Differential ability of proinflammatory and anti-inflammatory macrophages to perform macropinocytosis

Dar’ya S. Redka, Michael Gütschow, Sergio Grinstein, and Johnathan Canton

*Division of Cell Biology, Hospital for Sick Children, Toronto, ON M5G 1X8, Canada; †Pharmaceutical Institute, Pharmaceutical Chemistry I, University of Bonn, D-53121 Bonn, Germany; ‡Department of Biochemistry, University of Toronto, ON M5S 1A8, Canada; §Keenan Research Centre for Biomedical Science, St. Michael’s Hospital, Toronto, ON M5B 1W8, Canada

**Address correspondence to: Sergio Grinstein (sergio.grinstein@sickkids.ca).

ABSTRACT Macropinocytosis mediates the uptake of antigens and of nutrients that dictate the regulation of cell growth by mechanistic target of rapamycin complex 1 (mTORC1). Because these functions differ in proinflammatory and anti-inflammatory macrophages, we compared the macropinocytic ability of two extreme polarization states. We found that anti-inflammatory macrophages perform vigorous macropinocytosis constitutively, while proinflammatory cells are virtually inactive. The total cellular content of Rho-family GTPases was higher in anti-inflammatory cells, but this disparity failed to account for the differential macropinocytic activity. Instead, reduced activity of Rac/RhoG was responsible for the deficient macropinocytosis of proinflammatory macrophages, as suggested by the stimulatory effects of heterologously expressed guanine nucleotide-exchange factors or of constitutively active (but not wild-type) forms of these GTPases. Similarly, differences in the activation state of phosphatidylinositol 3-kinase (PtdIns3K) correlated with the macropinocytic activity of pro- and anti-inflammatory macrophages. Differences in PtdIns3K and Rho-GTPase activity were attributable to the activity of calcium-sensing receptors (CaSRs), which appear to be functional only in anti-inflammatory cells. However, agonists of PtdIns3K, including cytokines, chemokines, and LPS, induced macropinocytosis in proinflammatory cells. Our findings revealed a striking difference in the macropinocytic ability of pro- and anti-inflammatory macrophages that correlates with their antigen-presenting and metabolic activity.

INTRODUCTION

Macrophages can exist in a variety of functional states, often distinguished by their proinflammatory or anti-inflammatory properties.
localized primarily in endolysosomal compartments (Barton and Kagan, 2009; De Nardo, 2015). Macrophages can internalize pathogens, thereby delivering ligands to PRRs in intracellular compartments of PRRs, via two pathways: phagocytosis, which mediates the uptake of particulate targets, and macropinocytosis, a distinct process that enables the internalization of soluble signals. By trapping soluble PRR ligands released by pathogens, macropinocytosis enables early signaling by macrophages distant from the sites of infection.

Macropinocytosis, the uptake of extracellular fluid into large (>250 nm) vacuoles called macropinosomes, is an actin- and phosphatidylinositol 3-kinase (PI3K)-dependent process (Swanson, 2008; Kerr and Teasdale, 2009). In certain cell types, macropinocytosis can be triggered by growth signals or chemokines for nutrient uptake during development and cancer (Bloomfield and Kay, 2016), recycling of adhesion receptors during cell migration (Gu et al., 2011), or for membrane retrieval during neuronal growth (Holt et al., 2003). Such macropinocytosis will be referred to hereafter as the induced type. In dendritic cells and macrophages, however, where macropinocytosis is thought to play a role in PRR ligand uptake and antigen presentation (Swanson, 2008; Lim and Gleeson, 2011), macropinocytosis occurs continuously in the ostensible absence of stimulation by growth factors or chemokines. In nonpolarized human monocyte-derived macrophages (hMDMs), this type of “constitutive” macropinocytosis was shown to be mediated by calcium-sensing receptors (CaSRs; Canton et al., 2016).

Considering the different functional roles of proinflammatory and anti-inflammatory macrophages, we wondered whether macropinocytosis operates similarly in differentially polarized macrophages. To this end, we compared the constitutive macropinocytic activity of hMDMs polarized toward the proinflammatory (M1-like) phenotype using granulocyte macrophage colony-stimulating factor (GM-CSF) followed by interferon-γ (IFN-γ) and lipopolysaccharide (LPS), with that of anti-inflammatory (M2-like) cells generated using macrophage colony-stimulating factor (M-CSF) followed by interleukin-4 (IL-4). We also assessed whether macropinocytosis could be induced acutely in these cells by the addition of cytokines, chemokines, or LPS. Our analysis revealed striking differences in the macropinocytic activity of both polarization states, prompting us to investigate the molecular basis of this contrasting behavior.

RESULTS

Analysis of constitutive macropinocytosis in polarized macrophages

Dendritic cells and macrophages are known to carry out macropinocytosis constitutively, i.e., in the absence of ostensible stimulation (Sallusto et al., 1995; Lim and Gleeson, 2011; Canton et al., 2016). The effect of polarization on macropinocytic activity, however, has not been established. To this end, proinflammatory macrophages were generated by exposing human blood monocytes to GM-CSF for 5 d, followed by treatment with IFN-γ and LPS for an additional 2 d. These cells are referred to hereafter as GM/IFN-γ/LPS–cultured cells. Anti-inflammatory macrophages were induced by exposing monocytes to M-CSF, followed by IL-4 and are referred to as M/IL4–cultured cells. The occurrence of polarization was validated assessing the expression of two canonical proinflammatory cell-surface markers, CD40 and CD80, and two anti-inflammatory markers, CD163 and CD200R (Jagun et al., 2013; Vogel et al., 2014) by flow cytometry. On day 7, GM/IFN-γ/LPS–cultured cells had high levels of CD40 and CD80, whereas expression of these markers was low in M/IL4–cultured cells (Figure 1, Supplemental Figure S1, and Supplemental Table S1). Conversely, the anti-inflammatory markers CD163 and CD200R were clearly expressed in M/IL4–cultured cells yet were barely detectable in GM/IFN-γ/LPS–cultured cells. Measuring CD1a expression, we had confirmed previously that these protocols yield preparations devoid of dendritic cells (Canton et al., 2014).

The pro- and anti-inflammatory cells generated as above differed also in their appearance. In accordance with earlier findings (McWhorter et al., 2013), the GM/IFN-γ/LPS–cultured cells displayed a round and flattened (“fried-egg”) morphology, while M/IL4–cultured cells were elongated (Figure 2A).

To examine the ability of GM/IFN-γ/LPS– and M/IL4–cultured macrophages to perform macropinocytosis constitutively, we measured the uptake of tetramethylrhodamine (TMR)-labeled 70 kDa dextran in otherwise unstimulated cells. As shown in Figure 2B, M/IL4–cultured macrophages took up large amounts of the 70 kDa dextran, generating on average 46 macropinosomes (defined as vacuolar structures of diameter ≥0.8 µm) per cell in 15 min. In sharp contrast, macropinosomes were rarely found in the GM/IFN-γ/LPS–cultured macrophages; after a comparable incubation less than one macropinosome per cell was observed (Figure 2, B and C).

FIGURE 1: Cell-surface expression of pro- and anti-inflammatory markers in macrophages cultured under various conditions. Human blood monocytes were cultured for 5 d as described in Materials and Methods in either M-CSF (columns 1–4) or GM-CSF (columns 5–7), followed by 2 more days in either M-CSF (M) or GM-CSF (GM) alone, or in the additional presence of either IL-4 (M/IL4), IFN-γ (M/IFN-γ and GM/IFN-γ), or a combination of IFN-γ and LPS (M/IFN-γ/LPS and GM/IFN-γ/LPS). Cells were then lifted and labeled with either FITC-conjugated anti-human CD40 (a proinflammatory marker; A), APC-conjugated anti-human CD80 (a proinflammatory marker; B), Alexa-647-conjugated anti-human CD163 (an anti-inflammatory marker; C), or PE-conjugated anti-human CD200R antibody (an anti-inflammatory marker; D). Cell-surface labeling was determined by flow cytometry. Similarly labeled isotype-matched (κ) controls were used to subtract nonspecific labeling (shown in Supplemental Figure S1). Data are means (± SEM) of median fluorescence intensity per cell, minus the signal from the corresponding isotype control, normalized to the highest median fluorescence intensity per experiment, from 15,400 to 19,700 cells in three independent experiments of each type, with blood from three donors. Representative fluorescence intensity histograms are shown in Supplemental Figure S1. Probabilities of statistical significance for comparisons between all pairs of data sets are presented as a matrix in Supplemental Table S1.
It was shown previously that 70 kDa dextran is internalized almost exclusively via macropinosomes, as it is excluded by smaller-sized endocytic pathways such as clathrin-coated vesicles or caveolae (Li et al., 2015). This was confirmed to be the case also for nonpolarized human hMDMs, where uptake of the large dextran was virtually eliminated by latrunculin—the prototypical inhibitor of macropinocytosis—under conditions where the uptake of transferrin (a marker of clathrin-mediated uptake) was unaffected (Canton et al., 2016). To ensure that the probe used was entering the cells through fluid-phase, as opposed to adsorptive uptake, M/IL4-cultured cells were incubated with the labeled dextran for 15 min at 4°C, followed by its removal from the medium and incubation for an additional 15 min at 37°C. Under these conditions, uptake of dextran was negligible (Supplemental Figure S2, A and B). In addition, after coincubation at 37°C for 15 min, the small fluorescent molecule Lucifer Yellow, which is structurally different from dextran and hence unlikely to adsorb onto the same sites, filled the same large vacuoles as the 70 kDa dextran (Supplemental Figure S2, C and D). Together, these observations confirm that 70 kDa TMR-dextran is internalized via fluid-phase uptake, rather than via an adsorptive mechanism.

Both induced and constitutive forms of macrophage polarization were measured by ptdIns(3,4,5)P3 detection. To confirm that the vacuoles formed constitutively by the M/IL4-cultured cells were bona fide macropinosomes, we tested the effects of wortmannin (a PtdIns3K inhibitor) and latrunculin B. Both of these agents markedly reduced the number of macropinosomes in M/IL4-cultured macrophages (Figure 2D). Interestingly, while amiloride and its analogues are potent inhibitors of macrophage polarization in several systems (West, 1989; Dowrick et al., 1993; Canton et al., 2016), HOE-694 was without effect on M/IL4-cultured cells (Figure 2C), replicating the results obtained in other myeloid cells (Norbury et al., 1997; Canton et al., 2016).

**Macropinocytic activity of macrophages with intermediate phenotypes**

Differentiation of macrophages with GM/IFN-γ/LPS or M/IL4 gives rise to two extreme phenotypes: fully pro- and anti-inflammatory, respectively. Others have used less extreme culture conditions to polarize macrophages, some of which yielded intermediate phenotypes. We therefore assessed the macrophagic ability of cells subjected to a range of differentiation protocols (detailed in Materials and Methods), namely, 1) M-CSF alone (M); 2) M-CSF followed by IFN-γ (M/IFN-γ); 3) M-CSF followed by IFN-γ and LPS (M/IFN-γ/LPS); 4) GM-CSF only (GM); and 5) GM-CSF followed by IFN-γ (GM/IFN-γ). To verify the effects of these individual protocols on polarization we measured the expression of proinflammatory markers (CD40 and CD80; Figure 1, A and B) and anti-inflammatory markers (CD80 and CD200R; Figure 1, C and D; Jaguin et al., 2013; Vogel et al., 2014) on day 7 of differentiation. The expression of CD40 was comparatively low with M-CSF alone, but increased 4.0-fold and 7.4-fold if IFN-γ or a mixture of IFN-γ and LPS, respectively, were also present (Figure 1A; see Supplemental Table S1 for probabilities of statistical significance for pairwise comparisons). A similar pattern was observed with GM-CSF–cultured cells (Figure 1A). The expression of CD80 was elevated primarily in GM/IFN-γ/LPS– (left) and M/IL4– (right) macrophages. (A) GM/IFN-γ/LPS– (left) and M/IL4– (right) macrophages were pretreated with HOE 694 (10 µM, 30 min), latrunculin B (2 µM, 30 min), or wortmannin (100 nM, 15 min) and then incubated with fluorescently labeled 70 kDa dextran (TMR-dextran, 125 µg/ml) for 15 min at 37°C in calcium-containing medium. Cells were then washed, fixed, and imaged immediately by spinning disk confocal microscopy. The number of macropinosomes (i.e., TMR-positive vacuoles) per cell then was determined as described for C. Typical images (A, B) and quantifications (C, D; means ± SEM) are representative of 30–60 cells from three to six experiments of each type using blood from at least two different donors. Scale bars, 15 µm.

**FIGURE 2:** Differential macropinocytic activity and actin structure of GM/IFN-γ/LPS– vs. M/IL4–cultured macrophages. (A) GM/IFN-γ/LPS– (left) and M/IL4– (right) macrophages were fixed, permeabilized, and labeled with fluorescent phalloidin. (B) GM/IFN-γ/LPS– (left) and M/IL4– (right) macrophages were incubated with fluorescently labeled 70 kDa dextran (TMR-dextran, 125 µg/ml) for 15 min at 37°C in calcium-containing medium. Cells were then washed, fixed, and imaged immediately by spinning disk confocal microscopy. The number of macropinosomes (i.e., TMR-positive vacuoles) per cell then was determined as described for C. Typical images (A, B) and quantifications (C, D; means ± SEM) are representative of 30–60 cells from three to six experiments of each type using blood from at least two different donors. Scale bars, 15 µm.

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proinflammatory properties, while those cultured in M-CSF with or without IL-4 acquired (at least some) anti-inflammatory properties, which were in turn suppressed by GM-CSF. As anticipated, the combination of GM/IFN-γ/LPS yielded the most pronounced proinflammatory property, while M/IL4-cultured macrophages displayed the most consistent anti-inflammatory phenotype.

Next, we examined the macropinocytic activity of these differently polarized macrophages. As illustrated in Figure 3, both M- and M/IL4-cultured macrophages exhibited a considerably higher macropinocytic activity than the other preparations (e.g., 15- and 7-fold higher, respectively, than GM-cultured cells; see Table S2 for probabilities of statistical significance for pairwise comparisons). GM-CSF was insufficient to elicit macropinocytosis, while IFN-γ and/or LPS suppressed the stimulatory effect of M-CSF. It appears, therefore, that the macropinocytic phenotype is associated with an anti-inflammatory phenotype. To further analyze the mechanism underlying the differential macropinocytic activity, we therefore proceeded to use the two most contrasting phenotypes: GM/IFN-γ/LPS– versus M/IL4-cultured cells.

**Rho GTPases are more abundant and more active in M/IL4- than in GM/IFN-γ/LPS–cultured macrophages**

Morphological differences between the GM/IFN-γ/LPS– and M/IL4-cultured macrophages pointed to a distinct cytoskeletal arrangement, which was validated by staining F-actin with phalloidin (Figure 2A). Whereas GM/IFN-γ/LPS–cultured cells displayed a continuous band of cortical actin on their dorsal membrane and multiple ventral podosomes, M/IL4-cultured cells had florid dorsal ruffles and few podosomes. We reasoned that these profound differences may be attributable to differential content and/or activity of Rho-family GTPases, which dictate the assembly of F-actin. The abundance of Rac1, RhoA, RhoG, and Cdc42 was compared in the two polarization states by immunoblotting, normalizing the data to the actin content. Remarkably, M/IL4-cultured macrophages expressed higher levels of the four tested Rho GTPases, the difference being greatest for Rac1 (Figure 4A; see Supplemental Figure S3A for representative blots).

To determine whether the differential expression of the Rho GTPases resulted in their differential activity, the GTP-bound (i.e., active) form of each of the G proteins was measured using either a G-LISA (for Rac1, RhoA, and Cdc42) or a pull-down assay (for RhoG; Figure 4B and Supplemental Figure S3B). Rac1 and RhoG were significantly more active in M/IL4– than in GM/IFN-γ/LPS–cultured cells, while the differences detected in the case of RhoA and Cdc42 were not statistically significant (Figure 4B).

**Activation—but not overexpression—of Rac1 or RhoG confer constitutive macropinocytic activity to GM/IFN-γ/LPS–cultured macrophages**

Considering that M/IL4-cultured macrophages express substantially greater levels of Rac1 and to a lesser extent RhoA, RhoG, and Cdc42 (Figure 4A), we tested whether ectopic overexpression of any of the GTPases would confer the constitutive macropinocytotic ability to GM/IFN-γ/LPS–cultured cells. Fluorescently tagged versions of the wild-type forms of the GTPases were used to verify their expression following transient transfection of GM/IFN-γ/LPS–cultured macrophages. As shown in Figure 4C, none of the GTPases increased macropinocytosis significantly in the GM/IFN-γ/LPS–cultured cells. The lower abundance of Rho GTPases in the GM/IFN-γ/LPS–cultured cells is therefore unlikely to account for their reduced macropinocytic effectiveness.

We next considered whether insufficient activation rather than differential expression of the GTPases was responsible for the reduced macropinocytosis of GM/IFN-γ/LPS–cultured cells. This possibility was assessed expressing constitutively active (CA) forms of the GTPases; as before, fluorescently tagged constructs were used to identify transfectants. Transfection of GM/IFN-γ/LPS–cultured macrophages with either CA-Rac1 (Rac1-Q61L-GFP [green fluorescent protein]) or CA-RhoG (RhoG-G12V-CFP) resulted in a striking increase in macropinocytosis, exceeding the constitutive levels observed in the M/IL4-cultured macrophages (Figure 4, D and E). CA-Cdc42 (Cdc42-G12V-YFP) effected a much smaller, yet significant, increase, whereas CA-RhoA (RhoA-Q63L-GFP) was without effect (Figure 4E).

Together, these results indicate that availability of active Rac1 and/or RhoG, as opposed to the total level of expression of the GTPases, limit the ability of GM/IFN-γ/LPS–cultured cells to perform constitutive macropinocytosis. This notion was validated by heterologous overexpression of guanine nucleotide exchange factors (GEFs) that can activate the endogenous GTPases. To activate Rac1 we used T-cell lymphoma invasion and metastasis-inducing protein 1 (Tiam1), which has been shown to localize to the membrane in macrophages (Bohdanowicz et al., 2013). SH3-containing GEF (sGEF) was used to stimulate RhoG; of note, sGEF was shown to be involved in macropinocytosis in epithelial cells (Ellerbroek et al., 2004). As illustrated in Figure 4, D and E, both GEFs caused a remarkable increase in the number of macropinosomes, reminiscent of the effects of the CA mutants of Rac1 and RhoG. These results suggest that, regardless of the lower expression levels detected in GM/IFN-γ/LPS–cultured cells, the amount of Rac1 and RhoG expressed by GM/IFN-γ/LPS–cultured macrophages suffices to support active macropinocytosis.
PtdIns(3,4,5)P3 is less active in the GM/IFN-γ/LPS–cultured macrophages

The preceding data indicate that the ineffective activation of Rac1 and RhoG curtails the ability of GM/IFN-γ/LPS–cultured cells to form macropinosomes, suggesting insufficient GEF activity. Several Rho-family GEFs are recruited and/or activated by PtdIns(3,4,5)P3, and this phosphoinositol is essential for macropinocytosis. We therefore considered the possibility that reduced PtdIns(3,4,5)P3 generation may be responsible for the poor ability of GM/IFN-γ/LPS–cultured cells to perform constitutive macropinocytosis. To that end, we used a fluorescently labeled PH domain of Akt [PH(Akt)-GFP] to compare the abundance and distribution of PtdIns(3,4,5)P3 in GM/IFN-γ/LPS– and M/IL4-cultured cells. In M/IL4-cultured macrophages, the biosensor associated with the inner leaflet of the plasma membrane, making its fluorescence intensity significantly greater than that of the cytosol (Figure 5, A and B). That this reflected the constitutive presence of PtdIns(3,4,5)P3 was indicated by the effect of wortmannin, which caused the release of the membrane-associated PH(Akt)-GFP, increasing the cytosolic fluorescence intensity. In contrast, no preferential association of the probe with the membrane was discernible in most GM/IFN-γ/LPS–cultured cells.

The activity of PtdIns3K was also compared in the two types of macrophages by quantifying the phosphorylation of Akt, which is known to vary in proportion to the PtdIns(3,4,5)P3 content. Immunoblotting revealed that Akt is phosphorylated to a greater extent in M/IL4- than in C. difficile toxin B treatment (3 h in serum-free medium) was used to inhibit all four GTPases, i.e., as a negative control, in all G-LISA and RhoG.GTP pull-down assays. (C–E) GM/IFN-γ/LPS–cultured macrophages transfected with fluorescently tagged constructs of either wild-type (C) or constitutively active Rac1, RhoA, RhoG, or Cdc42, or of the Rac1 and RhoG GEFs Tiam1 and sGEF, as indicated (D, E). The specific constructs used were Rac1-GFP, RhoA-GFP, RhoG-CFP, Cdc42-GFP, Rac1-Q61L-GFP, RhoA-Q63L-GFP, RhoG-G12V-CFP, Cdc42-G12V-YFP, Tiam1-GFP, and sGEF-GFP. GM/IFN-γ/LPS–cultured macrophages transfected with GFP alone were used as a negative control, while M/IL4-cultured macrophages transfected with GFP alone were used as a positive control. After 24-h transfection, the cells were incubated with fluorescently labeled 70 kDa dextran (TMR-dextran, 125 µg/ml) for 15 min at 37°C, and washed, fixed, and imaged immediately (D); only transfected cells were selected for measurements of macropinocytosis, which was quantified (C, E) as the total volume of TMR-positive vacuoles per cell from 3D stacks using 3D particle analysis in ImageJ software, applying a lower particle volume threshold of 0.26 µm3. Typical images (D) and quantifications (C, E; means ± SEM) are representative of 20–50 cells from three to five independent experiments using blood from at least two separate donors. Scale bars, 15 µm.
We next measured the relative abundance of isoforms of the class I PtdIns3K to establish whether the increased activity of M/IL4- compared with GM/IFN-γ/LPS–cultured cells is attributable to differential expression of the enzyme. We quantified the amount of catalytic (p110α, p110β, p110δ, and p110γ) and regulatory (p85) subunits of the kinase in GM/IFN-γ/LPS– and M/IL4-cultured macrophages by immunoblotting (Figure 5D and Supplemental Figure S4B). Expression of p110γ subunit was significantly (~2.0-fold) higher in the M/IL4-cultured cells, while only smaller, statistically insignificant differences were noted for the p110α, p110β, p110δ, and p85 subunits (Figure 5D).

Recruitment of the p85 regulatory subunit of PtdIns3K to the plasma membrane induces macropinocytosis in GM/IFN-γ/LPS–cultured macrophages

The preceding results suggest that the reduced activity of PtdIns3K in GM/IFN-γ/LPS–cultured cells and their failure to perform constitutive macropinocytosis may result from the lower expression of the p110γ catalytic subunit of PtdIns3K. However, insufficient activation of its regulatory subunit could equally explain the observations. To distinguish between these alternatives, we used a rapamycin-inducible heterodimerization system to steer the p85 subunit of PtdIns3K to the plasma membrane (Figure 6A). GM/IFN-γ/LPS– and M/IL4-cultured macrophages were cotransfected with two constructs: 1) a domain of FK506-binding protein (FKBP) fused to the p85 subunit and a cyan florescent protein (CFP), and 2) the FKBP-rapamycin binding (FRB) domain of mechanistic target of rapamycin (mTOR) fused to the N-terminal 11 amino acids of Lyn kinase, which target the fusion protein to the plasma membrane. Before the addition of rapamycin, FKBP-p85-CFP was mostly cytosolic, but was translocated to the plasma membrane within minutes of the addition of rapamycin (Figure 6B for GM/IFN-γ/LPS–cultured cells; M/IL4-cultured cells not shown). The effect of translocation of p85 to the plasma membrane on the activity of PtdIns3K was verified using the biosensor of PtdIns(3,4,5)P3 [PH(Akt)-RFP]. In the GM/IFN-γ/LPS–cultured macrophages translocation of p85 to the plasma membrane resulted in a 2.5-fold increase in the phosphamemal phospho-Akt (Akt) biosensor.

GM/IFN-γ/LPS–cultured macrophages (Figure 5C and Supplemental Figure S4A), in good accordance with the results obtained with the PtdIns(3,4,5)P3 biosensor.
GM/IFN-γ/LPS–cultured cells to perform macropinocytosis. As illustrated in Figure 6, D and E, stimulation of PtdIns(3,4,5)P\(_3\) synthesis by rapamycin-mediated recruitment of p85 markedly activated the macropinocytic activity of GM/IFN-γ/LPS–cultured cells. Stimulation was also observed in M/IL4–cultured cells, although the net effect was not statistically significant due to the varying magnitude of the response. Rapamycin has been reported to depress macropinocytosis, which was quantified as described for Figure 2C (E). Representative images (D) and quantifications (means ± SEM; E) of 30–50 cells from three to five independent experiments. Scale bars, 15 µm.

FIGURE 6: Recruitment of the p85 domain of PtdIns3K to the plasma membrane increases macropinocytic activity of GM/IFN-γ/LPS–cultured macrophages. (A) Schematic showing the recruitment of p85-CFP to the plasma membrane via the rapamycin-inducible heterodimerization system. (B, C) GM/IFN-γ/LPS– and M/IL4–cultured macrophages were cotransfected with FKBP-p85-CFP, Lyn\(_{11}\)-FRB, and PH(Akt)-RFP. Transfected cells were imaged (B) 24 h posttransfection either before (left) and after (right) addition of rapamycin (1 µM, 15 min; the same cell is shown on the left and right), and the activation PtdIns3K was quantified as redistribution of PH(Akt)-RFP, a biosensor of PtdIns(3,4,5)P\(_3\), to the plasma membrane (C), quantified as described for PH(Akt)-GFP in Figure 5B. Typical images (B) and quantifications (C; means ± SEM) are representative of 20–30 cells from three to four independent experiments. (D, E) GM/IFN-γ/LPS– and M/IL4–cultured macrophages were cotransfected with FKBP-p85-CFP and Lyn\(_{11}\)-FRB. After a 24-h transfection, the cells were incubated with fluorescently labeled 70 kDa dextran (TMR-dextran, 125 µg/ml) for 15 min at 37°C, with or without concurrent treatment with rapamycin (1 µM). Cells were then washed, fixed, and imaged immediately (D). Cells positive for CFP were selected for measurements of macropinocytosis, which was quantified as described for Figure 2C (E). Representative images (D) and quantification (means ± SEM; E) of 30–50 cells from three to five independent experiments. Scale bars, 15 µm.

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Together, the preceding results indicate that, despite their lower expression, sufficient amounts of the catalytic subunits of PtdIns3K are present in GM/IFN-γ/LPS–cultured cells to support macropinocytosis, and that inadequate activation underlies their inability to form macropinosomes.

The constitutive activity of PtdIns3K in M/IL4–cultured macrophages is attributable to CaSR

We recently showed that constitutive macropinocytosis in nonpolarized hMDMs is stimulated by the calcium-sensing receptor, CaSR (Canton et al., 2016). This serpentine receptor is potentially capable of activating the p110\(_\gamma\) and p110\(_\delta\) subunits of PtdIns3K via a heterotrimeric G protein, likely G\(_i\). It was therefore conceivable that the constitutive macropinocytosis observed in macrophages polarized toward the anti-inflammatory phenotype was similarly mediated by CaSR. This possibility was tested by a) removing extracellular calcium (the ligand responsible for activation of CaSR); b) addition of NPS 2143, a specific CaSR antagonist; c) pretreatment of the cells with pertussis toxin, an inhibitor of G\(_i\); and d) addition of BIM 46187, a pan-G protein inhibitor. As reported in Figure 7A, every one of these treatments markedly inhibited the constitutive macropinocytic activity of M/IL4–cultured macrophages, consistent with the involvement of CaSR and the G\(_i\) protein in the process.

Because macropinocytic activity in polarized macrophages was proportional to the level of PtdIns(3,4,5)P\(_3\), we examined whether the constitutive PtdIns3K activity is indeed dictated by CaSR. To this end, the plasmalemmal levels of the PtdIns(3,4,5)P\(_3\) biosensor were compared in GM/IFN-γ/LPS– versus M/IL4–cultured macrophages, in the presence and absence of extracellular calcium (Figure 7B), and in the presence of the CaSR antagonist NPS 2143 (Figure 7C). The removal of external calcium markedly reduced plasmalemmal PtdIns(3,4,5)P\(_3\) in the M/IL4–cultured cells, but had only an insignificant effect in the GM/IFN-γ/LPS–cultured cells. Comparable results were observed using NPS 2143, which had a pronounced effect on M/IL4–, but not GM/IFN-γ/LPS–cultured cells. The specificity of NPS 2143 was confirmed by the observation that the antagonist had no effect in HEK 293 cells (unpublished data), which lack CaSR (Canton et al., 2016).

The preceding results suggest that CaSR is more active in M/IL4– than in GM/IFN-γ/LPS–cultured macrophages. We sought to
validate this conclusion by measuring other outcomes of CaSR stimulation. CaSR has been shown to couple to both G_i and G_o proteins (Saidak et al., 2009; Conigrave and Ward, 2013); G_i protein regulates adenylyl cyclase and PtdIns3K, whereas G_o protein activates PLC. By generating diacylglycerol, activation of PLC indirectly stimulates the production of PA. We therefore examined whether the plasmalemmal levels of PA were different between GM/IFN-γ/LPS- and M/IL4-cultured cells, and whether any observed differences were attributable to CaSR, which was assessed by omission of calcium. A biosensor of PA, GFP-2PABD (Bohdanowicz et al., 2013), was expressed in both cell types and its levels at the plasma membrane were normalized to those in the cytoplasm. M/IL4-cultured macrophages had 2.4-fold more plasmalemmal GFP-2PABD than did the GM/IFN-γ/LPS-cultured cells (Figure 7D). Although the difference did not reach statistical significance, it was clear that PA accumulation in the M/IL4-cultured cells was depressed by inhibition of PLC with U73122 (Figure 7D) and also with DAG Kinase Inhibitor II (Figure 7E). Most importantly, accumulation of GFP-2PABD in the membrane of M/IL4-, but not GM/IFN-γ/LPS-cultured cells required extracellular calcium, consistent with formation of PA via CaSR-induced stimulation of G_o.

Next, we examined whether the differential function of the CaSR in the two types of macrophages was due to differential expression; the absence of CaSR in GM/IFN-γ/LPS-cultured cells could readily explain our functional observations. Although available antibodies were not suitable for detection of endogenous CaSR protein by immunoblotting or immunofluorescence, message encoding the receptor was readily detectable in GM/IFN-γ/LPS-cultured cells; its abundance, determined by PCR, was not significantly different from that in M/IL4-cultured cells (Figure 7F). In addition, ectopic expression of CaSR tagged with a fluorescent protein failed to increase constitutive macropinocytosis in the GM/IFN-γ/LPS-cultured macrophages was determined by RT-PCR. Expression of GAPDH mRNA was used as a reference. The ethidium bromide gel is representative of four independent experiments. When normalized to GAPDH, the average intensity of bands corresponding to CaSR were not significantly different between GM/IFN-γ/LPS- and M/IL4-cultured cells (p = 0.26). Abbreviation: PTX, pertussis toxin.
Acute treatment with cytokines and chemokines induces macropinocytosis in GM/IFN-γ/LPS–cultured cells

While macropinocytosis is negligible in otherwise untreated GM/IFN-γ/LPS–cultured macrophages, we showed earlier that macropinosomes can be formed by these cells when the p85 regulatory subunit of PtdIns3K is forcibly recruited to the plasma membrane (Figure 6). This raised the possibility that receptors other than CaSR may induce macropinocytosis in GM/IFN-γ/LPS–cultured cells. To analyze this possibility we assessed the effects of several agonists on macropinosome formation, comparing their effectiveness in GM/IFN-γ/LPS– and M/IL4-cultured macrophages. These experiments were performed in nominally calcium-free medium, to obviate the contribution of CaSR to macropinosome formation in the M/IL4–cultured cells. When added at 250 ng/ml (a concentration 10-fold higher than that used during differentiation) M-CSF induced significant macropinocytosis in both GM/IFN-γ/LPS– and M/IL4–cultured macrophages, the effects being greater in the latter (Figure 8). Similar effects were observed when a high dose (250 ng/ml) of GM-CSF was added acutely. We also tested the acute response to LPS (500 ng/ml); in this instance, LPS was omitted from the culture during incubation with GM-CSF plus IFN-γ. As shown in Figure 8, LPS markedly stimulated macropinocytosis in both cell types. Lastly, we compared the effectiveness of two chemokines, CCL7 and CCL19, which have been shown to promote chemotaxis differently in pro- and anti-inflammatory macrophages (Xuan et al., 2015). CCL7 induced appreciable macropinocytosis in both GM/IFN-γ/LPS– and M/IL4–cultured cells, although the effect was of borderline significance in the proinflammatory cells (Figure 8). CCL19, on the other hand, induced considerable macropinocytosis in the GM/IFN-γ/LPS–, but not the M/IL4–cultured cells (Figure 8), consistent with its effects on chemotaxis (Xuan et al., 2015). Taken together, these findings support the conclusion that GM/IFN-γ/LPS–cultured macrophages possess a functional and responsive macropinocytic machinery but that, unlike M/IL4–cultured cells, do not perform macropinocytosis in the “resting” state because their constitutive activity of PtdIns3K is minimal.

**DISCUSSION**

In the present report, we show that blood-derived anti-inflammatory M/IL4–cultured macrophages actively perform constitutive macropinocytosis, while their proinflammatory GM/IFN-γ/LPS–cultured counterparts lack this ability. Differential macropinocytic ability was also observed in macrophages that were cultured under less extreme conditions, but still displayed distinct anti- and proinflammatory properties (Figures 1 and 3). The uptake of large dextrans by the M/IL4–cultured cells was actin and PtdIns3K dependent (Figure 4D), hallmarks of bona fide macropinocytosis (Swanson, 2008; Kerr and Teasdale, 2009). While M/IL4–cultured cells displayed considerably higher expression of Rho-family GTPases, notably Rac1 and RhoG that are thought to support macropinocytosis, this difference is unlikely to account for the inability of GM/IFN-γ/LPS–cultured cells to internalize the dextrans. This conclusion stems from the observation that ectopic overexpression of the wild-type form of the GTPases in GM/IFN-γ/LPS–cultured cells failed to restore macropinocytic activity (Figure 4C). In contrast, the constitutively active forms of Rac1 and RhoG strongly enhanced the macropinocytic activity of the GM/IFN-γ/LPS–cultured cells, reaching levels exceeding those seen in untreated M/IL4–cultured cells (Figure 4E). Moreover, the endogenous levels of the GTPases sufficed to stimulate macropinocytosis when GEFs for Rac1 or RhoG (Tiam1 and sGEF, respectively) were overexpressed. We therefore concluded that insufficient activation—as opposed to limited expression—of the GTPases accounts for the inactivity of GM/IFN-γ/LPS–cultured macrophages. This implies that the machinery downstream from the active Rho GTPases that is necessary for macropinocytosis is available and poised to respond in GM/IFN-γ/LPS–cultured cells.

Our experiments also showed that insufficient activation of the GTPases in GM/IFN-γ/LPS–cultured macrophages correlates with reduced accumulation of PtdIns(3,4,5)P3 by these cells, which was established using genetically encoded biosensors and also by assessing the phosphorylation state of Akt by immunoblotting (Figure 5, A–C). As in the case of the GTPases, the abundance of PtdIns3 kinases was greater in M/IL4– than in GM/IFN-γ/LPS–cultured cells, but their degree of activation—rather than their density—appeared to be the cause of the reduced level of PtdIns(3,4,5)P3 in the inflammatory cells. Forced recruitment of the catalytic subunit (Figure 6) confirmed this notion.

The elevated activity of PtdIns3K we observed in M/IL4–cultured macrophages is consistent with previous reports, which show that increased PtdIns3K function is associated with polarization of...
macrophages toward the anti-inflammatory, immunosuppressive M2-like phenotype (Wang et al., 2015; Kaneda et al., 2016). Conversely, reduction of PtdIns3K activity by genetic or pharmacological means leads to increased expression of M1 markers and an exaggerated inflammatory response to infection (Kaneda et al., 2016).

Our data further suggest that CaSR activity contributes to the constitutive formation of PtdIns(3,4,5)P$_3$ in the M/IL4-cultured cells. Heterotrimeric G proteins such as those that transduce the signals generated by CaSR can accomplish this feat, by activating the p110$_{\beta}$ or p110$_{\gamma}$ isoforms of PtdIns3K. In this regard, it is noteworthy that macropinocytosis can be induced in other systems (e.g., podocytes) by addition of free fatty acids, which activate G protein–coupled receptors (Chung et al., 2015). Notably, activation of one such receptor, GPR120, by free fatty acids has been shown to activate PtdIns3K through coupling to G$_{\gamma}$ protein (Oh et al., 2010). It is noteworthy that CaSR activity was not detected in GM/IFN-$\gamma$-LPS–cultured cells, despite the presence of mRNA encoding the receptor. The absence of suitable antibodies precluded determination of whether the protein was expressed and properly targeted. It is also conceivable that ancillary proteins required for CaSR function may be absent or inhibited in GM/IFN-$\gamma$-LPS–cultured cells. Consistent with this hypothesis, we were able to induce macropinocytosis in the absence of calcium in GM/IFN-$\gamma$-LPS–cultured macrophages by activating other receptors that can stimulate PtdIns3K (Figure 8).

Superficially, the impaired constitutive macropinocytic ability of GM/IFN-$\gamma$-LPS–cultured macrophages would appear at odds with the well-established stimulatory effect of LPS, the quintessential pyrogen, on macrophages and dendritic cells (West et al., 2004; Zanoni et al., 2011). The solution to this apparent paradox likely lies in the temporal course of events. Indeed, we were able to induce macropinocytosis by acute treatment with LPS (coincident with the 10-min exposure to dextran; Figure 8) of macrophages cultured with GM-CSF and IFN-$\gamma$ only; like GM/IFN-$\gamma$-LPS–cultured macrophages, such cells do not perform constitutive macropinocytosis (Figure 3). A similar acute response to LPS was observed in M/IL4-cultured macrophages in the absence of calcium (Figure 8). These results resemble the reported transient increase in macropinocytosis in response to acute (minutes) stimulation of dendritic cells by LPS, that results in improved antigen uptake. Prolonged stimulation by LPS (hours), however, is accompanied by a dramatic suppression of macropinocytosis (West et al., 2004). The latter situation is more akin to the conditions used in our studies where chronic (2 d) exposure to LPS and IFN-$\gamma$ was used to generate pro-inflammatory macrophages, as described by others (Cassatella et al., 1990; Jaguin et al., 2013; McWhorter et al., 2013; Derlindati et al., 2015; Kaneda et al., 2016).

The acute versus chronic effects of LPS, and indeed the differences in the behavior between proinflammatory and anti-inflammatory macrophages, have important functional implications. In immature dendritic cells, as well as in alternatively activated macrophages, macropinocytosis serves for antigen sampling, as the phagocytes patrol the tissues. Upon activation by either TLR ligands or proinflammatory cytokines, however, the cells turn to an antigen-presenting function. The resulting mature dendritic cells devote their attention to processing the previously captured antigens, and carrying them to the secondary lymphoid organs to be presented to lymphocytes (Liu and Roche, 2015). During this stage, further uptake of antigens is unnecessary and macropinocytosis may in fact be counterproductive, as wholesale membrane internalization would antagonize the retention of presented antigens at the cell surface (Pan et al., 2013; Liu and Roche, 2015), and because cell migration and macropinocytosis appear to be mutually exclusive (Veltman et al., 2014; Veltman, 2015). We propose that suppression of macropinocytosis in proinflammatory macrophages plays a similar role, inasmuch as proteins involved in antigen presentation are expressed preferentially in M1-like cells compared with M2-like cells (Derlindati et al., 2015; Weichhart et al., 2015; Kaneda et al., 2016).

In addition to antigen sampling, macropinocytosis also plays a role in cellular metabolism, a function first identified in amoebae that has been conserved during evolution. Growth factor–induced macropinocytosis in macrophages, for example, delivers amino acids to lysosomes, resulting in increased activation of mTOR (Yoshida et al., 2015). The above-mentioned stimulatory effect of free fatty acids on macropinosome formation (Chung et al., 2015) likely also has an anabolic purpose. In this context, it is worth noting that the polarization of macrophages is accompanied by marked changes in cellular metabolism (Ouimet et al., 2015; Mills and O’Neill, 2016). We are tempted to speculate that the suppression of macropinocytosis contributes to this metabolic switch.

**MATERIALS AND METHODS**

**Reagents**

TMR-conjugated 70,000 MW dextrans, rhodamine phalloidin, and Lucifer Yellow CH lithium salt were purchased from Life Technologies (Carlsbad, CA). NPS 2143, pertussis toxin, LPS, latrunculin B, rapamycin, and the anti-$\beta$-actin monoclonal antibody (mAb) were from Sigma-Aldrich (St. Louis, MO). Wortmannin was from Millipore (Etobicoke, ON, Canada). Paraformaldehyde (PFA; 16% wt/vol) was from Electron Microscopy Sciences (Hatfield, PA). Monoclonal mouse anti-Rac1, anti-RhoA, and anti-Cdc42 antibodies were from the respective G-LISA activation kits (Cytoskeleton, Denver, CO). Monoclonal mouse anti-RhoG antibody (1F3 B3E5) was from Millipore. Rabbit polyclonal anti-p110$_{\beta}$, anti-p110$_{\gamma}$, and anti-p85 antibodies were prepared as described previously (Backer et al., 1992; Yu et al., 1998; Hill et al., 2000). Rabbit monoclonal anti-p110$_{\beta}$ (D1Q7R), anti-p110$_{\gamma}$ (DSSD5), and anti-phospho-Akt(Ser473) (#9271) antibodies were from Cell Signaling Technology (Danvers, MA). FITC-conjugated anti-human CD40 antibody (SC3), FITC-conjugated anti-mouse immunoglobulin G1 (IgG1) $\kappa$ isotype (MOPC-21), Alexa-647-conjugated anti-human CD163 antibody (GHI/61), and Alexa-647-conjugated anti-mouse IgG1 $\kappa$ isotype (MOPC-21) were from BD Biosciences (San Jose, CA). APC-conjugated anti-human CD80 antibody (2010), APC-conjugated anti-mouse IgG1 $\kappa$ isotype (MOPC-21), PE-conjugated anti-human CD200R antibody (OX-108), and PE-conjugated anti-mouse IgG1 $\kappa$ isotype (MOPC-21) were from BioLegend (San Diego, CA). Human recombinant M-CSF, GM-CSF, IFN-$\gamma$, IL-4, CCL7, and CCL19 were from PeproTech (Rocky Hill, NJ). EGTA was from BioShop Canada (Burlington, ON, Canada). U73122 was from Tocris Bioscience (Etobicoke, ON, Canada). HOE 694 was a gift from H. J. Lang (Aventis Pharma, Frankfurt am Main, Germany).

The Superecliptic Phluorin-CaSR (SeP-CaSR) construct was kindly provided by Jeremy Henley (University of Bristol, Bristol, UK). The ELMO-GST construct was a gift from Keith Burridge (University of North Carolina, Chapel Hill). Sources of other plasmids were as follows: GFP (Clontech Laboratories); GFP-Rac1 (Michaelson et al., 2001); GFP-RhoA, GFP-Cdc42, and RhoA-Q63L-GFP were kindly provided by John H. Brumell; GFP-RhoG (Heo et al., 2006); Rac1-G61L-GFP (Michaelson et al., 2001); Cdc42-G12V-YFP (plasmid #11399; Addgene); RhoG-G12V-CFP was obtained from the laboratory of Tobias Meyer (Stanford University, CA); Tiam1-GFP (plasmid #20154; Addgene).
Addgene); sGEF-GFP (Ellerbroek et al., 2004); Ph-Akt-GFP (Marshall et al., 2001); Ph-Akt-RFP (Flanagan et al., 2010); Lyn11-FRB (Marshall et al., 2001); FKBP-p85-CFP (plasmid #20159; Addgene); and GFP-2PABD with nuclear export sequence (Schlam et al., 2013).

BIM 46187 was synthesized as described in Schmitz et al. (2014). The forward (5′-CAGGATATAAATTTTCCGTTGGT-3′) and reverse (5′-GTACTGGGAGATGAGTTC-3′) primers for the CaSR reverse transcription-PCR (RT-PCR) and the forward (5′-TTCCAATATGGT-3′) and reverse (5′-CATACCAGGAAATGAGCTTG-3′) primers for the GAPDH RT-PCR were purchased from Integrated DNA Technologies (Coralville, IA). The GeneJet RNA purification kit and One-Step RT-PCR kit were from Thermo Fisher Scientific (Waltham, MA). The Superscript VILO cDNA synthesis kit was from Life Technologies (Carlsbad, CA). Rac1, RhoA, and Cdc42 G-LISA Activation Assay kits (colorimetric based) were from Cytoskeleton (Denver, CO).

**Solutions**

The calcium-containing medium was HBSS from Wisent (1.26 mM CaCl2; catalogue number 311-515-CL; MultiCell, St. Bruno, Canada). The calcium-free medium contained 1 mM KH2PO4, 154 mM NaCl, 5.6 mM Na2HPO4, 2 mM MgCl2, 2 mM EGTA, and 10 mM glucose, pH 7.2 at 37°C.

**Macrophage cell isolation, differentiation, and culture**

Peripheral blood mononuclear cells were isolated from the blood of healthy donors by density-gradient separation with Lympholyte H (Cedarlane, Burlington, ON, Canada). Monocytes were separated from other mononuclear cells in the washed buffy coat by adherence to tissue-culture–grade dishes at a density of 4.0 × 107 mononuclear cells per dish. Monocytes were plated in RPMI 1640 supplemented with 10% heat-inactivated fetal bovine serum (FBS) in case they were specified, with TMR-conjugated 70 kDa dextran (125 µg/ml), plus the drug used during the pretreatment. Unless specified otherwise, cells were incubated at 37°C for 15 min and then washed, fixed with 4% PFA and imaged immediately. Imaging was performed by spinning disk confocal microscopy on an Axiovert 200 M equipped with a 63× objective and a separate 1.5× magnifying lens (Carl Zeiss, Oberkochen, Germany). The microscope was fitted with a piezo focus drive and diode-pumped solid-state lasers (440, 491, 561, 638, and 655 nm; Spectral Applied Research, Richmond Hill, ON, Canada). Micrographs were acquired using a charge-coupled device camera (Hamamatsu Photonics, Hamamatsu, Japan) under the control of Velocity software. Macropinosomes were then quantified from the acquired images using the measurement tool in the ImageJ software (Schindelin et al., 2012), either from areas of two-dimensional (2D) particles from projection of three-dimensional (3D) stacks or from volumes of 3D particle analysis from 3D stacks, as specified in the figure legends.

**Flow cytometry**

Flow cytometers were gently lifted from tissue-culture plates by incubation with PBS (Ca2+- and Mg2+-free) supplemented with 10 mM EDTA at pH 7.4 for 15 min at 37°C. Cells then were washed three times with 2% bovine serum albumin (BSA; in PBS) and resuspended in 2% BSA (in PBS) at a concentration of ~106 cells per 100 µl and labeled with fluorophore-conjugated primary antibodies at the dilution specified by the manufacturer. Cells were incubated for 1 h at 4°C, followed by three washes with cold 2% BSA in PBS and fixation with 2% PFA in PBS; 30 min. Cells then were passed through a cell strainer, run through an LSR II flow cytometer (BD Biosciences, Franklin Lakes, NJ), and analyzed using FlowJo software (Tree Star, Ashland, OR).

**Transfections**

For human macrophages, DNA transfections were performed using the Neon electroporation system (Life Technologies, Carlsbad, CA). The cells were suspended using Accutase (Life Technologies, Carlsbad) and washed three times. Transfections were performed using the manufacturer’s recommended protocol. Transfection efficiency varied from plasmid to plasmid (10–60%); however, only transfected cells were selected for analysis throughout the paper. For HEK293 adherent cells, transfections were performed with FuGENE 6 transfection reagent (Promega, Madison, WI) according to the manufacturer’s instructions.

**PCR**

To detect the expression of CaSR mRNA in macrophages, one-step RT-PCR was performed. RNA was purified from macrophages using the GeneJet RNA purification kit (Thermo Fisher Scientific) and used as a template for the generation of cDNA using the Superscript VILO cDNA synthesis kit (Life Technologies) and subsequent PCR amplification using the One-Step RT-PCR kit from Invitrogen according to the manufacturers’ instructions. The primary sequences of the forward and reverse primers are listed in the Reagents section. The amplification products were visualized by electrophoresis on agarose gels prestained with ethidium bromide.

**Macropinocytosis assay**

Cells were suspended using Accutase and plated on 18-mm coverslips 24 h before performing the macropinocytosis assay. In the case of drug treatments, cells were pretreated as follows: HOE 694 (10 µM, 30 min), latrunculin B (2 µM, 30 min), Wortmannin (100 nM, 15 min), pertussis toxin (0.1 µg/ml, 16 h), BIM 46187 (10 µM, 1 h), and NPS 2143 (10 µM, 30 min). After pretreatment, the medium was aspirated and replaced with the calcium-containing medium, unless otherwise specified, with TMR-conjugated 70 kDa dextran (125 µg/ml), plus the drug used during the pretreatment. Unless specified otherwise, cells were incubated at 37°C for 15 min and then washed, fixed with 4% PFA and imaged immediately. Imaging was performed by spinning disk confocal microscopy on an Axiovert 200 M equipped with a 63× objective and a separate 1.5× magnifying lens (Carl Zeiss, Oberkochen, Germany). The microscope was fitted with a piezo focus drive and diode-pumped solid-state lasers (440, 491, 561, 638, and 655 nm; Spectral Applied Research, Richmond Hill, ON, Canada). Micrographs were acquired using a charge-coupled device camera (Hamamatsu Photonics, Hamamatsu, Japan) under the control of Velocity software. Macropinosomes were then quantified from the acquired images using the measurement tool in the ImageJ software (Schindelin et al., 2012), either from areas of two-dimensional (2D) particles from projection of three-dimensional (3D) stacks or from volumes of 3D particle analysis from 3D stacks, as specified in the figure legends.

**Rac1, RhoA, and Cdc42 activation assays**

Rac1, RhoA, and Cdc42 activity was measured with separate colorimetric G-LISA Activation Assay kits (Cytoskeleton, Denver, CO) according to the manufacturer’s instructions. In summary, cells were lysed, cellular debris was removed, and the samples were added to the wells of the Rho G-LISA plate coated with Rho-GTP–binding protein on ice. The plate immediately was placed on a shaker for 30 min at 4°C. Then the plate was washed three times at room temperature, and the retained GTP-bound GTPase was probed with either an anti-Rac1, anti-RhoA, or anti-Cdc42 antibody, followed by a secondary antibody tagged with horseradish peroxidase. The signal was measured with a SpectraMax 190 absorbance microplate reader (Molecular Devices, Sunnyvale, CA) at 490 nm after incubation of the well with the detection reagent. The measured absorbance was normalized to total protein levels based on the standard curve obtained with proteins provided with the kit manufacturer.

**RhoG activation assay**

Differentiated macrophages (day 7) were washed three times with ice-cold PBS and lysed with 500 µl (per 100-mm dish) of 10 mM PtdIns(3,4,5)P3 in polarized macrophages | 63
MgCl$_2$, 150 mM NaCl, 50 mM Tris, pH 7.4, 1% Triton X-100, 1 mM PMSF, and protease inhibitor cocktail (EDTA-free, cOmplete, Roche). Lysates were cleared at 14,000 × g for 10 min. Supernatants were rotated for 30 min with ~10 µg ELMO-GST (GST fusion protein containing the full-length RhoG effector ELMO) conjugated to glutathione-Sepharose beads (GE Healthcare). Beads were washed in 50 mM Tris, pH 7.4, 10 mM MgCl$_2$, 150 mM NaCl, 1% Triton X-100, and protease inhibitor cocktail. Full downs and lysates were then subjected to SDS–PAGE (10%) and immunoblotted with an anti-RhoG antibody.

**Immunoblotting**

Cells were washed three times with ice-cold PBS and subsequently lysed using a small volume of ice-cold RIPA buffer supplemented with protease inhibitor cocktail. Protein concentration was measured using the bicinchoninic acid assay, and 50 µg of protein of each sample was loaded and separated by 10% SDS–PAGE. The protein was transferred to a polyvinylidene difluoride membrane and blocked in Tris-buffered saline (TBS) containing 0.05% Tween-20 (TBST) and 10% fat-free milk for 30 min. The primary antibody was added to the membrane in 5% fat-free milk in TBST for 1 h at room temperature and washed 3 times with TBST for 10 min each. The membrane was reblocked with 10% milk in TBST for 15 min and the HRP-conjugated secondary antibodies were added in 5% milk in TBST for 45 min. The membrane was washed three times with TBST for 10 min each, and the blot was visualized on the Odyssey Fc (LI-COR, Lincoln, NE). Band intensity was quantified using the ImageStudio Lite software.

**Data presentation and statistics**

Unless otherwise indicated, images are representative of at least three independent experiments, with cells from different donors. Data presented in the text and graphs are the means with standard error of at least three independent experiments, representing the number of total cells as indicated in the figure legends. Unpaired t-tests were used to establish the significance of experimentally observed differences. Microscopy images were analyzed using ImageJ software (Schindelin et al., 2012) as indicated in the text.

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