Memo–RhoA–mDia1 signaling controls microtubules, the actin network, and adhesion site formation in migrating cells

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ACTIN assembly at the cell front drives membrane protrusion and initiates the cell migration cycle. Microtubules (MTs) extend within forward protrusions to sustain cell polarity and promote adhesion site turnover. Memo is an effector of the ErbB2 receptor tyrosine kinase involved in breast carcinoma cell migration. However, its mechanism of action remained unknown. We report in this study that Memo controls ErbB2-regulated MT dynamics by altering the transition frequency between MT growth and shortening phases. Moreover, although Memo-depleted cells can assemble the Rac1-dependent actin meshwork and form lamellipodia, they show defective localization of lamellipodial markers such as α-actinin-1 and a reduced number of short-lived adhesion sites underlying the advancing edge of migrating cells. Finally, we demonstrate that Memo is required for the localization of the RhoA guanosine triphosphatase and its effector mDia1 to the plasma membrane and that Memo–RhoA–mDia1 signaling coordinates the organization of the lamellipodial actin network, adhesion site formation, and MT outgrowth within the cell leading edge to sustain cell motility.

Introduction

Cell motility is the result of a complex series of events that must be integrated both spatially and temporally (Ridley et al., 2003). Efficient motility requires the coordinated regulation of actin, microtubules (MTs), and adhesion sites throughout the migration process (Lauffenburger and Horwitz, 1996; Rodríguez et al., 2003). While investigating signaling pathways involved in ErbB2-driven breast carcinoma cell migration, we have identified Memo, a protein that does not belong to any known family of signaling molecules, as a novel ErbB2 effector (Marone et al., 2004). Memo knockdown cells showed reduced migration despite their preserved ability to extend lamellae/lamellipodia. Interestingly, Memo appears to be involved in MT outgrowth within cell protrusions (Marone et al., 2004). We have investigated the mechanisms underlying Memo’s function in ErbB2-driven motility in living breast carcinoma cell lines. We found that Memo regulates MT dynamics, formation of adhesion sites, and the organization of the lamellipodial actin network by contributing to localize the small G protein RhoA and its effector mDia1 at the plasma membrane.

Results and discussion

In migrating cells, MTs are usually organized radially with the minus ends anchored at the centrosome and the plus ends probing the cytoplasm. Because MT plus ends are dynamically unstable and switch abruptly between phases of growth, pause, and shortening (Mitchison and Kirschner, 1984), MT net growth depends on the tight regulation of MT dynamic instability. Upon heregulin–β1 (HRG)–induced ErbB2 activation, breast carcinoma cells extend wide membrane protrusions in the direction of cell displacement invaded by outgrowing MTs (Fig. 1 A). Incorporation of EGFP–α-tubulin into MTs allowed the tracking of individual MT plus ends in the lamellae/lamellipodia and the analysis of MT dynamics (Fig. 1 A and Video 1, available at

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Memo knockdown cells were still capable of forming protrusions of the same size and at the same rate as control cells (unpublished data). However, despite the fact that MTs were still dynamic (Video 2), they failed to enter lamellipodia (Fig. 1B). Analysis of plus end dynamics showed that in Memo knock-down cells, HRG treatment failed to decrease catastrophes or increase rescues (Fig. 1, C and E) and no longer affected the time MTs spent in the growth/shortening phases (Fig. 1, D and F) in SKBR3 and T47D breast carcinoma cells. Importantly, http://www.jcb.org/cgi/content/full/jcb.200805107/DC1). We found that HRG treatment did not affect MT growth or shortening rates (not depicted) but strongly decreased the frequency of transitions from growth or pause to shortening (catastrophes) and increased the frequency of transitions from shortening to growth or pause (rescues; Fig. 1, C and E). As a consequence, MTs spent more time growing than shortening, in contrast to what was observed in the absence of stimulation (Fig. 1, D and F).

Expression of Memo was knocked down via siRNA (Fig. S1). Memock knockdown cells were still capable of forming protrusions of the same size and at the same rate as control cells (unpublished data). However, despite the fact that MTs were still dynamic (Video 2), they failed to enter lamellipodia (Fig. 1B). Analysis of plus end dynamics showed that in Memo knock-down cells, HRG treatment failed to decrease catastrophes or increase rescues (Fig. 1, C and E) and no longer affected the time MTs spent in the growth/shortening phases (Fig. 1, D and F) in SKBR3 and T47D breast carcinoma cells. Importantly,
reexpression of Memo restored all ErbB2 effects on MTs (Fig. 1 G). Interestingly, Memo depletion did not significantly affect basal MT dynamics. Thus, Memo is an essential effector of ErbB2-regulated MT dynamics.

MTs have been shown to target adhesion complexes and promote their disassembly (Kaverina et al., 1999). Thus, we investigated the consequences of Memo knockdown on adhesion complexes formed by migrating cells, using paxillin as a marker. We observed that DsRed-paxillin was incorporated into two discrete populations of adhesions: small short-lived adhesion sites (mean lifetime of 12.3 ± 0.4 s; n = 70 individual adhesions in seven cells) underlying the advancing leading edge and large stable sites, the focal adhesions ([FAs] mean lifetime of 11.6 ± 1.2 min; n = 20 individual adhesions in four cells) deeper within the lamella, which derive from a subpopulation of the former (Fig. 2 A and Video 3, available at http://www.jcb.org/cgi/content/full/jcb.200805107/DC1). Interestingly, Memo knockdown resulted in a striking decrease in the number of small adhesion sites, whereas the number of FAs increased (Fig. 2, A and B; and Video 4). In accordance with the known role of MTs on FA turnover, we found that the lifetime of adhesion sites was increased in the absence of Memo (unpublished data). Reintroduction of Memo in knockdown cells restored small adhesions (Fig. 2 A). We propose that the reduction in short-lived adhesion sites that underlie membrane protrusions combined with the increased number of FAs that anchor cells to the substratum contributes to the motility defect in Memo knockdown cells.

Because of the known link between adhesion sites and actin filaments, we examined Memo’s impact on the assembly of the filamentous-actin network in migrating cells. Actin filaments assemble at the leading edge of migrating cells in a Rac1- and Arp2/3-dependent manner to form a branched network that pushes the cell membrane forward (Pollard and Borisy, 2003). EGFP-actin labeling was concentrated in several structures, including the cell cortical area and actin stress fibers, and appeared similar in control and Memo-depleted cells. Width of α-actinin-labeled lamellipodia (brackets) is reduced in Memo-depleted cells (middle) and restored upon Memo reexpression (right). Actin labeling is unchanged. Insets show enlargements of leading edges. (E and F) EGFP-actin (E) or EGFP-α-actinin-1 (F) fluorescence was bleached (arrows) within the lamellipodia of migrating cells, and fluorescence recovery was monitored over time (images from Videos 9 and 10, available at http://www.jcb.org/cgi/content/full/jcb.200805107/DC1). Actin fluorescence recovers from the cell membrane reflecting topical actin assembly, whereas α-actinin fluorescence recovers at a faster rate throughout the lamellipodia. Bars, 10 μm.
We have further explored the organization of the lamellipodial network using FRAP experiments. After photobleaching of EGFP-actin, the whole bleached area flowed back until disappearance in the lamella, with little recovery through diffusion (Fig. 2E and Video 9, available at http://www.jcb.org/cgi/content/full/jcb.200805107/DC1). This observation illustrates that the lamellipodial actin network is assembled at the cell membrane as a stable structure that is carried back by retrograde flow until it reaches the lamellipodium/lamella border, where it is disassembled (Ponti et al., 2004). Arp3 displayed the same

We observed that α-actinin–EGFP labeled a structure that overlapped with EGFP-actin–labeled lamellipodia. Interestingly, we observed that the α-actinin–labeled lamellipodia were almost 50% thinner in Memo knockdown relative to control cells (Fig. 2D, Fig. 4A, and Videos 7 and 8). Importantly, the width of α-actinin–labeled lamellipodia was restored to normal upon Memo reexpression (Fig. 2D and Fig. 4A). The effect was even more drastic for endogenous α-actinin-1, as immunolabeling appeared to be decreased throughout the lamellipodia of Memo-deficient cells (Fig. S1).

We have further explored the organization of the lamellipodial network using FRAP experiments. After photobleaching of EGFP-actin, the whole bleached area flowed back until disappearance in the lamella, with little recovery through diffusion (Fig. 2E and Video 9, available at http://www.jcb.org/cgi/content/full/jcb.200805107/DC1). This observation illustrates that the lamellipodial actin network is assembled at the cell membrane as a stable structure that is carried back by retrograde flow until it reaches the lamellipodium/lamella border, where it is disassembled (Ponti et al., 2004). Arp3 displayed the same
Expression of active mDia1 also restored formation of small adhesion sites (in 92% of the cells vs. 12% of Memo knockdown cells; \( n = 69 \) and 74, respectively; Fig. 3 C) and normal actinin-labeled lamellipodia (Fig. 4 A) in Memo knockdown cells. Because mDia1 is an effector of RhoA (Watanabe et al., 1997), we tested the role of RhoA in Memo signaling. Expression of moderate levels of an active form of RhoA (RhoV14) in Memo knockdown cells induced thick stress fibers and large FAs but also restored Memo functions: MT outgrowth (in 89% of the cells vs. 5% of Memo-depleted cells and 90% of control cells; \( n = 73, 61, \) and 87, respectively; Fig. 3 A), formation of small adhesion sites (in 87% of the cells vs. 8% of Memo-depleted cells and 92% of control cells; \( n = 78, 63, \) and 79, respectively; Fig. 3 C), and normal lamellipodial actin (Fig. 4 A). These results demonstrate that RhoA and mDia1 are important Memo effectors that control MTs, the actin network, and adhesion sites. Finally, we evaluated the impact of the Memo–RhoA–mDia1 signaling pathway on cell motility using time-lapse microscopy. We found that Memo knockdown decreased velocity of SKBR3 cells by 40–50% (Fig. 4 B). Reintroduction of Memo, active mDia1, or active RhoA in Memo knockdown cells restored normal cell velocity, revealing the significant contribution of the Memo–RhoA–mDia1 pathway to ErbB2-dependent breast carcinoma cell migration.

Activation of RhoGTPases is associated with their re-localization from the cytosol to the cell membrane (Kurokawa and Matsuda, 2005; Pertz et al., 2006). We investigated the possibility that Memo controls the recruitment of RhoA and mDia1 to the cell membrane. HRG treatment induced the recruitment of Memo from the cytosol to the plasma membrane (Marone et al., 2004). EGFP-mDia1, EGFP-RhoA, and the respective endogenous proteins were similarly recruited to lamellipodial membranes and ruffles (Fig. 5 A). RhoA and mDia1 labeling is not simply caused by increased cell volume at ruffles because cytoplasmic markers (e.g., the mCherry fluorescent protein) show limited labeling of ruffles (Fig. S2, available at 405M EMO – RHO A – M D IA 1 CONTROLS CELL MOTILITY • Zaoui et al. 405M EMO – RHO A – M D IA 1 CONTROLS CELL MOTILITY • Zaoui et al. 405M EMO – RHO A – M D IA 1 CONTROLS CELL MOTILITY • Zaoui et al. 405M EMO – RHO A – M D IA 1 CONTROLS CELL MOTILITY • Zaoui et al. 405M EMO – RHO A – M D IA 1 CONTROLS CELL MOTILITY • Zaoui et al. 405M EMO – RHO A – M D IA 1 CONTROLS CELL MOTILITY • Zaoui et al. 405M EMO – RHO A – M D IA 1 CONTROLS CELL MOTILITY • Zaoui et al.
In fact, the functional interaction between Memo and RhoA could be the consequence of a physical interaction, as bacterially expressed Memo interacted with purified RhoA in pull-down experiments. Interestingly, Memo interacted preferentially with the active forms of RhoA (GTP-bound and constitutively active L63QRhoA). In contrast, Memo did not bind GTP-loaded Cdc42 or Rac1, which is indicative of the specificity of the interaction (Fig. 5D). Thus, we propose that Memo retains active RhoA at the plasma membrane to allow spatially and temporally restricted activation of RhoA effectors such as mDia1 at the cell leading edge.

Importantly, Memo knockdown prevented recruitment of EGFP-RhoA or EGFP-mDia1 to the plasma membrane (Fig. 5A and Fig. S2). This was not caused by decreased ruffling activity, as other RhoGTPases such as Rac1 and Cdc42 are still associated with plasma membrane and ruffles in the absence of Memo (Fig. 5 B and Fig. S3). Biochemical analysis of lamellipodia- and cell body-enriched fractions (Cho and Klemke, 2002) confirmed that RhoA and Memo were enriched in the cell leading edge and that decreased expression of Memo led to a specific reduction of RhoA in lamellipodia (Fig. 5 C).

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If Memo’s main role is to localize RhoA to the plasma membrane, a membrane-targeted form of RhoA should compensate for the loss of Memo in Memo knockdown cells. Expression of a fusion protein between wild-type RhoA and the palmitoylation motif of RhoB (RhoA-CCCKVL; Michaelson et al., 2001) promoted constitutive membrane localization independently of Memo expression levels (Fig. 5E). Expression of RhoA-CCCKVL in Memo-deficient cells restored MT outgrowth (in 81% of the cells vs. 10% of RhoA-expressing cells; n = 65 and 61 cells, respectively) but also normal lamellipodial localization of α-actinin-1 and cell motility (unpublished data), showing that forcing RhoA to the plasma membrane recapitulates Memo’s functions. The data also suggest that Memo is not directly required for activation of RhoA because wild-type membrane-associated RhoA was fully functional in the absence of Memo.

It is noteworthy that Memo signals via RhoA but does not control known functions of RhoA such as formation of stress fibers or maturation of FAs (Burridge and Wennerberg, 2004). In fact, these are under the control of the Rho-associated kinase (ROCK), as Y27632 inhibition of ROCK activity drastically reduced stress fibers (not depicted) and FAs while preserving small peripheral adhesions (Fig. 3C), which is in striking contrast to mDia1 inhibition. These results are in line with earlier work showing that ROCK is involved in the formation of central but not peripheral adhesions (Totsukawa et al., 2004). Moreover, an active form of ROCK induced thick stress fibers and large FAs but failed to restore small adhesion sites or lamellipodial α-actinin labeling in Memo-depleted cells (unpublished data). Thus, Memo knockdown provides a method to distinguish between discrete pools of RhoA with distinct functions: one controlling cortical actin and initiation of adhesion sites via the mDia1 effector at the cell leading edge, and the other triggering ROCK-dependent stress fiber formation, adhesion maturation, and cell contraction within the cell body.

Although the role of the Rac1–Scar–Arp2/3 pathway for the assembly of the branched actin meshwork was clearly demonstrated (Pollard and Borisy, 2003), models of lamellipodia protrusion need to take into account evidences for the coexistence within the lamellipodium of kinetically distinct yet overlapping actin networks (Ponti et al., 2004) and the role of other types of regulators of actin assembly such as formins (Yang et al., 2007) and Ena/VASP proteins (Bear et al., 2002). We propose that mDia1, as an effector of Memo–RhoA, contributes to the organization of the lamellipodial actin network. mDia1 controls the lamellipodial localization of both α-actinin-1 and cortactin, suggesting that an alteration of the network organization allows actin-binding protein association to the actin network. We are exploring the possibility that, as shown for mDia2 recently (Yang et al., 2007), mDia1 controls the length or branching of actin filaments. This could lead to a modification of the structure of the actin network and allow α-actinin-1 localization and bundling or scaffolding activity in lamellipodia (Otey and Carpen, 2004). As a recent study suggested that myosin-dependent tension exerted on the lamellipodial network controls formation of adhesion sites (Giannone et al., 2007), changes in actin filament length, branching, or bundling could in turn affect the response of the actin network to tension and formation of peripheral adhesions. Thus, our study showing that the Memo–RhoA–mDia1 pathway controls the cytoskeleton and adhesion sites at the cell leading edge adds a layer of complexity to the signaling network that controls cell motility.

Materials and methods

Cell transfection and plasmid constructs

SKBR3 and T47D breast carcinoma cells grown in DMEM and 10% fetal calf serum were transfected by nucleofection (Amaxa) with 2 μg siRNA for Memo (nt 1,438–1,458; within the 3′ untranslated region), mDia1 (nt 132–152), or LacZ (nt 2,289–2,309; Invitrogen) and the following plasmid constructs: GFP-Memo, Myc-Memo6, EGFP–α-actinin (Clontech Laboratories, Inc.), DiRed–β-tubulin (provided by V. Homburger, Institut de Génomique Fonctionnelle, Montpellier, France), EGFP–α-actinin-1 (provided by J. Wehland, Helmholtz Centre for Infection Research, Braunschweig, Germany), paxillin-DiRed, paxillin-EGFP (provided by A. R. Horwitz, University of Virginia, Charlottesville, VA), EGFP-mDia1, Flag-mDia1L133 (provided by S. Narumiya, Kyoto University, Kyoto, Japan), EGFP-RhoA-CCCKVL, EGFP-RhoA (provided by M. R. Philips, New York University, New York, NY), EGFP-RhoA-V14, EGFP-Rac1, EGFP-Cdc42 (provided by M. A. Schwartz, University of Virginia, Charlottesville, VA), or EGFP–α-actinin-1 (11908; Addgene; provided by C. A. Otey, University of North Carolina, Chapel Hill, NC).

Analysis of MT dynamic instability

Cells grown on collagen-coated glass coverslips were placed in a double coverslip chamber maintained at 37°C and observed upon addition of 5 nM HRG (R&D Systems) using the 100× plan fluor NA 1.3 or plan Apochromat NA 1.4 objective of a fluorescence microscope (DMIRBE Microsystem [Leica]; Axiovert 200 [Carl Zeiss, Inc.]). 31 images per cell or 1-min intervals for 30 min. Background-corrected fluorescence intensity curves generated from photobleached areas of the same sizes and locations were sampled every 0.5 s for 42 s and corrected for overall photobleaching and were used to calculate h/2.

Quantification of adhesion sites and actin dynamics

Cells were observed as described in the previous paragraph except that images of paxillin-expressing cells were acquired at 4-s intervals for 2 min or 1-min intervals for 30 min. Background-corrected fluorescence intensity images were used to measure small adhesion sites and FA number and lifetime in lamellipodia and lamellae, respectively. MetaMorph software was used to measure the width of the actin network in the leading edge of migrating cells expressing EGFP-actin or EGFP–α-actinin-1. FRAP experiments were performed on a confocal microscope (LSM 510; Carl Zeiss, Inc.) with a 63× plan Apo 1.4 NA objective. EGFP fluorescence was eliminated by 30 bleach cycles at 100% intensity of the 488-nm argon laser. Recovery curves generated from photobleached areas of the same sizes and locations were sampled every 0.5 s for 42 s and corrected for overall photobleaching and were used to calculate h/2.

Motility assay

Cell motility was analyzed as described previously (Pourroy et al., 2006) except that pictures were collected for 150 min at 5-min intervals. Means of velocity were calculated using MetaMorph and Excel (Microsoft) software.

Pull-down assay, cell fractionation, and Western blotting

GST-RhoA, GST-Cdc42, GST-Rac1, and GST-RhoA L63 beads (Cytoskeleton, Inc.) were loaded with GDP or GTPγS according to the manufacturer’s instructions before overnight incubation with 2 μg of bacterially produced Memo and Western blot analysis. Cell body or lamellipodia-enriched fractions were obtained as described previously (Cho and Klemke, 2002) after the addition of 5 nM HRG to the lower chamber of a 3-μm pore Transwell (Costar) for 1 h. Antibodies directed against Memo (monoclonal antibody to amino acids 23–43), mDia1, Shc, Gsk3β (BD Biosciences), RhoA, RhoGD1a (Santa Cruz Biotechnology, Inc.), and Flag (Sigma-Aldrich) were used for Western blotting.

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Immunofluorescence microscopy

Cells grown on collagen I-coated coverslips were fixed in 4% formaldehyde and permeabilized in 0.2% Triton X-100 before the addition of antibodies directed toward Cdc42, RhoA, α-actinin-1 (Santa Cruz Biotechnology, Inc.), mDia1, Rac1, Arp3 (BD Biosciences), cortactin (provided by E. van Obbergen-Schilling, National Center for Scientific Research, Nice, France), and α-tubulin (provided by W. Krek, Swiss Federal Institute of Technology, Zurich, Switzerland). Secondary antibodies and Alexa Fluor 546 phallolidin were obtained from Invitrogen. DNA was counterstained with Hoechst dye (Sigma-Aldrich). Images were recorded with a microscope (ApoTome IMAGER Z1; Carl Zeiss, Inc.) with a 63×plan Apo 1.4 NA objective coupled to a camera (AxioCamMRm; Carl Zeiss, Inc.) and driven by AxioVision LE software (Carl Zeiss, Inc.).

Statistical analysis

Data are presented as mean ± SEM and were analyzed by Mann-Whitney, Kruskal-Wallis, or permutation tests using StatXact software (Cytel) as indicated. P < 0.05 was considered statistically significant.

Online supplemental material

Fig. S1 shows the efficiency of the Memo, mDia1, and RhoA siRNAs and the effect of Memo depletion on lamellipodial markers. Fig. S2 shows that RhoA labeling of the cell membrane is not simply by ruffling and that Memo is required for plasma membrane localization of endogenous RhoA and mDia1. Fig. S3 shows that Memo is not required for plasma membrane localization of Rac1 and Cdc42. Video 1 shows MT outgrowth upon HRG treatment. Video 2 shows the lack of MT outgrowth in Memo-depleted cells. Videos 3 and 4 show adhesion site dynamics in control and Memo-depleted cells. Videos 5–8 show actin and α-actinin-1 localization in control and Memo-depleted cells. Videos 9 and 10 show EGFP-actin and EGFP-α-actinin-1 FRAP. Online supplemental material is available at http://www.jcb.org/cgi/content/full/jcb.200805107/DC1.

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