**RESEARCH ARTICLE SUMMARY**

**IMMUNOLOGY**

**Bacteriophage trigger antiviral immunity and prevent clearance of bacterial infection**

Johanna M. Sweere, Jonas D. Van Belleghem, Heather Ishak, Michelle S. Bach, Medeea Popescu, Vivekananda Sunkari, Gernot Kaber, Robert Manasherob, Gina A. Suh, Xia Cao, Christiaan R. de Vries, Dung N. Lam, Payton L. Marshall, Maria Birukova, Ethan Katznelson, Daniel V. Lazzareschi, Swathi Balaji, Sundee G. Keswani, Thomas R. Hawn, Patrick R. Secor, Paul L. Bollyky*

**INTRODUCTION:** We have identified previously unsuspected, directly pathogenic roles for bacteriophage (phage) virions in bacterial infections. In particular, we report that internalization of phage by human and murine immune cells triggers maladaptive viral pattern recognition receptors and suppressed bacterial clearance from infected wounds.

**RATIONALE:** Bacteriophage are abundant at sites of bacterial infection, but their effect on mammalian immunity is unclear. To investigate this, we studied *Pseudomonas aeruginosa* (*Pa*), a major human pathogen associated with chronic wounds and other infections, and Pf, a filamentous phage produced by *Pa*. Notably, Pf is lysogenic and its production does not typically destroy its bacterial host, unlike the lytic phage used in phage therapy for bacterial infections. Previous work had suggested that Pf phage are important in the pathogenesis of *Pa* infections, although the underlying mechanisms were unclear. Here, we have examined the impact of Pf on *Pa* wound infections in humans and in animal models.

**RESULTS:** We report that Pf bacteriophage were present in 25 of 37 (68%) *Pa*-infected wounds in our cohort. Furthermore, wounds infected with Pf-positive strains were significantly older than wounds infected with Pf-negative strains, and Pf was more commonly found in chronic, nonhealing wounds. Consistent with this finding, in a murine wound infection model, Pf-positive strains of *Pa* required an average of 50 times fewer bacteria than Pf-negative strains to establish wound infections. Additionally, mice infected with Pf-positive strains of *Pa* exhibited greater morbidity and mortality than mice infected with Pf-negative strains.

Mechanistically, these effects were associated with endocytosis of Pf phage by mammalian immune cells, both in vivo and in vitro. We found that uptake of Pf phage resulted in the production of phage RNA, which, in turn, triggered Toll-like receptor 3 (TLR3)– and TIR domain-containing adapter-inducing interferon-β (TRIF)–dependent type I interferon production, the inhibition of tumor necrosis factor factor production, and the suppression of phagocytosis. These data suggest that a natural (unmodified) bacteriophage may be able to produce mRNA within human cells.

Consistent with a pathogenic role for Pf phage, we report that a vaccine against Pf phage protects against *Pa* wound infections. Passive immunization of mice with monoclonal antibodies against Pf was likewise effective in protecting against *Pa* infection by enhancing the opsonization of *Pa* bacteria.

**CONCLUSION:** These results reveal direct, pathogenic roles for phage virions in bacterial infections. Building upon these insights, we report that vaccination against phage virions represents a potential therapeutic strategy for the prevention of infections by antibiotic-resistant *Pa*. These findings may have broad utility and impact beyond the pathophysiology of chronic wound infections. *Pa* is a major pathogen in other clinical settings as well, including lung infections in cystic fibrosis. Moreover, many other Gram-negative bacteria, including *Klebsiella pneumoniae*, *Salmonella enterica*, *Vibrio cholerae*, and *Escherichia coli*, have the capacity to harbor similar filamentous phage (genus Inovirus).

Indeed, several of these phage are known to contribute to the virulence potential of their host bacteria. We propose that filamentous phage may be relevant to human interactions with a broad range of pathogenic and commensal bacteria and that these viruses may have profound, direct effects on human health and disease.

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**RESEARCH ARTICLE**

**IMMUNOLOGY**

**Bacteriophage trigger antiviral immunity and prevent clearance of bacterial infection**

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Bacteriophage are abundant at sites of bacterial infection, but their effects on mammalian hosts are unclear. We have identified pathogenic roles for filamentous Pf bacteriophage produced by *Pseudomonas aeruginosa* (Pa) in suppression of immunity against bacterial infection. Pf promote Pa wound infection in mice and are associated with chronic human Pa wound infections. Murine and human leukocytes endocytose Pf, and internalization of this single-stranded DNA virus results in phage RNA production. This triggers Toll-like receptor 3 (TLR3)– and TIR domain–containing adapter-inducing interferon-β (TRIF)–dependent type I interferon production, inhibition of tumor necrosis factor (TNF), and the suppression of phagocytosis. Conversely, immunization of mice against Pf prevents Pa wound infection. Thus, Pf triggers maladaptive innate viral pattern-recognition responses, which impair bacterial clearance. Vaccination against phage virions represents a potential strategy to prevent bacterial infection.

*Pseudomonas aeruginosa* (Pa) is a Gram-negative bacterium found in infections of wounds, pressure ulcers, and burns that is responsible for extensive morbidity and mortality (1–3). The World Health Organization recently categorized Pa as a “priority pathogen” with the greatest risk to human health (4). Pa clinical isolates are often antibiotic-resistant, and no approved vaccine against Pa is available. Alternative strategies to prevent Pa infections are desperately needed.

The clearance of Pa infections typically requires effective phagocytosis and production of tumor necrosis factor (TNF) (5–7). However, Pa exhibits multiple strategies that inhibit bacterial clearance (8), allowing opportunistic infections to be established (9).

At sites of infection, filamentous Pf bacteriophage (Pf phage) are produced in abundance by Pa (10–12). Unlike lytic bacteriophage, temperate phage such as Pf do not typically lyse their bacterial hosts. Instead, they integrate into the bacterial chromosome as a prophage that is passed on to daughter cells. When filamentous phage virions are produced, they are generally extruded without bacterial lysis (14, 15). We recently reported that Pf phage act as structural elements in Pa biofilms (12, 13) and that they contribute to reduced inflammation in acute murine lung infections through unclear mechanisms (13). Here, we investigated the possibility that Pf phage promote Pa infections through direct effects on mammalian immunity.

**Pf phage are found in human wounds and promote bacterial wound colonization, morbidity, and mortality in mice**

To determine whether Pf phage are found in chronic Pa infections, we sampled 111 patients with infected nonhealing wounds referred to the Infectious Disease Service at the Stanford Advanced Wound Care Center over a period of 20 months and identified 37 patients infected with Pa (table S1).

We first examined the incidence of Pf phage in this cohort. Using a quantitative polymerase chain reaction (PCR) assay to measure Pf phage (12), we found that 68% (25 out of 37) of Pa-infected human wounds in our study harbor Pf phage (Fig. 1, A to C). This is consistent with a previous report that 60% of 241 clinical Pa isolates were Pf phage–positive (16). Thus, Pf is frequently present in human Pa infections, including wounds.

By contrast, only 22% (2 out of 9) of wounds younger than 6 months were Pf-positive (Fig. 1D). In addition, patients infected with Pf-positive strains of Pa were younger than patients infected with Pf-negative strains (table S1). Thus, Pf is associated with chronic wounds in this cohort of human patients.

To experimentally assess the possibility that Pf phage promote Pa infections, we examined the ability of Pf-positive versus Pf-negative strains of Pa to establish wound infections in mice. To this end, we generated full-thickness, excisional wounds (20), allowed nascent wound eschars to form for 24 hours, and then inoculated these wounds with murine Pa strains PAO1, a bacterial wound isolate infected by Pf phage strain Pf4 (Fig. 1E and fig. S1A). In this model, we measured the presence or absence of bacterial infection 3 days after inoculation by quantifying luminescence relative to background signal (Fig. 1, F and G). On average, wounds infected with PAO1 contained ~109 copies of Pf4 phage per wound (fig. S1B). A strain of Pa lacking the Pf4 prophage, PAO1ΔPf4 (11), showed no differences in growth rate or luminescence emission compared to its isogenic parent PAO1 (fig. S1, C to E).

We then asked whether Pf4 phage contributes to bacterial pathogenesis. We examined the infection rate (the number of wounds with detectable and sustained Pa signal 72 hours after inoculation over the number of total wounds initially inoculated with Pa). Wounds deemed “infected” were those with luminescent bacterial signal above baseline. PAO1 and PAO1ΔPf4 differed significantly in their ability to establish wound infections in this model, with calculated IC50 values (bacterial dose at which 50% of wounds become colonized) of ~7.7 × 107 colony-forming units (CFU)/ml and ~3.8 × 106 CFU/ml, respectively (Fig. 1H and fig. S2A). In subsequent experiments, mice received an inoculum of 7.5 ± 2.5 × 106 CFU/ml Pa, unless otherwise noted. Inoculation with PAO1 led to significantly higher infection rates than PAO1ΔPf4 (Fig. 1I and fig. S2A), whereas the reintroduction of Pf4 phage to PAO1ΔPf4 established infection rates and bacterial burdens comparable to those of infections caused by wild-type PAO1 (Fig. 1J and fig. S2B). The supplementation of additional Pf4 to PAO1 did not affect the infection rate or bacterial burden (fig. S2, C and D). Thus, Pf4 phage reduce the number of bacteria required to establish an infection.

We then explored the impact of Pf4 phage on morbidity and mortality. As inoculation with neither PAO1 nor PAO1ΔPf4 at a dose of 7.5 ± 2.5 × 106 CFU/ml resulted in any mortality, we...
increased our inoculation dose to $10^7$ CFU/ml. This resulted in a 100% infection rate for both PAO1 and PAO1P4. In this setting, mortality was much higher in mice infected with PAO1 than in mice infected with PAO1P4 (Fig. 1K). Moreover, PAO1-infected mice that survived lost more weight than PAO1P4-infected mice (Fig. 1L). The bacterial burden over time was not statistically different at this higher inoculum (fig. S2, E and F). Thus, P4 phage promotes both the incidence and virulence of Pa wound infections in this model.

Stimulation with P4 phage triggers antiviral immunity and impaired bacterial clearance

Given the enhanced bacterial infection rate observed in the presence of P4 phage, we asked whether P4 phage might affect immune cell recruitment. However, analysis of wound leukocyte infiltrates by flow cytometry or histology showed no notable differences in either immune cell count or composition, granulature tissue, or epithelial gap (figs. S3, A to I, and S4, A to J).

We then considered whether P4 phage might improve bacterial infection rate through effects on phagocytosis and antibacterial cytokine secretion. To test this, we first measured the uptake of Pa by murine and human phagocytes in the presence or absence of P4. We found that the presence of P4 phage caused up to a 10× reduction in the number of bacteria internalized by murine bone marrow–derived dendritic cells (BMDCs) (Fig. 2A) and bone marrow–derived macrophages (BMDMs) (fig. S5A), as well as human U937 (Fig. 2B) and primary (fig. S5B) macrophages. The presence of P4 phage also reduced phagocytic clearance of clinical Pa isolates collected from human wounds (fig. S5C). Introduction of P4 phage to PAO1P4 was sufficient to inhibit phagocytic uptake (fig. S5D). Phagocytic killing was not affected by the presence of P4 (fig. S5E). Moreover, murine phagocytes incubated with purified P4 likewise engulfed fewer fixed Escherichia coli particles than phagocytes stimulated with an equivalent volume of phosphate-buffered saline (PBS) carrier (Fig. 2, C and D, and fig. S5, F and G). Phagocytosis was not affected by increased adsorption of bacteria or bacterial particles to the exterior of phagocytes in the presence of P4 (fig. S5, H and I). Thus, P4 acts directly on phagocytes to inhibit bacterial engulfment.

Because the cytokine TNF is critical for phagocytosis and Pa clearance (5, 6), we next asked whether P4 affected TNF production. We stimulated murine phagocytes with lipopolysaccharide (LPS) or algin-ate—proinflammatory bacterial products known to induce TNF production (21)—and then added purified P4 or an equivalent volume of PBS carrier. Although both LPS and alginate induced robust TNF secretion, costimulation with P4 reduced TNF in murine phagocytes (Fig. 2, E and F, and fig. S6A) and human primary monocytes (Fig. 2G).

Because the P4-mediated inhibition of TNF occurred early after stimulation with bacterial products (fig. S6B), we hypothesized that the addition of exogenous TNF could overcome the inhibitory effects of P4 on phagocytosis. Pre-treatment of phagocytes with TNF rescued phagocytic function in vitro despite the presence of P4 (Fig. 2H and fig. S6C) and reduced in vivo PAO1 wound infection (fig. S6D).

Notably, P4 did not affect phagocyte viability (fig. S7, A and B), and P4-mediated immune inhibition was not due to the presence of contaminating nucleic acids (fig. S7C), protein (fig. S7D), or endotoxin (fig. S7, E and F, and detected levels <0.05 endotoxin units (EU)/ml) in our preparations. In addition, mock phage preparations generated from the PAO1P4 strain with the same protocol used to isolate P4 from PAO1 did not impair TNF secretion (fig. S7, G and H) or phagocytosis (fig. S7, I and J), indicating that P4 and

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**Fig. 1. Pf phage promote Pa wound infection.** (A and B) Representative images of human Pa-infected wounds (A) negative and (B) positive for Pf. Scale bars: 5 mm. (C) Prevalence of Pf phage in infected human wounds. (D) Prevalence of Pf phage in human wounds younger (n = 9) and older (n = 28) than 6 months of age: two-tailed Fisher’s exact test. (E) The full-thickness wound infection model. (F) Representative images of murine wounds before (left) and after infection (middle) showing luminescent bacterial signal (right). Scale bars: 5 mm. (G) Luminescent signal reflecting wound bacterial burden after inoculation with $7.5 \times 10^2$ CFU/ml PAO1 (n = 14 wounds), the same dose used in (H) to (J).
not residual bacterial components conceivably present in our preparations were responsible for immune inhibition. Thus, TNF is implicated in the Pf4-mediated inhibition of phagocytic capacity.

Next, we wanted to define the mechanism of Pf4-mediated immune inhibition. BMDCs were stimulated with proinflammatory bacterial products such as alginate or LPS. We found that Pf4-mediated inhibition of TNF occurred in a concentration-dependent manner, which was maintained over time (Fig. 3, A and B). The Pf phage Pf1, but not the E. coli filamentous phage Fd1, also inhibited TNF production (Fig. 3, C and D) and phagocytosis (Fig. S8, A and B). Thus, this effect was common among Pf phage but was not universal across all inoviruses. Both LPS and alginate still triggered the nuclear translocation of nuclear factor κB (NF-κB) (Fig. S9, A to C), a key transcription factor associated with microbial-sensing pathways (22). Thus, phagocytes still sense these molecules in the presence of Pf4. TfNf mRNA expression levels likewise remained intact, despite the reduction in TNF protein (Fig. 3, E and F, and Fig. S8, C and D). Notably, the inhibition of TNF in this system did not begin until 240 min after stimulation following a modest, transient increase in TNF (Fig. S3C). This suggests that the inhibition of TNF was somewhat delayed relative to its induction. Thus, Pf4 inhibits TNF production otherwise induced in response to bacterial products. Furthermore, this inhibition occurs at the level of mRNA translation.

One factor that can suppress TNF production at the level of mRNA translation is type I interferon (IFN) (23, 24). Indeed, Pf4 treatment in the setting of inflammatory bacterial stimuli promoted the production of type I IFN (Fig. 3G). Moreover, in phagocytes generated from mice lacking either the type I IFN receptor (Ifnar1−/−) or both the type I and type II IFN receptors (Ifnar1−/− Ifnar2−/−), Pf4 treatment did not reduce TNF production (Fig. 3H and Fig. S8F) or impair phagocytosis (Fig. S8G) compared to wild-type controls. Thus, Pf4 reduces TNF production in a type I IFN-dependent manner.

Consequently, we hypothesized that Pf phage is recognized by antiviral pattern recognition receptors (PRRs) within the Toll-like receptor (TLR) family, which signal through the adapters TIR domain–containing adapter-inducing interferon-β (TRIF) or myeloid differentiation primary response 88 (MyD88) (25). Pf4-mediated suppression of TNF production (Fig. 4A) and phagocytosis (Fig. 4B) was abrogated in TIRβ−/− murine phagocytes but was maintained in MyD88−/− cells (Fig. 4C), suggesting that Pf phage requires TRIF to interfere with TNF translation. This is consistent with recent reports that the simultaneous triggering of MyD88–NF-κB and TRIF pathways leads to type I IFN transcription (26) and that TRIF-mediated signaling pathways affect TfNf mRNA translation in other settings (27).

To determine which specific antiviral PRR upstream of TRIF might mediate Pf-mediated immune inhibition, we generated bone marrow–derived phagocytes from mice deficient for either the antiviral sensors TLR3 and TLR9 or the anti-viral sensor TLR2 (as a control). We also generated phagocytes from mice lacking the single-stranded DNA (ssDNA) sensors cyclic GMP-AMP synthase (cGAS) or stimulator of interferon genes (STING) (28, 29), as Pf phage have ssDNA genomes. We measured TNF production by these cells in response to bacterial inflammatory stimuli alginate or LPS in the absence or presence of Pf4. Pf4 did not diminish TNF production or phagocytic capacity in Tlr3−/− phagocytes (Fig. 4, D and E, and fig. S10, A and B), nor did it up-regulate type I IFN production in Tlr3−/− cells (Fig. 4F). TNF suppression was intact in Tlr9−/−, Tlr2−/−, Cx30−/−, and Sting−/− phagocytes (fig. S10, C to E). Finally, Pf4 phage did not promote wound infection in Tlr3−/− mice (Fig. 4G). Thus, the canonical antiviral receptor TLR3 and the adapter TRIF are required for Pf4 to inhibit TNF production, which consequently promotes bacterial infection.

To ascertain which components of Pf4 trigger TLR3, we used a Tlr3-reporter cell line. There was robust TLR3 signaling in response to whole Pf1 and Pf4 phage (Fig. 4H). As TLR3 detects double-stranded RNA (dsRNA) rather than ssDNA, we added purified RNA from Pf4 phage genes to the Tlr3-reporter cells and observed TLR3 activation in response to several of them (Fig. 4I). Thus, Pf4 RNA can trigger TLR3. Finally, we stimulated human monocytes with whole Pf1 or Pf4 phage and subsequently detected RNA derived from various phage genes (Fig. 4J). Considering that ribonuclease (RNAse) treatment of purified Pf4 did not abrogate TNF inhibition (fig. 7C), phase internalization appeared to result in RNA synthesis, which, in turn, triggered TLR3 responses. This hypothesis is consistent with a previous report that used a related filamentous phase modified to facilitate cell entry and to express a reporter gene (30). However, our data suggest that a natural bacteriophage can produce mRNA within human cells. We cannot exclude the possibility that a small amount of viral RNA may have been inadvertently packaged within some phage particles. Nonetheless, Pf phage and Pf phage RNA can trigger TLR3 signaling.

Given that TLR3 is present within endocytic vesicles (31), we next asked whether Pf4 is internalized by endocytosis. Fluorescently labeled Pf4 was internalized by murine phagocytes with minimal adsorption (Fig. 5, A and B; fig. S11A; and movie S1). Moreover, within these cells, Pf4 phage colocalized with TLR3 (Fig. 5C). Transmission electron microscopy imaging of Pf4 within BMDCs demonstrated the presence of Pf4 in endosomal or lysosomal vesicles and in the cytosol (Fig. 5, D and E, and fig. S11B to E). Treatment with brefeldin A,
a vesicular transport inhibitor, greatly reduced Pf4 uptake (fig. S11F). Inhibitors of receptor-mediated endocytosis and microtubule assembly—wortmannin and nocodazole, respectively—also reduced Pf4 uptake, as did chlorpromazine, an inhibitor of clathrin-mediated endocytosis, although to a lesser extent (Fig. 5F). Thus, Pf4 is internalized by mammalian cells, consistent with reports of uptake of other phages (32–34). Additionally, this internalization occurs by endocytosis, and Pf4 encounters TLR3 in endosomes and lysosomes.

We then asked which cell types internalize Pf4. Flow cytometric analysis revealed that most phagocytes that internalized Pf4 were positive for CD14 (fig. S11G), a receptor that enhances TLR3 signaling (35). Pf4 was taken up by diverse cell types within mouse splenocytes and lymphocytes and wound-infiltrating immune cells, including B cells and DCs (Fig. 5, G and H, and figs. S11H and S12). Finally, various human peripheral blood mononuclear cells (PBMCs), including monocytes, DCs, and B cells, also internalized Pf4 phage (Fig. 5, I and J, and fig. S13).

Thus, we propose a model in which Pf phage are endocytosed by leukocytes, whereupon the Pf particles or RNA trigger TLR3 and TRIF-dependent viral PRRs, driving type I IFN production, inhibiting TNF production, and limiting phagocytosis. These effects result in impaired bacterial clearance and more frequent infection (Fig. 6).

**Antibodies directed against Pf4 prevent Pa colonization**

We next asked whether targeting Pf4 therapeutically might prevent Pa wound colonization. To this end, we identified a region of CoaB, the major coat protein of Pf phages, that is broadly conserved across 669 isolates of Pa (Fig. 7A). We then immunized mice with a peptide version of this epitope conjugated to keyhole limpet hemocyanin (KLH) (fig. S14A). This protocol successfully generated humoral immunity against Pa phage (fig. S14B).

Using our infection model, we next tested the impact of vaccination against Pf phage on the development of stable Pa infection. Compared to mock vaccination, vaccination against the CoaB epitope reduced the incidence of Pa wound infections by half (Fig. 7B).

Because immunization itself might influence the incidence of wound infections, we asked whether transfer of passive, humoral immunity against Pa phage was sufficient to prevent Pa infection. To this end, we administered monoclonal antibodies (mAbs) generated against the same CoaB peptide epitope incorporated into our vaccine to mice during bacterial inoculation (fig. S14, C and D). Transfer of these mAbs into wounded skin in the presence of Pa significantly reduced stable wound infection compared to treatment with isotype control antibodies (Fig. 7C). Thus, Pf phage is critically important during early wound infection, and humoral immunity against Pf is protective against Pa infection.

Given these pronounced effects, we then asked how antibodies directed against Pf are protective. Antibodies against Pf phage did not prevent Pf phage internalization by bone marrow–derived phagocytes (fig. S14E). Instead, mAbs directed against Pf promoted phagocytic engulfment of PAO1 but not PAO1Δp6 (Fig. 7D). This enhanced phagocytosis was abrogated upon addition of anti-CD16/CD32 Fc block (Fig. 7E and fig. S14F). Thus, antibody-mediated recognition of Pf phage facilitates Pa phagocytosis. Because Pf phage adhere to type IV pili on the surface of Pa (10), it is possible that antibodies to Pf promote opsonization of bacteria coated with Pf phage.

**Discussion**

These findings establish pathogenic roles for Pf bacteriophage in Pa infections. In particular, we find that internalized bacteriophage trigger maladaptive, antiviral responses that suppress bacterial clearance, resulting in the establishment of Pa wound infections. Consistent with this, 68% of human Pa wound infections in our cohort contained Pf, and the presence of this phage was associated with more chronic wounds. Our data reveal direct roles for phage virions in the pathogenesis of bacterial infection by suppressing antibacterial immunity.

These findings may have broad relevance beyond Pf phage and Pa wound infections. Pf phage are also abundant in Pa-associated respiratory infections (12) and are likely to be present in other Pa infections. Other phage can be internalized by mammalian cells (30, 32–34), and large numbers of phage are transcytosed by gut epithelial cells (36). We hypothesize that human cells may internalize phage produced by both commensal and pathogenic bacteria, and this direct interaction with immune cells could occur on a scale that substantially affects human health.

Consistent with a pathogenic role for Pf, both passive and active immunization against Pf phage protected against Pa infections. Because CoaB is highly conserved across Pf isolates, this strategy may have value in preventing infection with Pa. Other studies have reported that antibodies against bacteriophage clear phages and lead to a reduction of their antibacterial activity and worsening of infection (37). As other filamentous phage are implicated in bacterial colonization (38), immunization against phage virions may be a promising strategy to prevent other infectious diseases as well.

Our findings may complicate efforts to develop phage as a therapeutic option for antibiotic-resistant infections. However, to date, lytic phage have not been reported to suppress phagocytosis (39) or local inflammatory cytokine secretion (40). This suggests that temperate and lytic phage may differ in this regard. Lytic bacteriophages have
previously been reported to play an important, synergic role in the immunological clearance of bacterial pathogens (41). Interactions between mammalian hosts and their phageomes are likely to be dynamic and complex (42) and a promising area for further investigation.

In conclusion, these studies establish the importance of internalized bacteriophage in a bacterial infection and suggest that bacteriophages may have profound and direct effects on human health and physiology.

Materials and Methods

Mice

Mice were bred and maintained under specific pathogen-free conditions, with free access to food and water, in the vivarium at Stanford University. Mice that underwent surgery received additional 0.9% sodium chloride solution (Hospira, Cat. No. 0409-4888-10) and intraperitoneal injections of Suppical Pet Gel (Henry Schein Animal Health, Cat. No. 029908) and water, in the vivarium at Stanford University. Mice were bred and maintained under specific pathogen-free conditions, with free access to food and water, in the vivarium at Stanford University. Mice may have profound and direct effects on human health and physiology.

Jackson Laboratory-derived Ifnar−/− and Ifngr−/− mice for >10 generations. All experiments and animal use procedures were approved by the Institutional Animal Care and Use Committee at the School of Medicine at Stanford University.

Chemicals, antibiotics, and reagents

The following chemicals, antibiotics and reagents were used: lipopolysaccharide from *E. coli* O111: B4 (Sigma, Cat. No. LA439); alginic acid (Sigma, Cat. No. A09682); bovine serum albumin (BSA) (PeproTech, Cat. No. A0682); alginic acid (Sigma, Cat. No. L4391); alginate (Sigma, Cat. No. L4391); and alginate and Pf4 for 24 hours. (A) TNF production by WT and Tlr3−/− BMDMs stimulated with alginate and Pf4 for 48 hours. (B) E. coli–phrdo–positive cells in WT and Tlr3−/− Pf–stimulated BMDMs. (C) TNF production by WT and Myd88−/− BMDMs stimulated with alginate and Pf4 for 48 hours. (D) TNF production by WT and Tlr3−/− BMDMs stimulated with alginate and Pf4 for 24 hours. (E) E. coli–phrdo–positive cells in WT or Tlr3−/− BMDCs stimulated with Pf4. (F) Type I IFN production by BMDMs stimulated with LPS and Pf4 for 24 hours. (A) to (F) are each representative of *n* ≥ 3 experiments and depict mean with SEM of *n* = 3 replicates. Statistics: two-tailed Student’s t test. (G) Wound infection rate 3 days after infection in Tlr3−/− and WT mice inoculated with 7.5 ± 2.5 × 10^2 CFU/ml PAO1 or PAO1ΔP4. Summary of *n* = 2 experiments, *n* = 30 to 34 wounds/group. Statistics: two-tailed Fisher’s exact test. (H and I) Tlr3-reporter signal in response to (H) whole Pf1 and Pf4 phage; (I) RNA from Pf4 genes. (J) RNA detected in human monocytes stimulated with whole Pf1 or Pf4 phage for 24 hours. Summary of *n* = 2 (I and J) or *n* ≥ 3 (H) experiments with ≥ 3 replicates; depicted means are SEM.

*P. aeruginosa* strain PAO1 was used for all experiments unless stated otherwise. Isogenic phage-free strain PAO1AP4 is derived from strain PAO1, but PAO1AP4 lacks the genomic copy of Pf4 entirely (11). This strain can still be reinfected by Pf4. For one experiment, two clinical strains of *P. aeruginosa* isolated from infected human Pf-positive (patient 5) and Pf-negative (patient 320) wounds were used. In general, bacteria were prepared as follows. Frozen glycerol stocks were streaked on Luria–Bertani (LB) agar containing selective antibiotics (PAO1 and clinical strains: none; PAO1AP4: 10 μg/ml of gentamicin; luminescent strains: 100 μg/ml of carbenicillin and 12.5 μg/ml of kanamycin) and grown overnight at 37°C. An isolated colony was picked and grown overnight at 37°C in LB medium, pH 7.4 (for luminescent strains, broth contained 100 μg/ml of carbenicillin) under shaking, aerobic conditions. If Pf phage supplementation was required, aliquots were grown until mid-exponential phase and split in two. One split received ~1 × 10^6 PAO1 purified phage before overnight incubation. The next day, cultures were diluted to OD_600 (optical density at 600 nm) = 0.05 in 75 ml of LB medium and cultures were grown until the early exponential phase. OD_600 was measured and the required number of bacteria were calculated, washed, and readied for use in experiments.

Preparation of heat-killed bacteria

In brief, bacteria were prepared as follows. Frozen glycerol stocks were streaked on LB agar as described above. Individual colonies were grown in 5 ml of LB broth the next day for 2 hours to approximately 2 × 10^6 CFU/ml. The bacterial cultures were centrifuged at 6000g for 5 min,
and the pellet was washed in 1 ml of PBS three times. Finally, the pellet was resuspended in 1 ml of PBS and heated for 30 min at 90°C under shaking conditions. The preparation was checked for sterility by plating.

**Generation of bioluminescent bacteria**
Plasmid pUT-Tn5-EM7-lux-Km1 (43) was extracted from the *E. coli* Top10 strain using the QiaPrep Spin Miniprep Kit (Qiagen, Cat. No. 27104) according to the manufacturer’s instructions. The plasmid was used to transform CaCl2 competent PAO1 and its isogenic PA01ΔPf4 strain (11) as described previously (44). Restriction enzymes were used according to the vendor’s directions (New England BioLabs). Screening for PAO1 transformants was performed on LB agar plates supplemented with 100 μg/ml of ampicillin (Sigma, Cat. No. A958) and 50 μg/ml of kanamycin.

**Phage purification**
Three phage strains—two from *Pa* (Pf1, Pf4), and one from *E. coli* (Pfd1)—were studied. *Pa* strain K produced Pf1 and *Pa* strain PAO1 produced Pf4. All supernatants from PAO1ΔPf4 were prepared and diluted in sterile PBS in the same manner as other phage samples. Phage were harvested following treatment with 1 μg/ml of DNase I (Roche, Cat. No. 476728001) and polyethylene glycol 8000 (PEG-8000)—precipitated as described previously (45). Additional details about phage PEG precipitation can be found in the supplementary methods. For some experiments, we performed an additional cesium chloride purification step after PEG precipitation as described previously (22).

**Collection of wound swabs from human patients**
From 06/2016 to 02/2018, human patients visiting the Stanford Advanced Wound Care Center...
Infection without Pf phage
1) Recognition of bacterial ligands.
2) Bacterial ligands lead to NFκB translocation.
3) NFκB translocation leads to TNF secretion.
4) TNF stimulates bacterial phagocytosis.

Infection with Pf phage
1) Recognition of Pf phage RNA by TLR3-mediated TRIF signaling.
2) TRIF activation leads to type I interferon production.
3) Type I interferon inhibits TNF secretion and bacterial phagocytosis.

**Fig. 6. A model of Pf4-mediated inhibition of TNF and phagocytosis.** Bacterial ligands stimulate TNF production and phagocytosis in BMDCs. Pf4 gets taken up through endocytic pathways, where Pf RNA stimulates TLR3-mediated TRIF signaling, which induces IFN production. IFN inhibits TNF production and phagocytosis.

in Redwood City, California with open wounds were swabbed in duplicate over a 2.2-cm² area using Levine’s technique, using nylon-flocked wet swabs (Copan Diagnostics, Cat. No. 23-600-963). Swabs were collected in PBS and stored in −80°C before transport on dry ice. In the laboratory, the swabs in PBS were thawed and vortexed vigorously for 15 s. The contents were then aliquoted for quantification of *P. aeruginosa* gene and Pf prophage gene PAO717, as detailed below. Patients at the Wound Care Center were also swabbed for confirmation by diagnostic laboratory culture, the swabs in PBS were thawed and vortexed vigorously for 15 s. The contents were then accessed to assess renal function and glycemetic control.

Quantification of Pf phage
As several factors can produce plaques on bacterial lawns (other species of phage, pyocins, host defensins, etc.), we quantitated Pf phage using a qPCR assay as previously described (47). Additional details about Pf phage quantification can be found in the supplementary methods. Pf1 phage level was determined spectrophotometrically by measuring absorption, using the wave-lengths and extinction coefficients as previously described (49).

**In vivo murine full-thickness wound infection model**
Ten-to-twelve-week-old male mice were anesthetized, shaved, and received two dorsal excisional wounds as described previously (47). Mice were inoculated with 40 μl of luminescent bacteria per wound at the indicated doses 24 hours after wounding, and control mice were inoculated with sterile PBS. Mice were imaged daily for luminescent signal on the IVIS Spectrum (Perkin Elmer), the Ami HTX (Spectral Instruments Imaging), or the Lago-X (Spectral Instruments Imaging) at the Stanford Center for Innovation in In Vivo Imaging daily before takedown. Additional details on the surgical procedure and wound processing can be found in the supplementary methods.

**Calculation of IC50 for bacterial wound colonization**
Infection rates in percentage of total wounds inoculated were plotted on the y-axis, whereas inoculation doses were plotted on the x-axis. Inoculation doses were transformed to log form and a nonlinear regression curve fit function was executed using the Graphpad Prism “log(agonist) versus normalized response–Variable slope” function. This function produces a calculated EC50 value, which is equivalent to IC50 in this setting. The function also provides a r² that indicates goodness of fit of the nonlinear regression curve.

**Histology**
Wounds were harvested, bifurcated, fixed in 10% neutral buffered formalin, and embedded in paraffin. H&E staining and analysis of granulation tissue and epithelial gap was performed as previously described (47). CD45 staining was performed as previously described (48). Any sample that did not yield reliable counts due to sample quality was excluded from the analysis for that particular variable. All scoring was performed by an independent pathologist.

**Mouse wound, spleen, and lymph node harvesting and immunophenotyping**
Mouse wounds were excised with scissors, cut into pieces, and placed in RPMI containing 0.025 mg/ml Liberase TM Research Grade (Roche, Cat. No. 540127001), 50 μM 2-mercaptoethanol (Sigma, Cat. No. M3148), and 20 μM HEPEs (Teknova, Cat. No. 101446-740). Tissue was digested at 37°C for 2 hours, homogenized through a sterile 1-ml syringe (BD Biosciences, Cat. No. 3098659), and passed through a 70-μm cell strainer (Fisherbrand, Cat. No. 22063548). Cells were washed twice with PBS at 300g for 5 min and processed according to the flow cytometric protocol. Murine spleen and lymph node were harvested and treated as previously described (49). Cells were counted and used for assays as described.

**Flow cytometry**
For immunophenotyping, cells were lifted with cold PBS and processed according to our flow cytometric protocol using the antibodies found in table S3. Cells were processed and stained and flow cytometry was performed as previously described (49). Analysis for this project was done on LSR II (BD Biosciences) instruments in the Stanford Shared FACS Facility. Cells were divided into different leukocyte populations according to the gating schemes depicted in figs. S3, S12, and S13.
Inoculation with $7.5 \pm 2.5 \times 10^2$ CFU/ml PAO1 in mice vaccinated against CoaB coat protein, plating. Supernatant containing non-adherent

Harvested for use between days 7 and 9 after changing 5 and 7 days after plating. BMDCs were three days after plating, and 50% of media was fresh media containing cytokines was added

Cat. No. 14-8983-62). An equivalent amount of supplemented with 20 ng/ml of M-CSF (Invitrogen, 2 ng/ml of recombinant mouse IL-4 (Invitrogen, (STEMCELL Technologies, Cat. No. 78206.1) and (BMDCs), the media was supplemented with 10 μg/ml of IgG mAb clones 1 to 3 and the IgM clone 4, or PBS, immediately before bacterial inoculation. For some assays, 5 μg/ml of TruStain FcX (anti-mouse CD16/32) Fc block (BioLegend, Cat. No. 101320) was added before addition of the anti-CoaB mAbs.

To assess phagocytosis using pHrodo-labeled particles, cells were seeded at a density of $5 \times 10^5$ cells/well in a 96-well plate and stimulated with coated $1 \times 10^8$ Pf phage/ml, unless noted otherwise, for 2 hours at 37°C. Next, cells were incubated with 10 μg/ml of pHrodo Red E. coli BioParticles (Invitrogen, Cat. No. P35361) for 1 hour at 37°C, 5% CO₂ or for 1 hour at 4°C for adsorption controls. Cells were harvested by lifting with cold PBS and repeated pipetting, washed twice with flow cytometry buffer, and stained for live cells according to the general flow cytometric protocol before acquisition on a flow cytometer. For TNF rescue experiments, cells were stimulated with coated Pf phage and 25 ng/ml of TNF (Invitrogen, Cat. No. 34-8321-63) for 3 hours at 37°C before addition of the E. coli pHrodo particles.

Generation of murine bone marrow-derived phagocytes

Femurs were isolated from male or female mice of >10 weeks of age and bone marrow was flushed out using a 27 G PrecisionGlide needle (BD Biosciences, Cat. No. 303109). Cells were plated at $1 \times 10^5$ cells in 10 ml of media per Petri Dish (Fisherbrand, Cat. No. FB0875711). To generate bone marrow-derived dendritic cells (BMDCs), the media was supplemented with 10 ng/ml of recombinant mouse granulocyte-macrophage colony-stimulating factor (GM-CSF) (STEMCELL Technologies, Cat. No. 78206.1) and 2 ng/ml of recombinant mouse IL-4 (Invitrogen, Cat. No. 14-8041-62). To obtain bone marrow-derived macrophages (BMDMs), media was supplemented with 20 ng/ml of M-CSF (Invitrogen, Cat. No. 14-8983-62). An equivalent amount of fresh media containing cytokines was added three days after plating, and 50% of media was changed 5 and 7 days after plating. BMDCs were harvested for use between days 7 and 9 after plating, whereas BMDMs were used on day 6 after plating. Supernatant containing non-adherent cells was collected, and adherent cells were lifted by the addition of 10% Accutase in Accutase for 5 min. Both cell populations were pooled to obtain a mixture of mature and immature cells, washed in fresh media at 300 g for 5 min, and prepared for experiments.

Isolation and preparation of human immune cells

Human PBMCs were isolated from an LRS chamber obtained from a healthy donor from the Stanford Blood Center using Ficoll-Paque PLUS (GE Healthcare, Cat. No. 17-4440-02) or lymphoprep (Stemcell technologies, Cat no. 07851) according to the manufacturer’s instructions. Monocytes were isolated using the EasySep Human Monocyte Enrichment Kit according to the manufacturer’s instructions (STEMCELL Technologies, Cat. No. 19058). If differentiation into macrophages was required, monocytes were seeded at a concentration of $6.6 \times 10^5$ cells per ml and 25 ng/ml of recombinant GM-CSF (STEMCELL Technologies, Cat. No. 78206.1) was added to the cultures. Medium was supplemented every 2 to 3 days. After 7 to 9 days, macrophages were harvested by pipetting and the adherent cells were collected by subsequent treatment with 10% Accutase in Accutase. Cells were then centrifuged and re-suspended in RPMI, to be used in assays as described. U937 cells (ATCC CRL-1593.2) were differentiated into macrophages by overnight stimulation with 100 ng/ml Phorbol 12-myristate 13-acetate (Sigma, Cat. No. P1355).

Immune-cell culture conditions

Mouse BMDCs and human PMBCs and other immune cells were cultured in RPMI with 10% FBS, 100 IU of penicillin, 100 μg/ml of streptomycin, and 1 mM sodium pyruvate. Mouse BMDMs were cultured in RPMI with 10% FBS, 100 IU of penicillin, 100 μg/ml of streptomycin, and 20 ng/ml M-CSF (Invitrogen, Cat. No. 14-8983-62). Mouse BMDCs and BMDMs were cultured on sterile plates not treated for tissue culture (Falcon), whereas all other cells were cultured on sterile tissue culture-treated plates treated for tissue culture (Falcon). Cells were cultured at 37°C, 5% CO₂ in a 90% humidified atmosphere.

Phagocytosis assays

To determine phagocytic capacity with live bacteria, gentamicin protection assays and phagocytic killing assays were performed as described previously (50). BMDMs were inoculated with multitude of infection (MOI) of 50, whereas BMDCs and human phagocytes were inoculated with MOI = 10. For the assays with anti-CoaB mAbs, the seeded BMDMs were treated with 10 μg/ml of IgG mAb clones 1 to 3 and the IgM clone 4, or PBS, immediately before bacterial inoculation. For some assays, 5 μg/ml of TruStain FcX (anti-mouse CD16/32) Fc block (BioLegend, Cat. No. 101320) was added before addition of the anti-CoaB mAbs.

To assess phagocytosis using pHrodo-labeled particles, cells were seeded at a density of $5 \times 10^5$ cells/well in a 96-well plate and stimulated with coated $1 \times 10^8$ Pf phage/ml, unless noted otherwise, for 2 hours at 37°C. Next, cells were incubated with 10 μg/ml of pHrodo Red E. coli BioParticles (Invitrogen, Cat. No. P35361) for 1 hour at 37°C, 5% CO₂, or for 1 hour at 4°C for adsorption controls. Cells were harvested by lifting with cold PBS and repeated pipetting, washed twice with flow cytometry buffer, and stained for live cells according to the general flow cytometric protocol before acquisition on a flow cytometer. For TNF rescue experiments, cells were stimulated with coated Pf phage and 25 ng/ml of TNF (Invitrogen, Cat. No. 34-8321-63) for 3 hours at 37°C before addition of the E. coli pHrodo particles.

Murine leukocyte stimulation with bacterial pathogen-associated molecular patterns (PAMPs) and bacteriophage

Cells were seeded at a density of $5 \times 10^5$ cells/well in a 96-well plate and stimulated with various stimuli, including coated 100 μg/ml alginate, $1 \times 10^5$ phage particles/ml purified Pf6 from the PAO1 strain, Pf from the PAK strain or Fdl phage.
from *E. coli*, preparations from the PA01Δp4 strain, soluble 1 μg/ml LPS, or equivalent volumes of PBS as control. Cells were incubated at 37°C, 5% CO₂ for the indicated time points. Next, cells were centrifuged for 300g for 5 min, and supernatant was removed for protein analysis.

**TNF enzyme-linked immunosorbert assay (ELISA)**

To measure TNF protein levels in cell culture supernatants, we used an adapted version of the eBioscience protocol ELISA protocol using antibody pairs. The coating antibody was used at 2.5 μg/ml and the detection antibody was used at 1.25 μg/ml. Carrier-free recombinant mouse TNF (Invitrogen, Cat. No. 34-8321-63) was used for the standard. Absorbance was read on a Spark microplate reader (Tecan).

**TNF detection in human immune cells in response to bacterial PAMPs and Pf phase**

Isolated human monocytes were resuspended in RPMI supplemented with MEM non-essential amino acids (HyClone, Cat. No. SH30238.01), sodium pyruvate, penicillin–streptomycin, 200 mM L-glutamine, 50 mM 2-mercaptoethanol, and 10% heat-inactivated FBS. Monocytes (5 × 10⁶ cells/well) were seeded in a 24 well plate and stimultated for 15 hours with 10⁹ phage particles. Subsequently, 1 μg/ml of LPS was added for 4 hours. Supernatants were collected and used for cytokine profiling through Luminex by The Human Immune Monitoring Center at Stanford University.

**SDS–polyacrylamide gel electrophoresis (PAGE) of Pf4 phase preparations**

PEG-purified phase, crude supernatant from a PA01 culture, and supernatant from a PEG precipitation were assessed via SDS–PAGE electrophoresis using previously described protocols (22).

**Tnf mRNA quantitative PCR**

To quantify *Tnf* mRNA levels, BMDCs were cultured in a 24-well plate at a density of 0.5 × 10⁶ cells/well and left unstimulated or stimulated with 100 μg/ml of alginate or 1 μg/ml of LPS and/or 1 × 10⁹ Pf4/ml for the indicated time periods at 37°C, 5% CO₂. Cells were lysed with TRI Reagent (Sigma, Cat. No. 93289) before RNA isolation. RNA was isolated as previously described (57). Eight hundred nanograms of RNA from all samples was converted into cDNA using High-Capacity cDNA Reverse Transcription Kit (Applied Biosystems, Cat. No. 4368814) per the manufacturer’s instructions. TNF mRNA levels were quantified in a qPCR reaction containing 1× PowerUp SybrGreen (Applied Biosystems, Cat. No. A25776), 0.2 μM primers, and 40 ng of cDNA. See table S2 for the primers used. Reactions were run on a StepOnePlus Real-Time PCR system (Applied Biosystems, Cat. No. 4376600) using a modified Standard Program (50°C for 2 min; 95°C for 2 min.; 40 × (95°C for 10 s; 60°C for 25 s); melt curve). Relative fold expression was calculated using the ΔΔCt method with beta-actin (*Actb*) as the housekeeping gene.

**NF-κB nuclear translocation immunoblot**

BMDCs were seeded in 6-well plates at 1.5 × 10⁶ cells/well and stimulated with coated 100 μg/ml alginate and 5 × 10⁹ Pf4/ml for 30 min. Immunoblotting was then performed as previously described (52).

**NF-κB luciferase reporter assay**

RAW264.7 cells stably transfected with a NF-κB-dependent luciferase reporter (33) were plated at 5 × 10⁶ cells/well in Dulbecco's modified Eagle medium (DMEM) (HyClone, Cat. No. SH3024301) with 10% FBS, penicillin, and streptomycin in a 96-well tissue culture plate and incubated overnight at 37°C, 5% CO₂ to adhere. Media was removed and fresh media containing 1 μg/ml of LPS, 100 μg/ml of alginate, or an equivalent volume of PBS was added, followed immediately by the addition of 1 × 10⁹ Pf4/ml of purified Pf4 or an equivalent dilution of a phage purification prep from the PA01Δp4 strain. Cells were incubated for 9 hours at 37°C 5% CO₂ before lysis with Bright-Glo Reagent (Promega, Cat. No. E2610) and detection of relative light units of luminescence using a Spark microplate reader (Tecan). Values were normalized to relative light units of unstimulated cells.

**Intracellular TNF cytokine staining**

BMDCs were seeded at 2 × 10⁶ cells/well in a 24-well plate coated with 1 × 10⁹ Pf4/ml purified Pf4 and cultured for 15 hours at 37°C. Cells received a stimulus of 1 μg/ml of LPS and 3.0 μg/ml of Brefeldin A (Invitrogen, Cat. No. 00-4506) for 4 hours at 37°C to stimulate TNF production. Cells were lifted by cold PBS and processed according to our flow cytometric protocol, with some additional steps. After washing in flow cytometry buffer (3% FBS and 1 mM EDTA), cells were fixed with IC fixation buffer (Invitrogen, Cat. No. 00-8222-49), permeabilized with Permeabilization buffer (Invitrogen, Cat. No. 00-8333-56), and stained with 2 μg/ml of anti-mouse TNF antibody according to the manufacturer’s instructions. Cells were washed twice in flow cytometry buffer before acquisition.

**Type I IFN protein reporter assay**

ISRE-L929 cells (54) were seeded at a density of 1 × 10⁶ cells/well in a 96-well tissue culture plate in DMEM with 10% FBS, penicillin, and streptomycin, and cultured overnight at 37°C, 5% CO₂ to allow them to adhere. Media was removed, and cells were incubated with supernatant from BMDCs or BMDCMs stimulated with LPS or alginate and Pf4, as described elsewhere in the Methods section. To quantify type I IFN production, a twofold serially diluted standard curve of recombinant mouse IFN-β (BioLegend, Cat. No. 581309), starting at 1000 pg/ml, was used. ISRE-L929 cells were incubated with cells or protein standard for 9 hours at 37°C, 5% CO₂, and cell supernatant was removed and Bright-Glo Reagent (Promega, Cat. No. E2610) was added to lyse cells for 2 min. Relative luminescent units were detected using a Spark microplate reader (Tecan).

**HEK-Blue hTLR3 cells treated with whole Pf phase particle**

HEK-Blue hTLR3 cells (Invivogen, Cat. No. hkb-hitr3), cells in which native Tlr3 is expressed in endosomes, were grown to a confluency of at least 85%, as described by the manufacturer. Subsequently, cells were harvested and diluted to a concentration of 5 × 10⁶ cells/ml in HEK-Blue growth medium (DMEM supplemented with 10% FBS, penicillin–streptomycin, 100 μg/ml of noromycin, 30 μg/ml of blasticidin (Fisher Scientific, Cat. No. MT30100RB), 100 μg/ml of zeocin (Invivogen, Cat. No. ant-zn-1), 10 mM HEPES, and 10 μM MG-132 (Selleck Chemicals, Cat. No. S2619)). Cells were seeded in a flat-bottom 96-well plate at a final concentration of 5 × 10⁶ cells/well, and 20 μl of stimulus was added (PBS or 10⁹ phage/ml). Cells were incubated for 16 hours at 37°C with 5% CO₂, after which the media was replaced with 100 μl of HEK-Blue growth media. After 8 hours incubation, 100 μl of HEK-Blue Detection media (Invivogen, Cat. No. nb-det2) was added. After a final 16-hour incubation, discoloration of the media was measured at 620 nm using a Spark microplate reader (Tecan).

**Preparation of pure phage RNA from ssDNA**

In order to prepare pure phage RNA, ssDNA from PF1 (used because of higher purification yield) was extracted as previously described (55). Individual phage genes were amplified using specific primers (1 μM), to which a T7 promoter was added on the forward primer, using Ecomeq (Lucigen, Cat. No. 30035-1). See table S2 for the primer sequences. The PCR products were purified prior to RNA conversion using silicon column-bound guanidine thiocyanate purification. More details on this procedure can be found in the supplementary methods for more details. RNA was transcribed according to the manufacturer instructions (HiScribe T7 High Yield RNA Synthesis kit, New England Biosciences, Cat. No. E20408). RNA products were purified as described for the PCR products. Subsequently, the purified RNA was used in a HEK-Blue hTLR3 assay. Briefly, 20 μl of each RNA product (10 ng/μl) was added to wells in triplicate in a 96-well plate. Subsequently, 5 × 10⁶ HEK-Blue hTLR3 cells were added to each well, as described by the manufacturer, in HEK-Blue detection medium. After 16 hours of stimulation, discoloration of the media was measured at 620 nm using a Spark microplate reader (Tecan).

**Phage RNA production in a human monocyte cell line**

The human monocyte cell line U937 was grown to a concentration of 2 × 10⁶ cells/ml in RPMI with penicillin and streptomycin. Subsequently,
cells were transferred to RPMI with penicillin and streptomycin to a final concentration of 10^5 cells/ml. 10^5 cells/ml were stimulated with 10^9 phage. After 24 hours, the whole-cell suspension was transferred to 1 ml of TriReagent (Sigma) and stored overnight at -80°C. Total RNA was extracted using a RNeasy Minikit, as described by the manufacturer (Qiagen, Cat. No. 74106). Genomic DNA was removed from the total RNA using DNase I and incubating at 37°C for 15 min, followed by inactivation at 85°C for 15 min. Subsequently, cDNA was prepared using the RevertAid RT Reverse Transcription Kit (Thermo Fisher, Cat No. K1690). Additionally, as a negative control, the remaining RNA was diluted in the same ratio as for the cDNA synthesis. Both cDNA and RNA were used for qPCR, using SensiFast Sybr HIROX (Bioline, Cat no. BIO-92005), a 1:1 template, and forward and reverse primers (400 nM each). In total, 45 cycles were run (5 s at 95°C, 30 s at 60°C, 20 s at 72°C). See table S2 for primer sequences.

Fluorescent labeling of Pf phage

Purified Pf4 preparations were labeled with Alexa Fluor 488-labeled TFP ester (Molecular Probes, Cat. No. 107) following the manufacturer’s protocol. Following labeling, purified Pf4 viral particles were separated from unincorporated dye using PD10 Sephadex G-25 desalting columns (GE Healthcare, Cat. No. 17085101) according to the manufacturer’s instructions. The conjugate was then quantitated for Pf concentration using qPCR and appropriately diluted in PBS.

Uptake assays of labeled Pf phage by murine leukocytes

BMDCs or BMDMs were seeded at a density of 5 × 10^6 cells/well in a 96-well plate or 5 × 10^5 cells/well in a 24-well plate. Splenocytes and lymph node cells were seeded in 24-well plates at 1.5 × 10^6 cells/well and incubated with 1 × 10^9 Pf4/ml labeled Pf4 or soluble 1 × 10^9 Pf4/ml labeled Pf4 for 3 hours. Cells were washed, fixed with 10% neutral buffered formalin, and prepared for immunofluorescent microscopy according to the procedure outlined in the supplementary methods.

Transmission electron microscopy

For microscopy analysis, cells were seeded on glass 12-mm #1.5H precision coverslips (Thorlabs, Cat. No. KG15CH) coated with poly-L-lysine (Sigma, Cat. No. P8920) or 0.2% poly-L-ornithine (Sigma, Cat. No. P3655) according to the manufacturer’s instructions at a density of 1 × 10^6 cells/well in a 6-well plate. Cells were either stimulated with coated 1 × 10^9 Pf4/ml labeled Pf4 or soluble 1 × 10^9 Pf4/ml labeled Pf4 for 3 hours. Cells were prepared using a RNeasy Minikit, as described by the manufacturer. As an alternative method, the cells were噱熏 with 5×10^4 cells/well in a 96-well plate or 5×10^5 cells/well in a 24-well plate. Splenocytes and lymph node cells were transferred to RPMI with penicillin and streptomycin to a final concentration of 10^5 cells/ml. 10^6 cells/ml were stimulated with 100 µl/well of 1% BSA + 0.1% goat serum in PBS for 1 hour at room temperature. Blocking buffer was removed, and 100 µl/well of immunized mouse serum samples were then added diluted 1:10 in blocking buffer. Samples were incubated overnight at 4°C or 2 hours at room temperature. Plates were then incubated with 100 µl/well of 1-Step Ultra TMB ELISA Substrate and 100 µl/well of 0.05% Tween-20 in PBS between steps.

Anti-CoaB mAb purification

Hybridomas produced by immunizing mice with KLH-conjugated CoaB peptide and selecting monovalent antibodies, were generated by ImmunoPrecise Antibodies LTD (Victoria, Canada) upon commission. Binding affinities of the resulting antibodies were confirmed by indirect ELISA using peptide-BSA, free BSA, or human transferrin as irrelevant antigen control. For hybridoma growth conditions, see the supplementary methods. After adaptation, cells were expanded, and culture supernatant was collected. Supernatants from each culture were centrifuged at 2000g for 15 min at 4°C to remove debris and insolubles. Medium was filtered through a 0.22-µm filter and loaded into a Protein A Resin FF PrepPacked column (Genescript, Cat. No. L00680-51) according to the manufacturer’s instructions. Purified antibody clones were concentrated using a 100-kDa cut-off centrifugal filters (EMD Milipore, Cat. No. UFC910024) and resuspended in 0.01 M PBS (pH 7.4). The protein concentration of purified mAbs was determined by BCA assay (Pierce Thermo Scientific,

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tests were two-tailed. Depicted are means with error bars; statistical tests, Mann–Whitney tests, and Fisher’s exact tests were two-tailed. Depicted are means with error bars; statistical tests, Mann–Whitney tests, and Fisher’s exact tests were two-tailed. Depicted are means with error bars; statistical tests, Mann–Whitney tests, and Fisher’s exact tests were two-tailed. Depicted are means with error bars; statistical tests, Mann–Whitney tests, and Fisher’s exact tests were two-tailed. Depicted are means with error bars; statistical tests, Mann–Whitney tests, and Fisher’s exact tests were two-tailed. Depicted are means with error bars; statistical tests, Mann–Whitney tests, and Fisher’s exact tests were two-tailed. Depicted are means with error bars; statistical tests, Mann–Whitney tests, and Fisher’s exact tests were two-tailed. Depicted are means with error bars; statistical tests, Mann–Whitney tests, and Fisher’s exact tests were two-tailed. Depicted are means with error bars; statistical tests, Mann–Whitney tests, and Fisher’s exact tests were two-tailed. Depicted are means with error bars; statistical tests, Mann–Whitney tests, and Fisher’s exact tests were two-tailed. Depicted are means with error bars; statistical tests, Mann–Whitney tests, and Fisher’s exact tests were two-tailed. Depicted are means with error bars; statistical tests, Mann–Whitney tests, and Fisher’s exact tests were two-tailed. Depicted are means with error bars; statistical tests, Mann–Whitney tests, and Fisher’s exact tests were two-tailed. Depicted are means with error bars; statistical tests, Mann–Whitney tests, and Fisher’s exact tests were two-tailed. Depicted are means with error bars; statistical tests, Mann–Whitney tests, and Fisher’s exact tests were two-tailed. Depicted are means with error bars; statistical tests, Mann–Whitney tests, and Fisher’


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Data and materials availability: All data are available in the main text or the supplementary materials.

SUPPLEMENTARY MATERIALS

www.sciencemag.org/content/363/6434/eaat9691/suppl/DC1
Materials and Methods
Figs. S1 to S14
Tables S1 to S3
Movie S1

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Phage subverts immune response

*Pseudomonas aeruginosa* (*Pa*) is a multidrug-resistant Gramnegative bacterium commonly found in health care settings. *Pa* infections frequently result in considerable morbidity and mortality. Sweere *et al.* found that a type of temperate filamentous bacteriophage that infects and integrates into *Pa* is associated with chronic human wound infections. Likewise, wounds in mice colonized with phage-infected *Pa* were more severe and longer-lasting than those colonized by *Pa* alone. Immune cell uptake of phage-infected *Pa* resulted in phage RNA production and inappropriate antiviral immune responses, impeding bacterial clearance. Both phage vaccination and transfer of antiphage antibodies were protective against *Pa* infection.

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