Correlates of Immune Protection following Cutaneous Immunization with an Attenuated *Burkholderia pseudomallei* Vaccine

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Infections with the Gram-negative bacterium *Burkholderia pseudomallei* (melioidosis) are associated with high mortality, and there is currently no approved vaccine to prevent the development of melioidosis in humans. Infected patients also do not develop protective immunity to reinfection, and some individuals will develop chronic, subclinical infections with *B. pseudomallei*. At present, our understanding of what constitutes effective protective immunity against *B. pseudomallei* infection remains incomplete. Therefore, we conducted a study to elucidate immune correlates of vaccine-induced protective immunity against acute *B. pseudomallei* infection. BALB/c and C57BL/6 mice were immunized subcutaneously with a highly attenuated, Select Agent-excluded purM deletion mutant of *B. pseudomallei* (strain Bp82) and then subjected to intranasal challenge with virulent *B. pseudomallei* strain 1026b. Immunization with Bp82 generated significant protection from challenge with *B. pseudomallei*, and protection was associated with a significant reduction in bacterial burden in lungs, liver, and spleen of immunized mice. Humoral immunity was critically important for vaccine-induced protection, as mice lacking B cells were not protected by immunization and serum from Bp82-vaccinated mice could transfer partial protection to nonvaccinated animals. In contrast, vaccine-induced protective immunity was found to be independent of both CD4 and CD8 T cells. Tracking studies demonstrated uptake of the Bp82 vaccine strain predominately by neutrophils in vaccine-draining lymph nodes and by smaller numbers of dendritic cells (DC) and monocytes. We concluded that protection following cutaneous immunization with a live attenuated *Burkholderia* vaccine strain was dependent primarily on generation of effective humoral immune responses.

Melioidosis is a serious disease of humans caused by the Gram-negative facultative intracellular bacterium *Burkholderia pseudomallei* (1–3). The disease is endemic in Southeast Asia, northern Australia, and parts of South and Central America (1, 4–6). *B. pseudomallei* is being isolated increasingly in other parts of the world as well, likely due to greater awareness and surveillance for the organism (7, 8). Infection with *B. pseudomallei* can be contracted via several routes, including subcutaneous (s.c.) inoculation, inhalation, and likely ingestion (9, 10). The route of *B. pseudomallei* infection is correlated with the severity of infection, with inhalational infection generally associated with a more rapid disease course. Bacteremic infection is common with *B. pseudomallei*, and sepsis in melioidosis patients is associated with high mortality rates (11–13). In patients who develop chronic infection, the disease may manifest as disseminated abscesses in multiple sites, including the spleen, liver, joints, and central nervous system (CNS) (14–18). *B. pseudomallei* is inherently resistant to multiple classes of antibiotics, most notably aminoglycosides and some beta-lactam drugs (19), due to the expression of efflux pumps and PenA β-lactamase (20–23). A delay in diagnosis is often associated with treatment failures in patients with acute infection (24–27).

Currently, there is no approved vaccine for protection of humans against *B. pseudomallei* infection. A number of candidate vaccines have been developed and tested in animal models of melioidosis, and the state of melioidosis vaccines has been reviewed recently (28). Briefly, the most effective immunity to date has been achieved by use of live attenuated *B. pseudomallei* vaccines, including strains lacking *ilvL* (29, 30); *serC* (31); *aroB* (32); *purN*, *purM*, BPSS1509, *lipB*, and *pabB* (33); and *bipD* (34), and *aroC* (35) mutants.

Subunit vaccines have also been developed for immunization against *B. pseudomallei*, with the most promising protection to date being achieved with purified proteins such as BipD (34) and BipB and BipC (36); with recombinant proteins, including Hcp1, Hcp2, Hcp3, and Hcp6 (37); with lipopolysaccharide (LPS) (38); with purified flagellin (39); with LolC, PotF, and OppA nonmembrane protein (40); with outer membrane vesicles (41); and with recombinant Omp85 protein (42). However, the use of single antigen subunit vaccines for *Burkholderia* infection is unlikely to generate broad protective immunity against this very genetically diverse and unstable organism (35, 43). Effective immunization with subunit vaccines administered by the subcutaneous route has also not been reported.

While the risk of reversion to virulence is a primary concern with the use of live attenuated bacterial vaccines, the ability to induce rapidly broad protective immunity is a plus for this type of vaccine. Our group previously developed a highly attenuated strain of *B. pseudomallei* 1026b (strain Bp82), which was recently excluded from Select Agent regulations and which with Institutional Biosafety Committee approval can be used under biosafety level 2 (BSL-2) conditions (44). This ΔpurM strain of *B. pseud-
domallei was extensively tested in several different highly immuno-compromised animal strains, and reversion to virulence or persistence of the organism was not found (44). In addition, a purM deletion mutant of the K96243 strain of $B. \text{pseudomallei}$ was also highly attenuated and safe in animal studies (though this strain is not yet excluded from Select Agent regulations) (44). The ability of these Select Agent-excluded mutant strains of $B. \text{pseudomallei}$ to induce protective immunity from melioidosis has not been previously investigated in animal models. Nor has it been determined whether protection could be achieved by subcutaneous (s.c.) vaccine administration, which is a more practical route of immunization than the intranasal (i.n.) or intraperitoneal (i.p.) routes used in most prior studies of attenuated $B. \text{pseudomallei}$ vaccines.

Therefore, in the present study we intended to determine whether the Bp82 strain of $B. \text{pseudomallei}$ was capable of inducing protective immunity following cutaneous immunization. We also sought to elucidate immune mechanisms by which Bp82 immunization could induce protective immunity and to also understand how the Bp82 vaccine antigens were processed by antigen-presenting cells (APC) in lymph nodes (LN). Our findings indicate that the Bp82 vaccine is immunogenic following s.c. immunization and capable of inducing significant protection against acute inhaled challenge. Protective immunity was provided primarily by humoral immune responses. Therefore, these new insights into protective immune responses generated by live attenuated vaccines such as Bp82 should help guide the development of newer melioidosis vaccines and clinical evaluation of vaccine efficacy using immune correlates.

MATERIALS AND METHODS

Mice. Specific-pathogen-free 6- to 8-week-old female BALB/c and C57BL/6 mice were obtained from the Jackson Laboratory (Bar Harbor, ME). In addition, mutant mouse strains (on the C57BL/6 background), including TCRα/β−/−, uMT−/−, CD4−/−, and CD8−/− strains, were also purchased from Jackson. Mice were used for studies when they were between 8 and 12 weeks of age. Animals were housed in microisolator cages under pathogen-free conditions. All studies were conducted in animal biosafety level 3 (ABSL-3) facilities at the Rocky Mountain Regional Biosafety Laboratory and were approved by the Institutional Animal Care and Use Committee at Colorado State University.

Bacteria and culture conditions. $B. \text{pseudomallei}$ strain 1026b (P1026b) is a clinical isolate from a patient with septicemic melioidosis in Thailand (45, 46). Culture stocks were grown overnight in Luria-Bertani broth (LB) (BD Biosciences, San Jose, CA) at 37°C with shaking. Frozen stocks of $B. \text{pseudomallei}$ were prepared by adding 15% glycerol (Fisher BioReagent, Pittsburgh, PA) to the overnight culture and dividing the sample into 1-ml aliquots. Aliquots were stored at −80°C, and titers were determined prior to use. The construction and characterization of 1026b ΔpurM strain Bp82 have been previously described (44). A 114-bp fragment was deleted from the purM coding sequence, causing attenuation of the strain for the melioidosis animal infection model (44). Because Bp82 is a purine auxotroph, culture was done in LB medium supplemented with 0.6 mM adenine for optimal bacterial growth. For animal experiments, each strain was thawed just before use, and the bacteria were diluted to obtain the desired numbers of cells using sterile phosphate-buffered saline (PBS) (Sigma-Aldrich, St. Louis, MO).

Mouse vaccination. Mice were restrained in a tail bleed apparatus allowing access to the hind leg without the need for anesthesia. Mice were vaccinated s.c. in the hind leg with 5 × 10⁶ CFU Bp82, in 100 µL PBS. Mice received a booster immunization, 10 days after the first immunization.

In vitro stimulation assay for T cell immunity. Mice were euthanized, and single-cell suspensions from spleens were generated via mechanical disruption. Cells were filtered through a 70-µm nylon mesh screen (BD Biosciences, San Jose, CA) and treated with NH₄Cl to remove red blood cells. Spleen cells were plated in 24-well plates at a concentration of 2 × 10⁶ cells/ml and restimulated with antigens for 72 h. These antigens included heat-killed Bp82 (2 × 10⁶ cells/ml), Bp82 lysates (5 µg/ml), or recombinant GroEL (12.5 µg/ml) kindly provided by Katherine Brown, University of Texas Austin). Cell culture supernatants were collected and analyzed for gamma interferon (IFN-γ) and interleukin-17 (IL-17) production via enzyme-linked immunosorbent assay (ELISA), using kits according to the manufacturer’s instructions (R&D Systems, Minneapolis, MN).

Determination of antibody titers. Nunc Maxisorp 96-well plates (Thermo Fisher Scientific, San Jose, CA) were coated with heat-killed Bp82 (5 × 10⁶ bacteria/ml) in carbonate-bicarbonate buffer (pH 9.6) and incubated at 4°C overnight. Plates were washed with PBS and Tween 20 (0.05%) and blocked with PBS-Tween containing 5% nonfat dry milk for 2 h at room temperature. Sera were serially diluted (10-fold dilutions) in blocking buffer and incubated for 90 min at room temperature. All secondary antibodies were conjugated to horseradish peroxidase, including rat anti-mouse IgM and IgG (BD Biosciences, San Jose, CA), goat anti-mouse IgG2a (Abcam, Cambridge, MA), and rat anti-mouse IgG1 (BD Biosciences, San Jose, CA). These antibodies were diluted 1:2,000 in blocking buffer and were incubated on the plate for 1 h at room temperature. 3’,5’,5’-Tetramethylbenzidine (TMB) (Sigma-Aldrich, St. Louis, MO) substrate was used to develop the plates. The reaction was stopped by the addition of 50 µL of 1 N HCl. The absorbance at 450 nm was determined using a Thermo Multiskan EX spectrophotometer (Thermo Scientific, Rockford, IL).

Serum transfer studies. To obtain immune serum, naive BALB/c mice were vaccinated with Bp82 (5 × 10⁶ CFU) s.c. and boosted 10 days later. Two weeks after the boost, mice were anesthetized via intraperitoneal (i.p.) injection with 100 mg ketamine/kg of body weight plus 10 mg/kg xylazine diluted in sterile water, and blood was collected via terminal cardiac puncture. Blood was allowed to clot for 30 min at 4°C and then centrifuged to prepare serum, which was collected and stored at −80°C prior to use. Nonimmune serum was also collected from unvaccinated mice in a similar manner. Anti-$B. \text{pseudomallei}$ IgG titers were quantified via ELISA prior to transfer as previously described (47, 48). Recipient mice received 250 µL of immune serum or naive (control) serum given i.p. 1 day prior to challenge.

B. pseudomallei i.n. challenge model. For intranasal (i.n.) challenge, mice were anesthetized as described above. Bacteria were thawed just prior to use and diluted in sterile PBS (pH 7.4) for inoculation. Mice were challenged with approximately 5.0 × 10⁶ CFU of $B. \text{pseudomallei}$ (5 × 10⁶ CFU for BALB/c mice or 1.2 × 10⁶ CFU for C57BL/6 mice), in a total volume of 20 µL administered in sequential droplets on alternating nares. Actual infectious doses delivered to the mice were determined by plating the inoculum.

Determination of bacterial burden in organs. Lungs, liver, and spleen were harvested at 72 h postinfection in the acute challenge model and at 30 and 60 days postinfection for the chronic challenge model and placed in 4 ml of sterile PBS. Organs were homogenized using a Seward stomacher (Seward, Bohemia, NY). Homogenates (300 µL) were removed, placed in a 96-well plate, and serially diluted in sterile PBS, using 10-fold dilutions. Bacterial counts were determined by plating serial 10-fold dilutions on LB agar and placing them at 37°C, and CFU were enumerated after 48 h of incubation.

Tracking Bp82 uptake in vaccine-draining lymph nodes. To facilitate tracking the fate of Bp82 in vaccine-draining lymph nodes following immunization, Bp82 was engineered to express green fluorescent protein (GFP) (Bp82-gfp). Briefly, strain Bp82 (44) was labeled with an enhanced green fluorescent protein (eGFP) whose expression is driven by the constitutive $B. \text{pseudomallei}$ (5 × 10⁶ CFU for BALB/c mice or 1.2 × 10⁶ CFU for C57BL/6 mice), in a total volume of 20 µL administered in sequential droplets on alternating nares. Actual infectious doses delivered to the mice were determined by plating the inoculum.

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of the site-specific mini-Tn7 kanamycin (Kan) resistance element was performed by triparental conjugation of Bp82 with RH03/pTNS3 and RH03/pUC18T-mini-Tn7-Ps12-eGFP, and transfectants were selected on LB medium containing 1,000 μg/ml kanamycin and 0.6 mM adenine. Isolates were screened by PCR to identify single insertions in the glmS2-associated attTn7 site. An unmarked strain was obtained by Flp-mediated excision of the Kan resistance marker using pFLPe2. Bp82-gfp was injected into the left footpad (1.2 × 10⁸ CFU/mouse) in 50 μl PBS, and popliteal lymph nodes were harvested 10 h after injection and processed as previously described (50). Briefly, lymph nodes were harvested and placed in 5 ml of Hanks’ balanced salt solution (HBSS) and digested in collagenase D (Roche, San Francisco, CA) for 30 min at 37°C. Following digestion, the tissue was triturated using a glass pipette and filtered through a 70-μm nylon mesh screen (BD Biosciences, San Jose, CA).

Flow cytometry. Cells from spleen or lymph nodes were suspended in fluorescence-activated cell sorting (FACS) buffer (PBS with 2% fetal bovine serum [FBS] [Atlas Biologicals, Fort Collins, CO] and 0.05% sodium azide [Fisher Scientific, Philadelphia, PA]) and stained as previously described (31). To block nonspecific binding, cells were incubated at room temperature for 15 min in FACS block consisting of normal mouse serum (Jackson ImmunoResearch, West Grove, PA) and human IgG (Jackson ImmunoResearch, West Grove, PA), along with unlabeled anti-mouse CD16/32 (clone 93) (eBioscience, San Diego, CA) prior to staining. After blocking, cells were stained with the following antibodies: anti-Ly-6-G (phycoerythrin [PE], clone HK1.4), anti-Ly-6-C (PE-Cy7, clone RB6-8C5), anti-CD11c (biotin, clone N418), anti-CD11b (allophycocyanin Alexa 750, clone M1/70), anti-CD169 (Alexa Fluor 647, clone MOMA-1). The antibodies were purchased from eBioscience (San Diego, CA), BD Pharmingen (San Diego, CA), or AbD Serotec (Raleigh, NC). Cells were immunostained at room temperature for 30 min with antibodies diluted in FACS buffer. After the primary antibody incubation, cells were washed in FACS buffer and incubated with streptavidin-peridinin chlorophyll protein (PerCP) for 20 min at room temperature. Cell acquisition was done using a Gallios flow cytometer (Beckman Coulter, Brea, CA). Data were analyzed using FlowJo software (Tree Star, Ashland, OR).

Immunohistochemistry. Lungs and spleen were collected 48 h postinfection and fixed in 4% paraformaldehyde for 48 h. Tissues were paraffin embedded, cut into 5- to 7-μm sections, deparaffinized using EZ-DeWax solution (Biogenex Lab, San Ramon, CA), and transferred into PBS. Sections were blocked with blocking eraser (Biocare Medical, Concord, CA) for 5 min at room temperature and then incubated with appropriately diluted rabbit polyclonal anti-B. pseudomallei antibody (provided by D. Waag from USAAMRIID), and the slides were incubated overnight at room temperature. Slides were washed three times with PBS followed by incubation with anti-rabbit Cy3-conjugated secondary antibody (Millipore, Billerica, MA) for 1 h at room temperature. Slides were washed and counterstained with 4’,6-diamidino-2-phenylindole (DAPI) (0.5 μg/ml) for 2 min and coverslipped with Prolong Gold mounting medium (Invitrogen, Carlsbad, CA). Sections were examined using a Zeiss 510 confocal microscope and analyzed using Volocity software.

Statistical analyses. Statistical analyses were performed using GraphPad Prism5 software (La Jolla, CA). A nonparametric t test was used to analyze statistical differences between 2 groups. Comparisons between multiple groups were done using one-way analysis of variance (ANOVA), followed by Tukey’s multiple means comparison test. Survival differences were compared using Kaplan-Meier survival curves, followed by log rank test. Statistical significance was defined as P < 0.05.

RESULTS

Immunization with Bp82 protects against lethal B. pseudomallei challenge. To determine whether s.c. immunization with Bp82 was capable of generating protective immunity against lethal inhaled challenge with B. pseudomallei, BALB/c and C57BL/6 mice were immunized twice with Bp82 (5 × 10⁸ CFU per immunization) and then subjected to lethal i.n. challenge with B. pseudomallei strain 1026b. The immunizing dose of Bp82 selected for these studies was comparable to doses used in previous studies of live attenuated Burkholderia vaccines. Following challenge with B. pseudomallei, mice were monitored closely and euthanized when they met predetermined endpoints for signs of illness. We observed that unvaccinated mice (both C57BL/6 and BALB/c) reached morbidity endpoints within 3 days of challenge and were humanely euthanized. In contrast, there was 100% survival of vaccinated C57BL/6 mice (Fig. 1A) and 80% survival of vaccinated BALB/c mice, for at least 30 days after challenge (Fig. 1B) (P < 0.05 compared to unvaccinated mice). At 60 days postchallenge, Bp82 vaccination conferred 100% protection on C57BL/6 mice (Fig. 1A), whereas 60% of BALB/c mice were still alive and apparently healthy (Fig. 1B).

Vaccination with Bp82 reduces bacterial burden. The evidence that Bp82 is not detectable in mouse organs after 48 h of inoculation was previously considered by the study in reference 44, which challenged mice by the i.n. route with 6 × 10³ CFU of Bp82 or B. pseudomallei 1026b. After 48 h of infection, the authors found that Bp82 was below the limit of detection in the lung, liver, and spleen. Therefore, studies were done next to test the effects of Bp82 vaccination on the B. pseudomallei 1026b bacterial burden in key organs shortly after challenge. We observed that the bacterial burden in the lung and the spleen 72 h after B. pseudomallei
challenge was significantly reduced in Bp82-vaccinated animals compared to unvaccinated animals (Fig. 2). In addition, bacteria were not detected in the livers of Bp82-vaccinated animals, compared to the high bacterial burden in the livers of unvaccinated mice (Fig. 2). At 30 and 60 days after challenge, Bp82-vaccinated mice still had detectable B. pseudomallei in the spleen and small amounts in the lung and liver (data not shown).

Induction of antibody responses following Bp82 immunization. Serum from vaccinated mice was analyzed 10 days after the last immunization for induction of antibodies against Bp82 intact organisms. In the serum of vaccinated animals, significantly increased titers ($P < 0.05$) of antibodies against intact Bp82 were detected, compared to unvaccinated control animals (Fig. 3). Vaccinated animals mounted strong IgG responses (Fig. 3A), as well as IgM responses (Fig. 3B), against heat-killed Bp82. In addition, there was significant induction of IgG responses of the IgG1 and IgG2a isotypes (Fig. 3C and D, $P < 0.05$). Immunized animals also had significantly increased titers against antigens present in Bp82 lysates (data not shown). Immunization with Bp82 also induced antibody responses against certain immunogenic proteins of B. pseudomallei, including GroEL (data not shown), but not other antigens such as BimA (data not shown).

Induction of cellular immune responses following Bp82 immunization. The ability of the Bp82 vaccine to induce specific T cell responses was evaluated next. Spleens were harvested from immunized and nonimmunized mice 2 weeks after s.c. Bp82 vaccination. Single-cell suspensions of spleen cells were placed in triplicate wells of 96-well plates as noted in Materials and Methods and incubated with heat-killed Bp82, lysed Bp82, or recombinant GroEL protein. Incubation of spleen cells from immunized mice with heat-killed Bp82 generated production of significant increases in IFN-$\gamma$ and IL-17 production (Fig. 4A and B). However, incubation with Bp82 did not induce production of IL-10 from spleen cells (data not shown). Lysates of Bp82 or recombinant GroEL, however, did not induce significant cytokine production. Thus, immunization with the Bp82 vaccine induced cellular immune responses that appeared to be directed primarily toward surface determinants on Bp82.

FIG 2 Immunization reduces bacterial burden in vaccinated mice at 72 h after challenge. Bacterial burden in lung, spleen, and liver tissues was determined in naive and vaccinated mice ($n = 5$) 72 h after intranasal challenge with B. pseudomallei strain 1026b. Statistical comparisons between vaccinated and naive groups were done using a nonparametric $t$ test. *, $P < 0.05$. Similar results were obtained in one additional experiment.

FIG 3 Humoral immune responses in Bp82-vaccinated mice. Serum was prepared from BALB/c and C57BL/6 mice ($n = 5$) vaccinated twice s.c. with Bp82, and titers of antibodies to heat-killed Bp82 were determined by endpoint dilution ELISA. Titers of total IgG (A), IgM (B), IgG1 (C), and IgG2a (D) were expressed as the reciprocal of the endpoint dilution. Significant differences (**, $P < 0.05$; ***, $P < 0.005$; ****, $P < 0.0001$) were determined by a nonparametric Kruskal-Wallis test followed by Dunnett’s multiple comparison test. Data represent pooled data from two independent experiments.
Role of humoral immunity in vaccine-induced protection against *B. pseudomallei*. In previous studies where immune mechanisms of protection have been examined, humoral immunity was found to be a primary mediator of vaccine-induced immune protection against acute *B. pseudomallei* challenge (52–55). Therefore, the role of antibody-mediated protection was investigated in the Bp82 vaccine model. First, serum transfer experiments were conducted, by transferring serum from Bp82-vaccinated animals to naive animals, which were then subjected to challenge. These animals were also significantly protected from infection 3 days after challenge. Mice lacking all conventional T cells (TCRα/β−/−) were also immunized with the Bp82 vaccine and subjected to challenge. These results indicated that humoral immunity is an important, but incomplete, role in Bp82 vaccine-induced protection from *B. pseudomallei* infection.

Role of T cells in Bp82 vaccine-induced protection. Studies were done next to elucidate the role of T cells in vaccine-induced immunity to *Burkholderia* infection. Mice lacking CD4 T cells (CD4−/−) or CD8 T cells (CD8−/−) on the C57BL/6 background were vaccinated s.c. with the Bp82 vaccine and subjected to *B. pseudomallei* challenge. Survival times were compared to those of WT animals (Fig. 6). At 30 days after challenge, there was no statistical difference in survival times between CD8−/− or CD4−/− vaccinated animals and vaccinated WT mice (Fig. 6A and B). All animals from CD4−/− and TCRα/β−/− groups survived to at least 60 days after challenge and did not manifest signs of infection, whereas unvaccinated C57BL/6 mice were euthanized due to progressive infection 3 days after challenge. Mice lacking all conventional T cells (TCRα/β−/−) were also immunized with the Bp82 vaccine and subjected to challenge. These results also suggest that it is likely that T-cell-independent antibodies such as IgM antibodies may have played a major role in vaccine-induced protection.

**Delivery of Bp82 vaccine to draining LN.** Finally, studies were conducted to assess the uptake and trafficking of the Bp82 vaccine to vaccine-draining lymph nodes (LN). We recently demonstrated that following immunization with conventional adjuvanted vaccines, a significant and rapid influx of inflammatory monocytes into vaccine-draining lymph nodes occurs (36). Therefore, studies were conducted to determine whether the Bp82
vaccine elicited a similar leukocyte response in LN and to also identify potential antigen-presenting cells (APC) responsible for trafficking of Bp82 from the cutaneous inoculation site to the LN. Therefore, a gfp-expressing construct of Bp82 was used to track the early movement of Bp82 vaccine bacteria from the skin site to the nearest draining LN. For these studies, the vaccine was administered s.c. in the footpad and the popliteal LN response was monitored using flow cytometry.

We found that following s.c. administration of the Bp82 vaccine, there was a marked influx of CD11b⁺ Ly6G⁺ Ly6C⁻ neutrophils into the nearest draining LN, as well as a much smaller influx of CD11b⁺ Ly6G⁻ Ly6C⁺ monocytes (Fig. 7A and B). We also found that approximately 80% of all Bp82-gfp bacteria found in the LN after vaccination were contained within neutrophils (Fig. 7C). In addition, bacteria in the LN were also found within Langhans dendritic cells (DC) (10%), CD169⁺ subcapsular macrophages (3%), and inflammatory monocytes (3%), with very few non-cell-associated bacteria being detected.

To compare the relative efficiencies of uptake of Bp82-gfp by antigen-presenting cells in the LN, the percentages of each cell population containing Bp82 were calculated. We calculated that 22.12% of all neutrophils in the LN, 1.87% of all DC in the LN, 16.17% of subcapsular macrophages in the LN, and 11.62% of all...
inflammatory monocytes in the LN contained Bp82-gfp. Thus, neutrophils appeared to be the most efficient antigen-presenting cells at taking up and transporting Bp82 bacteria to the draining LN, and the majority of Bp82 bacteria in the LN were found within neutrophils.

**DISCUSSION**

A number of different live attenuated and subunit vaccines have been evaluated for their ability to generate protective immunity against *B. pseudomallei* (29, 30, 32–36, 38–42, 55). However, there is still relatively little known regarding the most effective methods of generating protection against melioidosis or the immune mechanisms of protection. In the present study, we demonstrated long-term (i.e., >60 days) protection against acute *B. pseudomallei* challenge following cutaneous immunization with a live attenuated strain of *B. pseudomallei*. Importantly, our study utilized a more conventional route of immunization (i.e., cutaneous immunization) than did previous studies, which have typically relied on intranasal or intraperitoneal routes of immunization. In addition, we found that the Bp82 vaccine did not require an adjuvant for activity (data not shown). Another important consideration is the fact that the Bp82 vaccine strain used in this study has been extensively evaluated for attenuation and safety. Thus, issues of reversion to virulence are potentially less of a concern with the purM deletion mutant of *B. pseudomallei*.

There remain substantial gaps in our knowledge regarding mechanisms of immune protection from melioidosis. The best protective immunity to date has been achieved using live attenuated *Burkholderia* vaccines, but the mechanisms of protection are not completely understood. There is some evidence that protective immunity is dependent on induction of innate immune responses by live bacterial vaccines (43, 57). Also, there is speculation that live attenuated vaccines generate better immunity due to prolonged antigen persistence in the host (33).

Protection generated by the Bp82 vaccine was found to be mediated almost entirely by antibodies, as revealed by serum transfer studies and by studies in B-cell- and T-cell-deficient mice. These findings are consistent with those of prior studies, where protection from inhaled *Burkholderia* challenge was found to be mediated largely by humoral immune mechanisms. However, these are the first studies to demonstrate that protection acquired by s.c. immunization is also antibody mediated and that routes of mucosal administration were not required to induce effective levels of immune protection against inhalational challenge.

Interestingly, our studies also revealed a minor role for T cells in vaccine-mediated protection. For example, while CD4<sup>+</sup>/CD8<sup>+</sup> animals did not have a defect in immune protection, mice that lacked both CD4 and CD8 T cells (i.e., TCRε/B<sup>−/−</sup> mice) did have a small, though not statistically significant, loss of immune protection following Bp82 immunization (Fig. 6C). While the Bp82 vaccine induced both IgG and IgM antibodies, the fact that CD4<sup>+</sup> mice were fully protected following immunization suggests an important potential role for IgM antibodies in mediating vaccine protection, as CD4<sup>−</sup> mice are generally unable to effectively produce antibodies of the IgG subclass (58).

Our findings indicate that administration of a live attenuated vaccine by the s.c. route is effective in generating systemic protection from bacterial challenge by a mucosal route. Though relatively high titers of anti-*Burkholderia* antibodies were detected in serum of vaccinated animals, it is not clear exactly where bacterial control by vaccine antibodies occurred. It is plausible to suggest that vaccination may protect from lethality mainly by blocking bacterial dissemination from the lungs to other sites (e.g., spleen and liver), rather than by neutralizing bacteria directly in the lungs. The protective antibody immune response in *Burkholderia* infection has been shown previously to be specific to the lipopolysaccharide (LPS) of *B. pseudomallei*, which promotes opsonic phagocytic killing (59, 60).

The response of antigen-presenting cells in LN to immunization with the live attenuated Bp82 vaccine was found to be quite different from the response to conventional adjuvanted vaccines, with the primary difference being the much more pronounced neutrophilic response in the case of mice immunized with the Bp82 vaccine. It is unlikely that neutrophils are able to present Bp82 antigens directly to T cells but much more likely that neutrophils containing vaccine bacteria hand the antigens off to professional antigen-presenting cells in the LN, including DCs, subcapsular macrophages, or monocytes. Indeed, it was recently observed for a live *Mycobacterium boris* BCG vaccine that the vaccine bacteria are taken up primarily by neutrophils but then later handed off to other antigen-presenting cells (e.g., DC) in the LN and peritoneal cavity (61). With the Bp82 vaccine, most of the bacteria were delivered to the LN in neutrophils, but the most avidly phagocytic antigen-presenting cells for Bp82 in the LN were found to be CD169<sup>+</sup> macrophages and inflammatory monocytes. Thus, it is likely that initial uptake of live attenuated vaccines such as Bp82 may be mediated by neutrophils, while the bacterial antigens may ultimately be delivered to macrophages and monocytes within the LN for presentation to T cells.

In summary, we have found that s.c. immunization with a highly attenuated strain of *B. pseudomallei* can generate significant protection against inhalational challenge with virulent *B. pseudomallei*. These studies suggest, therefore, that it may be possible to develop conventionally delivered vaccines (i.e., vaccines administered by the s.c. or intramuscular [i.m.] route) capable of generating effective humoral immune protection against melioidosis. While subunit vaccines are generally preferred over live attenuated bacterial vaccines, it may be possible as an interim measure for melioidosis protection to use an attenuated vaccine, particularly if vaccine administration is safe and easily accomplished. Our studies in mice indicated that vaccine site reactions did not occur (data not shown), suggesting that the whole-cell Bp82 vaccine did not induce significant local inflammation. Thus, there is reason for optimism that an effective subunit or nonliving bacterial cell vaccine for melioidosis that is safe and easily administered can be developed.

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