Y
ersinia pestis is a highly virulent pathogen and the causative agent of bubonic, septicemic, and pneumonic plague. Primary pneumonic plague caused by inhalation of respiratory droplets contaminated with Y. pestis is nearly 100% lethal within 4 to 7 days without antibiotic intervention. Pneumonic plague progresses in two phases, beginning with extensive bacterial replication in the lung with minimal host responsiveness, followed by the abrupt onset of a lethal proinflammatory response. The precise mechanisms by which Y. pestis is able to colonize the lung and survive two very distinct disease phases remain largely unknown. To date, a few bacterial virulence factors, including the Ysc type 3 secretion system, are known to contribute to the pathogenesis of primary pneumonic plague. The bacterial GTPase BipA has been shown to regulate expression of virulence factors in a number of Gram-negative bacteria, including Pseudomonas aeruginosa, Escherichia coli, and Salmonella enterica serovar Typhi. However, the role of BipA in Y. pestis has yet to be investigated. Here, we show that BipA is a Y. pestis virulence factor that promotes defense against early neutrophil-mediated bacterial killing in the lung. This work identifies a novel Y. pestis virulence factor and highlights the importance of early bacterial/neutrophil interactions in the lung during primary pneumonic plague.

KEYWORDS  plague, pneumonic plague, Yersinia, Yersinia pestis, bacterial GTPases, pathogenesis

Y
ersinia pestis is a highly lethal Gram-negative bacterium that causes bubonic, septicemic, and pneumonic plague. Inhalation of aerosols and respiratory droplets contaminated with Y. pestis results in primary pneumonic plague, a rapidly progressing pneumonia that is nearly 100% fatal without antibiotic treatment (1). Due to its ease of transmission by aerosol and potential to cause mass casualties in outbreak and bioterrorism scenarios, Y. pestis is categorized as a Tier 1 Select Agent by the Centers of Disease Control and Prevention (CDC), and a Category A Priority Pathogen by the National Institutes of Health (NIH). While often considered a disease of ancient times, plague continues to be a modern threat, and Y. pestis is endemic on every continent except Antarctica and Australia (2). For example, a 2017 plague epidemic in Madagascar saw approximately 2,600 suspected or confirmed cases of plague, of which roughly 1,700 were pneumonic plague (3).

Essential to Y. pestis virulence is the Ysc type 3 secretion system (T3SS). The Yersinia T3SS is a “needle-like” apparatus that directly injects bacterial effector proteins called Yersinia outer proteins, or Yops, into the cytosol of target host cells. Once injected, the Yop effectors function to prevent phagocytosis and limit innate immune signaling (4). Aside from the T3SS, few Y. pestis virulence factors have been shown to contribute to disease progression of pneumonic plague. As a result, there is a paucity of information...
regarding how *Y. pestis* is able to establish infection and replicate to high numbers in the lung prior to the onset of host inflammatory responses.

BPI-inducible protein A (BipA), also known as TypA, was first identified in *Salmonella enterica* serovar Typhimurium, and its expression was shown to be induced in the presence of human bactericidal permeability increasing protein (BPI) (5), an antimicrobial peptide (AMP) produced primarily by granulocytes (6, 7). BipA is classified as a translational GTPase with homology to elongation factor G (EF-G) (8, 9) that demonstrates differential ribosomal subunit binding in response to relative guanosine-5′-triphosphate (GTP) and guanosine tetraphosphate (ppGpp) concentrations (10). BipA is highly conserved, and is implicated in regulating virulence mechanisms at the translational level in a number of bacterial species, including *Escherichia coli*, *S. enterica*, and *Pseudomonas aeruginosa* (11). In *E. coli*, BipA plays a role in K5 capsule production (12), flagella-mediated motility (13, 14), BPI resistance (13), and attachment and effacement to epithelial cells (14). BipA contributes to macrophage uptake and survival in *S. enterica* serovar Typhi (15), and BPI resistance in *S. enterica* Typhimurium (16). In *P. aeruginosa*, BipA is involved in bacterial virulence in amoeba and nematode models, as well as adhesion, biofilm formation, resistance to phagocytosis, antibiotic resistance, and expression of the T3SS (17). While BipA has been shown to contribute to virulence mechanisms in these pathogens, there has been minimal characterization beyond initial in vitro analysis. The role of BipA in *Y. pestis* has yet to be investigated. In this study, we show that deletion of BipA results in decreased bacterial survival upon challenge with human BPI and primary human neutrophils. Further, we show that BipA facilitates bacterial survival in the lung, and contributes to lethality in a murine infection model of pneumonic plague. This work positions BipA as a novel *Y. pestis* virulence factor that promotes resistance to early neutrophil-mediated bacterial killing in the lung.

**RESULTS**

BipA is involved in the *Y. pestis* response to challenge with BPI. Previous work (18) identified 405 bacterial genes that were differentially expressed in *Y. pestis* isolated from the murine lung during infection compared to broth-grown bacteria. Of these, the gene *bipA* (or *YPO0026*) encoding the bacterial translational GTPase BipA was up-regulated 2.3-fold during pneumonic plague. Expression of BipA is known to be induced in the presence of BPI (5), an AMP released in primary granules by neutrophils (19–21). BipA, in turn, has been implicated in regulating expression of proteins, including known virulence factors, in a number of pathogens (8, 9, 11, 14, 17, 22–25). We sought to determine if the absence of BipA impacted expression of BPI-induced proteins in *Y. pestis*. We generated a strain of *Y. pestis* CO92 lacking *bipA* (*D*bipA) and evaluated total protein expression by liquid chromatography-tandem mass spectrometry (LC-MS/MS) in wild type and *D*bipA strains in the logarithmic phase of broth-grown culture and during incubation with BPI. This analysis revealed a number of proteins that were differentially expressed (at least 2-fold difference) in *D*bipA *Y. pestis* compared to wild-type *Y. pestis* under both conditions (Table S1 and S2 in the supplemental material), as well as altered regulation of an additional set of proteins in the presence of BPI. For example, we observed increased expression of the zinc import ATP-binding protein ZnuC and decreased expression of Der, putative type 6 secretion system components, and various uncharacterized proteins in *D*bipA *Y. pestis* compared to wild-type *Y. pestis* under both conditions (Table S1 and S2 in the supplemental material), as well as altered regulation of an additional set of proteins in the presence of BPI. For example, we observed increased expression of the zinc import ATP-binding protein ZnuC and decreased expression of Der, putative type 6 secretion system components, and various uncharacterized proteins in *D*bipA *Y. pestis* compared to wild-type *Y. pestis* under both conditions. During incubation with BPI, we saw differential regulation of additional proteins, including ClpB, YscJ, and a number of other uncharacterized proteins. Deletion of BipA did not appear to effect transcription of *znuC*, *YPO0502*, *psaA*, and *yoph*, each of which encode proteins that showed altered expression in the absence of BipA (Fig. S1). Similar to its function in other bacteria, our data suggest that BipA contributes to the bacterial response to challenge with BPI.

BPI is released from the granules of neutrophils and has high affinity for the lipid A component of bacterial lipopolysaccharide (LPS) (19). Binding of BPI to LPS inhibits the endotoxic activity of LPS, facilitates opsonization of bacteria, and has potent
bactericidal activity (6, 26, 27). BipA has been shown to contribute to bacterial resistance to BPI in *E. coli* and *S. enterica* Typhi (13, 16). We sought to determine if BipA contributes to *Y. pestis* resistance to BPI. To this end, we incubated wild-type *Y. pestis* strain CO92, ΔbipA, or the ΔbipA::bipA complement strain with a sub-MIC of BPI (40 µg/ml) at 37°C for 8 h, and enumerated CFU of each strain at 2, 4, and 8 h postinoculation (hpi). While deletion of *bipA* had no impact on *Y. pestis* growth in minimal medium (Fig. 1A), we observed a significant decrease in survival of *Y. pestis* lacking BipA by 8 hpi compared to wild-type and BipA-complemented *Y. pestis*, indicating a role for BipA in bacterial resistance to BPI (Fig. 1B). To determine if ΔbipA was attenuated in the presence of antimicrobial peptides in general, we also evaluated bacterial survival upon challenge with polymyxin B and LL-37. Deletion of *bipA* did not impact *Y. pestis* sensitivity to sub-MICs of polymyxin B or LL-37, suggesting that BipA may not contribute to resistance to all AMPs (Fig. 1C and D). Evaluating the MIC of each antimicrobial peptide for all three strains showed no difference in AMP resistance (Table S3). These results indicate that BipA contributes to the bacterial response to challenge with BPI.

**BipA contributes to resistance to neutrophil-mediated bacterial killing.** BPI is primarily produced by granulocytes such as neutrophils (6). We therefore sought to determine whether BipA contributes to bacterial defense against neutrophils. We inoculated primary human neutrophils with wild-type, ΔbipA, or ΔbipA::bipA *Y. pestis* CO92, and monitored bacterial survival for 8 h. We observed significantly decreased survival of ΔbipA *Y. pestis* compared to wild-type and ΔbipA::bipA *Y. pestis* in the presence of primary human neutrophils (Fig. 2A), indicating that BipA contributes to bacterial resistance to neutrophil-mediated bacterial killing in vitro. To determine if BipA contributes to resistance to killing by professional phagocytes in general, we also evaluated bacterial survival during incubation with primary human alveolar macrophages (hAM) and immortalized murine alveolar macrophages (MH-S) (Fig. 2B and C). *Y. pestis* encounters alveolar macrophages early during infection, and these initial host/
pathogen interactions are important to establishing infection in the lung. We did not observe a significant difference in $\Delta$D~bipA survival compared to wild type or complemented $\Delta$D~bipA during incubation with macrophages, suggesting that the contribution of BipA to bacterial survival may be specific to neutrophils. Gentamicin protection assays also revealed no difference in the ability of primary human neutrophils or MH-S cells to phagocytose wild-type, $\Delta$D~bipA, or $\Delta$bipA::D~bipA Y. pestis strains (Fig. S2).

The Yersinia pestis T3SS (encoded on the pCD1 plasmid) inhibits phagocytosis of bacteria, limits innate immune signaling (4, 28), and is essential for virulence. The Yersinia T3SS is critical for defense against phagocytosis by neutrophils (29–31). BipA was shown to regulate expression of the T3SS of P. aeruginosa (17), indicating that BipA-related phenotypes may result from the inability to effectively target host cells for T3SS. Though we did not see major differences in expression of the T3SS in the absence of BipA, we still sought to determine if reduced T3SS could explain increased $\Delta$D~bipA Y. pestis sensitivity to killing by primary human neutrophils. Initial efforts to quantify T3SS in primary neutrophils were unsuccessful, likely due to the rapid turnover and sensitivity of primary neutrophils to manipulation in culture. As an alternative approach, we incubated primary human neutrophils with wild-type, $\Delta$D~bipA, $\Delta$D~bipA::D~bipA, $\Delta$pcD1 lacking the T3SS, or $\Delta$D~bipA $\Delta$pcD1 Y. pestis CO92 and monitored bacterial survival. We observed a similar decrease in the viability of $\Delta$pcD1 and $\Delta$D~bipA $\Delta$pcD1 Y. pestis compared to the $\Delta$D~bipA strain in the presence of neutrophils (Fig. 3A). This finding does not confirm a link between BipA and T3SS, but does indicate it is possible that BipA-mediated resistance to neutrophil-mediated killing is due to its regulation of the T3SS. To investigate this further, we sought to determine if the absence of BipA impacted the effects of the T3SS on target neutrophils. The Y. pestis T3SS was recently shown to inhibit degranulation of neutrophils via the combined activity of YopE and YopH (30, 32). We therefore investigated the ability of Y. pestis $\Delta$D~bipA to inhibit neutrophil degranulation compared to wild-type Y. pestis as a read-out of T3SS. Primary human neutrophils were inoculated with wild-type, $\Delta$D~bipA, or $\Delta$pcD1 Y. pestis using an MOI of 10:1 bacteria to host cells, and neutrophil degranulation was evaluated by flow cytometric detection of CD63-PE-Cy7 as a marker of primary granule release (21). We observed no difference in CD63 fluorescence intensity between wild-type and $\Delta$D~bipA Y. pestis, indicating that loss of BipA did not impact the ability of Y. pestis to inhibit neutrophil degranulation by the T3SS (Fig. 3B). As expected, we observed increased CD63 fluorescence intensity consistent with neutrophil degranulation after incubation with $\Delta$pcD1 Y. pestis lacking the T3SS effectors and machinery. Gentamicin protection assays also showed similar resistance of $\Delta$D~bipA Y. pestis (compared to both wild-type and BipA-complemented Y. pestis) to phagocytosis by neutrophils (Fig. S2), suggesting similar T3SS function. Together, these data show that BipA likely promotes Y. pestis resistance to neutrophil-mediated killing via a T3SS-independent mechanism.
BipA promotes Y. pestis virulence and bacterial survival in a murine model of primary pneumonic plague. We next sought to determine if BipA contributes to virulence in vivo during primary pneumonic plague. Groups of 4- to 6-week-old C57BL/6 female mice were inoculated via the intranasal route with 10^3 CFU of wild-type, ΔΔbipA, ΔΔpCD1, or ΔΔbipA ΔΔpCD1 Y. pestis CO92 at an MOI of 1:1 bacteria to host cells. Surviving bacteria counts were determined at 2, 4, and 8 hpi by serial dilution and plating. Significance was calculated using two-way ANOVA; ns, no significant difference; **, P ≤ 0.005; ***, P ≤ 0.0005. (B) Human primary neutrophils were inoculated with wild-type, ΔbipA, or ΔpCD1 Y. pestis CO92 using an MOI of 10:1 bacterial to host cells for 1 h at 37°C. CD63 mean fluorescence intensity was measured by flow cytometry. Significance relative to mock-inoculated neutrophils was calculated using one-way ANOVA with Tukey's multiple-comparison test; ****, P ≤ 0.0001. For both experiments, error bars represent SD. Data are presented as pool of at least two independent experiments.

**FIG 3** Role of the T3SS in BipA-mediated resistance to neutrophils. (A) Human primary neutrophils were inoculated with wild-type, ΔΔbipA, ΔΔpCD1, or ΔΔbipA ΔΔpCD1 Y. pestis CO92 at an MOI of 1:1 bacteria to host cells. Surviving bacteria counts were determined at 2, 4, and 8 hpi by serial dilution and plating. Significance was calculated using two-way ANOVA; ns, no significant difference; **, P ≤ 0.005; ***, P ≤ 0.0005. (B) Human primary neutrophils were inoculated with wild-type, ΔbipA, or ΔpCD1 Y. pestis CO92 using an MOI of 10:1 bacterial to host cells for 1 h at 37°C. CD63 mean fluorescence intensity was measured by flow cytometry. Significance relative to mock-inoculated neutrophils was calculated using one-way ANOVA with Tukey's multiple-comparison test; ****, P ≤ 0.0001. For both experiments, error bars represent SD. Data are presented as pool of at least two independent experiments.

BipA promotes Y. pestis virulence and bacterial survival in a murine model of primary pneumonic plague. We next sought to determine if BipA contributes to virulence in vivo during primary pneumonic plague. Groups of 4- to 6-week-old C57BL/6 female mice were inoculated via the intranasal route with 10^3 CFU of wild-type, ΔΔbipA, or ΔΔbipA::bipA Y. pestis and survival was observed over time. Mice inoculated with ΔΔbipA Y. pestis showed increased survival compared to mice inoculated with wild-type and ΔΔbipA::bipA Y. pestis (Fig. 4A). To determine if BipA contributes to bacterial survival in the lung and/or dissemination to the spleen, bacterial burdens were quantified at 24 hpi (lungs only) and 48 hpi (lungs and spleen). While deletion of bipA did not impact bacterial growth in culture (Fig. 1A), animals inoculated with ΔΔbipA Y. pestis had significantly decreased bacterial burdens in the lung compared to animals infected with wild-type or BipA-complemented Y. pestis strains at both 24 and 48 hpi (Fig. 4B and C). While we did not observe a statistically significant difference in ΔΔbipA Y. pestis bacterial burdens in the spleen (Fig. 4D) compared to wild-type and ΔΔbipA::bipA Y. pestis-infected animals, we failed to detect measurable burdens in 6 out of 15 ΔΔbipA Y. pestis-infected animals, indicating a lack of bacterial dissemination from the lung. These data show that BipA promotes Y. pestis virulence in a murine model of primary pneumonic plague and contributes to bacterial survival in the lung during infection. Importantly, decreased survival of the bacterium was evident at 24 hpi, indicating that BipA likely contributes to early events facilitating survival of Y. pestis in the lung.

Our in vitro data suggest that BipA contributes to bacterial resistance to neutrophil-mediated killing via a mechanism independent of regulation of the T3SS. To determine if this is the case in vivo, we evaluated the T3SS in the lungs at 6 and 12 hpi, when bacterial burdens were roughly equivalent between wild-type and ΔΔbipA strains (Fig. 5A and E). To this end, C57BL/6 mice were inoculated with 5 × 10^4 CFU of wild-type, ΔbipA, or ΔΔbipA::bipA Y. pestis YopE-Bla reporter strains. These strains express wild-type YopE as well as a protein consisting of the first 100 amino acids of YopE fused to a truncated TEM β-lactamase gene (bla), and have been used extensively to detect the T3SS in vitro and in vivo (33–35). Upon incubation of infected cells with the fluorescent Bla substrate CCF2-AM, mammalian cells can be differentiated as Yop-injected (blue fluorescence) or uninjected (green fluorescence). At 6 and 12 hpi, there were no
significant differences in levels of Yop-injected cells in the bronchoalveolar lavage fluid (BALF) between wild-type CO92, ΔbipA, or ΔbipA::bipA Y. pestis infections (Fig. 6B and F). As BipA contributes to bacterial survival upon challenge with neutrophils in vitro, we quantified T3SS into neutrophils specifically. We observed no differences in ΔbipA Y. pestis Yop injection into neutrophils at 6 hpi or 12 hpi compared to wild-type and BipA-complemented Y. pestis (Fig. 5C and G). No difference in neutrophil influx into BALF was observed at 6 or 12 hpi with any strain (Fig. 5D and H). Together, these data indicate that BipA likely does not contribute significantly to the Y. pestis T3SS in vivo.

Neutrophils are responsible for limiting growth of Y. pestis lacking BipA. The finding that Y. pestis lacking BipA was more sensitive to neutrophil-mediated killing in vitro prompted us to investigate whether BipA promotes bacterial resistance to neutrophils in vivo. To test this, groups of mice were inoculated with 10^3 CFU of wild-type, ΔbipA, or ΔbipA::bipA Y. pestis CO92 and bacterial burdens were analyzed in the lung at 24 and 48 hpi. One cohort of ΔbipA-infected mice received 20 μg of anti-mouse Ly-6G antibody (αLy-6G) prior to and at the time of infection to deplete neutrophils (Fig. 6). At both 24 (Fig. 6A) and 48 hpi (Fig. 6B), depletion of neutrophils resulted in complete restoration of ΔbipA Y. pestis growth in the lung. Neutrophil depletion did not significantly enhance bacterial burdens of wild-type Y. pestis at 24 hpi or 48 hpi compared to untreated mice, indicating that ΔbipA Y. pestis is exceptionally susceptible to neutrophil-mediated killing in vivo. These results indicate that neutrophils control ΔbipA Y.
**DISCUSSION**

*Y. pestis* employs a potent arsenal of virulence factors to cause multiple manifestations of plague. While many virulence factors have been characterized, only a few have been shown to contribute to the pathogenesis of pneumonic plague specifically. The precise mechanisms underlying biphasic disease progression of pneumonic plague are unknown, and it is likely that a number of bacterial and host mediators of pathogenesis are yet to be identified. Antibiotic-resistant strains of *Y. pestis* have emerged in plague-endemic foci (36), warranting investigation of putative virulence factors that contribute to disease and might be exploited therapeutically. In this study, we investigated the role of a novel virulence factor, BipA, in the pathogenesis of primary pneumonic plague. We show that BipA confers resistance to BPI in vitro and contributes to bacterial resistance to neutrophil-mediated killing, likely via a T3SS-independent mechanism. We further demonstrate that BipA promotes *Y. pestis* survival in lungs and bacterial dissemination to the spleen. Importantly, depletion of neutrophils rescues growth of the ΔbipA *Y. pestis* strain in the lung, confirming the importance of early bacterial-neutrophil interactions in the lung. This is the first time BipA has been evaluated in *Yersinia*, and to our knowledge is the first study to demonstrate the importance of BipA in bacterial virulence in vivo using a murine model of infection.

There is little known about the precise role BipA plays in bacterial translation or...
virulence. Studies from multiple labs using *E. coli* in low temperature conditions have shown that BipA promotes efficient assembly of ribosomal proteins (22, 25). Furthermore, BipA is shown to have binding capabilities to the ribosome and 30S small ribosomal subunit in response to the relative abundance of GTP and ppGpp (9, 10, 23), suggesting a role in bacterial amino acid starvation responses (37). Other work used *in vitro* approaches to show that BipA has a role in regulating critical bacterial virulence mechanisms such as T3SS, adhesion, AMP resistance, motility, capsule expression, and survival and uptake into macrophages (13–15, 17). Further work is needed to determine specifically how BipA contributes to stress responses at the mammalian body temperature and changes in protein expression that we observed here in response to challenge with BPI.

Our data indicate that the presence of BipA may impact the translation of a myriad of proteins in response to BPI, and potentially *in vivo*. These include known proteins such as Der and ZnuC. Der (also known as YphC in *Bacillus subtilis*) is a tandem GTPase functioning in 50S ribosomal subunit biogenesis and is essential for growth in *E. coli* (38, 39). Our proteomic analysis showed a decrease in Der abundance in *Y. pestis* lacking BipA. ZnuC is the ATPase component of the zinc ion ATP-binding cassette (ABC) transporter system encoded by *znuABC*. Our proteomic analysis also suggested that the presence of BipA impacts the expression of a number of hypothetical proteins, including predicted type 6 secretion system (T6SS) components. The Hcp-like component encoded by *YPO0502* is implicated in autoagglutination (40) and has decreased expression in ΔbipA *Y. pestis* upon BPI challenge, according to our proteomic analysis. The precise roles of this and other predicted T6SS components in *Yersinia* have yet to be elucidated, but studies have suggested their roles in environment adaptation (41) and virulence (42). While it is unclear if BipA directly impacts translation of specific proteins, dysregulated expression of important virulence factors and metabolic proteins in its absence likely contribute to decreased viability in the presence of neutrophils and BPI, and may explain the decreased fitness of ΔbipA *Y. pestis in vitro and in vivo*.

The early stage of pneumonic plague has been described as “preinflammatory,” characterized by minimal host responsiveness despite rapid outgrowth of *Y. pestis* in the lung (43). The Ysc T3SS is essential for colonizing and maintaining infection in the lung, as it functions to inhibit phagocytosis (4) and various immune signaling

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**FIG 6** Neutrophils control ΔbipA *Y. pestis in vivo*. C57BL/6 mice (n=14 to 15 per group) were infected with 10⁵ CFU of either wild-type CO92 or ΔbipA *Y. pestis* via the intranasal route. αLy-6G-treated mice were given anti-mouse Ly-6G antibody via tail vein injection at one day prior to infection and on the day of infection. Bacterial burdens in the lung were determined at 24 (A) and 48 (B) hpi by homogenization, serial dilution, and plating. Significance was calculated using two-way ANOVA; ***, P ≤ 0.005; ****, P ≤ 0.0001. Error bars represent SD. Data are presented as a pool of three independent experiments.
processes (44) by primarily targeting alveolar macrophages in the lung early in infection (35). This interaction is further facilitated by adhesins such as Ail, Psa, and Pla that encourage close contact between Y. pestis and host cells targeted for T3SS (45–47). The resulting effect is the creation of localized pockets of immunosuppression. Despite this, it is becoming clear from our lab and others that an early wave of neutrophils can be detected within the first 24 h of pulmonary infection (35, 48) and Y. pestis must overcome this early onslaught. While several virulence factors have been identified and characterized in the later “proinflammatory” stage of pneumonic plague, where increased proinflammatory cytokines are expressed and increased immune cell infiltration occurs (43), it is unclear which play roles in establishing Y. pestis pulmonary infection. Here, we pose BipA as a novel virulence factor that aids in establishing Y. pestis pulmonary infection by contributing to resistance to early neutrophil infiltration into the lungs.

In summary, this work characterizes the role of the Y. pestis translational GTPase BipA in primary pneumonic plague. We show that BipA promotes resistance to neutrophil-mediated killing in a T3SS-independent manner. This effect is likely indirect, due to the impact of BipA on the expression of a number of proteins, including previously identified virulence factors, metabolic genes, and uncharacterized proteins in the face of host challenges during pulmonary infection. Future work will focus on identifying the mechanism of BipA-mediated protein expression and determining how BipA mediates resistance to neutrophil-mediated bacterial killing.

MATERIALS AND METHODS

Bacterial strains. Y. pestis strain CO92, CO92 ∆bipA, and CO92 ∆pCDJ were obtained from the lab of William Goldman (UNC-Chapel Hill). The ∆bipA Y. pestis strain was generated using a modified lambda red recombination described previously (49). Upstream and downstream sequences of bipA were amplified by PCR using the following primers: 5’-TTACTGCATTATGTTTGCAGGCAAT-3’, 5’-GAAG CAGGCTGCTGCAACACGTTTGTGGCTGTTGACCTATCCAGAATTTCTTCTTTGCATTGTTGATACTTAGGGC-3’ (upstream forward and reverse, respectively); 5’-GTCGACGGAGGATCCCCGGAATTTCTTCTTTGCATTGTTGATACTTAGGGC-3’, and 5’-CTGCTGCAA CTCAAGGATATCCC-3’ (downstream forward and reverse, respectively). Underlined sequence indicates tags complementary to sequence tags added to primers used to amplify a kanamycin resistance marker. PCR product of sequence immediately upstream and downstream of the target loci was then used with tags complementary to sequence tags added to primers used to amplify a kanamycin resistance marker. PCR product of sequence immediately upstream and downstream of the target loci was then used with tags complementary to sequence tags added to primers used to amplify a kanamycin resistance marker.

Antimicrobial peptide susceptibility tests. To determine the MIC of BPI, polymyxin B, and LL-37 susceptibility to BPI, polymyxin B, and LL-37 was prepared in 96-well plates with 2.5 mM CaCl2 at 37°C. Serial dilutions of BPI, polymyxin B, and LL-37 were prepared in 96-well plates containing 100 μl BHI and 2.5 mM CaCl2 per well. Aliquots of 105 CFU of wild-type (CO92), CO92 ∆bipA YopE-Bla, and CO92 ∆pCDJ YopE-Bla strains were generated as described previously (35, 51). Y. pestis strains were grown on brain heart infusion (BHI) agar (Difco) at 26°C for 2 to 3 days. Strains were confirmed by growth on Congo red plates, and PCR was performed to confirm the presence of the pgm locus and the pCD1 plasmid for wild-type and mutant strains. For infection, Y. pestis strains were inoculated into 10 ml BHI broth supplemented with 2.5 mM CaCl2 and incubated at 37°C for 12 to 16 h with constant shaking. Infections with fully virulent Y. pestis CO92 were performed in the UAMS Biosafety level 3 (BSL-3) facility. For bacterial growth curve analysis, optical density of Y. pestis cultured in BHI or PMH2 medium was obtained using a spectrophotometer at the indicated time points.

Cell culture and cell culture infections. MH-S cells were obtained from ATCC and were cultured in RPMI 1640 (Corning) medium containing 10% fetal bovine serum (FBS) (Gibco) and Antibiotic-Antimycotic (Gibco). Cell cultures were incubated at 37°C with 5% CO2. To perform bacterial killing assays, primary human neutrophils (Astarte Biologics), primary human alveolar macrophages (hAMs), and MH-S cells were cultured briefly in IMDM (Gibco), DMEM:F12K 1:1 (Corning), or RPMI 1640 (Corning) medium supplemented with 10% human AB sera or FBS. After determining cell density, cells were inoculated with Y. pestis using an MOI of 1:1 CFU bacteria to host cell. Cells were then incubated at 37°C with 5% CO2 for up to 8 h. At 2, 4, and 8 hpi, cells were scraped and 100 μl from each technical replicate was serially diluted and plated on BHI agar to determine CFU. Gentamicin protection assays were performed by inoculating cells with Y. pestis strains using an MOI of 1:1. After the indicated time points postinfection, cells were treated with 8 μg/ml gentamicin-containing (52) cell culture medium for 1 h. After gentamicin treatment, medium was aspirated from cells and replaced with sterile diH2O to lyse cells. Technical replicates were serially diluted and plated on BHI agar to determine levels of intracellular Y. pestis.

Antimicrobial peptide susceptibility tests. To determine the MIC of BPI, polymyxin B, and LL-37 needed to inhibit growth of Y. pestis strains, strains were cultured for 16 h in BHI broth supplemented with 2.5 mM CaCl2, at 37°C. Serial dilutions of BPI, polymyxin B, and LL-37 were prepared in 96-well plates containing 100 μl BHI and 2.5 mM CaCl2 per well. Aliquots of 105 CFU of wild-type (CO92), ∆bipA, or ∆bipA::bipA Y. pestis were inoculated into wells and incubated overnight at 37°C with shaking. The MIC was determined the following day by assessing the concentration of antimicrobial peptide needed to
inhibit Y. pestis growth. To determine susceptibility to subinhibitory concentrations of antimicrobial peptides, the Y. pestis strains were cultured for 16 h in BHI broth and then aliquots of wild-type, ΔbipA, and ΔbipA:ΔapY. pestis were subcultured in PMH2 (5%) medium with or without 40 μg/ml BPI (R&D Systems), 1 μg/ml polymixin B (Fisher), or 64 μg/ml LL-37 (Tocris Biosciences). Cultures were grown at 37°C with constant shaking and at 2, 4, and 8 hpi, 100 μl from each technical replicate was serially diluted and plated on BHI agar to determine CFU.

**Preparation of human alveolar macrophages.** To isolate human alveolar macrophages (hAMs), lungs obtained from anonymous donors through the Arkansas Organ Recovery Agency were lavaged using phosphate-buffered saline (PBS). Lavage fluid was centrifuged at 500 × g for 10 min and treated with red blood cell lysis buffer (0.15 M NH4Cl, 12 mM NaHCO3, and 0.1 mM EDTA) for 5 min. The remaining cells were washed with PBS and reconstituted in DMEM-F12K 1:1 (GE) supplemented with 10% FBS (Gibco) and Antibiotic-Antimycotic (Gibco). Cells were plated in 12-well plates and incubated at 37°C with 5% CO2 for 4 to 5 days. Cells were counted prior to infection.

**Animals, animal infections, and ethical approval.** All animal experiments were conducted with approval from the UAMS Institutional Animal Ethics Committee. Six- to 8-week old C57BL/6J female mice were obtained from Jackson Laboratories. Mice were provided with food and water ad libitum and maintained at 25 to 26°C with 40 to 70% humidity. To prepare mice for infection, mice were anesthetized using ketamine/xylazine injected via the intraperitoneal route. Once anesthetized, mice were inoculated via the intranasal route with lethal doses (10³ bacteria) of Y. pestis, Y. pseudotuberculosis, or control. Mice were sacrificed using a lethal dose of sodium pentobarbital injected via the intraperitoneal route. To determine bacterial burdens in lungs and spleens of mice, lungs and spleens were homogenized in 2 ml of 1% NaCl and plated on BHI agar to determine CFU. For neutrophil depletion, 20 μg Ly-6G antibody (clone 1A8, BioLegend) was injected via the intravenous route one day prior to and on the day of infection, as described previously (48).

**Neutrophil degranulation assay.** Neutrophils were isolated from human peripheral blood samples and degranulation was quantified as described previously (30). Briefly, 10⁶ isolated human neutrophils were infected in triplicate with 10⁶ CFU of wild-type, ΔbipA, or ΔpcD1 Y. pestis CO92 or mock-inoculated with PBS. Neutrophils and Y. pestis were incubated for 1 h at 37°C. After incubation, neutrophils were washed in PBS with 2% FBS, then stained for 25 min at 4°C with CD63-phycoerythrin-Cy7 (clone H5C6, Invitrogen) with PBS. Neutrophils and 10⁶ isolated human neutrophils were acquired using a Millipore Guava 6HT flow cytometer and analyzed with the InCyte EasyCyte v3.1 software.

**Preparation of BALF for flow cytometry.** Bronchoalveolar lavage fluid (BALF) was retrieved from mice postmortem. BALF was centrifuged for 5 min at 500 × g to pellet cells. Cells were resuspended in 3% FBS in PBS containing the following antibodies (1:500 dilution) for 30 min at 4°C to stain cell surface markers: CD45-phycocerythrin (clone 30-F-11, BD Biosciences), CD3-allophycocyanin-Cy7 (clone 17A2, BD Biosciences), CD11b-Alexa Fluor 700 (clone M1/70, BD Biosciences), CD11c-Brilliant Violet 786 (clone H3, BD Biosciences), F4/80-allophycocyanin (clone BM8, Invitrogen), and Ly-6G-phycocerythrin-Cy7 (clone 1A8, BD Biosciences). After antibody staining, cells were centrifuged at 500 × g for 5 min and the supernatant was removed. Cells were subsequently stained using 1 × CCF2-AM (Invitrogen) for 30 min at room temperature. Cells were pelleted at 500 × g for 5 min and resuspended in 2% formalin in PBS for fixation at 20 min at room temperature. After fixation, cells were pelleted and resuspended in 200 μl of 3% FBS in PBS for flow cytometry. The following cell populations were identified as described previously (35): alveolar macrophages (F4/80⁺CD11b⁻/hiCD11c⁻/lo), CD11b⁻/hi interstitial/macrophages (F4/80⁺CD11b⁻/hiCD11c⁻/hi), and CD45⁺CD11b⁻/loLy-6G⁻ lymphocytes (F4/80⁻CD11c⁻/loCD11b⁻/loLy-6G⁻).

**Proteomic analysis.** To obtain protein extracts from broth-grown Y. pestis, equal optical densities of Y. pestis CO92 and Y. pestis CO92 ΔbipA cultures were spun down at 10,000 × g for 5 min to pellet bacterial cells. To obtain bacterial protein extracts from Y. pestis incubated with BPI, Y. pestis was incubated with 40 μg/ml BPI or PBS for 30 min at 37°C. Bacteria were spun at 10,000 × g for 5 min to pellet bacteria. Bacterial pellets were resuspended in 1 × Laemmlle buffer (Bio-Rad) and boiled for 5 min at 96°C to lyse cells. Volumes of each cell lysate sample (normalized to CFU) were resolved via SDS-PAGE and stained with Coomassie brilliant blue to visualize protein bands. Protein gels were subsequently submitted to the UAMS Proteomics Core for in-gel trypsin digestion, MS/MS analysis, and protein identification and quantification. Spectral counts were exported from Scaffold (Proteome Software) to Microsoft Excel for further statistical analyses.

**Statistical analyses.** Statistical analyses were done using one-way and two-way analysis of variance (ANOVA) and log rank test where applicable. P values of < 0.05 are represented as *, of <0.005 as **, of <0.0005 as ***, and of <0.0001 as ****. All statistical analyses were performed using GraphPad Prism v7.04 software.

**SUPPLEMENTAL MATERIAL**

SUPPLEMENTAL material is available online only.

SUPPLEMENTAL FILE 1, PDF file, 0.1 MB.

SUPPLEMENTAL FILE 2, PDF file, 0.1 MB.
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