Phosphatidylinositol 4-kinase III beta regulates cell shape, migration, and focal adhesion number

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ABSTRACT Cell shape is regulated by cell adhesion and cytoskeletal and membrane dynamics. Cell shape, adhesion, and motility have a complex relationship and understanding them is important in understanding developmental patterning and embryogenesis. Here we show that the lipid kinase phosphatidylinositol 4-kinase III beta (PI4KIIIβ) regulates cell shape, migration, and focal adhesion (FA) number. PI4KIIIβ generates phosphatidylinositol 4-phosphate (PI4P) from phosphatidylinositol and is highly expressed in a subset of human breast cancers. PI4KIIIβ and the PI4P it generates regulate a variety of cellular functions, ranging from control of Golgi structure, fly fertility, and Akt signaling. Here, we show that loss of PI4KIIIβ expression decreases cell migration and alters cell shape in NIH3T3 fibroblasts. The changes are accompanied by an increase in the number of FA in cells lacking PI4KIIIβ. Furthermore, we find that PI4P-containing vesicles move to the migratory leading edge during migration and that some of these vesicles tether to and fuse with FA. Fusion is associated with FA disassembly. This suggests a novel regulatory role for PI4KIIIβ and PI4P in cell adhesion and cell shape maintenance.

INTRODUCTION

A culture of genetically identical NIH3T3 fibroblasts displays a striking visual diversity. A single microscopic field of view is populated with elongated cells, round cells, and yet others with extraordinarily complex geometries. Furthermore, some NIH3T3 cells are solitary while others cluster themselves into multicellular groups. Last, some fibroblasts are stationary while others are motile. These moving cells display marked differences in speed and directionality. This fascinating architectural and behavioral diversity is at the root of processes such as organismal development and patterning.

Cell shape has important implications in cell function (Bellas and Chen, 2014; Gilbert and Weaver, 2017). For example, the spreading of a cell in two-dimensional culture regulates both sensitivity to apoptosis and proliferative capacity (Chen et al., 1997). The spreading and shape of a cell results from a complex and dynamic interaction among its cytoskeleton, the plasma membrane, and the extracellular matrix (ECM) (Mogilner and Keren, 2009). In fish keratinocytes, which assume a hemispherical appearance in two-dimensional culture, cell shape is determined by a balance between the prosurvival force of actin polymerization and tension in the plasma membrane (Keren et al., 2008). In other cells, cell shape is regulated by lipid constituents of the plasma membrane (Wen et al., 2018) and by membrane-binding proteins that induce curvature (Nishimura et al., 2018). In migratory fibroblasts, which assume a wedge-shaped form, cell shape is additionally regulated by the presence and strength of adhesive structures at the cell periphery (Satulovsky et al., 2008). These adhesive structures provide sites for attachment to the ECM and the sensing of its mechanical properties. Cellular adhesions also serve as sites for the generation of contractile forces (Bosch-Fortea and Martin-Belmonte, 2018).

Perhaps the most important of mammalian adhesive structures are focal adhesions (FAs) (Burridge, 2017). FA are oval-shaped, multi-protein complexes, generally 2 μm wide and 3–10 μm long, coupling the actin cytoskeleton to the ECM via transmembrane integrin heterodimeric receptors (Parsons et al., 2010). During cell migration, FAs are created from smaller integrin-containing structures described as either focal complexes or nascent adhesions (Parsons et al., 2010). In a mature FA, the cytoplasmic tails of the integrin molecules are linked to actin filaments via adapter proteins.
such as talin, paxillin, zyxin, and vinculin (Kanchanawong et al., 2010). FA are sites of mechano-transduction and activate a diverse array of signal transduction pathways (Geiger et al., 2009). FA number and size are critical in the maintenance of migratory capacity (Kim and Wirtz, 2013a), cell spreading (Kim and Wirtz, 2013b), and cell shape (Chen et al., 2003; Mogilner and Keren, 2009).

In this report, we find that the lipid kinase phosphatidylinositol 4-kinase III beta (PI4KIIIβ) has an important role in regulating cell shape, migration, and FA number. A Golgi-resident enzyme, PI4KIIIβ is one of four mammalian proteins that generate phosphatidylinositol 4-phosphate (PI4P) from PI (Balla, 1998, 2013). PI4 kinases have been implicated in several human cancers (Waugh, 2012). PI4KIIIβ is a likely human oncogene based on its high expression in a subset of human breast tumors and its ability to co-operate with the Rab11a GTPase to activate signaling through Akt (Morrow et al., 2014). PI4KIIIβ and its homologues have multiple physiological roles, ranging from controlling Golgi structure in yeast to regulating male fertility and development in Drosophila (Godi et al., 1999; Polevoy et al., 2009). PI4KIIIβ also regulates in vitro morphogenesis of human breast cells (Pinke and Lee, 2011).

RESULTS

PI4P vesicles move to the migratory leading edge

To explore a role for PI4P in cell migration, we used two PI4P biosensors, GFP-FAPP1 and GFP-P4M, to study intracellular PI4P localization during NIH3T3 migration. The GFP-FAPP1 biosensor consists of GFP conjugated to the Pleckstrin Homology domain of FAPP1 (four-phosphate-adaptor protein 1) (Balla, 2007). The GFP-P4M biosensor consists of GFP conjugated to a single P4M domain containing residues 546–647 of the Legionella pneumophila SidM protein (Hammond et al., 2014). Both biosensors bind to PI4P directly. However, FAPP1 binds to PI4P only when complexed to the Arf-1 GTPase (Balla et al., 2005).

In migrating NIH3T3 cells, FAPP1 shows a broadly perinuclear distribution consistent with Golgi localization (Figure 1A; Supplemental Video S1). Large (0.8–1.6 μm diameter) and small vesicles (0.2–0.45 μm diameter) are visible. The large vesicles (lv) are predominantly perinuclear while the small vesicles (sv) localize to the cell periphery. PI4M shows a somewhat similar localization pattern with many lv in the perinuclear region. Different from FAPP1, many of these perinuclear vesicles appear hollow (hv) (Figure 1A; Supplemental Video S2). As is the case with FAPP1, there are sv in the cell periphery but these seem to be much more numerous in P4M-transfected cells. In direct contrast to FAPP1 and as observed previously in COS7 cells (Hammond et al., 2014), the P4M reporter localizes to plasma membrane structures. Some plasma membrane features at the migratory leading edge are reminiscent of ruffles (r), actin-rich structures that move away from the leading edge and derive from poor lamellar adhesion to the growth substrate (Borm et al., 2005). The difference between FAPP1 and P4M staining presumably reflects the requirement that FAPP1 interact with Arf-1-associated PI4P pools.

During migration, the lv of both P4M and FAPP1 show limited intracellular movement and remain generally perinuclear (Supplemental Videos S1 and S2). In contrast, the peripheral sv show a high degree of mobility (Supplemental Videos S1 and S2). In addition, we observe multiple P4M and FAPP1 sv moving to the migratory leading edge and then disappearing (Figure 1B; Supplemental Videos S3 and S4). This is consistent with vesicular delivery of PI4P-containing vesicles, both Arf-1 associated and Arf-1 free forms, to the plasma membrane. The absence of FAPP1 staining at the plasma membrane suggests that once the vesicle delivers the PI4P/Arf-1 cargo, either Arf-1 dissociates or PI4P is converted by PI4P kinases to another PI. In the case of P4M, the persistence of PI4P in some parts of the leading edge indicate that some of the membrane delivered PI4P remains as PI4P. In addition to their movement to the plasma membrane, many small P4M and FAPP1 vesicles show considerable directional freedom. Both retrograde and anterograde movements are observed (Supplemental Videos S3 and S4). However, the movement of sv is generally toward the migratory leading edge.

The movement of vesicles toward the migratory leading edge is mirrored in the visibly unequal intracellular distribution of both FAPP1 and P4M vesicles in the migratory leading edge compared with the trailing edge. To further explore this, we used automated image analysis to quantify the distribution of PI4P vesicles in migratory and nonmigratory cells (Supplemental Videos S5–S8 show representative cells). In migratory cells, we counted vesicles in both the leading and trailing edges. In stationary cells, we counted vesicles present in two opposing edges of the cell. In stationary cells there are a generally equal number of P4M and FAPP1 marked PI4P vesicles at two opposite edges (Figure 2B, open circles; Supplemental Videos S6 and S8). On the other hand, migratory cells show a greater number of vesicles at their leading edge compared with the trailing edge (Figure 2B, closed circles; Supplemental Videos S5 and S7). This is consistent with the idea that the transport of PI4P to the migratory leading edge is a part of the directional cell migration machinery.

PI4KIIIβ deletion decreases cell migration

To explore a role for PI4P in cell migration, we used CRISPR to delete the PI4KIIIβ gene in mouse NIH3T3 fibroblasts. PI4KIIIβ generates PI4P in the Golgi (Balla and Balla, 2006; Balla, 2013). We then tested two independent PI4KIIIβ-null lines for migratory capacity in wound healing assays. As shown in Figure 3A (left panel), the two CRISPR lines close a wound much slower that parental cells. Wild-type (WT) NIH3T3 cells close half the wound in ~18 h, while the null lines take ~25 h. This defect in wound closure is rescued by re-expression of WT PI4KIIIβ (Figure 3A, middle panel).

The best characterized function of PI4KIIIβ is the generation of PI4P from PI (Balla, 1998). However, PI4KIIIβ also interacts with the Rab11a GTPase and is part of the Rab11a-dependent pathway controlling endosome function (de Graaf et al., 2004; Polevoy et al., 2009; Burke et al., 2014). To determine which of these functions were involved in migration regulation, we expressed wt-PI4KIIIβ, kinase dead PI4KIIIβ (KD-PI4KIIIβ) (Zhao et al., 2000) or a PI4KIIIβ mutant that does not interact with Rab11a (N162A) (Burke et al., 2014) in a CRISPR line. These lines were then tested for migratory capacity. As shown in Figure 3A (right panel), a CRISPR line expressing wt-PI4KIIIβ or the Rab11a-binding mutant N162A had similar closure times to those of parental fibroblasts. On the other hand, KD-PI4KIIIβ was unable to rescue would closure kinetics. This indicates that the ability to generate PI4P, rather than Rab11a interaction, is required PI4KIIIβ-mediated control of migration. Figure 3B shows protein expression of the PI4KIIIβ-rescued cell lines.

PI4KIIIβ regulates stress fiber appearance

In yeast, the PI4KIIIβ homologue Pik1 is required for maintaining Golgi structure (Walch-Solimena and Novick, 1999; de Graaf et al., 2004). Because cell migration involves repositioning of the Golgi between the nucleus and the leading edge (Kupfer et al., 1982) and functional alterations of the Golgi affect migration (Xing et al., 2016; Ahat et al., 2019), we next investigated whether or not PI4KIIIβ depletion affected Golgi appearance. When
stained for markers of either the cis- or the trans-Golgi (GM130 and TGN46, respectively), NIH3T3 cells lacking PI4KIIIβ have Golgi that are visibly indistinguishable from WT cells (Figure 4A). Moreover, in scratch wound assays, PI4KIIIβ depletion does not visibly affect Golgi positioning to face the wound (Figure 4B). Microtubule structure also appears normal as does centrosome alignment away from the wound (Figure 4B). One the other hand, PI4KIIIβ depletion causes an increase in the number of cells with numerous stress fibers (Figure 4C). In a population of normal NIH3T3 cells, most cells show few stress fibers with only 3.1 ± 5% of the population having more than six fibers per cell. In contrast, the PI4KIIIβ-depleted lines have a much larger fraction (11.2 ± 8% and 18.9 ± 13%) in their population with high stress fiber content.
**PI4KIIIβ regulates cell shape**

In our studies of the cell cytoskeleton and cell migration, we noticed that cultures of the CRISPR-deleted NIH3T3 cells had a very different morphological appearance under phase than either parental or rescued cells. In our experience, most cultured NIH3T3 cells assume one of three broad shapes. The first is an elongated form (Figure 5A) and approximately half of WT NIH3T3 cells assume this shape (Figure 5A, right panel). The second most common shape is what we term “multidirectional.” These cells are roughly rectangular in shape with multiple pseudopodial protrusions. Approximately 25% of WT cells are of this type (Figure 5A, right panel). The remaining cells, which have a smaller, generally spherical appearance, we classified as “other.” The loss of PI4KIIIβ leads to a redistribution of cell shapes in both of the CRISPR lines. In freely migrating conditions, the number of elongated cells in the PI4KIIIβ-deleted cells decreases by almost 50% and the number of multidirectional cells more than doubles (Figure 5A, top panel). Similarly, cells present in the wound of a scratch migration assay show an increase in the number of multidirectional cells and a decrease in elongated ones (Figure 5A, bottom panel). Representative fields are shown in Figure 5B. As is the case with the wound healing assay, wt-PI4KIIIβ and the Rab11-binding mutant (N162A) were able to restore WT shape distribution to the CRISPR lines, while the KD-PI4KIIIβ did not (Figure 5C). Parental cells and CRISPR lines rescued with either WT PI4KIIIβ or N162A had 45–50% of cells as elongated, while the KD-rescued cells had nearly 35% as multidirectional, similar to the original CRISPR line. Similarly, parental cells and CRISPR lines rescued with either WT PI4KIIIβ or N162A had 25–35% of cells as multidirectional, similar to the original CRISPR line. The line of best fit is shown.
PI4KIIIβ regulates focal adhesions

To further explore the regulation of cell shape and migration by PI4KIIIβ, we investigated what cellular structures might be interacting with PI4P-containing vesicles. We reasoned that FA might be one such structure since FA-dependent adhesion is likely to be involved in cell shape, migration control, and stress fiber formation (Chen et al., 2003; Kim and Wirtz, 2013a). To investigate possible FA and PI4P interaction during cell migration, we simultaneously imaged them in migrating cells using fluorescent Talin and P4M reporters. At the migratory leading edge (Figure 6A; Supplemental Video S9), PI4P vesicles can be observed moving to FA. Vesicles dock with FA and then disappear, presumably delivering their cargo to FA and the PI4P becoming metabolized. Similar behavior occurs at FA of the trailing edge (Figure 6B; Supplemental Video S10).

To further explore delivery of PI4P vesicles to FA, we used a Bessel Lattice Light Sheet microscope (BLLSM) to image migrating NIH3T3 cells. The BLLSM illuminates a sample using a thin structured illumination beam and allows for long-term imaging with high axial and temporal resolution and with minimal photobleaching (Planchon et al., 2011). Using the BLLSM, we visualized 50 PI4P vesicle-FA fusion events in five independent migrating WT NIH3T3 cells at 3 s/vol. Cells were observed for ∼25 min each. As shown in Figure 6C, approximately two-thirds of fusion events were associated with disappearance of the FA within 3–6 s. This suggests that PI4P or cargo in PI4P vesicles is involved in FA disassembly. To test this idea,
we used an automated image analysis program to count FA in WT and CRISPR-deleted NIH3T3 cells (Horzum et al., 2014). As shown in Figure 7A, WT NIH3T3 cells \( (n = 60) \) had an average of 56.7 ± 17 FA per cell. CRISPR-deleted cells \( (n = 59) \), on the other hand, had significantly (t test, \( p < 1 \times 10^{-5} \)) more FA, averaging 84.6 ± 42 per cell. Rescue of the CRISPR line by PI4KIII\( \beta \) re-expression returned the FA number to WT levels (58.4 ± 24). Thus, the number of FA/cell is dependent, at least in part, on PI4KIII\( \beta \).

To assay FA signaling capacity in the CRISPR lines, we subjected cells to hyperosmotic stress by culturing them in 400 mM sucrose. Hyperosmotic stress causes cell shrinkage and activates FA kinase (FAK) autophosphorylation at tyrosine 397 (Lunn and Rozengurt, 2004) and Src-mediated phosphorylation of multiple FAK residues (Calalb et al., 1996; Westhoff et al., 2004). FAK hyperphosphorylation is part of Src-mediated FA disassembly (Calalb et al., 1996; Westhoff et al., 2004). As shown in Figure 7B, parental cells have strong FAK phosphorylation (Y-861) in response to sucrose treatment but PI4KIII\( \beta \)-deleted cells do not. This reduced phosphorylation is rescued by expression of wt-PI4KIII\( \beta \). Thus, loss of PI4KIII\( \beta \) allows for more FA per cell but these have impaired signaling consistent with attenuated disassembly.

**DISCUSSION**

The central findings of this paper are that the PI4KIII\( \beta \) is an important regulator of cell shape, cell migration, and FA disassembly. The best known function of PI4KIII\( \beta \) is the generation of PI4P from PI (Balla and Balla, 2006; Balla, 2013). In addition, PI4KIII\( \beta \) binds to the Rab11a GTPase in a kinase-independent manner (de Graaf et al., 2004; Polevoy et al., 2009; Burke et al., 2014) and is involved in Rab11a-dependent control of Akt signaling (Jeganathan et al., 2008; Morrow et al., 2014) and endosome function (de Graaf et al., 2004; Polevoy et al., 2009). In our case, a catalytically inactive version of PI4KIII\( \beta \) rescues neither the reduced migration nor the cell shape defects in PI4KIII\( \beta \)-deleted cells. On the other hand, a PI4KIII\( \beta \) mutant that has kinase activity but does not bind the Rab11a GTPase (Burke et al., 2014) does rescue motility and cell shape distribution to a level equivalent to WT. This indicates that control of cell migration and shape is independent of Rab11a interaction but requires PI4P generation.

Consistent with an importance for PI4P in migration, our live-cell imaging shows directed movement and fusion of PI4P-containing vesicles to the migratory leading edge and to FA. There are also more PI4P vesicles between the nucleus and the leading edge than between the nucleus and the trailing edge. These vesicles are likely to derive from the Golgi and therefore could be produced by both PI4KIII\( \beta \) and PI4KIII\( \beta \) (Balla, 1998) in the Golgi. PI4P is a precursor for two phosphoinositides, PI(4,5)P\(_2\) and PI(3,4,5)P\(_3\), that have a well-documented role in cell migration control (Schink et al., 2016).
FIGURE 5: Loss of PI4KIIIβ alters cell shape distribution. (A) Two independent lines of PI4KIIIβ-deleted cells have different population shape distributions (depicted in the right panels) in both freely migrating conditions and in the wound of a scratch. The fraction of elongated cells in either line of CRISPR cells is significantly \( p < 0.001, t \text{-test} \) lower than in a population of WT cells. Similarly, the fraction of elongated cells in either line of CRISPR cells is significantly \( p < 0.001, t \text{-test} \) higher than in a population of WT cells. Results are the mean and SD of triplicate independent measurements of at least 200 cells each from a minimum of 20 randomly selected fields. (B) Representative fields of each cell type. Scale bar is 20 μm. (C) Re-expression of either WT- or N162A-PI4KIIIβ restores elongated and multidirectional shape distribution to the CRISPR cells. Expression of KD-PI4KIIIβ does not rescue cell shape changes and their elongated of multidirectional composition is similar to the CRISPR control and significantly different from WT cells \( p < 0.05, t \text{-test} \). Results are the mean and SD of quadruplicate independent measurements of at least 200 cells from at least 20 randomly selected fields. (D) Distance traveled over time was monitored for 50 WT NIH3T3 cells with either elongated or multidirectional morphologies. Tracks for 10 cells of each type are shown. Multidirectional cells have a
significantly reduced average angular rotation that elongated cells ($p < 0.05$, t test). (E) Quantification of migration patterns in WT cells with either multidirectional or elongated morphologies. Mean angular rotation or velocity is presented as the mean and SD of 50 cells collected over at least 30 min. Angular rotation is significantly lower in the multidirectional population compared with the elongated one ($p < 0.05$, t test). Comparison between Angular Rotation and Velocity between WT and CRISPR lines. Mean angular rotation or velocity is presented as the mean and SD of 50 cells collected over at least 30 min.

Some of the delivered PI4P could be converted to PI(4,5)P$_2$ and PI(3,4,5)P$_3$ at the leading edge. While PI4P is in large intracellular molar excess over PI(4,5)P$_2$ and PI(3,4,5)P$_3$, localized pools of PI(4,5)P$_2$ in the plasma membrane that regulate ion transport are dependent on an influx of Golgi-derived PI4P (Dickson et al., 2014). We propose that pools of PI(4,5)P$_2$ and PI(3,4,5)P$_3$ regulating cell migration could similarly be dependent on PI4P transport from the Golgi.
FIGURE 6: Continued.
Our observation that PI4KIIIβ deletion alone produces defects in cell migration suggests that the PI4P produced by PI4KIIIβ is an important part of cell migration that is not duplicated by PI4KIIβ or other PI4P-generating enzymes. Silencing PI4KIIIβ in a breast tumor cell line has previously been shown to attenuate cell migration (Tokuda et al., 2014). Here we describe a similar role for PI4KIIIβ in the migration of a noncancer line. We believe that migratory deficiency in PI4KIIIβ-deleted cells is directly related to the changes of cell shape we observe. The majority of WT NIH3T3 cells display an extended and elongated morphology in culture. On the other hand, a population of PI4KIIIβ-deleted cells show fewer elongated cells and more with multiple pseudopodia. We term the multi-pseudopod phenotype “multidirectional” because the cells attempt to move in several directions at once. Elongated cells, on the other hand, generally move along a single axis. The net consequence of the two different types of motion in the different cells is that the multidirectional cells cover smaller distances than elongated ones because they are constantly making small angle turns. Thus, in a wound healing assay, a population of elongated cells will close the wound faster than multidirectional ones. It is important to note that, as a group, both elongated and multidirectional PI4KIIIβ-deleted cells have the same migratory parameters as their respective.

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**FIGURE 6:** PI4P Vesicles move to and fuse with FAs. (A) Visualization of PI4P and FAs in the leading edge of a migrating NIH3T3 cells. The time series indicates a single vesicle observed from the boxed inset. The fusing vesicle (white arrow) appears at 95 s. Scale bar in the large image is 10 μm and is 3 μm in the enlarged images. (B) Visualization of PI4P and FA in the trailing edge of a migrating NIH3T3 cells. The time series indicates a single vesicle seen from the boxed inset. The fusing vesicle (white arrow) appears at 175s. Scale bar in the large image is 10 μm and is 3 μm in the enlarged images. (C) BLLSM imaging of PI4P vesicles (green P4M reporter) fusing with FA (red Vinculin reporter) in a migrating cell. Histogram shows that the majority of fusions events (n = 50) cause FA destruction within 3–6 s. The bottom panels show enlarged fields of the top panel where two fusion events, one causing FA destruction and the other not, can be seen.
elongated and multidirectional counterparts in WT cells. They travel at the same velocity and have the same turning behaviors. We propose that deletion of PI4KIIIβ is not impairing the intrinsic migration machinery per se but rather is altering the relative balance of fast-moving and slow-moving cells in the overall population.

The increase in FA number we observe in PI4KIIIβ-deleted cells indicates that specific PI4KIIIβ cargo is necessary for normal FA disassembly. FA link the actin cytoskeleton to the ECM. During cell migration, FA are created immediately behind the migratory leading edge and are disassembled at later stages of migration (Parsons et al., 2010; Burridge, 2017). Our live-cell imaging indicates that PI4P-containing vesicles move to and fuse with FA at both the leading and the trailing edges. The majority of these fusion events are associated with rapid FA disassembly and we favor a model where PI4KIIIβ regulates FA number by directing the delivery of cargo that initiates FA disassembly. Further consistent with the idea that FA disassembly involves a PI4P-dependent process is the impaired FAK tyrosine phosphorylation in PI4KIIIβ-deleted cells. Hyperphosphorylation of FAK by Src is associated with movement of FAK away from FA and a decrease in FA number (Westhoff et al., 2004; Hamadi et al., 2010). Possibly, PI4P-containing vesicles could facilitate the movement of Src to FAK on FA to potentiate adhesion disassembly.

Another well-characterized process of FA disassembly is the endocytic destruction of a FA via microtubule-mediated delivery of dynamin (Ezratty et al., 2005, 2009). This activates FA destruction through clathrin-mediated endocytosis. We suggest that delivery of endocytic machinery to sites of FA may be dependent on PI4P-mediated transport. Alternatively, the transport of proteases such as calpain or MMP1 that destroy FA components (Franco et al., 2004; Takino et al., 2006) could be transported in or on PI4P vesicles.

We favor the idea that the increase in FA number by the loss of PI4KIIIβ is directly responsible for changes in cell shape and migration. The change in FA number is likely responsible for the increase in stress fiber appearance (Figure 4C). Stress fibers are long bundles of actin polymers that are typically linked to FA at one or both ends (Burridge and Guilloy, 2016). FA regulate migration because they mediate connection to the ECM and intracellular contractile forces (Parsons et al., 2010; Kim and Wirtz, 2013a). While some adhesion is necessary for cell migration, too many FA could impair migration because of excessive adhesion. FA have a reciprocal relationship with cell shape in that the positioning of adhesions determines cell shape but altering the cell shape itself modifies FA location in a cell (Chen et al., 2003; Lehnert et al., 2004). However, it is a possibility that the...
changes in cell shape, migration, and FA number we observe are regulated by independent PI4KIIIβ and PI4P-dependent pathways.

In summary, we have identified an import role for PI4KIIIβ in cell shape, migration, and adhesion. PI4KIIIβ is likely to be a human oncogene (Waugh, 2012; Morrow et al., 2014) and we propose that its role in cancer is related to its regulation of these processes. In the future it will be important to identify the cargo involved.

MATERIALS AND METHODS
Cell lines and culture
The NIH3T3 fibroblast cell line was obtained from ATCC (Manassas, VA). NIH3T3 cells were cultured in DMEM (Sigma Aldrich) supplemented with 10% fetal bovine serum (FBS) (Thermo Scientific), 1 mM sodium pyruvate (Thermo Scientific), and 1 mM penicillin and streptomycin (Thermo Scientific). Cells are passaged following treatment with 0.25% trypsin protease (GE Healthcare Life Sciences) and counted using TC20 cell counter (Bio-Rad). The PI4KIIIβ-targeted RNA sequences for CRISPR deletion of PI4KIIIβ exons 4-5 were 5′-CACGACCTGTAGTCCGGAAT-3′, 5′-GGCTCCCTACTCTGATCAG-3′, 5′-ATAAGCTCCTGCCCAGTCTC-3′ (Santa Cruz sc-430739).

Wound healing and cell tracking
For wound healing and cell tracking, cells were seeded at a density of 1 × 10^5 cells/ml in a 24-well ImageLock (EssenBio) plate containing a culture 2-insert well (ibidi) and incubated at 37°C at 5% CO2; 24 h later, the insert was removed. Images were acquired at a magnification of 10× every 30 min using the IncuCyte ZOOM Scratch Wound tool (EssenBio). For cell tracking, cells were seeded at 5000 cells/ml in a 24-well ImageLock plate (EssenBio) and images were acquired at a magnification of 10× using the IncuCyte ZOOM (EssenBio). Cells were individually tracked and analyzed in ImageJ.

Western blot
Cells were lysed in radioimmunoprecipitation assay buffer (Tris-HCl, pH 7.4, 50 mM; NaCl, 150 mM; NP-40 1%; sodium deoxycholate, 0.5%; sodium dodecyl sulfate, 0.1%; EDTA, 2 mM; sodium fluoride, 50 mM) supplemented with protease and phosphatase inhibitor cocktails (Roche, Mississauga, Canada). Protein concentrations were determined by Bradford protein assay (Bio-Rad, Mississauga, Canada). Loading buffer was added to 30 μg of protein lysate and resolved by SDS–PAGE. The protein was then transferred onto a polyvinylidene difluoride membrane (Millipore, Tokyo, Canada) and probed using antibodies for PI4KIIIβ (BD Biosciences 611817; Mississauga, Canada), pan-actin (Cell Signaling Technology 4968), tubulin (1:1000, Cell Signaling Technology 3873; Whitby, Canada), mouse α-FAK (1:200 Thermo Fisher #396500), rabbit α-Phospho-FAK (1:200 Thermo Fisher #44-426), as well as anti-rabbit HRP-linked (Cell Signaling Technology catalogue no. 7074). Bands were detected with a MicroChem Chemiluminescent system (DNR Bio-Imaging Systems, Toronto, Canada) and intensities were quantified by densitometry using GelQuant (DNR Bio-Imaging Systems).

Plasmids and transfections
P4M-GFP and mApple-Talin-N-10 were obtained from Addgene (51469 and 54951, respectively). FAPP-GFP was a gift from Tamas Balla. Cells were grown and transfected on μ-Dish 35-mm, high-wall dishes (ibidi). For individual P4M and FAPP1 experiments, cells were transfected with 1 μg of DNA mixed with Lipofectamine 2000 (Invitrogen) according to manufacturer’s protocol. For P4M and FA imaging, cells were transfected with 1.5 μg of P4M, 1.5 μg of Talin, and 12 μl of Lipofectamine 2000 reagent in 250 μl.

Sucrose stimulation of FAK phosphorylation
Proliferating NIH3T3 cells and derivatives thereof were serum starved for 2 h and then incubated for 10 min in serum-free DMEM media (Sigma) or stimulated for 10 min in serum-free media containing 400 mM sucrose (Sigma). Cells were collected and pelleted in ice-cold phosphate-buffered saline (PBS). Cell pellets were lysed in ice-cold RIPA buffer containing protease inhibitors (Complete, Millipore Sigma #11873580001) and phosphatase inhibitors (PhosSTOP, Millipore Sigma #4906845001) and frozen in 5× LSB.

Microscopy
For fixed-cell microscopy, NIH3T3 cells were seeded on High Precision 1.5H cover glass (Deckglaser) in 12-well cluster plates (Corning). Cells were fixed with 3.7% paraformaldehyde/PBS for 10 min, permeabilized for 10 min in 0.1% Triton X-100/PBS, and incubated for 1 h in 3% FBS/0.1% Triton X-100/PBS at room temperature. Rabbit α-GM130 (ThermoFisher PA5-95727) and mouse α-tubulin (Cell Signaling 3873S) or γ-tubulin (Abcam 27074) were diluted 1:300 in blocking buffer and incubated overnight at 4°C. Mouse α-Vinculin (Millipore Sigma V9131) was diluted 1:300 in blocking buffer and incubated on the cells for 2 h (RT). After triplicate PBS washes, Goat anti-Mouse IgG (H+L) Antibody, Alexa Fluor 647 (Thermo Fisher A21235) was diluted 1:500 in PBS and incubated on cells for 1 h (RT). Cells were mounted on microscope slides (Fisher Scientific 12-544-7) with EverBrite Mounting Medium with DAPI (Biotium 23002) and sealed with CoverGrip Coverslip Sealant (Biotium 23005). Epifluorescent images were acquired with a Zeiss AxioObserver Z1 or a Zeiss AxioObserver M2 microscope with a 63× Plan-Apochromat 1.4 NA oil objective and Zen Blue 2.3 software. Image processing was carried out using ImageJ. The identification of FA immunostained for vinculin were processed as described by Horzum et al. (2014).

Live-cell microscopy
Time-lapse series of cells in phenol-free DMEM were recorded at 37°C on a Zeiss LSM880-AxioObserver Z1 microscope equipped with 63× Plan Achromat (NA 1.4) oil objective. We used an AiryScan detector in FAST mode and the bandpass emission filter 495–620 nm. Live-cell time-lapse series were also captured using a Bessel Beam Lattice Light Sheet microscope equipped with detection objective (Sermrock; NA of 0.54). A 5-mm coverslip was loaded to a piezo stage and submerged in a 37°C water bath. Time-lapse series of 175 Z-planes and 350 vol were captured at 63× magnification with 5 Piezo Offset 50 and an interval of 0.4 μm. Raw data were assembled into .tif hyperstacks using ImageJ. Three-dimensional renders were then generated for analysis using Imaris.

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