteinase inhibitor/10 ml; Roche). Cells were swollen on ice for 30 min and lysed by 20 strokes through a tuberculin needle. Tonicity was restored with 0.25 M sucrose homogenized with 10 strokes of a tuberculin needle; a portion was held aside as “input.” The remainder was ultracentrifuged at 100,000 g for 90 min to generate a supernatant (S100G) containing cytosolic proteins for analysis by SDS-PAGE. To achieve a more concentrated cytosolic fraction for BN-PAGE analysis, we used a standard freeze-thaw protocol (Tansey, 2006). In brief, cells were washed twice with PBS**, scraped, and pelleted at 1,000 rpm for 3 min. The pellet was resuspended in Freeze/Thaw Lysis Buffer (600 mM KCl, 20 mM Tris-HCl, pH 7.5, 20% glycerol, 100 mM PMSF, 0.35 mg/ml Pepstatin A, 1 mg/ml aprotinin, and 2.3 mg/ml leupeptin) containing 200 µl per 10-cm dish). Cells were lysed in cycles of liquid nitrogen wash in ddH₂O. The membrane was blocked in TBST-5% milk for 1 h before proceeding with normal Western blot methods using the indicated antibodies (Cell Signaling). For immunoprecipitation, cells were harvested 1 h before processing for immunoprecipitation; IgG antibody (Roche) and precipitated with the indicated antibodies.

**Supplemental material**

The supplemental material for this manuscript provides important control experiments validating construct expression levels and typical binding partners (Fig. S1 A, B, and F), as well as experiments showing that the αCatKR3A mutant does not significantly alter F-actin binding (Fig. S1 C). We also present evidence that the αCat homodimer is not sensitive to inhibition of PI3K by wortmannin (Fig. S1 D) but that this treatment is associated with an increase in the cytosolic pool of αCat (Fig. S1, D and E). Given evidence that αCat homodimers can inhibit Arp2/3-dependent actin polymerization (Drees et al., 2005) and accumulation at the leading edge of cells (Benjamin et al., 2010), we also show evidence that Arp3 protein levels are not obviously altered by αCat forced dimerization (Fig. S1 G). Last, the videos included provide temporal context to the still images in the text (Video 1 with Figs. 1 F, 2 A, and 4 C; Video 2 with Figs. 2 G and 4 H; Video 3 with Fig. 4 A; and Video 4 with Fig. 7, A–E).

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**Author contributions:** Conceptualization: M.N. Wood and C.J. Gottardi; validation: A.S. Flozak; formal analysis: M.N. Wood, I. Singaram, W. Cho, C. Chung, and N. Ishiyama; investigation: M.N. Wood, N. Ishiyama, A.S. Flozak, and C. Chung; resources: A. Yemelyanov, M. Ikura, and W. Cho; writing original draft: M.N. Wood, W. Cho, and C.J. Gottardi; editing and review, M.N. Wood, C.J. Gottardi, and A.S. Flozak; visualization: M.N. Wood; supervision: C.J. Gottardi; funding acquisition: M.N. Wood, M. Ikura, W. Cho, and C.J. Gottardi.

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**Online supplemental material**

The supplemental material for this manuscript provides important control experiments validating construct expression levels and typical binding partners (Fig. S1 A, B, and F), as well as experiments showing that the αCatKR3A mutant does not significantly alter F-actin binding (Fig. S1 C). We also present evidence that the αCat homodimer is not sensitive to inhibition of PI3K by wortmannin (Fig. S1 D) but that this treatment is associated with an increase in the cytosolic pool of αCat (Fig. S1, D and E). Given evidence that αCat homodimers can inhibit Arp2/3-dependent actin polymerization (Drees et al., 2005) and accumulation at the leading edge of cells (Benjamin et al., 2010), we also show evidence that Arp3 protein levels are not obviously altered by αCat forced dimerization (Fig. S1 G). Last, the videos included provide temporal context to the still images in the text (Video 1 with Figs. 1 F, 2 A, and 4 C; Video 2 with Figs. 2 G and 4 H; Video 3 with Fig. 4 A; and Video 4 with Fig. 7, A–E).
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