ADF/cofilin promotes invadopodial membrane recycling during cell invasion in vivo

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Introduction

Basement membrane is a thin, dense, sheet-like matrix that underlies or surrounds most tissues (Hohenester and Yurchenco, 2013). Cell invasion through basement membrane occurs in many normal and pathological processes, including embryo implantation, gastrulation, leukocyte trafficking, and cancer metastasis (Rowe and Weiss, 2008; Hagedorn and Sherwood, 2011). Because of its small pore size and cross-linked organization, cells are thought to require specialized mechanisms to transmigrate basement membrane (Madsen and Sahai, 2010). In particular, protrusive F-actin-based membrane structures, termed invadopodia, have been proposed to be critical for breaching basement membrane (Condeelis and Segall, 2003). Although extensively studied in transformed cells and cancer lines in vitro, the regulation of invadopodia in vivo has been difficult to establish due to the challenge of examining invasion in an intact organism (Beerling et al., 2011; Saltel et al., 2011).

Anchored cell (AC) invasion in Caenorhabditis elegans is a simple model of basement membrane transmigration that combines genetic analysis and imaging of the invasive cell–basement membrane interface (Hagedorn and Sherwood, 2011). The AC is a specialized uterine cell that invades through basement membrane to connect the developing uterine and vulval tissues during development (Sherwood and Sternberg, 2003). We have recently shown that F-actin–based AC invadopodia presage loss of F-actin dynamics and an inability to breach basement membrane. Optical highlighting indicated that UNC-60A disassembles actin filaments at invadopodia. Surprisingly, loss of unc-60a led to the accumulation of invadopodial membrane and associated components within the endolysosomal compartment. Photobleaching experiments revealed that during normal invasion the invadopodial membrane undergoes rapid recycling through the endolysosome. Together, these results identify the invadopodial membrane as a specialized compartment whose recycling to form dynamic, functional invadopodia is dependent on localized F-actin disassembly by ADF/cofilin.
its concentration relative to actin and other actin-binding proteins (Carlier et al., 1999; Van Troys et al., 2008). Alterations in ADF/cofilin activity affect formation of cellular protrusions, including neurite and lamellipodia extensions, and influence cell migration (Chen et al., 2001; Gungabissoon and Bamberg, 2003; Ghosh et al., 2004; Hotulainen et al., 2005; Kiuchi et al., 2007). Consistent with a possible role in cell invasion, cofilin is highly expressed in numerous metastatic cancers and tumor cell lines and promotes invadopodia stability in rat MTLn3 mammary tumor cells (Yamaguchi et al., 2005; Wang et al., 2007). Despite many studies, it remains unclear whether ADF/cofilin proteins primarily regulate actin assembly or disassembly during protrusive cellular activity (Van Troys et al., 2008). Moreover, it is unknown if ADF/cofilin proteins regulate invadopodia in vivo.

We find here that the C. elegans ADF/cofilin orthologue UNC-60A is expressed within the AC, localizes to AC invadopodia, and is required for AC invasion. RNAi-mediated depletion of the unc-60a transcript caused a reduction in F-actin disassembly, resulting in a pronounced expansion in F-actin at AC invadopodia. Strikingly, loss of unc-60a caused a specific disruption in the trafficking of the invadopodial membrane from the endolysosome to the invasive cell membrane, inhibiting invadopodia formation. Together our findings indicate that ADF/cofilin is a critical regulator of invadopodia in vivo, and support a model where ADF/cofilin directs localized F-actin filament turnover that promotes the trafficking of invadopodial membrane to the plasma membrane.

Results

UNC-60A (ADF/cofilin) is a critical cell-autonomous regulator of AC invasion

Recent work has shown that invadopodia form within the C. elegans uterine AC and mediate basement membrane penetration as the AC invades the vulval tissue (Fig. 1 A; Hagedorn et al., 2013). To identify direct regulators of AC invadopodia, we examined genes identified in a whole-genome RNAi screen for AC invasion (Matus et al., 2010). Of these, we noted that loss of unc-60a, one of two isoforms of the C. elegans orthologue of the ADF/cofilin gene (McKim et al., 1994; Anyanful et al., 2004), resulted in one of the most penetrant AC invasion defects. An expanded analysis of RNAi-mediated knockdown of unc-60a confirmed this finding, revealing a 90% AC invasion defect (Fig. 1 B; n = 14/140 [10%] normal invasion, 54/140 [39%] partial invasion, 72/140 [51%] no invasion; see Materials and methods for scoring details). unc-60 (sul158) mutants, which are null for the muscle-specific isoform unc-60b, have normal AC invasion (Ono et al., 1999; Ono et al., 2003; Ziel et al., 2009), indicating that the unc-60a isoform regulates AC invasion.

To examine where unc-60a (ADF/cofilin) is expressed, we generated a translational GFP::UNC-60A fusion protein under the control of its endogenous promoter (unc-60 > GFP::unc-60a). At the time of invasion GFP::UNC-60A was present throughout the uterine tissue and at elevated levels in the AC (Fig. 1 C; 1.35-fold greater levels versus neighboring uterine cells; n = 11 animals examined). GFP::UNC-60A was also more enriched at the invasive cell membrane (1.29-fold enrichment vs. 0.75-fold for neighboring uterine cells; n = 11 animals examined). To determine if ADF/cofilin functions within the AC, we generated a construct in which the unc-60 (ADF/cofilin) transcript from the related nematode Caenorhabditis briggsae was expressed specifically in the AC (cdh-3 > GFP::Cbrunc-60a). The C. briggsae unc-60 and C. elegans unc-60a genes share 83% nucleotide identity over the conserved first four exons (nucleotides 1–427 of C. elegans unc-60a), which is below the 95% threshold required for RNAi targeting (Rual et al., 2007). cdh-3 > GFP::Cbrunc-60 restored invasion in animals treated with C. elegans unc-60a RNAi (Fig. 1 C; n = 72/86 [84%] normal invasion, 12/86 [14%] partial invasion, 2/86 [2%] no invasion). Hence, UNC-60A promotes AC invasion cell autonomously.

UNC-60A (ADF/cofilin) localizes to AC invadopodia

To visualize the subcellular localization of UNC-60A, we generated animals with AC-specific GFP::UNC-60A expression (cdh-3 > GFP::unc-60a). GFP::UNC-60A localized in punctate structures at the AC–basement membrane interface before invasion (Fig. 1 D). To determine whether these structures were invadopodia, we examined their localization relative to F-actin and basement membrane breaches. We observed a tight correlation between F-actin localization and UNC-60A enrichment at the AC–basement membrane interface (Fig. 1 D), as well as localization to the invadopodium that breached the basement membrane (5/5 animals; Fig. 1 E). Further, UNC-60A localization to the AC–basement membrane interface was dependent on the integrin INA-1/PAT-3, whose activity is required for invasion and promotes the localization of other invadopodia constituents (Fig. 1 F; Hagedorn et al., 2009, 2013). Thus, UNC-60A is a component of AC invadopodia.

UNC-60A promotes F-actin turnover at AC invadopodia

To determine whether UNC-60A regulates AC invadopodia, we examined F-actin within the AC after loss of unc-60a. In unc-60a (RNAi) animals we observed a twofold expansion in the total volume and amount of the F-actin network at the AC–basement membrane interface, suggesting UNC-60A promotes actin filament turnover (Fig. 2 A). Although F-actin remained polarized, it was no longer organized into discrete foci and instead formed larger structures that failed to penetrate basement membrane (Fig. 2 B). Compared with F-actin structures in wild-type ACs, which had a median diameter of 0.96 µm, in the absence of ADF/cofilin, F-actin structures had a median size of 1.90 µm (n = 30 structures from 10 animals for each; P < 0.0001). These larger F-actin structures were also more stable. In 20-min time-lapse movies of wild-type animals (n = 4 animals), the average lifetime of invadopodia was only 88 s (Fig. 2 D; n = 207 structures examined). In contrast, in unc-60a (RNAi) ACs (n = 10 animals), most F-actin structures (n = 43/51) persisted for the entire time-lapse (20 min; Fig. 2 C and Videos 1–3). This duration far exceeded that of invadopodia in wild-type animals that are associated with basement membrane breaching events (Fig. 2 D; average lifetime of 6.3 min; n = 10 invadopodia from 8 animals).
region of actin (F-actin and likely a small portion of G-actin) at the invasive cell membrane where invadopodia reside (Fig. 3 A). In control animals, optically highlighted actin began redistributing immediately and spread throughout the AC by 30 min, consistent with rapid F-actin turnover (Fig. 3, A and B; n = 5/5 ACs). In contrast, after loss of unc-60a, optically highlighted actin did not disperse significantly, even after 30 min (Fig. 3, A and B; n = 4/4). To test if the accumulation of F-actin resulted from reduced F-actin disassembly, we created animals that expressed an actin isoform tagged with Dendra2 in the AC. Dendra2 is a photoconvertible fluorophore that switches from green to red fluorescence after brief exposure to 405-nm light (Gurskaya et al., 2006). We treated the Dendra2::ACT-1 animals with either unc-60a RNAi or an empty vector control and then photoconverted a small
not enriched at the plasma membrane or the expanded F-actin network. Instead, these molecules were localized within intra-cellular vesicles (Fig. 4, B and E). Colocalization analysis of PI(4,5)P2 and MIG-2 indicated that these molecules were within overlapping vesicles (Fig. 4 C), suggesting there is coordinated trafficking of invadopodial membrane to the plasma membrane regulated by UNC-60A.

To better understand invadopodial membrane trafficking, we examined markers for the early (Rab5), late (Rab7), and recycling (Rab11) endosomes, and the endolysosome (LMP-1 and CUP-5; Balklava et al., 2007; Campbell and Fares, 2010; Humphries et al., 2011). This analysis revealed that the endolysosomal markers LMP-1 and CUP-5 were polarized at the invasive membrane (Fig. S2). Consistent with a normal role in trafficking through this compartment, the invadopodial membrane components

**Figure 2.** **UNC-60A regulates invadopodial membrane recycling from the endolysosome.** To further examine UNC-60A’s role in AC invadopodia, we analyzed the localization of other invadopodia components after loss of unc-60a. Consistent with localization to the actin cytoskeleton (Trichet et al., 2008), the actin regulator UNC-34 (Ena/VASP) was distributed throughout the large F-actin aggregates that formed after loss of unc-60a (Fig. 5, A and E). In contrast, the invadopodial membrane component PI(4,5)P2 and the membrane-associated Rac proteins MIG-2 and CED-10, which colocalize with F-actin at invadopodia in wild-type animals (Fig. S1), were not enriched at the plasma membrane or the expanded F-actin network. Instead, these molecules were localized within intracellular vesicles (Fig. 4, B and E). Colocalization analysis of PI(4,5)P2 and MIG-2 indicated that these molecules were within overlapping vesicles (Fig. 4 C), suggesting there is coordinated trafficking of invadopodial membrane to the plasma membrane regulated by UNC-60A.

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revealed over a twofold decrease in LMP-1::GFP recycling, indicating a reduction in trafficking (Fig. 5 A, Fig. S3, and Video 4). Time-lapse analysis of PI(4,5)P₂ also confirmed these vesicles were more static (Fig. 5 B and Video 5). We conclude that UNC-60A promotes dynamic invadopodial membrane recycling from the endolysosome to the plasma membrane.

Other polarized trafficking and secretion events are normal after loss of unc-60a
To determine whether the disruption in invadopodial membrane trafficking was an indirect effect from a general perturbation in vesicle trafficking or polarity, we examined markers of polarity and secretion. Notably, after depletion of unc-60a the integrin receptor INA-1/PAT-3 and the netrin receptor UNC-40 (DCC) were polarized normally to the AC’s invasive cell membrane (Fig. 6, A and D; Hagedorn et al., 2009; Ziel et al., 2009). Further, AC deposition of the matrix component hemicentin into the basement membrane (Fig. 6, B and D; Sherwood et al., 2005) and secretion of the EGF-like ligand LIN-3, which induces vulval development,
 UNC-60A (ADF/cofilin) regulates the invadopodial membrane. (A) In contrast to its tight localization at the invasive cell membrane in wild-type animals (arrowhead), GFP::UNC-34 (Ena/VASP) localizes to the expanded F-actin network (bracket) after unc-60a RNAi treatment. (B) The invadopodial membrane components PtdIns(4,5)P_2 (mCherry::PLCδ) and the Racs GFP::MIG-2 and GFP::CED-10 were no longer localized to the invasive cell membrane after loss of unc-60a and were found within intracellular vesicles (arrowheads) independent of the expanded F-actin (brackets; r-value reports average Pearson’s correlation coefficient for colocalization within the AC ± SEM). (C) The invadopodial membrane components PtdIns(4,5)P_2 and GFP::MIG-2 colocalized within the same vesicles (arrowheads) after loss of unc-60a [r-value reports average Pearson’s correlation coefficient for colocalization within the AC ± SEM]. (D) The endolysosome marker LMP-1::GFP localized to invadopodial structures and at the initial breach site (top, arrowhead). LMP-1::GFP colocalized with invadopodial membrane markers [GFP::MIG-2, GFP::CED-10, and PtdIns(4,5)P_2] in wild-type ACs (middle panels, arrowheads) and after loss of unc-60a within internalized vesicles (bottom, arrowheads). r-value reports average Pearson’s correlation coefficient for colocalization within the AC ± SEM. (E) Graphs show the volume of GFP::UNC-34 and the polarized localization of the invadopodial membrane components in wild-type (black) and unc-60a (RNAi) animals (orange; n ≥ 6 animals examined for each; *, P < 0.01; ***, P < 0.0001; Student’s t test; error bars indicate ±SEM). Bars, 5 µm.

Discussion

Invadopodia, protrusive F-actin–based membrane structures capable of ECM remodeling, were identified in transformed cells and cancer lines in vitro over 20 years ago (Chen, 1989). Despite intense interest, the regulation and function of invadopodia in vivo has been elusive (Beerling et al., 2011; Murphy and Courtneidge, 2011). Our data indicate that the C. elegans ADF/cofilin orthologue UNC-60A is a critical and specific regulator of invadopodia during AC invasion. We find that UNC-60A mediates F-actin disassembly at AC invadopodia, which is critical for normal invadopodia dynamics and the trafficking of invadopodial membrane and associated membrane components to the plasma membrane (Fig. 6 E).

Our live-cell imaging and site-of-action studies indicate that UNC-60A specifically localizes to AC invadopodia and functions within the AC to promote breaching the basement membrane. The increased accumulation of F-actin and dramatically decreased actin turnover after loss of unc-60a indicate that UNC-60A promotes F-actin disassembly at invadopodia, rather than nucleating new actin filaments. Our data add weight to the notion that in most cell types ADF/cofilin contributes to actin dynamics primarily by mediating actin filament disassembly (Hotulainen et al., 2005; Kiuchi et al., 2007; Okreglak and Drubin, 2007; Lai et al., 2008). In the C. elegans AC, F-actin disassembly might be...
necssary for breaking down existing AC invadopodia and forming new invadopodia through the recycling of actin monomers. Dynamic F-actin turnover at AC invadopodia is also likely critical to generate the protrusive force required to breach basement membrane. Consistent with this idea, ADF/cofilin proteins are thought to generate actin monomers to promote lamellipodial extensions in Cos and MCF7 cell lines and migration in NIH 3T3 and B16F1 cells (Hotulainen et al., 2005; Kiuchi et al., 2007).

Ultrastructural analysis and TIRF microscopy have indicated that invadopodia in tumor cells have a dynamic, morphologically unique membrane (Baldassarre et al., 2003; Artym et al., 2011). Studies on the trafficking of invadopodial membrane components in cell culture have focused on the membrane-anchored matrix metalloproteinase MT1-MMP (Poincloux et al., 2009). Intriguingly, recent work in tumor cells has indicated that MT1-MMP is recycled from the endolysosomal compartment to the plasma membrane (Hoshino et al., 2012; Yu et al., 2012). Our studies indicate that the invadopodial membrane is a unique compartment, which is actively recycled through the endolysosome in an UNC-60A (ADF/cofilin)–dependent manner (Fig. 5 E). Trafficking through the endolysosome might facilitate dynamic membrane addition to invadopodia required for active protrusions, similar to roles of endolysosome membrane addition in wound healing and neurite outgrowth (Reddy et al., 2001; Arantes and Andrews, 2006). Active membrane movement through the endolysosome may also be required for delivery of proteases to AC invadopodia (Yu et al., 2012).

Spatially restricted cofilin activity is necessary for vesicular trafficking of acetylcholine receptors to nascent postsynaptic sites as well as planar cell polarity proteins to apical membranes in the mouse embryo (Lee et al., 2009; Mahaffey et al., 2013). It is thought that localized F-actin turnover by ADF/cofilin facilitates vesicle targeting or fusion to the plasma membrane, perhaps by breaking the cortical actin barrier (Etizen, 2003; Lee et al., 2009; Mahaffey et al., 2013). Thus, targeted F-actin turnover by UNC-60A at AC invadopodia might direct trafficking of invadopodial membrane from the endolysosome to discrete sites to form invadopodia. Cofilin has also been implicated in other trafficking events within cells, including sorting at the trans-Golgi network and endocytic trafficking to the yeast vacuole (Okreglak and Drubin, 2007; von Blume et al., 2009). In the trans-Golgi, cofilin is proposed to mediate trimming of F-actin, which allows the transmembrane Ca2+ ATPase SPCA1 to import Ca2+ into the lumen to direct proper sorting (von Blume et al., 2011). During endocytic delivery of vesicles to the yeast vacuole, cofilin is required to disassemble F-actin from the endocytic vesicles and regenerate the actin monomer pool, both of which might be required for late endocytic targeting or fusion with the vacuole (Okreglak and Drubin, 2007). It is thus possible that cofilin might regulate aspects of invadopodial membrane targeting, fusion, or sorting at the endolysosome itself that are required for trafficking back to the invasive membrane to form invadopodia. Given the tight localization of cofilin to invadopodia at the invasive cell membrane, however, we favor the notion that cofilin has a direct role in targeting invadopodial membrane to nascent invadopodia. Taken together, our results establish a unique organizing function for UNC-60A (cofilin) at invadopodia, coordinating both protrusive F-actin formation and invadopodial membrane addition.

Vertebrate ADF/cofilin proteins are overexpressed in many cancers and mediate invadopodia maturation in an adenocarcinoma cell line (Wang et al., 2007). The results of our functional in vivo studies here strongly support the notion that ADF/cofilin proteins are conserved regulators of invadopodia in both normal and pathological processes. Further, the specific defect in AC invadopodia formation and strong perturbation in ability to invade suggests that targeting ADF/cofilin activity might be an effective strategy to block invasion in diseases such as cancer.
reported as full width at half-maximum (FWHM) was measured by imaging fluorescent beads (210 nm; channel 488) on both microscopy systems used in this study: (1) spinning-disk confocal; XY (lateral) 287 nm, Z (axial) 571 nm; and (2) laser-scanning confocal; XY (lateral) 333 nm, Z (axial) 772 nm. Acquired images were processed using ImageJ 1.40g (National Institutes of Health) and Photoshop (CS6 Extended; Adobe Systems, Inc.). 3D reconstructions were built from confocal Z-stacks, analyzed, and exported as .MOV files using IMARIS 7.4 (Bitplane, Inc.). Figures and graphs were constructed using Illustrator (CS6 Extended; Adobe Systems, Inc.). Videos were annotated using Photoshop.

Materials and methods

Strains and culture conditions
C. elegans were cultured as described previously (Brenner, 1974) and wild-type were strain N2. In the text and figures, linkage to a promoter is designated with “>” and linkages that fuse open reading frames with “::”.

The following alleles and transgenes were used: qyEx411 [zmp-1 > lmp-1::mCherry], qyEx403 [cdh-3 > GFP::cup-5], qyIs61 [cdh-3 > GFP::unc-34], qyEx237 [unc-60 > GFP::unc-60a], qyIs223 [cdh-3 > GFP::unc-60a], qyIs221 [cdh-3 > GFP::unc-60a], qyIs224 [cdh-3 > GFP::Cbrunc-60], qyIs219 [cdh-3 > GFP::PLCδPH], qyIs220 [cdh-3 > GFP::mig-2], qyIs67 [cdh-3 > unc-40::GFP], qyIs127 [cdh-3 > mCherry::PLCδPH], qyIs23 [cdh-3 > mCherry::GFP], qyEx282 [cdh-3 > Dendra2::act-1].

Microscopy, image acquisition, processing, and analysis
Images were acquired using a spinning-disk confocal microscope (CSU-10; Yokogawa Corporation of America) mounted on a microscope (AxioImager; Carl Zeiss) with a 100× Plan Apochromat objective (1.4 NA) and controlled by iVision software (Biovision Technologies). The point spread function (PSF) reported as full width at half-maximum (FWHM) was measured by imaging fluorescent beads (210 nm; channel 488) on both microscopy systems used in this study: (1) spinning-disk confocal; XY (lateral) 287 nm, Z (axial) 571 nm; and (2) laser-scanning confocal; XY (lateral) 333 nm, Z (axial) 772 nm.

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Scoring of AC invasion, fluorescence intensity, colocalization, and polarity

AC invasion was scored as previously described using DIC microscopy (Sherwood et al., 2005). In brief, animals were scored for invasion at the P6.p four-cell stage when basement membrane clearance is completed in wild-type animals. ACs were scored as “normal invasion” if the breach in the basement membrane was at least the width of the AC by the P6.p four-cell stage, “partial invasion” if there was a visible breach smaller than the width of the AC, and “no invasion” if there was no detectible breach in the basement membrane. For analysis of UNC-60A up-regulation in the AC, regions of interest were drawn in ImageJ around the AC and then a neighboring uterine cell in the same animal; mean fluorescence intensity of each cell was then determined. Up-regulation was calculated as the following ratio: [AC mean intensity – background]/[neighboring uterine cell mean intensity – background]. Colocalization analysis was performed on confocal
z-sections using the “Coloc” module in Imaris. In brief, a mask was first used to exclude all signal outside of the AC. Pearson’s correlation coefficient was then calculated from the GFP and mCherry fluorescence inside the AC (n ≥ 5 animals for each comparison). All images were acquired using identical settings. Polarity in wild-type animals and animals treated with unc-60a RNAi was determined using the ratio of the mean fluorescence intensity from a 5-pixel-wide line scan drawn along the invasive and apical/lateral membranes of the ACs and in some cases neighboring uterine cells expressing unc-60a > GFP:unc-60a::MG-2, GFP::CED-10, MPM-1:: GFP, mCherry::PLC8, mCherry::RAB-5, mCherry::RAB-7, and mCherry::RAB-11, GFP::CUP-5, PAT-3::GFP, and UNC-40::GFP. Numbers examined for each are indicated in the text or figure legends. Polarity was calculated as the following ratio: [invasive membrane mean intensity – background]/[apical/lateral membrane mean intensity – background].

Scoring of F-actin volume and dynamics
Quantitative F-actin measurements were performed as described previously (Hagedorn et al., 2009). In brief, confocal Z-stacks were used to make 3D reconstructions of F-actin networks in ACs expressing the F-actin probe mCherry::moeABD using Imaris 6.0 (Bitplane, Inc.). Surface renderings of mCherry::moeABD were created setting a threshold that outlined the dense F-actin network at the invasive membrane in wild-type ACs. This same thresholding was used in cdh-3 > mCherry::moeABD untreated and unc-60a RNAi-treated animals. Quantitative measurements were then made for the volume and amount of fluorescence intensity with these isosurfaces rendered (n > 10 animals for each). Similar methods were used to quantify GFP::UNC-34 and hemicentin::GFP volumes. Quantitative measurements of the total (sum) fluorescence of mCherry::moeABD present in the AC showed no change after treatment with unc-60a RNAi (n = 10 animals each; P = 0.91; Student’s t-test). AC invadopodia dynamics were quantified using the “Spots” module within Imaris, which facilitated the tracking of F-actin patches (cdh-3 > mCherry::moeABD) at the invasive membrane of the AC over time in wild-type and unc-60a RNAi-treated animals.

Optical highlighting and fluorescence loss in photobleaching (FLIP)
Imaging was conducted with a laser-scanning confocal microscope (LSM 510; Carl Zeiss) equipped with a 100× objective (1.4 NA). The PMT light detection wavelength was measured as follows: [ROI background]/[outside ROI background]. For FILP, selected ROIs (~1 µm) of the AC in the worms expressing cdh-3 > MPM-1::GFP were bleached every 30 s for 10 min at 100% power for 20 iterations, and images were taken before and after each bleaching interval. Quantitative data were obtained with ImageJ by normalizing background-subtracted mean intensities to t = 0 of the LMP-1::GFP signal at the invasive membrane adjacent to the bleached ROI. For each technique, identical settings were used to acquire images at all time-points and ≥5 animals were analyzed in each group.

RNAi
RNAi targeting pat-3 and unc-60a was delivered by RNAi feeding (Sherwood et al., 2005). To bypass embryonic lethality, synchronized L1-stage larvae were plated on unc-60a RNAi; and synchronized L2-stage larvae were plated on pat-3 RNAi as described previously (Hagedorn et al., 2009). The empty RNAi vector L4440 was used as the wild-type control.

Construction of GFP protein fusions
The GFP::unc-60a fusions were constructed by placing GFP immediately after the unc-60a start codon and 1.9 kb downstream of the stop codon was included as the 3’ UTR. The GFP::UNC-34 ampiclon was linked by PCR fusion to promoters, either endogenous (fragment encompassing 3.6 kb upstream of the unc-60a start codon, unc-60a > GFP::unc-60a) or AC specific (cdh-3 > promoter, cdh-3 > GFP) respectively. The Dendra2::act-1 fusion was constructed by cloning Dendra2 at the N terminus of a 1.7 kb genomic fragment of the C. elegans act-1 gene (fragment encompassing act-1 start codon and downstream 3’ UTR). The AC-specific cdh-3 > promoter was then cloned immediately upstream of the Dendra2 start codon. The Imp-1::GFP: GFP::cup-5, mCherry::rab-5, mCherry::rab-7, and mCherry::rab-11 reporters were amplified from plasmid DNA (see Table S2) and linked to the cdh-3 > promoter by PCR fusion. Constructs were co-injected with 50 ng/µl unc-119 rescue DNA 50 ng/µl pBSK, and 50 ng/µl EcoR1 cut salmon sperm DNA into unc-119[ed4] hermaphrodites. Extrachromosomal lines were established and integrated by gamma irradiation as described previously (Sherwood et al., 2005). See supplemental material for transgenic strains generated (Table S1) and primer sequences used (Table S2).

Site-of-action studies with C. briggsae unc-60a rescue
The GFP::CbrUNC-60 protein in the AC. The confirmed C. briggsae unc-60 RNA was not depleted by C. elegans unc-60a RNAi.

Statistical analysis
All statistical analysis was performed in JMP version 9.0 (SAS Institute). Figure legends specify which test was used.

Online supplemental material
Online supplemental material includes three figures, five videos corresponding to the time-lapse data presented in the main text figures, and two tables: Table S1 for extrachromosomal arrays and integrated strains and Table S2 for primer sequences. Online supplemental material is available at http://www.jcb.org/cgi/content/full/jcb.201312098/DC1. Additional data are available in the JCB DataViewer at http://dx.doi.org/10.1083/jcb.201312098.

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References

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