Macrolets: Outsized Extracellular Vesicles Released from Lipopolysaccharide-Stimulated Macrophages that Trap and Kill *Escherichia coli*

**HIGHLIGHTS**
- Macrolets, outsized extracellular vesicles, release from LPS-stimulated macrophages
- Macrolets are rich in tetraspanin proteins such as CD81, CD63, and CD9
- Macrolets capture and internalize *E. coli* bacteria within acidic compartments
- Macrolets kill *E. coli* by a mechanism associated with production of ROS and superoxide

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Macrolels: Outsized Extracellular Vesicles Released from Lipopolysaccharide-Stimulated Macrophages that Trap and Kill Escherichia coli

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SUMMARY
Macrophages release a variety of extracellular vesicles (EVs). Here we describe a previously unreported class of EVs that are released from macrophages in response to Escherichia coli endotoxin, lipopolysaccharide (LPS), that we have named "macrolels" since they are extruded as large "droplets" released from macrophages. Morphologically, macrolels are anuclear, bounded by a single lipid membrane and structurally dependent on an actin cytoskeleton. Macrolels are enriched in tetraspanins and separable on this basis from their parent macrophages. Macrolels are distinguished from classic exosomes by their larger size (10–30 μm), discoid shape, and the presence of organelles. Macrolels are rich in both interleukin 6 (IL-6) and interleukin 6 receptor (IL-6R), and are capable of trapping and killing E. coli in association with production of reactive oxygen species. Our observations offer insights into the mechanisms by which macrophage activities may be amplified in sites of infection, inflammation, and healing.

INTRODUCTION
Macrophages play a pivotal role in the initiation, resolution, and persistence of inflammation (Dalli and Serhan, 2017; Ginhoux and Jung, 2014; Varol et al., 2015). Well-documented functions of macrophages include the production and secretion of cytokines and chemokines, phagocytosis of pathogens or dead/dying cells, and elaboration of extracellular matrix metalloproteinases (Ariel and Serhan, 2012; Hundertmark et al., 2018; Tomlin and Piccinini, 2018; Wight et al., 2017; Wynn and Barron, 2010). In response to danger signals such as E. coli endotoxin (lipopolysaccharide, LPS), macrophages lay extracellular traps (Doster et al., 2018; Sharma et al., 2017) and produce extracellular vesicles (EVs) containing a variety bioactive molecules (e.g., proteins, carbohydrates, lipids, and nucleic acids) that can influence local inflammatory responses in tissue and lead to phenotypic change in target cells (Esser et al., 2010; Ismail et al., 2013; O’Neill and Quah, 2008). These observations provide evidence for a wide variety of mechanisms by which macrophages are able to sense changes in their surrounding microenvironment and have their signature functions amplified and coordinated.

With respect to EVs, four categories have been reported, including exosomes, microvesicles or microparticles, apoptotic bodies, and oncosomes (Akers et al., 2013; Dreyer and Baur, 2016). All but oncosomes are produced by macrophages and are emerging as potentially consequential mediators in communications between macrophages and other cell types (Lanyu and Feilong, 2019; Zhu et al., 2017). EVs produced by macrophages are small relative to the diameter of their parent cells, with sizes ranging from 50 to 100 nm for exosomes (Bhatnagar et al., 2007), 200 to 1,000 nm for microvesicles (Ismail et al., 2013), and 1,000 to 5,000 nm for apoptotic bodies (Zhu et al., 2017). Recent studies, however, also provided evidence that EV classes of larger dimensions may also be released by human primary monocyte-derived dendritic cells (Kowal et al., 2016), with separation based on their enriched expression of tetraspanins such as CD63, CD81, or CD9. More recently, it has been shown that malignant cells can produce larger EVs or oncosomes (1–10 μm) that have an organized cytoskeleton and contain organelles (Johnson et al., 2017) or, in some cases, can be detected in the circulation of patients with cancer (Vagner et al., 2018). These findings suggest that larger classes of EVs might be produced by other cell types, including the macrophage.

Here we report a class of large EVs (10–30 μm) produced by human and mouse macrophage cell lines and primary human monocytes transformed to macrophages ex vivo and released in response to...
stimulation by LPS. We have named these EVs "macrolets," since, as demonstrated below, they appear as large droplets released from "macrophages." Macrolets are distinguished from currently known extracellular traps and EVs based on their size, morphology, contents, and apparent mode of biogenesis. As has been reported for exosomes (Kowal et al., 2016), macrolets are enriched in tetraspanins CD63, CD81, and CD9; have an organized cytoskeletal structure, and contain organelles such as mitochondria, lysosomes, and secretory compartments. In response to LPS stimulation, macrolets produce cytokines such as interleukin-6 (IL-6) as well as interleukin-6 receptor (IL-6R). Moreover, we show that macrolets are capable of trapping microbes such as E. coli, in association with classic bactericidal functions such as vesicular acidification and production of reactive oxygen species. These findings offer insights into novel and potentially multifunctional vehicles by which individual macrophages may communicate with other cells and amplify its danger signals within a space of inflammation and healing.

RESULTS

Macrolets: Ousized Extracellular Vesicles Released by Macrophages in Response to E. coli Endotoxin (LPS)

Under light microscopy, in both static images and in time-lapse imaging, we observed release of large EVs from human THP-1 macrophages after exposure to LPS (100 ng/mL, 4 h) as shown in Figure 1A. They did not contain nuclei (i.e., were DAPI-negative). As shown in time-lapse recordings (Videos S1A and S1B) they first appeared as hyperdense droplets forming out of membrane and cytoplasm and, once extruded, rapidly expanded to form discoid particles. Based on these initial observations we named these outsized EVs "macrolets," large cell droplets released from macrophages.

Staining with phalloidin (Figure 1B) indicated that macrolets have an actin cytoskeleton providing structure both at the borders and in the interior. Under these conditions, macrolets were flattened, discoid structures (Figure 1C and Video S2), with thickness of a few micrometers and diameters ranging from 10 to 30 μm. Internal cytoskeletal organization was variable: a large proportion of macrolets have a well-organized cytoskeletal structure at their boundaries with relatively sparse interior staining (Figure S1A), whereas some macrolets had an outer "shell" and an interior "core" with somewhat fragmented staining (Figure S1B). In addition, macrolets were bounded by intact membrane: as shown in Figure 1D, the sides of the thin, discoid macrolet stain diffusely with the fluorescent, membrane-lipid reporter DiO'; DiOC18(3) (3, 3'-Dioctadecyloxacarbocyanine Perchlorate) (Vemula et al., 2014). Moreover, we found that macrolets are also rich in Annexin V (Figure 1E and Video S3). Annexin V is a Ca2+-dependent phospholipid-binding protein with high affinity for phospholipid phosphatidylserine (PS). Consistent with studies characterizing other EV proteins, Annexin V has been utilized for identification and purification of exosomes and microvesicles (Fitzner et al., 2011; Garzetti et al., 2014; Kowal et al., 2016; Pieters et al., 2019; Thery et al., 2018). Lastly, formation of macrolets was observed much less frequently in the absence of pre-treatment with LPS (Figures 1F and 1G).

To confirm macrolets were not a product of cell death, we induced apoptosis in THP-1 cells with etoposide (50 μM) for 4 h (Zhuang et al., 1998) and pyroptosis with a combination of LPS (100 ng/mL) and Nigericin (10 μM) (Bergsbaken et al., 2009; Broz, 2015; Cullen et al., 2015). As shown in Figure S2A, apoptosis was characterized by nuclear fragmentation and disruption of the cytoskeleton, which was distinct from anuclear macrolets with intact cytoskeletal structure. As shown in Figure S2B, pyroptosis was characterized by caspase-1 activation and associated with membrane and cytoskeletal disruption but no macrolets were detected as seen in cells treated with LPS alone (Figure S2C). In addition, we verified that the macrolets are produced by viable parent macrophages showing both Annexin V and PI negative staining (Figure S2D). These studies confirm that formation of the macrolets is an active process and not a by-product of two key pathways that lead to cell death.

Organelles and Membrane Markers of Macrolets

Considering their large size and structure, we asked whether macrolets contain subcellular organelles such as endoplasmic reticulum (ER), lysosomes, and mitochondria. In immunofluorescence studies (Figure 2A), calnexin, a principal protein-folding chaperone in the ER, was localized proximal to the nucleus in LPS-stimulated THP-1 cells; however, macrolets stained diffusely for calnexin, in contrast to exosomes that do not contain calnexin (Wen et al., 2017). Using immunofluorescent staining with the lysosomal markers Lamp-1 and the vital fluorescent reporter LysoTracker, we also detected lysosomes (Figures 2B and 2C). Presence of lysosomes was consistent with diffuse immunofluorescent staining for the vacuolar H⁺-ATPase (Figure S3), the principal proton transporter responsible for the acidification of the interior of lysosomes, autophagic vacuoles, and secretory compartments (Sun-Wada and Wada, 2013). Lastly, using the vital fluorescent
reporter MitoTracker Deep Red, we found macrolets also contained mitochondria (Figure 2D). These cyto-
logical surveys provide evidence that the macrolet contains organelles associated with signature macro-
phage functions such as protein secretion and degradation, as well as uptake and killing of bacteria.

To confirm these observations and further explore macrolet ultrastructure, transmission electron micro-
scopy (TEM) was performed. Profoundly different from macrophages as shown in Figures 2E and 2F,
Macrolets were organized into “core” and “shell” domains. Unlike the lipid bilayer membrane of the macrophage, the core of the macrolet was delimited by a single membrane encompassing the entire macrolet (Figure S4A). Moreover, mitochondria, vesicular structures, and autophagic vacuoles were also observed (Figure 2G) confirming our observations using vital reporters and immunofluorescent imaging. At high-power magnification (Figure S4B), interesting differences in mitochondrial morphology were observed showing that mitochondria located in macrophages were elongated, whereas those located in macrolets were shorter and fragmented, potentially reflecting differences in their accessibility to LPS or differential influences of LPS on metabolism (Kang et al., 1990, 1992).

To explore potentially common origins between macrolets and exosomes, we performed immunofluorescent imaging for tetraspanin proteins (Kowal et al., 2016). The tetraspanin CD63 localized predominantly at the cell membrane of the macrophage, and proximal to the outer border of the macrolet (Figure 3A). A similar staining pattern was observed for CD81 (Figure 3B); however, CD9 exhibited a relatively diffuse staining pattern within the macrolet (Figure 3C). To stratify macrolet populations based on expression levels of CD81 and CD63, we performed flow cytometry. Culture medium was collected from THP-1
macrophages after LPS stimulation and was centrifuged at 5,500 g for 25 min to pellet the insoluble fraction. As shown in Figure 3D, we first eliminated cell debris and the nucleus-positive population with 7-aminoactinomycin D (7-AAD) labeling and found that 85.6% of macrolets express both CD81 and CD63, 13.2% of macrolets are CD81+CD63-, 0.6% of them only express CD63 protein, and the remaining 0.6% of macrolets express neither CD81 nor CD63.

In addition, as shown in Figure 3D, we found only a very small number of macrolets in the media bathing LPS-stimulated macrophages, whereas great numbers were adherent to the culture plates. We made trials of several methods to detach macrolets from their culture dishes (trypsin EDTA, Accutase, and Accumax, which is the concentrated Accutase). Using these methods, we were only able to obtain fragments and not enough intact macrolets for separation by density gradients or by fluorescence-activated cell sorting.

**Functional Capabilities of Macrolets: Production of Pro-inflammatory Cytokines and Responses to Live E. coli**

As release of IL-6 and expression of the IL-6 receptor (IL-6R) are among the earliest and potentially actionable cytokine responses to trauma and infection (Namas et al., 2013; Netea et al., 2017; Prenissl et al., 2019; Zhang et al., 2014), we determined the relative expression of IL-6 and its receptor in an enriched population of macrophages after LPS stimulation and was centrifuged at 5,500 g for 25 min to pellet the insoluble fraction. As shown in Figure 3D, we first eliminated cell debris and the nucleus-positive population with 7-aminoactinomycin D (7-AAD) labeling and found that 85.6% of macrolets express both CD81 and CD63, 13.2% of macrolets are CD81+CD63-, 0.6% of them only express CD63 protein, and the remaining 0.6% of macrolets express neither CD81 nor CD63.

Figure 3. Tetraspanins Such as CD9, CD63, and CD81 Are Highly Expressed in Macrolets
(A) Representative immunofluorescent images of THP-1 macrophages treated with LPS (100 ng/mL) for 4 h showed that CD63 (green) was expressed on the cell membrane of THP-1 macrophages and the surface of macrolets (arrow); Alexa Fluor 594-phalloidin (F-actin; red); scale bar, 10 μm.
(B) Representative immunofluorescent images of THP-1 macrophages treated with LPS for 4 h showed that CD81 (green) is predominantly expressed on the cell membrane of THP-1 macrophages and the surface of macrolets (arrow); Alexa Fluor 594-phalloidin (F-actin; red); scale bar, 10 μm.
(C) Representative immunofluorescent images of THP-1 macrophages treated with LPS for 4 h showed that CD9 (green) is not expressed on the cell membrane of THP-1 macrophages or at the surface of macrolets (arrow); Alexa Fluor 594-phalloidin (F-actin; red); scale bar, 10 μm.
(D) Culture medium was collected from LPS-stimulated THP-1 macrophages (LPS 100 ng/mL, 4 h), and the medium was centrifuged at 5,500 g for 25 min to pellet the insoluble fraction prior to flow cytometry assays. To analyze macrolet population, the cell debris and clumped cells were eliminated with side and forward scatters (left panel), then 7-AAD staining was used to exclude dead cell bodies that contain either intact or damaged nucleus (middle panel), and the 7-AAD macrolet population was further separated with CD81-PE and CD63-FITC channels (right panel).
Figure 4. Macrolets Express both IL-6 and IL-6R and Are Able to Trap and Kill E. coli Bacteria

(A) Representative immunofluorescent images show that macrolets express IL-6 (gray). Alexa Fluor 594-phalloidin (F-actin; red); scale bar, 10 μm.

(B) Representative immunofluorescent images show that macrolets express IL-6R (green). Alexa Fluor 594-phalloidin (F-actin; red); scale bar, 10 μm.

(C) Representative immunofluorescent images show that IL-6R (green) is profoundly co-localized with ADAM-17 (red) in macrolets. Ten macrolets were randomly picked for fluorescence intensity and co-localization analysis using Imaris 9.0 software showing Pearson’s coefficient = 0.70 ± 0.09. Data were obtained and analyzed from three independent experiments.

(D) Representative images show that E. coli bioparticles were captured by both CD81-GFP+ macrophages and macrolets (arrow); scale bar, 10 μm.

(E) THP-1 macrophages were co-cultured with pH-sensitive E. coli (pHrodo) for 4 h. Confocal images were collected to assess the internalization of pHrodo E. coli in both macrophages and macrolets. Representative images show that pHrodo E. coli were taken up into acidic vacuoles in both macrophages and macrolets; scale bar, 10 μm.

(F) THP-1 macrophages stimulated with LPS were co-cultured with live E. coli (strain HB101) for 4 h. The viability of E. coli bacteria was assessed using propidium iodide staining (P.I., red; dead bacteria) and compared with SYTO-9 staining (green, total bacteria); scale bar, 10 μm. About 89.7% of bacteria engulfed by macrophages (n = 20) were dead, and 69.3% of bacteria captured by macrolets (n = 20) were dead. Data were obtained and analyzed from three independent experiments.

(G) THP-1 macrophages were stimulated with 100 ng/mL of LPS for 4 h, and the production of reactive oxygen species (ROS; red) and superoxide (green) was assessed using fluorescent staining. Representative images show that macrophages (left panel) and macrolets (right panel) were able to produce ROS and superoxide; scale bar, 10 μm.
of macrolets. Immunofluorescent studies illustrated that both IL-6 (Figure 4A) and IL-6R (Figure 4B) were detected in macrolets sourced from LPS-stimulated THP-1 cells.

The interaction of IL-6 with its receptor has two dimensions (Garbers and Rose-John, 2018): a classic signaling pathway, which refers to interactions with IL-6R (full-length) localized at the cell membrane, and the so-called trans-signaling pathway that refers to interactions with a soluble form of IL-6R and assembly of a soluble pro-inflammation complex with the ubiquitously expressed receptor subunit gp130. Formation of the soluble form results from cleavage of the full-length form, mediated by metalloproteinases such as ADAM-10 and ADAM-17 (Garbers et al., 2011; Schumacher et al., 2015). To demonstrate the capability of macrolets to generate the components of both pathways, we used immunofluorescent staining to show that ADAM-17 was highly expressed in macrolets sourced from LPS-stimulated THP-1 cells and localized in close proximity to IL-6R (Figure 4C). The Pearson's coefficient for the colocalization between ADAM-17 and IL-6R was 0.70 ± 0.09, suggesting these two proteins were predominantly located in the same compartments within the macrolets.

Based on their robust release and cytokine signals in response to soluble LPS, we hypothesized that macrolets would harbor mechanisms for trapping microbes such as E. coli, thereby amplifying the capabilities of the stimulated macrophage. To track the behavior of macrolets, we developed a stable THP-1 cell line expressing a green fluorescent protein-tagged CD81 (CD81-GFP). As shown in Figure S5, CD81-GFP expression localized to the cell surface in macrophages and proximal to the periphery in macrolets, with patterns similar to those observed for endogenous CD81. In addition, transduction of cells with the GFP-CD81 construct did not interfere with the discoid morphology of the macrolet as shown in Video S4.

Using GFP-CD81 constructs as tools for visualizing macrophages and macrolets, we were able to show that macrolets are able to capture E. coli particles (K-12 strain) conjugated with Texas Red (Figure 4D). In further studies, microbes were loaded with the pH-sensitive fluorescent reporter pHrodo Red E. coli bioparticles, which turn red when bacteria are internalized in acidic compartments such as phagolysosomes (Figure 4E). We found a similar degree of pHrodo Red E. coli bioparticles activated in macrophages (88.4%) and macrolets (83.3%), illustrating the biological activity of phagocytosis in macrolets. To test whether the bacteria trapped in macrolets were vulnerable to bactericidal activity, we co-cultured LPS-stimulated macrophages with live E. coli for 4 h and then measured bacterial viability in macrophages and macrolets using propidium iodide (P.I.) to stain for dead bacteria and SYTO-9 to stain for total bacteria. As shown in Figure 4F, we found that 89.7% of bacteria were killed after engulfment by macrophages, whereas 69.3% of E. coli captured by macrolets were not viable. Also, as shown in Figure 4G, similar to macrophages, macrolets were capable of generating both reactive oxygen species (ROS) and superoxide, which in conjunction with acidification, would be the likely mechanism of bacterial killing within a phagolysosome.

**Macrolets Are Detected in Other Models of Monocyte/Macrophage Function**

In a final set of studies, we found that macrolets were also detected following LPS stimulation of RAW 264.7 cells (Figure S6A), a mouse monocyte/macrophage lineage, and human primary monocytes transformed to M1 macrophages by pre-treatment with human recombinant macrophage colony stimulating factor (M-CSF) for 4 days, and then treated with LPS and interferon γ (IFN-γ) for additional 48 h (Figure S6B). From these studies it seems reasonable to infer that biogenesis of macrolets following LPS stimulation is not species specific, nor is it a property only of transformed cells.

**DISCUSSION**

Our studies provide in vitro evidence of a population of outsized (10–30 μm) EVs, released in response to LPS, that share attributes of smaller exosome populations but that also have capabilities attributed to a macrophage. To our knowledge, this is the first report providing details of their morphology, structure, composition, and potential range of functions. Our studies identify macrolet populations as distinct from exosomes and other EVs by size and the presence of organelles such as ER, mitochondria, and phagolysosomes. Also excluded by our studies is the possibility that they represent by-products of dying cells (Bergsbaken et al., 2009; Broz, 2015). Lastly, their morphology, delimitation by a lipid membrane, the lack of association with dying cells, and presence of organelles distinguish macrolets from extracellular traps despite the shared characteristic of microbe trapping (Brinkmann and Zychlinsky, 2012; Doster et al., 2018; Sharma et al., 2017).
The production of such "macrolets" by cells of the innate immune system has been suggested recently, through methods of separation and proteomic analysis (Kowal et al., 2016). Our studies demonstrate that macrolets are capable of producing pro-inflammatory cytokines such as IL-6 and IL-6R that have also been linked to the initial responses to infection, trauma, and shock (Garbers et al., 2011; Prenissl et al., 2019; Schumacher et al., 2015; Zhang et al., 2014). Strikingly, macrolets also express critical metallo-proteins such as ADAM-17, which is critical in the processing of the IL-6R and other extracellular signals that constitute the "sheddome" (Gutierrez-Lopez et al., 2011). In addition, we find that macrolets are capable of trapping and killing bacteria, in association with well-recognized bactericidal mechanisms such as phagosomal acidification and production of ROS. Underpinning their functional capabilities, macrolets contain organelles such as ER, mitochondria, phagosomes, and lysosomes, which suggests the potential for flexibility and a kind of sustainability in response to injury or infection. These general considerations suggest that macrolets may serve as extenders and amplifiers of macrophage functions—such as production of cytokines or phagocytosis of foreign material—in injured tissue or spaces such as peritoneum or pleura in response to infection or irritation. In addition, it offers the possibility that larger EV populations such as macrolets might serve as important modifiers and therapeutic targets for disease states driven by failed resolution of innate immunity (Serhan, 2017a, b; Serhan et al., 2018).

These observations raise three additional issues for discussion. First, we have found that macrolets, although present in small numbers under baseline conditions, are released in substantial numbers following stimulation with LPS. Among macrophages, the principal mechanism for recognizing and responding to LPS is the Toll-like receptor 4 (TLR4) pathway, which plays a pivotal role in the sensing and processing danger and harmful signals known as pathogen-associated molecular patterns (PAMPs) (Jaekal et al., 2007; Lu et al., 2008; Muta and Takeshige, 2001). Based on these considerations, it is likely that the biogenesis of macrolets is mediated by the TLR4 pathway, and the underlying mechanism needs to be further elucidated. Also, it would be of great interest to know whether macrolets can be released in response to other PAMPs or danger signals and whether macrolets formed under different stimuli carry different sets of bioactive molecules (e.g., miRNA, cytokines, and chemokines) to communicate different messages to target tissues and cells.

A second issue is the role of tetraspanins in facilitating macrolet function. Tetraspanins play potentially critical roles in the adhesion of a wide variety of bacterial species to host cells (Green et al., 2011; Hassuna et al., 2017). Both anti-CD63 monoclonal antibody and knockdown of CD63 in human monocyte-derived macrophages (MDM) greatly inhibited the binding capacity between Salmonella typhimurium and MDM (Hassuna et al., 2017). We found that tetraspanin proteins CD63, CD81, and CD9 were enriched in macrolets, suggesting that they may serve similar roles as adhesion molecules to capture pathogenic microbes. More broadly, tetraspanins have been implicated in biogenesis of exosomes, assembly of compartments, and selection of proteins such as ADAM-17 for recruitment and stabilization at the membrane (Andreu and Yanez-Mo, 2014; Gutierrez-Lopez et al., 2011; Perez-Hernandez et al., 2013; Tsukamoto et al., 2014). Our observations indicate that therapeutic targeting of exosomes through their expression of tetraspanins (Rana et al., 2012; Rana and Zoller, 2011; Yim and Choi, 2016) may need to take into account their enrichment in larger EV populations (Kowal et al., 2016) and the potential for unintended/off-target effects.

A third issue is the mechanism by which, following release, a macrolet may be transported to its site of activity as an extender/amplifier of the functions of the source macrophage. Although we have no direct evidence for motility of macrolets too far beyond the source macrophage, it is intriguing that they have a discoid morphology. By analogy with the erythrocyte, it is tempting to speculate that such a morphology implies a high surface to volume ratio (Diez-Silva et al., 2010), which may, in turn, facilitate large reversible elastic formation as the macrolet engages with its microenvironment. Our studies provide evidence that shape is maintained by an actin cytoskeleton, suggesting that this discoid shape may not be a response only to the two-dimensional platform of a microscope slide. Further studies would be needed to explore whether shape and biomechanical characteristics are observed in three-dimensional organ culture models and how they influence function.

Limitations of the Study
A principal limitation of this study is that most of our observations have been made in an in vitro system, using THP-1 macrophage-like cells exposed to a danger signal such as purified LPS. Extending these observations to more complex conditions of cell co-culture and in vivo models of infection and sepsis...
will provide more detailed information about the conditions leading to formation and release, longevity, and biological importance of this new class of EVs. At the cellular level, it will be of great interest to determine their mechanisms of biogenesis and release.

**METHODS**
All methods can be found in the accompanying Transparent Methods supplemental file.

**SUPPLEMENTAL INFORMATION**
Supplemental Information can be found online at https://doi.org/10.1016/j.isci.2020.101135.

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**AUTHOR CONTRIBUTIONS**
Conceptualization, W.D., and D.I.S.; Methodology and Investigation, W.D. and O.C.R.; Writing – Original Draft, W.D.; Writing – Review and Editing, W.D., S.L.K., and D.I.S.; Funding Acquisition, D.I.S.; Supervision, D.I.S.

**DECLARATION OF INTERESTS**
The authors declare no competing interests.

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Supplemental Information

Macrolets: Outsized Extracellular Vesicles

Released from Lipopolysaccharide-Stimulated Macrophages that Trap and Kill *Escherichia coli*

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Figure S1. Related to Figure 1

**Macrollets are outsized heterogeneous extracellular vesicles.** (A) THP-1 macrophages were treated with 100 ng/ml of LPS for 4 h, and macrophages and macrolets were fixed with 4% formaldehyde and stained with Alexa Fluor-594 phalloidin (F-actin; red) and DAPI (blue). The large proportion of macrolets have a well-organized structure with relatively sparse interior F-actin staining. (B) A small proportion of macrolets have an outer “shell” and an interior “core” with somewhat fragmented F-actin staining (arrow). Multiple images were collected (n>10), scale bar = 10 μm.
Figure S2. Related to Figure 1

Macrolets were not products from apoptosis or pyroptosis. (A) Apoptosis was induced with Etoposide (50 μM) for 4 h, and apoptotic cells were stained with phalloidin (F-actin; red) and DAPI (nuclear DNA; blue). A typical apoptotic cell (arrow) was characterized by morphological changes such as nuclear DNA fragmentation and cytoskeleton destruction, scale bar = 10 μm. (B) Pyroptosis was induced with 100 ng/ml LPS for 3 h, followed by 10 μM Nigericin for 1 h. Cells were fixed with 4% formaldehyde and then stained with caspase-1 (green), phalloidin (red) and DAPI (blue). Pyroptotic cells were characterized by cell body enlargement, cell membrane rupture, and cytoskeleton destruction. Multiple images (n=5) were collected, scale bar = 10 μm. (C) Cell integrity was well-maintained and macrolets were generated when THP-1 macrophages were treated with LPS alone. Multiple images (n=5) were collected, scale bar = 10 μm. (D) Annexin V+ macrolet (arrow) was released from a viable parent macrophage (Mö) with both Annexin V and PI negative labels. Multiple images (n=5) were collected, scale bar = 10 μm.
Figure S3. Related to Figure 2

**Macrolets contain vacuolar H⁺-ATPase.** A representative image shows that both macrophages and macrolets (phalloidin; red, arrow) have vacuolar H⁺-ATPase (V-ATPase B1; green). Multiple images (n=5) were collected. Scale bar = 10 μm.
Figure S4. Related to Figure 2

(A) Macrolets contain an interior core that is delimited by a single membrane. Transmission electron microscopic (TEM) image shows that a macrolet is composed of a central core structure containing a variety of organelles and is delimited by a single membrane (highlighted with a dotted red line). Scale bar = 2 μm. (B) Macrolets contain shorter fragmented mitochondria instead of elongated ones carried by macrophages. Transmission electron microscopic (TEM) images show that mitochondria (arrow) are present in macrolets and are fragmented (left side image), which is distinct from the elongated ones that are carried by macrophages (right side image), scale bar = 0.2 μm.
Figure S5. Related to Figure 4

Lentiviral overexpression of CD81-GFP did not interfere with the discoid morphology of the macrolets. CD81-GFP was stably expressed in THP-1 macrophages by the lentiviral transduction system. CD81-GFP expression is localized to the cell membrane of macrophages and exhibits the same expression pattern as endogenous CD81 protein as shown in Figure 3B. Scale bar = 10 μm.
Figures S6. Related to Figure 1

Mouse and human macrophages are capable of producing macrolets. (A) Mouse RAW 264.7 macrophages were treated with 100 ng/ml of LPS for 4 h, and cells were fixed with 4% formaldehyde and stained with Alexa Fluor 594 phalloidin (red) and DAPI (blue). Representative images show that multiple macrolets (arrow) were released upon LPS treatment, and F-actin\(^+\) macrolets are anuclear particles (DAPI\(^-\)). (B) Human primary CD14\(^+\) monocytes were differentiated into M1 macrophages. Representative images show that IL-6R (green) is expressed in F-actin\(^+\) macrolets (phalloidin; red) released from human primary M1 macrophages. Multiple images (n>5) were collected. Scale bar = 10 µm.
TRANSPARENT METHODS

Cells and cell culture

Human monocytic leukemia THP-1 cells were purchased from ATCC (TIB-202), and maintained in RPMI 1640 medium supplemented with 10% heat-inactivated fetal bovine serum containing 1% penicillin/streptomycin, 1×GlutaMax-I, 10 mM HEPES (pH 7.4), and 1× MEM-Non essential amino acids NEAA (Gibco). Mouse macrophage-like RAW264.7 cells were purchased from ATCC (TIB-71), and maintained in the same culture medium as for THP-1 cells. Human CD14+ primary monocytes (70035) were purchased from StemCell technology (Vancouver, Canada) and differentiated into M1 macrophages using a 6-day standard protocol as recommended by the manufacturer. In brief, primary monocytes were cultured in serum-free macrophage medium (StemCell, 10961) and plated in 24-well plates (5×10⁵ cells/well), then treated with 50 ng/ml of human recombinant macrophage colony-stimulating factor (M-CSF) for 4 days. The medium was replaced with fresh macrophage culture medium containing 10 ng/ml lipopolysaccharide (LPS, Escherichia coli O127:B8, Sigma-Aldrich, L5024) and 50 ng/ml IFN-γ for additional 2 days to differentiate monocytes into M1 macrophages.

Time-lapse image collection

Time-lapse images were collected using DeltaVision microscopy. In brief, THP-1 monocytes were placed in a 35-mm bottom slide dish (MetTek, P35G-0-20-C), and were differentiated with 50 ng/ml of phorbol 12-myristate 13-acetate (PMA, Sigma-Aldrich, P8139) for 48 h. The culture medium was replaced with pre-warmed Opti-MEM I supplemented with 10% FBS, and 100 ng/ml of LPS was added. Images were captured at 3-min intervals for total 4.5 h. The video was created using Volocity software.

Transmission electron microscopy

THP-1 cells (5×10⁵ cells/well) were cultured in a MatTek bottom glass slide (MetTek, P35G-1.5-14-C-GRID), then differentiated with PMA and stimulated with 100 ng/ml of LPS for 4 h. Samples were fixed by 2.5% glutaraldehyde and 2% paraformaldehyde in 0.1 M cacodylate buffer (pH 7.4) and further fixed in 1% osmium tetroxide in 0.1 M cacodylate buffer (pH 7.4) for 1 h. Samples were dehydrated in a graduated ethanol series, and embedded in LX112 (Ladd Research, Williston, VT). Thin sections (70 nm) were stained with uranyl acetate and lead citrate and images were captured using a JEOL JEM1400 Transmission Electron Microscope (JEOL USA Inc., Peabody, MA, USA).

Immunofluorescence staining and confocal microscopy

THP-1 cells (5×10⁴ cells/well) were cultured in 24-well plates containing a cover glass (Fisher Scientific, 12-545-83). After indicated treatments, cells were fixed in 4% methanol-free formaldehyde (Thermo Scientific Pierce, 28908) prepared in 1×PBS for 15 min. A containing lentiviral particles was collected and filtered with low-protein binding endotoxin-free 0.45 μm PES filters. Lentiviral particles were concentrated with the Lenti-X concentrator kit (Takara, 631231). THP-1 monocytes (1×10⁷ cells) were transduced with 20 μl of concentrated lentivirus in 2 ml culture medium containing 8 μg/ml of polybrene for 20 min at room temperature. The spinoculation protocol was used by centrifuging cell and virus mixture for 30 min at 800×g at 32°C. 48 h after lentiviral transduction, 0.5 μg/ml of puromycin was added for selecting CD81-GFP positive cell pools.
MitoTracker, LysoTracker, and DiO plasma membrane staining

Mitochondria and lysosomes were stained with 500 nM of MitoTracker Deep Red FM and 75 nM of LysoTracker Red DND-99 (ThermoFisher Scientific, L7528) following the manufacturer’s recommendation. Vybrant DiO cell-labeling solution (ThermoFisher Scientific, V22886) was used to stain membrane lipids. In brief, 5×10⁵ cells were seeded on 35-mm Bottom Slide Dishes, and after the indicated treatment cells were stained with DiO solution diluted in Opti-MEM medium for 15 min. Dishes were washed with 1×PBS 3 times, and the confocal images were collected using a Leica TCS SP8 laser scanning fluorescence microscope.

Flow cytometry

THP-1 monocytes were differentiated to macrophages with 50 ng/ml of PMA for 48 h, and then treated with 100 ng/ml of LPS for 4 h to induce macrolet production. Culture medium was centrifuged at 5,500 × g for 25 min to pellet the insoluble fraction. Pellets were washed with 1×PBS and re-suspended in 100 µl of flow cytometry buffer (1×PBS, 2% FBS, 0.09% sodium azide), and then labeled with CD81-PE (BioLegend, Cat#349505, clone 5A6, 1:100 dilution), CD63-FITC (BioLegend, Cat#363006, clone H5C6, 1:100 dilution), and 1 µl of 7-aminoactinomycin D (7-AAD, ThermoFisher Scientific, Cat#A1310, 1 mg/ml) on ice for additional 20 min in dark. Flow cytometry was performed using BD LSRFortessa and data were analyzed with FlowJo software (BD Biosciences).

Bacteria trapping and killing assays

For bacteria trapping assay, THP-1 cells (5×10⁵ cells/well) expressing CD81-GFP were plated on glass-bottom slides. Release of macrolets was induced with 100 ng/ml of LPS for 1 h, then E.coli (K-12 strain) bioparticles conjugated with Texas Red (ThermoFisher Scientific, Cat# E2863) were added to the culture medium for additional 3 h followed by Hoechst 33324 (NucBlue) nuclear staining for 15 min. The bacteria trapping was visualized with Leica confocal microscopy. For bacterial killing assays, THP-1 cells were cultured in antibiotic-free culture medium for three passages to wash away the residual antibiotics. THP-1 monocytes (5×10⁵ cells/well) were plated in 35-mm bottom slide dish, and differentiated into macrophages with 50 ng/ml of PMA for 48 h, then 5 µl of live E.coli bacteria (strain# HB101; OD 670 nm of 0.03) were incubated with LPS-stimulated THP-1 macrophages for 4 h, and bacterial killing capability was assessed using Live/Dead BacLight Bacterial Viability Kit with SYTO-9 and propidium iodide dyes (ThermoFisher Scientific, Cat# L7012).

Statistical analysis

Data presented are mean ± SD or SE as indicated in figure legends. The number for each experiment is noted in the figure legends. Student’s t test was used for statistical analysis, and a significant difference was determined at p value < 0.05.