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# Functional integrity of the contractile actin cortex is safeguarded by multiple Diaphanous-related formins

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The contractile actin cortex is a thin layer of filamentous actin, myosin motors and regulatory proteins beneath the plasma membrane crucial to cytokinesis, morphogenesis and cell migration. However, the factors regulating actin assembly in this compartment are not well understood. Using the Dictyostelium model system, we show that the three Diaphanous-related formins (DRFs) ForA, ForE and ForH are regulated by the RhoA-like GTPase RacE and synergize in the assembly of filaments in the actin cortex. Single or double formin-null mutants displayed only moderate defects in cortex function whereas the concurrent elimination of all three formins or of RacE caused massive defects in cortical rigidity and architecture as assessed by aspiration assays and electron microscopy. Consistently, the triple formin- andRacE-mutants encompassed large peripheral patches devoid of cortical F-actin and exhibited severe defects in cytokinesis and multicellular development. Unexpectedly, many forA<sup>-</sup>/H<sup>-</sup>/E<sup>-</sup> and racE<sup>-</sup>-mutants protruded efficiently, formed multiple exaggerated fronts and migrated with morphologies reminiscent of rapidly-moving fish keratocytes. In 2D-confinement, however, these mutants failed to properly polarize and recruit myosin II to the cell rear essential for migration. Cells arrested in these conditions displayed dramatically amplified flow of cortical actin filaments, as revealed by TIRF-imaging and iterative particle image velocimetry (PIV). Consistently, individual and combined, CRISPR/Cas9-mediated disruption of genes encoding mDia1 and -3 formins in B16-F1 mouse melanoma cells revealed enhanced frequency of cells displaying multiple fronts, again accompanied by defects in cell polarization and migration. These results suggest evolutionarily conserved functions for formin-mediated actin assembly in actin cortex mechanics.

actin cortex | formin | RhoGTPase | cell migration | cytokinesis

## Introduction

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The actin-rich cell cortex is required for cell shape remodeling in fundamental cellular processes such as cytokinesis, morphogenesis and cell migration (1). Cell motility is regulated by polarization, adhesion and cytoskeletal activities leading to sitespecific force generation, as exemplified by leading edge actin assembly and myosin-dependent rear contraction (2-4). Based on considerable variations of these activities in different cell types, this process is further subdivided into mesenchymal and amoeboid types of migration as two extremes of a wide spectrum (5). The slow mesenchymal type of motility is characterized by strong substrate adhesion and formation of prominent stress fibers as well as a protruding lamellipodium at the front (6), whereas fast amoeboid migration as exemplified by Dictyostelium cells is defined by weaker and more transient adhesions, a rounder cell shape, actin-rich protrusions or blebs in the front and myosindriven contraction in the rear (7, 8). However, migration and other processes involving cell shape remodeling as e.g. cytokinesis also require a thin, actin-rich cortex below the membrane.

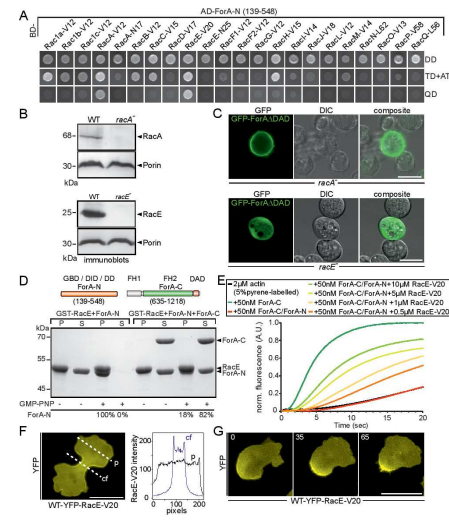
This cortex contains actin, myosin and associated factors assembling into a multi-component layer (9, 10), which is intimately linked to the membrane in a PI(4,5)P2-dependent manner by the ezrin, radixin and moesin (ERM)-family of proteins in animal cells (11, 12) and cortexillin (Ctx) in *Dictyostelium* (13–15). The function of this thin actin meshwork is comparable to cell walls in plants, yeast and bacteria, as it defines the cell's stiffness, resists external forces and counteracts intracellular, hydrostatic pressure (9, 16). However, as opposed to the static cell wall of plants and bacteria, the actin cortex of amoebae and animal cells has viscoelastic properties that can be remodeled in the timescale of seconds. Rapid F-actin rearrangements enable cells to promptly modify their shapes for fast adaptation to changes in extracellular environment (9, 16). Moreover, and as opposed to cells with rigid cell walls encaging them entirely, cell cortex constituents of motile eukaryotic cells are organized in gradients due to the asymmetry of positioning signals (17).

The physical properties of the cell cortex such as its tension and contractility likely impacting on plasma membrane dynamics are regulated by myosin motor activity as well as the arrangement and density of F-actin networks generated by distinct actinassembly machineries (9). In cells, actin polymerization is mostly initiated by Arp2/3 complex and formins (18). The Arp2/3 complex creates branches at the sides of preexisting mother filaments and generates a dense actin meshwork at the front of migrating cells (18, 19). Formins instead nucleate and elongate long and linear actin filaments (19). A major subgroup of the formin family is comprised by Diaphanous-related formins (DRFs), which are autoinhibited due to intramolecular interactions of the Diaphanous

#### Significance

The actin-rich cell cortex is a viscoelastic structure participating in a variety of cellular processes. However, the complete inventory of actin assembly factors driving its formation and knowledge about their specific contributions is still incomplete. We show here that functional integrity of the cell cortex in *Dictyostelium* and mammalian cells is backed up by multiple Diaphanous-related formins that are regulated by Rhosubfamily GTPases. These DRFs contribute to the generation of long actin filaments of the contractile actin cortex and are required for cell mechanics. Of note, these factors are excluded from Arp2/3 complex-nucleated networks, implying diversification of the cortex into functional subcompartments to segregate cortical actomyosin contraction in the rear or cleavage furrow ingression from actin-based protrusion in the front.

**Reserved for Publication Footnotes** 

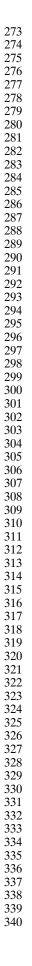


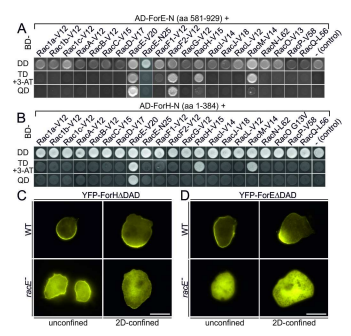
**Fig. 1.** ForA is regulated by the RhoA-homologue RacE. (*A*) The N-terminal domain of ForA (ForA-N) containing the GBD interacts specifically with the activated forms of RacA (RacA-V12) and RacE (RacE-V20) in the Y2H assay. Yeast was transformed with the indicated constructs and selected for the presence of prey and bait plasmids by growth on double-dropout (DD) media lacking leucine and tryptophan. Interactions were assayed by growth on stringent triple-dropout (TD) media additionally lacking histidine in the presence of 3 mM 3-AT and on quadruple-dropout (QD) media additional lacking histidine and adenine. AD, Gal4-activation domain; 3-AT, 3-amino-1,2,4-triazole; (*B*) Genetic elimination of RacA and RacE was confirmed by immunoblotting. Porin was used as a loading control. (C) Constitutively active ForA fused to GFP requires RacE for targeting to the cell cortex but localizes appropriately in the absence of RacA. Scale bars, 10 µm. (*D*) ForA constructs used for biochemical analyses. GBD, GTPase-binding domain; DID, diaphanous inhibitory domain; DD, dimerization domain; FH, formin homology domain; DAD, diaphanous autoinhibitory domain. Active RacE interacts directly with ForA-N and was able to partially release ForA-N from the autoinhibited ForA-N/ForA-C complex. GST-pulldown experiments with GMPPNP-loaded RacE are shown. (P) pellet; (S) supernatant. The numbers below indicate the relative amounts of ForA-N in P and S fractions. (*E*) Active RacE releases autoinhibition of the catalytically inactive ForA-N/ForA-C complex to promote actin assembly in pyrene assays in a concentration-dependent manner. (*F*) Active RacE N-terminally fused to YFP accumulates about 2-fold in the cell cortex of the cleavage furrow in 2D-confinement under agar. Abbreviations: cf, cleavage furrow; p, pole. Scale bar, 20 µm. (G) Images from lime-lapse movies correspond to Movie S1 and show that active RacE is enriched in the rear cell cortex of a polarized cell migrating under agar. Scale bar, 20 µm.

inhibitory domain (DID) with the Diaphanous autoregulatory domain (DAD) (20). DRF autoinhibition is commonly released by binding of activated Rho-family GTPases (21, 22), but can also be driven by Ras (23). As yet, both Arp2/3 complex and formins have been implicated in the generation of cortical actin in different cell types (24, 25). However, the precise quantitative contributions of Arp2/3 complex- and formin- generated filaments to this structure and their interplay in cortical functions are still elusive. Depletion of the formin mDia1 (Diaph1) in HeLa cells led to failure of cortex function in mitotic cell division, while depletion of Arp2/3 complex alone did not (24). Interestingly, the same study reported Arp2/3 complex inhibition to potentiate effects of mDia1 depletion, suggesting synergistic activities of mDia1 and Arp2/3 complex in the nucleation of cortical actin (24). AFM measurements indicated that cortical elasticity in HeLa and M2 

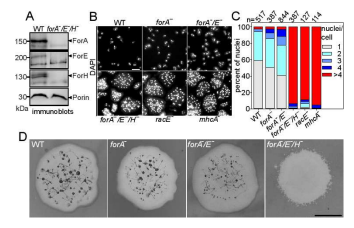
melanoma cells is mostly affected by the formin inhibitor SmiFH2 (25). In contrast, reducing Arp2/3 complex activity by CK666 did not appear to play a critical role (25). These data suggested central roles for formins in cell cortex mechanics, but need to be complemented by genetics, last not least to identify specific, contributing formins.

Previously, we established that the mDia1-related formin ForA in *Dictyostelium* cells prevents blebbing in the rear to assist protrusion at the front, in particular under mechanical stress (18). Here, we identify and characterize two additional mDia-related, cortical *Dictyostelium* formins, ForE and ForH, which synergize with ForA to safeguard cortex-dependent functions. Moreover, we extend our studies to mammalian formins mDia1 and -3, providing conclusive evidence that comparable pathways operate in higher eukaryotes.





**Fig. 2.** Active RacE interacts with two additional cortical formins. (*A-B*) ForE-N and ForH-N interact specifically with the active form of RacE (V20) in the Y2H assay. Yeast was transformed with the indicated constructs and selected for the presence of prey and bait plasmids by growth on double-dropout (DD) media lacking leucine and tryptophan. ForE-N additionally showed strong interaction with active RacF2 (V12). Interactions were scored by growth on stringent triple-dropout (TD) media in the presence of 3 mM 3-AT or quadruple-dropout (QD) media as outlined in Fig. 2. Both formins showed no genetic interaction using the dominant-negative RacE (N25) variant or empty AD plasmids as negative controls. AD, Gal4-activation domain; 3-AT, 3-amino-1,2,4-triazole; BD, Gal4-binding domain. (*C-D*) Constitutively active ForF and ForE localize in the cell cortex and rear of migrating WT cells. YFP-tagged variants of the formins were expressed in WT and *racE*<sup>-</sup> cells and analyzed by wide-field fluorescence microscopy at the conditions indicated. Scale bars, 10 µm.



**Fig. 3.** Elimination of all three cortical formins is detrimental for cell division and development. (A) Inactivation of the *forA*, *forE* and *forH* genes in the triple knockout mutant was verified by immunoblotting using specific formin sera. Porin was used as a loading control. (B) WT and the mutant cells indicated were grown for 48 h in shaken suspension at 150 rpm, subsequently seeded on glass cover slips, fixed and stained with DAPI to visualize the nuclei. Scale bar, 20 µm. (C) Quantification of nuclei in cells as shown in (B). n, number of analyzed cells. (D) Coincident elimination of ForA, ForE and ForH blocks development. WT or formin-deficient cells were transferred with a tooth pick onto a lawn of *K. aerogenes* on non-nutrient agar plates and monitored after 96-120 h of development. Scale bar, 0.5 mm.

#### Footline Author

#### Results

Active ForA accumulates in the cleavage furrow of dividing cells. Consistent with the mechanistic similarities between migration and cytokinesis, many proteins accumulating in the trailing edge, as for instance myosin II, cortexillin (Ctx), the functional homologue of ERM proteins in Dictyostelium, and IQGAPs, were also found in cleavage furrows and are known to regulate cytokinesis (26-28). Thus, we explored the localization of active ForA at different stages of the cell cycle. Active ForA is uniformly localized in the cell cortex of unpolarized interphase cells (17). In mitotic cells, active ForA remained evenly distributed in the cell cortex up to early anaphase, but subsequently began to relocalize to the cleavage furrow like IQGAP1 and Ctx I ("SI Appendix, Fig. S1 A and B"), strongly suggesting a critical function of formingenerated cortical actin in cytokinesis, as previously shown in a variety of cell types (29). However, when cultivated in petri dishes allowing adhesion of cells to the substratum, or even when exposed to high shear forces in shaken suspension culture, forAcells exhibited negligible defects in cytokinesis ("SI Appendix Fig. S1 C and D"). Of note, we have previously shown that either Ctx I and II or IQGAP1 and IQGAP2 had to be eliminated simultaneously to cause strong defects in cytokinesis, while single knockout mutants exhibited no or minor defects (28). Thus, the lack of a cytokinesis defect in forA cells suggested functional overlaps of ForA with one or multiple other cortical formins to safeguard this critical cellular function. In line with this view, cortical F-actin is still present in contractile regions of forA<sup>-</sup> cells, such as the trailing edge (17).

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369 ForA interacts with active form of the RhoA homologue RacE. 370 DRFs such as ForA are commonly assumed to be activated by 371 GTP-bound Rho-family GTPases. Dictyostelium cells lack canon-372 ical Cdc42 and Rho homologues, but express 20 Rac proteins, 373 some of which exert characteristics of Cdc42 and RhoA func-374 tions. Since appropriate ForA targeting and activation requires 375 concurrent interactions with PI(4,5)P2 and an active GTPase (17), 376 we employed yeast-two-hybrid (Y2H) analyses to systematically 377 screen all 20 Dictvostelium Racs for interaction with the N-378 terminal domain of ForA encompassing the GBD. Under the 379 most stringent growth conditions on selective media, ForA ge-380 netically interacted with constitutively active RacA and RacE, 381 while it failed to interact with dominant-negative variants of these 382 GTPases (Fig. 1A). RacA has not yet been characterized, but 383 contains a BTB domain at its C-terminus and lacks a classical 384 CAAX motive required for prenylation (30). Consistently, ectopi-385 cally expressed RacA fused to GFP localized ubiquitously in the 386 cytoplasm and was not enriched at the cell cortex (Fig. S2). Thus, 387 it appeared unlikely that it regulates recruitment and activation of 388 ForA at the cortex. RacE instead was previously implicated in reg-389 ulation of cortical tension and cleavage furrow progression (31). 390 Since Y2H analyses can occasionally generate ambiguous results, 391 we sought to corroborate these findings in an independent assay. 392 To this end, we generated genetic knockout cell lines devoid of 393 RacA and RacE in the AX2 wild type (WT) strain (Fig. 1B). Then, 394 we monitored localization of constitutively active ForA fused to 395 GFP in these knockouts. As shown in Fig. 1C, ForA was normally 396 targeted to the cell cortex in racA<sup>-</sup> cells, but failed to localize 397 to the cortex in *racE*<sup>-</sup> cells, strongly suggesting a physiologically 398 interaction between ForA and RacE. Consistently, the purified 399 N-terminus of ForA also physically interacted with GMP-PNP-400 loaded RacE in pull-down experiments (Fig. 1D). In addition, 401 constitutively active RacE was capable to release autoinhibition 402of an inactive formin sandwich complex formed by N- and C-403 terminal fragments of ForA in both pulldowns and pyrene assays 404 (Fig. 1 D and E). Finally, we monitored localization of active 405 RacE fused to YFP in cells in 2D-confinement, i.e. under a thin 406 sheet of agar. Unlike previous work that failed to detect active 407 RacE in the cleavage furrow of unconfined cells (32), the active 408

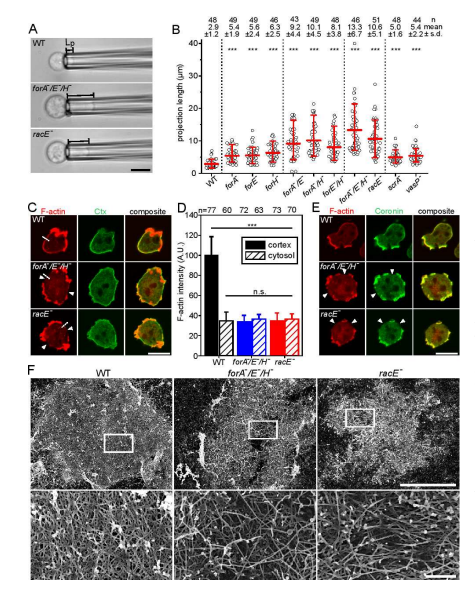


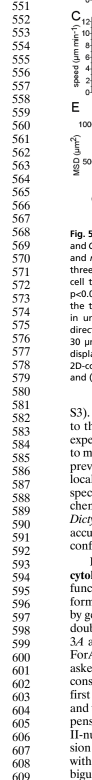
Fig. 4. Elimination of the three cortical formins or of RacE increasingly impairs cortical rigidity and disrupts the contractile actin cortex. (A) Projection length (Lp) of WT and the mutant cells indicated was determined by micropipette aspiration using a constant suction pressure of 500 Pa from time-lapse movies and correspond to Movie S2. Scale bar, 10 µm. (B) Quantitative analysis of the projection lengths of probed cells. n, number of analyzed cells, \*\*\*P < 0.001 by Mann-Whitney rank sum test. Statistical differences refer to WT. (C) Defects of the actin cortex in forA<sup>-</sup>/E<sup>-</sup>/H<sup>-</sup> and racE<sup>-</sup> mutants. Fixed WT and mutant cells were stained with a monoclonal Ctx antibody (green) to visualize PI(4,5)P2-containing membranes and F-actin with phalloidin (red). The white arrow heads indicate regions of low cortical density in the mutants. Scale bar, 10 µm. (D) Quantification of cortical and intracellular actin in WT and mutant cells. Average intensity profiles along 5 pixel wide lines as shown in (C) by the white dashed lines. n, number of analyzed cells; error bars, standard deviations (s.d). (E) Prominent cortical F-actin assemblies outside the breaches in forA /E /H and racE cells are identified as protrusions through containing coronin (green), particularly marking Arp2/3 complex-driven actin networks, and phalloidin-stained F-actin (red). Scale bar, 10 µm. (F) Representative SEM micrographs of detergentextracted WT and mutant cells (low magnification and high magnification of insets shown at top and bottom, respectively). Scale bars, 5 µm (overview) and 0.5 µm (insets).

GTPase accumulated about two-fold in the cleavage furrow as compared to pole regions upon 2D-confinement (Fig. 1F). Moreover, like active ForA, the GTPase was also markedly enriched in the rear cortex of migrating cells (Fig. 1G and ("SI Appendix, Movie S1"), substantiating the view that ForA is regulated by RacE.

Active RacE additionally interacts with cortical formins ForE and ForH. Based on the critical effect of RacE deficiency on cytokinesis, contractility and the regulation of ForA, and the fact that ForE constitutes the only known Rho-family GTPase in Dictyostelium with RhoA-like functions, we reasoned that additional formins may interact with the active GTPase and localize to the cell cortex to safeguard cortex functions. Thus, from the 10 formins expressed in Dictyostelium cells (33), we screened all four potential candidates expressed at the vegetative stage, referred to as ForB, ForE (dDia3), ForH (dDia2) and ForF (dDia1) with Rho- GTPases in the Y2H assay. ForI could be excluded from the screen because of lacking the regulatory GBD, and ForG was omitted due to its specific interaction with active Ras (23).. Strong and specific interactions with active RacE were identified for two of the four tested formins. ForE interacted with active variants of RacE and RacF2 (Fig. 2A). However, since RacF2 appears to carry out specific functions in macrocyst formation during the sexual cycle (34), we did not follow that lead. Unexpectedly, active RacE also interacted with ForH (Fig. 2*B*), previously shown to operate in filopodia formation (35).

Next, we examined the subcellular localization of constitutively active ForE and ForH variants lacking the DAD regions and fused to YFP in freely moving or 2D-confined WT and racE<sup>-</sup> cells. Consistent with previous work (35), ectopic expression of active ForH in WT cells triggered the formation of numerous filopodia, with the active formin being markedly enriched at the cell cortex and filopodial tips (Fig. 2C and ("SI Appendix, Fig. S3")). In 2Dconfinement, filopodia formation was strongly suppressed and active ForH accumulated in the rear cortex of migrating cells, resembling localization of active ForA under the same conditions (17). To our surprise, and as opposed to the entirely diffuse localization of active ForA in racE<sup>-</sup> cells, constitutively active ForH was still able to trigger filopodia formation and accumulate in the cell cortex of freely moving racE<sup>-</sup> mutant cells. In 2Dconfinement, however, the formin became largely cytosolic and failed to be incorporated into the rear.

Active ForE also markedly localized to the cortex of unconfined WT cells and to distal tips of filopodia (Fig. 2D and Fig.



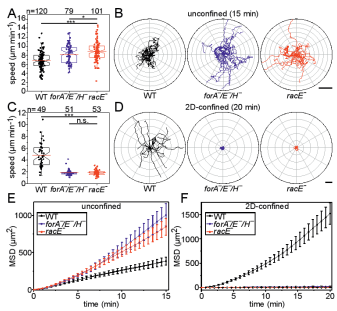
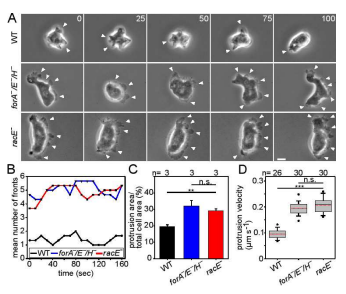


Fig. 5. For A'IE'/H' and racE' mutants cannot migrate in 2D-confinement. (A and C) Box plots summarizing the random migration speed of WT, For A'IE'/H' and racE' cells in (A) unconfined and (C) 2D-confinend conditions. At least three movies from three independent experiments were analyzed for each cell type. n, number of cells analyzed. n.s. non-significant, \* p<0.05, \*\*\* p<0.001 by Mann-Whitney rank sum test. (B and D) Radar plots showing the trajectories of 20 randomly migrating WT, For A'IE'/H' and racE' cells in unconfined and 2D-confinend conditions as indicated. Note the high directional persistence of the mutant cells in unconfined settings. Scale bars, 30 µm for (B) and 20 µm for (D). (E and F) Analysis of the mean square displacement of WT, For A'IE'/H' and racE' cells migrating in unconfined and 2D-confined conditions as indicated. Error bars represent s.e.m. n as in (A) and (C).

S3). In the *racE*<sup>-</sup> mutant, however, the formin was not targeted to the cortex and remained diffuse in the cytoplasm under both experimental settings, implying a requirement for RacE signaling to mediate appropriate subcellular targeting. Consistently, and as previously shown for RacE (36), active ForA, ForH and ForE localized in folate gradients after treatment with latrunculin B specifically on that portion of the plasma membrane facing lower chemoattractant concentrations ("SI Appendix, Fig. S4"). Thus, *Dictyostelium* cells express three RacE-regulated formins that accumulate in cell cortex and rear of cells migrating in 2D-confinement.

Elimination of the three cortical formins causes drastic cytokinesis and developmental defects. To uncover a potential functional redundancy of these three RacE-regulated, cortical formins, we eliminated them all either alone or in combination by gene disruption, to obtain a complete collection of single- and double-mutants as well as a cell line lacking all three formins (Fig. 3A and ("SI Appendix, Fig. S5")). Mutant lines devoid of either ForA or ForH have previously been described (17, 35). Next, we asked whether, or to which extent, cytokinesis is impaired after consecutive elimination of these cortical formins. For that, we first assayed cytokinesis of the for  $A^-$ -single, the for  $A^-/E^-$ -double and the for  $A^{-}/E^{-}/H^{-}$  -triple mutant at high stringency in shaken suspension, and compared effects obtained with RacE- and myosin II-null mutants known to exhibit strong defects in mitotic cell division under these conditions (37, 38). After fixing the cells together with DAPI, we quantified number of nuclei per cell as an unambiguous readout for cytokinesis defects. In cells harvested from shaken suspension after 48 h, the vast majority of WT cells, for A<sup>-</sup>-single, and for A/E-double mutants was mono- or bi-nucleated, although a few  $for A^{-}/E^{-}$ -double mutant cells also displayed three 



**Fig. 6.** For A<sup>-</sup>/E<sup>-</sup>/H<sup>-</sup> and racE<sup>-</sup> mutants form multiple and faster protruding fronts. (A) Gallery with stills from a phase-contrast time-lapse series of randomly migrating WT and mutant cells in unconfined settings corresponding to Movie S4 shows the recurring formation of multiple fronts (white arrow heads) in For A<sup>-</sup>/E<sup>-</sup>/H<sup>-</sup> and racE<sup>-</sup> mutants. Time is in seconds. Scale bar, 5 µm. (B) Quantification of the average number of protruding fronts in migrating WT and mutant cells as shown in (A). (C) Quantification of the ratio of protrusion area over total cell area in WT and indicated mutant cells. Error bars represent s.e.m. n.s. non-significant, \*\* p<0.01 by Mann-Whitney rank sum test.(D) Average protrusion velocities of fronts in WT and mutant cells. Boxes include 50% and whiskers 80% of all measurements, dots represent the 5<sup>th</sup>/95<sup>th</sup> percentile. n, number of cells analyzed. n.s. non-significant, \*\*\* p<0.001 by Mann-Whitney rank sum test.

or four nuclei (Fig. 3 *B* and *C*). By contrast, the *forA*<sup>-</sup>/*E*<sup>-</sup>/*H*<sup>-</sup>triple mutant exhibited a severe cytokinesis defect and was virtually indistinguishable from *racE*<sup>-</sup> and *mhcA*<sup>-</sup> mutants, since about 90% of these mutants developed highly multinucleated cells (Fig. 3 *B* and *C*). To exclude the possibility that a specific formin executes a predominant function in cytokinesis, we additionally performed these cytokinesis assays with all three combinations of double-mutant cells. Although mutant cells lines lacking ForH had a stronger tendency to form multinucleated cells, about 80% of all three formin double-null mutants still contained cells with only one or two nuclei ("SI Appendix, Fig. S6"). Thus, a severe cytokinesis defect was only manifested after inactivation of all three cortical formins.

Importantly, multicellular development also depends on contractility and cortical integrity. Myosin II mutants for instance cannot advance beyond the aggregation stage (39) and double mutants devoid of Ctx I/Ctx II known to tether cortical actin filaments to the membrane entirely fail to develop (40). Thus, we additionally compared multicellular development of WT cells, *forA* -single, *forA* /*E* -double, and the *forA* /*E* /*H* -triple mutant on bacterial lawns. Similar to WT, single- and double formin mutants were still able to advance through development and produce viable spores, although fruiting bodies of the *forA* /*E* double mutant already appeared considerably smaller (Fig. 4D). Notably, the *forA* /*E* /*H* -triple mutant was completely blocked in development and not even able to aggregate. Thus, consistent with their overlapping functions in cytokinesis, all three formins have to be eliminated simultaneously to abrogate morphogenesis.

nave to be eliminated simultaneously to abrogate morphogenesis.675ForA, ForE and ForH synergize in the maintenance of cortical integrity. To quantify the assumed synergistic role of these formins in cortical integrity in the absence of adhesion forces, and to directly compare their contributions to this with those of racE, we performed micropipette aspiration assays (MPA)675675676677678678678679678679670671672673673674679679680

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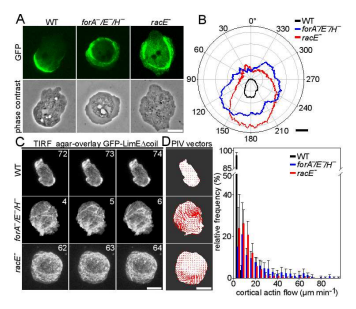
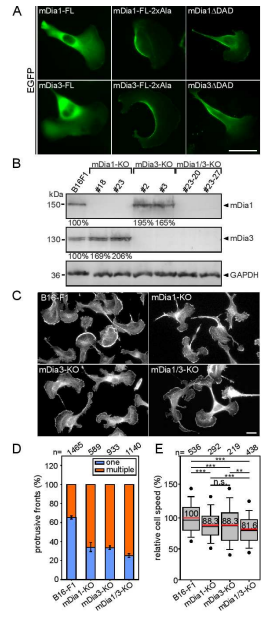


Fig. 7. ForA<sup>-</sup>/E<sup>-</sup>/H<sup>-</sup> and racE<sup>-</sup> mutants cannot polarize and exhibit a drastically increased cortical flow in 2D-confinement. (A) The characteristic localization of GFP-myosin II in the rear cortex of WT cells is abolished in  $for A^{-}/E^{-}/H^{-}$  and racE mutant cells illustrating a major defect in polarization. Still images from time-lapse movies correspond to Movie S6. Scale bar, 10 µm. (B) Quantification of the width of the cortical myosin layer in WT and mutant cells determined by rotational analysis of the fluorescence signal from cells as shown in (A). Radar plot shows the mean myosin II band thickness of the cell lines indicated. Scale bar, 0.5  $\mu$ m. (C) Still images from TIRF time-lapse movies of WT and mutant cells expressing the F-actin probe GFP-LimE $\Delta$ coil in 2D-confined conditions under agar. Time is in seconds. Scale bar, 10 µm. (D) PIV analyses of cortical actin flow in WT and mutant cells. Five consecutive frames recorded at 0.5 s intervals corresponding to Movie S7 were used for PIV analysis. The resulting vectors mark the mean actin velocity per frame (left). Distribution of actin flow velocities of the cell lines indicated reveal a strikingly increased actin flow in the mutants (right). Error bars are s.e.m..

of resuspended WT and mutant cells to measure their global mechanical resistance. To avoid secondary responses of the highly dynamic Dictyostelium cells to external suction pressure, we quantified the initial projection length (Lp) of cells captured from suspension at a constant pressure of 500 Pa in MPA assays. Serial elimination of the formins correlated with increasing defects of cortical rigidity and peaked in the  $for A^{-}/E^{-}/H^{-}$ -triple mutant with an average indentation length of 13  $\pm$  6.7 µm (mean±s.d.) as opposed to  $3 \pm 1.2 \ \mu m$  in WT cells (Fig. 4 A and B). In contrast to many WT cells,  $for A^{-}/E^{-}/H^{-}$  cells were unable to withdraw from the micropipette even at this comparably low suction pressure, and all of them were ultimately sucked into the pipette within 5-10 min ("SI Appendix, Movie S2"). RacE cells also exhibited a major defect of cortical rigidity with an Lp of  $11 \pm 5.1 \,\mu\text{m}$ , although the defect was slightly weaker as compared to the formin triple-knockout mutant. Unexpectedly, however, and albeit none of the analyzed cells was able to completely withdraw from the pipette, almost all cells (98%) resisted complete aspiration at 500 Pa within 10 min. Finally, we also measured the cortical properties of mutant cells lacking the Arp2/3-complex activator Scar and the actin filament elongator VASP. Cortex rigidity of scrA<sup>-</sup> and vasP<sup>-</sup> cells was also clearly impaired as compared to control. However, the contribution of Scar and of VASP was moderate, since measured Lp values were only in the range of the formin single-knockout mutants. Thus, in Dictyostelium Arp2/3 complex and VASP appear to contribute far less to mechanical rigidity of the cortex as compared to formins.

Next, we examined the distribution of cortical F-actin in fixed WT,  $forA^{-}/E^{-}/H^{-}$  and  $racE^{-}$  cells after phalloidin staining. Additionally, we labelled the specimens for PI(4,5)P<sub>2</sub>-binding Ctx, to



Fia. 8. Formation of multiple fronts and defects in polarization and migration in mDia1- and/or -3-deficient B16-F1 cells. (A) Subcellular localization of EGFP-mDia1 and -3 variants in B16-F1 cells migrating on laminin. While full-length (FL) mDia1 and -3 were cytosolic and largely excluded from protrusive fronts, constitutively active mDia1-FL-2xAla (M1182A and F1195A), mDia1<sup>A</sup>DAD, mDia3-FL-2xAla (M1057A and F1170A) and mDia3<sup>A</sup>DAD strongly accumulated in the rear cortex. Scale bar, 20 µm. (B) Immunoblotting of individual and combined mDia1 and mDia3 KO clones as indicated, GAPDH: loading control. Numbers below respective lanes indicate relative changes of mDia1 and -3 expression levels normalized to GAPDH. (C) Polarization defects and formation of multiple fronts in respective cell types, as revealed by phalloidin-staining. Note increasingly pronounced multiplefront phenotypes in single versus double (mDia1/3) KO cells. Scale bar, 10 µm. (D) Quantification of protrusive fronts (one versus multiple) from images as shown in C. Error bars, s.e.m. from at least six independent experiments, n, cell number. (E) Quantification of random migration of respective cell types on laminin. Boxes include 50% and whiskers 80% of all measurements, dots represent the 5th/95th percentile. n.s. non-significant, \*\* p<0.01, \*\*\* p<0.001 by Mann-Whitney rank sum test. n, number of tracked cells.

visualize the lipid gradient between front and rear in polarized *Dictyostelium* cells. In highly polarized WT cells, the bulk of Factin was concentrated in the leading edge and contained only 

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817 small amounts of Ctx, while the rear and lateral sides, encom-818 passing the thin layer of cortical actin, were strongly enriched 819 for Ctx (Fig. 4C). By contrast,  $for A^{-}/E^{-}/H^{-}$  and  $rac E^{-}$  mutants 820 were rounder overall and did not show the characteristic Ctx 821 differential. Most notably, large sections of the cortex in both 822 mutants were devoid of the cortical actin layer, while remaining 823 segments of the cortex still contained prominent F-actin assem-824 blies. Quantification of phalloidin fluorescence intensities across 825 the cortex in actin-deficient areas confirmed this view (Fig. 4D), 826 and was further substantiated by time-lapse imaging of WT and 827 mutant cells expressing the F-actin probe LimE∆coil-GFP ("SI 828 Appendix, Movie S3"). We hypothesized that the thin layer that is 829 missing in the mutants corresponds the contractile actin cortex, 830 whereas the remaining and prominent F-actin assemblies rep-831 resent Arp2/3 complex-driven F-actin structures such as leading 832 edges or endocytic cups. Thus, we labelled WT and mutant cells 833 with phalloidin and for the F-actin binding protein coronin, which 834 is a central constituent of Arp2/3 complex-mediated F-actin net-835 works (41). Consistent with the key function of Arp2/3 complex 836 in protrusion, coronin was strongly enriched in the leading edges 837 of WT cells (Fig. 4E). Notably, coronin was depleted from actindeficient regions, but co-localizing with the prominent F-actin assemblies in for A'/E'/H' and racE' mutants, strongly suggesting 838 839 840 that these structures are nucleated by Arp2/3 complex. Finally, 841 we explored the ultrastructural cortex architecture by scanning 842 electron microscopy (SEM) after detergent extraction of cells. As 843 opposed to the dense, cortical meshwork of WT cells with numer-844 ous overlapping filaments, elimination of the three formins or of 845 RacE caused marked differences in cortical actin organization, 846 including a lower filament density interspaced with large gaps 847 containing much fewer filaments with different geometry (Fig. 848 4F). 849

Cortical formins are essential for motility in 2D-850 confinement. Loss of ForA was previously shown to affect cell migration in unconfined and 2D-confined scenarios (17). 851 852 Thus, we analyzed random cell migration of freely moving and 853 2D-confined for A/E/H and racE mutants in phosphate buffer 854 (PB) employing phase-contrast time-lapse microscopy, and 855 compared migration rates of the mutants to that of WT cells. 856 Additionally, we also determined mean square displacement 857 (MSD) to discriminate locally restricted movement or wiggling of 858 cells from effective directional cell migration (42). Intriguingly, 859 elimination of ForA, ForE and ForH or of RacE even increased 860 the speed of randomly migrating, mutant cells in unconfined 861 environments to 8.2 ± 2.1  $\mu$ m·min<sup>-1</sup> (for A<sup>-</sup>/E<sup>-</sup>/H<sup>-</sup>) or 8.9 ± 2.1 862  $\mu m \cdot min^{-1}$  (racE<sup>-</sup>) (mean±s.d.) as compared to WT controls 863 with 6.8  $\pm$  1.8  $\mu$ m·min<sup>-1</sup> (Fig. 5 A and B). However, when 864 compressed under a sheet of agar,  $for A^{-}/E^{-}/H^{-}$  and  $racE^{-}$  mutants 865 were abrogated for migration (1.9  $\pm$  0.59 µm·min<sup>-1</sup> and 1.8  $\pm$ 866 0.4 µm·min<sup>-1</sup>), as assessed by tracking of the centroids when 867 compared to WT (4.8  $\pm$  1.9  $\mu$ m·min<sup>-1</sup>) (Fig. 5 C and D). 868 Consistent with their higher motility in unconfined settings, a 869 large proportion of both mutants cells were more directional 870 and had higher MSD values as compared to control (Fig. 5 E). 871 In marked contrast, the MSD values of both mutants virtually 872 dropped to zero in 2D-confinement, illustrating their inability 873 to migrate under agar, whereas WT cells were still able to 874 efficiently migrate under these conditions (Fig. 5F). These 875 findings substantiate the fundamental role of the contractile 876 actin cortex for cell migration in confinement. 877

Mutants lacking cortical formins or RacE form multiple fronts. Amoeboid cells such as *Dictyostelium* cells generally exhibit only weak adhesion to the substrate to allow for fast migration in unconfined settings. In *Dictyostelium* cells impaired in the cortical actin cytoskeleton, cell behavior or establishment and maintenance of cell shape are expected to be stronger affected by membrane tension. To test this hypothesis, we imaged freely mov-

885 ing for  $A^{-}/E^{-}/H^{-}$  and rac  $E^{-}$  mutants at high magnification by time-886 lapse phase contrast microscopy, and compared their activities to those of WT cells. WT cells were more spherical and typically 887 formed one or two protruding fronts in the form of pseudopods 888 or macropinosomes at a given time. By contrast,  $for A^{-}/E^{-}/H^{-}$  and 889 *racE* mutants were considerably flatter, as evidenced by strongly 890 reduced halos in phase-contrast images at their cell boundaries. 891 Notably, about 33% of for  $A^{-}/E^{-}/H^{-}$  cells and 19% of rac  $E^{-}$  cells in-892 termittently exhibited a fan-shaped, keratocyte-like morphology 893 and migrated with high, directional persistence, which was con-894 trasted by only 6% of highly directional WT cells ("SI Appendix, 895 896 Movie S4"). Remarkably, both mutants often developed multiple fronts exhibiting 5 or 6 lamellipodia-like pseudopods (Fig. 6 A 897 898 and B and ("SI Appendix, Movie S5"). The elimination of formins and of RacE also substantially increased the combined protrusion 899 area relative to total cell area in the mutants by more than 30% 900 to  $32.0 \pm 5.7\%$  (for A<sup>-</sup>/E<sup>-</sup>/H<sup>-</sup>) and  $29.2 \pm 2.0\%$  (racE<sup>-</sup>) as compared 901 to WT control (19.6  $\pm$  1.7%, Fig. 6C). Moreover, the protrusion 902 903 speed of the fronts in the mutants was about doubled to  $0.19 \pm$ 904  $0.03 \ \mu m \cdot sec^{-1} (for A^{-}/E^{-}/H^{-})$  and  $0.20 \pm 0.03 \ \mu m \cdot sec^{-1} (racE^{-})$  when 905 compared to WT control  $(0.10 \pm 0.02 \ \mu m \ sec^{-1})$  (Fig. 6D). Inter-906 estingly, inhibition of myosin II by blebbistatin had little effect on 907 the motility of  $ForA^{-}/E^{-}/H^{-}$  and  $racE^{-}$  mutant cells ("SI Appendix, 908 Fig. S7"), excluding augmented actomyosin contractility as direct 909 cause for these effects. Although RICM analyses revealed a larger 910 contact area of mutant cells ("SI Appendix, Fig. S8 A and B"), 911 they formed fewer actin foci, and these adhesion points were 912 significantly shorter lived than those in controls ("SI Appendix, 913 Fig. S8 C-E"). In line with these observations, the contact area 914 of multiple front- or keratocyte-shaped cells was inhomogeneous 915 and interspersed with less adhesive sections, as evidenced by 916 RICM ("SI Appendix, Movies S6 and S7"). Thus, the combination 917 of increased protrusive activity and decreased adhesiveness in the 918 mutants may explain the highly directional persistence in motility 919 assays. Finally, we noticed that the growth of initially formed 920 fronts in the mutants typically ceased when multiple, competing 921 protrusions were formed on the opposite side of the cell. In these 922 cases, initial fronts rapidly lost adhesion to the underlying surface 923 and were effectively retracted into the cell body ("SI Appendix, 924 Fig. S9 and Movies S 6 and S8"). 925

Polarity defects and dramatically increased cortical actin 926 flow in for  $A^{-}/E^{-}/H^{-}$  and rac  $E^{-}$  mutants. An intact contractile 927 actin cortex of amoeboid cells regulates cell migration in 2D-928 confinement by guiding hydrostatic pressure, created by acto-929 myosin contraction in the rear, to the front to promote leading 930 edge protrusion (17). Thus, we monitored myosin II and F-actin 931 representing the two main components of the contractile machin-932 erv in WT and mutant cells after mechanical stress in confinement 933 under agar. WT cells expressing fluorescently-tagged heavy chain 934 of myosin II were highly polarized, and the motor protein was 935 continuously concentrated in a compact, crescent-like sheet at 936 the rear cortex beneath the plasma membrane (Fig. 7A and 937 ("SI Appendix, Movie S9"). In striking contrast, for  $A^{-}/E^{-}/H^{-}$  and 938 racE<sup>-</sup> mutants remained highly unpolarized in 2D-confinement, 939 as evidenced by the aberrant circular localization of myosin II in 940 a band-like fashion along most of the cell periphery, albeit this 941 phenotype was slightly less prominent in  $racE^{-}$  cells (Fig. 7A). 942 Moreover, myosin II did not accumulate in a crisp band as in 943 WT cells, but was dispersed into multiple, string-like assemblies 944 in the mutants. Myosin II was also largely dislodged from the 945 membrane and was instead strongly enriched at an endoplasm-946 ectoplasm interface separating the organelle free area from the 947 cell interior (Fig. 7A). Consistently, time-lapse imaging of the 948 mutant cells revealed a highly erratic behavior of myosin II as-949 sociated with intense blebbing, substantiating the massive defects 950 in the contractile cell cortex of mutant cells ("SI Appendix, Movie 951 S9"). Quantification of myosin II band width as well as its radial 952

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953 distribution in WT and mutant cells in polarity plots corroborated 954 this view (Fig. 7B).

955 We then analyzed the dynamic behavior of cortical actin fila-956 ments at the ventral plasma membrane by total internal reflection 957 fluorescence (TIRF) microscopy in WT and mutant cells after 958 confinement under agar. In WT cells, cortical actin, visualized 959 by the F-actin marker LimE∆coil-GFP, was organized into a 960 delicate, filamentous web, interspaced with dynamic actin foci, 961 which accumulated most strongly in protruding fronts evidently 962 driven by Arp2/3 complex-mediated actin assembly (Fig. 7C and 963 ("SI Appendix, Movie S10")). A filamentous network was also 964 observed in the for  $A^{-}/E^{-}/H^{-}$  and rac  $E^{-}$  mutants. However, presum-965 ably due to the high membrane tension and their strong polar-966 ization defect in 2D-confinement, they did not form protruding 967 fronts. Moreover, as assessed from TIRF time-lapse imaging, the 968 dynamics of cortical actin filaments was drastically changed as 969 compared to control. These filaments were rapidly pulled into 970 the cell center in a process reminiscent of actin retrograde flow 971 in higher eukaryotes. Particle image velocimetry (PIV)-based 972 quantification of cortical actin flow confirmed this notion (Fig. 973 7D). The velocity distribution showed flows of up to 72  $\mu$ m·min<sup>-1</sup> 974 in racE<sup>-</sup> cells and almost up to 100  $\mu$ m min<sup>-1</sup> in the for A<sup>-</sup>/E<sup>-</sup>/H<sup>-</sup> 975 mutant as compared to the average flow with 1.48±1.08 µm·min<sup>-1</sup> 976 (mean±s.d.) in WT cells (Fig. 7E). Consistently, PIV analyses 977 further revealed that the region with fastest actin filament flows 978 overlap with myosin II-enriched regions in  $for A^{-}/E^{-}/H^{-}$  and  $rac E^{-}$ 979 mutants (Fig. 7 A and E and ("SI Appendix, Fig. S10"). Together, 980 these data strongly suggest that filaments nucleated by cortical 981 formins regulate subcellular myosin II localization and cortical 982 actin flow under mechanical stress. 983

Cell lines lacking murine mDia1 and -3 display phenotypes 984 indicative of conserved, cortical functions. To explore whether 985 our observations are generalizable to higher eukaryotes, we 986 analyzed mammalian formins in highly motile B16-F1 mouse 987 988 melanoma cells. Since mDia subfamily formins (1, 2, and 3) are regulated by RhoA (43), which is well established to drive 989 contractility, we focused on subcellular localization of EGFP-990 991 tagged mDia variants after ectopic expression in B16-F1 cells. Consistent with our previous findings (17), we found constitu-992 tively active mDia1 variants, i.e. mDia1<sup>A</sup>DAD as well as the 993 newly designed point mutant mDia1-FL-2xAla (M1182A and 994 995 F1195A), which is expected to release autoinhibition of the FL protein (44), to localize prominently in the cell rear, whereas 996 the full length, autoinhibited protein remained cytosolic (Fig. 997 8A). Virtually identical results were obtained with corresponding 998 mDia3 variants (Fig. 8A), whereas active mDia2 was not found 999 1000 in the cell rear, but mostly targeted to filopodia tips (45). To evaluate mDia functions in the mammalian cell cortex, we em-1001 ployed CRISPR/Cas9-mediated disruption of the genes encoding 1002mDia1 and -3, both individually and in combination in B16-F1 1003 cells. Loss of respective protein in independent, clonal cell lines 1004 was confirmed by immunoblotting (Fig. 8B). Interestingly, mDia1 1005 levels were evidently increased in both mDia3-KO cell lines, and 1006 vice versa, indicative of compensatory, regulatory mechanisms 1007 presumably serving to sustain sufficient levels of these cortical 1008 formins (Fig. 8B). Phalloidin stainings revealed defects in cell 1009 polarization as well as markedly increased frequencies in mDia1 1010 and -3 single mutants of cells forming multiple fronts, a phenotype 1011 that was even further increased in mDia1/3 double-KO cells 1012 (Fig. 8 C and D) and strikingly reminiscent of cortical formin 1013 pathway KOs in Dictyostelium. Next, we analyzed random cell 1014 migration of B16-F1 wildtype and mutant cells on laminin using 1015 time-lapse, phase-contrast microscopy. Interestingly, as opposed 1016 to Dictyostelium cells migrating without 2D-confinement, cell 1017 depolarization and apparent stimulation of the multiple front 1018 phenotype reduced the efficiency of the highly adhesive mode of 1019 melanoma cell migration, likely caused by inefficient protrusion 1020

in a productive, migratory direction. Specifically, whereas single 1021 mDia1 and -3 mutants displayed a moderate, but statistically 1022 significant reduction of migration rate in this assay (by 11.7%), re-1023 1024 moval of both mDia1 and -3 decreased migration even further (by 18.4%; Fig. 8*E*). Together, these data strongly suggest the RhoA-1025 effectors of the mDia formin subfamily, in particular mDia1 and 1026 -3 to exert functions in the actin cortex that are conserved in 1027 1028 evolution from Dictyostelium to mammals. 1029

# Discussion

Over thirty years ago, Bray and White postulated that cortical 1032 contractility may not only contribute to retraction of the trail-1033 ing edge, but also to ingression of the cleavage furrow during 1034 cytokinesis (46). Since then, localization and participation in 1035 both processes has been demonstrated for numerous cell cortex 1036 components including myosin II (47), Ctx (27), PTEN (48) and 1037 IQGAP family members (28). Here, we demonstrated that three DRFs, ForA, ForE and ForH act synergistically in the assembly of actin filaments in the contractile actin cortex as evidenced by the increasing severity of additive KO phenotypes, which directly correlated with a gradual decrease in mechanical cortex rigidity. We also showed that the active form of the Rho family GTPase RacE, binds to the GBD of ForA, releasing its autoinhibition to initiate actin assembly. RacE shows considerable sequence similarity with Rho proteins from other species and represents the closest homologue of mammalian RhoA in Dictyostelium (36, 49). The phenotype of  $racE^{-}$  cells was similar to that of the triple knockout DRF cells, including large cytokinesis defects in suspension (31) as well as large effects on development (36). This reinforces the idea that the DRFs are required for filament formation, and this is regulated by RacE. Moreover, RacE localizes to the cell rear as well as cleavage furrow in mitotic cells, and is essential for cortical localization of ForA and ForE.

Similar to the redundancy in cortical actin organization exhibited by DRFs in Dictyostelium, we also found stronger, complementary phenotypes after combined inactivation of the formins mDia1 and -3 in B16-F1 mouse melanoma cells. This redundancy of the microfilament system is a common phenomenon for essential cellular activities, and has been observed previously for the actin-crosslinking proteins  $\alpha$ -actinin and filamin (50) and for Ctx I and II (51) in Dictyostelium. While ForA, ForE and ForH act synergistically in actin filament assembly in the contractile actin cortex, it is likely that they also have isoform specific roles and can be activated by other signaling pathways. For example, we found that constitutively active ForH localized to the cortex in unconfined racE<sup>-</sup> cells, but remained largely cytosolic after compression in 2D-confinement, suggesting multiple interactors mediate its subcellular targeting. Of these, RacE is presumably important for ForH targeting under mechanical stress. Incidentally, ForH was recently also found as potential RacE interactor by mass spectrometry (52). In addition, active ForH and ForE also trigger filopodia formation, but as both cortex and filopodia constitute F-actin structures directly associated with the plasma membrane, these actin assembly factors may well participate in the formation of various membrane-associated structures entailing long, unbranched filaments.

The strong effects on cytokinesis and cortex-dependent func-1079 tions in for A/E/H are similar to those found in rac *E* cells. This 1080 indicates the phenotypic effect of racE gene elimination to derive 1081 from the lack of forA/E/H activation. Consistent with this, none 1082 of wild type formins (ForA, ForE or ForH) localized to the cortex 1083 as they are likely autoinhibited ("SI Appendix, Fig. S11") and 1084 (17). We also found that active RacE accumulates in the cleavage 1085 furrow of mitotic cells upon 2D-confinement, implying that it 1086 plays a role in regulating cytokinesis by recruitment and activation 1087 of cortical formins at this site. 1088

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Both  $for A^{-}/E^{-}/H^{-}$  and  $rac E^{-}$  mutants displayed flat morpholo-1089 1090 gies and formation of multiple, dynamic multi-directional protrusions along the cell contour. In agreement with recent work 1091 1092 analyzing enhanced expansion of lamellipodial networks upon reduction of plasma membrane load (2), and considering the-1093 oretical calculations of cortex mechanics (53), these data imply 1094 that cells harboring a compromised viscoelastic cell cortex may 1095 1096 experience reduced resistance to actin polymerization forces in protrusions. If correct, this could well cause the formation of am-1097 plified protrusions and multiple fronts observed in our mutants. 1098 1099 Multiple protrusion formation in Dictyostelium was accompanied by adhesion weakening and irregular detachment. Consequently, 1100 most unconfined  $for A^{-}/E^{-}/H^{-}$  and  $rac E^{-}$  cells, in some instances 1101 1102 resembling fan-shaped keratocytes, migrated faster and more directional than their wildtype controls. These phenotypes ap-1103 peared at the expense of the more amoeboid type of migration 1104 1105 usually seen in Dictyostelium.

Of note, the inhibition of myosin II by blebbistatin had little 1106 effect on the motility of keratocytes (54) as well as  $ForA^{-}/E^{-}/H^{-}$  and 1107 racE<sup>-</sup> mutant cells ("SI Appendix, Fig. \$7"). This indicates that the 1108 1109 migratory modes adopted by these cell types and/or experimental conditions are less dependent on cortical contractility than during 1110 1111 amoeboid or canonical, mesenchymal migration, and additionally mostly driven by the amplified network expansion activity at the 1112 leading edges of these cells (55). Interestingly, the migratory 1113 behavior of Dictyostelium cells can be switched from amoeboid 1114 to keratocyte-like by either decreasing PIP2 levels or increasing 1115 Ras/Rap signaling (56). Whether or not these phenotypes could 1116 relate to the molecular mechanisms described here will be an 1117 1118 exciting topic of future study. 1119

In 2-D confinement, *forA* /*E* /*H* and *racE* mutants are unable to localize myosin II properly to the cell cortex, polarize and migrate. Symmetry breaking obtained by an anisotropic distribution of components including myosin II drives both, rear retraction in directed cell migration and cytokinesis. How myosin II is localized to the cell cortex is still not fully understood, but has been proposed to include signaling and mechanical cues including myosin phosphorylation by MHCK-A, PTEN, Ctx I or talin in *Dictyostelium* (14, 48, 57, 58) or preferential binding to stretched actin filaments (59) In any case, a severely perturbed cortical actin cytoskeleton will likely interfere with myosin II positioning and activity through all these pathways. Our findings thus establish cortical formins as key to the establishment of polarity and properly regulated migration in both *Dictyostelium* and mammalian cells.

Consistent with the elimination of these formins or their activator RacE, SEM and live-cell TIRF imaging revealed cortical filament density in  $forA^{-}/E^{-}/H^{-}$  and  $racE^{-}$  cells to be reduced compared to WT, but raised the question as to which assembly factors are involved in generating the remaining cortical actin filaments. Our immunofluorescence results with fixed cells indicated these filaments to be primarily nucleated by remaining prominent actin assembly factors, such as Arp2/3 complex and VASP, although we cannot exclude at this stage the presence of

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filaments assembled by other formins, for instance ForG, known to cooperate with Arp2/3 complex in large scale endocytosis (23) or of ForF (dDia1), which homogenously accumulates in the entire pseudopod (60). 1157 1158 1159 1160

In 2-D confinement,  $for A^{-}/E^{-}/H^{-}$  and  $rac E^{-}$  mutants exhibited exceptionally fast centripetal flows of residual cortical filaments at velocities of 50-100 µm/min, which were contrasted by almost immobile cortical networks relative to the advancement of mi-1164 grating, wildtype cells. This suggests that in 2D-confinement, Dic-1165 tyostelium does not primarily use retrograde actin fluxes to drive 1166 force transmission during migration, as recently proposed for 1167 other cell types utilizing amoeboid motility (61, 62). However, the 1168 precise cause of the excessive cortical actin flows observed in the 1169 mutants remains to be clarified. An intact cortical cytoskeleton 1170 1171 harbors various transmembrane proteins and receptors, poten-1172 tially acting as barriers constraining lateral diffusion (63). Thus, we speculate that the increased cortical flows observed in our 1173 1174 mutants are caused by diminished viscosities of their perturbed 1175 cortical networks. 1176

Our initial characterization of genome-edited B16-F1 mouse melanoma mutants devoid of mDia1, mDia3 or both formins revealed striking similarities to the Dictyostelium system. Comparable to Dictyostelium ForA, ForE and ForH, both mammalian formins are regulated by Rho-subfamily proteins, which are ultimately linked to contractility. Active variants of mDia1 and -3, but not the autoinhibited full-length proteins localized prominently to the rear cortex in polarized B16-F1 cells, and the individual elimination of these formins triggered the formation of multiple protrusive fronts as well as substantial defects in polarization and migration. The fact that the phenotypes were noticeably amplified in double mutants reinforced the conclusion of their overlapping functions. This is consistent with a very recent study analyzing the contractile actin cortex in Sertoli cells of mouse seminiferous tubules (64). Loss of mDia1 and -3 in these cells compromised the cortical actin cytoskeleton leading to less dense F-actin meshworks ultimately resulting in impaired spermatogenesis. Taken together, our results suggest that formins are important in cell cortex establishment and maintenance, and that these functions are evolutionarily conserved across far distant organisms.

# **Materials and Methods**

A complete description of the methods is provided in SI Materials and Methods. This description includes construct generation, cell culture, transfections and establishment of *Dictyostelium* and B16-F1 mutants, protein purification, actin-assembly and pull-down assays, antibodies and immunoblots, imaging, aspiration assays, analyses of cell migration, Y2H assays, and quantification of actin flows and statistical analyses.

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