Antigen-Specific Cellular and Humoral Responses Are Induced by Intradermal *Mycobacterium leprae* Infection of the Mouse Ear


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Leprosy is caused by infection with *Mycobacterium leprae*. The immune response of leprosy patients can be highly diverse, ranging from strong cellular responses accompanied by an apparent deficit of *M. leprae*-specific antibodies to strong humoral responses with a deficit of cell-mediated responses. Leprosy takes many years to manifest, and this has precluded analyses of disease and immune response development in infected humans. In an attempt to better define development of the immune response during leprosy we have developed an *M. leprae* ear infection model. Intradermal inoculation of *M. leprae* into the ear supported not only infection but also the development of a chronic inflammatory response. The inflammatory response was localized, comprising a T-cell infiltration into the ear and congestion of cells in the draining lymph nodes. The development of local chronic inflammation was prevented by rifampin treatment. Importantly, and in contrast to subcutaneous *M. leprae* footpad infection, systemic *M. leprae*-specific gamma interferon and antibody responses were detected following intradermal ear infection. These results indicate the utility of intradermal ear infection for both induction and understanding of the immune response during *M. leprae* infection and the identification or testing of new leprosy treatments.
that experimental dermal infection may better mimic typical human infection (6–8). Our data indicate that *M. leprae* bacilli not only grow within the ears but also stimulate a rapid and prolonged local inflammatory response. The inflammatory response presents as a T-cell infiltrate within the ear and a local lymphadenopathy, both of which are limited by treatment with the antimycobacterial drug rifampin. In addition, and in contrast with mice infected in the footpad, mice infected in the ear demonstrate *M. leprae*-specific cellular responses. Our data indicate that *M. leprae* infection of the mouse ear provides a system with which to evaluate antileprosy treatments and analyze the development of *M. leprae*-induced inflammatory responses.

**MATERIALS AND METHODS**

*M. leprae* inoculations and rifampin treatment. Live *M. leprae* bacilli were obtained by heating bacilli at 70°C for 1 h and then quenching on ice. Mice were inoculated with bacilli in a volume of 10μl i.d. injection into the ear pinnae or subcutaneous (s.c.) injection into the footpad. To assess growth, both ears were harvested and the bacilli were enumerated by direct microscopic counting of acid-fast bacilli according to the method of Shepard and McRae (35). In treatment experiments, mice were injected intraperitoneally with 0.5 mg rifampin (Sigma) or phosphate-buffered saline (PBS) at 1, 2, and 3 weeks after infection.

Histology. Ears were fixed in formalin and sectioned. Slides were then stained with hematoxylin and eosin.

Cell preparations. Single-cell suspensions were prepared from the spleen and lymph nodes (LN; auricular, axillary, inguinal, and popliteal). Spleens and LN were disrupted between frosted slides and erythrocytes removed by lysis in 1.66% NH4Cl solution. Single-cell suspensions were also prepared from ears. Ears were collected, rinsed with 70% ethanol, and allowed to air dry. Ears were then split into dorsal and ventral halves and floated on 1 ml RPMI 1640 (BioWhittaker, Walkersville, MD) supplemented with Liberase CI (Sigma, St. Louis, MO) for 1.5 h at 37°C. Enzymatic digestion was stopped by the addition of 1 ml RPMI 1640 supplemented with 0.05% DNase (Sigma). Digests were homogenized in 50 μm Medicon filters using a MediMachine (BD Biosciences, San Jose, CA), and the resultant homogenate was filtered through a 70-μm Filcon filter (MVF Filters, Walkersville, MD) supplemented with Liberase CI (Sigma, St. Louis, MO) for 1.5 h at 37°C. Enzymatic digestion was stopped by the addition of 1 ml RPMI 1640 supplemented with 0.05% DNase (Sigma). Digests were homogenized in 50 μm Medicon filters using a MediMachine (BD Biosciences, San Jose, CA), and the resultant homogenate was filtered through a 70-μm Filcon filter (BD Biosciences). Ear, LN, and spleen cells were washed at least twice before enumerating mononuclear cells using a hemocytometer.

Flow cytometry. To stain for flow cytometry, cells were incubated with the anti-FcyRIIIR antibody 2.4G2 and then with combinations of fluorescently conjugated antibodies (all BD Biosciences) for 30 min on ice. Cells were then washed and analyzed with WinMDI 2.8.

**RESULTS**

Short-term assessment of inflammatory responses during *M. leprae* infection. To establish if ears can support *M. leprae* infection and be used to develop a more rapid system with which to investigate experimental leprosy, we inoculated mice in the ears with *M. leprae* bacilli. *M. leprae* bacilli could not be detected in ears collected 10 weeks after infection (detectable bacilli were recovered from only 12.5% ± 5% of the inoculated ears), whereas significantly more bacilli were counted in ears collected 20 weeks (62.5% ± 5%, *P = 0.028) or 30 weeks (90% ± 5%, *P = 0.123) after infection (Fig. 1A). Histological analyses of *M. leprae*-infected ears revealed that bacilli could be observed within the dermal layer, along with significant numbers of infiltrating cells (Fig. 1B). These data indicate that the ear can support *M. leprae* growth as well as a local immune response.

To investigate pathogenesis and immune responses during leprosy, we analyzed local inflammation during *M. leprae* ear infection in relatively short-term experiments (12 to 15 weeks, compared to 6 to 12 months required following footpad inoculation). Draining LN (DLN) from infected mice contained significantly more cells than distal LN from the same infected mice or LN from uninoculated mice (week 0), with DLN cell numbers progressively increasing during the early weeks of infection (Fig. 1B). In contrast to the DLN, the distal LN and spleens of infected and uninfected mice were similar in size and cell number (Fig. 1B). These data indicate that inoculation of *M. leprae* into the ear dermis stimulates a local inflammatory response.

To contrast the ear and footpad routes of infection, mice were infected by i.d. inoculation of bacilli into the ear or s.c. inoculation of bacilli into the footpad. Fifteen weeks later DLN cell numbers in mice inoculated via either route were increased (*P < 0.001 for either ear- or footpad-infected mice versus uninoculated mice; Fig. 1C). The increase, however, was greater following ear infection than footpad infection (*P = 0.007; Fig. 1C). Experiments were also conducted with an additional group of mice inoculated in the ears with heat-killed *M. leprae*. Although DLN from heat-killed *M. leprae*-inoculated mice were also enlarged, they contained significantly less...
cells than DLN from infected mice (Fig. 1C; P = 0.001). These observations indicate that infection of the ear stimulates a larger inflammatory response than infection of the footpad and that the magnitude of DLN hyperplasia is related to infection.

**DLN cell populations in infected mice.** To examine if the increase in DLN cell numbers in infected mice was due to the expansion of all LN cells or the selective expansion of a particular cell type, DLN cell composition was determined by flow cytometry. These investigations indicated that, compared with either distal LN from infected mice or LN from uninfected mice, there were no dramatic alterations in the DLN cell composition of *M. leprae*-infected mice (data not shown). Rather, the numbers of all cell types examined (αβ CD4 and CD8 T cells, dendritic cells, granulocytes, and macrophages) significantly increased during infection (Fig. 2). While the expansion of most cell types occurred early during infection and was maintained as infection progressed, the number of granulocytes peaked at 3 weeks following infection and then slowly declined (Fig. 2F). Vβ chain use within the T-cell populations was not altered (data not shown), demonstrating that selective outgrowth of particular T cells during infection did not occur. DLN cells capable of producing IFN-γ also increased in number during infection (Fig. 2G). These data indicate that *M. leprae* ear infection stimulates a prolonged expansion of cells, including IFN-γ-producing cells, within DLN.

**Cell infiltration of *M. leprae*-infected ears.** To more closely examine the site of infection, cells were extracted from *M. leprae*-infected ears. Infected ears yielded significantly more mononuclear cells than uninfected ears (Fig. 1A; P = 0.001). These observations indicate that infection of the ear stimulates a larger inflammatory response than infection of the footpad and that the volume of DLN hyperplasia is related to infection.

**FIG. 1.** Local lymphadenopathy following inoculation in each ear with *M. leprae* bacilli. (A) Mice were infected with 5 × 10⁶ *M. leprae* bacilli, and the numbers of *M. leprae* bacilli in 8 to 12 infected ears (four to six mice) at each time were determined. Each point represents the number of bacilli detected in an individual ear, with the bar representing the mean number of bacilli at each time and the dashed line indicating the limit of detection of this assay. (B) Hematoxylin- and eosin-stained sections of ears infected for 30 weeks. Some of the *M. leprae* bacilli are indicated by the arrows. (C) LN (each mouse with two DLN or six distal nodes) and spleen cell numbers are shown as the means for three mice per infected time or of eight total mice (two per time) to derive an uninfected value. *, P < 0.05; **, P < 0.01. (D) Additional mice were inoculated in the ears with 1 × 10⁶ heat-killed *M. leprae* (HKML) or in the footpads with 1 × 10⁶ live *M. leprae* bacilli, and DLN were examined 15 weeks later. Cell numbers are shown as the means and standard errors of the means for two groups comprising four to six DLN. Results are representative of three to five similar experiments.
cells increased as infection progressed, with CD4 T cells outnumbering CD8 T cells approximately 2 to 1 (Fig. 3). Many of the infiltrating T cells displayed an activated phenotype (CD44\textsuperscript{hi}; data not shown) and produced IFN-\gamma, but not IL-10, upon mitogen stimulation (Fig. 3A). The number of IFN-\gamma-producing cells in the ears increased as infection progressed (Fig. 3I). These results indicate that \textit{M. leprae} infection of the ear promotes an influx of IFN-\gamma-producing T cells and stimulates a persistent local inflammatory response.

FIG. 2. T-cell expansion within DLN during \textit{M. leprae} infection. Mice were infected in the ear with $1 \times 10^6$ live \textit{M. leprae} bacilli, LN were collected at various times thereafter, and single-cell suspensions were subjected to flow cytometry to determine and enumerate various cell types. (A) $\alpha\beta$ T cells (TCR\textsuperscript{\alpha\beta}); (B) CD4 T cells (CD4\textsuperscript{+}, TCR\textsuperscript{\alpha\beta}); (C) CD8 T cells (CD8\textsuperscript{+}, TCR\textsuperscript{\alpha\beta}); (D) dendritic cells (DC) (CD11c\textsuperscript{+}); (E) macrophages (F4/80\textsuperscript{+}); (F) granulocytes (granulo) (Ly6G/C\textsuperscript{+}); and (G) IFN-\gamma-producing cells (IFN-\gamma following PMA-ionomycin stimulation). Results are shown as the means and standard errors of the means for three mice per time and are representative of two individual experiments. *, $P < 0.05$; **, $P < 0.01$ (versus uninfected or distal LN).

FIG. 3. T cells infiltrate the ear during \textit{M. leprae} infection. Mice were infected in the ear with $1 \times 10^6$ live \textit{M. leprae} bacilli, and ears were collected at regular intervals to provide single-cell suspensions. Cell suspensions were subjected to flow cytometry. (A) Representative flow cytometry plots. Various cell types were enumerated as (B) total cells, (C) $\alpha\beta$ T cells (TCR\textsuperscript{\alpha\beta}), (D) CD4 T cells (CD4\textsuperscript{+}, TCR\textsuperscript{\alpha\beta}), (E) CD8 T cells (CD8\textsuperscript{+}, TCR\textsuperscript{\alpha\beta}), (F) dendritic cells (DC) (CD11c\textsuperscript{+}), (G) macrophages (F4/80\textsuperscript{+}), (H) granulocytes (Ly6G/C\textsuperscript{+}), and (I) IFN-\gamma-producing cells (IFN-\gamma following PMA-ionomycin stimulation). Results are shown as the means and standard errors of the means for three mice per time (results for uninfected mice are plotted at week 0) and are representative of two or three similar experiments. *, $P < 0.05$; **, $P < 0.01$ (versus uninfected mice).
Rifampin treatment prevents *M. leprae*-induced local inflammation. Having established that *M. leprae* infection of the ear causes local inflammation, we wanted to determine if this inflammation could be interrupted. Rifampin has antimycobacterial properties, and this drug is commonly used to treat leprosy (5, 14, 23). We hypothesized that rifampin treatment would prevent bacillus growth and therefore limit development of local inflammation. Mice were infected in the ears and treated with rifampin shortly after infection. In agreement with our hypothesis, although not significant in this experiment, fewer cells were recovered from DLN of infected, rifampin-treated mice than from DLN of infected, untreated mice (Fig. 4A). The DLN cell numbers for infected, rifampin-treated mice were similar to those for heat-killed *M. leprae*-inoculated mice (Fig. 4A). Similarly, when the ear cell infiltrates of these mice were examined, we found that infected, rifampin-treated mice had a significantly reduced T-cell infiltration compared to infected, untreated mice (Fig. 4B). Taken together, these data indicate that the local inflammation observed following *M. leprae* ear infection can be limited by antimycobacterial drug treatment.

**Infection of the ear induces specific anti-*M. leprae* responses.** Infection of the ear stimulated a larger local inflammatory response than infection of the footpad, suggesting that ear infection is more immune stimulatory than footpad infection. To further test this observation, spleen cells from the *M. leprae*-infected mice were incubated with *M. leprae* cell wall antigens and the culture supernatants analyzed. In these experiments, spleen cells from mice infected in the footpad did not exhibit *M. leprae*-specific IFN-γ recall responses 15 weeks after infection (Fig. 5A). In marked contrast, spleen cells from mice infected in the ear produced IFN-γ, but not IL-4, in response to *M. leprae* cell wall antigens 15 weeks after infection (Fig. 5A and data not shown). While spleen cells from mice inoculated in the ears with heat-killed *M. leprae* also exhibited IFN-γ recall responses, these responses were significantly lower than the responses of the infected mice (*P* < 0.01). These results demonstrate that, following ear infection but not footpad infection, specific anti-*M. leprae* cellular responses are induced. We explored the antigen specificity of these responses by stimulating cells from mice infected in the ears with the recombinantly expressed *M. leprae* antigen ML2028. Cells from mice infected in the ears, but not from mice infected in the footpad or inoculated with heat-killed *M. leprae*, secreted IFN-γ in response to ML2028 (Fig. 5A). Thus, infection in the ear primes antigen-specific T cells more easily than infection in the footpad.

Having demonstrated that mice infected in the ear with *M. leprae* develop local T-cell inflammation and systemic cellular responses, we questioned if these mice also developed anti-*M. leprae* humoral responses. Sera were collected during the early stages of infection, and antibody responses were assessed by...
ELISA. We could not detect IgM responses against PGL-I (data not shown), in agreement with previous results demonstrating that this *M. leprae* glycolipid is a poor immunogen in B6 mice (20a). Weak, but significant, IgG responses against *M. leprae* cell wall antigens and ML2028 were detected in sera from infected mice, and these responses increased as infection progressed (Fig. 5). Due to the low antibody responses, as measured by ELISA, after 6 weeks of infection, we also enumerated the number of long-lived ASC in the bone marrow of infected mice. More anti-ML2028-secreting plasma cells could be found in the bone marrow of infected mice than in that of uninfected mice (Fig. 5B). These data indicate that, in addition to T-cell priming, successful B-cell priming and antibody production occur following *M. leprae* infection of the ear.

**DISCUSSION**

Leprosy exhibits highly divergent immune responses and pathologies (28, 30). Little is known as to what influences the differential development of the strong cellular response of PB patients or the weak/absent cellular response of multibacillary patients. Animal models of leprosy could provide important insight, but the lack of animal models that demonstrate robust immune responses during *M. leprae* infection has limited the ability to understand leprosy pathogenesis. The *M. leprae* mouse footpad infection model was developed by Shepard in the 1960s to assess bacterial growth, but this model requires a significant length of time and demonstrates only minor pathology (32–34). Recent studies of experimental cutaneous leishmaniasis have indicated that i.d. inoculation of *Leishmania* major into the ear promotes disease symptoms (6–8). We report here that following *M. leprae* infection of the ear there is a marked DLN hyperplasia and a cell infiltration at the inoculation site, indicating the induction of inflammatory responses. In agreement, mice infected by i.d. inoculation in the ear also produce anti-*M. leprae* antibodies. Thus, following ear infection, both B- and T-cell responses are rapidly detected, indicating that the ear model can provide insight into leprosy pathogenesis and development of the immune response.

Mice infected by i.d. inoculation into the ear developed systemic IFN-γ responses against *M. leprae* antigens (ML2028 and *M. leprae* cell wall antigens). This was in marked contrast with mice infected by s.c. inoculation in the footpad, which did not demonstrate IFN-γ recall responses. Several recent studies...
investigating experimental cutaneous leishmaniasis have identified differences in the development of L. major infection when infection of the ear is compared with infection of the footpad (6–8). It is unclear why there are differences in the resulting immune responses of mice infected in these different sites. One possibility is that the presence of different antigen-presenting-cell populations in the ear and footpad results in different qualities of M. leprae antigen presentation (9), and this is worthy of further examination.

PB leprosy patients demonstrate well-defined skin lesions that, upon biopsy, demonstrate a granulomatous inflammation comprising more CD4 than CD8 T cells (25). Mice infected with M. leprae in the ears quickly developed a localized inflammatory response comprising DLN hyperplasia and recruitment of more CD4 T cells than CD8 T cells to the inoculation site. Although macroscopic lesions were not observed on M. leprae-infected ears, the cell types of the ear infiltrate are consistent with those of PB leprosy lesions. It remains to be determined if these cells form granulomas within the infected ears, but these results indicate that the ear infection model can be used to delineate the critical mediators of local inflammatory responses during leprosy.

The early treatment of M. leprae infection is critical for prevention of tissue and nerve damage. The ear infection model described here provides results more rapidly than the footpad model, which typically investigates bacillus growth and is currently used for the testing of M. leprae drug candidates (19). We have demonstrated that rifampin can be used to limit the local inflammation that develops following ear infection. It is likely that M. leprae ear infection can be used to more rapidly assess other antimycobacterial drugs that could be used alone or in conjunction with current MDT protocols for the treatment of leprosy. In addition, we have demonstrated immune responses to both crude M. leprae cell wall antigens and the recombinant antigen ML2028 following M. leprae ear infection. ML2028 induces the M. leprae homolog of M. tuberculosis Ag85B, a secreted antigen that has previously been demonstrated to confer protection against experimental leprosy infection (26). Thus, the ear infection system may be invaluable for the identification of additional M. leprae antigens that can be used in a vaccine to protect against M. leprae infection.

The results reported here demonstrate several immune parameters (DLN enlargement, ear infiltration, cellular responses, and antibody responses) that can be analyzed relatively rapidly following M. leprae ear infection and used to improve our understanding of the immune response during leprosy. These parameters will also be helpful for the identification and assessment of new treatments for leprosy.

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