Bispecific antibody targets multiple *Pseudomonas aeruginosa* evasion mechanisms in the lung vasculature

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*Pseudomonas aeruginosa* is a major cause of severe infections that lead to bacteremia and high patient mortality.

*P. aeruginosa* has evolved numerous evasion and subversion mechanisms that work in concert to overcome immune recognition and effector functions in hospitalized and immunosuppressed individuals. Here, we have used multilaser spinning-disk intravital microscopy to monitor the blood-borne stage in a murine bacteremic model of *P. aeruginosa* infection.

*P. aeruginosa* adhered avidly to lung vasculature, where patrolling neutrophils and other immune cells were virtually blind to the pathogen’s presence. This cloaking phenomenon was attributed to expression of Psl exopolysaccharide. Although an anti-Psl mAb activated complement and enhanced neutrophil recognition of *P. aeruginosa*, neutrophil-mediated clearance of the pathogen was suboptimal owing to a second subversion mechanism, namely the type 3 secretion (T3S) injectisome. Indeed, T3S prevented phagosome acidification and resisted killing inside these compartments. Antibody-mediated inhibition of the T3S protein PcrV did not enhance bacterial phagocytosis but did enhance killing of the few bacteria ingested by neutrophils. A bispecific mAb targeting both Psl and PcrV enhanced neutrophil uptake of *P. aeruginosa* and also greatly increased inhibition of T3S function, allowing for phagosome acidification and bacterial killing. These data highlight the need to block multiple evasion and subversion mechanisms in tandem to kill *P. aeruginosa*.

Introduction

*Pseudomonas aeruginosa* is an important opportunistic bacterial pathogen that causes life-threatening infections in critically ill and immunocompromised individuals. It is one of the most common causes of Gram-negative pneumonia in ventilated patients, is the principle pathogen responsible for persistent/chronic infections in the cystic fibrosis lung, and is a frequent colonizer of burn wounds, catheters, and medical devices (1). However, *P. aeruginosa* causes most deaths when it enters the bloodstream and disseminates. Specifically in the case of hospital-acquired pneumonia, *P. aeruginosa* has previously been reported to disrupt the lung barrier and enter into the bloodstream causing bacteremia, resulting in poor patient prognosis (2, 3). Further complicating these infections is the fact that *P. aeruginosa* is often multi-drug-resistant, likely contributing to its advanced innate resistance mechanisms. *P. aeruginosa* resistance has been reported for nearly all commonly used antibiotics (4–6). How *P. aeruginosa* avoids the innate immune system to survive in the bloodstream and disseminate to various organs is not well understood. In vitro, *P. aeruginosa* has the capacity to evade complement deposition, prevent neutrophil recruitment, and even have cytolytic effects on immune cells upon phagocytosis (7). However, which of these evasion or subversion mechanisms might work in vivo remains unknown. Visualization of the behavior of the pathogen and how it evades immunity when it gains access to the very dynamic systemic circulation has not yet been accomplished. Candidates contributing to successful subversion and/or evasion of the immune system in vitro include the type 3 secretion (T3S) injectisome and the unique Psl exopolysaccharide, respectively (8). The *P. aeruginosa* T3S injectisome is a major determinant of virulence, and its expression is frequently associated with acute invasive infections that have been linked to increased mortality in infected patients (9). The needle-like injectisome apparatus permits translocation of effector proteins from the bacterium into the host cell. To date, only 4 T3S effectors have been functionally characterized for *P. aeruginosa* (ExoY, ExoS, ExoT, ExoU), which is far fewer compared with other well-characterized T3S injectisomes (e.g., *Salmonella enterica* SPI-1 has 13; *Shigella* species have 25; ref. 9), potentially making *P. aeruginosa* more amenable for targeted therapeutic intervention. However, T3S expression on the bacterial surface is highly regulated and may only occur at certain stages during infection, making it difficult to inhibit this crucial virulence factor of *P. aeruginosa* (10). There have been some in vitro studies showing that T3S injectisome is expressed upon host cell contact, in low-calcium environments, and in acidic pH environments (10, 11). The T3S injectisome has been shown to aid *P. aeruginosa* colonization of the host in a variety of ways, including systemic spread of the pathogen and the sequestration of *P. aeruginosa* in nonacidic vacuoles in corneal epithelial cells (12–14). Psl exopolysaccharide, on the other hand, is a serotype-independent and abundantly expressed extracellular sugar polymer implicated in *P. aeruginosa* biofilm formation (15). Psl has been implicated in preventing complement deposition, which would prevent macro-
phage and neutrophil recognition of the bacteria (15, 16). Indeed, *P. aeruginosa* Psl prevented phagocytosis and oxidant production in neutrophils (16). However, to date, all of these studies have been performed in vitro settings. Therefore we sought to see the subversion or evasion of immune cells by *P. aeruginosa* in an in vivo model. We aimed to further understand the spatiotemporal mechanism of *P. aeruginosa* pathogenesis and delineate how we can use antibody-based therapeutics to optimize *P. aeruginosa* eradication.

In this study, we aimed to understand the neutrophil-*P. aeruginosa* axis in vivo during a bacteremic model of infection using multilaser spinning-disk intravitral microscopy to track the pathogen in the very dynamic bloodstream. We specifically looked at the blood-borne stage of infection because, regardless of the mode of entry of the pathogen, through the lung via pneumonia, through skin burns, or directly into the blood via a contaminant catheter, the majority of patients die from bacteremic sepsis (2). We also examined isogenic knockouts of Psl exopolysaccharide and PcrV component of the T3S injectisome. PcrV is necessary for T3S translocation; therefore mutant bacteria fail to have functional T3S and cannot deliver important toxins (10). *P. aeruginosa* was observed to be adequately eradicated from blood via extravascular macrophages in the liver (Kupffer cells). However, in stark contrast, the lung vasculature, which lacks extravascular macrophages, functioned as a safe haven and was subject to invasion by this pathogen via direct binding to the endothelium lining the blood vessels. Despite the presence of extravascular neutrophils patrolling the lung vasculature, these immune cells failed to detect and phagocytize *P. aeruginosa* binding to the endothelium, an event not seen with *Staphylococcus aureus*, *Streptococcus pneumoniae*, or *Escherichia coli* strains. While knockout of Psl exopolysaccharide expression greatly increased neutrophil-pathogen interactions through a complement-dependent mechanism, suboptimal extravascular bacterial killing activity was observed. Disruption of the T3S injectisome did not improve neutrophil capture of bacteria, but it significantly enhanced intracellular killing of *P. aeruginosa* in the few neutrophils that did take up the bacteria. Intriguingly, the bispecific mAb MEDI3902 (currently in phase IIb clinical testing; ref. 17), targeting PcrV and Psl, bound the surface of bacteria, enhancing neutrophil capture of the pathogen and thereby giving the antibody greater access inside the phagosome for subsequent inhibition of toxin injection, markedly improving eradication of *P. aeruginosa*. MEDI3902’s protective function was enhanced more than with administration of individual parenteral anti-PcrV and anti-Psl mAbs in combination. These data suggest that a multispecific sequential approach to disarm important coordinated bacterial virulence mechanisms is necessary for efficient immune cell clearance of *P. aeruginosa*. We suggest that making use of mAb binding to abundant Psl exopolysaccharide is advantageous to deliver more antibody intracellularly, thereby more efficiently inhibiting extravascular events mediated by the T3S injectisome.

**Results**

*The liver Kupffer cells catch and phagocytize P. aeruginosa optimally. Once P. aeruginosa has entered the bloodstream, 2 of the major organs that are involved in systemic bacterial infections are the liver and lung (18). We hypothesized that P. aeruginosa localization to the liver occurs because this organ has a resident population of extravascular macrophages known as Kupffer cells that reside in the sinusoids and are able to catch bacteria out of the mainstream of blood. Following i.v. infection of mice with *P. aeruginosa* that stably expresses GFP (*P. aeruginosa*-GFP), we observed that these Kupffer cells initially catch or capture very fast-moving bacteria (under high shear conditions) out of the mainstream of blood, an unnecessary step in vitro under static conditions but absolutely critical in vivo in rapidly flowing blood (19). After capture, the Kupffer cells then rapidly phagocytize *P. aeruginosa*-GFP (Supplemental Video 1; supplemental material available online with this article; https://doi.org/10.1172/JCI89652DS1). *P. aeruginosa* were selectively captured by Kupffer cells but did not bind to the sinusoidal endothelium or other cells (Supplemental Video 1). Within 3 hours after infection the GFP signal from *P. aeruginosa* was reduced by more than 50% of values seen at 15 minutes in the liver, and fewer Kupffer cells could be seen with bacteria in them (Figure 1A). The dissipation of the *P. aeruginosa*-GFP signal as noted by fewer Kupffer cells with bacteria at 3–4 hours (Figure 1A) suggests that the majority, but not all, of the bacteria are effectively killed in the liver. We previously observed that *S. aureus*-GFP actually replicated inside Kupffer cells and its fluorescent signal became much stronger as the colonies grew for 8–24 hours (20), whereas *P. aeruginosa* did not form larger colonies. Both Psl and PcrV systems have been implicated in potential immune evasion by *P. aeruginosa* (14, 16). We next used isogenic mutant strains lacking Psl expression (ΔpslA) and PcrV mutants lacking a functional T3S injectisome (ΔpcrV) to understand the role of these bacterial components during infection in vivo. Following i.v. infection of mice with WT-, ΔpslA-, or ΔpcrV-GFP strains, we found that all 3 strains were captured out of the mainstream of blood and phagocytized equally well by Kupffer cells in the liver (Figure 1, B and C). However, while the WT and ΔpslA strains were cleared equally well as noted by the dissipation of the GFP signal within Kupffer cells at 3–4 hours, the rate of Kupffer cell clearance for the ΔpcrV strain appeared to be much faster (Supplemental Figure 1A). At the same time point the number of Kupffer cells containing *P. aeruginosa*-GFP ΔpcrV was reduced by more than half compared with ΔpslA or WT strains (Supplemental Figure 1A). To confirm these results, livers from mice infected with the 3 strains were harvested, and CFU of *P. aeruginosa* revealed the greatest killing in the ΔpcrV strain, indicated by low CFU (Supplemental Figure 1B). These results are consistent with previous reports indicating that T3S-deficient strains are avirulent in preclinical animal infection models and are rapidly cleared by the host innate immune system (21). Prophylactic immunization with anti-Psl and anti-PcrV mAbs did not improve the initial capture of WT *P. aeruginosa* by liver Kupffer cells (Figure 1D). Liver bacterial burden of *P. aeruginosa* at 6 hours reflected what was observed in animals infected with mutant bacteria, where the antibodies against PcrV but not Psl alone expedited eradication (Supplemental Figure 2).*
These data suggest that unlike other pathogens, *P. aeruginosa* may be able to escape neutrophil recognition in the lung vasculature. Given the known presence of common pathogen-associated molecular patterns on the surface of *P. aeruginosa*, it was surprising to see a lack of neutrophil recognition after infection. As such, *P. aeruginosa* must employ a cloaking mechanism. The Psl exopolysaccharide has been previously postulated to restrict complement deposition onto bacterial membranes in vitro (16). Accordingly, we found that the Δ*pslA*, but not the Δ*pcrV*, bacteria were readily recognized and phagocytized by lung intravascular neutrophils with increased efficiency compared with WT *P. aeruginosa* (Figure 2C). These data were confirmed by ex vivo quantification of *P. aeruginosa*-GFP–positive neutrophils by flow cytometry (Figure 2D). To determine whether complement was responsible for increased uptake of the *psl* mutant by neutrophils, animals deficient in complement (data not shown). Using intravital microscopy, we observed a resident population of neutrophils as they moved through the lung vasculature, tethering to the endothelium and then letting go or briefly adhering or crawling in the pulmonary capillaries (Supplemental Video 2). Upon i.v. infection of animals with *P. aeruginosa*, the bacteria firmly adhered to the lung capillaries often in close proximity to the intravascular neutrophils (Figure 2A). However, this influx of *P. aeruginosa* failed to be recognized by the neutrophils, and it was extremely rare to see neutrophils interact with and phagocytose the adherent bacteria (Figure 2A and Supplemental Video 3). This was surprising and unexpected, as intravascular infection of animals with other bacteria such as *S. aureus*, *S. pneumoniae*, or *E. coli* resulted in similar adhesion of the bacteria to the pulmonary capillaries and immediate engulfment of bacteria by the neutrophils (Figure 2, A and B, and Supplemental Video 4).
come the evasion mechanisms of *P. aeruginosa* in the lung using therapeutic antibodies, mice were pretreated with mAbs targeting the Psl exopolysaccharide (anti-Psl) or the T3S injectisome PcrV component (anti-PcrV). In animals treated with the anti-Psl mAb, neutrophils recognized *P. aeruginosa* 3- to 5-fold above control IgG as indicated by neutrophils with internalized bacteria (Figure 3, A and B, and Supplemental Videos 5–7). These results are consistent with the increased neutrophil engagement and phagocytosis observed in experiments using Psl-deficient (ΔpslA) *P. aeruginosa* complement component 3 (C3−/−) were next examined. In these mice, neither WT, ΔpslA, nor ΔpcrV *P. aeruginosa* strains were recognized by neutrophils (Figure 2E). These data suggest that Psl prevents complement activation or deposition on the surface of WT *P. aeruginosa* and this is critical for this bacterium to escape recognition by the intravascular lung resident neutrophil population.

Anti-Psl antibodies enable acute recognition of *P. aeruginosa* by neutrophils, while anti-PcrV anticytotoxic activity allows acidification of endosomal compartments. To determine whether we could overcome the evasion mechanisms of *P. aeruginosa* in the lung using therapeutic antibodies, mice were pretreated with mAbs targeting the Psl exopolysaccharide (anti-Psl) or the T3S injectisome PcrV component (anti-PcrV). In animals treated with the anti-Psl mAb, neutrophils recognized *P. aeruginosa* 3- to 5-fold above control IgG as indicated by neutrophils with internalized bacteria (Figure 3, A and B, and Supplemental Videos 5–7). These results are consistent with the increased neutrophil engagement and phagocytosis observed in experiments using Psl-deficient (ΔpslA) *P. aeruginosa*...
PcrV was important for preventing incorporation of *P. aeruginosa* into acidic vacuoles in corneal epithelial cells (14). We therefore sought to determine whether blocking PcrV function would result in increased localization of ingested *P. aeruginosa* into acidified vacuoles in neutrophils. To evaluate this hypothesis, *P. aeruginosa*–GFP was stained with a pH-sensitive dye (pHrodo Red) that only fluoresces in acidic compartments (pH lower than 6). The very few *P. aeruginosa* that were internalized in neutrophils receiving isotype control antibodies were not in an acidic environment (Figure 2, C and D). Similar to PcrV-deficient *P. aeruginosa*, a subtle increase in *P. aeruginosa* internalization (also not significant) was observed in mice given the anti-PcrV antibody targeting the T3S injectisome (Figure 2, C and D, and Figure 3, A and B).

While no role was noted for T3S in the evasion of neutrophil recognition of *P. aeruginosa*, this may be due to the fact that T3S expression is largely triggered upon contact with host cells or perhaps after phagocytosis and incorporation into a more acidic environment (11). In fact a recent publication observed that PcrV was important for preventing incorporation of *P. aeruginosa* into acidic vacuoles in corneal epithelial cells (14). We therefore sought to determine whether blocking PcrV function would result in increased localization of ingested *P. aeruginosa* into acidified vacuoles in neutrophils. To evaluate this hypothesis, *P. aeruginosa–GFP* was stained with a pH-sensitive dye (pHrodo Red) that only fluoresces in acidic compartments (pH lower than 6). The very few *P. aeruginosa* that were internalized in neutrophils receiving isotype control antibodies were not in an acidic environment when...
assessed by intravitral microscopy (Figure 3, B–D). Moreover, neutrophils from anti-Psl mAb–treated mice caught more bacteria but, again, were not generally observed in a low-pH environment (Figure 3, B–D). In contrast, neutrophils from anti-PcrV–treated mice caught fewer bacteria; however, most of these internalized bacteria were contained within acidic compartments (Figure 3, B–D). These antibody data were similar to the results seen with the mutant strains of P. aeruginosa; more effective neutrophil processing was observed with the ΔpcrV P. aeruginosa than with the WT and Δpsl P. aeruginosa within neutrophil acidified phagosomes (Figure 3E).

Coordinated targeting of both Psl and PcrV optimizes P. aeruginosa killing. Targeting Psl with therapeutic antibodies led to increased phagocytosis but did not lead to more efficient processing of P. aeruginosa, while targeting PcrV allowed for optimal processing but yielded suboptimal phagocytosis. We therefore tested a recently described bispecific mAb targeting both Psl and PcrV, which was shown to exhibit synergistic protective activity in an acute pneumonia model (24). Dose-response studies clearly revealed more efficacy of this antibody versus anti-Psl and/or anti-PcrV on an equimolar basis in an acute pneumonia model, but how this antibody functioned remained poorly understood (24). We hypothesized that the dual inhibitory capacity would bind the abundant Psl molecule on extracellular P. aeruginosa, but upon host cell contact and/or internalization of the bacteria, which leads to expression of PcrV, the anti-PcrV motif of the antibody would inhibit the T3S injectisome and allow for proper processing of bacteria. After infection of mice, we observed a striking 8- to 10-fold increase in neutrophils with captured bacteria in the lungs of animals that received MEDI3902 versus mice that received the IgG control antibody before infection (Figure 4, A and B). These data were confirmed by ex vivo quantification of P. aeruginosa-GFP–positive neutrophils by flow cytometry from mice treated with anti-PcrV, anti-Psl, and MEDI3902 (Figure 4C). MEDI3902 resulted in the greatest amount of bacterial uptake by neutrophils in the lungs. In addition, we used 3D imaging to confirm that all bacteria were internalized rather than just attached to the surface of neutrophils (Supplemental Video 8). This enhanced recognition of P. aeruginosa was specific to the engineered MEDI3902 antibody, as a combination of parenteral anti-PcrV and anti-Psl mAbs, given to mice at molar equivalent dosages, did not cause antibody data were similar to the results seen with the mutant strains of P. aeruginosa (Figure 2, D and E).

C3 and Fcγ receptors are required for optimal anti-Psl- and MEDI3902-mediated neutrophil recognition of P. aeruginosa. We identified that C3 was important in mediating neutrophil recognition of Psl-deficient P. aeruginosa (Figure 2, D and E). To provide further insight into how MEDI3902 was functioning, the antibody was tested in both C3−/− and FcγR-deficient (Fcer1γ−/−) mice. Neutrophils of mAb-treated C3-deficient mice no longer showed enhanced phagocytosis of P. aeruginosa in any of the therapeutic mAb treatment groups compared with control IgG (Figure 5A; also compare with Figure 4C). Importantly, the same number of bacteria attached to the endothelium in both C3-deficient and WT mice (Figure 5B). mAb-mediated neutrophil phagocytosis in the lung vasculature of FcγR-deficient mice was also reduced (Figure 5C). These data indicate that both Fc receptors and C3 were essential for uptake of antibody-bound P. aeruginosa by lung neutrophils.

MEDI3902 specifically acts on pulmonary neutrophils to increase recognition of P. aeruginosa, resulting in decreased bacterial burden and increased survival of mice. We next assessed bacterial organ burden in the lungs of intravenously infected mice that were treated with the therapeutic antibodies. We observed a significant reduction in bacterial burden in lungs 6 hours after infection in anti-PcrV–treated animals compared with the anti-Psl– and control mAb–treated mice and a further 5-fold reduction with MEDI3902 versus the PcrV antibody (Figure 6A). This is consistent with our neutrophil data indicating more efficient phagocytosis and processing of internalized bacteria in MEDI3902-treated animals. Although the neutrophil data were compelling, it is possible that other immune cells also contributed to the improved efficacy of MEDI3902 over Psl antibody treatment. We therefore visualized all immune cells in the lung vasculature using CD45 staining during the first 2 hours of infection and identified 2 other cell types associated with P. aeruginosa–GFP that did not stain for Ly6G (Figure 6B). Gating on Ly6G− cells by flow cytometry, we found a very small population of eosinophils (Ly6G−, Siglec-F−, CD11b−) and monocytes (Ly6G−, Siglec-F−, CD11b+) that were able to phagocytose P. aeruginosa in the presence of either anti-Psl mAb or MEDI3902 (Figure 6C). However, P. aeruginosa uptake by eosinophils and monocytes was minute compared with that by
neutrophils and similar in anti-Psl- and MEDI3902-treated animals, making it unlikely that other immune cells contribute to the efficacy of MEDI3902 (Figure 6, B and C).

Uncontrolled inflammation (cytokine storm) can often contribute to the pathology associated with *P. aeruginosa* infections (25). Mice treated with antibodies against PcrV and Psl had cytokine and chemokine profiles similar to those of control IgG–treated mice. By contrast, MEDI3902–treated mice exhibited a significant reduction in all of the major proinflammatory cytokines, including TNF-α, IL-1β, and IL-6, and the chemokine KC/GRO compared...
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**Discussion**

P. aeruginosa is a highly adaptable opportunistic pathogen that can cause life-threatening infections in immunocompromised and critically ill patients. The T3S injectisome and the Psl exopolysaccharide are known to be key factors promoting acute and persistent infections (14, 16). Expression of the T3S components and Psl has been shown to be prevalent, with approximately 98% of isolates capable of expressing 1 or both of these antigens (25, 26). Indeed, antibodies targeting 1 or both (24) of these components have shown promising activity against different P. aeruginosa strain types in multiple preclinical infection models. However, until now a mechanism of action for how these antibodies were working poorly understood. Using multilaser spinning-disk microscopy to visualize P. aeruginosa infection in the lung and liver, we observed that P. aeruginosa evasion and subversion mechanisms predominate in the lung vasculature but not in liver, where intravascular macrophages, known as Kupffer cells, are capable of eradicating many of the bacteria. By initially using isogenic P. aeruginosa pslA or pcrV knockout strains stably expressing GFP that were deficient in Psl exopolysaccharide expression or had a dysfunctional T3S injectisome, respectively, we deduced the primary role of these pathogenic factors in vivo. In the lung vasculature, imaging in real time revealed that P. aeruginosa uses Psl to impair complement function, thereby remaining invisible to neutrophils and allowing colonization of the organ. By visualizing the host-pathogen response in vivo, we determined that P. aeruginosa effectively evaded neutrophils, suggesting that Psl plays a prominent role in preventing complement fixation in an active infection. We also demonstrate that anti-Psl antibodies acted as opsonins allowing neutrophils in the pulmonary microvasculature to bind and phagocytose P. aeruginosa. In addition, our data demonstrate that the mechanism of neutrophil recognition and phagocytosis of P. aeruginosa in the lung vasculature required activation of complement. Entirely consistent with these findings is a previous in vitro report stating that higher quantities of complement components were deposited on the Psl-deficient strain compared with the isogenic WT parent strain (16).

Interestingly, despite increased uptake of the pathogen in the presence of the anti-Psl mAb, we observed very little phagocytosis of the bacterium if it possessed a functional T3S. Our data are consistent with previous reports indicating that expression of the T3S system by some P. aeruginosa strain types helps the bacteria avoid intracellular destruction (14). A study looking at corneal epithelial cells that internalize P. aeruginosa showed that the T3S effector ExoS helped the bacteria evade acidic vacuoles once inside the epithelial cell (14). We hypothesize that a similar method of evasion is occurring once neutrophils phagocyte
Figure 6. MEDI3902 increases monocyte/eosinophil recognition of \textit{P. aeruginosa} and decreases lung bacterial load and inflammation. (A) Bacterial CFU were determined at 6 hours after i.v. infection with \textit{P. aeruginosa} in lung after prophylactic antibody treatment. \( n = 6 \). Error bars represent SEM. One-way ANOVA statistical analysis was performed on data points; multiple comparisons were performed between indicated columns. \(* P = 0.0255, \** P = 0.0109\). (B) Mice were stained for CD45 (blue) for all hematopoietic cells and Ly6G (red) for neutrophils. The percentage of neutrophil-negative but CD45-positive cells containing \textit{P. aeruginosa}–GFP was assessed at 1 hour after infection (number of FOV = 3, \( n = 3 \)). Error bars represent SEM. (C) Flow cytometry was performed to identify Ly6G-negative cells that contained \textit{P. aeruginosa}–GFP. Eosinophils are intermediate for Siglec-F and high for CD11b, while monocytes are Siglec-F negative and Cd11b high (\( n = 3, 3 \) FOV). Error bars represent SEM. One-way ANOVA statistical analysis was performed; \( P = 0.0667 \). (D) Whole lung homogenates were used to determine the cytokine/chemokine milieu 6 hours after infection with \textit{P. aeruginosa}. One-way ANOVA analysis was performed on the indicated columns. \( P < 0.05 \) is significant. Error bars represent SEM; \( * P = 0.0373, \** P = 0.0067, \*** P = 0.0003 \). (E) Survival analysis of mice prophylactically treated with therapeutic antibodies and then subsequently infected i.v. with \textit{P. aeruginosa} strain 6077. Mantel-Cox test was performed. There was a significant difference between MEDI3902-treated mice and IgG control mice (\( P < 0.0001 \)). There was a significant difference between anti-PcrV–treated mice and IgG control mice (\( P < 0.0001 \)). Finally, there was a significant difference between mice treated with a mixture of the antibodies (anti-Psl + anti-PcrV) and MEDI3902-treated mice (\( P = 0.0491 \)). Error bars represent SEM; \( n = 8 \) mice per group. All experiments were repeated 3 times unless otherwise indicated.
P. aeruginosa. In our model we propose that targeting Psl with the antibody increases recognition of the bacteria by neutrophils (Figure 7). However, inhibition of T3S secretion is required for optimal neutrophil killing of the bacteria (Figure 7). Lung neutrophils from mice receiving anti-PcrV enabled efficient processing of P. aeruginosa into low-pH compartments, but the number of neutrophil-bacterial interactions was very low in the absence of anti-Psl specificity (Figure 7). Indeed, bispecific mAb MEDI3902, which effectively and simultaneously targets Psl and PcrV, was by far the most effective therapeutic we tested.

In this bacteremic model, we also noted that anti-PcrV and anti-Psl antibodies in combination were not as effective as bispecific mAb MEDI3902, and neutrophils from mice treated with mAb mixture were not as effective in eradicating P. aeruginosa. With the in vivo imaging we were able to visualize which cells in the lung and liver phagocytized P. aeruginosa, and with our ability to visualize capture, phagocytosis, and processing of bacteria inside immune cells, we identify why the antibody combination within a single molecule is superior in its protective activity in comparison with the individual anti-Psl and anti-PcrV mAbs and the mAb mixture. Having both anti-Psl and anti-PcrV activities on a single molecule allows increased bacterial uptake (mediated by the anti-Psl component) by neutrophils while concentrating the colocated anti-PcrV on the bacterium; this was seen by the fact that we could detect antibody in MEDI3902-treated neutrophils but not in anti-PcrV–treated neutrophils (Supplemental Figure 4). The anti-PcrV activity of MEDI3902 is thus positioned to inhibit the T3S injectisome as soon as it is expressed, even if it is expressed only after bacteria are ingested intracellularly, thereby resulting in more effective phagosomal processing and eradication of P. aeruginosa.

Patient prognosis is known to be poor upon systemic dissemination of P. aeruginosa (26). This bacterium is able to enter the bloodstream of immunocompromised patients, often through i.v.
catheters during essential invasive interventions. In this study, we use a blood-borne P. aeruginosa infection model to assess important immune cell mechanisms in bacterial clearance from the host. In fact, we show that P. aeruginosa effectively bound pulmonary endothelium but not hepatic endothelium and evaded lung neutrophils but not liver macrophages (Kupffer cells). Interestingly, this correlated into 2 key phenotypes: (a) greater dissemination of P. aeruginosa from the vasculature into the lungs, and (b) increased mortality. In addition to the direct bacterial damage, “friendly fire” resulting from hyperactivation of the immune system in systemic infections causes further injury and organ failure. Multiforgan failure closely precedes death of the host. Interestingly, MEDI3902-treated mice had lower levels of many proinflammatory cytokines and chemokines at 6 hours after infection (Figure 6D) in the lung, which was not observed for the individual anti-PcrV and anti-Psl mAbs in comparison with the control IgG. It could be that MEDI3902 allows for effective clearance of P. aeruginosa by pulmonary neutrophils avoiding frustrated phagocytosis and cell lysis, leading to inappropriate release of many of the proteases and other noxious molecules that can injure the host. It could also be the ability of MEDI3902 to inhibit T3S injectisome function more effectively compared with the other antibodies that leads to less cytokine production by the host. It has previously been reported that T3S expression by P. aeruginosa is linked to increased IL-1β production (27), which could then induce the production of the many other cytokines and chemokines. In MEDI3902-treated mice, neutrophils appeared more active as evidenced by the ability of neutrophils to engulf multiple bacteria, an outcome not observed in mice treated with anti-PsI, anti-PcrV, or a mAb mixture. Our data suggest that this increased fitness is due to intracellular inhibition of the T3S injectisome.

In conclusion, whether it is during pneumonia leading to bacteremia or bacteremia leading to lung damage, the lung microvasculature is a battleground where neutrophils are critical in containing P. aeruginosa lung infections. Here we have visualized the lung of a live mouse during the first few hours of i.v. P. aeruginosa infection. Using intravital microscopy we found that under basal conditions, P. aeruginosa expressing Psl evades neutrophil recognition in this organ, and when phagocytosed the pathogen can prevent proper processing into low-pH phagolysosomes via a functional uninhibited T3S system. An antibody directed at both of these molecular targets allows enhanced neutrophil phagocytosis and efficient processing of P. aeruginosa and furthermore allows neutrophils to progressively capture and process more than a single bacterium. Precision immunotherapeutic mAbs offer several advantages in comparison with broad-spectrum antibiotics, including extended half-lives without disruption of the beneficial microbiome and without the potential to generate cross-resistance in on-targeted organisms. Given the diversity among P. aeruginosa strains, mAb-based approaches against multiple serotype-independent and highly conserved and pivotal pathogenic mechanisms, such as those targeted herein, may provide significant opportunity for clinical success. Currently, the MEDI3902 mAb is in phase 1b clinical trials where it is given to ventilated patients to prevent the development of pneumonia (17). From the novel mechanisms elucidated in this study, the success of MEDI3902 may also extend beyond treating for bacteremia per se, to treating patients with pneumonia prophylactically to decrease the likelihood of the bacteria entering the systemic circulation and disseminating to other organs.

Methods
Mice. Six- to 8-week-old C57BL/6 WT male or female mice were purchased from The Jackson Laboratory. All mice were housed under pathogen-free conditions and fed autoclaved rodent feed and water. B6.129S4-C3mIcr.J, referred to as C3–/– mice in this paper, were purchased from The Jackson Laboratory (catalog 3641). B6.129P2-FcRγ1mAmr N12, referred to as FcγR-deficient or FcεRγ1–/– mice in this paper, were purchased from Taconic (catalog 583). C3–/– and FcεRγ1–/– mice were always used in experiments at 6–8 weeks of age. All experiments were performed in male mice.

Intravitral microscopy. Multichannel spinning-disk intravital microscopy was used to image the mouse lungs. We used Ly6G clone IA8 antibodies (Biolegend) to identify neutrophils, F4/80 clone BM8 (eBioscience) to identify Kupffer cells, and CD31 to identify endothelium clone 390 (eBioscience). For intravitral experiments we used a very low concentration at 2 μg per mouse of each antibody.

Mice were anesthetized (10 mg/kg xylazine hydrochloride and 200 mg/kg ketamine hydrochloride) and body temperature maintained using a heating pad. The right jugular vein was cannulated to administer additional anesthetic and fluorescent dyes. For lung imaging, the trachea of the mouse was exposed and a small catheter threaded into the trachea, and this was used to attach the mouse to a small rodent ventilator (Harvard Apparatus). The mouse was placed on its side, and 3 ribs were removed in order to expose the lung. The lung was stabilized with a small suction chamber as described previously (28). Suction was maintained at 30 mmHg. Images were acquired with an upright microscope (BX51; Olympus) using a x20/0.95W NA water XLUM Plan Fl objective. The microscope was equipped with a confocal light path (WaveFx; Quorum) based on a modified Yokogawa CSU-10 head (Yokogawa Electric). A 512-by-512-pixel back-thinned electron multiplying charge-coupled device camera (C9100-13; Hamamatsu) was used for fluorescence detection. Simultaneous behavior of multiple cell types in the lung was assessed using 3 laser excitation wavelengths (488, 561, and 635 nm) (Cobolt AB) in rapid succession and visualized with the appropriate long-pass filters (Semrock). Velocity software (Improvision) was used to acquire and analyze images. For 3D reconstruction, a Z-stack of 1-μm slices was captured for each mouse at a rate of 10 frames per minute. Intravitral data were repeated with 3 strains of P. aeruginosa, PAO1, 6206, and 2259, as well as isogenic mutants deficient in Psl (Δpsl) or T3S (ΔpcrV) in the 6206 background.

For liver imaging, mice were anesthetized and cannulated as described above, and intravitral imaging was performed as previously described (19, 20).

Lung image quantification. Ten-minute videos with 3 fields of view were captured for each mouse at a rate of 10 frames per minute. Mice were infected in real time and imaged for 1 hour after infection. Bacterial fluorescence was significantly higher than background autofluorescence of untreated lungs. The total number of neutrophils was quantified for each video, and the number of neutrophils with bacteria was enumerated. The percentage of neutrophils with bacteria was then plotted. Only neutrophils that colocalized with bacteria for more than 30 seconds were considered to have “internalized” the
bacteria. Average bacterial phagocytosis by neutrophil was calculated for graphs; this was for the first 1–3 hours of infection. All videos were captured at the same speed and exposure time: 492 ms exposure at 10 frames per minute.

For pHrodo Red experiments, the number of neutrophils with bacteria was calculated, and the percentage of these that had red bacteria was subsequently calculated and graphed.

For Figure 4F, neutrophils with bacteria were tracked for 20 minutes, and the number of neutrophils containing bacteria was calculated; the percentage of these neutrophils containing bacteria that took up 1 or more other bacteria was then calculated and graphed.

**Liver image quantification.** For quantification of bacterial phagocytosis in the liver, 5 random fields of view with ×10 objective were selected before injection of bacteria. Acquisition of images was initiated, and after 1 minute of acquiring background images, fluorescent reporter bacteria were injected i.v. via tail vein. The “Find objects” function in Velocity software was used to identify individual captured bacteria by Kupffer cells (F4/80-positive cells in liver), when appropriate autofluorescent spots were subtracted from the final quantification.

For quantification of spinning-disk intravital images 3 hours after infection, computer-generated stitched images of 2 mm² were generated using the stitched image function in Velocity. Velocity software was used to calculate relative GFP fluorescence as measurement for the presence of *P. aeruginosa*-GFP in the liver. Spinning-disk intravital microscopy images of uninfected mice were used to determine the background fluorescence; *P. aeruginosa*-GFP was quantified with the same settings for all mutants and treatments. All videos and images were captured at the same speed and exposure time: 400 ms exposure at 3 frames per minute.

**Therapeutic antibody treatment.** Mice were prophylactically treated with therapeutic MEDI3902 antibody (bispecific mAb targeting Psl and PcrV), individual MEDI3902 parental mAbs targeting either Psl (Psl0096) or PcrV (V2L2-MD), or a parental mAb mixture at a dose of 1 × 10⁸ intravenously. Prophylactic treatment with therapeutic antibodies, mice were given 3.8 mg/kg of control IgG, anti-Psl, anti-PcrV, mAb mixture (3.8 mg/kg of each mAb), or the molar equivalent dosage of MEDI3902 (5 mg/kg) through the i.p. route. Twenty-four hours after prophylactic treatment with therapeutic antibodies, mice were given 1 × 10⁸ CFU of *P. aeruginosa* strain 6077 intravenously. Mice were monitored hourly for clinical symptoms; mice that lost more than 20% of their body weight were sacrificed using CO₂.

**Flow cytometry.** Mice were euthanized using CO₂. Lungs were perfused with 10 ml of ice-cold saline injected through the right ventricle of the heart. Perfused lungs were removed, minced, and placed in digestion buffer (PBS with 1 mg/ml collagenase type X1, 60 U DNase). Lungs were digested for 30 minutes and pushed through a 70-µm filter. Red blood cells were removed using ACK lysis buffer (Invitrogen), and samples were Fc blocked for 30 minutes. Samples were stained for 30 minutes with indicated antibodies. Samples were run using BD Canto and analyzed using FlowJo software. Flow cytometry was performed with PAO1-GFP and 6206-GFP strains. Bacterial strains used are indicated on the y axes of flow plots. The following antibodies were used in flow analysis: rat anti–mouse CD45 (clone 30-F11; eBioscience), rat IgG2a antibodies were used in flow analysis: rat anti–mouse CD45 (clone RA3-6B2; eBioscience), anti–mouse CD11c (clone N418; eBioscience), Armenian hamster IgG1 isotype control, and rat anti–mouse Siglec-F (E50-2440; BD Biosciences). All experiments were repeated with 3 biological samples 3 times. Isotype and fluorescence minus one (FMO) controls were used. Cells were gated based on total CD45-positive population, then on B220-negative and Ly6G-positive. Within the Ly6G-positive cells, percentage of PAO1-GFP-positive cells was quantified.

**Statistics.** All error bars represents SEM. One-way ANOVA was performed with multiple comparisons as described in the figure legends. All statistical analyses were performed using the GraphPad Prism program. Unless otherwise indicated, there were 4 mice in each group, and for intravital experiments 3 fields of view were gathered for each mouse. For flow cytometry each group was repeated with 3–4 mice. A P value of less than 0.05 was considered statistically significant.
Author contributions
AT and AD designed and performed experiments, analyzed data, and wrote the manuscript. BGJS, AEK, GW, PW, MEW, and ASN performed experiments. CKS contributed to experimental design and helped to write the manuscript. BRS contributed to experimental design. AD and PK conceptualized and directed the study, analyzed data, and wrote the manuscript.

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