Microvesicle-Mediated Communication Within the Alveolar Space: Mechanisms of Uptake by Epithelial Cells and Alveolar Macrophages

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Intra-alveolar microvesicles (MVs) are important mediators of inter-cellular communication within the alveolar space, and are key components in the pathophysiology of lung inflammation such as acute respiratory distress syndrome (ARDS). Despite the abundance of data detailing the pro-inflammatory effects of MVs, it remains unclear how MVs interact or signal with target cells in the alveolus. Using both in vivo and in vitro alveolar models, we analyzed the dynamics of MV uptake by resident alveolar cells: alveolar macrophages and epithelial cells. Under resting conditions, the overwhelming majority of MVs were taken up by alveolar macrophages. However, following lipopolysaccharide (LPS)-mediated inflammation, epithelial cells internalized significantly more MVs (p<0.01) whilst alveolar macrophage internalization was significantly reduced (p<0.01). We found that alveolar macrophages adopted a pro-inflammatory phenotype after internalizing MVs under resting conditions, but reduction of MV uptake following LPS pre-treatment was associated with loss of inflammatory phenotype. Instead, MVs induced significant epithelial cell inflammation following LPS pre-treatment, when MV internalization was most significant. Using pharmacological inhibitors, we interrogated the mechanisms of MV internalization to identify which endocytic pathways and cell surface receptors are involved. We demonstrated that epithelial cells are exclusively dependent on the clathrin and caveolin dependent endocytotic pathway, whereas alveolar macrophages predominantly engulf MVs via scavenger receptors whilst, epithelial cells internalize MVs via a phosphatidylserine/integrin receptor mediated pathway (specifically alpha V beta III), which can be inhibited with phosphatidylserine-binding protein (i.e. annexin V). In summary, we have undertaken a comprehensive evaluation of MV internalization within the alveolar space. Our results demonstrate that different environmental conditions can modulate MV internalization, with inflammatory stimuli strongly enhancing epithelial cell uptake of MVs and inducing epithelial cell...
activation. Our data reveal the unique mechanisms by which alveolar macrophages and epithelial cells internalize MVs thereby elucidating how MVs exert their pathophysiological effect during lung inflammation and injury. As MVs are potential novel therapeutic targets in conditions such as ARDS, these data provide crucial insights into the dynamics of MV-target cell interactions and highlight potential avenues for researchers to modulate and inhibit their pro-inflammatory actions within the alveolar space.

Keywords: extracellular vesicles, microvesicle internalization, alveolar space, intercellular communication, microvesicle processing

INTRODUCTION

Microvesicles (MVs) are cell membrane-circumscribed extracellular particles, carrying a variety of molecular cargo, such as proteins, receptors and nucleic acids (1–3) over a distance to remote cells (1–3). They provide an alternative yet essential pathway for inter-cellular communication (2, 4) and have been implicated in the pathophysiology of various inflammatory diseases (5–8). This has led to a considerable amount of interest regarding the role of MVs in inflammatory lung diseases such as acute respiratory distress syndrome (ARDS). Indeed, we have previously demonstrated that intra-alveolar MVs, particularly alveolar macrophage-derived MVs, are potent initiators of acute lung injury (ALI), mediated by molecular cargo packaged within them (9, 10) and it is now evident that MVs are key components in the pathophysiology of lung inflammation as well as potential novel therapeutic targets (9, 11–13).

Despite the abundance of data detailing the pro-inflammatory effects of MVs, it remains unclear how MVs interact or signal with target cells in the alveolus in vivo. Several mechanisms have been postulated based largely on extra-pulmonary models of inflammation (Figure 1): 1) via direct MV-cellular interaction forming a ligand-receptor complex (14); 2) through a paracrine fashion where MVs release their cargo near target cells, which subsequently act upon membrane receptors (15); 3) by endocytosis or internalization of MVs (and their cargo) by target cells (16, 17); and 4) through fusion of MVs with target cell membranes, thereby transferring their intra-vesicular cargo (18). Whilst some of these data may be applicable to the alveolar space, it is important to note that within the unique environment of the alveolus, MVs may have prolonged or enhanced effects as the alveolus acts a semi-closed environment, protected from the rapid dilution or washout of MVs by blood flow. Therefore, the dynamics of MV communication in the alveolus are likely to differ from more commonly studied compartments such as the circulation.

To date, studies on MV interactions with alveolar cells have focused on the process of endocytosis. Alveolar macrophages, as resident professional phagocytes, internalize MVs resulting in lung inflammation (11), while a recent study has shown that insulin-like growth factor-1 (IGF-1) augments particle engulfment function of non-professional phagocytic cells such as epithelial cells, resulting in enhancement of MV uptake by

FIGURE 1 | MV interaction with target cells. There are several hypotheses how MVs may interact with their target cells. Firstly this could be as simple MV and cell contact via a receptor via ligand-receptor interaction. Secondly this could occur through a paracrine fashion with MVs releasing soluble factors at a target cell site. MVs could undergo endocytosis, where they are internalized and then subsequently either undergo lysosomal degradation or activate endosomal signaling. Finally MV could fuse with their target cells resulting in fusion of 2 initially distinct membranes and subsequent release of contents into target cell cytoplasm.
airway epithelial cells in vitro (19). However these studies have focused on individual alveolar cells, predominantly in the in vitro setting and have not considered the alveolus as a whole, where alveolar macrophages and epithelial cells lie in close proximity to each other. This could lead to competition for uptake, which would not be apparent using isolated cells in vitro. Furthermore, it remains unknown whether differential uptake mechanisms/pathways occur between these different alveolar cells or if specific cellular populations preferentially take up MVs. Since MVs are a promising therapeutic avenue in lung inflammatory diseases such as ARDS, it is crucial to address these points but also to understand how MVs interact with these different cells in the alveolar environment.

In this study, we performed a comprehensive analysis of MV communication with target intra-alveolar cells in both in vivo and in vitro models, identifying key receptors and pathways involved in MV internalization in resting as well as in inflammatory conditions. We showed that MV uptake and internalization in the alveolar space is performed primarily by alveolar macrophages rather than epithelial cells. However, following exposure to lipopolysaccharide (LPS), epithelial cell uptake is significantly increased, enhancing epithelial cell inflammation, whilst alveolar macrophage internalization of MVs is reduced. Furthermore, we demonstrated clear differences in MV uptake mechanism between these two cell types, i.e. MV uptake by alveolar macrophages is predominantly a scavenger receptor mediated process, whilst epithelial cells rely upon integrin receptors. These data provide crucial mechanistic information and delineate potential means to interrupt MV-mediated signaling in the alveolus for therapeutic purposes.

**MATERIALS AND METHODS**

**Animal Experiments**

All protocols were approved by the Ethical Review Board of Imperial College London, carried out under the authority of the UK Home Office in accordance with the Animals (Scientific Procedures) Act 1986, UK and reported in compliance with the ARRIVE guidelines. One hundred and fifty-six male C57BL/6 mice (Charles River, Margate, UK), aged 10-14 weeks were used. Mice were housed in individual ventilated cages (maximum number of 5 per cage) and exposed to 12-hour light and dark cycles. All experiments were initiated and completed during the light cycle and no unexpected adverse effects were observed in any of the treatment groups.

**In Vitro MV Production and Fluorescent Labeling**

RAW 264.7 macrophages cells (Sigma-Aldrich, UK) were washed and pre-treated with 1μg/ml of ‘Ultrapure’ lipopolysaccharide (LPS) (in vivogen Toulouse, strain: E. coli O111:B4) for 1 hour to induce inflammatory conditions as previously described (9, 10). Cells were then stimulated with 3nM of ATP disodium salt (Bio-technne, UK) to induce release of ‘pro-inflammatory’ MVs. Supernatants were collected, centrifuged to remove cells (200g 10 minutes at 4°C) and then treated with 5μM of 1,12-Dioctadecyl-3,3,3’,3’-Tetramethylindodicarbocyanine Perchlorate (DiD) (ThermoFisher Scientific, UK) in Diluent C, in dark at room temperature for 7 minutes. DiD-labeled MVs were then pelleted (20,000g for 30 min at 4°C) and washed twice to remove unbound dye. The relative fluorescence of MVs was assessed using a fluorescence plate reader (Bio-tek FLX 800; Bio-tek instruments, USA), and a standardized amount of fluorescent MVs (25,000 relative fluorescence units (RFU), which corresponds to approximately 1 x 10⁶ MVs) was then added to our in vitro or in vivo models [Supplementary Figures 1, 2 (20)]. This dose of MVs was chosen as we have previously observed up to approximately 1 x 10⁶/ml of alveolar macrophage-derived MVs in bronchoalveolar lavage fluid (BALF) samples in i.t. LPS-induced ALI in mice at 1 hour (9).

MVs were identified by flow cytometry (CyAn™ ADP flow cytometer, Beckman Coulter, UK) as events under 1μm in size (forward scatter and side scatter with a trigger threshold of 0.01 were used to elucidate a 1μm gate that was delineated using sizing beads) and positive for specific surface marker CD11b (M1/70; Biolegend, CA) and DiD (Figure 2A). MVs were also enumerated using Accucheck counting beads (Invitrogen, Paisley, UK) as previously shown (9, 10, 20). Data were analysed using FlowJo software. All stained MV samples were also treated with 0.1% triton detergent in order to correctly differentiate MVs from non-vesicular antibody-bound events (21). The centrifugation and flow cytometry methods to isolate and characterise MVs in this study have been previously validated both by high-resolution imaging and electron microscopy (10).

**In Vivo MV Production**

In vivo-derived MVs were harvested from our LPS model of ALI as previously described (composed primarily of alveolar macrophage and epithelial cell-derived MVs) (9). In brief, mice were anesthetized (intraperitoneal ketamine 90mg/kg; xylazine 10mg/kg) and 20μg LPS in 50μl was instilled intratracheally (i.t.). After 1 hour, animals were euthanized and tracheostomized, BALF was obtained by flushing and gently aspirating 700μl of 0.9% saline in and out of the lungs via the endotracheal tube three times, and centrifuged to remove cells and larger particles (200g, 10mins at 4°C). Cell free supernatants were then stained with DiD as described above. DiD-labeled in vivo generated MVs were then pelleted (20,000g for 30 min at 4°C) and washed twice to remove unbound dye and other stimulatory factors.

**In Vivo Model of Intra-Alveolar MV Uptake**

Fluorescent-labeled RAW macrophage-derived MVs (25,000 RFU, resuspended in 50μl normal saline) or post-wash supernatant (50μl of supernatant following second wash step as control) was instilled i.t. into the lungs of randomly selected mice by an investigator blinded to the treatment groups (Supplementary Figure 1). In a separate set of experiments, in vivo derived MVs (25,000 RFU, obtained as described above) were also instilled i.t. into the lungs of randomly selected mice (Supplementary Figure 2). Either 1 or 4 hours after instillation, mice were euthanized, lungs were removed and mechanically disrupted in warm fixation buffer using a GentleMACS
dissociator [Miltenyi Biotec, Surrey, UK (20)]. Samples were then passed through 40µm sieves, washed and resuspended twice in flow cytometry buffer (2% fetal calf serum, 2mM EDTA and 0.1% sodium azide constituted in PBS) to yield a fixed single cell suspension. Uptake was then evaluated by assessing mean fluorescence intensity (MFI) of DiD in alveolar cells by flow cytometry. As described previously (9), alveolar macrophages were identified as CD45+ (clone 30-F11; Biolegend), CD11c+ (clone N418; eBioscience, CA), CD11b−, F4/80+ (BM8; eBioscience). Epithelial cells were identified as CD45−, CD31− (MEC 13.3; BD Bioscience, CA), and T1alpha+ (8.1.1; Biolegend) events (9). In some experiments, mice were pre-treated with i.t. 20ng LPS (in 50µl normal saline) or saline (50 µl) for 1 hour in order to assess the effect of underlying lung inflammation on MV uptake.

In Vitro Model of Intra-Alveolar MV Uptake
In order to model the intra-alveolar environment in vitro, we created a co-culture system comprised of primary alveolar macrophages and Murine Lung Epithelial (MLE-12) cells (ATCC, UK). MLE-12 cells were seeded overnight in a 24 well-plate at a density of 105 cells/well and primary alveolar macrophages [harvested by lung lavage from untreated mice as described previously (9)] were added to MLE cells for one hour in the ratio of 1:5 (22). DiD-labeled RAW macrophage-derived MVs or post-wash supernatant were then incubated in this in vitro model of the alveolus. After 1 or 4 hours, cells were detached using EDTA containing solution (Versene, Life Technologies), stained with the fluorescently conjugated antibodies against CD45, CD11c and T1α and DiD MFI of individual cells were assessed.

FIGURE 2 | In vivo uptake of MVs. (A) Flow cytometry plot of RAW macrophage derived MVs, demonstrating that DiD labeled events were also positive for CD11b, confirming their identity as macrophage-derived MVs. (B) These particles were readily visualized by confocal microscopy as DiD (red, top left panel), particles negative for nuclear materials (DAPI-, top-right), but positive for CD11b (green, bottom-left, co-localization shown in the bottom-right combined image) (n = 3). (C) DiD labeled RAW macrophage MVs (25,000 RFU) were instilled into the trachea of untreated mice and in vivo MV uptake by alveolar macrophages and epithelial cells was assessed by flow cytometry at 1 and 4 hours. Time 0 represents baseline auto-fluorescence, i.e. no MVs added. Alveolar macrophages significantly internalized the majority of MVs compared to alveolar epithelial cells at both time points [1 hour: alveolar macrophage 734 ± 135 RFU vs. epithelial cells 4.78 ± 0.57 RFU; 4 hours: alveolar macrophage 1153 ± 243 RFU vs. epithelial cells 5.21 ± 0.47 RFU (n = 5)]. (D) Alveolar macrophages, as compared to epithelial cells, internalized the majority of MVs at 1 hour after installation in vivo regardless of the MV phenotypes used: 1) in vitro-derived RAW macrophage MVs (i.e. from stimulated RAW macrophage culture); 2) in vivo-derived MVs, non-inflamed (intra-alveolar MVs harvested by lung lavage from untreated mice); and 3) in vivo-derived MVs, inflamed (intra-alveolar MVs harvested by lung lavage from LPS-treated mice). (n = 3-6, not significant by one way ANOVA).
Mechanisms of MV Uptake
In our in vitro alveolar co-culture model, we examined the surface expression of variety of scavenger/integrin receptors in each cell type via flow cytometry: MERTK (2B10C42; biolegend), TIM4 (RMt4-54; biolegend), Alpha V Beta III (2C9.G2; Biolegend), Alpha V Beta V (RMV-7; Biolegend), Macrophage Scavenger Receptor 1 (REA1A48; Miltenyi), MARCO, CD36 (HM36; Biolegend); CD68 (FA-11; Biolegend).

In separate experiments, to assess mechanism of MVs internalization, we pre-incubated the cells in the alveolar co-culture model with 2µM Cytochalasin D or 0.25mM Dynasore and then incubated them with DiD-labeled RAW MVs for 1 hour. In separate experiments, to visualize DiD-labeled MVs via confocal microscopy, MVs were placed on poly-L-ornithine coated coverslips to encourage adherence for 6 hours. Thereafter both MVs and cells were washed, fixed, permeabilized with 0.5% triton-X 100 and incubated with 3% bovine serum albumin (Sigma-Aldrich) for 30mins. Slides were then incubated with 5µg/ml T1 alpha (ab109059; Abcam, Cambridge UK) or 5µg/ml CD45 (ab23910; Abcam, Cambridge UK) and 5µg/ml T1 alpha (ab109059; Abcam, Cambridge UK) and then incubated with 5µg/ml T1 alpha (ab109059; Abcam, Cambridge UK) overnight in the dark at 4°C, followed by washing and incubation with secondary antibodies (1:1000) for 1 hour. After washing, slides were treated with 4',6-diamidino-2-phenylindole (DAPI) and an nuclear stain (1:10000) solution (pre-made) for 10mins. Coverslips were placed on slides with Mounting PermaFluor (ThermoFisher Scientific) and viewed using a Zeiss LSM880 NLO multiphoton confocal imaging system with Axio Observer 1 microscope. The objective lens used was a Plan Apochromat 40x/1.3 oil DIC UVVIS-IR. The imaging medium was oil and the temperature -20°C. The fluorochromes used were Alexa-Fluor 488 and Alexa-Fluor 594. CZI Images were acquired using Zen software, which were then exported as 16 bit Tiff images. No image processing software was used.

Statistical Analysis
Shapiro-Wilk normality tests were carried out (IBM SPSS). Comparisons between two data sets were performed using either paired T-tests or Wilcoxon Rank Sum test. All data was analyzed on GraphPad Prism and are expressed as Mean ± SD, or Median ± Interquartile range. A p<0.05 was defined as the minimum threshold for statistical significance.

RESULTS

In Vivo Uptake of MVs
We have previously shown that ‘pro-inflammatory’ macrophage-derived MVs are rapidly released within the alveolus in vivo and involved in the early pathogenesis of ALI (9). To model this effect, RAW macrophages were stimulated with LPS (inflammatory stimulus) followed by ATP (danger signal) to induce release of pro-inflammatory MVs, which were then labeled with the fluorescence lipophilic dye DiD (17) for detection by flow cytometry and confocal microscopy. A known fluorescent quantity of these MVs [25,000 RFU which corresponds to 1 x 10^6 MVs; we previously found up to approximately 1 x 10^8/ml of alveolar macrophage-derived MVs in BALF in i.t. LPS-induced ALI in mice at 1 hour (9)] was instilled into the trachea of untreated mice, and MV uptake by different alveolar cells was assessed by flow cytometric analysis of lung single cell suspensions. One hour after instillation, we found that alveolar macrophages rather than epithelial cells internalized the majority of MVs (Figure 2C), despite several studies previously demonstrating that epithelial cells rapidly take up MVs (19, 23, 24). This difference persisted over time, such that at 4 hours alveolar macrophage still internalized the majority of MVs (1153 ± 244 RFU for alveolar macrophages vs. 5.21 ± 0.47 RFU for epithelial cells).

To assess the uptake of the mixed populations of MVs released in vivo within the alveolus, we used intra-alveolar MVs harvested from an in vivo model of ALI. These MVs contained ‘mixed’ populations, predominantly composed of alveolar macrophage- and epithelial cell-derived MVs (with concentrations of ~1000 MVs/µL and ~650 MVs/µL respectively in the original BALF samples), and have significant pro-inflammatory activity (9). We also used intra-alveolar MVs harvested from untreated mice as a control, which are composed of both alveolar macrophage- and epithelial cell-derived MVs (~300 and ~180 MVs/µL in the original BALF samples), and devoid of inflammatory activity. Both types of MVs were labelled and instilled i.t. into another mouse (Supplementary Figure 2). Alveolar macrophages still internalized the overwhelming majority of these primary, in vivo-generated ‘inflamed’ or ‘non-inflamed’ MVs compared to epithelial cells 1 hour after installation (Figure 2D). Interestingly, there was a trend that epithelial cells internalized more in vivo derived MVs compared to RAW MVs but this did not reach statistical significance (epithelial uptake of RAW MV: 5.21 ± 0.47 RFU vs. non-inflamed in vivo MVs: 16.1 ± 12.0 RFU vs. inflamed in vivo MVs: 15.9 ± 2.9 RFU). Nevertheless, these data suggest that irrespective of MV population (either alveolar macrophage- or epithelial cell-derived MVs) and phenotype (inflamed or no-inflamed), alveolar macrophages internalize the majority of MVs within the alveolar space under normal physiological conditions, within the time frame of our observations.

In Vitro Uptake of MVs
In order to model the environment with the lungs, we created an in vitro alveolar system composed of primary alveolar macrophages (obtained from untreated mice) and murine lung epithelial (MLE-12) cells (22). This co-culture system allowed us to confirm our in vivo results and explore the dynamics and mechanisms of MV-mediated communication with alveolar cells in detail. DiD labeled RAW-derived MVs were incubated within this in vitro environment for 1 or 4 hours, and the uptake of MVs by individual cells was measured by flow cytometry and confocal microscopy. As occurred in vivo, substantial amounts of MVs...
were internalized by alveolar macrophages, after both 1 and 4 hours incubation time (Figure 3A), while epithelial cells did internalize MVs but this process occurred to a much lesser extent compared to alveolar macrophages (Figure 3B). As would be expected from this in vitro alveolar model (which is essentially a completely closed system), the amount of MVs recovered in the cell culture supernatant substantially decreased over time, consistent with these MVs being taken up by alveolar cells (Figure 3C). These findings were also confirmed by confocal microscopy (Figure 3D), where alveolar macrophages can be seen to internalize MVs.

Interestingly, we found that when we increased MV numbers in our co-culture system, alveolar macrophages appeared to become saturated and unable to internalize any further MVs. Instead, epithelial cells started taking up MVs to almost a similar extent as alveolar macrophages at 4 hours of incubation, presumably as there is less competition from alveolar macrophages (Supplementary Figure 3). There may also be increased uptake efficiency of the epithelial cells at higher MV concentrations. However, it is important to note that the number of MVs required to precipitate this effect (0.5 to 1.5 x 10^7/ml) far exceeds the number of macrophage MVs measured in BALF (~1 x 10^6/ml) in our in vivo models of ALI (9).

Effects of Inflammation on MV Internalization

Next, we assessed whether MV internalization in the alveolar space would differ during inflammation. DiD-labeled RAW macrophage-derived MVs were instilled into the trachea of mice that were pre-treated with either 20ng LPS or saline. We found that LPS pre-treatment caused a significant switch in MV uptake: alveolar macrophage internalization was reduced (~<50%, p<0.01) whilst epithelial cell uptake was increased (~>150%, p<0.01) (Figure 4A). This pattern was replicated in our in vitro model when the co-culture was pretreated with 1µg/ml LPS (Figure 4B). These findings demonstrated that inflammatory conditions substantively modulate MV uptake within the alveolar space, producing a partial shift of MV uptake/internalization from alveolar macrophages to epithelial cells.

We also assessed whether MV uptake caused a change in alveolar cell phenotype or activation status. Using ICAM-1 as a surrogate marker of cell activation (25, 26), we found that alveolar macrophages adopted a pro-inflammatory phenotype after internalizing RAW macrophage-derived MVs under resting conditions (p<0.01, Figure 5A). Interestingly, when uptake was reduced following pre-treatment with 1µg/ml LPS, the pro-inflammatory effect of MVs was no longer evident (Figure 5A).
Conversely, MVs only induced significant epithelial cell activation following pre-treatment with LPS, when MV internalization was most significant (Figure 5B).

Endocytosis of MVs

We then investigated the mechanisms of MV internalization in both alveolar macrophages and epithelial cells. Firstly, MV uptake experiments were carried out at 4°C which abolished MV uptake in both alveolar macrophages and epithelial cells, indicating that an active, energy dependent process is involved (Figure 6A). We next assessed the effect of inhibitors of endocytosis on MV uptake by each of these cells. Cytochalasin (generalized inhibitor of endocytosis) reduced alveolar macrophage uptake by 59.5%, and dynasore (cell-permeable inhibitor of the clathrin and caveolin dependent endocytosis pathway) inhibited macrophage uptake by 47.9% (Figure 6A). In contrast, epithelial cell uptake was inhibited by 62.7% by cytochalasin but completely blocked by dynasore (Figure 6A). The absolute effects of dynasore on epithelial cells, but not on alveolar macrophages, suggests that epithelial cells are exclusively dependent on the clathrin and caveolin dependent endocytotic pathway, whereas alveolar macrophage uptake may involve a significant phagocytic component.

Receptor-Mediated Mechanism of MV Uptake

To identify receptor-mediated mechanisms involved in MV internalization by alveolar cells, we examined the presence of cell surface receptors that have been implicated in the uptake of endogenous and foreign particles or apoptotic cells, and thus may be involved in MV uptake. In particular, we investigated the relative expression of integrin receptors which bind to phosphatidylserine (PS) such as MerTK (27, 28), alpha V beta III (29, 30) and alpha V beta V (31), and scavenger receptors such as macrophage scavenger receptor A (32), MARCO (33), CD36 (34) and CD68 (35). We also studied the expression of the integrin TIM4 since it is expressed on the surface of macrophages and has a role in MV and exosome internalization (36, 37). We found that alveolar macrophages express MerTK and Alpha V beta III integrin receptors whereas epithelial cells only express Alpha V beta III integrin receptor (Figure 6B). In addition, we detected a variety of scavenger receptors on alveolar macrophages including MARCO, CD36 and CD68, but expression of scavenger receptors was low or undetectable on epithelial cells (Figure 6C). Based on these profiles, we hypothesized that both integrin (MerTK and alpha V beta III) and scavenger receptors (MARCO, CD36, and CD68) were involved in MV uptake by alveolar macrophages, whereas...
epithelial cell uptake occurred principally via integrin receptors (specifically alpha V beta III).

To examine this hypothesis, we pre-treated our co-culture system with either polyinosinic acid (50µg/mL) (Class A scavenger receptor inhibitor) or annexin V [binds to PS expressed on MVs preventing integrin/PS mediated MV uptake (38, 39)] and assessed their effect on MV uptake by each cells (Figures 6D, E). As predicted, annexin V inhibited uptake by both alveolar macrophages and epithelial cells but had a greater effect on uptake by epithelial cells. Conversely, scavenger receptor inhibitor polyinosinic acid dramatically reduced alveolar macrophage internalization of MVs, with a lesser effect on epithelial cell uptake. The results suggest that integrin receptors play an essential role in MV uptake by epithelial cells, whereas scavenger receptors play a more predominant role in that by alveolar macrophages.

**DISCUSSION**

Elevated production of MVs is a common feature in the pathophysiology of alveolar inflammation, particularly in conditions such as ARDS (9) or COPD (8), yet there is a paucity of data describing MV communication or cellular uptake within the alveolus. This study demonstrates several novel findings detailing MV trafficking within the unique environment of the alveolar space, under both resting and inflammatory states. We have provided convincing data that the overwhelming majority of MV uptake is performed by alveolar macrophages under resting conditions, irrespective of MV phenotype or origin and inducing a pro-inflammatory phenotype in these cells. Although epithelial cells internalized MVs only to a limited extent, this was substantially enhanced by LPS priming, leading to significant MV-induced epithelial cell activation. Furthermore, our data indicate that MVs are internalized by distinct receptor-mediated pathways: alveolar macrophages predominantly internalize MVs via scavenger receptors, whilst epithelial cells endocytose MVs through a PS/integrin receptor mediated pathway.

Within the alveolus, the major resident cell populations directly exposed to the air space are alveolar macrophages and epithelial cells. Previous studies investigating MV mediated communication within the alveolus have been performed mostly in an *in vitro* setting, concentrating on MV uptake by either epithelial cells (19, 23, 24) or alveolar macrophages (11, 28) alone. This is the first study to analyze MV internalization by different alveolar cells *in vivo*, and we have shown that the degree of uptake by each of these alveolar cells is dependent on the environmental condition to which they are exposed. Under normal physiological conditions, alveolar macrophages internalize the overwhelming majority of MVs, inducing an inflammatory phenotype, while epithelial cells...
internalize only a very modest proportion of MVs. Hence, in non-inflamed alveoli, any pro-inflammatory effects of MVs seems to be mediated largely by alveolar macrophages. However, in reality, MVs are actively produced within the alveolus when cells are exposed to injury/inflammation causing release of danger associated molecular patterns such as ATP (40, 41). Therefore, any study examining MV pro-inflammatory signaling in the alveolus should rather be conducted under...
inflammatory stress to better simulate clinically-relevant disease states. We found that during inflammation, alveolar macrophage uptake of MVs is substantively reduced, with the pro-inflammatory effect of MVs upon macrophages being no longer evident. In sharp contrast, MV uptake by alveolar epithelial cells was markedly enhanced, inducing significant epithelial cell activation. We have previously observed a similar phenomenon of uptake shift of circulating MVs such that LPS injection reduced MV uptake by liver Kupffer cells but enhanced internalization of MVs by monocytes margined within the pulmonary vasculature, leading to significant pulmonary vascular inflammation (20). Previous studies have demonstrated the reduced expression of scavenger receptors such as CD36 and MARCO on the surface of inflamed/M1-polarized macrophages, potentially explaining this finding (42, 43).

The mechanisms detailing MV communication and interaction with target cells remain poorly understood and this is particularly true within the alveolus during inflammatory lung diseases. It is thought that this interaction occurs primarily through endocytosis (3, 23), which consists of a number of pathways including macropinocytosis, phagocytosis, clathrin-dependent, caveolin-dependent, or clathrin/caveolin-independent pathways. Alveolar macrophages and epithelial cells are likely to internalize MVs via different endocytosis pathways (44, 45), and we undertook a series of experiments to identify the mechanisms and receptors involved in MV uptake by these alveolar cells. Our confocal microscopy images demonstrated that MVs are internalized by alveolar macrophages rather than fusion with cell membranes. This internalization is an active process, and cytochalasin D, an actin polymerization inhibitor and a generalized suppressor of endocytosis (38, 46), reduced but did not completely abolish MV uptake in both cell types to a similar extent as previously reported (47). Dynasore, a dynamin inhibitor which inhibits both clathrin and caveolin-dependent endocytosis (48), had a much more profound effect on epithelial cells, effectively abolishing epithelial cell uptake. This suggests that there are mechanistic differences in MV uptake between the two cell types: epithelial cell uptake is dependent upon clathrin and caveolin-dependent endocytosis whereas alveolar macrophages may be more reliant upon other endocytic mechanisms (e.g. macropinocytosis and phagocytosis), which is consistent with recent data of endocytosis mechanisms of nanoparticle uptake by alveolar cells (49). Indeed, the different speed and capacity of uptake mechanism may explain to some extent why alveolar macrophages (via phagocytosis by forming large phagosomes) take up the majority of MVs in the alveolar space compared to epithelial cells (via clathrin and caveolin-dependent endocytosis by forming much smaller plasma membrane vesicles).

Since MVs express PS on their surface and integrins have been implicated in the uptake of apoptotic material via binding of PS, we investigated the expression of various integrin receptors in these alveolar cells. We found that alpha V beta III was the only integrin expressed by murine epithelial cells whilst alveolar macrophages expressed both MERTK and alpha V beta III. Inhibition of integrin/PS binding with annexin V substantially blocked epithelial cell uptake while alveolar macrophage uptake was moderately inhibited. We also investigated the expression of scavenger receptors on both cell types, and as expected, identified several on alveolar macrophages, particularly MARCO (class A) and CD36 (class B), both of which have been shown to play a prominent role in removing apoptotic material (50, 51) and particles (32, 52, 53), but not on epithelial cells. Polyinosinic acid, a known inhibitor of scavenger receptors particularly class A (33, 54), markedly reduced MV internalization in macrophages whilst having little effect on epithelial cells. These results suggest that integrin/PS binding (specifically via alpha V beta III) plays a crucial role in MV uptake by epithelial cells, while MV uptake by alveolar macrophages are largely dependent on scavenger receptors but integrin/PS binding may also play a role.

There are some caveats to our work. We used a membrane-bound dye DiD to measure MV uptake, but this dye may affect the functional structure of MVs. However, the use of fluorescent markers is the current gold standard method of visualizing MV internalization, and these stained MVs had similar pro-inflammatory effects to un-stained MVs used in our previous studies (9). There is also a possibility that DiD may leak from stained MVs and cells, falsely conveying uptake in recipient cells. However this is unlikely since alveolar macrophages were preferentially stained with DiD, compared to epithelial cells which would be uniformly stained if DiD was truly leaking from vesicle membranes. In addition to this, uptake was inhibited by low temperature, endocytosis and receptor-mediated uptake inhibition, making significant DiD leak or artefactual uptake of precipitated DiD by cells very unlikely. Furthermore, whilst we have presented a comprehensive evaluation of MV uptake mechanisms within the alveolar space, this was based on pharmacological inhibitor studies and more specific evidence would be required to confirm the molecular pathways. For example, scavenger receptors MARCO and CD36 have been implicated in EV uptake previously (55, 56), but it was beyond the scope of the study to precisely define whether these specific scavenger receptors were responsible for initiating MV endocytosis. Further studies, potentially using MARCO or CD36 knockout mice/SiRNA knockdown cells, would be prudent to elicit if alveolar macrophages were reliant on a particular scavenger receptor during MV internalization.

On the other hand, our data have a number of strengths as a study investigating MV interaction with target cells within the lungs. Firstly, we have investigated MV uptake in the alveolus as a whole, rather than just concentrating upon individual alveolar cells. We have comprehensively characterized MV processing in the alveolar space using both in vitro and in vivo models, while previous studies have just investigated internalization using in vitro models (19, 24). Furthermore, we have used MVs from different sources, both in vitro and in vivo generated, to demonstrate that alveolar macrophages take up the majority of MVs irrespective of their phenotype or origin (19, 23, 24). Finally, we have employed robust methodologies to reliably assess MV uptake by different cells in these models, e.g. reproducible standardization of administered MV doses using fluorescence, sensitive quantification of MV uptake by individual cells using flow cytometry, morphological confirmation of MV.
internalization by confocal microscopy, and mechanistic investigations into MV uptake using combined surface marker assessment/inhibition experiments.

In conclusion, this study demonstrates, for the first time, that alveolar macrophages internalize the majority of MVs within the alveolar space under resting conditions, but during intra-alveolar inflammation, MV uptake by epithelial cells is substantially increased, leading to MV-mediated alveolar epithelial activation. Furthermore we have established that alveolar macrophages and epithelial cells internalize MVs via contrasting mechanistic pathways: alveolar macrophages predominantly engulf MVs via scavenger receptors, whilst epithelial cells internalize MVs through a PS/integrin receptor-mediated endocytosis pathway. These data elucidate crucial mechanistic information describing how MVs can produce alveolar epithelial injury, exacerbating lung inflammation and have key implications when understanding MV signaling within the alveolar space and MV interactions with target cells. As potential novel therapeutic targets, these data highlight future areas of study by which researchers can modulate the actions of MVs within the alveolar space.

DATA AVAILABILITY STATEMENT

The original contributions presented in the study are included in the article/Supplementary Material. Further inquiries can be directed to the corresponding author.

REFERENCES

1. Wang J-G, Williams JC, Davis BK, Jacobson CM, Ting JP-Y, et al. Monocytic Microparticles Activate Endothelial Cells in an IL-1β-Dependent Manner. *Blood* (2011) 118(8):2366–74. doi: 10.1182/blood-2011-01-330878

2. Lee TH, D’Asti E, Magnus N, Al-Nedawi K, Meehan B, Rak J. Microparticles as Mediators of Intercellular Communication in Cancer—the Emerging Science of Cellular ‘Debris’. *Semin Immunopathol* (2011) 33(3):455–67. doi: 10.1007/s00281-011-0250-3

3. Ismail N, Wang Y, Dakhilallah D, Moldovan L, Agarwal K, Batte K, et al. Macrophage Microparticles Induce Macrophage Differentiation and mmp-223 Secretion. *Blood* (2013) 121(6):984–95. doi: 10.1182/blood-2011-08-374793

4. Islam MN, Das SR, Emin MT, Wei M, Sun L, Westphalen K, et al. Mitochondrial Transfer From Bone-Marrow-Derived Stromal Cells to Pulmonary Alveoli Protects Against Acute Lung Injury. *Nat Med* (2012) 18(5):759–65. doi: 10.1038/nm.2736

5. Hulsmans M, Holvoet P. MicroRNA-Containing Microparticles Regulating Inflammation in Association With Atherosclerotic Disease. *Cardiovasc Res* (2013) 100(1):7–18. doi: 10.1093/cvr/cvt161

6. Kniff-Dutmer E, Koerts J, Nieuwland R, Kalsbeek-Batenburg E, Van De Laar M. Elevated Levels of Platelet Microparticles Are Associated With Disease Activity in Rheumatoid Arthritis. *Arthritis Rheumatol* (2002) 46(6):1498–503. doi: 10.1002/art.10312

7. Reid V, Webster N. Role of Microparticles in Sepsis. *Br J Anaesth* (2012) 109(4):503–13. doi: 10.1093/bja/aes321

8. Soni S, Garnier JL, O’Dea KP, Koh M, Finney L, Tirlapur N, et al. Intra-Alveolar Neutrophil-Derived Microparticles Are Associated With Disease Severity in COPD. *Am J Physiol Lung Cell Mol Physiol* (2021) 320(1):L73–83. doi: 10.1152/ajplung.00099.2020

9. Soni S, Wilson MR, O’Dea KP, Yoshida M, Katbeh U, Woods SJ, et al. Alveolar Macrophage-Derived Microparticles Mediate Acute Lung Injury. *Thorax* (2016) 71(11):1020–9. doi: 10.1136/thoraxjnl-2015-208032

10. Soni S, O’Dea KP, Tan YY, Cho K, Abe E, Romano R, et al. ATP Redirects Cytokine Trafficking and Promotes Novel Membrane TNF Signaling via Microvesicles. *FASEB J* (2019) 33(5):6442–55. doi: 10.1096/fj.201802386R

11. Lee H, Zhang D, Wu J, Otterbein LE, Jin Y. Lung Epithelial Cell–Derived Microparticles Regulate Macrophage Migration via MicroRNA-17/221–Induced Integrin β1 Recycling. *J Immunol* (2017) 199(4):1453–64. doi: 10.4049/jimmunol.1700165

12. Takahashi T, Kobayashi S, Fujino N, Suzuki T, Ota C, He M, et al. Increased Circulating Endothelial Microparticles in COPD Patients: A Potential Biomarker for COPD Exacerbation Susceptibility. *Thorax* (2012) 67(12):1067–74. doi: 10.1136/thoraxjnl-2011-201395

13. Bacha NC, Blandinieres A, Rossi E, Gendron N, Nevo N, Leoutre S, et al. Endothelial Microparticles are Associated to Pathogenesis of Idiopathic Pulmonary Fibrosis. *Stem Cell Rev Rep* (2018) 14(2):223–35. doi: 10.1007/s12015-017-9778-5

14. Beyer C, Pisetsky DS. The Role of Microparticles in the Pathogenesis of Rheumatic Diseases. *Nat Rev Rheumatol* (2010) 6(1):21–9. doi: 10.1038/nrrheum.2009.229

15. Pizzirani C, Ferrari D, Chiozzi P, Adinolfi E, Sandona D, Savaglio E, et al. Stimulation of P2 Receptors Causes Release of IL-1β–Loaded Microparticles From Human Dendritic Cells. *Blood* (2007) 109(9):3856–64. doi: 10.1182/blood-2005-06-301377

16. Dasgupta SK, Le A, Chavakis T, Rumbaut RE, Thiagajaran P, Developmental Endothelial Locus-1 (Del-1) Mediates Clearance of Platelet Microparticles by the Endothelium. *Circulation* (2012) 125(13):1664–72. doi: 10.1161/CIRCULATIONAHA.111.068333

17. Tan YY, O’Dea KP, Soni S, Shah S, Patel BV, Takata M. Enhanced Recognition and Internalisation of Microparticles by Lung-Marginated, Ly-6chigh Monocytes During Endotoxaemia. *FASEB J* (2017) 31(Supplement):327.7. doi: 10.1096/fasebj.31.1_supplement.327.7

18. Parolini I, Federici C, Raggi C, Lugini L, Palleschi S, De Milto A, et al. Microenvironmental pH Is a Key Factor for Exosome Traffic in Tumor Cells. *J Biol Chem* (2009) 284(49):34211–22. doi: 10.1074/jbc.M109.04115
19. Han CZ, Juncadella IJ, Kinchen JM, Buckley MW, Klibanov AL, Dryden K, et al. Macrophages Redirect Phagocytosis by Non-Professional Phagocytes and Influenza Infected Cells. *Nature* (2016) 539(7630):570–4. doi: 10.1038/nature20141

20. O’Dea KP, Tan YY, Shah S, V Patel B, C Tatham K, Wilson MR, et al. Monocytes Mediate Homing of Circulating Microvesicles to the Pulmonary Vasculature During Low-Grade Systemic Inflammation. *J Extracell Vesicles* (2020) 9(1):170678. doi: 10.1080/20013078.2019.170678

21. Osteikoetxea X, Sódar B, Németh A, Szabó-Taylor K, Pálóci K, Vukman KV, et al. Differential Detergent Sensitivity of Extracellular Vesicle Subpopulations. *Org Biomol Chem* (2015) 13(8):9775–82. doi: 10.1039/C5OB01451D

22. Fritz JM, Dwyer-Nield LD, Malkinson AM. Stimulation of Neoplastic Mouse Lung Cell Proliferation by Alveolar Macrophage-Derived, Insulin-Like Growth Factor-1 Can Be Blocked by Inhibiting MEK and PI3K Activation. *Mol Cell* (2011) 10(1):76. doi: 10.1186/1476-4598-10-76

23. Bourdonnay E, Zaslona Z, Penke LRK, Speth JM, Schneider DJ, Przybranowski S, et al. Transcellular Delivery of Vesicular SOCS Proteins From Macrophages to epithelial Cells Blunts Inflammatory Signaling. *J Exp Med* (2015) 212(5):729–42. doi: 10.1084/jem.20141675

24. Schneider DJ, Speth JM, Penke LR, Wettlaufer SW, Swanson JA, Peters-Golden M. Mechanisms and Modulation of Microvesicle uptake in a Model of Alveolar Macrophage Communication. *J Biol Chem* (2017) 292(51):20897–910. doi: 10.1074/jbc.M117.792416

25. Sharma AK, Fernandez LG, Awad AS, Kron IL, Laubach VE. Growth Factor-1 Can be Blocked by Inhibiting MEK and PI3K Activation. *Lung Injury Requires MerTK*. *Front Immunol* (2017) 8:1014. doi: 10.3389/fimmu.2017.01097

26. Bachurski CJ, Pryhuber GS, Glasser SW, Kelly SE, Whitsett JA. Tumor Necrosis Factor-α Inhibits Surfactant Protein C Gene Transcription. *J Biol Chem* (1995) 270(33):19402–7. doi: 10.1074/jbc.270.33.19402

27. Kazeros A, Harvey B-G, Carolan BJ, Vanni H, Krause A, Crystal RG. Overexpression of Apoptotic Cell Removal Receptor MERTK in Alveolar Macrophages of Cigarette Smokers. *Am J Respir Cell Mol Biol* (2008) 39(6):674–57. doi: 10.1165/rcmb.2007-0306OC

28. Mohning MP, Thomas SM, Barthel L, Mould KJ, McCubrey AL, Frasch SC, et al. Phagocytosis of Microparticles by Alveolar Macrophages During Acute Lung Injury Requires MerTK. *Am J Physiol Lung Cell Mol Physiol* (2007) 293(1):L105–L113. doi: 10.1152/ajplung.00470.2006

29. Hoffmann PR, Ogden CA, Leverrier Y, Bratton DL, Daleke DL, Ridley AJ, et al. Phosphatidylserine (PS) Induces PS Receptor–Mediated Macropinocytosis and Promotes Clearance of Apoptotic Cells. *J Cell Biol* (2001) 155(4):649–60. doi: 10.1083/jcb.200108080

30. Oehlin TM, Haidaris CG, Lerner NR, Simpson-Haidaris PI. Integrin αvβ3 Mediated Endocytosis of Immobilized Fibrinogen by A549 Lung Alveolar Epithelial Cells. *Am J Respir Cell Mol Biol* (2001) 24(1):12–21. doi: 10.1165/ajrccm.24.1.199229

31. Pande P, Mosleh TA, Aust AE. Role of caveolae in the phagocytosis of apoptotic cells by macrophages. *J Cell Biol* (2001) 153(5):1141. doi: 10.1083/jcb.200111141

32. Palecanda A, Paulauskis J, Al-Mutairi E, Simms G, Suzuki H, et al. Role of the Scavenger Receptor MARCO in Alveolar Macrophage Binding of Unopsonized Environmental Particles. *J Exp Med* (1999) 189(9):1497–506. doi: 10.1084/jem.189.9.1497

33. Dodd CE, Pyle CJ, Glowinski R, Rajaram MV, Schlesinger LS. CD36-Mediated Endocytosis of Immobilized Fibrinogen by A549 Lung Alveolar Epithelial (A549) Cells. *Toxicol Appl Pharmacol* (2006) 210(1–2):70–7. doi: 10.1016/j.taap.2005.07.017

34. Franza A, Aggarwal P, Barsow EV, Kozlov SV, Dobrovolskaia MA. Recruitment of ATG7 for Autophagy during Lipid Vesicle Incorporation into Lipid Bilayers. *Proc Natl Acad Sci U S A* (2008) 105(20):6953–8. doi: 10.1073/pnas.0710099105

35. Pfeiler S, Thakur M, Grünauer P, Megens RT, Joshi U, Coletti R, et al. CD36-Triggered Cell Invasion and Persistent Tissue Colonization by Tumor Microvesicles During Metastasis. *FASEB J* (2019) 33(2):1860–72. doi: 10.1096/fj.201800985R

36. Soni et al. Microvesicle-Mediated Communication Within the Alveolar Space
56. Kanno S, Hirano S, Sakamoto T, Furuyama A, Takase H, Kato H, et al. Scavenger Receptor MARCO Contributes to Cellular Internalization of Exosomes by Dynamin-Dependent Endocytosis and Macropinocytosis. Sci Rep (2020) 10(1):1–12. doi: 10.1038/s41598-020-78464-2

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