Spatial regulation of MCAK promotes cell polarization and focal adhesion turnover to drive robust cell migration

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ABSTRACT The asymmetric distribution of microtubule (MT) dynamics in migrating cells is important for cell polarization, yet the underlying regulatory mechanisms remain underexplored. Here, we addressed this question by studying the role of the MT depolymerase, MCAK (mitotic centromere-associated kinesin), in the highly persistent migration of RPE-1 cells. MCAK knockdown leads to slowed migration and poor directional movement. Fixed and live cell imaging revealed that MCAK knockdown results in excessive membrane ruffling as well as defects in cell polarization and the maintenance of a major protrusive front. Additionally, loss of MCAK increases the lifetime of focal adhesions by decreasing their disassembly rate. These functions correlate with a spatial distribution of MCAK activity, wherein activity is higher in the trailing edge of cells compared with the leading edge. Overexpression of Rac1 has a dominant effect over MCAK activity, placing it downstream of or in a parallel pathway to MCAK function in migration. Together, our data support a model in which the polarized distribution of MCAK activity and subsequent differential regulation of MT dynamics contribute to cell polarity, centrosome positioning, and focal adhesion dynamics, which all help facilitate robust directional migration.

INTRODUCTION Cell migration is critical for embryonic development, immune response, wound healing, and pathological diseases such as tumor metastasis (Bravo-Cordero et al., 2012; Reig et al., 2014). It is a complex cellular behavior that integrates chemical and mechanical cues from polarized signaling networks, cellular adhesions, and the dynamic actin and microtubule (MT) cytoskeletal networks (Lauffenburger and Horwitz, 1996; Broussard et al., 2008; Kaverina and Straube, 2011; Ridley, 2015). Actin polymerization at the cell front drives the formation of protrusions that provide the force for cell locomotion, and actomyosin exerts contractile forces that move the cell body forward (Cramer et al., 1994). While the contribution of the actin cytoskeleton has been extensively explored, the roles of MTs are relatively understudied.

MTs contribute to multiple aspects of cell migration that are highly coordinated with each other. First, MTs contribute to the directional persistence of migration in several cell types by promoting the initiation and maintenance of cell polarity (Waterman-Stoer and Salmon, 1997; Mikhailov and Gundersen, 1998; Takesono et al., 2010; Kaverina and Straube, 2011; Zhang et al., 2014; Garcin and Straube, 2019). For example, disruption of MT dynamics by nocodazole or Taxol slows cell locomotion by inhibiting the formation of lamellipodia (Liao et al., 1995; Mikhailov and Gundersen, 1998). MT growth also forms a positive feedback loop with signaling molecules, such as the Rho family GTPase Rac1, which promotes actin polymerization and protrusion formation at the leading edge that facilitates cell polarization (Waterman-Stoer et al., 1999; Wittmann et al., 2003). Second, MTs spatially and temporarily regulate the assembly and disassembly of integrin-based focal adhesions (FAs) (Kaverina et al., 1998; Small et al., 2002; Broussard et al., 2008; Stehbens and Wittmann, 2012). FAs are transmembrane complexes that couple the actin cytoskeleton to the extracellular matrix (ECM),
and their dynamic turnover is essential for regulating cellular adhesive properties and intracellular force distribution that impact migration (Ridley et al., 2003). Nascent adhesions form near the protruding leading edge, and most undergo subsequent disassembly in the cell body and trailing edge for cell locomotion (Broussard et al., 2008; Vicente-Manzanares et al., 2009). Targeting MTs to FAs promotes FA disassembly, possibly mediated by cargo transport to and from the adhesion sites (Kaverina et al., 1999; Broussard et al., 2008; Efimov et al., 2008; Stehbens and Wittmann, 2012). Furthermore, MTs can contribute to migration by interacting with the actin cytoskeleton (Wadsworth, 1999; Akashi et al., 2014) and by transporting membrane vesicles to the leading edge (Kaverina and Straube, 2011).

Efficient cell movement requires an asymmetric distribution of MT dynamics along the migration axis, which are controlled by numerous MT dynamics regulators in the cell (Kaverina and Straube, 2011; Etienne-Manneville, 2013). Live imaging revealed that stable, long-lived MTs are enriched at the leading edge to support protrusion, whereas more dynamic MTs are found in the cell body and cell tail (Wadsworth, 1999; Ganguly et al., 2012). An example of a spatially regulated MT dynamics modulator in migrating cells is the MT destabilizer, stathmin/Op18, which is regionally active in the cell tail (Niethammer et al., 2004). Stathmin is inhibited by phosphorylation via the Rac1-PAK1 cascade, and the level of phosphorylation correlates with the level of cell migration in cancer cells (Wittmann et al., 2004; Liu et al., 2013). Another key MT destabilizer is MCAK (mitotic centromere-associated kinesin), which negatively regulates MT polymer levels in interphase (Kline-Smith and Walczak, 2002). MCAK is overexpressed in numerous cancer and tumor cells, and its expression level correlates with increased lymphatic invasion, metastasis, and poor prognosis (Nakamura et al., 2007; Sanhaji et al., 2011). In addition, knockdown of MCAK impairs migration of human bronchial epithelial cells transformed with mutant Ras to migrate and to invade Matrigel (Zaganjor et al., 2014). Previously, MCAK was shown to regulate migration in endothelial cells (Braun et al., 2014), where the authors proposed that a Rac1-Aurora A pathway selectively suppresses MCAK activity at the leading edge to promote regional MT growth (Braun et al., 2014). However, it is not known whether MCAK activity is spatially distributed in migrating cells or what steps of migration are affected by loss of MCAK. Here, we show that MCAK is needed for polarized protrusions to generate a leading edge. Our results from an MCAK Förster resonance energy transfer (FRET) biosensor suggest that MCAK activity is spatially distributed with higher activity at the trailing edge of cells, consistent with the observations of long stable MTs in the leading edge and more dynamic MTs in the cell body and trailing edge reported by others. In addition, overexpression of Rac1 is dominant to the loss of MCAK, suggesting that Rac1 acts downstream of or parallel to the MCAK pathway. Interestingly, inhibition of Aurora A kinase alters the lifetime of the MCAK biosensor, suggesting that Aurora A inhibition does alter MCAK conformation, but Aurora A inhibition does not cause a significant effect on migration in this cell type, suggesting an Aurora-independent aspect of MCAK function. We postulate a model in which MCAK is a key component of a feedback loop that amplifies the gradient of MT polymerization activity along the migration axis to generate a dominant protrusion during leading-edge migration, which highlights the importance of tightly regulated MT dynamics for proper cell migration.

RESULTS
MCAK contributes to cell migration in RPE-1 cells
To examine MCAK function in cell migration, we tested whether MCAK is required for migration in nontransformed retinal pigment epithelial cells (RPE-1). Using a transwell migration assay, knockdown of MCAK by RNA interference (RNAi) resulted in a significant decrease in the relative number of cells that migrated, and a similar reduction in migration was seen with MCAK CRISPR knockout cells (MCAK<sup>−/−</sup>) (Figure 1, A–C). As a parallel test, we performed a wound healing assay and found that MCAK knockdown cells were unable to close the wound as robustly as control cells (Figure 1, D and E) and had a 17% decreased velocity (Figure 1F). To test the specificity of the knockdown, we developed a knockdown/rescue strategy using a small interfering RNA (siRNA) generated against the 3′ UTR of MCAK. Knockdown of MCAK using this siRNA reduced wound closure to a similar extent as the siRNA to the coding region of MCAK (Supplemental Figure S1). Rescue with mEmerald-MCAK (em-MCAK) was not achieved because the lentiviral vector resulted in overexpression of MCAK, which also caused a decrease in wound healing (Supplemental Figure S1), consistent with previous studies (Braun et al., 2014). Together, these results indicate that MCAK contributes to robust cell migration in RPE-1 cells. While the MCAK knockout cells would be the ideal system to explore the effects of MCAK on migration, these cells were phenotypically unstable, as they became large and abnormally shaped after multiple passages. This is likely because depletion of MCAK increases lagging chromosomes that lead to genomic instability (Kline-Smith et al., 2004; Bakhroum et al., 2009a,b). Hence, we pursued the rest of the experiments using RNAi knockdown.

Loss of MCAK perturbs directional persistence and reduces FA turnover
To examine how MCAK knockdown impacts the movement of individual cells, we used time-lapse imaging to directly visualize migrating cells on a two-dimensional (2D) surface and quantified migration by semiautomatically tracking the nuclei over time. During the 4 h time window examined, most control RPE-1 cells established a polarized morphology with a stable protrusion and showed highly persistent directional movement typical of RPE-1 cells (Figure 2, A, top, and B, left). In contrast, ∼30% of cells with MCAK knockdown did not form or maintain a major protrusion but displayed numerous short-lived protrusions with excessive membrane ruffling (Figure 2A, bottom, and Supplemental Video 1). These defects in protrusion correlated with reduced migration distance (Figure 2B, right) and consequently resulted in a 40% decrease in the overall migration velocity (Figure 2C). It is possible that cell movement was decreased because the individual cells were unable to move effectively and/or because they did not move in a directed manner. We therefore calculated the instantaneous velocity of the cells during each 10 min imaging interval and found that MCAK knockdown caused a 26% reduction in the median instantaneous velocity (Figure 2D). To address whether the directional persistence of migration was also affected, we calculated the ratio of the displaced migration (Euclidean distance between the start- and endpoints) to the accumulated migration path over time and found a 24% reduction in the directional persistence (Figure 2E). Overall, these data demonstrate that MCAK promotes efficient movement in RPE-1 cells by both controlling the rate and persistence of movement.

Because our analysis of cell movement after MCAK knockdown revealed defects in cell morphology, and it is known that MTs are important for the establishment and/or maintenance of cell polarity (Schiff and Horwitz, 1980; Takesono et al., 2010; Kamath et al., 2014), we asked whether MCAK knockdown disrupts cell polarization. Cells were treated with control or MCAK siRNAs for 30 h and then were seeded at low density to allow cells to attach and spread for 20 h before being processed for immunofluorescence. Most
control cells showed a classic polarized morphology with a broad front protrusion and a narrow trailing edge (Figure 3A, left). In contrast, after MCAK knockdown, many cells displayed a smaller rounded shape without clear polarity (Figure 3A, right). In comparison to controls, the median cell area after MCAK knockdown decreased by 43% (Figure 3B), suggesting that cells may have defects in their ability to spread. To compare the cell shape, we measured the eccentricity or roundness of the two populations of cells. Linear objects have an eccentricity of 1, whereas a perfect circular object has an eccentricity of 0. Compared with control cells, MCAK knockdown cells were 20% less eccentric, indicating that they were less polarized (Figure 3C). Together, these results show that MCAK contributes to the establishment and/or maintenance of cell polarity, which is a key step for productive migration.

A key event in cell polarization is centrosome reorientation in which centrosomes relocate from the center of the cell to the front of the nucleus toward the direction of movement (Luxton and Gundersen, 2011). To test whether MCAK knockdown disrupted centrosome reorientation, we performed a scratch wound assay and quantified the relative position of the centrosome to the centroid of the nucleus (Figure 3, D and E). An angle of less than 90° represents a centrosome in front of the nucleus, whereas an angle of greater than 90° represents a centrosome behind the nucleus. In control cells, the median angle was 45°, and 70% of cells had centrosomes positioned with an angle of less than 90° (Figure 3F). In contrast, after MCAK knockdown, the median angle was 103°, and only 45% of cells had centrosomes positioned with an angle of less than 90°, suggesting that the defects in cell polarity after MCAK knockdown are coupled with defects in centrosome repositioning.

Our data suggest that the defects in motility after MCAK knockdown are in part due to the inability of the cells to form stable protrusions in the direction of migration. In addition, translocation of the cell body may be affected if the adherence of the cell to the substrate is also perturbed by loss of MCAK. Because MT dynamics influence the turnover of FAs (Stehbens and Wittmann, 2012), and MCAK controls MT dynamics, we postulated that the distribution or turnover of FAs might be affected after MCAK knockdown. We first utilized fixed cell analysis of cells after MCAK knockdown and qualitatively analyzed the distribution of MTs and FA using a scratch-wound assay in cells...
expressing paxillin–green fluorescent protein (GFP) (Supplemental Figure S2). Two hours after wounding, both control cells and cells with MCAK knockdown had a polarized morphology directed at the wound edge. MCAK knockdown cells had a visible increase in the staining intensity of FAs, suggesting that loss of MCAK increased the stability of FAs. To measure the dynamics of the FAs, we carried out time-lapse microscopy of these migrating cells using established methods to determine the rates of assembly and disassembly of FAs at the leading edge of cells (Figure 4, A and B, and Supplemental Video 2) (Theisen et al., 2012; Stehbens and Wittmann, 2014; Kenific et al., 2016). Briefly, the assembly phase of the paxillin-GFP fluorescence intensity profile was fitted with a logistic function (Figure 4B, green), the disassembly phase was fitted with a single exponential decay function (red), and the lifetime was determined as the amount of time the fluorescence intensity was above half the maximum derived from the assembly and disassembly (black arrow).

In comparison to controls, the median rate of FA assembly after MCAK knockdown decreased by 17% and the median rate of disassembly was reduced by 45%, resulting in a 52% increase in FA lifetime (Figure 4C). In addition, the fluorescence intensity profiles of

**FIGURE 2:** MCAK knockdown results in defective polarized 2D cell migration. (A) Representative DIC images of individual cells from the indicated times during 4 h time-lapse microscopy. Scale bar, 25 µm. (B) Pseudo-colored migration tracks of individual cells during the 4 h imaging. Cell position was determined by manually tracking the center of the nucleus over time. Each colored line represents a single cell, and the starting position was normalized to (0,0) on the axes. n = 60 cells for control and 61 cells for MCAK knockdown. (C, D) Quantification of the migration velocity of individual cells determined as the total distance traveled divided by the total time of migration (C) or as the instantaneous velocity of cells during each 10 min imaging interval (D). (E) Directional persistence was determined as the displacement divided by the total migration path. Data are represented as a Tukey box plot, in which the median (line), first and third quartiles (box), whiskers (±1.5 times the interquartile range), and mean (red +) are shown. Outliers are indicated as black dots. p values were determined by the two-tailed Mann–Whitney U test, *, p < 0.05, ***, p < 0.001.
centrosomes positioned in front of the nucleus toward the wound
nucleus (blue with the centrosome (green with asterisk) relative to the centroid of the nucleus. (E) Schematic illustrating the strategy for determining the position of the centrosome relative to the centroid of the nucleus from four independent experiments, with \( n = 296 \) cells for control and \( n = 502 \) cells for MCAK-KD. The horizontal line represents the median, and the whiskers represent the interquartile range.

**MCAK activity is spatially regulated in cells**

In migrating cells, MTs are more stable at the leading edge than in the cell body or trailing edge (Wadsworth, 1999; Ganguly et al., 2012), providing a framework for how cells coordinate the distribution of components by their MT and actomyosin cytoskeletons for persistent migration. Consistent with this idea, in migrating endothelial cells, it was proposed that MCAK activity is locally inhibited at the cell front by a Rac1-Aurora A pathway to stabilize MT growth (Braun et al., 2014). Aurora A kinase directly phosphorylates MCAK to inhibit its activity (Zhang et al., 2008). Similarly, Aurora B phosphorylation inhibits the activity of MCAK, which can be assayed by a conformational change using FRET (Andrews et al., 2004; Lan et al., 2004; Ohl et al., 2004; Ems-McClung et al., 2013; McHugh et al., 2019). To reveal the distribution of MCAK activity in migrating cells, we generated a fluorescence lifetime imaging microscopy (FLIM)-FRET biosensor based on its conformational regulation and performed FLIM on migrating cells in a wound healing assay. In this assay, active MCAK will have a shorter lifetime and inactive MCAK will have a longer lifetime (Figure 5A). Cells expressing either mEmerald-MCAK (em-MCAK) donor alone or the FLIM–FRET sensor mEmerald-MCAK-mCherry (em-MCAK-mCh) both had mEmerald fluorescence distributed throughout the cells (Figure 5B, left) with an enrichment of fluorescence intensity at the trailing edge relative to the leading edge (Supplemental Figure S3A). An analysis of the lifetimes revealed regional differences between the leading and trailing edges (Figure 5B, right). To compare the differences in lifetime at the leading edge versus the rear of the cell, we drew three 4 × 4 pixel area boxes at the front of the cell in the direction of the wound edge and at the base of the rear of the cell before the extended tail (Figure 5C). The lifetimes within these regions were averaged and then compared between the control and the sensor. At the leading edge of the cell, the MCAK sensor cells had an average lifetime similar to that of the control cells (Figure 5D, left), suggesting that MCAK is in an inactive state. In contrast, the average lifetime at the rear of the cell was significantly shorter in cells expressing the sensor (Figure 5D, right), suggesting that MCAK is in an active conformation at the trailing edge. These results are consistent with the idea that MCAK is more active in the tail than at the leading edge, which positions MCAK to play an important role in the regional distribution of MT dynamics for proper cell polarization and persistent migration.

FAs after MCAK knockdown often appeared more complex (Figure 4B). In control cells, most of the intensity profiles fit a bell-shape curve (56 of 60 in control), whereas 34% (20 of 58 in MCAK knockdown) of the FA intensity profiles in MCAK knockdown showed discontinuous disassembly profiles with more than one local maxima (Figure 4B, middle and right). These data suggest that these FAs may undergo cycles of aborted disassembly attempts, possibly due to the inability to sustain FA disassembly (Miranti and Brugge, 2002; Meenderink et al., 2010). Taken together, our data show that MCAK knockdown disrupts FA dynamics and suggest that MCAK-mediated MT dynamics are required to promote efficient FA turnover at the leading edge of migrating cells.

FIGURE 3: MCAK knockdown causes defects in cell polarization. (A) Representative images of individual cells that were grown for 20 h after either control or MCAK knockdown. Cells were stained with rhodamine-phalloidin to show the actin cytoskeleton. Scale bar, 50 \( \mu \)m. Quantification of the cell area (B) and eccentricity (C) of cells in A. \( n = 118 \) for control and 150 for MCAK knockdown. Data are represented as a Tukey box plot, in which the median (line), first and third quartiles (box), whiskers (±1.5 times the interquartile range), and mean (red + ) are shown. Outliers are indicated as black dots. Data are pooled from three independent experiments and were compared using the two-tailed Mann–Whitney U test, ****, \( p < 0.0001 \). (D) Representative control or MCAK knockdown images of cells at a wound edge fixed and stained to visualize MTs (magenta), centrosomes (green), and DNA (blue). The arrows point in the direction of cell movement toward the wound. Scale bar, 20 \( \mu \)m. (E) Schematic illustrating the strategy for determining the position of the centrosome (green with asterisk) relative to the centroid of the nucleus (blue with + ). Angles (\( \alpha \)) between 0 and 90° represent centrosomes positioned in front of the nucleus toward the wound edge, whereas angles >90° represent centrosomes positioned behind the nucleus. (F) Dot plots showing quantification of the angle of the centrosome relative to the centroid of the nucleus from four independent experiments, with \( n = 296 \) cells for control and \( n = 502 \) cells for MCAK-KD. The horizontal line represents the median, and the whiskers represent the interquartile range.
If MCAK activity is spatially regulated by Aurora A kinase, then we would predict that treatment of cells with an Aurora A kinase inhibitor would result in a decrease in the lifetime of the MCAK sensor at the leading edge of cells. Because Aurora A kinase is a negative regulator of MCAK activity, we postulate that the lifetimes in the presence of an Aurora A kinase would be shorter, indicative of the more active conformation of MCAK. We found, however, that treatment of cells with Aurora A kinase inhibitor resulted in an increase in the lifetime of the MCAK sensor at both the leading edge and the trailing edge of cells compared with control treatment (Figure 5E), suggesting either that MCAK was less active in the presence of the Aurora A inhibitor or that MCAK underwent a different conformational change, resulting in lower FRET. The Aurora A kinase inhibitor did not simply disrupt the FLIM properties of the mEmerald donor, as the inhibitor had no effect on the lifetimes of the em-MCAK donor at the leading or trailing edge of cells (Supplemental Figure S3B). The inhibitor also resulted in a decrease in the intensity of fluorescence of the sensor at both the leading and trailing edges, with a lesser effect on the control em-MCAK (Supplemental Figure S3A).

To ask whether the changes in the lifetime of the MCAK sensor by the Aurora A inhibitor correlated with changes in the migration of cells, we carried out a wound healing assay and found that there was no significant effect on the migration of the cells with Aurora A inhibition (Supplemental Figure S4, A and B). One possible explanation for the lack of an effect of the Aurora A inhibitor on cell migration is that Aurora A regulation of MCAK is most important during the initial polarization of the cells. To test this idea, we plated cells in the presence of Aurora A inhibitor and allowed them to adhere to the plate overnight and begin to establish polarity. In both control and Aurora A inhibitor–treated cells, the cell morphologies were similar (Supplemental Figure S4C), and both the area and eccentricity were indistinguishable from controls (Supplemental Figure S4D, left). As an alternative, we first allowed the cells to attach to the plates and then added the inhibitor for 4 h to more closely mimic the conditions utilized in the FLIM–FRET experiments (Supplemental Figure S4D, right). Similar to the cells incubated in inhibitor overnight, Aurora A inhibition had no effect on the area or the eccentricity of the cells as they established polarization, and both of these measurements were indistinguishable from those of the cells that adhered in the presence of inhibitor. In summary, our results show that inhibition of Aurora A alters the conformation of MCAK but does not result in defects in cell migration or cell polarity in RPE-1 cells.
Rac1 acts downstream from or controls pathways parallel to MCAK

Cell polarization is coordinated by the activation of Rac1, which promotes actin polymerization, protrusion formation, and membrane ruffling at the leading edge of migrating cells (Hall, 1998; Nobes and Hall, 1999). MT growth stimulates Rac1 activation (Waterman-Storer et al., 1999), which would suggest that increased MT stability from inhibition of MCAK might promote Rac1 activation and cause the MCAK knockdown phenotype we observed in cells. However, other studies have suggested that MCAK is downstream of Rac1 in which Rac1 signals through Aurora A–dependent inhibition of MCAK (Braun et al., 2014). To ask whether Rac1-induced cell protrusion is overridden by MCAK knockdown, we transiently expressed constitutively active (CA) GFP-tagged Rac1 in cells after MCAK knockdown and then examined the morphology of the cells as described above. Expression of GFP slightly lowered the area of control cells relative to previously observed control cells (Figure 3), which was likely due to differences in the timing of experiments, but did not alter the significant difference in eccentricity of control cells versus MCAK knockdown cells (Figure 6, A, top, B, and C). In contrast, expression of CA-GFP-Rac1 enhanced cell spreading as evidenced by broad lamellipodia in both control and MCAK knockdown cells (Figure 6A, bottom). Quantification of the morphological parameters confirmed that CA-GFP-Rac1–expressing cells showed dominant effects of increased cell area and reduced eccentricity, in both control and MCAK knockdown cells (Figure 6, B and C).
Our results raise the question of whether or how Aurora A might regulate MCAK activity during cell migration. In our system, we find that Aurora A inhibition has no effect on cell polarization or by enhanced MT growth that occurs in the absence of MCAK. These results also raise the possibility that MCAK is a key component of a feedback loop that mediates MT growth at the leading edge of migrating cells.

**DISCUSSION**

One emerging idea in the field of cell migration is that cytoskeletal dynamics need to be spatially controlled, but the molecular mechanisms demonstrating differences in local concentrations of proteins or a spatial distribution of their activity have not been well-established. Our work takes advantage of an MCAK FRET-based biosensor that reports conformational changes in MCAK that correlate with its activity (Ems-McClung et al., 2013). Our findings represent the first demonstration of spatial changes in MCAK activity at the leading and trailing edges of cells, which likely contribute to the localized differences in MT dynamics for polarized cell migration (Figure 7A). Abolishing MCAK activity disrupted cell migration by negatively affecting several important aspects, including cell polarization/protrusion formation, centrosome reorientation, and the turnover of FAs.

Our results evoke new thoughts about how localized MT dynamics are coupled to the formation of a stable leading edge. Loss of MCAK results in enhanced membrane ruffling and short-lived protrusions, which are likely due to the stabilization of MTs resulting from the lack of MCAK-induced MT depolymerization. Consistent with this idea, stabilizing MTs by Taxol treatment leads to the formation of membrane protrusions evenly distributed around the cell perimeter rather than mainly at the cell front (Schiff and Horwitz, 1980; Waterman-Storer and Salmon, 1999). Down-regulation of the MT destabilizer, stathmin/Op18, also enhances membrane ruffles in macrophage activation (Xu and Harrison, 2015), supporting the idea that global stabilization of MTs disrupts polarity. The enhanced formation of random protrusions and membrane ruffling upon the depletion of MCAK indicates that MCAK might be involved in regional suppression of signals to prevent the formation of extra protrusions around the periphery, thus ensuring a dominant lamellipodial protrusion for efficient migration. MT growth locally activates Rac1, which in turn promotes protrusion formation (Waterman-Storer et al., 1999); thus it is possible that MCAK knockdown results in enhanced global MT growth, which promotes Rac1 localization leading to short-lived protrusions and ruffling all over the cell cortex. In support of this idea, our data showed that most CA-Rac1-GFP transfected cells have broad lamellipodia in both control and MCAK knockdown cells, suggesting that Rac1 acts downstream of MCAK, although it is also possible that Rac1 acts in a parallel pathway. It is possible that the dramatic changes in the cell cortex are due to Rac1 effects on actin polymerization that simply override any potential changes in MT dynamics. However, others have also proposed that Rac1 is important in mediating MCAK-mediated MT dynamics. A previous study postulated that Rac1 was upstream of MCAK and worked through a Rac1-Aurora A pathway to locally inhibit MCAK (Braun et al., 2014). One model that could reconcile both observations is that there is a feedback loop in which Rac1 stimulates protrusion formation, which signals through Aurora A or other modulators to further locally inhibit MCAK, resulting in MT stabilization and persistent cell polarization. These stabilized MTs then may serve as tracks on which material is transported to the leading edge of cells to generate a single dominant lamellipodial protrusion.

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migration. This contrasts with previous work by the Myers group, who found that inhibition of Aurora A caused reduced cell migration, which correlated with faster MT dynamics at the leading edge and slower MT dynamics at the trailing edge of cells (Braun et al., 2014). In their model they propose that a major function of Aurora A is to phosphorylate MCAK at S196/S192 to limit its activity at the leading edge of cells. If this were true, then we would expect that Aurora A inhibition would decrease MT lifetime at the leading edge of cells. If this were true, then we would expect that Aurora A inhibition would decrease MT lifetime at the leading edge, the MT lifetimes at the trailing edge were longer. One possibility is that the effects of Au

Another surprise from our work is that Aurora A inhibition results in an increase in fluorescence lifetimes of our MCAK biosensor, resulting in the release of MTs from the centrosome (Ganguly et al., 2011a,b), so perhaps inhibition of Aurora A suppresses this MT release, which changes the organization of MTs in that region and prevents centrosome repositioning. In support of this idea, we also noted in earlier work that there was an accumulation of MTs in the perinuclear area after MCAK inhibition (Kline-Smith and Walczak, 2002), suggesting that MCAK may act on short MT plus ends near the centrosome or that MCAK activity may also be important at MT minus ends.

Another key component in migration is the formation and release of FAs, which provide traction for motility. We showed that MCAK knockdown increased leading-edge FA lifetime and disturbed FA turnover mainly by reducing the disassembly rate, resulting in more persistent FAs. It was shown previously that MT target endocytosis that may facilitate FA disassembly (Efimov et al., 2008; Efimov and Kaverina, 2009). MTs may provide tracks for the transport of cargoes that are involved in FA assembly and disassembly to and from the adhesion sites (Stehbens and Wittmann, 2012; Seetharaman and Etienne-Manneville, 2019). For instance, integrin-linked kinase is required for MT stabilization and caveolin transport to FAs (Wickstrom et al., 2010). Caveolin mediates endocytosis that may facilitate FA disassembly (Kiss, 2012; Stehbens and Wittmann, 2012). In addition, the MT-dependent transport and accumulation of matrix metalloproteases at FAs promote exocytosis that degrades ECM and consequently facilitates FA disassembly (Sakai et al., 2006; Wiesner et al., 2010). Because MCAK does not associate directly with FAs but rather with the plus ends of MTs (Moore et al., 2005) and our unpublished results), where it controls their dynamics (Kline-Smith and Walczak, 2002; Braun et al., 2014), phosphorylation of MCAK by Aurora B kinase at S196 (S192 in humans) reduces MCAK MT depolymerization activity and causes a corresponding conformational change in our MCAK FRET biosensor, resulting in longer fluorescence lifetimes (Ems-McClung et al., 2013). While S196 can also be phosphorylated by Aurora A kinase, S719 (S715 in humans) in the C-terminal tail of MCAK is also an Aurora A and Polo-like kinase phosphorylation site (Zhang et al., 2008; Zong et al., 2016). Furthermore, we showed previously that modification of this C-terminal domain was correlated with changes in the conformation of an MCAK FRET biosensor that do not affect MCAK activity but rather alter MCAK localization, resulting in a localized increase in MT depolymerization activity. In the present study, we did not find any major change in the distribution of MCAK with Aurora A inhibition. We did observe a decrease in fluorescence intensity at the leading and trailing edges, but whether this decrease correlates with a corresponding increase of MCAK elsewhere in the cell is not known. Finally, it is important to keep in mind that the turnover of phosphates on MCAK are incredibly fast (<5 s in mitosis) (Andrews et al., 2004) relative to the temporal resolution of the assays in the current study. Therefore, it is possible that there are multiple phosphorylation events that locally modulate MCAK conformation, and thus both its activity and its localization are affected as cells are undergoing directed migration. In this scenario, MCAK would be ideally suited to respond rapidly to changing signals in the cell to spatially modulate MT dynamics in multiple processes that require proper MT dynamics.

It is currently not clear whether the defects in centrosome position are a direct or indirect result of MCAK inhibition. In some cell types, movement of the centrosome during migration is correlated with the establishment of the leading edge. Because inhibition of MCAK enhances multiple protrusions and disrupts directional migration, it is possible that the failure to establish a single dominant protrusion consequently results in the failure to properly position the centrosome. Alternatively, overexpression of MCAK activates the release of MTs from the centrosome (Ganguly et al., 2011a,b), so perhaps inhibition of MCAK suppresses this MT release, which changes the organization of MTs in that region and prevents centrosome repositioning. In support of this idea, we also noted in earlier work that there was an accumulation of MTs in the perinuclear area after MCAK inhibition (Kline-Smith and Walczak, 2002), suggesting that MCAK may act on short MT plus ends near the centrosome or that MCAK activity may also be important at MT minus ends.

Another key component in migration is the formation and release of FAs, which provide traction for motility. We showed that MCAK knockdown increased leading-edge FA lifetime and disturbed FA turnover mainly by reducing the disassembly rate, resulting in more persistent FAs. It was shown previously that MT target endocytosis that may facilitate FA disassembly (Efimov et al., 2008; Efimov and Kaverina, 2009). MTs may provide tracks for the transport of cargoes that are involved in FA assembly and disassembly to and from the adhesion sites (Stehbens and Wittmann, 2012; Seetharaman and Etienne-Manneville, 2019). For instance, integrin-linked kinase is required for MT stabilization and caveolin transport to FAs (Wickstrom et al., 2010). Caveolin mediates endocytosis that may facilitate FA disassembly (Kiss, 2012; Stehbens and Wittmann, 2012). In addition, the MT-dependent transport and accumulation of matrix metalloproteases at FAs promote exocytosis that degrades ECM and consequently facilitates FA disassembly (Takino et al., 2006; Wiesner et al., 2010). Because MCAK does not associate directly with FAs but rather with the plus ends of MTs (Moore et al., 2005) and our unpublished results), where it controls their dynamics (Kline-Smith and Walczak, 2002; Braun et al., 2014),
we favor the idea that the increased MT stability generated by the loss of MCAK indirectly affects the transport of these other components along the MTs that interact with FAs. If we assume that MCAK knockdown does not change the level of other proteins in cells, an overall increase in the number of MTs may result in fewer cargoes per MT, which could lead to lower efficiency in disassembly of FAs. This “dilution effect” may also account for our observed decrease in the assembly rate.

Previously, it was reported that several MT plus tip tracking proteins, APC, CLASPs, and ACF7, in addition to their activity in MT remodeling, may mediate a physical link between the MT plus ends and FAs, and contribute to robust cell migration (Watanabe et al., 2004; Wen et al., 2004; Wu et al., 2008, 2011; Stehbens et al., 2014). MCAK also tracks MT plus ends (Moore et al., 2005; Honnappa et al., 2009). Plus-tip tracking proteins rely on polymerizing MTs to track MT ends; thus, the loss of MCAK activity on MT plus tips may cause a change in the composition, dynamics, and/or architecture of the protein complexes riding on growing MT tips due to increased polymerization that results in our observed FA defects.

The defects in FA dynamics after MCAK knockdown may also help explain the defects in cell movement in MCAK knockdown cells, as properly regulated FA dynamics are required for productive forward translocation of the cell body (Stehbens and Wittmann, 2012). Our analysis also showed that the overall decrease in migration after MCAK knockdown was due at least in part to the short-term movement of the cell body. If the cell body is more adherent to the substrate because of slowed FA turnover, this would result in failure of the cell body to be able to readily translocate forward. In addition, we also consistently observed “blebbing” in cells with MCAK knockdown that occurs around the edges of the adhered cell body. It has been proposed that bleb formation is driven by hydrostatic pressure generated in the cytoplasm by the contractile actomyosin cortex, which is myosin-II dependent (Paluch and Raz, 2013). MCAK regulation of MT dynamics has also been shown to be mechano-sensitive in a myosin-II dependent manner in 3D cultured HUVeC cells (D’Angelo et al., 2017), indicating cross-talk between the two pathways. Thus, it is possible that the “blebbing” phenotype we observed was caused by a change in myosin-II activity after MCAK knockdown. Perhaps this phenotype becomes more apparent in cells that are unable to generate a single leading protrusive edge due to changes in adhesion as well as changes in the contractile actomyosin cortex.

Taken together, our results support a model wherein MCAK contributes to overall cell migration at multiple levels (Figure 7). Cells require MCAK to be locally inhibited to allow for the establishment of a single protrusive front with stabilized MTs (Figure 7A). The stabilization of this front is further enhanced by the transport of Rac1 and other factors along the MTs to the leading edge of the cell. We propose that the population of MCAK, located at the plus tips of MTs, acts to ensure the dynamic turnover of these MT ends, generating a dynamic array of MTs, some of which can interact with FAs and stimulate their turnover, which will in turn allow for the cell body to translocate forward. Our data suggest that MCAK may also act on MTs near centrosomes, where it enhances proper centrosome positioning toward the leading edge of the cell. In this model, MCAK control of MT dynamics plays a critical role in a feedback loop that ensures the development of a dominant leading edge for directed cell migration (Figure 7B). This model also highlights how other key players, such as Rac1 and Aurora A, modulate MT dynamics through other pathways to provide additional levels of control for the precise regulation of cell migration.

MATERIALS AND METHODS

Cell culture and RNAi

Human retinal pigment epithelial cells immortalized with hTERT (hTERT-RPE-1; American Type Culture Collection) were cultured in DMEM or Opti-MEM (Invitrogen) supplemented with 10% fetal bovine serum, 2 mM l-glutamine, and 50 mg/ml penicillin/streptomycin at 37°C, 5% CO₂, in a humidified incubator. A stable RPE-1 cell line expressing paullin-GFP (RPE-1 PG6) was obtained from the laboratory of A. Straube (Theisen et al., 2012). HEK-293 cells were cultured in Life Technologies DMEM, high glucose, supplemented with GlutaMAX and pyruvate (Thermo Fisher), 10% fetal bovine serum (Corning), and 50 mg/ml penicillin/streptomycin at 37°C, 5% CO₂, in a humidified incubator. Aurora A inhibitor I was from Selleck Chemicals and used at 40 or 80 nM (Braun et al., 2014). Cell line identity was verified by short tandem repeat profiling.

For siRNA-mediated MCAK knockdown, 3 × 10⁵ RPE-1 cells were seeded in a six-well plate. After 24 h, cells were transfected with 10 nM RNAi oligonucleotides using Lipofectamine RNAiMax (Invitrogen) according to the manufacturer’s instructions. The following siRNAs (Dharmacon) were used: negative control-2 (5’-UGGUUAACALGUUUGUGUGA-3’), MCAK-2 RNAi (5’-GAUCCACGCAUGAAUGGUG-3’) (Hedrick et al., 2008), and a newly synthesized siRNA to the 3’ UTR of MCAK (ACUGAUGUUGUGUACUUAAAAAAA; Integrated DNA Technologies). Cells were examined at 48–60 h after knockdown depending on the specific type of experiment.

DNA constructs, transfection, and lentivirus production

Plasmids used for transfection include pEGFP-C1 (GenBank: U55763.1; Clonetech) and pcDNA3-EGFP-Rac1-Q61L (a gift from G. Bokoch; Addgene plasmid #12981) (Subauste et al., 2000). mEmerald-MCAK and mEmerald-MCAK-mCherry were cloned using Gibson assembly by fusing the cDNA of mEmerald, hMCAK, and/or mCherry into the backbone of pEGFP-C1 (Clonetech). To generate pLOVE-mEmerald-MCAK and pLOVE-mEmerald-MCAK-mCherry, mEmerald-MCAK and mEmerald-MCAK-mCherry sequences were cloned into the pENTR/D-TOPO backbone (Invitrogen) and then recombined into the pLOVE plasmid (Addgene #15948) according to the manufacturer’s instructions. For Rac1 overexpression, at 30 h post–RNAi transfection, cells were plated at ~2 × 10⁶ cells/ml on poly-L-lysine coverslips and allowed to attach for 24 h before DNA constructs were introduced into cells using Lipofectamine 3000 (Invitrogen) according to the manufacturer’s instructions. Cells were analyzed at 6 h post–DNA transfection.

Lentiviruses were produced by cotransfecting 1 µmol of pLOVE-mEmerald-MCAK or pLOVE-mEmerald-MCAK-mCherry with the packaging plasmids dRT-pMDLg/pRRE (Addgene #60488), pRSV-Rev (Addgene #12253), and pMD2.G (Addgene #12259) using Lipofectamine 3000 in HEK293T cells plated at 3.5 × 10⁶ cells/ml. Media was changed 6 h posttransfection, and media containing virus was collected at 24 and 48 h posttransfection, filtered, and stored at ~80°C in single-use aliquots.

For knockdown/rescue experiments, 2.5 × 10⁴ RPE-1 cells were seeded in a 24-well plate. After 24 h, cells were transfected with 10 nM RNAi oligonucleotides (3’ UTR) using Lipofectamine RNAiMax (Invitrogen) according to the manufacturer’s instructions. At 24 h post–RNAi transfection, the transfection media was removed and transduced with 12,500 infectious units of mEmerald-MCAK virus in DMEM complete with 0.1% polybrene. At 36 h posttransduction, a wound was generated in the center of the well with a P20 tip, and then the cells were imaged every 2 h on a BioTek Lionheart FX...
microscope for 12 h. At the end of the experiment (48 h posttransduction), the cells were trypsinized, counted, and diluted into sample buffer for Western blot analysis. All lentivirus expression levels were based on transduction conditions used in FLIM experiments that required levels of reporter expression to acquire sufficient counts for analysis (see more detail below).

**Generation of MCAK null mutant**

An MCAK-cell line was generated using CRISPR/Cas9-mediated homology-directed recombination (Ran et al., 2013). CRISPR target sites adjacent to the start codon were selected by CRISPR Design (http://crispr.mit.edu/), and guide RNAs (gRNAs) were designed and cloned into the pSpCas9n(BB)-2A-Puro (PX462) V2.0 plasmid (a gift from Feng Zhang; Addgene plasmid #62987). Homology arms containing the 5′-UTR, mEmerald, and the first exon and intron of MCAK genomic DNA were cloned into the pENTR/D-TOPO plasmid and 1 μg repair template with Lipofectamine 3000 and were selected with 5 μg/ml puromycin. Single cells were sorted, and positive cell clones were screened by PCR against the inserted genomic region and were further verified by DNA sequencing. The insertion of mEmerald before the first exon of MCAK abolished the gene product of MCAK, as determined by Western blot.

gRNA 1.
Sense oligo: 5′-AAA CAA TGG CCA TGG ACT CTG CGC C-3′
Antisense oligo: 5′-CAC CGG CGA CGA GTC CAT GGC CAT T-3′
gRNA 2.
Sense oligo: 5′-CAC CGA TCA AGA TTC ACC GCA GTA A-3′
Antisense oligo: 5′-AAA CCT ACT GCG TTG GAT CTT GAT C-3′
Homology arm 1.
F primer: 5′-CAC CGA GAA CGG GCC ATG ATG ACG ATG GCC GTT TT-3′
R primer: 5′-AAA ACC GCC ATC GTC ATC ATG GCC CGT TCT CGG TG-3′
Homology arm 2.
F primer: 5′-CTG CCC ATT CCA CTG AAA CAC AGG ATT TCT CCA A-3′
R primer: 5′-TTG GAG AAA TTC TGT GTA GCA TGG GCA G-3′

**Immunofluorescence and Western blotting**

For immunofluorescence, cells on coverslips were fixed for 20 min in 4% formaldehyde (Thermo Scientific #28908) in phosphate-buffered saline (PBS; 137 mM NaCl, 2.7 mM KCl, 10 mM Na2HPO4, 1.8 mM KH2PO4, pH 7.2) or with −20°C MeOH and permeabilized for 20 min with 1% Triton X-100. Cells were blocked in Abdil-Tx (TBS [20 mM KH2PO4, pH 7.5] with 0.1% Triton X-100, 2% bovine serum albumin, and 0.1% sodium azide) for 30 min at room temperature (RT). Cells were stained with primary antibodies for 1 h at RT or overnight at 4°C and stained with secondary antibodies and/or phallolidin-rhodamine (1:1000; Santa Cruz #362065) for 1 h at RT. DNA was stained with 10 μg/ml Hoechst for 20 min (Sigma-Aldrich). Coverslips were washed three times with TBS-Tx (TBS with 0.1% Triton X-100) between each step and mounted in ProLong Diamond (Invitrogen). The primary antibodies used were mouse anti-α-tubulin (1μg/ml; Sigma-Aldrich GTU-88), mouse DM1α (1:5000; Sigma-Aldrich T6199), or rat anti-tubulin YL1/2 (10 μg/ml, purified in house from the rat monoclonal cell line). Secondary antibodies used were DyLight488 donkey anti-mouse (1 μg/ml; Jackson ImmunoResearch #715-545-150), DyLight594 goat anti-rat (1.5 μg/ml; Jackson ImmunoResearch #112-515-175), or Alexa Fluor 594 goat anti-mouse (1 μg/ml; Invitrogen #A10239).

For Western blots, cells were collected, washed with PBS, and lysed in 2x sample buffer (0.125 M Tris, 4% SDS, 20% glycerol, 4% β-mercaptoethanol, and a trace amount of bromophenol blue, pH 6.8) at −5 × 105 to 1 × 106 cells/ml. Equal amounts of cell lysates were electrophoresed on 10% (vol/vol) SDS–PAGE gels and transferred to nitrocellulose (Schleicher & Schuell). Blots were incubated in blocking buffer (TBS with 0.1% Tween 20, 5% [wt/vol] nonfat dry milk) and probed with mouse DM1α (1:5000) and rabbit anti–hMCAK-154 (0.8 μg/ml) (Hedrick et al., 2008) diluted in Abdil-T (TBS with 0.1% Tween-20, 2% bovine serum albumin, and 0.1% sodium azide). Secondary antibodies were used at 1 μg/ml for goat anti-rabbit and sheep anti-mouse linked horseradish peroxidase (Invitrogen). Blots were developed with SuperSignal West Pico Chemiluminescent Substrate (Pierce).

**Transwell migration, wound healing, and random migration assays**

For transwell migration assays, 5 × 104 cells in serum-free DMEM were introduced to the upper chamber of an 8-μm-pore-size insert (BD Falcon), and DMEM supplemented with 10% fetal bovine serum was used as a chemoattractant in the lower chamber. Cells were allowed to migrate for 18 h before they were fixed in 4% formaldehyde in PBS for 30 min. Cells were stained with Giemsa (5% diluted in water; Sigma-Aldrich) overnight. The stained inserts were imaged with a 20× objective (0.60 NA; Plan Apo) using an EVOS FL Auto microscope (Life Technologies). The number of cells per image was quantified with a custom MATLAB code. Briefly, 12-bit images were converted to 8-bit grayscale images, local contrast adjusted, and converted to binary images. Adaptive filtering was used to remove noise, followed by a background cutoff, image fill and open, and a size cutoff of the nucleus. To separate adjacent cells, local maxima that approximately correspond to the cell nuclei were identified, transformed to local minima, and superimposed over the binary image. All source codes can be found at https://github.com/haizong-cell_morph_tracker. Three replicate inserts were assayed per condition in an experiment, and the mean of five representative fields per insert was calculated. The results of an individual experiment were averaged from the mean number of cells migrated for the three inserts, and then the assay was repeated on three biological replicate experiments. For ease of comparison, the cell number in MCAK knockdown or MCAK−/− was normalized to the control.

For wound healing assays, 70 μl of cells at 5–7 × 104 cells/ml were seeded into a Culture-Insert 2 Well (ibidi) and grown to confluency. After 24 h, the insert was removed, and images of the wound gap were acquired every 2 h over 12 h using a 20× objective (0.60 NA; Plan Apo) on the EVOS microscope. To quantify wound closure, the area (A) of the wound was measured manually using the measure function in Fiji. Alternatively, cells were seeded in individual wells of a 24-well dish with knockdown/rescue as described above or allowed to grow to confluency and then treated with control dimethyl sulfoxide or Aurora A inhibitor. Cells were wounded with a P-20 tip, rinsed two times with complete media, and then imaged every hour on a BioTek Lionheart FX microscope. The area of the wound was measured with a custom-written Cell Profiler algorithm. To compare between experiments, the wound recovery progress (%) was normalized as ΔA/A0 for each time point. To calculate the velocity of the wound front, the width of the wound was calculated by subtracting the value of the wound front from the value of the wound width, and then dividing the result by the time interval.
dividing the wound area by the height of the wound (image height). The wound width as a function of time was fitted with linear regression. The wound front velocity was determined as the slope of the linear fit divided by two.

For the random migration assay, at 30 h post--RNAi transfection, cells were plated at \(-2 \times 10^6/\text{ml}\) on poly-l-lysine–coated MatTek dishes and allowed to adhere for 18–24 h. The culture medium was supplemented with 20 mM HEPES (pH 7.2) before imaging. Time-lapse differential interference contrast (DIC) images were obtained every 5 min for 4 h using a 20x objective (0.75 NA; PlanApo) on a DeltaVision microscope (GE Healthcare) equipped with a CoolSNAP HQ charge-coupled device (CCD) camera (Photometrics) and controlled by Softworx software (Applied Precision/GE Healthcare) at 37°C in a humidified chamber. Only individual cells that did not go through mitosis or touch other cells were used for analysis. Cell migration tracks were quantified using a custom semiautomatic MATLAB code. The center of the nucleus of each individual cell over time was manually input, and the x, y coordinates of the nucleus were saved. The migration displacement was defined as the distance between the first \((t_1)\) and last \((t_2)\) time point and was calculated as the Euclidean distance \(d = \sqrt{(x_n-x_1)^2 + (y_n-y_1)^2}\). The migration velocity of individual tracked cells was determined as the total distance traveled divided by the total time as \(d/(t_2-t_1)\). The migration path \((p)\) of a cell was calculated as the sum of distance traveled between each frame, and the directional persistence was calculated as the displacement divided by the migration path \(d/p\) (Theisen et al., 2012).

**Cell morphological analysis and centrosome reorientation assay**

For cell morphology analysis, at 30 h post--RNAi transfection, cells were plated at \(-1–2 \times 10^4/\text{cells/ml}\) on poly-l-lysine–coated coverslips and allowed to adhere for 20 h before being processed for immunofluorescence or before being transfected with GFP or CARac1-GFP plasmids. For analysis, cells were segmented using a fluorescence intensity cutoff based on phalloidin-rhodamine or GFP intensity followed by a size cutoff as described above in the analysis of the Transwell assay. Cell area and eccentricity (the distance between the foci of the cell/major axis length) of the segmented cells were determined from the MATLAB function regionprops. Alternatively, cells were plated on poly-l-lysine–coated coverslips and allowed to adhere for 16–18 h. Cells were treated with Aurora A inhibitor either at the time of plating or for the last 4 h of adherence and then processed for immunofluorescence. For Aurora A inhibition experiments, cell area and eccentricity were determined via a custom-written Cell Profiler algorithm in which each image was opened sequentially, and the cell area was determined by a user-drawn freehand shape around the cell. Area measurements were exported to Microsoft Excel, and the graphs were assembled and statistics performed with GraphPad Prism.

For the centrosome reorientation assay, \(1.35 \times 10^5\) RPE-1 cells were plated in a six-well plate containing poly-l-lysine–coated coverslips and were grown for 24 h for full attachment. Cells were transfected for knockdown as described above. At 48 h post--RNAi when cells were grown to full confluency, a scratch wound was introduced to the cells with a 20 μl micropipette tip. Cells were allowed to migrate for 2 h before being processed for immunofluorescence as described above for DNA, centrosomes, and MTs. Single-plane images were collected with a 40x objective (1.0 NA; PlanApo) attached to a Nikon Eclipse 90i microscope equipped with a CoolSnap HQ CCD camera (Photometrics) and controlled by Metamorph (Molecular Devices). Images were processed in Fiji for illustration and the measurement of the wound edge and center of mass for the centrosomes and nuclei.

The angle \(\alpha\) between the centrosome and nucleus was determined in Excel using basic geometric principles (Figure 3E) and plotted in Prism as the median and interquartile range. Briefly, a triangle between the wound edge, centrosome, and nucleus can be defined by three lines: a line from the center of mass of the centrosome to the nucleus (a), the shortest line from the center of mass of the nucleus to the wound edge (b), which forms a right angle to the wound edge, and a line between the wound edge and centrosome (c). To calculate the angle \(\alpha\) in each cell, a line parallel to the wound edge was drawn and the angle and the x, y coordinates of the center of the line recorded. Then the x, y coordinates of the centroid of the centrosome and nucleus of each cell at the leading edge were determined by drawing an ellipse around the centrosome and then the nucleus. The lengths of lines a, b, and c were determined in Excel using geometry and applied to the law of cosines to determine the angle in radians \((\theta)\) and converted to degrees for angle \(\alpha\): \[\arccos\left(\frac{a^2 + b^2 - c^2}{2ab}\right) = \theta\]

**FA turnover assay and analysis in fixed cells**

For analysis of FAs in fixed cells, RPE-1 PG6 cells were plated at \(1 \times 10^6/\text{cells/ml}\) onto poly-l-lysine coverslips in a six-well plate 24 h before RNAi transfection. At 48 h post--RNAi transfection, a P20 Pipetman tip was used to create wound edges in the monolayer. After 2 h, cells were fixed with 4% formaldehyde in PHEM (60 mM Pipes, 25 mM HEPES, 5 mM EGTA [ethylene glycol-bis(2-aminoethyl)ether]-N,N,N',N'-tetraacetic acid), 1 mM MgCl, pH 6.9) with 0.2% glutaraldehyde for 20 min. Glutaraldehyde was quenched for 10 min with freshly prepared 0.5 mg/ml NaBH₄ in PBS. Coverslips were then stained to visualize MTs as described above. Images of cells along the wound edge were captured with a 60x objective (1.42 NA; PlanApo) attached to a Nikon Eclipse 90i microscope equipped with a CoolSnap HQ CCD camera (Photometrics) and controlled by Metamorph (Molecular Devices). A Z-stack was used to capture planes at 0.2 μm steps across a 5 μm range. Images were deconvoluted using AutoQuant software (Media Cybernetics).

For analysis of FAs in live cells, at 32 h post--RNAi transfection, RPE-1 PG6 cells (100 μl) at \(5 \times 10^5/\text{cells/ml}\) were seeded into a Culture-Insert 2 Well that was placed on 35-mm glass bottom dishes coated with 5 μg/ml fibronectin. Cells were allowed to adhere and grown to confluency for 18–20 h. HEPES (pH 7.2) was added to a final concentration of 20 mM before imaging, and 10–20 image fields were taken along the wound edge every 3 min for 2 h with a 60x objective (1.42 NA; PlanApo) and captured on a CoolSnap HQ2 camera using a DeltaVision Personal DV microscope.

Analysis of FA turnover was performed as previously described (Stehbens and Wittmann, 2014; Kenific et al., 2016). Only FAs that underwent full turnover from appearance to disappearance during the imaging window were analyzed. Two to five FAs were randomly chosen per cell. To track FAs, regions of interest (ROIs) were manually drawn around individual FAs using the Bezier ROI tool in NIS-Elements software (Nikon). The ROI was redrawn if the FA changed size or location over time. The fluorescence intensity of each FA in the fluorescence intensity curve, a three-frame running average was calculated and plotted as a function of time. The curve was used to

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calculate the FA assembly rate constant, the disassembly rate con-
stant, and the lifetime by curve fitting using the “Solver” add-in in
Excel (Microsoft). Assembly was the first phase of the fluorescence
intensity curve when intensity continuously increased and was fitted
with a logistic function. Disassembly was the latter portion of the
intensity plot when fluorescence intensity was decreasing and was
fitted with an exponential decay function. Some MCAK knockdown
cells showed a pause or a second disassembly phase before full
disassembly occurred. In these cases, the second disassembly curve
was used for fitting to determine the disassembly rate.

FLIM
For FLIM, RPE1 cells were plated at 1.25 × 10^4 cells/ml onto poly-lysine–coated MatTek dishes 24 h before infection and then imaged
at 48 h postinfection. One h before imaging, a P20 Pipetman tip was
used to create wound edges within the cell monolayer, and the cells
were rinsed three times with media before imaging.

FLIM was performed on a Leica SP8 laser scanning confocal mi-
croscope controlled by LAS software (Leica) and equipped with a
40 × 1.2 NA HCX PL APO CS water objective (Leica) and an at-
tached PicoQuant multidimensional time-correlated single-photon
counting system. MatTek dishes containing lentiviral infected cells
were placed in a stage-top humidified incubator maintained at 37°C
and 5% CO2 (Toki Hit). For confocal imaging, 256 × 256 pixel
images were acquired at 400 MHz with a zoom factor of 2, and
eMerald was excited using a white light laser (WLL) at 70% power
and the 488 nm laser line at 40 MHz, 70% intensity, and 250% gain.
For FLIM imaging, 256 × 256 pixel images were acquired at
200 MHz, and eMerald was excited by a 488 nm pulsed WLL line
at 20 MHz, 70% intensity, and 250% gain until the signal was >1000
photon counts per pixel. The lifetimes per pixel were determined
using SynPhoTime 64 software (PicoQuant) in which an ROI around
each cell was manually made by following the contour of the cell,
the cell ROI thresholded to 50 counts, the decay data binned 2 ×
+ixels, and fitted to a triexponential tail fit decay model that gave a
χ² close to 1 (Ems-McClung et al., 2020). The average amplitude-
adjusted lifetimes, τav/amp, were then calculated per pixel by fixing
background and individual lifetimes determined by the fitting. To
calculate the mean lifetime at the leading and trailing edges for
each cell, the confocal and FLIM images were overlaid, and three 4 × 4 pixel ROIs were drawn at the leading edge of the cell and three
at the rear of the cell body using the confocal image as a positional
reference. The mean lifetimes for the leading edge or rear of the cell
were calculated from each of the three ROIs from the FLIM image.
For all lifetime experiments, the lentiviral transduction conditions
were calculated from each of the three ROIs from the FLIM image.

Image processing and statistical analysis
Images in each experiment were taken with the same imaging set-
ings, and analyses were done on unprocessed images. Images were
processed in Fiji and assembled in Illustrator (Adobe). Data were
plotted in MATLAB, Prism (GraphPad Software), and graphs assem-
bled in Illustrator. Statistical analyses were performed using Prism. To
determine whether the data came from a normal distribution, a
D’Agostino–Pearson omnibus normality test was performed, and a
p < 0.05 was considered a nonnormal distribution. A normal distribu-
tion was assumed when the sample size was small. For comparing
two data sets, a two-tailed Student’s t test was used for data
distributed normally, and a two-tailed Mann–Whitney U test was
used for nonnormal data. For comparing two groups with normal
distribution and unequal variances, Welch’s t test was used. For
multiple comparisons, an ordinary one-way analysis of variance
(ANOVA) following by Tukey post-hoc test was used for both normal
and nonnormal distributions. A p < 0.05 was considered significant
for all tests

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