Neutrophil-targeted, protease-activated pulmonary drug delivery blocks airway and systemic inflammation

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Introduction

Intractable lung inflammation in patients with acute respiratory distress syndrome (ARDS), chronic obstructive pulmonary disease (COPD), and cystic fibrosis (CF) is characterized by sustained and non-resolving polymorphonuclear neutrophil (PMN) recruitment to the airways (1–4). Subsequent extracellular release and accumulation of PMN-derived proteases, such as neutrophil elastase (NE) and matrix metalloproteinase-9 (MMP-9), lead to the saturation of the antiprotease shield and unopposed proteolytic action, resulting in structural lung damage and decline in lung function (5, 6). Repackaging of released NE into extracellular vesicles (EVs) also leads to a pathological conformation that is inherently resistant to the lung antiprotease shield and promotes emphysema (7). The presence of these proteases in the airway milieu is linked to progression of structural lung damage (8–10). Thus, targeting of airway PMN granule and EV release is of significant therapeutic importance, with potential for broad clinical impact (11). However, to date, the standard of care relies on the prescription of antiinflammatories delivered systemically or on inhaled corticosteroids, which do not target effectively the neutrophilic component of the inflammatory response.

Pulmonary drug delivery presents a unique opportunity to target lower airway inflammation, which is often characterized by the massive recruitment of neutrophils from blood. However, specific therapies are lacking modulation of airway neutrophil function, and difficult challenges must be overcome to achieve therapeutic efficacy against pulmonary inflammation, notably drug hydrophobicity, mucociliary and macrophage-dependent clearance, and high extracellular protease burden. Here, we present a multistage, aerodynamically favorable delivery platform that uses extracellular proteolysis to its advantage to deliver nanoparticle-embedded hydrophobic drugs to neutrophils within the lower airways. Our design consists of a self-regulated nanoparticle-in-microgel system, in which microgel activation is triggered by extracellular elastase (degranulated by inflammatory neutrophils), and nanoparticles are loaded with Nexinhib20, a potent neutrophil degranulation inhibitor. Successful in vivo delivery of Nexinhib20 to the airways and into neutrophils promoted resolution of the inflammatory response by dampening neutrophil recruitment and degranulation, proinflammatory cytokine production in both airway and systemic compartments, as well as the presence of neutrophil-derived pathological extracellular vesicles in the lung fluid. Our findings showcase a new platform that overcomes challenges in pulmonary drug delivery and allows customization to match the proteolytic footprint of given diseases.
some drugs, such as Nexinhib20, a potent PMN degranulation inhibitor (15), also hampers direct delivery. Embedding of drugs into particles could therefore improve pulmonary delivery. However, efficient particle-aided drug delivery imposes paradoxical constraints on particle size: those with a geometric diameter between 1 and 5 μm are readily phagocytosed by lung macrophages, resulting in rapid clearance (16, 17), while only those smaller than 0.3 μm enable effective intracellular delivery (18, 19).

To overcome these challenges, we fabricated a nanoparticle-in-microgel (N-in-M) using a modified Michael addition during water-in-oil-emulsion method (20). The swellable microgel part of this multistage formulation provides the appropriate aerodynamic size for deep lung deposition in the polymer’s relaxed state and appropriate geometric size to avoid alveolar macrophage clearance in the swollen state. Meanwhile, a cross-linked NE-degradable peptide incorporated into the microgel backbone allows for the rapid degradation of the microgel in the presence of NE to release Nexinhib20-loaded nanoparticles. We demonstrate that these nanoparticles allowed for efficient delivery of Nexinhib20 to the airway and into PMNs, leading to substantial blockade of systemic and lung inflammation.

**Results**

*N-in-M degrades in NE-rich patient airway fluid, resulting in uptake of nanoparticles by PMNs.* Targeting PMNs in vivo, particularly in highly proteolytic microenvironments, is limited by our ability to effectively deliver drugs to these cells. The multistage N-in-M particle delivery system we designed allows for the controlled release of nanoparticles in regions of the lung with high concentration of proteolytic enzymes, specifically NE (Figure 1, A and B). The polymers used for both the nano and micro portions of the formulation were chosen because of their positive regulatory history and biodegradable properties. The swollen microgel geometric diameter is 3.9 μm (mean, 8 μm range), while encapsulated nanoparticles are under 200 nm (Supplemental Table 1 and Supplemental Figure 1; supplemental material available online with this article; https://doi.org/10.1172/jci.insight.131468DS1). Detailed physiochemical characterization of such microgels encapsulating representative polystyrene nanoparticles, and their biodistribution in naive mice, are reported elsewhere (21). To test the degradation of the multistage system cross-linked with a pulmonary disease-relevant NE-cleavable peptide, we imaged the N-in-M during exposure at 37°C to pathologically high levels of NE in CF airway supernatant (CF ASN). Within 30 minutes of exposure to NE-rich CF
ASN, 95% of the microgels had degraded (Figure 1C), which shows that our N-in-M system should quickly release nanoparticles if delivered to protease-rich airways.

Next, we assessed delivery of our N-in-M system to PMNs in vitro. This was accomplished in a PMN transmigration model (22), which uses the relevant biological milieu, i.e., CF ASN, as the apical fluid and has been shown previously to recapitulate the in vivo phenotype of PMNs found in CF patients’ airways (23). This model was also used to recapitulate PMN adaptation occurring in pediatric ARDS (24). Here, we measured uptake by airway PMNs of 60-nm fluorescent carboxylated polystyrene nanoparticles encapsulated within the NE-degradable microgels and observed that the degradation of the microgels in the presence of NE-rich CF ASN increased the percentage of PMNs positive for the nanoparticle by 5%–10% at 60 minutes, regardless of the transmigration milieu (chemoattractant control consisting of leukotriene B4 [LTB4] or CF ASN). When both PMNs’ transmigration and the N-in-M exposure occurred in the context of CF ASN, uptake of fluorescent nanoparticles by PMNs was increased (Figure 1D). Together, these data indicate that for efficient uptake of nanoparticles by airway PMNs, the NE-responsive microgel must degrade first. The additive effect between the acquired PMN phenotype, imprinted by the transmigration condition, and the incubation milieu with the N-in-M system suggests that targeting of airway PMNs by the N-in-M system would be more efficient in pathological settings (CF ASN) rather than during the normal course of inflammation (LTB4).

N-in-M administered to an acute neutrophilic inflammation model degrades, delivering representative fluorescent nanoparticles to airway PMNs. Next, we tested the efficacy of the N-in-M multistage particle system in an in vivo inflammation model characterized by high levels of extracellular NE (25). The N-in-M system consisting of microgels labeled with the near IR dye DyLight 650 loaded with representative blue (350/440) FluoSpheres nanoparticles was delivered intratracheally to mice treated with saline or LPS. The radiant efficiency in the excised lungs, corresponding to the fluorescence intensity of the microgel dye at 1, 6, and 22 hours after N-in-M delivery (3, 8, and 24 hours after LPS), showed a significant reduction at 6 hours after N-in-M delivery (Figure 2A), revealing a faster microgel degradation in mice treated with LPS than in saline-treated controls.

Concurrent with microgel degradation, we observed a reduction over time of blood PMN activation and PMN recruitment to the airways and subsequent release of NE-rich granules (reflected by surface CD63) and amount of soluble NE in the bronchoalveolar lavage (BAL) (Figure 2, B and C).

Figure 2. N-in-M in an acute neutrophilic inflammation model. (A) Representative in vitro imaging system (IVIS) images of excised mouse lungs with corresponding measures of radiant efficiency for 1, 6, and 22 hours for saline- and LPS-treated mice. (B) Increase in BAL NE levels for the LPS-treated mice. (C) Increased CD63 expression on BAL PMNs. (D) Percentage of BAL PMNs positive for nanoparticle only, N-in-M, and microgel only or negative for both. Box plots show median, interquartile range (IQR), and min to max. Statistical tests: Shapiro-Wilk followed by 2-way ANOVA (A) and Kruskal-Wallis (B and C). *P < 0.05, and ***P < 0.001 within time points indicated. n = 5 per group.
Thus, the short-term clearance of microgels during LPS treatment is preceded by and dependent upon the degradation of the NE-sensitive peptides cross-linking it. Notably, the population of BAL PMNs double positive for the microgel and nanoparticles showed increased surface CD63 expression, suggesting that fusion to the plasma membrane of NE-rich granules and release of their content in the extracellular milieu is concomitant with increased uptake of soluble material, as shown before (22). Last, analysis of the distribution of microgel and nanoparticle uptake showed that airway PMNs at 1 and 6 hours after delivery were primarily positive for nanoparticles alone, suggesting that microgels were degraded, allowing for nanoparticle endocytosis (Figure 2D). These findings showcase the efficacy of the N-in-M system for nanoparticle delivery to airway PMNs in a protease-rich environment in vivo.

N-in-M delivery of Nexinhib20 attenuates neutrophilic inflammation in vivo. To determine whether the N-in-M system could not only effectively deliver nanoparticles to PMNs but also modulate PMN-driven inflammation in vivo, Nexinhib20- and DiR-carrying PLGA nanoparticles were loaded into the NE-degradable microgels (N-in-M/Nex), with DiR, a near IR fluorescent dye serving as a tracer for nanoparticle uptake. To assess whether the N-in-M system with PLGA influenced the immune response, additional control groups for both saline- and LPS-treated mice were treated with vehicle DiR-only PLGA nanoparticles inside the NE-degradable microgels (N-in-M/Veh).

Although there was no detectable sign of neutrophilia or neutropenia in blood (Figure 3A), N-in-M/Nex reduced the percentage (Figure 3, B–E, and Supplemental Figure 2) and mean total number of PMNs in the lung lumen and lung parenchyma compared with the LPS-treated control group, by 13- and 4.3-fold, respectively. These results, in addition to the detection of DiR dye in airway PMNs (Figure 3F), suggests successful delivery of the drug to PMNs by the N-in-M system. Next, we investigated whether the reduction in PMN in the airways was reflective of a decrease in inflammatory poise. Analysis of critical mediator profiling in plasma and BAL confirmed the reduction in inflammatory signaling induced by treatment with N-in-M/Nex, both locally and systemically (Supplemental Figure 4).
To determine whether N-in-M/Nex could influence not only PMN numbers, but also their phenotype and function, active granule release was assessed by ELISA and flow cytometry. Despite similar PMN recruitment in LPS-alone controls, in N-in-M/Veh-alone controls, and in N-in-M/Veh LPS-treated mice, MMP-9 levels in the latter showed a 28-fold increase compared with the saline treatment group and a 5-fold increase compared with the LPS-alone group (Figure 4A). Delivery of Nexinhib20 abrogated this response in N-in-M/Nex mice, supporting the effectiveness of this potentially novel delivery system at not only counteracting LPS-induced inflammation but also dampening potential inflammatory responses induced by the delivery of the N-in-M system itself, likely caused by the detection of foreign particles or by delivery-associated tissue trauma. Most strikingly, concomitant with increased nanoparticle uptake by PMNs, surface CD63 expression on BAL PMNs (Figure 4B), reflective of NE release, was significantly lower for the Nexinhib20 formulation than all the other treatment groups. Likewise, N-in-M/Nex treatment lowered the amount of extracellular NE (Figure 4C) and presence of NE-rich EVs (Figure 4D), a pathological NE conformation we previously showed to be resistant to the lung antiprotease shield and thereby inducing emphysema (7). Last, the successful reduction of damaging enzymes in the extracellular compartment, with the dampening of the inflammatory response upon delivery of the N-in-M/Nex system, was reflected in the health of the lung tissue and reduction in inflammation-induced capillary leak, as reflected by IgM levels in the BAL fluid (Figure 4E).

Figure 4. Delivery of Nexinhib20 reduces PMN degranulation in the lung. (A) Treatment with Nexinhib20 reduced active release by BAL neutrophils of soluble MMP-9. Primary granule release, measured by surface CD63 (B), soluble NE (C), and EV-bound NE measured in the BAL (D), also showed that significant increases by LPS and control PLGA N-in-M were abrogated by Nexinhib20 delivery. (E) Reduction of the overall inflammatory response upon delivery of Nexinhib20 led to reduced capillary leakage measured by IgM levels in the BAL fluid. Saline was administered to the control group. Box plots show median, IQR, and min to max. Statistical tests: Shapiro-Wilk followed by 1-way ANOVA. *P < 0.05, **P < 0.01, ***P < 0.001, and ****P < 0.0001 between groups indicated. n = 5 per group.
Discussion

Together, our findings establish a potentially novel and specific delivery method for hydrophobic drug administration to airway PMNs. The protease-sensitive design of the N-in-M system not only allows the microgel to provide the needed aerodynamic size for deposition and geometric size to avoid rapid clearance (<1 hour), but it also exploits the high proteolytic microenvironment for controlled release of drug-loaded nanoparticles. Furthermore, encapsulation of the drug in the nanoparticle shields it and preserves its therapeutic potential. Although the system presented here was designed to be effective in the presence of NE, it presents ample opportunities for customization because the protease-sensitive peptides embedded in the microgel can be tailored to a plethora of cleavage sites specific to other enzymes. Moreover, the N-in-M system can be made to accommodate virtually any drug. Here, we chose to include Nexinhib20 because its main target is actually exocytosis and downstream NE release. Therefore, we accomplished efficient NE-dependent delivery of a drug targeting the machinery responsible for the release of NE itself, which led to the resolution of inflammation, eventually preventing further degradation of new microgels (i.e., self-limiting antiinflammatory therapy, as illustrated in the graphical abstract).

In addition to mucociliary clearance, another major challenge for inhaled drug delivery is that, because of progressive airway damage and heterogeneous worsening of pulmonary ventilation and perfusion matching, it is difficult to have adequate deposition of the drug into diseased areas of the lung (26). This remains a major problem for therapeutic development and has limited the effectiveness and safety of inhaled medications in chronic lung disease. Although we anticipate some limitation in delivery because of disease pathophysiology, our disease-targeted N-in-M system takes advantage of protease-specific drug release, which provides the ability to direct the active pharmaceutical ingredient primarily to areas of active inflammation and remodeling.

Another critical point in inhaled therapeutic development is the ability to tune the clinical response to provide benefit without incurring additional pathology or side effects (27). When using antiinflammatory agents, it is imperative to recognize the often narrow therapeutic windows, which may reduce healthy host responses, such as release of antimicrobial enzymes and neutrophil extracellular traps, while targeting progressive tissue damage. A benefit of the protease-responsive N-in-M system is that the release of the drug is self-limited by the loss of excessive protease activity. This approach allows for localized and measured delivery of antiinflammatory drugs to the diseased tissue, allowing for enhanced safety compared with nontargeted delivery of antiinflammatory agents to the whole lung.

Last, these results highlight the importance of appropriate drug selection when using foreign particles for therapeutic delivery. In this study, the addition of the empty N-in-M demonstrated some increase in inflammatory metrics (i.e., cell count, permeability), which was further exacerbated in the inflamed lung (LPS treatment). This is a key challenge in pulmonary delivery to diseased lungs. However, the delivery of the appropriate drug (Nexinhib20) in the N-in-M system completely reshaped the inflammatory landscape to the levels of the saline-treated group, confirming the effectiveness and safety of this system.

In conclusion, our findings establish the potential of N-in-M delivery systems to effectively target inflammation in pulmonary disorders characterized by unrelenting PMN influx and activation. Similar N-in-M systems may provide a relevant mechanism to strategically deliver a variety of therapeutics not only to the lung but also to other organs with high proteolytic burden (e.g., inflamed gut, joints, and tumors).

Methods

Nanoparticle and N-in-M fabrication. Therapeutics and/or dye were encapsulated in nanoparticles during fabrication using a single emulsion solvent evaporation method. Then, 100-nm Blue FluoSpheres (Thermo Fisher Scientific) or lyophilized PLGA nanoparticles were resuspended and encapsulated in 20% w/v elastase-responsive microgels using a Michael addition during water-in-oil emulsion. See details in Supplemental Methods.

In vitro transmigration experiments. Blood PMNs were isolated using Polymorphprep (Alere Technologies) as per the manufacturer’s protocol, washed, and then loaded on the transmigration chamber; transmigration and following incubations were performed using 100 nM of LTB4 (MilliporeSigma) in RPMI or CF ASN, prepared as previously published (22). The uptake of fluorescent microgels was then assessed 10 hours after transmigration by incubating PMNs recruited apically with microgels at a 1:10 ratio (PMN/microgel) in CF ASN or RPMI for 30 and 60 minutes and quantified using flow cytometry.

Flow cytometry. Cells from in vitro and in vivo assays were analyzed by multiparametric flow cytometry, as previously detailed (28), and gated as shown in Supplemental Figure 5. See Supplemental Methods for details and Supplemental Table 2 for antibodies’ characteristics.
**EV analysis.** EVs were isolated from the BAL of mice using bead-based pull-down with annexin V, as per the manufacturer’s protocol (SBI). Isolated EVs were stained with the EV marker ExoFITC (SBI) and anti-mouse NE (R&D Systems). EVs were analyzed by flow cytometry on a Cytoflex S (Beckman Coulter).

**Fluid assays.** NE and MMP-9 concentrations were measured by ELISA in mouse BAL fluid per the manufacturer’s protocol (R&D Systems). Inflammatory mediators in plasma and BAL fluid were measured using a multiplex chemoluminescent platform per the manufacturer’s protocol (MesoScale Diagnostic, V-PLEX mouse cytokines, 19 analytes). BAL fluid IgM concentration was measured by ELISA per the manufacturer’s protocol (Immunology Consultants Laboratory).

**Microscopy.** N-in-Ms were imaged on a PerkinElmer UltraVIEW VoX spinning disk confocal microscope with a Hamamatsu C9100-23b back-thinned EM-CCD camera and Nikon 100× NA-1.45 oil objective; lung sections were imaged with the Nikon 40× NA-1.3 oil objective. Microgel size was measured in the relaxed state using Volocity (PerkinElmer Inc.) and in the swollen state using MATLAB ‘‘imfindcircles’’; this does cap the lower size limit measured. Degradation was measured by kinetic imaging every minute using a BioTek Lionheart at 37°C using cell traps (Lu lab at the Georgia Institute of Technology) to hold the microgels in place (29).

**Animal studies.** Mice were treated with either saline or 100 μg of LPS. Then for the fluorescent uptake study, the N-in-M formulation was delivered 2 hours later and for the Nexinhib20 study 1 hour after injection. The time for N-in-M injection was shortened because at 2 hours after LPS mice had significant difficulty in aspirating the particle formulation. Mice used for analysis with an IVIS were given a low-alphafalfa feed diet to reduce fluorescence background. Excised lobes were imaged on an IVIS and analyzed by LivingImage software (Xenogen). Further analysis of excised lobes by immunofluorescent staining was performed per established protocols, as detailed in the Supplemental Methods.

**Statistics.** Data were analyzed in Prism (GraphPad). The Shapiro-Wilk test was used to assess normality (α = 0.05). For parametric data, we used 2-sided 1-way ANOVA and 2-way ANOVA (α = 0.05) with post hoc Tukey’s test with adjusted P value for multiple comparisons. For nonparametric data, Kruskal-Wallis (α = 0.05) with Dunn’s multiple-comparisons test with adjusted P value was used. The P value results of the statistical tests were compared with the α level. If the P value was less than the α level, then the null hypothesis was rejected, and the result was statistically significant.

**Study approval.** Blood was collected from healthy donors after informed consent was obtained according to an approved Emory University Institutional Review Board protocol. Animal studies were performed according to protocols approved by the IACUC at the University of Alabama Birmingham. Five-week-old female C57BL/6 littermates (Jackson Laboratories) were maintained in a pathogen-free facility and given sterile water and food ad libitum.

**Author contributions**
JCM, OAF, CM, AG, RT, and KR conceived the studies. JCM, OAF, CM, and XX conducted primary experiments and performed data analyses. DAFR, LV, and JL provided critical support to experiments. JCM, OAF, CM, AG, RT, and KR wrote the manuscript.

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