Introduction

Tight junctions, adherens junctions (AJs), and desmosomes promote adhesion between epithelial cells, initiate the assembly of the mechanical cytoskeleton linkage, and facilitate the formation of a polarized epithelial monolayer (Gumbiner, 1996). AJs initiate these processes and are essential for morphogenesis, wound healing, and the retention of cell polarity and tissue integrity (Perez-Moreno et al., 2003). In epithelia, AJ formation is mediated by the calcium-dependent homophilic binding of E-cadherin molecules on neighboring cells (Gumbiner, 1996). These interactions link adjacent cells and promote the nucleation of a cytoplasmic protein complex consisting of p120-, β-, and α-catenins, which bridges E-cadherin clusters and the actin cytoskeleton. The biological necessity of AJ proteins has been underscored by a high correlation between the malfunctioning of AJ proteins, E-cadherin in particular, and tumor metastasis (Kang and Massague, 2004).

Assembly of E-cadherin–based adherens junctions (AJ) is obligatory for establishment of polarized epithelia and plays a key role in repressing the invasiveness of many carcinomas. Here we show that type Iγ phosphatidylinositol phosphate kinase (PIPKIγ) directly binds to E-cadherin and modulates E-cadherin trafficking. PIPKIγ also interacts with the μ subunits of clathrin adaptor protein (AP) complexes and acts as a signalling scaffold that links AP complexes to E-cadherin. Depletion of PIPKIγ or disruption of PIPKIγ binding to either E-cadherin or AP complexes results in defects in E-cadherin transport and blocks AJ assembly. An E-cadherin germline mutation that loses PIPKIγ binding and shows disrupted basolateral membrane targeting no longer forms AJs and leads to hereditary gastric cancers. These combined results reveal a novel mechanism where PIPKIγ serves as both a scaffold, which links E-cadherin to AP complexes and the trafficking machinery, and a regulator of trafficking events via the spatial generation of phosphatidylinositol-4,5-bisphosphate.

During tumor progression, the E-cadherin gene can be functionally silenced or inactivated by distinct mechanisms (Nelson and Nusse, 2004). In addition to transcriptional repression by SIP-1, 8EF-1, Snail/Slug, E12/47, and Twist (Huber et al., 2005), posttranslational regulation of E-cadherin stability modulates its activity. Precisely tuned exocytic and endocytic pathways control the amount of E-cadherin residing on the plasma membrane (PM) and are important for modulation of E-cadherin function and AJ assembly (Bryant and Stow, 2004). Recent evidence suggests that Rab11 (Lock and Stow, 2005), p120-catenin, ARF6, tyrosine phosphorylation, and ubiquitination (D’Souza-Schorey, 2005) all control the trafficking and assembly of E-cadherin in mammalian cells. Additionally, transport of E-cadherin is regulated by the composition of the cadherin–catenin complex as well as the vesicular trafficking machinery (D’Souza-Schorey, 2005), where multiple adaptor and signaling proteins orchestrate trafficking specificity and efficiency.

Clathrin adaptor protein (AP) complexes are important in the sorting of cargoes containing dileucine or tyrosine-based sorting motifs (Bonifacino and Traub, 2003). In epithelial cells, AP1B is the unique isoform that mediates basolateral transport (Folsch et al., 1999; Folsch, 2005). Although AP1B is closely related to the more ubiquitously expressed form of AP1, AP1A,
HR-ECDT was overexpressed in HEK293 cells and immunoprecipitated. It preferentially binds the E-cadherin dimer in vivo. Myc-tagged ECDT or PIPKιγ associates with cadherin complexes. Immunoprecipitations (IP) and immunoblots (IB) were performed as indicated. NCD, N-cadherin; VECD, VE-cadherin. Normal mouse (mIgG) or rabbit IgG (rIgG) were used as controls. (C) PIPKIγ was important for association with PIPKIγ. A parallel dimeric construct had a greater binding affinity for PIPKIγ. (Fig. S1, C and D). The dimeric construct had a greater binding affinity for PIPKIγ. However, the mechanism by which PI4,5P2 generation is regulated to mediate these trafficking events has not been defined.

Recent studies have unveiled that the spatial targeting and temporal regulation of type I phosphatidylinositol phosphate kinases (PIPKIs) is a critical mechanism for PI4,5P2 generation (Ling et al., 2006). Here we show that in epithelial cells PIPKιγ targets to AJs by a direct interaction with the E-cadherin dimer. PIPKιγ regulates E-cadherin trafficking by acting as a scaffold between E-cadherin and AP complexes. We also demonstrate that localized generation of PI4,5P2 via these complexes is necessary for E-cadherin transport and AJ formation.

### Results

**PIPKιγ interacts with cadherins**

Upon examination of the basolateral membrane in polarized epithelial cells, we found that PIPKιγ colocalized with E-cadherin (Fig. 1 A) but not with occludin (not depicted). PIPKιγ also presented in a cytosolic vesicular compartment and partially colocalized with E-cadherin at this site (Fig. 1 A, arrows). These regions of colocalization were confirmed by constructing vertical sections of z-series images shown in Fig. 1 A, suggesting an interaction between PIPKιγ and a component of AJs. To examine this possibility, E-cadherin and PIPKιγ were immunoprecipitated. As shown in Fig. 1 B, PIPKιγ and E-cadherin associate in vivo, along with other cadherin-associated proteins, demonstrating that PIPKιγ associates with E-cadherin complexes.

N-cadherin and VE-cadherin also associate with PIPKιγ (Fig. 1 B), suggesting that PIPKιγ associates with the classical cadherin complexes.

PIPKιγ is predominantly expressed as two distinct splice variants, PIPKιγ635 and PIPKιγ661, which differ by a 26–amino acid C-terminal extension. HA-tagged PIPKιγ splice variants were expressed in human embryonic kidney 293 (HEK293) cells, and their association with the endogenous N-cadherin complex was analyzed. PIPKιγ635 and PIPKιγ661 both communoprecipitated with N-cadherin indistinguishably (Fig. S1 A, available at http://www.jcb.org/cgi/content/full/jcb.200606023/DC1), indicating that this association does not depend on the PIPKιγ661 C-terminal extension. However, the endogenous PIPKιγ associated with E-cadherin was indistinguishable in apparent molecular weight from PIPKιγ661, which is the predominant splice variant expressed in these cells. To ascertain whether this association was direct, in vitro GST pull-down assays were performed using recombinant GST-tagged PIPKιγ and His-tagged E-cadherin cytoplasmic domain (ECDT). ECDT showed specific binding to both GST-PIPKιγ635 and GST-PIPKιγ661, but not to GST alone (Fig. 1 C) or GST-PIP.Kια (Fig. S1 B).

E-cadherin molecules form lateral homodimers in vivo, and oligomer formation is critical for AJ assembly and stability (Patel et al., 2003). Consequently, we examined if dimerization was important for association with PIPKιγ. A parallel dimeric ECDT was constructed by inserting a heptad repeat (HR) peptide sequence between the His tag and ECDT (Ling et al., 2003; Fig. S1, C and D). The dimeric construct had a greater binding affinity for PIPKιγ compared with the monomeric protein.
either the p120-catenin (ECDγ E-cadherin was expressed in HEK293 cells and assessed for E-cadherin involves other AJ components, wild-type or mutated conserved domain in the type I classical cadherins (Fig. 2 C). Including amino acids 837–847 of E-cadherin that is a highly data demonstrate that PIPKIγ directly interacts within a region of PIPKIγ and p120-catenin with ~10-fold greater affinity compared with the monomer (Fig. 1 D and Fig. S1 F). Consistent with a previous paper (Huber et al., 2005), β-catenin bound the monomeric and dimeric E-cadherin C terminus with the same affinity.

To further determine whether PIPKIγ binding to E-cadherin involves other AJ components, wild-type or mutated E-cadherin was expressed in HEK293 cells and assessed for endogenous PIPKIγ association (Fig. 2 A). Elimination of either the p120-catenin (ECDΔp120; 76EED764 to AAA) or β-catenin (ECDΔβctn; ECD847, deletion of the last 35 amino acids) binding sites had no effect on PIPKIγ association. A chimera of truncated E-cadherin (deletion of the last 70 amino acids) fused to a truncated α-catenin that lacks the β-catenin binding site (Imamura et al., 1999) abrogated both β-catenin (not depicted) and PIPKIγ binding (Fig. 2 A). These results indicate that PIPKIγ binding to E-cadherin is independent of α-, β-, or p120-catenin and narrowed the PIPKIγ interaction region on E-cadherin to residues 837–847. To confirm this putative PIPKIγ binding site, the last 45 amino acids of E-cadherin were truncated (ECD836). This truncation resulted in ablation of both β-catenin and PIPKIγ binding (Fig. 2 B). These combined data demonstrate that PIPKIγ directly interacts within a region including amino acids 837–847 of E-cadherin that is a highly conserved domain in the type I classical cadherins (Fig. 2 C).

To examine whether PIPKIγ modulates E-cadherin function through this direct interaction, we introduced wild-type or p120/β-catenin binding site–deleted Myc-HR–ECDT into MDCK cells to compete with endogenous E-cadherin for PIPKIγ binding (Fig. S2 A, available at http://www.jcb.org/cgi/content/full/jcb.200606023/DC1). Expression of dimeric ECDTγ that specifically binds PIPKIγ resulted in a loss of AJs identified by E-cadherin staining and a cytosolic accumulation of PIPKIγ in cells, indicating that this phenotype is likely caused by sequestration of PIPKIγ (Fig. S2 B). Overexpression of PIPKIγ661 was sufficient to rescue the loss-of-AJ phenotype induced by wild-type Myc-HR–ECDT expression (Fig. S2 C). These data establish that a specific interaction between PIPKIγ and E-cadherin plays a key role in E-cadherin function and appears to be a limiting factor in AJ formation.

**PIPKIγ modulates AJ assembly by facilitating E-cadherin trafficking**

To further determine the functional role of PIPKIγ at AJs, we knocked down endogenous PIPKIγ expression using siRNAs. Although the cellular E-cadherin content was not changed (Fig. 3 C), loss of PIPKIγ caused a striking loss of E-cadherin from the PM with an apparent accumulation in a cytoplasmic compartment (Fig. 3, A and B). Upon loss of PM E-cadherin, the cells spread (Fig. 3 A) and underwent a morphological transition from a polarized epithelial to a more migratory mesenchymal-like phenotype (Fig. 3 D), supporting the requirement for PIPKIγ in E-cadherin–mediated AJ assembly.

In addition, MDCK stable cell lines were generated that inductively express HA-tagged wild-type (PIPKIγ661WT) or kinase-dead PIPKIγ661 (PIPKIγ661KD), the major endogenous PIPKIγ isoform associated with cadherins (Fig. S1 A). Expression of PIPKIγ661WT or PIPKIγ661KD was induced by removing doxycyclin from the growth media (Fig. 3 D). As shown in Fig. S3 A (available at http://www.jcb.org/cgi/content/full/jcb.200606023/DC1), upon PIPKIγ661KD expression, both E-cadherin PM targeting and AJ assembly appeared defective compared with parental or PIPKIγ661WT-expressing cells (Fig. S3 A). These cells formed E-cadherin–mediated cell–cell contacts much more slowly then parental cells when maintained at confluence (Fig. S3 A, 72 vs. 16 h). These observations are consistent with a dominant-negative effect for PIPKIγ661KD and also establish a requirement for PI4,5P₂ generation in AJ assembly.

Our combined results demonstrate a highly specific role for PIPKIγ in the assembly of E-cadherin–based AJs, possibly by modulating the trafficking of E-cadherin. It has been shown that depletion of extracellular calcium by EGTA results in a loss of E-cadherin homoligation, internalization of E-cadherin, disassembly of AJs, and cell scattering (Chitaev and Troyanovsky, 1998).

**Figure 2. Determination of the PIPKIγ-interacting region on E-cadherin.**

[A] Endogenous PIPKIγ was immunoprecipitated from HEK293 cells overexpressing wild-type [ECDwt] or the indicated mutant E-cadherin constructs. [B] Endogenous PIPKIγ was immunoprecipitated from HEK293 cells overexpressing wild-type or truncated E-cadherin constructs. The immunoprecipitates were analyzed as indicated. [C] A schematic model representing the predicted minimum region on E-cadherin required for PIPKIγ binding. TM, transmembrane domain; JMD, juxtamembrane domain; βctnBD, β-catenin binding domain. Regions corresponding to βctnBD in E-, N-, P-, and VE-cadherin were aligned and the conserved residues were highlighted.
Figure 3. **PIPmγ is required for E-cadherin assembly and trafficking.**

(A) MDCK cells were treated with scrambled (Scram.) or PIPmγ-specific siRNA (PIPmγ1). PIPmγ (green) and ECD (red) were visualized by indirect immunofluorescence. Bars, 10 μm. (B) PM or cytosolic fluorescence intensity of cells in A was quantified from six randomly picked fields (5 cells/field) representing three independent experiments were randomly counted. The intensity ratio was plotted using SigmaPlot 8.0. Error bars are ± SEM. (C) PIPmγ and E-cadherin protein levels in cells treated with scrambled (Scram.) or PIPmγ-specific siRNAs (PIPmγ1 and PIPmγ2) were shown by immunoblots. (D) Protein levels of endogenous and inducibly overexpressed PIPmγ (661KD) in stable MDCK lines were determined. Actin was used as a loading control. (E) After surface biotinylation, cells expressing the indicated proteins were lysed before or after exposure to 0.5 mM EGTA at 18°C to examine the original biotinylated E-cadherin (Ori. bio-ECD) and the internalized biotinylated E-cadherin (Intern. bio-ECD). Data was quantified and plotted from four independent experiments. (F) Calcium restoration subsequent to EGTA treatment induced E-cadherin recycling. The extent of this recycling was assessed in cells expressing the indicated proteins. Data were quantified and plotted from from three independent experiments. The values representing surface E-cadherin without calcium rescue were used as control. Error bars are ± SEM.

To explore possible modulation of E-cadherin trafficking by PIPmγ, the exocytic and endocytic trafficking of E-cadherin was quantitated in parental, PIPmγy661WT−, and PIPmγy661KD-expressing MDCK cells.
The direct interaction of PIPKIγ661 with μ1β was confirmed by direct binding of purified components (Fig. 4 A, left), and the in vivo association was established by coimmunoprecipitation (Fig. 4 A, right). PIPKIγ635 did not interact with either μ subunit (Fig. 4 A; Bairstow et al., 2006), indicating that the last 26 residues of PIPKIγ661 are required. In addition, in vitro binding of μ1β-adaptin stimulated the kinase activity of PIPKIγ661 (Fig. 4 D), whereas binding of the soluble ECDT (His-REDT) had no effect on PIPKIγ661 activity under these conditions (not depicted).

Epithelial cells typically express two variants of the AP1 complex, AP1A and AP1B. These AP1 complexes differ only in their μ subunits, μ1α and μ1β, which are almost 80% identical but target to distinct membrane compartments and have distinct functions (Folsch, 2005). To further define the interaction between PIPKIs and the AP1 complexes, we examined the interaction between PIPKIγ and both μ1α and μ1β in parallel experiments. As shown in Fig. 4 B (left), PIPKIγ661 binds to μ1α and μ1β indistinguishably in vitro. The primary binding site of the μ subunits is in the last 26 amino acids of PIPKIγ661 because the C terminus of PIPKIγ661 but not PIPKIγ635 bound the μ subunits (Fig. 4 C). A weak interaction between full-length PIPKIγ635 and the μ subunits was observed under less rigorous GST pull-down conditions, where detergent and BSA concentrations were decreased. This may be consistent with the observations by Krauss et al. (2006), which indicate that there may be a secondary μ subunit binding site in the kinase domain of the PIPKIs. However, in vivo the interaction is specific for PIPKIγ661 and only the interaction between μ1β subunit (AP1B) and PIPKIγ661 was detected (Fig. 4 B, right). The in vivo specificity of PIPKIγ661 for μ1β may be regulated by targeting to AP1B-positive membrane compartments via an interaction with other trafficking components, or the PIPKIγ661–AP1B interaction maybe specifically regulated by other mechanisms.

Endogenous E-cadherin associates with the PIPKIγ–AP1 complex (Fig. 5 A). This association was disrupted by internalization of E-cadherin triggered by extracellular calcium depletion (Fig. 5 B). When calcium was restored and E-cadherin recycling to the PM was triggered, E-cadherin reassembled into the PIPKIγ–AP1 complex (Fig. 5 B). To further examine the interactions between E-cadherin, PIPKIγ661, and AP1, GST pull-down assays were performed. Although there is no direct interaction between the μ1β subunit of AP1 and HR-ECDT, PIPKIγ661 was sufficient to link HR-ECDT to μ1β in a GST pull-down experiment (Fig. 5 C). PIPKIγ661 contains a Yxxφ sorting motif (Y644SPL; Bonifacino and Traub, 2003; Bairstow et al., 2006). The substitution of the tyrosine with phenylalanine in the sorting motif was reported to reduce binding to the μ subunits (Ohno et al., 1998; Aikawa and Martin, 2003; Bairstow et al., 2006). Concurrent with these results, the Y644F mutation diminished PIPKIγ661 binding to μ1β (Fig. 4 C and Fig. 5 C), and consequently the amount of E-cadherin C terminus pulled down by μ1β was considerably reduced (Fig. 5 C), indicating that the interaction between PIPKIγ661 and μ1β-adaptin is necessary and sufficient to link E-cadherin to the AP1B complex.

Because AP complexes play an important role in protein transport, our data suggest that PIPKIγ661 regulates E-cadherin trafficking via a direct interaction with and regulation of AP complexes. Such a model would require AP1B for E-cadherin transport to the PM. To address this hypothesis, we used LLC-PK1 cells, which do not express μ1β (i.e., are AP1B deficient). In LLC-PK1 cells, many basolateral proteins are mistargeted and cell polarity is disrupted (Folsch et al., 1999; Folsch, 2005). To assess the role of μ1β in E-cadherin transport, GFP–E-cadherin was expressed (Fig. 5 D). A small fraction of GFP–E-cadherin was able to translocate to the PM; however, the majority was observed in a perinuclear compartment, indicating inefficient transport of E-cadherin to the PM. In cells expressing...
PIPKIγ regulates E-cadherin transport by recruiting E-cadherin to AP1. (A) E-cadherin or AP1 were immunoprecipitated from polarized MDCK cells. (B) E-cadherin was immunoprecipitated from polarized MDCK cells, polarized MDCK cells treated with 2 mM EGTA for 20 min, or polarized MDCK cells treated with EDTA for 20 min followed by regular medium for 10 min. The immunoprecipitates were analyzed as indicated. (C) GST pull-down assays were performed using 1 μg of GST, GST-μ1B, His- and His-HR-ECDT (γ661), and His-HR-ECDT and analyzed by immunoblotting as indicated. (D) GFP-fused E-cadherin (ECD-GFP) was transiently introduced into parental (μ1B deficient) or μ1B-expressing LLC-PK1 cells using lipofectamine 2000 for 8 h and then visualized using fluorescence microscopy. Bar, 10 μm.

Figure 5. GFP–E-cadherin, there was a greatly enhanced recruitment of endogenous PIPKIγ to GFP–E-cadherin–containing compartments, which is consistent with the association between PIPKIγ and E-cadherin. Upon expression of μ1B in the LLC-PK1 cells, GFP–E-cadherin was efficiently targeted to sites of cell–cell adhesion, and endogenous PIPKIγ colocalized with E-cadherin at AJs (Fig. 5 D). The expression of μ1C, however, did not rescue E-cadherin trafficking to the PM (unpublished data). These data supported a model in which PIPKIγ associates with E-cadherin and this interaction is required for functional recruitment of AP1B to PIPKIγ via its interaction with the YSPL motif in the PIPKIγ661 C terminus.

PIPKIγ regulates E-cadherin transport by recruitment to AP1B compartments

The functional relationship between PIPKIγ and AP1B is reinforced by the observation that endogenous PIPKIγ and AP1 co-localized in vesicle compartments (Fig. 6 A). Both E-cadherin and PIPKIγ partially colocalized with γ-adaptin in cytoplasmic compartments after removal of calcium (Fig. 6 B, arrows), suggesting a functional link between E-cadherin, PIPKIγ661, and AP1 in E-cadherin trafficking. Interestingly, when E-cadherin recycling was triggered by replenishing calcium, we observed that overexpression of PIPKIγ661 enhanced the recruitment of AP1B to PM. In parental MDCK cells, AP1 showed typical perinuclear localization with a small fraction targeting to the PM. When PIPKIγ661 was expressed, AP1 targeted to the basolateral membrane where it colocalized with E-cadherin and PIPKIγ661 (Fig. 6 C). However, when PIPKIγ635 was expressed, AP1 organization was strikingly distinct, as it was concentrated in a central perinuclear compartment with no localization near the PM and little colocalization with E-cadherin (Fig. 6 C). In these cells, E-cadherin was largely trapped in the cytosol and was not efficiently targeted to the PM. In PIPKIγ661KD-expressing cells, AP1 weakly localized beneath the PM or showed strong colocalization of both E-cadherin and PIPKIγ661KD in a large perinuclear compartment, but there was little detectable PM E-cadherin (Fig. 6 C). These data again support a model where both the PIPKIγ–AP1 interaction and PIPKIγ kinase activity are necessary for recruitment of AP1 to the PM and the efficient trafficking of E-cadherin to the PM.

To explore this hypothesis, the internalization and recycling of E-cadherin in cells ectopically expressing PIPKIγ635 was first determined. As shown in Fig. 7 A, when internalization and recycling of E-cadherin was measured by surface biotinylation, overexpression of PIPKIγ635 had a dominant-negative effect and inhibited E-cadherin trafficking to and from the PM compared with parental cells. Again, these results support a functional role for the PIPKIγ661–AP1B interaction in modulation of E-cadherin trafficking. Consistent with this conclusion, both ectopically expressed GFP–E-cadherin (Fig. 7 B) and endogenous E-cadherin (Fig. 6 C) were sequestered in a cytosolic compartment in PIPKIγ635-overexpressing cells, displaying a phenotype similar to that observed when endogenous PIPKIγ was knocked down.

To further characterize these E-cadherin–containing vesicles in PIPKIγ635-expressing cells, we induced accumulation of transferrin receptor (TfnR) in the recycling endosome using an established approach (Sheff et al., 1999). As shown in Fig. 7 C (top), internalized E-cadherin in parental MDCK cells showed partial colocalization with both endogenous PIPKIγ and internalized TfnR, representing the recycling endosome, and colocalization among these three proteins was also observed (Fig. 7 C, arrows). Interestingly, overexpression of PIPKIγ635, which interacts with E-cadherin but not AP1B, blocked E-cadherin colocalization with the TfnR compartment, but E-cadherin did colocalize with PIPKIγ635 (Fig. 7 C, bottom). These data suggest that PIPKIγ661 mediates the transport of E-cadherin from the trans-Golgi network to the recycling endosome, which has been argued to serve as an intermediate between the trans-Golgi network and the basolateral PM (Ang et al., 2004; Folsch, 2005). Further, this data establishes that trafficking of E-cadherin to this compartment requires a functional interaction between PIPKIγ661 and AP1B.

If PIPKIγ serves as an adaptor between E-cadherin and AP complexes, one would expect that an E-cadherin mutant...
lacking or with diminished PIPKIγ binding would not be transported efficiently to the PM. A V832M germline mutation was identified in hereditary diffuse gastric cancer (Yabuta et al., 2002), which lacks the ability to mediate cell–cell adhesion or suppress invasion (Suriano et al., 2003). In these patients, the wild-type E-cadherin gene is repressed, and only the mutant is expressed.

Figure 6. PIPKIγ is functionally related with AP1 and E-cadherin trafficking in vivo. (A) Endogenous PIPKIγ and AP1 colocalize at vesicular compartments. PIPKIγ (FITC, green) and AP1 (Texas red, red) were visualized in MDCK cells by indirect immunofluorescence. (B) PIPKIγ and AP1 colocalize with the internalized E-cadherin. Parental MDCK cells were plated on cover slips and allowed to form AJs. They were then treated with 2 mM EGTA for 30 min. Cells were fixed and indirect immunofluorescence staining was performed using anti-PIPKIγ and anti-γ-adaptin, followed by Cy5-conjugated anti-rabbit IgG antibody and Texas red-conjugated anti-mouse IgG antibodies. Cells were then incubated with 0.5 μg/ml of normal mouse IgG at 37°C for 30 min and then with FITC-conjugated anti-E-cadherin antibodies. Arrows indicate the colocalization of AP1, internalized ECD, and PIPKIγ. (C) PIPKIγ661 recruits AP1 to functional sites of E-cadherin recycling. Parental or PIPKIγ isoform-overexpressing MDCK cells were treated with 2 mM EGTA for 30 min followed by complete medium for 10 min at 37°C to induce the recycling of E-cadherin back to the PM. Immunofluorescence staining was then performed to visualize AP1 (γ-adaptin), E-cadherin, and PIPKIγ. Both horizontal and vertical sections were collected to show precise subcellular localizations. Bars, 10 μm.

Figure 7. The interaction between PIPKIγ and AP1B is necessary for E-cadherin transport. (A) MDCK cells with or without PIPKIγ635 overexpression were used to analyze E-cadherin internalization and recycling by surface biotinylation as described. Immunoblot data from three independent experiments were quantified by ImageJ and plotted with SigmaPlot 8.0. Error bars are ± SEM. (B) Parental, PIPKIγ635-, or PIPKIγ661-expressing MDCK cells were transiently transfected with ECD-GFP. PIPKIγ was visualized by indirect immunofluorescence and ECD-GFP using direct fluorescence microscopy. (C, top) Parental MDCK cells were incubated at 20°C for 2 h, treated with 2 mM EGTA for 30 min, and then fixed and triple-labeled using monoclonal anti-CD71, FITC-conjugated anti-E-cadherin, and rabbit anti-PIPKIγ antibodies. (Bottom) MDCK cells with the expression of HA-PIPKIγ635 induced for 72 h were maintained at 20°C for 2 h. The cells were then fixed and stained using monoclonal anti-CD71, FITC-conjugated anti-E-cadherin, and rabbit anti-HA antibodies. Arrows indicate the colocalization of ECD, PIPKIγ, and the recycling endosome. Bars, 10 μm.
in the carcinomas (Yabuta et al., 2002). Interestingly, the V832M mutation lies in the PIPKIγ binding region. To determine whether this mutation impacts PIPKIγ binding, E-cadherin V832M was introduced into HEK293 cells and its association with PIPKIγ was quantified. This mutant showed a substantially reduced ability to bind PIPKIγ (Fig. 8 A). Consistent with published data (Suriano et al., 2003), β-catenin binding was normal (not depicted). The basolateral transport of this V832M mutation was also explored in both LLC-PK1::μ1β (unpublished data) and MDCK cells using GFP-fused E-cadherinV832M. As shown in Fig. 8 B, although the V832M mutant was visualized on the PM as reported by others (Suriano et al., 2003), a large accumulation of this E-cadherin mutant was observed in a cytosolic compartment. This phenotype was similar to that of wild-type E-cadherin observed in the PIPKIγ635-over-expressing cells (Fig. 7 B) or the LLC-PK1 cells deficient in μ1β (Fig. 5 D). Wild-type E-cadherin in LLC-PK1::μ1β (Fig. 5 D) and MDCK (Fig. 8 B) cells was transported efficiently to the basolateral membrane and little was visualized in the cytosol. This result is consistent with a requirement for an interaction between E-cadherin and PIPKIγ661 for normal trafficking of E-cadherin.

**Discussion**

Proteins that interact with the ECDT not only mediate the elemental functions of E-cadherin, such as AJ assembly, actin organization, and cell proliferation, but also regulate E-cadherin by modulating its expression and trafficking. Here we have shown that PIPKIγ directly binds to both E-cadherin and AP complexes. This dual interaction supports a mechanism for the highly regulated generation of PI4,5P2 to spatially drive the assembly of the trafficking machinery and to specifically control E-cadherin trafficking. These results reveal a novel mechanism where PIPKIγ661 functions as both scaffolding and signaling molecule during E-cadherin trafficking (Fig. 8 C). In this model, the AP complex interacts indirectly with the E-cadherin cargo via the PIPKIγ661 scaffold, which directly binds to AP complexes via a Yxxyb sorting motif in its C terminus. This represents a novel paradigm in which PIPKIγ661 serves as a cargo adaptor for AP complexes. Although PIPKIγ661 binds to both AP1A and AP1B indistinguishably in vitro, we found that PIPKIγ661 preferentially interacts with AP1B in vivo, and regulation of E-cadherin trafficking to the basolateral PM appears to be specific for AP1B. AP1A and AP1B both use Yxxyb sorting motifs for cargo recognition. However, despite the substantial sequence and structural homology of the μ1 subunits, the AP1 complexes are targeted to distinct compartments by an unknown mechanism (Folsch et al., 2003; Folsch, 2005). PIPKIγ661 might be specifically recruited to AP1B-containing membrane domains in vivo via an interaction with one or more other proteins.

PIPKIγ661 may also have additional lower affinity binding contacts with AP1B, as Krauss et al. (2006) recently reported that multiple PIPKIs bind to the μ2 subunit of AP2 complex via the kinase domain. In our hands, the YSPL motif of PIPKIγ661 was the preferential binding site for the μ subunits of AP1 and AP2 (Bairstow et al., 2006), but PIPKIγ635 did bind μ1- and μ2-adaptin subunits under less rigorous conditions (unpublished data), suggesting additional lower affinity interacting sites between the AP complexes and PIPKIγ. For E-cadherin trafficking, the YSPL motif of PIPKIγ661 is the key interaction with AP complexes and subsequent interactions may regulate kinase activity. As the kinase domains of the PIPKI isoforms are highly homologous, other isoforms of PIPKIs (e.g., PIPKIα) may interact with and regulate some AP complex–dependent trafficking events (Barbieri et al., 2001). These putative interactions could be through the conserved kinase domains or, like PIPKIγ661, bind to the μ subunit of AP2.

**Figure 8.** The interaction between PIPKIγ and E-cadherin is required for E-cadherin transport. (A) Indicated E-cadherin constructs were overexpressed in HEK293 cells, and their association with PIPKIγ was determined, quantified, and plotted from four independent experiments. Error bars are ± SEM. (B) MDCK cells were transfected with GFP-ECD(V832M). Cells were analyzed by fluorescence microscopy 8–10 h after transfection. Bar, 10 μm. (C) A model representing the mechanism for PIPKIγ661-mediated E-cadherin trafficking. The dual interaction of PIPKIγ661 with the E-cadherin dimer and the AP complex provides a mechanism for the specific regulation of E-cadherin trafficking. This dual interaction mediates the scaffold between AP complexes and E-cadherin to facilitate the assembly of this cargo into the trafficking machinery. In addition, the interaction between PIPKIγ661 and AP complexes ensures the localized generation of PI4,5P2, which regulates the function of AP complexes and other processes in E-cadherin trafficking.
PIPKIγ661, could be mediated by specific binding partners via interaction with variable regions of the PIPKI isoforms. Nevertheless, because PI4,5P2 is a key modulator of the recruitment and assembly of trafficking machinery (Simonsen et al., 2001; Roth, 2004), the localized generation of PI4,5P2 at sites where E-cadherin and other cargoes are assembled into the trafficking machinery is an indispensable step in this process. This finding suggests that any association between a PIPK and the trafficking machinery must be spatially and temporally regulated. The interaction between the PIPKIγ661 and the AP1B complex fits this criterion, as this association is detected when E-cadherin is recycled back to the PM but not when E-cadherin is being internalized (Fig. 5 B). This observation supports the concept of cellular signals coordinating the interactions between PIPKIγ661 and the AP complexes.

A dileucine motif in the juxtamembrane region of the ECĐT is required for basolateral sorting (Miranda et al., 2001), and this motif was proposed to be a cargo signal recognized by the β subunit of the AP1 complex (Rapoport et al., 1998). There is no solid evidence supporting this interaction at present. However, if this is true, the E-cadherin–PIPKIγ661–AP1B complex could be further stabilized via the interaction of the E-cadherin dileucine motif with the β subunit of AP1B. Alternatively, other trafficking components may recognize this motif and cooperate with PIPKIγ and AP1B to provide specificity.

The E-cadherin–PIPKIγ661–AP1B interaction serves as a foundational signal for exocytic targeting and basolateral sorting of E-cadherin. Indeed, internalized E-cadherin accumulated at the recycling endosome, as indicated by internalized TfnR. This compartment contained, in addition, endogenous PIPKIγ supporting a role for PIPKIγ in trafficking though this compartment. Consistent with this observation, the overexpression of PIPKIγ635, which binds E-cadherin but not μ1B (AP1B), blocked E-cadherin trafficking to the TfnR-positive compartment of the recycling endosome. E-cadherin did colocalize with PIPKIγ635, indicating that PIPKIγ interacts with E-cadherin in this compartment. The combined results demonstrate that the association of PIPKIγ661 with AP1B is require for E-cadherin trafficking through this compartment. The recycling endosome is a major site of AP1B, and this further supports our hypothesis that PIPKIγ functions as an adaptor in E-cadherin trafficking and facilitates E-cadherin transport to and from the recycling endosome via binding to AP1B and generation of PI4,5P2.

E-cadherin endocytosis can occur in a clathrin-dependent (Palacios et al., 2001; Ivanov et al., 2004) or independent manner (Paterson et al., 2003). Calcium removal stimulates E-cadherin endocytosis by the clathrin-dependent pathway (Ivanov et al., 2004). As there is no known Yxxφ sorting motif in the ECĐT, the interaction between PIPKIγ661 and E-cadherin may recruit AP2 for clathrin-dependent E-cadherin endocytosis. Additionally, Arf6 promotes E-cadherin internalization (Palacios et al., 2002) and has been shown to associate with and stimulate the activity of PIPKIγ (Aikawa and Martin, 2003, 2005). Arf6, in cooperation with PI4,5P2, was also shown to directly interact with and promote the recruitment of AP2 to the PM (Krauss et al., 2003; Paleotti et al., 2005). These combined results suggest that PIPKIγ661, E-cadherin, AP2, and Arf6 may cooperate to regulate E-cadherin internalization in epithelial cells. This would position PIPKIγ661 as a nexus between AP complexes and E-cadherin in endocytic recycling. Nevertheless, there is no direct evidence demonstrating that AP2 mediates the internalization of E-cadherin, and further investigation is needed to characterize the role of PIPKIγ in E-cadherin endocytosis.

Our data demonstrates that a loss of PIPKIγ in cultured epithelial cells results in the severe mistargeting of E-cadherin, suggesting a strong functional connection between PIPKIγ and E-cadherin. Interestingly, the PIPKIγ knockout mouse does not share the same phenotype as the E-cadherin knockout mouse (Larue et al., 1994; Di Paolo et al., 2004). This is not surprising, as the knockout phenotypes of other modifiers of E-cadherin function, such as p120-catenin, differ from that of the E-cadherin knockout as well (Pettitt et al., 2003; Davis and Reynolds, 2006). Considering the existence of multiple pathways for E-cadherin trafficking, the roles of these E-cadherin modifiers may only become apparent during the development of specific tissues in later stages of animal development.

Dimerization is an essential property of E-cadherin assembly driving AJ formation (Yap et al., 1997). The association of both PIPKIγ and p120 catenin with the E-cadherin dimer may be a mechanism to functionally regulate E-cadherin assembly and could promote AJ formation by stimulating E-cadherin clustering. Because PIPKIγ specifically binds to E-cadherin dimers, the in situ PI4,5P2 generation resulting from this interaction may drive other local complementary cellular events, such as actin reorganization (Janmey and Lindberg, 2004). Actin assembly is important not only in AJ assembly but also for E-cadherin internalization/exocytosis (D’Souza-Schorey, 2005). The association of PIPKIγ with E-cadherin may be crucial for downstream signaling, as Rac and phosphoinositide-3 kinase are activated by E-cadherin and both regulate the stability of AJs by modulating actin assembly (Noren et al., 2001; Yap and Kovacs, 2003; D’Souza-Schorey, 2005). Phosphoinositide-3 kinase requires PI4,5P2 for signaling, and Rho family small G proteins regulate some PIPKI isoforms (Fruman et al., 1998).

As a result, PIPKIγ may also regulate AJ assembly through local cooperation with phosphoinositide-3 kinase and small G protein signaling.

The generation of phosphoinositide messengers upon assembly of AJs has implications beyond simple control of E-cadherin trafficking. Because E-cadherin is a major suppressor of invasion of epithelial tumors, the cell biological data suggest that PIPKIγ may play a similar role. In exploring this possibility, we have discovered that a loss of E-cadherin correlates with a loss of PIPKIγ in human breast cancers (unpublished data). This finding supports a physiological role for PIPKIγ in assembly of E-cadherin junctions and potentially a role in progression of epithelial tumors.

Materials and methods

Constructs and antibodies
The C terminus of E-cadherin was amplified by PCR and constructed into normal or modified (Ling et al., 2003) pET128 to generate the His-tagged E-cadherin tail or HR-Ecadherin tail, which were then subcloned into pCMV-Myc vector (CLONTECH Laboratories, Inc.). Wild-type E-cadherin,
E-cadherinΔ120ctn, E-cadherinΔ3ctn, and E-cadherin/Δctn were provided by B. Gumbiner (University of Virginia, Charlottesville, VA). E-cadherin36 was amplified by PCR, and E-cadherinV832M was generated using the QuikChange II Site-Directed Mutagenesis kit (Stratagene) according to the manufacturer’s instructions. Both μ 1 and μ 1β constructs were provided by I. Mellman (Yale University, New Haven, CT). CDNAs encoding N-terminal truncated μ 1 and μ 1β (1-135 aa truncated) were amplified by PCR and subcloned into pET42. All of the PI PK constructs were created as described previously (Ling et al., 2002; 2003; Bairstow et al., 2006). Duplexes of siRNA oligos (for both human and mouse: aagctctataggcttcagttgac) were synthesized by Dharmacon.

Monoclonal antibodies for E-cadherin (recognizing the C terminus), N-cadherin, human β VE-cadherin, β-catenin, γ-adaptin, and FITC-conjugated anti-E-cadherin were purchased from Transduction Laboratories. The H68.4 monoclonal anti-CD71 (TfnR) antibody was purchased from BioGenex Inc. Polyclonal PI PKγ antibody was generated as described previously (Ling et al., 2002). Regular mouse and rabbit IgG and secondary antibodies were obtained from Jackson Immunoresearch Laboratories. Anti-β antibody was purchased from Covance. Anti-Myc and FITC-conjugated anti-Myc were obtained from Upstate Biotechnology. HRP-conjugated anti-GST antibodies were purchased from GE Healthcare. Anti-E-cadherin antibodies recognizing the ectodomain were purchased from Zymed Laboratories (monoclonal, for immunoblotting) and Sigma–Aldrich (rat monoclonal, for immunofluorescence).

Cell culture, transfection, immunofluorescence, and confocal microscopy

MDCK-Tet-Off cells (CLONTECH Laboratories, Inc.) and HEK293 cells were cultured in Dulbecco’s modified eagle medium (Mediatech, Inc.) supplemented with 10% FBS (Invitrogen). Lysates of human umbilical vein endothelial cells were obtained from E. Bresnick (University of Wisconsin-Madison, Madison, WI). MDCK cells were transfected using FuGENE 6 (Roche) for 48 h, and then 100 μg/ml hygromycin B was added to the medium to select stable clones and 10 mg/ml doxycycline was used to suppress PI PKγ expression. To induce expression, doxycycline was removed for 72 h. HEK293 cells were transfected using the calcium phosphate–DNA coprecipitation method for 48 h. For siRNA knockdown, MDCK cells in a 6-well plate were transfected twice at 48 h intervals using the calcium phosphate–DNA coprecipitation method. Cells were analyzed 48 h after the second transfection. LLC-PK1 cell lines were provided by I. Mellman and cultured as described previously (Folsch et al., 1999).

Indirect immunofluorescence and confocal microscopy were performed as described previously (Ling et al., 2002). For triple labeling, double labeled samples were blocked by 0.5 mg/ml of normal mouse IgG in 3% BSA/PBS at 37°C for 30 min, rinsed in PBS twice, and incubated with FITC-conjugated anti-E-cadherin or anti-Myc antibodies in 3% BSA/PBS at 37°C for 1 h. Confocal images were acquired using photomultiplier tubes through LaserSharp2000 (Carl Zeiss MicroImaging, Inc.) with a PlanApo ×100 oil objective (NA 1.4) on an inverted microscope (Eclipse TE2000; Nikon) with Radiance 2100 MP Rainbow (Bio-Rad Laboratories). Z series were acquired by sequentially scanning FITC, Texas red, or Cy3 channel at 0.3-μm steps. Single sections were exported to Photoshop CS2 (Adobe) for final image processing. Fluorescence intensity was quantified using ImageJ 1.62 (National Institutes of Health) and plotted using SigmaPlot 8.0.

Immunoprecipitation, GST pull-down assay, and immunoblotting

Cells were lysed in lysis buffer (50 mM Tris-Cl, pH 7.5, 150 mM NaCl, 0.5% NP-40, 1 mM EDTA, 1 mM PMSF, and 10% glycerol), and then used for immunoprecipitation (Ling et al., 2002). The immunocomplexes were separated by SDS-PAGE and analyzed. Unless indicated, immunoprecipitations were performed using 800 μl of cell lysate from one confluent 60-mm dish. One fourth of each precipitate and 20 μl of each lysate were analyzed. GST-tagged PI PKγ, PI PKγ35, and PI PKγ661, histagged E-cadherin, or HR-E-cadherin tail were purified from Escherichia coli, and GST pull-down assays were performed (Ling et al., 2003). One fourth of the GST beads for each pull down were loaded on the gel and 20 μl of each purified protein was loaded as an input control. Images were scanned and exported to Photoshop CS2 for final processing. Intensity of bands was quantified using ImageJ 1.62 and plotted using SigmaPlot 8.0.

Calcium depletion, surface biotinylation, and trafficking of E-cadherin

Cells were allowed to grow on coverslips for 72 h to reach confluence and were then incubated with 2 mM EGTA for 20 min before performing indirect immunofluorescence. Confluent MDCK cells grown in 24-mm-diam Transwells (Costar) were biotinylated by 1 mg/ml sulfo-NHS-SBiotin (Pierce Chemical Co.) and analyzed as described previously (Ie et al., 1999).

Cells were lysed in 500 μl of lysis buffer and one third of the precipitates were analyzed. Internalization of E-cadherin was induced by 0.5 mM EGTA at 18°C. To measure the recycling of E-cadherin, MDCK cells were treated with 2 mM EGTA for 40 min at 37°C and chased in normal medium, and then surface biotinylation was performed.

PI PKγ activity assay

Activity of 10 μg of purified recombinant PI PKγ proteins was assayed against 20 μg of Folsch Brain Extract III as previously described (Di Paolo et al., 2002). Kinase activity was quantified using Storm 840 (Molecular Dynamics) and plotted using SigmaPlot 8.0.

Online supplemental material

Fig. S1 shows that PI PKγ directly binds E-cadherin. Fig. S2 shows that the direct interaction with PI PKγ is important for E-cadherin assembly. Fig. S3 shows that functional PI PKγ is required for E-cadherin assembly. Fig. S4 shows that a modification of PI PKγ activity has no effect on global PI4,5P2 level. Online supplemental material, including Figs. S1–S4 and supplemental Materials and Methods, is available at http://www.jcb.org/cgi/content/full/jcb.200606023/DC1.

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