The focal adhesion scaffold protein Hic-5 regulates vimentin organization in fibroblasts

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ABSTRACT Focal adhesion (FA)-stimulated reorganization of the F-actin cytoskeleton regulates cellular size, shape, and mechanical properties. However, FA cross-talk with the intermediate filament cytoskeleton is poorly understood. Genetic ablation of the FA-associated scaffold protein Hic-5 in mouse cancer-associated fibroblasts (CAFs) promoted a dramatic collapse of the vimentin network, which was rescued following EGFP-Hic-5 expression. Vimentin collapse correlated with a loss of detergent-soluble vimentin filament precursors and decreased vimentin ST2/SB2 phosphorylation. Additionally, fluorescence recovery after photobleaching analysis indicated impaired vimentin dynamics. Microtubule (MT)-associated EB1 tracking and Western blotting of MT posttranslational modifications indicated no change in MT dynamics that could explain the vimentin collapse. However, pharmacological inhibition of the RhoGTPase Cdc42 in Hic-5 knockout CAFs rescued the vimentin collapse, while pan-formin inhibition with SMIFH2 promoted vimentin collapse in Hic-5 heterozygous CAFs. Our results reveal novel regulation of vimentin organization/dynamics by the FA scaffold protein Hic-5 via modulation of RhoGTPases and downstream formin activity.

INTRODUCTION Vimentin is a type III intermediate filament (IF; Huber et al., 2015) that is highly expressed in fibroblasts and other mesenchymal cells and is an integral component of the cytoskeleton. Due to the unique structural resiliency of IFs (Eriksson et al., 2009; Huber et al., 2015; Lowery et al., 2015), these proteins play key roles in maintaining the integrity of cells under high mechanical stress, such as during fibroblast-mediated contraction of the extracellular matrix (ECM; Eckes et al., 2000; Wei et al., 2008; Vuoriluoto et al., 2011; Kidd et al., 2014; Costigliola et al., 2017). In addition to their structural function and roles in response to mechanical stimuli, vimentin filaments also regulate cell shape, cell motility, and signal transduction (Eckes et al., 2000; Helfand et al., 2003; Guo et al., 2014; Chernoivanenko et al., 2015). Importantly, modifications in vimentin organization regulate its function in these cellular processes.

Vimentin assembles spontaneously into tetramers that oligomerize into unit-length filaments (ULFs). These assemble further via end-to-end annealing and lateral association to form filaments and filament networks (Engel et al., 1985). In response to extracellular...
cues, vimentin filaments can reorganize by exchange with filament precursors (tetramers and ULFs) transported by microtubule (MT)-and F-actin–associated motor proteins to support their interactions with other cytoskeletal elements (Leduc and Etienne-Manneville, 2017). Kinase-mediated phosphorylation slows subunit assembly or promotes filament disassembly, while phosphatase-mediated dephosphorylation opposes these effects, together regulating vimentin assembly and reorganization (Eriksson et al., 2004; Snider and Omary, 2014). For example, Rac1 activity within nascent lamellipodia promotes p21-activated kinase (PAK)-mediated vimentin phosphorylation and disassembly (Goto et al., 2002). This localized disassembly of vimentin allows Cdc42 activation of actin nucleation and actin branching enzymes including Arp2/3 and FMNL2/3 formins (Small et al., 2002; Helfand et al., 2011; Kage et al., 2017) and their localization to the cell membrane by Rac1-promoted cortactin recruitment (Weed et al., 2000), ultimately resulting in F-actin–driven force generation and cell membrane protrusion. Vimentin filaments also act as a template for growing MTs during cell polarization (Gan et al., 2016) and regulate organization of integrin-mediated traction forces required for directed cell migration (Costigliola et al., 2017). Therefore, impairment of either Rac1/PAK signaling or vimentin expression impedes directional migration as a result of impaired MT and actin organization in the absence of an intact vimentin cytoskeleton capable of responding to mechanical stimuli (Helfand et al., 2011). Despite increasing evidence of a crucial role for vimentin cross-talk with other cytoskeletal elements in integrating functional cellular responses to mechanical cues, knowledge of the underlying mechanisms for translating these mechanical cues into global vimentin rearrangements that participate in these cellular processes is incomplete.

Recent studies have identified integrin-based cell–ECM adhesions, known as focal adhesions (FAs), as regulators of IF dynamics (Burgstaller et al., 2010; Gregor et al., 2014; Scarpa and Mayor, 2016), although the mechanisms remain unclear. FAs form within lamellipodia of migrating cells, where integrin engagement with the ECM leads to the recruitment of adaptor proteins that link FAs to the F-actin cytoskeleton. The resulting FA–actin associations provide anchoring points on which the cell can exert traction forces for migration (Gardel et al., 2008, 2010; Blangy, 2017; Haage et al., 2018). Actin forms contractile bundles in association with nonmuscle myosin II (called stress fibers) and application of force by these FA-associated stress fibers helps to promote maturation of focal adhesions into fibblar adhesions (Pankov et al., 2000; Zamir et al., 2000; Goreczny et al., 2018). Conversely, MTs display increased catastrophe rates at FAs (Efimov et al., 2008) and deliver molecules involved in FA disassembly to the FA in a motor protein–dependent manner (Ezratty et al., 2005; Yue et al., 2014). Furthermore, perturbation of MTs with colchicine promotes the maturation of newly formed focal contacts into FAs, suggesting that MTs regulate FA maturation and lifetime negatively (Lloyd et al., 1977; Kavera et al., 1998). Interestingly, MTs have also been observed to elongate toward immature FAs in the cell periphery, where MT association with the immature FA can result in stabilization of these MTs (Kavera et al., 1998). Vimentin has been reported to interact directly with FAs via association with the cytoplasmic tails of ε2β1 and β3 integrins (Kreis et al., 2005; Kim et al., 2016) and indirectly via the linker protein plectin 1f (Bhattacharya et al., 2009; Burgstaller et al., 2010). Vimentin–FA interactions are largely associated with a subset of large, mature FAs (Burgstaller et al., 2010), while vimentin filament precursors (ULFs and tetramers) are present at immature focal complexes, suggesting that vimentin filament coupling to FAs may play a role in FA maturation (Terriac et al., 2017). While the interaction of F-actin and MTs with FAs has been extensively studied, the interactions between vimentin and FAs and the implications of these interactions on the regulation of vimentin organization and function remain poorly understood.

Hic-5 is an important FA adaptor/scaffold protein of the Paxillin family (Brown and Turner, 2004; Deakin and Turner, 2008; Deakin et al., 2009, 2012). We have previously shown that ablating Hic-5 in murine cancer-associated fibroblasts (CAFs) promotes F-actin cytoskeleton abnormalities, reduces cell contractility, and decreases the formation of central fibillary adhesions, thereby reducing extracellular fibronectin remodeling (Goreczny et al., 2017, 2018). Additionally, Hic-5 is up-regulated following TGF-β stimulation (Tumbarello and Turner, 2007) and is associated with the epithelial–mesenchymal transition (Shibanuma et al., 1994; Wang et al., 2008; Pignatelli et al., 2012). It is also important in maintaining an activated fibroblast phenotype (Dabiri et al., 2008) by promoting increased synthesis of ECM components, increased contraction of the ECM, and an increased proportion of mature FAs (Dabiri et al., 2008; Yue et al., 2014; Goreczny et al., 2017).

In the present study, we characterized a collapsed vimentin network and F-actin “hole” phenotype associated with Hic-5 ablation in CAFs, normal lung fibroblasts (LFs), and human foreskin fibroblasts (HFFs). The dynamics of the collapsed vimentin filament network was assessed by fluorescence recovery after photobleaching (FRAP) and Western blot analysis of total and phosphorylated vimentin levels in detergent-insoluble (filamentous) and detergent-soluble (filament precursors) cellular fractions. The phenotype with vimentin collapse and an F-actin hole was recapitulated in Hic-5 heterozygous (Hic-5 Het) CAFs following inhibition of formin activity, while rescue experiments suggest an important role for Hic-5 regulation of Cdc42 activity. These findings provide novel evidence that the FA-associated scaffold protein Hic-5 regulates vimentin dynamics and organization by modulating Rho family GTPases and the activity of formins downstream of these GTPases. Importantly, these data advance our limited knowledge of FA–IF cross-talk.

RESULTS

Hic-5 regulates vimentin organization in cancer-associated and normal fibroblasts

Communication between FAs and IFs has been recently implicated in regulation of FA dynamics and vesicular trafficking to modulate cell migration in normal mesenchymal cells and cells undergoing EMT at sites of tumor formation (Leube et al., 2015; Liu et al., 2015). Coupling of IFs to FAs has been shown previously to increase FA size and strength of adhesion to the matrix while decreasing FA turnover, but the factors that promote IF recruitment to or dissociation from FAs and the subsequent effects on IF organization and dynamics are not known (Tsuruta and Jones, 2003; Bhattacharya et al., 2009; Havel et al., 2015; Osmanagic-Myers et al., 2015). Based on recent studies from our laboratory indicating a role for Hic-5 in promoting FA maturation into fibblar adhesions in cancer-associated fibroblasts (CAFs; Goreczny et al., 2018), the evidence of a role for IFs in FA and fibbrilar adhesion maturation, and previous data that describe a reduction of FA–IF interactions when FA maturation is impaired (Burgstaller et al., 2010), we sought to investigate if Hic-5 has a role in mediating the FA–vimentin intermediate filament cross-talk implicated in these processes.

Hic-5-null CAFs were isolated from a constitutive Hic-5 knockout (KO) murine breast tumor model as previously described: Hic-5 KO mouse, mammary tumor virus-polymavirus Middle T antigen (MMTV-PyMT) strain (Goreczny et al., 2017, 2018). Hic-5
FIGURE 1: Hic-5 regulates vimentin organization in CAFs. (A, B) Representative images of the vimentin, actin, and microtubule cytoskeleton of Hic-5 Het and Hic-5 KO CAFs at 4- and 24-h postplating with (C, D) quantification of the percentage of cells with perinuclear collapse of vimentin and with reduction in centrally located actin stress fibers (actin hole; n = at least 60 cells/condition). (E) Vimentin collapse observed in Hic-5 KO CAFs was also quantified as an increased ratio of perinuclear to peripheral vimentin filaments (n = at least 60 cells/condition). (F) Total cell area and percentage of total cell area occupied by vimentin was decreased in Hic-5 KO CAFs (n = at least 75 cells/condition). (G) Images and (H, I) quantification of exogenous EGFP-Hic-5 rescue of vimentin collapse and the actin hole phenotype 4 h postplating (n = at least 41 cells/condition). All data are shown as the mean ± SEM and are collected from three independent experiments. **, p < 0.01; ***, p < 0.001; ****, p < 0.0001. Scale bar = 50 μm. All CAF experiments were from three unique Hic-5 Het CAF cell lines and one Hic-5 KO CAF cell line.

heterozygote (Het) CAFs were utilized in place of wild-type (WT) CAFs due to the absence of phenotypic differences between WT and Hic-5 Het CAFs and the higher abundance of Hic-5 Het animals available, as previously described (Goreczny et al., 2017). The organization of the three cytoskeletal systems was evaluated by confocal imaging. Visualization of vimentin, the principal IF in fibroblasts, MTs, and F-actin in primary Hic-5 Het and Hic-5 KO CAFs revealed striking differences in F-actin and vimentin cytoskeletal organization (Figure 1, A and B). The vimentin cytoskeleton of Hic-5 KO CAFs was collapsed into closely associated bundles of coaligned vimentin filaments, previously termed “cables” (Hollenbeck et al., 1989), that were restricted to the perinuclear region of the cell, with the collapsed vimentin appearing to coil around the nucleus in some cells. Organization of vimentin was analyzed at an early time point (4 h), when cells are still actively spreading, and a later time point (24 h) post cell plating, corresponding to a fully spread steady state. Vimentin organization was qualitatively classified as filamentous (normal distribution), partially collapsed, or collapsed (see Materials and Methods for details). Strikingly, 75–90% of Hic-5 KO CAFs versus 0–3% of Hic-5 Het CAFs were observed to have collapsed vimentin at 4 and 24 h (Figure 1C). Additionally, 72–83% of Hic-5 KO CAFs and 0–10% of Hic-5 Het CAFs were observed to have peripheral F-actin organization with a reduced amount of centrally located F-actin stress fibers (“F-actin hole” phenotype) at 4 and 24 h (Figure 1D), as previously reported (Goreczny et al., 2018). The vimentin collapse and exclusion of actin stress fibers from the area occupied by the collapsed vimentin (the “F-actin hole”) is consistent with previous observations of vimentin and actin reorganization following drug-induced MT depolymerization (Hollenbeck et al., 1989). However, MT filament organization (as visualized by α-tubulin staining) was not grossly different between Hic-5 Het and Hic-5 KO CAFs (Figure 1, A and B).

The increase in vimentin staining intensity resulting from compaction and perinuclear localization of vimentin filaments was confirmed with quantitative analyses. Hic-5 KO CAFs displayed a threefold higher ratio of perinuclear/peripheral vimentin mean fluorescence intensity (MFI) than Hic-5 Het CAFs at all time points, while the perinuclear/peripheral ratio of MT MFI was not significantly different between Hic-5 Het and Hic-5 KO CAFs (Figure 1E; see Materials and Methods for details).
and Methods for details on defining perinuclear and peripheral regions). Additionally, Hic-5 KO CAFs had a 50% reduction in the percentage of cell area occupied by vimentin compared with Hic-5 Het CAFs (Figure 1F), and the total cell area, as measured by F-actin staining, was reduced in Hic-5 KO CAFs (Figure 1F). The vimentin collapse was rescued by exogenous expression of EGFP-Hic-5 in Hic-5 KO CAFs, shifting the percentage of cells with a normal, filamentous distribution of vimentin from 20 to 80% and reducing the percentage of cells with an F-actin hole from 85 to 30% (4-h time point, Figure 1, G–I). Importantly, previous studies have associated vimentin collapse with disruption of the MT cytoskeleton or its associated motor proteins (Hollenbeck et al., 1989; Prahlad et al., 1998; Rathje et al., 2014) and elevated activity of the GTPases Cdc42 and/or Rac1 (Meriane et al., 2000; Chan et al., 2002). Based on the well-documented role of paxillin in regulating Rho family GTPase activity (Turner, 2000; Deakin et al., 2009), MT acetylation (Deakin and Turner, 2014) and paxillin’s functional cross-talk, mutual interacting proteins, and extensive structural homology with Hic-5 (Turner, 2000; Brown and Turner, 2004; Deakin and Turner, 2008), we hypothesized that depletion of paxillin may also induce a vimentin phenotype in fibroblasts. However, staining of paxillin KO mouse embryonic fibroblasts (MEFs) showed no indication of either a vimentin collapse or an F-actin hole (Supplemental Figure 1A), indicating that Hic-5 has a nonredundant role in regulating vimentin organization.

Despite their extensive use in the literature (Prahlad et al., 1998; Meriane et al., 2000; Chan et al., 2002; Rathje et al., 2014; Tsui et al., 2018), the methods for vimentin organization analysis described above remain time-inefficient and limited in capability, prompting our use of an automated image analysis pipeline to assess vimentin organization. The Broad Institute Cell Profiler 3.0 software (Carpenter et al., 2006; Jones et al., 2008) was utilized to create a workflow to analyze the distribution of vimentin within the cell via the program’s radial distribution module. Consistent with the manual measurements, both the heat map and the graph of vimentin mean fractional intensity, which depicts the distribution of vimentin fluorescence intensity in 16 equal-sized radial bins extending from the center of the nucleus to the outer edge of the cell, show a higher concentration of vimentin localized to a smaller region within the cell (perinuclear localization) in Hic-5 KO CAFs (Figure 2, A–C). Conversely, expression of EGFP-Hic-5 (but not EGFP vector) decreased perinuclear vimentin levels and extended the peak of the vimentin radial distribution further into the peripheral region of the cell (Figure 2, D–F).

Importantly, the CAFs used in these initial studies are fibroblasts converted to an active state within the tumor microenvironment, resulting in inherent modifications to their cytoskeleton that differ from their normal fibroblast counterparts. For example, CAFs often display increased actin stress fibers and elevated ε-smooth muscle actin expression (Rasalan and Vaheri, 2010). This causes increased cellular contractility, aiding in CAF-mediated remodeling of the ECM to promote tumor invasion (Rasalan and Vaheri, 2010; Albregues et al., 2015; Shiga et al., 2015; Kalluri, 2016; Goreczny et al., 2017; LeBlu and Kalluri, 2018). Due to these inherent cytoskeletal differences between CAFs and normal fibroblasts, we also assessed whether Hic-5 depletion induced vimentin collapse in normal lung fibroblasts (LFs) isolated from Hic-5 KO normal (nontumorigenic) mice and in normal human foreskin fibroblasts (HFFs) treated with small interfering RNAs (siRNAs) targeting Hic-5 expression (Figure 3). Hic-5 KO LFs showed a phenotype similar to that of Hic-5 KO CAFs with 68% of cells exhibiting vimentin collapse, as confirmed by a shift toward more perinuclear accumulation of vimentin visualized by automated radial distribution analyses. Additionally, 71% of these cells exhibited an F-actin hole (Figure 3, A–E). Following siRNA-mediated Hic-5 knockdown (KD) in HFFs (Hic-5 KO was 87 ± 5% and 67 ± 5% for Hic-5 siRNA 1 and Hic-5 siRNA 2, respectively; Figure 3, F–H), 61–63% of cells had collapsed vimentin (Figure 3I) associated with a shift in vimentin radial distribution (Figure 3, K and L) and 48–50% had an F-actin hole (Figure 3J). In contrast, RNA interference (RNAi)-mediated depletion of paxillin in HFFs (Deakin and Turner, 2011) failed to induce a vimentin collapse, as was also observed in paxillin KO MEFs (Supplemental Figure 1B). In summary, Hic-5, but not paxillin, is required for establishing and maintaining a filamentous, well-spread vimentin network and the presence of centrally located F-actin stress fibers in normal fibroblasts (LFs and HFFs) and CAFs.

Hic-5 is required for key vimentin phosphorylation events that maintain normal vimentin dynamics, independent of microtubules

We hypothesized that the perinuclear collapse of vimentin filaments observed in Hic-5 KO CAFs represents a shift of cellular vimentin into a hyperassembled state, with potential effects on its phosphorylation status. Importantly, the proportion of filamentous vimentin versus vimentin filament precursors (unit length filaments [ULFs] and tetramers) can be evaluated by fractionating the cell lysate with the detergent Triton X-100. Following centrifugation of the cell lysate, filamentous vimentin pellets (Triton X-100–insoluble material [TX100-insoluble]) and the filament precursors remain in the supernatant (Triton X-100–soluble material [TX100-soluble]); see Materials and Methods for details; Shea, 1990; Cheng et al., 2003; Vohnoutka et al., 2017). We therefore analyzed the TX100-insoluble versus TX100-soluble fractions of Hic-5 Het and Hic-5 KO CAFs for total vimentin and vimentin phosphorylated at several key epitopes known to regulate vimentin organization, including phospho-S72, 82, and 55 (Figure 4A; Tsujimura et al., 1994; Goto et al., 1998, 2003; Janosch et al., 2000; Yasui et al., 2001; Cheng et al., 2003; Eriksson et al., 2004; Yamaguchi et al., 2005; Li et al., 2006). Hic-5 KO CAFs plated for 4 or 24 h had levels of TX100-insoluble vimentin equivalent to that of Hic-5 Het CAFs (Figure 4, A and B). However, Hic-5 KO CAFs had severely reduced TX100-soluble vimentin (Figure 4, A and B). The loss of TX100-soluble vimentin, in the absence of increased TX100-insoluble vimentin observed in Hic-5 KO CAFs, indicates either increased proteolysis of TX100-soluble vimentin, as vimentin phosphorylation may modulate IF susceptibility to proteolysis (Zhu et al., 2011), or that the shift of the small population of TX100-soluble vimentin to TX100-insoluble vimentin was below the limit of detectable difference of Western blot analysis. Interestingly, only the TX100-insoluble fractions of Hic-5 Het and Hic-5 KO CAFs were positive for vimentin phospho-epitopes. Hic-5 KO CAFs showed a decrease in both phospho-S72 and phospho-S82 vimentin epitopes in the TX100-insoluble fraction but did not have altered phospho-S55 (Figure 4, C–E). Notably, paxillin KO MEFs did not phenocopy the loss of TX100-soluble vimentin observed in Hic-5 KO CAFs (Supplemental Figure 2A) and paxillin KO MEFs had a significantly higher percentage of total vimentin within the TX100-soluble fraction than Hic-5 KO CAFs (Supplemental Figure 2B). These data indicate that the majority of vimentin within Hic-5 KO CAFs is assembled into vimentin filaments, most likely due to a reduction in the phosphorylation level required to maintain nonassembled, dynamic vimentin. The decrease in the amount of TX100-soluble vimentin and vimentin phosphorylation in Hic-5 KO CAFs suggests an impairment in vimentin dynamics, which is regulated by these phosphorylation events (Eriksson et al., 2004; Snider and Omary, 2014).
FRAP analyses have been utilized previously to determine the dynamics of IFs in various cell types, with the time for fluorescence recovery in bleached regions of vimentin reported to be 2–8 min for 50% recovery and 11–23 min for full recovery (Yoon et al., 1998; Leduc and Etienne-Manneville, 2017), which is significantly longer than that for actin stress fibers (5–10 min; Hotulainen and Lappalainen, 2006; Campbell and Knight, 2007) or MTs (20 s–6.5 min; Saxton et al., 1984; Omelyanchuk and Munzarova, 2017). To assess possible differences in vimentin dynamics in the presence and absence of Hic-5, we performed FRAP of mCherry-vimentin in Hic-5 Het and Hic-5 KO CAFs (Figure 5; Supplemental Movie S1). The vimentin fluorescence in the bleached zone of Hic-5 Het CAFs recovered to 80% of the prebleach fluorescence intensity in 15 min (900 s), while the vimentin fluorescence of Hic-5 KO CAFs recovered only by 30% in 15 min (Figure 5, A and B). These percentage recoveries indicate that 20 and 70% of the bleached vimentin remained immobile during the bleach recovery period, which corresponds to fluorescence recovery rates of 5 and 2% per minute in Hic-5 Het and Hic-5 KO CAFs, respectively (Figure 5C). Ectopic expression of EGFP-Hic-5 was able to partially rescue the percentage of vimentin fluorescence recovery in Hic-5 KO CAFs to 62%, which corresponds to an immobile fraction of 38% and a
FIGURE 3: Hic-5 regulates vimentin organization in normal mouse and human fibroblasts. (A) Images and (B, C) quantification of percentage of Hic-5 Het and Hic-5 KO normal LFs with collapsed vimentin and an actin hole at 4 h postplating (n = at least 140 cells/condition). (D, E) Vimentin mean fractional intensity showing perinuclear vimentin localization in Hic-5 KO LFs (n = at least 66 cells/condition). (F) Images of HFFs following Hic-5 siRNA KD with (G, H) Western blot analysis of KD efficiency. (I, J) Increased percentage of HFFs with vimentin collapse and an actin hole following Hic-5 depletion (n = at least 102 cells/condition). (K) Corresponding heat maps of HFFs treated with Hic-5 siRNA and (L) graphical representation of this vimentin fluorescence distribution (n = at least 90 cells/condition). All data are shown as the mean ± SEM. Data for LFs (A–E) were collected from two independent experiments, while data from HFFs (F–L) were collected from four independent experiments. * p < 0.05; **, p < 0.01; ****, p < 0.0001. Scale bar = 50 μm. Assessment of HFF cells utilized one HFF cell line obtained from the ATCC, while LF experiments were carried out with two unique Hic-5 Het and two unique Hic-5 KO LF cell lines.
Hic-5 regulates vimentin organization and dynamics by modulating RhoGTPases and formin-mediated actin organization

Elevation of Cdc42 and Rac1 RhoGTPase activity has been shown to induce vimentin collapse in cultured fibroblasts (Meriane et al., 2000; Chan et al., 2002), and we have previously shown that Hic-5 KD in human MDA-MB-231 breast epithelial tumor cells alters RhoGTPase signaling (Deakin et al., 2009, 2012; Deakin and Turner, 2011). Importantly, the disruptions in actin organization, including the reduced centrally located actin stress fibers, impaired contraction of collagen gels, and reduction in mature FAs observed in Hic-5 KO CAFs (Figure 1; Engel et al., 1985;
FIGURE 5: Depletion of Hic-5 reduces vimentin dynamics. (A) Individual frames from confocal movies of Hic-5 Het and Hic-5 KO CAFs transfected with mCherry–vimentin subjected to FRAP analysis (Supplemental Movie S1) indicate a marked reduction in vimentin dynamics. Insets depict regions where mCherry–vimentin was bleached and recovery followed over 15 min (1 frame/30 s). (B, C) Quantification of % fluorescence recovery and rate of fluorescence recovery of mCherry–vimentin with measurements of fluorescence values normalized for whole-frame bleaching induced by repeated imaging (n = at least 8 cells/condition). (D) Individual frames of FRAP experiments in which Hic-5 KO CAFs were transfected with mCherry–vimentin and EGFP or EGFP-Hic-5 (Supplemental Movie S2). (E, F) Quantification of the % fluorescence recovery and rate of fluorescence recovery of mCherry–vimentin (n = at least 8 cells/condition). All data represent the mean ± SEM and are collected from three independent experiments. * p < 0.05; ****, p < 0.0001. Scale bar = 50 μm.
and Hic-5 have been shown to regulate migration by modulating (Eckes et al., 2013) three-dimensional matrix environments cancer-associated fibroblast migration in two- and three-dimensional environments.

Interestingly, both Cdc42 and Rac1 inhibition also decreased the percentage of cells with an F-actin hole from 62% to 16 and 41%, respectively (Figure 7C). Inhibition of Cdc42 activity resulted in a significant reduction in peak height and increase in peak width of vimentin radial distribution, indicative of vimentin reorganization (Eckes et al., 1998). In contrast, vimentin KD in our Hic-5 KO fibroblasts did not affect migration velocity but did reduce migration directionality and persistence (Supplemental Movie S8; Figure 8, E and F). Directionality and persistence were unaffected by Rac1 inhibition (Figure 8, D and E). Additionally, inhibition of Cdc42 increased the average major axis length (size) of Hic-5 KO cells, as indicated by the increased distance on the x axis displaying the vimentin radial distribution graph of Cdc42 inhibition (Figure 7E).

Decreased Cdc42/Rac1 activity has also been associated with increased RhoA activity (Kholmanskih et al., 2003; Zhang et al., 2007), which in turn promotes the up-regulation of actin stress fiber formation via multiple mechanisms, including activation of formin family members (Oakes et al., 2017). We chose to focus on investigation of formin-mediated actin polymerization because of the known role of formins in the formation/growth of FA-associated actin stress fibers (Burrage and Geiger, 2016; Livne and Geiger, 2016), coupled with our previous data showing that Hic-5 KO cells exhibit reduced centrally located actin stress fibers resulting from impaired tensin-1–mediated coupling of actin stress fibers with FAs during their maturation to fibrillar adhesions (Goreczny et al., 2017, 2018). Therefore, we hypothesized that Cdc42 inhibition may rescue the central stress fiber assembly and possibly the vimentin collapse observed in Hic-5 KO fibroblasts by increasing the activity of the opposing RhoA pathway and RhoA-activation of formins that promote FA-associated stress fiber formation (Burrage and Geiger, 2016; Livne and Geiger, 2016). To assess the role of formins, we investigated the effects of the panformin inhibitor SMIFH2 (30 μM) on vimentin organization in Hic-5 KO fibroblasts. Strikingly, treatment of Hic-5 KO fibroblasts with SMIFH2 resulted in both vimentin collapse in 94% of cells and the presence of an F-actin hole in 58% of cells (Figure 7, F–H). SMIFH2-induced vimentin collapse produced a substantial shift in vimentin radial distribution, with a significant increase in peak height, decrease in peak width, and decrease in major axis length, indicative of vimentin collapse and decreased cell size (Figure 7, I and J).

These data suggest that ablation of Hic-5 likely results in changes in RhoGTPase activity, including a shift in the balance of Cdc42 versus RhoA activity, which in turn may promote vimentin collapse and F-actin redistribution either directly or indirectly via regulation of formin activity.

RhoGTPase and formin perturbations of Hic-5 knockout cancer-associated fibroblasts increase the velocity of cancer-associated fibroblast migration in two- and three-dimensional matrix environments

Coordinated RhoGTPase-mediated regulation of MT, actin, and vimentin dynamics/organization is crucial to productive cell migration (Eckes et al., 2000; Eriksson et al., 2009; Ridley, 2015). Both paxillin and Hic-5 have been shown to regulate migration by modulating RhoGTPase activity, FA turnover, distribution of key polarity proteins, MT acetylation, and actin stress fiber formation (Deakin et al., 2009, 2012; Deakin and Turner, 2011, 2014; Goreczny et al., 2018). However, whether Hic-5’s role in regulating vimentin collapse alters cell migration has not been investigated. Initial observations revealed that Hic-5 KO fibroblasts were more motile and migrated at a higher velocity on fibronectin in two-dimensional environments (fibronectin-coated tissue culture dishes) than Hic-5 KO fibroblasts but did not display alterations in directionality (Figure 8, A and B, directionality not shown; Supplemental Movie S4). Ectopic expression of EGFP-Hic-5 in Hic-5 KO fibroblasts was able to reduce migration velocity in 2D (Figure 8, C and D; Supplemental Movie S5; trending toward significance). Similar Hic-5-dependent migration changes were also observed in fibroblasts migrating within a three-dimensional cell-derived matrix (3D-CDM) environment (Supplemental Figure 3, A–D; Supplemental Movies S6 and S7). Importantly, Hic-5 KO resulted in the opposite effect on migration from that seen after vimentin KO in normal fibroblasts, which has been previously shown to result in impaired directionality and decreased motility (Eckes et al., 1998).

We next investigated whether the phenotypes observed upon Cdc42 and RhoA inhibition of Hic-5 KO fibroblasts might be contributing to the motility differences observed in the Hic-5 KO fibroblasts. Inhibition of Cdc42 with Zcl278 reduced the migration velocity of Hic-5 KO fibroblasts (Figure 8, H and I; Supplemental Movie S9), and, unexpectedly, treatment of Hic-5 KO fibroblasts with the pan-formin inhibitor, SMIFH2 also reduced migration velocity (Figure 8, J–K; Supplemental Movie S10). This reduction in velocity likely indicates that panformin inhibition disrupts multiple formins required for actin-driven cell propulsion, while the effects of Hic-5 KO on formin activity are more specific and do not impair formin activity involved in actin-driven cell propulsion. These data indicate that Hic-5 KO–mediated vimentin collapse does not phenocopy the effects of vimentin KD on cellular motility, and the increased migration velocity of Hic-5 KO fibroblasts is due to altered Rho family GTPase activity, but not to formin-mediated actin stress fiber formation defects.

**DISCUSSION**

Vimentin organization has been shown to regulate a broad array of important cellular functions and biological processes such as polarized lamellipodia formation (Helfand et al., 2011), directional persistence of migrating cells (Gan et al., 2016), and maintenance of cell–cell junctions during collective cell migration in wound healing and development (Li et al., 2013; Menko et al., 2014; Scarpa and Mayor, 2016). Vimentin also associates with points of integrin attachment to the ECM (FAs), where it interacts with the cytoskeletal linker protein plectin 1f (Burgstaller et al., 2010; Castanon et al., 2013), the cytoplasmic tails of several integrins, and potentially other FA-associated proteins or other cytoskeletal elements (Kreis et al., 2005; Bhattacharya et al., 2009; Havel et al., 2015). These vimentin–FA interactions have been correlated with increased FA size, attachment strength to the ECM, and FA maturation (Tsurtua and Jones, 2003; Bhattacharya et al., 2009; Havel et al., 2015; Liu et al., 2015; Osmanagic-Myers et al., 2015). However, the proteins that mediate or promote these FA–vimentin interactions and the effects of these interactions on the vimentin cytoskeleton are poorly understood despite their clear implications for cellular migration, wound healing, metastasis, and many other important biological processes.

We report here a novel FA–vimentin signaling axis coordinated by the FA protein Hic-5. Ablation of Hic-5, but not its...
FIGURE 6: Hic-5 regulation of vimentin dynamics is independent of microtubules. (A–C) Western blot of Hic-5 Het and Hic-5 KO CAFs probed with antibodies against α-tubulin (total MT content of cells), acetylated MTs and tyrosinated MTs, accompanied by quantification normalized to GAPDH (data represent the mean ± SEM). (D) Individual frames of Hic-5 Het and Hic-5 KO CAFs transfected with EGFP-EB1 (MT plus end tip protein) and mCherry-EMTD (labels MTs) imaged with TIRF microscopy illustrating EB1 movement and localization to MT tips (Supplemental Movie S3). (E–J) Analysis of MT dynamics of Hic-5 Het and Hic-5 KO CAFs transfected with EB1-EGFP, imaged using TIRF microscopy over 3 min (1 frame/s), and analyzed using a MatLab particle tracker plug-in to track EB1 movement show increased MT growth but no change in overall dynamics in Hic-5 KO CAFs (>157,000 MT tracks/condition). Analyses are plotted as box-and-whisker graphs where the bounds of the box represent the interquartile range, the horizontal line within the box represents the median, and the whiskers represent the range of the data. Data are collected from three independent experiments. *, p < 0.05; **, p < 0.01. Scale bar = 50 μm.
close relative paxillin (Brown and Turner, 2004; Deakin and Turner, 2008), was shown to cause a MT-independent collapse of the vimentin cytoskeleton to the perinuclear region of the cell, as well as decreased dynamics and altered phosphorylation status (Figures 1–6; Supplemental Figures 1 and 2). Downstream of Hic-5, we have shown that elevated Cdc42 and, unexpectedly, decreased formin activity were the key causes of the vimentin collapse (Figure 7). These findings represent a previously unobserved pathway
Hic-5 modulates migration velocity independent of its effects on vimentin, but through its effects on RhoGTPases. Migration tracks (A, C, E, H, J) of individual cells from cultures of CAFs plated on two-dimensional culture plates coated with fibronectin are shown. Graphs of migration velocity of cells imaged over 16 h (1 image/10 min) are shown for each through which a FA scaffold protein exhibits cross-talk with the IF cytoskeleton and modulates its organization/dynamics as summarized in Figure 9.

The vimentin reorganization observed in Hic-5 KO CAFs represents a drastic alteration in the dynamics of vimentin and potentially its interaction with other proteins within the cell (Figures 1, 2, 4, and 5). Previous reports indicate that vimentin collapse is a two-step process in which vimentin filaments first undergo exacerbated filament packing to form thick vimentin cables, followed by reorganization of these vimentin cables into a perinuclear mass. Reorganization of the vimentin cables into a collapsed state, but not the formation of cables, is dependent on an intact actin cortex, which was hypothesized to generate the physical force required to mediate vimentin’s perinuclear localization (Hollenbeck et al., 1989). The cytoskeletal linker protein plectin1f was later found to be involved in physically linking vimentin to a subset of actin filaments, referred to as transverse arcs, that undergo retrograde flow and therefore exert a perinuclear drag on vimentin filaments (Jiu et al., 2015). Furthermore, siRNA depletion of plectin1f, or specific loss of transverse actin arcs by siRNA depletion of Tm4, in fibroblasts have been shown to promote “overspreading” of the vimentin network to the edge of the cell in circumstances when it would not typically extend this far, such as during cell migration (Jiu et al., 2015). Interestingly, these data suggest that plectin–vimentin interactions at actin arcs, and therefore the maintenance of actin arc formation and actin retrograde flow, are likely retained in the Hic-5 KO CAFs and thus may also contribute to the resulting vimentin collapse. Future analysis of the plectin1f distribution in Hic-5 KO cells is therefore warranted.

In the case of Hic-5 KO–mediated vimentin collapse, these findings are particularly interesting due to the actin abnormalities that were observed in conjunction with migration experiment (B, n = at least 60 cells/condition; D, n = at least 28 cells/condition; F, n = at least 47 cells/condition). Cells treated with inhibitors were imaged for 4 h at a frame rate of 1 image/10 min (I, n = at least 55 cells/condition; K, n = at least 85 cells/condition). Western blot was used to assess KD efficiency of vimentin in Hic-5 Het CAFs utilized for migration experiments (G, NT = nontarget siRNA and Vm siRNA = siRNA targeting vimentin transcripts; 35.5% ± 9.8%; p < 0.05). All data shown represent the mean ± SEM and are collected from at least three independent experiments. * p < 0.05; ***, p < 0.001.
vimentin collapse (F-actin holes; Goreczny et al., 2017, 2018). If actin retrograde flow of transverse arcs is involved in the Hic-5 KO-mediated vimentin collapse, this suggests that the formation of actin arcs or actin retrograde flow is not ablated in either Hic-5 KO CAFs (Figures 1 and 2) or formin inhibitor–treated CAFs (Figure 7), which is consistent with persistence of peripheral actin structures under both of these conditions. Hic-5 KO may selectively disrupt the formation or maintenance of central FA-associated stress fibers, which is consistent with its effects on FA maturation (Goreczny et al., 2018) and the presence of an F-actin hole in Hic-5 KO CAFs. In the case of pharmacological inhibition of formins, this does not inhibit 100% of their activity and the high nucleotide efficiency of several formins (Breitsprecher and Goode, 2013), coupled with the specificity of SMIFH2 for formin-mediated actin assembly without affecting activity of other actin nucleation and polymerization enzymes (Rizvi et al., 2009), would result in incomplete actin filament assembly inhibition. This could indicate that Hic-5 KO CAFs have reduced overall actin polymerization, which directly contributes to the F-actin hole, the reduced contractility of cells, the reduction in FA maturation, and the vimentin collapse observed in Hic-5 KO CAFs, without completely ablating actin retrograde flow. This is consistent with the presence of several different kinds of actin stress fibers in fibroblasts (dorsal, ventral, and transverse arcs) that all have different assembly mechanisms and separate regulatory pathways that control their formation and disassembly (Vallenius, 2013). Further perturbation of individual components of actin biology, such as the formation of transverse arcs, utilizing both chemical inhibitors and siRNA KD (such as KD of tropomyosin-4 to disrupt actin arc formation; Jiu et al., 2015), would be of great interest for future studies in order to further investigate this novel connection between formin activity and vimentin organization.

The causes of vimentin collapse observed in other studies are not fully understood, but several different contributing factors have been identified, including chemical perturbation of the MT cytoskeleton that promotes its disassembly (Hollenbeck et al., 1989; Rathje et al., 2014), inhibition of the motor protein kinesin or overexpression/up-regulation of the motor protein dynein (Prahial et al., 1998), and dysregulation of RhoGTPases and their downstream effectors (Meriane et al., 2000; Chan et al., 2002). In Hic-5 KO CAFs, we have shown that pharmacologic inhibition of Rac1 slightly reduces the percentage of cells with an F-actin hole, while inhibition of Cdc42 efficiently rescues both the vimentin collapse and the F-actin hole phenotype (Figure 7), suggesting that Hic-5 KO promotes vimentin collapse and the loss of central stress fibers via the elevation of Cdc42 activity. It is important to note that the Cdc42 inhibitor utilized for these experiments (Zcl278) has been reported to act through inhibition of the interaction of Cdc42 with one of its activators, the guanine nucleotide exchange factor intersectin-1 (Friesland et al., 2013). Interestingly, exogenous expression of intersectin-1 in human lung cancer cells has been associated with vimentin collapse (Jegathan et al., 2016). However, the regulation of RhoGTPases and their effects on cytoskeleton and cell shape may have opposing effects in epithelial cells and in fibroblasts (Deakin et al., 2012). Furthermore, Zcl278 binds directly to the site on Cdc42 where intersectin-1 binds, which affects the ability of intersectin-1 to activate Cdc42, but does not affect other functions of intersectin-1 that may otherwise induce vimentin collapse (Friesland et al., 2013). These findings could suggest that Hic-5 may have a direct role in intersectin-1–mediated activation of Cdc42. However, our data more strongly support a mechanism in which Hic-5 KO leads to reduced RhoA activity, in turn promoting increased Cdc42 and Rac1 activity and a corresponding decrease in the activity of certain formins, leading to vimentin collapse (Figure 9).

As discussed, the actin abnormalities and subsequent rescue of both actin and vimentin abnormalities following Cdc42 inhibition in Hic-5 KO CAFs are consistent with impaired signaling of the opposing RhoA pathway. In particular, GTP-bound RhoA is known to bind to several formins crucial for FA-associated stress fiber formation and contribute to their conversion from an autoinhibited state to an

![FIGURE 9: Hic-5 regulates vimentin and actin organization via its modulation of RhoGTPases and downstream formin activity. Depletion of the focal adhesion adaptor protein Hic-5 in CAFs and normal fibroblasts results in altered vimentin phosphorylation and vimentin dynamics that are associated with collapse of the vimentin cytoskeleton. Additionally, loss of Hic-5 decreases centrally located actin stress fibers (F-actin hole). These cytoskeleton abnormalities perturb migration dynamics increasing migration velocity of CAFs. Both changes in migration dynamics and abnormalities in vimentin and actin organization are rescued by inhibition of the RhoGTPase Cdc42, while treatment of Het Hic-5 CAFs with the pan formin inhibitor SMIFH2 promotes vimentin collapse and the loss of centrally located actin stress fibers, but not increased migration velocity. These data indicate cross-talk between FAs, vimentin, and actin that may regulate the transition from a promigratory fibroblast (low Hic-5), in which a collapsed vimentin network may allow the cells to navigate through a complex ECM, to a procontractile phenotype (high Hic-5), where increased actin stress fibers and an extended vimentin cytoskeleton promote matrix remodeling.](image-url)
active state. This suggests that Hic-5 KO may reduce the activity of several formins via a reduction in RhoA activity and a corresponding elevation in Cdc42/Rac1 activity (Lammers et al., 2005; Rose et al., 2005; Kuhn and Geyer, 2014). Furthermore, vimentin collapse associated with increased Cdc42/Rac1 signaling was previously shown to be mediated by the activation of two of their downstream targets, PAK and p70 S6 kinase, which phosphorylate vimentin at S72, S55, and multiple other sites (Chan et al., 2002). In Hic-5 KO CAFs, we observed changes in the level of phosphorylation of several key vimentin phosphorylation sites known to regulate its assembly, including a decrease in S72 and S82 phosphorylation, but no difference in phosphorylation of S55 (Figure 4). This shift in vimentin phosphorylation likely contributes to both the loss of soluble vimentin (Figure 4) and the reduction in vimentin dynamics (Figure 5).

Interestingly, the vimentin phosphorylation epitopes were detected in the TX100-insoluble fraction of vimentin, but not the TX100-soluble fraction, in both Hic-5 Het and Hic-5 KO CAFs. We hypothesize that this is in part due to the common use of phosphatase inhibitors before cell lysis, fractionation, and the assessment of vimentin phosphorylation in studies that observe soluble phosphorylated vimentin. These phosphatase inhibitors artificially stabilize vimentin phosphorylation and elevate the levels of TX100-soluble vimentin (Erikkson et al., 2004). It has previously been hypothesized that in cells not treated with phosphatase inhibitors, phosphatases would likely display a preference for soluble vimentin subunits in order to maintain the low levels of TX100-soluble vimentin normally observed in the absence of phosphatase inhibitors/pharmacological enrichment of vimentin phosphorylation (Erikkson et al., 2004). Therefore, in the absence of phosphatase inhibitors, phosphatase preference for soluble vimentin subunits would result in low levels of soluble vimentin phosphorylation and would promote reincorporation of soluble subunits into vimentin filaments over accumulation of the soluble subunits. In addition, other studies have neglected to investigate vimentin phosphorylation in CAFs, which are likely to have more TX100-insoluble vimentin than normal fibroblasts due to their activated, contractile status, associated with their growth within the tumor ECM, which displays a much higher rigidity than normal ECM (Murray et al., 2014). It will be of interest to further probe which specific kinases are acting downstream of Hic-5 to regulate vimentin organization and to perform a comprehensive phosphoproteomic analysis of Hic-5–dependent regulation of vimentin phosphorylation. Subsequent use of kinase inhibitors and/or site-directed mutagenesis of key vimentin phosphorylation consensus sites may be successful in addressing their respective roles in Hic-5–dependent organization of the vimentin network. However, to the best of our knowledge, there is no single phospho site mutation of vimentin known to cause network collapse, while mutation of one or more phospho sites in keratin IFs has been shown to promote their collapse, suggesting a complex interrelationship (Kakade et al., 2016; Sawant et al., 2018).

The dysregulation of Rho GTTPases in Hic-5 KO CAFs may also stimulate the changes in fibroblast migration observed on two- and three-dimensional substrates (Figure 8; Supplemental Figure 3). RhoA promotes the formation of actin stress fibers, which, along with actin retrograde flow, have been shown to promote migration in mesenchymal cells (Anderson et al., 2008; Vallienius, 2013; Katsumo et al., 2015; Inagaki and Katsumo, 2017). However, Hic-5 KO CAFs migrate faster (Figure 8, A and B) than their Hic-5 Het counterparts despite having reduced actin stress fibers (Figure 1) as well as impaired RhoA signaling, likely due to elevation of Cdc42 (Figure 7). Increased migration velocity in cells with small numbers of actin stress fibers has been noted in several additional cell types including immune cells (macrophages and leukocytes; Friedl and Wolf, 2003, 2010). Surprisingly, depletion of RhoA/B in macrophages also increased their migration velocity and phosphoryosyn light chain activity, a component of actin stress fibers that mediates their contractility (Friedl and Wolf, 2010). These seemingly conflicting observations are the result of different modes of cell migration (mainly amoeboid versus mesenchymal) that employ unique profiles of actin dynamics and FA composition, lifetime, and interaction with the ECM to facilitate cell migration (Abercrombie et al., 1970a–c; Laufenberger and Horwitz, 1996; Friedl and Wolf, 2003, 2010; Ridley et al., 2003; Madsen and Sahai, 2010; Sixt, 2012). Cancer cells can switch between amoeboid and mesenchymal modes of migration (referred to as plasticity), and this has been shown to be reciprocally regulated by Hic-5 and paxillin (Deakin and Turner, 2011; Gulvady et al., 2018). While fibroblasts have not been observed to exhibit a similar amoeboid mode of migration, the regulation of cancer cell plasticity by Hic-5 does support a role for this protein in modulating the type of migration a cell employs. This is likely mediated through Hic-5 RhoGTTPase-dependent modulation of actin dynamics and possibly through RhoGTTPase cross-talk with the vimentin cytoskeleton, as observed in a recent study connecting RhoGTTPase modulation of vimentin organization to integrin α6β4 signaling and cellular migration dynamics in transformed epithelial cells (Colburn and Jones, 2018).

In normal fibroblasts, RhoA–stimulated stress fiber formation is precisely balanced with Cdc42/Rac1 activation of PAK and other kinases, which promotes efficient turnover of FAs (Rane and Minden, 2014). In Hic-5 KO CAFs, we have shown that FA turnover is accelerated (Goreczny et al., 2018) and that inhibition of Cdc42 rescues cytoskeletal abnormalities observed in Hic-5 KO CAFs. Additionally, vimentin association with FAs has been shown to increase FA size and lifetime (Burgstaller et al., 2010), which suggests that Hic-5 localization to the FA may regulate FA lifetime both through its previously shown effects on FA association with actin stress fibers and through a novel function of recruiting vimentin to FAs via modulation of RhoGTTPase activity. Thus, our results support a role for Hic-5 in regulating the velocity of fibroblast migration via modulating RhoGTTPase activity and recruitment of vimentin to the cell periphery, where vimentin may interact with FAs to promote their maturation.

Here, we have identified key signaling events regulated by Hic-5 that mediate dramatic changes in vimentin organization. Nevertheless, it is also likely that physical interactions involving Hic-5 at FAs play a role in vimentin organization and F-actin–vimentin crosstalk. For example, Hic-5 could spatially promote activation of Cdc42 via its recruitment of guanine nucleotide exchange factors (GEFs) and GTTPase activating proteins (GAPs), including its binding partners βPIX/GIT/VAV2 (Yu et al., 2010). Indeed, vimentin has also been shown to interact with VAV2 (Havel et al., 2015). Alternatively, Hic-5 could serve as a physical linker of vimentin to FAs, possibly via either direct interactions with tensin (Goreczny et al., 2018) or potentially plectin1f, both of which are known to associate with fibrillar adhesions, a mature subset of FAs involved in fibronectin fibrillogenesis and thereby ECM remodeling (Burgstaller et al., 2010; Goreczny et al., 2018). However, any such additional physical interactions appear not to be essential, as pharmacologic Cdc42 inhibition rescues vimentin collapse in the absence of Hic-5 (Figures 7 and 9). Although less likely, Hic-5 could also play a role in regulating the activity or location of these signaling proteins, by binding or sequestering them in the cytosol. Last, due to vimentin collapse following acute treatment with siRNA
targeting Hic-5 expression, it is unlikely that Hic-5 regulates vimentin organization via its well-documented roles in regulation of global changes in gene expression (Heitzer and DeFranco, 2006). Future studies investigating which of the many domains of Hic-5 (Brown and Turner, 2004; Gulvady et al., 2018) are required for regulating vimentin organization would be of value in further elucidating the molecular pathways involved.

In conclusion, the data described herein identify a novel role for Hic-5 as a regulator of vimentin organization and dynamics via its modulation of RhoGTPase and formin activity. Hic-5 also regulates the switch from a migrating fibroblast (low Hic-5 expression) to a more contractile fibroblast with reduced migration and increased contraction and matrix remodeling of the ECM (high Hic-5 expression). Collapsed vimentin, associated with low levels of Hic-5 expression (KD and KO), may represent a unique cytoskeletal organization that allows a promigratory phenotype of fibroblasts through a dense ECM network where vimentin expression has been shown to impair migration velocity (Patteson et al., 2018). Similarly, the relationship between RhoGTPase-mediated regulation of formins and vimentin collapse represents a signaling axis that may also contribute significantly to the transition of fibroblasts from a migratory state to a contractile state at the site of wound healing or within the tumor stroma.

MATERIALS AND METHODS

Cell lines, reagents, chemicals, and antibodies

CAFs were isolated from tumors of Hic-5 Het (+/−) and Hic-5 KO (−/−) PyMT mice as previously described (Goreczny et al., 2017). Hic-5 Het and Hic-5 KO normal lung fibroblasts (LFs) were isolated from Hic-5 Het and Hic-5 KO mice using a slightly modified version of the protocol used to isolate CAFs (Goreczny et al., 2017) where the lungs were isolated as previously described (Seluanov et al., 2010). Human foreskin fibroblasts (HFFs) were obtained from the American Type Culture Collection (ATCC), and paxillin knockout (−/-) mouse embryonic fibroblasts (KO MEFs) (Hagel et al., 2002) were a generous gift from S. Thomas (Harvard Medical School, Cambridge, MA).

CAFs and normal LFs were cultured in a 1:1 mixture of DMEM:Ham's F12, while all other cell lines were cultured in 100% DMEM. All media were supplemented with 10% fetal bovine serum (Atlanta Biologicals; vol/vol), 2 mM l-glutamate, 10 I.U./ml penicillin, 10 µg/ml streptomycin, and 1 mM sodium pyruvate. Cells were maintained in a 37°C humidified incubator with 5% CO2, 10 µg/ml streptomycin, and 1 mM sodium pyruvate. Cells were plated onto fibronectin-coated glass coverslips (12 mm × 1.5 mm) for 4 h before fixation unless otherwise stated (cells were replated on fibronectin for subsequent analysis). KD efficiency was routinely assessed by immunoblotting and was found to be 87 ± 5% and 67 ± 5% for Hic-5 siRNA 1 and 2, respectively, and 35.5% ± 9.8% for vimentin siRNA. RNAi-mediated depletion of Paxillin (KD) in HFFs was assessed by immunofluorescence staining for Paxillin and cytoskeletal arrangement was analyzed in cells with paxillin staining (control) versus without paxillin staining (KD).

Oligomers used in this study included nontargeting control siRNA ("NT siRNA"; Dharmacon), a pool of four siRNA targeting mouse vimentin transcripts ("Vm siRNA"; Dharmacon SMARTpool ON-TARGETplus siRNA L-061596-01), one of two siRNA targeting human Hic-5 transcripts (Hic-5 siRNA 1 and Hic-5 siRNA 2; Deakin and Turner, 2011; Pignatelli et al., 2012; Gulvady et al., 2018) or siRNA targeting human paxillin transcripts (Deakin and Turner, 2011). Hic-5 siRNA oligonucleotides were purchased from Life Technologies (catalogue number AM16106) and their sequences were as follows: Hic-5–1, 5′-GGAGCUGGUAGACUGAUAUG-3′; Hic-5–2, 5′-GGACCAUGCUAGAGAUGAAG-3′. Paxillin siRNA oligonucleotides were purchased from Ambion with the following sequence: paxillin–2, 5′-UGUGGGACCCUUCUUGGUG-3′.

For overexpression experiments, cultures were transfected using Lipofectamine 3000 (Thermo Fisher Scientific) according to the manufacturer's instructions, except for the DNA amount. Cells were grown for 24 h following plating of 50,000 cells per condition and then transfected for 24 h with the appropriate construct. For vimentin collapse and migration rescue experiments, Hic-5 KO CAFs were transfected with 5 µg EGFP or EGFP-Hic-5 (Gulvady et al., 2018). For rescue of vimentin FRAP, Hic-5 KO CAFs were transfected with 4 µg mCherry-vimentin (a generous gift from Michael Davidson [Addgene plasmid # 55158; http://n2t.net/addgene:55158; RRID:Addgene_55158]) and 4 µg of either EGFP vector or EGFP-Hic-5. For assessment of microtubule dynamics, Hic-5 Het or Hic-5 KO CAFs were transfected with either 3 µg EB1-EGFP (JBD13; Addgene plasmid number: 39299; gift from Tim Mitchison and Jennifer Tnrauer, Harvard Medical School) or with both 3 µg EB1-EGFP and 3 µg enconsin microtubule binding domain–2x mCherry (EMTD-mCherry; Addgene plasmid number: 26742; Miller and Bement, 2009). Finally, cells were replated and allowed to adhere for 4 h on the appropriate substrate for subsequent downstream analyses (see below for more details).

Immunofluorescence microscopy and analysis

Cells were plated onto fibronectin-coated glass coverslips (12 × 1.5 mm) for 4 h before fixation unless otherwise stated (cells were spread for 2 or 24 h where indicated). All fibronectin-coated surfaces for cell plating were prepared by incubation with 10 µg/ml of 633 (Thermo Fisher Scientific) were used as secondary antibodies for all immunofluorescence experiments.

The following pharmacological inhibitors were also utilized: the Cdc42 inhibitor Zcl278 (Cayman Chemicals; 50 µM), the Rac1 inhibitor NSC23766 (EMD Millipore; 100 µM) and the pan-formin inhibitor SMIFH2 (Calbiochem; 30 µM).

RNAi and transfections

Hic-5 and vimentin protein expression was depleted in HFFs and CAFs, respectively, via transfection with siRNA. Cells were plated onto plastic tissue culture dishes, allowed to adhere for 24 h, and then transfected with 10 µM siRNA using RNAiMAX (Life Technologies) transfection reagent per the manufacturer's instructions. After 24 h, transfection complexes were removed by media change and cells were incubated for an additional 24 h in growth media to allow adequate depletion of transcripts and proteins. Following KD, cells were replated on fibronectin for subsequent analysis. KD efficiency was routinely assessed by immunoblotting and was found to be 87 ± 5% and 67 ± 5% for Hic-5 siRNA 1 and 2, respectively, and 35.5% ± 9.8% for vimentin siRNA. RNAi-mediated depletion of Paxillin (KD) in HFFs was assessed by immunofluorescence staining for Paxillin and cytoskeletal arrangement was analyzed in cells with paxillin staining (control) versus without paxillin staining (KD).
fibronectin diluted in phosphate-buffered saline (PBS) containing magnesium and calcium for 24 h at 4°C. Following cell adherence, cells were washed once in prewarmed PHEM buffer (pH 7.2; 120 mM PIPES, 50 mM HEPES, 20 mM ethylene glycol-bis(β-aminoethoxy) ether)-N,N′,N″,N‴-tetraacetic acid [EGTA], 4 mM magnesium sulfate) and then fixed with 4% (wt/vol) paraformaldehyde (PFA) in PBS (pH 7.2) at 37°C for 15 min. Following fixation, free aldehyde groups were quenched for 15 min with 100 mM glycine in PBS. Cells were then permeabilized and blocked by incubation in 3% bovine serum albumin (BSA) diluted in PBS containing 0.25% (vol/vol) Triton X-100 and incubated overnight at 4°C with primary antibodies diluted in blocking buffer. Next, cells were washed three times with PBS + 0.1% Tween-20 (vol/vol) and stained for 1 h with the appropriate secondary antibodies (as well as DAPI and phalloidin where indicated) diluted in blocking buffer. Finally, cells were washed three times with PBS + 0.1% Tween-20 and mounted on glass slides with Gelvatol (10% [wt/vol] polyvinyl alcohol, 20% [vol/vol] glycerol, 0.5 M Tris-HCl, pH 8.5). Cells were imaged using a Leica SP8 laser scanning confocal microscope with a Plan Apochromat 63x/1.4 NA oil objective or a Zeiss Axioskop2 plus microscope fitted with a Q-Imaging ExiBlue charge-coupled device camera using an Apochromat 40x/0.75 NA oil objective. For analyses of vimentin collapse and the presence of an F-actin hole, described below, a minimum of three independent experiments with at least 20 cells per experiment were quantified (except for normal LFs, of which a minimum of 66 cells were analyzed from two independent experiments).

Cells were analyzed for vimentin collapse and for the presence of an F-actin hole. Vimentin collapse was visually defined by two identifying characteristics: 1) collapsed vimentin had higher-intensity staining than filamentous vimentin due to bundling into closely associated vimentin filaments called thick vimentin cables (Hollenbeck et al., 1989) and 2) the collapsed vimentin occupied a smaller area within the cell than filamentous vimentin. By observing these characteristics, cells were qualitatively assessed as having one of three vimentin phenotypes: filamentous (normal distribution), partially collapsed, or collapsed vimentin. Cells with a filamentous distribution of vimentin exhibited a normal perinuclear cage of vimentin surrounding the nucleus and an array of filaments extending toward the periphery of the cell. In cells with collapsed vimentin, the network of peripheral vimentin filaments is lost and replaced by the formation of thick vimentin cables localized to the perinuclear region of the cell. Cells with partially collapsed vimentin also displayed a perinuclear localization of vimentin, but with some normal vimentin filaments remaining extended toward the cell periphery. Additionally, cells were visually assessed and classified as having a normal F-actin distribution or as having a reduced number of centrally located actin stress fibers compared with their corresponding controls (Goreczny et al., 2017). These analyses were expressed as percentages of cells with an F-actin hole.

Qualitative differences in vimentin distribution were also confirmed with the following quantitative measurements using FIJI software: a ratio of peripheral vimentin mean fluorescence intensity (MFI) to peripheral vimentin MFI and the percentage of total cell area occupied by vimentin. For regional fluorescence measurements, F-actin staining with phalloidin was used to define the total area of the cell (actin mask). Four points located at 50% of the distance from the edge of the nucleus to the outer edge of the actin mask were marked, and each point was connected to define the perinuclear region of the cell. In cells in which regions of the nucleus contacted the outer edge of the cell, the perinuclear mask was drawn to the edge of the nucleus at points of nucleus–cell membrane contact. The peripheral region of the cell was then defined as the area of a cell extending from the edge of the actin mask to the edge of the perinuclear region of the cell. The areas occupied by actin (total cell area) and vimentin were measured using the threshold function in FIJI and these values were expressed as the % total cell area occupied by vimentin: (area occupied by vimentin + area occupied by actin) x 100.

Additionally, a high-throughput automated analysis pipeline for measuring vimentin collapse was designed in Cell Profiler 3.0 to facilitate quantification of large image sets and to provide a publicly available tool for future analyses of vimentin organization. The manual analyses of vimentin collapse described above and utilized in Figure 1 was utilized to train and validate the analysis pipeline. The software was configured to identify a mask of the area occupied by the actin signal and a mask of the area occupied by the vimentin signal on a per-cell basis. The area occupied by the nucleus was removed from both masks. Adjustments in the range of fluorescence values used to define a positive signal (presence of actin or vimentin) were made on a per-experiment basis. The accuracy of mask identification was manually confirmed for each cell using the outline overlay output of the Cell Profiler pipeline. Outliers in which the analysis pipeline was not able to accurately identify the actin and vimentin masks were removed from the data analysis output. Additionally, for analysis of vimentin radial distribution in cells transfected with EGFP vector or EGFP-Hic-5, a minimum threshold of total EGFP fluorescence was set to select for EGFP-expressing cells only.

Fluorescence values for vimentin were then collected from within the vimentin mask and the radial distribution function of Cell Profiler was used to display their distribution across the total cell area (actin mask). The vimentin distribution was calculated across 16 bins defined by dividing the area from the center of the nucleus to the edge of the actin mask into 16 segments of equal area. Vimentin mean fractional intensity was used to measure the amount of vimentin within each bin. This measurement is calculated as the average fluorescence intensity per pixel at a given distance from the nucleus, normalized to the total intensity of the cell. As fluorescence is generally proportionate to target abundance, this normalized per-pixel measure corresponds to the relative protein concentration as a function of distance from the nucleus. Results were displayed as a heatmap of vimentin distribution from the Cell Profiler output and as a line graph of the average of each bin’s mean fractional vimentin intensity versus the bin distance from the nucleus. Bin sizes represent the distance of the outer edge of each bin from the center of the nucleus (µm) and were calculated on a per-condition basis for each experiment using the major axis length parameter calculated by Cell Profiler and the following formula: (major axis length + 16) x (bin number).

Inhibitor treatments
For inhibitor treatments, cells were incubated at 37°C for 2 h post-plating before treatment with ZR278, NSC23766, or SMIFH2 for 4 h as indicated. Cells treated with inhibitors for migration experiments were imaged during the 4-h treatment, while cells treated with inhibitors for immunofluorescence experiments were fixed following the 4-h inhibitor treatment. All inhibitor treatments were compared with a vehicle control, dimethyl sulfoxide (DMSO), for assessment of their cellular effects.

Triton fractionation of cell lysates
The amount of assembled filamentous vimentin versus vimentin filament precursors was assessed by fractionation of Hic-5 Het and Hic-5 KO CAF lysates with the detergent Triton X-100 (TX100). Cells were grown on fibronectin-coated plastic tissue culture dishes for 4
or 24 h before cellular protein was harvested by lysis in 1X cell lysis buffer (50 mM Tris-HCl, pH 7.2, 5 mM EGTA, 120 mM sodium chloride, 10 mM magnesium chloride, 1% TX100, 1X protease inhibitor cocktail from Roche Diagnostics, 1X PhosStop from Roche Diagnostics, and 1 mM phenylmethylsulfonyl fluoride [PMSF]). Cell lysates were centrifuged at 10,000 x g for 10 min at 4°C and the supernatant was transferred to a new tube (Triton-soluble vimentin filament precursors) and the pellet (Triton-insoluble filamentous vimentin) was resuspended in 1X cell lysis buffer with 8M urea and 1% SDS. The TX100-soluble fraction of each sample was normalized according to total protein content as determined by a Pierce bicinchoninic acid protein assay kit (Thermo Fisher Scientific) and TX100-insoluble fractions were normalized based on the soluble fractions’ protein concentration. All fractions were diluted using 4X Laemmli buffer with 2.5% (vol/vol) β-mercaptoethanol. Samples were boiled for 5 min and then run on SDS–PAGE gels with 15 µg total protein or 25% of the total lysate volume (for samples lysed directly into sample buffer, such as samples used for confirmation of knockdown) loaded per lane. Proteins were then transferred for 1 h to nitrocellulose membranes. Nitrocellulose membranes were blocked with 3% BSA in Tris-buffered saline with 0.1% Tween-20 (TBST) and incubated with primary antibodies diluted in 1.5% BSA in TBST overnight at 4°C. Next, nitrocellulose membranes were washed three times with TBST and incubated with the horseradish peroxidase (HRP)-conjugated secondary antibodies (anti-mouse IgG or anti-rabbit IgG) diluted 1:10,000 in 1.5% BSA-TBST for 1 h at room temperature. After three TBST washes, immunoblots were visualized with enhanced chemiluminescence substrate (SuperSignal West; Thermo Fisher Scientific) and imaged using a ChemiDoc MP (BioRad). Relative band intensities were measured using FIJI software and normalized to each sample’s corresponding GAPDH loading control. TX100-insoluble fractions were normalized to the GAPDH loading control of their corresponding TX100-soluble fraction.

Fluorescence recovery after photobleaching
Hic-5 Het and Hic-5 KO CAFs transfected with mCherry–vimentin with EGFP or mCherry–vimentin with EGFP/EGFP-Hic-5 and plated onto fibronectin-coated glass bottom tissue culture dishes (MatTek, No. 1.5) were utilized for live-cell total internal reflection fluorescence (TIRF) microscopy. Cells transfected with both EB1-EGFP and EMTB-mCherry were used to confirm localization of EB1 to the tips of MTFs, while cells transfected with only EB1-EGFP were used for analysis of MT dynamics using the u-track 2.2 plug-in for Matlab created by the Danuser Lab at UT Southwestern Medical Center (Jaqaman et al., 2008; Applegate et al., 2011; Ng et al., 2012). Cells were imaged 4 h postplating using a Plan Apochromat 100×/1.4 NA objective and a Nikon Eclipse TE2000-E multimode TIRF microscope equipped with an environmental chamber maintained at 37°C. During imaging, cells were maintained in live-cell imaging medium as described above. Each cell was imaged for 3–5 min at 1-s intervals for single-channel imaging and 5.5-s intervals for two-channel imaging using 488- and 550 nm-lasers. MT dynamic parameters showing a statistical difference between Hic-5 Het and KO CAFs (growth lifetime, growth length, and frequency of termination), as well as three additional key measurements of MT dynamics that did not show a statistical difference (number of nucleation events, growth speed, and dynamics) were selected from the software output to represent MT dynamics under these two conditions. Five cells each from three separate experiments (15 cells per condition) were quantified for a minimum of 150,000 EB1 tracks per condition.

Three-dimensional cell-derived matrix generation and migration analysis in two and three dimensions
Migration of Hic-5 Het and Hic-5 KO CAFs was assessed on both two-dimensional tissue culture dishes and three-dimensional cell-derived matrices (CDMs). 3D-CDMs were generated using HFFs as described previously (Deakin and Turner, 2011). For live-cell imaging and migration analyses, cells were plated for 2 h and then imaged using an HCX Plan Fluorat 10×/0.30 NA objective and a Nikon TE2000 microscope equipped with a humidified environmental chamber maintained at 37°C with 5% CO₂. Images were acquired at 10-min intervals in four or five fields per well. Inhibitor-treated cells were imaged for 4 h following addition of the inhibitor/DMSO. Cells that remained in frame and did not contact other cells throughout the duration of the movie were selected for analysis using the FIJI manual tracker plug-in to track cell movement and to calculate migration velocity and directionality of each cell.

Statistical analyses
Two-tailed unpaired Student’s t tests were performed using Excel for comparison between two experimental groups for most experiments, except where stated otherwise. One-tailed unpaired Student’s t tests were utilized to assess whether treatments with an expected direction of effect promoted either increased (formin inhibition) or decreased (Hic-5 overexpression, Cdc42 inhibition) migration velocity. For radial distribution of vimentin, the p value was calculated from a nested mixed-effects model. Both models include a fixed intercept, slope, and curvature (a parabolic fit) and an image-stratified random effect on the intercept and the slope. The nested model adds an interaction between experimental condition and curvature, allowing statistical comparison of shifts in the peak of the curve representing vimentin fluorescence radial distribution. Models were fitted using the Ime4 library and p values were derived from the analysis of variance function in both the statistical programming language and environment R (Bates et al., 2015). Data sets
were acquired from three or more independent experiments, except where otherwise indicated. *, p < 0.05; **, p < 0.01; ***, p < 0.001; and ****, p < 0.0001. Data are graphed as averages with error bars indicating SEM, except for MT dynamics data, which are graphed as a box-and-whisker plot where the bounds of the box represent the interquartile range, the horizontal line within the box represents the median, and the whiskers represent the range of the data.

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