Oral Live Vaccine Strain-Induced Protective Immunity against Pulmonary Francisella tularensis Challenge Is Mediated by CD4⁺ T Cells and Antibodies, Including Immunoglobulin A

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Francisella tularensis is an intracellular gram-negative bacterium and the etiological agent of pulmonary tularemia. Given the high degrees of infectivity in the host and of dissemination of bacteria following respiratory infection, immunization strategies that target mucosal surfaces are critical for the development of effective vaccines against this organism. In this study, we have characterized the efficacy of protective immunity against pneumonic tularemia following oral vaccination with F. tularensis LVS (live vaccine strain). Mice vaccinated orally with LVS displayed colocalization of LVS with intestinal M cells, with subsequent enhanced production of splenic antigen-specific gamma interferon and of systemic and mucosal antibodies, including immunoglobulin A (IgA). LVS-vaccinated BALB/c mice were highly protected against intranasal (i.n.) SCHU S4 challenge and exhibited significantly less bacterial replication in the lungs, liver, and spleen than mock-immunized animals. Depletion of CD4⁺ T cells significantly abrogated the protective immunity, and mice deficient in B cells or IgA displayed partial protection against SCHU S4 challenge. These results suggest that oral vaccination with LVS induces protective immunity against i.n. challenge with F. tularensis SCHU S4 by a process mediated cooperatively by CD4⁺ T cells and antibodies, including IgA.

Francisella tularensis is an intracellular gram-negative bacterium that can cause acute pneumonic disease in humans (18, 54). F. tularensis can be classified into several subspecies, including Francisella tularensis subsp. tularensis (type A), Francisella tularensis subsp. holarctica (type B), “Francisella tularensis subsp. novicida,” and Francisella tularensis subsp. mediasiatica (55). The ease of aerosol dissemination and the ability to cause pneumonic disease by inhalation of as few as 10 organisms of a type A strain have made this organism a potential biothreat agent (48). An attenuated strain of F. tularensis subsp. holarctica (type B), the live vaccine strain (LVS), has been evaluated for protection of humans and animals (14, 48). Parenteral administration of LVS to humans by scarification has been shown to provide protection against intradermal (i.d.) challenge with type A but afforded minimal protection from SCHU S4 by a process mediated cooperatively by CD4⁺ T cells and antibodies, including IgA.

Protective immunity against F. tularensis requires the efficient induction of cellular immunity, including T cells, and gamma interferon (IFN-γ) induction (16, 17, 52, 63). Moreover, evidence for the role of antibodies (26, 41, 44, 45, 53), and particularly immunoglobulin A (IgA) (4), in mucosal immunity against Francisella infection has been accumulating. IgA is the principal immunoglobulin isotype involved in the inhibition of bacterial attachment and the neutralization of viruses at mucosal surfaces (31). Moreover, serum IgA and secretory IgA have been shown to suppress inflammatory pathology by reducing inflammatory cytokine production or the oxidative burst (21, 37, 60). Thus, a targeted vaccination regimen that induces cellular and mucosal immunity in the respiratory compartment may be highly beneficial in defense against an F. tularensis type A strain.

In this study, we examined various mechanisms that underlie protective immunity induced by oral LVS vaccination against murine pulmonary tularemia. Mice vaccinated orally with LVS were remarkably protected against subsequent intranasal (i.n.) or i.d. challenge with the F. tularensis type A strain SCHU S4. The significant protection conferred by oral LVS immunization was reflected in reductions in the degrees of bacterial replication and dissemination following pulmonary challenge.
The oral vaccination regimen induced splenic antigen-specific IFN-γ responses and serum IgG2α responses. Moreover, orally vaccinated mice produced LVS-specific feral and respiratory secretory IgA. The respiratory protection conferred by oral LVS vaccination was partially dependent on B cells and on IgA production and required the presence of CD4+ T cells.

**MATERIALS AND METHODS**

**Bacteria.** Francisella tularensis LVS (lot 703-0303-016) was obtained from Rick Lyons at the University of New Mexico, and F. tularensis subsp. tularensis (strain SCHU S4) was obtained from the Centers for Disease Control and Prevention. The bacteria were grown at 37°C in Trypticase soy broth (TSB) or on Trypticase soy agar (TSA), each supplemented with 0.1% (wt/vol) cysteine (25). mCherry-expression plasmid pKEK1124 was then electroporated into LVS as described previously (33). Bacteria were grown at 37°C in TSB-cysteine plus 10 μg/ml of tetracycline (Sigma-Aldrich, St. Louis, MO). LVS expressing mCherry was visualized by fluorescent microscopy (wavelength, 587 to 610 nm) using a Axioskop 2 Plus microscope (Zeiss, Thornwood, NY) and by confocal microscopy (with a 510 Meta laser scanning confocal microscope [Zeiss]).

**Vaccination.** BALB/c and C57BL/6 mice were purchased from the National Cancer Institute (Bethesda, MD), BALB/c IFN-γ−/− mice (11) and C57BL/6 IFN-γ−/− mice were generated as described previously (37), and the mCherry expression plasmid pKEK894 (65), and the mCherry expression plasmid pKEK1124 was then electroporated into LVS as described previously (33). Bacteria were grown at 37°C in TSB-cysteine plus 10 μg/ml of tetracycline (Sigma-Aldrich, St. Louis, MO). LVS expressing mCherry was visualized by fluorescent microscopy (wavelength, 587 to 610 nm) using a Axioskop 2 Plus microscope (Zeiss, Thornwood, NY) and by confocal microscopy (with a 510 Meta laser scanning confocal microscope [Zeiss]).

**Vaccination and challenge.** Mice were anesthetized with 3% isoflurane by use of a rodent anesthesia machine (Harvard Apparatus, Holliston, MA) (40, 41). Mice were either vaccinated orally, using a 22-gauge, 25-mm-long, 1.25-mm-tip feeding needle (Fine Science Tools Inc., Foster City, CA) (20), with 10^6 CFU of LVS in 200 μl of phosphate-buffered saline (PBS) or mock immunized with PBS alone. We have determined the 50% lethal dose (LD50) of LVS administered orally to be approximately 10^6 CFU. Vaccinated mice were rested for the indicated periods and challenged i.n. with 50, 100, or 500 LD50 of SCHU S4 (LD50, <10 CFU [50, 56]) in 25 μl PBS at 3 or 8 weeks after oral LVS vaccination. For i.d. challenge, mice were injected at the base of the tail with either 100 or 500 LD50 of F. tularensis SCHU S4 in 50 μl of PBS. Some mice received a second oral vaccination boost (10^6 CFU) of LVS 8 weeks after the initial oral vaccination. Mice were challenged i.n. with SCHU S4 after 4 weeks. The actual vaccination and challenge doses administered in each experiment were determined by dilution plating on TSA plus cysteine. Animals were monitored daily for morbidity and mortality.

**Splenocyte culture for analysis of cytokine production.** Mice were either immunized orally with 10^7 CFU of LVS or mock immunized with PBS alone and were euthanized 14 days after immunization. Splenocytes were then collected. Single-cell suspensions were prepared and cultured (1 × 10^6 cells/well) for 72 h in Dulbecco’s modified Eagle medium supplemented with 10% (vol/vol) fetal bovine serum (FBS) with or without 10^5 or 10^6 CFU of UV-inactivated LVS. Bacteria were inactivated by exposure to a 30-W short-wavelength UV light source for 15 min at a distance of 15 cm. The inactivation was confirmed by the absence of bacterial growth on TSA-plus-cysteine plates. Some cells were also cultured with the unrelated antigen hen egg lysozyme (HEL). Culture supernatants were harvested for IFN-γ, interleukin 2 (IL-2), and IL-4 analysis by enzyme-linked immunosorbent assay (ELISA) as described previously (40, 41).

**Detection of antibody and isotype levels by ELISA.** Three weeks after oral LVS vaccination, mice were bled and sera prepared. For collection of bronchoalveolar lavage (BAL) fluid, the mice were sacrificed and the trachea were intubated using a 0.58-mm (outer diameter) polyethylene catheter (Becton Dickinson, Sparks, MD). The lungs were then lavaged twice with Hank’s balanced salt solution (Invitrogen, Carlsbad, CA). The recovered BAL fluid (1 ml) was centrifuged at 9,500 × g for 7 min at 4°C, and the supernatant was stored at −70°C until use. For analysis of fecal supernatants, 0.1 g of fresh fecal pellets was collected and dissolved in 1 ml of PBS containing protease inhibitor cocktail (Roche Applied Science, Indianapolis, IN), and supernatants were collected by centrifugation. Microtiter plates were coated overnight with 10^6 CFU of UV-inactivated LVS in sodium bicarbonate buffer (pH 9.5), washed with PBS containing 0.3% Brij 35 (Sigma), and blocked for 2 h at room temperature with PBS containing 5% FBS and 0.1% Brij 35 as described previously (44). Serial dilutions of serum (starting at a 1:50 dilution), undiluted BAL fluid, or fecal supernatants were added to wells and incubated at room temperature for 2 h. The plates were then washed and incubated for an additional 2 h with goat anti-mouse total Ig, IgG1, IgG2a, IgA, and IgM conjugated to horseradish peroxidase (Southern Biotechnology Associates, Birmingham, AL). After incubation, the plates were washed, and a tetracycline benzidine substrate (BD Biosciences, San Diego, CA) was added for color development. Absorbance at 630 nm was measured using an ELISA microplate reader (Bio-Tek Instruments). The reciprocal serum dilutions corresponding to 50% maximal binding were used to obtain titers. However, because of the large dilution involved in the procedures for collection of BAL fluid and fecal supernatants, these samples were tested undiluted, and results were reported as absorbance units. No binding of immune serum was detected in plates coated with the unrelated antigen HEL.

**M-cell and LVS containing.** Groups of BALB/c mice were immunized orally with 10^6 CFU of mCherry-labeled LVS. After 90 min, mice were euthanized, and sections of small intestine were removed, embedded in freezing medium (Triangle Biomedical Sciences, Durham, NC), and stored at −80°C. Cryosections (thickness, 5 μm) were prepared, fixed with 4% paraformaldehyde at 4°C for 1 h, and blocked for an additional 1 h in PBS containing 10% FBS. Sections were stained at room temperature for 1 h with the fluorescein isothiocyanate-conjugated lectin Ulex europaeus agglutinin I (UEA-1) (20 μg/ml; Sigma) to visualize M cells, and with Hoechst stain (Sigma). Images were acquired using a 510 Meta laser scanning confocal microscope (Zeiss) and were analyzed using Imaris software (Bitplane, Salt Lake, PA).

**CD4+ T-cell depletion.** The hybridoma cell line GK1.5 (36) was purchased from the ATCC and grown in HyQ serum-free medium (HyClone) supplemented with decreasing amounts (20% to 1.25%) of FBS to produce an anti-CD4 neutralizing antibody. Ammonium sulfate precipitation was performed on cell culture supernatants to produce a purified antibody, and a Bradford assay was performed to determine the protein concentration by using known concentrations of bovine serum albumin (Fisher Scientific) as standards and an ELISA plate reader (Bio-Tek Instruments, Winooski, VT). A rat Ig (Sigma-Aldrich) was used as an isotype control. BALB/c mice were either immunized orally with 10^6 CFU of LVS or mock immunized (PBS). Three weeks later, mice were injected intraperitoneally (i.p.) with either 0.25 μg of a monoclonal anti-CD4 antibody or a purified antibody, and a Bradford assay was performed to determine the protein concentration by using known concentrations of bovine serum albumin (Fisher Scientific) as standards and an ELISA plate reader (Bio-Tek Instruments, Winooski, VT). The oral LVS vaccine induced splenic antigen-specific IFN-γ responses in BALB/c mice by days 10 and 15, respectively (Fig. 1), and with Hoechst stain (Sigma). Images were acquired using a 510 Meta laser scanning confocal microscope (Zeiss) and were analyzed using Imaris software (Bitplane, Salt Lake, PA).

**Statistical analyses.** SigmaStat (Systat Software Inc., San Jose, CA) was used to perform all the tests of significance. Statistical analysis for survival experiments was performed using the Kaplan-Meier test, and the Student t test was used to determine differences in cytokine and antibody production. All data are reported as the mean ± standard error from each experimental animal group and are representative of at least two independent experiments.

**RESULTS**

**LVS Vaccination.** Oral LVS vaccination induces systemic and mucosal cellular and humoral immune responses. BALB/c mice were orally challenged with escalating doses (10^2 to 10^4 CFU) of LVS and were monitored daily for morbidity and mortality. These analyses revealed that the LD50 of LVS administered orally was approximately 10^4 CFU (data not shown). A challenge dose of 10^3 CFU of LVS delivered orally did not induce apparent morbidity in either BALB/c or C57BL/6 mice but induced 100% mortality in BALB/c IFN-γ−/− and C57BL/6 IFN-γ−/− mice by days 10 and 15, respectively (Fig. 1), consistent with previous studies demonstrating the importance of IFN-γ production in the initial control of parental LVS infection (1, 15, 32).

In addition, antigen-specific cell-mediated and humoral responses were characterized in BALB/c mice following oral LVS vaccination. Splenocytes were removed at day 14 after
oral immunization, and single cells were stimulated with 10^4 or 10^5 CFU of UV-inactivated LVS. These analyses revealed that antigen-specific IFN-γ production was significantly (P < 0.001) increased in a dose-dependent manner (0.326 ± 0.15 ng/ml and 0.762 ± 0.77 ng/ml, respectively) in splenocytes from mice orally vaccinated with LVS (Fig. 2A). Moreover, the cervical lymph node cells from orally vaccinated mice also produced an appreciable IFN-γ response in culture, in a dose-dependent manner (data not shown). In contrast, there was negligible induction of IFN-γ production in cells from mock-vaccinated mice and in cells from both groups that were incubated with the unrelated antigen HEL or with medium alone. Moreover, antigen-specific IL-2 production was also detected upon oral vaccination, but IL-4 production was not detectable in any of the cell cultures (data not shown).

Mice were bled 21 days following oral vaccination, and sera were analyzed for anti-LVS antibody responses. As shown in Fig. 2B, there was significant induction of total (3.948 ± 224; P < 0.001) and IgG2a (1.993 ± 641; P < 0.015) serum antibodies. In contrast, there was minimal induction of LVS-specific IgG1 and IgA in the sera. Anti-LVS antibody responses in mock-vaccinated mice were negligible. No binding of immune serum was detected in plates coated with HEL. To further determine the effects of oral LVS vaccination at inductive and distal mucosal sites (13), we examined antibody responses in fecal supernatants and BAL fluids collected from immunized animals. Oral LVS immunization induced significant (P < 0.001) IgA production but minimal IgM production in the intestines (Fig. 3A). No LVS-specific IgG was detected in fecal supernatants (data not shown). In the respiratory compartments of orally vaccinated animals, LVS induced significant levels of antibodies, including total antibodies, IgG1, IgG2a, and IgM (P < 0.001), and IgA (P < 0.015) (Fig. 3B). As expected, there was negligible induction of antibodies in the fecal supernatants and BAL fluids of mock-vaccinated mice. Collectively, these results demonstrate the efficacy of the oral route of vaccination in inducing antigen-specific systemic and mucosal cellular and humoral immune responses.

M cells located in the follicle-associated epithelium of intestinal tissue play an important role in the sampling and transport of antigens for processing and the initiation of immune responses (10, 28). To determine whether orally delivered LVS localizes to M cells following vaccination, we administered mCherry-labeled LVS and examined the gastrointestinal tract for the presence of these bacteria by confocal laser scanning microscopy. As shown in Fig. 4, M cells (green) were identified by staining with UEA-1 (34) and were apparent in the crypts of the small intestine. Interestingly, orally administered LVS cells expressing mCherry (red) were visible within the small intestine after 90 min and colocalized (yellow) to M cells. These results suggest that orally administered LVS may be trafficking to M cells for the initiation of mucosal immune responses.

Oral LVS vaccination induces robust protection against pulmonary and i.d. challenges with F. tularensis SCHU S4. To determine the efficacy of oral LVS immunization in conferring protective immunity, BALB/c mice were vaccinated with LVS (10^4 CFU) and were challenged i.n. or i.d. with F. tularensis SCHU S4 3 weeks later. As shown in Fig. 5A, mice orally vaccinated with LVS exhibited significant protection (100% survival with 126 CFU and 80% survival with 580 CFU) against i.n. pulmonary challenge during the monitoring period of 1 month. Additionally, oral LVS vaccination induced 100% pro-
tection against both infectious doses following SCHU S4 challenge administered i.d. (Fig. 5C), another common route of infection. All vaccinated mice exhibited minimal loss of body weight following type A bacterial challenge (Fig. 5B and D), and as expected, mock-vaccinated mice showed a rapid decline in body weight and succumbed to the infection by day 6 after challenge.

Since the duration of protective immunity against virulent type A strains induced by LVS immunization has been shown to be short (9, 30, 59, 61), we examined the extent of protection conferred by the oral vaccination regimen in this study. Given that oral LVS vaccination induced protective immunity against pulmonary SCHU S4 challenge after 3 weeks (Fig. 5), we evaluated the protective efficacy at 8 weeks postvaccination. Mice vaccinated orally with LVS (10³ CFU) were rested for 8 weeks and subsequently challenged i.n. with SCHU S4. As shown in Fig. 6A, immunized mice exhibited 33% and 50% survival rates (at 86 and 375 CFU, respectively) at this extended time following oral LVS vaccination, indicating a waning of protective immunity over time, with no significant difference in the survival rate between mice that received either dose. Therefore, some immunized mice were boosted with LVS (10³ CFU) orally after 8 weeks, rested for an additional 4 weeks, and challenged i.n. with SCHU S4. As shown in Fig. 6B, mice receiving a secondary boost of LVS orally were highly protected (80% survival with 86 CFU and 60% survival with 375 CFU) against i.n. challenge with SCHU S4. These analyses reveal that maintenance of long-term protective immunity by oral LVS vaccination may require additional booster immunizations.

To further evaluate the efficacy of the oral vaccination regimen, mice vaccinated with LVS (10³ CFU) or PBS were euthanized following pulmonary SCHU S4 challenge, and bacterial replication in the lungs, liver, and spleen was examined. As shown in Fig. 7A, small, comparable numbers of bacteria were recovered from the target organs of both LVS- and mock-
vaccinated animals 1 day after challenge. However, by days 3 and 4, mock-vaccinated animals exhibited extensive bacterial replication within the lungs (~10⁶ CFU), liver (10⁵ to 10⁷ CFU), and spleen (10⁵ to 10⁷ CFU). Moreover, as shown in Fig. 5A and B, the mock-vaccinated mice exhibited a significant loss of body weight and rapidly succumbed to the infection. In contrast, animals orally vaccinated with LVS exhibited significantly (P < 0.05) lower levels of recoverable viable bacteria in the target organs than mock-immunized animals during this initial period and up to day 14 (Fig. 7A). In parallel, histological analyses of the lungs were performed on both sets of animals at day 3 post-SCHU S4 pulmonary challenge. These experiments revealed that mock-vaccinated, SCHU S4-challenged mice exhibited minimal signs of cellular infiltration, with otherwise normal lung architecture (Fig. 7BI), like that of naive animals (Fig. 7BIII). The lung sections of mice orally vaccinated with LVS and challenged with SCHU S4 were generally comparable to those of mock-vaccinated animals, with the exception of foci of peribronchiolar mononuclear lymphocytic infiltration (Fig. 7BII). Collectively, these results demonstrate the efficacy of the oral vaccination route with LVS at inducing effective control of SCHU S4 replication and dissemination, presumably via the initiation of an early cellular influx into the primary site of infection.

Pulmonary immunity against SCHU S4 challenge is mediated by CD4⁺ T cells and antibodies, including IgA. Since the protective immunity against SCHU S4 challenge after oral LVS vaccination correlated with the early infiltration of mononuclear lymphocytes to the sites of infection, and given the demonstrated role of CD4⁺ T cells and IFN-γ in the control of Francisella infections (16, 63), we also examined the role of CD4⁺ T cells in orally vaccinated mice by treatment with an anti-CD4 neutralizing antibody (36), i.e. injection of the neutralizing anti-CD4 antibody markedly depleted splenic CD4⁺ T cells (0.5% of total splenocytes after treatment) in contrast to injection of a control rat Ig (17.6% of total splenocytes after treatment) (data not shown). As shown in Fig. 8, depletion of antigen-specific CD4⁺ T cells following pulmonary challenge with SCHU S4 (80 CFU) had a pronounced effect on the survival (25%) of the vaccinated animals in comparison to that of vaccinated animals not receiving the CD4⁺ T-cell depletion treatment (87%) or vaccinated mice injected with a control rat Ig (87%). As expected, all mock-immunized animals succumbed to the infection by day 6. These results suggest the significant contribution of CD4⁺ T cells to the protective immunity induced by oral LVS vaccination.

Apart from cellular immunity, antibodies have been shown to play a role in immunity against Francisella (26, 41, 44, 45, 53). Elevated levels of serum and mucosal antibodies were detected in mice after oral LVS vaccination (Fig. 3 and 4). IgG2a is the major murine isotype involved in the opsonization and phagocytosis of bacteria (2), while IgA has been shown to be the principal immunoglobulin isotype involved in the inhibition of bacterial attachment and the neutralization of viruses at mucosal surfaces (31). Moreover, there is evidence to suggest that IgA is required for the effective priming of T cells and the development of Th1 type immunity (3). To elucidate the role of humoral immunity in protection, μMT (B-cell-defi-
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Oral LVS vaccination was shown to induce fecal and respiratory antibodies, including IgA, and CD4^+^ T cells cooperatively play an important role in the induction of effective pulmonary immunity to *F. tularensis* challenge following oral LVS vaccination.

**DISCUSSION**

Vaccination strategies that effectively induce mucosal immunity and control *F. tularensis* growth and pneumonic and systemic disease are important considerations for the development of effective vaccines against tularemia. We have now reported, further validating the findings of Chen and colleagues (30), that the oral route of vaccination using LVS effectively induces mucosal and systemic immunity and confers significant protection against both respiratory and i.d. challenges with the type A strain *F. tularensis* S4. Moreover, the protective efficacy of the oral vaccination regimen may involve the antigen-sampling mechanisms of M cells within the intestinal tract and appears to be mediated by CD4^+^ T cells and antibodies, including IgA.

Recent studies have suggested that the oral route of vaccination with LVS may be preferential for inducing protective pulmonary immunity against human-virulent *F. tularensis* (30). Specifically, Chen and colleagues have shown that BALB/c mice vaccinated orally with 10^9^ CFU of LVS exhibited lower bacterial burdens than sham-immunized animals (30). Although protection in their study was seen with challenges up to 10^9^ CFU of LVS in PBS or mock immunized; 3 weeks later, they were challenged i.n. with either 50 CFU (A) or 102 CFU (B) of *F. tularensis* SCHU S4; then they were monitored daily for survival. Differences in survival between immunized wild-type and B-cell-deficient mice at a challenge dose of 50 CFU were significant at a *P* value of <0.01 (statistical power, 0.996 with an alpha of 0.50). Results are representative of two separate experiments.

**FIG. 8.** Contribution of LVS-specific CD4^+^ T cells to protective immunity against SCHU S4 challenge. Groups of BALB/c mice (*n* = 8) were either vaccinated orally with 10^9^ CFU of LVS or mock immunized with PBS. Mice were rested for 3 weeks and received i.p. injections of either an anti-CD4 neutralizing antibody, a control rat Ig, or PBS at day −2, day −1, the day of i.n. challenge with 80 CFU of SCHU S4 (day 0), and every subsequent third day. Mice were monitored daily for morbidity and mortality. Differences in survival between vaccinated mice receiving rat Ig and those receiving anti-CD4 antibody treatment were significant at a *P* value of 0.0185 (statistical power, 0.996 with an alpha of 0.50). Results are representative of two separate experiments.

**FIG. 9.** Contribution of antibodies, including IgA, to protective immunity against SCHU S4 challenge. (A and B) Groups (*n* = 6) of wild-type and B-cell-deficient (μMT) C57BL/6 mice were either vaccinated orally with 10^8^ CFU of LVS or mock immunized with PBS. Three weeks later, mice were challenged i.n. with either 50 CFU (A) or 102 CFU (B) of *F. tularensis* SCHU S4; then they were monitored daily for survival. Differences in survival between immunized wild-type and B-cell-deficient mice at a challenge dose of 50 CFU were significant at a *P* value of <0.01 (statistical power, 0.996 with an alpha of 0.50). Differences in survival between LVS- and mock-immunized mice were significant at a *P* value of <0.001 (statistical power, 0.996 with an alpha of 0.50). (C) Groups (*n* = 6) of wild-type and IgA^−/−^ C57BL/6 mice were either immunized orally with 10^9^ CFU of LVS in PBS or mock immunized; 3 weeks later, they were challenged i.n. with 98 CFU of *F. tularensis* SCHU S4. Differences in survival between LVS-immunized wild-type and IgA^−/−^ mice were significant at a *P* value of 0.0179 (statistical power, 0.996 with an alpha of 0.50). Results are representative of two separate experiments.

Oral LVS vaccination was shown to induce fecal and respiratory IgA responses (Fig. 3) that may contribute to protective immunity against mucosal pathogens. To evaluate the contribution of IgA to protective immunity, we vaccinated IgA^−/−^ and wild-type C57BL/6 × 129 mice orally with LVS and challenged these animals i.n. with 98 CFU of SCHU S4. IgA^−/−^ mice exhibited 50% survival, while 87% of the corresponding wild-type animals were protected against bacterial challenge (Fig. 9C). As expected, all mock-immunized animals in both experiments succumbed to the infection by day 8 after challenge (Fig. 9). These results further suggest that IgA may contribute significantly to the protective immunity mediated by antibodies in general. Collectively, these results suggest that
50 CFU i.n. and 20 CFU by aerosol exposure, mice challenged i.n. with higher inocula were not protected by oral LVS vaccination. The primary differences between the previously reported study and this study include the oral vaccination doses (10^6 CFU versus 10^7 CFU, respectively) of LVS and the type A strains (FSC33/snMF [strain FSC033] versus SCHU S4, respectively) used for challenges. Various studies (12, 49, 61) have shown significant differences in the virulence of LVS that may arise from culture with different media or different growth conditions, as well as from differences in the source of the strain. The strain of LVS used by Chen et al. was acquired from a source different from that used for this study, which may account for the differences in the oral LD50 between the studies. Nevertheless, evidence from both independent studies clearly indicates the feasibility of the oral route of vaccination in inducing significant respiratory immunity against virulent *F. tularensis* type A strains. Given the successful history of oral vaccines for humans, this immunization route may be highly viable for inducing both systemic and mucosal immune protection against *F. tularensis*. For example, oral vaccination with a *Salmonella enterica* delivery system expressing *Yersinia pestis* F1 and V antigens has been reported to be protective against bubonic and pneumonic plague (62). The efficacy of the oral vaccination regimen in protection against pulmonary tularemia may be a result of the effective delivery of vaccine antigens to M cells, which are located in the follicle-associated epithelium of Peyer's patches (10, 28, 29). M cells have been shown to play an important role in the sampling and uptake of luminal antigens (46) and to play a role in the release of costimulatory signals for effective induction of T- and B-cell proliferation (42). In this regard, Kiyono and colleagues (38) have recently shown the feasibility of targeting vaccine antigens to the M-cell-specific carbohydrate moiety as a highly effective strategy for inducing mucosal immunity. To this end, the uptake of microorganisms and microparticles from the small intestine may occur both through the M cells of Peyer's patches (35) and through intestinal villous M cells, described recently (24), as well as by an alternative mechanism of villous transepithelial passage, originally termed persorption (57). Early studies have shown that oral administration of suspensions of a large variety of different solid particles, the size of microorganisms and larger, to animals and human volunteers resulted in passage in less than an hour from the small intestine through the lymphatic and portal systems to the peripheral blood and a variety of body organs (57). Given that larger inocula of bacteria may result in greater systemic spread and induce some degree of morbidity themselves, the size of the immunizing oral LVS inoculum may be an important consideration and may affect the protective efficacy of the vaccination.

Effective mucosal defenses have been shown to be mediated by both cell-mediated and humoral mechanisms that operate in concert at major portals of entry for microorganisms (6). In the respiratory system, distinct mechanisms may be involved in the clearance of bacteria from the upper airways and deeper alveolar spaces. To this end, phagocytic cells such as macrophages and neutrophils may be involved in the removal of microorganisms that reach the deeper alveolar spaces by cognate interaction with antibodies through Fc-receptor-mediated processes (43). Whereas infection with SCHU S4 provokes a minimal inflammatory response in the lungs early after pneumonic challenge, as seen in this study and others (5, 8), the lungs of orally vaccinated and challenged mice exhibited an increase in the number of inflammatory cells, which were primarily lymphocytic. This influx of lymphocytes, which was evident only in vaccinated mice, may have contributed to the effective local control of bacterial replication. Moreover, depletion of antigen-specific CD4+ T cells at the time of infection remarkably abrogated the protective effects of oral LVS vaccination, indicating the importance of this cell type and of the production of cytokines such as IFN-γ for optimal bacterial clearance and protection against i.n. *Francisella* challenge. Oral LVS vaccination induced significant levels of antibodies in the respiratory compartment. LVS-mediated protection against pulmonary SCHU S4 challenge was also partially abrogated in the absence of B-cell and IgA expression. Antibodies have been shown by us (41, 44) and others (26, 53) to play an important role in the control of pulmonary *Francisella* infection. The mechanisms by which antibodies may facilitate the control of bacterial replication may include the neutralization of infectious organisms and Fc-receptor-mediated killing (44, 45). Both of these mechanisms may act in concert during an infection and limit the early dissemination of the organism, given that *Francisella* bacteremia occurs both in intra- and extracellular phases (19, 64).

The protection conferred by oral LVS vaccination began to wane by 2 months. Both the magnitude of the antibody responses and that of the antigen-specific cell-mediated IFN-γ response in vaccinated mice were reduced by factors of 2 and 4, respectively, by 2 to 3 months postimmunization (H. J. Ray and B. P. Arulananandam, unpublished data). A similar waning of LVS-mediated immunity against pulmonary tularemia has been reported previously (9, 30, 59, 61), following immunization by different routes. Given that the correlates of protective immunity against SCHU S4 have yet to be defined, the question of the long-term efficacy of LVS vaccination in the mouse model remains to be resolved. However, we have now shown that an additional boost of LVS given orally can be used to maintain protective immunity for an extended period.

In summary, with the significant interest in the development of a licensed vaccine for use against *F. tularensis*, consideration also has to be given to routes of delivery that induce optimal immunity at sites of infection. The advantages of an oral vaccine include (i) the ease of delivery, (ii) the possibility of fewer adverse effects than those with parenteral injection of dead whole or subunit vaccines, and (iii) the effective induction of both systemic and mucosal immunity, particularly in the upper respiratory system. While LVS continues to be used only to treat certain at-risk individuals, it may be unlikely to be licensed for use in the general population with the current level of understanding of the exact conditions under which it was generated, the mutations responsible for its attenuation, and the residual (dose-dependent) morbidity and mortality (39). However, LVS is a useful organism to be used in animal models for the evaluation of immune mechanisms that confer protective immunity, particularly against the virulent type A *Francisella* strains. Further studies to determine the efficacy of the oral vaccination route with defined attenuated *Francisella* vaccine strains and in other animal models of pulmonary tularemia are warranted and are currently under development.
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