REPORT



Cell migration orchestrates migrasome formation by shaping retraction fibers

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Migrasomes are recently discovered vesicle-like structures on retraction fibers of migrating cells that have been linked with transfer of cellular contents, shedding of unwanted materials, and information integration. However, whether and how the cell migration paradigm regulates migrasome formation is not clear. Here, we report that there are significantly fewer migrasomes in turning cells compared with straight persistently migrating cells. The major insight underlying this observation is that as the cells elongate, their rear ends become narrower, subsequently resulting in fewer retraction fibers during impersistent migration. In addition to migration persistence, we reveal that migration speed positively corelates with migrasome formation, owing to the derived length of retraction fibers. Substantiating our hypothesis, genetically removing vimentin compromises cell migration speed and persistence and leads to fewer migrasomes. Together, our data explicate the critical roles of two cell migration patterns, persistence and speed, in the control of migrasome formation by regulating retraction fibers.

Introduction

Migrasomes are recently discovered cellular organelles with vesicle-like morphology that mediate migracytosis, a cell migration-dependent mechanism for releasing cellular contents (Ma et al., 2015). It has been reported that migrasomes govern critical cellular processes including mitochondrial quality control (Jiao et al., 2021), cell-cell communication (Jiang et al., 2019), and lateral transfer of mRNA and proteins (Zhu et al., 2021). When cells crawl on extracellular substrates, retraction fibers (RFs) are pulled out of the plasma membrane from the cells' rear ends (Taylor and Robbins., 1963). Migrasomes subsequently emerge on the RFs. When cells migrate away, the RFs break, and migrasomes are left behind (Ma et al., 2015; Wu et al., 2017).

Directed migration is an important behavior of cell motility, which can be characterized quantitatively by their velocity and persistence, describing the rate of migration and the continuity in directional migration, respectively (Petrie et al., 2009; Gail and Boone, 1970). In addition to external regulation, such as chemotaxis in response to soluble cues (Andrew and Insall, 2007), durotaxis in response to mechanical signals (Lo et al., 2000), and electrotaxis in response to electric fields (Zhao, 2009), the directional cell migration can also be regulated by intrinsic clues, the fundamental mechanisms of which are associated with the Rho family of small GTPases (Raftopoulou and Hall, 2004) and the ECM receptor-integrin family (Caswell and Norman, 2006).

Previous studies revealed that several factors contributed to migrasome formation. Tetraspanin 4 (TSPAN4), abundant in the membrane of migrasomes (Zhao et al., 2019; Ma et al., 2015), together with cholesterol, are necessary and sufficient for migrasome formation (Huang et al., 2019). Correct pairing of integrins with ECM proteins determines the migrasome formation (Wu et al., 2017). A chemical screen has identified Rhoassociated coiled coil-containing protein kinase 1 (ROCK1) as a regulator for migrasome formation (Lu et al., 2020). Notably, migrasome formation strictly depends on cell migration. Blocking migration using the myosin II inhibitor blebbistatin or by modulating cell adhesions inhibits migrasome formation (Ma et al., 2015). However, how exactly distinct cell migration

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patterns contribute to migrasome formation has barely been addressed.

Vimentin is one of the most well-studied components of intermediate filaments (IFs; Ivaska et al., 2007). Recent studies have revealed that vimentin IFs interact and regulate cytoskeletal dynamics that drive cell motility. Vimentin controls actin stress fiber through GEF-H1 and RhoA (Jiu et al., 2017) and/or modulates lamellipodia formation (Helfand et al., 2011) to facilitate cell migration. Vimentin can also template microtubule networks (Gan et al., 2016) and/or strengthen focal adhesions (Bhattacharya et al., 2009) to enhance cell migration. Nevertheless, its correlation with biogenesis and function of migrasomes has not been previously reported.

In this study, we witness that cells form fewer migrasomes when making turns, and reveal that the reason is the narrower cell rear ends and fewer derived RFs during turning. In addition to persistence, cell migration speed is a critical aspect to confine migrasome formation by controlling the length of derived RFs. Depletion of vimentin IFs cause cell migration deficiency and consequently leads to fewer migrasomes, which verified our hypothesis, that migrasome formation is directly correlated with cell migration persistence and speed.

Results and discussion

There are fewer migrasomes formed when a migrating cell turns

Random migration occurs when a cell receives a nondirectional motogenic signal in the absence of any external guiding factor (Stoker and Gherardi, 1991); thus cells are not always going straight during random migration but quite often change direction by different degrees. To experimentally explore whether persistence is a key factor for cell migration-mediated migrasome formation, we started live-cell imaging experiment by examining mouse fibroblast L929 cells with stable expression of TSPAN4-GFP to label migrasomes (Fig. 1 A; and Video 1). Because migrasomes were found to grow on RFs hours after the initiation of RF formation (Wu et al., 2017), we quantified the average migrating distance of the cell (41.7 \pm 10.4 μ m; mean \pm SD) from the emergence of RFs to the appearance of migrasomes (Fig. S1, A and B; and Video 2), and decided to take regions 45 µm away from the cell rear body for measurements to ensure that migrasomes have enough time to form.

We next made an outline of this migrating cell and identified the central line of the migrating track at 240 min, when the cell completed turning and continued to migrate straight forward for >45 μ m (Fig. S1 C). The turning region (T-region, 40 × 40- μ m square) was subsequently defined (Fig. 1 A, region of interest [ROI] 1). A square of the same size, which was along the straight RFs before turning, is denoted as the persistent region (P-region), the center of which was 50 μ m away from the center of T-region to avoid overlapping (Fig. 1 A, ROI 2). It appears that there were only two migrasomes in the T-region, compared with 12 in the P-region in this example. A clue for an intuitive causal factor of this effect is RFs, where migrasomes grow. By using the ridge detection plugin in ImageJ (Steger, 1998; Schindelin et al., 2012), we were able to calculate the total RFs by counting the linear segments between each intersection, and found that there are more RFs in the P-region than in the T-region (Fig. 1 B).

We next wondered how RFs directly derived from the cell body (defined as "parental RFs") and the cell rear end (where the parental RFs grow) are reformed while the cell makes turns (Fig. 1 C). Thus, the turning phase (T-phase) was defined as cells migrating in the T-region (0–120 min), and the straightforward phase (P-phase) is the subsequent 120 min of rectilinear motion (Fig. 1 D). We identified that the cell rear end indeed became wider and parental RFs subsequently increased when the cell moved from T-phase to P-phase (Fig. 1 E), suggesting that a narrower rear end leads to fewer RFs derived from the cell body.

It is well known that cell shapes and polarity are constantly changing during migration (Bodor et al., 2020; Gómez-Moutón and Mañes, 2007; Zhong and Ji, 2013). To explain whether the morphology transformation could be the reason for the smaller rear end during turning, we defined three parameters: the distance along the migrating direction via centroid (Fig. 1 C, axis a), the distance perpendicularly to a via centroid (Fig. 1 C, axis b), and the distance from cell centroid to cell rear (Fig. 1 C, a2). Therefore, the elongation of the cell can be calculated by the ratio of a versus b, and the deformation of the cell rear can be calculated by the ratio of a2 versus a. Consistent with our speculation, this cell was more elongated, and the cell rear part became attenuated (longer and more narrow) while turning (Fig. 1 F). Together, the data from this turning cell indicate that the cell rear end and the RFs derived during cell turning are critical factors for migrasome formation.

Turning cells lead to narrower rear ends, fewer RFs, and consequently fewer migrasomes

To evaluate whether there is indeed a tight correlation between cells turning and migrasome formation, we classified cell turnings into three categories: sharp turning (ST), mild turning, and continuous turning (Fig. 2 A). Among them, ST and mild turning were defined when the turning angles are less or more than 90°, respectively. To measure the turning angles, we first performed binary processing for the TSPAN4-GFP L929 cells, then identified the centerlines of the migration tracks; the turning angles were measured based on the centerlines (Fig. S1 D).

To boost the discrepancy of migrasomes between straightforward migrating cells and turning cells, we decided to choose ST cells for further analysis and used the same strategy in Fig. 1 to define the T-region and P-region (Fig. 2, B and C). The results from numbers of ST cells further supported the conclusion that the reduced number of migrasomes results from the narrower rear ends and subsequently fewer RFs while cells change directions (Fig. 2, D–F). To further validate this finding, we observed cells cultured on a surface coated with laminin 511 (Humphries et al., 2006; Wu et al., 2017), another ECM protein, and found that there were fewer migrasomes at the T-region in ST cells (Fig. S1 E). It was noted that RFs on laminin 511 appeared not to extend as well as on fibronectin, which may due to the lower pairing of this ECM protein to specific integrins (Wu et al., 2017).

How do migration directionality and RFs regulate migrasome formation? As we observed, the location of migrasomes on RFs

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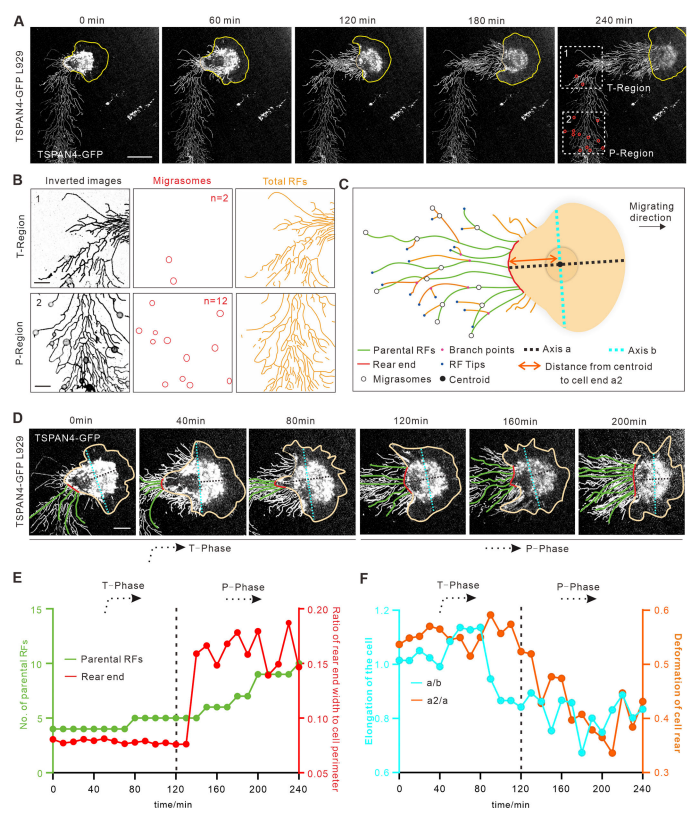


Figure 1. **Migrasomes form less when a migrating cell turns. (A)** Representative images from a time-lapse video of a L929 cell expressing TSPAN4-GFP. Yellow lines mark the outline of the cell, and white boxes indicate ROIs of T-region and P-region. Scale bar, 20 μ m. **(B)** Analysis of TSPAN4-GFP signals in T-phase and P-phase depicted in A. Images were inverted, and migrasomes and RFs were masked on the right panel. *n* in middle panel represents the number of migrasomes. Scale bars, 10 μ m. **(C)** Schematic diagram of a migrating cell. Different migration-associated components and indexes are defined in the diagram. **(D)** Enlarged time-lapse image of the cell shown in A, with rear end and parental RFs indicated as red and green lines, respectively. Axis *a* and axis *b* are indicated with cyan and black dashed lines, respectively. Scale bars, 10 μ m. **(E)** Quantification of parental RFs (green lines) and the ratio of rear end width to cell perimeter (red lines) of the migrating cell in A. **(F)** Quantification of the length ratio of *a* to *b* (cyan) and *a*2 to *a* (orange) in each time point of the migrating cell in A. Black dashed line in E and F indicates the time cutoff point between T-phase and P-phase. *n* = 25 time points.

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Cell migration controls migrasome formation

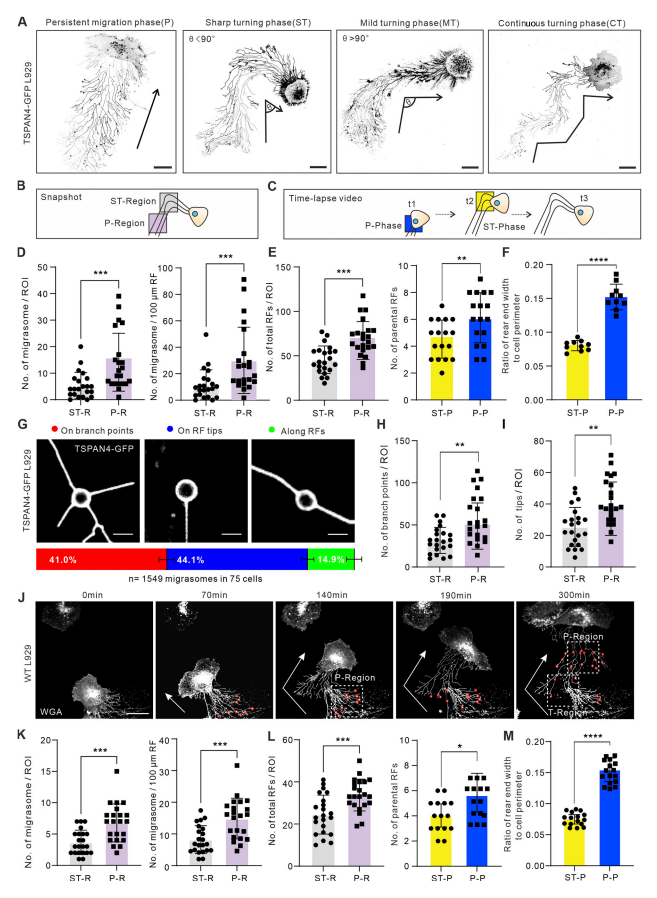


Figure 2. Cell turning leads to narrower rear ends and consequently fewer RFs. (A) Examples of different migration patterns of TSPAN4-GFP L929 cells. Black arrows indicate the migration direction. Θ indicates the turning angle. Scale bars, 20 μm. (B) Schematic diagram of the analyzed region for quantifying

the number of migrasomes (per ROI and per 100 μ m) and total RFs. ST-region, sharp turning region. (**C**) Schematic diagram of the time point for calculating the number of parental RFs and rear ends. *t*1, *t*2, and *t*3 are ordinal chronological points of the cell. (**D**) Quantification of the number of migrasomes per ROI (T-region or P-region) or per 100 μ m of RFs within the ROI in ST cells. *n* = 22 cells. (**E**) Quantification of total RFs (*n* = 22) and parental RFs (*n* = 17) within the ROI in T-phase and P-phase of the ST cell. (**F**) Quantification of the ratio of rear end width to cell perimeter within the ROIs in T-phase and P-phase of the ST cells. *n* = 10 cells. (**G**) Three types of migrasome localizations in RF networks and their corresponding proportions in TSPAN4-GFP L929 cells. Scale bars, 5 μ m. Quantification is from *n* = 1,549 migrasomes in 75 cells. (**H**) Quantification of the branch points within the defined ROIs in T-phase and P-phase of the ST cells. *n* = 22 cells. (**I**) Quantification of the tips within the defined ROIs in T-phase and P-phase of the ST cells. *n* = 22 cells. (**I**) Quantification of the tips within the defined ROIs in T-phase and P-phase of the ST cells. *n* = 22 cells. (**J**) Representative images from a time-lapse video of WT L929 cell. White arrows indicate the migration direction; red circles indicate migrasomes, and white boxes indicate ROIs of T-region and P-region. Scale bar, 20 μ m. (**K**) Quantification of the number of migrasomes per ROI (T-region or P-region), or per 100 μ m of RFs within the ROI in ST cells. *n* = 15 cells. (**L**) Quantification of the ratio of rear end width to cell perimeter within the ROI in ST cells. *n* = 12 cells. (**L**) Quantification of the number of migrasomes per ROI (T-region or P-region), or per 100 μ m of RFs within the ROI in ST cells. *n* = 12 cells. (**L**) Quantification of the number of total RFs and parental RFs within the ROI in T-phase and P-phase of the ST cell. *n* = 22 cells in left panel and *n* = 15 cells. In equivab

could be divided into three categories: on the branch points, on the tips, and along the linearized RFs (Figs. 2 G and S1 F). We measured the proportion of each case during random cell migration, and the calculation indicates that >80% of migrasomes grow de novo either on the intersections or on the tips of RFs (Fig. 2 G and Video 3). We further tested whether cell turning affects the organization of the RF network by manually quantifying the branch points and RF tips and found that sharp turnings indeed reduced the numbers of branches and tips (Fig. 2, H and I).

A recent study revealed that TSPAN4 promotes individual migrasome biogenesis (Huang et al., 2019). To exclude the effect of TSPAN4 on migrasomes, we repeated the experiments in WT L929 cells, which also form a fair amount of migrasomes. Fluorescently tagged wheat germ agglutinin was used to stain and track migrasome formation (Chen et al., 2019; Fig. 2 J and Video 4). The average migrating distance from the emergence of RFs to the appearance of migrasomes in WT L929 cells was measured (29.7 \pm 7.4 μ m; Fig. S2, A and B; and Video 5), and thus regions ≥30 µm away from the cell rear body were measured. Similar Tand P-regions were defined, and the same statistical tests were performed as in Fig. 2, D-F. Consistent with the results in TSPAN4-GFP L929 cells, migrasomes decreased during T-phase compared with straight migrating phase (Fig. 2 K), via reshaping RFs and rear ends (Fig. 2, L and M). Taken together, these data support the concept that cell migration persistence is a critical factor for migrasome formation via regulating widths of rear ends and numbers of RFs.

There are more migrasomes formed when cells migrate more persistently and faster

Cell migration speed is another critical aspect to confine migration behavior. We noticed that cell migrating speeds vary a lot even in the same Petri dish, and two straight-moving examples with slower and faster speed, respectively, are shown in Fig. 3 A (Video 6). Our measurement indicated that the speeds of straightforward-migrating TSAPN4-GFP L929 cells ranged from 15 to 80 μ m/h, with an average speed of 30.5 μ m/h (Fig. 3 B).

We then quantified the number of migrasomes in these straight-moving cells. It is interesting to note that the number of migrasomes per cell scaled up with the speed of cell migration (Fig. 3 C), yet the amount of migrasomes remained at a similar level when quantified by migrasomes derived per $100-\mu m$ length of RFs (Fig. 3 D), indicating that the changes in the number of migrasomes are associated with the length of RFs. We then compared the correlation between cell migration speed and the length of RFs and found that the faster-migrating cells generated significantly longer RFs (Fig. 3 E). In contrast, the average number of parental RFs derived from the cell body showed no direct proportion with cell migration speed (Fig. 3 F), suggesting that the influence of cell migration velocity on migrasome formation is mainly due to the relative length of derived RFs. Moreover, more branch points and tips were accompanied by longer RFs generated with faster speed (Fig. 3 G). Not surprisingly, similar results were found in WT L929 cells (Fig. 3, H–N; and Video 7). Collectively, these data suggest that migration speed is another general mechanism that is critical for migrasome formation via regulating the length of derived RFs.

Persistence and speed are critical cell migration parameters for migrasome formation

To verify that the persistence and speed of cell migration, in general, modulate the formation of migrasomes, we reproduced experiments in human gastric carcinoma cell line MGC803 and normal rat kidney cell line NRK. Migrasomes appear when cells migrate 29.7 \pm 7.4 and 28.7 \pm 6.0 µm away in MGC803 and NRK cells, respectively (Fig. S2, A and B; and Video 6). The T- and P-regions were thus defined 30 µm away from the cell rear body. Representative live-cell imaging revealed consistently that more migrasomes were formed when cells migrated more persistently and faster (Fig. 4, A, B, D, E, F, and H; and Videos 8 and 9), which is tightly associated with rear end width, number of parental RFs, and RF length (Fig. 4, C, D, G, and H). Together, results from WT L929, MGC803, and NRK provided more evidences to support the generalization of our findings.

Vimentin IFs control migrasome formation by regulating directed cell migration

The essential roles of cytoskeleton networks, including actin, microtubules, and IFs, in cell migration have been well characterized (Leduc and Etienne-Manneville, 2015; Gardel et al., 2010; Etienne-Manneville, 2013; Jiu et al., 2015). To regulate cell migration patterns by intrinsic factors and confirm whether they affect migrasome formation, we planned to manipulate the cytoskeleton system.

To find an optimal target, we first visualized the subcellular distribution of three cytoskeleton networks. It is not surprising



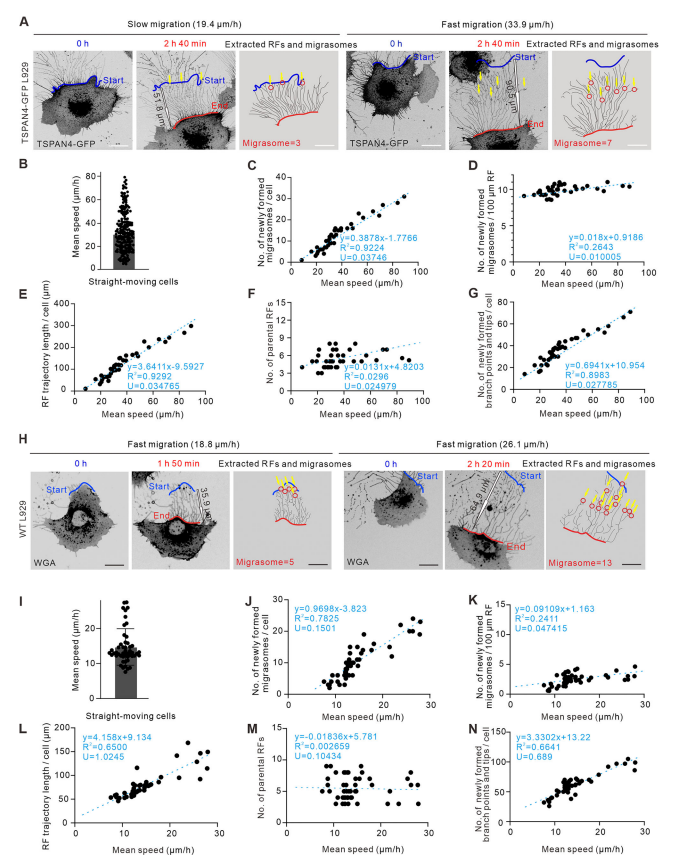


Figure 3. Migrasomes form more when cells migrating faster regardless of their migration persistence. (A) Representative images from time-lapse videos of an L929 cell expressing TSPAN4-GFP. The left panel shows an example of a slow persistent migrating cell, and the right panel shows an example of a fast persistent migrating cell; red and blue lines indicate the start and end positions of the migrating cell, respectively; yellow arrows indicate the position of

migrasome formation; and black outline arrows indicated the displacement of the migrating cell. Scale bar, 20 μm. (B) Quantification of average speed of straight-moving cells in the same culture condition. n = 200 cells. (C) Quantification of the number of newly formed migrasomes per cell in straight-moving cells. n = 39 cells. (D) Quantification of the number of newly formed migrasomes per 100- μ m RF trajectory in straight-moving cells. n = 39 cells. (E) Quantification of the RF trajectory length per cell in straight-moving cells. n = 39 cells. (F) Quantification of the number of parental RFs per cell in straightmoving cells. *n* = 39 cells. (G) Quantification of the number of newly formed branch points and tips per cell in straight-moving cells. *n* = 39 cells. The fitting lines are indicated by the blue dashed line, and the fitting equations are listed in each panel from C to G, with goodness of fit R² and uncertainty of the fitting slope U to define the correlation coefficient. (H) Representative images from time-lapse videos of WT L929 cells. The left panel shows an example of a slow persistent migrating cell, and the right panel shows an example of a fast persistent migrating cell; red and blue lines indicated the start and end position of the migrating cell, respectively; red circles indicate migrasomes; yellow arrows indicate the position of migrasome formation; and black outline arrows indicated the displacement of the migrating cell. Scale bar, 20 μm. (I) Quantification of average speed of straight-moving WT L929 cells in the same culture condition. n = 41 cells. () Quantification of the number of newly formed migrasomes per cell in straight-moving cells. n = 43 cells. (K) Quantification of the number of newly formed migrasomes per 100-μm RF trajectory in straight-moving cells. n = 44 cells. (L) Quantification of the RF trajectory length per cell in straight-moving cells. n = 39 cells. (M) Quantification of the number of parental RFs per cell in straight-moving cells. n = 41 cells. (N) Quantification of the number of newly formed branch points and tips per cell in straight-moving cells. n = 41 cells. The fitting lines are indicated by blue dashed line, and the fitting equations are listed in each panel from C-G and J-N with goodness of fit R² and uncertainty of the fitting slope U to define the correlation coefficient. The data for quantification in C–G and |-N| are from n = 3 independent experiments.

that actin filaments are localized on both RFs and migrasomes, while microtubules are mainly on migrasomes (Fig. S3, A and B). We then visualized the localization of the major IF protein vimentin by exogenous expression of vimentin-mCherry (Fig. 5 A) and endogenous staining with antibody (Fig. 5 B). In both cases, vimentin showed no localization on RFs, and more importantly, no expression on migrasomes.

Next, three independent CRISPR/Cas9 vimentin knockout (KO) lines and accordingly three full-length rescue lines were established in TSPAN4-GFP L929 cells (Fig. 5 C), and one line from each condition was chosen for further analysis. Vimentin depletion did not affect cell size and proliferation according to our measurements (Fig. S3, C-E). Importantly, the size and TSPAN4 fluorescence intensity of individual migrasomes remain comparable among WT and three KO cell lines (Fig. S3, F-H), indicating that vimentin is not involved in migrasome biogenesis per se.

To confirm that vimentin depletion shows defects in the cell migration pattern, we carried out cell random migration and wound healing assays. Straightness is an index to generally describe the turning extent of cells and was measured by the ratio between cell migration displacement and track length (Gorelik and Gautreau, 2014; Fig. 5, D and E). Turning frequency was measured by quantifying turning events per 100- μ m RFs. Depletion of vimentin significantly decreased straightness, turning frequency, and the speed of cell migration over long time lapses, which could be partially recovered with full-length vimentin (Fig. 5, D–H; Fig. S3, I–L; and Video 10). These results confirm that vimentin is involved in regulating cell migration patterns, as previously observed in other cells (Li et al., 2020; Ostrowska-Podhorodecka et al., 2021).

By snapshot quantification, there are far fewer migrasomes on the vimentin KO background regardless of whether the calculation was implemented by per cell or per 100- μ m RFs (Fig. 5, I and J). Moreover, vimentin KO cells showed fewer RFs and defective cell rear ends (Fig. 5, K and L). Together, these results show that the effective contribution for migrasome formation by vimentin depends strongly on its effect on directed cell migration.

The complexity and heterogeneity of cell migration paradigms have frustrated efforts to understand how migrasome formation is related to cell migration, even though this interconnectedness is understood to play an essential role. In this regard, our work provides far-reaching effects on future study relevant to migrasome formation (Fig. 5 M). A migration cycle is accompanied by dramatic cytoskeletal and membrane rearrangement. Previous work revealed that rear retraction is controlled by membrane tension sensing and subcellular traction force redistribution, which leads to dramatic cell shape deformation to coordinate cell migration (Hetmanski et al., 2019; Zhong and Ji, 2013). Cells become elongated during turning in our results (Fig. 1, D and F), which lead to narrower rear ends, from which RFs grow. Therefore, RFs are tightly associated with migration pattern.

Migrasomes tend to generate on stiffer RFs where TSPAN4 and cholesterol accumulate (Huang et al., 2019). The numbers of branch points and tips are mildly reduced when cells make turns (Fig. 2, G–I). It is thus tempting to speculate that impersistent migration disrupts the topology-based rigidity distribution on RFs, and TSPAN4 and cholesterol on RFs are subsequently rearranged.

Vimentin has been reported to regulate cell migration in various cell types, such as fibroblasts (Vakhrusheva et al., 2019; Sliogeryte and Gavara, 2019), cancer cells (Xuan et al., 2020; Schoumacher et al., 2010), leukocytes (Nieminen et al., 2006), and astrocytes (De Pascalis et al., 2018). Importantly, vimentin localized on neither migrasomes nor RFs compared with actin and microtubule networks (Fig. 5, A and B). Unlike microtubules and actin filaments, vimentin filaments uniquely provide cells with a hyperelastic network that regulates cell stretchability and resilience (Hu et al., 2019). Thus, it is tempting to speculate that migrasomes are stiff because of the enrichment of TSPAN4 and cholesterol (Huang et al., 2019), whereas they may lack stretchability and resilience compared with the cell body because of the absence of vimentin IFs.

Cell migration plays a key role in organ formation, tissue regeneration, wound healing, cancer metastasis, and many



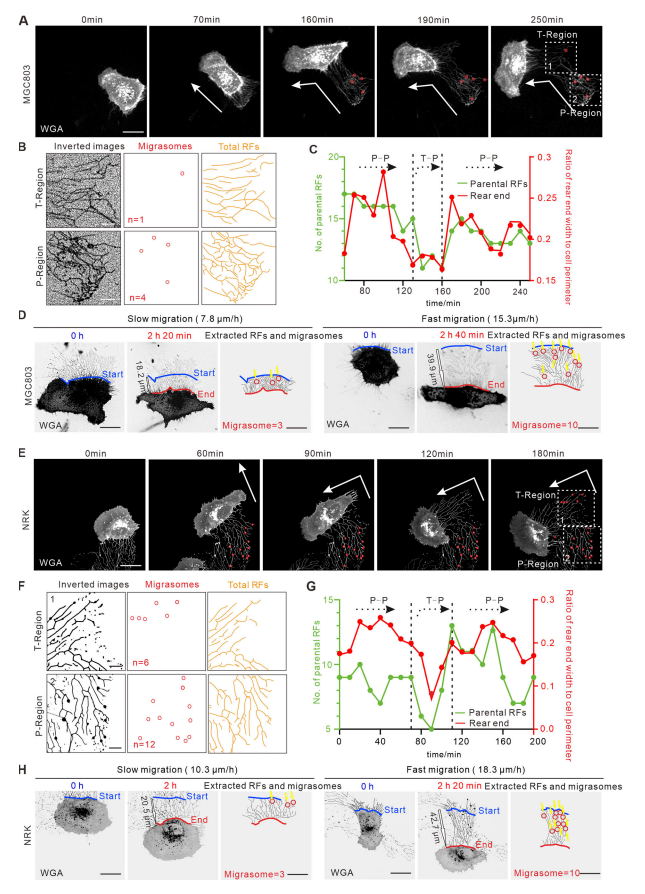


Figure 4. Persistence and speed of cell migration regulate migrasomes formation in MGC803 and NRK cells. (A and E) Representative images from a time-lapse video of WT MGC803 cell (A) and NRK cell (E). White arrows indicate the migration direction, red circles indicate migrasomes, and white boxes

indicate ROIs of T-region and P-region. Scale bar, 20 μm. (**B and F**) Analysis of RFs and migrasomes in T-region and P-region depicted in A and E, respectively. Images were inverted, and migrasomes and RFs were masked on the right. *n* in middle panel represents the number of migrasomes. Scale bars, 10 μm. (**C and G**) Quantification of the number of parental RFs (green lines) and the ratio of rear end width to cell perimeter (red lines) of the migrating cell in A and E, respectively. (**D and H**) Representative images from time-lapse videos of WT MGC803 cells (D) and NRK cells (H). The left panel shows an example of a fast persistent migrating cell. The red and blue lines indicate the start and end position of the migrating cell, respectively; yellow arrows indicated the position of migrasome formation; and black outline arrows indicated the displacement of the migrating cell. Scale bar, 20 μm.

diseases (Kurosaka and Kashina, 2008; Amini et al., 2019; Dekoninck and Blanpain, 2019; Villalobo and Berchtold, 2020). Persistent cell migration is beneficial to cell chemotaxis in tissue and organ formation, whereas impersistent cell migration allows cells, especially immune-associated cells, to better sense local environmental stimuli and signaling molecules (Worbs et al., 2017). That migration behavior controls migrasome formation potentiates a spatiotemporal dynamic regulatory function of migrasomes during various physiological and pathological processes. Our work has made a quantitative analysis of the association: that is, migration patterns affect the topology of RFs and thus affect migrasome formation, which is critical in current models of cell migration-generated migrasomes.

Materials and methods

Cell culture and transfection

Mouse fibroblast L929 cells, normal rat kidney epithelial cells (NRK cells), and human stomach (gastric) cancer MGC803 cells were kindly given by Dr. Li Yu (Tsinghua University, China). 293T cells were kindly given by Dr. Jing Zhong (Institute Pasteur of Shanghai, Chinese Academy of Sciences, China). All cells were maintained in high glucose (4.5 g/l) DMEM (06-1055-57-1A; Biological Industries) supplemented with 10% FBS (10270-106; Gibco), 100 U/ml penicillin, 100 μ g/ml streptomycin, and 4 mM L-glutamine (complete DMEM) at 37°C in a humidified atmosphere with 5% CO₂. Transient transfections were performed with JetPrime (Polyplus-transfection) according to the manufacturer's instructions using a DNA:JetOPTIMUS ratios of 1:1.25 and 24-h incubation per assay.

Vimentin CRISPR/Cas9 KO cell line and rescue cell line generation

As previously described, vimentin KO cells were generated using CRISPR/Cas9 methods (Jiu et al., 2015) based on lenti-CRISPRv2 vector (98290; Addgene) with two targets. Primers for vimentin target 1 were 5'-CACCGGAATGACTGCAGGGTGCTTT-3' and 5'-AAACAAAGCACCCTGCAGTCATTCC-3'; primers for vimentin target 2 were 5'-CACCGAGCTCCGGTTGGAGCTGGGGC-3'and 5'-AAACGCCCAGCTCCAACCGGAGCTC-3'. CRISPR clones were cultivated for 2 wk with 3 μ g/ml puromycin before selecting clones with no discernible vimentin protein expression by Western blotting. Rescue cell lines were constructed by lentivirus. Briefly, lentivirus encoding vimentin-mCherry was used to establish L929 TSPAN4-GFP vimentin KO and vimentinmCherry cell lines. 293T cells were cotransfected with the following vectors: pLVX-vimentin-mCherry, which was constructed by our lab using pLVX (125839; Addgene), HIV-1 packaging plasmid (psPAX2, 12260; Addgene), and a vesicular stomatitis virus glycoprotein expression vector (pMD2.G, 12259; Addgene). The rescue cell lines were selected by puromycin pressure.

Cell migration assay

Cells were collected from culture dishes using trypsin-EDTA and resuspended in complete DMEM at a concentration of 2×10^7 cells/ml. 20 µl cell suspension was added to the device inlet. The dish was pretreated for 3 h with 10 $\mu g/ml$ fibronectin diluted in PBS at 37°C. Cells were allowed to adhere and spread overnight. Before imaging with an Olympus IX73 inverted microscope with an UplanFL 10×/0.3 objective (Olympus), cells were incubated with Hoechst 33342 for 10 min and washed twice with PBS, and medium was replaced with complete DMEM. Images were acquired by Olympus software CellSens Dimension v1.18. Average migration velocity and directional migration duration were quantified by tracking nucleus movement using Imaris Spots plugins (Bitplane). The recording was set as every 7.5 min for 12 h. Only cells that did not collide with one another were selected for measurements. 15-µm estimated xy diameter, 30-µm maximum distance, and 3-frame maximum gap sizes were set for analyzing in Imaris.

Wound healing assay

Cells were seeded in a fibronectin-coated (10 μ g/ml) 6-well cell culture plate with a cell density of 25,000/cm² and cultivated at 37°C in 5% CO₂ overnight. Subsequently, the cell monolayers were scratched with a sterile 0.2-ml pipette tip to create linear wounds and washed with PBS to remove detached cells. Cells were incubated in complete DMEM and observed on an Olympus IX73 inverted microscope with an UplanFL 4×/0.13 objective (Olympus). Images were acquired by Olympus software CellSens Dimension v1.18. With ImageJ (National Institutes of Health), the wound areas between cells were measured and plotted at each time point. From the slope of the plot, a simple calculation yielded the cell migration rate in micrometers per hour (Jonkman et al., 2014). If we take *A* as the area of the gap and *l* as the length of the gap, the cell migration rate is given as

$$V_{\text{mingration}} = \frac{dA}{2l * dt}$$

Western blot

All cell lysates were prepared by washing the cells once with PBS and scraping them into radioimmunoprecipitation assay lysis buffer (50 mM Tris, pH 7.4, 150 mM NaCl, 1% Triton X-100, 1% sodium deoxycholate, and 0.1% SDS) supplemented with 1 mM PMSF, 10 mM DTT, 40 µg/ml DNase I, and 1 µg/ml of leupeptin,



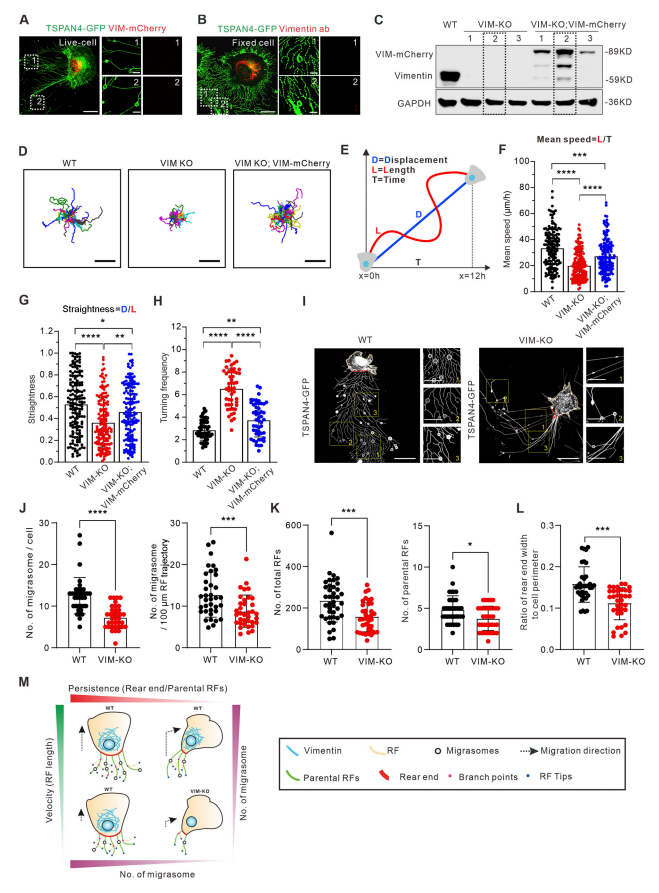


Figure 5. Vimentin regulates migrasome formation by modulating cell migration velocity and persistence. (A) Representative snapshots of live cells show the subcellular distribution of exogenously expressed vimentin-mCherry and TSPAN4-GFP in L929 cells. The white dotted outlines represent ROIs

containing RFs and migrasomes. Scale bars, 10 and 2 μ m in image and magnified view, respectively. (**B**) Representative immunofluorescence images show the subcellular distribution of endogenous vimentin with respect to TSPAN4-GFP. The white dotted outlines represent ROIs containing RFs and migrasomes. Scale bars, 10 and 2 μ m in image and magnified view, respectively. (**C**) Western blot analysis of the extracts of WT, VIM-KO, and VIM-KO; VIM-mCherry TSPAN4-GFP expressing L929 cells. The dashed line indicates the cell line we chose for further experiments. (**D**) Cell random migration trajectory of WT, VIM-KO, and VIM-KO; VIM-mCherry TSPAN4-GFP-expressing cells within 12 h. *n* = 15 cells. Scale bars, 100 μ m. (**E**) Schematic diagram for the quantification of cell migration speed and straightness. (**F**) Quantification of the mean speed of TSPAN4-GFP WT, VIM-KO, and VIM-KO; VIM-mCherry cells. *n* = 150 cells. (**G**) Quantification of the straightness of TSPAN4-GFP WT, VIM-KO, and VIM-KO; VIM-mCherry cells. *n* = 50 cells. (**I**) Snapshots of live TSPAN4-GFP-expressing WT and VIM-KO L929 cells. Yellow squares indicate the magnified fields, and red lines mark the rear end. Scale bars, 20 and 10 μ m in image and magnified view, respectively. (**J**) Quantification of the number of migrasomes per cell and per 100- μ m RF trajectory in TSPAN4-GFP WT and VIM-KO L929 cells. *n* = 36 cells. (**K**) The number of total RFs and parental RFs in TSPAN4-GFP-expressing WT and VIM-KO L929 cells. *n* = 36 cells. (**M**) Schematic model of cell migration pattern to orchestrate migrasome formation. The data for quantification in F–H and J–L are from *n* = 3 independent experiments. The data are presented as mean \pm SD (*t* test in J–L; one-way ANOVA in I–K). *, P < 0.05; **, P < 0.01; ***, P < 0.001; ****, P < 0.0001. Source data are available for this figure: SourceData F5.

pepstatin, and aprotinin. All preparations were conducted at 4°C. Protein concentrations were determined with BCA Protein Assay kit (P0012; Beyotime) and equal amounts of total cell lysates were mixed with Laemmli sample buffer (LSB), boiled, and run on 12.5% SDS-PAGE gels. Proteins were transferred to nitrocellulose membrane with Trans-Blot Turbo transfer system (Bio-Rad) using Mini TGX gel transfer protocol. Membrane was blocked in 5% BSA for 1 h at RT. Primary and secondary antibodies were diluted into fresh blocking buffer overnight at 4°C and 1 h at RT, respectively. Proteins were detected from the membranes with SuperSigna West Femto Maximum Sensitivity Substrate (Thermo Fisher Scientific). Antibodies were used with the following dilutions in primary antibody dilution buffer (PO023A; Beyotime): vimentin rabbit monoclonal D21H3 antibody (dilution 1:1,000; 5741; Cell Signaling Technology), GAPDH rabbit polyclonal antibody (dilution 1:5,000 for WB; G9545; Sigma-Aldrich). HRP-linked anti-rabbit IgG antibody (dilution 1:5,000; 7074V; Cell Signaling Technology). Chemiluminescence was measured after resin with Western blotting ECL (180-501; Tanon).

Immunofluorescence microscopy

Cells were fixed with methanol for 5 min at -20° C, washed three times with PBS, and permeabilized with Saponin in PBS for 5 min. Cells were then blocked in PBS supplemented with 5% BSA. Vimentin rabbit monoclonal D21H3 antibody (dilution 1:200; 5741; Cell Signaling Technology) and Alexa Fluor 568 goat anti-rabbit IgG (H + L; dilution 1:1,000; A11011; Invitrogen) were applied to cells and incubated at RT for 2 h. All images were acquired by Olympus software CellSens Dimension v1.18 on an Olympus SpinSR10 Ixplore spinning disk confocal microscope equipped with an UplanApo 100×/1.5 oil objective (Olympus). The pixel size was optimized to achieve the maximum resolution, which was calculated to be 65 nm.

Live-cell imaging

For live-cell imaging, 35-mm glass-bottom dishes (MatTek Corp.) were coated with 10 μ g/ml fibronectin (PHE0023; Gibco) in PBS for \geq 3 h at 37°C, washed with PBS twice, and immersed in complete DMEM without phenol red (01-053-1A; Biological Industries) before seeding of cells. For labeling actin and microtubule cytoskeleton, cells were incubated with 0.2 μ M SiR-Actin (CY-SC001; Cytoskeleton) and SiR-Tubulin (CY-SC002; Cytoskeleton) for 6 h, respectively.

Time-lapse images of cells with transient transfection were acquired with the Olympus CellSens Dimension system, consisting of an Olympus SpinSR10 Ixplore spinning disk confocal microscope and a Yokogawa CSU-W1 confocal scanner. Appropriate filters, heated sample environment (37° C), controlled 5% CO₂, and UplanApo 100×/1.5 oil objective (Olympus Corp.) were used. The recording was set as every 10 min for 12 h, and one focal plane was recorded for all live-cell videos.

Statistical analysis

Statistical data analyses were performed with Excel (Microsoft) and Prism v8 (GraphPad Software). For the data with normal distribution, Student's two-sample unpaired *t* test was used. If data did not follow a normal distribution, Mann–Whitney *U* test for two independent samples was conducted. One-way ANOVA followed by a Tukey's post hoc test was used to evaluate differences between three or more groups. Statistical analysis was obtained with *t* test or *U* test for Fig. 2, D–F, H, I, K–M; Fig. 5, J–L; and Fig. S3 E and with one-way ANOVA for Fig. 5, F–H; and Fig. S3 C, G, H, J, and L. Significance is indicated by asterisks: *, P < 0.05; **, P < 0.01; ***, P < 0.001; ****, P < 0.0001.

For analyzing the number of migrasomes, parental RFs, branch points and tips, we manually counted in the defined ROI views or cells. The position of centroid is the geometric center of the cell and equal to the arithmetic mean position of the uniformly distributed imaging pixel points within the spreading area of the cell. Therefore, the position of the centroid of the cell can be calculated with the following formula:

$$x_{c} = \frac{x_{1} + x_{2} + \dots + x_{n}}{n},$$

 $y_{c} = \frac{y_{1} + y_{2} + \dots + y_{n}}{n},$

where (x_c, y_c) are the coordinates of the centroid of the cell, *n* is the number of pixel points in the cell, and (x_i, y_i) are the coordinates of the ith pixel within the cell.

For analyzing the number of migrasomes per 100 μ m, migrasome number ($N_{\rm migrasome}$) and parental RFs number ($N_{\rm P-RF}$) per ROI in snapshot or per cell in live cells were manually counted. The average length of parental RF trajectories (L; μ m) per ROI in snapshot or per cell in live cells were calculated by "line measurement" in ImageJ. This gave us the number of migrasomes per 100- μ m RFs as No. of migrasomes per 100 μ m = $\frac{N_{\text{migrasome}} \times 100}{L \times N_{\text{P-RF}}}$.

For analyzing the number of total RFs, "freehand selection" in ImageJ was used to identify the ROI of RFs, and appropriate line width was determined (5 pixels in this case) to make sure that all RFs were recognized. The numbers of recognized linear fragments between intersections of RFs were then calculated by "rigid detection" plugin in ImageJ (Steger, 1998) to obtain the number of total RFs.

For analyzing the ratio of rear end width versus cell perimeter, "free-hand line" of ImageJ was used to manually outline the rear ends and the cell perimeter, and the lengths were subsequently measured.

To estimate the linear regression in Fig. 3, C–G and J–N, we used "regression analysis" in Microsoft Excel (Office16) to fit the data with the linear equation: $y = ax \pm b$. *a* indicates the slope. The calculated fitting equation, goodness of fit (R^2), and uncertainty of the fitting slope (U) are indicated in the figures. R^2 tending to 1 indicates a better linear correlation between two sets of variables. The uncertainty of the slope was reflected by the range of slope fluctuation U, indicating the confidence interval for the slope ($a \pm U$).

Online supplemental material

Fig. S1 shows details on the data analysis. Fig. S2 shows the quantification of cell displacement from RF formation to migrasome appearance in WT L929, MGC803, and NRK cells. Fig. S3 shows that vimentin deficiency leads to defective cell migration and abnormal migrasome formation. Video 1 shows a turning TSPAN4-GFP L929 cell. Video 2 shows a migrating TSPAN4-GFP L929 cell, which indicates the average time from the emergence of RFs to the appearance of migrasomes. Video 3 shows migrasome formation in TSPAN4-GFP L929 cell-derived RFs. Video 4 shows a turning WT L929 cell. Video 5 shows cell displacement from RF formation to migrasome appearance in WT L929, MGC803, and NRK cells. Video 6 shows two persistent migrating (slow and fast) TSPAN4-GFP L929 cells. Video 7 shows two persistent migrating (slow and fast) L929 cells. Video 8 shows a turning MGC803 cell and a turning NRK cell. Video 9 shows two persistent migrating (slow and fast) MGC803 and NRK cells, respectively. Video 10 shows cell random migration in WT, VIM-KO, and VIM-KO; VIM-mCherry TSPAN4-GFP expressing cells.

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Supplemental material

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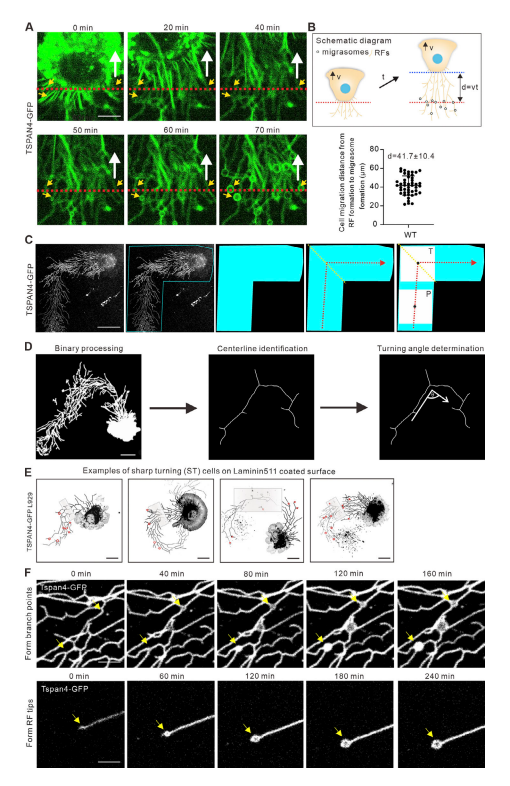


Figure S1. Data analysis details. (A) Representative images from a time-lapse video indicate the time from RF derivation to migrasome formation in TSPAN4-GFP L929 cells. The white arrow indicates the direction of cell migration, the red dashed line denotes the position of cell rear at initial time point, and the yellow arrows indicated the position of migrasome appearance. Scale bars, 10 μm. (B) Schematic diagram and quantification of cell displacement from RF formation to migrasome appearance. Data are presented as mean \pm SD; n = 48 cells. (C) Process diagram to determine the ROI of T-phase and P-phase in a sharp-turning TSPAN4-GFP L929 cell. Blue region marks the ROI of turning cell, yellow lines mark the position of turning, red lines mark the trajectory of migration, black dots mark the center of squares, and white squares mark the ROI of T-phase and P-phase. Scale bars, 20 µm. (D) Schematic diagram of the steps to define the turning angle. Scale bars, 20 µm. (E) Examples of sharp-turning TSPAN4-GFP L929 cells grown on laminin 511-coated surface. Gray boxes indicate the T-regions, and red circles indicate the migrasomes. Scale bars, 20 μm. (F) Representative images from time-lapse videos in TSPAN4-GFP L929 cells visualizing the migrasome formation process on branch points and tips of RFs. Yellow arrows indicate the position where migrasomes formed. Scale bars, 10 µm.

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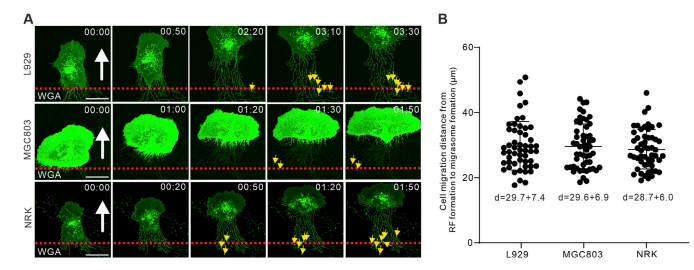


Figure S2. **Cell migration displacement from RF formation to migrasome appearance.** (A) Representative images from time-lapse videos indicate the time from RF derivation to migrasome formation in WT L929, MGC803, and NRK cells. The white arrows indicated the direction of cell migration, the red dashed lines denote the position of cell rear at initial time point, and the yellow arrows indicate the position of migrasome appearance. Scale bars, 20 μ m. (B) Quantification of cell displacement from RF formation to migrasome appearance. Data are presented as mean ± SD; *n* = 54 cells in L929, 53 cells in MGC803 and 53 cells in NRK.

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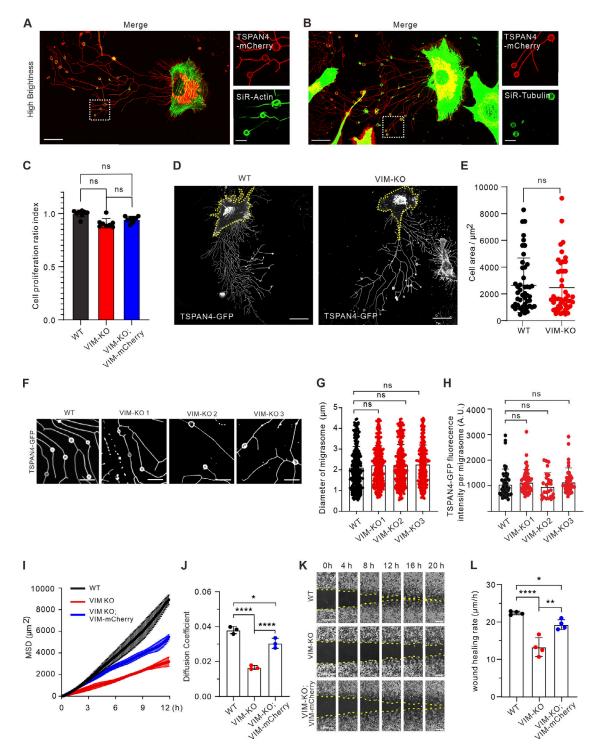


Figure S3. Vimentin deficiency leads to defective cell migration and abnormal migrasome formation. (A) Representative images from a time-lapse video of TSPAN4-mCherry L929 cells incubated with SiR-actin. Magnified panels are the white box in the left panel. Scale bars, 20 μ m. (B) Representative images from a time-lapse video of TSPAN4-mCherry L929 cells stained with SiR-tubulin. Magnified panels are the white box in the left panel. Scale bars, 20 μ m. (C) Analysis of CCK8 assay in TSPAN4-GFP WT, VIM-KO, and VIM-KO; VIM-RES cells. (D) Examples of snapshots of live TSPAN4-GFP WT and VIM-KO L929 cells. Yellow dashed lines indicate the cell outlines. Scale bars, 20 μ m. (E) Quantification of the cell area in TSPAN4-GFP WT and VIM-KO L929 cells. *n* = 47 cells. (F) Examples of RF-associated migrasomes in TSPAN4-GFP WT and VIM-KO L929 cells. *n* = 343 migrasomes in 23 cells. (H) Quantification of the TSPAN4-GFP intensity per migrasome in TSPAN4-GFP WT and VIM-KO L929 cells. *n* = 50 migrasomes in 11 cells. (I) Mean square displacement (MSD) analysis of representative trajectories per condition. *n* = 3 regions including 15 trajectories. (K) Examples of wound healing assay in TSPAN4-GFP WT, VIM-KO, and VIM-RS cells. Scale bars, 100 μ m. (L) Quantification of wound healing migration rate per condition. *n* = 4 independent experiments. The data for quantification in C, E, and G–L are from *n* = 3 independent experiments. Data are presented as mean ± SD (*t* test in E; one-way ANOVA in C, G, J, and K). *, P < 0.05; **, P < 0.001.

Video 2. **Time-lapse video of a migrating TSPAN4-GFP L929 cell, which indicates the distance from the emergence of RFs to the appearance of migrasomes.** Duration of the video is 100 min, and the recording time interval is 10 min/frame. The display rate is 4 frames/s. Scale bar, 20 μm.

Video 3. **Time-lapse videos of migrasome formation in TSPAN4-GFP L929 cell-derived RFs.** Migrasomes tended to generate on branch points and tips. Duration of the video is 240 min, and the recording time interval is 10 min/frame. The display rate is 5 frames/s. Scale bar, 10 μm.

Video 4. **Time-lapse video of a turning WT L929 cell.** Duration of the video is 300 min, and the recording time interval is 10 min/frame. The display rate is 4 frames/s. Scale bar, 20 µm.

Video 5. **Time-lapse video of migrating L929, MGC803, and NRK cells, which indicate the distance from the emergence of RFs to the appearance of migrasomes.** Duration of the videos are 200, 110, and 100 min, respectively. The recording time interval is 10 min/frame. The display rate is 4 frames/s. Scale bar, 20 µm.

Video 6. **Time-lapse video of two persistent migrating (slow and fast) TSPAN4-GFP L929 cells.** The recording time interval is 7.5 min/frame. The red line indicates the start position, and the blue line indicates the end position. Yellow arrows indicate the position of migrasome formation. The mean speed during the recording time is labeled above the images. The display rate is 6 frames/s. Scale bar, 20 µm.

Video 7. **Time-lapse video of two persistent migrating (slow and fast) L929 cells.** The recording time interval is 10 min/frame. The red line indicates the start position, and the blue line indicates the end position. Yellow arrows indicate the position of migrasome formation. The display rate is 4 frames/s. Scale bar, 20 µm.

Video 8. **Time-lapse video of a turning MGC803 cell and a turning NRK cell.** Duration of the video is 250 min and 190 min, respectively. The recording time interval is 10 min/frame. The display rate is 4 frames/s. Scale bar, 20 µm.

Video 9. **Time-lapse video of two persistent migrating (slow and fast) MGC803 cells and NRK cells.** The recording time interval is 10 min/frame. The red line indicates the start position, and the blue line indicates the end position. Yellow arrows indicate the position of migrasome formation. The display rate is 4 frames/s. Scale bar, 20 µm.

Video 10. **Time-lapse video of cell random migration in WT, VIM-KO, and VIM-KO; VIM-mCherry TSPAN4-GFP-expressing cells within 12 h.** Duration of the video is 720 min, and the recording time interval is 10 min/frame. Color-coded bar from blue to red indicates the speeds ranging from 0 to 80 μm/h. The display rate is 7 frames/s. Scale bar, 100 μm.