Protein kinase A activity is regulated by actomyosin contractility during cell migration and is required for durotaxis

Andrew J. McKenzie\textsuperscript{a,b,\dagger}, Kathryn V. Svec\textsuperscript{a,b,\dagger}, Tamara F. Williams\textsuperscript{a,b}, and Alan K. Howe\textsuperscript{a,b,c,*}

\textsuperscript{a}Department of Pharmacology, \textsuperscript{b}University of Vermont Cancer Center, and \textsuperscript{c}Department of Molecular Physiology and Biophysics, University of Vermont, Burlington, VT 05405

\textbf{ABSTRACT} Dynamic subcellular regulation of protein kinase A (PKA) activity is important for the motile behavior of many cell types, yet the mechanisms governing PKA activity during cell migration remain largely unknown. The motility of SKOV-3 epithelial ovarian cancer (EOC) cells has been shown to be dependent both on localized PKA activity and, more recently, on mechanical reciprocity between cellular tension and extracellular matrix rigidity. Here, we investigated the possibility that PKA is regulated by mechanical signaling during migration. We find that localized PKA activity in migrating cells rapidly decreases upon inhibition of actomyosin contractility (specifically, of myosin ATPase, Rho kinase, or myosin light-chain kinase activity). Moreover, PKA activity is spatially and temporally correlated with cellular traction forces in migrating cells. Additionally, PKA is rapidly and locally activated by mechanical stretch in an actomyosin contractility-dependent manner. Finally, inhibition of PKA activity inhibits mechanically guided migration, also known as durotaxis. These observations establish PKA as a locally regulated effector of cellular mechanotransduction and as a regulator of mechanically guided cell migration.

\textbf{INTRODUCTION} Cells sense, respond to, and contribute to the mechanical properties of the extracellular matrix (ECM) by exerting actomyosin-dependent contractile force on integrin-based adhesive contacts and sensing countertension through mechanoochemical systems (Bershadsky \textit{et al.}, 2003; Janmey and McCulloch, 2007; Schwartz, 2010; Levayer and Lecuit, 2012; Schiller and Fassler, 2013; Ringer \textit{et al.}, 2017; Weinberg \textit{et al.}, 2017). Integrin engagement and clustering initiate the generation of cellular forces through actomyosin contractility (Chrzanowska-Wodnicka and Burridge, 1996; Choquet \textit{et al.}, 1997; Balaban \textit{et al.}, 2001), which in turn promotes maturation and strengthening of adhesive contacts, thereby providing countertension to the force of protrusive actin polymerization within leading-edge lamellipodia and large-scale contractility to pull the cell body in the direction of migration (Riveline \textit{et al.}, 2001; Parker \textit{et al.}, 2002; Prager-Khoutorsky \textit{et al.}, 2011; Wolfenson \textit{et al.}, 2011; Plotnikov \textit{et al.}, 2012; Plotnikov and Waterman, 2013; Roca-Cusachs \textit{et al.}, 2013; Lintz \textit{et al.}, 2017). The distribution of contractile forces within a migrating cell generates subcellular areas with varying degrees of intracellular tension, countered both by internal cytoskeletal scaffolds and by matrix adhesions (Ingber, 1997; Schwartz, 2010). Intracellular contractile forces regulate myriad aspects of cell migration (Vicente-Manzanares \textit{et al.}, 2011; Levayer and Lecuit, 2012; Plotnikov \textit{et al.}, 2012; Pasaperla \textit{et al.}, 2015; Schiffhauer and Robinson, 2017), and during the migration of many cell types, traction force tends to be highest within the leading edge, often within the lamella just behind actively protruding lamellipodia (Berether-Hahn and Luers, 1998; Dembo and Wang, 1999; Lo \textit{et al.}, 2000; Beningo \textit{et al.}, 2001; McKenzie \textit{et al.}, 2018).
dynamics (Howe, 2004, 2011), and localized activation of PKA signaling, facilitated by A-kinase anchoring proteins (AKAPs), is necessary for the motile behavior of numerous cell types (Howe et al., 2005; Lim et al., 2007, 2008; Paulucci-Holthauzen et al., 2009; Zhang et al., 2010; McKenzie et al., 2011; Tkachenko et al., 2011; Takahashi et al., 2013; Deming et al., 2015; Sinha et al., 2015). However, the mechanisms controlling PKA activation during migration remain unclear.

Previously, we showed that efficient migration of SKOV-3 human ovarian cancer cells requires localized PKA activity within the leading edge (McKenzie et al., 2011). More recently, we demonstrated that SKOV-3 cell migration is also governed by the mechanical microenvironment; specifically, that cell contractility and migration positively correlate with ECM stiffness and that directional increases in ECM tension promote SKOV-3 cell durotaxis (McKenzie et al., 2018). In the present work, we explore the possibility that localized PKA activity in migrating cells might be regulated by mechanical signaling and cell–matrix tension.

RESULTS

Protein kinase is activated at the leading edge during cell migration but not at the periphery during cell spreading

In an attempt to facilitate the investigation of the regulation of PKA within the leading edges of migrating cells, we performed live-cell Förster resonance energy transfer (FRET) microscopy using pmAKAR3, a genetically encoded PKA biosensor consisting of a sensor cassette (a PKA-specific substrate domain and a flanking phosphoamino acid–binding FHA1 domain) located between ECFP and an EYFP variant (cpV-E172), and a C-terminal CAAX box (derived from K-Ras) for targeting to the plasma membrane (Allen and Zhang, 2006). PKA-mediated phosphorylation of the pmAKAR3 substrate domain promotes intramolecular binding by the FHA1 domain, juxtaposition of the fluorophores, and increased FRET. Mutation of the PKA phosphorylation site in pmAKAR3 from Thr to Ala (pmAKAR3T34A) ablates the baseline FRET signal and renders the biosensor unresponsive to pharmacological elevation of cAMP by treatment with forskolin and IBMX (to activate adenylyl cyclase and inhibit phosphodiesterases, respectively; Supplemental Figure S1A). Importantly, it is this biosensor that has been used and characterized most extensively to describe and investigate localized PKA activity in migrating cells (Lim et al., 2008; Paulucci-Holthauzen et al., 2009; McKenzie et al., 2011; Tkachenko et al., 2011). We first used pmAKAR3 to compare the PKA activity in the leading edges of migrating SKOV-3 cells and in the peripheries of spreading SKOV-3 cells, shortly after plating onto fibronectin-coated surfaces, as this periphery is often used as a model for the migratory leading edge (e.g., Giannone et al., 2007). Consistent with prior reports (McKenzie et al., 2011), robust and dynamic PKA activity was seen within the leading edge of migrating cells (Figure 1, B and C; Supplemental Movie S1). No FRET signal was observed in cells expressing the phosphoresistant pmAKAR3T34A mutant biosensor (Supplemental Figure S1B; Supplemental Movie S2), confirming that the leading-edge signal from the wild-type sensor is indeed due to biosensor phosphorylation. In contrast to the robust activity at the leading edge, PKA activity in the peripheries of spreading cells was very low (Figure 1, A and C; Supplemental Movie S1). This was somewhat surprising, given that prior investigations have reported, using biochemical methods, some degree of activation of PKA early upon integrin engagement and during cell spreading (Whittard and Akiyama, 2001; Howe et al., 2002). It is important to reiterate, here, that the pmAKAR3 biosensor used throughout this study (and others; Lim et al., 2008; Paulucci-Holthauzen et al., 2009; McKenzie et al., 2011; Tkachenko et al., 2011) is targeted to the plasma membrane and thus reports only membrane-proximal events, and therefore might not detect other pools of PKA that might be activated during spreading. That notwithstanding, the exceedingly low PKA peripheral activity suggests that spreading cells are not a suitable experimental system for studying regulation of PKA within the leading edge.

Leading-edge protein kinase activity events are spatially and temporally distinct from sites of focal adhesion formation

Over the course of additional experimentation and optimization of conditions for visualizing localized PKA activity during migration, we noticed that activation of PKA at the leading edge was common to many modes of migration under a variety of culture conditions; for example, signaling events were seen in the presence or absence of serum, before and after serum starvation, and with and without supplemental growth factor, albeit with differences in signaling event morphology and kinetics (Supplemental Movie S3). This suggested that, rather than being instigated by soluble cues from the cell culture media, the mechanisms governing localized PKA activity during migration might be governed by something intrinsic to cells as they move.

One process common to most, if not all, modes of migration is the engagement of integrins by the ECM and the subsequent formation of adhesive complexes such as focal contacts and focal adhesions (Mostafavi-Pour et al., 2003). Importantly, engagement of integrins has also been implicated in activation of PKA (O’Connor and Mercurio, 2001; Whittard and Akiyama, 2001; Goldmann, 2002; Howe et al., 2002, 2005; Howe, 2004; Alenghat et al., 2009; Tkachenko et al., 2011). We therefore investigated whether there were any spatiotemporal links between leading-edge PKA signaling events and the formation of focal adhesion. For this, we imaged cells coexpressing pmAKAR3 and mCherry–paxillin to visualize PKA activity and focal adhesion dynamics, respectively. Paxillin was used as a marker of focal adhesions because it recruits to focal adhesions early in the maturation process and remains throughout the entirety of the adhesion lifetime (Zaidel-Bar et al., 2003). Visual assessment of numerous images suggested no close or obvious spatial correlation between leading-edge PKA events and paxillin-containing focal adhesions (Figure 2A). To further assess any spatiotemporal correlation between focal adhesion dynamics and PKA activity, leading-edge focal adhesions were tracked over time and an interrogation region of interest (ROI) 50% larger than the area of the adhesion was used to track pmAKAR3 FRET ratios under the same cellular regions (Figure 2B). Paxillin displayed full adhesion life cycles including assembly, peak, and disassembly phases (Figure 2C, red line), while PKA activity showed little, if any, variation in the same regions. Additionally, ROIs were placed around areas of dynamic leading-edge PKA activity events and used to measure corresponding mCherry–paxillin intensities (Figure 2D). Interestingly, there was no apparent covariation between the intensity of mCherry–paxillin and peak PKA activity events (Figure 2E). Similar results were seen when mCherry–FAK was used as a distinct focal adhesion marker (Supplemental Figure S2). Integrin-containing complexes are known to generate myriad signals—most notably, a local increase in tyrosine phosphorylation and phosphotyrosine-dependent protein–protein interactions (Humphries et al., 2019). Thus, to ensure that the lack of coincidence between PKA events and focal adhesion dynamics was not due to an inability to detect adhesion-associated signaling events, cells were cotransfected with mCherry–paxillin and YFP–diSH2 (Kirchner et al., 2003), a phosphotyrosine reporter comprising yellow fluorescent protein fused to two tandem Src SH2 domains.
FIGURE 1: PKA activity is increased at the leading edge of migrating cells but not at the periphery of spreading cells. SKOV-3 cells expressing pmAKAR3 were plated onto fibronectin-coated glass coverslips and imaged via FRET microscopy 10 min (A) or 4 h (B) after plating. Representative pseudocolored FRET/CFP images of spreading (A) and migrating (B) cells are shown (bar = 25 µm). FRET ratios of each image are scaled from 1.0 to 2.5 (cool to warm color). (C) A linescan analysis of the change in centroid-to-front FRET ratio was performed on the spreading (left) and migrating (right) cells from A and B. Kymographs represent the FRET ratio along the linescans shown in Supplemental Movie S1 over the course of 1 h. Images were acquired every 60 s. Kymograph scale bars represent 5 µm on the X axis and 5 min on the Y axis.

that bind phosphotyrosine. As expected, we saw strong covariation of paxillin intensity and the intensity of YFP-dSH2 during the assembly, peak, and disassembly of focal adhesions (Figure 2F). Quantification of Pearson's correlation coefficients showed a strong positive correlation between mCherry-paxillin and YFP-dSH2, but no correlation between pmAKAR3 FRET ratios and mCherry-paxillin (Figure 2G). While we cannot rule out a role for small and/or transient focal complexes that are below our threshold of detection, these results demonstrate that leading-edge PKA dynamics is not spatiotemporally correlated with the onset, maturation, or dissolution of mature focal adhesions in migrating cells. This suggests that leading-edge PKA activity is regulated through a mechanism dependent on, but downstream of and spatially removed from, integrin-mediated focal adhesion formation.

In addition to spanning the plasma membrane, integrins and their dependent adhesion complexes can both regulate and be regulated by membrane order and lipid raft dynamics in complex ways (Leitinger and Hogg, 2002; Gagnoux-Palacios et al., 2003; Del Pozo, 2004; del Pozo et al., 2004, 2005; Fabbri et al., 2005; Gaus et al., 2006; Vassilieva et al., 2008; Norambuena and Schwartz, 2011; Wang et al., 2013; Head et al., 2014; Sun et al., 2016). Importantly, PKA also has distinct functions in lipid rafts (Golub and Caroni, 2005; Ruppelt et al., 2007; Delint-Ramirez et al., 2011; Raslan and Naseem, 2015), and work using distinctively targeted PKA biosensors has demonstrated differential regulation of PKA in bulk plasma membrane as compared with lipid rafts (Depry et al., 2011). Therefore, to determine whether a spatiotemporal correlation might exist between adhesion dynamics and leading-edge PKA activity in a distinct membrane microdomain, we visualized PKA dynamics in migrating cells using the lipid raft–targeted biosensor LynAKAR4 (Depry et al., 2011). Similarly to the activity seen with pmAKAR3, dynamic PKA activity was also observed in the leading edges of LynAKAR4-expressing cells, albeit with discernibly distinct patterns and dynamics (Supplemental Figure S3A; Supplemental Movie S4). Notably, these distinct raft–associated PKA activity events also did
FIGURE 2: Leading-edge PKA activity does not correlate with focal adhesion dynamics. (A) Migrating SKOV-3 cells coexpressing pmAKAR3 and mCherry–paxillin were plated onto fibronectin-coated imaging dishes, and representative near-simultaneous paxillin (Pxn) and pseudocolored FRET/CFP (pmAKAR3) images from live-cell microscopy experiments are shown. Arrow indicates location of peak FRET/CFP signal and corresponding location on mCherry–paxillin image. The leading edge is magnified in the insets, shown with cell outlines plotted for reference (bar = 10 µm). (B) Images of a single paxillin-containing focal adhesion during assembly, peak, and disassembly (top) with overlapping pseudocolored FRET/CFP (bottom) are shown. (C) The changes in either paxillin fluorescence intensity or FRET ratios were plotted over time. (D) Images of dynamic PKA activity and the corresponding paxillin images are shown with ROI plotted as a reference. (E) The changes in either FRET ratios or paxillin fluorescence intensity were plotted over time. (F) YFP-dSH2 and paxillin intensities were tracked over time during focal adhesion assembly, peak, and disassembly. (G) Pearson’s coefficients of the covariance of paxillin with either YFP-dSH2 (Pxn/dSH2) or pmAKAR3 (Pxn/pmAKAR3) during focal adhesion assembly, peak, and disassembly are shown as mean ± SEM (Pxn/dSH2, n = 10; Pxn/pmAKAR3, n = 13; * p < 0.001 for each phase of focal adhesion lifetime).

Leading-edge protein kinase activity is regulated by actomyosin contractility

In further consideration of what might regulate PKA activity during migration, we began to consider a cellular characteristic that is dependent on integrin-mediated adhesion, intrinsic to many cells across many modes of migration, and not typically associated with cell spreading—namely, actomyosin-dependent cellular contractility (Bershadsky et al., 2003; Wakatsuki et al., 2003; Zhang et al., 2008; Wolfenson et al., 2011; Plotnikov et al., 2012). Intracellular contractile forces regulate diverse aspects of signaling and cytoskeletal dynamics during cell migration (Vicente-Manzanares et al., 2011; Levayer and Lecuit, 2012; Plotnikov et al., 2012; Pasapera et al., 2015; Schiffhauer and Robinson, 2017) and, importantly, these forces tend to be highest within the leading edge (Bereiter-Hahn and Luers, 1998; Dembo and Wang, 1999; Lo et al., 2000; Beningo et al., 2001; McKenzie et al., 2018), placing them in the correct subcellular location to affect PKA activity.

To investigate whether cellular contractility affected PKA dynamics, we imaged PKA activity in live cells before and after addition of blebbistatin, an inhibitor of the myosin II-ATPase. Upon addition of blebbistatin, localized and dynamic activity of PKA at the leading edge rapidly and significantly diminished (Figure 3, A and B; Supplementary Movie S5), suggesting that this localized activity is dependent on actomyosin contractility. Exposure of blebbistatin to blue light (≤488 nm), however, can inactivate the compound and generate cytotoxic free radicals (Kolega, 2004; Sakamoto et al., 2005), although attenuation of light intensity and rapid diffusion of “fresh” drug from the media bath into cells often allow its utility in live-cell imaging experiments using blue light excitation (Hotulainen and Lappalainen, 2006; Burnette et al., 2008; Aratyn-Schaus and Gardel, 2010; Myers et al., 2011). Nonetheless, to ensure that blebbistatin-mediated inhibition of leading-edge PKA activity was due to inhibition of contractility and not cytotoxicity, we repeated this experiment with other contractility
inhibitors. Initial attempts at imaging pmAKAR3-expressing cells before and after addition of para-aminoblebbistatin, a nontoxic blebbistatin derivative that is supposedly nonfluorescent (Varkuti et al., 2016), gave uninterpretable results due to significant yellow fluorescence of the drug inside cells—an observation confirmed in nontransfected cells (Supplemental Figure S4). We therefore employed compounds that are devoid of phototoxicity and optical artefacts and inhibit actomyosin contractility (Supplemental Figure S5) through distinct mechanisms; specifically, through inhibition of Rho-kinase (using H-1152, Fasudil/HA-1077, or Y-27632) or myosin light-chain kinase (using ML-7), both of which contribute to the phosphorylation and activation of the myosin light chain (MLC) of myosin II to promote contractility (Amano et al., 1996; Totsukawa et al., 2000). Treatment of migrating, pmAKAR3-expressing cells with each of these compounds lead to a rapid and significant decrease in leading-edge PKA activity (Figure 3C; Supplemental Movie S6; Supplemental Figure S6). Leading-edge PKA activity visualized by LynAKAR4 showed a similar dramatic reduction upon inhibition of Rho-kinase by Y-27632 (Supplemental Figure 3C), further supporting the observation that leading-edge PKA activity is dependent on actomyosin contractility.

Inhibition of actomyosin contractility is associated with discrete morphological effects, including cessation of leading-edge dynamics and eventual dissolution of focal adhesions, so it is possible that the loss of PKA is coupled to one of those events. To better assess the kinetics of loss of PKA activity upon inhibition of contractility, we generated maps of PKA activity (top) and edge velocity (bottom) within a 10-µm band along the leading edge before and after treatment with blebbistatin (Blebb). (E) SKOV-3 cells expressing mCherry–paxillin, mCherry–zyxin, or pmAKAR3 (FRET) migrating on fibronectin-coated glass dishes, or cells plated on 125 kPa fluorescent nanosphere-functionalized hydrogels (traction force) were treated with 25 µM blebbistatin (or with 0.1% vol/vol DMSO; Ctrl) at time = 0. Capturing images every 60 s, the fluorescence intensity of paxillin or zyxin within focal adhesions, PKA activity (via the pmAKAR FRET signal), or traction force was measured, normalized to values at time = 0, and plotted. The graph depicts mean values ± SEM (n_paxillin-Blebb = 211 adhesions from seven cells; n_zyxin-Blebb = 156 adhesions from five cells; n_FRET = 30 linescans from six cells; n_zyxin-Ctrl = 125 adhesions from five cells; n_FRET-Ctrl = 35 linescans from seven cells; n_traction_force = average traction from six cells). The inset shows the first 5 min of the time course at higher resolution; these data were used to calculate the one-phase half-life (t1/2) or apparent t1/2 decay values for each signal, using GraphPad Prism.

**FIGURE 3:** Actomyosin contractility regulates leading-edge velocity and PKA activity. (A) A migrating SKOV-3 cell expressing pmAKAR3 was plated on a fibronectin-coated glass-bottomed imaging dish and monitored by live-cell microscopy before and after treatment with 25 µM blebbistatin (Blebb). Representative pseudocolored FRET/CFP images (corresponding to the boxed region in Supplemental Movie S3) are shown. (B) A maximum-intensity projection of cumulative PKA activity in 10 images (2 min apart) before and after treatment with blebbistatin. (C) Representative images of pmAKAR3-expressing SKOV-3 cells migrating on fibronectin-coated dishes before (0 min) and 30 min after treatment with 0.1% vol/vol dimethyl sulfoxide (DMSO), 1 µM H-1152, or 10 µM fasudil. The leading edge is magnified in the insets (scale bar = 10 µm). (D) QuimP11 software (Bosgraaf and Van Haastert, 2010) was used to generate maps of PKA activity (top) and edge velocity (bottom) within a 10-µm band along the leading edge before and after treatment with blebbistatin (Blebb). (E) SKOV-3 cells expressing mCherry–paxillin, mCherry–zyxin, or pmAKAR3 (FRET) migrating on fibronectin-coated glass dishes, or cells plated on 125 kPa fluorescent nanosphere-functionalized hydrogels (traction force) were treated with 25 µM blebbistatin (or with 0.1% vol/vol DMSO; Ctrl) at time = 0. Capturing images every 60 s, the fluorescence intensity of paxillin or zyxin within focal adhesions, PKA activity (via the pmAKAR FRET signal), or traction force was measured, normalized to values at time = 0, and plotted. The graph depicts mean values ± SEM (n_paxillin-Blebb = 211 adhesions from seven cells; n_zyxin-Blebb = 156 adhesions from five cells; n_FRET = 30 linescans from six cells; n_zyxin-Ctrl = 125 adhesions from five cells; n_FRET-Ctrl = 35 linescans from seven cells; n_traction_force = average traction from six cells). The inset shows the first 5 min of the time course at higher resolution; these data were used to calculate the one-phase half-life (t1/2) or apparent t1/2 decay values for each signal, using GraphPad Prism.
contractility would be expected to disrupt that signaling. Specifically, if PKA activity were dependent on signaling from intact focal adhesions, we would predict that disassembly of focal adhesions would precede loss of PKA activity. Thus, we assessed the kinetics of inhibition of PKA activity relative to focal adhesion disassembly by measuring the rate of blebbistatin-induced loss of fluorescent signal intensity of two focal adhesion markers: zyxin, a mechanosensitive protein that leaves focal adhesion rapidly upon loss of contractility, and paxillin, which leaves “relaxed” focal adhesions much more slowly and with apparent zero-order kinetics (Zaidel-Bar et al., 2003; Wolfenson et al., 2011; Lavelin et al., 2013). As expected, paxillin intensity within focal adhesions decreased slowly but steadily after blebbistatin treatment (Figure 3E), with an apparent half-life of 52.69 ± 3.35 min, while zyxin intensity decreased rapidly and exponentially, with a half-life of 7.63 ± 0.66 min (Figure 3E; Supplemental Movie S7), consistent with prior observations of the higher dependence of zyxin on mechanical forces for residence within focal adhesions (Lele et al., 2006; Hirata et al., 2008; Pasapera et al., 2010; Wolfenson et al., 2011; Lavelin et al., 2013). Interestingly, PKA activity decreased even more rapidly after blebbistatin treatment than focal zyxin intensity (Figure 3E), with a half-life of ∼50 s (0.83 ± 0.09 min). These observations are consistent with our observation of the lack of spatiotemporal correlation between focal adhesion dynamics and peripheral PKA signaling events and suggest a closer relative coupling of PKA to the contractile state of the cell. With this in mind, we then determined the kinetics of loss of cellular traction force after blebbistatin treatment using traction force microscopy on cells adhering to fibronectin-coated polyacrylamide hydrogels functionalized with fluorescent nanospheres (McKenzie et al., 2018). As expected, cellular traction force decreased very rapidly after addition of blebbistatin, with a $t_\frac{1}{2}$ of 0.37 ± 0.05 min (Figure 3E and Supplemental Movie S8). While this $t_\frac{1}{2}$ value is likely to be erroneously high, given that the rate of image acquisition for these experiments (1 frame/min) results in an image interval that far exceeds the apparent $t_\frac{1}{2}$, it is consistent with published reports (Pasapera et al., 2010; Wolfenson et al., 2011; Lavelin et al., 2013). More importantly, this decrease was far more rapid than the loss of paxillin or even zyxin from focal adhesions but only slightly more rapid than loss of PKA activity. Collectively, these data demonstrate that, during cell migration, regulation of PKA activity within the leading edge is kinetically coupled to and dependent on actomyosin contractility.

**Spatial distribution of cellular traction forces and protein kinase activity during cell migration**

If leading-edge PKA were regulated by actomyosin contractility, then one might expect PKA activity and cellular contractile forces to be coincident in migrating cells. To investigate this correlation directly, SKOV-3 cells expressing pmAKAR3 and migrating on nanosphere-functionalized hydrogels were analyzed by simultaneous imaging of PKA to the contractile state of the cell. With this in mind, we then determined the kinetics of loss of cellular traction force after blebbistatin treatment using traction force microscopy on cells adhering to fibronectin-coated polyacrylamide hydrogels functionalized with fluorescent nanospheres (McKenzie et al., 2018). As expected, cellular traction force decreased very rapidly after addition of blebbistatin, with a $t_\frac{1}{2}$ of 0.37 ± 0.05 min (Figure 3E and Supplemental Movie S8). While this $t_\frac{1}{2}$ value is likely to be erroneously high, given that the rate of image acquisition for these experiments (1 frame/min) results in an image interval that far exceeds the apparent $t_\frac{1}{2}$, it is consistent with published reports (Pasapera et al., 2010; Wolfenson et al., 2011; Lavelin et al., 2013). More importantly, this decrease was far more rapid than the loss of paxillin or even zyxin from focal adhesions but only slightly more rapid than loss of PKA activity. Collectively, these data demonstrate that, during cell migration, regulation of PKA activity within the leading edge is kinetically coupled to and dependent on actomyosin contractility.

**Protein kinase is locally activated by acute mechanical stretch in an actomyosin-dependent manner**

Given the demonstrated requirement of cellular tension for localized activation of PKA during migration, we wondered whether PKA might be locally activated by acute increases in cellular tension. Recently, it was shown that cellular mechanosensing is mediated not only by substrate rigidity but also by substrate deformation strain energy (Panzetta et al., 2019). Thus, to test acute mechanical activation of PKA, cells expressing pmAKAR3 were plated onto fibronectin-coated hydrogels, and the hydrogel under individual cells was stretched (perpendicular to the axis of cell migration) with a microneedle (Supplemental Figure S5; Figure S5A; Svec et al., 2019). This directional stretch produces a linear, inhomogeneous strain field (Supplemental Figure S8, A and B) between the cell and the probe. If one simplifies the elastic hydrogel to a series of equivalent springs, Hooke’s Law ($F \propto k \times \Delta x$) dictates that a pulling force ($F_R$) applied to a spring results in a displacement ($\Delta x$) that is proportional to the spring’s stiffness ($k$). Thus, the magnitude of the restoring force (i.e., the force required to restore the stretched spring to its original length) is proportional to how far the spring is stretched from its original length; that is, $F_R = -k \times \Delta x$. Because, in a locally stretched gel, the displacement increases with the proximity to the pulled microprobe (e.g., $\Delta x_1 < \Delta x_2 < \Delta x_3$; Supplemental Figure S5, A and B), so then the restoring force increases with the proximity to the pulled microprobe ($F_{R1} = -k \times \Delta x_1 < F_{R2} = -k \times \Delta x_2 < F_{R3} = -k \times \Delta x_3$; Supplemental Figure 8C). At the cellular level, this increased force or countertension perceived as a stiffer substrate. Specifically, as a cell probes this gradient (e.g., from a to b in Supplemental Figure S8A), either a “constant” cell-generated contractile force would produce less and less gel movement, or the cell would have to exert higher force in order to move the gel the same distance. In other words, because the restoring force increases in the direction of the pull, the apparent rigidity—as perceived by the cell—increases. The reader is referred to an elegant description of this in the original report of durotaxis (Lo et al., 2000).

Application of directional stretch revealed a rapid (i.e., within 20 s), robust, and localized increase in PKA activity in the direction of stretch in both pmAKAR3-expressing cells (Figure 5, B–D; Supplemental Movie S9) and LynAKAR4-expressing cells (Supplemental Figure 3D), but not in cells expressing the phosphoreistant
pmAKAR3 biosensor (Supplemental Figure S9). Indeed, acute stretch appeared to “reorient” leading-edge PKA activity, as the increased activity seen proximal to stretch at the leading edge was often accompanied by decreased activity in other areas of the leading edge (Figure 5, C–E; Supplemental Movie S9). This activation was completely inhibited in cells coexpressing mCherry fused to the PKA-inhibitor protein (mCh-PKI; McKenzie et al., 2011; Figure 5F), confirming the PKA specificity of the response. Importantly, acute mechanical activation of PKA was also completely inhibited in cells treated with blebbistatin before stretch (Figure 5F). These results show that acute mechanical stimulation can activate PKA in a manner that requires actomyosin contractility.

Protein kinase activity is required for durotaxis in SKOV-3 cells

Previously, we have shown that efficient migration in SKOV-3 cells is dependent upon PKA activity (McKenzie et al., 2011). More recently, we showed that the migration of these cells is strongly influenced by the mechanical microenvironment and that these cells exhibit durotaxis, or mechanically guided migration toward regions of increased ECM rigidity and/or cell–matrix tension (McKenzie et al., 2018). Given the current observations connecting cellular tension and stretch to activation of PKA, we investigated whether the durotactic migration of SKOV-3 might be similarly dependent on PKA activity. To this end, control cells or cells expressing mCh-PKI were cultured to migrate on fibronectin-coated hydrogels, subject to directional stretch as described above, and monitored for durotactic response (Svec et al., 2019). While control cells exhibited robust durotaxis in response to acute directional stretch, inhibition of PKA activity dramatically decreased durotactic efficiency (Figure 6, A–C; Supplemental Movie S10). Taken together, these observations demonstrate that PKA activity is mechanically regulated during cell migration, is activated upon acute mechanical cell stretch, and is required for durotaxis.

DISCUSSION

The mechanosensitivity of cAMP/PKA signaling is well supported by the literature. For example, the cAMP cascade was the first mechanosensitive signaling cascade ever to be described (Rodan et al., 1975). Also, the levels of cAMP and PKA activity have been shown to vary as a function of the mechanical tension of fibroblasts embedded in collagen gels (He and Grinnell, 1994). Furthermore, direct application of mechanical force through integrin-mediated adhesive contacts rapidly activates cAMP and PKA signaling and PKA-dependent transcription (Meyer et al., 2000; Goldmann, 2002; Alenghat et al., 2009). Finally, mechanical signals from fluid flow activate PKA in an ECM-specific manner (Funk et al., 2010). While these and other studies firmly establish PKA as a mechanoresponsive target, it is important to note that they all assessed global rather than subcellular activation of cAMP/PKA and did not follow PKA dynamics in migrating cells.
Here, we demonstrate a link between cellular tension and regulation of subcellular PKA activity during cell migration. We show that leading-edge PKA activity correlates with the spatial distribution of cellular traction forces and that disruption of actomyosin contractility uncouples the spatial correlation between cellular forces and PKA activity. We establish that leading-edge PKA activity is regulated by cellular tension and show that locally applied mechanical forces elicit localized increases in PKA activity. Furthermore, we demonstrate that PKA activity is required for the durotactic response to directional mechanical cues, thereby expanding this enzyme’s well-established role as a regulator of other modes of cell migration (Howe et al., 2005; Lim et al., 2008; Paulucci-Holthauzen et al., 2009; McKenzie et al., 2011; Tkachenko et al., 2011). An intriguing recent report studying cells migrating from open to confined two-dimensional spaces showed that PKA activity was down-regulated during this transition in a manner dependent on Piezo1-mediated Ca\(^{2+}\) influx and, importantly, that leading-edge PKA activity appeared to increase upon treatment of cells with blebbistatin (Hung et al., 2016). These latter observations are in direct contrast to the results reported here, which demonstrate a decrease in PKA activity upon treatment not only with blebbistatin, but also with other “upstream” inhibitors of actomyosin contractility (Y-27632, H-1152, fasudil/HA1077, and ML7; Figure 3). The reasons for this disparity are currently unknown but may include differences in cell lines and culture conditions. The prior study saw increased PKA activity after blebbistatin treatment (in both confined and unconfined cells) using CHO-K1 cells and a derivative line expressing the \(\alpha_4\) integrin (CHO-\(\alpha_4\)WT), adhering to surfaces coated with 20 \(\mu\)g/ml

**FIGURE 5:** PKA is activated upon acute mechanical stretch and is required for SKOV-3 cell durotaxis. (A) Schematic of technique using a glass microneedle to impart acute directional stretch on individual cells by deformation of the underlying hydrogel (see text for details). (B) An SKOV-3 cell expressing pmAKAR3, plated on a hydrogel coated with fibronectin for 4 h and then imaged by FRET microscopy, before and 30 s after application of mechanical stretch by a microneedle in the direction indicated by the arrow. Warmer colors correspond to higher PKA activity as assessed by FRET ratio. (C) Time course of FRET ratio indicating PKA activity in an SKOV-3 cell expressing pmAKAR3 before and after application of mechanical stretch. Immediately before the 2:00 min mark, stretch was applied directly to the right of the panel. Deformation of the cell is highlighted by the outline of the cell before the stretch overlaid at 2:00 min. (D) Line scan analyses in the direction of (a) and orthogonal to the axis of stretch (b) show a unique increase in PKA activity along the axis of stretch after stimulation. (E) FRET ratio images were sectioned into quadrants a through d, depicted by dotted lines, for quantification of directional response. The change in PKA activity before and after stretch (\(\Delta\)FRET) of SKOV-3 cells expressing pmAKAR3 was calculated in quadrants proximal (quadrant a), orthogonal (quadrants b and d), or contralateral (quadrant c) to the stretch (mean ± SEM; n = 6). (F) Change in PKA activity before and after stretch (\(\Delta\)FRET) in quadrant a of SKOV-3 cells coexpressing pmAKAR3 and mCh-PKI (shown), coexpressing pmAKAR3 and mCherry, or SKOV-3 cells expressing only pmAKAR3 but pretreated with 25 \(\mu\)M blebbistatin (Blebb) for 10 min (mean ± SEM; n = 9 or 5 cells for control or mCh-PKI cells, respectively; * \(p < 0.005\) [Student’s t test]).
fibronectin. These are notable, yet modest differences compared with those in the current work, which uses SKOV-3 cells adhering to fibronectin at 10 μg/ml.

While it will be important to keep these differences in mind for future studies, it is perhaps more important to appreciate that, together, the two reports firmly establish PKA as a target for mechanotransduction during migration. Moreover, given the importance of mechanotransduction in regulating cell adhesion and motility (Schwartz and DeSimone, 2008; Roca-Cusachs et al., 2013; Schiller and Fassler, 2013) and the long and growing list of adhesion-related and cytoskeletal targets for PKA (Howe and Juliano, 2000; Howe et al., 2002, 2005; Howe, 2004, 2011; Lim et al., 2007; Tkachenko et al., 2011; Yeo et al., 2011; Takahashi et al., 2013; Ithychanda et al., 2015; Nagy et al., 2015; Robertson et al., 2015; Chavez-Vargas et al., 2016; Gau et al., 2019; Tonucci et al., 2019), the demonstration that PKA activity can be dynamically and locally regulated by cell tension provides an important new axis of regulation. Though the current work establishes the connection between leading-edge PKA activity and actomyosin contractility, the exact molecular mechanism coupling cellular tension to localized PKA activity remains to be elucidated. A plausible hypothesis is that canonical activators of the cAMP/PKA pathway (G-protein coupled receptors, adenylyl cyclases, or phosphodiesterases) are locally and mechanically regulated during cell migration. Importantly, GPCRs are well-established mediators of mechanotransduction (Storch et al., 2012), and experiments imparting tension across fibronectin-coated scaffolding structures (e.g., actin microfilaments, focal adhesions) are principal mediators of mechanical signaling. However, the relative kinetics of PKA inactivation and focal adhesion disassembly suggests that PKA is regulated by a tension-dependent aspect/characteristic of intact focal adhesions (e.g., actin microfilaments, focal adhesions) are principal mediators of mechanical signaling. However, the relative kinetics of PKA inactivation and focal adhesion disassembly suggests that PKA is regulated by a tension-dependent aspect/characteristic of intact focal adhesions (e.g.,}

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**FIGURE 6:** PKA is required for SKOV-3 cell durotaxis. (A) An SKOV-3 cell plated onto a 25-kPa fibronectin-coated hydrogel for 4 h was monitored by live-cell phase-contrast microscopy before and after durotactic stimulation (in the direction indicated by the arrow); images captured 1 min before (Pre) application of stretch, one minute after application (Stretch), and every 20 min thereafter are shown. (B) Control cells or cells expressing mCherry–PKI (PKI) were cultured and stimulated to invoke durotaxis as described for panel A. Protrusion/retraction analysis maps (PRAMs) were generated from manually thresholded and outlined phase-contrast images taken 1 min before and 1 min after stretch (Stretch) and 1 min and 45 min after stretch (45°) to identify regions of protrusion, retraction, and overlap (green, red, and black, respectively). (C) Durotactic efficiency (see Materials and Methods for details) was calculated for control and mCherry–PKI-expressing cells (n = 8 for each condition; *p < 0.01).
Scott et al., 2013) and the myriad known and potential substrates for PKA present in various cellular regions and structures involved in cell migration (discussed above), it would be of considerable interest (and complexity) to evaluate PKA activity using a series of biosensors targeted to distinct migration-associated domains or structures to identify structure-specific dynamics and substrates, and thus more fully dissect the contributions of PKA to cell motility.

**MATERIALS AND METHODS**

Reagents and cell culture

Human epithelial ovarian cancer (SKOV-3) cells were purchased from the American Type Culture Collection, authenticated at the UVM Advanced Genomic Technology Core, and maintained in a humidified incubator at 37°C containing 5% CO2 in DMEM supplemented with 10% fetal bovine serum. All cell lines in the laboratory were checked for mycoplasma monthly by DAPI (4’6-diamidino-2-phenylindole) staining and every 4–6 mo using a commercial kit (MycoAlert, Lonza). Cells were trypsinized and split 1:5 every 3–4 d to avoid reaching confluence. Cells were transfected using Fugene6 (Promega) according to the manufacturer’s protocol. In brief, cells were plated into 35-mm dishes at 60–70% confluence the day before transfection so that they were 75–85% confluent at the time of transfection. Fugene6 and Opti-MEM were warmed to room temperature (RT) and 6 μl of Fugene6 was diluted into 100 μl of Opti-MEM, vortexed, and incubated for 5 min at RT. A total of 1.5 μg of plasmid DNA was added to the diluted Fugene6 at 1:4 ratio (1.5 μg DNA/6 μl Fugene6), vortexed, and incubated for 15 min at RT. The transfection solution was then added dropwise to cells and images were acquired 48 h posttransfection.

Blebbistatin, fasudil, H-1152, and ML7 were purchased from Tocris. Acrylamide and N,N-methylenebisacrylamide were purchased from National Diagnostics. Tetramethylmethylenediamine (TEMED), ammonium persulfate (APS), and bovine serum albumin (BSA), along with other sundry chemicals, were purchased from Sigma (St. Louis, MO). Plasmids used in this work included PKA biosensor pmAKAR3 (Allen and Zhang, 2006) from Jin Zhang (Johns Hopkins University); pYFP-dSH2 from Benny Geiger (Weizmann Institute of Science); and pmCherry-FAK and pRFP-zyxin from A. J. McKenzie, K. V. Svec, et al., 2013; and the myriad known and potential substrates for PKA present in various cellular regions and structures involved in cell migration (discussed above), it would be of considerable interest (and complexity) to evaluate PKA activity using a series of biosensors targeted to distinct migration-associated domains or structures to identify structure-specific dynamics and substrates, and thus more fully dissect the contributions of PKA to cell motility.

**Fabrication of polyacrylamide hydrogels**

Acrylamide hydrogels with a Young’s elastic modulus of ~25 kPa were fabricated essentially as described previously (McKenzie et al., 2019). Briefly, cleaned 25 mm–diameter round glass coverslips were derivatized with the heterobifunctional cross-linker sulfo-SANPAH as described previously (Tse and Engler, 2010; McKenzie et al., Svec et al., 2019). For routine studies, activated gels were functionalized with 20 μg/ml fibronectin at 37°C for 45 min. For traction force microscopy studies, 0.2 μm red fluorescent carboxy-modified latex microspheres (Invitrogen, F8810) were conjugated to the gel surface by incubating a sonicated suspension of the beads (1:200 in 50 mM HEPES, pH 8.5) on the gels for 30 min. The gels were rinsed three times with 50 mM HEPES (pH 8.5) to remove all unattached beads and then incubated with 20 μg/ml fibronectin (diluted in 50 mM HEPES, pH 8.5) at 37°C for 45 min. The gels were postfixed with 0.5% glutaraldehyde for 1 h at RT and quenched in NaBH4 before cells were plated in complete media. Coated gels were washed 3 × 5 min in PBS and either used immediately or stored at 4°C for up to 1 wk. In some experiments, hydrogels were cast in a similar manner directly onto the surface of glass-bottomed imaging dishes (Delta T; Biophtechs) instead of 25-mm coverslips.

**Live cell imaging**

Cells transfected with plasmids encoding pmAKAR3 were cultured overnight in serum-free DMEM + 40 ng/ml epidermal growth factor (EGF), trypsinized, soybean trypsin inhibitor added, pelleted, and resuspended in DMEM 1% BSA + 25 or 40 ng/ml EGF. These cells were plated on fibronectin-coated coverslips and incubated for ~4 h before imaging at low density to induce migration. Cells plated on hydrogels for durotaxis were incubated overnight in complete media then rinsed twice in modified Ringer’s buffer without phosphate (10 mM HEPES; 10 mM glucose; 155 mM NaCl; 5 mM KCl; 2 mM CaCl2; 1 mM MgCl2). All cells were reseeded modified Ringer’s buffer supplemented with 25 or 40 ng/ml EGF for imaging. Coverslips were mounted in a chamber (Attofluor; ThermoFisher) before imaging. Culture temperature was maintained at 35–37°C with hot air (ASI 400 Air Stream; Nevtek). Cultures of hydrogels cast in imaging dishes were mounted, warmed, and imaged in a suitable temperature controller (Delta T; Biophtechs).

**Durotaxis assay**

Cells were seeded on fibronectin-coated gels and maintained on the microscope as above. Cells were manipulated with a glass microneedle as described previously (Wang et al., 2001; McKenzie et al., 2018; Svec et al., 2019). Briefly, micropipettes were fashioned from borosilicate glass capillaries (18150-4 or TW150-4; World Precision Instruments) on a two-stage pipette puller (Pul-2; World Precision Instruments). A Narishige MF-900 microforce was used to form the micropipette tip into a hooked probe with a rounded end to engage the polyacrylamide hydrogels without tearing. The probe was mounted on a micromanipulator (Leitz or Narishige) and lowered onto the gel surface ~20 μm away from a cell and pulled 20 μm in a direction orthogonal to the cell’s long axis. Quantification of response to stretch was calculated using custom ImageJ Protrusion-Retrac tion Analysis Mapping (PRAM) macros designed to calculate the percentage of cell area protruding, retracting, and overlapping between any two given frames of time-lapse images (Deming et al., 2015). A positive durotactic response was defined as a >50% increase in the protrusion index (protrusion area/ [protrusion area + overlap area]) in the direction of stretch over 45 min, and durotactic efficiency was defined as the number of cells showing a durotactic response divided by the number of cells pulled.

**Förster resonance energy transfer imaging and analysis**

The FRET-based PKA activity biosensor pmAKAR3, consisting of a sensor cassette (i.e., a PKA-specific substrate domain and a flanking
phosphoamino acid-binding FHA1 (forkhead-associated domain-1) located between ECFP and an EYFP variant (circularly permuted Venus csV-E172; Allen and Zhang, 2006), and a C-terminal CAAX box (derived from K-Ras) for targeting to the plasma membrane, was imaged in SKOV-3 cells as previously described (McKenzie et al., 2011). Briefly, 48 h after transfection, cells were rinsed twice and maintained in a HEPES-buffered saline solution containing (in mM) 134 NaCl, 5.4 KCl, 1.0 MgSO₄, 1.8 CaCl₂, 20 HEPES, and 5 d-glucose (pH 7.4) without serum, unless otherwise specified. Cells were imaged on a Nikon Eclipse TE-2000E inverted microscope with a 60×/1.4NA Plan Apo oil-immersion objective lens using the appropriate fluorophore-specific filters (Chroma Technology, Rockingham, VT) and an Andor Clara charged coupled device camera (Andor Technologies, South Windsor, CT) controlled by Elements (Nikon) software. CFP, YFP, and FRET images were acquired with (400–700)-ms exposures and 2×2 binning for each acquisition at 60-s intervals, unless otherwise noted in the figure legends. Images in each channel were subjected to background subtraction, and FRET ratios were calculated using either the Biosensors FRET ImageJ plug-in (Hodgson et al., 2010) or a slightly modified mathematical protocol as described elsewhere (Broussard et al., 2013). Pseudocolor images were generated using a custom-written ImageJ look-up table.

Cell spreading and migration assays
To monitor cell spreading, cells were prepared and cultured as previously described (Howe et al., 2002). Briefly, cells were serum-starved overnight, trypsinized, quenched with 1 mg/ml soybean trypsin inhibitor, washed via centrifugation (50 × g for 5 min), resuspended in DMEM 1% BSA, and rocked for 1 h before being plated on fibronectin-coated (10 µg/ml) glass-bottomed imaging dishes. The cells were allowed to settle to the bottom of the dish for 10 min at 4°C before imaging as described below. Similar conditions were used to monitor migrating cells, with the exception that the cells were allowed to adhere, spread, and begin migrating for 4 h at 37°C before imaging. Cells were imaged in Ringer’s buffer.

Correlating edge velocity and protein kinase activity
Corrected FRET ratio time-lapse movies were fed to the Quantitative Imaging of Membrane Proteins (QuimP11) package (http://go.warwick.ac.uk/bretscher/quipm) software, which analyzed edge dynamics and calculated edge velocity. Additionally, the software generated two-dimensional morphodynamic plots of edge velocities along the cell edge over time and computed autocorrelation coefficients of edge dynamics. Once edge dynamics was analyzed, the corrected FRET ratio images were analyzed by QuimP11 to sample the FRET ratios within 10 µm of the cell edge. The software generated two-dimensional heat maps of PKA activity along the cell edge over time and calculated cross-correlation coefficients at different time lags to determine when peak PKA activity events were occurring in relation to peak protrusion events. The ImageJ plug-in, QuimP11, was also used to display time-coded depth stacks to depict cell movement over time in a single image.

Focal adhesion analysis
Cells were transfected with plasmids encoding focal adhesion markers (mCherry–paxillin, mCherry–FAK, or RFP–zyxin) and either pmAKAR3 or dSH2–YFP to visualize focal adhesions and PKA activity or tyrosine phosphorylation. Images were acquired at 60-s intervals. Focal-adhesion pixel intensities and lifetimes were calculated by manually thresholding adhesions and measuring pixel intensities, adhesion assembly and disassembly rates, and lifetimes of individual adhesions in time-lapse movies. To quantify the spatiotemporal correlation between focal adhesion dynamics and PKA activity, leading-edge focal adhesions were tracked over time and an interrogation region of interest (ROI) that was 50% larger than the area of the adhesion was used to track pmAKAR3 FRET ratios in the same cellular regions. Pearson’s correlation coefficients were generated using the Intensity Correlation Analysis ImageJ plug-in, and average values are represented as mean ± SEM. LynAKAR4 FRET and mCherry–Paxillin signals were analyzed by overlaying the thresholded peak signals (≥85% of maximum).

Traction force microscopy
TFM was performed essentially as described previously (McKenzie et al., 2018). Briefly, cells were plated on polyacrylamide gels that were surface-conjugated with 0.2-µm red fluorescent latex microspheres as described above. Cells were adhered to the hydrogels overnight in complete media and were washed twice and maintained in HEPES-buffered saline as described above. Coverslips were mounted in imaging chambers as above and fluorescent bead images were captured through a 20× Plan Apo objective on a Nikon Eclipse TE-2000E inverted microscope as described above. Bead images were acquired before and after cells were cleared by the addition of trypsin/EDTA (0.5%). Cell outlines were generated using either the YFP image from cells expressing pmAKAR3 or a transmitted light image in cases where cells were not transfected. Bead images were registered to correct for any stage drift; then the movement of individual microspheres between image pairs was calculated using particle image velocimetry (PIV) and the Young’s elastic modulus of the polyacrylamide hydrogels (25 kPa) was used to calculate traction forces using Fourier transform traction cytometry (FTTC; Marinovich et al., 2012; Tseng et al., 2012). The mean traction force within the cell was used to generate the average cellular traction and the mean of the maximum traction forces was used to generate the average maximum traction force generation.

Traction force microscopy/Förster resonance energy transfer correlation analysis
To correlate cellular traction forces and PKA activity, both readings were captured simultaneously and analyzed independently. Once heat maps of both PKA activity and traction forces were generated, standard image correlation analysis was performed. This analysis was made possible because the two signals are rendered as 8-bit gray-scale images and higher pixel intensity corresponds with either higher PKA activity or higher traction forces. Lookup tables are assigned to the images after analysis for ease of interpreting biosensor and TFM data. Mander’s correlation coefficients and intensity correlation quotients were generated using the Intensity Correlation Analysis ImageJ plug-in, and average values are represented as mean ± SEM. Intensity correlation quotient (ICQ) analysis has been described in detail elsewhere. In brief, ICQ reflects the ratio of the number of positive (Ai-aI)(Bi-bI) values to the total number of pixels in the region of interest, where a and b are the means of each signal intensity’s values Ai and Bi. ICQ values from -0.05 to +0.05 indicate random (noncovariant) signals, 0.05–0.1 indicate weak covariance, and >0.1 indicates strong covariance.

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