Corneal ulceration as a result of bacterial infection is a potentially devastating disease which may lead to permanent scarring of the cornea and loss of visual acuity or vision. The pathogenesis is considered to be multifactorial and includes numerous bacterial proteases, toxins, and other virulence factors as well as mediators produced by a host’s own inflammatory responses (17, 32). Pseudomonas aeruginosa is a frequently isolated pathogen from bacterial keratitis and accounts for 70% of soft contact lens-associated cases (31). Once infection is initiated it is often difficult to control because of its progressive nature and/or the possible resistance to antibiotics of the infecting bacteria. Even if the infection responds to antibiotics, inflammation can persist. Polymorphonuclear leukocytes (PMNs) are the major inflammatory cells that migrate into the corneal stroma early after the onset of infection (16). Although PMNs are required for the removal of viable bacteria from the tissue, their continued presence may lead to extensive corneal damage.

Protective mechanisms against bacterial infection may include recruitment of phagocytic cells, specific B- and T-cell responses, and the presence of antigen-specific antibodies. Previous studies using passive transfer of monoclonal antibodies to outer membrane proteins of P. aeruginosa and immune sera produced during corneal infection have shown that passive immunization can provide partial protection against infection (26, 38). Similarly, active immunization with lipopolysaccaride and elastase can protect the cornea to some degree against bacterial infection (19). Immunization via nonocular routes (subcutaneous and intraperitoneal) with peptide antigens of herpes simplex virus has been shown to protect mice against corneal challenge with herpes simplex virus (14). These studies suggest that considerable protection can be achieved by manipulating the formulation of vaccines and immunization routes and schedules. However, effector mechanisms of immunity against P. aeruginosa infection in the eye remain poorly understood. Thus, understanding effector mechanisms can help in designing strategies for better management of sight-threatening corneal inflammation.

Cytokines play an important role in inflammatory and immune responses. They have both beneficial and detrimental influences. Various cytokines have been shown to enhance immunoglobulin A (IgA) antibody responses, especially the immunosuppressive cytokines interleukin-4 (IL-4), IL-10, and transforming growth factor beta (7). IL-5 and IL-6 induce IgA-committed B cells to terminally differentiate into IgA plasma cells (3). Synthesis and secretion of the secretory component is stimulated by tumor necrosis factor alpha and -beta, IL-1α, and IL-1β (15). On the other hand, proinflammatory cytokines produced during bacterial infection regulate PMN recruitment by inducing chemokines. Recent studies have shown that IL-1β and macrophage inflammatory protein 2 (murine IL-8 homolog) are major cytokines involved in the direct and indirect recruitment of PMNs (18, 29). Inocorneal infections with P. aeruginosa, the host’s own inflammatory response is primarily derived from stimulated PMNs (32), and
the inappropriate production of inflammatory cytokines possibly contributes to corneal damage. Effective immunization should protect the host not only by facilitating effective removal of bacteria but also by controlling the inflammatory process through appropriate cytokine expression and release.

The purpose of this study was to evaluate the various routes (ocular topical [OT], oral, nasal, and intra-Peyer’s patch [IPP]) that can provide significant protection against *P. aeruginosa* keratitis. Further, we attempted to define the mechanisms involved in protection against acute bacterial ocular infections.

**MATERIALS AND METHODS**

**Animal model.** Sprague-Dawley (inbred) rats of 10 to 12 weeks of age were used in this study. Eye swabs were taken from each rat for bacteriological culture prior to the study, and rats that were not carrying *P. aeruginosa* were used. Baseline measurements of corneal integrity that included slit lamp biomicroscopy were performed on all rats.

**Bacterial strain and growth conditions.** The cytotoxic strain 6206 of *P. aeruginosa* was used. Strain 6206 was isolated from a human corneal ulcer and classified as a cytotoxic strain on the basis of its interaction with corneal epithelial cells in vitro (8). Bacteria were grown in 10 ml of tryptone soy broth (Oxoid Ltd., Sydney, Australia) overnight at 37°C, harvested and washed three times in sterile phosphate-buffered saline (PBS), and resuspended in PBS prior to use.

**Vaccine.** Vaccine was prepared by exposing *P. aeruginosa* strain 6206 (2 × 10^10 CFU/ml) to 1% (wt/vol) paraformaldehyde (Sigma Chemical Co., Sydney, Australia) in PBS (pH 7.4) for 2 h at 37°C. After incubation, bacteria were washed three times in sterile PBS. For oral, nasal, and OT immunization, paraformaldehyde-killed bacteria were suspended in PBS to a concentration of 2 × 10^10 CFU/ml. Paraformaldehyde-killed bacteria emulsified at a 1:1 ratio with incomplete Freund’s adjuvant (Pierce, Sydney, Australia) were used to immunize rats via their intestinal Peyer’s patches.

**Immunization.** The primary mucosal immunization protocols were described elsewhere (9). In this study the following four immunization schedules were included: (i) combined IPP-OT immunization, (ii) combined oral-OT immunization, (iii) combined nasal-OT immunization, and (iv) OT immunization only. The OT immunization was included because local booster doses have been shown to be necessary for an optimal response in other systems (36). For each immunization group, 16 rats (3 animals for histology, 3 for enzyme-linked immunosorbent assays [ELISAs] and bacterial counts, 3 for PMN quantitation, 3 for lymphocyte proliferation assay [mesentric lymph nodes] and antigen-specific antibody detection [blood and tears], and 4 for mRNA quantitation) were used at each time point. Test groups were anesthetized by inhalation of isoflurane (Cenvet, Sydney, Australia).

**i) Peyer’s patch immunization.** The delivery procedure involved performing a laparotomy to expose the small intestine and delivery of a small volume (50 µl) of the inoculum subserosally to each Peyer’s patch. The incision was closed by suturing the abdominal wall and skin.

(ii) **Oral immunization.** Daily doses (2 × 10^10 CFU/ml) of paraformaldehyde-killed bacteria suspended in PBS were administered on days 1 to 5 and then days 10 to 14 in a 0.5-ml volume via an infant feeding tube.

(iii) **Nasal immunization.** Killed bacteria (2 × 10^10 CFU/ml) suspended in PBS were administered on days 1 to 3 and then days 7 to 10 intranasally in a volume of 0.2 ml.

(iv) **OT booster dose.** The tear fluid was blotted from the corner of the eye, and 5 µl of vaccine (paraformaldehyde-killed bacteria suspended in PBS) was delivered onto the corneal surface on the 7th day after completion of oral and nasal immunization and 14 days after IPP immunization.

**Animal infection.** After completion of the immunization schedule and 7 days postbooster, rats were anesthetized and the left and right corneas were scratched using a 26-gauge needle. Left scratched corneas were challenged topically with 2 × 10^4 live bacteria (*P. aeruginosa* strain 6206) in a 5-µl dose, while the right eyes served as scratch controls. **Clinical examination.** Anesthetized animals were examined at 4, 8, and 24 h and 3, 5, and 7 days postinfection using a slit lamp biomicroscope to grade the severity of infection. The following anterior segment variables were assessed: (i) corneal infiltrate density, grades 0 to 4, where 0 corresponds to none, 1 corresponds to very slight (irid detail visible), 2 corresponds to slight (irid detail partly obscured), 3 corresponds to moderate (irid detail not visible), and 4 corresponds to severe (opaque); (ii) depth of infiltrates, 0 to 100%, where 100% means the full corneal thickness shows infiltrates; (iii) extent of infiltrates, 0 to 100%, where 100% corresponds to full corneal coverage; (iv) epithelial defect size, 0 to 4 mm, where 4.0 mm means full epithelial loss; (v) epithelial defect depth, 0 to 100%, where 100% means a defect involving the full epithelial thickness; and (vi) edema severity, 0 to 4, where 0 corresponds to none, 1 corresponds to very slight, 2 corresponds to slight, 3 corresponds to moderate, and 4 corresponds to severe.

The anterior chamber reaction was graded on the basis of cells (grades 0 to 4), flare (grades 0 to 4), fibrinotic membrane presence or absence, hypopyon presence or absence, and hyphema presence or absence. A composite corneal disease score was derived from the sum of the first five variables and a maximum total corneal score would be the total of each grade for each variable (i.e., 20).

**Antigen-specific IgG and IgA detection by ELISA.** Animals were examined for an antibody response for 3 weeks after immunization. Eye wash or blood samples were collected each week to monitor the effect of the vaccine. Rats were bled from the lateral tail vein once per week after immunization to detect IgG in serum. Tears were collected by washing eyes with 20 µl of PBS (pH 7.4) to detect ocular IgA. Specific antibody to *P. aeruginosa* was measured by ELISA. Bacterial antigen was prepared from bacteria grown overnight on 10 nutrient agar plates. Cells were collected, washed, and resuspended in 5 ml of PBS. Suspended bacteria were sonicated using a small probe assembly. Sonication (Branson Sonifier 250; Branson Ultrasonics Corp., Danbury, Conn.) was performed with the amplitude set at 6 µ for 3 cycles of 30 s each one. Sonicated bacteria were centrifuged at 10,000 × g for 15 min, and the supernatant was used as a crude polyclonal antigen. ELISA plates were coated by adding 100 µl of polyclonal antigen diluted 1:1,000 in carbonate and bicarbonate buffer (pH 9.6) and were incubated at 4°C overnight. After 4 cycles of incubation and blocking with blocking buffer (5% skim milk and 0.05% Tween 20), Diluted serum or eye wash (control sera, 1:100; immune sera, 1:1000; control eye wash, 1:10; and immune eye wash, 1:100) was added in 100-µl volumes. IgG and IgA were probed using peroxidase-conjugated goat anti-rat IgA and IgG (Pharmingen, Sydney, Australia). Antibody present in samples was detected by adding color substrate tetramethyl benzidine, and the reaction was detected at 405 nm.

**Bacterial enumeration.** Clearance of *P. aeruginosa* from infected corneas was monitored by assessing the number of viable bacteria in whole eye homogenates at 4, 8, and 24 h and 3, 5, and 7 days postinfection. Small aliquots (20 µl in duplicate) of serial dilutions were plated onto nutrient agar plates. Plates were incubated for 18 h at 37°C. Results were expressed as the mean CFU/cornea ± standard errors of the means (SEMs).

**PMN quantitation.** Samples were assayed for myeloperoxidase (MPO) activity as previously described (13). Briefly, the whole eye collected at various time points (4, 8, and 24 h and 3, 5, and 7 days) was homogenized in 1 ml of hexadeyl trimethylammonium bromide (HTAB) buffer (0.5% HTAB in 50 mM phosphate buffer, pH 6.0) and sonicated for 10 s in an ice bath. The samples were freeze-thawed three times and centrifuged at 8000 × g for 20 min. Supernatant (0.1 ml) was mixed with 2.9 ml of 50 mM phosphate buffer (pH 6.0) containing 0.167 mg of O-dianisidine hydrochloride per ml and 0.0005% hydrogen peroxide. The change in absorbance at 460 nm was monitored continuously for 5 min in a spectrophotometer (Unicam; Selby Bioscience, Sydney, Australia). One unit of MPO activity was determined to be equivalent to approximately 2 × 10^5 PMNs/ml (4).

**Lymphocyte proliferation assay.** The lymphocyte proliferation assay was performed as described by Kyd et al. (21). Briefly, lymphocytes were obtained by passing mesenteric lymph nodes through a steel sieve and washing them in cold, sterile PBS supplemented with calcium, magnesium (CSL Biosciences, Sydney, Australia), 5% fetal calf serum, 100 µg of penicillin/ml, 100 µg of streptomycin/ml, and 0.25 µg of amphotericin B/ml (CSL Biosciences). Viable cells were counted by trypan blue exclusion. Cells were resuspended in culture medium RPMI 1640 (CSL Biosciences) containing HEPES (pH 7.2), 5 × 10^-5 M J-mercaptoethanol (ICN, Sydney, Australia), 2 mM L-glutamine, 5% fetal calf serum, and penicillin, streptomycin, and amphotericin B (as described above) at a final concentration of 10^5 cells/ml. Polyvalent antigen was diluted in culture medium in a 10-fold dilution series and filter sterilized. The cell suspension and antigen were cultured in triplicate in a final volume of 0.2 ml/well. Lymphocyte proliferation was determined by [³H]-thymidine incorporation (Amersham, Sydney, Australia) incorporation for the last 8 h of a 4-day culture by counting radioactivity in a scintillation counter. Results were calculated by subtraction of the background counts (radioactivity from the geometric means [counts] of triplicate wells).

**Histopathology of rat corneas.** Rats were sacrificed at 4, 8, 24 h and 3, 5, and 7 days postinfection using a 2.5% sodium pentobarbitone (Vetalar, Sydney, Australia) in 0.1 M sodium cacodylate (pH 7.4) at 4°C for 4 h. Fixed tissues were washed three times with PBS and dehydrated in graded ethanol (30, 50, 70, and 90%). Tissues were left at least 1 day in the infiltrating solution (90% ethanol and historesin at a 1:1 ratio) before they were embedded in Historesin Plus (Leica,
digested with T1 nuclease and proteinase K. Protected fragments were purified probe at 56°C overnight. After completion of hybridization, the samples were homogenizing the whole rat eye in sterile PBS. Homogenates were centrifuged at kits (R & D Systems, Minneapolis, Minn.). Samples for ELISA were prepared by circaed animals at different time points using commercially available ELISA measured in ocular homogenates of challenged eyes of immunized and nonim- software (Bio-Rad, Sydney, Australia). RNA was isolated using standard methods of phenol-chloroform extraction and ethanol precipitation from homogenized eyes. Concentration was detected by measuring the absorbance at 260 nm. Various cytokines were detected using a multiprobe RNase protection assay (Pharmingen). Briefly, a mixture of 32P-labeled antisense riboprobe was generated from a cytokine template. Total RNA isolated from whole rat eyes was hybridized with 32P-labeled ribo- probe at 56°C overnight. After completion of hybridization, the samples were digested with T1 nuclease and proteinase K. Protected fragments were purified by phenol-chloroform extraction followed by ethanol precipitation. Protected hybridized RNA samples were air dried and reconstituted in 2 μl of loading buffer, and the samples were resolved on a 4.5% polyacrylamide sequencing gel. After completion, the gel was transferred onto filter paper, dried, and exposed to X-ray film (Kodak X-omat; Sigma-Aldrich) overnight at −70°C. Film was then developed and bands were identified by comparing molecular weights to a cytokine template (rCK-1). Relative quantities were determined using Multi-analyst analysis of variance tests to assess the differences in cytokine gene and protein expression in the corneas of immunized and nonimmunized animals infected with P. aeruginosa. In addition, Pearson’s correlations were sought between bacterial clearance and/or PMN recruitment and the levels of cytokines. Mean differences were considered significant when P was ≤0.05.

**RESULTS**

Clinical Examination. (i) Nonimmunized animals. Control nonimmunized rats challenged with P. aeruginosa strain 6206 developed a predominantly edematous response at 24 h postchallenge. A single peripheral ring infiltrate covered 50 to 75% (grade 3) of the corneal diameter, and 75% (grade 3) of the stroma was involved, with moderate to severe density (grade 3.5). Ulceration involved up to 25% (grade 1) of the corneal epithelial thickness. The anterior chamber reaction was moderate, and there was moderate conjunctival redness. The composite corneal score for the severity of disease was 10.5 ± 2.1 (Fig. 1). At 7 days postchallenge, the severity (6.5 ± 1.2) of the disease was reduced.

(ii) Oral immunization. The corneas of 25 to 50% of the immunized animals were clear at 24 h postchallenge. Infected corneas showed complete or incomplete ring infiltrates at the periphery, with moderate densities (grade 3). Infiltrates involved 40 to 50% (grades 2 to 2.5) of the stromal thickness and 50% of the corneal diameter (grades 2 to 2.5), with overlying epithelial defects. There was a mild to moderate anterior chamber response and some hypopyon was seen. The composite score for the severity of the disease was 8.0 ± 1.5. At 7 days
postchallenge, the severity of the disease was reduced (5.2 ± 1.2) (Fig. 1).

(iii) Nasal immunization. At 24 h postchallenge, 75% of the animals showed clear, healthy corneas and infected animals showed a few focal and diffuse infiltrates and no epithelial defects. The composite score for the severity of disease was 5.5 ± 1.2. At 7 days postchallenge, the corneas of nasally immunized rats appeared normal (Fig. 1).

(iv) IPP immunization. The corneas of most IPP-immunized animals (50 to 75%) were normal 24 h after challenge with strain 6206. Infected corneas were edematous, a few focal stromal infiltrates covered 25% (grades 1.5 to 2.0) of the corneal diameter, and 40% of infected corneas had stromal involvement (grades 2 to 2.5) with mild densities (grades 2 to 2.5). There was no epithelial defect present. In these animals an anterior chamber examination revealed a fibrinous reaction (grades 2 to 3). The composite score for the severity of disease was 6.5 ± 1.5. At 7 days postchallenge, the corneas appeared normal (Fig. 1).

Histological examination. (i) Nonimmunized (control) animals. There was massive PMN infiltration streaming from the limbus and conjunctiva to the mid-periphery (densely packed) of the corneal stroma and fewer PMNs in the central cornea at 24 h postchallenge with strain 6206 in nonimmunized animals. The PMNs were lined up at the Descemet’s membrane. Bacteria could be seen at the wound site and throughout the stroma. The epithelial defect was present (Fig. 2). At 7 days postchallenge, the infiltrates were still present in diffuse and focal patches and bacteria could not be seen in the corneal stroma.

(ii) Oral immunization. The corneas of immunized rats that developed infection (50 to 75%) after challenge with strain 6206 showed PMN infiltration, with PMN streaming from the limbus to the periphery of the corneal stroma. A moderate epithelial defect was present (Fig. 2). At 7 days postchallenge, the infiltrates were still present in diffuse and focal patches and bacteria could not be seen in the corneal stroma.

(iii) Nasal immunization. The corneas of intranasally immunized rats showed diffuse infiltration throughout the corneal stroma. The epithelium was intact (Fig. 2). At 7 days postchallenge, the corneal histology appeared normal.

(iv) IPP immunization. Immunized animals (25 to 50%) challenged with strain 6206 showed focal patches of infiltration in the stroma at 24 h postchallenge. Infected corneas were
edematous, and no epithelial defect was present (Fig. 2). At 7 days postchallenge, very few infiltrates were seen in the corneal stroma.

Evidence for the presence of antigen-specific antibody in tear fluid and serum of immunized rats. The antibody response following immunization was measured by ELISA. Antigen-specific IgA antibodies in tears measured with ELISA. (A) Antigen-specific IgA antibodies in tears measured by ELISA. (B) Antigen-specific IgG antibodies in serum. Each value represents the mean ± SEM for samples from three rats in each group. The ELISA titer corresponds to the absorbance at 405 nm. Mean differences were considered significant (+) when \( P < 0.05 \). OI, orally immunized; NI, nasally immunized; IPP, IPP immunized.

Evidence for rapid bacterial clearance in immunized groups. Viable counts of the infected eye from immunized and nonimmunized animals were performed at 4 h postchallenge and continued for up to 7 days. All immunized groups showed rapid clearance of bacteria. Significantly lower numbers of bacterial cells were present in nasally \( (P = 0.03) \), IPP- \( (P = 0.045) \), and orally \( (P = 0.048) \) immunized animals at 24 h postchallenge than in nonimmunized animals. Bacteria could not be recovered from nasally immunized groups by day 3, and by day 5 all immunized groups lacked recoverable bacteria. Bacterial cells could not be cultured from clinically clear corneas of IPP- orally, and nasally immunized rats (Fig. 4).

Effect of immunization on PMN infiltration. The MPO activity in experimental groups was calculated by subtracting the MPO activity of the normal eye (21 ± 3.05). Comparison of MPO activity in immunized and nonimmunized animals showed significantly higher levels \( (P < 0.05) \) in all three immunized groups at 4 h post challenge which were significantly diminished \( (P < 0.05) \) at 24 h post challenge compared to nonimmunized rats. The levels of PMNs in nonimmunized rats peaked at 24 h postchallenge and remained elevated for up to 7 days (Fig. 5).

Enhanced antigen-specific lymphocyte proliferation in immunized animals. Lymphocytes isolated from mesenteric lymph nodes from immunized and nonimmunized rats were cultured with killed bacteria (at 1:10 and 1:100 antigen dilutions) to assess the levels of antigen-specific lymphocyte responses. Antigen-specific proliferation was significantly higher \( (1:100 \text{ dilution}, P < 0.0001) \) in immunized groups than in nonimmunized rats. Lymphocytes isolated from nasally and IPP-immunized animals showed significantly higher \( (P < 0.001) \) proliferation in the presence of killed bacteria than in orally immunized animals (Fig. 6).

Differential profile of cytokine mRNA expression in immunized groups. The rCK-1 template with multiple probes (IL-1\(\beta\), IL-4, IL-5, IL-6, IL-2, and IL-10) was used to detect mRNA in immunized and nonimmunized groups. Immunized groups...
showed differential mRNA expression compared to the nonimmunized control group.

(i) Nonimmunized (control) rats. Various cytokines were present in the corneas of immunized and control rats infected with strain 6206. Transcripts of IL-1β and IL-4 were highly upregulated, while IL-6 was upregulated to a lesser extent, at 24 h postchallenge compared to immunized groups. IL-10 was present in significantly lower \( (P < 0.0003) \) levels than in immunized groups. Transcripts of IL-2 and IL-5 were not detected at any time points.

(ii) Oral immunization. There was upregulation of IL-1β mRNA expression at 24 h postchallenge compared to other immunized (nasal and IPP) groups. IL-4 and IL-10 mRNA showed similar expression patterns to those of other immunized animals. Similar to controls, IL-2 and IL-5 mRNAs were not detected.

(iii) Nasal immunization. The expression profile of cytokine mRNA in nasally immunized rats differed from those of both nonimmunized and orally immunized animals. Transcripts of IL-2, IL-5, and IL-10 were upregulated, while IL-1β and IL-4 mRNA were expressed at significantly lower \( (IL-1β\), \( P < 0.0002\); IL-4, \( P < 0.003) \) levels at 24 h postchallenge than in control nonimmunized animals. Expression of IL-6 mRNA was below the detection limit at any time point.

(iv) IPP immunization. Rats immunized through IPP had a similar pattern of cytokine mRNA expression to those that were immunized nasally, except for IL-6 expression. IPP-immunized rats showed increased expression of IL-2, IL-5, and IL-10 mRNA and decreased expression of IL-1β, IL-4, and IL-6 at 24 h postchallenge compared to controls. Unlike nasally immunized rats, IPP-immunized animals expressed both IL-5 and IL-6 mRNA (Fig. 7).

Effect of immunization on cytokine protein secretion. The protein levels were not determined for all cytokines (those probed for mRNA) due to the limited availability of reagents for rats.

(i) Nonimmunized controls. In nonimmunized rats, neutrophil chemoattractant CINC-1 protein levels were significantly lower \( (P < 0.04) \) early (4 h) during the infection and were significantly higher \( (P < 0.03) \) later (24 h) during the infection than those of immunized rats. Expression of CINC-1 protein remained high up to 7 days postinfection. The amount of IL-1β protein gradually increased and peaked at 24 h postchallenge and remained high up to 7 days postinfection. IL-6 protein also peaked at 24 h, diminished drastically at 3 days postinfection, and remained low up to 7 days postchallenge. Nonimmunized rats showed high levels of IL-4 protein which peaked at 24 h \( (P < 0.03) \) postchallenge and remained high up to 7 days postinfection.

(ii) Oral immunization. Expression of CINC-1 protein was significantly higher \( (P < 0.04) \) early during the infection (4 and 8 h) and significantly lower \( (P = 0.013) \) by 24 h postinfection than that of nonimmunized control rats. IL-1β protein levels were low throughout the period of infection compared to those of nonimmunized rats. The levels of IL-6 protein were significantly higher \( (P < 0.033) \) in orally immunized rats at 4 h postchallenge than those of nonimmunized rats. The pattern of protein expression was reversed at 8 h postinfection, with IL-6 protein levels increasing dramatically in nonimmunized animals. IL-10 and IL-4 proteins showed a biphasic pattern, with the first peak appearing at 4 to 8 h and the second at 3 days postinfection.

(iii) Nasal immunization. The protein secretion pattern of CINC-1 and IL-1β was similar to those of orally and IPP-immunized rats. IL-6 protein was below the limit of detection. IL-4 and IL-10 proteins were present late during the infection. IL-2 protein was present at most time points but at very low levels.

(iv) IPP immunization. The pattern of CINC-1 protein secretion was the same as in orally or nasally immunized rats. IL-1β and IL-6 levels were low throughout the period of infection compared to nonimmunized rats, except for the levels of IL-6 at 4 h \( (P < 0.033) \). IL-4 protein was highly upregulated at 24 h postinfection and diminished thereafter. Unlike in nasally immunized rats, IL-10 protein showed a biphasic pattern peaking very early (4 to 8 h) and late (5 days) during the infection (Fig. 8). IL-2 protein was present at all time points at very low levels.
DISCUSSION

Our study showed that the route of immunization affects the severity and persistence of microbial keratitis. Immunization has the potential to modulate the inflammatory response to an infection. This modulation includes the production of chemical signals, cytokines and chemokines, with recruitment and activation of cells involved in clearing the infection (29). This study has demonstrated that immunization changes the kinetics of PMN infiltration, with immune groups having more rapid recruitment and resolution of PMNs in the cornea than the nonimmune group. Associated with this was a more rapid clearance of bacteria, differences in the levels of cytokines expressed and produced, and reduced adverse pathology. In particular, the IPP and intranasal immunization regimes with an OT boost provided the best protection from corneal ulceration.

CINC-1 is a potent activator and attractant of neutrophils (27). Increased CINC-1 levels were detected earlier (4 to 8 h) postinfection in immunized rats than in nonimmunized rats, with all groups peaking at 24 h postchallenge. However, despite the earlier increased production of CINC-1 in the immunized groups, the peak levels of CINC-1 were significantly lower in the immunized groups and also decreased far more rapidly. The changes in the CINC-1 levels corresponded to the recruitment and resolution profiles of the PMNs. The rate of PMN recruitment in other disease settings has been associated with early bacterial clearance, such as enhanced respiratory clearance of nontypeable Haemophilus influenzae following mucosal immunization (5, 10). Persistence of PMNs in the nonimmune animals during the later stages of infection may contribute to corneal scarring and perforation.

For the PMN response to be beneficial rather than detrimental, a rapid resolution of PMN infiltration must occur. Immunization of rats against P. aeruginosa corneal infection achieved a rapid resolution of PMN infiltrates. In addition to the modulation of CINC-1 levels, there were reduced levels of the proinflammatory cytokines (IL-1ß and IL-6) and similar or higher levels of the cytokines associated with immunosuppres-
sive or IgA antibody responses (IL-10 and IL-4). Balanced expression of proinflammatory and anti-inflammatory cytokines in the immunized animals compared to the overwhelming proinflammatory cytokine response in the nonimmune group may control inflammation by regulating not only inflammatory cell recruitment but also IgA secretion.

Clearance of bacteria from the ocular surface is presumed to involve the combined actions of PMNs and secretory IgA. Immunization induced significantly elevated levels of antigen-specific IgA in tears and IgG in serum. The role of antigen-specific antibodies in protection against corneal infection is controversial, with correlation between the presence of antibody and protection not always being clearly defined. A recent study has shown that secretory IgA can significantly inhibit P. aeruginosa binding to wounded mouse cornea in vitro, thereby protecting against keratitis. One of the mechanisms by which IgA antibodies may prevent bacterial colonization is by specifically interacting with bacterial adhesins required for binding to mucosal tissue (24). IgA is capable of potentiating the function of innate antibacterial factors and interacting with mucosal phagocytic cells and lymphocytes (25). Oral immunization with Acanthamoeba spp. antigen mixed with cholera

FIG. 8. Kinetics and levels of cytokine protein secretion quantified by ELISA in whole rat eyes of nonimmunized (C) and immunized rats challenged with cytotoxic P. aeruginosa strain 6206. The results are presented as means ± SEMs from three experiments, with three animals in each group at each time point. Mean differences were considered significant (*) when P was ≤0.05.
toxin induces the production of parasite-specific IgA in mucosal secretions and prevents corneal infection (23). Although antigen-specific IgG antibody appears to be important for opsonophagocytosis (34), a correlation between the presence of opsonizing antibodies and protection in vivo has not been clearly determined to be an essential mechanism of effective immunity (33, 35). There is also evidence that suggests that systemically derived IgG may also be capable of conferring protection in the cornea (28). In addition to measuring significant titers of antigen-specific IgA in tears, we have demonstrated the presence of a group of IgA-enhancing Th2-type cytokines (IL-4, IL-5, IL-6, and IL-10) which may provide an environment for preferential immunoglobulin class switching for IgA in the eye.

Previous studies using a rat model for pulmonary *P. aeruginosa* infection have shown that mucosal immunization significantly alters the profile of inflammatory cytokines produced in response to infection (5). Other evidence also suggests that nasal and IPP immunization with mucosal adjuvant induces dominant Th2 responses in nasal-associated lymphoid tissue and Peyer’s patches (12, 39). This study has shown that the route of immunization changes the profile of cytokine expression during *P. aeruginosa* corneal infection, with the most significant differences appearing in the nasal and IPP immunization groups. Expression of IL-2 and IL-5 were especially altered, with nasally immunized rats expressing high levels of IL-5 and baseline levels of IL-6 mRNA, with corresponding baseline levels of IL-6 protein. In contrast, orally immunized rats showed no IL-5 expression but had high IL-6 expression and secretion, while IPP immunization resulted in the upregulation of both IL-5 and IL-6. IL-5 and IL-6 are known to differentially influence the B-1 and B-2 lineage of plasma cells (2). Collectively, the data suggest that nasally immunized animals may be producing IgA plasma cells of B-1 lineage, which are IL-5 dependent and IL-6 independent (2), whereas orally immunized animals may be producing predominantly cells of B-2 lineage. B-1 cells are physically and functionally unique B cells producing antibodies to bacterial antigens such as lipopolysaccharide and phospholipase C (1). B-1 cells mainly reside in mucosal effector tissues, while conventional IgA+ B-2 cells reside in mucosal inductive sites (39). Nasal-associated lymphoid tissue functions as a primary inductive site for IgA antibody in tears by contributing triggered IgA-committed B cells to the lacrimal gland (22). A recent study (30) has shown that a high frequency of IgA-committed B-1 cells occurs in the lacrimal gland (an effector site).

A role for T cells and cytokines produced by activated T cells in protection from ocular bacterial infections has not been demonstrated previously. Nasally and IPP-immunized rats induced antigen-specific lymphocyte responses, providing evidence that an antigen-specific T-lymphocyte response was induced by immunization and that these lymphocytes migrated from the site of immunization. Immunologically specific T cells recruit neutrophils in an antigen-dependent and dose-dependent fashion (6). Cytokines released by activated T cells may direct the activity of nonspecific effector cells (21, 37). All of these studies have shown the involvement of T cells and cytokines in respiratory disease models. Evidence that supports the relevance of a CD4+ Th1- versus Th2-type immune response was presented in a study that used a mouse *P. aeruginosa* keratitis model. Data from this study suggest that Th2-responsive mice regulate inflammatory cellular infiltration more efficiently by downregulating the inflammatory response, which in turn results in less corneal stromal damage (11, 20). Further studies are required to define the importance of a T-cell response in protection against ocular infection.

This study has demonstrated that the immunization route modulates the inflammatory response to ocular *P. aeruginosa* infection, thus affecting the severity of keratitis and adverse pathology. The results show that immunization affects the rate of bacterial clearance and alters the profile of cytokines produced in response to ocular infection, with nasal immunization resulting in the most significant level of protection. The results suggest that the degree of protection afforded by immunization may depend upon the rapid recruitment of PMNs, the induction of antigen-specific IgA, and the balanced production of proinflammatory and immunosuppressive cytokines and that T-cell responses may influence these events.

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**REFERENCES**


