Macrophage filopodia, finger-like membrane protrusions, were first implicated in phagocytosis more than 100 years ago, but little is still known about the involvement of these actin-dependent structures in particle clearance. Using spinning disk confocal microscopy to image filopodial dynamics in mouse resident Lifeact-EGFP macrophages, we show that filopodia, or filopodia-like structures, support pathogen clearance by multiple means. Filopodia supported the phagocytic uptake of bacterial (Escherichia coli) particles by (i) capturing along the filopodial shaft and surfing toward the cell body, the most common mode of capture; (ii) capturing via the tip followed by retraction; or (iii) sweeping mode of capture; (ii) capturing via the tip followed by retraction; or (iv) sweeping mode of capture. In addition, filopodia supported the uptake of zymosan (Saccharomyces cerevisiae) particles by (i) providing fixation, (ii) capturing at the tip and filopodia-guided actin anterograde flow with phagocytic cup formation, and (iii) the rapid growth of new protrusions. To explore the role of filopodia-inducing Cdc42, we generated myeloid-restricted Cdc42 knock-out mice. Cdc42-deficient macrophages exhibited rapid phagocytic cup kinetics, but reduced particle clearance, which could be explained by the marked rounded-up morphology of these cells. Macrophages lacking Myo10, thought to act downstream of Cdc42, had normal morphology, motility, and phagocytic cup formation, but displayed markedly reduced filopodia formation. In conclusion, live-cell imaging revealed multiple mechanisms involving macrophage filopodia in particle capture and engulfment. Cdc42 is not critical for filopodia or phagocytic cup formation, but plays a key role in driving macrophage lamellipodial spreading.

Phagocytosis is a specialized form of endocytosis, which requires localized actin polymerization to engulf large particles (>~0.5 μm), and initially involves particle binding to phagocytic receptors, which triggers phagocytic cup formation through the activation of kinases, such as Syk, and Rho GTPases, such as the Cdc42 and Rac subfamilies (1). Although phagocytosis has been investigated in-depth, the involvement of filopodia (finger-like projections containing bundled actin filaments (2)) in actually capturing and clearing particles has not been conclusively demonstrated, except for the historical observations of Metschnikoff (3) and more recent work using brightfield microscopy (4–7).

Young et al. (4) observed using time-lapse differential interference contrast (DIC) imaging that Escherichia coli expressing invasin, a transmembrane protein of Yersinia pseudotuberculosis, could enter Hep-2 (HeLa-derived) cells via filopodia. Using another approach, coating of magnetic microbeads with invasin, Vonna et al. (5) found that the adhesion of beads to filopodial tips induced pulling toward the cell body. Similarly, Kress et al. (6) reported that filopodia act as “phagocytic tentacles” and pulled IgG-coated beads in an optical trap with discrete steps, suggesting that a motor protein was involved. More recently, Möller et al. (7) reported that macrophages can use filopodia to lift membrane-bound bacteria, thereby enabling a lamellipodial protrusion to engulf the target in a so-called hook-and-shovel mechanism. In this study, we looked at phagocytosis from a new angle. We isolated resident peritoneal macrophages from transgenic Lifeact-EGFP mice (8) and used time-lapse spinning disk confocal microscopy to visualize the involvement of filopodial dynamics in the clearance of fungal and bacterial particles. Lifeact-EGFP mice ubiquitously express the F-actin-binding (17-amino acid) peptide Lifeact fused to the 238-amino acid fluorescent protein EGFP (9). The filopodia-inducing proteins Cdc42 and Myo10 have both been implicated in phagocytosis (10, 11). Using knock-out mouse models, we explored the roles of these proteins in macrophage phagocytosis.

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This article contains supplemental Videos S1–S9.

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Results

Capture of zymosan particles by filopodial tentacles

Initially, we used superresolution-structured illumination microscopy (SR-SIM) to confirm that Lifeact-EGFP is expressed in resident peritoneal macrophages and localizes to F-actin (Fig. 1, A and B). Whereas Lifeact-EGFP and the high-affinity F-actin probe phalloidin, conjugated to Alexa Fluor 594, clearly co-localized, Lifeact-EGFP additionally accumulated in the nucleus. Similar to Lifeact-EGFP, we previously observed that EGFP alone also accumulated in the nuclei of resident peritoneal macrophages (isolated from LysM-EGFP (Lyz2tm1.1Graf) mice (12)). SR-SIM imaging provided striking detail of the actin architecture in macrophages (Fig. 1C). Instead of a dense branching network, typical for the lamellipodial membrane protrusions of fibroblast-like and epithelial cells (13, 14), SR-SIM imaging showed nicely that macrophage lamellipodial membrane protrusions, as well as the cell body, are rich in arcs of actin bundles orientated parallel to the cell periphery. In addition, we observed that one or more prominent filopodial tentacles, or even a cluster of tentacles, typically extend from the cell periphery.

Using 2-channel spinning disk confocal microscopy, we found that the tips of filopodial tentacles were capable of capturing fluorescent zymosan particles (Fig. 2, A–C; supplemental Videos S1–S3). Filopodial extensions could also be observed arising from closed phagocytic cups in snapshots of phagocytic events captured by fixation and scanning electron microscopy (Fig. 2D). While performing spinning disk confocal microscopy experiments, we invariably observed dynamic actin “hot spots” within the actin ring (corresponding to the phagocytic cup) surrounding ingested zymosan particles, and we speculated that these spots may correspond to transient filopodial tentacles emerging from the phagocytic cup. Instead, these hot spots, which accompany each uptake event, were variously located beneath, alongside, and above zymosan particles, and presumably serve as local pools of actin.

We assumed that zymosan uptake in our assays was triggered via the recognition of β-glucans by Dectin-1 (15), although both TLRs (Toll-like receptors) (16) and the MR (mannose receptor) (17, 18) have been implicated in zymosan phagocytosis. To assess the contribution of these receptors to zymosan phagocytosis in our assays, we compared the phagocytic activity of wild-type (WT) versus Toll-like receptor-2/4 double knock-out (dKO), mannose-receptor knock-out, or Dectin-1 knock-out macrophages. Using zymosan particles labeled with the pH-sensitive dye pHrodo, which becomes brighter at decreasing pH (Fig. 3, A and B), we could measure phagosome acidification, a readout for particle internalization. One caveat of this approach is that a subset of macrophages exhibits weak phagosome acidification (for example, see Fig. 3C). Nevertheless, using this approach, we could confirm that Dectin-1 is the primary recep-

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**Figure 1. SR-SIM imaging of actin filaments in macrophages.** A, PCR genotyping of offspring derived from crossing heterozygous Lifeact-EGFP and wild-type (WT) mice. B, resident peritoneal macrophage isolated from a Lifeact-EGFP mouse. After fixation and permeabilization, the cell was stained with Alexa Fluor 594 (AF594)-conjugated phalloidin, an F-actin probe, and imaged by SR-SIM. The single channel images have been inverted. Scale bar, 10 μm. C, SR-SIM (inverse) image of fixed and permeabilized mouse macrophages, stained with Alexa Fluor 488 (AF488)-conjugated phalloidin. The image, obtained via an oil-immersion ×63/NA 1.4 objective lens, is an overlay of 3 z-slices taken at 220 nm steps. Scale bar, 10 μm.
tor mediating both the binding and internalization of zymosan particles (Fig. 3D).

Multiple modes of filopodial tentacles in bacterial particle clearance

Next, we investigated the involvement of filopodial tentacles in the capture of E. coli particles (Fig. 4). Time-lapse spinning disk confocal microscopy revealed that filopodial tentacles were highly effective at clearing particles beyond the cell body. Once captured by the tips of filopodia, particles were retracted toward the cell body and internalized, or, alternatively, the sweeping of a tentacle was effective at bringing particles toward the cell body (Fig. 4, A and B; supplemental Videos S4 and S5). The growth of new filopodial tentacles, fed by a pool of actin, could be observed in time-lapse recordings (Fig. 4A; supplemental Video S4). In addition to retraction and sweeping, the surfing of particles along the shaft of filopodial tentacles was frequently observed (Figs. 4C and 5, A and B; supplemental Videos S6–S8). Particles surfed (moved) along filopodia with a velocity of 1–2 μm/min (Fig. 5C). Combinations of surfing and retraction were also frequently observed (supplemental Video S9). As illustrated in Fig. 5D, surfing and combinations of surfing and retraction accounted for the majority of filopodia-mediated E. coli particle uptake events. We speculated that actin retrograde flow may drive the surfing of particles (see "Discussion"). Using kymograph analyses we found that the velocity of actin retrograde flow (treadmilling) in Lifeact-EGFP macrophages was in the range 2–4 μm/min (Fig. 5E), consistent with previous reports (19, 20).

Cdc42-deficient macrophages

To investigate the role of Cdc42 in macrophage filopodial tentacle formation and phagocytosis we generated myeloid-restricted Cdc42 (conditional) knock-out (cKO) mice (Fig. 6). Conditional deletion was confirmed by Western blot analysis of lysates from F4/80+ cells (macrophages), purified by cell sorting (Fig. 6A). Spinning disk confocal microscopy of living cells (not shown in Fig. 6, but see Fig. 13D) and superresolution imaging of fixed cells stained with an F-actin probe, Alexa Fluor 488-conjugated phalloidin, showed that Cdc42-deficient macrophages had a severe cell spreading defect (Fig. 6B), reminiscent of Myo9b-deficient macrophages, which have overactive RhoA activity (21). Indeed, Cdc42-deficient macrophages were resistant to cell spreading, even in the presence of the Toll-like receptor 4 ligand lipopolysaccharide (LPS), used to activate cells, as shown in Fig. 6C and quantified in Fig. 6D.
where the projected cell area is used as an index of spreading. Unstimulated Cdc42 cKO macrophages exhibited less filopodia per cell compared with WT cells (Fig. 6E). However, LPS stimulation for 90 min increased the number of filopodia per cell in both WT and Cdc42 cKO cells, such that there was no significance difference between activated (+LPS) WT and Cdc42

**Figure 3. Dectin-1 mediates zymosan binding and internalization.** A, fluorescence emission spectra of pHrodo-conjugated zymosan particles suspended in buffers with different pH values, ranging from pH 4 to 8. Particles were labeled using the succinimidyl ester of pHrodo. B, top panel: DIC and fluorescence images of pH-sensitive pHrodo-conjugated zymosan particles superfused with media buffered to different pH values, as indicated. Bottom panel: time-lapse images showing the ingestion and acidification of pHrodo-conjugated zymosan particles. C, DIC image, with superimposed red fluorescence, of two wild-type (WT) macrophages, which have ingested multiple pHrodo-conjugated zymosan particles. Red fluorescence indicates acidification of phagosomes. Scale bar, 10 μm. D, box plots of the number of attached (left) and red fluorescent (right) pHrodo-conjugated zymosan particles in WT, Toll-like receptor 2/4 double knock-out (TLR2/4 dKO), mannose receptor knock-out (MR/−/−), and Dectin-1 knock-out (Dectin-1/−/−) macrophages. Cells, seeded in fibronectin-coated μ-slide I chambers, were incubated with particles for 15 min before washing. *, p < 0.05; Kruskal-Wallis one-way analysis of variance on ranks (at the 0.05 level of significance).

**Figure 4. Multiple modes of filopodia-mediated capture and ingestion of bacterial particles.** A, time-lapse images (15 × 15 μm) showing the capture of an Alexa Fluor 594-conjugated E. coli (AF594-E. coli) particle (red) by a filopodium of a mouse resident peritoneal Lifeact-EGFP macrophage, followed by retraction toward the cell body and the growth of a new filopodial tentacle. B, time-lapse images (15 × 15 μm) showing the retraction of a filopodium with an E. coli particle on its tip, as well as the clearance of E. coli particles by the sweeping action of another filopodial tentacle. C, time-lapse images (20 × 20 μm) showing the fortuitous transfer of an E. coli particle from the filopodium of one cell to the filopodium of another cell. The particle is pulled toward the cell body by the first filopodium, then, following transfer, it appears to “surf” along the second filopodium.
Filopodia and phagocytosis

Figure 5. Filopodia-mediated uptake of bacterial particles. A, time-lapse images (20 × 20 μm) showing captured E. coli particles surfing along filopodia toward the cell body. B, another example of E. coli particles surfing along a filopodium (images are 20 × 20 μm). C, box plot of the velocity of E. coli particles surfing along filopodia. D, normalized frequency (all events = 100%) of different modes in filopodia-mediated phagocytic events. E, measurement of actin retrograde flow in Lifeact-EGFP macrophages by kymograph analysis.

In addition, the median length of filopodia was shorter in unstimulated Cdc42 cKO macrophages, compared with WT, and LPS stimulation significantly increased filopodia length in both WT and Cdc42 cKO macrophages (Fig. 6F). We could not detect the putative filopodia-inducing Rho GTPases RhoJ, RhoQ, RhoD, or RhoF in mouse macrophages using RT-PCR and Western blot analyses (Fig. 7, A and B). Thus, Cdc42 is presumably the predominant filopodia-inducing Rho GTPase in macrophages (22), and, in Cdc42-deficient cells, the paucity of filopodia, but not cell spreading, is compensated by LPS stimulation.

Using time-lapse spinning disk confocal microscopy and fluorescent labeling of the cell membrane with Alexa Fluor 488-conjugated anti-F4/80 antibodies, we imaged the engulfment of large (diameter, 5.19 μm) mouse IgG-coated polystyrene beads by macrophages (Fig. 8). Most beads were taken up after passive contact with the cell membrane. The kinetics of phagocytic cup formation was measured by tracking XY, XZ, and YZ views of the cells (Fig. 8). Surprisingly, Cdc42-deficient macrophages were highly efficient at engulfing large particles (Fig. 8A), and the rate of cup closure was faster compared with WT cells (Fig. 8, B–D). However, the start of phagocytic cup formation may have been underestimated in Cdc42-deficient cells due to the highly ruffled surface of these rounded-up cells (Fig. 9A). In contrast, the initiation of phagocytic cup formation could be more easily detected in spread-out WT macrophages, especially when particles landed on lamellipodia (Fig. 8B).

Nevertheless, whether first or second phagocytic half-cup kinetics were measured (Fig. 8D), Cdc42 cKO macrophages were faster at enveloping large beads. Notably, we occasionally observed that filopodia could haul in large particles before phagocytic cup formation was initiated (Fig. 8D), but these events were excluded from the above analyses in Fig. 8D.

Although Cdc42 cKO macrophages had faster phagocytic cup kinetics than WT cells, mutant cells engulfed less IgG-coated polystyrene beads and zymosan particles in end-point assays (Fig. 10A), presumably due to less frequent contacts (see “Discussion”). SR-SIM imaging of Cdc42-deficient macrophages, which were fixed while engulfing zymosan particles, revealed filopodia projecting from phagocytic cups (Fig. 10B).

Aside from phagocytosis (23), Cdc42 is implicated in chemotactic navigation (24–26). Using real-time chemotaxis assays, we found that Cdc42-deficient macrophages had greatly impaired chemotactic navigation and modestly decreased median cell velocity (Fig. 10, C and D). Chemotaxis assays were performed in a complement C5a gradient (target concentration, 20 nM) and a uniform concentration of LPS (0.1 μg/ml).

Myo10 is implicated in phagocytosis (11), especially in the engulfment of large particles, and it is thought to act downstream of Cdc42 in the generation of filopodia (27). We confirmed deletion of Myo10 in homozygous Myo10 reporter knock-out (Myo10tm2/tn2) mice using Southern and Western blot analyses, as well as PCR analysis (Fig. 11A).3 In contrast to Cdc42-deficient macrophages, Myo10 knock-out macrophages exhibited no defects in cell velocity or chemotactic navigation (Fig. 11B). Moreover, there were no impairments in phagocytic cup formation and the ingestion of large particles (Fig. 11, C–E). Consistent with the time-lapse imaging data, no differences in the uptake of particles could be detected in end-point assays (Fig. 11F). To test whether Myo10-deficient macrophages have impaired filopodia formation we performed dynamic assays because filopodia cannot be unambiguously distinguished from retraction fibers arising from lamellipodial membrane retraction (Fig. 12A). We challenged macrophages with fluorescent E. coli particles, without prestimulation with LPS, and counted newly extended filopodia over a 20-min recording period (Fig. 12, B–D). Filopodia were formed by Myo10-deficient macrophages, but the total number per cell was greatly reduced compared with wild-type cells (Fig. 12D).

In another series of experiments, we compared the rates of filopodia formation in WT, Myo10tm2/tn2, and Cdc42 cKO macrophages during time-lapse recordings, with or without

pretreatment with LPS (1 μg/ml for 45 min) (Fig. 13, A–D). Consistent with the findings in Fig. 12, filopodia formation was markedly impaired in Myo10-deficient macrophages. Moreover, comparison of the images in Fig. 13, C (Myo10<sup>−/−</sup>/LysM-Cre) and D (Cdc42 cKO), underscore the spreading defect of Cdc42-deficient macrophages (see also Fig. 6), as well as the importance of distinguishing retraction fibers from filopodial protrusions, especially in the case of Myo10-deficient cells.
Phagocytosis, the uptake of large particles, requires the force generated by actin polymerization to drive plasma membrane extensions around a target particle and form a phagocytic cup, followed by phagosome formation (28–30). However, before a particle can be ingested it first needs to come in contact with a phagocyte. In 2010, Flannagan *et al.* (31) reported that particle-phagocytic receptor interactions were not passive, stochastic events, but, instead, macrophages continually probe the environment using Rac-dependent membrane protrusions. Filopodia, finger-like structures containing actin bundles, are highly suitable for probing the environment (2), and can be induced by “switching on” Cdc42 via Cdc42 guanine nucleotide exchange factors (22). As proof of principle, stud-
ies using optical traps (tweezers) to present microbeads to macrophages showed that filopodia can pull beads (6), or, alternatively, beads can surf along filopodia toward the cell body (32). We asked what are the roles of filopodial dynamics in practice when primary macrophages are challenged with fungal or bacterial particles? In addition, we investigated the roles of the filopodia-inducing proteins Cdc42 and Myo10 in phagocytosis.

Figure 9. High-resolution image of a Cdc42-deficient (Cdc42 cKO) macrophage and filopodium-mediated capture of a large bead by a WT cell. A, scanning electron microscopy image of a Cdc42-deficient macrophage. Scale bar, 4 μm. B, time-lapse images showing capture of an IgG-coated polystyrene bead by a filopodium, followed by phagocytic cup formation. Scale bar, 10 μm.

Figure 10. Impaired particle clearance and chemotactic navigation of Cdc42-deficient macrophages. A, end-point assays showing the number of mouse IgG-coated, 5.19-μm diameter polystyrene beads (labeled with Alexa Fluor 594-conjugated secondary antibodies) or Alexa Fluor 594-conjugated zymosan particles ingested by WT and Cdc42 cKO macrophages, respectively, after 15 min incubation, followed by wash steps. B, snapshots of zymosan ingestion by Cdc42 cKO macrophages showing an extending phagocytic cup (left image) and filopodial tentacles emerging from a closed phagocytic cup (right image). The white arrows indicate filopodia. Macrophages were fixed, permeabilized, stained with Alexa Fluor 488-conjugated phalloidin, and imaged by SR-SIM. Scale bars, 5 μm. C, migration plots of WT and Cdc42 cKO macrophages in a chemotactic complement C5a gradient. D, summary box plots of velocity and chemotactic efficiency (chemotaxis index). *, p < 0.05; Mann-Whitney U test (n = 75 each group; 3 independent experiments).
We found that resident Lifeact-EGFP macrophages could capture moderately large (~3 μm diameter) zymosan particles beyond the cell body using filopodial tentacles. Particle binding triggered the flow of actin and membrane toward the particle, rather than simple retraction, such that filopodial tentacles widened and could support phagocytic cup formation, followed by internalization and new tentacle growth. A similar outward flow of the membrane, forming a pedestal, has been observed when a zymosan particle, initially held by a micropipette, is brought into contact with a neutrophil, held by a second micropipette (33). In contrast, no membrane protrusive effect was observed when neutrophils were presented with antibody-opsonized beads (33). Zymosan is a glucan prepared from yeast (*Saccharomyces cerevisiae*) and is recognized by TLR2/TLR6 (Toll-like receptor 2/Toll-like receptor 6) heterodimers and Dectin-1 (34, 35), and possibly also mannose receptors (36).

Using macrophages isolated from TLR2/TLR4 dKO, Dectin-1−/− and MR−/− mice, we confirmed that mouse macrophage
zymosan binding and phagocytosis are primarily mediated by Dectin-1. We infer that Dectin-1, akin to Fcγ-receptors, transduces phagocytic signaling through an immunoreceptor tyrosine-based activation motif-like domain (37), and evokes anterograde actin flow and phagocytic cup formation when a filopodium makes contact with a zymosan particle (see schematic diagram, Fig. 14A).

Filopodial tentacles were highly efficient at fishing out E. coli particles beyond the cell body. We observed four main mechanisms (in order of frequency): capture and surfing along the filopodium, sweeping actions, combinations of surfing and retraction, and simple retraction (see schematic summary, Fig. 14B). In contrast to studies using optical traps (6, 32), we did not bring particles to macrophage protrusions, but, rather, macrophages were required to seek and capture targets. Notably, time-lapse imaging was important to distinguish the formation and dynamics of filopodia from the formation of retraction fibers. On the one hand, the relative contribution of surfing to particle uptake may have been inflated because retraction fibers were variably present at the start of time-lapse recordings. On the other hand, retraction fibers are structurally related to filopodia and similarly contain parallel, bundled actin (38). Furthermore, Svitkina et al. (38) demonstrated using time-lapse imaging that filopodia and retraction fibers are interconvertible structures. E. coli particles surfed along filopodia with velocities of 1–2 μm/min, marginally slower than measured rates of retrograde flow (2–4 μm/min). Assuming that the surfing of bound particles is driven by actin retrograde flow, a mechanism linking particles to actin filaments is required. Unconventional myosins could fulfill this role, along the lines nicely illustrated in Fig. 3 of the review article by Mermall et al. (39), or possibly the ERM (ezrin, radixin, and moesin) protein family could play a role (40).

In the pioneering study by Nobes and Hall (22), the injection of constitutively active Cdc42, together with inhibitors of Rac and Rho, into Swiss 3T3 (embryonic fibroblast-derived) cells was shown to induce filopodial tentacles remarkably similar to those we observed in macrophages, whereas this effect was much weaker in the presence of Rac activity. It is now known that Cdc42 stimulates actin nucleation by the actin-related protein-2/3 (Arp2/3) complex via WASP (Wiskott-Aldrich syndrome protein), which is predominantly expressed in hematopoietic cells (41). The Cdc42-WASP-Arp2/3 pathway is thought to be important for the initiation of filopodia formation, but, importantly, other proteins can induce filopodia, such as RIF (RhoF), mammalian Diaphanous-like (mDia) formins, and Myo10 (2). In relationship to this point, we could not detect filopodia-inducing Rho GTPases, other than Cdc42, in macro-

Figure 12. Deletion of Myo10 decreases nascent filopodia formation. A, time-lapse recording of a fluorescently labeled Myo10-deficient (Myo10tm2/tm2) macrophage showing the appearance of retraction fibers following the retraction of a lamellipodial membrane protrusion. Scale bar, 10 μm. B, formation of bona fide filopodia (indicated by white arrows) by a Myo10-deficient macrophage challenged with E. coli particles (for clarity, only the green (membrane) channel is shown). The inset shows the overlay of green (membrane) and red (E. coli particles) channels. C, filopodia formation by a wild-type (WT) macrophage. Scale bar, 10 μm. D, summary box plot showing the number of nascent filopodia formed by WT (n = 34) versus Myo10 knock-out (n = 64) macrophages over a 20-min recording period. *, p < 0.05; Mann-Whitney U test (n = 2 independent experiments).
phages, although mDia1, mDia2 (42), and Myo10 (Fig. 11) are expressed in macrophages. Remarkably, macrophages from myeloid-restricted Cdc42 knock-out mice had an aberrant rounded-up morphological phenotype, which was not rescued by activation with a chemotactrant (complement C5a) or the TLR4 ligand LPS (lipopolysaccharide). This overt phenotype suggests that, at least in the case of macrophages, the Cdc42-WASP pathway, rather than the Rac-WAVE (WASP family verprolin homologs) pathway, plays a predominant role in stimulating Arp2/3 activity and lamellipodial spreading. Less surprisingly, we found that chemotactic navigation and cell velocity were decreased in Cdc42 conditional knock-out macrophages, compared with WT cells, in line with observations in neutrophils (24) and dendritic cells (25), as well as earlier work using dominant-negative constructs (26). Indeed, Yang et al. (43) recently showed in ele-

Figure 13. Paucity of nascent filopodia in Myo10 knock-out macrophages. A, time-lapse images of WT macrophages, prestimulated with LPS for 45 min, showing the formation of new finger-like membrane protrusions (filopodia). Scale bar, 10 μm. B, time-lapse 10 × 30-μm images cropped from the cell shown in panel A. Image intensity has been increased to better visualize (thin) filopodia, indicated by white arrows. C, image of four Myo10-deficient (Myo10tm2/tm2) macrophages (scale bar, 10 μm) stimulated with LPS. Enlarged views of one of the cells is shown at two different time points. Note the formation of filopodia (white arrows) and retraction fibers (the right membrane perimeter (marked red) at t = 0 s has been superimposed on the same cell at t = 470 s). D and E, Cdc42 cKO macrophages (scale bar, 10 μm) and 15 × 25-μm cropped view (scale bar, 10 μm) at different time points. White arrows indicate filopodia. F, summary data showing number of new filopodia over a 10-min recording period formed by macrophages without LPS (−LPS) or with LPS (+LPS) pretreatment. *, p < 0.05; Kruskal-Wallis one-way analysis of variance on ranks (at the 0.05 level of significance) (n = 30–42 cells per group; 2 independent experiments).
Filopodia and phagocytosis

Figure 14. Schematic summary. A, zymosan phagocytosis in mouse macrophages is driven by Dectin-1 receptors. When a zymosan particle is engaged by a filopodium, Dectin-1 presumably signals via its immunoreceptor tyrosine-based activation motif (ITAM)-like domain to induce outward actin flow and phagocytic cup formation. Following ingestion, nascent filopodia grow upon the closed phagocytic cup. B, the TLR4 (Toll-like receptor 4) ligand LPS (lipopolysaccharide), derived from the cell wall of Gram-negative bacteria such as *E. coli*, induces cell spreading and filopodia formation within 60 min of application. Spreading is markedly impaired in Cdc42-deficient macrophages, whereas Myo10-deficient macrophages have normal spreading, but reduced filopodia formation. Myo10 harbors pleckstrin homology domains, which promote binding to PIP2 (phosphatidylinositol 3,4,5-trisphosphate)-rich membranes. PIP2 production is catalyzed by PI3K (phosphoinositide 3-kinase), activated by various pathways, including TLR4 signaling. Time-lapse imaging reveals multiple modes in which filopodia support particle capture and ingestion, including retraction of the tip-bound particle (red spot) toward the cell body, surfing of an attached particle along the shaft of a filopodium (presumably driven by actin retrograde flow), combinations of surfing and retraction, and sweeping actions. C, summary table. Both Myo10 and Cdc42 are redundant for phagocytic cup formation, whereas Cdc42, but not Myo10, is important for chemotactic navigation.

Experimental procedures

Mice

Lifeact-EGFP mice were generated as previously described by Riedl et al. (8). To generate myeloid-restricted Cdc42 knock-out mice, Cdc42 (Cdc42fl/fl) mice, described by Wu et al. (45) and kindly supplied by Cord Brakebusch, were crossed with LysM-Cre mice (referred to as Lyz2tm1(cre)Ifo mice by the Mouse Genome Informatics database), kindly supplied by Irmgard Förster (Heinrich-Heine-Universität, Düsseldorf). Heterozygous Myo10 reporter knock-out (Myo10^{+/m2}) mice were obtained from the Wellcome Trust Sanger Institute (Hinxton, Cambridge, United Kingdom). Phenotypic analysis of these mice is documented in Bachg et al. (46), as Carsten...
Filopodia and phagocytosis

Kirschning (Universität Duisburg-Essen), and mannose receptor knock-out (MR−/−) mice were provided by Sven Burgdorf (LIMES (Life & Medical Sciences) Institut, Bonn). Dectin-1-deficient (Dectin-1−/−) mice were kindly provided by Gordon Brown (University of Aberdeen) and Laura E. Layland (Universitätsklinikum Bonn). Mice were bred in specific pathogen-free conditions and used in accordance with local animal experimentation guidelines. All procedures were approved by the University of Münster animal care and use committee.

Genotyping

PCR for genotyping the Cdc42 conditional knock-out mouse line was performed in two phases. First, touchdown PCR was performed using the following thermocycling protocol: 94 °C for 7 min, then 10 cycles of 94 °C for 30 s, 65 to 55 °C (starting at 65 °C and subtracting 1 °C each cycle) for 30 s, and 72 °C for 45 s. This was followed by 35 cycles of 94 °C for 30 s, 55 °C for 30 s, and 72 °C for 45 s. The following primer sequences (5' → 3') were used: Cdc42 forward, TCTGCCATCTACATA-CAC; Cdc42 reverse, ATGTAGTGCTGTCCATGG. The product sizes were 160 bp (wild-type allele) and 300 bp (floxed Cdc42 allele). The LysM-Cre allele was detected using the following primer pair (product size, 470 bp): forward (Cre-F), CCAATTATTGACCGTACC; reverse (Cre-R), TATAT-CCTGGCAGGATGTC. PCR genotyping for the Lifeact-EGFP mouse line was performed using a hot start (94 °C) and the following thermocycling protocol: 35 cycles of 94 °C for 30 s, 70 °C for 30 s, and 72 °C for 60 s. The following primer sequences (5' → 3') were used for the Lifeact-EGFP allele (product size, 265 bp): EGFP forward, GCACGACTTCTT-CGAAGTCCGATG; EGFP reverse, GCGGATCTGAGA-GTTACCTGTGAGCC. As a positive control, a primer pair giving a 466-bp product was used: Fabp2 (fatty acid-binding protein 2, intestinal) forward, CCTCCGGAGACGAGCCA-TTAAGGTGTCA; Fabp2 reverse, TAGAGCTTTTGCCA-CATCACAGGTC. Antibodies

Resident peritoneal macrophages were purified by cell sorting using rat anti-mouse F4/80 antibodies conjugated to Alexa Fluor 488 (5:200 dilution; MCA497A488, AbD Serotec, Germany, or MF48020, Molecular Probes, Life Technologies). The following antibodies were used for Western blot analyses: mouse anti-CD64 (clone 264-30, BD Pharmingen), rabbit anti-RhoQ (also known as anti-TC10; 1:1000 dilution; catalogue number, PA1-1061, Cell Signaling Technology), rabbit anti-Cdc42 (mouse; 1:1000 dilution; catalogue number 11A11, Cell Signaling Technology), rabbit anti-RhoJ (used at 1:500 dilution; catalogue number, ab57584, Abcam), rabbit anti-RhoF (used at 1:500 dilution; catalogue number, ab81024, Abcam) and goat anti-RhoD (1:500 dilution; catalogue number, sc-27880, Santa Cruz Biotechnology). Mouse monoclonal anti-β-actin antibodies (clone AC-15; catalogue number, A1978, Sigma-Aldrich, Germany), diluted 1:2000, were used to control loading in Western blot analyses. The following secondary antibodies were used: horseradish peroxidase-conjugated goat anti-mouse IgG polyclonal antibodies (115-035-003; Jackson ImmunoResearch Laboratories), horseradish peroxidase-conjugated goat anti-rabbit IgG polyclonal antibodies (111-035-003; Jackson ImmunoResearch Laboratories), and horseradish peroxidase-conjugated rabbit anti-goat IgG polyclonal antibodies (sc-2922; Santa Cruz Biotechnology), and horseradish peroxidase-conjugated rabbit anti-goat IgG polyclonal antibodies (sc-2922; Santa Cruz Biotechnology). Plasmids and cloning

To generate positive controls for Western blot analyses, the coding sequences of "mouse" RhoJ, RhoQ, and RhoF were subcloned into the expression vector pIRE2-DsRed2 (Clontech), which contains an internal ribosomal entry site (IRES) and an expression reporter (DsRed2), which encodes the red fluorescent protein DsRed2. The parent plasmids containing the genes of interest were pBluescript-Rho (insert accession number, AK003482; Source BioScience), pCMV6-RhoQ (insert accession number, BC056363; OriGene Technologies), pEGFP-RhoD (plasmid 23236; Addgene), and pEGFP-RhoF (gift from Harry Mellor). Successful insertion of the respective mouse Rho GT-Pase genes into the vector pIRE2-DsRed2 (yielding pIRE-RhoJ, pIRE-RhoQ, pIRE-RhoD, and pIRE-RhoF) was confirmed by DNA sequencing.

Isolation and handling of resident peritoneal macrophages

Mice were killed by an overdose of isoflurane in air, and the peritoneal cavity was lavaged via a 24-gauge plastic catheter (B. Braun, Melsungen, Germany) using 2 × 4.5 ml of ice-cold Hank's balanced salt solution without Ca2+ or Mg2+ (Thermo Fisher Scientific). After centrifugation (300 × g for 6.5 min), cells were resuspended in bicarbonate-free RPMI 1640 medium containing 20 μg/ml Heps (Biochrom AG, Berlin, Germany), and supplemented with 10% heat-inactivated fetal calf serum (FCS), 100 units/ml of penicillin, and 100 μg/ml of streptomycin (pH 7.4). The cells were seeded into fibronectin-coated μ-slide I chambers or μ-slide chemotaxis chambers (Ibidi, Martinried, Germany), and placed in a humidified incubator (37 °C). After 2 h, μ-slide I chambers were filled with 1 ml of RPMI 1640 medium, containing 10% FCS and antibiotics, and incubated overnight at 37 °C with 5% CO2. Experiments were performed after switching back to bicarbonate-free RPMI 1640 medium containing 20 μg/ml Heps. Alternatively, cells were seeded onto fibronectin-coated glass coverslips or glass bottom dishes.

Cell sorting

Mouse peritoneal macrophages were labeled with Alexa Fluor 488-conjugated anti-F4/80 antibodies, washed, and resuspended in autoMACS Running Buffer (130-091-221; Miltenyi Biotec), containing phosphate-buffered saline, 2 mM EDTA, and 0.5% bovine serum albumin (pH 7.2). Cell sorting and cell analyses were performed using a BD FACSAria II (or FACSARia III) cell sorter (BD Biosciences, San Jose, CA).

Western blot analyses

Macrophages and NIH-3T3 cells, transfected with Rho GT-Pase constructs, were lysed using Cell Lysis Buffer (9803; Cell Signaling Technology) containing: 20 mM Tris-HCl (pH 7.5), 150 mM NaCl, 1 mM Na2EDTA, 1 mM EGTA, 1% Triton X-100, 2.5 mM sodium pyrophosphate, 1 mM β-glycerophosphate, 1 mM Na3VO4 (phosphatase inhibitor), and 1 μg/ml of leupeptin (protease inhibitor), supple-
Labeled with Alexa Fluor 488-conjugated anti-F4/80 antibodies (Viano, Italy). Z-stacks of Lifeact-EGFP macrophages or cells were acquired for 10 min with 0.1% Tween 20 followed by overnight incubation (4 °C) with primary antibodies. For detection, horseradish peroxidase-conjugated secondary antibodies were used in combination with SuperSignal West Pico chemiluminescence substrate (Perbio, Bonn, Germany).

Two-dimensional chemotaxis assays

Cells obtained by peritoneal lavage of a single mouse were resuspended in 125–175 μl of medium, and 10 μl of the suspension was seeded into the narrow (1000 × 2000 × 70 μm) channel of an uncoated (IbidiTreat) μ-slide chemotaxis chamber (Ibidi). The narrow channel (observation area) connects two 40-μl reservoirs. After 3 h, the chemotaxis chamber was filled with bicarbonate-free RPMI 1640 medium containing 10% FCS, 100 ng/ml of lipopolysaccharide and antibiotics. Next, 15 μl of medium containing chemoattractant (C5a) and 0.003% Patent Blue V (blue dye) was drawn into one of the reservoirs. The final concentration of C5a was 20 nM. The observation area was imaged by phase-contrast microscopy via a ×10 (NA 0.3) objective lens. The blue dye served as a visual indicator of gradient formation, and we have previously confirmed that it does not affect cell migration. Images were captured every 2 min for 14 h, and cell migration tracks between 6 and 12 h were analyzed with ImageJ software (National Institutes of Health) using a manual tracking plug-in and the chemotaxis and migration tool from Ibidi. Twenty-five randomly selected cells were manually tracked in each chemotaxis experiment.

Spinning disk confocal microscopy

Two- or three-dimensional time-lapse imaging of living macrophages was performed using an UltraVIEW Vox three-dimensional live cell imaging system (PerkinElmer Life Sciences, Rodgau, Germany) coupled to a Nikon Eclipse Ti inverted microscope. The system incorporated a Yokogawa (Japan) CSU-X1 spinning disk scanner, a Hamamatsu (Japan) C9100-50 EM-CCD camera (1000 × 1000 pixels), and Volocity software. Cells were imaged via a Nikon ×60 (NA 1.49) oil immersion objective lens, and the temperature was maintained at 37 °C using an Okolab all-in-one stage incubator (Okolab, Otta- viano, Italy). Z-stacks of Lifeact-EGFP macrophages or cells labeled with Alexa Fluor 488-conjugated anti-F4/80 antibodies were obtained using steps in the range 0.5 to 1 μm. Focal drift was circumvented using the Nikon Perfect Focus System, which tracks the position of the coverslip in the z axis by reflecting infrared light (870 nm) from the coverslip and detecting it via a CCD (charge-coupled device) sensor.

DIC and fluorescence imaging

In selected experiments, time-lapse DIC and fluorescence microscopy was performed using a Zeiss inverted microscope (AxioObserver) equipped with a temperature-controlled incubator (incubator XL S, Zeiss) and controlled by AxioVision software (Zeiss). Images were captured via a ×63 (NA 1.4) oil-immersion objective lens and charge-coupled device camera (AxioCam MRm, Zeiss).

Phagocytosis assays

Time-lapse imaging of phagocytic events was performed after introducing particles to macrophages seeded in fibronectin-coated μ-slide I chambers, which have a channel volume of 100 μl. Stock solutions of BioParticles (Thermo Fisher Scientific) were prepared at 20 mg/ml in Dulbecco’s PBS and diluted 1:75 in live cell imaging medium, which consisted of bicarbonate-free RPMI 1640 medium containing 20 mM Hepes and 1 mM N-(2-mercaptoethyl)glycine, a free radical scavenger. The suspension of BioParticles in live cell imaging medium was sonicated to disperse the particles, and 100 μl was pipetted into a μ-slide I chamber before starting time-lapse imaging via spinning disk confocal microscopy. The following BioParticles were used: Alexa Fluor 594-conjugated E. coli (K-12 strain) particles (E23370; Thermo Fisher Scientific), Alexa Fluor 594-conjugated zymosan (polysaccharide prepared from the cell wall of S. cerevisiae (bakers’ yeast), catalogue number Z23374; Thermo Fisher Scientific) and pHrodo-conjugated zymosan, prepared using unlabeled zymosan (Z2849; Thermo Fisher Scientific) and the succinimidyl ester of pHrodo red (P36600; Thermo Fisher Scientific), which has absorbance and emission maxima of 560 and 585 nm, respectively. In addition, phagocytosis assays were performed using mouse IgG-coated polystyrene particles (MsGPX-50-5; Spherotech), which had a mean diameter of 5.19 μm. The 0.5% (w/v) stock solution was diluted 1:10 before use, and the beads were rendered red fluorescent by incubation with Alexa Fluor 594-conjugated goat anti-mouse IgG antibodies (ab150116; Abcam), followed by a wash step.

Scanning electron microscopy

Macrophages, seeded on fibronectin-coated glass coverslips, were fixed for 40 min with ice-cold Hepes-buffered (pH 7) medium containing 2% paraformaldehyde and 2.5% glutaraldehyde. As a pre-critical drying step, the cells were dehydrated by exposure, each time 15 min, to increasing concentrations of ethanol (30, 50, 70, 90, and 100%). The transitional fluid ethanol (31 °C and 73.8 bar). Critical point drying was achieved by subsequently decreasing the pressure at constant temperature (31 °C). The dried sample was sputter coated with platinum and imaged using an Hitachi S5000 scanning electron microscope.

Superresolution structured illumination microscopy

Superresolution-structured illumination microscopy was performed using an Elyra S.1 inverted microscope system (Carl Zeiss MicroImaging, Germany), controlled by ZEN 2011 SP2 software (black edition; Zeiss). Cells, seeded on mouse fibronectin-coated coverslips or glass bottom dishes, were imaged via an oil-immersion Plan Apo ×63 (NA 1.4) objective lens and images were captured with an Andor iXon EM-CCD. The following lasers and filters (in parentheses) were used: 488 nm (NIR), 532 nm, 561 nm, 633 nm, and 647 nm. The following lasers and filters (in parentheses) were used: 488 nm (NIR), 532 nm, 561 nm, 633 nm, and 647 nm.
Filopodia and phagocytosis

nm (BP 495–550 nm + LP 750 nm) and 561 nm (BP 570–620 nm + LP 750 nm). Five grating positions and 5 phase shifts were used for each z-slice.

Statistics

Normality and homoscedasticity were tested using the Shapiro-Wilk and Levene tests, respectively. A one-way analysis of variance was used to test for statistical differences at the 0.05 level of significance. When the assumed conditions of normality and homogeneity of variance were not fulfilled, as in most cases, we used the non-parametric Mann-Whitney U test or Kruskal-Wallis one-way analysis of variance on ranks (at the 0.05 level of significance). Post hoc multiple comparisons were made using Dunn’s method. Statistical analyses were performed using SigmaPlot (version 12) software (Systat Software), SPSS Statistics 22 (IBM Corporation), or Origin 2015 SR2 (OriginLab), and data are presented as box plots or mean ± S.E.


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References

Multiple roles of filopodial dynamics in particle capture and phagocytosis and phenotypes of Cdc42 and Myo10 deletion
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